141. Fluxional Behaviour of Isocyanide Derivatives of Dodecacarbonyltetr airidium

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The fluxional behaviour of $[Ir_4(CO)_{11}(t-BuNC)]$ **(1)** and $[Ir_4(CO)_{11}(ArNC)]$ (Ar = 4-methoxyphenyl, 4-tolyl, phenyl, 4-chlorophenyl, 4-nitrophenyl; $2-5$) has been investigated by 2D and variable-temperature ¹³C-NMR. These techniques give new evidence of the processes responsible for CO exchange, namely two successive merry-gorounds causing Cotton-like scrambling and a CO rotation about one metal center.

Introduction. – The synthesis of two series of isocyanide derivatives of $\text{[Ir}_{4}(\text{CO})_{12}$, namely $[\text{Ir}_4(\text{CO})_{12-x}\text{L}_x]$ (x = 1–4, L = t-BuNC, MeNC) has been reported by *Stunz* and *Shapley* [1]. Carbonyl scrambling in $[Ir_4(CO)_{11}(t-BuNC)]$ (1) has been studied by these authors through variable-temperature 13C-NMR without simulation. Two mechanisms were proposed for the dynamic processes of lowest activation energy, consisting of two merry-go-rounds taking place, simultaneously, on two different faces of the Ir₄ tetrahedron.

We have re-examined the fluxionality of *1,* together with that of five new isocyanide derivatives $[Ir_4(CO)_{11}(ArNC)]$ (Ar = 4-methoxyphenyl, 2; 4-tolyl, 3; phenyl, 4; 4chlorophenyl, 5 ; 4 -nitrophenyl, 6) by ¹³C-NMR. The distinction between possible mechanisms is now based on dynamic connectivities derived from 2D-NOESY spectra and backed up by simulation of the variable-temperature ¹³C-NMR spectra up to complete coalescence. The ArNC ligands in $2-6$ were selected for their different γ parameters [2] to examine a possible effect of ligand basicity on the fluxional processes.

Results and Discussion. – The direct reaction of $[Ir_4(CO)_{12}]$ with RNC (R = t-Bu, Me) in the presence or absence of Me, NO in THF gives low yields of $[Ir_4(CO)_{12}](RNC)$, $(x = 1, 2)$. We have found that the reaction of $[\text{Ir}_4(CO)_{11}][\text{NEt}_4]$ with AgBF₄ and 1 mol-equiv. of isocyanide in CH₂Cl₂ gives better yields of $[Ir_4(CO)_{11}(RNC)]$ (88-96%) under milder conditions and without need of prep. TLC.

Cluster **1** adopts an unbridged *C,* structure in the solid state [3]. This structure is also adopted by compounds 1–6 in solution as only IR absorptions in the region (2140–2200) $\rm cm^{-1}$) appropriate for terminal isocyanide ligands and in the range (2000–2100 cm⁻¹) suitable for terminal CO ligands are observed.

The I3C-NMR spectra of **1-6** display in each case one single carbonyl resonance around 310 K. The limiting low-temperature 13 C-NMR spectra show five resonances,

 $\sum_{i=1}^{n}$ Taken from the doctoral dissertation of *A. O.*, Université de Lausanne.

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each of relative intensity two, and one resonance of unit relative intensity, all of which appear in the terminal CO region (see *Exper. Part*). The signal of relative intensity one is assigned to the unique carbonyl ligand f in pseudo-trans-position with respect to the isonitrile ligand. Inspection of the COSY ¹³C-NMR spectra of samples enriched with I3CO *(ca.* **30%;** *e.g. Fig. 1* for I3C-enriched **1)** leads to the assignment of the signal of

Fig. I. *COSY I3C-NMR spectrum of "C-enriched*

relative intensity two, and devoid of couplings to carbonyl ligands *b.* It is indeed well known that in carbonyltetrairidium clusters, the only CO's which couple significantly are in relative pseudo-*trans*-positions (in particular, geminal $^{2}J(CO,CO)$ are very small) [1]. The remaining four signals with cross peaks have identical vicinal *3J(C0,CO)* coupling constants (11.4 \pm 0.8 Hz) and cannot be individually assigned; hence they are labelled *I*, 2, *4,* and **5** in order of decreasing chemical shift. Among the 24 virtually possible assignments, only eight are consistent with the couplings *1,4* and *2,5.*

