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NHC complex [Ni(6-Mes)(PPh3)Br]+ William J. M. Blackaby, a Sara Sabater, a Rebecca C. Poulten, Michael J. Page, a Andrea Folli, *D** Vera Krewald, *D** Mary F. Mahon, *a Damien M. Murphy, *D** Damien M. Mu Emma Richards ** and Michael K. Whittlesev ** **

Mono- and dinuclear Ni(1) products formed upon

bromide abstraction from the Ni(1) ring-expanded

Bromide abstraction from the three-coordinate Ni(i) ring-expanded N-heterocyclic carbene complex $[Ni(6-Mes)(PPh_3)Br]$ (1; 6-Mes = 1,3-bis(2,4,6-trimethylphenyl)-3,4,5,6-tetrahydropyrimidin-2-ylidene) with TIPF₆ in THF yields the T-shaped cationic solvent complex, [Ni(6-Mes)(PPh₃)(THF)][PF₆] (2), whereas treatment with NaBAr F_4 in Et₂O affords the dimeric Ni(i) product, [{Ni(6-Mes)(PPh₃)}₂(μ -Br)][BAr F_4] (3). Both 2 and 3 act as latent sources of the cation [Ni(6-Mes)(PPh₃)]⁺, which can be trapped by CO to give [Ni(6-Mes)(PPh₃)(CO)]⁺ (5). Addition of $[(Et_3Si)_2(\mu-H)][B(C_6F_5)_4]$ to **1** followed by work up in toluene results in the elimination of phosphine as well as halide to afford a co-crystallised mixture of [Ni(6-Mes)(η^2 -C₆H₅Me)][B $(C_6F_5)_4$ (4), and $[6MesH\cdots C_6H_5Me][B(C_6F_5)_4]$. Treatment of 1 with sodium salts of more strongly coordinating anions leads to substitution products. Thus, NaBH₄ yields the neutral, diamagnetic dimer [{Ni(6-Mes)}₂(BH₄)₂] (6), whereas NaBH₃(CN) gives the paramagnetic monomeric cyanotrihydroborate complex $[Ni(6-Mes)(PPh_3)(NCBH_3)]$ (7). Treatment of 1 with $NaO^tBu/NHPh_2$ affords the three-coordinate Ni(i)amido species, [Ni(6-Mes)(PPhz)(NPhz)] (8). The electronic structures of 2, 5, 7 and 8 have been analysed in comparison to that of previously reported 1 using a combination of EPR spectroscopy and density functional theory.

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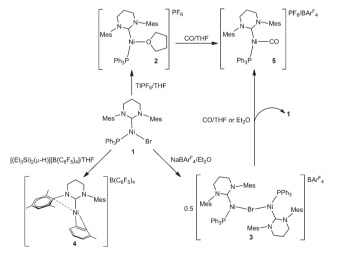
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Introduction

In a very recent review, Lin and Power referred to Ni(1) as a "... "rare" oxidation state of growing importance. 1,2 In terms of monodentate ligands, the early dependence on tertiary phosphines to stabilise Ni(1)^{3,4} has largely been superseded by the use of N-heterocyclic carbenes (NHCs) and these have facilitated the isolation of a wide range of fully characterised four-, three- and even two-coordinate Ni(1) species.^{5,6}

Over the last few years, we have used so-called ringexpanded NHCs (RE-NHCs; carbenes with ring sizes >5) for the preparation of three- and two-coordinate Ni(1) complexes with interesting stoichiometric⁷ and catalytic chemistry,⁸ as well as novel magnetic properties.9 In all cases, the starting point for our chemistry has been the three-coordinate species

1582301 and 1584193. For ESI and crystallographic data in CIF or other elecproducts.



Scheme 1

[Ni(RE-NHC)(PPh₃)Br]. The first of these to be prepared, [Ni (6-Mes)(PPh₃)Br] (1, Scheme 1), has continued to be the focus of much of our attention as it tends to yield readily isolable

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Herein, we describe the stoichiometric reactivity of 1 with a range of bromide abstracting agents to afford seven new Ni(1) complexes. Five of these are monomeric (cationic as well as neutral) and their adoption of T- or Y-shaped structures has been probed using DFT calculations.

Results and discussion

Bromide abstraction from 1 by [Tl]+, NaBArF4 and $[(Et_3Si)_2(\mu-H)][B(C_6F_5)_4]$

We have previously shown that the addition of free 6-Mes to 1 results in transfer of the bromide ligand to the outer-sphere to give the two-coordinate, cationic product [Ni(6-Mes)₂]Br.⁹ Initial efforts to abstract bromide from 1 with more typical halide abstractors such as AgX reagents (X = BF₄, NO₃, OTf) yielded only mixtures of products containing the pyrimidinium salt [6-MesH]X and the plating out of what appeared to be metallic nickel. However, when 1 was treated with TlPF₆ in THF, the three-coordinate cationic THF complex [Ni(6-Mes) (PPh₃)(THF)[[PF₆] (2) was isolated as a pale yellow solid in 85% yield (Scheme 1).

The X-ray crystal structure of 2 (Fig. 1) revealed a distorted T-shaped geometry at the Ni(1) centre, with C-Ni-P and C-Ni-O angles of 156.58(4)° and 103.74(4)° respectively. In contrast to the precursor complex 1, the Ni-C_{6-Mes} bond length was slightly elongated (1.9601(12) Å cf. 1.942(2) Å), although the Ni-P bond was unchanged. The Ni-O distance of 2.0603(9) Å was intermediate between those reported for the neutral β-diketiminato species $[L^{R}Ni(THF)]$ $(L^{R} = [HC(C(^{t}Bu)NC_{6}H_{3}(^{i}Pr)_{2})_{2}]^{-}, 2.000(1)$ \mathring{A})^{11–13} and the cationic, bis-THF complex [(THF)₂Ni $(CNAr^{Mes2})_3[OTf](Ar^{Mes2} = 2,6-(2,4,6-Me_3C_6H_2)_2C_6H_3; 2.174(2) Å,$ 2.1935(19) Å).14

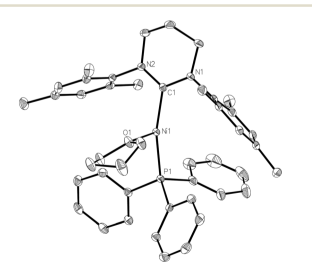


Fig. 1 Molecular structure of the cation in 2. Ellipsoids are shown at 30% probability. Hydrogen atoms have been omitted for clarity. Selected bond lengths (Å) and angles (°): Ni(1)-C(1) 1.9601(12), Ni(1)-P(1) 2.2117(3), Ni(1)-O(1) 2.0603(9), C(1)-Ni(1)-P(1) 156.58(4), C(1)-Ni(1)-O(2) 103.73(4), P(1)-Ni(1)-O(2) 99.69(3).

The ¹H NMR spectrum of 2 displayed a series of broad resonances between ca. δ 17-0 which could not be integrated. As the signals for the bound THF could not be assigned, we were unable to establish spectroscopically the lability of the THF ligand. However, X-ray crystallography repeatedly revealed the presence of THF following recrystallization of 2 from a number of solvents (CH₂Cl₂, C₆H₅F, C₆H₆) suggesting that the THF cannot be easily dissociated from the nickel.

The formation of TlBr as a side-product in the synthesis of 2 proved problematic, as even following multiple recrystallisations, complete removal was not always achievable. This manifested itself in EPR spectra of 2 (ESI†), but more obviously in reactions with CO (vide infra). Fig. 2 shows the EPR spectrum of a 'clean' sample of complex 2 (Fig. 2d). The spin Hamiltonian parameters of the EPR spectra of all of the species shown in Fig. 2 are listed in Table 1, and are discussed further below.

