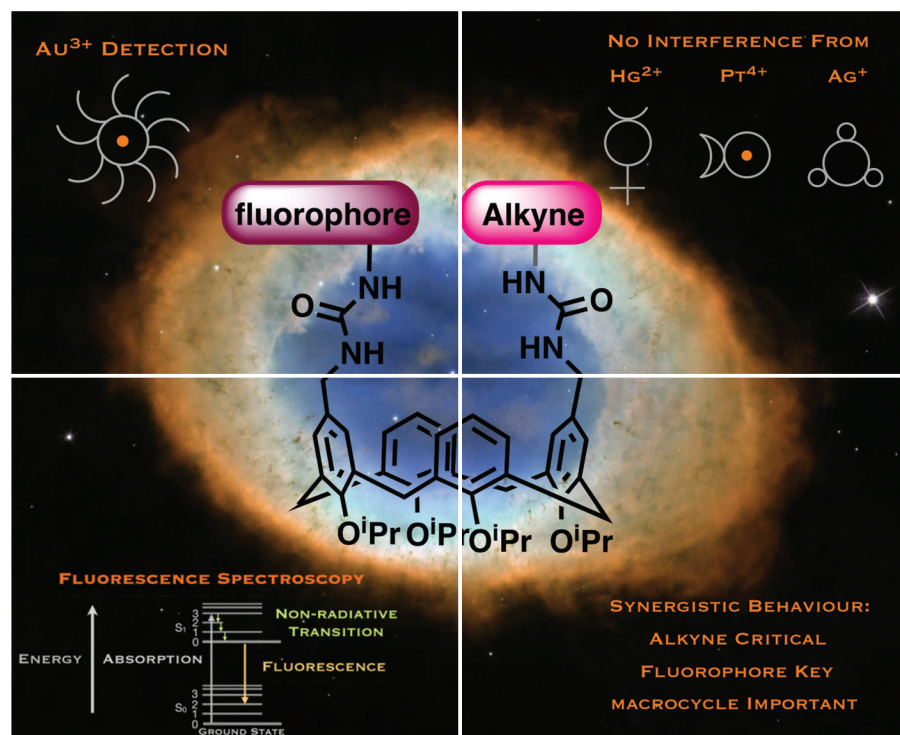


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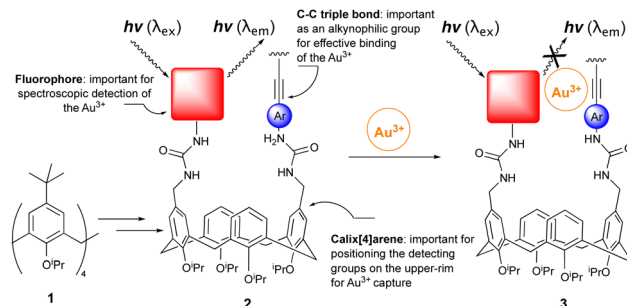
Upper-rim functionalised calix[4]arenes for chemoselective Au³⁺ detection†Sean P. Bew,^a Sunil V. Sharma^b and William H. Gardiner^cA unique, readily synthesised, upper-rim 1,3-difunctionalised calix[4]arene is reported; equipped with a fluorophore and an alkyne it mediates the efficient chemoselective detection of Au³⁺. Its ability to detect Au³⁺ is not perturbed by Au⁺ or excess competing and/or contaminating metal e.g. platinum, cadmium, mercury, silver, sodium, magnesium or potassium cations.

Introduction

Owing to its operational simplicity and sensitivity, fluorescence spectroscopy is a powerful method for metal cation detection. Appending fluorescent and metal recognition ‘antennas’ onto easily synthesised and readily functionalised calix[4]arenes affords motifs with the potential to chemoselectively detect and differentiate between metal cations.¹ A wide variety of metal detectors have previously been developed. Indeed, many are utilized in diverse applications across the breadth of science and includes chemistry, biology, biochemistry, and medicine. A typical, but non-exhaustive list of uses includes detecting polluting toxic metals² e.g., lead, mercury, cadmium, or the development of motifs for cell microscopy and the *in vivo* location of biologically important metal ions such as iron, calcium, zinc, sodium, potassium and copper³ as well as the non-invasive imaging of calcium in the human brain.⁴

