A fundamental assumption of the dynamical density functional theory (DDFT) of colloidal systems is that a grand-canonical free-energy functional may be employed to generate the thermodynamic driving forces. Using one-dimensional hard rods as a model system, we analyze the validity of this key assumption and show that unphysical self-interactions of the tagged particle density fields, arising from coupling to a particle reservoir, are responsible for the excessively fast relaxation predicted by the theory. Moreover, our findings suggest that even employing a canonical functional would not lead to an improvement for many-particle systems, if only the total density is considered. We present several possible schemes to suppress these effects by incorporating tagged densities. When applied to confined systems, we demonstrate, using a simple example, that DDFT necessarily leads to delocalized tagged particle density distributions, which do not respect the fundamental geometrical constraints apparent in Brownian dynamics simulation data. The implication of these results for possible applications of DDFT to treat the glass transition are discussed.

1. INTRODUCTION

During the past decade of liquid-state research, the classical dynamical density functional theory (DDFT) has proven to be a versatile and reliable tool for describing the dynamics of interacting colloidal particles in a wide variety of situations. Building upon the success of equilibrium DFT [1,2], the dynamical theory enables first-principles calculation of the inhomogeneous density $\rho(r,t)$ generated in response to a time-dependent external potential field $V_{\text{ext}}(r,t)$ [3]. Within the DDFT framework, a nonvanishing particle flux arises solely from gradients in a local chemical potential $\mu(r,t)$, derived from an equilibrium free-energy functional. The theory has been successfully applied to many problems, including relaxation to equilibrium from a given nonequilibrium initial state [3], the early and intermediate stages of spinodal decomposition [4], and systems for which the time-dependence of $V_{\text{ext}}(r,t)$ drives the system into nonequilibrium steady or stationary states [5–9]. In recent years, the original formulation of DDFT has been extended to treat both systems and situations of ever increasing complexity. These more recent developments have lead to a better understanding of the influence of both nonpotential fields (e.g., mechanical [10–13], thermal [7,14]) and particle geometry (see, e.g., Ref. [15]) on diffusive colloidal dynamics.

Despite the undeniable success of the DDFT in robustly capturing the qualitative features of $\rho(r,t)$ for many problems of interest, the theory rests upon two fundamental assumptions, both of which remain to be either systematically evaluated or improved upon. The first of these is the so-called adiabatic approximation, namely the assumption that the two-body correlations may be calculated from the instantaneous one-body density using equilibrium statistical mechanical relations. The second major assumption is that the nonequilibrium chemical potential $\mu(r,t)$ generating the particle dynamics can be identified with the functional derivative of a grand-canonical free energy. Combining these two approximations yields a closed equation for the one-body density, which does not require explicit knowledge of the higher-order correlations.

In the present work we investigate the validity of employing a grand-canonical density functional to treat many-body effects in DDFT. Problems can be anticipated in confined systems with small particle number, for which the choice of ensemble strongly influences the equilibrium density profiles. More generally, use of the grand-canonical ensemble becomes questionable for situations where the density field is strongly localized in space and contains only few particles. This can occur either as a direct consequence of minima in $V_{\text{ext}}(r,t)$ or as a transient, which may occur along the natural diffusive trajectory of $\rho(r,t)$ through the space of density functions. Transient localization occurs quite naturally when considering the time-evolution of the density from sharply defined initial conditions, for which the positions of all $N$ particles, or a subset thereof, are known precisely. The individual density peaks of particles sharply located at $t = 0$ remain well separated for short times and are normalized to unity (each peak contains a single particle).

The above considerations become even more pertinent when considering potential applications of grand-canonical DDFT to describe dynamic arrest and glass formation. By tagging the density field of a single particle in a dense hard-sphere liquid, it has recently been shown [16] that the tagged density (the self part of the van Hove function) can exhibit a two-step relaxation within DDFT, leading to dynamic arrest at sufficiently high-volume fractions. Similar behavior was also found for particles interacting via a Gaussian pair potential [17]. However, due to the use of an approximate free-energy functional, it remains unclear whether the observed slow dynamics arise purely from the presence of (possibly spurious) metastable minima in the free energy, or whether it is a genuine physical prediction of the DDFT which would persist even when using an exact equilibrium functional. The work of Hopkins et al. [16,17] raises a fundamental question: Can a theory that involves only the one-body density field capture, even in principle, the transition to a nonergodic state?

A major difficulty in answering the above question for realistic glass forming systems in two or three dimensions is the absence of an exact equilibrium free-energy functional.
It thus becomes difficult to disentangle errors in the dynamical theory from those induced by an approximate free energy. For this reason we focus on the simplest nontrivial model for which the grand-canonical free-energy functional is known exactly, namely a system of hard rods in one dimension \[18,19\]. Despite its simplicity, the hard-rod model has one feature in common with glassy states occurring in higher dimensional systems: both exhibit a partitioned phase space in which physically meaningful averages must be taken over a subset of phase space that is dynamically accessible. In glasses, this partitioning occurs spontaneously as a thermodynamic control parameter crosses some critical value, whereas for hard rods the reduced phase space is always present, simply as a result of ordering the particles on a line. The confinement of a given rod by its neighbors to the left and right presents the prototypical glassy “cage” and serves as a useful reality check for theories aiming to describe dynamical arrest in higher dimensions.

