Direct measurement of thermophoretic forces

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We study the thermophoretic motion of a micron sized single colloidal particle in front of a flat wall by evanescent light scattering. To quantify thermophoretic effects we analyse the nonequilibrium steady state (NESS) of the particle in a constant temperature gradient perpendicular to the confining walls. We propose to determine thermophoretic forces from a ‘generalized potential’ associated with the probability distribution of the particle position in the NESS. Experimentally we demonstrate, how this spatial probability distribution is measured and how thermophoretic forces can be extracted with 10 fN resolution. By varying temperature gradient and ambient temperature, the temperature dependence of the Soret coefficient $S_r(T)$ is determined for $r = 2.5 \mu m$ polystyrene and $r = 1.35 \mu m$ melamine particles. The functional form of $S_r(T)$ is in good agreement with findings for smaller colloids. In addition, we measure and discuss hydrodynamic effects in the confined geometry. The theoretical and experimental technique proposed here extends thermophoresis measurements to so far inaccessible particle sizes and particle solvent combinations.

1 Introduction

When colloidal particles dispersed in a liquid are exposed to a temperature gradient, they are subjected to thermophoretic forces which drives them towards one side of the gradient. Which side is favored depends on the ambient temperature and the details of particle solvent interactions.1,3 Thermophoresis has been employed for instance in thermal field flow fractioning to separate colloidal particles6 or in microscale thermophoresis to study protein interactions.4 Furthermore, in hydrothermal pore model systems, a combination of thermophoresis and convection led to an extreme accumulation of nucleotides, RNA and DNA and it is likely that this mechanism played a key role in the evolutionary building up of more complex structures.5

To quantify thermophoresis of colloids, usually the stationary distribution of a particle ensemble governed by the interplay between thermophoresis in a temperature gradient and Brownian diffusion in a concentration gradient is analyzed by different optical techniques reviewed e.g. in ref. 6. This way the Soret coefficients of polystyrene particles of up to $r = 1 \mu m$ radius have been characterized.7–9 For larger particles unfeasible long equilibration times and sedimentation effects restrain these ensemble based methods. Even for polystyrene, where sedimentation can be minimized by matching the density of particles and solvent, micron sized particles are notoriously difficult to measure.6

Here we propose a new strategy to characterize thermophoresis of larger ($r > 1 \mu m$) particles based on a single particle trajectory measurement. The probe particle is held by optical tweezers in a slit geometry over which a stable temperature gradient is applied. By adapting the strength of these optical tweezers, gravity or buoyancy effects can be compensated for rendering our method essentially independent of particle weight. This overcomes sedimentation problems and enables measurements with particles of larger sizes or composed of higher density materials like PMMA or melamine.

2 Experimental section

As experimental method we use total internal reflection microscopy (TIRM, see ref. 10 and 11 for a review). It is a single particle evanescent light scattering technique capable of measuring the trajectory of a spherical colloidal particle performing its Brownian motion in vicinity of a flat wall. In equilibrium TIRM is well established as a sensitive technique to measure double layer interactions, van der Waals forces and other particle wall interactions. The new idea to characterize thermophoresis is to apply TIRM in a non-equilibrium steady state (NESS) given by a constant temperature gradient and to develop a suitable scheme for data analysis in a NESS.

As sketched in Fig. 1a, an evanescent field decaying into the solvent is created by total internal reflection of a laser beam ($\lambda = 658 \text{ nm}$) at a substrate solvent interface. A single colloidal particle near the substrate will scatter light from this evanescent field. The scattered light intensity is then monitored through microscope optics ($50 \times$, NA = 0.55 objective) by a
coating serves as an (≈80%) optical absorber for green laser light (\(\lambda = 532\) nm) while the TIRM detection wavelength (\(\lambda = 658\) nm) and larger wavelength of optical tweezers are >95% transmitted. Thus the upper part of the cell above the probe particle can be heated by a slightly focused laser beam (≈ 200 \(\mu\)m beam waist) and the temperature at the absorber is tunable by laser power.

In each sample cell the temperature increase due to laser heating is estimated by the onset of water evaporation i.e. bubble formation at high laser powers (≈ 1 W). During this procedure care was taken not to superheat the water by using probe or spacer particles as nucleation sites. The procedure was crosschecked by determining the demixing temperature (34.0 °C) of a critical water 2,6-lutidine mixture in an identically build cell, where due to the spinodal demixing process superheating is impossible.13 Good agreement between both temperature estimates was found. Depending on the actual cell in use, heating of 0.10 to 0.15 K mW⁻¹ at the upper cover glass is typical for the experiments.

