Basicity of Coordinated Pyrazine and Bonding Interactions with $[M^{II}(CN)_5]^{3-}$ Fragments (M = Fe, Ru, Os)

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Introduction

Modification of the basicity of pyrazine (pz) upon coordination to [MX₅] fragments was related to the back-bonding capabilities of the latter species. Saturated coligands such as ammines change the p K_a of free pzH⁺ (0.6) to 2.5 and 7.4 for M = Ru(II)² and Os(II)³, respectively. The p K_a values depend on the bonding properties of the coligands for a given metal, as shown by the upward and downward shifts for [Os^{II}A₅pzH]³⁺ (A = ammine) when one of the A ligands is substituted by a π -donor (chloride) or a π -acceptor (dinitrogen) ligand, respectively.

A current interest exists in establishing the bonding trends for the $[M^{II}(CN)_5L]^{n-}$ series (M = Fe, Ru, Os).⁵⁻⁸ The pK_a values quoted for L = pzH^+ in the iron (0.065) and ruthenium (0.4) complexes were composite values, associated to competitive and successive equilibria on pyrazine and cyanides. These results disagreed with NMR measurements, suggesting that iron could be more strongly back-bonding than ruthenium,⁷ a result substantiated later for the L = dimethylpyrazine derivatives.⁹

After the synthesis of the $[Os^{II}(CN)_5pz]^{3-}$ ion, 8a it seemed worthwhile to extend the basicity studies and revisit the titration experiments with the iron and ruthenium analogues. Using an adequate acid—base description and a chemometrical methodology, 10 we present pK_a values for the mono- and diprotonated species in the three complexes, disclosed UV–visible spectra and reliable separate pK_a values for the pz- and cyanide-protonated species. Complementary NMR measurements support the analysis of the bonding trends for the $[M^{II}(CN)_5pz]^{3-}$ species.

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Experimental Section

Salts of the Pentacyano-L Complexes. $Na_3[FeCN)_5pz] \cdot 4H_2O$, $K_3[Ru(CN)_5pz] \cdot xH_2O$, and $Na_3[Os(CN)_5L] \cdot xH_2O$ (L=py, pz, dmpz, N-methylpyrazinium (mepz⁺)) were prepared as previously described. All other chemicals were of analytical grade. NMR spectra were recorded on a Bruker AC200 spectrometer. The pK_as for the different protonation equilibria were determined spectrophotometrically with a Hewlett-Packard 8452A instrument. Solutions containing weighed amounts of the corresponding salts of the $[M(CN)_5pz]^{3-}$ ions (around 10^{-5} M) and varying concentrations of HCl were prepared (pH range 0-7, I=1 M, NaCl). The UV—vis spectra were obtained quickly after adding the acid and diluting to volume, using a 10.00 cm quartz cuvette. Repetitive scans were performed on each solution to ensure stability under acid conditions.

An adequate characterization of the species in solution was achieved by using the SPECFIT software package. It consists of a global least-squares fitting routine for equilibrium and kinetics experiments that uses factor-analysis decomposition methods;¹⁰ this was useful to obtain the single species' spectra along with the equilibrium constants.

Results and Discussion

Figure 1 shows the titration spectra for the [M^{II}(CN)₅pz]³-complexes. For the three metals the behavior is similar, showing multistep protonation: Figure 1a shows the spectra of solutions of Na₃[Fe(CN)₅pz]·4H₂O. The MLCT band at 452 nm⁶ decays for decreasing pH's, while new bands develop at 408 and 636 nm, with isosbestic points at 272, 321, 418, and 544 nm (range 7.0–2.0). In the pH range 2.0–0.0 the isosbestic points disappear and the maxima of both bands shift to lower wavelengths (378 and 556 nm). The ruthenium (and osmium) complexes display similar behavior. Figure 1b (1c) shows that when going from pH 7 to 2 the MLCT bands at 372⁷ (384)^{8a} nm decay, being replaced by two absorptions at 342 (345) and 504 (514) nm. Upon further acidification these bands are replaced by new absorptions at 322 (315) and 444 (460) nm.