The paper will be structured as follows: In Sec. II we first outline the microscopic dynamics under consideration, review the standard formulation of DDFT, and define the functional to be employed. In Sec. III we consider the diffusion of a single rod and various methods by which grand-canonical contributions can be eliminated. In Sec. IV we identify the importance of tagging the individual density profiles. In Sec. V we present numerical results for various test cases involving several interacting rods. Finally, in Sec. VI we provide a discussion of our results and identify some future challenges.

II. FUNDAMENTALS

A. Microscopic dynamics

We consider a system of \( N \) colloidal particles in a time-dependent external potential \( V_{\text{ext}}(r,t) \) interacting via the spherically symmetric pair potential \( \phi(r) \). The total potential energy is given by

\[
U_N(r_i,t) = \sum_i V_{\text{ext}}(r_i,t) + \sum_{i<j} \phi(r_i - r_j). \tag{1}
\]

As the individual colloidal trajectories are stochastic, it is appropriate to adopt a probabilistic description of the particle motion. The probability distribution of particle positions is described by the Smoluchowski equation \[20\]:

\[
\frac{\partial P(t)}{\partial t} + \sum_i \nabla_i \cdot \mathbf{j}_i = 0, \tag{2}
\]

where \( P(t) \equiv P(\{r_i\},t) \). The fact that Eq. (2) takes the form of a continuity equation expresses the conservation of particle number. Neglecting the influence of solvent-induced hydrodynamic interactions (see Ref. [21] for a discussion of the implications of this approximation) the probability flux of particle \( i \) is given by

\[
\mathbf{j}_i = -D_0(\nabla_i - \beta \mathbf{F}_i)P(t), \tag{3}
\]

with \( \beta = 1/k_BT \). The force \( \mathbf{F}_i \) on particle \( i \) is generated from the total potential energy [Eq. (1)] according to \( \mathbf{F}_i = -\nabla_i U_N \).

B. Dynamic density functional theory

Integration of Eq. (2) over all but one of the particle coordinates leads to an exact coarse-grained expression for the one-body density profile

\[
\frac{\partial \rho(r_1,t)}{\partial t} = -\nabla_1 \cdot \mathbf{j}(r_1,t), \tag{4}
\]

where the particle flux involves an integral over the nonequilibrium two-body density

\[
\mathbf{j}(r_1,t) = \Gamma k_B T \nabla_1 \rho(r_1,t) + \Gamma \rho(r_1,t) \nabla_1 V_{\text{ext}}(r_1,t) + \Gamma \int dr_2 \nabla_1 \phi(r_{12}) \rho^{(2)}(r_1,r_2,t), \tag{5}
\]

with mobility \( \Gamma = D_0/k_BT \). Equation (4) is the first in a hierarchy of equations for the \( n \)-body density. In equilibrium, the flux is identically zero and Eq. (4) reduces to the first member of the Yvon-Born-Green hierarchy [22].

The integral term in Eq. (5) may be approximated as an explicit functional of the one-body density using the methods of equilibrium DFT. The Helmholtz free energy is split into three contributions,

\[
\mathcal{F} = \mathcal{F}_{\text{id}} + \mathcal{F}_{\text{ex}} + \int dr_1 V_{\text{ext}}(r_1) \rho(r_1), \tag{6}
\]

where \( \Lambda \) is the thermal de Broglie wavelength and the ideal gas free energy is known exactly:

\[
\mathcal{F}_{\text{id}}[\rho(r_1)] = \int dr_1 \rho(r_1) [\ln(\Lambda^3 \rho(r_1)) - 1]. \tag{7}
\]

The excess free-energy functional \( \mathcal{F}_{\text{ex}}[\rho(r_1)] \) contains all information regarding the interparticle interactions and is connected to the average interaction force via the grand-canonical sum rule \[1\],

\[
\int dr_2 \nabla_1 \phi(r_{12}) \rho(r_2|r_1) = \nabla_1 \frac{\delta \mathcal{F}_{\text{ex}}[\rho(r_1)]}{\delta \rho(r_1)}, \tag{8}
\]

where we have introduced the conditional distribution \( \rho(r_2|r_1) \equiv \rho^{(2)}(r_1,r_2)/\rho(r_1) \); i.e., the average number density at \( r_2 \) given a particle is fixed at \( r_1 \).

The assumption that Eq. (8) remains valid in nonequilibrium is the so-called adiabatic approximation. It is equivalent to assuming that the pair density \( \rho^{(2)}(r_1,r_2,t) \) relaxes instantaneously to the equilibrium pair-density corresponding to the current one-body density \( \rho(r_1,t) \). The total particle flux [Eq. (5)] may thus be written in the form

\[
\mathbf{j}(r_1,t) = -\Gamma \rho(r_1,t) \nabla_1 \mu(r_1,t), \tag{9}
\]

where the nonequilibrium chemical potential is given by

\[
\mu(r_1,t) = \frac{\delta \mathcal{F}[\rho(r_1,t)]}{\delta \rho(r_1,t)}. \tag{10}
\]

Combining Eqs. (4), (9), and (10) yields the familiar form of the DDFT equation of motion:

\[
\frac{\partial \rho(r_1,t)}{\partial t} = \nabla_1 \left\{ \Gamma \rho(r_1,t) \nabla_1 \frac{\delta \mathcal{F}[\rho(r_1,t)]}{\delta \rho(r_1,t)} \right\}. \tag{11}
\]

In standard applications of DDFT the free-energy \( \mathcal{F} \) entering Eq. (11) is a grand-canonical quantity. Equation (11) thus
describes the evolution of the density between two grand-canonical states, subject to the constraint that the average particle number is conserved.

