# Fluorescent PET (Photoinduced Electron Transfer) sensors as potent analytical tools

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Fluorescent sensors are an important part of the analytical scientist's toolbox. The use of fluorescent PET (Photoinduced Electron Transfer) sensors has seen particular growth in recent times. This Critical Review discusses recent growth areas in fluorescent PET sensors by emphasizing the modular features of the 'fluorophore-spacer-receptor' design. The occurrence of the dipicolylamine receptor in PET sensor designs is critically examined as a case in point.

## Introduction

Analytical science continues to be enriched by the principles of supramolecular chemistry.1,2 Hardly a week goes by without a new fluorescent PET (Photoinduced Electron Transfer) sensor being announced. What are these sensors and how do they operate? How did they evolve? What are the reasons for their widespread development? This Critical Review briefly addresses these questions before tracing the recent lineage of a single subfield of fluorescent PET sensors and presenting some highlights in the field from the past year.

## Molecular engineering design

A single simple picture (Fig. 1) encapsulates the design of fluorescent PET sensors. The 'fluorophore-spacer-receptor' format is a rational combination of three components. The rationale is contained in Fig. 2a and 2b. In its 'off' state, excitation of the fluorophore component of the sensor produces an electron transfer from the receptor to the fluorophore as one possibility. In other words, the excited state energy of the fluorophore needs to be sufficient to provide both the reduction potential of the fluorophore and the oxidation potential of the receptor. This is a thermodynamic condition first derived by Weller.<sup>3</sup> In its 'on' state, excitation of the fluorophore results in fluorescence only because the PET process is arrested by the arrival of the analyte at the receptor site. The arrest can be easily comprehended by considering H+ as the analyte. H+ electrostatically attracts the electron which increases the oxidation potential of the



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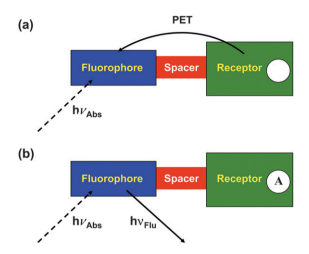
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Fig. 1 The 'fluorophore-spacer-receptor' format of fluorescent PET sensors.



**Fig. 2** (a) An electron transfer from the analyte-free receptor to the photo-excited fluorophore creates the 'off' state of the sensor. (b) The electron transfer from the analyte-bound receptor is blocked resulting in the 'on' state of the sensor.

analyte-bound receptor to the point that the thermodynamics for PET are no longer favourable.<sup>4,5</sup>

These ideas can also be expressed with the aid of molecular orbital energy diagrams (Fig. 3a and 3b). We note the bracketing of the receptor HOMO by the frontier orbitals of the fluorophore in the 'off' state of the sensor and the stabilization of the analyte-bound receptor's HOMO to lie below the fluorophore's HOMO in the 'on' state. Fig. 3a and 3b allow us to deduce an even simpler criterion for PET sensor design: PET occurs if the oxidation potential of the receptor is smaller in magnitude than that of the fluorophore. The opposite applies in the 'on' state of the sensor. This rule of thumb is very useful practically, even though several approximations are involved. More accurate treatment of PET processes are available for the interested reader. MO energies and related redox potentials are increasingly used by PET sensor designers. 9-12

The availability of a quantitative design criterion is common in engineering but rare in chemistry. The case of fluorescent PET sensors is a rare example of molecular engineering design. Just like houses and cars, molecular PET sensors can now be designed and built for a variety of individual purposes.

Each of the three components within the 'fluorophore-spacer-receptor' format deserves the designer's attention. The analyte to be sensed determines the choice of receptor. The reciprocal of the binding constant for the receptor-analyte interaction determines the median analyte concentration to be sensed. Consideration needs to be given at this stage to the selectivity of the receptor towards the analyte and against anticipated levels of potential interferents.

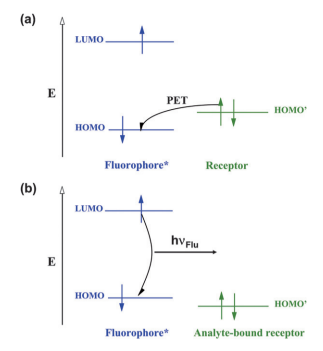


Fig. 3 Molecular orbital energy diagrams which show the relative energetic dispositions of the frontier orbitals of the fluorophore and the receptor in (a) the analyte-free situation and (b) the analyte-bound situation.

