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Adjustable coordination of a hybrid

phosphine-phosphine oxide ligand in

Thuy Minh Dau, a Benjamin Darko Asamoah, Andrey Belyaev, a

luminescent Cu, Ag and Au complexes†

A potentially tridentate hemilabile ligand, $PPh_2-C_6H_4-PPh(O)-C_6H_4-PPh_2$ (P^3O), has been used for the construction of a family of bimetallic complexes $[MM'(P^3O)_2]^{2+}$ (M=M'=Cu (1), Ag (2), Au (3); M=Au, M'=Cu (4)) and their mononuclear halide congeners $M(P^3O)$ Hal (M=Cu (5–7), Ag (8–10)). Compounds 1–10 have been characterized in the solid state by single-crystal X-ray diffraction analysis to reveal a variable coordination mode of the phosphine-oxide group of the P^3O ligand depending on the preferable number of coordination vacancies on the metal center. According to the theoretical studies, the interaction of the hard donor P=O moiety with d^{10} ions becomes less effective in the order Cu>Ag>Au. 1–10 exhibit room temperature luminescence in the solid state, and the intensity and energy of emission are mostly determined by the nature of metal atoms. The photophysical characteristics of the monometallic species were compared with those of the related compounds $M(P^3)$ Hal (11–16) with the nonoxidized ligand P^3 . It was found that in the case of the copper complexes 5–7 the P^3O hybrid ligand introduces effective non-radiative pathways of the excited state relaxation leading to poor emission, while for the silver luminophores the P=O group leads mainly to the modulation of luminescence wavelength.

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Introduction

Transition metal complexes, containing multidentate ligands with electronically different functionalities, have been very actively studied due to their versatile coordination chemistry and diverse reactivity. A major stimulus, which determines substantial progress observed in the area over the past few decades, is related to the notable success of such compounds in homogeneous catalysis^{1–3} and activation of small molecules, design of dynamic systems^{5,6} and sensing. The hybrid ligands, which comprise strongly and weakly bound moieties, were defined as hemilabile ones by Jeffrey and Rauchfuss in 1979. The labile functions of such ligands under favourable conditions are capable of easy and reversible coordination—

Among a variety of potentially bifunctional ligands, molecules with phosphorus donors as strongly bound anchors have been most widely used for the preparation of numerous late transition metal compounds. In particular, considerable interest has been focused on the ligands, combining soft (P atom) and hard donors (N or O atoms). ^{2,3,10,11} The corresponding P–N (phosphine–amine/oxazoline/nitrile) and P–O (phosphine–ether/phosphine oxide) complexes of Ni, Pd, Co, Au, and Cu demonstrate impressive catalytic activity in many industrially important reactions, *e.g.* olefin oligo- and polymerization (including ring opening metathesis polymerization), ¹² amination, ^{13,14} hydrogenation, ¹⁵ annulation, ¹⁶ hydroformylation, ¹⁷ allylation, ¹⁸ Suzuki^{3,19} and Heck²⁰ coupling.

Alternatively, hemilabile phosphines have been extensively exploited in the construction of supramolecular systems, which exhibit geometry changes controlled by a weak-link approach (WLA).^{6,21} This general strategy developed by Mirkin resulted in a number of functional compounds, the behaviour of which can be effectively switched *via* binding of an incoming small guest molecule or ion.²²

The mixed phosphine-phosphine oxide (P-PO) ligands have been less studied in comparison with other bifunctional

dissociation to provide a vacancy on the metal center, accessible to accommodate a suitable substrate.

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phosphine congeners and are represented mainly by a family of bis(phosphine) monoxides (BPMO). The relatively limited development of this class of P-PO compounds could be attributed to a more challenging synthesis of unsymmetrical oxide derivatives in comparison with the preparation of the parent phosphines and the low selectivity of direct partial oxidation of the di- or oligophosphines. In this respect, a recently reported tridentate PP(O)P ligand, obtained *via* quite a facile route, a pathway to pincer POP complexes, which remain very poorly explored.

In our previous studies we have been extensively using the di-, tri- and tetraphosphines to support various coordination complexes and cluster compounds of coinage metals, which exhibit tunable photoluminescence characteristics and stimuli-responsive features.²⁵⁻²⁷ However, the employment of mixed P-PO type ligands for the design of luminescent transition metal compounds has been virtually neglected.²⁸ Motivated by this lack of investigations, we attempted to study the coordination chemistry of a hybrid phosphine-phosphine oxide $PPh_2-C_6H_4-PPh(O)-C_6H_4-PPh_2$ (P^3O) with respect to Cu subgroup metals. Utilizing potentially variable denticity of the ligand, which is able to adjust to a preferable coordination number of a given ion, herein we present a series of novel mono- and dinuclear d¹⁰ coinage complexes, their photoemissive properties and a computational analysis of the electronic structures to rationalize the photophysical behavior.

Results and discussion

Synthesis and structural characterization

In the course of our recent studies of d^{10} metal cyanide complexes supported by a triphosphine ligand, $PPh_2-C_6H_4-PPh-C_6H_4-PPh_2$ (P^3), P^3 it was observed that heating a solution of a dinuclear complex $[(P^3)Ag-CN-Ag(P^3)]^+$ on air resulted in some degradation and formation of a novel compound, crystallographically characterized as $[(P^3O)_2Ag_2]^{2^+}$. The product contains the partially oxidized ligand $PPh_2-C_6H_4-PPh(O)-C_6H_4-PPh_2$ (P^3O), which was apparently formed upon interaction with oxygen at elevated temperatures. Due to the low yield and slow rate of this transformation, we prepared independently the diphosphine–phosphine oxide P^3O and investigated its coordination behavior with respect to coinage metal ions.

The ligand, $PPh_2-C_6H_4-PPh(O)-C_6H_4-PPh_2$ (P^3O), was reported earlier as a side product, obtained in 3% yield. ³⁰ Later a similar compound, $P(iPr)_2-C_6H_4-PPh(O)-C_6H_4-P(iPr)_2$, was obtained *via* a more practical route in 73% yield. ²⁴ Following the latter method, P^3O was synthesized from (2-bromophenyl)diphenylphosphine ³¹ in good yield (Scheme 1). The ³¹P NMR spectrum of P^3O displays two resonances (31.9 and -13.6 ppm, $^3J_{PP}$ 12 Hz) with 1:2 integral relative intensities that are consistent with the proposed structure.

Dinuclear complexes $[M_2(P^3O)_2]^{2+}$. The bimetallic compounds $[M_2(P^3O)_2]^{2+}$ (M = Cu (1), Ag (2)) were obtained reacting the P^3O ligand with stoichiometric amounts of the corresponding metal salts (Scheme 2). The complex $[Au_2(P^3O)_2](PF_6)_2$ (3)

Scheme 1 Synthesis of the hybrid ligand P^3O .

was prepared by a slightly modified procedure, which involves treatment of AuCl(tht) (tht = tetrahydrothiophene) with P^3O and subsequent removal of chloride ions with AgPF₆. The heterometallic congener [AuCu(P^3O)₂](PF₆)₂ (4) was obtained from the reaction mixture containing the equimolar amounts of homometallic complexes 1 and 3. The resulting compounds were isolated as colorless (1–3) or pale yellow (4) crystalline materials, stable to air and moisture in the solid state.

The crystal structures of 1-4 were determined by XRD analysis and are shown in Fig. 1 and 2, and selected structural parameters are given in Tables S4 and S5.† In the dication 1 the copper centers adopt a distorted tetrahedral coordination geometry. Both P^3O ligands are bound to the metal ions in a tridentate mode via the terminal P atoms and the O=PPh moiety to form a chair-like P3OCu2OP3 framework with the oxygen atoms in nearly symmetrical µ2-bridging positions between two Cu ions (Cu-O distances are 2.121 and 2.198 Å). In this structural pattern each of the oxygen centers donates both available electron pairs to complete the coordination vacancies at the metals. The copper-oxygen bond lengths are comparable to the values found for the previously reported Cu(I) complexes with coordinated phosphine-oxide function. 13,32 The P=O bond in 1 (1.520 Å) however is slightly longer than those in the abovementioned compounds (<1.503 Å)^{13,32} and in 2-4 (1.491, 1.494 and 1.507/1.506 Å, respectively; see Table 1) presumably due to the bridging coordination mode of oxygen and therefore more effective π -back donation from the metals to $\pi^*(P=O)$ orbitals. The intramolecular Cu···Cu separation of 3.279 Å is noticeably longer than the sum of van der Waals radii (2.80 Å) and the range of common metallophilic Cu-Cu bonds.^{33,34} Together with the saturated coordination environment of Cu(1) ions these structural parameters point to the absence of appreciable cuprophilic interactions.

