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First-in-class hydrazone-pyrazoline sensors for selective detection of Zn²⁺, Cd²⁺, and Hg²⁺ in aqueous environments†

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Six first-in-class hydrazone-pyrazoline fluorescent sensors that selectively detect Zn^{2+} , Cd^{2+} , and Hg^{2+} at three distinct wavelengths in aqueous environments are reported. These novel hybrid sensors are synthesized in just two steps from inexpensive, commercially available materials, enabling rapid generation and screening for desirable photophysical properties. Lead sensor **P1** was validated for *in situ* detection of group 12 metals in river and pond water using a portable, low-cost device. Crucially, this system enabled rapid, naked-eye detection of toxic metals directly in the environment without the need for expensive instrumentation or mains electricity. These findings establish a new direction in pyrazoline sensor research and set a high benchmark for future sensor development.

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Introduction

The first three metals in group 12 of the periodic table are zinc, the second most abundant transition metal in the human body critical to life,1 cadmium, a highly toxic industrial pollutant linked to numerous cancers,2 and mercury, with wellestablished neurotoxicity.3 The European Union (EU) maximum drinking water limits for cadmium and mercury are 44 nM and 5 nM respectively.4 The World Health Organisation (WHO) has a recommended zinc drinking water limit of 46 μM.5 The regular monitoring of group 12 metals in aqueous environments such as drinking water and external water courses is paramount. The ability to distinguish between Zn²⁺, Cd²⁺, and Hg²⁺ in aqueous environments is a major challenge requiring expensive and complex equipment not widely available, for example, ICP-MS. Fluorescence spectroscopy offers several advantages6 over traditional techniques including high selectivity, low limits of detection (LoD), and rapid analysis. Fluorescent sensors are "turned on" when the analyte increases fluorescence emission $(\lambda_{em})^7$ or "turned off" when the analyte decreases λ_{em} .8 Inhibition of photoinduced electron transfer (PET)9 or the chelation enhancement effect (CHEF)10 can increase λ_{em} . Decreased λ_{em} may result from fluorescence resonance energy transfer (FRET)11 or chelation-enhanced quenching effect (CHEQ).12 While numerous fluorescent sensors have been reported for Zn2+ and Cd2+,13 few can differentiate between these metals at different λ_{em} .¹⁴ The pyrazoline scaffold15 (blue in Fig. 1) is highly advantageous due to

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its unique fluorescence properties16 and modular structure from readily accessible chalcone precursors. 17 Chalcones, while utilized as precursors for pyrazolines, have also been reported to display useful sensing properties.18 Pyridine is a wellestablished metal chelator often integrated into fluorescent sensors for Zn²⁺, Hg²⁺ and Fe³⁺ detection. 19 Simple pyrazoline A displayed a "turn on" response only in the presence of Zn²⁺ and Cd2+;20 B is a "turn on" sensor21 for Fe3+/Fe2+ whereas C displayed a "turn off" response in the presence of Fe³⁺/Fe²⁺. 21 Pyrazole D (red in Fig. 1)22 demonstrated that an additional acetyl group greatly improved Zn2+/Cd2+ selectivity. A-D reveal that structural complexity is not essential for complex functionality20-22 highlighting the potential of these scaffolds in sensor development. During a recent pyrazoline synthesis study (see ESI S1†) side product P1 was identified (Scheme 1), isolated, and discovered as a "turn on" sensor for Zn^{2+} at λ_{em} 560 nm and Cd²⁺ at 510 nm and a "turn off" sensor for Hg²⁺ at $\lambda_{\rm em}$ 460 nm. This unexpected side product is a highly attractive first-in-class hydrazone-pyrazoline multi-analyte group 12 sensor in aqueous environments.

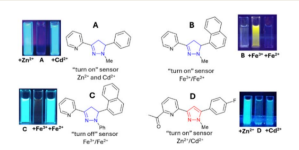


Fig. 1 Pyrazoline and pyrazole sensors; images reproduced from ref. 20–22 with permission from RSC.

