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High-pressure observation of elusive iodoplumbic acid in different hydronium-hydrate solid forms†‡

Szymon Sobczak,^a Athena M. Fidelli,^b Jean-Louis Do,^b
George P. Demopoulos,^c Audrey Moores,^b Tomislav Friščić^{b,d} and
Andrzej Katrusiak^a

High-pressure and high-temperature isochoric crystallization combined with single-crystal X-ray diffraction revealed the proposed, but previously never demonstrated, hydronium acid hydrates of $[\text{PbI}_3]^-$, the closest forms of the elusive iodoplumbic acid. Depending on the pressure range, the reaction of PbI_2 and aqueous concentrated hydriodic acid under isochoric conditions in a diamond anvil cell held between 0.11 and 1.20 GPa produces two hydrated acids with compositions $[\text{H}_3\text{O}][\text{PbI}_3] \cdot n\text{H}_2\text{O}$ ($n = 3, 4$). Comprised of polymeric one-dimensional double-chain PbI_3^- anions, analogous to those seen in archetypal lead-halide perovskites such as RbPbI_3 and CsPbI_3 , these hydrates offer the first observation of the iodoplumbic acid progenitor of hybrid lead perovskites. We also reveal the 'hidden' polymorph of lead(II) iodide, adopting a three-dimensional $\text{Pb}-\text{I}$ bonded network, contrasting with the prototypic ambient-pressure layered PbI_2 structure.

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Introduction

The emergence of photovoltaic lead halide perovskites APbI_3 , where A = alkaline metal, ammonium or organoammonium cation has sparked extensive studies on the existence of iodoplumbic(II) acid, HPbI_3 .^{1,2} In most cases, lead halide perovskites are composed of polymeric PbI_3^- anions, such as NH_4PbI_3 ,³ $[\text{CH}_3\text{NH}_3]\text{PbI}_3$,^{4–9} or CsPbI_3 ,^{10–12} which can be synthesized *via* traditional¹³ as well as unconventional methods.^{14–16} The proposed HPbI_3 would be the simplest member of this compound class and the progenitor of hybrid, as well as inorganic lead(II) perovskites. There were several attempts to propose a computational prediction of the most probable HPbI_3 structure, which suggested that it forms a 3-dimensional perovskite structure of

space-group symmetry $\text{Pm}\bar{3}\text{m}$, consisting of the PbI_3^- -polyanionic framework encapsulating hydrogen inside the cubic cages.¹⁷ Whereas iodoplumbic acid was proposed as a precursor for the synthesis of hybrid perovskites in 2015 by Zhao *et al.*,¹⁸ the existence of an acid with composition HPbI_3 has remained controversial.^{19–28} The groups of Kanatzidis and of Hillebrecht^{29,30} have shown that the postulated HPbI_3 precipitate obtained from *N,N*-dimethylformamide (DMF) solutions of PbI_2 and hydriodic acid (HI) is actually the dimethylammonium hybrid perovskite $[\text{N}(\text{CH}_3)_2\text{H}_2][\text{PbI}_3]$. Moreover, Daub and Hillebrecht demonstrated that the reaction of PbI_2 with concentrated (57% w/w) aqueous HI can produce two forms of hydrated iodoplumbic acid.³⁰ One form exhibits the composition $[\text{H}_3\text{O}]_{2x}[\text{Pb}_{1-x}\text{I}_2] \cdot (2 - 2x)\text{H}_2\text{O}$ (**1**) ($x \approx 0.23$), and is based on two-dimensional (2-D) anionic CdI_2 -type sheets with approximate composition $[\text{Pb}_3\text{I}_8^{2-}]_n$ (Fig. 1a). The second reported form of hydrated iodoplumbic acid exhibits the composition $(\text{H}_3\text{O})_2\text{Pb}_3\text{I}_8 \cdot 6\text{H}_2\text{O}$ (**2**) (Fig. 1b), and is based on one-dimensional (1-D) polyanionic tapes of $[\text{Pb}_3\text{I}_8^{2-}]_n$.

Compound **2** was reported to be the first product of either crystallization of PbI_2 from concentrated aqueous HI, or of the gas–solid reaction between PbI_2 and HI vapours. When exposed to open air, **2** quickly transforms into **1**. Anions in both **1** and **2** are separated by layers of water molecules containing hydronium ions. Overall, these prior studies suggest that an inorganic acid based on the PbI_3^- anion does not exist, and that the only accessible forms of iodoplumbic acid are hydronium salts of the $\text{Pb}_3\text{I}_8^{2-}$ anion.

We now report the synthesis and observation of a hydronium salt of $[\text{Pb}_2\text{I}_6]^{2-}$ achieved under unconventional, isochoric con-

^aDepartment of Materials Chemistry, Faculty of Chemistry, Adam Mickiewicz University, Uniwersytetu Poznańskiego 8, 61-614 Poznań, Poland.
E-mail: katran@amu.edu.pl; https://twitter.com/szymon_sobczakhttps://twitter.com/DMCh_AMU

^bDepartment of Chemistry, McGill University, 801 Sherbrooke St West, Montreal, QC, H3H 0B8, Canada. E-mail: audrey.moores@mcgill.ca; <https://twitter.com/MooresResearch>https://twitter.com/JLD_Chemist<https://twitter.com/TomislavFriscic>

^cDepartment of Materials Engineering, McGill University, 801 Sherbrooke St West, Montreal, QC, H3A 0C5, Canada. E-mail: george.demopoulos@mcgill.ca

^dSchool of Chemistry, University of Birmingham, Edgbaston, Birmingham B15 2TT, UK. E-mail: t.friscic@bham.ac.uk