In an attempt to circumvent the problem of TlBr contamination, 1 was reacted instead with NaBArF4 in THF. No bromide abstraction resulted. However, a reaction between 1 and NaBAr^F₄ took place upon changing the solvent to Et₂O, affording large orange crystals. These proved to be of the unusual cationic, mono-bromide bridged dimer, [{Ni(6-Mes) $(PPh_3)_2(\mu-Br)[BAr_4]$ (3, Scheme 1) rather than 2. The structure of 3 (Fig. 3) comprised of two {Ni(6-Mes)(PPh₃)} fragments and a close to symmetrically Ni bound µ-bromide ligand (Ni1-Br1 2.3803(6) Å, Ni2-Br1 2.3688(6) Å) in an arrangement distorted from linearity (∠Ni1-Br1-Ni2 167.64(3)°). This is adopted presumably to minimise the steric demands of the ligand substituents in the solid-state. Unsurprisingly, these steric demands also preclude the ligands from eclipsing each other relative to the Ni...Ni axis. Thus, there is 46.8(1)° angle between the mean planes containing atoms Br1, Ni1, P1, C1 and Br1, Ni2, P2, C41, respectively. The Ni...Ni separation exceeds 4.7 Å.

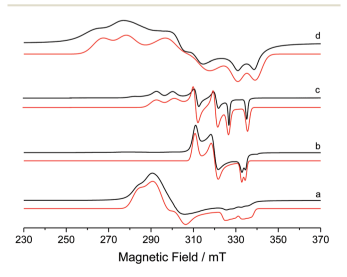


Fig. 2 Experimental (black) and simulated (red) X-band CW EPR spectra of (a) 1, (b) 5, (c) 8 and (d) 2 in frozen THF solution at 140 K.

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Table 1 Spin Hamiltonian parameters for $[Ni(6-Mes)(PPh_3)X]^{0/+}$ dissolved in THF: experimental and computed g-tensors, and phosphorus and bromide (indicated by #) superhyperfine coupling constants (MHz), with Euler angles derived from DFT calculations

	g values				Euler angles/rad			A values/MHz			Euler angles/rad			
	g_1	g_2	g_3	$g_{\rm iso}$	α	β	γ	$\overline{A_1}$	A_2	A_3	$a_{\rm iso}$	α	β	γ
Ni(6-N	1es)(PPh ₃)Br] 1												
Expt	2.050	2.265	2.365	2.227	-2.270	2.619	1.643	184 -6 [#]	$194 \\ -27^{\#}$	250 70 [#]	$209 \\ 12^{\#}$	$0.035 \\ 1.580^{\#}$	$1.580 \\ 1.566$	$1.936 \\ -1.328$
DFT	2.055	2.252	2.285	2.197	-2.276	2.246	1.628	173 -19 [#]	173 -21 [#]	204 51 [#]	$\begin{array}{c} 183 \\ 4^{\#} \end{array}$	0.176 0.093 [#]	1.590 1.578 [#]	$1.968 \\ 1.859$
Ni(6-M	1es)(PPh ₃)(THF)∏PI	F ₆] 2											
Expt DFT	2.025 2.013	2.210 2.315	2.490 2.389	2.242 2.239	1.497 -1.580	2.540 1.203	1.458 1.622	292 -7	210 -15	419 -19	$307 \\ -14$	0.646 1.700	2.969 1.635	-0.474 -0.436
Ni(6-M	1es)(PPh ₂)(CO)][PF ₆	15											
Expt	2.035	2.121	2.185	2.114	-3.114	3.064	-1.605	21	29	48	33	1.738	1.560	1.287
DFT	2.044	2.117	2.155	2.105	3.100	3.051	-1.654	14	24	42	26	1.728	1.558	1.235
Ni(6-N	1es)(PPh3)(NCBH ₃)]	7											
Expt	2.028	2.225	2.373	2.209	1.176	2.554	-1.575	260	300	260	273	-0.815	1.427	1.780
DFT	2.020	2.286	2.303	2.203	-1.950	1.051	1.568	185	187	220	197	-0.865	1.423	1.778
[Ni(6-N	1es)(PPh ₃)(NPh ₂)] 8												
Expt	2.050	2.150	2.290	2.163	-1.459	0.209	0.877	230	265	300	265	1.621	1.602	-1.216
DFT	2.059	2.162	2.238	2.152	-1.513	0.227	0.930	248	249	280	259	1.554	1.610	-1.190

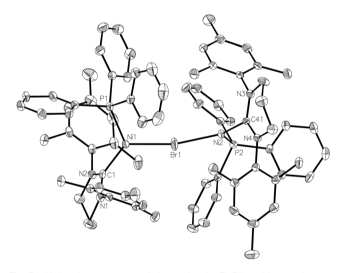


Fig. 3 Molecular structure of the cation in 3. Ellipsoids are shown at the 30% probability level. Hydrogen atoms have been omitted for clarity. Selected bond lengths (Å) and angles (°): Ni(1)–C(1) 1.937(4), Ni(1)–P(1) 2.2172(12), Ni(1)–Br(1) 2.3803(6), Ni(2)–C(41) 1.935(4), Ni(2)–P(2) 2.2186(11), Ni(2)–Br(1) 2.3688(6), C(1)–Ni(1)–P(1) 124.15(11), C(1)–Ni(1)–Br(1) 122.16(11), P(1)–Ni(1)–Br(1) 113.68(4), Ni(1)–Br(1)–Ni(2) 167.64(3).

3 exhibited a paramagnetic NMR spectrum in Et_2O solution, as well as a room temperature magnetic moment (μ_{eff} , Evans method) of 2.51 μ_B , consistent with the presence two Ni(1) centres. Density functional theory (DFT) calculations based on the crystal structure coordinates with optimised hydrogen atom positions afforded Mulliken spin populations of 0.78 at each Ni centre, thus correctly representing the formal Ni(1) oxidation states. Broken-symmetry DFT predicted antiferromagnetic coupling between the two nickel centres,

with a medium strength negative exchange coupling constant (density functional dependent: TPSSh: -97.6 cm⁻¹, B3LYP: -76.1 cm⁻¹, PBE0: -64.3 cm⁻¹, M06: -69.3 cm⁻¹). This coupling appears weak enough to allow significant population of the high-spin state at room temperature: indeed, a Boltzmann population distribution analysis showed that *ca.* 40% of the triplet state would be populated at 300 K (ESI†).

EPR spectroscopy revealed that, at least in diethyl ether solution, 16 the two Ni(1) centres were [Ni(6-Mes)(PPh_3)Br] (1) and (solvated) [Ni(6-Mes)(PPh_3)]^+. Thus, the CW EPR spectrum (140 K, frozen Et_2O glass; ESI†) clearly contained signal intensity from 1, as well as a second Ni(1) centre. The similar profile of this second species to that of 2 suggests it is the diethyl ether complex, [Ni(6-Mes)(PPh_3)(OEt_2)]^+. Dissociation of 3 was also supported by the appearance of [Ni(6-Mes)(PPh_3)(CO)][BAr^F_4] (5-BAr^F_4, vide infra) by IR spectroscopy following treatment of the dimer with CO in Et_2O solution.

Treatment of **1** with an equimolar amount of $[(Et_3Si)_2(\mu-H)][B(C_6F_5)_4]^{17}$ in fluorobenzene resulted in an instantaneous colour change from yellow to red. Upon layering with toluene, light green crystals formed, which consisted of a co-crystallised mixture of the Ni(I) toluene salt, [Ni(6-Mes) $(\eta^2-C_6H_5Me)][B(C_6F_5)_4]$ (4), and $[(6-MesH)\cdots C_6H_5Me)[B(C_6F_5)_4]$.

