Aalborg Universitet



## Structural Origins of the Enhancement in Ionic Conductivity of a Chalcogenide Compound by Adding Agl

Qiao, Wanchun; Qiao, Ang; Tao, Yunhang; Gu, Shaoxun; Yue, Yuanzheng; Tao, Haizheng

Published in: ChemElectroChem

DOI (link to publication from Publisher): 10.1002/celc.201902163

Publication date: 2020

Document Version Accepted author manuscript, peer reviewed version

Link to publication from Aalborg University

Citation for published version (APA): Qiao, W., Qiao, A., Tao, Y., Gu, S., Yue, Y., & Tao, H. (2020). Structural Origins of the Enhancement in Ionic Conductivity of a Chalcogenide Compound by Adding Agl. *ChemElectroChem*, 7(7), 1567–1572. https://doi.org/10.1002/celc.201902163

#### **General rights**

Copyright and moral rights for the publications made accessible in the public portal are retained by the authors and/or other copyright owners and it is a condition of accessing publications that users recognise and abide by the legal requirements associated with these rights.

- Users may download and print one copy of any publication from the public portal for the purpose of private study or research.
  You may not further distribute the material or use it for any profit-making activity or commercial gain
  You may freely distribute the URL identifying the publication in the public portal -

#### Take down policy

If you believe that this document breaches copyright please contact us at vbn@aub.aau.dk providing details, and we will remove access to the work immediately and investigate your claim.

# FUNDAMENTALS & APPLICATIONS

## ANALYSIS & CATALYSIS, BIO & NANO, ENERGY & MORE

# **Accepted Article**

Title: Structural origins of the enhancement in ionic conductivity of a chalcogenide compound by adding Agl

Authors: Wanchun Qiao, Ang Qiao, Yunhang Tao, Shaoxuan Gu, Yuanzheng Yue, and Haizheng Tao

This manuscript has been accepted after peer review and appears as an Accepted Article online prior to editing, proofing, and formal publication of the final Version of Record (VoR). This work is currently citable by using the Digital Object Identifier (DOI) given below. The VoR will be published online in Early View as soon as possible and may be different to this Accepted Article as a result of editing. Readers should obtain the VoR from the journal website shown below when it is published to ensure accuracy of information. The authors are responsible for the content of this Accepted Article.

To be cited as: ChemElectroChem 10.1002/celc.201902163

Link to VoR: http://dx.doi.org/10.1002/celc.201902163



WILEY-VCH

www.chemelectrochem.org

# Structural origins of the enhancement in ionic conductivity of a chalcogenide compound by adding Agl

Wanchun Qiao, <sup>[a]</sup> Ang Qiao, <sup>[a]</sup> Yunhang Tao, <sup>[b]</sup> Shaoxuan Gu, <sup>[a]</sup> Yuanzheng Yue, <sup>[a,c]</sup> Haizheng Tao \*<sup>[a]</sup>

**Abstract:** Through a chemo-mechanical milling process, we prepare a highly conductive  $(1.1 \times 10^{-3} \text{ S} \cdot \text{cm}^{-1})$  amorphous  $0.5\text{Agl} \cdot 0.5\text{Ag}_3\text{PS}_4$  conductor. Detailed structural characterizations indicate that the significantly higher ionic conductivity in the amorphous  $0.5\text{Agl} \cdot 0.5\text{Ag}_3\text{PS}_4$  compared to  $\text{Ag}_3\text{PS}_4$  can be ascribed to the formation of mixed polymeric anions {[PS4]ml<sub>h</sub>} around Ag<sup>+</sup> ions. Through heat-treatment at 370 °C for 20 minutes, the room temperature ionic conductivity of the  $0.5\text{Ag}_3\text{PS}_4$  conductor is further enhanced by about 4 times. This enhancement can be ascribed to the following two aspects: 1) the existence of residual amorphous phase with higher ionic conductivity; 2) the connection of the fast ionic conductive interfaces between the precipated Ag<sub>7</sub>PS<sub>6</sub> nano-crystals and the residual amorphous phase. This work reveals the key roles of both disorder and interface in improving the ionic conductivity of solid state electrolytes.

### Introduction

Compared to the liquid electrolytes currently used in commercial batteries, solid electrolytes have obtained extensive attention mainly owing to their highly applicable perspectives in stability (non-volatility),<sup>[1-4]</sup> safety (non-explosiveness)<sup>[5-9]</sup> and device fabrication (easy shaping, patterning and integration).<sup>[10-13]</sup> As an important class of solid-state electrolytes, Ag-based ionic conductors have undergone several breakthroughs in recent years in enhancing the room temperature ionic conductivity.<sup>[14-17]</sup> In addition, the prototypes of all-solid-state batteries based on the silver fast ionic conductor were also reported.<sup>[18-21]</sup> How ever, to fulfill the requirements of commercial applications, the performances of such devices still need to be improved.

