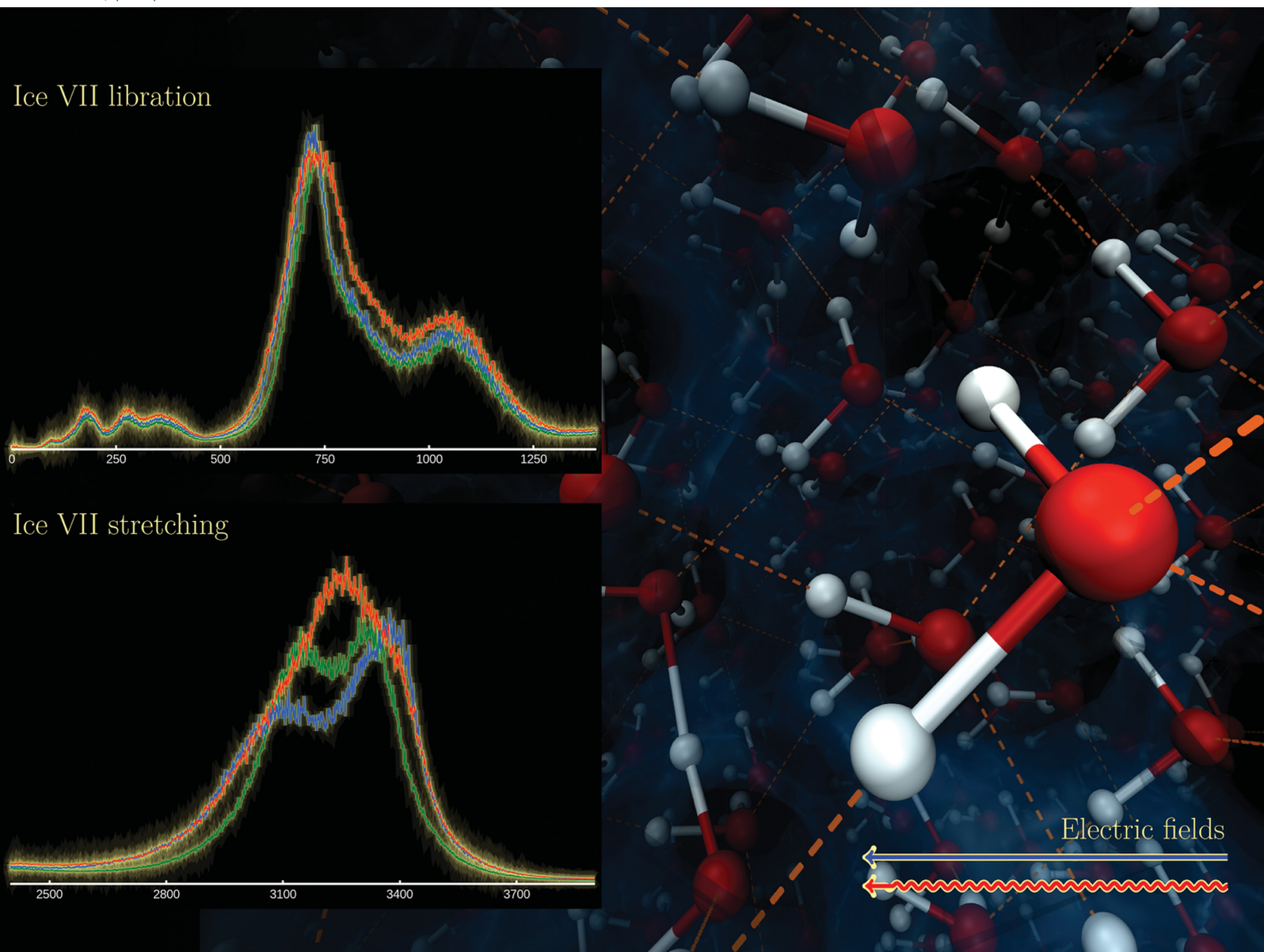


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PAPER

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Dielectric properties of ice VII under the influence of time-alternating external electric fields

