

Cite this: *Chem. Sci.*, 2025, 16, 16751

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# Multinuclear $\text{Cu}_n\text{S}_m$ clusters encapsulated by aromatic micelles as aqueous red-to-NIR phosphorescent ink

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Sulfur-bridged copper clusters are important biocomponents, yet their artificial analogues have rarely been studied *in solution* due to poor solubility and stability. Here we report the preparation of red-to-near-infrared (NIR) phosphorescent solutions from multinuclear  $\text{Cu}_n\text{S}_m$  clusters upon encapsulation by aromatic micelles in water. For instance, whereas a 2-mercapto-6-methylpyridine-based  $\text{Cu}_6\text{S}_6$  cluster shows no solution-state emission owing to its insolubility in common solvents, the encapsulated cluster emits strong red-to-NIR phosphorescence ( $\Phi = 34\%$ ;  $\lambda = 550\text{--}850\text{ nm}$ ) with high stability, even under aerobic and ambient conditions. Similar host-guest complexes are also obtained from analogous  $\text{Cu}_6\text{S}_6$  clusters, displaying substituent-dependent red-to-NIR emission in water. The present method is applicable to larger  $\text{Cu}_{12}\text{S}_6$  and smaller  $\text{Cu}_4\text{S}_4$  clusters to generate aqueous, red, yellow, and green emissive solutions. Notably, the resultant host-guest solution can be used as aqueous colorless ink, for potential security applications, which exhibits strong emission when painted on paper in the red-to-NIR region upon UV-light irradiation.

Received 6th June 2025  
Accepted 1st August 2025DOI: 10.1039/d5sc04122h  
rsc.li/chemical-science

## Introduction

Multinuclear metal clusters embedding heteroatoms exhibit varied structures with intriguing physicochemical properties, which cannot be found in related clusters without heteroatoms as well as mononuclear metal complexes.<sup>1</sup> For instance, sulfur-bridged metal clusters, composed of  $\text{Fe}_n\text{S}_m$  and  $\text{Cu}_n\text{S}_m$  cores (e.g.,  $n = 2\text{--}8$  and  $m = 1\text{--}8$ ),<sup>2</sup> have been known as key biocomponents with unique redox ability, protected in protein cavities (Fig. 1a). Synthetic multinuclear  $\text{Cu}_n\text{S}_m$  clusters have also attracted interest, due to their characteristic photophysical properties and reactivities with tunability.<sup>1,3</sup> However, except for biological systems, the majority of such artificial  $\text{Cu}_n\text{S}_m$  clusters are insoluble in water-organic solvent mixtures as well as water, which interferes with their green chemical processes and biomedical applications.<sup>4</sup> The intrinsic instability of the Cu-S bonds in solution also restricts their usability. To overcome these drawbacks, water-soluble multinuclear clusters have been generally prepared by utilizing hydrophilic thiol-based ligands attached covalently (Fig. 1b)<sup>5</sup> and by encapsulation in alkyl-based micelles (Fig. 1c).<sup>6</sup> Since these usual methods are inapplicable to  $\text{Cu}_n\text{S}_m$  clusters, owing to the facile oxidation/reduction of the Cu(I) moieties,<sup>4b,10c</sup> their water-solubilization and resultant solution-state strong emission remain elusive so far. To expand  $\text{Cu}_n\text{S}_m$ -based cluster chemistry in water by a new

method, we herein report (i) the facile preparation of aqueous solutions of  $\text{Cu}_6\text{S}_6$  clusters upon encapsulation by aromatic micelles (Fig. 1d). (ii) The obtained host-guest solutions display strong red-to-near-infrared (NIR) phosphorescence (up to  $\Phi = 34\%$ ) with high stability under ambient conditions. (iii) Similarly, various host-guest complexes are prepared using analogous  $\text{Cu}_6\text{S}_6$  clusters, showing wide red-to-NIR emission

