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Collective dynamics effect transient subdiffusion of inert tracers in flexible gel networks

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Abstract

Based on extensive Brownian dynamics simulations we study the thermal motion of a tracer bead in a cross-linked, flexible gel in the limit when the tracer particle size is comparable to or even larger than the equilibrium mesh size of the gel. The analysis of long individual trajectories of the tracer demonstrates the existence of pronounced transient anomalous diffusion. From the time averaged mean squared displacement and the time averaged van Hove correlation functions we elucidate the many-body origin of the non-Brownian tracer bead dynamics. Our results shed new light onto the ongoing debate over the physical origin of steric tracer interactions with structured environments.

Keywords: anomalous diffusion, gel network, van Hove correlation

1. Introduction

Modern single particle tracking technology \cite{1} unveils the non-Brownian stochastic motion of submicron tracers or fluorescently labelled macromolecules in a variety of complex liquids. Anomalous diffusion of the form \( \langle r^2(t) \rangle \approx t^\alpha \) with \( 0 < \alpha < 1 \) \cite{2} was observed in the...
cytoplasm and membrane [3–5] of living biological cells, as well as in dense polymer or protein solutions [6]. Causes for the observed subdiffusion may be the crowded state of the environment [7] or sticking effects between the tracer and the environment [8]. Another important origin for anomalous diffusion is the subtle interaction between the tracer particle and the dynamic confines of a structured, gel-like environment with a well defined mesh size [9].

Mesh-like, structured environments are a defining property in various systems: biological cells are equipped with a characteristic mechanical network consisting of actin and other biofilaments through which submicron particles diffuse or are actively transported [10]. Inside eukaryotic nuclei, biomolecules diffuse through the complex chromatin network [11]. Particles in a biofilm move through a flexible, porous bacterial matrix [12], and in cellular tissues the intercellular space is filled with a mesh-like extracellular matrix [13]. In novel clinical diagnosis tools pathogens diffuse in a hydrogel [14].

How do particles diffuse through a flexible, fluctuating mesh such as a hydrogel? When the particles are much smaller or larger than the typical mesh size, they diffuse normally or are fully immobilized [9]. We here consider the physically interesting case when the size of the particle is comparable to the typical mesh size (figure 1). From extensive Brownian dynamics simulations we elucidate the micro- and mesoscopic principles behind the complex tracer dynamics. The time averaged mean squared displacement (TAMSD) and the van Hove cross-correlation function (TAHCF) demonstrate the occurrence of significant tracer subdiffusion (figure 2(a)) and cooperative mesh fluctuations, allowing relatively large particles to move in the gel albeit at massively reduced diffusivity.

The tracer motion in a flexible mesh can be viewed as a convolution of the Brownian self-dynamics of the tracer and the thermal agitation of the interacting gel particles. Important ingredients characteristic for the tracer-gel system are the steric obstruction of the tracer by the
mesh [15], tracer-gel interactions [16], and the elastic response of the network itself [17]. In an effective continuum view, these features translate into the local viscoelasticity [18], which is probed in microrheology experiments [19]. In this picture, memory effects in the dynamics cause an anomalous frequency dependence of the generalized friction coefficient and hence, expressed in terms of the generalized Langevin equation, an anomalous diffusion of the tracer. Remarkably, distinct anomalous tracer diffusion was found in numerous experiments [9, 16, 20–22] and simulations [23]. To explain this non-Brownian dynamics in the gel two scenarios are typically invoked: (i) trapping of the tracer due to attractive tracer-gel interactions, that effect transient binding to the gel network [16, 20, 24, 25], such that the tracer intermittently follows the gel’s Rouse dynamics [20] or remains transiently immobilized [24]. (ii) Sterical hindrance within a fractal organization of the accessible volume [9, 16, 21–23, 26]. In these approaches the gel is represented by impenetrable beads (static [23] or dynamic [26]) on a lattice. In static networks

