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Azo and imine functionalized 2-naphthols: promising supramolecular gelators for selective detection of Fe³⁺ and Cu²⁺, reactive oxygen species and halides†

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Azo and imine functionalised 2-naphthols **1–3** have been established as low molecular weight supramolecular gelators (LMWGs). Hydrogen bonding and π – π stacking between the naphthalene units contribute a weak force for gelation. The gels have sufficient mechanical strength and are responsive to multiple ions. While the gel of **1** in CH₃CN:H₂O (1:1, v/v) is responsive to Fe³⁺ and Cu²⁺ ions, under identical conditions the gel of **2** is sensitive to Fe³⁺ ions involving a gel-to-sol transition. The imine compound **3** in its gel state (DMSO:H₂O; 1:1 v/v) displays selective interaction with Fe³⁺ ions over a series of other metal ions through a color change of the gel state from yellow to dark brown. Compound **3**, in this context, is an LMWG that selectively detects Fe³⁺ and also Fe²⁺ in the presence of an oxidising agent. Oxidation of Fe²⁺ by reactive oxygen species (ROS) and its subsequent colorimetric effect on the gel of **3** finds application in ROS detection. Moreover, the Fe³⁺ ion-induced colored gel of **3** successfully senses F[–] ions visually from other halides. For real-life application, a test strip of **3** for the detection of these analytes and also its use as a rewritable display material and combinational logic gate system are demonstrated.

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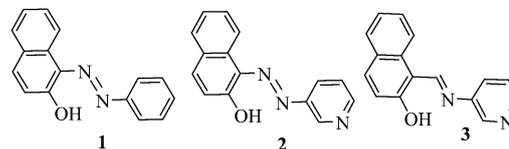
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Introduction

In recent years, the design and synthesis of stimuli responsive low molecular weight gelators (LMWGs) has been of foremost interest in molecular recognition and material science because of their wide range of potential applications in biomedicine, drug delivery and sensing *etc.*¹ Gels are viscoelastic semisolid materials formed by caging a large volume of solvent into the intermolecularly linked three-dimensional architecture of a gel matrix through several non-covalent interactions like hydrogen bonding, π – π stacking, van der Waals interactions *etc.*² Different functional entities of compounds of this class play a crucial role in establishing such non-covalent interactions leading to self organization. Of the different functional architectures available, salen-type imine and azo phenol motifs receive attention in molecular recognition because of their binding properties towards cations and anions.³ Several approaches in developing stimuli responsive organogelators based on these motifs are known.⁴ Intramolecular H-bonding and extensive

π – π stacking between the aromatic rings assist the self aggregation of these molecules. Compounds of this class also exhibit several interesting properties like AIE (aggregation induced emission), photochromism, thermochromism and solvatochromism *etc.*⁵ In order to exploit such a class of compounds in metal ion sensing through a gel–sol methodology, we report here some naphthol-based azo and imine compounds **1–3** which exhibit gelation in semi-aqueous solvents. While the gel of **1** was responsive to both Fe³⁺ and Cu²⁺ ions, the gel state of **2** was only sensitive to Fe³⁺ ions, involving a reversible gel–sol transition. The imine compound **3** showed selective visual sensing of Fe³⁺ ions *via* a noticeable color change from yellow to dark brown in the gel state.



Among the different transition metals, the sensing and detection of Fe³⁺ and Cu²⁺ ions are important because of their biological relevance. Cu²⁺ ions serve as a catalytic cofactor for a variety of metalloenzymes,⁶ though an excess of it can damage several body functions such as gastrointestinal disturbance and liver or kidney damage.⁷ On the other hand, the Fe³⁺ ion is an essential element in living cells, performing vital roles in the

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function of haemoglobin or other cellular metabolism processes.⁸ A deficiency of Fe³⁺ can lead to liver damage, anaemia, Alzheimer's disease, Parkinson's disease and cancer.⁹ Again, sensing probes for Fe³⁺ ions have often suffered from interference by Fe²⁺ or other cations.¹⁰ So, the specific detection of these metal ions is crucial. Although several fluorimetric and colorimetric sensing probes for Fe³⁺ ions have been reported in the literature,^{10,11} until now there have only been a few reports of their detection involving gel media.^{10i,j} Thus, the scope for the development of a new supramolecular gelator for the selective recognition of Fe³⁺ ions is exceedingly demanding.

Results and discussion

Synthesis

Compounds 1–3 were synthesized according to Scheme 1.¹² A diazo coupling reaction of 2-naphthol with aniline and 3-amino pyridine afforded compounds 1 and 2, respectively. On the other hand, Schiff base formation between 3-aminopyridine and 2-hydroxy-1-naphthaldehyde in dry benzene introduced compound 3 in appreciable yield.

Gelation study, aggregation and morphology of gels

The gelation tendencies of compounds 1–3 were investigated in a number of pure organic solvents and solvent mixtures. All the compounds were prone to form gels in semi-aqueous solvents rather than in pure organic solvents (Table 1S, ESI[†]). In polar solvents like THF, DMSO or DMF they remained soluble, but in the presence of water they readily formed aggregations leading to gelation. While the azo dyes 1 and 2 formed gels in CH₃CN or CH₃CN:H₂O solvent systems, the imine analogue 3 was unable to undergo self-association under similar conditions. Compound 3 formed a gel from DMSO:H₂O (1:1, v/v). There was a noticeable difference in their minimum gelation concentrations (mgc) and gel melting temperatures (*T*_{gel}).

For our study, we focused on the CH₃CN:H₂O (1:1, v/v) gels of 1 and 2, and the DMSO:H₂O (1:1, v/v) gel of 3 due to their lower mgc values for gelation. Both azo dye-based gelators 1 and 2 underwent gelation in CH₃CN:H₂O (1:1, v/v) to give red and orange colored gels, respectively. Gelator 2 exhibited low mgc and high *T*_{gel} values (mgc = 8 mg mL⁻¹ and at mgc *T*_{gel} = 42 °C) compared to 1 (mgc = 9 mg mL⁻¹ and at mgc *T*_{gel} = 38 °C) thereby highlighting the subtle role of pyridine in 2. In our opinion, pyridine forms hydrogen bonds with water and introduces

a cross-linked network in solution to form a gel. Furthermore, the gel forming abilities of the pyridyl conjugates 2 and 3 in THF:H₂O compared to the phenyl analogue 1 that remained in solution, supported the participation of pyridyl nitrogen in gelation.