Fig. 2. *NOESY* ^{*I3C-NMR spectrum of ¹³C-enriched* **1** *in CD₂Cl₂ a) at 215 K* (mixing time 50 ms); *and* b) *at 225 K*} (mixing time **65.5** ms)

The NOESY¹³C-NMR spectrum of 1 at 215 K (Fig. 2a) gives the dynamic connectivities $I \leftrightarrow b \leftrightarrow 4$ and corresponds to the exchange matrix of a first merry-go-round of 6 CO's around either the Ir(1)-Ir(2)-Ir(3) face (for 4 of the 8 possible assignments) or the Ir(2)-Ir(3)-Ir(4) (for the remaining ones). The NOESY spectrum at 225 K *(Fig. 26)* shows 4 additional cross peaks which lead to the dynamic connectivities $1 \leftrightarrow 2 \leftrightarrow b \leftrightarrow 5 \leftrightarrow 4$ which corresponds to a second merry-go-round taking place on any of the equivalent Ir(1)-Ir(3)-Ir(4) or Ir(1)-Ir(2)-Ir(4) faces. The eight possible assignments are in agreement with such a mechanism. A merry-go-round of 5 CO's has never been observed when the ground state of the Ir₄ cluster contains a face bearing a radial L ligand, 3 edge-bridging, and two terminal CO's [4]. The proposed *Scheme* shows that the intermediate of the second merry-go-round in $[Ir_4(CO)_{11}(RNC)]$ contains a radial RNC ligand. However, as

the ground state has here all terminal ligands, the only condition to be satisfied in the fluxional process is to end up with a state of equal energy with the same pseudo-trans relationship between RNC and carbonyl ligand *f* and without constrains on the choice of the face for the merry-go-round. This condition doesn't necessarily imply the participation of the RNC ligand in the CO site exchanges.

At *ca.* 270 K, a third process involving carbonyl ligand f in a 3 CO's exchange around one metal center $(\text{Ir}(4))$ is taking place.

Line-shape analysis [5] of the variable-temperature ¹³C-NMR spectra was carried out using the following *Kubo-Sack* matrix elements: $(I,I) = (4,4) = -k_1/2$ $-k_2/2$, $(5,5) = -k_2 - k_3/2$, $(5,f) = k_3/2$, $(f,f) = -k_3$, $(f,5) = k_3$, with k_1, k_2, k_3 = rate constant in s⁻¹ of the first and the second merry-go-round processes and of the rotation of 3 CO's at one metal center, respectively. An example of the obtained and simulated spectra is given in *Fig. 3.* $(1,2) = (2,b) = (b,5) = (4,5) = k_2/2$, $(1,b) = (b,4) = k_1/2$, $(2,2) = -k_2$, $(b,b) = -k_1 -k_2$,

A simulation using matrix elements derived from the mechanism proposed by Stunz *et al.* [l] failed to reproduce variable-temperature I3C-NMR spectra of **1.** However, other possible CO site exchange mechanisms should also be considered. First, one should take into account a process involving bridging of the isocyanide ligand, since edge-bridging RNC geometries have been observed in di- and trinuclear complexes [6]. Bridging of RNC on the Ir(1)-Ir(2) or Ir(1)-Ir(3) edge involved in merry-go-rounds around the Ir(1)-Ir(2)-Ir(4) or Ir(1)-Ir(3)-Ir(4) faces, respectively, gives an exchange matrix different from that corresponding to the observed connectivities of the second merry-go-round *(Fig. 26).* This hypothetical matrix failed to reproduce experimental spectra.

Second, we have seen that the mechanism proposed for the lowest-energy process *(Fig. 2a)* is a merry-go-round through a CO-bridged intermediate having RNC either in axial or in apical position. When the ground state of Ir_4 clusters has 3 edge-bridging CO's defining a basal plane, as in $[Ir_4(CO)_{11}(PEt_3)]$ [7], the relative energies of the 3 possible isomers follow the sequence $L(axial) > L(adial) > L(apical)$. If we assume that the same sequence of energy levels is applicable to the postulated intermediates of the first merrygo-round in $[Ir_4(CO)_{11}(t-BuNC)]$ (1), this same process could be due to a merry-go-round on the Ir(1)-Ir(2)-Ir(3) face, followed by another on the Ir(2)-Ir(3)-Ir(4) face. This alterna-

