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Supramolecular gels in cyanide sensing: a review

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Supramolecular low molecular weight gels have potential in many areas from medicine to optoelectronics. Of the different applications, visual sensing of chemical analytes using gels is a fairly new concept in materials chemistry research. Among various analytes, cyanide is considered the most threatening to the environment and human life because of its acute toxicity. Generally, the detection of cyanide involves the use of sophisticated and expensive instrumentation, which complementary entails highly trained personnel to operate it. In contrast, a gel-based visual detection technique provides an advantage as it is simple, cost-effective, and instrument-free whilst the sensing event can be executed either by observing a phase transformation or by a naked eye detectable color change of the gel. Further, as the gel formation requires a high concentration of gelator $(\geq 10^{-3}$ M), for the same chemical compound, the gel phase interactions often exhibit better selectivity for the ionic guests than in the solution state. Despite these facts, limited attempts have been made to synthesize gel-based cyanide sensors. In this review, we discuss an up-to-date summary of various reports on cyanide responsive gels emphasizing the approaches, design principles, and reaction mechanisms. We also highlight the advantages, limitations, and challenges as well as the necessity of further exploration of gels in this domain. **REVIEW Symptons**
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Introduction

Low molecular weight supramolecular gels are important soft materials that draw attention in both academic and industrial research. These are formed by the self-assembly of small organic

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molecules leading to network structures that immobilize solvents. Despite having large amounts of liquids, these materials behave as viscoelastic solids.1–7 The solid nature arising from the network is maintained by the tandem effects of various weak non-covalent interactions, such as hydrogen bonding, π -stacking, ionic interactions, hydrophobic interactions $etc.⁸⁻¹⁵$ As the interactions are individually weak and reversible, it is possible to tune and control the gel properties, which enables one to explore them in many areas including catalysis, tissue engineering, and environmental remediation.15–29

Nowadays, there is a great deal of interest in designing 'smart gels', where the 'smart' nature arises from excellent responsiveness of the gels towards various external stimuli such as pH, light, redox, ions, ultrasound, mechanical stress $etc.³⁰⁻⁴²$ Exposure of gels to external stimuli results in changes in molecular level interactions that bring various macroscopic changes in properties like the shape and mechanical and optical properties of gels. $43-55$ Stimuli responsiveness notably extends the scope of applications of gels in the fields of sensing, actuators, drug delivery, optoelectronics etc.⁵⁶⁻⁷⁹ Of the various stimuli, ionic analytes dramatically change the chemical as well as optical properties of $gels⁸⁰⁻⁸⁸$ and thus there is immense scope for developing gelation-based visual sensors.⁸⁹⁻¹⁰⁰ Exploitation of gels in sensing involves the interaction of the externally added ionic analytes with the gelator molecules, which produces multiple responses of gelators (Fig. 1). The most common is the destruction of the network structure, which results in gel-to-sol conversion. Alternatively, upon interaction, a change in the gel network that leads to a gel-to-gel transition can occur. Thus ion-responsive gels deserve **Proview** Proview Proview Proview Proview and internalitie scheme.

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Fig. 1 Cartoon representing different modes of cyanide sensing by supramolecular gels.

attention not only for their attributes in visual detection of ionic analytes but also to adapt their material properties.

Out of several ionic analytes, sensing and detection of cyanide (CN-) ions demands considerable attention as they are considered to be the most threatening for the environment and human life. $101-103$ Cyanide is mostly produced as industrial waste. It is extensively used in the textile, paper, and plastic industries as well as in gold and silver extraction processes.^{104,105} Cyanide can also be released from biological processes of bacteria, fungi, and algae and even from human activity such as cigarette smoking.¹⁰⁵⁻¹⁰⁷ However, CN⁻ is extremely toxic to the human body. It forms a stable complex with cytochrome c oxidase leading to inhibition of enzyme activity and thereby decreases the oxygen supply to the cell, which ultimately leads to cellular asphyxiation.^{108,109} Accumulation of CN⁻ causes disorder in the nervous system and respiratory problems and eventually leads to death within a few minutes.^{110,111} By considering the adverse effect of cyanide on human health, the World Health Organization (WHO) has fixed the maximum acceptable level of cyanide concentration in drinking water at 0.2 ppm.¹¹² Therefore, sensing and detection of cyanide ions is of utmost importance.

Generally, detection of ionic analytes is performed in the solution state at a concentration of $\leq 10^{-5}$ M by using different instrumental techniques, which show changes in absorbance, emission, surface plasmon resonance, electrochemical properties, circular dichroism etc.¹¹³⁻¹²⁶ Most of these detection techniques involve the use of sophisticated and expensive instruments and complementarily entail highly trained personnel to operate them. On the contrary, a gel-based visual detection technique provides an advantage as it is simple, cost-effective, and mostly instrument-free (in some cases a normal UV-light source is used), whilst the sensing event can be executed either by observing a phase transformation or by a naked eye detectable color change of the gel. Further, gel formation requires a high concentration of gelator ($\geq 10^{-3}$ M). In many cases, for the same chemical compound, the gel phase interactions exhibit better selectivity for the ionic guests than in the solution state.¹²⁷⁻¹³¹

Scrutiny of the literature reveals that although several artificial receptors have been synthesized for sensing of CN⁻ ions in solution, $132-148$ very few design-based molecules are explored

for their visual detection involving supramolecular gels. This is due to a lack of understanding of the gelling behavior of gelators with proper structural parameters.^{16,149-158} Additional trouble relates to the correct use of CN⁻-specific binding sites in the gelators to show selectivity in the sensing process. In the context of the application of self-assembled supramolecular structures, a number of reviews are available on gel-based visual detection of ionic analytes. $^{89-100}$ However, discussion on CN $^-$ responsive gels is hardly incorporated. As the use of gels for visual detection of CN^- is a fairly new phenomenon, a detailed and up-to-date summary of work in this domain is highly demanding. From this perspective, the current review focusing on design-based CN⁻responsive supramolecular gels is well-timed and long overdue.