As a distinguished solid state electrolyte, the hightemperature  $\alpha$ -phase AgI exhibits the super-ionic conductivity higher than 1 S·cm<sup>-1</sup>, which is comparable with the liquid ones. How ever, upon cooling to 147 °C, the  $\alpha$ -phase AgI will experience a structural transformation into the inferiorly conductive  $\beta$ - and  $\gamma$ -phase AgI, therefore restricting its uses. Various attempts have been performed to preventing the hightemperature  $\alpha$ -phase AgI from transforming into the low -

[a] W. C. Qiao, Prof. H. Z. Tao, Prof. S. X. Gu, Dr. A. Qiao. State Key Laboratory of Silicate Materials for Architectures, Wuhan University of Technology, Wuhan 430070, China E-mail: <u>thz@whut.edu.cn</u>
[b] Y. H. Tao

Wuhan Britain-China School, Wuhan 430034, China

[c] Prof. Y. Z. Yue

Department of Cherristry and Bioscience, Aalborg University, DK-9000 Aalborg, Denmark. E-mail: yy@bio.aau.dk phase.

For example, using a traditional melt-quenching method, at room temperature, the high-temperature  $\alpha$ -phase AgI can be kept in an AgI-Ag<sub>2</sub>O-B<sub>2</sub>O<sub>3</sub> glass matrix.<sup>[22]</sup> How ever, the high-temperature  $\alpha$ -phase AgI will vanish upon a heating cycle.

Through a chemo-mechanical milling process, an enhancement of three orders of magnitude in ionic conductivity Ag<sub>3</sub>PS<sub>4</sub> sample compared to the for the amorphous corresponding crystal phase was recently achieved by our group.<sup>[13]</sup> In this work, through the addition of Agl to Ag<sub>3</sub>PS<sub>4</sub>, we prepared the amorphous 0.5Agl-0.5Ag<sub>3</sub>PS<sub>4</sub> sample via the highenergy ball-milling method, and this sample exhibited the ionic conductivity of  $1.1 \times 10^{-3}$  S cm<sup>-1</sup>. By subjecting the amorphous 0.5Agl-0.5Ag<sub>3</sub>PS<sub>4</sub> to a certain degree of heat-treatment, a nanocrystal containing composite was obtained, which displayed an room temperature ionic conductivity of 3.9×10<sup>-3</sup> S·cm<sup>-1</sup>, i.e., approximately 4 times higher than that of the not heat-treated sample. By conducting structural characterizations, we observed and analyzed the atomic-scale structural evolutions from the crystalline to the amorphous 0.5Agl 0.5Ag<sub>3</sub>PS<sub>4</sub> phase and further to the nano-crystalline composite. This enabled us to reveal the structural origins of the enhancement in ionic conductivity induced by both amorphization and heat-treatment.

### Results

To determine its thermodynamic characteristics, calorimetric curves of the as-prepared 0.5Ag $I \cdot 0.5$ Ag $_3$ PS $_4$  pow der are show n in Fig.1. In the first up-scan curve, besides three clear exothermic peaks located at 137 °C, 197 °C, and 370 °C respectively, we can also see four smaller exothermic responses at 103, 260, 302 and 333 °C respectively.

According to the calorimetric responses of the as-prepared specimen, we selected the highest exothermal peak temperature of 370 °C as the annealing temperature. Utilizing an impedance analyzer, the comparison of complex impedance curves for both the as-synthesized sample and the isothermally heat-treated one are made in Fig.2. All curves display one semicircle in the high frequency region and a straight line in the low frequency region. The semicircle reflects the ionic conductivity  $\sigma$  (S·cm<sup>-1</sup>) of the bulky materials, while the straight line is related to the characteristics of interface betw een the specimen and eletrode.

