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Rotational and translational self-diffusion in concentrated suspensions of permeable particles

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Abstract

In our recent work on concentrated suspensions of uniformly porous colloidal spheres with excluded volume interactions, a variety of short-time dynamic properties were calculated, except for the rotational self-diffusion coefficient. This missing quantity is included in the present paper. Using a precise hydrodynamic force multipole simulation method, the rotational self-diffusion coefficient is evaluated for concentrated suspensions of permeable particles. Results are presented for particle volume fractions up to 45%, and for a wide range of permeability values. From the simulation results and earlier results for the first-order virial coefficient, we find that the rotational self-diffusion coefficient of permeable spheres can be scaled to the corresponding coefficient of impermeable particles of the same size. We also show that a similar scaling applies to the translational self-diffusion coefficient considered earlier. From the scaling relations, accurate analytic approximations for the rotational and translational self-diffusion coefficients in concentrated systems are obtained, useful to the experimental analysis of permeable-particle diffusion. The simulation results for rotational diffusion of permeable particles are used to show that a generalized Stokes-Einstein-Debye relation between rotational self-diffusion coefficient and high-frequency viscosity is not satisfied.

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I. INTRODUCTION

The rotational and translational self-diffusion of interacting colloidal and macromolecular particles suspended in a low-molecular-weight solvent is the subject of ongoing research both experimentally and theoretically [1, 2]. Originally, self-diffusion in dilute systems was studied, however the center of interest has shifted since to concentrated dispersions where solvent-mediated many-particle hydrodynamic interactions (HIs) are of central importance. An example, of biological relevance, is self-diffusion of proteins and other macromolecules in the crowded environment of a cell [3].

Two central quantities quantifying the configuration-averaged influence of HIs on the suspensions dynamics are the concentration-dependent short-time rotational and translational self-diffusion coefficients D_r and D_t , respectively. At zero particle concentration, these quantities reduce to the single-particle diffusion coefficients D_0^r and D_0^t . For solvent-impermeable colloidal hard spheres with stick hydrodynamic surface boundary conditions, the single-particle coefficients are given by

$$D_0^{r,\text{hs}} = \frac{k_B T}{8\pi\eta_0 a^3} \tag{1}$$

$$D_0^{t,\text{hs}} = \frac{k_B T}{6\pi\eta_0 a} \,, \tag{2}$$

with η_0 the Newtonian solvent shear viscosity, Boltzmann constant k_B , temperature T and hydrodynamic particle radius a. The influence of the HIs at non-zero concentrations gives rise to values for D_r and D_t smaller than their respective values D_0^r and D_0^t at infinite dilution. The short-time coefficients describe self-diffusion on the time scale $t \ll a^2/D_0^t$, but with t large enough that solvent and particle velocity correlations have decayed. On the colloidal short-time scale, the concentration dependence of D_r and D_t is determined by averaging the HIs with the equilibrium particle distribution.

Self-diffusion coefficients in colloidal suspensions have been determined experimentally by a variety of techniques. The mean-squared displacement (MSD) of partially solvent-index matched suspensions of colloidal spheres [4, 5] has been measured as a function of time using dynamic light scattering (DLS), with D_t determined from the initial slope of the MSD.

For the vast majority of systems where this specialized index-matching technique is not applicable, D_t may be inferred, to decent accuracy according to theory and simulation [6–8], from a first cumulant analysis of the scattered light electric field autocorrelation function, probed at a scattering wavenumber larger than the peak location of the static structure factor where the structure factor attains the value one [9–11]. Translational long-time self-diffusion coefficients not considered in the present work can be determined using, e.g., forced Rayleigh scattering [12, 13], fluorescence recovery after photobleaching (FRAP) [2], and fluorescence correlation spectroscopy [14].

Experimental studies of rotational colloidal self-diffusion are based on techniques which can distinguish different particle orientations. Methods which have been used for this purpose are depolarized dynamic light scattering (DDLS) on optically anisotropic particles [15], and nuclear magnetic resonance [16]. More recently developed techniques applicable to a larger variety of systems include time-resolved phosphorescence anisotropy [17, 18] and polarized FRAP [19, 20] measurements. The latter methods have been carried out using fluorophore-labeled colloidal particles. Most of the published experimental results deal with self-diffusion properties of monodisperse colloidal systems. However, experimental and theoretical work has been also performed on rotational diffusion in colloidal mixtures, in particular for binary systems where one component (the tracer) is very dilute [17, 18, 20, 21]. In addition, DDLS measurements of the rotational diffusion of tracer spheres in a polymeric solution have been used to infer viscoelastic properties from a frequency-dependent generalized Stokes-Einstein-Debye (GSED) relation [22].

