Measuring Electric Fields and Noncovalent Interactions Using the Vibrational Stark Effect

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CONSPECTUS: Over the past decade, we have developed a spectroscopic approach to measure electric fields inside matter with high spatial (<1 Å) and field (<1 MV/cm) resolution. The approach hinges on exploiting a physical phenomenon known as the vibrational Stark effect (VSE), which ultimately provides a direct mapping between observed vibrational frequencies and electric fields. Therefore, the frequency of a vibrational probe encodes information about the local electric field in the vicinity around the probe. The VSE method has enabled us to understand in great detail the underlying physical nature of several important biomolecular phenomena, such as drug−receptor selectivity in tyrosine kinases, catalysis by the enzyme ketosteroid isomerase, and unidirectional electron transfer in the photosynthetic reaction center. Beyond these specific examples, the VSE has provided a conceptual foundation for how to model intermolecular (noncovalent) interactions in a quantitative, consistent, and general manner.

The starting point for research in this area is to choose (or design) a vibrational probe to interrogate the particular system of interest. Vibrational probes are sometimes intrinsic to the system in question, but we have also devised ways to build them into the system (extrinsic probes), often with minimal perturbation. With modern instruments, vibrational frequencies can increasingly be recorded with very high spatial, temporal, and frequency resolution, affording electric field maps correspondingly resolved in space, time, and field magnitude.

In this Account, we set out to explain the VSE in broad strokes to make its relevance accessible to chemists of all specialties. Our intention is not to provide an encyclopedic review of published work but rather to motivate the underlying framework of the methodology and to describe how we make and interpret the measurements. Using certain vibrational probes, benchmarked against computer models, it is possible to use the VSE to measure absolute electric fields in arbitrary environments. The VSE approach provides an organizing framework for thinking generally about intermolecular interactions in a quantitative way and may serve as a useful conceptual tool for molecular design.

1. MOTIVATION AND BACKGROUND

Much of contemporary chemistry is concerned with noncovalent interactions, as predicted by J.-M. Lehn nearly a quarter-century ago.1 Noncovalent interactions form the basis of molecular recognition, enabling matter to self-organize and emerge into complex structures. For these reasons, it is the recurring leitmotif of molecular biology2 (nucleic acid base-pairing, receptor−ligand specificity, protein folding, enzyme catalysis, membrane biophysics) and is increasingly exploited in new frontiers of synthesis based on self-assembly (DNA origami, crystal engineering,3 reticular/framework materials,4 and hybrid materials).

Despite the importance and ubiquity of noncovalent (intermolecular) interactions, it has been challenging for chemists to identify the right language to describe them. The dissociation constant ($K_D$) is certainly a common descriptor, but what it possesses in generality, it lacks in molecular insight. Dissociation constants do not identify what portions of the molecules are responsible for an interaction. Moreover, they depend on macroscopic parameters (temperature and pressure) and are difficult to relate to (or even calculate from5) the...
microscopic arrangement of molecules in a given system. These limitations pose challenges for the chemist interested in predicting properties of noncovalent interactions, issues at the heart of numerous synthetic challenges such as drug design. Conventionally, chemists address this problem by extending the concept of the covalent bond to describe noncovalent interactions, for example, hydrogen bonds (H-bonds), halogen bonds (X-bonds), π-stacking, cation–π bonds, charge-transfer interactions. However, these terms for specific interactions are often based on arbitrary geometric criteria (indeed, there is still debate as to what “counts” as a H-bond7), and their ability to explain or predict energetics is limited to ranges or ballpark values.

In our view, the usage of bonding concepts to describe intermolecular interactions is problematic because it belies the fact that most of these interactions are electrostatic (can be explained well without orbitals or electron densities) and because it ignores the nonspecific interactions (e.g., dipole–dipole, dipole-induced dipole) that can be just as energetically significant as the specific interactions attached special labels. What is needed is a model for intermolecular interactions that does not depend on assigning labels or cutoffs, applies equally to specific and nonspecific interactions, and is quantitative and microscopic. We believe the electric field, a fundamental concept from physics, is a conceptual tool that meets all these criteria.

Figure 1A illustrates the simple principles governing the interaction between a dipole and an external field: if the dipole is not aligned with the field, it is in a configuration of high energy; the field exerts a torque and rotates the dipole to align with it to give the lowest energy configuration. The energy depends on the dipole magnitude, the field magnitude, and their relative orientation as $U = -\mu \cdot \vec{F}$ where $\vec{F}$ denotes electric field and $\mu$ denotes dipole moment. Now consider a solute molecule (represented as a green circle) solvated by an aqueous environment (Figure 1B). In a chemical picture that consists of atoms and orbitals, a quantum mechanical calculation is needed to determine the interaction energy between solute and environment. However, many simple molecules can be represented as a point dipole (and complex molecules as a collection of dipoles) and the other molecules in the environment can be viewed as creating an electric field (represented with red field lines, Figure 1C) through their own charges, dipoles, induced dipoles, etc. In this picture, the interaction between a molecule and its surrounding environment can be recast as an interaction between a dipole and an electric field. This picture is quantitative and holds as long as there is no (or little) covalent character to the interaction.

By electric field, we mean specifically the electric field due to the environment, that is, not that due to the atoms that are part of the same molecule as the one that is said to experience the field. Interactions between atoms in the same molecule must be treated quantum mechanically. In this definition, atoms on an isolated molecule (as in the gas phase) experience zero electric field. Although arbitrary on some level, this definition amounts to choosing a useful reference state since our primary goal is to use electrostatic concepts to describe intermolecular interactions.