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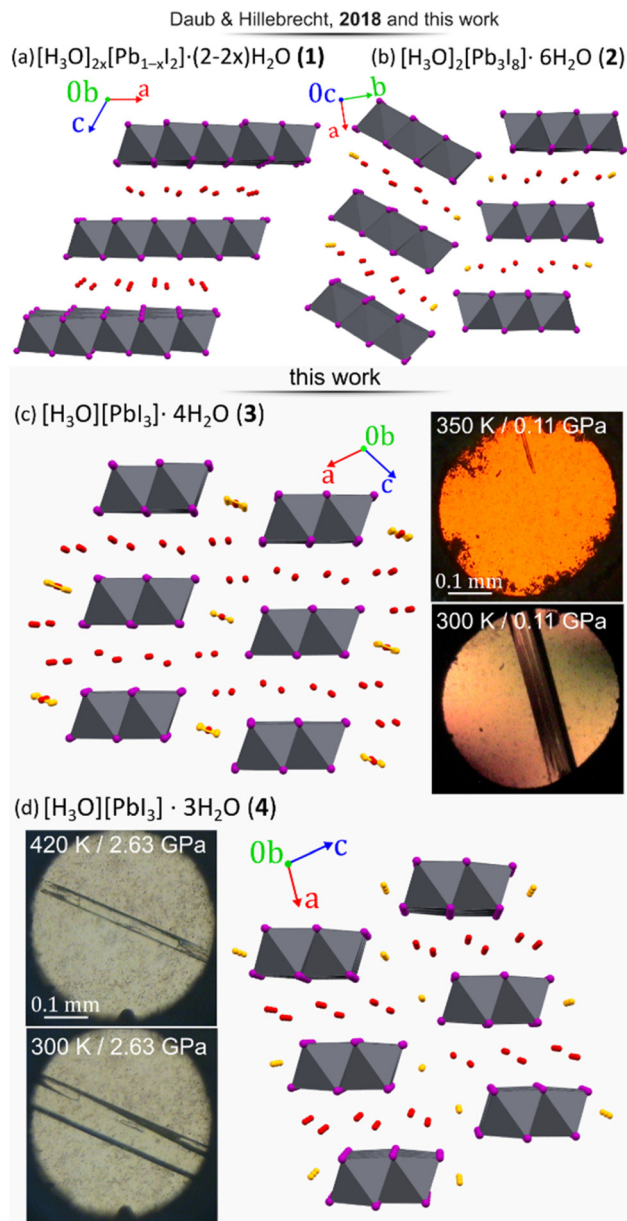


Fig. 1 Four forms of hydrated iodoplumbic acid obtained by reaction of PbI_2 and concentrated aqueous HI : (a) $[\text{H}_3\text{O}]_{2x}[\text{Pb}_{1-x}\text{I}_2] \cdot (2-2x)\text{H}_2\text{O}$ (**1**, $x \approx 0.23$); (b) $(\text{H}_3\text{O})_2\text{Pb}_3\text{I}_8 \cdot 6\text{H}_2\text{O}$ (**2**);³⁰ as well as new materials herein synthesized under high-pressure (*p*) and -temperature (*T*) conditions: (c) $(\text{H}_3\text{O})\text{PbI}_3 \cdot 4\text{H}_2\text{O}$ (**3**) and (d) $(\text{H}_3\text{O})\text{PbI}_3 \cdot 3\text{H}_2\text{O}$ (**4**). The proposed sites of H_3O^+ cations are marked in orange. Photographs show crystals **3** and **4** *in situ* grown under pressure.

ditions of elevated temperature and pressure. Specifically, we show that crystallization of PbI_2 from concentrated aqueous HI provides, at pressure above 0.11 GPa, access to hydronium of 1-D double-chain polyanions with composition PbI_3^- . Based on the chemical and structural composition of the anion, which are identical to those in RbPbI_3 , CsPbI_3 etc., and the absence of any ammonium or metal cations in the system, the herein reported structures can be considered a significant step towards the observation and understanding of iodoplumbic acid, a historically elusive entity.

Results and discussion

With HPbI_3 apparently inaccessible by solution techniques, our study has focused on less conventional reaction environments based on introduction of mechanical energy in the form of either ball milling (mechanochemistry)^{31–33} or by high-pressure chemistry. Our first exploration of reactivity between PbI_2 and concentrated aqueous hydroiodic acid HI (aq) was done mechanochemically, by milling of the two components in the stoichiometric ratio of $\text{PbI}_2/\text{HI} = 1.5 : 1$ (corresponding to the 3 : 8 stoichiometric ratio of Pb to I, respectively). Milling produced a bright yellow powder that, upon powder X-ray diffraction (PXRD) analysis, matched the previously reported **1** (see ESI†). Increasing the amount of HI (aq) in the milling reaction to produce a 1 : 3 respective stoichiometric ratio of PbI_2 and HI led to the formation of **2** in a pure form (see ESI†). Consistent with previous work,³⁰ upon standing in air **2** slowly transforms into **1** and, subsequently, into the orange solid PbI_2 .