From a simulation methods viewpoint, short-time rotational self-diffusion in monodisperse colloidal systems of non-permeable spheres with excluded volume interactions was studied, e.g., using lattice-Boltzmann (LB) [23], Stokesian dynamics [24], and accelerated Stokesian dynamics (ASD) simulations [7]. Moreover, virial expansion results of varying accuracy have been derived for the rotational self-diffusion coefficient up to quadratic order in the concentration [15, 25–28].

While there has been no theoretical work so far on rotational self-diffusion in concentrated suspensions of porous particles, other transport properties of porous particles have been studied, including the high-frequency shear viscosity [29–32], and to first order in concentration the mean sedimentation velocity [33].

In our earlier work on the short-time dynamics of concentrated suspensions of uniformly porous particles, a broad spectrum of dynamic properties has been calculated, including the hydrodynamic function [8, 34] and sedimentation coefficient [8], translational self-diffusion coefficient [8, 34], and the high-frequency-limiting shear viscosity η_{∞} [35, 36]. These simulation studies were amended by the derivation of easy-to-use approximate analytic expressions of good accuracy, notably a generalized Saito formula for the shear viscosity [36], and a spherical annulus model approximation for η_{∞} [36], and to first order in concentration also for D_t and D_r [37]. Additionally, precise values for the first-order virial coefficients of D_r and D_t corresponding to two-body HIs have been obtained [37].

In all these studies on permeable particles, the solvent flow inside the spheres is described by the Debye-Büche-Brinkman (DBB) equation [38, 39], and the particles are assumed to interact directly by excluded volume (i.e., hard-sphere type) forces. Our simplifying particle model is specified by two parameters only, namely the particle volume fraction $\phi = (4\pi/3)na^3$, where *n* is the number concentration, and the ratio, *x*, of the particle radius, *a*, to the hydrodynamic penetration depth, κ^{-1} , inside a permeable sphere. Large (low) values of *x* correspond to weakly (strongly) permeable particles. Typical values for *x* in permeable-particle systems, such as core-shell particles, are in the range of $x \sim 30$ or larger [40]. While a specific intra-particle structure is ignored in the model, it is generic in the sense that a more complex internal hydrodynamic structure can be approximately accounted for in terms of a mean permeability. Porous-particle systems of current interest include dendrimers [41–43], microgel particles [44–46], core-shell particles [40, 47–49], and star-like polymers of lower functionality [50].

The present work complements our earlier analysis of the short-time dynamics in concentrated suspensions of uniformly permeable spheres by giving simulation results and a theoretical analysis of the short-time rotational diffusion coefficient not considered so far at non-dilute concentrations. On employing the multipole simulation method of a very high accuracy [51] encoded in the HYDROMULTIPOLE program package [28], we calculate the short-time rotational self-diffusion coefficient, $D_r(x, \phi)$, as a function of x and ϕ . Our results cover the full range of porosities, with the volume fraction extending up to 0.45. In combination with recently obtained tabulated values for the first-order virial coefficients of $D_r(x, \phi)$ and $D_t(x, \phi)$ [37], and precise HYDROMULTIPOLE simulation results for $D_t(x, \phi)$ obtained earlier [8, 34], we show that both $D_r(x, \phi)$ and $D_t(x, \phi)$ can be scaled, in the whole range of permeabilities and volume fractions, to the self-diffusion coefficients $D_r^{hs}(\phi) = D_r(\infty, \phi)$ and $D_t^{hs}(\phi) = D_t(\infty, \phi)$ of non-permeable hard spheres with stick boundary conditions and the same size. From these scaling relations, accurate analytic expressions for $D_r(x, \phi)$ and $D_t(x, \phi)$ are obtained. We expect these expressions to be useful in the experimental data analysis of diffusion measurement on permeable particle systems. The present simulation results for $D_r(x, \phi)$, and known results for $\eta_{\infty}(x, \phi)$, are used to show the violation of a GSED relation between $D_r(x, \phi)$ and $\eta_{\infty}(x, \phi)$, amending our earlier study of similar GSE relations in [35].

The paper is organized as follows: Sec. II provides the theoretical background on shorttime self-diffusion of permeable particles. Furthermore, it includes our simulation results for rotational self-diffusion. The scaling relations allowing to map permeable to non-permeable hard-sphere systems are discussed in Sec. III. In Sec. IV, we complete the scaling relations by providing simple expressions for the scaling functions for non-permeable hard spheres. We also discuss the special case of non-permeable hard spheres in comparison to earlier simulations and experimental work. In Sec. V, we demonstrate the violation of the GSED relation. In our conclusions in Sec. VI, we explicitly write convenient expressions for $D_r(x, \phi)$ and $D_t(x, \phi)$ which should prove useful in practical applications.

