Metal Halide Perovskite and Perovskite-like Materials Through the Lens of Ultra-wideline $^{35/37}\text{Cl}$ NMR Spectroscopy

Diganta Sarkar,\textsuperscript{a} Riley W. Hooper,\textsuperscript{a} Abhoy Karmakar,\textsuperscript{a} Amit Bhattacharya,\textsuperscript{a} Arkadii Pominov,\textsuperscript{a} Victor V. Terskikh,\textsuperscript{b} and Vladimir K. Michaelis*\textsuperscript{a}

\textsuperscript{a}Department of Chemistry, University of Alberta, Edmonton, Alberta T6G 2G2, Canada
\textsuperscript{b}Metrology, National Research Council of Canada, Ottawa, Ontario K1A 0R6, Canada

*Corresponding author: vladimir.michaelis@ualberta.ca

ABSTRACT

With their exceptional optoelectronic features, metal halide perovskites (MHPs) are pushing the next wave of energy–related materials research. Heretofore, most solid–state nuclear magnetic resonance (NMR) investigations have focused on readily accessible nuclei. In contrast, the halogen environments have been avoided due to their challenging quadrupolar nature. Here, we report a rapid $^{35/37}\text{Cl}$ NMR strategy for MHPs, halide double perovskites (HDPs) and perovskite–inspired (PI) materials embracing ultra–wideline acquisition approaches at moderate and ultrahigh magnetic fields. The observed quadrupolar NMR parameters ($C_Q$ and $\eta$), supported by GIPAW–DFT computations, provide an analytical fingerprint revealing distinct features for chemically unique Cl environments sensitive to ion mixing, dimensionality, cell volume and Cl coordinating polyhedra. Moreover, we report resolution between two nearly identical and two distinct Cl environments of 3D and 2D Cs–based lead halide perovskites, respectively. These results reveal a strategy for a routine and robust spectroscopic approach to analyze local Cl chemical environments in metal halide perovskites that can be extended broadly to other halogen containing semiconductors.

Toc Graphic
Metal halide perovskites (MHPs) are emerging as an important class of semiconducting solids that exhibit impressive optical and electrical properties. Among them, lead halide perovskites (LHPs) are competing at the frontier of next–generation energy materials, leading in commercial advances for solar cells, LEDs, lasers, photodetectors, X–ray detectors, batteries, and photocatalytic applications. Recently, LHPs have reached power conversion efficiencies of 25.5%, surpassing that for the “gold standard” silicon solar cell. The three–dimensional (3D) LHPs are expressed using the general formula APbX₃ (A = Cs⁺, MA⁺ (methylammonium), FA⁺ (formamidinium); X = Cl⁻, Br⁻, I⁻), where the A–site cation resides in the cuboctahedral voids of a corner–sharing 3D network of [PbX₆]⁴⁻ octahedra. The zero–dimensional (0D) and two–dimensional (2D) LHPs are recent entrants, providing enhanced material stability, photoluminescent properties, and charge carrier recombination dynamics compared to their 3D analogues. Cesium–based 0D and 2D LHPs can be described by the general formula Cs₄PbX₆ and CsPb₂X₅, respectively, where 0D Cs₄PbX₆ consists of isolated [PbX₆]⁴⁻ octahedra surrounded by Cs⁺ and 2D CsPb₂X₅ has layers of face sharing [PbX₆]⁶⁻ side–bicapped trigonal prisms with Cs⁺ sandwiched between the layers.

Environmental concerns related to lead and phase instability remain as commercialization challenges for LHP–based photovoltaic technologies. Inevitably, group 14 cousin materials, viz., tin and germanium halide perovskites (GHPs) are gaining further attention in the search for lead–free alternatives, while halide double perovskites (HDPs) [A₂B'(III)B"(I)X₆] consisting of corner–sharing alternate [B'(III)X₆]³⁻ and [B"(I)X₆]⁵⁻ octahedra with A–site cations residing in the cuboctahedral voids have shown improved ambient stability, excellent bandgap tunability and appealing light–emission properties. Additionally, perovskite–inspired A₂B(IV)X₆ (B = Sn⁴⁺, Pd⁴⁺) compounds, being air–stable and environment–friendly with tailorable optoelectronic capabilities, may provide an alternative direction. Their structure (cubic, Fm̅3m) can be described as an ABX₃ perovskite structure with every other B–site removed. The crystal structures of several MHPs studied here are shown in Figure 1.

