1. Introduction

Over the past few decades, traditional techniques like atomic absorption spectroscopy (AAS), inductively coupled plasma atomic emission spectroscopy (ICP-OES) voltammetry and colorimetric techniques have been used for the detection of Fe$^{3+}$. Nonetheless, these techniques require advanced equipment, time-consuming sample preparation procedures and trained professionals. Fluorescence techniques have been extensively applied by research groups to avoid these disadvantages. A number of fluorescent probes for selective sensing of Fe$^{3+}$ ion have been developed. As Fe$^{3+}$ possesses paramagnetic nature in its 3d orbital it leads to fluorescence quenching. Most of the early reported probes exhibited turn-off (fluorescence quenching) response because of the paramagnetic nature of Fe$^{3+}$ ion that restricts its application in biological systems. Turn on signals are dominant in biosensing compared to turn-off ones because of their good processing of signals in biological media.

Highly optically active metal chalcogenide/quantum dots (QDs) have been reported as fluorescence based sensors/receptors with excellent stability and photophysical properties. Due to their tunable optical properties they have gained importance in numerous fields such as photovoltaics and bioimaging and as sensors. In the early years Jiang et al. reported a gallic acid modified nanometer sized alumina micro-column separations and were able to detect up to 52.1 μM Fe$^{3+}$ in real human blood samples using ICP-MS. With time, in 2019 Mohammadi et al. developed a MBTBA-Fe$_3$O$_4$@SiO$_2$ nanocomposite fluorescent ligand capable of sensing Fe$^{3+}$ with a detection limit of 43.09 nM. Further Nibu et al. reported a dual responsive colorimetric/fluorescent turn-on sensor with detection limit up to 7.49 μM concentration. All these probes were for iron detection in water samples. Reports on detection of metal ions other than Fe$^{3+}$ in blood samples can also be found. In 2009, Jung et al. reported a BODIPY-functionalized magnetic silica nanoparticle fluorescent receptor for probing Pb$^{2+}$ in children’s blood and the same group in 2010 reported a nitrobenzene-functionalized Ni@SiO$_2$ core/shell magnetic nanoparticles as a fluorogenic chemosensor that removed about 96% of Cu$^{2+}$ in human blood. Subsequently in 2011 a macrocyclic dioxotetraamine probe of two-photon excited chemosensor was designed by Liu et al. for determination of copper ions with a detection limit of 0.007 μM in human blood serum samples based on the principle of ICT. Bandyopadhyay et al. reported a fluorescent ligand to determine the concentration of inorganic phosphate and detected about 1.82 mM of inorganic phosphate in chicken serum. Recently, Vishaka et al. designed a colorimetric chemosensor for the sensing of Cu$^{2+}$ with a detection limit of 1.31 μM in human blood serum samples. For the first time in 2016 Wei et al. reported a Fe$_3$O$_4$@ZnO based fluorescent chemosensor for the detection of Fe$^{3+}$ in human blood serum samples with 92.6–108.4% recovery and no reports since then have been reported for such chemosensing of Fe$^{3+}$ in blood. Interference of other transition metal ions, like Al$^{3+}$, Pb$^{2+}$, Hg$^{2+}$, Cu$^{2+}$ with Fe$^{3+}$ results in poor selectivity and...
sensitivity of the chemosensor. However many colorimetric and fluorometric chemosensors are described in the literature for intracellular imaging of cells. Most of the reported probes cannot be used in live cells due to their cellular toxicity and many of them are susceptible to be affected by the background fluorescence since they possess a shorter emission wavelength below 550 nm.

Xanthene and its derivatives with their spirocyclic structure being responsible for off and on fluorescence when the specific ion unbinds and binds to the probe respectively have been reported for detection of Fe$^{3+}$ ions in water samples based on this mechanism. Still most of the reported probes show some fragility and insufficiency like interference of other metal ions and poor sensitivity. However, the present work on such probes for Fe$^{3+}$ detection in human blood samples is first of its kind. Turn-on fluorescent probes favor high selectivity, sensitivity and anti-interference, hence it demands the synthesis of new fluorescent probes for the detection of Fe$^{3+}$ ions. Herewith, we report an excellent biocompatible rhodamine 6G-derivative fluorescent probe named RG5NC as a turn-on fluorescent sensor with high selectivity and sensitivity (in nM range) towards Fe$^{3+}$ in the existence of various other metal ions. The probe exhibited negligible cytotoxicity.

