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PAPER

Anomalous, non-Gaussian tracer diffusion in crowded two-dimensional environments

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Abstract

A topic of intense current investigation pursues the question of how the highly crowded environment of biological cells affects the dynamic properties of passively diffusing particles. Motivated by recent experiments we report results of extensive simulations of the motion of a finite sized tracer particle in a heterogeneously crowded environment made up of quenched distributions of monodisperse crowders of varying sizes in finite circular two-dimensional domains. For given spatial distributions of monodisperse crowders we demonstrate how anomalous diffusion with strongly non-Gaussian features arises in this model system. We investigate both biologically relevant situations of particles released either at the surface of an inner domain or at the outer boundary, exhibiting distinctly different features of the observed anomalous diffusion for heterogeneous distributions of crowders. Specifically we reveal an asymmetric spreading of tracers even at moderate crowding. In addition to the mean squared displacement (MSD) and local diffusion exponent we investigate the magnitude and the amplitude scatter of the time averaged MSD of individual tracer trajectories, the non-Gaussianity parameter, and the van Hove correlation function. We also quantify how the average tracer diffusivity varies with the position in the domain with a heterogeneous radial distribution of crowders and examine the behaviour of the survival probability and the dynamics of the tracer survival probability. Inter alia, the systems we investigate are related to the passive transport of lipid molecules and proteins in two-dimensional crowded membranes or the motion in colloidal solutions or emulsions in effectively two-dimensional geometries, as well as inside supercrowded, surface adhered cells.

1. Introduction

The cytoplasmic fluid of living cells is a superdense medium [1–3], in which biomacromolecules occupy volume fractions reaching well above 30% [4–8]. This macromolecular crowding (MMC) affects the diffusion of larger passive molecules, endogenous as well as artificially introduced submicron tracer particles, and cellular components [9]. One of the central observations is the existence of transient but often very extended anomalous diffusion [10, 11] with the sublinear scaling

$$\langle \mathbf{r}^2(t) \rangle \simeq K_\beta t^\beta \quad (1)$$

of the mean squared displacement (MSD) of the diffusing particles with the anomalous diffusion exponent in the subdiffusive range $0 < \beta < 1$ [9, 12]. Here K_β is the generalised diffusion coefficient with units $\text{cm}^2/\text{sec}^\beta$. Subdiffusion in the crowded cytoplasm of living cells was observed for fluorescent smaller proteins [13, 14], labelled polymeric dextrane [15] and messenger RNA [1, 16], chromosomal loci and telomeres [16, 17], as well as submicron endogenous granules [18–20] and viruses [21]. Subdiffusion was also reported for the motion of tracer particles in artificially crowded environments [22–30]. We note that active transport processes in living cells may lead to superdiffusion with $1 < \beta < 2$ [31–35].

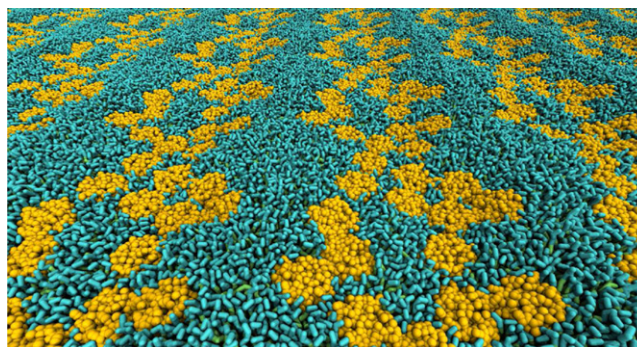


Figure 1. Protein crowded lipid bilayer membrane, illustration from coarse grained simulations. The lipids (small blue molecules) move in an array of proteins (large yellow objects) creating local confinement and bottlenecks for the passage of the lipids. This causes pronounced anomalous diffusion of lipids and proteins [43]. Figure courtesy Matti Javanainen.

Similarly to the volume crowding, cellular membranes are highly crowded interfaces with up to 25% of the total area [36] occupied by various membrane proteins, ion channels, and pores [37]. By fluorescent correlation spectroscopy and single particle tracking methods anomalous diffusion of the form (1) was measured for both membrane embedded proteins [37–39] and lipid molecules [40]. These experimental findings are corroborated by molecular dynamics as well as coarse grained simulations [41–46]. In model membranes as the one shown in figure 1 the embedded proteins may form aggregates or show only weak mutual attraction, depending on the chemical structure of the lipids and proteins [43]. As illustrated in figure 1 the proteins may form cage-like environments and bottlenecks for the lipid molecules, effecting pronounced anomalous diffusion [43]. The effective mobility in effectively two-dimensional, crowded colloidal systems or emulsions is also of current interest, including the existence of jamming effects [47].

Particle diffusion in crowded and structured three-dimensional environments has been in the focus of a number of simulations [48–55] and theoretical studies [9, 49, 56, 57]. In membrane systems similar structural elements are effected by the cellular cytoskeleton supporting the membrane [37, 39]. The observed anomalous diffusion in such systems is addressed by various generalised stochastic processes [58–60]. Specifically in an environment of densely packed, freely moving crowders the tracer diffusion follows Brownian motion at sufficiently long times [53], whereas for crowders confined by a potential and for static crowders the tracer diffusion features an extended—albeit ultimately transient—subdiffusive regime [53, 61].

The current interest in crowding comes from experimental evidence of polydisperse mixtures of crowding proteins and spatial variation of the apparent diffusion coefficient in the bacterial [62–64] and eukaryotic [65, 66] cytoplasm, see also [8, 53, 55]. Moreover, the distribution of crowders in the cell was shown to be rather heterogeneous, effecting a faster mobility of small tracer proteins near the cell nucleus of surface adhered cells [65]. We also mention the spatially dependent protein diffusion in the cytoplasm and nucleus [66], the dynamical heterogeneity of the cell cytoplasm [3], and a position dependent viscosity in bacteria [67]. These properties of the cell cytoplasm impose severe restrictions on the rates of biochemical reactions [68, 69] including polymer dynamics [70–73] and particle diffusion [53, 55, 61]. As discussed below our two-dimensional model should remain valid for the diffusion of large tracers in surface adhered, superdensely crowded cells.

In lipid bilayer membranes heterogeneities are also common. Thus cholesterol is known to lead to aggregate [74] and supramolecular complex formation [75]. This controls the preference for different membrane phases [76–78]. Demixing of lipid components and pattern formation may occur [79]. The formation of such domains by electrostatic lipid-proteins interactions was demonstrated by simulations recently [80]. Moreover, microdomains of up to 200 nm in size, so-called lipid rafts, are believed to function as organising centres for molecular assembly [81].

