Diffusiophoretic and diffusioosmotic velocities for mixtures of valence-asymmetric electrolytes

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Diffusiophoresis and diffusioosmosis are electrokinetic phenomena where relative motion is induced between a charged surface and a surrounding electrolyte due to a concentration gradient of ions. In the literature, a relative velocity between a surface and the electrolyte has been derived for a valence-symmetric \( z^+/z^- \) electrolyte. In this article, we reformulate the governing equations in a convenient form based on a systematic generalization of the nonlinear Poisson-Boltzmann equations in the limit of a thin double layer, which allows us to derive results for diffusiophoretic and diffusioosmotic velocities for a mixture of electrolytes with a general combination of cation and anion valences. We find that the relative motion depends significantly on ion valences. We also provide analytical approximations for the diffusiophoretic and diffusioosmotic velocities and discuss their accuracy and applicability. Further, we tabulate diffusiphoretic velocities for some common cases, which highlights the importance of asymmetry in cation and anion valences. Finally, we discuss the validity of our assumptions and the importance of effects such as finite ion size, dielectric decrement, and surface conduction for typical experimental conditions.

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I. INTRODUCTION

Several physical scenarios involve an electrolyte in the vicinity of a charged surface, such as (1) charging and discharging of double layers, which is the underlying process in electrochemical capacitors [1,2], (2) electrophoresis and electroosmosis, which are electrokinetic phenomena where relative motion is induced between the surface and the electrolyte due to the application of an external electric field [3–5], and (3) diffusiophoresis and diffusioosmosis, which are electrokinetic phenomena where relative motion is induced between the surface and the electrolyte due to a concentration gradient of ions [5,6]. In this article, we focus on diffusiophoresis and diffusioosmosis, which are observed in processes such as dialysis [7], sedimentation and centrifugation [8], and the motion of colloidal particles in microchannels [9–16]. Moreover, reports in the literature have utilized diffusiophoresis and diffusioosmosis for applications in water treatment [10], fabric cleaning [17], self-propelling “active” swimmers [18–20], and measurement of the surface zeta potential [21], among others.

Diffusiophoresis is defined as the movement of a charged particle while the surrounding electrolyte is stationary. In contrast, diffusioosmosis occurs when the electrolyte is moving while the surface is stationary. In their classical work, Prieve, Anderson, and others derived an expression for...
the relative velocity for a valence-symmetric electrolyte, or a $z = z$ electrolyte, such as NaCl ($z = 1$) and CaSO$_4$ ($z = 2$) [5,6,22]. They showed that the diffusiophoretic (or diffusioosmotic) velocity is the sum of an electrophoretic component and a chemiphoretic component. In their derivation, a restriction is not imposed on the potential drop across the double layer or zeta potential $\psi_D$. However, their analyses require knowledge of the potential variation inside the double layer, first derived by Gouy for a $z = z$ electrolyte [23], to arrive at the expression for the diffusiophoretic velocity. Though there has been an increase in interest to derive results for a mixture of electrolytes with a general combination of cation and anion valences [24–26], a complete solution of the potential inside the double layer is not readily available for a mixture of valence-asymmetric electrolytes. Some authors report an expression for a particle’s diffusiophoretic velocity in a mixture of electrolytes where the potential distribution can be evaluated [24,27], although their results are valid only in the Debye-Hückel limit of $|\psi_D| \ll 1$, where $\psi_D$ is the dimensionless zeta potential, $e$ is the charge on an electron, $k_B$ is the Boltzmann constant, and $T$ is the temperature. However, typical experiments include $|\psi_D| \lesssim 100$ mV or $|\psi_D| \lesssim 4$ [21]. Therefore, it is necessary to go beyond the limit of $|\psi_D| \ll 1$.

Recently we demonstrated that unequal cation and anion valences influence diffuse-charge relations for an electrolyte near a charged surface [26]. Here we extend the analysis to calculate the diffusiophoretic and diffusioosmotic velocities. First, we reformulate the classical derivation of the diffusiophoretic velocity and show, perhaps surprisingly, that the detailed profile of the electric potential within the double layer is not necessary to determine the diffusiophoretic velocity. We derive velocities for a general mixture of ions with valence $z_i$, without imposing any restriction on $\psi_D$, in the limit of thin double layers. We show that the relative velocity can be significantly influenced by the valence $z_i$. Our results are in agreement with those previously reported in the Debye-Hückel limit [24,27], and we also report simplified expressions valid for large zeta potentials. For the case of a single electrolyte with the same valence, we recover the original solution highlighting the generality of our approach.

We derive our main results in the limit of thin double layers and do not consider effects such as finite ion size [26,28,29], dielectric decrement [26,30,31] and surface conduction [32–34]. Therefore, we discuss experimental conditions where our analysis is applicable and detail scenarios where the inclusion of the aforementioned effects should be considered.

II. PHYSICAL ORIGIN OF DIFFUSIOPHORESIS AND DIFFUSIOOSMOSIS

We first summarize the physical mechanism that gives rise to diffusiophoresis and diffusioosmosis. For convenience, we describe only the mechanism of diffusioosmosis. A schematic of diffusioosmosis is provided in Fig. 1 where a mixture of electrolytes is in contact with a charged surface. The valence of an $i$th ion is denoted as $z_i$. Diffusioosmotic motion of the electrolyte is produced when a gradient in ion concentration is present far away from the surface, or $c_i(x)$ (a more rigorous definition of $c_i(x)$ is provided in the next section). Depending on the charge of the surface, either cations or anions are attracted towards the surface, and the concentration of the attracted ion increases significantly near the wall. The length scale of this diffuse-charge region is given by the Debye length, $\lambda_D$. Most studies assume a single electrolyte with equal valences of cations and anions, while we focus on the general case of a mixture of multiple electrolytes with arbitrary $z_i$.

The diffusioosmotic velocity is commonly described as the sum of an electrophoretic component and a chemiphoretic component. We first focus on the electrophoretic component. Due to a gradient in the electrolyte concentration far away from the surface, ions diffuse towards the region of low concentration. However, different mobilities of cations and anions give rise to an electric field to maintain a zero flux of net charge. Since the double layer has a net charge, this electric field creates an electrostatic force inside the double layer. Therefore, the fluid far away from the surface moves so as to generate the shear forces in the double layer that balance the electrostatic forces, similar to an electrophoretic system. In contrast, chemiphoretic movement of the fluid far away from the
Fig. 1. Sketch and notation for the derivation of the diffusioosmotic velocity. The valence of the $i$th ion is denoted as $z_i$, where $z_i > 0$ for a cation and $z_i < 0$ for an anion, here illustrated in shorthand by $\oplus$ and $\ominus$, respectively. The concentration gradient of ions in the $x$ direction induces a relative motion between the electrolyte and the surface. The surface charge density is assumed to be constant and is denoted as $q$ (the schematic shows the scenario of $q > 0$).

