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$[Br_4F_{21}]^-$ – a unique molecular tetrahedral interhalogen ion containing a μ_4 -bridging fluorine atom surrounded by BrF_5 molecules†

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The reaction of [NMe₄][BrF₆] with an excess of BrF₅ leads to the compound [NMe₄][Br₄F₂₁]·BrF₅. It features molecular $[(\mu_4-F)(BrF_5)_4]^-$ anions of tetrahedron-like shape containing central μ_4 -bridging F atoms coordinated by four BrF₅ molecules. It is the most BrF₅-rich fluoridobromate anion by mass. Quantum-chemical calculations showed that the μ_4 -F-Br bonds within the anion are essentially ionic in nature. The compound is the first example where F atoms bridge μ_4 -like neither to metal nor to hydrogen atoms. It was characterized by Raman spectroscopy and by single-crystal X-ray diffraction. The latter showed surprisingly that its crystal structure is related to the intermetallic half-Heusler compound and structure type MgAgAs.

Introduction

The halogen fluorides form a class of chemically most reactive compounds, some of them even exceeding the reactivity of F2 itself. BrF3, BrF5, and ClF3 are hypergolic oxidizers that may react vigorously, sometimes explosively, with many organic and inorganic substrates and materials if conditions are not controlled very carefully.1,2 If non-oxidizable starting materials are used, such violent reactions are not to be expected, the respective halogen fluoride is not decomposed and may act as a ligand. Towards Lewis acids, such as BF₃, AsF₅ or SbF₅, the halogen fluorides can act as fluoride ion donors, forming the corresponding fluoridohalonium cations,3,4 whereas towards Lewis basic compounds, such as the alkali metal fluorides, many halogen fluorides can act as a fluoride ion acceptors, forming fluoridohalogenate anions.5,6 Currently known poly halogen and interhalogen anions are summarized in Table S1.† Currently known poly (inter-) halogen cations are listed in Table S2.† Besides numerous "mononuclear" halogen fluoride ions, only a few examples have been discovered, where halogen fluoride molecules and their derived ions are linked by bridging F atoms forming "oligonuclear" halogen fluoride ions. To the best of our knowledge, only two cations, namely $[Br_2F_8]^+$ and $[Br_3F_8]^{+,7}$ and seven anions of this type are currently known. $[(\mu_3-F)(ClF)_3]^-,^8$ These $[(\mu_3-F)(ClF_3)_3]^{-,9}$

F)(BrF₃)₂]⁻,¹⁰⁻¹² [(μ_3 -F)(BrF₃)₃],¹¹ two different isomers of the [Br₄F₁₃]⁻,¹³ [(μ_3 -F)(BrF₅)₃]⁻,¹⁴ and [(μ_3 -F)(IF₅)₃]⁻.^{15,16} All those "oligonuclear" anions consist of a central μ - or μ_3 -bridging fluoride ion acting as a linker between two or three halogen fluoride moieties. Depending on the isomer, the [Br₄F₁₃]⁻ anion formally consists of a [(μ_3 -F)(BrF₃)₃]⁻ anion with a further BrF₃ unit attached to one of the terminal F atoms or of a [(μ -F)(BrF₃)₂]⁻ anion which is linked to two further BrF₃ units *via* two different terminal F atoms.¹³

Tetracoordinated F atoms are present in ionic compounds such as CaF_2 . In contrast, only a few complexes with μ_4 -F atoms are known in Y, V, Mo, and Al compounds. The Molecular ions containing fluorine atoms that are tetracoordinated by non-metal atoms are generally very scarce, with the tetrahydrogen pentafluoride ion, $[(\mu_4$ -F)(HF)₄]⁻, The likely being the most prominent example. We recently reported the compound Cs $[Br_3F_{16}]$, which contained the first fluoride-bridged oligonuclear Br(v) anion, $[(\mu_3$ -F)(BrF₅)₃]⁻. It was obtained in the reaction of CsF with an excess of BrF₅. By variation of the counter ion using KF or RbF instead of CsF as a starting material, we did not succeed in the isolation of other compounds containing fluoride-bridged fluoridobromate(v) anions, as of yet, but obtained the corresponding $[BrF_6]$ ⁻ salts. Which is salted the corresponding $[BrF_6]$ salts. Heading the salted the corresponding $[BrF_6]$