Fig. **3.** a) *Experimental and* b) *simulated variable-temperature I3C-NMR spectra of "C-enriched* **1** *in CD,CI,*

tive mechanism agrees with the observed NOESY **I3C-NMR** spectra of **1** at 225 K *(Fig. 2b),* if second-order peaks are taken into account. Therefore, a quantitative treatment of the 2D spectrum was needed. Recently, *Orrell* and coworkers *[8]* have proposed a general method for evaluating rate constants in complex exchange networks from two-dimensional NOESY NMR spectra. We have used their D2DNMR computer program capable of performing signal intensity to exchange rate calculation to distinguish between first- and second-order peaks. Including, as input data, the number of exchanging sites **(6),** their relative populations, and the experimental 2D signal intensities based on integration of all 2D peaks, a kinetic matrix was derived (see *Supplementary Material)* whose elements are the rate constants of the first-order exchange process. This matrix was equivalent to that of the proposed second merry-go-round, but not to that of the above alternative mechanism.

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| in $[Ir_4(CO)]_1(RNC)$ at 298 K | | | | | | | | |
|---------------------------------|--------------------|----------------------|------------------------|---------------------|------------------|------------------------|------------------------|----------------------|
| R | | $\varDelta G^{\neq}$ | $\varDelta G_2^{\neq}$ | ΔG_2^{\neq} | | $\varDelta G_1^{\neq}$ | $\varDelta G_2^{\neq}$ | $\varDelta G_2^{\#}$ |
| $\mathbf{1}$ | $t - Bu$ | 47.5 | 48.6 | 60.7 | 4 phenyl | 49.4 | 49.9 | 60.8 |
| | 2 4-methoxyphenyl | 48.8 | 49.8 | 59.3 | 5 4-chlorophenyl | 50.6 | 51.6 | 60.6 |
| | $3 \quad 4$ -tolyl | 49.0 | 50.4 | 59.9 | 6 4-nitrophenyl | 49.8 | 53.2 | 62.4 |

Table. *Free Enthalpies of Activation* ΔG^{\neq} (± 0.4 kJ mol⁻¹) *of the 3 Fluxional Processes in [Ir4(CO),,(RNC)] at 298 K*

The observed fluxional behaviour of cluster compounds **2-6** is the same as that of **1,** and can be viewed as a CO site exchange through two consecutive merry-go-rounds, followed by the rotation of 3 CO's around one metal center. The energy barriers are given in the *Table.*

The basicity of the non-CO ligand has no significant effect on the $\Delta G^*_{\gamma_{\rm 98}}$ of the exchange processes, as also observed in $[Ir_4(CO)_{11}X]^-$ (X = Br, I) and $[Ir_4(CO)_{11}(PAr_3)]$ systems [9]. The rotation of apical CO's is also the highest activation energy process in the halogeno and phosphine clusters which have a different ground-state geometry.

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Experimental Part

1. *General.* IR spectra: *Perkin-Elmer 880* spectrophotometer equipped with a data station; 0.1-mm CaF₂ cells previously purged with N₂. ¹³C-NMR spectra: *Bruker WH 360* (90.55 MHz); in CD₂Cl₂ at normal atmospheric pressure; δ in ppm, $J(C, C)$ in Hz from COSY spectra. 2D ¹³C-NMR spectra: COSY experiment in CD₂Cl₂ at 180 K, 256 t_1 increments with 2-K transients, spectral width 900.90 Hz in the F_2 domain and 450.45 Hz in the F_1 domain; NOESY experimets in CD₂Cl₂ at 215 and 225 K, mixing time 50 and 65.5 ms, resp., 512 increments with 2-K transients, spectral width 1199.0 Hz in the *F2* domain and 599.52 **Hz** in the *Fl* domain; a squared sine bell was applied in both domains prior to *Fourier* transformation.

2. *Isocyanide Ligands.* Commercially available t-BuNC was used *(Merck)* without further purification. The other ligdnds were synthesised starting from the corresponding formamides and following the method proposed by *Ugi* and coworkers [10]. When formamides were not commercially available (4-chloro-, 4-nitro-, and 4-methoxyphenyl-substituted ones), they were obtained from the corresponding amines according to [I I]

3. Monosubstituted Complexes. A soln. of $[Ir_4(CO)_{11}][NEt_4]$ [12] (200 mg, 0.15 mmol) and the isocyanide ligand (0.15 mmol) was vigorously stirred, after addition of $AgBF₄$ (32.1 mg, 0.165 mmol) for 1 h at 263 K. The mixture containing solid AgI was filtered through a 15×3 cm column packed with silica gel (70–230 mesh) and eluted with cold CH₂Cl₂. The yellow filtrate was evaporated and the residue crystallised from CH₂Cl₂/hexane 1:1 at 248 K.