Surface generates shear forces inside the double layer that balance the pressure forces inside the double layer, which arise due to variation in electrical force along the $x$ direction. Depending on the charge at the surface and the relative diffusivities of cations and anions, the electrophoretic and chemiphoretic components can be either in the same direction or in the opposite direction.

III. ANALYSIS OF A GENERAL MIXTURE OF ELECTROLYTES

A. Governing equations

We assume that the liquid far away from the stationary surface in Fig. 1 contains multiple ions where $z_i > 0$ for a cation and $z_i < 0$ for an anion. Here $x$ denotes the coordinate along the surface and $y$ is normal to the surface. The surface charge density $q$ is assumed to be constant and is related to the potential drop across the double layer, denoted here as $\psi_D$ (Fig. 1). The incompressible fluid flow is governed by the Stokes equations with an electrostatic body force, Gauss’s law of electrostatics, and the Nernst-Planck equations for the flux of ions. We define $u = u_x(x, y)e_x + u_y(x, y)e_y$ as the velocity field, $p(x, y)$ as pressure, $\mu$ as the fluid viscosity, $\rho_e$ as the charge density, $\psi(x, y)$ as the electrical potential, $c_i(x, y)$ as the concentrations of the $i$th ion, $j_i = j_{ix}(x, y)e_x + j_{iy}(x, y)e_y$ as the fluxes of the $i$th ion, and $D_i$ as the diffusivity of the $i$th ion. Then the governing equations are

$$ \nabla \cdot u = 0, \quad (1a) $$
$$ -\nabla p + \mu \nabla^2 u - \rho_e \nabla \psi = 0, \quad (1b) $$
$$ -\nabla \cdot (\varepsilon \nabla \psi) = \rho_e = e \sum_i z_i c_i, \quad (1c) $$
$$ \nabla \cdot j_i = 0, \quad (1d) $$
$$ j_i = -D_i \nabla c_i + u c_i - \frac{D_i z_i e c_i}{k_B T} \nabla \psi. \quad (1e) $$

In the description used above, the electrical permittivity $\varepsilon$ is assumed constant.

Our objective is to solve for the diffusioosmotic velocity $u_{DO}$ parallel to the surface, which develops at the outer boundary of the double layer. We first nondimensionalize the equations. We define a diffusivity scale $D^*$ and an ion concentration scale $c^*$, which are typical values of ion diffusivities and ion concentration in bulk. There are two relevant length scales in our system. First, there is the length at which the ion concentration changes in bulk, which we denote here as $a^*$. The
second relevant length scale is the Debye length \( \lambda_D = \sqrt{\frac{2aT}{e^2c^2}} \). We define the thermal potential as 
\[
\psi_T = \frac{k_BT}{e}, \quad \delta = \frac{2aT}{e^2c^2},
\]
and a dimensionless surface charge \( \tilde{Q} = \frac{q}{e\lambda_D} \).

We nondimensionalize the equations by introducing the dimensionless quantities \( X = \frac{x}{\lambda_D}, \quad Y = \frac{y}{\lambda_D}, \quad C_i = \frac{z_iC_i}{\varepsilon\psi_1}, \quad D_i = \frac{D_i}{\varepsilon\lambda_D^2}, \quad U = \frac{\alpha'\mu}{e^2c^2}\tilde{j}, \quad j_i = \frac{\alpha'\varepsilon}{e^2c^2}\tilde{j}, \quad P = \frac{\lambda^2}{e^2c^2}p, \quad \Psi = \frac{\psi}{\psi_1}, \quad \alpha = \frac{e\sqrt{\lambda_D}}{\muD^2}. \) We note that \( \alpha \) is a characteristic Péclet number for the flow outside the double layer and the typical value of \( \alpha = O(10^{-1}) \). Thus, Eqs. (1) reduce to

\[
\begin{align*}
\hat{\nabla} \cdot U &= 0, \\
-\hat{\nabla}P + \delta \hat{\nabla}^2U + \hat{\nabla}^2\Psi \hat{\nabla}\Psi &= 0, \\
-\hat{\nabla}^2\Psi &= \sum_i z_iC_i, \\
\hat{\nabla} \cdot j_i &= 0, \\
j_i &= -\delta^{-1}D_i\hat{\nabla}C_i + \alpha UC_i - \delta^{-1}D_iz_iC_i\hat{\nabla}\Psi,
\end{align*}
\]

where \( \hat{\nabla} = \lambda_D^{-1}\nabla = e_x\frac{\partial}{\partial x} + e_y\frac{\partial}{\partial y} \) is the rescaled gradient operator.

B. The thin double-layer approximation

We analyze the system of Eqs. (2) in the limit of \( \delta \to 0 \), for which the solution is thus divided into two regions: the double-layer region and the bulk region \([5,6,35]\). For a formal treatment of these regions through asymptotic analysis, we refer the readers to Schnitzer and Yariv \([33]\).

1. Analysis in the double layer

Inside the double layer, since \( x = O(a^*) \) and \( y = O(\lambda_D) \), the dimensionless \( X = O(1) \) and \( Y = O(1) \). We now use order-of-magnitude analysis to simplify Eqs. (2a)–(2e). Using Eq. (2a), we conclude \( \frac{U_x}{U_y} = O(\delta) \). Similarly, Eq. (2b) suggests that \( \frac{\partial U}{\partial y} = O(\delta) \). In the limit of \( \delta \to 0 \), these relations suggest that velocity and species flux in the \( Y \) direction are negligible when compared to their values in \( X \) direction. However, Eq. (2e) suggests an opposite relation between \( j_y \) and \( j_x \). Assuming \( C_i = O(1), \psi = O(1) \) and \( \alpha \lesssim O(1) \), we can estimate \( \frac{U_x}{U_y} = O(\delta^{-1}) \). Therefore, for the order-of-magnitude predictions between Eq. (2e) and Eq. (2d) to be consistent with each other and the condition of zero normal flux at \( Y = 0 \), the dominant terms in \( j_y \) from Eq. (2e) must vanish. Thus, the diffusion and electromigration normal to the surface are in balance (\( \delta^{-1}D_i\frac{\partial C_i}{\partial y} = 0 \)), which upon integration yields

\[
C_i = C_{i\infty}\exp[-z_i(\Psi - \Psi_{\infty})],
\]

where we have defined \( C_{i\infty}(X) \) and \( \Psi_{\infty}(X) \) as the concentration of an \( i \)th ion and the electrical potential, respectively, far away from the surface, or in the bulk. For convenience, we define \( \Psi(X, Y) = \Psi_0(X, Y) + \Psi_{\infty}(X) \), to get

\[
C_i = C_{i\infty}\exp(-z_i\Psi_0).
\]

Equation (4) is the well-known Boltzmann distribution. We note that \( \Psi_0(X, 0) = \Psi_D \) is, by definition, the surface (zeta) potential relative to the bulk.

