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Unconventional Electron-Deficient Multicenter Bonds in AIO₃ Perovskites

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eventually to a cubic perovskite, with IO_6 units, in other AIO_3 compounds. We demonstrate that the PIP process induces a change in the chemical bonding from the resonant delocalized I–O bonds in IO_3^- polyanions toward the unconventional I–O electrondeficient multicenter bonds (EDMBs) in AIO_3 cubic perovskites. The process of EDMB formation in the cubic perovskites agrees with the recently proposed unified theory of multicenter bonding and contradicts previous assumptions that considered these bonds to be impossible in valence electron-rich elements, such as chalcogens and halogens. Interestingly, our results suggest that (i) the formation of the cubic and slightly distorted ABX_3 and BX_3 perovskites, with A, B, and X being main-group elements, at high pressure is driven by the formation of 3D EDMBs due to the PIP process of the BX_3 units (monomers) leading to the formation of regular BX_6 units; and (ii) unconventional EDMBs could be already present at room conditions in the cubic or slightly distorted ABX_3 and BX_3 perovskites, with A, B, and X being main-group elements. The presence of unconventional EDMBs could explain the extraordinary properties of these perovskites.

1. INTRODUCTION

The perovskite (PV) structure can be found in ABX_3 and BX_3 compounds and is perhaps the single most versatile ceramic host since appropriate changes in composition can transform the most significant electro-ceramic dielectric phase in the industry, BaTiO₃, into metallic conductors, superconductors, catalysts, sensors, lasers, magnetoresistive/multiferroic/thermoelectric materials, and even high-performance photovoltaic materials, such as lead halide PVs.¹⁻⁴ The rich diversity of chemical compositions and properties of PVs results in "high-tech" applications, which have made PVs to be dubbed *inorganic chameleons*.

PVs are also very important materials from a fundamental perspective, and still, they represent an exciting challenge in terms of chemical bonding. If we consider ABX_3 PVs with A, B, and X being main-group elements, e.g. $CsPbI_3$, it is clear that there is mainly an ionic A-X bonding. However, it must be noticed that a certain covalent character has been observed in

the A-X bonding of certain PVs when A is a group-15 element (e.g., Pb in PbTiO₃).⁵ On the other hand, the nature of the chemical B-X bonding, which forms the ReO₃-type octahedral framework in which A atoms are inserted,⁶⁻⁸ is a mystery to be unraveled in both ABX_3 and BX_3 PVs of main-group elements.

Claims for a mixture of classical ionic, covalent, and even metallic bonding have been proposed to be present in B-X bonds to explain the extraordinary properties of ABX_3 and BX_3 PVs.^{9,10} Recently, the debate regarding the chemical B-X bonding in PVs has been renewed since Wuttig and co-workers have suggested that lead halide PVs, like CsPbI₃, feature an

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Figure 1. Crystal structure of $CsIO_3$ in (a) the R3m phase at 0 GPa and (b) the $Pmn2_1$ phase at 20 GPa. Cs, I, and O atoms are plotted in gray, pink, and red, respectively. Short and long bonds are noted as solid and dashed orange lines. The IO₃ trigonal pyramids can be observed. The perspective of both structures has been selected to highlight their similitudes.

unconventional new type of two-center-one-electron (2c-1e) bonding different from the classical (covalent, ionic, and metallic) ones, dubbed as metavalent bonding.¹¹ Noteworthy, this new brand bonding is the same that is supposed to explain the extraordinary properties of phase change materials.¹² In this context, Wuttig and co-workers have suggested that oxide perovskites, like BaTiO₃, do not have this kind of unconventional bonding (suggesting that the Ti–O bond is of covalent type).¹¹

Contrarily, other researchers consider that the metavalent bonding is simply the century-old electron-rich multicenter bonding (ERMB), also known as the hypervalent bonding or the three-center-four-electron (3c-4e) bonding,¹³ and propose that the extraordinary properties of phase change materials is explained in terms of the ERMBs.

To complete the picture, Manjón and co-workers, who have proposed a unified theory of multicenter bonding,^{14,15} have suggested that the unconventional bonding of phase change materials and lead halide PVs is neither the metavalent bonding nor the ERMB, but just the also century-old electron-deficient multicenter bonding (EDMB) occurring in solids with electron-rich elements.^{14,15} These EDMBs are just a linear or quasi-linear concatenation of three-center-two-electron (3c-2e) bonds in which a single electron is present between two atoms; i.e., a 2c-1e bond, as shown by Wuttig and co-workers.

The proposal of Manjón and co-workers that the unconventional EDMB is present in phase change materials and lead halide PVs is based on different bonding descriptors and the behavior of those descriptors with increasing electron density.^{14,15} One of the bonding descriptors is the combination of the number of electrons shared (ES) and the normalized number of electrons transferred (ET) between two atoms, as obtained from the Quantum Theory of Atoms in Molecules.¹⁶ In the unified theory of multicenter bonding, Manjón and coworkers have shown that the ES and ET values are different for the two multicenter bondings (ERMBs and EDMBs), so they can be located in different regions of the ES vs ET map.^{14,15} Several arguments of Manjón and co-workers demonstrate that metavalent bondings are the old EDMBs and not the old ERMBs: (i) the region in the ES vs ET map for the EDMB is the same already attributed to the new metavalent bonding and completely different to that for the ERMB; and (ii) the two multicenter bonds can be fully distinguished because they have

a different formation mechanism. Despite both multicenter bonds share a common origin, their formation proceeds through different mechanisms.

The unified theory of multicenter bonding also suggests that the increase in electron density, either by applying highpressure (HP) conditions, reduction (i.e., injection of electrons in a system), or substitution of elements by their heavier analogs, helps to form unconventional multicenter bonds. Both ERMBs and EDMBs can be formed by a polymerization process, in which multicenter bonds are formed from original primary and secondary bonds. The primary bonds are usually iono-covalent bonds, although we have recently shown that pressure-induced EDMBs can be formed in AX₃ compounds when primary bonds are ERMBs.¹⁷ On the other hand, secondary bonds are usually noncovalent interactions related to the presence of stereochemically active lone electron pairs (LEPs), typically leading to asymmetric structures at room pressure (RP).¹⁸⁻²³ Importantly, ERMBs and EDMBs have a different formation mechanism because the formation of the ERMB involves the transformation of the nonbonding electrons present in the LEP of secondary bonds into bonding electrons in the ERMB, so the charge for the new bonds is provided by a LEP. On the contrary, the formation of the EDMB is mainly due to the charge transfer from the primary iono-covalent bond to the secondary noncovalent interaction. In other words, the charge for the new EDMB is provided by the strong bond, and, in general, the LEP becomes stereochemically inactive during the formation of the EDMB, so the nonbonding electrons of the LEP remain as nonbonding electrons when the EDMB is formed.^{14,15}

The present work aims to show that EDMBs do occur in main-group PVs, in particular in some oxide PVs, such as those formed by AIO_3 (A = K, Rb, Cs, Tl, NH₄) compounds at HP. For this purpose, we have selected CsIO₃, a compound with a strongly distorted PV structure, and conducted a joint experimental and theoretical study of this compound under compression which is complemented with ab initio calculations in related AIO_3 compounds. The choice of CsIO₃ to conduct this study is justified because it is a member of the AIO_3 family and this family was theoretically predicted to gradually become closer to the cubic PV structure under compression.²⁴ In particular, RbIO₃ was shown to progressively transform from the distorted rhombohedral PV structure (space group (SG)

R3*m*, No. 160) to the perfect cubic PV structure (SG $Pm\overline{3}m$, No. 221) at moderate pressures (above 35 GPa).

