# UNVIERSITY OF CALIFORNIA SAN DIEGO

# Anisotropic Wetting Property of Superhydrophobic Surfaces and Electrokinetic Flow on Liquid-Filled Surfaces

# A Dissertation submitted in partial satisfaction of the requirements for the degree Doctor of Philosophy

in

Engineering Sciences (Mechanical Engineering)

by

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2019

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#### ACKNOWLEDGEMENTS

I would like to acknowledge Professor Prabhakar Bandaru for his support as the chair of my committee and in my research endeavors. I'm grateful to his patience in my Ph. D period and his valuable help and advice on my career development. I would also like to thank my lab mates for their help in my research. I would also like to acknowledge my committee member who provided precious suggestions in my research and support for my career.

I would like to thank my parents for their support and patience in my Ph. D studies. I would also like my thank my friends for their help both in my study and life.

Chapter 2, in full, is a reprint of the materials as it appears in Fan, B. and Bandaru, P.R., Anisotropy in the hydrophobic and oleophobic characteristics of patterned surfaces, Appl. Phys. Lett, 111, 261603, 2017. The dissertation author was the primary investigator and author of this paper.

Chapter 3, in full, is a reprint of the materials as it appears in Fan, B., Bhattacharya, A. and Bandaru, P.R., Enhanced voltage generation through electrolyte flow on liquid-filled surfaces, Nature Communications, 9, 4050, 2018. The dissertation author was the primary investigator and author of this paper.

Chapter 4, in full, is a reprint of the materials as it appears in Fan, B. and Bandaru, P.R., Modulation of streaming potential and slip characteristics in electrolyte flow over liquid-filled surfaces, Langmuir, 2019, 35, 18, 6203 – 6210. The dissertation author was the primary investigator and author of this paper.

Chapter 5, in full, is a reprint of the materials as it appears in Fan, B. and Bandaru, P.R., Tensorial modulation of electrokinetic streaming potentials on air and liquid filled surfaces, Langmuir, 2019, 35, 46, 14812 – 14817. The dissertation author was the primary investigator and author of this paper.

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Fan, B. and Bandaru, P.R., Modulation of streaming potential and slip characteristics in electrolyte flow over liquid-filled surfaces, *Langmuir*, 2019, 35, 18, 6203 – 6210.

Fan, B., Bhattacharya, A. and Bandaru, P.R., Enhanced voltage generation through electrolyte flow on liquid-filled surfaces, *Nature Communications*, 9, 4050 (2018).

Fan, B. and Bandaru, P.R., Anisotropy in the hydrophobic and oleophobic characteristics of patterned surfaces, *Appl. Phys. Lett*, 111, 261603 (2017).

#### ABSTRACT OF THE DISSERTATION

# Anisotropic Wetting Property of Superhydrophobic Surfaces and Electrokinetic Flow on Liquid-Filled Surfaces

by

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#### Doctor of Philosophy of Engineering Science (Mechanical Engineering)

University of California San Diego, 2019

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Understanding the wetting property of rough surface is critical in guiding droplets and novel superhydrophobic surface design. The Cassie-Baxter model and Wenzel model are always used to describe the totally non-wetting and completely wetting states, however, there were few discussions about the intermediate state. Through measuring the contact angles of groove patterned surfaces in different groove orientations, the anisotropic wetting properties of groove patterned superhydrophobic surface were investigated. The degree of water penetration into the grooves was experimentally observed and it was found that the degree of water penetration was different with groove orientations, which would affect the corresponding contact angle. Besides guiding droplets, superhydrophobic surfaces are also very important in microfluidic due to their ability to generate fluid slip and flow enhancement. After a deeper understanding

of the wetting property of groove patterned superhydrophobic surface, I further investigated its important role in microfluidics. In this dissertation, I mainly focus on electrokinetics on groove patterned surface and liquid-filled slippery surfaces, a new kind of surface by filling low surface tension oil into the interstices of groove patterned surfaces. I experimentally measured the streaming potential on flat parylene surface, airfilled groove patterned surface and liquid-filled surfaces and compared their effects in streaming potential enhancement. The liquid-filled surfaces were shown to be able to enhance the generated streaming potential due to its slippery property and liquid-oil interface charges. As the electrokinetic on liquid-filled surfaces is a new phenomenon, the underlying physics is still not clear. I further investigated the influences of filled oil properties and groove orientation on streaming potentials and fluid slip. Oils with different densities, viscosities, dielectric constant, conductivities and surface tensions were filled into the interstices of groove patterned surfaces to make different types of liquid-filled surfaces. The streaming potentials on liquid-filled surfaces with different oils were experimentally measured. An empirical relationship between streaming potential and oil properties was found and the effects of electrical properties, such as interface charge density and dielectric constant of filled oil, on fluid slip were also studied. Finally, the groove orientation was varied to study the tensorial effects on streaming potential. Through both streaming potential measurement and theoretical analysis, it was found that the streaming potential at  $45^{\circ}$  was always smaller than the arithmetic mean of those at  $0^{\circ}$  and  $90^{\circ}$ , and the pressure gradient in the transvers direction generated by tensorial effects was important in the streaming potential modification. My work will be important in guiding droplets, flow patterning, lab-on-chip devices and the development of electrokientic based power sources.

# **Chapter 1 Introduction**

#### 1.1 Wetting models on rough surface

When a water droplet contacts with a solid surface, there exists a thermodynamic equilibrium state of the three phases that are liquid phase, vapor phase and solid phase and form a contact angle ( $\theta_c$ ) as shown in Figure 1.1. Through the force balance at the contact line or triple line, the contact angle can be calculated using the Young equation<sup>1</sup>:

$$\gamma_{SA} - \gamma_{SL} - \gamma_{LA} \cos\theta_c = 0 \tag{1.1}$$

 $\gamma_{SA}$  is the solid-air interface energy,  $\gamma_{SL}$  is the solid-liquid interface energy and  $\gamma_{LA}$  is the liquid-air interface energy.



**Figure 1.1** (a) Liquid droplet on flat surface and the formation of contact angle, (b) Cassie and (c) Wenzel state on rough surface.

When the liquid droplet contacts with a rough surface, the Cassie and Wenzel model are usually used to describe the wetting state of the substrate. The Cassie state, indicating in Figure 1.1(b), describes the completely non-wetting state at which the liquid droplet will stay above the pattern. The Wenzel state in Figure 1.1(c) describes the totally wetting state and the liquid will completely fill into the interstices of the rough surface.

#### **1.4 Electrokinetics**

When electrolyte contacts with a solid surface, there will form an electrochemical double layer (EDL) structure due to the mismatch of dielectric constant ( $\varepsilon$ ). Generally, the electrolyte with a larger dielectric constant flows over a material with smaller dielectric constant, the solid surface will be negatively charged<sup>2</sup>. At the electrolyte-solid interface, the negative charged solid interface will attract positive counterions near the solid surface and form the doubly layer structure. The first layer will be stern layer which consists fixed positive ions adhered to the solid surface and those positive ions can not move with electrolyte. The second layer is diffuse layer which contains mostly positive ions which can move with the fluid flow. The characteristic thickness of EDL is defined as Debye length  $\lambda_D$  as indicated in Figure 1.2. The  $\lambda_D$  =  $\sqrt{\frac{\varepsilon_0 \varepsilon_r k_B T}{2N_A(ze)^2 l}}$ ,  $\varepsilon_0$  is the dielectric constant of free space,  $\varepsilon_r$  is the relative dielectric constant of the electrolyte,  $k_B$  is Boltzmann constant, T is the temperature,  $N_A$  is Avogadro's number, z is the number of charge of the ion, *e* is the elementary electron charge and *I* is the concentration, the  $\lambda_D \sim \frac{9.6}{\sqrt{I}}$  (the units of  $\lambda_D$  and *I* are nm and mM) for electrolytes composed of ions with single charge. Under a pressure driven flow, say the pressure drop is  $\Delta P = P_2 - P_1$ , as indicated in Figure 1.2, the positive ions in the diffuse layer will move with the fluid and accumulate at the right end of the channel, due to the ion accumulation, an imbalance between the ion concentration is built up between the ends of the channel which can induce a potential difference termed as streaming potential<sup>2</sup>. The generated streaming potential,  $V_s$ , is related to the dielectric constant  $\varepsilon$  (= $\varepsilon_r \varepsilon_0$ ), viscosity  $\eta$ , conductivity  $\sigma$ , the zeta potential  $\zeta$  of the surface and the pressure drop  $\Delta P$ through<sup>2</sup>:

$$V_{S} = \frac{\varepsilon \zeta}{\eta \sigma} \Delta P \tag{1.2}$$

The derivation of Eqn. 1.2 is based on the following assumptions<sup>2</sup>: (a) negligible surface conductivity, (b) thin EDL, (c) the Debye-Huckel assumption that the wall potential is of the order of ~ 26 mV (= $k_BT/e$ , where  $k_B$  is the Boltzmann constant ~ 8.6 × 10<sup>-5</sup> eV/K, T is the temperature at ~ 300 K in the ambient, and e is the elementary electronic charge = 1.602 × 10<sup>-19</sup> C). As electrokinetic flow can

convert mechanical energy to electricity and there is no moving parts in the structure, it is promising to be developed as microchannel battery<sup>3</sup>, the power sources for MEMS system and wearable devices.



**Figure 1.2** Electrokinetic flow on a flat surface under a pressure drop  $\Delta P (=P_2 - P_1)$ .  $\zeta$  is the zeta potential which is the potential at the shear plan where fluid starts to move.

#### **1.3 Motivation**

Superhydrophobic surfaces which are inspired by lotus leaf and other plant leaves that can not get wet by water are important in both static and dynamic fluid phenomena and have essential practical applications, such as self-clean surface, anti-icing, oil-water separation and drag reduction<sup>4</sup>. To make superdydrophobic surfaces, the traditional method is to make micro/nano scale roughness that will trap air to increase the water contact angle and decrease the tendency of the binding of water to the surface. The wetting property of water on rough surfaces are important in superhydrophobic design. As discussed previously, the Cassie model and Wenzel model are usually used to describe the wetting status of liquid droplet on rough surface. However, there are other conditions that these two models don't consider. For example, both of these two models don't consider the effect of anisotropic property of the roughness on the wetting property. Besides, these two models only describe the two extreme wetting states that are completely wetting and non-wetting states, and there were few discussions abut the intermediate wetting state between these two models. To have a better understanding of the liquid wetting states on rough surfaces, more work should be done to consider the anisotropic property of the superhydrophobic surface and the partial wetting state to make modifications of current Cassie and Wenzel model.

Besides studying the static liquid wetting on superhydrophobic surface, my work also extended to the continuous and dynamic flow on superhydrophobic due to the important role of superhydrophobic surface in fluid drag reduction and flow enhancement. In electrokinetic field, traditional researches were mainly on flat surface which introduced non-slip boundary condition at the electrolyte-solid interface and yielded low streaming potential<sup>3</sup>, which limited the applications of electrokinetic as practical power sources. Then it was proposed that using superhydrophobic surface can introduce fluid slip for streaming potential enhancement<sup>5</sup>. Since the electrical potential decays away from the surface, the further the shear plane is from the substrate boundary the smaller the equivalent  $\zeta$  and the obtainable  $V_s$ . One possible method to enhance the  $\zeta$  is then to introduce fluid slip at the substrate surface, where the shear plane is moved as close as possible to the substrate. In this case, the zero velocity boundary condition at the surface (y=0) is replaced with a Navier slip condition:  $u_s$  (y=0)=  $b \frac{\partial u (y=0)}{\partial y}$ ) with *b* as the slip length (a length below the surface at which the fluid effectively has a zero velocity)<sup>6.7</sup>. The  $\zeta$  is modified by the introduction of the Navier condition to<sup>8</sup>:

$$\zeta = \psi_o (1 + \frac{b}{\lambda_D}) \tag{1.3}$$

Hence, a larger slip (implying a larger *b*) relative to a  $\lambda_D$  for a given solution (with a fixed  $\varepsilon$ ,  $\sigma$  and  $\eta$  could be beneficial in harnessing large  $V_s$  – through a closer coupling of the fluid velocity profile to the charged double layer. In this context, it was previously indicated<sup>9</sup> that relatively large values of *b* of the order of ~ 6 mm may indeed be obtained in superhydrophobic surfaces fabricated through a roll-to-roll processing methodology<sup>10</sup>. As  $\lambda_D \sim \frac{9.6}{\sqrt{l}}$ ,  $\lambda_D$  is around 9.6 nm with 1 mM NaCl, and a consequent  $b/\lambda_D$  of ~ 625, one may be led to expect up to three orders of magnitude larger  $\lambda_D$ . However, there have been theoretical suggestions that the superhydrophobic surfaces may not yield the predicted degree enhancement due to the non-charged liquid-air interfaces introduced by superhdrophobic surfaces<sup>11</sup>. Most of the work on

electrokinetics on superhydrophobic surfaces was theoretical and computational analysis, experimental efforts are necessary to investigate the practical situation. In my dissertation, I will also mainly focus on the experimental study on electrokinetics on superhydrophobic surfaces and will also come up with the solution to streaming potential enhancement.

# Chapter 2 Anisotropy in the hydrophobic and oleophilic characteristics of patterned surfaces

#### 2.1 Abstract

A significant difference in the wetting angles of water and oil was observed on patterned substrates, combining interstitial spaces along with hydrophobic solid surfaces, as a function of orientation. The difference was ascribed to a modification of the liquid – interstice interfacial surface energy, due to different degrees of penetration of the liquid. A roughness metric, related to the ratio of the areas related to the extent of which the liquid infiltrates the interstice to the geometrically determined area is proposed. The study has implications in modulating surface slip behavior and would be of importance in guiding liquid droplets, e.g., in electrophoresis.

#### **2.2 Introduction**

The extent of wettability of a surface is dependent on the underlying substrate surface energies ( $\gamma$ ). Considering the substrate-wetting liquid energy ( $\gamma_{SL}$ ), the substrate-ambient energy ( $\gamma_{SA}$ ), and the liquid ambient (*e.g.*, air) energy ( $\gamma_{LA}$ ), two related well-known parametric measures<sup>1</sup> are the spreading parameter, *S*, and the contact/wetting angle:  $\theta$  - determined through the Young relation, *i.e.*,

$$S = \gamma_{SA} - (\gamma_{SL} + \gamma_{LA}) \tag{2.1a}$$

$$\theta = \cos^{-1}[(\gamma_{SA} - \gamma_{SL})/\gamma_{LA}]$$
(2.1b)

The extension to predict the wetting behavior of rough or patterned surfaces has been done through the Cassie-Baxter/*fakir* model: Figure 2.1(a) or the Wenzel (W) model: Figure 2.1(b), which broadly deal with two extremes related to complete exclusion (/inclusion) of the interstices in the solid substrate by the liquid, respectively. In this letter, we investigate the plausible intermediate regime, *i.e.*, Figure 2.1(c), where there would be partial liquid penetration into patterned substrates: Figure 2.1(d). The parameters of relevance in the Cassie-Baxter and Wenzel models are the respective area fractions of the interstice surface area ( $\phi_I$ ) and the solid surface area ( $\phi_s$ ), where  $\phi_I + \phi_s = 1$ , and a somewhat arbitrarily defined roughness parameter: r, as the ratio of the total surface area to the projected/apparent surface area, which would of course depend on the measurement scale. For instance, the Cassie-Baxter model relates the *overall* wetting angle ( $\theta_{net}$ ) to the wetting angle on the air in the interstice (*i.e.*,  $\theta_{int}$ ) and that in the constituent solid (*i.e.*,  $\theta_{sol}$ ) as:

$$\cos\left(\theta_{\text{net}}^{F}\right) = \phi_{I} \cos\left(\theta_{\text{int}}\right) + \phi_{s} \cos\left(\theta_{\text{sol}}\right)$$
(2.2a)

Alternately, the Wenzel model indicates a roughness (*r*) related  $\theta^{W}_{net}$ , through:

$$\cos\left(\theta^{W}_{\text{net}}\right) = r \cos\left(\theta_{\text{sol}}\right) \tag{2.2b}$$

Per recommendations<sup>12</sup> on the proper use of Eqn. 2.2, we did not assume that the liquid-vapor interface was smooth, and consider liquid penetration into the troughs of the rough surface<sup>13</sup>. An effective medium approach (EMA)<sup>9,10</sup> was implicitly assumed to average the water penetration, whereby the extent of the wetting liquid is much larger compared to the underlying texture, and the perturbations to the liquid drop in the interstice could be considered small.