Next, we focus on Eq. (2c): since \( X \) and \( Y \) are both \( O(1) \) in the Debye layer, \( \hat{\nabla}^2 = \delta^2\frac{\partial^2}{\partial X^2} + \delta^2\frac{\partial^2}{\partial Y^2} \approx \frac{\partial^2}{\partial Y^2} \), leading to

\[
\frac{\partial^2\Psi}{\partial Y^2} = \frac{\partial^2\Psi_0}{\partial Y^2} - \sum_i z_iC_i.
\]
Substituting Eq. (4) in Eq. (5) and integrating once while utilizing the matching condition $\frac{\partial \Psi_0}{\partial Y}|_{Y \to \infty} = 0$ [33], we get

$$\left( \frac{\partial \Psi_0}{\partial Y} \right)^2 = 2 \sum_i C_{i\infty} g_i(\Psi_0),$$

(6)

where we define

$$g_i(\Psi_0) \equiv \exp(-z_i \Psi_0) - 1.$$  

(7)

We note that $g_i(\Psi_0)$ is the excess concentration of the $i$th ion in the double layer relative to the bulk.

The typical solution strategy at this stage is to integrate Eq. (6) to find the potential $\psi(X, Y)$. However, analytical solutions are possible only for $z$; electrolytes or in the limit of small potentials. Below we develop an alternative method that circumvents this step to address the general case of electrolyte mixtures.

Gauss’s law at the surface, i.e., $y = 0$, defines the surface charge $q = -\varepsilon \frac{\partial \psi}{\partial y}|_{y = 0}$, which in dimensionless variables is $Q = -\frac{\partial \psi}{\partial Y}|_{Y = 0}$. We utilize Eq. (6) to relate $Q$ to $\psi_D$ as

$$Q = -\frac{\partial \psi}{\partial Y}|_{Y = 0} = \text{sgn}(\psi_D) \left[ 2 \sum_i C_{i\infty} g_i(\psi_D) \right]^{1/2}.$$  

(8)

Since we have solved for $C_i$ [Eq. (4)] and $\frac{\partial \psi}{\partial Y}$ [Eq. (6)], we now focus on Eq. (2b) and utilize an order-of-magnitude analysis. In the $Y$ direction, the ratio of shear stress to the electrostatic term is $O(\delta U_Y) \ll 1$, and therefore, to satisfy Eq. (2b), the pressure term balances the electrostatic term, or

$$\frac{\partial P}{\partial Y} = \frac{\partial^2 \psi_0}{\partial Y^2} \frac{\partial \psi_0}{\partial Y}.$$  

(9)

One integration in $Y$, along with $P(X, Y \to \infty) = 0$ (choosing the bulk pressure as a reference) and the matching condition $\frac{\partial \Psi_0}{\partial Y}|_{Y \to \infty} = 0$, yields

$$P(X, Y) = \frac{1}{2} \left( \frac{\partial \Psi_0}{\partial Y} \right)^2 = \sum_i C_{i\infty} g_i(\Psi_0).$$  

(10)

Physically, Eq. (10) states the well-known idea that the osmotic pressure inside the double layer is higher than that in the bulk due to the electric field inside the double layer. Therefore, $\sum_i C_{i\infty} g_i(\Psi_0)$ is a measure of the rescaled energy density inside the double layer.

Given the form of Eq. (10) in terms of $\Psi_0$, it is useful to change independent variables to express $P(X, Y) = \tilde{P}(C_{i\infty}(X), \Psi_0(X, Y))$; note that there is no loss of generality in this transformation within the thin Debye layer approximation used here. Thus, we may write

$$\frac{\partial P}{\partial X} \bigg|_Y = \sum_i \frac{\partial \tilde{P}}{\partial C_{i\infty}} \bigg|_{\Psi_0} \frac{dC_{i\infty}}{dX} + \frac{\partial \tilde{P}}{\partial \Psi_0} \bigg|_{C_{i\infty}} \frac{\partial \Psi_0}{\partial X},$$  

(11)

$$\frac{\partial P}{\partial Y} \bigg|_X = \sum_i \frac{\partial \tilde{P}}{\partial C_{i\infty}} \bigg|_{\Psi_0} \frac{dC_{i\infty}}{dY} + \frac{\partial \tilde{P}}{\partial \Psi_0} \bigg|_{C_{i\infty}} \frac{\partial \Psi_0}{\partial Y} = \frac{\partial \tilde{P}}{\partial \Psi_0} \bigg|_{C_{i\infty}} \frac{\partial \Psi_0}{\partial Y}.$$  

(12)

Comparing Eq. (12) with Eq. (9), we obtain

$$\frac{\partial \tilde{P}}{\partial \Psi_0} \bigg|_{C_{i\infty}} = \frac{\partial^2 \Psi_0}{\partial Y^2}.$$  

(13)
Utilizing Eq. (13) in Eq. (11) then results in
\[
\frac{\partial P}{\partial X} \bigg|_Y = \sum_i \frac{\partial P_i}{\partial C_{i\infty}} \bigg| \frac{dC_{i\infty}}{dX} + \frac{\partial^2 \Psi_0}{\partial Y^2} \frac{\partial \Psi_0}{\partial X}.
\] (14)

Now, we focus on the X-direction balance of Eq. (2b). Since both X and Y are O(1) in the Debye layer, Eq. (2b) can be approximated as
\[
\frac{\partial P}{\partial X} = \frac{\partial^2 U_X}{\partial Y^2} + \frac{\partial^2 \Psi_0}{\partial Y^2} \frac{\partial \Psi_0}{\partial X}.
\] (15)
Substituting Eq. (14) in Eq. (15) yields
\[
\frac{\partial^2 U_X}{\partial Y^2} = -\frac{\partial^2 \Psi_0}{\partial Y^2} \frac{\partial \Psi_0}{\partial X} + \sum_i \frac{\partial P_i}{\partial C_{i\infty}} \bigg| \frac{dC_{i\infty}}{dX}.
\] (16)

To the best of our knowledge, the form of Eq. (16) has not been noted previously. Equation (16) provides insights into the mechanism for motion as it clearly separates the contributions from the bulk electric field (electrophoresis), which is proportional to \(\frac{d\Psi_0}{dX}\), and the electric field energy inside the double layer (chemiphoresis), which is proportional to \(\frac{dC_{i\infty}}{dX}\).