Here we address two aspects of crowding: the finite size of the tracer and heterogeneous distributions of crowders (HDC) in a two-dimensional circular domain. This domain could represent a region of a protein crowded suspended model membrane or a membrane domain within a larger lipid bilayer. Similarly it could represent a fluidic device with a mixture of smaller and larger colloidal particles or emulsion droplets. In the space between the outer boundary and the centre—representing, for instance, a large receptor domain—we place either homogeneous or heterogeneous distributions of monodisperse crowders, as shown in figure 2. We then simulate the motion of a finite sized tracer through a static crowder configuration. Given the separation of mobilities between lipids and proteins in membranes or between small colloids and large, squeezed droplets in fluidic setups, the assumption of a static crowding background appears a good approximation. The implications

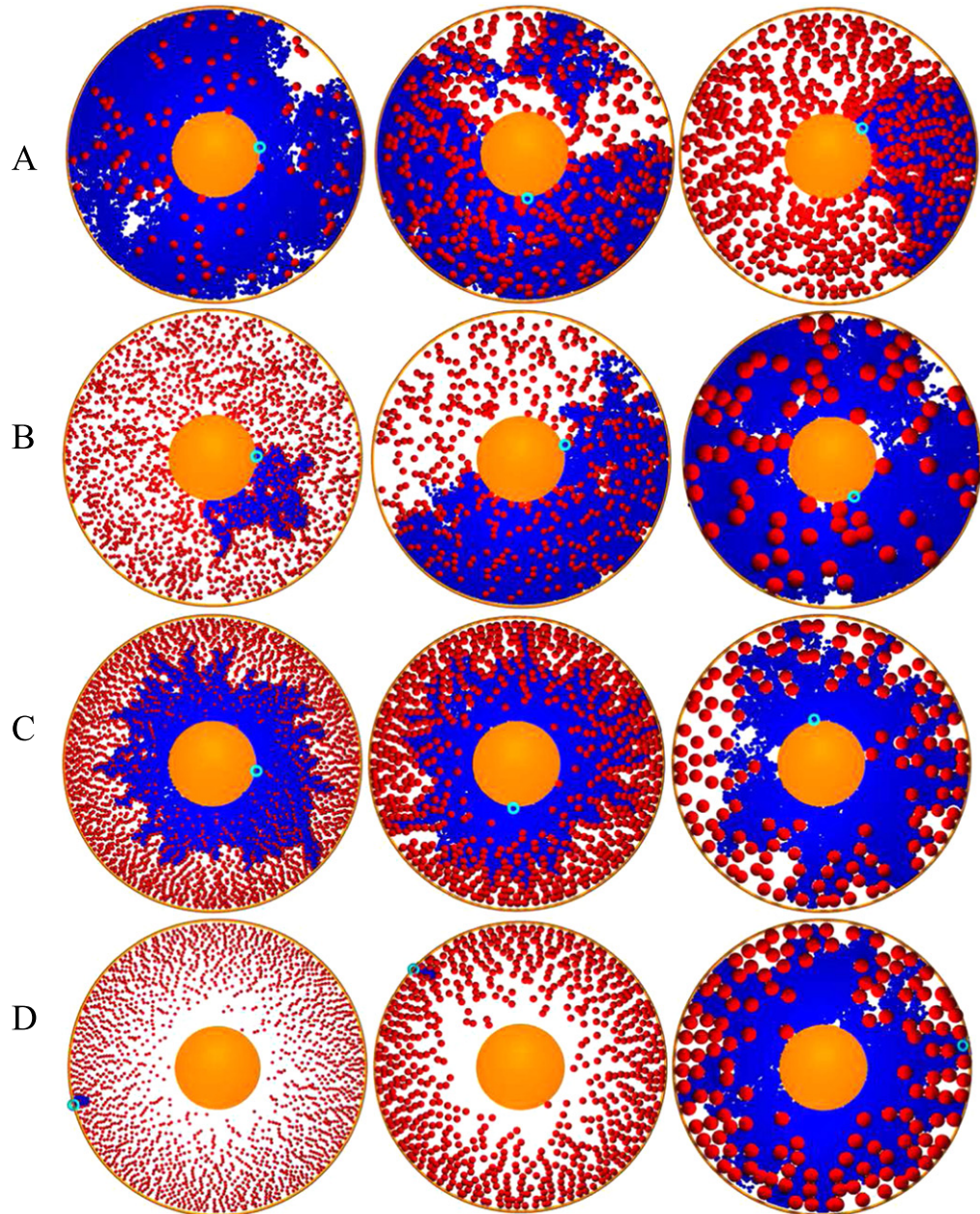


Figure 2. Diffusion profiles for homogeneous (panels (A) and (B)) and heterogeneous ((C) and (D)) monodisperse crowders. In panel (A) the fraction of crowders increases from left to right: $\phi = 0.05, 0.2, 0.3$, and their size is fixed to $R_c = 2$. In panel (B) the size of the crowders grows from left to right: $R_c = 1, 2, 5$, while the crowding fraction is fixed at $\phi = 0.2$. For HDC in panels (C) and (D), the crowding fraction $\phi(r)$ exhibits the linear growth (2). The centre represented by the excluded disk shown in orange represents a receptor domain or similar structure. The diffusion is in-out (i.e., from the centre to the domain boundary) in panel (C) and out-in (from the boundary to the centre) in panel (D). The initial tracer position is shown by the small blue circle. The crowder radius in panels (C) and (D) grows from left to right: $R_c = 1, 2, 5$. The regions of the domain visited by the tracer up to the diffusion time $T = 10^4$ are depicted in blue. For a single tracer diffusing in the domain with HDC, significant asymmetries in the particle spreading and sampling of the space may arise at higher crowding fractions.

of polydisperse mixtures of crowders is beyond the scope of this paper, see the discussion section. We investigate the two biologically relevant scenarios of in-out (from centre to domain boundary) versus out-in (from domain boundary to centre) tracer diffusion, finding fundamental differences. From extensive simulations we determine the particle distribution for different crowding environments and study the particle dynamics in terms of the ensemble and time averaged MSDs. We further analyse the non-Gaussianity of trajectories, the van Hove correlation function, as well as the behaviour of the survival probability of the tracer.

The paper is organised as follows. In the next section we set up the model, discuss the simulation procedure and the data analysis. In sections 3 and 4 we describe the main results for the homogeneous and heterogeneous cases, respectively, and compare them to theoretical models. Further analyses of both cases in terms of the non-

Gaussianity parameter and the van Hove function are shown in section 5, while the survival probability is discussed in section 6. In section 7 we draw our conclusions and discuss some applications of our results.

2. Model, simulations scheme, and data analysis

We consider a model domain in the form of a planar circular annulus between the centre, an excluded region of radius a , and the domain boundary at radius R , see figure 2. The space between the boundary and the centre is filled with *static* monodisperse crowders of radius R_c . We consider two cases: a homogeneous distribution of crowders with a prescribed crowding (area) fraction ϕ , and HDC with a linear radial gradient

$$\phi(r) = \phi(a) + \frac{\phi(R) - \phi(a)}{R - a}(r - a) \quad (2)$$

for $a < r < R$. As we show below, equation (2) leads to a transient subdiffusion of the tracer motion from the centre to the boundary, emerging due to an increasing density of crowders near the boundary as evidenced by figures 2(C), (D). The present approach is complementary to our recent studies of heterogeneous diffusion processes [82–87] with a deterministic space dependent diffusivity. Similar to the findings in those studies the explicit HDC considered here gives rise to a radially varying particle diffusivity in the domain, to non-Brownian diffusion, and non-Gaussian statistics of particle displacements, see below. The methodology reported here further develops the model of tracer diffusion in lattices with periodically distributed crowders examined by us recently [61].

In the simulations the crowders are placed at random positions in the domain without overlap. The initial tracer position is well in between neighbouring crowders. The highest crowding fraction is around 30%. When computing the mean time averaged MSD we typically average over $M = 10^2$ random configurations of crowders. This disorder average is taken in addition to the average over individual trajectories in a given, quenched crowder configuration.

