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Ambidentate bonding and electrochemical implications of pincer-type pyridylidene amide ligands in complexes of nickel, cobalt and zinc†

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Pincer-type tridentate pyridyl bis(pyridylidene amide) (pyPYA₂) ligand systems were coordinated to the Earth-abundant first row transition metals nickel, cobalt and zinc. A one-pot synthesis in water/ethanol afforded octahedral homoleptic bis-PYA complexes, [M(pyPYA₂)₂](PF₆)₂, whereas five-coordinate mono-PYA dichloride complexes, [M(pyPYA₂)Cl₂], were obtained upon slow addition of the ligand to the metal chlorides in DMF. Electrochemical measurements further revealed a facile oxidation of the metal centers from Ni²⁺ to Ni⁴⁺ and Co²⁺ to Co³⁺, respectively, while the Zn²⁺ system was redox inactive. These experiments further allowed for quantification of the much stronger electron donor properties of neutral *N,N,N*-tridentate pyPYA₂ pincer ligands as compared to terpy. Remarkably, *ortho*-PYA pincer ligands feature amide coordination to the metal center via oxygen or nitrogen. This ambidentate ligand binding constitutes another mode of donor flexibility of the PYA ligand system, complementing the resonance structure dynamics established previously. NMR spectroscopic and MS analysis reveal that the *meta*-PYA ligand undergoes selective deuteration when coordinated to cobalt. This reactivity suggests the potential of this ligand as a transient proton reservoir for HX bond activation and, moreover, indicates the relevance of several resonance structures and therefore supports the notion that *meta*-PYA ligands are mesoionic.

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Introduction

The large-scale production of bulk chemicals requires the utilization of inexpensive materials, and many industrially relevant processes rely on heterogeneous catalysts based on Earth-abundant metals. Prominent examples are the iron oxide catalysts used in the Haber–Bosch process,^{1–3} nickel catalysts in steam reforming for the production of syngas^{4–6} or RANEY® nickel catalysts utilized in hydrogenation reactions.^{7,8} In contrast, the development of homogeneous catalysts has long focused on complexes based on platinum group and coinage metals.⁹ These noble metals feature excellent reactivity and stability properties, though their high price¹⁰ and scarcity^{11,12} as well as the high toxicity¹³ demand for more sustainable solutions and have led to extensive research activities aiming for the replacement of noble metals by first row transition metals.¹⁴ As a consequence, iron catalysis has made major progress over the last years,¹⁵ including for example the develop-

ment of highly efficient hydrogenation catalysts that are competitive with the best ruthenium catalysts.¹⁶ Likewise manganese(i) pincer complexes have been explored as attractive alternative to established ruthenium- and iridium-based catalysts in (de)hydrogenation as well as in C–C and C–X bond-forming reactions.¹⁷ Also nickel complexes have gained much attention recently, *e.g.* as low-cost replacement for palladium in cross-coupling reactions.¹⁸

We have recently developed a new pincer platform based on pyridylidene amide (PYA) ligands (**A**, Fig. 1), which imparted high catalytic activity to ruthenium centers in transfer hydrogenation catalysis.^{19,20} Based on the relatively hard yet formally neutral amide donor site of these PYA ligands, we sought to expand the application of this ligand system towards first row transition metals in view of providing a more sustainable alternative to noble metals. PYA ligands have been shown to induce attractive catalytic properties to iridium, ruthenium^{21–26} and palladium²⁷ complexes, in parts by stabilizing high-valent oxidation states and in other parts due to their unique electron donor flexibility.^{28–30} Despite these attractive features, no first row transition metal complexes with PYA ligands have been reported, but analogous pyridylidene amine (PYE) ligands have been coordinated to nickel centers by the groups of Johnson³¹ and Douthwaite³² (**B** and **C**, Fig. 1). Here we report nickel, cobalt, and zinc complexes con-

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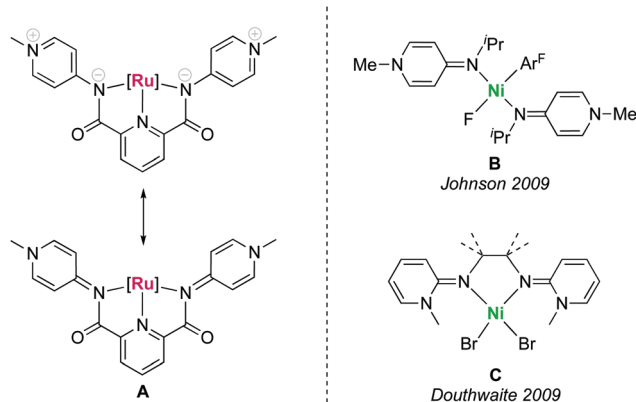


Fig. 1 Pincer pyridylidene amide (PYA) ruthenium complex A with two limiting PYA resonance structures, and pyridylidene amine (PYE) nickel complexes B and C reported earlier.

tanting the pyridyl bis(PYA) (pyPYA_2) pincer ligand and disclose some unique reactivity properties, including the linkage isomerism between *N*- to *O*-coordination of the PYA ligand as well as isotope exchange reactions at the *meta*-PYA unit that support a mesoionic notion of this ligand.

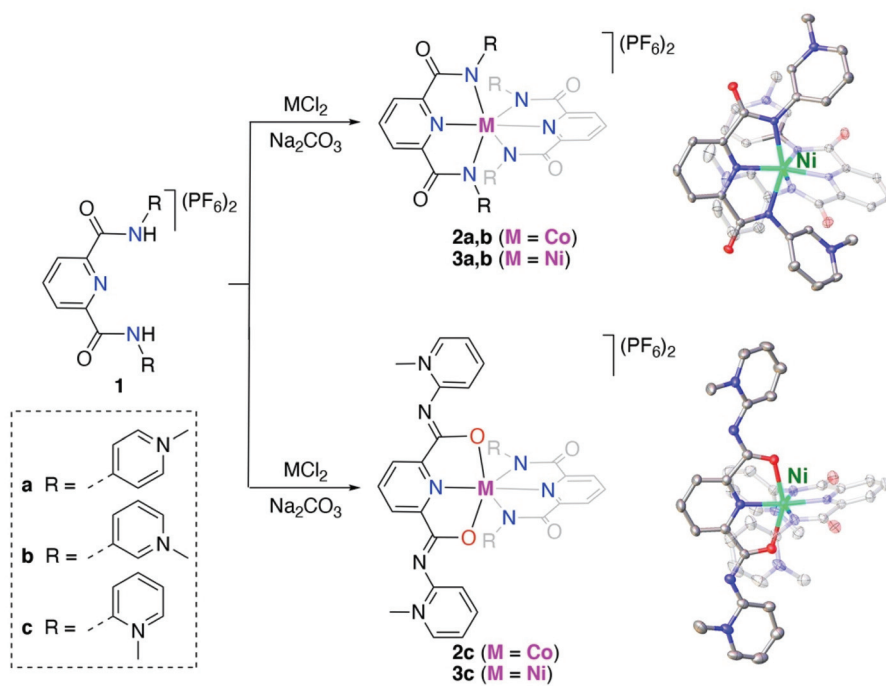
Results and discussion

Synthesis and characterization of bis(PYA-pincer) complexes

The known *para*- and *meta*-pyPYA₂ ligand precursor salts **1a** and **1b** were prepared according to standard procedures.^{19,33}

Similarly, the *ortho*-pyPYA₂ ligand precursor **1c** was prepared from 2-aminopyridine and 2,6-pyridinedicarbonyl dichloride followed by selective methylation of both aminopyridine sites and anion metathesis. Notably, methylation required an excess of MeOTf (5 eq.) to be quantitative, while the use of MeI, which was sufficient for methylation of the *para* and *meta* analogues, resulted in only partial methylation. Complexation to cobalt and nickel was accomplished by reacting the dicationic ligand salts **1a–c** with the corresponding chloride salt (CoCl_2 , NiCl_2) in the presence of Na_2CO_3 as a base (Scheme 1). Complexation under these conditions consistently afforded the octahedral homoleptic complexes $[\text{M}(\text{pyPYA}_2)_2](\text{PF}_6)_2$ ($\text{M} = \text{Co}$, **2a–c**; $\text{M} = \text{Ni}$, **3a–c**) even when a large excess of the metal salt was used. This reactivity pattern contrasts the analogous ruthenation, which exclusively afforded mono-ligated $[\text{Ru}(\text{pyPYA}_2)(\text{MeCN})_3]^{2+}$ complexes. Complexes **2** and **3** are paramagnetic and NMR silent, their quantitative elemental composition, structure and purity was therefore confirmed by CHN microanalysis, single crystal X-ray diffraction, and HR ESI-MS. The latter showed the monocationic complex fragments $[\text{M} - \text{PF}_6]^+$, *e.g.* at 898.1733 amu for **2a** (calcd 898.1733).

