A naphthoquinone based colorimetric probe for real-time naked eye detection of biologically important anions including cyanide ions in tap water: experimental and theoretical studies

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A naphthoquinone based colorimetric sensor (N) bearing hydrazone receptors in aqueous media was developed and its recognition properties towards biologically important anions in DMSO–water mixture (9 : 1) were investigated using spectroscopic methods. The hydrazone based receptors showed selectivity towards anions (F\textsuperscript{−}, OH\textsuperscript{−}, CN\textsuperscript{−} and AcO\textsuperscript{−}), through naked eye observable colour changes, from green to light blue (F\textsuperscript{−}, CN\textsuperscript{−} and AcO\textsuperscript{−}) and violet (OH\textsuperscript{−}). The colour changes were concomitant with spectral changes. The sensor could also detect the presence of fluoride ions in commercially available toothpastes, through remarkable colour and spectral changes. In addition, test paper strips prepared from N were able to detect the presence of cyanide (KCN) and hydroxide (NaOH) in tap water. The study was complimented by density functional theory computations to have more insight in the interaction between N and the anions.

1. Introduction

The development of sensors and receptors for anion detection has been receiving great attention in recent years, because of their extensive roles in the biological, chemical and environmental fields. This technique falls among other optical methods that are proving to be convenient for ion monitoring and detection. Colorimetric detection is mostly popular due to its naked eye detectable changes, achieved without resorting to expensive spectroscopic instruments. Thus, colorimetric sensors do offer several advantages over conventional instrumental methods such as atomic absorption spectroscopy, voltammetry, inductively coupled plasma emission, X-ray photoelectron spectroscopy or mass spectroscopy, due to their simplicity, high sensitivity and selectivity, perform real time-monitoring, cost effective, as well as their naked eye observable colour changes.

Fluoride (F\textsuperscript{−}) and acetate (AcO\textsuperscript{−}) ions play a significant role in environmental and physiological processes. For instance, the presence acetate ion in physiological systems is mostly significant in acetyl coenzymy activities, and it is heavily used in the manufacturing industry for products such as dyes, plastics, paints and paper. On the other hand fluoride ion has been primarily used in the dental care industry and it can be applied in osteoporosis treatment. However, in excessive amounts it is greatly toxic to the biological tissue which can lead to fluorosis and osteosclerosis and cause many serious neurodegenerative diseases. Correspondingly, cyanide is one of the most toxic anions and the World Health Organization has set its maximum permissive level in drinking water at 1.9 \textmu M. Despite a huge number of anion sensors published in literature, there are very limited modules which can detect cyanide ions, simultaneously with fluoride, acetate and hydroxide ions. Moreover, there is still a very huge gap in industry for practical applications of chemosensor modules published in literature. Therefore, there is still a great need to develop more effective, cost effective, aqueous based and user friendly sensors, in order to continuously monitor the levels of these anions in different systems.

Herein, we report the synthesis of a remarkable naphthoquinone based colorimetric sensor N (Scheme 1) with hydrazone derivative signaling units, and its application towards anion sensing. The design of N is based on the rich-in-\pi-electron aromatic framework which is potentially likely to result in an excellent sensor, especially when coupled with a powerful electron withdrawing group (EWG) of the nitro group (\texttextit{NO2}), resulting in a moiety with strong electronic coupling, from the receptor to the reporting unit. Subsequently, N is found highly selective and sensitive to four anions being F\textsuperscript{−}, OH\textsuperscript{−}, AcO\textsuperscript{−} and CN\textsuperscript{−} in DMSO–H\textsubscript{2}O (9 : 1) among several anions. The four anions induce significant changes, both in the absorption spectrum and colorimetric changes upon molar additions of the three anions to N in DMSO–H\textsubscript{2}O (9 : 1). Furthermore, for

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practical applications, test papers were also prepared and used in samples, providing remarkable results, which were able to detect cyanide ion-contaminated tap water through colour changes. The sensor was also applied in real sample analysis of commercially available toothpastes (Colgate and Aqua-fresh) which are known to contain fluoride ions.