In 2, the equivalent silver ions adopt a highly distorted tetrahedral arrangement of the ligand sphere with Ag···Ag distance (4.475 Å) significantly exceeding the values typical for argentophilic bonding.³⁵ Each metal center is connected to two terminal P-donors and to the O atom of the phosphine oxide fragment. In contrast to 1, oxygen occupies a visibly unsymmetrical position between the Ag ions, being predominantly coordinated to only one metal (Ag–O distances are 2.457 and 2.885 Å) that is indicative of terminal oxygen coordination in 2 in contrast to the bridging one in 1. The Ag–O bond length of 2.457 Å fits in the range found for other structurally characterized phosphine-oxide silver complexes,³⁶ which are quite rare. However, the tendency of silver(I) to have a tetracoordinate geometry results in a visible π-interaction of the metal ions with adjacent phenyl rings of the O—PPh group

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Scheme 2 Synthesis of complexes 1-10.

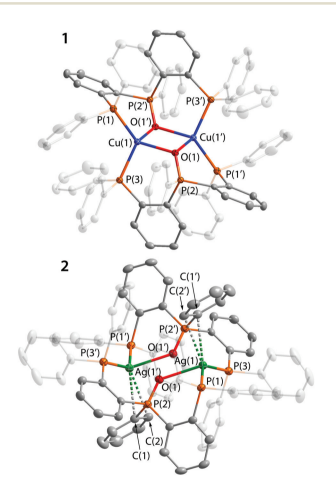


Fig. 1 Molecular views of complexes 1 and 2. Thermal ellipsoids are shown at the 50% probability level. Counterions and H atoms are omitted for clarity. Symmetry transformations used to generate equivalent atoms (') in 1: 1-x, -y, 1-z; in 2: 2-x, 2-y, 2-z.

(see Fig. 1). The distances Ag(1)–C(1) and Ag(1)–C(2) are 2.954 and 2.822 Å, respectively, that point to a moderately weak Ag– η^2 (C=C) bonding³⁷ as the effective interaction of this type requires a characteristic bond length less than 2.9 Å.³⁸

The dimeric complex 3 contains two gold(I) ions, which are held together by the P^3O ligands. The phosphorus atoms

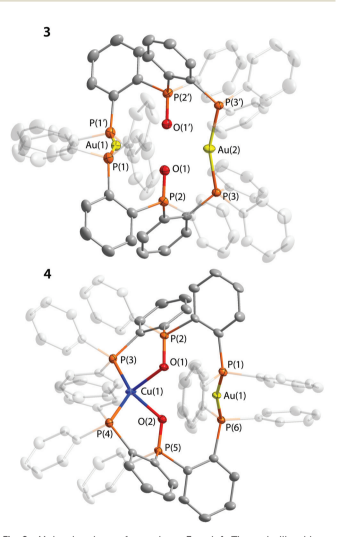


Fig. 2 Molecular views of complexes 3 and 4. Thermal ellipsoids are shown at the 50% probability level. Counterions and H atoms are omitted for clarity. Symmetry transformations used to generate equivalent atoms (') in 3: 1-x, y, 0.5-z.

of the PPh_2 functions form a two-coordinate environment of the metal centers, typical of Au(i) compounds.³⁹ The deviation of the P-Au-P angles (164.3 and 159.8°) from the ideal value

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Table 1 Properties of the electron density at the selected bond critical points (BCP) according to the quantum theory of atoms in molecules (QTAIM) analysis of the bimetallic complexes 1-4

	BCP	ρ^a (e Å ⁻³)	$ V /G^b$	$E_{\rm INT}^{\ c} \left({\rm kJ \ mol}^{-1} \right)$	$\delta(A, B)^d$	$q^{e}\left(\mathbf{M}\right)$
1	Cu(1)···O(1)	0.325	1.04	-87.8	0.30	0.501
	Cu(1)····O(2)	0.310	1.02	-82.2	0.28	
	Cu(2)····O(1)	0.310	1.02	-82.2	0.28	
	$Cu(2)\cdots O(1)$	0.325	1.04	-87.8	0.30	
	P(2)-O(1)	1.374	1.35	-741.1	0.77	
	Cu–P	0.532	1.41	-120.1	0.67	
2	Ag(1)···O(1)	0.210	1.03	-42.9	0.20	0.400
	$Ag(1)\cdots O(2)$	0.219	1.01	-46.3	0.21	
	$Ag(2)\cdots O(1)$	0.219	1.01	-46.3	0.21	
	$Ag(2)\cdots O(1)$	0.210	1.03	-42.8	0.20	
	P(2) - O(1)	1.441	1.33	-808.2	0.82	
	Ag-P	0.481	1.36	-94.8	0.63	
3	Au(1)···O(1)	0.174	0.99	-29.9	0.21	0.062(Au1)
	$Au(1)\cdots O(2)$	0.174	0.99	-29.9	0.21	0.024(Au2)
	$Au(2)\cdots O(1)$	0.111	0.97	-15.8	0.11	` ′
	$Au(2)\cdots O(1)$	0.111	0.97	-15.8	0.11	
	P(2)-O(1)	1.466	1.33	-831.9	0.85	
	Au-P	0.716	1.69	-148.3	0.88	
4	Cu(1)···O(1)	0.336	1.04	-92.1	0.32	0.514(Cu)
	$Cu(1)\cdots O(2)$	0.336	1.04	-92.1	0.32	0.014(Au)
	$Au(2)\cdots O(1)$	0.116	0.99	-17.2	0.12	, ,
	$Au(2)\cdots O(1)$	0.116	0.99	-17.2	0.12	
	P(2)-O(1)	1.427	1.34	-792.8	0.82	
	Cu-P	0.524	1.39	-118.8	0.65	
	Au-P	0.730	1.73	-149.6	0.87	

^a Local electron density at the BCP. ^b Ratio of potential energy density and kinetic energy density. ^c Interaction energy between two interacting atoms. d Delocalization index between A and B (bonding) atoms. Total atomic charge of metal atoms.

of 180° could be attributed to some weak Au-O interactions in the solid state (Au-O distances are 2.986 and 2.745 Å), which are much less efficient than M-O bonds in 1 and 2, reflecting generally a higher coordination number of copper(1) and silver(1) than that of gold(1)40 and a lower affinity of Au(1) centers to hard electron donors. Due to the stereochemical arrangement of 3 the large Au···Au separation of 4.477 Å suggests no intramolecular metal-metal bonding.41 Interestingly, a mononuclear arrangement was suggested for the gold(I) complex $[Au(i-Pr-P^3O)]^+$ with a closely related ligand (i-Pr- $P^3O = (o-iPr_2P-(C_6H_4))_2P(O)Ph$) on the basis of the computationally optimized structure and NMR spectroscopic data.42

In the heterobimetallic complex 4 a tridentate mode of P^3O provides two types of ligand spheres ("P2O2" and "P2"), which successfully saturate four and two coordination vacancies of Cu(I) and Au(I) ions, respectively. The structural parameters of 4 (Table S5†) are somewhat comparable with the related values determined for homometallic congeners 1 and 3.

The ESI-MS of the dimeric 1 and 2 display dominating signals at m/z 709.1 and 753.1, respectively, which correspond to the monocations as a result of fragmentation under the conditions of the ESI experiment. The spectra of complexes 3 and 4, however, show the signals of the doubly charged dinuclear cations at m/z of 843.1 and 776.1. The observed isotopic patterns completely match the calculated ones for the proposed

 $[Cu(P^3O)]^+$, $[Ag(P^3O)]^+$, $[Au_2(P^3O)_2]^{2+}$ and $[AuCu(P^3O)_2]^{2+}$ stoichiometry (Fig. S1, ESI†). It has to be noted that in the case of 4 additional strong signals assigned to the presence of 1 and 3 were detected. This denotes the formation of a mixture of 1, 3 and 4 upon dissolving 4, which was also confirmed by NMR spectroscopic studies (vide infra).

