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Local Coordination Environments and Vibrational Dynamics of Protons in Hexagonal and Cubic Sc-Doped BaTiO₃ Proton-Conducting Oxides

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ABSTRACT: The proton local coordination environments and vibrational dynamics associated with the two order of magnitude change in proton conductivity in hydrated forms of hexagonal and cubic structured BaTiₓ₋ₓScₓO₃Hₓ (0.16 < x < 0.7) were investigated using optical spectroscopy, neutron spectroscopy, and first-principles calculations. Whereas the cubic structure compositions display a single proton site, we show that protons occupy three distinct sites in compositions exhibiting the hexagonal structure. The principal site is characterized by interoctahedral hydrogen bonds, while two additional low occupancy sites are similar to those in the cubic structure, with classic intraoctahedral geometry. Furthermore, the proton hydrogen bond strength increases with decreasing scandium doping level. We infer from this that the stronger, more energetic hydrogen bonds in the hexagonal structure, resulting from proton sites with lower symmetry (lower multiplicity), are predominantly responsible for the significant reduction in macroscopic conductivity between cubic and hexagonal BaTi₁₋ₓScₓO₃Hₓ materials, rather than simply the absolute number of protons. Our findings are highly relevant to the field, clarifying the advantages of high-symmetry structures with high-multiplicity proton sites to favorable properties in ceramic proton-conducting oxides.

INTRODUCTION

Proton-conducting oxides find uses in many energy-related applications, including membranes for steam methane reforming,¹² steam electrolysis,¹³ and hydrogen separation,¹ or as electrolyte materials in a solid-state proton-conducting fuel cell (PCFC).⁶−¹⁵ The development of the PCFC technology depends on the discovery of novel materials with high proton conduction in the intermediate temperature regime, T = 200–600 °C. Acceptor-doped perovskites, based on barium zirconate, BaZrO₃, and barium cerate, BaCeO₃, are currently the most studied electrolyte candidates, due to good thermal and chemical stability in addition to high proton conductivity.¹¹−¹⁷ Materials based on titanates, such as acceptor-doped BaTiO₃,¹⁹−²⁶ and Ti-doped Ba₂In₂O₅,²⁵−²⁷ are less well-studied, yet show great promise. Of particular interest to this work is the solid solution of scandium doping into barium titanate,²² BaTi₁₋ₓScₓO₃Hₓ (BTSSX where x is the scandium doping x expressed as a percentage from 0–100), assuming ideal substitution of Sc⁴⁺ for Ti⁴⁺.

The BTSSX solid solution (0 < x < 70) exhibits a structural phase transition from a 6H hexagonal perovskite to a cubic perovskite phase with increasing x. In hexagonal materials, cations distribute over two sites: corner-sharing MO₆ octahedra and face-sharing M₂O₉ polyhedra where M = Sc and Ti. Only corner-sharing MO₆ octahedra occur in the cubic variants.²⁵ In all cases, the oxygen-deficient materials fully hydrate under humid conditions, yielding BaTi₁₋ₓScₓO₃Hₓ, which are good proton conductors, reaching a maximum conductivity of 2 × 10⁻³ S cm⁻¹ at 600 °C for the x = 0.7 composition.²⁴ The proton conductivity in the hexagonal materials is at least 2 orders of magnitude lower with respect to cubic materials. There is also a high activation energy for bulk proton conduction of about 0.77−0.80 eV.²⁵,²⁶

Recent further structural work, using neutron powder diffraction (NPD) on BTS20 and BTS70 compositions,⁶⁷ shows cation ordering in the hexagonal structure composition, with scandium only occupying the corner-sharing MO₆ octahedra, implying an upper limit at X = 33 (x = 0.33) before transition to the cubic structure. No long-range metal order occurs in the cubic structure. In the dehydrated, host materials, oxygen vacancies localize in the shared face of the M₂O₉ polyhedra in the hexagonal structure and show no long-
range order in the cubic structure. Figure 1 shows the hydrated hexagonal and cubic structures from room temperature NPD by Torino et al., which demonstrate significantly different proton sites for the two structural types. Additionally, in situ dehydration of BTS50 and BTS70, which are both cubic, shows significantly different dehydration rates and response of the lattice parameter as a function of temperature during the dehydration process, indicative of differing hydrogen bond strengths in the two materials. The structural investigations suggest that the proton site geometry in the hexagonal phase negatively influences proton mobility, in addition to the reduced number of protons present per formula unit. It also suggests a secondary contribution to the proton mobility related to the total number of protons and their interaction with the oxygen sublattice.