2. Experimental section

2.1 Materials and apparatus

All reagents and solvents were commercially available and of analytical grade. They were used without further purification unless stated. UV/vis spectroscopy was performed with PerkinElmer Lambda 35 spectrophotometer in a standard 3.0 ml quartz cuvette with 1 cm path length. FT-IR spectra were obtained on a PerkinElmer Spectrum 100 FTIR spectrometer. $^1$H NMR spectra were recorded on a Varian Mercury VX-300 MHz spectrometer in $d_6$-DMSO, steady state excitation and emission spectra were obtained on the Molecular Device spectraMax M2. Elemental analyses (C, H and N) were carried out on a PerkinElmer 240C analytical instrument. All the measurements were carried out at ambient temperature (25 °C).

2.2 Synthesis of sensor N

The sensor (N) was synthesized according to Scheme 1, using the approach of green chemistry via a simple, low cost and one-step-reaction Schiff base condensation reaction. To the ethanol solution (20 ml) of 2,4-dinitrophenylhydrazine (2 mmol), while magnetically stirred, was charged with the ethanolic solution (20 ml) containing about 1,4-napthoquinone (1 mmol), drop-wise. The resulting solution mixture was catalyzed with 5 ml of acetic acid and refluxed for 3 h, which resulted in a light brown precipitate being formed. The product was filtered out while the solution was hot, and washed repeatedly with warm ethanol. The product was re-crystallized in ethanol to yield a light brown precipitate (N) as outlined in Scheme 1. Yield: 43%. $^1$H NMR (300 MHz, DMSO-$d_6$) $\delta$ 9.92 (s, NH), 8.79 (d, $J = 2.7$ Hz, 2H, ArNO$_2$), 8.22 (dd, $J = 9.7$, 2.7 Hz, 2H, ArNO$_2$), 7.92 (dd, $J = 5.8$, 3.3 Hz, 4H, Ar), 7.67 (d, $J = 9.7$ Hz, 2H, Ar). Elemental analysis calc (%) for C$_{22}$H$_{14}$N$_8$O$_8$: C, 50.97; H, 2.72; N, 21.62; O, 24.69; found: C, 51.27; H, 2.54; N, 20.93; O, 24.69. The UV-vis spectrum of N exhibits a band at 357 nm ($\varepsilon = 1.74 \times 10^4$ M$^{-1}$ cm$^{-1}$).

2.3 General procedures for UV-vis and fluorescence experiments

All UV-vis spectra were recorded in DMSO-$H_2O$ on a PerkinElmer Lambda 35 spectrometer after the addition of tetrabutylammonium salts while keeping the sensor N concentration constant (1 x 10$^{-5}$ M). Tetrabutylammonium salt (TBA) of anions (F$, Cl^-$, Br$^-$, I$, AcO^-$, H$_2$PO$_4^-$, ClO$_4^-$, CN$^-$) were used for UV-vis experiments. Subsequently, all fluorescence spectra were recorded on a Molecular Devices SpectraMax M2 upon the addition of TBA salts while keeping the concentration of N constant (1 x 10$^{-5}$ M). TBA salts of anions (F$, Cl^-$, Br$^-$, OH$^-$, I$, AcO^-$, H$_2$PO$_4^-$, ClO$_4^-$, CN$^-$) were used for fluorescence experiments.