Figure 2. (a) TAMSD for tracers of diameter \(\sigma_t = 3, 4, 5\) (top to bottom) versus lag time \(\Delta\), and \(\varphi_D = 20\) (circles) and \(\varphi_D = 60\) (triangles). Intermediate-asymptotics \(\overline{\delta^2} \approx \Delta^\mu\) (full lines) shown for \(\mu = 0.85, 0.8,\) and \(0.4\), for increasing tracer size. Dashed and full black lines denote a linear scaling and a plateau. Each curve is a time average over \(T = 10^4\) and additional average over 10 independent realizations. Inset: magnification of the last decade of \(\Delta\) in the case of \(\sigma_t = 3\). The transition from the subdiffusive \(\overline{\delta^2} \approx \Delta^\mu\) (green line) to diffusive \(\overline{\delta^2} \approx \Delta\) (dashed black line) regime is nicely visible. (b) Gel correlation \(g_{\text{gel}}(r)\) for \(\varphi_D = 60\) (blue dashed) and \(\varphi_D = 20\) (full black). Inset: distribution of bond length (b), face diagonal (fd), and body diagonal (bd) for both stiffnesses.
simulations showed that isolated, randomly positioned beads effect non-transient subdiffusion, whereas tracer diffusion in the presence of local bead clusters in the form of randomly distributed, condensed cubes is only transiently anomalous [23]. The debate on the dominant mechanism in real gels remains open [16]. Notably, both of the above mechanisms are only explained on a heuristic level and a better physical understanding is needed.

2. Model

While lattice representations of gels are a computationally efficient approach for the modelling of tracer diffusion [23, 26], they suffer from the severe localization of the gel despite its exposure to a thermostat. Moreover, static gels introduce spurious artifacts in the tracer dynamics, as quenched configurational disorder of obstacles introduces a persistent memory in the motion of a Brownian particle at all obstacle densities [27]. As shown here, the tracer motion is inherently related to collective soft modes of local expansion and contraction of the thermalized mesh. Such modes cannot be appropriately captured in lattice models, calling for more realistic models with fully coupled, many-body tracer-gel dynamics.

We consider a gel of $N_g = 7 \times 7 \times 7$ connected beads with a simple cubic unit cell and a freely diffusing tracer bead (figure 1). Tracer and gel beads are impenetrable via the repulsive part of the Lennard-Jones potential. Nearest neighbour gel beads are connected via Morse springs. The size $a$ of the unit cell, the equilibrium centre-to-centre distance between nearest neighbours beads, is 4 times the gel bead diameter $\sigma_g$, and the gel density is $\rho = a^{-3}$. We use three tracer bead diameters, $\sigma_t = 3\sigma_g$, $4\sigma_g$ and $5\sigma_g$. The average face and body diagonals (fd and bd) of the unit cell are then equal to $5.66\sigma_g$ and $6.93\sigma_g$. Thus, the smallest tracer comfortably passes through a static mesh, while the middle size is already similar to the value $4.66\sigma_g$ ($= fd - \sigma_g$). The largest bead still fits the unit cell as set by $bd - \sigma_g$. However, it can no longer pass through the static mesh. We express distances in units of $\sigma_g$, energy in units of thermal energy $k_B T$ and time in units of $\sigma_g \epsilon_g$, where $\epsilon_g$ is the energy scale of the repulsive LJ potential. The tracers’ diffusion coefficient is given by the Stokes–Einstein relation, and the gel stiffness is set by the width $\zeta = 0.6$ and depth $\varphi_D$ of the Morse potential. We consider two depths resulting after shifting due to the cut-off in the potential minima at $-7.5$ and $-22.9 k_B T$. The two values represent a moderately soft and a rather stiff gel (see figure 2(b)). The dynamics of both tracer and gel beads are governed by overdamped Langevin equations with Gaussian white noise. After extensive equilibration we simulate trajectories of length $T = 10^4$. More details on the model and simulations are found in the appendix.

