

A two-co-ordinated gold(I) loop $[\text{Au}(\text{dpdo})]\text{ClO}_4$ [dpdo = 1,8-bis(diphenylphosphino)-3,6-dioxaoctane] as a luminescence light switch for substrate binding reactions

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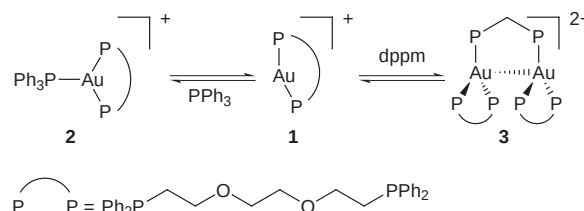
Reaction of $[\text{Au}(\text{tht})\text{Cl}]$ (tht = tetrahydrothiophene) with dpdo [dpdo = 1,8-bis(diphenylphosphino)-3,6-dioxaoctane] afforded $[\text{Au}(\text{dpdo})]^+$ **1** which was isolated as a perchlorate salt; binding of a phosphine moiety to **1** in solution triggered a strong photoluminescence.

A 'molecular light switch' effect, *i.e.* the conversion of a non-emissive metal complex to a strongly emissive one through substrate binding reactions, is of increasing importance in molecular recognition and sensory materials research. The notable examples in this area are $[\text{Ru}(\text{bpy})_2(\text{dppz})]^{2+}$ [bpy = 2,2'-bipyridine, dppz = dipyrro(3,2-*a*:2',3'-*c*)phenazine] for DNA binding and a cyclodextrin derivative with a europium aza crown for incorporation of aromatic hydrocarbons.^{1,2} In these two cases, the external substrates (DNA and aromatic hydrocarbons) bind to the molecular sensors through non-covalent hydrophobic interactions. Our attention has focused on mononuclear two-co-ordinate gold(I) complexes which are known to be non-emissive but display strong photoluminescence upon interaction with added nucleophiles.^{3–6} In this context, the gold(I) loop $[\text{Au}(\text{dpdo})]\text{ClO}_4$ [**1**] ClO_4 [dpdo = 1,8-bis(diphenylphosphino)-3,6-dioxaoctane] (Scheme 1) is of interest. This complex is isostructural to $[\text{Au}(\text{PPh}_3)_2]^+$ but the chelating nature of the dpdo ligand imparts stability and disfavors phosphine ligand dissociation from the Au atom.⁴ We herein report that photoluminescence is 'switched on' through substrate binding reactions to cation **1**. Among the nucleophiles studied, including SCN^- and CN^- , the light switching-on effect is unique to phosphine ligands, suggesting that **1** is a good sensor for phosphine detection even at 10^{-5} mol dm⁻³ concentration.

Reaction of $[\text{Au}(\text{tht})\text{Cl}]$ (tht = tetrahydrothiophene) with a stoichiometric amount of dpdo in dichloromethane at room temperature readily afforded **1**, which was isolated as the perchlorate salt.† The structure of [**1**] ClO_4 has been established by an X-ray crystal analysis.§ As shown in Fig. 1, the complex cation **1** displays a loop-like structure with the dpdo ligand

adopting a *trans*-chelating geometry. The Au atom is two-co-ordinated and adopts a nearly linear configuration [P(1)–Au(1)–P(2) angle of 172.2(1)°]. The Au–O distances of 3.14 and 3.23 Å indicate insignificant interaction between the Au and oxygen atoms.