The X-ray structure of the metal-containing cation is shown in Fig. 4. The metrics of the coordinated toluene ligand revealed short Ni–C24/C25 distances (2.054(3) Å and 2.092(3) Å), intermediate Ni–C23/C26 distances (2.152(3) Å, 2.202(3) Å) and two substantially longer interactions (Ni–C27 2.271(3) Å, Ni–C28 2.241(3) Å), consistent with an η^2 rather than η^6 bound arene ligand. To overcome electron deficiency, this then formally 13-electron nickel centre exhibits a close interaction with the *ipso-*C of one of the mesityl rings (Ni–C5 2.525(2) Å; *cf.* Ni–C14 3.367(2) Å). Comparable stabilising close contacts have been

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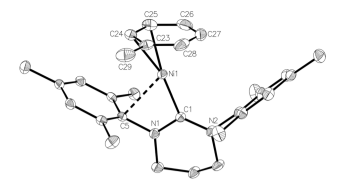


Fig. 4 Structure of the metal-containing, cationic component in compound 4. Ellipsoids are shown at the 30% probability level. Hydrogen atoms have been omitted for clarity. Selected bond lengths (Å) and angles (°): Ni(1)-C(1) 1.914(2), Ni(1)-C(23) 2.152(3), Ni(1)-C(24) 2.054(3), Ni(1)-C(25) 2.092(3), Ni(1)-C(26) 2.202(3), Ni(1)-C(27) 2.271(3), Ni(1)-C(28) 2.241(3), N(1)-C(1)-Ni(1) 104.72(15), N(2)-C(1)-Ni(1) 134.13(17).

seen in other coordinatively unsaturated metal complexes bearing bulky NHCs. 19

Further analysis revealed that the toluene ligand lies almost parallel to one of the fluoroaryl ligands of the $[B(C_6F_5)_4]^-$ anion. A value of 9.9° for the angle between the relevant least-squares aromatic ring planes, 3.78 Å for the centroid–centroid distance between these rings and 3.27 Å for the shortest distance from the centroid of one ring to the mean plane of the other support the presence of offset π – π stacking and additional π -stabilisation of the complex.

The ratio of 4: toluene stabilised pyrimidinium cation varied from one synthesis to another. The 'best' ratio, determined crystallographically, in terms of optimising the percentage of nickel complex yielded, was 65:35. This was achieved by (i) performing the complete reaction in a glovebox and (ii) washing the $[(Et_3Si)_2(\mu\text{-}H)][B(C_6F_5)_4]$ five times with hexane and drying overnight. Although the irreproducibility in yield of 4 frustrated efforts to further characterise the complex, the synthetic approach was validated by isolation of the corresponding mesitylene analogue $[Ni(6\text{-}Mes)(\eta^2\text{-}C_6H_3Me_3)][B(C_6F_5)_4]$ (ESI†) through reaction of 1 with $[(Et_3Si)_2(\mu\text{-}H)][B(C_6F_5)_4]$ in C_6H_5F , followed by crystallisation from $C_6H_5F/\text{mesitylene}$. Again co-crystallisation with pyrimidinium salt $([(6\text{-}MesH)\cdots C_6H_3Me_3][B(C_6F_5)_4])$ was found.

Synthesis and characterisation of [Ni(6-Mes)(PPh₃)(CO)]⁺

Exposure of a THF solution of 2 to 1 atm CO led to an almost instantaneous colour change from yellow to pale green. Removal of the CO atmosphere after *ca.* 1 min, followed by recrystallization of the residue from THF/hexane gave light green/yellow crystals of the Ni(I) carbonyl complex, [Ni(6-Mes) (PPh₃)(CO)][PF₆] (5-[PF₆]). The EPR spectrum of 5-[PF₆] (Fig. 2b) and the corresponding DFT calculations of the spin Hamiltonian parameters is discussed in further detail below.

The X-ray crystal structure (Fig. 5) revealed a similarly distorted T-shaped geometry to that of 2 (\angle C-Ni-P = 151.93(9)°). Both the Ni-CO bond length (Ni1-C23 = 1.787(3) Å) as well as

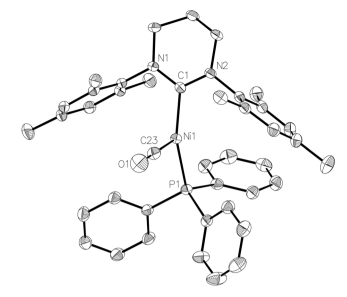


Fig. 5 Molecular structure of the cation in 5. Ellipsoids are shown at the 30% probability level. Hydrogen atoms have been omitted for clarity. Selected bond lengths (Å) and angles (°): Ni(1)-C(1) 1.943(3), Ni(1)-C(23) 1.787(3), Ni(1)-P(1) 2.2374(8), C(23)-O(1) 1.132(4), C(1)-Ni(1)-P(1) 151.93(9), C(1)-Ni(1)-C(23) 106.24(13), P(1)-Ni(1)-C(23) 101.76(11), O(1)-C(23)-Ni(1) 178.0(3).

the $\nu({\rm CO})$ in the IR spectrum (2032 cm $^{-1}$) showed good agreement with the few other (predominantly neutral) reported Ni(i)–CO complexes. $^{20-23}$ As expected, 5-[PF $_6$] displayed broad, paramagnetic $^1{\rm H}$ and $^{13}{\rm C}$ NMR spectra, although upon subjecting a THF solution to 1 atm $^{13}{\rm CO}$, we observed the rapid appearance of an isotopically enhanced carbonyl resonance in the $^{13}{\rm C}$ NMR spectrum at δ 198.1, consistent with reversible coordination of the carbonyl ligand. 22 Since the SOMO is an orbital with antibonding character between the Ni centre and the CO ligand, one would expect a weakened Ni–CO bond, and it may be this that facilitates the facile exchange with $^{13}{\rm CO}.^{22,23}$

As aforementioned, the presence of residual TlBr in samples of 2 was apparent from reactions with CO, particularly after prolonged periods. At times >1 min, the initial pale green solution of 5-[PF₆] became orange, and then over ca. 48 h, yellow. A small number of crystals were isolated from this yellow solution and structurally characterised as the Ni(II) salt, [Ni(6-Mes)(PPh₃)(CO)Br][PF₆] (ESI†).

Reactions of 1 with NaBH₃X (X = H, CN) and NaNPh₂

Efforts to abstract the bromide ligand from 1 using sodium salts of more coordinating anions generated the Ni(i) products 6-8 shown in Scheme 2.

NaBH₄ addition to a yellow THF suspension of 1 in the presence of EtOH rapidly generated a green solution, from which dark green crystals of the dimeric borohydride complex, $[\{Ni(6\text{-Mes})\}_2(\mu\text{-BH}_4)_2]$ (6) were isolated in 90% yield. The X-ray crystal structure of 6 is shown in Fig. 6 and revealed asymmetry across the $\{Ni_2B_2\}$ moiety. Thus, B1 is closer to Ni2 than to

Mes NaBH₄, THF/EtOH NHPh₂/NaO'Bu, THF

NaBH₄, THF/EtOH NHPh₂/NaO'Bu, THF

NaBH₃(CN) NHPh₂/NaO'Bu, THF

NaBH₃(CN) NHPh₂
Nes NHPh₃ NH NHPh₂
Nes NHPh₃ NH NHPh₂
NABH₃(CN) NHPh₃
NABH₃(CN) NHPh₂
NHPh₃(CN) NHPh₃(CN) NHPh₂
NHPh₃(CN) NHPh₂
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NHPh₃(CN) NHPH

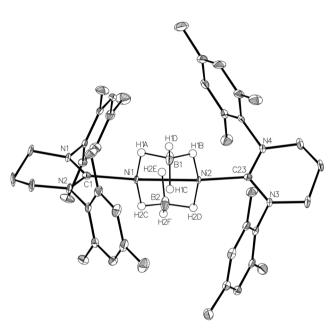


Fig. 6 Molecular structure of compound 6. Ellipsoids are displayed at 50% probability. The minor disordered component of C25 and hydrogen atoms (with the exception of those bonded to boron centres) have been omitted for clarity.

Ni1, and the reverse situation prevails for B2 (Ni1···B1 2.180(2), Ni1···B2 2.143(3), Ni2···B1 2.144(2), Ni2···B2 2.181(3) Å). Overall, the data suggest a rare $\mu^2,\eta^1:\eta^1$ coordination mode 24,25 for the borohydride based on B1, with H1C being equidistant from both metal centres (Ni1-H1C 2.16(4) Å; Ni2-H1C 2.11(4) Å). The B2 based borohydride has a similar coordination mode once experimental errors are taken into consideration. However, H2E may be closer to Ni1 (1.92(4) Å) than to Ni2 (2.12(4) Å), which would indicate a tendency towards an even more unusual $\mu^2,\eta^2:\eta^1$ coordination mode. In an effort to further probe the bonding of the borohydrides, a neutron dataset was collected, but a phase transition hampered acquisition of any additional insights (see Experimental). IR spectroscopy provided little in the way of diagnostic characterisation of any particular coordination mode, as only a single,

broad $\nu(B-H)$ absorption band was measured at 2378 cm⁻¹ in KBr.