The design strategy for CN^- -responsive LMWGs

Design of small molecule-based anion responsive gelators is an important aspect of supramolecular chemistry. Anion responsive gelators typically contain an anion binding site. As the anions are basic in nature, two types of interactions are common in the gels: electrostatic interactions with oppositely charged centers (e.g. metal ions) and hydrogen bonding interactions with several functionalities such as hydroxyl, urea, amide, sulphonamide etc.^{83,86,93,159-164} However, unlike other anions such as F^- and ACO^- , $CN^$ behaves as a good nucleophile.¹³⁴ However, its basicity and nucleophilicity strongly depend on the nature of the working solvent. Thus, depending upon the gelation conditions and binding sites, CN⁻-responsive gels can exhibit different types of interaction. Apart from the binding site, the rest of the gelator's backbone is manipulated with hydrophobic or hydrophilic groups of various kinds^{11,152,165–174} (e.g., cholesterol, long alkyl chains, amino acid derivatives, aromatic π -surfaces etc.) that assist self-aggregation of the molecule in solution.

Depending on the nature of the interaction, CN⁻-responsive gelators are broadly classified into three categories:

- (i) Metal-anchored CN⁻ sensory gels
- (ii) H-bonding motif-based CN⁻ sensory gels
- (iii) Reaction-based CN⁻ sensory gelators

Herein, we summarize these three categories by giving up-todate details. In the discussion, advantages, limitations, challenges, and future possibilities of cyanide-responsive LMWGs are also highlighted.

(i) Metal-anchored CN^- sensory gels

Organic ligands which upon coordination of metal ions show gelation involving various weak forces belong to the class of metal-anchored gelators. Gels derived from these gelators are of growing interest in materials chemistry.^{41,51,96,174–176} In the present context, metal-anchored gelators draw attention because of the presence of metal ions that exhibit electrostatic interactions with anions. In designing these gelators, suitable metal ion binding sites are incorporated in the structural backbones. Generally, pyridine, acylhydrazone and salicylimine segments, etc. are introduced as metal ion binders. When cyanide is added externally, instead of

Fig. 2 Schematic representation of two different approaches (Type I and Type II) of cyanide sensing involving metal anchored gels associated with changes in emission.

interacting with the gelator, cyanide binds to the metal ion, leading to scavenging of the metal ions from the gel matrix. Depending upon the nature of the gelators, two different scenarios may appear (Fig. 2). In one case, the compound is a nongelator, but forms a gel in the presence of suitable metal ions. The scavenging of the metal ions by cyanide ions from this gel results in the collapse of the gel network, leading to a return to the solution state. In another case, if the gelator is capable of forming a self-supported gel even in the absence of the metal ion, metal displacement by cyanide drives the system to achieve the original attributes of the gelator. Consequently, a gel-to-gel transition is accomplished with a change in physicochemical properties such as the color, emission, thermal stability, mechanical properties, etc. of the gel (Fig. 2). Interestingly, the selectivity of metal-anchored gels towards cyanide can be controlled just by altering the nature of the metal ions.

Typical examples of Type I of Fig. 2 are compounds 1a and **1b** reported by the Damodaran group (Fig. 3). 177 Both these mono-N-oxide pyridyl amides behaved as nongelators in water; however, stable gelation was achieved for both 1a and 1b in the presence of either $ZnCl₂$ or $CdCl₂$ (1:2 metal: ligand ratio). Single-crystal X-ray diffraction studies confirmed that while coordination of the metal ions involved the pyridyl nitrogens, the N-oxide moieties were engaged in hydrogen bonding with the amide groups in both gels. Although all the metallogels displayed almost similar morphology (needle-type fibrous) under a scanning electron microscope, they behaved differently when exposed to various anions. The anion-responsive nature of the metallogels significantly depends upon the nature of the metal ion present in the gel architecture. While the Zn-gels of both 1a and 1b showed multiple responses towards CN^{-} , F^{-} and I^- , the Cd-gels of both 1a and 1b selectively recognize CN^-

Fig. 3 (a) Chemical structures of compounds 1a, 1b and 2. (b) Change in the emission of the $2\cdot$ Hg²⁺ gel (HgG) in the presence of CN $^-$. The inset presents the color change of the ${\bf 2} \cdot$ Hg $^{2+}$ gel (HgG) in the presence of CN $^-$. Reproduced from ref. 178 with permission from the Royal Society of **Chemistry**

ions, exhibiting gel-to-sol transitions. Halides such as F^- , Cl^- , Br⁻ and I⁻ did not affect the gel network, whilst CN⁻, being a stronger ligand, scavenged the Cd^{2+} -ions, making the gelator molecules free from the polymeric gel network and resulting in sol formation (the concentration of all anions was 2 equiv.).

Unlike compounds 1a and 1b, Yin et al. reported a compound, 2 (an example of Type II in Fig. 2), which was capable of self-assembling into a stable supramolecular polymer material in DMSO without the assistance of any metal ions.¹⁷⁸ The gel formation was primarily driven by intermolecular hydrogen bonding involving the hydroxyl groups and the van der Waals forces exerted by the long alkyl chains. Such assembled structures were further stabilized by the acylhydrazone group, which locks the molecular conformation by forming intramolecular hydrogen bonding, which in turn facilitates π - π stacking of the phenyl rings. In the structure, the acylhydrazone functionality additionally served as a metal-binding site. Diffusion of Hg^{2+} ions into the gel network resulted in the conversion of the organogel into a 2Hg^{2+} metallogel in which the emission of the ligand at 475 nm was almost quenched. The Hg^{2+} ion binding by the acylhydrazone of 2 was confirmed by infrared resonance spectra, which showed shifting of the stretching frequencies at 1583, 3192, and 1639 $\rm cm^{-1}$ for the –C $=$ O, –NH and –C $=$ N groups to 1581, 1641, and 3197 cm^{-1} , respectively. It was proposed that Hg^{2+} binding involving the -C=O, -NH and -C=N groups induced charge transfer and quenched the fluorescence of the Hg-gel. This Hg-gel was further employed in anion sensing involving competitive binding interactions between metal ions, anions, and gelators. Among various anions $(F^-, Cl^-, Br^-, I^-, I^+)$ AcO⁻, H₂PO₄⁻, HSO₄⁻, N₃⁻, SCN⁻, S²⁻, ClO₄⁻ and CN⁻), the Hg-metallogel showed strong yellow fluorescence selectively in the presence of CN⁻ ions and thereby executed its visual sensing in the gel–gel state (Fig. 3). The gradual addition of CN- (0–0.8 equiv.) to the 2Hg^{2+} gel resulted in demetallation and restoration of the

original spectra of organogel 2. The similar microstructure of the CN--treated metallogel to the original DMSO gel of 2 again confirmed the CN⁻-induced demetallation reaction. The detection limit of the 2·Hg²⁺ gel for CN $^-$ was reported as 3.72 \times 10 $^{-9}$ M, quite a lot lower than the standard of the WHO.