Desired colours for excitation and emission help in the choice of fluorophore. For instance, intracellular studies using glass microscopy optics will preclude the use of excitation wavelengths below 340 nm. Tissue experiments will prefer these wavelengths to be in the red region.

The ease of sensor synthesis dictates the choice of spacer, but the more fundamental determinant is that the spacer must be short enough to permit reasonably fast PET rates in the 'off' state of the sensor. <sup>13–15</sup> Even virtual spacers can be used provided that other means, such as sterically-induced orthogonalization, <sup>16</sup> maintains the separation between the fluorophore and the receptor.

Let us consider an example of simple pH sensing which will serve as a foundation for a case study (see below). Fig. 4 shows two of the main options available to the excited PET sensor 1.4 In order to apply the approximate Weller equation, we note that the excited state energy of the anthracene fluorophore is 3.0 eV.<sup>17</sup> Its reduction potential is -2.0 V (vs. sce). The oxidation potential of the receptor can be estimated from that of triethylamine (+1.0 V).17 As the transiting electron falls through these potentials, the corresponding energies are 2.0 eV and 1.0 eV respectively (Fig. 5a). So the approximate  $\Delta G$  for PET is 0.0 eV.<sup>17</sup> This gives a sufficiently fast PET rate to overcome fluorescence  $(k_{\rm PET} \gg k_{\rm Flu})$ . The same result can be obtained by the rule of thumb when we note that the oxidation potential of the receptor is +1.0 V as above and that the oxidation potential of the anthracene fluorophore is +1.0 V. So  $\Delta G_{PET}$  is 0.0 eV again. When we consider the H+-bound amine receptor of 1, its oxidation potential rises to an immeasurably high value.  $\Delta G_{\rm PET}$ becomes a large positive number and fluorescence dominates.

Fig. 4 De-excitation pathways open to the photo-excited fluorescent PET sensor 1.

Another more recent example shows how these thermodynamic arguments help even when fluorescent PET sensors are constructed without a covalent linkage between the fluorophore and the receptor. Self-assembly of the trianionic fluorophore 2 within the cavity of the tetracationic receptor 3 produces the 'off' state of the sensor system. 18 The reduction potential of the receptor can be estimated from that of dimethyl viologen (-0.3 V). The oxidation potential of 2 is 0.8 V and its excited state energy is 3.0 eV (Fig. 5b). 19 Fast PET is made possible by significantly negative  $\Delta G_{\text{PET}}$  value (-1.9 eV). Displacement of 2 from its complex with 3 by guanosine triphosphate (GTP) occurs due to increased synergistic effects, such as electrostatics and  $\pi$ -stacking inside the cavity. Since the fluorophore 2 is now distanced from the receptor 3 (besides somewhat less favourable thermodynamics for PET) fluorescence is switched 'on'. Sensing of GTP is thus enabled.

The validity of such molecular engineering makes PET sensors<sup>20</sup> an important segment of research in molecular devices.<sup>21</sup>

### Early history

The first case fitting the above description of a fluorescent PET sensor was Wang and Morawetz's dibenzylamine compound (4; as seen later in Fig. 7b).<sup>22</sup> It contained a small fluorophore requiring excitation in the deep ultraviolet, spaced with a methylene group from an aliphatic amine receptor. Naturally, H<sup>+</sup> was the chosen target, though it was also engaged with Zn<sup>2+</sup> and also reacted with acetic anhydride. Several cases<sup>23–28</sup> followed with

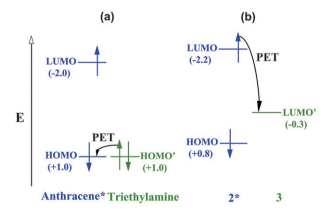


Fig. 5 The relative energetic dispositions of the frontier orbitals of the fluorophore and the receptor in the analyte-free situation for PET sensor systems (a) 1 and (b)  $2 \cdot 3$ . Redox potentials (V vs. sce) are given in parentheses as a measure of molecular orbital energies. The energies are not to scale.