6 adds to the surprisingly few examples of structurally characterised nickel borohydride complexes, ^{26–28} in amongst which there is just a single example of a bridging borohydride species²⁹ and a Ni(1)–BH₄ complex.^{27,30}

The 2.4218(4) Å separation of the two Ni centres is suggestive of a Ni–Ni bond, high which explains the diamagnetism of the complex in solution. The borohydride groups appear fluxional, with just a single broad resonance apparent in both the H ($ca. \delta -5.1$) and Hg{1H} ($ca. \delta -30$) NMR spectra. The proton signal sharpened slightly with Hg decoupling, but remained broad even down to 198 K.

NaBH₄ was replaced by NaBH₃(CN) in an effort to prepare a Ni–Ni dimer analogous to 6 but with an asymmetric and potentially simpler coordination mode. Instead, the paramagnetic, monomeric Ni(i) complex, [Ni(6-Mes)(PPh₃)(NCBH₃)] (7, Fig. 7) was formed. Cyanotrihydroborate complexes remain (like their [BH₄]⁻ counterparts) extremely rare for nickel,³² and unknown for Ni(i). Trigonal planar 7 exhibited a Ni–N bond length of 1.924(2) Å which, although shorter than that reported in [(tren)Ni(μ -NCBH₃)]₂²⁺ (tren = 2,2',2"-triaminoethylamine),^{32a} is consistent with values reported for a number of monomeric cobalt derivatives.³³ The EPR spectrum of 7 (see ESI†) confirms the paramagnetism of this complex, but is again poorly resolved due to overlapping intensity originating from the precursor complex 1.

We have previously reported that 1 reacts with NaO^tBu to provide a low yielding route to the Ni(0) product, [Ni(6-Mes) (PPh₃)₂].³⁴ A repeat of this reaction in the presence of diphenylamine³⁵ afforded the three-coordinate Ni(1) amido

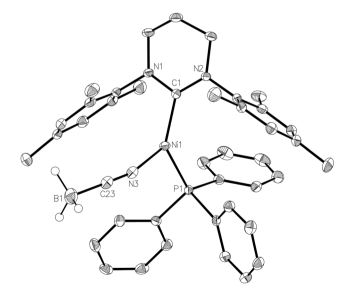


Fig. 7 Molecular structure of compound 7. Ellipsoids are displayed at 30% probability. The minor disordered component of C3 and hydrogen atoms (with the exception of those bonded to boron) have been omitted for clarity. Selected bond lengths (Å) and angles (°): Ni(1)-C(1) 1.957(2), Ni(1)-P(1) 2.2201(6), Ni(1)-N(3) 1.924(2), C(1)-Ni(1)-P(1) 125.14(7), C(1)-Ni(1)-N(3) 132.38(9), P(1)-Ni(1)-N(3) 102.47(7).

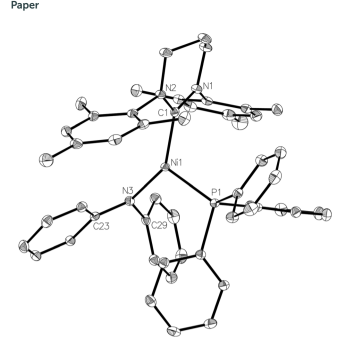


Fig. 8 Molecular structure of 8. Ellipsoids are shown at 30% probability. Hydrogen atoms and the minor disordered component of C3 have been omitted for clarity. Selected bond lengths (Å) and angles (°): Ni(1)–C(1) 1.9612(14), Ni(1)–P(1) 2.2337(4), Ni(1)–N(3) 1.9350(12), C(1)-Ni(1)–P(1) 109.25(4), C(1)–Ni(1)–N(3) 142.47(5), P(1)-Ni(1)–N(3) 108.13(4).

complex, [Ni(6-Mes)(PPh₃)(NPh₂)] (8), as a deep-red solid that could be isolated in high (72%) yield.

The X-ray crystal structure of the complex (Fig. 8) revealed a more acute C-Ni-P angle (109.25(4)°) than that found in the starting bromide complex (117.01(6)°), which most likely results from the need to alleviate steric clashes between the P- and N-bound phenyl groups. Indeed, replacing the phenyl groups with methyl groups in silico and fully relaxing the geometry showed that steric effects play a role in shaping the geometry of 8: in the case of the (hypothetical) [Ni(6-Mes)(PPh₃)(NMe₂)] complex 8-Me with a less bulky NMe₂ ligand, the C-Ni-N angle decreased by 6.7° while the P-Ni-C angle increased by 6.5°. Concomitantly, the Ni-N bond decreased from 1.94 Å in the crystal structure to 1.84 Å in 8-Me (fully relaxing the crystal structure of 8 results in a bond length of 1.90 Å). Sterics may also account for the non-planarity of the amido group (dihedral angle between Ni1, N3, C23 and C29 of ca. 160°), as well as the elongation of the Ni-N distance (1.9350(12) Å) compared to those in either $[Ni(d^tbpe)\{N(2,6^{-i}Pr_2C_6H_3)H\}]$ (1.881(2) Å; $d^tbpe =$ ${}^{t}\text{Bu}_{2}\text{P(CH}_{2})_{2}\text{P}{}^{t}\text{Bu}_{2})^{36}$ or $[\text{Ni}(\text{PPh}_{3})_{2}\{\text{N}(\text{SiMe}_{3})_{2}\}]$ (1.88(1) Å). 4,37

Electronic structure analysis of complexes 1, 2, 5, 7 and 8

Inspection of the angles around Ni in complexes 1, 2, 5, 7 and 8 (as well as 8-Me) shows that the L-Ni-P angle varies least across all complexes (<10°, Fig. 9). Complexes 1 and 7 can be classed as Y-shaped (\angle C-Ni-L, \angle P-Ni-C > \angle L-Ni-P, Fig. 9), whereas complexes 2 and 5 are T-shaped (\angle P-Ni-C > \angle C-Ni-L, \angle L-Ni-P angles, see Fig. 9). Complex 8 represents a more sym-

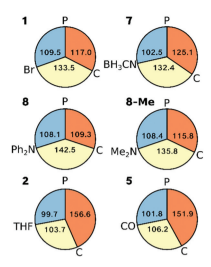


Fig. 9 Pie diagrams representing the angles around the Ni ion in the crystal structures for 1, 2, 5, 7, 8 and the geometry optimised structure of 8-Me; orange: $\angle P-Ni-C$, yellow: $\angle C-Ni-L$, blue: $\angle L-Ni-P$, where L stands for the respective ligand.

metric Y-shaped case with a larger C–Ni–L angle than in **1** and 7 (142.5° vs. 133.5°, 132.4°) and a smaller P–Ni–C angle (109.2° vs. 117.0°, 125.1°). Although one may also view complex **8** as a T-shaped complex with \angle C–Ni–L as the largest angle, this appears to be purely due to steric and not electronic effects, as is seen by **8-Me** adopting a more Y-shaped geometry.

The overall geometric changes in fully relaxing the crystal structures are small (see ESI†). Most importantly, the striking consistency of the Ni–P and Ni–C bond lengths in the crystal structures across the series (variation <0.03 Å and <0.02 Å, respectively) is preserved upon geometry optimisation (variation <0.03 Å and <0.04 Å, respectively: ESI†).

