



Cite this: *Phys. Chem. Chem. Phys.*,
2025, 27, 13601

Proton delocalization in short hydrogen bonds assembling HSeO_4^- anions into supramolecular adducts†

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Four hydrogen-bonded complexes of selenic acid with N-heterocycles (pyridine, 4,4'-bipyridil, quinoline and 2,2,6,6-tetramethylpiperidine) were studied in the crystalline state by single crystal X-ray diffraction, Fourier-transform infrared spectroscopy, and density functional theory with periodic boundary conditions. In all cases short $\text{SeO}-\text{H}\cdots\text{OSe}$ hydrogen bonds (≤ 2.61 Å) were found, either 'isolated' ones or within infinite chains. The coherent quasi-adiabatic proton transfer pathways were computed providing broad asymmetric single-well or (low-barrier) double-well potentials with significantly delocalized protons. The ground state vibrationally-averaged proton positions are noticeably shifted from the equilibrium ones towards hydrogen bond centres, and for asymmetric low-barrier double wells, a proton transfer in the first vibrationally excited state is established. The computed O–H stretching frequencies lie in the range of 1600–3050 cm^{-1} and are in semi-quantitative agreement with experiment. Moreover, in case of low-barrier double-well potentials, a rather exotic H/D isotope effect, namely, a higher vibrational frequency for the O–D stretching than for the O–H stretching, is predicted. The mutual influence of neighboring $\text{SeO}-\text{H}\cdots\text{OSe}$ and $\text{SeO}\cdots\text{H}-\text{N}$ bonds affecting the bridging proton position in both bonds has also been addressed, indicating the vivid geometric cooperativity in complexes with 'isolated' $\text{SeO}-\text{H}\cdots\text{OSe}$ hydrogen bonds.

Received 28th March 2025,
Accepted 6th June 2025

DOI: 10.1039/d5cp01211b

rsc.li/pccp

Introduction

Inorganic and hybrid organic–inorganic hydrogen sulphates (HSO_4^-) and hydrogen selenates (HSeO_4^-) salts are studied for their proton conductivity or, when they form non-centrosymmetric crystals, for their nonlinear optical properties.¹ The phenomenon of proton conductivity is efficient in the presence of an organized hydrogen bond (H-bond) pattern that sometimes assembles two or more anions into supramolecular adducts. For hydrogen selenates, the proton conductivity is facilitated by the formation of H-bonds between neighbouring selenate moieties assembled into dimers or infinite chains. For instance, it was established that in the dimeric selenates of alkali metals, the superprotonic conductivity at

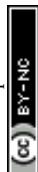
relatively low temperatures involves proton transfer along the $\text{SeO}-\text{H}\cdots\text{OSe}$ bond as well as proton hops between neighbouring selenate dimers over the dynamically disordered network of H-bonds.² Both processes are believed to occur *via* phonon-assisted proton tunneling.^{3–7}

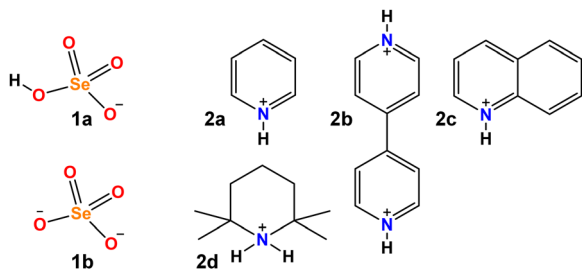
Anion $\cdots$ anion self-assembly is frequently directed by H-bond,⁸ and this is typically the case for HSO_4^- , H_2PO_4^- , and HCO_3^- anions. Recently, it has been described that also σ -hole interactions can drive anion $\cdots$ anion self-assembly,⁹ and even hybrid motifs have been reported where both H-bonds and σ -hole bonds contribute to the formation of anion $\cdots$ anion supramolecular adducts.¹⁰ In the Cambridge Structural Database, a non-minor fraction of the structures containing the HSeO_4^- anion shows the presence of anion $\cdots$ anion adducts formed under H-bond control. For instance, $\text{SeO}-\text{H}\cdots\text{OSe}$ bonds assemble HSeO_4^- anions into supramolecular infinite chains in guanidinium,¹¹ 2-amino-5-chloropyridinium,¹² 2-amino-3-nitropyridinium,¹³ 4-benzylpyridinium,¹⁴ 1,2,4-triazolium¹⁵ and imidazolium¹⁶ hydrogen selenates, as well as in divalent metal hydrogen selenates $\text{M}(\text{HSeO}_4)_2$ ($\text{M} = \text{Mg}, \text{Zn}, \text{Mn}$).¹⁷ Anion $\cdots$ anion dimers are instead formed by (ureido)guanidinium,¹⁸ di-*n*-butyl-ammonium¹⁹ and tetra-*n*-alkyl-ammonium¹ hydrogen selenates, as well as by alkali metal hydrogen selenates encapsulated within the cavity of 18-crown-6 ether: $[\text{18-crown-6}\cdot\text{M}]\text{HSeO}_4\cdot\text{H}_2\text{O}$ ($\text{M} = \text{Rb}, \text{Cs}$).²⁰

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† Electronic supplementary information (ESI) available: Data on experimental and computed crystal structures; QTAIM analyses, calculated potentials and geometries of hydrogen bonds; computed IR features. CCDC 2417665, 2417668, 2417669, 2417697, 2417671, 2417672 and 2417674. For ESI and crystallographic data in CIF or other electronic format see DOI: <https://doi.org/10.1039/d5cp01211b>





Scheme 1 Structural formulas of components in obtained crystals.

While the number of structures containing the HSeO_4^- anion is rather small to draw statistically reliable conclusions, it can be surmised that the HSeO_4^- anion is prone to self-assemble under H-bond control.

The resulting H-bond network plays a key role in the proton conductivity of such materials. In turn, the proton conductivity depends on H-bond strength, on the mutual coupling of neighbouring H-bonds, on the degree of proton delocalization in each H-bond and on the other factors that might alter the proton position. A detailed insight into any and all of these factors would afford useful information for the rational design and functional optimization of the materials containing the $\text{HSeO}_4^- \cdots \text{HSeO}_4^-$ supramolecular synthon. In this paper, we aimed to focus on the degree of proton delocalization within the $\text{O}-\text{H} \cdots \text{O}$ bond and the resulting facilitation of proton transfer along it. The study was started by preparing four hybrid organic–inorganic hydrogen selenates **1a-2a**, **1a-2b**, **1a-1b-2c** and **1a-2d** (see Scheme 1). Single crystal X-ray analyses showed the presence of the $\text{SeO}-\text{H} \cdots \text{OSe}$ bonds in all four salts, proving that H-bond can drive the self-assembly of HSeO_4^- anions when the cation (a protonated nitrogen base) is either aromatic or aliphatic.

The paper is organized as follows. Firstly, we describe the X-ray structures of the studied crystals. Secondly, we turn to the density functional theory (DFT) calculations with periodic boundary conditions (PBC) in order to construct a model for investigation of proton dynamics. Thirdly, we calculate the quasi-adiabatic potentials for proton transfer in order to describe the effects of proton delocalization. Fourthly, we predict anharmonic vibrational frequencies of the $\text{SeO}-\text{H}$ stretching modes and the corresponding H/D isotope effects on them, which might be used to characterize the potential of the bridging hydron (proton or deuteron) transfer, and discuss the experimental Fourier-transform infrared (FT-IR) spectra. Finally, we concisely outline cooperativity effects of the coupled $\text{SeO}-\text{H} \cdots \text{OSe}$ and $\text{SeO} \cdots \text{H}-\text{N}$ bonds and the interdependence of bridging proton positions in both bonds.

Results and discussion

X-ray: H-bond network in crystal packing

The pyridinium hydrogen selenates **1a-2a** and the 4,4'-bipyridinium analogue **1a-2b** crystallize in the monoclinic $C2/c$ space group and their overall crystal packing show some

similarities (see Fig. 1). Cations and anions are held in their position by the electrostatic attraction between their opposite charges and by a network of the $\text{N}-\text{H} \cdots \text{O}$ and $\text{C}-\text{H} \cdots \text{O}$ bonds involving the pyridinium moiety. As expected, the former interactions are shorter than the latter, the $\text{N} \cdots \text{O}$ separations of the $\text{N}-\text{H} \cdots \text{O}$ bonds are 2.717 Å in **1a-2a** and 2.644 Å in **1a-2b**, while the mean values of the $\text{C} \cdots \text{O}$ separations of the $\text{C}-\text{H} \cdots \text{O}$ bonds are 3.343 Å and 3.362 Å, respectively. Importantly, in both **1a-2a** and **1a-2b** crystals, the HSeO_4^- anions are assembled into infinite H-bonded supramolecular chains. As H-bonding occurs between anions, *i.e.* entities with the same net charge, such hydrogen bonds were called ‘anti-electrostatic’ in the literature.²¹ Close contacts between anions and cations contribute to disperse the anion negative charge, allowing for short $\text{SeO}-\text{H} \cdots \text{OSe}$ bonds: 2.610 Å in **1a-2a** and 2.605 Å in **1a-2b**.

