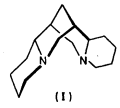
J.C.S. Dalton

Complexes of Some First-row Transition Elements with (-)-Spartein

By S. F. Mason* and R. D. Peacock, Chemistry Department, King's College, Strand, London WC2R 2LS

Complexes of cobalt(II), nickel(II), copper(II), and zinc(II), of the general formula [M(spartein) X_2], where X = CI-, Br-, or I-, have been prepared and characterised. Magnetic moments, electronic spectra, and the circular dichroism absorption (down to 5000 cm⁻¹) of [Ni(sp)Cl₂] are reported. The electronic structures of the cobalt(II) and nickel(II) complexes are discussed in terms of an effective C_{2v} chromophoric symmetry. Complexes of manganese-(III) and iron(III) have been obtained, but are less well characterised.

THE tetracyclic tertiary diamine, (—)-spartein, with the absolute configuration, 1,2 (6R, 7S, 9S, 11S), acts as a bidentate chelating ligand in the conformation (I). Complexes of (I) with organo-lithium and magnesium derivatives 3-5 and with zinc 6 have been postulated in the asymmetric synthesis of allenes 3,4 and carbinols.4-6 On the n.m.r. time-scale ligand-exchange in the dialkylmagnesium complexes of (I) is found to be slow.7 The biological activity 8 of (—)-spartein has been ascribed 7 to the sequestration of magnesium, and possibly other metal ions, by (I). In the present work we report that (-)-spartein (sp) forms complexes of the type, [M(sp)X₂], $X = Cl^-$, Br^- , or I^- , with cobalt(II), nickel(II), copper(II), and zinc(II), which are stable in ethanol and chloroform, and complexes with iron(III) and maganese(III) which are stable only in the solid state.



RESULTS AND DISCUSSION

The analytical results (Table 1) indicate that the complexes of the divalent metal ions are formulated

TABLE 1
Analytical results (%) a and magnetic moments

	~				[Leff/
	С	H	N	Halogen	B.M.
Ni(sp)Cl ₂	49.3(49.6)	7.0(7.2)	7.6(7.7)	18.1(19.5)	3.69
$Ni(sp)Br_{2}$	40.0(38.9)	5.8(5.8)	$6 \cdot 4 (6 \cdot 2)$	$35 \cdot 4(35 \cdot 3)$	3.66
Ni(sp)I2	33.3(32.9)	5·0(4·8)	5.4(5.1)	45·8(46·4)	3.47
Co(sp)Cl ₂	49.6(49.5)	7.2(7.2)	7.7(7.7)	17.0(19.5)	4·12 b
$Co(sp(Br_2))$	39.8(39.9)	5.9(5.8)	6.4(6.2)	37.5(35.5)	4·15 b
$Co(sp)I_2$	32.7(32.9)	4.9(4.8)	5.3(5.1)	45.6(46.4)	4·16 b
Cu(sp(Cl ₂	48.7(49.6)	7.0(7.1)	7.6(7.6)	20.6(19.2)	1.91
$Zn(sp)Cl_2$	48.3(48.6)	7.0(7.1)	7.5(7.6)	18.5(19.1)	
$Mn(sp)Cl_3$	39.9(45.5)	6.5(6.6)	$6 \cdot 4(7 \cdot 1)$	23.9(26.9)	
Fe(sp)Cl ₃	42.9(45.4)	6.7(6.6)	6.9(7.1)	24·1(26·5)	

^a Calculated values in parentheses. ^b Corrected for temperature independent paramagnetism.

 $[M(sp)X_2]$, $X = Cl^-$, Br^- , or I^- . The stoicheiometry is analogous to that of the dialkylmagnesium complexes ⁷

¹ F. Bohlmann, D. Schumann, and C. Arndt, *Tetrahedron*

Letters, 1965, 2705.

² S. Okuda, H. Kataoka, and K. Tsuda, Chem. Pharm. Bull.

Tokyo, 1965, 13, 491.

³ H. Nozaki, T. Aratani, and R. Noyori, Tetrahedron Letters, 1968, 2087.

⁴ H. Nozaki, T. Aratani, T. Toraya, and R. Noyori, Tetra-hedron, 1971, 27, 905.

with (I) and reflects the steric constraints upon the coordination of two (—)-spartein molecules to a metal ion. Iron(II) and maganese(II), under the same preparative conditions as used for the other metal ions, give complexes which analyse more satisfactorily for [M(sp)Cl₃]

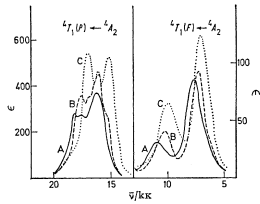


FIGURE 1 Electronic spectra in chloroform solution of A, [Co(sp)Cl₂]; B, [Co(sp)Br₂]; and C, [Co(sp)I₂]

than $[M(sp)Cl_2]$ (Table 1), suggesting an oxidation to the metal(III) state. The manganese complex may be polymeric, since it is insoluble in water and organic solvents. The iron complex dissolves sparingly in chloroform with some decomposition and, accordingly, the magnetic moment of the complex in this solvent, which was low and not well reproduced, is unreliable and is not reported. The presence of iron(III) in the complex is supported by the electronic spectrum which is typically that of a d^5 metal ion.