The tracer particle has a fixed unit radius $\sigma/2$. We set the outer and inner domain radii to $R = 100$ and $a = 30$, respectively. The radii are measured in terms of the length scale $\sigma = 6$ nm, see below. For most cases of HDC we set $\phi(a) = 0.01$ and $\phi(R) \approx 0.3$. The latter value is considerably smaller than the percolation threshold in two dimensions. However, due to the finite size of the diffusing particles, the effective crowding fraction ϕ_{eff} is higher than for a point-like tracer. For instance, for $R_c = 5$ and a unit tracer radius one gets for a domain with $R = 100$ and $a = 30$ that $\phi_{\text{eff}} \approx 0.43$; this is already much closer to the percolation threshold of $\phi \approx 0.59$.

The Weeks–Chandler–Andersen repulsive potential given by the 6–12 Lennard–Jones (LJ) potential $E_{\text{LJ}}(r)$ with the standard cutoff distance r_{cut} is used to parameterise the repulsion between the tracer and crowders

$$E_{\text{LJ}}(r) = 4\epsilon \left[\left(\frac{\sigma}{r} \right)^{12} - \left(\frac{\sigma}{r} \right)^6 + \frac{1}{4} \right] \quad (3)$$

for $r < r_{\text{cut}} = 2^{1/6}\sigma$, and $E_{\text{LJ}}(r) = 0$ otherwise [88–90]. We do not consider here attractive tracer-particle interactions, see the discussion section. We evaluate the LJ based interactions in our two-dimensional system by calculating the centre-to-centre distance between interacting discs. We represent the LJ potential in polar coordinates, independent of the space dimension. The potential of the form (3) is often used to simulate the properties of two-dimensional fluids [91–93]—see also [94] for higher dimensions. When implementing the repulsive part of the LJ potential in our simulations, we calculate the force in one dimension and then update the coordinates x and y independently. The details of the system behaviour, however, can evidently depend on the procedure of truncation of the LJ potential, see, for instance, [89, 90] for an example of two and three-dimensional LJ fluids. Some of these differences for systems of repulsive discs can, however, be reconciled in terms of an effective particle diameter [92].

We simulate the dynamics of the mid point $\mathbf{r}(t)$ of the tracer via the Langevin equation

$$m \frac{d^2 \mathbf{r}(t)}{dt^2} = - \sum_j \nabla \left[E_{\text{LJ}} \left(|\mathbf{r} - \mathbf{R}_j| - \left(R_c + \frac{\sigma}{2} \right) \right) + E_{\text{LJ}} \left(|\mathbf{r}| - \left(a + \frac{\sigma}{2} \right) \right) \right] + E_{\text{LJ}} \left(\left(R - \frac{\sigma}{2} \right) - |\mathbf{r}| \right) - \xi \mathbf{v}(t) + \mathbf{F}(t), \quad (4)$$

where m is the mass of the tracer particle, ξ is the friction coefficient experienced by the tracer particle, $\mathbf{v}(t)$ is its velocity, and \mathbf{R}_j is the static position of the j th crowder. Finally $\mathbf{F}(t)$ represents a Gaussian δ -correlated noise with zero mean and covariance

$$\langle \mathbf{F}_j(t) \cdot \mathbf{F}_k(t') \rangle = 2\delta_{j,k} \xi k_B \mathcal{T} \delta(t - t'). \quad (5)$$

The inertial term in equation (4) gives rise to a short time ballistic dynamics, as shown below (see [61] for more details). In the simulations we set $\epsilon = k_B \mathcal{F} = 1$, $m = 1$, and $\xi = 1$ corresponding to moderate damping [95–97]; the solution viscosity in these units is 1. The reader is also referred to the studies [55, 98] regarding the scale dependent viscosity known to exist in the cell cytoplasm. The diffusivity of a tracer in an uncrowded environment is

$$D_0 = k_B \mathcal{F} / \xi = 1. \quad (6)$$

The unit time step of simulations corresponds to the physical time $\tau = \sigma \sqrt{m / (k_B \mathcal{F})}$. For the chosen radius of σ , given a mass characteristic for the molecular weight of biomolecular crowders of a similar size of about 68 kDa [55, 99, 100], the physical time scale corresponds to $\tau \approx 1$ nsec. We employ the Verlet velocity algorithm with the time step $\delta t = 0.01$ to integrate the stochastic equation (4).

To characterise the diffusion behaviour we evaluate the time averaged MSD

$$\overline{\delta^2(\Delta)} = \frac{1}{T - \Delta} \int_0^{T-\Delta} [\mathbf{r}(t + \Delta) - \mathbf{r}(t)]^2 dt \quad (7)$$

for individual particle trajectories $\mathbf{r}(t)$. Here T is the observation time and Δ is the lag time defining the width of the window slid along the trajectory. This definition is standard in single particle tracking experiments [1, 12, 60, 101]. Given N measured trajectories the ensemble average is

$$\langle \overline{\delta^2(\Delta)} \rangle = \frac{1}{N} \sum_{i=1}^N \overline{\delta_i^2(\Delta)}. \quad (8)$$

For quenched HDC we also calculate the disorder average over M realisations of a crowding environment (compare [102, 103]),

$$\langle \overline{\delta^2(\Delta)} \rangle = \frac{1}{M} \sum_{j=1}^M \langle \overline{\delta^2(\Delta)} \rangle_j. \quad (9)$$

The ensemble averaged MSD is also computed as double average over N tracer trajectories for each crowders distribution and M crowders distributions

$$\langle \mathbf{r}^2(t) \rangle = \langle \overline{[\mathbf{r}(t) - \mathbf{r}(0)]^2} \rangle = \frac{1}{MN} \sum_{j=1}^M \sum_{i=1}^N [\mathbf{r}_i(t) - \mathbf{r}_i(0)]_j^2. \quad (10)$$

3. Homogeneous crowding: ensemble and time averaged MSDs

We start with the analysis of the tracer diffusion among static HDC, figures 2(A), (B). Figure 3 shows the ensemble averaged MSD $\langle \mathbf{r}^2(t) \rangle$ and the local MSD scaling exponent

$$\beta(t) = \frac{d[\log \langle \mathbf{r}^2(t) \rangle]}{d[\log t]}, \quad (11)$$