For the present work, the DDFT equation is solved numerically using finite difference techniques. The time integration is performed using the Euler method and the spatial derivatives with central differences. The integrations required for the convolutions are evaluated using the trapezoidal rule, which in one dimension can be calculated very efficiently due to the finite range of the weight functions.

C. The Percus hard-rod functional

For our numerical investigations we will employ the exact excess Helmholtz free-energy functional of hard rods in one dimension [18,19]. For an $m$ component mixture of rods the functional is given by

$$F_{\text{ex}}[\rho(x)] = \int dx \Phi(n_x),$$

(12)

where the free-energy density, $\Phi = -n_0(x) \ln[1 - n_1(x)]$, is a function of a set of weighted densities,

$$n_x = \sum_{i=1}^m \int dx' \rho_i(x') \omega_i(x - x'),$$

(13)

where $\rho_i(x)$ is the density profile of species $i$. The weight functions are characteristic of the particle geometry, with $\omega_i^0(x) = \Theta(|x| - d_i/2)$ and $\omega_i^0(x) = \delta(|x| - d_i/2)/2$, where $d_i$ is the length of rod species $i$.

III. SINGLE-PARTICLE DIFFUSION

We begin by considering a single rod, whose diffusion is governed by the diffusion equation, resulting in a Gaussian density distribution. The diffusion equation can be recovered from the DDFT equation of motion [Eq. (11)] in the case of vanishing interactions; i.e., $F_{\text{ex}} = 0$. In Fig. 1 we compare the results of DDFT Eq. (11) using the Percus functional with the exact Gaussian result for the relaxation of the density profile from the sharp initial condition $\rho(x) = \delta(x)$. The inset to Fig. 1 shows the corresponding mean-squared displacement (MSD) as a function of time. The DDFT reproduces not only the expected Gaussian form of the density profile nor the linear increase of the MSD as a function of time. Only at long times does the slope of the MSD approach unity as the local density becomes very small for all $x$ and the ideal gas term starts to dominate the free energy [Eq. (6)]. The deficiency of the theory lies in the nonvanishing contribution from the excess free energy, which leads to an effective self interaction of the density field and consequent enhanced relaxation rate.

Original points out by Marconi and Tarazona [3], employing a grand canonical functional effectively puts the system into contact with an unphysical particle reservoir, such that even for $\langle N \rangle = 1$ the density distribution contains additional contributions from states with $N = 0, 2, 3, \ldots$. The interaction term thus does not vanish, as it should for a single particle, because states with $N > 1$ naturally incorporate interparticle interactions. The first step toward an improved theory is thus to eliminate, or at least reduce, the interaction between the physical particle and the reservoir.

A. Canonical correction

In Refs. [23] and [24], González et al. proposed a scheme by which canonical equilibrium density profiles can be expanded in terms of grand-canonical averages. The method can be interpreted as a formal expansion of the canonical density profile in inverse powers of $\langle N \rangle$. Using this expansion, it was found possible to systematically correct the grand-canonical density profiles predicted by equilibrium DFT to achieve an improved description of a hard-sphere fluid confined in a small spherical cavity. Generalizing the arguments presented in Refs. [23,24], the canonical average of an arbitrary function of the particle positions $A(r_i)$ may be expressed in terms of grand-canonical averages. To second order the expansion is given by

$$\langle A \rangle^c = \langle A \rangle + f_1(A) + f_2(A) + \cdots,$$

(14)

where $\langle \cdot \rangle$ is a grand-canonical average and $\langle \cdot \rangle^c$ is a canonical average. The correction terms are given by

$$f_1(A) = -\frac{1}{2} \langle N - \langle N \rangle \rangle^2 \frac{\partial^2 A}{\partial \langle N \rangle^2},$$

$$f_2(A) = -\frac{1}{6} \langle N - \langle N \rangle \rangle^3 \frac{\partial^3 A}{\partial \langle N \rangle^3},$$

$$f_2(A) = -\frac{1}{2} \langle N - \langle N \rangle \rangle^2 \frac{\partial^2 f_1}{\partial \langle N \rangle^2}.$$

Due to the scaling of the partial derivatives in Eq. (14) the terms $f_1$ and $f_2$ are of order $\langle N \rangle^{-1}$ and $\langle N \rangle^{-2}$, respectively. The utility of the expansion Eq. (14) lies in its rapid convergence, even for very small values of $\langle N \rangle$ [23,24].
Returning to the diffusion of a single particle, we now seek to use Eq. (14) to suppress unwanted grand-canonical contributions to the dynamics of $\rho(r,t)$. At any given time we can use the instantaneous density profile $\rho(r,t)$ to construct an effective external potential,

$$V_{\text{eff}}(r,t) = -\beta^{-1}\ln[\rho(r,t)] - c^{(1)}[\rho(r,t)] + \ln(z),$$

where the fugacity term $\ln(z)$ is a physically irrelevant constant. The one-body direct correlation function,

