Metal Complexes with Cis α Topology from Stereoselective Quadridentate Ligands with Amine, Pyridine, and Quinoline Donor Groups

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Though the principles governing quadridentate topology and metal stereochemistry have been known for some time, the cis α topology has been little exploited in designing catalysts for asymmetric reactions. Investigation of the inorganic chemistry of labile metal cis α complexes was undertaken as a prelude to exploring their potential to serve as catalysts for a variety of different reactions. The synthesis of a series of first row transition metal complexes of quadridentate ligands with ethylenediamine (en) and S-propylenediamine (S-pn) backbones that have been alkylated at nitrogen with either pyridine (py) or quinoline (qn) donor groups as well as with noncoordinating benzyl (Bn) or pentafluorobenzyl (F₅Bn) groups was undertaken. The steric and electronic properties vary throughout the ligand series, en(Bn)py, **1**, $en(F_5Bn)py$, **2**, $S-pn(F_5Bn)py$, **3**, and $S-pn(F_5Bn)qn$, **4**. These ligands were reacted with MCl_n salts (n = 2, M = Mn, Fe, Co, Ni, Cu, Zn; n = 3, M = Fe) to generate, in most cases, octahedral complexes with the targeted cis a topology. UV/vis, NMR, IR, cyclic voltammetry (CV), and conductivity analysis are described for the metal compounds. X-ray structural analysis of [Cu{en(F₅-Bn)py{Cl]Cl reveals a five coordinate square pyramidal geometry. Single or major diastereomers were obtained for all diamagnetic Zn(II) complexes as well as for Co(III) analogues that were prepared by oxidation of Co(II) species using Br_2 as the oxidant. Electronic differences among ligands are reflected in the oxidation potentials of the respective metal complexes as determined by CV, with fluorinated systems showing greater resistance to oxidation, as expected.

Introduction

Recently there has been a resurgence of interest in chiral coordination compounds with nitrogen ligands¹⁻⁴ for their potential as asymmetric catalysts in both small molecule and polymer synthesis. For example, chiral metal salen analogues have been utilized as asymmetric oxygen¹⁻⁸ and nitrogen atom transfer catalysts^{5,9,10} and as catalysts for the ring opening of epoxides (Figure 1).^{11,12} Complexes of oxazoline ligands, such as pybox, are effective Lewis acid catalysts and promote cyclopropanation and aziridination reactions.^{13–21} Osmium–

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Figure 1. Examples of coordination compounds with chelating nitrogen donor ligands used in catalysis.

pybox

X

pyridine diimine

alkaloid systems have been developed into highly selective catalysts for dihydroxylation^{22–24} and aminohydroxylation.²⁵ Interestingly, simple coordination compounds are also playing key roles as catalysts in contemporary polymer chemistry. Atom

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10.1021/ic990475v CCC: \$18.00 © 1999 American Chemical Society Published on Web 11/09/1999 transfer radical polymerization (ATRP) of styrene and acrylate monomers²⁶ utilizes Cu(I) complexes with bipyridine,^{26–28} chelating amine,^{29–31} and Schiff base ligands,^{32–34} whereas metal complexes with diimine ligands figure prominently in recent advances in a polyolefin synthesis, with many showing high catalytic activity.^{35–37} Chiral variants of some of these systems have been employed in an attempt to control polymer tacticity.³⁸

Though systematic catalyst tuning can be difficult for asymmetric complexes possessing C_1 symmetry, numerous derivatives of the dissymmetric C₂-symmetric quadridentate salen¹⁻⁸ and bi- or tridentate oxazoline-based systems¹³⁻²¹ are known. Upon coordination, these ligands typically adopt planar topologies in which carbon stereocenters on the ligand framework generate a chiral array at the reactive center. Though much success has been achieved with these planar catalysts, comparatively little has been reported about the reactivity of nonplanar chiral chelates despite the fact that methods for controlling linear quadridentate chelate topology and metal absolute configuration have been known for some time. Many studies have been performed using nitrogen donor ligands,39,40 with mixed pyridylamine chelates figuring prominently.⁴¹ Structural features and physical characterization of many inert and labile metal pyridylamine systems have been investigated.⁴²⁻⁴⁹ This family of ligands has been exploited in bimetallic enzyme

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Figure 2. Quadridentate chelate topologies.

models $^{50-54}$ and as catalysts for redox processes $^{55-58}$ and olefin polymerization. 59

By the appropriate selection of metal chelate ring sizes, donor groups, and chiral centers on the ligand backbone, it is possible to generate ligands that result in major, if not single diastereomers upon coordination to metal ions. Since five-membered chelate rings typically possess bite angles of less than 90°, cumulative angle strain is minimized when chelate rings are positioned out of plane with respect to each other.⁶⁰ Others have pointed out that "B strain," namely unfavorable in-plane nonbonding interactions between α -hydrogens on pyridyl residues, also disfavor the planar orientation.⁴³ Thus, quadridentate chelates with three consecutive five-membered metal chelate rings preferentially adopt cis topologies, either cis α or cis β , over trans structures (Figure 2). For quadridentates in which the two internal donor groups are secondary amines, cis β complexes or mixtures of cis α and cis β isomers are typically obtained.^{41,43,50,61-64} However, when these internal donor groups are tertiary amines⁵⁰ or sulfides, $^{65-67}$ cis α structures have been observed. It is assumed that apical binding is energetically preferable to the cis β in-plane chelation, since the terminal

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Figure 3. Comparison of cis α and ansa metallocene complexes.







Figure 4. Topological and conformational chirality in quadridentate chelates.

donor group and the substituent on N (or lone pair on S) are oriented anti to each other in the cis α arrangement.⁴⁷ It is interesting to compare cis α complexes and ansa metallocenes, 70-73 which are similar in topology. They differ in that strapped metallocene ligands bind "face on" via cyclopentadienyl (Cp) and its analogues, with substituents projecting out and away from the reaction wedge for many derivatives. In contrast, the terminal donor groups on cis α pyridylamine ligands bind "edge on" and R groups point more directly at substrate binding sites (Figure 3).

While these principles may be exploited to impose a nonplanar C_2 -symmetric cis α topology, other features must be incorporated to make the ligand stereoselective and, in some cases, even stereospecific.41 In many chelating ligands, the introduction of one or more chiral centers on the internal chelate ring backbone has been sufficient to ensure that a single diastereomer is obtained. For example, when the sexadentate chelate S-propylenediamine tetraacetic acid (S-PDTA) binds to metal ions, the methyl substituent assumes a pseudoequatorial position on the puckered five-membered chelate ring. This preference fixes the conformational chirality of the metal propylenediamine ring as δ , which in turn, controls the way in which the terminal donor groups wrap to give exclusively the Δ absolute configuration about the metal center. 68,69 A similar kind of diastereoselection has been observed in amine-based quadridentate chelates prepared from enantiomerically pure chiral diamines (Figure 4).⁴¹

Taking these concepts and precedent with related systems into account, we have prepared a series of quadridentate ligands (Figure 5) and their first-row transition metal complexes. This study was undertaken as a prelude to screening these complexes



Figure 5. Quadridentate pyridylamine ligands.

as catalysts for a variety of reactions. The first goal was to identify convenient and versatile synthetic routes to ligand targets. Ethylenediamine (en) and the chiral S-propylenediamine (S-pn) were employed as the backbones for comparison.^{46,74} Ligands based on the bulkier R.R-diaminocyclohexane and their complexation chemistry are the subject of a future report.⁷⁵ For the S-pn system, ligands with pyridyl, 3, and quinolyl, 4, donor groups were prepared to vary the steric bulk in the terminal donor positions of the quadridentate. To determine whether subtle differences in the electronic nature of substituents might influence physical properties, both -CH2C6H5, 1,76 and $-CH_2C_6F_5$, 2, analogues of the achiral en backbone ligand were synthesized. Benzyl groups were also chosen for their steric bulk and their potential to improve solubility in organic solvents. Since many reactions are promoted by Lewis acidic metal centers, electron withdrawing fluorine substituents were introduced to generate more electron deficient complexes. Moreover, since pyridine and quinoline groups are π acidic, these donors are also expected to enhance Lewis acidity. Depending upon the particular metal ion, the steric features of the ligand, and the donor strength of the counterions, six, five, or four coordinate structures could be obtained upon reaction of quadridentates with divalent metal halide salts, MX₂. Ultimately, complexes prepared from metal halides may be further activated to coordinate Lewis basic substrates by exchanging the halides for triflates, OTf⁻, hexafluoroantimonates, SbF₆⁻, hexafluorophosphates, PF₆⁻, tetraphenylborates, BPh₄⁻, or other weakly coordinating counterions via metathesis with the respective silver salts. The synthesis of these ligands and many of their firstrow transition metal complexes, as well as structural and physical characterization, are discussed below.

Experimental Section

General Considerations. All reagents and solvents were used as received from commercial sources (Aldrich, Acros, Strem) unless otherwise indicated. Pyridinecarboxaldehyde was distilled under vacuum prior to use. THF was dried and purified on alumina columns.77 Cyclic voltammetric measurements were made either with a Bioanalytical Systems, Inc., model CV-27 or a Bioanalytical Systems model CV-

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50W instrument on dichloromethane solutions that contained 0.1 M tetra-n-butylammonium hexafluorophosphate (TBAH) as the supporting electrolyte. A glassy carbon electrode and a Pt-wire electrode were utilized. $E_{1/2}$ values, determined as $(E_{p,a} + E_{p,c})/2$, were referenced to the aqueous Ag/AgCl electrode at room temperature and are uncorrected for junction potentials. Under our experimental conditions the ferrocenium/ferrocene couple is at $E_{1/2} = 0.47$ V vs Ag/AgCl. Voltammograms shown in Figure 8 were referenced vs a nonaqueous (CH₃CN) Ag/AgCl electrode. Under these conditions the ferrocene couple is observed at $E_{1/2} = 0.24$ V. Conductivity measurements were performed at room temperature on 1 mM acetonitrile solutions using a YSI model 35 conductance meter. Infrared spectra were recorded as mineral oil (Nujol) mulls supported on NaCl plates in the region 4000-600 cm⁻¹ using a Nicolet Impact 400 Fourier transform spectrometer. Electronic absorption spectra were recorded on Hewlett-Packard diode array spectrophotometers, either model HP8452A (200-800 nm) or model HP8453 (200-1100 nm). ¹H and ¹³C NMR spectra were recorded at 300 and 75 MHz, respectively, on either a GE QE-300 or a GN-300 spectrometer. Elemental microanalyses were performed on a Perkin-Elmer model 2400 Series II CHNS/O analyzer or were determined by Atlantic Microlab, Inc., Norcross, GA.

