Effects of soft interactions and bound mobility on diffusion in crowded environments: a model of sticky and slippery obstacles

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Abstract. Crowded environments modify the diffusion of macromolecules, generally slowing their movement and inducing transient anomalous subdiffusion. The presence of obstacles also modifies the kinetics and equilibrium behavior of tracers. While previous theoretical studies of particle diffusion have typically assumed either impenetrable obstacles or binding interactions that immobilize the particle, in many cellular contexts bound particles remain mobile. Examples include membrane proteins or lipids with some entry and diffusion within lipid domains and proteins that can enter into membraneless organelles or compartments such as the nucleolus. Using a lattice model, we studied the diffusive movement of tracer particles which bind to soft obstacles, allowing tracers and obstacles to occupy the same lattice site. For sticky obstacles, bound tracer particles are immobile, while for slippery obstacles, bound tracers can hop without penalty to adjacent obstacles. In both models, binding significantly alters tracer motion. The type and degree of motion while bound is a key determinant of the tracer mobility: slippery obstacles can allow nearly unhindered diffusion, even at high obstacle filling fraction. To mimic compartmentalization in a cell, we examined how obstacle size and a range of bound diffusion coefficients affect tracer dynamics. The behavior of the model is similar in two and three spatial dimensions. Our work has implications for protein movement and interactions within cells.

1 Introduction

The diffusion of macromolecules in crowded environments is generally slowed relative to the uncrowded case, and particle motion can undergo transign anomalous subdiffusion [1]. The motion of lipids or macromolecules within biological membranes can be affected by crowding [2–9], because the membrane contains both macromolecules and inhomogeneities in membrane composition [10, 11]. In the cell interior, macromolecules, organelles and other cellular structures can inhibit motion, or in contrast, enhance sampling of non-crowded regions [12]. Biological crowders can also contain interaction sites which further modify the macromolecular motion [13]. The kinetics and equilibrium behavior of interactions between mobile proteins can be modified by crowding [14, 15]. The magnitude of the effects of crowding on macromolecular motion and reactions is important to determine the limiting rate of biological processes such as signaling receptor activation.

Because of its biological importance, the effects of crowding on diffusion and macromolecular interactions have seen significant experimental and theoretical work [16, 17]. Lattice gas models have been used to demonstrate the effects of crowding [18, 19], binding [20, 21], and repulsion [22] on the diffusion of tracer particles. These effects—including transient anomalous diffusion at short times and hindered normal diffusion at long times—have been studied for both immobile [23] and mobile obstacles [24, 25].

Although most theoretical work has focused on anomalous diffusion in crowded systems made up of impenetrable obstacles with attractive or repulsive surfaces [18–20, 22, 26], there is growing evidence of the importance of soft compartments and barriers in biological systems. In membranes, lipids can be only partially excluded from lipid rafts or domains. When they do interact, they can still diffuse within them [27–30]. Lipid motion can be hindered, though not stopped, near α -synuclein protein aggregates [31]. For all of these cases, theoretical considerations of a twodimensional system should include the effects of the soft interaction potentials and bound-state mobility.

Inside the cell, intrinsically disordered or lowcomplexity domains can act as soft obstacles or wells, with rapid diffusion within the wells. Membrane-less organelles spontaneously form from low-complexity domain proteins. They are typically highly dynamic assemblies [32], which show fast intra-particle diffusion times, and allow rapid entry and exit of constituents Proteins which interact with intrinsically [33]. disordered proteins can still diffuse during the binding interaction [34, 35]. This effect may be particularly pronounced in the central channel of the nuclear pore complex, which contains a high density of binding sites on intrinsically disordered domains. Recent simulation work suggests that the disordered protein binding pockets can exchange on transport factors [35], providing a clear mechanism for mobility while bound to an obstacle. Particles are weakly excluded from individual disordered protein chains due to the lowering of the polymer chain entropy [36], but are expected to allow other macromolecules to enter, and pass through, the space that they occupy. The increasingly recognized importance of proteins which are intrinsically disordered or contain low-complexity domains within their assemblies warrants a more careful consideration of the differences between the previously well-studied models, in which binding immobilizes the bound species, and a model which includes soft interactions and obstacles or barriers in which the bound species may remain mobile.

Motivated by the biological importance of binding interactions which can retain mobility of the bound particle, we studied a minimal model with bound tracer mobility (figure 1). In our model, tracer particles move on a 2D or 3D lattice in the presence of immobile obstacles, to which the tracers can bind. A primary distinction between our model and many others that consider binding or adhesion is that others typically consider adhesion between a tracer and an adjacent hard obstacle, in which there is no overlap between tracers and a hard obstacles core [20, 22, 24]. Here, obstacles are soft: tracer particles can overlap with obstacles, with an energy penalty (or gain) ΔG upon moving to a lattice site occupied by an obstacle. Unlike previous work modeling lipid rafts, we closely examine the dependence on binding, instead of just pure exclusion or free entry into lipid regions [4].

