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Solvent vapour-responsive structural transformations in molecular crystals composed of a luminescent mononuclear aluminium(III) complex

Solvent vapour-induced structural transformations of molecular crystals composed of [Al(sap)(acac)(H<sub>2</sub>O)]·(solvent) (solvent = Me<sub>2</sub>CO, MeCN, DMSO) have been demonstrated. This solvent vapour-responsive system is formed using a versatile mononuclear aluminium(III) complex that exhibits strong yellow emission in the solid state. Our results also suggest the versatility of the hydrogen-bonded dimer unit [Al(sap)(acac) (H<sub>2</sub>O)]<sub>2</sub> as a molecular building block unit for the development of an advanced switching system associated with structural rearrangements.

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### Introduction

Molecular materials that respond to external stimuli have attracted significant attention owing to their potential applications in sensors and memory devices. These external stimuli, such as heat, light, and pressure, induce functional changes in the materials, including alterations in magnetic,<sup>1-9</sup> luminescent,<sup>10-14</sup> and dielectric properties.<sup>15-18</sup> In recent years, several molecules responsive to external stimuli and their functional control have been reported within a group of compounds referred to as "soft crystals".<sup>19</sup> Among them, molecules responsive to solvent vapours are particularly intriguing as they allow reversible functional control in the solid state, without the need for recrystallization from solution.<sup>20-35</sup> In conventional molecular crystals, unlike rigid frameworks such as metal-organic frameworks (MOFs) and porous coordination polymers (PCPs),<sup>36-44</sup> the removal or desorption of solvent molecules from the pores can induce structural changes, such

## Solvent vapour-responsive structural transformations in molecular crystals composed of a luminescent mononuclear aluminium(III) complex<sup>+</sup>

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Investigations into the construction of functional molecular crystals and their external stimuli-induced structural transformations represent compelling research topics, particularly for the advancement of sensors and memory devices. However, reports on the development of molecular crystals constructed from discrete mononuclear complex units and exhibiting structural transformations *via* the adsorption/desorption of guest molecules are scarce. In this study, we synthesised three molecular crystals composed of [Al(sap)(acac)(H<sub>2</sub>O)]-(solvent) (H<sub>2</sub>sap = 2-salicylideneaminophenol, acac = acetylacetonate, solvent = Me<sub>2</sub>CO (Al·Me<sub>2</sub>CO), MeCN (Al·MeCN), or DMSO (Al·DMSO)), and demonstrated solvent vapour-responsive reversible crystal-to-crystal structural transformations in Al·Me<sub>2</sub>CO and Al·MeCN. For Al·DMSO, exposure to DMSO vapour led to the formation of DMSO-coordinated compound [Al(sap)(acac)(DMSO)], indicating an irreversible structural transformation. This solvent vapour-responsive system incorporates a luminescent mononuclear aluminium(III) complex ( $\lambda_{max} = 539-552$  nm,  $\phi_{em} = 0.07-0.27$ ) as the molecular building unit for the porous-like framework. Therefore, we synthesised a new functional molecular material and a potential molecular building unit that facilitates guest fixation through hydrogen-bonding.

as shrinkage or collapse, which may then lead to closing of the pores. This typically prevents subsequent reversible re-adsorption. However, reversible guest adsorption and desorption, along with functional control, have been reported in several systems comprising discrete molecules.<sup>44–56</sup> For example, Ito *et al.* reported solvent vapour-induced reversible structural rearrangements and luminescent colour changes in crystals composed of a mononuclear gold(1) isocyanide complex.<sup>57</sup> However, intentionally incorporating guest molecules, such as solvent molecules, and designing molecules that exhibit reversible adsorption and desorption characteristics can be highly challenging.