The tendency of three-coordinate transition metal d9 complexes to form either T- or Y-shaped geometries is due to the Jahn-Teller effect, thus lifting orbital degeneracy $(d_{xy}, d_{x^2-y^2})$ at the ideal D_{3h} symmetry (Fig. 10a). MO theory predicts that the SOMO in a T-shaped d⁹ complex will be of $d_{x^2-y^2}$ character, whereas in a Y-shaped d^9 complex, it will be of d_{xy} character (Fig. 10a), in agreement with the dominant character of the DFT-calculated orbitals (Fig. 10b). For Ni(1) complexes, this was most recently discussed by the groups of Holland and Lee, 13,21 and prior to that, by Pietrzyk.³⁸ Holland and co-workers²¹ rationalised the formation of T- vs. Y-shaped complexes with a charge donation analysis (natural bond orbital analysis).39 Their findings indicated that a T-shape is inherently favoured by d9 complexes, but a Y-shape can result when there is increased donation of charge from the ligands to the metal centre, thus effectively partially reducing the metal centre. In the present case, the analysis of Mulliken⁴⁰ and Chelpg⁴¹ charges did not reveal a clear connection between charges and geometry. Likewise, the Mulliken spin populations on the Ni ion do not show a significant variation across the series (ESI†).

The CW X-band EPR spectra of complexes 1, 2, 5 and 8 were shown in Fig. 2. The resulting spin Hamiltonian parameters,

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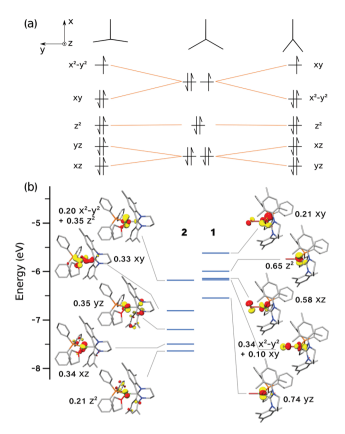


Fig. 10 (a) Schematic molecular orbitals diagrams expected for threecoordinate d^9 complexes, left to right: T-shaped, ideal D_{3h} , Y-shape; a larger ligand field splitting would lead to crossings of MO energy levels. (b) Quantitative MO diagrams for complexes 1 (right) and 2 (left) based on energies and characters of spin-up orbitals.

notably the g-tensor and A(31P)-tensor components were extracted by simulation, and are listed in Table 1. All spectra display a rhombic g profile, with one component (g_1) close to the free spin value of g_e (2.0023), indicating that there is considerable $3d_{z^2}$ character in the SOMO. The large Δg shifts observed for the $g_{2,3}$ parameters result from the large spinorbit coupling constant for Ni (ζ_{Ni^+} = 565 cm⁻¹).

The considerably broadened linewidths mainly arise from g-strain effects and not fully resolved superhyperfine coupling to the ³¹P nucleus of the PPh₃ ligand. In the case of complex 1, an additional hyperfine broadening of ca. 60 MHz is present. This is in the order of the largest A_3 value for the Br nucleus (50.69% ⁷⁹Br, 49.31% ⁸¹Br; both possessing nuclear spin 3/2), hence impeding resolution of the quartets arising from the coupling of the unpaired electron to this nuclear spin. The DFT-derived parameters are also listed in Table 1 and are in reasonable agreement with the experimentally determined values. All complexes, with the exception of 5, display large, predominantly isotropic superhyperfine coupling to the ³¹P nucleus, in good agreement with the calculated values.

The relative orientations of the g- and A-tensors for the cationic complex 5 are shown in Fig. 11, alongside the spin density. As a comparison, g- and A-tensor orientations and spin densities derived from the DFT calculations of EPR parameters for starting complex 1 are given in Fig. 11b (corresponding figures for complexes 2, 7 and 8 are given in the ESI†).

As already mentioned, the ³¹P superhyperfine interaction is almost entirely isotropic, therefore an explanation for the much smaller HFC in the case of complex 5 compared to starting complex 1 (see $a_{iso}(^{31}P)$ in Table 1) can be found by simply looking at the overall spin density on the ³¹P nuclei, neglecting the relative orientations of the A(31P) frames in each of the complexes (isotropic interaction is orientation independent). As the insets in Fig. 11 clearly show, there is a significantly less spin density on the ³¹P nucleus of 5 when compared to 1, which readily explains the much lower hyperfine interaction found experimentally and computationally. In fact, the spin density on the ³¹P nucleus of 5 is so small that two of the principal values of the A(31P) tensor for this complex are smaller than the overall broadening caused by g-strain effects and are completely unresolved at X-band. Only the $A_3(^{31}P)$ component of the tensor is visible at X-band. In the spectrum in Fig. 2, the

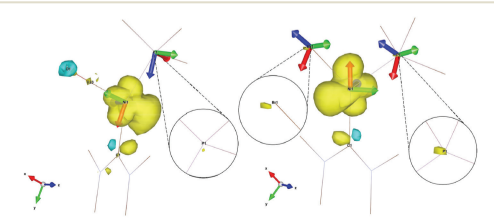


Fig. 11 Spin density contour plot with g- and A-frames for [Ni(6-Mes)(PPh₃)(CO)][PF₆] (5, left) and [Ni(6-Mes)(PPh₃)Br] (1, right). Ligands are truncated for clarity.

 $A_3(^{31}P)$ component is found on g_1 because of almost exact and complete alignment between the A_z axis and the g_x axis, as deducible by Fig. 11. The extent of anisotropy and rhombicity for 5 is much lower than for the other complexes, and is similar to that reported previously for $[L^{Me}Ni(CO)]$ ($L^{Me} = [HC(C(Me)NC_6H_3(^iPr)_2)_2]^-)$, with g values of 2.01, 2.17 and 2.19 and T-shape geometry.

Notably, the calculated ³¹P superhyperfine coupling for the two cationic complexes (2 and 5) are an order of magnitude smaller than for the series of neutral complexes reported. Whilst the experimental and calculated values for 5 are in reasonable agreement, the experimentally observed hyperfine for 2 does not match the DFT-derived values and bears closer resemblance to the neutral complexes; currently, we do not have an explanation for this observation.

Fig. 12 shows the computed SOMO d-character of the different complexes as a function of the difference between the largest and the two smaller angles (e.g. (\angle C-Ni-Br) – (\angle P-Ni-C) – (\angle Br-Ni-P) for 1, denoted as $\Delta\Delta$ (bond angle)), taken as an index for the deviation from ideal D_{3h} symmetry. Noticeable in Fig. 12a is the smaller difference in d_{xy} and $d_{x^2-y^2}$ contributions to the SOMO for complexes 1 and 7 (closest examples to D_{3h} symmetry where d_{xy} and $d_{x^2-y^2}$ are degenerate), and the increasingly higher $d_{x^2-y^2}$ character (simultaneously to d_{xy} contributions approaching zero) when moving away from ideal D_{3h} symmetry towards T-shape symmetry. Both these observations seem to be in good agreement with what was described

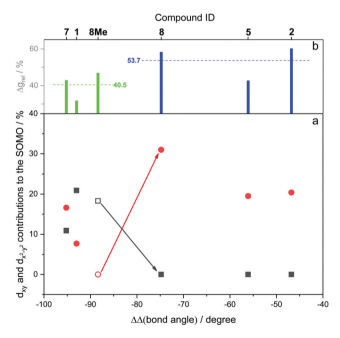


Fig. 12 (a) d_{xy} and $d_{x^2-y^2}$ contributions to the spin up SOMO orbital as a function of the difference between the largest and the two smaller angles ($\Delta\Delta$ (bond angle)). Black squares refer to d_{xy} and red circles to $d_{x^2-y^2}$. Empty symbols refers to d orbital contributions from fully optimised structures rather than X-ray crystal structures. (b) $\Delta g_{\rm rel}$ values as a function of the double bond angle variation. Average $\Delta g_{\rm rel}$ values for Y-and T-shape geometries are also reported.

previously and represented in Fig. 9. Very interesting is the case of complex 8, which as we noted above may be regarded as a T-shape complex with ∠C-Ni-L as the largest angle. However, a fully geometry optimised version of the same complex where the amido phenyl substituents were replaced by methyl substituents (8-Me, vide supra) showed angles that are similar to the Y-shape complexes 1 and 7. Our interpretation is that this compound is electronically inclined to be a Y-shape (similar to the other neutral compounds of the present series), however large steric strain pushes the amido group towards the carbene ligand, thus geometrically distorting it towards a T-shape. Orbital distribution and coordination geometry should reflect the shape and magnitude of the g tensor associated with the paramagnetic centre. In Fig. 12b, experimental $\Delta g_{\rm rel}$, a parameter used to evaluate the shape of the diagonalised g tensor and calculated according to eqn (1), is also reported as a function of the $\Delta\Delta$ (bond angle).

$$\Delta g_{\rm rel} = \frac{|g_3 - g_2|}{|g_3 - g_1|} \times 100 \tag{1}$$

It can be seen that an increase in the $d_{x^2-y^2}$ contribution to the SOMO corresponds to a shift of the g_2 value away from g_3 towards g_1 , and indeed for T-shape complexes g_2 is closer to g_1 than to g_3 , highlighting a geometry induced shape shifting of the g tensor.

