Polarization-Dependent Heterodyne-Detected Sum-Frequency Generation Spectroscopy as a Tool to Explore Surface Molecular Orientation and Ångström-Scale Depth Profiling

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ABSTRACT: Sum-frequency generation (SFG) spectroscopy provides a unique optical probe for interfacial molecules with interface-specificity and molecular specificity. SFG measurements can be further carried out at different polarization combinations, but the target of the polarization-dependent SFG is conventionally limited to investigating the molecular orientation. Here, we explore the possibility of polarization-dependent SFG (PD-SFG) measurements with heterodyne detection (HD-PD-SFG). We stress that HD-PD-SFG enables accurate determination of the peak amplitude, a key factor of the PD-SFG data. Subsequently, we outline that HD-PD-SFG can be used not only for estimating the molecular orientation but also for investigating the interfacial dielectric profile and studying the depth profile of molecules. We further illustrate the variety of combined simulation and PD-SFG studies.

I. INTRODUCTION

The arrangement of interfacial atoms and molecules in a nanometer thickness region governs the material properties, atmospheric chemistry, chemical reactions as well as (bio)-molecular processes. Understanding these processes requires knowledge of the structure of the molecules at interfaces. Among a number of the surface-specific techniques, including atomic force microscopy, X-ray spectroscopy, tip-enhanced Raman spectroscopy, and second-harmonic generation spectroscopy, vibrational sum-frequency generation (SFG) spectroscopy is a unique tool, because it allows probing the molecules at the soft matter interfaces with molecular specificity.

Vibrational SFG spectroscopy is a second-order nonlinear optical technique, and its signal is generated by the infrared (IR) and visible pulses. The SFG signal is enhanced when the IR frequency matches the frequency of vibrational mode, providing molecular specificity. The observable is the complex $\chi^{(2)}$ spectrum, where the imaginary and real parts of $\chi^{(2)}$ spectrum represent the absorption and dispersion of the molecular response, respectively. The even-order response excludes the contribution from the centrosymmetric medium, i.e., from the bulk. As such, SFG allows us to probe the interfacial molecular responses selectively, not only at solid interfaces but also at soft matter interfaces.

By carrying out SFG measurements at different polarization combinations of the IR, visible, and SFG beams, one can obtain information on the orientation of the molecular moiety of proteins, water, and organic compounds. Furthermore, recent studies showed that polarization-dependent (PD)-SFG has the potential to address the nature of interfacial dielectric medium and can provide depth information on the interfacial molecules. The analysis of PD-SFG requires the ratio of the peak amplitude in the Im $\chi^{(2)}$ spectra at different polarization combinations. An essential technique for accurate estimation of the Im $\chi^{(2)}$ peak amplitudes is the heterodyne detection of SFG (HD-SFG) signal, because HD-SFG can directly access the Im $\chi^{(2)}$ spectrum, unlike the conventional, homodyne-detection of SFG, which provides the $|\chi^{(2)}|^2$ spectrum. In particular, accurate phase determination in HD-SFG allows us to obtain the peak amplitudes.

In this perspective, we explain the fundamentals of the PD-HD-SFG spectra analysis and outline the research topics that the PD-HD-SFG spectroscopy can explore together with the theoretical modeling. In Section II, we introduce the principles...
II. POLARIZATION-DEPENDENT HETERODYNE-DETECTED SFG (PD-HD-SFG)

II.A. Principles of PD-HD-SFG. A number of the papers account for the measurement and processing of the HD-SFG data. Here, we explain how to take the different polarizations into account. Let us assume that we have the measured SFG spectra of the sample and the z-cut quartz (denoted as $\chi^{(2)}_{abc,\text{measured sample}}$ and $\chi^{(2)}_{abc,\text{measured quartz}}$, respectively) at the abc polarization for abc = spp, ppp, and sps, where abc polarization represents the a-polarized SFG, b-polarized visible, and c-polarized IR beams. The effective SFG spectra at the abc polarization is given by:

$$\chi^{(2)}_{abc,\text{eff}} = \frac{\chi^{(2)}_{abc,\text{measured sample}}}{\chi^{(2)}_{abc,\text{measured quartz}}} r_{\text{sample,abc}} r_{\text{quartz,abc}} \chi^{(2)}_{abc,\text{eff,quartz}}.$$  

(1)