By introducing the sterically more demanding $[NMe_4]^+$ ion as a counterion, the reaction favors the formation of the bulkier $[(\mu_4\text{-F})(BrF_5)_4]^-$ ion instead of the smaller $[(\mu_3\text{-F})(BrF_5)_3]^-$ anion that is formed in the Cs compound. Here we present the first interhalogen compound, $[NMe_4][Br_4F_{21}]\cdot BrF_5$, in which a central fluorine atom is coordinated simultaneously by four halogen fluoride molecules forming a molecular $[(\mu_4\text{-F})(BrF_5)_4]^-$ anion. The compound was characterized by single-crystal X-ray diffraction and Raman spectroscopy and was investigated by quantum-chemical methods.

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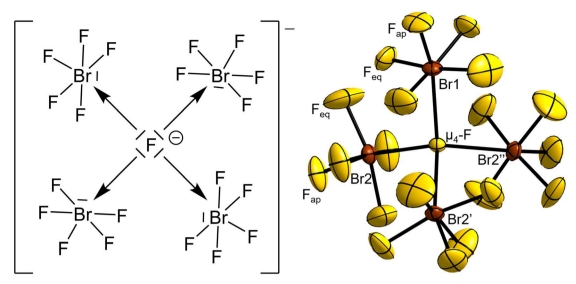


Fig. 1 Valence structural formula of the $[(\mu_4-F)(BrF_5)_4]^-$ anion and its molecular structure in the salt $[NMe_4][Br_4F_{21}] \cdot BrF_5$. Displacement ellipsoids are shown at 50% probability level at 100 K. Symmetry-equivalent, disordered F atoms bound to Br1 are omitted for clarity. Selected bond lengths (Å) and angles (°): μ_4 -F-Br1 2.474(3), μ_4 -F-Br2 2.5447(14), Br1-F $_{ap}$ 1.668(8), Br2-F $_{ap}$ 1.696(3), Br1-F $_{eq}$ 1.685(13) to 1.864(16), Br2-F $_{eq}$ 1.743(5) to 1.783(4), Br1- μ_4 -F-Br2 111.01(7), Br2- μ_4 -F-Br2 107.89(7), μ_4 -F-Br1-F $_{ap}$ 166.6(5), μ_4 -F-Br2-F $_{ap}$ 160.26(18). Symmetry codes: 'y, z, x, "z, x, y.

Results and discussion

Synthesis

The compound tetramethylammonium μ_4 -fluorido-tetrakis(pentafluoridobromate(v))—bromine pentafluoride(1/1), [NMe_4] [Br_4F_{21}] \cdot BrF_5, was obtained from the reaction of [NMe_4][BrF_6] with an excess of BrF_5. Therefore, [NMe_4][BrF_6] was cooled to -196 °C, BrF_5 was distilled onto the solid, and the reaction mixture was allowed to warm up to room temperature giving a colorless solution. The solution was stored at -36 °C. After several hours, colorless cube-shaped single crystals of [NMe_4] [Br_4F_{21}] \cdot BrF_5 had formed. The reaction can be described with (Eq. 1).

$$[NMe_4][BrF_6] + 4BrF_5 \xrightarrow{BrF_5, -36\,{}^{\circ}C} [NMe_4][Br_4F_{21}] \cdot BrF_5 \ \ (Eq. \ 1)$$

Crystals of the title compound are not stable in ${\rm BrF}_5$ solution at room temperature and decompose to the starting materials. If crystals are warmed slightly above room temperature in the absence of excess ${\rm BrF}_5$, then they decompose under evolution of brown vapors of ${\rm Br}_2$. The left-overs were not characterized further.

