I. INTRODUCTION

Confined water plays an integral role in the structure and function of an enormous range of nanoscale systems. One notable class of systems is mesoporous silica,1−11 of which the powder MCM41 is a prominent example. These materials are widely used as a solid support for catalytic groups in heterogeneous reactions, in which water often acts as a key reactant or solvent.12 In addition to providing a high density of active sites by virtue of a large surface area to volume ratio, the surface of the nanoscale silica pores has a dramatic impact on the behavior of the water. For example, confinement of water in MCM41 has enabled scientists to explore the dynamics of deeply supercooled water below its homogeneous nucleation temperature,3,13 providing the first evidence of a fragile-to-strong dynamic crossover hypothesized to exist in bulk water.

The structure of an MCM41 particle (Figure 1A) consists of uniform cylindrical pores (variable from 1.5 to 10 nm) that pack into a hexagonal 2D array along a particular direction. In the orthogonal direction, the pores extend in a parallel fashion for the study of water H-bond dynamics in confined systems. These methods monitor fluctuations in hydrogen bond network configurations through variations in the orientation and frequency of a vibrational mode.5,20−22 Although water is the molecule of interest, direct study of the hydroxyl stretch of
water in MCM41 with PSPP and 2D IR experiments poses several experimental issues. One unavoidable problem is the short vibrational lifetime (<2 ps) of the water hydroxyl stretching mode.\(^\text{21}\) The resulting low signal-to-noise ratio at long time delays limits the observation of slower dynamics which are often the main signature of the confinement. Other disadvantages include pronounced heating artifacts\(^\text{23}\) and nonexistent 2D IR spectral advantages include pronounced heating artifacts\(^\text{23}\) and nonexistent 2D IR spectral stretching mode.\(^\text{21}\) The resulting low signal-to-noise ratio at long time delays limits the observation of slower dynamics which are often the main signature of the confinement. Other disadvantages include pronounced heating artifacts\(^\text{23}\) and nonexistent 2D IR spectral advantages include pronounced heating artifacts\(^\text{23}\) and nonexistent 2D IR spectral stretching mode.\(^\text{21}\) The resulting low signal-to-noise ratio at long time delays limits the observation of slower dynamics which are often the main signature of the confinement. Other disadvantages include pronounced heating artifacts\(^\text{23}\) and nonexistent 2D IR spectral advantages include pronounced heating artifacts\(^\text{23}\) and nonexistent 2D IR spectral

The present work combines PSPP and 2D IR measurements with molecular dynamics (MD) simulations to analyze time and space-resolved dynamics in water confined in MCM41 with \(\sim 2.4\) nm diameter pores. The selenocyanate anion (SeCN\(^-\)) was selected as a probe of the surrounding water dynamics because it avoids many of the issues associated with studying the hydroxyl stretch of water directly, yet its spectral diffusion has been demonstrated to closely track the time scales of water in the bulk liquid.\(^\text{22}\) An empirical frequency map has been developed for SeCN\(^-\) in water,\(^\text{22}\) providing a route to calculate its spectral diffusion in confined water. The long CN stretch vibrational lifetime (\(\sim 36\) ps) permitted the observation of complete orientational relaxation and spectral diffusion in the aqueous phase of MCM41 for the first time, although not without the need for several experimental improvements. Powdered solids are challenging to study with PSPP and 2D IR experiments, as these techniques are susceptible to scattered light from the pulses that generate the signal.\(^\text{24,25}\) The scattered light was reduced in this study using a combination of methods, ranging from the sample preparation procedure to a newly developed phase cycling scheme. In the 2D IR experiments, a previously reported method of combining mid-IR frequency-domain pulse shaping techniques\(^\text{26,27}\) and a polarization filter\(^\text{17}\) was successfully employed to reveal water—anion hydrogen bond dynamics in the MCM41 pores. A new phase cycling procedure was developed for the PSPP experiment, which improved the quality of data acquired in the scatter prone “parallel” polarization configuration. The strong transition dipole of the CN stretch of SeCN\(^-\) also facilitated the discrimination of the desired third-order signals from scatter.

II. EXPERIMENTAL PROCEDURES

II.A. Particle Size Reduction. MCM41 was purchased from Sigma-Aldrich. The as-received powder generated considerable scatter in the optical experiments. Because the experiments were performed in transmission mode, forward scattering of the excitation pulses into the signal detection direction (probe pulse direction, see below)\(^\text{27,28}\) posed an issue. The first strategy used to combat scatter was to reduce the average particle size of the MCM41 sample. It is well-known that when the wavelength and particle are comparable in size, the amplitude of scattering increases with the size of the scattering particle.\(^\text{29}\) A sedimentation procedure, outlined in the Supporting Information, was used to select smaller particles from the as-received powder. Briefly, MCM41 was placed on top of a column of water. Following 10 min, the top fraction, \(\sim 8\) in. or 20 cm of water, was vacuum filtered, and the particles were collected for later use. The duration and length of the column in principle selected particles with diameters less than \(\sim 40\ \mu m\). The entire procedure was repeated multiple times to compensate for the low yield of smaller particles.

II.B. Surface Area, Pore Size Distribution, and Pore Volume of MCM41. The as-received and smaller particle (see section II.A) MCM41 samples were characterized with N\(_2\) (g) adsorption and desorption isotherms (Figure S1A) acquired at the Soft & Hybrid Materials Facility (SMF) at the Stanford Nano Shared Facilities (SNSF). The area of the mesopore-free surface was assessed by generating BET plots\(^\text{30}\) (Figure S1B) from the adsorption curves in Figure S1A. The mean specific BET surface area, \(S_{\text{BET}}\), was found to be 960 \(\pm 30\) m\(^2\)/g from four independently prepared samples. The pore diameter distributions for each sample (Figure S1C) were obtained from BJH analysis\(^\text{17}\) of the sorption isotherm curves in Figure S1A and are displayed in Figure S1C. Analysis of the adsorption or desorption curves gave identical results. The narrow
The inner diameter of the smaller spacer was 8 mm. The primary mesopore volume was also obtained from a 3D-simulated silica pore model used in the simulations (2.4 nm). The homogeneity of the material was further confirmed in the experiments, with high reproducibility observed for both the rotational and spectral diffusion measurements made on several samples. The primary mesopore volume was also determined to be 0.8 ± 0.1 cm³/g. Further details on the methods and calculations are provided in the Supporting Information.

II.C. Sample Hydration. Details of the sample hydration are provided in the Supporting Information. Solutions and powders described in this section were prepared and manipulated under an N₂(g) atmosphere. Briefly, completely dry MCM41 powder (of smaller particle size) was gently mixed in 0.3 M KSeCN/D₂O solution. The particles were filtered to remove bulk solution from the sample. Variable degrees of hydration were obtained sample-to-sample following the filtration step, and it was found that the measured dynamics were very sensitive to the amount of solution in the sample. To obtain reproducible dynamics, each sample was equilibrated overnight in a home-built hydration chamber (Figure S2). The D₂O vapor pressure in the hydration chamber was maintained at 78% or 100% RH by continuously bubbling N₂(g) through a saturated NaCl/D₂O or pure D₂O reservoir and flowing the vapor over the powder. The % RH was chosen to lie beyond the capillary condensation region in the room-temperature water sorption isotherm of MCM41 (P/P₀ = 0.06) to fully hydrate the uniform primary mesopores (D = 2.5 ± 0.2 nm) in the sample. The water sorption isotherm has noticeable hysteresis between the adsorption and desorption curves. We note that if the equilibration vapor pressure is too close to the hysteresis loop, markedly different hydration states can be obtained depending on the initial hydration status of the sample. The simulated dynamics were compared to the samples hydrated at 100% RH. The 78% RH data are tabulated in the Supporting Information.