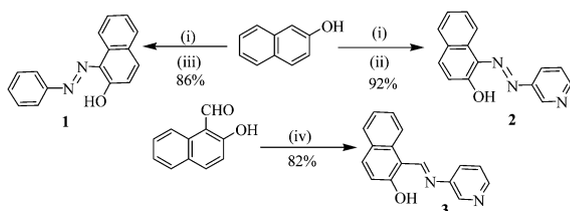
In FTIR study (Fig. 1S, ESI[†]), the stretching signals of phenolic –OH (at 3421 cm⁻¹ and 3418 cm⁻¹ for 1 and 2, respectively) became broad. The stretching frequency for the azo –N=N– bond (at 1547 cm⁻¹ and 1578 cm⁻¹ for 1 and 2, respectively) also shifted downfield and appeared as an overlapped band in the broad region 1634 cm⁻¹. This suggested the breaking of some intramolecular H-bonding with the simultaneous making of intermolecular linkages involving the solvent as mediator during gelation. In the case of 3, the stretching signal for the imine –C=N bond (1621 cm⁻¹) also moved to the higher region by 23 cm⁻¹ on changing from an amorphous to a gel state (Fig. 1S, ESI[†]) and we thereby inferred a similar situation to that proposed for 1 and 2. The aggregates are stabilized by a π -stacking force, occurring among the naphthalenes of the gelators. Intramolecular hydrogen bonding between the imine or azo nitrogen with the –OH group locks the molecular conformation into a planar arrangement and thereby results in extensive π - π stacking between the naphthalene rings, which governs the gel formation.^{4a}

All the gels exhibited thermally activated gel-sol phase transition. A sharp increase in *T*_{gel} was observed with an increase in gelator concentration (Fig. 2S, ESI[†]).

The morphologies of the gels were characterised by SEM images that revealed closely packed flakes for 1 and rod like fibrous networks for 2 and 3 (Fig. 1). In our opinion, as π - π stacking between the naphthalene units is the only stabilizing force for gelation of 1, a layer-by-layer arrangement of a large number of flakes is observed. In comparison, intermolecular hydrogen bonding involving pyridyl nitrogen atoms in 2 and 3 could possibly lead to the formation of rod-like fibers in the gel architectures.

To understand the molecular aggregation, UV-vis and fluorescence spectra of compounds 1–3 in the solution and gel states were recorded and compared. The absorptions for compound 1, noted at 314 nm, 410 nm and 482 nm in solution, were red shifted to 326 nm, 415 nm and 490 nm, respectively in the gel state. In addition, a weak shoulder at 580 nm in the gel state corroborated the aggregation. A red shift of 5 nm of the absorption peak at 410 nm during gelation indicated π - π stacking (J aggregate)¹³ between the naphthalene rings (Fig. 2a). For compound 2, significant broadening of the absorption in the region 350–470 nm and for compound 3 a red shift of the absorption band at 432 nm by 33 units in the gel states compared to their sol states (Fig. 3S, ESI[†]) supported the stacking interaction.

In the fluorescence spectra, emissions centered at 380 nm, 375 nm and 362 nm for 1, 2 and 3 in the sol states, due to their excited state keto forms through ESIPT (excited state intramolecular proton transfer), were quenched and suffered red shifts by 10 nm, 8 nm and 32 nm in the gel states, respectively (for 1: Fig. 2b, for 2 and 3: Fig. 3S, ESI[†]). This was attributed to the existence of π - π stacking among the naphthalenes in the gel matrixes.¹⁴



Scheme 1 NaOH, 0–5 °C, 15 min; (ii) 3-amino pyridine, NaNO₂, HCl, 0–5 °C, 30 min; (iii) aniline, NaNO₂, HCl, 0–5 °C, 30 min; (iv) 3-amino pyridine, benzene, reflux, 4 h.

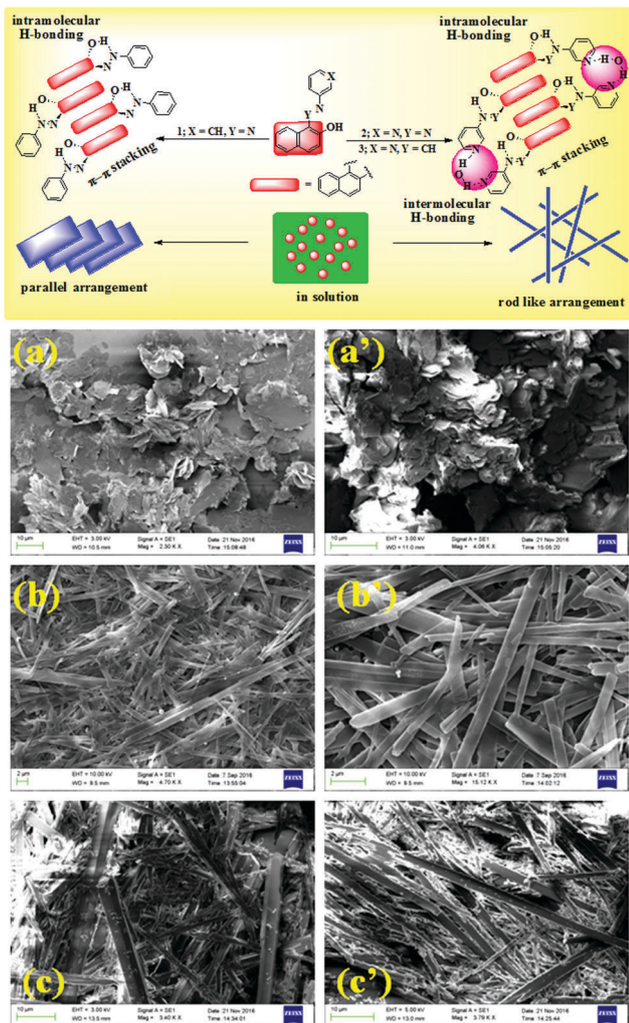


Fig. 1 Probable modes of interaction of **1**, **2** and **3** in making aggregations (top) and SEM images of xerogels of **1** (a and a'), **2** (b and b') and **3** (c and c') (bottom). [Gels were prepared from CH₃CN : H₂O (1 : 1, v/v) for **1** and **2**. For **3** it was DMSO : H₂O (1 : 1, v/v).]