It is worth mentioning that acylhydrazones are widely used for synthesizing aggregation-induced fluorescent gels. $93,179-181$ The five-membered ring formed by the intramolecular hydrogen bonding of the acylhydrazone locked the molecular conformation in a planar arrangement and enables the aromatic groups to stack with one another in a directional way. Such locking of molecules increases the rigidity and thereby inhibits fluorescence quenching by restricting intramolecular rotation as well as inhibiting photo-induced electron transfer. Consequently, an increase in the gelator concentration shows significant enhancement of the emission of the gel compared to the monomeric solution. Furthermore, a number of metallogels can be prepared from such a self-supported gelator by introducing various metal ions inside the gel network. Depending upon the nature of the metal ions, these metallogels can exhibit various morphology, photophysical properties and anion responsiveness during gel– gel transitions. Such systems are attractive in developing multi analyte optical sensor arrays.¹⁸²⁻¹⁸⁵ Provide Materials Chemistry Frontiers

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Lin et al. introduced gelator 3 containing the same acylhydrazone moiety as a cation binder and a quinoline unit as a better aromatic capping group and fluorescence signalling unit than the phenyl of 2 (Fig. 4).186 The organogel of 3 in DMF showed aggregation-induced strong brilliant blue emission. In the presence of 0.5 equiv. of various metal ions such as Cu^{2+} , Fe^{3+} , Hg^{2+} , and Cr^{3+} , the strong emission at 500 nm was almost quenched and the corresponding non-fluorescent metallogels were obtained. On the other hand, the

Fig. 4 (a) Structure of gelator **3**. (b) Change in emission of the $3 \text{ } Cu^{2+}$ gel (CuG, 0.8% in DMF) in the presence of CN^- and other anions (F⁻, Cl⁻, Br⁻, I^- , AcO⁻, H₂PO₄⁻, N₃⁻, SCN⁻, ClO₄⁻ and S²⁻). (c) Change in the color of different metallogels in the presence of 1 equiv. of various anions. Reproduced from ref. 186 with permission from the Royal Society of Chemistry.

diffusion of Zn^{2+} ions into the organogel of 3 resulted in an almost 40 nm red shift of the emission spectra and the color of the gel changed to yellow. When the gels were treated with anions, while the 3 Cu^{2+} and 3 Fe^{3+} gels selectively responded to CN^- and recognized it via fluorescence turn-on, the Hg²⁺ and Cr³⁺-gels of 3 were nonresponsive to the same. Rather they showed fluorescence emission in the presence of SCN^- and S^{2+} ions, respectively. On the other hand, while I $^-$ quenched the strong yellow emission of the 3 \cdot Zn $^{2+}$ gel by the heavy atom effect, CN^- complexation with Zn^{2+} retrieved the original blue color of the gel. Importantly, the $3 \cdot Cu^{2+}$ and $3 \cdot Fe^{3+}$ gels could selectively sense CN⁻ ions from a mixture with other anions (multi-analyte conditions) with limits of detection of 1.0 \times 10⁻⁷ M and 1.0 \times 10⁻⁵ M, respectively. Based on these observations, a fivemembered sensor array was created which could identify different anions in water with high accuracy and sensitivity. Moreover, these metallogels were explored in the construction of erasable security display materials. Materials Chemistry Frontiers

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Sun et al. undertook a different strategy in developing CN⁻responsive gels. They incorporated two different metal ions inside the gel network, so that if CN^- could scavenge one metal ion then the gel state might still persist in the presence of the second metal ion (Fig. 5). They introduced *L*-glutamic acid Schiff base 4, which was incapable of forming a gel in a variety of organic solvents (Fig. 6).¹⁸⁷ However, stable yellow and green colored gels were obtained in DMSO either by direct treatment with $Zn(OAc)₂$ and $Cu(OAc)₂$, respectively, or by treating the solutions obtained with $\text{Zn}(\text{ClO}_4)_2$ and $\text{Cu}(\text{ClO}_4)_2$ with NaOAc. The Zn₄ and Cu₄ metallogels showed different responses under UV-light. While the Zn-4 gel exhibited strong, brilliant blue fluorescence at 457 nm, the Cu-4 gel was non-fluorescent. Following these observations, they prepared a green-colored Zn–Cu composite gel $(Zn^{2+}: Cu^{2+}: 4 = 1:1:1)$ either by adding Cu^{2+} to the Zn-4 gel or by diffusing Zn^{2+} into the Cu-4 gel. However, the properties of the Zn–Cu \cdot 4 gels were invariant to the sequence of metal ion addition. The Zn–Cu-4 gel could recognize CN⁻ ions with specific selectivity over other anions by exhibiting a naked eye detectable color change from green to yellow, and a change from a non-fluorescent to a blue colored gel under UV-luminescence. In this process, the detection limit of the Zn–Cu \cdot 4 gel for CN $^{-}$ ions was 1.6 \times 10 $^{-6}$ M. From UV-vis, fluorescence, and mass spectra it was confirmed that the properties of the CN⁻⁻treated Zn-Cu-4 gel were identical to

Fig. 5 Cartoon representing the mechanism of cyanide sensing involving a two metal ion composite gel.

Fig. 6 Structures of compounds 4 and 5.

those of the original Zn \cdot 4 gel. They suggested that CN⁻ ions competitively bound to Cu^{2+} to form stable $[Cu(CN)_x]^{n-}$ complexes, while the Zn^{2+} remained coordinated with 4 and maintained the gel structure. Further addition of Cu^{2+} again enabled the restoration of the non-fluorescent Zn-Cu-4 gel. An off-on-off fluorescence switch was constructed by altering the addition of CN^{-} and Cu^{2+} , which allowed detection for at least four cycles without any substantial loss of the fluorescence efficiency of the Zn–Cu-4 gel.

