

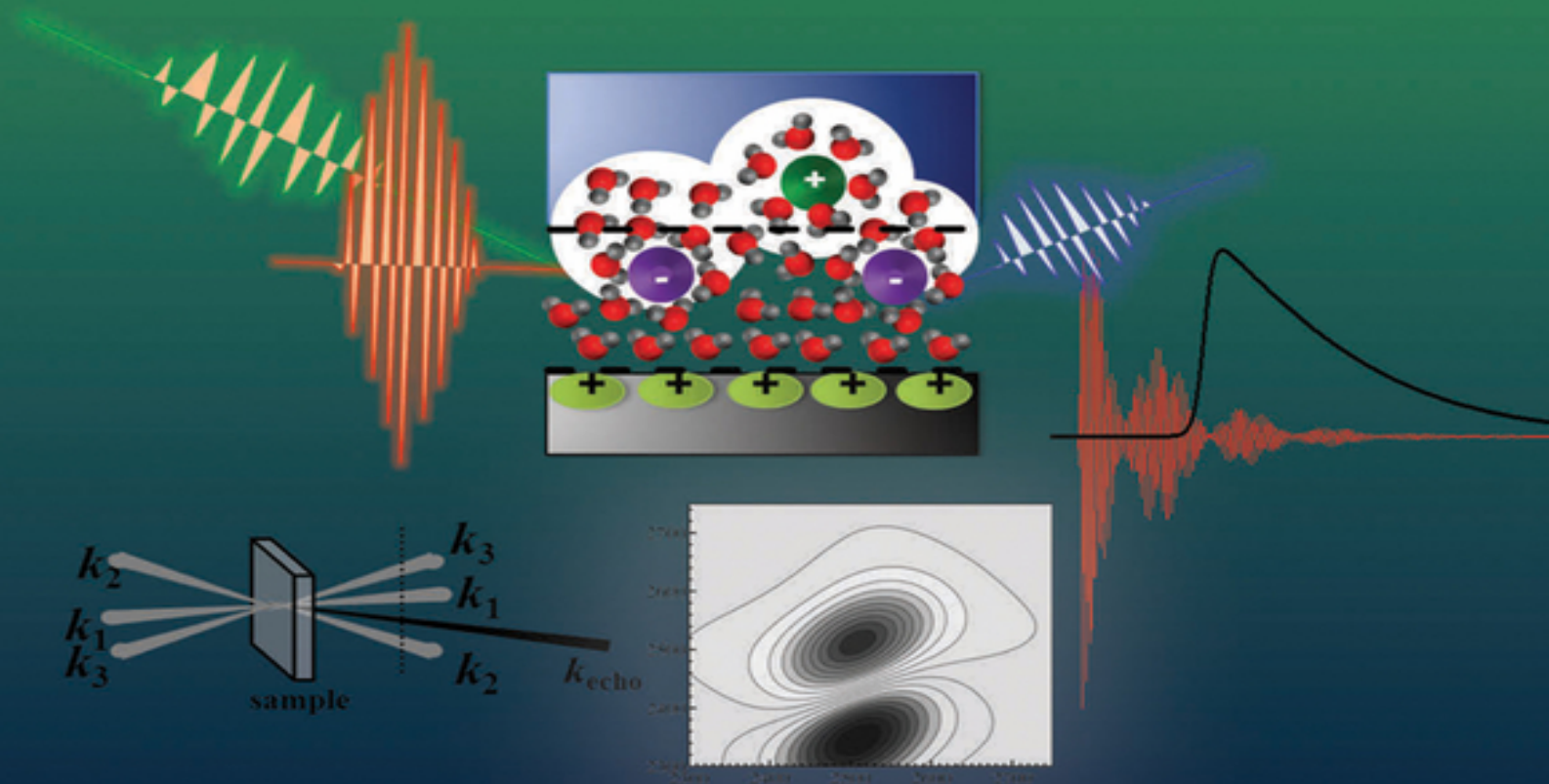
**Wiley Series on Electrocatalysis and
Electrochemistry**

Andrzej Wieckowski, Series Editor

VIBRATIONAL SPECTROSCOPY AT ELECTRIFIED INTERFACES

Edited by Andrzej Wieckowski, Carol Korzeniewski and Björn Braunschweig

With a Foreword by Masatoshi Osawa, Hokkaido University



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Wiley Series on Electrocatalysis and Electrochemistry

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Preface to the Wiley Series on Electrocatalysis and Electrochemistry

This series covers recent advances in electrocatalysis and electrochemistry and depicts prospects for their contribution into the present and future of the industrial world. It aims to illustrate the transition of electrochemical sciences from its beginnings as a solid chapter of physical chemistry (covering mainly electron transfer reactions, concepts of electrode potentials, and structure of electrical double layer) to the field in which electrochemical reactivity is shown as a unique chapter of heterogeneous catalysis, is supported by high-level theory, connects to other areas of science, and includes focus on electrode surface structure, reaction environment, and interfacial spectroscopy.