Rheological study

A detailed rheological study reported the viscoelastic properties of the gels. In this experiment, the strain sweep and frequency sweep gave sufficient information about the deformation and flow characteristics of the compounds in the gel states (Fig. 2c and d). To compare the results, we prepared the three gels at a common concentration of 15 mg mL⁻¹. The change in storage modulus (G') and loss modulus (G'') of the gels was recorded against increasing strain at a fixed frequency (ω) of 1 Hz (Fig. 2c). The gels were mechanically strong ($G' > G''$) over a wide range of applied strain; however, beyond a certain strain, a sharp fall in both G' and G'' resulted in a crossover point (critical strain) and the gels were disrupted. The crossover points were detected at 3.3%, 3.7% and 31.2% of strain for the gels of **1**, **2** and **3**, respectively (Table 1). Disintegration of the gel of **3** at a higher critical strain suggested its greater strain bearing capability.^{1f} In the frequency sweep experiment (Fig. 2d) at a constant strain of 0.5%, for all the gels, both G' and G'' were

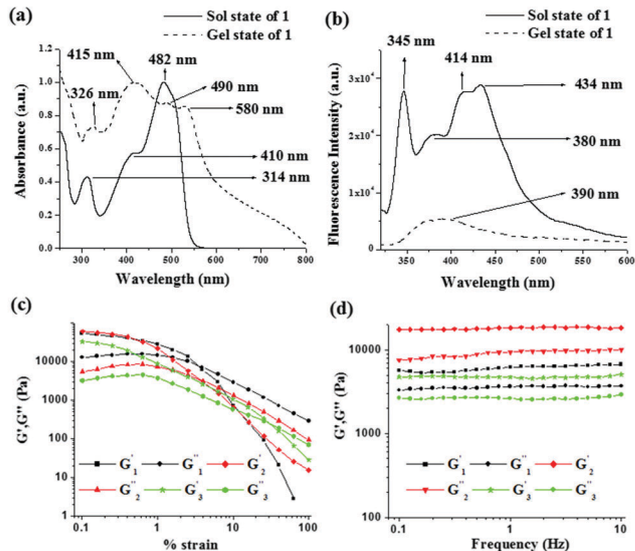


Fig. 2 Comparison of normalized UV-vis and fluorescence spectra ($\lambda_{\text{ex}} = 310$ nm) of the sol and gel states [solution was prepared in CH₃CN : H₂O (1 : 1, v/v)] (a and b) and rheological study of all the gels: (c) amplitude sweep and (d) frequency sweep experiments.

Table 1 Summary of the rheological properties of the gels of **1–3** in different solvents

Gelators ^a	Critical strain (%)	G_{av}' (Pa)	G_{av}'' (Pa)	$\tan \delta$ ($G_{\text{av}}''/G_{\text{av}}'$)
1	3.3	6063	3579	0.59
2	3.7	18 170	9122	0.50
3	31.2	4578	2655	0.58

^a Gels of **1** and **2** were obtained from CH₃CN : H₂O (1 : 1, v/v) and the gel of **3** was prepared from DMSO : H₂O (1 : 1, v/v) due to precipitation in aq. CH₃CN.

frequency independent over the entire range studied and the values of G' were always higher than those of G'' (almost doubled) at any frequency. This confirmed that all the gels were elastic semisolid and stable over the entire region studied. Importantly, the gel of **2** exhibited a higher storage modulus (G') compared to the gels of **1** and **3** (almost 3–4 times higher, Table 1). Analysis of $\tan \delta$ values ($\tan \delta = G''/G'$; 0.59, 0.50 and 0.58 for the gels of **1**, **2** and **3**, respectively) revealed that the gel of **2** exhibited higher mechanical strength (lower $\tan \delta$) compared to the other two gels.^{1g}

Metal ion responsive behavior of the gels

The stimuli responsive natures of the gels were investigated by adding aqueous solutions of different metal salts (as perchlorate salts, $c = 0.2$ M) in 1 equiv. amount on the top of the gels. In the case of **1**, only Cu²⁺ and Fe³⁺ ions disintegrated the gel and validated their visual sensing. While Cu²⁺ ions disintegrated the gel within 1 h, Fe³⁺ ions took 3 h for complete rupturing of the gel. Under identical conditions, other cations did not show any phase change (Fig. 3a).

In the case of **2**, only Fe³⁺ ions over the other metal ions examined disintegrated the gel within 1 h (Fig. 3b) and thus the gel state of **2** was more selective than the gel of **1**.

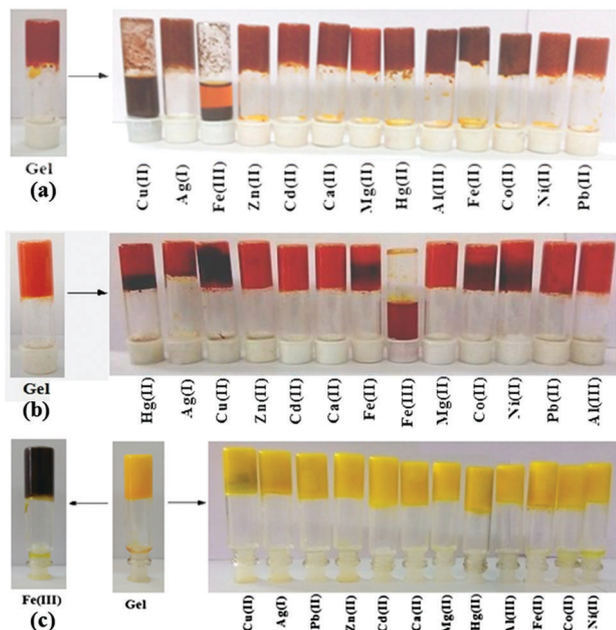


Fig. 3 Photograph showing the changes in the gel state of (a) **1**, (b) **2** and (c) **3** (at mgc) upon the addition of different metal ions.

To become acquainted with the sensitivity of the gel state for metal ion detection, 0.5 equiv. amounts of the metal ions instead of 1 equiv. amounts were used in the study and the same observations were noted. Only the gel to sol conversion was achieved in a longer time (Fig. 4S, ESI[†]). To be precise, a detailed study, as shown in Fig. 4S (ESI[†]), reveals that the gels of **1** and **2** were interactive with Cu²⁺ and Fe³⁺ ions when they were treated with the metal ions with a concentration as low as 0.02 M for **1** and 0.03 M for **2**. Below these concentrations, there was gel formation.

The recovery of the gels from Cu²⁺ and Fe³⁺ ion-induced broken gels was tested upon separate additions of EDTA, acetylacetone, and 1-dodecanethiol (Fig. 5S and 6S, ESI[†]). In both cases, EDTA retrieved the gels. In contrast, while 1-dodecanethiol recovered the gel from Cu²⁺-induced broken gel by complexing Cu²⁺ ions, under similar conditions the Fe³⁺-induced broken gel remained unaffected. Similarly, acetyl acetone only regenerated the gel from an Fe³⁺-induced broken gel through strong chelation of Fe³⁺. These observations are worthwhile for distinguishing these metal ions.

Like the gels of **1** and **2**, the metal ion responsive nature of the DMSO:H₂O gel of **3** was verified using the same metal ions. None of the metal ions ruptured the gel when added in 1 equiv. amount (as perchlorate salts, *c* = 0.2 M). Only in the presence of Fe³⁺ ions, the yellow colored gel turned into dark brown without showing any phase change (Fig. 3c). Upon addition of Fe³⁺ ions, the yellowish gel slowly changed to light brown and finally to dark brown after 24 h (Fig. 7S, ESI[†]). It is worth mentioning here that the dark brown colored metallogel was obtained instantly when the gel was prepared by adding an aqueous solution of Fe³⁺ ions to the DMSO solution of **3** by maintaining the required ratio of solvents. Thorough experiments

pointed out that the presence of 0.03 equiv. of Fe³⁺ ions (*c* = 0.003 M) was the optimum at which the gel of **3** was capable of changing its color from yellow to dark brown (Fig. 8S, ESI[†]). Fe²⁺ ions did not bring about this sharp color change in the gel.