Conclusions

Treatment of the three-coordinate Ni(1) complex [Ni(6-Mes) (PPh₃)Br] (1) with a variety of bromide abstracting reagents has yielded a series of new mono- and dinuclear nickel products. Of most interest are the three-coordinate d⁹ complexes, 2, 5, 7 and 8, of general formula [Ni(6-Mes)(PPh₃)X]^{0/+} that distort from ideal D_{3h} symmetry by forming either T-shaped or Y-shaped geometries. These structural differences manifest in different electronic structure characteristics, namely that the SOMO for a T-shape complex is expected to be of $d_{x^2-y^2}$ character, whereas for a Y-shape complex, it will be of d_{xy} character. Electron paramagnetic resonance spectroscopy was used to derive spin Hamiltonian parameters for this series of threecoordinate Ni(1) complexes, which showed that all complexes have a rhombic g-tensor profile and that the 31P superhyperfine couplings are predominantly isotropic. The much lower magnitude of 31P superhyperfine coupling constants observed for the CO-containing complex 5 was explained with a smaller spin density found at the phosphorus ligand as predicted by density functional theory calculations. The overall computed spin densities in this series are polarised differently for the Yand T-shaped complexes, namely with a larger lobe trans to the phosphine ligand in the former case as opposed to a larger lobe trans to the ligand in the latter case. This directly affects the shape and magnitude of the g-tensor: while all complexes have a rhombic **g**-tensor with $g_1 \ll g_2 < g_3$, a larger $d_{x^2-y^2}$ contribution to the SOMO shifts g_2 closer to g_1 .

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Experimental

General considerations

All manipulations were carried out using standard Schlenk, high vacuum and glovebox techniques. Solvents were purified using an MBraun SPS solvent system (hexane, Et₂O) or under a nitrogen atmosphere from sodium benzophenone ketyl (benzene, THF). C₆D₆ and THF-d₈ were vacuum transferred from potassium. NMR spectra were recorded on Bruker Avance 400/500 NMR and Agilent 500 MHz spectrometers and referenced to solvent signals as follows: benzene (1 H, δ 7.16; 13 C $\{^{1}H\}, \delta$ 128.0), THF $(^{1}H, \delta$ 3.58; $^{13}C\{^{1}H\}, \delta$ 67.6); $^{31}P\{^{1}H\}, \text{ exter-}$ nally to 85% H_3PO_4 (δ 0.0); ¹⁹F, externally to CFCl₃ (δ 0.0). analyses were performed by Microanalysis Ltd, Okehampton, Devon, UK. 1 and $[(Et_3Si)_2(\mu-H)][B(C_6F_5)_4]$ were prepared according to literature methods.8,17

$[Ni(6-Mes)(PPh_3)(THF)][PF_6](2)$

A THF (10 mL) solution of TlPF₆ (95 mg, 0.27 mmol) was added to a J. Young's resealable ampoule containing 1 (163 mg, 0.23 mmol) and the beige suspension was stirred for 2 h. This was cannula filtered, the filtrate concentrated to half volume and hexane (10 mL) added slowly to form a pale yellow precipitate. This was isolated by cannula filtration, recrystallised from THF/hexane and dried in vacuo. Yield: 138 mg (85%). ¹H NMR (500 MHz, THF- d_8 , 298 K): δ 16.9 (br s), 9.9 (br s), 6.0 (br s), 5.3 (br s), 3.1 (br s); anal. calcd for $C_{44}H_{51}N_2OF_6P_2Ni$ (858.48): C 61.56%, H 5.99%, N 3.26%; found: C 61.39%, H 5.85%, N 3.18%; μ_{eff} (Evans method, THF, 298 K): 2.2 μ_{B} .

$[{Ni(6-Mes)(PPh_3)}_2(\mu-Br)][BAr^F_4](3)$

1 (100 mg, 0.14 mmol) and NaBAr^F₄ (130 mg, 0.15 mmol) were dissolved in Et₂O (10 mL) and the solution stirred for 16 h in a J. Young's resealable ampoule. The solution was concentrated, filtered and layered with pentane (10 mL) to form orange crystals. Yield: 125 mg (81%). Anal. calcd for C₁₁₂H₉₈BN₄F₂₄P₂BrNi₂ (2225.96): C 60.43%, H 4.44%, N 2.52%; found: C 60.07%, H 4.68%, N 2.36%; μ_{eff} (Evans method, Et₂O, 298 K): $2.5\mu_{\rm B}$.

$[Ni(6-Mes)(PPh_3)(CO)][PF_6](5)$

To a degassed THF solution (0.5 mL) of 2 (20 mg, 0.02 mmol), 1 atm of CO was added to the stirring solution. An immediate colour change to dark yellow/green occurred, and after 1 minute the solution was reduced to dryness. The residue was extracted into THF (0.5 mL), filtered and layered with hexane (2 mL) to produce green crystals. Yield: 15 mg (79%). ¹H NMR (500 MHz, THF- d_8 , 298 K): δ 16.9 (br s), 10.7 (br s), 9.9 (br s), 8.4 (s), 7.7 (br s), 7.3 (s), 7.0 (s), 6.4 (br s), 5.3 (br s), 4.9 (br s), 3.9 (br s), 3.4 (br s), 2.5 (br s), 2.3 (br s), 2.3 (br s); IR (ν , cm⁻¹: THF): 2035 (CO); IR (ν , cm⁻¹: KBr): 2030 (CO); anal. calcd for C₄₁H₄₃N₂OF₆PNi₂ (814.39): C 60.47%, H 5.32%, N 3.44%; found: C 60.45%, H 5.58%, N 2.96%; μ_{eff} (Evans method, THF, 298 K): $1.8\mu_{\rm B}$.

$[{Ni(6-Mes)}_2(\mu-BH_4)_2]$ (6)

1 (100 mg, 0.138 mmol) and NaBH₄ (16 mg, 0.423 mmol) were placed in a J. Young's resealable ampoule in THF (10 mL). EtOH (1 mL) was added and the suspension was stirred for 5 min at room temperature to give a dark green solution. The volatiles were removed in vacuo, the residue extracted into benzene (2 × 10 mL) and evaporated to dryness. The green crude was washed with hexane (2 × 10 mL) at 195 K to give 6 as a green solid. Yield: 47 mg (90%). Crystals suitable for X-ray diffraction were obtained by slow diffusion of hexane into a concentrated THF solution of 6. ¹H NMR (THF-d₈, 400 MHz, 298 K): δ 6.71 (s, 8H, CH), 3.10 (t, 8H, ${}^{3}J_{HH}$ = 5.7 Hz, NCH₂), 2.37 (s, 12H, CH_3), 2.02 (br s, 28H, CH_3 and NCH_2CH_2), -5.72 (br s, 8H, B H_4); ¹³C{¹H} NMR (THF- d_8 , 100 MHz, 298 K): δ 213.0 (s, NCN), 144.6 (s, N-ipso-C), 136.2 (s, o-C), 136.1 (p-C), 129.7 (CH), 45.2 (NCH₂), 22.7 (CH₂), 21.7 (CH₃), 18.6 (CH₃); ¹¹B NMR (THF d_8 , 128 MHz, 298 K): δ –32.0 (br s); IR (ν , cm⁻¹: KBr): 2378 (BH₄); anal. calcd for C₄₄H₆₄B₂N₄Ni₂ (787.98): C 67.06%, H 8.19%, N 7.11%; found: C 66.97%, H 8.28%, N 6.98%.