2. Experimental

The materials and methods used for synthesis of probe (RG5NC) and its Fe$^{3+}$ complex can be found in ESI 1–9 where Schemes 1 and 2 explains the reactions involved. Details of preparation of stock solution for spectral determination, absorption and emission studies are explained in ESI 10 and 11.

2.1 Detection of Fe$^{3+}$ in real samples and human blood samples

Industrial waste water was collected from paper and plastic industry from Harohalli Industrial Area, Bangalore rural, Karnataka, India. Human blood samples were collected from Arunodaya Polyclinic, Harohalli, Bangalore and samples were digested as per the established standard protocol and the obtained blood serum was used for further studies. Details of the certified reference material is attached in ESI 12.

2.2 Cell viability assessment

The NIH 3T3 mouse embryonic fibroblast cells (ATCC) were used for the cell culture experiments for testing cytocompatibility of the composite. The cells were grown in 25 cm$^2$ culture flasks with DMEM, supplemented with 10% FBS and 1% antibiotic/antimycotic solution. Trypsinization of confluent cells in the culture flask was carried out using trypsin (0.25% trypsin–EDTA solution). The cells were counted using a hemocytometer (Improved Neubauer cell counter). Tissue culture polystyrene (TCPS) well plates were used as control in all the cell culture experiments.

The cellular viability after incubation of cells with different Fe complex concentration was quantified by MTT assay. In this assay, formazan crystals are formed when metabolically active cells react with MTT salt by the activity of oxidoreductase enzymes in the mitochondria. The concentration of the color formed by the formazan is directly proportional to the percentage of viable cells. Before adding MTT solution to the cell culture dishes, the cells were washed with PBS in order to remove any probe/complex present in the cell culture solution. Additionally, after incubation of cells with MTT solution, formed formazan crystals were dissolved in DMSO followed by centrifugation and OD was taken of the supernatant to avoid any interference during OD measurement. For MTT assay, 10,000 cells were seeded in each well of 96 well plate and incubated with Fe complex with various concentrations. After 24 h and 72 h of incubation, 0.5 mg mL$^{-1}$ of the MTT solution was added to each well and incubated for a period of 2 h. The formazan crystals formed were later solubilized in DMSO and the optical density was recorded at 570 nm using an ELISA plate reader (Tecan InfiniteVR M1000 PRO). The MTT assay was repeated at least three times.

The cellular morphology and intracellular fluorescence after uptake of Fe complex were observed using fluorescence microscope (INCell Analyzer 6000, GE Healthcare Life Sciences, USA). The cells were grown in 24 well plate with Fe complex or ligand for 24 h and thereafter fixed with 3.7% formaldehyde in 1× PBS for 20 min and Hoechst 33258 dye was used to stain the nucleus of the cells. The cells were stained only using Hoechst 33258 dye which binds DNA of the cells and gives blue fluorescence. The fluorescence in green and red channels by cells is resulting in cellular uptake of the complex which is absent in the cells incubating without complex. All the experimental data, obtained using MTT assay are expressed as mean ± standard deviation (SD) and were analyzed by one-way ANOVA (SPSS 16.0) for the calculation of significance level of the experimental data. The differences were considered statistically significant, when $p \leq 0.05$.

3. Results and discussions

3.1 Selectivity study

The selectivity of RG5NC in MeCN solution was examined with different metal ions (like Al$^{3+}$, Ca$^{2+}$, Zn$^{2+}$, Cu$^{2+}$, Ni$^{2+}$, Cd$^{2+}$, Pb$^{2+}$, Fe$^{3+}$, Cr$^{3+}$, Hg$^{2+}$ and Co$^{2+}$). As shown in Fig. 1a, after coordination of Fe$^{3+}$ with RG5NC, the probe exhibited notably strong and high selective ‘OFF–ON’ absorption at $\lambda_{\text{max}} = 529$ nm. The free probe remained colorless when no Fe$^{3+}$ ions were added into the solution. The solution color changed from colorless to orangish pink upon addition of Fe$^{3+}$ ions into the RG5NC solution (inset Fig. 1b). Whereas, the other metal ions did not incite any visible color change. These results designate that RG5NC shows excellent selectivity for Fe$^{3+}$ in MeCN solution.

