Cite this: Chem. Sci., 2022, 13, 6830
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Received 26th February 2022
Accepted 2nd May 2022
DOI: 10.1039/d2sc01200f
rsc.li/chemical-science

## Introduction

Transition metal complexes in which an arsolyl ligand ' $\mathrm{AsC}_{4} \mathrm{R}_{4}$ ' acts as a pentahapto cyclopentadienyl mimic are limited to $\left[\mathrm{M}(\mathrm{CO})_{3}\left(\eta^{5}-\mathrm{AsC}_{4} \mathrm{Ph}_{4}\right)\right](\mathrm{M}=\mathrm{Mn}, \mathrm{Re})$ and a small number of structurally characterised ferrocene analogues $\left[\mathrm{Fe}\left(\eta^{5}-\mathrm{AsC}_{4} \mathrm{Me}_{n}{ }^{-}\right.\right.$ $\left.\left.\mathrm{H}_{4-n}\right)_{2}\right](n=0,2,4),{ }^{1}$ all of which involve $d^{6}$-pseudo-octahedral metal centres. This is in contrast to the chemistry of $\eta^{5}$-phospholyls which is richly diverse and well-charted across the entire periodic system. ${ }^{2}$ Of late, the question of intermetallic polarcovalent (dative) bonding has received considerable attention, ${ }^{3}$ but finds its origin in the early observation that $d^{8}$ -$\left[\mathrm{Co}(\mathrm{CO})_{2}\left(\eta-\mathrm{C}_{5} \mathrm{H}_{5}\right)\right]$ forms a Lewis base/acid adduct with $\mathrm{HgCl}_{2} .{ }^{4}$ While championing the concept of "Metal Bases par excellence" ${ }^{5}$ Werner placed particular emphasis on group 9 complexes of the form $\left[\mathrm{ML}_{2}\left(\eta-\mathrm{C}_{5} \mathrm{H}_{5}\right)\right]$, recalling the archetypal Lewis basicity of $\left[\mathrm{Co}(\mathrm{CO})_{2}\left(\eta-\mathrm{C}_{5} \mathrm{H}_{5}\right)\right]$. Given that $\eta^{5}$-phospholyls may on occasion display $P$-centred nucleophilicity, ${ }^{6}$ and that Mathey has established the viability of late transition metal $\eta^{5}$-phospholyls, e.g., $\left[\mathrm{Co}(\mathrm{CO})_{2}\left(\eta^{5}-\mathrm{PC}_{4} \mathrm{Ph}_{2} \mathrm{H}_{2}\right)\right]$, , we have considered whether currently unknown arsolyl complexes of late transition metals with higher d-occupancies might also be viable. Specifically, we were intrigued to explore whether these might also serve as Lewis bases towards other metal centres and to what extent the arsenic donor, being typically less nucleophilic than in phosphorus congeners, might augment, support, or competitively compromise resulting metal-metal bonding. Accordingly, we report herein the isolation of the first late transition metal $\eta^{5}$ arsolyl complexes and demonstrate their proclivity towards bridge-assisted metal-metal bond formation.

[^0]Heating $\left[\mathrm{Co}_{2}(\mathrm{CO})_{8}\right]$ and a selection of $A s$-phenyl arsoles (1ae, Scheme 1) in refluent THF or $n$-hexane provides the highly airsensitive arsolyl complexes $\left[\mathrm{Co}(\mathrm{CO})_{2}\left(\eta^{5}-\mathrm{AsC}_{4} \mathrm{R}_{4}\right)\right]$ (2a-e) in modest yields following strictly anaerobic chromatography. Efforts to increase the isolated yields of 2a-e with extended reaction times or increased reaction temperatures were unsuccessful, however slightly increased yields were obtained by instead employing the more reactive As-chloro arsoles (see $\mathrm{ESI} \dagger$ ). The precise fate of the As-substituent during these reactions was not evident from the significant quantities of intractable materials also produced. The cleavage of an As-Ph bond from an arsole has on one previous occasion been observed in the reaction of $\left[\mathrm{Mn}_{2}(\mathrm{CO})_{10}\right.$ ] with $\mathrm{PhAsC}_{4} \mathrm{Me}_{2} \mathrm{H}_{2}$, albeit under rather more forcing conditions $\left(150{ }^{\circ} \mathrm{C}, 4\right.$ hours). ${ }^{1 c}$ Sub-optimal yields notwithstanding, the syntheses of 2a-e were highly reproducible, affording complexes $2 \mathbf{a}$ and $\mathbf{2 b}$ as orange solids or 2c-e as dark orange-red liquids at ambient temperature; the latter group underwent substitution with trimethylphosphite in toluene at $100{ }^{\circ} \mathrm{C}$ to provide the bright orange crystalline complexes $\left[\mathrm{Co}(\mathrm{CO})\left\{\mathrm{P}(\mathrm{OMe})_{3}\right\}\left(\eta^{5}-\mathrm{AsC}_{4} \mathrm{R}_{4}\right)\right](\mathbf{2 f - h})$ in high yield.