In our previous study, we reported a solvent vapor-responsive molecular system consisting of a mononuclear complex of type [M(sap)(acac)(solvent)] (where  $M = Fe^{III}$  or  $AI^{III}$ ,  $H_2sap =$ 2-salicylideneaminophenol, acac = acetylacetonate, solvent = MeOH, DMSO, or pyridine), with coordination sites for solvent molecules.<sup>58,59</sup> In this system, molecular arrangement and associated functionality can be controlled through solvent vapor exposure-induced substitution reactions of coordinating solvents. In addition, intentional incorporation of solvent vapor responsiveness into this molecular system is possible, potentially enabling the development of highly functional molecular materials by combining it with functionality arising from metal ions and aggregated structures. As a part of this research, we report the synthesis of new solvent vapour-respon-

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<sup>†</sup>Electronic supplementary information (ESI) available: PXRD patterns, IR spectra, crystallographic data, crystal structures, luminescence properties, TGA, DFT and TD-DFT calculations. CCDC 2337339–2337341. For ESI and crystallographic data in CIF or other electronic format see DOI: https://doi.org/10.1039/ d4dt00747f

#### Paper

sive molecular crystals of  $[Al(sap)(acac)(H_2O)]$  (solvent) (solvent) = Me<sub>2</sub>CO, MeCN, or DMSO), their crystal-to-crystal structural transformations, and their photo-luminescence properties. The hydrogen-bonded dimers of  $[Al(sap)(acac)(H_2O)]$  units formed a one-dimensional (1D) channel structure, securing solvent molecules within the pores through hydrogen-bonding. The channel structures and lattice solvents exhibited reversible recovery upon exposure to solvent vapours (Scheme 1).

### **Results and discussion**

### Syntheses and single-crystal structural analyses

[Al(sap)(acac)(MeOH)] was synthesised according to our previously reported method. <sup>58,59</sup> Single crystals of [Al(sap)(acac) (H<sub>2</sub>O)]·(Me<sub>2</sub>CO) (Al·Me<sub>2</sub>CO), [Al(sap)(acac)(H<sub>2</sub>O)]·(MeCN) (Al·MeCN), and [Al(sap)(acac)(H<sub>2</sub>O)]·(DMSO) (Al·DMSO) were obtained by recrystallising [Al(sap)(acac)(MeOH)] from Me<sub>2</sub>CO, MeCN, and DMSO, respectively (see Experimental section for details). All compounds were then characterised using elemental analysis, single-crystal X-ray diffraction (XRD), powder X-ray diffraction (PXRD), and Fourier-transform infrared (FT-IR) spectroscopy (Fig. S1 and S2, ESI†).

The single-crystal X-ray structures of Al·Me<sub>2</sub>CO, Al·MeCN, and Al-DMSO were obtained at 173 K, and the crystallographic data are displayed in Table S1 (ESI<sup>†</sup>). Al·Me<sub>2</sub>CO crystallised in the orthorhombic Pbca space group with an asymmetric unit containing one lattice Me<sub>2</sub>CO molecule (Fig. 1a and Fig. S3 and S4, ESI<sup>†</sup>). The structure of Al·Me<sub>2</sub>CO was found to be similar to the previously reported structure of the aluminium (III) complex [Al(sap)(acac)(MeOH)].<sup>59</sup> The aluminium(III) centre possesses a NO<sub>5</sub>-donor coordination sphere with a distorted octahedral geometry. The metal centre has a coordinated NO3donor set equatorially arising from the sap and acac ligands, while the axial positions are occupied by O and O' donors from acac and  $H_2O$ , respectively. Each [Al(sap)(acac)(H\_2O)] molecule exhibits stereoisomers and forms dimers with neighbouring isomers by complementary hydrogen-bonding between coordinated water molecules and phenol oxygen atoms from the sap ligands (O(2)-O(5) = 2.645(2) Å) (Fig. 2 and Fig. S5a, ESI<sup>†</sup>). Each dimer interacts with neighbouring dimers by CH- $\pi$ , CH-O and CH-N interactions, leading to a supramo-



**Scheme 1** Structural transformations for [Al(sap)(acac)(H<sub>2</sub>O)] (solvent).



Fig. 1 Single-crystal structures of (a)  $[Al(sap)(acac)(H_2O)] \cdot (Me_2CO)$ (Al·Me<sub>2</sub>CO), (b)  $[Al(sap)(acac)(H_2O)] \cdot (MeCN)$  (Al·MeCN), and (c)  $[Al(sap)(acac)(H_2O)] \cdot (DMSO)$  (Al·DMSO) at 173 K. Colour code: C, grey; H, white; N, blue; O, red; Al, pink; S, yellow.