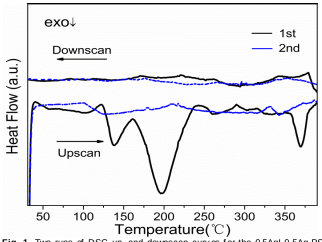


Fig. 1. Two runs of DSC up- and downscan curves for the  $0.5 Agl \cdot 0.5 Ag_3 PS_4$  powder prepared via milling at 800 rpm for 10 hrs. Up-and downscan rates are 10 °C min  $^{\circ}.$ 

The electrical resistances of the sample at various temperatures can be respectively found by reading the difference of the two intercepts of the semicircle on Z axis, i.e., by reading the diameter of the semicircle.  $\sigma$  can be obtained from the equation:

$$\sigma = \frac{L}{RS} \tag{1}$$

where *L* (cm) is the thickness of the plate, *R* ( $\Omega$ ) is the sample resistance, and *S* (cm<sup>2</sup>) stands for the area of the electrodes. From Fig. 2 and equation (1), the room temperature ionic conductivity of the as-synthesized  $0.5AgI \cdot 0.5Ag_3PS_4$  powder was enhanced by about 4 times through the isothermal treatment at 370 °C for 20 min, i.e., from  $1.1 \times 10^3$  to  $3.9 \times 10^3$  S·cm<sup>-1</sup>. This enhancement is associated with the drop of the activation energy (*E*<sub>a</sub>) from 0.295 eV to 0.244 eV as a result of isothermal heat-treatment. As shown in Fig.2, *E*<sub>a</sub> can be calculated by fitting the obtained  $\sigma$  (Fig.3) to the Arrehnian equation, <sup>[23,24]</sup>:

$$T = Aexp(-\frac{E_a}{kT}) \tag{2}$$

where  $\sigma$  is the total conductivity of the solid conductor at the temperature *T*, *k* is the Boltzmann constant, and *A* is the pre-exponential factor.



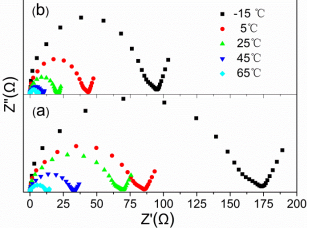


Fig. 2. Temperature dependence of the complex impedance for (a) the assynthesized amrophous  $0.5Agl\cdot0.5Ag_3PS_4$  powder and (b) the heat-treated one. The heat-treatment procedure is as follows: the as-synthesized powder was heated up to 370 °C at 5 °C min^1, then held at 370 °C for 20 min, finally cooled down to room temperature by turning off the furnace.

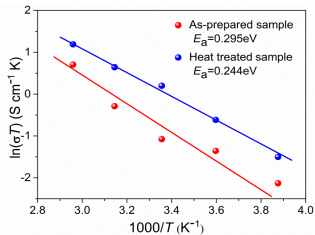


Fig. 3. Arrhenius plots of the ionic conductivity for both the as-synthesized amorphous 0.5AgI·0.5Ag $_3$ PS $_4$  powder and the sample heat-treated at 370 °C for 20 min.

To investigate the atomic-scale origin of these calorimetric responses upon heating and the mechanism of the enhancement of the ionic conductivity, several structural characterization techniques were utilized. XRD patterns were obtained for both the as-prepared 0.5Agl-0.5Ag<sub>3</sub>PS<sub>4</sub> powder and those treated at 150 °C, 250 °C and 370 °C, respectively, for 20 min. In addition, Raman and <sup>31</sup>P solid NMR spectra of the as-prepared 0.5Agl-0.5Ag<sub>3</sub>PS<sub>4</sub> powder and the heat-treated one at 370 °C for 20 min were shown in Figs. 5 and 6 to probe the structural difference of the two samples. Finally, to identify the structural difference betw een the as-prepared 0.5Agl-0.5Ag<sub>3</sub>PS<sub>4</sub> pow der and the previously reported amorphous Ag<sub>3</sub>PS<sub>4</sub> pow der<sup>[13]</sup>, Ag K-edge XANES and EXAFS spectra were also obtained as seen in Fig.7.

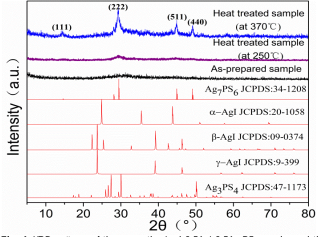


Fig. 4. XRD patterns of the as-synthesized 0.5Agl-0.5Ag<sub>3</sub>PS<sub>4</sub> powder and the heat-treated ones. The heat-treated protocol is as follows: the as-synthesized powder was heated at 5 °C min<sup>-1</sup> to a temperature (250 °C or 370 °C), then holding at this temperature for 20 min, finally cooling to room temperature through turning off the power.

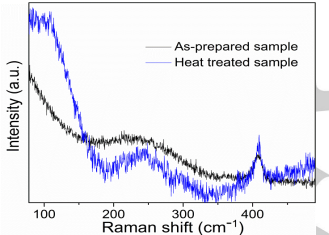
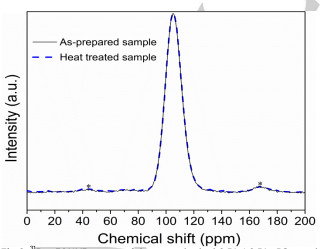
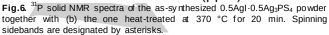


Fig.5. Raman spectra of the as-synthesized  $0.5Ag10.5Ag_3PS_4$  powder (black) together with the one heat-treated at 370 °C for 20 min (blue).





