Direct Measurement of Competing Quantum Effects on the Kinetic Energy of Heavy Water upon Melting

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ABSTRACT: Even at room temperature, quantum mechanics plays a major role in determining the quantitative behavior of light nuclei, changing significantly the values of physical properties such as the heat capacity. However, other observables appear to be only weakly affected by nuclear quantum effects (NQEs); for instance, the melting temperatures of light and heavy water differ by less than 4 K. Recent theoretical work has attributed this to a competition between intra- and intermolecular NQEs, which can be separated by computing the anisotropy of the quantum kinetic energy tensor. The principal values of this tensor change in opposite directions when ice melts, leading to a very small net quantum mechanical effect on the melting point. This Letter presents the first direct experimental observation of this phenomenon, achieved by measuring the deuterium momentum distributions $n(p)$ in heavy water and ice using deep inelastic neutron scattering (DINS) and resolving their anisotropy. Results from the experiments, supplemented by a theoretical analysis, show that the anisotropy of the quantum kinetic energy tensor can also be captured for heavier atoms such as oxygen. (The iceberg image in the Table of Contents and Abstract graphics was used with permission of the NOAA’s National Ocean Service, 2012 (http://commons.wikimedia.org/wiki/File:Iceberg-_NOAA.jpg.).)

SECTON: Liquids; Chemical and Dynamical Processes in Solution

The structure and dynamics of liquid water are directly influenced by quantum mechanics, not only in terms of the electronic structure and chemical bonding but also at the level of the nuclear motion. So-called nuclear quantum effects (NQEs) include zero-point energy, tunnelling, isotope effects in the thermodynamic properties, and, what is most relevant to the present work, large deviations from the classical, Maxwell–Boltzmann behavior of both the average nuclear kinetic energy $\langle E_N \rangle$ and the momentum distribution $n(p)$.

Even though NQEs are very large (the zero-point energy content of an O–H stretching vibration is in excess of 200 meV), it is often the case that their net effect on macroscopic properties is relatively small. For instance, the melting temperatures of light and heavy water differ by less than 4 K, and the boiling temperatures differ by just 1 K. Recent theoretical analyses have suggested that this could stem from a partial cancellation between quantum effects in the intra- and intermolecular components of the hydrogen bond, so that the net effect is small even if the individual contributions are large. In particular, the competition between quantum effects can be seen very clearly when decomposing the changes in the quantum kinetic energy of protons and deuterons along different molecular axes.

The mechanism that underlies the competition between changes in the different components of the quantum kinetic energy can be understood by considering it as an analogy a two-level quantum system with an environment-dependent off-diagonal coupling $\beta$. A small change in the coupling $\Delta \beta$, arising from a phase transition or some other change in the environment of the system, will shift its eigenvalues by the same amount proportional to $\Delta \beta$, but in opposite directions. Even though this picture is clearly oversimplified, it is consistent with a diabatic state model of the hydrogen bond, it demonstrates that the notion of competing quantum effects is nothing exotic, and explains why it returns in many circumstances in the study of water and other hydrogen-bonded systems.

Competing quantum effects have in fact been identified in a diverse variety of simulations, and it seems entirely plausible...
that they are at the root of the explanation for why many of the properties of water depend only weakly on isotopic composition. As discussed in the Supporting Information (SI), the change of many thermodynamic properties upon isotopic substitution can be related to changes in the quantum kinetic energy. Under a few simplifying assumptions, one can, for instance, relate the change in the quantum kinetic energy of the D atoms when heavy water melts, $\Delta_{\text{h2o}}E_k(m_D, T_{\text{m2}}(m_D))$, to macroscopic thermodynamic properties such as the entropy of melting of light water, $\Delta_{\text{h2o}}S(m_D)$, and the change in the melting temperature upon isotopic substitution

$$\Delta_{\text{h2o}}E_k(m_D, T_{\text{m2}}(m_D)) \approx \frac{\Delta_{\text{h2o}}S(m_D)}{2(m_D/m_H - 1)} [T_{\text{m2}}(m_H) - T_{\text{m2}}(m_D)]$$

Equation 1 predicts a change in kinetic energy per D atom of $\Delta_{\text{h2o}}E_k = -0.5$ meV. This is a tiny value, less than 0.5% of the total kinetic energy of the D atoms at room temperature. A direct experimental investigation of the competing quantum effects at play here should reveal whether this happens because of a cancellation or because the environment of a D atom changes very little upon melting.