The ³¹P{¹H} NMR spectra of 1 and 2 display two signals each and agree with idealized symmetrical structures of these complexes. The low field resonances at 42.0 ppm (1) and 36.1 ppm (2) can be assigned to the O=PPh moieties. The high field signals (-11.3 ppm and -1.3 for 1 and 2, respectively) of double integral intensities correspond to the metalcoordinated PPh₂ groups, which become essentially equivalent in solution. The presence of well resolved signals of the two isotopomers with $P^{-107, 109}$ Ag couplings ($J(P^{-109}Ag)$ 473 Hz and $J(P^{-107}Ag)$ 412 Hz) in the high field resonance of 2 additionally confirms coordination of both diphenylphosphine functions to silver(I) ions. The ¹H NMR data of 1 and 2 are also in accordance with the symmetrical arrangement of the complexes in solution. The corresponding spectra (Fig. S2 and S3†) are very much alike, showing poorly resolved resonances of the aromatic protons that prevent their complete assignment, but the number of the signals and their relative intensities indicate the equivalence of the P^3O ligands in fluid medium or possible dissociation of the dimers into the monomeric species, as suggested by the ESI-MS measurements.

Complex 3 demonstrates higher rigidity in solution compared to the Cu and Ag relatives. Its phosphorus NMR spectrum shows 3 resonances with 1:1:1 relative intensities (Fig. 3). The low field multiplets (37.5 and 32.5 ppm, dd $J_{\rm PP}$ ca. 6 Hz) from the nonequivalent PPh₂ moieties testify to the retaining of the structure found in the crystal state, in which two types of terminal phosphorus atoms are identified. The P centres apparently become different due to the chiral twisting of the 16-membered metallocycle, induced by the arrangement of the phenyl rings to minimise intramolecular repulsion. The proton NMR of 3 (Fig. S4†) is in line with this structural arrangement and supports the unsymmetrical coordination of P^3O to the gold centers.

NMR investigation of 4 revealed that in solution it produces a mixture of three compounds existing in dynamic equilibrium. The $^{31}P\{^{1}H\}$ NMR spectrum of 4 (Fig. 3) exhibits two sets of signals, which can be easily assigned to compounds 1 and 3 by comparing them with spectroscopic patterns obtained for the individual homometallic species. The third suite of resonances of equal intensities is completely compatible with the structure of heterometallic cation 4, and contains three signals, corresponding to phosphine oxide (45.3 dd, J_{PP} 11.2, 10.7 Hz) and diphenylphosphine groups coordinated to gold (42.8 s) and copper (–12.3 br) ions.

The freshly dissolved recrystallized complex 4 shows the presence of a dominant heterometallic form found in the crystal state, and gradually growing amounts of the homometallic complexes 1 and 3, until the ratio of the species becomes roughly equimolar in ca. 2 h (room temperature). Recrystallization of the mixture results in recovery of a uniform crystalline material of 4, as confirmed by the XRD analysis of several crystals. Additionally, the crystalline form of 4 demonstrates unique photophysical characteristics different from those of 1 and 3 (see below) that clearly indicate the difference in the nature of these solid materials and supports phase purity of 4. A similar behavior of heterometallic phosphine coinage metal complexes, which involves migration of the metal ions and appearance of multicomponent mixtures in solution, was described earlier by us for some other multidentate ligand complexes. 26,29,43

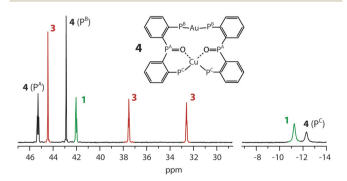


Fig. 3 162 MHz $^{31}P{^{1}H}$ NMR spectrum of the equilibrated mixture of complexes formed upon dissolution of 4 (dmso- d_{6} , 298 K). The relative molar ratio of 1:3:4 is ca. 1:1:1.

The nature of the intramolecular interactions was investigated *via* topological charge density analysis utilizing the Quantum Theory of Atoms in Molecules (QTAIM). Table 1 presents selected properties of the electron density at the corresponding bond critical points for the bimetallic complexes 1–4. The interaction energies, which were calculated to be half of the potential energy density according to the method of Espinosa *et al.*, ⁴⁴ show a clear trend in the strength of the M···O interactions, Cu > Ag > Au. The difference in the bond strength is most pronounced in the heterobimetallic complex 4, where the interaction energies are –92 kJ mol⁻¹ and –17 kJ mol⁻¹ at the Cu···O and Au···O bond critical points, respectively. The strength of the interaction is directly proportional to the amount of local electron density at the bond critical point.

However, even though the interaction energies are rather notable, at least in the case of copper complex 1, the M···O interactions are mostly non-covalent in nature, which can be seen in the ratio of the potential energy density and kinetic energy density, $|V|/G \sim 1$, and in the small amount of shared electrons given by the delocalization index, $\delta(A, B)$. Because of the non-covalent nature of the M···O interactions, the original properties of the P(2)-O(1) interaction are only slightly changed, when compared to the values in the free ligand. For example, the ρ , |V|/G, E_{INT} and $\delta(A, B)$ values are 1.500 e Å⁻³, 1.32, -868.2 kJ mol⁻¹, and 0.88, respectively, for the freely optimized P^3O . On the other hand, the weaker contacts between metal and oxygen were found to increase the phosphorusmetal interaction, thus leading to an opposite trend in the M-P BCPs, Au > Ag > Cu, which can be seen in Table 1, where average values at the M-P interactions are presented for complexes 1-4. As can be seen in Fig. S5,† no metal-metal interactions were found in the bimetallic compounds. It should be noted that although the optimized geometry of 2 was found to be more symmetrical than the experimental one, for which the charge density analysis led to two stronger Ag···O interactions and two weaker ones, this did not change either the nature of the Ag...O contacts or the trend in the strength of bonding in the bimetallic species.

Mononuclear halide complexes $M(P^3O)X$ and $M(P^3)X$. The copper(1) compounds of general formula $Cu(P^3O)X$, X = Cl (5), Br(6), I(7) can be easily obtained by treating the corresponding halide with a stoichiometric amount of the P^3O ligand (Scheme 2). The synthesis of $Ag(P^3O)X$ congeners (X = Cl (8), Br (9), I (10)) was carried out using a slightly modified procedure that involved the coupling of P^3O with $AgPF_6$, followed by the addition of the respective NBu_4X salt (Scheme 2).

Complexes 5–10 were isolated by crystallization as colourless to pale yellow materials in good yields (Scheme 2). The XRD analysis revealed an essentially similar geometry for these mononuclear compounds (Fig. 4 and S6†). The structural motif implies a quasi-tetracoordinate environment of the metal ions, which involves the regular metal–phosphine/halide bonding⁴⁵ along with some metal–O—PPh interactions (Table S6†).

The weakness of phosphine oxide coordination to Cu and Ag ions evidently determines trigonal planar geometry of the **Dalton Transactions**

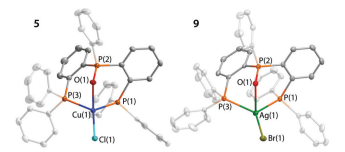


Fig. 4 Molecular views of complexes 5 and 9. Thermal ellipsoids are shown at the 50% probability level. H atoms are omitted for clarity.

P₂X ligand at the metal centers that is similar to that found for the congener Ag(i-Pr-P³O)X. 42 The Cu-O distances particularly in 5 and 6 are visibly longer than those in 1, 4 and other related copper species, 13,32 but are comparable to the value reported for the $[Cu(\kappa^2-DPEphos)(\kappa^2-DPEphosO)]^+$ complex.⁴⁶ In the silver species 9 and 10 the Ag-O separations are similarly longer than those in 2 and congener compounds, ³⁶ which points to a rather insignificant interaction of the O=PPh moiety with silver ions.

The solution NMR spectroscopic data of the halide complexes 5-10 are in complete agreement with the molecular arrangement found in the crystal. Their ³¹P{¹H} spectra display two signals with a 1:2 intensity ratio, which correspond to the P-oxide fragment and the metal-coordinated equivalent PPh2 groups, respectively (see the Experimental section).