II.D. Sample Cell Preparation and Linear-IR Spectroscopy. The sample cell preparation used in the linear and nonlinear-IR experiments is detailed in the Supporting Information and briefly described here. The sample cell is similar to one described previously. The path length was set to 25 µm with two concentric polytetrafluoroethylene (PTFE) spacers with outer diameters of 25.4 and 13 mm, respectively. The inner diameter of the smaller spacer was ∼8 mm. The MCM41 powder, prepared according to section II.C, and an index matching fluid (paraffin oil) were placed in the center of the smaller spacer. The region between the inner and outer spacers was filled with paraffin oil. The spacers were sandwiched between two CaF₂ windows with 25.4 mm diameters and 3 mm widths. Fourier transform IR (FT IR) spectra of the KSeCN/D₂O containing MCM41 were measured with a Thermo Scientific Nicolet 6700 FT IR spectrometer purged with air scrubbed free of CO₂ and H₂O. The absorption spectrum of the silica powder containing solely D₂O with identical spacer thicknesses was measured and subtracted from the resonant sample spectrum.

II.E. 2D IR Spectroscopy. The 2D IR spectrometer used in this work has been described in detail in past publications. The spectrometer is configured in a pump–probe geometry and features a mid-IR pulse shaping system in the excitation beam path (see Figure S3A). 2D IR and PSPP spectroscopies are resonant, third-order time-domain experiments. In 2D IR, three pulsed electric fields interact with a material through its third-order susceptibility. Following three interactions, the induced polarization in the material emits a fourth field known as the vibrational echo signal. In the pump–probe geometry, phase matching dictates that the signal is emitted collinearly with pulse 3, the probe pulse. In addition to stimulating the echo emission, pulse 3 also serves as the local oscillator for heterodyne detection of the echo signal. In the limit of very short pulses (semi-impulsive limit), the polarization is proportional to the response function of the molecular system. The response function is dependent on the dynamics of the ensemble. Time-resolved dynamics are obtained from the data in the form of two-point time correlation functions (TCFs). The pulse sequence is shown in Figure 1C. The measured third-order response functions are 3D functions of the time delays, t₁ = t, t₂ = Tω and t₃. A more straightforward manner to visualize and analyze the response functions is to perform a 2D Fourier transform on the t₁ and t₃ axes to generate a 2D IR spectrum, S(ωᵣ,ωₚ,ω₃). A 2D IR spectrum is a correlation plot obtained from the data in the form of two-point time correlation functions (TCFs).

Qualitatively, 2D IR works as follows. There are four pulses in the pulse sequence (Figure 1C). Three input pulses generate the nonlinear echo, which gives rise to the emission of a fourth pulse, the vibrational echo. The first two pulses label and store the initial frequencies, which form the horizontal axis of the 2D spectrum. As the water structure evolves, the intermolecular interactions between water molecules and the vibrational probes change. This change in local and mesoscopic distance scale interactions changes the frequencies of the vibrational probes. The third pulse and echo read out the final frequencies, which form the vertical axis. As Tω increases, the water structure interacting with the probes has more time to evolve, leading to more frequency change and therefore more change in the 2D IR line shape. The Tω dependence of the 2D line shape is analyzed to extract the time dependence of the water structure.

II.F. Polarization Selective Pump–Probe Spectroscopy with Scatter Reduction. PSPP spectroscopy can be regarded as a special case of 2D IR spectroscopy where the time delay, t₁, separating the first two pulses is fixed to zero (see Figure 1C). The first two sample-field interactions occur simultaneously with the arrival of a single pump pulse. The third interaction occurs when the probe pulse arrives an interval, t (t₂ = Tω in the echo experiment, Figure 1C), later. The PSPP signal is the transient difference in the probe intensity with the pump pulse on versus off. To acquire the signal, the pump is chopped at 500 Hz, or half the laser repetition rate, with an acousto-optic modulator (AOM). The
signal is measured in two polarization configurations termed “parallel” and “perpendicular”. The probe is linearly polarized at 0° (in the plane of the optical table), and the pump is linearly polarized at 45° relative to the probe. After the sample, a polarizer in a computer-controlled rotation mount resolves the signal at +45° or −45°, which correspond to parallel, S∥(t), and perpendicular, S⊥(t), configurations, respectively.

In the measurement of a scattering sample, the total integrated intensity on the array detector,23 for laser shots where the pump is on, can be written

\[ S_{on} \propto |E_{p}] + E_s + sE_p|^2 \]

\[ = S_{off} + |E_p|^2 + E_p^*E_s + s^*E_pE_p^* + s^*E_sE_p^* + \ldots \quad (c.\ c.) \]

where \( E_p \), \( E_s \), and \( sE_p \) refer to probe, signal, and scattered pump fields (\( s \) is a complex scatter coefficient), respectively, \( S_{off} = |E_p|^2 \), and the small homodyne signal, \( |E_p|^2 \), has been neglected. The explicit time dependence of the terms has been omitted. The signal of interest is the third term on the second line of eq 1. As discussed in the Supporting Information, the homodyne scatter term \( |E_p|^2 \) is independent of \( t \) and can be straightforwardly accounted for by measuring the offset in the data at negative \( t \) delays (probe arriving before pump). The heterodyne scatter terms (final two terms on the second line of eq 1) not only vary in amplitude but also oscillate at the optical frequency as \( t \) is scanned (Figure S3B). To remove the heterodyne scatter, a four-shot scatter removal phase cycle (eq S7) was used to acquire the data. In addition to this, for each delay \( t \), the delay was scanned over 200 fs, i.e., \( t - \epsilon \rightarrow t + \epsilon \) (\( \epsilon = 100 \) fs), using the pulse shaper. As this scanning range is comparable to the optical pulse duration, the pump–probe signal level is essentially invariant; while the heterodyned scatter exhibits an oscillation at the optical frequency (Figure S3B), with the desired pump–probe signal representing the average of the oscillatory signal. Thus, the pump–probe signal of interest and the heterodyned scatter can be clearly distinguished in this procedure. The acquired data were averaged over six optical periods to remove residual scatter fringes present following phase cycling.