The picture in Figure 1C gives us a quantity (the electric field) that we can attach to a given environment (a protein, a solvent, a host) and tells us the energetic value associated with inserting a molecule (a ligand, a solute, a guest) into that environment at a particular position and orientation. More than just being an abstract concept though, the electric field can be measured using the vibrational Stark effect (VSE, section 2), enabling comparisons between very different systems (section 3), and can be easily computed with molecular mechanics models (section 2.3).

2. THE VIBRATIONAL STARK EFFECT

2.1. Vibrational Stark Spectroscopy (VSS)

Consider the molecular potential energy curve for a diatom such as CO (Figure 2A). These potentials are anharmonic, and one consequence of this is that when a molecule goes to higher vibrational energy levels, its bond lengths slightly ($\Delta d$). Note that the magnitude of $\Delta d$, $|\Delta d|$ or simply $d$, denotes the CO bond length, and its direction, $\hat{d}$, denotes the orientation of the CO bond axis. If this diatom possess a charge separation (q), that is, it possesses a permanent dipole moment, then its dipole moment will be slightly larger in the first vibrational excited state compared with the ground state: $\mu_0 = qd$, $\mu_1 = q(d + \Delta d)$. Infrared (IR) and Raman spectroscopies probe the energy gap between the ground and first excited vibrational state. Since these states have (slightly) different dipoles, they will be (de)stabilized (slightly) differently by an external field, $\vec{F}_{\text{ext}}$ (Figure 2B), depending on the orientation between the dipole and the field. Therefore, the applied electric field will produce a shift in the vibrational transition energy that is linear with the difference in the dipole moments of the two vibrational states, $\Delta \mu$: $\Delta \tilde{\nu} = -\Delta \mu \cdot \vec{F}_{\text{ext}}$. A vibration’s difference dipole is directly related to the vibrational frequency’s sensitivity to an external electric field, and so it is referred to as the Stark tuning rate. Vibrational Stark tuning rates are measured to be around 0.5–2
The unit corresponds to the frequency shift (in cm\(^{-1}\)) that would accompany the application of a unit electric field (1 MV/cm) projected along the vibrational axis. These tuning rates imply vibrational difference dipoles of 0.03–0.12 D (i.e., 1 D = 16.8 cm\(^{-1}\)/(MV/cm)). A polyatomic molecule has a more complex electrostatic and vibrational structure. However, if we limit our attention to high-frequency vibrations of specific functional groups (such as C=O or C≡N stretches), we find that they are largely decoupled from the rest of the molecule and behave similarly to the diatom (called one-dimensional behavior).8 This implies that \(\Delta \mu\) for such a vibrational transition is colinear with the vibrational mode’s bond axis.

Because the difference dipole arises from mechanical anharmonicity, it can be calculated from experimental quantities by finding \(q\) (for a diatom, simply \(\frac{\mu_0}{d}\)) and multiplying by \(\mu d\) (related to the difference in the gas-phase 1 ← 0 and 2 ← 1 transition energies). Values obtained from this kind of simple calculation are shown in Table S1 in the Supporting Information. Difference dipoles can also be calculated with \textit{ab initio} methods. Experimentally, the difference dipole can be measured in two different ways. The most straightforward method consists of accurately measuring the molecule’s dipole moment in the ground and vibrationally excited states and taking the difference; the accuracy required limits the scope of this approach to a few small gaseous molecules. A more general approach is vibrational Stark spectroscopy (VSS), in which external fields are applied to molecules and the effect on the vibrational spectrum is recorded (see Table S1, Supporting Information for examples and comparisons).9

In VSS (Figure 3A), a molecule containing the vibration of interest is first immobilized by embedding it in a polymer film or dissolving it in a glass-forming solvent that is rapidly cooled by immersion into liquid nitrogen. The glassy matrix fills a transparent capacitor, in which parallel plates are displaced ca. 20 \(\mu\)m. Voltages on the order of 1–2 kV are achievable before dielectric breakdown, resulting in applied electric fields as large as \(\sim 1\) MV/cm.9 The measurement consists of acquiring the IR spectrum of the sample in the presence and absence of the applied electric field. By analyzing the small differences between the spectra, the vibration’s difference dipole can be determined.9,10 For most vibrations that we have studied, the
frequency responds to the external field in a linear fashion. In the difference spectrum, this effect manifests as a second derivative line shape (the interested reader should consult refs 9 and 10 for details on this analysis).

Stark tuning rates have now been determined for many vibrations: O−H,11 N−H and S−H,12 C==O,13,14 C≡N,15−17 azide,17 and C−D.17 In VSS, all of these vibrations respond to an external field in a mostly linear fashion. There is however one major limitation to VSS that complicates the experiment’s interpretation, and that is the local field effect.9,18 What we measure is not \( \Delta \mu_f \) but \( \Delta \mu_{ff} \), where \( f \) (the local field correction factor) is a scalar whose value we estimate to be \( \sim 2 \). How we arrived at this estimate and its importance requires a lengthier and more technical discussion and is provided as a Supporting Information to this Account.