### Discussion

Calorimetric responses

In this work, we observed several fascinating phenomena in the DSC curves of the as-prepared  $0.5AgI\cdot0.5Ag_3PS_4$  sample (Fig. 1), in strong contrast to the amorphous  $Ag_3PS_4$  reported in our previous w ork.<sup>[25]</sup>

First, as confirmed by the XRD patterns of the heat-treated ones, the 0.5Ag10.5Ag3PS4 sample still remains amorphous after heat-treatment at 250 °C for 20 min. Therefore, two distint exothermic peaks occur at 137 °C and 197 °C respectively during the first upscan (Fig. 1), which could be due to the two the stages of relaxation of distorted a morphous 0.5Agl-0.5Ag<sub>3</sub>PS<sub>4</sub>. According to the structural characterizations, the ionic bonding occurs in the 0.5Agl-0.5Ag<sub>3</sub>PS<sub>4</sub> sample owing to the presence of Ag<sup>+</sup>, I, and  $[PS_4]^{3-}$  ions. The exothermic peaks at 137 °C could be attributed to the relaxation of the distorted Ag{[PS4]mln} polyhedral units, whereas the peak at 197 °C could be assigned to the relaxation of the distorted [PS4] tetrahedral units. These assignments were also reported for the as-prepared amorphous Ag<sub>3</sub>PS<sub>4</sub>.<sup>[25]</sup>

econd, according to the XRD pattern of the sample heattreated at 370 °C for 20 min (Fig. 4), the exothermic peak at 370 °C (Fig. 1) should be assigned to the formation of the single  $Ag_7PS_6$  crystal, in contrast to the precipitated crystalline  $Ag_3PS_4$ phase in the milling-induced amorphous  $Ag_3PS_4$  powder. In addition, the crystallization peak for the as-prepared amorphous 0.5Ag1.0.5Ag\_3PS\_4 powder is much smaller than the relaxation peaks (Fig. 1), and this scenario is different from that of the amorphous  $Ag_3PS_4$ , where the crystallization peak is stronger than the relaxation peak. This difference could be due to the enhanced structural disorder and heterogeneity upon adding Ag1.

Finally, similar to the case for the Ag\_3PS\_4 amorphous sample,  $^{[25]}$  a glass transition cannot be detected prior to the crystallization peak for the amorphous 0.5Agl-0.5Ag\_3PS\_4 pow der.

Atomic-scale structural origin of the enhancement in ionic conductivity

Compared to the as-prepared amorphous Ag<sub>3</sub>PS<sub>4</sub> pow der,<sup>[13]</sup> the addition of Ag1 to Ag<sub>3</sub>PS<sub>4</sub> following the stoichiometric ratio of 0.5Ag1·0.5Ag<sub>3</sub>PS<sub>4</sub> leads to a large jump of the room temperature (298 K) ionic conductivity, i.e., an increase from  $8.5 \times 10^{-4}$  S·cm<sup>-1</sup> to  $1.1 \times 10^{-3}$  S·cm<sup>-1</sup> (Fig. 2). This enhancement could be mainly ascribed to the drop in the activation energy (*E*<sub>a</sub>) from 0.320 eV for the as-prepared Ag<sub>3</sub>PS<sub>4</sub> amorphous pow der to 0.295 eV for the 0.5Ag1·0.5Ag<sub>3</sub>PS<sub>4</sub> amorphous pow der.

Now the central question arises: what is the atomistic origin of the enhancement in ionic conductivity by adding AgI? To answer this question, the comparison in the Ag K-edge XANES and EXAFS spectra between the as-prepared amorphous Ag<sub>3</sub>PS<sub>4</sub> and 0.5Ag10.5Ag3PS<sub>4</sub> powders are made in Fig. 7. It turns out that only a small difference between the two samples

can be seen in the Ag K-edge XANES spectra, implying the changes in the coordinating surroundings around Ag<sup>+</sup> ions. Furter information about the nearest neighbor coordinating species can be deduced from the peak located at 2.7 Å near the Ag-S peak in the Ag K-edge EXAFS spectra (Fig. 7b), implying a formation of Ag-I bonds upon adding AgI. This suggests that the nearest neighbor surroundings of Ag<sup>+</sup> ions from [PS<sub>4</sub>] single anions transform into the {[PS<sub>4</sub>]<sub>m</sub> h<sub>i</sub>} mixed polymeric anions. Based on the model presented elsew here,<sup>[26,27]</sup>