Here, we present a detailed characterization of proton vibrational dynamics, to investigate the different contributions, in four compositions of hydrated BaTi$_{1-x}$Sc$_x$O$_{2+δ}$H$_{x}$ across both structure types, with $x = 0.16$, 0.20 (hexagonal) and $x = 0.50$ and 0.70 (cubic), hereafter referred to as BTS16, BTS20, BTS50, and BTS70 respectively. We report results from Raman spectroscopy, inelastic neutron scattering (INS), and infrared (IR) spectroscopy, which give access to different regions of the vibrational spectra: the lattice dynamics, the hydroxide wag bands, $\delta$(O−H), and the hydroxide stretch bands, $\nu$(O−H). We also report results from first-principles density functional theory (DFT) calculations, based on structures determined by NPD. The purpose of the study is to evaluate the impact of the scandium content and crystal structure, i.e., the influence of the phase transition, on the number and relative occupation of distinct proton sites in BaTi$_{1-x}$Sc$_x$O$_{2+δ}$H$_x$ to obtain insights into the relationship between these properties and the proton conductivity.

**METHODS**

**Sample Preparation.** All samples were prepared using solid-state reaction methods described in detail in references 27 and 28. The starting materials were BaCO$_3$ (Alfa Aesar, 99.8%), TiO$_2$ (Sigma-Aldrich, 99.8%), and Sc$_2$O$_3$ (Alfa Aesar, 99.9%). Final sintering temperatures for the four samples were 1425 °C for BTS16, 1410 °C for BTS20, and 1500 °C for BTS50 and BTS70. After the sintering, the samples were vacuum-dried in a furnace at 500 °C for 96 h, inside a quartz tube, under dynamic vacuum at 1 × 10$^{-6}$ mbar. Sample hydration in a flow-tube furnace assembly used nitrogen (Air Liquide Alphagaz 1, containing 6 ppm of H$_2$O) as the carrier gas, after bubbling through distilled water. See references 27 and 28 for details of the hydration process and precharacterization of the samples by diffraction and thermogravimetric techniques.

**Infrared Spectroscopy.** IR spectroscopy measurements were performed in diffuse reflectance mode using a Thermo Scientific Nicolet 6700 FT-IR spectrometer equipped with a Collector II accessory for diffuse reflection measurements. IR spectra were collected on the four samples (ca. 50 mg each) at room temperature, over the energy range 1200−4000 cm$^{-1}$. The energy resolution over that interval was set to 2 cm$^{-1}$. The spectrum of a rough Au mirror was used as the reference spectrum. Absorbance-like spectra were derived by taking the logarithm of the ratio between the reference and sample spectra. All measurements were performed at room temperature.

**Raman Spectroscopy.** Raman spectra were measured on the four samples at 300 K, using a Renishaw InVia Reflex spectrometer. A 532 nm laser was used as the excitation source and the laser beam was focused on the sample through a ×50 objective lens (NA = 0.5). The power of the laser beam impinging on the sample was approximately 1 mW. The light backscattered from the sample was then collected through the same objective lens and was dispersed by a 2400 l/mm grating prior to the detection by a CCD camera.

**Inelastic Neutron Scattering.** The INS experiments were performed on the IN1 Lagrange spectrometer at the Institut Laue-Langevin, France, and on the TOSCA and MAPS spectrometers at the ISIS Facility, U.K.