**Figure 1.** Crystallographic structures of lead–based 3D (a) FAPbCl₃, (b) MAPbCl₃, (c) CsPbCl₃, 2D (d) CsPb₂Cl₅, and 0D (e) Cs₄PbCl₆; (f) CsGeCl₃, (g) Cs₂Bi/InAgCl₆ (B' = Bi or In), and (h) Cs₂SnCl₆.
Though standard characterization tools, such as diffraction techniques, are traditionally used to characterize the average long–range structures of MHPs, complementary techniques are often needed to provide further insight into local structure, vacancies, dynamics, and nanodomains.\textsuperscript{28–30} Solid–state nuclear magnetic resonance (NMR) spectroscopy is a robust analytical method that may be used to inform on short– to medium–range structure and ion dynamics which has been shown to be a highly sensitive probe to understand the structure–property relationships displayed by MHPs.\textsuperscript{31–35} Typically, all the constituents of MHPs have sufficiently abundant and NMR–sensitive isotopes,\textsuperscript{34} which makes them perfect candidates for NMR analyses. Among all NMR–active nuclei in Cs–based MHPs, \textsuperscript{133}Cs (nuclear spin, $I = 7/2$, quadrupolar ($I > 1/2$) but with a small quadrupole moment such that quadrupolar effects are negligible) and \textsuperscript{207}Pb ($I = 1/2$) are the most intensely studied because of their relative accessibility compared to more difficult nuclei, and have contributed significantly to understanding of phase transitions,\textsuperscript{36,37} chemical exchange reactions,\textsuperscript{38,39} and phase segregation.\textsuperscript{40,41} On the other hand, quadrupolar halogen nuclei ($\textsuperscript{35}Cl, \textsuperscript{79}/\textsuperscript{81}Br, \textsuperscript{127}I$) are extremely useful in characterizing unique chemical environments, however challenging as resolution and sensitivity is often hampered due to second–order quadrupolar broadened lineshapes.\textsuperscript{42} The resulting quadrupolar interactions can be described by the nuclear quadrupole coupling constant ($C_Q$) and the asymmetry parameter ($\eta$). Recent halogen NMR and nuclear quadrupolar resonance (NQR) studies have demonstrated their utility in probing chemical structure and dynamics in solids.\textsuperscript{34,35,42–47} Amidst the quadrupolar halogen nuclei, \textsuperscript{35}Cl has a moderate quadrupole moment ($Q_m$) and high natural abundance (N.A.), making it a less challenging halogen nucleus for NMR studies compared to \textsuperscript{79}/\textsuperscript{81}Br and \textsuperscript{127}I (Table 1). While \textsuperscript{37}Cl is complementary to \textsuperscript{35}Cl, it is more challenging due to its lower gyromagnetic ratio and N.A.\textsuperscript{42,45} This approach is lending support to further expansion of the NMR toolbox into these exotic NMR active nuclei, as even \textsuperscript{73}Ge NMR spectroscopy has been shown to provide sensitivity of the octahedral distortion in CsGeX\textsubscript{3} (X = Cl, Br, I) perovskites.\textsuperscript{48}

<table>
<thead>
<tr>
<th>Halogen Nucleus</th>
<th>$I$</th>
<th>N.A. (%)</th>
<th>$Q_m$ (mb)</th>
<th>$\varepsilon$ / %</th>
</tr>
</thead>
<tbody>
<tr>
<td>\textsuperscript{35}Cl</td>
<td>3/2</td>
<td>75.78</td>
<td>−81.65</td>
<td>9.79</td>
</tr>
<tr>
<td>\textsuperscript{37}Cl</td>
<td>3/2</td>
<td>24.22</td>
<td>−64.35</td>
<td>8.15</td>
</tr>
<tr>
<td>\textsuperscript{79}Br</td>
<td>3/2</td>
<td>50.69</td>
<td>313.0</td>
<td>25.0</td>
</tr>
<tr>
<td>\textsuperscript{81}Br</td>
<td>3/2</td>
<td>49.31</td>
<td>261.5</td>
<td>27.0</td>
</tr>
<tr>
<td>\textsuperscript{127}I</td>
<td>5/2</td>
<td>100.0</td>
<td>−696.0</td>
<td>20.0</td>
</tr>
</tbody>
</table>