II. SHORT-TIME ROTATIONAL SELF-DIFFUSION: THEORY AND RESULTS

Like in our earlier work on the dynamics of permeable particle systems [8, 34–37], we employ a model of uniformly permeable spheres of radius a, dispersed in a Newtonian fluid of viscosity η_0 . The low-Reynolds number incompressible flow inside and outside the spheres is described, respectively, by the Stokes [52, 53] and Debye-Bueche-Brinkman [38, 39] equations

$$\eta_0 \nabla^2 \boldsymbol{v}(\boldsymbol{r}) - \eta_0 \kappa^2 \chi(\boldsymbol{r}) \left[\boldsymbol{v}(\boldsymbol{r}) - \boldsymbol{u}_i(\boldsymbol{r}) \right] - \nabla p(\boldsymbol{r}) = 0.$$
(3)

Here, \mathbf{v} and \mathbf{p} are the fluid velocity and pressure, respectively, and κ^{-1} is the hydrodynamic penetration depth. The characteristic function, $\chi(\mathbf{r})$, is equal to one for the field point \mathbf{r} inside any of the spheres and zero outside. The skeleton of a particle *i*, centered at \mathbf{r}_i , moves rigidly with the local velocity $\mathbf{u}_i(\mathbf{r}) = \mathbf{U}_i + \boldsymbol{\omega}_i \times (\mathbf{r} - \mathbf{R}_i)$, determined by the translational and rotational velocities \mathbf{U}_i and $\boldsymbol{\omega}_i$, respectively. The fluid velocity and stress change continuously across a particle surface.

The short-time rotational self-diffusion coefficient of a quiescent, isotropic system is given in frame-invariant notation by [21, 28]

$$D_r = \frac{k_B T}{3} \left\langle \frac{1}{N} \sum_{i=1}^N \operatorname{Tr} \boldsymbol{\mu}_{ii}^{rr}(\mathbf{X}) \right\rangle, \qquad (4)$$

where $\mathbf{X} = {\mathbf{r}_1, \cdots, \mathbf{r}_N}$ is the configuration of $N \gg 1$ sphere centers, and Tr denotes the trace operation. The hydrodynamic mobility tensor, $\boldsymbol{\mu}_{ii}^{rr}(\mathbf{X})$, linearly relates the torque acting on a particle *i* to its rotational velocity, for zero forces and torques exerted on the other particles. For the present model system, the average $\langle \cdots \rangle$ is taken over an equilibrium distribution of non-overlapping spheres, consistent with the periodic boundary conditions used in our simulations. Our numerical calculation of $D_r(x, \phi)$ makes use of Eq. (4).

The coefficient D_r is a function both of x and ϕ . At infinite dilution, Eq. (4) reduces to [54, 55]

$$D_0^r(x) = \frac{k_B T}{8\pi\eta_0 a^3 \left[1 + \frac{3}{x^2} - \frac{3\coth x}{x}\right]}.$$
 (5)

Note here that $D_0^r(x) > D_0^{r,hs}$ unless $x = \infty$.

We have calculated $D_r(x, \phi)$ to high precision using a hydrodynamic multipole method corrected for lubrication [28, 51, 56, 57], and encoded in the HYDROMULTIPOLE program package extended to permeable spheres. The hydrodynamic particle structure enters into the HYDROMULTIPOLE method only through a single-particle friction operator, whose form is known for a variety of particle models [54–56]. The details of the simulation method are given elsewhere [8]. The values for D_r presented in the following have been determined from equilibrium configuration averages over typically N = 256 particles in a periodically replicated cubic simulation box, using 100 initial random configurations for each set of parameters. This gives a statistical relative error of less than 0.001. In our multipole expansion method used for the rotational mobility tensor in Eq. (4), the multipole order, L, was truncated usually at L = 3. To gain high-precision data, extrapolations to $L \rightarrow 8$ were made, leading to an accuracy in D_r better than 1%. The calculated values for $D_r(N)$ using the periodic simulation box with N particles are not critically dependent on the system size, since $D_r(N = \infty) - D_r(N)$ scales with the particle number like 1/N. This system size dependence is similar to that of the high-frequency viscosity, $\eta_{\infty}(x,\phi)$, of permeable particles. The latter was calculated in earlier work [29, 36].

Table I lists our high-precision simulation results for $D_r(x, \phi)$, for volume fractions up to $\phi = 0.45$. Values of the inverse (reduced) penetration depth x are considered from a very small value $x \sim 5$, characteristic of highly permeable particles, up to $x = \infty$ characteristic of dry particles with stick surface boundary conditions.

TABLE I: Simulation results for the normalized short-time rotational self-diffusion coefficient $D_r(x,\phi)/D_0^r(x)$.