Next, we integrate Eq. (16) to evaluate the diffusioosmotic slip velocity \(U_{DO} = U_X(X, Y \to \infty)\). Integration of the electrophoretic component is straightforward. We first use Eq. (10) to recognize that \(\frac{dP_i}{dC_{i\infty}} = g_i(\Psi_0)\), which on substitution into Eq. (16) yields
\[
\frac{\partial^2 U_X}{\partial Y^2} = -\frac{\partial^2 \Psi_0}{\partial Y^2} \frac{\partial \Psi_0}{\partial X} + \sum_i g_i(\Psi_0) \frac{dC_{i\infty}}{dX}.
\] (17)

We integrate this equation once across the double layer subject to the condition that Y derivatives vanish as \(Y \to \infty\) to obtain
\[
\frac{\partial U_X}{\partial Y} = -\frac{\partial \Psi_0}{\partial Y} \frac{\partial \Psi_0}{\partial X} + \sum_i \left\{ \int_Y^\infty \int_X^\infty g_i(\Psi_0(X, Y')) dY' \right\}.
\] (18)

Since the chemiphoretic contribution to the velocity distribution depends on \(Y'\) only through the excess potential \(\Psi_0(X, Y')\), it is convenient to change the integration variable from \(Y'\) to \(\Psi'\) to find
\[
\frac{\partial U_X}{\partial Y} = -\frac{\partial \Psi_0}{\partial Y} \frac{\partial \Psi_0}{\partial X} + \sum_i \left\{ \int_\Psi^\Psi_0 \int_0^{\Psi_0} g_i(\Psi') \left( \frac{\partial \Psi'}{\partial Y} \right)^{-1} d\Psi' \right\}.
\] (19)

Similarly, we integrate again across the double layer and utilize the boundary conditions on velocity and potential, i.e., \(U_X(X, 0) = 0\), and \(U_X(X, \infty) = U_{DO} = \Psi_0(X, 0) = \Psi_D\), and \(\Psi_0(X, \infty) = 0\) to get
\[
U_{DO} = \frac{d\Psi_0}{dX} \Psi_D + \sum_i \left\{ \int_0^{\Psi_0} \int_0^{\Psi_0} g_i(\Psi') \left( \frac{\partial \Psi'}{\partial Y} \right)^{-1} d\Psi' \right\} \left( \frac{\partial \Psi_0}{\partial Y} \right)^{-1} d\Psi_0.
\] (20)

Finally, we substitute the expression for \(\frac{\partial \Psi_0}{\partial Y}\) from Eq. (6) in Eq. (20) to obtain
\[
U_{DO} = \frac{d\Psi_0}{dX} \Psi_D - \frac{1}{2} \sum_i \left( \int_0^{\Psi_0} \left[ \left( \frac{\partial \Psi_0}{\partial Y} \right)^{-1} d\Psi_0 \right] \right) \left[ \sum_j C_{j\infty} g_j(\Psi) \right]^{\Psi_0} \frac{dC_{j\infty}}{dX},
\] (21)
where we recall that \(g_i(\Psi) \equiv \exp(-z_i \Psi) - 1\), as given by Eq. (7). Equation (21) is a statement relating the diffusioosmotic velocity to the excess ion concentration \(g_i(\Psi)\) and the energy density of the double layer \(\sum_i C_{i\infty} g_i(\Psi)\), albeit a complicated one. To relate \(\frac{d\Psi_0}{dX}\) and \(\frac{dC_{i\infty}}{dX}\), we now analyze Eqs. (2) in the bulk.
2. Analysis in bulk

Now, we assume that there is no external electric field applied to the system. We also invoke the condition that the surface conduction effect is negligible, i.e., the flux of ions in the direction perpendicular to the surface can be neglected [33]. Hence, in the bulk, the dependence on \( V \) vanishes, or \( C_i = C_{i\infty}(X) \), \( P = 0 \), and \( \Psi = \Psi_{\infty}(X) \). In the bulk, \( x = O(\alpha^*) \) or \( X = O(1) \). Thus, for \( \delta \to 0 \), Eq. (2c) yields the electroneutrality condition

\[
\sum_i z_i C_{i\infty} = 0. \tag{22}
\]

Moreover, since there is no externally applied electric field in diffusioosmosis, the net current in the bulk should be zero, \( \sum_i z_i J_{iX} = 0 \) [22,32]. This condition allows us to relate \( \Psi_{\infty} \) to \( C_{i\infty} \) by substituting the expression for \( J_{iX} \) from Eq. (2e) and summing over the ions to find

\[
- \sum_i z_i D_i \frac{dC_{i\infty}}{dX} + \alpha U_{DO} \sum_i z_i C_{i\infty} - \frac{d\Psi_{\infty}}{dX} \sum_i z_i^2 D_i C_{i\infty} = 0. \tag{23}
\]

Using Eq. (22), Eq. (23) yields [24,36]

\[
\frac{d\Psi_{\infty}}{dX} = - \frac{\sum_i D_i z_i^2 C_{i\infty}}{\sum_i D_i z_i^2 C_{i\infty}}. \tag{24}
\]

We note that Eq. (24) is derived by ignoring surface conduction effects, and in the absence of externally applied electric fields. However, for large \( \Psi_D \), the surface conduction effect could be important [32,33]. In Sec. IV we summarize typical scenarios encountered in experiments and discuss when the surface conduction effect is likely to be significant. A more comprehensive and rigorous theoretical analysis of this effect for a mixture of multiple ions is feasible within the current framework and should be carried out in future studies.

