Molecular theory of Langevin dynamics for active self-diffusiophoretic colloids

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ABSTRACT
Active colloidal particles that are propelled by a self-diffusiophoretic mechanism are often described by Langevin equations that are either postulated on physical grounds or derived using the methods of fluctuating hydrodynamics. While these descriptions are appropriate for colloids of micrometric and larger size, they will break down for very small active particles. A fully microscopic derivation of Langevin equations for self-diffusiophoretic particles powered by chemical reactions catalyzed asymmetrically by the colloid is given in this paper. The derivation provides microscopic expressions for the translational and rotational friction tensors, as well as reaction rate coefficients appearing in the Langevin equations. The diffusiophoretic force and torque are expressed in terms of nonequilibrium averages of fluid fields that satisfy generalized transport equations. The results provide a description of active motion on small scales where descriptions in terms of coarse grained continuum fluid equations combined with boundary conditions that account for the presence of the colloid may not be appropriate.

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I. INTRODUCTION
Active matter systems can take many forms, ranging from those whose active agents are micro-organisms or synthetic colloids to active materials and gels, among many others. Since active matter is not at equilibrium, its properties often differ markedly from its equilibrium analogs, and this fact has prompted investigations that explore the mechanisms by which such systems function and their possible applications.

Here, we consider active colloidal particles that are self-propelled through a diffusiophoretic mechanism where chemical reactions, maintained out of equilibrium, take place on a catalyst that is asymmetrically distributed on the colloid and produce concentration gradients in reactants and products. Interactions of the colloid with chemical species under these nonequilibrium conditions give rise to fluid flows in the vicinity of the colloid as a consequence of momentum conservation, leading to propulsion of the active particle.

Active colloidal particles with micrometer sizes are frequently considered in experiments17–21 so that continuum descriptions of the fluid in which they move are adequate; however, on this length scale, thermal fluctuations cannot be neglected. As a result, stochastic descriptions, usually in the form of Langevin equations, are used to describe the motions of these particles. In its simplest form, the Langevin equation that describes the evolution of the velocity \( v \) of an active colloidal particle with mass \( M \) propelled by a self-diffusiophoretic mechanism is written as:

\[
M \frac{dv}{dt} = F_{sd} - \zeta v + F_{fl},
\]

where \( \zeta \) is a friction coefficient, \( F_{fl} \) is a random force, and the new ingredient that distinguishes this equation from that for simple equilibrium Brownian motion is \( F_{sd} \), the diffusiophoretic force. Under most conditions, the inertial term on the left-hand side of Eq. (1) can be neglected for micrometric particles in condensed
phases, and the overdamped limit of this equation is sufficient. The expression for the diffusiophoretic force, or the corresponding diffusiophoretic velocity, \( V_{ad} = F_{ad}/\zeta \), in overdamped descriptions, is often simply postulated or derived from continuum models of the fluid subject to boundary conditions that account for coupling to the colloid.

On smaller nanometer or even angstrom scales, continuum descriptions will break down since the dimensions of fluid particles may no longer be negligible on the scale of the colloid size. In these cases where the molecular nature of the fluid manifests itself in the vicinity of the colloid, it is difficult to describe fluid–colloid interactions through boundary conditions. Active colloids with linear dimensions on the order of a few tens of nanometers have been studied in the laboratory.\(^{23,24}\) While motions of these very small active particles are dominated by thermal noise, the characteristics of active motion persist and are observable. In addition, molecular dynamics simulations of very small active dimer colloids with linear dimensions of a few nanometers exhibit features of active motion due to catalytic chemical reactions on part of their surface. Even for these very small particles, the local fluid velocity fields, obtained by extensive averaging to remove thermal noise effects, show flow patterns that are characteristic of self-diffusiophoresis. This feature is reminiscent of the fluid velocity fields observed in early molecular dynamics simulations of tagged particle motion that lead to a long-time power law decay of velocity correlations.\(^{7,26}\) Such collective solvent motions contribute to the values of diffusion coefficients and form the microscopic basis for Stokes law relating the frictional force on the colloid to the viscosity of the solvent.\(^{27,33}\) In a similar way, the microscopic flow fields seen in the vicinity of tiny active particles point to the presence of coupling to fluid collective modes with hydrodynamic character and the operation of a diffusiophoretic mechanism on molecular scales.

In order to study Brownian motion on very small scales where continuum descriptions break down, a molecular perspective must be adopted, and molecular derivations of Langevin equations for inactive colloidal particles have been carried out. Perhaps the most complete description is that of Mazur and Oppenheim,\(^{21,22}\) where the statistical properties of the noise are determined for a massive Brownian particle in an equilibrium bath. Such derivations have been extended to situations where the fluid in which the Brownian particle moves is subjected to constraints that drive it out of equilibrium.\(^{33,34}\)

Similarly, to study active motion on very small scales, a molecular description is needed where the particulate nature of the solvent is taken into account, and assumptions on the large relative colloid to solute size are relaxed. In this paper, we present a molecular derivation of the Langevin equations that describe the translational and rotational dynamics of a rigid active self-diffusiophoretic colloidal particle in a nonequilibrium environment. Since the system must be out of equilibrium for active motion to take place, we make use of a statistical mechanical formulation that accounts for the constraints that maintain the system in a nonequilibrium state.

A Langevin description of the translational and orientational dynamics of a colloidal particle is obtained from the equations of motion for the entire system by projecting out the bath degrees of freedom. Because the bath is in a nonequilibrium state, a time-dependent projection operator formalism is required, where the projection operator averages dynamical variables over a nonequilibrium bath density that depends conditionally on the presence of a fixed colloid. The nonequilibrium density is expressed in terms of a local equilibrium density containing time-dependent local thermodynamic fields conjugate to microscopic hydrodynamic density fields. The conjugate fields are defined self-consistently by constraint conditions that require the nonequilibrium averages of the hydrodynamic densities to be given exactly at all points in the system by averages over the local equilibrium density.

In addition, since our description is fully microscopic, we show how to include catalytic reactive dynamics in a way that treats the reactive chemical species at a molecular level. The resulting generalized Langevin equations serve the dual functions of describing active diffusiophoretic dynamics on molecular scales and providing microscopic expressions for the transport properties that enter Langevin descriptions on larger scales.

Section II specifies the system comprising the colloid and its fluid environment, gives an expression for its Hamiltonian, and presents the Liouville equations that govern its evolution. Chemical species are defined in Sec. III in terms of microscopic reaction coordinates and species variables that depend on the internal coordinates of the reactive molecules. The densities and constraints that characterize and determine the nonequilibrium state of the system are presented in Sec. IV. The derivation of the generalized Langevin equations using nonequilibrium time-dependent projection operator methods is given in Sec. V, while in Sec. VI, it is shown how these general equations yield the Langevin equations for the linear and angular momenta of the active colloid. The diffusiophoretic force and torque that are responsible for the active motion are further discussed in Sec. VII, and Sec. VIII provides a link between the microscopic and continuum theories. Section IX gives the conclusions of this study. Additional details of the calculations are presented in Appendices A–C.

II. SYSTEM AND DYNAMICS

The physical system considered here consists of a single rigid colloid of arbitrary mass distribution and total mass \( M \) immersed in a multi-component fluid of molecules of mass \( m \).\(^{35}\) The fluid is composed of reactive molecules dilutely dispersed in a solvent in contact with reservoirs that isothermally feed and remove species from the system at boundaries that are spatially distant from the colloid. A typical configuration of the physical system consists of \( N_b \) molecules of the reactive species \( R \) and \( N_S \) solvent molecules \( S \), with \( N_S \gg N_b \). These fluid species are denoted by \( \nu \in \{S, R\} \). Each reactive molecule \( i \) with total mass \( m_i \) contains \( n_a \) chemically bound atoms with masses \( \{m[k] = 1, 2, \ldots, n_a\} \) and nuclear positions and momenta \( \mathbf{x}_i^R = (r_{i1}, p_{i1}), \ldots, (r_{in_a}, p_{in_a}) \) = \( (\mathbf{x}^R, \mathbf{p}^R) \). The coordinates and momenta of the collection of the \( N_b \) reactive molecules are denoted by \( \mathbf{x}_b^R = (\mathbf{x}_1^R, \ldots, \mathbf{x}_N^R) = (\mathbf{r}_b^R, \mathbf{p}_b^R) \). While the solvent molecules, also taken to have mass \( m \) for simplicity, can be described in a similar way, their internal degrees of freedom will play no role in this work, and only their center of mass positions and momenta will be considered, \( \mathbf{x}_S = (r_{N_b+1}, p_{N_b+1}), \ldots, (r_{N_b+N_S}, p_{N_b+N_S}) = (\mathbf{r}_S, \mathbf{p}_S) \).

The spherical colloid has a total of \( n_t \) catalytic \( C \) and noncatalytic \( N \) sites on its surface. The distribution of these sites on the surface is left arbitrary at this point and may be chosen to describe active colloids with various properties. For instance, if the catalytic sites are
confined to one hemisphere, the colloid is a Janus particle, while hollow silica colloids whose catalytic activity is due to enzymes attached to their surfaces often have an irregular distribution of catalytic regions. Although it is feasible to treat the internal motions of the components of the colloid to allow for energy exchange between the internal degrees of freedom of the colloid and the surrounding fluid molecules, we will assume that the small and rapid internal fluctuations of the positions of the components of the colloid around their equilibrium values are not physically important in an isothermal system. For the rigid model, the positions of the sites relative to the center of the colloid, \( S'(R) \), are at a fixed distance from the center of the colloid and can be written as \( S'(R) \equiv S^0 - R = \mathbf{A}^T \cdot \mathbf{S}^0 - R \), where \( \mathbf{A}^T \) is a rotation matrix known as the attitude matrix that converts vectors between the body-fixed and laboratory frames of reference and \( \mathbf{S}^0 \) are constant vectors specifying the location of a site relative to the center of the colloid in the body-fixed frame. The rotation matrix \( \mathbf{A}^T \) and its inverse \( \mathbf{A} \) are specified by a set of arbitrary Euler orientational angles \( \mathbf{\Theta} \). In general, the active colloid need not be significantly larger than the solvent in which it moves. Figure 1 shows the type of active colloid, reactive molecules with an internal structure, and structureless solvent molecules comprising the system under study.