Here, $r_{\text{sample,abc}}$ and $r_{\text{quartz,abc}}$ are the reflectivity coefficients of the local oscillator signal at the z-cut quartz and the sample surfaces for the a-polarized SFG beam frequency, respectively. $\chi^{(2)}_{abc,\text{eff,quartz}}$ is the effective susceptibility of the z-cut quartz. Assuming the crystal coordinate is overlapped with the lab coordinate, then $\chi^{(2)}_{abc,\text{eff,quartz}}$ is given by:

$$|\chi^{(2)}_{abc,\text{eff,quartz}}| = 2 \cos \beta_{IR} L_{jy} (\omega_{SFG}) L_{yy} (\omega_{Vis}) L_{xx} (\omega_{IR}) \chi^{(2)}_{abc,\text{eff,quartz}}.$$  

(2)

The beam configuration of the SFG measurement is displayed in Figure 1(a). Note that the dielectric constant of the interfacial layer, i.e., interfacial dielectric constant, $\epsilon'$ is known and should be obtained from the model calculation. The beam configuration of the SFG measurement is displayed in Figure 1(a). Note that the dielectric constant of the interfacial layer, i.e., interfacial dielectric constant, $\epsilon'$ is known and should be obtained from the model calculation. The beam configuration of the SFG measurement is displayed in Figure 1(a). Note that the dielectric constant of the interfacial layer, i.e., interfacial dielectric constant, $\epsilon'$ is known and should be obtained from the model calculation. The beam configuration of the SFG measurement is displayed in Figure 1(a). Note that the dielectric constant of the interfacial layer, i.e., interfacial dielectric constant, $\epsilon'$ is known and should be obtained from the model calculation.
\[
(\chi^{(2)}_{ppp})_{\text{eff}} \approx -L_{xx}(\omega \text{FG})L_{xx}(\omega \text{Vis})L_{zz}(\omega \text{IR})\cos \beta_{\text{SGF}}\cos \beta_{\text{Vis}} \\
\times \sin \beta_{\text{IR}}(\chi^{(2)}_{xxz}) + L_{zz}(\omega \text{FG})L_{xx}(\omega \text{Vis})L_{zz}(\omega \text{IR}) \\
\times \sin \beta_{\text{SGF}}\sin \beta_{\text{Vis}}\sin \beta_{\text{IR}}(\chi^{(2)}_{zzz})
\]

\[
(\chi^{(2)}_{ppp})_{\text{eff}} = L_{yy}(\omega \text{SGF})L_{zz}(\omega \text{Vis})L_{yy}(\omega \text{IR})\sin \beta_{\text{IR}}(\chi^{(2)}_{xyy})
\]

Note that eq 9 is valid when the \(\chi^{(2)}_{xxz}\) and \(\chi^{(2)}_{zzz}\) components are negligibly small compared to the \(\chi^{(2)}_{xxz}\) and \(\chi^{(2)}_{zzz}\) components. For a case where \(x\) and \(y\) axes are indistinguishable (as for liquid interfaces), we have \(\chi^{(2)}_{yyz} = \chi^{(2)}_{xyy}\). Under the situation that the vibrational relaxation is slower than the rotational motion of vibrational chromophores (slow-motion limit), the peak amplitudes of the vibrational modes with the C\(_{\text{oo}}\) symmetry group (such as the free O–H group of water, and C–H and C==O stretches of formic acid) in the \(\chi^{(2)}_{yyz}\) and \(\chi^{(2)}_{xyy}\) spectra (denoted as \(A^{(2)}_{yyz}, A^{(2)}_{xyy}, A^{(2)}_{zzz}\), and \(A^{(2)}_{xzz}\), respectively) are connected via the following forms:

\[
\frac{A^{(2)}_{yyz}}{A^{(2)}_{xyy}} = \frac{(1 - r)(\cos \theta) - (\cos^3 \theta)}{(1 + r)(\cos \theta) - (1 - r)(\cos^3 \theta)}
\]

\[
\frac{A^{(2)}_{zzz}}{A^{(2)}_{xzz}} = \frac{(1 + r)(\cos \theta) - (1 - r)(\cos^3 \theta)}{2r(\cos \theta) + 2(1 - r)(\cos^3 \theta)}
\]

where \(r\) denotes the depolarization ratio for the target vibration, and \(\theta\) represents the angle formed by the molecular axis and surface normal. Equations 11 and 12 are the key equations for orientational analysis and depth analysis.

The depolarization ratio of \(r\) has been obtained either from the Raman data\(^{41-43}\) or from \(ab\) \(initio\) calculations.\(^{44-46}\) The recent development of \(ab\) \(initio\) calculation allows us to obtain the \(r\) value directly.