Crystalllographic analysis revealed ambidentate coordination of the *ortho*-pyPYA₂ ligand in **2c** and **3c**, as one of the ligands is bound to the metal center in the typical $\kappa^3\text{-}N,N,N$ bonding mode, while the other ligand binds with the amide through oxygen in a $\kappa^3\text{-}O,N,O$ coordination mode (*cf.* Scheme 1, Table S1, Fig. S1†). Both *meta*-pyPYA₂ ligands in complexes **2b** and **3b** show the expected $\kappa^3\text{-}N,N,N$ coordination through the amide nitrogens and the central pyridine as observed for related dianionic pyridine dicarboxamidate



Scheme 1 Synthesis of Co(II) and Ni(II) bis-pyPYA₂ pincer complexes **2a–c** and **3a–c** in $\text{H}_2\text{O}/\text{EtOH}$ (1:1 v/v). Crystal structures of complex cations **3b** and **3c**; all ellipsoids at 50% probability level (hydrogen atoms and co-crystallized solvent molecules omitted for clarity).



systems.^{34–36} Such a linkage isomerism of PYA ligands has been proposed,³⁷ yet never established so far³⁸ and presents a further type of flexibility of the PYA system. Notably, the same *ortho*-pyPYA₂ ligand displayed the classic κ^3 -*N,N,N*-coordination mode when bound to ruthenium(II) in [Ru(*ortho*-pyPYA₂)(MeCN)₃](PF₆)₂ (Fig. S2†). Likewise, pyridine dicarboxamide systems with *ortho*-dimethylated aryl substituents show exclusively κ^3 -*N,N,N*-bonding.^{35,39} The κ^3 -*O,N,O* coordination mode is unprecedented and presumably a consequence of the Lewis acidity of the first row transition metal centers and their ensuing oxophilicity, combined with the unique donor flexi-

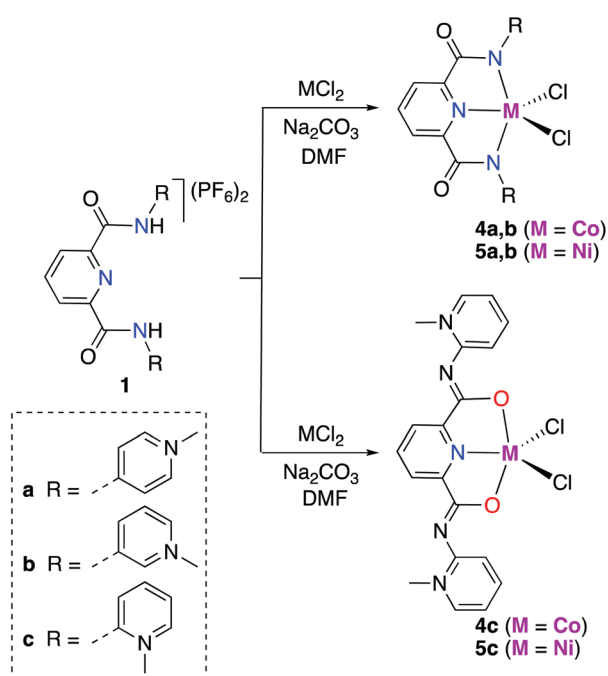
bility of the formally neutral and sterically constrained *ortho*-PYA donor site, and distinguishes these ligands from the more commonly used pyridine carboxamide systems.

Synthesis and characterization of mono(PYA-pincer) complexes

The synthesis of cobalt and nickel mono-PYA complexes, [M(pyPYA₂)Cl₂] complexes **4a–c** (M = Co) and **5a–c** (M = Ni; Scheme 2), was accomplished by slow addition of a filtered DMF solution containing the ligand and Na₂CO₃ to a DMF solution containing the metal chloride. The slow addition of ligand as well as the choice of solvent played a crucial role. Initial attempts to control the reaction of the PYA ligands **1a–c** with MCl₂ by quenching the reaction at early reaction times or by using a large excess of the metal salt (up to 10 eq.) invariably resulted in the formation of the homoleptic [M(pyPYA₂)₂]²⁺ compounds **2** and **3**.

Formation of complexes **4** and **5** was established by CHN microanalysis as well as HR ESI-MS, which showed the monocationic complex fragments [M – Cl]⁺, *e.g.* at 441.0409 amu for **4a** (calcd 441.0403). The set of cobalt complexes **4a–c** was furthermore characterized by single crystal X-ray diffraction analysis (Fig. 2). The complexes feature a strongly distorted trigonal bipyramidal arrangement ($\tau_5 = 0.45$ –0.49),⁴⁰ which is in good agreement with the reported analogous structure of [Co(terpy)Cl₂] (terpy = terpyridine).⁴¹ The *para*- and *meta*-PYA pincer ligands in complexes **4a,b** bind to the metal center in a κ^3 -*N,N,N* coordination mode whereas the *ortho*-PYA pincer ligand in **4c** is again κ^3 -*O,N,O* coordinated as observed for the bis-PYA pincer complexes. Notably, steric constraints should be less relevant when the PYA pincer ligand is bound to the CoCl₂ fragment than in bis-PYA pincer complexes, indicating an electronic stabilization of the PYA coordination through oxygen.

The *para*- and *meta*-PYA pincer ligands in complexes **4a,b** are almost planar, *i.e.* the dihedral angle between the PYA heterocycles and central pyridine ring are only 8.5°, 9.8° in **4a** and



Scheme 2 Synthesis of Ni(II) and Co(II) [M(pyPYA₂)Cl₂] complexes **4a–c** and **5a–c** by slow addition of deprotonated ligand at room temperature.

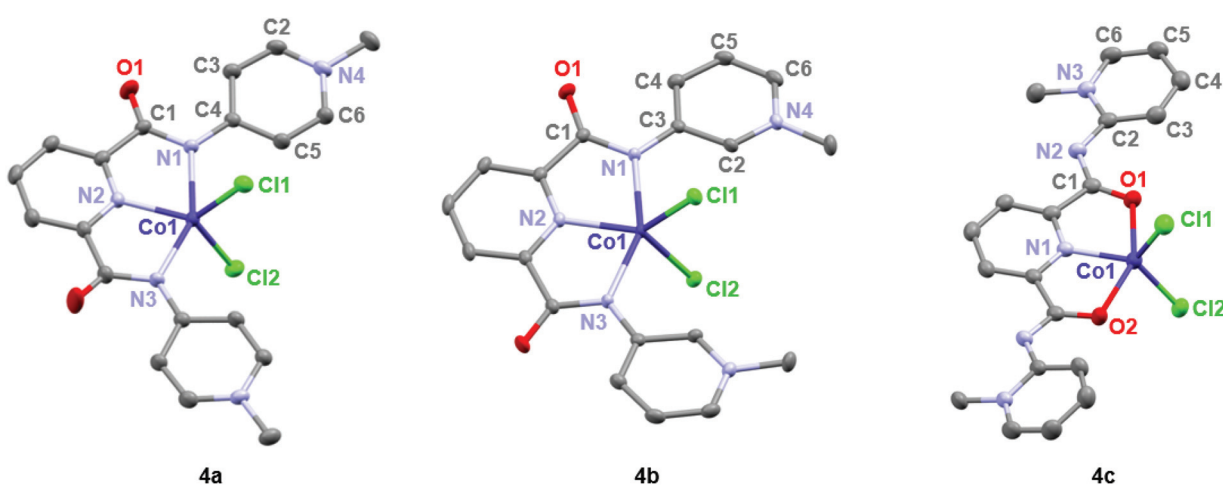


Fig. 2 ORTEP representations of mono-pyPYA₂ cobalt complexes **4a–c**; all ellipsoids at 50% probability level (hydrogen atoms and co-crystallized solvent molecules omitted for clarity).



Table 1 Selected bond lengths (Å) and angles (°) of mono-pyPYA₂ complex cations **4a–c**

	4a	4b	4c	
Co1–N1	2.227(2)	2.185(1)	Co1–O1	2.217(2)
Co1–N2	2.020(2)	2.005(1)	Co1–N1	2.018(1)
Co1–N3	2.217(2)	2.185(1)	Co1–O2	2.225(2)
Co1–Cl1	2.315(1)	2.330(1)	Co1–Cl1	2.249(1)
Co1–Cl2	2.296(1)	2.330(1)	Co1–Cl2	2.262(1)
N1–C1 ^a	1.360(3)	1.347(2)	N2–C1 ^a	1.331(3)
C1–O1 ^a	1.231(2)	1.244(2)	C1–O1 ^a	1.254(2)
N1–C3/4 ^a	1.373(2)	1.391(2)	N2–C2 ^a	1.353(3)
C3–C4 ^a	1.412(3)	1.406(2)	C2–C3 ^a	1.405(3)
C4–C5 ^a	1.419(3)	1.388(2)	C3–C4 ^a	1.368(4)
C5–C6 ^a	1.364(3)	1.366(2)	C4–C5 ^a	1.393(4)
C6–N4 ^a	1.345(3)	1.349(2)	C5–C6 ^a	1.353(3)
N4–C2 ^a	1.353(2)	1.342(2)	C6–N3 ^a	1.356(3)
C2–C3 ^a	1.365(3)	1.399(2)	N3–C2 ^a	1.375(3)
N1–Co1–N3	153.1(1)	154.1(1)	O1–Co1–O2	152.0(2)
θ PYA–pyr ^b	8.5 & 9.8	10.4 & 10.4	θ PYA–pyr ^b	11.3 & 25.8
τ_5^c	0.46	0.49	τ_5^c	0.45

^a Average of two bonds. ^b Dihedral angle θ between the PYA heterocycles and the pyridyl-amide plane. ^c See ref. 40.