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The high-pressure solid phase of water known as ice VII has recently attracted a lot of attention when its presence was detected in large exoplanets, their icy satellites, and even in Earth's mantle. Moreover, a transition of ice VII to the superionic phase can be triggered by external electric fields. Here, we investigate the dielectric responses of ice VII to applied oscillating electric fields of various frequencies employing non-equilibrium *ab initio* molecular dynamics. We focus on the dynamical properties of a dipole-ordered ice VII structure, for which we explored external-field-induced electronic polarisation and the vibrational spectral density of states (VDOS). These analyses are important for the understanding of collective motions in the ice-VII lattice and the electronic properties of this exotic water phase.

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Introduction

Ice VII is one of the 18 currently-known phases of water, which, in contrast to everyday hexagonal ice Ih, crystallises to a cubic body-centered lattice under high pressure, above 2 GPa at room temperature.^{1–3} Although such conditions might seem to be extreme, ice VII plays an important role in the geophysics of solar-system planets. For example, the conditions in mantles of large planets like Uranus and Neptune allow not only for ice-VII formation but also its transition to a conductive superionic phase.^{4–7} High-pressure ice phases could be present on the icy moons of Saturn and Jupiter too.^{8,9} Recently, even the ice VII presence deep in the Earth's mantle has been discovered by its detection in the natural diamond crystal.¹⁰

A good deal of well-warranted attention has been paid to superionic ice phases when water molecules's intact-molecular structure is lost by rapid dissociation of covalent O–H bonds, and the hydrogen protons diffuse freely amongst the crystallographically-placed oxygen atoms, which retain the Wyckhoff lattice positions. The transition of ice VII to a superionic phase is usually initiated by increasing the pressure to critical values where the potential barrier for proton transfer between two neighboring oxygen atoms is low enough to allow symmetric proton exchange to become possible,^{11–13} enhanced by significant nuclear quantum effects.^{14–17} Finally, the protons themselves become fully diffusive upon a further pressure increase as the double-welled O–H···O potential transforms

to a symmetric O···H···O profile, where the individual protons cannot be assigned to particular oxygen atoms anymore.^{18–21} This phenomenon is intimately connected with changes in proton diffusivity and conductivity.^{22–24}

The above-mentioned transitions occur at extreme > 50 GPa pressures and relatively high temperatures. Nevertheless, proton diffusion could be also initiated by externally-applied static electric fields – allowing to lower the critical “proton-hopping-onset” pressure to 10–15 GPa range, as we recently demonstrated by non-equilibrium *ab initio* molecular dynamics (NE-AIMD) simulations.²⁵ Static electric fields are known to align water dipoles, and, eventually, facilitate the breaking of O–H bonds when their intensities reach the electro-dissociation threshold. Saitta *et al.*²⁶ studied these effects on liquid water by DFT-based non-deterministic simulations, establishing this field-threshold intensity of $\sim 0.35 \text{ V \AA}^{-1}$, whilst Casone *et al.*²⁷ showed that in ice Ih and ice XI these limits are considerably lower – slightly above 0.25 V \AA^{-1} and 0.22 V \AA^{-1} , respectively. These external-field intensities, although strong enough to induce molecular-dipole alignment and the consequent covalent-bond breaking, are of the order of $\sim 15\%$ of the intrinsic electric-field strengths in these condensed-matter states of solid/liquid water, which are within the range of 1.5 to 2.5 V \AA^{-1} in these condensed water phases.²⁸ Recently, as mentioned in passing above, we investigated the static-field effect on dipole-ordered ice VII structure under a pressure of 10 GPa and temperature 410 K.²⁵ Under these conditions, the lower boundary of field intensity required to initiate collective proton-hopping processes was found to be 0.33 V \AA^{-1} .