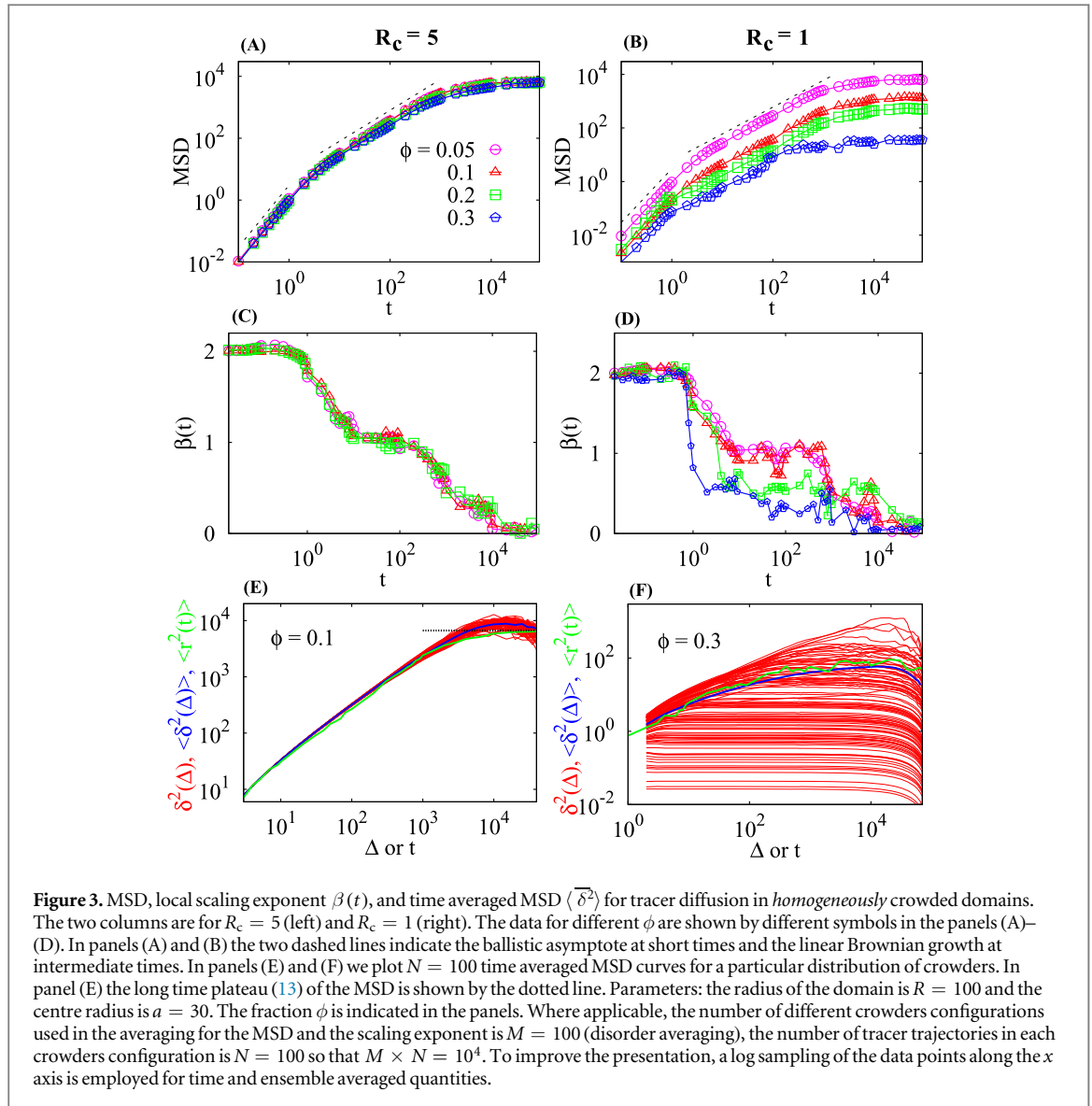
when the crowding fraction ϕ and the crowder radius R_c are varied. For relatively large crowders the initial standard ballistic growth of the MSD (underdamped particle motion, see also [61]) crosses over to a quite prolonged Brownian regime with $\beta \approx 1$, figures 3(A), (C). For the smaller crowders, after about the same physically expected relaxation time of $t \sim 1$, a region of subdiffusion emerges, compare panels (B) and (D) in figure 3 for the behaviour of the MSD and the scaling exponent with panels (A) and (C).

The ballistic regime in the tracer displacement extends to about the same time scale

$$t_{\text{ball}} \lesssim m / \xi = 1, \quad (12)$$

both for normal and subdiffusive tracer behaviour at later times, as we checked in the simulations (results not shown). At even later times the tracer motion starts to be affected by confinement due to the outer reflecting membrane boundary positioned at $r = R$, and the MSD begins to saturate to a plateau, as expected. Concurrently the exponent $\beta(t)$ tends to zero. This drop is solely due to the HDC in our model: for the tracer diffusion in dense mixtures of purely repulsive and uniformly smeared crowders, the diffusion was shown to be nearly Brownian [55]. We also expect that the short time ballistic behaviour of the MSD and the peculiar features of the repulsive tracer-crowder interaction potential do not measurably affect the long time scaling behaviour of the tracer MSD.

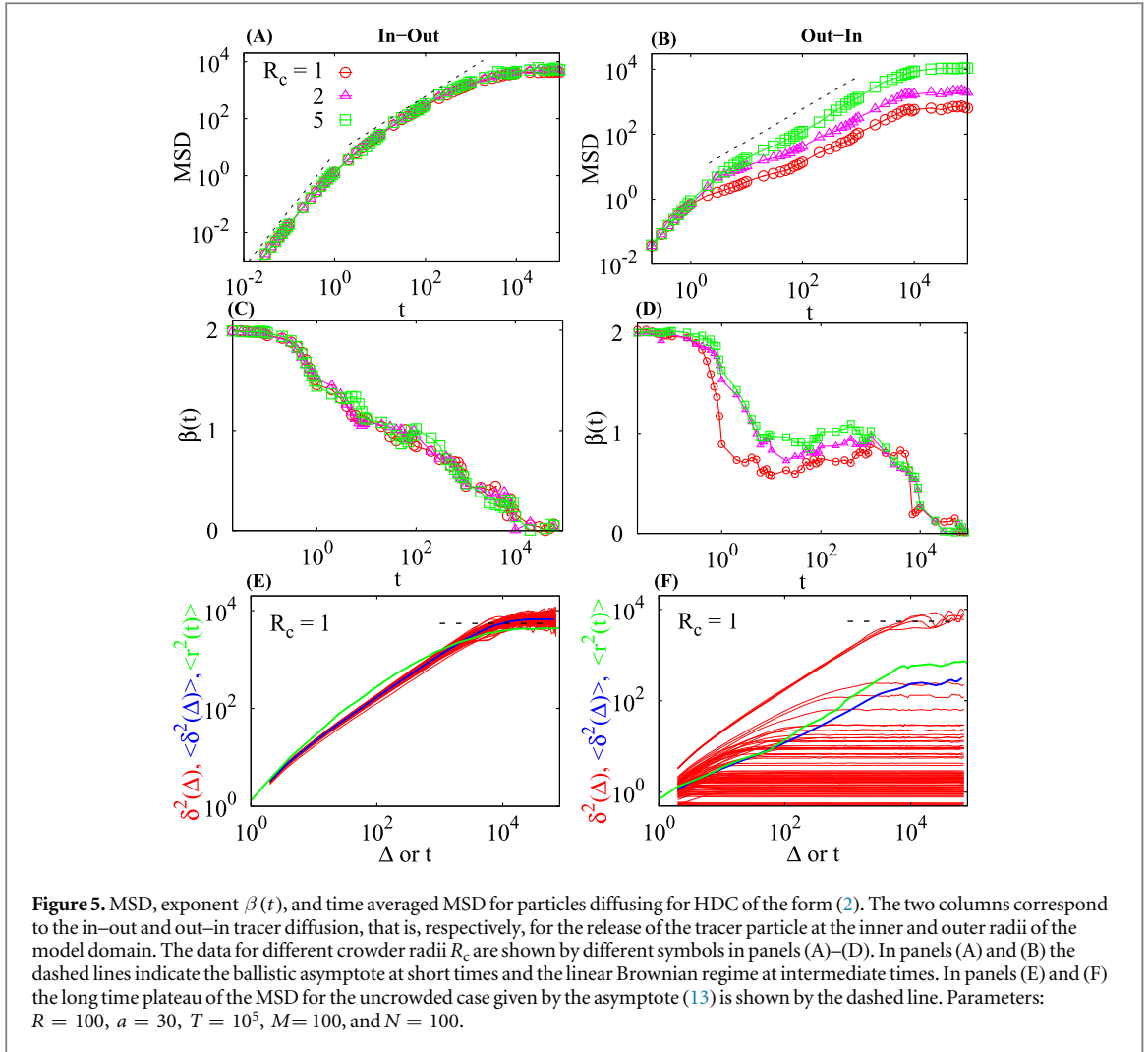
The effect of ϕ on the MSD behaviour is illustrated in panels (A) and (B) of figure 3 for large and small crowders, respectively. For a tracer of unit size the larger crowders do not appear to create substantial obstruction for the simulated MMC fractions. Even at relatively large ϕ values the MSD only marginally diminishes with increasing ϕ , see panel (A). In other words, small tracers always manage to manoeuvre around large void spaces remaining between *large* crowders (figure 2). In contrast, relatively *small* crowders at identical