When reacting [NMe₄]F directly with an excess of neat BrF₅, a very powerful oxidant, the utmost care must be taken. The reaction is hardly controllable and often results in explosions even at temperatures as low as -196 °C (see Fig. S1†). The metathesis reaction between Cs[BrF₆] and [NMe₄]F, as reported by Christe and coworkers,²⁴ however, offers an elegant and dependable method for obtaining [NMe₄][BrF₆], see (Eq. 2).

$$Cs[BrF_6] + [NMe_4]F \xrightarrow{MeCN} [NMe_4][BrF_6] + CsF \downarrow$$
 (Eq. 2)

[NMe₄][BrF₆] is stable at room temperature, not shock sensitive,²⁴ and did not exhibit any violent reaction tendencies

when exposed to BrF5. Further details on the synthesis are given in the ESI.†

Crystal structure of [NMe₄][Br₄F₂₁]·BrF₅

Single-crystal X-ray structure determination showed [NMe₄] [Br₄F₂₁]·BrF₅ to crystallize in the cubic space group $P2_13$ (no. 198, $b^{27}a^5$, cP192) with the lattice parameter a=13.389(4) Å, V=2400(2) Å³, Z=4 at T=100 K. The structure consists of [(μ_4 -F)(BrF₅)₄]⁻ fluoridobromate(v) anions (Fig. 1), [NMe₄]⁺ cations and a BrF₅ molecule of crystallization per formula unit.

The crystal structure presents a borderline scenario between twinning and/or disorder affecting the cation, a BrF $_5$ unit of the anion, and the BrF $_5$ molecule of crystallization. These three reside on the threefold rotation axis of the space group and are therefore disordered by symmetry. A corresponding reduction of symmetry by eliminating the threefold rotation axis leads to the orthorhombic space group $P2_12_12_1$ and permits a crystallographic description of the structure as a three-component twin. However, the cubic unit cell metric results in too few symmetry-independent reflections to refine the structure properly in the orthorhombic crystal system and to construct a satisfying structural model. Therefore, the choice of the cubic space group $P2_13$ with a description of the disorder lead to the clearly superior structure model. Further crystallographic details are given in the ESI.†

The central μ_4 -F atom (4a, .3.) of the anion is surrounded tetrahedron-like by BrF $_5$ moieties. One BrF $_5$ moiety is 2.474(3) Å distant from the μ_4 -F atom with its Br atom residing on a 4a position with 3. symmetry, and therefore is affected by disorder over three positions. The remaining three symmetry-equivalent BrF $_5$ units reside on the 12a position (site symmetry 1), with a μ_4 -F-Br distance of 2.5447(14) Å.

For comparison, the observed distances for the two $\mu_3\text{-}$ bridged fluoridohalogenate(v) anions $[Br_3F_{16}]^{-14}$ and $[I_3F_{16}]^{-16}$

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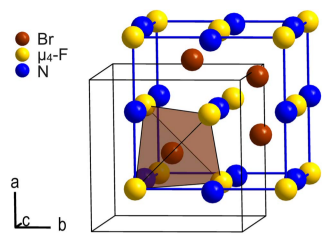


Fig. 2 Section of the crystal structure of $[NMe_4][Br_4F_{21}] \cdot BrF_5$ showing the arrangement of the μ_4 -F atoms of the anions (yellow), the N atoms of the $[NMe_4]^+$ cations (blue), and the Br atoms of the BrF $_5$ molecules of crystallization (brown) according to the MgAgAs structure type. The pseudo-face centered cubic unit cell is shown in blue. It is shifted by +(1/4, 1/4, 1/4) with respect to the origin of the original unit cell shown in black. Atoms are shown as spheres with arbitrary radii.

are almost equal with those present here, with X– μ_3 -F distances of 2.462(2) Å for X = Br and 2.477 to 2.514 Å for X = I, respectively. The Br– μ_4 -F–Br angles of the $[Br_4F_{21}]^-$ anion are 3 × 111.01(7)° and 3 × 107.89(7)°, showing a small distortion from the ideal tetrahedral angle with 109.5°.

The atomic distances of the Br to the terminally bound F atoms within the BrF5 units closely agree with those determined for pure BrF5 at 100 K.23 The bond lengths between Br2 and the equatorial F atoms, Br2-F_{eq}, range from 1.743(5) to 1.783(4) Å and are therefore essentially similar within their tripled standard uncertainties with those in pure BrF5 that range from 1.744(3) to 1.779(3) Å. The bond length of the apical Br-F bond, Br2-F_{ap}, is slightly longer with 1.696(3) Å compared to 1.686(2) Å for the one in pure BrF5, however, there is an overlap of the two bond lengths considering their tripled standard uncertainties. For the disordered BrF₅ unit, the Br1-F_{ap} bond length is 1.668(8) Å and the Br1-F_{eq} bond lengths range from 1.685(13) to 1.864(16) Å. This broader range and the larger standard uncertainties are caused by the disorder mentioned above. From a chemical point of view, the intramolecular atomic distances and angles of the BrF₅ units should differ only insignificantly among themselves, which is also confirmed by our quantumchemical calculations (see below).