We have also synthesized the non-oxidized triphosphine halide compounds $M(P^3)X$ (M = Cu, X = Cl (11), Br (12), I (13); $M = Ag, X = Cl (14), Br (15), I (16), P^3 = bis(2-diphenylphosphino$ phenyl)phenyl phosphine) to compare their photophysical characteristics with those of $M(P^3O)X$ complexes 5-10. The species 11 and 14-16 were employed earlier in the fabrication of luminescent membranes and electroluminescent devices, 47

and 14 was characterized spectroscopically and crystallographically. 48 Complexes 11, 12, 15 and 16 were studied by means of XRD analysis (Fig. S7†), which revealed that 11, 12, and 15 are isomorphous to 14 (space group $P2_1/n$) with small alterations of the unit cell parameters. Additionally, $M(P^3)X$ compounds closely resemble the related cyanide complexes $M(P^3)CN$ we described recently.²⁹ Complex **16** crystallizes in the $P2_1/c$ type space group with somewhat different cell dimensions, but its molecular geometry is virtually identical to that of 11, 12, and 15 (Fig. S6†). The tetracoordinate ligand sphere of the metal ions in $M(P^3)X$ compounds is completed by the tridentate phosphine P^3 and the halide X that is in line with previous reports.^{29,48} The selected structural parameters are listed in Table S7.† The NMR spectra are consistent with the structural data and indicate that all the P donors are bound to the corresponding metal centers (see the Experimental section).

Selected mononuclear halides were also optimized and the intramolecular interactions were investigated. The compounds studied were copper complexes 7 and 13, and silver complexes 10 and 16. The results of the QTAIM analysis are given in Table S8 of the ESI.† The relative trends in the properties of the electron density in the monomeric compounds were essentially the same as those observed in the bimetallic species, the Cu···O interaction with the P^3O ligand being stronger than the corresponding Ag...O contact. The same trend was also found for $M\cdots P(2)$ interactions with the P^3 ligand. Interestingly, in the complexes with P^3 ligands, the Cu···P(2) is similar to the other Cu-P interactions, but the Ag-P in 16 is notably weaker, which becomes apparent from the larger optimized distance.

Solid state photophysical properties

The title compounds do not demonstrate appreciable photoluminescence in solution, and their photophysical behavior was investigated in the solid state only. The relevant data are listed in Table 2 and the spectroscopic patterns are shown in Fig. 5-7 and S8-S10.†

Table 2 Solid state photophysical properties of 1-4 and 7-16

	λ_{ex} , nm		λ_{em} , nm		$ au_{ m obs}{}^a, \mu { m s}$. 0/	, h -1	, c -1
	298 K	77 K	298 K	77 K	298 K	77 K	$\Phi_{ m em},\%$ 298 K	$k_{\rm r}^{\ \ b},{\rm s}^{-1}$	$k_{\rm nr}^{\ c}$, s ⁻¹
1	364	365	604	604	0.35	174.1	0.5	1.4×10^{4}	2.0×10^{6}
2	332	324	500	485	10.9	2354.3	5.9	5.0×10^{3}	8.5×10^4
3	330	325	548	564	6.7	82.3	28.9	4.3×10^{4}	1.0×10^{5}
4	360	363	538	546	5.5	777.5	25.8	4.7×10^{4}	1.3×10^{5}
7	394	362, 380	621	624	0.5	30.7	0.8	1.7×10^{4}	2.1×10^{6}
8	310, 372	340	480	495	64.8	482.0	17.7	2.7×10^{3}	1.3×10^{4}
9	362	340	467	464	19.0	529.9	7.5	3.9×10^{3}	4.7×10^{4}
10	362	353	488	486	38.1	95.7	20.4	5.4×10^{3}	2.1×10^{4}
11	370	363, 397	517	520	9.0	565.8	21.0	2.3×10^{4}	8.7×10^{4}
12	370	363, 397	507	523	11.7	794.4	36.3	3.1×10^{4}	5.4×10^{4}
13	370	311	504	515	10.5	136.9	37.2	3.5×10^{4}	6.0×10^{4}
14	335, 367	329, 375	521	535	27.0	595.7	39.5	1.5×10^{4}	2.2×10^{4}
15	335, 395	329, 375	521	535	25.6	537.7	40.5	1.6×10^{4}	2.3×10^{4}
16	335, 396	329, 375	513	525	13.5	179.9	38.9	2.8×10^4	4.5×10^4

^a Average emission lifetimes for 1, 4 and 7 for the two-exponential decay determined using the equation $\tau_{av} = (A_1\tau_1^2 + A_2\tau_2^2)/(A_1\tau_1 + A_2\tau_2)$, where A_i is the weight of the *i*-exponent. bk_r were estimated using $\Phi/\tau_{\rm obs}$. ${}^ck_{\rm nr}$ were estimated using $k_r(1-\Phi)/\tau_{\rm obs}$.

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Fig. 5 Normalized solid state excitation (dotted lines) and emission (solid lines) spectra of 1–4 at 298 K.

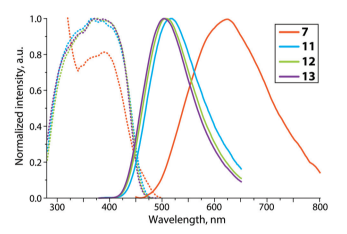


Fig. 6 Normalized solid state excitation (dotted lines) and emission (solid lines) spectra of copper complexes 7 and 11–13 at 298 K.

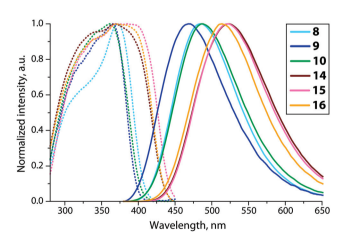


Fig. 7 Normalized solid state excitation (dotted lines) and emission (solid lines) spectra of copper complexes 8–10 and 14–16 at 298 K.

The dinuclear copper (1) and silver (2) complexes are weakly luminescent at room temperature and exhibit broad emission peaks centered at 604 ($\Phi_{\rm em}$ 0.5%) and 500 nm

($\Phi_{\rm em}$ 5.9%), respectively. Cooling the samples to 77 K doesn't affect the emission maximum of 1 and causes some blue shift for 2 (Table 2, Fig. S7†) that allows excluding for these species a phenomenon of thermally activated delayed fluorescence, for which a small red shift of the emission band is typically observed upon lowering the temperature. 34,49 The goldcontaining compounds show the emission bands at 548 (3) and 538 (4) nm with considerably larger quantum efficiencies ($\Phi_{\rm em}$ 28.9 and 25.8%), which is reflected by increased radiative rate constants (k_r) with respect to the rates of radiationless deactivation ($k_{\rm nr}$, Table 2). The effect evidently results from larger spin orbit coupling in gold complexes and a higher ISC rate compared to analogous compounds of the first and second transition row metals. The lifetimes for 1-4 are found to be in the microsecond domain at 298 K and show a dramatic growth upon lowering the temperature, reaching the value of 2354.3 µs (2, 216-fold increase) that is comparable to the demeanor of some [Ag(diphosphine)₂]⁺ compounds.⁵⁰ The emission profile of 4 does not reveal any detectable shoulders to evidence the presence of homometallic complexes 1 and 3 (vide supra), which were identified in the solution of 4 by ESI-MS and NMR spectroscopy. This observation additionally supports the very selective crystallization of heterobimetallic complex 4.