$\phi \setminus x$	5	10	20	30	50	100	∞
0.05	0.995	0.987	0.980	0.977	0.973	0.970	0.967
0.15	0.983	0.958	0.934	0.922	0.911	0.901	0.888
0.25	0.968	0.925	0.881	0.860	0.839	0.820	0.796
0.35	0.951	0.886	0.820	0.788	0.757	0.729	0.690
0.45	0.932	0.842	0.753	0.711	0.669	0.629	0.576

III. SCALING SELF-DIFFUSION OF PERMEABLE TO NON-PERMEABLE PARTICLES

From analyzing the numerical data for the rotational self-diffusion coefficient in Table I, we have found an interesting scaling of permeable to non-permeable spheres of the same size. In addition, we found that a similar scaling is valid for the translational self-diffusion coefficient. Therefore, results for both quantities will be given in this section. We start from a brief comparison of $D_r(x, \phi)$ to $D_t(x, \phi)$.

The simulation results for $D_r(x, \phi)$ from table I are depicted in the left panel of Fig. 1 using symbols. For comparison, the right panel of Fig. 1 shows the corresponding simulation results for $D_t(x, \phi)$ taken from [8]. For permeable particles, the fluid is allowed to penetrate so that the strength of the HIs is decreasing with increasing permeability, i.e., decreasing x. This is the reason for the larger values of D_r and D_t at larger permeabilities. Our results show that the effect of HIs on D_r is weaker than on D_t , i.e., for a given x and ϕ , the reduction of the self-diffusion coefficient relative the infinite dilution value is smaller for rotational diffusion.

The numerical results for $D_r(x, \phi)$ and $D_t(x, \phi)$ plotted in Fig. 1 have significantly different slopes at small volume fractions ϕ . On the other hand, these slopes are well-reproduced by the first-order virial coefficients, $\lambda_r(x)$ and $\lambda_t(x)$, defined by the following relations,

$$\frac{D_r(x,\phi)}{D_0^r(x)} = 1 + \lambda_r(x)\phi + \mathcal{O}(\phi^2)$$
(6)

$$\frac{D_t(x,\phi)}{D_0^r(x)} = 1 + \lambda_t(x)\phi + \mathcal{O}(\phi^2), \qquad (7)$$



FIG. 1: Rotational (left) and translational (right) self-diffusion coefficients, $D_r(x,\phi)/D_0^r(x)$ and $D_t(x,\phi)/D_0^t(x)$, as functions of ϕ , for values of x as indicated. Symbols: simulation results. Solid lines: interpolated r.h.s. of Eqs. (13-14).

evaluated in Ref. [37] and listed in Table II. The single-particle rotational diffusion coefficient, $D_0^r(x)$, has been already given in Eq. (5), and the translational one has the form given in [38, 39],

$$D_0^t(x) = \frac{k_B T}{6\pi\eta_0 a} \left(1 + \frac{1}{x \coth x - 1} + \frac{3}{2x^2} \right).$$
(8)

TABLE II: First-order virial terms, $\lambda_r(x)$ and $\lambda_t(x)$, of the rotational and translational self-diffusion coefficients [37].

	5	10	20	30	50	100	∞
λ_r	-0.097	-0.236	-0.376	-0.442	-0.505	-0.561	-0.631
λ_t	-0.569	-1.060	-1.416	-1.550	-1.661	-1.746	-1.832

Therefore, the idea is to introduce the following scaling functions,

$$u_r(x,\phi) = \left(\frac{D_r(x,\phi)}{D_0^r(x)} - 1\right) \frac{1}{\lambda_r(x)},\tag{9}$$

$$u_t(x,\phi) = \left(\frac{D_t(x,\phi)}{D_0^t(x)} - 1\right) \frac{1}{\lambda_t(x)}.$$
 (10)

For all values of $x \ge 5$, the functions $u_r(x, \phi)$ and $u_t(x, \phi)$ do practically not depend on x, i.e. they are permeability-independent. Indeed, as shown in Fig. 2, the curves for u_r and u_t as functions of ϕ collapse on the corresponding curves for the non-permeable solid spheres, i.e.

$$u_r(x,\phi) \approx u_r(\infty,\phi),$$
 (11)

$$u_t(x,\phi) \approx u_r(\infty,\phi),$$
 (12)

with a relative error less than 3% for $x \ge 10$.



FIG. 2: The functions $u_r(x, \phi)$ and $u_t(x, \phi)$ are practically independent of x. Symbols: simulation results for the indicated values of x. Solid lines: spline fit interpolations of $u_r(\infty, \phi)$ and $u_t(\infty, x)$.