The $F_{ap}\text{-Br-}F_{eq}$ angles of the ordered BrF_5 units of the anion lie in the range from $83.1(3)^\circ$ to $84.7(3)^\circ$, while for the disordered BrF_5 unit and BrF_5 molecule of crystallization they are observed around 83° to 85° . As observed for pure BrF_5 , the Br atoms are located slightly below a virtual plane spanned by the four surrounding F_{eq} atoms by 0.126(7) Å in the case of Br1 and 0.190(3) Å for Br2, respectively. The $\mu_4\text{-F-Br-}F_{ap}$ angles are $166.6(5)^\circ$ for Br1 and $160.26(18)^\circ$ for Br2 and can be interpreted as an indication for the extent of the stereochemical activity of the free electron pair located on the Br atoms. The more the $\mu_4\text{-F-Br-}F_{ap}$ angle deviates from 180° , the stronger the

stereochemical influence of the lone pair. In the octahedral $[BrF_6]^-$ anion, no stereochemical effect of the lone pair is observed, $^{16,23,25-27}$ while it is also present in the $[(\mu_3\text{-F})(BrF_5)_3]^-$ anion, as evidenced in the $\mu_3\text{-F-Br-F}_{ap}$ bond angle of $165.0(3)^\circ$, also deviating from 180° . The stereochemical effect of the free electron pair is even more apparent in the analogous $[(\mu_3\text{-F})(IF_5)_3]^-$ anion, whose $\mu_3\text{-F-I-F}_{ap}$ angles are in the range from 141.4° to 149.0° . The stereochemical effect of the free electron pair is even more apparent in the analogous $[(\mu_3\text{-F})(IF_5)_3]^-$ anion, whose $\mu_3\text{-F-I-F}_{ap}$ angles are in the range from 141.4° to 149.0° .

Except for Coulomb interaction between the [NMe₄]⁺ cations and the [Br₄F₂₁] anions we observe, due to the disorder, no additional intermolecular interactions such as hydrogen bonds in the crystal structure of [NMe₄][Br₄F₂₁]·BrF₅. C-H···F hydrogen bonds are however likely present as H···F distances are overall in the proper range.28 Each [Br4F21] anion is octahedrally surrounded by [NMe₄]⁺ cations, and vice versa. Thus, cations and anions are arranged in a cubic close packing, respectively, in which all octahedral voids are occupied by counterions. The ionic part of the crystal structure is therefore related to the NaCl structure type. The BrF5 molecules of crystallization occupy half of its tetrahedral voids and are arranged tetrahedron-like. So, the overall arrangement of these building blocks corresponds to the intermetallic half-Heusler compound and structure type MgAgAs. Fig. 2 shows the pseudo-face centered cubic unit cell in blue and the arrangement of the N atoms of the cations, the μ_4 -F atoms of the anions, and the Br atoms of the BrF5 molecules of crystallization corresponding to the MgAgAs structure type.

Quantum-chemical calculations on the crystal structure of $[NMe_4][Br_4F_{21}] \cdot BrF_5$

The lattice parameters and atomic positions of the crystal structure were optimized by using hybrid density functional methods (DFT-PBE0/TZVP level of theory, see ESI† for the computational details). Since the structure model in space group P2₁3 derived from the X-ray structure determination includes disordered atoms, the structure was optimized in the orthorhombic subgroup $P2_12_12_1$ so that a description of the structure without disorder was possible. A harmonic frequency calculation for the optimized structure confirmed it as a true local minimum on the potential energy surface. The optimized lattice parameters deviate less than 2.3% from the lattice parameter determined by X-ray diffraction and bond lengths and angles are in accordance with the experiment. The calculated four μ_4 -F-Br distances of the $[Br_4F_{21}]^-$ anion are in the range of 2.456 to 2.577 Å compared to the experimentally determined distances of 2.474(3) and 2.5446(14) Å. The calculated Br-F_{ap} bond lengths range from 1.704 to 1.708 Å, while the calculated $Br-F_{eq}$ bond lengths are in the range from 1.777 to 1.798 Å. Compared to the experimentally determined bond lengths, the agreement is significantly better for the BrF₅ units that are not affected by disorder, as expected. For these, the Br- F_{ap} distance was determined to be 1.696(3) Å and the Br- F_{eq} distances are 1.742(5) to 1.783(4) Å. However, the experimentally determined Br-F distances for the disordered parts, which are 1.668(8) Å for the Br-F_{ap} bond and 1.685(13)-1.864(16) Å for the Br-F_{eq} bonds, are of lower precision and therefore spread