$$c^{(1)}(r_1) = -\frac{\beta V_{\text{ex}}[\rho]}{\delta \rho(r_1)},$$

is evaluated using the instantaneous density. When employed in an equilibrium calculation, the potential $V_{\text{eff}}(r,t)$ will, by construction, yield the equilibrium results for $\rho(r,t)$ and all other quantities accessible to DFT. The adiabatic instantaneous density profile

$$\rho_{\text{in}}(r,t) = \int d^3r_2 \nabla_1 \phi(r_2) \rho_{\text{gc}}^{(2)}(r_1,r_2),$$

where the subscript (gc) makes explicit the fact that the pair-density inside the integral is a grand-canonical average. Using Eq. (8) yields

$$I(r_1,t) = -k_B T \rho(r_1,t) \nabla_1 c^{(1)}(r_1,t).$$

Expressed in this way, it is rather natural to employ the expansion Eq. (14) to approximate the required integral in terms of known grand-canonical quantities:

$$\int d^3r_2 \nabla_1 \phi(r_2) \rho_{\text{gc}}^{(2)}(r_1,r_2) = I(r_1,t) + f_1[I(r_1,t)] + f_2[I(r_1,t)] + \cdots$$

When calculated to all orders, the corrected interaction force should be zero for a single particle. We now proceed to explicitly calculate the first two correction terms in Eq. (19). The practical implementation of our scheme is as follows: at each time step in the numerical integration of Eq. (2), the potential terms appearing in Eq. (19) are then calculated numerically by performing equilibrium DFT calculations in the presence of a fixed $V_{\text{eff}}(r_1,t)$. The series Eq. (19) is then evaluated to the desired order and used to generate the density distribution at the next time step. Grand-canonical contributions to the time evolution of the density $\rho(r,t)$ can thus be suppressed on the fly in order to provide a more realistic description of the trajectory through the space of density functions.

In Fig. 1 we show the time evolution of the density of a single particle corrected using Eq. (19) to both first and second order. The series converges very rapidly and the second-order results are virtually indistinguishable from the exact Gaussian function, despite the fact that $\langle N \rangle = 1$. Similar rapid convergence has been observed in the static case [24].

### IV. MULTIPLE-PARTICLE DIFFUSION

We have now established that a systematic suppression of the grand-canonical contributions to the dynamics indeed leads to improved results, at least for the trivial case of a single particle. However, application of the same procedure to systems with $N > 1$ reveals an additional complication.

Consider first the equilibrium average for a single infinite potential well in the canonical ensemble with $N = 1$ and in the grand-canonical ensemble with $\langle N \rangle = 1$ [see Figs. 2(a) and 2(b)]. The average value of a quantity is the average over the value of this quantity for all microstates appropriate to the given ensemble. As a result, the average density profile for canonical and grand-canonical cases are different because of the additional microstates arising from coupling to an external particle reservoir.

The situation is similar to DDFT applied to transiently localized particles that have not had enough time to diffuse far away from their starting position. The average interaction force is a grand canonical average, and the potential wells discussed above are manifest in the effective external potential [Eq. (15)]. In the grand-canonical case the microstates with more than one particle in the well give rise to a nonvanishing interaction, which leads to the erroneous average interaction force, which does not vanish even if the two density peaks are far away from each other. The interaction force on a particle resulting from number

![FIG. 2. (Color online) Schematic illustration of the microstates that contribute to canonical and grand-canonical averages for particles in one and two infinite potential wells. (a) Canonical average $N = 1$; (b) grand-canonical average $\langle N \rangle = 1$; (c) canonical average $N = 2$; (d) grand-canonical average $\langle N \rangle = 2$. The second and third microstate in (c) are unphysical in the context of DDFT for localized particles and lead to a overestimation of the interactions.](http://doc.rero.ch)
fluctuations in the $V_{ij}(r,t)$ well generated by the particle can therefore be interpreted as a self interaction.

For larger numbers of particles and wells, each well is coupled to a reservoir formed by the other wells. As $N$ increases, the canonical and grand-canonical average become increasingly similar and the canonical correction decreases in magnitude, rendering it useless for the suppression of the self interaction.

What is required is a method to prevent the averaging procedure from including microstates states with more than one particle per localized density peak.

A. Tagged particle approach

An improved description can be achieved by viewing the $N$-particle system as an $N$-component mixture, in which each component corresponds to a single particle. This allows us to precisely locate particles, even after their densities started overlapping.

The DDFT for an arbitrary $m$-component mixture is a set of $m$-coupled equations,

$$\frac{\partial \rho_i(r_1,t)}{\partial t} = \nabla_1 \left[ \Gamma_0(r_1,t) \nabla_1 {\delta F_1}[\rho_1] \right],$$

where $i$ labels the species. By tagging each individual particle, Eq. (20) constitutes a set of $m$ partial differential equations in $D + 1$ dimensions ($D$ space dimensions and time) for the time-dependent one-body density fields. On the other hand, the Smoluchowski equation [Eq. (2)] for such a system is a partial differential equation in $D^m + 1$ dimensions for the probability distribution. A numerical solution of the Smoluchowski equation is therefore very demanding and only possible for very few particles.

Second, in real applications, one is not forced to tag each particle individually, and Eq. (20) offers complete flexibility as to which subsets of particles are associated with distinct species. For example, in the dynamic test particle calculations of Ref. [16], only a single particle was tagged and the rest of the system was treated as a second component.

