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Introduction

Metal halide organic–inorganic perovskites represent an actively growing class of multipurpose materials. Classical 3D hybrid halide perovskites are compounds of the general formula ABX₃, where A is an organic cation, B is a divalent cation (typically Pb²⁺, Sn²⁺, Ge²⁺) and X is halogen. They attract attention mainly as semiconductors and have been recently investigated for their ability to serve as active elements in solar cells,^{1–5} lasers,^{6,7} photodetectors,^{8,9} light emitting diodes,^{10,11} for water splitting^{12,13} *etc.* One of the most studied and applied hybrid perovskites nowadays is CH₃NH₃PbI₃ because of the outstanding combination of relatively simple and low-cost fabrication techniques, tunability of band gap through chemical modification,¹⁴ high carrier mobility, long lifetime, long diffusion length, and high absorption coefficient, *etc.*¹⁵

A relatively easy way to modify the chemical composition of perovskites is through the introduction of different organic cations and inorganic anions, which lead to obtaining hundreds

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Hybrid organic-inorganic perovskites as microwave radiation switches[†]

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Hybrid organic–inorganic perovskites are an intensively growing class of functional materials that attract much attention due to their various practical and important functionalities. In this paper, we show the ability of two hybrid perovskites, namely $CH_3NH_3PbI_3$ and $(C_5H_{11}NH_3)_2PbI_4$, to act as phase transition (PT) microwave switches. Microwave switching was studied as a consequence of PTs which occur at technologically attractive temperatures – slightly above room temperature: 330 K for $CH_3NH_3PbI_3$ and 320 K for $(C_5H_{11}NH_3)_2PbI_4$. It was found that, at selected frequencies, $(C_5H_{11}NH_3)_2PbI_4$ is characterized by a very high transmission value of -0.4 dB which changes to -4.4 dB upon transition to a high-temperature phase. On the contrary, the transmission of $CH_3NH_3PbI_3$ at room temperature is deficient (-25 dB) and increases just slightly upon transformation into a high-temperature phase. The microwave absorption changes by 38% for $(C_5H_{11}NH_3)_2PbI_4$ and by 0.6% for $CH_3NH_3PbI_3$ at selected frequencies. Notably, absorption in $(C_5H_{11}NH_3)_2PbI_4$ changes from 9.9% at room temperature to 48.2% above the temperature of the PT, which is a five-fold increase. Our measurements show that $CH_3NH_3PbI_3$ is rather suitable for application as microwave absorber, while $(C_5H_{11}NH_3)_2PbI_4$ can be efficiently used as an active material in microwave switches.

of compounds, belonging to this class.¹⁶ Besides 3D hybrid perovskites, 2D analogues of this class have gained widespread popularity. These perovskites feature 2D halometallic layers separated by bulky organic cations.¹⁷ 2D hybrid perovskites are much more common than the 3D ones due to the large variety of suitable cations that can support 2D structures.¹⁸

A lot of hybrid organic–inorganic perovskites demonstrate temperature-induced structural PTs, which are mostly associated with rearrangement of organic cations or tilting of BX₆ octahedra.¹⁸ Importantly, many physical properties of hybrid perovskites are associated with PTs, for example, dielectric transitions and ferroelectric transitions.¹⁹ At the same time, phase-transition hybrid perovskites can potentially be used as molecular switches of very different functionalities.

It has been repeatedly shown that hybrid organic–inorganic perovskites can serve as efficient absorbers of microwave radiation.^{20–22} However, their ability to switch microwave radiation upon a PT has majorly remained outside the scope of attention of researchers. There are several reports on the change of permittivity of hybrid perovskites in the GHz range^{23,24} upon PT which show prospects of microwave switching.

One of the challenges of today's industries that utilize microwave radiation (5G, SatCom applications, *etc.*) is the development of efficient reconfigurable components that can be tuned to support variable frequency bands and communication standards. In highly integrated devices, which have to



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Paper

support multiple communication standards simultaneously, a reconfigurable filter can help reduce the total number of filters which would benefit the miniaturization of devices. Switching with smart materials is considered to be an efficient alternative to classical techniques for microwave switching, such as p-i-n diodes,^{25,26} microelectromechanical systems,^{27,28} field effect transistors,^{29,30} etc. For example, successful switching has been shown by dielectric anisotropy control in liquid crystals.³¹ Additionally, there are numerous examples of microwave switching by PTs in germanium halogenides³² and vanadium dioxide.33,34 Our team has recently shown the ability of Fe^{II} coordination compounds with thermally-induced spin crossover to operate as efficient microwave switches.^{35,36} In this paper we study capabilities of two hybrid organic-inorganic perovskites to switch microwave radiation: 3D semiconductive perovskite CH₃NH₃PbI₃ (C₁PbI) and a 2D perovskite (C₅H₁₁NH₃)₂PbI₄ (C₅PbI).³⁷