3. Analysis of the tracer dynamics

The gel–gel position correlation

$$g_{gg}(r) = \frac{1}{4\pi r^2 \rho} \int_0^T \sum_{i<j} \delta \left( \left| r_i^g(t) - r_j^g(t) \right| - r \right) dt,$$

see figure 2(b), confirms the simple cubic structure of the gel: the first three peaks correspond to $a$, fd and bd, where the latter is not clearly observed in the softer gel due to the more pronounced thermal fluctuations. The length fd is smeared out over the interval 4–7 causing a larger effective gel bead size. Comparing the structural features of the gel in the presence and
absence of the tracer we see no appreciable difference, indicating a large effective tracer
dilution, that is, the length-scale of deformation is small compared to size of the system.

We use the TAMSD at lag time $\Delta$ over a trajectory of length $T$,

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

(2)
to characterize the tracer dynamics. For a wide parameter range $\overline{\delta^2}$ exhibits a distinct transient
subdiffusion regime between the initial free diffusion and asymptotic long time diffusion
(figure 2(a))\(^5\), in agreement with similar simulations [28]. The stiffness only affects the values
of the short and long time diffusivities and the time window of the intermediate-asymptotic
subdiffusion regime (circles and triangles of same colour in figure 2(a)).

Evidently a pronounced subdiffusive power-law regime $\overline{\delta^2} \approx \Delta^\mu$ emerges for $\Delta > 1$, and
the anomalous diffusion exponent $\mu$ delicately depends on the tracer size but, remarkably, is
almost independent of the gel stiffness, except for a terminal scaling for the largest tracer which
completely localizes within the unit cell. Intuitively, a smaller tracer, which comfortably passes
through a unit cell surface in a static gel, diffuses faster in a stiffer gel due to the smaller
effective size (smaller fluctuations around the equilibrium position) of the gel beads. Clearly,
as the tracer diameter reaches 4 the situation is reversed: now the effective size of the gel beads
prevents an unobstructed passage of the tracer and larger gel fluctuations allow passage events.
We stress that $\overline{\delta^2}$ is a weighted average of squared relative displacements along the trajectory.
Hence, $\overline{\delta^2} < a$ does not necessarily mean that the tracer remains localized within the same
unit cell.

To gain a deeper understanding we study the TAHCF

$$G_t(\chi, \Delta) = \frac{1}{T - \Delta} \int_0^{T - \Delta} \frac{\delta[|r(t + \Delta) - r(t)| - \chi]}{4\pi\chi^2} dt,$$

(3)

where $4\pi\chi^2 G_t(\chi, \Delta)$ is the probability density to find the tracer at time $\Delta + t'$ at a distance
between $\chi$ and $\chi + d\chi$ from its position at time $t'$, averaged over all instantaneous positions
along the trajectory. The usual ensemble van Hove self correlation function (HCF) measures
position correlations at time $t$ with the initial position at time $t = 0$ over an ensemble of
independent realizations. The TAHCF measures correlations between positions at different
times along the same trajectory (similar to the difference between the MSD and the TAMSD)\(^6\).
A broad or multimodal HCF denotes an inherently heterogeneous tracer population, while the
same features in the TAHCF reveal temporally heterogeneous dynamics. The TAHCF can be
thought of as a distributional analysis of single particle traces.

Figure 3 shows the results of $4\pi\chi^2 G_t(\chi, \Delta)$. Strikingly, for $\sigma_i = 3$ and 4 already at time
lags $\Delta \sim 2 - 5$, where the relative displacement along the trajectory is mostly $\chi \lesssim a$, we find
that the tracer is occasionally displaced over several $a$ and hence visits several new cells. Thus,
while it spends most of the time fluctuating within the confines of a cell it occasionally ventures
into new cells. The stiffness of the gel does not affect the qualitative dynamics beyond the fact
that transport is more efficient in stiffer gels if the tracer is small enough and vice versa for

\(^5\) For $\sigma_i = 4$ terminal normal diffusion is reached approximately at $\Delta \sim 150 - 200$ while in the case of $\sigma_i = 5$ a
clear crossover to linear scaling was not observed after averaging over a trajectory of length $T = 10^4$. For $\Delta > 100$
the results become progressively noisier due to deteriorating statistics in the time average.