3. Results and discussion

3.1 Colorimetric analysis of N with anions

In order to establish whether there is any chemical interaction between N and all respective anions, a series of prepared anionic solutions (1 x 10$^{-3}$ M) in DMSO-$H_2O$ were added dropwise, each separately, to 3 ml of N (1 x 10$^{-5}$ M) in a similar solution (DMSO-$H_2O$), at room temperature. The resulting observations were recorded as indicated (Fig. 1). Upon dropwise addition of the CN$^-$, F$^-$ and AcO$^-$, all as tetrabutylammonium salts (TBA), a visible colour change was observed from the original colour of N (light-greenish) to light blue (Fig. 1). However, when the OH$^-$ (TBA) was added dropwise to the solution of N, the colour changed to violet (Fig. 1). The intensities of colour depended on the concentration of the N in the solution, ranging from intense (high concentration) to relatively weaker or lighter colours at lower concentrations. Moreover, these anions are all electronegative in nature, which are likely to interact with electropositive protons (–NH) in the structure, however, they all have different affinities towards N due to differences in association constants with the proton group in the structure (N), normally with F$^-$ strongly associated with the protons due to its smaller size and high electronegative than others. In contrast; other competitive anions such as Cl$^-$, Br$^-$, I$, H_2PO_4^-$, and ClO$_4^-$ were completely non-responsive as there were no significant colour changes observed upon their addition of each to the solution of N.
3.2 UV-vis studies of N

The absorption spectra of N in DMSO–H₂O were characterized by two prominent absorption bands at 356 nm and 637 nm, attributed to internal-charge transfer (ICT) of the nitrophenyl groups and the naphthyl moiety in the middle, respectively. Moreover, N displayed a fluorescent emission in visible light region characterized by two weak vibronic peaks, with maxima at 416 nm and 438 nm respectively. The weakened emission is suspectedly attributed to the presence of electron withdrawing NO₂ groups in the structure.

3.3 Photophysical properties of N towards CN⁻ and other anions

UV-vis titrations were conducted in order to investigate spectra behaviors of N upon interacting with different anions (F⁻, Cl⁻, Br⁻, I⁻, AcO⁻, H₂PO₄⁻, ClO₄⁻, CN⁻) in DMSO–H₂O and confirm the colorimetric activities observed in the section above. Subsequently, upon the molar addition of CN⁻ to N, a remarkable change in spectra activities were observed, where an initial absorption band at 356 nm, due to the n → π* transitions, gradually decreased significantly, at the same time experiencing a hypsochromic shift, while the band at 637 nm, attributed to π → π* transitions, increased in intensity (Fig. 2a), upon complexation with an incoming cyanide ion. Moreover, a new band centered at 445 nm appeared gradually with a bathochromic intensity increase. The isosbestic point observed is a clear indication of two species in one system at equilibrium. A 3-dimensional plot (Fig. 2b) of the titrations of N with CN⁻ clearly shows the variations of the spectra and their gradual shifts as explained above. The gradual spectral changes experienced with the molar additions of the cyanide ion, were complimentary with the colorimetric activities observed. The summary of spectral data is provided (Table 1) detailing the molar absorptivities (with and without the addition of CN⁻), the fluorescence quantum yield and the Stokes shift (the difference in wavelength of the band maxima of absorption and emission).

Accordingly, the Job plot (Fig. 3a) has given a clue that the interaction ratio between N and CN⁻ is predominantly 1 : 2 (N : 2CN) with the possibility of fractional existence of a 1 : 1 interaction ratio, which is closely in agreement with theoretical prediction as shown (Fig. 4a), interacting through the hydrazone protons. The titration profile (Fig. 3b) of N with CN⁻ is also in agreement with the Job plot results, as the interaction seemed to reach equilibrium upon the addition of 1 to 2 molar of the toxic anion. Furthermore, similar changes were also observed in UV-vis spectra of N upon molar additions of F⁻ and AcO⁻ (Fig. 5a and b). Nevertheless, the addition of Cl⁻, Br⁻, I⁻, H₂PO₄⁻ and ClO₄⁻, did not result in any significant response (Fig. 4b).