For further analysis, heat distribution within the cell was modelled and computed by finite elements methods (COMSOL multiphysics using heat transfer model) taking into account the heat conductance and capacity of prism, substrate, water layer, particle, coverglass and 1 mm air above the cell. The copper housing of the cell was modelled as constant temperature boundary condition while for laser heating a Gaussian heating power profile within the SiO layer was assumed. The numerical results show that the temperature drop within the cell is not uniform due to the heat conductivity of the particle \(\kappa_p\) being different from that of the solvent \(\kappa_s\). Within the confined geometry, we determine the temperature gradient over the probe particle from the temperature difference between the particle’s top and bottom boundary divided by its diameter. For the case of a \(r = 2.5\) μm polystyrene bead (\(\kappa_p = 0.13\) W K⁻¹ m⁻¹) in water (\(\kappa_s = 0.6\) W K⁻¹ m⁻¹), as studied in Section 5, we find that the temperature gradient within the particle is a factor 1.34 steeper than in an identical cell without particle. This factor is very similar to the renormalization factor \(3\kappa_s/(2\kappa_s + \kappa_p)\) established for the temperature gradient in bulk systems (cf. eqn (24) of ref. 2). Also for melamine particles (\(\kappa_p = 0.45\) W K⁻¹ m⁻¹), our numerical COMSOL simulations of the cell reveal a renormalization factor of 1.09, very close to the bulk value \(3\kappa_s/(2\kappa_s + \kappa_p)\).

Furthermore, within the parameter range of our experiments, we find only a weak dependence (≤5%) of this renormalization factor on particle size and \(z\)-position, indicating that the temperature distribution over the particle surface in our cell geometry is very similar to the one under bulk conditions. In the data analysis, we therefore neglect that weak size and position dependency of the temperature field. For determining the temperature difference over the sample cell, it has to be considered that the heat conduction of the sapphire substrate is not infinite. As a consequence, the temperature of the substrate at the sapphire–water interface \(T_S\) is not independent of laser heating and temperature gradient. According to our simulations, the change of \(T_S\) is, depending on details of cell design, about 1/3 of the total temperature rise. This is taken into

For thermophoresis measurements, a stable temperature gradient has to be established in the sample cell. This is accomplished by a thin cell design (Fig. 1b) that allows considerable temperature gradients at only moderate temperature changes. It consists of a 1 mm thick sapphire substrate to ensure good thermal coupling to a heat sink, colloidal silica particles of 7.7 μm diameter as spacers to define the cell height \(L_s\), and a 1 mm thick cover glass coated with about 1.8 μm SiO layer on the inner side of the cell. The thermally evaporated SiO

![Sketch of experimental set up](image-url)
3 Model and theory

In this section we outline the standard phenomenological description of thermophoresis and adapt it to the confined geometry and particle wall interactions relevant for the present experiment. In particular we demonstrate how to extract thermophoretic forces and the Soret coefficient \( S_T \) from single particle measurements.

We describe the probability density \( p = p(x, y, z, t) \) for finding a particle at position \((x, y, z)\) in the sample cell at time \( t \) by the Fokker–Planck equation \( \frac{\partial p}{\partial t} + \nabla J = 0 \), with the probability current \( J = (J_x, J_y, J_z) \). In the present experimental setup, \( J \) contains contributions from deterministic forces (particle–wall interactions and external forcing by optical tweezers and gravity), thermophoretic drift and thermal noise effects. We define the coordinate system in such a way, that the \( z \)-axis is oriented along the temperature gradient (i.e. perpendicular to the sapphire substrate) and its origin is located at the sapphire–water interface (see Fig. 1b). Thus the two walls (sapphire substrate and cover glass) with their different temperatures are located at \( z = 0 \) and \( z = L \) and the \( x \)-\( y \) plane is parallel to the substrate. Since the optical tweezers confine the particle motion in the \( x \) and \( y \) directions, and we are only interested in the motion perpendicular to the wall (\( z \) direction), we can integrate out the \( x \) and \( y \) components in the Fokker–Planck equation to obtain

\[
\frac{\partial p}{\partial t} + \frac{\partial J_z}{\partial z} = 0, \tag{1}
\]

where now \( p = p(z, t) \) and \( \frac{\partial J_z}{\partial z}|_{z=0} = \frac{\partial J_z}{\partial z}|_{z=L} = 0 \) (reflecting boundary conditions).

