



Cite this: *Phys. Chem. Chem. Phys.*, 2024, 26, 13675

Systematic study of ionic conduction in silver iodide/mesoporous alumina composites 2: effects of silver bromide doping†

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In our preceding paper (Y. Fukui *et al.*, *Phys. Chem. Chem. Phys.*, 2023, **25**, 25594–25602), we reported a systematic study of the Ag⁺-ion conducting behaviour of silver iodide (AgI)-loaded mesoporous aluminas (MPAs) with different pore diameters and AgI-loading ratios. By optimising the control parameters, the Ag⁺-ion conductivity has reached 7.2×10^{-4} S cm⁻¹ at room temperature, which is more than three orders of magnitude higher than that of bulk AgI. In the present study, the effect of silver bromide (AgBr)-doping in the AgI/MPA composites on Ag⁺-ion conductivity is systematically investigated for the first time, using variable-temperature powder X-ray diffraction, differential scanning calorimetry, and electrochemical impedance spectroscopy measurements. The AgBr-doped AgI/MPA composites, AgI-AgBr/MPA, formed a homogeneous β/γ -AgI-structured solid solution (β/γ -AgI_{ss}) for the composites with AgBr \leq 10 mol%, above which the composites underwent a phase separation into β/γ -AgI_{ss} and face-centred cubic AgBr solid solutions (AgBr_{ss}). The onset temperature of the exothermic peaks attributed to the transition from α -AgI-structured solid-solution phase to β/γ -AgI_{ss} or AgBr_{ss} decreased with increasing the AgBr-doping ratio. The room-temperature ionic conductivity of the AgI-AgBr/MPA composites exhibited a volcano-type dependence on the AgBr-doping ratio with the highest value (1.6×10^{-3} S cm⁻¹) when the AgBr content was 10 mol%. This value is more than twice as high as that of the highest conducting AgI/MPA found in our previous study.

Received 21st February 2024,
 Accepted 17th April 2024

DOI: 10.1039/d4cp00744a

rs.li/pccp

Introduction

Solid-state ionic conductors have been the subject of intense research as potential greener alternatives to their liquid counterparts for electrolytes in diverse electrochemical devices.^{1–7} Despite their inevitably reduced performance, solid-state electrolytes are essential for solving safety problems caused by liquid leakage, flammability, and corrosion. To date, the major drawback of such systems is their high-temperature operation, which impedes quick start-up and shutdown of cycles and expedites device degradation. Thus it is evident that a low-temperature operation is highly demanded for state-of-the-art electrochemical devices.^{8,9}

Ions in ionic solids are essentially immobile due to the electrostatic interactions between the oppositely charged ions. Among them, a silver ion with a 4d¹⁰ electron configuration is

particularly promising as a carrier ion in solid-state ionic conductors, primarily owing to its soft acidic nature.¹⁰ In particular, silver iodide (AgI) with a low room-temperature ionic conductivity (*ca.* 10^{-7} S cm⁻¹) exhibits an extraordinary high ionic conductivity (*ca.* 1 S cm⁻¹) in the high-temperature α -phase ($T > 147$ °C), where the melted Ag⁺ ions migrate in the immobile I⁻ sublattice.^{11–15} It is known that the reduction in the size of AgI particles leads to the stabilisation of the highly ion-conducting α -phase by virtue of the high surface energy; for example, the transition temperature from the α -phase to the low-temperature β - and γ -phases is significantly lowered to 37 °C for AgI nanoparticles with a diameter of 6.3 nm.^{16–19} However, organic polymers embedded with the AgI nanoparticles to prevent their aggregation apparently disturb the long-range Ag⁺-ion migration, which is critical for their practical use in the field of solid-state ionics. Another way to increase the room-temperature ionic conductivity of AgI is the introduction of interstitial Ag⁺ ions near the surface of the β/γ -AgI particles (even if AgI is not in the α -phase), because the formation energy of Frenkel pairs comprised of an interstitial Ag⁺ ion and a Ag⁺-ion vacant site decreases with decreasing particle size.²⁰ In addition, the incorporation of the AgI particles into insulating mesoporous metal oxides, such as

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† Electronic supplementary information (ESI) available: Fig. S1–S13. See DOI: <https://doi.org/10.1039/d4cp00744a>



alumina,^{21–32} silica,^{33–39} zeolite,⁴⁰ and zirconia,³³ is beneficial for enhanced ionic conductivity as a consequence of the space-charge layer formed at the heterogeneous interface.^{41–46}