A different approach to introduce mechanical energy to a reaction is in the form of hydrostatic pressure in a diamond anvil cell (DAC),^{34–40} allowing the exploration of otherwise inaccessible thermodynamic coordinates and formation of new products.^{39–42} As a reactor, the DAC represents an almost perfect closed system confining the reaction to the volume of $\sim 0.02 \text{ mm}^3$ between two diamond culets and a steel gasket.⁴¹ For each of the high-pressure reactions, a saturated solution of PbI_2 in HI (aq) was loaded in the DAC and isothermally compressed at 297 K. At 0.11 GPa, a polycrystalline mass precipitated that, upon subsequent isochoric recrystallization led to a colorless elongated prism-shaped crystal suitable for structure determination by single-crystal X-ray diffraction (SCXRD). After increasing pressure to 0.20 GPa, the unit-cell dimensions were determined (see ESI Table S1†) and the next SCXRD measurement at 0.5 GPa revealed a novel hydrated hydronium salt of composition $(\text{H}_3\text{O})\text{PbI}_3 \cdot 4\text{H}_2\text{O}$ (**3**). The structure of **3** consists of 1-D anionic PbI_3^- tapes of edge-sharing PbI_6 -octahedra running along the crystallographic *b*-axis (Fig. 1c). The PbI_6 -octahedra in the PbI_3^- polyanion are distorted, exhibiting four different lengths of Pb–I bonds around each metal ion: 3.046 (2), 3.174(6) (twice), 3.250(6) (twice), and 3.3663(19) Å. These Pb–I bond length distances are very similar to those observed in diguanidinium tetraiodoplumbate⁴³ and other 6-coordinated lead(II) complexes (see ESI, Fig. S10†), confirming the retention of the Pb^{2+} oxidation state. Importantly, the PbI_3^- anionic tapes in **3** are identical to those in $\text{NH}_4\text{CdCl}_3 \cdot 2\text{H}_2\text{O}$, hybrid and lead iodide perovskites CsPbI_3 , RbPbI_3 , as well as related hydrates $\text{NH}_4\text{PbI}_3 \cdot 2\text{H}_2\text{O}$, $\text{CsPbI}_3 \cdot 2\text{H}_2\text{O}$, and $\text{CH}_3\text{NH}_3\text{PbI}_3 \cdot \text{H}_2\text{O}$.^{44–49} The structure of the anion makes compound **3** the first example of a direct hydronium-based acid analogue of these well-known lead perovskite solids. Crystallographic parameters for **3** are distinct from those of previously reported³⁰ **1** or **2** (Fig. 1c and Table 1).

Each $[\text{PbI}_3]_n^-$ polyanion requires a counter hydronium cation for the charge balance. Although the strong scattering of X-rays by heavy lead and iodine atoms hinders the precise

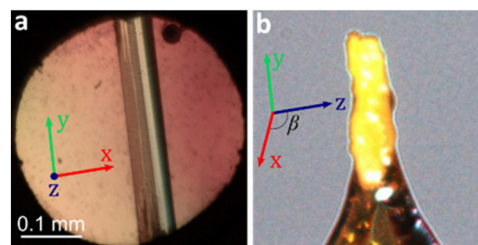
Table 1 Selected crystallographic data for compounds **1–4** and β -PbI₂. The structure of **1** was re-determined at ambient conditions on a crystal that was obtained by single-crystal-to-single-crystal decomposition of a crystal of **3** and recovered from the DAC

Compound	1 ^a	2 ³⁰	3	4	β -PbI ₂
Formula	[H ₃ O] _{0.40} [Pb _{0.6} I ₂] $\cdot$ 1.6H ₂ O	(H ₃ O) ₂ Pb ₃ I ₈ $\cdot$ 6H ₂ O	(H ₃ O)PbI ₃ $\cdot$ 4H ₂ O	(H ₃ O)PbI ₃ $\cdot$ 3H ₂ O	PbI ₂
<i>P</i> (GPa)	0.0001	0.0001	0.5	2.63	2.05
<i>T</i> (K)	300	300	300	300	320
Space group	<i>C2/m</i>	<i>Pbam</i>	<i>I2/m</i>	<i>P2₁/m</i>	<i>C2/m</i>
<i>a</i> (Å)	7.8946(10)	10.033(3)	16.205(2)	9.57(3)	14.06(4)
<i>b</i> (Å)	4.5598(3)	30.126(7)	4.5170(1)	4.513(3)	4.4560(12)
<i>c</i> (Å)	11.1985(18)	4.5610(10)	17.184(14)	12.98(11)	10.540(6)
β (°)	118.023(19)	—	111.50(4)	95.9(6)	93.08(12)
<i>V</i> (Å ³)	355.859	1378.66(6)	1170.3(10)	558(5)	654.9(17)
<i>D</i> _{calc} (g cm ^{−3})	4.638	4.290	3.666	3.883	7.013

^aThe herein determined structure [H₃O]_{2x}[Pb_{1−x}I₂] $\cdot$ (2 − 2x)H₂O, *x* = 0.20(2), is analogous to that previously reported³⁰ with *x* = 0.23.

location of hydrogen atoms in X-ray diffraction, the location of oxygen atoms and their shortest distances can discriminate the corresponding water molecules and hydronium ions. It is known that the hydronium cations form with iodide anions strong, charge-assisted O $\cdots$ I bonds, significantly shorter than those involving water molecules. In **3**, the [Pb₂I₆]^{2−} polyanions are separated by tapes of hydrogen-bonded water molecules (Fig. 1c and 2b) connected into an extended honeycomb-like structure through O–H $\cdots$ O hydrogen bonds with O $\cdots$ O distances of 2.63(4), 2.68(6) and 2.79(7) Å. The tapes propagate in the crystallographic *b*-direction and are periodically interrupted by iodoplumbate(n) anions, forming I $\cdots$ O contacts that are significantly shorter than the 3.54 Å non-bonding distance expected from the sum of van der Waals radii⁵⁰ of O (1.50 Å) and I (2.04 Å) atoms, indicating the formation of the charge-assisted O–H $\cdots$ I bonds. The structure of **3** also exhibits 1-D arrays of oxygen atoms (with O $\cdots$ O distances of 2.41(6) Å) separate from the hydrogen-bonded honeycomb tapes, which are located between pairs of polymeric PbI₃[−] anions (Fig. 2b). At 0.50 GPa, these oxygen atoms are situated at notably short I $\cdots$ O distances of 3.2(2) Å, that are commensurate with charge-assisted⁵¹ O–H $\cdots$ I hydrogen bonding, tentatively indicating the location of H₃O⁺ ions.

