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Coligand role in the NHC nickel catalyzed C-F bond activation: investigations on the insertion of bis(NHC) nickel into the C-F bond of hexafluorobenzene†

Maximilian W. Kuntze-Fechner,^a Hendrik Verplancke, ^b Lukas Tendera,^a Martin Diefenbach,^b Ivo Krummenacher,^{ac} Holger Braunschweig, ^b ^{ac} Todd B. Marder, ^b ^{ac} Max C. Holthausen ^b *b and Udo Radius ^{ac} *a

The reaction of $[Ni(Mes_2Im)_2]$ (1) $(Mes_2Im = 1,3-dimesityl-imidazolin-2-ylidene)$ with polyfluorinated arenes as well as mechanistic investigations concerning the insertion of 1 and $[Ni(^iPr_2|m)_2]$ (1^{ipr}) $(^iPr_2|m)_2$ (1^{ipr}) diisopropyl-imidazolin-2-ylidene) into the C-F bond of C_6F_6 is reported. The reaction of 1 with different fluoroaromatics leads to formation of the nickel fluoroaryl fluoride complexes trans-[Ni(Mes₂Im)₂(F)(Ar^F)] $(Ar^F = 4-CF_3-C_6F_4 2, C_6F_5 3, 2,3,5,6-C_6F_4N 4, 2,3,5,6-C_6F_4H 5, 2,3,5-C_6F_3H_2 6, 3,5-C_6F_2H_3 7)$ in fair to good yields with the exception of the formation of the pentafluorophenyl complex 3 (less than 20%). Radical species and other diamagnetic side products were detected for the reaction of 1 with C_6F_{6} , in line with a radical pathway for the C-F bond activation step using 1. The difluoride complex trans- $[Ni(Mes_2lm)_2(F)_2]$ (9), the bis(aryl) complex trans- $[Ni(Mes_2lm)_2(C_6F_5)_2]$ (15), the structurally characterized nickel(i) complex trans-[Ni¹(Mes₂Im)₂(C₆F₅)] (11) and the metal radical trans-[Ni¹(Mes₂Im)₂(F)] (12) were identified. Complex 11, and related [Ni^I(Mes₂Im)₂(2,3,5,6-C₆F₄H)] (13) and [Ni^I(Mes₂Im)₂(2,3,5-C₆F₃H₂)] (14), were synthesized independently by reaction of trans-[Ni(Mes₂Im)₂(F)(Ar^F)] with PhSiH₃. Simple electron transfer from 1 to C_6F_6 was excluded, as the redox potentials of the reaction partners do not match and [Ni(Mes₂Im)₂]⁺, which was prepared independently, was not detected. DFT calculations were performed on the insertion of $[Ni(Pr_2|m)_2]$ (I^{ipr}) and $[Ni(Mes_2|m)_2]$ (1) into the C-F bond of C_6F_6 . For I^{ipr} , concerted and NHC-assisted pathways were identified as having the lowest kinetic barriers, whereas for 1, a radical mechanism with fluoride abstraction and an NHC-assisted pathway are both associated with almost the same kinetic barrier.

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Introduction

Fluorinated organic compounds have exceptional properties that are being exploited in many applications including materials, pharmaceuticals and agrochemicals. The development of methods to introduce fluorinated aromatic building blocks selectively into organic molecules is thus of fundamental interest in many areas of chemical research. One strategy for such transformations is the selective activation and subsequent

functionalization of C-F bonds of readily available fluoroorganic compounds such as fluoroaromatics. The challenge here is the selective cleavage of very stable C-F bonds.² We have recently established a protocol for the transformation of commercially available fluoroaromatics via a selective C-F defluoroborylation process to obtain polyfluorinated arylboronic esters,3 which may be further used in late stage functionalization, for example in Suzuki-Miyaura cross-coupling reactions.4 Defluoroborylation of polyfluoroaromatics can be achieved by a thermal $[Ni(Mes_2Im)_2]$ -catalyzed $(Mes_2Im = 1,3$ dimesityl-imidazolin-2-ylidene) transformation fluoroarenes into fluoroaryl boronic acid pinacol esters via C-F bond activation and transmetalation with bis(pinacolato) diboron (B₂pin₂) as the boron source (see Scheme 1).^{3a} Various arenes with different degrees of fluorination were converted into their corresponding boronate esters in this way. One particularly interesting finding of our study was that activation of the C-F bond by the nickel(0) complex is fast at ambient temperature. This step yields the oxidative addition product

^aInstitute for Inorganic Chemistry, Julius-Maximilians-Universität Würzburg, Am Hubland, 97074 Würzburg, Germany. E-mail: u.radius@uni-wuerzburg.de

^bInstitute for Inorganic and Analytical Chemistry, Goethe-Universität Frankfurt, Maxvon-Laue-Strasse 7, 60438 Frankfurt, Germany. E-mail: max.holthausen@chemie. uni-frankfurt.de

Institute for Sustainable Chemistry & Catalysis with Boron, Julius-Maximilians-Universität Würzburg, Am Hubland, 97074 Würzburg, Germany

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Scheme 1 Thermal borylation of fluoroarenes with B_2pin_2 mediated by $[Ni(Mes_2lm)_2]$ *via* the oxidative addition product *trans*- $[Ni(Mes_2-lm)_2(F)(Ar^F)]$ as the resting state of the catalysis.

trans-[Ni(Mes₂Im)₂(F)(Ar^F)] (Ar^F = fluoroaryl), which represents the resting state in the catalytic cycle. The subsequent defluor-oborylation step with B₂pin₂ is the rate determining step and requires elevated temperatures. A boryl complex trans-[Ni(Mes₂Im)₂(Bpin)(Ar^F)], a likely intermediate, was never observed and stoichiometric reactions of trans-[Ni(Mes₂Im)₂(-F)(Ar^F)] with B₂pin₂ led directly to the formation of Ar^F-Bpin. This finding implied that reductive elimination is very fast and that [Ni(Mes₂Im)_n(Bpin)(Ar^F)], once formed, will eliminate Ar^F-Bpin immediately (Scheme 1).^{5a}

As an alternative to the thermally-induced C-F bond activation and subsequent borylation of fluoroarenes, we have recently developed a process that employs visible-light photocatalysis, which has emerged as a powerful tool in organic synthesis.6 Our highly selective and general photocatalytic C-F borylation protocol3b employs a rhodium biphenyl complex as triplet sensitizer combined with the nickel catalyst [Ni(Mes₂-Im)₂ (1) for the C-F bond activation step and the defluoroborylation process. This Rh/Ni tandem catalyst system operates with visible light (400 nm) and achieves the highly selective borylation of a wide range of polyfluoroarenes with B2pin2 at room temperature in excellent yields. Both procedures, the thermal and photochemical defluoroborylation, work well for partially fluorinated aromatics but fail, or afford only low yields, perfluoroaromatics such as hexafluorobenzene octafluorotoluene.

Utilizing the dinuclear complex $[Ni_2(^iPr_2Im)_4(\mu-(\eta^2:\eta^2)-COD)]$ ($^iPr_2Im = 1,3$ -diisopropyl-imidazolin-2-ylidene) or the ethylene complex $[Ni(^iPr_2Im)_2(\eta^2-C_2H_4)]^8$ as sources of $[Ni(^iPr_2Im)_2]$ (1^{iPr}), we previously found that both readily undergo C-F bond insertion with a wide variety of per- and polyfluoroaromatics on a time scale suitable for catalysis (exemplarily shown for C_6F_6 in Scheme 2). 4c,k,9 Mechanistic investigations 9a of the insertion process were performed using

$$\bigcap_{\substack{|P_{\Gamma}|N\\|P_{\Gamma}|N\\|P_{\Gamma}|N\\|P_{\Gamma}|N\\|P_{\Gamma}|N}} \bigcap_{CH_{2}} \bigcap_{CH_{2}CH_{R}} \bigcap_{F} \bigcap_{I=1}^{F} \bigcap_$$

Scheme 2 Stoichiometric C–F bond activation of C_6F_6 using sources of $[Ni(^lPr_2|m)_2] \ 1^{lPr}$.

the ethylene complex $[Ni(^iPr_2Im)_2(\eta^2-C_2H_4)]$ as nickel precursor. Ethylene exchange at the $[Ni(^iPr_2Im)_2(\eta^2-C_2H_4)]$ complex with hexafluorobenzene and octafluoronaphthalene occurs at low temperatures (-80 °C and -30 °C, respectively; Scheme 2). Subsequent insertion reactions occur at higher temperatures (0 °C and 20 °C, respectively) to form the *trans*- $[Ni(^iPr_2Im)_2(-F)(C_6F_5)]$ and *trans*- $[Ni(^iPr_2Im)_2(F)(C_{10}F_7)]$ fluoroaryl fluoride complexes. We studied the C–F bond activation kinetics and, based on the decay rates of the octafluoronaphthalene complex $[Ni(^iPr_2Im)_2(\eta^2-C_{10}F_8)]$ determined by variable-temperature NMR spectroscopy, we derived an activation enthalpy of $\Delta H^{\ddagger} = 27.7 \pm 1.9 \text{ kcal mol}^{-1} (\Delta S^{\ddagger} = 8.8 \pm 6.0 \text{ cal K}^{-1} \text{ mol}^{-1}).$

We report herein on the reactivity of 1 with polyfluorinated arenes. We compare the results with those of earlier studies on C-F bond activation processes using nickel complexes with sterically less demanding NHCs, employing iPr instead of Mes substituents, i.e., using $[Ni(^{i}Pr_{2}Im)_{2}](\mathbf{1}^{ipr})$ as the nickel source. We demonstrate that the complex of the small NHC ligand ⁱPr₂Im favors a concerted oxidative addition proceeding through an $\eta^2(C,C)$ intermediate in reactions with fluoroarenes to yield trans-[Ni^{II}(NHC)₂(F)(Ar^F)] complexes, whereas the complex of the larger Mes₂Im ligand leads to fluorine atom abstraction to yield [NiI(NHC)2(F)] and a phenyl radical. For both mechanisms, competitive NHC-assisted pathways are found which account for the formation of diamagnetic products by a C-F bond activation step across the Ni-C_{NHC} bond. These NHC-assisted pathways play an important role for complexes of both sterically demanding and less bulky NHC ligands, and should thus be of general importance and widely applicable for the reactivity of NHC-stabilized transition metal complexes.

Results and discussion

C-F bond activation of fluoroaromatics

To gain insight into the C–F bond activation process using $[Ni(Mes_2Im)_2]$ (1), we first investigated stoichiometric reactions of perfluorotoluene, perfluorobenzene, perfluoropyridine and the partially fluorinated arenes pentafluorobenzene, 1,2,3,5-tetrafluorobenzene and 1,3,5-trifluorobenzene with 1 (see Scheme 3). We monitored the reactions by 1H and $^{19}F\{^1H\}$ NMR spectroscopy and observed a significant effect of the degree of fluorination on both reaction rate and yield. Reactions of 1 with hexafluorobenzene and octafluorotoluene proceed within seconds at room temperature, whereas the reactions with tetra-and pentafluorobenzene take minutes to complete. With 1,3,5-trifluorobenzene, full conversion of 1 takes weeks at room temperature (see ESI, Fig. S1†), but can be accelerated at 80 $^{\circ}$ C in thf to reach completion after 5 days.

