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## Highly tunable fluorinated trispyrazolylborates $[\text{HB}(3\text{-CF}_3\text{-5-}\{4\text{-RPh}\}\text{pz})_3]^-$ ( $\text{R} = \text{NO}_2, \text{CF}_3, \text{Cl}, \text{F}, \text{H}, \text{OMe}$ and $\text{NMe}_2$ ) and their copper(I) complexes†

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The ethene and carbon monoxide adducts of copper(I) with seven trispyrazolylborate ligands ( $[\text{HB}(3\text{-CF}_3\text{-5-}\{4\text{-RPh}\}\text{pz})_3]^-$ ;  $\text{R} = \text{NO}_2$  (**4a**),  $\text{CF}_3$  (**4b**),  $\text{Cl}$  (**4c**),  $\text{F}$  (**4d**),  $\text{H}$  (**4e**),  $\text{OMe}$  (**4f**) and  $\text{NMe}_2$  (**4g**)) were synthesized and characterized. The ligands were synthesized from their corresponding pyrazoles and sodium tetrahydridoborate and were obtained as solvent adducts of their sodium salts after workup. When the pyrazole with the most electron-withdrawing substituent ( $\text{R} = \text{NO}_2$ ) is used the asymmetric ligand  $[\text{HB}(3\text{-CF}_3\text{-5-}\{4\text{-NO}_2\text{Ph}\}\text{pz})_2(3\text{-}\{4\text{-NO}_2\text{Ph}\}\text{-5-}\text{CF}_3\text{pz})]^-$  (**4a'**) is formed as the major product. Copper(I) complexes with ethene or CO as a co-ligand were prepared in good yields and were structurally characterized using  $^1\text{H}$  NMR,  $^{13}\text{C}$  NMR and infrared spectroscopy. Single crystal X-ray crystallography analyses revealed the structures of  $\text{Na4a}'$ ,  $\text{Na4b}$ , four copper ethene complexes and four copper carbonyl complexes. The structures of the copper(I) complexes show  $\text{Cu}^1$  ions in pseudo-tetrahedral coordination environments consisting of three nitrogen atoms of the trispyrazolylborate ligand and the carbonyl or  $\eta^2$ -coordinated ethene ligands, with nearly identical coordination environments around the  $\text{Cu}^1$  ion. The compound  $[\text{Na}(\text{4a}')(\text{H}_2\text{O})]_n$  crystallizes as one-dimensional chains with intermolecular  $\text{Na}\cdots\text{O}_2\text{N}$  interactions. The sodium ions were found in severely distorted octahedral geometries with three nitrogen atoms from the trispyrazolylborate ligand, one aqua ligand and two oxygen atoms from the nitro group of an adjacent molecule. The compound  $[\text{Na}_2(\text{4b})_2(\mu\text{-H}_2\text{O})_2]$  crystallizes as a centrosymmetric water-bridged dimer: two five-coordinate square-pyramidal sodium ions each are coordinated facially by three nitrogen atoms from a trispyrazolylborate ligand and two bridging water ligands. Below the base of the pyramidal structure one intermolecular and two intramolecular  $\text{Na}\cdots\text{F}$  short contacts are present. The  $^1\text{H}$  and  $^{13}\text{C}$  NMR spectra of the copper–ethene complexes show signals for the ethene ligands in the range of 4.84–4.96 ppm and 84.9–86.8 ppm respectively. The infrared spectra of the carbonyl complexes show CO stretching frequencies in the range of 2096–2120  $\text{cm}^{-1}$ . Both the NMR signals for the ethene ligands and infrared signals for the carbonyl ligands were found to show good correlations with the Hammett  $\sigma_p$  parameters of the substituents on the phenyl rings of the ligands.

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## Introduction

Trispyrazolylborates (commonly referred to as scorpionate ligands<sup>1</sup>) form a highly versatile class of ligands that were pioneered by Trofimenko in the 1960s. They have since been developed from the simple unsubstituted  $\text{Tp}^-$  into ligands of increasing complexity and scope.<sup>2–7</sup> The great diversity in steric and electronic properties available in trispyrazolylborate

ligands allows for the optimization of complexes for specific purposes such as catalysis and biomimetic structural and functional models. The properties of trispyrazolylborate ligands can be changed by systematic modification of the substituents on the pyrazolyl groups or, though much less common, on the central boron atom. At present a wide variety of the facially coordinating tridentate ligands is known with substituents ranging from alkyl groups to aryl groups (both aromatic like phenyl groups and heteroaromatic such as the thiienyl group) as well as phosphines, esters and amides.<sup>8–13</sup> Due to the structurally and electronically similar properties of the pyrazole and imidazole rings trispyrazolylborate ligands offer an interesting avenue towards the modelling of biological systems that incorporate multiple histidine residues coordinating facially to the metal center in the active site. A metal that is encountered

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often in such sites is copper, in enzymes and proteins such as hemocyanin, cytochrome-*c* oxidase, ceruloplasmin, superoxide dismutase, laccase and many others.<sup>14</sup> Copper offers advantages besides being biologically relevant; the ethene and carbonyl complexes of copper(i) offer relatively simple systems to characterize the steric and electronic properties of novel trispyrazolylborate ligands. Such complexes have a (pseudo) tetrahedral coordination environment in which a single copper(i) ion is facially coordinated by a single trispyrazolylborate ligand and one ethene or carbonyl ligand which allows for unambiguous characterization of the steric and electronic properties of the trispyrazolylborate ligand. Unfortunately comparison of the properties of trispyrazolylborate ligands typically results in their differences being ascribed to a combination of steric and electronic factors without detailed knowledge of the contributions of each factor. To facilitate the comparison of the steric and electronic factors a series of isosteric, purely electronically varied trispyrazolylborates would be useful. By keeping the steric properties of the binding pocket around the metal center constant and varying only the electronic properties of the trispyrazolylborate ligands a spectrochemical series of isosteric trispyrazolylborate ligands could be created. Such a spectrochemical series of trispyrazolylborate ligands could be used to separately estimate the optimal electronic properties of a trispyrazolylborate ligand in a catalytic or biomimetic system prior to optimization of the steric factors. To create a spectrochemical trispyrazolylborate ligand series strict regiochemical control during ligand synthesis is required. The coordination environment around the metal must vary as little as possible while the electronic properties of the ligand are varied. In this regard trispyrazolylborate ligand synthesis offers a convenient handle in that the 3-positions of the pyrazolyl rings are typically occupied by the bulkiest substituent. Exceptions to this situation occur if the pyrazole also carries a particularly electron-withdrawing substituent like a trifluoridomethyl group, in which case the electron-withdrawing group assumes the 3-position instead. The resulting electron-withdrawing trispyrazolylborate ligands have a special place in copper(i) chemistry because they allow for the synthesis and isolation of otherwise thermally and oxidatively unstable copper(i) complexes with ligands like ethene and carbon monoxide. Examples of such ligands include  $[Tp^{(CF_3)_2}]^-$ ,  $[Tp^{(CF_3)_3}]^-$ ,  $[Tp^{CF_3, Ph}]^-$ ,  $[Ttz^{(CF_3)_2}]^-$  and  $[Tp^{CF_3}]^-$ .<sup>15-18</sup> The  $[Tp^{CF_3, R}Cu(C_2H_4)]$  complexes of these ligands are air stable as are the corresponding carbonyl complexes which have infrared CO stretching frequencies close to that of free CO indicating particularly electron-poor copper(i) centers. Both effects are unusual for copper(i) trispyrazolylborate complexes and can be ascribed to the electron-withdrawing effects of the fluorinated substituents on the pyrazole moieties. The trifluoridomethyl groups thus tune the electronic properties of the ligand to a great extent while adding only a limited amount of steric bulk and a chemically inert binding pocket around the copper(i) center. With the aim of synthesizing isosteric, electronically different trispyrazolylborate ligands the ligand  $[Tp^{CF_3, Ph}]^-$  in particular is interesting because of its potential

**Table 1** Schematic representation of the ligands and complexes described in this paper, including the Hammett  $\sigma_p$  values of the substituents on the *para*-positions of the phenyl groups<sup>a</sup>

R	$\sigma_p$ <sup>b</sup>	Ligand	$[Tp^{CF_3, 4-RPh}CuL]$	
			L = C <sub>2</sub> H <sub>4</sub>	L = CO
NO <sub>2</sub> <sup>c</sup>	0.78	4a/4a'	5a'	6a'
CF <sub>3</sub>	0.54	4b	5b	6b
Cl	0.23	4c	5c	6c
F	0.06	4d	5d	6d
H	0	4e	5e	6e
OMe	-0.27	4f	5f	6f
NMe <sub>2</sub>	-0.83	4g	5g	6g
<b>5a-g and 6a-g</b>				

<sup>a</sup> The third pyrazole ring is schematically shown as [N=N]. <sup>b</sup> Taken from ref. 18. <sup>c</sup> The major product is an asymmetric isomer, indicated with a', see text.

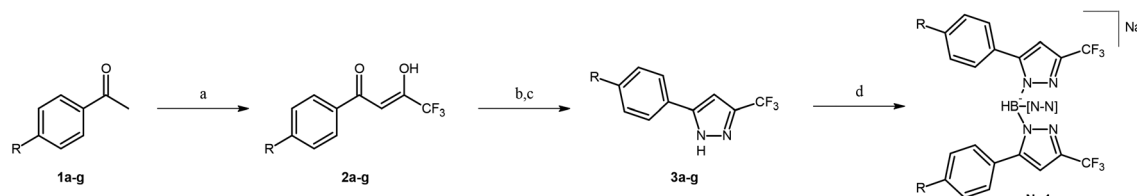
for chemical modification of the phenyl rings while retaining the structure of the binding pocket surrounding the copper(i) center. Delocalization in the aromatic rings allows for efficient charge transfer from substituents on the distal phenyl rings to the copper(i) ion while conformational differences between the resulting complexes are limited by the steric bulk of the crowded phenyl groups. Particularly substitution on the *para*-position of the phenyl rings would result in charge transfer without significant structural modification to the ligands. Even though the substituents would be placed relatively far away from the copper(i) ions the charges of the substituents could be transferred quite effectively by resonance effects in the conjoined aromatic rings. In this work the effect of *para*-substitution on the phenyl rings of  $[Tp^{CF_3, Ph}]^-$  was examined; by judicious selection of electron donating and withdrawing substituents a spectroscopic series was created (Table 1). Substituents were chosen that offer a broad range of electronic properties (based on their Hammett  $\sigma_p$  parameters) and are tolerant to the strongly reducing conditions required for the synthesis of these trispyrazolylborate ligands.<sup>19</sup> To study the upper limit of electron withdrawing effects the particularly electron withdrawing nitro substituent was also included despite its sensitivity to reducing conditions.

## Results

### Synthesis

The ligands used in this work were synthesized from 4'-substituted acetophenones (**1a-g**) in a multistep procedure (see Scheme 1). The 4-substituted acetophenones were converted into 4'-substituted benzoyl trifluoroacetates (**2a-g**) in a Claisen condensation with ethyl trifluoroacetate in THF or diethyl ether. Typically the use of diethyl ether gave higher yields and less side products but the more polar THF was sometimes required to keep reaction mixtures from becoming too viscous.



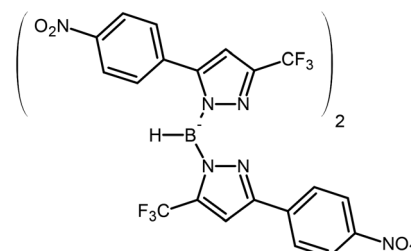


**Scheme 1** (a)  $\text{CF}_3\text{COOEt}$ ,  $\text{KO}^\text{t}\text{Bu}/\text{NaOEt}$ ,  $\text{Et}_2\text{O}$ , rt; (b)  $\text{N}_2\text{H}_4 \cdot \text{H}_2\text{O}$ ,  $\text{EtOH}$ , reflux; (c) 3 N  $\text{HCl}$ , reflux; (d)  $\text{NaBH}_4$ , Ar either at 180 °C (melt reaction) or in refluxing 4-methylanisole.

The 4'-substituted benzoyltrifluoridoacetates were then converted into the corresponding pyrazoles (**3a–g**) by a two-step condensation reaction with hydrazine hydrate which resulted in product mixtures comprising the desired pyrazole and (partially) hydrated intermediates which were dehydrated by refluxing in dilute hydrochloric acid; without the additional dehydration step the reaction is typically incomplete. Alternative methods were less successful; reflux in toluene under Dean–Stark conditions required much longer reaction times and vacuum thermolysis frequently led to co-sublimation of incompletely dehydrated products. The pyrazoles can be purified readily by means of vacuum sublimation to yield pure and dry products that can be used immediately for ligand synthesis.