Table 1. Comparison of NMR properties of the quadrupolar ($I > 1/2$) halogen nuclei.\textsuperscript{42}
In this letter, we determine unique $^{35}$Cl (and $^{37}$Cl for CsPbCl$_3$) quadrupolar NMR parameters for leading hybrid/inorganic and multidimensional metal halide perovskites, double perovskites, and perovskite–inspired materials. We also demonstrate that rapid acquisition ($\sim 30$ min) of wideline NMR spectra is efficient and practical. We further discuss the highly receptive nature of $^{35}$Cl NMR spectroscopy, making it suitable to elucidate distinct Cl chemical environments that are sensitive to their polyhedron geometry and coordination. In addition, we discuss the practicality of resolving multiple Cl environments and detecting low Cl concentrations ($< 5\%$). Lastly, we demonstrate that a quantum chemical computational approach using gauge–including projector augmented waves$^{49}$ assisted density functional theory (GIPAW–DFT) reasonably predicts the $^{35}$Cl quadrupolar parameters making it a useful complementary technique.

![Figure 2. Non–spinning $^{35}/^{37}$Cl NMR spectra of 3D LHPs acquired at 21.1 T at room temperature. (a) Experimental WURST–CPMG (black) and simulated (blue) $^{35}$Cl NMR spectra of 3D (a) CsPbCl$_3$ and hybrid (b) MAPbCl$_3$ and (c) FAPbCl$_3$. The asterisk (*) and the dagger (†) denote the presence of PbCl$_2$ and an unidentified impurity in MAPbCl$_3$ and FAPbCl$_3$, respectively. (d) WURST–echo (violet) and quadrupolar–echo (red) $^{35}$Cl NMR spectra of CsPbCl$_3$ showing resolution between two nearly identical Cl sites with slightly different Pb–Cl–Pb bond angles ($\angle$); individual simulations of the $\angle \sim 161^\circ$ (brown) and $169^\circ$ (green) sites are shown above the overall simulated spectra (blue), and an enlarged view of the highlighted low–frequency shoulder is displayed. (e) Experimental quadrupolar–echo (red) and simulated (blue) $^{37}$Cl NMR spectra of CsPbCl$_3$ displaying resolution between the previously mentioned Cl sites, with the corresponding simulations shown with the assignments of $\angle \sim 161^\circ$ and $169^\circ$ sites indicated in the figure (brown and green, respectively). The highlighted low–frequency shoulder is enlarged.

Non–spinning $^{35}$Cl NMR spectra of 3D APbCl$_3$ ($A =$ Cs$^+$, MA$^+$, and FA$^+$) at 21.1 T, shown in Figure 2, display second–order quadrupolar–broadened line shapes due to the
distinct two-coordinate linear or near-linear Cl chemical environments present in these materials. The hybrid species MAPbCl$_3$ was synthesized using two synthetic routes, inverse temperature crystallization (ITC) and mechanochemical synthesis (MCS) (Figure S1a). All other bulk LHPS were prepared by MCS, and their NMR parameters are discussed here. MAPbCl$_3$ crystallizes in a cubic space group $Pmar{3}m$ (Figure S1a), with each Cl atom covalently bonded$^{50}$ with two Pb atoms in a collinear geometry.$^{51}$ The $^{35}$Cl NMR spectrum of MAPbCl$_3$ (ITC) at 7.05 T was acquired using a Hahn–echo$^{52}$ with 13 variable offset cumulative spectra (VOCS)$^{53}$ steps (7.5 h/step) as well as a wide-band, uniform rate, and smooth truncation Carr–Purcell–Meiboom–Gill (WURST–CPMG)$^{54,55}$ pulse sequence with five VOCS steps (25 min/step) which offered a ~47-fold savings in time (Figure S4).