To understand the effects of bound mobility, we first consider the limits of 'sticky obstacles', in which tracers are immobile while bound, and 'slippery obstacles', in which tracers are mobile while bound. We use lattice Monte Carlo methods to explore a range binding energy and obstacle filling fraction. We also examine the effects of semi-sticky obstacles—i.e., intermediate bound diffusion coefficient—and obstacle



Figure 1. Model schematic. Tracers (colored circles) hop on a lattice of empty sites (white squares) and obstacles (gray squares). Tracer binding with a soft interaction potential allows them to overlap with obstacles (top). For sticky obstacles, the only allowed moves of a bound tracer are to empty sites (unbinding). For slippery obstacles, tracers can hop to other obstacles while remaining bound. Arrows denote possible moves and P the probability that a given move is accepted.

size effects, which demonstrates how diffusion is altered in a crowded environment with compartments with different properties—such as a cell [21]—is altered. Our results demonstrate how binding and bound-state motion independently impact particle dynamics, including long-time normal diffusion and anomalous diffusion. Bound tracer mobility increases the long-time diffusion coefficient, reduces the transient anomalous time, and eliminates caging for all times typically observed above the percolation threshold. These results demonstrate that mobility of bound particles can benefit biological systems by allowing mobility even in highly crowded environments.

2 Model

Our model seeks to build on stochastic lattice-gas models that have been important to understanding tracer dynamics in the presence of immobile and mobile hard obstacles [18], anomalous subdiffusion [19], and effects of binding on diffusion [20]. Saxton showed that the tracer diffusion coefficient drops to zero at the percolation threshold, the critical concentration of obstacles at which a continuous path of vacancies through which a tracer can move no longer exists. Above this percolation threshold, diffusion is anomalous at long times. The effects of tracer and obstacle size [26, 37, 38] and adhesion and repulsion to sites adjacent to obstacles [22] on transient subdiffusion and long-time diffusion have been studied. Extensions



Figure 2. Mean-squared displacement $\langle r^2 \rangle$ divided by time delay t as a function of time delay t for (a) impenetrable obstacles, (b) repulsive slippery obstacles ($\Delta G = 2$), (c) repulsive sticky obstacles ($\Delta G = 2$), and (d) attractive sticky obstacles ($\Delta G = -2$). Different colors correspond to different filling fraction ν . Curves with non-zero slope indicate anomalous diffusion, and the horizontal asymptote indicates the long-time diffusion coefficient. Each curve represents an average over tracers, independent time windows, and obstacle configurations.

to mobile obstacles which interact with each other have demonstrated how obstacle clustering dynamics can influence the diffusivity of tracers [39]. Numerically exact methods for calculating diffusion coefficients using the Nernst-Einstein relation [40, 41] and Markov chains [42] have been implemented as a different approach to analyzing these systems; the Nernst-Einstein approach can lower the computational cost of measuring diffusion coefficients for lattice gases [37]. Protein motion in polymer networks has been studied using random-walk and self-avoiding-chain models for immobile [43] and mobile [24, 44] hard chains. Studies of chains with binding sites found that modeling chain dynamics allowed a mapping onto randomly distributed obstacles with an effective volume, and showed how sliding along a defined chain can effect tracer dynamics [44, 45]. In some previous work, the effects of binding and sliding while bound were entangled because both effects were encoded by a single parameter [44, 45]. Domains with different diffusion coefficients and sizes—to model lipid rafts—have been studied, but the analysis only included total or no exclusion although it was noted that binding effects could play a large role [4].

In our model, tracer particles undergo a random walk on a square lattice and interact with immobile obstacles. The interaction is characterized by a binding free energy; for simplicity, we neglect any additional



Figure 3. Top panel: Illustration of fitting procedure, showing $\langle r^2 \rangle / t$ vs time delay t for simulation data (blue), line fitted to horizontal asymptote (red dashes), line tangent to point of maximum absolute slope of the curve (red dash-dots), and anomalous time t_a (black dots) for different parameters. Bottom panel: instantaneous scaling exponent α vs time delay t. (a, d) Slippery obstacles with $\Delta G = 1$, $\nu = 0.95$: normal diffusion occurs for all measured time. (b, e) Sticky obstacles with $\Delta G = 2$, $\nu = 0.45$. (c, f) Sticky obstacles with $\Delta G = -5$, $\nu = 0.50$.

activation barrier. The characteristic binding free energy change of a tracer that hops from an empty site to an obstacle site is ΔG (in units where $k_BT = 1$). We consider both attractive ($\Delta G < 0$) and repulsive ($\Delta G > 0$) obstacles. We use the Metropolis algorithm [46] to accept or reject candidate binding (probability P_B) and unbinding (probability P_U) events. Each tracer occupies a single site lattice site, but the obstacle size is varied to represent domains of characteristic size. Obstacles are squares with sides of length l_{obst} , measured in units of the lattice spacing.