Selected spectroscopic data for 2a-e and other germane cobalt $(\mathrm{I})$ dicarbonyl complexes are presented in Table 1. We note that $\nu_{\mathrm{CO}}$ frequencies for 2a-e fall between those for $\left[\mathrm{Co}(\mathrm{CO})_{2}\left(\eta^{5}-\mathrm{C}_{5} \mathrm{H}_{5}\right)\right]$ and $\left[\mathrm{Co}(\mathrm{CO})_{2}\left(\eta^{5}-\mathrm{C}_{5} \mathrm{Me}_{5}\right)\right]$, being comparable to those for $\eta^{5}$-phospholyl cobalt dicarbonyl complexes


Scheme 1 Synthesis of $\eta^{5}$-arsolyl complexes of cobalt. (i) $\left[\mathrm{CO}_{2}(\mathrm{CO})_{8}\right]$, $\Delta$, THF or $n$-hexane. (ii) $\mathrm{P}(\mathrm{OMe})_{3}, 100^{\circ} \mathrm{C}$, toluene.

Table 1 Selected spectroscopic data for complexes prepared in this work, and some previously reported complexes for comparison

| Entry | Complex | ${ }^{13} \mathrm{C} \delta_{\mathrm{C}-\alpha}{ }^{a}$ | ${ }^{13} \mathrm{C} \delta_{\mathrm{C}-\beta}{ }^{a}$ | ${ }^{13} \mathrm{C} \delta_{\mathrm{CO}}{ }^{a}$ | $\nu_{\mathrm{CO}}{ }^{\text {b }}(\mathrm{sym})$ | $\nu_{\mathrm{CO}}{ }^{b}($ asym $)$ | $k_{\mathrm{CO}}{ }^{c}$ |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| 2 a | $\mathrm{Co}(\mathrm{CO})_{2}\left(\eta^{5}-\mathrm{AsC}_{4} \mathrm{Ph}_{4}\right)$ | 137.2 | 135.1 | 203.8 | 2026 | 1978, $1970{ }^{\text {d }}$ | 16.13 |
| 2b | $\mathrm{Co}(\mathrm{CO})_{2}\left(\eta^{5}-\mathrm{AsC}_{4} \mathrm{Ph}_{2} \mathrm{H}_{2}\right)$ | 138.1 | 94.9 | 203.0 | 2032 | 1985, 1976 ${ }^{\text {d }}$ | 16.23 |
| 2c | $\mathrm{Co}(\mathrm{CO})_{2}\left(\eta^{5}-\mathrm{AsC}_{4} \mathrm{Me}_{4}\right)$ | $118.9^{e}$ | $113.1{ }^{e}$ | $205.1^{e}$ | 2019 | 1967 | 16.02 |
| 2d | $\mathrm{Co}(\mathrm{CO})_{2}\left(\eta^{5}-\mathrm{AsC}_{4} \mathrm{Et}_{4}\right)$ | $128.4{ }^{e}$ | $118.8{ }^{e}$ | $205.3{ }^{e}$ | 2017 | 1965 | 15.99 |
| 2 e | $\mathrm{Co}(\mathrm{CO})_{2}\left\{\eta^{5}-\mathrm{AsC}_{4}\left(\mathrm{SiMe}_{3}\right)_{2} \mathrm{Me}_{2}\right\}$ | $122.2{ }^{e}$ | $119.8{ }^{e}$ | $204.5{ }^{e}$ | 2015 | 1962 | 15.95 |
| 2 f | $\mathrm{Co}(\mathrm{CO})\left\{\mathrm{P}(\mathrm{OMe})_{3}\right\}\left(\eta^{5}-\mathrm{AsC}_{4} \mathrm{Me}_{4}\right)$ | 114.3 | 109.8 | 206.8 | 1947 |  | 15.39 |
| 2 g | $\mathrm{Co}(\mathrm{CO})\left\{\mathrm{P}(\mathrm{OMe})_{3}\right\}\left(\eta^{5}-\mathrm{AsC}_{4} \mathrm{Et}_{4}\right)$ | 124.4 | 115.3 | 207.3 | 1945 |  | 15.36 |
| 2 h | $\mathrm{Co}(\mathrm{CO})\left\{\mathrm{P}(\mathrm{OMe})_{3}\right\}\left\{\eta^{5}-\mathrm{AsC}_{4}\left(\mathrm{SiMe}_{3}\right)_{2} \mathrm{Me}_{2}\right\}$ | 118.7 | 116.1 | 205.5 | 1941 |  | 15.30 |
|  | $\mathrm{Co}(\mathrm{CO})_{2}\left(\eta^{5}-\mathrm{C}_{5} \mathrm{H}_{5}\right)^{11}$ | $84.5{ }^{e}$ |  | $205.6{ }^{e}$ | $2033{ }^{f}$ | $1972{ }^{f}$ | 16.17 |
|  | $\mathrm{Co}(\mathrm{CO})_{2}\left(\eta^{5}-\mathrm{C}_{5} \mathrm{Me}_{5}\right)^{11}$ | $96.7^{e}$ |  | $207.9^{e}$ | 2011 | 1949 | 15.81 |
|  | $\mathrm{Co}(\mathrm{CO})_{2}\left(\eta^{5}-\mathrm{C}_{5} \mathrm{Ph}_{5}\right)^{8}$ | No data |  |  | 2000 | 1945 | 15.69 |
|  | $\mathrm{Co}(\mathrm{CO})_{2}\left(\eta^{5}-\mathrm{C}_{5}\left\{\mathrm{CH}_{2} \mathrm{Ph}\right\}_{5}\right)^{8}$ | No data |  |  | 2020 | 1960 | 15.97 |
|  | $\mathrm{Co}(\mathrm{CO})_{2}\left(\eta^{5}-\mathrm{PC}_{4} \mathrm{Ph}_{2} \mathrm{H}_{2}\right)^{9}$ | No data |  |  | 2030 | 1980 | 16.21 |
|  | $\mathrm{Co}(\mathrm{CO})_{2}\left(\eta^{5}-\mathrm{PC}_{4}{ }^{t} \mathrm{Bu}_{2} \mathrm{H}_{2}\right)^{10}$ | 136.2 | $91.9$ | $204.0$ | 2023 | 1968 | 16.06 |
|  | $\mathrm{Co}(\mathrm{CO})\left\{\mathrm{P}(\mathrm{OMe})_{3}\right\}\left(\eta^{5}-\mathrm{PC}_{4}{ }^{t} \mathrm{Bu}_{2} \mathrm{H}_{2}\right)^{10}$ | 134.7 | 90.5 | 208.0 | No data reported |  | - |