Results and discussion

Phase transitions

C₁PbI and **C₅PbI** were chosen due to the temperatures of PTs. While both perovskites display multiple PTs, we focused our attention in this work on microwave switching during the transitions that occur at technologically attractive temperatures – slightly above room temperature. PTs in both the obtained samples were studied using differential scanning calorimetry (Fig. 1). **C₁PbI** displayed a temperature-induced PT at 333 K in heating mode and at 327 K in cooling mode. According to the previously reported data, this transition corresponded to the change in crystallographic space group from *I*4/*mcm* to *Pm*3*m* at low and high temperature, respectively.³⁸ In the case of **C₅PbI**, PT occurred at 323 K upon heating and 311 K upon cooling. According to Billing *et al.*, this PT corresponded to the structural change, namely space group *P*2₁/*a* at low-temperature changes to



Fig. 1 DSC curves of C_1PbI (a) and C_5PbI (c) showing temperatureinduced PT associated with structural changes. Crystal structures of C_1PbI (b) and C_5PbI (d) showing different packing before and after PT. Crystal structures are depicted using CIF files provided in ref. 37 and 39 for C_1PbI and C_5PbI , respectively.



Fig. 2 Optical reflectance measurements for C_5Pbl , displaying a color change associated with PT.

Pbca at high temperature.³⁷ PT in C_5 PbI was additionally monitored by following optical reflection (Fig. 2), which showed similar temperatures of transition. Such an experiment was not possible for C_1 PbI because of its black color which essentially does not change upon PT.

Microwave transmission and reflection

When a microwave radiation falls on a material, the incident energy is divided into three parts: energy that is reflected back, transmitted through, and absorbed by a material.⁴⁰

To fully characterize the way C_1PbI and C_5PbI interacted with microwave radiation, both the microwave transmission and reflection were measured using a scalar network analyzer. The S_{11} parameter corresponded to the reflection of microwave radiation at port 1 which occurred due to impedance mismatch. When S_{11} is equal to 0 dB, complete reflection from port 1 occurred, while S_{11} values of -10 dB or lesser corresponded to a low reflection, indicating good impedance matching. S_{21} parameter characterized microwave transmission from port 1 to 2: S_{21} value of 0 dB indicated a 100% transmission without losses, while all the values below 0 dB indicated losses of the signal.³⁴

Powdered samples of C_1PbI and C_5PbI were pressed inside the Ka-band rectangular hollow waveguide. The spectral variations of microwave transmission and reflection have been measured in 26.0–37.5 GHz frequency ranges at variable temperatures. C_1PbI at room temperature (291 K) was characterized by very low transmission values, which ranged in between (-24.5) and (-29.5) dB depending on frequency (Fig. 3a). Such low values meant that microwave radiation was essentially not transmitted through the sample. Microwave transmission through C_1PbI increased slightly upon heating to the temperatures of PT; it was found to lie in the range of (-17.2)-(-25.7) dB at 348 K. Temperature variation in the microwave transmission for C_1PbI at 30.0 GHz is shown in Fig. 3b.

Microwave transmission measurements for C_5 PbI revealed a completely different result. Very high values of transmission (Fig. 3c) were observed at room temperature. Importantly,



Fig. 3 Spectral variation in microwave transmission (dB) for C_1PbI (a) and C_5PbI (c) with temperature obtained in a matched waveguide. Extracted values of transmission vs. temperature for C_1PbI (b) and C_5PbI (d) are given for selected frequencies. These curves demonstrate a microwave transmission change upon PT.

microwave transmission was -0.4 dB at 35.3 GHz and 293 K, which indicated almost complete transmission of radiation through the sample with minimal losses. An abrupt decrease in the transmission was observed upon heating to 338 K (Fig. 3d). S_{21} equaled -4.4 dB in the high-temperature phase, which indicated considerable rise of absorption and/or reflection by the sample. Some slight differences in transition temperatures and hysteresis width observed in microwave transmission and DSC experiments were associated with the settling mode of S_{21} measurements. Large-sized samples and the waveguide itself required thermalization for 10–15 min at each temperature before the measurements. Some small temperature oscillations altered the hysteresis in the measured curve. Small kinetic effects could also be suggested.