While increasing the pressure up to 1.20 GPa does not affect the crystals of **3**, releasing the pressure to 0.1 MPa quickly leads to their transformation into **1** (Fig. 3). This

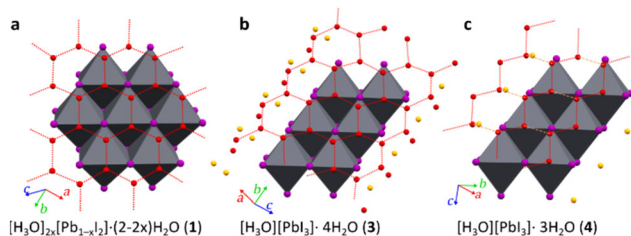
**Fig. 3** Single crystal of **3**: (a) as grown in the DAC and (b) after being recovered to ambient conditions and transforming to **1**, mounted on a nylon loop. Crystal axes are indicated.

chemical transformation takes place in a single-crystal-to-single-crystal manner, as shown by X-ray diffraction on the crystal recovered from the DAC (see ESI†), which revealed a clear matrix relationship between the lattices for the starting crystal structure of H₃OPbI₃ $\cdot$ 4H₂O (**3**) and the daughter phase **1**:

$$\begin{pmatrix} -1/3 & 0 & 1/3 \\ 0 & -1 & 0 \\ 2/3 & 0 & 1/3 \end{pmatrix} \begin{pmatrix} a_3 \\ b_3 \\ c_3 \end{pmatrix} = \begin{pmatrix} a_1 \\ b_1 \\ c_1 \end{pmatrix},$$

where *a*₁, *b*₁, *c*₁ and *a*₃, *b*₃, *c*₃ are sets of unit-cell vectors for **1** and **3**, respectively (Table 1).

The quality of the crystal recovered to 0.1 MPa permitted the SCXRD measurement of lattice dimensions and structure refinement, which revealed monoclinic symmetry of space group *C2/m* (Table 1, Fig. 2a, also ESI†). The structure was found to be highly pseudo-symmetric (for details see ESI†) and similar to the trigonal structure previously reported for **1**.³⁰ The two determinations of this layered [H₃O]_{2x}[Pb_{1−x}I₂] $\cdot$ (2 − 2x)H₂O structure are consistent (Fig. 1a and 2a), except for a somewhat lower *x* = 0.20(2) value resulting from our least-squares refinement, compared to *x* = 0.23.³⁰ The difference, we believe, indicates the possibility of **1** to adopt a wider range of Pb : I stoichiometric compositions. The observed highly topotactic transformation⁵² from **3** to **1** requires a transition from 1-D to 2-D polyanions, in which some of the water and HI molecules leave the structure, probably by diffusion, while the

**Fig. 2** Fragments of crystal structures of (a) **1** herein re-determined³⁰ from a crystal obtained by transformation of **3** at 0.1 MPa; (b) **3**; and (c) **4**, all viewed perpendicular to the honeycomb pattern of water molecules (red-dotted lines, orange-dashed lines in **4** mark the longest O $\cdots$ O distances). The oxygen-atom sites suggested for H₃O⁺ ions are shown in orange.

edge-sharing connectivity of PbI_6 -octahedra is preserved in both materials.

Above 1.2 GPa and above 420 K, a clearly distinct pink-coloured crystalline material (Fig. 4a and b) different from compound **3**, is formed. Subsequent SCXRD at 320 K and pressures above 2.05 GPa revealed a 'hidden' polymorph of PbI_2 , herein termed $\beta\text{-PbI}_2$. This novel form of $\beta\text{-PbI}_2$ (Table 1, Fig. 4c, d, also ESI†) of the monoclinic space group $C2/m$ and unprecedented for PbI_2 polymorphs exhibits a 3-D framework of alternating six- and seven-coordinated Pb^{2+} cations bridged by iodide ions, in stark contrast to the well-known 2-D layered structure with six-coordinated Pb^{2+} cations herein termed $\alpha\text{-PbI}_2$.^{53,54} The $\beta\text{-PbI}_2$ displays a rare property of reverse solubility, as we observed the growth of $\beta\text{-PbI}_2$ crystals on increasing the temperature, and their dissolution on cooling of the DAC. The isochoric conditions in the DAC imply that the increased temperature results either in the increased pressure (for the overall positive thermal expansion of all components in the DAC chamber) or the pressure drops (for the overall negative thermal expansion). According to our knowledge on numerous high-pressure crystallizations reported in the literature, the temperature increase always enhanced dissolution of the solid compounds. There are very few compounds becoming less soluble with increasing temperature (e.g. Li_2SO_4); likewise, few compounds for which the reverse solubility with the increase of pressure were reported. The formation of $\beta\text{-PbI}_2$ is consistent with high-pressure effect increasing the coordination numbers due to stronger compression of anions than cations.^{42,55,56} Recently, the high-pressure phases (phase II and III) of the 2H-type PbI_2 polytype, first identified by Bridgman *via* isothermal compression,^{58,59} were characterized.⁵⁷ Phase II (space group $P3m1$) crystallizes above 0.58 GPa as a two-dimensional 4H polytype, while in phase III (orthorhombic $Pnma$), stable above 2.6

