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Platinum has been inserted into corroles for the first time and three oxidized Pt^{IV}(corrole^{•2-})ArAr⁷ complexes have been structurally characterized. The Soret maxima of these complexes exhibit an unusually strong dependence on the meso-aryl substituents on the corrole, indicating aryl \rightarrow corrole $^{\bullet 2-}$ charge transfer character in these transitions.

As trianionic ligands with contracted $N₄$ cores, corroles sustain a great deal of unique coordination chemistry relative to porphyrins.¹ Within this area, heavy element corroles are of particular interest as optical sensors, near-IR dyes, phosphors, and organic light emitting diodes.² The size mismatch between the corrole N_4 cores and the large ionic radii of the lower oxidation states of the 5d elements, however, poses formidable challenges for metal insertion. As of today, Hf , $\mathrm{^{3}}$ W, $\mathrm{^{4}}$ Re, $\mathrm{^{5}}$ Ir, $\mathrm{^{6}}$ and Au , 7 corroles have been synthesized, whereas others such as osmium and platinum corroles are still to be reported. Here we present unambiguous evidence, including multiple singlecrystal X-ray analyses, for the formation of platinum corroles.

Insertion of Pt into corroles proved to be extraordinarily challenging. A large number of commercially available Pt precursors, each tested with a wide selection of solvents, failed to yield isolable Pt corrole derivatives. In the end, microwave irradiation of the commercially unavailable tetranuclear platinum acetate complex $[Pt(OAc)_2]_4$ 2HOAc⁸ in benzonitrile at 140–150 °C for 2 hours led to low but reproducible yields (\sim 6%) of diamagnetic Pt(IV) corroles. Notably, aerobic conditions were critical to

the success of the reaction; strict exclusion of oxygen did not result in Pt-containing products. Based on MALDI-TOF mass spectrometry and ¹H NMR spectroscopy, the products could be formulated as $Pt\{T(p-X-Ph)C\}(o/m/p-C_6H_4CN)(PhCN)$, where ${T(p-X-Ph)C}^{3-}$ is a *meso*-triarylcorrole trianion with aryl *para* substituents $X = CF_3$, H, CH₃, and OCH₃ and the axial benzonitrile-derived aryl ligand may be bound through the o-, m_z , or p-carbon, relative to the CN group (Fig. 1). Thus, C–H activation of benzonitrile resulting in a $Pt(w)$ -aryl center has been critical to the synthesis of stable platinum corroles. Electrospray ionization mass spectrometric studies indicated the N-bound benzonitrile ligand in these $Pf(w)$ complexes to be labile. Preliminary X-ray analysis of the putative $Pt\{T(p-CF_3-Ph)C\}(m-C_6H_4CN)(PhCN)$ complex indicated extensive disorder involving benzonitrilederived ligands; despite some evidence of five-coordinate Pt{T(p -X-Ph)C}($o/m/p$ -C₆H₄CN), much of the material consisted of various regioisomers of Pt{T(p -X-Ph)C}($o/m/p$ -C₆H₄CN)₂, which formally correspond to a $Pt(v)$ oxidation state. Thus, a second C-H activation, involving the N-coordinated benzonitrile, had taken place during crystallization. Air-stable, oxidized platinum complexes Pt{T(p -X-Ph)C}($o/m/p$ -C₆H₄CN)(p -C₆H₄CH₃) could be more **COMMUNICATION**
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Fig. 1 Synthesis of six-coordinate Pt^{IV} –Ar corroles and oxidized Pt^{IV} – corrole^{•2-}-ArAr' derivatives.

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reliably obtained by treating the $Pt(w)$ complexes with an aryl-Grignard reagent (Fig. 1). Because of the low yields, only the $m-C_6H_4CN$ regioisomers were fully characterized for all the corroles, whereas the p -C₆H₄CN isomer could be characterized for only one of the corroles. Fortunately, three oxidized Pt corroles yielded X-ray quality crystals, affording full structural characterization.

X-band EPR spectra were recorded for the oxidized Pt corroles in the solid state (Fig. 2), in solution $(2:1 \text{ CH}_2\text{Cl}_2)$: toluene) at RT, and in a frozen glass at 73 K obtained from the same solution. The complexes all gave strong signals with g-values \sim 2.00, consistent with the ligand radical formulation Pt(corrole $^{\bullet 2-})$ ArAr'. The spectra could be simulated with fairly narrow linewidths (Lorentzian, fwhh = 3 G) and a slight g-anisotropy, but notably without any 195 Pt hyperfine coupling.⁹ Upon dissolution in a mixture of CH_2Cl_2 and toluene at RT, the spectra changed profoundly (Fig. 3); the bandwidth became nearly twice as large, with concomitant appearance of fully resolved hyperfine coupling to 195 Pt. Freezing of the solution at 73 K had only a slight effect on the spectrum (Fig. 3). The much narrower signals in the solid state may be interpreted as an unusually well-behaved example of ''exchange narrowing''.¹⁰ This is in agreement with the crystal structures, which indicate partial corrole π -stacking with relatively short interplanar distances of about 3.4 Å. The observed g-value range of 1.997–2.011 agrees with literature values for corrole-based radical complexes.¹¹ Although well-resolved in solution, the hyperfine coupling to Pt is too small for the radical to be Pt-centered. Communication Werel complexes with an ay-1 Three of the solidar articles. Digramate on the systems of the box of the MCD-Calinette is likened to the systems of the Calinette is likened in the system of the system of the s