The pyrazoles were converted into the trispyrazolylborate ligands by heating in the presence of  $\text{NaBH}_4$  (Scheme 1). A reaction temperature of 180 °C was used in a solvent-free reaction as reported for the synthesis of **Na4e** except in the cases of **Na4a** and **Na4f** for which 4-methylanisole was used as a solvent.<sup>16</sup> In the case of **Na4f** the solvent was used because the melting point of **3g** is too high to let the reaction proceed smoothly at 180 °C and higher temperatures led to excessive scorching. For **Na4a** the solvent-free synthesis resulted in the formation of large amounts of byproducts, whereas stepwise heating in 4-methylanisole facilitated the formation of the monopyrazolylborate and bispyrazolylborate intermediates at lower reaction temperatures before raising the reaction temperature to 180 °C. This was not possible without the use of a solvent as the melting point of **3a** is too high (171 °C). As evident from the  $^1\text{H}$  and  $^{13}\text{C}$  NMR spectra of the sodium salt and the complexes of  $[\mathbf{4a}]^-$  the nitro-substituted ligand formed almost exclusively as the asymmetric species in which two of the pyrazole rings have the trifluoridomethyl group in the 3-position and one pyrazole ring is attached “in reverse” with the trifluoridomethyl group in the 5-position (see Scheme 2). This asymmetric isomer will be referred to as **4a'**.

All trispyrazolylborate ligands were isolated as solvent adducts of their sodium salts, typically incorporating solvents such as acetone, THF or diethyl ether in stoichiometric ratios. Instances of mixed solvent adducts were also observed such as in  $\text{Na4d} \cdot (\text{Et}_2\text{O})_{0.5}(\text{acetone})_{0.5}$ . Examples of similar solvent adducts in other fluorinated trispyrazolylborate ligands have been described.<sup>17,20,21</sup> **Na4b** formed as the water adduct even when recrystallized in the presence of suitable solvents or



**Scheme 2** The asymmetric structure of the major fraction  $[\mathbf{4a}']^-$ .

mixtures thereof. All ligand salts exchanged some solvent with adventitious water upon standing in air.

The ethene and carbonyl copper(i) complexes (respectively **5a'**, **5b–g** and **6a'**, **6b–g**, see Table 1) were synthesized by stirring the sodium salts of the trispyrazolylborate ligands **Na4a'**, **Na4b–g** and  $\text{CuI}$  in DCM in an atmosphere of ethene or carbon monoxide. It was found that if  $\text{CuI}$  was added to an ethene or carbon monoxide saturated solution of the ligand in DCM the synthesis could be performed with minimal further use of Schlenk techniques. Workup consisted of filtration to remove  $\text{NaI}$  and evaporation of the solvent *in vacuo* to yield the complexes as white powders which were recrystallized from DCM/pentane to yield the pure products. Most of the isolated ethene or carbonyl copper(i) complexes are air stable for at least six months in the solid state. In solution in dichloromethane **5a'**, **6a'**, **5g** and **6g** slowly turn green over the course of hours with associated loss of the ethene or carbonyl ligands (as evident from their NMR spectra) if exposed to air. Complexes **5b–g** and **6b–g** show good stability in the presence of air, light and moisture and can be manipulated without special precautions. The complexes **5a'** and **6a'** were found to be less stable over time and had to be kept under argon to avoid decomposition. Aside from DCM the complexes were found to be slightly soluble in *n*-pentane, *n*-hexane and cyclohexane and to have good solubility in benzene, toluene, THF, 1,4-dioxane, chloroform and acetonitrile. A general trend appears to be that the more polar complexes (those incorporating methoxy and dimethylamino substituents) are less soluble in the more apolar solvents although even **5g** and **6g** are somewhat soluble in *n*-hexane. Coordinating solvents (acetonitrile, acetone and THF) cause decomposition of the complexes over

**Table 2** Selected bond distances (Å) and angles (°) for **Na4a'** and **Na4b**

Bond distances (Å)	Na4a'	Na4b	Bond angles (°)	Na4a'	Na4b
Na1–N12	2.4731(15)	2.4595(12)	O1W–Na1–O1W'		82.03(4)
Na1–N22	2.4351(15)	2.4440(12)	O1W–Na1–N12	99.18(6)	104.30(4)
Na1–N32	2.5737(15)	2.4355(12)	O1W–Na1–O231'	86.28(6)	
Na1–O1W	2.2581(16)	2.4512(12)	O1W–Na1–O232'	67.35(5)	
Na1–O1W'		2.4274(12)	O1W–Na1–N12		100.89(4)
N11–N12	1.3566(19)	1.3605(14)	O1W'–Na1–N32		99.69(4)
N21–N22	1.3603(18)	1.3589(15)	O1W–Na1–N22	103.78(6)	104.57(4)
N31–N32	1.3551(19)	1.3590(15)	N22–Na–N32	81.30(5)	73.79(4)
Na1…F12	3.625(2)		N12–Na–F13'		162.1(3)
Na1…F13'		2.930(12)	O231'–Na1–O231'	46.64(4)	
Na1…F21'	4.144(1)				
Na1…F23		3.197(4)			
Na1…F33		3.116(2)			

the course of hours to days; decomposition is slowest for the most Lewis acidic complexes.

### Single crystal X-ray crystallography

Colorless single crystals of **Na4a'**, **Na4b**, **5b**, **5c**, **5d**, **5f**, **6b**, **6c**, **6d** and **6f** were obtained by slow evaporation of DCM solutions at –20 °C and were characterized by single crystal X-ray crystallography. Crystallographic data are given in Tables S1–S3.† Selected distances and bond angles are given in Table 2 for the sodium salts of **4a'** and **4b**, in Table 3 for the ethene complexes, and in Table 4 for the carbonyl complexes. Projections of the structures of **Na4a'** and **Na4b** are shown in Fig. 1 and 2. As representative examples of the copper(I) complexes projections of the structures of **5b** and **6b** are shown in Fig. 3; projections of the other structures are provided in the ESI.†

**Na4a'** crystallized as a coordination polymer with bridging  $\text{NO}_2\cdots\text{Na}^+$  interactions (with bidentate  $\kappa^2\text{-O},\text{O}'$  coordination). The sodium ions are in a severely distorted octahedral coordination sphere comprising three nitrogen atoms from the ligand **4a'**, two oxygen atoms from a nitro group of an adjacent complex and a water molecule, which is stabilized by hydrogen bridges to the nearby nitro group oxygen atoms. The Na–N distances range from 2.4351(15) to 2.5737(15) Å, the Na–O<sub>aqua</sub> distance is 2.25581(16) Å and the Na–O<sub>2N</sub> distances are 2.5291 (14) and 2.8633(14) Å. The severely distorted octahedral coordination sphere on the sodium ion in **Na4a'** appears to be complemented by an intramolecular Na…FC contact between the sodium ion and a nearby trifluoromethyl group. The upper limit to what may be considered as short contacts (and thus potentially dative bonds) between a hard donor/acceptor pair such as sodium ions and fluorine atoms in fluoridocarbons is somewhat vague as it depends on the ionic and covalent radii that are used. Using the ionic radius of six-coordinate  $\text{Na}^+$  ions reported by Shannon *et al.* (1.02 Å) Plenio *et al.* suggested an upper limit to the interatomic distance between  $\text{Na}^+$  ions and fluorine atoms in fluoridocarbons at 3.07 Å.<sup>22,23</sup> It was further pointed out that Na…FC contacts typically have shallow potential wells, which means small outside forces such as those present in crystals because of packing can cause significant

**Table 3** Selected bond distances (Å) and angles (°) for complexes **5b**, **5c**, **5d** and **5f**<sup>a</sup>

Bond distances (Å)				
	<b>5b</b>	<b>5c</b>	<b>5d</b>	<b>5f</b>
C=C <sub>ethene</sub>	1.357(4)	1.311(14)	1.350(3)	1.351(8)
Cu–C <sub>ethene</sub>	2.034(2)	2.019(9)	2.0428(16)	2.054(5)
	2.051(5)	2.035(9)	2.0455(16)	2.046(5)
Cu–N12	2.0450(15)	2.094(3)	2.0255(14)	2.026(4)
Cu–N22	2.0369(15)	2.070(3)	2.0164(13)	2.046(4)
Cu–N32	2.2465(16)	2.137(4)	2.2964(13)	2.224(4)
N11–N12	1.3626(19)	1.356(5)	1.3685(17)	1.360(5)
N21–N22	1.3619(19)	1.355(5)	1.3648(17)	1.363(6)
N31–N32	1.3631(19)	1.363(5)	1.3682(17)	1.358(5)
N11–B1	1.553(2)	1.558(5)	1.553(2)	1.564(5)
N21–B1	1.554(2)	1.561(5)	1.552(2)	1.539(6)
N31–B1	1.545(2)	1.537(6)	1.545(2)	1.551(6)

### Bond angles (°)

	<b>5b</b>	<b>5c</b>	<b>5d</b>	<b>5f</b>
C–Cu–C	38.79(11)	37.7(4)	38.55(7)	38.5(2)
Cu–C=C	71.3(2)	70.5(5)	70.83(10)	70.5(3)
	69.9(2)	71.8(5)	70.62(10)	71.1(3)
N12–Cu1–N22	92.55(6)	89.58(14)	90.77(5)	94.64(15)
N22–Cu1–N32	87.03(6)	90.60(13)	90.77(5)	88.25(14)
N12–Cu1–N32	89.81(5)	90.17(13)	85.68(5)	87.01(14)
C1–Cu1–N12	149.26(9)	126.2(4)	115.04(7)	148.49(18)
C1–Cu1–N22	112.92(9)	111.5(3)	147.96(7)	110.75(18)
C1–Cu1–N32	107.70(9)	135.8(4)	108.95(7)	111.30(18)
C2–Cu1–N12	111.18(9)	111.8(3)	151.13(6)	110.68(18)
C2–Cu1–N22	147.41(11)	148.9(3)	111.15(6)	144.25(19)
C2–Cu1–N32	114.02(12)	110.8(3)	111.57(7)	116.93(18)

<sup>a</sup> Bond distances and bond angles are only provided for one of the two crystallographically independent molecules.

changes to the interatomic distance between the sodium ion and the fluorine atom. The distances observed for the Na…FC contacts in **Na4a'** are 3.625(2) Å and 4.144(1) Å, considerably longer than the upper limit of 3.07 Å even when crystal packing effects are taken into account. The proximity of the trifluoromethyl groups is therefore not considered to constitute



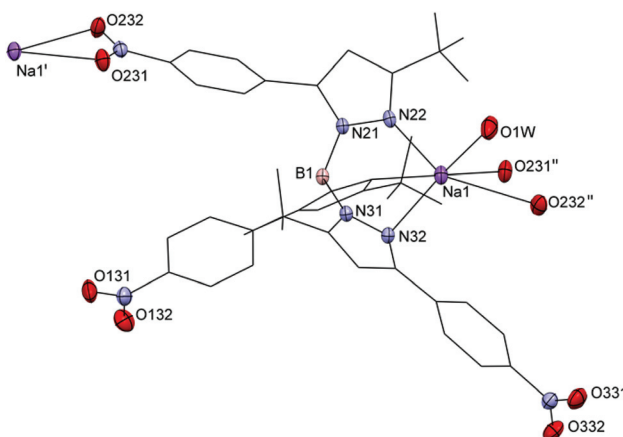
**Table 4** Selected bond distances (Å) and angles (°) for complexes **6b**, **6c**, **6d** and **6f**<sup>a</sup>

Bond distances (Å)			
	<b>6b</b>	<b>6c</b>	<b>6d</b>
C1–O11	1.124(2)	1.126(4)	1.123(8)
Cu1–C1	1.7998(19)	1.800(3)	1.791(6)
Cu–N12	2.0642(15)	2.069(2)	2.0552(19)
Cu–N22	2.0496(15)	2.056(2)	2.062(2)
Cu–N32	2.0505(15)	2.057(3)	2.075(2)
N11–N12	1.3571(19)	1.354(3)	1.364(2)
N21–N22	1.3623(19)	1.358(3)	1.367(3)
N31–N32	1.3625(19)	1.360(3)	1.358(3)
N11–B1	1.548(2)	1.553(3)	1.553(2)
N21–B1	1.557(2)	1.549(3)	1.554(3)
N31–B1	1.553(2)	1.546(4)	1.554(3)

Bond angles (°)			
	<b>6b</b>	<b>6c</b>	<b>6d</b>
Cu1–C1–O11	176.0(2)	177.4(3)	180.0
C1–Cu1–N12	118.66(8)	122.17(11)	125.13(5)
C1–Cu1–N22	129.95(8)	127.99(11)	121.77(12)
C1–Cu1–N32	126.11(7)	124.75(13)	125.74(11)
N12–Cu1–N22	90.60(6)	90.14(9)	90.19(7)
N22–Cu1–N32	88.85(6)	90.58(8)	90.23(7)
N32–Cu1–N12	91.85(6)	90.37(8)	90.31(8)

<sup>a</sup> Bond distances and bond angles are only provided for one of the two crystallographically independent molecules.