Deep inelastic neutron scattering (DINS), or neutron compton scattering (NCS) at high momentum and energy transfers (that is, and $h\omega$), is an experimental technique that is particularly well suited to probe the quantum behavior of atomic nuclei, by directly measuring $n(p)$. The results from DINS experiments have stimulated the development of improved theoretical methods for evaluating the proton momentum distribution, as well as their application to benchmark systems, with a close interplay between theory and experiment. One can infer the anisotropy of the particle momentum distribution from DINS experiments even in cases when only the spherically averaged $n(p)$ is available. In water, this provides insight into the local environment of the proton and can help elucidate the nature of hydrogen bonding, the structure of hydration shells, and the effects of confinement. Indeed this information can be seen as the direct experimental counterpart of the decomposition of the quantum kinetic energy along molecular axes, which has been used so successfully to unravel competing quantum effects in simulations.

The focus of DINS studies has recently broadened to consider also heavier atoms, which, although challenging because of their less-pronounced quantum nature, promise a more comprehensive picture of the underlying physics. Theoretical calculations demonstrate a sizable excess of kinetic energy for the oxygen atoms in ice, relative to the classical value. This kinetic energy excess shows a clear dependence on the chemical environment and on the microscopic structure. A direct, accurate measurement of the kinetic energy of the oxygen atoms could, for instance, shed light on recent findings that indicate an increased localization of the oxygen in heavy water compared to that in light water, as the extension of high-information data on the liquid of the oxygen atoms in heavy water.

At present, the only instrumentation suitable to perform measurements of $n(p)$ in condensed matter systems is the VESUVIO spectrometer, which operates on a dedicated beamline at the pulsed neutron source ISIS (Rutherford Appleton Laboratory, U.K.). The instrument uses neutrons with incident energies in the range of 1–800 eV and relies on the fact that, at sufficiently high momentum transfers, any scattering process can be described within the impulse approximation (IA). This implies that the neutron scatters from a single atom, with conservation of the total kinetic energy and momentum of the neutron and the atom. In the IA regime the inelastic neutron scattering cross section is related in a simple way to $n(p)$. The neutron scattering function $S_N(q, \omega)$ is

$$\frac{h^4}{m} S_N(q, \omega) = \int n(p) \delta(y - p \cdot q) \, dp$$

where $(q, \omega)$ are the wave vector and energy transfers, $m$ is the mass of the atom being struck, $y = (m/h\omega)(\omega - (h\omega/2m))$ is the particle momentum along the q direction, and $I_N(q, \omega)$ is the neutron Compton profile (NCP) (for consistency with previous literature and ease of notation we write the momentum as a wave vector).

When the sample is isotropic, the particle momentum distribution only depends on the modulus of $p$, and the $q$ direction is immaterial; therefore, the NCP is simply $I_N(q, \omega) = 2\pi \delta(\omega - \omega_\text{p}(p)) dp$. This ideal peak profile is broadened by finite-$q$ correction terms $\Delta(y_q)$, as discussed in the SI, by convolution with the instrumental resolution function $R(y_q)$; therefore, the experimental NCP, $F(y_q)$, is

$$F(y, q) = [I_N(y, \omega) + \Delta(y, q)] \otimes R(y, q)$$

One reasonable (and also insightful) way to extract the physical information content from the experimental $F(y_q)$ profile is to assume that the underlying $n(p)$ arises from the spherical average of an anisotropic Gaussian distribution

$$4\pi p^2 n(p) = \int d^3p \exp \left[ \frac{p_x^2}{2\sigma_x^2} - \frac{p_y^2}{2\sigma_y^2} - \frac{p_z^2}{2\sigma_z^2} - \frac{p_x^2 + p_y^2 + p_z^2}{2m} \right]$$

This expression involves three parameters, the variances $\sigma_x^2$, $\sigma_y^2$, and $\sigma_z^2$, which are related to three effective principal frequencies $\omega_\alpha$ by $\sigma_\alpha^2 = \omega_\alpha^2/m$. In the present study, this approach has been used to interpret DINS data acquired on heavy water in the solid at 274 K and in the liquid at 280 and 300 K.