For the three metals, the factor analysis displays *only* three colored species, consistent with a two-step protonation scheme:

$$[M^{II}(CN)_5pz]H_2 \xrightarrow{K_2} [M^{II}(CN)_5pz]H \xrightarrow{K_1} [M^{II}(CN)_5pz]$$

The concentration profiles depending on [H⁺] are given by

$$[M^{II}(CN)_5pz] = \frac{C_0K_1K_2}{[H^+]^2 + K_2[H^+] + K_1K_2}$$
(1)

$$[M^{II}(CN)_{5}pz]H = \frac{C_{0}[H^{+}]K_{2}}{[H^{+}]^{2} + K_{2}[H^{+}] + K_{1}K_{2}}$$
(2)

$$[M^{II}(CN)_5pz]H_2 = \frac{C_0[H^+]^2}{[H^+]^2 + K_2[H^+] + K_1K_2}$$
(3)

Leaving K_1 and K_2 as fitting parameters, we retrieve the spectra of the three species (displayed as insets in Figure 1) and the following results for p K_1 : 1.84 \pm 0.05 (Fe), 1.44 \pm 0.05 (Ru), 1.88 \pm 0.04 (Os). For p K_2 we obtain 0.4 \pm 0.1 (Fe), 0.3 \pm 0.6 (Ru), $-0.2 \pm$ 0.6 (Os).

These pK_1 and pK_2 values do not reflect clean, separated protonation steps at pz or cyanide ligands. The *two* new MLCT

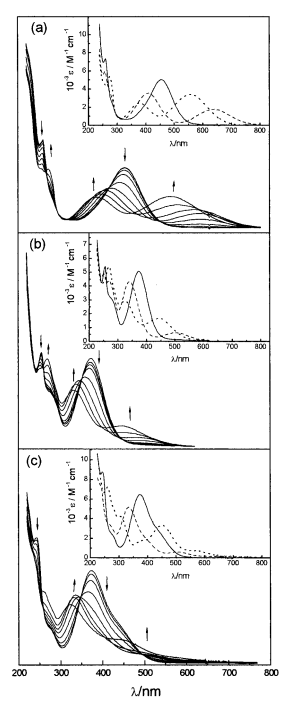


Figure 1. Spectrophotometric acid—base titration of the $[M^{II}(CN)_5pz]^{3-}$ complexes ((a), (b), (c) correspond to Fe, Ru, Os, respectively), c = ca. 1×10^{-5} M, I = 1 M (NaCl), pH range = 0-7 (HCl). Arrows indicate the changes observed on increasing acidity of the medium. Insets: Spectra obtained for the un-, mono-, and diprotonated species (full, dashed, dotted lines, respectively).

bands in the monoprotonated species spectra show that *both* pz and cyanide are affected by the *first* protonation step.

$$\begin{split} [M^{II}(CN)_5pzH]^{2-} & \xrightarrow{K_{MI.H}} [M^{II}(CN)_5pz]^{3-} + H^+ \\ \\ [M^{II}(CN)_4(CNH)pz]^{2-} & \xrightarrow{K_{HMI.}} [M^{II}(CN)_5pz]^{3-} + H^+ \end{split}$$

The concentration ratio of monoprotonated species is pH independent, hence behaving as a pure monoprotonated moiety, with a *global* acidity constant K_1 , related to the *individual* K_{HML}

Table 1. Absorption Maxima (nm), Disclosed Individual pK_a Values, and ¹H NMR Chemical Shifts for $[M(CN)_{spz}]^{3-}$ Systems

	Fe	Ru	Os
$\lambda_{\text{max}}([M(CN)_5pz]^{3-})^a$	452 (5011)	372 (5040)	384 (6435)
$\lambda_{\text{max}}([M(\text{CN})_5\text{pzH}]^{2-})^b$	636 (6900)	504 (6540)	514 (12700)
$\Delta E (\mathrm{cm}^{-1})^c$	6400	7040	6600
$\lambda_{\text{max}}([(\text{HNC})\text{M}(\text{CN})_4\text{pz}]^{2-})^d$	408 (4710)	342 (4750)	345 (5380)
$\lambda_{\text{max}}([(\text{HNC})\text{M}(\text{CN})_4\text{pzH}]^-)$	556	444	460
$\lambda_{\text{max}}([(\text{HNC})_2\text{M}(\text{CN})_3\text{pz}]^-)$	378	322	315
pK_{MLH^e}	1.24 ± 0.07^{f}	0.41 ± 0.06^{f}	0.53 ± 0.05
pK_{HML}^e	1.71 ± 0.09	1.40 ± 0.06	1.86 ± 0.04
$\delta(CH)\alpha^g$	9.09^{h}	8.72, $dd^{h,i}$	9.11, dd ^j
δ (CH) β ^g	8.24^{h}	8.45, $dd^{h,i}$	8.36, dd ^{<i>j</i>}