2.2. Vibrational Stark Effects to Probe Electric Fields

The VSE method treats the difference dipole as a sensitivity parameter that enables subsequently measured IR frequency shifts to be interpreted in terms of changes in electric field (Figure 3B). For example, if a vibration on a ligand (e.g., the C≡O stretch of heme−CO) has a Stark tuning rate \( |\Delta \mu_f|/f = 6 \text{ cm}^{-1}/(\text{MV/cm}) \) and its vibrational frequency shifts by \( 28 \text{ cm}^{-1} \) to the red when the vibrational probe is surrounded by a protein environment relative to a nonpolar solvent, then we would assert it experiences an electric field (projected along the CO bond axis) more stabilizing by 28 MV/cm in the binding site \( \Delta \mu_{ff} \). This information could provide a physical basis for (some of) the driving force for the ligand to bind to the protein. Similarly, frequency shifts accompanying a mutation (which can also be similarly large) can be interpreted in terms of changes in protein electric field due to the mutation (Figure 3B).20 To interpret these frequency shifts in proteins, structural data is needed to determine the orientation of the vibrational probe \( \Delta \mu_{ff} \) with respect to the protein field.

These assertions make two assumptions: first, that the frequency’s linear dependence on field, which is only shown from VSS for field perturbations of order 1 MV/cm, is also the case for the larger perturbation caused by the binding pocket; second, that the vibrational frequency shift due to changes in the environment can be fully attributed to an electric field effect. In the next section, we show how computational methods helped us test these critical assumptions.

Chemists are accustomed to interpreting vibrational frequency shifts as changes in bonds’ force constants (i.e., \( \psi = (2\pi)^{-1}(k/m)^{1/2} \)),
arising from changes in either bond order or bond strength. In applying the VSE to interpret frequency shifts associated with intermolecular interactions, we assume that the force constants (and intramolecular properties in general) are constant as the environment changes; for instance, the VSE would not explain the frequency shift between free CO and heme—CO. In practice, this assumption can be experimentally tested because a frequency shift due to a change in force constant would also result in a change in the difference dipole (measured by VSS) and anharmonicity (measured by overtone spectroscopy or 2-D IR). We find for many vibrations that frequencies can shift substantially among different environments without a noticeable change in the difference dipole, ruling out the changing force constant model; some vibrations involving hydrogen (S–H and N–H) are a notable exception.12

2.3. Solvatochromic Calibration

In VSS, when we take the IR spectrum of a solute in the glassy matrix, the peak frequency is very different from that molecule’s peak frequency in the gas phase, even before the external electric field is turned on. To a large extent, this shift is also a Stark effect, except it arises from the field created by the matrix16,21 (i.e., the surrounding molecules’ dipoles as illustrated by Figure 1C) instead of by an external voltage. Building on this concept, we can imagine a much easier experiment than VSS, wherein we vary the field on a vibrational probe not with a voltage source but by dissolving it in solvents of various polarities.22,23 Liquid solvents will exert an electric field (termed the solvent reaction field) in proportion to how polar the solvent is (a more polar solvent molecule will possess larger dipoles)24,25 and how polar the solute is (a more polar solvent will force the solvent to organize less randomly around it).25,26 However, solvent electric fields cannot be precisely controlled by an experimentalist, and so ascribing them specific values requires a model.

Dielectric constants and empirical polarity scales are useful ways to characterize solvents, but they are difficult to relate to the system’s microscopic structure. We have demonstrated that solvent-induced vibrational frequency shifts are effectively Stark effects. In one test of this, we found that a vibrational probe’s sensitivity to solvent variation (i.e., the frequency shift across a given range of solvents) is directly proportional to the probe’s difference dipole (Figure 4A,B). For instance, acetonaphone’s C==O vibration has approximately twice the Stark tuning rate of benzonitrile’s C==N (Figure 4A shows two schematic Stark spectra). We would then predict that acetonaphone’s band would shift twice as much as benzonitrile’s in response to solvatochromatic effects, which indeed turns out to be the case (shown schematically in Figure 4B), so Δνsub is proportional to |ΔFsub|/|ΔFsub| This relationship was also obtained for (1) two isopotologues of phenol, Ph—OH and Ph—OD,14 (2) a set of aminobenzonitrile positional isomers,27 and (3) a set of carbonyls with various degrees of conjugation (unpublished results). The proportionality is more quantitative when considering similar solutes. This test is important because it is independent of the values of the solvent fields.

A second test that is more stringent but also more model-dependent involves correlating the solvent-by-solvent frequency variation for an individual solute against the average solvent field calculated for each solvent, as shown in Figure 4C.28,29 The strategy we have generally adopted for these calculations is molecular dynamics (MD) simulations, though Poisson–Boltzmann and the Onsager models can give similar results with some caveats.28 For some solutes, these curves are linear over the full range of electric fields spanned by solvent variation and include solvents that can participate in specific interactions (H-bonds and X-bonds) with the vibrational probe. This behavior has been observed for several molecules’ C==O vibrations (acetophenone’s is shown in Figure 4C) and O==H vibrations. Solvent fields (order tens of MV/cm) can be large relative to the largest achievable external fields in a capacitor (order 1 MV/cm) because of the short distances on which molecular interactions act.

The solvent field-frequency relationships affirm the two earlier assumptions about the VSE: that its linearity can extend over a wider domain than that directly observed in VSS and that it is sufficient to explain almost all variation in frequency across solvents with very different chemical compositions and H-bonding capacities.14 The field-frequency curve additionally allows us to map the observed frequency in an arbitrary environment to the average absolute electric field.

The extra word “average” is important because solvent molecules’ positions fluctuate, so the instantaneous solvent electric field for a given configuration can be very different from the mean of the field distribution sampled by the solvation dynamics. The peak position of a vibrational probe only conveys information about the ensemble-average electric field; on the other hand, the line width of a vibrational probe’s band is related to the dispersion in electric fields associated with an environment (i.e., its inhomogeneity).