However, a major issue is that these models do not explicitly consider the anisotropy of the underlying substrate texture. The wetting angle related to the placement of a liquid drop (say, of radius,  $R \sim 0.3$  cm) on a patterned surface (*e.g.*, with 20 µm (=*w*) ridges, with a  $\phi_I = 0.5$ ) would not be expected, through use of the *fakir* and the Wenzel models to be a function of the orientation of the ridges. However, experimentally we did notice a change, with a larger contact angle when the drop would be translated perpendicular to the ridges compared to an orientation parallel to the ridges: Figure 2.2. The two contact angles imply differing radii of curvature for the impregnated liquid parallel and perpendicular to the ridges<sup>1</sup>.



Figure 2.1 Schematic images of the wetting behavior, as indicated through the (a) Cassie-Baxter based fakir model, corresponding to the total absence of impregnation of the interstices by the liquid state, and the (b) Wenzel model, corresponding to the total impregnation by the liquid. I discuss here (c) the intermediate regime, corresponding to partial penetration of the interstices, (d) SEM image of the morphology of the ridged surfaces used for the experiments.

Much previous work on the anisotropy of wetting on textured surfaces<sup>14</sup> has not quantitatively estimated the penetration of liquid into interstices<sup>15</sup>. Measurements on sub-micron scale triangular grooves indicated<sup>16</sup> a state where the water completely penetrates the grooves. It was also discussed<sup>17</sup> that a water droplet may be suspended on patterns (modified with hydrophobic coatings<sup>18</sup>) and does not directly contact the substrate. Along with contact angle hysteresis<sup>19</sup>, a proposal that the contact angle decreases as penetration progresses was posited<sup>12</sup>. A related aspect of interest concerns the wetting dynamics, *e.g.*, in low Reynolds number (*Re*) flow. The *fakir* and the Wenzel models are often implicitly assumed in fluid flow<sup>20</sup> for understanding the efficacy of hydrophobic/super-hydrophobic surfaces in reducing fluid drag and inducing slip<sup>21</sup>, through reduced shear stress at the moving liquid-air interface. It is then the aim of this paper to examine the influence of anisotropy related to the substrate<sup>22</sup> on the wetting characteristics in static and quasi-static scenarios and lay out the physical rationale behind liquid penetration into the grooves.

#### 2.3 Methods

The tested Parylene-C coated patterned surfaces (on Si substrates) were patterned through standard photolithography. Briefly, coated photoresist on a Si wafer (*n*-type, <110>, and thickness of 500 µm) was patterned, developed and subject to reactive ion etching (RIE) to a trench depth of about 95 µm. The photoresist was then removed and the surface was coated with 1 µm thick Parylene-C. The ridged pattern surface morphology is shown in Figure 2.1(d). The average width (*w*) and height (*h*) was ~ 18 µm and ~ 95 µm, respectively, at a distance (*d*) of 18 µm. We define an interstice fraction:  $\phi_I$ , as the ratio [= d/(d + w)], with distances measured through scanning electron microscopy. Two different substrate patterns, lithographically designed for  $\phi_I = 0.5$  and  $\phi_I = 0.75$ , were used. To probe the related anisotropic wetting characteristics, a liquid drop (*i.e.*, 10 µL water, with  $\gamma_{LA}$  of 72.8 mJ/m<sup>2</sup>, and 3 µL oil: Krytox<sup>®</sup> GPL104<sup>23</sup>, with  $\gamma_{LA}$  of 18 mJ/m<sup>2</sup>) was placed on the Parylene-C coated patterned substrates (using a reported<sup>24</sup>  $\gamma_{SV}$  of 46.2 mJ/m<sup>2</sup>), and the contact angles observed in two orthogonal orientations at 20°C using Ramé-Hart Model 190 Contact Angle Goniometer. For comparison, the contact angle of water drops on the Parylene-C coated substrate (flat and unpatterned) was ~ 94.6<sup>0</sup>, with a corresponding  $\gamma_{SL}$  of 51 mJ/m<sup>2</sup> estimated through Eqn. 2.1(b). Detailed contact angle measurements as a function of orientation are indicated through Figure 2.2 and listed in Table 2.1.



Figure 2.2 Schematics of the top views of a water droplet, on a ridged surface, (a) parallel, and (b) perpendicular, to the ridges. (c) and (d) indicate the corresponding edge-on shapes, respectively.

**Table 2.1** The measurements of the contact angle, as a function of the orientation of the ridges. The values in the brackets indicate the corresponding  $\gamma_{SL}$  (=  $\gamma_{SL}$ , *mod*), in units of mJ/m<sup>2</sup>.

		$\phi_s = 0.5$	$\phi_{s} = 0.25$
	$\rightarrow$	111.3° (72)	124.9° (87)
	1	147.0° ( <i>107</i> )	148.5° ( <i>108</i> )

#### 2.4 Results and Discussions

We denote the *S*-*L* energies on the patterned surfaces as  $\gamma_{SL,mod}$ , as *modified* from the values obtained from those on unpatterned Parylene-C coated surfaces. The shapes of the droplets were broadly elliptical, with major axis along the ridge/groove, indicative of macroscopic preferential wetting in this direction. In all cases, the underlying pattern texture enhanced the degree of hydrophobicity, compared to the planar substrate, as indicated by an increase of the wetting angle to greater than 94.6<sup>o</sup>. It was noted that the contact angles on the patterned surfaces differ significantly, *e.g.*, by ~ 36<sup>o</sup>, (the difference between 147<sup>o</sup> in the perpendicular orientation *vs.* the  $111^0$  in the parallel direction) and correlated to the variation in the estimated  $\gamma_{SL}$  (=  $\gamma_{SL}$ , *mod*) from ~ 107 mJ/m<sup>2</sup> to ~ 72 mJ/m<sup>2</sup>.

Generally, when a liquid drop is placed on a patterned surface a small deflection/bowing ( $\delta$ ) of the liquid into the interstices (*I*) is expected based on gravitational considerations<sup>1</sup>: Figure 2.1(c). We consider the surface energy related to the liquid over the interstices as  $\gamma_{LI}$ , *e.g.*,  $\gamma_{LI} = \gamma_{LA}$ , if the interstice is mostly air (*A*). Here, for a liquid drop of radius *R*, and an interstitial length of *d*, from elementary geometrical considerations,  $\delta \sim d^2/8R$ . Such deflection is also favored on energetic considerations<sup>25</sup>, when  $\gamma_{LI}$  is smaller/comparable to the  $\gamma_{SL}$ . However, considering gravitational forces in addition to surface tension,  $\delta$  would be proportional to  $R^2/l_c^2$ , and the lateral spread ( $l_{spread}$ ) on the substrate would be proportional to  $R^2/l_c$  where  $l_c = \sqrt{\gamma/\rho g}$  is the capillary length, with  $\rho$  as the liquid density and *g* the acceleration due to gravity. Consequently, a larger  $\delta$  or  $l_{spread}$  would be favored by a larger/heavier drop and a smaller  $\gamma$ . Then, for a given drop size, the drop infiltrates into the interstice or spreads to lower the net  $\gamma$ . For a large liquid drop on top of a patterned surface, where R >> w and *d*, the net  $\gamma$ , could be formulated through:

$$\gamma_{SL, mod} = \phi_s \gamma_{SL} + \phi_I^{\prime} \gamma_{LI} \tag{2.3}$$

Here,  $\phi_I$  is related to the extent to which the liquid penetrates the interstice and may not necessarily be equal to the geometrically patterned  $\phi_I$ . While we designed our patterns: see Figure 2.1(d), with a  $\phi_I$  (= $\phi_s$ ) = 0.5, a smaller  $\phi_{LI}$  (/  $\phi_{LA}$ ) would implicate a larger  $\phi_I$  when other parameters in Eqn. 2.3 are fixed. The computed  $\gamma_{SL,mod}$  would then be a better metric to estimate the spreading (*S*) instead of the  $\gamma_{SL}$ .

A measure of drop spreading<sup>26</sup> could occur due to a very small pressure gradient, additional to the  $l_{spread}$ . When a liquid drop is moved *parallel* to the direction of the ridges, a given unit of the drop is constantly in contact with the underlying solid *or* interstitial region, during the entire trajectory of the motion. However, when the liquid drop is moved perpendicular to the direction of the ridges, the drop unit alternates contact between the underlying solid *and* the interstitial region. Consequently, in the latter case, the moving liquid drop does not benefit from penetrating further into the interstice, as in the former situation.

It would also be expected that a larger air fraction would increase the penetration and further decrease the extent of wetting on top of the patterned surface, manifesting a larger contact angle: Table 2.I. It is pertinent to note that that such impregnation of the liquid into the interstices yields an overall lower macroscopic wetting of the surface of the patterned substrate<sup>27</sup>.

Using Eqn. 2.3, with a computed  $\gamma_{SL, mod}$  of ~ 72 mJ/m<sup>2</sup> (from Table 2.1),  $\phi_s = 0.5$ ,  $\gamma_{SL}$  of 51 mJ/m<sup>2</sup>, and  $\gamma_{LI} \sim 72.8$  mJ/m<sup>2</sup>, we obtain a  $\phi_I^{-}$  of 0.64: Table 2.2(a). Alternately, with  $\gamma_{SL, mod}$  of ~ 107 mJ/m<sup>2</sup>, and  $\phi_s = 0.5$ ,  $\phi_I^{-}$  is ~ 1.12: Table 2.2(a). It is to be noted that as  $\gamma_{LI} > \gamma_{SL}$ , the liquid would be in intimate contact with the solid surface. Then, the  $\phi_s$  and the related  $\gamma_{SL}$  would not change and be equal to the geometrical/patterned value. The solid-liquid interfacial area could then be essentially considered flat. In both cases, the obtained  $\phi_I^{-}$  is greater than the patterned  $\phi_I$  of ~ 0.5. The corresponding  $\phi_I^{-}$  values, with a  $\phi_s$  (or a  $\phi_I$  of ~ 0.75) of ~ 0.25, were 1.02 and 1.31 – see Table 2.2(a), for the parallel and perpendicular orientation, respectively. Generally, spread of liquid into the groove would be expected to since  $\gamma_{SL}$  (~ 51 mJ/m<sup>2</sup>) is less than  $\gamma_{LI}$  (~ 73 mJ/m<sup>2</sup>); this would be in addition to the gravity induced penetration into the interstice.

**Table 2.2** The values of the fraction:  $\phi_I$ , corresponding to the extent to which the liquid penetrates the interstices of the ridged surfaces, for (a) water, and (b) oil, as estimated from Eqn. 2.3.

(a)			
		$\phi_s = 0.5$	$\phi_s = 0.25$
	$\rightarrow$		
		0.64	1.02
	ſ	1.12	1.31



Correlating the contact angle measurements in Table 2.1 and the corresponding  $\phi_l$  in Table 2.2(a), we can find that larger water penetration will result in larger  $\phi_l$  and contact angle, and the anisotropy of wetting angles can be attributed to the different degrees of water penetration in different directions. We also directly observe water penetration via confocal microscopy (using a Photron FASTCAM camera): Figure 2.3 and inset. The side-view images, on the bottom left and right of Figure 2.3(b) indicate both non-uniform and partial penetration of the liquid (red dotted lines), as well as full penetration into the interstices, respectively. Our studies preclude evaporation related considerations<sup>28</sup>.



**Figure 2.3 (a)** Confocal microscopy imaging of a water drop indicates a penetration of liquid into the interstices, of ~ 6  $\mu$ m as indicated in the inset. The red circle relates to the scan profile of the inset, where the vertical axis is related to the depth of the interstice, while the horizontal axis represents the distance along the interstices, above the patterned surface, (b) The side view of a water drop placed on the patterned substrate. The bottom left inset indicates the presence of water (partial penetration) in the interstices, through the reflecting contrast (from the white patches encompassed by the dotted red lines) and corresponding to the scan profile in the inset of (a). The bottom right inset indicates the infiltration of water (full penetration) into the interstices.

The obtained  $\gamma_{SL mod}$  values, together with the degree of penetration of the liquid into the interstices, are intermediate to the solid substrate surface energies (~ 46.2 mJ/m<sup>2</sup>) and the summation of the surface energies of the solid substrate and the liquid of ~ 119 mJ/m<sup>2</sup> (= 72.8 mJ/m<sup>2</sup> + 46.2 mJ/m<sup>2</sup>), where water was completely penetrating the interstices and tending to avoid the top solid substrate. The anisotropic tendencies in the underlying substrate modulate the extent of wetting/hydrophobic character.

We additionally tested the influence of anisotropy through the placement of Krytox<sup>®</sup> GPL104 oil on the patterned substrates, with a patterned  $\phi_s$  of 0.5. It was noted that the contact angles parallel and perpendicular to the ridges, was ~ 21.0° and ~ 30.1° respectively. For reference, the contact angle of the oil drop on the unpatterned Parylene-C coated surface was ~ 24.1°, implying a  $\gamma_{SL}$  of 29.6 mJ/m<sup>2</sup>, from Eqn. 2.1(b). While the smaller angles indicate a greater oleophilic character for the patterned surfaces, we estimate from Eqn. 2.3, correspondingly higher  $\phi_I$ ° of ~ 0.8 and ~ 0.9, respectively, as in Table 2.2(b) – indicating again an area enhancement of the oil in the interstice. Contrasting the time-dependent behavior of the water and the oil on the patterned surface: Figure 2.4, while the shape of water did not change much with time with little spreading, the oil drop spreads along and into the grooves quickly. A film propagates in advance of the oil drop indicating the enhanced oleophilic character due to the ridges.



**Figure 2.4** The time-dependent spreading of (a) water and (b) Krytox GPL104 oil, on the ridged surface. While the shape of the water drop was relatively stable, the oil drop spreads quickly into the grooves, with a film (outlined in a dotted red line) advancing ahead of the drop.

# **2.5 Conclusion**

Based on the results of our measurements and analyses, it is concluded that the anisotropy of wetting is related to the liquid penetration, where a larger effective air fraction implies larger penetration, yielding a larger contact angle. In most studies<sup>29</sup> related to the influence of patterned surfaces on inducing hydrophobic behavior, the degree of liquid penetration into the interstices has not been much considered<sup>6</sup>. The ratio of the estimated  $\phi_1^{T}$  to the  $\phi_1$  (which may be larger or smaller than unity) would enable the definition of a roughness metric, in contrast with the traditional<sup>1</sup> definition: Eqn. 2.2(b). We have indicated that a larger penetration, with a larger interfacial area/contact angle and lower shear would be of significance in modulating hydrophobic and surface slip behavior. The concomitant traversal direction dependent penetration of the liquid into the interstices may also be related to the system size as well as a shear-dependent effective slip length<sup>30</sup>. Related considerations would also be of importance in guiding liquid droplets, *e.g.*, in electrophoretic applications<sup>31</sup>, the design of drag-reducing super-hydrophobic surfaces<sup>6,21</sup>, *etc.* Future work would focus on the study of liquid flow dynamics as a function of the degree of anisotropy of the underlying patterned surfaces.

# 2.6 Acknowledgement

This chapter, in full, is a reprint of the materials as it appears in Fan, B. and Bandaru, P.R., Anisotropy in the hydrophobic and oleophobic characteristics of patterned surfaces, Appl. Phys. Lett, 111, 261603, 2017. The dissertation author was the primary investigator and author of this paper. The authors are grateful for support from the National Science Foundation (NSF: CMMI 1246800 and CBET 1606192). We also appreciate the assistance of Prof. James Friend, Shuai Zhang, and Dr. Jonathan Yu for help with imaging and confocal microscopy.

# Chapter 3 Enhanced voltage generation through electrolyte flow on liquidfilled surfaces

#### 3.1 Abstract

The generation of electrical voltage through the flow of an electrolyte over a charged surface may be used for energy transduction. Here, we show that enhanced electrical potential differences (i.e., streaming potential) may be obtained through the flow of salt water on liquid-filled surfaces that are infiltrated with a lower dielectric constant liquid, such as oil, to harness electrolyte slip and associated surface charge. A record-high figure of merit, in terms of the voltage generated per unit applied pressure, of 0.043 mV Pa<sup>-1</sup> is obtained through the use of the liquid-filled surfaces. In comparison with air-filled surfaces, the figure of merit associated with the liquid-filled surface increases by a factor of 1.4. These results lay the basis for innovative surface charge engineering methodology for the study of electrokinetic phenomena at the microscale, with possible application in new electrical power sources.