[Ni(6-Mes)(PPh₃)(NCBH₃)] (7)

1 (100 mg, 0.138 mmol) and NaBH₃(CN) (14 mg, 0.222 mmol) were placed in a J. Young's resealable ampoule in THF (10 mL). EtOH (1 mL) was added and the suspension was stirred for 5 min at room temperature to give a pale orange solution. The solution was reduced to dryness and the residue extracted into benzene (2×10 mL). Upon removal of the benzene, the orange residue was washed with EtOH (2 × 10 mL) to give 7 as a pale yellow solid. Yield: 50 mg (53%). Crystals suitable for X-ray diffraction were obtained by slow diffusion of hexane into a concentrated solution of 7 in THF at 238 K. ¹H NMR (500 MHz, C₆D₆, 298 K): δ 24.8 (br s), 12.4 (br s), 10.1 (s), 8.8 (br s), 5.5 (br s), 4.4 (br s), 3.9 (s), 1.6 (s); ¹¹B NMR (500 MHz, C₆D₆, 298 K): δ –20.2; $\mu_{\rm eff}$ (Evans method, THF, 298 K): 1.9 $\mu_{\rm B}$. Repeated attempts to determine elemental microanalysis on crystalline samples of the complex consistently gave low %C values (e.g. elemental analysis calcd (%) for C41H46N3PBNi: C, 72.28; H, 6.81; N, 6.16; found C, 69.98; H, 6.75; N, 5.77).

$[Ni(6-Mes)(PPh_3)(NPh)_2]$ (8)

1 (100 mg, 0.138 mmol), NaO^tBu (17 mg, 0.171 mmol), PPh₃ (36 mg, 0.138 mmol) and NHPh₂ (26 mg, 0.152 mmol) were placed in a J. Young's resealable ampoule in THF (10 mL). The suspension was stirred for 10 min at room temperature to give a dark red solution. The volatiles were removed in vacuo, the residue extracted into Et₂O (2 × 10 mL) and evaporated to dryness. The orange residue was washed with cold pentane $(2 \times 10 \text{ mL})$ to give 8 as a bright red solid. Yield: 80 mg (72%). Crystals suitable for X-ray diffraction were obtained by slow diffusion of pentane into a concentrated solution of 8 in diethyl ether at 238 K. 1 H NMR (500 MHz, C₆D₆, 298 K): δ 19.2 (br s), 8.2 (br s), 7.5 (br s), 7.1 (s), 6.9 (s), 6.8 (br s), 6.0 (br s), 5.1 (br s), 3.0 (br s), -15.6 (br s), -20.0 (s); μ_{eff} (Evans method, THF, 298 K): $2.0\mu_{\rm B}$. The sensitivity of the complex precluded all attempts to determine elemental microanalysis.

X-ray crystallography

Paper

Using Mo(K α) radiation, single crystals of compounds 2, 3 and 7 were analysed using an Agilent Xcalibur diffractometer, while datasets for 4 and 5, as well as 2a (vide infra; ESI \dagger only) were collected on a Nonius kappaCCD machine. An Agilent Supernova diffractometer was used to study 6, 8 and [Ni(6-Mes)(η^2 -C₆H₃Me₃)][B(C₆F₅)₄] (ESI \dagger only) using Cu(K α) radiation. All experiments were conducted at 150 K, with the exception of that for 6, which was achieved at 100 K. Details of the data collections and refinements are given in Table 2. The structures were uniformly solved using SHELXS, ⁴² and refined using full-matrix least squares in SHELXL⁴³ via the Olex-2⁴⁴ software suite. Only noteworthy refinement details follow.

A small amount of racemic twinning was accounted for in the refinement of 2a (ESI† only). This structure represents a P2₁ polymorph of compound 2, the latter solving in space group P2₁/c. In 3, the asymmetric was seen to contain one anion, one cation and one molecule of diethyl ether. While the cation and solvent were both ordered, disorder prevailed for four of the [BAr₄^F]⁻ trifluoromethyl substituents. In particular, the fluorine atoms attached to C88, C111 and C103 were each modelled over two sites in 65:35, 55:45 and 65:35 ratios, respectively, while the entire CF₃ group containing C87 exhibited 65:35 disorder. In 3 (and all subsequent structures containing disordered [BAr₄^F]⁻ trifluoromethyl groups) C-F and F...F distances within each disordered region were restrained to being similar in the final least squares. In addition, the ADPs for fractional occupancy atoms were also restrained, to assist convergence.

The cation in the asymmetric unit of 4 also fell prey to disorder. In particular, there is a 50:50 ratio of the tolyl-Ni-carbene moiety present *versus* the tolyl---pyrimidinium pair, the latter being stabilised by a C-H--- π interaction. In 5, the asymmetric unit was seen to comprise one cationic nickel containing species, one $[PF_6]^-$ anion and one THF molecule. The crystal was small, which contributed to weak diffraction at higher Bragg angles. Hence, data were truncated to a θ value of 24.7°.

The borohydride hydrogen atoms in the structure of compound 6 were readily located and refined with a common U_{iso} in each [BH₄] moiety. No distance restraints employed. C25 was modelled for 87:13 disorder, and the minor component of this atom was refined isotropically. A data collection was also performed on this compound, at room temperature (designated 6a, ESI†), in which the asymmetric unit was seen to consist of one half of a dimer molecule, wherein the metal centres and carbene carbon atoms were noted to coincide with a crystallographic 2-fold rotation axis. This necessarily means that the apical NHC carbons (C3 and C15) are each disordered in a 50:50 ratio. This disorder precluded addition of the hydrogen atoms bound to C2 and C14 using the riding model; hence, they were omitted from the refinement. The borohydride hydrogens were located, and refined without restraints, but their credibility is somewhat questionable given their associated U_{iso} values and the overall atomic displacement parameters. The reason for implementing a room temperature data collection for **6a** was to resolve a phase transition that arose in the course of a neutron experiment conducted on **6**, using VIVALDI, at the ILL. The rationale for doing a neutron experiment arose because, at 100 K, the borohydride moieties appeared to coordinate unsymmetrically to the nickel centres. Unfortunately, during cooling at the neutron source, the large crystals cracked. This ultimately resulted in collection of a neutron data set at room temperature, which suggested a different space group (C2/c) to that for the structure determined at 100 K using X-rays $(P2_1/c)$.

This phase transition, from a diffraction perspective, results in averaging the electron density that arises from the borohydrides across the sample and, overall, the ambient temperature neutron data did not afford any additional insight into the bonding subtleties which the experiment aimed to probe.

The asymmetric unit in 7 was seen host to one molecule of THF in addition to one molecule of the nickel complex. C3 in the latter was equally disordered over two sites, and the four chemically equivalent C–C distances involving C3/C3A were restrained to being similar in the final least squares. Three of the five atoms in the solvent were also refined to take account of 75:25 disorder. Once again, the chemically equivalent distances involving fractional occupancy atoms in this moiety were restrained to being similar, and ADP restraints were also incorporated to assist convergence.

In addition to one molecule of the complex, the asymmetric unit in 8 was noted to contain one molecule of guest diethyl ether.

Analysing the crystal structure of $[Ni(6\text{-Mes})(\eta^2-C_6H_3Me_3)][B(C_6F_5)_4]$ (ESI† only) was nothing short of excruciating, and it involved three data collections, some 350 refinements and approximately 10 data integrations. The issue appears to be that the compound is undergoing a phase transition. A more detailed treatment is presented in the ESI.†

Crystallographic data for all compounds have been deposited with the Cambridge Crystallographic Data Centre as supplementary publications CCDC 1578636–1578644, 1582301 and 1584193 for 2, 3, 4, 5, 6, 7, 8, 2a (ESI†), [Ni(6-Mes)(η^2 -C₆H₃Me₃)][B(C₆F₅)₄] (ESI†), 6a (ESI†) and [Ni(6-Mes)(PPh₃)(CO)Br][PF₆] (ESI†) respectively.