A similar method was followed by Lin et al. to explore gelator 5 as a CN^- sensor in gel-gel mode (Fig. 6).¹⁸⁸ Unlike 4, compound 5 formed stable organogels in various solvents such as dimethyl sulfoxide, propanol, ethanol, n-butanol, and iso-amyl alcohol. Among these solvents, 5 showed the highest gel-to-sol transition temperature and lowest minimum gelation concentration (0.4% w/v, 10 mg mL^{-1} = 1%) in ethanol. On the addition of calcium perchlorate to the ethanol gel of 5, a stable $Ca²⁺$ -coordinated metallogel was formed, accompanied by strong brilliant blue coloration under UV-light. Initially, they examined the competitive coordination of Ca^{2+}/Cu^{2+} with 5 by adding Cu^{2+} to the Ca-metallogel, which turned the fluorescent Ca-metallogel into a non-fluorescent Ca–Cu-metallogel through replacement of the Ca^{2+} from the acylhydrazone binding core. This Ca-Cu-metallogel showed a selective response for $CN^$ among various anions by exhibiting turn on emission of the gel. It was suggested that, because of the Group 11 element, Cu^{2+} showed stronger coordination ability with acylhydrazone than that of group 2 element Ca^{2+} and replaced it from the metal binding site. Further addition of CN⁻ showed higher affinity for $Cu²⁺$ ions and scavenged them from the acylhydrazone binding core, which allowed Ca^{2+} ions to engage with the acylhydrazone group and restored the brilliant blue emission of the gel.

Metal coordination followed by demetallation using CN^{-} is apparently an indirect method of sensing where CN⁻ ions do not interact with the original gelator. Although this could be an effective strategy for CN⁻ sensing with a very low detection limit, however, the reported systems encountered several drawbacks. The success of such gels considerably depends on many factors such as the self-assembly ability of the gelators in the presence of chemical guests, the role of the metal ions, the nature of the anions taken in the study, the concentration of the chemical analytes, etc. Metallogels can show additional interactions with other anions as well. For instance, the Cd-gels of both 1a and 1b developed by the Damodaran group, apart from CN⁻, also showed a gel-to-sol phase transition in the presence of a very high concentration of I^- (5 equiv.).¹⁷⁷ Hence, these gels are

effective in CN⁻ sensing only at a low concentration of cyanide. Similarly, the Zn-Cu-4 metallogel additionally showed interactions with S^{2-} and Cys and recognized them by fluorescence quenching.¹⁸⁷ However, the major issue is that the construction of CN⁻ responsive metallogels is really very complicated and often relies on trial-and-error methods. For example, gelator 2 was also capable of forming a metallogel with $Fe³⁺$. However, unlike the Hg²⁺-gel, the Fe³⁺-gel remained silent for CN⁻ ions but exhibited strong emission with $H_2PO_4^{-178}$ In the same line, gelator 3 reported by Lin et al. could form stable gels in the presence of a number of metal ions such as Cu^{2+} , Fe^{3+} , Hg^{2+} , Cr^{3+} , and Zn^{2+} , of which only the $3 \cdot Cu^{2+}$ and $3 \cdot Fe^{3+}$ gels showed selective interactions with CN^{-186} While the $3 \cdot Zn^{2+}$ gel showed interactions with both CN $^-$ and I $^-$, the Hg²⁺·3 and Cr³⁺·3 metallogels were non-interactive with CN^- and showed responses to SCN^- and S^{2-} , respectively. Recently, the same research group introduced compound 6, which could form a series of organogels in the presence of various metal ions (Fig. 7).189 Among the metallogels, only the Al–Fe and Al–Cu gels could execute selective visual detection of CN⁻ by fluorescence turn-on. Other metallogels either remained silent in the presence of cyanide or showed multiple responses with anions. The same is also true for tripodal gelator 7, for which only the $Ni²⁺$ -gel behaved as a cyanide sensor (Fig. 7).¹⁹⁰ Further, it is not always true that gels **Paview** Published in CX" sensing only at a low conventration of cyanide, containing the same metal tone was between big statistical Sometical Sometical Sometical Sometical Sometical Sometical Sometical Sometical Sometica

Fig. 7 (a) Structures of compounds 6 and 7. (b) Gel-based sensor array of metallogels of 6 in the presence of various anions. Reproduced from ref. 189 with permission from the Royal Society of Chemistry.

containing the same metal ions would show similar responses towards cyanide. The responsive behavior of these gels entirely depends on the nature of the coordination site and hence the outcome is quite unpredictable. For example, while the Hg^{2+} -gel of 2 validates selective visual sensing of cyanide, the same metallogel of 3 was nonresponsive to CN⁻. A similar comparison can be made by considering the contrasting behavior of the Cu^{2+} -metallogels of 3 and 7. In this context, the detail of cyanide sensing by metal anchored gels is provided in Table 1. Such unpredictable and inconsistent performance makes it complicated and more difficult to design cyanide sensing metallogelators. Hence, there was a real need to consider other approaches for cyanide sensing which allowed design of new molecules that interact directly with cyanide ions either through hydrogen bonding or via a chemical reaction.

(ii) H-Bonding motif-based CN⁻ sensory gels

H-Bonding motif-based cyanide sensory gels typically contain various hydrogen bond donor functionalities such as hydroxyl, amide, sulphonamide, urea, acylhydrazone etc. on the gelator skeleton. During the interaction, they can form strong intermolecular hydrogen bonds with the incoming cyanide ions and produce changes in the gel properties (Fig. 8). Again, cyanide sensing can induce either a gel-to-sol or a gel-to-gel phase transformation, which is unpredictable. Hydrogen bonding with CN⁻ often leads to deprotonation of the corresponding donor hydrogen, which in turn produces optical changes. In order to achieve a better spectroscopic response during the sensing process, the hydrogen bonding groups can be connected directly with aromatic surfaces, so that the effective π -conjugation length can be increased after deprotonation. Apart from the H-bonding core, the rest of the gelator design follows similar approaches as discussed earlier.

Compound 8 is an example of a hydrogen bondingfunctionalized cyanide-responsive gelator (Fig. 9), which was capable of forming stable gels in organic solvents like ethylene glycol and benzyl alcohol as well as in a mixture solvent like DMSO/water $(8:2, v/v).^{191}$ Both the intermolecular H-bonding involving the amide groups and the π - π interactions between the aromatic moieties were responsible for gel formation. The DMSO–water gel was comprised of entangled and twisted fibers of several micrometers as observed by a scanning electron microscope. It could also form a stable gel with CN⁻ with a fibril structure. The CN⁻-gel showed lower thermal stability than the original gel of 8. Hence, the DMSO–water gel of 8 was capable of detecting CN⁻ ions in a gel-gel fashion by exhibiting a change in the gel properties. However, the gelator simultaneously showed similar changes in the presence of other anions like F^- , AcO⁻ and H₂PO₄⁻, and hence the interaction with CN⁻ was not selective. By correlating proton NMR and solution phase interactions studies, it was suggested that solvation of these anions in the prescribed solvent made it a less efficient competitor for the gelator –NH groups and partly influenced the intermolecular hydrogen bonding so that the gel maintained its structure.