10.4° in **4b** respectively (Table 1), which is distinctly different when compared to the correspondent homoleptic complex variation [Co(*meta*-pyPYA₂)₂](PF₆)₂ which show a 55° twist between the PYA and central pyridyl planes. This high planarity may hint to a significant π -contribution of the Namide–C_{PY}A bonds (N1–C3/4) in both complexes. In such a model, the amide nitrogen is acting predominantly as neutral L-type N-donor ligand, presumably as a consequence of the higher electron density at the formally neutral metal center in complexes **4** and **5** compared to the dicationic center in the bis (pincer) complexes **2** and **3**. In agreement with a more pronounced L-type and π -acidic binding mode of the amide nitrogen, the *para*-PYA heterocycles of complex **4a** reveal a pronounced diene-imine resonance form with localized double and single bonds (average C2–C3 and C5–C6 1.365(4) Å, average C3–C4 and C4–C5 bonds (1.415(7) Å; Table 1). This bond alteration implies considerable contribution of the imine resonance structure of the PYA unit (*cf.* bottom structure of **A** in Fig. 1).

The κ^3 -O,N,O coordination mode of the *ortho*-PYA pincer ligand in mono-pyPYA₂ complex **4c** suggests this complex to be an intermediate *en route* to the formation of the heteroleptic bis-pyPYA₂ pincer complex **2c**. The stronger coordination of

the O,N,O donor set may then accommodate a second PYA pincer ligand in the more typical κ^3 -N,N,N mode.

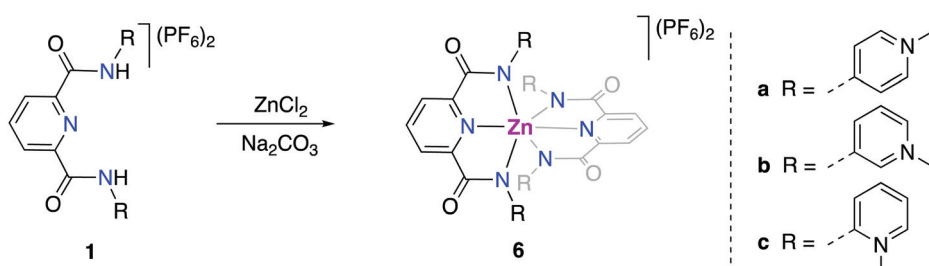
Dynamic linkage isomerism of the *ortho*-PYA-pincer ligand system

For better spectroscopic analysis in solution, the analogous zinc(II) complexes [Zn(pyPYA₂)₂](PF₆)₂ **6a–c** were prepared by reacting ligands **1a–c** with 0.5 eq. ZnCl₂ under mildly basic conditions (Scheme 3). In contrast to the paramagnetic and NMR-silent octahedral Co(II) and Ni(II) complexes **2** and **3**, complexes **6a–c** are diamagnetic and allow for NMR spectroscopic characterization. Ligand coordination to the Zn^{II} center resulted in the loss of the NH amide proton resonances and in a marked upfield shift of the aromatic H_{PY}A proton resonances, *e.g.* from δ_H = 9.76 (H⁶_{PY}A) in *meta*-PYA ligand **1b** to δ = 8.43 in complex **6b**.

All zinc complexes **6a–c** feature one distinct set of PYA ligand proton resonances, including N–CH₃ (δ_H around 4), pyridyl (δ_H 8.25–8.70) and pyridylidene (δ_H 7.08–8.43) signals. This symmetric arrangement indicates that both pyPYA₂ ligands in each zinc complex are coordinated in the same mode, most presumably the typical κ^3 -N,N,N-coordination based on the structures of the cobalt and nickel analogues and previously described structures.^{19,42–45}

The symmetric ligand bonding in **6c** containing *ortho*-pyPYA₂ ligands contrasts the solid state bonding established for the analogous nickel and cobalt complexes **2c** and **3c** comprised of a mixture of κ^3 -N,N,N and κ^3 -O,N,O pincer coordination. NMR spectroscopic analysis of complex **6c** at low temperature indicated decoalescence of the signals into two sets in a 9:1 integral ratio, indicating the presence of two species (Fig. 3). Based on the six inequivalent aromatic resonances, the major species was assigned to the homoleptic bis- κ^3 -N,N,N-coordinated pincer isomer. The minor isomer features at least ten inequivalent proton signals, which were therefore attributed to mixed κ^3 -N,N,N/ κ^3 -O,N,O-coordination isomer (*cf.* structures of **2c**, **3c**). These measurements therefore suggest that at room temperature, complex **6c** undergoes rapid κ N- κ O-amide isomerization.

The dynamic process of this complex was investigated by variable temperature ¹H NMR spectroscopic experiments in CD₂Cl₂ in the +20 to –80 °C temperature range (Fig. S3†). Decoalescence of most of the pyridyl and PYA proton resonances was observed around –40 °C, apart from the PYA H⁶

**Scheme 3** Synthesis of [Zn(pyPYA₂)₂](PF₆)₂ complexes **6a–c**. (a) 0.5 eq. ZnCl₂; 3 eq. Na₂CO₃ in MeCN.

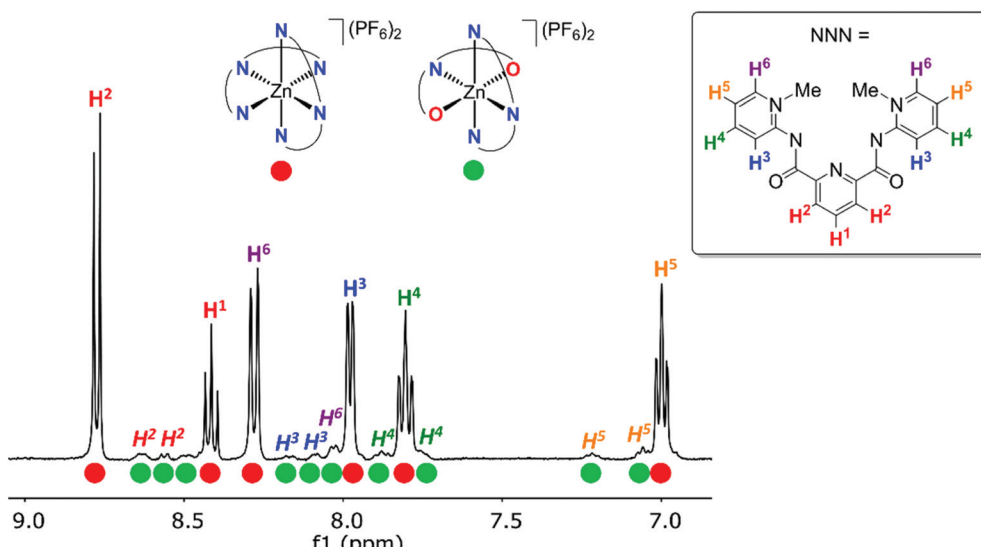


Fig. 3 ^1H NMR spectrum of $[\text{Zn}(\text{ortho-pyPYA})_2](\text{PF}_6)_2$ **6c** in CD_2Cl_2 at $-80\text{ }^\circ\text{C}$, revealing two different complexes in 9 : 1 ratio. The major species was assigned to the bis-($\kappa^3\text{-N,N,N}$)-coordination isomer (red circles), the minor species to the mixed $\kappa^3\text{-N,N,N/}\kappa^3\text{-O,N,O}$ -coordination isomer (green circles).

signal, which undergoes the largest shift and features a coalescence temperature T_c around $-20\text{ }^\circ\text{C}$. Based on these values and applying the Eyring equation, an average activation energy $\Delta G^\ddagger = 11.4 \pm 0.3\text{ kcal mol}^{-1}$ was calculated for this PYA linkage isomerism (Table S2†).^{46,47} A similar behavior was noted in CD_3CN (Fig. S4†), though due to the higher melting point of this solvent, the decoalescence was not resolved. Nonetheless, these experiments suggest that the dynamic bonding is not significantly depending on the solvent.