The indirect-geometry spectrometer IN1 Lagrange was used to measure about 5 g of the BTS20 and BTS70 samples, respectively, placed in cylindrical hollow aluminum cells. Using a combination of monochromator crystals, the spectra were collected over the energy range $\hbar \omega$ = 0−500 meV, and with increased statistics with the Cu(220) monochromator over the extended $\delta$(O−H) mode region, $\hbar \omega$ = 50−200 meV. The indirect-geometry spectrometer TOSCA was used to complement the data set by measuring over the extended $\delta$(O−H) mode region the BTS16 (5.9 g), BTS20 (5.0 g), BTS50 (2.1 g) and BTS70 (5.2 g) samples, which were placed in flat aluminum cells and sealed with indium wires. The indirect-geometry spectrometer MAPS was used to access the extended $\nu$(O−H) mode region, $\hbar \omega$ = 300−600 meV, for the four samples and with the same sample cells used on TOSCA. Spectra were collected with incident neutron energies of $E_i$ = 650 meV (600 Hz) and $E_i$ = 300 meV (400 Hz), respectively,
where the number in parentheses refers to the Fermi chopper frequency. All the measurements were performed at $T < 20$ K. Reduction and standard corrections of the data were performed using the LAMP$^{29}$ software for the IN1 Lagrange data, and the MANTID$^{30}$ software for the TOSCA and MAPS data.

Computational Details. DFT calculations were performed using the projector augmented wave (PAW) method$^{31,32}$ as implemented in the VASP$^{33,34}$ software together with the PBE$^{35}$ exchange-correlation functional. For the cubic BaTiO$_3$, a supercell containing 40 atoms constructed from a 2 $\times$ 2 $\times$ 2 stacking of primitive unit cells was used in most calculations. We denote this the defect cell. The introduction of a proton in the defect cell implies a defect concentration of 12.5%. The same defect cell was used for the calculation of the combined defect of a proton and a scandium atom in various configurations. For hexagonal BaTiO$_3$, the unit cell containing 30 atoms was used as the defect cell implying a defect concentration of 16.7%. For the defect cell a Monkhorst–Pack $k$-point mesh of 6 $\times$ 6 $\times$ 6 for the cubic and a Γ-centered 6 $\times$ 6 $\times$ 2 for the hexagonal cell were used together with a plane wave cutoff energy of 700 eV. The calculations were converged to energies within $10^{-7}$ eV for the electronic structure and to forces within $10^{-3}$ eV/Å for the ionic relaxations. The eigenmodes were obtained using the frozen phonon method in the harmonic approximation as implemented in phonopy. $^{36}$ Calculations of the force constants of the dynamical matrix were conducted in the defect cell.

RESULTS AND DISCUSSION

Infrared Spectroscopy. The IR spectra of the four samples are shown in Figure 2. The main feature of the spectra is the band in the region 350–450 meV, characteristic of the $\nu$(O–H) modes in hydrated oxides and particularly in hydrated acceptor-doped cubic perovskite oxides.$^{37-40}$ The overall downward shift of the band in comparison with the free O–H molecular ion $\nu$(O–H) energy, at about 450 meV,$^{41}$ is attributed to the presence of hydrogen bonds.$^{42}$ The large width of the band is then rationalized by a distribution in strength of the hydrogen bonds, underpinning the coexistence of numerous proton local environments. The asymmetry of the band toward lower energy may indicate the presence of a distinct second population of proton sites, or it may be an effect of the increased IR cross-section of the increasingly polarized O–H unit when its hydrogen bond is strengthened. The main $\nu$(O–H) band is centered at about 420 meV for BTS50 and BTS70, 410 meV for BTS20, and 405 meV for BTS16, which denotes an overall strengthening of the hydrogen bonds in the hexagonal phases.

The spectra of the BTS16 and BTS20 samples show a more defined $\nu$(O–H) band than the cubic counterparts. They also contain minor peaks at about 270 and 418 meV (triangle up and down in Figure 2). This indicates that, while the spread of hydrogen bond environments may be lower in the hexagonal phases with respect to the cubic phases, additional distinct proton sites are present.

Raman Spectroscopy. The Raman spectra of the four samples are shown in Figure 3. The presence of bands in the

Raman spectra of cubic BTS50 and BTS70, which should be forbidden based on the group theory analysis of the crystal structure with perfect cubic symmetry ($Pm\bar{3}m$ space group), is rationalized by symmetry breaking at the local scale induced by the occupational disorder between Sc and Ti over the B-site of the ABO$_3$ perovskite structure, and by the presence of oxygen vacancies. This local disorder effect on Raman modes is also commonly observed for the cubic phase of undoped BaTiO$_3$ (BTO) ($Pm\bar{3}m$ space group), which is usually transformed from the tetragonal phase ($P4mm$) at ambient pressure and room temperature after applying high pressures ($\sim$2–3.5 GPa) or temperatures ($\sim$395–405 K).$^{44-46}$ The energies of the modes are extracted by peak fit analysis of the Raman spectra with Gaussian functions, and reported in Table 1.