Parameters of MAPbCl$_3$ (MCS), extracted from the $^{35}$Cl WURST–CPMG spectrum, were $C_Q = 16.34$ MHz, $\eta = 0$, and isotropic chemical shift ($\delta_{iso}$) of 155 ppm. Likewise, FAPbCl$_3$, which is isostructural (Figure S1a) with a slightly larger unit cell volume$^{56}$ due to the larger FA$^+$ cation, has experimentally determined parameters of $C_Q = 17.46$ MHz, $\eta = 0$, and $\delta_{iso} = 160$ ppm. The effect of A-site mixing was investigated by acquiring a $^{35}$Cl WURST–CPMG spectrum of FA$_{0.50}$MA$_{0.50}$PbCl$_3$ (Figure S1a) at 7.05 T, and a $C_Q = 16.55$ MHz, $\eta = 0$, and $\delta_{iso} = 200$ ppm were determined (Figure S5f). The stable CsPbCl$_3$ assumes a lower-symmetry orthorhombic structure ($Pnma$) (Figure S1b) with Cl atoms coordinated to two lead atoms in two different non-collinear geometries (Pb–Cl–Pb bond angles, $\angle$ of ~161° and 169°, with a similar Pb–Cl bond distance of ~2.8 Å).$^{57}$ The WURST–CPMG spectrum fails to resolve these two slightly different Cl environments and the average NMR parameters for the observed $^{35}$Cl signal are $C_Q = 15.60$ MHz, $\eta = 0$, and $\delta_{iso} = 155$ ppm (Figure 2a). Utilizing a $^{35}$Cl WURST–echo$^{55}$ and quadrupolar–echo$^{58,59}$ experiment allowed us to resolve two nearly identical Cl sites with similar parameters $\eta = 0$ and $\delta_{iso} = 150$ ppm, and $C_Q = 15.57$ and 15.71 MHz for the $\angle$ ~161° and 169° environments, respectively (Figure 2d), supported by GIPAW–DFT computations. This observation was confirmed in a quadrupolar–echo experiment using $^{37}$Cl NMR. The two Cl sites were resolved with $C_Q \approx 12.27$ and 12.38 MHz for $\angle$ ~161° and 169°, respectively, having the same $\eta = 0$ and $\delta_{iso} = 150$ ppm (Figure 2e). The 1.27–fold reduction in the quadrupole coupling constant for both $^{37}$Cl sites is consistent with the ratio of their respective quadrupolar moments (Table 1). To obtain resolved spectra efficiently, one need not obtain the full spectrum using an echo technique as shown here. Having determined the positions of the horns in these spectra via WURST–CPMG, higher resolution spectra of just these two regions can be acquired using an echo technique. The results obtained for CsPbCl$_3$ are comparable with the recent study by Piveteau et al. where bulk CsPbCl$_3$ (monoclinic, $P2_1/m$) was found to have $C_Q = 15.48$ MHz and $\eta = 0.43$ These three results make it apparent that all these structures contain axially symmetric Cl electric field gradient (EFG) tensors where $V_{ZZ}$ is aligned along the direction of Pb–Cl bonds and CsPbCl$_3$ has the smallest $C_Q$. In fact, the formula unit cell volume ($V/Z$) shrinks from FAPbCl$_3 >$ FA$_{0.50}$MA$_{0.50}$PbCl$_3$ (Table S5) > MAPbCl$_3 >$ CsPbCl$_3$, influencing the EFG about Cl, whereby the associated $C_Q$ decreases in a near linear ($R^2 \sim 0.9$, Figure S2) relationship, suggestive of a secondary effect on the magnitude of the quadrupole
coupling, *vide infra*. In addition, the NMR spectra of these 3D LHPs were also recorded at a lower magnetic field of 7.05 T for comparative analysis (Figure S5 and S6).

**Figure 3.** Experimental (black) and simulated (blue) non–spinning $^{35}$Cl NMR spectra using the WURST–CPMG pulse sequence acquired at 7.05 T and room temperature. Lead–based (a) 0D Cs$_4$PbCl$_6$ (top), 2D CsPb$_2$Cl$_5$ (middle), and 3D CsPbCl$_3$ (bottom). The individual simulations for two different Cl environments in CsPb$_2$Cl$_5$ are shown above the simulated spectra for three–coordinate Cl ([3]$^{35}$Cl in brown) and four–coordinate Cl ([4]$^{35}$Cl in green). Lead–free (b) CsGeCl$_3$ (top), Cs$_2$BiAgCl$_6$ and Cs$_2$InAgCl$_6$ (middle), and Cs$_2$SnCl$_6$ (bottom). The asterisks (*) in (b) denote signals from an unidentified source.

Extending beyond the 3D LHPs, Figure 3a shows the non–spinning $^{35}$Cl NMR spectra acquired at 7.05 T of the 0D and 2D non–hybrid LHPs, found in trigonal ($R3c$)$^{60}$ and tetragonal ($I4/mcm$)$^{17}$ crystal symmetries (Figure S1b), respectively. The 0D Cs$_4$PbCl$_6$ perovskite has Cl atoms enclosed within a distorted Cs$_5$Pb quasi–octahedron$^{60}$ with the smallest $^{35}$Cl quadrupole coupling constant ($C_Q = 8.50$ MHz) within this family of compounds, with $\eta = 0.20$ and $\delta_{iso} = 150$ ppm (Figure 3a). The two distinct environments around Cl atoms, viz., three–coordinate ClPb$_3$ ([3]$^{35}$Cl) and square planar ClPb$_4$ ([4]$^{35}$Cl)$^{17}$ are clearly resolved at moderate (Figure 3a) and high (Figure S6d) magnetic fields with assignments supported by GIPAW–DFT computations. The [4]$^{35}$Cl in a planar environment has a $C_Q = 14.10$ MHz, $\eta = 0$, and $\delta_{iso} = 190$ ppm, while a decrease in the local coordination environment to [3]$^{35}$Cl in three-coordinate geometry yields NMR parameters of $C_Q = 9.00$ MHz, $\eta = 0.40$ and $\delta_{iso} = 170$ ppm. This decrease is attributed to the increased local symmetry about the Cl. A comparison of the NMR spectra recorded at 21.1 and 7.05 T for the low dimensional LHPs is shown in Figures S5 and S6. The unique
NMR parameters for the LHPs with different dimensions (above) warrant a closer look into other perovskite and perovskite–like metal chlorides.