The application of the full correction series with an infinite number of terms to such a tagged system corresponds to a physically sensible average, which suffers from neither particle fluctuations nor from the combinatorial effects discussed above. However, for large numbers of species and higher orders the canonical correction series becomes rather unwieldy, but progress can be made by only keeping some of the terms. This can be demonstrated by considering a pair of interacting particles, each carrying a distinct species label.

To first order, the canonical correction for such a binary mixture is given by

$$\langle A \rangle^c = \langle A \rangle^{gc} + f_1^{(1)}(A) + f_1^{(00)}(A) + f_1^{(01)}(A),$$

where the correction terms are given by

$$f_1^{(1)}(A) = -\frac{1}{2} \frac{\partial \langle N_1 \rangle}{\partial \mu_1} \frac{\partial^2 A}{\partial \langle N_1 \rangle},$$

$$f_1^{(00)}(A) = -\frac{1}{2} \frac{\partial \langle N_0 \rangle}{\partial \mu_0} \frac{\partial^2 A}{\partial \langle N_0 \rangle},$$

$$f_1^{(01)}(A) = -\frac{\partial \langle N_0 \rangle}{\partial \mu_1} \frac{\partial^2 A}{\partial \langle N_0 \rangle \partial \langle N_1 \rangle}.$$
As we have shown in Sec. III, this corresponds to a first-order suppression of the unphysical self-interaction, as each component consists of only one particle. Neglecting the mixed terms of the full canonical correction series leads to a system where each component does not interact with itself but only with the other components.

The conclusion which we draw from this simple test is that a suppression of the self-interaction in a system of $N$-tagged density peaks, in which each peak represents a separate species and starts from sharp initial conditions, results in a reasonable approximation to the true Brownian dynamics at short and intermediate times. However, the prohibitive numerical demands of performing the canonical transformation for more than a few particles (especially if higher orders of correction are required) make desirable an alternative scheme for eliminating the self-interaction.

B. Eliminating the self-interaction

1. Widom-Rowlinson model

A well-established model for which particles of the same species do not interact is the Widom-Rowlinson (WR) model [25,26]. An approximate density functional for the $m$-component WR model has been derived [27], which provides a reasonable description of both the bulk phase behavior and structural correlations. We propose to identify each species of an $N$-component WR model with an individual particle. Employing the Widom-Rowlinson functional in this fashion guarantees the absence of self-interactions but treats the interactions in an approximate fashion as the functional is not exact. In equilibrium with finite particle number (e.g., in confined geometries) the Percus and tagged WR functionals will yield different results. Due to the absence of unphysical self interactions, the equilibrium density profiles of the tagged WR model should lie closer to the results of Brownian dynamics simulation than those obtained from the Percus functional.

In one dimension, the approximate excess free energy of the WR model is given by Ref. [27],

$$F_{\text{ex}}^{\text{WR}}(\rho_i) = \int dx \Phi(x) \quad \Phi = \sum_{j=0}^{m} n_j \phi_j,$$

with weighted densities $n_0$ and $n_3$ for each component of the mixture. $\phi_j$ are the first derivatives of the 0D excess free energy with regard to $n_j$ and given by

$$\phi_j = \ln \left( 1 - m + \sum_{j=1}^{m} \exp(z_j) \right) - z_j.$$

The fugacities $z_j$ can be calculated from the implicit equation

$$z_j \exp(z_j) = \left[ 1 - m + \sum_{j=1}^{m} \exp(z_j) \right] n_j.$$

The one-particle direct correlation function required to calculate the average interaction force [Eq. (18)] is obtained by functional differentiation of the excess free energy.

2. Subtraction of the self-energy

A simpler, albeit ad hoc method of suppressing the self-interaction is to first calculate the excess free energy of the full $N$ component mixture using the Percus functional [Eq. (12)] and then subtract the individual excess free energy of each density peak. The remainder thus provides an approximation to the desired excess free energy arising from interaction between different species. This prescription amounts to employing the "subtraction" functional

$$F_{\text{ex}}^{\text{sub}}(\rho_i) = F_{\text{ex}}(\rho_i) - \sum_i F_{\text{ex}}(\rho_i)$$

in the DDFT Eq. (20). While the ansatz Eq. (25) is not justified from fundamental principles, it nevertheless has a certain physical appeal. In particular $F_{\text{ex}}^{\text{sub}}$ vanishes for the case of a single particle and becomes exact for many particles in the low density limit. A similar approach was taken in Ref. [16], in which an explicit self-interaction term of a tagged particle was omitted from the Ramakrishnan-Yussouff functional [28] in order to recover the exact single-particle diffusion.

V. NUMERICAL RESULTS FOR SEVERAL RODS

A. Free expansion

In order to compare the dynamics generated by the WR functional [Eq. (22)] with those generated by the subtraction functional [Eq. (25)], we consider the free diffusion of five rods from sharp initial conditions. The initial delta function distributions are each separated by 2.5 particle diameters (corresponding to a volume fraction of around 0.4). This choice ensures that the density fields have not entered the low-density regime for the intermediate times at which overlap between neighboring density peaks becomes significant. The present test, which is similar to that considered in Ref. [3], thus constitutes a genuine test of the functional Eqs. (22) and (25) beyond the second virial level. In Fig. 4 we show the time evolution of the total density,

$$\rho(x,t) = \sum_{i=1}^{m} \rho_i(x,t),$$

with $m = 5$, for three different times generated using the Percus Eq. (12), WR Eq. (22), and subtraction Eq. (25) functionals in the multicomponent DDFT Eq. (20).