(tert-Butyl isocyanide)undecacarbonyltetrairidium ([Ir₄(CO)₁₁(t-BuNC)]; 1): For spectral characteristics, see [I].

Uuhcacarbonyl(4-methoxyphenyl isocyanide)tetrairidium **(Ir4(CO)l,(4-MeOC6H4NC)];** 2): **1R** (CH,CI,, 298 K): 2091m, 2060vs,2038s, 2018s, 2009m (CO); 2149m (CN). I3C-NMR (CD,CI,, 190 K): 158.7 *(J* = 11.4, *I);* 156.9 $(J = 11.2, 2)$; 155.7 *(b)*; 155.3 $(J = 11.4, 4)$; 155.0 $(J = 11.6, 5)$; 154.4 *(f)*. Anal. calc. for C₁₉H₇Ir₄NO₁₂ (1210.14): C 18.86, H 0.58, N 1.16; found: C 19.20, H 0.59, N 1.17.

Undecacarbonyl(4-tolyl isocyanide) tetrairidium ([$Ir_4({\rm CO})_{11}(4\textrm{-MeC}_6\rm H_4NC)$]; 3): IR (CH₂Cl₂, 298 K): 2092m, 2058vs, 2037s, 2016s. 2007m (CO); 2165m (CN). "C-NMR (CD2C1,, 190 K): 158.6 *(1);* 156.9 (2); 155.7 *(h);* 155.3 (4); 154.9(5); 154.3(f); same J's as in 2. Anal. calc. for C₁₉H₇I_{r4}NO₁₁(1194.1): C 19.11, H 0.59, N 1.17; found: C 19.34, H 0.6, N 1.18.

Undecacarbonyl(phenyl isocyanide)tetrairidium ([Ir₄(CO)₁₁(PhNC)]; **4**): **IR** (CH₂Cl₂, 298 K): 2092m, 2058vs, 2037s, 2016s, 2008m(CO); 2164m (CN). "C-NMR(CD,CI,, 190K): 158.5 *(1);* 156.9 (2); 155.7(6); 155.3 (4);154.9 (5); 154.4 (f); same *J*'s as in 2. Anal calc. for C₁₈H₅H₅I_{T4}NO₁₁ (1180.1): C 18.32, H 0.43, N 1.19; found: C 18.70, H 0.44, N 1.19.

Undecacarbonyl(4-chlorophenyl isocyanide) tetrairidium ([Ir₄(CO)₁₁(4-ClC₆H₄NC)]; **5**): **IR** (CH₂Cl₂, 298 K): 2092m, 2059vs, 2038s, 2016m (CO); 2161s, 2008m (CN). ¹³C-NMR (CD₂Cl₂, 190 K): 158.3 (1); 156.9 (2); 155.6 *(b);* 155.1 *(4); 154.9 (5); 154.3 <i>(f);* same J's as in 2. Anal calc. for C₁₈H₄ClIr₄NO₁₁ (1214.6): C 17.8, H 0.33, N 1.16; found: C 18.5, H 0.34, N 1.16.

Undecacarbonyl(4-nitrophenyl *isocyanide*) *tetrairidium* ($[\text{Ir}_{4}(\text{CO})_{11}(4-\text{NO}_{2}\text{C}_{6}\text{H}_{4}\text{NC})]$; 6): IR (CH₂Cl₂, 298 K): 2093m, 2059vs, 2036s, 2014s, 2006m (CO); 2183m (CN). ¹³C-NMR (CD₂Cl₂, 190 K): 157.8 (1); 156.9 (2); 155.4 *(b)*; 155 *(4)*; 154.8 *(5)*; 154.4 *(f)*; same *J*'s as in **2**. Anal calc. for C₁₈H₄Ir₄N₂O₁₃ (1225.1): C 17.65, H 0.33, N 2.29; found: C 17.88, H 0.34, N 2.27.

Supplementary Material. – The kinetic matrices for the D2DNMR program is available from *R. Roulet* upon request.

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