Ligand Syntheses. 1,6-Diphenyl-2,5-bis(2-methylpyridyl)-2,5-diazahexane, en(Bn)py (1). Ethylenediamine (2.1 mL, 0.031 mol) was stirred with 2-pyridinecarboxaldehyde (6 mL, 0.063 mol) in CH₂Cl₂ (40 mL) containing molecular sieves for 2 h. The reaction was then filtered and concentrated to give the Schiff base as a pale yellow oil: 7.10 g, 95%.78,79 The diimine (7.10 g, 0.030 mol) was dissolved in MeOH (50 mL), NaBH₄ (2.48 g, 0.066 mol) was added, and the reaction was stirred for 30 min at room temperature. Aqueous HCl (3 M, 24 mL) was added slowly with stirring, and then the MeOH was removed via rotary evaporation. Aqueous NaOH (20%) was added until the solution reached pH > 10. The product was extracted with CH_2Cl_2 (4 \times 50 mL), was washed with H₂O (2 \times 50 mL) and brine (50 mL), and then was dried over Na₂SO₄. Following filtration and concentration, the diamine was obtained as a yellow oil: 6.0 g, 83%.^{42,80} Alkylation was effected by dissolving the diamine (0.742 g, 3.06 mmol) in dry THF (50 mL) and deprotonating with NaH (0.476 g, 19.8 mmol). The mixture was cooled to 0 °C and was stirred for ~20 min prior to addition of benzyl bromide (1.1 mL, 9.3 mmol). The reaction was allowed to warm to room temperature slowly and then was stirred at that temperature for an additional 12 h. Water (15 mL) and then HCl (3 M) were added until the solution reached pH = 2. The acidic mixture was concentrated in vacuo to remove THF and then was washed with Et₂O (3 \times 20 mL). The aqueous solution was made basic (pH > 10) through addition of NaOH (10%), was saturated with NaCl, and then was extracted with EtOAc (4 \times 20 mL). The solution was dried over Na₂SO₄, filtered, and concentrated to yield the product, 1, as an offwhite solid: 1.22 g, 80%. The ligand may be further purified by silica gel flash chromatography with EtOAc. ¹H NMR (CDCl₃, 300 MHz): δ 2.68 (s, 4H), 3.57 (s, 4H), 3.70 (s, 4H), 7.11 (m, 2H), 7.25 (m, 10H), 7.45 (d, J = 8.1 Hz, 2H), 7.57 (m, 2H), 8.48 (d, J = 5.0, 2H).

1,6-Bis-(pentafluorophenyl)-2,5-bis(2-methylpyridyl)-2,5-diazahexane, en(F₅Bn)py (2). The ligand 2 was prepared by the method described for 1 using α -bromo-2,3,4,5,6-pentafluorotoluene instead of benzyl bromide. In the final work up, after the aqueous layer was made basic, the alkylated product was extracted with CH₂Cl₂ instead of EtOAc. The ligand, 2, was obtained as a white crystalline solid: 2.44 g, 87%. ¹H NMR (CDCl₃, 300 MHz): δ 2.66 (s, 4H), 3.74 (s, 4H), 3.76 (s, 4H), 7.14 (m, 2H), 7.34 (d, J = 7.9 Hz, 2H), 7.60 (m, 2H), 8.48 (d, J = 4.3 Hz, 2H).

3S-Methyl-1,6-bis(pentafluorophenyl)-2,5-bis(2-methylpyridyl)-2,5-diazahexane, S-pn(F_5Bn)py (3). L-Alanine methyl ester hydrochloride was converted to the free base by passing a MeOH solution

(400 mL) of the HCl salt (21.33 g, 0.153 mol) through a column of strongly basic Amberlite 402 anion-exchange resin (225 g, 17% capacity): 10.84 g, Yield: 69% by NMR integrations. The resulting ester was reacted with NH3 in a saturated MeOH solution to form the amide, which was subsequently reduced with BH3'THF and worked up as described by Miller et al.⁸² to produce S-propylenediamine as its HCl salt. S-propylenediamine•2HCl (0.980 g, 6.66 mmol) was crushed into a powder and then was suspended in CH₃CN (50 mL). Triethylamine (18 mL, 0.13 mol) and 2-pyridinecarboxaldehyde (1.3 mL, 0.014 mol) were added, and the reaction was allowed to stir for ~ 15 h at 25 °C. After this time, H₂O (30 mL) was added and the aqueous layer was extracted with CH_2Cl_2 (5 × 50 mL). Combined CH_2Cl_2 layers were dried over Na₂SO₄, filtered, and concentrated in vacuo to give the crude Schiff base product as a yellow orange oil: 1.50 g, 89%. ¹H NMR (CDCl₃, 300 MHz): δ 1.39 (d, J = 6.1 Hz, 3H), 3.89 (m, 3H), 7.28 (m, 2H), 7.70 (m, 2H), 7.97 (m, 2H), 8.36 (s, 1H), 8.40 (s, 1H), 8.61 (m, 2H). The resulting diimine was reduced with NaBH₄ in MeOH by the general procedure described above for 1: 1.14 g, 75%. $^{83}\ ^1\mathrm{H}$ NMR (CDCl₃, 300 MHz): δ 1.11 (d, J = 6.1 Hz, 3H), 2.71–2.56 (br m, 2H), 2.84 (m, 1H), 4.01-3.83 (br m, 4H), 7.14 (m, 2H), 7.33 (d, J = 7.3 Hz, 2H), 7.62 (m, 2H), 8.49 (d, J = 4.9 Hz, 2H). Alkylation of the resulting diamine was effected by the method described for 1 using α -bromo-2,3,4,5,6-pentafluorotoluene in place of benzyl bromide. In the final workup, after the aqueous layer was made basic, the alkylated product was extracted with Et₂O instead of EtOAc. The crude ligand, 3, was obtained as a viscous brown oil which solidified upon standing: 2.37 g, 87%. Typically the crude ligand **3** was not purified prior to reaction with metal chloride salts. However further purification may be effected by dissolving the crude ligand in Et₂O and adding hexanes to precipitate a viscous brown impurity. After filtration through Celite and concentration, the resulting residue may be recrystallized from Et₂O/hexanes to yield 3 as a beige powder. ¹H NMR (CDCl₃, 300 MHz): δ 1.07 (d, J = 6.6 Hz, 3H), 2.43 (m, 1H), 2.73 (m, 1H), 2.98 (q, J = 6.6 Hz, 1H), 3.70 (m, 8H), 7.11 (m, 2H), 7.39 (d, J = 7.8 Hz, 2H), 7.59 (m, 2H), 8.42 (d, J = 4.2 Hz, 1H), 8.47 (d, J = 5.1 Hz, 1H).

3S-Methyl-1,6-bis(pentafluorophenyl)-2,5-bis(2-methylquinolyl)-2,5-diazahexane, S-pn(F5Bn)qn (4). Crushed S-pn·2HCl was suspended in CH₃CN (54 mL), and then Et₃N (18 mL) and 2-quinoline carboxyaldehyde (2.22 g, 14.0 mmol) were added. The reaction was stirred for 15 h at 25 °C. H₂O (20 mL) was added, and the solution was extracted with CH_2Cl_2 (4 × 40 mL). Combined organic layers were washed with H₂O (2 \times 40 mL) and with saturated brine (2 \times 40 mL) and then were dried over Na2SO4. After filtration and concentration via rotovap, the crude Schiff base product was obtained as a brownish solid. The crude solid was dissolved in a minimal amount of CH₂Cl₂, and hexanes were added until a dark brown residue precipitated. The remaining supernatant was filtered through Celite and then was concentrated in vacuo just to the point when crystallization commenced. After being chilled at 0 °C for 2 h, the pale yellow crystalline solid was collected by filtration, was washed with a minimal amount of hexanes, and then was dried in vacuo: 1.48 g, 62%. ¹H NMR (CDCl₃, 300 MHz): δ 1.45 (d, J = 5.5 Hz, 3H), 4.00 (m, 3H), 7.55 (m, 2H), 7.72 (m, 2H), 7.82 (d, J = 7.9 Hz, 2H), 8.08 (s, 1H), 8.11 (s, 1H), 8.16 (m, 4H), 8.57 (s, 1H), 8.61 (s, 1H). The diimine was reduced with NaBH₄ in MeOH by the general procedure described for 1 to give the crude diamine as a tan oil in essentially quantitative yield. ¹H NMR (CDCl₃, 300 MHz): δ 1.17 (d, J = 6.1 Hz, 3H), 3.02–2.68 (br m, 3H), 4.24-4.05 (br m, 4H), 7.52-7.47 (m, 4H), 7.66 (m, 2H), 7.77 (d, J = 8.6 Hz, 2H), 8.02 (d, J = 3.6 Hz, 2H), 8.09 (d, J = 3.6 Hz, 2H)2H). The diamine was subsequently alkylated with α -bromo-2,3,4,5,6pentafluorotoluene in THF in the presence of NaH by the standard procedure described for 1. In the final workup, after the aqueous layer was made basic, the alkylated product was extracted with Et₂O instead of EtOAc. The product, 4, was obtained as a brittle yellow solid: 2.19

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g, 77%. ¹H NMR (CDCl₃, 300 MHz): δ 1.12 (d, J = 6.9 Hz, 3H), 2.51 (m, 1H), 2.84 (m, 1H), 3.08 (m, 1H), 3.78 (m, 8H), 7.49 (m, 4H), 7.68 (m, 2H), 7.76 (m, 2H), 7.99 (m, 4H).