The scope of this series ranges from electrocatalysis (practice, theory, relevance to fuel cell science and technology) to electrochemical charge transfer reactions, biocatalysis, and photoelectrochemistry. While individual volumes may appear quite diverse, the series promises updated and overall synergistic reports providing insights to help further our understanding of the properties of electrified solid-liquid systems. Readers of the series will also find strong reference to theoretical approaches for predicting electrocatalytic reactivity by such high-level theories as density functional theory. Beyond the theoretical perspective, further vehicles for growth are such significant topics such as energy storage, syntheses of catalytic materials via rational design, nanometer-scale technologies, prospects in electrosynthesis, new instrumentation, and surface modifications. In this context, the reader will notice

that new methods being developed for one field may be readily adapted for application in another.

Electrochemistry and electrocatalysis have both benefited from numerous monographs and review articles due to their depth, complexity, and relevance to the practical world. The Wiley Series on Electrocatalysis and Electrochemistry is dedicated to present the current activity by focusing each volume on a specific topic that is timely and promising in terms of its potential toward useful science and technology. The chapters in these volumes will also demonstrate the connection of electrochemistry to other disciplines beyond chemistry and chemical engineering, such as physics, quantum mechanics, surface science, and biology. The integral goal is to offer a broad-based analysis of the total development of the fields. The progress of the series will provide a global definition of what electrocatalysis and electrochemistry are now, and will contain projections about how these fields will further evolve in time. The purpose is twofold, to provide a modern reference for graduate instruction and for active researchers in the two disciplines, as well as to document that electrocatalysis and electrochemistry are dynamic fields that are expanding rapidly, and are likewise rapidly changing in their scientific profiles and potential.

Creation of each volume required the editors involvement, vision, enthusiasm, and time. The Series Editor thanks each Volume Editor who graciously accepted his invitation. Special thanks go to Ms. Anita Lekhwani, the Series Acquisitions Editor, who extended the invitation to edit this series to me and has been a wonderful help in its assembling process.

ANDRZEJ WIECKOWSKI
Series Editor

Foreword

Despite extensive efforts, the electrochemical interface, where central processes in electrochemical reactions occur, had long been a black box until researchers started to shed light on it in the 1960s. At the beginning, the light used was mostly that in the visible region, but information obtainable by visible light was very limited, and the use of vibrational spectroscopy, which can provide detailed information on molecules, was strongly desired. In 1974, an innovation was made in spectroelectrochemistry by the application of Raman spectroscopy. It is well known that the Raman study of pyridine adsorbed on silver electrodes led to the discovery of surface-enhanced Raman scattering (SERS). The application of infrared reflection absorption spectroscopy (IRAS) to electrochemical interfaces in 1980 also was a great achievement. Surface-enhanced infrared absorption (SEIRA), an effect similar to SERS, was discovered in the same year, and the first observation of sum frequency generation (SFG) from monolayers on solid surfaces was made in 1986, although their applications to electrochemical interfaces were somewhat delayed. During the last four decades, these surface vibrational spectroscopy techniques have been advanced greatly owing to the improvement in instrumentation and the development of experimental techniques, which are still further developing year by year.

Vibrational spectroscopy is a powerful tool to identify molecules and to study their structures and reactions, as it is often mentioned that *vibrational spectra are letters from molecules*. It is also the case at surfaces and interfaces. It provides us information on adsorbed structures and orientations of molecules. Spectra are sensitive to changes

in the environment at interfaces, from which we can obtain deeper insight into chemistry and physics at the interfaces. In situ, time-resolved monitoring of reactions taking place at surfaces and interfaces is also possible. Owing to these advantages, they have gained wide application, from fundamental electrochemistry to many other related fields of science and technology, including surface science, heterogeneous electrocatalysis, energy conversion, biochemistry, nanotechnology, and sensors. However, each technique has strong and weak points. Difficulty in interpretation of the obtained spectra is another problem. For appropriate use of the techniques and for correct interpretation of spectra, sufficient fundamental understanding of the techniques is required.

This book, which features all the aforementioned four surface vibrational spectroscopy techniques and their applications to recent research topics, will provide fundamental information for nonspecialists and an up-to-date account of recent advances in this field for specialists.

