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Ammonia-modulated reversible gel-solution phase transition and fluorescence switch for a salicylhydrazide-based metal-organic gel†

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Based on a multi-dentate salicylhydrazide-derived ligand, a Zn-contained metal-organic gel that could stand for months has been prepared. The driving force within the supramolecular assembly was discussed by studying the gelation conditions, optical spectra, dynamic rheology and mass spectra. These studies revealed that both the coordination bonds and hydrogen bonds participate in the supramolecular assembly. In addition, electrostatic force also played as a key role in stabilizing the gel phase. The charge nature of the colloidal particle is negative charged. Consequently, the xerogel shows significant adsorption behavior that favors positively charged dye molecules, such as methyl violet and methylene blue. The title gel exhibits gelation-induced fluorescence enhancement property. When the gel was exposed to ammonia vapor, a gel-solution transition occurred, accompanied by a great weakening of fluorescence. The phase transition and fluorescence switching were found to be reversible for at least seven cycles. The mechanism involved was studied and discussed.

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Introduction

In the last few decades, metal-organic gels (MOGs) have attracted much attention in the applications of optical materials, sensing, self-healing, tredox, chiral materials, drug delivery, catalysis, adsorption and separation etc. The advantages of MOGs include the rational design of the organic ligand, the properties of metal ions, and the processability of soft materials. By adopting the concept of supramolecular self-assembly to fabricate a MOG, coordination bonding is of key importance. The organic ligand always plays the role of a low molecular weight gelator (LMWG). By exploring the reported functional MOGs, the structures of LMWGs share common characteristics as follows: (i) multiple coordination sites for multiple metal bindings, sometimes chelated; (ii) certain function groups for molecular

Herein, a multidentate salicylhydrazide derivative $\rm H_6L$ (Scheme 1) was synthesized and studied as a LMWG to form a Zn-contained MOG. The protons of the hydrazine and phenolic hydroxyl groups in salicylhydrazide ligand are apt to be released when $\rm H_6L$ reacts with transition metal ions in the DMF solution. The typical coordination mode reported is edge sharing 5-5-6 chelated rings (Scheme 1). This chelated

Scheme 1 The structure of H_6L (up) and the popular coordination mode and hydrogen bonding mode (down).

interaction, such as amides for hydrogen bonding, ¹² aromatic groups for $\pi \cdots \pi$ stacking, ¹³ aliphatic groups for hydrophobic-hydrophilic interaction; ¹⁴ (iii) precise balance between the structural rigidity and flexibility to benefit both the minimum gelation concentration (MGC) and tunable properties. ^{10,15}

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State Key Laboratory of Structural Chemistry, Fujian Institute of Research on the Structure of Matter, Chinese Academy of Sciences, Fuzhou, Fujian 350002, PR China \dagger Electronic supplementary information (ESI) available: Part I: experimental details including synthesis of the ligand H_6L , preparation of Gel2/1, dye adsorption tests, cycling experiments of the fluorescence sensing for NH $_3$ and measurement details. Part II: synthesis and characterization of H_6L and Gel2/1 including the 1H NMR and mass spectra, IR spectra, dynamic rheology and high resolution mass spectrum. See DOI: 10.1039/c7ra02551c

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coordination geometry is fairly stable in either solution or crystalline phase.17 Secondly, as shown in Scheme 1, the salicylhydrazide group is a good candidate for hydrogen bonding in the supramolecular self-assembly. Thirdly, the two salicylhydrazides groups are spaced by a 4,4'-biphenyl oxide group. Thus the ligand might exhibit certain flexibility. In our previous study,18 when the two salicylhydrazide groups are spaced by a 2,2'-biphenyl group, the ligand adopts typical 5-5-6 chelated coordination geometries with cupric ion. By the linkage of 2,2'biphenyl group, there exists a common trinuclear coordination unit in several crystal structures.

In this paper, the gelation of the ligand H₆L was observed by the involvement of Zn2+ ion in DMF solution, showing interesting properties, such as gelation-induced fluorescence enhancement, reversible gel-solution transition, fluorescence switch, and selective dye adsorption. The characterization of the gel, as well as the intrinsic mechanism of properties were studied and discussed.