III. SIMULATION DETAILS

III.A. System Modeling. The dynamics of SeCN− in D2O confined in nanoscale silica pores were investigated using MD simulations based on a combination of previously developed models. In particular, the fixed-charge force field for SeCN− previously obtained from a combination of electronic structure calculations and comparisons to experiments in bulk D2O was used.22 The parameters, developed using the SPC/E water model15 for D2O, are given in Table 1 of ref 22. In brief, the key elements of the model are a symmetric charge distribution \( q_{Sa} = q_{N} = -0.55; q_{C} = 0.1 \) and a larger Lennard-Jones radius for Se (4.0 Å) compared to that of C and N. The SeCN− is treated as a rigid molecule, while the D2O bonds and angles are fixed using the SHAKE algorithm.43 This model was shown to give excellent agreement with the measured reorientational anisotropy from PSP experiments in bulk D2O.22

The silica pore model (Figure 1B) is based on those previously developed by Gulmen and Thompson.44,45 In this approach, the pore is generated by first heating a simulation cell of crystalline SiO2 to 8000 K to generate molten silica. A cylinder that interacts only through a Lennard-Jones potential is then slowly grown in the center of the simulation cell, displacing the liquid SiO2 to form a pore. The simulation temperature is subsequently reduced to room temperature at a rate fast enough to prevent crystallization (e.g., 50 K/ps). This process results in an amorphous silica pore that is not functionalized. To provide the typical –OH termination of mesoporous silica, a hydrolysis reaction is simulated by choosing elongated Si–O bonds near the pore surface and adding an –OH to the Si and an H to the O atom; this procedure ensures charge neutrality in a straightforward way. Changing the parameters used to identify the Si–O bonds can modify the surface hydroxyl density. The pore model used in the present simulations is ~2.4 nm in diameter with 2.5 OH/\( \text{nm}^2 \) (Figure 1B).

The interactions of the pore with the D2O liquid and SeCN− solute are through Lennard-Jones and Coulombic interactions, with the parameters for the pore atoms given in Table 1 (\( \text{O}_\text{h} \)).

Table 1. Silica Pore Force Field Parameters

<table>
<thead>
<tr>
<th>atom</th>
<th>(Å)</th>
<th>(kcal/mol)</th>
<th>(e)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Si</td>
<td>2.500</td>
<td>0.00010</td>
<td>2.10</td>
</tr>
<tr>
<td>O\text{ bonded}</td>
<td>2.700</td>
<td>0.45686</td>
<td>−1.05</td>
</tr>
<tr>
<td>O\text{ hydrogen}</td>
<td>3.070</td>
<td>0.16995</td>
<td>−0.95</td>
</tr>
<tr>
<td>H</td>
<td>1.295</td>
<td>0.00037</td>
<td>0.425</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>angle</th>
<th>(kcal/mol/rad(^2))</th>
<th>(deg)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Si–O–H</td>
<td>24.499325</td>
<td>118.5</td>
</tr>
<tr>
<td>HO–Si–OH</td>
<td>38.155673</td>
<td>118.5</td>
</tr>
</tbody>
</table>

indicates a bridging, or siloxane, oxygen); Lorentz–Berthelot mixing rules were used. The pore atoms are held rigid except for the hydroxyls of the silanol, Si–OH, and geminal, Si(OH)\text{2}, groups at the pore surface, for which bonds are held fixed by the SHAKE algorithm but bond and dihedral angles are flexible (Table 1). For the geminal groups, no intramolecular dihedral potential is used, but the 1–4 (O–Si–O–H) and 1–5 (H–O–Si–O–H) Lennard-Jones and Coulombic interactions are included, scaled by a factor of 0.5 and 1, respectively. The charges used in the present simulation are taken from the ClayFF model46 and are larger in magnitude than those previously used to successfully simulate confined liquids,45,47,48 including vibrational spectra.49,50 This choice is derived from comparisons of the present simulation results and measurements (vide infra). The interactions were calculated with a cutoff of radius 12 Å and the long-range electrostatics were accounted for with the damped-shifted force method51 with a damping parameter of 0.2.

The simulation was prepared by carrying out grand canonical Monte Carlo (GCMC) simulations in which water molecules are inserted and removed from the pore to equalize the chemical potential with that of bulk liquid water; this provides the equilibrium density in the pore corresponding to complete filling (100% RH). These GCMC calculations were completed using our previously developed pore charges.44,45 The SeCN− solute was inserted by replacing one of the water molecules. This gives 448 water molecules and one solute in the pore. While this is a lower concentration than used in the experiments, both are in the dilute limit with no effect from interactions between solutes. Moreover, the K+ counterion present in the experiments is not included because no effects of ion pairing, e.g., a blue-shifted shoulder in the linear-IR spectrum, were observed.\textsuperscript{62,63} The results presented are obtained from ten 18 ns trajectories in the NVT ensemble.
with the temperature maintained at 298 K by a Nosé–Hoover thermostat,\textsuperscript{54,55} with a time constant of 100 fs. The time step is 1 fs, and configurations are written every 12 fs following a 2.5 ns equilibration stage, with velocity rescaling used to maintain the temperature over the first 0.5 ns. Each trajectory differs in the initial velocities, randomly selected from a Boltzmann distribution. In most cases, error bars on MD-derived data were calculated from block averaging (with each of the 10 trajectories as a block) and are reported as 95% confidence intervals using the Student’s t-distribution.\textsuperscript{56} For multi-exponential fits, the errors in the amplitudes and time scales are reported as a single standard deviation from five blocks (each consisting of an average of two trajectories); this approach is adopted to address issues of heterogeneity, i.e., the sensitivity of the time correlation function decay to the fraction of time SeCN\textsuperscript{−} spends in the interfacial layer in a given trajectory.

III.B. Spectroscopic Modeling. The vibrational spectra of the CN stretching mode of SeCN\textsuperscript{−} were calculated using the empirical frequency map developed previously for simulations in bulk D\textsubscript{2}O.\textsuperscript{22} The map is based on density functional theory calculations of the vibrational frequency in clusters extracted from MD simulations in the bulk liquid. Each cluster is centered on the SeCN\textsuperscript{−} solute with nine waters explicitly considered and remaining waters within 10 Å of the center-of-mass included as point charges. The field used in the map is that from all waters within this 10 Å distance using the SPC/E model charges.\textsuperscript{57} The map parameters are given in Table 2 of ref 22; the frequency is found to vary linearly with the field.

In the confined environment, there is a contribution to the field from the pore atoms, and because the map is derived from SeCN\textsuperscript{−} in bulk D\textsubscript{2}O, there is a choice about how those contributions should be calculated. The results presented are based on using the charges for all atoms within 10 Å of the SeCN\textsuperscript{−} center of mass, but we found only modest differences when the field from all pore atoms was used instead.

IV. RESULTS AND DISCUSSION

IV.A. Linear-IR Spectrum. The linear-IR spectra of SeCN\textsuperscript{−} (CN stretch) in bulk D\textsubscript{2}O (black solid curve) and D\textsubscript{2}O confined in the silica pore (red solid curve) are compared in Figure 2. Nanoscale confinement has no effect on the band center and full width at half-maximum (fwhm), which are 2075 and 33 cm\textsuperscript{−1}, respectively, for both systems (Table 2). This result is analogous to the surprising invariance of the water OH stretch spectrum upon confinement in silica glasses with different pores sizes\textsuperscript{2} and in MD simulations of water confined in a silica pore model similar to that used here.\textsuperscript{10} The simulations revealed that the line shapes for OH groups rapidly approached the bulk spectrum as they moved only a small distance away from the surface, beginning at as little as 3 Å between the hydroxyl H atom and nearest pore O atom. In other words, only OH groups that were hydrogen bonded to the pore exhibited perturbed frequencies.