Recent years have witnessed a sharp increase in the use of gold in developing new methodology based on, for example, homogeneous and heterogeneous catalysis. It is, however, not the sole preserve of chemists. As noted, its potential has been recognized, explored, and unlocked in biology, medicine, and physics. For example, Au₅₅ clusters ([Au₅₅(Ph₂PC₆H₄SO₃)₁₂]¹²⁻) are used to visualize cell component⁵ and in medicine, Tauredon, Solganal, Sanocrycin, and Auranofin are used to fight asthma, HIV, malaria, lesions, cancer, and arthritis. Thus, although gold is in many respects ‘compatible’ it can have adverse effects on the environment,⁶ and in biological systems Au³⁺ metallodrugs can be reduced to Au⁺.⁷ For these

reasons and due to its increased use, the development of new modalities and protocols for differentiating Au³⁺/Au⁺ are urgently required. Ideally, these must be of relatively low cost, easily synthesised, chemoselective and, importantly, they should not be perturbed by other metal contaminants e.g., mercury, silver, platinum, and cadmium. No reaction-based fluorescent probes for gold detection have been developed which proceed *via* a ‘turn off’ mechanism and use readily synthesised calix[4]arenes derived from **1** (Scheme 1).⁸ Exploiting our expertise and knowledge of calix[4]arenes and their upper-rim chemical functionalization⁹ we were intrigued by the possibility of transforming scaffold **1** into a chemospecific Au³⁺ metal-binder (**2**) which specifically presents, facilitates and, subsequently, reports its presence *via* adjacent alkyne and dansyl ‘antenna’ (cf. **3**, Scheme 1). Attracted by the chemical and operational simplicity of our system, the straightforward synthesis of entities based on **3** uses easily generated starting material **1** and proceeds *via* a simple, readily observable ‘turn-off’ mechanism (*i.e.*, **2** → **3**) when the metal is bound/detected (Scheme 1). With strategies to generate upper-rim functionalised calix[4]arenes in hand, key to our proposed application is the π -aurophilic alkyne which chemoselectively ‘captures’ the Au³⁺. Substantiating this as a mode of action, Ultimoto and Fukuda reported internal alkynes react readily

Scheme 1 Overview of Au³⁺ detection (**3**) using calix[4]arene **2**.^aSchool of Chemistry, UEA, Norwich Research Park, Norwich, NR4 7TJ, UK.
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St Andrews, Scotland, UK^cEvotec UK Ltd, Innovation Drive, Milton Park, Abingdon, Oxfordshire, OX14 4RZ, UK† Electronic supplementary information (ESI) available. See DOI: <https://doi.org/10.1039/d3qo00021d>

Research Article

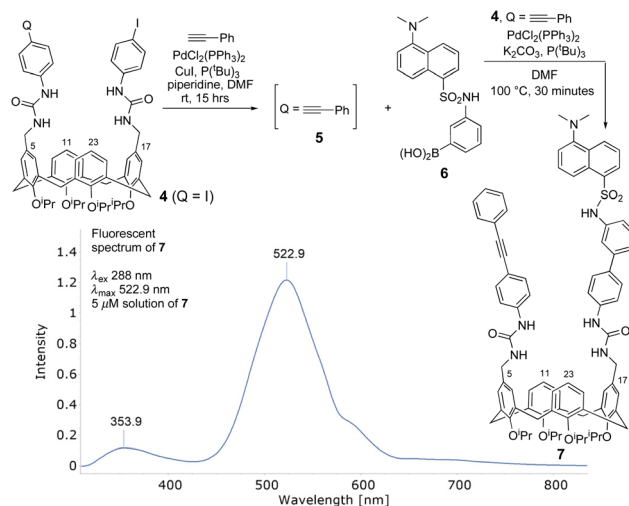
with a catalytic source of Au^{3+} in the form of sodium tetrachloroaurate which in aqueous methanol transforms an alkyne into a ketone (17, Scheme 3).¹⁰ With the gold 'captured' the conformational mobility of the calix[4]arene¹¹ helps facilitate the fluorescent dansyl group to move closer and interact with the metal whilst 'reporting' its presence *via* fluorescent quenching (Scheme 1).

With this rationale in mind we were also convinced of the merits of generating entities based on 2 due to the widespread use of the calix[4]arene motif in alternate metal detection regimes which, when dovetailed with the known ability of gold salts to mediate efficient internal nonradiative relaxation decay, results in fluorescent quenching (FQ).¹² Supporting our approach to gold detection are the abundant applications of the calix[4]arene motif in metal recognition but, surprisingly, the lack of any specific calixarene-based Au^{3+} modality. Thus, although there are many examples of *cone*-confined calix[4]arenes with lower-rim metal cation sensing properties,¹³ this fact contrasts sharply with the handful of known metal sensing upper-rim functionalized calix[4]arenes.¹⁴

Results and discussion

We wanted to synthesize upper-rim functionalized 2 with three structural components, namely an alkyne, dansyl and calix[4]arene macrocycle, and test it in a proof-of-concept study. Initiating this required an efficient and straightforward route to bis-5,17-*N*-aryleurea derived calix[4]arene 4. Two synthetic routes were envisaged. The first used commercially available 4-iodophenyl isocyanate and the easily synthesized bis-5,17-aminomethylene derived *cone*-confined calix[4]arene (not shown).¹⁵ Their reaction in THF at ambient temperature afforded 4 in a 90% yield which was, after methanol trituration, pure enough to use in the next step. An alternative ionic hydrogenation route¹⁶ used triphenylsilane, TFA and two commercially available starting materials *i.e.*, bis-5,17-diformylcalix[4]arene and 4-iodophenylurea. This afforded 4 in an unoptimised 63% yield. Installing the alkyne (4 → 5) was straightforward. A Sonogashira coupling between 4 and 1 equivalent of phenylacetylene was catalysed by tri-*tert*-butylphosphine (20 mol%), copper(I) iodide (20 mol%) and bis (triphenylphosphine)palladium(II) dichloride (7 mol%).