Our group has recently reported the effect of the pore size of mesoporous alumina (MPA) and the filling level of AgI into MPA on Ag⁺-ion conductivity.⁴⁷ We found that the AgI/MPA composite, in which the pore space was nearly fully occupied by β/γ -AgI, exhibited the highest room-temperature ionic conductivity when MPA has an average pore size of 7.1 nm. Notably, the ionic conductivity, $7.2 \times 10^{-4} \text{ S cm}^{-1}$, is more than three orders of magnitude higher than that of the bulk β/γ -AgI (*ca.* $1.5 \times 10^{-7} \text{ S cm}^{-1}$).¹⁶ In this study, we focused on the solid-solution system with homovalent substitutions of immobile I[−] ions, because the substitution efficiently produces a local distortion in the lattice to induce the Frenkel defects.^{48–73} In fact, the bulk AgI–AgBr solid solutions showed a significant increase in ionic conductivity compared with that of the parent AgI, which motivated us to investigate the effect of the homovalent substitution on the Ag⁺-ion conductivity of the AgI/MPA composites. To the best of our knowledge, the simultaneous introduction of AgI and AgBr into mesoporous materials has not yet been achieved. This paper presents the dependence of the room-temperature structural and ion-conducting properties of AgI/AgBr-loaded MPA composites (hereafter AgI–AgBr/MPA) on the AgBr-doping level, where a MPA with the highest conducting AgI/MPA composite in ref. 47 (pore diameter: 7.1 nm) was utilised. In addition, we examined the variable-temperature structural, calorimetric, and ion-conducting properties, especially for the purpose of the assessment of the thermal stability of each phase in the AgI–AgBr/MPA composites.

Experimental section

Materials

In this study, we used a commercially available MPA (Sigma-Aldrich Japan) with a pore diameter of 7.1 nm and a pore volume of $0.455 \text{ cm}^3 \text{ g}^{-1}$, which were estimated based on the N₂ gas adsorption measurements. The MPA was dried at 500 °C for 6 h under N₂ atmosphere prior to use. AgI (99.0%, Kojima Chemicals Co., Ltd) and AgBr (97.0%, Wako Pure Chemical Industries, Ltd) were dried at 200 °C for 6 h under N₂ atmosphere prior to use. Adequate amounts of dried silver halides (AgX) and MPA were co-ground in an agate mortar under low-humidity conditions in a N₂-filled glovebox (volumetric moisture content: <0.35%) to yield the volumetric filling ratio of approximately 90%. Each mixed powder was pressed at 500 MPa for 1 min to obtain a compressed pellet with a diameter of 2.5 mm, followed by heat treatment at 600 °C for 20 h to introduce AgX (melting point: 552 °C for X = I and 432 °C for X = Br) into the MPA pores.

Characterization

Room-temperature powder X-ray diffraction (PXRD) measurements were performed with a Bruker D8 ADVANCE instrument

using Cu K α radiation ($\lambda = 1.5418 \text{ \AA}$), whereas variable-temperature PXRD measurements were performed with a Rigaku SmartLab 3 kW instrument using Cu K α radiation. The temperature was controlled using an Anton Paar DHS900 attachment at a rate of $5 \text{ }^\circ\text{C min}^{-1}$. Rietveld refinements to evaluate the ratio of the phases with α -AgI, β/γ -AgI, and AgBr-like structures were performed with fixed atomic coordinates. Scanning electron microscope (SEM) observation and energy dispersive X-ray spectroscopy (EDS) analysis were performed with a JEOL JSM-IT500HR at an accelerating voltage of 2 kV for SEM and 20 kV for EDS. N₂ gas sorption isotherms were measured at 77 K with a MicrotracBEL BELSORP-mini II volumetric adsorption system. The thermal properties were characterised by differential scanning calorimetry (DSC) thermograms using a NETZSCH DSC 3500 Sirius or a Rigaku Thermo plus EVO2 DSC8231 instrument equipped with N₂ cryostatic cooling. The composites, which were sealed in an aluminium pan, were heated from room temperature to 200 °C followed by cooling to 0 °C at a heating and cooling rate of $5 \text{ }^\circ\text{C min}^{-1}$. The ionic conductivities of the compressed pellets were measured by the AC impedance technique with an applied voltage of 100 mV using a Solartron SI 1260 Impedance/Gain-phase Analyzer and 1260A Dielectric Interface in the frequency range of 1 MHz to 0.1 Hz. Both sides of the pellets were painted with gold paste (Tokuriki, 8560-1A) and measurements were performed in a temperature-controlled chamber ESPEC ST-110. The impedance was determined from the first real-axis touch-down point in the Nyquist plot.

Results and discussion

The AgBr-doped AgI/MPA composites, AgI–AgBr/MPA, were prepared by co-grinding of AgI : AgBr (100 : 0–70 : 30 molar ratio) with MPA (pore diameter: 7.1 nm; Fig. S1, ESI[†]) followed by heat treatment at 600 °C (Fig. S2 and S3, ESI[†]; hereafter denoted as **Br0–Br30**, respectively; Table 1). All the peaks observed in the PXRD patterns of the mixtures before heat treatment can be assigned to pristine β -AgI, γ -AgI, or AgBr phases (Fig. 1(a)), and the relative intensity of the AgBr peaks steadily increases with increasing AgBr content. The heat treatment led to the broadening of the PXRD peaks due to the reduction in the crystallite size of AgI/AgBr (Fig. 1(b)), which is indicative of the incorporation into the mesopores of MPA.⁴⁷ To confirm the incorporation of AgI/AgBr into the MPA pores, EDS mappings of the AgI–AgBr/MPA composite with AgBr = 10 mol% (*i.e.*, **Br10**) were performed before and after the heat