enhancement in ionic conduction (originating from mixed iodine and oxygen environments) upon adding AgI to an oxide glass system could be ascribed to the increase in the I-Ag distance, since oxygens draw Ag<sup>+</sup> ions, and thereby increase the average I-Ag distance. In this work, the rise in ionic conductivity could be also ascribed to the extension of the pathways for ionic transport due to the formation of  $\{[PS_4]_m l_n\}$  mixed polymeric anions. The asymmetry of the nearest neighbor surroundings of Ag<sup>+</sup> ions facilitates the migration of Ag<sup>+</sup> ions.

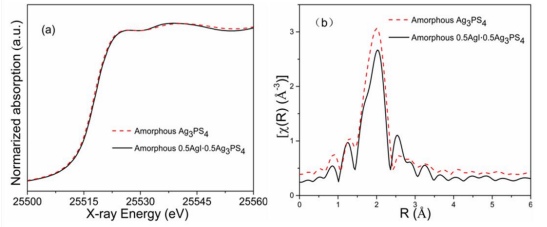


Fig.7. XANES (a) and EXAFS spectra in R-space (b) for Ag K-edge of the as-synthesized amorphous Ag<sub>3</sub>PS<sub>4</sub> and 0.5Ag1-0.5Ag<sub>3</sub>PS<sub>4</sub> powders.

The further enhancement (about 4 times) in room temperature ionic conductivity upon heat-treatment at 370 °C for 20 min for the amorphous  $0.5AgI \cdot 0.5Ag_3PS_4$  powder could be attributed to the following aspects. First, different from the precipitated Ag<sub>3</sub>PS<sub>4</sub> crystal phase in the as-synthesized Ag<sub>3</sub>PS<sub>4</sub> amorphous powder, Ag<sub>7</sub>PS<sub>6</sub> crystal phase (JCPDS 41-0866) appears in the heat-treated amorphous 0.5AgI-0.5Ag<sub>3</sub>PS<sub>4</sub> pow der as confirmed by the XRD pattern shown in Fig. 4. It has been reported that the ionic conductivity of Ag7PS6 crystal is  $1.5 \times 10^{-6} \ {\rm S} \cdot {\rm cm}^{-1},^{\rm [28]}$  which is much low er than those of both the amorphous  $0.5AgI \cdot 0.5Ag_3PS_4$  pow der  $(1.1 \times 10^{-3} \text{ S} \cdot \text{cm}^{-1})$  and the heat-treated amorphous  $0.5AgI \cdot 0.5Ag_3PS_4$  pow der  $(3.9 \times 10^{-3})$ S·cm<sup>-1</sup>). According to the stoichiometry of the 0.5Agl-0.5Ag<sub>3</sub>PS<sub>4</sub> amorphous powder, given the complete precipitation of the Ag<sub>7</sub>PS<sub>6</sub> crystal phase, the composition of the residual amorphous phase should be  $AgPS_2 I_2$ , and the latter exhibits a room temperature ionic conductivity of about 1.1~3.3×10<sup>-3</sup> S·cm<sup>-</sup> <sup>1</sup>.<sup>[29,30]</sup> This value is also lower than that of the heat-treated amorphous  $0.5Agl \cdot 0.5Ag_3PS_4$  pow der  $(3.9 \times 10^{-3} \text{ S} \cdot \text{cm}^{-1})$ . Thus, another question arises: What else causes the further enhancement of the room temperature ionic conductivity for the heat-treated amorphous 0.5AgI 0.5Ag<sub>3</sub>PS<sub>4</sub> pow der?

Based on the investigations on the Li<sup>+</sup> solid-state electrolyte,<sup>[31-33]</sup> the enhancement in ionic conductivity might be realized through the formation of a certain crystal phase with a low er ionic conductivity in the amorphous sample. This

anomalous behavior could be clarified by considering the role of interface between the crystalline and the amorphous phases in influencing the ionic conductivity of the composite, e.g.,through the formation of percolative conductive pathways, the aggregation of transferring ions, and space charges. According to a previous study,<sup>[34]</sup> two possibilities might explain the anomalous phenomenon. On one hand, the severe lattice deformations could lead to the enhanced ionic transportation in the interfacial region. On the other hand, the rise of transferring ions in quantity in the interfacial regions could also contribute to the enhancement in ionic conductivity. Therefore, when the size of the crystallites is reduced to nano-scale, the interconnected interfacial regions benefit the formation of percolative pathways for ionic transportation in the composite. Such a strong enhancement in ionic conductivity caused by interfacial effects has been observed in many nano-crystals containing composites.[35,36]

In addition, due to the higher binding energy of P-S bonds, the polymerized anions [PS<sub>4</sub>] tetrahedra are preserved in the asprepared amorphous  $0.5AgI\cdot0.5Ag_3PS_4$  powder and the heat-treated one, deduced from the similar characteristic Raman peaks for [PS<sub>4</sub>] tetrahedra I<sup>[13,37-39]</sup> (Fig. 5) and the similar <sup>31</sup>P NMR spectra (Fig. 6).