Fig. 2 Dimeric structure of Al·Me<sub>2</sub>CO constructed by hydrogenbonding. Blue-dashed lines represent the hydrogen-bonding. Colour code: C, grey; N, blue; O, red; Al, pink.

lecular framework and a 1D channel structure along the b axis (Fig. 3a and Fig. S6a, Table S2, ESI<sup>†</sup>). The lattice Me<sub>2</sub>CO molecules present in the channel pores are stabilised by hydrogenbonding with the coordinated water molecules (O(5)-O(6) =2.845(2) Å) (Fig. S7a, ESI†). Compared to the previously reported crystal structure of the non-porous [Al(sap)(acac) (MeOH)] complex,<sup>59</sup> the bidirectional hydrogen-bonding of the coordinated water molecules in Al·Me2CO contributes to the fixation of guest molecules and the formation of a porous framework. Al-MeCN has an identical structure to that of Al·Me<sub>2</sub>CO and also forms dimers through hydrogen-bonding (O(2)-O(5) = 2.654(3) Å) (Fig. 1b and Fig. S8, ESI<sup>†</sup>). Therefore, Al-MeCN crystallises in the orthorhombic space group Pbca with an asymmetric unit containing one lattice MeCN molecule and forms a 1D channel structure along the *b* axis (Fig. 3b and Fig. S6b, S9, Table S3, ESI<sup>†</sup>). The lattice MeCN molecules



**Fig. 3** Molecular packing diagrams of (a) **Al·Me<sub>2</sub>CO**, (b) **Al·MeCN**, and (c) **Al·DMSO** viewed along the *b*-axis. All hydrogen atoms are omitted for clarity. Colour code: C, grey; N, blue; O, red; Al, pink; S, yellow.

present in the channel pores are stabilised by hydrogenbonding with the coordinated water molecules (N(2)-O(5) =2.886(2) Å) (Fig. S5b and S7b, ESI<sup>†</sup>). Al·DMSO has a similar molecular structure to that of Al·Me<sub>2</sub>CO and Al·MeCN; however, it crystallises in the monoclinic space group  $P2_1/n$ with an asymmetric unit containing one lattice DMSO molecule (Fig. 1c and Fig. S10, S11, ESI<sup>+</sup>). The intermolecular interactions observed in Al·DMSO are also similar to those observed in Al·Me<sub>2</sub>CO and Al·MeCN, resulting in the formation of a 1D channel structure along the b axis (Fig. 3c and Fig. S5c, S6c, Table S4, ESI<sup>†</sup>). The decrease in the symmetry of the crystal system in Al-DMSO can be attributed to the intermolecular interactions involving the low-symmetry DMSO molecule (Fig. S7c, ESI<sup>†</sup>). The results of Hirshfeld  $d_{\text{norm}}$  surface analysis for Al·Me<sub>2</sub>CO, Al·MeCN and Al·DMSO are shown in Fig. S12,† displaying the observed intermolecular interactions visibly. Thermogravimetric analysis (TGA) was used to determine the number of lattice solvents in Al·Me<sub>2</sub>CO, Al·MeCN, and Al-DMSO and their thermal stabilities. The TGA results for Al·Me<sub>2</sub>CO and Al·MeCN showed 17.7% and 15.1% weight

losses corresponding to one water molecule and one solvent molecule (Me<sub>2</sub>CO and MeCN) at temperatures above 110 °C, respectively, thus corroborating the results of single-crystal X-ray diffraction (Fig. S13a and b, ESI†). Moreover, the de-solvated compound [Al(sap)(acac)] exhibited high thermal stability at temperatures below 250 °C. In contrast, a two-step weight loss of 22.2% was observed for Al-DMSO at 145 and 230 °C, corresponding to one water molecule and one DMSO molecule. Then, further weight loss was observed, which was attributed to the decomposition of Al-DMSO (Fig. S13c, ESI†). Thermogravimetric mass spectrometry (TG-MS) results for Al-DMSO indicate that both the observed first and second steps can be attributed to the de-solvation of a water molecule and a DMSO molecule, respectively.