In contrast to external static electric fields, where structural and energetic rearrangement typically occurs more rapidly, the investigation of the effects of time-varying external fields

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requires significantly longer simulation times, as one needs to simulate at least several periodic field oscillations to gain enough data for meaningful statistical analyses.^{28–31} This is usually a limiting factor for first-principles studies where the potential-energy and wavefunction evaluation of reasonably large systems is computationally demanding. However, by leveraging the so-called second-generation CPMD method,^{32,33} which speeds up the computations by lowering convergence thresholds of the energy and forces calculation in Born–Oppenheimer-type dynamics whilst utilising canny extrapolation techniques and Langevin stochastic thermostating, the propagation of relatively large system to sub-nanosecond times become possible. Using rigorous Berry-phase treatment of electric-field effects on systems within periodic boundary conditions, we have demonstrated good performance of this procedure for non-equilibrium *ab initio* molecular dynamics (NE-AIMD) simulation of both static and oscillating electric fields on liquid water,²⁸ clathrate hydrates,³⁴ ice VII,^{25,35} and, very recently, also on fully-solvated interfaces of hematite,³⁶ where we were able to study external-field responses on 0.1-ns time scales.

Here, bearing in mind open questions of the effect of oscillating electric fields, as opposed to static ones,^{25,35} on the behaviour of ice VII, notably vibrational behaviour,³⁵ we study the effects of oscillating fields on ice-VII structure to compare their influence on this hydrogen-bonded, pressurised, condensed phase of water with previously explored static-electric-field responses.^{25,35} A root-mean-square (r.m.s.) field intensity 0.2 V \AA^{-1} was chosen – well below the intramolecular-dissociation limit of static fields (0.33 V \AA^{-1}),²⁵ to focus on the dynamical dielectric responses of structural motions and electronic density. In this way, we investigate by means of auto-correlation functions and by computing and analyzing the vibrational densities of states, the system response to oscillating electric fields, as we discuss further in the text.

Computational details

Density-Functional Theory (DFT) calculations performed within this study were based on the accurate van der Waals density functional (vdW-DF) with the opt-B88 exchange–correlation part^{37,38} and DRSSL nonlocal correlation correction describing the dispersion interactions.^{39,40} This functional was demonstrated to capture well the properties of water, hexagonal ice Ih, weakly-bonded networks of clathrate hydrates as well as the high-pressure structure of ice VII,^{25,28,34,35,41–46} and we have highlighted before the importance of choosing appropriate functionals in condensed-phase systems.⁴⁷ We applied Goedecker–Teter–Hutter (GTH) pseudopotentials,⁴⁸ triple-zeta polarised basis set (TZV2P), and plane-wave cutoff 350 Ry to describe the system potential within the CP2K software,⁴⁹ as justified by our previous calculations of water-based systems.^{25,28,34–36}

The system was propagated by *ab initio* molecular dynamics (AIMD) using the second-generation CPMD technique.^{32,33} This

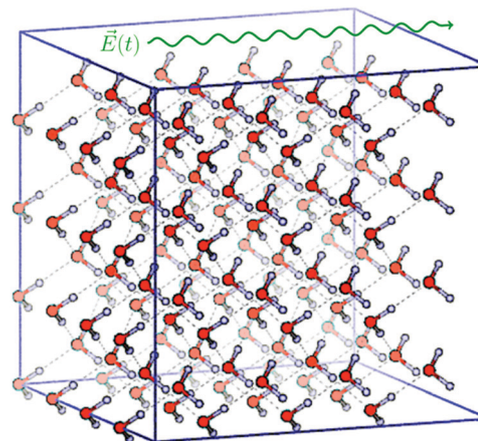


Fig. 1 Structure of proton-ordered ice VII model used in the simulations. The cubic supercell of side length 12.62 \AA contains 128 water molecules. Direction of the applied oscillating electric field is indicated by green arrow.