3. Final result

We now combine the results from the double-layer analysis and the bulk region. Substituting Eq. (24) in (21) yields

\[
U_{DO} = - \frac{1}{2} \sum_i \left( \frac{dC_{i\infty}}{dX} \right) \int_0^{\Psi_D} \left\{ \int_0^{\Psi_0} \frac{g_i(\Psi') \left[ \sum_j C_{j\infty} g_j(\Psi') \right]^{-\frac{1}{2}} d\Psi'}{\left[ \sum_j C_{j\infty} g_j(\Psi_0) \right]^{\frac{1}{2}}} \right\} d\Psi_0, \tag{25}
\]

where \( g_i(\Psi_0) \equiv \exp(-z_i \Psi_0) - 1 \) and \( C_{i\infty}(X) \) needs to satisfy the bulk electroneutrality condition, i.e., \( \sum_i z_i C_{i\infty} = 0 \). We also recall that \( \Psi_D \) is dependent on \( Q \) through Eq. (8).

Equation (25) is our main result for the diffusiophoretic velocity as a function of the zeta potential \( \Psi_D \) of the surface and the bulk ion concentration fields. To the best of our knowledge, the form of this equation has not been reported previously for the case of a general mixture of ions. We emphasize that for the evaluation of \( U_{DO} \), the expression of \( \Psi_0(X, Y) \) is not necessary, and integrating terms containing \( g_i(\Psi_0) \) is sufficient to evaluate \( U_{DO} \), as evident from Eq. (25).

It is clear from Eq. (25) that for a single salt, both the electrophoretic and chemophoretic components would be proportional to \( \frac{d\ln C_{i\infty}}{dX} \). Moreover, since \( X = O(1) \) and \( C_{i\infty} = O(1) \), the electrophoretic and chemophoretic terms are both \( O(\Psi_D) \). Last, the diffusiophoretic velocity \( U_{DP} \) is equal and opposite to the value of \( U_{DO} \) predicted in Eq. (25), i.e., \( U_{DP} = -U_{DO} \), in the limit of \( \frac{b^*}{b^r} \to 0 \), where \( b^* \) is the diameter of the particle.

C. The limit of \(|\Psi_D| \ll 1 \)

We now discuss some limiting cases for which the integrals involved in \( U_{DO} \) can be evaluated explicitly. For small potentials (\(|\Psi_D| \ll 1 \)), a Taylor series expansion shows that \( g_i(\Psi_0) = -z_i \Psi_0 + \ldots \)
\[ \frac{\varepsilon}{2} \Psi_0^2 - \frac{\varepsilon}{6} \Psi_0^3 + O(\Psi_0^4). \]

Substituting this expression into Eq. (25) and using the electroneutrality condition, we get

\[
U_{DO} = -\frac{\sum_i \mathcal{D}_i z_i dC_\infty}{\sum_i \mathcal{D}_i z_i^2 C_\infty} \Psi_D - \frac{\sum_i z_i^2 dC_\infty}{\sum_i z_i^2 C_\infty} \Psi_D^2 + \frac{5}{24} + O(\Psi_D^3).
\]

Equation (26) provides a convenient expression to evaluate \(U_{DO}\) with an order of accuracy \(O(\Psi_D^3)\).

If we neglect the cubic term, our result is consistent with previous reports \([24,27]\), where the authors arrived at their result by utilizing the potential description \(\Psi_0(X, Y)\) in the linear limit (the Debye-Hückel approximation) \([24]\). We also observe that at \(O(\Psi_D^3)\), the diffusioosmotic velocity depends explicitly on the difference of magnitude between the anion and cation valences (through \(z_i^3\)), which is an effect that is not captured by the Debye-Hückel theory.

**D. The limit of \(|\Psi_D| \gg 1\)**

We now develop useful approximations for the case of large potentials \(|\Psi_D| \gg 1\). For \(\Psi_D > 0\), in any given mixture of ions, we identify the largest anion valence by \(z_- = -\max_i(|z_i^-|)\), and approximate \(g_i(\Psi_0) \approx \exp(-z_i^- \Psi_0)\). Since, in the chemiphoretic term, \(z_-\) appears in the exponential, only anions \((i)\) with \(z_i^- = z_-\) will contribute significantly to summations involved in the chemiphoretic portion of \(U_{DO}\). Thus, we sum only over salts with anion valence \(z_-\), indicated below with a (−) over the sum. The analysis for \(\Psi_D < 0\) is similar: we identify \(z_+ = \max_i(z_i^+)\) and sum only over salts with cation valence \(z_+\), indicated below with a (+) over the sum.

Next, we observe from Eq. (25) that evaluating integrals involves the limit from 0 to \(\Psi_0\) or 0 to \(\Psi_D\). The approximation of \(g_i(\Psi_0)\) above is inaccurate near the lower integration limit and can be shown, as a result, to overestimate the integrals by a subdominant \(O(1)\) term for \(|\Psi_0| \gg 1\). We correct for this error by modifying the lower limit of the integral in Eq. (25) from 0 to \(\Psi_\ell\), where \(\Psi_\ell\) is an \(O(1)\) parameter whose value we discuss later. As we will show, the introduction of \(\Psi_\ell\) leaves the leading-order asymptotic behavior of \(U_{DO}\) unchanged for \(|\Psi_D| \gg 1\), while allowing us obtain a significant improvement in the numerical accuracy of our predictions for moderate zeta potentials \(|\Psi_D| \gtrsim 1\).

Thus substituting (1) \(g_i(\Psi_0) \approx \exp(-z_i^- \Psi_0)\) for \(\Psi_D \gg 1\), (2) \(g_i(\Psi_0) \approx \exp(-z_i^+ \Psi_0)\) for \(-\Psi_D \gg 1\), and (3) modifying the lower limits of integration from 0 to \(\Psi_\ell\) in Eq. (25), we obtain

\[
U_{DO} = \left\{ \begin{array}{ll}
-\frac{\sum_i \mathcal{D}_i z_i dC_\infty}{\sum_i \mathcal{D}_i z_i^2 C_\infty} \Psi_D + \sum_i^{(+)} \frac{dC_\infty}{C_\infty} (\Psi_\ell + \Psi_D - \frac{2\exp[z_+ (\Psi_\ell + \Psi_D)/2 - 1]}{z_+^2}), & -\Psi_D \gg 1 \\
-\frac{\sum_i \mathcal{D}_i z_i dC_\infty}{\sum_i \mathcal{D}_i z_i^2 C_\infty} \Psi_D - \sum_i^{(-)} \frac{dC_\infty}{C_\infty} (\Psi_D - \Psi_\ell + \frac{2\exp[-z^- (\Psi_D - \Psi_\ell)/2 - 1]}{-z^-}), & \Psi_D \gg 1. \end{array} \right.
\]

To leading order, observe that the diffusioosmotic velocity is linear in \(\Psi_D\) for \(|\Psi_D| \gg 1\). The next order is independent of \(\Psi_D\) but depends on the \(O(1)\) parameter \(\Psi_\ell\), for which we determine an approximation in the next section. In Sec. V we detail the accuracy of the Debye-Hückel limit and the \(|\Psi_D| \gg 1\) approximations. We reiterate that our analysis assumes that finite ion size, dielectric decrement, and surface conduction effects are negligible, which may become important for \(|\Psi_D| \gg 1\), as we detail below.