A significant datum in Figure 4C is the point for water, which produces a considerably larger bathochromic shift than other solvents, and is also calculated to exert proportionally larger electric fields. This finding is suggestive that the H-bonding interaction is amenable to an electrostatic description, and that we can think of its greater strength relative to a dipole–dipole interaction in terms of its capacity to generate a greater electric field. This feature is central to our picture that electric fields serve as a unifying concept to compare specific and nonspecific interactions on an equal footing.

We arrived at this conclusion using the carbonyl vibration’s simple frequency variation (that is, a “pure” linear Stark effect). This property is not universal. For instance, a great deal of work on the VSE has focused on the nitrile probe because of its appearance in an uncluttered window in the mid-IR (ca. 2200 cm⁻¹), the many methods now available for introducing nitriles into proteins,30 and its common occurrence in drugs.31 However, once we started to use solvatochromism to benchmark vibrational probes, a problem came into focus, as exemplified by benzonitrile (Figure 4D). When dissolved in water, its CN stretching frequency falls to the blue of what it is in hexane, a staggering departure from the expected behavior (the dotted line).32 This effect, referred to as the “H-bonding contribution” to the nitrile frequency shift, has led to confusing results.33,34 We dedicated considerable effort to dissecting the H-bonding component from the electrostatic component of the frequency shift;55 however, it is no longer clear to us whether the nitrile’s complicated frequency shifts are necessarily limited to H-bonding per se. Rather, these shifts likely report indirectly on a more complex electrostatic sensitivity beyond the first-order electric field term.23 Similarly, the Fe–C==O frequency in CO–myoglobin was found not to conform to a purely linear Stark effect;56 in this case, the frequency was also modulated by changes in the electronic structure of the weak Fe–C bond. In hindsight, nitriles and CO–myoglobin constitute a cautionary tale for future work: in order to establish quantitative
mappings between frequency measurements and electric fields, it is imperative that the vibrational probe be benchmarked against reference data before interpreting its frequencies in more complicated systems. As of this writing, carbonyl probes have proven to be the most compliant by this standard.

3. ELECTRIC FIELDS AND A BROADER VIEW OF NONCOVALENT INTERACTIONS

3.1. Chemical Systems

Noncovalent interactions and molecular environments come in many different forms. In all cases, the electric field provides a metric of the environment’s interaction strength and enables us to make broad comparisons across highly diverse environments (Figure 5).

On the weakest end, we encounter solvation forces associated with nonpolar solvents. These are limited to induced dipole effects and give rise to electric fields in the regime of 0 to −10 MV/cm. Note that the negative sign implies a favorable electric field that lowers the molecule’s energy. It is important to point out that induced dipole effects are not captured by most force fields, because they require explicit incorporation of polarizability. As such, the data in Figure 4C,D (which used a nonpolarizable AMBER-like force field) associate hexane with a solvent field near zero.

As we move to polar solvents, the average electric field increases to the range of −20 to −40 MV/cm, reflecting the additional contributions of permanent dipoles, which can create larger electric fields than induced dipoles. In general, a more polar environment will consist of molecules with greater permanent dipoles, and so exert correspondingly larger electric fields.

In addition to these simple interactions, we can also consider the electrostatics associated with numerous specific interactions. For our purposes, a specific interaction has two main criteria: (1) it is local; that is, it involves one specific molecule in the environment as opposed to the environment as a whole, and (2) it enforces a specific geometry between the interacting partners. Weak H-bonds constitute a class of specific interactions whose definition and scope have expanded tremendously in the past two decades. Two examples of this category are C−H hydrogen bonds (where C−H acts as a H-bond donor) and π−H bonds (where a π-ring acts as a H-bond acceptor). Previous work of ours interrogated the π−H bond formed between phenol’s OH group and benzene derivatives. It was found that benzene exerts an electric field around −20 MV/cm onto phenol’s O−H bond (presumably, through its quadrupole moment), an effect that explains most of the binding energy of the interaction; furthermore, the interaction could be systematically varied by changing benzene’s substituents, and the changes in electric fields and binding constants were well explained by the VSE. Standard H-bond donors (such as O−H and N−H bonds) possess larger dipole moments than C−H, and as such can exert larger electric fields and interact more favorably. The key feature about H-bonds that allow them to exert larger electric fields than the similar dipole−dipole interaction is that hydrogen is the smallest atom, so the dipole associated with a (heavy atom)−H bond can get much closer to other molecules (e.g., acetone in Figure 5). This proximity is very significant for the electric field because the field due to a dipole attenuates as the distance cubed.

A significant instance of the standard H-bond is the one donated by water molecules. Because the water molecule is small, liquid water possesses a very high density of H-bond donors, facilitating the formation of multiple simultaneous H-
bonds. For these reasons, water is very commonly at the extreme of many solvatochromic series. We would classify electric fields as being "large" if they exceed the average field exerted by water. At ambient pressures and temperatures, liquid water achieves possibly the highest density of polar atoms and H-bond donors, so large fields require an environment that can (1) squeeze atoms closer together than they ordinarily would be or (2) force molecules to stay confined to a particular configuration that optimizes the electric fields. In chemical systems, these criteria are most usually met through certain intramolecular architectures. Proteins also possess the ability to tightly control H-bonding geometry and compress interatomic distances to achieve large electric fields, as discussed in the next section. It is important to point out that in these strong H-bonds, the distinction between a molecule and its environment can break down because interatomic separations are close to those found between atoms within molecules. In these cases, the interaction can exhibit decidedly quantum features, such as nuclear delocalization and electronic covalency, which are not captured in the electric field picture (Figure 1C).