The salt **1a-1b-2c** crystallizes in the triclinic $P\bar{1}$ space group. The cation–anion assembly is secured by a tight net of the $\text{N}-\text{H} \cdots \text{O}$ and $\text{C}-\text{H} \cdots \text{O}$ bonds as in **1a-2a** and **1a-2b**, while the anion–anion assembly is different from that of the two hydrogen selenates discussed above. The HSeO_4^- and SeO_4^{2-} anions **1a** and **1b** form H-bonded dimers. Two such dimers further couple back-to-back *via* two water molecules bridging the SeO_4^{2-} units (see Fig. 2). Interestingly, the $\text{O} \cdots \text{O}$ separation in the $\text{SeO}-\text{H} \cdots \text{OSe}$ supramolecular synthon which forms the anion dimers in **1a-1b-2c** is 2.538 Å long, namely, it is even shorter than in **1a-2a** and **1a-2b**. This is rather counterintuitive as it may be expected that the Coulombic repulsion in an anion–anion adduct is greater than in anion–anion adducts. However, this distance is similar to those reported for dimers formed by doubly negatively-charged anions such as

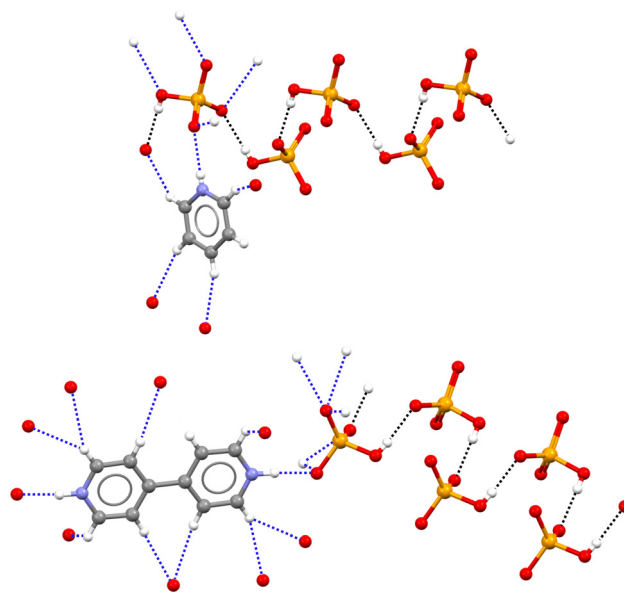


Fig. 1 Partial view of the H-bonded anion chains in **1a-2a** (top) and **1a-2b** (bottom); one cation unit is also shown. H-bonds formed by the cation and one anion unit are indicated; those assembling the supramolecular anion chain are black dotted lines; those between cation and anion units are blue dotted lines. Colour code: grey, carbon; whiteish, hydrogen; red, oxygen; indigo, nitrogen; orange, selenium.



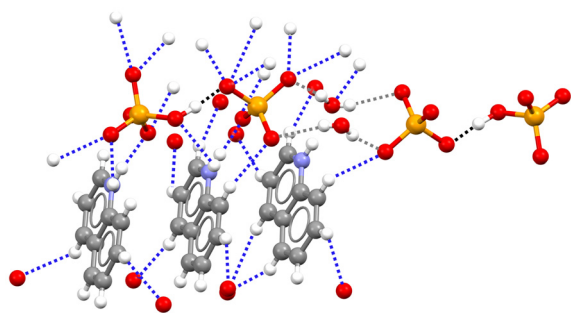


Fig. 2 Partial view of the crystal packing in **1a-1b-2c** showing two anion...dianion dimers linked by two bridging water molecules. Three quinolinium units are shown, which are involved in the tight net of H-bonds with both **1a** anion and **1b** dianion; H-bonds forming anion...dianion dimers are black dotted lines, anion...water H-bonds are grey dotted lines, other H-bonds are blue dotted lines. Colour code as in Fig. 1.

HPO_4^{2-} or HAsO_4^{2-} .²¹ The short distance observed in **1a-1b-2c** salt is probably related to the pronounced acidic character of HSeO_4^- ($\text{p}K_a \sim 1.9$) in combination with the marked basic behaviour of SeO_4^{2-} ($\text{p}K_b \sim 12$).

Of note, in these three structures, π -stacking interactions between aromatic moieties favour the formation of distinct cationic and anionic regions, and thus the anion...anion pairing.

Similar to the three salts described above, in the 2,2,6,6-tetramethylpiperidinium hydrogen selenate **1a-2d** (monoclinic space group $P2_1/n$), a tight network of C-H...O and N-H...O bonds contributes to locking cations and anions in their position. The ability of HSeO_4^- unit to act as both donor and acceptor of H-bonds, *i.e.* its amphoteric character, is robust enough to drive the self-assembly of two HSeO_4^- units into a supramolecular dimer with 'dangling' SeO-H bond. A water molecule further bridges the two HSeO_4^- units of this dimer and an H-bonded cyclic array is produced. Two such cyclic arrays are linked together *via* their respective water molecules (see Fig. 3). The O...O separation is 2.590 Å in the anion...anion H-bond, while the mean value of the O...O separations in the anion...water H-bonds is 2.748 Å.

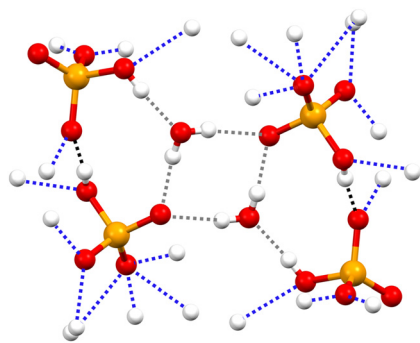


Fig. 3 Partial view of the anion supramolecular assemblies in **1a-2d**. H-bonds formed by anions with another anion, with a cation, and with water are black, blue, and grey dotted lines, respectively. Colour code as in Fig. 1.

X-ray crystallographic data can hardly provide a realistic description of the degree of delocalization of the hydrogen atoms involved in the anion... (di)anion H-bonds. As the quantum nature of proton plays profound role in the H-bonding and proton transfer processes, a firm cooperation between experimental and theoretical methods is required in order to achieve deeper insight into both phenomena. In the following subsections we describe the results of our DFT computations with PBC for the reported structures aiming to understand the bridging proton dynamics and cooperativity effects of neighbouring H-bonds.

Fully optimized crystal structures: comparison with experiment

The optimized crystal structures stay very similar to the experimental ones and their overall bonding motif is preserved (visual comparison between X-ray and DFT data is given in Fig. S1–S4 in ESI†). All four crystals are composed of large unit cells with low point group symmetry: 4 symmetry operators for **1a-2a** and **1a-2b**, 2 symmetry operators for **1a-1b-2c** and **1a-2d**. All structures are held by various non-covalent interactions exhibiting broad range of lengths and strengths: (i) dominating O-H...O and N-H...O interactions, (ii) π -stacking-type dispersive interactions between protonated aromatic N-heterocycles (pyridine, 4,4'-bipyridil and quinoline in **1a-2a**, **1a-2b**, and **1a-1b-2c**, respectively), and (iii) numerous additional C-H...O contacts. The SeO-H...OSe bonds are fairly linear (the OHO angles cover the 169°–176° range) and are formed along the directions of the expected lone pair localizations on the proton-accepting O atoms.

The equilibrium O...O separation of 2.512–2.544 Å can be considered as the boundary between moderate and short strong H-bonds.²² A similar O...O separation of 2.54 Å was observed in the crystal structures of H-bonded dimeric selenates of alkali metals: experimentally in $\text{Cs}_3\text{L}(\text{SeO}_4)_2$ (L = H/D) by Matsui *et al.*^{6,7} and computationally in $\text{Rb}_3\text{H}(\text{SeO}_4)_2$ by Pavlenko *et al.*⁵

In **1a-2a** and **1a-2b**, the infinite chains of HSeO_4^- anions are assembled by the O-H...O bonds along the crystallographic *b* axis. Together with cations, they form a herringbone structure (see Fig. 4); in the crystals, neighbouring herringbone structures are antiparallel. **1a-1b-2c** and **1a-2d** feature 'isolated' H-bonds between selenate moieties. These bonds are formed between SeO_4^{2-} and HSeO_4^- (**1a-1b-2c**) or between two HSeO_4^- ions (**1a-2d**). In both structures there are also H-bonds with water molecule that occurs once per the irreducible fragment (see Fig. 5).

The lattice parameters for the minimum-energy and X-ray crystal structures of **1a-2a**, **1a-2b**, **1a-1b-2c** and **1a-2d** along with percent changes between them are listed in Table 1. Although the packing and structural arrangement is generally well reproduced, the optimized cell volumes are noticeably smaller than the experimental ones (net contraction from 5.9% for **1a-2d** to 9.5% for **1a-2b**). The factors affecting this can be arranged into the following clauses.