To make the anion–ligand interaction stronger, the sulfonamide group was introduced by Hu et al. in designing anion

Fig. 8 Mode of interaction of hydrogen bonding motif-based gelators with cyanide.

responsive gels. They synthesized compound 9a, which formed a green fluorescent gel in DMSO (Fig. 9). 127 In the structure, they additionally incorporated a urea functionality in close proximity to the sulphonamide to strengthen the ligand–anion interaction through participation in hydrogen bonding. However, initially, the urea group promotes self-aggregation via bifurcated hydrogen bonding.^{86,192} Furthermore, the nongelling behavior of 9b in comparison to 9a also endorsed the supportive role of hydrophobic interactions in gelation. The response of organogel 9a towards CN⁻ led to easy-to-discern changes in the fluorescence color. On exposure to CN^{-} , the organogel underwent a gel-to-sol transition into a homogeneous non-emissive solution.

Instead of a long alkyl chain, the sulphonamide functionality has been attached to the cholesteryl group to synthesize the gelator. Conceptually, cholesterol with a large hydrophobic surface can stimulate molecular self-assembly via hydrophobic interactions.172–174 Ghosh et al. established the cholesterol-appended sulfonyl hydrazone derivative 10 as a naked-eye anionic-sensor (Fig. 10).¹²⁸ Employing intermolecular H-bonding involving the sulfonamide moieties and hydrophobic interactions between the cholesterol units, compound 10 formed a stable gel from

Fig. 10 (a) Chemical structure of gelator 10. (b) Change in absorbance of 10 $[c = 2.50 \times 10^{-5}$ M, taken in DMSO-H₂O $(1:1 \text{ v/v})$ upon addition of 40 equiv. of different anions ($c = 1.0 \times 10^{-3}$ M, taken in DMSO). (c) Visual sensing of cyanide by 10 (c = 20 mg mL⁻¹) in DMSO-H₂O (1:1 v/v) involving gel to sol phase transformation: (a) CN^- , (b) F^- , (c) AcO⁻, (d) $H_2PO_4^-$, (e) Cl^- , (f) Br^- , (g) Γ , (h) HSO₄⁻ and (i) NO₃⁻. Reproduced from ref. 128 with permission from The Royal Society of Chemistry (RSC) on behalf of the Centre National de la Recherche Scientifique (CNRS) and the RSC.

DMSO–H2O. The SEM image of the xerogel shows a tiny rod-like fibrous structure in the aggregated state. Among different anions, basic anions like CN^{-} and F^{-} interact with the sulfonamide –NH and disrupt the hydrogen bond-assisted molecular aggregation through deprotonation, causing gel-to-sol transformation. Other weakly basic anions were practically noninterfering in such an event. However, a relatively better response was observed for CN^- than F^- ions. The presence of 2 equiv. of CN⁻ caused the complete destruction of the gel within 30 min, while F^- ions took almost 2 h to bring a similar change. Importantly, in the presence of less than 2 equiv. of $F^$ the gel was virtually stable, while gel-to-sol transformation was achieved with one equiv. of CN⁻ ions. Further, to discriminate these two anions, different metal ions $(Ca^{2+}$ and $Fe^{3+})$ were used as chelating agents. Only Ca^{2+} ions successfully differentiated CN^- from F⁻ ions. The scavenging of F⁻ ions by Ca²⁺ ions from the medium retrieved gelation. Compound 10 also exhibited strong interactions with CN^{-} , showing ratiometric changes in the absorption spectra. Upon successive addition of CN⁻ ions (up to 40 equiv.), the initial absorption band at 282 nm gradually decreased along with the simultaneous increase of a new absorption band at 330 nm, leading to a clear isosbestic point at 312 nm. For F⁻ ions, such a ratiometric response was very weak. CN^{-} ions displayed relatively stronger binding than F⁻ ions in 1:1 stoichiometric fashion $(2.12 \times 10^3 \text{ M}^{-1}$ and 1.04×10^3 M⁻¹ for CN⁻ and F⁻ ions, respectively) with a moderate detection limit (7.96 \times 10⁻⁵ M). As suggested, the F⁻ ion due to its smaller size was more hydrated in water compared to CN⁻ and remained less free to interact with gelator 10, while the CN^- ion being less hydrated could display its basic nature and thereby showed a relatively sharp and better response. Thus, apart from the gel-to-sol conversion of 10 in the presence of CN^- and F^- ions in different time frames, use of Ca^{2+} ions and comparison of absorption spectra and binding constant data were important in discriminating CN^{-} from F^- ions. **Proview Published on 2021 Published on 2022 Published on**

Like sulfonyl hydrazone, acylhydrazones can also interact with anions by hydrogen bonding involving –NH and therefore are utilized in sensor design. The designed gelators 11a and 11b differ in boron substitution and therefore showed different self-assembly attributes as well as anion sensing properties (Fig. 11).¹⁹³ Compound 11a being more hydrophilic than 11b exhibited gelation in the presence of water (acetonitrile–water, 4 : 6, v/v), whilst hydrophobic molecule 11b self-assembled in a chloroform-methanol $(1:9, v/v)$ mixture. In the sensing process, while the gel of 11a remained intact after the addition of CN^- and F^- ions, the organogel of 11b turned into a solution in the presence of both the anions. It was explained by the idea that the presence of the two hydroxyl groups on the boron center provides extra stability to the gel by increasing the number of hydrogen bond contacts and thereby prevents the gel from rupturing in the presence of anions. On the contrary, compound 12 showed a rapid gel-to-sol phase transition with CN⁻ in dioxane (Fig. 11).¹⁹⁴ Strong hydrogen bonding of -NH with CN⁻ followed by deprotonation broke the yellow gel into a dark orange sol instantly. The gel was retrieved by adding Hg^{2+}

Fig. 11 (a) Structures of compounds 11a, 11b and 12. (b) Sol-gel phase transformation of the dioxane gel of 12 under different conditions. Reproduced with permission from ref. 194. Copyright 2019 American Chemical Society.

ions to the broken gel. Importantly, Hg^{2+} ions scavenge $CN^$ ions from the medium and allow the molecules to reassemble. However, the recovered gel showed a decrease in mechanical properties.

