Halogen Effects on Ordering and Bonding of CH$_3$NH$_3^+$ in CH$_3$NH$_3$PbX$_3$ (X = Cl, Br, I) Hybrid Perovskites: A vibrational spectroscopic study

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ABSTRACT: This study reports Raman and infrared spectra of hybrid organic-inorganic MAPbX₃ perovskites (MA = CH₃NH₃, X = Cl, Br, I) and their mixed-halide derivatives. Raman spectra were recorded at three laser wavelengths (514, 785 and 1064 nm) under on- and off-resonance conditions, as well as at room temperature and 100 K. Use of different excitation wavelengths allowed the unambiguous acquisition of ‘true’ Raman spectra from the perovskites, without degradation or photo-induced structural changes. Low frequency PbX vibrational modes were thoroughly identified by comparison of Raman and far-IR results. Red Raman frequency shifts for almost all MA vibrations from 200-3200 cm⁻¹, particularly intense for the torsional mode, were observed towards heavy halide derivatives, indicative of strengthening the interaction between halides and the organic cation inside the inorganic cage. Different MA-X bonding schemes are evidenced by torsional mode pairs emerging at the orthorhombic phase. MAPbBr₃ was further characterized by variable temperature Raman measurements (100-295 K). Broadening of the MA rocking mode slightly above the tetragonal I to II phase transition is connected with disorder of the MA cation. Our results help understanding perovksite materials properties (ferroelectric domain formation, anomalous hysteresis) and their use as efficient light absorbers in solar cells.
1. INTRODUCTION

Recently, there has been an increase of research into perovskite materials \(^1\)\(^-\)\(^6\) leading to a steep rise in cell performances to a certified value of 21 %\(^7\) within a few years. Similar developments took decades for other photovoltaic technologies to achieve. Even though perovskites are a large family of compounds which all share the common stoichiometry ABX\(_3\), most of the recent attention has been given to the hybrid organic-inorganic methyl ammonium lead halide perovskites (MAPbX\(_3\), MA = methyl ammonium, Pb = lead, X = halide) and in particular the iodide derivative MAPbI\(_3\). Structure and bandgap in ABX\(_3\) perovskites can be easily modified by changing the cation, the metal or the halide. Partial substitution of the iodide in MAPbI\(_3\) results in the iso-structural MAPbI\(_{3-x}\)Br\(_x\) with a blue-shifted bandgap, which is favorable for multi-junction devices and increases chemical stability\(^8\)\(^-\)\(^11\). However, the photo-instability of the excited state of MAPbI\(_{3-x}\)Br\(_x\) results in segregation of the material into micro-domains for \(x > 0.6\)\(^12\). Addition of chloride during perovskite crystallization has been shown to improve charge transport\(^13\),\(^14\) even though its exact role remains unclear\(^15\),\(^16\). Besides the influences of chemical composition on optoelectronic performance and stability, temperature-dependent structural changes play a pivotal role as well. The low-temperature orthorhombic phase of MAPbI\(_3\) is intrinsically more stable compared to the tetragonal phase obtained at room temperature\(^10\) and charge-carrier dynamics become more confined and show excitonic behavior in the orthorhombic phase\(^17\). This indicates that temperature-induced structural changes can tip the scales from an exothermic towards endothermic dissociation enthalpy and from free charge-carrier movement to excitonic confinement. Thus tuning of the perovskite composition or structure can be used in order to enhance stability and performance of perovskite solar cells, both of which are investigated in this study.
X-ray diffraction has been widely used to determine the crystal structure of MAPbX$_3$ perovskite materials after their discovery in 1978$^{18,19}$. However, it can be difficult to resolve the positions of lighter atoms like nitrogen and carbon in the presence of the heavier lead and iodine. The structural orientation of the MA cation within the inorganic framework has therefore been studied in more detail with NMR spectroscopy$^{20}$ and neutron scattering$^{21,22}$. These studies were able to resolve the orientation of MA cations and show an increasing disorder when moving from the orthorhombic phase towards higher temperature phases alongside a weakening of the bonding between the NH$_3$ in the MA and the halides$^{21,22}$. Alignment of the MA cations can yield a net dipole that is energetically favorable in the tetragonal phase and is further stabilized in the orthorhombic perovskite phase. It has been suggested that this MA alignment can induce polar-ferroelectric characteristics in the material in low symmetry phases$^{23}$. Structural dynamics within the perovskite structure have important implications on the operation mechanism of perovskite solar cells. They are thought to assist charge separation$^{24}$, tune the band gap$^{25,26}$, improve open-circuit voltage$^{27}$ and affect hysteresis during current-voltage measurements$^{28}$. These versatile implications show the importance of a detailed study of MA dynamics with Raman and IR spectroscopies.

Perovskite halides have been examined before with mid-IR absorption experiments$^{29-32}$ which give valuable information about MA dynamics, hydrogen bonding and water infiltration inside the perovskites, but do not cover the important low-frequency range where vibrations of the inorganic lead halide network of corner sharing PbX$_6$ octahedra are mainly active. Despite the suitability of the infrared absorption technique for obtaining the vibrational modes in light sensitive systems, only limited far-IR data for single halide perovskite compounds have been reported up to now$^{33,34}$. Besides, reports of Raman measurements on MAPbI$_3$ perovskites$^{23,35-40}$
have so far been limited to resonant or near resonant conditions with typical excitation wavelengths of 514 nm, 532 nm or 785 nm. However, this may be problematic because of the high absorbance and photo-sensitivity of the material. Moreover, the Raman spectrum of MAPbI₃ at low frequencies resembles the PbI₂ precursor spectrum, which makes it difficult to distinguish between pristine and (partially) decomposed samples, as we pointed out recently. Ledinsky et al. reported on photo-induced decomposition during resonant Raman measurements and emphasized the use of laser intensities far below the values that would usually be employed. Apart from the photo-induced decomposition, MAPbI₃ photo-excitation induces a range of other effects that can alter its structure and therefore the experimental outcome.