The bonding mode in the bis-(pyPYA₂) complexes was further analyzed by solid state infrared (IR) spectroscopy of the pure compounds. The ligand precursor **1a** features three characteristic absorption bands for the amide carbonyl and PYA-imine units in the $1580\text{--}1700\text{ cm}^{-1}$ region (Table 2, Fig. S5–S11†).⁴⁸ Upon metalation, the two lower bands around 1590 and 1645 cm^{-1} remain essentially unchanged, while the

high energy band at 1706 cm^{-1} shifts characteristically to 1617 ($\pm 1\text{ cm}^{-1}$) presumably due to metal coordination to the amide unit (Fig. 4a). Likewise, the strong absorption above 1700 cm^{-1} for ligand precursors **1b** and **1c** shifts to lower energy upon metalation, though assignments are more complicated as all bands shift considerably (Fig. 4b and Fig. S5†). It is worth noting that the shift is independent of the metal center and that the IR spectra are essentially identical for Co(II), Ni(II), and Zn(II) complexes for the *para*- and *meta*-pyPYA₂ ligand even in the low frequency area. This high similarity suggests an identical coordination mode for all three metal complexes. For the *ortho*-pyPYA₂ series, however, subtle differences are noted in the $1550\text{--}1650\text{ cm}^{-1}$ range which distinguish the Zn from the Ni and Co systems (Fig. 4c). These differences may point to a different ligand behavior, which may be associated with the $\kappa^3\text{-N,N,N}$ vs. $\kappa^3\text{-O,N,O}$ linkage isomerism established

Table 2 Characteristic IR stretch frequencies of all compounds in the carbonyl/imine range between $1580\text{--}1700\text{ cm}^{-1}$ and redox potentials of all compounds

Entry	Ligand motif	Bonding partner	$\nu_{\text{C=O/C=N}} [\text{cm}^{-1}]$	$E_{1/2} (\text{M}^{\text{II/III}})^a [\text{V}]$	$E_{1/2} (\text{M}^{\text{III/IV}})^a [\text{V}]$
1	<i>para</i> -PYA	H	1a	1593, 1646, 1706	—
2	<i>para</i> -PYA	Co	2a	1586, 1617, 1642	0.20 (90)
3	<i>para</i> -PYA	Ni	3a	1587, 1617, 1645	1.20 (80)
4	<i>para</i> -PYA	Zn	6a	1584, 1618, 1646	—
5	<i>meta</i> -PYA	H	1b	1695	—
6	<i>meta</i> -PYA	Co	2b	1567	0.01 (80)
7	<i>meta</i> -PYA	Ni	3b	1564	1.03 (80)
8	<i>meta</i> -PYA	Zn	6b	1566	—
9	<i>ortho</i> -PYA	H	1c	1583, 1637, 1721	—
10	<i>ortho</i> -PYA	Co	2c	1597, 1628	0.40 (100)
11	<i>ortho</i> -PYA	Ni	3c	1598, 1627	1.34 (100)
12	<i>ortho</i> -PYA	Zn	6c	1587, 1630	—

^a Potentials in MeCN vs. SSCE at 100 mV s^{-1} scan rate using the Fc^+/Fc couple as standard; $E_{1/2} = +0.43\text{ V}$ and $(n\text{Bu}_4\text{N})\text{PF}_6$ as electrolyte, $0\text{--}2.2\text{ V}$ sweep range; $\Delta E_p = |E_{pc} - E_{pa}|$ [mV] in parenthesis.



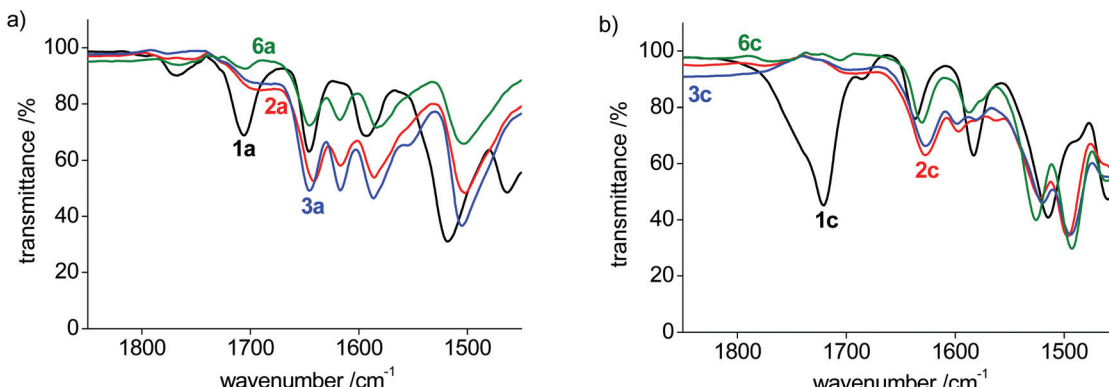


Fig. 4 IR spectra of (a) *para*-pyPYA₂ pincer complex series; (b) *ortho*-pyPYA₂ pincer complex series.

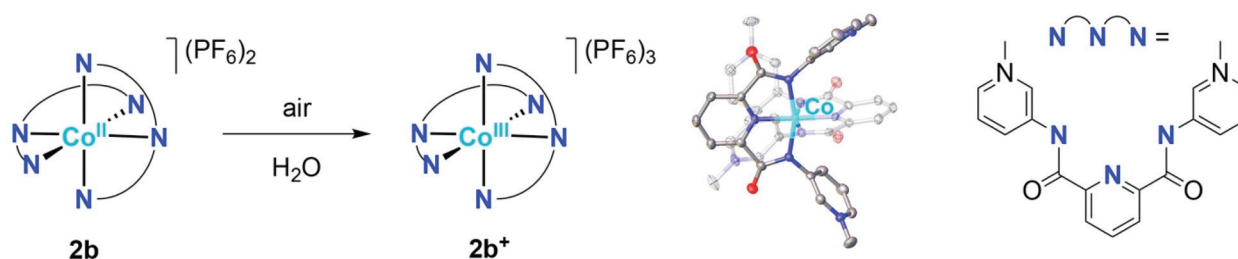
for the $[\text{Zn}(\text{ortho-pyPYA}_2)_2](\text{PF}_6)_2$ complex **6c** by NMR spectroscopy. IR spectroscopy therefore does not conclusively indicate whether this linkage isomerism is exclusively occurring at Zn^{2+} , or whether it is more general for *ortho*-pyPYA₂ ligands and also takes place at Co^{2+} and Ni^{2+} . The exclusive $\kappa^3\text{-O},\text{N},\text{O}$ bonding in the solid state of the latter two complexes (*cf.* structures of **2c**–**5c**) might be induced by lower solubility or a preferential crystal packing of the mixed coordination isomer, leading to a dynamic kinetic resolution of the two isomers through crystallization.

Redox properties of homoleptic bis-PYA pincer complexes

The orange *meta*-pyPYA₂ cobalt(II) complex **2b** oxidized to the green Co(III) analogue **2b**⁺ under aerobic conditions in a saturated aqueous solution (Scheme 4). The oxidation state of the cobalt center in **2b**⁺ was unambiguously identified by crystallographic analysis due to the 0.2 Å shorter Co–N bond distances compared to **2b** as well as the presence of three PF_6^- counter ions. This oxidation under mild conditions indicates strong donor properties of the PYA pincer ligand in these cobalt and nickel complexes. Therefore, the electrochemical properties of the complexes were investigated by cyclic voltammetry.

Cobalt complexes **2a**–**c** show reversible redox processes in the 0 to +0.5 V potential range that were assigned to metal-centered Co^{2+} to Co^{3+} oxidations (all potentials *vs.* Ag/AgCl; Fig. 5). The octahedral complex $[\text{Co}(\text{terpy})_2](\text{PF}_6)_2$ ⁴⁹ was also included to compare the donor strength of the formally neutral triden-

tate PYA pincer ligands with a well-studied reference compound. The reversibility of the $\text{M}^{\text{II/III}}$ oxidation processes in complexes **2** and **3** is supported by the linear relationship of anodic (I_{pa}) and cathodic peak currents (I_{pc}) *vs.* the square root of the scan rate (Fig. S12†). This linearity together with the small peak potential difference $\Delta E_p < 105$ mV (Table 2) indicates a diffusion limited electron transfer process.^{50–52} The oxidation potential is dependent on the PYA substitution pattern and is lowest for cobalt coordinated to the *meta*-PYA pincer ligand ($E_{1/2} = 0.01$ V in **2b**) and increases with *para*-PYA ($E_{1/2} = 0.20$ V in **2a**)⁵³ and even more with *ortho*-PYA ($E_{1/2} = 0.40$ V in **2c**). It is noteworthy that $[\text{Co}(\text{ortho-pyPYA}_2)](\text{PF}_6)_2$ complex **2c** features two consecutive redox processes in the potential range of 0.3–0.6 V, which might be caused by possible linkage isomerism in solution and the presence of two coordination isomers (exclusively $\kappa^3\text{-N},\text{N},\text{N}$ -coordinated ligand and mixed coordination mode, *cf.* NMR studies of analogous *ortho*-PYA zinc complex above). These half wave potentials indicate an increase of ligand donor strength along the series *ortho*-PYA < *para*-PYA < *meta*-PYA, a trend that has been previously observed for $[\text{Ru}(\text{pyPYA}_2)\text{L}_3]^{2+}$ and $[\text{IrCp}^*(\text{phenyl-PYA})\text{L}]^+$ complexes.^{19,54} This trend points to a strongly π -basic character of the *meta*-PYA donor, while the donor properties of the *ortho*-PYA unit are probably attenuated by the vicinity of the formally positively charged pyridinium nitrogen. When compared to the redox properties of $[\text{Co}(\text{terpy})_2]^{2+}$ ($E_{1/2} = 0.32$ V)⁵⁵ the *ortho*-PYA pincer ligand is less donating than terpy,