The compounds of the AIO₃ family have been profusely studied because they show ferroelectric and nonlinear optical properties, $^{25-30}$ related to their strongly distorted PV structures. $^{28,31-36}$ The strong distortion of the IO₆ octahedra in AIO₃ compounds at RP results in molecular compounds with IO₃ units and a much smaller symmetry than the cubic PV structure. KIO₃ crystallizes in a triclinic noncentrosymmetric (space group (SG) P1, No. 1) structure³¹⁻³³ and NH₄(IO₃) does it in an orthorhombic (SG *Pna2*₁, No. 33) structure.³⁴ Less distorted phases are found in AIO₃ (A = Rb, Cs, Tl) compounds, which crystallize in the rhombohedral (SG *R3m*, No. 160) structure at RP.^{28,35,36} Noteworthy, the rhombohedral R3*m* phase is also a high-temperature phase of KIO₃ at RP.²⁹

In the cubic PV structure, all the B-X bonds of the BX_6 units have the same bond length; however, in the strongly distorted PV structure of CsIO₃, there are two I–O bond lengths (see Figure 1a): one short I–O bond length, which corresponds to three delocalized resonant (covalent-like) bonds that define a trigonal pyramidal IO₃ unit, and one long I–O bond length, which corresponds to three interactions that can be considered as secondary noncovalent bonds in a strongly distorted IO₆ octahedron (indeed an IO₃₊₃ unit).

To our knowledge, the AIO₃ family of distorted PVs has not been studied in deep under compression, with only one experimental study on KIO3³⁷ and one theoretical study on RbIO₃²⁴ being available. In both works, a pressure-induced symmetrization (PIS) of the distorted PV structure was found to be driven by the regularization of the distorted IO₆ units. In KIO₃, two pressure-induced phase transitions were observed at 7 and 14 GPa under hydrostatic conditions. Under nonhydrostatic conditions, these transitions occurred at significantly lower pressures (4 and 10 GPa).³⁷ The first HP phase of KIO₃ was identified as a rhombohedral structure (SG R3, No. 146). On the other hand, RbIO₃ was theoretically predicted to transform from the rhombohedral to the cubic PV structure (SG $Pm\overline{3}m$, No. 221) beyond 35 GPa.²⁴ Notably, a PIS of the distorted IO₆ units has also been observed in several metal iodates,^{38–43} as recently reviewed.⁴⁴

In relation to the PIS of the distorted IO_6 units in iodates, several recent studies on metal iodates have evidenced a pressure-induced change in chemical I–O bonding with the increase in I coordination (hypercoordination).^{38–41} Specifically, the delocalized resonant (covalent-like) I–O bonds in IO_3 units at RP have been suggested to transform into EDMBs at HP.⁴¹ We have to note in passing that some of us previously referred to EDMBs as metavalent bonds in earlier studies on metal iodates under compression;^{38–40,44} this was due to the claimed electron-deficient character of the metavalent bond just before it was clarified by Manjón and co-workers that the metavalent bonds are 2c-1e EDMBs that result from the linear (or quasi-linear) concatenation of 3c-2e bonds in solids with electron-rich elements.^{14,15,45}

In this work, we present a joint HP experimental and theoretical study of the distorted PV structure of $CsIO_3$ to study the change in chemical I–O bonding during the PIS of the crystalline structure of $CsIO_3$. For this study, we have combined experimental X-ray diffraction (XRD) and Raman scattering (RS) measurements, ab initio calculations, and a detailed electron topology analysis in several members of the AIO_3 family. Our results reveal that $CsIO_3$ undergoes several

pressure-induced phase transitions driven by the pressureinduced polymerization (PIP) process of the IO₃ units, which are the monomers that become linked upon pressure increase. As a consequence of this process, the original rhombohedral phase (SG R3m, No. 160) undergoes a phase transition to an orthorhombic structure (SG Pmn2₁, No. 31). In addition, a second HP phase with a tetragonal structure (SG P4/nmm, No. 129) is theoretically predicted beyond 45 GPa. We demonstrate that the PIP process leads to a progressive PIS of the structure and a change in the chemical I-O bonding with the increase in I coordination. In particular, we show that two of the three short delocalized resonant I–O bonds of IO₃ units at the R3m phase at RP progressively transform (together with two secondary noncovalent bonds) into two EDMBs as pressure increases and the tetragonal PV phase (with IO₅₊₁ units) is formed in CsIO₃. Our findings align with the recently developed theory of multicenter bonding^{14,15} and contradict previous assumptions that considered EDMBs to be impossible in valence electron-rich elements, such as chalcogens and halogens. Interestingly, our findings suggest that the cubic PV structure in ABX₃ and BX₃ compounds with main-group elements, e.g., in AIO₃ compounds at HP, is likely characterized by the presence of EDMBs among the regular BX_6 units that result from the polymerization of BX_3 units.

2. EXPERIMENTAL DETAILS

2.1. Synthesis and Characterization at Room Pressure. Cesium iodate (CsIO₃) powders were synthesized from the reaction of cesium carbonate (Cs_2CO_3) and iodic acid (HIO_3) . All starting materials, including Cs₂CO₃ (99.9%) and HIO₃ (99.5%), were purchased from Sigma-Aldrich and used without further purification. Cs_2CO_3 , with a molar mass of 325.82 g/mol, was weighed to achieve a 0.004 M concentration, corresponding to 1.31 g. HIO_3 , with a molar mass of 175.91 g/mol, was similarly prepared at a 0.004 M concentration, equating to 0.704 g. Each substance was dissolved separately in 15 mL of distilled water and stirred for 10 min. The solutions were then combined to produce a total volume of 30 mL and mixed for an additional 10 min. The resulting mixture was transferred to a 40 mL Teflon-lined stainless-steel autoclave and heated in an electric oven at 220 °C for 48 h. The obtained powder was washed several times with distilled water to remove impurities and then dried in an electric oven at 100 °C for 24 h.

The purity of the samples was determined via powder XRD analysis performed using a Rigaku Ultima IV diffractometer in the Bragg– Brentano configuration with CuK α radiation ($\lambda = 1.54060$ Å) at room temperature. Data were collected over a 2θ range of $10-70^{\circ}$ with a step size of 0.02° and a fixed counting time of 1 s per step. A single phase corresponding to the rhombohedral phase of CsIO₃ was found, thus confirming the purity of the synthesized samples.

2.2. Characterization at High Pressure. HP experiments on powders of CsIO₃ were conducted utilizing a membrane-style diamond-anvil cell having diamonds with anvils measuring 350 μ m in diameter. Stainless-steel gaskets preindented to a thickness of 40 μ m, with a 120- μ m diameter hole at the center, were employed. As a pressure-transmitting medium, we used a mixture of methanol, ethanol, and water with a ratio 16:3:1, which ensures quasi-hydrostatic conditions up to ca. 10 GPa.⁴⁶ In all experiments, the diamond-anvil cell loading was performed avoiding sample bridging between the diamonds.⁴⁷

2.2.1. HP-XRD Experiments. We performed two HP-XRD experiments in CsIO₃. One was carried out at the Xpress beamline of Elettra Synchrotron Radiation Facility up to 34.9(1) GPa and the other was conducted up to 40.6(1) GPa at the BL04-MSPD beamline of ALBA synchrotron.⁴⁸ At Elettra (ALBA) we used a monochromatic wavelength of 0.4957 Å (0.4246 Å) and a DECTRIS-PILATUS-3 S 6-M (Rayonix SX165) detector. At Elettra (ALBA) the X-ray beam was focused down and to a spot of 50 μ m × 50 μ m (20 μ m × 20 μ m) in

size. The sample–detector distance and the detector geometry were calibrated using a LaB₆ standard and the pressure was measured using the signal of Cu powder loaded together with the sample and the equation of state (EOS) reported by Dewaele et al.⁴⁹ The two-dimensional diffraction images collected with the detectors were processed to create intensity versus 2θ XRD patterns utilizing DIOPTAS.⁵⁰ The analysis of these data was performed by means of Rietveld refinements conducted with the FullProf suite.⁵¹

The reason for the two HP-XRD measurements was that there were reasonable doubts regarding the nature of the first HP phase above 14 GPa. Initially, we were working with two possibilities (a monoclinic phase and an orthorhombic one). Both were very similar and none of them was theoretically predicted since calculations of enthalpy curves of the two phases were on top of the original R3m phase. Therefore, we decided to see if a second experiment could give us slightly better results than the first one and clarify which of the two HP candidate phases was the most likely one. At the end, we concluded that the orthorhombic one was the best candidate in reasonable agreement with Raman scattering experiments.