As mentioned earlier, mixed-halide perovskites have been shown to segregate into micro-domains under illumination, presumably caused by an instability of the photo-excited state. A recent study by Gottesman et al. showed reversible structural modifications of MAPbI₃ in the Raman spectrum by excitation under resonant conditions. Experimental results are often backed by computational studies, which give valuable information. Nevertheless, these calculations are based on several assumptions and produce complex spectra with many superimposed lines especially in the low-frequency range. Also, small uncertainties involved in calculations can result in very large offsets in the resulting spectrum, making peak assignment ambiguous.

The current work investigates MAPbX₃ structural dynamics for all halide derivatives by combining far-IR absorption and Raman scattering and addresses most of the critical issues raised by the literature up to now. For the first time we perform off-resonance Raman experiments and manage to obtain unambiguous Raman spectra avoiding decomposition or structural changes in the material. Measurements have also been carried out on the low-
frequency Raman and IR active modes of the inorganic PbX\textsubscript{6} octahedra. Variable temperature comparative Raman measurements show characteristic spectroscopic changes of the MA torsion and rocking vibrations which have implications on the ordering and bonding of MA inside the inorganic framework. Understanding the fundamental material properties is the key to the construction of perovskite solar cells with improved efficiency and stability.

2. EXPERIMENTAL METHODS

2.1 MATERIALS AND DEPOSITION OF FILMS

The methylammonium halide salts (MAI, MABr and MACl) were synthesized by drop-wise neutralization of an ice-bath cooled solution of 24 mL methylamine (Sigma-Aldrich) in 100 mL of ethanol with the corresponding hydrogen-halide (10 mL of 57 wt% HI in water; 14.6 mL of 48 wt% HBr in water; 15 mL of 37 wt% HCl in water; all Sigma-Aldrich). The solvent was removed in a rotary evaporator and the powder was recrystallized from ethanol. Upon slow cooling of the ethanol solution white crystals were formed. Washing with diethylether and drying at 70 °C for 6 h was performed afterwards. The single halide perovskite (MAPbI\textsubscript{3}, MAPbBr\textsubscript{3} and MAPbCl\textsubscript{3}) precursor solutions were prepared by mixing a 1:1 stoichiometric ratio of lead(II) halide (PbI\textsubscript{2}, 99 %; PbBr\textsubscript{2}, 99.99 %; PbCl\textsubscript{2}, 99.99 %; all Sigma-Aldrich) with the corresponding methylammonium halide at a 1 mol/L concentration in DMSO (≥ 99.9 %; Sigma-Aldrich ) for 1 h at 60 °C. For the mixed halide perovskite precursor solutions, the respective ratios of single halide solutions were mixed and stirred for 1 h. Synthesis of the precursor salts was performed in the fume hood, while the mixing of the perovskite precursor solutions, films deposition as well as annealing were done in an Argon-filled glovebox (H\textsubscript{2}O and O\textsubscript{2} concentration < 1 ppm).
For planar films the precursor solution was deposited on a microscope glass slide. For mesoporous films, a dispersion of Al₂O₃ nanoparticles (20 wt% in 2-propanol; Sigma-Aldrich) was spin-coated onto glass slides (2000 rpm for 60 s) and annealed at 500 °C for 30 min. All perovskite solutions were spin-coated onto the flat or mesostructured substrates at 2000 rpm for 60 s and then annealed on a hotplate at 100 °C for 30 min. Samples for far-IR characterization were prepared by depositing mesoporous perovskite films on a 0.8 mm thick high density polyethylene (HDPE) to improve surface wettability. In order to enhance the perovskite Raman scattering volume, measurements were performed on powder samples taken by scratching the films.

2.2 CHARACTERIZATION

The perovskite films were structurally characterized using a Siemens D-500 diffractometer, that operates in Bragg-Brentano geometry with Cu Kα1 radiation (λ = 1.5406 Å) and Cu Kα2 radiation (λ = 1.5444 Å). Data were collected over the angular range 5 ° ≤ 2θ ≤ 100 ° counting for 1 s at each step of 0.05 ° in detector position. Unit-cell refinements were performed with GSAS package⁴⁴.

UV-Vis Diffuse Reflectance (R) spectra were acquired on a Hitachi 3010 spectrophotometer using an integrating sphere with 60 mm diameter and BaSO₄ as a reference. The data were transformed into Kubelka-Munk absorbance F(R) = (1−R)²/2R.

The IR transmittance measurements were run at normal incidence on a Bruker Vertex 80v vacuum spectrometer using HDPE-Alumina films as a reference. Each spectrum was averaged over 200 scans with a 4 cm⁻¹ resolution. The far-infrared (FIR) beamsplitter and detector used
offered a working range of ca. 60 cm\(^{-1}\) to 680 cm\(^{-1}\). The transmittance spectra were converted to absorbance and normalized to 1.

Raman measurements with excitation at 514 nm and 785 nm were performed with a Renishaw In-Via Reflex dispersive micro-Raman spectrometer with <2 cm\(^{-1}\) resolution. The laser beam was focused on a spot of 1 µm in diameter with a x100 magnification lens. Rayleigh scattering was rejected with 50 cm\(^{-1}\) or 110 cm\(^{-1}\) cut-off dielectric edge filters, and analysis of the scattered beam was performed on a 250 mm focal length spectrometer along with suitable diffraction gratings (1800 lines/mm for visible and 1200 lines/mm for NIR) and a high-sensitivity CCD detector. Cryostatic Raman measurements at low temperature (LT) were done under inert atmosphere in a THMS600PS Linkam cell by focusing the laser beam with a x50 long focal distance lens. Raman spectra under off-resonance conditions for all samples were acquired with a Bruker RFS100 FT-Raman spectrometer equipped with a Nd-YAG laser emitting at 1064 nm. Each spectrum was averaged over 100 scans or 200 scans with 4 cm\(^{-1}\) resolution, while the effective working range was ca. 100 cm\(^{-1}\) to 3500 cm\(^{-1}\). All RT Raman measurements were performed in air and at different laser powers to assure that the intensities used did not induce changes in the spectra. The measured Raman spectra present a strong elastic scattering background in the low-frequency (LF) spectral range. In order to enhance the Raman features at this LF range where very informative PbX and MA vibrations are active, and correctly determine spectral characteristics (frequency, linewidth and line shape), the spectra were temperature-reduced using the expression of eq. (1):\(^{45}\):