III. IMPACT OF INTERFACIAL DIELECTRIC CONSTANT

Here, we examine the impact of the interfacial dielectric constant by using the SFG spectra at the air–water interface based on our recent report.\(^{29}\) The \(\chi^{(2)}\) and \(\chi^{(2)}_{\text{eff}}\) spectra are displayed in parts b and c of Figure 1, respectively. The signs of the responses contained in the Im \(\chi^{(2)}\) spectra reflect the absolute orientation of molecular moieties; for the O–H stretch, for example, the positive (negative) sign of the peak indicates that the O–H group points \(\text{up to the air (down to the bulk water)}\).\(^{47,48}\) Specifically, the Im \(\chi^{(2)}_{\text{eff}}\) spectrum shows the positive ~3700 cm\(^{-1}\) peak and the negative band ranging from 3000 to 3500 cm\(^{-1}\), indicating that the danging O–H group points toward the air, while the hydrogen-bonded O–H group points toward the bulk.\(^{49-53}\)

To convert the \(\chi^{(2)}_{\text{eff}}\) and \(\chi^{(2)}_{\text{eff}}\) spectra into the \(\chi^{(2)}_{\text{eff}}\) and \(\chi^{(2)}_{\text{eff}}\) spectra, two approaches have been commonly adopted to describe the interfacial dielectric constant \(\varepsilon\): fully embedded model (Lorentz model) and half-embedded model (Slab model).\(^{54}\) In the Lorentz model, where the vibrational chromophores are fully solvated (Figure 1d), \(\varepsilon = \infty\). In the half-embedded model, the vibrational chromophores are half-solvated (Figure 1e), and the interfacial dielectric constant can be expressed by \(\varepsilon = \frac{\varepsilon(\varepsilon + 5)}{2(\varepsilon + 1)}\). Parts f and g of Figure 1 display Im \(\chi^{(2)}_{\text{eff}}\) and Im \(\chi^{(2)}_{\text{eff}}\) spectra with the fully- and half-embedded model of \(\varepsilon\). The comparison between the simulated and experimental Im \(\chi^{(2)}\) spectra indicates that neither the Lorentz model nor the Slab model provides satisfactory agreement. The Lorentz model overestimates the 3700 cm\(^{-1}\) peak amplitude in the Im \(\chi^{(2)}_{\text{eff}}\) spectrum, while the Slab model underestimates the amplitude of the negative 3100–3500 cm\(^{-1}\) band in both Im \(\chi^{(2)}_{\text{eff}}\) and Im \(\chi^{(2)}_{\text{eff}}\) spectra. Such disagreement between the experimental data and simulation data indicates that the choice of models has a strong impact on the concluded molecular response inferred from experimental Im \(\chi^{(2)}\) spectra.

Next, we examine the impact of the model of the interfacial dielectric constant \(\varepsilon\) on the values of \(A^{(2)}_{yyz}/A^{(2)}_{xyy}\) and \(A^{(2)}_{zzz}/A^{(2)}_{xzz}\) in the C–H stretch mode (~2900 cm\(^{-1}\)) region by assuming that a single formic acid molecule is located at the air–water interface, and thus, the dielectric profile is governed by water.\(^{55}\) Here, we set the incident angles of 64° and 50° for visible and IR beams, respectively, and the visible wavelength of 800 nm, eqs 11 and 12 can be recast as:

\[
A^{(2)}_{yyz}/A^{(2)}_{xyy} = 0.83A^{(2)}_{eff,yy}/A^{(2)}_{eff,xyy}
\]

in the half-embedded model, while they are given by:

\[
A^{(2)}_{zzz}/A^{(2)}_{xzz} = 0.78A^{(2)}_{eff,zz}/A^{(2)}_{eff,xzz}
\]

in the fully-embedded model. The comparison of eqs 13 and 14 and eqs 15 and 16 clearly shows that \(A^{(2)}_{yyz}/A^{(2)}_{xyy}\) is much less sensitive to the choice of model to describe the interfacial dielectric constant, than \(A^{(2)}_{zzz}/A^{(2)}_{xzz}\).