In writing the sums over particles, it is convenient to define indicator functions \( \Theta^i_{\nu} \) where \( \Theta^i_{\nu} = 1 \) if molecule \( i \) is species \( \nu \) and \( \Theta^i_{\nu} = 0 \) otherwise. Using this notation to determine whether molecule \( i \) is a solvent molecule or a reactive solute, the nuclear Hamiltonian for a system with \( N \) fluid molecules may be written as

\[
H = \frac{P^2}{2M} + K_{\text{rot}} + \sum_{i=1}^{N} \Theta^i_{\nu} \left( \frac{P^2_i}{2m_i} + \sum_{k=1}^{N} \Theta^k_{\nu} H_{mi} \right) + U_i(r^{N_i}, r^{N_i}_m) + U_1(R, r^{N_i}, r^{N_i}_m).
\]

This Hamiltonian is the sum of the translational and rotational kinetic energies of the colloid, the kinetic energies of the centers of mass of the \( N_s \) solvent molecules, and the sum of the reactive molecule Hamiltonians,

\[
H_{mi} = \sum_{k=1}^{n_i} \left( \frac{P^{(k)}_{\nu m}^2}{2m_k} + V_m(r^{a}_{nu}) \right).
\]

where \( V_m(r^{a}_{nu}) \) is the potential function for the nuclei in chemically bonded molecule \( i \). Interactions among the fluid molecules are given by \( U_0 \), while \( U_1 \) describes the interactions of the fluid particles with the colloid.

In the laboratory frame, the time derivative of the relative site position vector \( S'(R) = \mathbf{A}^T \cdot \mathbf{S}^0 - R \) is given in terms of the angular velocities \( \omega \) by \( \dot{S}'(R) = \omega \times S'(R) = \mathbf{\dot{\Theta}} \cdot \nabla \mathbf{A}^T \cdot \mathbf{A} \cdot (\mathbf{S}^0 - R) \), from which one finds that the angular velocities are related to time derivatives of the angles by \( \omega = N^T \cdot \mathbf{\dot{\Theta}} \), where the elements of the matrix \( N \) are

\[
N_{ab} = \frac{1}{2} \epsilon_{bca} A_{ac} \nabla_{\Theta} A_{bd}.
\]

Here, \( \epsilon_{bca} \) is the Levi-Civita symbol, and the Einstein convention of a sum over repeated indices has been used. The rotational kinetic energy of the colloid is

\[
K_{\text{rot}} = \frac{1}{2} \mathbf{\dot{\Theta}} \cdot \mathbf{I}_m \cdot \mathbf{\dot{\Theta}} = \frac{1}{2} \mathbf{\dot{\Theta}} \cdot \mathbf{M} \cdot \mathbf{\dot{\Theta}},
\]

where \( \mathbf{I}_m \) is the moment of inertia tensor in the laboratory frame and the matrix \( \mathbf{M} = N \cdot \mathbf{I}_m \cdot N^T \). Defining the generalized momentum \( \mathbf{P} \) conjugate to the angles \( \mathbf{\Theta} \) as \( \mathbf{P} = \partial K_{\text{rot}} / \partial \mathbf{\dot{\Theta}} = \mathbf{M} \cdot \mathbf{\dot{\Theta}} \), the total Hamiltonian in Eq. (2) for the system with colloidal phase space coordinates \( X = (R, P, \Theta, \Pi) \) can now be written as

\[
H = \frac{P^2}{2M} + \frac{1}{2} \Pi \cdot \mathbf{M}^{-1} \cdot \Pi + H_0,
\]

which defines \( H_0 \), the bath Hamiltonian in the presence of the fixed colloidal particle. It will play a central role in the development that follows.

The bath Hamiltonian \( H_0 \) contains the \( V_m, U_0 \), and \( U_1 \) potential functions. While the potential function for the chemically bonded atoms in a molecule, \( V_m(r^{a}_{nu}) \), is generally a many-body potential, we assume that the non-bonded interactions between the atoms in different molecules and those between the atoms in a molecule and the solvent molecules are pair-wise additive. Consequently, we can

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**FIG. 1.** Illustrating showing the components of the system: The \( N_s \) structureless solvent particles \( S \) are represented by yellow spheres (the solvent is a dense fluid but only a few solvent molecules are shown in this visualization), the \( N_s \) reactive molecules \( R \) of species \( A \) (red) and \( B \) (blue) are composed of \( n_x \) atoms, and the colloid possesses both the catalytic (red) and noncatalytic (blue) sites. In this graphic, the interaction sites are configured to represent a spherical colloid comprised of irregularly distributed catalytic and noncatalytic sites similar to that in enzyme-powered active colloids, but other geometries and distributions can be considered. In this figure, we show the distance \( r^{(b)}_{(k)i} \) of nucleus \( k \) in molecule \( i \) of species \( A \) from the colloid site \( a \). This nucleus interacts with the colloid site \( a \) through the intermolecular potential \( V_{bi}(r_{(k)i}) \), where \( b \) denotes whether the site is catalytic or noncatalytic of \( N \). A similar notation is used for reactive species \( B \), while the distance of molecule \( m \) of the chemically inert solvent \( S \) from colloid site \( \beta \) is \( r^{\beta}_{m} \).
The time evolution of a dynamical variable \( B(x_{N_i}, x_{N_{m}}, X) \) is given by the Liouville equation,

\[
\partial_t B(t) = -\{H, B(t)\} = i\mathcal{L} B(t),
\]

where \( i\mathcal{L} \), the Liouville operator for the evolution of the entire system, is defined in terms of the Poisson bracket of the Hamiltonian and the dynamical variable. It can be written as \( i\mathcal{L} = i\mathcal{L}_c + i\mathcal{L}_o \), the sum of the Liouvillian for the colloid, \( \mathcal{L}_c \), and the Liouvillian for the bath in the presence of the fixed colloid, \( \mathcal{L}_o \). The Liouvillian for the colloid is

\[
i\mathcal{L}_c = \frac{P}{M} \cdot \nabla R - \nabla R \cdot U_{\perp} \cdot \nabla P + i\mathcal{L}_{rot}.
\]

The rotational part of the Liouvillian operator \( \mathcal{L}_{rot} \) can be decomposed into an operator for the free rotation of a rigid body and an operator for the orientationally dependent interactions,

\[
i\mathcal{L}_{rot} = i\mathcal{L}_{rot, f} - i\mathcal{L}_{rot, t} \cdot \nabla \theta \cdot U_{\perp} \cdot \nabla P + \Pi \cdot M^{-1} \cdot \nabla \theta - \nabla \theta \cdot K_{rot} \cdot \nabla P.
\]

The Liouvillian operator \( i\mathcal{L}_{rot, f} \) for the free rotation of a rigid body has the property that \( i\mathcal{L}_{rot, f} L = 0 \), where \( L = I_{m} \cdot \omega = N^{-1} \cdot \Pi \) is the angular momentum of the colloid. The torque on the colloid, \( T \), is given by the time derivative of the angular momentum vector,

\[
T = \dot{L} = \nabla \theta \cdot U_{\perp} \cdot \nabla P \left( N^{-1} \cdot \Pi \right) = -N^{-1} \cdot \nabla \theta \cdot U_{\perp}.
\]
that separates regions where the metastable chemical species lie.\(^{44}\)

In order to describe the change in the reaction dynamics when the reactive molecules interact with the colloid, it is useful to introduce a second scalar reaction coordinate that is the distance of the center of mass of the reactive molecule from an active site on the colloid, \(r_i^a(r_i^m) = |r_i - S_i| = |r_i - S_i(R)|\) as defined earlier, but now, the center of mass of a reactive molecule is given by \(r_i = \sum_{a} (m_a r_{i,a})\). The free energy along the vectorial reaction coordinate \((\xi_i(r_i^m), r_i^a(r_i^m))\) can be defined as \(W(\xi, r^a) = -\beta^{-1} \ln(P(\xi, r^a)/P_{eq})\), with the probability density of specified numerical values of the reaction coordinates, \((\xi, r^a)\), given by

\[
P(\xi, r^a) = \langle \delta(\xi_i(r_i^m) - \xi)\delta(r_i^a(r_i^m)) - r^a \rangle_i,
\]

where the angle brackets denote an average over the local nonequilibrium distribution defined below [Eq. (29)] and \(P_{eq}\) is a uniform probability density. The free energy \(W(\xi, r^a)\) has the form shown schematically in Fig. 2.

The species variables may be defined in terms of \((\xi_i(r_i^m))\) as

\[
\theta_i^\gamma(\xi_i) = \Theta_i^\gamma \delta H_j(\xi_i(r_i^m)),
\]

where \(H_j(\xi_i(r_i^m))\) restricts molecular configurations to species \(\gamma \in \{A, B\}\): \(H_A(\xi_i(r_i^m)) = H(\xi_i - \xi_r(r_i^m))\) and \(H_B(\xi_i(r_i^m)) = H(\xi_i(r_i^m) - \xi_r)\), with \(H\) being a Heaviside function. The local number density of reactive molecules at a field point \(r\) with origin at the center of the colloid is given by

\[
N_B(r) = \sum_{i=1}^{N} \Theta_i^\gamma \delta(r_{ic} - r),
\]

and it can be partitioned into the sum of the local number densities of the A and B species at this field point, \(N_B(r) = N_A(r) + N_B(r)\), where

\[
N_i(r) = \sum_{i=1}^{N} \Theta_i^\gamma \delta(r_{ic} - r).
\]

These densities are important quantities for the specification of the nonequilibrium state of the system and enter the reaction–diffusion equation for the system. The fluxes of these species densities in the presence of a fixed colloid are given by

\[
\dot{N}_i(r) = i \mathcal{L}_0 N_i(r) = \nabla_r \cdot j_i(r),
\]

where the local reaction rate and the number density fluxes of species \(\gamma\) are

\[
\dot{j}_i(r) = \sum_{\gamma} \Theta_i^\gamma \eta^{-1} \mathcal{P} \delta(r_{ic} - r).
\]

IV. NONEQUILIBRIUM STATE OF THE FLUID

In the microscopic theory of Brownian motion in an equilibrium system developed by Mazur and Oppenheim,\(^{12}\) the Langevin equation is obtained by applying a projection operator formalism in which the projection extracts the average of dynamical variables

\[\text{FIG. 2.} \text{ The upper panels of the plots } W(\xi, r^a) \text{ as a color-coded function of } \xi \text{ and } r^a. \text{ It shows the potential wells corresponding to the metastable } A \text{ and } B \text{ species separated by a free energy barrier. The lower panels show how } W(\xi, r^a) \text{ varies with } \xi \text{ at two chosen values of } r^a. \text{ The upper right panel is for an } r^a \text{ value where the reactive molecule is far from the colloid and } W(\xi, r^a) \text{ has a double-well structure with deep wells separated by a high barrier, while the lower left panel is for an } r^a \text{ value where the molecule interacts with the colloid and the barrier separating the two metastable states is low, and the reaction is much more likely that in the bulk fluid. The numbers on the axis labels are simply guides to illustrate the changes in the well depths and barrier heights.}\]
over the equilibrium bath density $\rho_e$ in the presence of a fixed colloid. This density is stationary under the Liouville operator $i\mathcal{L}_0$ for a system in which the colloid is held fixed.