MMC fractions yield severe reductions of the diffusive tracer motion, figure 3(B). This observation is consistent with the dramatic differences of the spatial space exploration patterns in panels (B) of figure 2. The magnitude of the MSD plateau attained for larger ϕ values is much smaller than expected for an annulus without crowders, compare with equation (13) below. This fact is due to the intricate *labyrinthine* environment formed by small crowders for diffusion of the same tracer. For small crowders the value of β is reduced significantly and at much earlier times for larger ϕ , see panels (C) and (D) of figure 3. These features obviously strongly depend on the specific quenched environment, in which the particle motion occurs, resulting in a high degree of irreproducibility of the tracer diffusion for different realisations of the disorder. This is also quantified in panel (F) of figure 3 and leads to substantially larger uncertainties in the scaling exponent $\beta(t)$ computed from the MSD curves. The strong effect of small crowders on the MSD and its scaling exponent at intermediate times and the minor effect of the large crowders is our first main result.

We now turn to the analysis of the time averaged MSD obtained from averaging over $M = 10^2$ tracer trajectories for a *single* crowder configuration. As shown in panel (E) of figure 3 for *large* crowders with $R_c = 5$ and small MMC fractions ϕ the spread of $\overline{\delta^2}$ is quite small. At later times—when the tracer motion starts to be influenced by the outer boundary—the average $\langle \overline{\delta^2}(\Delta) \rangle$ approaches a plateau which has about twice the amplitude of the MSD plateau, see panel (E) in figure 3. Note that due to the relatively small domain size used in the simulations, the moderate trajectory lengths, and the presence of randomly distributed crowders this plateau is not as distinct as, for instance, for the deterministic, confined heterogeneous diffusion processes, compare figure 4 in [86] and figure 8 in [87].

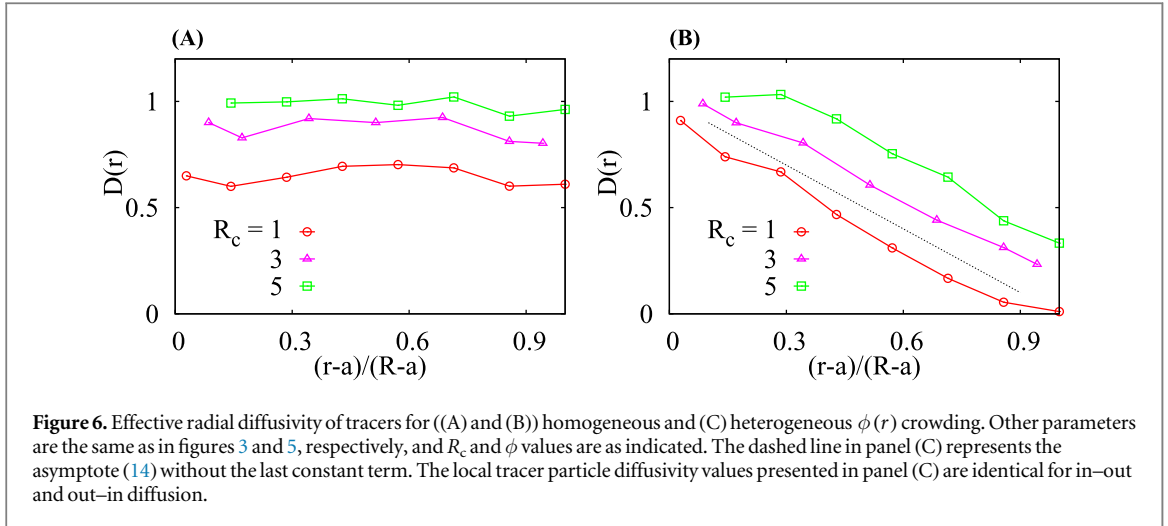
We recall that for the uncrowded case the long time (plateau) values of the ensemble and time averaged MSDs are related to the inner and outer radii in two-dimensions as



MSD is also observed when compared to the homogeneous case, see figure 5(E). The scatter of $\overline{\delta^2}$ decreases when longer trajectories are analysed (results not shown). This behaviour is expected and is realised for several ergodic and nearly ergodic processes [60, 105]. The magnitude of the amplitude scatter of $\overline{\delta^2}$ in panels (E) of figures 3 and 5 is comparable to that of Brownian motion [60, 106, 107] as the tracers explore a less crowded space over time.

In the opposite case of out–in diffusion (panel (D) of figure 2) we observe that for relatively high ϕ of small crowders a finite size tracer often cannot even leave the vicinity of the boundary, giving rise to prolonged trapping events in this confined area. This leads to a large proportion of low amplitude, nearly constant $\overline{\delta^2}$ curves, corresponding to the red curves in figure 5(F). The mean time averaged $\langle \overline{\delta^2}(\Delta) \rangle$ is in many cases dominated by several successful fast translocation events of tracers from the domain boundary to the centre. In the long time limit the asymptote (13) is thus not valid in this situation. The tracer localisation and the dominance of one or few extreme tracer trajectories in the mean $\langle \overline{\delta^2}(\Delta) \rangle$ is also a rather common feature of stochastic processes in the presence of well pronounced traps as well as in ageing stochastic processes [60, 108]. The distinctly different behaviour between out–in and in–out diffusion in the system of HDC is the third main results of this study.

Since the crowding fraction in equation (2) grows from the centre towards the boundary, the in–out diffusion for HDC is expected to be subdiffusive [84]. In this scenario the tracers are progressively trapped closer to the domain periphery. Here, however, we observe the formation of a radial percolation in the circular domain: the tracers are not allowed to penetrate beyond some critical radius that features a particular critical density of crowders, an effect that is crucially related to the *finite size* of both the crowders and the tracer particle. Similar effects of local confinement are naturally observed for single particle trajectories of tracer diffusion in random percolation systems [103]. The initial period of the tracer diffusion that occurs in the region with low crowder density is naturally reproducible and leads to a small spread of $\overline{\delta^2}$ curves, as shown in figure 5(E).



In contrast, for the out–in diffusion the tracer starts in the region of the highest concentration of crowders and diffuses into regions containing less and less crowders. The associated initial tracer localisation events give rise to stalling $\overline{\delta^2}$ curves with a very large spread, see figure 5(F). Early trapping in this scenario leads to the emergence of a longer plateau region in the corresponding time averaged MSD curve. The mean $\langle \overline{\delta^2} \rangle$ computed over an ensemble of N trajectories is rather imprecise because of the dominance of few extreme events.

We checked that for homogeneously distributed crowders the radial tracer diffusivity is approximately constant (panels (A) and (B) of figure 6), as it should be. As the size of the crowders decreases at a constant ϕ , the effective viscosity of the solution increases, as follows from the decreasing normalised diffusion coefficient plotted in figure 6(B). Also, the solution becomes effectively more viscous with increasing MMC fraction ϕ , see panel (A) of figure 6. In turn, for HDC given by equation (2) the radial tracer diffusivity is a decreasing and nearly linear function of the distance from the centre (panel (B) of figure 6),

$$D(r) \approx D_0 \left(1 - 0.78 \frac{r-a}{R-a} \right) + \text{const.} \quad (14)$$

The slope 0.78 is nearly independent on the crowder radius R_c . This linear dependence of the instantaneous diffusivity decrease in the domain is a consequence of the linear increase of the crowders density $\rho(r)$, equation (2). This linear dependence of $D(r)$ in equation (14) is our fourth main result.