(i) The studied crystals are held together by networks of H-bonds, in which heavy atoms are subject to displacements under the influence of weak intermolecular forces. In calculations, these forces are reproduced with varying degrees of



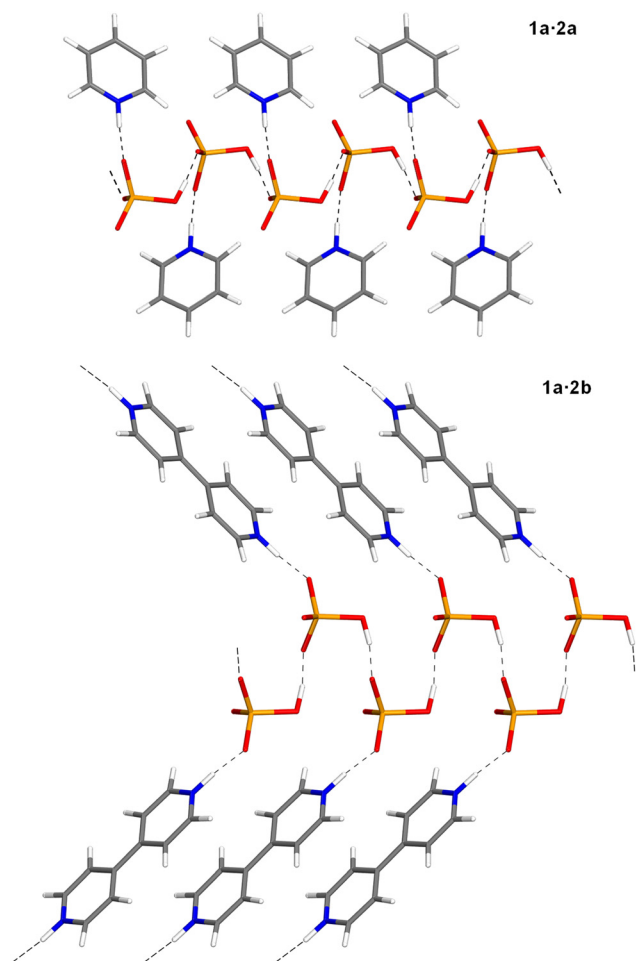


Fig. 4 Optimized structures of **1a-2a** and **1a-2b** (for clarity, chain fragments are shown). Colour code: grey, carbon; whiteish, hydrogen; red, oxygen; blue, nitrogen; orange, selenium.

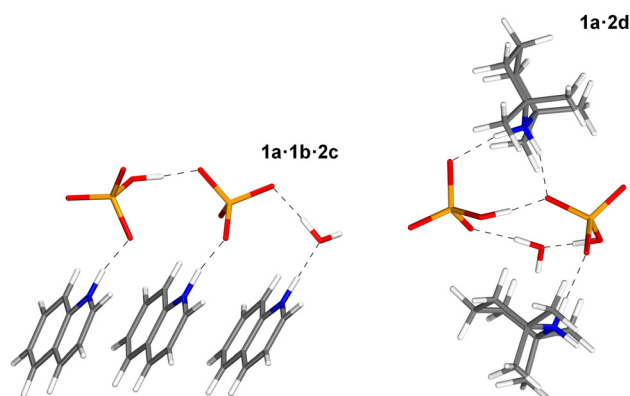


Fig. 5 Optimized structures of **1a-1b-2c** and **1a-2d** (for clarity, irreducible fragments are shown). Colour code as in Fig. 4.

precision and thus the optimized H-bond lengths are among the most fluxional structural parameters. Thus, for **1a-2a**, the largest unit cell dimension change is observed along the *b* axis, which is mostly parallel to the chain of O–H...O bonds. In

contrast, for **1a-2b**, the largest unit cell dimension change is due to adjustments of N–H...O bonds, which extend along the *c* axis (see Table 1). Nevertheless, the full geometry optimization should be imperative as it ensures the optimized crystal structures to become true minima on the potential energy surface (PES).

(ii) Our DFT calculations were performed at an effective temperature of 0 K neglecting the presence of zero-point motions (for re-introduction of these motions in the SeO–H...OSe bonds, *vide infra*), whereas the experimental crystal structures were obtained at 298 K and thus the atomic positions are affected by the non-negligible thermal diffuse scattering effects. As a consequence, theoretical prediction of the crystal volume is expected to be smaller with respect to the experimental one, as was shown by the estimated thermal expansion for various systems, ranging between 1 and 8% from 0 K to room temperature.^{23–25} Acknowledging that single crystal X-ray diffraction may not be particularly tailored for proton localization, we recollected some of the structures at 100 K for further confirmation of the data obtained at 298 K and showcasing the unit cell volume contraction (see Tables S1–S7, ESI†). Of note, **1a-2b** undergoes transition into a different polymorphic phase at 100 K, where the supramolecular motif of the H-bonded HSeO₄[−] infinite chains is fully maintained. This observation confirms the robustness of the synthon and hence the reliability of proton localization in this motif.

(iii) As the B3LYP functional in general poorly models weak long-range van der Waals interactions,^{26,27} this deficiency is compensated here by a mandatory inclusion of *a posteriori* empirical corrections for dispersive forces *via* the Grimme's approach.^{28–30} These corrections, however, are known for their propensity to overestimate long-range attractive forces (see, e.g., recent ref. 31 and 32), possibly causing an underestimation of the lattice parameters in comparison with experiment. Further system-dependent tuning of the dispersion parameters could help to achieve results closer to experimental ones as demonstrated by Civalleri *et al.*;³³ nevertheless, this was not pursued here in order to present a systematic study with consistent parameters across all the studied systems.

In this light, the observed deviations of the cell parameters can be considered acceptable.

QTAIM analysis of the SeOH...OSe bonds

For charge-assisted H-bonds, the topological properties of the electron density and various functions derived from it at the bond critical point (BCP) of the (3, −1) type are of particular interest as they provide valuable information on the very chemical nature of the interatomic interaction.³⁴ For **1a-2a**, **1a-2b**, **1a-1b-2c** and **1a-2d**, the values of the electron density and its Laplacian, ρ_{bcp} and $\nabla^2\rho_{\text{bcp}}$, the local electron kinetic, potential and total energy densities, G_{bcp} , V_{bcp} , and H_{bcp} , and the bond degree index, $H_{\text{bcp}}/\rho_{\text{bcp}}$, at the SeO–H...OSe intermolecular BCPs are listed in Table 2.

The SeO–H...OSe bonds are characterized by $0.10 > \rho_{\text{bcp}} > 0.06$ a.u. and $\nabla^2\rho_{\text{bcp}} > 0$ a.u., which corresponds to the transition region separating the closed-shell (non-covalent)

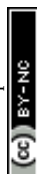


Table 1 Comparison between single-crystal X-ray data (the variance for the last digit is given in parentheses) and fully optimized structures of **1a-2a**, **1a-2b**, **1a-1b-2c** and **1a-2d**. Lattice parameters are given in Å ($\text{\AA}^3$ for the cell volume), angles and degrees. Δ refers to the difference between theory and experiment (in %)

	1a-2a			1a-2b			1a-1b-2c			1a-2d		
	X-ray	B3LYP-D3	Δ	X-ray	B3LYP-D3	Δ	X-ray	B3LYP-D3	Δ	X-ray	B3LYP-D3	Δ
SG ^a	Monoclinic $C2/c$			Monoclinic $C2/c$			Triclinic $P\bar{1}$			Monoclinic $P2_1/n$		
NIA ^b	18			17			68			75		
TNA ^c	72			68			136			300		
<i>a</i>	19.8196(2)	19.5524	−1.3	18.0562(2)	18.0958	+0.2	9.91309(16)	9.66092	−2.5	15.70227(9)	14.97982	−4.6
<i>b</i>	6.26820(10)	5.94464	−5.2	5.15783(5)	5.03361	−2.4	10.3638(2)	10.2068	−1.5	10.12297(5)	9.79767	−3.2
<i>c</i>	13.2167(2)	12.6145	−4.6	16.9795(2)	16.2880	−4.1	14.6046(3)	14.1464	−3.1	16.96831(9)	17.32988	+2.1
α	90.0	90.0		90.0	90.0		76.9481(16)	75.6802	−1.6	90.0	90.0	
β	109.0590(10)	106.7255	−2.1	115.2773(14)	117.5110	+1.9	89.2209(14)	91.8124	+2.9	102.9938(6)	103.5649	+0.6
γ	90.0	90.0		90.0	90.0		71.1833(16)	70.4968	−1.0	90.0	90.0	
<i>V</i>	1551.95	1404.18	−9.5	1429.91	1315.86	−8.0	1380.70	1264.04	−8.4	2628.11	2472.51	−5.9

^a Space group. ^b Number of irreducible atoms. ^c Total number of atoms in the unit cell.

and shared (covalent) interatomic interactions.^{37–39} On the one hand, relatively high values of ρ_{bcp} indicate a significant overlap of the closed shells of hydrogen and oxygen atoms.^{40,41} On the other hand, positive values of $\nabla^2\rho_{\text{bcp}}$ are a signature of non-covalent interactions.⁴² These findings are in accordance with the relative values of the local electron kinetic and potential energy densities, as well as the bond degree index. Namely, inequalities $2 > |V_{\text{bcp}}|/G_{\text{bcp}} > 1$ and $H_{\text{bcp}}/\rho_{\text{bcp}} < 0$ are also indicative of the character of the H-bonds, intermediate between covalent and non-covalent interactions.^{37,43} This is further confirmed by the H-bond energies, E_{OHO} (see Fig. 6), estimated from the G_{bcp} values.^{35,36} E_{OHO} values are around 70 kJ mol^{-1} and are close to each other: the span is only 6.5 kJ mol^{-1} . This allows us to firmly classify such H-bonds as moderately strong ones. For completeness, the 2D maps of ρ and $\nabla^2\rho$ around the $\text{SeO} \cdots \text{H} \cdots \text{OSe}$ bonds for all four studied systems in their equilibrium geometries are shown Fig. S5 and S6 (ESI†).