The probability current \( J_z \) along the temperature gradient can be written as

\[
J_z = \frac{f z}{\gamma} - p D_T \frac{\partial T}{\partial z} - D \frac{\partial p}{\partial z}. \tag{2}
\]

The force \( f \) in the first term contains all particle wall interactions, light forces from the optical tweezers and gravity; \( \gamma \) is the viscous friction coefficient. The second term is the standard phenomenological ansatz for the thermophoretic drift being proportional to the temperature gradient with \( D_T \) as a phenomenological coefficient quantifying the thermophoretic effects.\(^\text{2,6,14}\) The last term represents thermal diffusion with Einstein’s diffusion constant \( D = k_B T / \gamma \). The details connected with the specific form of the thermal noise term with the space-dependent diffusion coefficient being written in front of the gradient \( \partial \partial z \) are discussed in the Appendix.

In the following, we are interested in the non-equilibrium steady state (NESS) where thermophoretic effects are balanced by external forces and diffusion currents so that \( J_z = 0 \). The current-free solution of eqn (2) reads

\[
p \text{NESS}(z) = \frac{1}{\lambda} \exp \left[ \int_0^z \left( \frac{f z}{k_B T} - \frac{D_T}{D} \frac{\partial T}{\partial z} \right) \, dz' \right], \tag{3}
\]

where we need to keep in mind that \( f, T, D \) and, in general, also \( D_T \) depend on position \( z \). The normalization constant \( \lambda \) is chosen such that \( \int_0^L p(z) \, dz = 1 \).

Following the procedure in ref. 15, we define the generalized “pseudo-potential”\(^\text{15,16}\) as the negative logarithm of the steady-state distribution,

\[
\Phi(z) := -\ln p \text{NESS}(z) = -\int_0^z \left( \frac{f z}{k_B T} - \frac{D_T}{D} \frac{\partial T}{\partial z} \right) \, dz' + \Phi_0. \tag{4}
\]

The offset \( \Phi_0 = \ln \lambda \) is due to normalization; it is irrelevant for the spatial dependence and thus will be omitted in the following. The generalized potential is composed of two space-dependent contributions

\[
\Phi(z) = \Phi_f(z) + \Phi_T(z). \tag{5}
\]

The first part

\[
\Phi_f(z) = -\int_0^z \frac{f z}{k_B T} \, dz' \tag{6}
\]

is related to the conservative forces \( f \) on the particle. It simplifies considerably for small temperature gradients. Writing

\[
T(z) = T_s + \theta z \tag{7}
\]
with the substrate temperature $T_s = T(z = 0)$ and temperature gradient $\theta = \partial T/\partial z \approx (T(z = L) - T_s)/L$, and assuming that $|T(z = L) - T_s|/T_s \ll 1$, the term $1/k_BT_s$ to lowest order in $\theta L/T_s$. Thus for small temperature gradients $\theta L/T_s \ll 1$, the generalized potential $\Phi_T$ is directly determined by the equilibrium potential $V(z) = -\int_0^z f(z') \dd z'$, 

$$\Phi_T(z) = V(z) \frac{1}{k_BT_s}. \quad (8)$$

This relation is exact in case of an equilibrium situation, i.e. where there are no temperature variations over the sample cell, $\theta = 0$.

The second part in eqn (4),

$$\Phi_T(z) = \int_0^z \frac{f(z)}{D} \frac{\partial T}{\partial z'} \dd z', \quad (9)$$

can be interpreted as the generalized potential of a “thermophoretic force”

$$f_z = -k_BT \frac{\partial \Phi_T}{\partial z} = -k_BT \frac{D_T}{D} \frac{\partial T}{\partial z}. \quad (10)$$

A simplification based on a small gradient expansion as we performed it for $\Phi_T$ is not easily possible due to the a priori unknown temperature dependence of the coefficient $D_T/D$.