To study the effects of tracer particle motion while bound, we considered the limits of perfectly sticky and slippery obstacles (fig. 1), as well as the intermediate 'semi-slippery' case. In all models, obstacles are soft, so that tracers overlap with obstacles when bound. For sticky obstacles, no hopping between obstacle sites can occur, but tracers can exit an obstacle in any direction that would move the tracer to an unoccupied site. For slippery obstacles, tracers can hop between adjoining obstacles while remaining bound. In the limit of perfectly slippery obstacles, in which bound motion is identical to unbound motion, there is no difference in hopping rates between free and bound tracers. For semi-slippery obstacles, we vary the bound diffusion coefficient.

2.1 Simulation methods

In our kinetic Monte Carlo scheme, at each time step a tracer attempts a move in a randomly chosen direction. Moves from **empty** \rightarrow **empty** are always accepted, **empty** \rightarrow **obstacle** moves are accepted with probability min $(e^{-\Delta G}, 1)$, **obstacle** \rightarrow **empty** moves are accepted with probability min $(e^{\Delta G}, 1)$, and **obstacle** \rightarrow **obstacle** moves are always accepted/rejected if obstacles are slippery/sticky (fig. 1); for semi-slippery obstacles, the acceptance probability is $D_{\text{bound}}/D_{\text{free}}$. If a tracer's move is rejected, it remains immobile for that time step. We assume noninteracting tracers.

Initially, obstacles were uniformly randomly placed on the lattice, at the specified filling fraction, without overlaps. Next, tracers were randomly placed on obstacles and empty sites at their equilibrium occupancy, as determined by the filling fraction of obstacles ν , and binding energy ΔG . The fraction of tracers on obstacles is proportional to the obstacle filling fraction times the Boltzmann factor, $\nu e^{-\Delta G}$, while the fraction of tracers on empty sites is proportional to the fraction of empty sites, $(1 - \nu)$. The equilibrium fraction of tracers on obstacles of size 1 is then

$$f_o = \frac{\nu e^{-\Delta G}}{\nu e^{-\Delta G} + (1 - \nu)}.$$
(1)

Using an initial fraction of tracers bound to obstacles determined from f_o avoids the time required for binding equilibration in the simulations, ensuring that mean-squared displacement measurements are independent of a time origin.

We performed 2D simulations with 200 tracers on a 256 × 256 periodic lattice for $10^5 - 10^{7.5}$ time steps, with recording interval of 10 - 100 steps. For each parameter set (determined by filling fraction and binding energy), we averaged over 96 separate obstacle configurations. We varied ν from 0 to 1 and ΔG from -5 to 10. 3D simulations used similar parameters with a 256 × 256 × 256 periodic lattice. In the semi-slippery case, we varied the ratio of bound to free diffusion coefficient $D_{\text{bound}}/D_{\text{free}}$ between 0 (perfectly sticky) and 1 (perfectly slippery) in steps of 0.2, for binding energies $\Delta G = 1, 2, 3, \infty$ for two filling fractions, $\nu = 0.3$ and 0.6. When varying obstacle size, we used square obstacles with the length of a side, l_{obst} , equal to odd values from 1 to 15.

2.2 Trajectory analysis

We determined tracer mean-squared displacement (MSD) as a function of time delay by averaging over all tracers, 100 randomly selected independent time origins, and obstacle configuration. For long time delays for which 100 independent time intervals were not available, we averaged over the maximum number of independent time intervals. As previously mentioned, averaging over time windows improves our statistics; note that the time origins are not unique, since the placement of tracers in their equilibrium binding distribution ensures that there is no initial binding equilibration time. We have verified that that are no aging effects [47, 48], *i.e.*, MSD measurements that depends on simulation time, in our model (data not shown).

We sought to quantify the effects of binding and obstacle filling fraction on tracer mobility. In systems with either purely Fickian diffusion or particular obstacle geometry, the mean-squared displacement grows as a power law in time:

$$\langle r^2 \rangle = 2dDt^{\alpha},\tag{2}$$

where $\langle r^2 \rangle$ is the ensemble-, time-origin-, and obstacleconfiguration-averaged mean-squared displacement, dis the spatial dimension, D is the diffusion coefficient, α is the diffusion scaling exponent, and t is the time delay. This fractional diffusion equation has been studied extensively [49], both because it emerges from certain microscopic theories and as a means to quantify anomalous random walks. Fractional diffusion has been experimentally measured in cells, using fluorescence recovery after photobleaching [50], fluorescence correlation spectroscopy [51], and singleparticle tracking [52]. For hard obstacles, α reflects the non-homogeneity and fractal structure of a cluster. In this case, α can be thought of as a measure of a local landscape, in which obstacles have the possibility of trapping a tracer and introducing memory effects into the system. The value of α does not quantify the time it takes to escape a trapping cage; but $\alpha < 1$ suggests the possibility that the landscape can cage tracers. In the $\alpha \rightarrow 0$ limit, a tracer is fully caged, and the $\alpha \rightarrow 1$ limit represents Fickian diffusion.

However, many systems have more complex dynamics that are not power law. For example, tracer dynamics can be transiently anomalous: subdiffusive on short time scales and Fickian on longer time scales (fig. 2b). The dynamics can be quantified using a phenomenological approximation in which the exponent α is treated as time dependent [19,26,37,38, 42,53]. Thus, $r^2 \sim t^{\alpha}$ holds only over particular time scales.