In order to provide a more complete picture of the non-covalent interactions in the studied systems, we additionally performed a comparative QTAIM analysis of the anion $\cdots$ cation and anion $\cdots$ water interactions within the H-bond networks, as well as the π -stacking type $\text{C} \cdots \text{C}$ and $\text{C} \cdots \text{N}$ dispersive interactions between aromatic cation units (see Fig. S7 and the following discussion in ESI†).

Delocalization of bridging hydrons within the $\text{SeOH(D)} \cdots \text{OSe}$ bonds

In this subsection we focus on the bridging proton quantum delocalization within the $\text{SeO} \cdots \text{H} \cdots \text{OSe}$ bonds, additionally exploring the changes in the H-bond geometry and bridging particle delocalization, caused by proton–deuteron substitution.

Here, we consider the proton delocalization as the result of its motion in an effective potential along a proton transfer reaction coordinate. The latter was chosen for the following considerations.

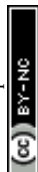
The bridging proton position and the heavy atom distance are closely linked to each other,⁴⁴ which is manifested, for example, by the quite strong coupling between the $\nu(\text{O} \cdots \text{H})$ and $\nu_{\text{e}}(\text{OH} \cdots \text{O})$ vibrational modes.⁴⁵ This means that neither the

Table 2 QTAIM parameters (in a.u.) at the intermolecular BCPs of the $\text{SeO} \cdots \text{H} \cdots \text{OSe}$ bonds and estimated corresponding bond energies E_{OHO} (in kJ mol^{-1}) = $0.429 G_{\text{bcp}}$ (see ref. 35 and 36) for **1a-2a**, **1a-2b**, **1a-1b-2c** and **1a-2d**

	1a-2a	1a-2b	1a-1b-2c	1a-2d
ρ_{bcp}	0.0728	0.0725	0.0829	0.0715
$\nabla^2\rho_{\text{bcp}}$	0.1401	0.1437	0.1363	0.1454
G_{bcp}	0.0592	0.0595	0.0650	0.0592
V_{bcp}	−0.0834	−0.0830	−0.0959	−0.0820
$ V_{\text{bcp}} /G_{\text{bcp}}$	1.4085	1.3959	1.4754	1.3858
H_{bcp}	−0.0242	−0.0235	−0.0309	−0.0228
$H_{\text{bcp}}/\rho_{\text{bcp}}$	−0.3323	−0.3249	−0.3727	−0.3193
E_{OHO}	66.69	66.99	73.18	66.68

displacement of the proton with fixed heavy atoms, nor the displacement of heavy atoms with fixed $\text{O} \cdots \text{H}$ distance are good candidates for the reaction coordinate. A more realistic choice is given by so-called quasi-adiabatic pathway⁴⁶ along which the light proton position is adjusted ‘on a fly’ to the instantaneous heavy atom positions, in a sort of Born–Oppenheimer separation of nuclear from electronic motions. Thus, we modelled the bridging proton transfer by changing the $\text{O} \cdots \text{H}$ distance r_1 in a step-wise fashion and optimizing all the other geometric parameters, including the $\text{H} \cdots \text{O}$ distance r_2 . As the calculations were performed with PBC, all symmetry-related bridging protons were displaced simultaneously. Such process can be called coherent or synchronous proton transfer (previously studied experimentally, e.g., by Fillaux *et al.*^{47–52}) and in the case of the H-bonded HSeO_4^- infinite chains mentally visualized as an infinite Newton’s cradle. To the best of our knowledge, this is the first time such an approach has been employed in the solid-state simulations.

Previously, the anharmonic O–H stretching in periodic systems was treated as a non-adiabatic bridging particle transfer with O–H distance as a pure normal coordinate decoupled from the manifold of all other modes (mostly in crystals and slabs, see, e.g., ref. 53–60). A more sophisticated – albeit prohibitively demanding – treatment of the correlation between the O–H and $\text{O} \cdots \text{O}$ distances *via* solving a two-dimensional vibrational problem (see, e.g., ref. 61–65) is beyond the thrust of this paper.



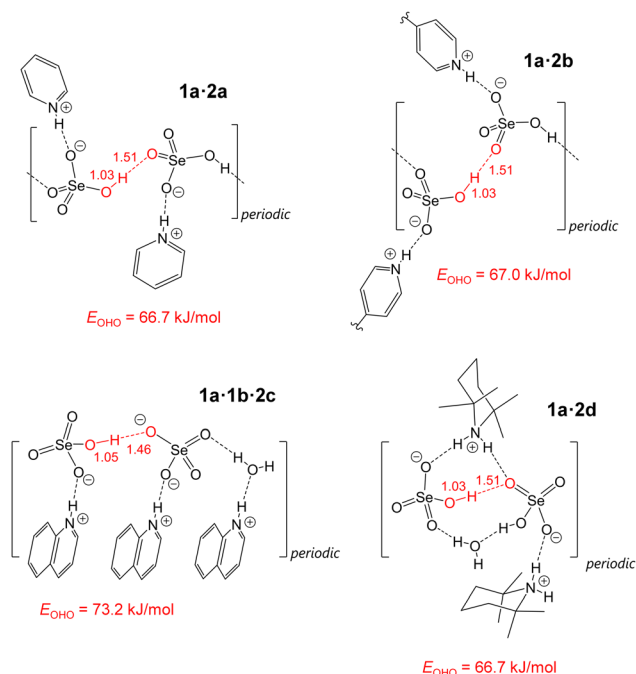


Fig. 6 The optimized H-bond interatomic distances (in Å) for **1a-2a**, **1a-2b**, **1a-1b-2c** and **1a-2d**. The values of the SeO-H...OSe bond energies, E_{OHO} , estimated from the local electron kinetic energy density values at BCP, G_{BCP} , are given below the structures.

In the following, the resulting reaction was expressed as a proton motion along the $q_1 \equiv 0.5(r_1 - r_2)$ coordinate, which is accompanied by the corresponding adjustments of the overall O...O distance, $q_2 \equiv (r_1 + r_2)$. The calculated 1D potentials, $U(q_1)$, along the q_1 coordinate for **1a-2a**, **1a-2b**, **1a-1b-2c** and **1a-2d** are shown in Fig. 7 and 8 together with the zero-point energy (ZPE) and the first excited state energy levels, $\epsilon_{|0\rangle}$ and $\epsilon_{|1\rangle}$ (hereafter, we refer to them as $|0\rangle$ and $|1\rangle$ vibrational states, respectively), squares of corresponding vibrational wave functions, $\Psi_{|0\rangle}^2(q_1, L)$ and $\Psi_{|1\rangle}^2(q_1, L)$, $L = \text{H/D}$, as well as

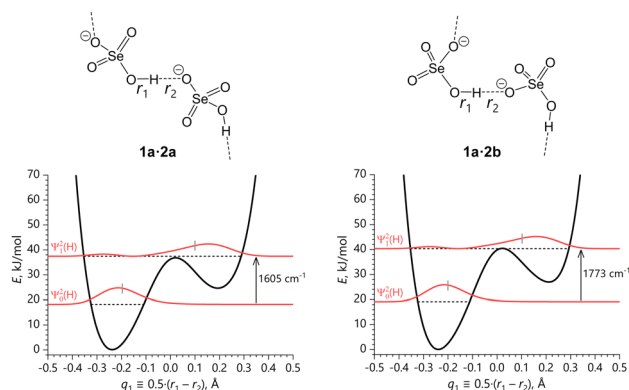


Fig. 7 Asymmetric double-well potentials of the coherent quasi-adiabatic transfer of the bridging protons within the infinite SeO-H...OSe bond chains along the q_1 coordinate with the corresponding $|0\rangle$ and $|1\rangle$ vibrational states (horizontal dashed lines at $\epsilon_{|0\rangle}$ and $\epsilon_{|1\rangle}$) and fundamental transition frequencies pertaining to **1a-2a** and **1a-2b**. Short grey vertical bars indicate the vibrationally-averaged positions of the proton q_1 coordinate.

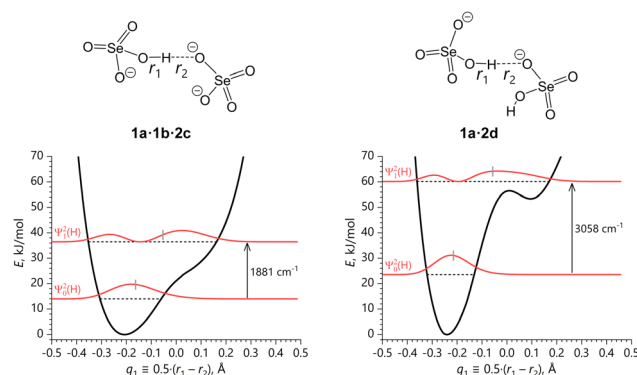


Fig. 8 Asymmetric single-well potentials of the coherent quasi-adiabatic transfer of the bridging protons within the 'isolated' SeO-H...OSe bonds along the q_1 coordinate with the corresponding $|0\rangle$ and $|1\rangle$ vibrational states (horizontal dashed lines at $\epsilon_{|0\rangle}$ and $\epsilon_{|1\rangle}$) and fundamental transition frequencies pertaining to **1a-1b-2c** and **1a-2d**. Short grey vertical bars indicate the vibrationally-averaged positions of the proton q_1 coordinate.

fundamental transition frequencies, ν_{01} , for O-H stretching. For more on solution of the vibrational problem see DFT calculations subsection. Note that ZPE values correspond to only one vibrational mode, namely, OH/OD stretching mode in the irreducible part of the unit cell. Similar plots including the data for bridging deuterons are shown in Fig. S8 and S9 (ESI[†]), while the geometrical parameters of the H-bonds and relative total energies per unit cell at each q_1 point are summarized in Tables S8–S11 (ESI[†]).