**IV. VALIDITY OF DERIVED PREDICTIONS**

In this section, we address the validity of the assumptions we invoked during our analysis. Specifically, we discuss the following assumptions: (A) thin double layer, (B) ions are point charges.

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TABLE I. Summary of physical parameters and relevant dimensionless groups from recent experiments on diffusiophoresis that justify our assumptions. To estimate the values, \( d_{\text{ion}} = 3 \text{ Å} \) is assumed for the diameter of an ion (see Sec. IV B) and \( \gamma = 10 \text{ [M]^{-1}} \) is assumed for the dielectric decrement factor for an ion (see Sec. IV C).

<table>
<thead>
<tr>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>( c^* )</td>
<td>50 [mM]</td>
<td>50 [mM]</td>
<td>2 [mM]</td>
<td>1 [mM]</td>
</tr>
<tr>
<td>( a^* )</td>
<td>60 [μm]</td>
<td>100 [μm]</td>
<td>500 [μm]</td>
<td>100 [μm]</td>
</tr>
<tr>
<td>( b^* )</td>
<td>200 [nm]</td>
<td>500 [nm]</td>
<td>60–1000 [nm]</td>
<td>500 [nm]</td>
</tr>
<tr>
<td>( \lambda_D )</td>
<td>1.3 [nm]</td>
<td>1.3 [nm]</td>
<td>6.7 [nm]</td>
<td>9.5 [nm]</td>
</tr>
<tr>
<td>( \frac{\lambda_D}{a^*} )</td>
<td>( 2.2 \times 10^{-5} )</td>
<td>( 1.3 \times 10^{-5} )</td>
<td>( 1.3 \times 10^{-5} )</td>
<td>( 9.5 \times 10^{-5} )</td>
</tr>
<tr>
<td>( \frac{\lambda_D}{b^*} )</td>
<td>( 6.7 \times 10^{-3} )</td>
<td>( 2.6 \times 10^{-3} )</td>
<td>( 6.7 \times 10^{-3} ) to ( 1.1 \times 10^{-1} )</td>
<td>( 5.2 \times 10^{-3} )</td>
</tr>
<tr>
<td>( \frac{1}{\zeta} \ln \left( \frac{1}{d_{\text{ion}} c^*} \right) )</td>
<td>7</td>
<td>7</td>
<td>10</td>
<td>11</td>
</tr>
<tr>
<td>( \frac{1}{\zeta} \ln \left( \frac{\Psi_D}{\gamma c^*} \right) )</td>
<td>5</td>
<td>5</td>
<td>8</td>
<td>9</td>
</tr>
<tr>
<td>( 2 \ln \left( \frac{1}{\lambda_D} \right) )</td>
<td>10</td>
<td>12</td>
<td>4–10</td>
<td>8</td>
</tr>
</tbody>
</table>

(assumed implicitly through the Nernst-Planck equations; see below for details), (C) electrical permittivity is constant, and (D) surface conduction can be ignored. To facilitate this discussion, we compiled data from some of the previously published experimental studies on diffusiophoresis and estimated the physical parameters along with the relevant dimensionless groups; see Table I.

A. Thin double layer

The common thin double-layer assumption is central to the validity of our derived relation in Eq. (25). Typically, experiments are conducted on diffusiophoresis (and not diffusioosmosis), i.e., the scenario when charged particles move under the presence of a salt gradient while the electrolyte is stationary. In diffusiophoresis, there are three length scales: (1) particle size \( b^* \), (2) Debye length \( \lambda_D \), and (3) the length scale over which the imposed ion concentration decays \( a^* \). To utilize Eq. (25) for describing experiments, in addition to \( a^* = \delta \ll 1 \), \( \frac{\lambda_D}{b^*} \ll 1 \) also needs to be satisfied. Based on previously published experimental studies in Table I, we observe that \( \lambda_D = 1–10 \text{ nm} \), \( a^* = 100–500 \text{ μm} \), and \( b^* = 60–1000 \text{ nm} \). The table shows that \( \frac{\lambda_D}{a^*} \gtrsim O(10^{-4}) \) and \( \frac{\lambda_D}{b^*} \gtrsim O(10^{-2}) \). Therefore, the thin double-layer approximation is applicable to typical experimental scenarios. However, we note that when \( b^* \lesssim 100 \text{ nm} \), \( \frac{1}{\zeta} \ln \left( \frac{1}{d_{\text{ion}} c^*} \right) \gtrsim O(10^{-1}) \), and there might be some deviations from the velocities predicted by Eq. (25).

B. Ions are point charges

Our analysis utilizes the Nernst-Planck equations, i.e., Eqs. (2d)–(2e), to solve for the diffusioosmotic and diffusiophoretic velocities. The Nernst-Planck description assumes that ions are point charges and the volume fraction of ions can be ignored when compared to the volume fraction of solvent [28]. However, for large \( c^* \) and \( \Psi_D \), the concentration of surface-attracted ions inside the double layer can be large, and finite ion size effects could become important [26,28,29]. The dimensionless number that governs this effect is \( d_{\text{ion}}^3 c^* \), where \( d_{\text{ion}} \) is the diameter of an ion. Physically, this parameter is the dimensionless volume fraction of ions in the bulk. The finite ion size effects are not significant when \( |\Psi_D| \lesssim \frac{1}{\zeta} \ln \left( \frac{1}{d_{\text{ion}} c^*} \right) \) [26]. To estimate this value, it is crucial to estimate the value of \( d_{\text{ion}} \). Typically, \( d_{\text{ion}} \approx 1 \text{ Å} \), although previous studies have argued that the ion diameter should include the hydration shell, modifying this estimate to \( d_{\text{ion}} \approx 3 \text{ Å} \) [28]. Therefore, we assumed that \( d_{\text{ion}} = 3 \text{ Å} \) to estimate values. Table I shows that finite ion sizes are not significant.
for $|\Psi_D| \lesssim 7$. Since most experiments are performed for particles with $\Psi_D \lesssim 4$, we expect that finite ion size effects are not significant for diffusiophoretic and diffusioosmotic velocity predictions.