Therefore, the short-time self-diffusion coefficients in suspensions of permeable particles are well approximated by the following expressions,

$$\frac{D_r(x,\phi)}{D_0^r(x)} \approx 1 + \lambda_r(x)u_r(\infty,\phi), \qquad (13)$$

$$\frac{D_t(x,\phi)}{D_0^t(x)} \approx 1 + \lambda_t(x)u_t(\infty,\phi).$$
(14)

In Fig. 1, the solid, continuous lines are not just mere fits to the simulation data, but represent the expressions in Eqs. (13-14), i.e. the outcome of the interesting scaling behavior of the short-time self-diffusion of permeable particles. The error made in using Eqs. (13-14) instead of the precise simulation values, is at most 1% for rotational and 3% for translational self-diffusion. To complete the analysis, we need to specify the scaling functions for the non-permeable solid spheres. This will be done in the next section.

IV. SELF-DIFFUSION COEFFICIENTS OF NON-PERMEABLE SPHERES

We will now use the existing data for non-permeable hard spheres to construct simple approximate expressions for the scaling functions $u_r(\infty, \phi)$ and $u_t(\infty, \phi)$.

We start with a comparison between our present simulation results for $D_r^{\rm hs}(\phi) = D_r(\infty, \phi)$ and $D_t^{\rm hs}(\phi) = D_t(\infty, \phi)$ for vanishing permeability, and a selection out of a large body of published experimental (see, e.g., [4, 6, 15, 16, 18]) and simulation (see, e.g., [7, 23, 24, 29, 58– 61]) data on impermeable hard spheres.

Related to this comparison, we note first that numerical results of varying precision have been published for the first and second virial coefficients of the rotational [15, 25–27] and translational [26, 27, 62] self-diffusion coefficients. High-precision second-order virial expansion results,

$$\frac{D_r^{\rm hs}}{D_0^{r,\rm hs}} = 1 - 0.631\phi - 0.726\phi^2 + \mathcal{O}(\phi^3), \qquad (15)$$

$$\frac{D_t^{\rm hs}}{D_0^{t,\rm hs}} = 1 - 1.8315\phi - 0.219\phi^2 + \mathcal{O}(\phi^3), \qquad (16)$$

have been obtained by Cichocki et al. [28], using a lubrication correction for the three-body HIs contributions.

Regarding rotational diffusion, Fig. 3(a) shows the comparison of our data with Lattice-Boltzmann [23] and ASD [7] simulation results, and DDLS experimental data [15] for optically anisotropic fluorinated polymer particles. The rotational diffusion coefficient as a function of ϕ has a concave shape, different from that for D_t which is weakly convex. Our simulation data for non-permeable particles agree well with the ASD result. The LB data at large ϕ are somewhat smaller. The key message conveyed by Fig. 3(a) is that the secondorder virial result for $D_r^{\rm hs}(\phi)$ in Eq. (15) describes the simulation and experimental data remarkably well for all volume fractions up to the freezing transition value 0.49 [7, 23], indicating that higher-order virial coefficients are small or mutually cancel out. Therefore, for constructing a simple approximation for $u_r(\infty, \phi)$ from Eq. (9), it is sufficient to take as $D_r(\infty, x)$ the 2nd-order virial expansion in Eq. (15). In this way, the rotational scaling function is approximated by

$$u_r(\infty,\phi) \approx \phi + 1.151\phi^2, \qquad (17)$$

with an accuracy of 1.5% or better relative to our simulation data.



FIG. 3: (a) Rotational and (b) translational self-diffusion coefficients of non-permeable hard spheres with stick boundary conditions, as functions of ϕ . Compared in (a) are our HYDROMULTIPOLE data (labeled HYDRO) for $D_r^{\rm hs}/D_0^{\rm hs}$ with SD [7] and LB [23] simulation results, and DDLS experimental data [15] for optically anisotropic particles. In (b), we compare the HYDROMULTIPOLE data for $D_t^{\rm hs}(\phi)/D_0^{\rm t,hs}$ with ASD simulation results [7], force multipole calculations by Ladd [58], and DLS experimental data by Segre et al. [6]. Solid lines: 2nd-order virial expansion results, in (a) according to Eq. (15), and in (b) according to Eq. (16).

In Fig. 3(b), we compare our simulation data for $D_t^{\rm hs}(\phi)/D_0^{t,\rm hs}$ with ASD simulation [7] and force multipole calculation results [58], and with DLS experimental data [6]. The figure shows that the translational second-order virial expression in Eq. (16) for $D_t^{\rm hs}$ noticeably underestimates the simulation and experimental data when ϕ is larger than 0.3.