For non-power-law dynamics, we can apply equation 2 locally, with a phenomenological, time-varying exponent. Then $\alpha(t)$ is defined by local fitting to the the logarithm of $\frac{\langle r^2 \rangle}{t}$:

$$\log\left(\frac{\langle r^2 \rangle}{t}\right) = \log\left(2dD\right) + (\alpha(t) - 1)\log\left(t\right).$$
(3)

so that $\alpha(t) - 1$ is the local slope of the $\frac{\langle r^2 \rangle}{t}$ versus t curve on a log-log plot. As seen in figs. 2 and 3, the instantaneous effective α varies with delay time. Thus, a power-law MSD scaling with time, such as can arise from fractional Brownian dynamics, does not encompass the complexity of our crowded diffusion model, as has been found previously [8, 38].

At short time, our model typically exhibits anomalous diffusion. However, in some conditions, the short-time behavior is diffusive, with an intermediate anomalous regime. We defined α_{\min} as the minimum instantaneous value of α (the most anomalous exponent). We characterized the transition between short- or intermediate-time anomalous diffusion and long-time normal diffusion by the time scale t_a , determined as the intersection of the horizontal longtime asymptote of $\frac{\langle r^2 \rangle}{t}$ with a line tangent to the point of the maximum rate of decrease of this curve (fig. 3b,c). We found that this transition time could be robustly determined for a wide range of diffusion coefficients and anomalous behavior. We denote t_a the anomalous time. Qualitatively, it is the crossover time from short-time subdiffusion to long-time Fickian diffusion. While α_{\min} characterizes how trapped a tracer is, t_a quantifies how long it takes a tracer to escape a cage.



Figure 4. Sticky obstacles of size 1 in 2D. (a, d) Diffusion coefficient D^* , (b, e) anomalous time t_a , and (c, f) minimum scaling exponent α_{\min} as a function of obstacle filling fraction ν for positive (top) and attractive (lower) binding energy. Note that points for $\Delta G = 10$ are partially hidden behind $\Delta G = \infty$. The approximate locations of the critical occupancies ν^l and ν^u are indicated with gray dotted lines.

We defined the long-time Fickian diffusion coefficient as

$$D = \lim_{t \to \infty} \frac{\langle r^2 \rangle}{2dt}.$$
(4)

All diffusion coefficient measurements are expressed in terms of the scaled diffusion coefficient $D^* = \frac{D}{D_0}$, where $D_0 = \frac{l^2}{2d\tau}$ is the diffusion coefficient in the absence of obstacles, where l is the distance between lattice sites (here defined to be 1), and τ the time interval between steps (also set to 1).

In some cases, we were unable to determine all of D^* , α_{\min} , and t_a . For some parameter sets, the slope of $\frac{r^2}{t}$ vs. t on a log-log plot approached a nonzero constant, indicating that diffusion was anomalous over all measured time delays, so that the Fickian diffusion coefficient was not well-defined. For other parameter sets, the $\frac{r^2}{t}$ versus t curve did not reached a clear asymptote during the simulation time. We therefore could not determine D^* , but could measure α_{\min} . When tracer diffusion was normal over all or nearly all measured time delays, neither α_{\min} nor t_a were well-defined, but D^* could be measured.

3 Sticky soft obstacles

We initially focus on the limit of perfectly sticky obstacles of size 1, to determine the effects of stickiness, filling fraction, and binding energy on tracer motion. We varied parameters over a wider range for the 2D model, with a comparison to 3D results for some parameter sets.



Figure 5. The two types of percolation threshold in our lattice model: the lower critical occupancy ν^{l} (left) and the upper critical occupancy ν^{u} (right). For the lower critical occupancy, which is the standard percolation threshold, the percolating network is the obstacles. At the upper critical occupancy, the percolating network is the *interface* between two or more obstacles. The barrier to tracer motion is shown as a black lines; obstacle-obstacle boundaries which cannot be crossed by a tracer in the sticky model are shown in red. Without binding, tracers cannot pass through the lower percolating network. If they can bind, tracers can 'hop through' the lower percolation barrier with or without bound motion.

For sticky obstacles, the motion of a bound tracer to an adjacent obstacle is prohibited. This could occur, for example, because the net free energy cost of binding to an obstacle is a result of an attractive binding interaction, with a high free energy barrier to moving to an adjacent site. Here, we consider the limit that the free energy cost of moving to an adjacent obstacle is so large that it approaches infinity. This situation provides an important point of comparison to explicitly test the effects of bound-state diffusion on tracer behavior.

We separately consider repulsive and attractive obstacles (fig. 4). Note that we include the case $\Delta G = 0$, that is, where the binding interactions are neither attractive nor repulsive, but still block moves to adjacent obstacles. We define the lower critical occupancy ν^l as the filling fraction at which the diffusion is non-Fickian for all time scales for impenetrable obstacles ($\Delta G = \infty$). In the limit of a hard repulsive obstacle, D^* decreases with filling fraction, and approaches zero at the percolation threshold expected for hard obstacles on a square lattice, $\nu^l \approx 0.4$ [54], where t_a diverges [18]. The lower critical occupancy is the percolation threshold, at which there is no longer a continuous path of empty sites (fig. 5).