method allows us to use relatively long integration time steps – 1 fs in our case – and obtain long enough trajectories (100 ps) needed for reliable statistical analyses, bearing in mind the applied-field periods – 50, 150, 250, and 500 GHz frequencies for the NE-AIMD case, with r.m.s. magnitude of 0.2 V \AA^{-1} . A zero-field, equilibrium run was also carried out. Numerical stability of these simulations is ensured by application of ASPC corrector⁵⁰ and Langevin thermostat with stochastic parameter γ set to 0.005. The model was studied in canonical (NVT) ensemble at temperature 410 K and fixed cubic cell dimensions with side length 12.62 \AA accommodating 128 water molecules in $2 \times 2 \times 2$ supercell shown in Fig. 1. This setup corresponds to a pressure of 10 GPa. Although the ice-VII water phase is characterised by hydrogen disorder, the ordered structure with water dipoles aligned to the *x*-direction of the cell was used as an initial structure for the simulations to bring the system into the ferroelectric state that would be induced by applied electric fields, as this was used in our previous NE-AIMD studies of electric-field effects on ice VII.^{25,35}

Results and discussion

Dielectric materials, like water and its ice phases, interact with external electric fields primarily through their constituent molecular dipoles, which tend to align with local electric fields. Macroscopically, this effect is manifested by the polarisation of the system, and the response, within the linear regime, is generally described by the frequency dependent dielectric constant also known as permittivity. At the microscopic level, the dipoles not only rearrange their orientation, at least in part, with the field direction by molecular rotation (so-called orientational polarisation), but their magnitude can also be enhanced due to underlying structural changes (*e.g.*, bond stretches, angle bending, or intra-molecular rotations) and the connected charge-density redistributions (*i.e.*, electronic polarisation). These electronic changes are, in general, much

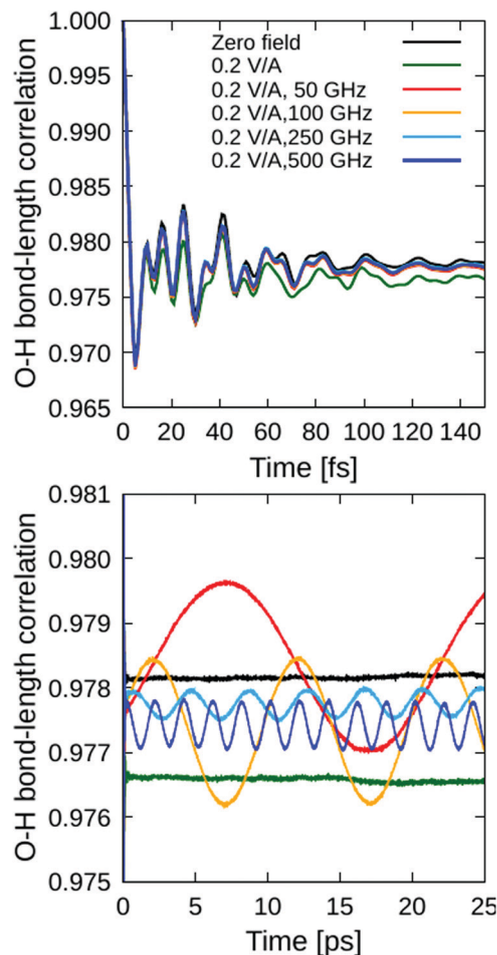


Fig. 2 Autocorrelation function (ACF) of O–H bond stretching vibrations shown at a short-time scale of the first 150 fs (left) and long-timescale of 25 ps (right). Field effects of static as well as alternating applied electric fields are compared with the zero-field ACF curve (plotted in black).

faster than the orientational effects. In the present study, we focus on the electronic-polarisation contributions to ice VII's dielectric response by investigating the effects of homogeneous, time-alternating, harmonic electric fields acting along the direction where the water molecular dipoles have been pre-oriented by electric-field effects.^{25,35}