GPa, each Pb atom is coordinated by 9 I-atoms similarly as in the orthorhombic PbCl_2 -structure.⁵⁷ However, both phases II and III contrast with the complex structure of $\beta\text{-PbI}_2$, which requires *in situ* recrystallization and cannot be obtained by cooling/heating or compressing another phase; hence $\beta\text{-PbI}_2$ is referred to as a 'hidden phase'. In this respect, PbI_2 is similar to imidazole where the β phase can be accessed only by high-pressure recrystallization.⁶⁰

Another crystalline phase (**4**) was obtained either by recrystallizing **3** above 1.2 GPa or by spontaneous recrystallization of $\beta\text{-PbI}_2$ in the DAC below 350 K (Fig. 1d, also ESI†). Below 1.2 GPa, at 320 K, pink $\beta\text{-PbI}_2$ dissolves and colourless needle-like crystals of **4** appear. Recrystallization by mild temperature oscillation produced a diffraction-quality single crystal of **4** (Fig. 1d and Table 1).

Compound **4** was found to exhibit the formula $(\text{H}_3\text{O})\text{PbI}_3 \cdot 3\text{H}_2\text{O}$, again based on polymeric anions with edge-sharing PbI_6 -octahedra identical to those in **3** and a variety of lead halide perovskites, but with a lower content of crystallization water. The lower content of water in **4** compared to **3** is consistent with shorter $\text{I}\cdots\text{I}$ contacts at higher pressure. In the case of compound **4**, also four distinct lengths of Pb–I bonds are present in the anions: 2.81(4), 3.186(12) (twice), 3.232(11) (twice), and 3.34(4) Å. These bond lengths are consistent with lead again adopting the Pb^{2+} oxidation state (Fig. S10†), implying the anion composition PbI_3^- .

In **4**, oxygen atoms of water molecules form ribbons (about 10 Å wide). Within the ribbons shorter hydrogen bonds [$\text{O}\cdots\text{O}$ distances of 2.73(9) Å] are arranged into three 1-D zigzag chains, interconnected by weaker hydrogen bonds [$\text{O}\cdots\text{O}$ distance of 3.1(2) Å] into a strongly distorted honeycomb motif. The ribbons separate the adjacent pairs of PbI_3^- polyanions. Additionally, there are also oxygen atoms not involved in any $\text{O}\cdots\text{O}$ contacts commensurate with hydrogen bonds, but they are each close to iodine atoms of PbI_3^- anions, with $\text{O}\cdots\text{I}$ distances of 3.187(12) (twice), 3.232(11) (twice), 2.80(4) and 3.33(4) Å, at a pressure of 2.63 GPa and a temperature of 300 K. As indicated above, the hydronium cations and water molecules (Fig. 2c) were discriminated according to the $\text{O}\cdots\text{PbI}_3^-$ distances, shorter for the charge-assisted $\text{OH}^+\cdots\text{I}$ bonds.

The appearance of the herein described series of structures comprising the α - and β -forms of PbI_2 , as well as hydrated iodo-plumbic acids **1–4** can be rationalized through the interplay of effects related to the intercalation of $\text{HI}(\text{aq})$ and to high-pressure conditions. Specifically, the observation of different structures at different conditions outlines several stability regions in the preference p - T diagram (Fig. 5), where the low-pressure end-member is $\alpha\text{-PbI}_2$ with a 2-D layer structure, and the high-pressure end-member is $\beta\text{-PbI}_2$, in which some of the Pb^{2+} cations become 7-coordinated to form a 3-D network.

All members of the series, exhibiting either 1-D, 2-D or 3-D structures, contain the common motif of edge-sharing PbI_6^- octahedra (Fig. 1 and 2), based on Pb–I bonds that are by far and large the least compressed elements constituting the scaffolds of the structures (Table 1). In contrast, the $\text{I}\cdots\text{I}$ contacts between PbI_2 sheets or the iodo-plumbate(II) anions are

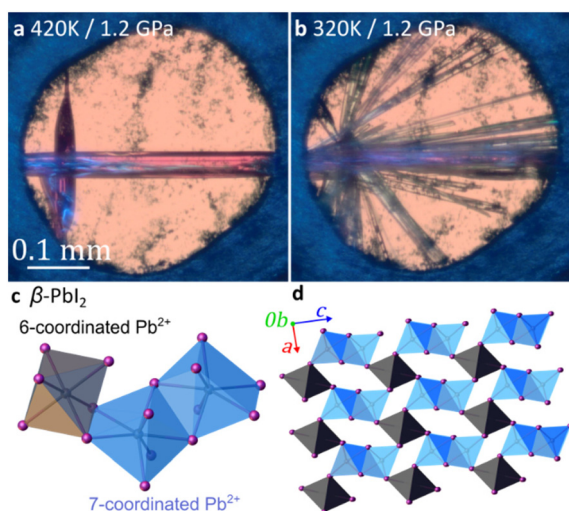


Fig. 4 The 3-D network polymorph of PbI_2 . Pink single crystals of $\beta\text{-PbI}_2$ at 1.2 GPa and: (a) 420 K and (b) 320 K covered by a bundle of needle crystals of **3**. Views of the $\beta\text{-PbI}_2$ structure: (c) 6- and 7-coordinated Pb^{2+} cations shown in grey and blue, respectively, and (d) the 3-D network viewed along the crystallographic b -axis.