Table 1 Composition ratios of AgX (X = Br or I) in AgI–AgBr/MPA composites and their notations

Notation	AgI (mol%)	AgBr (mol%)
Br0 ⁴⁷	100	0
Br1	99	1
Br5	95	5
Br10	90	10
Br20	80	20
Br30	70	30



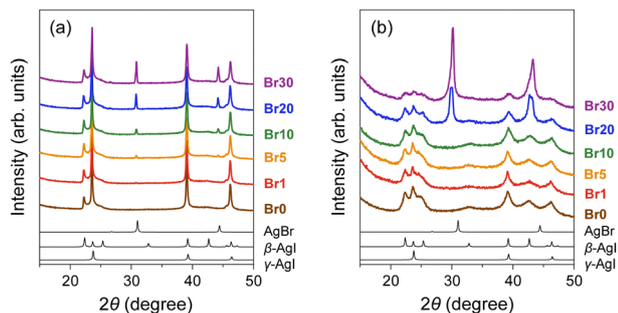


Fig. 1 PXR D patterns of AgI-AgBr/MPA composites with various AgI:AgBr molar ratios (a) before and (b) after heat treatment at 600 °C for 20 h (brown: **Br0**, red: **Br1**, orange: **Br5**, green: **Br10**, blue: **Br20**, purple: **Br30**). Black lines show the simulated patterns of AgBr, β -AgI, and γ -AgI (from the top).

treatment. Whereas a clear distinction between Al and Ag/I/Br was found in the mapping images taken before heat treatment (Fig. 2(a)), the composite exhibited a homogeneous elemental distribution after heat treatment (Fig. 2(b)). Notably, the SEM image showed that the surface morphology of the MPA remained intact during heat treatment (Fig. S4, ESI[†]).

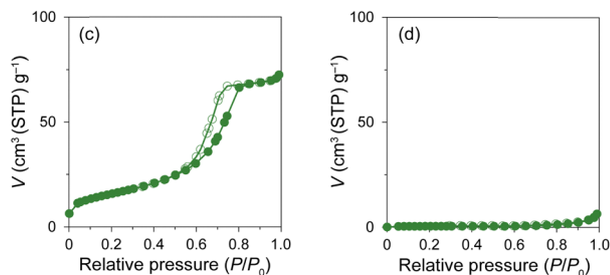
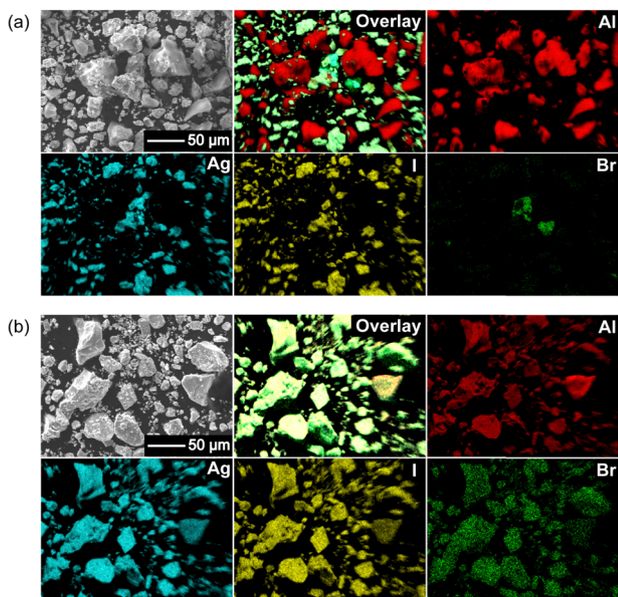


Fig. 2 SEM images and EDS mappings (overlay, Al, Ag, I, and Br) of **Br10** (a) before and (b) after heat treatment. N_2 gas adsorption (closed circles) and desorption (open circles) isotherms of **Br10** (c) before and (d) after heat treatment.

Furthermore, N_2 gas sorption measurements demonstrated that the mesopores, which are responsible for the significant uptake at approximately $P/P_0 = 0.7$ (Fig. 2(c)), disappeared during heat treatment (Fig. 2(d)) as a consequence of the successful incorporation of AgI/AgBr into the mesopores of MPA. To the best of our knowledge, the simultaneous incorporation of AgI and AgBr into mesoporous materials have been reported first time in this study.