However, active motion can take place only under nonequilibrium conditions, and the constraints that drive the system out of equilibrium must be specified. If the system is maintained out of equilibrium by an externally imposed reservoir, the entropy production of the fluid is nonzero at all times, and the bath density $\rho_b$ does not equilibrate to $\rho_e$ but instead evolves according to the Liouville equation for the bath in the presence of a fixed colloid,

$$\partial_t \rho_b(t) = -i\mathcal{L}_0 \rho_b(t).$$

To implement the constraints imposed by the external reservoirs, we use the statistical mechanical theory for transport processes in systems out of equilibrium. The nonequilibrium state of the fluid is determined by a set of conjugate fields that couple to the following local fluid fields: the number density $N_i(r)$ of the reactive solute species defined in Eq. (18), the total number density of fluid molecules,

$$N(r) = \sum_{i=1}^{N} \delta(r_i - r),$$

that is equal to the sum of solvent and solute densities, $N(r) = N_s(r) + N_d(r) + N_b(r)$, where the solvent density is

$$N_s(r) = \sum_{i=1}^{N} \Theta_i^s \delta(r_i - r),$$

the total momentum density of the centers of mass of the solvent and solute molecules,

$$g_s(r) = \sum_{i=1}^{N} p_i \delta(r_i - r),$$

and the energy density of the fluid particles in the presence of the colloid,

$$E_N(r) = \sum_{i=1}^{N} \left[ \Theta_i^s \frac{p_i^2}{2m} + \Theta_i^s H_{m_i} + U_0 + U_0 \right] \delta(r_i - r).$$

We further assume that the system is isothermal with temperature $T$ although the formulation can be generalized to accommodate temperature variations. Note that the constraints are applied to the species densities $N_i(r)$ and total number and momentum densities. They are not applied to the total reactive molecule density $N_b(r)$ since we are primarily interested in situations where the species densities are maintained out of equilibrium. We then consider the set of fluid fields,

$$A(r) = \{N_i(r), N(r), g_s(r), E_N(r)\},$$

and corresponding conjugate fields,

$$\phi_A(r, t) = \{\beta \rho_b(r, t), \beta \mu_b(r, t), -m v^2(r, t)\},$$

where $\beta = 1/(k_B T)$, with $k_B$ being Boltzmann’s constant. The local relative chemical potential of species $i$ is $\mu_i(r, t) = \mu_i(r, t) - \mu_0(r, t)$, while $v(r, t)$ is the local fluid velocity field. The approach can be generalized to include mode coupling contributions by expanding the set of variables to include all nonlinear products of the slowly varying fields.

The local nonequilibrium distribution function may be written as

$$\rho_L(r, t) = \frac{\prod_i (N_i^0 h_i^0)^{-1} e^{A(r)} \phi_A(r, t)}{\text{Tr}[\prod_i (N_i^0 h_i^0)^{-1} e^{A(r)} \phi_A(r, t)]},$$

where $\phi_A(r, t) = \int dr A(r) \cdot \phi_A(r, t)$, and $\lambda \in \{S, A, B\}$. The trace operation includes an integration over phase space and a sum over particle numbers and types,

$$\text{Tr}[\ldots] = \prod_{\lambda} \sum_{N_\lambda=0}^{\infty} \int dN_\lambda d^2 x_{ \lambda, 0} \ldots.$$

The values of the conjugate fields $\phi_A(r, t)$ are chosen such that the local nonequilibrium averages of the $A(r)$ variables in the presence of the colloid are given by their exact nonequilibrium averages in the presence of a fixed colloid,

$$\langle A(r) \rangle = \text{Tr}[\rho_L(r) A(r)] = \text{Tr}[\rho_b(t) A(r)] \equiv \langle A(r) \rangle_i.$$

Both $\rho_b(t)$ and $\rho_b(t)$ depend parametrically on the fixed position $r$ and orientation $\theta$ of the colloidal particle, explicitly through the interaction potential in the Hamiltonian and through the thermodynamic conjugate fields $\phi_A(r, t)$.

The local equilibrium distribution function $\rho_b(t)$ can be generalized to incorporate additional higher order conjugate fields that couple to nonlinear products of the hydrodynamic densities. The additional conjugate fields are important when considering the dynamics of multilinear densities in nonequilibrium systems where the densities can exhibit long range correlations. However, for linear densities of hydrodynamic fluid fields, the additional conjugate fields provide only small mode coupling corrections that can be neglected to a good approximation.

To study the self-diffusiophoretic motion of the colloid, the solute chemical potentials can be given specified values far from the particle to describe a nonequilibrium scenario in which fuel and product species are fed in or removed from the system using external reservoirs. In this circumstance, the fluid velocity field vanishes far from the colloid, and there are no net fluid flows, although fluid flows are produced in the vicinity of the colloid as part of the diffusiophoretic mechanism.

V. DERIVATION OF GENERALIZED LANGEVIN EQUATION

Preparatory to obtaining the equations of motion for the linear and angular momenta of the colloid, we first consider how the Langevin equation for a general function $V$.

$$\frac{d}{dt} D(t) = i\mathcal{L} D(t) + e^{i\Omega} i\mathcal{L} D(0).$$

The generalized Langevin equation is obtained from Eq. (32) by projecting out the bath degrees of freedom so that their effects are incorporated into frictional and random forces. In order to project out the dependence on the bath variables, we make use of the
time-dependent projector $P(t)$ defined by its action on an arbitrary function $f$:

$$P(t)f = \text{Tr}[\rho_p(t)f],$$

and its complement, $Q(t) = 1 - P(t)$. The adjoint of the projector $P(t)$ is $P^\dagger(t)$, defined by $P^\dagger(t)f = \rho_p(t)Tr[f]$. Following usual generalizations, the generalized Langevin equation is obtained by rewriting the propagator $U(0, t) = \exp\left(i\mathcal{L}t\right)$ in an equivalent form involving the time-ordered propagator $U_0(0, t) = \mathcal{T}_+ \exp\left(f_0(t)dt, i\mathcal{Q}(t_1)\right)$, where $\mathcal{T}_+$ is a time ordering operator that orders operators in an increasing order of their time argument. As shown in Appendix A, the evolution operators $U(0, t)$ and $U_0(0, t)$ are related by

$$U(0, t) = U(0, t)P(t) + Q(t)U_0(0, t) - \int_0^t dt_1 U(0, t)(\partial_t P(t_1))U_0(t, t_1) + \int_0^t dt_1 U(0, t_1)P(t)\mathcal{L}Q(t_1)U_0(t, t_1).$$

Inserting this expression for $U(0, t) = \exp\left(i\mathcal{L}t\right)$ into the equation of motion [32], we obtain

$$\frac{d}{dt}D(t) = e^{i\mathcal{L}t}P(t)D(t) + \frac{F_0(t)}{\mu} - \int_0^t dt_1 e^{i\mathcal{L}t_1}(\partial_t P(t_1))K_D(t_1, t) + \int_0^t dt_1 e^{i\mathcal{L}t_1}P(t_1)i\mathcal{K}D(t_1, t),$$

where we have defined $D = i\mathcal{L}D$ and

$$K_D(t_1, t_2) = \mathcal{Q}(t_1)U_0(t_1, t_2)\mathcal{Q}(t_2)\mathcal{D} = \mathcal{Q}(t_1)U_0(t_1, t_2)\mathcal{Q}(t_2)\mathcal{D},$$

and made use of the relation $P(t_1) + Q(t_1) = 1$ in writing the third term on the right-hand side of Eq. (35). The fluctuating force is given by $F_0(t) = K_D(0, t)$.

The integral terms in Eq. (33) can be evaluated as shown in Appendix B, and using these results, the generalized Langevin equation for $D(t)$ reads

$$\frac{d}{dt}D(t) = \text{Tr}[\rho_p(t)D(t)]X(t) + \frac{F_0(t)}{\mu} + \int_0^t dt_1 \left( -\frac{P(t_1)}{M} M_1(t, t_1) + \nabla_\mu(t_1) - \Pi(t_1)^T \cdot M_1(t, t_1) \right),$$

where we defined

$$M_1(t_1, t) = \text{Tr}[\nabla_\mu(t_1)K_D(t_1, t)],$$

$$M_2(t_1, t) = \text{Tr}[\rho_p(t_1)F(t_1)K_D(t_1, t)],$$

$$M_3(t_1, t) = \text{Tr}[\nabla_\mu(t_1)K_D(t_1, t)],$$

$$M_4(t_1, t) = \text{Tr}[\rho_p(t_1)K_D(t_1, t)].$$

but have not indicated the dependence of these quantities on $X(t_1)$ in Eq. (37). The matrix $M(t)$ in Eq. (37) corresponds to the mass-weighted kinetic matrix $\mathcal{M}$ defined by Eq. (5) evaluated at the fixed colloidal position and orientation at time $t_1$.

A. Approximate form of Langevin equation: Brownian motion scaling

Following the theory of Brownian motion [32], when $M \gg m$, it is useful to introduce scaled momenta, $p^* = \mu p$ and $\Pi^* = \mu \Pi$, where $\mu = (m/M)^{1/2}$ is a small parameter that gauges the magnitude of the colloidal momenta. The corresponding scaled colloidal Liouvillian is $i\mathcal{L}^* = \mu i\mathcal{L}^*$. The above results, along with the action of $\exp(i\mathcal{L}^* t)$, allow us to write the generalized Langevin equation for the colloid in scaled colloidal coordinates as

$$\frac{dp^*(t)}{dt} = -\Pi^* (t_1) \cdot M_1(t_1)^{-1} \cdot M_3(t_1, t) - \nabla_{\Pi^*} \cdot M_4(t_1, t).$$

Again, we have not indicated the dependence of the matrices of transport coefficients $M$ on $X$ for simplicity.

The relation between the exact and local nonequilibrium distributions, $\rho_p(t)$ and $\rho_l(t)$, respectively, is given in Eq. (C22). The fields $\mu(t, r)$ and $\nu(t, r)$ in this equation are assumed to be slowly varying in space, so we may associate a small parameter $\epsilon_0$ that gauges the size of the gradients of these fields. In addition, we assume that the reactions are rare events and associate another small parameter $\epsilon_1$ that gauges the magnitude of the reactive flux. Equation (C22) also contains a term $Q_\mu(t)F(t) \cdot \nu(t, r)$. The $Q_\mu$ projector removes the contributions to $F(t)$ that are proportional to the species and total number densities, leaving only contributions to the force that arise from internal molecular degrees of freedom. Neglecting such contributions, we have $\rho_p(t) = \rho_l(t) + O(\epsilon_0, \epsilon_1)$, and we can replace $\rho_p(t)$ by $\rho_l(t)$ in evaluating the $M$ functions, however, since $\text{Tr}[\rho_p(t)D]$ scales as $\mu$, this replacement cannot be made in this term.