Figure 2. IR spectra of BTS70 (blue), BTS50 (purple), BTS20 (orange), and BTS16 (yellow). The 276–300 meV region of the spectra are omitted due to a strong absorption of the Au reference; interpolated dashed lines are instead shown as guide for the eyes. The peak at ca. 180 meV (squares) is assigned to a carbonated impurity.$^{43}$ The spectra have been corrected by subtraction of a linear background and normalization to the integral of the $\nu$(O–H) band (350–450 meV) and are vertically offset for clarity.

Figure 3. Raman spectra of BTS70 (blue), BTS50 (purple), BTS20 (orange), and BTS16 (yellow), over the lattice mode region. The spectra have been corrected by the subtraction of a linear background, have been normalized to unity, and are vertically offset for clarity.
Overall, we observe minor differences between the Raman spectra of the two cubic samples, BTS50 and BTS70. The double peak at 35–45 meV in BTS70 appears as a single peak at 40 meV in BTS50. The highest-energy band at ~95 meV appears as a narrower band with a single peak in BTS70, but as a wider doublet band in BTS50. These differences in the band shapes and positions may arise from the changes in the local B–O bond lengths and/or angles of BO₆ octahedral units as a result of increasing the concentration of the larger Sc³⁺ dopant, whose ionic radius is about 23% larger than the Ti⁴⁺ one. Note that the spectral differences may be less likely related to the mass effect since the masses of Sc³⁺ (45 u) and Ti⁴⁺ (48 u) are comparable. Nonetheless, the Raman spectra of BTS50 and BTS70 are in general similar, which indicates that the local structures and associated lattice dynamics of these materials are almost the same. Interestingly, the Raman spectra of cubic BTS50/BTS70 and cubic BTO exhibit some similarities, as characterized by two main groups of bands in the spectral ranges of <50 and ~50–75 meV, respectively, except for the absence of bands at ~75 meV in cubic BTO.44–46

The Raman spectra of the two hexagonal samples, BTS16 and BTS20, are also similar to each other, except for the ~2 meV downward shift of the two bands at 75 meV and at 96 meV on increasing the Sc concentration. Specifically, the band at 74.2 meV for BTS20 (76.5 meV for BTS16) is one of the key Raman mode features for hexagonal perovskites oxides.47–49 Moreover, the Raman spectra of BTS16 and BTS20 in the spectral range of <90 meV are very similar to the Raman spectrum of hexagonal BTO, which is mainly characterized by three strong bands at 19.5, 27.9, and 79.7 meV.49

In summary, both cubic and hexagonal phases of Sc-doped BTO show an additional band feature in the high energy regions (>75 and >90 meV) for the cubic and hexagonal phases, respectively, in comparison with the corresponding phases of undoped BTO, which suggests that the local disorder specifically due to the Sc doping effect induces Raman-active modes. Regardless of different crystal structures, this high energy band exhibits a general energy downward shift upon increasing the Sc concentration from BTS16 to BTS70. This implies that these high-energy modes in the four samples studied may originate from the same type of local disorder that is virtually independent of the structural phases but it is primarily influenced by the amount of Sc dopant.

Indeed, qualitatively similar spectra are recorded in the system BaZrₓTi₁₋ₓO₃ with 0.5 ≤ x ≤ 1.0, where the B-site is shared by Zr⁴⁺, similar in size to Sc³⁺ (72 and 74 pm, respectively, when hexacoordinated).50,51 An intense band at ~90 meV is observed and discussed in the context of the cubic to tetragonal phase transition, where the symmetric Ti–O–Ti unit in the cubic phase becomes asymmetric in the tetragonal phase, Ti–O–Ti, with longer and shorter Ti–O bonds. This ~90 meV band is assigned to an A₁(LO) mode, the stretch of the shorter Ti–O bonds. It is also observed in orthorhombic BaTiO₃, but with a weaker intensity.52 In BTS50 and BTS70, the occupational disorder between Sc and Ti over the B-site leads necessarily to a number of asymmetric B–O–B units and, thus, a distribution of B–O bond lengths.