The two cases of **1a-2a** and **1a-2b** (see Fig. 7) featuring infinite H-bond chains are very similar to each other and exhibit bridging proton delocalization in the asymmetric double-well potential, where two energy minima are separated by a barrier which is barely as high as the first vibrationally excited level. The potentials do not show two wells of equal depth because of the asymmetry in crystal packing of selenates and cations. The shape of $\Psi_{|0\rangle}^2(q_1, \text{H})$ indicates that in the ground state the bridging proton is localized in the left potential well ($q_1 < 0$), while in the first excited state it is localized in the right well ($q_1 > 0$), though the $\Psi_{|1\rangle}^2(q_1, \text{H})$ is quite broad, has a node and a second maximum at negative q_1 .

From the energy difference between lowest two energy levels, *i.e.* ν_{01} frequencies, it follows that the $|0\rangle$ states, being localized in the deeper well, are by far the largest contributors in thermal equilibrium at temperatures around 300 K: the Boltzmann populations of $|0\rangle$ states are $p_{|0\rangle} = 0.9995$ for **1a-2a** and $p_{|0\rangle} = 0.9998$ for **1a-2b**, *i.e.* the populations of $|1\rangle$ states remain negligible. At the same time, it means that vibrational excitation (optical phonon excitation) could lead to multiple synchronous proton transfers within the crystal, without large amplitude librations of the SeO₄ tetrahedra and molecular cations that would exchange the roles of proton-donating and proton-accepting oxygen atoms (*i.e.* without the 'flip' of the double-well potential itself). Furthermore, the density distributions of the first excited states, $\Psi_{|1\rangle}^2(q_1, \text{H})$, are predominantly localized in the shallower well in proximity of the barrier top in both cases, which means that energy of the $|1\rangle$ states is high

enough to allow for energy-free ‘cross’ transitions between the two possible off-centre sites, as suggested by the non-negligible probability distribution in the deeper well. A similar description is valid for bridging deuterons (see Fig. S8, ESI†), though the vibrationally excited wave function $\Psi_{11}^2(q_1, D)$ is more smeared over two local minima.

We propose therefore that the bridging hydrons in **1a-2a** and **1a-2b** ought not to be regarded as moving independently in individual double-well potentials, but as participating in a collective mode that is promoted by a vibrational perturbation. This mode is the coherent hydron transfer from the more stable configuration $\text{SeO-H(D)} \cdots \text{OSe}$ to the vibrationally excited configuration $\text{SeO} \cdots \text{H(D)-OSe}$. The latter also meets the spatial periodicity of the crystal lattice and has a limited lifetime.

The $\text{SeO-H} \cdots \text{OSe}$ bond in **1a-1b-2c** is shorter than in **1a-2a** or **1a-2b** and the barrier between the two potential minima is absent, making the proton transfer potential an asymmetric single-well shown in Fig. 8, left. In comparison with **1a-2a** and **1a-2b**, the broader potential minimum corresponds to a more delocalized $|0\rangle$ state with wider $\Psi_{10}^2(q_1, H)$ distribution along the q_1 coordinate. Despite the absence of the second potential minimum, due to the asymmetry of the potential the average proton position in the first excited state is substantially shifted towards the accepting oxygen atom, and the most probable proton position (*i.e.* the maximum of $\Psi_{11}^2(q_1, H)$) is located virtually in the H-bond centre, making it effectively a central-symmetric H-bond with a shared proton. A similar observation remains true for the bridging deuterons (see Fig. S9, left, ESI†), though the domain spanned by the vibrationally excited wave function $\Psi_{11}^2(q_1, D)$ is more contracted as opposed to the case of **1a-2a** and **1a-2b**.

Finally, the $\text{SeO-H(D)} \cdots \text{OSe}$ bond potential in **1a-2d** incorporates the features outlined in the preceding three potentials, possessing a second local minimum – though an utmost shallow – as in **1a-2a** and **1a-2b**, but a rather steep potential shape along the q_1 coordinate, making the $\Psi_{11}^2(q_1, H)$ distribution akin to the one in the single-well potential in **1a-1b-2c** (see Fig. 8, right). This leads to the dramatically high ν_{01} frequency of the O–H stretching mode featuring pronounced negative anharmonicity⁶⁶ in comparison with the shallower harmonic potential, and to the narrowest $\Psi_{10}^2(q_1, H)$ distribution in the series (*vide infra*).

Let us now summarize how the motions of the three atoms forming OLO bridges ($L = \text{H/D}$) influence the effective H-bond geometry. The four studied crystals vary from each other in the H-bond length, which is determined by the crystal packing and the positioning of the counterions. However, there are also three important common features.

(i) Account for hydron delocalization yields quantum-mechanically induced contraction and symmetrization of the $\text{SeO-L} \cdots \text{OSe}$ bond (see Fig. 9 and Table 3, Table S12, ESI†). The heavy atom distance decreases by 0.05–0.09 Å, while the bridging proton shifts *ca.* 0.08 Å closer to the H-bond centre.

(ii) Due to significant anharmonicity of the potentials, the main effect of replacing deuteron over proton is a shift of the $\Psi_{10}^2(q_1, D)$ maxima towards the energy minimum (see grey vertical bars in

Fig. S8 and S9, ESI†), leading to the negative primary geometric isotope effect (primary GIE):

$$\Delta\langle q_1 \rangle = \langle q_1^D \rangle - \langle q_1^H \rangle < 0,$$

and to the small positive *secondary* geometric isotope effect (secondary GIE or ‘Ubbelohde effect’^{67,68}):

$$\Delta\langle q_2 \rangle = \langle q_2^D \rangle - \langle q_2^H \rangle > 0.$$

In other words, deuteration shortens the ‘covalent’ bond and lengthens the ‘H-bond’, making ODO bridge slightly longer and more asymmetric than OHO one.

(iii) As quantum-mechanical property of deuteron is suppressed in comparison with proton, the full width at half maximum (FWHM) for $\Psi_{10}^2(q_1, D)$ is smaller than that for $\Psi_{10}^2(q_1, H)$, *i.e.* deuteron is less delocalized than proton.

In the following section we discuss the optical spectral manifestations of the delocalized $\text{SeO-H(D)} \cdots \text{OSe}$ bridging protons (deuterons) on the basis of the calculated anharmonic stretching frequencies, and compare them with the experimental FT-IR spectra.

Anharmonicity of the vibrational O–H(D) frequencies

The broad and strongly anharmonic shapes of the bridging hydron potentials (some with low barriers and shallow second minima) lead to the following three interesting phenomena.

(i) Extreme difference between harmonic and anharmonic $\nu(\text{O-L})$ frequencies. On the one hand, the bottom wells of the calculated potentials are rather broad, even for the narrowest one in **1a-2d**, so already the DFT-calculated harmonic O–L stretching frequencies, $\nu_{\text{harm}}(\text{O-L})$, are low enough to indicate that the intermolecular $\text{SeO-L} \cdots \text{OSe}$ bonds in all considered crystals are quite strong,^{44,69} in accordance with the equilibrium geometries and QTAIM analysis. On the other hand, taking into account the anharmonicity lowers the frequencies

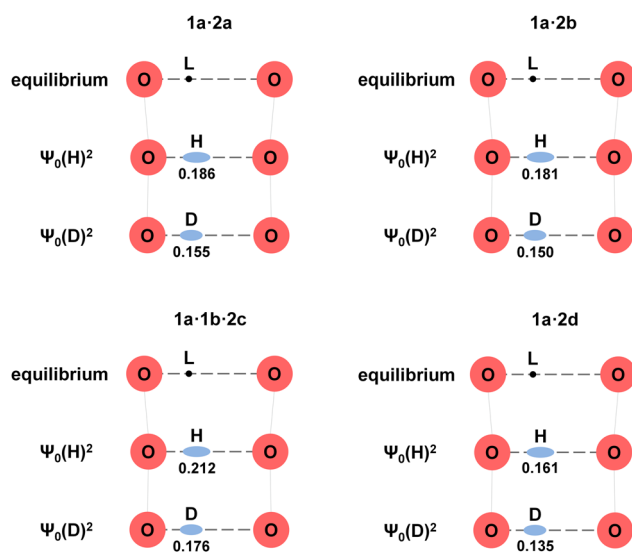


Fig. 9 Effects of the bridging hydron delocalization on the $\text{SeO-L} \cdots \text{OSe}$ ($L = \text{H/D}$) bond geometry in **1a-2a**, **1a-2b**, **1a-1b-2c** and **1a-2d**. The FWHM for the $\Psi_{10}^2(H)$ and $\Psi_{10}^2(D)$ distributions is given in Å.