The desired mono-phenylacetylene-derived 5 (Scheme 2) was used 'as is' in the next step [an estimated 15% of the corresponding di-phenylacetylene adduct (not shown) was also formed]. A subsequent Suzuki–Miyaura coupling with 3-(dansylamino)phenylboronic acid (6) catalysed by bis(triphenylphosphine)palladium(II) dichloride (5 mol%), tri-*tert*-butylphosphine (10 mol%) and potassium carbonate afforded, after chromatographic purification, the desired product 7. We propose the detection capabilities of 7 (*vide infra*) rely on the capacity of the dansyl group and π -aurophilic alkyne to 'communicate' with the gold only when all three are in close proximity (*cf.* 3, Scheme 1). Clearly, it was important to verify the presence of the dansyl and alkyne groups, as well as the con-



Scheme 2 Synthesis of upper-rim bis-5,17-functionalised calix[4]arene 7 from bis-5,17-*N*-urea-derived calix[4]arene 4.

formation of the calix[4]arene. Frustrated by its reluctance to afford crystals suitable for X-ray diffraction, we analyzed the ¹³C-NMR data associated with the bridging methylene's which, Mendoza *et al.* report, allow the conformation of a calix[4]arene to be established.¹⁷ The ¹³C-NMR signal at 31 ppm confirmed 7 resided in a *cone*-conformation. Additional physicochemical studies using FT-IR and mass spectrometry verified the dansyl and alkyne groups were present. A fundamental reason for incorporating and using the dansyl group is its strong fluorescence signal and long emission wavelength. Probing the photophysical properties of 7 a 5 μM solution had an excitation absorption maximum at 288 nm (λ_{ex}) which generated a strong fluorescent emission signal (λ_{em}) with a λ_{max} of 522.9 nm. A weaker signal was observed at 353.9 nm (Scheme 2). Confirming the signal at 522.9 nm was broadly like other λ_{em} of alternative upper- and lower-rim dansyl-derived calix[4]arenes our value is in general agreement with literature reports of λ_{max} at 538 nm and 520–575 nm respectively.¹⁸ Recording additional spectra of 7 every two minutes for 45 minutes we established the emission (λ_{em}) intensity remained constant. This validated the fluorophore was photochemically stable.

It was important to assess if 7 was susceptible to non-chemoselective metal mediated FQ. That is, does it generate false positives? Screening individual solutions of Na^+ , K^+ , Cs^+ , Ca^{2+} and Mg^{2+} salts for their effect, if any, on the strong λ_{em} signal would help answer this question. Adding 1–5 equivalents of aqueous sodium chloride (2.5 μM) afforded almost identical 'before' and 'after' λ_{em} spectra, even after 3 hours (*cf.* Scheme 2 with Fig. 1 in the ESI†). Similarly, 1–5 equivalents of a mixture of Li^+ , Na^+ , K^+ , Cs^+ , Ca^{2+} and Mg^{2+} salts (all at 2.5 μM) had little effect. There are reports that Group 12 metal cations FQ the dansyl group. For example, Bartsch *et al.* described a calix[4]arene appended on the lower-rim with two dansyl groups as a fluorogenic Hg^{2+} selective extractant which underwent FQ.¹⁹ Screening individual solutions of Group 12 salts [cadmium



acetate, mercury(II) chloride, hydrogen tetrachloroplatinate and silver tetrafluoroborate] had little effect on the photophysical properties of **7**.

Indeed, only with >15 equivalents and extended mixing times was partial FQ observed (see ESI†). Further stress testing its photophysical properties 1–5 equivalents of a mixture of Group 2 (Mg^{2+} and Ca^{2+}), Group 10 (Pt^{4+}) and Group 12 (Cd^{2+} and Hg^{2+}) salts had minimal effect on **7**'s λ_{max} , even after 3 hours (Fig. 1).

Au^{3+} and Au^+ are strong Lewis acids, indeed both can activate alkynes to nucleophilic attack with weak nucleophiles *e.g.*, water or simple alcohols.²⁰ Clearly, if **7** was not able to chemoselectively differentiate Au^{3+} and Au^+ its use would be severely limited. With this in mind we were delighted adding 5 equivalents of chloro(triphenylphosphine)gold(I) to **7** had little impact on the λ_{max} , even after 3 hours (Fig. 2).