**Fig. 1** Projection of part of the structure of  $[\text{Na}(4\text{a}')(\text{H}_2\text{O})]_n$  with displacement ellipsoids plotted at the 50% probability level at 110(2) K. For clarity major parts of the ligand are shown in wireframe and hydrogen atoms have been omitted. Symmetry operations ' =  $\frac{1}{3} - x, y - \frac{1}{2}, \frac{2}{3} - z$ ' and '' =  $[\frac{2}{3} - x, y + \frac{1}{2}, \frac{2}{3} - z]$ .

actual bonding between the sodium ion and the fluorine atoms but merely dipolar interactions.

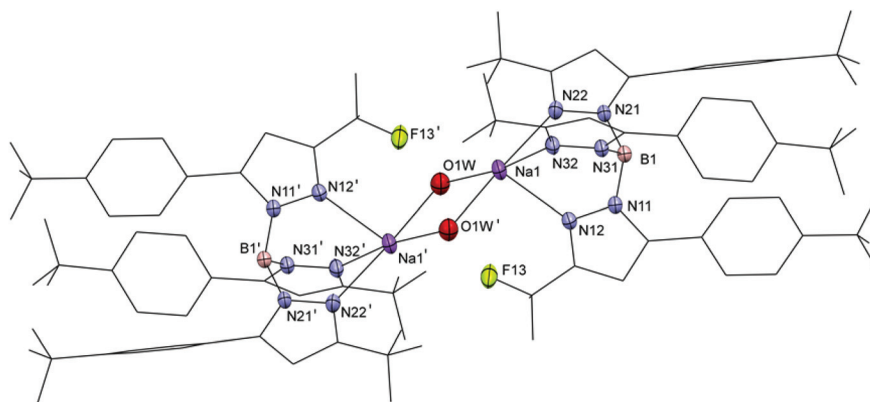
**Na4b** crystallized as a dimer comprising two sodium ions in distorted square-pyramidal coordination geometries. The sodium ions are each coordinated facially by three nitrogen atoms of the trispyrazolylborate ligand and by two bridging water molecules ( $\text{O}_{\text{aqua}}$ ). The Na–N distances range from

2.4355(12) Å to 2.4595(12) Å and the Na– $\text{O}_{\text{aqua}}$  distances are 2.4274(12) Å and 2.4512(12) Å. *trans* to the apical position are one intermolecular and two intramolecular Na…FC short contacts, at distances of respectively 2.930(12), 3.197(4) and 3.116(2) Å. With the foregoing considerations regarding Na…FC short contacts in mind it appears reasonable to consider both the intermolecular and intramolecular Na…FC distances in the structure of **Na4b** as short contacts.

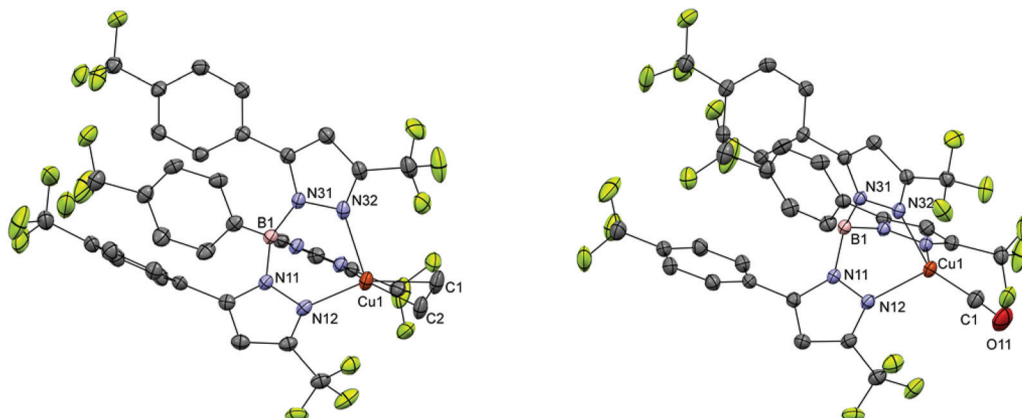
The crystal structures of **Na4a'** and  $[\text{Na}(4\text{e})(\text{H}_2\text{O})]_n$  contain non-bridging water ligands while the crystal structures of **Na4b** and  $[\text{Na}_2(\text{Tp}^{\text{CF}_3, \text{CH}_3})_2(\mu\text{-H}_2\text{O})_2]$  are dinuclear and contain bridging water molecules. The Na– $\text{O}_{\text{aqua}}$  distance in the crystal structure of **Na4a'** of 2.2558(16) Å is comparable to the distance observed in  $[\text{Na}(4\text{e})(\text{H}_2\text{O})]_n$  (2.245(2) Å) but shorter than the distance in  $[\text{Na}_2(\text{Tp}^{\text{CF}_3, \text{CH}_3})_2(\mu\text{-H}_2\text{O})_2]$  (2.417(2) Å), which is closer to the Na– $\text{O}_{\text{aqua}}$  distance found in **Na4b** (2.4274(12) Å and 2.4512(12) Å).<sup>16</sup>

Complexes **5b**, **5c**, **5d**, **5f**, **6b**, **6c**, **6d** and **6f** all have highly similar coordination environments around the copper(I) center. In every case the copper(I) center is coordinated by three nitrogen atoms from the trispyrazolylborate ligand and either the carbon atom of the carbonyl ligand or the ethene ligand in the typical  $\eta^2$  coordination. Bond lengthening or contraction in the carbonyl and ethene ligands as a result of  $\pi$ -backbonding interactions are not evident from the bond lengths observed in the crystal structures and are assumed to be obscured by crystal packing effects. The Cu–N distances in the ethene complexes (**5b**, **5c**, **5d** and **5f**) range from 2.0164(13) to 2.2964(13) Å with in all cases two shorter bonds (2.0164(13)–2.094(3) Å) and one longer bond (2.173(4)–2.2964(13) Å); such asymmetry in the Cu–N bond lengths was also observed in other  $[\text{Cu}^{\text{I}}(\text{Tp}^{\text{R}, \text{R}'})(\text{C}_2\text{H}_4)]$  complexes.<sup>24,25</sup> The Cu–N bond lengths in the carbonyl complexes (**6b**, **6c**, **6d** and **6f**) fall in the range 2.038(2) Å–2.075(2) Å and do not feature the two ranges of Cu–N bond lengths observed in the ethene complexes. The unequal Cu–N bond lengths in the ethene complexes are attributed to the symmetry of ethene. Ethene ligands have a two-fold symmetry while the carbonyl ligands have full rotational symmetry. This means that while the carbonyl copper complexes can retain the approximate threefold symmetry of the trispyrazolylborate ligand, whereas the ethene complexes cannot. The resulting pseudo-twofold symmetry observed in the solid state of the ethene complexes means that  $\pi$ -backbonding into the  $\pi^*$  orbitals of the ethene ligands can not occur with equal contributions from all of the pyrazole rings but mostly from two of the three rings. The resulting discrepancy in charge transfer from the rings to the  $\pi^*$ -orbitals of the ethene ligand explains the presence of two short Cu–N bonds and one long Cu–N bond in complexes **5b**–**g**. The compounds **6b**, **6c** and **6f** crystallized with pseudo threefold rotational symmetry with Cu–C–O angles ranging from 176.0(2) to 178.3(3)°. Complex **6d** crystallized with proper threefold rotational symmetry. The elongated thermal ellipsoid for the oxygen atom of the carbonyl ligand suggests that this ligand is most likely disordered as the C≡O bond may not be perfectly aligned along the threefold axis. The deviation from





**Fig. 2** Projection of the structure of  $[\text{Na}_2(4\mathbf{b})_2(\mu\text{-H}_2\text{O})_2]$  with displacement ellipsoids plotted at the 50% probability level at 150(2) K. For clarity major parts of the ligand are shown in wireframe and hydrogen atoms have been omitted. Symmetry operation ' =  $[-x, 1 - y, -z]$ .



**Fig. 3** Projections of the structures of **5b** (left) and **6b** (right) with displacement ellipsoids plotted at the 50% probability level at 110(2) K. For clarity hydrogen atoms have been omitted. Projections of the structures of the other ethene and carbonyl complexes are available in the ESI.†

rotational symmetry in **6b**, **6c** and **6f** is small and might be the result of crystal packing effects.

### NMR spectroscopy

The  $^1\text{H}$  and  $^{13}\text{C}$ -NMR spectra of compounds **5a'**, **5b-g** and **6a', 6b-g** were recorded in deuterated DCM as it was found to be the only solvent in which the ethene complexes are all soluble and air-stable over longer periods of time (more than 30 minutes).  $\text{CDCl}_3$ , acetone- $d_6$ , THF- $d_8$ , benzene- $d_6$  and  $\text{DMSO-}d_6$  caused solutions of the copper ethene compounds to turn green after some time. The coordinating solvents  $\text{DMSO-}d_6$ , acetone- $d_6$ , THF- $d_8$  and benzene- $d_6$  (a  $\pi$ -donor solvent) all appeared to be competing with the ethene ligands as peaks for free ethene grow over time on  $^1\text{H}$  NMR. The complexes with the less Lewis basic trispyrazolylborate ligands are more stable in solution. Complexes **5b-g** gave very similar spectra with no evidence of anisotropy in the signal for the ethene ligand at room temperature. Considering the asymmetry observed in the crystal structures of complexes **5b-g** splitting of the ethene peaks or at least some peak broadening could be expected. The absence of

such effects appears to indicate fluxional behavior of the ethene ligand within the NMR timescale. In the NMR spectra of **5b-g** the ethene ligands were detected as sharp singlets with chemical shifts between 4.81 and 4.96 ppm which are upfield from the chemical shift of free ethene (5.40 ppm in  $\text{DCM-}d_2$ ) but not as far upfield as the signals observed in complexes of non-fluorinated trispyrazolylborate ligands like  $[\text{Tp}^{\text{Me}_2}\text{Cu}(\text{C}_2\text{H}_4)]$  (4.41 ppm in  $\text{DCM-}d_2$ ).<sup>26</sup> The ethene protons of **5a'** showed as a singlet at 4.44 ppm which was assigned to the asymmetric isomer; a small singlet at 4.99 ppm in the same spectrum was tentatively assigned to the ethene protons of the complex **5a** incorporating the symmetric isomer **[4a]**<sup>-</sup>. The ratio of the integrals of the ethene protons of the symmetric **5a** vs. the asymmetric species **5a'** was approximately 1 : 20. In all ethene complexes peak broadening in the presence of free ethene was not observed, indicating that the exchange of coordinated ethene, if at all, does not occur within the NMR time scale. Solutions of all ethene complexes in  $\text{DCM-}d_2$  show no peak for free ethene even upon prolonged standing. The  $^1\text{H}$ -NMR signals for the borohydrides were



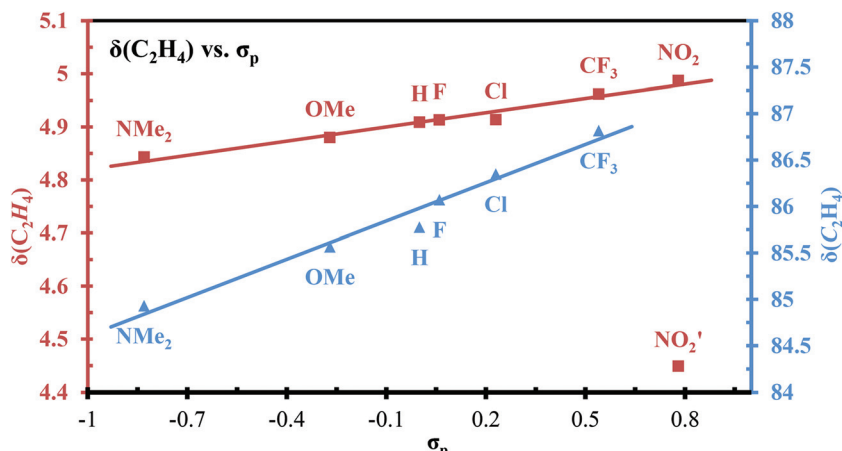


Fig. 4  $^1\text{H}$  chemical shifts (squares, red trend line.  $5\text{a}'$  was not included in the trend line) and  $^{13}\text{C}$  chemical shifts (triangles, blue trend line) of the ethene ligands vs. the Hammett  $\sigma_p$  parameters of the substituents on the trispyrazolylborate ligands in complexes  $5\text{a-g}$  and  $5\text{a}'$ .

observed in the sodium compounds  $\text{Na4a-g}$  as well as the carbonyl and ethene complexes as broadened singlets with chemical shifts in the range 4.3–4.9 ppm. In principle the resonances of the borohydrides should be present as overlapping quartets and smaller septets as a result of splitting by  $^{10}\text{B}$  ( $I = 3$ ) and  $^{11}\text{B}$  ( $I = 3/2$ ). Typically, however, these resonances are observed as (broadened) quartets only in relatively symmetric environments. In less symmetric environments the quadrupoles of the boron nuclei are more pronounced and the borohydride resonances broaden to broad singlets without distinguishable splitting as is the case in this work.<sup>27</sup>

In the  $^{13}\text{C}$ -NMR spectra of complexes  $6\text{a}'$  and  $6\text{b-g}$  no signals were observed for the CO ligands even after increasing the relaxation delay; we ascribe the lack of signals for CO to the low natural abundance of  $^{13}\text{CO}$  and peak broadening.<sup>28,29</sup> The shifts of the resonances trispyrazolylborate ligands show only small differences between the ethene and the carbonyl complexes. For complexes  $5\text{b-g}$  the change in the  $^1\text{H}$ -NMR and  $^{13}\text{C}$ -NMR shifts of the ethene protons and carbon atoms respectively is linearly correlated with the  $\sigma_p$  values of the *para*-substituents on the phenyl groups ( $R^2 = 96\%$  for  $^1\text{H}$  and  $R^2 = 97\%$  for  $^{13}\text{C}$ ; see Fig. 4).