To extend beyond lead–based perovskites, the chlorine environments of the GHP CsGeCl₃ and HDPs Cs₂BiAgCl₆ and Cs₂InAgCl₆ (Figure S1a) were studied via ³⁵Cl NMR spectroscopy (Figure 3b). The non–collinear geometry around the Cl atoms in CsGeCl₃ (rhombohedral, R₃m) with a Ge−Cl−Ge bond angle ~173° gives rise to an asymmetry parameter, η = 0.02, with a larger C_Q = 23.65 MHz and a sizeable change in the isotropic chemical shift (δ_iso = 850 ppm). In the case of the HDPs, both Cs₂BiAgCl₆ and Cs₂InAgCl₆ (cubic, Fm̅3m) have linear Cl environments (η = 0) with Ag−Cl−Bi or Ag−Cl−In coordination. The replacement of the B−site cation with a combination of B′ (Bi³⁺ or In³⁺) and B” (Ag⁺) cations in Cs₂BiAgCl₆ and Cs₂InAgCl₆ results in a significant impact on the ³⁵Cl NMR spectra with δ_iso values increasing to 550 and 950 ppm, respectively; deviation from linear symmetry results in increased C_Q values to 22.85 and 26.65 MHz, respectively. These results expand the Cl quadrupole coupling constants for two–coordinate linear (or near–linear) Cl species ranging here between 15 and 27 MHz depending on the medium−range B−site structure. The perovskite–inspired Cs₂SnCl₆ (cubic, Fm̅3m) compound has Cl atoms that are singly coordinated to Sn, resulting in the largest ³⁵Cl C_Q of 32 MHz (η = 0, δ_iso = 560 ppm) reported in this study (Figure S8).

Table 2. Experimental and GIPAW–DFT computed ³⁵Cl NMR parameters of metal halide compounds.

<table>
<thead>
<tr>
<th>Compound</th>
<th>CIF Ref.</th>
<th>CN</th>
<th>C_Q^{a,b} (MHz)</th>
<th>η^b</th>
<th>δ_iso^{a,b} (ppm)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td>±</td>
<td>±</td>
<td>±</td>
</tr>
<tr>
<td>Cs₂SnCl₆</td>
<td>[64]</td>
<td>1</td>
<td>32.0(3)c</td>
<td>−33.54</td>
<td>0.00</td>
</tr>
<tr>
<td>Cs₂InAgCl₆</td>
<td>[62]</td>
<td>2</td>
<td>26.65(7)</td>
<td>−29.02</td>
<td>0.00(1)</td>
</tr>
<tr>
<td>Cs₂BiAgCl₆</td>
<td>[63]</td>
<td>2</td>
<td>22.85(6)</td>
<td>−25.34</td>
<td>0.00(1)</td>
</tr>
<tr>
<td>CsGeCl₃</td>
<td>[61]</td>
<td>2</td>
<td>23.65(6)</td>
<td>−25.30</td>
<td>0.02(1)</td>
</tr>
<tr>
<td>FAPbCl₃</td>
<td>[56]</td>
<td>2</td>
<td>17.46(5)</td>
<td>−18.47</td>
<td>0.00(1)</td>
</tr>
<tr>
<td>MAPbCl₃</td>
<td>[51]</td>
<td>2</td>
<td>16.34(6)</td>
<td>−18.24</td>
<td>0.00(1)</td>
</tr>
<tr>
<td>CsPbCl₃</td>
<td>[57]</td>
<td>2</td>
<td>15.57(4)</td>
<td>−16.52</td>
<td>0.00(1)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>15.71(3)</td>
<td>−16.92</td>
<td>0.00(1)</td>
</tr>
<tr>
<td>Cs₂PbCl₅</td>
<td>[17]</td>
<td>3</td>
<td>9.00(7)</td>
<td>−8.59</td>
<td>0.40(2)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>14.10(5)</td>
<td>15.34</td>
<td>0.00(1)</td>
</tr>
<tr>
<td>Cs₄PbCl₆</td>
<td>[60]</td>
<td>6</td>
<td>8.50(6)</td>
<td>−8.92</td>
<td>0.20(2)</td>
</tr>
<tr>
<td>CsCl (ref.)</td>
<td>[65]</td>
<td>8</td>
<td>0.00(1)</td>
<td>0.00</td>
<td>0.00(1)</td>
</tr>
</tbody>
</table>

