

CORRECTION

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rsc.li/materials-a**Correction: Designing a descriptor for the computational screening of argyrodite-based solid-state superionic conductors: uniformity of ion-cage size**Byeongsun Jun ^a and Sang Uck Lee ^{*ab}Correction for 'Designing a descriptor for the computational screening of argyrodite-based solid-state superionic conductors: uniformity of ion-cage size' by Byeongsun Jun *et al.*, *J. Mater. Chem. A*, 2022, 10, 7888–7895, DOI: [10.1039/d1ta10964b](https://doi.org/10.1039/d1ta10964b).

In the original article, the bottom part of Fig. 2 is missing. The corrected Fig. 2 is shown below:

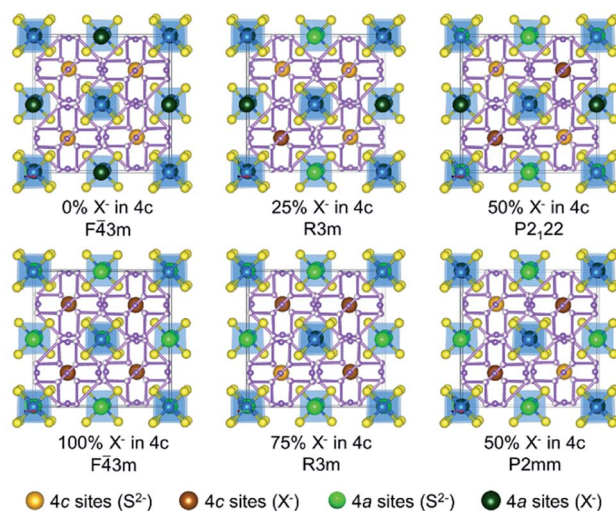


Fig. 2 Configurations of single anion (S^{2-} and X^{-}) distribution at 4a and 4c sites of Li_6PS_5X structure. Yellow and blue balls make up the PS_4^{3-} tetrahedron. Orange and brown (green and dark green) balls indicate S^{2-} and X^{-} single anions at 4c (4a) sites, respectively.

The Royal Society of Chemistry apologises for these errors and any consequent inconvenience to authors and readers.

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