spacers ranging from trimethylene to none at all. The latter situation occurred due to the sterically-enforced twisting of an aniline receptor from an anthracene fluorophore. The generality of the sensing principle was established with a set of related cases<sup>28–36</sup> carrying different fluorophores (or phosphors), spacers and receptors. Reviews of this phase are available.<sup>4,5,37,38</sup>

# Current uptake

As may be expected, a principle that is general, flexible and extensible tends to be put to use by people seeking solutions to various analytical problems. A sensing principle would naturally be applied to target various analytes in various situations. A commercially successful example which measures blood components like H<sup>+</sup>, Na<sup>+</sup>, K<sup>+</sup> and Ca<sup>2+</sup> deserves a special mention.<sup>39</sup> There is a growing body of work where PET sensors are operating within living cells.<sup>40,41</sup> The current situation is perhaps best shown graphically. Fig. 6a and 6b show the sources for fluorescent PET sensors and switches around the world as deduced from the literature. Some of these laboratories may have produced a single publication in this field or several dozen. It is clear that research in fluorescent PET sensors is now a delocalized activity.

#### A case study: dipicolylamine-based sensors

It is educational to track how a single avenue of fluorescent PET sensors has evolved. It illustrates how different people considering different problems can exploit a single structural motif. Consider di(2-picolyl)amine {IUPAC name: 2-pyridinemethanamine, *N*-(2-pyridinylmethyl)-} which is a popular receptor<sup>42</sup> for d-block cations among coordination chemists. All of the structures discussed are contained within Fig. 7a and 7b.

Though 5 (Fig. 7a) was a previously known compound,  $^{43}$  S. A. de Silva *et al.*  $^{44}$  were the first to recognize its 'fluorophore–spacer–receptor' format, its PET potential and its di(2-picolyl)-amine receptor  $^{42}$  for  $Zn^{2+}$ . Indeed a strong  $Zn^{2+}$ -induced switching 'on' of fluorescence (fluorescence enhancement factor, FE = 77) is seen in acetonitrile. No wavelength shift of the emission band is seen as befits a PET system. The replacement of the terminal methyl groups of 1 by 2-pyridyl units will make the  $\Delta G_{\rm PET}$  only slightly more positive. The thermodynamic conditions for cation sensing discussed in a previous paragraph are maintained. More recent crystallographic and computational studies have also backed up the  $Zn^{2+}$ -binding of  $5.^{45}$ 

The modularity of the PET sensing system **5** can now be exploited in many ways and we track structural mutations where the dipicolylamine unit is conserved. The number of laboratories that have followed this path is remarkable, spurred on by the

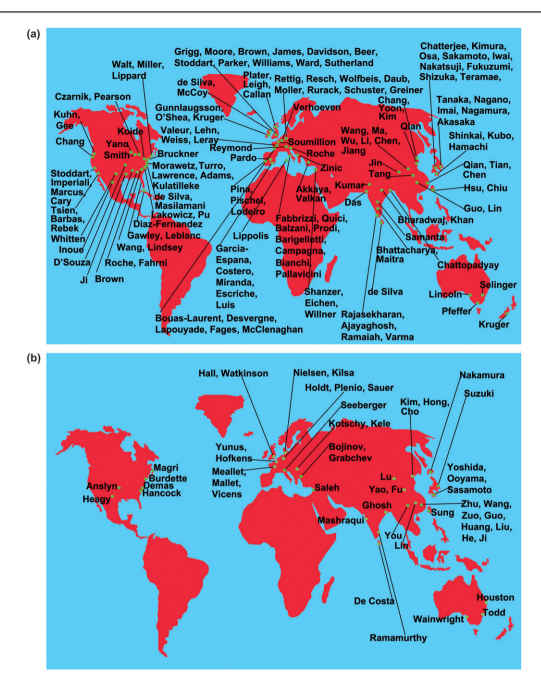


Fig. 6 (a,b) Sources of fluorescent PET sensors. Only the names of corresponding authors from the literature are given. The corresponding authors will be happy to receive evidence of errors and omissions so that future versions of the maps can be improved.