It is obvious, however, that in thermal equilibrium with $\partial T/\partial z = \theta = 0$ we have $f_z = 0$ and thus $\Phi_T = 0$ (again, up to an irrelevant constant). Using eqn (5) and (8), we find $\Phi = V(z)/k_BT_s$ and, finally, from eqn (4)

$$V(z) = -k_BT \ln p_{eq}(z), \quad (11)$$

when substituting the NESS density $p_{NESS}$ by its equilibrium counterpart $p_{eq}$ as the stationary distribution reached under thermal equilibrium conditions. Because the equilibrium density $p_{eq}$ is given by the Boltzmann factor $\exp(-V/k_BT_s)$, this confirms that the general non-equilibrium approach eqn (4) is consistent with equilibrium statistical mechanics.\(^{15}\)

## 4 Data evaluation

As already mentioned in Section 2, the probability density $p(z)$ for finding the particle at distance $z$ from the substrate can be extracted from the TIRM scattering intensities. Performing such a TIRM measurement without applying a temperature gradient, allows to deduce the potential $V(z)$ via eqn (11) to quantify particle–wall interactions and the external forces due to the optical tweezers and gravity.

In our experiments, the temperature gradient $\theta$ is below 1 K $\mu$m$^{-1}$ so that $\theta L/T_s \leq 0.025$ (with $L = 7.7 \mu$m and $T_s = 300$ K) is indeed negligibly small. As shown above [see eqn (8)], such small temperature variations do not interfere with the potential forces due to $V(z)$. We can therefore determine the “thermophoretic potential” $\Phi_T$ from the stationary distribution $p_{NESS}$ in a given (weak) temperature gradient by using [cf. eqn (4), (5) and (8)]

$$\Phi_T(z) = \Phi(z) - \Phi(z) = -\ln p_{NESS}(z) - \frac{V(z)}{k_BT_s}. \quad (12)$$

A central quantity for characterizing thermophoretic effects is the Soret coefficient $S_T = D_T/D$. According to eqn (10) it is related to the “thermophoretic force” and the generalized “thermophoretic potential” by

$$S_T = -\frac{f_z}{k_BT \partial T/\partial z} = -\frac{\partial \Phi_T/\partial z}{\partial T/\partial z}. \quad (13)$$

The Soret coefficient can therefore be measured from the stationary particle distribution $p_{NESS}$ observed in a weak temperature gradient $\theta$ (with $\theta L/T_s \ll 1$) by making use of the relations (12), (13), after $V(z)$ has been determined in an independent equilibrium measurement [eqn (11)]. We remark that Parola and Piazza in ref. 14 suggested a relation analogous to eqn (13) for determining the Soret coefficient, however derived directly from hydrodynamic forces on the particle due to thermophoretic effects and without taking into account other conservative forces.

## 5 Results and discussion

Typical TIRM measurements for different temperature gradients are shown in Fig. 2a. Without temperature gradient, the equilibrium potential $V(z)/k_BT_s = \Phi(z)$ (black squares) is obtained. For $z < 30$ nm it exhibits a steep repulsion which is due to screened Coulomb interactions between the negatively charged surfaces of sapphire substrate and sulfate terminated polystyrene particle. For larger distances up to about 150 nm a potential well of 0.8 $k_BT$ depth is observed. This can be attributed to attractive van der Waals forces. Towards larger distances the potential has a constant slope that reflects the buoyancy of the particle and additional light forces of optical tweezers. The entire potential is well fitted by the function $\Phi_{th}(z) = 16.50 \exp(-z/10 \text{ nm}) - 6.53 \exp(-z/21.91 \text{ nm}) + 3.05 \text{ nm}^{-1}z + 0.35$ displayed as orange line in Fig. 2a.