The main point though that emerges from Figure 5 is that whereas strong specific interactions can produce larger electric fields than are possible with nonspecific interactions, there is considerable overlap between the weaker end of the specific interactions and the stronger end of the nonspecific interactions. Therefore, most intermolecular interactions can be unified and described quantitatively using the electric field language. The electric field becomes less rigorous when quantum effects are important; however even in such cases, we (and others) have found the electric field to be a useful and even semiquantitative proxy for estimating the energy of these partially covalent interactions.

3.2. Biochemical Systems

Native proteins are highly organized forms of matter and so the environments within their active sites or binding pockets are controlled with atomic precision. Given this, what electric fields would we expect to find inside them? This question was what motivated us to develop the vibrational Stark effect method (as well as techniques for incorporating vibrational probes) starting about 15 years ago, though there had been other less direct strategies developed a decade earlier to infer fields indirectly based on $pK_a$ or redox potential shifts. Recent work based on deploying vibrational probes into the hydrophobic core of ribonuclease S via unnatural amino acids found that electric fields in the protein interior were on the low end; for instance, the C=O bond of a para-acetyl-phenylalanine probe records an average field of $-19$ MV/cm (Figure 6A). Though less simple to connect with absolute electric fields, the frequencies of nitrile probes incorporated into ribonuclease $S$ and the interior of other proteins are generally similar to the frequencies that those probes exhibit when dissolved in low-to-medium polarity solvents.

One feature that immediately stands out as distinct is that IR bands of probes in protein interiors tend to be very narrow. To a first approximation, the line width of a vibrational Stark probe is related to the span of electric fields that the probe samples (i.e., inhomogeneous broadening). Because proteins are more
structured and less free to reorganize than bulk solvents, our probes generally experience smaller ranges of electric fields when inserted inside proteins.33,44

In our studies on ketosteroid isomerase (Figure 6B), we used a substrate-like inhibitor to probe the electric field that the enzyme exerts onto the chemical bond (a carbonyl group) that undergoes a charge rearrangement in the reaction mechanism. The combination of strong H-bonds furnished by key active site residues and the oriented dipoles across the whole enzyme scaffold give rise to an unprecedentedly large field of around \(-140 \text{MV/cm}, \) much larger than the average field the inhibitor experiences in aqueous solution (as seen by the spectrum’s red-shift in Figure 6B).55 This extreme electric field seems to be the physical realization of an intricate catalytic apparatus because studies on a series of mutants demonstrated a direct relationship between electric field strength and enzyme catalytic power.56 The narrow line width (much narrower than water) possibly reflects the active site’s ability to keep the steroid molecule locked into a very specific orientation in the active site, thus maintaining an extreme and functionally important electric field at equilibrium.

4. OUTLOOK

The VSE provides a direct mapping between vibrational frequency and local electric field. Since the former is an experimental observable and the latter can be calculated (or related to chemical intuition), the VSE provides an approach to translate experimental data on chemical systems into a simple and general language for describing and quantitating intermolecular interactions. Vibrational probes are intrinsic to many molecules; since they can consist of just two atoms, they can also be extrinsically added to systems with minimal perturbation. Vibrational frequencies can be measured with many modalities and can also be finely resolved both spatially and temporally. Therefore, from an experimental perspective, the VSE approach should be applicable to a host of physical and biological problems. Moreover, electric fields can be calculated with many different models and levels of theory, from back-of-the-envelope point-charge calculations to \textit{ab initio} quantum calculations. This further increases the VSE’s generality since many systems are more amenable to one kind of model over others.

It is worth briefly mentioning that in previous literature, other frameworks were applied to interpret vibrational frequency shifts, in terms of changes in bond strength or electron polarization; however, the results summarized here (and elsewhere11–17,19,22,23,27,56,43) have demonstrated that electrostatics provide a consistent way to interpret vibrational frequency shifts. This paradigm shift prompts the need to reevaluate the interpretation of vibrational spectra in earlier literature.

Finally, because the electric field picture provides a natural meeting point for computation, experiment, and chemical intuition, we think it may prove to be a useful conceptual tool for framing problems in molecular design. This matter remains to be seen for the next generation of VSE research.

ASSOCIATED CONTENT

Supporting Information

More technical discussion of the local field effect. This material is available free of charge via the Internet at http://pubs.acs.org.


Addendum to
Measuring Electric Fields and Non-covalent Interactions Using the Vibrational
Stark Effect

on
The Local Field Correction Factor

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The local field effect is a fundamental phenomenon that accompanies the
application of an external electric field to a dielectric material, which in Vibrational Stark
Spectroscopy (VSS) is the frozen glass that contains the sample molecule. When a
capacitor is empty, the electric field between the plates is homogeneous and everywhere
well described by the simple relation $V = \frac{|F_{\text{ext}}|d}{\varepsilon}$, where $V$ is the voltage applied between
plates separated by a distance, $d$ (both accurately known). When the capacitor is filled
with a dielectric medium, the equation still holds in the macroscopic sense (i.e., for the
spatially-averaged electric field) but the microscopic position-dependent field will be
inhomogeneous due to the fact that the external field creates an inhomogeneous
polarization in the medium, which in turns exerts its own electric field. As a result, the
actual electric field at the position where the vibrational chromophore resides is not
simply $|F_{\text{ext}}| = \frac{V}{d}$; it is said to experience the local field, which for small external fields
varies linearly with the external field ($\vec{F}_{\text{local}} = f\vec{F}_{\text{ext}}$), where $f$ (the local field correction
factor, in general a tensor, but well approximated by a scalar) is greater than unity. The
consequence of this is that in VSS, what we measure is not $|\Delta\mu|$, but $|\Delta\mu|f$. The local
field effect cannot be easily dispensed, since the matrix (which creates the local field
effect) is essential for the VSS experiment. Note that the local field effect enters into
many forms of spectroscopy and its evaluation and complications have a long history.