\(^6\) In fact, $\overline{\delta^2}$ is the second moment of $G_t(\Delta, |\delta t|)$. 

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larger tracer size. The largest tracer, which can only escape a cell at an extreme gel fluctuation, occasionally visits other cells at a much later point $\Delta \sim 50$ and only for the softer gel. The stiffer gel does not allow sufficiently large fluctuations on the observed time scale and the tracer remains localised in the same cell, fluctuating around its mean position (figure 3 right panel). Possibly, an escape may occur at longer time scales. How can we reconcile the intermediate anomalous scaling for all tracer sizes with the different statistics of relative displacements?

4. Collective gel-tracer dynamics

The escape from a mesh cell is slow on time scales of the anomalous diffusion regime, effecting a local violation of the central limit theorem in this regime. What is the microscopic mechanism? Are the tracer’s cell-to-cell passages simply governed by an averaged obstruction effect and thus decoupled from the gel dynamics, or by many-body tracer-gel interactions? We address this question via the tracer-gel TAHCF

$$G_{tg}(r, \Delta) = \frac{C^{-1}}{T - \Delta} \int_0^{T - \Delta} \sum_{i=1}^{N_g} \delta\left(\|\mathbf{r}_i(t + \Delta) - \mathbf{r}(t)\| - r\) \right) dt,$$

where the sum $i$ includes all gel beads of coordinate $\mathbf{r}(t)$ and $C^{-1}$ normalizes the probability density to observe a gel bead infinitely far away from the initial tracer position to unity at any lag time, i.e. it corresponds to the ratio of the uncorrected correlator and the number density $\rho$ of beads. $G_{tg}(r, 0)$ is the static gel-tracer pair correlation function. Equation (4) effectively measures the memory loss of the environment of the instantaneous tracer location and thus reflects the dynamic correlations of the gel beads and the tracer. As $\lim_{\Delta \to \infty} G_{tg}(r, \Delta) = 0$ while $\lim_{r \to \infty} G_{tg}(r, \Delta) = \rho$, we omit the prefactor $4\pi r^2$ to more clearly observe the relative motions of the gel beads. As for the self-part, the meaning of $G_{tg}(r, \Delta)$ differs from the ensemble picture. The present weighted average along the tracer trajectory integrates out all effects of the tracer

Figure 3. $4\pi r^2 G_{tg}(r, \Delta)$ for tracer sizes $\sigma = 3$ (left), $\sigma = 4$ (middle), $\sigma = 5$ (right). Black dashed lines represent the stiffer gel. Each curve is a time average over $T = 10^4$ with additional average over 10 independent realizations.
motion such that $G_{tg} (r, \Delta )$ truly measures relative motions of gel beads. We expect that the instantaneous tracer position is dominated by statistically likely configurations and ‘transition states’ (locations within a cell face) have a smaller statistical weight. Gel beads should thus be found at characteristic distances from the tracer.

Figure 4 shows results for $G_{tg} (r, \Delta )$ for different tracer sizes. The first peak corresponding to nearest gel beads shows a high initial correlation due to frequent collisions. Entropic effects smear out the peaks around their mean (we typically expect to find gel beads in the shaded areas if the tracer is moved along the bd up to contact with a bead). Since the gel structure on average is well defined the tracer mostly moves along ‘paths of free volume’ while constantly colliding with gel beads, resembling hopping transport [29]. According to figure 4, for $\Delta < 1$ gel beads diffuse rather freely, effecting the broadening of the main peak, which also becomes more symmetric. Similar trends hold for the second peak corresponding to next-nearest neighbours. On time scales $\Delta \gtrsim 1$, for small and intermediate tracer sizes some of the nearest gel beads exhibit excursions towards $r \to 0$ made possible by exceeding tracer fluctuations within a cell. Concurrently, below $\Delta \sim 15$, the peak positions of the nearest and next-nearest neighbours drift towards a larger $r$, indicating cooperative tracer-cage fluctuations. At these time scales the inter-peak distance remains constant, suggesting that bond lengths are maintained near their