The magnetic moment (1.91 B.M.) of $[Cu(sp)Cl_2]$ falls within the range (1.8—2.0 B.M.) expected 9 for a mononuclear copper(II) complex. The electronic spectrum consists of one band at 7.7 kK (ϵ 260) and another at 12.4 kK (ϵ 270), which is consistent with a quasi-tetrahedral structure. A similar structure for the $[Co(sp)X_2]$ complexes is indicated by their magnetic moments (Table 1) and electronic spectra (Figure 1). The absorption band intensities are too large for octahedral coordination and the magnetic moments are appreciably

⁶ M. Guetté, J. P. Guetté, and J. Capillon, Tetrahedron Letters, 1971, 2863.

⁷ G. Fraenkel, C. Cottrell, J. Ray, and J. Russell, Chem. Comm., 1971, 273.

A. K. Reynolds in, 'The Alkaloids,' vol. V, ed. R. M. F. Manske, Academic Press, New York, 1955, pp. 123 and 197.
D. E. Billings and B. J. Hathaway, Co-ordination Chem.

⁹ D. E. Billings and B. J. Hathaway, Co-ordination Chem Rev., 1970, 5, 143.

⁵ H. Nozaki, T. Aratani, and T. Toraya, Tetrahedron Letters, 1968, 4097.

227 1973

larger than expected 10 for either square planar (2·1— 2.9 B.M.) or five-co-ordinate (1.9—2.5 B.M.) complexes of cobalt(II). By analogous criteria the magnetic moments (Table 1) and electronic spectra (Figure 2) of the [Ni(sp)X₂] complexes show that these also have a tetrahedral structure.

Models of the (-)-spartein complex with divalent transition metal ions indicate that the tetrahedral symmetry of the chromophore is lowered not only by the presence of two different pairs of ligand atoms but

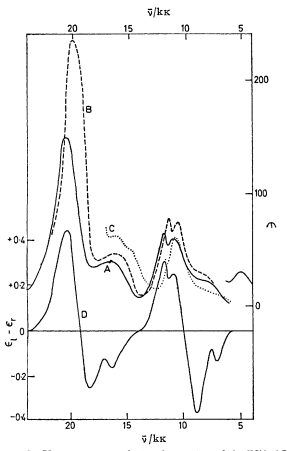


Figure 2 Upper curves: electronic spectra of A, [Ni(sp)Cl₂]; B, [Ni(sp)Br₂]; and C, [Ni(sp)I₂] in CHCl₃ solution and 4—6 kk in a KBr matrix ($\epsilon \times 5$); and lower curve, D, the c.d. spectrum of [Ni(sp)Cl₂] in CHCl₃ solution

also by the steric constraints of the diamine ligand. For the metal-nitrogen bond distance of 1.95 Å, found in the dichloro-di(p-toludine)cobalt(II) complex, 11 the N-M-N bond angle in [M(sp)Cl₂] is ca. 95° and if that angle has the tetrahedral value the metal-nitrogen bond distance is reduced to ca. 1.7 Å. The two sets of ligand atoms and the bond-length or bond-angle distortion reduce the effective chromophoric symmetry to C_{2v} in which the degeneracy of the T_1 and T_2 tetrahedral delectron states of the metal ion is completely lifted.

Three components are found in a number of the band

systems in the electronic spectra of the cobalt(II) and nickel(II) complexes, $[M(sp)X_2]$ (Figures 1 and 2). These spectra are similar to those of the corresponding complexes 12 [M(tmdm)X₂], where tmdm is a tetramethyl-N,N',N',N'-1,2- or -1,3-diamine, but are more fully resolved. The circular dichroism (c.d.) spectrum of the [Ni(sp)Cl₂] complex (Figure 2) reveals two component transitions, one at 9.0 kK and the other at 18.5 kK, which are weak and overlaid in the corresponding absorption spectrum. Each of these two components arises from a transition which is electric dipole forbidden but magnetic dipole allowed, with A_2 direct-product symmetry in C_{2v} . The 3T_1 tetrahedral ground term of nickel(II) gives the C_{2v} components, 3A_2 , 3B_1 , and 3B_2 , whereas the lowest excited term, 3T_2 , gives components with ${}^{3}A_{1}$, ${}^{3}B_{1}$, and ${}^{3}B_{2}$ symmetry. Accordingly the magnetic-dipole transitions represent ${}^3B_1 \longleftrightarrow {}^3B_2$ intercombinations, the one at 9.0 kK from the 3T_2 manifold and the other at 18.5 kK from the ${}^3T_1(P)$ tetrahedral term. The electronic ground state of [Ni(sp)Cl₂] may have ${}^{3}B_{2}$ and not ${}^{3}B_{1}$ symmetry, as shown in the approximate energy-level diagram of this complex (Figure 3), and in that event the excited ${}^{3}B_{2}$ states are relabelled