To complement this experimental study, we have also performed some new ab initio computer simulations of heavy water and ice, using the same density functional theory (DFT) framework as that described in ref 13. Tests with different basis sets and the inclusion of dispersion corrections produced no qualitative changes in the results. NQEs were incorporated using the PGLE technique, which combines the path integral formalism with a correlated-noise Langevin equation, thereby enabling fully converged results for room-temperature water to be obtained with as few as six path integral beads.

The conventional way to extract the particle momentum distribution from the path integral formalism involves opening the path and is computationally very demanding. A simpler alternative is to assume that the momentum distribution can be modeled as a multivariate Gaussian as in eq 4 and to use the eigenvalues of the quantum kinetic energy tensor $\hbar^2(p, p_i)/2m$ to estimate the principal components of this distribution. The only difficulty here lies in the fact that in the liquid, the orientations of the water molecules change with time; therefore,
one cannot simply average the centroid virial estimator to obtain the anisotropic kinetic energy tensor.

Here, we compare two different ways around this difficulty. One is to perform a running average of the kinetic energy estimator, the so-called “transient anisotropic Gaussian” (TAG) approximation. For this, we used a triangular averaging window of 100 fs, which has previously been shown to give converged results for light water. Another possibility is to assume that the principal axes of the kinetic energy tensor will have a fixed orientation relative to the molecular geometry. One can then perform a mean-square displacement (MSD) alignment of the instantaneous configuration of each water molecule to a reference structure, rotating the kinetic energy estimator into the molecular reference frame and computing its average and its eigenvalues. We will show that the two approaches give results that are consistent with one another and that they enable a direct comparison with the DINS experiment.

Figure 1 reports examples of the experimental detector-averaged NCP for D and O, $F(y,q)$, together with the best fits obtained with an isotropic and a multivariate Gaussian ansatz for $n(p)$. The angle-averaged $F(y,q)$ is obtained by averaging over the detectors in the range of 32°–66° for D and those between 130° and 163° for O. Data were not symmetrized. This figure provides a graphical representation of the overall quality of both the data and the fit (see also the SI). Clearly, the multivariate Gaussian profile provides a better fit to the experimental data than an isotropic Gaussian.

In the upper panels of Figure 2, we compare the experimental and theoretical $n(p)$’s for liquid D$_2$O at 300 K. There is a near-perfect agreement between theory and experiment in the case of D. The discrepancy is larger in the case of O but comparable with the error bar and much smaller than the deviation from a classical, Maxwell–Boltzmann distribution.

The lower panels show that the discrepancy between theory and experiment is more pronounced when one focuses on the anisotropy of the distribution. The TAG and MSD approximations are consistent with each other, as discussed in more detail in the SI. It is interesting that, despite the noticeable differences in the individual values of $\langle E_n \rangle$, the theoretical and experimental $n(p)$’s for D are almost indistinguishable. However, the theoretical and experimental $n(p)$’s for O, which involve a larger discrepancy in the total $\langle E_n \rangle$ but smaller discrepancies in the individual components, show a more evident difference. Because of the averaging in eq 4, the computed $n(p)$ depends only weakly on how the kinetic energy components are distributed, but in a more pronounced way on the total kinetic energy.

The relative insensitivity of $n(p)$ to the partitioning of $\langle E_n \rangle$ into three principal components justifies the use of either the TAG or the MSD approach to estimate the anisotropy of the kinetic energy tensor. However, this insensitivity also means that extracting the anisotropy from the spherically averaged $n(p)$ is an ill-conditioned problem. For this reason, the analysis of the experimental data typically yields larger relative errors in the individual components of the kinetic energy than in the total.

