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Recognizing the Less Explored “Active Solid”—“Moving Liquid” Interfaces in Bio/Chemical Sensors
Nikhil Bhalla*

ABSTRACT: Bio/chemical sensors possess a plethora of advantageous features that have proven to be invaluable tools in detecting and monitoring biomolecules, facilitating advancements in healthcare, environmental monitoring, food safety, and more. However, when it comes to their routine use in continuous fluid flow conditions, an intricate web of solid—liquid interfacial phenomena emerges, which requires a deep understanding of the sensor surface and fluid interactions. These interfacial phenomena encompass a broad spectrum of physical, chemical, and biological processes, other than the actual detection, that influence the sensor’s response. In this context, perhaps exploring a new theme “active solid”—“moving liquid” interface will unleash the full potential of bio/chemical sensors in any flow-based application.

KEYWORDS: solid—liquid interfaces, biosensors, transducers, continuous flow, plasmonics, electrochemistry

Performing bio/chemical sensing in continuous fluid flow conditions can alter the dynamics of a chemical reaction under investigation, leading to variations in reaction rates or binding activity at different fluid flow rates. On the other hand, the transducer element of a bio/chemical sensor is intrinsically sensitive to external stimulus applied on the sensor surface. This means that any bio/chemical sensor will reveal changes in its output signal when disturbed by liquid, in static or continuous flow condition, that is in contact with the sensor surface. Therefore, it is important to isolate the physical effects, such as friction from the liquid, liquid-slips, contact electrification, or local temperature changes that could be occurring on the solid surface of the sensor, other than the actual bio/chemical detection event. Moreover, such physical effects are immensely augmented in the dynamic state of the sensor, i.e., when the fluid is continuously moving on the surface, and therefore such a system requires a much deeper understanding of effects associated with moving fluid/flow rates on the sensor response.

To isolate such effects and especially in continuous fluid flow conditions, it is absolutely essential to investigate the solid—liquid interfacial phenomenon arising due to the interaction of the sensor surface with the liquid in contact with it. However, a couple of questions which then arise are (1) how can one isolate the effects of this solid–liquid interaction on the surfaces of bio/chemical sensors and (2) are there mathematical or physical models in literature that one should incorporate in the equations of sensors which will exclusively relate output of the sensor to the bio/chemical detection activity? To address both of these questions, the development of new models that quantify the interaction between the sensor surface and the moving liquid is imperative. Once these new models are established, the equations describing the transduction mechanisms of bio/chemical sensors must be modified (to incorporate the effect of moving fluid) for yielding true detection signals for the bio/chemical event happening under the continuous fluid flow conditions.

Traditionally, several routinely used continuous fluid flow based bio/chemical sensors such as quartz crystal microbalance (QCM) and surface plasmon resonance (SPR) have neglected the effect of continuous moving fluid on their surfaces. As a result, these sensors have reported different results for parameters which indicate binding kinetics. This is because wavelength/frequency changes within such sensors are interpreted exclusively with mass change on the sensor surface, irrespective of the flow conditions. The repercussion of this slender cognizance of solid–liquid interfacial phenomenon is manifested in the form of overestimation/underestimation of mass which binds on the surface of the sensor. There are several examples of various transducers which are reported to overestimate the amount of analyte detected when correlated with the equations/models that govern their signal response.

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Nevertheless, recent works have discovered events such as “liquid slip” previously totally ignored in the response of QCM sensor.

Along similar lines, it remains to be understood if such effects exist in a range of electrochemical and optical sensors that measure binding kinetics in continuous flow conditions of the fluid.

**IMPACT OF IGNORING THE MOVING FLUID AND SENSOR SURFACE INTERACTION**

There is a profound impact of neglecting the effect of continuous moving fluid in sensors, and it may urge researchers to reassess the analyte’s “fraction bound” on the sensor surface when it is considered proportional to the sensor signal. This is because most of the analyte fraction bound is presented in the form of $K_d/K_{on}/K_{off}$ which are measured in continuous flow conditions. Here, $K_{on}$ is a constant used to indicate how fast the analyte binds to its target, while $K_{off}$ is a constant used to specify how fast the analyte dissociates from its target. The $K_d$ is simply a ratio of $K_{on}/K_{off}$ which typically quantifies the affinity or strength of binding of a single analyte to its target.