Preparation of Metal Complexes. Metal complexes were prepared by reaction of alcohol (EtOH or MeOH) solutions of the appropriate metal chloride salt with an alcohol solution of the ligand. A typical reaction scale is as follows: ligand (0.15 mmol); metal chloride salt (0.16 mmol); alcohol (3 mL total). For the less soluble ligands, **2** and **4**, a minimal amount of methylene chloride was sometimes added to facilitate dissolution. In cases where the complexes precipitated from the reaction solution, they were collected by filtration. Otherwise the solutions were concentrated in vacuo and the residues were purified by recrystallization. Specific purification procedures and deviations from this standard method are indicated below for the respective compounds.

Metal Complexes of en(Bn)py (1). [Mn{en(Bn)py}Cl₂]·0.5CH₂Cl₂. The reaction solution was concentrated in vacuo. The resulting residue was recrystallized from CH₂Cl₂/Et₂O to give the Mn(II) complex as a white microcrystalline solid. Yield: 74%. $\Lambda_M = 0.20 \ \Omega^{-1} \ mol^{-1} \ cm^2$. Anal. Calcd for C_{28.5}H₃₁N₄Cl₃Mn: C, 57.93; H, 5.29; N, 9.48. Found: C, 57.61; H, 5.68; N, 9.30.

$$\label{eq:charge} \begin{split} & [\text{Fe}\{\text{en}(Bn)py\}Cl_2]\text{-}0.5CH_2Cl_2. \ \text{The CH}_2Cl_2/\text{EtOH reaction mixture} \\ & \text{was concentrated in vacuo. The resultant yellow residue was recrystal$$
 $lized from CH_2Cl_2/hexanes to give canary yellow needles. Yield: 82%. \\ & \Lambda_M = 0.13 \ \Omega^{-1} \ \text{mol}^{-1} \ \text{cm}^2. \ \text{Anal. Calcd for } C_{28.5}H_{31}N_4Cl_3\text{Fe: C, 57.84;} \\ & \text{H, 5.28; N, 9.47. Found: C, 58.23; H, 5.44; N, 9.43.} \end{split}$

[Co{en(Bn)py}Cl₂]·0.5CH₂Cl₂. The Co(II) compound was recrystallized by vapor diffusion of Et₂O into a CH₂Cl₂/Et₂O solution of the complex to yield a purple microcrystalline solid. Yield: 85%. $\Lambda_M = 5.6 \ \Omega^{-1} \ \text{mol}^{-1} \ \text{cm}^2$. Anal. Calcd for C_{28.5}H₃₁N₄Cl₃Co: C, 57.54; H, 5.25; N, 9.42. Found: C, 57.14; H, 5.60; N, 9.32.

[**Zn**{en(**Bn**)**py**}**Cl**₂]. The Zn(II) compound was recrystallized by vapor diffusion of Et₂O into a CH₂Cl₂/Et₂O solution to yield a pale yellow microcrystalline solid. Yield: 66%. $\Lambda_{\rm M}$ = 1.7 Ω⁻¹ mol⁻¹ cm². ¹H NMR (CDCl₃, 300 MHz): δ 2.34, 2.57 (system AB, $J_{\rm AB}$ = 4.4 Hz, 4H), 3.18, 4.96 (system AB, $J_{\rm AB}$ = 14.4 Hz, 4H), 3.42, 4.64 (system AB, $J_{\rm AB}$ = 13.7, 4H), 7.12 (br s, 4H), 7.22 (d, J = 6.39, 2H), 7.35 (m, 6H), 7.41 (t, 2H), 7.81 (t, 2H), 9.73 (s, 2H). ¹³C NMR (CDCl₃, 75 MHz): δ 43.8, 54.4, 58.3, 123.3, 123.6, 127.9, 128.1, 131.0, 131.9, 138.5, 149.6, 154.1.

Metal Complexes of en(F₅Bn)py (2). [Mn{en(F₅Bn)py}Cl₂]-0.5CH₃CH₂OH. The Mn(II) complex was prepared from MnCl₂ and the tetra-HCl salt of the ligand, **2**, in EtOH solution. White needles precipitated from the EtOH solution. Yield: 97%. $\Lambda_M = 1.4 \ \Omega^{-1} \ mol^{-1}$ cm². Anal. Calcd for C₂₉H₃₃N₄O_{0.5}F₁₀Cl₂Mn: C, 46.36; H, 3.09; N, 7.46. Found: C, 46.27; H, 3.04; N, 7.26.

[Fe{en(F₅Bn)py}Cl₂]PF₆. The Fe(III) complex precipitated from EtOH solution upon addition of excess NH₄PF₆ (3 equiv). The resulting yellow residue was recrystallized from CH₃CN/EtOH to yield a canary yellow microcrystalline solid. Yield: 62%. $\Lambda_M = 97 \ \Omega^{-1} \ mol^{-1} \ cm^2$. Anal. Calcd for C₂₈H₂₀N₄PF₁₆Cl₂Fe: C, 38.50; H, 2.31; N, 6.41. Found: C, 38.42; H, 2.46; N, 6.16.

[Fe{en(F₅Bn)py}Cl₂]·CH₂Cl₂. The EtOH reaction mixture was concentrated in vacuo, and the resulting crude Fe(II) compound was recrystallized from CH₂Cl₂/hexanes to give powdery canary yellow needles. Yield: 92%. $\Lambda_{\rm M} = 0.8 \ \Omega^{-1} \ {\rm mol}^{-1} \ {\rm cm}^2$. Anal. Calcd for C₂₉H₂₂N₄F₁₀Cl₄Fe: C, 42.78; H, 2.72; N, 6.88. Found: C, 42.45; H, 2.92; N, 6.65.

[Co{en(F₃Bn)py}Cl₂]·0.5CH₃CH₂OH·0.5CH₂Cl₂. The Co(II) complex was obtained as lavender needles by concentration of the EtOH reaction mixture, followed by recrystallization of the resulting residue by evaporation of CH₂Cl₂ from a CH₂Cl₂/EtOH solution. Yield: 90%. $\Lambda_{\rm M} = 7.6 \ \Omega^{-1} \ {\rm mol}^{-1} \ {\rm cm}^2$. Anal. Calcd for C_{29.5}H₂₄N₄O_{0.5}F₁₀Cl₃Co: C, 44.10; H, 3.03; N, 7.02. Found: C, 44.33; H, 3.00; N, 7.03. (Note: The ¹H NMR spectrum obtained after oxidation of this complex to Co(III) confirms the presence of these associated solvents. This spectrum is provided as part of the Supporting Information.)

[Ni{en(F₅Bn)py}Cl₂]·0.5CH₃CH₂OH. The Ni(II) complex was prepared from NiCl₂ and the tetra-HCl salt of the ligand. It precipitated as mint green needles from the EtOH reaction medium. Yield: 79%. $\Lambda_M = 22 \ \Omega^{-1} \ mol^{-1} \ cm^2$. Anal. Calcd for C₂₉H₂₃N₄O_{0.5}F₁₀Cl₂Ni: C, 45.76; H, 2.96; N, 7.49. Found: C, 46.05; H, 2.79; N, 7.48.

[Cu{en(F₅Bn)py}Cl₂]·0.5CH₃OH. The Cu(II) compound was obtained as a mint green microcrystalline solid after slow evaporation of CH₂Cl₂ from a CH₂Cl₂/MeOH solution of the complex. Yield: 93%. $\Lambda_M = 54 \ \Omega^{-1} \ mol^{-1} \ cm^2$. Anal. Calcd for C_{28.5}H₂2N₄O_{0.5}F₁₀Cl₂Cu: C, 45.46; H, 2.95; N, 7.44. Found: C, 45.54; H, 2.99; N, 7.13.

[**Zn**{en(F₃Bn)py}Cl₂]·**CH**₃OH. This complex was obtained as white needles. Yield: 84%. $\Lambda_{\rm M} = 5.1 \ \Omega^{-1} \ {\rm mol}^{-1} \ {\rm cm}^2$. Anal. Calcd for C₂₉H₂₄N₄OF₁₀Cl₂Zn: C, 45.14; H, 3.14; N, 7.27. Found: C, 45.42; H, 3.01; N, 7.14. ¹H NMR (CDCl₃, 300 MHz): δ 2.23, 2.49 (system AB, $J_{\rm AB} = 11 \ {\rm Hz}$, 4H), 3.36 (d, $J = 15.4 \ {\rm Hz}$, 2H), 3.48 (m, 2H), 4.83 (d, $J = 14.6 \ {\rm Hz}$, 2H), 5.05 (d, $J = 14.3 \ {\rm Hz}$, 2H), 7.30 (d, $J = 7.7 \ {\rm Hz}$, 2H), 7.47 (t, 2H), 7.87 (t, 2H), 9.75 (d, $J = 4.6 \ {\rm Hz}$, 2H). ¹³C NMR (CDCl₃, 75 MHz): δ 42.4, 45.5, 57.3, 123.7, 124.0, 135.4, 139.2, 142.6, 143.6, 147.0, 149.5, 153.6.

Metal complexes of S-pn(F₅Bn)py (3). [Mn{S-pn(F₅Bn)py}Cl₂]. The Mn(II) complex precipitated from the EtOH reaction solution as a white microcrystalline solid. Yield: 85%. $\Lambda_M = 2.0 \ \Omega^{-1} \ mol^{-1} \ cm^2$. Anal. Calcd for C₂₉H₂₂N₄F₁₀Cl₂Mn: C, 46.92; H, 2.99; N, 7.55. Found: C, 46.69; H, 3.28; N, 7.35.