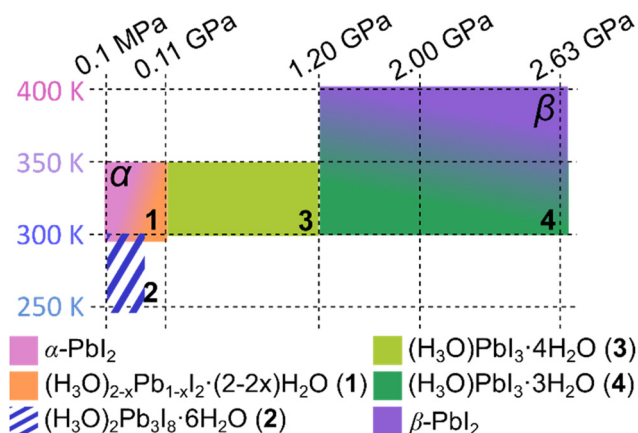


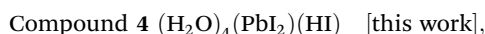
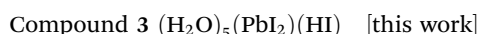
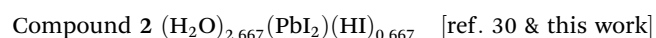
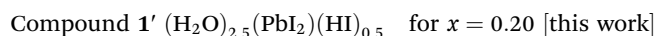
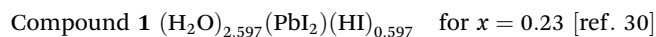
Fig. 5 The p/T preference diagram for the PbI_2 and HI(aq) system, with the end members being the previously known 2-D $\alpha\text{-PbI}_2$ and the herein reported 3-D $\beta\text{-PbI}_2$. Acids **1** and **2** are obtained by milling and vapor-solid reactions, whereas **1**, **3**, **4** and $\beta\text{-PbI}_2$ result from high-pressure synthesis in the DAC.

expected to be considerably weaker and most likely to be affected by pressure, temperature and overall chemical environment. The structures **1–4** can be seen as resulting from the $\alpha\text{-PbI}_2$ structure through intercalation of water and H_3O^+ from HI(aq) , resulting in $\text{O-H}\cdots\text{I}$ bonds between polyanionic sheets and tapes, which prevent their contacts involving iodine atoms.

The observation of compounds **1–3** at pressures up to 1.2 GPa occurs by progressive insertion of hydronium ions from HI(aq) into the PbI_2 structure, leading to the formation of an anion with PbI_3^- composition. At 1.2 GPa a partial desorption of water is observed, leading to the formation of compound **4** that maintains the structure and composition of the PbI_3^- anions. This chemical process first requires a dissolution of **3** and it is consistent with our previous general observation that recrystallizations conducted above 1 GPa often destabilize hydrates.^{60–62} Ultimately, exposing the system to still higher pressure and temperature completely prevents water intercalation into the structure of PbI_2 and leads to the formation of a 3-D structure through creation of new Pb–I bonds and 7-coordinated lead(II) ions.

The effect of temperature and pressure on the aqueous solutions of $\alpha\text{-PbI}_2$ and HI can be represented in the chemical formula reported by Daub and Hillebrecht:³⁰ $(\text{H}_2\text{O})_\delta(\text{PbI}_2)_\epsilon(\text{HI})_\zeta$.

According to this formula, the stoichiometry of compounds **1–4** can be represented as:



where x is the Pb^{2+} deficit parameter.³⁰ The formulae above illustrate the stoichiometric relations observed for the syn-

theses of compounds **1**, **2**, **3** and **4**. When related to the amount of PbI_2 in the ambient-pressure compounds **1** and **2**, the number of moles of HI and of H_2O is higher in **3** and **4**, which can be rationalized by a smaller compression of the Pb–I bonded polyanionic skeletons, compared to considerably softer $\text{OH}\cdots\text{O}$ and $\text{OH}\cdots\text{I}$ interactions. Consequently, more water molecules are needed for separating the polyanions. On the transition from **3** to **4**, the polyanions are compressed closer, their separation decreases and the space available for the corrugated layers of H-bonded H_2O and $[\text{H}_3\text{O}]^+$ is reduced. This reduced space can accommodate fewer H_2O molecules in **4**, whereas the contents of hydronium cations H_3O^+ equilibrate the charge of the polyanions through the neutral-charge condition. The above formulae also illustrate that more HI is needed for the formation of high-pressure products **3** and **4** compared to the gas-phase reaction leading to compound **2**. Under ambient conditions, the increased presence of HI leads to the formation of ribbon polyanions in **2** capable of accommodating more H_2O molecules and H_3O^+ cations around them, compared to the layer polyanions formed in **1**. In the reaction to compound **3**, the higher pressure decreases the distances and further increases electrostatic interactions, which in turn can be better stabilized by compressed contacts between I^- anions and H_3O^+ cations. These charge-assisted contacts are shorter by about 0.5 Å compared to those in ambient-pressure compound **2**. Accordingly, the formation of smaller polyanions increases the surface where the negative charge is distributed on terminal I atoms, hence resulting in the presence of wider triple-chain polyanionic ribbons $[\text{Pb}_3\text{I}_8]_n^{2-}$ in **2** and narrower double-chain polyanionic ribbons $[\text{Pb}_2\text{I}_6]_n^{2-}$ in **3** and **4**.