aSign of C_Q were not determined experimentally. bError representation is given as, e.g., 17.46 ± 0.05 MHz as 17.46(5) ppm. cReference.35

Unlike past NMR studies on the B−sites of Pb and Sn where chemical shift ranges correspond to distinct coordination geometries for these I = 1/2 nuclei, the ³⁵Cl chemical shifts of LHPs are rather crowded within a 40 ppm range. Together with the fact that each NMR signal spans thousands of ppm, this makes it nearly impossible to distinguish between multiple signals. It is therefore impractical to assess structural changes based on δ_iso alone. However, the ³⁵Cl quadrupolar parameters (C_Q and η) are
distinct for each compound (multidimensional LHPs, GHP, HDPS, PI) revealing that the unique EFG for each Cl site may be used to elucidate its unique coordination environment and geometry. Furthermore, while we are aware that overlap between the $C_Q$ values for $n$–dimensional ($n = 0, 2$ and $3$) LHPs can occur, discerning these unique sites in mixtures may be overcome by collecting and analyzing NMR spectra acquired at multiple–field strengths (Figure S7) or using both $^{35}\text{Cl}$ and $^{37}\text{Cl}$ probing nuclei (Figure 2).

![Figure 4](image_url)

**Figure 4.** (a) Comparison of the experimental and GIPAW–DFT computed $^{35}\text{Cl}$ $C_Q$ values for MHPs listed in Table 2. (b) Experimental $C_Q$ values for distinct $^{35}\text{Cl}$ chemical polyhedra of the samples with different coordination numbers and symmetry. The highly symmetric tetrahedral ($T_d$) and octahedral ($O_h$) geometry with $C_Q = 0$ MHz are displayed as reference.

A series of quantum chemical computations using GIPAW–DFT were performed starting from their corresponding crystallographic information files (CIF) to assess the effectiveness of computing $^{35/37}\text{Cl}$ NMR parameters in MHPs (Table 2). An area where further research efforts are needed, as we build a training set here, is to establish a predictive tool to screen NMR/NQR parameters to identify unique solutions when exploring ion substituted materials, unique chemical environments, or vacancies. Figure 4a shows that there is good agreement for the computed $|C_Q|$ values of MHPs when compared with experiment (slope $\sim 1$, $R^2 \sim 0.99$). The computed $\delta_{iso}$ values give rise to a slope of $2.92$ with an $R^2$ of $0.94$ for multidimensional LHPs (Figure S9a) and deviates further when the $\delta_{iso}$’s for GHP, HDPS and PI materials are considered. The agreement in the quadrupolar coupling constants reinforce the growing importance of experimental NMR and theoretical computations as complementary tools to assist our understanding of the X–site in these materials. From an *in–silico* perspective, the computed chlorine NMR parameters are sensitive to local structure and dimensionality. Furthermore, the ability to accurately calculate NQR parameters is important since the signals can be very difficult to locate. The computed $\eta$ values for the two hybrid LHPs (Figure S9b) deviate from the expected trend due to insertion of fixed–position hydrogen atoms within the input files. In reality, the organic molecules occupying the A–site (FA, MA) are dynamic, which result in discrepancies in their calculated $\eta$ values (Table S7 and S8).\textsuperscript{51,56} Figure 4b shows the general trend observed between Cl coordination number and $C_Q$ for all compounds studied here including CsCl, which has a $C_Q$ of zero due to its spherically
symmetric Cl environment. Examination of the local Cl environment for each perovskite reveals a series of unique polyhedra, where their local coordination environment and degree of polyhedral distortion guide the magnitude of the $^{35}$Cl $C_Q$. A similar effect has been reported for $^{23}$Na NMR spectra using a point charge model, whereby the degree of deviation from spherical symmetry and coordination results in sizeable changes to $C_Q$. The computed EFG ellipsoids around the $^{35}$Cl atoms of all the perovskites are displayed in Figure S10, where the largest principal component of the EFG tensor ($V_{ZZ}$) for all Cl atoms is aligned along their metal−Cl bonds except for four−coordinate Cl ($^{14}$Cl) in CsPb$_2$Cl$_3$.