However, the enhanced absorption on the red edge of the bulk spectrum is absent in the confined spectrum (Figure 2). The asymmetric bulk line shape has been attributed to the frequency dependence of the CN transition dipole moment, $\mu_{01}$, in D\textsubscript{2}O, which results from the variation in H-bond number and strength across the line.\textsuperscript{22} The disappearance of the red wing results in a symmetric line shape in the pore. Insight can be gained from the simulations by computing, in addition to the linear-IR absorption spectrum, the frequency distribution, $P(\omega) = \langle \delta(\omega - \omega_{01}) \rangle$, and $P(\omega)$ weighted by the transition dipole moment squared, i.e., the spectral density, $P_{\mu} (\omega) = \langle |\mu_{01}|^2 \delta(\omega - \omega_{01}) \rangle$. The calculated quantities in the bulk liquid and silica pore systems are compared in Figure 3A. The simulated IR spectrum (blue curve) shows a minimal (<1 cm\textsuperscript{−1}) shift in the band center relative to that in bulk D\textsubscript{2}O (black curve), in agreement with the measurements. However, the modeling predicts significant broadening of the spectrum relative to that in the bulk liquid, with a fwhm that is 60% larger (Table 2). This broadening appears to be a consequence of both greater heterogeneity in the distribution of CN frequencies (red dashed curve), as shown in Figure 3A, for which the fwhm increases by 30% relative to that in bulk D\textsubscript{2}O (Table 2), and reduced motional narrowing. It is not fully clear in which of these aspects the modeling of the CN frequency is deficient, though based on the dynamical results below, it appears to be the former.

The simulated spectra can be calculated as a function of the SeCN\textsuperscript{−} position in the pore, based on the distance of the C atom from the nearest pore O atom, $d$, at $t = 0$ in the calculation of the response function.\textsuperscript{22} This is shown in Figure 3B. These results show a systematic red shift in the CN frequency as the molecule approaches the pore surface. At the same time, the line width increases as $d$ decreases. At the largest distance from the pore wall, $d = 10$ Å, the spectrum is

---

**Figure 2.** Linear-IR spectrum of SeCN\textsuperscript{−} in bulk D\textsubscript{2}O (black curve) and in D\textsubscript{2}O confined in the silica pore (red curve). Broadening on the red side of the spectrum is noticeably diminished in the pore.

**Table 2. Parameters for Line Shapes and Distributions**

<table>
<thead>
<tr>
<th>line shape/distribution</th>
<th>sample</th>
<th>center (cm\textsuperscript{−1})</th>
<th>fwhm (cm\textsuperscript{−1})</th>
</tr>
</thead>
<tbody>
<tr>
<td>expt $I(\omega)$</td>
<td>bulk</td>
<td>2074.7 ± 0.1</td>
<td>32.9 ± 0.1</td>
</tr>
<tr>
<td></td>
<td>pore</td>
<td>2075.1 ± 0.7</td>
<td>32.6 ± 0.7</td>
</tr>
<tr>
<td>sim $I(\omega)$</td>
<td>bulk</td>
<td>2075</td>
<td>38.5</td>
</tr>
<tr>
<td></td>
<td>pore</td>
<td>2074</td>
<td>61</td>
</tr>
<tr>
<td>$P(\omega)$</td>
<td>bulk</td>
<td>2077</td>
<td>59.6</td>
</tr>
<tr>
<td></td>
<td>pore</td>
<td>2077</td>
<td>78</td>
</tr>
<tr>
<td>$P_{\mu}(\omega)$</td>
<td>bulk</td>
<td>2072</td>
<td>58.8</td>
</tr>
<tr>
<td></td>
<td>pore</td>
<td>2071</td>
<td>77</td>
</tr>
</tbody>
</table>

\textsuperscript{*}Centers and fwhm’s for the measured (expt) and simulated (sim) FTIR spectra, $I(\omega)$, frequency distributions, $P(\omega)$, and spectral densities, $P_{\mu}(\omega)$, for 0.3 M SeCN\textsuperscript{−}/D\textsubscript{2}O in bulk solution and confined in the silica pore. The parameters for the bulk liquid are reproduced from ref 22.
Figure 3. (A) Simulated linear-IR spectrum (blue curve), frequency distribution (red dashed curve), and spectral density (violet dashed curve) of SeCN\(^-\) in D\(_2\)O confined in the silica pore; the simulated bulk IR spectrum (black curve) is shown for comparison. (B) Distance-dependent IR spectra (dashed curves) for SeCN\(^-\) in D\(_2\)O confined in the pore are compared to the total pore (blue curve) and bulk (black curve) IR spectra.

Figure 4. (A) Calculated probability of the total number of H-bonds donated to SeCN\(^-\) by D\(_2\)O and silanols (black points) and the number of axial H-bonds (red points) defined as \(\theta_{\text{CNOD}} \geq 120^\circ\). Results in the silica pore (solid lines) are compared to those for the bulk solution (dashed lines). (B) Average number of total (black circles) and axial (red circles) H-bonds donated to SeCN\(^-\) as a function of the C atom distance to the nearest pore O atom, \(d\). (C) Averaged CN stretching frequency as a function of the number of total (black points) and axial (red points) H-bonds. Results in the pore (solid lines) are compared to those for the bulk solution (dashed lines). (D) Average CN stretching frequency as a function of \(d\).

These values are slightly less than the average of 3.4 total H-bonds found in bulk D\(_2\)O. In contrast, the average number of axial H-bonds is largest next to the pore surface, oscillates between \(d = 4\) and 5 Å, and converges to 0.88 at larger distances. This gives the overall average number of axial H-bonds in the pore as 0.89 compared to 0.97 in bulk solution. Thus, the simulations show that confinement slightly inhibits the formation of H-bonds.

The effect of the H-bonding arrangement on the CN stretching frequency is illustrated in Figure 4C, where the average frequency is plotted versus the number of axial and total H-bonds. The results are quite comparable to those found in bulk D\(_2\)O and consistent with the picture described above for the frequency shifts; i.e., increasing the number of axial H-bonds increases the CN frequency while a greater number of total H-bonds decreases it. For a given number of total (axial) H-bonds, SeCN\(^-\) in the pore has a lower (higher) frequency compared to in the bulk solution. This suggests that the lack of a frequency shift in the IR spectrum upon confinement in silica pores results from the cancellation of different effects.