It is well-known that these reaction rate constants are dependent on the conditions such as the buffer pH, temperature of bioentities and the buffer, and the composition of the buffer itself. For instance, experiments conducted with Biacore SPR system at different temperatures (ranging from 20 °C to 37 °C) yield different binding rates after the SPR sensorgrams are analyzed. The conundrum of such a sensor outcome is to precisely isolate the effects of temperature on the transducer response from the thermodynamic effect of the binding interaction. In parallel, if the transducer is sensitive to some of the aforementioned conditions such as the pH and buffer content (type of ions in the buffer), the response of the sensor becomes more complicated to reveal the exclusive contribution of the binding event. As mentioned earlier, such effects are amplified in the dynamic state of the sensor, i.e., when fluid is moving, which requires a much deeper understanding of fluid flow rate effects on the sensor response.

Significant backlash of limited understanding of such solid—liquid interfacial phenomenon on sensor surfaces has also led to neglect of fluid flow rates in determination of binding constants within the academic community. For example, within the experiments conducted in the work of MacDonald et al., a flow rate of 30 μL·min$^{-1}$ for SPR measurements and a flow rate of 50 μL·min$^{-1}$ for QCM measurements was used to elucidate binding events. Thereafter, both SPR and QCM results were directly compared with each other in the context of binding events. In this case, unless the flow rates were kept the same, signals from SPR and QCM cannot be directly compared without caution. Additionally, some works in the past have also claimed enhanced sensor sensitivity by simply observing larger...
sensor signal output at specific flow rates without considering the exclusive effect of flow on the sensor surface.11

Another major impact of not considering the physical effects of continuous flow rate is that different industries selling the same chemical (especially in the case of proteins, enzymes) may report different \( K_d/K_{on}/K_{off} \) values in their data sheets without providing much information on the fluid flow rates and the type of instrument used for computing the binding constants. Typically, a molecule may not have different binding constants when evaluated with the same instruments unless tested in different flow rates. This may result in academic researchers or other clients of the company buying reagents which show higher binding affinity. This might be beneficial to a company from a commercial perspective, but researchers buying the reagents may want to investigate how binding constants are shared in Table 2.

### Table 2. Definitions of Solid–Liquid Interfacial Processes

<table>
<thead>
<tr>
<th>Term</th>
<th>Definition</th>
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<tbody>
<tr>
<td>1. Friction</td>
<td>Damping force or a dissipative force due to interplay of viscosity of the liquid and the roughness of the solid surface.</td>
</tr>
<tr>
<td>2. Shear force</td>
<td>Force acting on the moving fluid parallel to the solid surface.</td>
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<tr>
<td>3. Liquid slip/slip length</td>
<td>The slip length, more often used to describe the liquid slip is the distance beyond the solid–liquid interface where the liquid velocity linearly extrapolates to zero.</td>
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<tr>
<td>4. Contact electrification</td>
<td>A phenomenon whereby solid surfaces become electrically charged upon contact with the liquid.</td>
</tr>
<tr>
<td>5. Fluid inertia</td>
<td>The solid like behavior of the moving liquid’s first few layers when in contact with the solid.</td>
</tr>
<tr>
<td>6. Surface tension</td>
<td>The virtue of liquid at rest to shrink into the minimum surface area possible when in contact with the solid surface.</td>
</tr>
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In the context of the aforementioned bio/chemical sensing in continuous fluid flow conditions, a value proposition for introduction of a new term for sensor surfaces, i.e., “active solids” instead of simply solids, is created; for details, see Figure 1 and Table 1. This is because a sensor surface is in an active state compared to a simple solid without any external sensor. See Figure 1a and b, demonstrating a simple solid (inactive state) in contact with liquid in static and moving state, respectively. In contrast, Figure 1c shows an electrically active solid, Figure 1d,e shows optically vibrating SPR/LSPR solid, respectively, and a mechanically vibrating active solid in the form of QCM is shown in Figure 1f. Note that in Figure 1c–f the interface between the active solid and the moving liquid is also shown. The moving liquid requires a completely new recognition of solid–liquid interfacial interactions in the fluid dynamics community where solids are usually modeled in a recessive state, unlike solids within transducers which are in active state (such us in a vibrating state). Henceforth, scrutinizing these effects at “active solid”—“moving liquid” interface should yield development of new models/equations which will precisely govern how bio/chemical sensors operate in flow. This may as well lead to a new research field of “active solid–moving liquid interface”. Some physical effects occurring at the interface of the “active solid”—“moving liquid” interface are shared in Table 2.