A small temperature-dependent drift of microwave transmission was noted in the regions above and below PT. In order to investigate temperature effects on S_{21} that were not associated with PT, this parameter was studied for $(C_4H_9NH_3)_2PbBr_4$ $(C_4 PbBr)$ above room temperature. The material crystalized in Pbca space group at room temperature and was isostructural to the high-temperature form of C₅PbI.^{41,42} DSC measurements revealed the absence of any PT in the studied temperature range for C₄PbBr (Fig. S1, ESI[†]). Temperature dependent measurements of microwave transmission (Fig. S2, ESI†) showed a slight gradual decrease from -3.6 dB at 301 K to -4.6 dB at 348 K (defined at 35.3 GHz). Such a drift of S_{21} with temperature was possibly associated with slight changes of permittivity with temperature in this sample.43 Simultaneously, C₅PbI S₂₁ switching was intensified by structural deformations due to the PT and the observed decrease in transmission significantly exceeded pure temperature-related effects.



Fig. 4 Spectral variation of microwave reflection (dB) of C_1PbI (a) and C_5PbI (d) with temperature. Reflection vs. temperature plots for C_1PbI (b and c) and C_5PbI (e and f) at selected frequencies are plotted.

As for the microwave reflection, S_{11} spectra of **C₁PbI** were characterized by the presence of reflection bands with minima at 28.3 and 32.5 GHz (Fig. 4a). Transition to the high-temperature phase was followed by a slight change in the peaks' intensity while their position remained almost unchanged. Depending on the frequency, either a decrease or an increase in reflection could be achieved (Fig. 4b and c).

For example, S_{11} was -3.52 dB at 30 GHz and room temperature (293 K). This value increased slightly upon heating with transition to the high-temperature (348 K) phase; it reached -3.32 dB. If analysis was performed at 27 GHz, reflection at room temperature equaled -3.7 dB. This value decreased to -4.0 dB under heating.

Similar to microwave transmission, reflection measurements revealed more significant changes for C_5PbI upon PT. This perovskite was characterized by the presence of a S_{11} band at 34.6 GHz. During the transition to high-temperature phase, a significant shift of band's position towards lower frequencies (up to 33.0 GHz) was observed. Such shift can indicate the increase of permittivity of this material upon PT. Either a decrease or increase in reflection can be observed for C_5PbI depending on the operating frequency. For example, S_{11} equals -9.1 dB at 35.4 GHz and 303 K, indicating low reflection from the sample. This correlates well with very high transmission at the same frequency and temperature. With the transition to the



Fig. 5 Absorption of microwave radiation by C_1PbI (a) and C_5PbI (b) at selected fixed frequencies. The dependencies show that C_5PbI can effectively switch microwave radiation, while C_1PbI should be rather used as a microwave absorber.

high-temperature phase (338 K), S_{11} increases to -5.4 dB at this frequency. An inverse effect was observed at 32.5 GHz: S_{11} decreased from -6.2 to -9.5 dB upon transition from the low-temperature (303 K) to the high-temperature phase (338 K).

Effective microwave absorption value of materials can be described by the following equation:^{44,45}

$$A_{\rm eff} = \frac{1 - T - R}{1 - R},\tag{1}$$

where T is transmission and R is reflection.

Changes in effective absorption upon PT for C_1PbI and C_5PbI at selected frequencies are shown in Fig. 5. The absorption for C_1PbI sample was approx. 99.6% at 303 K (at 30.0 GHz). During the temperature-induced PT, this value decreased slightly and reached 99.0%. These observations show that C_1PbI is rather suitable for application as a microwave absorber, while its switching ability associated with a PT above room temperature is poor.

At the same time, effective absorption of C_5PbI sample was 9.9% at 303 K. This parameter drastically increases with a transition to the high-temperature phase and reaches 48.2% at 338 K. Importantly, the microwave radiation absorption in C_5PbI increases by a factor of five, which means that C_5PbI can be prospective material for microwave switches.

Such a strong difference in absorption between the two investigated HOIPs can be associated with the difference in their crystal structures. The 3D packing of C_1PbI is known to be responsible for its semiconducting properties, which, in turn, induce strong microwave absorption of this material. Simultaneously, C_5PbI with its 2D crystal structure does not display any



Fig. 6 Dependence of complex permittivity (ϵ' and ϵ'') of C_1PbI (a) and C_5PbI (b) on frequency. Cole–Cole plots for C_1PbI (c) and C_5PbI (d) in 26–38 GHz frequency range.

semiconducting properties, and, thus, its microwave absorption is considerably lower. At the same time, during structural transition to the high-temperature phase of C_5PbI , an increase in cell volume and, consequently, a decrease in the material's density occurs. This effect, in turn, may facilitate orientation of dipoles, leading to an increase in microwave absorption due to dielectric relaxation mechanisms.