[†] Electronic supplementary information (ESI) available. See DOI: https://doi.org/10.1039/d5ay00639b

Scheme 1 Unexpected synthesis of side product P1 from C1.

Experimental

Chemicals and instruments

Chemicals, solvents and reagents were purchased from commercial sources and used without further purification. PE refers to petroleum ether, bp 40–60 °C. Spectroscopy was performed with CHROMASOLV® gradient grade acetonitrile for HPLC, ≥99.9%, from Sigma-Aldrich. The metal complexes used in this study were LiCl, NaCl, KCl, CaCl₂, MgCl₂, CuCl₂, NiCl₂, ZnCl₂, CdCl₂, RuCl₃, CoCl₂, MnCl₂, PbCl₂, ZnCl₂ and HgI₂.

UV/vis spectroscopy was performed on an Agilent Cary5000 using quartz cuvettes with a 1 cm pathlength using HPLC grade MeCN, in the 250–500 nm range with 0.2 s dwell time. Detector switchover occurred at 350 nm. FTIR spectroscopy was performed on a Bruker VERTEX 70 spectrometer. Fluorescence spectroscopy was performed on an Edinburgh Instruments FLS1000 with a xenon excitation source, 2 nm bandwidths for both excitation and emission monochromators, a scan speed of 1 nm and a dwell time of 0.2 s. Fluorescence quartz cuvettes with a 1 cm pathlength were used throughout with HPLC grade MeCN. A 100 watt 365 nm Analytik Jena high intensity UV lamp was used to image the sensors.

Synthesis of chalcones C1-C4

The following general procedure was followed (see the ESI† for more detailed information), using a method adapted from a previous synthesis, 22 6 mmol 2,6-acetylpyridine was added to a stirred solution of 3.0 mmol aldehyde in MeOH followed by the addition of 3.0 mmol NaOH and stirring was continued at room temperature. After 24 hours the precipitate was filtered, washed with copious amounts of cold $\rm H_2O$ collected and dried to afford the desired chalcone without further purification.

Synthesis of hydrazone-pyrazolines P1-P6

The following general procedure was followed: 4.0 mmol of 2-hydrazinopyridine was added to a stirred solution of 1.0 mmol of required chalcone in 50 mL MeOH and heated to 60 °C. After 24 hours, the solvent was removed under reduced pressure, 100 mL H_2O was added and it was extracted into 3 \times 50 mL ethyl acetate. The ethyl acetate fractions were combined and the solvent was removed under reduced pressure to give an oil which was then purified by column chromatography with 6:4 ethyl acetate: petroleum ether to afford the desired product.

Results and discussion

Chalcones C1-C4 were prepared using a previous procedure²² in acceptable yields (13-60%). A direct chalcone to hydrazone-

Scheme 2 Synthesis of hydrazone-pyrazoline series P1-P4.

pyrazoline synthesis was developed to give novel compounds **P1-P4** (Scheme 2) in good yield (26-61%).

This two-step synthesis from inexpensive commercially available starting materials unlocks a new class of previously unexplored pyrazoline sensors. Furthermore, the modular nature of the pyrazoline scaffold enables a high degree of modification unlocking a deeper understanding of their sensing properties. With four novel hydrazone-pyrazolines in hand, we investigated their photophysical properties against the group 12 metals using well-established protocols.²²⁻²⁴ UV/vis studies confirmed that P1 undergoes chelation with both Zn²⁺ and Cd²⁺ with the formation of a new band at 285 and 290 nm respectively (Fig. 2A and B). A ¹H NMR titration study for P3 in the presence of the group 12 metals indicated broadening and downfield movement for multiple signals (Fig. 2C for P3 with Hg²⁺). The broad singlet at 8.54 ppm attributed to the hydrazone proton H^a shifted to 9.31 ppm on the addition of 2.0 equivalents Hg²⁺. Significant shifts in the remaining protons were also observed; for example, the pyridine signal from Hb moved from 7.79 ppm to 7.99 pm and the aryl singlet for H^c moved from 6.54 to 6.68 ppm. This is a well-established