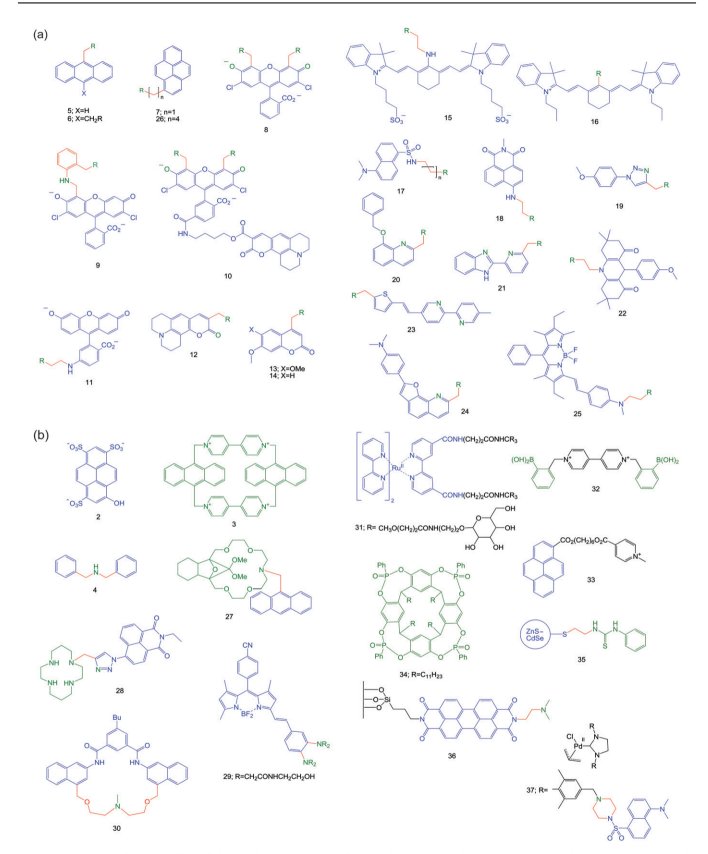
need for monitoring the neurophysiology of  $Zn^{2+}$  and its role in degenerative disease.  $^{40,41,46}$ 

The extension of PET systems by adding an extra 'spacer–receptor' component can lead to improvements in FE values.<sup>4,47</sup> This is seen when we go from 5 to 6,<sup>48,49</sup> with the requirement that each receptor captures  $Zn^{2+}$ . However, ethanol:water (1:1) was needed as the solvent.

The hydrophobicity of the anthracene fluorophore would hamper the use of 5 for monitoring Zn<sup>2+</sup> in the cytosol. Hydrophobic PET sensors can localize in intracellular membranes and lose their ion-sensing ability.<sup>12</sup> Therefore, replacement of anthracene by hydrophilic heterocyclic fluorophores is logical,

and is discussed in the very next paragraph. On the other hand, the hydrophobicity of **5** or its cousin  $7^{50}$  can be exploited for  $Zn^{2+}$  measurement in nanospaces adjoining membranes. The fluorophore is expected to embed in detergent micelles with the more hydrophilic receptor being accessible to water neighbouring the membrane<sup>51,52</sup> and any  $Zn^{2+}$  therein. In the event, a strong  $Zn^{2+}$ -induced FE value of 7 is found for 7 in neutral Tween 20 micelles.<sup>50</sup>

The mutation of **5** is easiest to see in **8**,<sup>53</sup> where the tricyclic anthracene has become the tricyclic fluorescein. The chloro substituents serve to reduce **8**'s response to H<sup>+</sup> in physiological conditions.<sup>54</sup> Sensor **8** gives a Zn<sup>2+</sup>-induced FE value of 2 in



**Fig. 7** (a) Structural formulae of the dipicolylamine-based sensors discussed in the case study. In all these cases, the R group represents the di(2-pyridylmethyl)amino moiety. (b) Structural formulae for the compounds highlighted from the past year. Following on from Fig. 2a and 2b, the following colour scheme is employed in both Fig. 7a and 7b: fluorophores in blue, spacers in red and receptors in green. When atoms in the fluorophore can also ligate to the analyte, these are shown in green.