Early work of ours found that $f$ is reasonably constant across typical matrices, but
was agnostic to its value (we expressed difference dipoles and all key quantities as
products with $f$). While we often proposed estimates like $1.1 < f < 1.3$, our most recent
work and reasoning (described in the following) suggests $f$ is closer to 2. This remains
one of the less well-resolved issues of Stark spectroscopy that also impacts many other
areas of spectroscopy, and our hope is that this discussion will help stimulate further
work on comparing simple systems in the gas and condensed phases.

We begin by defining terms, and discussing the similarities and differences
between four experimental methods (rows 2–5 in Table S1) and ab initio methods (row 1) available for determining difference dipoles. We next discuss the available results (Table S1) where these methods have been applied to a set of six model compounds. Finally, we provide a short review of previously published literature on the subject.

Table S1. Stark tuning rates of six compounds based on five methods

| method of finding $|\Delta \mu|$ | molecule | C=O | C≡N<sup>©</sup> | H--C≡N | H--F | acetone | acetonitrile |
|---------------------------------|----------|-----|----------------|--------|------|---------|-----------|
| 1. ab initio methods            |          |     |                 |        |      |         |           |
| 0.437<sup>a</sup>              |          | 0.18<sup>a</sup> | -0.02<sup>b</sup> | 0.817<sup>c</sup> | 0.44<sup>d</sup> | 0.27<sup>d</sup> |
| 2. based on anharmonicity       |          |     |                 |        |      |         |           |
| 0.429<sup>e</sup>              |          | 0.18<sup>e</sup> | —              | 0.735<sup>f</sup> | —    | 0.19<sup>g</sup> |
| 3. based on experimental $\mu_i - \mu_j$ | |     |                 |        |      |         |           |
| 0.421<sup>h</sup>              |          | —   | -0.078<sup>i</sup> | 0.792<sup>l</sup> | —    | —       |
| 4. measured by vibrational Stark spec. | |     |                 |        |      |         |           |
| 0.7<sup>k</sup>                |          | 0.32<sup>e</sup> | 0.71, 0.82<sup>l</sup> | —      | 0.76<sup>k</sup> | 0.43<sup>l</sup> |
| 5. slope fit to solvatochromism | |     |                 |        |      |         |           |
|                                 |          | —   | —              | —      | 0.34<sup>m</sup> | 0.22<sup>n</sup> |

<sup>g</sup> frequencies from Reimers, J. R.; Hall, L. E. J. Am. Chem. Soc. 1999, 121, 3730–3744 and electric fields borrowed from work on benzonitrile.
<sup>m</sup> frequencies from Kolling, O. W. Trans. Kansas Acad. Sci. 1996, 99, 161–166 and electric fields borrowed from work on acetophenone.

**Definition.** The field-dependence of a vibrational transition is assumed to be captured by the equation $	ilde{v} = -\frac{1}{2} \hat{F} \cdot \Delta \alpha \cdot \hat{F} - \Delta \tilde{\mu} \cdot \hat{F} + \tilde{v}_0$, where $\Delta \alpha$ is the difference polarizability tensor, and $\Delta \tilde{\mu}$ is the difference dipole vector. Assuming 1-dimensional behavior to the oscillator, the only component of the field that is relevant is that which is parallel to the internuclear axis of the vibration in question, in which case the equation simplifies to $	ilde{v} = -\frac{1}{2} |\Delta \alpha| F^2 - |\Delta \tilde{\mu}| F + \tilde{v}_0$, where $|\Delta \alpha|$ is the quadratic Stark tuning rate.
and $|\Delta \mu|$ is the linear Stark tuning rate. As mentioned in the text, this equation assumes that the electric field dominates the frequency variation, and excludes scenarios in which electric field gradients or other higher-order electrostatic effects are significant.\(^3\)

Moreover, we note the derivative of this equation, which corresponds to the instantaneous change in frequency with respect to field, is $\Delta \bar{\mu}_{\text{full}} = -|\Delta \alpha| |F| - |\Delta \bar{\mu}|$. This equation indicates that the linear Stark tuning rate has two parts: a field-independent component, $|\Delta \bar{\mu}|$, which we will refer to as the \textit{intrinsic} difference dipole; and a field-dependent component, $|\Delta \alpha| |F|$, which we will refer to as the induced difference dipole. In other words, the previous equation can be rewritten as: $\Delta \bar{\mu}_{\text{full}} = -|\Delta \bar{\mu}_{\text{ind}}| - |\Delta \bar{\mu}_{\text{int}}|$

\textbf{Four measures of difference dipole.} Referring to Table S1, there are four independent ways of experimentally assessing a vibration’s difference dipole. Each of these measures is based on different models and comes with its own set of assumptions. Additionally, we critically assess for each experiment \textit{what is actually measured}.

The most fundamental measure of a vibration’s difference dipole is to simply take the difference in molecular dipole moment between the ground and excited vibrational states in the gas phase (method 3). This experiment does not involve external or internal fields of any kind, and directly measures $|\Delta \bar{\mu}_{\text{int}}|$. The high accuracy needed in the dipole moments (on the order of parts per thousand) limits this method to a few small molecules. We would consider comparisons between this measure (method 3) and Stark spectroscopy in the condensed phase (method 4) an ideal test of the local field effect; however, we have been able to find only scant gas-phase data of this kind (see row 3).