Since $\nabla_{\mu(t)}p^*(t) = \beta (F_c - (F_c)_c)p^*(t)$ and $\nabla_{\mu(t)}p^*(t) = -\beta (\nabla_\mu U_{l} - \langle \nabla_\mu U_{l} \rangle)p^*(t)$, in this approximation, we have

$$M_1(t_1, t) = \text{Tr}[(\nabla_\mu(t_1)p^*(t_1))K_D(t_1, t)],$$

$$M_2(t_1, t) = \beta \text{Tr}[(\nabla_\mu(t_1))F(t_1)K_D(t_1, t)],$$

$$M_3(t_1, t) = \text{Tr}[\nabla_\mu(t_1)p^*(t_1)K_D(t_1, t)],$$

$$M_4(t_1, t) = \text{Tr}[\rho_p(t_1)K_D(t_1, t)].$$

Noting that $\nabla_\mu U_l = -N \cdot T$ and $\Pi^* \cdot M_1^{-1} \nabla_\mu U_l = -L \cdot I_m^{-1} \cdot T$, the last two terms of Eq. (42) can be written in terms of the scaled angular momentum $L^*$ as

$$-\Pi^* (t_1) \cdot M_1(t_1)^{-1} \cdot M_3(t_1, t) - \nabla_{\Pi^*} \cdot M_4(t_1, t) = (-\beta L^* (t_1) \cdot I_m^{-1} + \nabla_{L^*} \cdot M_{TD}(t_1, t)),$$

where $M_{TD}(t_1, t) = \text{Tr}[\rho_l(t_1)TK_D(t_1, t)]$. We also let $M_{TD}(t_1, t) = M_2(t_1, t)$. 
Using the equations given above, we can write \( M_{3D} \) and \( M_{T2D} \) more explicitly in the form of friction kernels,

\[
M_{3D}(t_1,t_2) = \left( \langle F_c - \langle F_c \rangle \rangle Q(t_1) U_0(t_1) (D - \langle D \rangle) \right)_{t_1,t_2},
\]

\[
M_{T2D}(t_1,t_2) = \left( \langle T - \langle T \rangle \rangle Q(t_1) U_0(t_1) (D - \langle D \rangle) \right)_{t_1,t_2},
\]

(46)

where \( U_0(t_1,t) \) is now taken to be the projected evolution operator with \( \rho_0 \) replaced by \( \rho \) and \( i\mathcal{L} \) by \( i\mathcal{L}_0 \) neglecting higher order \( \mu \) contributions.

VI. LANGEVIN EQUATIONS FOR LINEAR AND ANGULAR MOMENTA

Taking the dynamical variables \( D = P \) and \( D = L \), noting the fact that \( U_0(t_1,t) \approx \text{e}^{-L(t-t_1)}(1 + O(\mu) + O(\varepsilon_h)) \) and retaining only the lowest order terms of the small parameters \( \mu, \varepsilon_h \), and \( \varepsilon_s \) in Eq. (46), we obtain the coupled Langevin equations for the translational and rotational motion of the colloid in the unscaled coordinates,

\[
\frac{d}{dt} P(t) = \text{Tr}[\rho(t)F_c](R(t),\theta(t)) + F_B(t)
\]

\[
- \int_0^t dt_1 \frac{P(t_1)}{M} \cdot M_F(t_1,t)
\]

\[
- \int_0^t dt_1 \beta L(t_1) \cdot I_m(t_1)^{-1} \cdot M_T(t_1,t)
\]

(47)

and

\[
\frac{d}{dt} L(t) = \text{Tr}[\rho(t)T](R(t),\theta(t)) + T_B(t)
\]

\[
- \int_0^t dt_1 \frac{P(t_1)}{M} \cdot M_F(t_1,t)
\]

\[
- \int_0^t dt_1 \beta L(t_1) \cdot I_m(t_1)^{-1} \cdot M_T(t_1,t)
\]

(48)

where we used the notation \( F_B^p = F_B \) and \( F_B^b = T_B \) for the random force and torque. The generalized Langevin equation for the linear momentum takes the form of an ordinary Langevin equation by changing variables \( t' = t - t_1 \), taking \( P(t' - t_2) \approx P(t) \) on the fast time scale of the force correlation decay, and defining the friction tensor by

\[
\zeta = \beta \int_0^\infty dt' M_F(0,t')
\]

\[
= \beta \int_0^\infty dt' \langle (F_c - \langle F_c \rangle)(F_c - \langle F_c \rangle) \rangle.
\]

(49)

In a similar approximation, the rotational friction tensor \( \zeta_r \) can be defined as

\[
\zeta_r = \beta \int_0^\infty dt' M_T(0,t')
\]

\[
= \beta \int_0^\infty dt' \langle (T - \langle T \rangle)(T - \langle T \rangle) \rangle
\]

(50)

with analogous expressions for the cross coupling friction tensors \( \zeta_{ir} \) and \( \zeta_{ir} \) that couple translational and rotational motion.

Setting \( P = MV \), when the translational and rotational motions decouple, the Langevin equations take the final form

\[
\frac{d}{dt} V(t) = \text{Tr}[\rho_b(t) F_c](R(t),\theta(t)) - \zeta V(t) + F_B(t).
\]

(51)

\[
\frac{dL(t)}{dt} = \text{Tr}[\rho_b(t)T](R(t),\theta(t)) - \zeta_r L(t) \cdot I_m(t)^{-1} + T_B(t).
\]

(52)

where we used \( \zeta_r = \zeta_1 I \) and \( \zeta_r = \zeta_1 I \). For a spherical rotor, \( I_m(t) = I_1 U \) is diagonal and independent of time. In this case, the angular momentum is \( L(t) = I_1 \omega(t) \), and an equation of the Langevin form can be written for the angular velocity \( \omega(t) \).

VII. DIFFUSIOPHORETIC FORCE AND TORQUE

The mean force and torque in the Langevin equations, \( \text{Tr}[\rho_b(t) F_c](R(t),\theta(t)) \) and \( \text{Tr}[\rho_b(t)T](R(t),\theta(t)) \), respectively, are responsible for the active translational and rotational motions of the colloid. In the absence of constraints that drive the system out of equilibrium, both of these quantities vanish, and Langevin equations reduce the standard forms that describe the Brownian dynamics of inactive colloids.

The constraints described by the \( \phi_A(r,t) \) fields can be applied in various ways to specify the nonequilibrium state. For a self-diffusiophasic colloid, a simple constraint is the specification of the values of chemical potentials of the \( A \) and \( B \) species far from the colloid. To study more general aspects of diffusiophasic colloidal motion, the gradients of these chemical potentials could also be specified. Under such constraints, the fluid velocity fields vanish far from the colloid, although, as noted earlier, the active motion of the colloid will generate local variations of the concentration and velocity fields in the vicinity of the colloid as part of the diffusiophasic mechanism. In this section, we consider the forms that the diffusiophasic force and torque take under such constraints.

A. Force

Using momentum conservation, the force on the colloid can be written in terms of the local force on the fluid, \( F_c = -\int dr F_i(r) \), given in Eq. (C.241), as

\[
F_c = \sum_{a} \int dr \sum_{b=1}^{N} \sum_{k=1}^{C} \Theta^a_b \left( \nabla_r V_{SL}(r^a) \right) \delta(r_a - r)
\]

(53)

and \( F^a_r \) can be written in terms of the local solvent and \( r^{a\alpha} \)-dependent reactive molecule densities as

\[
F^a_r = \int dr \sum_{b=1}^{N} \sum_{k=1}^{C} \Theta^a_b \left( \nabla_r V_{SL}(r^a) \right) N_b(r) + \int dr \int dr^{\alpha} \sum_{b=1}^{N} \Theta^a_b \left( \nabla_r V_{SL}(r^a) \right) N_b(r, r^{\alpha}).
\]

(54)

The local \( r^{a\alpha} \)-dependent reactive molecule density is defined by

\[
N_b(r, r^{\alpha}) = \sum_{i=1}^{N} \Theta^a_b \delta(r_a - r) \delta(r^{a\alpha} - r^{\alpha}).
\]

(55)
The expression for the force on the colloid in Eq. (54) involves $N_b(r, r^{α})$ and not the $A$ and $B$ species densities that enter the constraint conditions in Eq. (31). We can rewrite it in terms of $N_b(r)$ using projectors that project $N_b(r, r^{α})$ onto the species densities. We let $p(r||r^α)$ be the conditional probability density of the molecular coordinates $r^{α}$, given a distance $r$ of the center of mass of the molecule from the colloid center, and define a projector $P_r$ and its complement $Q_r = 1 - P_r$, by

$$P_r f(r^{α}) = \frac{\gamma_r(r^{α}) p(r||r^α) f(r^{α})}{\int dr^{α} \gamma_r(ξ(r^{α})) p(ξ(r^{α})|r)},$$

$$Q_r f(r^{α}) = \int dr^{α} \gamma_r(ξ(r^{α})) p(ξ(r^{α})|r) - P_r f(r^{α}),$$

where $p(r||r^α) = \frac{\gamma_r(r^{α}) p(r||r^α) f(r^{α})}{\int dr^{α} \gamma_r(ξ(r^{α})) p(ξ(r^{α})|r)}$.