^a Values in parentheses are the molar absorbances ϵ (M⁻¹ cm⁻¹). ^b Values in parentheses are estimated molar absorbances (see text). ^c Energy difference between the absorption maxima for [M(CN)₅pz]³⁻ and [M(CN)₅pzH]²⁻. ^d Molar absorbances calculated according to eq 5. ^e This work, I=1 M, 25 °C. ^f 0.065 (Fe); 0.4 (Ru), in ref 7. ^g δ (ppm) in D₂O vs TMS. Data for free pz: 8.69. Data for [Co^{III}(CN)₅pz]²⁻: 8.99(α), 8.72(β). ⁷ h Reference 7. ⁱ A₂X₂ system, with ³J_{HαHβ} = 3.3 Hz and ⁵J_{HαHβ} = 1.3 Hz. ^j A₂X₂ system, with ³J_{HαHβ} = 3.0 Hz and ⁵J_{HαHβ} = 1.5 Hz.

and $K_{\rm MLH}$ by

$$K_1 = \frac{K_{\text{MLH}}K_{\text{HML}}}{K_{\text{MLH}} + K_{\text{HML}}} \tag{4}$$

The individual species contribute to the observed spectra according to

$$\epsilon_{\text{obs}}(\lambda) = \frac{K_{\text{HML}}\epsilon_{\text{MLH}}(\lambda) + K_{\text{MLH}}\epsilon_{\text{HML}}(\lambda)}{K_{\text{MLH}} + K_{\text{HML}}}$$
(5)

In other words, even if protonation can take place at *two* nonequivalent positions, the spectrophotometric titration experiment reveals *one* "monoprotonated" species, whose spectrum is a combination of the corresponding cyano- and pz-protonated ones. Analogously, the spectra of the "diprotonated" species are combinations of those of $[(HNC)M(CN)_4pzH]^-$ and $[(HNC)_2M(CN)_3pz]^-$.

Independent values of pK_{MLH} and pK_{HML} can be calculated from the pK_1 values by estimation of the molar absorbances of the $[M^{II}(CN)_5pzH]^{2-}$ species (Table 1).¹¹ The reliability of the global pK_a values is supported by the analysis of the resolved spectra (insets, Figure 1). The MLCT bands of the $[M^{II}(CN)_5pz]^{3-}$ species split into two separate absorptions upon protonation. We assign the bathochromically and hypsochromically shifted bands to the $[M^{II}(CN)_5pzH]^{2-}$ and $[(HNC)M^{II}(CN)_4pz]^{2-}$ complexes, respectively.⁶ The calculated molar absorbances are greater (pz) and smaller (cyanide) than for the corresponding unprotonated species, as expected.⁶

Table 1 shows that the energy differences for the unprotonated and pz-protonated MLCT bands (ΔE) are around 6000–7000 cm⁻¹, but depend significantly on M, namely: $\Delta E(\text{Fe}) < \Delta E(\text{Os}) < \Delta E(\text{Ru})$. In the limit of no mixing between the M and $\pi^*(L)$ orbitals, the ΔE values should be independent of M, corresponding to the energy differences of $\pi^*(L)$ for the pz and pzH⁺ ligands. The observed ΔE values are indicative of backbonding, decreasing in the sense Fe > Os > Ru. The sensitivity of the MLCT band energy to changes in L (or M) decreases

⁽¹¹⁾ The molar absorbances for the [M^{II}(CN)₅pzH]²⁻ species were interpolated by plotting log(E_{MLCT}) (the energy of the MLCT bands) against the molar absorbance of the corresponding absorptions for a series of [M^{II}(CN)₅L]ⁿ⁻ complexes (M = Fe, Ru, Os). Then the values of ϵ for the [M^{II}(CN)₄(CNH)pz]²⁻ species were obtained (eq 5).

when the back-bonding interaction between M and $\pi^*(L)$ increases. Besides, inspection of the individual p $K_{\rm MLH}$ (Table 1) and of the chemical shifts for the β -protons in the pz-ring^{12,13} indicate the same trend. All this evidence suggests a decrease in back-donation to pz when going from Fe to Os and Ru.