Recently, two acylhydrazone derived LMWGs 13 and 14 have been introduced for cyanide sensing (Fig. 12). While compound 13 exhibited a gel-to-sol transition in the presence of cyanide in $DMSO: H₂O$ $(1:1, v/v)$ with a color change from white to yellow,¹⁹⁵ the DMF : H₂O (1 : 1, v/v) gel of 14 recognized cyanide ions in a gel-to-gel fashion involving a rapid color change from yellow to saffron.¹⁹⁶ ¹H NMR studies confirmed that both the compounds undergo deprotonation of the acyl hydrazone –NH in the presence of cyanide in semi aqueous medium. Interestingly, the color change of the cyanide treated gel of 14 was reversible in the presence of a proton source, enabling the reusability of the material.

To control the molecular assembly, –OH groups have been incorporated near the acylhydrazone moiety in different ways to achieve structures 15–17. The –OH group is involved in intramolecular hydrogen bonding with the adjacent imine nitrogen and provides rigidity to the molecules as discussed above. Importantly, in some cases, the six-membered hydrogenbonded network becomes so stable that it prevents deprotonation

Fig. 12 Structures of compounds 13 and 14.

Fig. 13 (a) Structures of compounds 15 and 16. (b) Fluorescence response of organogel 16 (WJG) to the presence of various anions. Reproduced from ref. 198 with permission from The Royal Society of Chemistry (RSC) on behalf of the Centre National de la Recherche Scientifique (CNRS) and the RSC.

of –OH with anions. Compound 15 is a classic example, which forms a gel in n -BuOH/H₂O (9:1, v/v) driven by the hydrogen bondmediated rigidity of the molecules (Fig. 13). 197 The gel, when treated with various anions, underwent a gel-to-gel transition in the presence of CN⁻ ions, exhibiting blue-shifted emission in the fluorescence spectra. The detection was also confirmed visually through a color change of the gel from green-yellow to blue fluorescence. However, this cyanide sensing suffered from poor selectivity as the gel showed a similar response to S^{2-} as well. Interestingly, by spectroscopic studies, it was corroborated that during interaction with anions, the –OH remained intact while the –NH group underwent deprotonation and produced remarkable spectroscopic changes of the gels. On the other hand, the glycerol gel of 16 undergoes deprotonation of both the phenolic -OH and acylhydrazone -NH with CN^{-} and shows visual sensing through a color change of the gel from yellow-green to blue (Fig. 13).198 This color change was associated with a blue shift in the emission of the gel from 516 nm to 498 nm. However, X-ray diffraction pattern analysis of the CN⁻ treated gel revealed peaks at 2θ = 25.95° and 27.90° corresponding to d-spacing of 3.43 Å and 3.19 Å, which indicated that the $\pi-\pi$ stacking between the naphthyl rings of 16 and 16 CN⁻ was not influenced even after the deprotonation process.

Our group, for the first time, explored a pyridoxal-based LMWG 17 in anion sensing (Fig. 14).¹⁹⁹ Structural analysis revealed that, while several hydrogen bond donor and acceptor entities such as phenolic –OH, alcoholic –OH, amide –NH and pyridyl ring nitrogen assist the molecular assembly through hydrogen bonding, the aryl ether segments are involved in π -stacking. A combination of all these weak forces resulted in the formation of stable gels from various solvents such as DMSO, DMF, $DMSO/H₂O$, and $DMF/H₂O$. When the DMSOgel of 17 was subjected to common basic anions and halides, the gel showed a strong affinity toward basic anions such as F^- , A c $O⁻$ and $CN⁻$ ions. These anions caused deprotonation of the phenolic –OH and the amide –NH, for which the self-assembly of 17 was destroyed and the gel was completely transformed into a solution in different times (for F^- : 3 h, CN^- : 3.5 h and ACO^- : 6.5 h) according to the basicity of the corresponding anions ($F^ >$ $CN^ >$ AcO⁻). Regeneration of the gel states after the addition of water to the anion-induced broken gels further supported the deprotonation phenomena. To discriminate F^- , A c $O⁻$ and $CN⁻$ ions, different chelating agents were introduced

Fig. 14 (a) Structures of compounds 17, 18a and 18b. (b) Phase transformation of $18a$ in toluene in the presence of different anions: (a) CN^{-} , (b) F^- , (c) AcO⁻, (d) H₂PO₄⁻, (e) Cl⁻, (f) Br⁻, (g) I⁻, (h) NO₃⁻ and (i) HSO₄⁻. (c) Chemical reversibility of the anion-induced sols and discrimination between F⁻ and CN⁻ ions. Reproduced from ref. 200 with permission from The Royal Society of Chemistry (RSC) on behalf of the Centre National de la Recherche Scientifique (CNRS) and the RSC.

to the anion-induced broken gels. The addition of $BF₃$ to the broken gels retrieved the gel states in all cases. Addition of $Ca(CIO₄)₂$ could recover the F⁻-induced broken gel only and thereby differentiate F^- from AcO⁻ and CN⁻ ions visually. The use of other chelating agents such as $Cu(CIO₄)₂$ and AgClO₄ was also found to be ineffective to differentiate $\rm{A} \rm{c} \rm{O}^-$ and $\rm{C} \rm{N}^-$ ions. Hence, gelator 17 acts as a cyanide sensor but with poor selectivity.

Two structurally similar diaminomaleonitrile-based gelators 18a and 18b have been recently reported by us as excellent cyanide-indicators where the diaminomaleonitrile core as the anion-binding site was attached to the cholesterol unit via an aromatic linker (naphthalene/benzene) (Fig. 14).²⁰⁰ A small change in the π -surface (naphthalene/benzene) significantly influenced the gelation behaviors and gel properties (mgc, thermal stability, morphology, mechanical behavior, stimuli responsiveness etc.). The naphthyl analogue 18a exhibited gelation in a greater number of solvents compared to the benzene analogue 18b, while in common solvents 18a formed a gel with relatively low mgc and improved thermal and mechanical properties. The naphthyl derivative 18a exhibited self-healing behavior and also acted as an injectable material, while the benzene analog 18b did not. However, both the compounds showed identical anion-responsiveness. The toluene and 1,2-dichlorobenzene gels of 18a and 18b respectively sense F⁻ and CN⁻ anions by showing rapid gel-to-sol transformation through

the deprotonation of the diaminomaleonitrile $-NH₂$ group. Other basic anions and halides virtually remained inert to such phase changes and thereby validated the sensing of F^- and CN^- ions. The addition of MeOH to the broken gels recovered gelation with no virtual changes in the gel properties. Importantly, F^- and CN^- ions were distinguished by adding external chelating agents like $Fe³⁺$ ions to the broken gels where only the scavenging of F⁻ ions through Fe F_6^{3-} formation rendered gelation. Further, in solution (acetonitrile) both the compounds displayed a similar trend and recognized them by exhibiting ratiometric absorption spectral changes with 1:1 stoichiometric binding to F^- and CN^- ions.