Results and discussion

The synthesis and characterization of H₆L were described in the ESI.† Typical gelation could be observed in 1 mL DMF solution that contains H_6L (0.0125 mmol) and $Zn(OAc)_2$ (0.0250 mmol). When the concentration is higher, gelation occurs but serving mash-like gel; when the concentration is lower, the resultant looks like viscous fluid. Consequently, the MGC of the gel is ca. 1.2 wt%. In this concentration, transparent gel phase could be observed when the Zn/L molar ratio equal to 0.5/1, 1/1, 2/1 and 3/1 respectively (labelled as Gelx/1 respectively, x = 0.5, 1, 2, 3) as shown in Fig. 1a. When the ratio is 4/1, no gelation occurs. In the Gelx/1 series, Gel2/1 shows the most stable gel phase that could maintain uniform gel phase for months, while Gel3/1 shows liquid extrusion after standing for several days, Gel0.5/ 1 and Gel1/1 exhibit too soft phase that easily deformed upon staying overnight.

As discussed above, the salicylhydrazide group tends to fully deprotonated to adopt the thermodynamic stable 5-5-6 chelated rings when it coordinates with transition metal ion.16 For Gel3/ 1, the Zn/L ratio is equal to the charge ratio of L^{6-} and Zn^{2+} ion, so there is good chance for the coordination assembly to occur

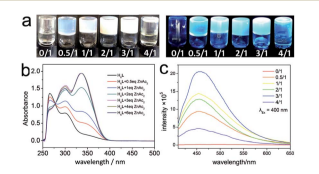


Fig. 1 (a) Photos of gelation test in different Zn/L molar ratios under daylights (left) and UV-lights (right). The UV-Vis absorption spectra (b) and fluorescence spectra (c) of the samples with diverse Zn/L molar ratio.

in the system. The liquid extrusion effect of Gel3/1 suggests the coordination bonding is too strong to maintain an open network for trapping solvent molecules. When Zn/L ratio is lower, such as 2/1, the amount of metal ion is not enough for the coordination with L^{6-} . Excess H_6L may form hydrogen bonding to participate the supramolecular assembly. Considering the Gel2/1 is stable for months, the synergistic effect of coordination bonding and hydrogen bonding in Gel2/1 benefits the gel phase stabilization. When the Zn/L ratio is lower than 2/1, such as 0.5/1 and 1/1, the resulting gels become too soft to trap the solvent molecules, which suggests the hydrogen bonding assembly is not strong enough for obtaining the supramolecular network. On the other hand, when Zn/L ratio is 4/1, the remnant zinc ion might exist as solvated ion that brings in excess free positive charged particles, which make the system become solution. Thus the colloid particles in the gel may be negative charged.

To characterize the gel system, UV-Vis absorption spectra were taken. The spectrum of H₆L (red curve in Fig. 1b) shows typical absorption bands for $\pi \to \pi^*$ and $n \to \pi^*$. Upon the addition of Zn²⁺ ion (Zn/L: 0.5/1-3/1), the charge transfer band (280-400 nm) greatly strengthen, which suggests the coordination assembly occurs for Gelx/1 (x = 0.5, 1, 2, 3). When the equivalent of zinc ion increases from 0.5 to 3 eq., the absorbance at 339 nm increases. When the zinc ion' equivalent increases to 4 eq. or 6 eq., the absorbance at 339 nm doesn't change. These studies indicate the absorption band at 339 nm should be attributed to coordination bonding. IR spectra for zinc acetate, H₆L and Gel2/1 (Fig. S4 in ESI†) were be recorded to study the coordination between H₆L and zinc ion. The 3309 $(\nu_{\rm OH})$ and 3046 $(\nu_{\rm NH})$ cm⁻¹ peaks of H₆L greatly weaken in the curve of Gel2/1, which indicates the proton loss of the H₆L in Gel2/1. By comparing with the spectra of zinc acetate, H₆L and Gel2/1 in the range of 1800–1200 cm⁻¹, the red shift of $\nu_{C=O}$ (from 1666 cm $^{-1}$ in H₆L curve to 1650 cm $^{-1}$ in Gel2/1 curve, the $\nu_{\rm C=O}$ in zinc acetate curve is at 1536 cm⁻¹) suggests the coordination between H₆L and zinc ion may occur. MALDI-MS (Matrix Assisted Laser Desorption Ionization Mass) spectrum for Gel2/1 reveals the possible self-assembly fashion. As shown in Fig. 2a, there exists wave-like spectrum line in the range of m/

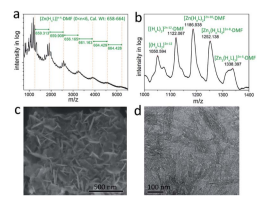


Fig. 2 (a and b) MALDI-MS spectra in the range of m/z 750–5500 (a) and 1000-1400 (b) for Gel2/1. (c and d) FESEM (c) and TEM (d) micrograph of Gel2/1.