Bearing this in mind, let us now discuss how DINS can provide a direct verification of the concept of competing quantum effects in water. Table 1 collects all of the present experimental and theoretical results together in a compact form. The agreement between the total deuterium kinetic
energy obtained by DINS and that by simulation is almost perfect. The change in kinetic energy between the liquid at 300 and 280 K is much smaller than the drop in classical thermal energy, which is consistent with the deuteron being almost completely frozen in its vibrational ground state. There is also agreement with previous experiments at T = 292 K within their (much larger) error bar.39

The most interesting results in Table 1 concern the behavior of the momentum distribution in heavy water upon freezing. When going from the liquid to the solid, the DINS data show substantial changes in ⟨E_2⟩_D (associated with motion perpendicular to the plane of the water molecule3,13) and in ⟨E_4⟩_D (associated with motion parallel to the covalent O−H bond). However, the two components change in opposite directions, leading to a much smaller change in the total kinetic energy, which is not statistically significant given the experimental error bars. The increase in ⟨E_2⟩_D is a signature of the more hindered librations in the solid phase, while the decrease of ⟨E_4⟩_D signals a weakening of the covalent bond, which is consistent with the red shift of the stretching peak observed in the vibrational spectroscopy of ice.40 These observations therefore provide a direct experimental verification of the competition between quantum effects resolved along different molecular axes.

Simulations predict the same qualitative effect on the different components of ⟨E_2⟩_D; ⟨E_4⟩_D decreases upon freezing, but ⟨E_2⟩_D increases, leaving almost no change in the total kinetic energy. Performing simulations of the liquid at 280 K and of both the liquid and the solid at 274 K allows us to infer that these effects are due to the phase transition and not the 6 K temperature drop. Note that our simulations show no sign of an increase in quantum kinetic energy upon supercooling, confirming previous theoretical results for light water.41 The present experiments were deliberately well performed into the stable solid and liquid phases of heavy water in order to focus on the experimental signature of competing quantum effects upon melting without interference from the more controversial effects that have been observed in DINS measurements on supercooled water.40,41

While experiment and theory agree on the qualitative observation of a competition between quantum effects upon melting, there are quantitative discrepancies that deserve further comment. For one thing, our DFT results predict ΔE_T ≈ 0, whereas simple thermodynamic arguments predict ΔE_T ≈ −0.5 meV (see eq 1). As discussed in the SI, a simple, empirical water model1 yields predictions that are in agreement with the macroscopic thermodynamic data, which is perhaps unsurprising given that this empirical model accurately describes the change in melting temperature upon isotope substitution.42 While it is remarkable that an ab initio calculation can get so close to the correct result, it is clear that DFT has not yet reached the level of accuracy necessary to obtain a quantitative description of isotope effects. The DINS experiments also have not reached the exquisite level of accuracy that is necessary to discern such a minute change in the total kinetic energy. Currently, the overall sensitivity of DINS measurements allows one to infer values for proton ⟨E_p⟩ within 2 meV uncertainty43 and similar if not higher uncertainty for the heavier masses, D and O. Although changes on the order of 0.5 meV are beyond the current sensitivity of the instrument, we have demonstrated that one can nevertheless gain insight into the competition of effects that leads to a small kinetic energy change by resolving the anisotropy of the kinetic energy tensor. The quantitative differences between DINS and PIGLET on the individual components of the kinetic energy, however, indicate that at present, this insight is only qualitative.

Table 1 also presents the results for the oxygen momentum distribution. While there is good qualitative agreement between theory and experiment, we observe a discrepancy of almost 10% in the total kinetic energy, which may stem from shortcomings of the modeling or from the analysis of the experimental data, which is made harder by the weaker signal given by oxygen and by the partial overlap between the F(νd) peak of the O and that of the Cu can. Nevertheless, the analysis captures even the comparatively weak anisotropy of the oxygen atom kinetic energy, demonstrating how promising it is to extend DINS to heavy atoms. We anticipate that analysis,44 software, and instrument upgrades planned on VESUVIO in the near future will enable a greater precision in the simultaneous measurement of light and heavy atoms, enabling one to access quantitative as well as qualitative information on the particle momentum distribution.

ASSOCIATED CONTENT

Supporting Information

Additional details on the relation between the change in kinetic energy upon melting and macroscopic thermodynamic observables, on the analysis of the experimental data, on the technical aspects of the simulations, and on the results obtained with an empirical force field. This material is available free of charge via the Internet at http://pubs.acs.org.

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