# **3.2 Introduction**

The motion of an aqueous electrolyte, such as salt water, over a surface may be harnessed for the generation of electrical voltage<sup>32</sup>. Indeed, the exploration of such electrokinetic phenomenon, and their possible use for energy conversion, have a long history, extending over the past two centuries (from Morrison & Osterle<sup>33</sup> and cited references). The relevant potential difference, termed as streaming potential  $(V_s)$ , arises due to the relative motion of charged<sup>34</sup> species in the electrolyte, with respect to the fluid channel substrate with residual charges<sup>31</sup>. While superhydrophobic (SH) surfaces<sup>6</sup> have been posited to increase the ion velocity and the resultant potential difference, such enhancement has not been observed to date, due to the inability of the air in the SH surfaces to hold charge<sup>35</sup>. Here, we monitor the flow and measure the  $V_s$  of higher dielectric constant ( $\varepsilon$ ) NaCl based electrolyte (consisting of Na<sup>+</sup> and Cl<sup>-</sup> dissolved in water) over a

lower  $\varepsilon$  solid substrate/surface with an induced negative charge<sup>2,36</sup>, in a microfluidic channel<sup>37</sup>. The electrically compensating positive counter-ions reside within the adjacent electrical double layer (EDL)<sup>38</sup>, which consists of an inner layer with fixed counter-ions adsorbed onto the surface and a diffuse layer with mobile counter-ions, with a thickness of the order of the Debye length ( $\lambda_D$ ). On application of a pressure difference ( $\Delta P$ ) across the two ends of the channel, say, through a mechanical pump: Figs. 3.1(a) - (c), the electrolyte flow would be mainly constituted from the mobile counter-ions in the diffuse layer, with the consequent charge separation yielding a streaming potential ( $V_s$ ), proportional to the  $\varepsilon$  ( $=\varepsilon_0\varepsilon_r -$  with  $\varepsilon_0=$  8.854. 10<sup>-12</sup> C<sup>2</sup> N<sup>-1</sup> m<sup>-2</sup> as the free space permittivity, and  $\varepsilon_r$  as the relative permittivity of electrolyte *e.g.*, ~ 80 for 0.1 mM NaCl solution), the zeta potential ( $\zeta$ ) at the edge of the shear plane where the mobile ion motion occurs, and varying inversely with the dynamic viscosity of the electrolyte ( $\eta$ ), and bulk electrolyte conductivity ( $\kappa$ ), as described through the Helmholtz-Smoluchowski model<sup>8,36</sup>:

$$V_S = \frac{\epsilon \zeta}{\eta \kappa} \Delta P \tag{3.1}$$

The  $\zeta$  is considered close to the substrate – electrolyte liquid interface, and is significant in that it determines the electrical potential that may be utilized<sup>2,36</sup> for the  $V_s$ . In the flowing electrolyte itself, there is a close to exponential decay of the surface potential over<sup>38</sup>  $\lambda_D$ . It should be noted that Eqn. 3.1 is based<sup>39</sup> on the assumptions of negligible surface (/substrate) conductivity and a very small EDL thickness<sup>25</sup>, with Poiseuille flow of the electrolyte. Moreover, there is an implicit assumption of the *no-slip* boundary condition<sup>40</sup>, where liquid adjacent to the substrate wall has a zero velocity with a finite flow velocity only at a certain distance (corresponding to the shear plane) into the fluid. Consequently, the  $\zeta$  would be related to the electrical potential at the edge of the shear plane in electrokinetic flows over smooth surfaces<sup>36,39</sup>. Most work on harnessing the  $V_s$  through electrokinetic effects, to date, has been concerned with fluid flow over smooth surfaces (where the scale of roughness is smaller than  $\lambda_D$ ), and consequently very small potential differences of the order of 18 mV may be predicted and obtained<sup>8</sup>, in correspondence to Eqn. (3.1), *i.e.*, for 0.1 mM NaCl, with  $\varepsilon_r \sim 80$ ,  $\eta \sim 10^{-3}$  Pa · s,  $\zeta \sim 25$  mV,  $\kappa \sim 10^{-3}$  S m<sup>-1</sup>, with  $\Delta P \sim 1000$  Pa.



**Figure 3.1 Electrokinetics over patterned and fluid filled surfaces.** The phenomena were investigated (a) through monitoring the streaming potential ( $V_s$ ) of salt water, under pressure driven flow, in a microchannel (of length 11.8 cm, width of 0.9 cm and height of ~250 µm). The upper surface of the channel was PDMS (poly-dimethyl siloxane) while the lower fluid-solid surface, was engineered with the fluid (f) such as oil or air. The velocity (u) profile with fluid slip, due to the pressure difference ( $P_2 - P_1$ ) applied by syringe pump is indicated, (**b**) The schematic of the flow arrangement in the microchannel and its (**c**) experimental realization, wherein the top and the bottom surfaces were separated by a silicone rubber spacer to adjust the microchannel height, and Ag/AgCl electrodes were inserted in reservoirs at either end to measure the  $V_s$ . While (**d**) liquid flow over an *AFS* (air filled surface) yields a finite slip velocity ( $u_s$ ), and slip length (b) the (**e**) flow of an electrolyte such as salt water yields a finite  $V_s$  due to surface charge (indicated as negative, here) induced influence on the electrolyte ions. (**f**) The  $V_s$  can be made significantly larger when the air is replaced by a water-immiscible liquid, such as oil yielding a liquid filled surface (*LFS*).

With the objective of obtaining significantly larger  $V_s$ , we indicate briefly the principles of our approach, which first involved the modulation of the effective  $\zeta$ , through introducing fluid slip via groove patterned surfaces (air/liquid filled surface (*AFS/LFS*)) and modifying surface charges at the substrate surface. It has previously been considered, based on molecular dynamics based simulations<sup>41</sup>, that slip may mobilize the Stern layer, significantly enhancing the  $\zeta$ . The broad concept is that the shear plane/surface of shear (at which the electrolyte flow velocity is zero) is to be moved as close as possible to the substrate. As the surface electrical potential decays away from the substrate, the closer the shear plane is to the substrate boundary the larger the  $\zeta$  with concomitantly increased  $V_s$ . Here, the zero velocity boundary condition at the surface (*y*=0) would be replaced with a Navier slip condition:  $u_s (y=0)=b \frac{\partial u (y=0)}{\partial y}$ ) with *b* as the slip length<sup>20,42</sup>,  $u_s$  as the slip velocity: Fig. 3.1(d). The larger the *b*, the greater the fluid velocity at the surface by a factor of  $(1 + \frac{b}{\lambda_D})$ . A related apparent increase<sup>43</sup> in the  $\zeta$  and the  $V_s$  may then be consequently obtained<sup>44</sup>. As much relatively large values of *b* of the order of 10 µm were thought to be obtained through the use of superhydrophobic (SH) surfaces<sup>10</sup>, and given a  $\lambda_D$  of 9.6 nm with 1 mM NaCl, a  $b/\lambda_D$  of ~ 1000 is estimated, yielding large  $V_s$ . While non-uniform surface conduction<sup>45</sup>, *e.g.*, as related to slip *to* no-slip transitions, may reduce such an enhancement a finite *b* would yet contribute to  $V_s$ , and may be aided by surface charge induced forces.

Our experiment results show that the  $V_s$  of NaCl electrolyte over *LFS* is increased by 1.4 times compared with that of electrolyte over *AFS*, which is due to the harness of both the slip and surface charge at liquid-liquid interface.

#### 3.3 Methods

Design considerations and fabrication of air filled surfaces and liquid filled surfaces. The critical parameters in our channel surface design are (a) structural, *i.e.*, the interstitial fraction of *AFS/LFS*:  $\phi$  - the ratio [= d/(d + w)], with an average ridge width (*w*) and spacing (*d*), as indicated in Fig. 3.2(a), (b) dimension, the channel length (*L*) and channel width (*W*) should be much larger than channel height (*H*), then the channel can be treated as two infinite parallel plates and the flow is Poiseuille flow, (c) pressure gradients, *i.e.*, over a range of 0-1000 Pa, with higher pressures yielding greater streaming potentials (*V<sub>s</sub>*), and (d) the nature of the filling fluid, easier for oil with low surface tension to penetrate into the grooves. For instance, we have used in our electrokinetic flow experiments,  $\phi \sim 0.5$ , 0.9 cm wide, 11.8 cm long and 250 µm high channel, with a pressure difference of ~ 1200 Pa, using GPL oil as a filling liquid, to generate maximal *V<sub>s</sub>*.

The *LFS* were fabricated through filling oil into the grooves of a patterned surface, fabricated through photolithography and dry-etching. A Si wafer (n-type, <110>, thickness 500 µm) was cleaned with acetone and IPA, rinsed by DI water, and subsequently baked at 180 °C for 5 minutes. Negative photoresist

(NR9-3000) was coated on the Si wafer (3000 rpm for 40 s) and baked at 120 °C for 60 s. The pattern was defined through a mask (using the EVG 620) and developed (RD6) for 1 minute. Then the resist patterned wafer was subject to dry etching (using Oxford Plasmalab 100 RIE/ICP) to yield a trench depth (h) ~ 95  $\mu$ m. The photoresist was then removed (using RR2 photoresist remover) over ~ 12 hours and any residues was further removed (using PVA TePla PS100 at 120 sccm, 200 W for 90 s). The grooved surface was coated with Parylene-C (using PDS 2010 Parylene Coater) to a thickness of ~600 nm.

For *LFS*, oils, *e.g.*, Dupont Krytox GPL 104, Castor oil, *etc.* was filled into the grooves of the patterned surface. The air-/liquid- patterned surfaces were imaged using environmental scanning electron microscopy (ESEM): FEI/Phillips XL ESEM and the FEI Quanta FEG 250 ESEM. We have indicated, *e.g.*, in Figure 3.2(c) that a lower  $\phi$  - the ratio of the average groove width to the overall period, yields a larger *V<sub>s</sub>*.

Choice of applied pressure and related flow rate. The electrolyte flow rate in the channel, constituted from rectangular plate geometry<sup>46</sup>, was modulated through a syringe pump: Figure 3.1(c), and was transduced to an applied pressure ( $\Delta P$ ) based on the Poiseuille relation:  $\Delta P = \frac{12\eta LQ}{wh^3}$  (*Q* is flow rate, *L* is channel length of ~ 11.8 cm, and *h* the channel height of ~ 250 µm). The  $\Delta P$  over the channel length (where the other end was left open to atmosphere) motivated the pressure gradient driven flow, and for the experiment was in the range of 200 to1200 Pa, as measured through a manometer (UEI EM152 Dual Differential Input Manometer). It was observed that when the  $\Delta P$  was larger (/smaller) than 1200 Pa (/200 Pa), that the generated streaming potential *V<sub>s</sub>* was not stable.

**Choice of liquid to fill the interstices in the liquid filled surfaces.** For the fabrication of the *LFS*, the filling liquid (*e.g.*, Krytox<sup>®</sup> GPL 104 and castor oil) was chosen primarily on the basis of immiscibility with water/electrolyte. Additional criteria would be a low surface tension - for better penetration into the interstices, low viscosity and a smaller dielectric constant – to reduce electrolyte-oil interfacial shear stress. We observe that the parylene coated Si surface was oleophilic (/hydrophobic), and that the chosen liquid or oil could penetrate the interstices easily: Figure 3.3(a).

#### **3.4 Results**

**Voltage generation on air filled surfaces.** To further understand the issues related to electrokinetics over surfaces with slip, we conducted experiments - see Figs. 3.1(b) and 1(c), where the electrolyte flow was driven in a microfluidic channel (of length: 11.8 cm, width: 0.9 cm and height: 250  $\mu$ m) by a syringe pump with  $\Delta P$  varied in the range of 0 - 1200 Pa to measure the  $V_s$  over ridged pattern surface morphology: termed as *AFS* (air filled surface): Fig. 3.1(e). The substrate material of *AFS* is silicon, coated by 600 nm hydrophobic parylene-C. The patterns were characterized through an interstitial/air fraction:  $\phi_{air}$ , through the ratio [= d/(d + w)], with an average ridge width (w) and spacing (d): Fig. 3.2(a). We report results on samples with  $\phi_{air} = 0.5$ ,  $d = w = 18 \ \mu$ m, for  $\phi_{air} = 0.75$ ,  $d = 27 \ \mu$ m and  $w = 9 \ \mu$ m.



**Figure 3.2 Streaming potential of air-filled surfaces. (a)** A scanning electron microscope (SEM) micrograph of the grooved pattern surfaces with silicon as the substrate covered by parylene-C (thickness  $\approx 600$  nm), parameterized on the basis of the solid surface width ( $w = 18 \mu$ m), groove width ( $d = 18 \mu$ m), and pattern height ( $h \approx 95 \mu$ m), and the air fraction:  $\phi_{air} [= d/(d + w) = 0.5]$ , and a solid fraction:  $\phi_s = 1$ - $\phi_{air}$ . The flow direction is perpendicular to the grooves and indicated by the arrow. Scale bar is 100  $\mu$ m. (b) The typical response of streaming potential ( $V_s$ ) was measured, for a  $\phi_s = 0.5 AFS$ , through a syringe pump driven pressure, *e.g.*, with an applied pressure of 440 Pa, 0.1 mM L<sup>-1</sup>NaCl here. The red dotted line indicated the average flow velocity in the channel. The measured voltage difference between a baseline value ( $V_o$ ) and a final value ( $V_f$ ) was considered as the  $V_s$ . (c) The measured  $V_s$  (0.1 mM NaCl solution) on an AFS (*i.e.*, Air\_{0.5}, the subscript indicating the  $\phi_{air}$ ) scales linearly with the applied pressure (in the range of 200-1200 Pa), in accord with the Helmholtz-Smoluchowski model (Eqn. 3.1) with a lower value for  $V_s$  obtained for a smaller (/larger)  $\phi_s$  ( $/\phi_{air}$ ) and rationalized as due to the absence of surface charges, and contributing zeta potential  $\zeta$ , in the air regions. (d) The  $V_s$  scales with the salt water electrolyte conductivity and related concentration ( $I_c$ ) through a log [ $I_c$ ]/ ( $I_c$ ] variation for both AFS (Air\_{0.5}) and parylene coated flat surface (*Flat*) (concentration varying from 0.1mM L<sup>-1</sup> to 5 mM L<sup>-1</sup> at 200 Pa for (d)).

The consequent voltage-time trace: Fig. 3.2(b), was used to determine the  $V_s$  – monitored through a high resistance (>10 GΩ) voltmeter (Keithley 2700) connected across the Ag/AgCl electrodes placed at either ends of the channel: Fig. 3.1(c). However, a significant enhancement was not obtained for *AFS*, with an estimated<sup>6</sup> *b* of ~ 10 µm. It was interesting to note that a lower  $V_s$  was obtained for a larger  $\phi_{air}$  (*i.e.*, smaller  $\phi_s$ ) for which a larger *b* would be expected: Fig. 3.2(c). Moreover, very little increase over a flat unpatterned surface was obtained comparing *Air*<sub>0.5</sub> and *Flat*: Fig. 3.2(d). The lower  $V_s$  of *AFS* may be rationalized as due to the absence of a surface charge in the air regions<sup>25</sup>, which overwhelms the contribution of the slip.

While the data here seems to indicate the general validity of Eqn. 3.1, in terms of a linear  $V_s - \Delta P$ plot, the slope of the obtained experimental curves would then be directly connected to the  $\varepsilon$ ,  $\zeta$ ,  $\eta$ , and the  $\kappa$ . Such parameters are then intimately connected, *i.e.*, while independent variation would be difficult to discern experimentally, theoretical or computational input (which is outside the purview of the present experimental work) may be able to yield guidance for explicit dependence. The net  $V_s$  has been construed to be related to a weighted summation of  $\zeta$  in the no-slip region (over the solid fraction of the surface, *i.e.*,  $\zeta_{NS}$ ) and the slip region (*i.e.*, over the air of the *AFS*, *i.e.*,  $\zeta_S$ ) <sup>35</sup>. The very small  $\zeta_S$  would adversely affect the  $V_s$  and as seen through our experiments, overwhelms the slip length related contribution. Related correspondence of the obtained  $V_s$  to the  $\zeta$  as well as the electrolyte conductivity, which is proportional to  $I_c$  (the counter-ion concentration) was determined through a log  $[I_c]/[I_c]$  dependence: Fig. 3.2(d).

Increased voltage on liquid filled surfaces. Subsequently, we deployed patterned surfaces where the air pockets were infiltrated by a water immiscible liquid, *e.g.*, oil, of a higher density compared to the NaCl electrolyte: Fig. 3.3(a). Such liquid filled surfaces (*LFS*) were hypothesized to yield a finite  $\zeta_S$  at the oil interfaces, in addition to a non-zero *b*. The *LFS* were fabricated through replacing the air in groove patterned arrays by water-immiscible liquid, such as oil, and may be designed to exhibit various degrees of fluid slip<sup>27,47,48</sup>, as well as electrical voltages,– significantly larger than on air-filled or SH surfaces. The oil filling was found to be reliable and stable.