\begin{figure}
\centering
\includegraphics[width=\textwidth]{figure4.png}
\caption{TAHCF for different tracer sizes and lag times, for $\varphi_D = 20$ and (a) $\sigma_t = 3$, (b) $\sigma_t = 4$, and (c) $\sigma_t = 5$. For clarity the plots are divided into two panels for different $\Delta$. Dashed lines denote the positions of vertices for a static gel if the tracer sits in the centre of a cell. In the shaded areas we would find the vertices if the tracer is displaced diagonally up to the point of contact with a gel bead. Corresponding results for $\varphi_D = 60$ are shown in figure 5. Each curve represents a time average over $T = 10^4$ and 10 independent realizations.}
\end{figure}
equilibrium values and cage fluctuations are in fact mediated by collective displacements. Strikingly, on time scales $15 \leq \Delta < 55$ (and up to 75 for $\sigma_t = 4$) both peaks simultaneously move towards smaller $r$ while the value at $r \to 0$ simultaneously increases, indicating cell compression. At even longer $\Delta$, the first and second peak move towards smaller and larger $r$, indicating that the tracer particle escaped the cell\textsuperscript{7}. We stress that $G_{n}^{r\tau}(r, \Delta)$ is a weighted average over escape and blocking events, such that it is natural to expect that the result will contain a progressively smaller contribution of escaped configurations as the tracer size grows. This is exactly what is observed here. Moreover, as the tracer size grows the magnitude of fluctuations decreases, indicating that the tracer locally stiffens the network\textsuperscript{8}. The same qualitative features are also observed for the stiffer gel with the sole exception that for the largest tracer the position of the second peak does not move at all while the position of the first peak still oscillates at long times. This suggests localized fluctuations within the same cell, in agreement with the results for $G_{n}^{r\tau}(r, \Delta)$. As shown in figure 5, due to the higher gel stiffness the peaks are more pronounced, as expected. A distinct feature is indeed the strong localization of

\textsuperscript{7} For $\sigma_t = 3$ we even observe a recoil at $r \to 0$, suggesting that as the particle leaves the cell the originally innermost gel beads are entropically pulled back as they have changed their topological position with respect to the tracer. Moreover, the peaks at $r \to 0$ are a consequence of the fact that the spherical volume element diverges.

\textsuperscript{8} This is similar to the effect of fillers in rubber networks [30].
the largest tracer due to the fact that the gel stiffness is too large to allow sufficient cage-opening fluctuations needed for a particle to escape a cell on the investigated time scale.

5. Discussion

To rationalize these observations, we note that the gel-bond fluctuations are much faster than the cell-to-cell tracer motion. Hence, it is reasonable to assume that the gel is locally equilibrated corresponding to a free energy minimum, i.e. gel beads are on average at equilibrium bond-distance concurrent with extensive mutual fluctuations of nearby bead pairs\(^9\). In such a case larger local rearrangements in the gel necessarily have to be collective, otherwise they would be thermodynamically penalised. However, such collective fluctuation events demand many-body correlations and thus come at an entropic cost and are thereby statistically less probable. Hence, correlated fluctuations of gel bead positions on the many-body level may effect the slow cage-opening dynamics and thereby the transient subdiffusion of tracers with size comparable to the mesh size.