With these assignments the three C_{2v} components of the ${}^{3}T_{1}(P)$ nickel(II) term in [Ni(sp)Cl₂] are located at 16.5, 18.5, and 20.5 kK (Figure 2). The c.d. spectrum shows that rotational strengths of the individual components, R_i , follow the sum-rule, $\Sigma_i R_i = 0$, as expected for the effect of a weak dissymmetric field, due to the carbon and hydrogen atoms of (-)-spartein, superimposed upon the relatively strong C_{2v} perturbation. At lower energies the doublet observed in both the absorption and the c.d. spectrum at 12.0 and 11.2 kk is ascribed to transitions to the mixed C_{2v} states, ${}^{3}A_{2}$ and ${}^{1}A_{2}$, arising from the tetrahedral nickel(II) terms, 3A2 and ¹E, respectively. Two of the components of the tetrahedral 3T_2 nickel(II) term are identified by the c.d. bands at 9.0 and 7.2 kk, and the third may be the weak band at 5 kk (ε 6) observed with [Ni(sp)Cl₂] in a KBr matrix (Figure 2). Alternatively the 5 kk band may represent the highest-energy component of the ground tetrahedral ${}^{3}T_{1}(F)$ term. A large splitting of that term (ca. 5 kK) is required to accommodate the observed spectra of the [Ni(sp)X₂] complexes (Figures 2 and 3) to values of the ligand field (Dq) and electron-repulsion (B) parameters found for analogous pseudo-tetrahedral nickel(II) complexes (Table 2). A C_{2v} splitting of the ground ${}^3T_1(F)$ term of a similar magnitude (4.8 kK) has been obtained 13 from a polarised single-crystal study of the analogous distorted tetrahedral complex, [Ni(Ph₃P)₂Cl₂].

The polarised single-crystal spectrum of the isostruc $tural\ cobalt (II)\ complex,\ [Co(Ph_3P)_2Cl_2],\ also\ reveals$ large C_{2v} splittings of the tetrahedral T_1 d-electron terms of the metal ion.¹⁴ The three C_{2v} components of the 4T_1 (P) term, centred on 15.4 kK, are split by 3.5 kK, and the corresponding components of the ${}^{4}T_{1}$ (F) term

B. N. Figgis and J. Lewis, *Progr. Inorg. Chem.*, 1964, 6, 37.
 T. Malinovskii, *Kristallografiya*, 1957, 2, 734.
 L. Sacconi, I. Bertini, and F. Mani, *Inorg. Chem.*, 1967, 6,

¹⁸ R. J. Fereday, B. J. Hathaway, and R. J. Dudley, J. Chem. Soc. (A), 1970, 571.

228 J.C.S. Dalton

at 8.4 kK are split by 4.3 kK.14 In the latter case the middle component at 8.0 kK (ε 25) is weak 14 relative to its neighbours at 6.4 kK (ϵ 110) and 10.7 kK (ϵ 60), and in the corresponding ${}^{4}T_{1}$ (F) manifold of the [Co(sp)X₂] complexes, where the splitting is smaller (3.0—3.5 kK). only two components are observed in the solution spectra (Figure 1), the middle component presumably

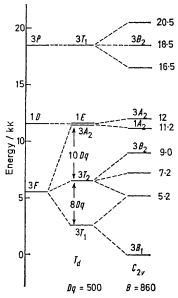


FIGURE 3 Partial energy level diagram for [Ni(sp)Cl₂]

being overlaid. The splitting of the 4T_1 (P) term of the [Co(sp)X₂] complexes in the 15—18 kK region (Figure 1) is similarly smaller (2 kK) than in [Co(Ph₃P)₂Cl₂]. Again the central of the three components has the smallest intensity (Figure 1), as also in the case 14 of $[Co(Ph_3P)_2Cl_2].$

TABLE 2 Values of Dq and B for pseudotetrahedral nickel(II) complexes

	$Dq/\mathrm{cm^{-1}}$	B/cm^{-1}	Ref.
Ni(sp)CI ₂	500	860	a
Ni(sp)Br ₂	470	830	а
$Ni(Ph_3P)_2Cl_2$	489	824	\boldsymbol{b}
Ni(NCO) ₄ 2-	460	810	c
NiCl,2-	345	734	d

^a This work. ^b R. J. Fereday, B. J. Hathaway, and R. J. Dudley, J. Chem. Soc. (A), 1970, 571. ^e D. Forster and D. M. L. Goodgame, J. Chem. Soc., 1964, 2790. ^d G. B. Smith, C. H. Liu, and T. R. Griffith, J. Amer. Chem. Soc., 1964, 86, 4796.