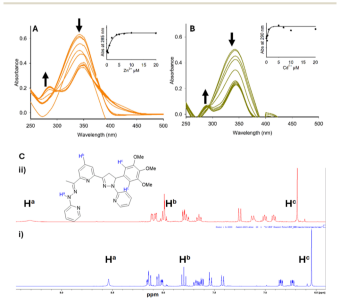


Fig. 2 UV/vis study of P1 (20 μ M, MeCN) with 0–20 equivalents of Zn²⁺ (panel A) and Cd²⁺ (panel B). ¹H NMR studies of P3 (9 μ M, deuterated MeCN), (panel C) (i) P3 only and (ii) +2.0 equivalents of Hq²⁺.

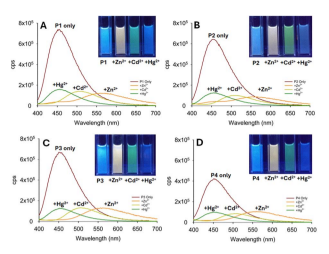
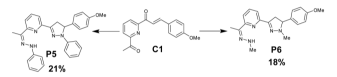


Fig. 3 Fluorescence studies of P1–P4 (A–D) 15 μ M, 7:3 MeCN: H₂O at $\lambda_{\rm ex}$ 285 nm, on addition of 5.0 equivalents of the indicated metal; cps is counts per second. Cuvette images were obtained under the irradiation of a 100 W $\lambda_{\rm ex}$ 365 nm lamp.



Scheme 3 Synthesis of hydrazone-pyrazoline P5-P6.

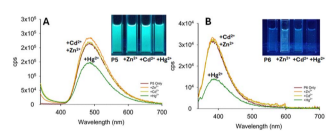


Fig. 4 Fluorescence studies of P5 (A) and P6 (B) 15 μ M, 7:3 MeCN: H₂O at λ_{ex} 285 nm; cps is counts per second. Cuvette images were obtained under the irradiation of a 100 W λ_{ex} 365 nm lamp.

observation with previous pyrazoline sensors undergoing chelation. $^{20-23,24a,b}$

The fluorescence response of **P1** (φ_f 4.0%, τ 3.4 ns) was confirmed with the formation of new emission bands at 560 nm with Zn^{2^+} (φ_f 1.0%, τ 14 ns) and 510 nm with Cd^{2^+} (φ_f 1.0%, τ 13 ns) in 7:3 MeCN: H_2O (Fig. 3A). On addition of Hg^{2^+} a "turn off" response at λ_{em} 460 nm (φ_f < 1.0% τ 4.0 ns) is observed. Of note is that the response is retained in an aqueous solution demonstrating the real-world application of this sensor. A similar fluorescence profile was obtained with **P2–P4** suggesting that substitution on the aryl ring was well-tolerated a range of both electron donating and electron withdrawing substituents. To determine the importance of hydrazone, two further pyrazolines were synthesised; **P5** using phenyl hydrazine and **P6** with methyl hydrazine (Scheme 3). **P5** displayed no

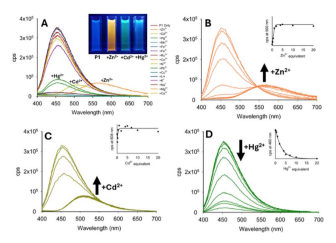


Fig. 5 (A) Fluorescence studies of P1 15 μ M, 7:3 MeCN: H₂O at λ_{ex} 285 nm. (Panels B–D) Titration studies for Zn²⁺, Cd²⁺ and Hg²⁺, respectively; cps is counts per second. Cuvette images were obtained under the irradiation of a 100 W λ_{ex} 365 nm lamp.