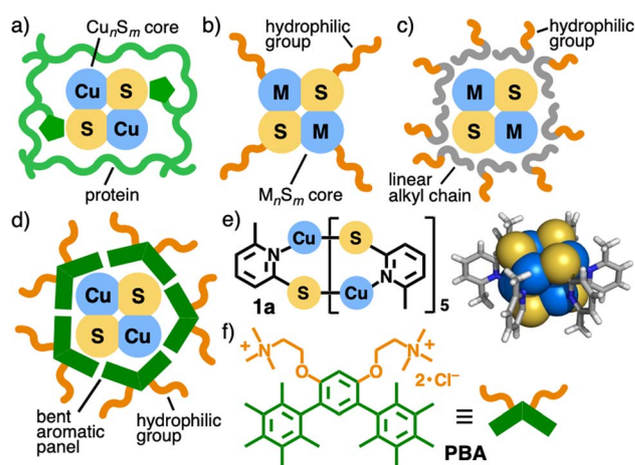


Fig. 1 Preparation of aqueous  $\text{Cu}_n\text{S}_m$  or  $\text{M}_n\text{S}_m$  cluster-based solutions, using (a) a protein cavity, (b) hydrophilic thiol ligands and (c) an alkyl micelle (except for  $\text{M} = \text{Cu}$ ), and (d) an aromatic micelle (this work). (e)  $\text{Cu}_6\text{S}_6$  cluster **1a** and its crystal structure.<sup>14,15</sup> (f) Bent aromatic amphiphile PBA.

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depending on the substituents. (iv) This method enables other multicopper clusters, with a  $\text{Cu}_{12}\text{S}_6$  and  $\text{Cu}_4\text{I}_4$  core, to generate red, yellow, and green emissive complexes in water. Furthermore, (v) the obtained host-guest solution can be used as red-to-NIR phosphorescent ink on paper without visible color under room light, for potential security applications.

Unlike multicopper clusters without heteroatoms,  $\text{Cu}_n\text{S}_m$  clusters show strong phosphorescent emission *in the solid state*, whereas this feature is lost to a large extent *in solution*.<sup>14,4,8</sup> Mercaptopyrindine-based  $\text{Cu}_6\text{S}_6$  clusters (e.g., Fig. 1e),<sup>9</sup> as a representative example, are poorly soluble in common organic solvents as well as insoluble in water, without laborious functionalization. Larger  $\text{Cu}_{12}\text{S}_6$  clusters with bisphosphine ligands<sup>10</sup> are soluble in organic solvents yet suffer from thermal instability and air-sensitivity. For the efficient encapsulation of such multicopper clusters with dimensions of 1–2 nm, here we employed bent aromatic amphiphile **PBA**, bearing two pentamethylbenzene panels and two trimethylammonium groups (Fig. 1f).<sup>11,12</sup> Its spontaneous and quantitative assembly generates aromatic micelle  $(\text{PBA})_n$  ( $n \approx 5$ ) in water at room temperature. Thanks to the hydrophobic effect and multiple  $\text{CH}-\pi$  interactions in the adaptable cavity, the aromatic micelle displays wide-ranging host ability toward planar and large aromatic compounds as well as metal-complexes with adequate stability,<sup>11,13</sup> unlike conventional alkyl micelles. On the other hand, like other host compounds reported previously,<sup>7</sup> its encapsulation ability toward sterically demanding, multinuclear  $\text{M}_n$  clusters ( $n > 3$ ) with low solution-state stability has not been revealed to date.

## Results and discussion

### Encapsulation of multinuclear $\text{Cu}_6\text{S}_6$ clusters by aromatic micelles

Bent aromatic amphiphile **PBA** displayed superior encapsulation ability toward 2-mercapto-6-methylpyridine-based  $\text{Cu}_6\text{S}_6$  cluster **1a** in water, as compared with typical alkyl amphiphiles. Colorless solid **PBA** (0.7 mg, 1.0  $\mu\text{mol}$ ) and pale yellow solid **1a** (0.6 mg, 0.5  $\mu\text{mol}$ ) were ground with a mortar and pestle for 1 min.<sup>14</sup> Water addition (1.0 mL) to the mixture, centrifugation, and filtration gave rise to a clear colorless solution including  $(\text{PBA})_n \cdot (\text{1a})_m$  in a nearly quantitative manner based on **PBA** (Fig. 2a). The UV-visible spectrum showed new shoulder bands at 300–420 nm, besides a prominent band for **PBA** (250–300 nm), which indicates the successful water-solubilization of  $(\text{1a})_m$  upon encapsulation (Fig. 2b). The absorption bands of  $(\text{1a})_m$  within  $(\text{PBA})_n$  were significantly blue-shifted ( $\Delta\lambda \approx 130$  nm), as compared to that of solid **1a**, suggesting the suppression of cluster aggregation and the isolation of a single cluster. The obtained, aqueous host-guest solution is stable enough under ambient conditions (*i.e.*, room temperature and light) for >1 week (Fig. 6c and S27b), as discussed later and at elevated temperature (e.g., 80 °C; Fig. S27c). The concentration of **1a** solubilized in  $\text{H}_2\text{O}$  upon encapsulation was calculated to be 0.1 mM by UV-visible and ICP-AES studies (1.0 mM based on **PBA**; Fig. S29). The core size and composition of the product were estimated to be 2.6 nm and 10 : 1 **PBA/1a**, respectively, by