The mononuclear copper chloride and bromide complexes 5 and 6 are not luminescent either at 298 K or at the temperature of liquid nitrogen. Their iodide congener 7 however exhibits weak orange emission (621 nm, $\Phi_{\rm em}$ 0.8%, Fig. 6). A comparison of 7 with non-oxidized complexes $Cu(P^3)X$ (11-13) reveals the dramatic effect of changing the phosphine PPh group to phosphine oxide O=PPh on the photophysical properties of copper halides. Unlike 7, the $Cu(P^3)X$ species demonstrate a moderately strong blue-green luminescence (504–517 nm, $\Phi_{\rm em}$ 21–37%), which is only slightly sensitive to the nature of the X ligand (Fig. 6 and S9†). The significant decrease of emission energy for 7 with respect to 11-13 can be tentatively assigned to a possibly larger flexibility of the $Cu(P^3O)X$ framework, which could facilitate the formation of a planar metal geometry in the MLCT excited state. The latter, as generally accepted, is able to provide nonradiative ways of relaxation.51

The two families of silver complexes Ag(P^3O)X (8–10) and Ag(P^3)X (14–16) display a very different phosphine ligand effect on the photophysics of the solid materials compared to the copper relatives. The triphosphine compounds 14–16 show rather intense room temperature emission with quantum yields of around 40% (Table 2). The broad bands with maxima in the range 513–521 nm are slightly red shifted with respect to the corresponding Cu species 11–13 at 298 K (Fig. 7), and in a similar way experience a small decrease of luminescence energies at 77 K (Fig. S10†). The emission bands for oxygen-modified complexes 8–10 show a visible hypsochromic shift of 25–54 nm, accompanied by at least 2-fold drop of intensities, which, however, is not as drastic as in the case of copper congeners. The observed difference of the photophysical characteristics between the

series 8-10 and 14-16 could be attributed to apparently weak interaction of Ag-O=PPh2 and thus formally a lower coordination number of silver ions in $Ag(P^3O)X$ complexes that

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evidently affects the frontier orbitals and the energies of electronic transitions.

In order to rationalize the difference in photophysical behavior between the complexes with P^3O ligands and P^3 ligands, we studied the emission properties by optimizing the first excited triplet state for copper compounds 7 and 13 as well as for silver complexes 10 and 16. The calculated emission wavelengths are well consistent with the experimental spectra for the monometallic compounds. However, the emission energies were considerably overestimated in the case of the bimetallic complexes, even though the general trend could be correctly represented. Obviously, the DFT method was not able to describe the electron distribution in the triplet state for the dimeric compounds. This can be seen as an example of complex 1 in Fig. S11,† which shows the HOMO and LUMO orbitals of the singlet state, where the MOs are evenly distributed in the vicinity of both copper atoms, leading to correct absorption energies. On the contrary, in the triplet state the singly occupied MOs are very unsymmetrically expanded around different metals, leading to a less stable triplet state, and hence overestimated emission energies. Because of this inconsistency we focused on the electronic features of the monometallic complexes. The appearance of the singly occupied molecular orbitals (SOMOs, HSOMOs) is compared along with the HOMOs of the S₀ state in Fig. 8 and 9. In the singlet ground state, the HOMO orbitals mainly consist of a combination of metal d-orbitals with the halogen p-orbital. There is a notable difference between the complexes with different ligands, since in the ground state, the oxygen and P(2) orbitals of the P^3O ligand do not participate in HOMO (complexes 7 and 10), which leads to its higher energy compared to the complexes 13 and 16, where p-orbitals of all phosphorus

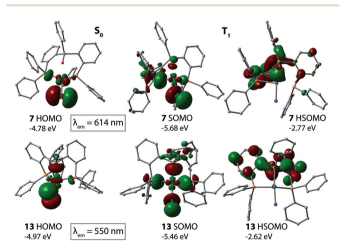


Fig. 8 The appearance of the highest occupied orbitals in the singlet ground state and the first excited triplet state of copper complexes 7 and 13. The emission wavelength was estimated from the total energy difference of the states.

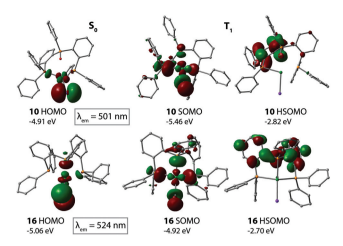


Fig. 9 The appearance of the highest occupied orbitals in the singlet ground state and the first excited triplet state of silver complexes 10 and 16. The emission wavelength was estimated from the total energy difference of the states.

atoms of P^3 interact with the metal center and contribute to номо.

When one electron is excited from HOMO, the remaining electron forms a SOMO orbital (singly occupied molecular orbital), the energy of which is considerably lowered in the case of the oxygen containing ligand P^3O , as much as 0.90 eV in 7 and 0.55 eV in 10. In both cases, the stabilization results from the substantial increase of the contribution of metal d-orbitals in SOMO, which was not observed in the ground state $(18\% \to 31\% \text{ in 7 and } 11\% \to 21\% \text{ in 10})$. On the contrary, 16 showed slight destabilization of the energy of SOMO compared to the singlet state HOMO, probably due to decreasing contribution from the iodide ligand, which is also observed, though not so extensively, for other compounds upon excitation. The full fragment analysis of the frontier MOs is presented in Table S9.† This discrepancy can also account for the different trends in the emission energies between the series of copper and silver compounds.

The highest singly occupied MO (HSOMO) is rather similar in all cases; it is formed as a combination of the phenylene ring orbitals. Because of the larger flexibility of the framework with the oxygen containing ligand, the optimized geometry in the triplet state was less symmetrical for P3O-containing complexes than that found for their P^3 -based relatives. Therefore, the HSOMO orbital is less evenly distributed along the ligand. The rather notable contribution of oxygen p-orbitals leads to the localization of HSOMO more in the central part of the ligand, which has a small stabilizing effect. However, this has a minimal effect on the energetics of the HSOMOs and, consequently, on the photophysical characteristics. Basically, the emission can be assigned to the MXLCT type for all mononuclear compounds. Nevertheless, the variations in the metal-ligand interactions in the series with P^3O and P^3 ligands lead to the observed differences in luminescence behavior.

Conclusions

Paper

The phosphine-phosphine oxide hybrid ligand P^3O , which shows variable binding ability with respect to d10 coinage metal ions, was used for the preparation of a series of novel mono- and dinuclear complexes. The combination of soft (P) and hard (O) donor functions allows for an efficient adaptation of the ligand environment to the preferable coordination number of the Cu^I, Ag^I and Au^I centers that preserves the nuclearity of the title dicationic compounds $[MM'(P^3O)_2]^{2+}$ (M = M' = Cu (1), Ag (2), Au (3); M = Au, M' = Cu (4)). The nature of metal-ligand interactions was elucidated using the OTAIM computational approach, which revealed a clear trend in the strength of the M···O bonds, Cu > Ag > Au. A similar tendency was estimated for the neutral monometallic halide complexes $M(P^3O)$ Hal (M = Cu (5-7), Ag (8-10)). The studied complexes exhibit weak to moderate room temperature photoluminescence in the solid state except 5 and 6, which are virtually not emissive even at 77 K. The bimetallic compounds demonstrate quantum efficiency up to 28.9% (3, $\lambda_{\rm em}$ = 548 nm), while the mononuclear ones reach a $\Phi_{\rm em}$ value of 20.4% (10, $\lambda_{\rm em}$ = 488 nm). The effect of partial phosphine oxidation on the photophysical properties was analyzed comparing the $M(P^3O)$ Hal species with their $M(P^3)$ Hal congeners. For the copper complexes the presence of the phosphine-oxide group is apparently detrimental, leading to a dramatic decrease of quantum yields that is accompanied by a 117 nm bathochromic shift (7 vs. 13). On the contrary, for silver complexes the emission intensity is much less influenced by the introduction of oxygen, but the wavelength shows a substantial blue shift (up to 54 nm for 9 vs. 15) that illustrates a convenient possibility of fine tuning the luminescence characteristics of this class of inorganic material.

Experimental

General comments

(2-Bromophenyl)diphenylphosphine, ³¹ bis(2-diphenylphosphinophenyl)phenyl phosphine (*P*³)²⁷ and the complex AuCl(tht) (tht = tetrahydrothiophene)⁵² were synthesized according to published procedures. Tetrahydrofuran (THF) was distilled over Na-benzophenone ketyl under a nitrogen atmosphere prior to use. Other reagents and solvents were used as received. The solution ¹H, ³¹P{¹H} and ¹H-¹H COSY NMR spectra were recorded on a Bruker Avance 400 spectrometer. Mass spectra were recorded on a Bruker micrOTOF 10223 instrument in the ESI⁺ mode. Microanalyses were carried out in the analytical laboratory of the University of Eastern Finland.