Fig. 2 Solid state, RT X-band EPR spectrum of Pt(TPC)(m -C₆H₄CN)(p -C₆H₄CH₃). Simulation offset for clarity. Simulation parameters: $q = 1.999$, 2.007, 2.011, value of $A(^{195}Pt)$ used = 0.8×10^{-3} cm $^{-1}$, Lorentzian bandshape, fwhh = 3.0 G.

Fig. 3 Left: RT, solution, X-band EPR spectrum of Pt(TPC)(m -C₆H₄CN)-(p -C₆H₄CH₃). Simulation offset for clarity. Simulation parameters: $g = 2.005$, $A(^{195}Pt)$ = 5.9 \times 10^{-3} cm $^{-1}$, Lorentzian bandshape, fwhh = 5.6 G. Right: frozen glass, 73 K spectrum of same solution. Simulation parameters: $q = 2.005$, $A(^{195}Pt) = 5.9 \times 10^{-3}$ cm⁻¹, Lorentzian bandshape, fwhh = 5.7 G.

Three of the oxidized Pt complexes $Pt(TPC)(m-C₆H₄CN)$ - $(p - C_6H_4CH_3)$, Pt{T(p-CH₃-Ph)C}(p-C₆H₄CN)(p-C₆H₄CH₃), and $Pt\{T(p-CH_3-Ph)C\}(m-C_6H_4CN)(p-C_6H_4CH_3)$ proved amenable to single-crystal X-ray analysis. All the structures refined well, giving R_1 [$I > 2\sigma(I)$] values ~3%. No evidence was found for counterions, positive or negative, consistent with the Pt^{IV} –corrole e^{2} – formulation mentioned above. Selected bond distances are listed in Table 1 and a representative thermal ellipsoid plot is shown in Fig. 4. As for many metallocorroles, $12,13$ the corrole macrocycles in all three complexes are almost perfectly planar. The short Pt–N distances of approximately 1.95 Å are consistent with a $Pf(v)$ oxidation state and the sterically constrained nature of the corrole N_4 core, while the axial Pt–C distances of about 2.1 Å are typical for unconstrained Pt^{IV} –C bond distances.

Fig. 5 depicts key results from DFT calculations on the model complexes Pt(corrole)(Ph)(PhCN) (C_s) and Pt(corrole)Ph₂ (C_{2v}) . The optimized structural parameters are in excellent agreement with those observed experimentally (Table 1). To a first approximation, the spin density of Pt(corrole)Ph₂ corresponds to a corrole b_1 radical (in terms of C_{2v} irreps), which resembles a porphyrin a_{2u} -type radical,¹⁴ *i.e.*, the spin density is largely localized on the three *meso* carbons and the four nitrogens and to a lesser extent on the direct C_6-C_6 linkage. The Pt does not carry a significant amount of spin density, thereby ruling out any degree of Pt(V) character. Closer examination of Fig. 5 indicates that the corrole carries only about two-thirds of the total molecular spin density; the remaining onethird of the spin density is evenly divided between the two formally anionic phenyl ipso carbons. Thus, the HOMO is not a pure corrole b_1 HOMO, but has a certain amount of aryl character as well.

Table 2 presents redox potentials for $Pt\{T(p-X-Ph)C\}(m-C_6H_4CN)$ -(PhCN), the oxidized Pt{T(p -X-Ph)C}(m -C₆H₄CN)(p -P)C} complexes

Table 1 Pertinent bond distances for Pt $\{T(p\text{-CH}_3\text{-Ph})C\}(p\text{-C}_6H_4CN)\text{-}$ $(p - C_6H_4CH_3)$, Pt(TPC)(m-C₆H₄CN)(p-C₆H₄CH₃) and Pt{T(p-CH₃-Ph)C}- $(m - C_6H_4CN)(p - C_6H_4CH_3)$