As was the case for an isolated particle (see Fig. 1), the short-time relaxation of the density peaks generated by the Percus functional is too fast when compared with the Brownian dynamics simulation data. In contrast, the relaxation of the total density predicted by both the WR and subtraction functionals is in almost perfect agreement with the simulation results. This good level of agreement supports our argument that the self-interaction is the primary source of error induced by a grand-canonical generating functional, at least for short and intermediate times. Close inspection of the data reveals that the subtraction functional describes the simulation data slightly better than the WR functional, but the difference is marginal. However, if the initial packing of the rods is denser, so that larger local densities occur, the WR functional becomes less reliable and predicts an unphysically fast relaxation, while
2. Relaxation through a highly correlated state

We now consider a further test case for which rods with a length of 1.6 relax from sharp initial conditions in a periodic external potential \( \beta V_{\text{ext}}(x) = -2 \sin(2\pi x) \). The initial positions are chosen such that a particle is located at every second potential minimum. This situation was originally suggested in Ref. [3] as a toy model for the study of slow dynamics. The external potential and rod length are constructed in such a way that two rods cannot be simultaneously at the bottom of neighboring potential minima. Transport of particles between neighboring minima thus requires a correlated motion of all \( N \) rods and it is the rarity of such events which leads to a long relaxation time. For long times, an equilibrium solution is reached in which each potential well is equally populated.

In Fig. 5 we compare the time evolution of the total density from standard DDFT employing the Percus functional [Eq. (12)] and from a tagged particle calculation employing the subtraction functional [Eq. (25)] with Brownian dynamics data.

For short times, the relaxation generated by the Percus functional is faster than that of the subtraction functional, in keeping with the intuition obtained from the case of a single free particle (see Sec. III). Surprisingly, for later times the relaxation of the subtraction functional profile overtakes that of the Percus profile, arriving more quickly at the equilibrium distribution in which, on average, half a rod is located in each well. Use of the Percus functional thus yields better agreement with the simulation data than the supposedly superior subtraction functional. However, the good performance of the standard theory arises from a fortuitous cancellation of errors, which can be appreciated by looking at the time evolution of a single tagged density peak.

In Fig. 6 we show the time evolution of the tagged density corresponding to the third rod (labeling from left to right in Fig. 5). The other tagged densities evolve identically with time. Inspection of the profiles for \( t = 5 \) reveals the strange behavior of the tagged density from the Percus functional. Physically, it is reasonable to expect that the density peak will first diffuse into the neighboring wells before spreading further to the next-nearest-neighbors, and so on. However, the Percus DDFT predicts that the density in the next-nearest-neighbor well grows more rapidly than that in the neighboring well. For intermediate times, the unphysically large density that has built up in the two next-nearest-neighbor wells then pushes back on the central peak and slows its decay (i.e., generates a component of the self interaction that tends to confine the remaining density in the central peak).

In contrast to the Percus functional, the subtraction functional has a strongly reduced self interaction and does not suffer from an unphysical decay of the tagged density. Unfortunately, the improved description provided by use of the subtraction functional destroys the error cancellation presented by the Percus functional profile and thus relaxes much faster than the simulation data. The relative relaxation rate of the two
approaches can be appreciated from the inset to Fig. 6, which shows the difference in total density between neighboring potential minima.

The important message that emerges from this test case is that a qualitatively good description of the total density relaxation is not sufficient to conclude that the DDFT is really capturing correctly the underlying dynamics of the tagged density profiles.

C. Cage dynamics

In two recent papers, Hopkins et al. [16,17] have shown that standard grand-canonical DDFT (in three-dimensions) can reproduce the two-step relaxation of the self part of the van Hove function [22] characteristic of glass-forming systems. At sufficiently high volume fractions, a divergent relaxation time was identified. In these studies, the one-component Gaussian-core [17] and hard-sphere [16] models were formally viewed as a two-component mixture consisting of \( N - 1 \) particles of species \( d \) and a single particle of species \( s \), thus tagging an arbitrary particle. A sharp initial condition was taken for particle \( s \) and the corresponding equilibrium distribution for the density field of the \( d \)-component

\[
\rho_d(r,0) = \delta(r), \quad (27)
\]

where \( \rho_b \) is the bulk density.

Interpretation of the results of Refs. [16,17] was complicated by the fact that an approximate quadratic density functional was employed in the DDFT equation. It remains an open question whether the observed slow dynamics, which arise from a minimum in the equilibrium free energy, are genuinely associated with the glass transition, or rather an indication of freezing within the theory. We think that some light can be shed on this issue by considering the very simple case of two rods confined between impenetrable walls. The unique ordering of the rods on the line restricts the kinetically accessible phase space. As this is a characteristic feature of arrested states in general, we believe that a robust account of the equilibrium tagged density distributions of two rods confined to a slit is a necessary prerequisite for any DDFT claiming a realistic description of dynamic arrest in higher-dimensional systems.