To check whether the counter anions of the Fe³⁺-salt had any role, salts other than perchlorate, such as FeCl₃ and Fe₂(SO₄)₃, were used in the study. Observations in all cases were identical. However, when an aqueous solution of K₃Fe(CN)₆ was used, no change in the color of the gel state was observed due to the non-existence of free Fe³⁺ ions (Fig. 9S, ESI[†]). These findings not only ruled out the role of counter anions but also indicated the key function of free Fe³⁺ ions in bringing a color change to the gel state. As in the case of **1** and **2**, the yellow color of the original gel of **3** was recovered upon addition of an aqueous solution of EDTA (Fig. 10S, ESI[†]). Thus compound **3**, comprising a naphthalene-salicylideneaniline architecture, acted as a new supramolecular gelator to selectively detect Fe³⁺ ions by the naked eye from other various metal ions tested.

Since most Fe³⁺-sensors suffer from interference by Fe²⁺ ions,^{10d-f} the development of a selective, sensitive and efficient probe for Fe³⁺ ions is a great challenge. Gelator **3** solves this problem by giving a different coloration of the gel in the presence of Fe³⁺ and Fe²⁺ ions (Fig. 8S and 9S, ESI[†]).

The spectacular color change of the gel state of **3** in the presence of Fe³⁺ ions can presumably be attributed to the subtle change in the molecular structure of the gelator/ligand upon complexation of Fe³⁺ that augments the charge transfer occurring between the ligand and the metal ions.¹⁵ The SEM image of the Fe³⁺-**3** gel exhibited clusters of small sized rods (Fig. 11S, ESI[†]) which is in contrast to the morphology of the gel of **3** (Fig. 1c). Even the thermal and mechanical properties of the Fe³⁺-induced colored gel of **3** varied from those of the simple gel of **3**. A slight increase in *T*_g (4 °C) was recorded at the mgc of **3** in the presence of 1 equiv. of Fe³⁺ ions. In a dynamic amplitude sweep experiment (at a fixed frequency ω = 1 Hz) of 3-Fe³⁺ gel, the crossover point was detected at 1.99% strain (for the gel of **3** it was 31.2%) which indicated a decrease in the strain bearing capability and elastic nature of the gel in the presence of Fe³⁺ ions (Fig. 12S, ESI[†]). However, in a frequency sweep experiment (at a constant strain of 0.5%), the 3-Fe³⁺ gel was found to exhibit a higher storage modulus (*G'*) compared to the gel of **3** (almost 1.3 times higher), which suggested an increase in its mechanical strength.

A UV-vis spectral study of the gel of **3** itself and in the presence of Fe³⁺ was performed. As can be seen from Fig. 13S (ESI[†]), the absorption bands at 382 nm (assigned to the n- π^* transition of the imine bond) and 465 nm (for the keto-amine form of the Schiff base) of the gel of **3** underwent a marked change in the presence of Fe³⁺. The addition of Fe³⁺ showed the disappearance of the peak at 382 nm followed by the appearance of a 5 nm blue shifted peak at 460 nm. This observation enables us to presume that there is a difference in the population of the equilibrium forms of the imine and keto-amine forms in the gel of **3** when Fe³⁺ ions are present.

An FTIR study of compounds **1-3** in their gel states was performed in the absence and presence of the said metal ions.

As can be seen from Fig. 39S (ESI[†]), the azo motifs in **1** and **2**, which appeared at 1558 cm⁻¹ and 1578 cm⁻¹ respectively, moved to lower stretching frequencies in the presence of metal ions (for **1**: 1539 cm⁻¹ and 1545 cm⁻¹ in the presence of Fe³⁺ and Cu²⁺ ions respectively; for **2**: 1552 cm⁻¹ in the presence of Fe³⁺ ions). Similarly, in the case of **3** the stretching frequency for the imine bond at 1622 cm⁻¹ in the gel state became broad in the presence of Fe³⁺ ions and it was difficult to interpret. However, in all cases, the C–O bond stretch moved to lower regions to different extents in the presence of the metal ions (Fig. 14S, ESI[†]). This information demonstrates that the imine or azo group in the compounds participates in interaction with the metal ions either individually or collaboratively with the phenolic-OH.

Metal ion interaction in solution

UV-vis study. In order to understand the solution phase interactions, we performed UV-vis and fluorescence titrations of **1**, **2** and **3** ($c = 2.50 \times 10^{-5}$ M) with the said metal ions (as perchlorate salt; $c = 1.0 \times 10^{-3}$ M) in CH₃CN:H₂O (1:1, v/v). During titration with different metal ions, the absorption spectra of all the compounds were influenced by Fe³⁺ ions alone (Fig. 15S, ESI[†]). Upon titration with Fe³⁺ ions (up to 40 equiv.), for compound **1**, the absorption at 482 nm was gradually decreased followed by an increase in intensity of the band at 314 nm, resulting in an isosbestic point at 410 nm (Fig. 4a), which clearly indicated the complexation of Fe³⁺ ions with **1**. Except for Cu²⁺ ions (up to 5 equiv.) which brought about a weak ratiometric response (Fig. 16S, ESI[†]), other metal ions did not significantly affect the absorption spectrum of **1** (Fig. 15Sa, ESI[†]). In our opinion, such a weak interaction of **1** with Cu²⁺ ions was sufficient to disturb the gel state.

Like **1**, a ratiometric spectral change for pyridyl azo analogue **2** with Fe³⁺ was observed (Fig. 4b), where all the other metal ions were found to be non-interacting (Fig. 15Sb, ESI[†]).

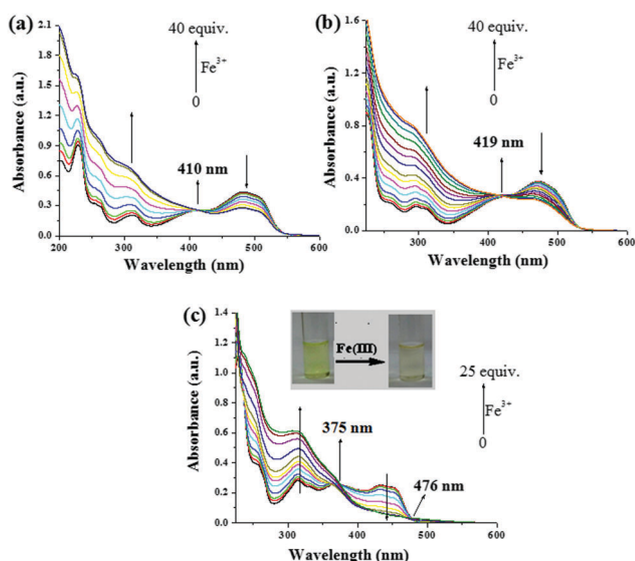


Fig. 4 Change in absorbance of (a) **1**, (b) **2** and (c) **3** ($c = 2.50 \times 10^{-5}$ M) upon addition of Fe³⁺ ions ($c = 1.0 \times 10^{-3}$ M) in CH₃CN:H₂O (1:1, v/v). Inset of 'c' represents the color change in the presence of Fe³⁺ ions.