These reactions can be performed in thf, toluene or hexane at room temperature and lead, in each case, to the insertion of the nickel complex into the C-F bond of the fluoroarene to form the nickel fluoroaryl fluoride complexes *trans*-[Ni(Mes₂Im)₂(-F)(Ar^F)] (Ar^F = 4-CF₃-C₆F₄ 2, C₆F₅ 3, 2,3,5,6-C₅F₄N 4, 2,3,5,6-C₆F₄H 5, 2,3,5-C₆F₃H₂ 6, 3,5-C₆F₂H₃ 7) in fair to good isolated yields. Notably, however, the reaction with C₆F₆ yields less than 20% of the pentafluorophenyl complex 3. Higher temperature, different solvents (thf, toluene, hexane) or added [NMe₄]F does

Scheme 3 The reactions of $[Ni(Mes_2lm)_2]$ (1) with (a) octafluorotoluene, (b) hexafluorobenzene, (c) perfluoropyridine, (d) pentafluorobenzene, (e) 1,2,3,5-tetrafluorobenzene and (f) 1,3,5-trifluorobenzene to give the complexes trans- $[Ni(Mes_2lm)_2(F)(4-CF_3-C_6F_4)]$ (2), trans- $[Ni(Mes_2lm)_2(F)(C_6F_5)]$ (3), trans- $[Ni(Mes_2lm)_2(-F)(2,3,5,6-C_6F_4N)]$ (4), trans- $[Ni(Mes_2lm)_2(-F)(2,3,5,6-C_6F_4N)]$ (5), trans- $[Ni(Mes_2lm)_2(-F)(2,3,5,6-C_6F_4N)]$ (6) and trans- $[Ni(Mes_2lm)_2(-F)(3,5-C_6F_2H_3)]$ (7), respectively. Isolated yields are given.

not seem to affect the yield of the insertion product 3. Complexes 2–7 were characterized by elemental analysis, 1 H, 19 F 1 H 1 H 10 A and 13 C 1 H 10 NMR spectroscopy (see ESI 1). In the 19 F 1 H 10 NMR spectra of these complexes, the resonances of the nickel-bound fluoride ligand were observed in the typical range between -361.9 and -333.1 ppm. Within the series presented (see ESI, Table S1 1), the NMR shift of this resonance depends on the degree of fluorination of the fluoroaryl ligands, *i.e.*, an increase of the degree of fluorination of the aryl ligand leads to an upfield shift of the Ni–F resonance.

Crystals of 3, 4, and 5 suitable for X-ray diffraction were obtained from saturated solutions of these compounds either in pentane or hexane at -30 °C (Fig. 1, Table 1; see also ESI Fig. S34-S36 and Table S2†). The crystal structure of 6 was published previously.3a All complexes of the type trans- $[Ni(Mes_2Im)_2(F)(Ar^F)]$ $(Ar^F = C_6F_5 \ 3, \ 2,3,5,6-C_5F_4N \ 4, \ 2,3,5,6-C_5F_6N \$ C₆F₄H 5, 2,3,5-C₆F₃H₂ 6) adopt a square planar structure with a trans arrangement of the NHC ligands. An increasing degree of fluorination of the fluoroaryl ligand leads to a slight shortening of the Ni-F bond lengths (Ni-F: 6: 1.874(2) Å, 5: 1.856(2) Å, 4: 1.859(2) Å, 3: 1.844(2) Å), while the distances of the nickel center to the fluoroaryl ligand become gradually longer (Ni-C3: 6: 1.854(5) Å, 5: 1.896(3) Å, 4: 1.883(3) Å, 3: 1.944(5) Å). We assume that both the upfield shift of the Ni-F ¹⁹F NMR resonance and the shortening of the Ni-F bond lengths with increasing degree of aryl fluorination are indications of stronger Ni-F bonding.

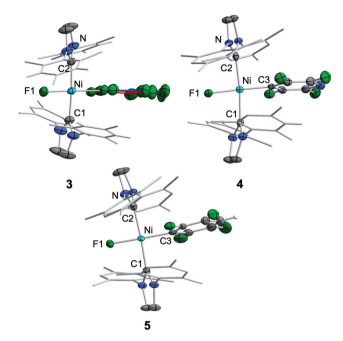


Fig. 1 Molecular structures of trans-[Ni(Mes₂Im)₂(F)(C₆F₅)] (3) (top left), trans-[Ni(Mes₂Im)₂(F)(2,3,5,6-C₅F₄N)] (4) (top right) and trans-[Ni(Mes₂Im)₂(F)(2,3,5,6-C₆F₄H)] (5) (bottom) in the solid state (ellipsoids drawn at the 50% probability level). Hydrogens atoms, with exception of the proton at the fluoroaromatic of 5, are omitted for clarity.

As the low yield of trans- $[Ni(Mes_2Im)_2(F)(C_6F_5)]$ (3) is in sharp contrast with the results we obtained previously for the reaction of $[Ni_2]^i Pr_2 Im]_4 (\mu - (\eta^2 : \eta^2) - COD)$ or $[Ni]^i Pr_2 Im]_2 (\eta^2 - C_2 H_4)$ with C₆F₆, we decided to take a closer look at the corresponding reaction using [Ni(Mes₂Im)₂] (1). Performing the stoichiometric reaction of 1 with C₆F₆ in an NMR tube in C₆D₆ led to an immediate color change from dark-violet, the color of concentrated complex 1, to orange after addition of C₆F₆ at room temperature. A quantitative conversion of 1 was achieved after 5 min as monitored by ¹H NMR spectroscopy (see ESI, Fig. S2†). However, the spectroscopic yield determined by 19F{1H} NMR spectroscopy after 5 min at room temperature, vs. a Ph-F containing capillary as internal standard, revealed the formation of 3 in approximately 17% yield and, in addition, the formation of small amounts of fluoride-containing side products (see ESI, Fig. S3†). Even after 72 h at room temperature, no increase in the spectroscopic yield of 3 was observed. In further control experiments, neither the use of an excess of 1 (2.85 equiv.) nor C_6F_6 (2.5 equiv.) increased the yield of 3 substantially. These experiments demonstrate that the low isolated yield of 3 is not a problem of the isolation process for this complex, but rather an intrinsic problem associated with its formation and the C-F bond activation step. Low temperature NMR experiments (-50 °C to +20 °C) revealed that a nickel fluoride resonance at -358 ppm appeared for this reaction in the $^{19}F\{^{1}H\}$ NMR spectrum already at -50 °C (see ESI, Fig. S4†), but also that, at these temperatures, all resonances are significantly broadened in the ¹H NMR spectrum of the reaction mixture (see ESI, Fig. S5†). Although we previously observed some line

Table 1 Crystallographic data for compounds 1, 3, 4, 5, 6, 8, 9, 11, 13, 14, [Ni^I(6-Mes)₂][Br]^{11a} and [Ni^I(P^IPr₃)₂(C₆F₅)]²²

	d Ni–C1/C2	d Ni–C3 $_{(ArF)}$	d Ni-F	∠C1–Ni–C2	\angle NHC(C1): NHC(C2)
$[Ni(Mes_2Im)_2] 1$	1.827(6)	_	_	176.4	53.0
E (= /=3	1.830(6)				
3	1.923(3)	C3: 1.882(7)	1.844(2)	175.6(1)	37.07(2)
	1.922(3)	C3': 1.944(5)			
4	1.923(3)	1.883(3)	1.859(2)	174.3(1)	36.01(2)
	1.920(3)				
5	1.921(2)	1.896(3)	1.856(2)	176.4(1)	33.81(1)
	1.924(2)				
6	1.912(3)	1.854(5)	1.874(2)	176.7(1)	31.65(2)
	1.912(3)				
8	1.894(3)	_	_	174.5(1)	57.99(1)
	1.894(3)				
9	1.903(3)	_	F1: 1.845(2)	178.5(1)	53.34(1)
	1.902(3)		F2: 1.823(2)		
11	1.923(2)	1.984(3)	_	159.8(8)	82.37(1)
	1.923(2)				
13	1.930(2)	1.987(3)	_	157.3(8)	82.11(1)
	1.930(2)				
14	1.918(1)	C3: 1.869(1)	_	159.5(5)	82.46(8)
	1.917(1)	C3': 2.046(1)			
[Ni ^I (6-Mes) ₂][Br]	1.939(3)	_	_	179.3(1)	57.99(1)
	1.941(3)				
$[\mathrm{Ni}^{\mathrm{I}}(\mathrm{P}^{\mathrm{i}}\mathrm{Pr}_{3})_{2}(\mathrm{C}_{6}\mathrm{F}_{5})]$	P1: 2.243(5)	1.973(2)	_	P1-Ni-P2	_
	P2: 2.233(5)			145.2(2)	

broadening for the *N*-alkyl groups of the related complex *trans*-[Ni($^{\rm i}$ Pr $_2$ Im) $_2$ (F)(C $_6$ F $_5$)], 9a which arose due to hindered rotation of the NHC ligand about the Ni–C axis, all resonances observed for the reaction of 1 with C $_6$ F $_6$ are involved in the broadening. This led to the assumption that radical species are involved in the process. Subsequent EPR experiments were performed at $-203~^{\circ}$ C for the reaction of 1 with C $_6$ F $_6$ which confirmed the presence of metal-centered radicals in the mixture.

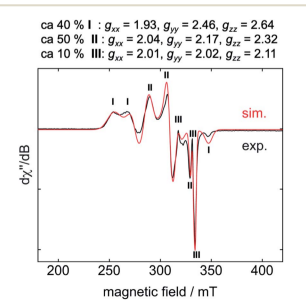


Fig. 2 EPR spectrum (-203 °C) of the reaction mixture of 1 with C_6F_6 after 5 s at -78 °C in thf.

For EPR spectroscopic investigations, **1** and C_6F_6 were combined in an EPR tube with thf at -78 °C and the sample was frozen immediately in liquid nitrogen. The EPR tube containing the frozen reaction mixture was transferred to the cooled EPR cavity at -203 °C and a spectrum was recorded. The resulting EPR spectrum displays a superposition of resonances of three different products, of which **I** and **II** represent the two dominant species (Fig. 2, **I**: 40%, **II**: 50%, **III**: 10%).

Cyclic voltammetry results exclude a simple electron transfer from 1 to C_6F_6 as the origin of radical generation in the reaction mixture (see ESI, Fig. S6†), as 1 shows a reversible oxidation/reduction associated with a redox potential of -2.03 V for the redox-couple Ni^0/Ni^I , and an irreversible oxidation at 0.14 V for the redox-couple Ni^I/Ni^{II} . Although the reduction of C_6F_6 at -2.87 V is irreversible, we exclude simple one electron transfer because of the large separation of 0.84 V.