2.2.2. HP-RS Experiments. We performed one HP-RS experiment in powder $CsIO_3$ up to 31.2 GPa. RS signal was excited with a 633 nm laser (with a power of less than 10 mW) in backscattering geometry using a Horiba Jobin Yvon LabRAM HR UV microspectrometer. The microspectrometer is equipped with a thermoelectrically cooled multichannel charge-coupled device detector and a 1200 grooves/mm grating that allows a spectral resolution better than 2 cm⁻¹. Pressure was determined by the ruby luminescence method.⁵² Raman peaks were analyzed with a Voigt profile fixing the Gaussian line width (1.6 cm⁻¹) to the experimental setup resolution.

3. THEORETICAL DETAILS

All theoretical calculations in this paper were performed using density functional theory (DFT) as implemented in the Vienna Ab initio Simulation Package (VASP).53-56 The electronic structure was computed using PAW potentials⁵⁷ with Cs (5s², $5p^6$, $6s^1$), I ($5s^2$, $5p^5$) and O ($2s^2$, $2p^4$) electrons treated as valence electrons. The exchange-correlation term was computed using the PBEsol functional⁵⁸ and the plane waves basis was expanded up to a 560 eV kinetic energy cutoff. The sampling of the Brillouin-zone (BZ) was converged with Γ centered Monkhorst–Pack⁵⁹ grids employing adequate meshes $8 \times 8 \times 4$ for the trigonal (*R3m*) phase, $6 \times 8 \times 6$ for the orthorhombic $(Pmn2_1)$ phase, and $6 \times 6 \times 8$ for the tetragonal (P4/nmm) phase. The cell parameters and atomic positions for the different structures were fully optimized for a range of volumes by calculating the forces on atoms and the stress tensor. In terms of the resulting optimized configurations, the forces on atoms were smaller than 0.003 eV/Å, and the deviation of the stress tensor components from the diagonal hydrostatic form was less than 0.1 GPa with a convergence of the total energy within 10^{-6} eV.

The phonon properties were computed by using the supercell finite-displacement method implemented in the Phonopy package,⁶⁰ with VASP being used as the second-order force calculator. Supercells were expanded up to $2 \times 2 \times 2$ for both *R3m*, *Pmn2*₁, and *P4/nmm* phases, enabling the exact calculation of frequencies at the zone center (Γ) and inequivalent zone-boundary wavevectors, which were then interpolated to obtain phonon-dispersion curves.

A density-based approach grounded in the Quantum Theory of Atoms in Molecules (QTAIM) was employed to analyze the electron density topology of $CsIO_3$ in the different phases. Quantum ESPRESSO (version 6.5)⁶¹ was used for this analysis, in conjunction with Wannier90^{62,63} and CRITIC2 programs.⁶⁴ Single-point calculations were performed at the VASP equilibrium geometries, using the same uniform k-point

grids mentioned above. A plane-wave cutoff of 100 Ry and a density cutoff of 400 Ry were consistently applied. Normconserving pseudopotentials for the Kohn-Sham states and PAW data sets for the all-electron density were sourced from the pslibrary.⁶⁵ Delocalization index (DI) calculations, used to determine the number of electrons shared (ES) between two atoms as $2 \times DI$, were conducted using a Wannier transformation as described in ref 66. The renormalized number of electrons transferred between two atoms (ET) for I-O bonds was calculated as the Bader charge of the I atom divided by the nominal valence (5+). In addition, this value is divided by 3 to account for the different structural multiplicities of I and O atoms (being the charge of I atoms distributed into 3 O atoms in the IO₃ units according to the formula unit CsIO₃) in all phases. All the structures were visualized by VESTA program.⁶⁷ We have to note that our density-based calculations provide a consistent picture of chemical bonding, which is similar to that provided by orbitalbased calculations.^{15,68} We have also provided ELF calculations (using VASP) to further analyze the electron density topology. Note that ELF calculations with PAW pseudopotentials using VASP are not all-electron but they contain a partial information on core electrons and have been used in previous works to study phase change materials which have the EDMBs we deal with in this work.^{69,70}

4. RESULTS AND DISCUSSION

4.1. Structural Behavior under Compression. XRD patterns of CsIO₃ at RP (10^{-4} GPa) were assigned to the rhombohedral R3m structure, as in a previous work.²⁸ The assignment is supported by the small residuals of the Rietveld refinements and by the R-values obtained (see Figures S1 and S2 in Section 1 in the Supporting Information (SI)), which support a good fit of the structural model to the data (see Table S1 in SI). The R3m structure can be described as a distorted pseudocubic structure because $c/a \sim \sqrt{3/2}$ and the structure is composed of CsO₁₂ polyhedra and IO₃ trigonal pyramids with three oxygen atoms in the base and an iodine atom in the vertex. The stereochemically active LEP of iodine points to the opposite direction of the base (along the [001]direction), featuring a well-ordered alignment. Interestingly, a cubic $Pm\overline{3}m$ structure was identified for CsIO₃ at RP by Goldschmidt et al. in an early study.⁷¹ However, this is not the correct structure of our sample since the cubic Pm-3m structure does not exhibit first-order Raman-active modes while our CsIO₃ sample exhibits 13 experimental Raman-active modes (see discussion in Section 2 in SI) that are consistent with the assignment to the trigonal structure.

Two HP-XRD experiments at the ALBA and Elettra synchrotron facilities were performed up to a maximum pressure of 40.6 GPa. Both experiments show evidence of a reversible pressure-induced phase transition from the low-pressure (LP) rhombohedral R3m structure to the HP orthorhombic $Pmn2_1$ structure (see discussion in Section 1 in SI). Rietveld refinements of XRD patterns above 14 GPa can be better explained with the $Pmn2_1$ structure than with the R3m phase (see discussion of Figure S3 in SI). The HP phase can be considered a distorted version of the LP phase. The structural information on the orthorhombic $Pmn2_1$ structure at 14.4(1) GPa is given in Table S2 in SI. It must be stressed that the existence of the $R3m \rightarrow Pmn2_1$ phase transition is consistent with the results of our Raman experiments (see the

following section) and with the results of our DFT calculations, which show that both phases have the same enthalpy, within the accuracy of calculations, for the pressures of interest for this study.⁷²

From the XRD patterns, we obtained the pressure dependence of the unit-cell parameters (Figure 2a) and unit-



Figure 2. Pressure dependence of the unit-cell parameters (a) and volume (b) of the LP R3m and HP $Pmn2_1$ phases. Solid (empty) symbols are used for the ALBA (Elettra) experiments. Black (color) symbols are used for the LP (HP) phase. Solid (dashed) lines correspond to results from DFT calculations for the LP (HP) phase. Unit-cell parameters are identified in the figure. For the HP phase, we have plotted $\sqrt{3}b$ to facilitate the comparison with the LP phase. The red line is the experimental EOS described in the text. The unit-cell volume of the HP phase was renormalized to facilitate the comparison with the LP phase. The inset shows the pressure dependence of the c/a ratio in the LP phase.

cell volume (Figure 2b) of the two phases of CsIO₃. The compression is anisotropic, being the *c*-axis more compressible than the *a*-axis in the hexagonal unit-cell of the R3*m* phase. This explains the peak splitting we described when discussing Figures S1 and S2 in SI. The anisotropic behavior of the structure can be better appreciated in the decrease of the c/a ratio at HP and its deviation from the $\sqrt{3/2}$ value (see inset of Figure 2b). Figure 2b shows that our calculations reproduce the behavior of the experimental c/a ratio of the R3*m* phase and the unit-cell parameters of the R3*m* and Pmn2₁ phases, which point to the convergence of the *a* and *c* lattice parameters of the HP phase above 45 GPa. This result

confirms the hypothesis of the expected PIS of the structure of CsIO₃. Data of the experimental and theoretical equation of state (EOS) of CsIO₃ under compression are provided in Section 1 in SI. It must be stressed that CsIO₃ is one of the most compressible iodates,⁴⁴ indeed as compressible as $Sr(IO_3)_2HIO_3^{41}$ and as $KIO_3^{.37}$

Rietveld refinement has allowed us to obtain the evolution of the experimental atomic parameters of both R3m and $Pmn2_1$ phases of CsIO₃ up to 40 GPa. As observed in Figure S4 in SI, there is a nice agreement between experimental and theoretical atomic parameters in the $Pmn2_1$ phase from 14.4 to 40 GPa. Importantly, our theoretical calculations up to 70 GPa show that the atomic parameters of the $Pmn2_1$ phase evolve toward values compatible with a tetragonal phase (SG P4/*nmm*, No. 129) above 45 GPa. The structural parameters of the P4/*nmm* phase at 52 GPa are summarized in Table S3 in SI. Figure 3 shows a picture of the tetragonal P4/*nmm* phase of CsIO₃ expected above 45 GPa and a comparison with the orthorhombic $Pmn2_1$ phase.