The UV-vis spectrum of the imine analogue **3** that exhibited four absorption bands at 314 nm, 368 nm, 432 nm and 452 nm, was also influenced by Fe³⁺ ions. Upon the gradual addition of 25 equiv. amounts of Fe³⁺ ions, a ratiometric change with two isosbestic points at 375 nm and 476 nm associated with a noticeable color change from light green to a faint pale brown color was observed (Fig. 4c). Other metal ions, except Cu²⁺, did not show any considerable change in absorption (Fig. 15Sc, ESI[†]). Although Cu²⁺ brought about a change in the absorption spectra, the light green color of the solution remained unchanged (Fig. 17S and 18S, ESI[†]).

Fluorescence study. In the fluorescence study, although compounds **1** and **2** exhibited significant quenching of emission in the presence of Fe³⁺ ions, they did not show any preference for the metal ions (Fig. 19S–22S, ESI[†]). Job analysis¹⁶ revealed a 1:1 binding interaction of **1** and **2** with Fe³⁺ and Cu²⁺ ions (Fig. 23S and 24S, ESI[†]). Benesi–Hilderband plots¹⁷ from the UV and fluorescence for **1** and **2** with these metal ions also showed linearity and indicated a 1:1 binding stoichiometry (Fig. 25S–27S, ESI[†]). Further confirmation was achieved from mass analysis (Fig. 28S, ESI[†]). The detection limits¹⁸ for Fe³⁺ and Cu²⁺ ions were found to be moderate (Fig. 29S and 30S, ESI[†]). In this regard, Table 2S (ESI[†]) summarizes the findings.

Compound **3** demonstrates ESIPT (excited state intramolecular proton transfer).⁵ The enol form of salicylideneaniline **3** involves intramolecular proton transfer from the oxygen to the nitrogen atom to give the keto form.¹⁹ In CH₃CN, compound **3** exhibited two emission bands at 343 nm and 432 nm corresponding to the monomeric enol form and stacked naphthalene excimer emissions, respectively. However, in CH₃CN:H₂O (1:1, v/v) it exhibited emissions at 346 nm and 434 nm with reduced intensities for monomer and excimer emissions, respectively. Over time the monomer emission at 362 nm progressively increased, showing a noticeable color change from almost colorless (faint green color) to light green (Fig. 31S, ESI[†]). This is happened due to the slow disintegration of the aggregation of **3** and the emission reached a plateau region on standing at room temperature for 11 h (Fig. 32S, ESI[†]).

A fluorescence titration of **3** with the said metal ions was, therefore, performed 11 h after the preparation of the solution. Upon titration, Fe³⁺ ions resulted in a ratiometric change by showing a decrease in emission at 362 nm, with the simultaneous generation of a new peak at 408 nm. On the gradual addition of Fe³⁺ ions (up to 10 equiv.) the new emission at 408 nm for chelation enhanced fluorescence (CHEF) was progressively increased with a clear isoemissive point at 368 nm (Fig. 5a). Strong chelation causes intramolecular bond rotation and stops the nonradiative decay. Like Fe³⁺, Al³⁺ and Cu²⁺ ions also induced similar spectral changes but to lesser extents, as realized from the bar plot in Fig. 5b. In the event, Fe²⁺ ions did not change the emission (Fig. 33S, ESI[†]).

Fe³⁺ ions interacted with **3** in 1:2 [G:H] stoichiometric fashion, as is evident from the Job plot (Fig. 34S, ESI[†]). This was further supported by mass spectroscopy (Fig. 35S, ESI[†]). The non-linear fitting of the UV-vis and emission titration data (Fig. 36S and 37S, ESI[†]) gave values of the binding constant

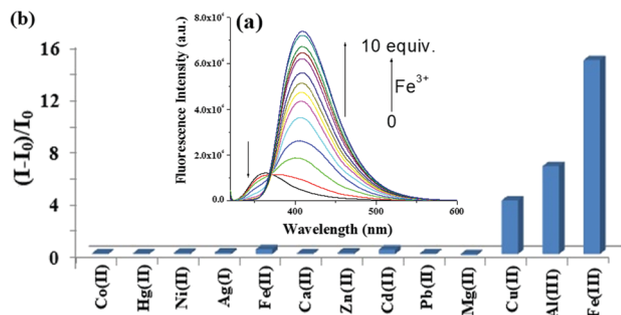


Fig. 5 (a) Fluorescence titration of **3** ($c = 2.50 \times 10^{-5}$ M) with Fe³⁺ ($c = 1.0 \times 10^{-3}$ M) and (b) change in fluorescence ratio ($\lambda_{\text{ex}} = 310$ nm) of **3** ($c = 2.5 \times 10^{-5}$ M) at 408 nm upon addition of 10 equiv. amounts of different metal ions ($c = 1.0 \times 10^{-3}$ M) in CH₃CN : H₂O (1 : 1, v/v).

which were found to be higher than those of the Fe³⁺ complexes of **1** and **2** (Table 2S, ESI[†]). The detection limit for Fe³⁺ ions was determined to be 6.67×10^{-6} M and 9.76×10^{-7} M from UV-vis and fluorescence, respectively (Fig. 36S and 37S, ESI[†]).

To understand the interaction further, ¹H NMR of **1**, **2** and **3** with the said metal ions were recorded in CDCl₃, but no measurable change in chemical shift was observed (Fig. 38S and 39S, ESI[†]). The signal for phenolic -OH in each case was retained even in the presence of the metal ions, thereby indicating the existence of strong intramolecular hydrogen bonding.

Applications of gel **3**

(a) **Recognition of ROS.** As compound **3** discriminates between Fe³⁺ from Fe²⁺ ions in both gel and solution phases, we proceeded with its practical application in detecting *in situ* generated Fe³⁺ ions *via* the Fenton reaction. In the presence of either H₂O₂ or Fe²⁺ ions, the emission spectrum of **3** was not changed significantly (Fig. 6A). But there was a noticeable change in emission when compound **3** was mixed with H₂O₂ and Fe²⁺-salt. As expected, upon the gradual addition of H₂O₂ (up to 20 equiv.) to a solution of **3** containing Fe²⁺ ions, a significant increase in emission at 408 nm with a color change from light green to colorless was observed (Fig. 6B). Similar changes in the absorption spectrum of **3** were observed (Fig. 40S, ESI[†]).