**Figure 3.3 Enhanced streaming potential at liquid filled surfaces.** Enhanced streaming potential ( $V_s$ ) was obtained by (**a**) filling the grooves with salt water immiscible oil, *e.g.*, Krytox GPL 104, (scale bar is 100 µm) (**b**) a significantly larger  $V_s$  was obtained for an GPL 104 oil filled surface (*i.e.*, *GPL*<sub>0.5</sub>, with the subscript now indicating the interstitial fraction of oil), compared to one with air ( $Air_{0.5}$ ) and even compared to a *Flat* solid substrate (all measured using 0.1 mM L<sup>-1</sup> NaCl solution with applied pressure as 1200 Pa). (**c**) The  $V_s$  could be larger or smaller compared to the control *AFS*, depending on the choice of the liquid in the *LFS*, *e.g.*, GPL oil and castor oil (*Castor*<sub>0.5</sub>), respectively.

Additionally, the use of oil with a lower  $\varepsilon_r$  compared to the flowing electrolyte would mean that the electric field from the wall is effectively propagated. Through the use of Krytox GPL 104 oil ( $\varepsilon_r \sim 2.1$ ,  $\eta \sim 340$  cP), a significant enhancement of the  $V_s$  by 50%, to around ~ 52 mV, compared to that of *Air*<sub>0.5</sub> was observed: Fig. 3.3(b), attesting to the utility of the *LFS*. However, it was noted that the obtained  $V_s$  critically depends on the choice of the oil in the *LFS*. Broadly, a larger  $\varepsilon_{oil}$  seems to yield a lower  $V_s$ . For instance, a factor of two smaller value was obtained for a castor oil ( $\varepsilon_r \sim 4.7$ ,  $\eta \sim 312$  cP) filled *LFS*: Fig. 3.3(c). The enhanced van der Waals energy of interaction<sup>1</sup> between the oil and the electrolyte, proportional to the dielectric constants, and the related mutual electric polarizabilities causes a larger friction. The consequent movement of the shear plane away from the solid substrate would result in a lower zeta potential and  $V_s$ .

The electrolyte flow over the *LFS* as well as the adjacent solid surface was parameterized<sup>46</sup> through the use of a material dependent surface charge density ( $\sigma$ ) related to  $\zeta = \frac{\lambda_D \sigma}{\epsilon}$  and fluid slip at the respective interfaces. From literatures, for the top surface of the channel (constituted from poly-dimethyl siloxane: PDMS),  $\sigma_{PDMS}^{49} \sim 18$  mC m<sup>-2</sup>, while for the bottom patterned surfaces, we use  $\sigma_{oil}^{50} \sim 1.8$  mC m<sup>-2</sup> and  $\sigma_{parylene}^{51} \sim 3.6$  mC m<sup>-2</sup>. The  $\sigma$  and the velocity slip is inhomogeneous along the length of the channel, but constant along the channel width. The no-slip hydrodynamic boundary condition was assumed to hold true at all solid-electrolyte interfaces, while a Navier slip boundary condition over the flowing electrolyte-oil interface was assumed, with a finite slip velocity:  $u_s$  and slip length: *b*. For a hydrophobic surface, the *b* may be approximated<sup>6</sup> to be of the order of *d*, *i.e.*, ~ 18 µm. A correspondence to a Cassie state-like channel flow (with no electrolyte penetration into the grooves of the patterned surface) for *LFS* surface may be seen. It should be noted that simulations considering the continuity of tangential shear stress across the electrolyte-oil interface also obtained the  $V_s$  very close (within 1 %) to the results obtained through an assumption of a slip length.

**Modeling of electrolyte flow over the liquid filled surfaces.** We partition<sup>39</sup> the pressure gradient driven electrolyte flow, following experiment, over the *LFS* as the superposition of (i) hydrodynamic flow, with fluid slip over electrically neutral surfaces, and (ii) an electrokinetic flow, with no-slip fluid flow over flat electrically charged surfaces: Figs. 3.4(a) – (c). A multiphysics model coupling the Nernst-Planck-Poisson (N-P-P) equation with the Stokes Equation, was deployed to determine the volumetric charge density profiles. The velocity profile combined with the simulated volumetric charge density profile along the channel, as indicated in Fig. 3.4 (d), arises due to the difference of ionic concentrations between the counter-ions and co-ions. The resultant horizontal electric field, is responsible for the observed streaming potential (*V<sub>s</sub>*). The calculated average electrolyte velocity of 6 cm s<sup>-1</sup> is in agreement with that expected from Poiseuille flow along the channel, with the estimated *u<sub>s</sub>* at the oil-electrolyte interface of ~ 2 cm s<sup>-1</sup> at  $\Delta P \sim 1200$  Pa. Additionally, and given the relatively large groove widths, the influence of the surface conductivity on the obtained results would be relatively small<sup>44</sup>.



Figure 3.4 Parametrization and modeling of charged fluid-solid surfaces. (a) The surface charge density ( $\sigma$ ) of an *AFS* (air filled surfaces) varies between zero (at the air) and a finite value at the solid, while a nominally planar *LFS* (liquid filled surfaces) maintains a larger value of the  $\sigma$ . The electrokinetics on the *LFS* was considered in terms of a superposition of (b) hydrodynamic flow, with fluid slip over electrically neutral surfaces and (c) an electrokinetic flow, with no-slip fluid flow over electrically charged surfaces, and solving the coupled Nernst-Planck-Poisson (NPP) equation along with the Stokes Equation, to determine the (**d**) velocity profile as well as the volumetric charge density ( $\rho$ ) profiles. The insets indicate the computed variation of the  $\rho$  at the top PDMS surface and the bottom *LFS*.

#### **3.5 Discussion**

Excellent agreement of the simulations with experimentally observed values of the streaming potential was observed through considering a reduced  $\varepsilon_r \sim 28$ , over a distance of  $\sim 3\lambda_D$  proximate to the *LFS* and an  $\varepsilon_r \sim 78$  beyond - correspondent to the bulk solution. Such aspect is in accord with the use of different values of the dielectric constant for the inner and outer regions of a solid -solution interface<sup>39</sup>. The *LFS*, consisting of both oil as well as solid surfaces, may yield an uneven shear plane with fascinating implications for novel electrokinetic phenomena<sup>52,53</sup>, such as localized concentration polarization<sup>44</sup>, *etc*. Our work also indicates that a heterogeneous surface could be patterned through the use of discrete surface charge density or applied potentials, mimicking a superhydrophobic surface, and making a connection with electro-wetting applications, as well as with significant literature on the modeling and parameterization of liquid flows on striped surfaces exhibiting alternate regions of slip and no-slip, with<sup>54,55</sup> and without<sup>30</sup> surface charge.

A more detailed investigation of several issues related to the interplay of enhanced surface charge as well as increased slip velocity at the flowing electrolyte - *LFS* interface would then be warranted. It has

been indicated, for instance, that an increased  $V_s$  would result due to the larger ion convection currents<sup>56</sup>. Additionally, viscous effects may be playing a major role<sup>57,58</sup>. As the coefficient in Eqn. (3.1), intimately connects the  $\varepsilon_r$ , and the  $\eta$ , a decrease in the former may not be readily deconvoluted from an increase in the latter experimentally. In our numerical simulations, an increase in the  $\eta$  by a factor of ~ 2.5 over the equivalent length scale (from a bulk value of 0.89 cP to 2.19 cP) was necessary to obtain the experimentally observed streaming potentials, assuming that the  $\varepsilon_r$  is ~ 78.

Our experiments have deep scientific implications underlying fluid flow interfacing with both air and liquid as well as fluid shear at a surface, and electrokinetic phenomena as related to the localized variation of the zeta potential and nonlinearity. We anticipate that our work would revitalize research related to the fabrication of alternate electrical voltage/power sources from liquid flow over charged surfaces. Other anticipated applications extend to electrophoretic applications in biological separations (cell transport, manipulation<sup>59</sup>, and interactions<sup>60</sup>), as well as voltage sources for lab-on-a-chip applications<sup>61</sup>.

#### **3.6 Conclusion**

In summary, our work has experimentally demonstrated the largest figure of merit thus far<sup>62–65</sup> to the best of our knowledge, with primary focus on methodologies related to enhance the streaming potential  $(V_s)$  per unit pressure difference  $(\Delta P)$ , through the use of liquid filled surfaces (also see Supplementary Note 6 for considerations related to overall electrokinetic efficiency). The use of the *LFS* yields a factor of 1.4 larger figure of merit compared to that obtained through the use of the *AFS*. It has been shown that larger voltages, through a measured streaming potential, may be achieved through careful engineering of the coupled electric field and fluid flow. The application of the related increase in the electrokinetic energy conversion efficiency would need further optimization of the fluidic and electrical impedances, in concert with the streaming conductance, as matched to an appropriate load<sup>66</sup>. Concomitantly, unipolar transport (where for example, either Na<sup>+</sup> or Cl<sup>-</sup> ions are transported<sup>67</sup>) through electrical double layer overlap<sup>46,68</sup> in nanoscale channels, may be coupled with LFS to yield much larger voltages, comparable to that of batteries<sup>69</sup>.

#### 3.7 Acknowledgement

This chapter, in full, is a reprint of the materials as it appears in Fan, B., Bhattacharya, A. and Bandaru, P.R., Enhanced voltage generation through electrolyte flow on liquid-filled surfaces, Nature Communications, 9, 4050, 2018. The dissertation author was the primary investigator and author of this paper. The authors are grateful for support from the National Science Foundation (NSF: CMMI 1246800 and CBET 1606192). We also appreciate the assistance of Prof. J. Friend and P. Chen for help with imaging and Dr. S. Rubin for discussions.

#### 3.8 Supplementary materials

## **3.8.1** Geometrical parameters for the construction of air filled surfaces (*AFS*) and liquid filled surfaces (*LFS*):

For the influence of surface geometry alone, we compare the results of *Flat* (unpatterned surface), and the air-filled surfaces (*AFS*): *Air*<sub>0.5</sub> and *Air*<sub>0.75</sub>, as indicated in Figures 3.2(c) and (d). The air in the *AFS* can reduce the friction between the flowing liquid electrolyte and patterned surface which helps in the increase of the streaming potential ( $V_s$ ). However, a non-charged liquid-air interface will have no contribution to streaming potential<sup>34,35</sup>. Consequently, the  $V_s$  of *Air*<sub>0.5</sub> is only a little larger than compared to the *Flat* case and the  $V_s$  of *Air*<sub>0.75</sub> is even smaller, presumably due to an increase of non-charged liquid-air areas.

The aspect of nano- and micro-scale roughness was considered and found to be of much less importance compared to the groove width. The nominal roughness of the parylene coated surface was found to be  $R_a = 1.25$  nm  $\pm 0.19$  nm (as determined through Dektak 150 Surface Profilometer) and the obtained

streaming potential was insensitive to variations around such values. Generally, a roughness of < 6 nm has minimal effects on the slip length and flow rate<sup>70</sup>, and can be considered hydrodynamically smooth<sup>71</sup>.



**Supplementary Figure 1** The contact angle of (a) water, (b) GPL 104, and (c) castor oil on parylene coated flat Silicon surface were measured using Ramé-Hart Model 190 Contact Angle Goniometer. The measured results are 94.6°, 24.1°, 28.8° for water, GPL 104 and castor oil, respectively. It is then clear that parylene is hydrophobic (/oleophilic). (d) shows that the GPL 104 oil as well as the castor oil is immiscible with water.

Supplementary Table 1 | Physical properties of filled fluids (*data from* DuPont<sup>®</sup> Performance Lubricants, 2015)

Oils	Density (g/cm³)	Kinematic Viscosity (cm²/s)	Dynamic Viscosity (mPa·s)	Dielectric Constant	Surface Tension (mN/m)
Krytox <sup>®</sup> 104	1.93	1.77	340	2.1	18
Molivera Organics	0.96	3.25	312	4.7	39
Castor Oil					

#### 3.8.2 The pressure dependence of $V_s$ and the numerical estimation of the surface potential ( $\zeta$ )

In the chosen pressure range, that the  $V_s$  was linearly proportional to the applied  $\Delta P$ , following the Helmholtz-Smoluchowski relation:  $V_s = \frac{\epsilon \zeta}{\eta \kappa} \Delta P$ , as shown in Figures 3.2(c), 3.3(b), and 3.3(c). From the

slope of the  $V_s - \Delta P$  curve, we estimate the surface potential ( $\zeta$ ) at the edge of the shear plane, assuming an the electrolyte permittivity:  $\varepsilon (= \varepsilon_o \varepsilon_r)$  with  $\varepsilon_r$  as the relative permittivity, *e.g.*, ~ 80 for 0.1 mM NaCl solution),  $\eta$ : the dynamic viscosity of the electrolyte (~ 10<sup>-3</sup> Pa·s), and a bulk electrolyte conductivity ( $\kappa \sim$ 10<sup>-3</sup> S/m), which are listed in Supplementary Table 2, below.

Sample	R <sup>2</sup>	Slope (mV/Pa)	ζ (mV)	
Flat	0.9972	0.03081	44.2	
<i>Air</i> <sub>0.5</sub>	0.9997	0.03117	46.1	
<i>Air</i> <sub>0.75</sub>	0.9963	0.0286	41.0	
$GPL_{0.5}$	0.9996	0.04392	65.6	
Castor <sub>0.5</sub>	0.9963	0.02418	34.7	

Supplementary Table 2 | Linear fitting results and ζ

#### 3.8.3 Estimates of the overall slip length ( $b_{eff}$ ) from averaged surface potential ( $\zeta$ )

The  $\zeta$  values listed in Supplementary Table 2, may be considered an averaged/effective value(=  $\zeta_{eff}$ ) considering the top (*PDMS*) and bottom (the *LFS*, constituted from the *parylene* and the *oil* in the *interstices*) surfaces of the channel. Based on the effective medium approach (EMA)<sup>35</sup>, we consider a weighted summation of the  $\zeta_{PDMS}$ ,  $\zeta_{parylene}$ , and  $\zeta_{interstice}$  as follows:

$$\zeta_{eff} = 0.5\zeta_{PDMS} + 0.5\zeta_{parylene} \tag{S3-1}$$

$$\zeta_{eff} = 0.5\zeta_{PDMS} + \frac{b_{eff}}{\lambda_D} \left( 0.25\zeta_{parylene} + 0.25\zeta_{interstice} \right)$$
(S3-2)

The reported  $\zeta_{parylene}$  is 17.5 mV at 10 mM NaCl solution<sup>72</sup>, consequently  $\zeta_{parylene}$  was estimated to be ~ 35 mV for the 0.1 mM electrolyte used in our studies, based on the relation<sup>36</sup>  $\zeta$ ~ log[*I*]. The  $\zeta_{PDMS}$  was estimated to be ~ 53.4 mV based on the first equation using  $\zeta_{eff} = 44.2$  mV for *Flat*. Such an estimate for the  $\zeta_{PDMS}$  seems to be in reasonable accord with literature, considering that there may be many affecting factors<sup>36,73</sup>, *e.g.*, counter-ion type, pH of the solution and temperature. The  $\zeta_{interstice} = 0$  mV for air<sup>35</sup>. Then, the  $b_{eff}$  for  $Air_{0.5}$  can be estimated as 66.5 nm based on the second equation. Given that<sup>74</sup> the  $\zeta_{oil}$  is ~ 69 mV, the effective slip length  $b_{eff}$  for the  $GPL_{0.5}$  : would be ~ 44.9 nm. The  $b_{eff}$  can also be estimated for  $Air_{0.25}$  and  $Castor_{0.5}$  using the same method and all the results are listed in Supplementary Table 3.

Sample	<i>b<sub>eff</sub></i> (nm)
Air <sub>0.5</sub>	66.5
<i>Air</i> <sub>0.25</sub>	98.0
$GPL_{0.5}$	44.9
Castor <sub>0.5</sub>	9.2

Supplementary Table 3 | Estimates of the *effective* slip length: b<sub>eff</sub>

The  $b_{eff}$  of  $Air_{0.75}$  is larger compared to that of  $Air_{0.5}$  and maybe related to the larger amount of air, in the former. The  $b_{eff}$  of *GPL oil* is smaller due to a larger viscosity, compared to air, while the  $b_{eff}$  of *castor oil* is even smaller due to a larger dielectric constant.