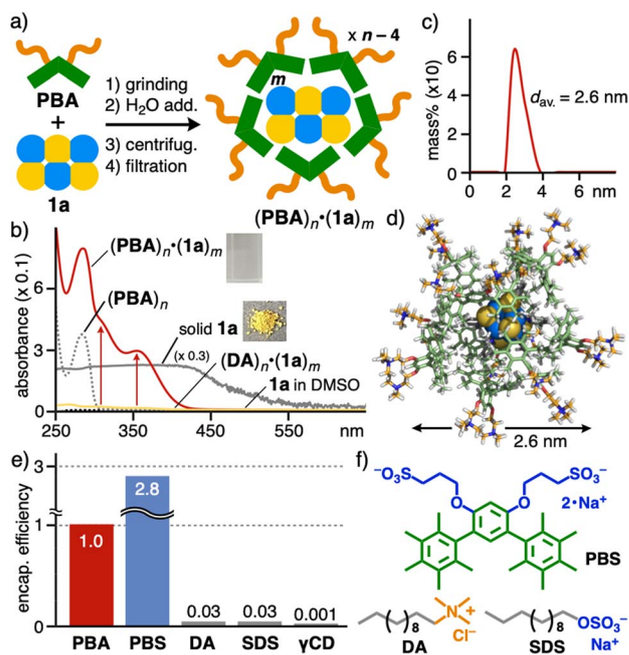


Fig. 2 (a) Preparation of an aqueous solution of the host-guest complex  $(\text{PBA})_n \cdot (\text{1a})_m$ . (b) UV-visible spectra ( $\text{H}_2\text{O}$ , r.t., 1.0 mM based on **PBA**) of  $(\text{PBA})_n \cdot (\text{1a})_m$ ,  $(\text{DA})_n \cdot (\text{1a})_m$ , and  $(\text{PBA})_n$  in  $\text{H}_2\text{O}$ , **1a** in DMSO (<10  $\mu\text{M}$ , saturated), and solid **1a**. (c) DLS chart ( $\text{H}_2\text{O}$ , r.t.) of  $(\text{PBA})_n \cdot (\text{1a})_m$ . (d) Optimized structure of  $(\text{PBA})_{10} \cdot \text{1a}$ . (e) Relative encapsulation efficiency of various amphiphiles and  $\gamma\text{CD}$  toward **1a** in water. (f) Aromatic and alkyl amphiphiles studied herein.

the DLS (Fig. 2c) and UV-visible analyses (Fig. S29). On the basis of these experimental results and the crystal structure of **1a**,<sup>15</sup> molecular modeling studies indicated the formation of  $(\text{PBA})_{10} \cdot \text{1a}$ , with a core diameter of 2.6 nm, as the major product (Fig. 2d). The present efficient encapsulation most probably stems from the hydrophobic effect and multiple  $\text{CH}-\pi$  interactions between **PBA** and **1a**.

To compare encapsulation efficiency, alkyl amphiphiles **DA** and **SDS**, bent aromatic amphiphile **PBS** with anionic hydrophilic groups (Fig. 2f), and  $\gamma$ -cyclodextrin ( $\gamma\text{CD}$ )<sup>16</sup> were employed under the same conditions. From the UV-visible spectra of the resultant host-guest complexes (Fig. S20), the cluster-based band intensities ( $\lambda = 353$  nm) revealed low efficiencies with dodecyltrimethylammonium chloride (**DA**), sodium dodecyl sulfate (**SDS**), and  $\gamma\text{CD}$  (<0.03-fold) yet high efficiency with **PBS** (2.8-fold), relative to that of **PBA** (Fig. 2e). These results revealed the unusability of the conventional micellar and tubular hosts as well as supported the importance of host-guest  $\text{CH}-\pi$  interactions for the efficient preparation of the **1a**-based host-guest solution.