According to the above-mentioned explanations, the mechanisms of the enhancement in ionic conductivity are schematically illustrated in Fig. 8.

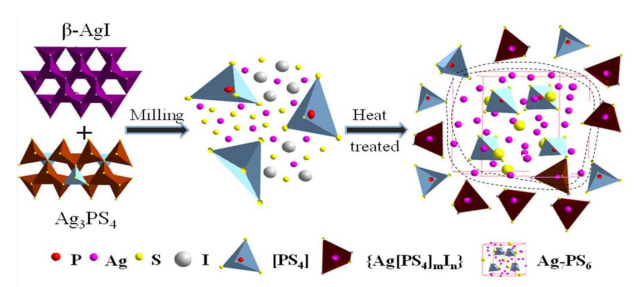


Fig.8. Schematic diagrams of the structural evolution from the initial crystals, to amorphous materials upon milling, and finally to nano-crystal composites obtained by heat-treatment at 370 °C for 20 min.

### Conclusions

We synthesized a new kind of the Ag-based fast ionic conductor, i.e., the amorphous 0.5Agl-0.5Ag<sub>3</sub>PS<sub>4</sub>, via a chemomechanical ball milling. Compared to the Ag<sub>3</sub>PS<sub>4</sub> amorphous powder, the addition of equal molar AgI to Ag<sub>3</sub>PS<sub>4</sub> strongly enhances the room temperature ionic conductivity, i.e., up to  $1.1 \times 10^{-3}$  S cm<sup>-1</sup> for the amorphous 0.5Ag $1 \cdot 0.5$ Ag $_3$ PS $_4$  pow der. Further enhancement of about 4 times was realized upon the heat-treatment of the amorphous 0.5AgI-0.5Ag<sub>3</sub>PS<sub>4</sub> powder at 370 °C for 20 min. The detailed structural characterizations allow ed us to reveal the atomistic origin of the ionic conductivity enhancement upon adding AgI to the amorphous Ag<sub>3</sub>PS<sub>4</sub>, i.e., the formation of  $\{[\mathsf{PS}_4]_m\,\mathsf{I}_h\}$  mixed polymeric anions around  $\mathsf{Ag}^+$ ions in the amorphous 0.5AgI-0.5Ag<sub>3</sub>PS<sub>4</sub> powder plays a crucial role. We found the precipitation of Ag<sub>7</sub>PS<sub>6</sub> crystal phase in the heat-treated amorphous  $0.5AgI \cdot 0.5Ag_3PS_4$  pow der. Finally, based on the detailed structural analysis, we established the mechanisms of the enhancement in ionic conductivity for both the amorphous 0.5Agl-0.5Ag<sub>3</sub>PS<sub>4</sub> powder and the heat-treated one. This work will be useful for the design and production of the silver-based super-ionic electrolytes.

### **Experimental Section**

#### Sample preparation

The amorphous Ag<sub>3</sub>PS<sub>4</sub> and 0.5Ag<sub>1</sub>·0.5Ag<sub>3</sub>PS<sub>4</sub> samples were prepared through a chemo-mechanical milling equipment (Pulverisette-7 premium line; Fritsch, Idar-Oberstein, Germany) at 800 rpm for 10 hours. Reagent-grade Ag<sub>1</sub>, Ag<sub>2</sub>S and P<sub>2</sub>S<sub>5</sub> (all from Aladdin, Shanghai, China) crystalline pow ders were used as starting materials. Then these pow ders were loaded into a 20ml ZrO<sub>2</sub> pot with  $\phi$  3 mm ZrO<sub>2</sub> balls according to the ~1:15 weight ratio between the sample and balls. The detailed procedure was reported elsew here.  $^{[13, \ 40, \ 41]}$ 

#### Characterizations

To detect the calorimetric responses of the as-prepared  $0.5AgI\cdot 0.5Ag_3PS_4$  powder and the appropriate heat-treated parameters, two rund of DSC up- and downscan at 10 °C min<sup>-1</sup> were conducted by utilizing a differential scanning calorimeter (DSC) (Netzsch STA449 F1, Netzsch, Selb, Germany).