While significant care was taken in removing the oil from the surface in the use of the *LFS*, there is a possibility that the oil may partially cover the parylene surface. Consequently, the effective medium approach (EMA) aspect may indeed be considered an approximation. Previously, we used  $\phi_{parylene} = 0.25$  and  $\phi_{interstice} = 0.25$  (assuming that parylene area was not covered by oil) and obtained an effective slip length ( $b_{eff}$ ) of ~ 44.9 nm for *GPL*<sub>0.5</sub>. However, even with  $\phi_{parylene} = 0.1$  and  $\phi_{interstice} = 0.4$  (assuming that 80% of the LFS was covered by oil), the  $b_{eff}$  is ~ 37.5 nm, a change of ~ 16%. Consequently, the use of the EMA seems to be a reasonable approximation.

#### 3.8.4 Pressure-driven electrokinetic flow with no hydrodynamic slip

To describe electrolyte flow, we modeled through self-consistent finite element simulations (COMSOL<sup>®</sup>), the local fluid velocity at a point (u), the respective ionic species concentration ( $c_i$ ), and

electric potential ( $\phi$ ). The relevant mass conservation equations, and the Stokes equation were coupled with the Nernst-Planck-Poisson (*N-P-P*) models, as described below.

Let i=1, 2 represent the ionic species present in the electrolyte, *e.g.*, i=1 refers to Na<sup>+</sup>, while i=2, refers to the Cl<sup>-</sup> species. The flux conservation equation<sup>75</sup> of each ionic species, *i*, is

$$\frac{\partial n_i}{\partial t} = \nabla \cdot \boldsymbol{J}_i \tag{S4-1}$$

where  $J_i$  is the ionic flux vector and  $n_i$  is the number density of ions per unit volume.  $n_i = c_i * N_{Av}$  where  $c_i$  is the concentration of each ionic species in moles/L (M) and  $N_{Av}$  is the Avogadro constant = 6.02 x10<sup>23</sup> /mole. For steady state flow:  $\nabla \cdot J_i = 0$ .

The Nernst-Planck (*N-P*) equation<sup>32,39,76</sup>: Eqn. (S4-2) below, considers the motion of ions in a fluid as a function of fluxes due to convection, diffusion, as well as ion migration under the effects of electrostatic forces.

$$\boldsymbol{J}_{\boldsymbol{i}} = n_{\boldsymbol{i}}\boldsymbol{u} - D_{\boldsymbol{i}}\nabla n_{\boldsymbol{i}} - \frac{z_{\boldsymbol{i}}e \, n_{\boldsymbol{i}}D_{\boldsymbol{i}}}{k_{B}T}\nabla \,\boldsymbol{\phi}$$
(S4-2)

 $D_i$  is the respective species diffusivity, *i.e.*,  $D_{Na^+} = 1.13 \ge 10^{-9} \le m^2 \le and D_{Cl^-} = 1.0 \ge 10^{-9} \le z_i$  is the valence of each ionic species.  $z_{Na^+} = 1$  and  $z_{Cl^-} = -1$ ,  $k_B$  is the Boltzmann constant = 1.38 \xeta 10^{-23} J/K and T = 298 K. The *u* is obtained through the Stokes' equation<sup>40</sup>: Eqn. (S4-3),

$$-\nabla p + \mu \nabla^2 \boldsymbol{u} + \rho_f \nabla \phi = 0 \tag{S4-3}$$

Here, pressure gradient driving the flow  $(\nabla p)$ , the viscous force  $(\mu \nabla^2 u)$ , with  $\mu$  as the dynamic viscosity) and the electrical body force  $(=\rho_f \nabla \phi)$ , with  $\rho_f$  as the volumetric space charge density in the electrolyte due to the presence of free charges, *e.g.*, counter-ions near the surface, are all considered. The electric potential ( $\phi$ ) is obtained from the Poisson (*P*) equation<sup>77</sup>:

$$\nabla \cdot (\epsilon_0 \epsilon_r \nabla \phi) = \rho_f \tag{S4-4}$$

 $\epsilon_0$  is = 8.854·10<sup>-12</sup> C<sup>2</sup>/Nm<sup>2</sup>- the permittivity of free space and  $\epsilon_r$  is the relative permittivity of the species near the wall. The  $\phi$  from Eqn. (S4-4) and the  $n_i$  from Eqn. (S4-2) are coupled through:

$$\rho_f = \sum_{i=1}^2 z_i e \, n_i \tag{S4-5}$$

Eqns. (S4-2), (S4-3), (S4-4), and (S4-5) are solved self-consistently, and used to obtain (i) u, and (ii)  $\phi$  and related  $n_i$ , at any point (x,y) in the channel. Here, x is along the direction of fluid flow along the length of the channel, and y is the vertical coordinate – along the height of the channel. The u yields the local velocity, and is averaged along the width of the channel for the average velocity. The  $V_{s,j}$  developed between any two points  $(x_j, y_j)$  and  $(x_{j+1}, y_j)$  along the channel is then computed from  $|\phi(x_j, y_j) - \phi(x_{j+1}, y_j)|$ . The net  $V_s$  was aggregated from such individual contributions along the length of the channel. The mesh size, for the finite element simulations, was chosen to be 5 nm at both the top and bottom walls outwards.

Boundary Conditions: For the N-P equation (Eqn. S4-2), the bulk concentration  $c_o (= c_{Na^+} = c_{Cl^-}) = 0.1$  mM, was kept constant for both the electrolyte ion species at both inlet and outlet, as well as at the centerline of the channel, assumed to be far outside the electrical double layer (EDL), as the Debye length ( $\lambda_D = \sqrt{\frac{\epsilon_o \epsilon_r k_B T}{e^2 c_i}}$ ) was estimated to be ~ 30 nm. The difference of ionic concentration between counter-ions and co-ions, yields a net charge profile density along the height of the channel. The flux normal to top and bottom walls  $J_i = 0$ .

For the Poisson equation (Eqn. S4-4), the surface charge density ( $\sigma$ ) at the top wall<sup>49</sup>:  $\sigma_{PDMS} = -18$  mC/m<sup>2</sup>, while at the bottom surface, (i) at the oil<sup>74</sup>-electrolyte interface is  $\sigma_{oil} = -1.8$  mC/m<sup>2</sup>, and (ii) for the parylene<sup>51</sup>-electrolyte interface is  $\sigma_{Parylene} = -3.7$  mC/m<sup>2</sup>. The  $\sigma$  values are related to the surface electrical field. The outlet was set at zero potential (/ground). It was assumed that the electric field due to EDL and that due to generated streaming potential were completely decoupled. The  $\phi$  at the center-line of flow, in the middle of the channel, *i.e.*,  $\phi$  (y = H/2) = 0 and far away from the EDL at the surface was set to zero.

For the Stokes equation (Eqn. S4-3), the inlet pressure was set to 1200 Pa, while the outlet was set to zero. The no-slip hydrodynamic boundary condition was assumed to hold true at all interfaces.

#### 3.8.5 Hydrodynamic flow with no surface charge but slip at the oil-electrolyte interface

Here, the Stokes Equation (without any electrical body force) is the governing equation

$$-\nabla p + \mu \nabla^2 u = 0 \tag{S5-1}$$

The inlet pressure was set to 1200 Pa, while the outlet was set to zero. A Navier slip boundary condition:  $u_s (y=0) = b \frac{\partial u (y=0)}{\partial y}$  over the flowing electrolyte-oil interface was assumed, with a finite slip velocity:  $u_s$ and slip length: *b*). For a hydrophobic surface the *b* may be approximated<sup>29,78</sup> to be at least *d*, *i.e.*, ~ 18 µm. **3.8.6 Considerations related to the figure of merit (in mV/Pa) and energy conversion efficiency** 

Our focus, in the presented work, was less on the energy conversion efficiency and *more* on methodologies to enhance the streaming potential ( $V_s$ ) per unit pressure difference ( $\Delta P$ ), *i.e.*,  $\left(\frac{V_s}{\Delta P}\right)$ . Indeed, our work has experimentally demonstrated the *largest figure of merit*, thus far, in terms of the voltage generated per unit applied pressure, in comparison to previous experimental work<sup>79–82</sup>.

More specifically, the fluid flow to electrical conversion efficiency  $(Eff.) = P_{out} / P_{in}$ , where the output power:  $P_{out} (= \frac{1}{4} V_s \cdot I_s)$ ,  $I_s (= V_s \frac{A\sigma}{L})$  is the streaming current, with  $A (= w \cdot h)$  as the cross-sectional area of the channel of width: w, and height: h, and  $P_{in} = Q \cdot \Delta P$ , Q is the flow rate  $(=\frac{Gh^3}{12\eta})$ , with G as a constant pressure gradient, and  $\eta$  the viscosity.

Then, the  $Eff.=\frac{3V_S^2\sigma\eta w}{\Delta PLGh^3}=3\left(\frac{V_S}{\Delta P}\right)\left(\frac{V_S}{L}\right)\left(\frac{\eta\sigma}{G}\right)\left(\frac{w}{h^2}\right)$ . For a given electrolyte concentration (fixed  $\sigma$ ) and flow velocity (a given u), <u>both</u>  $\left(\frac{V_S}{\Delta P}\right)$  and  $\left(\frac{w}{h^2}\right)$  are important. While the former, *i.e.*,  $\left(\frac{V_S}{\Delta P}\right)$  was considered in detail in our work, the geometrical factor:  $\left(\frac{w}{h^2}\right)$  may be significantly enhanced through the use of nanoscale diameter channels (small h) with the overlap of double layers, as seen for example, in the work by van der Heyden, *et al, Nanoletters*, vol. 6, p. 2232, (2006). We used channels of the order of 250  $\mu$ m in

height over which most of the electric field is zero and a finite field is obtained only close to the channel surfaces of the order of 0.1  $\mu$ m (the Debye length). Consequently, the estimated energy conversion efficiencies in our work is quite small of the order 10<sup>-3</sup> %, employing the computational methodology indicated in van der Heyden, *et al, Nanoletters,* vol. 6, p. 2232, (2006). More specifically, the efficiency values are ~ 7.7 \cdot 10<sup>-4</sup> % (through the use of the oil in a *LFS*), ~ 3.8 \cdot 10<sup>-4</sup> % (on an *AFS*), and ~ 3.5 \cdot 10<sup>-4</sup> % (on a flat unpatterned substrate).

An important point to note is that the energy conversion efficiency may be improved by more than a factor of two through the use of LFS compared to the flat/AFS substrates. We may expect that the efficiency of nanochannels may be further enhanced by using *LFS*, and will be one of the focus areas of future work.

## Chapter 4 Modulation of streaming potential and slip characteristics in electrolyte flow over liquid-filled surface

#### 4.1 Abstract

A significant enhancement in the streaming potential ( $V_s$ ) was obtained in experiments considering the flow of electrolyte over liquid filled surfaces (*LFS*), where the grooves in patterned substrates are filled with electrolyte immiscible oils. Such *LFS* yield larger  $V_s$  (by a factor of 1.5) compared to superhydrophobic surfaces, with air-filled groves, and offer tunability of electrokinetic flow. It is shown that the density and viscosity, surface tension as well as the dielectric constant of the filling oil, in the *LFS*, determines  $V_s$ . Relating a hydrodynamic slip length to the obtained  $V_s$  offers insight into flow characteristics, as modulated by the liquid interfaces in the *LFS*.

#### **4.2 Introduction**

Electrokinetic flows, considering the movement of electrolyte relative to another charged surface, are important for understanding the effects of charge accumulation and dispersion, with applications ranging from power generation<sup>83–85</sup> to biochemical separations<sup>31,86</sup>. For instance, in pressure driven flows, the flow of excess ions in the electrical double layer (EDL) near the surface, would generate a streaming current, and in the absence of such current – an ion accumulation related streaming potential ( $V_s$ ). The exploration of the use of this phenomenon for energy sources has a long history<sup>87,88</sup>, and has also received much attention recently, from a micro- and nano-fluidics pestpective<sup>46,83,89</sup>. For example, the overlap of the EDLs in nanoscale channels could lead to unipolar flows mimicking battery-like behavior. However, considering that flow and throughput restrictions are a major constraint at the nanoscale<sup>90</sup>, we focus our work as presented here on engineered microscale channels for obtaining high  $V_s$ .

Fluid flows over smooth surfaces with concombinant no-slip conditions, yield low streaming currents and potential. It has then been indicated that enhanced electroosmotic mobility: M, may be obtained through the use of patterned<sup>91,92</sup> or superhydrophobic (SH) surfaces<sup>5</sup>, in both laminar and turbulent flows. The basis is that the aiding of the counter-ion motion near the surface would enhance the electroosmotic current and the related  $V_s$ . From the Helmholtz-Smoluchowski relation<sup>32</sup>,  $M = \frac{\varepsilon \zeta}{\eta}$ , where  $\varepsilon$  (=  $\varepsilon_o \varepsilon_r$ , with  $\varepsilon_o = 8.854 \cdot 10^{-12} \text{ C}^2/\text{Nm}^2$  is the vacuum permittivity and  $\varepsilon_r$  is the relative permittivity) is the dielectric constant of the electrolyte,  $\zeta$  is the zeta potential, and  $\eta$  is the viscosity. However, such an increased mobility is based on the requirement that the no-shear/slipping surfaces have a significant charge density, and of a similar magnitude and sign as that of the no-slip surface in fluidic channels.



**Figure 4.1** Large (a) slip velocities  $(u_s)$ , as well as (b) electrokinetic streaming potentials  $(V_s = V_1 - V_2)$  may be generated through Poiseuille type electrolyte flow (under a pressure difference  $\Delta P = P_1 - P_2$ ) over a liquid filled surface (*LFS*), with interstices/grooves filled with liquids of varying physical parameters, such as the density, dielectric constant, viscosity, *etc.* The positive and negative charges are indicated by the red and the blue circles. A larger  $V_s$  may be generated due to a larger  $\zeta$  (the zeta potential).

Traditionally, SH surfaces were fabricated through the use of air to promote slip, *e.g.*, in patterned<sup>29,93</sup> or non-smooth surfaces<sup>7,9,10</sup>, with a flat air-electrolyte interface. Through much insight, it was determined that only a charged liquid-air interface could enhance the streaming potential<sup>94–96</sup>. It was found, for instance, that the  $V_s$  for flow over uncharged liquid-air interface was similar to that of the homogeneously charged smooth surface for low EDL thicknesses and  $\zeta$  <sup>94,95</sup>, and may be smaller compared to that obtained from a smooth surface with high zeta  $\zeta$  <sup>96</sup>. There is also considerable ambiguity on the

electrical character of the air-electrolyte interface, as to whether there is a charge at all *to* yielding an electrical potential due to residual OH<sup>-</sup> ions.

We circumvent such ambiguity in our work using liquid (of an appropriate  $\varepsilon_r$ ) filled into the interstices of patterned surfaces, fabricated to mimic SH character. Consequently, a definitive charge density would be ensured at the flowing electrolyte slipping surface<sup>97</sup>, promoting the conditions for obtaining large streaming potentials: Fig. 4.1. The shear plane is generally understood as the location at which the fluid starts to move and would be expected to be located at the liquid-liquid interface and the solid-liquid interface. Fig. 4.1(a) emphasizes the flow profile (incorporating the slip velocity:  $u_s$ , as well as the associated slip length: *b*) over a liquid filled surface (*LFS*) <sup>98</sup>, while Fig. 4.1(b) depicts the associated electrokinetic flows over the *LFS*. In this study, we investigate the characteristics of the *LFS*, which may be used for enhancing the magnitude of the associated  $V_s$ . We also show that the use of such surfaces enables a simultaneous decrease of the fluid drag<sup>99–102</sup>, and increased flow<sup>103,104</sup> coupled with an increase in the  $V_s$ .

#### 4.3 Experiment section

**Fabrication and characterization of hybrid surfaces: the** *AFS* **and the** *LFS*. A reference air filled surface (*AFS*) – consisting of rectangular ridges and troughs, was fabricated through photolithography processes and is shown in Fig. 4.2(a). Briefly, a Si wafer (n-type, <110>, thickness 500µm, from University Wafer) was cleaned with acetone, IPA, and rinsed by DI water, then baked at 180° C for 5 minutes. Negative photoresist NR9-3000 was coated on Si wafer at 3000 rpm for 40 s and then baked at 120° C for 60 s. The specific pattern was defined through a mask aligner (EVG 620) and developed (using RD6) for 1 minute. Subsequently, the resist patterned wafer was subject to dry etching (using Oxford Plasmalab 100 RIE/ICP) to yield a trench depth (*h*) ~ 95µm. The etched Si wafer was then put in RR2 photoresist remover over ~ 12 hours and any resist residues further removed (using PVA TePla PS100 at 120 sccm, 200 W for 90 s).