\[
I_{\text{red}}(\omega) = I(\omega) \frac{1}{n(\omega,T)+1}, \quad n(\omega,T) = \left(\exp^{\frac{\hbar \omega}{kBT}} - 1\right)^{-1} = \left(\exp^{\frac{1.44 \cdot \omega}{T}} - 1\right)^{-1}, \quad (1)
\]
where $I(\omega)$ and $I_{\text{red}}(\omega)$ are the measured and reduced Raman intensity, respectively. $n(\omega, T)$ is the Bose-Einstein statistic factor, $h$ and $k_B$ are the Planck and Boltzmann constants, respectively, $c$ is the speed of light, $\omega$ is the Raman shift in cm$^{-1}$ and $T$ is the temperature in K. The analysis of the reduced spectra was done by fitting them with mixed Lorenzian-Gaussian functions using the Wire software from Renishaw. Linewidths were corrected for instrumental broadening.

3. RESULTS AND DISCUSSION

3.1 STRUCTURAL AND OPTICAL CHARACTERIZATION

Schemes of the perovskites structure at the three different structural modifications (cubic, tetragonal and orthorhombic) together with transition temperatures are shown in Figure 1a. Powder-XRD patterns of the area of interest for the whole set of MAPbX$_{3-x}$Y$_x$ materials, shown in Figure 1b, confirm the presence of the perovskites as the main phase. In addition, certain amounts of admixtures were present, in particular PbI$_2$ for the I-containing samples (for the full patterns see Figure S1 in the Supporting Information). MAPbCl$_3$, MAPbBrCl$_2$, MAPbBr$_2$Cl and MAPbBr$_3$ were indexed as cubic systems, whereas MAPbBrI$_2$, MAPbBrI$_2$ and MAPbI$_3$ were indexed as tetragonal systems, in accordance to ref. [8]. A regular trend in the unit-cell expansion with substitution of Cl by Br and I was observed. The determined lattice parameters of the tetragonal MAPbI$_3$ ($a = b = 8.861(2)$ Å, $c = 12.653(4)$ Å) are in agreement with the literature values of $a = b = 8.849(2)$ Å, $c = 12.642(2)$ Å$^{47}$, within the 3σ estimated standard deviation. The lattice parameters are given in Table S1 of Supporting Information.
Figure 1: (a) The different phases of the MAPbX₃ perovskite, which transform upon changing temperature or halide composition; halides: large purple spheres, Pb: large grey spheres and MA: interstitial. Transition temperatures into tetragonal II MAPbBr₃ are in brackets. (b) Powder XRD patterns of the range of interest from all halide perovskites examined in this study. (c) UV-Vis spectra (where F(R) is the Kubelka-Munk function) of perovskite films which change in color (from dark brown to white) and show a shift in band gap energy (from 1.61 to 3.06 eV) as lighter halides are introduced into the structure.

Substitution of heavy halogen atoms with lighter ones in MAPbX₃₋ₓYₓ results in films with tunable color from dark brown to white, as shown by the photographs of the prepared films inserted in Figure 1c. In order to analyze the light harvesting of the films over the solar spectrum diffuse reflectance measurements were carried out. In Figure 1c, the absorption F(R) data are plotted in terms of \((F(R)h\nu)^2\) vs. \(h\nu\), which is the standard analysis for materials having a direct band gap. Band gap values \(E_g\) were found by the intersections of the slope with the energy-axis and are shown upon the graphs, and further listed in Table S2 where they are compared with the Raman excitation energies. The values are slightly blue-shifted compared to literature values. Blue-shifted absorption onsets were reported recently. The shift can be caused by a ferroelectric domain wall induced band gap reduction and has been observed for mesoporous...
scaffold architectures. The influence of annealing temperature on band gap is another possible explanation for this observation. In general, the reported band gaps for MA lead perovskites can vary significantly. The excitonic peaks appearing in the spectra for lighter halide derivatives are presumably caused by an increase in the exciton binding energies.

3.2 LOW FREQUENCY Pb-X VIBRATIONAL MODES: FAR-IR AND MICRO-RAMAN RESULTS AT ROOM TEMPERATURE

The PbX	extsubscript{6} octahedra of the perovskite halides belong to the O	extsubscript{h} point group symmetry and have two IR active modes, ν	extsubscript{3} and ν	extsubscript{4}, with T	extsubscript{1u} symmetry and three Raman active modes ν	extsubscript{1}, ν	extsubscript{2} and ν	extsubscript{5}, with A	extsubscript{1g}, E	extsubscript{g} and T	extsubscript{2g} symmetry, respectively. Splitting of the doubly and triply degenerate modes is expected due to distortion of the octahedra, a) by interaction between the methylammonium and certain halogen atoms, and b) by the occupation of the octahedral corners with different halide atoms in the complex halide derivatives. According to theoretical calculations, the splitting is considerably larger for the Cl based perovskites (Cl has the strongest electronegativity). In the present study, and for the first time in the literature, both the IR and Raman low-frequency Pb-X vibrational modes have been observed for all members of the MAPbX	extsubscript{3} family.