Bis(2-(diphenylphosphino)phenyl)phenyl phosphine oxide (P^3O). The synthesis was carried out under a nitrogen atmosphere. A solution of 2-(bromophenyl)diphenylphosphine (2.0 g, 5.8 mmol) in THF (20 ml) was cooled to -78 °C, and a 1.6 M solution of n-BuLi (4.0 ml, 6.4 mmol) was added dropwise within 10 min. The resulting yellow solution was stirred

for 1 hour at this temperature and treated dropwise with OPPh₂ (0.50 g, 2.56 mmol). The reaction mixture was then stirred below -70 °C for 2 hours, and then overnight at room temperature. The reaction was quenched with methanol (2 ml) and evaporated. The amorphous creamy residue was washed with methanol (3 × 20 ml), dried, dissolved in dichloromethane and passed through a pad of silica gel (150 mesh, 2 × 5 cm) and evaporated to dryness. Additional washing with methanol (10 ml) gave a colorless solid of sufficient purity (1.33 g, 80%). 31 P{ 1 H} NMR (CD₂Cl₂, 298 K, δ): 31.9 (t, 3 J_{PP} 12 Hz, 1P, P(O)Ph), -13.6 (d, 3 J_{PP} 12 Hz, 2P, PPh₂). 1 H NMR (CD₂Cl₂, 298 K, δ): 7.67 (dd, J_{PH} 11.9 Hz, J_{HH} 7.9 Hz, 2H), 7.39–7.55 (m, 7H), 7.19–7.39 (m, 16H), 7.05–7.19 (m, 8H). Anal. calc. for C₄₂H₃₃OP₃ (%): C 78.01; H 5.14. Found: C 77.65; H 4.82.

 $[Cu(P^3O)]_2(PF_6)_2$ (1). $Cu(NCMe)_4PF_6$ (43.6 mg, 0.117 mmol) was dissolved in acetone (5 ml) and added to a solution of P^3O (75.6 mg, 0.117 mmol) in dichloromethane (5 ml). The colorless reaction mixture was stirred for 2 h and evaporated to dryness. An amorphous solid was recrystallized by a gas-phase diffusion of diethyl ether into acetone solution of 1 to afford colorless crystalline material (90 mg, 90%). ES MS (m/z): $[CuP^3O]^+$ 709.1 (calc. 709.1). $^{31}P\{^1H\}$ NMR (dmso- d_6 , 298 K, δ): 42.0 (br, 2P, P(O)Ph), -11.3 (br, 4P, PPh₂), -144.6 (sept, 2P, PF₆). ¹H NMR (dmso- d_6 , 298 K, δ): 7.69 (m, 2H, para-H Ph-PO), 7.58 (m, 8H, meta + ortho-H Ph-PO), 7.48 (m, 4H, meta-H C_6H_4), 7.46 (m, 4H, meta-H C_6H_4), 7.39 (t, 4H, meta-H Ph), 7.33–7.51 (m, 24H), 7.29 (dd, J_{HH} 7.4 and 7.6 Hz, 8H, meta-H Ph), 7.03 (dd, J_{PH} 11.9 Hz, J_{HH} 6.2 Hz, 8H, ortho-H Ph), 6.95 (ddd, J_{PH} 11.8 Hz, J_{HH} 7.8 and 3.8 Hz, 4H, ortho-H C₆H₄). Anal. Calc. for Cu₂C₈₄H₆₆O₂P₈F₁₂ (%): C 58.99; H 3.89. Found: C 58.51; H 4.12.

[Ag(P^3O)]₂(CF₃SO₃)₂ (2). Prepared analogously to 1 using AgCF₃SO₃ (28.4 mg, 0.111 mmol) and P^3O (71.5 mg, 0.111 mmol) (76 mg, 76%). ES MS (m/z): [Ag P^3O]⁺ 753.1 (calc. 753.1). ³¹P{¹H} NMR (dmso- d_6 , 298 K, δ): 36.1 (t, ³ $J_{\rm PP}$ 5 Hz, 2P, P(O)Ph), -1.3 (two d, $J({\rm P}^{-109}{\rm Ag})$ 473 Hz, $J({\rm P}^{-107}{\rm Ag})$ 412 Hz, 4P, PPh₂). ¹H NMR (dmso- d_6 , 298 K, δ): 7.59 (t, $J_{\rm HH}$ 7.4 Hz, 2H, para-H Ph-PO), 7.44 (dm, 4H, meta-H Ph-PO), 7.39 (m, 4H, meta-H C₆H₄), 7.31 (m, 4H, ortho-H Ph-PO), 7.30 (m, 4H, meta-H C₆H₄), 7.27-7.47 (m, 36H), 7.22 (dd, $J_{\rm PH}$ 12.1 Hz, $J_{\rm HH}$ 6.2 Hz, 8H, ortho-H Ph), 6.91 (ddd, $J_{\rm PH}$ 12.2 Hz, $J_{\rm HH}$ 7.5, 3.6 Hz, 4H, ortho-H C₆H₄). Anal. Calc. for Ag₂C₈₆H₆₆O₈S₂P₆F₆ (%): C 57.16; H 3.68; S 3.55. Found: C 57.15; H 4.17; S 3.34.

[Au(P^3O)]₂(PF₆)₂ (3). AuCl(tht) (32.4 mg, 0.101 mmol) and P^3O (65.4 mg, 0.101 mmol) were dissolved in dichloromethane (10 ml), and a solution of AgPF₆ (25.6 mg, 0.101 mmol) in acetone (5 ml) was added. A flaky colorless precipitate of AgCl instantly formed. The reaction mixture was stirred for 30 min in the absence of light. AgCl was removed by filtration, and the resulting colorless solution was evaporated. An amorphous residue was thoroughly washed with diethyl ether and recrystallized by a gas-phase diffusion of diethyl ether into acetone solution of 3 at 278 K to give a colorless crystalline material (71 mg, 72%). ES MS (m/z): [Au₂(P^3O)₂]²⁺ 843.1 (calc. 843.1). ³¹P{¹H} NMR (dmso- d_6 , 298 K, δ): 44.4 (s, 2P, P(O)Ph), 37.5

(dd, J_{PP} ca. 6 Hz, 2P, PPh₂), 32.5 (dd, J_{PP} ca. 6 Hz, 2P, PPh₂), -144.6 (sept, 1P, PF₆). ¹H NMR (dmso- d_6 , 298 K, δ): 8.16 (dd, J_{HH} 7.4, 7.6 Hz, 1H), 7.96 (dd, J_{HH} 7.4, 7.6 Hz, 1H), 7.79 (dm, J_{HH} 7.4 Hz, 1H), 7.76 (dm, J_{HH} 7.4 Hz, 1H), 7.60–7.71 (m, 5H), 7.53 (t, J_{HH} 7.8 Hz, 2H), 7.42 (m, 2H), 7.35 (m, 4H), 7.29 (dd, J_{HH} ca. 7.4 Hz, 1H), 7.09 (m br, 4H), 6.80-6.93 (m, 5H),

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6.38-6.54 (m, 6H). Anal. calc. for $Au_2C_{84}H_{66}O_2P_8F_{12}$ (%): C 51.03; H 3.37. Found: C 50.94; H 3.50.

 $[AuCu(P^3O)_2](PF_6)_2$ (4). 3 prepared in situ as described above from AuCl(tht) (14.9 mg, 0.046 mmol) and P^3O (30.0 mg, 0.046 mmol) was added to a solution of 1 (prepared from $Cu(NCMe)_4PF_6$ (17.2 mg, 0.046 mmol) and P^3O (30.0 mg, 0.046 mmol)) in dichloromethane (5 ml). The reaction mixture was stirred for 2 hours, the solvents were evaporated and the solid residue was recrystallized by a gas-phase diffusion of diethyl ether into an acetone/methanol (3:1 v/v mixture) solution of 4 at 278 K to give a yellow crystalline material (72 mg, 84%). $^{31}P\{^{1}H\}$ NMR (dmso- d_6 , 298 K, δ): the spectrum corresponds to the mixture of the 3 complexes, see discussion; 45.3 (dd, J_{PP} 11.2, 10.7 Hz, 1P, P(O)Ph), 44.4 (3), 42.8 (s, 1P, Au-PPh₂), 42.0 (1), 37.5 (3), 32.5 (3), -11.3 (1), -12.3 (br s, 1P, PPh₂), -144.6 (sept, 1P, PF₆). ES MS (m/z): [AuCu(P^3O)₂]²⁺: 776.12 (calc. 776.13). Anal. calc. for AuCuC₄₂H₃₃OP₄F₆ (%): C 54.72; H 3.61. Found: C 54.56; H 3.65.