Job’s plot has hinted that the binding ratio between N and the respective anions (F⁻, OH⁻, AcO⁻ and CN⁻) is at 1 : 2, with anions binding via hydrogen bonding of the –NH group, well known with similar receptors. However, there exists a possibility

| Sensor | λmax, abs peaks (nm), without CN⁻ | ε (M⁻¹ cm⁻¹) without CN⁻ | λmax, abs peaks (nm), with CN⁻ | ε (M⁻¹ cm⁻¹) with CN⁻ | λmax, em (nm) (Φₚ) | Stokes shift (nm) |
|--------|----------------------------------|---------------------------|-------------------------------|-------------------------|-------------------|-------------------|
| N      | 356; 637                         | 18 000; 8700              | 320; 637                      | 13 300; 13 600          | 415; 436 (0.58)   | 79                |

Fig. 2 UV-vis absorption spectra of N in DMSO–H₂O (1 × 10⁻⁶ M) upon the molar addition (0–3 equiv.) of (a) CN⁻. (b) A 3-D plot of the spectra.

Table 1 Spectral data of N and interaction with CN⁻
of deprotonation once excess anions are added, liberating the proton from the –NH group to form the HCN entity as shown (Scheme 2), leaving the imine group in a radical form. Thus, the proposed interaction mechanisms is based on hydrogen bonding through the –NH group, followed by deprotonation when an anion is in excess. The deprotonation mechanism is however taking place in sequences via intermediate steps, afore attaining the final structural conformation, which is kinetically and thermodynamically governed by the fundamental principles of the rates of chemical reactions.

3.4 Photophysical properties of N towards OH$^-$

Moreover, the molar addition of OH$^-$ to N, resulted in another slightly different behavior from the above anions (CN$^-$, F$^-$ and AcO$^-$), both in terms of colour (Fig. 1) and spectral changes (Fig. 7). The initial intense peak at 357 nm decreased in intensity with a hypsochromic shift effect (at 320 nm), with a new peak simultaneously forming at 445 nm due to the complexation of N with OH$^-$. While the peak at 637 nm has experienced increase in intensity, the isosbestic point at 408 nm clearly indicates that two species existed at equilibrium in one system. A better view of spectral changes upon the molar addition of OH$^-$ are displayed in the 3-dimensional plot (Fig. 7b), which clearly shows how individual spectrum varies with one another. The Job plot of N with OH$^-$ shows the admixture of possibly a 1 : 1 and 1 : 2 (N : OH) or even more complex than this (Fig. 6), presumably due to the interaction nature of the hydroxide ions with N, which is mainly not only through hydrogen bonding, but are also rich in hydrogen atoms (OH). As a result, even the colorimetric activities (green to violet) observed were slightly different from other ions detected (CN$^-$, F$^-$, AcO$^-$), green to light blue (Fig. 1). Moreover, the titration profile (inset, Fig. 7a) clearly indicates the dynamics of the formation of the complexed N-OH moiety, concomitant with the disappearance of N. In Table 2, the summary of spectral data of N are provided detailing the absorption wavelengths and molar absorptivities (with and without the addition of OH$^-$).

![Fig. 3](image-url) (a) The Job plot and (b) titration profile, of N with CN$^-$.  

![Fig. 4](image-url) (a) The proposed binding mechanism of (a) N to 2CN$^-$ (1 : 2) and (b) titration spectra of other anions.
3.5 Fluorescence studies of N upon interaction with anions (CN⁻)

The fluorescence property of sensor N towards its selective anions was investigated in DMSO–H₂O by fluorescent molar titrations. The free sensor N exhibits two emission maxima at 415 nm and 436 nm upon excitation at 400 nm. However, when 0–3 equiv. of CN⁻ (tetrabutylammonium cyanide) was added to the solution of N, a concomitant increase in fluorescence intensity was observed (Fig. 8). The fluorescence intensity of N peaked at 17, however, when the molar addition of CN⁻ (0–3 equiv.) was gradually introduced, there was a slight enhancement in the fluorescence intensity to about 40, ascribed to structural deprotonation of N (from NH–CN to N⁻ + HCN), after which no matter how much of CN⁻ was added afterward, no significant change could be observed. The molar addition using other anions did not induce any significant change in the emission intensity. Thus, the enhanced emissive properties of N may be ascribed to structural deprotonation (from NH–CN to N⁻ + HCN) upon the addition of CN⁻, which is may not be the case upon adding any other anion.