H-Bonding motif-based gelators are simple in design (Table 2). They are advantageous over metallogels in terms of cost-effectiveness and ease of handling. However, the H-bond mediated detection technique still encounters several limitations. Basic anions like F^- , AcO⁻, H₂PO₄⁻ etc. frequently interfere in the sensing process.128,191,193,194,197,199,200 Interference from basic anions makes this kind of gelators less selective and hence less effective. In many cases, it demands different chelating agents or solvent optimization to show $CN^$ selectivity for such multi-responsive gels, which makes this strategy a little complicated.^{128,200} In some cases, even the chelating agents are inefficient to discriminate CN^- from other anions.¹⁹⁹ These limitations necessitated the development of reaction-based gelators where specific chemical reactions with CN^- ions at the reaction center could minimize the interference of other anions.

In this context, it is mentionable that most of the hydrogen bonding-motif based compounds were subjected to anion binding studies in solution. As in solution molecules remain discrete compared to a gel, solution-phase studies showed different selectivity for anions, and even higher selectivity to

Table 2 List of H-bonding motif-based gelators explored in cyanide sensing

cyanide in some cases.^{127,128} This indicates that the self-assembly nature of the gelators has a critical role in the responsive behavior.

(iii) Reaction-based CN^- sensory gelators

Cyanide responsive reaction-based gelator probes (dosimetric gelators) differ from the H-bonding motif-based sensors only in the reaction centre (binding site), where instead of reversible hydrogen bonding, cyanide forms a permanent covalent bond (Fig. 15). The rest of the gelator segments follow almost similar design strategies. Dicyanovinyl, oxazole, etc. and even an activated imine or carbonyl group can serve as a potential acceptor for cyanide.¹³⁴ The higher nucleophilicity of cyanide compared to other anions in some specific solvents provides an advantage to react irreversibly with the organic gelator molecules. Other anions remain silent to these reactions. This makes such gelators significantly superior to other types of gelators in terms of selectivity. Another advantage is their fast response time, which allows real-time monitoring of the sensing processes. Cyanideinduced chemical changes of the gelator substantially alter the self-assembly of the molecules and often induce a gel-to-sol transition. Usually these reaction centers are either part of the chromophoric units or different segments connected to the chromophores and thus, due to permanent chemical modifications, there are always notable spectroscopic changes for characterization. Power **Process**

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The design of dosimetric sensors is a fairly new domain of research in gel chemistry. Our group has been dedicatedly working on this topic. We have reported dosimetric probes for $Hg^{2+}, ^{201-204}$ perborate,²⁰⁵ sulfide,²⁰⁶ and hydrazine²⁰⁷ for their selective recognition involving sol-to-gel conversion. In the same line, we have shown with compounds 19 and 20 how the functional group modification of the gelator can lead to cyanide sensing selectively (Fig. 16).²⁰⁸ Both the compounds were based on a cholesterol-linked phenyl substituent, a

Fig. 15 Mode of interaction of reaction-based gelators with CN^{-} .

Fig. 16 (a and b) Phase transformation of the organogel of 19 in the presence of various anions along with the proton NMR changes of 19 in the presence of CN^- ions. (c and d) Phase transformation of the organogel of 20 in the presence of various anions along with the proton NMR changes of 20 in the presence of CN^- ions. For (b and d), from left to right: (a) CN⁻, (b) F⁻, (c) AcO⁻, (d) H₂PO₄⁻, (e) Cl⁻, (f) Br⁻, (g) I⁻, (h) HSO₄⁻, (i) CLO_4^- and (j) NO_3^- . Reproduced with permission from ref. 208.

common self-assembling unit. However, while gelator 19 is comprised of an oxime unit as a hydrogen bond donor, gelator 20 possesses a dicyanovinyl moiety as a Michael acceptor for CN⁻. The change in functional group in the gelators has a significant impact on the gelation, gel properties, and morphology; however, the main difference lies in their anion sensing behavior. The oxime analogue 19, due to the free –OH group, interacted with multiple basic anions like CN^{-} , F^{-} , and AcO⁻ and displayed a gelto-sol transition within 1 h with no selectivity. The phase transitions were attributed to the anion-induced deprotonation of the oxime –OH. In contrast, the dicyanovinylated gelator 20 showed a gel-tosol transition within 15 min selectively in the presence of CN^- over a series of anions and demonstrated visual sensing. Such a high degree of selectivity was ascribed to the Michael type addition of CN^{-} to the dicyanovinyl moiety in 20, which in turn established the

Fig. 17 (a) Structures of compounds 21 and 22. (b) Probable mechanism of cyanide sensing by 22. Reproduced with permission from ref. 210.

necessity of designing dosimetric sensors for CN⁻ ions over hydrogen bonding mediated recognition. The cyanide adduct formation was characterized by proton NMR spectra. In ¹H NMR, the upfield chemical shift of the vinylic proton (Ha) from 7.59 ppm to 4.26 ppm endorsed the nucleophilic addition of CN⁻ ions to 20. Similarly, gelator 21 exhibited a selective response to CN⁻ involving the same chemical reaction and recognized it through the gel-to-sol transition in $CH₃CN$ with a vivid color change from orange-yellow to deep red (a response time of 2 h in the presence of 2 equiv. of cyanide) (Fig. 17).²⁰⁹ Not only in the gel state, gelators 20 and 21 also showed similar trends in solution and selectively recognized CN^- over other anions by exhibiting ratiometric changes in the absorption spectra.

An activated ester can also serve as a cyanide acceptor. Compound 22 underwent gelation in DMF: H_2O (2:1, v/v) through extensive $\pi-\pi$ stacking of the naphthalene rings (Fig. 17). 210 The gel was found to be anion-responsive because of the phenolic -OH. CN⁻ ions over a series of other anions resulted in a rapid gel-to-sol transition (within 8 h) with a colour change from yellow to brown. The sensing mechanism involved cyanide-induced deprotonation of the phenolic –OH followed by nucleophilic addition to the activated ester. ¹H NMR, FTIR, and HRMS studies were conducted to establish CN^- adduct formation via nucleophilic addition to the ester carbonyl of 22.