For finite binding free energy in our model, Fickian diffusion can still occur above the percolation threshold ν^l because soft binding allows tracers to 'hop through' single obstacles via binding and unbinding. Without soft binding of the type we consider, obstacle percolation would prevent a tracer from moving between vacancy clusters. In other words, tracers that start in an area caged by obstacles are stuck there. With soft binding, tracers that start in a cage can hop onto an obstacle and then hop off into a new vacancy cluster. For soft binding interactions and sticky obstacles, there is an upper critical occupancy $\nu^{u} \approx 0.72$ at which the long-time diffusion coefficient approaches zero irrespective of binding energy (fig. 4). Above ν^{u} , tracers become caged regardless of the binding kinetics. Therefore, there is a different type of percolating network above the upper critical occupancy: the percolation of the inter-obstacle boundary (fig. 5). At the upper critical occupancy, there is a second adjacent obstacle preventing the tracer from 'hopping through.' Note that as expected, the transition time t_a appears to diverge on the approach to the upper critical occupancy (fig. 4). We are unaware of a theoretical value for this percolating density, but our results suggest its approximate value is 0.72 in 2D (fig. 4).

Intermediate repulsive binding energy leads to intermediate behavior, as expected. For strong repulsion, e.g., $\Delta G = 5$, D^* remains small, though

clearly non-zero, up to the upper critical occupancy, while t_a monotonically increases until it diverges at ν^u .

Anomalous dynamics appear in the slope of $\langle r^2 \rangle / t$ on a log-log plot. The most anomalous behavior occurs when the scaling coefficient α reaches its smallest value, α_{\min} . We find that α_{\min} decreases with filling fraction and binding energy (fig. 4c). Adding more obstacles and increasing the repulsion causes greater hindrance of tracer motion. We note that $\alpha_{\min} \approx 0.7$ near ν^l for impenetrable obstacles, as found previously [4, 19]. Finite repulsive binding energy leads to a smaller exponent ($\alpha_{\min} < 0.7$) than the infinite case at filling fraction above ν^l . For lower values of ΔG , the scaling coefficient does not go to zero at the upper critical occupancy ν^u . Note that the sharp cutoff with filling fraction occurs because we did not collect data past ν^u .

Sticky obstacles with attractive binding interactions show a more rapid falloff in the diffusion coefficient and larger anomalous time (fig. 4). The upper critical density $\nu^u \approx 0.72$ is in the same vicinity as for $\Delta G > 0$. The dependencies of the diffusion coefficient on filling fraction for positive and negative binding energy are similar for low magnitude of the binding energy, but the diffusion coefficient falls off more rapidly with filling fraction for highly attractive obstacles. This occurs because an attractive obstacle confines a tracer in one position until it escapes, while a repulsive obstacle only impedes tracer motion for one time step. Therefore, repulsive interactions require several obstacles to transiently confine a tracer, while a single attractive obstacle can cause confinement. Note that we did not include large attractive binding free energy in our analysis.

For attractive obstacles, α_{\min} is independent of binding energy over the range we studied (fig. 4). The characteristic time for a tracer to unbind from an attractive obstacle depends on the binding energy, leading to the energy-dependent variation in the anomalous time we observe. However, it is properties of the obstacle arrangement, rather than of binding, which determine the shape of the MSD curve, and therefore the α_{\min} . The minimum anomalous exponent occurs when tracers are, on average, confined to a cage formed by inter-obstacle boundaries and single-site wells. Therefore, the minimum anomalous exponent is approximately the same for all binding energy, but varies with filling fraction.

We note that the sticky soft obstacle model studied here does not simply map to the impenetrable obstacles at a lower effective obstacle filling fraction. Such a mapping cannot be made because tracers can 'hop through' single obstacles via binding, while never being able to hop between obstacles. Sticky obstacles allow for move attempts—and blocks—that would



Figure 6. Sticky obstacles of size one in 3D. (a) Diffusion coefficient D^* , (b) anomalous time t_a , and (c) minimum scaling exponent α_{\min} as a function of obstacle filling fraction. The approximate locations of the critical occupancies ν^l and ν^u are indicated with gray dotted lines.



Figure 7. Slippery obstacles in 2D. (a, d) Diffusion coefficient D^* , (b, e) anomalous time t_a , and (c, f) minimum scaling exponent α_{\min} as a function of obstacle filling fraction ν for positive (top) and attractive (lower) binding energy. The approximate locations of the critical occupancies ν^l and ν^u are indicated with gray dotted lines.

never be attempted in the impenetrable case.