This effect is also present for BTS16 and BTS20 to a lesser degree, as only the (2a) B-sites are concerned by the occupational disorder. By analogy, we thus tentatively assign the ~95 meV band, present in the four samples, to A₁(LO) stretch modes of short B–O bonds. The fwhm of the ~95 meV band would then be related to the spread of the B–O bond length distribution. The larger fwhm of the ~95 meV band in

Table 1: Mode Energies, in meV, Extracted by Peak Fit Analysis of the Raman Spectra

|       | BTS70  | BTS50  | BTS20  | BTS16  |
|-------|--------|--------|--------|--------|
| 1     | 19.8(1) | 21.5(4) | 21.3(2) | –      |
| 2     | 24.7(3) | 28.3(5) | 28.7(2) | 27.4(3) |
| 3     | 35.4(1) | 38.7(3) | 37.8(2) | 37.5(3) |
| 4     | 43.8(1) | 45.8(4) | 44.5(5) | 45.0(2) |
| 5     | –      | –      | 47(1)  | 47.4(2) |
| 6     | 53.7(2) | 54.4(1) | 52.0(6) | 52.6(1) |
| 7     | 63.9(2) | 63.3(1) | 63.2(1) | 64.3(1) |
| 8     | 72.7(2) | 71.3(1) | 74.2(1) | 76.5(1) |
| 9     | 85.3(4) | 88.7(4) | 86.5(3) | 86.9(2) |
| 10    | 94.5(2) | 96.2(3) | 94.9(3) | 96.9(7) |
| 11    | –      | 100.8(4) | 102.0(2) | 102.7(2) |
| 12    | 114.0(3) | –      | 111.1(2) | –      |

*Standard error is indicated in parentheses. Main peaks are indicated with a single prime.*

Figure 4. (a) IN1 Lagrange data. $S(Q, \omega_B)$ of BTS70 (blue) and BTS20 (red). (b) TOSCA data. $S(Q, \omega_B)$ of BTS70 (blue), BTS50 (purple), BTS20 (red), and BTS16 (yellow) over the δ(O–H) mode region. Spectra are offset for clarity. (c) MAPS data. $S(\omega_B)$ of the four samples over the δ(O–H) mode region (same color code as for part b), obtained by integration of $S(Q, \omega_B)$ over the interval $Q = [0–12] \text{Å}^{-1}$. Spectra are vertically offset for clarity.
BTS50 than in BTS70 is then consistent with the higher expected disorder in BTS50. The relatively lower intensity of the ∼95 meV band in BTS16 and BTS20 than in the cubic phases may then be due to the reduced number of disordered B-sites. Furthermore, an additional band at ∼94 meV is also observed in BaZrTi1−xO3 for Ti-rich compositions (x ≤ 0.6), and is interpreted as a signature of the relaxor phase. This may correspond in BTSX to the band observed at 100.8, 102.0, and 102.7 meV for BTS50, BTS20, and BTS16, respectively.

Inelastic Neutron Scattering. The S(Q, hω) of the BTS20 and BTS70 samples, measured on IN1 Lagrange, are shown in Figure 4a. The spectrum of the BTS70 sample shows strong similarities to other hydrated acceptor-doped cubic perovskite oxides. It is characterized by three broad bands at 95−120 meV (A), 190−240 meV (B), and 380−440 meV (C), assigned, by comparison to said isostructural materials, to the fundamental δ(O−H) modes, overtone of the δ(O−H) modes, and fundamental υ(O−H) modes, respectively. The broad nature of the bands is attributed to a wide range of compounds to inhomogeneous broadening caused by a distribution in the strength of the hydrogen bonds. By comparison, the spectrum of the hexagonal BTS20 sample shows several well-defined peaks over the δ(O−H) region (Figure 4b), at hω = 98, 104, 118, 128, and 137 meV, which suggest the presence of at least three distinct proton environments. Note the overall blue-shift of the δ(O−H) band, and associated red-shift of the υ(O−H) band, indicating in overall stronger hydrogen bonds in the hexagonal than in the cubic phase.