${ }^{a} \mathrm{C}_{6} \mathrm{D}_{6}$ solution unless otherwise stated, ppm downfield from $\mathrm{SiMe}_{4}, 25{ }^{\circ} \mathrm{C}$; the labels $\alpha$ and $\beta$ refer to ring-carbon positions with respect to the heteroatom (where applicable). ${ }^{b} n$-Hexane solution unless otherwise stated, $\mathrm{cm}^{-1}, 25{ }^{\circ} \mathrm{C}$. ${ }^{c}$ Cotton-Kraihanzel force constant in N cm ${ }^{-1}$. ${ }^{d}$ Resolution of the doubly degenerate E vibrational mode is observed in $n$-hexane for these complexes $c f$., $\nu_{\mathrm{CO}}\left(\mathrm{CH}_{2} \mathrm{Cl}_{2}\right): \mathbf{2 a} 2022,1963 \mathrm{~cm}{ }^{-1} ; \mathbf{2 b}$ 2027, $1972 \mathrm{~cm}^{-1} .{ }^{e} \mathrm{CDCl}_{3}$ solution, $25{ }^{\circ} \mathrm{C} .{ }^{f}$ Cyclohexane solution.
reported by Mathey. ${ }^{7}$ The ${ }^{13} \mathrm{C}\left\{{ }^{1} \mathrm{H}\right\}$ NMR shifts for the CO ligands in 2a-e mirror the trends in $\nu_{\mathrm{CO}}$ frequencies with a gradual shift to higher frequency and are not dissimilar to those for $\left[\mathrm{Co}(\mathrm{CO})_{2}\left(\eta^{5}-\mathrm{C}_{5} \mathrm{R}_{5}\right)\right](\mathrm{R}=\mathrm{H}, 204.7 ; \mathrm{R}=\mathrm{Me}, 207.9 \mathrm{ppm})$ and $\left[\mathrm{Co}(\mathrm{CO})_{2}\left(\eta^{5}-\mathrm{PC}_{4}{ }^{t} \mathrm{Bu}_{2} \mathrm{H}_{2}\right)\right](204.0 \mathrm{ppm}) .{ }^{7}$ Comparison of ${ }^{13} \mathrm{C}\left\{{ }^{1} \mathrm{H}\right\}$ NMR data of the ring-carbon nuclei to those of the corresponding free arsoles 1a-e (ESI $\dagger$ ) reveals a shift to low frequency of $20-30 \mathrm{ppm}$. The molecular structures of $2 \mathbf{2 a}, 2 \mathbf{b}, 2 \mathbf{f}$, and $2 \mathbf{h}$