$Pt\{T(p-CH_3-Ph)C\}$ - $(p-C_6H_4CN)(p-C_6H_4CH_3)(p-C_6H_4CH_3)$		$Pt(TPC)(m-C6H4CN)$ - $Pt\{T(p-CH3-Ph)C\}$ -		$(m-C_6H_4CN)(p-C_6H_4CH_3)$	
Pt1-N1 $Pt1-N2$ $Pt1-N3$ $Pt1-N4$ $Pt1-C_{ax}(41)$ $Pt1-C_{ax}(48)$	1.948(2) 1.9698(19) 1.9693(19) 1.9470(19) 2.119(2) 2.129(2)	$Pt1-N1$ $Pt1-N2$ $Pt1-N3$ $Pt1-N4$	$1.942(3)$ Pt1-N1 $1.965(3)$ Pt1-N2 $1.970(3)$ Pt1-N3 $1.945(3)$ Pt1-N4	Pt1–C _{ax} (38) 2.119(3) Pt1–C _{ax} (41) Pt1–C _{ax} (45) 2.114(3) Pt1–C _{ax} (48)	1.951(2) 1.973(2) 1.976(2) 1.944(2) 2.128(2) 2.123(3)

Fig. 4 Thermal ellipsoid plot for Pt(TPC)($m - C_6H_4CN$)($p - C_6H_4CH_3$) with ellipsoid probabilities at 40%. H-atoms and solvent molecules have been omitted for clarity.

Fig. 5 Selected BP86-D3/STO-TZ2P results on the model complexes Pt(corrole)(Ph)(PhCN) (C_s) and Pt(corrole)Ph₂ (C_{2v}): (a) distances (black, Å) and Mulliken spin populations (blue); (b) spin density plots for Pt(corrole)Ph₂: top and side views; (c) the HOMO of Pt(corrole)Ph₂: top and side views.

in $CH₂Cl₂$. Note that, whereas the first oxidation potentials vary little among the different classes of metallocorroles, the reduction potentials vary considerably. Compared with Cu corroles, the Pt(w) and Au(m) corroles undergo reduction at considerably substantially more negative potentials.^{7c} Thus, $Pt(w)$ and $Au(m)$ corroles exhibit relatively large electrochemical ''HOMO–LUMO gaps'' (i.e., the algebraic difference between the first oxidation and reduction potentials) – \sim 1.4 eV for Pt(IV) and \sim 2.2 eV for Au(III). These two metal ions appear to be strongly stabilized by the trianionic corrole ligands, which would explain the resistance to reduction. In contrast, the oxidized Pt{T(p -X-Ph)C} $(m-C_6H_4CN)(p-C_6H_4CH_3)$ complexes, like Cu triarylcorroles, are readily reduced at approximately 0.1 ± 0.1 V (vs. SCE), as expected on the basis of their corrole $^{\bullet 2-}$ radical character.¹⁷

A number of metallocorrole families such as Cu, MnCl and FeCl¹⁵ meso-triarylcorroles exhibit strongly substituent-sensitive

electronic absorption spectra, with the Soret maximum shifting sensitively as a function of substituents on the *meso*-aryl groups. These have been analyzed for copper triarylcorroles with TDDFT calculations and ascribed to so-called $hyper$ character, i.e., phenyl-to-metal charge transfer (CT) character mixing into the Soret transitions.¹⁷ For CrO, MoO, Ag and Au triarylcorroles on the other hand the Soret maxima are relatively independent of substituents on the *meso*-aryl groups.^{7c,18} Against this backdrop, the Soret maxima of Pt corroles were found to be substituent-sensitive or -insensitive, depending on the overall oxidation level of the complexes. As shown in Fig. 6 and Table 3, the Pt(IV) complexes Pt $\{T(p-X-Ph)C\}(m-C_6H_4CN)$ -(PhCN) exhibit substituent-insensitive Soret maxima, whereas the oxidized Pt{T(p -X-Ph)C}{ m -C₆H₄CN)(p -C₆H₄CH₃) series exhibits some of the strongest meso substituent effects observed for metallocorroles. These observations suggest that meso-aryl substituent sensitivity occurs precisely in those cases where the corrole has substantial corrole e^{2} character; this is the case for Cu, FeCl, MnCl and the oxidized Pt corroles. For the other metallocorroles, where the corrole is relatively innocent, the Soret maxima are substituent-insensitive. Open Access Article 2012 and the published on 10 AM-

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Compared with other 5d metallocorroles such as Au and Ir corroles, which are comparatively unreactive, platinum corroles have long been of interest on account of their potential for significant axial reactivity vis-à-vis small-molecule activation. Here we have presented the first unambiguous proof of platinum insertion into the corrole macrocycle, including three single-crystal X-ray structures. Two series of complexes have been prepared in low yields: the six-coordinate $Pt(w)$ series $Pt\{T(p-X-Ph)C\}(m-C_6H_4CN)(PhCN)$ and the oxidized series $Pt\{T(p-X-Ph)C\}(m-C_6H_4CN)(p-C_6H_4CH_3)$. Ongoing research on Pt corroles focuses on developing higher-yielding syntheses and on detailed studies of C–H activation and other reactivity. Whether the compounds exhibit significant biological activity, particularly anticancer activity, remains an exciting question for the future.

 a Irreversible feature is noted at -1.43 eV.

Fig. 6 Electronic absorption spectra of the Pt($|v\rangle$ (top) and oxidized Pt($|v\rangle$ (bottom) series.

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