[Fe{S-pn(F₅Bn)py}Cl₂]·0.5CH₂Cl₂. The EtOH reaction mixture was concentrated in vacuo. The resulting residue was dissolved in CH₂Cl₂ and precipitated from hexanes to produce a canary yellow powdery solid. Yield: 78%. $\Lambda_M = 0.5 \ \Omega^{-1} \ \text{mol}^{-1} \ \text{cm}^2$. Anal. Calcd for C_{29.5}H₂₃N₄F₁₀Cl₃Fe: C, 45.10; H, 2.95; N, 7.13. Found: C, 45.17; H, 3.26; N, 7.14.

 $\label{eq:cost} \begin{array}{l} \textbf{[Co{S-pn(F_5Bn)py}Cl_2]. The Co(II) complex precipitated from the EtOH reaction solution as a lavender microcrystalline solid. Yield: 82%. \\ \Lambda_M = 7.0 \ \Omega^{-1} \ mol^{-1} \ cm^2. \ Anal. \ Calcd \ for \ C_{29}H_{22}N_4F_{10}Cl_2Co: \ C, 46.67; \\ H, \ 2.97; \ N, \ 7.51. \ Found: \ C, \ 46.28; \ H, \ 3.33; \ N, \ 7.16. \end{array}$

[Ni{S-pn(F_5Bn)py}Cl₂]. The Ni(II) complex precipitated from the EtOH reaction solution as a mint green microcrystalline solid. Yield: 34%. $\Lambda_M = 30 \ \Omega^{-1} \ mol^{-1} \ cm^2$. Anal. Calcd for $C_{29}H_{22}N_4F_{10}Cl_2Ni$: C, 46.69; H, 2.97; N, 7.51. Found: C, 46.53; H, 3.31; N, 7.33.

[Cu{*S*-pn(F₅Bn)py}Cl₂]·0.5CH₂Cl₂. The Cu(II) compound precipitated as a lime green powder from a CH₂Cl₂/Et₂O solution. Yield: 71%. $\Lambda_{\rm M} = 92 \ \Omega^{-1} \ {\rm mol}^{-1} \ {\rm cm}^2$. Anal. Calcd for C_{29.5}H₂₃N₄F₁₀Cl₃Cu: C, 44.66; H, 2.92; N, 7.06. Found: C, 44.54; H, 3.27; N, 7.30.

[**Zn**{*S*-**pn**(**F**₅**Bn**)**py**{**Cl**₂]. The Zn(II) complex precipitated from the EtOH reaction solution as a white microcrystalline solid. Yield: 67%. $\Lambda_{\rm M} = 36 \ \Omega^{-1} \ {\rm mol}^{-1} \ {\rm cm}^2$. Anal. Calcd for C₂₉H₂₂N₄F₁₀Cl₂Zn: C, 46.27; H, 2.95; N, 7.44. Found: 46.13; H, 3.24; N, 7.24. Major isomer: ¹H NMR (CDCl₃, 300 MHz) δ 0.66 (d, *J* = 7.3 Hz, 3 H), 2.12 (d, *J* = 11.6 Hz, 1H), 2.47 (m, 1H), 2.87 (m, 1H), 3.34 (d, *J* = 6.1 Hz, 1H), 3.39 (d, *J* = 5.5 Hz, 1H), 3.45 (d, *J* = 14.0 Hz, 1H), 4.05 (d, *J* = 14.0, 1H), 4.76 (d, *J* = 16.5 Hz, 1H), 4.98 (m, 2H), 5.21 (d, *J* = 14.0 Hz, 1H), 7.30 (d, *J* = 7.9 Hz, 1H), 7.37 (d, *J* = 7.3 Hz, 1H), 7.46 (m, 2H), 7.89 (m, 2H), 9.72 (m, 2H); ¹³C NMR (CDCl₃, 75 MHz) δ 11.7, 39.5, 43.0, 50.8, 51.3, 55.8, 57.2, 123.6, 123.8, 124.1, 139.1, 149.6, 149.7, 153.4, 154.4

Metal Complexes of *S*-pn(F₅Bn)qn (4). [Co{*S*-pn(F₅Bn)qn}Cl₂]· 0.5CH₂Cl₂. The Co(II) complex was obtained as a periwinkle microcrystalline solid after precipitation from CH₂Cl₂/Et₂O. Yield: 70%. Λ_M = 43 Ω^{-1} mol⁻¹ cm². Anal. Calcd for C_{37.5}H₂₇N₄F₁₀Cl₃Co: C, 50.67; H, 3.06; N, 6.30. Found: C, 50.78; H, 3.45; N, 6.23.

[**Zn**{*S*-**pn**(**F**₅**Bn**)**qn**}**Cl**₂]. The Zn(II) compound was obtained as a beige solid upon concentration of the EtOH/CH₂Cl₂ solution. Yield: 46%. $\Lambda_{\rm M} = 29 \ \Omega^{-1} \ {\rm mol}^{-1} \ {\rm cm}^2$. Major isomer: ¹H NMR (CDCl₃, 300 MHz) δ 1.09 (d, *J* = 5.5 Hz, 3H), 2.99 (d, *J* = 15.3 Hz, 1H), 3.65 (m, 1H), 3.87 (m, 1H), 4.00 (d, *J* = 19.5 Hz, 1H), 4.09 (d, *J* = 15.9 Hz, 1H), 4.26 (d, *J* = 9.16 Hz, 1H), 4.32 (d, *J* = 12.8 Hz, 1H), 4.54 (d, *J* = 15.3 Hz, 1H), 4.79 (d, *J* = 15.3 Hz, 1H), 5.22 (m, 2H), 6.96 (m, 1H), 7.31 (m, 2H), 7.73 (m, 4H), 7.90 (d, *J* = 7.9 Hz, 1H), 8.03 (m, 1H), 8.25 (d, *J* = 7.9 Hz, 1H), 8.34 (d, *J* = 7.9 Hz, 1H), 9.69 (d, *J* = 6.7 Hz, 1H); ¹³C NMR (CDCl₃, 75 MHz) δ 13.2, 42.4, 46.2, 49.8, 55.8, 56.2, 59.6, 104.6, 108.8, 121.2, 122.7, 126.2, 127.0, 127.3, 128.0, 128.3, 129.7, 130.3, 135.6, 138.9, 139.6, 139.9, 140.9, 143.6, 144.0, 145.3, 147.2, 157.6, 159.7.

Oxidation of Co(II) Complexes. Co(II) complexes were oxidized with H_2O_2 according to the procedure described by Fenton et al.⁴⁶ Oxidations with Br₂ were performed by the procedure described below for [Co{en(Bn)py}Cl₂], and then ¹H NMR spectra were recorded. If





necessary, the solutions were warmed to dissolve the complexes prior to addition of bromine. Chemical shifts attributable to the major isomers evident in the ¹H NMR spectra of the bromine reactions are tabulated below. Note: In all cases, minor products characterized by resonances at \sim 9.88–9.95 ppm are present. It is estimated that these comprise up to 15–20% of the reaction mixture as determined by integration.

[Co{en(Bn)py}Cl₂] Oxidation. [Co{en(Bn)py}Cl₂] (8.7 mg, 0.015 mmol) and CDCl₃ (0.6 mL) were combined in an NMR tube. Bromine (0.050 mL, 0.97 mmol) was added, and a ¹H NMR spectrum of the resultant red-brown solution was recorded. ¹H NMR (CDCl₃, 300 MHz): δ 2.85 (s, 4H), 3.35 (m, 4H), 4.48 (m, 2H), 5.13 (m, 2H), 7.34 (m, 4H), 7.55 (m, 6H), 7.62 (m, 2H), 7.72 (m, 2H), 8.09 (m, 2H), 9.64 (d, *J* = 5.8 Hz, 2H).

[Co{en(F₅Bn)py}Cl₂] Oxidation. ¹H NMR (CDCl₃, 300 MHz): δ 2.97 (s, 4H), 3.42 (d, J = 14.2 Hz, 2H), 3.93 (d, J = 15.0 Hz, 2H), 4.72 (d, J = 15.4 Hz, 2H), 5.21 (d, J = 15.4 Hz, 2H), 7.76 (m, 4H), 8.17 (m, 2H), 9.68 (d, J = 5.8 Hz, 2H). This spectrum is provided as part of the Supporting Information.

[Co{*S*-pn(F₅Bn)py}Cl₂] Oxidation. ¹H NMR (CDCl₃, 300 MHz): δ 1.27 (m, 3H), 2.34 (m, 1H), 2.62 (d, J = 14.2 Hz, 1H), 3.32–2.96 (br m, 2H), 3.49 (m, 1H), 4.02 (d, J = 16.6 Hz, 1H), 5.01–4.67 (br m, 3H), 5.12 (d, J = 13.9 Hz, 1H), 5.45 (d, J = 16.2 Hz, 1H), 7.94–7.73 (br m, 4H), 8.20 (m, 2H), 9.64 (m, 2H).

X-ray Structure Determination. A thin green plate of dimensions $0.32 \times 0.11 \times 0.48$ mm was used for all X-ray experiments. The data collection was carried out on a Rigaku AFC6S diffractometer at -120 °C using Mo K α radiation ($\lambda = 0.710$ 69 Å). Unit cell dimensions were determined by applying the setting angles of 25 high-angle reflections. Intensities of three standard reflections were monitored during the data collection showing no significant variance. The intensities were corrected for absorption by using ψ scans of several reflections. The transmission factors ranged from 0.76 to 1.00. The structure was solved by direct methods (SIR92).⁸⁴ Calculations were performed on a Silicon Graphics Indigo 2 Extreme computer by employing the teXsan 1.7 software.85 Full-matrix least-squares refinement with anisotropic thermal displacement parameters for the Cu, Cl, and F atoms yielded a final R of 0.057 ($R_w = 0.078$). An inspection of a difference Fourier map indicated the presence of three weak peaks corresponding to a partially populated ethanol molecule. The nonhydrogen atoms of this molecule were refined isotropically with the occupancy of 0.5. The final difference map was essentially featureless with the highest peak of 0.59 $e/Å^3$.