5. Non-Gaussianity parameter and van Hove correlation function

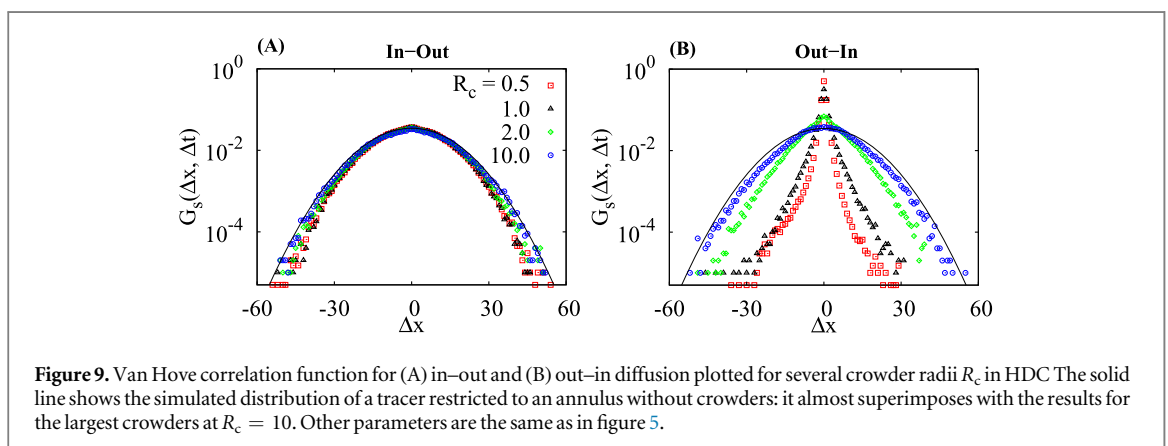
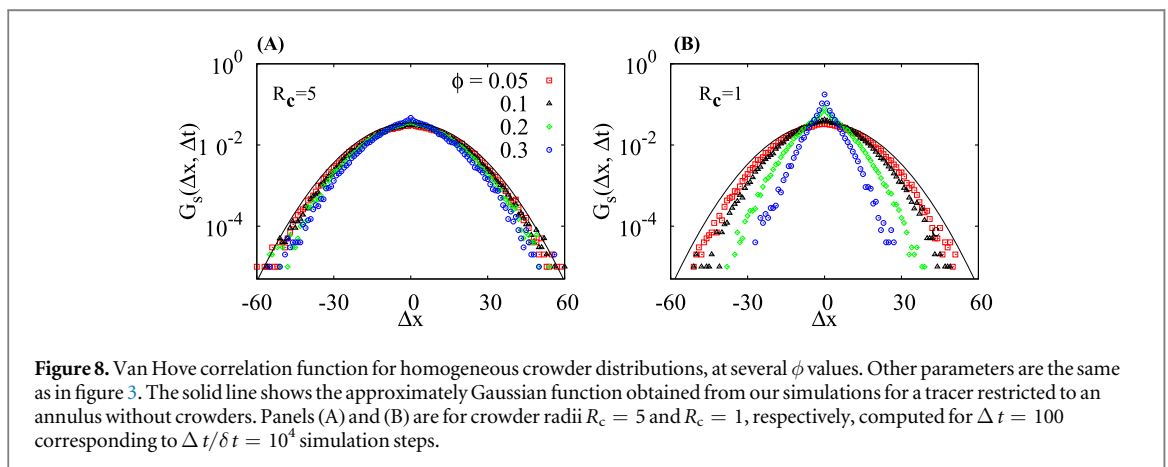
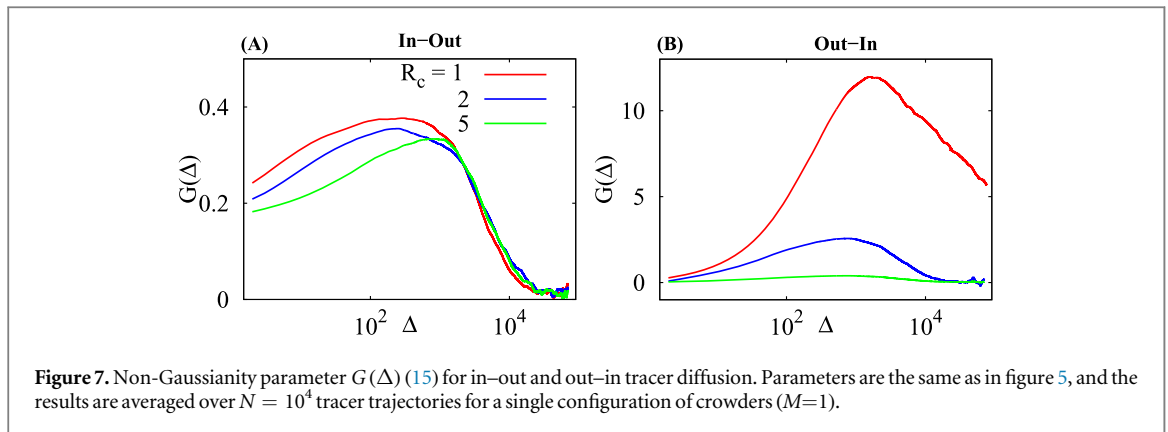
Following reference [61] we now proceed to evaluate the experimentally relevant non-Gaussianity parameter for the tracer diffusion in our crowded environment. Similar to the ergodicity breaking parameter EB [60, 109] it contains the fourth order moment of the tracer time averaged MSD. Namely, in two-dimensions we have [9, 110]

$$G(\Delta) = \frac{\langle \overline{\delta^4(\Delta)} \rangle}{2 \langle \overline{\delta^2(\Delta)} \rangle^2} - 1. \quad (15)$$

We find that for in–out diffusion the non-Gaussianity parameter assumes moderate values for shorter lag times Δ while it becomes close to zero for longer lag times, figure 7(A). This is a typical long time behaviour of ergodic tracer diffusion, compare figure 3(a) in [110]. Indeed, as we show in figure 5(E), in the long time limit the ensemble and time averaged MSDs differ simply by the above mentioned factor of 2 for HDC. In contrast for out–in diffusion the non-Gaussianity parameter attains substantially larger values. This feature is likely due to the highly non-reproducible trajectories of the tracer motion and prolonged localisation events near the domain boundary at high crowder concentration.