In Fig. 7 we compare the equilibrium total density from DDFT with simulation data for two rods confined by the potential

\[
\beta V_{\text{ext}}(x) = e^{\frac{x}{4.5}} + e^{\frac{4.5-x}{3.5}} \quad (29)
\]

with decay length \( l = 0.2 \). This gives a volume fraction around 0.2. For this situation it can easily be shown that the canonical equilibrium density of the left particle is given by

\[
\rho_l(x) = e^{-\beta V_{\text{ext}}(x)} \int_{x+d}^{\infty} dx' e^{-\beta V_{\text{ext}}(x')}. \quad (30)
\]

The primary observation to be made from Fig. 7 is that the Percus functional generates a packing structure at the wall with a well developed peak and a dip. The dip is absent in the exact profile and the peak is less pronounced. It should be recalled that the Percus functional profiles represent the exact grand-canonical result for a slit with \( \langle N \rangle = 2 \). The peak is slightly underestimated by the WR functional [Eq. (22)] and entirely absent from the subtraction functional profile. Overall the WR functional shows the best level of agreement with the simulation data for the total density \( \rho(x) \).

On the basis of the total density profiles shown in Fig. 7, it is tempting to conclude that the WR functional provides an acceptable description of the equilibrium density distribution. Indeed, this is true if one is interested only in the total density...
show the tagged densities corresponding to the same situation \( \rho \). DFT approaches yield identical equilibrium tagged profiles and the simulation data is dramatic. All of the rods can be well described using DDFT. However, the constrained order of the rods is encoded in the hierarchy of correlation functions capturing a true nonergodic transition. The constrained order of the rods is encoded in the hierarchy of correlation functions (see Sec. II B) in a complicated way, so one cannot expect that the DDFT employed in Refs. [16,17] is capable of capturing this most elementary “caging” dynamic, we find it unlikely that the DDFT employed in Refs. [16,17] is capable of capturing a true nonergodic transition. The constrained order of the rods is encoded in the hierarchy of correlation functions (see Sec. II B) in a complicated way, so one cannot expect that truncating this hierarchy preserves this property.

VI. DISCUSSION

We have analyzed the shortcomings of DDFT for the description of localized density distributions using a simple one-dimensional model of hard rods. This model has the advantage that the exact grand-canonical equilibrium functional is known, thus removing the possibility of dynamical artifacts arising from use of an approximate functional. It is well known among DDFT practitioners that the standard theory predicts a relaxational dynamic that is systematically too fast when benchmarked against the Brownian dynamics simulation. By employing a formal expansion of the canonical average, we conclude that this deficiency is due to the unphysical self-interaction arising from grand-canonical coupling to a reservoir.

In situations where few particles are strongly confined, the presence of a self-interaction leads to equilibrium density profiles that differ from those obtained in Brownian dynamics simulation. We anticipate that the self-interaction will become relevant for large systems in cases for which large unbalanced forces are present, e.g., during the relaxation of large-density gradients in a nonequilibrium profile. The corrections in the force resulting from removal of the self-interaction will modify the relaxation timescale and slow the relaxation relative to standard grand-canonical DDFT.

The only method by which the self-interaction can be removed is to individually tag the density field of each particle and then employ either the canonical expansion series, a Widom-Rowlinson-type model, or ad hoc subtraction of undesirable contributions to the excess free-energy functional. Of these three possibilities, the latter proved to be both the most reliable and simplest to implement.

Once the self-interaction has been dealt with appropriately, we find that the free expansion of any number of interacting rods can be well described using DDFT. However, the considered test case that focused on the collective motion of rods (see Sec. V B) demonstrated that the previously reported good agreement between DDFT and simulation [3] is due to cancellation of errors. Removal of the self-interaction corrects the unphysical aspects of the tagged density relaxation, with the consequence that the results for the relaxation of the total density become worse (even faster than standard DDFT). The fact that DDFT clearly has difficulty in describing the relaxation through highly correlated states raises suspicions about its ability to capture the slow dynamics of dense systems.

Although our one-dimensional model provides only a crude description of a real fluid, it nevertheless enables some of the subtleties associated with a partitioned phase space to be investigated within a simple setting. Motivated by the recent work of Hopkins et al. [16,17], we presented the case of two confined rods as a toy model for the cage effect, whereby particles in a glass are localized by the geometrical constraints of their neighbors. Given that the very simple geometrical constraints on the tagged particle densities are not respected (see Fig. 8), it would be remarkable if the same theory turned out to be capable of describing the spontaneous localization associated with the glass transition in two- and three-dimensions. One possibility is that the particular functional employed in Refs. [16,17], namely the quadratic Ramakrishnan-Yussouff functional [28], is somehow able to compensate for the errors arising from a grand-canonical treatment of the density. However, such a scenario would seem to require a highly nontrivial cancellation of errors.

By considering test cases in which finite numbers of rods are confined to a slit, we now strongly suspect that any theoretical approach that is closed on the level of the one-body density, such as DDFT, will be unable to describe localization of the tagged density fields (at least when employing the exact equilibrium functional). When working solely with the one-body density, effective interactions between the average quantities \( \rho_i(r) \) are implemented, whereas Brownian dynamics considers interactions between the density operators \( \hat{\rho}_i(r) = \sum_j \delta(r - r_j) \) before taking the average. This mean-field treatment of the interaction between tagged density fields appears to make the tunneling of tagged density into geometrically
forbidden regions unavoidable. We note that these limitations do not necessarily pose a problem for the established density functional theory of freezing [29]. In such calculations, the crystal is identified as a periodic variation of the total density field and no claim is made to identify any specific particle: the tagged densities are fully delocalized throughout the sample. To arrive at a theory that respects the geometrical constraints on the tagged density fields, it is necessary to go beyond a density-based description and consider explicitly the dynamics of the higher-order density correlators, i.e., going beyond the simplest adiabatic approximation. Indeed, the importance of improving upon the standard adiabatic approach was recently identified by Haataja et al. [30] as one of the most important open problems in DFT. A fundamental question is whether a finite-order truncation of the dynamical hierarchy [of which Eqs. (4) and (5) constitute the first member] can capture tagged density localization.