With a reliable synthesis in hand and robust photophysical data confirming little evidence of FQ with either single or mixtures of metal salts, it was essential to establish if **7** was able to 'capture, sense and report'. Adding a solution of sodium tetrachloroaurate to a 5 μM solution of **7**, an almost instantaneous reduction in the λ_{max} signal was observed (Fig. 3). Increasing the equivalents from 1 to 2, 3 and 4 induced further fluorescent quenching and a reduction in the λ_{max} intensity (Fig. 3). The lack of FQ with **7** in a protic solvent when presented with Group 10 or 12 metal salts contrasts sharply with observations using the gold salt (Group 11); this

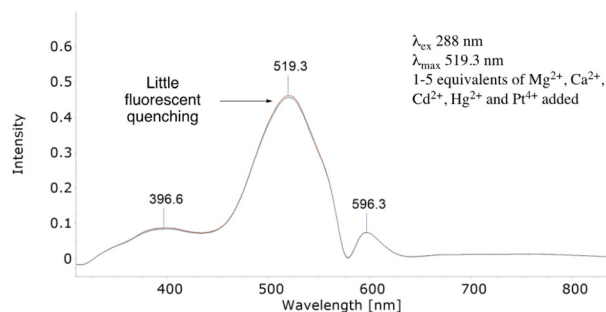


Fig. 1 The fluorescent spectrum above of **7** outlines how a mixture of Group 2, 10 and 12 metal salts had little FQ effect on **7**.

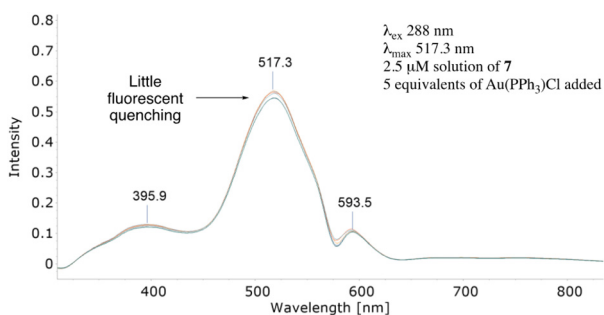


Fig. 2 Fluorescent spectrum of **7** with excess chloro(triphenylphosphine)gold(I) after 3 hours.

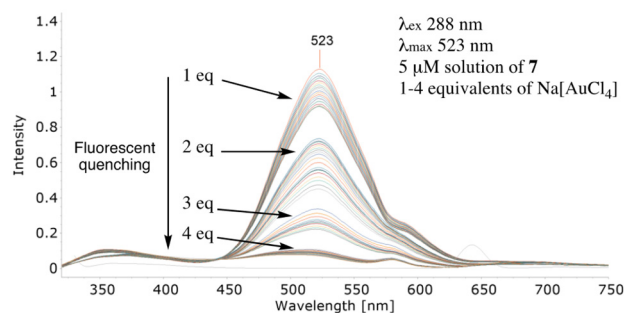


Fig. 3 Efficient fluorescent quenching of **7** appended with an upper-ribose dansyl and phenylacetylene group.

suggests key to FQ is the formation of significant and favourable interactions between the alkyne on **7** and the highly charged Au^{3+} .

Evidently, bifunctional calix[4]arene **7** readily and efficiently detected the gold salt in the absence of, potentially, competing metal salts. Critical to its wider application however is its ability to retain this property when presented with a mixture of metal salts. So, do externally added salts inhibit or complicate the use of **7**? Generating individual solutions of (2.5 μM) sodium chloride, potassium hexafluorophosphate, calcium chloride, magnesium chloride and cesium chloride (see ESI†) as well as a mixture (2.5 μM) of lithium chloride, sodium chloride, potassium hexafluorophosphate, cesium chloride, magnesium chloride and calcium chloride their addition (5 equivalents each) to **7** and, subsequently, sodium tetrachloroaurate (0–5 equivalents) allowed us to compare 'contaminated' (Fig. 4) with a 'non-contaminated' gold salt solution (Fig. 3). Fig. 4 clearly demonstrates the strong λ_{max} signal at 520.3 nm was quenched in a manner comparable to 'pure' sodium tetrachloroaurate. Further probing and testing the robustness of a 'contaminated' system employed a mixture of 20 equivalents of cadmium acetate, mercury(II) chloride and hydrogen tetrachloroplatinate. Comparing Fig. 4 with Fig. 5 their similarities validated the efficient FQ by Au^{3+} to be unaffected by extraneous, potentially, competing salts.