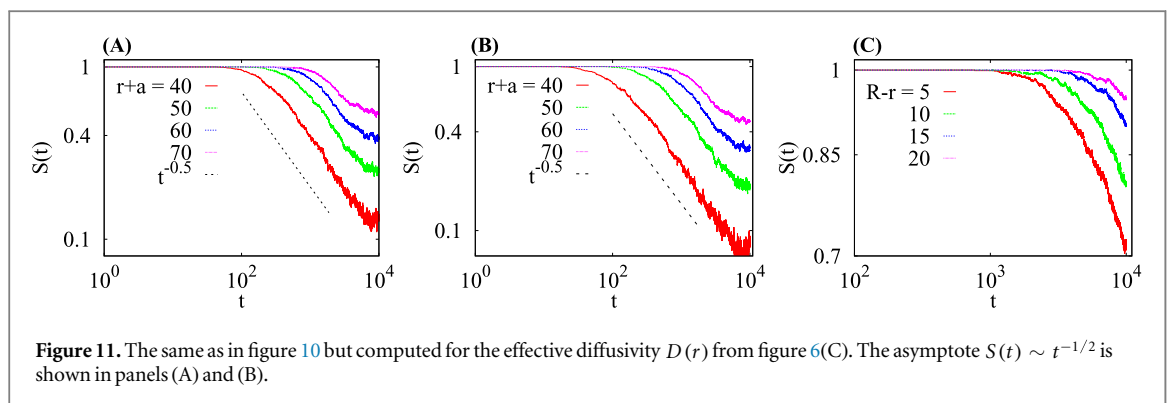
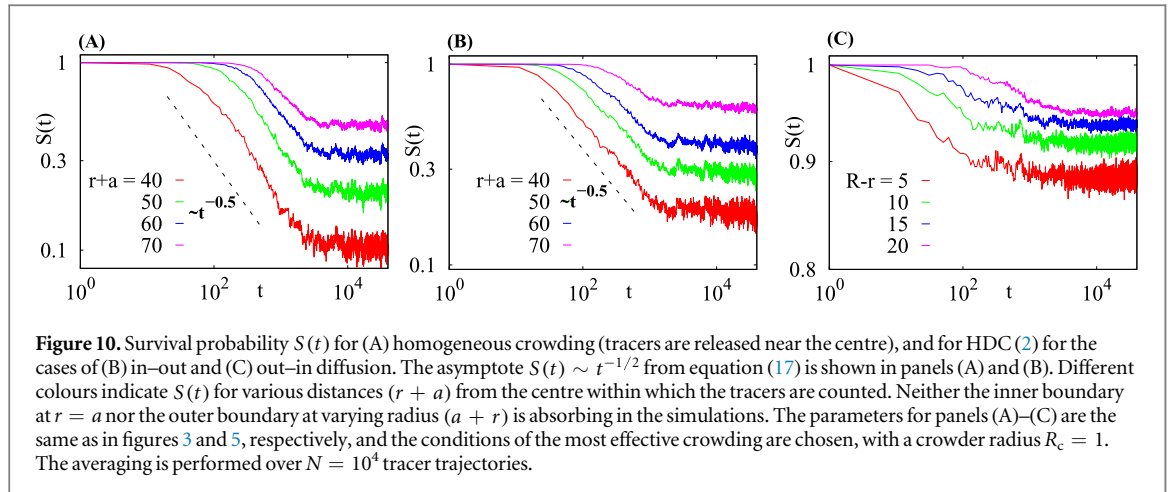
The trapping as well as the non-Gaussianity of the tracer diffusion can also be characterised by the van Hove correlation function $G_s(\Delta x, \Delta t)$ describing the probability that a particle moves a distance Δx during time Δt [111, 112],

$$G_s(\Delta x, \Delta t) = \frac{1}{N} \sum_{i=1}^N \delta(x_i(\Delta t) - x_i(0) - \Delta x), \quad (16)$$



where N is the number of tracers used for averaging. For a system of hard spheres this function simply corresponds to the Gaussian propagator [113] governed by the diffusion equation.

Figure 8 shows the behaviour of $G_s(\Delta x, \Delta t)$ for tracer diffusion in HDC. We observe that for relatively large crowders the probability of trapping events of the tracer particle is quite low and the distribution of tracer displacements remains close to Gaussian for all ϕ studied, as it should (see panel (B) in figure 8). In contrast, for small crowders the non-Gaussianity of the van Hove function becomes quite pronounced, in particular for larger ϕ , figure 8(A). A faster decay of the tracer displacements at high ϕ and for small crowders is a consequence of tracer caging by crowders and anomalously slow diffusion. The almost exponential distributions for high crowding fraction in panel (A) of figure 8 compare well with the experimentally measured step size distributions for polymer diffusion on nano-patterned surfaces presented in figure 4 of [114] as well as for liposome diffusion in nematic solutions of actin filaments in figure 3(C) of [115]. HDCs for tracer diffusion cause a separation of



particles into slow and fast populations (compare [83]), reflected in a *cusp* of the particle distribution near the origin and longer than Gaussian tails for large tracer displacements.

For HDC the behaviour of the van Hove function is illustrated in figure 9. We observe that for in–out diffusion the distribution remains approximately Gaussian for all crowder radii R_c in our simulations. This corresponds to the rather small non-Gaussianity parameter and quite limited spread of $\overline{\delta^2}$ curves. In contrast, for out–in diffusion among small crowders $G_s(\Delta x, \Delta t)$ becomes progressively non-Gaussian. It features a pronounced cusp near $\Delta x = 0$ describing a prevalence of small displacements characteristic for subdiffusion in rather confined conditions. The detailed behaviour of the non-Gaussianity parameter and the van Hove correlation function represents the fifth main result of the present study.

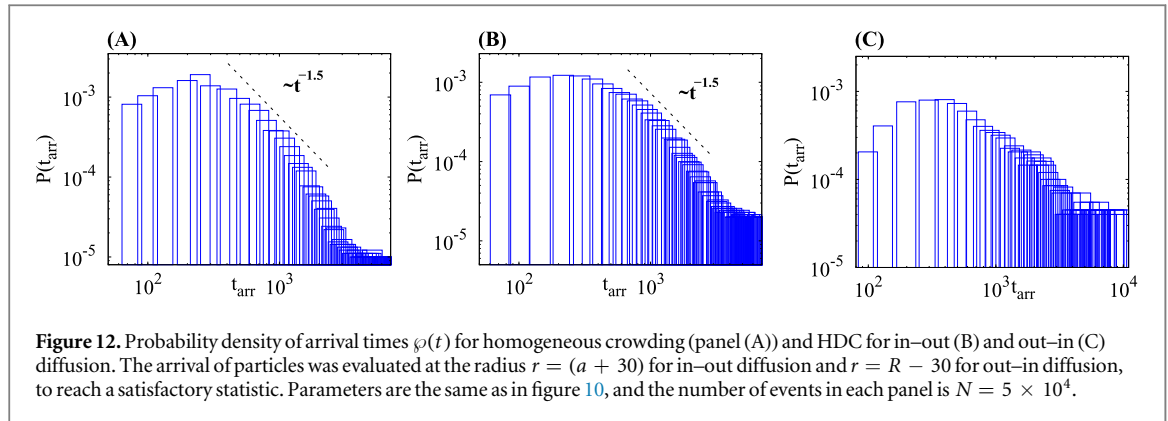
6. Survival probability

First passage time statistics are important to describe cellular processes, for instance, to quantify the diffusion limit of reactions triggered by incoming, diffusing molecules [116–118]. Similarly, the anomalously diffusive motion of membrane channels is implicated in the function of centres responsible for the exchange of larger particles across membranes [37]. In the present model for a membrane domain we focus on the survival behaviour of particles arriving to the boundary from the centre surface (in–out case) or arriving to the centre surface from the domain boundary (out–in case). To examine this behaviour we consider the survival probability $S(t)$ that a tracer started either at the centre or at the boundary, and does not attain a distance ($a + r$) away from the centre up to time t . This standard definition of the survival probability is directly related to the probability density of the first arrival time $\wp(t)$ of the tracer to that distance [116–118].