Fig. 1 Molecular structures of (a) 2a and (b) $2 f$ ( $50 \%$ displacement ellipsoids, arsenolyl ring and phosphite substituents simplified). Selected distances [ $\AA$ ] and angles [ ${ }^{\circ}$ ]: 2 a As1-Co1 2.427(4), As1-C3 $1.900(2)$, C3-C4 1.443(3), C4-C5 1.427(3), C5-C6 1.439(3), C6-As1 1.915(2), Co1-C1 1.746(3), Co1-C2 1.747(3), C3-As1-C6 84.52(8), As1-C3-C4 114.32(1), C3-C4-C5 113.65(2), C4-C5-C6 112.84(2), C5-C6-As1 114.38(1), C1-Co1-C2 90.82(1); 2f As1-Co1 2.408(5), As1-C5 1.910(3), C5-C6 1.430(4), C6-C7 1.423(4), C7-C8 1.427(5), C8-As1 1.903(3), Co1-C1 1.719(3), Co1-P1 2.097(9), C5-As1-C8 84.12(1), As1-C5-C6 114.25(2), C5-C6-C7 113.30(3), C6-C7-C8 113.47(3), C7-C8-As1 114.56(2), C1-Co1-P1 93.49(1). (c) Intersection of C5-C6-C7-C8 (blue) and P1-Co-C1-C2 (red) planes for 2 f at $89.7^{\circ}$ with As, C5, C6, C7, C8 centroid in green.
were crystallographically determined, with two representative examples, $\mathbf{2 a}$ and $2 \mathbf{f}$, depicted in Fig. 1 (for $\mathbf{2 b}$ and 2 h see $\mathrm{ESI} \dagger$ ).

The structural models for $2 \mathbf{2 a}, 2 \mathbf{b}, \mathbf{2 f}$ and 2 h all confirm the targeted $\eta^{5}$-arsolyl coordination. In $2 \mathbf{a}$ and $2 f$ the ligands are almost symmetrically disposed with respect to the vertical plane which bisects the $\eta^{5}$-arsolyl ring, whereas for $2 \mathbf{b}$ and $2 \mathbf{h}$ these are rotated to a position slightly offset from the arsenic-ring centroid vector (ESI $\dagger$ ). Consistent with the difference in the covalent radii of carbon $(0.76 \AA)$ and arsenic $(1.19 \AA)$, the latter is in each case very slightly displaced $\left(3-5^{\circ}\right)$ from the mean plane defined by the heterocycle carbon atoms, though less than found in the free arsoles (1a: $\left.10.2^{\circ} ; \mathbf{1 b}: \mathbf{7 . 1 0}{ }^{\circ}\right) .^{\mathbf{1 0}}$ The geometry of the metal and $\eta^{5}$-arsolyl rings in $2 \mathbf{a}, \mathbf{2 b}, \mathbf{2 f}$ and $2 \mathbf{h}$ are of a distorted pentagonal pyramid. The $\mathrm{C}_{\alpha}-\mathrm{As}-\mathrm{C}_{\alpha}$ angles at the arsenic vertices are significantly contracted $\left(84-87^{\circ}\right)$ from that of an idealised pentagon $\left(108^{\circ}\right)$ while the remaining $\mathrm{C}-\mathrm{C}-\mathrm{C}$ angles are in the range $111-114^{\circ}$, and comparable to those found in $\left[\mathrm{Co}(\mathrm{CO})_{2}\left(\eta^{5}-\mathrm{C}_{5} \mathrm{R}_{5}\right)\right]\left(\mathrm{R}=\mathrm{Me}, \mathrm{Ph}, \mathrm{CH}_{2} \mathrm{Ph}\right) .{ }^{8}$

The results of computational interrogation of the model 'parent' compound $\left[\mathrm{Co}(\mathrm{CO})_{2}\left(\eta^{5}-\mathrm{AsC}_{4} \mathrm{H}_{4}\right)\right]\left(2_{\mathrm{As}}^{\prime}\right)(\mathrm{DFT}: \omega \mathrm{B} 97 \mathrm{X}-\mathrm{D} / 6-$ 31G*/LANL2D $\zeta$; ESI $\dagger$ ) are summarised in Fig. 2 alongside those for $\left[\mathrm{Co}(\mathrm{CO})_{2}\left(\eta^{5}-\mathrm{C}_{5} \mathrm{H}_{5}\right)\right]\left(2_{\mathbf{C H}}^{\prime}\right)$, and the hypothetical pnictogenyl analogues $\left[\mathrm{Co}(\mathrm{CO})_{2}\left(\eta^{5}-\mathrm{AC}_{4} \mathrm{H}_{4}\right)\right]\left(\mathrm{A}=\mathrm{N} 2_{\mathbf{N}}^{\prime}, \quad \mathrm{P} 2_{\mathbf{p}}^{\prime}, \mathrm{Sb} 2_{\mathbf{S b}}^{\prime}\right)$. The HOMO-1 is in all cases substantially derived from the Co $-\mathrm{d}_{z^{2}}$ orbital and readily corroborates the known nucleophilic behaviour of $2_{\mathbf{C H}}^{\prime}$. For $2_{\mathbf{C H}}^{\prime}$ this is, however, effectively the only orbital that is geometrically disposed to allow the complex to function as a Lewis base since the HOMO is involved with cyclopentadienyl binding. This is also the case with the HOMO of the pnictogenolyl examples however the orbital substantially protrudes radially from the ring. The HOMO-1 involves substantial contribution from the pnictogen orbital such that both the HOMO and HOMO-1 (and also HOMO-2) contribute to a prominent region of electron density localised over these atoms which is reflected in the electrostatic potential map for $2_{\text {As }}^{\prime}$ and