Scheme 4 Oxidation of $[\text{Co}(\text{meta-pyPYA}_2)_2](\text{PF}_6)_2$ **2b** in H_2O under aerobic conditions, to the Co(III) complex **2b**⁺. Crystal structure of complex cation **2b**⁺; all ellipsoids at 50% probability level (hydrogen atoms and co-crystallized solvent molecules omitted for clarity).



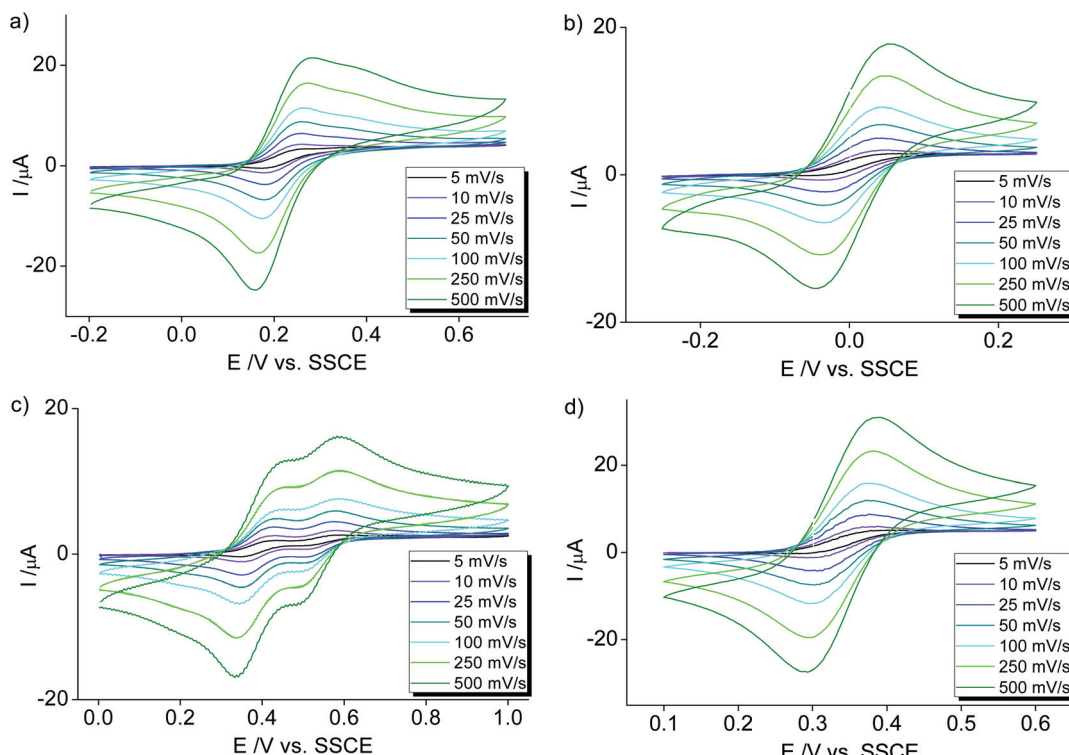


Fig. 5 Cyclic voltammetry measurements of homoleptic $[\text{Co}(\text{pyPYA}_2)_2](\text{PF}_6)_2$ complexes in MeCN: (a) **2a** $[\text{Co}(\text{para-pyPYA}_2)_2](\text{PF}_6)_2$; (b) **2b** $[\text{Co}(\text{meta-pyPYA}_2)_2](\text{PF}_6)_2$; (c) **2c** $[\text{Co}(\text{ortho-pyPYA}_2)_2](\text{PF}_6)_2$; (d) $[\text{Co}(\text{terpy})_2](\text{PF}_6)_2$. Conditions: 10 mL solvent, 1 mM sample and 100 mM $(n\text{Bu}_4\text{N})\text{PF}_6$ as supporting electrolyte at 5–500 mV s⁻¹ scan rate (potentials vs. Ag/AgCl using the Fc^+/Fc couple as standard; $E_{1/2} = +0.4319$).

while the *para* and even more so the *meta* analogue are stronger donors.

The metal-centered nature of the oxidation process was confirmed by redox analyses of the Zn analogues **6a–c**. These complexes do not show any redox activity in the 0–2.2 V potential window (Fig. S14†), indicating that the coordinated PYA pincer ligands are redox inactive within this range. Likewise, the three pyridinium salt ligand precursors **1a–c** do not show any oxidation process up to 2 V and feature an irreversible reduction processes around -1 V (Fig. S13†). This reduction is dependent on the PYA substitution pattern and easiest for *ortho*-PYA system **1c** ($E_{\text{pc}} = -0.98$ V), intermediate for *meta*-PYA **1b** ($E_{\text{pc}} = -1.22$ V) and at lowest potential for the *para* analogue **1a** ($E_{\text{pc}} = -1.33$ V).

Electrochemical characterization of the nickel complexes supports the trend in donor strength of the PYA pincer ligands as observed for the Co series. Thus, the Ni^{3+} oxidation state was stabilized better by the *meta*-PYA pincer ligand ($E_{1/2} = 1.03$ V in **3b**) than the *para* analogue ($E_{1/2} = 1.20$ V in **3a**) or the *ortho*-pyPYA₂ system ($E_{1/2} = 1.34$ V in **3c**; Fig. 6a–c). In addition, the stronger donor sets also provided access to Ni^{IV} complexes *via* a second reversible oxidation at higher potential ($E_{1/2} = 1.72$ V and 1.54 V for complexes **3a** and **3b**, respectively). Notably, $[\text{Ni}(\text{terpy})_2](\text{PF}_6)_2$ as reference compound is oxidized at considerably higher potential ($E_{1/2} = 1.65$ V),⁵⁵ suggesting weaker donation of the terpy ligand than even the least basic *ortho*-pyPYA₂ ligand.

H/D exchange in *meta*-PYA sites

A closer investigation of the homoleptic cobalt complex **2b** with *meta*-PYA pincer ligands revealed intriguing ligand-based reactivity. Crystallization of this complex in D₂O under aerobic conditions resulted in a mixture of orange and green crystals. The latter are a product of spontaneous oxidation of the metal center and furthermore revealed selective deuterium incorporation into the PYA heterocycle, as demonstrated by MS and NMR analyses. Material crystallized from D₂O showed the *m/z* signals for the mono-, di-, and tri-cationic species at 8, 4 and 2.66 amu higher, respectively, than samples of **2b**⁺ crystallized from H₂O (Table S3†). Formation of **2b**⁺-D₈ was further supported by the absence of the PYA resonances at $\delta_{\text{H}} = 8.11$ and 8.25 in the ¹H NMR spectrum of the complex generated in D₂O (Fig. S15†). Based on NOE spectroscopy as well as the pertinent coupling constants, these resonances were assigned to H²_{PY} and H⁶_{PY}, respectively, *i.e.* the positions *ortho* to the PYA nitrogen. These data indicate a highly selective H/D exchange. Interestingly, the isomeric complexes **2a** and **2c** containing *para*- and *ortho*-PYA pincer ligands, respectively do not undergo any detectable deuteration or oxidation (MS, NMR) when exposed to D₂O, presumably because of the higher oxidation potential of these complexes (*vide supra*). The selective deuterium incorporation in the PYA 2- and 6-position of **2b** may be rationalized by considering resonance forms **B** and **C** of the *meta*-PYA ligand system, which imply a formally neutral imine donor site and opposite charges within the pyridylidene