This is the conditional probability density with the internal molecular complement $P_r$ molecule from the colloid center, and define a projector $P_r$ and its complement $Q_r = 1 - P_r$, by

$$P_r f(r^{α}) = \frac{\gamma_r(r^{α}) p(r||r^α) f(r^{α})}{\int dr^{α} \gamma_r(ξ(r^{α})) p(ξ(r^{α})|r)}$$

The expression for the force on the colloid in Eq. (54) involves $N_b(r, r^{α})$ and not the $A$ and $B$ species densities that enter the constraint conditions in Eq. (31). We can rewrite it in terms of $N_b(r)$ using projectors that project $N_b(r, r^{α})$ onto the species densities. We let $p(r||r^α)$ be the conditional probability density of the molecular coordinates $r^{α}$, given a distance $r$ of the center of mass of the molecule from the colloid center, and define a projector $P_r$ and its complement $Q_r = 1 - P_r$, by

$$P_r f(r^{α}) = \frac{\gamma_r(r^{α}) p(r||r^α) f(r^{α})}{\int dr^{α} \gamma_r(ξ(r^{α})) p(ξ(r^{α})|r)}$$

$$Q_r f(r^{α}) = \int dr^{α} \gamma_r(ξ(r^{α})) p(ξ(r^{α})|r) - P_r f(r^{α})$$

is the conditional probability density with the internal molecular complement $P_r$ molecule from the colloid center, and define a projector $P_r$ and its complement $Q_r = 1 - P_r$, by

$$P_r f(r^{α}) = \frac{\gamma_r(r^{α}) p(r||r^α) f(r^{α})}{\int dr^{α} \gamma_r(ξ(r^{α})) p(ξ(r^{α})|r)}$$

$$Q_r f(r^{α}) = \int dr^{α} \gamma_r(ξ(r^{α})) p(ξ(r^{α})|r) - P_r f(r^{α})$$

where $T^a$ is the contribution to the total torque from interaction site $α$ on the colloid. Noting that the $Θ$-dependence of the interaction potential arises from the relative position $S^α(R) = S^α - R = A^α(Θ) \cdot S^α$ of the interaction site from the center of the colloid, we have

$$T^a = -N^{-1} \sum_{i=1}^{C} \Theta^b_1 \sum_{k=1}^{C} \Theta^r_1 \nabla_{r^α} V_{ab} \cdot \nabla_{r^α} V_{ab} (r^α_i)$$

$$+ \Theta^s_1 \sum_{k=1}^{C} \Theta^r_1 \nabla_{r^α} V_{ab} \cdot \nabla_{r^α} V_{ab} (r^α_i)$$

$$= N^{-1} \sum_{i=1}^{C} \Theta^b_1 \sum_{k=1}^{C} \Theta^r_1 \nabla_{r^α} S^α (R) \cdot \nabla_{r^α} V_{ab} (r^α_i)$$

$$+ \Theta^s_1 \sum_{k=1}^{C} \Theta^r_1 \nabla_{r^α} S^α (R) \cdot \nabla_{r^α} V_{ab} (r^α_i).$$

From the definition of the $N$ matrix in Eq. (4) with the use of the identity $e^{iαa} e^{ib} = δ_αδ_ab - δ_αδ_b$, we find that

$$e^{iα} N_{ab} = \frac{1}{2} (A_{αa} V_{ab} A_{β} - A_{αb} V_{ab} A_{β})$$

$$= A_{αb} V_{ab} A_{β}.$$

Inserting $N_b(r, r^{α}) = P_b N_b(r, r^{α}) + Q_b N_b(r, r^{α})$ into Eq. (54), we get

$$F^a = \sum_{b=1}^{C} \int dr N_b(r) \sum_{i=1}^{C} F^a_{ib}(r) + \Delta F^a$$

where $F^a_{ib}(r) = Θ^a \sum_{i=1}^{C} \nabla_{r^α} V_{ab}(r^α_i)\cdot p_y(r||r^α)$.

$$\Delta F^a = \sum_{b=1}^{C} \Theta^b_1 \int dr \sum_{i=1}^{C} \nabla_{r^α} V_{ab}(r^α_i)\cdot Q_b N_b(r, r^{α}).$$

Using $(N(r))_i = \sum_j (N_j(r))_i$ and the notation introduced in Eq. (31) where $(N(r))_i = n_i(r, t)$ and $(N_j(r))_i = n_j(r, t)$, the diffusiophoretic force may now be written as

$$\text{Tr}[p_b(t) F^a] = \int dr \left[ \sum_{b=1}^{C} (F^a_{ib}(r) - F^a_{ib}) \right]p_y(r||r^α) + \sum_{b=1}^{C} F^a_{ib}(r) n(r, t) + \text{Tr}[p_b(t) ΔF^a]$$

B. Torque

A similar calculation can be carried out for the torque starting from the expression

$$T = -N^{-1} \cdot \nabla_θ \sum_{a=1}^{n_a} U^a_θ$$

$$= - N^{-1} \cdot \nabla_θ U^a_θ = \sum_{a=1}^{n_a} T^a,$$

where $T^a$ is the contribution to the total torque from interaction site $α$ on the colloid. Noting that the $Θ$-dependence of the interaction potential arises from the relative position $S^α(R) = S^α - R = A^α(Θ) \cdot S^α$ of the interaction site from the center of the colloid, we have

$$T^α = -N^{-1} \sum_{i=1}^{C} \Theta^s_1 \sum_{k=1}^{C} \Theta^r_1 \nabla_{r^α} V_{ab} \cdot \nabla_{r^α} V_{ab} (r^α_i)$$

$$+ \Theta^s_1 \sum_{k=1}^{C} \Theta^r_1 \nabla_{r^α} V_{ab} \cdot \nabla_{r^α} V_{ab} (r^α_i)$$

$$= N^{-1} \sum_{i=1}^{C} \Theta^s_1 \sum_{k=1}^{C} \Theta^r_1 \nabla_{r^α} S^α (R) \cdot \nabla_{r^α} V_{ab} (r^α_i)$$

$$+ \Theta^s_1 \sum_{k=1}^{C} \Theta^r_1 \nabla_{r^α} S^α (R) \cdot \nabla_{r^α} V_{ab} (r^α_i).$$

Since $A_{α} \cdot A_{β} = 0$.

Considering

$$\left( N^{-1} \right)_{ab} \nabla_{r^α} S^α (R) = -\left( N^{-1} \right)_{ab} \nabla_{r^α} A_{αb} S^α (R)$$

and taking the relation above into account, we get

$$\left( N^{-1} \right)_{ab} \nabla_{r^α} S^α (R) = -\left( N^{-1} \right)_{ab} \nabla_{r^α} A_{αb} S^α (R)$$

$$= -\left( N^{-1} \right)_{ab} \nabla_{r^α} S^α (R).$$

Using this relation in Eq. (64), we find the simple result

$$T^α = S^α (R) \cdot \left[ \sum_{i=1}^{C} \sum_{k=1}^{C} \Theta^s_1 \Theta^r_1 \nabla_{r^α} V_{ab}(r^α_i) + \Theta^s_1 \sum_{k=1}^{C} \nabla_{r^α} V_{ab}(r^α_i) \right]$$

$$= S^α (R) \cdot F^α.$$

The average of the diffusiophoretic torque then adopts a form that is analogous to that for the diffusiophoretic force,

$$\text{Tr}[p_b(t) T^α] = \int dr \left[ \sum_{a=1}^{n_a} n_i(r, t) \sum_{b=1}^{C} (F^a_{ib}(r) - F^a_{ib}) \right] + \sum_{a=1}^{n_a} T^a \cdot \text{Tr}[p_b(t) ΔF^a]$$

$$= \sum_{a=1}^{n_a} T^α \cdot \text{Tr}[p_b(t) F^a]$$

where $T^α = S^α (R), \Theta (t)$ are the positions of the sites $α$ relative to the center of the colloid at time $t$.

C. Contributions to force and torque

The diffusiophoretic force and torque in Eqs. (62) and (67) have several contributions. The first two contributions involve the local
equilibrium averages of the species and total density fields whose values are fixed by the constraints to give the exact nonequilibrium values of these quantities. These average fields may be determined from the solutions of the generalized hydrodynamic equations they satisfy. The last terms still retain the averages over the exact nonequilibrium density.

The terms involving $\text{Tr}[\rho_b(t)\Delta F_c]$ are expected to be small. While the introduction of the species densities $N_s(r)$ accounts for nonequilibrium effects through the reaction, the projected microscopic reactive molecule density $Q_sN_s(r, r')$ that enters $\text{Tr}[\rho_b(t)\Delta F_c]$ accounts for a nonequilibrium effect in the internal molecular degrees of freedom induced by the reaction. While such nonequilibrium effects can be taken into account, they are not a dominant effect and are expected to be small in most situations.

With these approximations $\text{Tr}[\rho_b(t)\mathcal{F}_c] = \langle \mathcal{F}_c \rangle$, and $\text{Tr}[\rho_b(t)T] = \langle T \rangle$, the diffusiophoretic force and torque are given by

$$\langle \mathcal{F}_c \rangle = \int dr \left[ \sum_{b,N} C \mathcal{F}_{sb}(r) n(r,t) \right. + \left. \sum_{b} \left( C \mathcal{F}_{sb}(r) - \mathcal{F}_{sb}(r) \right) \right] n_{b,r}(t), \quad (68)$$

$$\langle T \rangle = \sum_{a=1}^n S^a(t) \int dr \left[ n(r,t) \left( C \mathcal{F}_{sb}(r) - \mathcal{F}_{sb}(r) \right) \right] + \sum_{i} n_i(r,t) \left( C \mathcal{F}_{sb}(r) - \mathcal{F}_{sb}(r) \right) \right] \quad (69)$$

and their evaluation requires a knowledge of the local nonequilibrium averages of the hydrodynamic fields, which we consider below.

### D. Reaction-diffusion equations for species densities

The generalized hydrodynamic equations for the nonequilibrium averages $A(r, t)$ of slowly varying densities $A(r)$ of microscopic variables can be derived by noting that

$$\partial_t A(r, t) = \text{Tr}[\partial_t \rho_b(t)A(r)] = -\text{Tr}[i\mathcal{L}_0 \rho_b(t)A(r)] = \text{Tr}[\rho_b(t)\mathcal{J}_0 A(r)], \quad (70)$$

Using the relation between $\rho_b(t)$ and the local equilibrium density $\rho_b(t)$ established in Appendix C, the hydrodynamic equations assume the form

$$\partial_t A(r, t) = \langle \dot{A}(r) \rangle + f_{\mathcal{J},A}(r) - \int dt_1 \langle \mathcal{F}_{\mathcal{J},A}(r, t_1) \mathcal{F}_{\mathcal{J},A}(r') \rangle \bar{\phi}_A(r', t_1), \quad (71)$$

where

$$U_Q(t, t, t) = T_0 \exp \left\{ \int_0^t dt \mathcal{Q}_A(t) \mathcal{L}_0 \right\},$$

which follows by taking the Hermitian conjugate of Eq. (C13), and the random force is given by $f_{\mathcal{J},A}(r, t) = \text{Tr}[\rho_b(0) \mathcal{F}_{\mathcal{J},A}(r, 0, t)]$. From the general expression (71), a set of coupled equations for local nonequilibrium species densities and total number and momentum densities can be written that depend on their corresponding conjugate fields. The solutions of these equations can then be inserted into the expressions for the diffusiophoretic force and torque to complete the calculation of the quantities. To illustrate how to carry out this program, consider the equation of motion for the average species number density fields $n_i(r, t)$. For simplicity, we suppose that the Péclet number is small, $Pe = V_b R_i / D_i \ll 1$, so that advective effects can be neglected. Here, $R_i$ is the colloid radius, and $D_i$ is the diffusion coefficient of species $i$. When advective effects are small, the evolution of the number densities is independent of the fluid flow field. In this case, the evolution equation reads

$$\partial_t n_i(r, t) = f_{\mathcal{J},n_i}(r, t)$$

$$- \int dt_1 \langle \mathcal{F}_{\mathcal{J},n_i}(r, t_1) \mathcal{F}_{\mathcal{J},n_i}(r') \rangle \bar{\phi}_{n_i}(r', t_1), \quad (73)$$

where we have used the summation convention and

$$\mathcal{F}_{\mathcal{J},n_i}(r, t, t_1) = U_Q(t, t, t_1) \mathcal{Q}_A(t) \mathcal{L}_0 N_s(r) \quad = U_Q(t, t) \mathcal{F}_{\mathcal{J},n_i}(r), \quad (74)$$

with the random force given by $f_{\mathcal{J},n_i}(r, t) = \text{Tr}[\rho_b(0) \mathcal{F}_{\mathcal{J},n_i}(r, 0, t)]$. The random force vanishes if the initial condition is the local equilibrium distribution and will be neglected here.