For further scrutiny, complex **1** was oxidized by adding ferrocenium tetrafluoroborate in thf at room temperature to a suspension of **1** in thf. A few min after addition of the ferrocenium salt the metal-centered radical [Ni(Mes₂Im)₂][BF₄] (8) precipitated as an off-white solid (83% isolated yield, Scheme 4), which is only sparingly soluble in common organic solvents. The Ni^I complex **8** was characterized by ¹¹B{¹H} and ¹⁹F{¹H} NMR spectroscopy in acetonitrile (decomposition occurs after

$$\underbrace{ \begin{bmatrix} \text{Mes} \\ \text{N} \\ \text{N} \end{bmatrix} }_{\text{Mes}} \underbrace{ \begin{bmatrix} \text{Mes} \\ \text{Fe} \\ \text{N} \end{bmatrix} }_{\text{Mes}} + \underbrace{ \begin{bmatrix} \text{Mes} \\ \text{Fe} \\ \text{RT} \end{bmatrix} }_{\text{RT}} \underbrace{ \begin{bmatrix} \text{Mes} \\ \text{N} \\ \text{N} \end{bmatrix} }_{\text{N}} \underbrace{ \begin{bmatrix} \text{Mes} \\ \text{Mes} \\ \text{N} \end{bmatrix} }_{\text{N}} \underbrace{ \begin{bmatrix} \text{Mes} \\ \text{N} \end{bmatrix} }_{\text{RT}} \underbrace{ \begin{bmatrix} \text{Mes} \\ \text{N} \\ \text{N} \end{bmatrix} }_{\text{N}} \underbrace{ \begin{bmatrix} \text{Mes} \\ \text{N} \end{bmatrix} }_{\text{N}} \underbrace{ \begin{bmatrix} \text{M$$

Scheme 4 Synthesis of [Ni(Mesalm)a][BFa] (8)

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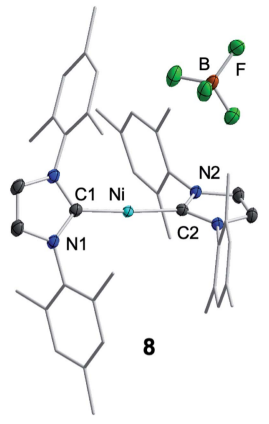


Fig. 3 Molecular structure of $[Ni(Mes_2Im)_2][BF_4]$ (8) in the solid state (ellipsoids drawn at the 50% probability level). Hydrogens atoms are omitted for clarity.

some time) and IR spectroscopy, elemental analysis and high-resolution mass spectroscopy. The $^{11}\mathrm{B}\{^1\mathrm{H}\}$ and $^{19}\mathrm{F}\{^1\mathrm{H}\}$ NMR spectra revealed an intact counter anion $[\mathrm{BF_4}]^-$ (see ESI, Fig. S64†). Crystals of 8 suitable for X-ray diffraction (Fig. 3, Table 1; see also ESI, Table S2 and Fig. S37†) were obtained by slow evaporation of a saturated solution of 8 in a 1 : 1 toluene/ethanol mixture under an argon atmosphere at room temperature. The X-ray crystal structure reveals a nearly linear alignment of the NHC ligands with slightly elongated Ni–C distances compared to those of the starting material 1.

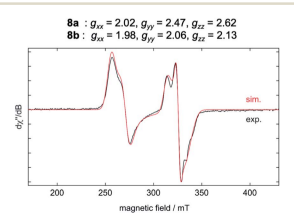


Fig. 4 EPR spectrum of 8 in the solid state at -203 °C with NBu₄Br.

The results of the EPR spectroscopic investigations¹⁰ performed on solid-state samples of [Ni(Mes₂Im)₂][BF₄] (8) are shown in Fig. 4. The general insolubility of 8 precluded determination of its magnetic moment by the Evans method. The spectrum reveals two sets of signals, i.e., 8a ($g_{xx} = 2.02$, $g_{yy} =$ 2.47, $g_{zz} = 2.62$; 70%) and **8b** ($g_{xx} = 1.98$, $g_{yy} = 2.06$, $g_{zz} = 2.13$; 30%), both in line with nickel-centered radicals. Note that in previous studies of two other homoleptic two-coordinate cationic d^9 -nickel(I) complexes, [Ni(6-Mes)₂][Br] (6-Mes = 1,3bis(2,4,6-trimethylphenyl)-3,4,5,6-tetrahydropyrimidin-2ylidene) and $[Ni(P^tBu_3)_2][Al(OC(CF_3)_3)_4]$, no EPR signals were observed.11 To obtain further insight, we performed density functional theory (DFT) calculations on the molecular geometries, electronic structures and EPR parameters (g tensors) of a variety of potential candidates for 8 (PBE0-D/pcSseg-2, see the ESI for details†).12

The DFT-optimized, D_2 -symmetric geometry of the $[Ni(Mes_2Im)_2]^+$ radical cation of 8 agrees very well with the X-ray structure (Fig. 5; *e.g.*, $d_{Ni-C} = 1.89$ Å, exp: 1.894(3) Å). In the 2A electronic ground state, the spin density is localized at the metal center, with the unpaired electron residing in an s/d_2 -type orbital (Fig. 5a).

The calculated g values for the radical cation $[Ni(Mes_2Im)_2]^+$ ($g_{xx} = 2.01$, $g_{yy} = 2.65$, $g_{zz} = 2.98$), computed under gas-phase conditions, strongly differ from the experimental data with a maximum deviation of 0.36 (8a) and 0.85 (8b; see Table 2 and ESI, Table S3†). However, computations in the presence of the counter ion result in further structural motifs with impact on

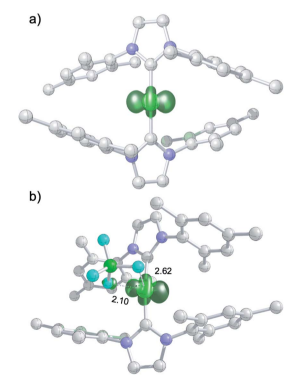


Fig. 5 (a) Spin density plot for $[Ni(Mes_2lm)_2]^+$; (b) molecular structure of 8_{DFT}^{1} showing $Ni-F^{BF_4-}$ contacts (isovalue $\pm~0.0075~a_0^{-3}$; lengths of Ni-F contacts in Å; hydrogen atoms not shown).

Table 2 Experimental and DFT calculated g tensors for species 8

		g tensor components					
Compound	g_{xx}	g_{yy}	g_{zz}				
8a	Exp. (solid state)	2.02	2.47	2.62			
8b	Exp. (solid state)	1.98	2.06	2.13			
[Ni(Mes ₂ Im) ₂] ⁺ (gas phase)	DFT	2.01	2.65	2.98			
8 _{DFT} ^{1a}	DFT	2.03	2.50	2.59			
a DFT-optimized structure with Ni– $F^{\mathrm{BF_4}^-}$ contacts.							

the computed g tensors (see ESI, Fig. S7†). A Ni–F contact with the counter ion in $\mathbf{8_{DFT}}^1$ (Table 2, entry 4, Fig. 5b) results in g tensor components closely corresponding to those of $\mathbf{8a}$ (maximum deviation: 0.03), while no species matching the EPR parameters of $\mathbf{8b}$ were identified in our computational exploration. However, none of the EPR signatures detected for the electrochemically-formed complex $\mathbf{8a}$ appeared during the reaction of $\mathbf{1}$ with C_6F_6 (Fig. 2) and, in light of our CV results, it is unlikely that the $[Ni(Mes_2Im)_2]^+$ cation is involved here.

We then focused on identifying the byproducts of the reaction of 1 with C₆F₆. Stoichiometric reaction of 1 with C₆F₆ in thf overnight at room temperature led to a very small amount of a dark-green precipitate which was removed by filtration. After removal of all volatiles from the filtrate, the residue was washed with a large amount of hexane to extract the C-F bond activation product. The yellow residue, which remained after washing, was identified as the difluoride complex trans-[Ni(Mes₂Im)₂(F)₂] (9) by elemental analysis, X-ray diffraction and ¹H, ¹⁹F{¹H} and ¹³C {1H} NMR spectroscopy (see ESI†). Most significantly, the fluoride resonance, detected as a singlet at -560 ppm in the 19 F $\{^{1}$ H $\}$ NMR spectrum, is shifted ca. 200 ppm to higher field compared to those of the mono-fluoride complexes 2-7 (-333 ppm to -362 ppm, vide supra). A similar high-field shifted fluoride resonance was also observed for the phosphine-stabilized platinum complex [Pt(PiPr₃)₂(F)₂] (-455.9 ppm) compared to

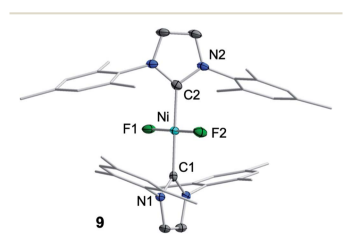


Fig. 6 Molecular structure of trans-[Ni(Mes₂Im)₂(F)₂] (9) in the solid state (ellipsoids drawn at the 50% probability level). Hydrogens atoms are omitted for clarity.

Scheme 5 Synthesis of $[Ni(Mes_2Im)_2(I)_2]$ (10) and $[Ni(Mes_2Im)_2(F)_2]$ (9).

[Pt(PPh₃)₂(F)(C₆H₅)] (-107.6 ppm).¹³ Crystals of **9** suitable for X-ray diffraction (Fig. 6, Table 1; see also ESI, Table S2 and Fig. S38†) were obtained after storing a saturated solution of the complex at room temperature in C₆D₆. Crystallographic analysis revealed a square planar coordination environment about the Ni^{II} center with a *trans*-arrangement of NHC and fluoride ligands.

An independent sample of complex **9** was synthesized in 38% yield by fluorination of [Ni(Mes₂Im)₂(I)₂] (**10**) using an excess (2.5 equiv.) of silver(1) fluoride in CH_2Cl_2 at 0 °C (Scheme 5). Complex **10** was synthesized by reaction of **1** with I₂, isolated in 80% yield and characterized by elemental analysis, and 1H and $^{13}C\{^1H\}$ NMR spectroscopy (see ESI†). Interestingly, the resonance of the carbene carbon atoms is almost unaffected by substitution of the fluoride by the more electropositive

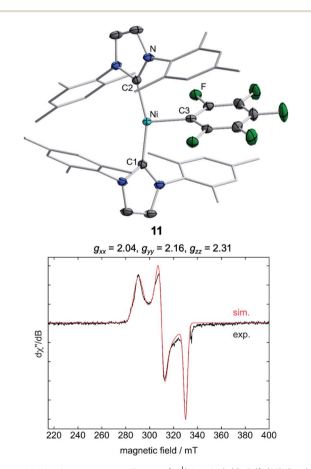


Fig. 7 Molecular structure of trans-[Ni¹(Mes₂Im)₂(C₆F₅)] (11) (top) in the solid state (ellipsoids drawn at the 50% probability level) and EPR spectrum at $-203\,^{\circ}$ C of the isolated compound 11 (bottom). Hydrogen atoms are omitted for clarity.

iodide ligand, and was detected at 176.5 ppm (cf. [Ni(Mes₂-

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 $[Im]_2(F)_2$ (9): 174.6 ppm).

Thus, $[Ni(Mes_2Im)_2(F)_2]$ (9) was clearly identified as one of the side products of the reaction of 1 with C_6F_6 . This complex is formed in low yield (17%) but in an amount similar to that of the insertion product trans- $[Ni(Mes_2Im)_2(F)(C_6F_5)]$ (3). The amounts of complexes 9 and 3 total ca. 40% when the reaction of 1 with C_6F_6 is performed at room temperature, and thus the majority of the products formed in this reaction is still unaccounted for.