From the experimental and theoretical pressure dependence of the atomic parameters, we have calculated the pressure dependence of the experimental and theoretical I-O bond lengths in the three phases $(R3m, Pmn2_1, and P4/nmm)$ of $CsIO_3$ (Figure 4a). As observed, the 3-fold-degenerated short delocalized resonant I-O bond of the LP phase (intramolecular bonds inside the IO₃ units) shows a slight increase in bond length with increasing pressure up to 14 GPa, while the 3-fold-degenerated long secondary noncovalent I···O bond of the LP phase (intermolecular bonds between neighbouring IO_3 units) show a considerable decrease (14%) in bond length in the same pressure range; i.e., an astonishing average change of around 1% per GPa. Therefore, IO₃ units exhibit a strong tendency to form IO₆ units, as would be expected in a cubic or slightly distorted PV structure. The increase in symmetry of the IO_6 units (as all the I–O bonds tend to equalize) is not completed at 14 GPa and continues above this pressure in the Pmn21 phase. In this HP phase, the 3-fold-degenerated short and long I-O bond lengths (six in total) of the LP phase split into one single and one 2-fold-degenerated short and long I-O bond distances (six in total). The single short and long I-O bond lengths (two in total) are along the b-axis of the orthorhombic phase, while the 2-fold-degenerate short and long I–O bond lengths (four in total) are in the *ac*-plane of the orthorhombic phase. Interestingly, the 2-fold-degenerated short and long distances in the ac-plane tend to become equal as pressure increases; i.e., four of the original six bonds in the LP phase tend to the same bond length in the HP Pmn2₁ phase. Our calculations, which nicely reproduce the experimental results, predict that the mentioned PIS occurs beyond 45 GPa once the HP tetragonal P4/nmm phase is formed (see black dashed lines in Figure 4a). On the contrary, the single short I–O and long I···O bond distances along the *b*-axis of the $Pmn2_1$ phase, which tend to equalize in the pressure range between 14 and 25 GPa, display an opposite tendency above 25 GPa and remain as short and long bonds in the P4/nmm phase (see squares and solid lines in Figure 4a). This results in the formation of IO_{5+1} units in the tetragonal P4/*nmm* phase as shown in Figure 3b.

As commented in the Introduction, the PIS of the crystalline structure of $CsIO_3$ was expected due to a theoretical work in isostructural $RbIO_3$ which suggested that the full PIS would end with the formation of the cubic PV structure (*Pm-3m*, S.G. No. 221) around 35 GPa.²⁴ Instead, our calculations predicted



Figure 3. Crystal structure of CsIO₃ in (a) the HP orthorhombic $Pmn2_1$ phase at 14.4 GPa (see data of Table S2) and (b) the HP tetragonal P4/nmm phase at 52 GPa (see data of Table S3). The perspective of both structures has been selected to highlight their similitudes. The orthorhombic phase at 14.4 GPa still exhibits IO₃ trigonal pyramidal units since there are very distorted IO₆ units with three long I···O bond distances (dashed orange lines) along the *a*, *b*, and *c* axes. The tetragonal phase still exhibits slightly distorted IO₆ octahedra that can be described as square pyramidal IO₅₊₁ units since there is a long I···O distance (dashed orange line) along the *c*-axis. Note that all I–O distances in the *ab* plane of the tetragonal phase are equal and slightly larger than the short I–O bond along the *c*-axis; therefore, the tetragonal structure is a slightly distorted (quasi-cubic) perovskite structure.



Figure 4. (a) Pressure dependence of experimental (symbols) and theoretical (lines) I–O bond distances in the different phases of CsIO₃. Circles (triangles and squares) correspond to the R3m ($Pmn2_1$) phase. Solid (open) symbols correspond to the short and long distances in both R3m and $Pmn2_1$ phases. Solid and dashed lines correspond to theoretical bonds with different pressure behavior. Dashed vertical lines indicate the pressures for the experimental (first) and theoretical (second) phase transitions. (b) Pressure dependence of theoretical ES values of the I–O bond distances in the different phases of CsIO₃. To help in the interpretation of the theoretical data we have used the same symbols as in (a) for the R3m and $Pmn2_1$ phases. For the P4/nmm phase solid (open) green diamonds are plotted to distinguish between short (long) bonds.

that a full PIS of the structure in $CsIO_3$ does not occur even at 70 GPa, although a full PIS at higher pressures cannot be discarded.

In summary, experimental and theoretical structural information in CsIO₃ under compression suggests the presence of two pressure-induced phase transitions up to 50 GPa, the first one (from rhombohedral to orthorhombic) around 14 GPa and the second one (from orthorhombic to tetragonal) above 45 GPa. The transitions occur without appreciable changes in the unit-cell volume; however, it does not exist a group-subgroup relationship either between the R3m and $Pmn2_1$ space groups or between the $Pmn2_1$ and P_4/nmm space groups; thus indicating that the transitions could not be of second order⁷³ and are likely weak first-order transitions. The fact that there is no bond breaking/formation at the transitions supports a displacive mechanism for these transitions, which is consistent with their reversibility.⁷⁴ These phases occur before the possible phase transition toward the cubic PV phase above

70 GPa. It must be noted that the PIS following the rhombohedral-orthorhombic-tetragonal-(cubic) sequence of phase transitions in CsIO₃ agrees with that expected for the formation of the cubic PV structure, as observed in BaTiO₃ with increasing temperature (see Figure 6.2 in ref 4) and also similar (in this case the rhombohedral-orthorhombic phases are reversed) to that predicted for LaGaO₃ under compression.⁷⁵ We have also to note that the rhombohedral-to-orthorhombic phase transition near 14 GPa is perhaps induced by non-hydrostatic conditions since our calculations of both phases on hydrostatic conditions indicate that both phases have the same enthalpy in the whole pressure range studied.⁷²

The orthorhombic-tetragonal phase transition above 45 GPa is a phase transition resulting in a PIS of the IO_6 octahedra which leads to a 5 + 1 coordination for I atoms; i.e., IO_{5+1} units are formed in the tetragonal PV phase. The slightly distorted PV structure of the tetragonal phase in CsIO₃ at 52 GPa can be noted if we compare the I–O bond distances in IO_{5+1} units

(with one I–O bond at 1.824 Å, four at 1.948 Å, and one much longer at 2.205 Å) with those of the TiO₅₊₁ units present in the tetragonal distorted PV phase of BaTiO₃ (with one Ti–O bond at 1.86 Å, four at 2.00 Å and one longer at 2.17 Å) as reported at page 469 in ref 3. Similar BO_{5+1} units are also present in other tetragonal PV phases, as that of T*-Nd_{2-y-z}Ce_ySr_zCuO₄ (see Figure 6.10 in ref 4).

