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Fast and versatile thermo-osmotic flows with a pinch of salt†

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Thermo-osmotic flows – flows generated in micro and nanofluidic systems by thermal gradients – could provide an alternative approach to harvest waste heat. However, such use would require massive thermo-osmotic flows, which are up to now only predicted for special and expensive materials. Thus, there is an urgent need to design affordable nanofluidic systems displaying large thermo-osmotic coefficients. In this paper, we propose a general model for thermo-osmosis of aqueous electrolytes in charged nanofluidic channels, taking into account hydrodynamic slip, together with the different solvent and solute contributions to the thermo-osmotic response. We apply this model to a wide range of systems by studying the effects of wetting, salt type and concentration, and surface charge. We show that intense thermo-osmotic flows can be generated using slipping charged surfaces. We also predict for intermediate wettings a transition from a thermophobic to a thermophilic behavior depending on the surface charge and salt concentration. Overall, this theoretical framework opens an avenue for controlling and manipulating thermally induced flows with common charged surfaces and a pinch of salt. COMMUNICATION

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Due to the increasing world energy consumption and the need for new clean energies, waste heat harvesting is a major challenge for decades to come. Some of the most common difficulties in harvesting waste heat come from the small temperature differences between the source and the environment $(**50** °C)¹$ as well as from the need to use rare, expensive and often toxic thermoelectric materials.² Alternatively, thermo-osmotic flows (generated at liquid–solid interfaces by temperature gradients) can be used to transform waste heat into electricity via a turbine³ or to pump water for desalination.^{4,5} Thermo-osmotic flows also directly generate electric current on charged surfaces $6,7$

by advecting the charge of the electric double layer (EDL) appearing in the liquid to screen the surface charge.⁸ Historically, the first experimental results on thermo-osmosis were published by Lippmann⁹ and Aubert¹⁰ at the beginning of the 20th century. Since then, a broad literature has been devoted to the study of aqueous solutions and various membranes, both from experiments^{11–15} or molecular dynamics (MD) simulations.^{16–20} Some disagreements have been reported for aqueous electrolytes, specifically in the flow direction (toward the hot side, so-called thermophilic flow, or toward the cold side, so-called thermophobic flow) for similar systems, 2^{1-24} or on the relation between the flow amplitude and surface charge. 13 Although with exceptions, a thermophobic flow is generally expected for hydrophobic membranes and a thermophilic flow for hydrophilic membranes.^{25,26} Such differences cannot be understood by the classical theory²⁷ developed by Derjaguin and Sidorenkov $11,28$ and by Ruckenstein for thermophoresis.29 This theory, based on the electrostatic enthalpy of the EDL, predicts that the flow is controlled by the electric surface charge, and always goes to the hot side. Other authors have recently proposed a different understanding in terms of irreversible thermodynamics by taking into account solvent contribution.¹⁵ However, their description only explains qualitatively the flow direction and does not provide a microscopic description of the effects of fluid wetting properties, salt concentration and electric surface charge.

Thermo-osmosis has seen a renewed interest due to the massive thermo-osmotic responses predicted by the use of novel materials, such as soft nanochannels,³⁰ carbonnanotubes^{5,18,31} or graphene,¹⁷ together with novel experiments by Bregulla et $al.^{24}$ who first reported a microscale manifestation of thermo-osmotic flows. Thermo-osmotic flows could, in particular, be boosted by the slip boundary condition (BC) that describes the velocity jump v_s at the interface by a general expression first proposed by Navier: $32,33$

$$
\nu_{\rm s} = b \frac{\partial \nu}{\partial z} \big|_{z=z_{\rm s}},\tag{1}
$$

where z_s corresponds to the shear plane position³⁴ and b denotes the slip length. 35 The role of interfacial hydrodyn-

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Following this work, we propose in this communication an analytical framework with the objective to predict thermoosmosis of aqueous electrolytes confined by charged surfaces, extendable to thermoelectricity and thermophoresis. Solvent contribution and electrostatic ionic interactions are shown to play the leading role along with hydrodynamic slip. We apply the model to a wide range of systems by varying the wetting interaction, salt type and concentration, and surface charge. We report large thermo-osmotic responses, comparable to the highest responses predicted for special systems as inferred from previous simulations, $5,16,17,31$ as well as a change of sign in the flow direction. Such a change of sign cannot be predicted by only considering electrostatic interactions and can be crucial in order to interpret the different experimental results reported in the literature.