Finally, the grooved surface was coated with Parylene-C (using PDS 2010 Parylene Coater) to a thickness of ~ 600 nm. The basis *AFS* was defined through an air fraction,  $\phi_a = \frac{d}{d+w}$  - the ratio of the trough width (*d*) over one period length (*d* + *w*). The surfaces were imaged using environmental scanning electron microscopy (ESEM): FEI/Phillips XL ESEM and the FEI Quanta FEG 250 ESEM, respectively: Fig. 4.2(b): *AFS*, and Fig. 4.2(c): *LFS*.



**Figure 4.2** A liquid filled surface (*LFS*) was fabricated through (**a**) lithographic procedures to synthesize a surface with ridges and grooves, and subsequent deposition of a uniform parylene film. The grooves are filled with a liquid (immiscible with the electrolyte and with low surface tension) and subsequently the *LFS* is planarized. (**b**) The surface, as indicated through a scanning electron microscope (SEM) image is defined through the width of the trough (*d*) and ridge (*w*), as well as the height (*h*). (**c**) An image of a practical *LFS*, indicating grooves filled with oil.

Due to the oleophilicity of the  $AFS^{105}$ , oils immiscible with water and with low surface tension, *e.g.*, Dupont Krytox GPL 104 (General Purpose Lubricant) - a fluorinated synthetic oil, castor oil, and synthetic motor oils (0W-20 and 20W-50), can be filled into the troughs of the patterned surface to yield a *LFS*, located at the bottom of the micro-fluidic channel. The physical parameters for the oils used in the *LFS* are listed in Table I. The oil was spread evenly on the surface via soft brush and any excess oil was removed from the top of the *LFS*. Detailed microscopic examination, *e.g.*, Fig. 4.2(c), did not indicate any entrapped bubbles in the grooves. The oil filling was found to be reliable and is stable for the *LFS*, *e.g.*, compared to the *AFS*. For instance, the surface energy of the *LFS*:  $E_{LFS} \sim 0.5 \times \gamma_{pary} + 0.5 \times \gamma_{oil} \sim 32 \text{ mJ/m}^2$  is an order of magnitude smaller compared to that of the *AFS*:  $E_{AFS} \sim r \times \gamma_{pary} \sim 290 \text{ mJ/m}^2$  with r as the roughness factor (= 6.3, from the ratio of the total area to the projected area<sup>105</sup>). The excellent liquid-interface stability was also considered previously considering surface-immobilized lubricants<sup>106</sup>.

**Table 4.1** The parameters of the oils (the density:  $\rho$ , kinematic viscosity:  $\eta$ , dynamic viscosity:  $\mu$ , dielectric constant:  $\varepsilon$  and the surface tension:  $\gamma$ , of the liquids filling the interstices of the liquid filled surfaces (*LFS*).

Oils	ρ (g/cm <sup>3</sup> )	η (cm²/s)	µ (mPa·s)	3	$\gamma (mJ/m^2)$
Krytox <sup>®</sup> 104	1.93	1.77	340	2.1	18
Castor Oil	0.96	3.25	312	4.7	39
Motor Oil	0.86	1.19	102	2.4	30
0W-20					
Motor Oil	0.86	5.76	495	2.4	30
20W-50					

Setup for measurement of the streaming potential ( $V_s$ ). The experimental setup is shown in Fig. 4.3(a) with a schematic: Fig. 4.3(b). A microchannel (11.8 cm in length, 9 mm in width, and 255 µm in height), was constituted from a *top surface* (constituted from a polycarbonate coated with silicone), and a bottom *AFS* or *LFS*. These two surfaces are separated by a silicone rubber spacer for sealing and adjusting the height of the channel. For the streaming potential ( $V_s$ ) measurement, electrolyte, *i.e.*, salt water (in the concentration range of 0.1 mM – 10 mM), preparing by dissolving NaCl into DI water, was pumped by a syringe pump through a tube into the microchannel at a constant flow rate, and provides a pressure gradient. The pressure drop for the experiment was in the range of 200 Pa to 1200 Pa, with lower and higher pressures

yielding unreliable values for the  $V_s$ . The voltages – related to the  $V_s$ , were measured (using a Keithley 2700 Multimeter with input resistance >10 G $\Omega$ ) through Ag/AgCl electrodes inserted into the two end reservoirs located at the inlet and outlet of the channel. The  $V_s$  for each applied pressure was measured six times. Given that the channel length (*L*) and width (*W*) are much larger than channel height (*h*), Poiseuille flow is assumed. The  $V_s$  was measured in flow at different orientations with respect to the patterned substrate to consider various tensorial aspects<sup>95,107</sup>.



**Figure 4.3.** The apparatus used for the measurement of the streaming potential, (a) consisting of a syringe pump connected by a tube to the microchannel to provide a constant flow rate to the electrolyte and a pressure gradient, and a (b) schematic of the placement of the electrodes (non-polarizable Ag/AgCl type) at the ends to measure the streaming potential ( $V_s$ ) related voltage (*e.g.*, the), along the microchannel.

**Measurement of the Pressure.** The electrolyte flow rate in the channel, constituted from rectangular plate geometry<sup>46</sup>, was modulated through a syringe pump: Fig. 4.3(a), and was transduced to an applied pressure ( $\Delta P$ ) based on the Poiseuille relation:  $\Delta P = \frac{12\eta LQ}{h^3}$  (*Q* is flow rate, *L* is the channel length of ~ 11.8 cm, and *h* the channel height of ~ 250 µm). The  $\Delta P$  over the channel length (where the other end was left open to atmosphere) motivated the pressure gradient driven flow. For the experiment, the flow rates were chosen as 0.0275 mL/s (corresponding to 200 Pa), 0.055 mL/s (440 Pa), 0.0825 mL/s (670 Pa), 0.11 mL/s (900 Pa) and 0.1375 mL/s (1130 Pa). Additionally, the calculated pressure drop was checked using a manometer (UEI EM152) by inserting the measurement ports into the two reservoirs of the channel and corresponded very closely to the values estimated from the Poiseuille relation.



**Figure 4.4** The contact angles measured on a (a) water-parylene, and (b) water - GPL 104 interface at ~94.6° and ~ 68.9°, respectively. These values were used for estimating the corresponding  $\gamma_{SL}$  from given  $\gamma_{SA}$  (= 46 mJ/m<sup>2</sup> for parylene) and  $\gamma_{LA}$  (= 72 mJ/m<sup>2</sup>).

**Table 4.2** The measured contact angles for oils and parylene and calculated interface energy  $\gamma_{SL}$ .

Substrate	Contact angle	$\gamma_{SL}$ (mJ/m <sup>2</sup> )	
Krytox <sup>®</sup> 104	68.86°	-7.97	
Castor Oil	74.39°	19.63	
Motor Oil 0W-20	82.93°	21.14	
Motor Oil 20W-50	81.01°	18.75	
Parylene-C	94.6°	51	

Measurement of the contact angle. The electrolyte-oil interfacial energy will influence the interface shape and adhesion force, and modifies the interface flow profile and the  $V_s$ . From the Young relation:

$$\gamma_{SA} - \gamma_{SL} - \gamma_{LA} \cos\theta = 0 \tag{4.1}$$

with  $\gamma_{SA}$ ,  $\gamma_{SL}$ ,  $\gamma_{LA}$  as the substrate-air, substrate-liquid and liquid-air surface energy, respectively,  $\theta$  is the substrate-liquid contact angle. The contact angles of electrolyte (0.1mM NaCl solution) with parylene and Si substrate with different oil films were measured using Ramé-Hart Model 190 Contact Angle Goniometer at 20°C, for estimating the corresponding  $\gamma_{SL}$  from given  $\gamma_{SA}$  and  $\gamma_{LA}$ : Fig. 4.4. The estimated  $\gamma_{SL}$  values for the various oils are listed in Table 4.2 (with  $\gamma_{SA} = 46 \text{ mJ/m}^2$  for parylene and  $\gamma_{LA} = 72 \text{ mJ/m}^2$ ).

#### 4.4 Results and discussion

Pressure induced flow on the *LFS*, and related measurement of the streaming potential. We report measurements of the  $V_s$  with 0.1 mM electrolyte concentration, where maximal values of the streaming potential were determined: Fig. 4.5. Here, the observed variation arises from the Helmholtz – Smoluchowski relation:  $V_s = \frac{\varepsilon \zeta}{\eta \kappa_b} \Delta P$ , in terms of the electrolyte dielectric constant ( $\varepsilon$ ), the surface zeta potential ( $\zeta$ ), electrolyte viscosity ( $\eta$ ) bulk electrolyte conductivity ( $\kappa_b$ ) and pressure drop in the channel ( $\Delta$ P). Assuming relatively constant  $\varepsilon$  and  $\eta$  for the electrolyte, the  $\kappa_b$  is proportional to concentration (I) <sup>2,36</sup>, while the  $\zeta \approx c1 * \log[I] + c2$  – where c1 and c2 are constants. The overall effect<sup>36</sup> is that the  $V_s$  decreases with the I.



**Figure 4.5** Variation of the streaming potential ( $V_s$ ) for air filled surface (*AFS*), and the GPL oil filled *LFS* and castor oil filled *LFS* at different flowing electrolyte concentrations: *I*, (0.1 mM, 1 mM and 10 mM) at 1200 Pa at transverse direction. The  $V_s$  is fit to a relation of the form ~ log [*I*]/[*I*] with  $R^2 = 0.993$  for GPL oil filled *LFS*,  $R^2 = 1$  for castor oil filled *LFS*, and  $R^2 = 1$ , for the *AFS*.

A typical response of the measured voltage as a function of flow is indicated in Fig. 4.6(a). The voltage increases when the electrolyte was driven through the channel and subsequently attains a steady value  $V_f$ . The difference between  $V_f$  and the initial voltage:  $V_0$  (in the absence of flow) was defined to be the measured streaming potential:  $V_s$ . The variation of the  $V_s$  with applied pressure is shown in Fig. 4.6(b) for *both* the AFS and *LFS* (with four different filling liquids) at transverse direction. Generally, a linear variation between the  $V_s$  and the  $\Delta P$ , following the Helmholtz-Smoluchowski relation<sup>32</sup> is observed.



**Figure 4.6 (a)** A typical measured voltage response indicating the voltage increase under the action of an applied pressure of 1200 Pa causing the flow of 0.1 mM NaCl solution. The periodic action of the syringe pump is responsible for the increase (flow) and decrease (no flow/Flow Pause) of the measured voltage. The corresponding velocity is indicated on the right-hand side. (b) A linear variation of the streaming potential ( $V_s$ ) as a function of the applied pressure ( $\Delta P$ ), for various filling liquids/oils, in the *LFS*. The GPL is a fluorinated synthetic oil while the 0W-20 and 20W-50 are synthetic motor oils. It is to be noted that a larger  $V_s$  is obtained using GPL and other oils compared to air in the interstices, *i.e.*, an air-filled surface (*AFS*).

**Possible influence of oils on the measured**  $V_s$ . The selected oils, which serve as lubricants in the *LFS*, vary in terms of the dielectric constant ( $\varepsilon$ ), dynamic viscosity ( $\eta$ ), electrical conductivity ( $\kappa$ ), surface tension ( $\gamma$ ) and density ( $\rho$ ) – with the individual properties as listed in Table 4.1. As the  $V_s$  may be related to the electrolyte-oil interface shape and influenced by many of such parameters, we first aim to obtain empirical relationships of such an influence through the Buckingham II theorem<sup>110</sup>. While the given *LFS* system is quite complex, as related to the large number of parameters and the multi-physics aspects, the motivation was to make sense of the results through dimensional analysis. Consequently, at given flowing liquid, groove fraction and groove orientation, one may obtain related non-dimensionalized groups, *i.e.*,

$$\Pi_0 = \frac{V_s}{\eta^{-0.5} \kappa^{-0.5} \gamma}$$
(4.2a)

or

$$\Pi_1 = \frac{\rho}{\varepsilon^{-1} \eta^3 \kappa \gamma^{-2}} \tag{4.2b}$$

may be obtained. From fitting the above, with  $V_s$  from experiment and the values of oil properties, an empirical relationship  $\Pi_0 = F(\Pi_1)$  and fit ( $R^{2} \sim 0.82$ ) was obtained: **Fig. 4.7**. A larger (/smaller)  $\phi_{oil}$  may be expected to increase (/decrease) the polarizability and increase (/decrease) friction at the oil-water interface<sup>1</sup> and yield a decreased (/enhanced)  $V_s$ . A possible correspondence between the  $\eta$  and the  $\varepsilon_r$  was indicated earlier<sup>111</sup>.

For a better understanding of the obtained  $V_s$  variation, we considered the flow profile variation for an *LFS* taking into account *both* the electrical attributes, such as  $\varepsilon$ ,  $\kappa$ , as well as the nominal fluidic parameters, such as the  $\rho$ ,  $\eta$ , and  $\gamma$ , of both the impregnated liquid in the *LFS* as well as at the electrolyte – *LFS* interfaces.



**Figure 4.7.** Experimentally derived correlations between the obtained streaming potential ( $V_s$ ) on *LFS* as a function of non-dimensional parameters, through considering the viscosity ( $\eta$ ), electrical conductivity ( $\kappa$ ), surface tension ( $\gamma$ ), the density ( $\rho$ ) and the dielectric constant ( $\varepsilon$ ) of the filling liquid.

# The influence on slip, as a function of electrical parameters of the filling liquid, in *LFS*. One manifestation of a liquid impregnated surface, akin to the *LFS* discussed in this work, was a slippery liquid infused porous surface<sup>27</sup>. There, the wettability in terms of enhancing the liquid slip was considered. The

slip length (*b*), as determined through the slip velocity ( $u_s$ ) and the surface shear rate  $(\frac{\partial u}{\partial y}|_{y=0})$  from Navier boundary condition:  $u_s = b \frac{\partial u}{\partial y}|_{y=0}$ , would be dependent on the filling liquid properties<sup>99,104</sup>. Considering electrical characteristics, we indicate the inverse relation of the  $V_s$  to the dielectric constant of oil:  $\varepsilon_{oil}$ . For a hybrid surface, such as the LFS, consisting of both solid parylene and liquid oil, an effective slip length  $b_{eff}$  may be defined.

Using the relations in Belyaev & Vinogradova<sup>112</sup> and Nizkaya<sup>113</sup> *et al*, we calculated the slip length of the *LFS* with transverse grooves where we consider the groove orientation, groove fraction and viscosity ratio. The local slip length for transverse groove, considering the viscosity of the electrolyte ( $\eta_{elect}$ ) and the oil ( $\eta_{oil}$ ) and a geometry-dependent function:  $\beta$ , was estimated through<sup>113</sup>:

$$b = w \frac{\eta_{elect}}{\eta_{oil}} \beta \tag{4.3}$$

For deep grooves, as in our case (with a height: *h* of ~ 95 µm, and a width: *w* of 18 µm, *i.e.*, with  $h/w \ge 1$ ),  $\beta \approx 0.12$ . The effective slip length ( $b_{eff}$ ) was then estimated to be:

$$b_{eff} = \frac{L}{2\pi} \frac{\ln\left[\sec\left(\frac{\pi\phi}{2}\right)\right]}{1 + \frac{L}{2\pi b_1^{\gamma}} \ln\left[\sec\left(\frac{\pi\phi}{2}\right) + \tan\left(\frac{\pi\phi}{2}\right)\right]}$$
(4.4)

The calculated  $b_{eff}$  (/b) for GPL oil and castor oil filled *LFS* are then ~ 2.5 nm (/6.4 nm) and ~ 2.7 nm (/6.9 nm), respectively. The  $b_{eff}/b$  ratio is then ~ 0.4, corresponding closely<sup>114</sup> to the expected value of ~ 0.5 (the area fraction), when  $b/L \ll 1$ ,  $b_{eff}^{\perp,\parallel} \simeq \phi b$ . If we consider the charge effects at the gas/oil sectors on flow transport<sup>114</sup>, then the electroosmotic mobility for transversely oriented groove was shown to be:

$$M_{\perp} = \frac{\varepsilon \zeta}{\eta} \left[ 1 + \frac{b_{eff}}{b} \left( \frac{q_2}{q_1} (1 + kb) - 1 \right) \right]$$
(4.5)

From literature, the surface charge density is ~ -3.6 mC/m<sup>2</sup> (=  $q_I$ ) for parylene<sup>51</sup> and ~ -1.8 mC/m<sup>2</sup> (= $q_2$ ) for oil<sup>74</sup>. Through  $V_s$  measurements on flow of 0.1 mM NaCl solution (with  $k \sim (1/30)$  nm<sup>-1</sup> =  $1/\lambda_D$ , where  $\lambda_D$  is the Debye length) on parylene and PDMS coated *flat* substrates, and  $V_s = \frac{\varepsilon \zeta}{\eta \sigma} \Delta P$ , we estimated

that  $\zeta_{parylene} \sim 31.7 \text{ mV}$  and  $\zeta_{PDMS} \sim 26.9 \text{ mV}$ . Then, the *enhanced* zeta potential for bottom surface  $\zeta_{bottom,en}$ , considering the correspondence<sup>114,115</sup> of the electroosmotic mobility to the  $V_s$  is:

$$\zeta_{bottom,en} = \zeta_{bottom} \left[ 1 + \frac{b_{eff}}{b} \left( \frac{q_2}{q_1} (1 + kb) - 1 \right) \right]$$
(4.6)

and the concomitant  $\zeta$  for the channel is:

$$\zeta_{channel} = 0.5\zeta_{PDMS} + 0.25\left(\zeta_{parylene} + \zeta_{oil}\right) \left[1 + \frac{b_{eff}}{b} \left(\frac{q_{oil}}{q_{parylene}}\left(1 + kb\right) - 1\right)\right]$$
(4.7)

Here, the effective zeta potential of the rectangular channel was defined through  $\zeta_{eff}(=\frac{1}{2}\zeta_{bottom} + \frac{1}{2}\zeta_{top})$ , and for the  $\zeta_{bottom}$ , a weighted summation<sup>94</sup> of  $\zeta_{parylene}$  and  $\zeta_{oil}$  was assumed. We then estimate that  $\zeta_{GPL} \sim 214$  mV and  $\zeta_{castor} \sim 66$  mV from the above relation. The larger  $V_s$  obtained with the use of GPL oil in the *LFS* compared with that of the use of castor oil could be tentatively ascribed to the larger  $\zeta_{GPL}$ .