4.2. Vibrational Behavior under Compression. A selection of the measured RS spectra of CsIO₃ at different pressures is shown in Figure S6 in SI. A total of 13 Raman modes have been experimentally detected in the range up to 1000 cm⁻¹, in agreement with those previously reported at RP.⁷⁶ These results are consistent with group theory predictions for the rhombohedral R3m phase. The different Raman-active modes observed in the LP phase are provided in Table S4 in SI and discussed in Section 2 in SI. Modes of the LP phase are observed up to 16.4 GPa and changes in the Raman spectrum are observed between 13.5 and 16.4 GPa (see Figure S6), in agreement with the phase transition detected by HP-XRD experiments. The increase in the number of Raman modes observed above 13.5 GPa is consistent with a decrease in the symmetry of the crystal structure at the phase transition. Again, this is in agreement with what is observed in HP-XRD experiments.

According to group theory,⁷⁷ the orthorhombic $Pmn2_1$ phase (SG No. 31, Z = 2) of CsIO₃ has 30 vibrational modes, comprising 3 acoustic modes (A₁+ B₁+ B₂) and 27 optical modes (8A₁ + 6A₂ + 5B₁ + 8B₂). All optical modes are both Raman- and infrared (IR)-active except for the A₂ modes, which are silent ones. This means that both the A₁, B₁, and B₂ modes exhibit TO-LO splitting. Therefore, a total of 42 Raman- and IR-active modes are expected in the $Pmn2_1$ phase of CsIO₃. In contrast, in our RS measurements, we have only detected a total of 13 Raman modes.

Figure 5 shows the experimental and theoretical pressure dependence of the wavenumber of the Raman-active modes in both R3m and Pmn2₁ phases of CsIO₃ up to 30.2 GPa. Experimental data (symbols) are compared to theoretical calculations (lines) which include the TO-LO splitting. Our calculations show that while TO-LO splitting is considerable for E modes, the splitting is negligible for A modes. In the R3m phase, there is a good agreement between experiments and calculations both for the wavenumbers of the first-order Raman-active modes (within 5% accuracy) and their pressure dependence. While the assignment of the symmetries of the Raman-active modes of the LP phase is relatively easy (see discussion in Section 2 in SI), the symmetries of the experimental Raman-active modes of the HP phase of CsIO₃ are considerably more difficult to assign (see discussion in Section 2 in SI where a tentative assignment is proposed in Table S5 in SI). In any case, the number of Raman modes and their pressure dependence is consistent with the $Pmn2_1$ phase at least between 15 and 25 GPa.

The most interesting result for our purpose is the observed softening of the high-wavenumber modes in both R3m and $Pmn2_1$ phases of CsIO₃, at least up to 25 GPa, which leads to the almost complete closing of the phonon gap between the stretching and bending modes of the IO₃ units at 30 GPa. Noteworthy, the negative pressure coefficient of the high-wavenumber stretching modes in iodates has been already reported and explained by the increase of the short I–O bond distance and the loss of charge of these short primary delocalized resonant (covalent-like) I–O bonds.^{38–41} Also





Figure 5. Pressure dependence of the Raman modes of $CsIO_3$ up to 30.2 GPa. Symbols (lines) correspond to results from experiments (DFT calculations). Circles (triangles) correspond to the *R3m* (*Pmn2*₁) phase. Up to 15 GPa, red and blue colors correspond to A₁ and E modes of the *R3m* phase, respectively, while the green color represents modes that likely are not first-order modes. Above 15 GPa, red, green, and black colors correspond to A₁, B₁, and B₂ modes of the *Pmn2*₁ phase. Solid lines correspond to TO phonons while dashed lines correspond to the LO phonons. Note that LO phonons of A₁ modes in the *R3m* phase and of B₁ modes in the *Pmn2*₁ phase are not observed, especially at low wavenumbers, due to the negligible LO-TO splitting.

noteworthy, the soft phonon behavior upon approaching the cubic PV phase is consistent with what has been observed in many compounds.⁵

4.3. Chemical Bonding under Compression. The loss of charge of the short primary I–O bonds in both R3m and Pmn2₁ phases of CsIO₃ indicated by the softening of the highwavenumber modes in both phases suggests a pressureinduced change in chemical bonding in this compound. The change in chemical bonding is confirmed by the decrease in the number of electrons shared (ES) between two atoms (see theoretical details in Section 3 in the SI), according to QTAIM,¹⁶ plotted in Figure 4b. The ES values of the short I-O bonds in IO_3^- units of the R3m phase are larger than 2 at RP. This value is consistent with the delocalized resonant (covalent-like) character of the I–O bonds in IO_3^- units. The delocalized resonant bonding in IO₃⁻ units is similar to that of well-known NO3⁻ units, with the difference that there is a resonance between one single covalent bond and two double covalent bonds in IO₃⁻ units and a resonance between two single covalent bonds and one double covalent bond in the NO₃⁻ units due to the larger number of valence electrons in halogens than in pnictogens.

The delocalized resonant I–O bonding in IO_3^- units is also consistent with the I–O bond lengths. Notice that the bonds in the IO_3^- units in CsIO₃ have bond lengths of around 1.9 Å at RP. This value is intermediate between the larger single



Figure 6. Electron localization function (ELF) isosurfaces and ELF values along the different I–O bonds of CsIO₃ for the R3*m* phase at 0 GPa (a, e), the $Pmn2_1$ at 20 GPa (b, f) and 40 GPa (c, g), and the P4/nmm phase at 60 GPa (d,h). In (a), (b), (c), and (d), yellow isosurfaces correspond to the lone electron pairs (LEP) around I and O atoms, the short bonds are depicted with wide bars in red and magenta colors, and the long bonds are indicated with dashed blue lines.

covalent I–O bond and the shorter double covalent I–O bond, estimated to be 1.96 and 1.86 Å from the covalent radii,

respectively. 78,79 These values are in turn consistent with reported values in other iodates. $^{42,43,80-84}$

Interestingly, the ES values of the short primary I–O bonds decrease at the same rate as the ES values of the long secondary I···O bonds increase in both R3m and $Pmn2_1$ phases. This concomitant ES decrease of the short bonds and the ES increase of the long bonds can be interpreted as a multicenter interaction in which there is a charge transfer from the primary, short I–O bonds toward the secondary, long I···O bonds. The charge transfer is a consequence of the *trans influence* of the secondary bond into the primary bond that occurs as a prior step (stage 2 of 3) to the formation of multicenter bonds (stage 3) in the process of formation of multicenter bonds according to the recent unified theory of multicenter bonding.^{14,15}

According to the mentioned theory, the charge transfer ends once the two I–O bond types (short and long) equalize and form the multicenter bonds (stage 3).^{14,15} From that point on, all I–O bond lengths tend to decrease with pressure, and the soft high-wavenumber modes disappear; i.e., high-wavenumber vibrational modes (related to stretching vibrations) show a positive pressure coefficient. This is exactly what happens once the *P4/nmm* phase crystallizes above 45 GPa, as shown by our theoretical simulations (see Figure S7 in SI).

According to our ES calculations, the formation of the tetragonal P4/nmm phase above 45 GPa leads to the formation of 2D electron-deficient multicenter bonds (EDMBs) in the ab plane of the tetragonal PV structure (see Figure 3b) that come from the equalization of short and long bonds in the ac plane of the orthorhombic PV structure (see Figure 3a). The EDMBs are characterized by ES values close to 1, unlike the ERMBs, which typically show values of ES well above 1.4.^{14,15} In the case of the I–O EDMBs within the IO_{5+1} units of the P4/nmm phase, the ES value is ca. 1.25 because of the partial resonant bonding still present in this phase. As already explained, this resonant bonding explains why the I-O bonds within the IO₃ units of the R3m phase have ES values well above 2 (typical of resonant bonds). In addition to the two EDMBs, there are two additional bonds along the *c*-axis in the IO_{5+1} units of the P4/nmm phase. The fifth I–O bond, which is the shortest I-O bond, still preserves part of the delocalized resonant character already present at the LP phase, while the sixth I-O bond is the longest one and still preserves a partial noncovalent character similar to those of the LP phase. Note that the bond length of the four I-O EDMBs in the *ab* plane of the P4/nmm phase in CsIO₃ is ca. 1.95 Å, which is similar to the value of the long bonds present in I_2O_5 units⁸³ and I_4O_{12} units.⁸⁴ This suggests the possibility of observation of EDMBs in some iodates at RP; a feature worthy to be studied in future works.