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z 1000–5000 with regular intervals that closed to the molecular weight of $[\mathrm{Zn}(\mathrm{H}_n\mathrm{L})]^{n-4}\cdot\mathrm{DMF}$ (0 < n < 6, calc. wt: 658–664). The strongest peaks in the range of m/z 1000–1400 could be attributed to the assembly of two $\mathrm{H}_n\mathrm{L}$ (0 < n < 6) molecules with certain equivalents of Zn^{2+} ion and DMF molecule (Fig. 2b). Consequently, the self-assembly between $\mathrm{H}_6\mathrm{L}$ and Zn^{2+} ion involved in Gel2/1 is assumed to follow a step-by-step style driven by the synergistic effect of coordination assembly and hydrogen bonding. The microscopic morphology of Gel2/1 is studied by FESEM and TEM, showing thin sheets in nanoscale (Fig. 2c and d).

As shown in Fig. 1a, blue light emission could be observed by keeping Gelx/1 (x = 0.5, 1, 2, 3) sample under UV light in the dark room. By using fluorescence spectrometer, the gel samples could emit fluorescence under the excitation of 400 nm (Fig. 1c). The emission intensity increases along with the Zn/L from 0.5/1 to 3/ 1. While for the Zn/L = 4/1 sample, the emission intensity greatly weaken along with the destruction of gel phase, which indicates gelation-induced fluorescence enhanced effect. Further dynamic rheology performed on Gel2/1 suggests the gel has a certain rigidity, with storage modulus G' about 10 times larger than the loss modulus G'' within the frequency 0.01–10 Hz (Fig. S5 in ESI†). In this context, the gelation-induced fluorescence enhanced effect could be due to the rigidification upon gelation that slows down non-radiative decay processes. 10a Besides that, rheological measurements also shows a small critical strain (0.1%), indicating Gel2/1 may be an electrostatically stabilized system. To study the charge property of Gel2/1, NaCl, MgCl₂ and AlCl₃ aqueous solution were layered onto the gel phase. After leaving the testing gel in room temperature for one hour, the gel which treated with AlCl₃ aqueous solution completely dissolved into clear solution, the gel that treated with MgCl2 aqueous solution partly dissolved. While the NaCl solution seems to do not influence the gel status. In that case, the colloidal particle in Gel2/1 should be negative charged. The electrostatic force plays an important role to stabilize the gel system.

The relationship among the supramolecular structure, phase status and photoluminescence property were studied by introducing ammonia vapour. On one hand, ammonia vapour is a kind of basic gas that could influence the coordination bonding in some extent, and be easy to evaporate to allow a promising reversible modulation. On the other hand, NH₃ molecule is a good candidate for hydrogen bonding. As shown in Fig. 3a, upon exposing in the ammonia vapour, the gel (labelled as Gel^O, where subscribe "O" means original) becomes transparent light yellow solution (labelled as Solution^{C1}, where subscribe "C1" means the first cycle), and the fluorescence greatly weakens. After the NH₃ evaporates naturally, transparent gel phase reforms in two days, and the photoluminescence enhances again (labelled as Gel^{C1}). Emission spectra for Gel2/1 sample were recorded to study this phenomenon. As shown in Fig. 3b, the initial emission spectrum for original Gel2/1 (Gel^O) exhibits strong peak at 447 nm. After the phase transition completes, the emission intensity of the solution with the involvement of NH₃ (Solution^{C1}) is closed to zero (grey curve in Fig. 3b). Repeated tests for 7 cycles of Gel2/1 sample suggest that the fluorescence on and off effect is reversible along with