### Infrared spectroscopy

The IR spectra of complexes  $5\text{a}'$ ,  $5\text{b-g}$  and  $6\text{a}'$ ,  $6\text{b-g}$  were recorded in the solid state at  $1\text{ cm}^{-1}$  resolution. Complexes  $6\text{a}'$ ,  $6\text{b-g}$  have CO stretching frequencies in the range  $2096$ – $2120\text{ cm}^{-1}$ , close to that of free CO ( $2143\text{ cm}^{-1}$ ). This indicates that the compounds have relatively Lewis-acidic copper(I) ions compared to other, similar, copper(I) carbonyl complexes like  $[\text{Tp}^{\text{Me}_2}\text{CuCO}]$  ( $2066\text{ cm}^{-1}$ ) and  $[\text{Tp}^{\text{Ph}}\text{CuCO}]$  ( $2080\text{ cm}^{-1}$ , see Table 5 for more examples). The CO stretching frequencies correlate linearly ( $R^2 = 95\%$ ) with the  $\sigma_p$  values of the substituents on the phenyl rings for compounds, with the exception of  $6\text{a}'$  and  $6\text{g}$  (see Fig. 5). For complex  $6\text{a}'$  this is likely because the structure of  $[\text{4a}']^-$  is asymmetric and thus poorly comparable to the other ligands in the series, in

analogy with the ethene resonances in the NMR spectra of compound  $5\text{a}'$ . The CO stretching frequency observed in complex  $6\text{g}$  is significantly higher than the value predicted based on the trend line in Fig. 5 (approximately  $2076\text{ cm}^{-1}$ ) for which no obvious explanation is available. Possibly unusual crystal packing effects cause the CO stretching frequency to be different than predicted, but as all attempts to grow crystals failed this hypothesis remains untested. Unfortunately infrared measurements on a sample of  $6\text{g}$  in DCM were inconclusive as no clear signal was observed until evaporation caused solid  $6\text{g}$  to precipitate. CO stretching typically shows itself in infrared spectroscopy as sharp absorptions due to the symmetry of the vibration. One possible explanation for the absence of an obvious CO stretching peak in solution is a lowering of the symmetry of the bonding environment surrounding the carbonyl ligand. Such lowering of local symmetry could be the result of out-of-axis vibrations of the carbonyl ligand, somewhat similar to motion of the ethene ligands in the NMR experiments and presumably have similarly low barriers. We surmise that liberation of the carbonyl ligand from the copper ion is considerably more pronounced in solution compared to the solid state and that as a result peak broadening causes the weak signal to be drowned out by noise.

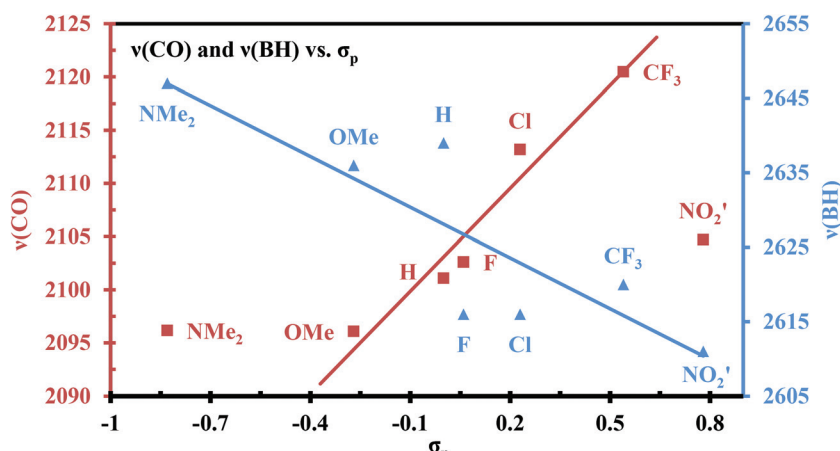
The B–H moieties of the trispyrazolylborate ligands feature IR stretching frequencies that are typically found as small, broad peaks around  $2600\text{ cm}^{-1}$  (see Table 5 and Fig. 5) in Lewis-basic trispyrazolylborate complexes. In order to compare the B–H stretching frequencies of ligands  $4\text{a-g}$  the carbonyl complexes  $6\text{a}'$  and  $6\text{b-g}$  and other, structurally similar, carbonyl complexes were compared. The pseudo-threefold axial symmetries observed in such carbonyl complexes mean that the comparison is not likely to be distorted by local asymmetry. The B–H stretching frequencies of complexes  $6\text{a}'$  and  $6\text{b-g}$  were observed in the region  $2611$ – $2647\text{ cm}^{-1}$ . The B–H stretching frequencies show considerably less linearity vs. the  $\sigma_p$  parameters of the substituents of the trispyrazolylborate ligands



**Table 5** Selected  $^1\text{H}$  and  $^{13}\text{C}$  chemical shifts and bond lengths for the compounds **5a–g** and **6a–g**,  $\text{L} = \text{C}_2\text{H}_4$  or  $\text{CO}$ 

Compound	$\text{C}_2\text{H}_4$ <sup>a</sup> (ppm)		$\nu_{\text{CO}}$ (cm $^{-1}$ )	$\nu_{\text{BH}}$ <sup>f</sup> (cm $^{-1}$ )	$\text{C}=\text{C}$ (Å)	$\text{C}\equiv\text{O}$ (Å)	Ref.
	$^{13}\text{C}$	$^1\text{H}$					
Free $\text{C}_2\text{H}_4/\text{CO}$	123.2	5.40	2143		1.3384(10)	1.13078(9)	30, 31
$[\text{Tp}^{(\text{CF}_3)_2}\text{CuL}]$	89.1 <sup>b</sup>	4.96 <sup>b</sup>	2137	2634	1.325(9)	1.110(5)	18, 20, 32
$[\text{Cu}(4\text{a}')\text{L}]$ <sup>d</sup>	86.4	4.44 (4.99 <sup>e</sup> )	2105	2611			This work
$[\text{Cu}(4\text{b})\text{L}]$	86.8	4.96	2120	2620	1.342(3)	1.124(2)	This work
$[\text{Cu}(4\text{c})\text{L}]$	86.4	4.91	2113	2616	1.340(9)	1.126(4)	This work
$[\text{Tp}^{(\text{CF}_3,\text{Me})}\text{CuL}]$			2107	2575		1.122(3)	33
$[\text{Cu}(4\text{d})\text{L}]$	86.1	4.91	2103	2616	1.350(3)	1.123(8)	This work
$[\text{Tp}^{(\text{CF}_3,\text{Ph})}\text{CuL}]$	85.8	4.91	2101	2639	1.30(1)	1.113(5)	This work, 16
$[\text{Tp}^{\text{CF}_3}\text{CuL}]$	85.8 <sup>c</sup>	4.89 <sup>c</sup>	2100	2507	1.34(1)	1.126(5)	18, 20
$[\text{Cu}(4\text{f})\text{L}]$	85.6	4.81	2096	2636	1.351(8)	1.130(4)	This work
$[\text{Cu}(4\text{g})\text{L}]$	84.9	4.84	2096	2647			This work
$[\text{Tp}^{\text{Me}}\text{CuL}]$		4.41	2066	2500	1.329(9)		26, 34
$[\text{Tp}^{\text{Ph}}\text{CuL}]$	81.6 <sup>b</sup>	3.53	2080	2635	1.381(19)	1.08(1)	8, 35

<sup>a</sup> DCM- $d_2$ . <sup>b</sup>  $\text{CDCl}_3$ . <sup>c</sup>  $\text{C}_6\text{D}_6$ . <sup>d</sup> Asymmetric isomer. <sup>e</sup> Symmetric isomer. <sup>f</sup> As observed in the Cu(I) CO complexes.



**Fig. 5** Infrared CO stretching frequencies (red squares) and BH stretching frequencies (blue triangles) of complexes **6a'**, **6b–g** vs. the Hammett  $\sigma_p$  parameters of the substituents on the trispyrazolylborate ligands. The red linear trend line was calculated including all points except **6a'** and **6g** ( $R^2 = 0.95$ ). The blue linear trend line was calculated including all complexes **6a'**, **6b–g** ( $R^2 = 0.74$ ).

**4a'** and **4b–g** ( $R^2 = 0.74$ ) than the CO stretching frequencies. Intriguingly, compared to the unexpectedly high CO stretching frequency the B–H stretching frequencies of **6a'** and **6g** appear almost exactly on the trend line in Fig. 5. The most Lewis-basic trispyrazolylborate ligands have the highest B–H stretching frequencies as donation of electron density towards the borohydride increase the strength of the hydridic bond.

## Discussion

In this work seven new fluorinated trispyrazolylborate ligands ( $[\text{Tp}^{\text{CF}_3,4-\text{RPh}}]^-$ , R =  $\text{NO}_2$ ,  $\text{CF}_3$ , Cl, F, H,  $\text{MeO}$  and  $\text{NMe}_2$ ) and their copper(I) complexes with carbon monoxide and ethene were prepared. The ligands form an isosteric spectrochemical series based on the electronic properties of the substituents placed on the 4-position of the phenyl rings. The pyrazoles

**3a–g** were readily prepared from ethyl trifluoridoacetate and 4'-substituted acetophenones using a Claisen condensation, followed by cyclization with hydrazine. The synthesis of the trispyrazolylborate ligands requires fine-tuning of the conditions depending on the specific pyrazole using solventless reactions whenever possible; 4-methylanisole was used as a solvent if temperature control was required or the melting point of the pyrazole was too high. To the best of our knowledge  $[\text{4a}']^-$  is only the second example in the literature of a trispyrazolylborate ligand to include the synthetically challenging nitro group, the first being  $[\text{Tp}^{\text{NO}_2}]^-$ .<sup>36</sup>

The trispyrazolylborate ligands studied in this work all formed as their symmetric isomers with the trifluoromethyl groups in the 3-positions of the pyrazole rings, except  $[\text{4a}']^-$  which formed as a mixture of symmetric and asymmetric species. The major product was the asymmetric species in which one of the pyrazole rings is connected to the boron

center with the nitrogen atom adjacent to the phenyl ring. The regioselectivity usually observed in the synthesis of trispyrazolylborate ligands is a result of the relative steric and electronic properties of the substituents on the 3 and 5 positions of the pyrazole, which influence the nucleophilicity of the nitrogen atoms. Typically the bulkiest substituent assumes the 3 positions of the pyrazole rings in the trispyrazolylborate anion. However, if one of the substituents is considerably more electron withdrawing than the other it will assume the 3 position even if the other substituent is bulkier. This regioselectivity is pronounced in ligands with clearly sterically or electronically differentiated substituents like  $[\text{Tp}^{\text{Ph}, \text{Me}}]^-$ ,  $[\text{Tp}^{\text{CF}_3, \text{Me}}]^-$  and  $[\text{Tp}^{\text{CF}_3}]^-$  but breaks down when the steric and electronic differences are small such as in  $[\text{Tp}^{\text{iPr}, \text{Me}}]^-$  which formed a 4:1 mixture of symmetric and the asymmetric isomers.<sup>37</sup> Evidently the strongly electron withdrawing nitro group in  $[\text{4a}']^-$  causes the electronic disparity between the trifluoridomethyl groups and phenyl rings to diminish sufficiently to shift the thermodynamic equilibrium of the system so that the asymmetric isomer is favored over the symmetric isomer. The absence of the formation of asymmetric isomers in the other ligands in the series puts an upper limit on the  $\sigma_p$  parameter of the substituents that can be used to synthesize symmetric  $[\text{Tp}^{\text{CF}_3, 4\text{R-Ph}}]^-$  ligands between +0.54 ( $\text{CF}_3$ ) and +0.78 ( $\text{NO}_2$ ).