Storing the concentrated hexane mother liquor of the extract from the isolation of 9 (vide supra) for 3 days at -30 °C led to crystallization of the remaining C-F bond insertion product trans-[Ni(Mes₂Im)₂(F)(C₆F₅)] (3) and a novel nickel(1) complex trans- $[Ni^{I}(Mes_{2}Im)_{2}(C_{6}F_{5})]$ (11) as yellow (3) and orange (11) crystals, respectively, which were manually separated in a glovebox (see ESI, Fig. S8†). The paramagnetic compound 11 was characterized by elemental analysis, EPR spectroscopy and Xray diffraction. Determination of the room-temperature magnetic moment of 11 in solution (Evans method) gave a μ_{eff} value of 1.80 $\mu_{\rm B}$, which is consistent with the presence of one unpaired electron. The molecular structure (Fig. 7, top, Table 1; see also ESI, Table S2 and Fig. S39†) and the EPR spectrum (Fig. 7, bottom) of 11 confirm that this complex is a threecoordinate nickel(1) radical. Simulation of the EPR spectrum of 11 gave a g tensor of $g_{xx} = 2.04$, $g_{yy} = 2.16$ and $g_{zz} = 2.31$, which was also observed in the EPR spectrum of the crude reaction mixture of 1 and C_6F_6 (Fig. 2). With the experimentally obtained g tensors and the molecular structure of the radical

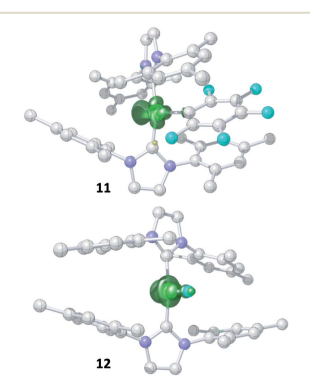


Fig. 8 Spin density plots for trans-[Ni¹(Mes₂Im)₂(C₆F₅)] (11) (top) and trans-[Ni¹(Mes₂Im)₂(F)] (12) (bottom) (isovalue 0.0075 a_0^{-3} ; hydrogen atoms are omitted for clarity).

Scheme 6 One-electron oxidative addition of C_6F_6 to $[Ni(Mes_2lm)_2]$ (1) to yield the metal radicals $trans-[Ni^l(Mes_2lm)_2(C_6F_5)]$ (11) and $trans-[Ni^l(Mes_2lm)_2(F)]$ (12).

species **11** in hand, we carried out computational studies of the electronic properties of complex **11** and a likely radical counterpart from the reaction of **1** and C_6F_6 , $[Ni^I(Mes_2Im)_2(F)]$ (**12**) (Fig. 8). Both complexes **11** and **12** would be the result of a one-electron oxidative addition reaction of two equiv. of **1** with one equiv. C_6F_6 (Scheme 6).

Molecular geometries, electronic structures and EPR parameters (g tensors) were thus calculated for the metal radicals trans-[Ni^I(Mes₂Im)₂(C₆F₅)] (11) and trans-[Ni^I(Mes₂Im)₂(F)] (12) (Fig. 8) in order to connect the experimentally observed EPR spectra from the reaction mixture of 1 and C₆F₆ (Fig. 2), the EPR spectra of the isolated compound 11, and the corresponding isotropic g tensor components with the assigned structure of 11 (Fig. 7).

According to DFT calculations, complexes **11** and **12** are C_2 -symmetric doublet ground state species. The spin density is located at the metal center and the unpaired electron resides in an s/d₂-type orbital, yielding ²A electronic ground states (Fig. 8). Calculated and experimental g tensor components are in good agreement for species **11**, with a maximum difference of 0.03 in g_{zz} . With the largest deviation being 0.08 for **12**, the agreement is still reasonable (Table 3).

To provide further evidence for the existence of *trans*- $[Ni^I(Mes_2Im)_2(C_6F_5)]$ (11) and *trans*- $[Ni^I(Mes_2Im)_2(F)]$ (12), we attempted to synthesize these complexes independently. The

Table 3 Comparison of experimental and calculated g tensors for species ${\bf 11}$ and ${\bf 12}$

		g tensor components b		
Compound	DFT/Exp ^a	g_{xx}	g_{yy}	g_{zz}
11	Exp. (isol.) ^c	2.04	2.16	2.31
	Exp. (react. mix.) d	2.04	2.17	2.32
	DFT	2.06	2.17	2.29
12	Exp. (react. mix.) d	1.93	2.46	2.64
	DFT	2.01	2.42	2.57

^a The experimental *g*-tensor components are reorganized in ascending order from g_{xx} to g_{zz} . ^b EPR parameter have been calculated using DFT. The calculated values are rounded to match the number of digits of the experimental values. ^c Exp. (isol.): see Fig. 7 (bottom). ^d Exp. (react. mix.): see Fig. 2 I and II (I corresponds to compound 12; II corresponds to compound 11).

Scheme 7 Synthesis of the metal radicals $[Ni^{1}(Mes_{2}Im)_{2}(C_{6}F_{5})]$ (11), $[Ni^{1}(Mes_{2}Im)_{2}(2,3,5,6-C_{6}F_{4}H)]$ (13) and $[Ni^{1}(Mes_{2}Im)_{2}(2,3,5-C_{6}F_{3}H_{3})]$ (14).

reaction of [Ni(Mes₂Im)₂][BF₄] (8) with CsF led to a mixture of two complexes, which we were not able to separate. One of them was identified via ¹⁹F{¹H} NMR spectroscopy as trans-[Ni(Mes₂Im)₂(F)₂] (9) (¹⁹F{¹H} NMR resonance at -560 ppm), and the resulting mixture reveals an EPR resonance with g tensors ($g_{xx} = 2.05$, $g_{yy} = 2.42$, $g_{zz} = 2.61$) which are close to the g-tensors calculated for trans-[Ni^I(Mes₂Im)₂(F)] (12). We are thus confident that the second metal radical obtained in the reaction mixture is the monofluoride complex trans-[Ni^I(Mes₂Im)₂(F)] (12).

The complex trans-[Ni^I(Mes₂Im)₂(C₆F₅)] (11) as well as related trans-[Ni^I(Mes₂Im)₂(2,3,5,6-C₆F₄H)] (13) and trans-[Ni^I(Mes₂Im)₂(2,3,5-C₆F₃H₂)] (14) can be synthesized from the reaction of trans-[Ni(Mes₂Im)₂(F)(Ar^F)] (Ar^F = C₆F₅ 3, 2,3,5,6-C₆F₄H 5, 2,3,5-C₆F₃H₂ 6) with PhSiH₃ (Scheme 7, see also ESI Fig. S9 and S10†).¹⁴

The metal radicals were characterized by elemental analysis, IR and EPR spectroscopy as well as single-crystal X-ray diffraction. All compounds are stable in the solid state as well as in solution for several days. If the reactions are performed in an NMR tube and followed by ¹H and ¹⁹F{¹H} NMR spectroscopy (see ESI; Fig. S9 and S10†), the resonances for the Mes₂Im, pentafluorophenyl and fluoride ligands vanish, indicating the formation of a paramagnetic species. For complexes of the type trans-[Ni(NHC)₂(H)(Ar^F)], we expect hydride resonances in the region of ca. −13 ppm in the ¹H NMR spectrum, ^{9b,d} and a strong absorption in the IR spectrum in the region between 1600 and 2200 cm $^{-1}$ 15 (we expect the Ni-H stretch to be at ca. 1850 cm $^{-1}$ based on DFT calculations). However, such signals were absent for 11, 13 and 14. Thus, although complexes of the type trans-[Ni^I(Mes₂Im)₂(Ar^F)] cannot easily be distinguished from the corresponding hydride complexes trans-[Ni^I(Mes₂Im)₂(H)(Ar^F)] by X-ray diffraction (see below), we are confident that 11, 13 and 14 are the metal radicals. Crystals of trans-[Ni^I(Mes₂Im)₂(C₆F₅)] (11), trans-[Ni^I(Mes₂Im)₂(2,3,5,6-C₆F₄H)] (13) and trans- $[Ni^{1}(Mes_{2}Im)_{2}(2,3,5-C_{6}F_{3}H_{2})]$ (14) suitable for X-ray diffraction (Fig. 9, Table 1; see also ESI Table S2 and Fig. S39-S41†) were obtained by storing saturated solutions of these compounds either in pentane or hexane at -30 °C. Complexes 11-13 adopt a distorted T-shaped structure, in which the NHC ligands occupy mutually trans positions. Due to the absence of the fluoride ligand, 11, 13 and 14 exhibit shortened Ni-C distances to the fluoroaryl ligand and reduced C1-Ni-C2 angles compared to nickel(II) complexes 3, 4, 5 and 6, which is also a further

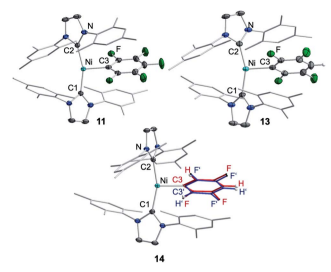


Fig. 9 Molecular structures of trans-[Ni¹(Mes₂Im)₂(C_6F_5)] (11) (top left), trans-[Ni¹(Mes₂Im)₂(2,3,5,6- C_6F_4 H)] (13) (top right) and trans-[Ni¹(Mes₂Im)₂(2,3,5- C_6F_3 H₂)] (14) (bottom) in the solid state (ellipsoids drawn at the 50% probability level). Hydrogens atoms (with exception of the protons at fluoroarene rings) are omitted for clarity. Because of disorder of the fluoroaryl ligand of 14, the ligand is represented by a ball and stick model in two different colors for clarity.

indication of the absence of a metal hydride. The data is in line with the data observed for $[Ni^I(P^iPr_3)_2(C_6F_5)]$ reported by Johnson and co-workers previously (Table 1, see also ESI Table S2†). EPR spectra of compounds 11, 13 and 14 were recorded in frozen thf solutions and reveal similar g tensors for the complexes, which are in good agreement with the calculated parameters (see ESI, Fig. S11–S13 and Table S4†).

Thus, the reaction of 1 with C₆F₆ affords the insertion product trans-[Ni(Mes₂Im)₂(F)(C₆F₅)] (3) in approximately 20% isolated yield, the difluoride complex trans-[Ni(Mes₂Im)₂(F)₂] (9) in approximately 17% isolated yield, the three-coordinate nickel(1) metal radicals trans-[Ni^I(Mes₂Im)₂(C₆F₅)] (11) (isolated yield: 10%), trans-[Ni^I(Mes₂Im)₂(F)] (12) (not isolated), and a small amount of a decomposition product, i.e., a dark green precipitate which was not characterized. Trans-[Ni^I(Mes₂Im)₂(-F)] (12) was not isolated; it was only observed in the EPR spectra after 5 s at -78 °C, and the signals vanish after about 10 s during the course of the reaction. Further investigation of the hexane mother liquor of the reaction of 1 and C₆F₆ revealed that the bis(aryl) nickel(II) complex $[Ni(Mes_2Im)_2(C_6F_5)_2]$ (15) remains in solution and was identified in the reaction mixture by ¹⁹F{¹H} NMR spectroscopy. The radical species **11** and **12** were identified by EPR spectroscopy in a frozen thf solution at -78 °C (Fig. 2). The diamagnetic products trans-[Ni(Mes₂- $\operatorname{Im}_{2}(F)(C_{6}F_{5})$] (3), trans- $\left[\operatorname{Ni}(\operatorname{Mes}_{2}\operatorname{Im})_{2}(F)_{2}\right]$ (9), and trans- $[Ni(Mes_2Im)_2(C_6F_5)_2]$ (15) were identified by NMR spectroscopy (see Fig. S14 of the ESI†).