Far-IR spectra obtained for all MAPbX	extsubscript{3} derivatives are shown in the range of 50-300 cm	extsuperscript{-1} in Figure 2a. Broad spectral lines are observed in all cases, justified by the polycrystalline character of the materials. Analysis of spectral components as well as spectrum of the HDPE-Alumina reference are shown in Figure S2. The spectra for MAPbI	extsubscript{3} present strong IR-activity below 120 cm	extsuperscript{-1}, which is mainly due to internal PbI	extsubscript{3} vibrations as shown in recent density functional perturbation theory calculations. The spectra of the MAPbBr	extsubscript{x}I	extsubscript{3-x}Br	extsubscript{x} series are analyzed with...
two component bands, which are straightforwardly attributed to the $\nu_3$ asymmetric stretching (103-125 cm$^{-1}$) and $\nu_4$ asymmetric bending (66-76 cm$^{-1}$) vibrations of PbX$_6$ octahedra. For the Cl containing samples, the higher frequency band is split into two component-bands, which are well described by mixed Lorenzian-Gaussian character. For the pure Cl perovskite, a very broad band is observed which includes both the $\nu_4$ band, as a low frequency shoulder, as well as the split $\nu_3$ band at higher frequencies. The peak positions of the deconvoluted $\nu_3$ and $\nu_4$ modes (two peaks for the Cl containing perovskites) are marked upon Figure 2a. In Figure 2b we plot the frequency of the two modes versus the halogen order. For the estimation of the $\nu_3$ frequency for the mixed halides, a weight average of the two $\nu_3$ related component bands was taken into account. First of all we should note that the ratio $\nu_4/\nu_3$ remains stable in the 0.60-0.64 range. The only exception is the pure Cl perovskite for which it is slightly larger and equal to 0.67, presumably due to uncertainties in the fitting of the corresponding broad far-IR band. The fixed $\nu_4/\nu_3$ ratio verifies self-consistency of the analysis, and the ratio values match the corresponding ratios for several other compounds quite well (e.g., the ratio for the [PbCl$_6$]$^{2-}$ anion from ref. [Error! Bookmark not defined.] is 0.54).
Figure 2 (a) Far-IR spectra for the various perovskite derivatives measured at RT. The band marked by PE is due to the polyethylene substrate. Peak frequencies of the \( \nu_3 \) and \( \nu_4 \) bands obtained by fitting the spectra (see Figure S2) are marked with dashed and solid lines, respectively. The values are noted for the end perovskite members, MAPbI\(_3\) and MAPbCl\(_3\). For Cl based perovskites the \( \nu_3 \) band is split into two component bands, \( \nu_{3a} \) and \( \nu_{3b} \). (b) Frequency of \( \nu_3 \) (average) and \( \nu_4 \) modes vs. halide composition. Dashed lines show predictions based on the harmonic oscillator model.

Both \( \nu_3 \) and \( \nu_4 \) modes shift to higher frequencies upon introduction of lighter halides in the structure. The \( \nu_3 \) and \( \nu_4 \) frequencies depend strongly on the reduced mass \( \mu \) of the Pb-X bond and follow the model for a harmonic oscillator \( \nu = (\frac{1}{2\pi})\sqrt{k/\mu} \), with the crude assumption of the same bond strength \( k \) for all halides. The reduced mass is \( \mu^{-1} = m_{Pb}^{-1} + m_{X}^{-1} \) where \( m_{X} \) for the
composite perovskite halides was calculated by the weight average of the halogen atomic masses in their chemical formula. The model calculations have been done by weighting the \( \nu_3 \) and \( \nu_4 \) frequencies against those observed for MAPbBr\(_3\) and are shown with dashed lines in Figure 2b. These results verify that the low-frequency IR active vibrations are of the inorganic Pb-X type and lack noticeable influence from the MA cation. A similar behavior which consolidates our argument is observed by analyzing the frequency shifts of the Sn-X modes in the analogous \([SnX_6]\)^{2-} octahedral system, irrespective of the vibration mode\(^{52}\).

Complementary to the far-IR spectra, low frequency Raman spectra were obtained above 50 cm\(^{-1}\) with 785 nm laser line excitation. The spectra are presented in Figure 3a for all the perovskite compounds apart from the pure MAPbI\(_3\) which showed only a strong photoluminescence (PL) response. The vibrational modes of the PbX\(_6\) octahedra were observed below 250 cm\(^{-1}\). At 270 cm\(^{-1}\) for MAPbBrI\(_2\) and at higher frequencies for the rest of the compounds, a broad and well-formed band was observed. This band is due to the torsion of MA units and will be discussed in detail in the next section.

The Raman spectra below 230 cm\(^{-1}\) involve vibrations of Pb-X bonds consisting of heteropolar ionic/covalent interactions in the inorganic framework and show very broad multi-bands which are not easily resolved into components. The general trend is that all these measured bands increase in frequency by moving towards light halides following the corresponding behavior shown in the far-IR spectra. The multi-band character of the Raman features is clearer for the mixed perovskite derivatives and is justified considering the lifting in degeneracy of modes. As already discussed, three Raman bands due to the Pb-X vibrations of the octahedral structure are expected in the LF range. Among them the symmetric stretching mode \( \nu_1 \) is expected to be the highest in frequency, followed with the asymmetric stretching mode \( \nu_2 \) and the asymmetric
deformation $\nu_5^{52}$. Since it is difficult to fit the spectra, we attempt an analysis based on the ratios of the frequencies between the various inorganic vibrational modes from the literature. Thus according to data in Ref [Error! Bookmark not defined.], the ratios $\nu_1/\nu_3 = 1.07$, $\nu_2/\nu_1 = 0.74$ and $\nu_5/\nu_1 = 0.49$ are observed for $[\text{PbCl}_6]^{2-}$ while similar values are found for other systems, too, such as for $[\text{SnX}_6]^{2-}$ and $[\text{SbX}_6]^{2-}$. Thus, based on our $\nu_3$ values from our far-IR analysis we have predicted the mean frequencies of the corresponding Raman bands. The respective calculated values of the three Raman modes are reported in Figure 3a and qualitatively describe the shifts observed as a function of the halide change. By extending the calculations for MAPbI$_3$, the mean frequencies for the three Raman active modes are predicted to be $\nu_1 = 110$ cm$^{-1}$, $\nu_2 = 81$ cm$^{-1}$ and $\nu_5 = 54$ cm$^{-1}$ (see also Figure 3a).