First, we explored the molecular O–H bond stretching field response by calculating normalised autocorrelation functions (ACFs) of the \vec{R}_{OH} vectors oriented along with these bonds as

$$c(t) = \langle \vec{R}_{\text{OH}}(t_0) \cdot \vec{R}_{\text{OH}}(t_0 + t) \rangle / \langle \vec{R}_{\text{OH}}(t_0) \cdot \vec{R}_{\text{OH}}(t_0) \rangle. \quad (1)$$

where $\langle \dots \rangle$ means ensemble average. The bond ACFs, averaged over 50 000 time origins t_0 along the computed (NE-)AIMD trajectories, are shown in Fig. 2. Because there is no reorientation effect, the water molecules are only allowed to liberate (or “wiggle”) in their equilibrium positions, and even these motions are considerably hindered by a strong hydrogen-bond network typical for this high-pressure ice phase; unsurprisingly, the correlation functions are close to unity, indicating strong correlations.

The fast variations at short times (within 140 ps) that can be seen in the upper panel of Fig. 2 correspond to O–H vibrations, which form the stretching band in the vibrational spectrum spanning approximately 2600 to 3700 cm^{-1} range, as we discuss further below. The fast vibrations are only slightly affected by $0.2 \text{ V \AA}_{\text{RMS}}^{-1}$ oscillating fields with 50, 150, 250, and 500 GHz frequencies, and the corresponding bond ACFs are very similar to the one obtained from zero-field reference simulation. For comparison, we also analysed the effect of a static electric field to the same magnitude (0.2 V \AA^{-1}), which, in stark contrast to the oscillating fields, affects the bond ACF considerably (*cf.* green curve in Fig. 2). This is natural, because static electric fields tend to “stretch” the bonds (with the water oxygen atom carrying a negative partial charge due to electron-density localisation thereat, whilst the hydrogen atoms are electron-deficient and positively charged); this suppresses anti-symmetric vibrations. However, oscillating fields change their polarisation in time, *ipso facto*, and their average effect is small – in particular at higher frequencies when the system

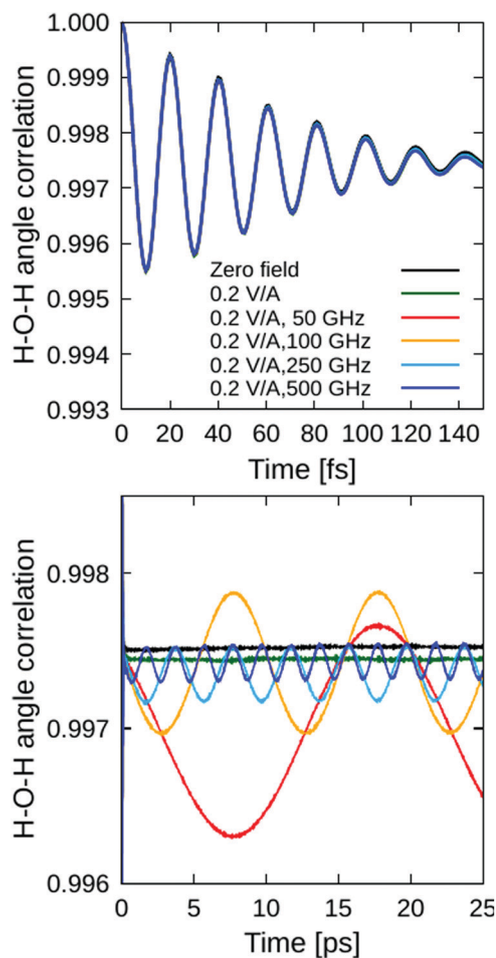


Fig. 3 Autocorrelation function (ACF) of H–O–H valence-angle bending vibrations shown at a short-time scale of the first 150 fs (left) and long-timescale of 25 ps (right). Field effects of static as well as alternating applied electric fields are compared with the zero-field ACF curve (plotted in black).