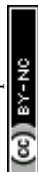


Table 3 Comparison of the SeO–H···OSe bond interatomic distances (in Å) in **1a-2a**, **1a-2b**, **1a-1b-2c** and **1a-2d** according to the single-crystal X-ray data and B3LYP-D3 calculations

	B3LYP-D3 (equilibrium distances)				B3LYP-D3 ($\langle 0 \rangle$ vibrationally-averaged distances)		
	X-ray	O···O	O···O	O–H	H···O	$\langle \text{O} \cdots \text{O} \rangle$	$\langle \text{O} \cdots \text{H} \rangle$
1a-2a	2.610	2.544	1.034	1.510	2.518	1.064	1.457
1a-2b	2.605	2.543	1.032	1.511	2.512	1.059	1.461
1a-1b-2c	2.538	2.512	1.049	1.463	2.492	1.084	1.411
1a-2d	2.590	2.537	1.028	1.509	2.523	1.050	1.479

even further (see $\nu_{\text{anarm}}(\text{O} \cdots \text{L})$ in Table 4); these values were obtained from Ugliengo's ANHARM code⁷⁰ as described in the ESI†, below Table S13. However, these values are still higher than those obtained using the full shape of the hydron stretching potentials (see $\nu_{01}(\text{O} \cdots \text{L})$ in Table 4). This is understandable, since the position of the first excited level is strongly influenced by the shape of the stretching potential far from the equilibrium, *i.e.* the overall shape of the potential on the other side of the H-bond being apparently the strongest influence.

It is of interest to overlay the calculated vibrational frequencies with the experimental FT-IR absorption spectra (see Fig. S10–S13, ESI†), recorded for powder samples of four studied crystals between 4000 and 400 cm^{-1} at 298 K (the region 1900–2500 cm^{-1} is omitted due to the strong absorption of the diamond UATR crystal). In the range 3500–1500 cm^{-1} , all spectra show several broad absorptions of selenate OH groups, counteraction NH groups, as well as water molecules (in case of **1a-1b-2c** and **1a-2d**). The positions and shapes of many of these bands are hard to discern, especially for exceedingly broad bands of OH groups involved in short and strong H-bonds.

Nevertheless, the overall shapes of the experimental spectra (more precisely, the part appearing due to the stretching vibrations of bridging protons) are reasonably consistent with the calculated vibrational frequencies, as shown in Fig. S10–S13 (ESI†): for visual clarity, Lorentzian lineshapes with arbitrary integral intensities were placed as red and blue traces at $\nu_{01}(\text{O} \cdots \text{H})$ and $\nu_{\text{anarm av.}}(\text{N} \cdots \text{H})$ wavenumbers, respectively.⁷¹

For each system the $\nu_{\text{anarm av.}}(\text{N} \cdots \text{H})$ values were calculated as mean ones over ungerade $\nu_{\text{anarm av.}}(\text{N} \cdots \text{H})$ frequencies, computed for all unique SeO···HN bonds: one in **1a-2a** and **1a-2b**, three in **1a-1b-2c**, and four in **1a-2d**, see Table S14 (ESI†).

Table 4 (An)harmonic treatment of the O–L (L = H/D) stretching frequencies (in cm^{-1}) of A_g symmetry within the SeO–L···OSe bonds in **1a-2a**, **1a-2b**, **1a-1b-2c** and **1a-2d**

	1a-2a	1a-2b	1a-1b-2c	1a-2d
$\nu_{\text{harm}}(\text{O} \cdots \text{H})$	2581	2616	2373	2742
$\nu_{\text{anarm}}(\text{O} \cdots \text{H})$	1806	1863	1814	2231
$\nu_{01}(\text{O} \cdots \text{H})$	1605	1773	1881	3058
$\nu_{\text{harm}}(\text{O} \cdots \text{D})$	1894	1918	1752	2000
$\nu_{\text{anarm}}(\text{O} \cdots \text{D})$	1454	1501	1408	1721
$\nu_{01}(\text{O} \cdots \text{D})$	1650	1828	1484	2638
$\Delta\nu(\text{O} \cdots \text{H})^a$	–201	–90	67	828
$\Delta\nu(\text{O} \cdots \text{D})^a$	196	327	76	917

^a $\Delta\nu(\text{O} \cdots \text{L}) \equiv \nu_{01}(\text{O} \cdots \text{L}) - \nu_{\text{anarm}}(\text{O} \cdots \text{L})$.

As expected from the equilibrium distances, the anharmonic N–H stretching frequencies decrease in the **1a-2d** → **1a-2a** → **1a-2b** → **1a-1b-2c** sequence. Judging from the experimental band shapes in the region 3000–2500 cm^{-1} , the NH band is probably strongly structured, especially for **1a-2a** and **1a-2d**. Apart of that, evidently, the bands of the $\nu(\text{N} \cdots \text{H})$ modes are superimposed with the characteristic and quite intensive bands of $\nu(\text{C} \cdots \text{H})$ modes (2750–3150 cm^{-1} region) and, in the spectra of **1a-1b-2c** and **1a-2d**, also by the $\nu(\text{O} \cdots \text{H})$ bands of structural water molecules (3250–3650 cm^{-1} region).

The presence of extremely broad OH band manifests itself mostly as a raised background around 2000 cm^{-1} and below, matching reasonably well (semi-quantitatively) with the region where our calculations of delocalised protons place the $\nu_{01}(\text{O} \cdots \text{H})$ transitions, shifting to lower frequencies in the **1a-2d** → **1a-1b-2c** → **1a-2b** → **1a-2a** order. It is worth noting that for **1a-2a**, **1a-2b**, and **1a-2d**, some of the band broadening likely stems from the Fermi resonance between the $\nu_{01}(\text{O} \cdots \text{H})$ and $\gamma_{02}(\text{O} \cdots \text{H})$, $\delta_{02}(\text{O} \cdots \text{H})$ modes (*i.e.* between the fundamental of stretching vibration and the overtone of the bending modes), which was shown by Ikeda *et al.* to be a common phenomenon for OHO bonds with the length of ~ 2.6 Å in a solid state.^{72,73}

The most unexpected result is obtained for **1a-2d**. The spectrum shows a unique asymmetric absorption band around 3150–2500 cm^{-1} with complex structure and the absorption maximum at 3019 cm^{-1} , which matches well with the calculated the $\nu_{01}(\text{O} \cdots \text{H})$ value of 3058 cm^{-1} , arising from one of the SeO–H···OSe stretching modes with negative anharmonicity, meaning that both $\nu_{\text{harm}}(\text{O} \cdots \text{H}) = 2742$ cm^{-1} and $\nu_{\text{anarm}}(\text{O} \cdots \text{H}) = 2231$ cm^{-1} frequencies are considerably lower. On the contrary, the calculated the $\nu_{\text{anarm}}(\text{O} \cdots \text{H})$ frequency for the SeO–H···O_{water} stretching (1720 cm^{-1}) in **1a-2d** seems to be in a good agreement with the experimental spectrum, see Table S15 and Fig. S13 (ESI†).

Thereby, we conclude that a reasonable agreement between observed and calculated vibrational stretching frequencies of the short and moderately strong SeO–H(D)···OSe bonds requires thorough account of the bridging proton (deuteron) transfer dynamics extending virtually from one oxygen site to another.

For reference, all computed harmonic wavenumbers, IR and Raman intensities, and band assignments are summarized in Tables S16–S20 (ESI†). The corresponding IR and Raman spectra are visualized in Fig. S14–S17 (ESI†).

(ii) Absence of the correlation between O–L vibrational frequencies and O···L distances. It can be considered a general and well-accepted knowledge that a close-to-linear correlation exists between structural and vibrational properties of O–L bonds: the longer is the equilibrium O···L distance, the smaller is the O–L stretching frequency.^{74–76}

In our case, such correlation seems to hold for calculated harmonic frequencies ($\nu_{\text{harm}}(\text{O} \cdots \text{L})$ vs. $\langle \text{O} \cdots \text{L} \rangle$; see Fig. 10a). It means that regardless of whether there is a second local minimum on the stretching potential or not, the local minimum pertaining to the equilibrium geometry becomes broader (shallower) for larger O···L equilibrium distances (shorter



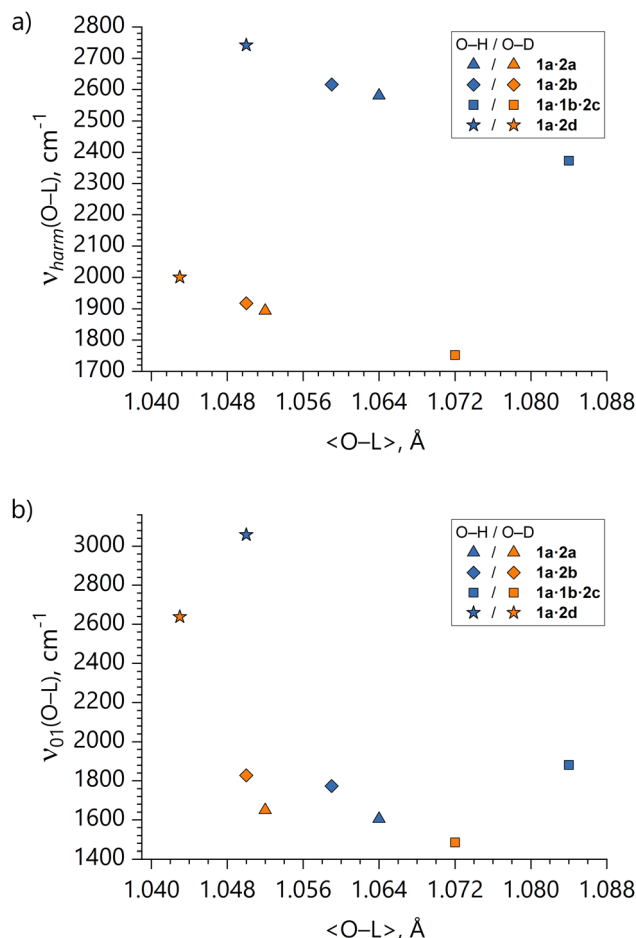


Fig. 10 Dependencies of the (a) harmonic $\nu_{\text{harm}}(\text{O-L})$ and (b) anharmonic $\nu_{01}(\text{O-L})$ stretching frequencies of A_g symmetry on the vibrationally-averaged (O-L) distances in **1a-2a**, **1a-2b**, **1a-1b-2c** and **1a-2d**. The picture does not change qualitatively if one uses equilibrium O-L distances instead of vibrationally-averaged ones.