To expand our study to less fluorinated systems, we reacted **1** with pentafluorobenzene. After 48 h at room temperature, the $^{19}F\{^1H\}$ and ^{19}F NMR spectra recorded in C_6D_6 reveal the formation of the C–F bond activation product *trans*-[Ni(Mes₂-Im)₂(F)(C₆F₄H)] (5), the nickel difluoride complex [Ni(Mes₂-Im)₂(F)₂] (9), and the corresponding bis(aryl) nickel(II) complex

[Ni(Mes₂Im)₂(C₆F₄H)₂] (see ESI, Fig. S15†). Furthermore, an EPR spectrum of the frozen reaction mixture of **1** with pentafluorobenzene in thf recorded after 5 s at -78 °C (see ESI, Fig. S16†) revealed resonances for three different products, one of which is in accordance with *trans*-[Ni¹(Mes₂Im)₂(F)] (**12**) and another has the same *g* tensor as observed for isolated [Ni¹(Mes₂Im)₂(C₆F₄H)] (**13**). Thus, the reaction of **1** with C₆F₅H also follows a radical reaction mechanism akin to the reaction of **1** with C₆F₆ below.

Mechanistic investigations

Experimental investigations and DFT studies reported previously 9a for the reaction of $[Ni_2(^iPr_2Im)_4(\mu-(\eta^2:\eta^2)-COD)]$ and $[Ni(^iPr_2Im)_2(\eta^2-C_2H_4)]$, used as source of $[Ni(^iPr_2Im)_2]$ ($\mathbf{1}^{ipr}$), with C_6F_6 suggested a concerted mechanism for the insertion of $\mathbf{1}^{ipr}$ into the C–F bond, and no indications for radical reactivity were obtained. As presented above, however, paramagnetic complexes clearly emerge in the reaction of $\mathbf{1}$ and C_6F_6 . To obtain further insight, we performed a quantum-chemical investigation (COSMO(THF)-PBE0-D/def2-TZVP, for details see ESI †) 17 on the reaction pathways of C_6F_6 with $[Ni(Mes_2Im)_2]$ ($\mathbf{1}^{ipr}$). and with the sterically less encumbered $[Ni(^iPr_2Im)_2]$ ($\mathbf{1}^{ipr}$).

C–F bond activation in the latter reaction commences with the formation of a rather stable 16-electron η^2 adduct between $\mathbf{1}^{ipr}$ and C_6F_6 (I1, Scheme 8; see ESI, Fig. S17†). The DFT-optimized geometry of I1 is in good agreement with the structure of the closely related complex $[Ni(^iPr_2Im)_2(\eta^2-C_{10}F_8)]^{.9a}$. Three distinct reaction pathways are then possible. First, direct oxidative addition of the C–F bond to the nickel atom proceeds through TS1 to yield the *trans* product $\mathbf{3}^{ipr}$ with an effective activation barrier of $\Delta^{\ddagger}G = 23$ kcal mol⁻¹ relative to I1 (see ESI, Fig. S18†). Alternative formation of the corresponding *cis*-

[Ni($^{\rm i}$ Pr₂Im)₂(F)(C₆F₅)] (I2) and subsequent isomerization is kinetically disfavored ($\Delta^{\dagger}G^{\rm eff}=27$ kcal mol⁻¹, see ESI Fig. S19 and S20†), as is dissociation of an NHC ligand ($\Delta G^{298}=28$ kcal mol⁻¹, see Fig. S32†).

Second, NHC ligand cooperativity (see ESI; Fig. S21 and S23†) opens a kinetically competitive pathway to the *trans*-product 3^{ipr} , that is, addition of the C–F bond across the Ni–C^{NHC} bond through **TS2** to yield intermediate **I3**, in which coordination of the fluorinated NHC–F ligand to the nickel atom involves a bridging C–N bonding interaction. In **TS2**, the C_{aryl}–F bond of 1.93 Å is strongly elongated compared to C₆F₆ (C_{aryl}–F bond: 1.32 Å) and **TS1** (C_{aryl}–F bond: 1.77 Å), while NHC–F bond formation is hardly visible (C/F distance: 2.40 Å). From **I3**, fluoride migration onto the nickel ion (**TS3**, with a low barrier of $\Delta^{\ddagger}G$ = 15 kcal mol⁻¹) leads to 3^{ipr} with an overall barrier of $\Delta^{\ddagger}G^{eff}$ = 24 kcal mol⁻¹. Third, homolytic C–F bond cleavage involves an effective barrier of $\Delta^{\ddagger}G^{eff}$ = 31 kcal mol⁻¹ (**TS4**) and, hence, radical abstraction is kinetically disfavored here (see ESI, Fig. S22†).

C–F bond activation with the sterically more congested Mes-NHC complex 1 shows marked differences. Formation of the η^2 -C₆F₆ adduct **I5** (see ESI, Fig. S24†) is now endergonic by 12 kcal mol⁻¹, and consecutive oxidative C–F bond addition *via* **TS5** ($\Delta^{\ddagger}G^{\rm eff}=21$ kcal mol⁻¹, see ESI; Fig. S25†) leads to the *cis*-product **I6**. We attribute the endergonicity of the η^2 -C₆F₆ adduct formation (**I5**, $\Delta\Delta G=28$ kcal mol⁻¹ compared to the exergonic formation of **I1**) mainly to the increased steric demand of the mesityl groups. A trajectory to the *trans*-product is precluded by the steric demand of the mesityl substituents. NHC dissociation to yield [Ni(Mes₂Im)(η^6 -C₆F₆)] and subsequent insertion into the C–F bond is associated with a large barrier ($\Delta^{\ddagger}G^{\rm eff}=34$ kcal mol⁻¹, see ESI; Fig. S32 and S33†) and is

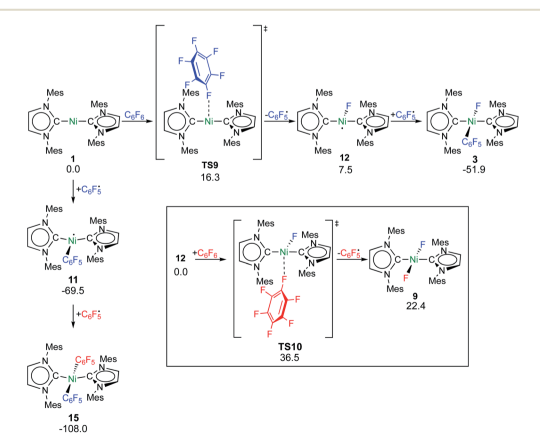
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Scheme 8 Calculated pathways for the C-F bond activation of C_6F_6 with $\mathbf{1}^{ipr}$ (ΔG^{298} in kcal mol^{-1}).

Calculated pathways for the heterolytic C-F bond cleavage of C_6F_6 by 1 and further reaction steps (ΔG^{298} in kcal mol⁻¹).

irrelevant here. Note that an alternative adduct formation stabilized by π -stacking interactions between C_6F_6 and one of the NHC mesityl substituents, 18 such as I7 (see ESI; Fig. S26†), is also endergonic and less favorable than I5. Furthermore, a "concerted" NHC-assisted process as in the ⁱPr system does not exist. We found a multi-step sequence for the mesityl system

instead (Scheme 9 and ESI; Fig. S27†), commencing with heterolytic C-F bond cleavage in I5, which exhibits a partially reduced C_6F_6 fragment $(q_{NPA}(C_6F_6) = -0.69)$. The fluoride anion expelled from the nickel coordination sphere is loosely held within the cleft formed by the mesityl substituents in I8. A similar stabilizing association of a fluoride anion by the methyl



Scheme 10 Calculated pathways for the homolytic C-F bond cleavage of C_6F_6 by 1 and further radical reaction steps (ΔG^{298} in kcal mol $^{-1}$; energies of TS10 and 9 are given relative to $12 + C_6F_6$).

groups of mesityl substituents has been reported by Macgregor *et al.* for the C–F bond activation step in hydrodefluorination reactions. Formation of the *trans*-product 3 from here involves binding to the carbene carbon atom and subsequent F-shift onto the Ni center. The overall path involves a low effective barrier of 16 kcal mol⁻¹ (TS6).

Fluorine radical abstraction to yield C₆F₅ and radical complex 12 via TS9 is slightly endergonic and exhibits a barrier of 16 kcal mol⁻¹ (Scheme 10 and ESI, Fig. S26 and S28†). Recombination of C₆F₅ and 12 to 3 then provides a large thermodynamic driving force. Alternative addition of C₆F₅ to the initial complex 1 to yield radical species 11 is also a highly exergonic process (-69.5 kcal mol⁻¹), as well as addition of a second equivalent of C_6F_5 to yield 15 (-108.0 kcal mol⁻¹). Endergonic formation of difluoride complex 9 from 12 and another equiv. of C₆F₆, can be compensated by consumption of C₆F'₅; however, a second fluorine abstraction step is prevented by the high kinetic barrier of 37 kcal mol⁻¹ via **TS10** (see ESI; Fig. S29†). The mechanism for the formation of 9 remains obscure to us thus far. We compute the ligand exchange reaction $3 + 3 \rightarrow 9 + 15$ to be exceedingly endergonic (25.7 kcal mol⁻¹), and also the disproportionation reactions of radicals 11 and 12 yielding 1 + 15 (31.0 kcal mol⁻¹) or 1 + 9, (14.9 kcal mol⁻¹), are unlikely to contribute to the formation of 9 (see ESI, Fig. S30†). A dinuclear complex [{Ni(Mes₂Im)₂}₂(μ- $(\eta^2:\eta^2)$ -C₆F₆)], which would be an intermediate for an oneelectron oxidative addition, is too high in energy to be considered (35 kcal mol⁻¹, see ESI; Fig. S31†). Hence both, the radical pathway and the NHC-assisted multistep pathway represent kinetically competitive C-F bond activation steps in the reaction with $[Ni(Mes_2Im)_2]$ (1).