EPR spectroscopy

Samples for EPR measurements were prepared under an N_2 atmosphere in a glovebox. A solution of each complex was prepared by dissolving ca. 4 mg of 1–3, 5, 7 and 8 in 200 μ L of dry THF (in all cases, a small quantity of dry toluene was also added to improve the quality of the polycrystalline glass formed in frozen solution, and thereby enhance the quality of the EPR spectra). The solutions were transferred to an EPR tube, sealed in the glove box and then cooled to 77 K before rapid transfer to the pre-cooled EPR cavity. The X-band CW EPR measurements were performed on a Bruker EMX spectrometer utilizing an ER4119HS resonator, 100 kHz field modulation at 140 K.

Table 2 Crystal data and structural refinement details for compounds 2-8

	2	3	4	5	6	7	8
Empirical formula	C ₄₄ H ₅₁ F ₆ N ₂ NiOP ₂	C ₁₁₆ H ₁₀₈ BBrF ₂₄ N ₄ Ni ₂ OP ₂	C ₅₃ H _{36.5} BF ₂₀ N ₂ Ni _{0.5}	C ₄₅ H ₅₁ F ₆ N ₂ NiO ₂ P ₂	C ₄₄ H ₆₄ B ₂ N ₄ Ni ₂	C ₄₅ H ₅₄ BN ₃ NiOP	C ₅₆ H ₆₃ N ₃ NiOP
Formula weight	858.52	2300.14	1121.50	886.53	788.03	753.40	883.77
Crystal system	Monoclinic	Triclinic	Monoclinic	Monoclinic	Monoclinic	Monoclinic	Monoclinic
Space group	$P2_1/c$	$P\bar{1}$	C2/c	$P2_1/c$	$P2_1/c$	$P2_1/n$	$P2_1/n$
a/Å	16.0749(1)	12.9050(4)	31.5490(3)	14.3080(3)	14.17049(12)	14.2120(4)	18.99247(14)
b/Å	14.8045(1)	17.3278(5)	10.32800(10)	16.4910(3)	20.24524(15)	16.5650(4)	11.96944(9)
$c/ ext{Å}$	19.0391(1)	25.2732(6)	28.8100(3)	18.8500(4)	14.83697(15)	18.0386(6)	20.99341(16)
<i>α</i> /°	90	75.082(2)	90	90	90	90	90
β/°	111.435(1)	84.432(2)	92.977(1)	95.145(1)	91.0848(8)	108.838(3)	91.1551(7)
γ/°	90	87.444(2)	90	90	90	90	90
$U/\text{Å}^3$	4217.55(5)	5434.1(3)	9374.73(16)	4429.80(15)	4255.74(6)	4019.2(2)	4771.45(6)
Z	4	2	8	4	4	4	4
$\rho_{\rm calc}/{\rm g~cm}^{-3}$	1.352	1.406	1.589	1.329	1.230	1.245	1.230
μ/mm^{-1}	0.598	0.836	0.341	0.574	1.343	0.560	1.219
F(000)	1796.0	2364.0	4540.0	1852.0	1688.0	1604.0	1884.0
Crystal size/mm ³	$0.36\times0.31\times0.19$	$0.845 \times 0.77 \times 0.563$	$0.2 \times 0.1 \times 0.1$	$0.2 \times 0.1 \times 0.1$	$0.25\times0.25\times0.25$	$0.621 \times 0.378 \times 0.062$	$0.236 \times 0.157 \times 0.048$
2θ range for data collection/°	5.7 to 54.97	6.908 to 54.968	7.078 to 54.872	7.294 to 49.404	15.77 to 144.026	6.814 to 54.968	6.214 to 146.89
Index ranges	$-20 \le h \le 20$	$-16 \le h \le 16$	$-40 \le h \le 40$	$-16 \le h \le 16$	$-17 \le h \le 15$	$-13 \le h \le 18$	$-23 \le h \le 22$
	$-19 \le k \le 19$	$-22 \le k \le 17$	$-13 \le k \le 13$	$-19 \le k \le 19$	$-24 \le k \le 19$	$-21 \le k \le 20$	$-14 \le k \le 14$
	$-24 \le l \le 24$	$-32 \le l \le 32$	$-37 \le l \le 37$	$-22 \le l \le 22$	$-18 \le l \le 18$	$-23 \le l \le 22$	$-24 \le l \le 26$
Reflections collected	95 090	46 948	75 915	41 528	59 289	38 736	67 012
Independent reflections, $R_{\rm int}$	9656, 0.0282	24 033, 0.0376	10 665, 0.0610	7457, 0.0463	8330, 0.0847	9218, 0.0347	9580, 0.0510
Data/restraints/parameters	9656/0/511	24 033/121/1365	10 665/0/701	7457/128/565	8330/0/511	9218/25/502	9580/0/567
Goodness-of-fit on F^2	1.089	1.023	1.122	1.053	1.017	1.028	1.027
Final R_1 , w R_2 $[I > = 2\sigma(I)]$	0.0284, 0.0809	0.0635, 0.1396	0.0503, 0.1086	0.0435, 0.1070	0.0528, 0.1401	0.0472, 0.1128	0.0343, 0.0828
Final R_1 , w R_2 [all data]	0.0370, 0.0830	0.1297, 0.1745	0.0852, 0.1192	0.0555, 0.1154	0.0580, 0.1457	0.0703, 0.1267	0.0400, 0.0860
Largest diff. peak/hole/e Å ⁻³	0.60/-0.36	1.19/-0.93	0.51/-0.30	0.52/-0.48	0.68/-0.64	0.88/-0.94	0.39/-0.29

Computational details

All density functional theory (DFT) calculations were carried out with ORCA (version 4.0.0.2).45 The geometries were taken from crystallographic refinements, either optimising only the positions of the hydrogen atoms or fully relaxing the geometry. The geometry optimisations used the BP86 density functional,46 making use of the zeroth order relativistic correction ZORA retaining onecenter terms.⁴⁷ The scalar-relativistically recontracted versions of Ahlrich's triple-zeta quality basis sets (ZORA-def2-TZVP) were used on all atoms except carbon and hydrogen for which ZORA-def2-SVP basis sets were used. 48 The resolution of the identity (RI) approximation and the auxiliary basis SARC/J were used.49 The integration accuracy was increased to 7.0, the grid was set to 7 in ORCA nomenclature, and 'tight' SCF criteria were used. The optimisations considered solvent effects through the conductor-like polarisable continuum model, with the solvents as indicated in the experimental part. 50 Dispersion effects were taken into account with Grimme's D3BJ model including Becke-Johnson damping.51 Mulliken spin populations were inspected to confirm convergence to the targeted electronic structure.

Broken-symmetry DFT calculations used the functionals TPSSh, 52 B3LYP, 53 PBE0, 54 M06L, 55 additionally making use of the chain-of-spheres approximation (RIJCOSX) and using the 'flipspin' feature in ORCA to generate the initial guess for the broken-symmetry solution, with otherwise unchanged calculation setups.⁵⁶ The exchange coupling constants were taken directly from the ORCA output, using the definition by Yamaguchi.⁵⁷ For the calculation of EPR parameters, it was found that calculations with a different family of basis sets gave superior results. Generally, the IGLO-II basis set was used on all atoms, with CP for Ni and aug-pc-3 for Br, 58 in conjunction with the PBE0 density functional and the RIJCOSX approximation as for the BS-DFT calculations, making use of the AutoAux feature in ORCA. The grid sizes were set to Grid6 and GridX9 in ORCA nomenclature, with increased grids (7) on the Ni ion and all directly bound atoms as well as the nitrogen atoms in the carbene ligand. The spin-orbit mean field operator (SOMF(1X)) was used, and the origin for the g-tensor was taken at the centre of the electronic charge.⁵⁹ All tensor orientations, spin densities and molecular orbitals depicted and discussed in the main text and the ESI† are derived from calculations at this level of theory.

Conflicts of interest

There are no conflicts of interest to declare.

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