### Strong emission of aqueous host-guest complexes

The aqueous host-guest solution of  $(\text{PBA})_n \cdot (\text{1a})_m$  in hand showed the strongest emission under ambient conditions, among the tested host-guest solutions. No solution-state emission was detected from **1a** in any common solvents (e.g., DMSO and  $\text{CH}_2\text{Cl}_2$ ; Fig. 3a, c, and S22a), owing to its



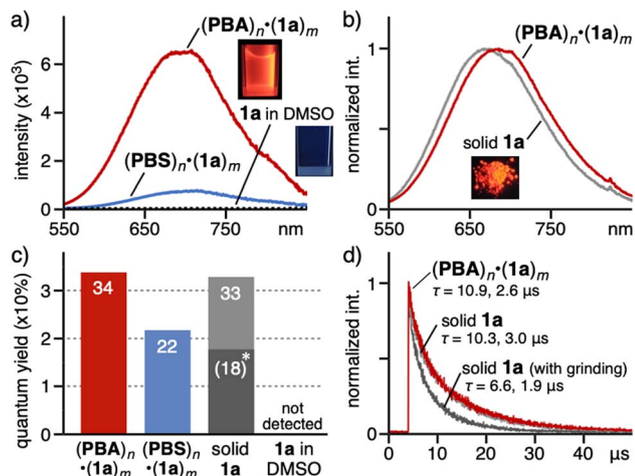


Fig. 3 (a) Emission spectra (r.t.,  $\lambda_{ex} = 320$  nm, 1.0 mM based on amphiphiles) of  $(PBA)$  or  $(PBS)_n \cdot (1a)_m$  in  $H_2O$  and  $1a$  in DMSO ( $<10 \mu M$ , saturated), and (b) the comparison with that of solid  $1a$ . (c) Emission quantum yields ( $H_2O$ ,  $\lambda_{ex} = 320$  nm) of  $(PBA)$  or  $(PBS)_n \cdot (1a)_m$ , solid  $1a$  (\*: after grinding), and  $1a$  in DMSO. (d) Emission lifetimes ( $\lambda_{ex} = 340$  nm) of  $(PBA)_n \cdot (1a)_m$  in  $H_2O$  ( $\lambda_{det} = 685$  nm) and solid  $1a$  (with/without grinding,  $\lambda_{det} = 695/670$  nm).

insolubility. In contrast, the solution of  $(PBA)_n \cdot (1a)_m$  displayed intense red emission, derived from the triplet-state cluster center,<sup>9b</sup> with a broad band at  $\lambda_{max} = 685$  nm and a quantum yield ( $\Phi$ ) of 34% upon light irradiation at 320 nm, under aerobic conditions.<sup>14</sup> Under the same conditions, anionic derivative  $(PBS)_n \cdot (1a)_m$  also showed red emission with slightly lower efficiency ( $\Phi = 22\%$ ), most probably owing to side chain-induced photo deactivation. Nearly no band was observed in the emission spectrum of  $(\gamma CD)_n \cdot (1a)_m$  in water (Fig. S20),<sup>14</sup> due to the poor uptake efficiency. The emission band of  $(PBA)_n \cdot (1a)_m$  in water was slightly red-shifted ( $\Delta\lambda_{em} = +15$  nm) relative to that of solid  $1a$  (Fig. 3b), in contrast to their absorption bands (Fig. 2b). Remarkably, the emission efficiency is comparable to that of solid  $1a$  ( $\Phi = 33\%$ ) yet much higher than that of ground solid  $1a$  ( $\Phi = 18\%$ ; Fig. 3c and S24),<sup>14,17</sup> even including the grinding process. The emission lifetime analysis of  $(PBA)_n \cdot (1a)_m$  in water elucidated its phosphorescence ( $\tau = 10.9$  and  $2.6 \mu s$ ; Fig. 3d), which is usually largely quenched under aerobic conditions. The lifetime is slightly longer than that of solid  $1a$  ( $\tau = 10.3 \mu s$ ).<sup>18</sup> These experimental and theoretical structural analyses (Fig. 2d) indicated that the present unusual results are mainly derived from the host-based steric shielding effect against oxygen through tight host-guest interactions.