It may also be necessary to consider the issue of *fixed* charges and *mobile* charges on the slipping interfaces<sup>109</sup>. In Maduar<sup>109</sup>, *et al*, a parameter  $\mu$  was used to modify the Navier slip boundary conditions, with  $\mu = 0$  for fully *mobile* charges and with  $\mu = 1$ , for *fixed charges*. If there is adsorption of hydroxyl ions, for example, the liquid-oil interface would be charged, and correspondent  $V_s$  is modified to:

$$V_s = \frac{\varepsilon \zeta}{\eta \sigma} (1 + \mu b_{eff} k) \Delta P \tag{4.8}$$

Consequently, another reason for the difference in the  $V_s$  between GPL- and castor oil- filled *LFS* may be due to differing  $\mu$ , and that  $\mu_{GPL} > \mu_{castor}$  - due to the smaller dielectric constant of GPL oil ( $\varepsilon \sim 2.1$ ) compared to castor oil ( $\varepsilon \sim 4.7$ ). Considering the continuity of tangential shear stress across the electrolyte (with  $\eta$  of ~ 1 mPa·s) - oil (with  $\eta$  of ~ 300 mPa·s) interface, it was assumed that the slip is along the oil-electrolyte interface.

An alternate understanding of the influence of charges, may be manifested through a charge density ( $\sigma$ ) difference:  $|\sigma_{parylene} - \sigma_{oil}|$ , which is expected to yield a longitudinal electric field ( $E_x$ ) as well as a transverse electric field  $(E_y)$ , the former of which drives ion motion across the surface: Fig. 4.8. Here, where x is in the direction of fluid flow and the and y is the orthogonal direction- transverse to the direction of the pressure driven flow. It may be expected that with a large (/small)  $\sigma_{oil}$ , that the screening of the charge into the flowing electrolyte occurs over a smaller (/larger) distance yielding a larger (/smaller)  $\rho_s$  and  $\zeta$ . The shear plane is then closer (/further), implying smaller (/larger)  $u_s$  and b. We estimated, for the parylene coated Si surface:  $\sigma_{parylene}$  at - 3.6 mC/m<sup>2</sup> and that of the oil:  $\sigma_{Oil}$  at - 1.8 mC/m<sup>2</sup>. The resultant  $E_y$  would be of the form:  $E_{y, parylene} \sim \sigma_{parylene} / \varepsilon_{elec}$  and  $E_{y, oil} \sim \sigma_{oil} / \varepsilon_{elec}$ , respectively. If we assume no residual charge at the oil-electrolyte (elec) interface, then from the continuity of electrical displacement and electrical fields, across the interface: (i)  $\varepsilon_{oil} = \varepsilon_{v,oil} = \varepsilon_{elec} E_{v,elec}$ , and (ii)  $E_{x,oil} = E_{x,elec}$ . Here, the capillary number  $(Ca) = \eta v/\gamma$ , with  $\eta$  the dynamic viscosity, v as the characteristic velocity, and the surface tension ( $\gamma$ ), was estimated to be of the order of 10<sup>-3</sup>, which indicates that the surface tension dominates over the viscous forces. A similar inference could be made through estimating the Bond number (Bo) – which relates the difference of the density of the two phases (electrolyte and oil) over a characteristic length, say, the groove width, with respect to the surface tension. Consequently, the pressure driven flow induced viscous force will not significantly deform the liquid-liquid interface<sup>116</sup>.



**Figure 4.8** The schematic of electrokinetic flow over *LFS* surface with varying surface charge density ( $\sigma$ ). As  $\sigma_{oil}$  is less than  $\sigma_{parylene}$ , the surface conductivity  $\kappa_s$  is smaller for the oil regions. Consequently, driven surface currents: *J* (both parallel: in the *x*-direction, and perpendicular: in the *y*-direction, to the *LFS*) serve to enhance the slip and the slip length (*b*). The *b* was correlated to the healing length  $L_H \sim \kappa_s/\kappa_b$  (the ratio of the surface to the bulk electrical conductivities), the Debye length ( $\lambda_D$ ),  $\eta_{elec}$ , as well as the  $\varepsilon_{oil}$  and  $\sigma_{oil}$ , through analysis and experiment.

The deployment of a *hybrid* bottom surface, constituted from both the liquid and the solid with the concomitant differing surface charge densities may be expected to yield concentration polarization<sup>44</sup> as well as non-uniform conduction<sup>117</sup> effects over the surface. In electroosmotic flows, the ions contributing to the surface current ( $J_x \sim \kappa_s E_x$ , with  $\kappa_s$  being the surface conductivity) in the direction of the flow are replenished through an ion current from the bulk liquid (of conductivity:  $\kappa_b$ ) over a length scale (termed a healing length:  $L_H$ ) and due to a normal component of the electric field ( $E_y$ ). From such ion conservation considerations:  $\kappa_s E_x \sim \kappa_b E_y L_H$ . If  $E_x$  and  $E_y$  are of comparable magnitudes, then the  $L_H$  is of the order of  $\kappa_s/\kappa_b$ .

For flow over the LFS, it may be expected that the  $E_x$  or the  $E_y$  may contribute to  $u_s$  and the resultant surface current:  $J_{s,x} \sim \rho_s u_s$ , where  $\rho_s$  is the *mobile* surface charge density. From analogy to the arguments presented above, for  $E_y$  induced current dominated case,  $\kappa_b E_y L_H \sim \rho_s u_s$ , implying, for a given shear rate  $(\partial u/\partial y)$  in the absence of nonlinear effects<sup>6</sup>:

$$b \sim \frac{\kappa_b \sigma_{oil} L_H}{\varepsilon_{elec} \rho_s} \tag{4.9a}$$

However, this relation can not explain the experiment results that the slip length is larger for filled oil with smaller dielectric constant. Alternately, if the longitudinal electrical fields, say  $E_{x, elec}$  (= $E_{x, oil}$ ) are more responsible for the fluid slip, then  $\kappa_s E_{x, elec} \sim \rho_s u_s$ , indicating:

$$b \sim \frac{\kappa_s \sigma_{oil}}{\varepsilon_{oil} \rho_s} \tag{4.9b}$$

Here, a larger *b* or a  $u_s$  also seems to be facilitated through a higher  $\sigma_{oil}$ . Additional electrostatic considerations, *e.g.*, related to the screening in the electrolyte, must also be considered. It was previously indicated, for instance<sup>74</sup>, that the magnitude of the associated surface charge density may be independent of oil type, and mostly dependent on the aqueous phase/electrolyte. Indeed, testing with several oils indicates that the correlations indicated in Eqns. (4.2a) and (4.2b), are valid. As we use non-polar oils in the *LFS*, the molar conductivity is negligible and the Walden rule relating  $\eta$  to the  $\sigma$  is not applicable<sup>118</sup>.

At the shear plane, from considering an equivalence of the shear stress:  $\tau$  (~ $\eta \partial u/\partial y$ ) to  $u_s$ , a direct correlation between the *b* and the  $\eta$  of the electrolyte may be deduced. Indeed, such a correlation and scaling has been previously indicated for liquid slip on air<sup>119</sup>, through a "gas cushion" model<sup>29</sup>. From the previously deduced  $b \sim 1/\varepsilon_{oil}$ , we now obtain a significant correlation for the oil-electrolyte interface, that:

$$b \sim \frac{\eta_{elec}}{\varepsilon_{oil}} \tag{4.10}$$

The expression above posits a relation between the measured  $V_s$  and the flow characteristics as modulated by the oil-electrolyte interface/s in the *LFS*. Consequently, aspects of the surface that promote b, e.g., geometry<sup>29</sup>, air bubbles<sup>6</sup>, surface interactions, *etc.*, may all now be utilized for obtaining enhanced  $V_s$ .

#### **4.5** Conclusion

In summary, we have indicated that enhanced streaming potentials  $(V_s)$  may be obtained through electrolyte flows over liquid filled surfaces. It was observed that the  $V_s$  was sensitive to the character of the liquid, in terms of the density viscosity, dielectric constant, in the *LFS*. Moreover, the magnitude and variation of the pressure at the ridge-groove interface seems to be important in determining the enhancement of the  $V_s$  over that obtained in air-filled grooves and surfaces. The influence of such local perturbations merit further study, *e.g.*, the modification of the flow profile close to *LFS* surface<sup>120</sup>, due to the unequal interface energy between electrolyte-parylene (*ridge*) and electrolyte-oil (groove). A correlation of the slip length (*b*) to the  $V_s$  yields a better understanding of the hydrodynamics inherent in electrokinetic flows. The presented study indicates a path forward<sup>94</sup> in terms of generating large streaming potentials, through fluid flow in microchannels with a *hybrid* surface – consisting of both solid and oil interfacing with an aqueous electrolyte, moving beyond smooth walled nanoscale channels<sup>46</sup>. Our results and those expected from extending our investigations may be utilized to gain greater insights with applications ranging from energy conversion<sup>46,66,83,85,87</sup> to transport in lab on a chip devices<sup>59,60</sup> and nanosystems<sup>121</sup>.

#### 4.6 Acknowledgement

This chapter, in full, is a reprint of the materials as it appears in Fan, B. and Bandaru, P.R., Modulation of streaming potential and slip characteristics in electrolyte flow over liquid-filled surfaces, Langmuir, 2019, 35, 18, 6203 – 6210. The dissertation author was the primary investigator and author of this paper. The authors are grateful for support from the National Science Foundation (NSF: CBET 1606192). We also appreciate the assistance of Prof. J. Friend and P. Chen for help with imaging.

### Chapter 5 Tensorial Modulation of Electrokinetic Streaming Potentials on Air and Liquid Filled Surfaces

#### 5.1 Abstract

Textured surfaces, comprised of grooves filled with air, e.g., air-filled surfaces (AFS), or with liquid, e.g., liquid-filled surfaces (LFS), significantly influence fluid flows and the related electrokinetic streaming potential (Vs). Here, electroosmotic mobility related tensorial effects on the Vs were experimentally investigated. A significant modulation of the Vs, as high as 100%, due to transverse pressure gradients, was demonstrated. The study yields insights into understanding geometrical effects in electrolyte flows with implications to the establishment of local electric fields, energy generation, and biological separations.

#### **5.2 Introduction**

Electrokinetic flows, considering the movement of electrolyte relative to another charged surface, are important for understanding the effects of charge accumulation and dispersion with applications ranging from power generation<sup>83–85</sup> to biochemical separations<sup>31,86</sup>. There are two major related phenomena in such flows, under a pressure difference across the microchannel, *i.e.*, (i) where the motion of ions in the electrical double layer (EDL) near a charged surface generates an electrical streaming current ( $I_s$ ), and (ii) under open-electrical circuit conditions – where a potential difference, termed the streaming potential ( $V_s$ ) is measured, due to charge separation. The  $V_s$  may be particularly enhanced in micro- and nano-scale fluidics<sup>46,83,89</sup>, *e.g.*, through using through the overlap of EDLs in nanometer size pores/channels which may enable unipolar flow and new battery-like voltage sources. Our group has pioneered the use of liquid filled surfaces (*LFS*) to simultaneously enhance the  $V_s$  as well as to reduce frictional drag<sup>100,101,123</sup> between the flowing electrolyte and the surface. Here, the hydrophobic character of the *LFS* enhances flow, through inducing a finite slip

velocity<sup>66</sup>, and reduces the potential drop that contributes to a reduction of the  $V_s$ . Going beyond such preliminary measurements, we discuss in this work, how the *AFS/LFS* geometry especially the groove orientation, may be related to the changes in the  $V_s$ , which has been typically considered through the Helmholtz-Smoluchowski (H-S) equation, and derived on homogeneous surface<sup>8,36</sup>:

$$V_{\rm S} = \frac{\varepsilon \zeta}{\eta_e \sigma} \Delta P \tag{5.1}$$

Here,  $\varepsilon$  (=  $\varepsilon_o \varepsilon_r$ , with  $\varepsilon_o$ =8.854·10<sup>-12</sup> C<sup>2</sup>/Nm<sup>2</sup> is the vacuum permittivity and  $\varepsilon_r$  is the relative permittivity) is the permittivity of the electrolyte,  $\zeta$  is the zeta potential of the surface,  $\eta_e$  is the electrolyte viscosity, and  $\sigma$  is the electrolyte conductivity. Relevant to SH surfaces<sup>5</sup> is an enhanced ion mobility<sup>32</sup>, M=  $\frac{\varepsilon\zeta}{\eta_e}$ , predicated on the requirement<sup>94,96,124</sup> that the surfaces ensuring fluid slip have a significant charge density with a similar magnitude and sign as that of the no-slip surface. However, when the surface is uncharged/partially charged, the magnitude of the  $V_s$  could be diminished compared to a homogeneously charged smooth surface <sup>94,124</sup>, as was also confirmed through our own experimental work. The *LFS* also serve to circumvent the ambiguity related to electrolyte-air interfaces through filling liquid (of a definite  $\varepsilon_r$ ) into the interstices of patterned surfaces (**Figure 5.1**) and ensure a definitive charge density. While preliminary experiments stated that a 50% enhancement in the *V<sub>s</sub>* over the use of traditionally used air-filled surfaces (*AFS*) could be obtained, such improvement seems to hinge critically on the physico-chemical attributes of the filling liquid<sup>125</sup>, as well as on geometrical parameters such as the orientation of the surface with respect to the applied pressure gradient.

Typically, the influence of geometrical parameters of the substrate has been considerably discussed in the context of wetting<sup>1</sup>, *e.g.*, with respect to anisotropy<sup>105</sup>, as well as its influence<sup>30</sup> on the slip length (*b*), *e.g.*, the relative variation for stripe and post geometries<sup>29</sup>. A generalization of the underlying concepts to describe surface anisotropy through slip and related interfacial mobility tensors<sup>107</sup> has yielded insights into the modulation of flows, say, on hydrophobic surfaces<sup>112</sup>. Considering that the electroosmotic mobility (M) may also be formulated as a tensor, insights into charged liquid flows on heterogenous geometries<sup>115</sup> were obtained into electroosmotic slip, with respect to modification of fluid flow patterns, such as flow reversal<sup>114</sup>. The degree of alignment of the texture, *e.g.*, misaligned grooves, was shown to generate transverse shear, also leading to complex flow patterns and mechanisms for mixing in directions transverse to the nominal pressure driven flow axes<sup>126</sup>. All such intriguing characteristics of flow have been studied from a theoretical and computational perspective and are subject to experimental verification. Moreover, the specific impact of such tensorial generalizations, *e.g.*, of the *b* and *M*, have not been discussed with respect to their influence on the  $V_s$ . In this paper, we first report and discuss the modulation of the  $V_s$  brought about through groove orientation variation of the substrate and studied the rotation angle at which yielded the smallest  $V_s$ .



**Figure 5.1** (a) Electrolyte flow (driven through a syringe pump yielding a pressure gradient) in a microchannel generates an electrokinetic streaming potential (Vs) that may be measured by a voltmeter. The patterned parylene substrate either has air in the grooves, i.e., an air-filled surface (AFS), or (b) liquid in the grooves, i.e., a liquid filled surface (LFS), SEM image.