How the SeCN\(^-\) position in the pore affects the vibrational frequency is examined in Figure 4D, where the average frequency is plotted as a function of the distance, \(d\), from the nearest pore O atom. The frequency increases monotonically with \(d\) from 2074 cm\(^{-1}\) at \(d = 3\) Å to 2080 cm\(^{-1}\) at \(d = 10\) Å, though the change at small \(d\) is not well resolved due to the large uncertainties. This trend is not fully explained by the results in Figure 4B\(_c\). The convolution of those results would suggest the transition frequency would be the largest at the interface, reach a minimum at \(d = 5\) Å, and then increase with \(d\) thereafter; because the frequency is more sensitive to the total number of H-bonds, this is the dominant contribution. These differences suggest that the electrostatic environment at the interface cannot be understood solely in terms of H-bonding to the SeCN\(^-\) anion.
IV.B. PSPP Spectroscopy: Orientational Relaxation.

To analyze the orientational relaxation dynamics of the SeCN$^-$ anion in confined D$_2$O, the reorientational anisotropy was calculated from the PSPP signals with

$$r(t) = \frac{S(t) - S(0)}{S(t) + 2S(0)}$$

(2)

and from the simulations as

$$r(t) = \frac{2}{5} C_2(t) = \frac{2}{5} \langle 3 \ell(t) \cdot \ell(0) \rangle$$

(3)

where $C_2(t)$ is the orientational correlation function, $P_2$ is the second-order Legendre polynomial, $\ell(t)$ is the transition dipole moment unit vector at time $t$, and $\langle ... \rangle$ represents an ensemble average. The measured (solid points) and simulated (solid curves) $r(t)$ are displayed in Figure 5A. It is important to note that the form of eq 2 is only strictly true for a 3D isotropic system.$^{38,59}$ Assuming an average spherical particle size of $\sim 10$ μm, we estimated that within the $\sim 100$ μm waist and $\sim 25$ μm focal depth of the incident infrared beams there are $> 10^3$ particles that are randomly oriented. Therefore, the system can be regarded as macroscopically isotropic, and eq 2 can be used to extract $C_2(t)$.

The cylindrical pore structure of MCM41 may lead to a preferred orientation of SeCN$^-$, particularly near the silica surface. This possibility was addressed through the simulations by calculating the probability distribution of the SeCN$^-$ as a function of the C atom distance from the pore wall. This is shown in Figure 6 along with corresponding results for the water atoms. The results show that SeCN$^-$ is preferentially found away from the pore interface, with a broad distribution between $d = \sim 3.5$ and $\sim 10$ Å. This is a result of the strong affinity of water for the pore surface, which is indicated by the strongly peaked probability distributions for the water atoms at small $d$. At larger distances, the distributions for the water sites behave linearly, reflecting a uniform density convoluted with the linearly decreasing volume available as the pore center is approached. It is in this relatively unstructured water outside the interfacial layer that the SeCN$^-$ is predominantly located.

Quantitatively, the interfacial water layer can be roughly defined as $d < 4$ Å, and the solute is outside of this region 95% of the time. Because of this preference for the center of the pore, the SeCN$^-$ has a nearly uniform angular distribution.

In Figure 5A, it is seen that the calculated orientational relaxation in the simulated pore (black curve) agrees very well with the experimental measurements (red points). Excellent agreement was also obtained in the bulk case,$^{22}$ and those data (black points) are reproduced in Figure 5A for comparison.

The amplitude of the fast inertial fluctuations in $\ell(t)$ can be assessed by summing the amplitudes of multiexponential fits to the data (Table 3), which are associated with the observable dynamics, and calculating the deviation from 0.4 (eq 3). The amplitude of the inertial component is $\sim 0.02$ (within error) out of 0.4 total for both systems, representing a small contribution to the total orientational randomization. Thus, confinement perturbs the amplitudes.
and time scales of the diffusive motions of the anion, which lead to significantly slower orientational relaxation.

A useful measure of the change in dynamics, which weights each time scale component by its normalized amplitude, is the integrated correlation time, defined by

$$\tau_i = \frac{1}{A_i} \int_0^\infty A(t) \, dt$$

(4)

where $A(t)/A_i(0)$ represents a normalized correlation function, in this case $C_i(t)$. The integrated correlation time describing the reorientation of SeCN$^-$ is seen to increase by a factor of ~1.7 between the bulk (3.8 ps) and confined liquid (6.3 ps) (Table 3). This substantial slowing is remarkable considering the modest change in the linear-IR spectrum (Figure 2).

Similar results have been obtained for neat water confined in mesoporous silica.$^5,10$ Namely, the IR spectrum in the OH stretching region was found, in both simulations and measurements, to be insensitive to confinement, while the same simulations show dramatically slowed, nonexponential reorientational dynamics. Because of the considerably shorter OH vibrational lifetime (compared to SeCN$^-$), the complete water orientational relaxation is not readily measured. The anisotropy decay of SeCN$^-$ in the silica pore (red solid points, Figure 5A) is described well with a triexponential fit (Table 3, row 3). The data show fast time scales with magnitudes and amplitudes similar to the bulk time scales (Table 3, row 2), although modestly slower. The major difference is a smaller amplitude, long time scale ($t_3 = 25 \pm 8$) that is absent in the bulk liquid. In Figure 5B, the simulated SeCN$^-$ anisotropy is decomposed according to the distance of the solute from the pore wall (defined by $d$, the C atom distance to the nearest pore O atom) at $t = 0$. These data show that the orientational relaxation is slowest directly near the surface and gradually accelerates as the molecule moves further into the center of the pore. This is most notable in the decay at longer times which, as noted above, exhibits significant retardation relative to that in the bulk liquid. The results in Figure 5B show that this is primarily associated with SeCN$^-$ molecules that are in the interfacial layer of the pore. However, the change in the rotational dynamics is continuous as a function of the SeCN$^-$ distance from the pore surface, distinct from the assumption of two-state, or core−shell, models. Indeed, $C_i(t)$ can be fit well to a triexponential form for all distances except the largest considered ($d = 10 \AA$) where a biexponential fit is sufficient. It is interesting, however, that even at this distance the orientational relaxation is slightly slower than that in the bulk liquid, a result that is indicative of the small pore size.

It is interesting to further consider the distance dependence of $C_i(t)$ in the context of the typical models applied to confined liquids. Foremost among these is the two-state, or core−shell, model,$^{45,60–63}$ in which it is assumed that the property of interest in the interfacial layer (the "shell") is distinctly different than that in the pore interior (the "core"). Mathematically, it can be represented, for $C_i(t)$, as

$$C_2_{shell}(t; d) = C_{2,core}(t) + [C_{2,shell}(t) - C_{2,core}(t)]\theta(\Delta - d)$$

(5)

where $C_{2,core}(t)$ and $C_{2,shell}(t)$ are the orientational correlation functions characteristic of the interior and interfacial regions, respectively, $\Delta$ is the width of the interfacial layer, and $\theta$ is the Heaviside step function. It can be seen from Figure 5B that this does not even qualitatively describe the distance dependence of the orientational correlation function. The behavior of $C_i(t; d)$ with $d$ is more quantitatively examined in Figure 7,

![Figure 7](https://doi.org/10.1021/acs.jpcc.9b00593)

**Figure 7.** Simulated $C_i(t; d)$ as a function of the distance, $d$, between the C atom and nearest pore O atom evaluated at four delay times: 1.2 ps (black circles), 2.4 ps (red circles), 3.6 ps (blue circles), and 4.8 ps (violet circles). Dashed lines of the same color show fits to an exponential decay (eq 6).

where the values of $C_i(t; d)$ at particular times are plotted versus $d$. This behavior is clearly not that of a step function change from the interfacial layer to the core. Rather, as shown in Figure 7, the value at each time is well described by a decaying exponential

$$C_{2,exp}(t; d) = C_{2,core}(t) + [C_{2,shell}(t) - C_{2,core}(t)]e^{-(d-d_0)/\xi}$$

(6)

where $d_0$ is a reference distance taken to be 3 Å. However, these exponential fits yield different values of the length scale, $\xi$, for each time. The change in $\xi$ is ~20%. This change reflects the multieponential time dependence of $C_i(t)$ and indicates that the distance dependence for each amplitude and time scale is not the same.