Aqueous host-guest solutions were prepared from  $Cu_6S_6$  clusters  $1b-e$  and  $1f$  with various substituents (*i.e.*,  $R = H, CH_3, Br,$  and  $CF_3$ ) at the 5-position and quinoline rings, respectively (Fig. 4a). In the same manner as  $(PBA)_n \cdot (1a)_m$ , the grinding and filtration protocol using solids  $PBA$  ( $2.0 \mu mol$ ),  $1b-f$  ( $1.0 \mu mol$ ), and  $H_2O$  ( $2.0 mL$ ) led to the formation of host-guest complexes  $(PBA)_n \cdot (1b-e)_m$  and  $(PBA)_n \cdot (1f)_m$  as colorless and pale yellow aqueous solutions, respectively.<sup>14</sup> The UV-visible spectra of the resultant solutions showed new shoulder bands at 300–420 nm for bound  $1b-e$  and a new broad band at  $\lambda_{max} = 395$  nm for

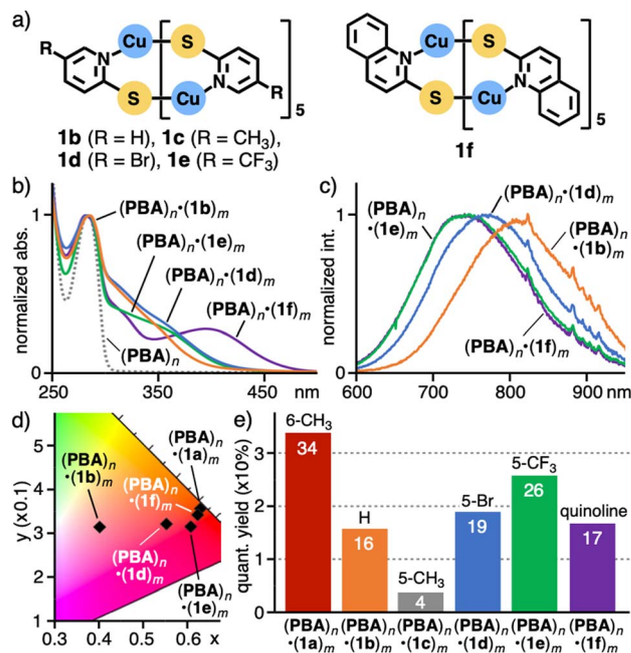


Fig. 4 (a)  $Cu_6S_6$  clusters  $1b-f$  with various substituents. (b) UV-visible spectra ( $H_2O$ , r.t., 1.0 mM based on  $PBA$ ) of host-guest complexes  $(PBA)_n \cdot (1b)$  or  $1b-f)_m$  and (c) their emission spectra ( $\lambda_{ex} = 320$  nm). (d) CIE diagram ( $H_2O$ , r.t.,  $\lambda_{ex} = 320$  nm) of  $(PBA)_n \cdot (1a-f)_m$  in water and (e) their emission quantum yields.

bound  $1f$  (Fig. 4b), which confirmed the successful water-solubilization of  $(1b-f)_m$  upon encapsulation. Emission spectra of complexes  $(PBA)_n \cdot (1b-f)_m$  in water were red-shifted ( $\Delta\lambda_{em} = +55$  to  $+120$  nm), relative to that of  $(PBA)_n \cdot (1a)_m$  (Fig. 4c), due to reduced steric hindrance between the cluster ligands, as suggested by DFT calculation (Fig. S50).<sup>18</sup> In particular, non-substituted  $(PBA)_n \cdot (1b)_m$  ( $R = H$ ) showed the largest shifted red-to-NIR emission at 805 nm ( $\Delta\lambda_{em} = +120$  nm) among the tested complexes. The emission quantum yields of host-guest complexes  $(PBA)_n$  including clusters  $1b-f$  were moderate (16–26% except for  $1c$  (4%); Fig. 4e) and their emission color was widely tunable depending on the small substituents (Fig. 4d and S18). These yields were again higher than those of ground solids  $1b-f$  without  $PBA$  (1.1 to 17-fold; Fig. S24 and S25), because of the loss of their crystallinity and the generation of undesired intermolecular contact in the solids.