To attain further thorough analysis on the selectivity of RG5NC for different metal ions, the change in the fluorescence intensity upon the addition of metal ions under the same condition was also examined by using fluorescence spectra. The fluorescence spectra of RG5NC in MeCN exhibited no fluorescence at 553 nm for free probe. Upon addition of Fe$^{3+}$ to the RG5NC solution, a significant fluorescence enhancement was
observed as shown in Fig. 1b. All other metal ions did not display any fluorescence enhancement under similar conditions. These aspects indicate that opening of RG5NC spirrolactam ring is due to the Fe$^{3+}$ induced delocalization of xanthene moiety by chelation enhanced fluorescence (CHEF) mechanism and this reveal that RG5NC is a highly selective fluorescence chemosensor for Fe$^{3+}$ and can be applied to biological sensing. The fluorescence quantum yield were calculated to be 2% and 78% in the absence and presence of Fe$^{3+}$ ions (300 μM) respectively with rhodamine 6G as standard ($\Phi_F = 0.95$ in ethanol).²⁰

To gain more insight into the binding behavior of RG5NC with Fe$^{3+}$, fluorescence titrations of Fe$^{3+}$ against RG5NC was monitored in MeCN solution. As seen in Fig. 1c, the free probe RG5NC did not show any fluorescence at 553 nm. Upon addition of Fe$^{3+}$, it leads to a remarkable increase in the emission intensities and reaches maximum at 300 μM. Since our main focus of the study was for sensing of biological samples and bio imaging it was necessary that the sensor is acceptable for physiological pH, hence, we evaluated the fluorescence response of RG5NC in presence and absence of Fe$^{3+}$ at different pH values ranging from 1 to 11 as shown in Fig. 1d. The fluorescence OFF–ON worked well in the pH range 6 to 9, suggesting that the sensor RG5NC could be easily used for determining Fe$^{3+}$ in biological samples at physiological pH.

The limit of detection for this fluorescent probe was evaluated. Each spectrum was recorded at $\lambda_{ex/em} = 550/570$ nm. As shown in Fig. 2a, the Fe$^{3+}$ concentration was varied over the range of 0.10–20 μM. The calculated detection limit of Fe$^{3+}$ is 8.2 × 10⁻⁹ M with a good linear regression ($R = 0.992$).²⁰

Further, binding constant of RG5NC with Fe$^{3+}$ was calculated using a Benesi–Hildebrand plot.²¹ The results in Fig. 2b shows a plot of $1/(I - I_0)$ vs $1/[Fe^{3+}]$ and yields a binding constant ($K_a$) value as $3.3 \times 10^4$ M⁻¹ ($R^2 = 0.992$) suggesting the sensor binding ability between the probe and Fe$^{3+}$ ion. The complexation ratio of RG5NC with Fe$^{3+}$ was also explained in MeCN using Job's plot. The molar concentration of Fe$^{3+}$ was varied from 0 to 0.9 in a solution containing [Fe$^{3+}$] + [RG5NC] and the total concentration of RG3NC with Fe$^{3+}$ was 20 μM. The obtained results suggests that the fluorescence intensity reached a maximum value when the molar fraction of Fe$^{3+}$ is 0.5 (Fig. 2c), indicating that the Fe$^{3+}$ complexes with RG5NC is in 1 : 1 binding ratio.

In order to further investigate the selectivity of RG5NC for Fe$^{3+}$, its selectivity for Fe$^{3+}$ in presence of other competitive metal cations were examined under same conditions. Fig. 3a
depicts changes in the emission spectra of RG5NC. The detection was experimented in the presence/absence of other competitive metal cations. These results evidently indicate the noninterference of other competitive metal cations during selective sensing of Fe$^{3+}$. One of the important parameters in developing a novel probe for practical and on field applications is its reversibility. The reversible interaction between RG5NC and Fe$^{3+}$ was confirmed by the addition

![Image of fluorescence spectra](image)

Fig. 2 (a) Fluorescence spectra of RG5NC with various concentration of Fe$^{3+}$ in MeCN solution varying from 0–20 μM (b) Benesi–Hildebrand plot to evaluate binding constant. (c) Job’s plot with a total concentration of [RG5NC] + [Fe$^{3+}$] = 20 μM.