To further assess the sensitivity and utility of ultra−wideline $^{35}$Cl NMR spectroscopy, the influence of Cl environments in cesium mixed−halide perovskites (CsPb(Cl$_x$Br$_{1-x}$)$_3$; $x = 0.03$, 0.33, and 0.67) were surveyed (Figure S1b). A 3% Cl substituted−CsPbBr$_3$ compound was detectable via the ultra−wideline method at 7.05 T. Fitting the $^{35}$Cl NMR spectrum revealed an experimental $C_Q$ of 16.5 MHz ($\eta = 0$ and $\delta_{iso} = 250$ ppm) which is an increase of ~1 MHz from the parent CsPbCl$_3$ phase (Figure 5b, top). This sensitivity test demonstrates the potential of this method in detecting Cl additives at low atomic %, such as methylammonium chloride (MACl) in MHP−based solar cells that is a common additive for improved stability and efficiency. Two additional mixed−halide CsPb(Cl$_x$Br$_{1-x}$)$_3$ ($x = 0.33$ and 0.67) perovskites were investigated and display sharp high frequency horns of the overall $^{35}$Cl quadrupolar broadened central transition (Figure 5b). The poor resolution renders the challenge to determine accurate $C_Q$ values, however it is still possible to perform a crude fit, enabling to track changes in

Figure 5. (a) The dependence of $C_Q$ values with the formula unit cell volume ($V/Z$; $Z = 4$) for the mixed−halide CsPb(Cl$_x$Br$_{1-x}$)$_3$ series. $C_Q$ values for $x = 0.33$ and 0.67 were obtained with respect to their $V/Z$ values from the linear equation ($C_Q = 0.0402(V/Z) + 8.5384$) constructed from two end points $x = 0.03$ and 1.0. (b) Comparison of non−spinning $^{35}$Cl NMR experimental (black) and simulated (blue; for $x = 0.03$ and 1.0) spectra of CsPb(Cl$_x$Br$_{1-x}$)$_3$ using WURST−CPMG pulse sequence recorded at 7.05 T and room temperature. The predicted $C_Q$ values for $x = 0.33$ and 0.67 were used to generate the simulated spectra shown by the dotted lines in red and green, respectively, using an average $\delta_{iso} = 200$ ppm and $\eta = 0$. The shift of the left horns towards higher frequency with a decrease in Cl concentration ($x$) are indicated with an arrow (orange) in the enlarged view on left in (b).
the quadrupolar coupling that is associated with an expansion of the unit cell, leading to changing the Cl chemical environment.

Similar to the mixed A–site cation alloys (FA\textsubscript{x}MA\textsubscript{1−x}PbCl\textsubscript{3}; x = 0, 0.50, and 1.0), where the \( C_Q \) displayed a linear dependence of the volume per formula unit (V/Z) (Figure S2), the influence of V/Z on \( C_Q \) as Cl substitutions occur was examined for the CsPb(Cl\textsubscript{3−x}Br\textsubscript{x})\textsubscript{3} series. Assuming a similar linear relationship between \( C_Q \) and V/Z (\( C_Q = 0.0402(V/Z) + 8.5384 \)) as the halide ions exchange for CsPbCl\textsubscript{3} and CsPb(Cl\textsubscript{0.03}Br\textsubscript{0.97})\textsubscript{3}, the \( C_Q \) values of mixed–halide perovskites with Cl concentrations x = 0.33 (16.34 MHz) and x = 0.67 (16.15 MHz) were extracted with respect to their refined V/Z values (Table S5) as shown in Figure 5a. The predicted \( C_Q \)'s were used to simulate the \(^{35}\text{Cl} \) NMR spectra and show reasonable agreement with our experimental results (Figure 5b). Therefore, the dominant structural parameter driving the unique \(^{35}\text{Cl} \) quadrupole couplings is indeed the coordination polyhedron as this would influence the electric field gradient; however, a secondary influence related to changes in the unit cell volume (contraction/expansion) further influences the quadrupole parameter as ions exchange. Unfortunately, the hybrid MAPb(Cl\textsubscript{3−x}Br\textsubscript{x})\textsubscript{3} (x = 0.25, 0.50 or 0.75) series suffers from poor sensitivity, inhibiting analysis. This unexpected outcome may be caused by shortened spin–spin relaxation (\( T_2 \)) characteristics that could arise from the dynamics at the X– (halide migration) or A–site (MA/FA mobility), or both\textsuperscript{31,72} These dynamic considerations are inhibited within the CsPbX\textsubscript{3} series due to the smaller unit cell that results when the smaller Cs\textsuperscript+ resides in the A–site leading to octahedral tilting of the B–site\textsuperscript{31,39,72}.