The $\mathcal{F}_{\mathcal{J},n_i}(r, t, t_1)$ functions evolve on a short time scale $\tau_m$ in view of the projected dynamics. Consequently, the time-ordered evolution operator $U_Q(t, t, t)$ can be simplified by replacing the projectors $\mathcal{Q}_A(t)$ by $\mathcal{Q}_A(t)$ so that $U_Q(t, t, t) \approx e^{\mathcal{Q}_A(t) \mathcal{L}_0}$. Using this approximation and making the substitution $t_1 = t - \tau$ in the integral, we have

$$\int_0^t dt_1 \langle \mathcal{F}_{\mathcal{J},n_i}(r, t_1) \mathcal{F}_{\mathcal{J},n_i}(r') \rangle \bar{\phi}_{n_i}(r', t_1)$$

$$\equiv \left[ \int_0^\infty dt \left\{ \left( e^{\mathcal{Q}_A(t) \mathcal{L}_0} \mathcal{F}_{\mathcal{J},n_i}(r) \right) \mathcal{F}_{\mathcal{J},n_i}(r') \right\} \right] \bar{\phi}_{n_i}(r', t), \quad (75)$$

where in the last line, we replaced the upper limit $t$ in the integral by infinity for $t \gg \tau_m$ and replaced $\bar{\phi}_{n_i}(r', t - \tau)$ by $\bar{\phi}_{n_i}(r', t)$.

Using Eqs. (19) and (20) to obtain $\mathcal{F}_{\mathcal{J},n_i}(r) = \mathcal{I}^A_{n_i}(r) - \nabla \cdot j_{n_i}(r)$, along with Eq. (75), and neglecting cross coupling between reaction and diffusion, the generalized reaction–diffusion equation (73) can be written as

$$\partial_t n_i(r, t) = -L_{\mathcal{I},n_i}(r, r') \bar{\phi}_{n_i}(r', t)$$

$$+ \nabla \cdot L_{\mathcal{J},n_i}(r, r') \bar{\phi}_{n_i}(r', t), \quad (76)$$

where the reaction and diffusion transport terms are

$$L_{\mathcal{I},n_i}(r, r') = \int_0^\infty dt \left( j_{n_i}(r, t^+) \right)^A_{n_i}(r') \right\}, \quad (77)$$

$$L_{\mathcal{J},n_i}(r, r') = \int_0^\infty dt \left( j_{n_i}(r, t^+) \right)^A_{n_i}(r') \right\}, \quad (78)$$
and $\tau^*$ is used to denote evolution by projected dynamics. Since the chemical species are dilute in the solvent, the constraint condition [Eq. (70)] relating the nonequilibrium species densities $n_y(r, t)$ at time $t$ to the conjugate fields $\phi(r, t)$ can be inverted to the leading order in the fugacities of the dilute species. As a result, the chemical potentials can be written as $
abla \mu_y(r, t) = \nabla \gamma + k_B T \ln(n_y(r, t)/n_0)$, and substitutions into Eq. (76) yield closed equations for these local species density fields in the presence of the fixed colloid. For our self-diffusiophoretic colloid, these equations should be solved subject to constraints on the concentration fields at the boundaries where the system is in contact with reservoirs with fixed chemical concentrations. Although the solution of the fluid equations depends on where the colloid is located relative to the reservoirs, the behavior of the fluid densities in the vicinity of the colloid is determined by the local microscopic interactions of the fluid particles. An analogous treatment can be applied to the equations for the total number and momentum density fields. These transport equations, along with the expressions given above for the diffusiophoretic force and torque and colloid friction, provide a fully microscopic Langevin description of active self-diffusiophoretic dynamics.

VIII. RELATION TO CONTINUUM THEORIES

For large colloidal particles, it is appropriate to use the equations of continuum fluid mechanics to describe the dynamics of the fluid and to account for interactions of the fluid fields with the colloid through boundary conditions.\textsuperscript{9,10,36} Such descriptions are widely used for active colloidal particles with micrometer dimensions but will break down when the colloid size is no longer large compared to that of the fluid species, roughly in the lower portion of the nanometer range. In the low Reynolds number regime where inertia is unimportant, the computation of the velocity of an active colloidal particle propelled by a self-diffusiophoretic mechanism proceeds by solving the Stokes equation for the fluid velocity field and reaction–diffusion equation for the solute species subject to boundary conditions on the colloid surface.

As an example, consider a spherical Janus colloid with radius $R$ and orientation vector $\hat{n}$ pointing from the noncatalytic $N$ to catalytic $C$ hemispheres. The reaction $A \rightleftharpoons B$ occurs on its catalytic face, and the diffusion coefficients of these solute species are equal. Assuming perfect stick boundary conditions for the velocity field on the colloid yields the following equation for the self-diffusiophoretic velocity:\textsuperscript{2,23,57}

$$V_{sd} = \frac{k_B T}{\eta} \sum_y \Lambda_y (1 - \hat{n} \cdot \nabla n_y^*(r))^S$$

with $\Lambda_y = \frac{f_R}{\kappa} (r - R) \left( e^{-\beta V_{yc}(r)} - 1 \right)$, where $V_{yc}$ is the solute–colloid interaction potential, $\eta$ is the shear viscosity, and $\hat{n}$ is the unit normal to the surface. The surface average is defined as $
abla^S = \frac{1}{V} \int dr \sum_y \nabla n_y^*(r)$ for simplicity, we have assumed that the $y = A, B$ species particles interact with the both faces of the Janus particle with the same potential although this assumption need not be made. The $S$ superscript on the species density indicates that it is for the fluid phase outside the interaction zone where $\delta$ is the range of the solute–colloid potentials. The second equality in Eq. (79) used $\Lambda_A + \Lambda_B = \Lambda$. The result (79) is obtained by using the Navier–Stokes equations for the fluid velocity and the aforementioned reaction–diffusion equations, but the key features of this result can already be identified in the different contributions to the diffusiophoretic force in Eq. (68). As in the continuum calculation, we may assume that the solute and solvent species are structureless and that the interactions do not depend on the $C$ or $N$ labels of the surface sites. With these approximations, the contributions of the solute species $y$ to Eq. (68) can be evaluated as follows:

$$\int dr \sum_y F_{ny}(r) n_y(r) = -k_B T \int dr \sum_y \left( \nabla e^{-\beta V_{yc}(r)} \right) n_y^*(r, R)$$

$$= 4\pi k_B T \Lambda (1 - \hat{n} \cdot \nabla n_y^*(r))^S$$

where we have used the fact that $F_{ny}(r) = \nabla_r V_{ny}(r)$ for this simplified model and $n_A + n_B = n_0$ to write the third equality. Since the fluid–colloid intermolecular forces vanish outside the interaction zone, the integration is restricted to $R < r \leq R + \delta$. An approximate computation leading to the intermediate steps in Eq. (80) can be obtained by taking the density field in the interaction region as $n_y(r) = e^{-\beta V_{yc}(r)} n_y^*(r, R)$. Furthermore, we have taken $\hat{n}$ to be the polar axis for the integration, and $\mu = \cos \theta$. The last equality follows from the use of the identity

$$(1 - \hat{n} \cdot \nabla n_y^*(r))^S = \frac{\hat{n}}{R} \int_1^R \mu d \mu n_y^*(R, \theta).$$

In the overdamped regime, the diffusiophoretic velocity is related to the diffusiophoretic force by $V_{sd} = \zeta^{-1} (F_{ny})$. Taking the translational friction coefficient for stick boundary conditions, $\zeta = 6\pi \eta R$, we obtain an expression comparable to the known result (79) in the continuum limit.

To go beyond this simplified model, the complete calculation would require the use of the Navier–Stokes equations, which can also be deduced in the present formalism, as discussed in Sec. VII D, and which is coupled to the Langevin equations for the colloidal particle. The behavior of the fluid densities in the vicinity of the colloid is determined by the local microscopic interactions of the fluid particles with the colloid. These fluid densities typically exhibit rapid variations and structural and dynamic correlations near the colloid due to strong interactions of solute particles at short distances from interaction sites on the colloid. The determination of the appropriate boundary conditions that account for the complicated surface structure and dynamics in the fluid induced by the colloidal interactions requires a detailed analysis of the generalized reaction–diffusion and hydrodynamic equations in the interaction zone around the colloid. Through such analyses, a quantitative link between the present microscopic description and continuum treatments of self-diffusiophoresis for large colloidal particles can be made.
IX. CONCLUSION

The molecular-level derivation of the Langevin equations given in Eqs. (47) and (48) for an active particle whose propulsion arises from a diffusiophoretic mechanism allows one to assess the domain of validity of Langevin descriptions of such active systems that are often proposed on phenomenological grounds. The generalized Langevin equations incorporate features that become important on small length and time scales. These include the static structural correlations among fluid species and the active particle that complicate descriptions in which the effect of the colloid on the fluid is incorporated into boundary conditions, explicit treatment of both the solvent and solute species and their interactions with the active particle, and memory effects that enter because the time scales of the dynamics are not as well separated as when the active particle is orders of magnitude larger than the solvent species. They also provide a molecular basis for the active colloid Langevin equations obtained through treatments based on fluctuating chemohydrodynamics.

The diffusiophoretic force and torque in the Langevin equations are important quantities that differentiate these Langevin equations from those for ordinary Brownian motion. They contain contributions that depend on the local nonequilibrium averages of species density fields expected from continuum calculations; however, these fields themselves satisfy generalized hydrodynamic and reaction–diffusion equations, as seen in Eqs. (71) and (76). In addition, they have contributions that involve full nonequilibrium averages of the reactive molecules that cannot be expressed separately in terms of the fuel and product species densities.