![Image of selectivity and sequential addition](image)

Fig. 3 (a) Selectivity of RG5NC for Fe$^{3+}$ in presence of other metal cations. (b) Fluorescence spectra of RG5NC after sequential addition of Fe$^{3+}$ and 1:1 AcO$^-$ solution.
of AcO⁻ ions into the [RG5NC–Fe³⁺] complex. The experimental results exhibited color change from orangish red to colorless upon introduction of AcO⁻ ions forming iron acetate. Concurrently, about 95% of the fluorescence intensity was quenched. Then the fluorescence intensity was originally regained on addition of Fe³⁺ into the mixture. This process was repeated at least five times to suggest the reversibility of probe (Fig. 3b).

4. Cytocompatibility and cell imaging studies

The cytocompatibility of [RG5NC–Fe³⁺] complex was confirmed with NIH 3T3 cells in vitro. No significant difference was observed in the cell viability up to 80 μM concentration of [RG5NC–Fe³⁺] complex in the media (after 1 day of incubation) when compared to the control (Fig. 4). However, at higher concentration (100 μM) the viability of cells was less compared to control. Interestingly, this difference in the viability disappeared when cells were allowed to grow with the [RG5NC–Fe³⁺] complex for 3 days. Initially, intracellular stress is induced because of higher concentration of [RG5NC–Fe³⁺] complex. However, with time, cells recover and start growing like control even at higher concentration of [RG5NC–Fe³⁺] complex.

The main aspect of the study was to detect the presence of Fe³⁺ in NIH 3T3 mouse embryonic fibroblast cells in vitro. Two different sets of cells, one were incubated with [RG5NC–Fe³⁺] and the other treated with just the probe. The latter set of cells did not show any fluorescence signal (Fig. 5) while the other exhibited fluorescence. Additionally the intensity of the signal was observed to increase with increase in concentration of the [RG5NC–Fe³⁺] uptake. The cells could efficiently uptake the probe and fluoresce illustrating its ability to act as sensor to detect Fe³⁺ in vitro.

![Fig. 4](image.png)

NIH 3T3 cell viability grown with Fe complex. Error bars represent mean ± SD. Asterisk shows the significant difference at p ≤ 0.05 with respect to control.

![Fig. 5](image.png)

Representative fluorescence images of NIH 3T3 cells after 24 h grown with 20 μM RG5NC (a) bright field, (b) red channel (c) green channel; 100 μM (d) bright field, (e) red channel (f) green channel; 20 μM [RG5NC–Fe³⁺] complex (g) bright field, (h) red channel (i) green channel and 100 μM Fe complex (j) bright field, (k) red channel (l) green channel. Scale bar is 50 μm.
4.1 Detection of Fe$^{3+}$ in real samples

Further, we tested the applicability of this probe RG5NC for the detection of Fe$^{3+}$ ions in real samples. We have used this probe to detect Fe$^{3+}$ in industrial effluents, tap water, and Iron tablet (Irozorb) and Iron syrup (Orofex XT). In all the real samples significant fluorescence dependence on concentration was observed with a good linear response. Importantly, trace levels of metal ions could be detected using this probe RG5NC which ensured the potential analysis of Fe$^{3+}$ in real sample sources. The obtained results were also validated by AAS and ICP-OES methods respectively. It is important to mention that testing of commercially available Iron syrup and Iron tablet for Fe$^{3+}$ was examined using this probe. Each sample was analyzed with three of their replicates. The estimated detection limits have been tabulated in Table 1.

4.2 Response of RG5NC towards human blood serum

We have investigated the use of the probe towards the detection of Fe$^{3+}$ ion in blood serum samples. In order to investigate the sensitivity of the probe RG5NC towards Fe$^{3+}$ similar titration protocol was followed. Even in high serum condition the probe showed effective detection response towards Fe$^{3+}$. A linear change in the emission with good linear regression at 553 nm was observed. The control experiment was also performed to study the interaction of the probe RG5NC with human blood serum under similar conditions. The obtained values detected from this method matched well with the mM values of Fe$^{3+}$ ion given in certified reference material (CRM). The results were also validated by AAS and ICP-OES methods respectively. The experiments were conducted in triplicates. The determined Fe$^{3+}$ content in human blood samples have been tabulated in Table 2.