### Crystal-to-crystal structural transformations

Owing to the porous-like nature and high thermal stabilities of Al·Me<sub>2</sub>CO and Al·MeCN, we investigated their solvent vapourresponsive structural transformations. When the Me<sub>2</sub>CO and MeCN-containing complexes Al·Me2CO and Al·MeCN were annealed at 100 °C for 1 h, their initial PXRD patterns changed (Fig. 4). These patterns were found to be identical and corresponded to the de-solvated five-coordinated compound [Al(sap)(acac)]. When the de-solvated compounds were exposed to Me<sub>2</sub>CO or MeCN vapour for 1 h, the PXRD patterns changed again (Fig. 4 and S14, ESI<sup>†</sup>). These PXRD patterns corresponded to the original patterns for Al·Me<sub>2</sub>CO and Al·MeCN, respectively. This result is in accordance with the coordination of water molecules upon exposure to solvent vapour, resulting in structural rearrangements to the 1D channel structure. The TGA results for Al·reMe<sub>2</sub>CO and Al·reMeCN showed 17.3% and 14.1% weight losses, corresponding to adsorbed water molecules and solvent molecules (Me<sub>2</sub>CO, MeCN), respectively (Fig. S15, ESI<sup>†</sup>). Notably, the observed PXRD transformations and de-sorption/adsorption process were found to be reversleading ible, crystal-to-crystal to structural transformations.<sup>60-63</sup> Unlike the results of Al·Me<sub>2</sub>CO and Al·MeCN mentioned above, Al·DMSO exhibited different structural transformations. When the DMSO-containing complex Al·DMSO was annealed at 200 °C for 1 h, the initial PXRD pattern changed to those of de-solvated compound [Al(sap) (acac)]. After exposing the de-solvated compound to DMSO vapour for 1 day, the PXRD pattern changed again. However, the obtained pattern corresponded to that of the DMSO-coordinated compound [Al(sap)(acac)(DMSO)]<sup>59</sup> rather than to Al·DMSO (Fig. 4c). This result indicates that the structural transformation to [Al(sap)(acac)(DMSO)] takes precedence over that to Al·DMSO under DMSO vapour. This trend is in accordance with the spontaneous DMSO vapour-induced coordinated solvent molecule exchange from [Al(sap)(acac)(MeOH)] and [Al (sap)(acac)(EtOH)] to [Al(sap)(acac)(DMSO)], as previously reported by us.59 The TGA result for Al·reDMSO supported the above result and showed one-step weight loss of 19.3%, corresponding to the coordinated DMSO molecule (Fig. S15c, ESI<sup>+</sup>). Thus, Al·DMSO exhibited irreversible structural transformations.

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Fig. 4 PXRD patterns for the de-solvated compounds [Al(sap)(acac)] and those obtained after exposure to (a)  $Me_2CO$ , (b) MeCN, and (c) DMSO vapours.

#### Luminescence properties

We also investigated the luminescence properties of the aluminium(III) complexes.<sup>64–67</sup> The solid-state excitation and emission spectra of Al·Me<sub>2</sub>CO, Al·MeCN, and Al·DMSO obtained at 298 K are shown in Fig. S16† and Fig. 5 (Table S5, ESI†), respectively. At 298 K, the crystalline solid samples of Al·Me<sub>2</sub>CO, Al·MeCN, and Al·DMSO exhibited strong yellow emission. The emission spectra of Al·Me<sub>2</sub>CO, Al·MeCN, and Al·DMSO were similar and showed unstructured broad emission bands. However, their maxima ( $\lambda_{max}$ ) were slightly different; 539 nm for Al·Me<sub>2</sub>CO, 540 nm for Al·MeCN, and 552 nm for Al·DMSO. Emission quantum yield ( $\Phi_{em}$ ) measure-