An interesting feature was observed for gelator 22 in that, unlike 20 and 21, its selectivity could be controlled in solution either by adjusting its concentration or by changing the solvent composition. A fluorescence study of 22 in CH₃CN containing 1% CHCl₃ ($c = 2.5 \times 10^{-5}$ M) enabled the detection of CN⁻ ions by fluorescence turn-on but it suffered from less selectivity due to interference of F⁻ and $HP_2O_3^{-7}$. However, an increase in the

Fig. 18 Photograph showing the colour changes of 22 (c = 1.0×10^{-3} M) in the presence of equiv. amounts of various anions (c = 5.0 \times 10^{-3} M) with time in CH₃CN containing 4% CHCl₃. Adapted with permission from ref. 210.

concentration of 22 in solution $(c = 1.0 \times 10^{-3} \text{ M})$ brought about selectivity towards CN^{-} with a distinguished colour change of the solution from yellow to rose-red (Fig. 18). It was proposed that initially, due to the presence of large amounts of basic anions, rapid formation of naphthoxide ions favoured excited-state charge transfer, which with time was repressed by the keto–enol equilibrium in solution. Close proximity of –NH of the keto form (Fig. 17b) then triggered CN^- addition to the ester carbonyl of 22 and produced a distinguishable color change from F⁻ and $HP_2O_7^{\;3-}$. Similarly, the solution-phase interactions of 22 with the same anions in $DMF/H₂O$ (2:1, v/v) showed considerable changes in absorbance and fluorescence selectively to CN^- ions only at higher cyanide concentration, consistent with the gel phase observations.

Despite several advantages, chemodosimetric gelators also have a few drawbacks (Table 3). One major disadvantage of such probes is that the chemical reaction brings about permanent changes of the gelators. As a consequence, the scope for reuse is limited. In some cases, the rate of cyanide addition is very slow at ambient temperature and heating of the gel sample is required to drive the chemical reaction. Fang et al. presented 2-(hexadecylthio)oxazolo[4,5-b]phenazine 23 as a cyanide sensor (Fig. 19). 211 The phenazine derivative formed a yellow fluorescent gel in DMSO. When aqueous solutions of anions were added to the gel at room temperature, no change in fluorescence was noticed in any case even after a considerable time. However, when the mixtures were heated above the gel melting temperature and then cooled down again, a nonfluorescent gel appeared only for CN⁻. At high temperature, CN⁻ addition to 23 destroyed the π - π interactions, which led to

Fig. 19 (a) Chemical reaction involved between 23 and CN^- and (b) change in the emission spectra of the DMSO–water gel of 23 in the presence of increasing amounts of CN⁻. The inset presents the corresponding change in the color of the gel. Adapted with permission from ref. 211.

Fig. 20 (a) Suggested self-assembly mechanism for the gel formation of **24** in DMSO/H₂O and (b) a proposed mechanism for the interaction of 24 with CN⁻. Reproduced with permission from ref. 212.

steady quenching of the emission during a sol-to-gel transition. The gel showed an extraordinary detection limit of 4.18 \times 10^{-10} M for CN^- ions.

In a recent study, supramolecular gelator 24, composed of indolin-2-one and quinoline moieties with a long chain N-alkyl substituted amide group, has been reported to form a stable and durable gel from DMSO/H₂O $(1:1).^{212}$ The addition of water is explained to enhance the solvophobic interactions, which act as a driving force for the gel (Fig. 20a). The SEM image revealed the closely spaced plate-like morphology of the xerogel. Addition of aqueous solutions of various anions

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including CN⁻ as their sodium salts to a DMSO solution of 24 furnished a color change of the gel. In the presence of $CN^$ ions, the orange colored gel changed into a dark purple color. It was suggested that, at first, compound 24 undergoes a pseudo-Michael attack by CN^{-} ions, followed by a ring-closing reaction. Subsequent proton shift and tautomerization reactions finally generate the conjugate anion 24a, responsible for the dark purple color (Fig. 20b). Thus gelator 24 was used as an extremely selective and fast chemosensor for the detection of CN⁻ ions in environmental water resources.

Conclusions and future outlook

Gel-based sensing probes have huge potential in materials chemistry research.80–100 This review provides an up-to-date report describing cyanide sensing in the gel phase. Attention has been given to the approaches where the hydrogen bonding and nucleophilic character of cyanide are encountered to show gel-to-sol as well as gel-to-gel phase transformation for its detection and sensing. The solution-phase interactional approaches for cyanide sensing are well-reviewed.^{132–139} Indeed, limited gelators are available for cyanide detection. Most of the cyanide-responsive gels are serendipitous. This is probably due to the uncertainty of the responsive behaviour of a molecule, which is an inherent challenge in the field of gel chemistry.^{149,213–216} Hence, study of the structure– property relationship of gelators is highly desirable. 217 **Materials Chemistry Frontiers** www.colonisty

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Gel-based cyanide sensors are extremely useful, particularly as accessing sophisticated instrumentation and trained manpower is limited. The study so far performed in this domain is mainly connected with the detection of cyanide involving only phase transformation and observable color changes. Other properties like swelling–deswelling etc. are not investigated. Polymeric gels have been reported to show changes in gel volume in the presence of cyanide.²¹⁸ It would be interesting to see if small molecule gelators respond in a similar way. If the results are positive, it is then possible to construct cyanide responsive actuators. Furthermore, unlike other design-based sensors, low molecular weight gelators are rarely explored in in vivo detection of cyanide, $219-221$ and hence designing fluorophore tagged gelators is highly demanding for constructing bioimaging probes.⁶³ As the majority of cyanide sources belong to industrial waste, emphasis should also be put on adsorbing and separating cyanide from cyanide contaminated water. Importantly, although some designed gelators show quite a low detection limit compared to the prescribed limit of the WHO, their sensitivities for cyanide are not so high in most cases. Alongside this, cyanide mostly exists in aqueous systems, while most of the gelators are studied in pure organic solvent. Hence, discovery of super-gelators capable of forming hydrogels at extremely low concentrations is highly essential for improving the sensitivity of detection.

Conflicts of interest

There are no conflicts to declare.

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