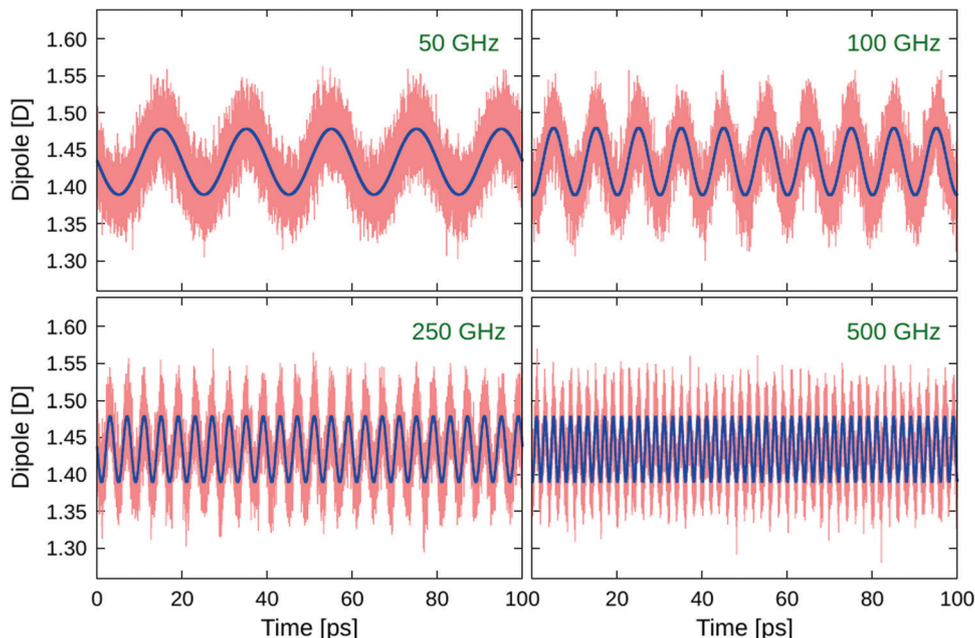


Fig. 4 Response of the mean water dipole (red curve) obtained as the collective dipole divided by the number of water molecules along the 100 ps MD trajectories affected by oscillating electric fields of RMS magnitude 0.20 V \AA^{-1} and frequencies 50, 100, 250, and 500 GHz. The dipole time series are fitted by harmonic functions (blue curves).

has less time to respond. To witness these time-varying effects more clearly, we magnify the long-time tail of the bond ACFs in the lower panel of Fig. 2.

Besides bond stretching *per se*, the applied electric fields affect also the bending motions of the water-valence angles. To detect these effects, we computed the angle autocorrelation function analogous to eqn (1), where, instead of the \vec{R}_{OH} vector, the valence H–O–H angle ϑ_{HOH} was correlated. The calculated ACFs are shown in Fig. 3. Obviously, the external-field effects on the bending motions are much weaker than on the previously-discussed bond stretching. The fast variations in the short-time region (*cf.* upper panel of Fig. 3) are manifested by the vibrational spectral band between *circa* 1300 and 2000 cm^{-1} , and are practically not affected at all by the applied fields of 0.2 V \AA^{-1} in intensity. The field-induced perturbations are detectable only in the long-time tail of the ACFs (lower panel of Fig. 3), where it can be clearly seen that higher frequencies lead to smaller effects on average, as the system does not have enough time to adapt to rapidly-varying field periods; note that 50 GHz corresponds to a field period of 20 ps, whilst the 500 GHz field has a period of only 2 ps.