Thus far, the alkyne was assumed to be an integral component of **7**, confirming this was essential. Hydrogenation of **7** afforded the alkyne-free hydrogenated 'diphenylethane' adduct

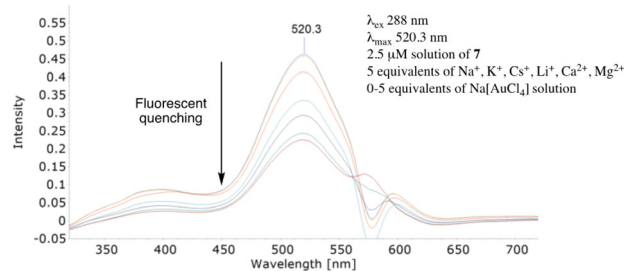


Fig. 4 Fluorescent quenching of **7** in the presence of excess Li^+ , Na^+ , K^+ , Cs^+ , Mg^{2+} and Ca^{2+} salts.



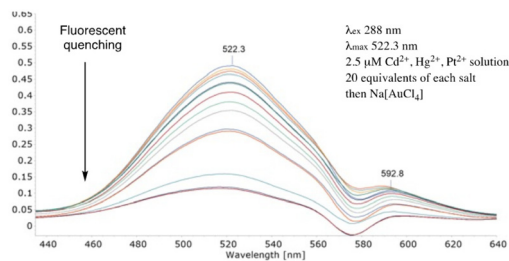


Fig. 5 Fluorescent quenching of **7** in the presence of 20 equivalents each of Cd^{2+} , Hg^{2+} and Pt^{4+} salts.

(not shown). Acquiring its fluorescent spectrum after 30 equivalents of sodium tetrachloroaurate had been added verified it did not undergo FQ. This result substantiates the key role the alkyne has in, presumably, 'capturing' and acting as an 'aurophile' bringing the metal into close proximity to the dansyl group generating 'alkyne- Au^{3+} -dansyl' interactions (*cf.* **3**) which, ultimately, initiate FQ. Seemingly, removing its ability to form a gold-alkyne complex is the first step since, by a process of elimination, the alkyne-free system seems unable to form gold-dansyl interactions which, if formed, would still, presumably, result in FQ.

Although the exact role the macrocycle occupies is not clear, its presence is evidently important when combined with the two upper-rim substituents. Testing for a 'macrocycle' effect 'linear' macrocycle-free **8** was generated from readily available starting materials. With its urea and dansyl modalities (Fig. 6) matching those of its cyclic 'cousin', **8** afforded the perfect opportunity to evaluate the importance of the calix[4]arene. Adding increasing equivalents (up to 20) of sodium tetrachloroaurate a slight reduction in FQ was observed. The propensity of sulfonamide groups to bind cationic metal centers is known.²¹ Thus, it seems plausible the observed FQ is associated with intermolecular functional group interactions derived from two molecules of **8** combining to form a weak 'dansyl- Au^{3+} -dansyl' complex like **9** (Fig. 6). The ability of upper-rim appended urea-derived calix[4]arenes to bind anions is well documented.²² Aqueous solutions of sodium tetrachloroaurate are known to undergo rapid equilibrium with water generating hydroxytrichloroaurate and chloride.²³ With its two upper-rim derived urea's **7** has the potential to generate multiple hydrogen-bonds and bind chloride (**10**) near the alkyne and dansyl gold binding motifs. Unfortunately, quantifying this using NMR and mass spectrometry proved inconclusive. Likewise generating crystals of **10** suitable for X-ray diffraction was unsuccessful. Our thoughts focused instead on the possibility that **7** when reacted with tetraethylammonium tetrafluoroborate affords a hydrogen bonded BF_4^- complex (*cf.* **11**) with the tetraethylammonium cation in close proximity to the tetrafluoroborate anion, a consequence of electrostatic attraction.

We anticipated the formation of **11** would reduce gold π -alkyne co-ordination and, consequently, diminish FQ. Mindful of reports outlining *N*-tetraalkylammonium cations residing in, largely unsubstituted upper-rim derived calix[4]

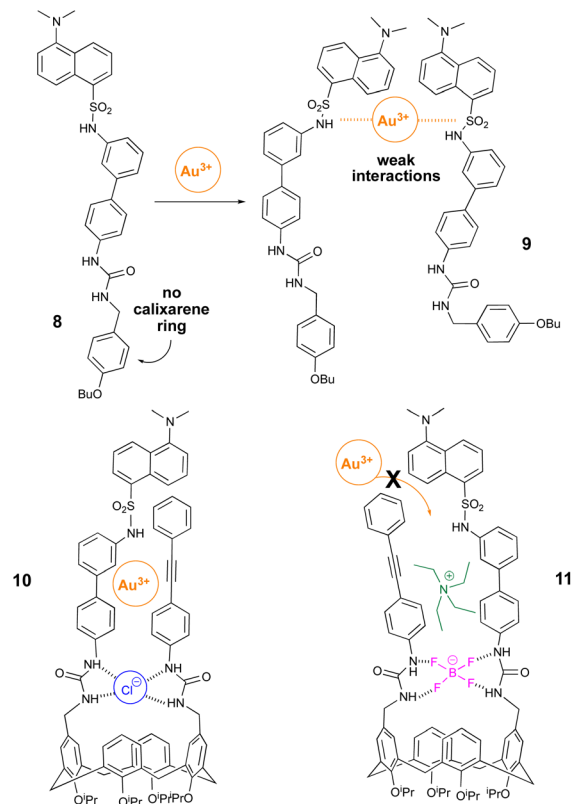


Fig. 6 Examples of motifs generated to probe the role of the calix[4]arene (**8**) and the two urea's (**10** and **11**).