The results presented here are consistent with results on small reverse micelles.$^{63}$ The core−shell model provides a
reasonable description of water orientational relaxation dynamics for large reverse micelles, in which the diameter of the water nanopool is greater than \( \sim 6 \) nm. For large reverse micelles, the water in the core displays bulk water dynamics and bulk IR spectra of the hydroxyl stretching mode. Water molecules directly interacting with the interface show substantially different spectra and dynamics than the bulk-like water core. However, even for large reverse micelles with a substantial bulk water center, there is a transition region between the interfacial dynamics and the bulk dynamics of the core. For reverse micelles with water nanopools less than \( \sim 4 \) nm, orientational relaxation measurements display no evidence of a distinct core. Like the experiments presented here, for small reverse micelles the data cannot be separated into core and shell components. However, simulations show that there is a change in the dynamics as the distance from the interface increases. The size of the nanopores studied here is in the range of the size of the water nanopools in reverse micelles that do not fit the core–shell model.

IV.C. 2D IR Spectroscopy: Spectral Diffusion. Nanoscale confinement in porous silica results in a significant change in the collective and single molecule orientational relaxation of water. Thus, it is also anticipated to have a large influence on the rearrangement of the water H-bond network. The time scales of these motions in bulk water have been the focus of many experimental and theoretical studies. However, direct study of water in the context of nanometer-size silica pores with nonlinear-IR spectroscopy has been limited by its short-lived vibrationally excited state, which prevents observation of the slower dynamics occurring in the confined environment. Furthermore, scattered light from the silica framework poses a nontrivial challenge to acquire and extract reliable time-dependent line shapes from mesoporous silica powders.

The spectral diffusion of SeCN\(^-\) in D\(_2\)O was recently analyzed in detail and shown to be remarkably similar to the measured spectral diffusion of HOD in D\(_2\)O. Water spectral diffusion is characterized by two main time scales. The first (\( \sim 0.4 \) ps) originates from structural fluctuations associated with local H-bond configurations. The final time scale (\( \sim 1.4 \) ps), which is within error, the same for HOD and SeCN\(^-\), is associated with the concerted intermolecular rearrangements that randomize the H-bond network around the probe. The rotational dynamics of the SeCN\(^-\) molecule were also shown to contribute negligibly to its own spectral diffusion through the absence of reorientation induced spectral diffusion (RISD). SeCN\(^-\) spectral diffusion in water is therefore mainly determined by H-bond network structural dynamics.

To quantify the nature of water H-bond dynamics in MCM41, measurements and simulations of SeCN\(^-\) spectral diffusion were performed in the water-filled silica pores. The full range of dynamics were measured with complete removal of scatter by acquiring spectra in an (XXYY) configuration with a four-shot phase cycle sequence. Representative spectra for \( T_w = 0.5 \) ps and \( T_w = 30 \) ps are shown in Figure 8. A clear change in the shape of the 0 \( \rightarrow \) 1 transition (red, positive going peak) is observed as spectral diffusion proceeds with increasing \( T_w \). The time-dependent 0 \( \rightarrow \) 1 band shape was analyzed using the center-line-slope (CLS) method as was described for SeCN\(^-\) in D\(_2\)O. The CLS decay is shown (red points) in Figure 9 alongside the data for the bulk system (black points) reproduced from the previously published study. Consistent with predictions, the H-bond network dynamics in the silica pore follows the CLS decay of the bulk system.

![Figure 8](image1.png)  
**Figure 8.** Measured 2D IR spectra of SeCN\(^-\) in D\(_2\)O confined in the silica pore. The spectrum is noticeably rounder and less tilted at the longer waiting time (\( T_w = 30 \) ps), indicating that spectral diffusion is more complete.

![Figure 9](image2.png)  
**Figure 9.** CLS decays (spectral diffusion) of SeCN\(^-\) in D\(_2\)O (black points) and in D\(_2\)O confined in the silica pore (red points). The respective simulated CLS decays are shown as dashed lines of the same color.
pore are substantially slower than in bulk $\text{D}_2\text{O}$. The integrated correlation time (eq 4), $\tau_f$, slows from 0.97 ps (bulk) to 2.4 ps (pore) (Table 4). The results indicate that deceleration of water structural relaxation around $\text{SeCN}^-/\text{D}_2\text{O}$ (~2.5-fold, Table 4) is more substantial than deceleration of $\text{SeCN}^-$ reorientation (~1.7-fold, Table 3) between the confined and bulk liquid.

In a manner consistent with the behavior of $\text{SeCN}^-$ reorientation in the silica pore, the time scales of the fast spectral diffusion components, $\tau_f$ and $\tau_2$, are analogous to the bulk spectral diffusion (Table 4). The significant change is the appearance of an additional slow spectral diffusion component in the pore, $\tau_3 = 17 \pm 1$ ps, which is absent in the bulk liquid and more than an order of magnitude longer than any time scale observed in the bulk system. The spectral diffusion comparison is consistent with a picture of dramatically slower H-bonding dynamics near the silica surface and faster dynamics further away, which are still slower than in bulk water.

The measured CLS and linear absorption line shape were obtained to use the frequency–correlation function function (FFCF).$^{39,40}$ The FFCF was modeled with the Kubo ansatz$^{37}$

$$\text{FFCF} = C_n(t) \langle \delta \omega_0(0) \rangle^2 = \langle \delta \omega(t) \delta \omega(0) \rangle = \sum_i \Delta_i^2 \exp[-t/\tau_i]$$

where $C_n(t)$ is the normalized FFCF and the frequency fluctuation, $\delta \omega(t) = \omega(t) - \langle \omega \rangle$, is the difference between the probe’s instantaneous and average frequency, $\langle \omega \rangle$ and $\langle \omega(t) \rangle$, respectively. The ith component of the FFCF is characterized by a frequency fluctuation amplitude, $\Delta_i$, and time constant, $\tau_i$. Components in the homogeneous limit satisfy $\Delta_i \tau_i \ll 1$. In this case, $\Delta_i$ and $\tau_i$ cannot be independently determined. In the homogeneous limit, the Kubo line shape function$^{37,39,40}$ generates a Lorentzian whose width $\Gamma = \Delta \tau/\pi = 1/\tau_i T_2^\tau$ is narrower than the frequency distribution ($\Delta \tau/\pi \ll \Delta$, motionally narrowed) and inversely proportional to the pure dephasing time, $T_2^\tau$. The observed homogeneous dephasing time is more precisely ($T_2^\tau \sim 1 = (T_2^\tau)^{-1} + (T_w)^{-1} + (3T_\text{re})^{-1}$, where $T_1$ and $T_w$ are the vibrational and orientational relaxation times, respectively.$^{39}$ Components with $\Delta \tau_i \gg 1$ are in the inhomogeneous limit.$^{37,39,40}$ In this limit, $\Delta_i$ is the standard deviation of the ith Gaussian component, and the total inhomogeneous line width is given by convolution: fwhm