The change in the I–O bonds in CsIO₃ across its successive pressure-induced crystalline phase transitions can be better understood by the analysis of the electron localization function (ELF). At the left of Figure 6a–d, we show the ELF isosurfaces in the three crystalline structures at four different pressures, while at the right of Figure 6e–h we show the corresponding ELF values along the different I–O bonds present in each phase. In the R3m phase at 0 GPa, the I atom shows a basin (corresponding to the iodine LEP) oriented along the axis perpendicular to the plane defined by the three O atoms of the IO₃ units. At this pressure, the O atoms show a basin which is intermediate between a toroidal and quasi-circular basin around each O atom (corresponding to the electrons forming part of the LEPs and the delocalized electrons of the resonant bonding). From the point of view of the IO₆ units, the R3m

phase is characterized by two different bonds, the short primary 3-fold degenerated intramolecular I-O bonds of the IO₃ units and the long secondary 3-fold degenerated intermolecular I-O interactions between the IO3 units. In Figure 6e, the ELF values of the short primary I-O bonds at RP show a weak minimum between I and O atoms (with a value of ca. 0.7 around the value of 0.5 for the normalized distance). This shape of the ELF profile (without a welllocalized minimun) is not expected for a covalent bond (with a well-localized minimum) and is consistent with the delocalized resonant (covalent-like) character of the I-O bonds in the R3m phase at RP. On the other hand, the ELF value of the long secondary I–O interactions exhibits a marked minimum between I and O atoms (around 0.56) whose value close to 0.1 evidences the weak noncovalent character of the secondary interaction at RP.

In the $Pmn2_1$ phase at 20 GPa, the I atom shows a LEP is still oriented as in the R3m phase (Figure 6b), while all O atoms show more defined toroidal basins oriented perpendicular to the more defined linear I-O-I bonds (the I-O-I angle changes from 168° at RP to 175° at 20 GPa). From the point of view of the IO₆ units, there are four different I-O bonds in this phase, the short and long bonds along the b axis and the 2-fold degenerate short and long bonds in the *ac* plane; however, the difference between the short/long bonds along the b axis and in the ac plane cannot be distinguished (see Figure 4a). Consequently, they show ELF values (Figure 6f) similar to those of the R3m phase, with the difference that the long bonds are stronger than at RP (the minimum of the ELF changes from ca. 0.1 at RP to ca. 0.3 at 20 GPa). In the Pmn2₁ phase at 40 GPa, the I atom shows a LEP reoriented along the secondary I-O bond in the b-axis, while the LEPs of all O atoms in the ac plane show almost a perfect toroidal form (Figure 6c). From the point of view of the IO_6 units, there are four different I–O bonds in this phase at 40 GPa, the short and long bonds along the b axis and the 2-fold degenerate short and long bonds in the ac plane. The ELF value of the short and long bonds along the b axis (axial bonds) still show the same resonant delocalized and noncovalent characters as at 20 GPa, respectively (see red and blue symbols in Figure 6g). On the other hand, the short and long bonds in the ac plane (equatorial bonds) show ELF values that tend to approach each other; i.e., there appears a minimum in the short bonds (consistent with a weakening of the intramolecular bonds), and the ELF value at the minimum of the long bonds increases (showing a strengthening of the intermolecular bonds).

Finally, in the P4/nmm phase at 60 GPa, the I atom still shows a LEP oriented along the secondary I–O bond in the *c*axis, while the LEPs of all O atoms show a toroidal basin (Figure 6d). The toroidal basins are totally symmetric (corresponding to the six electrons of three LEPs) for the O atoms located in the *ab* plane, while are asymmetric for the O atoms along the c axis. This can be explained from the point of view of the IO₆ units because there are three different I-O bonds in this phase, the short and long bonds along the c-axis and the 4-fold degenerate bond in the *ab*-plane (see Figure 4a). The ELF value of the short and long bonds along the *c*-axis still shows the same delocalized resonant and noncovalent character as at 0 GPa, respectively, although the strength of the secondary bond has notably increased. Interestingly, the four-degenerate bond in the *ab*-plane shows intermediate ELF values between those of the primary delocalized resonant bonds and the secondary noncovalent bonds (Figure 6h).



Figure 7. 2D map of the number of electrons shared (ES) vs the normalized number of electrons transferred (ET). The ES and ET values of the I– O bonds in the different phases of $CsIO_3$ at different pressures are indicated. When several similar bonds are present the average ES and ET values are represented.

These ELF values correspond to the EDMBs formed in the ab plane; therefore, it can be speculated that these are the ELF values that will be reached by all six I–O bonds (forming three EDMBs along the three directions of space) in the regular IO_6 unit of the cubic PV phase of CsIO₃ once PIS takes place along the *c*-axis of the tetragonal phase at much higher pressures. In fact, an almost total PIS has been found in our calculations for $RbIO_3$ (see Figure S8 in SI). This compound goes from the R3m phase at RP toward the quasi-cubic R-3m phase at 60 GPa. At that pressure, all bonds are equal up to the fourth decimal place, so the simulated phase can be considered as a very subtle distortion of the cubic Pm-3m phase. In RbIO₃, the almost total PIS leads to the formation of EDMBs in all three directions, so the LEP corresponding to the I atom is now symmetrically distributed around I atoms as it corresponds to an inactive LEP, whose (s-type) electrons do not participate in bonding. At the same time, the LEPs of O atoms along the three spatial directions show perfectly symmetric toroidal basins in the plane perpendicular to the I–O–I bond lines.

It must stressed that square pyramidal PbBr₅ units, similar to the units found in the P4/*nmm* phase of CsIO₃, have been recently found in 1D chains of lead bromide at RP.⁸⁵ The Pb– Br bonds in the square basis of the pyramidal unit are longer (average 3.04 Å) than the Pb–Br bond perpendicular to the square basis (2.71 Å). This suggests that this last bond is a covalent bond, while those of the square basis are EDMBs similar to those found in the PbBr₆ units of the *Pnma* phase in the distorted PV RbPbBr₃ (average ca. 3.05 Å).⁸⁶

The pressure-induced evolution of the I–O bonds in the distorted PV CsIO₃ across its successive crystalline phase transitions can be monitored using the ES vs ET map (Figure 7). For this purpose, the Bader charges, along with the ES and ET values for the short I–O bonds in various phases and pressures are provided in Table S6 in SI. In the rhombohedral R3m phase, the three short I–O bonds at 0 GPa are located slightly above the red region, indicating their delocalized resonant (covalent-like) character. In the orthorhombic $Pmn2_1$

phase, the short I–O bonds remain within the red region but shift closer to the green region (where EDMBs are located) as pressure increases (see different values for the three short bonds at 20 and 40 GPa). In the tetragonal *P4/nmm* phase at 60 GPa, the ES values of the two types of short I–O bonds behave differently: the bond along the *c* axis remains in the red region, while those in the *ab* plane move into the green region. This means that the short I–O bonds in the *ab* plane within the IO_{5+1} octahedra are no longer covalent bonds and have transformed into EDMBs.