#### Efficient emission and stabilization of larger and smaller multinuclear clusters within aromatic micelles

Larger/smaller multinuclear clusters such as  $Cu_{12}S_6$  cluster **2** with four bis(diphenylphosphino)pentane ligands and  $Cu_4I_4$  clusters **3a** and **3b** (Fig. 5a) were encapsulated by aromatic micelles in water and the resultant aqueous solutions emitted moderate red and strong yellow/green phosphorescence, respectively. In a similar manner to  $(PBA)_n \cdot (1a)_m$ , host-guest complex  $(PBA)_n \cdot (2)_m$  was obtained as a nearly colorless aqueous solution, using  $PBA$  ( $2.0 \mu mol$ ), **2** ( $0.3 \mu mol$ ), and  $H_2O$  ( $2.0 mL$ ), in an optimized ratio. The UV-visible and emission spectra showed a new broadened shoulder band (300–500 nm; Fig. 5b)



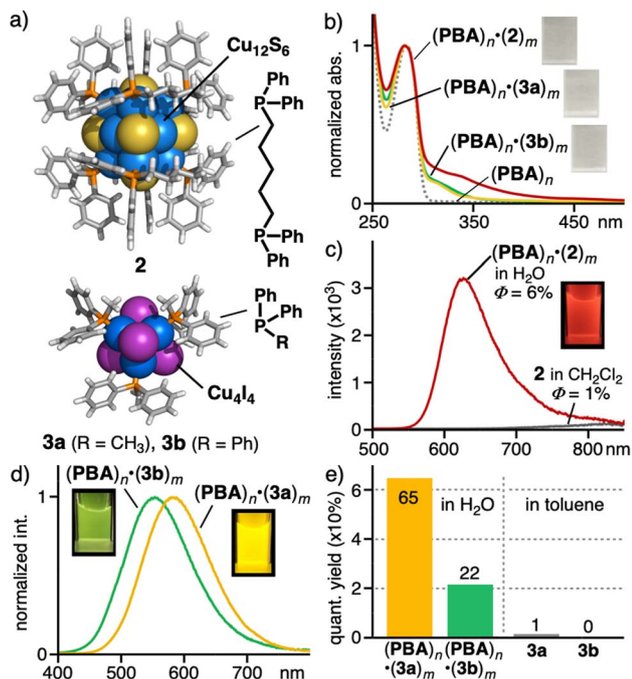


Fig. 5 (a)  $\text{Cu}_{12}\text{S}_6$  cluster **2** and  $\text{Cu}_4\text{I}_4$  clusters **3a** and **3b**. (b) UV-visible spectra ( $\text{H}_2\text{O}$ , r.t., 1.0 mM based on PBA) of  $(\text{PBA})_n \cdot (\text{2})_m$  and  $(\text{PBA})_n \cdot (\text{3a})_m$  and  $(\text{PBA})_n \cdot (\text{3b})_m$ . (c) Emission spectra (r.t.,  $\lambda_{\text{ex}} = 320$  nm, 1.0 mM based on PBA) of  $(\text{PBA})_n \cdot (\text{2})_m$  in  $\text{H}_2\text{O}$  and **2** in  $\text{CH}_2\text{Cl}_2$  (<0.1 mM, saturated). (d) Emission spectra ( $\text{H}_2\text{O}$ , r.t.,  $\lambda_{\text{ex}} = 320$  nm, 1.0 mM based on PBA) of  $(\text{PBA})_n \cdot (\text{3a})_m$  and  $(\text{PBA})_n \cdot (\text{3b})_m$  and (e) their emission quantum yields ( $\lambda_{\text{ex}} = 320$  nm; **3a** and **3b** in toluene (0.1 mM)).