Fig. 2 Frontier orbitals of interest for $\left[\mathrm{Co}(\mathrm{CO})_{2}\left(\eta^{5}-\mathrm{AC}_{4} \mathrm{H}_{4}\right)\right]\left(2_{\mathrm{A}}^{\prime}: \mathrm{A}=\right.$ CH, N, P, As, Sb; DFT: $\omega$ B97X-D/6-31G*/LANL2D $\zeta$ ). Inset $=$ electrostatic potential map for $\left[\mathrm{Co}(\mathrm{CO})_{2}\left(\eta^{5}-\mathrm{AsC}_{4} \mathrm{H}_{4}\right)\right] 2_{\text {As }}^{\prime}$ indicating site for charge-controlled electrophilic attack. Condensed Fukui functions (natural populations) for electrophilic attack: $f^{-}(\mathrm{Co})=0.113, f^{-}(\mathrm{As})=$ 0.251 .
the condensed Fukui functions for both arsenic and cobalt (Fig. 2 inset). Furthermore, on descending group 15, the pnictogen A-p $p_{z}$ orbital increasingly contributes and this is accompanied by a monotonic increase in the energy the HOMO, HOMO-1 and HOMO-2 orbitals which should manifest as an increase in the basicity of not only the metal but also the pnictogen. This is intriguingly counterintuitive in that the basicity, nucleophilicity and strength of pnictogen coordination generally decreases for simple pnictanes $\mathrm{AR}_{3}$ traversing from P to $\mathrm{Sb} .{ }^{12}$ Compared to phospholyl and arsolyl complexes, $\eta^{5}$-stibolyl complexes are rarer still, being limited to three ferrocene analogues, but clearly worthy of further study, not least because of the onset of secondary bonding for the heavier pnictogens. ${ }^{13}$

To explore the possibility of metal-metal bond-formation, the representative 2 c was chosen, commencing with mercuric chloride by analogy with the prototypical and monomeric adduct $\left[\mathrm{Co}\left(\mathrm{HgCl}_{2}\right)(\mathrm{CO})_{2}\left(\eta-\mathrm{C}_{5} \mathrm{H}_{5}\right)\right] .^{4}$ The reaction of 2c with $\mathrm{HgCl}_{2}$ in acetone rapidly results in precipitation of the poorly soluble yellow dimer $[2 \mathbf{c} \cdot \mathrm{HgCl}(\mu-\mathrm{Cl})]_{2}(3)$ in high yield (Scheme 2 and Fig. 3).

The dimeric formulation follows from HR-ESI-MS data, which are devoid of ions due to dissociated $2 \mathbf{c}$, in addition to crystallographic analyses of two isomers that differ in the $\mu: \sigma-$ $\eta^{5}$-arsolyl rings adopting mutually syn or anti positions with


Scheme 2 Bridge-assisted formation of dative bonds from cobalt to gold(I), mercury(II) and platinum(II). (i) $\mathrm{HgCl}_{2}$. (ii) cis-[Pt( $\left.\mathrm{C}_{6} \mathrm{~F}_{5}\right)_{2}($ hex $\left.)\right]$ (hex $=1,5$-hexadiene). (iii) $\left[A u\left(C_{6} F_{5}\right)(T H T)\right]$ ( $\mathrm{THT}=$ tetrahydrothiophene).
respect to the rhomboidal $\mathrm{Hg}_{2}(\mu-\mathrm{Cl})_{2}$ core. Thus, yellow needles of anti-3 (major) and orange prisms of syn-3 (minor) slowly crystallise together from solutions of 3 in acetone stored at $-30^{\circ} \mathrm{C}$.