For this reason, we need a more precise expression for $D_t(\infty, \phi)$ than the 2nd-order virial expansion in Eq. (16). We have found that our simulation data are approximated with a 0.4% accuracy by the following expression for the scaling function $u_t(\infty, \phi)$, defined in Eq. (10),

$$u_t(\infty, \phi) \approx \phi + 0.12 \phi^2 - 0.65 \phi^3.$$
 (18)

The term $\phi + 0.12\phi^2$ follows from the virial expansion in Eq. (16), and the coefficient of the third order term, $-0.65 \phi^3$, has been obtained by fitting to the numerical data for $u_t(\infty, \phi) - \phi - 0.12\phi^2$, in the range $0 \le \phi \le 0.45$.

V. GENERALIZED STOKES-EINSTEIN-DEBYE RELATION

We proceed with the discussion of a generalized short-time GSED relation. Having obtained in this paper precise numerical data for $D_r(x, \phi)$, and taking values of $\eta_{\infty}(x, \phi)$ tabulated in [36], we are in the position to test the validity of the following short-time GSED relation

$$\frac{D_r(x,\phi)}{D_0^r(x)} \frac{\eta_\infty(x,\phi)}{\eta_0} \stackrel{?}{\approx} 1, \qquad (19)$$

between the rotational self-diffusion coefficient, and the high-frequency viscosity of permeable particles. The validity of generalized Stokes-Einstein relations such as the present one is an important issue in microrheological studies where one tries to infer rheological properties more easily from diffusion measurements. The GSED relation in Eq. (19) was shown before to be violated for suspensions of non-permeable neutral and charged particles [18, 63]. Here, we ask the same validity question for permeable particle systems.

In Fig. 4, the GSED relation is examined for different values of x. If valid, all curves should collapse on a single horizontal line of unit height. One notices from the figure that the GSED relation is significantly violated for $x \ge 30$, and volume fractions $\phi > 0.15$ where the particles are significantly correlated. Thus, a rotating particle experiences its neighborhood not just as a structureless medium characterized by the viscosity $\eta_{\infty}(x, \phi)$.



FIG. 4: The generalized Stokes-Einstein-Debye relation between $D_r(x, \phi)$ and high-frequency viscosity $\eta_{\infty}(x, \phi)$ is not satisfied. Solid lines are interpolating spline fits to our simulation results (symbols).

The GSED relation for rotational diffusion is more strongly violated than its translational counterpart. As shown in [35], $(D_t/D_0^t) \times (\eta_{\infty}/\eta_0)$ increases practically linearly in ϕ , even for non-permeable particles, whereas in Fig. 4 a pronounced non-linear increase is observed.

VI. CONCLUSIONS

Using the HYDROMULTIPOLE simulation method, the short-time rotational self-diffusion coefficient, $D_r(x, \phi)$, of uniformly permeable spheres was calculated to high precision as a function of permeability and volume fraction.

An interesting scaling relation was found between $D_r(x, \phi)$ and the corresponding coefficient, $D_r(\infty, \phi)$, of non-permeable, solid spheres of the same size, where the permeability enters only through the first-order rotational virial coefficient. A similar scaling was found for translational self-diffusion.

The combination of the scaling relations with accurate 2nd-order and 3rd-order concentration expansion results in Eqs. (17-18) for $D_r(\infty, \phi)$ and $D_t(\infty, \phi)$, respectively, has led us to the expressions

$$\frac{D_r(x,\phi)}{D_0^r(x)} \approx 1 + \lambda_r(x)\phi \ (1+1.151\phi) \ , \tag{20}$$

$$\frac{D_t(x,\phi)}{D_0^t(x)} \approx 1 + \lambda_t(x)\phi \left(1 + 0.12\phi - 0.65\phi^2\right), \qquad (21)$$

for the permeability-dependent self-diffusion coefficients. In combination with table II for λ_r and λ_t , these are convenient expressions useful in diffusion measurement analysis of permeable particle systems, and as input to theories of long-time dynamic properties. The accuracy of these expressions is better than 1.5% for rotational and 3.5% for translational self-diffusion, for the whole range of volume fractions $\phi \leq 0.45$ provided $x \geq 5$. We expect the expressions to be useful in the experimental analysis of self-diffusion, to gain a quick estimate of the mean porosity in concentrated systems. Moreover, they can serve as short-time inputs into theoretical methods of calculating frequency-dependent and long-time diffusion properties, such as in mode-coupling and dynamic density functional theory methods.

The simulation results for $D_r(x, \phi)$, and recent results for $\eta_{\infty}(x, \phi)$, were used to scrutinize the validity of a generalized Stokes-Einstein-Debye relation in its dependence on permeability. We found this relation to be significantly violated for non-dilute suspensions, unless the permeability is unrealistically large. The GSED test for porous particles presented in this paper complements earlier GSE performance tests [35] of different short-time diffusion properties. Of all considered GSE relations, only the one for the cage diffusion coefficient can claim a certain validity when applied to neutral porous particles [35]. However, also this relation becomes invalid when the particles are significantly charged [7].