The characteristics of the fluorescence enhancement are presided by the removal of the n → π* transitions which are normally known to mask the π → π* transition, which are mainly responsible to the emission behaviors. This is another

![Figure 5](image_url)  
Fig. 5  The UV-vis titration spectra of N in DMSO–H₂O with 3 equiv. of (a) AcO⁻ and (b) F⁻.

![Scheme 2](image_url)  
Scheme 2  Proposed binding mechanism of N on interaction with anions (CN⁻ ion) in DMSO–H₂O.
indication that colorimetric activities of the sensor upon the addition of CN$^-$/CO$^-$ are accompanied by electronic changes of the sensor, thereby affecting the HOMO and LUMO interactions significantly. Thus, N can be used as a “turn-on” fluorescence sensor for CN$^-$ detection in DMSO$-$H$_2$O. Moreover, it is strongly believed that the presence of the –NO$_2$ electron withdrawing group (EWG) has some anti-fluorescence properties, thus weak emission intensity is observed on N.

The stoichiometry investigation of sensor N and the selective anions, Job plots were produced for each anion. The solutions were prepared ranging from 10 ml sensor : 0 ml anion to 1 ml sensor : 9 ml anion and from these settings the Job’s plots were obtained (Fig. 5), it the interaction ratios for AcO$^-$/CO$^-$, OH$^-$ and F$^-$ were found to be 1 : 1 while that of CN$^-$ was 1 : 2.

4. Computational calculations

In order to further understand the electronic transitions, the structures of the uncomplexed (N) and complexed (NF) sensors in gas phases were calculated on DFT method [B3LYP/6-31G**] with Spartan’14.$^{33-38}$ The calculations have indicated that there is a significant decrease in the energy gap (HOMO–LUMO) from the uncomplexed to complexed (Table 3). The decrease in energy gap is in agreement with the characteristics of the change in spectra of N, upon complexing with F$^-$, a bathochromic effect. Subsequently, the energy levels of both HOMO and LUMO have decreased upon the molar addition of F$^-$ to N, with the LUMO decreasing more significantly than the HOMO, which is normally associated with a more stable set of orbitals.

![Fig. 6](image)

*Fig. 6* The Job plot of N with (a) OH$^-$, (b) AcO$^-$ and (c) F$^-$ in DMSO$-$H$_2$O.

![Fig. 7](image)

*Fig. 7* UV-vis absorption spectra of N in DMSO$-$H$_2$O (1 $\times$ 10$^{-5}$ M) upon the molar additions (0–3 equiv.) of (a) OH$^-$, (b) A 3-D plot of the spectra.

| Sensor | $\lambda_{\text{max}}$, abs peaks (nm), without OH$^-$ | $\epsilon$ (M$^{-1}$ cm$^{-1}$) without OH$^-$ | $\lambda_{\text{max}}$, abs peaks (nm) with OH$^-$ | $\epsilon$ (M$^{-1}$ cm$^{-1}$) with OH$^-$ |
|--------|---------------------------------|-----------------|-----------------|-----------------|
| N      | 356; 637                        | 18 000; 8700    | 319; 637        | 13 300; 13 600  |

Table 2  Spectral data of N and interaction with OH$^-$
The chemical interaction of N with anions has resulted in structural distortion, for instance, upon complexing with F⁻, saw the structure shortens in length, from 180.180 Å to 17.924 Å (N₅ to N₆), decreasing in length with 0.256 Å. Furthermore, the structure has also experienced angular distortion through C₂N₃N₅ (or C₅N₄N₆) of 146.39° (Fig. 10) after the addition of F⁻, initially from 150.02° of N, signaling the hydrogen bonding interaction between the host (N) and the guest (F⁻). The Mulliken’s atomic charge at the nitrogen of the NH group changed from 0.457 to 0.375 upon the NF adduct formation. Normally, the change of the negative environment is ascribed to the internal charge transfer (ICT) between the host and the guest, which is responsible for lowering down the HOMO–LUMO energy levels, through the stabilization of π* excited state of N by charge delocalization. The HOMO–LUMO gaps of NF have decreased (Fig. 9) and this translates the decrease in the molecular energy and thus stability of the complexed entity. Lowering of the HOMO–LUMO gap is responsible for absorption to be redshifted to longer wavelength and thus colorimetric activities are observed.