3.1 Sticky soft obstacles in 3D

We extended our study of single-site sticky repulsive obstacles to three dimensions, to determine whether the spatial dimension plays a key role in the tracer behavior (fig. 6). The results are qualitatively the same as the 2D model (fig. 4). However, in 3D, the lower and upper critical occupancies appear at higher filling fraction: a higher obstacle filling fraction is required to percolate a 3D lattice. The anomalous time is also typically smaller in 3D. For soft sticky obstacles, increasing the spatial dimension does not change the qualitative features of our model, but does shift the critical occupancies and anomalous time.

4 Slippery soft obstacles

When obstacles are perfectly slippery, bound tracers can hop to adjacent obstacles without penalty. Our model of perfectly slippery obstacles contains an occupancy-energy inversion symmetry: the dynamics are invariant to changing the filling fraction by switching obstacles and empty sites $(\nu \rightarrow 1 - \nu)$ while



Figure 8. Slippery obstacles of size 1 in 3D. (a) Diffusion coefficient D^* , (b) anomalous time t_a , and (c) minimum scaling exponent α_{\min} as a function of obstacle filling fraction ν . The approximate locations of the critical occupancies ν^l and ν^u are indicated with gray dotted lines.

simultaneously switching the sign of the binding energy $(\Delta G \rightarrow -\Delta G)$. In other words, a low filling fraction of attractive obstacles is equivalent to a high filling fraction of repulsive barriers (fig. 7).

Slipperv obstacles remove the obstacle percolation threshold for all measured binding energies (fig. 7). The curves for $\Delta G = 10$ for the repulsive slippery obstacles qualitatively resemble the sticky case (fig. 4), because the diffusion coefficient approaches zero for However, for slippery obstacles, the $\nu \approx 0.4.$ anomalous time increases, but does not diverge, at the percolation threshold, and then decreases at larger filling fraction. For slippery obstacles with finite ΔG , one can always find a time after which the system displays normal diffusion. Slipperv obstacles lead to non-monotonic behavior: for large enough ν , the diffusion coefficient increases and anomalous time decreases. For high obstacle filling fraction, binding increases tracer mobility, because they can hop along the percolating network of obstacles. Similarly, the minimum exponent is non-monotonic with filling fraction.

4.1 Slippery soft obstacles in 3D

As for sticky obstacles, we examined tracer motion with single-site slippery obstacles in three dimensions (fig. 8). The results are qualitatively the same as the 2D model (fig. 7), with typically smaller anomalous time.

4.2 Comparison of sticky and slippery obstacles in 2D

The limits of perfectly sticky and slippery obstacles are most similar at low filling fraction (fig. 9). In general, slippery obstacles lead to exponents closer to one (less anomalous) than do sticky obstacles, because tracers are not caged by the obstacle-obstacle interface. Even for relatively small values of the binding energy $(|\Delta G| \leq 3)$ and intermediate filling fraction, sticky and slippery obstacles lead to significantly different tracer dynamics (fig. 9). Slippery obstacles, on which motion can occur for high obstacle filling fraction, allow normal diffusion with coefficients comparable to those for low filling fraction. This effect may be important to explain the rates of a number of biological processes that are diffusion-limited, including transcriptional regulation and nucleo-cytoplasmic transport.

5 Semi-slippery obstacles

Having compared the limits of perfectly sticky $(D_{\text{bound}} = 0)$ and slippery $(D_{\text{bound}} = D_{\text{free}})$ obstacles, we now study intermediate cases. We varied the bound diffusion coefficient for repulsive binding energy $\Delta G = 1, 2, 3, \infty$ and filling fraction $\nu = 0.3$ and 0.6. For finite binding energy, increasing D_{bound} increases the long-time diffusion coefficient (fig. 10). This effect is larger for higher filling fraction and lower binding energy, when tracers spend more time bound. Varying D_{bound} has little effect on the anomalous time at low filling fraction, because t_a is already near the threshold at which we can accurately measure it. However, increasing D_{bound} decreases t_a at higher filling fraction, because tracers can more quickly escape obstacles when their bound diffusion coefficient is larger. Similarly, varying D_{bound} has little effect on α_{\min} at low ν , but does make diffusion less anomalous at higher filling fraction, because increasing bound motility reduces tracer caging.

6 Varying obstacle size

We varied the length of the obstacles l_{obst} , while maintaining their square shape. Increasing the obstacle size (with filling fraction fixed) clusters obstacles. Since



Figure 9. Comparison of models with slippery repulsive obstacles (solid lines), sticky repulsive obstacles (dashed lines), and hard repulsive obstacles (purple dashed line). (a) Diffusion coefficient D^* , (b) anomalous time t_a , and (c) minimum scaling exponent α_{\min} as a function of obstacle filling fraction ν . The gray dotted lines are indicating the approximate locations of critical occupancies ν^l and ν^u .