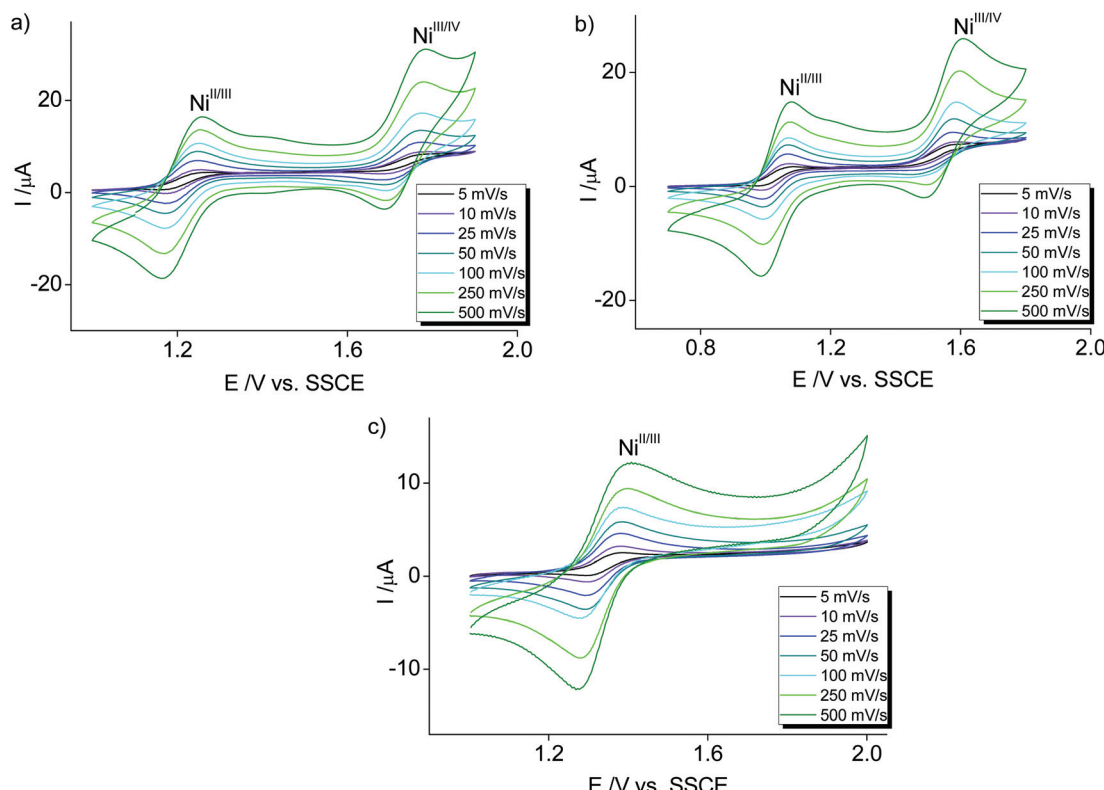
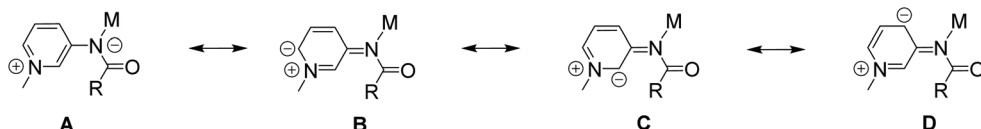


Fig. 6 Cyclic voltammetry measurements of homoleptic $[\text{Ni}(\text{pyPYA}_2)_2](\text{PF}_6)_2$ complexes in MeCN: (a) $3\mathbf{a}$ $[\text{Ni}(\text{para-pyPYA}_2)_2](\text{PF}_6)_2$; (b) $3\mathbf{b}$ $[\text{Ni}(\text{meta-pyPYA}_2)_2](\text{PF}_6)_2$; (c) $3\mathbf{c}$ $[\text{Ni}(\text{ortho-pyPYA}_2)_2](\text{PF}_6)_2$. Conditions: 10 mL solvent, 1 mM sample and 100 mM $(\text{nBu}_4)_2\text{PF}_6$ as supporting electrolyte at 5–500 mV s^{-1} scan rate (potentials vs. SSCE using the Fc^+/Fc couple as standard; $E_{1/2} = +0.4319$).



Scheme 5 Limiting resonance structures of the mesoionic *meta*-PYA ligand site. The negative charge is localized on the amide nitrogen and in 2-, 4- and 6-position of the PYA heterocycle.

heterocycle (Scheme 5). Reversible deuteration/protonation at the partially negatively charged positions would account for the observed selectivity and indicates that resonance forms **B** and **C** are contributing in addition to resonance form **A**. This contribution of several resonance structures therefore validates *meta*-PYA systems as mesoionic compounds.⁵⁶

Conclusions

Here, we have shown that PYA pincer ligands form stable complexes with nickel, cobalt and zinc, which expands the application potential of this ligand beyond platinum group metals to first row metals. The substitution pattern of the PYA heterocycle influences the binding mode of the pincer ligand to these Earth-abundant transition metals. *meta*- and *para*-pyPYA₂ ligand systems are exclusively κ^3 -*N,N,N*-coordinated, whereas the *ortho*-methylated PYA pincer ligand forms either κ^3 -*O,N,O*-coordinated

mono-pyPYA₂ compounds or bis-pyPYA₂ complexes with a mixed κ^3 -*O,N,O*/ κ^3 -*N,N,N* coordination mode. According to electrochemical analyses, the donor strength of the PYA ligand systems decrease in the order *meta*-PYA > *para*-PYA > *ortho*-PYA. The mesoionic *meta*-PYA pincer cobalt complex displays unique reactivity, since the 2- and 6-positions of the pyridinium heterocycle are selectively deuterated in D₂O under mild conditions. This deuteration suggests partial delocalization of the negative charge and therefore mesoionic properties of the *meta*-PYA unit. Moreover, this reactivity may become potentially useful for metal-ligand cooperative HX bond activation catalysis.

Experimental

General

All reagents were commercially available and used as received. Ligands **1a** and **1b** have been prepared according to previously



described methods.¹⁹ Unless specified otherwise, ¹H NMR spectra were recorded at 25 °C on Bruker spectrometers operating at 300 MHz. Chemical shifts (δ in ppm, coupling constants J in Hz) were referenced to residual solvent signals. Assignments are based on homo- and heteronuclear shift correlation spectroscopy. Purity of bulk samples of the complexes has been established by NMR spectroscopy, and when possible by elemental analysis. Elemental analyses were performed at DCB Microanalytic Laboratory using a Thermo Scientific Flash 2000 CHNS-O elemental analyzer. High-resolution ESI mass spectrometry was carried out with a Thermo Scientific LTQ Orbitrap XL (ESI-TOF). Cyclic voltammograms were recorded using an Autolab PGSTAT101 from Metrohm in MeCN solutions: 10 mL solvent, 1 mM sample and 100 mM tetrabutylammonium hexafluorophosphate (*n*Bu₄N)PF₆ as supporting electrolyte. Solutions were deaerated with argon gas for 10 min prior to each run. The scan rate was 100 mV s⁻¹. Redox potentials were measured using a Pt-button working electrode, an Ag/AgCl reference electrode and a Pt-wire auxiliary electrode and are tabulated *versus* a ferrocene internal standard. The Fe⁺/Fe couple is 0.43 V *vs.* Ag/AgCl in 0.1 M (*n*Bu₄N)PF₆ MeCN solutions.⁵⁷ ATR-FTIR spectra were recorded with a Jasco FT-IR-4700 spectrometer with a Jasco ATR Pro One unit containing a diamond crystal.

Synthetic procedures

Compound 1c. The bis-amide *N*²,*N*⁶-bis(pyridin-2-yl)pyridine-2,6-dicarboxamide has been prepared according to literature procedures.³³ Methylation was carried out under nitrogen atmosphere in 1,2-dichloroethane (50 mL) at 80 °C under vigorous stirring for 3 days with 5 eq. MeOTf (1.64 g; 10 mmol) in respect to the bis-amide (640 mg; 2 mmol). The reaction was quenched with MeOH (5 mL) and the product precipitated upon addition of Et₂O (50 mL). It was purified by filtration and repetitive washing with Et₂O (3 × 50 mL), yielding the bis-methylated OTf ligand salt **pre-1c** (1.03 g; 79% yield). The final product was obtained after anion exchange to PF₆, which was achieved upon addition of 4 eq. NH₄PF₆ (1.04 g; 6.3 mmol) to a solution of **pre-1c** (1.03 g; 1.6 mmol) in 30 mL of a warm (70°) MeCN/water mixture (1 : 2). The product crystallized upon slow cooling to room temperature, was filtered and washed with small portions of cold water (3 × 20 mL) and dried under vacuum. **1c** was obtained as white crystalline material (0.73 g; 72%). ¹H NMR (300 MHz, MeCN-*d*₃) δ 10.45 (s, 2H, NH), 8.64–8.51 (m, 6H, H_{PyA}, H_{pyr}), 8.46–8.36 (m, 3H, H_{PyA}, H_{pyr}), 7.87–7.79 (m, 2H, H_{PyA}), 4.29 (s, 6H, N-CH₃). ¹³C NMR (75 MHz, MeCN-*d*₃) δ 163.1 (CO), 148.1 (C_{Py}), 148.0 (C_{PyA}), 147.9 (CH_{PyA}), 146.5 (CH_{PyA}), 141.9 (CH_{pyr}), 128.8 (CH_{pyr}), 125.1 (CH_{PyA}), 124.7 (CH_{PyA}), 45.4 (N-CH₃). HR ESI-MS: *m/z* calculated for C₁₉H₁₉O₂N₅F₆P [M – PF₆]⁺ = 494.1151; found: 494.12. Elemental analysis: anal. calculated for C₁₉H₁₉F₁₂N₅O₂P₂: C: 35.70; H: 3.00; N: 10.95. Found: C: 36.28; H: 2.95; N: 11.03.