The PXR D peaks of the composites with $AgBr \leq 10$ mol% (*i.e.*, **Br0–Br10**) were assigned only to β - or γ -AgI phases, whereas the PXR D patterns of the composites with $AgBr > 10$ mol% (*i.e.*, **Br20** and **Br30**) exhibited additional peaks that were assigned to the face-centred cubic (fcc) AgBr phase (Fig. 1(b)). Therefore, the composites with $AgBr \leq 10$ mol% form a homogeneous solid solution with β/γ -AgI_{ss}, whereas those with $AgBr > 10$ mol% underwent a phase separation into the aforementioned β/γ -AgI_{ss} and another solid-solution phase in which AgI was dissolved in the AgBr lattice (hereafter AgBr_{ss}). Furthermore, the formation of the solid solution of AgI in the AgBr lattice is evident from the shift of the (200) reflection of AgBr_{ss} to the lower angle side than that in parent bulk AgBr (*e.g.*, 29.9° for **Br20** vs. 30.9° for parent bulk AgBr) as a reflection of the partial substitution of Br atoms with larger I atoms in the AgBr lattice (ionic radius: 1.96 Å for Br⁻, 2.19 Å for I^{-74,75}). On the other hand, the PXR D peaks of β/γ -AgI_{ss} were barely shifted by the heat treatment, which can be attributed to the fact that the amount of AgBr dissolved in AgI is less than that of AgI dissolved in AgBr as observed in bulk AgI-AgBr mixed crystals.^{52,55,73,76,77}

To investigate the effect of AgBr-doping on the thermal behaviour, the DSC profiles of the AgI-AgBr/MPA composites were measured during the heating and cooling processes. Focusing on the cooling process, two exothermic peaks were observed for the composites with $AgBr \leq 10$ mol% (*i.e.*, **Br0–Br10**; Fig. 3(a)–(d)). Given that the composites remain a single β/γ -AgI_{ss} phase in this AgBr content region at room temperature (Fig. 1(b)), both DSC peaks can be assigned to the structural phase transition from the α - to β/γ -phases of AgI_{ss}. In our previous paper,⁴⁷ we identified that the high- and low-temperature DSC peaks observed for the parent AgI/MPA composite (*i.e.*, **Br0**) are associated with the bulk-like AgI occupying the core and AgI near the surface of the pores, respectively. Therefore, it is possible that the same assignment holds for AgI-AgBr/MPA composites. For the composites with $AgBr > 10$ mol% (*i.e.*, **Br20** and **Br30**), only one exothermic peak, possibly corresponding to the low-temperature side, was observed, as shown in Fig. 3(e) and (f). Because the heating scan of the DSC profile of **Br20** exhibited two distinct endothermic peaks assignable to the β/γ -AgI_{ss}-to- α -AgI_{ss} and AgBr_{ss}-to- α -AgI_{ss} (α -AgI_{ss}: solid-solution phase with α -AgI structure) at *ca.* 110 and 170 °C, respectively,⁷⁰ it is likely that the corresponding exothermic DSC peaks merge into the broad peak in the cooling process. This is reinforced by the very broad temperature range of the phase transitions observed in the temperature-dependent PXR D measurements (see below). Furthermore, the



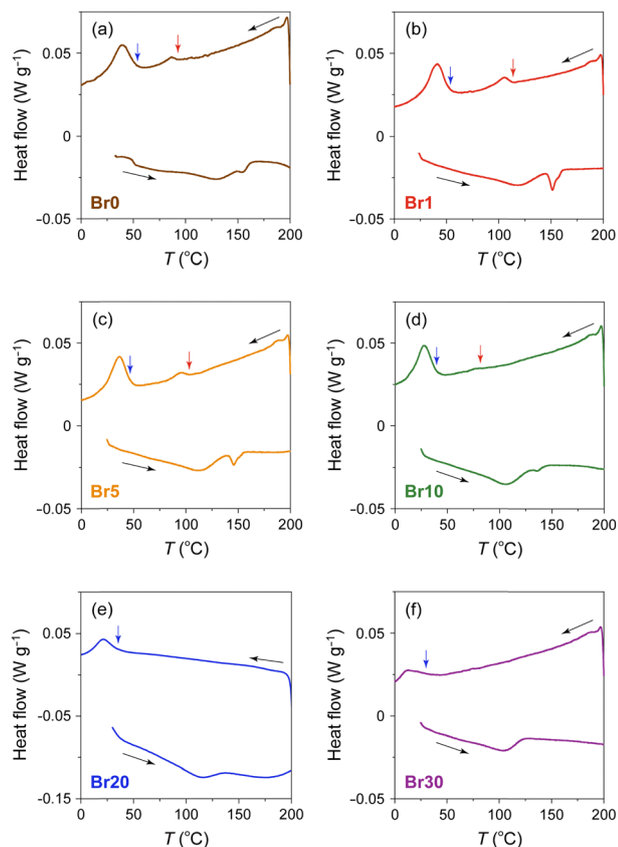


Fig. 3 DSC profiles of AgI–AgBr/MPA composites with various AgI:AgBr molar ratios ((a) **Br0**, (b) **Br1**, (c) **Br5**, (d) **Br10**, (e) **Br20**, and (f) **Br30**). The red and blue arrows indicate the onset temperatures of the transition on the high- and low-temperature sides, respectively, in the cooling process.

transition enthalpy comparable to those of the composites with $\text{AgBr} \leq 20 \text{ mol}\%$ (Fig. 4(a)) supports the assignment. On the other hand, the transition enthalpy of **Br30** is significantly lower than the others, which may be associated with the significant increase in the AgBr_{ss} -to- $\alpha\text{-AgI}_{\text{ss}}$ transition temperature as observed in bulk AgI–AgBr mixed crystals (*ca.* 200 °C for $\text{AgBr} = 30 \text{ mol}\%$).^{52,55,73,76,77} The incomplete transition to $\alpha\text{-AgI}_{\text{ss}}$ in the measured temperature range was verified by the temperature-dependent PXRD measurements (see below).