The S(hω) over the δ(O−H) mode region of the four samples, measured on TOSCA, are shown in Figure 4b. The two cubic samples, BTS50 and BTS70, have similar spectra, showing a broad band centered at about hω = 110 meV, discussed previously. There is no argument, based on the structure or the spectra, which suggests that the nature of the proton sites may be different between these two materials. The spectra of the two hexagonal samples, BTS16 and BTS20, are also similar overall, showing distinct peaks. The major difference relates to the relative intensity of these peaks, associated with the relative occupation of the proton sites. The increase of the high-energy δ(O−H) peak, at about 138 meV, indicates a higher population of strongly hydrogen-bonded protons in BTS16 than in BTS20.

The S(hω) over the extended υ(O−H) modes region of the four samples, measured on MAPS, are shown in Figure 4c. The spectra are obtained by integration of the S(Q, hω) data over the interval Q = [0−12] Å−1. The broad bands in the interval 350−450 meV are tentatively assigned to the fundamental υ(O−H) modes, and the bands at hω > 500 meV to combinations between fundamental wag and stretch modes (multiphonon contribution). The fundamental υ(O−H) modes are found at about 420 meV for BTS50 and BTS70, and 405 and 400 meV for BTS20 and BTS16, respectively, which confirms the overall stronger hydrogen bonds in the hexagonal phases. It is also consistent with the greater amount of strongly hydrogen-bonded protons in BTS16 than in BTS20. Note the presence of a low energy shoulder on the υ(O−H) band of BTS50 at about 380 meV, which may indicate that additional proton sites are present in BTS50 with respect to BTS70.

First-Principles Calculations. We have performed first-principles lattice dynamics calculations of pristine BaTiO3 with a cubic and hexagonal structure. The phonon eigenvectors at the Γ-point for the two structures are reported in Tables S1 and S2 in the Supporting Information, respectively.

The energies of the δ(O−H) and υ(O−H) bands are calculated in the relaxed structure of cubic BaTiO3 where a single proton, OH−, is added to the vicinity of one of the (formerly equivalent) oxygen atoms. The hydroxide band energies are calculated at 71, 116, and 455 meV, which correspond, with respect to the Ti−O−Ti plane, to the in-plane wag motion (δ(O−H)∥), out-of-plane wag motion (δ(O−H)⊥), and stretch motion (υ(O−H)), respectively. Considering the effect of Sc, we calculated several structures containing both OH− and ScO−. The most stable structure is found with the scandium and the proton in close proximity. Note that the configuration with Sc and OH as second nearest neighbor is also rather likely, with an energy only 42 meV higher. The hydroxide band energies are calculated at 71, 113, and 452 meV, which suggests that the proton vibrational dynamics are largely unaffected by the presence of Sc.

For the hexagonal phases, several structures with distinct proton configurations are calculated (see Figure 5). The geometry of the hydrogen bonds in the three most stable structures, their calculated vibrational energies, and the energy difference from the most stable structure, are reported in Table 2. The H(1) configuration is found the most stable, and is characterized by an interoctahedral hydrogen bond with a hydrogen bond angle close to 180º, in agreement with our previous work.27 Despite the relatively long hydrogen bond length, and because of its linear geometry, the calculated vibrational energies indicate that the hydrogen bond strength is superior to that of the cubic compound. The H(2) and H(3) configurations are associated with a hydrogen bond geometry comparable to the proton configuration in the cubic structure, on the edge of the TiO6 octahedra, yet also with stronger hydrogen bonds. Other structures, such as the one marked H(4) in Figure 5, involving hydrogen bond patterns on a face

![Figure 5](https://dx.doi.org/10.1021/acs.jpcc.0c01705)
Table 2. Geometry of the Covalent Bonds (CB) and Hydrogen Bonds (HB) and Proton Vibrational Energies for the Three Most Stable Proton Configurations in the Calculation of Hexagonal BaTiO$_3$.

|            | H(1) | H(2) | H(3) |
|------------|------|------|------|
| CB length [Å] | 0.98 | 1.03 | 1.02 |
| HB length [Å] | 2.04 | 1.94 | 1.90 |
| HB angle [deg] | 178  | 140  | 145  |
| $\delta$(O−H) [meV] | 94   | 114  | 118  |
| $\delta$(O−H)$_\parallel$ [meV] | 134  | 130  | 125  |
| $\nu$(O−H) [meV] | 427  | 409  | 411  |
| $\Delta\varepsilon$ [meV] | 0    | 317  | 355  |

The difference in energy from the most stable configuration is also indicated.