Discussion

It is now well established that nickel(0) complexes with phosphine, carbene, and even some nitrogen ligands undergo C-F oxidative addition with perfluoroarenes to yield complexes trans-[Ni(L)₂(F)(C₆F₅)].^{1,9,20} Although the lack of clean kinetics for many of the C-F oxidative additions indicate complex mechanistic scenarios, there were strong indications that the conversion of C₆F₆ to the aryl fluoride complex follows the same type of mechanism as observed for typical C-H activation reactions of benzene. It has been demonstrated, for nickel NHC and phosphine complexes, that the first stage of C-F oxidative addition is the η^2 -coordination of the fluoroarene. ^{1g,9a,21,22} The introduction of fluorine substituents on the arene results in a lower lying LUMO, which renders the fluorinated arene a better electron acceptor compared to H-arenes and makes the reaction of electron-poor C₆F₆ with an electron-rich, suitable nickel precursor more exothermic. The fluoroarene of $[Ni(L)_2(\eta^2-C_6F_6)]$ is ene-diene distorted, and the arene fluoride substituents are bent out of the plane, as observed for I1 and I5. Subsequent C-F oxidative addition is strongly exothermic for $trans-[Ni(^{i}Pr_{2}Im)_{2}(F)(C_{6}F_{5})] (\Delta G^{298} = -57 \text{ kcal mol}^{-1}) \text{ and } trans [Ni(Mes_2Im)_2(F)(C_6F_5)] (\Delta G^{298} = -52 \text{ kcal mol}^{-1}).$ Computational studies reported previously94,23 of the reaction pathways have supported the idea of concerted mechanisms involving

a σ-complex as a three-center transition state between the C₆F₆ carbon and fluorine atoms and the transition metal atom. The transition state structures typically show limited elongation of the C-F bond and interaction of the electron-rich transition metal ion with the C-F σ^* orbital leads to C-F bond breaking and formation of the M-C and M-F bond. We have demonstrated now for $[Ni_2(^iPr_2Im)_4(\mu-(\eta^2:\eta^2)-COD)]$ and the related [Ni(ⁱPr₂Im)₂] (1^{ipr}) synthon complexes that C-F bond activation of C₆F₆ occurs via both a concerted and an NHC-assisted pathway, as both are associated with very similar kinetic barriers of $\Delta^{\ddagger}G^{\text{eff}} = 23 \text{ kcal mol}^{-1}$ for the concerted and of $\Delta^{\ddagger}G^{\text{eff}}$ $= 24 \text{ kcal mol}^{-1}$ for the NHC-assisted pathway. This situation will probably change if other substrates with other leaving groups, such as partially fluorinated arenes, fluoropyridines or other aryl halides, are involved in the reaction with the nickel complex; however, our calculations demonstrate that both reaction paths are feasible, at least for fluoroarenes.

The direction of the concerted oxidative addition in **TS1** to give the *trans* product is rather unusual. For the oxidative addition of A–B to d^{10} -ML $_2$ the important orbital interactions of the transition state are those between the filled $\sigma(A–B)$ orbital and the empty d_{σ} -type orbital of the metal, leading to electron donation from A–B to the metal center, and a second interaction between the filled d_{π} -orbital of the metal and the $\sigma^*(A–B)$, leading to electron transfer from the metal to the ligand. Strong back-donation will lead to fission of the A–B bond. This back-donation is strongest if A–B lies within the bent- d^{10} -ML $_2$ plane and the $\sigma^*(A–B)$ orbital can interact with the $d_{x^2-y^2}$ orbital (actually a d–p hybrid orbital), which is pointing at the two ligands L. 24

However, it was also shown previously that concerted oxidative addition reactions may take place through a nonplanar transition state structure even for non-polar substrates with dihedral angle between ML2 and M(A-B) planes larger than 70°.25 It was demonstrated that this nonplanar transition state is connected to the planar product on the singlet surface and suggested that steric rather than electronic factors are responsible for the nonplanar transition state structure. Martin et al.,25c for example, calculated at the B3LYP/LanL2DZ-level of theory a nonplanar transition state for the oxidative addition of C_6H_5 -I to [Pd(dmpe)] (dmpe = bis{dimethylphosphino}ethane), in which the P-Pd-P and C-Pd-I planes are almost perpendicular to one another. Another example was provided by Jones et al.25d for the oxidative addition of the C-CN σ-bond of organonitriles to the low-valent nickel complex [Ni(dmpe)]. The C-C-N plane of the transition state (calculated at the B3LYP/6-31G(d,p)-level of theory), which leads to C-CN bond cleavage, is rotated by 38° relative to the P-Ni-P plane.

The $\eta^2(C,C)$ -bonded complex $[Ni(^iPr_2Im)_2(\eta^2-C_6F_6)]$ (I1) is also the crucial reaction intermediate for the NHC-assisted pathway. The key step here is the addition of the C–F bond across the Ni– C_{NHC} bond and, thus, the unoccupied NHC p_{π^-} orbital plays a central role for this pathway as intramolecular fluoride acceptor. Fluoride transfer from the arene to the NHC leads to a η^2 -fluoro-imidazolyl intermediate (I3; Scheme 8) which rearranges with a second fluoride transfer step from the NHC to the nickel atom to give *trans*- $[Ni(^iPr_2Im)_2(F)(C_6F_5)]$ (3^{ipr}).

A phosphine-assisted process has been proposed before for the C-F bond activation of pentafluoropyridine with [Ni(PR₃)₂], based on the experimental observation of an unusual selectivity for the insertion into the 2-position of C5NF5 and on DFT calculations.26 However, another study performed on the reaction of pentafluoropyridine with [Ni(PEt₃)₂] suggested that pathways other than a concerted oxidative addition or a phosphine-assisted pathway account for the unusual selectivity.²⁷ The detailed experimental analysis of the reactivity of a [Ni(PEt₃)₂] precursor with perfluoropyridine demonstrated the formation of a mononuclear adduct [Ni(PEt₃)₂(η^2 -C₅F₅N)], of dinuclear adducts $[{Ni(PEt_3)_2}_2](\mu-(\eta^2:\eta^2)-C_5F_5N)]$, some of which exhibit C-F bond activation, and a nickel(1) radical species [Ni(PEt₃)₂(2-C₅F₄N)]. Other heteroatom-assisted C-F bond activation processes have also been proposed for other metals mainly including boryl or silyl moieties.28

Despite precedent in the oxidative addition of other aryl carbon-halide bonds to nickel,29,30 there is only little experimental evidence for the involvement of radicals in C-F bond activation processes. It is known that some polyfluoro pyridines react with [Ni(PR3)2] to yield EPR-active complexes as likely intermediates,16,27 and some studies on C-F bond activation have shown unusual products with highly-fluorinated arenes that may be indicative of radical pathways. 16,22,31 However, the clear identification of radical intermediates has not been possible so far and alternate mechanisms cannot be ruled out. Although DFT calculations were performed to examine the traditional concerted oxidative addition and phosphineassisted pathways for C-F bond activation, radical pathways involving Ni(I) intermediates were rarely considered computationally.

Thus, the reaction of 1 with different fluoroarenes leads to nickel insertion into the C-F bond to give the nickel fluoroaryl fluoride complexes trans-[Ni(Mes₂Im)₂(F)(Ar^F)], but EPR spectroscopy also provided evidence that at least three paramagnetic species are intermediates or products of the reaction of C₆F₆ with 1. We provide evidence that simple electron transfer from [Ni(Mes₂Im)₂] (1) to C₆F₆, often considered as the first step in radical oxidative additions at nickel,29 is unlikely to occur. The redox potentials are not in line with intermolecular electron transfer to yield [Ni(Mes₂Im)₂]⁺ and C₆F₆⁻ and the EPR resonance of [Ni(Mes₂Im)₂]⁺, which has been established for the authentic complex [Ni(Mes₂Im)₂][BF₄] (8), was not detected in the reaction mixture. Furthermore, many diamagnetic and radical products of the reaction of [Ni(Mes₂Im)₂] (1) to C₆F₆ were identified, namely the insertion product trans-[Ni(Mes₂Im)₂(- $F(C_6F_5)$ (3), the difluoride complex trans- $[Ni(Mes_2Im)_2(F)_2]$ (9), the bis(aryl) complex trans-[Ni^{II}(Mes₂Im)₂(C₆F₅)₂] (15), the nickel(1) complex trans-[Ni^I(Mes₂Im)₂(C₆F₅)] (11), and the metalcentered radical trans-[Ni^I(Mes₂Im)₂(F)] (12). DFT calculations performed on the reaction of [Ni(Mes₂Im)₂] (1) with C₆F₆ explain the occurrence of the radical species observed. Both an NHC-assisted and a radical process are kinetically equally favored routes for this reaction. Fluorine radical abstraction from C₆F₆ by **1** is associated with a barrier of only 16 kcal mol⁻¹ and subsequent radical recombination steps provide the thermodynamic driving force required.

Matsubara et al. and Louie et al. reported the clean isolation of T-shaped three-coordinate radical species $[Ni^{I}(NHC)_{2}(X)](X =$ Cl, Br, I; NHC = Mes₂Im, Dipp₂Im) from the reaction of [Ni(NHC)₂] with aryl halides. 30a,b,d We have demonstrated earlier that $[Ni_2(^1Pr_2Im)_4(\mu-(\eta^2:\eta^2)-COD)]$, a source of $[Ni(^1Pr_2Im)_2]$ (1^{ipr}), reacts cleanly with aryl chlorides to yield the nickel(11) complexes trans-[Ni(NHC)2(Cl)(Ar)].32 Our calculations show now that a trajectory to the trans-product by a concerted oxidative addition is precluded for [Ni(Mes₂Im)₂] (1) (and most probably also for [Ni(Dipp₂Im)₂]) by the steric demand of the mesityl substituents. As a consequence, other pathways such as electron transfer and radical abstraction must occur which are responsible for a limited or altered reactivity of complex [Ni(Mes₂Im)₂] (1) and analogues containing even more bulky Naryl substituents compared to complexes of sterically less demanding NHCs. However, fluoride abstraction occurs for the reaction of 1 and C_6F_6 even at -78 °C to yield trans-[Ni^I(Mes₂- $[Im]_2(C_6F_5)$ (11) and trans- $[Ni^I(Mes_2Im)_2(F)]$ (12). The latter is, in contrast to the complexes of the heavier homologues, very reactive and has defied thus far isolation. In turn, the complexes $trans-[Ni^{I}(Mes_{2}Im)_{2}(C_{6}F_{5})]$ (11), $[Ni^{I}(Mes_{2}Im)_{2}(2,3,5,6-C_{6}F_{4}H)]$ (12) and $[Ni^{I}(Mes_{2}Im)_{2}(2,3,5-C_{6}F_{3}H_{2})]$ (13) seem to be much more stable than [Ni^I(NHC)₂(C₆H₅)] and have been synthesized and characterized. The increased stability of [Ni^I(Mes₂- $Im)_2(2,3,5,6-C_6F_4H)$] (12) can be explained by the increased Ni-CAr bond strength of the fluoroaryl ligand with respect to C_6H_5 .33

Nelson and Maseras³⁴ reported computational investigations of the reaction of [Ni(NHC)2] complexes with aryl halides Ph-X (X = Cl, Br, I) and demonstrated that steric effects determine the mechanism. Small NHC ligands (NHC = Me₂Im^{Me}) favor concerted oxidative addition via a $\eta^2(C,C)$ π -coordinated intermediate leading to trans-[Ni^{II}(NHC)₂(X)(Ar)] complexes whereas larger NHC ligands (e.g. NHC = Mes₂Im) lead to halide abstraction to form [Ni^I(X)(NHC)₂] and a phenyl radical. We confirm here, by means of experiment and theory, that [Ni(NHC)₂] complexes of sterically less demanding NHCs favor the reaction with fluoroarenes via a concerted oxidative addition proceeding through an $\eta^2(C,C)$ intermediate, and that for the bulkier NHC Mes₂Im, C-F bond activation is achieved more easily by fluorine atom abstraction. However, for both mechanisms, we found an NHC-assisted pathway which is competitive, that accounts for the formation of diamagnetic products by a C-F bond activation step across the Ni-C_{NHC} bond. NHCassisted pathways play an important role for complexes of both sterically demanding and less bulky NHC ligand. We believe that this dual reaction pathway concept, including NHCassisted reaction pathways, should be of general importance and widely applicable for the reactivity of NHC transition metal complexes.