The ³¹P NMR spectrum of **1** shows a singlet at δ 35.2 (CD₂Cl₂, room temperature). Addition of stoichiometric amounts of PPh₃ or dppm gradually shifts the phosphorus resonances to values closer to that of free PPh₃ or dppm. Cooling the solution to –70 °C leads to three singlets corresponding to **1**, species **2** or **3** and free PPh₃ or dppm (Scheme 1). Attempts to isolate the gold–phosphine species **2** by addition of diethyl



Scheme 1 Equilibria between cation **1** and PPh₃ or dppm in solution

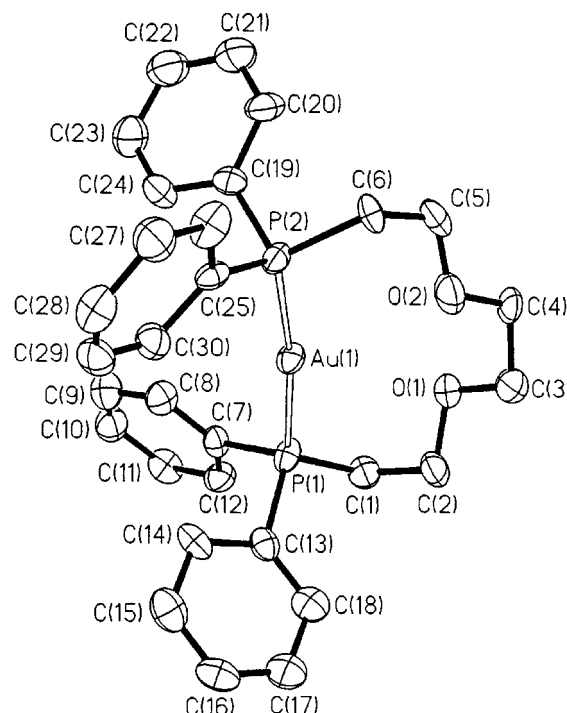


Fig. 1 A perspective drawing of complex cation **1**. Hydrogen atoms are omitted for clarity. Selected bond lengths (Å) and bond angles (°): Au(1)–P(1) 2.328(6), Au(1)–P(2) 2.293(6), Au(1)···O(1) 3.23, Au(1)···O(2) 3.14; P(1)–Au(1)–P(2) 172.2(1)

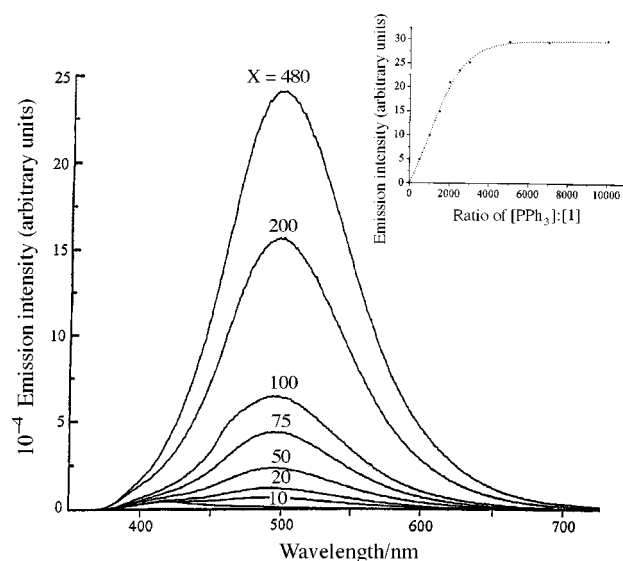
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‡ A mixture of $[\text{Au}(\text{tht})\text{Cl}]$ (0.32 g, 1.0 mmol) and dpdo (0.89 g, 1.0 mmol) in dichloromethane (50 ml) was stirred at room temperature for 2 h. The solvent was removed *in vacuo* leaving a white residue. Metathesis of the crude product with LiClO_4 (0.11 g, 1.0 mmol) in methanol (5 ml) afforded a colorless crystalline solid. Crystals of [**1**] ClO_4 were obtained by diffusion of diethyl ether into an acetonitrile solution (yield 0.51 g, 65%) (Found: C, 45.87; H, 4.15. Calc. for $\text{C}_{30}\text{H}_{32}\text{AuClO}_6\text{P}_2$: C, 46.02; H, 4.12%).