Values of Dq and B consistent with the observed solution spectra of the [Co(sp)X₂] complexes are recorded in Table 3, together with the values of the spin-orbit coupling parameter, λ , derived from the measured magnetic moments after correction for temperature independent paramagnetism. The Dq and Bvalues place the 4T_2 (F) term some 5 kK above the ground ${}^{4}A_{2}$ state of the [Co(sp)X₂] complexes. In a KBr matrix

these complexes give a weak and broad absorption in the 4.5-5.55 kk region which is not present in the i.r. spectrum of the corresponding [Zn(sp)X₂] complex. The Dq and B values obtained in both the $[Ni(sp)X_2]$ (Table 2) and the $[Co(sp)X_2]$ series (Table 3) are rather large, exceeding in the cobalt(II) case the values proposed ¹⁶ for the tetrahedral [CoN₄] chromophore in [Co-(benzimidazole)₂]. However, the π -bonding effects of an aliphatic diamine ligand are small, and possibly the

TABLE 3 Values of Dq and B for pseudotetrahedral cobalt(II) complexes

Dq/cm^{-1}	B/cm^{-1}	— λ/cm ⁻¹	Ref.
510	690	164	a
490	690	175	a
470	680	174	a
510	720	128	\boldsymbol{b}
510	580		с
43 0	700		d
312	710	145	e
	510 490 470 510 510 430	510 690 490 690 470 680 510 720 510 580 430 700	510 690 164 490 690 175 470 680 174 510 720 128 510 580 430 700

^a This work. ^b M. Goodgame and F. A. Cotton, J. Amer. Chem. Soc., 1962, **84**, 1543. ^c C. Simo and S. Holt, Inorg. Chem., 1968, **7**, 2655. ^d J. Ferguson, J. Chem. Phys., 1959, **32**, 528. ^e F. A. Cotton, D. M. L. Goodgame, and M. Goodgame, Amer. Chem. Chem. Chem. 20, 1964, **199**, 4600. game, J. Amer. Chem. Soc., 1961, 83, 4690.

steric constraints of the (-)-spartein molecule give rise to a shortened metal-nitrogen bond distance in the complex, producing a larger ligand field.

EXPERIMENTAL

Metal halide (1 mmol) was dried over silica gel under vacuum for a few hours and quickly dissolved in boiling absolute ethanol (50 ml) to which triethylorthoformate (10 ml) had been added. The solution was boiled until the metal ion was completely dehydrated (5 min for the chlorides and up to 20 min for the bromides and iodides) when a solution of (-)-spartein (1.5 mmol) in absolute ethanol (50 ml) and triethylorthoformate (10 ml) was added. The mixture was boiled for a few minutes and filtered. It was sometimes necessary to extract the large amount of insoluble material produced with ethanol to obtain the co-precipitated complex. The ethanol solutions were cooled to 0 °C when the complexes crystallised. They were re-crystallised from boiling ethanol. A similar procedure was adopted with FeCl₂ and MnCl₂, oils being produced in each case. Complete removal of the ethanol and scratching caused these to solidify. No complexes could be obtained using the nitrate, perchlorate, or thiocyanate of nickel(II) nor could chromium-(III) chloride be made to form a complex with (—)-spartein.

Absorption spectra were obtained on a Cary 17 spectrophotometer in EtOH or CHCl₃ solutions or in a KBr disc. C.d. spectra were obtained on a Jouan II instrument and on an instrument designed in this laboratory.17 Magnetic moments were by the n.m.r. method,18 the complex being dissolved in a CHCl3-CDCl3 mixture and using CHCl3 as the external reference.

We thank the S.R.C. for a Fellowship (to R. D. P.).

[2/2018 Received, 25th August, 1972]

C. Simo and S. Holt, Inorg. Chem., 1968, 7, 2655.
 B. N. Figgis, Trans. Faraday Soc., 1960, 56, 1553.
 M. Goodgame and F. A. Cotton, J. Amer. Chem. Soc., 1962, 84, 1543,

¹⁷ R. Grinter, M. J. Harding, and S. F. Mason, J. Chem. Soc. (A), 1970, 667.

18 D. F. Evans, J. Chem. Soc., 1959, 2003.