The first term accounts for electrostatic interactions according to Debye–Hückel theory. The fitted Debye length of 10 nm is attributed to the counterions in the thin sample cell. The second term describes the van der Waals attraction by an empirical exponential formula given in eqn (5) of ref. 17. The last two terms incorporate light pressure and gravity. In the following $\Phi_{th}(z)$ is used to subtract $\Phi_T(z)$ from $\Phi(z)$ in the presence of temperature gradients [cf. eqn (12)]. The result is shown in Fig. 2b for $z \geq 30$ nm, i.e. the position of the potential minimum. The pure thermophoretic potentials $\Phi_T$ obtained at different temperature gradients are, within errors, linear functions of distance. This means that the probe particle experiences a constant thermophoretic force within the distance range sampled. For bulk measurements this is certainly expected. However, in the vicinity of a surface it is worthwhile to discuss the different contributions to $f_z$ (eqn (10)), in particular due to the hydrodynamic wall effects altering the diffusion coefficient $D(z)$.
Close to a wall the hydrodynamic friction coefficient $\gamma$ of a spherical colloidal particle is changed drastically (it even becomes anisotropic) compared to the bulk value $\gamma_0 = 6\pi\eta r$ ($\eta$ being the fluid viscosity) given by Stokes’ solution. By Einstein’s relation $D(z) = k_BT(z)/\gamma(z)$ also the Brownian diffusion coefficient for diffusion perpendicular to the wall $D(z)$ acquires a pronounced distance dependence. Exploiting, as before, that the temperature variations over the sample cell are small in our experiments, $\beta L/T_0 \ll 1$, we can neglect the space-dependence of temperature and find in lowest order $D(z) = k_BT_0/\gamma(z)$. The effects of hydrodynamic corrections in the friction and (normal) diffusion coefficient close to the walls can therefore be considered to be unaffected by the (weak) temperature gradients. The theoretical prediction for the normal diffusion coefficient is displayed in Fig. 3 as orange line. It has been calculated by Brenner’s formula and its extensions for a thin slit between two walls, which represents the actual experimental situation. The inset predicts that in the experimental slit geometry diffusion is at maximum only 23% of the bulk diffusion coefficient $D_0 = k_BT_0/(6\pi\eta r)$.

Experimentally, the spatially resolved diffusion coefficient can be extracted from dynamical analysis of TIRM-data according to a procedure described in ref. 21. It is shown as symbols in Fig. 3. For all temperature gradients data are in remarkable agreement. This proves that the Brownian motion is completely independent of thermophoretic effects. Since $\partial\Phi_T/\partial z$ is constant within the experimentally accessible distance range, it follows from eqn (10) that $D_T(z)$ has the same distance dependence as $D(z)$ and the Soret coefficient $S_T = D_T(z)/D(z)$ is constant i.e. independent of $z$.

The magnitude of the thermophoretic force, i.e. the negative slopes of the fitted (black) lines in Fig. 2b, also depends linearly on the temperature gradient, as shown in the inset of Fig. 2b. This experimental observation confirms the usual assumption that thermophoretic velocities (and forces) are proportional to the temperature gradient and $S_T$ is independent of the temperature gradient. It also imparts an impression on the magnitude of thermophoretic forces which are here in the range of about 20–100 fN.

While $S_T$ does not depend on temperature gradient, a characteristic of thermophoresis is its pronounced dependence on the absolute temperature. It has been shown that for many substantially different dispersed systems, like polystyrene nanoparticles, Lysozyme micelles, DNA etc., the temperature dependence of the Soret coefficient follows a common empirical fitting formula introduced by Iacopini and Piazza,

$$S_T(T) = S_T^0 \left[1 - \exp\left(\frac{T^*-T}{T_0}\right)\right]$$

(14)

with system specific constants $S_T^0$, $T^*$ and $T_0$.

Fig. 4 shows the temperature dependence of $S_T$ for probe particles of different material. As an example of particles that due to their size and strong sedimentation (density $\rho_{MF} = 1.51 \text{ g cm}^{-3}$) are difficult (if not impossible) to access with other
techniques, we present data for $r = 1.35 \mu m$ melamine particles (triangles) with carbonyl terminated surface. Qualitatively, they follow the temperature dependence given by eqn (14) with a remarkably high $T^* = 43.5 \degree C$. Hence at room temperature they have a negative $S_T$ and belong to the rare group of thermophilic particles.

$S_T(T)$ for $r = 2.5 \mu m$ polystyrene particles also accords with eqn (14). Even the temperature of sign reversal is in good agreement with previous results for considerably smaller polystyrene particles of 53 nm to 253 nm radius where $T^* = 12 \degree C \pm 285.2 K$ is reported. Comparing the absolute values of $S_T$ for the $r = 253 \text{ nm}$ particles in these measurements and for the 10 times larger polystyrene particles in the present experiment, we find in our experiments that $S_T$ is two orders of magnitude larger. This certainly exceeds the linear prediction for the size dependence of $S_T$ found for smaller particles measured in ref. 9 and expected from theories of thermophoresis based on a flow field around the particle. While in our experiments no salts or surfactants where added, dispersions of ref. 9 were treated with Triton X100 surfactant and experiments where performed in a density matched H$_2$O : D$_2$O mixture containing 1 mM Tris-HCl pH 7.8 buffer and 10 mM NaCl. These different condition certainly question the comparability of both measurements even though $T^*$ known to depend sensitively on the particle-solvent interface is quite similar.