IV. MULTIDIRECTIONAL ANALYSIS AND (MULTIDIRECTIONAL) ORIENTATION DISTRIBUTION

IV.A. Principles of Orientational Analysis. As is seen in eqs 11 and 12, one can obtain \(\langle \cos \theta \rangle/\langle \cos^3 \theta \rangle\) from either \(A^{(2)}_{yyz}/A^{(2)}_{xyy}\) or \(A^{(2)}_{zzz}/A^{(2)}_{xzz}\) while it is highly recommended to use \(A^{(2)}_{yyz}/A^{(2)}_{xyy}\) rather than \(A^{(2)}_{zzz}/A^{(2)}_{xzz}\) for obtaining \(\langle \cos \theta \rangle/\langle \cos^3 \theta \rangle\), because \(A^{(2)}_{zzz}/A^{(2)}_{xzz}\) is practically insensitive to the interfacial dielectric constant, as is discussed in Section III. Since the quantity \(\langle \cos \theta \rangle/\langle \cos^3 \theta \rangle\) does not directly provide the physical insights into the orientation, one may want to convert \(\langle \cos \theta \rangle/\langle \cos^3 \theta \rangle\) to \(\langle \theta \rangle\). To do so, one needs to assume the orientational distribution function \(f(\theta_1, \ldots, \theta_n)\). The ensemble average of B is given by:

\[
B = \frac{\int_{0}^{\infty} f(\theta_1, \ldots, \theta_n)\sin \theta_1 \cdots \sin \theta_n \sin \theta_1 \cdots d\theta_n}{\int_{0}^{\infty} f(\theta_1, \ldots, \theta_n)\sin \theta_1 \cdots \sin \theta_n \cdots d\theta_n}
\]

Below, we consider a one-dimensional orientational distribution function \(f(\theta)\) for simplicity. So far, four distinct orientation distribution functions have been assumed: the rectangular function (eq 18),\(^{54-56}\) the Gaussian-shaped function (eq 19),\(^{57-59}\) the delta function (eq 20),\(^{15,19,60-62}\) and the exponential decay function (eq 21):\(^{55}\)

\[
f_{\text{ex}}(\theta) = \begin{cases} 1 & \text{for } 0 \leq \theta \leq \theta_b \\ 0 & \text{for } \theta_b < \theta < \pi \\ \end{cases}
\]
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Figure 2. (a) $\langle \cos \theta \rangle / (\cos^3 \theta)$ vs. $\langle \theta \rangle$ based on different distribution functions $f_G(\theta)$, $f_E(\theta)$, $f_D(\theta)$, and $f_f(\theta)$, $f_{2f}(\theta)$ employs $\sigma = 15^\circ$. (b) Orientational distribution of the free O–H group of water with respect to the surface normal at the air–water interface obtained from the classical MD simulation. Adapted with permission from ref 45. Copyright 2018 American Physical Society. (c) Orientational distributions of C–H stretch of formic acid $^{22}$ and C–H symmetric stretch of methanol $^{24}$ and acetonitrile. $^{26}$ Definition of the angle $\theta$ formed by the molecular axis and the surface normal. Adapted with permission from refs 22 (Copyright 2022 AIP Publishing), 46 (Copyright 2017 Royal Society of Chemistry), and 64 (Copyright 2015 American Chemical Society), respectively.

$$f_G(\theta) = \frac{1}{\sqrt{2\pi}\sigma_G^2}e^{-(\theta - \theta_0)^2/2\sigma_G^2}$$

$$f_D(\theta) = \delta(\theta - \theta_0)$$

$$f_E(\theta) = e^{-\theta/\theta_c}$$

Because the choice of the orientational distribution function critically affects the inferred molecular orientation, $^{45,46,52}$ as illustrated in Figure 2a, it is important to check the shape of $f(\theta)$ with MD simulations (with multiple force field models in the classical MD simulation) $^{45}$ or accurate MD techniques including ab initio MD (AIMD). $^{22,63}$ In fact, the shapes of $f(\theta)$ computed by MD simulation indicates that it is challenging to predict the functional form of $f(\theta)$.

Although these functions are approximations, and none of the functions can be used universally to obtain the interfacial molecules’ angular information, we have the following pointers based on previous studies.

1. Rectangular ($f_R(\theta)$) and exponential decay ($f_E(\theta)$) functions are the approximations on the basis of the broadness of the distribution, while the Gaussian ($f_G(\theta)$) and delta ($f_D(\theta)$) functions focus on the center angle.

2. $f_G(\theta)$ and $f_D(\theta)$ seem more appropriate for some large molecules at the liquid interfaces $^{22,56}$ or self-assembled molecules on solid surfaces $^{59}$ while the use of $f_E(\theta)$ and $f_f(\theta)$ appears more suitable for modeling the angle distribution of a small molecule $^{22,45}$.