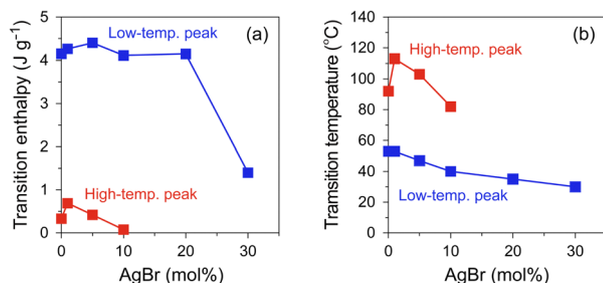


Fig. 4 AgBr-doping ratio dependence of (a) transition enthalpy and (b) transition temperatures determined by DSC profile in the cooling process (red: high-temperature peak, blue: low-temperature peak).

As shown in Fig. 4(b), the onset temperatures of the peaks decreased with increasing AgBr-doping ratio. The rapid decrease in the transition temperatures of the composites with $\text{AgBr} \leq 10 \text{ mol}\%$ can be primarily attributed to the lattice distortion induced by AgBr-doping, which causes the increase in the Frenkel defect concentration to stabilise the high-temperature phase. Furthermore, the phase separation observed in the composites with $\text{AgBr} > 10 \text{ mol}\%$ may suppress the $\alpha\text{-AgI}_{\text{ss}}$ -to- $\beta/\gamma\text{-AgI}_{\text{ss}}$ transition in the cooling process.

Variable-temperature PXRD measurements of the AgI–AgBr/MPA composites were performed to understand their phase transition behaviour from a structural perspective. Fig. 5(a) shows the PXRD patterns of **Br10** during heating and cooling processes. The as-prepared composite demonstrated the PXRD peaks assignable to $\beta/\gamma\text{-AgI}_{\text{ss}}$ at 30 °C with a trace of AgBr_{ss} ($2\theta = 30.3^\circ$), which disappeared after cyclic heating and cooling treatments implying the metastable nature of the phase in this composite. On heating, the $\beta/\gamma\text{-AgI}_{\text{ss}}$ peaks began to disappear at *ca.* 100 °C with the concurrent appearance of the $\alpha\text{-AgI}_{\text{ss}}$ peak, and eventually became invisible above *ca.* 150 °C. The phase transition was more explicit when the ratio of each phase determined by Rietveld analysis is plotted as a function of temperature (Fig. 5(b)). Refinements were performed under the assumption that only I atoms occupy the halogen sites in AgI_{ss} , whereas only Br atoms occupy the sites in AgBr_{ss} . In the subsequent cooling process, the $\alpha\text{-AgI}_{\text{ss}}$ phase was maintained down to *ca.* 60 °C, below which it underwent a reversible transformation to the $\beta/\gamma\text{-AgI}_{\text{ss}}$ phase (Fig. 5(c)). It is readily apparent that the structural transition occurs at which an intense peak was observed in the DSC profile (Fig. 3(d)), supporting the above identification of the phase transition. In addition, we measured the temperature dependence of the PXRD patterns of **Br20** (Fig. 5(d)) and **Br30** (Fig. 5(g)), which underwent a phase separation into $\beta/\gamma\text{-AgI}_{\text{ss}}$ and AgBr_{ss} at room temperature. The molar ratios of the AgI/AgBr phases were estimated to be 33 : 67 for **Br20** and 14 : 86 for **Br30** based on the Rietveld refinement, indicating that a significant level of I atoms was incorporated in AgBr_{ss} as mentioned above. For **Br20**, the gradual disappearance of both phases entailed the appearance of the $\alpha\text{-AgI}_{\text{ss}}$ phase in the temperature range of 110–140 °C for $\beta/\gamma\text{-AgI}_{\text{ss}}$ and 110–180 °C for AgBr_{ss} (Fig. 5(e)), which are possibly associated with the broad endothermic DSC peaks at approximately 110 and 170 °C, respectively, in the heating process. In the cooling process, the $\alpha\text{-AgI}_{\text{ss}}$ phase was reversibly transformed to the $\beta/\gamma\text{-AgI}_{\text{ss}}$ and AgBr_{ss} phases (Fig. 5(f)), although there was only one exothermic DSC peak at *ca.* 40 °C. Considering that gradual phase transition processes were observed in the PXRD measurements, the DSC peaks assigned to the two transitions may have merged into a broad peak as mentioned above. The temperature-dependent PXRD data of **Br30** bear a few similarities to **Br20**, along with several striking differences. As shown in Fig. 5(h), the $\beta/\gamma\text{-AgI}_{\text{ss}}$ phase in **Br30** disappeared at *ca.* 120 °C in the heating process, which is associated with the broad endothermic DSC peak at approximately 110 °C. The corresponding $\alpha\text{-AgI}_{\text{ss}}$ -to- $\beta/\gamma\text{-AgI}_{\text{ss}}$ transition, which is responsible for the DSC peak with the onset