In experiments with $r = 1.25 \mu m$ polystyrene particles (data not shown) measured under same conditions as the $r = 2.5 \mu m$ particles, we find a 4–5 times smaller $S_T$ which corresponds to a nonlinear scaling of $S_T$ with particle radius. This contrasts with the majority of experiments reported in the literature (see ref. 6 for a review), but a quadratic size dependence for $S_T$ is found in ref. 7 and 8. Such a quadratic dependence is theoretically supported by thermodynamic arguments in a parameter range where $S_T r^2 \ll 1$. However, in our experiments the combination of $S_T$, particle size and temperature gradients $S_T r^2 \approx 5…100$ largely exceeds the applicability of this theory.

Finally, if we presume a driving mechanism for thermophoretic motion that creates a flow field around the particle and in bulk results in a linear size dependence of $S_T$, it is to be expected, that the pronounced influence of confining surfaces on hydrodynamics shown in Fig. 3 will not only influence the Brownian diffusion but also the thermophoretic propulsion itself. On a length-scale of the order of the particle radius, i.e. for distances considerably larger than those sampled in Fig. 2, this would lead to a spatial variation of thermophoretic forces. If this is indeed the case, comparison with bulk measurements and furthermore the concept of $S_T$ (which should be a particle property, independent of distance from boundaries) is inappropriate in confined geometries. A distance dependence of thermophoretic forces could also explain measurements showing a size dependence of $S_T$ which is stronger than linear within this hydrodynamic concept. These issues will be in the focus of our further research.

6 Conclusions

In conclusion we have demonstrated the applicability of TIRM in NESS-systems and pointed out a suitable scheme for data analysis. The origin of nonequilibrium is a temperature gradient that drives a thermophoretic motion of micron sized colloidal particles. Thermophoretic forces have been defined and directly measured with a precision in the 10 fN range. In good qualitative agreement with existing measurements for smaller polystyrene particles we measured the temperature dependence of $S_T$ for polystyrene particles. Influences of the confined geometry on $D$, $D_m$ and $S_T$ were discussed. This might have important consequences for the applicability of thermophoresis in microfluidic devices. Furthermore we demonstrated the potential of the method to characterize micron sized particles with higher densities with respect to their thermophoretic properties by determining $S_T(T)$ for melamine particles in agreement with the empirical formula by Iacopini and Piazza. The new technique not only makes a larger variety of micro-particles accessible to thermophoretic measurements, but also opens up a route for generalization to different solvents like alcohols, hydrocarbons and other unpolar liquids. This might allow to study the dependence of thermophoresis on detailed particle solvent interaction in the near future.

Appendix

In this Appendix, we discuss the details associated with formulating overdamped Brownian motion in an inhomogeneous thermal environment with space-dependent diffusion coefficient in the context of thermophoresis. For simplicity we restrict ourselves to one spatial dimension which is denoted by $z$ (as in the main text). We furthermore omit any deterministic...
forces, so that the probability current $J_z$ for the particles in a dilute suspension consists of a thermophoretic part proportional to the gradient of temperature $T = T(z)$ and a diffusive part proportional to the gradient of the probability density $p = p(z, t)$,

$$J_z = -pD_T \frac{\partial T}{\partial z} - D \frac{\partial p}{\partial z}. \quad (15)$$

This description represents the standard form used in the thermophoretic literature,$^{28,30,34}$ based on the reciprocal formulation of heat and particle currents driven by temperature and density gradients.$^{37}$ The so-called “thermal diffusion coefficient” $D_T$ essentially quantifies the effects of the thermal gradient on a thin layer at the particle–solvent interface, where interfacial tension gradients parallel to the temperature gradient drive thermophoretic particle motion.$^{28}$ The strength of the diffusion term is given by the diffusion coefficient $D = k_T/T\gamma$.