With the present paper on self-diffusion in combination with earlier simulation results for other dynamic properties such as the hydrodynamic function and viscosity, and the development of accurate analytic approximations for these properties [8, 35–37], we have obtained an essentially complete description of the short-time dynamics of uniformly permeable particles with no-overlap interactions.

Dispersions of spherical particles with more complex internal hydrodynamic structure, such as core-shell particles, and different direct interactions, will be the subject of a future study.

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- [3] T. Ando and J. Skolnick, PNAS **107**, 18457 (2010).
- [4] W. van Megen, S.M. Underwood, and I. Snook, J. Chem. Phys. 85, 4065 (1999).
- [5] A. Brand, H. Versmold, and W. van Megen, J. Chem. Phys. 110, 1283 (1999).

P.N. Pusey, in *Liquids, Freezing, and the Glass Transition*, edited by J. P. Hansen, D. Levesque, and J. Zinn-Justin, Elsevier, Amsterdam, (1991).

^[2] G. Nägele, J.K.G. Dhont, and G. Meier in *Diffusion in Condensed Matter*, edited by P. Heitjans and J. Kärger, Springer, Berlin (2005).

- [6] P. N. Segrè, O. P. Behrend and P. N. Pusey, Phys. Rev. E 52, 5070 (1995).
- [7] A.J. Banchio and G. Nägele, J. Chem. Phys. **128**, 104903 (2008).
- [8] G.C. Abade, B. Cichocki, M.L. Ekiel-Jeżewska, G. Nägele, and E. Wajnryb, J. Chem. Phys. 132, 014503 (2010).
- [9] P.N. Pusey, J. Phys. A: Math. Gen. 11, 119 (1978).
- [10] P. Holmqvist and G. Nägele, Phys. Rev. Lett. 104, 058301 (2010).
- [11] M. Heinen, P. Holmqvist, A.J. Banchio and G. Nägele, J. Appl. Cryst. 43, 970 (2010).
- [12] W.D. Dozier and P.M. Chaikin, J. Phys. (Paris) 46-C.3, 165 (1985).
- [13] R. Simon, T. Palberg, and P. Leiderer, J. Chem. Phys. **99**, 3030 (1993).
- [14] A. Wilk, J. Gapinski, A. Patkowski, and R. Pecora, J. Chem. Phys. 121, 10794 (2004).
- [15] V. Degiorgio, R. Piazza, and R.B. Jones, Phys. Rev. E 52, 2707 (1995).
- [16] J. Kannetakis, A. Tölle, and H. Sillescu, Phys. Rev. E 55, 3006 (1997).
- [17] G.H. Koenderink, H. Zhang, M.P. Lettinga, G. Nägele and A.P. Philipse, Phys. Rev. E 64, 2001 (2001).
- [18] G.H. Koenderink, H. Zhang, D.G.A.L. Aarts, M.P. Lettinga, A.P. Philipse, and G. Nägele, Faraday Discuss. 123, 335 (2003).
- [19] M.P. Lettinga, G.H. Koenderink, B.W.M. Kuipers, E.Bessels, and A.P. Philipse, J. Chem. Phys. 120, 4517 (2003).
- [20] G.H. Koenderink, Ph.D. thesis: Rotational and translational diffusion in colloidal mixtures, Utrecht University, The Netherlands (2003).
- [21] H. Zhang and G. Nägele, J. Chem. Phys. 117, 5908 (2002).
- [22] E. Andablo-Reyes, P. Diaz-Leyva, and J.L. Arauz-Lara, Phys. Rev. Lett. 94, 106001 (2005).
- [23] M.H.J. Hagen, D. Frenkel and C.P. Lowe, Physica A 272, 376 (1999).
- [24] R.J. Phillips, J.F. Brady, and G. Bossis, Phys. Fluids **31**, 3462 (1988).
- [25] C. Urdaneta, D.M. Jones and M. Muthukumar, J. Chem. Phys. 91, 5127 (1989).
- [26] H.J.H. Clercx and P.P.J.M. Schram, Physica A 174, 293 (1991).
- [27] H.J.H. Clercx and P.P.J.M. Schram, J. Chem. Phys. 96, 3137 (1992).
- [28] B. Cichocki, M.L. Ekiel-Jeżewska and E. Wajnryb, J. Chem. Phys. 111, 3265 (1999).
- [29] G. Mo and A. S. Sangani, Phys. Fluids 6, 1637 (1994).
- [30] A.A. Potanin and W.B. Russel, Phys. Rev. E 52, 730 (1995).
- [31] S.L. Elliot and W.B. Russel, J. Rheol. 42, 361 (1998).