In order to study the mechanism of microwave absorption in C_1PbI and C_5PbI , complex dielectric permittivity of these materials was measured as a function of frequency at room temperature (Fig. 6a and b). It can be seen that the ε' values of both the materials displayed minor changes depending on frequency. The real part of C_1PbI permittivity was 3.47 at 26 GHz; it slightly dropped with an increase in frequency and reached 3.37 at 38 GHz. The imaginary part of C_1PbI permittivity ity was 0.16 at 26 GHz, which changed to 0.28 and 0.24 at 32.4 and 38.0 GHz, respectively. The values found in this experiment were close to previously shown ones for C_1PbI in 2–18 GHz range at room temperature.⁴⁶

The real part of permittivity for C_5 PbI was close to the value obtained for C_1 PbI: ε' of this perovskite was 3.29 at 26 GHz, 3.39 at 34.0, and 3.33 at 38.0 GHz. At the same time, ε'' values of C_5 PbI were lower than measured values for C_1 PbI. The imaginary part of permittivity for pentylammonium perovskite was 0.16 at 26 GHz, ε'' first dropped with a frequency increase and reached its minimum value of 0.06 at 33 GHz. With further increase in frequency, ε'' slightly increased again and reached 0.16 at 38 GHz. These dependences of the complex permittivity on frequency cannot be described *via* any simple model, *i.e.*, more information on the mechanism of dielectric relaxation is necessary.

Absorption of microwave radiation in such materials is typically realized due to dielectric losses caused by dipole orientation polarization. In case of the studied materials, free organic cations in the pores of inorganic frameworks could serve as efficient polarization centers. Those dipoles re-orient

under the influence of thermal motion upon removal of external electromagnetic field; such process can be described as dipolar polarization relaxation.⁴⁰ In such systems, dependence between ε' and ε'' can be described by the Debye theory:

$$\left(\varepsilon' \frac{\varepsilon_{\mathrm{s}} + \varepsilon_{\infty}}{2}\right)^2 + \left(\varepsilon''\right)^2 = \left(\frac{\varepsilon_{\mathrm{s}} - \varepsilon_{\infty}}{2}\right)^2,$$

where ε_{∞} is the relative permittivity at infinite frequency and ε_{s} is the static permittivity. Cole-Cole semicircles, which can be observed in $\varepsilon' vs. \varepsilon''$ dependence shown in Fig. 6c and d, represent the Debye relaxation process. Importantly, each semicircle describes one relaxation process. C₁PbI demonstrated three semicircles in the investigated frequency range, while in the case of C₅PbI, only one typical Cole-Cole semicircle could be observed. Therefore, a higher number of relaxation processes and higher values of ε'' with similar values of ε' explain the higher absorptivity of C₁PbI.

Transmission and reflection spectra obtained in VNA measurements at room temperature for C₁PbI and C₅PbI are shown in Fig. S4 (ESI[†]). While the values observed for C₅PbI were similar for scalar and vector measurements, S₂₁ parameters observed for C₁PbI in the second experiment were lower. The observed distinctions were associated with different types of waveguides used in two measurements (rectangular waveguide for scalar temperature-dependent experiments and coaxial waveguide for vector measurements) in which significantly different dispersion and attenuation were realized. Additionally, dilution with PMMA polymer, which was used for fabrication of toroidal samples for VNA, could also influence transmission/reflection values.

To the best of our knowledge, the current work was the first study of temperature-dependent microwave transmission/ reflection of hybrid perovskites, which exhibited PT. There have been several previous studies on temperature-dependent permittivity of 2D²⁴ perovskites at 9.5 GHz and 3D²⁴ perovskites at 90 GHz, where PT were observed. In both cases, an increase in microwave permittivity was observed upon heating, with a transition occurring from a more ordered to a less ordered state. Also, there were multiple reports on significant absorption of microwave frequency radiation by hybrid perovskites at room temperature.^{21,22,46,47} Thus, changes in permittivity in the microwave range upon PT and high absorption ability, in combination with a huge variety of accessible hybrid organic-inorganic perovskites, made this class of materials very attractive for use in different microwave technologies. Considering the low number of studies published on hybrid perovskites interaction with microwave radiation and the promising results obtained in the current work, further investigations in this direction are worth performing.

Experimental

Materials

Lead iodide was purchased from Acros organics, pentylamine, buthylamine and methylammonium iodide were purchased from UkrOrgSyntez Ltd and used as received.