physiological media and permits fluorescence microscopy in living cells. However, the residual fluorescence response to H<sup>+</sup> is still quite large, since scavenging of Zn<sup>2+</sup> does not noticeably reduce the fluorescence. Nevertheless, **8** was the vanguard of a strong programme<sup>55</sup> which included cases like **9** and **10**. Sensor **9**<sup>56</sup> locates the dipicolylamine unit somewhat remote from the fluorophore. However, the aniline NH group is just one methylene unit away from the fluorophore so that PET would occur at a reasonably rapid rate.<sup>57</sup> The NH unit joins in Zn<sup>2+</sup>-binding along with the dipicolylamine receptor. Interestingly, another strong program begun in 2000<sup>58</sup> has a similar juxtaposition of an aniline NH and a dipicolylamine, as well as a fluorescein fluorophore, *e.g.* sensor **11**.<sup>59</sup>

Sensor 10<sup>60</sup> contains 8 as the Zn<sup>2+</sup>-responsive component with green emission of dichlorofluorescein as the output. It also carries an aminocoumarin fluorophore whose blue emission is Zn<sup>2+</sup>-independent, which is released by hydrolysis of the ester bond of 10 by intracellular esterases. Virtually the same aminocoumarin fluorophore is present in 1261,62 (though closer to the receptor) and so the Zn<sup>2+</sup>-independent emission intensity might be expected. However, the positioning of the dipicolylamine near the coumarin carbonyl oxygen allows the latter to participate in Zn<sup>2+</sup>-binding. Since this push-pull fluorophore has an internal charge transfer (ICT) excited state whose  $\delta^-$  pole lies at the carbonyl oxygen, Zn<sup>2+</sup> causes an emission red-shift due to the electrostatic attraction. Alkoxycoumarin fluorophores when coupled to amine receptors have suitable PET thermodynamics. 63,64 Cases like 1361,62 and 1465 show Zn2+-induced FE values of 22 (in methanol) and 8 (in acetonitrile) respectively.

While blue (*e.g.* coumarin) and green (*e.g.* fluorescein) emissions remain the workhorses of fluorescent sensor research, redemitting fluorophores are sought after for intracellular and tissue studies due to easy transmission of light. Zn<sup>2+</sup> sensors 15<sup>66</sup> and 16<sup>67</sup> address this need. PET has been achieved with cyanine fluorophores when coupled with electron-rich anilinic receptors, <sup>68</sup> but is harder to produce with the dipicolylamine receptor. Sensor 16 succeeds at this with a Zn<sup>2+</sup>-induced FE value of 7 (in water) whereas 15 only shows wavelength shifts in fluorescence excitation spectra typical of ICT excited states, *e.g.* Tsien's classical Ca<sup>2+</sup> sensors.<sup>69</sup>

The blue-green region is also represented by several more Zn<sup>2+</sup> sensors 17–21.<sup>70–75</sup> All of them show significant Zn<sup>2+</sup>-induced FE values, though some of them involve the fluorophore's participation in the Zn<sup>2+</sup>-coordination sphere. System 22<sup>76</sup> is clearly of the 'fluorophore–spacer–dipicolylamine' format but PET appears to be thermodynamically unfavourable. So, 22 is highly emissive to begin with and only the quenching metal ion Cu<sup>2+</sup> signals its presence. Zn<sup>2+</sup> has no effect.

Another blue-green-emitting  $Zn^{2+}$  sensor  $23^{77}$  is distinguished by operating over a large dynamic range. Besides the dipicolylamine receptor, 23 also contains a 2,2'-bipyridine as a low-affinity binding site which is engaged at high  $Zn^{2+}$  concentrations. This incurs positive charging and planarization of the bipyridine rings and so the ICT character of the excited push-pull fluorophore is enhanced with a significant red-shift. The switching 'on' of the blue emission and that of the green emission occur at two different  $Zn^{2+}$  concentration ranges.

The conjugation of an electron-rich aminophenyl substituent with a furoquinoline unit increases its reduction potential to

weaken PET activity of 24.<sup>78</sup> However, this produces a push-pull fluorophore with considerable ICT character in its excited state. The quinoline nitrogen atom of the fluorophore clearly participates in the Zn<sup>2+</sup>-coordination sphere, so that the ICT character of the excited state increases, resulting in a Zn<sup>2+</sup>-induced red-shift from green to orange. This allows ratiometric measurement of Zn<sup>2+</sup> in living cells.