Figure 10 shows $S(t)$ for homogeneous and heterogeneous MMC. For both homogeneous case and in–out diffusion for HDC we observe at intermediate times

$$S(t) \simeq t^{-1/2}. \quad (17)$$

In turn, for out–in diffusion in HDC $S(t)$ has only a very weak dependence on the diffusion time t . As the distance ($a + r$) increases the survival probability starts to follow the scaling behaviour at later times, figure 10. Similar scaling relations were obtained for subdiffusive heterogenous diffusion processes with a diffusivity



$D(r) = D_0 A / (A + r^2)$ for in-out diffusion of particles [84]. We also simulated the tracer diffusion in the annulus without crowders but with the effective local radial diffusivity $D(r)$ presented in figure 6(C). In these simulations, the same scaling law (17) for $S(t)$ was obtained, compare figures 10 and 11.

Figure 10 shows that the survival probability appears to saturate to finite values, instead of decaying to zero. The non-zero limiting value of $S(t)$ is related to the disorder averaging and should be equal to the fraction of HDC in which the circle of radius $a + r$ is not accessible for a finite size tracer. In fact, for such configurations the survival probability is 1, and it is then weighted by the fraction of these configurations. The behaviour of $S(t)$ is the sixth main result of this study.

The probability density $\varphi(t)$ of arrival times of the tracer particle at the inner and outer boundaries of a given domain for the particle diffusion in homogeneous and heterogeneous mixtures of crowders is shown in figure 12. It reveals a decay

$$\varphi(t) \propto t^{-3/2} \quad (18)$$

at intermediate time scales when the scaling for $S(t)$ is very clear; namely, at times $t \sim 10^2 \dots 3$. The observed scaling of $\varphi(t)$ is thus consistent with the general relation $\varphi(t) = -dS(t)/dt$ known between the two quantities [118]. A better statistic and larger domain size might, however, be needed in simulations to unambiguously validate the precision of the scaling laws (17) and (18) and the regimes of their temporal validity.

7. Discussion and conclusions

The dynamics of the lateral mobility of individual lipid molecules in a lipid bilayer as well as of membrane embedded proteins is of fundamental importance for the function of biological membranes. For instance, the lipid demixing to form domains and patterns or protein complex formation are considered to be elementary prerequisites for membrane associated processes. Here we studied a schematic two-dimensional model system for such processes: small particles (the lipids or small embedded molecules) with a finite size diffuse in a static array of large particles (the crowding proteins or protein complexes) at different crowding densities and crowder distributions. We observed several remarkable effects that are of interest from a physical point of view but will also impact the study of (quasi) two-dimensional crowded systems such as lipid bilayer membranes or fluidic systems such as confined emulsions or colloids.

The spatial heterogeneity of physical systems is often modelled through space-dependent diffusivities $D(x)$. These approaches include deterministic prescribed shapes of $D(x)$ [82–87] corresponding to the systematic variation of the local particle diffusivity observed in living cells [65, 67]. Concurrently $D(x)$ can be taken as a spatially and temporally random quantity [119, 120]. Both kinds of processes may lead to non-stationary anomalous diffusion exhibiting ageing even at long observation times [60]. Our extensive Langevin dynamics simulations of tracer diffusion in a finite heterogeneous environment reveal transient anomalous diffusion with a number of features that are similar to these idealised stochastic models.

For point-like diffusers one would expect that smaller crowders impede the diffusion less, as the tracer can always pass through even small gaps between the crowders. In turn, the point tracer would have to navigate around larger crowders and thus be affected more severely. As we showed here the opposite effect occurs for a tracer of a finite size, comparable to the size of the crowders. In this case small crowders significantly hamper the spreading of the tracers as compared to larger crowders at the same crowding fraction ϕ . Moreover, the van Hove correlation function acquires an exponential shape at higher ϕ and small crowders, whereas this function is nearly Gaussian and quite insensitive to the crowding fraction for large crowders. While one would expect this effect to be somewhat mitigated by a finite yet small mobility of the crowders recent simulations studies of

protein crowded lipid bilayers indicates that the motion is very similar to that of particles diffusing in static obstacle arrays [43].

Concurrent to this effect we observe for larger, homogeneously distributed crowders an extended Brownian regime of the ensemble averaged MSD, whose magnitude is almost independent of the crowding fraction within the investigated range. The time averaged MSD in this case is also highly reproducible. For smaller crowders, however, anomalous diffusion effects in the ensemble averaged MSD occur and become more severe as ϕ increases. Additionally, individual time averaged MSD curves demonstrate the early immobilisation of the tracer particles in the quenched landscape, corresponding to the crossing of a local percolation threshold for the tracer motion.

Similar effects were observed for HDCs. Due to the deterministic gradient of the crowder distribution, we distinguished the two scenarios of in–out and out–in diffusion. The in–out case resembles in many aspects the homogeneous case: the spread of the time averaged MSD is small, the van Hove correlation function is close to Gaussian and weakly depends on the crowder size, and the survival probability exhibits the characteristic square root decay at intermediate times. In contrast, the out–in case is marked by a highly non-Gaussian diffusion of the tracer: high values of the non-Gaussianity parameter and a pronounced cusp in the van Hove correlation function near $\Delta x = 0$ reveal a prevalence of small displacements. This is characteristic for subdiffusive processes. Moreover, the survival probability exhibits a very slow decay and tends to saturate at large values reflecting the dominance of crowder configurations that block the tracer near the release point at the domain boundary. As a consequence, the computed averages over tracer trajectories are dominated by few successful translocation events.

Could some of the observations of our two-dimensional system also be relevant for the diffusion of particles in the cytoplasm of biological cells? Many eukaryotic cells are often adhered onto surfaces and reveal a quasi two-dimensional, *fried egg* shape [21, 31, 65, 84]. The vertical cell thickness may be as small as several microns or less, as compared to tens of microns in the lateral dimension of non-adhered cells. This enables the tracking of tagged diffusing particles for longer distances, for instance, by conventional fluorescence microscopy when the particles need to stay close to the focal plane [121–123]. Still, we emphasise that for relatively small tracer particles (such as small cellular proteins), the cell thickness even in its flat regions is much larger than the average inter-crowder separations. However, larger diffusing particles may be impeded by crowders such as vesicles, often found in great abundance in certain cell types such as insulin producing cells [20] or amoeba [35]. In these *supercrowded* [35] cases our results could well be applicable albeit with a renormalised tracer particle size to accommodate for some possible overlap due to the third dimension.

In the present study we neglected hydrodynamic interactions between the tracer and the crowders. In three-dimensional systems this may affect the long time diffusion [2, 50, 124–126]. The slow, $1/r$ decay of this hydrodynamic coupling implies that a diffusing particle is impacted by crowders from a finite distance that helps avoiding to collide with the crowders. In the two-dimensional membrane system motivating our study the lipid molecules themselves represent a fluid background in which the crowding proteins are embedded. Water molecules are typically expelled from the bilayer and only wet its surfaces.

It will be of interest to extend our results to three-dimensional crowded situations, in particular, to the superdense [1] cytoplasm of biological cells. There, the crowding induced changes in the diffusion of signalling molecules is expected to impact gene regulation and metabolism [127] and also alter active transport properties [21, 33, 34, 128]. Other extensions of the current model include polydisperse static and dynamic crowders [129–131], cylindrical domains mimicking a typical bacterial shape [55, 132], specific and non-specific tracer-crowder interactions [55, 61], three-dimensional computer simulations, a variable tracer size, as well as deformable and non-spherical crowding particles [133, 134].

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