General procedure for compounds 2/3a–c

The respective amounts of the ligands **1a–c** (150 mg, 0.235 mmol, 1 eq.), Na₂CO₃ (75 mg, 0.705 mmol, 3 eq.) and

the MCl₂·6H₂O salts (28 mg, 0.117 mmol, 0.5 eq.) were dissolved in a H₂O/EtOH (1 : 1) mixture (10 mL) and stirred at r.t. for 30 min. The solvents were evaporated under reduced pressure. The crude product was redissolved in MeCN, cooled down to –18 °C and filtered over a pad of Celite to remove sodium salt residues. The final products were obtained upon precipitation/crystallization from Et₂O.

Compound 2a: [Co(*para*-PYA)₂](PF₆)₂. According to general procedure starting from **1a** and CoCl₂·6H₂O, which afforded **2a** as light orange solid (80 mg, 66%). ESI-MS: *m/z* calculated for [M – PF₆]⁺ = 898.1733; found: 898.1733. Elemental analysis: anal. calculated for C₃₈H₃₄CoF₁₂N₁₀O₄P₂: C: 43.73; H: 3.28; N: 13.42. Found: C: 43.50; H: 3.03; N: 13.85.

Compound 2b: [Co(*meta*-PYA)₂](PF₆)₂. According to general procedure starting from **1b** and CoCl₂·6H₂O, which afforded **2b** as orange solid (120 mg, 98%). ESI-MS: *m/z* calculated for [M – PF₆]⁺ = 898.1733; found: 898.1738. Elemental analysis: anal. calculated for C₃₈H₃₄CoF₁₂N₁₀O₄P₂·H₂O: C: 42.99; H: 3.42; N: 13.19. Found: C: 43.05; H: 4.20; N: 13.44.

Compound 2b⁺: [Co(*meta*-PYA)₂](PF₆)₃. A saturated solution of complex **2b** in H₂O or D₂O was left standing for 3 days which resulted in the formation of green crystals. The solvent was removed and the crystals washed with cold water and Et₂O and dried in air to give **2b⁺** (in H₂O) or **2b⁺–d₈** (in D₂O).

2b⁺: ESI-MS: *m/z* calculated for [M – PF₆]⁺ = 1043.1375; found: 1043.1393. ¹H NMR (300 MHz, MeCN-*d*₃): δ 8.39 (t, ³J_{HH} = 7.7 Hz, 2H, H_{PyA}), 8.27–8.22 (m, 4H, H⁶_{PyA}), 8.15 (d, ³J_{HH} = 7.7 Hz, 4H, H_{PyA}), 8.11 (s, 4H, H²_{PyA}), 7.62–7.56 (m, 8H, H^{4/5}_{PyA}), 4.09 (s, 12H, N-CH₃) ppm.

2b⁺–d₈: ESI-MS: *m/z* calculated for [M – PF₆]⁺ = 1051.1877; found: 1051.1883. ¹H NMR (300 MHz, MeCN-*d*₃): δ 8.39 (t, ³J_{HH} = 7.7 Hz, 2H, H_{PyA}), 8.15 (d, ³J_{HH} = 7.7 Hz, 4H, H_{PyA}), 7.61–7.56 (m, 8H, H^{4/5}_{PyA}), 4.10 (s, 12H, N-CH₃) ppm.

Compound 2c: [Co(*ortho*-PYA)₂](PF₆)₂. According to general procedure starting from **1c** and CoCl₂·6H₂O, which afforded **2c** as red solid (115 mg, 94%). ESI-MS: *m/z* calculated for [M – PF₆]⁺ = 898.1733; found: 898.1724. Elemental analysis: anal. calculated for C₃₈H₃₄CoF₁₂N₁₀O₄P₂: C: 43.73; H: 3.28; N: 13.42. Found: C: 42.22; H: 2.57; N: 13.51. Despite several attempts, no better microanalytical data were obtained. The irregular values may be associated with partial oxidation of the compound.

Compound 3a: [Ni(*para*-PYA)₂](PF₆)₂. According to general procedure starting from **1a** and NiCl₂·6H₂O, which afforded **3a** as bright yellow solid (78 mg, 64%). ESI-MS: *m/z* calculated for [M – PF₆]⁺ = 897.1754; found: 897.1730. Elemental analysis: anal. calculated for C₃₈H₃₄NiF₁₂N₁₀O₄P₂·0.5H₂O·0.5MeCN: C: 43.66; H: 3.43; N: 13.71. Found: C: 43.40; H: 3.03; N: 14.15.

Compound 3b: [Ni(*meta*-PYA)₂](PF₆)₂. According to general procedure starting from **1b** and NiCl₂·6H₂O, which afforded **3b** as dark yellow solid (115 mg, 95%). ESI-MS: *m/z* calculated for [M – PF₆]⁺ = 897.1754; found: 897.1758. Elemental analysis: anal. calculated for C₃₈H₃₄NiF₁₂N₁₀O₄P₂·1.5H₂O: C: 42.64; H: 3.48; N: 13.09. Found: C: 42.61; H: 4.09; N: 13.35.

Compound 3c: [Ni(*ortho*-PYA)₂](PF₆)₂. According to general procedure starting from **1c** and NiCl₂·6H₂O, which afforded **3c** as yellow/green solid (75 mg, 61%). ESI-MS: *m/z* calculated for

$[M - PF_6]^+$ = 897.1754; found: 897.1737. Elemental analysis: anal. calculated for $C_{38}H_{34}NiF_{12}N_{10}O_4P_2\cdot H_2O\cdot MeCN$: C: 43.58; H: 3.57; N: 13.98. Found: C: 43.14; H: 3.09; N: 14.04.

General procedure for compounds 4/5a–c

The respective amounts of the ligands **1a–c** (102 mg, 0.16 mmol, 1 eq.) and Na_2CO_3 (45 mg, 0.42 mmol, 3 eq.) were dissolved in DMF (4 mL) and stirred for 20 min at r.t. In a second vial the $MCl_2\cdot 6H_2O$ salts (38 mg, 0.16 mmol, 1 eq.) were dissolved in DMF (4 mL). The ligand solution was filtered and slowly added to the metal solution. The combined reaction phase was stirred for another 2 h at r.t. and the final products were obtained upon precipitation/crystallization from Et_2O and $MeCN/Et_2O$.

Compound 4a: $[Co(para-PY)_2Cl_2]$. According to general procedure starting from **1a** and $CoCl_2\cdot 6H_2O$, which afforded **4a** as light green solid (33 mg, 43%). ESI-MS: m/z calculated for $[M - Cl]^+$ = 441.0403; found: 441.0409. Elemental analysis: anal. calculated for $C_{19}H_{17}Cl_2CoN_5O_2$: C: 47.82; H: 3.59; N: 14.68. Found: C: 47.91; H: 3.31; N: 15.04.

Compound 4b: $[Co(meta-PY)_2Cl_2]$. According to general procedure starting from **1b** and $CoCl_2\cdot 6H_2O$, which afforded **4b** as turquoise solid (35 mg, 46%). ESI-MS: m/z calculated for $[M - Cl]^+$ = 441.0403; found: 441.0407. Elemental analysis: anal. calculated for $C_{19}H_{17}Cl_2CoN_5O_2\cdot 0.5MeCN$: C: 48.26; H: 3.74; N: 15.48. Found: C: 48.21; H: 3.36; N: 15.30.

Compound 4c: $[Co(ortho-PY)_2Cl_2]$. According to general procedure starting from **1c** and $CoCl_2\cdot 6H_2O$, which afforded **4c** as forest green solid (43 mg, 56%). ESI-MS: m/z calculated for $[M - Cl]^+$ = 441.0403; found: 441.0406. Elemental analysis: anal. calculated for $C_{19}H_{17}Cl_2CoN_5O_2$: C: 47.82; H: 3.59; N: 14.68. Found: C: 47.85; H: 3.55; N: 14.76.