Conclusions

We present herein a detailed account of the C-F bond activation of polyfluoroaromatics, especially of C_6F_6 using the nickel(0) complex $[Ni(Mes_2Im)_2]$ (1). The reaction of 1 with different fluoroarenes leads to insertion of nickel into the C-F bond of the

fluoroarene to give the nickel(II) complexes trans-[Ni(Mes₂- $[Im]_2(F)(Ar^F)$ $[Ar^F = 4-CF_3-C_6F_4 2, C_6F_5 3, 2,3,5,6-C_6F_4N 4, 2,3,5,6-C_6F_6]$ C_6F_4H 5, 2,3,5- $C_6F_3H_2$ 6, 3,5- $C_6F_2H_3$ 7) in good to fair yields with the exception of the formation of the pentafluorophenyl complex trans-[Ni(Mes₂Im)₂(F)(C₆F₅)] (3) (less than 20%). Whereas the C-F bond activation process of C₆F₆ using [Ni(iPr₂Im)₂] (1ipr) follows a concerted or NHC-assisted mechanism to give the insertion product via η^2 -coordinated intermediates, metal radical species were detected for the reaction of 1 with C₆F₆. EPR spectroscopy provided evidence that at least three paramagnetic products are intermediates or products of this reaction. The experiments reveal that simple electron transfer from [Ni(Mes₂Im)₂] (1) to C₆F₆ is unlikely to occur as (i) the redox potentials do not match for an electron transfer between $[Ni(Mes_2Im)_2]$ (1) and C_6F_6 to give $[Ni(Mes_2Im)_2]^+$ and $C_6F_6^-$, and (ii) the EPR resonance for $[Ni(Mes_2Im)_2]^+$, as established for the stable, isolated complex [Ni(Mes₂Im)₂][BF₄] (8), was not detected in the reaction mixture. Several other byproducts were identified aside from the insertion product 3, namely the difluoride complex trans-[Ni(Mes₂Im)₂(F)₂] (9), the complex trans-[Ni^{II}(Mes₂Im)₂(C₆F₅)₂] (15), structurally-characterized nickel(1) complex trans-[Ni^I(Mes₂- $Im)_2(C_6F_5)$] (11) and the metal radical trans- $[Ni^I(Mes_2Im)_2(F)]$ (12). Complex 11 and related complexes [Ni^I(Mes₂Im)₂(2,3,5,6- C_6F_4H)] (13) and $[Ni^I(Mes_2Im)_2(2,3,5-C_6F_3H_2)]$ (14) were synthesized and characterized independently from the reaction of trans-[Ni(Mes₂Im)₂(F)(Ar^F)] with PhSiH₃.

DFT calculations were performed on the insertion of $[Ni(^{i}Pr_{2}Im)_{2}]$ (1) and $[Ni(Mes_{2}Im)_{2}]$ (1) into the C-F bond of C₆F₆, which explain the formation of radical species for the reaction with [Ni(Mes₂Im)₂] (1). For [Ni(ⁱPr₂Im)₂] (1^{ipr}), the crucial reaction intermediate is an η²(C,C)-bonded complex $[Ni(^{1}Pr_{2}Im)_{2}(\eta^{2}-C_{6}F_{6})]$, from which two favorable pathways with almost identical barriers, i.e., a concerted oxidative addition pathway and a NHC-assisted pathway, lead to the formation of trans-[Ni(1 Pr₂Im)₂(F)(C₆F₅)]. For [Ni(Mes₂Im)₂] (1), an NHCassisted and a radical pathway were identified with similar kinetic barriers. Fluorine atom abstraction from C₆F₆ at $[Ni(Mes_2Im)_2]$ (1) occurs *via* end-on attack of C₆F₆, while the key intermediate for the NHC-assisted pathway is the $\eta^2(C,C)$ intermediate $[Ni(Mes_2Im)_2(\eta^2-C_6F_6)].$ The NHC-assisted pathway can be interpreted as heterolytic C-F bond cleavage to yield ionic intermediates trans-[Ni(Mes₂Im)₂(C₆F₅)]⁺F⁻, in which the fluoride anion is stabilized within the sphere of the trans-[Ni(Mes₂Im)₂(C₆F₅)]⁺ cation. Several fluoride transfer steps, i.e., migration to the NHC, NHC rotation, and fluoride transfer to the metal cation lead to the formation of trans- $[Ni(Mes_2Im)_2(F)(C_6F_5)]$ (3).

Conflicts of interest

The authors declare no conflict of interest.

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References

- 1 (a) N. M. Doherty and N. W. Hoffmann, Chem. Rev., 1991, 91, 553-573; (b) W. R. Dolbier, J. Fluorine Chem., 2005, 126, 157-163; (c) K. Uneyama, Organofluorine Chemistry, Wiley-Blackwell, Oxford, U.K., 2006; (d) S. Purser, P. R. Moore, S. Swallow and V. Gouverneur, Chem. Soc. Rev., 2008, 37, 320-330; (e) A. D. Sun and J. A. Love, Dalton Trans., 2010, 39, 10362-10374; (f) P. Jeschke, Pest Manage. Sci., 2010, 66, 10-27; (g) E. Clot, O. Eisenstein, N. Jasim, S. A. Macgregor, J. E. McGrady and R. N. Perutz, Acc. Chem. Res., 2011, 44, 333-348; (h) S. A. Johnson, J. A. Hatnean and M. E. Doster, Prog. Inorg. Chem., 2011, 57, 255-352; (i) M. F. Kuehnel, D. Lentz and T. Braun, Angew. Chem., Int. Ed., 2013, 52, 3328-3348; (j) L. Keyes and J. A. Love, Aromatic C-F Activation: Converting Fluoroarenes to Useful Building Blocks, in C-H and C-X Bond Functionalization: Transition Metal Mediation, ed. X. Ribas, RSC, Cambridge, U.K., 2013, pp. 159-192; (k) J. Wang, M. Sanchez-Rosello, J. L. Acena, C. del Pozo, A. E. Sorochinsky, S. Fustero, V. A. Soloshonok and H. Liu, Chem. Rev., 2014, 114, 2432-2506; (l) J. Weaver and S. Senaweera, Tetrahedron, 2014, 70, 7413-7428; (m) Y. Zhou, J. Wang, Z. Gu, S. Wang, W. Zhu, J. L. Acena, V. A. Soloshonok, K. Izawa and H. Liu, Chem. Rev., 2016, 116, 422-518; (n) D. E. Yerien, S. Bonesi and A. Postigo, Org. Biomol. Chem., 2016, 14, 8398-8427; (o) O. Eisenstein, J. Milani and R. N. Perutz, Chem. Rev., 2017, 117, 8710-8753. 2 (a) K. Daasbjerg, J. Chem. Soc., Perkin Trans. 2, 1994, 1275-
- 2 (a) K. Daasbjerg, J. Chem. Soc., Perkin Trans. 2, 1994, 1275–1277; (b) Y.-R. Luo, Comprehensive Handbook of Chemical Bond Energies, CRC Press, Boca Raton, FL, 2007; (c) H. Amii and K. Uneyama, Chem. Rev., 2009, 109, 2119–2183; (d) T. Ahrens, J. Kohlmann, M. Ahrens and T. Braun, Chem. Rev., 2015, 115, 931–972.
- 3 (a) J. Zhou, M. W. Kuntze-Fechner, R. Bertermann, U. S. D. Paul, J. H. J. Berthel, A. Friedrich, Z. Du, T. B. Marder and U. Radius, *J. Am. Chem. Soc.*, 2016, **138**, 5250–5253; (b) Y.-M. Tian, X.-N. Guo, M. W. Kuntze-Fechner, I. Krummenacher, H. Braunschweig, U. Radius, A. Steffen and T. B. Marder, *J. Am. Chem. Soc.*, 2018, **140**, 17612–17623.
- 4 (a) J. W. J. Kennedy and D. G. Hall, Boronic Acids, Wiley-VCH, Weinheim, 2005; (b) Boronic acids: preparation, applications in organic synthesis and medicine, ed. D. G. Hall, Wiley-VCH, Weinheim, Germany, 2006; (c) T. Schaub, M. Backes and U. Radius, J. Am. Chem. Soc., 2006, 128, 15964–15965; (d) C. Kleeberg, L. Dang, Z. Lin and T. B. Marder, Angew. Chem., Int. Ed., 2009, 48, 5350–5354; (e) I. A. Mkhalid, J. H. Barnard, T. B. Marder, J. M. Murphy and J. F. Hartwig, Chem. Rev., 2010, 110, 890–931; (f) A. Suzuki, Angew. Chem., Int. Ed., 2011, 50, 6722–6737; (g) J. F. Hartwig, Chem. Soc. Rev., 2011, 40, 1992–2002; (h) A. Ros, R. Fernandez and J. M. Lassaletta, Chem. Soc. Rev.