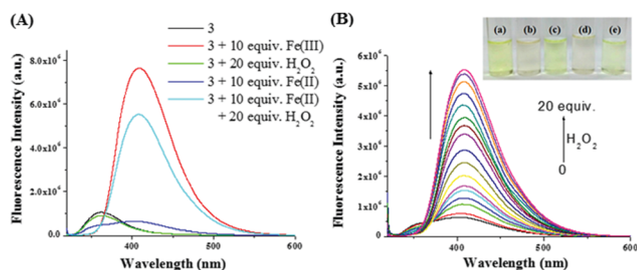


Fig. 6 Emission spectra of (A) **3** ($c = 2.50 \times 10^{-5}$ M) in the presence of Fe³⁺, Fe²⁺, H₂O₂ and a mixture of Fe²⁺ and H₂O₂ ($c = 1.0 \times 10^{-3}$ M) and (B) **3** ($c = 2.50 \times 10^{-5}$ M) + 10 equiv. of Fe²⁺ ($c = 1.0 \times 10^{-3}$ M) in the presence of 20 equiv. of H₂O₂ ($c = 1.0 \times 10^{-3}$ M) in CH₃CN : H₂O (1 : 1, v/v); Inset images show the naked eye color change of **3** [from left to right, (a) **3** and **3** in the presence of (b) Fe³⁺, (c) Fe²⁺, (d) a mixture of Fe²⁺ and H₂O₂ and (e) H₂O₂].

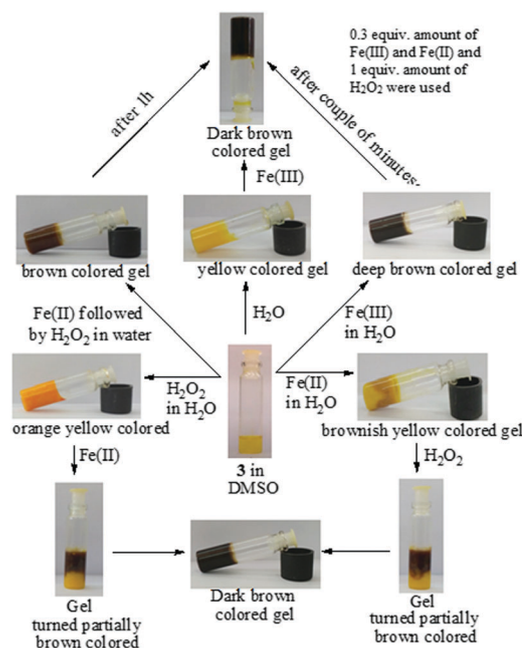


Fig. 7 Chemical responsiveness of the yellow colored DMSO : H₂O (1 : 1, v/v) gel of **3** under different conditions.

The Fenton reaction in the gel state was also performed successfully. A thorough experiment, in this context, is represented in Fig. 7. We performed several parallel experiments using Fe³⁺ and Fe²⁺-salts in 0.3 equiv. amounts ($c = 0.03$ M in H₂O). In the study, compound **3** was taken in DMSO and the gels were investigated in the presence of an aqueous solution of either Fe²⁺ ions or H₂O₂. In such cases, the gels were orange or brown colored. In contrast, the addition of an aqueous solution of Fe³⁺ ions to a DMSO solution of **3** yielded a deep brown colored gel. This was attained, in another way, when Fe²⁺ and H₂O₂ were added together to a DMSO solution of **3**. This implied the *in situ* oxidation of Fe²⁺ into Fe³⁺ and its subsequent effect on the gel in bringing about a color. To the best of our knowledge, this is the first report of utilizing the Fenton reaction condition in the gel state for discriminating Fe³⁺ from Fe²⁺ ions.

Since compound **3** successfully performed the Fenton reaction condition both in gel and solution states, we then explored it for the sensing of reactive oxygen species (ROS) which are expected to oxidize the Fe²⁺ into Fe³⁺ ions. Although numerous fluorimetric sensing probes for the recognition of reactive oxidants in solution have been reported in the literature, very few supramolecular gelators are known which have validated their visual sensing.²⁰

In this study, compound **3** (11 mg mL⁻¹) was taken in DMSO and then an aqueous solution of Fe²⁺ ions was added to it followed by the addition of different ROS like H₂O₂, TBHP, mCPBA, perborate and perchlorate (as a Na-salt), iodate and periodate (as a K-salt). The solutions were kept undisturbed for a while. Only H₂O₂ was found to be efficient in bringing about a dark brown coloration of the gel (Fig. 8). Thus, this simple experiment becomes pretty handy in the selective visual recognition of H₂O₂ from other ROS.

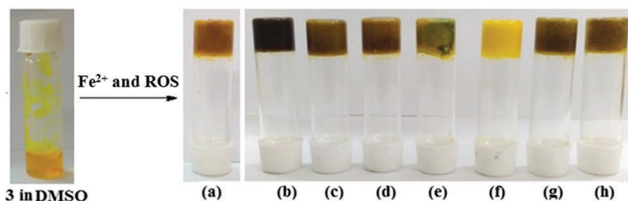


Fig. 8 Photograph showing the gel formation of **3** (11 mg mL^{-1}) in DMSO:H₂O (1:1, v/v) in the presence of (a) Fe²⁺ (0.1 equiv. $c = 0.03 \text{ M}$ in water) and Fe²⁺ with (b) H₂O₂, (c) TBHP, (d) mCPBA, (e) BO₃⁻, (f) IO₄⁻, (g) ClO₄⁻ and (h) IO₃⁻ (1 equiv. in water).

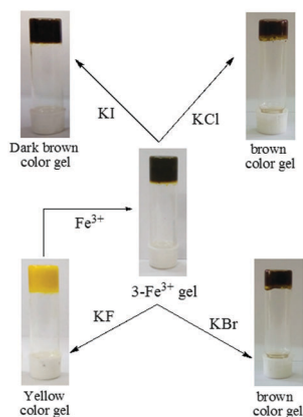
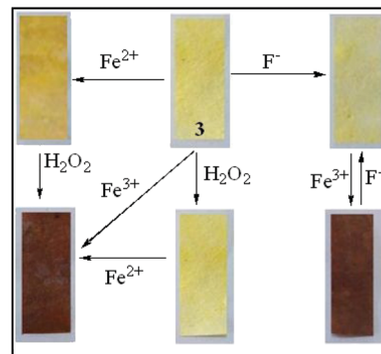


Fig. 9 Response of the **3-Fe**³⁺ gel towards KF, KBr, KCl and KI.