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Preface

Electrified interfaces play an important role in many phenomena. Electric fields that develop at junctions between different phases can align molecules and ions into configurations that greatly influence the physical and chemical nature of the interface. The molecular structure of charged interfaces impacts many practical processes, including energy conversion in batteries, solar cells, and fuel cells, corrosion at solid surfaces, chemical reactions over oxide particles in the earth and atmosphere, biochemical transformations, signal transduction in chemical sensors, and heterogeneous catalytic reactions, to name a few.

Surface vibrational spectroscopy techniques probe the structure and composition of interfaces at a molecular level. Their versatility and typically nondestructive nature often enable in situ measurements of operating devices and monitoring of interface-controlled processes under reactive conditions. This book highlights modern applications of Raman, infrared, and nonlinear optical spectroscopy in the study of charged interfaces.

Early chapters in the book provide a glimpse into the breadth of systems that can be investigated through the use of nonlinear optical techniques. Properties of interfacial water, ions, and biomolecules at charged dielectric, metal oxide, and electronically conductive metal catalyst surfaces, as probed by nonlinear optical techniques, are discussed in Part I. In addition to examples of practical experimental interest, the chapters guide readers to the latest in measurement and instrumental techniques. Part II includes coverage of Raman spectroscopy from the standpoint of sensitive approaches for detection of biomolecules at solid-liquid interfaces and the use of photon depolarization

strategies to elucidate molecular orientation at surfaces. Part III reports on wide-ranging systems from small fuel molecules at well-defined surfaces to macromolecular complexes as building blocks of functional interfaces in devices that have applications in chemical sensing and electric power generation. These interfaces are amenable for infrared spectroscopy due to versatile sampling methods, that is, specular and diffuse reflectance, polarization-modulation, and total internal reflection modes.

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Part One

Nonlinear Vibrational Spectroscopy

Chapter 1

Water Hydrogen Bonding Dynamics at Charged Interfaces Observed with Ultrafast Nonlinear Vibrational Spectroscopy

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1.1 Introduction

The question of how charged species affect water structure and dynamics is relevant to many applications in chemistry, biology, geology, and industry. Biological systems are often crowded aqueous environments filled with proteins, membranes, vesicles, and other structures that often rely on the presence of ions for stability and proper functioning [1-6]. The ion-water interface is critical for ion exchange resins [7, 8], heterogeneous catalysis [9-11], electrochemistry [12], as well as processes involving mineral dissolution [13, 14] and ion adsorption [15, 16]. Because the behavior of water in the presence of ions impacts a wide range of technical and scientific fields, a great deal of literature over the years has been dedicated to studying the aqueous

solvation of ions and the properties of water at charged interfaces. Studies that have examined ion-water interfaces have employed x-ray and neutron diffraction [17-19], Raman spectroscopy [20], ultrafast infrared spectroscopy [21-26], Fourier transform infrared (FTIR) spectroscopy [20, 27], and other spectroscopic techniques [15, 28, 29]. Theoretical models [30, 31], molecular dynamics (MD) simulations [32-35], and Monte Carlo (MC) calculations [20] have also been employed. While simulations can provide some insight into the underlying dynamics, most experimental techniques only provide steady-state data. Here we utilize ultrafast infrared spectroscopy to examine the hydrogen bonding dynamics of water at several types of charged and uncharged interfaces.

Ultrafast infrared spectroscopy has been shown to be a powerful technique for elucidating dynamics in water-ion systems [21-26], other hydrogen bonding systems [36-43], protein environments [44-52], and systems that undergo chemical exchange [25, 53-57]. Here, we apply ultrafast infrared pump-probe and two-dimensional infrared (2D IR) vibrational echo spectroscopic techniques to examine the dynamics of water when it is confined in nanoscopic environments and interacting with interfaces. The question is whether the nature of confinement or the chemical composition of the interface most significantly influences the dynamics. To explore this question, the dynamics of water at charged and neutral interfaces in reverse micelles are compared. In addition, water in ionic solutions is investigated. Some water molecules are hydrogen bonded to ions, while others are hydrogen bonded to water molecules. These are in equilibrium, with water molecules bound to ions switching and becoming bound to water molecules, and vice versa. Using 2D IR chemical exchange spectroscopy, we determine the exchange time required for a water hydroxyl initially hydrogen bonded to an anion to

switch to being hydrogen bonded to another water molecule.