O...O distances). However, the $\nu_{01}(\text{O-L})$ vs. $\langle\text{O-L}\rangle$ distance correlation is broken (see Fig. 10b). This is due to the fact that transition from the ground state to the first excited state dramatically broadens the potentials especially in case of low-barrier double-well systems **1a-2a** and **1a-2b**, and thus the energy gap between the vibrational levels for these systems is not as large as for the effectively single-well systems **1a-1b-2c** and **1a-2d**. It means that caution ought to be exercised when interpreting experimental vibrational frequencies: for systems with asymmetric double-well potentials the larger O-L vibrational frequency does not necessarily mean a shorter O-L distance.

(iii) Anomalous H/D isotope effect on $\nu_{01}(\text{O-H})$. In cases of low-barrier double-well potentials in **1a-2a** and **1a-2b**, we have predicted a rather exotic H/D isotope effect, *viz.* the higher $\nu_{01}(\text{O-D})$ frequency than the $\nu_{01}(\text{O-H})$ frequency, see Fig. S8 (ESI[†]) and Table 5. Here the isotopic ratio is $\gamma = \nu_{01}(\text{O-H})/\nu_{01}(\text{O-D}) \sim 0.97$, in a good agreement with the previously reported record-low values of $\gamma \sim 0.90$ for several systems with the low-barrier O-H...O bonds exhibiting stretching frequency in the range of

Table 5 Isotopic frequency ratios $\gamma = \nu_{01}(\text{O-H})/\nu_{01}(\text{O-D})$ for the SeO-L...OSe bonds in **1a-2a**, **1a-2b**, **1a-1b-2c** and **1a-2d**

	1a-2a	1a-2b	1a-1b-2c	1a-2d
γ	0.97	0.97	1.27	1.16

ca. 1200–1650 cm^{-1} .⁷⁷ The reason for the anomalous H/D isotope effect could be rationalized as follows: for **1a-2a** and **1a-2b**, approximately at the position of the $|1\rangle$ state the $U(q_1)$ potential becomes significantly broader, which makes the vibrationally excited OH and OD levels closer to each other than the ground levels. As a result, the $\epsilon_{|1\rangle} - \epsilon_{|0\rangle}$ energy difference for D becomes larger than that for H. This phenomenon is further discussed in ESI[†], below Fig. S8.

Cooperativity effects of the SeO-H...OSe and SeO...H-N bonds

Above, we have considered the pivotal features of the SeO-H...OSe bond, but have not yet addressed the question of how the proton transfer in it affects the H-bonding with counterions. In other words, in this subsection we consider the H-bond cooperativity effects (see, *e.g.*, recent ref. 78–82). The OHO and OHN bonds are interconnected through the SeO_4 tetrahedra. As a result, upon the $\text{SeO-H}\cdots\text{OSe} \rightleftharpoons \text{SeO}\cdots\text{H-OSe}$ transfer, the geometries of the $\text{SeO}\cdots\text{H-N}$ bonds are also adjusted ‘on the fly’. The displacement directions of the N-heterocycle molecular cations are shown in Fig. S18 and S19 (ESI[†]), and the explanation is the following.

In the case of ‘isolated’ OHO bonds in **1a-1b-2c** and **1a-2d**, the proton transfer in the OHO bond implies deprotonation of one SeO_4 tetrahedron and protonation of the neighbouring one. The resulting changes of charges on oxygen atoms determine the direction of the NH proton donor displacement. Thus, the cation shifts closer to the SeO_4 tetrahedron gaining an additional negative charge, and, *vice versa*, shifts away from the SeO_4 tetrahedron, whose (formal) charge increases by unity. The resulting range of changes in the N...O interatomic distances is *ca.* 0.10–0.15 Å.

In the case of infinite SeO-H...OSe bond chains in **1a-2a** and **1a-2b**, the proton transfer does not change the SeO_4 tetrahedron charge: deprotonation on its one side is compensated by protonation on the opposite side. As a result, the adjustment of the $\text{SeO}\cdots\text{H-N}$ bond geometry is rather trifling: the N...O interatomic distances vary by *ca.* 0.02–0.03 Å. Furthermore, these changes are influenced by additional non-covalent interactions in the crystal structure, *viz.* weak C-H...O contacts between the aromatic protons of the cations and various oxygen atoms of the nearest SeO_4 tetrahedra. As a result, the direction of the cation shift may seem intuitively unexpected. For example, for **1a-2b** the symmetrization of the OHO bond weakens the OHN bond.

A more detailed description of the cooperativity effects of the SeO-H...OSe and SeO...H-N bonds is provided in the ESI[†], below Fig. S19.

Conclusions

In this paper, we have endeavoured to shed light on the salient, but largely unexplored issue of the bridging proton (deuteron)



delocalization in the infinite and 'isolated' short and strong SeO–H(D)···OSe bonds of hydrogen selenate salts in the crystal-line state, employing the novel strategy for anharmonic treatment of the O–H(D) stretching modes, which implies the quasi-adiabatic potential energy calculations with a subsequent numerical solution of the vibrational Schrödinger equation for the bridging hydron. The obtained results can be summarized as the following.

(i) The bridging proton (deuteron) delocalizations in SeO–H(D)···OSe bonds are caused by the zero-point vibrations within broad and shallow stretching potentials, ranging from an asymmetric low-barrier double well to an asymmetric single well or high-barrier double well. The shape of the potential sensitively responds to the change in the O...O distance: its contraction by only a few hundredths of Å leads to the disappearance of potential barrier and to a greater hydron delocalization in the ground state. In turn, the O...O distance could be adjusted by the placement of OHN-bonded counterions due to the mutual coupling of OHO and OHN bonds.

(ii) The calculated equilibrium OH distances are 0.03–0.05 Å longer than those obtained by the automatic placement of hydrogen atoms in X-ray refinement procedure. In turn, the vibrationally averaged OH distances are longer still, by additional 0.03–0.05 Å. The quantum delocalization of hydrogens, estimated from the width of the vibrational wave function, is in the order of 0.2 Å. Since the zero-point energy depends on the nuclear masses, one observes an isotope dependence of the H-bond distances, *i.e.* the H-bond strength is (slightly) modified.

(iii) The O–H stretching frequencies lie in the range of 1600–3050 cm^{−1}, which is substantially determined by the width of the potential at the height of the first vibrationally excited level. Because of this, in case of low-barrier double-well potentials the frequency of the fundamental transition for the lighter proton is lower than that for the heavier deuteron by *ca.* 50 cm^{−1}, *i.e.* $\nu_{01}(\text{O–H}) < \nu_{01}(\text{O–D})$. In the experimental IR spectra, the OH/NH stretching bands are very broad and sometimes barely distinguishable from the background. Because of it, instead of using experimental IR spectra to validate calculations, one could use calculations to assign ambiguous experimental IR spectra in the region of broad OH/NH stretching bands.

(iv) An optical phonon excitation to the first excited state dramatically affects the OHO/ODO bridging hydron position. In the case of 'isolated' bonds featuring effectively single-well potentials, the hydron is shifted towards the H-bond centre, making it almost equally shared between two oxygen atoms. In the case of infinite chains of H-bonded selenates with double-well potentials, the hydron is almost entirely transferred across the H-bond centre. One can thus deduce that the degree of the excited-state hydron transfer in hydrogen selenates can be fine-tuned by changing the separation of oxygen sites, which, in turn, is controlled by the counterions.

In conclusion, an ordered H-bond network in the hydrogen selenate salts introduces strong anharmonic effects in the SeO–H(D)···OSe bonds, which are reflected in the hydron position and delocalization extent, in H-bond vibrational spectral properties, and in the peculiarities of hydron transfer. We were able

to account for these anharmonic effects only by utilizing the entire shape of the hydron stretching potential extending far from the equilibrium, all the way to another oxygen site.

Experimental

Materials and methods

All compounds were purchased from commercial suppliers (Sigma-Aldrich, TCI America, and Merck) and used without further purification.

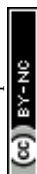
Crystal sample preparation. All the salts were prepared according to the following procedure. Aqueous solutions containing equimolar amounts of selenic acid and the correspondent base were mixed together under constant stirring. The solutions were left to slowly evaporate in a clear borosilicate vial at room temperature. Single crystals suitable for X-ray diffraction were produced within two weeks.