Theoretical framework

The thermo-osmotic response of a liquid–solid interface is quantified by the thermo-osmotic coefficient M_{to} , defined from the relation $v_{\text{to}} = M_{\text{to}}(-\nabla T/T)$, where $\nabla T/T$ is the relative temperature gradient parallel to the wall and v_{to} is the generated thermo-osmotic velocity in the bulk region (where the liquid does not interact with the wall). 27 In ref. 17, the authors propose a modification to the classical Derjaguin theory²⁸ and show that, in order to take into account the hydrodynamic BC, the thermo-osmotic response coefficient can be expressed as (see the ESI†):

$$
M_{\text{to}} = \frac{1}{\eta} \int_{z_0}^{\infty} (z - z_{\text{s}} + b) \delta h(z) \text{d}z, \tag{2}
$$

where η is the liquid viscosity, z is the distance to the surface, δh is the enthalpy excess density, b is the slip length defined from eqn (1), and z_s corresponds to the shear plane position. One can account for the presence of slip or a stagnant layer close to the wall by introducing an effective slip length b_{eff} = $b - z_s$ (see Fig. 1a). When $b_{eff} \ge 0$ (slip situation), the velocity profile does not vanish in the water slab and therefore the integral in eqn (2) should be performed from the wall position considered at zero, $z_0 = 0$. On the contrary, if $b_{\text{eff}} < 0$ (stagnant layer situation), then b_{eff} identifies with the size of a stagnant layer present at the liquid–solid interface, where the liquid velocity vanishes. In this case, the stagnant layer does not contribute to the integral in eqn (2) and consequently $z_0 = -b_{\text{eff}}$.

With regard to the enthalpy excess density δh , the standard approach^{28,29} assumes that it is mostly determined by the electrostatic enthalpy of ions in the EDL, $\delta h_{el}(z) = \rho_e(z)V(z)$ + $p(z)$, where ρ_e is the charge density, V is the local electric potential and p is the pressure. Using the Poisson equation ρ_e = $-e d_z^2 V$ (assuming a constant solvent permittivity ε) and con-

Fig. 1 (a) Schematics of the effective slip length b_{eff} as a function of the slip length b and the shear plane position z_s . We distinguish between the slip situation ($b_{\text{eff}} > 0$) and the stagnant layer situation ($b_{\text{eff}} < 0$). (b) Modelled system for the measures of water enthalpy excess density and slip, together with the normalized density profiles of water and ions (total density of cations and anions) for an uncharged surface, with z the distance to the wall. (c) Water enthalpy excess density δh_{wat} profiles, with z the distance to the wall, for different wetting angles θ , controlled by the interaction energy between the liquid and the solid atoms ε _{LS}. (d) Water contribution to the thermo-osmotic response coefficient, $M_{\rm{to}}^{\rm{wat}}$ for different wettings, determined based on water enthalpy excess and slip length computed from MD simulations; stars correspond to $M_{\rm{to}}^{\rm{wat}}$ < 0 and circles to $M_{\text{to}}^{\text{wat}} > 0$.

sidering the mechanical equilibrium along the z direction, dp $rac{\mathrm{d}p}{\mathrm{d}z} = -\rho_e \frac{\mathrm{d}V}{\mathrm{d}z}$, δh_{el} is then usually expressed in terms of the electrostatic potential as:

$$
\delta h_{\rm el}(z) = -\varepsilon V(z) \frac{\mathrm{d}^2 V}{\mathrm{d} z^2} + \frac{\varepsilon}{2} \left(\frac{\mathrm{d} V}{\mathrm{d} z} \right)^2. \tag{3}
$$

Accordingly, δhel vanishes outside the EDL, whose thickness is given by the Debye length $\lambda_{\rm D}$, controlled by the salt concentration.⁸ The corresponding contribution to the thermoosmotic response, $M_{\text{to}}^{\text{el}}$, can be then computed analytically within the mean-field Poisson–Boltzmann framework, 37 considering a semi-infinite channel (see the ESI†).

Although the model proposed by Derjaguin et al. can predict M_{to} experimental orders of magnitude under certain conditions, 24 it fails to describe the amplitude of the responses reported in the literature, $5,16,17,31$ the thermoosmotic response reported for weakly charged membranes,¹³

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and the experimental discrepancies observed in M_{to} sign.²¹⁻²⁴ Aside from the electrostatic ionic interactions, other contributions to δh can be important. Such contributions are related to the solvent (water in the present work), ion solvation, and water dipole orientation in the electric double layer. After comparing all the different contributions to M_{to} (see the ESI†), the two main ones are (in the case of symmetric salts such as NaCl or KCl):

$$
M_{\text{to}} \simeq M_{\text{to}}^{\text{wat}} + M_{\text{to}}^{\text{el}^*},\tag{4}
$$

related to the solvent enthalpy excess density δh_{wat} (Fig. 1c and d) and to a modified electrostatic term δh_{el}^* , accounting for the depletion of the ions in the vicinity of the wall (see the density profiles in Fig. 1b).