Results and Discussion

Ligand Synthesis. Ligands 1-4 were prepared from the diamines by condensation with aldehydes, followed by reduction of the resulting imines to secondary amines, and subsequent alkylation at the secondary nitrogen centers to give the desired quadridentate chelates (Scheme 1). Ligands are designated in

the following manner: diamine(alkyl group)terminal donor group. The diamine backbone is indicated first (en or S-pn), followed by the noncoordinating alkyl group given in parentheses (Bn or F_5Bn) and then the terminal donor group (py = pyridine; qn = quinoline). The Schiff base precursor to ligands 1 and 2 was prepared by stirring ethylenediamine with 2 equiv of pyridinecarboxaldehyde in CH₂Cl₂ containing molecular sieves. Filtration and concentration yielded the diimine products in high yield. While this approach works well for en backbone ligands, since S-pn is isolated as its di-HCl salt via an asymmetric synthesis starting from L-alanine methyl ester hydrochloride,⁸² analogous Schiff base syntheses using S-pn to make ligands 3 and 4 required the prior liberation of the free base. This can be achieved using either concentrated basic solutions or a basic ion exchange column. Since these methods for generating the S-pn free base are somewhat tedious and they result in significant losses of product, a direct route was devised that involved the reaction of S-pn·2HCl with the desired aldehyde in the presence of excess base, i.e., triethylamine, in acetonitrile solution. After workup, the desired S-pn Schiff bases were obtained in good yield.

The Schiff bases were then reduced with sodium borohydride in methanol to yield quadridentates with secondary diamines at the internal donor positions.^{86,87} This approach is more convenient than other reported methods using diborane⁸³ or reductions with Zn/HOAc42 or Pd/C.88 Reduction products were obtained relatively cleanly and in high yield. The final step of ligand synthesis involves alkylation of the secondary nitrogen centers to produce tertiary amines. Though reductive amination using formaldehyde and sodium cyanoborohydride works well for methylation,⁴⁷ this route is not effective for larger, bulkier aldehydes. Instead it was discovered that reaction of the 2° diamines with NaH in THF, followed by addition of benzyl bromide, C₆H₅CH₂Br, or α-bromo-2,3,4,5,6-pentafluorotoluene, C₆F₅CH₂Br, resulted in the efficient formation of the desired alkylated products.⁸⁹ Most quadridentate ligands were used as prepared for making complexes. In certain cases, they were purified by chromatography or by conversion to their tetra-HCl salts followed by recrystallization.

Synthesis and Characterization of Metal Complexes. Metal complexes were prepared by combining alcohol solutions of the free ligands with those of the appropriate metal chloride, MCl_n (n = 2, M = Mn, Fe, Co, Ni, Cu, Zn; n = 3, M = Fe). In a few cases (M = Mn and Ni), following literature precedent, the tetra-HCl salt of the en(F₅Bn)py ligand was utilized in place of the free base. Since the S-pn(F₅Bn)qn ligand, 4, showed very limited solubility in methanol or ethanol, it was dissolved in a warm CH₂Cl₂/EtOH solvent mixture prior to addition of ethanol solutions of the metal salts. Immediate color changes indicative of complexation were observed in all cases. For certain ligandmetal combinations, namely for S-pn(F₅Bn)py with M = Co, Mn, Ni, and Zn and $en(F_5Bn)py$ for M = Mn and Ni, complexes precipitated directly from the alcohol reaction medium. Soluble complexes were isolated by concentration in vacuo followed by crystallization using either vapor diffusion or layering techniques with the indicated two solvent systems. Crude

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products were typically redissolved in either CH₂Cl₂ or CH₃-CN and were precipitated using ethanol, diethyl ether, or, more rarely, hexanes. When it proved difficult to obtain crystalline materials, complexes were sometimes isolated as analytically pure powders by rapid precipitation from similar solvent combinations. Details pertaining to specific metal complexes are provided in the Experimental Section. In general it was our experience that complexes of Mn, Fe(II), Co, and Ni tended to be easier to isolate as crystalline or powdery solids than complexes of Zn, Cu, and Fe(III) throughout the ligand series. This was especially true for Cu and Fe(II) complexes of the more bulky quinoline bearing ligand, **4**, which appeared to be unstable. Certain iron complexes of related ligands are unstable as well.⁵⁹

It should be noted that it was not possible to prepare the entire series of metal complexes for each of the ligands. Often analytical data for isolated materials did not correlate well with predicted structures. Especially puzzling was the fact that though many complexes of en(Bn)py, 1, yielded beautifully crystalline materials, it was difficult to determine the compositions of these samples based upon CHN analytical data. This could be due to partial dissociation of halide ligands and the presence of fractional amounts of solvent(s) associated with complexes in the solid state, to solvento species, or to the presence of minor impurities that proved difficult to remove. Though we were unable to obtain complexes of satisfactory purity with certain ligand and metal combinations after screening a variety of different solvent systems and purification techniques, it may still be possible to achieve pure and stable quadridentate compounds if different metal salts (e.g. other halides, acetates) and solvents are employed. Only those samples for which elemental analyses correlate well with predicted products are included in the Experimental Section. Since diamagnetic Zn-(II) complexes provide valuable information about metal complex structure, ¹H NMR spectral data in CDCl₃ solution are tabulated for all Zn complexes, regardless of analytical purity. Likewise, ¹H NMR spectral data for Co(III) complexes generated in situ from Co(II) precursors are also provided. Metal complexes were routinely characterized by conductivity measurements, electronic absorption and IR spectroscopy, and cyclic voltammetry.

The FT-IR spectra of the ligands and complexes all contain major bands at approximately 1655, 1525, 1503, and 1124 cm^{-1} . Free ligands 1-3 also exhibit pyridyl ring vibrations at $\sim 1585-$ 1590 cm⁻¹. These shift to higher energies upon complexation $(\sim 1601 - 1611 \text{ cm}^{-1})$. By comparison, for the free quinoline ligand, 4, an absorption appears at 1596 cm^{-1} and this shifts very little upon complexation (e.g. Co complex: 1603 cm⁻¹), which may correlate with weaker binding to this bulkier ligand. Complexes that precipitate with associated alcohol solvent typically exhibit OH stretches in the \sim 3400-3050 cm⁻¹ range. The Fe(III) complex of 2 shows a peak at 843 cm⁻¹, characteristic of the PF_6^- counterion. These observations are consistent with what has been observed previously for related complexes.43 Additional details about the structure and properties of quadridentate metal complexes are provided below and are organized according to metal ion in reverse order from Zn to Mn.

Zinc Complexes. All Zn(II) complexes are white or pale yellow in color with UV/vis spectra exhibiting bands in the UV region attributable to ligand absorptions (Table 1). ¹H and ¹³C NMR spectra, combined with conductivity data, reveal considerable information about the structure of Zn(II) complexes of the quadridentate ligand series. In chloroform solution, either single or major diastereomers (>95%) were obtained for Zn(II)

 Table 1. UV/Vis Spectral Data (250-800 nm Range) for Ligands and Complexes in CH₃CN Solution

compd	wavelength (nm) ^a
en(Bn)py	262 (6300), 270 sh, 300 sh
en(F ₅ Bn)py	262 (8500), 270 sh, 300 sh
S-pn(F ₅ Bn)py	262 (7800), 270 sh, 300 sh
S-pn(F ₅ Bn)qn	263 (9000), 303 (6000), 308 sh,
	316 (7300)
$Mn\{en(Bn)py\}Cl_2$	262 (8400), 269 sh
$Mn\{en(F_5Bn)py\}Cl_2$	262 (3400), 270 sh, 320 sh
$Mn{S-pn(F_5Bn)py}Cl_2$	264 (7200), 270 sh
Fe{en(Bn)py}Cl ₂	258 (10 000), 413 (1100)
$Fe\{en(F_5Bn)py\}Cl_2$	260 (8400), 338 (1900), 402 (1200)
$Fe{S-pn(F_5Bn)py}Cl_2$	260 (7900), 341 (1600), 398 (1200)
$[Fe{en(F_5Bn)py}Cl_2]PF_6$	256 (19 000), 265 sh, 300 sh, 376 (420)
$Co{en(Bn)py}Cl_2$	258 (7100), 270 sh, 300 sh, 510 sh,
	536 (24), 660 sh
$Co{en(F_5Bn)py}Cl_2$	260 (6400), 270 sh, 300 sh, 510 sh,
	540 (33), 640 sh
$Co{S-pn(F_5Bn)py}Cl_2$	260 (8300), 270 sh, 300 sh, 536 (38),
	660 sh
$Co{S-pn(F_5Bn)qn}Cl_2$	292 sh, 297 sh, 305 (7400), 317 (7600),
	588 (210), 633 sh, 666 sh,
	685 (240)
$Ni\{en(F_5Bn)py\}Cl_2$	258 (9400), 270 sh, 398 sh, 638 (16)
$Ni{S-pn(F_5Bn)py}Cl_2$	258 (16 000), 270 sh, 400 sh, 625 (23)
$Cu\{en(F_5Bn)py\}Cl_2$	262 (11 000), 290 (4500), 794 (97)
$Cu{S-pn(F_5Bn)py}Cl_2$	261 (20 000), 289 sh, 460 (75),
	794 (260)
$Zn\{en(F_5Bn)py\}Cl_2$	264 (6000), 270 sh
$Zn{S-pn(F_5Bn)py}Cl_2$	264 (6300), 270 sh

^{*a*} ϵ_{max} values in parentheses (M⁻¹ cm⁻¹); sh = shoulder.

complexes of all the ligands. Structural differences among Zn-(II) complexes correlate with the degree of steric bulk surrounding the metal center. Our work with ligands 1-4 and others of this type⁷⁵ has revealed that all three features of the ligand—the diamine backbone, the pendant alkyl substituent, and the apical donor group—play a role.