- [32] P. A. Nommensen, M.H.G. Duits, D. van den Ende, and J. Mellema, Phys. Rev. E 59, 3147 (1999).
- [33] S.B. Chen and A. Cai, J. Colloid Interface Sci. 217, 328 (1999).
- [34] G.C. Abade, B. Cichocki, M.L. Ekiel-Jeżewska, G. Nägele, and E. Wajnryb, Phys. Rev. E. 81, 020404(R) (2010).
- [35] G.C. Abade, B. Cichocki, M.L. Ekiel-Jeżewska, G. Nägele, and E. Wajnryb, J. Phys.: Condens. Matter 22, 322101 (2010).
- [36] G.C. Abade, B. Cichocki, M.L. Ekiel-Jeżewska, G. Nägele, and E. Wajnryb, J. Chem. Phys. 133, 084906 (2010).
- [37] B. Cichocki, M.L. Ekiel-Jeżewska, G. Nägele and E. Wajnryb, submitted to Phys. Fluids (2011).
- [38] H.C. Brinkman, Proc. R. Dutch Acad. Sci. 50, 618, 821 (1947).
- [39] P. Debye and A.M. Bueche, J. Chem. Phys. 16, 573 (1948).
- [40] M.H.G. Duits, P. A. Nommensen, D. van den Ende, and J. Mellema, Colloids Surf. A 183-185, 335 (2001).
- [41] C. N. Likos, S. Rosenfeldt, N. Dingenouts, M. Ballauff, P. Lindner, N. Werner, and F. Vögtle, J. Chem. Phys. 117, 1869 (2002).
- [42] S. Huißmann, A. Wynveen, C.N. Likos, and R. Blaak, J. Phys.: Condens. Matter 22, 232101 (2010).
- [43] Z. Adamczyk, B. Jachimska, and M. Kolasinska, J. Colloid Interface Sci. 273, 668 (2004).
- [44] S. Pyett and W. Richtering, J. Chem. Phys. 122, 034709 (2005).
- [45] T. Eckert and W. Richtering, J. Chem. Phys. 129, 124902 (2008).
- [46] C.A. Coutinho, R.K. Harrinauth, and V.K. Gupta, Colloids Surf. A 318, 111 (2008).
- [47] G. Petekidis, J. Gapiński, P. Seymour, J. S. van Duijneveldt, D. Vlassopoulos and G. Fytas, Phys. Rev. E 69, 042401 (2004).
- [48] M. Zackrisson, A. Stradner, P. Schurtenberger and J. Bergenholtz, Langmuir 21, 10835 (2005).
- [49] J. Masliyah, G. Neale, K. Malysa, and T.G.M. van de Ven, Chem. Eng. Science 42, 245 (1987).
- [50] C. N. Likos, H. Löwen, M. Watzlawek, B. Abbas, O. Jucknischke, J. Allgaier, and D. Richter, Phys. Rev. Lett. 80, 4450 (1998).
- [51] B. Cichocki, B.U. Felderhof, K. Hinsen, E. Wajnryb, and J. Bławzdziewicz, J. Chem. Phys. 100, 3780 (1994).

- [52] J. Happel and H. Brenner, Low Reynolds Number Hydrodynamics, Martinus Nijhoff, Dordrecht (1986).
- [53] S. Kim and S.J. Karrila, Microhydrodynamics: Principles and Selected Applications, Butterworth-Heinemann, London (1991).
- [54] B.U. Felderhof and J.M. Deutch, J. Chem. Phys. 62, 2391 (1975).
- [55] R. Reuland, B.U. Felderhof and R.B. Jones, Physica A 93, 465 (1978).
- [56] B. Cichocki, B. U. Felderhof and R. Schmitz, PhysicoChemical Hydrodynamics 10, 383 (1988).
- [57] B. Cichocki, R.B. Jones, R. Kutteh and E. Wajnryb, J. Chem. Phys. 112, 2548 (2000).
- [58] A.J.C. Ladd, J. Chem. Phys. **93**, 3484 (1990).
- [59] A. Sierou and J.F. Brady, J. Fluid Mech. 448, 115 (2001).
- [60] A.J. Banchio and J.F. Brady, J. Chem. Phys. 118, 10323 (2003).
- [61] J.F. Brady, Curr. Opin. Colloid Interface Sci. 1, 472 (1996).
- [62] D.M. Jones, M. Muthukumar and S.M. Cohen, J. Chem. Phys. 90, 7542 (1989).
- [63] A.J. Banchio, G. Nägele and J. Bergenholtz, J. Chem. Phys. 111, 8721 (1999).