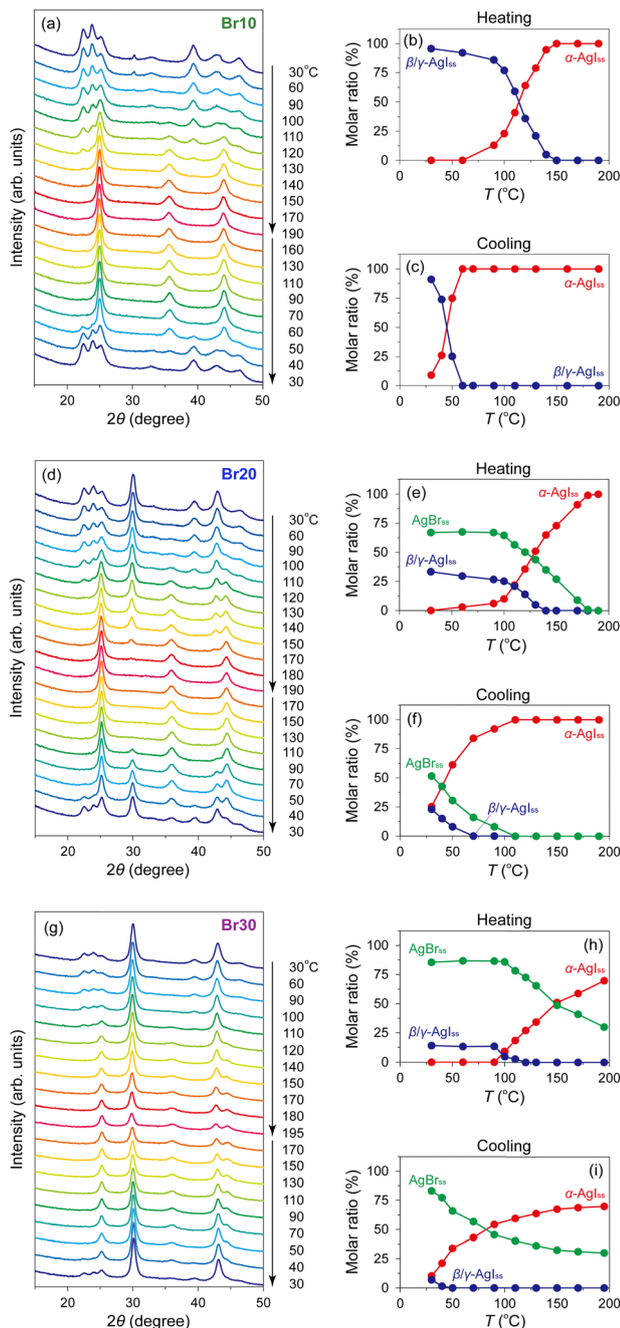


Fig. 5 Variable-temperature PXRD patterns of (a) **Br10**, (d) **Br20**, and (g) **Br30** in the heating and cooling processes. Temperature dependences of the molar ratios of α - (red) and β/γ - (blue) AgI_{ss} phases of **Br10** in the heating (b) and cooling (c) processes and the molar ratios of α - AgI_{ss} (red), β/γ - AgI_{ss} (blue), and AgBr_{ss} (green) phases of **Br20** and **Br30** in the heating ((e) and (h)) and cooling ((f) and (i)) processes. The ratios were determined by the Rietveld analyses.

temperature at 30 °C, was observed at approximately 40 °C in the cooling process (Fig. 5(i)). It is noteworthy that the AgBr_{ss} phase in **Br30** remained stable even at 195 °C, which must cause the observed low transition enthalpy (Fig. 4(a)). The phase diagram of the AgI - AgBr /MPA composites, in which the phase boundaries were defined when the molar ratio of

β/γ - AgI_{ss} or AgBr_{ss} phase reaches 50% relative to that at 30 °C, resembles that of bulk AgI - AgBr mixed crystals⁵⁵ with a slight shift towards a lower temperature (Fig. S5, ESI†).

The ionic conductivity (σ) of the AgI - AgBr /MPA composites with various $\text{AgI}:\text{AgBr}$ molar ratios was measured using AC impedance spectroscopy (Fig. 6 and Fig. S6, ESI†). The room-temperature value ($\sigma_{25\text{ }^\circ\text{C}}$) lies in the order of 10^{-4} - 10^{-3} S cm^{-1} , which is significantly higher than that of bulk AgI (*ca.* 1.5×10^{-7} S cm^{-1}).¹⁶ As shown in Fig. 6, the composites with $\text{AgBr} \leq 10$ mol% exhibited an Arrhenius-type ionic conduction up to approximately 140 °C, whereas those with $\text{AgBr} > 10$ mol% showed a moderate continuous increase in σ with increasing temperature up to the highest measured temperature (200 °C) without any significant slope change. Considering the fact that the composites with $\text{AgBr} > 10$ mol% formed a phase separation between β/γ - AgI_{ss} and AgBr_{ss} at room temperature, it is possible