The good correlation between the chemical shifts of the ethene protons in complexes **5b-g** vs.  $\sigma_p$  of the substituents on the phenyl rings of the ligands was used to predict the chemical shift of the ethene protons of the symmetric complex **5a**. Based on the trend line in Fig. 4 and the  $\sigma_p$  parameter of the nitro group the signal for the symmetric complex **5a** was predicted to be around 4.98 ppm and indeed a weak singlet was found at 4.99 ppm. Comparison of the integrals of the ethene protons in the symmetric and asymmetric complexes gave an approximate 1:20 ratio. Unfortunately attempts to locate the infrared CO stretching frequency of the symmetric isomer of **6a** using the  $\sigma_p$  parameter of the nitro group and the trend line in Fig. 5 were unsuccessful as the predicted CO stretching frequency for the symmetric isomer **6a** at  $2128 \text{ cm}^{-1}$  would be obscured by the much stronger absorption at  $2105 \text{ cm}^{-1}$  of the asymmetric isomer **6a'**.

More difficult to explain is the unexpectedly high CO stretching frequency observed for **6g**, which was predicted to be at  $2076 \text{ cm}^{-1}$  but instead was found at  $2096 \text{ cm}^{-1}$ , the same value as found for **6f**. It appears implausible that the CO stretching frequencies of **6f** and **6g** are equal as an effect of equal amounts of  $\pi$ -backbonding interactions as the  $^1\text{H}$  and  $^{13}\text{C}$  chemical shifts of the ethene ligands in **5f** and **5g** almost exactly conform to the predicted values. Unfortunately the higher than expected CO stretching frequency of **6g** cannot be conclusively explained by crystal packing effects as infrared spectroscopy on **6g** in solution was inconclusive and all attempts at crystallization failed.

With the exceptions of **6a** and **6g** the predictability of the chemical shifts of the ethene protons in the ethene complexes and CO stretching frequencies in the carbonyl complexes with respect to the  $\sigma_p$  parameters of the substituents of the phenyl

rings in the trispyrazolylborate ligands is excellent. The ability of the ligands to propagate the electron donating or withdrawing effects of the substituents on the phenyl rings over as many as eight bonds and almost one nanometer, is remarkable. The explanation for these long range effects is found in the ability of the ligands to propagate charges by means of resonance structures. Further underscoring the importance of resonance effects is the observation that the B-H stretching frequencies observed in the carbonyl complexes **6a'** and **6b-g** are much less predictable than the CO stretching frequencies because the nitrogen atom adjacent to the borohydride is not part of these resonance structures. We conclude that indeed the  $\pi$ -backbonding ability of the copper(i) centers can be modified extensively through substitution of the trispyrazolylborate ligands without significantly affecting the steric properties of the binding pocket surrounding the metal center.

## Conclusions

In this work the synthesis and characterization of copper(i) complexes comprising a spectrochemical series of trispyrazolylborate ligands and ethene or carbon monoxide have been described. The ethene and carbonyl complexes described in this work are essentially isostructural around their copper(i) centers while their electronic properties vary significantly and predictably based on the Hammett  $\sigma_p$  parameters of the substituents placed on the trispyrazolylborate ligands. The spectroscopic data reveal good correlations between the chemical shifts of the ethene protons and CO stretching frequencies vs. the Hammett  $\sigma_p$  parameters. Notable exceptions to the, otherwise predictable, compounds in this work were the ligand  $[\text{4a}']^-$  and the carbonyl complex **6g**. The ligand  $[\text{4a}']^-$  formed almost exclusively as an asymmetric isomer with one of the pyrazole rings counter-rotated so as to have its 4'-nitrophenyl substituent in the 3-position instead of the trifluoridomethyl group. We attribute the asymmetry of  $[\text{4a}']^-$  to the strongly electron withdrawing properties of the nitro group. Copper complex **6g** shows a surprisingly high CO stretching frequency. That this aberrant result is not due to trend-breaking properties of the ligand is evident from the good correlation between the predicted and found values in the corresponding ethene complex **5g**. The results presented in this work underscore the usefulness of combining the use of more than a single ancillary ligand probe with the systematic study of this promising and important class of ligands.

## Experimental

### General information

All manipulations of air-sensitive compounds were performed in an atmosphere of purified argon gas using standard Schlenk techniques. All solvents were purchased from commercial sources and reagent grade. Solvents used for air-sensitive manipulations were dried and deaerated using a



PureSolv MD 5 Solvent Purification System and stored on 3 Å molecular sieves under argon. When appropriate, glassware was flame dried *in vacuo* immediately prior to use.  $^1\text{H}$  and  $^{13}\text{C}$  NMR spectra were recorded on a Bruker DPX300 spectrometer (300 MHz for  $^1\text{H}$  and 75.44 MHz for  $^{13}\text{C}$ ); Bruker DMX400 spectrometer (400 MHz for  $^1\text{H}$  and 100.6 MHz for  $^{13}\text{C}$ ); Bruker Avance AV500 spectrometer (500 MHz for  $^1\text{H}$ , 160 MHz for  $^{11}\text{B}$  and 126 MHz for  $^{13}\text{C}$ ) or Bruker Avance 600 (600 MHz for  $^1\text{H}$ , 193 MHz for  $^{11}\text{B}$  and 151 MHz for  $^{13}\text{C}$ ). Chemical shifts are given in ppm and referenced using the deuterated solvents as internal references for  $^1\text{H}$  and  $^{13}\text{C}$ .  $^{38}\text{C}$   $^{13}\text{C}$  spectra were recorded using  $^1\text{H}$ -decoupling. Elemental analyses were performed using a Perkin Elmer 2400 series II CHNS/O analyzer or by the Microanalytical laboratory Kolbe in Germany. IR spectra were recorded on a Perkin Elmer UATR Two FT-IR spectrometer set to a resolution of 1  $\text{cm}^{-1}$ . HRMS spectra were recorded on a Thermo Scientific LTQ Orbitrap XL high resolution FT-MS system in MeCN. Intermediates **2a**, **2b**, **2d** and **2f** were synthesized according to a literature procedure using  $\text{KOt-Bu}$  in THF instead of sodium in methanol.<sup>39</sup> **2d** was purified by conversion to the copper(II) diketonate instead of column chromatography as described below.

### Single crystal X-ray crystallography

All reflection intensities were measured at 110(2) K or 150(2) K (only for **Na4b**) using a SuperNova diffractometer (equipped with Atlas detector) with Cu K $\alpha$  radiation ( $\lambda = 1.54178 \text{ \AA}$ ) under the program CrysAlisPro (Versions 1.171.36.32/1.171.37.31 Agilent Technologies, 2013/2014). The same program CrysAlisPro was used to refine the cell dimensions and for data reduction. All structures were solved with the program SHELXS-2013/2014 (Sheldrick, 2008) and were refined on  $F^2$  with SHELXL-2013/2014.<sup>40</sup> Analytical numeric absorption corrections based on a multifaceted crystal model were applied using CrysAlisPro. The temperature of the data collection was controlled using the system Cryojet (manufactured by Oxford Instruments). The H atoms were placed at calculated positions (unless otherwise specified) using the instructions AFIX 43 with isotropic displacement parameters having values 1.2 times  $U_{\text{eq}}$  of the attached C atoms. The H atoms attached to C1/C2, C3/C4 (ethene), and to B1 and B2 were found from difference Fourier maps, and their atomic coordinates were refined freely. The H atoms attached to the water molecules (only for **Na4a'** and **Na4b**) and to the B atoms were found from difference Fourier maps, and their coordinates were refined freely using the DFIX instructions. For **5b**, **5c**, **5d** and **5f**, the C-H and H $\cdots$ H distances of the ethene molecules were restrained using the DFIX instructions. Their isotropic temperature factors were fixed for the ethene molecules (1.2 times  $U_{\text{eq}}$  of the attached C atoms).

**Na4a'.** The structure is ordered. CCDC 1430370 contains the supplementary crystallographic data for **Na4a'**.

**Na4b.** The structure is partly disordered. The six  $-\text{CF}_3$  groups are found disordered over two or three orientations. All occupancies factors can be retrieved from the cif file. When crystals of **Na4b** were flash cooled from RT to 110 K, the

crystal shattered (most likely due to a solid–solid phase transition), resulting to poor quality diffraction. When crystals were cooled from RT to 150 K, the crystals remained stable. CCDC 1430371 contains the supplementary crystallographic data for **Na4b**.

**[Cu(4b)(C<sub>2</sub>H<sub>4</sub>)] (5b).** The structure is mostly ordered. The asymmetric unit contains two crystallographically independent molecules. One of the two crystallographically independent ethene molecules is found to be disordered over two orientations, and the occupancy factor of the major component of the disorder refines to 0.849(5). As the H atoms of the minor component could not be retrieved *via* difference Fourier map, the H atoms of the major component were constrained to have full occupancies. CCDC 1430362 contains the supplementary crystallographic data for **5b**.

**[Cu(4c)(C<sub>2</sub>H<sub>4</sub>)] (5c).** The structure is mostly ordered. The asymmetric unit contains two crystallographically independent molecules. One of the two crystallographically independent ethene molecules is found to be disordered over two orientations, and the occupancy factor of the major component of the disorder refines to 0.502(7). The crystal is racemically twinned, and the BASF scale factor refines to 0.510(14). CCDC 1430363 contains the supplementary crystallographic data for **5c**.

**[Cu(4d)(C<sub>2</sub>H<sub>4</sub>)] (5d).** The structure is ordered. CCDC 1430364 contains the supplementary crystallographic data for **5d**.

**[Cu(4f)C<sub>2</sub>H<sub>4</sub>] (5f).** The structure is mostly ordered. The asymmetric unit contains two crystallographically independent molecules. One of the two crystallographically independent ethene molecules is found to be disordered over two orientations, and the occupancy factor of the major component of the disorder refines to 0.56(3). The positions of the H atoms for the disordered ethene molecule are most likely smeared out, and cannot be retrieved reliably from the data collected. The crystal is racemically twinned, and the BASF scale factor refines to 0.43(3). The riding model AFIX 93 could not be used as this will put the H atoms along the plane defined by Cu2, C3 and C4, which is chemically impossible (the H atoms must be approximately located in the plane perpendicular to the plane defined by Cu2, C3 and C4). CCDC 1430365 contains the supplementary crystallographic data for **5f**.

**[Cu(4b)CO] (6b).** The structure is mostly ordered. One  $-\text{CF}_3$  group is disordered over two orientations. The occupancy factor of the major component of the disorder refines to 0.742(15). CCDC 1430366 contains the supplementary crystallographic data for **6b**.

**[Cu(4c)CO] (6c).** The structure is ordered. The structure was pseudo-merohedrally twinned. The twin relationship is defined by (101/0–10/00–1), which corresponds to a twofold axis along  $\mathbf{a}^*$ . The BASF scale factor refines to 0.3341(9). CCDC 1430367 contains the supplementary crystallographic data for **6c**.

**[Cu(4d)CO] (6d).** The structure is ordered. The Cu complex is found at sites of threefold axial symmetry, and only one third of the complex is crystallographically independent. The



absolute configuration was established by anomalous-dispersion effects in diffraction measurements on the crystal. The Flack and Hooft parameters refine to  $-0.009(6)$  and  $-0.010(6)$ , respectively. CCDC 1430368 contains the supplementary crystallographic data for **6d**.

**[Cu(4f)CO] (6f).** The structure is mostly ordered. One  $-CF_3$  group is disordered over two orientations. The occupancy factor of the major component of the disorder refines to  $0.879(14)$ . The crystal that was mounted on the diffractometer was non-merohedrally twinned. The twin relationship corresponds to a twofold axis around  $0.9995a^* + 0.0115b^* + 0.0288c^*$ . The BASF scale factor refines to  $0.5073(14)$ . CCDC 1430369 contains the supplementary crystallographic data for **6f**.

### Ligand and complex synthesis

**5-(4-Nitrophenyl)-3-(trifluoridomethyl)-1*H*-pyrazole (3a).** **2a** (19.6 g, 75 mmol) was suspended in 250 mL EtOH and the suspension was cooled to  $0\text{ }^\circ\text{C}$  using an ice bath. Hydrazine hydrate (3.9 mL, 80 mmol) was added dropwise with vigorous stirring. After stirring at  $0\text{ }^\circ\text{C}$  for 10 minutes the ice bath was removed and the reaction was stirred at room temperature for approx. 30 minutes, the reaction was then heated to reflux for approx. 17 hours. The EtOH was then evaporated *in vacuo* to a volume of approx. 100 mL. The solution was cooled in an ice bath to  $0\text{ }^\circ\text{C}$  and 20 mL 37% HCl was added dropwise with vigorous stirring. The solution was left to stir for 1 hour and then poured into 600 mL cold water, the resulting suspension was filtered and the residue was washed on the filter with 1 L water. The solids were dried *in vacuo* and then purified by vacuum sublimation ( $215\text{ }^\circ\text{C}$ ,  $10^{-4}$  atm) to yield the product as yellow needles. Yield 14.0 g (73%). M.p. darkened around  $150\text{ }^\circ\text{C}$ , melted  $169\text{--}171\text{ }^\circ\text{C}$ .  $^1\text{H}$  NMR (400 MHz, Acetone-*d*<sub>6</sub>)  $\delta$  13.53 (s, 1H), 8.35 (d, *J* = 8.6 Hz, 2H), 8.12 (d, *J* = 8.6 Hz, 2H), 7.31 (s, 1H).  $^{13}\text{C}$  NMR (75 MHz, Acetone-*d*<sub>6</sub>)  $\delta$  148.71, 144.53 (q, *J* = 37.6 Hz), 143.30, 135.14, 127.47, 125.27, 122.56 (q, *J* = 267.7 Hz), 103.70.