### C. Electrical permittivity is constant

We also assumed that the electrical permittivity $\varepsilon$ is constant. However, inside the double layer, the electrical permittivity can decrease due to a large concentration of ions. This effect is known as the dielectric decrement, and, similar to finite ion size, it also becomes significant at large $c^*$ and $\Psi_D$ [26,30,31]. Specifically, this effect is not significant when $|\Psi_D| \lesssim \frac{1}{7} \ln (\frac{c^*}{\gamma})$ [26], where $\gamma$ is the dielectric decrement factor and $\varepsilon_w$ is the dielectric constant of the solvent. In our analysis, we assume that the solvent is water where $\varepsilon_w = 78$. To estimate the validity, we assumed $\gamma = 10 \ [M]^{-1}$ [30]. We show in Table I that for $|\Psi_D| \lesssim 5$, this effect can be ignored. Therefore, a change in the dielectric constant is unlikely to be significant for typical diffusiophoretic experiments.

### D. Surface conduction

As mentioned previously, within the Poisson-Boltzmann framework, for large $\Psi_D$, a more careful treatment of the bulk region is required to account for the effect of surface conduction where the currents from the double layer can leak into the bulk region [32,33]. This effect is not significant for $|\Psi_D| \lesssim 2 \ln (\frac{1}{\delta})$ [33]. Table I demonstrates that this effect may not be significant for $|\Psi_D| \lesssim 10$ and would likely not influence typical experiments in diffusiophoresis and diffusioosmosis.

In addition to the above mentioned assumptions, we note that we have not considered specific ion interactions in our analysis. This effect could be taken into account by assuming a Stern layer near the surface and specifically treating ion interactions [37], which will be pursued in our future work.

### V. ONE ELECTROLYTE WITH ASYMMETRIC VALENces

We now discuss the results from Eq. (25) for the case of a single electrolyte. In this section, we also tabulate diffusiophoretic velocities for some common valence-asymmetric electrolytes such as CaCl₂, Na₂SO₄, H₂SO₄, and others. Since we focus on the case of a single electrolyte in this section, for further discussion, we omit the $i$ subscript from $C_{i\infty}, g_i(\Psi_0)$ etc.

#### A. Exact solution

For a solution consisting of a single electrolyte, to maintain electroneutrality, we assume that there is one cation with valence $z_+$ and one anion with valence $z_-$. Moreover, to satisfy electroneutrality in the bulk, we write $C_{+\infty} = -z_-C_{-\infty}$ and $C_{-\infty} = z_+C_{+\infty}$. Utilizing Eqs. (7) and (25), we obtain

$$U_{DO} = -\beta \Psi_D - \frac{1}{2} \int_0^{\Psi_0} \frac{[z_- \exp(-z_+ \Psi') + z_+ \exp(-z_- \Psi') - z_+ + z_-]^{1/2}}{[z_- \exp(-z_+ \Psi_0) + z_+ \exp(-z_- \Psi_0) - z_+ + z_-]^{1/2}} d\Psi'.$$

(28)

where $\beta = \frac{D_+ - D_-}{z_+D_+ - z_- D_-}$. We note that the value of $\beta$ here is consistent with the idea of a junction potential that is generally suggested for asymmetric diffusivities [38,39]. We now evaluate Eq. (28) in different limits. For one electrolyte and $z_+ = -z_-$, $z$, Eq. (28) reduces to the well-known result [6,27,35]

$$U_{DO} = -\frac{d \ln C_{\infty}}{dX} \left\{ \frac{1}{2} \beta \Psi_D + \frac{4}{z^2} \ln \left[ \cosh \left( \frac{z \Psi_D}{4} \right) \right] \right\}.$$

(29)
B. Analytical approximations

In the limit of $|\Psi_D| \ll 1$, Eq. (26) or Eq. (28) reduces to

$$\frac{U_{DO}}{d \ln C_\infty}{dx} = -\beta \Psi_D - \frac{\Psi_D^2}{8} - (z_+ + z_-) \frac{\Psi_D^3}{216} + O(\Psi_D^4).$$  

(30)

Equation (30) shows that the electrophoresis term is affected by valence through the definition of $\beta$. The quadratic terms of the chemiphoretic contribution is independent of the valence, whereas the cubic correction is influenced by the difference between the (unsigned) valences of the cation and anion. For $|\Psi_D| \gg 1$, Eq. (27) reduces to

$$\frac{U_{DO}}{d \ln C_\infty}{dx} = \begin{cases} -\beta \Psi_D + \left( \frac{\Psi_D + \Psi_\ell}{z_+} \right)^{-1} & -\Psi_D \gg 1 \\ -\beta \Psi_D - \left( \frac{\Psi_D - \Psi_\ell}{z_-} \right)^{-1} & \Psi_D \gg 1. \end{cases}$$  

(31a)

(31b)

Therefore, as $|\Psi_D|$ increases, the impact of asymmetry in cation and anion valences on the chemiphoretic contribution also increases. Comparing the approximation (31) with the exact solution (29) for large zeta potentials, we find that the two expressions agree identically at leading (linear) order in $\Psi_D$. Subdominant terms in both expressions are $O(1)$; requiring agreement at this order determines the parameter

$$\Psi_\ell = \begin{cases} z_+^{-1}(4 \ln 2 - 2), & -\Psi_D \gg 1 \\ -z_-^{-1}(4 \ln 2 - 2), & \Psi_D \gg 1. \end{cases}$$  

(32a)

(32b)

With these choices for $\Psi_\ell$, (31) and (29) differ only by a terms exponentially small in $|\Psi_D|$. We note that the $\Psi_\ell$ values determined above are formally valid for a single $z_\pm$ electrolyte, though we expect them to provide reasonable approximations for more complicated situations such as mixtures of valence-asymmetric electrolytes (see Fig. 2 and the Appendix). We reiterate that the leading-order behavior of our approximation Eq. (31) for large potentials remains independent of $\Psi_\ell$.