![Figure 3](image.png)

**Figure 3:** (a) Low frequency Raman spectra obtained by excitation at 785 nm. Peak frequencies of $\nu_1$ (solid line marks), $\nu_2$ (dashed line marks) and $\nu_5$ (dotted line marks) noted upon the graph are estimated from the experimentally recorded frequency of the $\nu_3$ IR band and modes
frequency ratios from the literature (for details see text). Raman spectra of the perovskite halide derivatives obtained by excitation at (b) 1064 nm, (c) 785 nm and (d) 514 nm, at RT.

3.3 VARIABLE EXCITATION MICRO-RAMAN SPECTRA OF ALL PEROVSKITE DERIVATIVES

Extensive Raman investigation with emphasis on MA vibrations was done by performing measurements in a broad frequency range (100 cm\(^{-1}\) to 3500 cm\(^{-1}\)) using three different excitation lines at 514, 785 and 1064 nm. This allowed data acquisition both at resonance (Res) as well as off-resonance conditions (Off-res). An overview of materials and excitation wavelengths is shown in the ESI (Table S2). In general, the majority of resonance Raman measurements obtained from the MAPbX\(_3\) perovskites carries limited information on the vibrational properties of the materials. The main reason is that the strong absorption of the perovskites, when excited within the bandgap, can induce damage or phase transformation in the materials. This is an issue especially for MAPbI\(_3\) with a phase transition from tetragonal to pseudo-cubic near room-
temperature, at 330 K. Despite keeping the laser power density very low, by working under resonance or near resonance conditions the Raman signals are frequently masked by the presence of a strong PL background signal. Furthermore, temporary PL signals which change with illumination time were observed here in Br-I mixed halide perovskites and are attributed to phase segregation, as previously reported by Hoke et al. 

Figures 3b-d show an overview of the Raman spectra for all the perovskite films obtained under different excitation wavelengths, 1064 nm (Figure 3b), 785 nm (Figure 3c) and 514 nm (Figure 3d). In the case where the Raman features are masked by strong PL signals, the corresponding spectra are not presented. Thus, for MAPbI$_3$, we include only the spectrum obtained at 1064 nm excitation in Figure 3b, and no spectra in Figure 3c and Figure 3d, because, under near bandgap or resonance excitation, the Raman signal is either covered by a strong PL background or it is affected by the quick degradation of the sample showing spectroscopic changes during the measurement. MAPbCl$_3$ on the other hand shows no apparent change in its Raman spectrum, for all the excitation sources used. Hence, excitation at 1064 nm (Figure 3b) ensures off-resonance conditions in all cases allowing a clear and unambiguous overview of the full MA spectrum for all halide perovskite derivatives and no spectral changes were observed that could be related to humidity or light exposure of the material. Furthermore, the quality of the spectra is excellent mainly due to the fast acquisition in the FT technique and the materials tolerating high laser power densities at 1064 nm excitation, which compensate well for the inherently weaker scattering efficiency vs. the increase of the excitation wavelength (inverse fourth power law applies).
The Raman spectra reported in Figures 3b-d show a large number of MA bands which present significant changes in their spectral characteristics (shifts, widths, intensities) with halide composition. An overview of the frequencies of the measured Raman bands and their mode assignments can be found in Table 1. Almost all MA vibrational modes present a blue shift from MAPbI₃ towards MAPbCl₃ (see Figure S4) in accordance with the IR data in Ref [Error! Bookmark not defined.] which indicates strong NH₃–X interaction. In particular, the mode that is most sensitive to the halide content is the high frequency $\nu_{15}$-$\nu_{6}$ (NH₃⁺) mode which positively shifts abruptly from MAPbI₃ to MAPbI₂Br by 33 cm⁻¹ and then up to MAPbCl₃ by another 14 cm⁻¹. Significant positive shifts also occur in all other modes related to NH₃ vibrations. This is in line with recent NMR results⁵³, which have shown that only the amine end of the MA group interacts strongly with the inorganic network. Nevertheless, the exact N-H bond lengths in MAPbX₃ cannot be accurately determined even with neutron diffraction methods due to the disorder in the MA group⁵³, thus the full understanding of the MA vibrational frequency shifts (e.g. the large $\nu_{15}$ shift for MAPbI₃ compared to the other perovskites) and the exact nature of the MA-halide interaction is not possible.

Table 1: Frequencies (in cm⁻¹) and assignments of vibrations numbered in series for MA lead derivatives MAPb[X], where [X] is the halogen composition. Modes from $\nu_6$ and above were obtained from Raman spectra with 1064 nm excitation and are due to MA vibrations. Modes $\nu_3$ and $\nu_4$ were obtained from far-IR spectra. $\nu_1$, $\nu_2$, and $\nu_5$ records are estimated frequencies and match well with experimental Raman spectra at low frequencies. Corresponding values for MAPbI₃ are shown in parenthesis to indicate that Raman experimental data are not available in this work.

<table>
<thead>
<tr>
<th>Mode/[X]</th>
<th>I₃</th>
<th>I₂Br</th>
<th>IBr₂</th>
<th>Br₃</th>
<th>Br₂Cl</th>
<th>BrCl₂</th>
<th>Cl₃</th>
<th>Assignment</th>
</tr>
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</table>

Characteristic internal vibrational modes of MA appear at high frequencies, just below 1000 cm\(^{-1}\) (C-N stretching), at 1400 cm\(^{-1}\) to 1600 cm\(^{-1}\) (CH\(_3\) and NH\(_3\) bending) and at around 3000 cm\(^{-1}\) (CH\(_3\) and NH\(_3\) stretching) and are well studied for methylammonium halides\(^{30,55}\). The most interesting vibrational modes are the rocking vibration, \(v_7\), and low-frequency torsion, \(v_6\), of MA around the C-N axis. The MA rocking mode, \(v_7\), appears in between 911 cm\(^{-1}\) to 923 cm\(^{-1}\) and sharpens, increases in intensity and red-shifts when moving towards heavier halides.