Zinc complexes of ligands with ethylenediamine backbones and pyridyl donor groups, en(Bn)py, 1, and en(F₅Bn)py, 2, adopt C_2 -symmetric structures in CDCl₃ solution as evidenced by their NMR spectra. The ¹H and ¹³C spectra for [Zn{en(F₅Bn)py}-Cl₂] in CDCl₃ are shown in Figure 6. This, combined with the fact that these complexes are nonconducting even in acetonitrile solution, is consistent with octahedral, $\operatorname{cis} \alpha$ structures in which both chlorides are coordinated. The Zn(II) complex of the S-pn backbone ligand, S-pn(F₅Bn)py, **3**, is also present as a major diastereomer in CDCl₃ solution. However, other species are also evident in trace amounts (\sim 5%). Since 3 bears a methyl group on the internal chelate ring backbone, the major isomer adopts a cis α topology but the complex is not C_2 symmetric. Most of the resonances in the aromatic region of the ¹H and ¹³C spectra are relatively insensitive to the methyl group on the diamine backbone. Differences are greater in benzylic and aliphatic regions; here all proton and carbon signals for [Zn{S-pn(F₅-Bn)py{Cl₂] are unique. Aside from expected differences arising from the unsymmetrical S-pn backbone, Zn complexes of 1-3give rise to very similar spectra. This suggests that they all possess the same cis α topology for the sole or major isomers.

In addition to varying the diamine backbone, it was also of interest to us to see how differences in steric demand at the terminal donor positions of the quadridentate ligand influence metal geometry. Thus, a Zn(II) complex of *S*-pn(F₅Bn)qn, **4**, was prepared for comparison. The ¹H NMR of the Zn complex of *S*-pn(F₅Bn)qn in chloroform solution suggests that one major isomer (>95%) is formed. However, the spectrum is complex, and all protons exhibit unique resonances. It is also interesting



Figure 6. ¹H (CD₃CN) and ¹³C (CDCl₃) NMR spectra of $Zn\{en(F_5-Bn)py\}Cl_2$ ·MeOH.

to note that the ¹H NMR spectrum of this qn complex shares considerable overlap in the benzylic and aliphatic regions with a minor isomer evident in the spectrum of $[Zn{S-pn(F_5Bn)py}]$ -Cl₂]. Both complexes exhibit resonances at \sim 9.7 ppm which, in our experience, seems to correlate with six coordinate structures. These observations, and comparison with data for cis β Co(III) complexes and ¹³C spectra of analogous bimetallic Zn complexes with cis α and cis β topologies, $^{41,43,50,61-64}$ strongly point to a cis β assignment for the major isomer of $[Zn{S-pn(F_5Bn)qn}Cl_2]^{.50}$ It should be noted, however, that both a nonideal cis- α structure, as was suggested by Rieger et al.⁵⁹ for a ferrous complex of a related but less bulky en(Me)qn ligand, and a five coordinate structure with one donor group dissociated, analogous to that seen for [Cu{en(F₅Bn)py}Cl₂] in the solid state (Figure 7), are also consistent with the observed conductivity and NMR data.

Unlike the Zn(II) complexes of the en backbone ligands, those of S-pn ligands 3 and 4 give rise to more complex ¹H NMR spectra upon dissolution in CD₃CN. New species emerge that are characterized both by the appearance of resonances in the 8-9 ppm and 3.5-4.6 ppm regions and the diminution of the downfield resonance at ~9.7 ppm. Moreover, these complexes conduct to a small extent ($\Lambda_M \sim 30 \ \Omega^{-1} \ mol^{-1} \ cm^2$) in CH₃CN solution. These observations suggest that a chloride ligand dissociates and either a five coordinate structure or a chlorosolvento species comprises a fraction of these samples. In summary, changes as subtle as the addition of a methyl substituent to the ethylenediamine backbone lead to observable differences in product distributions for Zn complexes of this family of quadridentate ligands. Whereas en-based ligands form single isomers, the S-pn systems show greater structural diversity in both polar and nonpolar solvents. This, combined with additional steric bulk at the terminal donor positions of these



Figure 7. ORTEP drawing and labeling scheme for $[Cu\{en(F_5Bn)-py\}Cl_2]$. Ellipsoids are at 30% probability. Hydrogens are omitted for clarity.

ligands (qn vs py), may force a cis β topology in Zn complexes of the *S*-pn(F₅Bn)qn ligand.

Copper Complexes. Copper complexes of the quadridentate ligands were typically isolated as green solids. Structural variation throughout the series of ligands for Cu(II) complexes is indicated by differences in color and conductivity in solution and by geometry in the solid state. In acetonitrile solution, complexes that are nonconducting or only slightly conducting (i.e. chlorides are coordinated in the inner sphere) tend to be green in color. Those in which chlorides are dissociated in a portion of the material are often blue-green or teal in color, whereas complexes from which the chloride ligands are removed using silver salts (e.g. $Ag(SbF_6)$) generally form blue solutions in both polar (CH₃CN, MeOH) and noncoordinating (CH₂Cl₂) solvents.42,75,90-93 Likewise, complexes become more blue in color as the electronic and steric features of the ligand are varied. For example, acetonitrile solutions of Cu(II) complexes of en-(Bn)py are green and nonconducting and those of the more electron deficient en(F₅Bn)py and S-pn(F₅Bn)py ligands are teal and slightly conducting, whereas a related ligand, R,R-cn(F₅-Bn)py with the bulkier *1R*,2*R*-diaminocyclohexane backbone,⁷⁵ is a 1:1 electrolyte and blue in color. These observations correlate well with predicted trends in the energy of d-d bands as the geometry at the Cu(II) center changes for a given ligand set75 and may be compared to Cu complexes of related ligands.43,94,95 Transitions are typically lower in energy (green) for distorted octahedral structures whereas square planar geometries typically give rise to higher energy transitions (blue). As is common for Cu(II) complexes, multiple Cu(II) species are likely present in acetonitrile solution as evidenced by complex UV/vis spectra and conductivities between values that would be expected for 1:1 electrolytes and nonconducting samples. This flexibility within the coordination sphere of Cu complexes has previously been described as the "plasticity effect".96 Moreover, as was seen in the solid state for [Cu{en- $(F_5Bn)py$ Cl₂, it is also possible that both chlorides are bound

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Table 2. Crystallographic Data for Cu{en(F₅Bn)py}Cl₂

chem formula	$C_{29}H_{23}N_4O_{0.5}F_{10}Cl_2Cu$
IW	/39.96
space group	$P2_1/a$ (No. 14)
a, Å	15.509(6)
b, Å	10.936(4)
c, Å	17.491(5)
β , deg	96.53(3)
V, Å ³	2947(1)
Ζ	4
T, °C	-120
λ, Å	0.710 69
$\rho_{\rm calcd}, {\rm g} {\rm cm}^{-3}$	1.71
μ (Mo K α), cm ⁻¹	10.17
$R(F_{o})^{a}$	0.057
R_{w}^{a}	0.078

a R =	$(\Sigma F_{\rm o} -$	$ F_{\rm c})/\Sigma F_{\rm o} ;$	$R_{\rm w} = \left[(\sum w(F_{\rm o} $	$- F_{\rm c})^2/\Sigma w(F_{\rm o})^2]$
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Table 3. Selected Bond Lengths (Å) and Bond Angles (deg) for $Cu\{En(F_5Bn)py\}Cl_2$

(a) Bond Lengths				
Cu-Cl(1)	2.519(3)	Cu-Cl(2)	2.369(3)	
Cu-N(1)	2.018(8)	Cu-N(2)	2.267(8)	
Cu-N(3)	2.425(7)	Cu-N(4)	2.034(8)	
(b) Bond Angles				
Cl(1)-Cu-Cl(2)	104.5(1)	Cl(1)-Cu-N(1)	89.4(2)	
Cl(1)-Cu-N(2)	89.2(2)	Cl(1)-Cu-N(4)	95.5(2)	
Cl(2)-Cu-N(1)	95.6(2)	Cl(2)-Cu-N(2)	164.8(2)	
N(1) - Cu - N(4)	171.6(3)	N(2) - Cu - N(4)	95.3(3)	

to give nonconducting solutions but that a nitrogen donor group may be dissociated to produce five coordinate structures overall.

The ORTEP diagram of the green Cu(II) complex, [Cu{en-(F₅Bn)py{Cl₂], is shown in Figure 7. Crystallographic parameters are provided in Table 2, and selected bond lengths and bond angles are given in Table 3. The especially long Cu-N(3) distance of 2.425(7) Å suggests that a five coordinate square pyramidal structure with an N₃Cl₂ donor set (an N(1)-N(2)-N(4)-Cl(2) base and a Cl(1) apical group) best describes the geometry of this compound in the solid state. Goodwin and Lions also proposed dissociation of a chelate donor group from Cu(II) centers to explain their observations with a related quadridentate ligand.⁴² This structure may be contrasted with [Cu{en(H)py}Cl]⁺, a five-coordinate Cu(II) complex with an N₄Cl donor set,⁹⁷ and also with a six coordinate Cu(II) complex of tn(H)py (tn = trimethylenediamine), a 5–6–5 chelate ring system with planar chelate coordination (i.e. N₄Cl₂ donor set).98,99 These solid-state examples illustrate that minor variations in the ligand can result in considerable structural differences in Cu complex geometry.96 Though crystallographic analysis of metal complexes of chiral backbone ligands could provide valuable insights into the topology and stereospecificity of these systems, it proved difficult for us to grow crystals suitable for analysis using the S-pn backbone ligands, 3 and 4.