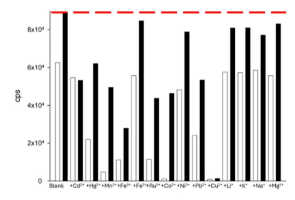


Fig. 6 Competition assay for P1 with Zn²+ in the presence of a range of interferents. The white bar represents the sensor (20 $\mu M,~7:3$ MeCN: H₂O) with 5.0 equiv. of the indicated cation; the black bars are the same with 5.0 equiv. of Zn²+ after equilibrating for 2 min. λ_{ex} 285 nm with λ_{em} 560 nm; the red line indicates the "turn on" response with Zn²+ only, and cps is counts per second.

distinguishing features suggesting that the pyridine ring in P1–P4 was critical for group 12 selectivity. P6 with methyl units displayed insignificant fluorescence on addition of the group 12 metals suggesting that an aromatic unit was required. The contrast between P1 and P5 is striking confirming that pyridine units are essential for group 12 selectivity (Fig. 4).

P1 was selected as the lead compound and submitted for further investigation. A titration study confirmed λ_{em} at 560 nm (yellow emission) which peaked with 5.0 equivalents of Zn^{2+} (Fig. 5B). A Zn^{2+} limit of detection (LoD) of 6.4 μM was calculated which is well below the WHO recommended drinking water limit. A similar study with Cd^{2+} confirmed λ_{em} at 510 nm (green emission) which reached a plateau with 5.0 equivalents of Cd^{2+} (LoD 2.4 μM). The 50 nm difference in λ_{em} between Zn^{2+} and Cd^{2+} is significant with very few examples reporting such a distinction between two group 12 metals. To our surprise Hg^{2+} generated an opposite response, a "turn off" response, with a decrease in λ_{em} at 460 nm, Hg^{2+} LoD 10.0 μM. These LoD

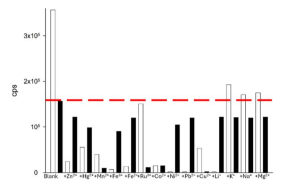


Fig. 7 Competition assay for P1 with Cd²⁺ in the presence of a range of interferents. The white bar represents the sensor (20 µM, 7:3 MeCN: H₂O) with 5.0 equiv. of the indicated cation; black bars are the same with 5.0 equiv. of Cd^{2+} after equilibrating for 2 min. λ_{ex} 285 nm with λ_{em} 500 nm.

values agree with those of previously reported pyrazoline sensors for Zn2+ and Cd2+ (ESI S6†). Competition assays were performed to determine the response of P1 in the presence of a range of interferents (Fig. 6 for **P1** with Zn²⁺).

Several paramagnetic metals including Fe³⁺, Ru³⁺, Co²⁺, and noticeably Cu2+ disrupted the "turn on" response to Zn2+ at 560 nm (Fig. 6). This is a common phenomenon for pyrazolines with paramagnetic metals. 13b,c,22,23 Interestingly, the biological group 1 and 2 metals, Li⁺, K⁺, Na⁺ and Mg²⁺, resulted in very minor disruption suggesting that P1 could be used for the monitoring of Zn2+ in living systems high in these metals, an active research area.13a,26 A similar profile was observed with Cd2+; paramagnetic metals Mn2+, Ru3+, Co2+ and Cu2+ hampered λ_{em} 510 nm but a good response remained with the group 1 and 2 metals (Fig. 7). A further assay of P1 with Hg²⁺ showed that paramagnetic metals disrupted the "turn off" response (Fig. 8). Under these conditions, the group 1 and 2 metals did not significantly influence the "turn off" response to Hg²⁺ suggesting that **P1** could be a useful sensor for the "turn