(b) Recognition of F⁻ from other halides. The complexation of Fe³⁺ by F⁻ ions is a well-known fact and it can be easily employed in the present case for the selective detection of F⁻ ions by Fe³⁺ impregnated gel **3**. In reality, while the dark brown colored Fe³⁺ ion impregnated metalogel was treated with different potassium halides, only F⁻ among other halides retrieved the original color of the gel of **3** by complexing the Fe³⁺ ions from the gel matrix (Fig. 9). A further addition of Fe³⁺ ions to this yellow colored gel caused a dark brown coloration.

(c) Test kit application, utilization as rewritable display material and formation of combinational logic gate. Besides this, for real-life applications in the qualitative detection of Fe³⁺, H₂O₂ and F⁻ ions, test kits were prepared by immersing filter papers in the DMSO:H₂O solution of **3** and then drying them in air (Fig. 10). When the yellow colored **3**-coated test kit was plunged deep into the Fe³⁺ solution a couple of times and dried in the air, it turned an intense brown color. Under identical conditions no color changes of the test kits were observed for Fe²⁺ or H₂O₂. Then we put the test kit into the Fe²⁺ solution first, and then into the H₂O₂. Here we also got similar deep brown colored kits. Again, the brown color kit turned to a pale yellow color in the presence of F⁻ ions. These results indicated that compound **3** can be used as a suitable practical sensor for multi-analyte recognition.

Recovery of the yellow colored kit upon fluoride treatment enabled us to establish compound **3** as rewritable display material, as shown in Fig. 41S (ESI⁺). Interestingly, writing on



Inputs			Output
Fe(III)	Fe(II)	H ₂ O ₂	
0	0	0	0
0	0	1	0
0	1	0	0
1	0	0	1
0	1	1	1
1	0	1	1
1	1	0	1
1	1	1	1

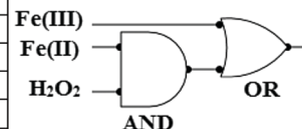


Fig. 10 Photographs of the test kits with **3** ($c = 0.001 \text{ M}$), for detecting Fe³⁺, H₂O₂ and F⁻ ($c = 0.01 \text{ M}$) (top) and representation of the color changes in the gel state and fluorescence changes of **3** based on different inputs for the combinational logic gate (bottom).

the Fe³⁺ coated brown colored strip with a brush dipped into an aqueous solution of F⁻ resulted in a yellowish written image. The written image was detectable by the naked eye, as there was a sharp color change. However, the written image could be wiped out by washing with a solution of Fe³⁺ ions and regenerated upon treatment with F⁻ solution.

Furthermore, compound **3** could also be represented as a molecular system exhibiting combinational logic gate properties based on three inputs (Fe³⁺, Fe²⁺ and H₂O₂) and one output (emission of **3** at 408 nm; see Fig. 6) (Fig. 10). In order to represent the corresponding changes in emission to a logic gate mimicking circuit, the absence and presence of Fe³⁺, Fe²⁺ and H₂O₂ were considered as inputs 0 and 1, respectively. And similarly, the output was considered as 0 in the case of no change in fluorescence intensity and 1 in the case of fluorescence enhancement at 408 nm. Among all the combinations, the three input signals (000, 010 and 001) exhibited no change in the fluorescence intensity of **3** at 408 nm, and thereby displayed the output as "OFF" with the readout signal as 0, whereas any other combination showed fluorescence enhancement of **3** at 408 nm and thereby displayed the output as "ON" with the AND logic gate, and the output of the AND logic obtained with Fe³⁺ build-up the OR logic gate. Similar logic gate mimicking could also be constructed in the gel phase, based on the same inputs and the formation of a dark brown colored gel as the output signal (readout signal as 1). Thereby compound **3** could be demonstrated to be a molecular system which could respond to a three input one output combinational logic gate mimicking circuit in solution as well as in gel states.

Conclusions

In conclusion, the gelling properties and the stimuli responsive nature of azo and imine functionalized 2-naphthols **1–3** have been investigated in detail for the first time. Compounds **1–3** exhibit gelation from various solvents. While the gel of **1** from CH₃CN/H₂O was found to be responsive to Fe³⁺ and Cu²⁺ ions, the gel state of **2** under identical conditions was only responsive to Fe³⁺ ions, *via* a reversible gel–sol phase transition. On the other hand, the imine analogue **3** turned out to be an excellent system compared to **1** and **2** for its various functions, such as (i) selective visual detection of Fe³⁺ ions through a color change from yellow to dark brown in the gel-to-gel state, (ii) visual discrimination of Fe³⁺ and Fe²⁺ ions in solution phase, (iii) selective visual recognition of ROS like H₂O₂ in the gel phase, (iv) visual discrimination of F[−] from other halides, (v) as a ratiometric fluorescence sensing probe in detecting Fe³⁺ ions, of comparable performance with the reported systems in the literature (Table 3S, ESI[†]).^{10,21}

System **3** was furthermore of practical use and in this regard, several features were established: (i) the feasibility of a Fenton reaction in both solution and gel states, (ii) the preparation of test kits for several analytes (Fe³⁺, H₂O₂ and F[−]) and (iii) the formation of a combinational logic gate system.

Experimental

Materials

2-Naphthol, 3-aminopyridine, NaNO₂, HCl and ROSs were purchased from Spectrochem and were carefully handled. Perchlorate salts of cations used in the study were purchased from Sigma-Aldrich. All solvents used in the synthesis were purified, dried and distilled as required. Solvents used in the NMR experiments were obtained from Aldrich. Thin layer chromatography was performed on Merck precoated silica gel 60-F₂₅₄ plates. ¹H and ¹³C NMR spectra were recorded using a Bruker 400 MHz instrument using TMS as the internal standard. High resolution mass data were acquired by the electron spray ionization (ESI) technique on a XEVO GS-2 QTOF Waters mass spectrometer. FTIR measurements of all the compounds and dried gels (xerogels) were carried out using a PerkinElmer L120-00A spectrometer (ν_{\max} in cm^{−1}) using a KBr cell and KBr pellets, respectively. Scanning electron microscopy (SEM) images were obtained on an EVO LS-10 ZEISS instrument. Fluorescence and UV-vis studies were performed using a Horiba Fluoromax 4C spectrofluorimeter and a Shimadzu UV-2450 spectrophotometer, respectively.