<table>
<thead>
<tr>
<th>(v_1)</th>
<th>(110)</th>
<th>114</th>
<th>124</th>
<th>134</th>
<th>142</th>
<th>163</th>
<th>183</th>
<th>(v_4) (Pb-X) this work</th>
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<tr>
<td>(v_2)</td>
<td>(81)</td>
<td>85</td>
<td>92</td>
<td>99</td>
<td>105</td>
<td>121</td>
<td>135</td>
<td>(v_s) (X-Pb-X) this work</td>
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<tr>
<td>(v_3)</td>
<td>103</td>
<td>107</td>
<td>116</td>
<td>125</td>
<td>133</td>
<td>152</td>
<td>171</td>
<td>(v_{as}) (Pb-X) this work</td>
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<tr>
<td>(v_4)</td>
<td>65</td>
<td>69</td>
<td>72</td>
<td>76</td>
<td>82</td>
<td>91</td>
<td>114</td>
<td>(\delta_{as}) (X-Pb-X) this work</td>
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<td>(v_5)</td>
<td>(54)</td>
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<td>61</td>
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<td>90</td>
<td>(\delta_s) (X-Pb-X) this work</td>
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\(v\): stretching; \(\delta\): bending; \(\rho\): rocking; \(\tau\): torsion; \(s\):symmetric; \(as\):asymmetric
Cl→Br→I (Figure 3b-d, Figure S4); the same trend was found for other modes (e.g., the v_{12}, δ_{s}(NH_{3}^{+}) mode). This behavior can be attributed to orientational ordering of MA for which v_{7}, having the E symmetry^{56}, is a good indicator. The effect can be analyzed by the calculation of the correlation times (τ_c) from the peak full width at half maximum (FWHM)^{29}, which are shown in Figure 4a, according to eq. 2:

\[ \tau_c = \frac{1}{2 \pi c \text{ FWHM}} \quad (2) \]

where c is the velocity of light.

**Figure 4:** (a) FWHM of the v_{7} rocking mode averaged over measurements obtained at RT with different laser lines (solid squares – standard deviation < 5 cm\(^{-1}\)) as well as at LT (100 K) obtained at 785 nm excitation (open circles) (b) Raman frequency of the torsional mode averaged over measurements obtained at RT with different laser lines (solid squares standard
deviation < 3 cm\(^{-1}\)) and corresponding results at LT (100 K) obtained at 785 nm (open circles). Solid line guides the eye for halogen dependence of the RT data and is extrapolated either sides with dashed lines.

The results show correlation times below 0.15 ps for the \(v_7\) rocking vibration for all perovskite derivatives. \(\tau_c\) gradually decreases from 0.15 ps to 0.1 ps from MAPbI\(_3\) up to MAPbBrCl\(_2\) and drops to 0.05 ps for MAPbCl\(_3\). Measurements of MA rocking vibration dynamics can be compared with early IR absorption data\(^{29}\) obtained from large perovskite crystals, where the trend of correlation time decrease from heavy to light halides is also followed, as well as with recent results on very high quality thin films prepared with the dual source evaporation method.

In both these literature works, the \(\tau_c\) values for MAPbI\(_3\) were substantially larger. Thus, the shorter correlation time in our samples is attributed to their polycrystalline character.

The \(v_6\) mode mainly consists of an NH\(_3\) motion and has an \(A_2\) symmetry which makes it forbidden in IR but permitted in Raman\(^{55}\). This mode is easily accessed in all Raman experiments and is analyzed in a straightforward way as it is not overlapping with other modes. For MAPbCl\(_3\)\(^{54}\) the mode is observed at 488 cm\(^{-1}\) and presents an extremely large red frequency shift upon halide substitution, moving down to 249 cm\(^{-1}\) for MAPbI\(_3\) (see Figure 4b and Table 1). This large spread of resonance frequencies for MA is unusual for a torsional mode and indicates intense MA deformation and a strong interaction of MA with the inorganic cage. Fitting of \(v_6\) for different halide derivatives gives a continuous shift from MAPbI\(_3\) to MAPbBr\(_2\)Cl, but jumps to higher wavenumbers for perovskites with large amounts of Cl, as shown in Figure 4b. According to recent theoretical studies\(^{23}\), different mechanisms like molecular deformation and dipolar orientational ordering of the MA as well as strength of the NH\(_3\)-X interaction are strongly
affected by the nature of the halide in the perovskite, and result in large shifts in the frequency of the MA torsion. Among them, changes in NH$_3$-X interaction upon halide substitution is the most prominent reason for the large frequency shift of the $\nu_6$ mode, considering that it affects the frequencies of all other MA modes, too, in the same manner, even though to a much lesser extent. Preliminary calculations performed with the same methodology reported by Brivio at al.$^{38}$ confirm the observed increase of the torsional mode frequency for perovskites with Br and Cl with respect to iodine. This is due to a halide specific NH$_3$–X interaction, and to a confinement effect. In fact, lighter halide results in a smaller cavity inside the inorganic cage that uplifts the frequencies of the molecular modes, and reduces the distance between hydrogen and halide, amplifying the resulting shift. In Figure S5 we attempt a correlation of the torsional mode peak frequency at room-temperature (RT) against the measured band gap for the various perovskite halides. Halide substitution results in vast changes of the materials optical properties, which can significantly affect their vibrational frequencies as also proposed before$^{30}$ by correlating the red shifts of MA vibrations in the Cl→Br→I order to the increase in the polarizability of the dielectric medium (Lorentz-Lorenz shift).