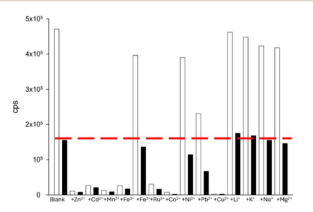


Fig. 8 Competition assay for P1 with Hg2+ in the presence of a range of interferents. The white bar represents the sensor (20 μ M, 7:3 MeCN: H₂O) with 5.0 equiv. of the indicated cation; the black bars are the same with 5.0 equiv. of Hg²⁺ after equilibrating for 2 min. λ_{ex} 285 nm with λ_{em} 460 nm; the red line indicates the "turn off" response with Hg²⁺ only.

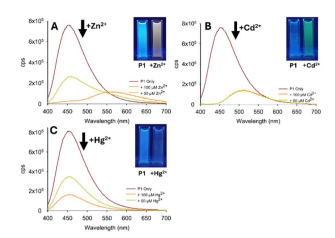


Fig. 9 Group 12 metal triggered fluorescence response of P1 (20 μ M, 7:3 MeCN: H_2O) in tap water; Zn^{2+} (panel A), Cd^{2+} (panel B) and Hg^{2+} (panel C). Cuvette images under the irradiation of the 100 W $\lambda_{\rm ex}$ 365 nm lamp

on" detection of Zn2+ at 560 nm and the "turn off" detection of Hg²⁺ at 460 nm in biological systems.

To determine the real-world application of P1 in large-scale monitoring, several studies of a variety of water sources were conducted including drinking water (tap and mineral water see Fig. 9 and ESI S5†). A strong response to 100 μM Zn²⁺ in tap water (Fig. 9) with a visible colour change from blue to yellow was observed at λ_{ex} 365 nm (Fig. 9A inset). A similar study with Cd²⁺ in tap water produced a more noticeable response at 50 μM Cd^{2+} (Fig. 9B) and a green colour change under λ_{ex} 365 nm irradiation (Fig. 9B inset). The "turn off" response to Hg2+ was detectable in tap water (Fig. 9C) with a reduction in λ_{em} at 460 nm with both 50 μ M and 100 μ M Hg²⁺. Mineral water (see ESI S5†) yielded a comparable result. The visible colour change on the addition of Zn²⁺ (yellow), Cd²⁺ (green) and Hg²⁺ (blue) suggests that P1 could form a prototype for a low-cost and portable field-based device. A low-cost prototype was constructed consisting of a portable excitation source and a suitable dark box (ESI S8†).25 Samples from two environmental

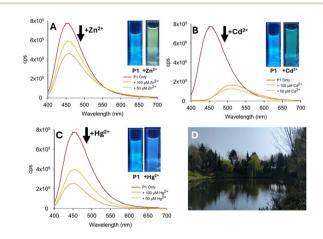


Fig. 10 Group 12 metal triggered fluorescence response of P1 (20 μM, 7:3 MeCN: H_2O) in pond water; $+Zn^{2+}$ (panel A), $+Cd^{2+}$ (panel B), +Hg²⁺ (panel C) and location (D).

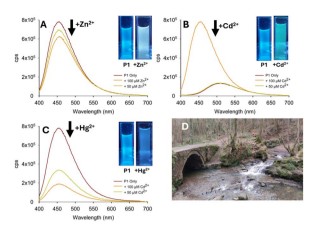


Fig. 11 Group 12 metal triggered fluorescence response of P1 (20 μ M, 7:3 MeCN: H₂O) in river water; +Zn²⁺ (panel A), +Cd²⁺ (panel B), +Hg²⁺ (panel C) and location (D).

locations (river and pond water) were taken and analysed in the laboratory and *in situ* using a portable device (Fig. 10 for pond water).