arene cavities,²⁴ we considered it unlikely the bulky tetraethylammonium would occupy the calix[4]arene cavity of **7**; a fact further compounded by its sterically encumbered upper-rim substituents. Stirring **7** with 5 equivalents of tetraethylammonium tetrafluoroborate afforded, presumably, the urea N-H bonded BF_4^- with its Et_4N^+ counterion (**11**, Fig. 6). Although attempts to substantiate the formation of **11** *via* X-ray diffraction were unsuccessful when the gold salt was added sequentially (>30 equivalents), a relatively small drop in the λ_{max} was observed (ESI⁺). This indicates the metal is largely blocked from interacting with the triple bond and dansyl group of **11**. Accordingly, we surmise from **8** and **11** that two urea groups are key parts of the architecture of **7**. These preliminary results establish the importance of the alkyne, with this in mind we wanted to probe this further exploring the stereoelectronic effect associated with *reducing* or *increasing* alkyne electron-density and, consequently, its ability to interact with the Au^{3+} . Our assumption focused on the premise that *reducing* alkyne electron density (**12**) would *reduce* its interaction with the metal and, as such, afford *reduced fluorescent quenching*. Of course, on this basis, incorporating *electron-rich* **13** the opposite would be expected; that is, *increased fluorescent quenching*. The synthesis of *para*-nitrophenyl-derived **12** and *para*-methoxyphenyl **13** followed the same general procedure outlined in Scheme 2 in 56% and 68% yields respectively.



Subjecting separate solutions of metal-free **12** and **13** to fluorescent analysis ($2.5 \mu\text{M}$, $\lambda_{\text{ex}} 288 \text{ nm}$, $\lambda_{\text{max}} 520.7 \text{ nm}$) established, as anticipated, the λ_{em} of **12** to have *reduced* fluorescence intensity (Fig. 8) whilst electron-rich **13** exhibited *increased* intensity (Fig. 9). Sequential addition of 1, 2, 3, 4 and 5 equivalents of aqueous sodium tetrachloroaurate afforded, under otherwise identical conditions, a greater *reduction* in the fluorescence intensity of *electron-rich* **13**. Accordingly, we

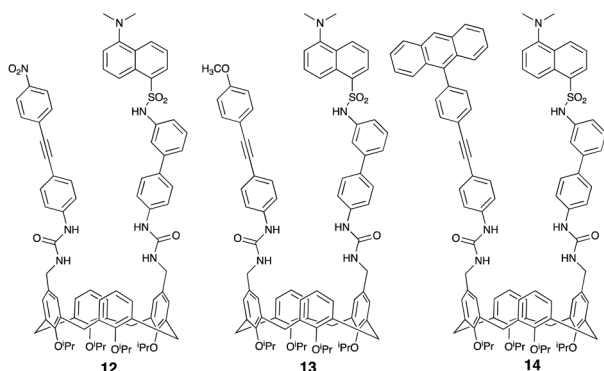


Fig. 7 Dansyl- and urea-derived *cone*-calix[4]arenes synthesised and appended with an electron-poor (**12**), electron-rich (**13**) or an anthracene-derived alkyne (**14**).

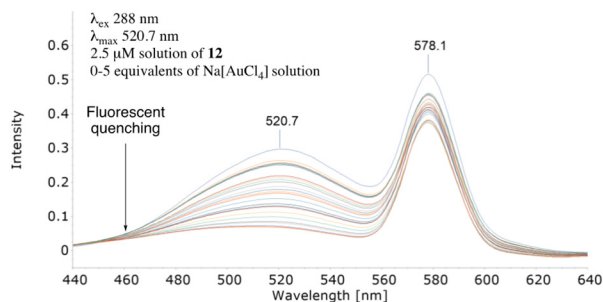


Fig. 8 Effect on the metal-mediated fluorescent quenching properties of **12** when a conjugated electron-withdrawing *para*-nitrophenyl group is appended.

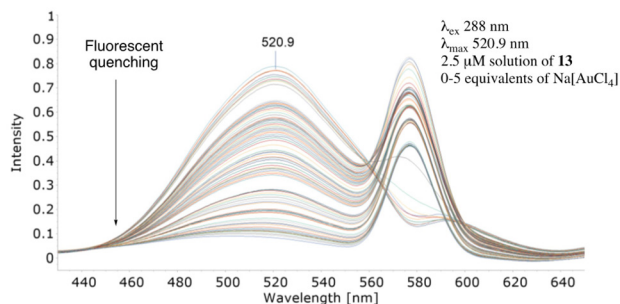


Fig. 9 Effect on the metal-mediated fluorescent quenching properties of **13** when a conjugated electron-donating *para*-methoxyphenyl group is appended.