**3-(Trifluoridomethyl)-5-[4-(trifluoridomethyl)phenyl]-1*H*-pyrazole (3b).** **2b** (13.3 g, 56 mmol) was dissolved in 26 mL EtOH and 2.2 mL 37% HCl and the solution was cooled to  $0\text{ }^\circ\text{C}$  with an ice bath. Hydrazine hydrate (1.35 mL, 27.9 mmol) was then slowly added. When the addition was complete the ice bath was removed and the reaction was allowed to stir at room temperature for 20 minutes, it was then heated to reflux for approximately 20 hours. The solvent was removed *in vacuo* and the resulting solids were purified by vacuum sublimation ( $140\text{ }^\circ\text{C}$ ,  $10^{-4}$  atm) to yield the product as a white solid. Yield 5.73 g (77%). M.p.  $143\text{--}145\text{ }^\circ\text{C}$ .  $^1\text{H}$  NMR (300 MHz, DMSO-*d*<sub>6</sub>)  $\delta$  13.58 (s, 1H), 7.82–7.55 (m, 4H), 6.74 (s, 1H).  $^{13}\text{C}$  NMR (100 MHz, Acetone-*d*<sub>6</sub>)  $\delta$  149.24, 143.44 (q, *J* = 38.1 Hz), 134.36, 131.37 (q, *J* = 34.6), 130.48, 125.12, 124.07 (q, *J* = 272.5 Hz), 121.36 (q, *J* = 269.3 Hz), 106.38 (q, *J* = 2.3 Hz), 86.82.

**5-(4-Chloridophenyl)-3-(trifluoridomethyl)-1*H*-pyrazole (3c).** In a flame-dried 500 mL round bottom flask (kept under argon) potassium *tert*-butoxide (13.5 g, 120 mmol) was suspended in 125 mL dry Et<sub>2</sub>O under a dry atmosphere. The sus-

pension was cooled to  $0\text{ }^\circ\text{C}$  on an ice bath and ethyl trifluoroacetate (14.2 mL, 119 mmol) was then added dropwise. 4'-Chloridoacetophenone (13 mL, 100 mmol) in 125 mL dry Et<sub>2</sub>O was then slowly dropped into the first suspension with vigorous stirring. The ice bath was removed and the reaction was stirred at room temperature for 30 minutes, the reaction was then warmed to  $50\text{ }^\circ\text{C}$  and stirred overnight. The mixture was added to 250 mL 0.5 M HCl and separated. The aqueous fraction was extracted twice with Et<sub>2</sub>O ( $2 \times 125$  mL). The combined organic fraction was then washed with water ( $1 \times 200$  mL) and brine ( $1 \times 200$  mL) and dried over Na<sub>2</sub>SO<sub>4</sub>. The solvent was removed *in vacuo* to yield **2c** as a yellow solid. Yield 23.4 g (93%).  $^1\text{H}$  NMR (400 MHz, CDCl<sub>3</sub>)  $\delta$  14.57 (s, 1H, enol-OH), 7.87 (d, *J* = 8.8 Hz, 2H), 7.47 (d, *J* = 8.8 Hz, 2H), 6.53 (s, 1H).  $^{13}\text{C}$  NMR (100 MHz, CDCl<sub>3</sub>)  $\delta$  185.00, 177.42 (q, *J* = 36.6 Hz), 140.75, 131.39, 129.50, 129.02, 117.22 (q, *J* = 283.5 Hz), 92.39 (q, *J* = 2.1 Hz).

**2c** (23.0 g, 93.4 mmol) was dissolved in 100 mL *n*-PrOH and hydrazine hydrate (5.0 mL, 103 mmol) was added dropwise. The reaction was heated to reflux for 2 hours. The reaction was then cooled to  $0\text{ }^\circ\text{C}$  in an ice bath. 10.5 mL 37% HCl was added after which the ice bath was removed and the reaction was heated to  $100\text{ }^\circ\text{C}$  for 30 minutes. The reaction was cooled down to room temperature and the mixture was diluted with 600 mL cold water. The product was collected by filtration and washed with water (approx. 1 L). The product was dried *in vacuo* and purified by vacuum sublimation ( $160\text{ }^\circ\text{C}$ ,  $10^{-4}$  atm) to yield the product as a white solid. Yield 20.4 g (88%). M.p.  $151\text{--}153\text{ }^\circ\text{C}$ .  $^1\text{H}$  NMR (300 MHz, CDCl<sub>3</sub>)  $\delta$  13.09 (s, 1H), 7.46 (d, *J* = 8.6 Hz, 2H), 7.38 (d, *J* = 8.5 Hz, 2H), 6.64 (s, 1H).  $^{13}\text{C}$  NMR (75 MHz, CDCl<sub>3</sub>)  $\delta$  144.55, 143.45 (q, *J* = 37.2 Hz), 135.62, 129.59, 126.87, 126.26, 120.97 (q, *J* = 268.9 Hz), 101.23 (q, *J* = 1.6 Hz).

**5-(4-Fluoridophenyl)-3-(trifluoridomethyl)-1*H*-pyrazole (3d).** **2d** (13.25 g, 56.6 mmol) was dissolved in 100 mL EtOH and cooled in an ice bath to  $0\text{ }^\circ\text{C}$ . Hydrazine hydrate (3.25 mL, 67 mmol) was added dropwise. When the addition was complete the ice bath was removed and the reaction was stirred at room temperature for 1 hour and then heated to reflux overnight. The solvent was removed *in vacuo* to approx. 25 mL. The solution was cooled to  $0\text{ }^\circ\text{C}$  in an ice bath and slowly 20 mL 37% HCl was added. The ice bath was removed and the reaction was heated to reflux for 1 hour after which the reaction mixture was poured into 150 mL cold water. The product was filtered off, dried *in vacuo* and purified by vacuum sublimation ( $100\text{ }^\circ\text{C}$ ,  $10^{-4}$  atm) to yield the product as an off-white solid. Yield 10.95 g (85%). M.p.  $115\text{--}118\text{ }^\circ\text{C}$ .  $^1\text{H}$  NMR (300 MHz, CDCl<sub>3</sub>)  $\delta$  11.32 (s, 1H), 7.53 (dd, *J* = 8.8, 5.1 Hz, 2H), 7.13 (t, *J* = 8.6 Hz, 2H), 6.67 (s, 1H).  $^{13}\text{C}$  NMR (100 MHz, acetone-*d*<sub>6</sub>)  $\delta$  163.45 (d, *J* = 250.1 Hz), 144.64, 143.53 (q, *J* = 38.6 Hz), 127.65 (d, *J* = 8.4 Hz), 124.18, 121.05 (q, *J* = 268.7 Hz), 116.51 (d, *J* = 22.1 Hz), 101.01 (q, *J* = 1.8 Hz).

**5-(4-Methoxyphenyl)-3-(trifluoridomethyl)-1*H*-pyrazole (3f).** **2f** (22.0 g, 89.3 mmol) was dissolved in 250 mL EtOH and cooled to  $0\text{ }^\circ\text{C}$  in an ice bath. Hydrazine hydrate (5.0 mL, 103 mmol) was added dropwise with vigorous stirring. The



reaction was stirred at 0 °C for 30 minutes and then heated to reflux overnight. The EtOH was removed *in vacuo* to a volume of approx. 100 mL and the solution was cooled to 0 °C in an ice bath. 10 mL 37% HCl was slowly added and the reaction was heated to reflux for 1 hour after which the mixture was poured into 500 mL cold water. The product was collected by filtration and was recrystallized from EtOH/water at -20 °C to yield transparent needles. Yield 19.66 g (91%). M.p. 134–136 °C. <sup>1</sup>H NMR (400 MHz, CDCl<sub>3</sub>) δ 11.55 (s, 1H), 7.49 (d, *J* = 8.7 Hz, 2H), 6.95 (d, *J* = 8.7 Hz, 2H), 6.63 (s, 1H), 3.85 (s, 3H). <sup>13</sup>C NMR (100 MHz, CDCl<sub>3</sub>) δ 160.50, 145.04, 143.62 (q, *J* = 38.4 Hz), 127.20, 121.36 (q, *J* = 268.9 Hz), 120.83, 114.71, 100.36 (q, *J* = 2.0 Hz), 55.50.

**N,N-Dimethyl-4-[3-(trifluoridomethyl)-1*H*-pyrazol-5-yl]aniline (3g).** In a flame-dried 250 mL round bottom flask (kept under argon) potassium *tert*-butoxide (8.5 g, 75.8 mmol) was suspended in 70 mL dry Et<sub>2</sub>O. The suspension was cooled to 0 °C on an ice bath and ethyl trifluoroacetate (9.0 mL, 75.6 mmol) was slowly added. 4-Acetyl-*N,N*-dimethylaniline (8.16 g, 50 mmol) was then added in small scoops with vigorous stirring resulting in a clear solution after the last scoop was added. When the addition was complete the reaction was stirred at 0 °C for 45 minutes, the ice bath was then removed and the reaction was allowed to warm to room temperature and stirred until TLC (silica, 5% MeOH in DCM) showed complete consumption of the starting material (approx. 90 minutes). The reaction mixture was then diluted with 30 mL Et<sub>2</sub>O and 100 mL 1 M HCl and the layers were separated. The aqueous layer was extracted with Et<sub>2</sub>O (1 × 50 mL, 2 × 25 mL) and the combined organic layers were washed with water (2 × 25 mL) and brine (1 × 25 mL). The organic fraction was then dried over Na<sub>2</sub>SO<sub>4</sub> and evaporated to dryness *in vacuo* resulting in a red oil that crystallized upon standing to yield 2g as red needles. Yield 12.7 g (98%). <sup>1</sup>H NMR (400 MHz, CDCl<sub>3</sub>) δ 15.27 (s, 1H, *enol-OH*), 7.86 (d, *J* = 9.1 Hz, 2H), 6.71 (d, *J* = 9.1 Hz, 2H), 6.43 (s, 1H), 3.11 (s, 6H). <sup>13</sup>C NMR (75 MHz, Acetone-*d*<sub>6</sub>) δ 187.41, 174.47 (q, *J* = 35.1 Hz), 155.7, 131.24, 119.50, 118.84 (q, *J* = 281.8 Hz), 112.12, 91.15 (q, *J* = 2.3 Hz), 40.08.

**2g** (12.7 g, 50 mmol) was dissolved in 125 mL EtOH and cooled to 0 °C in an ice bath. Hydrazine hydrate (2.7 mL, 56 mmol) was added dropwise, when the addition was complete the ice bath was removed and the reaction was stirred at room temperature for 30 minutes before the reaction was heated to reflux for approx. 17 hours. The solvent was evaporated *in vacuo* and 100 mL 3 M HCl was added. The solution was heated to reflux for 5 minutes, cooled to room temperature and diluted with 500 mL aqueous saturated NaHCO<sub>3</sub> solution and 100 mL DCM. The layers were separated and the aqueous fraction was extracted with DCM (2 × 50 mL). The combined organic fractions were washed with water (2 × 100 mL) and brine (1 × 100 mL) and dried over Na<sub>2</sub>SO<sub>4</sub>. The solvent was removed *in vacuo* and the product was recrystallized from hot MeOH/water to yield a white solid. Yield (10.2 g, 80%). M.p. 179–181 °C. <sup>1</sup>H NMR (300 MHz, DMSO) δ 12.03 (s, 1H), 7.45 (d, *J* = 8.9 Hz, 2H), 6.73 (d, *J* = 8.9 Hz, 2H), 6.62 (s, 1H), 3.00 (s, 6H). <sup>1</sup>H NMR (300 MHz, Acetone-*d*<sub>6</sub>) δ 12.86 (s, 1H), 7.66 (d, *J* =

9.1 Hz, 2H), 6.85–6.78 (m, 3H), 3.00 (d, *J* = 1.6 Hz, 6H). <sup>13</sup>C NMR (75 MHz, CDCl<sub>3</sub>) δ 151.07, 145.70, 143.79 (q, *J* = 35.7 Hz), 126.79, 121.54 (q, *J* = 268.9 Hz), 115.91, 112.44, 99.50, 40.35.