To conclude, we demonstrate the merit of \(^{35}/^{37}\text{Cl} \) NMR spectroscopy as a fast and sensitive structural probe for characterizing unique Cl coordination geometries in various MHPs. Distinct EFG tensors for \(^{35}\text{Cl} \) sites in these MHPs exhibit a sizeable range of \( C_Q \) values between 8.50 to 32 MHz, which have been shown to extend to over 40 MHz for other Cl–containing systems, determined using moderate field strengths\textsuperscript{71–75}. We highlight that the range of chemical shifts of 40 ppm for LHPs is very small with respect to the experimental spectra that span thousands of ppm in these materials due to sizable second–order quadrupolar broadening, thereby making \( C_Q \) a more efficient and sensitive parameter in discriminating specific Cl chemical environments. Resolution between two nearly identical crystallographic Cl environments of 3D Cs–based lead halide perovskite using both \(^{35}\text{Cl} \) and \(^{37}\text{Cl} \) NMR reveals the profound capabilities of this approach to discern between two similar Cl environments. The \(^{35}\text{Cl} \) NMR spectrum for the 2D lead halide perovskite reveals two crystallographically distinct Cl environments at moderate fields. In general, \( C_Q \) decreases with increasing Cl coordination number and is influenced by the degree of polyhedral distortion. Overlapping Cl environments may be resolved using a combination of multiple–field strengths, isotope selection, and NMR experimental methods. These methods can be further extended to materials with low Cl substitution or cation/anion modifications. Finally, the GIPAW–DFT results helped reliably estimate the quadrupolar NMR parameters in Cl–containing MHPs. This will be used to optimize experimental wideline acquisition parameters, to suggest assignments as well as to narrow the search space in NQR applications.
ASSOCIATED CONTENT
Supporting Information
Additional discussion about experimental techniques and GIPAW−DFT computations with Tables S1–S11 and Figures S1–S10 are available in the Supporting Information (PDF).

AUTHOR INFORMATION

Corresponding Author
Vladimir K. Michaelis – Department of Chemistry, University of Alberta, Edmonton, Alberta T6G 2G2, Canada
Email: vladimir.michaelis@ualberta.ca

Authors
Diganta Sarkar – Department of Chemistry, University of Alberta, Edmonton, Alberta T6G 2G2, Canada
Riley W. Hooper – Department of Chemistry, University of Alberta, Edmonton, Alberta T6G 2G2, Canada
Abhoy Karmakar – Department of Chemistry, University of Alberta, Edmonton, Alberta T6G 2G2, Canada
Amit Bhattacharya – Department of Chemistry, University of Alberta, Edmonton, Alberta T6G 2G2, Canada
Arkadii Pominov – Department of Chemistry, University of Alberta, Edmonton, Alberta T6G 2G2, Canada
Victor V. Terskikh – Metrology, National Research Council of Canada, Ottawa, Ontario K1A 0R6, Canada

Notes
The authors declare no competing financial interests.

ACKNOWLEDGEMENTS
We acknowledge the Natural Science and Engineering Research Council of Canada (DG and CREATE−ATUMS), Canada Research Chairs Program, Canada Foundation for Innovation and the Government of Alberta for research support. D.S. is supported by an Alberta Innovates Graduate Fellowship. R.W.H. is supported by an NSERC CGSD3, and Alberta Innovates Graduate Fellowship. Access to the 2.11 T NMR spectrometer was provided by the National Ultrahigh−Field NMR Facility for Solids (Ottawa, Canada), a national research facility funded by a consortium of Canadian Universities and managed by the National Research Council Canada (http://nmr900.ca). The authors thank Dylan G. Tkachuk for assistance with GIPAW−DFT computations and Yingjie He for performing an SEM measurement.
REFERENCES


(21) Schileo, G.; Grancini, G. Lead or No Lead? Availability, Toxicity, Sustainability and


(60) Cenzual, K.; Gelato, L. M.; Penzo, M.; Parthé, E. Overlooked Trigonal Symmetry in Structures