General procedure for the synthesis of $CuX(P^3O)$ (X = Cl, Br, I) complexes (5-7)

CuX (0.123 mmol) and $P^{3}O$ (80 mg; 0.123 mmol) were suspended in dichloromethane (10 ml). The reaction mixture was stirred for 5 hours to give a transparent light yellow solution. The solvent was evaporated and the solid residue was recrystallized.

 $CuCl(P^3O)$ (5). Recrystallized by a gas-phase diffusion of diethyl ether into its acetone solution at 278 K to give a yellow crystalline material (85 mg, 92%). ³¹P{¹H} NMR (dmso-d₆, 298 K, δ): 39.2 (s, 1P, P(O)Ph), -14.1 (s, 2P, PPh₂). ¹H NMR (dmso- d_6 , 298 K, δ): 7.56–7.65 (m, 5H), 7.44–7.55 (m, 8H), 7.30-7.43 (m, 10H), 7.25 (dd, J_{HH} ca. 7.7 Hz, 4H), 6.92 (m, 6H). Anal. calc. for CuC₄₂H₃₃OP₃Cl (%): C 67.65; H 4.46. Found: C 67.65; H 4.39.

 $CuBr(P^3O)$ (6). Recrystallized by a gas-phase diffusion of diethyl ether into its acetone/methanol solution (1:1 v/v mixture) at 278 K to give a yellow crystalline material (86 mg, 89%). $^{31}P\{^{1}H\}$ NMR (dmso- d_{6} , 298 K, δ): 39.6 (br s, 1P, P(O)Ph), -14.5 (br s, 2P, PPh₂). ¹H NMR (dmso- d_6 , 298 K, δ): 7.64 (m, 1H), 7.44-7.59 (m, 12H), 7.31-7.44 (m, 10H), 7.25 (dd, J_{HH} ca. 7.6 Hz, 4H), 6.97 (m, 4H), 6.89 (m, 2H). Anal. calc. for CuC₄₂H₃₃OP₃Br (%): C 63.85; H 4.21. Found: C 63.82; H 4.18.

 $CuI(P^3O)$ (7). Recrystallized by a gas-phase diffusion of diethyl ether into its acetone/methanol (5:2 v/v mixture) at 278 K to give a yellow crystalline material (93 mg, 90%). ³¹P{¹H} NMR (dmso- d_6 , 298 K, δ): 37.63 (br s, 1P, P(O)Ph), -14.27 (br s, 2P, PPh₂). ¹H NMR (dmso- d_6 , 298 K, δ): 7.65 (m, 1H), 7.44–7.55 (m, 12H), 7.37–7.43 (m, 4H), 7.35 (dd, $J_{\rm HH}$ ca. 7.2 Hz, 6H), 7.25 (dd, J_{HH} ca. 7.5 Hz, 4H), 7.00 (m, 4H), 6.86

(m, 2H). Anal. calc. for CuC₄₂H₃₃OP₃I (%): C 60.26; H 3.97. Found: C 59.98; H 4.00.

General procedure for the synthesis of $AgX(P^3O)$ (X = Cl, Br, I) complexes (8-10)

AgPF₆ (25.3 mg, 0.1 mmol) was dissolved in acetone (5 ml) and a solution of the corresponding tetrabutyl ammonium halide NBu₄X (0.1 mmol) in dichloromethane (10 ml) was added. The suspension was stirred for 10 min in the absence of light. Then P^3O (64.6 mg, 0.1 mmol) was added to the reaction mixture, which was stirred for 1 hour to give a colorless transparent solution. The solvents were evaporated and the solid residue was recrystallized by a gas-phase diffusion of diethyl ether into a dichloromethane solution of a crude complex at 278 K to afford a colorless crystalline material.

 $AgCl(P^3O)$ (8). Yield 56 mg, 71%. $^{31}P\{^1H\}$ NMR (CD₂Cl₂, 298 K, δ): 35.9 (t, J_{PP} 5.6 Hz, 1P, P(O)Ph), -1.5 (br d, $J_{\rm PAg}$ 348 Hz, 2P, PPh₂). ¹H NMR (CD₂Cl₂, 298 K, δ): 7.53 (m, 1H), 7.11-7.44 (m, 30H), 6.99 (m, 2H). Anal. calc. for AgC₄₂H₃₃OP₃Cl (%): C 63.86; H 4.21. Found: C 63.45; H 4.23.

 $AgBr(P^3O)$ (9). Yield 72 mg, 86%. ${}^{31}P{}^{1}H{}$ NMR (CD₂Cl₂, 298 K, δ): 36.0 (t, J_{PP} 5.3 Hz, 1P, P(O)Ph), -2.0 (br d, J_{PAG} 354 Hz, 2P, PPh₂). ¹H NMR (CD₂Cl₂, 298 K, δ): 7.53 (m, 1H), 7.11-7.45 (m, 30H), 6.99 (m, 2H). Anal. calc. for AgC₄₂H₃₃OP₃Br (%): C 60.45; H 3.99. Found: C 60.47; H 4.19.

 $AgI(P^3O)$ (10). Yield 69 mg, 78%. ${}^{31}P\{{}^{1}H\}$ NMR (CD₂Cl₂, 298 K, δ): 35.9 (t, J_{PP} 4.7 Hz, 1P, P(O)Ph), -2.75 (br d, J_{PAG} ca. 222 Hz, 2P, PPh₂). 1 H NMR (CD₂Cl₂, 298 K, δ): 7.53 (m, 1H), 7.11-7.44 (m, 30H), 6.99 (m, 2H). Anal. calc. for AgC₄₂H₃₃OP₃I (%): C 57.23; H 3.77. Found C 57.48; H 3.99.

General procedure of $CuXP^3$ (X = Cl, Br, I) complexes (11–13)

CuX (0.123 mmol) and P^3 (77.5 mg, 0.123 mmol) were suspended in dichloromethane (10 ml). The reaction mixture was stirred overnight to give a transparent light yellow solution. The solvent was evaporated and the solid residue was recrystallized by a gas-phase diffusion of diethyl ether into a dichloromethane solution of a crude complex at 278 K to give a yellow crystalline material.

CuCl P^3 (11). Yield 73 mg, 81%. ${}^{31}P{}^{1}H{}$ NMR (dmso- d_6) 298 K, δ): -6.3 (br d, J_{PP} 162 Hz, 2P, PPh₂), -13.3 (t, J_{HH} 162 Hz, 1P, PPh). ¹H NMR (dmso- d_6 , 298 K, δ): the signals are broadened 7.9 (m, 4H), 7.75 (m, 2H), 7.60 (m, 2H), 7.05-7.54 (m, 21H), 6.45 (m, 4H). Anal. calc. for CuC₄₂H₃₃P₃Cl (%): C 69.14; H 4.56. Found: C 68.81; H 4.52.

CuBr P^3 (12). Yield 86 mg, 88%. ${}^{31}P{}^{1}H{}$ NMR (dmso- d_6 , 298 K, δ): -7.2 (br d, J_{PP} ca. 150 Hz, 2P, PPh₂), -14.1 (br t, J_{PP} ca. 150 Hz, 1P, PPh). ¹H NMR (dmso- d_6 , 298 K, δ): 7.85 (m, 4H, PPh₂), 7.74 (m, 2H, C₆H₄), 7.61 (dd, J_{HH} ca. 7.5 Hz, 2H, C_6H_4), 7.51 (t, J_{HH} ca. 7.5 Hz, 3H), 7.28-7.46 (m, 11H), 7.09-7.25 (m, 8H), 6.50 (m, 4H, PPh₂). Anal. calc. for CuC₄₂H₃₃P₃Br (%): C 65.17; H 4.30. Found C 64.79; H 4.31.

 $CuIP^3$ (13). Yield 93 mg, 90%. ${}^{31}P{}^{1}H$ NMR (dmso- d_6 , 298 K, δ): -7.9 (br unresolved, 2P, PPh₂), -13.61 (br unresolved, 1P, PPh). ¹H NMR (dmso- d_6 , 298 K, δ): the signals are broadened 7.76 (m, 4H), 7.61 (dd, J_{HH} ca. 7.4 Hz, 2H), 7.54 (t, J_{HH} 7.4 Hz,

2H), 7.11-7.45 (m, 21H), 6.59 (m, 4H). Anal. calc. for CuC₄₂H₃₃P₃I (%): C 61.44; H 4.05. Found: C 61.22; H 4.06.