- 2014, 43, 3229–3243; (*i*) S. K. Bose and T. B. Marder, *Org. Lett.*, 2014, 16, 4562–4565; (*f*) S. K. Bose, A. Deissenberger, A. Eichhorn, P. G. Steel, Z. Lin and T. B. Marder, *Angew. Chem., Int. Ed.*, 2015, 54, 11843–11847; (*k*) J. Zhou, J. H. J. Berthel, M. W. Kuntze-Fechner, A. Friedrich, T. B. Marder and U. Radius, *J. Org. Chem.*, 2016, 81, 5789–5794; (*l*) Y. P. Budiman, A. Friedrich, U. Radius and T. B. Marder, *ChemCatChem*, 2019, 11, 5387–5396; (*m*) Z. Liu, Y. P. Budiman, Y. Tian, A. Friedrich, M. Huang, S. A. Westcott, U. Radius and T. B. Marder, *Chem.-Eur. J.*, 2020, DOI: 10.1002/chem.202002888.
- (a) M. S. Cheung, F. K. Sheong, T. B. Marder and Z. Lin, *Chem.-Eur. J.*, 2015, 21, 7480–7488; (b) L. Kuehn, M. Stang,
 S. Würtemberger-Pietsch, A. Friedrich, H. Schneider,
 U. Radius and T. B. Marder, *Faraday Discuss.*, 2019, 220, 350–363.
- 6 (a) J. R. Chen, X. Q. Hu, L. Q. Lu and W. J. Xiao, Acc. Chem. Res., 2016, 49, 1911–1923; (b) A. M. Mfuh, J. D. Doyle, B. Chhetri, H. D. Arman and O. V. Larionov, J. Am. Chem. Soc., 2016, 138, 2985–2988; (c) A. M. Mfuh, V. T. Nguyen, B. Chhetri, J. E. Burch, J. D. Doyle, V. N. Nesterov, H. D. Arman and O. V. Larionov, J. Am. Chem. Soc., 2016, 138, 8408–8411; (d) K. Chen, S. Zhang, P. He and P. Li, Chem. Sci., 2016, 7, 3676–3680; (e) K. Chen, M. S. Cheung, Z. Lin and P. Li, Org. Chem. Front., 2016, 3, 875–879; (f) A. M. Mfuh, B. D. Schneider, W. Cruces and O. V. Larionov, Nat. Protoc., 2017, 12, 604–610; (g) H. Yi, G. Zhang, H. Wang, Z. Huang, J. Wang, A. K. Singh and A. Lei, Chem. Rev., 2017, 117, 9016–9085.
- 7 C. Sieck, M. G. Tay, M. H. Thibault, R. M. Edkins, K. Costuas, J. F. Halet, A. S. Batsanov, M. Haehnel, K. Edkins, A. Lorbach, A. Steffen and T. B. Marder, *Chem.–Eur. J.*, 2016, **22**, 10523–10532.
- 8 (a) T. Schaub and U. Radius, Chem.-Eur. J., 2005, 11, 5024–5030; (b) T. Schaub, M. Backes and U. Radius, Organometallics, 2006, 25, 4196–4206; (c) P. Fischer, T. Linder and U. Radius, Z. Anorg. Allg. Chem., 2012, 638, 1491–1496; (d) A. P. Prakasham and P. Ghosh, Inorg. Chim. Acta, 2015, 431, 61–100; (e) A. A. Danopoulos, T. Simler and P. Braunstein, Chem. Rev., 2019, 119, 3730–3961.
- 9 (a) T. Schaub, P. Fischer, A. Steffen, T. Braun, U. Radius and A. Mix, J. Am. Chem. Soc., 2008, 130, 9304–9317; (b) T. Schaub, M. Backes and U. Radius, Eur. J. Inorg. Chem., 2008, 2680–2690; (c) T. Schaub, P. Fischer, T. Meins and U. Radius, Eur. J. Inorg. Chem., 2011, 3122–3126; (d) P. Fischer, K. Götz, A. Eichhorn and U. Radius, Organometallics, 2012, 31, 1374–1383; (e) U. S. D. Paul and U. Radius, Chem.-Eur. J., 2017, 23, 3993–4009; (f) M. W. Kuntze-Fechner, C. Kerpen, D. Schmidt, M. Häring and U. Radius, Eur. J. Inorg. Chem., 2019, 1767–1775.
- 10 W. R. Hagen, Dalton Trans., 2006, 4415-4434.
- 11 (a) R. C. Poulten, M. J. Page, A. G. Algarra, J. J. Le Roy,
 I. Lopez, E. Carter, A. Llobet, S. A. Macgregor,
 M. F. Mahon, D. M. Murphy, M. Murugesu and
 M. K. Whittlesey, J. Am. Chem. Soc., 2013, 135, 13640–13643; (b) M. M. Schwab, D. Himmel, S. Kacprzak,
 V. Radtke, D. Kratzert, P. Weis, M. Wernet, A. Peter,

- Z. Yassine, D. Schmitz, E. W. Scheidt, W. Scherer, S. Weber, W. Feuerstein, F. Breher, A. Higelin and I. Krossing, *Chem.-Eur. J.*, 2018, 24, 918–927.
- 12 Molecular structures of the radical species discussed in this section were obtained at the RI-PBE-D/def2-SVP/W06 level of DFT. The PBE0-D functional and the pcSseg-2 basis set, which is specifically optimized for the calculation of magnetic properties, were employed to obtain EPR parameters (PBE0-D/pcSseg-2).
- 13 (a) P. Nilsson, F. Plamper and O. F. Wendt, *Organometallics*, 2003, 22, 5235–5242; (b) A. Yahav, I. Goldberg and A. Vigalok, *Inorg. Chem.*, 2005, 44, 1547–1553.
- 14 (a) R. K. Marat and A. F. Janzen, Can. J. Chem., 1977, 55, 3845–3849; (b) R. Damrauer, R. A. Simon and B. Kanner, Organometallics, 1988, 7, 1161–1164; (c) A. Kunai, T. Sakurai, E. Toyoda and M. Ishikawa, Organometallics, 1996, 15, 2478–2482.
- 15 J. F. Hartwig, Organotransition Metal Chemistry: From Bonding to Catalysis, University Science Books, Mill Valley, CA, 2010, ch. 3.8.
- 16 J. A. Hatnean, M. Shoshani and S. A. Johnson, *Inorg. Chim. Acta*, 2014, 422, 86–94.
- 17 Optimized molecular structures and thermal/vibrational contributions to Gibbs free energies were obtained at the PBE0-D/def2-SVP level of DFT. The def2-TZVP basis set and the COSMO solvation model were employed to obtain improved relative energies (COSMO(THF)-PBE0-D/def2-TZVP).
- 18 (a) A. Hori, Arene-Perfluoroarene Interactions in Coordination Architectures, in *The Importance of Pi-Interactions in Crystal Engineering: Frontiers in Crystal Engineering*, ed. E. R. T. Tiekink and J. Zuckerman-Schpector, John Wiley & Sons, Chichester, UK, 2012, pp. 163–185; (b) T. Dahl, *Acta Chem. Scand.*, 1988, 42, 1–7; (c) J. C. Collings, K. P. Roscoe, R. L. Thomas, A. S. Batsanov, L. M. Stimson, J. A. K. Howard and T. B. Marder, *New J. Chem.*, 2001, 25, 1410–1417; (d) J. C. Collings, K. P. Roscoe, E. G. Robins, A. S. Batsanov, L.-M. Stimson, J. A. K. Howard, S. J. Clark and T. B. Marder, *New J. Chem.*, 2002, 26, 1740–1746; (e) Y. P. Budiman, A. Friedrich, U. Radius and T. B. Marder, *ChemCatChem*, 2019, 11, 5387–5396.
- S. A. Macgregor, D. McKay, J. A. Panetier and M. K. Whittlesey, *Dalton Trans.*, 2013, 42, 7386–7395.
- 20 M. E. Doster and S. A. Johnson, Angew. Chem., Int. Ed., 2009, 48, 2185–2187.
- 21 T. Braun, L. Cronin, C. L. Higgitt, J. E. McGrady, R. N. Perutz and M. Reinhold, *New J. Chem.*, 2001, 25, 19–21.
- 22 S. A. Johnson, E. T. Taylor and S. J. Cruise, *Organometallics*, 2009, **28**, 3842–3855.
- 23 M. Reinhold, J. E. McGrady and R. N. Perutz, *J. Am. Chem. Soc.*, 2004, **126**, 5268–5276.
- 24 (a) R. H. Crabtree, *The Organometallic Chemistry of the Transition Metals*, Wiley-VCH, New York, 6th edn, 2014, ch.
 6; (b) J. F. Hartwig, *Organotransition Metal Chemistry: From Bonding to Catalysis*, University Science Books, Mill Valley, CA, 2010, ch. 6–7; (c) T. A. Albright, J. K. Burdett and

Edge Article Chemical Science

- M.-H. Whangbo, *Orbital Interactions in Chemistry*, Wiley & Sons, Hoboken, N. J., 2nd edn, 2013, ch. 19.6; (*d*) S. Niu and M. B. Hall, *Chem. Rev.*, 2000, **100**, 353–405; (*e*) K. C. Lam, T. B. Marder and Z. Lin, *Organometallics*, 2007, **26**, 758–760; (*f*) L. P. Wolters and F. M. Bickelhaupt, *Struct. Bonding*, 2016, **167**, 139–162.
- 25 (a) S. Sakaki, N. Mizoe, Y. Musashi, B. Biswas and M. Sugimoto, J. Phys. Chem. A, 1998, 102, 8027–8036; (b) W. Guan, F. B. Sayyed, G. Zeng and S. Sakaki, Inorg. Chem., 2014, 53, 6444–6457; (c) A. Sundermann, O. Uzan and J. M. L. Martin, Chem.-Eur. J., 2001, 7, 1703–1711; (d) T. A. Atesin, T. Li, S. Lachaize, W. W. Brennessel, J. J. Garcia and W. D. Jones, J. Am. Chem. Soc., 2007, 129, 7562–7569.
- 26 (a) L. Cronin, C. L. Higgitt, R. Karch and R. N. Perutz, Organometallics, 1997, 16, 4920-4928; (b) A. Nova, S. Erhardt, N. A. Jasim, R. N. Perutz, S. A. Macgregor, J. E. McGrady and A. C. Whitwood, J. Am. Chem. Soc., 2008, 130, 15499-15511; (c) A. Nova, M. Reinhold, R. N. Perutz, S. A. Macgregor and J. E. McGrady, Organometallics, 2010, 29, 1824-1831; (d) E. Jiao, F. Xia and H. Zhu, Comput. Theor. Chem., 2011, 965, 92-100.
- 27 J. A. Hatnean and S. A. Johnson, *Organometallics*, 2012, 31, 1361–1373.
- 28 See for example: (a) M. Aizenberger and D. Milstein, *Science*,
 1994, 265, 359–361; (b) R. J. Lindup, T. B. Marder,
 R. N. Perutz and A. C. Whitwood, *Chem. Commun.*, 2007,
 3664–3666; (c) T. Braun, F. Wehmeier and K. Altenhöhner,
 Angew. Chem., Int. Ed., 2007, 46, 5321–5324; (d)

- M. Teltewskoi, J. A. Panetier, S. A. MacGregor and T. Braun, *Angew. Chem., Int. Ed.*, 2010, 49, 3947–3951.
- 29 T. T. Tsou and J. K. Kochi, *J. Am. Chem. Soc.*, 1979, **101**, 6319–6332.
- 30 (a) S. Miyazaki, Y. Koga, T. Matsumoto and K. Matsubara, Chem. Commun., 2010, 46, 1932–1934; (b) K. Zhang, M. Conda-Sheridan, S. R. Cooke and J. Louie, Organometallics, 2011, 30, 2546–2552; (c) A. Manzoor, P. Wienefeld, M. C. Baird and P. H. M. Budzelaar, Organometallics, 2017, 36, 3508–3519; (d) C.-Y. Lin and P. P. Power, Chem. Soc. Rev., 2017, 46, 5347–5399.
- 31 S. A. Johnson, N. M. Mroz, R. Valdizon and S. Murray, *Organometallics*, 2011, **30**, 441–457.
- 32 (a) T. Zell, M. Feierabend, B. Halfter and U. Radius, J. Organomet. Chem., 2011, 696, 1380–1387; (b) T. Zell, P. Fischer, D. Schmidt and U. Radius, Organometallics, 2012, 31, 5065–5073; (c) L. Kuehn, D. G. Jammal, K. Lubitz, T. B. Marder and U. Radius, Chem.–Eur. J., 2019, 25, 9514–9521.
- 33 (a) M. E. Evans, C. L. Burke, S. Yaibuathes, E. Clot, O. Eisenstein and W. D. Jones, J. Am. Chem. Soc., 2009, 131, 13464–13473; (b) E. Clot, M. Besora, F. Maseras, C. Mégret, O. Eisenstein, B. Oelckers and R. N. Perutz, Chem. Commun., 2003, 490–491; (c) E. Clot, C. Mégret, O. Eisenstein and R. N. Perutz, J. Am. Chem. Soc., 2009, 131, 7817–7827.
- 34 D. J. Nelson and F. Maseras, *Chem. Commun.*, 2018, 54, 10646–10649.