Although the changes in bond lengths and valence angles are small and their mean values ($R_{\text{OH}} = 1.00 \text{ \AA}$, $\vartheta_{\text{HOH}} = 105.20 \text{ deg}$) are not affected by the applied oscillating fields, there is a considerable field response of the collective dipole moment $\vec{M} = \sum \vec{\mu}_i$, which is the sum of the individual molecular dipoles $\vec{\mu}_i$. The dipoles were calculated semiclassically using Hirshfeld point charges derived from the charge density obtained by DFT. The collective dipole, normalised on the number of water molecules in the studied systems, is tracked in Fig. 4 for all four frequencies of the oscillating electric fields

studied here. The response is clearly harmonic, following the applied $E_0 \cos(\omega t)$ field. For clarity, we fitted the collective-dipole time series by harmonic functions

$$f(t) = A \cos(\omega t) + B \sin(\omega t) \quad (2)$$

which are shown in Fig. 4 as the blue curves on top of the detected red signal. The fitted amplitudes are *ca.* 4 D for all applied frequencies which roughly corresponds to a dielectric constant value of 35. This value is significantly lower than the experimentally reported static permittivity of 112 because we focus on the electronic polarisation here in relatively high-frequency region, and disregard the orientational effects which are contributing to the measured dielectric constant as well. Obviously, the computed value might be also affected by the quality of the applied DFT functional. In the present case, this is known to reproduce well the structural properties of water-based systems, such as radial distribution functions or vibrational spectra; however, admittedly, its accuracy to describe the dielectric response of these materials is as-yet unexplored.

Finally, we analysed the field effects on the vibrational spectra, which characterise collective motions of the water molecules in the ice VII structure. We computed the vibrational density of states (VDOS) by Fourier-transforming the velocity autocorrelation function of the hydrogen atoms

$$I(\omega) \propto \int \langle \vec{v}_{\text{H}}(t_0) \cdot \vec{v}_{\text{H}}(t_0 + t) \rangle e^{-i\omega t} dt. \quad (3)$$

The resulting spectra are shown in Fig. 5 for all applied fields. The three characteristic bands typical for water-based systems are clearly distinguishable – the librational band with the low-frequency modes up to *ca.* 450 cm^{-1} and the dominant peaks in the frequency range from 500 cm^{-1} and 1270 cm^{-1} , bending

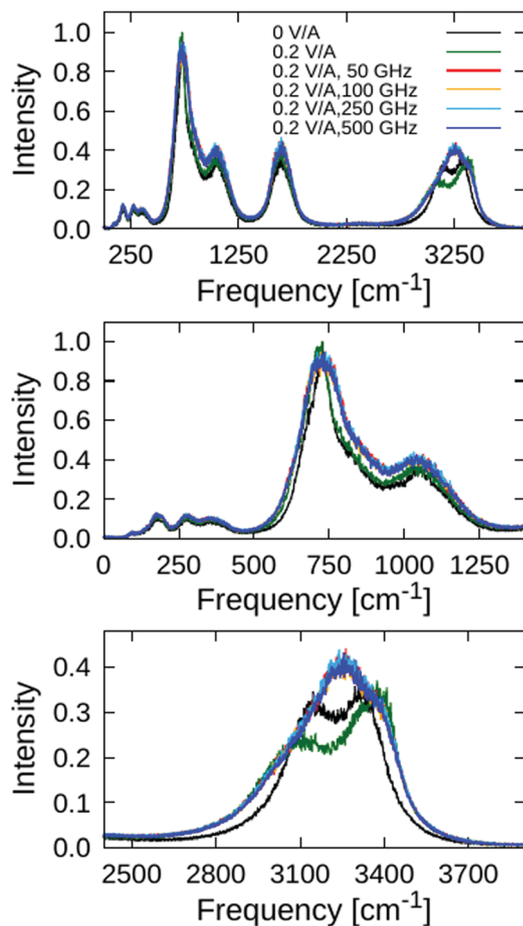


Fig. 5 Vibrational density of states (VDOS) obtained by Fourier transform of the velocity autocorrelation function. Effect of the static and oscillating electric fields of magnitude 0.2 V \AA^{-1} is shown compared to the zero-field spectrum (labeled as 0 V \AA^{-1}). The whole spectrum is plotted in the upper panel while the librational and stretching bands are shown in detail in the middle and lower panels, respectively.