Table 3 shows the vibrational energies observed in the INS spectra of cubic BTS70 and hexagonal BTS16 and the calculated $\delta$(O−H) and $\nu$(O−H) bands for the cubic and hexagonal defect cells. Overall, we observe a good agreement between the experimental and calculated proton vibrational energies for the $\delta$(O−H) bands, while the $\nu$(O−H) energies are overestimated in the calculations, especially for the cubic structure. Note that the structural model considered here is BaTiO$_3$, i.e. scandium-free (or a single-scandium atom for the cubic structure), which leads to pronounced disparities in terms of cell parameters between calculation and experiment, which could explain the differences in proton vibrational energies. Nonetheless, the tentative association of the experimental $\delta$(O−H) vibrational energies of BTS16 with the calculated ones, shown in Table 3, leads us to a description of the hexagonal BTEX where at least three distinct proton configurations may coexist: the H(1) configuration characteristic of the hexagonal structure, and the H(2) and H(3) configurations remnant of the cubic structure. Comparing the spectra of hexagonal BTS16 and BTS20, we observe that the band at 139 meV, assigned to the H(1) $\delta$(O−H)$_\parallel$ mode, has increased intensity when the content of Sc is lower, i.e., when the material is further away from the phase transition, which supports our interpretation.

Regarding the effect of the scandium content across the entire substitution range, we observe with increasing Sc content an overall increase of the $\nu$(O−H) band energy, i.e., a decrease in hydrogen bond strength. This effect is likely related to the large difference in ionic radius between Ti$^{4+}$ and Sc$^{3+}$, 60 and 74 pm, respectively. As scandium substitutes preferentially in the (2a) sites (cf. Figure 5), and since we have calculated that OH$^+_n$ and Sc$^{3+}$ attract each other, we propose the following structural description. In the hexagonal phases, increasing the scandium content leads to an increase of the cubic-like proton environments, and a decrease of their hydrogen bond strength, until the phase transition to the cubic structure. In the cubic structure, remnants of hexagonal-type hydrogen bond configurations may still be present in Ti clusters (see the low-energy $\nu$(O−H) shoulder of BTS50 on Figure 4c), which disappear with increasing Sc content. As the Ti−OH−Ti/Sc pattern becomes less likely, the hydrogen bond strength stabilizes at moderate-

![Figure 6](image-url)
to high Sc contents. Considering the superior bulk proton conducting properties of the cubic phases with respect to the hexagonal phases, by about 2 orders of magnitude,25 we infer that the presence of hexagonal-type proton-configurations, originating from Ti-rich domains, is unfavorable for the proton-conducting properties.

This result thus clearly highlights the benefits of high-symmetry structures with high-multiplicity sites to accommodate high proton conductivity in BaTi$_{1-x}$Sc$_x$O$_3$H$_x$ and, by extension, to other related proton-conducting oxides.

**CONCLUSIONS**

We have shown that the main factor impacting the local coordination environment and vibrational dynamics of protons in hydrated BaTi$_{1-x}$Sc$_x$O$_3$H$_x$ is the crystal structure, as protons occupy a single site in the cubic phase ($x=0.5$ and 0.7) and three distinct sites in the hexagonal phase ($x=0.16$ and 0.2). The main proton site in the hexagonal phase directly originates from the hexagonal arrangement and is characterized by interoctahedral hydrogen bonds, while the two additional sites are remnants of the cubic structure, with classic intraoctahedral geometries. The impact of the scandium content, besides on the absolute number of protons, is predominantly due to the cubic-to-hexagonal phase transition, rather than being due to the absolute number of protons, is predominantly due to the presence of stronger, more energetic hydrogen bonds in the hexagonal structure, which also involves proton sites with lower symmetry (lower multiplicity) than in the cubic structure. This insight showcases the favorable role of high-symmetry crystal structures and high-multiplicity proton sites to promote high proton conductivity in Sc-doped BaTiO$_3$ and, by extension, to other proton-conducting oxides.

**ASSOCIATED CONTENT**

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acs.jpcc.0c01705.

Eigenvectors from the lattice dynamics calculations of cubic and hexagonal BaTiO$_3$ (PDF)

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