$$= 2[2 \ln(2)]^{1/2} \Delta_{\text{total}}$$

where $\Delta_{\text{total}} = (\sum \Delta_i^2)^{1/2}$. The convolution of the line shapes associated with each component in eq 7 produces the linear (1D) absorption line shape. The correct FFCF will reproduce the linear absorption line shape and the CLS decay and is confirmed by reproducing the observables through response function calculations$^{37,39,40}$ of the 1D and 2D line shapes. The FFCF parameters obtained from the spectra in Figure 2 and corresponding CLS decays in Figure 9 are provided in Table 5.

The simulated CLS (dashed red curve), obtained from the simulations as previously described,$^{22}$ is compared to the measured results in Figure 9 along with that previously obtained for $\text{SeCN}^-$ in bulk $\text{D}_2\text{O}$ (dashed black curve). The simulations, in agreement with the experimental data, find that the confinement leads to the appearance of an additional long time scale, beyond those observed in bulk solution, that is estimated to be ~60 ps but with significant uncertainty. This time scale is ~3 times longer than that observed in the measurement, indicating that the simulation modeling of the frequency is reasonable but could be improved. However, the frequency map used is linear so that the normalized FFCF obtained is independent of the map parameters and equal to the autocorrelation function of the electric field at the C atom along the $\text{SeCN}^-$ axis. Thus, in this description, the FFCF is determined by the dynamics of those electrostatic fluctuations in the solute environment, even when the frequency is not quantitatively modeled.

The distance dependence of the FFCF, $C_n(T_w)$, is shown in Figure 10A. It shows qualitatively different behavior than that of the orientational correlation function in Figures 5B and 7, namely the FFCFs for distances near the interface do not differ much but are distinct from those obtained when the solute is in the pore interior. This is suggestive of two-state behavior as expressed in eq 5. This is verified by examining the dependence of the FFCF on the position of the $\text{SeCN}^-$ in the pore at short times, which is shown in Figure 10B. Indeed, the distance-dependent FFCF behaves as a (smoothed) Heaviside step function, which can be seen by the agreement with fitting to the function

<table>
<thead>
<tr>
<th>sample</th>
<th>$A_1$ (ps)</th>
<th>$\tau_1$ (ps)</th>
<th>$A_2$</th>
<th>$\tau_2$ (ps)</th>
<th>$A_3$</th>
<th>$\tau_3$ (ps)</th>
<th>$δω$ (ps)</th>
</tr>
</thead>
<tbody>
<tr>
<td>bulk</td>
<td>0.22 ± 0.02</td>
<td>0.1 ± 0.1</td>
<td>0.30 ± 0.04</td>
<td>1.3 ± 0.2</td>
<td>0.97 ± 0.05</td>
<td>2.4 ± 0.3</td>
<td></td>
</tr>
<tr>
<td>pore</td>
<td>0.49 ± 0.05</td>
<td>0.8 ± 0.2</td>
<td>0.19 ± 0.02</td>
<td>3.1 ± 0.3</td>
<td>0.04 ± 0.01</td>
<td>17 ± 1</td>
<td></td>
</tr>
<tr>
<td>simulations bulk</td>
<td>0.27 ± 0.01</td>
<td>0.6 ± 0.1</td>
<td>0.11 ± 0.01</td>
<td>1.4 ± 1.2</td>
<td>6.9 ± 0.2</td>
<td>2.3 ± 0.3</td>
<td></td>
</tr>
<tr>
<td>simulations pore</td>
<td>0.28 ± 0.01</td>
<td>1.7 ± 0.5</td>
<td>0.15 ± 0.01</td>
<td>10 ± 5</td>
<td>0.06 ± 0.01</td>
<td>60 ± 35</td>
<td></td>
</tr>
</tbody>
</table>

“Multieponential fit parameters to the CLS for SeCN$^-$/D$_2$O in bulk solution and confined in the silica pore. The $A_i$ and $\tau_i$ are the amplitude and correlation time constant of the ith component. The $δω_i$ are integrated correlation times (eq 4).

<table>
<thead>
<tr>
<th>sample</th>
<th>$T_2$ (ps)</th>
<th>$Γ$ (cm$^{-1}$)</th>
<th>$Δ_1$ (cm$^{-1}$)</th>
<th>$τ_f$ (ps)</th>
<th>$Δ_2$ (cm$^{-1}$)</th>
<th>$τ_2$ (ps)</th>
<th>$Δ_3$ (cm$^{-1}$)</th>
<th>$τ_3$ (ps)</th>
</tr>
</thead>
<tbody>
<tr>
<td>bulk</td>
<td>1.4 ± 0.1</td>
<td>8.7 ± 0.4</td>
<td>9.4 ± 0.5</td>
<td>0.5 ± 0.1</td>
<td>10.2 ± 0.9</td>
<td>1.3 ± 0.2</td>
<td>3.1 ± 0.3</td>
<td>17 ± 1</td>
</tr>
<tr>
<td>pore</td>
<td>2.3 ± 0.6</td>
<td>5 ± 1</td>
<td>12.2 ± 0.7</td>
<td>0.8 ± 0.2</td>
<td>6.8 ± 0.5</td>
<td>3.1 ± 0.3</td>
<td>3.1 ± 0.3</td>
<td>17 ± 1</td>
</tr>
<tr>
<td>simulations bulk</td>
<td>19.0 ± 0.1</td>
<td>0.089 ± 0.001</td>
<td>13.2 ± 0.1</td>
<td>0.6</td>
<td>6.0 ± 0.1</td>
<td>1.4</td>
<td>5.6 ± 0.1</td>
<td>7.1 ± 0.1</td>
</tr>
<tr>
<td>simulations pore</td>
<td>29.8 ± 0.9</td>
<td>0.157 ± 0.010</td>
<td>16.9 ± 1.0</td>
<td>1.7 ± 0.5</td>
<td>11.6 ± 1.5</td>
<td>10 ± 5</td>
<td>7.1 ± 2.6</td>
<td>60 ± 35</td>
</tr>
</tbody>
</table>