Nickel Complexes. Nickel complexes were isolated as light green paramagnetic solids. All show very similar UV/vis spectra in CH₃CN and CH₂Cl₂ solutions with bands at ~630 and ~1080 nm typical for octahedral systems (Tables 1 and 4).¹⁰⁰ It should be noted that an additional band is present at ~400 nm as a

Table 4. Comparison of UV/Vis/NIR Spectral Data (250-1100 nm Range) for Nickel, Cobalt, and Iron Complexes of en(F₅Bn)py and *S*-pn(F₅Bn)py Ligands in Methylene Chloride Solution

compd	wavelength (nm)		
Fe{en(F5Bn)py}Cl2	336 (1740), 407 (1420) ^a		
$Fe{S-pn(F_5Bn)py}Cl_2$	335 (604), 406 (527) ^{a}		
$Co{en(F_5Bn)py}Cl_2$	518 sh, 542 (37), 635 sh, 1086 (5)		
$Co{S-pn(F_5Bn)py}Cl_2$	521 sh, 543 (37), 633 sh, 1084 (5)		
$Co{S-pn(F_5Bn)qn}Cl_2$	300 sh, 307 (8400), 320 (8200),		
	540 sh, 601 (95), 613 (97),		
	635 (110), 669 (100), 809 sh		
$Ni\{en(F_5Bn)py\}Cl_2$	402 (38), 642 (20), 1078 (11)		
Ni{S-pn(F5Bn)py}Cl ₂	401 (41), 646 (21), 1083 (8)		

^a Long tail across visible region.

shoulder on a much stronger UV band in the spectra of all Ni complexes. Energies of absorptions for the Ni(II) complex of weaker field fluorinated ligands 2 and 3 appear at slightly lower energies than those for [Ni{en(Bn)py}Cl₂]. Satisfactory analytical data were not obtained for the green crystalline [Ni{en-(Bn)py{Cl₂] complex; however, spectra were recorded for comparison. The following d-d transitions were observed: CH₃-CN, 394 sh, 629 nm; CH₂Cl₂, 393 sh, 631, 1062 nm. Though the complex [Ni{en(Bn)py}Cl₂] is nonconducting in acetonitrile solution ($\Lambda_{\rm M} = 1.3 \ \Omega^{-1} \ {\rm mol}^{-1} \ {\rm cm}^2$), Ni complexes of fluorinated ligands, 2 and 3, are slightly conducting in acetonitrile solutions ($\Lambda_{\rm M} = 22$ and 33 Ω^{-1} mol⁻¹ cm², respectively). This suggests that chloride ligands dissociate from the Ni center in a portion of the sample in this polar solvent. A five coordinate Ni(II) species or six coordinate solvento-chloro or aquo-chloro species are all consistent with the observed conductivity data. No dramatic changes are observed in acetonitrile spectra as compared with those in methylene chloride that might be attributed to these new species that are formed. UV/vis spectra of Ni(II) complexes of related but stronger field secondary diamine ligands such as en(H)py with a variety of different ancillary ligands and counterions have been previously recorded for solids and in a variety of different solvents.⁴³ For these complexes, many of which are likely to be $cis \alpha$ or even dimeric structures, d-d bands shift toward higher energies as expected (580 and 1000 nm for en(H)py vs 630 and 1080 nm for complexes of 2 and 3).

Cobalt Complexes. Cobalt complexes of ligands 1-3 were isolated as lavender microcrystalline solids. Electronic absorption spectra in acetonitrile solution reveal absorptions at approximately 510, 538, and 650 nm indicative of six coordinate high-spin d⁷ Co(II) centers for ligands 1-3 (Table 1).¹⁰¹ Additional bands are present in the near-IR region at ~ 1085 nm (CH₂Cl₂) which are consistent with this assignment (Table 4). Conductivity data also support this assertion, as all complexes are essentially nonconducting ($\Lambda_M < 10 \ \Omega^{-1} \ mol^{-1} \ cm^2$ in CH₃-CN). By comparison, the Co(II) complex of the more bulky S-pn(F₅Bn)qn ligand, **4**, is slightly more bluish in color in the solid state. This is consistent with what has been seen previously for Co(II) complex with a related sterically bulky quinoline ligand.⁵⁹ Higher intensity bands at 588 and 685 nm ($\epsilon = 210$ and 240 L mol⁻¹ cm⁻¹, respectively) in the UV/vis spectrum (CH₃CN) of this quinoline complex along with a molar conductivity, $\Lambda_M = 43 \ \Omega^{-1} \ mol^{-1} \ cm^2$, suggest that chloride ligands are dissociated for a fraction of the sample. Most likely, mixtures of six and five coordinate species are present in

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solution.¹⁰² Weak features present in the UV/vis spectrum of this complex in acetonitrile solution are more pronounced in the spectrum run in methylene chloride solution (Tables 1 and 4).

Though Co(II) complexes are paramagnetic, their oxidation to diamagnetic Co(III) systems allows for structural characterization by ¹H NMR and for comparison of these quadridentate ligand systems with analogous Co(III) systems.^{44–48} A variety of different methods have been reported for the generation of Co(III) complexes with these types of quadridentate chelates, reaction with hydrogen peroxide in ethanol solution being the most common of these.⁴⁶ Unfortunately, our attempts to employ this standard preparation with ligands 1-4 met with little success. Reaction of CoCl₂, en(Bn)py, and hydrogen peroxide vielded a blue-green solid, as expected by analogy with related systems; however, it did not precipitate readily from EtOH solution and was isolated by concentration instead. Inspection of products by ¹H NMR revealed chemical shifts consistent with what has been previously reported for analogous C_2 symmetric cis α systems⁴⁶ though resonances were broad. Hydrogen peroxide reactions with Co(II) chloride and the S-pn(F₅Bn)qn ligand, 4, also exhibited the expected color changes; a green solid was obtained after partial concentration and precipitation with additional EtOH. However, as with the en(Bn)py ligand, 1, only broad resonances were observed, indicative of the presence of paramagnetic impurities. The Co(II) complex of $en(F_5Bn)py$, 2, turned brown upon reaction with H_2O_2 , and reaction of Co(II) with S-pn(F₅Bn)py and H₂O₂ resulted in the precipitation of the unreacted lavender Co(II) complex.

Lack of success with the standard methods prompted us to explore other common oxidants.¹⁰³ Reactions with Ag(OTf) did not generate significant quantities of the desired Co(III) products; however, reaction of Co(II) complexes with the stronger oxidant, bromine, did yield diamagnetic Co(III) products in all cases except for with the bulky ligand S-pn(F₅-Bn)qn, 4. The ¹H NMR spectrum of the oxidation product of $[Co{en(F_5Bn)py}Cl_2]$ is provided as Supporting Information. It clearly reveals that a C_2 -symmetric cis α isomer was obtained. Cobalt complexes of other ligands also formed major isomers under these reaction conditions; however, all spectra bear evidence of at least one other minor product as well (<20% of the total sample as determined by integration). It should be noted that Br₂ may not be an innocent oxidant in these reactions. The bromide ion formed could exchange with the chloro compounds to generate new mixed-halide complexes. Moreover, it has been reported previously that dichlorocobalt(III) complexes with analogous quadridentate ligands are highly susceptible to hydrolysis; they rapidly form chloro-aquo species that are 2:1 electrolytes.46,104 These various factors complicate product analysis and make it difficult to determine whether minor isomers are ancillary ligand byproducts, topological isomeric impurities, or some combination of the two.

Iron Complexes. Iron(II) complexes of $en(F_5Bn)py$, *S*-pn- $(F_5Bn)py$, and en(Bn)py were isolated as canary yellow solids. Iron complexes of the bulky quinoline ligand, **4**, were not amenable to synthesis by our standard preparations. Solutions turned orange and then dark brown upon reaction of the ligand with ferrous chloride, suggesting ready oxidation to Fe(III) products. Others have reported that the iron complex of a related nonfluorinated ligand, namely en(Bn)qn, was also susceptible to degradation in the presence of water and other solvents.⁵⁹ Though some investigators have used this sensitivity to oxidation to synthetic advantage, no such Fe complexes were isolated with the quinoline ligand, 4. In contrast, the Fe(II) complexes of 1-3showed no evidence of sensitivity to oxidation in solution or in the solid state. This is consistent with the observations of Hazell et al., who have noted that quadridentates with tertiary amine donors help to stabilize the lower Fe(II) oxidation state whereas ligands with stronger field secondary amines in the internal donor positions are sometimes prone to oxidative dehydrogenation, thus forming purple products.¹⁰⁵ For en(F₅Bn)py, an Fe-(III) complex was also readily prepared by reaction of the ligand with FeCl₃ followed by addition of NH₄PF₆ to precipitate the product, [Fe{en(F₅Bn)py}Cl₂]PF₆, as canary yellow plates. This complex is a 1:1 electrolyte in acetonitrile solution, consistent with an octahedral geometry at the metal center. The electronic absorption spectrum of this complex is similar to that observed for a related Fe(III) complex, [Fe(en(Me)py)Cl₂]ClO₄.⁵² Likewise, the cis β Fe(III) complex [Fe{tn(H)py}Cl₂]ClO₄ (tn = trimethylenediamine) prepared by Busch and co-workers was isolated as yellow prisms.¹⁰⁶ The other ligands were not amenable to Fe(III) complex synthesis; brown solids were often obtained. Though no structural information was obtained for these paramagnetic species, Fe(II) and Fe(III) complexes of related quadridentate ligands with pyridyl or quinoline and 3° donors all possess cis α structures in both monometallic and bimetallic complexes.^{51,59,107} Complexes previously prepared by others have found application as bioinorganic models^{51,52,107} and catalysts, 59,108 and they have been used in studies of spin equilibria in diisothiocyanate complexes.86,87

Manganese Complexes. Reaction of MnCl₂ with quadridentate ligands **1**–**4** produced off-white paramagnetic solids in high yield. White Mn(II) complexes of analogous ligands have been described previously as intermediates in the synthesis of bimetallic complexes⁵³ including many mixed valence species intended to model biological systems or for studying magnetic interactions between bridging ligands.^{109–112} However, no characterization of these intermediates was provided. Related bimetallic Mn complexes have been utilized as oxidation catalysts.¹⁰⁸ As expected for high-spin d⁵ systems, electronic absorption spectra of manganese complexes show ligand bands but no absorptions attributable to d–d transitions. A magnetic moment of 6.1 $\mu_{\rm B}$ was determined for [Mn{en(Bn)py}Cl₂] using the Evans NMR method. This is somewhat higher than the spin only value of 5.9 $\mu_{\rm B}$ calculated for this d⁵ system.