Table 1 also shows the pK_{HML} values for the cyanidemonoprotonated species, with Os > Fe > Ru. Now the osmium center appears as the most basic, probably reflecting the synergistically behaved, strong σ - and π -interactions between osmium and cyanides, dominant in the overall bonding scheme. 8a Note that this is also the order of NMR chemical shifts for the pz α -protons. $^{12-14}$

Although none of our measurements gives a quantitative approach to individual σ - or π - contributions, we consider that π -interactions are determinant of the Fe-Os-Ru order found for the p K_a values, electronic and NMR (β) shifts of bound pzH⁺. The π -interaction with pz appears as the strongest for iron, suggesting a dominant influence of the more favorable energy match between d_{Fe} and π^*_{pz} orbitals over the lesser overlapping capability of the 3d orbitals.¹⁵

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- (13) Upfield chemical shifts for the β -protons and downfield shifts for the α-protons seem to be characteristic of N-heterocyclic ligands in negatively charged MX₅ moieties. 12 A second-order influence of the σ - π interactions also operates at the α -positions; however, the β (and γ) positions show to be more sensitive to π -back-bonding, because of attenuation of the σ -interactions with distance. For $[Co^{III}(CN)_5pz]^{2-1}$ (no back-bonding and lower overall charge than the others), both α and β -protons shift downfield (Table 1), as was the case with the 2-Mepz complexes;¹² thus, the upfield shifts for the Fe, Ru, and Os pz-complexes seem to be really traced to the π -interactions.
- (14) As a check of consistency, we also made titrations with other [Os(CN)₅L]ⁿ⁻ complexes where only cyanide was protonable. The results for $L = py (pK_a 2.33)$ and $L = mepz^+ (pK_a = 1.05)$, together with the pz value, show that the cyanide-basicity decreases with the π -acceptor capability of L, as expected (Supporting Information, Figure
- (15) The values of E for the $[M^{III}, II(CN)_5pz]^{2, 3-}$ redox couples are: 0.55 (Fe); 60.96 (Ru), cf. Henderson, W. W.; Shepherd, R. E. Inorg. Chem. **1985**, 24, 2398; 0.78 (Os). 8a The extent of back-donation, Fe \geq Ru, was previously proposed for the dmpz-complexes.9 After measuring the [Os(CN)₅dmpzĤ]²⁻ complex, and using data taken from Figure 1

The insets of Figure 1 (dotted lines) show the spectra of the diprotonated species, which are mixtures of the [MII(CN)₄-(CNH)pzH]⁻ and [M^{II}(CN)₃(CNH)₂pz]⁻ ions. Their corresponding bands are both shifted to lower wavelengths compared to the associated monoprotonated ions (dashed lines), as expected. The global pK_2 values are also consistent, although a calculation of individual pK values (around zero) is associated to large

As a conclusion, the basicities at the exposed end of pz or cyanides are slightly dependent on M; the p K_a order Fe > Os > Ru, agrees with the NMR results. The pK_a values for M = Ru and Os are significantly smaller than those found for the [M^{II}(NH₃)₅pz]²⁺ ions (see above),^{2,3} reflecting the competitive influence of the cyanide coligands, as adressed by our recent calculations on the pz-basicities for the [Ru(CN)₅pz]³⁻ and [Ru-(NH₃)₅pz]²⁺ aqueous species, and the increased Mulliken population on the π^*_{pz} orbital for the latter compound.¹⁶

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Supporting Information Available: Figures showing spectrophotometric acid-base titrations of $[Os(CN)_5L]^{n-}$ ions $(L = py, mepz^+)$ and $[M(CN)_5 dmpz]^{3-}$ ions (M = Fe, Ru, Os). This material is available free of charge via the Internet at http://pubs.acs.org.

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in ref 9, we found 2.68 (Fe), 1.51 (Ru), and 1.99 (Os) for the pK_a values i.e., the same order of basicities as found with the pz-complexes. The numbers for Fe and Ru are slightly different from those of ref 9, due to the adequate consideration of simultaneous cyanide protonation in the present work (Supporting Information, Figure S2).

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