and a relatively sharp band ( $\lambda_{\text{max}} = 625$  nm,  $\Phi = 6\%$ ; Fig. 5c), respectively, attributed to bound cluster **2**. In contrast, the solution of **2** in  $\text{CH}_2\text{Cl}_2$  showed a very weak emission band ( $\lambda_{\text{em}} = >650$  nm,  $\Phi = 1\%$ ; Fig. 5c), derived from its decomposed clusters through aerobic oxidation.<sup>10a,c,19</sup> Colorless aqueous solutions of host-guest complexes  $(\text{PBA})_n \cdot (\text{3a})_m$  and  $(\text{PBA})_n \cdot (\text{3b})_m$  were also obtained by the treatment of PBA with **3a** bearing four methyl-diphenylphosphines or **3b** bearing four triphenylphosphines,<sup>14,20</sup> in a manner similar to the preparation of  $(\text{PBA})_n \cdot (\text{2})_m$ . These solutions displayed strong yellow emission ( $\lambda_{\text{max}} = 580$  nm,  $\Phi = 65\%$ ) and moderate green one ( $\lambda_{\text{max}} = 550$  nm,  $\Phi = 22\%$ ), respectively (Fig. 5d and 5e), while their absorption spectra were comparable (Fig. 5b). The high quantum yield most likely stems from the steric restriction of the ligand rotation on **3a** within  $(\text{PBA})_n$  (Fig. S41).<sup>14</sup> Whereas clusters **3a** and **3b** are soluble in toluene (Fig. S38a and S38b), the resultant solutions provided quite poor emission properties under aerobic conditions (e.g.,  $\Phi = 0$ –1%; Fig. 5e, S38a and S38b), in sharp contrast to their host-guest complexes.<sup>21</sup> Within the aromatic micelles in solution, thus, the cluster structures of **2** and **3a** and **3b** were effectively stabilized against air and non-radiative relaxation, respectively (Fig. S33 and S37),<sup>14,22</sup> again due to the host-based shielding effect.

### Applications as security ink

Host-guest complex  $(\text{PBA})_n \cdot (\text{1a})_m$  in hand showed a potential application as security ink to realize anti-counterfeiting.<sup>23</sup> Like the

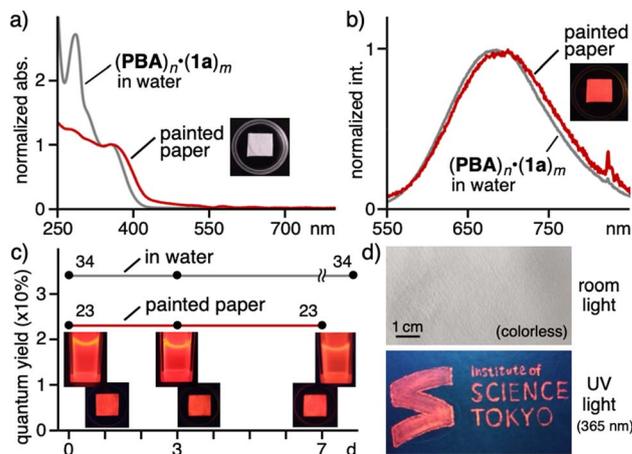


Fig. 6 (a) UV-visible and (b) emission spectra (r.t., 1.0 mM based on PBA,  $\lambda_{\text{ex}} = 320$  nm) of  $(\text{PBA})_n \cdot (\text{1a})_m$  painted on paper and in  $\text{H}_2\text{O}$ . (c) Time-dependent, emission quantum yields (r.t., 1.0 mM based on PBA,  $\lambda_{\text{ex}} = 320$  nm) of  $(\text{PBA})_n \cdot (\text{1a})_m$  in  $\text{H}_2\text{O}$  (undegassed) and painted on paper (stored in  $\text{N}_2$ ). (d) Photographs of a university logo and name, drawn with an aqueous  $(\text{PBA})_n \cdot (\text{1a})_m$  solution, taken under room light (top) and UV light (365 nm, bottom).