In method 2, the difference dipole is inferred on the basis that it is due to mechanical anharmonicity (see Figure 2 from main text).\(^4,5\) To populate this model with experimental quantities, we need measures of: (1) the increase in the bond length expectation value going from the ground to the first excited vibrational state, and (2) the charge separation of the vibration. The former can be determined analytically from the “experimental anharmonicity” ($\bar{v}_{0-1} - \bar{v}_{1-2}$) given a model anharmonic potential function (such as the Morse or the cubic potentials); the latter, for a diatomic molecule, is given simply by the ground state molecular dipole and bond length. Similar concepts can be applied to polyatomic molecules.\(^6\) If the relevant data are obtained from gas-phase experiments, the difference dipole obtained will correspond to $|\Delta \bar{\mu}_{\text{int}}|$; however, this method \textit{assumes} that only mechanical anharmonicity is responsible for the difference
dipole.

In Stark spectroscopy (method 4 in the table), a small external electric field is exerted onto a molecule that is already experiencing a considerably larger internal (matrix) field due to the frozen glass surrounding it. In other words, the Stark tuning rate that is measured corresponds to the derivative of the field-frequency curve evaluated at a field equal to the matrix field. Moreover, the local field effect introduces an extra uncertainty as to what electric field the external charges exercise locally where the vibrational chromophore is located. Therefore, in Stark spectroscopy one measures \( \langle |\Delta \mu_2| \rangle_{\text{matrix}} + \langle |\Delta \mu_{\text{int}}| \rangle_{\text{int}} \). Note that all the Stark experiments in Table S1 were conducted in a frozen 2-methyl tetrahydrofuran matrix, except CN\(^-\) (isolated in KBr\(^4\)). The value of the difference dipole as determined by vibrational Stark spectroscopy does not invoke any assumptions per se, but due to the local field effect and to the matrix-induced contribution, it corresponds to a different expression \( \langle |\Delta \mu_2| \rangle_{\text{matrix}} + \langle |\Delta \mu_{\text{int}}| \rangle_{\text{int}} \) than that of methods 2 and 3 \( \langle |\Delta \mu_{\text{int}}| \rangle_{\text{int}} \).

In solvatochromism experiments (method 5), the Stark effect is measured indirectly and assumes an accurate model for the average reaction fields the vibration experiences in a series of solvents. In this experiment, there is no local field effect (because no external fields are applied); however, unlike Stark experiments, the field differences between two solvents are large. Therefore, the frequency shift does not report on the instantaneous difference dipole at a given field: whatever quadratic dependence exists over the interval gets averaged over. Therefore, the slopes of solvatochromism curves correspond to \( \langle |\Delta \mu_2| \rangle_{\text{matrix}} + \langle |\Delta \mu_{\text{int}}| \rangle_{\text{int}} \) where \( \langle F \rangle \) is the average solvent electric field over the interval. However, since no external fields are operative, the local field factor does not enter.

In summary, we expect tuning rates from solvatochromism (method 5) to be slightly larger (due to the matrix-induced contribution) than the more fundamental determinations (methods 2 and 3), and the tuning rate from Stark spectroscopy (method 4) to be larger still (due to the local field effect).

\textbf{Ab initio methods.} A number of methods have been proposed over the years to calculate vibrational Stark tuning rates through \textit{ab initio} quantum calculations – either by applying an electric field \textit{in silico} or by determining field-free parameters that are interrelated to the Stark properties. Values are relatively sensitive to method and basis set, leading to results that vary widely in quality. In Table S1, the first four entries
(the small molecules) have had their tuning rates calculated at very high levels (\(a\) CCSD(T)/aug-cc-pV5Z; \(b\) QCISD/aug-cc-pV5Z; \(c\) QCISD/6-311++G(3df,2pd)) and are likely accurate to within 10% of \(|\Delta \vec{\mu}_{\text{int}}|\). A larger body of \textit{ab initio} work has been carried out with density functional theory (DFT); however, it has been known\(^1\),\(^15\) that single-reference methods tend to overestimate difference dipoles, by as much as 50%. High-level multi-reference techniques are difficult to perform on molecules of more than a few atoms, and so DFT-calculated values for acetone and acetonitrile are reported in Table S1. We estimate that these overestimate \(|\Delta \vec{\mu}_{\text{int}}|\) by ca. 20–30%.

**The diatomic molecules: CO, CN\(^{-}\), HF.** It is immediately clear that for the molecules CO, CN\(^{-}\), and HF, the \textit{ab initio} difference dipoles are in good agreement with the two fundamental experimental determinations (methods 2 and 3), which also correspond to \(|\Delta \vec{\mu}_{\text{int}}|\). The strong agreement between methods 1 and 2 implies that, at least for these small molecules, mechanical anharmonicity is sufficient to explain the difference dipoles.

On the other hand, the difference dipoles measured for CO and CN\(^{-}\) by vibrational Stark spectroscopy are significantly larger (1.6-fold and 1.8-fold) than the consensus values from methods 1–3. The larger value from method 4, which corresponds actually to \((|\Delta \alpha| F_{\text{matrix}} + |\Delta \vec{\mu}_{\text{int}}|) f\), could in principle be due either to an induced difference dipole or to the local field effect. From \textit{ab initio} calculations of CO, \(|\Delta \alpha| = 0.032 \text{ Å}^3 \) or 0.0018 cm\(^{-1}\)/(MV/cm)\(^2\). If we multiply this by a conservative estimate of the matrix field in frozen 2-methyl tetrahydrofuran (30 MV/cm), we would obtain a value for \(|\Delta \vec{\mu}_{\text{int}}|\) of 0.05 cm\(^{-1}\)/(MV/cm). However, the actual difference between \(|\Delta \vec{\mu}|\) from method 4 and the other methods is 0.27 cm\(^{-1}\)/(MV/cm). So, the induced difference dipole only accounts for at most 20% of the discrepancy; it is concluded that the majority of the discrepancy arises instead from the local field effect. A similar arguments holds for CN\(^{-}\), for which a high level calculated value of \(|\Delta \alpha|\) is also available.