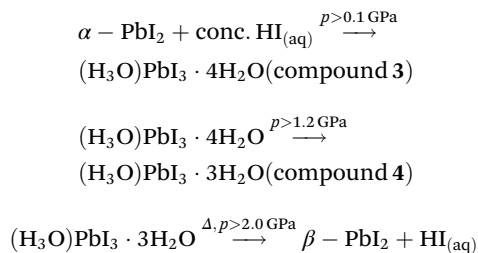
The herein observed structures of hydrated iodoplumbic acids **1–4** resemble some of those recently observed for solvates of popular methylammonium (CH_3NH_3^+) and formamidinium ($\text{CH}(\text{NH}_2)_2^+$) iodoplumbates with N,N -dimethylformamide (DMF), S,S -dimethylsulfoxide (DMSO) and/or water.⁴⁹ Such solvates have attracted significant attention as intermediates in the formation,^{63–65} or products of moisture-induced degradation, of hybrid perovskite thin films highlighting the importance of the current findings.^{66,67} Specifically, structures containing polymeric $\text{Pb}_3\text{I}_8^{2-}$ anions, reminiscent of the ones found in **2**, were observed as solvates for both methylammonium and formamidinium iodoplumbate with DMF,^{68,69} and in the former case also in a solvate with DMSO.⁷⁰ While the PbI_3^- anionic double-chain motifs observed in **3** and **4** are also found in several hydrates of alkaline metal iodoplumbates (II), it is also found in the hydrate and an alternative DMF solvate of $\text{CH}_3\text{NH}_3\text{PbI}_3$.^{49,69} In these two structures, the anionic PbI_3^- double chains are aligned in parallel and separated by protonated cations and solvent molecules (water, DMF), and form arrangements broadly similar to the crystal structures of **3** and **4**, where the anions also lie aligned in parallel and are separated by hydronium cations and water molecules. However, compounds **3** and **4** are not strictly isostructural to any of these previously reported solvated structures.

Other types of anions, not yet observed in our high-pressure studies, have been observed in solvated forms of methyl-

ammonium and formamidinium iodoplumbates, notably a dihydrate of the methylammonium salt based on monomeric octahedral PbI_6^{4-} ions,⁷¹ as well as DMF^{72} and DMSO^{73} solvates of the methylammonium salts containing polymeric PbI_5^- anions composed of corner-sharing PbI_6 -octahedra. Importantly in the context of the present work, the crystal structure⁶⁸ of the DMF solvate of $\text{CH}(\text{NH}_2)_2\text{PbI}_3$ presents an alternative form of the PbI_3^- anion, in the form of simple chains formed by face-sharing PbI_6 -octahedra. While our studies have so far not revealed any evidence for such a phase based on hydronium ions, we believe they suggest the possibility of another, structurally distinct, class of iodoplumbic acid hydrates.

Conclusion

In summary, high-energy isochoric syntheses in a DAC revealed the first structures of the so far elusive and controversial iodoplumbic acid, in the form of a hydrated hydronium salts of $[\text{PbI}_3]^-$ achievable at elevated temperature and pressure, providing new experimental information relevant for the previous speculations regarding possible HPbI_3 structure. In that respect, the iodoplumbic acids synthesized under high-pressure resemble other inorganic acids that are known only in the hydrated form, such as HAuCl_4 or HICl_4 .^{74,75} The herein obtained hydronium-based acids are based on 1-D double-chain of composition PbI_3^- anions, analogous to those found in the archetypal, well-known lead perovskites RbPbI_3 , CsPbI_3 , $\text{NH}_4\text{PbI}_3 \cdot 2\text{H}_2\text{O}$, or $\text{CsPbI}_3 \cdot 2\text{H}_2\text{O}$.^{44–48} The composition and structure of anions distinguish these high-pressure materials from iodoplumbic acids made by recrystallization of PbI_2 from aqueous hydriodic acid,³⁰ and render them the so far first observed examples of iodoplumbic acid: a material of fundamental significance as the formal inorganic progenitor of the highly popular class of hybrid perovskite materials. The herein described series of hydrated iodoplumbic acids constitutes a family of closely related compounds with those previously reported at standard pressure,³⁰ and their high-pressure synthesis can be described through the following equations:



These processes can be rationalized through intercalation of water molecules and hydronium ions between negatively charged lead iodide fragments, resulting in $\text{O}-\text{H} \cdots \text{I}$ and $\text{O}-\text{H}^+ \cdots \text{I}^-$ hydrogen bonds that reduce electrostatic repulsion and prevent close contacts between the anions. This system exhibits high sensitivity to external stimuli, as evidenced by the observation of so far four iodoplumbic acids, up to high

pressures (>2 GPa) and temperatures (>350 K) at which intercalation is no longer favoured and a new form of lead(II) iodide ($\beta\text{-PbI}_2$) appears (see the equation above). The $\beta\text{-PbI}_2$ structure is the unique polymorph of lead(II) iodide and, in contrast to the previously reported phases,⁵⁷ it is based on the 3-dimensional network of edge-sharing PbI_6 and PbI_7 octahedra.

Overall, this work provides new fundamental chemical information about lead-iodide perovskite solids, viewed from the perspective of high-pressure and high-temperature conditions. The role of investigated $\text{PbI}_2/\text{HI}(\text{aq})$ system as a key intermediate in the synthesis of perovskite-based solar cells and its use as a model for probing novel perovskite materials can unveil novel pathways in the renewable energy industry. This exploration underscores the criticality of further research into iodoplumbic-based materials applications, potentially catalysing breakthroughs in the design and performance of next-generation photovoltaic technologies. In line with the high potential of high-pressure techniques in generating new material structures,^{39,40,42,76} this work highlights crystallization from the high-energy DAC environment as a simple and straightforward means to discover new phases, even in compositionally simple systems such as PbI_2 that has been extensively studied^{53,54} for almost a century. Whereas the herein described structures of hydrated iodoplumbic acids are so far accessible only at high pressures, and therefore not of immediate importance for the construction of devices, they provide a fundamental advance in the understanding of this important family of materials and a new high-pressure perspective revealing unexpected structures important for the debate on the existence of iodoplumbic acids.