The trend of chemical bonds in CsIO₃ and RbIO₃ under compression shown in Figures 6, 7 and S8 suggests that in the cubic Pm-3m phase of AIO₃ compounds the ES value of all I-O bonds would be the same, and similar to those already present in the *ab* plane in the tetragonal PV phase of CsIO₃. In other words, all I–O bonds in the regular IO₆ octahedra in the hypothetical cubic PV phase would be EDMBs. This result is confirmed in our calculations of the ES values of the quasicubic R-3*m* phase of RbIO₃ at 60 GPa (see Table S7 in SI). Note that this result is consistent with the claim of Wuttig and co-workers, who found metavalent (electron-deficient) bonds in the simulated cubic PV structure of CsPbX₃ (X = F, Cl, Br, I) lead halides.¹¹ In this regard, we recall that we consider that metavalent bonds do not exist since they are linear (or quasilinear) combinations of concatenated 3c-2e EDMBs, as explained in the unified theory of multicenter bonding.^{14,15}

The unified theory of multicenter bonding allows us to explain the tendency to equalization of short and long I–O bonds (Figure 2a) and the softening of the high-wavenumber vibrational modes (Figure 5) in the R3m and $Pmn2_1$ phases of CsIO₃ since there is a multicenter interaction taking place in the whole pressure range from RP to 45 GPa. In fact, the decrease of all bonds (Figure 4a) and the hardening of the high-wavenumber vibrational modes (Figure 57) above 50 GPa indicates that this multicenter interaction finishes above 45 GPa when EDMBs are formed in the *ab* plane of the tetragonal phase (stage 3). Note that the charge transfer from



Figure 8. Pressure dependence of the bandgap (a), dielectric constant (b), and Born effective charges of Cs (c), I (d), and O (e) atoms in the different phases of $CsIO_3$. A metallization is predicted at pressures close to that of the $Pmn2_1$ -P4/nmm phase transition; i.e., upon formation of EDMBs. Similarly, an extraordinary increase of the dielectric constant and Born effective charges occur near the phase transition, as it occurs in pnictogens, chalcogens, and phase change materials when EDMBs are formed.

the short I–O bonds toward the long I \cdots O bonds, shown in Figure 4b, indicates that the multicenter bonds have an electron-deficient character as suggested by the multicenter bonding theory.^{14,15}

We have also to note that the multicenter interaction should persist along the c-axis of the tetragonal phase as pressure increases beyond 50 GPa because the PIS is not completed along this axis; however, our calculations show that the multicenter interaction is no longer present in the P4/nmm phase of CsIO₃. We may speculate that the lack of trans influence of the secondary bond (along the c-axis of the tetragonal phase) into the primary bond (along the same caxis) is caused by steric effects related to the large volume of the Cs atom that avoids further contraction of the structure along the c-axis. Note that such an effect is not observed in RbIO₃ (Figure S8). In fact, the lack of multicenter interaction along the c-axis in the P4/nmm phase of CsIO3 is already observed along the *b*-axis of the $Pmn2_1$ phase above 20-25 GPa. Above this pressure range, it occurs the separation of the degenerated short and long bonds into the bonds of the ac plane (that will become EDMBs) and the short and long bonds along the *b*-axis (that will remain as covalent and noncovalent interactions, respectively). This separation leads to a contraction of the short bond along the b axis (see blue squares in Figure 4a) resulting in an upturn in the corresponding vibrational modes, which show a sudden increase in wavenumber (see experimental and theoretical vibrational modes around 700 and 600 cm⁻¹, respectively, in Figures 5 and S7). Moreover, this contraction of the short bond along the *b*-axis of the $Pmn2_1$ phase is consistent with the

expected contraction of the delocalized resonant I–O bond (see red line in Figure S5 in SI whose extrapolation to RP leads to a value close to 1.90 Å as the I–O bonds at RP).

4.4. Electronic Behavior under Compression. It is noteworthy that the formation of EDMBs in the tetragonal PV phase of CsIO₃ is consistent with a pressure-induced change in the simulated electronic properties of the distorted PV (Figure 8). In particular, an increase of the average dielectric constant, ε , and the Born effective charges, Z^* , are observed due to the pressure-induced "metallization", related to the closing of the bandgap, taking place between 40 and 50 GPa. Note that our calculations, which underestimate the bandgap, predict the closing around 43 GPa, so the experimental bandgap closure is expected at higher pressures. A similar evolution of these properties was observed in pnictogens, chalcogens, and related phase change materials under compression upon the formation of EDMBs.^{12,14,15,45,87} In fact, high values of dielectric constants, Born effective charges, and low thermal conductivity have been discussed for many PVs, especially for BaTiO₃. Note that metallization (closing of bandgap) is predicted when EDMBs are formed in the *ab* plane of the *P4/nmm* phase and that the values of dielectric constant and Born effective charges along the *c*-axis of the *P*4/*nmm* phase do not show high values near the phase transition unlike what happens to the components in the *ab* plane. It must be noted that the metallic or quasi-metallic behavior of the tetragonal PV phase of CsIO₃ is similar to that of the tetragonal (P4/mmm) PV phase of La_{0.5}Ba_{0.5}CoO₃⁸⁸ and the ordered tetragonal (P4/ *mmm*) PV phase of LaBaCo₂O₆ which also show CoO₅ units⁸⁵ (see also chapter 4 in ref 3).

As a final comment regarding the comparison of perovskite CsIO₃ and phase change materials, it is interesting to note that the pressure-induced hypercoordination of I atoms in CsIO₃ is similar to that found in SnSe. In the orthorhombic Pnma phase (SG No. 62) of SnSe at RP, Sn atoms are 3-fold coordinated as I in the R3m of CsIO₃. Around 10 GPa, SnSe undergoes a phase transition toward the orthorhombic *Cmcm* (or *Bbmm*) phase (SG No. 63)^{90,91} and the *Cmcm* phase is characterized by square pyramidal $SnSe_5$ units similar to those IO_{5+1} of Figure 3b. An increase of the short covalent Sn-Se bonds in the *ab* plane occurs in the *Pnma* phase, while the covalent Sn-Se bond along the c axis compresses in a normal way.⁹⁰ A softening of the high-wavenumber vibrational modes related to Sn-Se bonds in the *ab* plane is also observed.⁹¹ A change is observed once the Cmcm phase is reached. All Sn-Se bonds compress in this phase and all vibrational modes show a positive pressure coefficient. This change has been ascribed to the formation of EDMBs in the square plane of the SnSe₅ unit, while a covalent Sn-Se bond (the shortest bond) still persists along the *c*-axis of the *Cmcm* phase.¹⁵

4.5. Trends in AlO₃ Compounds under Compression. It has been already suggested that phase transitions in the distorted PV structures of AIO_3 (A = K, Rb, Cs, Tl) compounds would occur at smaller pressures under non-hydrostatic conditions than under hydrostatic conditions, as indeed found in KIO_3 .³⁷ In this context, a change in the pressure coefficients of the high-wavenumber phonons was already observed in KIO_3 above 14 GPa³⁷ despite our calculations for this compound in the *R*3*m* phase do not support a PIS under hydrostatic conditions until ca. 35 GPa (see Figure 9). Note that the I–O bond distances in the simulated *R*3*m* phases of AIO_3 (A = K, Rb, Cs, Tl) compounds show a PIS in all of them (see Figure 9).



Figure 9. Pressure dependence of the simulated short (intramolecular) I–O and long (intermolecular) I···O bond distances in the simulated R3m phases of several AIO₃ (A = K, Rb, Cs, and Tl) compounds.

Our results confirm that AIO_3 (A = K, Rb, Cs, Tl, NH₄) compounds undergo a PIS and tend to form the cubic PV structure at HP. The complete PIS does not occur in CsIO₃, which remains with a tetragonal *P4/nmm* structure even up to 70 GPa. However, the complete PIS happens already at 60 GPa for RbIO₃, which shows a quasi-cubic rhombohedral *R*-3*m* phase (Figure S8). Therefore, we conclude that RbIO₃ and

TlIO₃ are excellent candidates to verify experimentally the formation of EDMBs during the formation of the PV structure at HP. It can be observed that there is a correlation between the PIS pressure of AIO₃ compounds and the ionic size radius, r, of the A cation ($r_{\rm K} < r_{\rm Rb} \approx r_{\rm Tl} < r_{\rm Cs}$).⁹² This result supports our idea that the A cation acts as a stabilizing unit of the PV structure and that the large ionic size of Cs is behind the lack of full PIS of CsIO₃ as previously commented.