Fig. 5 Emission spectra of  $Al \cdot Me_2CO$  (red),  $Al \cdot MeCN$  (blue), and  $Al \cdot DMSO$  (black) in the solid state at 298 K.

ments in the solid state revealed strong emissions from Al·Me<sub>2</sub>CO (0.27), Al·MeCN (0.22), and Al·DMSO (0.07) at 298 K. The observed differences in the photophysical properties of Al·Me<sub>2</sub>CO, Al·MeCN, and Al·DMSO can be attributed to the differences in their crystal structure and their resulting intermolecular interactions. The emission spectra of Al·Me<sub>2</sub>CO in Me<sub>2</sub>CO, Al·MeCN in MeCN and Al·DMSO in DMSO at 298 K  $(1.0 \times 10^{-5} \text{ M})$  are shown in Fig. S18 (Fig. S17 and Table S6, ESI<sup>†</sup>). The emission maxima for Al·Me<sub>2</sub>CO, Al·MeCN, and Al-DMSO were 523, 526, and 527 nm, respectively. The emission quantum yields for Al·Me<sub>2</sub>CO, Al·MeCN, and Al·DMSO in solution at 298 K were 0.47, 0.42, and 0.76, respectively, revealing stronger emissions than those in the solid state. The emission intensity of Al·DMSO was significantly higher than those of Al·Me<sub>2</sub>CO and Al·MeCN. This can be attributed to the coordination of the DMSO molecule in the aluminium(m) complex instead of the water molecule. This coordination decreases the nonradiative rate associated with the OH vibrations in water molecules. The previously reported DMSOcoordinated aluminium(m) complex, [Al(sap)(acac)(DMSO)], exhibits strong green emission even in the solid state ( $\Phi_{\rm em}$  = 0.42, at 298 K).<sup>59</sup> This result suggests that replacing the coordinated water molecules with other coordinating solvents decreases the nonradiative rates.

### DFT and TDDFT calculations

Density functional theory (DFT) and time-dependant DFT (TDDFT) calculations for the [Al(sap)(acac)(H<sub>2</sub>O)] (Al) unit were performed to investigate the relationship between its structures and emission origins. The highest occupied molecular orbital (HOMO) and lowest unoccupied molecular orbital (LUMO) energies for Al are shown in Fig. 6 (Tables S7 and S8, ESI†). The singlet excited states of Al were also calculated using TDDFT. The lowest energy of the spin-allowed electronic vertical transition for Al [ $\Delta E(S_0-S_1)$ ] was 2.93 eV (424 nm). The intraligand transition occurs from the  $\pi$  HOMO delocalised in the sap ligand to the  $\pi^*$  LUMO delocalised in the sap ligand. In addition, the ligand-to-ligand charge transfer (LLCT) occurs from the HOMO delocalised in the sap ligand to the LUMO+1 delocalised in the acac ligand. These results indicate that the



Fig. 6 HOMO and LUMO of  $[Al(sap)(acac)(H_2O)]$  at the experimental X-ray geometries.

S<sub>1</sub> states of Al include singlet ligand-centred (<sup>1</sup>LC) and <sup>1</sup>LLCT excited states. The above results are similar to those reported for [Al(sap)(acac)(MeOH)].59 We also performed DFT and TDDFT calculations for the hydrogen-bonded dimer of [Al(sap) (acac)(H<sub>2</sub>O)]<sub>2</sub> ([Al]<sub>2</sub>). The HOMO and LUMO orbitals for [Al]<sub>2</sub> are very similar with those of Al (Fig. S19 and Tables S9, S10, ESI<sup>†</sup>). Comparing the molecular orbital energies of Al and [Al]<sub>2</sub>, dimerisation via hydrogen bonding leads to the stabilisation of the HOMO energy. As a result, the lowest energy of the spin-allowed electronic vertical transition for  $[AI]_2 [\Delta E(S_0 - S_1)]$ was 3.05 eV (406 nm). Thus, the differences in intermolecular interactions observed in Al·Me<sub>2</sub>CO, Al·MeCN, and Al·DMSO are reflected in the photophysical properties. In particular, because the molecular orbital of the water molecule is included in the HOMO in Al, the short hydrogen bond between the coordinated water molecule and DMSO (O(5)-O(6) = 2.722(8) Å) may affect the red-shifted emission bands of Al·DMSO.