#### 5.3 Methods

**Fabrication of Geometrically Textured Surfaces.** The groove patterned AFS/LFS used in the experiment was fabricated by photolithography and have been previously reported in chapter 3 and 4. Here, we report on the obtained results from surfaces constituted from a given groove width, w, groove period, L (= d + w), with d as the lateral length of the solid surface, oriented at an angle  $\theta$  (in the range of  $-90^{\circ}$  to  $90^{\circ}$ ), with respect to the flow induced by an applied pressure gradient: Figure 5.2, also see right bottom inset to Figure 5.3(a). Here,  $\theta$  is the rotation angle needed to align the coordinate axes (x, y) fixed on the

AFS or LFS to the coordinate axes oriented to the frame in which the liquid electrolyte is flowing (x', y') due to a uniaxial pressure gradient, as indicated in the bottom right inset to Figure 3(a). For sample groove orientation parallel to the flow,  $\theta = 90^{\circ}$ , while for groove orientation transverse to the flow,  $\theta = 0^{\circ}$ . The air fraction,  $\phi a = w/L$ . For the LFS, the grooves were filled (replacing the air) with oils (e.g., Dupont Krytox GPL 104 and castor oil) of low surface energy and immiscible with the aqueous electrolyte (NaCl in deionized water). We ensured the robustness of the LFS, i.e., preventing the drainage of the filling liquid under external shear flow<sup>127</sup> through visual examination of the interface<sup>125</sup>. Moreover, it was previously noted that the filling liquid could be retained indefinitely in the grooves if L was less than a critical length<sup>127</sup>: L $\infty$ . As for the tested experimental conditions, the L $\infty$  was estimated to be of the order of millimeters.



**Figure 5.2** Differing orientation of the grooves with respect to the flow (from left to right). The grooves are oriented (a) perpendicular to the flow ( $\theta = 0^\circ$ , top), (b) parallel to the flow ( $\theta = 90^\circ$ , bottom), or (c) at an intermediate angle ( $\theta = 45^\circ$ , middle). The voltages are measured in the horizontal (i.e., through  $V_{x'}$ ) and vertical (i.e., through  $V_{y'}$ ) directions.

**Measurement of the Streaming Potential.** The Vs experiments were carried out through the measurement of the voltage under the flow of electrolyte at a constant flow rate, driven by an applied pressure difference ( $\Delta P$ ), using a setup as in the schematic of Figure 5.1(a). Electrolyte (typically, NaCl

dissolved in water) of concentration 0.1 mM (chosen for obtaining reliable and maximal Vs) was flowed into the microfluidics chamber (of length ~11.8 cm, width ~0.9 cm, and height ~255  $\mu$ m) using a syringe pump. Poiseuille flow conditions were assumed for converting the flow rate to an equivalent pressure. The differential pressure was in the range of 200 to 1200 Pa and was calibrated using a manometer (UEI EM152). The chosen range of pressure yielded stable and reproducible Vs in the microchannel. The channel surfaces were constituted from an upper surface (silicone coated onto polycarbonate) and a bottom test surface, which was of the AFS or the LFS type. More experimental details on the voltage measurement and flow arrangement have been previously discussed in chapter 3 and 4. The Vs was measured six times at each applied pressure and the average value was used.



**Figure 5.3** Measured streaming potential (Vs) over an (a) AFS, (b) castor oil filled LFS, and (c) GPL oil filled LFS, as a function of applied pressure, at three different orientations of the grooves with respect to the flow. The dotted lines are a guide to the eye and connect the measurements. The top left insets in panels a–c are a schematic of the grooves. The bottom right inset of (a) defines the  $\theta$ , the rotation angle needed to align the coordinate axes (x, y) fixed on the AFS/LFS grooves to the coordinate axes oriented to the frame in which the liquid electrolyte is flowing (x', y'). While the  $V_s(\theta = 0^\circ)$  is generally larger for all the AFS and LFS, each panel (a–c) represents different relative variation of the Vs at  $\theta = 45^\circ$  and  $\theta = 90^\circ$ .

#### 5.4 Results and Discussion

We consider the anisotropy of electrokinetic effects through the electroosmotic mobility tensor M(which is related the ratio of the streaming potential gradient/related electric field: E, to the pressure gradient  $\Delta P$ ). *i.e.*,  $E = -M \frac{1}{\sigma} \Delta P$ . Considering  $\theta$  as the rotation angle needed to align the coordinate axes (x, y) fixed on the AFS or LFS with respect to the axes parallel (x') and perpendicular (y') to the flow direction - as indicated in **Figure 5.3**, we obtain<sup>128,129</sup>:

$$\boldsymbol{M}' = \begin{bmatrix} M_x \cos^2\theta + M_y \sin^2\theta & (M_x - M_y)\sin\theta\cos\theta \\ (M_x - M_y)\sin\theta\cos\theta & M_y \cos^2\theta + M_x \sin^2\theta \end{bmatrix}$$
(5.2)

We then obtain the electric field in the x' direction as:

$$E_{x'} = -\frac{1}{\sigma} \left[ \left( M_x \cos^2\theta + M_y \sin^2\theta \right) \frac{dP}{dx'} + \left( M_x - M_y \right) \sin\theta \cos\theta \frac{dP}{dy'} \right]$$
(5.3)

The corresponding  $V_s$ , over a groove period (L), is the corresponding integral of the  $E_x$ .

$$V_{s} = \int_{0}^{L} \frac{1}{\sigma} \left[ \left( M_{x} \cos^{2}\theta + M_{y} \sin^{2}\theta \right) \frac{dP}{dx'} + \left( M_{x} - M_{y} \right) \sin\theta \cos\theta \frac{dP}{dy'} \right] dx'$$
(5.4)

For instance, at representative values of  $\theta$ , we obtain:

$$V_s(0^\circ) = \int_0^L \frac{1}{\sigma} M_x \frac{dP}{dx'} dx'$$
(5.5a)

$$V_{s}(45^{\circ}) = \int_{0}^{L} \frac{1}{\sigma} \left[ \frac{1}{2} \left( M_{x} + M_{y} \right) \frac{dP}{dx'} + \frac{1}{2} \left( M_{x} - M_{y} \right) \frac{dP}{dy'} \right] dx'$$
(5.5b)

$$V_{s}(90^{\circ}) = \int_{0}^{L} \frac{1}{\sigma} M_{y} \frac{dP}{dx'} dx'$$
(5.5c)

Several interesting aspects are evident through **Eqns. 5.2 - 5.4**, *e.g.*, the use of a pressure gradient implying<sup>128</sup> components of both longitudinal and transverse to the flow direction - from tensorial considerations, the fluid flux transforms covariantly<sup>130</sup>, *i.e.*, in the same way as the basis vectors, aligning the flow with the grooves<sup>131</sup>. Due to the existence of the side walls of the channel, the consequent recirculation<sup>128,131</sup> yields transverse components, with either positive/negative gradients  $\frac{dP}{dyr}$ . Since the experimental applied pressure is along x', the generated transverse  $\frac{dP}{dyr}$  will be positive. From an electrokinetic point of view, only when  $M_x=M_y$ , would the off-diagonal terms in Eqn. (5.2) be zero and yield a  $V_s$  ( $\theta = 45^\circ$ ) as the arithmetic mean of the  $V_s$  obtained for  $\theta = 0^\circ$  and  $\theta = 90^\circ$ . For instance, we obtain:

$$\frac{1}{2}[V_s(0^\circ) + V_s(90^\circ)] - V_s(45^\circ) = -\frac{L}{2\sigma}(M_x - M_y)\frac{dP}{dy'}$$
(5.6)



**Figure 5.4** Calculated eigenvalues of the electrokinetic mobility tensor (M), i.e., Mx and the My, see eq 2 in the text, estimated from streaming potential (Vs) measurements for an AFS, castor oil filled LFS, and GPL oil filled LFS (at 0.1 mM electrolyte concentration,  $\sigma = 0.001$  S/m,  $\frac{dP}{dx'} = 7627$  Pa/m).

Such considerations may be used to explain the related experimental observations in **Figure 5.3**. The estimates of the  $M_x$  and the  $M_y$  were estimated experimentally through measuring the streaming potential for two orientations of the grooves with respect to the flow direction, *i.e.*,  $M_x$  from  $V_s$  ( $\theta = 0^\circ$ ) using **Eqn. 5.5(a)**, and  $M_y$  from  $V_s$  ( $\theta = 90^\circ$ ) using **Eqn. 5.5**©. The respective values are listed in **Figure 5.4**. Since  $M_x - M_y$  is negative,  $\frac{dP}{dy'}$  is positive, the  $V_s$  of 45° is smaller than the arithmetic mean of that of 0° and 90°.


**Figure 5.5.** Variation of the Vs with groove orientation angle ( $\theta$ ) for (a) an AFS ( $\theta_{min} \approx 45.9^{\circ}$  with corresponding  $V_{s,min} \approx 25.1 \text{ mV}$ ) (b) castor oil filled LFS ( $\theta_{min} \approx 52.2^{\circ}$  with corresponding  $V_{s,min} \approx 10.8 \text{ mV}$ ), and (c) GPL oil filled LFS ( $\theta_{min} \approx 67.5^{\circ}$  with corresponding  $V_{s,min} \approx 18.9 \text{ mV}$ ). The Vs was measured at 900 Pa with the flow of 0.1 mM NaCl. Tensorial effects and transverse pressure gradients were invoked to explain the observed variations and the theoretical Vs minima (orange star).

Actually, 
$$\frac{dP}{dy'}$$
 is a function of  $\theta$  and is in the form<sup>128</sup> of  $\frac{dP}{dy'}(\theta) = c * sin\theta cos\theta$ . The values of  $\frac{dP}{dy'}$ 

at 45° was estimated using Eqn. (5.6), based on the obtained values of  $M_x$  and  $M_y$  and the measured  $V_s$  at  $\theta = 0^\circ$ , 45° and 90°, and  $\frac{dP}{dy'}(\theta)$  could be obtained. The dependence of  $V_s$  on  $\theta$  was derived via plugging  $\frac{dP}{dy'}(\theta)$  into Eqn. (5.4). It was observed that the variation of the  $V_s(\theta)$  seems to exhibit a minimum, say at  $\theta_{min}$  shown in **Figure 5.5**. From Table I, Eqns. (4) and (5), and a negative  $\frac{dP}{dx'}$  (as the pressure decreases in the flow direction), it may be deduced that such variation would essentially arise from the  $\frac{dP}{dy'}$  term – which may be negative or positive and is dependent on  $\theta$ . It was found that the  $\theta_{min}$  was less sensitive to a change of  $M_x$  or  $M_y$  but is decreasing with  $\frac{dP}{dy'}$  and is set to 45° when  $\frac{dP}{dy'}$  is large enough, and  $V_s(\theta)$  exhibits a  $sin^2(2\theta)$  form. This is because that the  $V_s(\theta)$  is composed of the diagonal term contribution and off-diagonal term contribution. The diagonal term in equation (5.5) is  $\int_0^L [\frac{1}{\sigma}(M_x \cos^2\theta + M_y \sin^2\theta) \frac{dP}{dx'}]dx'$  which monotone decreases with  $\theta$ , while the off-diagonal term  $\int_0^L [\frac{1}{\sigma}(M_x - M_y)sin\theta \cos\theta \frac{dP}{dy'}]dx'$  is minimum at  $\theta = 45^\circ$ . When the magnitude of  $\frac{dP}{dy'}$  increases, the off-diagonal term will become more dominant, and  $\theta_{min}$  will approach to  $45^\circ$  with increasing  $\frac{dP}{dy'}$ .

## **5.5** Conclusion

In summary, we have indicated geometrically induced modulation of the Vs obtained through electrolyte flows over specifically oriented air and liquid filled surfaces, described through tensorial considerations. Such engineering could find application in the establishment of local potentials and electric fields. Further experiments on electrolyte flow in microchannels with engineered hybrid surfaces would yield insights into electrokinetics beyond smooth walled channel flow. Such tunability of the Vs arises may be related to a variation in the local and effective slip characteristics as well as the surface charge density of the AFS and LFS and the presented work offers new avenues into exploring such effects. Moreover, related investigations would yield insights into the generation of transverse pressure gradients due to anisotropic surfaces, surface heterogeneities and roughness, with implications to the efficiency of energy conversion and lab on a chip devices.

## 5.6 Acknowledgement

This chapter, in full, is a reprint of the materials as it appears in Fan, B. and Bandaru, P.R., Tensorial modulation of electrokinetic streaming potentials on air and liquid filled surfaces, Langmuir, 2019, 35, 46, 14812 – 14817. The dissertation author was the primary investigator and author of this paper. The authors are grateful for support from the National Science Foundation (NSF: CBET 1606192). We also appreciate the assistance of Prof. J. Friend and P. Chen for help with imaging.

## **Chapter 6 Future Work**

Though there is a long history for electrokinetics, the development of micro/nano fabrication makes electrokinetic phenomenon revive and there are still a lot to be investigated, both in fundamental research and practical application. Especially, the electrokinetics on liquid-filled surface is a quite new multi-physics coupling phenomenon, which make it worthy to continue to study this phenomenon. Here, I proposed the following future directions.

Influences of geometrical factors of liquid-filled surfaces on  $V_s$ . The geometrical parameters, such as groove width, groove depth and groove fraction, play an important role in tuning the fluid slip characters and zeta potential ( $\zeta$ ) of LFS, which will influence the generated  $V_s$ . I tried to vary the groove width, groove depth and groove fraction with different values and measured the corresponding  $V_s$ . Generally speaking, the  $V_s$  of LFS increases with groove depth and decreases with groove width and groove fraction. However, it is difficult to find a quantitative relationship between the measured  $V_s$  and these geometrical parameters. In the future, simulation and theoretical analysis will be performed to find the quantitative relationship.

Electrokinetics on LFS with 2D pattern. Till now, all the work on electrokinetics on LFS focused on the 1D groove geometry. 2D geometries, such as post and hole geometry, have different fluid slip characteristics compared with groove pattern. For example, the fluid slip for strip geometry scales with solid fraction  $as^{29} b_{strip} \sim -Llog(\phi_s)$ , while the post geometry scales  $as^{29} b_{post} \sim \frac{L}{\sqrt{\phi_s}}$ . In addition, the interfacial fluid dynamics and robustness of the LFS will also be changed. In post geometry, the filled oil in the interstices will be interconnected and can flow in two directions, while for hole geometry, the filed oil inside the holes will be isolated and confined in the holes. The difference in oil movement between groove, post and hole geometries will greatly influence the electrolyte slip, electrolyte-oil interface dynamics, the robustness of LFS and the induced electrokinetics. In the future, expansion to electrokinetics on LFS with 2D geometry will be performed both experimentally and theoretically to investigate the corresponding fluid slip, streaming potential, etc, and those results will be compared with those of 1D geometry.

**Nanoscale electrokinetics.** I have pioneered in studying electrokinetic flow in microscale in my Ph. D period. The nanofluidic phenomena will be far different from microscale flow due to the dominant surface force effects and electrical double layer overlap. Nano electrokinetics is important in energy conversion, single molecule analysis and water purification. It may be possible to obtain voltages of the order of 1 volt through the use of nanoscale fluid channels. The fluid transport in *LFS* composed nanochannel and interfacial fluid dynamics induced by the effects of liquid-liquid/solid interface energy of the hybrid *LFS* will be investigated both experimentally and theoretically. In addition, I would also like to theoretically study the ion transport in the electrolyte flowing over *LFS* with nonhomogeneous surface charge density and interface energy. The streaming potential and energy conversion efficiency will also be experimentally measured and theoretically analyzed based on the ion transport behavior to provide theoretical guidance for designing electrokinetic power source. Besides the fundamental study of the underlying physics of nano electrokinetics on *LFS*, the practical nanoscale electrokinetics based power source will be developed and the energy conversion efficiency will be optimized based on the theoretical guidance to facilitate the practical application of electrokinetics.

**Electricity harvesting through solar desalination process.** A novel solar heat localization method using porous solar absorber has been used to enhance the evaporation efficiency. The aim has been to utilize solar energy in novel ways for water purification and desalination. Porous solar absorbers are usually composed by an array of micro/nano channels and EDL can form at the interface of salt water and solar absorber inner wall surfaces. Through nanoscale channels or tuning the surface charge density, only positive or negative ions can go through the channels. This charge separation process, which is similar to the working mechanism of electrokinetics, can prevent the formation of salt crystal and generate electricity

at the same time. Due to the imbalance of positive/negative ion concentration between the bottom and top sides of the porous solar absorber, the desalination system can generate electricity to harvest the electrokietic energy at the same time which would make better use of the incident solar energy.

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