At two different concentrations, 50 μ M and 100 μ M of Zn²⁺, Cd2+, and Hg2+, a fluorescence response was triggered analogous to the drinking water study. The decrease in λ_{em} with Zn^{2+} (Fig. 10A) was less than expected possibly due to additional interferents present in the environment, possibly sediment or bacteria (Fig. 10A). In the presence of Cd^{2+} the change in λ_{em} is noticeable and a visible change to green is observed (Fig. 10B inset). A decrease in λ_{em} at 460 nm was observed on addition of Hg²⁺ (Fig. 10C inset) confirming that **P1** can distinguish Cd²⁺ from Hg²⁺ in the field that can be observed with the naked eye. A similar result was obtained in river water (Fig. 11). This study confirmed that P1 can be used outside of a laboratory environment with naked eye detection of the group 12 metals in both stagnant pond and free flowing river waters. To confirm the robustness of P1, a reversibility study was performed which confirmed that P1 can be used for multiple cycles for Zn²⁺ detection (Fig. 12A). The proposed binding mechanism of P1 with Zn²⁺ (Fig. 12B) is 1:1 and agrees with previous studies on A-D and the Job plot (ESI S7†). 20-23 While hydrazone units have been incorporated into multi-analyte fluorescent sensors previously,27 P1-P4 are the first reported sensors combining hydrazone and pyrazoline units which distinguish Zn²⁺, Cd²⁺, and Hg^{2+} at different λ_{em} in aqueous environments.

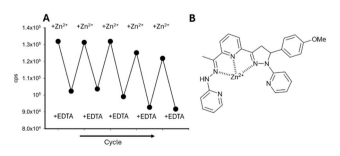


Fig. 12 A reversibility study for P1 in the presence of multiple cycles of Zn^{2+} and EDTA (panel A) and the proposed binding mode of P1 with Zn^{2+} (panel B).

First-in-class sensors
 Aqueous environments
 Can distinguish Zn²+/ Cd²+/ Hg²+
 Naked eye detection
 Validated for rapid use in rural areas
 Sets a new benchmark

Pyridine required for Group 12 selectivity

Fig. 13 Summary of P1-P6; EDG is the electron donating group and EWG is the electron withdrawing group.

Conclusions

Six first-in-class hydrazone-pyrazoline sensors were synthesised with P1-P4 displaying excellent selectivity towards group 12 metals. The aryl ring was well-tolerated to both electron donating and withdrawing substituents; however, substitution of pyridine for a phenyl unit in P5 abolished group 12 selectivity (Fig. 13). Substitution of the aryl ring on hydrazone and pyrazoline for methyl units in P6 significantly disrupted the fluorescence intensity. P1 was selected for further investigation and while a few heavy metals were significant interferents to the fluorescent response for P1, its ability to detect group 12 metals in real-world samples was confirmed. Combining P1 with a lowcost portable device enabled rapid visual detection by the naked eye of the 12 metals in rural environments without expensive apparatus or mains electricity. The group 1 and 2 metals exhibited very little influence on the fluorescence response indicating that P1 could also be a highly useful sensor for the selective detection of Zn²⁺ and Hg²⁺ in biological systems; for example cell culture and living systems.26 This highly unexpected discovery establishes a new subset of pyrazoline sensors, hydrazone-pyrazoline hybrid sensors. These hybrid sensors composed of both hydrazone and pyrazoline functional units set a high benchmark for future group 12 sensors. Further work is underway to transform this discovery into a second generation of sensors that retain the group 12 metal selectivity of P1 but with improved properties (increased λ_{em} intensity and reduced interference). The high tolerance of the aryl R¹ group to substitution enables the attachment of these sensors to solid substrates as reported previously by Magri et al.28 This provides valuable applications for enhanced sensor design and molecular logic gate development.29 These are priority objectives for ongoing work and will be reported in due course.

Data availability

The data supporting this article have been included as part of the ESI.†

Author contributions

Alexander Ciupa designed, synthesized, characterised, and performed all spectroscopy studies and authored the manuscript.

Conflicts of interest

There are no conflicts to declare.

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