It is important to notice that the PIS is associated with a pressure-induced polymerization (PIP) process of the IO_3 units. The PIP process of the IO_3 units (monomers) leads to the formation of concatenated IO_{5+1} units along two spatial directions (in CsIO₃) or IO_6 units along three spatial directions (in other AIO₃ compounds). Consequently, the polymerization of the IO_3 units results in the formation of EDMBs at HP in both the tetragonal and (quasi-) cubic phases of AIO₃ compounds.

A simple way of understanding the PIP process of IO₃ units to yield IO₆ units in AIO₃ compounds at HP is shown in Figure 10. To avoid complications related to the change of space group of the crystalline structure and bond angles, etc., we could assume that AIO₃ compounds at RP have a cubic unit cell of lattice parameter a (a_0 at RP) in which isolated IO₃ units (monomers) exhibit three short covalent I–O bonds (of bond length *d*) directed along the *a*, *b*, and *c*-axes, respectively. This means that all O–I–O angles are 90°. Let us also assume for simplicity that the IO₃ units are characterized by short primary covalent I–O bonds with bond length $d = a_0/3$ at RP. This configuration means that each I atom shows also three long secondary noncovalent I-O interactions (along each of the three axes) with bond length $d' = 2a_0/3$ at RP. This situation is described in Figure 10a, where the two types of I-O bonds lead to asymmetric IO₆ octahedra. Now let us imagine that at HP this cubic cell compresses (the lattice parameter *a* decreases) and we observe that *d* enlarges while *d*' compresses until both I–O bond lengths are d = d' = a/2 at a certain pressure (the phase transition pressure, $P_{\rm T}$). This situation is described in Figure 10b, where there is only one type of bond resulting in symmetric IO₆ octahedra characteristic of the PV structure obtained after the PIP process of IO₃ units. To understand the PIP process of IO₃ units from the point of view of the symmetrization of the two types of I-O bonds, the behavior of the d and d' values (in relation to a) with increasing pressure is shown in Figure 10c. In this simplified way, we can understand the PIP process that transforms all isolated IO₃ units (belonging to asymmetric IO₆ units) into concatenated IO₃ units (belonging to symmetric IO_6 units). This PIP process explains how the cubic PV structure is obtained in AIO₃ compounds under compression.

In summary, all AIO₃ (A = K, Rb, Cs, Tl, NH₄) compounds crystallize at RP in strongly distorted perovskite structures characterized by IO₃ units with delocalized resonant (covalentlike) I–O bonds, resulting from the strong distortion of IO₆ polyhedra due to the strength of the iodine LEP. As pressure increases, the LEP becomes progressively less active, as found in many studies.^{17–23} Our experiments and calculations show that AIO₃ (A = K, Rb, Cs, Tl, NH₄) compounds tend to symmetrize under compression, forming cubic PVs or slightly distorted PVs. These structures have regular or slightly distorted IO₆ units with EDMBs; i.e., 2c–1e I–O bonds that come from the PIP process of the IO₃ units leading to IO₅₊₁ and IO₆ units via concatenated linear or quasi-linear 3c–2e bonds in 2D and 3D, respectively. Therefore, the PV structure



Figure 10. Simplified scheme of the pressure-induced polymerization (PIP) process of IO₃ units in AIO₃ compounds leading to the IO₆ units of the cubic perovskite (PV) structure. Big blue atoms represent I atoms and small red atoms represent O atoms. To focus on the PIP process of IO₃ units, *A* atoms (inside the cubes) are not shown. (a) Room pressure (RP): Cubic structure of a strongly distorted PV with highly asymmetric IO₆ units due to the existence of three short (solid) and three long (dashed) I–O bonds characteristic of isolated IO₃ units. (b) High pressure: Cubic PV structure showing symmetric IO₆ units with six equal I–O bond lengths. (c) Pressure dependence of the *a* lattice parameter and the three short (*d*) and three long (*d*') I–O bond distances (in terms of *a*). The value of *a* at RP, *a*₀, has been taken as 4 Å, and the *d* and *d*' values at RP have been taken as $a_0/3$ and $2a_0/3$, respectively. P_T indicates the phase transition pressure at which the cubic PV structure is reached and the PIP process ends. Note the different compression of bond lengths before and after P_T ; i.e., before and after EDMBs in IO₆ units are formed.

to be formed in AIO_3 compounds at HP can be considered as a PIP process of the IO_3 units to form IO_6 units. That polymerization involves a change in chemical bonding from delocalized resonant (covalent-like) I–O bonds in the IO_3 units toward 3D I–O EDMBs in the IO_6 units. In CsIO₃, the tetragonal perovskite is found above 45 GPa with well-defined 2D I–O EDMBs in the *ab*-plane. Along the *c*-axis a weakened covalent bond and strengthened noncovalent bond still persist in this tetragonal phase and the cubic PV structure is not observed even up to 70 GPa likely due to steric repulsions related to the size of Cs atom. On the contrary, full PIS leading to a cubic PV structure is expected in AIO_3 compounds with A^+ cations showing a smaller ionic radius than Cs.

5. CONCLUSIONS

Through a joint HP experimental and theoretical study of the distorted perovskite $CsIO_3$, we have shown that the rhombohedral (R3m) phase at room conditions undergoes two pressure-induced phase transitions up to 70 GPa. The first transition occurs around 14 GPa, yielding an orthorhombic phase ($Pmn2_1$), while the second one, leads to a tetragonal perovskite phase (P4/nmm). These transformations precede a potential transition to the cubic (Pm-3m) perovskite structure that could happen at pressures well above 70 GPa.

Our study reveals a gradual transformation of the initial IO_3 units of CsIO₃ into slightly distorted IO_{5+1} units as the tetragonal (quasi-cubic) phase is approached; i.e., a pressureinduced polymerization of the IO₃ units is observed that leads to I hypercoordination. Concomitantly, the I–O bonds change from the short primary delocalized resonant (covalent-like) bonds in IO₃ units toward electron-deficient multicenter bonds (EDMBs) as the polymerization proceeds under compression. In other words, I–O bonds become 2c–1e EDMBs in IO₅₊₁ units of the tetragonal PV phase of CsIO₃ due to the progressive polymerization of IO₃ units. The equalization of bond distances and the hardening of the originally soft highwavenumber Raman-active modes, once the bond distances equalize, give support to the formation of EDMBs, in agreement with the recently proposed theory of multicenter bonding.^{14,15}

The results of this work can be extrapolated to the whole family of AIO_3 (A = K, Rb, Cs, Tl, NH₄) compounds, where EDMBs will be developed at high pressure as the cubic perovskite structure is approached upon polymerization of IO_3 units, being RbIO₃ and TlIO₃ ideal candidates to observe the EDMBs in the three directions of space once the complete pressure-induced symmetrization of the perovskite structure occurs due to the complete pressure-induced polymerization process of the IO₃ units.

The results of this work, together with previous findings indicating that lead halide perovskites feature electron-deficient metavalent bonds¹¹ (here understood as EDMBs^{14,15}), suggest that the cubic perovskite ABX_3 structure of main-group elements and their slightly distorted variants is fundamentally governed by a combination of ionic bonds (between the *A* cation and *X* anions) and EDMBs (between the *B* cation and *X*

anions). It has been considered that oxide perovskites, like $BaTiO_3$, do not show this type of bonding.¹¹ This could lead to think that oxide perovskites in general do not show this type of bonding. In contrast, we have shown here that this unconventional bonding is present in AIO_3 perovskites. This unconventional bonding interplay may account for many of the extraordinary properties found in main-group perovskites, which cannot be explained within the framework of classic (covalent, ionic, and metallic) bonding. We hope that our findings will stimulate more studies on the chemical bonding in other perovskites (w/o main-group elements) to evaluate the extent to which the concept of EDMB can be applied in perovskites.

ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acs.chemmater.5c00877.

Details of the structural, vibrational, and chemical properties of $CsIO_3$ and $RbIO_3$ under compression; equations of state of the different phases of $CsIO_3$; pressure dependence of experimental and theoretical atomic parameters in the orthorhombic phase; and tables with structural, vibrational, and chemical data of the different phases of $CsIO_3$ and $RbIO_3$ (PDF)

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Notes

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