Interestingly, the signal intensity of $\nu_6$ is much stronger for single halide perovskites (see Figure 3b-d) and its linewidth is not very broad, ranging from 40 to 60 cm$^{-1}$. In the mixed halide perovskites a big drop in the $\nu_6$ intensity takes place which goes along with extensive linewidth broadening, reaching values up to 150 cm$^{-1}$. A general peak broadening can be caused in several ways. A decrease in long-range order and crystallinity typically yields broader peaks. Unresolved mode-splitting also causes an increase of peak width and is connected to possible asymmetries of the mode. Such mode splitting can be enhanced by micro-structural inhomogenities due to an uneven distribution of a halide in the perovskite. This effect has been reported before for
MAPbI$_{3-x}$Br$_x$, which forms micro-domains upon crystallization that transform into a single-phase within a few weeks$^{57}$ but also segregates again upon illumination$^{12}$.

The informative torsional and rocking vibrations were further examined by comparative low-temperature Raman measurements in order to elucidate their behavior by lowering the symmetry of the perovskite structure.

3.4 LOW TEMPERATURE RAMAN MEASUREMENTS

Another series of Raman experiments was performed for all perovskites at 100 K which is well below the phase transition of all materials into the orthorhombic phase$^{46}$. Characteristic reduced spectra at low temperature are shown in Figure 5a obtained with excitation at 785 nm. The inset shows off-resonance spectra of MAPbBrCl$_2$ and MAPbCl$_3$ using 514 nm excitation. Peak frequencies of the strongest MA vibrations are given in Table S3. Mode splitting of the low-frequency modes was observed due to the lowering of the crystal symmetry. All materials transform from tetragonal to orthorhombic in the temperature range of 148.8-171.5 K$^{58}$. Splitting was more evident for lighter halide derivatives, in accordance to early results for MAPbCl$_3$$^{54}$. In the case of heavier derivatives, mode splitting was only partially observed due to the shifting of the majority of the modes below the experimentally accessed frequency range. Furthermore, apart from narrowing and blue shifting of the MA high frequency bands above 950 cm$^{-1}$, no clear MA mode splitting was evidenced upon decrease of the temperature. A similar effect has been reported before for mid-IR data below the phase transition temperature$^{29}$ as well as for Raman in other perovskite systems$^{59}$ implying that the crystal deformation does not drastically affect the high frequency vibrations of the MA units.
Figure 5: (a) Raman measurements of MAPbX$_{3-x}$Y$_x$ perovskites at 100 K obtained with 785 nm excitation. Intensity of the top three spectra below 200 cm$^{-1}$ is scaled down by 3 for better vision of the spectra. Inset shows spectra obtained with 514 nm excitation. (b) Raman spectra of MAPbBr$_3$ (785 nm) vs. temperature variation from 100 K (LT) to 295 K (RT).
It is remarkable that at LT the rocking vibrations \( \nu_7 \) and \( \nu_9 \) at 920 and 1250 cm\(^{-1}\) present considerable linewidth narrowing. Thus, the \( \nu_7 \) rocking mode width decreases to about 20 cm\(^{-1}\) for all perovskite derivatives (see Figure 4a) and the \( \nu_7 \) correlation time increases from below 0.15 ps at RT to above 0.25 ps at 100 K, which implies MA dipolar ordering\(^{29}\), in accordance with quasi-elastic neutron scattering results\(^{22}\).

Further insight into the behavior of the \( \nu_6 \) torsion mode can be extracted from the low-temperature (LT) data. While the measurements at RT showed a jump to higher wavenumbers for the MAPbBrCl\(_2\) and MAPbCl\(_3\) perovskites, at LT pairs of peaks appeared (see spectra in Figure 5a and analysis in Figure 4b) that line up with modes on either side of the gap, suggesting that those peaks in fact represent two distinctive vibrational bands. The appearance of two torsional bands at low-temperature is in line with early studies for MAPCl\(_3\)^{54}. Doubling of the torsional modes for the low symmetry orthorhombic phase and linewidth broadening might be due to different \( \text{NH}_3\)-X bonding schemes of the organic molecule within the hybrid\(^{60}\). On the other hand, the frequencies of the torsional modes at low temperature show small, mostly blue shifts relative to those at RT.

Another important aspect of the torsional mode is its significant broadening in the orthorhombic phase. The unit cell of this phase contains four different MA units and according to our preliminary calculations, the frequencies of respective modes are split in four due to slightly different local environments. This split is in the order of few cm\(^{-1}\) and therefore too small to be resolved but contributes in significant linewidth broadening. Moreover, as proved and observed in molecules\(^{61,62}\) or complex structures\(^{63}\), Van-der-Waals forces that are enhanced at low temperature, affect the stiffness of the bonds and influence the non-harmonic terms of the vibration that lead to a larger broadening of the peak. This effect strongly depends on the
position and the geometry of the system and could justify the different behavior below the transition temperature. In previous work\textsuperscript{64} it has been reported an anomalous temperature behavior of a low frequency Raman peak in NH\textsubscript{4}Br around phase transition. This anomaly results in asymmetric peak shape and anomalous broadening of the signal. These effects are addressed and explained with a long range order of the NH\textsubscript{4} cations. We can imagine that a similar mechanism, regarding the MA ordering, could influence the linewidths at LT of the torsional mode in the perovskite halides too.

In addition to fixed LT measurements at 100 K, variable temperature Raman measurements from 295 K to 100 K have been carried out for MAPbBr\textsubscript{3}. During this temperature ramp the material was transformed progressively from cubic to the tetragonal I phase (236.3 K), the tetragonal II phase at 154.0 K and the orthorhombic phase at 148.8 K\textsuperscript{46}. Recorded Raman spectra are shown in Figure 5b. MAPbBr\textsubscript{3} was chosen as the intermediate representative of our halide series and because of its high purity (see XRD in Figure S1), the fact that the full temperature dependence has not been reported before in the literature (MAPbCl\textsubscript{3} has been studied in ref. \textsuperscript{[Error! Bookmark not defined.]} and MAPbI\textsubscript{3} Raman signal is covered by a strong PL background at 785 nm excitation) and its significance as light absorber for the preparation of very stable perovskite solar cells\textsuperscript{8-11,65}.