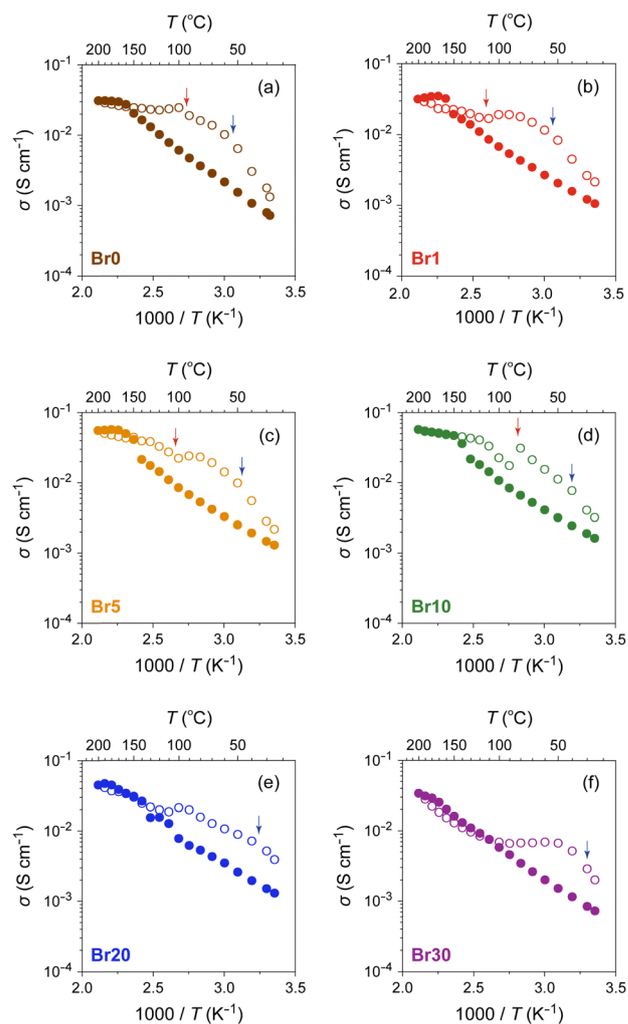


Fig. 6 Temperature dependence of σ for AgI - AgBr /MPA composites with various $\text{AgI}:\text{AgBr}$ molar ratios ((a) **Br0**, (b) **Br1**, (c) **Br5**, (d) **Br10**, (e) **Br20**, and (f) **Br30**) in heating (closed circles) and cooling (open circles) processes. The red and blue arrows indicate the onset temperatures of the transition on the high- and low-temperature sides, respectively, in the DSC cooling process (Fig. 3).



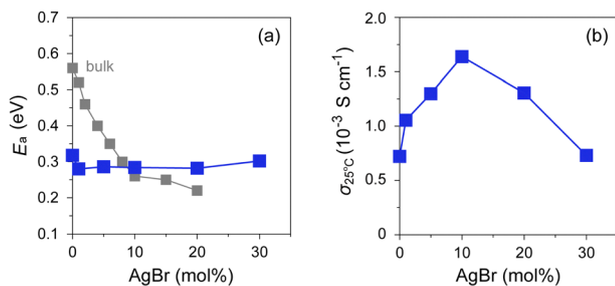


Fig. 7 Plots of (a) E_a and (b) $\sigma_{25^\circ\text{C}}$ against AgBr-doping ratio for AgI–AgBr/MPA composites represented by closed blue squares. Closed grey squares represent the values of bulk AgI–AgBr mixed crystals.^{49–51,78}

that the phase separation suppresses the discontinuous change in σ as observed for bulk AgI, which must be reflected in a very gradual increase in the ratio of α -AgI_{ss} phase on heating (Fig. 5(e) and (h)). The activation energy (E_a) values estimated by fitting the data up to 130 °C to the Arrhenius equation, $\sigma T = A \exp(-E_a/k_B T)$, where A is the pre-exponential factor and k_B is the Boltzmann constant, are shown in Fig. 7(a). The E_a values of the bulk AgI–AgBr mixed crystals reported by Shahi and Wagner are also shown for comparison.^{49–51,78} The E_a value of AgX is generally expressed as $E_a = H_f/2 + h_m$, where the migration energy of Frenkel defects (h_m) is smaller than the formation enthalpy of Frenkel defects (H_f).^{50,79} Thus the lower E_a value of **Br0** (0.32 eV) than that of bulk AgI (0.56 eV) can be attributed to the decreased H_f owing to the formation of AgI nanoparticles in the mesopores of MPA and the decreased h_m owing to the space charge layer formed at the interface with MPA. The E_a values of the AgI–AgBr/MPA composites remained relatively unchanged with the AgBr content, whereas the values of the bulk AgI–AgBr mixed crystals steadily decreased with increasing AgBr content (Fig. 7(a)). Apparently, the decrease in E_a in the bulk AgI–AgBr mixed crystals arose from the decrease in H_f with increasing AgBr content due to the lattice distortion induced by the partial substitution of I with Br in the AgI lattice. At present, the factor that keeps the E_a values nearly constant in the AgI–AgBr/MPA composites is unclear. However, the tendency seems to be associated with the fact that the 2-fold increase in $\sigma_{25^\circ\text{C}}$ with 10 mol% AgBr doping in the AgI–AgBr/MPA composite (see below) is significantly low compared with the increase in three orders of magnitude in the bulk AgI–AgBr mixed crystals. The σ value at 200 °C was the highest for the composite with AgBr = 10 mol% (*i.e.*, **Br10**; $5.7 \times 10^{-2} \text{ S cm}^{-1}$), above which the value exhibited a gradual decrease with increasing AgBr content ($3.4 \times 10^{-2} \text{ S cm}^{-1}$ for **Br30**) (Fig. S7, ESI[†]), primarily due to the incomplete transition to the highly ion-conducting α -phase.