One half of each of the dimeric structures of both anti-3 and syn-3 in the solid state is crystallographically unique due to the centre of the $\mathrm{Hg}_{2}(\mu-\mathrm{Cl})_{2}$ unit coinciding with either an inversion centre (anti-3 in $P 2_{1} / n$ ) or twofold rotation axis (syn-3 in C2/c). The coordination polyhedra of the $\mathrm{Hg}^{\mathrm{II}}$ atoms are strikingly different in each isomer: anti-3 features severely distorted trigonal bipyramidal mercury with the arsenic and cobalt atoms


Fig. 3 The molecular structures of (a) syn-3 and (b) anti-3 (methyl groups simplified, $50 \%$ displacement ellipsoids). Selected distances [Å] and angles [ ${ }^{\circ}$ ]: (a) syn-3 As1-Hg1 2.7268(9), Co1-Hg1 2.620(1), Hg1Cl1 2.491(2), Hg1-Cl2 2.598(2), Hg1-Cl2' 2.708(2), As1-Hg1-Co1 $55.86(3), \mathrm{Hg} 1-\mathrm{Cl} 2-\mathrm{Hg} 1^{\prime} 91.87(6), \mathrm{Cl} 2-\mathrm{Hg} 1-\mathrm{Cl} 2^{\prime} 87.62(6), \mathrm{Cl} 1-\mathrm{Hg} 1-$ Cl 2 100.71(6). (b) Anti-3 As1-Hg1 2.6334(6), Co1-Hg1 2.6702(9), Hg1Cl1 2.390(1), Hg1-Cl2 2.778(1), Hg1-Cl2' 2.777(1) As1-Hg1-Co1 $56.27(2), \mathrm{Hg} 1-\mathrm{Cl} 2-\mathrm{Hg}^{\prime}$ 84.37(4), $\mathrm{Cl} 2-\mathrm{Hg} 1-\mathrm{Cl} 2^{\prime} 95.63(4), \mathrm{Cl} 1-\mathrm{Hg} 1-$ $\mathrm{Cl} 2,100.59(5) . i=$ crystallographic inversion centre.
assuming pseudo-axial and -equatorial positions, respectively ( $\tau_{5}=0.89$ ), whereas for $s y n-3$ the more sterically congested mercury geometry more closely approaches a square-based pyramid ( $\tau_{5}=0.62$ ) with the non-bridging chlorides occupying the eclipsing apices. The As- Hg bond distances are somewhat shorter by $c a .0 .1 \AA$ in anti-3 (2.633(6) $\AA$ ) than syn-3 $(2.727(9) \AA$ ), while the Co-Hg bond distances of $2.670(9) \AA$ (anti) and $2.620(1) \AA(s y n)$ are essentially equivalent within crystallographic precision limits. The latter pair are somewhat longer than the sum of covalent radii for the individual elements (2.58 $\AA$ ) and the separation $(2.578(4) \AA)$ observed for $\left[\mathrm{Co}\left(\mathrm{HgCl}_{2}\right)(-\right.$ $\left.\mathrm{CO})_{2}\left(\eta^{5}-\mathrm{C}_{5} \mathrm{H}_{5}\right)\right]$, ${ }^{4}$ by virtue of the increased coordination number at mercury. The slight elongation of the $\mathrm{Co} \rightarrow \mathrm{Hg}$ interaction here is almost certainly a geometric compromise to accommodate the $\mathrm{Hg}^{\text {II }}$ centre within the disparate coordination spheres of the $\mathrm{As}^{\mathrm{III}}$ and $\mathrm{Co}^{\mathrm{I}}$ donors, rather than indicating any noteworthy electronic phenomena beyond non-directional spodium bonding. ${ }^{14}$ From a valence-bond perspective, 2c may be considered to serve as a neutral, 4-electron bidentate ligand with a somewhat narrow bite-angle ( $c a .55-56^{\circ}$ ). Solution infrared data for each isomer (after manual separation of crystals) resulted in spectra identical to that of the bulk sample of 3 obtained above, i.e., it appears that upon dissolution in acetonitrile, syn-3 isomerises to anti-3 and the latter conformation is the natural condition of the adduct. Given both isomers have identical QSAR volumes $\left(536 \AA^{3}\right)$, the polarity of $\operatorname{syn}-3$ (dipole $=$ 23.5 D cf. 0 for anti-3) possibly plays a role in its crystal formation.

Though their Lewis acidity is well-documented, neither $\left[\mathrm{Hg}\left(\mathrm{CF}_{3}\right)_{2}\right]$ nor $\left[\mathrm{Hg}\left(\mathrm{C}_{6} \mathrm{~F}_{5}\right)_{2}\right]$ provided any evidence of detectable adduct formation with $2 \mathbf{c}$. Coordination of $2 \mathbf{c}$ to divalent platinum(II) could however be demonstrated in its reaction with cis$\left[\mathrm{Pt}\left(\mathrm{C}_{6} \mathrm{~F}_{5}\right)_{2}(\right.$ hex $\left.)\right]$ (hex $=\eta^{2}: \eta^{2}$-1,5-hexadiene) in $\mathrm{CH}_{2} \mathrm{Cl}_{2}$ to provide after anaerobic chromatography a single isolable