All high frequency MA modes above 950 cm\textsuperscript{-1} present a ‘normal’ behavior with line narrowing and blue shift upon temperature decrease. Important observations are made by following the temperature dependence of the spectra below 950 cm\textsuperscript{-1} where peculiar behavior of the MA torsional and rocking modes at 320 cm\textsuperscript{-1} and 925 cm\textsuperscript{-1} are observed upon moving from the cubic (295 K) to the orthorhombic phase (100 K).
Figure 6 plots the temperature dependence of the linewidths for the ‘abnormal’ torsional \( \nu_6 \) and rocking \( \nu_7 \) modes in comparison to the CN stretching \( \nu_8 \) mode which shows a ‘normal’ behavior. The slope of linewidth against temperature shows a change for \( \nu_7 \), at about 175 K, slightly above the tetragonal I-II phase transition at 154 K. This transition temperature is close to the temperature of a discontinuous decrease of the real part of the dielectric permittivity\(^{58}\) as well as of significant changes in the H\(^1\)-NMR spectra and relaxation time\(^{66}\). According to this early H\(^1\)-NMR result, the tetragonal I to tetragonal II phase transition affects the alignment of MA with freezing of the C-N axis.

![Temperature dependence of linewidths for the torsional \( \nu_6 \), rocking \( \nu_7 \) and CN stretching \( \nu_8 \) vibrations of MAPbBr\(_3\). Solid line represent fitting of \( \nu_7 \) linewidths by eq. 1 with b=0. Dotted lines mark temperatures of phase transitions.](image)

**Fig. 6** Temperature dependence of linewidths for the torsional \( \nu_6 \), rocking \( \nu_7 \) and CN stretching \( \nu_8 \) vibrations of MAPbBr\(_3\). Solid line represent fitting of \( \nu_7 \) linewidths by eq. 1 with b=0. Dotted lines mark temperatures of phase transitions.

The temperature dependence of the linewidth of the \( \nu_7 \) rocking mode in Figure 6, was used to calculate the activation energy (\( E_a \)) of the order-disorder process, according to equation (3)\(^{67-69}\).

\[
FWHM(T) = a + bT + c \exp(-Ea / RT) \quad (3)
\]
where \( a, b, c \) are fitting parameters and \( R \) is the gas constant. The linear term describes the anharmonicity factor and the exponential term the increase of the linewidth due to the thermal reorientational motions. We fitted the FWHM(T) data of \( \nu_7 \) in Figure 6 with eq. (3), neglecting the linear term which has a small contribution and ended up with an estimation of the activation energy of \( E_a = 7.5 \pm 1.0 \text{ kJ/mol} \) which resembles the energy barrier in the disorder of the MA units. This value is very similar to that found for the rocking vibration of \( \text{NH}_3 \) in \([\text{Mn(NH}_3)_6]^{2-}\text{(ClO}_4)_2\). As shown in Figure 6, the reorientational motions are not significantly disturbed near the phase transition temperature which is indicative that the order-disorder of the organic molecule does not contribute significantly in the phase transition mechanism. Among different vibrational modes, the main torsional mode at about 320 cm\(^{-1}\) is the only one that shows line broadening instead of narrowing upon decreasing the temperature (see Figure 6). This broadening occurs progressively but shows a steep rise below the transition to the orthorhombic phase. This interesting behavior, which was also observed for other perovskite halides as discussed before, can be justified by the emergence of new unresolved torsional bands and long range MA order in the low symmetry phase.

4. CONCLUSIONS

In summary, our study gives a complete overview of the vibrational properties of single and mixed MAPbX\(_3\) (\( X = \text{Cl, Br, I} \)) perovskites. Measurement in off-resonance conditions, at room- and low-temperatures, and a comparison of the Raman with far-IR spectra allow an unambiguous attribution of inorganic and organic modes and their shifts against halide constitution and temperature. Using three different excitation wavelengths for the Raman measurements, a significant influence of beam-induced degradation on the spectra of this material class can be ruled out for the first time in the literature. The low frequency range is dominated by the modes
of the inorganic anion and the frequency shifts versus halide content is governed by the Pb-X reduced mass. Insight into structural dynamics of the MA cation has been given where specific vibrations such as the NH$_3$ rocking mode indicate ordering of MA at low temperatures. Amongst different modes, the MA torsion is particularly interesting since it spans a broad frequency range from 249 cm$^{-1}$ to 488 cm$^{-1}$ by altering the halogen content. This behavior is attributed to changes in the interaction between the NH$_3$ head of MA and the halide. Furthermore, the MA torsional band shows broadening and splitting at 100 K in the orthorhombic phase, which indicates possible different MA-X bonding schemes. We hope that these findings encourage a broader usage of vibrational spectroscopy for inorganic-organic hybrid perovskites, and aid a better understanding of their properties, which rank them among the most efficient solar cell active materials.

ASSOCIATED CONTENT

**Supporting Information Available**: Table S1: Lattice parameters from XRD data for MAPbX$_3$ perovskites; Table S2: Band gaps and laser excitation energies for the Raman and PL investigation of perovskite materials; Table S3: Overview of main MA Raman frequency peaks above 900 cm$^{-1}$, measured at 100 K.

Figure S1: Full powder XRD patterns of all halide perovskites; Figure S2: Fitting analysis of the far-IR absorption spectra; Figure S3: Spectra showing shifting of PL during measuring MAPbI$_2$Br under resonance (514 nm) excitation, indicating phase segregation; Figure S4: Raman frequency shifts of strongest MA modes above 900 cm$^{-1}$ for various halogen contents in MAPbX$_3$; Figure S5: The frequency of the $\nu_6$ torsional mode at RT against the band gaps for all measured MAPbX$_3$ derivatives.
This information is available free of charge via the internet at http://pubs.acs.org

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Author Contributions

The manuscript was written through contributions of all authors. All authors have given approval to the final version of the manuscript. ‡ These authors contributed equally.

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REFERENCES


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