Defining the characteristic depletion length as d_{ℓ} , one can effectively account for this effect by imposing a vanishing enthalpy excess in the interfacial region where there are no ions: $\delta h_{\text{el}}^* = \delta h_{\text{el}}$ for $z > d_e$ and $\delta h_{\text{el}}^* = 0$ otherwise. With regard to the water contribution, δh_{wat} can be computed as the sum of the different atomic contributions, $\delta h_{\text{wat}} = \delta h_{\text{O}} + \delta h_{\text{H}}$, where the atomic enthalpy density for an element i is defined as:

$$
\delta h_i(z) = [\delta u_i(z) + \delta p_i(z)] n_i(z), \qquad (5)
$$

where $dA(z) = A(z) - A_{bulk}$, with A_{bulk} denoting the bulk value of the physical property A , u_i denoting the energy per atom, p_i denoting the stress per atom (a practical difficulty with measuring this term is discussed in the ESI[†]), and n_i denoting the atomic number density profile.

To compute the solvent term δh_{wat} and the hydrodynamic BC as a function of wetting, we ran MD simulations using the LAMMPS package.³⁸ The system consisted of an aqueous electrolyte (constituted by SPC/E water³⁹ and NaCl, such that the bulk salt concentration was $n_s \sim 1$ M, following ref. 40), confined between generic uncharged Lennard–Jones (LJ) walls or graphene (see Fig. 1b and details in the ESI†). From these simulations with uncharged walls, we confirmed that in the case of symmetric salts the solute enthalpy, even at large concentrations, did not affect the total enthalpy profile and thus the enthalpy excess density is controlled by the solvent for neutral surfaces (see the ESI†).

The solid wall atoms were frozen and the oxygen–solid (LS) interactions were varied between the hydrophobic and hydrophilic values given in ref. 40 for LJ walls, corresponding to contact angles on uncharged surfaces of $\theta \sim 140^{\circ}$ and $\theta \sim 50^{\circ}$, respectively (the values for θ and b_{eff} can be found in the ESI†). In Fig. 1c, one can observe the typical shape of the δh_{wat} profile for different wettings. We note that the most hydrophilic situation ($\theta \sim 50^{\circ}$) is considered a no-slip situation with $b = 0.0$ Å, corresponding to a stagnant layer ($b_{\text{eff}} < 0$). For simplicity, we also did not take into account in our model the coupling between the surface charge and $slip.⁴¹$ Using the proposed analytical framework, we explored a range of experimentally accessible values for the surface charge density Σ and the salt concentration n_s : Σ was varied between -1 and

−300 mC m⁻² and $n_s \in \{10^{-4}, 1\}$ M corresponding to a Debye length $\lambda_{\text{D}} \in \{0.3, 30\}$ nm.

The objective of this communication is to present a general simple model, and in that regard, some approximations are applied in order to explore a broad range of parameters. Nevertheless, the validity of the approximations we use is consistent with the range of parameters we explored, such as the choice of a lower boundary for λ_D comparable to the size of water's first absorption layer (where water solvent properties should be accounted for in the calculations, and solvation and water properties should not be considered separately) and the upper boundary for Σ , under which the mean-field Poisson–Boltzmann description should remain valid³⁷ (see the ESI†).

Results and discussion

From eqn (4), we expect that M_{to} is controlled by the competition between water and electrostatic contributions, depending on wetting, Σ and n_s (or analogously λ_D). In Fig. 2, the total thermo-osmotic response is represented for all the wettings considered, together with the water contribution $M_{\text{to}}^{\text{wat}}$ (which, by construction, is independent of λ_D and Σ) and the modified electrostatic contribution $M_{\text{to}}^{\text{el}*}$ (which is only weakly affected by wetting, mostly through the change in b_{eff}). We observed from the figure the rich behavior resulting from that competition, where the water term mostly dominates for the most hydrophobic surfaces ($\theta \ge 110^{\circ}$), while for the less hydrophobic surfaces ($\theta \lesssim 110^{\circ}$), the electrostatic contribution can dominate for the larger λ_D . We can also see a large variation in M_{to} values for different wettings, ranging from 10⁻⁹ to 10⁻⁷ m² s⁻¹ for the most hydrophobic case. Communication