“See section IV.C for parameter definitions.
The simulated dynamics in a model silica pore were compared to the measured TCFs which, in contrast to the simulations, contain no spatial information. Comparisons of the spatially integrated dynamics from the simulations to the measured TCFs informed further adjustments to the pore model, leading to improved agreement with the measured dynamics. A new decay component was observed in both the rotational and spectral diffusion TCFs, \( C_{\omega}(t) \) and \( C_{\omega}(t; d) \), which in each case was significantly slower than the longest time scales in the bulk liquid (~6-fold and ~13-fold slower, respectively). These results indicate that the water H-bond dynamics are considerably perturbed upon confinement in the silica pore. On average, SeCN\(^{-}\) forms slightly fewer H-bonds in the hydrated pore (3.2) than in bulk water (3.4). This appears to be related to the strong affinity of water for the hydroxyl-terminated pore, which results in increased water layering near the surface. In response to this interfacial solvent structuring, SeCN\(^{-}\) tends to avoid the interfacial water \((d < 4 \text{ Å})\) and resides in the more uniform water in the pore interior. However, the retarded solvent dynamics propagate out from the surface into the center of the pore, in turn dramatically slowing reorientation and spectral diffusion of SeCN\(^{-}\) relative to bulk solution. This suggests that the significantly retarded dynamics observed in this system are likely modest compared to that in the interfacial layer, a conclusion that is supported by the distance dependence of the dynamics observed in the simulations. In the future, the interfacial dynamics might be more directly probed in experiments using solutes that possess a greater affinity for the silica surface.

Further analysis of the simulated TCFs revealed an unexpected difference in the evolution of \( C_{\omega}(t) \) and \( C_{\omega}(t; d) \) with distance from the pore surface. The \( C_{\omega}(t) \) dynamics accelerate exponentially with distance from the surface, indicating that the orientational relaxation cannot be separated into distinct “core” and “shell” states for the small pore diameter studied here.\(^{75,74}\) This is consistent with studies on very small water nanopools in reverse micelles.\(^{75,76}\) In contrast, the \( C_{\omega}(t; d) \) dynamics do transition as a (smoothed) step function from slower “shell” dynamics to accelerated “core” dynamics, in agreement with one of the conditions of a two-state model. The point of transition occurs at \( d \approx 6 \text{ Å} \), halfway between the surface and center of the pore. This behavior indicates that the H-bonding degrees of freedom that couple to the vibrational frequency fall off more rapidly with distance than those which influence reorientation. It is important to note, however, that bulk spectral diffusion dynamics do not appear to exist in the “core” region, which is in opposition to a common secondary assumption of two-state models.

V. CONCLUDING REMARKS

The detailed nature of water H-bond dynamics in MCM41 was elucidated by combining measurements and simulations of SeCN\(^{-}\) vibrational spectroscopy in the water-filled ~2.4 nm silica pores. The study of SeCN\(^{-}\) as a vibrational probe in the IR experiments offers several advantages over probing water directly. Foremost among these are the strong transition dipole and long vibrational lifetime of the CN stretching mode, which together permit the observation of complete rotational and spectral diffusion dynamics with high signal-to-noise. The combination of selecting smaller silica particles, index matching the sample, and implementing a new phase-cycling procedure aided in eliminating artifacts due to light scattering from the observables.

Figure 10. (A) Simulated FFCF as a function of the distance, \( d \), between the C atom and nearest pore O atom (dashed curves) compared to the total pore (black solid curve) and bulk solution (black dashed curve) results. (B) Simulated FFCF evaluated at four delay times: 1.2 ps (black circles), 2.4 ps (red circles), 3.6 ps (blue circles), and 4.8 ps (violet circles). Dashed lines of the same color show fits to a smoothed step function (eq 8).

\[
C_{\omega}(t; d) = C_{\omega,\text{core}}(t) + \left[ C_{\omega,\text{shell}}(t) - C_{\omega,\text{core}}(t) \right] \times \frac{\tanh[(\Delta + d_0 - d)/\alpha] + 1}{2}
\]

where \( d_0 \) is taken as 3 Å and \( \alpha \) is a fitting parameter that governs the smoothness of the step function. This qualitative difference in the distance dependence of the FFCF compared to the orientational correlation function is striking. It indicates that the spectral diffusion is governed by relatively local interactions, while SeCN\(^{-}\) reorientation is influenced by more long-range effects. This is consistent with the bulk dynamics,\(^{22}\) where the FFCF is observed to decay on a time scale of 1.3 ps, which is faster than that of 4.5 ps found for reorientation and also more rapid than the established time scale of ~3 ps for H-bond exchanges in water.\(^{72}\) This demonstrates that the FFCF and \( C_{\omega}(t) \) are probing different aspects of the H-bond dynamics.

\( C_{\omega}(t; d) \) dynamics do transition as a (smoothed) step function from slower “shell” dynamics to accelerated “core” dynamics, in agreement with one of the conditions of a two-state model. The point of transition occurs at \( d \approx 6 \text{ Å} \), halfway between the surface and center of the pore. This behavior indicates that the H-bonding degrees of freedom that couple to the vibrational frequency fall off more rapidly with distance than those which influence reorientation. It is important to note, however, that bulk spectral diffusion dynamics do not appear to exist in the “core” region, which is in opposition to a common secondary assumption of two-state models.
In conclusion, this work has shown that water experiences a range of structural and dynamic changes, some subtle and others substantial, upon confinement in nanoscale silica pores. The qualitatively different distance dependence observed for orientational relaxation and spectral diffusion, two properties sensitive to the H-bond dynamics of the confined water, revealed unexpected spatial complexity. Although considerable detail has been illuminated, certain aspects of this system remain incompletely described. In this study, the simulations quantitatively reproduced the measured orientational relaxation dynamics, but the agreement obtained for the spectral diffusion was not as good. Identifying properties of the simulation that lead to the overestimation of inhomogeneous diffusion, two properties sensitive to the H-bond dynamics of the conformation of H-bonds and their dynamics merits further study.

ACKNOWLEDGMENTS

The funders had no role in the study design, data collection, and analysis, decision to publish, or preparation of the manuscript. We thank Jun Nishida for his constructive comments on the manuscript.

REFERENCES


ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acs.jpcc.9b00593.

Experimental details; Figures S1–S4 and Tables S1–S3 (PDF)

AUTHOR INFORMATION

Corresponding Authors

*Phone 650 723-4446; e-mail fayer@stanford.edu.
*Phone 785 864-3980; e-mail wthompson@ku.edu.

ORCID

Ward H. Thompson: 0000-0002-3636-6448
Michael D. Fayer: 0000-0002-0021-1815

Notes

The authors declare no competing financial interest.

ACKNOWLEDGMENTS

This work was funded by the Division of Chemical Sciences, Geosciences, and Biosciences, Office of Basic Energy Sciences of the U.S. Department of Energy through Grant DE-FG03-84ER13251 (S.A.Y., J.Y.S., and M.D.F.) and the NSF-EPA Networks for Sustainable Design and Synthesis (NSMDS) for funding support under NSF Grant CHE-1339661 (W.H.T.). Additional support of the 2D IR instrument and for M.D.F. was provided by Air Force Office of Scientific Research Grant FA9550-16-1-0104. S.A.Y. acknowledges the support from a Stanford Graduate Fellowship. The simulations were performed at the University of Kansas Center for Research Computing (CRC). We thank Jun Nishida for his constructive comments on the manuscript.