Compound 5a: $[Ni(para-PY)_2Cl_2]$. According to general procedure starting from **1a** and $NiCl_2\cdot 6H_2O$, which afforded **5a** as yellow/orange solid (31 mg, 41%). ESI-MS: m/z calculated for $[M - Cl]^+$ = 440.0424; found: 440.0410. Elemental analysis: anal. calculated for $C_{19}H_{17}Cl_2NiN_5O_2\cdot 0.5H_2O$: C: 46.96; H: 3.73; N: 14.41. Found: C: 47.29; H: 4.13; N: 14.75.

Compound 5b: $[Ni(meta-PY)_2Cl_2]$. According to general procedure starting from **1b** and $NiCl_2\cdot 6H_2O$, which afforded **5b** as yellow solid (39 mg, 51%). ESI-MS: m/z calculated for $[M - Cl]^+$ = 440.0424; found: 440.0407. Elemental analysis: anal. calculated for $C_{19}H_{17}Cl_2NiN_5O_2$: C: 47.85; H: 3.59; N: 14.68. Found: C: 47.68; H: 3.70; N: 14.41.

Compound 5c: $[Ni(ortho-PY)_2Cl_2]$. According to general procedure starting from **1c** and $NiCl_2\cdot 6H_2O$, which afforded **5c** as green solid (42 mg, 55%). ESI-MS: m/z calculated for $[M - Cl]^+$ = 440.0424; found: 440.0410. Elemental analysis: anal. calculated for $C_{19}H_{17}Cl_2NiN_5O_2$: C: 47.85; H: 3.59; N: 14.68. Found: C: 47.72; H: 3.73; N: 14.31.

General procedure for compounds 6a–c

The respective amounts of ligands **1a–c** (100 mg, 0.16 mmol, 1 eq.) were reacted with Na_2CO_3 (85 mg, 0.80 mmol, 5 eq.) in $MeCN$ (5 mL) for 1 h at room temperature in a vial. Then a 0.2 M $ZnCl_2$ solution in $MeCN$ (470 μL , 0.09 mmol, 0.55 eq.) was

added to the ligand solution. The vial was sealed and heated up to 50 °C for 16 h under constant stirring. The reaction mixture was cooled to r.t. and filtered through Celite. A white solid was precipitated from Et_2O (20 mL). The solid was collected by filtration over a fritted funnel, rinsed with Et_2O (3 \times 5 mL) and dried.

Compound 6a: $[Zn(para-PY)_2](PF_6)_2$. According to general procedure starting from **1a** and giving complex **6a** as a white solid (47 mg, 54% yield). 1H NMR (300 MHz, $MeCN-d_3$): δ 8.47–8.24 (m, 3H, H_{PY}), 7.77 (d, $^3J_{HH}$ = 7.3 Hz, 4H, H_{PY}), 7.22 (d, $^3J_{HH}$ = 7.3 Hz, 4H, H_{PY}), 3.88 (s, 6H, $N-CH_3$) ppm. ^{13}C NMR (75 MHz, $MeCN-d_3$): δ 166.73 (CO), 165.14 (C_{PY}), 149.89 (C_{Py}), 143.73 (CH_{Py}), 143.35 (CH_{Py}), 125.90 (CH_{Py}), 120.89 (CH_{Py}), 46.13 (N-CH₃) ppm. ESI-MS: m/z calculated for $[M - PF_6]^+$ 903.1698; found: 903.1674. Elemental analysis: anal. calculated for $C_{38}H_{34}F_{12}N_{10}O_4P_2Zn\cdot 2H_2O$: C: 42.02; H: 3.53; N: 12.90. Found: C: 42.03; H: 3.18; N: 12.80.

Compound 6b: $[Zn(meta-PY)_2](PF_6)_2$. According to general procedure starting from **1b** and giving complex **6b** as a white solid (46 mg, 55% yield). 1H NMR (300 MHz, $MeCN-d_3$): δ 8.43 (s, 2H, H_{PY}), 8.29–8.18 (m, 3H, H_{PY}), 7.88 (d, $^3J_{HH}$ = 5.9 Hz, 2H, H_{PY}), 7.77 (d, $^3J_{HH}$ = 8.5 Hz, 2H, H_{PY}), 7.38 (dd, $^3J_{HH}$ = 8.5, 5.9 Hz, 2H, H_{PY}), 3.97 (s, 6H, $N-CH_3$) ppm. ^{13}C NMR (75 MHz, $MeCN-d_3$): δ 165.61 (CO), 151.40 (C_{PY}), 150.59 (C_{Py}), 143.43 (CH_{Py}), 140.48 (CH_{Py}), 139.66 (CH_{Py}), 137.10 (CH_{Py}), 127.66 (CH_{Py}), 124.93 (CH_{Py}), 48.95 (N-CH₃) ppm. ESI-MS: m/z calculated for $[M - PF_6]^+$ 903.1698; found: 903.1680. Elemental analysis: anal. calculated for $C_{38}H_{34}F_{12}N_{10}O_4P_2Zn\cdot 2H_2O$: C: 42.02; H: 3.53; N: 12.90. Found: C: 41.95; H: 3.23; N: 12.75.

Compound 6c: $[Zn(ortho-PY)_2](PF_6)_2$. According to general procedure starting from **1c** and giving complex **6c** as a white solid (72 mg, 80% yield). 1H NMR (300 MHz, $MeCN-d_3$): δ 8.70 (d, $^3J_{HH}$ = 7.8 Hz, 2H, H_{PY}), 8.41 (t, $^3J_{HH}$ = 7.8 Hz, 1H, H_{PY}), 8.11 (dd, $^{3/4}J_{HH}$ = 6.5, 1.7 Hz, 2H, H_{PY}), 8.06 (d, $^3J_{HH}$ = 8.7 Hz, 2H, H_{PY}), 7.85 (dt, $^{3/4}J_{HH}$ = 7.9, 1.7 Hz, 2H, H_{PY}), 7.08 (dt, $^3J_{HH}$ = 6.8, 1.4 Hz, 2H, H_{PY}), 4.02 (s, 6H, $N-CH_3$) ppm. ^{13}C NMR (75 MHz, $MeCN-d_3$): δ 166.38 (CO), 159.57 (C_{PY}), 150.23 (C_{Py}), 144.20 (CH_{Py}), 143.86 (CH_{Py}), 143.51 (CH_{Py}), 127.03 (CH_{Py}), 122.96 (CH_{Py}), 118.0 (CH_{Py}), 43.60 (CH₃) ppm. ESI-MS: m/z calculated for $[M - PF_6]^+$ 903.1698; found: 903.1673. Elemental analysis: anal. calculated for $C_{38}H_{34}F_{12}N_{10}O_4P_2Zn$: C: 43.47; H: 3.26; N: 13.34. Found: C: 42.99; H: 3.16; N: 13.11.

Crystal-structure determination

Suitable single crystals of **2b–c**, **3b–c**, and **4a–c** were mounted in air at ambient conditions and measured on an Oxford Diffraction SuperNova area-detector diffractometer⁵⁸ using mirror optics monochromated Mo K α radiation (λ = 0.71073 Å) and Al filtered.⁵⁹ The unit cell constants and an orientation matrix for data collection were obtained from a least-squares refinement of the setting angles of reflections. Data reduction was performed using the CrysAlisPro⁵⁸ program. The intensities were corrected for Lorentz and polarization effects, and a numerical absorption correction based on Gaussian inte-



gration over a multifaceted crystal model was applied. The structure was solved by direct methods using SHELXT,⁶⁰ which revealed the positions of all non-hydrogen atoms of the title compounds. The non-hydrogen atoms were refined anisotropically. All H-atoms were placed in geometrically calculated positions and refined using a riding model where each H-atom was assigned a fixed isotropic displacement parameter with a value equal to $1.2U_{\text{eq}}$ of its parent atom ($1.5U_{\text{eq}}$ for methyl groups and water). Refinement of the structure was carried out on F^2 using full-matrix least-squares procedures, which minimized the function $\sum w(F_{\text{o}}^2 - F_{\text{c}}^2)^2$. The weighting scheme was based on counting statistics and included a factor to down-weight the intense reflections. All calculations were performed using the SHELXL-2014/7⁶¹ program in OLEX2.⁶² The PF_6^- anions in the crystals of complex **2b**, **2c**, and **3b** are disordered as indicated by the high residual electron density peak near its P-atom. No conclusive disorder model was found for **2b** and **3b**, while for **2c**, disorders about two sites were restrained by SHELX SIMU and RIGU instructions. Further details are given in Tables S4–S6.† Crystallographic data for all structures have been deposited with the Cambridge Crystallographic Data Centre (CCDC) as supplementary publication numbers 2015875 (**2b**), 2015876 (**2b**†), 2015877 (**2c**), 2015878 (**3b**), 2015879 (**3c**), 2015880 (**4a**), 2015881 (**4b**), 2015882 (**4c**), and 2015883 ($[\text{Ru}(\text{ortho-pyPYA}_2)(\text{MeCN})_3](\text{PF}_6)_2$).†

Conflicts of interest

The authors declare no conflict of interest.

Acknowledgements

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