### Conclusions

In conclusion, we demonstrated the solvent vapour-induced structural transformations of molecular crystals composed of  $[Al(sap)(acac)(H_2O)]$  (solvent) (solvent = Me<sub>2</sub>CO (Al·Me<sub>2</sub>CO), MeCN (Al·MeCN), DMSO (Al·DMSO)). Interestingly, this solvent vapour-responsive system was formed using a versatile mononuclear aluminium(m) complex that exhibits strong yellow emission in the solid state ( $\lambda_{max} = 539-552$  nm,  $\Phi_{em} =$ 0.07-0.27, at 298 K). This study presents the first example of a solvent vapour-responsive luminescent porous-like crystal constructed from a mononuclear aluminium(III) complex. Our results also suggest the versatility of the activated five-coordinated compound [Al(sap)(acac)], which possesses a metal open site, as a molecular building block unit for the development of an advanced switching system associated with structural rearrangements. For example, these structural rearrangements can be induced by not only solvent vapour but also by the sublimation of the coordination compound. Further development of a novel solvent vapour-responsive system using the

activated Al(m) unit [Al(sap)(acac)] and the hydrogen-bonded unit [Al(sap)(acac)(H<sub>2</sub>O)] is currently underway in our laboratory.

### Experimental

#### General

All reagents and solvents were obtained from Tokyo Kasei Co. and Wako Pure Chemical Industries and were of reagent grade; they were used without further purification. All reactions were carried out under ambient atmosphere.

### Syntheses

Preparation of  $H_2$ sap (=2-salicylideneaminophenol) ligand and [Al(sap)(acac)(MeOH)].  $H_2$ sap and [Al(sap)(acac)(MeOH)] were synthesised according to the method we described previously.<sup>58,59</sup>

[Al(sap)(acac)(H<sub>2</sub>O)]·(Me<sub>2</sub>CO) (Al·Me<sub>2</sub>CO). Al·Me<sub>2</sub>CO was obtained by recrystallizing [Al(sap)(acac)(MeOH)] from Me<sub>2</sub>CO and allowing the solution to stand for a few days during which time yellow block crystals formed. They were collected by suction filtration, washed with small amount of diethyl ether, and dried in air. Anal. Calc. for Al·Me<sub>2</sub>CO (C<sub>21</sub>H<sub>24</sub>NAlO<sub>6</sub>): C, 61.01; H, 5.85; N, 3.39. Found: C, 61.02; H, 5.65; N, 3.29%. IR (KBr/cm<sup>-1</sup>): 1696, 1653, 1560, 1539, 1471, 1288, 1277, 1267, 1228, 1180, 1149, 1134, 1126, 1034, 926, 839, 762, 746.

[Al(sap)(acac)(H<sub>2</sub>O)]·(MeCN) (Al·MeCN). Al·MeCN was obtained by recrystallizing [Al(sap)(acac)(MeOH)] from MeCN and allowing the solution to stand for a few days during which time yellow block crystals formed. They were collected by suction filtration, washed with small amount of diethyl ether, and dried in air. Anal. Calc. for Al·MeCN ( $C_{20}H_{21}N_2AlO_5$ ): C, 60.60; H, 5.34; N, 7.07. Found: C, 60.20; H, 5.43; N, 7.13%. IR (KBr/cm<sup>-1</sup>): 2258, 1653, 1651, 1620, 1471, 1389, 1352, 1290, 1277, 1269, 1180, 1163, 1126, 1003, 839, 773, 766, 744, 561.