**Sodium hydridobis[3-(trifluoridomethyl)-5-(4-nitrophenyl)-pyrazol-1-yl][3-(4-nitrophenyl)-5-(trifluoridomethyl)-pyrazol-1-yl]borate (Na4a).** **3a** (3.00 g, 11.67 mmol) and NaBH<sub>4</sub> (126 mg, 3.33 mmol) are mixed in 10 mL 4-methylanisole and were placed under argon in a flame-dried Schlenk vessel fitted with a flame dried glass tube (30 cm). The mixture was heated to 120 °C for 2 hours to form the di-substituted borate species, bubbling was observed using an oil bubbler and became sporadic after approx. 90 minutes. The temperature was then raised to 180 °C for three hours. After approx. 1.5 hours bubbling stopped but BH<sub>4</sub><sup>-</sup> was still observed on <sup>1</sup>H NMR, after 3 hours BH<sub>4</sub><sup>-</sup> was no longer observed. The solvent was then removed *in vacuo* to yield the product as a brown oil. The product was purified by extensive trituration with Et<sub>2</sub>O to yield the product as an off-white solid which was recrystallized from DCM/pentane. Yield 1.11 g (40%). M.p. 225 °C (decomposition). <sup>1</sup>H NMR (500 MHz, CD<sub>2</sub>Cl<sub>2</sub>) δ 8.19 (d, *J* = 8.4 Hz, 2H), 8.05 (d, *J* = 8.3 Hz, 4H), 7.77 (d, *J* = 8.4 Hz, 2H), 7.32 (d, *J* = 8.3 Hz, 4H), 6.97 (s, 1H), 6.66 (s, 2H), 4.59 (bs, 1H). <sup>13</sup>C NMR (75 MHz, acetone-*d*<sub>6</sub>) δ 150.17, 148.75, 148.06, 147.82, 143.21 (q, *J* = 36.8 Hz), 140.64, 139.37, 138.63 (q, *J* = 38.1 Hz), 130.98, 127.09, 124.57, 123.50, 123.15 (q, *J* = 267.6 Hz), 121.73 (q, *J* = 269.0 Hz), 106.79 (q, *J* = 3.0 Hz), 105.73 (q, *J* = 1.8 Hz). HRMS (ESI neg.) *m/z* calcd For [M<sup>-</sup>] (=C30H16BF9N9O6^-) 780.11674 found 780.11778.

**Sodium hydrotris[3-(trifluoridomethyl)-5-(4-{trifluoridomethyl}phenyl)pyrazol-1-yl]borate (Na4b).** **3b** (7.75 g, 27.7 mmol) and NaBH<sub>4</sub> (0.202 g, 5.35 mmol) were mixed and placed under argon in a flame-dried Schlenk vessel fitted with a flame-dried condenser (not cooled with water). The mixture was heated 200 °C for 4 hours, **3b** subliming in the glassware was occasionally molten using a heat gun to return it to the reaction mixture. The remaining pyrazole was removed from the product by extensive vacuum sublimation at 160 °C. The product was dissolved in DCM and filtered over celite to remove insoluble byproducts. The solvent was removed *in vacuo* to yield the product as a white powder. Yield 3.59 g (77%). M.p. 212–214 °C. <sup>1</sup>H NMR (300 MHz, acetone-*d*<sub>6</sub>) δ 7.37 (d, *J* = 8.4 Hz, 6H), 7.23 (d, *J* = 8.4 Hz, 6H), 6.74 (s, 3H), 4.47 (bs, 1H). <sup>13</sup>C NMR (75 MHz, acetone-*d*<sub>6</sub>) δ 150.38, 143.75 (q, *J* = 36.7 Hz), 136.43, 131.05, 125.36 (m), 122.97 (q, *J* = 267.7 Hz), 125.25 (q, *J* = 271.8 Hz), 105.34 (q, *J* = 1.9 Hz). HRMS (ESI neg.) *m/z* calcd For [M<sup>-</sup>] (=C33H16BF18N6^-) 849.12366 found 849.12078.

**Sodium hydrotris[3-(trifluoridomethyl)-5-(4-chloridophenyl)-pyrazol-1-yl]borate (Na4c).** **3c** (5.00 g, 20.3 mmol) and NaBH<sub>4</sub> (0.247 g, 6.53 mmol) were placed in a flame-dried Schlenk vessel fitted with a flame-dried glass tube (30 cm) and placed under argon. The mixture was heated at once to 180 °C and held at this temperature for 2.5 hours. The condenser was then replaced with a cold finger and vacuum was applied (10<sup>-4</sup> atm). The pyrazole was left to sublime out of the product over-



night. When no more of the pyrazole deposited on the cold finger the cold finger was removed and the product was dissolved in THF (approx. 50 mL) and filtered over celite to remove insoluble byproducts. The THF was evaporated *in vacuo* and the resulting solids were washed with boiling heptane (4 × 20 mL) and Et<sub>2</sub>O (1 × 20 mL) on a glass frit leaving the pure product as a 1:1 mixture of the Et<sub>2</sub>O and THF adducts. Yield 3.2 g (62%). M.p. 192 °C (decomposition). <sup>1</sup>H NMR (400 MHz, CDCl<sub>3</sub>) δ 7.00 (d, *J* = 8.4 Hz, 6H), 6.77 (d, *J* = 8.4 Hz, 6H), 6.44 (s, 3H), 4.57 (bs, 1H), 3.92 (ddd, *J* = 6.5, 4.2, 2.7 Hz, 2H, coord. THF), 3.55 (q, *J* = 7.0 Hz, 2H, coord. Et<sub>2</sub>O), 2.02–1.96 (m, 2H, coord. THF), 1.24 (t, *J* = 7.0 Hz, 3H, coord. Et<sub>2</sub>O). <sup>13</sup>C NMR (100 MHz, Acetone-d<sub>6</sub>) δ 150.38, 143.25, 134.56, 131.69, 131.07, 128.52, 121.44, 104.67. HRMS (ESI neg.) *m/z* calcd For [M<sup>-</sup>] (=C30H16BCl3F9N6^-) 747.04459 found 747.04770.

**Sodium hydrotris[3-(trifluoridomethyl)-5-(4-fluoridophenyl)-pyrazol-1-yl]borate (Na4d).** **3d** (5.00 g, 21.7 mmol) and NaBH<sub>4</sub> (0.267 g, 7.06 mmol) were placed in a flame-dried Schlenk vessel fitted with a flame dried glass tube (30 cm) and placed under argon. The mixture was heated to 180 °C for 3 hours and 20 minutes. The temperature was lowered to 140 °C and the condenser was replaced with a cold finger, leftover pyrazole was removed by vacuum sublimation. When no more pyrazole deposited on the cold finger the residue was dissolved in toluene (50 mL) and filtered over celite to remove insoluble byproducts. The toluene was evaporated *in vacuo* and the remaining solids were dissolved in acetone and evaporated to dryness. Yield 1.72 g (53%). M.p. 190 °C (decomposition). <sup>1</sup>H NMR (400 MHz, CDCl<sub>3</sub>) δ 6.88 (dd, *J* = 8.6, 5.4 Hz, 6H), 6.72 (t, *J* = 8.6 Hz, 6H), 6.44 (s, 3H), 4.61 (bs, 1H), 2.34 (s, 6H, coord. acetone). <sup>13</sup>C NMR (100 MHz, CDCl<sub>3</sub>) δ 220.47 (coord. acetone), 162.75 (d, *J* = 248.1 Hz), 150.18, 142.70 (q, *J* = 36.6 Hz), 131.60 (d, *J* = 8.3 Hz), 127.79 (d, *J* = 3.3 Hz), 122.20 (d, *J* = 267.8 Hz), 114.74 (d, *J* = 21.7 Hz), 103.84 (q, *J* = 1.5 Hz), 65.98 (coord. THF), 31.31 (coord. acetone), 23.06 (coord. THF). HRMS (ESI neg.) *m/z* calcd for [M<sup>-</sup>] (=C30H16BF12N6^-) 699.13324 found 699.13033.

**Sodium hydrotris[3-(trifluoridomethyl)-5-(4-methoxyphenyl)-pyrazol-1-yl]borate (Na4f).** **3f** (6.867 g, 28.35 mmol) and NaBH<sub>4</sub> (0.346 g, 9.15 mmol) were placed in a flame-dried Schlenk vessel fitted with a flame-dried glass tube (30 cm) and placed under argon. The mixture was heated to 180 °C for 3 hours. The product was dissolved in boiling toluene (50 mL) and filtered, the filtrate was diluted with 200 mL heptane and left to stand while the product crystallized. Filtration yielded a white powder. Recrystallization from acetone yielded the product as its acetone adduct. Yield 3.75 g (53%). M.p. 206 °C (decomposition). <sup>1</sup>H NMR (300 MHz, CDCl<sub>3</sub>) δ 6.85 (d, *J* = 8.6 Hz, 6H), 6.52 (d, *J* = 8.6 Hz, 6H), 6.42 (s, 3H), 4.69 (bs, 1H), 3.79 (s, 9H), 2.33 (s, coord. acetone). <sup>13</sup>C NMR (75 MHz, CDCl<sub>3</sub>) δ 212.24 (coord. Acetone), 159.40, 151.06, 142.34 (q, *J* = 36.2 Hz), 131.26, 122.37 (q, *J* = 268.0 Hz), 113.08, 103.10 (q, *J* = 1.6 Hz), 55.14, 31.27 (coord. acetone). HRMS (ESI neg.) *m/z* calcd for [M<sup>-</sup>] (=C33H25BF9N6O3^-) 735.19320 found 735.19424.

**Sodium hydrotris[3-(trifluoridomethyl)-5-(4-{N,N-dimethylamino}phenyl)pyrazol-1-yl]borate (Na4g).** **3g** (6.00 g, 23.5 mmol) and NaBH<sub>4</sub> (0.286 g, 7.56 mmol) were suspended in 20 mL 4-methylanisole in a flame dried Schlenk flask fitted with a flame-dried glass tube (30 cm) and placed under argon. The mixture was heated to reflux for 4 hours. The mixture was allowed to cool to room temperature and 200 mL petroleum ether was added. The diluted mixture was stirred in an ice bath for approx. 30 minutes and then filtered. The filtrate was washed with MeCN (6 × 25 mL) and then dissolved in acetone (50 mL). The acetone was removed *in vacuo* to yield the product as a white powder. The product formed was the acetone adduct. Yield 1.80 g (28%). Decomposed around 230 °C. <sup>1</sup>H NMR (300 MHz, Acetone-d<sub>6</sub>) δ 6.82 (d, *J* = 8.9 Hz, 6H), 6.46 (s, 3H), 6.33 (d, *J* = 8.9 Hz, 6H), 4.93 (bs, 1H), 2.93 (s, 18H). <sup>13</sup>C NMR (75 MHz, CDCl<sub>3</sub>) δ 152.86, 150.81, 142.83 (q, *J* = 36.1 Hz), 131.28, 123.37 (q, *J* = 267.4 Hz), 120.35, 112.07, 103.18 (q, *J* = 1.8 Hz), 40.22. HRMS (ESI neg.) *m/z* calcd for [M<sup>-</sup>] (=C36H34BF9N9^-) 774.28810 found 774.28907.

#### General method for synthesis of the copper(i) ethene complexes (5a', 5b–g)

Na4a' or Na4b–g (50 mg) was dissolved in 5 mL dry, degassed DCM. The solution was bubbled with ethene for 2 minutes. CuI (1.05 eq.) was then added and the solution was saturated/bubbled with ethene for another 2 minutes. The flask was then stoppered and left to stir at medium speed overnight to form a white suspension. The suspension was filtered using a syringe filter (0.45 µm PTFE) and the solvent was removed *in vacuo* to yield the products as white powders. The complexes could be further purified by recrystallization from DCM/pentane at –20 °C under an ethene atmosphere to yield colorless blocks except for 5a' which was brown.

**[Cu(4a')(C<sub>2</sub>H<sub>4</sub>)]** (5a'). Yield 42 mg (83%), yellow solid. <sup>1</sup>H NMR (300 MHz, CD<sub>2</sub>Cl<sub>2</sub>) δ 8.37 (d, *J* = 8.9 Hz, 2H), 8.07 (d, *J* = 8.9 Hz, 4H), 7.76 (d, *J* = 8.9 Hz, 2H), 7.36 (d, *J* = 8.9 Hz, 4H), 6.87 (s, 1H), 6.72 (s, 2H), 4.54 (bs, 1H), 4.44 (s, 4H). <sup>13</sup>C NMR (126 MHz, CD<sub>2</sub>Cl<sub>2</sub>) δ 152.46, 148.84, 148.67, 148.42, 143.52 (q, *J* = 38.4 Hz), 139.00 (q, *J* = 39.1 Hz), 138.60, 136.99, 131.11, 129.57, 124.32, 123.36, 121.23 (q, *J* = 269.3 Hz), 120.05 (q, *J* = 269.7 Hz), 106.76 (q, *J* = 2.5 Hz), 86.37. Elemental analysis calc. (%) for C<sub>32</sub>H<sub>20</sub>BCuF<sub>9</sub>N<sub>6</sub>O<sub>6</sub>·1.5H<sub>2</sub>O·1.0DCM (found): C 40.29 (41.20), H 2.56 (2.8), N 12.81 (12.32).