aqueous solution, its painting on paper (i.e., cellulose filter paper) was colorless under room light, only with a weak absorption band in the visible region in the UV-visible diffuse reflectance spectrum (Fig. 6a). In contrast, the painted paper displayed red-to-NIR emission with strong intensity ( $\Phi = 23\%$ ) upon UV-light irradiation under ambient conditions ( $\lambda_{\text{ex}} = 320$  nm, Fig. 6b). An emission band was clearly observed in the red-to-NIR region ( $\lambda_{\text{max}} = 700$  nm) from the paper in the spectrum, which is comparable to that of  $(\text{PBA})_n \cdot (\text{1a})_m$  in water. The present ink provides several advantages: (i) environmentally benign water can be used as the solvent and the host-guest solution is odorless, (ii) the aggregation-caused visible absorption (<560 nm) from solid **1a** is fully suppressed upon encapsulation, (iii) the small particle size ( $\sim 3$  nm) excludes light scattering on the paper, which interferes with invisibility, (iv) the red-to-NIR emission from the solution remains intact ( $\sim 100\%$  retention), without any visible coloration, even after standing for more than 1 week (at least 10 d) even under ambient, aerobic conditions (Fig. 6c and S27b), (v) the emissivity of the painted paper is also maintained virtually ( $\sim 100\%$  retention) after 1 week under anaerobic conditions (Fig. 6c)<sup>24</sup> and after 10 min at elevated temperature (i.e., 80 °C; Fig. S44c), and (vi) after washing with organic solvents (i.e., methanol), the red-to-NIR emission is fully retained ( $\sim 100\%$ ; Fig. S45 and S46a).<sup>24</sup> Accordingly, names and logos could be facilely drawn on paper (Fig. 6d). The potential security inks with red-to-NIR emission and large Stokes shifts ( $\Delta\lambda > 340$  nm) are rare so far, without the use of rare-earth elements and noble metals (e.g., Nd, Yb, Ir, and Pt).<sup>25</sup>

## Conclusions

We have accomplished the preparation of aqueous red-to-NIR phosphorescent solutions of multinuclear  $\text{Cu}_n\text{S}_m$  clusters upon facile encapsulation for the first time. Once



mercaptopyridine-based  $\text{Cu}_6\text{S}_6$  clusters, which hardly emit by themselves in solution, were encapsulated by aromatic micelles in water, the resultant host-guest solutions exhibited strong red-to-NIR phosphorescent emission. Unlike previously reported aliphatic micelles as well as coordination cages, the present micelle efficiently encapsulated various multinuclear clusters, such as  $\text{Cu}_{12}\text{S}_6$  and  $\text{Cu}_4\text{I}_4$  clusters, generating aqueous phosphorescent solutions with various emission colors. Colorless yet strong red-to-NIR emissive features, besides high water, air, and thermal stability as well as large scale synthesis, established the present host-guest solutions as potential security ink. Further preparation of multinuclear cluster-based solutions (e.g., large heteronuclear Au, Mn, and Pt clusters) via this encapsulation strategy would provide new materials and catalytic applications in water.

## Author contributions

K. T., Y. T., and M. Y. designed the work, carried out the research, analyzed the data, and wrote the paper. M. Y. is the principal investigator. All authors discussed the results and commented on the manuscript.

## Conflicts of interest

There are no conflicts to declare.

## Data availability

CCDC 2420528 and 2420529 contain the supplementary crystallographic data for this paper.<sup>15</sup>

The experimental procedures and analytical data are available in the SI. See DOI: <https://doi.org/10.1039/d5sc04122h>.

## Acknowledgements

This work was supported by JSPS KAKENHI (Grant No. JP22H00348/JP23K17913/JP25K01783). PXRD measurements were performed with the help of Hiroto Toyoda and Prof. Toshiyuki Yokoi (Science Tokyo). K. T. acknowledges JST SPRING (Grant No. JPMJSP2106) and JSPS Research Fellowship for Young Scientists (Grant No. 25KJ1257). The theoretical calculations were performed using computers in the Research Center for Computational Science, Okazaki, Japan (24-IMS-C060 and 25-IMS-C062).

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- 14 See the SI. As a scale-up experiment, a 100 mL solution of  $(PBA)_n \cdot (1a)_m$  in water (0.2 mM based on PBA) was also obtained by a similar grinding protocol. The photophysical properties (Fig. S28 and S49) were comparable to those of the 1 mL scale (Fig. 2b and 3a). The ICP-AES analysis of  $(PBA)_n \cdot (1a)_m$  (0.1 mM based on PBA) also indicated the same PBA/1a ratio, with a Cu concentration of 4.09 ppm ( $\approx 0.01$  mM of 1a). The photophysical properties of  $(PBA)_n \cdot (1a)_m$  were less effective against its concentration, likely due to the isolation effect.
- 15 The single crystals of 1a were obtained from the slow diffusion between an acidic acetone/H<sub>2</sub>O solution of 2-mercapto-6-methylpyridine and an acidic CH<sub>3</sub>OH solution of CuSO<sub>4</sub>·5H<sub>2</sub>O for 11 d at r.t. (Fig. S2; CCDC-2420528).<sup>14</sup>
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