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A striking result from Fig. 2 is the transition for intermediate wettings from a thermophobic behavior $(M_{\text{to}} > 0)$ at high salt concentrations (small λ_D) to a thermophilic behavior (M_{to} < 0) at low salt concentrations; see for instance, θ ∼ 110°. In agreement with previous predictions, 24 the electrostatic contribution $M_{\text{to}}^{\text{el}*} < 0$ yields a thermophilic behavior independent of the sign of the surface charge. In contrast, the water term exhibits a change of sign when varying the wetting (see Fig. 1d and the ESI†). Such change of sign of $M_{\text{to}}^{\text{wat}}$ happens at $\theta \sim 110^{\circ}$ and thus, for $\theta \gtrsim 110^{\circ}$, sgn $(M_{\text{to}}^{\text{wat}}) = -\text{sgn}(M_{\text{to}}^{\text{el}*})$ resulting in a change of sign of M_{to} when λ_{D} is such that $|M_{\text{to}}^{\text{out}}| = |M_{\text{to}}^{\text{el}*}|$. Although this change of behavior happens for all $\theta \gtrsim 110^{\circ}$, for the most hydrophobic cases it takes place for λ_D values higher than the ones considered in this study and corresponding to extremely low salt concentrations. Even so, within our parameter range, we can still observe for $\theta \sim 130^{\circ}$ a decrease of the total response for high $\lambda_{\rm D}$, which goes against the standard expectation and can only happen if water and electrostatic contributions have opposite signs. In contrast, for the less hydrophobic cases (*e.g.* $\theta \sim 90^{\circ}$), $sgn(M_{to}^{wat}) = sgn(M_{to}^{e l*})$ and M_{to} does not change sign for any λ_D value.

It is interesting to note that a similar change of sign has been found in the context of thermophoresis experiments. $42-44$

Fig. 2 Thermo-osmotic response coefficient M_{to} (solid lines) as a function of the Debye length for different wettings and surface charges. In all the graphs, the two main contributions, water M $_{\rm{to}}^{\rm{wat}}$ (dash-dotted lines) and modified electrostatic M $_{\rm{to}}^{\rm{el}*}$ (dashed lines), are also represented. While M $_{\rm{to}}^{\rm{el}*}$ is always negative, the sign of M $_{\rm{to}}^{\rm{w}}$ depends on wetting such that the total response can be thermophilic or thermophobic depending on wetting, surface charge and Debye length.

This change of sign is commonly attributed to the so-called thermopotential ψ_0 .⁴⁵ Such a thermopotential appears when cations and anions have different mobilities and when the channel boundary conditions impose a vanishing flux of each ion type in the bulk. 45 The thermopotential generates an electro-osmotic flow that can go against the thermo-osmotic flow and reverse the total flow direction. Nevertheless, ψ_0 should disappear by allowing ionic fluxes through the channel, and as a consequence, the change of sign would disappear. By introducing the water contribution to the thermoosmotic response, we propose a more fundamental understanding of such change of sign, which should persist independently of the boundary conditions on the fluxes through the channel.

The proposed $M_{\text{to}}^{\text{el}*}$ and $M_{\text{to}}^{\text{wat}}$ decomposition allows us to obtain an order of magnitude in agreement with the experimental results of M_{to}^{24} in the order of 10^{-10} – 10^{-9} m² s⁻¹. Such an agreement is especially significant for hydrophilic surfaces and in the stagnant layer situation (see the ESI†), typical of experiments due to the presence of imperfections in the solid surface, when $M_{\text{to}}^{\text{wat}}$ decreases and $M_{\text{to}}^{\text{el}*}$ may dominate for a broader range of Debye lengths. Because $M_{\text{to}}^{\text{wat}}$ increases when increasing the slip, one interesting surface is the one constituted by graphene, with an effective slip length of b_{eff} = 538.77Å, which we obtained in MD simulations for NaCl aqueous solution at room temperature (see the ESI†). From the left part of Fig. 3, one can observe a significant increase in both electrostatic and water M_{to} contributions, resulting in a large value of the total response $(M_{\rm to} \sim 10^{-6} \; \rm m^2 \; \rm s^{-1})$ for this interface. Because $M_{\text{to}}^{\text{el}}$ does not vary significantly with wetting and the order of magnitude of the total response is given by the water contribution, one can expect $M_{\text{to}} \sim M_{\text{to}}^{\text{wat}}$ for graphene. In the right part of Fig. 3, one can see how $M_{\text{to}}^{\text{wat}}$ is affected by the effective slip. From this figure, one can observe that a large M_{to} value may be obtained for very slipping systems (as CNT, where slip values of $b \sim 300$ nm have been reported at room temperature for a tube radius of $R \sim 15$ nm (ref. 46)), although it is important to note that the presence of a stagnant layer or defects in the surface (resulting in smaller