The ion-conducting behaviour in the subsequent cooling process was considerably different from that in the heating process for each composite (Fig. 6). For the composites with AgBr \leq 10 mol%, the temperature at which the valley-like σ behaviour was observed (*ca.* 100 °C) corresponds approximately to the onset temperature of the exothermic DSC peaks on the high-temperature side (red arrows). Considering the fact that the structural transformation from α -AgI_{ss} to β/γ -AgI_{ss} with an

expanded lattice (*e.g.*, 64.0 \AA^3 per formula unit for α -AgI_{ss} at 110 °C vs. 67.8 \AA^3 per formula unit for β -AgI_{ss} at 30 °C in **Br10**) occurs at the core of the mesopores below the temperature, the flattening or upturn of σ observed in the temperature range may be attributed to the connection of the Ag⁺-ion conducting percolation pathways^{26,80–82} formed by the α -AgI_{ss} region owing to the expanded lattice on cooling.³¹ Similarly, the valley-like behaviour of σ was observed at approximately 100 °C for the composites with AgBr > 10 mol%, although there is no DSC peak in the temperature range. However, given that the variable-temperature PXRD measurements of **Br20** and **Br30** exhibited a nearly constant or increasing unit cell volume of the α -AgI_{ss} phase on cooling (Fig. S8, ESI[†]), the expanded lattice may be the primary cause of the non-Arrhenius-type conduction observed in the cooling process. We confirmed that the room-temperature PXRD pattern remained unchanged after the cyclic heating and cooling processes (Fig. S9, ESI[†]).

As shown in Fig. 7(b), the AgBr-doping ratio dependence of $\sigma_{25^\circ\text{C}}$ showed a volcano-type correlation with a maximum at AgBr = 10 mol% (*i.e.*, **Br10**). It is evident that the σ value ($1.6 \times 10^{-3} \text{ S cm}^{-1}$) is more than twice as high as that of the highest conducting AgI/MPA (*i.e.*, **Br0**; $7.2 \times 10^{-4} \text{ S cm}^{-1}$) reported in our previous paper.⁴⁷ An increase in $\sigma_{25^\circ\text{C}}$ in the low AgBr-doping range is possibly related to the increase in Ag⁺ migration caused by an increased concentration of Frenkel defects due to the lattice distortion. A decrease in $\sigma_{25^\circ\text{C}}$ in the high AgBr-doping range is caused by phase separation as evidenced by the PXRD measurements, which disturbs the Ag⁺-ion diffusion at the phase boundary in the composites (Fig. S10, ESI[†]).

Conclusions

In this study, our research interest is centred on the variable-temperature structural, calorimetric, and Ag⁺ ion-conducting properties of AgI/AgBr-loaded mesoporous alumina with various AgBr doping ratios. We found that the ionic conductivity of the AgI–AgBr/MPA composite with AgBr = 10 mol% exceeded $10^{-3} \text{ S cm}^{-1}$ at room temperature; the value, $1.6 \times 10^{-3} \text{ S cm}^{-1}$, is more than twice as high as that of the highest ion-conducting AgI/MPA found in our previous study ($7.2 \times 10^{-4} \text{ S cm}^{-1}$).⁴⁷ An increase in the AgBr content facilitates the formation of Frenkel pairs owing to lattice distortion, and the formed interstitial Ag⁺ ions lead to enhanced ionic conductivity. A further increase in the AgBr content causes the decrease in ionic conductivity due to the occurrence of phase separation. Solid-solution composite systems have the great advantage of using various combinations of homovalent anions and various selections of porous materials, which would allow for the rational design of highly ion-conducting composites.

Conflicts of interest

There are no conflicts to declare.



Acknowledgements

This work was supported by the ACCEL program (JPMJAC1501) of the Japan Science and Technology Agency (JST) and JSPS KAKENHI Grant Numbers JP20H02708 and JP20H05623. Variable-temperature PXRD measurements were conducted at the University of Tokyo, supported by “Advanced Research Infrastructure for Materials and Nanotechnology in Japan (ARIM)” of the Ministry of Education, Culture, Sports, Science and Technology (MEXT), Grant Number JPMXP1223UT0224.

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