Reverse micelles consist of a water pool surrounded by a layer of surfactant molecules and are often used as model systems for confined environments. The surfactant molecules are terminated by a hydrophilic head group that can be either charged or neutral. These hydrophilic head groups face in toward the water pool while the alkyl (hydrophobic) tails of the surfactant are suspended in a nonpolar organic phase. A schematic of a reverse micelle utilizing the surfactant Aerosol-OT, or AOT [sodium bis(2-ethylhexyl) sulfosuccinate], is shown in [Figure 1.1](#). The AOT surfactant ([Fig. 1.2](#)) forms spherical monodispersed reverse micelles that have been well characterized. The size of the AOT reverse micelles can be easily controlled by varying the amounts of starting materials according to the w_0 parameter: $w_0 = [\text{H}_2\text{O}]/[\text{surfactant}]$ [58–60]. AOT can yield sizes of $w_0 = 0$ (essentially dry reverse micelles) all the way up to $w_0 = 60$, which has a water pool diameter of 28 nm and contains $\sim 350,000$ water molecules [61]. Isooctane is a common solvent used as the nonpolar phase of AOT reverse micelle systems, but other solvents such as carbon tetrachloride, cyclohexane, and benzene can also be used with minimal changes in water pool size for a given w_0 [62]. A recent study has shown that the identity of the nonpolar phase has no effect on the water pool dynamics [63].

[Figure 1.1](#) Illustration of the reverse micelle interior. The bulk water core is surrounded by a layer of interfacial water. The total water pool diameter is denoted by d . The hydrophilic AOT head groups face in toward the water pool while the alkyl tails are suspended in the organic phase. The sodium counterions are dispersed in the water pool, but they generally reside close to the head group interface.

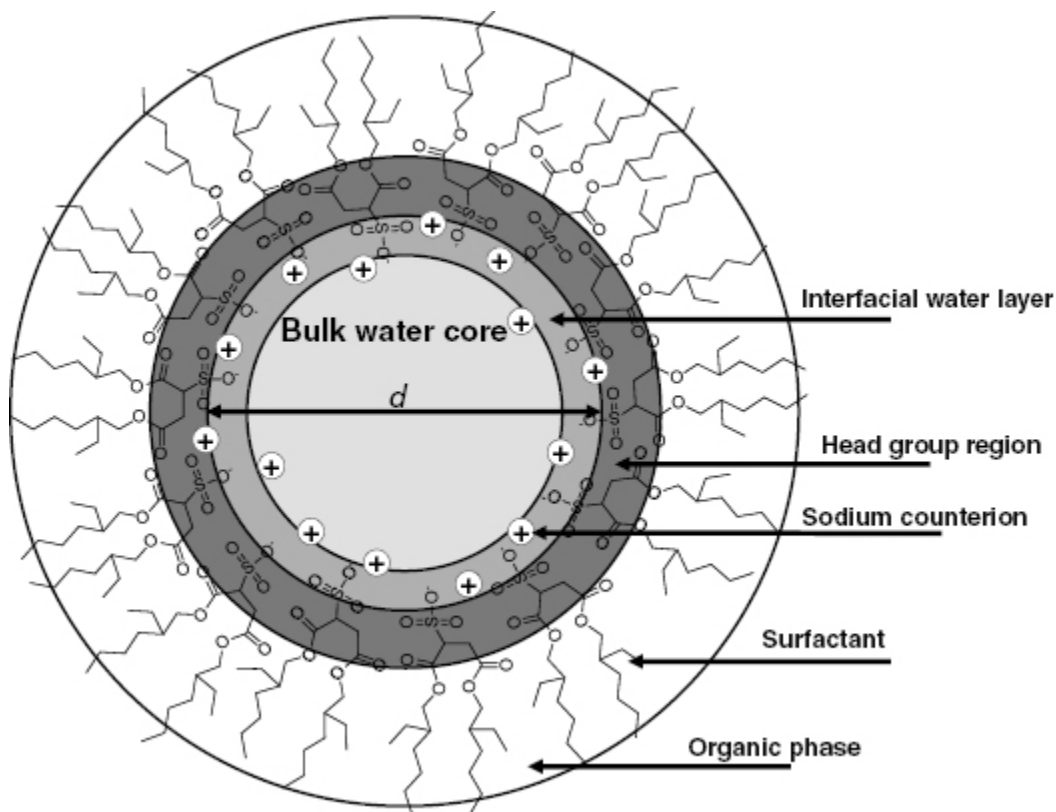
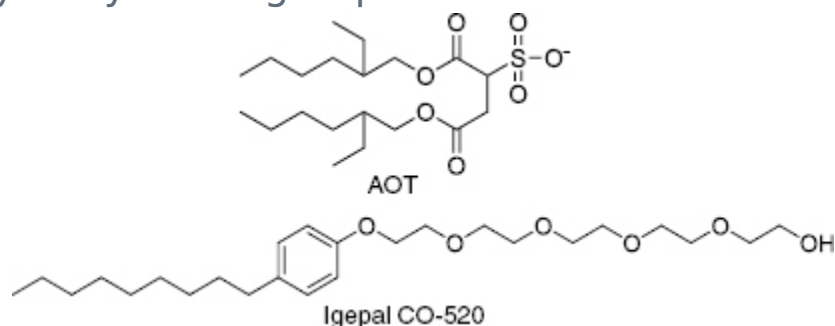