amorphous nature of The the as-synthesized 0.5Agl-0.5Ag<sub>3</sub>PS<sub>4</sub> samples, as well as the precipitated crystal phases in the heat-treated samples, were determined by an Xray diffractometer (Rigaku, RU-200B, Tokyo, Japan) with Cu Kα radiation at a voltage of 40 kV and a current of 40 mA, and the X-ray diffraction (XRD) patterns were recorded in the 20 range of 5-80°. In addition, structure investigation was conducted using a HORIBA LabRAM HR Evolution Raman microscope, in which a 633 nm laser with a typical power lower than 5 mW was used to avoid the laser-induced damage. Furthermore, the nearest neighboring environments of Ag in both the as-synthesized Ag<sub>3</sub>PS<sub>4</sub> and the 0.5Ag1 0.5Ag<sub>3</sub>PS<sub>4</sub> sample were characterized by the Ag K-edge X-ray absorption near edge structure (XANES) and the extended X-ray absorption fine structure (EXAFS) spectra. The details about the measurement can be found elsew here.<sup>[13,42]</sup> Finally, the local coordination surroundings of P for both the as-synthesized and the heat-treated 0.5AgI 0.5Ag<sub>3</sub>PS<sub>4</sub> samples were probed by the <sup>31</sup>P MAS NMR experiments. These measurements were conducted at 120 MHz on a Bruker Avance III 400 spectrometer furnished with a 4 mm probe. The <sup>31</sup>P chemical shifts are referenced to 85% H<sub>3</sub>PO<sub>4</sub> aqueous solution ( $\delta = 0$  ppm).

To characterize the electrochemical performances, the assynthesized pow ders were dry-pressed vertically into plates with a diameter of 1.28 cm and with a height of about 0.1 cm. Through magnetron sputtering technique, the two sides of the obtained plates were coated with a layer of platinum films. The AC impedance spectra of both the as-synthesized and the heat-treated samples were obtained using an impedance analyzer (Solartron, 1260A) in the frequency range from 0.1 Hz to 10 MHz using an amplitude voltage of 50 mV across the sample. These measurements were carried out in the temperature range of -15 °C to 65 °C under an atmospheric condition. The ionic conductive performance was analyzed by using the Z-View impedance software.

### Acknowledgements

This work was financially supported by NSFC (Nos.51772223, 51372180). We are grateful to Xiujian Zhao and Xianghua Zhang (Wuhan University of Technology) for valuable discussions. W.C. Qiao and A. Qiao contributed equally to this work. In addition, we also acknowledge the provision of synchrotron access to Beamline BL14W1 at the Shanghai Synchrotron Radiation Facility.

Keywords: Solid electrolyte; Ionic conductivity; EXAFS; Highenergy ball-milling

[1]. M. Armand, J. M. Tarascon, Nature. 2008, 451, 652-657.

- [2]. J. M. Tarascon, M. Armand, Nature. 2001, 414, 359-367.
- [3]. W. Hou, X. Guo, X. Shen, K. Amine, H. Yu, J. Lu, *Nano Energy*. **2018**, *52*, 279–291.

[4]. K. H. Park, D. Y. Oh, Y. E. Choi, Y. J. Nam, L. Han, J. Kim, H. Xin, F. Lin, S. M. Oh, Y. S. Jung, *Adv. Mater.* **2016**, *28*, 1874–1883.

[5]. J. Janek, W. G. Zeier, Nat. Energy. 2016, 1, 1-4.

[6]. J. B. Goodenough, K. Park, J. Am Chem Soc. 2013, 135, 1167-1176.

[7]. F. Strauss, T. Bartsch, L. Biasi, A. Y. Kim, J. Janek, P. Hartmann, T. Brezesinski, ACS Energy Lett. 2018, 3, 992–996.

[8]. T. Asano, A. Sakai, S. Ouchi, M. Sakaida, A. Miyazaki, S. Hasegawa, Adv. Mater. 2018, 1803075, 1–7.

[9]. K. Takada, Acta Mater. 2013, 61, 759-770.

[10]. M. Park, X. Zhang, M. Chung, G. B. Less, A. Marie, J. Power Sources. 2010, 195, 7904–7929.

[11]. M. Wakihara, Mater. Sci. Eng.. 2001, R33, 109-134.

[12]. K. Zaghib, P. Charest, A. Guerfi, J. Shim, M. Perrier, K. Striebel, J. Power Sources. 2004, 134, 124–129.