Another important feature that emerges from the microscopic derivation is that all transport and dynamical diffusiophoretic factors have microscopic expressions in terms of Green–Kubo correlation functions. This permits one, at least in principle, to determine these quantities directly from molecular dynamics simulations. The translational and rotational friction tensors in Eqs. (49) and (50), respectively, can be computed by numerically evaluating autocorrelation functions of the force and torque imparted on the fixed colloid by the fluid under nonequilibrium conditions. Similarly, the transport coefficients in Eqs. (77) and (78) that enter the reaction–diffusion equation (76) in the presence of the colloid are amenable to evaluation by simulation. In particular, since the reactive species are treated at a molecular level that explicitly accounts for the dynamics of the nuclei comprising the molecules, the activated rate processes that take place on the colloid (or in the fluid) can be described in terms of suitable reaction coordinates whose specific forms depend on the reaction mechanism, and reaction rates can be computed using molecular dynamics employing rare event sampling methods for these slow processes.12 Thus, all transport properties are specified in molecular terms through correlation functions whose infinite-time integrals are well-defined since the dynamics involves a fixed colloidal particle12 or projected time evolution.

It is simple to extend the formalism presented here to describe thermophoretically active colloids in the presence of an external temperature gradient or to incorporate reactive events that are not iso-enthalpic. In addition, while most of the presentation in this paper considered a rigid colloid, the development is not restricted to this specific kind of active particle. The active particle may be any molecule or molecular aggregate with internal degrees of freedom so that the generalized Langevin equations presented can form a basis for the analysis of molecular simulations and experiments dealing with reactive dynamics in the presence of nonequilibrium constraints on molecular scales.

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APPENDIX A: EVOLUTION OPERATOR

The full system evolution operator satisfies the following equation:

$$\partial_t U(0,t) = U(0,t)i\mathcal{L},$$  \hspace{1cm} (A1)

while the time-ordered projected evolution operator $U_Q(0,t)$ satisfies

$$\partial_t U_Q(0,t) = U_Q(0,t)i\mathcal{L}Q(t),$$  \hspace{1cm} (A2)

whose formal solution can be written as

$$U_Q(0,t) = T. \exp\left(\int_0^t dt_i \mathcal{L}Q(t_i)\right),$$  \hspace{1cm} (A3)

where $T.$ is a time ordering operator that orders operators in an increasing order by their time argument.

The relation between these propagators can be established as follows: The evolution operators $U(0,t)$ and $U_Q(0,t)$ have the property $U(t_1,t_2)U(t_2,t_3) = U(t_1,t_3)$, with an analogous expression for $U_Q(t_1,t_2)$. To establish the relation between these operators, we let $U(0,t) = G(t)U_Q(0,t)$ so that the operator $G(t) = U(0,t)U_Q^{-1}(0,t)$, Its initial value is $G(0) = 1$. Since $U_Q(0,t)U_Q^{-1}(0,t) = 1$ by definition, $U_Q^{-1}(0,t)$ satisfies

$$\partial_t U_Q^{-1}(0,t) = -i\mathcal{L}Q(t)U_Q^{-1}(0,t).$$  \hspace{1cm} (A4)

Differentiation of the definition of $G(t)$ yields

$$\partial_t G(t) = U(0,t)i\mathcal{L}U_Q^{-1}(0,t) + U(0,t)\partial_t U_Q^{-1}(t,0),$$

$$= U(0,t)i\mathcal{L}G(t)U_Q^{-1}(0,t),$$  \hspace{1cm} (A5)

which, after integration, gives

$$G(t) = 1 + \int_0^t dt_i U(0,t_1)i\mathcal{L}G(t_1)U_Q^{-1}(0,t_1)$$
$$+ 1 + \int_0^t dt_i (\partial_t U(0,t_1))G(t_1)U_Q^{-1}(0,t_1).$$  \hspace{1cm} (A6)

Using this result, we may then obtain $U(0,t)$ as

$$U(0,t) = U_Q(0,t) + \int_0^t dt_i (\partial_t U(0,t_1))G(t_1)U_Q(t_1,t)$$
$$= U_Q(0,t) + \int_0^t dt_i \partial_t (U(0,t_1))G(t_1)U_Q(t_1,t)$$
$$- \int_0^t dt_1 U(0,t_1)\partial_t G(t_1)U_Q(t_1,t),$$  \hspace{1cm} (A7)

which can be rearranged to give Eq. (34) in the main text.
APPENDIX B: REDUCTION OF INTEGRAL TERMS IN EQ. (35)

Let $I$ denote the integral terms in Eq. (35),

$$ I = \int_0^t dt_1 e^{i\mathcal{L}t_1} \left[ \mathcal{P}(t_1) i \mathcal{L} \mathcal{K}_D(t_1, t) - (\partial_t \mathcal{P}(t_1)) \mathcal{K}_D(t_1, t) \right]. \quad (B1) $$

We have

$$ -(\partial_t \mathcal{P}(t_1)) \mathcal{K}_D(t_1, t) = \text{Tr} \left[ [i \mathcal{L} \rho_0(t_1)] \mathcal{K}_D(t_1, t) \right] $$

$$ = -\text{Tr} \left[ \rho_0(t_1) i \mathcal{L} \mathcal{K}_D(t_1, t) \right] $$

$$ = -\mathcal{P}(t_1) i \mathcal{L} \mathcal{K}_D(t_1, t). \quad (B2) $$

Using this result, along with $i \mathcal{L} - i \mathcal{L}_0 = i \mathcal{L}_1$, Eq. (B1) can be written as

$$ I = \int_0^t dt_1 e^{i\mathcal{L}t_1} \mathcal{P}(t_1) i \mathcal{L} \mathcal{K}_D(t_1, t) $$

$$ = \int_0^t dt_1 e^{i\mathcal{L}t_1} \text{Tr} \left[ \rho_0(t_1) \left( \frac{P}{M} \nabla R + F_c \cdot \nabla p + i \mathcal{L} \rho_0(t_1) \right) \right] \mathcal{K}_D(t_1, t). \quad (B3) $$

In addition,

$$ \text{Tr} \left[ \rho_0(t_1) \left( \frac{P}{M} \nabla \mathcal{K}_D(t_1, t) \right) \right] = -\frac{P}{M} \text{Tr} \left[ (\nabla \rho_0(t_1)) \mathcal{K}_D(t_1, t) \right] $$

$$ \text{Tr} \left[ \rho_0(t_1) i \mathcal{L} \mathcal{K}_D(t_1, t) \right] = -\Pi^\top \cdot M^{-1} \cdot \text{Tr} \left[ \nabla \rho_0(t_1) \mathcal{K}_D(t_1, t) \right] $$

$$ -\nabla \Pi \cdot \text{Tr} \left[ \rho_0(t_1) \nabla \mathcal{K}_D(t_1, t) \right]. \quad (B4) $$

since $\text{Tr} \left[ \rho_0(t_1) \mathcal{K}_D(t_1, t) \right] = 0$. We then have

$$ I = \int_0^t dt_1 e^{i\mathcal{L}t_1} \left\{ -\frac{P}{M} \cdot \mathcal{M}_1(t_1, t) + \nabla p \cdot \mathcal{M}_2(t_1, t) \right. $$

$$ -\Pi^\top \cdot M \cdot \mathcal{M}_3(t_1, t) - \nabla \mathcal{M}_4(t_1, t) \right\} $$

$$ + \int_0^t dt_1 \left\{ -\frac{P(t_1)}{M} \cdot \mathcal{M}_1(t_1, t) + \nabla p(t_1) \cdot \mathcal{M}_2(t_1, t) \right. $$

$$ -\Pi(t_1)^\top \cdot M^{-1} \cdot \mathcal{M}_3(t_1, t) - \nabla \mathcal{M}_4(t_1, t) \right\}, \quad (B5) $$

where $\mathcal{M}_1, \ldots, \mathcal{M}_4$ are defined in Eq. (38). Use of this expression yields Eq. (37).

APPENDIX C: $\rho_0(t)$ AND $\rho_L(t)$ DENSITIES

An explicit relation between $\rho_0(t)$ and $\rho_L(t)$ is required in order to express average values in a convenient form. For this purpose, we consider a projection operator

$$ \mathcal{P}(t)f = \left\{ f C(r_1) \right\}_0 \cdot \left( \mathcal{C}^\top \right)^{-1} (r_1, r_2) \cdot C(r_2) \quad (C1) $$

and its complement $\mathcal{Q}(t) = 1 - \mathcal{P}(t)$. The adjoint of this projector is defined by

$$ \mathcal{P}^\dagger(t) f = \text{Tr} [f C(r_1)] \cdot \left( \mathcal{C}^\top \right)^{-1} (r_1, r_2) \cdot C(r_2) \rho_L(t) \quad (C2) $$

and its complement $\mathcal{Q}^\dagger(t) = 1 - \mathcal{P}^\dagger(t)$. The vector $C(r) = \{1, \hat{A}(r)\}$ is expressed in terms of the deviations $\hat{A}(r) = \mathcal{A}(r) - \langle \mathcal{A}(r) \rangle$, of the fields $\mathcal{A}(r)$ in Eq. (27). Using this notation, we observe that 1 is not a field extending over space but a single number, while $\hat{A}(r)$ is a field variable; hence, we can write

$$ \mathcal{P}^\dagger(t) f = \text{Tr} [f \rho_L(t) $$

$$ + \text{Tr} [f \hat{A}(r_1)] \cdot \left( \mathcal{A}^\top \right)^{-1} (r_1, r_2) \cdot \hat{A}(r_2) \rho_L(t). \quad (C3) $$

Taking $f = \rho_0(t)$ and using the fact that $\text{Tr}[\rho_0(t) \mathcal{A}(r)] = \langle \mathcal{A}(r) \rangle$, we obtain $\mathcal{P}^\dagger(t) \rho_0(t) = \rho_L(t)$. We may then write $\rho_0(t) = \rho_L(t) + \mathcal{Q}^\dagger(t) \rho_0(t)$.