Figure 1.2 Molecular structures for AOT and Igepal CO-520. AOT (top) is terminated by a charged sulfonate head group with a sodium counterion while Igepal (bottom) has a neutral hydroxyl head group.



As shown in [Figure 1.2](#), AOT has a sulfonate head group with a sodium counterion. The head group region of the reverse micelle therefore creates a charged interface that surrounds the water pool. The sodium ions will generally reside in a region close to the interface. [Figure 1.1](#) illustrates the regions of a reverse micelle. When the total water pool diameter, d , is sufficiently large (≥ 4.6 nm) the reverse

micelle can support a core of water with bulklike properties. Below we will discuss how far perturbations from the charged sulfonate interfacial region extend into the water pool and what happens to the water dynamics as the size of the water pool changes in size. The chemical identity of the surfactant layer can be changed by using a neutral surfactant molecule called Igepal CO-520 ([Fig. 1.2](#)). Igepal is terminated with neutral hydroxyl head groups, so the interfacial water molecules will be exposed to a very different chemical surface compared to the AOT reverse micelle system. To what extent changes in surfactant identity, particularly charged versus neutral head group regions, and reverse micelle size affect water dynamics will be described.

Water dynamics are investigated through the processes of orientational relaxation, spectral diffusion, and vibrational relaxation, which can be measured with ultrafast infrared vibrational spectroscopy. These observables report on how the hydrogen bond network of water evolves and rearranges over time. The hydroxyl stretch of water is monitored during the experiments and is used as a reporter for hydrogen bond dynamics. During vibrational relaxation, vibrational energy dissipates by transferring into a combination of low-frequency modes, such as torsions and bath modes [64, 65]. Energy must be conserved during this process. Certain pathways that facilitate vibrational relaxation in one system may or not be present in a different system. Thus, vibrational relaxation is extremely sensitive to local environments. Orientational relaxation measures how quickly water molecules reorient by monitoring the direction of the transition dipole of the hydroxyl stretch. Molecular reorientation is involved in water hydrogen bond exchange, which leads to global hydrogen bond network reorganization [66, 67]. Bulk water consists of an extended network of hydrogen bonds that are continually rearranging and

exchanging with one another. According to the theory of Laage and Hynes, water molecules exchange hydrogen bonds via a jump reorientation mechanism that involves concerted motions of water molecules in the first and second solvation shells [66, 67]. The mechanism proceeds when a molecule in the second solvation shell of another water molecule moves in toward the first solvation shell. In order to swap hydrogen bonds with the approaching water from the second solvation shell, a water molecule must pass through a five-coordinate transition state and then undergo a large-amplitude rotational motion (or “jump”). The jump allows it to switch one of its hydrogen bonds to the approaching water molecule. These large-amplitude jumps change the orientation of the transition dipole. Solutes and interfaces (such as the surfactant shell of the reverse micelles) can disrupt the jump reorientation mechanism, thus slowing down the process of reorientation [23, 68–72]. Both vibrational and orientational relaxation can be measured with ultrafast infrared pump-probe spectroscopy.

Ultrafast 2D IR vibrational echo spectroscopy is used to measure spectral diffusion of the water hydroxyl stretch. The linear infrared absorption spectrum of the hydroxyl stretch is very broad due to a large distribution in the lengths and strengths of hydrogen bonds. At the beginning of the 2D IR experiment, a hydroxyl will vibrate at a certain frequency, but due to dynamic structural evolution of the system, that frequency will change over time. This process of frequency evolution is known as spectral diffusion and reports on how quickly water molecules sample different structural environments.

In addition, 2D IR vibrational echo chemical exchange spectroscopy is used to examine how quickly a water hydroxyl bound to an anion will switch to being hydrogen bonded to a neighboring water hydroxyl. This process is illustrated schematically in [Figure 1.3](#). A model system for