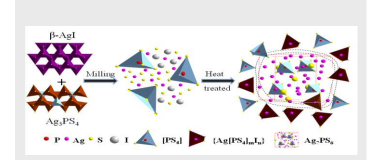
- [13]. A. Qiao, H. Tao, Y. Yue, J. Non-Cryst. Solids. 2019, 521, 119476.
- [14]. K. Terabe, T. Hasegawa, T. Nakayama, M. Aono, *Nature*. **2005**, *4*33, 47–50.
- [15]. H. Zhang, T. Tsuchiya, C. Liang, K. Terabe, *Nano Lett.* **2015**, *15*, 5161–5167.
- [16]. C. Lin, E. Zhu, J. Wang, X. Zhao, F. Chen, S. Dai, J. Phys. Chem C. 2018, 122, 1486–1491.
- [17]. R. Makiura, T. Yonemura, T. Yamada, M. Yamauchi, R. Ikeda, H. Kitagawa, K. Kato, M. Takata, *Nat. Mater.* **2009**, *8*, 476–480.
- [18]. P. Boolchand, W. J. Bresser, Nature. 2001, 410, 1070-1073.
- [19]. R. Murugaraj, G. Govindaraj, S. Ramasamy, *J. Power Sources.* 2002, 112, 184–190.
- [20]. B. K. Heep, K. S. Weldert, Y. Krysiak, T. W. Day, W. G. Zeier, U. Kolb, G. J. Snyder, W. Tremel, *Chem Mater.* 2017, *29*, 4833–4839.
- [21]. C. Lin, C. Rüssel, S. Dai, Prog. Mater. Sci. 2018, 93, 1-44.
- [22]. M. Tatsumisago, Y. Shinkuma, T. Minami, Nature. 1991, 354, 217-218.
- [23]. Z. Zhang, J. H. Kennedy, J. Electrochem Soc. 1993, 140, 2384–2390.
- [24]. W. Yao, S.W. Martin, Solid State Ionics. 2008, 178, 1777–1784.
- [25]. A. Qiao, H. Tao, Y. Yue, J. Am Ceram Soc. 2017, 100, 968–974.
- [26]. P. Mustarelli, C. Tomasi, A. Magistris, J. Phys. Chem B. 2005, 109, 17417–17421.
- [27]. S. Adams, J. Swenson, Phys. Rev. B. 2000, 63, 1-11.
- [28]. Z. M. Zhang, J. H. Kenned, J. Electrochem Soc. 1993, 140, 2384-2390.
- [29]. H. Tao, S. Yan, P. Wang, A. Qiao, C.N. Patent **2016**, 106057276A.
- [30]. H. Tao, P. Wang, A. Qiao, C. Zhong, Y. Yue, C.N. Patent 2015, 104752756A.
- [31]. F. Su, Solid State Ionics. 1982, 7, 37-41.
- [32]. L. Q. Chen, L. Z. Wang, G. C. Che, G. Wang, Z. R. Li, *Solid State Ionics*. **1984**, *14*, 149-152.
- [33]. U. M. Gundusharma, E. A. Secco, Solid State Ionics. 1990, 44, 47–50.
- [34]. S. Adams, K. Hariharan, J. Maier, Solid State Ionics. 1995, 75, 193-201.
   [35]. P. Knauth, J. M. Debierre, G. Albinet, Solid State Ionics. 1999, 121, 101–
- 106.
- [36]. G. Albinet, J. M. Debierre, P. Knauth, C. Lambert, L. Raymond, *Eur. Phys. J. B.* **2001**, *22*, 421–427.

[37]. I. P. Studenyak, V. O. Stefanovich, M. KranjEecb, D. I. Desnica, Y. M. Azhnyukd, G. S. Kovacsa, V. V. Panko, *Solid State Ionics*. **2003**, 95, 221–225.

- [38], S. Jörgens, A. Mewis, Solid State Sci. 2007, 9, 213–217.
   [39], H. Andrae, R. Blachnik, J. Alloy. Comp. 1992, 189, 209–215.
- [40]. Y. Katayama, S. Hama, T. Yanagihara, M. Hayashi, U.S. Patent **2011**, 20110049745A1.
- [41]. S. Hama, M. Hayashi, U.S. Patent 2013, 8, 556, 179.
- [42]. M. Newville, J. Synchrotron Radiat. 1998, 8, 322-324.

## WILEY-VCH

## ARTICLE



W.	С.	Qiao,	H. Z.	Tao*

Page No. – Page No.

Structural origins of the enhancement

in ionic conductivity of a

chalcogenide compound by adding

Agl

The back cover picture reveals the mechanisms of the enhancement in ionic conductivity for the present 0.5AgI-0.5Ag<sub>3</sub>PS<sub>4</sub> amorphous pow der together with the heat-treated one are schematically given.