5. Fast track detection of Fe$^{3+}$ using test strips

Assuring purity of drinking water and consumable food materials in remote areas is a challenging task where laboratory facilities are not available. Hence portable test strips were prepared for quick on-field detection of Fe$^{3+}$ ions. For this purpose test strips were soaked in MeCN solution of RG5NC and then dried in air before they were used for detection of Fe$^{3+}$ in water. A distinct color change was observed immediately upon dipping the test strips in Fe$^{3+}$ ion solution (Fig. 6). Presence of Fe$^{3+}$ selectively changed the test-strips color to orangish.

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Table 1 Determination of Fe$^{3+}$ in real samples

<table>
<thead>
<tr>
<th>Samples</th>
<th>Fe$^{3+}$ found by AAS method (M)</th>
<th>RSD (%)</th>
<th>Fe$^{3+}$ found by present method (M)</th>
<th>SE</th>
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<tr>
<td>Iron syrup</td>
<td>$9.12 \times 10^{-5}$</td>
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<td>1.62</td>
<td>$6.28 \times 10^{-4}$</td>
<td>1.53</td>
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<td>Tap water</td>
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<td>2.54</td>
<td>$1.55 \times 10^{-4}$</td>
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Table 2 Determination of Fe$^{3+}$ in various human blood serum samples using RG5NC

<table>
<thead>
<tr>
<th>Samples</th>
<th>Fe$^{3+}$ found by ICP_OES method (M)</th>
<th>RSD (%)</th>
<th>Fe$^{3+}$ found by present method (M)</th>
<th>SE</th>
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<tr>
<td>CBRM1</td>
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<td>3.64</td>
<td>$5.67 \times 10^{-2}$</td>
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<tr>
<td>CBRM2</td>
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<tr>
<td>CBRM3</td>
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<tr>
<td>BS1</td>
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<td>$5.87 \times 10^{-2}$</td>
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<td>BS2</td>
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<td>$5.85 \times 10^{-2}$</td>
<td>1.84</td>
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*CBRM: certified blood reference material, BS: blood serum samples.*
red. This newly developed sensor system provides an alternative method to confirm the nature as well as the extent of metal ion induced toxicity in natural water sources.

A comparison of the applicability and analytical section of this probe with some of the previous reports in terms of their solubility and detection limit is shown in Table 3.

### Table 3 Comparison of our probe performance with various other reported probes

<table>
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<tr>
<th>Probe</th>
<th>Detection limit</th>
<th>Reference</th>
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<tr>
<td>1.42 \times 10^{-6} \text{ M}</td>
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<tr>
<td>0.03 \times 10^{-6} \text{ M}</td>
<td>32b</td>
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<td>0.29 \times 10^{-6} \text{ M}</td>
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<td>17 \times 10^{-8} \text{ M}</td>
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<td>1.18 \times 10^{-8} \text{ M}</td>
<td>32e</td>
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<tr>
<td>8.2 \times 10^{-9} \text{ M}</td>
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6. Conclusion

The synthesized probe showed an outstanding sensitivity and selectivity for \( \text{Fe}^{3+} \) in human blood serum samples and excellent uptake by cells for imaging applications. The detection limit was about 8.2 nM. Spectral changes were observed to be reversible with respect to spirolactam ring opening. Confocal laser scanning microscopy experiments showed that RG5NC could be used to detect \( \text{Fe}^{3+} \) in live cells. The probe sensitivity of metal ion recognition was investigated in human blood serum. The detection as well as the discrimination of this toxic metal ion was also achieved in real samples at nano molar level. In addition, we have also demonstrated this detection of \( \text{Fe}^{3+} \) ions using portable test strips. The significance of this study lies in applying such fluorescent probes for rapid on-field detection of \( \text{Fe}^{3+} \) in human blood serum and live cells.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

The authors acknowledge Dr Vijayalakshmi S, Scientist, IGCAR, Kalpakkam, Tamilnadu, India for constant support and timely
help. The BRNS, DAE, India for financial assistance (Project Sanction No. 37(2)14/06/2014-BRNS) and the Institute of Excellence, Vijnana Bhavana, University of Mysore, India for providing the Nuclear Magnetic Resonance (NMR) and Liquid Crystal Mass Spectrometry (LC-MS) facility also TUV India Private Limited, Bengaluru, India for providing ICP-OES instrumentation facility.

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