Experimental section

High-pressure experiments

For the high-pressure experiments a liquid solution of PbI_2 dissolved in concentrated aqueous hydroiodic acid (HI , 57% by weight) was loaded to a modified Merrill-Bassett diamond anvil cell (DAC). The DAC anvils were supported directly on the steel discs with conical windows (the culet size was 0.8 mm, type 1A diamonds and gasket was made of 0.15 mm thick austenitic steel (type 25-6MO) with the hole diameter 0.45 mm).⁴¹ Pressure in the DAC was determined by the ruby fluorescence (R1 ruby line) shift with a photon control spectrometer affording an accuracy of 0.02 GPa.⁷⁷ Throughout all experiments, the hydrostatic conditions in the DAC were routinely inspected visually by microscopic observations (*cf.* General microscopy in ESI† for details), by checking the width of the R1 and R2 ruby fluorescence peaks and the widths of SCXRD reflections as well as by searching the background of X-ray images for the presence of diffraction events other than those from the sample crystal. By these means we can be confident that all experiments were conducted in hydrostatic conditions. We attempted high-pressure syntheses and recrystallizations above 3 GPa, however most of the DAC chamber contents froze and its melting required the temperatures approaching the re-

sistance of the steel parts of the DAC and caused difficulties in controlling precisely the temperature and pressure, as required for the growth of single crystals.

Diffraction data were collected at 295 K for **1**, **3** and **4** and at 320 K for β -PbI₂, by using a KM-4 CCD diffractometer with the graphite-monochromated MoK α radiation. The DAC was centered by the gasket-shadow method.⁷⁸ The CrysAlisCCD and CrysAlisRED programs were used for collecting the data, determination of the UB-matrices, initial data reduction, and L_p correction. Reflection intensities were corrected for the DAC and sample absorption; the gasket shadowing and the reflections of diamond-anvils were eliminated.⁷⁹ All structures were solved by direct methods, and refined with anisotropic displacement parameters, with programs ShelXS and ShelXL using Olex2 interface.^{80,81} For re-determination of the structure of **1**, the positions of water hydrogen atoms were estimated from the molecular geometry and consistently with the pseudo-hexagonal hydrogen-bonding pattern in the structure. The water molecules were then refined using a rigid-group model, with the isotropic temperature factors of the hydrogen atoms (U_{iso}) constrained to be 1.5 times the U_{eq} of the corresponding oxygen atom. Details of structure refinements and crystal data are given in Table S1.† Crystallographic data in CIF format have been deposited with the Cambridge Structural Data Centre, under deposition codes 2071140–2071144.†

High-pressure synthesis of **3** & **4**

Single crystals of **3** and **4** were obtained in isochoric conditions: after the polycrystalline mass precipitated, the DAC was heated using a heat gun until all but one grain dissolved. Then the single crystal grew as the DAC was cooled slowly to room temperature, where the pressure was remeasured before and after the X-ray diffraction experiment. The progress and experimental details on growing the single crystals of **3** and **4** are shown in Fig. S1 and S2.†

High-pressure synthesis of β -PbI₂

Single crystals of β -PbI₂ were obtained similarly to those of **3** and **4**, however due to the phase transformation at room temperature at 1.2 GPa, the single crystal grown at 2.05 GPa was kept at 320 K and the X-ray diffraction experiment was performed at this temperature.

Ball milling experiments

Milling synthesis of 1: For the ball milling synthesis, 0.461 g (1 mmol) of PbI₂ was added to one half of a zirconia (ZrO₂) 10 mL jar, followed by the addition of 67 μ L (0.5 mmol) of aqueous hydroiodic acid (HI, 57% by weight) and one 3.5 g zirconia ball. The jar was carefully sealed, placed on a MM400 mixer mill and the reaction mixture was milled for 30 minutes at a frequency of 30 Hz. Reaction completion afforded a bright yellow product, which was left to dry in dark for 1 h. The resulting solid product was scraped off the jar walls and subjected to powder X-ray diffraction (PXRD) analysis, revealing the formation of compound **1**. Compound **1** can be also isolated when performing the 1:1 stoichiometric

reaction by milling 0.230 g (0.5 mmol) of PbI₂ with 67 μ L (0.5 mmol) of aqueous hydroiodic acid (HI, 57% by weight), respectively. **Milling synthesis of 2:** For the ball-milling synthesis 0.230 g (0.5 mmol) of PbI₂ was added to one half of a zirconia (ZrO₂) 10 mL jar, followed by the addition of 201 μ L (1.5 mmol) of aqueous hydroiodic acid (HI, 57% by weight) and one 3.5 g zirconia ball. The jar was carefully sealed and placed on a MM400 mixer mill and the reaction mixture was milled for 30 minutes at an oscillation rate of 30 Hz. Reaction completion afforded a bright yellow product, which was left to dry in dark overnight. The resulting product scraped off the jar walls was identified as compound **2** by PXRD.

Data availability

Description of pseudosymmetry in **1**, experimental procedures and detailed crystallographic data of **3**, **4** and β -PbI₂, ball-milling experiments, and additional data are located within the ESI.†

Author contributions

All high-pressure experiments were performed by S. S., ball milling by A. M. F. and J.-L. D. Research was coordinated and supervised by A. K., T. F., A. M. and G. P. D. All authors discussed the results, contributed to writing the manuscript and commented on it.

Conflicts of interest

The authors declare no conflict of interest.

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