These results enable us to draw a full physical picture for the emergence of transient anomalous diffusion in flexible gels. For tracer sizes comparable to the gel mesh the probability of collision with the gel is very high. The rate-determining step is the cell-to-cell passage of a tracer, demanding cage opening fluctuations, which were shown to be collective. Open cage states are thus entropic bottlenecks and cause slow tracer motion on intermediate time scales and violation of the central limit theorem. A finite characteristic escape time \(\tau_{esc}\) restores normal diffusion at long time scales with an effective diffusion coefficient \(D_{eff} \approx \frac{d^2}{\tau_{esc}}\). Tracers considerably larger then the mesh size are expected to be completely localized in a cell, as seen in experiment [9]. At this point, a few remarks are in order. First, on a time scale on which correlations of the gel bead positions have all died out, the model gel system behaves as an array of ordered static beads and the gel does not exhibit any features of glassy dynamics. This is not necessarily the case in real gels [32]. Second, even though hydrodynamic interactions are believed to be effectively screened in such a dense system [33], direct solvent effects might nevertheless become important.

To conclude, extensive Brownian dynamics simulations of tracer particles in a flexible, thermally agitated gel in conjunction with a single trajectory analysis of the tracer dynamics and many-body correlations demonstrate that the distinct transient subdiffusion, frequently observed in experiments and simulations, is effected by the collective fluctuations of both the tracer and the vicinal gel beads. On the scale of the typical escape time of the tracer from a unit cell the dynamics of nearby gel beads turns out to be appreciably correlated. We here obtained compelling evidence identifying the physical origin of the anomalous diffusion, namely, collective fluctuations in the entangled tracer-gel dynamics representing an entropic bottleneck and thereby slowing down the tracer dynamics. This physical insight obtained from the time average van Hove correlation functions provides a significantly improved understanding of the elusive problem of tracer-gel interaction in the experimentally important limit when the tracer size is comparable to the typical mesh size. On a continuum level, these findings translate into the viscoelastic properties of the gel and thus establish a link with the insight obtained from microrheology [19]. We are confident that these results will inspire new experimental and theoretical investigations of tracer motion in structured matrices.

\(^9\) A similar situation is encountered in hydrogen-bond dynamics in water around a hydrophobic particle; see [31].
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Appendix

We consider a gel of $7 \times 7 \times 7$ connected beads forming a simple cubic lattice with periodic boundary conditions and a freely diffusing tracer bead. The gel beads are allowed to move off-lattice in continuous space. Each gel bead interacts with other gel beads as well as with the tracer with the repulsive part of the shifted Lennard-Jones (LJ) potential

$$V_{ij}^{LJ}(r_{ij}) = 4\epsilon_{ij} \left[ \left( \frac{\sigma_{ij}}{r_{ij}} \right)^{12} - \left( \frac{\sigma_{ij}}{r_{ij}} \right)^{6} \right] \Theta \left( 2^{1/6}\sigma_{ij} - r_{ij} \right) + \epsilon_{ij}, \quad (A.1)$$

where we use a two-index notation for compactness. In expression (A.1) $i$ and $j$ for a specific pairwise interaction take on the values $t$ for the tracer particle and $g$ for gel beads. $r_{ij} \equiv |\mathbf{r}_i - \mathbf{r}_j|$ is the physical separation between $i$ and $j$. Using combining rules, $\sigma_{ij} = \frac{1}{2}(\sigma_i + \sigma_j)$ are the separations at zero unshifted potential, i.e. $\sigma_{ii}$ is the diameter of species $i$ ($\sigma_g$ or $\sigma_t$ in the notation of the main text). $\epsilon_{ij}$ (with $\epsilon_{gg}$ equalling $\epsilon_g$ used in the main text) sets the respective energy scales, and $\Theta(x)$ is the step function. Each gel bead is additionally connected with its six nearest neighbours with a Morse potential (figure A.1)

$$V_{ij}^{M}(r_{ij}) = \varphi_D \left( \epsilon^{-2\zeta\phi} - 2\epsilon^{-\zeta\phi} \right) \Theta \left( r_c - r_{ij} \right) - V_s, \quad (A.2)$$

where $\zeta = \sqrt{\frac{k_c}{2\varphi_D}}$ determines the width of the potential. $\varphi_D$ is the potential depth and thus sets the energy scale, $k_c$ is the force constant at the potential minimum, and $d^r \equiv r_{ij} - r_{eq}$, and $r_{eq}$ is the equilibrium separation. $r_c$ is a cutoff distance (taken here as $1.4a$) and $V_s = V_M^G(r_c)$. In general, $\zeta$ controls the width of the potential well. The inclusion of LJ repulsion in Morse-bound