Synthesis

1-(Phenyldiazanyl)naphthalen-2-ol (1)^{12a}. 1.0 g of aniline (10.75 mmol) was dissolved in 4 mL of 3 (M) hydrochloric acid and cooled in an ice bath to 0–5 °C. 6 mL of freshly prepared 1 (M) NaNO₂ solution was added to this while maintaining the temperature below 5 °C and then immediately proceeding to the next step. 2-Naphthol (1.70 g, 11.82 mmol) was dissolved in 16 mL of 1 (M) sodium hydroxide solution, and cooled in an ice

bath with stirring for 15 min. The diazonium salt solution previously prepared was then added dropwise to it and stirred for a further 30 min under cold conditions. The pH of the reaction mixture was then adjusted to 7 and the crude product was filtered. It was then washed with water several times. Recrystallisation from EtOH–H₂O (1 : 10 v/v) afforded an orange red colored pure product **1** (2.29 g, yield 86%, mp 126 °C). ¹H NMR (CDCl₃, 400 MHz): δ 16.29 (s, 1H), 8.59 (d, 1H, *J* = 8 Hz), 7.75 (m, 3H), 7.62 (d, 1H, *J* = 8 Hz), 7.58 (t, 1H, *J* = 8 Hz), 7.50 (t, 2H, *J* = 8 Hz), 7.42 (t, 1H, *J* = 8 Hz), 7.32 (t, 1H, *J* = 8 Hz), 6.89 (d, 1H, *J* = 8 Hz); ¹³C NMR (CDCl₃, 100 MHz): 172.1, 144.6, 140.1, 133.5, 130.0, 129.6, 128.8, 128.6, 128.0, 127.4, 125.7, 124.8, 121.7, 118.5 (one carbon not found due to overlap); FTIR (KBr) ν cm^{−1}: 3421, 1617, 1496, 1447, 1206, 1142; HRMS (TOF MS ES⁺): calcd 249.1022 (M + H)⁺, found 249.1033 (M + H)⁺.

1-(Pyridin-3-ylidiazanyl)naphthalen-2-ol (2)^{12b}. Compound **2** was prepared by following the same procedure as was followed for **1**. In this case, 3-aminopyridine (1.0 g) yielded a red colored pure compound **2** (2.43 g, yield 92%, mp 136 °C). ¹H NMR (CDCl₃, 400 MHz): δ 15.78 (s, 1H), 8.85 (s, 1H), 8.44–8.43 (m, 2H), 7.97 (d, 1H, *J* = 8 Hz), 7.64 (d, 1H, *J* = 8 Hz), 7.51–7.47 (m, 2H, *J* = 8 Hz), 7.34–7.31 (m, 2H, *J* = 8 Hz), 6.77 (d, 1H, *J* = 8 Hz); ¹³C NMR (CDCl₃, 100 MHz): 171.5, 148.0, 141.4, 141.3, 140.7, 133.1, 130.7, 129.1, 128.7, 128.1, 126.1, 124.3, 124.1, 121.8 (one carbon not found due to overlap); FTIR (KBr) ν cm^{−1}: 3418, 2922, 2851, 1619, 1507, 1213, 1155. HRMS (TOF MS ES⁺): calcd 250.0975 (M + H)⁺, found 250.0961 (M + H)⁺.

1-((Pyridin-3-ylimino)methyl)naphthalen-2-ol (3)^{12c}. 2-Hydroxy-1-naphthaldehyde (1.46 g, 8.50 mmol) and 3-amino pyridine (1.0 g, 10.63 mmol) were refluxed in dry benzene for 4 h. The progress of the reaction was monitored by TLC. After completion of the reaction, the benzene was evaporated off completely, and the product was washed with diethyl ether several times to give the pure compound **3** in 82% yield (1.73 g, mp 184 °C). ¹H NMR (CDCl₃, 400 MHz): 14.95 (s, 1H), 9.48 (s, 1H), 8.68 (s, 1H), 8.57 (d, 1H, *J* = 4 Hz), 8.18 (d, 1H, *J* = 8 Hz), 7.89 (d, 1H, *J* = 8 Hz), 7.79 (d, 1H, *J* = 8 Hz), 7.70–7.67 (m, 1H), 7.60–7.56 (m, 1H), 7.44–7.39 (m, 2H), 7.20 (d, 1H, *J* = 8 Hz); ¹³C NMR (CDCl₃, 100 MHz): 166.1, 158.6, 147.6, 143.6, 142.9, 136.4, 132.8, 129.4, 128.2, 127.7, 124.0, 123.8, 120.5, 119.1, 109.3 (one carbon not found due to overlap); FTIR (KBr) ν cm^{−1}: 3414, 3048, 2921, 1621, 1561, 1482, 1329, 1182, 1022; HRMS (TOF MS ES⁺): calcd 249.1022 (M + H)⁺, found 249.1009 (M + H)⁺.

Gelation test and SEM imaging

The required amounts of the compounds were first dissolved in suitable organic solvents and slightly warmed to form a homogeneous solution. The required amount of water was added to the solution and it was kept at room temperature undisturbed. Gel formation was tested by the inversion of the vial method. A sample of gel for SEM imaging was dried under vacuum and then coated with a thin layer of gold metal.

Determination of gel–sol transition temperature (*T*_{gel})

The gel-to-sol transition temperature (*T*_{gel}) was defined as the temperature at which the gel melted and started to flow.

T_{gel} was measured by the dropping ball method. In this test, a small glass ball was carefully placed on the top of the gel to be tested, which was present in a test tube. The tube was slowly heated in a thermostated oil bath until the ball fell to the bottom of the test tube. The temperature at which the ball reaches the bottom of the test tube is taken as the T_{gel} of that system.

General procedures for fluorescence and UV-vis titrations

Stock solutions of the compounds were prepared in $\text{CH}_3\text{CN}/\text{H}_2\text{O}$ (1:1, v/v) at a concentration of 2.50×10^{-5} M. Stock solutions of metal ions were also prepared in the same solvent at a concentration of 1.0, put in the cuvette and different metal ions were individually added in different amounts. Upon addition, the change in emission of the compounds was recorded. The same stock solutions were used to perform the UV-vis titration experiments in the same way.

Binding constant determination¹⁷

A Benesi-Hildebrand plot was adopted to determine the binding constant value using the expression: $I_0/(I - I_0) = [\epsilon_{\text{M}}/(\epsilon_{\text{M}} - \epsilon_{\text{C}})](K_{\text{a}}^{-1}C_{\text{g}}^{-1} + 1)$, where ϵ_{M} and ϵ_{C} represent the molar extinction coefficients for the receptor and the complex, respectively, at a selected wavelength, I_0 denotes the emission intensity of the free receptor at that specific wavelength and C_{g} is the concentration of the guest. The measured absorbance $I_0/(I - I_0)$ as a function of the inverse of the guest concentration fits a linear relationship, indicating 1:1 stoichiometry of the receptor-guest complex. The ratio of the intercept to the slope was used to determine the binding constant K_{a} . Instead of fluorescence intensities, the respective absorbance of the receptor was also used to obtain the binding constant values.

Calculation of detection limit¹⁸

The detection limit was calculated using the fluorescence and UV-vis titration data. The emission and absorbance of the compounds were measured 5 times, and the standard deviation of blank measurement was achieved. To obtain the slope, emission intensities and absorbance values were plotted against concentrations of metal ions. The detection limits were calculated using the equation: detection limit = $3\sigma/k$, where σ is the standard deviation of blank measurement, and k is the slope.

Conflicts of interest

There are no conflicts to declare.

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