**[Cu(4b)(C<sub>2</sub>H<sub>4</sub>)]** (5b). Yield 49 mg (91%). <sup>1</sup>H NMR (400 MHz, CD<sub>2</sub>Cl<sub>2</sub>) δ 7.22 (d, *J* = 8.4 Hz, 6H), 7.00 (d, *J* = 8.3 Hz, 6H), 6.69 (s, 3H), 4.96 (s, 4H), 4.43 (bs, 1H). <sup>13</sup>C NMR (100 MHz, CD<sub>2</sub>Cl<sub>2</sub>) δ 149.24, 143.44 (q, *J* = 38.2 Hz), 134.36, 131.37 (q, *J* = 33.8 Hz), 130.48, 125.12, 124.07 (q, *J* = 272.4 Hz), 121.36 (q, *J* = 269.3 Hz), 106.38 (q, *J* = 2.4 Hz), 86.82. Elemental analysis calc. (%) for C<sub>35</sub>H<sub>20</sub>BCuF<sub>18</sub>N<sub>6</sub> (found): C 44.68 (44.63), H 2.14 (2.01), N 8.93 (8.91).

**[Cu(4c)(C<sub>2</sub>H<sub>4</sub>)]** (5c). Yield 47 mg (94%). <sup>1</sup>H NMR (600 MHz, CD<sub>2</sub>Cl<sub>2</sub>) δ 7.02 (d, *J* = 8.5 Hz, 6H), 6.80 (d, *J* = 8.5 Hz, 6H), 6.61 (s, 3H), 4.92 (s, 4H), 4.47 (bs, 1H). <sup>13</sup>C NMR (151 MHz, CD<sub>2</sub>Cl<sub>2</sub>) δ 149.57, 143.14 (q, *J* = 38.0 Hz), 135.49, 131.41, 129.26, 128.49, 121.45 (q, *J* = 269.0 Hz), 105.93 (q, *J* = 2.5 Hz), 86.35. Elemental



analysis calc. (%) for  $C_{32}H_{20}BCl_3CuF_9N_6$  (found): C 45.74 (46.37), H 2.40 (2.77), N 10.00 (9.87).

**[Cu(4d)(C<sub>2</sub>H<sub>4</sub>)] (5d).** Yield 36.6 mg (73%). <sup>1</sup>H NMR (300 MHz, CD<sub>2</sub>Cl<sub>2</sub>)  $\delta$  6.95–6.83 (m, 6H), 6.74 (t,  $J$  = 8.5 Hz, 6H), 6.60 (s, 3H), 4.91 (s, 4H), 4.54 (bs, 1H). <sup>13</sup>C NMR (126 MHz, CD<sub>2</sub>Cl<sub>2</sub>)  $\delta$  163.24 (d,  $J$  = 249.8 Hz), 149.65, 142.95 (q,  $J$  = 37.9 Hz), 131.86 (d,  $J$  = 8.3 Hz), 126.88 (d,  $J$  = 3.4 Hz), 121.41 (q,  $J$  = 268.7 Hz), 115.13 (d,  $J$  = 21.6 Hz), 105.92 (q,  $J$  = 2.4 Hz), 86.07. Elemental analysis calc. (%) for  $C_{32}H_{20}BCuF_{12}N_6\cdot H_2O$  (found): C 47.52 (47.48), H 2.74 (2.57), N 10.39 (10.23).

**[Cu(4f)(C<sub>2</sub>H<sub>4</sub>)] (5f).** Yield 43.5 mg (86%). <sup>1</sup>H NMR (400 MHz, CD<sub>2</sub>Cl<sub>2</sub>)  $\delta$  6.81 (d,  $J$  = 8.7 Hz, 6H), 6.56 (s, 3H), 6.51 (d,  $J$  = 8.7 Hz, 6H), 4.88 (s, 4H), 4.62 (bs, 1H), 3.78 (s, 9H). <sup>13</sup>C NMR (100 MHz, CD<sub>2</sub>Cl<sub>2</sub>)  $\delta$  160.29, 150.69, 142.51 (q,  $J$  = 37.6 Hz), 131.47, 123.22, 121.67 (q,  $J$  = 268.8 Hz), 113.55, 105.25 (q,  $J$  = 2.5 Hz), 85.57, 55.47. Elemental analysis calc. (%) for  $C_{35}H_{29}BCuF_9N_6O_3$  (found): C 51.01 (50.83), H 3.47 (3.53), N 10.12 (10.16).

**[Cu(4g)(C<sub>2</sub>H<sub>4</sub>)] (5g).** Yield 40 mg (79%). <sup>1</sup>H NMR (500 MHz, CD<sub>2</sub>Cl<sub>2</sub>)  $\delta$  6.75 (d,  $J$  = 8.7 Hz, 6H), 6.51 (s, 3H), 6.26 (d,  $J$  = 8.7 Hz, 6H), 4.84 (s, 4H), 4.77 (bs, 1H), 2.92 (s, 18H). <sup>13</sup>C NMR (126 MHz, CD<sub>2</sub>Cl<sub>2</sub>)  $\delta$  151.56, 150.56, 142.41 (q,  $J$  = 37.5 Hz), 130.98, 121.82 (q,  $J$  = 268.6 Hz), 118.42, 111.46, 104.57 (q,  $J$  = 2.6 Hz), 84.94, 40.12. Elemental analysis calc. (%) for  $C_{38}H_{38}BCuF_9N_9\cdot 0.2C_5H_{12}$  (found): C 53.20 (53.40), H 4.62 (4.61), N 14.32 (14.07).

### General method for the synthesis of the copper(I)-CO complexes (6a', 6b-g)

Na4a' or Na4b-g (50 mg) was dissolved in dry, degassed DCM. The solution was bubbled with argon for 2 minutes and placed in a glass-lined autoclave (volume 12 mL). CuI (1.05 eq.) was added and the autoclave was closed and purged with dry nitrogen gas. The autoclave was then pressurized with carbon monoxide to at least 5 atm and left to stir at room temperature for 20 hours. The autoclave was then bubbled with dry nitrogen gas to remove unreacted carbon monoxide. The resulting white suspension was filtered using a syringe filter (0.45  $\mu$ m, PTFE) and the solvent was removed *in vacuo* to leave behind the product as a white powder except in the case of 6a' which was brown.

**[Cu(4a')(CO)] (6a').** Yield 36 mg (56%). <sup>1</sup>H NMR (300 MHz, CD<sub>2</sub>Cl<sub>2</sub>)  $\delta$  8.40 (d,  $J$  = 8.5 Hz, 2H), 8.09 (d,  $J$  = 8.5 Hz, 4H), 7.91 (d,  $J$  = 8.4 Hz, 2H), 7.35 (d,  $J$  = 8.4 Hz, 4H), 6.91 (s, 1H), 6.72 (s, 2H), 4.40 (bs, 1H). <sup>13</sup>C NMR (100 MHz, CD<sub>2</sub>Cl<sub>2</sub>)  $\delta$  151.85, 149.09, 148.89, 148.66, 143.50 (q,  $J$  = 38.1 Hz), 139.41 (q,  $J$  = 39.5 Hz), 138.09, 136.57, 131.11, 129.05, 124.53, 123.49, 121.12 (q,  $J$  = 269.2 Hz), 119.97 (q,  $J$  = 269.5 Hz), 107.27, 106.33. IR (cm<sup>-1</sup>): 2611 (w, BH stretching), 2105 (s, CO stretching). Elemental analysis calc. (%) for  $C_{31}H_{16}BCuF_9N_9O_7$  (found): C 42.71 (42.88), H 1.85 (1.97), N 14.46 (14.34).

**[Cu(4b)(CO)] (6b).** Yield 36 mg (86%). <sup>1</sup>H NMR (300 MHz, CD<sub>2</sub>Cl<sub>2</sub>)  $\delta$  7.25 (d,  $J$  = 8.4 Hz, 6H), 6.99 (d,  $J$  = 8.4 Hz, 6H), 6.69 (s, 3H), 4.29 (bs, 1H). <sup>13</sup>C NMR (75 MHz, CD<sub>2</sub>Cl<sub>2</sub>)  $\delta$  149.02, 143.00 (q,  $J$  = 37.8 Hz), 133.62, 131.20 (q,  $J$  = 33.7 Hz), 130.13, 124.82 (tt,  $J$  = 3.9, 1.9 Hz), 123.70 (q,  $J$  = 277.1 Hz), 120.87 (q,

$J$  = 269.3), 105.24 (q,  $J$  = 1.8 Hz). IR (cm<sup>-1</sup>): 2620 (w, BH stretching), 2120 (s, CO stretching). Elemental analysis calc. (%) for  $C_{34}H_{16}BCuF_{18}N_6O$  (found): C 43.40 (43.20), H 1.71 (1.79), N 8.93 (8.83).

**[Cu(4c)(CO)] (6c).** Yield 45 mg (90%). <sup>1</sup>H NMR (300 MHz, CD<sub>2</sub>Cl<sub>2</sub>)  $\delta$  7.04 (d,  $J$  = 8.3 Hz, 6H), 6.79 (d,  $J$  = 8.5 Hz, 6H), 6.61 (s, 3H), 4.33 (bs, 1H). <sup>13</sup>C NMR (75 MHz, CD<sub>2</sub>Cl<sub>2</sub>)  $\delta$  149.71, 143.06 (q,  $J$  = 37.8 Hz), 135.66, 131.38, 128.85, 128.54, 121.31 (q,  $J$  = 269.2 Hz), 105.17 (q,  $J$  = 2.0 Hz). IR (cm<sup>-1</sup>): 2616 (w, BH stretching), 2113 (s, CO stretching). Elemental analysis calc. (%) for  $C_{31}H_{16}BCl_3CuF_9N_6O$  (found): C 44.31 (44.43), H 1.92 (1.99), N 10.00 (9.95).

**[Cu(4d)(CO)] (6d).** Yield 34 mg (72%). <sup>1</sup>H NMR (300 MHz, CD<sub>2</sub>Cl<sub>2</sub>)  $\delta$  6.89 (dd,  $J$  = 8.9, 5.3 Hz, 6H), 6.76 (dd,  $J$  = 9.0, 8.5 Hz, 6H), 6.59 (s, 3H), 4.45 (bs, 1H). <sup>13</sup>C NMR (75 MHz, CD<sub>2</sub>Cl<sub>2</sub>)  $\delta$  163.41 (d,  $J$  = 247.5 Hz), 149.86, 142.92 (q,  $J$  = 37.8 Hz), 131.93 (d,  $J$  = 8.5 Hz), 126.54 (d,  $J$  = 3.5 Hz), 121.34 (q,  $J$  = 268.9 Hz), 115.28 (d,  $J$  = 21.8 Hz), 105.25 (q,  $J$  = 1.8 Hz). IR (cm<sup>-1</sup>): 2616 (w, BH stretching), 2103 (s, CO stretching). Elemental analysis calc. (%) for  $C_{31}H_{16}BCuF_{12}N_6O\cdot 0.2C_5H_{12}$  (found): C 47.73 (48.03), H 2.30 (2.30), N 10.44 (10.65).

**[Cu(4f)(CO)] (6f).** Yield 45 mg (89%). <sup>1</sup>H NMR (300 MHz, CD<sub>2</sub>Cl<sub>2</sub>)  $\delta$  6.79 (d,  $J$  = 8.6 Hz, 6H), 6.55 (s, 3H), 6.52 (d,  $J$  = 8.7 Hz, 6H), 4.50 (bs, 1H), 3.79 (s, 9H). <sup>13</sup>C NMR (75 MHz, CD<sub>2</sub>Cl<sub>2</sub>)  $\delta$  160.07, 150.46, 142.24 (q,  $J$  = 37.3 Hz), 131.11, 122.45, 121.18 (q,  $J$  = 269.0 Hz), 113.23, 104.12 (q,  $J$  = 1.8 Hz), 55.17. IR (cm<sup>-1</sup>): 2636 (w, BH stretching), 2096 (s, CO stretching). Elemental analysis calc. (%) for  $C_{34}H_{25}BCuF_9N_6O_4$  (found): C 49.38 (49.11), H 3.05 (3.14), N 10.16 (9.99).

**[Cu(4g)(CO)] (6g).** Yield 51 mg (86%). <sup>1</sup>H NMR (300 MHz, CD<sub>2</sub>Cl<sub>2</sub>)  $\delta$  6.74 (d,  $J$  = 8.8 Hz, 6H), 6.50 (s, 3H), 6.28 (d,  $J$  = 8.8 Hz, 6H), 4.66 (bs, 1H), 2.92 (s, 18H). <sup>13</sup>C NMR (75 MHz, CD<sub>2</sub>Cl<sub>2</sub>)  $\delta$  151.32, 150.34, 142.14, 130.62, 117.65, 111.12, 103.41 (q,  $J$  = 1.8 Hz), 40.15. CO and CF<sub>3</sub> could not be resolved due to the low solubility of the complex. IR (cm<sup>-1</sup>): 2647 (w, BH stretching), 2096 (s, CO stretching). Elemental analysis calc. (%) for  $C_{37}H_{34}BCuF_9N_9O\cdot H_2O$  (found): C 50.27 (50.63), H 4.10 (4.24), N 14.26 (13.90).

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