IV.B. Free O–D Group of Water at Air–D$_2$O Interface. Here, we explain how an orientational analysis can be done using PD-HD-SFG, by revisiting the orientation of the free O–D group at the air-D$_2$O interface. $^{45}$ Note that we used $A_{xx}^{(2)}/A_{zzz}^{(2)}$ rather than $A_{yy}^{(2)}/A_{zzz}^{(2)}$, because the $A_{yy}^{(2)}$ contribution is extremely small. $^{41,45,57}$ Figure 3a shows the Im $\chi^{(2)}_{yy}$ and Im $\chi^{(2)}_{zz}$ spectra at the air-D$_2$O interface. Both spectra commonly show the dangling O–D stretch mode at $\sim$2730 cm$^{-1}$ and antisymmetric mode of water molecules with one weak donor bonded O–D at $\sim$2650 cm$^{-1}$. To extract the free O–D contributions in the Im $\chi^{(2)}_{yy}$ and Im $\chi^{(2)}_{zz}$ spectra (denoted as $A_{yy}^{(2)}$ and $A_{zzz}^{(2)}$), respectively, we fitted Gaussian lineshapes to the spectra. These fits provide the amplitude ratio of $A_{yy}^{(2)}/A_{zzz}^{(2)}$ of $\sim$0.42 for the dangling O–D stretch mode. Through eq 12 with $r = 0.15$, $^{45}$ we obtained $\langle \cos \theta / (\cos^3 \theta) \rangle \approx 1.52$. When we use $f_E(\theta)$ for the orientational distribution (eq 21), which resembles the distributions obtained from the MD simulation (see Figure 2b), $^{45}$ $\langle \cos \theta / (\cos^3 \theta) \rangle \approx 1.52$ provides $\langle \theta \rangle \approx 59^\circ$, as is seen in Figure 3b.

The broad exponential distribution function for the free O–D group shows that $\sim$20% of the free O–D groups at the air-D$_2$O interface point down to the bulk. The presence of the free O–D groups pointing down to the bulk can be ascribed to capillary waves causing surface roughness. While on the top and bottom...
of the capillary wave, a free O–D group typically points up, on the slope of the capillary wave, the free O–D groups have the tendency to point down. A snapshot of an MD simulation clearly captures this behavior of the free O–D group (Figure 3c). As such, due to the surface nanoroughness, the distribution of the free O–D groups becomes much broader and exponential shape.

IV.C. Formic Acid Molecule at the Air–Water Interface.

As is seen in Figure 2c, the molecular distribution of acetonitrile and formic acid molecules at the air–water interface cannot be described by any of the functions given in eqs 18–21. How should we extract the molecular orientation from the PD-HD-SFG data? The complicated distribution function often arises from the competing driving forces to stabilize the molecular structure at interfaces. For formic acid, the two oxygen atoms and one hydrogen atom generate the competing driving forces, i.e., multiple types of hydrogen bonds with water molecules. In such a case, a multidimensional orientational distribution function (or joint-probability function) should be considered, rather than an orientational distribution function as a function of a single orientation parameter. One can determine the multidimensional distribution function via the multimode SFG probe. Below, we outline how to extract the molecular orientation using the multidimensional orientational distribution functions by focusing on formic acid molecules at the air–water interface.

Figure 4. (a, b) Im $\chi^{(2)}_{yyz}$ and Im $\chi^{(2)}_{yzy}$ spectra in the C–H stretch mode (a) and C=O stretch mode (b) regions. The dotted lines represent the Gaussian lineshapes obtained from the fit, while the filled area represents the sum of the two Gaussians. (c, d) $A^{(2)}_{zzz}/A^{(2)}_{yzy}$ vs $\theta_{E,CH}$ and $\theta_{E,CO}$ for the C–H stretch mode (c) and the C=O stretch mode (d). The rainbow 3D curves represent the numerical data based on eq 11, while the gray planes represent the experimental values. (e) Lines obtained from the crossing of rainbow 3D curves and gray planes in parts c and d. The dotted lines represent the experimental error. (f) The 2D orientational distributions inferred from the crossing point of part e. (g) 2D orientational distribution obtained from the AIMD simulation. (h) Schematic of the average orientation of a formic acid molecule at the air–water interface. The blue arrow represents the surface normal. The black and green arrows represent the C → H and C → O vectors, respectively. Reprinted with permission from ref 22. Copyright 2022 AIP Publishing.
We assume that the multidimensional orientational distribution function for the formic acid molecule can be given as

\[ f(\theta_{\text{CH}}, \theta_{\text{CO}}) = \exp\left(-\frac{\theta_{\text{CH}}}{\theta_{\text{CH},\text{eff}}}ight) \exp\left(-\frac{\theta_{\text{CO}}}{\theta_{\text{CO},\text{eff}}}ight) \times g(\theta_{\text{CH}}, \theta_{\text{CO}}) \]

by assuming an exponential decay function, where \( \theta_{\text{CH}} \) is a parameter determining the steepness/width of the exponential decay function and \( g(\theta_{\text{CH}}, \theta_{\text{CO}}) \) represents the geometric constraint which \( \theta_{\text{CH}} \) and \( \theta_{\text{CO}} \) should satisfy. The term \( g(\theta_{\text{CH}}, \theta_{\text{CO}}) \) is needed because the orientations of the C–H group and C==O group are not independent for a formic acid molecule; the intramolecular H–C==O angle is \( \sim 120^\circ \).