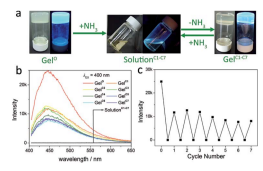


Fig. 3 (a) Scheme of the reversible gel-solution phase transition. (b) The emission spectra of Gel2/1 sample in gel-solution phase transition that induced by NH_3 gas. The colourful curves represent samples of Gel° and Gel° the grey curves represent samples of Solution C1-C7. (c) The relationship between fluorescence intensity at 447 nm and the cycle number of phase transition of Gel2/1 sample.

the phase transformation. However, it should be pointed out that the fluorescence intensity of the gel in cycle 1–7 (Gel^{C1-C7}) was lower than the original gel sample (Gel^{O}), *ca.* 51–31%, which might be due to the residue NH₃ in the gel sample.

To discuss the possible mechanism of the phase transition, the role of ammonia becomes important. It was found that the complete phase transition from gel to solution required only 5 minutes. When the Gel2/1 has been in the ammonia vapour for just 1 minute, the gel phase becomes so soft that closed to the colloidal sol phase. The UV-Vis absorption spectrum for three states, (1) the gel, (2) solution and (3) the intermediate soft colloidal sol, were recorded. As shown in Fig. S6,† the absorption spectra for state 1–3 are similar, which indicate ammonia vapour may not disrupt the coordination of H_6L to zinc ion significantly.

Furthermore, HRMS-ESI spectra for state 3 and state 4 (in this case, the initial gel solution was allowed to stay for 5 min after mixing reactants, then diluted in acetonitrile for mass study) were tested. As shown in Fig. S7† (state 3), there are anion $m/z = 1051.2930 ([H_6L \cdot H_5L]^- \text{ calc. } 1051.2904), \text{ anion } m/z =$ 1113.2063 ($[Zn \cdot H_5L \cdot H_4L]^-$ calc. 1113.2039), indicating the ligands can aggregate with each other or aggregate with Zn2+ in the presence of ammonia. Fig. S8 and S9† for state 4 suggest that small aggregates already exist in the first 5 min of gelation. There are cation m/z = 1115.2157 ([Zn·H₆L·H₅L]⁺ calc. 1115.2190), cation $m/z = 1579.4511 ([3(H_6L) \cdot H]^+ \text{ calc.})$ 1579.4544), anion m/z = 1014.2029 (z = 3) ([6Zn·5(H₃L)· $2(H_2O)$ ³⁻ calc. 1014.7386). By comparing the HRMS-ESI spectra of state 3 and 4, it is speculated that the introduction of ammonia may hinder the assembly of small aggregates into larger one. In other words, the supramolecular network in Gel2/ 1 that dealt with ammonia would be cut into pieces that no longer hold the solvent molecules. In that case, the gel-solution phase transition was observed.

To explore the application of Gel2/1, four charged dye molecules were tested by the adsorption of the xerogel sample for Gel2/1. The experimental details were described in ESI-1.3.† As shown in Fig. 4, the solution of positively charged methyl violet and methylene blue exhibit significant adsorption behavior upon the xerogel of Gel2/1, while, there is no evidence

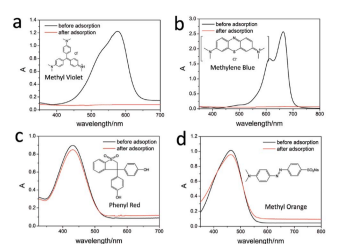


Fig. 4 The selective adsorption behavior for Gel2/1 in the presence of methyl violet (a), methylene blue (b), phenyl red (c) and methyl orange (d), respectively.

for the negatively charged phenyl red and methylene orange solution be absorbed by the xerogel. The results shown that the xerogel could significant absorb positive charged dye molecules, which is consistent with the electrolyte experiments that discussed above.

Conclusions

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In summary, a metal-organic gel based on Zn^{2+} and salicylhydrazide-derived ligand $\mathrm{H_6L}$ has been developed, showing significant gelation-induced fluorescence enhancement. Discussion around the driving force of supramolecular assembly inspires the study of modulation over material phase and photoluminescence. Ammonia vapour is found to successfully change the material phase from gel to solution. Reversible phase transition and fluorescence on–off switch were observed by the introduction and evaporation of $\mathrm{NH_3}$. The mechanism of the phase transition is discussed.

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