### Table 3  The summary of HOMO–LUMO orbitals and the energy gaps of N in the absence and presence of F⁻

|       | N (eV) | NF (eV) |
|-------|--------|---------|
| LUMO  | -3.17  | -7.09   |
| HOMO  | -5.61  | -7.91   |
| Gap   | 2.44   | 0.82    |

Fig. 8  Fluorescence spectra of N in DMSO–H₂O (1 × 10⁻⁵ M) upon titration with CN⁻ (0–3 equiv.).

**Fig. 9**  HOMO and LUMO diagrams of N in the presence and absence of F⁻.

5. Real sample analysis  

The effectiveness of sensor N towards anions, particularly F⁻, was further investigated by employing quantitative analysis using commercial tooth pastes (two). The toothpaste sample solution was prepared as 20 mg ml⁻¹ in 1 ml of H₂O. The toothpaste aqueous solution was added to N (1 × 10⁻⁵ M in DMSO–H₂O) dropwise, resulting in color change from green to light blue (Fig. 11c), at ambient temperature. In addition, concomitant with the colour change, the UV-vis titrations of N with tooth pastes generated parallel spectra, hinting to the presence of F⁻ (Fig. 11a and b). Thus, N is highly selective and...
sensitive to F\textsuperscript{−}, even when dissolved in water, which is very ideal in real time analysis.

In order to consolidate that F\textsuperscript{−} was responsible for the observed spectral and colorimetric changes in the toothpastes (Fig. 11a and b), three more titrations were carried out using different fluoride-containing counter ions (NH\textsubscript{4}F, KF and CaF\textsubscript{2}), different from TBA, with spectra displayed (Fig. 12a–c). Upon the molar additions of fluoride containing solutions (each at

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**Fig. 10** Optimized structure showing the positions of atoms.

**Fig. 11** UV-vis spectra of N (1 × 10\textsuperscript{−5} M) in DMSO–H\textsubscript{2}O with (a) Aquafresh, (b) Colgate, both dissolved in aqueous solutions and (c) visual colour change of N (1 × 10\textsuperscript{−5} M) upon addition of drops of toothpaste solution (20 g ml\textsuperscript{−1}) in H\textsubscript{2}O.
a time) to N, the spectra from the three graphs were characterized by the gradual decrease in absorption at 355 nm, concomitant with an increase in the absorption band at 510 nm. The isobestic points were clearly visible at 405 nm in both cases, signaling the formation of the N-F adjunct, upon the molar addition of F\(^-\) to N. These results were all in agreement with the spectra from the two toothpastes analyzed (Fig. 11a and b), as well as those of F\(^-\) (Fig. 5b), which is evidently suggesting that the presence of fluoride is responsible for the spectral changes, rather than the counter ions TBA (Fig. 4b). Notably, the molar additions of these solutions to N resulted in weaker interactions as displayed by spectra intensities, even when larger molar equivalents (20 equiv.) were used, the action suspected ascribed to the poor solubility of these compounds (NH\(_4\)F, KF and CaF\(_2\)) in DMSO–H\(_2\)O, and also the high association constants of F\(^-\) to these counter ions.

6. Competitiveness studies

In order to explore the possibilities of cross-detection of sensed anions by N, competitive discrimination studies were carried out, in different sequences (Table 4). It is evident that F\(^-\) was more associated with the sensor than any other sensed anion (Fig. 13), followed by OH\(^-\). Upon adding 5 equiv. of all anions listed in Fig. 13 in one cuvette (one after another), only F\(^-\) and OH\(^-\) persisted defying interferences from the other anions. This is in total agreement with the UV-vis titration behaviors of the two respective ions (Fig. 5a and 7a).