Integration of the Smoluchowski equation over all but two of the particle coordinates leads to an equation of motion for the pair density involving a weighted integral over the three-particle correlations. Specifically, one is required to evaluate integrals of the form

\[
\int dr_3 \left[ \rho^{(3)}(r_1, r_2, r_3) \rho^{(2)}(r_1, r_2) \right] [-\nabla \Phi(r_{13})].
\]  

(31)

The first factor in square brackets can be identified as the conditional probability to find a particle at \( r_3 \), given that there are particles at positions \( r_1 \) and \( r_2 \). The integral thus represents the average force acting on a particle at \( r_1 \) due to interactions mediated by the other particles (an analogous integral provides the indirect force on a particle at \( r_2 \)). Making an adiabatic approximation, the integral Eq. (31) may be replaced by \( \nabla_1 c^{(1)}(r_1), c^{(1)}(r_1) \) is the one-body direct correlation function at \( r_1 \) calculated in the presence of both the physical external field and a test particle fixed at \( r_2 \) (hence the parametric dependence upon \( r_2 \)). This leads to

\[
\frac{\partial \rho^{(2)}(r_1, r_2)}{\partial t} = \nabla_1 \cdot \left[ \rho^{(2)}(r_1, r_2) \nabla_1 \left( \frac{\delta F}{\delta \rho(r_1)} \right) \right] + \nabla_2 \cdot \left[ \rho^{(2)}(r_1, r_2) \nabla_2 \left( \frac{\delta F}{\delta \rho(r_2)} \right) \right].
\]  

(32)

Combining Eqs. (4), (5), and (32) leads to a closed theory for the dynamics of the one- and two-body density in the form of a pair of coupled first-order (in time) differential equations. A similar equation was employed in Ref. [31] [see their Eq. (9)] in which the three-body density was treated using a superposition approximation. Crucially, in Eq. (32) the free-energy functional to be differentiated contains an external field consisting of the physical external potential of interest, \( V_{\text{ext}}(r) \), plus a "test" particle held fixed at either position \( r_1 \) or \( r_2 \), as indicated by the subscript on the functional derivative. Equation (32) is still "adiabatic," in the sense that three- and higher-body correlations are determined from the nonequilibrium \( \rho(r, t) \) and \( \rho^{(2)}(r_1, r_2, t) \) using equilibrium statistical mechanical relations. Nevertheless, this extended theory goes beyond Eq. (11), as the pair density is no longer tied to the instantaneous value of the density and relaxes instead on a finite timescale. Note that the formal elimination of \( \rho^{(2)}(r_1, r_2) \) from the coupled equations generates, in principle, an equation for \( \rho(r_1) \) alone, which includes memory effects [32]. In practice, however, the nonlinearity of the equations does not allow an explicit form for the memory kernel to be obtained and memory effects remain implicit to the coupled system of equations.

In equilibrium, Eq. (32) predicts that the pair correlations are those obtained from a test-particle calculation using the chosen free-energy functional. The exact single-particle dynamics are recovered using the initial condition \( \rho^{(2)}(r_1, r_2, 0) = 0 \). More generally, the correct normalization of the initial pair density, \( \int dr_1 \int dr_2 \rho^{(2)}(r_1, r_2, 0) = N(N - 1) \), is preserved throughout the time evolution, which is not the case when applying the standard adiabatic approximation, i.e., calculating equilibrium pair correlations in the presence of the effective potential [Eq. (15)]. Although we defer a more detailed investigation of Eq. (32) to future work, preliminary calculations indicate the results for the nonequilibrium one- and two-body density of hard rods are considerably improved, relative to standard DDFT. Of particular interest will be the predictions of Eq. (32) for the density profile of confined systems in the limit of long times.

Apart from our preliminary investigations of Eq. (32), the only explicit test of the adiabatic approximation of which we are aware was performed using continuous-time Monte-Carlo simulations of the Potts model subject to Glauber dynamics [33]. In this study, initial simulation configurations were prepared that reproduced the equilibrium one-body occupation number profile but with nonequilibrium correlations between the occupation numbers. During the simulated time-evolution the relaxation of higher-order correlation functions caused the one-body profile to drift first out of equilibrium, passing through a sequence of transient intermediate states, before returning back to equilibrium at long times. These findings clearly cannot be reproduced by theories based on \( \rho(r_1, t) \) alone, as no distinction can be made between states with the same one-body profile but different higher-body correlations. This issue may prove to be important when considering systems with slow dynamics for which the one-body density remains constant during gradual structural relaxation processes.

ACKNOWLEDGMENTS

We thank M. Schmidt and A. J. Archer for stimulating discussions. Funding was provided by the Swiss National Science Foundation.