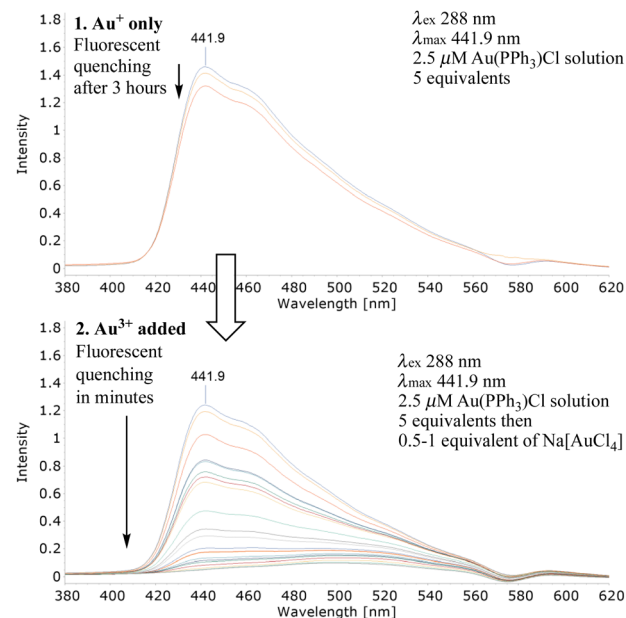


Fig. 10 (Top) limited effect of excess chloro(triphenylphosphine)gold(I) on the fluorescence properties of 9-(4-ethynylphenyl)anthracene-derived **14** after 3 hours. (Lower) efficient and very quick fluorescent quenching of **14** when 0.5-1 equivalents of sodium tetrachloroaurate are added to a mixture of **14** and excess chloro(triphenylphosphine)gold(I).

surmise increasing electron-density on the alkyne resulted, as anticipated, in augmented Au^{3+} sensitivity.

Preliminary work established **7** did not detect Au^+ (Fig. 2). However, it is known that alkyne activation using Au^+ or Au^{3+} co-ordination ($\pi \rightarrow \sigma^*$) is facilitated if the electrophilic, cationic intermediate is stabilised *via* extensive delocalisation.²⁵ Having synthesised extensively conjugated 9-(4-ethynylphenyl)anthracene-derived **14** (Fig. 7) from **4** in 2 steps (58% overall yield) we wanted to probe its ability to display enhanced sensitivity to Au^{3+} in the presence of Au^+ . Mixing solutions of **14** ($2.5 \mu\text{M}$) and 5 equivalents of chloro(triphenylphosphine)gold(I), the emission spectrum was acquired over 3 hours; little change was observed *viz.* Fig. 10. To the *same* solution containing the Au^+ was added 1 equivalent of sodium tetrachloroaurate (*cf.* 7). An almost immediate reduction in the λ_{em} and rapid, fluorescent quenching was observed (Fig. 10).

Mechanism of fluorescent quenching

Our preliminary results using **7-8** and **12-14** substantiate the importance of the 'triad' comprising the calix[4]arene, the alkyne and the fluorophore (*cf.* 3). Focusing on the two key properties outlined below, we wanted to rationalize the observed efficient fluorescent quenching of the fluorophore based on:

- The ability of a calix[4]arene bearing an *N*-sulphonylcarboxamide *e.g.*, **7**, **12-14** to interact with Au^{3+} .
- The propensity of the alkyne to act as an aurophile inducing electron-charge transfer from the alkyne to the metal



which stimulates alkyne polarization and subsequent reaction with a nucleophile.

Interestingly, although alkyne coordinated *linear* Au⁺ complexes have been known since the 1970's, alkyne π -systems comprising *square-planar* Au³⁺ have, only recently, been verified.²⁶ Nevertheless, invoking the findings of Belanzoni *et al.*²⁷ we propose Au³⁺-alkyne coordination proceeds *via* a classically described dominant σ -component coupled with smaller, but significant, π -back-donation (Dewar–Chatt–Duncanson model). Resulting in significant polarization of the π -alkyne electrons the Au³⁺ induces strong triple bond polarization and increased alkyne reactivity; this facilitates nucleophilic attack by, for example, a protic solvent (15 \rightarrow 16, Scheme 3).²⁸ Supporting the proposed formation of (*E*)-vinylgold(III) species 16 we note, with interest, the identification *via* X-ray diffraction of 19, see crystal structure 20 (Scheme 3). Oxyauration of *N*-(propargyl)benzamide 18 using Au³⁺ activates the C–C triple bond (Scheme 3) which initiates a 5-*exo-trig* cyclisation and the formation of 19 with the Au³⁺ in a square-planar complex.²⁹ As noted earlier, support for ketone formation (17, Scheme 3) was from a 1991 report by Ultimoto and Fukuda who described how a wide variety of structure and function diverse alkynes