[Al(sap)(acac)(H<sub>2</sub>O)]·(DMSO) (Al·DMSO). Al·DMSO was obtained by recrystallizing [Al(sap)(acac)(MeOH)] from DMSO and allowing the solution to stand for a few days during which time yellow block crystals formed. They were collected by suction filtration, washed with small amount of diethyl ether, and dried in air. Anal. Calc. for Al·DMSO ( $C_{20}H_{24}NSAlO_6$ ): C, 55.42; H, 5.58; N, 3.23. Found: C, 55.05; H, 5.71; N, 3.12%. IR (KBr/cm<sup>-1</sup>): 1651, 1637, 1622, 1560, 1508, 1473, 1458, 1390, 1348, 1290, 1269, 1178, 1149, 1028, 998, 929, 841, 762, 744, 669, 604, 579, 548.

**Preparation of de-solvated compound [Al(sap)(acac)].** De-solvated crystals of [Al(sap)(acac)] were prepared by annealing crystals of **Al·Me<sub>2</sub>CO** and **Al·MeCN** at 100 °C for 1 h, and **Al·DMSO** at 200 °C for 1 h, respectively.

#### Solvent-vapour exposure experiments

A small amount of the crystalline sample chosen from de-solvated crystals [Al(sap)(acac)] was added to a small Petri dish. The Petri dish was placed in a larger closed Petri dish together with a Kimwipe and required solvent (Me<sub>2</sub>CO or MeCN or DMSO) as the vapor source. The closed system was left to stand for 1 h or 1 day at room temperature (25 °C) (Fig. S14†).

### Physical measurements

**General procedures.** Elemental analyses (C, H, N) were performed on a PerkinElmer 2400II CHN analyzer. Infrared (IR) spectra measurements were performed on a HORIBA FT-730 instrument equipped with KBr pellet method. TG and TG-MS measurements were performed on a Bruker TG-DTA 2010SA/ MS 9610.

Single crystal and powder X-ray diffraction. The single crystal X-ray diffraction data for all compounds were recorded on a Bruker D8 QUEST diffractometer employing graphite monochromated Mo Kα radiation generated from a sealed tube ( $\lambda = 0.7107$  Å). Data integration and reduction were undertaken with APEX3. Using Olex2 software, the structure was solved with the SHELXT structure solution program using Intrinsic-Phasing Methods and refined with the SHELXL refinement package using Least Squares minimization. Hydrogen atoms were included in idealized positions and refined using a riding model. Powder X-ray diffraction data (PXRD) for all compounds were collected on a Rigaku MiniFlex II ultra (40 kV/15 mA) X-ray diffractometer using Cu Kα radiation ( $\lambda = 1.5406$  Å) in the 2θ range of 5°–30° with a step width of 3.0°.

**Luminescence property measurements.** Emission and excitation spectra were measured using a SHIMADZU RF-6000 spectrofluorometer. Emission quantum yields in crystalline powder solid were recorded using an integrating sphere (ISR-100,  $\lambda_{ex}$  = 430 nm). UV-Vis spectra were measured using a JASCO V-550 spectrophotometer.

**Theoretical calculations.** The optimized structures at ground states of  $[Al(sap)(acac)(H_2O)]$  and  $[Al(sap)(acac)(H_2O)]_2$  were obtained using the density functional theory (DFT) method with the B3LYP functional<sup>68,69</sup> and 6-31G(d)<sup>70</sup> basis sets. The molecular structure determined by X-ray crystallographic analysis was used as the starting structure for the optimization. Time-dependent density functional theory (TD-DFT)<sup>71</sup> measurements using the hybrid B3LYP functional (TD-B3LYP) were used to obtain the electronic transition energies which also included an account of electron correlation. All calculations were carried out using the GAUSSIAN 09 program.<sup>72</sup>

## Author contributions

F. K. and M. T. planned and directed the project; A. Y. and M. G. synthesised the compounds and carried out all the measurements. Y. T. assisted during XRD and luminescence measurements and analyses. F. K. drafted the manuscript and all authors contributed to revising the manuscript.

## Data availability

The data supporting this article have been included as part of the ESI.†

## Conflicts of interest

There are no conflicts to declare.

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