Electrochemistry. Cyclic voltammetry is a useful technique for assessing the purity of samples and for probing the electronic features of metal complexes. Methylene chloride solutions of the complexes and ligands were investigated using a glassy carbon working electrode, an aqueous silver/silver chloride reference electrode, and tetra-*n*-butylammonium hexafluorophosphate (TBAH) as the supporting electrolyte. Data are collected in Table 5. Free diamine ligands undergo oxidation

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 Table 5. Cyclic Voltammetric Data^a

compd	$E_{\rm p,a}\left({\rm V}\right)$	$E_{\rm p,c}\left({\rm V}\right)$	$E_{1/2}(V)$	$\Delta E_{\rm p}^{\ b} ({\rm mV})$
en(Bn)py	+1.05			
en(F ₅ Bn)py	+1.08			
S-pn(F ₅ Bn)py	+1.28	-1.24		
S-pn(F ₅ Bn)qn	+1.06			
$Mn\{en(Bn)py\}Cl_2$	+0.95	+0.86	+0.91	90
$Mn\{en(F_5Bn)py\}Cl_2$	+1.08	+1.00	+1.04	90
$Mn{S-pn(F_5Bn)py}Cl_2$	+1.09	+1.00	+1.04	90
$Fe\{en(Bn)py\}Cl_2$	+0.40	+0.32	+0.36	85
$Fe\{en(F_5Bn)py\}Cl_2$	+0.48	+0.41	+0.44	75
$Fe\{en(Bn)py\}Cl_2^c$	+0.18	-0.019	+0.082	202
$Fe\{en(F_5Bn)py\}Cl_2^c$	+0.26	+0.15	+0.20	112
$[Fe{en(F_5Bn)py}Cl_2]PF_6$	+0.46	+0.39	+0.42	70
$Fe{S-pn(F_5Bn)py}Cl_2$	+0.54	+0.45	+0.49	95
$Co\{en(Bn)py\}Cl_2$	+0.77	$+0.12^{d}$		
$Co{en(F_5Bn)py}Cl_2$	+0.86	$+0.26^{d}$		
$Co{S-pn(F_5Bn)py}Cl_2$	+0.89	$+0.26^{d}$		
$Co{S-pn(F_5Bn)qn}Cl_2$		fea	tureless	
$Ni\{en(F_5Bn)py\}Cl_2$	+1.40	+1.34	+1.37	65
$Ni{S-pn(F_5Bn)py}Cl_2$	+1.46	+1.36	+1.41	97
$Cu\{en(F_5Bn)py\}Cl_2$	+0.06	-0.11	-0.02	160
$Cu{S-pn(F_5Bn)py}Cl_2$	+1.25	-0.09^{e}		
$Zn\{en(F_5Bn)py\}Cl_2$		fea	tureless	
$Zn{S-pn(F_5Bn)py}Cl_2$	+1.28			

^{*a*} Instrumentation: BAS model CV-27 electrochemical analyzer. Working electrode: glassy carbon. Reference electrode: aqueous Ag/ AgCl. Concentration: 0.5 mM. Supporting electrolyte: 0.1 M TBAH in CH₂Cl₂. Scan rate: 200 mV/s. $E_{1/2}$ for the ferrocenium/ferrocene couple is at +0.47 V vs Ag/AgCl under these experimental conditions. ^{*b*} ΔE_p is defined here as $E_{p,a} - E_{p,c}$. ^{*c*} As in (a) with the following exceptions: Instrumentation: BAS model CV-50W electrochemical analyzer. Reference electrode: Ag/AgCl in CH₃CN. Scan rate: 100 mV/s. $E_{1/2}$ for the ferrocenium/ferrocene couple is at +0.23 V vs Ag/ AgCl under these experimental conditions. ^{*d*} Product wave. ^{*e*} Additional minor features present at +0.56 (anode) and +0.52 and +0.68 (cathode).

at high positive potentials (1.05-1.28 V), and for the S-pn(F₅-Bn)py ligand, 3, a reduction is also observed. It was of interest to us to determine which metal complexes exhibit reversible oxidations, as this could be important for certain catalytic reactions. For manganese, iron, and certain Cu complexes, quasireversible waves were observed. Manganese(II-III) couples appear at $E_{1/2} \sim 0.91 - 1.04$ V, whereas oxidation of Fe(II) to Fe(III) is more facile under similar experimental conditions ($E_{1/2}$ $\sim 0.36-0.49$ V). While the CV of the Cu(II) complex of the bulky S-pn(F₅Bn)qn ligand, 4, contains a quasireversible oxidation, other copper complexes are not as well behaved. A series of weak oxidations and reductions are evident at different potentials, including ones at -0.43 V (vs ferrocene) indicative of free copper. This may be contrasted with the findings of Urbach and co-workers on related copper complexes, albeit with different solvent systems.94 For Ni(II) complexes, weak quasireversible processes are evident at high potentials (\sim 1.4 V). The Ni(III) oxidation state is typically highly unstable; often Ni(II)/Ni(III) oxidations occur in the range of +1 to +1.5 V.^{113–116} As is expected for a d¹⁰ system, Zn complexes are essentially featureless over the potential range under investigation.

For catalytic systems, it is desirable to be able to tune the electron richness of the metal center. Thus, determining whether, and to what extent, complexes of the fluorinated systems 2-4



Figure 8. Comparison of the cyclic voltammograms for (a) [Fe{en-(Bn)py}Cl₂] and (b) [Fe{en(F₅Bn)py}Cl₂].

are more difficult to oxidize than analogous complexes of the nonfluorinated en(Bn)py ligand, **1**, is also important. The expected trends are clearly evident in $E_{1/2}$ values for Mn and Fe complexes of ligands **1** and **2** that are identical except for perfluorination of the phenyl rings in **2**. For Mn, the fluorinated ligand is harder to oxidize by ~130 mV whereas $\Delta E_{1/2} = 80$ mV for the two Fe complexes in this series. Voltammograms of [Fe{en(Bn)py}Cl₂] and [Fe{en(F₅Bn)py}Cl₂] that were recorded versus a nonaqueous Ag/AgCl reference electrode are compared in Figure 8.

Steric features of the ligands also influence how easily metal centers are oxidized. Few differences are observed between complexes of en(F₅Bn)py, 2, and S-pn(F₅Bn)py, 3; however, complexes of the bulky quinoline ligand may exhibit rather different electrochemistry, as is dramatically demonstrated for cobalt complexes. Complexes of the structurally similar ligands 1-3 undergo irreversible oxidations at 0.77-0.89 V; however, the voltammogram recorded for the Co complex of 4 is featureless throughout. The bulky quinoline ligand is less able to accommodate the contracted Co(III) oxidation state, perhaps due to unfavorable steric interactions between apical quinoline groups and equatorial chloro ligands that are undoubtedly necessary for stabilizing the octahedral geometry preferred by Co(III). Thus, $[Co{S-pn(F_5Bn)qn}Cl_2]$ resists oxidation, both chemically, using common oxidants such as Br₂ or H₂O₂, and electrochemically.

Conclusion

These investigations further demonstrate how readily the steric and electronic features of the amino-diimine family of ligands and their complexes may be varied. This, combined with the fact that appropriately designed ligands form C_2 -symmetric cis α topologies and single (or major) diastereomers upon complexation to metal ions should make them well suited for

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applications in asymmetric catalysis. While metals such as Fe, Co, and Mn seem to adopt similar geometries throughout most of the ligand series, other metals such as Zn, Cu, and, to some extent, Ni are far more sensitive to the steric and electronic features of the ligand and can assume six, five, or even four coordinate structures. These differences could be exploited in reaction optimization. Moreover, varying the ancillary ligands from halides to other species, or replacing some or both of them with noncoordinating counterions, opens up even more possibilities for tailoring reactivity. Once complexes are proven viable catalysts for specific transformations, considerably more work is merited on these systems. Devising alternate routes to complexes that were difficult to access by standard preparations reported herein and further varying the steric and electronic features of these ligands, as well as more detailed characterization, are all worth pursuing. Recently we have utilized Cu complexes of this family of ligands as catalysts for the controlled polymerization of methyl methacrylate.¹¹⁷ We have also seen moderate enantioselectivities using Cu complexes of chiral quadridentate ligands as catalysts for the Mukaiyama aldol

reaction.⁷⁵ Rieger has described their potential to serve as catalysts for olefin polymerization.⁵⁹ Yet these few examples barely begin to explore the potential of cis α complexes as catalysts. Numerous other reactions including oxidations, polymerizations, and Lewis acid-catalyzed transformations are ripe for exploration.

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Supporting Information Available: A ¹H NMR spectrum (300 MHz, CDCl₃) for [Co{en(F₅Bn)py}Cl₂]Br⁻¹/₂CH₃CH₂OH⁻¹/₂CH₂Cl₂ and IR spectral data for ligands and complexes measured as Nujol mulls (2 pages). An X-ray crystallographic file in CIF format for [Cu{en(F₅Bn)py}Cl₂]. This material is available free of charge via the Internet at http://pubs.acs.org.

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