X-ray diffraction analysis. Data collections were performed at 300 K and 100 K using a XtaLAB Synergy diffractometer equipped with a HyPix detector. Unit cell refinement and data reduction were performed using CrysAlisPro 1.171.41.98a. Structures were solved by direct methods using SHELXT and refined by full-matrix least-squares on F^2 with anisotropic displacement parameters for the non-H atoms using Olex2. Absorption correction was performed based on multiscan procedure.^{83–86} X-ray structures were visualized using ball and stick representation built in Mercury software (version 2022.3.0).⁸⁷

FT-IR analysis. FT-IR spectra were collected at room temperature using a Nicolet Nexus FT-IR spectrometer equipped with UATR unit (a diamond crystal). Spectra represent an average of 64 scans recorded in a single beam mode with a 2 cm^{−1} resolution and corrected for the background.

DFT calculations. All quantum-chemical calculations for **1a-2a**, **1a-2b**, **1a-1b-2c** and **1a-2d** were performed with PBC using LCAO approach implemented in the CRYSTAL17 software (version 1.0.2).⁸⁸

The B3LYP^{89,90} functional with the two-body D3(BJ)-damping approach^{28,29,91} for the long-range dispersion correction was employed. All-electron BSSE-corrected GTO basis sets POB-TZVP for Se, O, N, C and H atoms (latest 2019 versions⁹² taken from the CRYSTAL web page⁹³) were used. The chosen combination of the DFT functional and the atomic basis set has been proven to reproduce the H-bond geometric features in a solid state with sufficient accuracy, see, *e.g.*, recent ref. 94–98. Calculations of the Coulomb and Hartree–Fock exchange series were done with the accuracy controlled by a set of overlap tolerances, which were taken to be (10^{−9}, 10^{−9}, 10^{−9}, 10^{−9}, 10^{−18}).⁹⁹ The maximum order of multipolar expansion^{100–102} in the long-range zone for the electron–electron Coulomb interaction was set to 6. Calculation of the DFT exchange–correlation contribution over the unit cell volume was done with the XLGRID DFT grid specification (a pruned grid comprising 75 radial points and five angular shells with a maximum of 974 angular points generated through the Gauss–Legendre radial



quadrature and Lebedev two-dimensional angular point distributions, respectively^{99,103}), chosen for its accuracy, which is better than 0.0001% for the integrated charge density; tolerances for the DFT density and the DFT grid weight were taken to be 10^{-10} , 10^{-18} . Integration in the Brillouin zone was performed using the Monkhorst–Pack scheme¹⁰⁴ for a $12 \times 12 \times 12$ k -point grid, corresponding to 476, 476, 868 and 518 reciprocal space points at which the Hamiltonian matrix was diagonalized for **1a-2a**, **1a-2b**, **1a-1b-2c** and **1a-2d**, respectively. Tolerance on energy controlling the SCF convergence (employing the DIIS extrapolator technique^{105,106}) for geometry optimizations and phonon frequencies calculations was set to 10^{-8} and 10^{-10} Hartree, respectively.

Experimental single-crystal X-ray structures at 298 K were used as the starting point for the DFT calculations. The lattice parameters and atomic centres were optimized with tight convergence criteria (maximum/RMS forces and displacements of atoms smaller than 0.000075/0.000050 a.u. and 0.000225/0.000150 a.u., respectively) and symmetry constraints imposed by a monoclinic space group $C2/c$ for **1a-2a** and **1a-2b**, a monoclinic space group $P2_1/n$ for **1a-2d**, and a triclinic space group $P\bar{1}$ for **1a-1b-2c**. Analytical energy gradients^{107–109} within a quasi-Newton approach¹¹⁰ combined with the BFGS^{111–115} algorithm for Hessian updating were used. Consistency between the optimized geometries and the aforementioned overlap tolerances for integrals' evaluation was ensured by the FINALRUN option with the value of 4.⁹⁹

For optimized structures, phonon frequencies within the harmonic approximation were calculated numerically at the Γ point using the direct mode-following (frozen-phonon) method.¹¹⁶ For deriving the dynamical matrix, the 0.006 Å displacement was chosen both for the two-point (central difference) approach for **1a-2a** and **1a-2b** and for the one-point (right difference) approach for **1a-1b-2c** and **1a-2d**.⁹⁹ The analytical IR¹¹⁷ and Raman^{118–120} intensities for normal modes were calculated from the dipole moment and polarizability derivatives determined using CPHF approach¹²¹ for constructing the Born charge and Raman tensors, respectively. Integrated Raman intensities were normalized to a maximum value of 1000. The spectra were constructed adopting a pseudo-Voigt functional form with the default VOIGT and DAMPFAC variables of 1.0 (pure Lorentzian functions) and 8.0 (FWHM used for the spectra), respectively. Vibrational mode assignments were based on visual inspection of the corresponding atomic displacements.

The SeO–H···OSe bridging proton transfer pathways (potentials) were calculated in the quasi-adiabatic manner by stepwise elongating/contracting the frozen O–H/H···O distance with an increment of 0.02–0.04 Å. All the other degrees of freedom (internal coordinates) were optimized at each step employing the conjugate gradient BKTRNSF2 algorithm⁹⁹ with the fixed lattice parameters and integrals' classification (FIXINDEX directive)⁹⁹ pertaining to the corresponding optimized structures. To secure the optimization stability, the maximum coordinate change allowed between steps was constrained to 0.05 a.u. Such a procedure envisages the anharmonic coupling

in the potential energy between the $\nu_{\text{O}}(\text{OO})$ and $\nu(\text{OH})$ modes, and ensures physically sound modelling of coherent transfer of the symmetry-related bridging protons without reducing the unit cell symmetry. The four unique 1D pointwise potentials thereby calculated were subjected to polynomial fitting in order to get a suitable form for further processing.

Assuming perfect fulfilment with the Born–Oppenheimer approximation (*i.e.* when the only conceivable source of the isotope effect on the H-bond characteristics is the isotope mass dependence of the vibrational motion), the Schmidt's Wave-Packet MATLAB package^{122,123} was employed for numerical solution of the one-dimensional vibrational time-independent Schrödinger equations both for protons and deuterons with the calculated proton transfer potentials $U(q_1)$ represented as the grids of 1024 potential energy values, evenly tempered within the corresponding range of the q_1 coordinate:

$$\left(-\frac{\hbar^2}{2\mu} \frac{\partial^2}{\partial q_1^2} + U(q_1)\right) \Psi_{|n\rangle}(q_1, L) = \varepsilon_{|n\rangle} \Psi_{|n\rangle}(q_1, L),$$

where $L = H/D$, $\mu = m(L) \times m(\text{O})/m(L) \times m(\text{O})$, and $\Psi_{|n\rangle}(q_1, L)$ being a single variable function. The vibrational problem was solved by diagonalizing the Hamiltonian 1024×1024 matrices based on the Light-Carrington discrete variable representations^{124,125} for spatial discretization by means of the fast Fourier transform. This yielded a series of vibrational energy levels, $\varepsilon_{|n\rangle}$, and corresponding wave functions $\Psi_{|n\rangle}(q_1, L)$. The vibrationally-averaged bridging hydron positions were evaluated as an expectation value of the q_1 coordinate, *i.e.* as $\langle \Psi_{|n\rangle}(q_1, L) | q_1 | \Psi_{|n\rangle}(q_1, L) \rangle$.

Quantum-topological analysis (Quantum Theory of Atoms in Molecules analysis; QTAIM)^{126,127} of periodic electron density was performed using TOPOND14¹²⁸ module implemented in the CRYSTAL17 code. The following electron density features at the SeO–H···OSe intermolecular BCPs were considered: the values of (i) the electron density, ρ_{bcp} , (ii) the Laplacian of the electron density, $\nabla^2 \rho_{\text{bcp}}$, and (iii) the local electron kinetic, potential and total energy densities, G_{bcp} , V_{bcp} and H_{bcp} . The two-dimensional ρ and $\nabla^2 \rho$ distribution maps around the SeO–H···OSe bonds were plotted with the CRYSPLOT¹²⁹ tool.

Manipulation and visualization of the four crystal structures was performed with the BIOVIA's Materials Studio software.¹³⁰ Graphs were plotted with the Origin software.¹³¹

Data availability

The authors confirm that the data supporting the findings of this study are available within the article and ESI,[†] where the authors have cited additional ref. 132–152. Deposition numbers 2417665 and 2417668 (for **1a-2a**), 2417669 and 2417697 (for **1a-2b**), 2417671 (for **1a-1b-2c**), 2417672 and 2417674 (for **1a-2d**) contain the supplementary crystallographic data for this paper. These data are provided free of charge by the joint Cambridge Crystallographic Data Centre and Fachinformationszentrum Karlsruhe Access Structures service.



Conflicts of interest

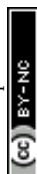
There are no conflicts to declare.

Acknowledgements

This work was financially supported by the Russian Science Foundation (RSF) grant no. 23-13-00095. Quantum-chemical calculations were performed at the Computing Centre of St. Petersburg University Research Park (<https://www.cc.spbu.ru/>). We would like to express our gratitude to Dr A. V. Bandura for illuminating discussions.

Notes and references

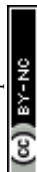
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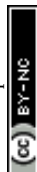
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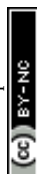
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