Fig. 4 The molecular structure of 4 (methyl and pentafluorophenyl groups simplified, $50 \%$ displacement ellipsoids, Pt TBPY-5-12-C enantiomer in non-centrosymmetric $P 2_{1} 2_{1} 2_{1}$ space group). Selected distances [Å] and angles [0]: As1-Pt1 2.5185(8), Co1-Pt1 2.996(1), Pt1C12 2.053(8), Pt-C18 2.077(7), As1-Pt1-Co1 53.04(3), As1-Pt1-C18 169.58(2), Co1-Pt1-C18 137.24(2).
orange compound in modest yield. The complex with three $\nu_{\mathrm{CO}}$ absorptions at 2084, 2053 and $2014 \mathrm{~cm}^{-1}$ and the observation of six resonances in the ${ }^{19} \mathrm{~F}$ NMR (ESI $\dagger$ ) confirmed the presence of two chemically inequivalent $\mathrm{C}_{6} \mathrm{~F}_{5}$ groups inconsistent with a simple ' $2 \mathbf{c} \cdot \operatorname{Pt}\left(\mathrm{C}_{6} \mathrm{~F}_{5}\right)_{2}$ ' adduct. The X-ray diffraction analysis (Fig. 4) confirmed it to be $\left[2 \mathrm{c} \cdot\right.$ cis- $\left.\operatorname{Pt}(\mathrm{CO})\left(\mathrm{C}_{6} \mathrm{~F}_{5}\right)_{2}\right]$ (4) arising from CO sequestration.

The structure of 4, like anti-3 and syn-3, comprises 2c acting as a bidentate ligand (chelate bite $=c a .53^{\circ}$ ) to a five-coordinate ${ }^{\prime} \mathrm{Pt}(\mathrm{CO})\left(\mathrm{C}_{6} \mathrm{~F}_{5}\right)_{2}$ ' centre. Putting aside the $\mathrm{Co} \rightarrow \mathrm{Pt}$ interaction, the platinum adopts a square-planar geometry (inset, Fig. 4). The $\operatorname{Pt}(\mathrm{II})$ centre is slightly displaced from the ligand donor mean plane (trans angles $169.58(2)^{\circ}, 174.58(3)^{\circ}$ ) towards the dative $\mathrm{Co} \rightarrow \mathrm{Pt}$ interaction which is remarkably long (Co1-Pt1: $2.966(1) \AA)$, falling between the sums of covalent $(2.62 \AA)$ and van der Waals (ca. $4.01 \AA$ ) radii for these elements. The As1-Pt1 bond length of $2.519(8) \AA$ is also significantly elongated relative to more conventional platinum-bound arsines e.g., $\left[\mathrm{PtCl}_{x}\left(\mathrm{C}_{6}-\right.\right.$ $\left.\left.\mathrm{F}_{5}\right)_{2-x}(\mathrm{~L})_{n}\right]\left(n=1,2 ; x=0,1 ; \mathrm{L}=\mathrm{CH}_{2}\left(\mathrm{AsPh}_{2}\right)_{2}, \mathrm{C}_{2} \mathrm{H}_{4}\left(\mathrm{AsPh}_{2}\right)_{2}:\right.$ 2.34-2.43 $\AA$ ). ${ }^{15}$ Despite being apparently rather weak, it is this interaction which completes the square coordination plane.

Further exemplifying the bidentate nature of 2c, combination with one equivalent of $\left[\mathrm{Au}\left(\mathrm{C}_{6} \mathrm{~F}_{5}\right)(\mathrm{THT})\right]$ (THT $=$ tetrahydrothiophene) leads to the isolation of a yellow solid of deceptively simple composition " $2 \mathbf{c} \cdot \mathrm{Au}\left(\mathrm{C}_{6} \mathrm{~F}_{5}\right)$ " that is actually the salt $\left[(2 \mathrm{c})_{2} \mathrm{Au}\right]\left[\mathrm{Au}\left(\mathrm{C}_{6} \mathrm{~F}_{5}\right)_{2}\right]$ (5, Fig. 5). The crystal structure of 5 reveals bidentate coordination of two 2 c units to a $\mathrm{Au}^{+}$cation with near to coplanar coordination of the Co and As donors ( Au sits $0.22 \AA$ above the $\mathrm{Co}_{2} \mathrm{As}_{2}$ mean plane). The 2c 'ligands' are