C. Validity of analytical approximations

We now present the variation of $U_{DO}$ by numerically integrating the exact expression Eq. (28) for some typical $z_+ z_-$ electrolytes, and discuss the accuracy and applicability of approximations Eqs. (30)–(31). The properties of common electrolytes are summarized in Table II, and typical results for $U_{DO}$ are shown in Fig. 2. We find that though Eq. (30) is derived for $|\Psi_D| \ll 1$, it is a good approximation of the exact result from Eq. (28) for $|\Psi_D| \lesssim 1$; see Fig. 2(a). However, we discover that Eq. (30) with accuracy up to $O(\Psi_D^3)$ is often closer to the numerical results as compared to Eq. (30) with accuracy only up to $O(\Psi_D^2)$, depending on the value of $\beta$, $z_+$, and $z_-$. Therefore, the improvements due to higher-order corrections are restricted for larger $|\Psi_D|$ and Eq. (30) should be utilized for $|\Psi_D| \lesssim 1$. In contrast, Eq. (31) gives a more accurate predication for larger $|\Psi_D|$ and is best utilized for $|\Psi_D| \gtrsim 4$, though in many cases can also provide good approximations for smaller potentials; see Fig. 2(b). We note that before utilizing Eq. (31), it should be ensured that effects such as finite ion size, dielectric decrement, and surface conduction are not significant. The results in Fig. 2(b) were generated using the values of $\Psi_\ell$ provided in Eq. (32). We discuss the sensitivity of our results on the choice of $\Psi_\ell$ in the Appendix. We emphasize that $z_+$ and $z_-$ significantly influence $U_{DO}$, as evident from Eq. (31) and Figs. 2(a) and 2(b).
D. Some common $z_+; z_-$ electrolytes

We now present a comparison of $U_{DO}$ for different electrolytes, and the results are summarized in Fig. 3. We first compare HCl and H$_2$SO$_4$. Since the cation is H$^+$ in these two electrolytes, $\beta$ values are large and positive in both cases. For small $|\Psi_D|$, the ratio of electrophoretic and chemiphoretic contributions is $\frac{\beta}{\Psi_D}$; see Eq. (30). Therefore, for HCl and H$_2$SO$_4$, the two contributions are cooperative for $\Psi_D > 0$ and competitive for $\Psi_D < 0$. Next, we compare NaCl and Na$_2$SO$_4$. Though the cation for both of these electrolytes is the same, due to the smaller diffusivity of SO$_4^{2-}$ as compared to Cl$^-$, and because the valence of SO$_4^{2-}$ is higher, there is a considerable difference between the $\beta$ values of the two electrolytes (see Table II). We find that since $\beta$ is negative for NaCl, for $\Psi_D > 0$, the electrophoretic and chemiphoretic components compete with each other. In contrast, since the electrophoretic effect is weak for Na$_2$SO$_4$, for $\Psi_D > 0$, the diffusioosmotic velocity of Na$_2$SO$_4$ is larger in magnitude as compared to NaCl. However, the opposite is true for $\Psi_D < 0$, where for NaCl, the electrophoretic and chemiphoretic components are similarly signed, and thus the case of NaCl has a larger magnitude of diffusioosmotic velocity as compared to Na$_2$SO$_4$. A similar comparison can be drawn between CaCl$_2$ and Mg(HCO$_3$)$_2$. These examples demonstrate that valence-asymmetric electrolytes can be utilized for control over diffusioosmotic and diffusiphoretic processes and may influence other electrokinetic phenomena.

<table>
<thead>
<tr>
<th>Electrolyte</th>
<th>$z_+$</th>
<th>$z_-$</th>
<th>$D_+ [10^{-9} \text{ m}^2 \text{s}^{-1}]$</th>
<th>$D_- [10^{-9} \text{ m}^2 \text{s}^{-1}]$</th>
<th>$\beta$</th>
</tr>
</thead>
<tbody>
<tr>
<td>HCl</td>
<td>1</td>
<td>-1</td>
<td>9.31</td>
<td>2.03</td>
<td>0.64</td>
</tr>
<tr>
<td>H$_2$SO$_4$</td>
<td>1</td>
<td>-2</td>
<td>9.31</td>
<td>1.06</td>
<td>0.72</td>
</tr>
<tr>
<td>NaCl</td>
<td>1</td>
<td>-1</td>
<td>1.33</td>
<td>2.03</td>
<td>-0.21</td>
</tr>
<tr>
<td>Na$_2$SO$_4$</td>
<td>1</td>
<td>-2</td>
<td>1.33</td>
<td>1.06</td>
<td>0.08</td>
</tr>
<tr>
<td>CaCl$_2$</td>
<td>2</td>
<td>-1</td>
<td>0.79</td>
<td>2.03</td>
<td>-0.34</td>
</tr>
<tr>
<td>Mg(HCO$_3$)$_2$</td>
<td>2</td>
<td>-1</td>
<td>0.70</td>
<td>1.18</td>
<td>-0.18</td>
</tr>
</tbody>
</table>

TABLE II. Values of $z_+; z_-, D_+, D_-$, and $\beta$ for some common electrolytes at 25°C. The ion diffusivities are taken from Ref. [27].
VI. CONCLUSIONS

In this article, we presented general results for diffusioosmotic and diffusiophoretic velocities for a mixture of valence-asymmetric electrolytes with arbitrary $z_i$. Equation (25) includes the effect of valence asymmetry for a mixture of electrolytes, and Eq. (28) describes the dependence for a single electrolyte. We also presented the approximate solutions in the limit of $|\Psi_D| \ll 1$ and $|\Psi_D| \gg 1$; see Eqs. (26), (27), (30), and (31). We demonstrated that asymmetry in electrolyte valence is a useful parameter to tune the diffusiophoretic and diffusioosmotic motions. Our analysis will motivate future experimental studies on diffusiophoresis and diffusioosmosis using mixture of electrolytes and multivalent salts. Future theoretical studies in this area could focus on including the effects of finite ion size, dielectric decrement, and surface conduction in the analysis to further generalize diffusiophoretic and diffusioosmotic velocity relations.

ACKNOWLEDGMENTS

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APPENDIX: THE CHOICE OF $\Psi_\ell$

We now discuss the dependence of $U_{DO}$ on $\Psi_\ell$ when Eq. (31) is utilized to evaluate $U_{DO}$. We find that $\Psi_\ell$ from Eq. (32) provides the best approximation to the numerical solution as compared to other values; see Fig. 4. However, all values of $\Psi_\ell$ between 0 and 1 work reasonably well, but for larger values of $\Psi_\ell$, such as $\Psi_\ell = 5$, the deviation from the numerical solution is significant.