vibrations between *ca.* 1300 cm^{-1} and 2000 cm^{-1} , and the stretching modes spanning spectral region from 2600 cm^{-1} to 3700 cm^{-1} . In general, the spectral features are more affected by the external static electric fields rather than the oscillating ones, comparing the fields of the same intensity 0.2 V \AA^{-1} . Further, the bending vibrations alternating the valence H–O–H angle of the water molecules are practically unaffected by the applied fields, whilst the largest effects are detectable on the spectral parts corresponding to H–O bond stretching vibrations, in accord with the changes in related autocorrelation functions discussed above. External static electric fields promote the asymmetric (ν_3) O–H stretch band and suppresses the symmetric (ν_1) band, which leads to considerable asymmetry in the stretching modes, which is enhanced further by growing the field intensity, as we analysed in ref. 35. On the other hand, oscillating fields tend to “smear” the difference between ν_1 and ν_3 bands, which are merged into one broad peak because the harmonic fields promote both symmetric and asymmetric motions, on average. Changes in the librational band are

visible, although less dramatic. These alternating fields somewhat broaden these spectral peaks and increase their intensities slightly compared to the zero-field reference, as the external-field oscillation promote the molecular vibrations and “wagging” around their equilibrium positions. However, these effects are relatively small in the dipole-oriented ice VII structure, which is well organised and the promotion of the thermal movements is restricted by active thermostating. Finally, we computed the IR spectra directly from dipole-autocorrelation functions

$$I(\omega) \propto \omega^2 \int \langle \vec{M}(t_0) \cdot \vec{M}(t_0 + t) \rangle e^{-i\omega t} dt. \quad (4)$$

However, except for the different intensities of the whole librational and stretching band, which is typical for computed VDOS and IR spectra (see for example Fig. 2 in ref. 35), the observed field effects are consistent with the above discussion. These small effects are expectable and well within the linear-response régime for this system.

Conclusions

We employed non-equilibrium *ab initio* MD based on an accurate correlation-corrected DFT potential and the second-generation CPMD procedure to investigate the dielectric response of dipole-oriented ice VII under pressure 10 GPa and temperature 410 K to oscillating external electric fields of 0.2 V \AA^{-1} RMS intensity and frequencies 50, 100, 250, and 500 GHz. The dynamical responses were analysed by tracking of the collective dipole of the system, computing of autocorrelation functions for H–O bond stretching and H–O–H angle bending, and total VDOS. In contrast to a static electric field of the same intensity, the effects of the time-alternating fields are relatively weak, although nevertheless detectable. Although static fields in general lead to electro-freezing when they have large enough intensities to restrain the water dipoles, and they can even initiate the ice VII transition to the superionic phase by promoting the H–O dissociation, as we showed in ref. 25, the impact of oscillating fields is quite different. The harmonic time variations of the field vector in a given direction promote librational motions and smear the contrasts between the symmetric and anti-symmetric stretching vibrations. There is relatively little difference in response to fields of the presently-explored frequency range, and their effects are clearly distinctive from those induced by the static field of the same intensity, meaning that the 50 GHz can be already characterised as a high-frequency field for ice VII, at least for the presently-gauges properties in terms of system response. Investigation of lower-frequency-field effects towards the lower-frequency end of the microwave spectrum (*e.g.*, 10 GHz and lower) at the DFT level is unfortunately hampered by the need to compute correspondingly longer MD trajectories, close to, and beyond, nanosecond durations. Still, advances in computer hardware mean that deterministic second-generation CPMD will be accessible in the short- to medium-term for such NE-AIMD explorations.

Conflicts of interest

There are no conflicts to declare.

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