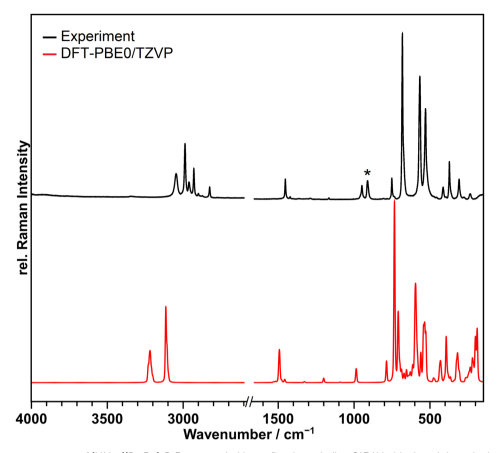


Fig. 3 Observed Raman spectrum of [NMe₄][Br₄F₂₁]·BrF₅ suspended in perfluorinated oil at 213 K in black and the calculated spectrum at the DFT-PBE0/TZVP level of theory in red. No bands were observed or calculated in the range from $1600-2700 \, \mathrm{cm}^{-1}$, so it was omitted in the Figure. The band marked with an asterisk originates form BrO₂F, which is a hydrolysis product formed during the Raman measurement.²⁹ The calculated spectrum in red shows many bands not observed in the recorded one. This is because of the lower resolution of the recorded spectrum.

around the calculated values. A comparison of other calculated and experimentally determined atomic distances and angles is available in Tables S6 and S7.†

Vibrational spectroscopy

A Raman spectrum of the compound was recorded at 213 K and is shown in Fig. 3 in black in comparison to the calculated spectrum at the DFT-PBE0/TZVP level of theory for the solid-state structure in red.

The bands above 700 cm $^{-1}$ belong to the [NMe $_4$] † cation, and those below 700 cm $^{-1}$ to the anion and the BrF $_5$ molecule of crystallization, however, the latter two could not be clearly separated. The bands from 529 to 682 cm $^{-1}$ are due to the Br–F stretching vibrations, those from 235 to 413 cm $^{-1}$ are assigned to wagging, scissoring and umbrella modes. Table S8 \dagger contains the observed and calculated band positions and their detailed assignments.

With the exception of two points the recorded spectrum agrees with the calculated one in which the frequencies are slightly overestimated as anharmonic effects are not considered.³⁰ Additional bands not reproduced by the frequency calculations are observed in the range of 2800–2970 cm⁻¹. However, these are assignable to second-order bands of the

[NMe $_4$]⁺ ion. 31,32 The band marked with an asterisk at 911 cm $^{-1}$, does not originate from the title compound and also increased in intensity during the measurement. It can be assigned to the symmetric $\nu(Br-O)$ stretching vibration of BrO_2F , which is obtained as a hydrolysis product of the title compound. 29 The other bands expected for BrO_2F are significantly weaker in intensity and overlap with the bands of the title compound, so that they cannot be observed. Contact of the sample with the atmosphere causing hydrolysis could not be completely excluded during the transfer of the cold sample into the spectrometer. Additional details of the Raman setup and sample preparation are given in the ESI.†

Intrinsic bonding orbital analysis

Quantum-chemical calculations on the isolated $[(\mu_4\text{-F})(\text{BrF}_5)_4]^-$ anion and its hypothetical homologues $[(\mu_4\text{-F})(\text{ClF}_5)_4]^-$ and $[(\mu_4\text{-F})(\text{IF}_5)_4]^-$ were conducted at the DFT-PBE0/def2-TZVP level of theory with a consecutive analysis of the bonding situation using intrinsic bonding orbitals (IBOs).³³ The structures of the molecular anions $[(\mu_4\text{-F})(XF_5)_4]^-$, with X = Cl-I, were all optimized for the gas phase at 0 K in point group S_4 and were confirmed to be true local minima on the potential energy surfaces by means of harmonic frequency calculations. The