The Im \( \chi_{yyz}^{(2)} \) and Im \( \chi_{zzz}^{(2)} \) spectra of the C–H and C==O stretch modes of interfacial formic acid are displayed in parts a and b of Figure 4, respectively. From these spectra, we determined the ratio of \( A_{yyz}^{(2)}/A_{zzz}^{(2)} \) to be 0.60 \( \pm \) 0.01 for the C–H stretch mode and 0.36 \( \pm \) 0.01 for the C==O stretch mode. By using eqs 11, 17, and 22, we can determine the parameter of \( \theta_{\text{CH},\text{eff}} \) and \( \theta_{\text{CO},\text{eff}} \). Parts c and d of Figure 4 display the \( A_{yyz}^{(2)}/A_{zzz}^{(2)} \) values calculated for various \( \theta_{\text{CH},\text{eff}} \) and \( \theta_{\text{CO},\text{eff}} \) via eqs 11, 17, and 22 (rainbow curves), as well as the experimentally determined \( A_{yyz}^{(2)}/A_{zzz}^{(2)} \) (gray planes). The crossing lines of the rainbow curves and gray planes in parts c and d of Figure 4 represent the conditions that \( \theta_{\text{CH},\text{eff}} \) and \( \theta_{\text{CO},\text{eff}} \) should satisfy in the C–H and C==O stretch modes, respectively. By coupling these crossing curves, one can find a crossing point (Figure 4e).

The orientational distribution obtained from the above-mentioned procedure is displayed in Figure 4f, and shows good agreement with that obtained from the AIMD simulation data (Figure 4g). This good agreement demonstrates that the multimode coupling scheme can accurately predict the orientation of the formic acid molecules. The obtained distribution functions provide \( \theta_{\text{CH},\text{eff}} = 56 \pm 5^\circ \) and \( \theta_{\text{CO},\text{eff}} = 124 \pm 5^\circ \). The summary of the trans-conformation of the interfacial formic acid molecule is shown in Figure 4h. The multimode PD-HD-SFG technique using the multidimensional orientational distribution provides a universal approach for obtaining the interfacial molecular orientation. This method can also be applied to the biomolecules by probing the different moieties of the amino acid unit.

V. Å-SCALE DEPTH INFORMATION MEDIATED BY INTERFACIAL DIELECTRIC CONSTANT

Above, we learned that we can obtain the \( \chi_{yyz}^{(2)} \), \( \chi_{zzz}^{(2)} \), and \( \chi_{yyz}^{(2)} \) spectra from the measured \( \chi_{yyz,\text{exp}}^{(2)} \), \( \chi_{zzz,\text{exp}}^{(2)} \), and \( \chi_{yyz,\text{exp}}^{(2)} \) spectra via eqs 8–10. On the other hand, the peak amplitudes in the Im \( \chi_{yyz}^{(2)} \), Im \( \chi_{zzz}^{(2)} \), and Im \( \chi_{yyz}^{(2)} \) spectra of \( A_{yyz}^{(2)} \), \( A_{zzz}^{(2)} \), and \( A_{yyz}^{(2)} \), respectively, are not independent; \( A_{yyz}^{(2)} \), \( A_{zzz}^{(2)} \), and \( A_{yyz}^{(2)} \) are related via eqs 11 and 12. Now, let us focus on the \( A_{zzz}^{(2)} \) value. The value for \( A_{zzz}^{(2)} \) can be obtained using two different routes; one route is to acquire the \( A_{yyz}^{(2)} \) and \( A_{zzz}^{(2)} \) values from the Im \( \chi_{yyz}^{(2)} \) and Im \( \chi_{zzz}^{(2)} \) spectra and sequentially obtain the \( A_{zzz}^{(2)} \) value based on eqs 11 and 12. The other route is to obtain the \( A_{zzz}^{(2)} \) value directly from the Im \( \chi_{yyz}^{(2)} \) spectra. The values of \( A_{zzz}^{(2)} \) obtained from these routes are not necessarily identical, because they depend on the choice of the interfacial dielectric constant (\( \varepsilon' \)). Inversely, through the comparison of \( A_{zzz}^{(2)} \) one has access to the interfacial dielectric constant. As such, one can determine \( \varepsilon' \) through the matching of
two $A_{zz}^{(2)}$ values. Note that the same analysis can also be done with a focus on the $A_{yy}^{(2)}$ and $A_{yz}^{(2)}$ values. Next, we explain how to explore the depth information from our recent work.\textsuperscript{30}