Furthermore, to study cross the detection behaviors of the sensed anions in the presence of N closely, competitive studies were carried out through addition of (sensed) anions added in different sequences (Table 4), in order to determine which of one of them stands out without distraction from others. Upon running out four different addition sequences in Table 2, it turned out that the discrimination patterns are dependent on which ion was added first and last. The spectrum displayed with high absorbance intensity is always the one added last in the sequence (Fig. 14a–d). This translates into the conclusion that, the first anion to be added is distracted by the subsequent following anions, and gets displaced by an immediate incoming anion, such that the characteristics of the last added anion dominates the spectra.

7. Applications of N on test papers

Testing paper strips were prepared from a printing paper which was dipped in 1 \(\times\) \(10^{-3}\) M solution of N in DMSO–H\(_2\)O (Fig. 14) for 24 hours. The paper strips were left to dry for another 24 hours at room temperature. The 10 ml solutions of KCN, NaOH, KF and Zn(AcO)\(_2\) (1 \(\times\) \(10^{-3}\) M) in tap water were prepared as a result (Fig. 15). The dry prepared test strip was dipped instantly (2 seconds at most), in each of the solutions, and allowed to dry on the white sheet of paper. Upon dipping the N-soaked strips in each solution, the results showed that cyanide and hydroxide ions induced intense colour changes on the paper strips (Fig. 15), from original yellow to dark bluish (CN\(^-\)) and dark brownish (OH\(^-\)) colours, instantaneously, detectable by naked eye. The colours of the litmus test papers may not be distinctively clear in the diagram, but cyanide (KCN) contaminated water gives a dark bluish colour, while hydroxide (NaOH) gives a dark brownish colour, when the paper is wet. Thus the distinction can be distinctively observed when the paper strips are dipped and removed immediately, when wet. Moreover, for simplicity and real-time applications, the preparation of test papers was prepared on normal printing papers, just to highlight that this can be done on a very simple substrate, readily

| Table 4 | Addition sequences of competing anions to N (5 equiv. for each addition) in different orders in DMSO–H\(_2\)O\(^a\) |
|---------|----------------------------------------------------------------------------------|
| Sequences | N → F → AcO → CN → OH |
| 1 | N → CN → AcO → OH → F |
| 2 | N → OH → AcO → CN → F |
| 3 | N → AcO → OH → F → CN |
| 4 |

\(^a\) F= fluoride ion, AcO = acetate ion, CN = cyanide ion, OH = hydroxide ion.
available even in rural and remote areas. The resulting tests were all complementary in terms of the colorimetric effects on the test papers. However, the result from the test paper strips were of high importance, that cyanide, in particular could be detected in aqueous solutions from ionic compound (e.g. common salts) sources.

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**Fig. 13** The combined titrations of N (1 x 10^{-5} M) vs. anions (5 equiv.) in DMSO–H₂O in different sequences.

**Fig. 14** The competitive nature of discriminated anions (5 equiv. each) added in four different sequences in DMSO–H₂O at 1 x 10^{-5} M of N.

**Fig. 15** N-soaked testing litmus paper, immersed in tap water contaminated with different contaminants as indicated in the diagram.
8. Conclusion

In summary, a highly stable sensitive chemosensor N was synthesized in a single step using Schiff base reaction and tested for sensitivity toward anions using the colorimetric sensor concept at a micro level in organic medium using water soluble DMSO–H2O mixture. Amongst all anions, N displayed a strong affinity of sensor towards AcO−, F−, OH− and CN− ions, as confirmed spectroscopically by UV-vis titration and naked eye observations; with a dramatic colour change from light green to light blue for AcO−, F− and CN− and a light green to violet/purple for OH−. The mode of interactions of sensor N with anions was determined to be predominantly 1 : 1 for the three detectable anions and 1 : 2 for CN− using the job’s plot method. The selectivity is greatly based on charge–charge interactions, and the involvement of the N–H group’s hydrogen bonds. The unique binding properties can have a greater utility in the development of new anion sensors with enhanced binding affinity and substrate specificity.

Conflicts of interest

There are no conflicts to declare.

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