§ Crystal data for [**1**] ClO_4 : $\text{C}_{30}\text{H}_{32}\text{AuClO}_6\text{P}_2$, $M = 782.9$, monoclinic, space group Cc , $a = 15.605(1)$, $b = 13.473(1)$, $c = 15.962(1)$ Å, $\beta = 117.12(1)^\circ$, $U = 2987.0(2)$ Å³, $Z = 4$, $\mu(\text{Mo-K}\alpha) = 5.164$ mm⁻¹, no. of unique reflections = 3567, no. of reflections with $F > 4.0\sigma(F) = 2888$, $R = 0.033$, $R' = 0.038$, $T = 294$ K. CCDC reference number 186/1024.

Table 1 Photophysical data for degassed solutions of [1]ClO₄* in the presence of different substrates measured at room temperature

Substrate (concentration/ mol dm ⁻³)	Solvent	$\lambda_{\text{max}}^{\text{em}}/\text{nm}$ (excited at 328 nm)	$\tau/\mu\text{s}$
—	CH ₃ CN	—	—
—	DMF	468	—
PPh ₃ (10 ⁻⁵ –10 ⁻¹)	CH ₃ CN	510	4.7
dppm (10 ⁻⁴)	CH ₃ CN	610	10.8
Et ₂ S ($\leq 10^{-2}$)	CH ₃ CN	—	—
NaSCN ($\leq 10^{-2}$)	CH ₃ CN	—	—
NaCN ($\leq 10^{-2}$)	DMF	474	—

* Concentration of [1]ClO₄ in acetonitrile $\approx 1 \times 10^{-5}$ mol dm⁻³.**Fig. 2** Room temperature emission spectra of [1]ClO₄ in the presence of PPh₃ in degassed acetonitrile solution. Molar ratio of PPh₃:1 = X:1. Insert: the graph shows a plot of the emission intensity of 1 + PPh₃ vs. the molar concentration of PPh₃:1 (concentration of [1]ClO₄ = 10⁻⁵ mol dm⁻³)

ether to an acetonitrile solution were unsuccessful and the starting complex [1]ClO₄ was recovered. Other nucleophiles such as Et₂S, SCN⁻ and CN⁻ with concentrations 100 times higher than that of PPh₃ do not show any notable effect on the ³¹P NMR spectrum of [1]ClO₄ in acetonitrile.

The photophysical data of [1]ClO₄, and its solutions in the presence of different substrates, are listed in Table 1. The complex is non-emissive in the solid state and in degassed acetonitrile solution. However, addition of PPh₃ ($\geq 10^{-5}$ mol dm⁻³) immediately triggers an intense yellow photoluminescence at 510 nm with a long lifetime ($\tau = 4.7 \mu\text{s}$). We

suggest that the emission comes from a three-co-ordinated gold–phosphine species **2** that is in equilibrium with **1** and PPh₃ (Scheme 1). As expected, the emission intensity increases with further addition of PPh₃ but reaches a plateau value at high PPh₃ concentrations (Fig. 2 and insert). The estimated equilibrium constant of the PPh₃-binding reaction is not large ($32 \pm 3 \text{ dm}^3 \text{ mol}^{-1}$), and hence the dramatic photoluminescence enhancement is likely due to a high emission quantum yield of the three-co-ordinated gold(i) phosphine species. Enhancement of emission has also been observed with dppm (*ca.* 10 molar equivalents). However, the emission maximum is red-shifted to 610 nm ($\tau = 10.8 \mu\text{s}$). Presumably, it is due to the bonding interaction between adjacent AuP₃ units (Scheme 1). Other nucleophiles X (X = Et₂S, SCN⁻ or CN⁻) (≈ 1000 molar equivalents), do not lead to any notable ‘molecular light switch’ effects. This may be attributed to low formation constants and/or low emission quantum yields of the three-co-ordinated AuP₂X species.

In this work, we have shown that photoluminescence is triggered through binding of phosphine substrates to the coordinatively unsaturated Au^I. Compared with [Au(PPh₃)₂]⁺, the structure of **1** is more robust and hence the complex can be exploited as a sensitive spectroscopic probe for compounds with a phosphine moiety.

Acknowledgements

We acknowledge support from The University of Hong Kong, the Hong Kong Research Grants Council and the Croucher Foundation.

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Received 31st March 1998; Communication 8/02464B