General procedure of $AgXP^3$ (X = Cl, Br, I) complexes (14–16)

AgPF₆ (25.3 mg, 0.1 mmol) was dissolved in acetone (5 ml) and a solution of the corresponding tetrabutyl ammonium halide NBu₄X (0.1 mmol) in dichloromethane (10 ml) was added. The suspension was stirred for 10 min in the absence of light. Then P^3 (63.0 mg, 0.1 mmol) was added to the reaction mixture, which was stirred for 1 hour to give a colorless transparent solution. The solvents were evaporated and the solid residue was recrystallized by a gas-phase diffusion of diethyl ether into a dichloromethane solution of a crude complex at 278 K to afford a colorless crystalline material.

AgClP³ (14). Colorless crystalline material (57.2 mg, 74%). ³¹P{¹H} NMR (CD₂Cl₂, 298, δ): -8.3 (dd, J_{PP} 209 Hz, J_{PAg} 272 and 306 Hz, 2P, PPh₂), -27.40 (td, J_{PP} 209 Hz, J_{PAg} 121 Hz, 1P, PPh). ¹H NMR (CD₂Cl₂, 298, δ): 7.65 (m, 4H), 7.23-7.37 (m, 17H), 7.18 (dd, J_{HH} 7.0 and 7.5 Hz, 4H), 7.03-7.12 (m, 4H), 6.94 (m, 4H). Anal. calc. for AgC₄₂H₃₃P₃Cl (%): C 65.18; H 4.29. Found: C 64.81; H 4.20.

 $AgBrP^3$ (15). Colorless crystalline material (67.0 mg, 82.0%). $^{31}\text{P}\{^{1}\text{H}\}$ NMR (CD₂Cl₂, 298, δ): –9.2 (br dd, J_{PP} 210 Hz, J_{PAg} 280 Hz, 2P, PPh₂), -28.14 (br td, $J_{\rm PP}$ 210 Hz, $J_{\rm PAg}$ 1180 Hz, 1P, PPh). ¹H NMR (CD₂Cl₂, 298, δ): 7.66 (m, 4H), 7.22–7.36 (m, 17H), 7.18 (dd, J_{HH} ca. 7.5, 4H), 7.07 (m, 4H), 6.96 (m, 4H). Anal. calc. for AgC₄₂H₃₃P₃Br (%): C 61.63; H 4.06. Found: C 61.33; H 4.08.

 $AgIP^3$ (16). Colorless crystalline material (73.7 mg, 85.0%). $^{31}P\{^{1}H\}$ NMR (CD₂Cl₂, 298, δ): AB₂ spin system -11.0 $(J_{PP} 207 \text{ Hz}, J_{PAg} 285 \text{ and } 247 \text{ Hz}, 2P, PPh_2), -29.2 (J_{PP} 207 \text{ Hz},$ $J_{\rm PAg}$ 125 and 109 Hz, 1P, PPh). ¹H NMR (CD₂Cl₂, 298, δ): 7.64 (m, 4H), 7.21–7.34 (m, 8H), 7.19 (dd, J_{HH} 6.9 and 7.5 Hz, 4H), 7.10 (dd, J_{HH} 7.8 and 8.2 Hz, 2H), 7.04 (m, 2H), 6.99 (m, 4H). Anal. calc. for AgC₄₂H₃₃P₃I (%): C 58.29; H 3.84. Found: C 57.91; H 3.96.

X-ray structure determination

The crystals of 1-12, 15, and 16 were immersed in cryo-oil, mounted in a Nylon loop, and measured at a temperature of 120 K. The X-ray diffraction data were collected with Bruker SMART APEX II or Bruker Kappa Apex II Duo diffractometers using MoK α radiation ($\lambda = 0.71073$ Å). The APEX2⁵³ program package was used for cell refinements and data reductions. The structures were solved by direct methods using the SHELXS-2013/2014⁵⁴ programs with the WinGX⁵⁵ graphical user interface. A semiempirical absorption correction (SADABS)⁵⁶ was applied to all data. Structural refinements were carried out using SHELXL-2013/2014.54

The PF₆⁻ counterion in 1 was disordered over two positions and was refined with occupancies 0.78/0.22. The displacement parameters of the fluorine atoms in both components were constrained to be equal and were restrained, so that their U_{ii} components approximate isotropic behaviour.

The contribution of the missing solvent to the calculated structure factors in complexes 1, 3, 4, and 8 was taken into

account by using a SQUEEZE routine of PLATON.57 The missing solvent was not included in the cell content.

The crystal of 8 was of low quality due to the partial loss and disorder of crystallization solvent molecules. Therefore, high-quality refinement could not be achieved and only the structural data of 8 are presented in the ESI†.

All H atoms in 1-12, 15, and 16 were positioned geometrically and constrained to ride on their parent atoms, with C-H = 0.95-0.99 Å and $U_{\rm iso}$ = 1.2-1.5 $U_{\rm eq}$. (parent atom). The crystallographic details are summarized in Tables S1-S3.† CCDC 1484959-1484962, 1484966-1484970, 1484973, 1484974, 1484978 (1-12), 1484979 (15), and 1484980 (16) contain the supplementary crystallographic data for this paper.

Photophysical measurements

The steady-state emission and excitation spectra of complexes **1–16** in the solid state at room temperature and at 77 K were recorded on a FluoroMax 4 and a FluoroLog 3 Horiba spectrofluorimeters. The xenon lamps (300 W and 450 W) were applied as light sources to excite luminescence. Two LEDs (maxima of emission at 340 and 390 nm) were used in pulse mode to pump luminescence in lifetime measurements at room temperature (pulse width 1.2 nm, repetition rate 100 Hz to 10 kHz). A pulse laser, DTL-399QT "Laser-Export Co. Ltd" (maximum of emission at 351 nm, 50 mW, pulse width 6 ns, repetition rate 1 kHz); a digital oscilloscope, Tektronix DPO3034 (bandwidth 300 MHz); a MUM monochromator (LOMO, interval of wavelengths 10 nm); and a photomultiplier tube from Hamamatsu were used for lifetime measurements at 77 K. The absolute emission quantum yield of the powders was determined using a FluoroLog 3 Horiba spectrofluorimeter and a Quanta-phi integration sphere.

Computational details

All models were fully optimized with the Gaussian 09 program package⁵⁸ at the DFT level of theory. The hybrid density functional PBE0⁵⁹ was utilized together with the basis set consisting of the def2-TZVPPD⁶⁰ effective core potential basis set with the triple-zeta-valence basis set with two sets of polarization and diffuse basis functions for Cu, Ag, and Au atoms, the standard all-electron basis set 6-31G(d,p) for C and H atoms, and a slightly more flexible basis set 6-311+G(d) for the interacting P and O atoms. Halogens were described with the def2-TZVPPD basis, which for iodine includes also ECP for the core electrons. To study the intramolecular interactions of the complexes in detail, we performed topological charge density analysis with the QTAIM (Quantum Theory of Atoms in Molecules)⁶¹ method, which allowed us to access the nature of the bonding via calculating different properties of the electron density at the bond critical points. The analysis was performed with the AIMAll program⁶² using the wave functions obtained from the DFT calculations.

Emission properties were studied by optimizing the corresponding models in the triplet state, and studying the changes in the appearance of the frontier molecular orbitals. Emission wavelengths were estimated by the total energy difference of the molecules in T_1 and S_0 electronic states, which seriously underestimates the emission wavelength in the bimetallic complexes 1–4, but is able to reproduce reasonable values for monomer complexes.

Single molecules were used as models for structures 1–4, 7, 10, 13 and 16. The geometries of all models were fully optimized in the singlet and triplet electronic states. The counteranions were not included for the cationic bimetallic models 1–4. The optimized coordinates of all calculated compounds are included in the ESI.† Generally, the optimized geometries were very well in line with the experimental crystal structures, except for the silver dimer 2, which was somewhat symmetrized in the computational optimization, and therefore, all computational properties were also calculated for the non-optimized experimental structure.

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