The $\varepsilon'$ information can be connected with the averaged depth position of the vibrational chromophores.\textsuperscript{16,40,67} Let us consider the situation where the vibrational chromophores are located at $z = z'$. Here, a cavity containing the vibrational chromophores is embedded in the medium with the dielectric function of $\varepsilon'$ (see Figure 5a). The calculation of the local field correction\textsuperscript{16} leads to the expression of the interfacial dielectric constant $\varepsilon'$ at position $z'$:\textsuperscript{16}

\[
\varepsilon' = \left\{ \begin{array}{ll}
\frac{1}{1 + \frac{1}{3}(1 - \frac{\varphi}{\pi})(\varepsilon - 1)} & \text{for } z' \leq -r \\
\frac{2 + \frac{1}{3}(1 - \frac{\varphi}{\pi})(\varepsilon - 1) - r}{2 + \frac{1}{r}(1 - \frac{\varphi}{\pi})(\varepsilon - 1)} & \text{for } -r < z' \leq r,
\end{array} \right.
\]

\text{for } z' > r
\]

(23)

where $\varphi$ is the angle between the surface normal and the vector pointing from the center of the sphere to the crossing point of the dielectric interface and the sphere’s surface. $r$ is the radius of the vibrational chromophore. $z = 0$ denotes the location where the chromophore experiences $\varepsilon' = \varepsilon = (e + 5)/(4e + 2) = \pi/2$.\textsuperscript{16} Note that $\varphi = 0$ and $\pi/2$ provide the interfacial dielectric constants within the Lorentz and Slab models, respectively.\textsuperscript{16,66,69} Equation 23 links the interfacial dielectric function $\varepsilon'$ with the depth position $z$. The variation of the dielectric constant described in eq 23 is displayed in Figure 5b, together with the MD simulation data.\textsuperscript{40} Despite the simplicity of the embedded model, it captures the trend that $\varepsilon'$ varies with the depth position $z$ on a $\sim$ 5 Å-scale. As such, one can get the Å-scale depth information from the PD-HD-SFG data.

As an example, we consider the depth location of formic acid molecules at the interface of air with a water/formic acid mixture, and vary the formic acid concentration ($x_{FA}$). We chose formic acid as a benchmark molecule for demonstrating the validity of this scheme, because the C–H stretch mode can be easily assigned to the C–H group of formic acid, unlike the −CH$_3$ group where the amplitude of the C–H mode is modulated by the Fermi resonance of the overtone of the H–C–H bending mode and C–H stretch mode.\textsuperscript{70–72} The measured Im ($\chi_{zz}^{(2)}$)$_{eff}$, Im ($\chi_{pp}^{(2)}$)$_{eff}$ and Im ($\chi_{pp}^{(2)}$)$_{eff}$ spectra are presented in parts c–e of Figure 5f, respectively. The two approaches to reach the $A_{zz}^{(2)}$ value outlined above are shown in Figure 5f. The matching of the $A_{zz}^{(2)}$ value provides the average depth of the vibrational chromophores. The matching points of the $A_{zz}^{(2)}$ values are marked by “X” in Figure 5f. This figure indicates that the average depth of the C–H stretch chromophore moves from the air region to the bulk region by $\sim$0.9 Å when the concentration of the formic acid changes from 2.5% to 10% molar fraction at the air–water/formic acid mixture. The trend that the C–H stretch vibrational chromophores of formic acid moves to the bulk with increasing $x_{FA}$ is consistent with the AIMD simulation (Figure 5g). This result demonstrates that the PD-HD-SFG can capture the depth information with sub-Å-resolution.