Our main concern in this Appendix is the specific form of this diffusion term. In thermophoresis, $T$ and thus $\gamma$ and $D$ are position-dependent. It is therefore not clear that $-D\partial p/\partial z$ is the “correct” form to describe particle diffusion in such inhomogeneous thermal environments or if additional “spurious” drift terms would have to be added in eqn (15).$^{28-34}$ This problem is per se not related to thermophoretic mechanisms as sketched above, but has a different origin rooted essentially in the mathematical description of Brownian motion. It is connected to the so-called Itô–Stratonovich dilemma for the stochastic Langevin equation associated with eqn (15).$^{35,38}$ In fact, as fluid viscosity changes with temperature, the friction coefficient $\gamma$ depends on position $z$. As an example, in our experimental setup, where we measure thermophoresis close to the substrate surface in a slit geometry (see Fig. 1), $\gamma$ even acquires an additional effective dependence on the wall distance $z$ by hydrodynamic effects$^{28}$ In other words, variations of $D$ with position $z$ are the result of separate dependencies of $T$ and $\gamma$ on $z$. It is well-known$^{28,29,32,34,38}$ that under such conditions, the correct form of the diffusion term is $-(1/\gamma)\partial(k_T T p)/\partial z$, so that we would have to write

$$J_z = -pD_T \frac{\partial T}{\partial z} - \frac{1}{\eta} \frac{\partial (k_T T p)}{\partial z}. \quad (16)$$

with a different thermophoretic coefficient $D_T$ as compared to eqn (15). This particular form of the diffusion term results from performing the overdamped limit in the full-fledged particle equations of motion including the velocity degrees of freedom (after the white noise limit has been performed first).

We can easily bring eqn (16) into the form eqn (15) by identifying $D_T = D_T + k_T/\gamma$. Based on an alternative physical reasoning, the additional “correction” term $k_T/\gamma$ has been discussed in ref. 40 to be related to the osmotic pressure in dilute suspensions.

The corresponding difference between the Soret coefficients $S_T = D_T/D$ and $D_T/D$ is $1/T$. Typical values of $S_T$ for colloidal beads are of the order of 0.1 K$^{-1}$ to 1 K$^{-1}$$^{[1, 2, 9]$, (or even several ten or hundred per Kelvin for micron-sized particles as measured here), so that around room temperature this difference is expected to be negligibly small, except maybe in temperature regions close to the sign change of $S_T$.

We finally remark that a third variant for the diffusion term has been discussed briefly in ref. 14, namely $-\partial(Dp)/\partial z$. In principle, this prescription amounts to yet another definition for the thermophoretic coefficient in the probability current

$$J_z = -pD_T \frac{\partial T}{\partial z} - \frac{\partial (Dp)}{\partial z}. \quad (17)$$

However, as we have pointed out above, this form of the diffusion term is not correct if $\gamma$ depends on position $z$; the difference to the correct form $-(1/\gamma)\partial(k_T T p)/\partial z$ in eqn (16) is an unphysical drift-like term $p(D/\gamma)(\partial\gamma/\partial z)$. In case the friction coefficient $\gamma$ depends on space only due to changes of fluid viscosity with temperature, this term may be hidden in $D_T$, as by comparison with eqn (16) we can write $D_T = D_T - (D/\gamma)(\partial\gamma/\partial T)$, or, comparing with eqn (15), $D_T = D_T + k_T/\gamma - (D/\gamma)(\partial\gamma/\partial T)$. For water around room temperature the corresponding “correction” $-(1/\gamma)(\partial\gamma/\partial T)$ in the Soret coefficient is of the order of 0.02 K$^{-1}$ and thus for colloidal particles probably negligibly small in most cases.

Summarizing, for modeling thermophoresis in dilute particle suspensions, one should either use eqn (15) or (16) for the probability current, while eqn (17) is physically questionable. The deviation between eqn (15) and (16) is compensated by different definitions of the “thermal diffusion coefficient” $D_T$ vs. $D_T$. They differ in the contribution $k_T/\gamma$, which is related to the temperature dependence of the osmotic pressure in dilute suspensions.$^{38}$ Given, however, that the formulation of thermophoretic effects is connected to the reciprocity of heat and mass flow in temperature and density gradients,$^{37}$ the standard representation eqn (15) is preferable.

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