Figure 10. Semi-slippery obstacles in 2D. Varied the bound diffusion coefficient from the sticky $D_{\text{bound}} = 0$ to slippery $D_{\text{bound}} = D_{\text{free}}$ limit for single site $l_{\text{obst}} = 1$ obstacles. Top panel: (a, d) Diffusion coefficient D^* , (b, e) anomalous time t_a , and (c, f) minimum scaling exponent α_{\min} as function of D_{bound} for low filling fraction $\nu = 0.3$ (top) and high filling fraction $\nu = 0.6$ (bottom).

in our model the binding penalty occurs only for **empty** \rightarrow **obstacle** moves, increasing the size of obstacles effectively reduces the number of binding sites: more obstacle sites are interior to obstacles, rather than on their perimeter. For sticky obstacles with $l_{obst} = 1$, tracers can easily hop through cages, since their bound motion is only blocked by an obstacle-obstacle interface. Increasing the obstacle size guarantees that individual obstacles will contain an obstacle-obstacle interface, which makes it less likely that tracers can hop through neighboring obstacles (fig. 11). Increasing obstacle size at fixed filling fraction also increases the typical distance between obstacles. These changes alter obstacle percolation effects: ν^l and ν^u depend on l_{obst} .

6.1 Sticky obstacles of varying size

First, we examined tracer dynamics on sticky obstacles of variable size (fig. 12). Qualitatively, large sticky obstacles have a soft surface (binding can occur on surface sites, although hops along the surface are still blocked), but a hard core (interior sites are inaccessible). A significant change in dynamics occur



Figure 11. Cartoon showing size effects for sticky and impenetrable obstacles. Red lines indicate borders between obstacles that cannot be crossed by a tracer.

when l_{obst} increases above 1. Any obstacle with $l_{obst} > 1$ is fundamentally different from $l_{obst} = 1$, because larger obstacles are guaranteed to contain sites with an adjacent obstacle site. Increasing l_{obst} prevents hopping across the interior of any one obstacle, which can hinder tracer motion. The cages are thus more robust. Tracers can still hop across corners, unlike in the case of a purely repulsive interaction (fig. 11).

The dependence of tracer dynamics on binding energy changes upon increasing the obstacle size above 1 (fig. 12). For size-one obstacles, particles can hop through a single obstacle, and so lower binding energy leads to higher long-time diffusion coefficient. In contrast, with larger obstacles, high binding energy leads to an increased diffusion coefficient. With higher repulsion, a tracer is less likely to bind to the surface of an obstacle where it can get stuck. Thus, for larger obstacles, higher repulsion can facilitate motion.

For $l_{obst} > 3$, increasing obstacle size increases the cage size, and so the long-term diffusion coefficient and the anomalous time both increase smoothly, in agreement with previous work on impenetrable obstacles [26]. The anomalous time increases with l_{obst} above 3, because the effective cage size increases: tracers take longer to explore a cage to escape. For $l_{obst} \geq 3$ and small filling fraction, the size dependence is roughly energy independent. The dynamics are dominated by blocked **obstacle** \rightarrow **obstacle** moves, rather than by the energy dependence of **empty** \rightarrow **obstacle** moves. For low filling fraction, α_{\min} remains > 0.9, suggesting that obstacle caging effects are minimal.

Next, we examined a higher packing fraction $\nu = 0.6$, chosen because it is between ν^l and ν^u for size-1 obstacles in 2D. The effects of obstacle size on percolation are significant, leading to larger changes in behavior than for $\nu = 0.3$. As $l_{\rm obst}$ increases, obstacles are on average spaced farther apart, which increases ν^l .

In contrast, the upper critical concentration is

more complicated, because now each obstacle contains within it obstacle-obstacle interfaces. The upper critical concentration decreases below 0.6 for $l_{obst} = 3$, and therefore the dynamics are anomalous at all times; t_a diverges and D^* goes to zero. Above $l_{obst} = 3$, the upper critical concentration increases with increasing obstacle size. For $l_{\rm obst} = 5$, $\nu^u > 0.6$, leading to long-time Fickian diffusion. Here t_a decreases with $l_{\rm obst}$, because the time required for a tracer escape a cage is not dominated by the cage size (as it was for low ν), but by the time needed to find a gap between cages. As l_{obst} increases, the gaps become larger on average, lowering the escape time. Overall, above $l_{obst} = 5$, the behavior is only mildly dependent on either obstacle size or binding energy, making the long term diffusivities primarily a function of the filling fraction.

6.2 Slippery obstacles of varying size

Understanding the effects of variable obstacle size on tracer motion is more straightforward for the case of slippery obstacles, because the difference between edge and interior obstacle sites is eliminated (fig. 13). In the perfectly slippery limit, increasing l_{obst} effectively lowers the number of binding sites: tracers experience the binding energy change only when binding to obstacle edge sites, but can move freely through obstacle interior sites. Therefore, D^* and α_{\min} increase with obstacle size, an effect that is larger for higher filling fraction, because obstacle overlaps at high filling fraction lower the fraction of obstacles that impede motion and cage tracers. In nearly all cases, t_a increases with obstacle size, because the effective cage size grows. The exception occurs for impenetrable obstacles, where increasing l_{obst} increases the size of vacancies between cages, allowing caged tracers to escape more quickly.



Figure 12. Size effects for sticky obstacles in 2D. (a, d) Diffusion coefficient D^* , (b, e) anomalous time t_a , and (c, f) minimum scaling exponent α_{\min} as a function of obstacle filling fraction ν for $\nu = 0.3$ (top) and $\nu = 0.6$ (lower).