Fig. 5 The molecular structure of 5 (methyl and pentafluorophenyl groups simplified and hydrogen atoms omitted for clarity, four crystallographically independent molecules in the unit cell, 50\% displacement ellipsoids). Selected distances [Å]: As1-Au1 2.5215(6), As2-Au1 2.5205(6), As1-Au2 3.1660(6), Co1-Au1 2.6793(9), Co2-Au1 2.6731(8). Selected angles [deg]: As1-Au1-Co1 57.89(2), As2-Au1Co2 57.67(2), As1-Au1-As2 178.99(2), Co1-Au1-Co2 167.70(3). (b) View orthogonal to the $b$-axis showing extended chain of $(\cdots A u \cdots A s)_{\infty}$ interactions (3.0746(6)-3.2122(6) Å).
asymmetrically disposed about the $\mathrm{Au}^{+}$cation, being transposed and offset by $c a .22^{\circ}$ when viewed along the As1-Au1-As2 vector. The As-Au distances are equivalent (mean $2.525 \AA$ ), with slightly longer Co-Au distances (mean $2.650 \AA$ ). The unit cell of 5 contains four crystallographically independent [(2c) $\left.)_{2} \cdot \mathrm{Au}\right]$ $\left[\mathrm{Au}\left(\mathrm{C}_{6} \mathrm{~F}_{5}\right)_{2}\right]$ pairs (ESI $\dagger$ ) with extended packing of loosely parallel columns of alternating $\left[(2 \mathbf{c})_{2} \cdot \mathrm{Au}\right]^{+}$cations and $\left[\mathrm{Au}\left(\mathrm{C}_{6} \mathrm{~F}_{5}\right)_{2}\right]^{-}$ anions. The nature of their arrangement differs slightly between pairs, with notably short $\mathrm{As} \cdots\left[\mathrm{Au}\left(\mathrm{C}_{6} \mathrm{~F}_{5}\right)_{2}\right]^{-}$interactions (between the arsolyl arsenic and $\left[\mathrm{Au}\left(\mathrm{C}_{6} \mathrm{~F}_{5}\right)_{2}\right]^{-}$gold atoms). These distances are, however, not equal, and capriciously vary between $3.0746(6) \AA$ (shortest) and 3.2122(6) $\AA$ (longest).

Computational interrogation ( $\omega$ B97X-X/6-31G*/LANL2D $\zeta$ ) of the model complexes $\left[\operatorname{CoPt}\left(\mu-\mathrm{AsC}_{4} \mathrm{H}_{4}\right)(\mathrm{CO})_{3}\left(\mathrm{CF}_{3}\right)_{2}\right]\left(4^{\prime}\right)$ and $\left[\mathrm{Co}_{2} \mathrm{Au}\left(\mu-\mathrm{AsC}_{4} \mathrm{H}_{4}\right)_{2}(\mathrm{CO})_{4}\right]^{+}\left(5^{\prime}\right)$ (detailed in the ESI $\dagger$ ) returns core geometries close to those of 4 and the cation of 5 . This analysis reveals molecular orbitals of interest (see ESI Fig. S12-S14 $\dagger$ ) that account for the geometrical features of note. For $\mathbf{4}^{\prime}$, the HOMO and HOMO-7 comprise significant overlap of the Co $-\mathrm{d}_{z^{2}}$ orbital with platinum, supporting a Co $\rightarrow$ Pt description (Löwdin bond orders As $-\mathrm{Pt} / \mathrm{Co}-\mathrm{Pt}=0.75 / 0.41$ ). For $5^{\prime}$ not only does the HOMO-16 adhere to the view of dative $\mathrm{Co} \rightarrow \mathrm{Au}^{+}$bonding (Löwdin bond orders As-Au/Co-Au = 0.78/0.56), but an orbital (HOMO-20) has a topology suggestive of the arsenolyl serving as a $\pi$-acceptor from gold, a feature also present in the MO scheme for $\mathbf{4}^{\prime}$. Furthermore, the LUMO which has substantial arsenic character would appear to account for the association of the $\left[\mathrm{Au}\left(\mathrm{C}_{6} \mathrm{~F}_{5}\right)_{2}\right]^{-}$anion at this point which underpins the extended polymeric assembly, though it is unlikely that this persists in solution.

## Conclusions

The first (eight) examples of late transition metal $\eta^{5}$-arsolyl complexes have been obtained with one example being then employed to explore the possibility of both the metal and arsenic serving as donors to $\mathrm{Pt}^{\mathrm{II}}, \mathrm{Au}^{\mathrm{I}}$ and $\mathrm{Hg}^{\mathrm{II}}$ centres. Although the individual interactions might appear weak, when both act in concert novel bridge-assisted heterometallic assemblies arise with intriguing features.

## Data availability

Crystallographic data for structurally characterised compounds have been deposited at the Cambridge Crystallographic Data Centre under CCDC 2130783-2130790 and can be obtained from https://www.ccdc.cam.ac.uk. Spectroscopic data for all new compounds are provided in the ESI accompanying this paper (https://doi.org/10.1039/d2sc01200f).

## Author contributions

RMK was responsible for the conceptualisation and execution of the experimental research, the acquisition and critical analysis of the characterisational data and compilation of the original draft. AFH was responsible for funding acquisition, project administration, validation and refinements to the manuscript.

## Conflicts of interest

There are no conflicts to declare.

## Acknowledgements

We gratefully acknowledge the Australian Research Council (DP190100723 and DP200101222) for funding.

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    $\dagger$ Electronic supplementary information (ESI) available: Synthetic, computational and experimental procedures. CCDC 2130783-2130790. For ESI and crystallographic data in CIF or other electronic format see https://doi.org/10.1039/d2sc01200f