Fig. 4 Selected resonance valence structural formulae of the $[(\mu_4-F)(BrF_5)_4]^-$ anion. Numbers next to the atoms indicate their calculated partial charges. The result of our IBO analysis shows that the one on the left with an essentially ionic Br-F bond is to be preferred.

calculated μ_4 -F-Br bond length in the $[(\mu_4$ -F)(BrF₅)₄]⁻ anion is 2.576 Å and agrees with the experimentally determined ones of $3 \times 2.5447(14)$ and 2.474(3) Å. The IBO analysis showed that none of the Br atoms contribute to the μ_4 -F-Br bond with more than 1%, which indicates a purely ionic bonding situation, and suggests the description of the μ_4 -F atom as a F⁻ anion. For comparison: In a gas-phase NaF molecule, in which the bond should be highly ionic, the contribution of the F atom is 96%. In the H₂ molecule, which has a purely covalent bond, the contribution of each H atom is 50%. The partial charge of the μ_4 -F atom is -0.77 e, which is significantly more negative than that of the F atoms in a BrF5 unit with partial charges between -0.41 and -0.51 e. The μ_4 -F atom is even more ionic than the μ_3 -F atom in the $[(\mu_3$ -F)(BrF₅)₃]⁻ anion with -0.73 e.¹⁴ The partial charges of the μ_4 -F atoms in the hypothetical anions [(μ_4 -F)(ClF₅)₄]⁻ and [(μ_4 -F)(IF₅)₄]⁻, calculated to be -0.76 and -0.77e, respectively, are similar to the one observed for $[(\mu_4$ F)(BrF₅)₄]⁻. The partial charges of the Br atoms of the $[(\mu_4$ $F(BrF_5)_4$ anion and the previously reported $[(\mu_3-F)(BrF_5)_3]^$ anion are almost the same with +2.38 e and +2.37 e, respectively.14

The high negative partial charge of the μ_4 -F atom as well as the high positive partial charge of the Br atoms of the $[(\mu_4$ -F)(BrF₅)₄]⁻ anion strengthen the suggestion of describing the μ_4 -F atom as an F⁻ anion and the μ_4 F-Br bonds as essentially ionic (see Fig. 4).

Conclusions

The compound $[NMe_4][Br_4F_{21}] \cdot BrF_5$ was obtained by reacting $[NMe_4][BrF_6]$ with an excess of BrF_5 . Surprisingly, its crystal structure is closely related to the intermetallic half-Heusler compound and structure type MgAgAs, as the cations, anions and BrF_5 molecules of crystallization are ordered as in MgAgAs. In the $[(\mu_4 \cdot F)(BrF_5)_4]^-$ anion, the $\mu_4 \cdot F$ atom is surrounded by four BrF_5 molecules in a tetrahedron-like shape. It is the first example of a F atom μ_4 -bridging to neither metal nor H atoms. IBO analyses of the anion showed that the covalence of the μ_4 -F-Br interaction is negligibly small and therefore the bond is best described as ionic. The recorded Raman spectrum of $[NMe_4]$

 $[Br_4F_{21}] \cdot BrF_5$ agrees with the quantum-chemically calculated one for the solid state. The tetrahedral $[(\mu_4\text{-F})(BrF_5)_4]^-$ and the propeller-shaped $[(\mu_3\text{-F})(BrF_5)_3]^-$ anions are so far the only known polynuclear anions of BrF_5 . We have currently no experimental evidence for the existence of the putative dinuclear $[(\mu\text{-F})(BrF_5)_2]^-$ anion.

Data availability

Upon request from the authors.

Author contributions

Martin Möbs: planning and conducting the experiments, main data acquisition and interpretation, quantum-chemical calculations and interpretation of the results, manuscript preparation. Tim Graubner: quantum-chemical calculations and interpretation of the results, manuscript preparation. Antti J. Karttunen: quantum-chemical calculations, CRYSTAL basis set development, manuscript preparation. Florian Kraus: project supervision, data interpretation, manuscript preparation.

Conflicts of interest

The authors declare no conflicts of interest.

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