A recent example of such a study is demonstrated by Payam et al. Here, the authors discover a “missing liquid slip” in the
vibrating solid–liquid systems such as the QCM-based biosensor (Figure 2a shows their system) where traditionally

frequency and dissipation changes were interpreted exclusively with mass change on the sensor surface, irrespective of the flow conditions. To some extent the study also argued the origin of the liquid slip by correlating the slip to a large extent with the flow rate of the liquid, rather than the surface wetting conditions. This was in contrast with the understanding of the liquid slip, where it is exclusively associated with the degree of surface wetting. For instance, a hydrophobic surface is well-known to provide a higher slip than a hydrophilic surface. While the experiments and theory within their work agreed to this concept, the work clearly demonstrated that the origin of slip is attributed more to the movement of the liquid. It can be concluded that with a change in the flow rate of the liquid, the slip changes to a much larger extent. The wetting condition then is simply a slip modulation factor. The second important finding was that the properties of the solid did not fully determine the slip of the Newtonian liquids. This means that hydrophobic surfaces are not slip-free. Slip values (in magnitude) comparable to hydrophobic surfaces can exist on active/vibrating solid surfaces at flow rates which are well within the laminar regime, as shown in Figure 2b. This finding is important for interpreting bio/chemical sensing results in flow. Additionally, the effects of liquid slip on energy dissipation and inertia were elucidated. A nonlinear solid-like behavior of the liquid was demonstrated in terms of inertial length. Essentially, the work has put forward the case that the first few layers of the liquid on a solid have mechanical characteristics like a solid, and this so-called “inertial length” increases as the flow rate increases (meaning more liquid layers are solid-like at higher flow rates). Such effects would completely change the perspective of how bio- and chemical sensor responses are interpreted in flow. Overall, the findings presented a new mathematical expression for liquid slip in the form of a complex number. The real part of the expression demonstrates the effect of flow (modulation in liquid properties), and the condition of the active solid surface (modulation of solid properties) is reflected by the imaginary part of the complex number.

CONCLUDING REMARKS

Going forward, more of such works are required to understand active solid-moving liquid interactions. This will eventually revolutionize the current biosensing measurements, where most often the sensor response is associated with a single stimulus. Furthermore, it will also provide more informed evidence of binding kinetics of biomolecules in flow instead of directly associating the sensor response (including the sensitivity of the sensor) with binding events, irrespective of the flow conditions. The holistic impact of such studies on the bio/chemical sensing community will be reflected in presenting a much stronger case for the adoption of bio/chemical sensors as reliable and routine tools in widespread applications ranging from healthcare, environmental, and energy domains.

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Notes

The author declares no competing financial interest.

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Figure 2. Liquid slip in active solid−moving liquid interface: (a) shows an experimental setup consisting of vibrating solid, microfluidic system, readout, and control electronics for studying effect of continuous fluid flow on the quartz crystal microbalance. (b) shows how the magnitude of liquid slip length changes with change in the fluid flow rates when the surface of QCM is treated with MPTS-octyl silane, octyl silane, MPTS, and plasma. It also includes the untreated QCM surface. Both subfigures are reprinted with permission from ref 6. Copyright (2022) Springer Nature.