The story doesn't end here. Some of these Zn<sup>2+</sup> sensors have led to another valuable line of research. It started with the filling of the unsaturated coordination sphere of dipicolylamine-Zn2+ (mentioned above) with anionic ligands such as phosphates. When di-receptor cases such as 6 are studied in neat water, binding of two Zn<sup>2+</sup> ions cannot be achieved under experimental conditions unless the mutual repulsion between the two metal centres is reduced by inserting a bridging anion. Therefore, PET suppression and fluorescence switching 'on' requires the presence of Zn2+ as well as the phosphate anion. Phosphorylation of tyrosines in peptides can be neatly signalled in this way. Of course, dephosphorylation can be followed fluorimetrically too. 79,80 Sensor 6 with Zn2+ also switches 'on' when uridine 5'-diphosphate is made available from uridine 5'-diphosphate glycoside during glycosyl transfer to sugar derivatives catalyzed by glycosyltransferases.81 The latter enzymes are measurable in a label-free manner. Sensor 6 with Zn2+ also lights up when phosphatidylserine is brought to the outer face of cell membranes when the cell is ready to die. Thus, apoptosis can be detected in a simple way.82

Phosphates can also be detected (though in acetonitrile solution) by the quenching of the emission of the  $Zn^{2+}$  complex of **25** by a factor of  $70.^{83}$  Sensor **25** itself is poorly emissive and  $Zn^{2+}$  causes an FE value of 5. A PET mechanism is probable. The coordination sphere of the single dipicolylamine- $Zn^{2+}$  is made up by the aniline nitrogen nearby and, importantly, also by phosphate which reduces the positive charge density so that the  $Zn^{2+}$ -induced PET suppression is weakened. The ICT nature of the push-pull fluorophore is clearly signalled by the  $Zn^{2+}$ -induced blue-shift.

Compound 8<sup>53</sup> is re-incarnated in the form of its di-Zn<sup>2+</sup> complex for pyrophosphate sensing.<sup>84</sup> An FE value of 3 is achieved. It appears that, as seen perhaps more strongly for 6,<sup>79</sup> the full binding of both dipicolylamine sites and the full PET suppression is only achieved when the pyrophosphate bridges the two Zn<sup>2+</sup> centres. In this case, pyrophosphate is of just the right length. Pyrophosphate bridging two Zn<sup>2+</sup>-dipicolylamine units is also found in the ground state dimer of 26<sup>85</sup> (indicated by redshifts in the absorption spectrum) which leads to a corresponding red-shifted emission compared to the pyrene monomer.

It is clear that the first PET sensing experiment on  $5^{44}$  keeps on putting out new shoots – a testament to the flexibilities arising from the modular nature of PET sensors.

#### Some highlights from the past year

Besides the sustained success of Zn<sup>2+</sup> sensing with PET sensors carrying bis(picolyl)amine receptors, recent cases also include macrocycles. An oxa-bridged version of older azacrown etherbased PET sensors,<sup>29</sup> **27** (Fig. 7b),<sup>86</sup> contains an amine PET donor and an anthracene fluorophore. It produces a Zn<sup>2+</sup>-induced FE of 100 but does not respond to alkali cations.

Interestingly,  $Na^+$  in sufficient concentration displaces  $Zn^{2+}$  and decreases the FE value of **27**, as the smaller ion coordinates to the oxygen atoms rather than the nitrogen.

Cases based on tetraazamacrocycle receptors are also useful for  $Zn^{2+}$  sensing.<sup>87</sup> A new example is **28**,<sup>88</sup> where the triazole group also plays a receptor role to cause selective binding of  $Zn^{2+}$  even over  $Cd^{2+}$ , with a  $Zn^{2+}$ -induced FE value of 6. Interestingly, it detects the intracellular flux of  $Zn^{2+}$  during cell apoptosis.

In spite of the successes concerning selective Zn<sup>2+</sup> sensing, Cd<sup>2+</sup> was an interferent in many of these cases. The rarity of intracellular Cd<sup>2+</sup> has been the saviour in this regard. Selective Cd<sup>2+</sup> sensing, which would be useful in studies of Cd<sup>2+</sup> toxicity, has been a harder nut to crack.<sup>89-91</sup> PET sensor **29** with a polyamide receptor provides a neat solution even within HeLa cells.<sup>92</sup> A Cd<sup>2+</sup>-induced FE value of 100 is attained. Steric hindrance between the receptor arms can produce a virtual spacer so that the usual PET behaviour can occur.