Figure 13. Size effects for slippery obstacles in 2D. (a, d) Diffusion coefficient D^* , (b, e) anomalous time t_a , and (c, f) minimum scaling exponent α_{\min} as a function of obstacle filling fraction ν for $\nu = 0.3$ (top) and $\nu = 0.6$ (lower).

7 Conclusion

In this paper, we have studied a lattice model of tracer particles that diffuse and experience crowding due to immobile obstacles. While most previous work has considered hard (impenetrable) obstacles, we consider soft (penetrable) obstacles characterized by a binding free energy that allows tracers to overlap with obstacles. We also consider the effects of varying the tracer mobility while bound, including the limiting cases of 'sticky' obstacles (which immobilize bound tracers) and 'slippery' obstacles (which allow full tracer mobility), as well as the intermediate regime between the two.

In some cases, diffusion crowded media leads to dynamics that are anomalous $(r^2 \sim t^{\alpha})$ with a constant α [1]. However, our system typically does not give a power-law dependence of the MSD on time delay; this has been seen by others [8, 38]. As a result, we quantified a long-time diffusion constant (D^*) , the timescale on which the systems transitions from anomalous to Fickian (t_a) , and the minimum instantaneous anomalous exponent (α_{\min}) .

Our results demonstrate the key differences between sticky and slippery obstacles. For sticky obstacles, increasing the obstacle filling fraction decreases the diffusion coefficient and increases the degree of anomalous diffusion. Above an upper critical occupancy $\nu^u \approx 0.72$ in 2D, diffusion becomes anomalous at all times, independent of binding energy. In the sticky case, the minimum anomalous exponent, α_{\min} monotonically decreases with filling fraction, because adding more obstacles creates more cages in which tracers become transiently confined.

For slipperv obstacles, by contrast, tracers always reach normal diffusion after a sufficiently long time; even increasing the filling fraction above the percolation threshold does not eliminate tracer For nonzero binding free energy, we find motion. a novel non-monotonic dependence of D^* on filling fraction: increasing the filling fraction away from zero introduces binding sites that slow tracer diffusion, but for sufficiently high filling fraction, bound mobility allows tracer motion along clusters of obstacles. The anomalous exponent decreases with binding energy magnitude, but varies non-monotonically with filling fraction. For low filling fraction, α_{\min} decreases as more obstacles are added, because binding transiently traps tracers on isolated obstacles. For sufficiently high density, diffusion becomes more normal when tracers hop along clusters of obstacles while bound.

For intermediate 'semi-slippery' obstacles, we demonstrate that in the crossover from from sticky to slippery behavior, D^* , α_{\min} , and t_a vary smoothly. Increasing bound diffusion always makes the diffusion coefficient larger and the diffusive motion less anomalous.

We varied obstacle size to examine how relatively large obstacle 'domains' affects tracer motion in our model. For sticky obstacles, increasing obstacle size above 1 led to a sharp jump in tracer properties. This occurs because larger obstacles always contain interior obstacle sites, which are inaccessible to tracers in the sticky model. For large obstacles, increasing repulsive binding energy tends to increase the tracer diffusion coefficient, because tracers spend less time trapped in a binding site.

For slippery obstacles, perimeter and interior obstacle sites are both accessible, which means that varying obstacle size has effects that are easier to understand intuitively. The diffusion coefficient and anomalous exponent increase with obstacle size, because larger obstacles lead to a fewer obstacle-empty boundaries. The effect of obstacle size on t_a varied with filling fraction, due to competing effects on increasing cage size and increasing gaps between cages.

Our models separately represent effects of soft interactions (through the binding energy) and boundstate motion (through obstacle stickiness/slipperiness). Sticky and slippery obstacles show dramatically different tracer dynamics, even at short time and low filling fraction. Slippery obstacles lead to a diffusion coefficient which varies non-monotonically with filling fraction, with high values at both high and low obstacle densities. As the filling fraction increases from zero, the particles are more and more inhibited by obstacles. However, as the obstacle density increases, particles which bind can more easily move between obstacles. This may describe transport factor motion within the nuclear pore complex, where transport factors can slide on the disordered FG Nups [35]. Therefore, biological systems may use soft interactions and slippery obstacles to allow particle diffusion, even in the highly crowded cellular interior.

Our work highlights how soft interactions and bound-state mobility can dramatically change tracer These effects are relevant to biological motion. systems, ranging from membrane-less organelles to lipid rafts. Although most previous theoretical work on crowded diffusion has focused on the anomalous exponent, these biological examples highlight the importance of changes in the diffusion coefficient. For example, proteins which do not passage through the nuclear pore complex on biologically relevant time scales (minutes to hours) cannot have biological effects, and so the speed of passage is the fundamentally important biological quantity. The long-time diffusion coefficient varies dramatically in our model between hard obstacles, soft sticky obstacles and soft slippery obstacles (figure 9). Thus, the effective permeability of obstacles and the degree to which bound particles can diffuse can be used by cells to tune macromolecular motion.

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