Finally, we note that the probed region for the depth profile where the interfacial dielectric constant varies is $|z| < \sim 2$ Å at the aqueous solution interface, while the SFG active region is at least $|z| < 5$ Å.\textsuperscript{35–25} As such, the probed region for the depth profile is thinner than the SFG active region. When wider probed region is required, using a novel technique to probe the nanometer scale depth profiling through the SFG and difference frequency generation spectra is available.\textsuperscript{16}

VI. MD SIMULATION AS A TOOL FOR CRITICAL CHECK OF EXPERIMENTAL RESULT

Above, we outlined that several assumptions are required to interpret the SFG data. However, most of these assumptions cannot be accessed from the experimental side, meaning that computational support would greatly help.\textsuperscript{40} Computing the (multidimensional) orientational distribution from MD simulations is an essential guide for calculating the orientation of interfacial molecules, as seen above. Comparing the estimated depth from SFG measurement with the depth profile obtained from MD simulations is also very beneficial in guaranteeing the accuracy of the signal. Below, we explain the use of MD simulation for two cases.

MD simulations have been used for computing spectra, allowing us not only to interpret vibrational spectroscopy data\textsuperscript{52,77–80} but also to check the accuracy of experimental data\textsuperscript{53,81} and modeling.\textsuperscript{82,83} Moreover, simulations can provide powerful support when making assumptions for the analysis of experimental data.\textsuperscript{55,46} The typical flow for computing the SFG spectra is displayed in Scheme 1. The IR and/or Raman spectra are first calculated, ensuring the accurate modeling of the vibrational frequency and (transition) dipole moment/(transition) polarizability by comparing with experimental data. Subsequently, the researchers tackle the SFG spectra simulation by using the frequency, dipole, and polarizability modeling developed for IR and Raman calculation. A typical drawback of this approach is that it is difficult to identify the origin of the discrepancy when the simulated SFG spectra differ from the experimental data. In this approach, the discrepancy of the spectra arises not only from the force field model used for MD simulation but also from the modeling of the dipole moment and polarizability used for computing spectra. PD-HD-SFG provides another route to compare the experimental SFG spectra with the MD simulation data.\textsuperscript{4,26,84} The flow of comparing the PD-HD-SFG data with the simulation data is described in Scheme 2. In this scheme, we
can compare the experimental SFG data with the simulation data without performing the SFG spectra calculation, allowing us to skip computing the time evolution of the dipole moment and polarizability during the simulation. On the other hand, one should carefully pick the SFG-active species,\textsuperscript{85} for which one can calculate the orientational distribution and depth of the molecules from other criteria.

**VII. FUTURE OUTLOOK**

The PD-HD-SFG technique can be applied to explore the molecular-scale structure of liquid−liquid, liquid−solid, and air−solid interfaces. It opens the door to access the depth-related information in these interfaces. For example, it is interesting to understand how deeply the water molecules are in the oil subphase at the water−oil interface. Theoretically, it has been proposed that interfacial water forms a “finger-like” structure when ion transport occurs,\textsuperscript{90,91} but it has not been investigated experimentally, for lack of appropriate techniques. Furthermore, the technique can be used for identifying the SFG response of the hydroxyl group and their role in the wetting transparency.\textsuperscript{80,92}

The depth profiling and orientational analysis through the PD-HD-SFG technique could be used for identifying the 3D structure of interfacial peptides or proteins, but this may require isotopic labeling of specific parts of the peptide or protein; all the amide modes in the peptide or protein backbones contribute to the SFG signal, making the individual position and orientation of the amide groups ambiguous. To resolve individual amide groups, isotopic labeling of the target peptide and protein would be needed.\textsuperscript{27,93,94} Combining PD-HD-SFG with the isotopic labeling is on the horizon.

**VIII. CONCLUSION**

In this Perspective, we explained how the PD-SFG technique can be used for understanding not only the molecular orientation but also the Å-scale depth profiling of molecules. Moreover, the technique can provide information on the interfacial dielectric constant profile. For these analyses, HD-SFG spectra with accurate phase determination are essential. Although the HD-SFG technique was first developed over 10 years ago, this technique has been rarely measured at the polarization combination other than $sp$ and has seldom been used for the analysis of interfacial molecular orientation. The HD-SFG measurement on the $sp$, $ps$, and $pp$ polarization combination and chiral polarization\textsuperscript{95,96} is on the horizon. Furthermore, such a PD-SFG technique has not been combined with the time-resolved SFG technique,\textsuperscript{84,97−99} except for some studies.\textsuperscript{99,100−102} Founding a theoretical basis for time-resolved PD-HD-SFG and its demonstration will be an interesting next challenge for the SFG community.

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**Funding**

Open access funded by Max Planck Society.

**Notes**

The authors declare no competing financial interest.

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ACKNOWLEDGMENTS

We would like to thank Kris Strunge, Sho Imoto, Johannes Hunger, Tatsuhiro Ohto, and Masanari Okuno for fruitful discussions. This project was financially supported by the MaxWater initiative of the Max Planck Society.

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