![Figure A.1. The two Morse potentials used in the simulations. See text for details.](image-url)
neighbours is necessary to prevent potential overlaps caused by rare, but still possible, exceedingly large thermal fluctuations where the Morse potential might not be sufficiently stiff.

The particle positions are governed by the overdamped Langevin equations

\[ \frac{d\mathbf{r}_i(t)}{dt} = \beta D_i \sum_{j \neq i} \mathbf{F}_{ji} + \sqrt{2D_i} \xi_i(t), \]  

(A.3)

where \( D_i \) is the diffusion coefficient of the tracer or gel beads, \( \beta^{-1} = k_B T \), and

\[ \mathbf{F}_{ji} = -\nabla_{r_{ij}} U_{ij}(r_{ij}) \]  

(A.4)

is the force exerted on particle \( i \) by particle \( j \), and equals the sum of the negative gradients of the Lennard-Jones potential (A.1) and the Morse potential (A.2), respectively. \( \xi_i(t) \) is a delta-correlated Gaussian noise with zero mean and component-wise variance

\[ \langle \xi_k(t) \xi_{i'}(t') \rangle = \delta(t - t') \delta_{k,i}. \]  

(A.5)

Since hydrodynamic interactions are screened in dense systems [33] we neglect them in our model. We use dimensionless units and express distances in units of the gel bead diameter \( \sigma_g \), energies in units of \( \beta^{-1} \) and diffusivities in units of \( \sigma_g/\sqrt{\epsilon_g} \), thus fixing the time unit \( t_1 = 1 \). We take \( \epsilon_g = \epsilon_{gg} = 1.2 \) and \( \epsilon_{gt} = \epsilon_g \). We choose \( r_{eq} = 4 \), which also fixes the gel unit cell size \( a = r_{eq} \). Finally, we consider various sizes \( \sigma_t \) of the tracer particle, \( q_D \) and spring stiffness \( k_e \).

While we performed simulations using a wide span of different parameters we here focus on two representative cases and fix \( \zeta = 0.6 \) and use \( q_D = 20 \) to describe the softer gel, and \( q_D = 60 \) to describe the stiffer gel. The value of \( \epsilon_g \) is essentially irrelevant as long as it is not too big, because it only affects the softness of the gel-tracer collision. Moreover, after testing for several values we choose \( r_c = 1.4d_g^c \) for the cutoff of the Morse potential. A further increase of this cutoff does not affect the results, but naturally makes the simulation slower. The obtained shifted potentials are shown in figure A.1. Such a choice of Morse parameters represents a well structured gel (see figure 2(b)), with roughly 5\% of bonds dissociated on average. The resulting effective stiffness of the gels is best seen from the fluctuations of the unit cell parameters (figure 2(b)). These cases in fact suffice to demonstrate the qualitative physical picture of transient subdiffusion.

The scale of the free tracer diffusivity is set by the Stokes–Einstein relation, \( D \approx 1/\sigma_t \). The Langevin equations are solved with the Euler method with integration step \( 10^{-4} \) under periodic boundary conditions. A small integration step is needed because of the high collision frequency in cases where the tracer is comparable to the mesh size. The initial configuration is chosen randomly to be slightly displaced from the energetically minimal configuration, not allowing significant overlap nor significant bond stretching. After extensive equilibration time of \( 10^4 \), trajectories of length \( 10^4 \) are simulated and analyzed. The overall time to simulate and analyze a single trajectory of length \( T = 10^4 \) was 5–6 weeks of single-CPU time on a computer cluster, and our study thus reaches the current computational limit.

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