Moreover, it would seem then that the difference in the Stark tuning rates between methods 2 and 4, previously believed to reflect to electronic anharmonicity,\(^2\),\(^5\)–\(^7\) is more parsimoniously explained by the local field effect. At present, we have been unable to obtain a vibrational Stark spectrum of HF, although that data could serve as an additional test of these claims, especially since its intrinsic difference dipole is quite large.

**The case of HCN.** HCN exhibits very different vibrational Stark properties than
the other small molecules.\textsuperscript{17} Its $|\Delta \mu_{\text{int}}|$, measured directly in the gas phase (method 3), is very small and of opposite sign compared to the other molecules in Table S1 (its dipole moment is smaller in the vibrational excited state). High-level \textit{ab initio} methods reproduce the sign inversion and small size of HCN’s difference dipole, although quantitative agreement is not achieved. In contrast, the difference dipole measured in vibrational Stark spectroscopy in a 2-methyl tetrahydrofuran matrix is of normal sign and 10 times larger. These observations are best explained as an induced effect, that is $|\Delta \mu_{\text{int}}| \approx 0$, but $|\Delta \alpha|$ is appreciable and so $|\Delta \mu|$ is significantly greater than zero when HCN is in the presence of a matrix field. This behavior is essentially a vibrational equivalent to the electronic Stark effects of carotenoid molecules, which possess difference dipoles of much greater magnitude when subjected to the large matrix field present inside the photosynthetic LHII antenna protein.\textsuperscript{18} In principle, HCN could provide another useful means to approach the local field effect, because its difference dipole is described by the simple expression $|\Delta \mu| = |\Delta \alpha| F_{||}$. This analysis would require a reliable estimate for the difference polarizability however, which is not presently available (experimental difference polarizabilities, manifested as first-derivative contributions to the Stark spectrum are obscured and difficult to quantify in molecules with large difference dipoles; and the \textit{ab initio} value differs by two orders of magnitude from experiment).

\textbf{The larger molecules: acetone, acetonitrile.} The vibrational difference dipoles of larger molecules are amenable to investigation by vibrational Stark spectroscopy, solvatochromic variation, and \textit{ab initio} methods (albeit at a lower level of theory). Data using Method 3 in general are not available, and analyses based on mechanical anharmonicity are possible, though are more complicated since inter-mode coupling must be accounted for; a calculation of this kind has been carried out on acetonitrile. For acetonitrile, methods 2 and 5 are in good agreement. Since solvatochromism probes $(|\Delta \alpha| F_{\text{matrix}} + |\Delta \mu_{\text{int}}|)$, and the agreement implies a small value for $|\Delta \alpha|$. The \textit{ab initio} difference dipole is slightly larger (by $\sim 30\%$), although this is expected because it is a DFT calculation. Stark spectroscopy, which probes $(|\Delta \alpha| F_{\text{matrix}} + |\Delta \mu_{\text{int}}|) f$, gives a difference dipole that is twice as large as methods 2 and 5. Since $|\Delta \alpha|$ is small, the difference is most likely due to $f$, as concluded for the smaller molecules. Very similar findings are obtained for acetone as well.
Summary. In light of these lines of evidence, it is worth reflecting on why the assumption that the local field effect is small ($f \approx 1$) has been so persistent. Our reflections here are not comprehensive, but we will point out a few major pitfalls that have likely promulgated misunderstanding.

Earlier work of ours asserted that solvatochromism curves give tuning rates that match the values from Stark spectroscopy assuming $f = 1$. Regrettably, however, this was based on an incorrect use of the Onsager reaction field equation that was missing a factor of $4\pi/3$.\(^9,10\) (Unfortunately, this error has been propagated by other workers). The use of post hoc dielectric factors used to re-scale electric fields calculated by MD simulations to force better agreement with Stark tuning rates determined by vibrational Stark spectroscopy (assuming $f \approx 1$) have also served to obscure the various sources of disagreement, more likely due to the local field correction.\(^19,20\)

DFT, because it overestimates difference dipoles by 20–50%, can give values that qualitatively agree (accidentally) with vibrational Stark spectroscopy assuming $f \approx 1$. Also, the DFT study undertaken by Brewer and Franzen\(^16\) used an incorrect approach to calculate difference dipoles (by separating geometry distortion from anharmonicity) that gave false confidence $f$ is close to unity. A proper ab initio treatment of the vibrational Stark effect was developed by Reimers and Hush;\(^14\) however, their methods are very computationally demanding and so have not been broadly adopted.

On the other hand, a study undertaken in our lab by Saggu et al. showed that the vibrational frequency shifts in the O–H bond of phenol induced by complex formation with a π H-bond acceptor were consistent with the Stark tuning rate from Stark spectroscopy assuming $f \approx 1$.\(^21\) Similar observations were obtained for O–D, N–H, and S–H.\(^22\) The evidence so far then would seem to suggest that $f \approx 1$ for bonds that involve hydrogen, while $f \approx 2$ is more suitable for “heavy” vibrations (e.g., vibrations for which the participation of hydrogen in the mode is low). We do not have a general explanation for these phenomena at the present time, but believe that more gas phase measurements (akin to row 3 in Table S1) may shed further light on this subtle issue.

References