readily react in the presence of 2 mol% sodium tetrachloroaurate and aqueous methanol affording ketones in excellent yield. Importantly, and with reference to our observed lack of FQ with chloro(triphenylphosphine)gold(I), when Ultimoto incorporated gold(I) potassium dicyanoaurate [KAu(CN)₂] the reaction completely failed to mediate any alkyne hydration.¹⁰ As the dansyl group on 7 is slightly acidic (pK_a 3.8 and 9.9) partial sulphonamide dissociation in methanol would afford a dansyl anion which we propose has the potential to undergo complex formation with the closely located, upper-rim bound Au³⁺. Thus, overall and consistent with known mechanisms of transition-metal ion-induced fluorescent quenching the ligand-immobilized excited dansyl fluorophore is quenched *via* a photoinduced electron transfer (PET) mechanism from the excited dansyl group which is in close proximity to the π -[LAu]-bound complex (15 \rightarrow 16, Scheme 3).

Conclusions

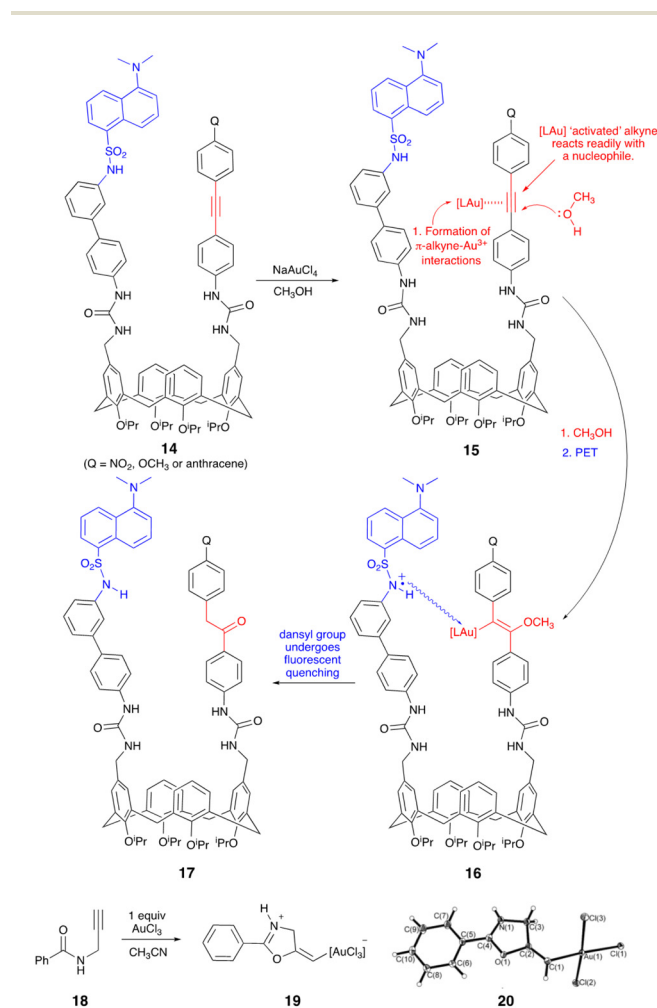
We report preliminary, first examples of simple, effectively designed macrocycles which when appended with a fluorophore and an alkyne react efficiently *via* a chemoselective aurophilic process. Our study demonstrates the synergistic behaviour of an alkyne which, when dovetailed with a dansyl fluorophore and a calix[4]arene, co-operatively and chemoselectively differentiates Au³⁺ in the presence of Au⁺. Of note, the presence of sodium, potassium, cesium, magnesium or calcium salts as well as potentially alkyne-activating Group 12 metals such as cadmium, mercury and platinum salts do not interfere with the aurophilic properties of 7. Given the generality of our synthetic approach, the utility/high demand for the compounds described it is our opinion this work has significant potential for the development of new sensing platforms with applications in the environment and or biology. Furthermore, due to the overarching simplicity and straightforward synthetic methodology outlined in this proof-of-concept study it is our hope this work will pave the way for future advances in supramolecular-based modalities with real-time target applications. In summary, we consider the work outlined herein to be of general interest to, not only, macrocyclic, organic and synthetic chemistry-based academics but will also garner the attention of industrial and analytical chemists working at the frontiers of biological, pharmaceutical, agrochemical, analytical and materials chemistry.

Author contributions

SPB undertook project conceptualization, supervision and project administration. SVS and WHG undertook the methodology and reaction investigation studies.

Conflicts of interest

There are no conflicts to declare.



Scheme 3



Acknowledgements

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