Organic chemical reactions rather than ion-coordination can also lead to PET-based fluorescence switching 'on', as seen in 30.93 When solubilized in aqueous Tween 20 micellar solution, its amine group serves as the PET donor so that the fluorophore's emission is 'off'. The dangerous alkylating agent chloromethyl methyl ether forms a quaternary ammonium product which prevents PET and fluorescence is switched 'on'. Related cases<sup>94</sup> are known.

As we saw with the 2·3 system, <sup>18</sup> covalent linking of a fluorophore and a PET-active receptor is not necessary for a working PET sensing system. There is no PET within 31, a tris(2,2'-bipyridyl)Ru(II) lumophore which is connected to a mannose-capped dendrimer. <sup>95</sup> Lectins, those exquisite sugarbinding proteins, rely on polyvalency. Therefore, lectins such as Concanavalin A associate strongly with 31. However, this does not produce a luminescence switch unless an extra component is provided. The extra component 32 contains arylboronic acid groups for sugar binding and a 4,4'-bipyridinium unit as a PET acceptor. In the absence of lectin, 31 and 32 associate so that PET takes place and the luminescence is 'off'. The addition of Concanavalin A displaces 32 from the mannose caps and a resultant increase in luminescence occurs.

Though perhaps not as grand as lectin sensing, synthetic macrocycles can also be sensed *via* PET schemes. A pyrene fluorophore and a pyridinium PET acceptor can be discerned within 33.96 The phosphonate-bridged resorcinarene 34 can engulf the pyridinium unit just like simpler resorcinarenes do.97 Upon complexation, PET becomes energetically unfavourable, causing an increase in fluorescence.

As seen in the previous pages, PET sensors usually consist of molecular lumophores. An important extension to this idea has been reported<sup>98</sup> where the lumophore component represents a quantum dot, the darling of nanotechnology. This particular quantum dot is a ZnS–CdSe core–shell structure. Sensor **35** possesses a thiourea receptor which binds to CH<sub>3</sub>CO<sub>2</sub><sup>-</sup> with two hydrogen bonds and this increases the reduction potential of the receptor, thus enhancing PET to the lumophore.

We finish off with two cases where 'fluorophore-spacer-receptor' systems are aimed at monitoring intermediates in catalytic cycles in solution. For instance, a transient Lewis acid centre could bind a receptor to switch 'on' the fluorescence of a PET sensor. This would be a valuable analytical tool. Example

36°9 attacks this problem at the single molecule level since the required sensitivity of similar sensors based on perylenediimide fluorophores has been demonstrated. Though this ambitious goal is yet to be achieved, the H<sup>+</sup>-induced switching 'on' of fluorescence is demonstrated by negating the PET donor amine by protonation. The single molecule studies show this pH dependence, though the presence of relatively long-lived dark states at low pH leading to 'blinking' is a complication.

The dimethylaminonaphthalenesulfonylamide fluorophore undergoes PET from neighbouring amines as seen in the case of 17.101 A similar 'fluorophore-spacer-receptor' motif can be found within 37. This motif is tagged to an N-heterocyclic carbene Pd(II) complex with the intent of monitoring its catalytic activity in a Suzuki coupling reaction between an aryl bromide and a boronic acid. As the reaction progresses, the halide ion product quenches 37's fluorescence due to the heavy atom effect. Such monitoring of a product formation is not the main point in the present context of monitoring of catalytic intermediates. However, the smaller fluorescence loss observed upon addition of base to prepare the active Pd(0) catalytic species before the addition of the aryl halide is potentially more interesting. If the mopping up of trace Brønsted acids can be ruled out, a 'steppingstone' mechanism could be imagined, where an electron is transferred from Pd(0) to the fluorophore via the amine receptor to quench fluorescence, but not from Pd(II).

#### Conclusion

The preceding pages have summarized a few of the current growth areas in fluorescent PET sensors where problems in analytical science are being attacked. We hope the quantitative design basis of PET sensors where molecules can be viewed as engineering objects will appeal to bright analytical minds so that more of them will join in this venture. With such a combination of forces, even more analytical solutions will emerge from the versatile fluorescent PET system as the days go on.

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