Applying this projector to Eq. (22), we have

$$ \mathcal{P}^\dagger(t) \partial_t \rho_0(t) = \text{Tr} [(\partial_t \rho_0(t) \mathcal{C}(r_1)] \cdot \left( \mathcal{C}^\top \right)^{-1} (r_1, r_2) $$

$$ \cdot \mathcal{C}(r_2) \rho_L(t) $$

$$ = \text{Tr} [(\partial_t \rho_L(t) \mathcal{C}(r_1)] \cdot \left( \mathcal{C}^\top \right)^{-1} (r_1, r_2) $$

$$ \cdot \mathcal{C}(r_2) \rho_L(t). \quad (C4) $$

Since the local nonequilibrium distribution function may be written as

$$ \rho_L(t) = \frac{\prod_i (N_i \hbar N_{\pi i})^{-1} e^{i C(r_i) \phi_c(r_i)}}{\text{Tr} \left[ \prod_i (N_i \hbar N_{\pi i})^{-1} e^{i C(r_i) \phi_c(r_i)} \right]}, \quad (C5) $$

with $\phi_c = (0, \phi_A)$, we have

$$ \partial_t \rho_L(t) = (\partial_t \mathcal{C}(r_i) \cdot \mathcal{C}(r_i) \rho_L(t) $$

$$ = (\partial_t \mathcal{C}(r_i) \cdot \hat{A}(r) \rho_L(t) \quad (C6) $$

and

$$ \mathcal{P}^\dagger(t) \partial_t \rho_0(t) = (\partial_t \mathcal{C}(r_i) \cdot \mathcal{C}(r_i) \rho_0(t) = \partial_t \rho_L(t). \quad (C7) $$

From this result, we can write

$$ \partial_t \rho_0(t) = -i \mathcal{L}_0 \mathcal{P}^\dagger(t) \rho_0(t) + \mathcal{Q}^\dagger(t) \rho_0(t) $$

$$ = -i \mathcal{L}_0 \rho_L(t) - i \mathcal{L}_0 \mathcal{Q}^\dagger(t) \rho_0(t) \quad (C8) $$

and

$$ \mathcal{Q}^\dagger(t) \partial_t \rho_0(t) = \partial_t \rho_0(t) - \partial_t \rho_L(t) = \partial_t \mathcal{Q}^\dagger(t) \rho_0(t). \quad (C9) $$

Using Eqs. (C8) and (C9), we have

$$ \partial_t \mathcal{Q}^\dagger(t) \rho_0(t) = -\mathcal{Q}^\dagger(t) \partial_t \rho_0(t) - \mathcal{Q}^\dagger(t) i \mathcal{L}_0 \mathcal{Q}^\dagger(t) \rho_0(t). \quad (C10) $$

To solve this equation, we introduce the projected propagator $U^\dagger_{\mathcal{Q}^\dagger}(0, t)$ that is the solution of the evolution equation

$$ \partial_t U^\dagger_{\mathcal{Q}^\dagger}(0, t) = -\mathcal{Q}^\dagger(t) i \mathcal{L}_0 U^\dagger_{\mathcal{Q}^\dagger}(0, t) \quad (C11) $$

and its inverse $(U^\dagger_{\mathcal{Q}^\dagger})^{-1}(0, t)$ whose evolution is given by

$$ \partial_t (U^\dagger_{\mathcal{Q}^\dagger})^{-1}(0, t) = (U^\dagger_{\mathcal{Q}^\dagger})^{-1}(0, t) \mathcal{Q}^\dagger(t) i \mathcal{L}_0 U^\dagger_{\mathcal{Q}^\dagger}(0, t). \quad (C12) $$

Formally, the solution of Eq. (C11) can be written as

$$ U^\dagger_{\mathcal{Q}^\dagger}(0, t) = \mathcal{T} \exp \left[ -\int_0^t dt_1 \mathcal{Q}^\dagger(t_1) i \mathcal{L}_0 \right]. \quad (C13) $$
where the time ordering operator $T_t$ orders operators with smaller time argument to the right of operators of larger time argument.

Defining an operator $G(t) = (U_{Q_A}^t)^{-1}(0, t) Q_A^t(t)\rho_0(t)$ and using the above results, its differential equation is given by

$$\partial_t G(t) = -(U_{Q_A}^t)^{-1}(0, t) Q_A^t(t)iL_0p_L(t). \tag{C14}$$

Integration of this equation gives

$$G(t) = Q_A^0(0)\rho_0(0) - \int_0^t dt_1 \left( (U_{Q_A}^t)^{-1}(0, t_1) Q_A^t(t_1)iL_0p_L(t_1) \right), \tag{C15}$$

from which we find

$$Q_A^t(t)\rho_0(t) = U_{Q_A}^t(0, t) Q_A^0(0)\rho_0(0) - \int_0^t dt_1 \left( U_{Q_A}^t(t_1) Q_A^t(t_1)iL_0p_L(t_1) \right). \tag{C16}$$

It follows that

$$\rho_b(t) = \rho_L(t) + U_{Q_A}^t(0, t) Q_A^t(0)\rho_0(0) - \int_0^t dt_1 \left( U_{Q_A}^t(t_1) Q_A^t(t_1)iL_0p_L(t_1) \right). \tag{C17}$$

This equation may be written in another form by using $-iL_0p_L(t) = -\hat{A}(r) \cdot \phi_b(r, t) p_L(t)$ and the fact that if $Q_A^t(0)\rho_0(0)$ is zero along with $Q_A^t(0)l_0\alpha(\hat{r})p_L(t) = 0$ since $\phi_b(r, t) = -\beta$. We have

$$\rho_b(t) = \rho_L(t) + U_{Q_A}^t(0, t) Q_A^t(0)\rho_0(0) - \int_0^t dt_1 \left( U_{Q_A}^t(t_1) Q_A^t(t_1)iL_0p_L(t_1) \right) \tag{C18}$$

where the use of the identity $Q_A^t(0)iL_0p_L(t) = (Q_A(t_1)iL_0p_L(t_1)$ for some function $f$ has made sense.

The fluxes in Eq. (C18) are $\tilde{N}_f(r)$ in Eq. (19) and

$$\tilde{g}_{A}(r) = -\nabla \cdot \hat{r} \cdot \gamma + F_i(r), \tag{C19}$$

where the fluid stress tensor is

$$\tau(r) = \sum_{i=1}^N \left[ \frac{P_i - \frac{1}{2} \sum_{j=1}^N \phi_{ij} \phi_{ij} r_i \nabla \cdot V_{ij}}{m} + \phi_{ik}^{(r)} \phi_{jk}^{(r)} \sum_{k=1}^m \nabla \cdot V_{ij} (r_i^{(k)} - r_j^{(k)}) \right] \delta(r - r). \tag{C20}$$

in the small gradient approximation, and the local force on the fluid is

$$F_i(r) = -\sum_{i=1}^N \sum_{k=1}^{m_i} \phi_{ik}^{(r)} \nabla \cdot V_{il}(r^p) \tag{C21}$$

We may then write Eq. (C18) for an isothermal system as

$$\rho_0(t) - \rho_0(t) + U_{Q_A}^t(0, t) Q_A^t(0)\rho_0(0) - \beta \int_0^t dt_1 U_{Q_A}^t(t_1) (Q_A(t_1)\rho_0(0) \star \mu_r(t, t_1) + Q_A(t) \gamma \rho_r(t) \nabla \cdot \rho_r(t) + Q_A(t) \gamma \rho_r(t) \nabla \cdot \rho_r(t_1) \star \mu_r(t, t_1) + Q_A(t) \gamma \rho_r(t) \nabla \cdot \rho_r(t_1) \star \mu_r(t, t_1) \tag{C22}$$

which is the relation we sought. The initial condition term will decay on a molecular time scale in view of the projected evolution. In addition, if the initial condition is $\rho_0(t) = \rho_0(0)$, this term is identical zero. Thus, we can neglect it in the computation. In addition, since $f^\beta(r) \star f^\beta_0(r) = -f^\beta_0(r)$, we can express the reactive contribution in terms of the chemical affinity $A(r, t) = \mu_A(r, t) - \mu_0(r, t)$ as

$$Q_A(t_1)\rho_0(t) \star \mu_r(t, t_1) = Q_A(t_1)\rho_0(t) \star A(r, t). \tag{C23}$$

**DATA AVAILABILITY**

The data that support the findings of this study are available from the corresponding author upon reasonable request.

**REFERENCES**

1. S. Ramaswamy, Annu. Rev. Condens. Matter Phys. 1, 323 (2010).
2. T. Vicsek and A. Zafeiris, Phys. Rep. 517, 71 (2012).
3. S. Aronson, C. R. Phys. 14, 518 (2013).
4. T. Elgeti, R. W. G. Gompper, J. Reprogram Phys. 73, 336601 (2015).
5. E. Fodor and M. C. Marchetti, Physica A 304, 106 (2018).
6. A. Zottl and H. Stark, J. Phys.: Condens. Matter 28, 235001 (2016).
7. S. Dukhin and B. V. Derjaguin, in *Surface and Colloid Science*, edited by E. Matijevic (Wiley, New York, 1974), Chap. 3, Vol. 7.
8. J. L. Anderson, Ann. N.Y. Acad. Sci. 469, 166 (1986).
9. J. L. Anderson, Annu. Rev. Fluid Mech. 31, 61 (1989).
10. J. L. Anderson, M. E. Lowell, and D. C. Prieve, J. Fluid Mech. 117, 107 (1982).
11. R. Golestanian, T. B. Liverpool, and A. Ajdari, Phys. Rev. Lett. 94, 220801 (2010).
12. R. Kapral, J. Chem. Phys. 138, 020901 (2013).
13. P. H. Colberg, S. Y. Reigh, B. Robertson, and R. Kapral, Acc. Chem. Res. 47, 3504 (2014).
14. C. Bechinger, R. D. Leonardo, H. Lowen, C. Reichhardt, G. Völpe, and G. Volpe, Rev. Mod. Phys. 88, 045006 (2016).
15. G. Oshanin, M. N. Popescu, and S. Dietrich, J. Phys. A: Math. Theor. 50, 134001 (2017).
16. H. Stark, Acc. Chem. Res. 51, 2681 (2019).
17. J. Wang, *Nanomachines: Fundamentals and Applications* (Wiley VCH, Weinheim, Germany, 2013).
18. W. Wang, W. Duan, S. Ahmed, T. E. Mallouk, and A. Sen, Nano Today 8, 531 (2013).
19. S. Sánchez, L. Soler, and J. Katuri, Angew. Chem., Int. Ed. 54, 1414 (2015).
20. F. Wong, K. K. Dey, and A. Sen, Annu. Rev. Mater. Res. 46, 407 (2016).
21. M. Alarcón-Correa, D. Walker, T. Qiu, and P. Fischer, Eur. Phys. J.: Spec. Top. 225, 2241 (2016).
22. P. Gaspar and R. Kapral, Adv. Phys. X 4, 1620480 (2019).
23. P. Gaspar and R. Kapral, J. Chem. Phys. 148, 134104 (2018).
24. B. C. Lee, M. Alarcón-Correa, C. Miksch, K. Hahn, J. G. Gibbs, and P. Fischer, Nano Lett. 14, 2407 (2014).
25. L. C. K. A. Abdelmohsen, F. Peng, Y. Tu, and D. A. Wilson, J. Mater. Chem. B 2, 2395 (2014).
While a rigid colloidal particle is considered here, it is straightforward to replace it by a molecule or molecular aggregate with internal degrees of freedom.

For example, an active colloid partially coated with maleate isomerase will catalyze the cis-trans isomerization from maleate to fumarate. The dihedral angle expressed in Cartesian coordinates can serve as a scalar reaction coordinate to characterize these metastable species.

Although the domain of validity of these equations is more restricted than that of the generalized Langenin equation (37) because of the assumption \( M \gg m \), even for colloids with linear dimensions of a few nanometers, there can be a large mass disparity between the colloid and solvent particles.