Synthesis

Methylammonium lead iodide CH₃NH₃PbI₃ (C₁PbI) was obtained according to a slightly modified procedure offered by Weber et al. in 1978.⁴⁸ Two grams of PbI₂ (4.3 mmol, 1 eq.) was dissolved in 6 ml of concentrated HI with heating. Thereafter, 828 mg (5.16 mmol, 1.2 eq.) of CH₃NH₃I was added to 1 ml of HI. The mixture was transferred to an oven and kept at 45 °C for 10 h. During this time, a black precipitate was formed. After that, the mixture was centrifuged and dried in air. Yield was 30% (800 mg).

Pentylammonium lead iodide (C5H11NH3)2PbI4 (C5PbI) was obtained according to a slightly modified procedure offered by Billing et al. in 2007.³⁷ A gram of PbI₂ (2.2 mmol, 1 eq.) was dissolved in 3 ml of concentrated HI with heating. Thereafter, 452 mg (5.28 mmol, 2.4 eq.) of $C_5H_{11}NH_2$ in 1 ml of HI was added. Bright-orange precipitate was formed immediately. The mixture was cooled down, centrifuged and dried in air. The yield was 95% (1.86 g).

Butylammonium lead bromide $(C_4H_9NH_3)_2PbBr_4$ (C₄PbBr) was obtained according to a slightly modified procedure offered by Karunadasa et al. in 2017.⁴² PbBr₂ (750 mg, 2.0 mmol, 1 eq.) was dissolved in 2 ml of concentrated HBr with heating. Thereafter, 350 mg (4.8 mmol, 2.4 eq.) of C₄H₉NH₂ was added to 1 ml of HBr. White crystals were formed after cooling the solution, collected, dried on air and ground for further experiments. Yield was 70% (740 mg).

Measurements

Optical reflectance measurements were conducted using a Linkam DSC 600 temperature-controlled microscope stage cryostat and Optica SZM-1 stereomicroscope equipped with a Sigeta UCMOS 1300 camera. The processing of images was performed with ImageJ software. Reflectance measurements were performed at a heating/cooling rate of 10 K min⁻¹.

Differential scanning calorimetry measurements were performed with a Linkam DSC 600. DSC profiles were recorded with a heating/cooling rate of 10 K min⁻¹.

Microwave transmission/reflection measurements were carried out with a P2-65 scalar microwave network analyzer operating in the Ka frequency band. An analyzer was equipped with a hollow rectangular waveguide (7.20 mm \times 3.40 mm). The powdered samples of C₁PbI and C₅PbI were pressed into rectangular pellets inside the waveguide by applying ca. 20 MPa pressure (d = 6.00 mm). The waveguide filled with a perovskite sample was heated with an external thermostat for temperature-dependent measurements. The experiments were conducted in a settling mode at which the waveguide with the sample inside was thermalized for 10-15 min at each temperature value before collecting the measurements. Narrow thermal intervals for microwave measurements were selected for each perovskite from the preceding optical and DSC experiments.

Complex dielectric permittivity was measured using Keysight PNA N5227A vector network analyser. For this experiment, C₁PbI and C₅PbI were mixed with 50% of inert poly(methyl methacrylate) matrix and cut as coaxial rings of toroidal shape with an outer diameter of 1.85 mm and inner diameter of 0.82 mm. Thickness of both samples was 6 mm. Permittivity measurements of PMMA matrix are given in Fig. S3 (ESI⁺).

The PXRD patterns were acquired on Shimadzu XRD-6000 diffractometer using Cu-K α radiation (10–50 °C range, 0.05 °C steps) (Fig. S5 and S6, ESI[†]). The measurements revealed some differences in peak intensities in comparison to simulated PXRD patterns. Such observations can be explained by preferential orientation in the samples. However, the same way of sample preparation used for both PXRD and microwave transmission/reflection experiments suggested the possibility that orientation of samples in the two experiments was similar.

Conclusions

Here we show the ability of hybrid organic-inorganic perovskites to change their microwave absorption during the temperature-induced structural PT. Methylammonium lead iodide displayed very high absorption around room temperature, which decreased by less than 1% upon PT, showing that this perovskite was rather suitable for use as an absorber, but not a switch. Pentylammoniun lead iodide displayed moderate absorption at room temperature, which increased five-fold upon PT, which highlighted this material as an efficient microwave switch. These results make an important contribution to the creation of preliminary dataset of microwave absorbing and switching properties of hybrid organic-inorganic perovskites.

Author contributions

Olesia I. Kucheriv, Volodymyr V. Zagorodnii, Vilen L. Launets investigation, Viktor V. Oliynyk and Igor O. Fritsky - data curation, Il'ya A. Gural'skiy - conceptualization and supervision. All authors contributed to the writing of this manuscript.

Conflicts of interest

There are no conflicts to declare.

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