Fig. 3 Thermo-osmotic response coefficient for graphene walls. Left: total response as a function of the Debye length for different surface charges as in Fig. 2. Right: water contribution $M_{\rm to}^{\rm wat}$ as a function of the effective slip length. The dash-dotted line indicates the b_{eff} value obtained from MD simulations. For graphene walls, $M_{\text{to}}^{\text{wat}}$ and M_{to} are always negative, corresponding to a thermophilic flow.

 b_{eff}) may decrease the large predicted thermo-osmotic response, down to 10^{-9} m² s⁻¹.

Conclusions

We proposed here an analytical framework aimed at predicting the thermo-osmotic response of aqueous electrolytes for a wide range of nanofluidic systems and experimental conditions. While the standard picture relates the response to the ion electrostatic enthalpy in the electrical double layer close to charged walls, we show firstly that this contribution to the interfacial enthalpy may be negligible when compared to the water contribution for a broad range of parameters, and secondly that it should be slightly lowered due to the depletion of ions from the solid surface.

The competition between the modified electrostatic and water contributions and the impact of the hydrodynamic

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boundary condition leads to a rich phenomenology that we have illustrated here. First, our theory predicts a higher thermo-osmotic response at low λ_D than the one expected from only considering the electrostatic contribution. Second, we predict a thermophobic flow for hydrophobic systems and a thermophilic flow for hydrophilic systems, in agreement with the typical tendencies observed in the experiments.^{15,25,26} Third, for intermediate wettings, the proposed model also predicts a transition between a thermophobic behavior at low salt concentrations to a thermophilic behavior at high salt concentrations. Such a transition has also been observed in thermophoresis experiments $42-44$ and is commonly attributed to the existence of a thermopotential which is, however, limited to particular boundary conditions imposing no ionic fluxes in the bulk liquid. In contrast, our interpretation of the change of sign is more general and independent of the nanofluidic channel boundary conditions, and opens the way to manipulate thermally induced nanoscale flows with a pinch of salt. Finally, we predict intense thermally induced flows for slipping systems, with orders of magnitude comparable to the ones reported from MD simulations of water thermo-osmosis in CNTs^{5,31} or on uncharged planar walls.^{16,17} Such large flows require a significant interfacial enthalpy excess, which we have shown can be obtained for a wide range of wettings, and a large slip length, favored by hydrophobicity. While very hydrophobic materials can pose practical issues in waste heat harvesting with nanoporous membranes, large slip lengths have also been reported for mildly hydrophobic carbon-based materials.46,47 Promising hydroelectric energy conversion performance is predicted for such materials, 41 and our model suggests that thermoelectric energy conversion should also be excellent. Communication www.becomes

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The importance of solvent contribution in thermo-osmosis of aqueous electrolytes, together with a modification of the classical electrostatic term, opens the way to several perspectives. With respect to the electrostatic term, a more accurate description of thermo-osmosis should take into account spatial heterogeneities of the dielectric and viscosity profiles at the interface.⁴⁸⁻⁵⁰ In addition, when salt ions adsorb specifically to the surface, e.g. iodide for NaI, the ion-size-dependent hydrophobic solvation term should be considered, e.g. through the modified Poisson–Boltzmann framework described in ref. 40 and 51. Regarding the water term, it is left to determine the impact of the surface charge and its distribution on water contribution to the response. In addition, one could consider more realistic surfaces than the apolar walls described in this study 52 and, by following the same methodology, we propose to establish the effect of different charge distributions on the total thermo-osmotic response through their effect on the enthalpy excess and on the $slip.^{41}$ Besides, one should take into account the limits of considering pure water simulations as an approximation of the water enthalpy contribution. For high concentrations, steric effects should be accounted for and ions can affect water viscosity.⁵³ Nevertheless, such effects correspond to extreme n_s values³⁷ and they should not understate one of the main results of this communication: the great

 M_{to} value found for slipping surfaces. Finally, it is straightforward to extend the current model to the Debye-overlap regime (when the system height is smaller than λ_D), as long as the Poisson–Boltzmann framework still holds. In the same line, one can also follow the same procedure here proposed to predict the thermoelectric^{6,54,55} and thermodiffusive⁵⁶ response, with promising applications for electricity production from waste heat or to refine large-scale continuum descriptions.57 Overall, our predictions call for future experimental verification and could be exploited for the design of innovative solutions for heat harvesting applications.

Conflicts of interest

There are no conflicts to declare.

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