Multinuclear NMR Spectroscopy of the Tetrahedral Uranium(1V) Complex U(BH₃CH₃)₄

E. Gamp,^{††} R. Shinomoto,^{†§} N. Edelstein,*[†] and B. R. McGarvey*^{*}

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The temperature dependence of the ¹H, ¹¹B, and ¹³C NMR spectra of T_d U(BH₃CH₃)₄ in solution is reported. The paramagnetic shifts are interpreted as originating purely from spin delocalization mechanisms with no contribution from the metal-orbital dipolar interaction. It is shown that the temperature dependence of both IH shifts (bridging and terminal protons) is identical with that calculated from a polarization theory which assumes the shift is proportional to the average value of electron spin in the inner 5f orbitals. The proportionality constant is -5.64 MHz for the bridging protons and -0.59 MHz for the terminal protons. The temperature dependences of ¹¹B and ¹³C shifts are found to depart significantly from that predicted by the polarization theory with the largest deviations shown by the ¹¹B shifts. It is shown how those deviations can be accounted for by postulating a second spin delocalization through direct covalency involving molecular orbitals formed from the uranium **5f** orbitals and ligand **s** and p orbitals.

Introduction

The spectroscopy of f-block tetraborohydrides is of great interest because of the high molecular symmetry of the $M(BH_3-R)_4$ unit $(M = Hf, Zr, Th, Pa, U, Np, Pu; R = H, CH₃).¹$ Only pure solid Th; Pa-, and $U(BH_4)_4$ are of lower than cubic symmetry, having a polymeric structure. All the other compounds contain molecular units of T_d symmetry with each of the four tetrahedrally coordinated borohydride ligands bound to the metal via three hydrogen bridges with the fourth hydrogen or the methyl group pointing out along the threefold axis of the complex.

The optical spectra,2 paramagnetic susceptibility, and *EPR* spectra of uranium and neptunium tetraborohydrides have recently been analyzed in terms of a parametrized Hamiltonian, with the full $5f^2(5f^3)$ basis.³ Within this basis, optical and magnetic properties could not be satisfactorily explained by the same set of Hamiltonian parameters without the introduction of orbital reduction factors. Furthermore, it was shown that *J* mixing by the crystal field in these compounds cannot be neglected.

Paramagnetic shifts in the NMR spectrum of the coordinated ligands can be a powerful tool to examine interactions between the metal f electrons and the ligands.⁴ Here again, a high molecular symmetry is important in that pseudocontact (dipolar) shifts, which are sometimes hard to separate from the spin delocalization shifts, are zero in T_d symmetry.

The only NMR spectra of $U(\bar{IV})$ compounds with cubic molecular symmetry reported to date are those of $U(\text{cp})_4$ (cp = cyclopentadienide),⁵ U(NCS)₈,⁶ and U(BH₄)₄.⁷ Although the last is a polymer in the solid state, tetrahedral molecular units were found in solution with rapid exchange on the 'H NMR time scale between terminal and bridging hydrogens. Replacement of the terminal hydrogen by a methyl group prevents this exchange and at the same time increases the delocalization range for the spin density originating at the central ion.

In this paper we report paramagnetic shifts from ${}^{1}H, {}^{11}B,$ and ¹³C NMR spectra of $U(BH_3CH_3)_4$ and interpret them in terms of spin delocalization from the central ion onto the ligands.

Theory of the Paramagnetic Shift

The theoretical parts of papers on paramagnetic NMR shifts contain a confusing variety of definitions and sign conventions.^{4,6-10} We will therefore give a short survey of the necessary equations that were used in the interpretation of the paramagnetic shifts presented in this paper.

¹¹University of Windsor.

The paramagnetic shift, $\Delta H/H$, of the signal under consideration is measured relative to a diamagnetic reference of similar structure (in the present case $Th(BH_3CH_3)_4$. The shift is given in ppm, and we define shifts to higher field as positive (the opposite direction has been used in the past, especially in ¹H and ¹³C NMR). Kurland and McGarvey¹⁰ have shown that the paramagnetic shift can be calculated from the equation

$$
\Delta H/H_i = (kTq)^{-1} \sum_{\Gamma n, \Gamma m} e^{-i\Gamma/kT} \langle \Gamma n | \mu_i | \Gamma m \rangle \langle \Gamma m | A_{N_i} / g_N \beta_N | \Gamma n \rangle - q^{-1} \sum_{\Gamma n, \Gamma' m(\Gamma' = \Gamma)} Q_{\Gamma \Gamma'} \langle \Gamma n | \mu_i | \Gamma' m \rangle \langle \Gamma' m | A_{N_i} / g_N \beta_N | \Gamma n \rangle
$$
 (1)

in which

$$
q = \sum_{\Gamma} e^{-\epsilon \Gamma / kT} \tag{2}
$$

$$
Q_{\Gamma\Gamma'} = (e^{-\epsilon_{\Gamma}/k} - e^{-\epsilon_{\Gamma}/k}) / (\epsilon_{\Gamma} - \epsilon_{\Gamma'})
$$
\n(3)

$$
\mu_i = -\beta_e (L_i + g_e S_i) \tag{4}
$$

 $A_{N_i}/g_N\beta_N = (8\pi/3)g_e\beta_e\sum_j \delta(r_j) (s_i)_j + g_e\beta_e\sum_j [3(\vec{s}_f\vec{r}_j)(r_i)_j - r_j^2(s_i)_j] + 2\beta_e\sum_j r_j^{-3}(l_i)_j$ (5)

In the above equations the Γ and Γ' indices label energy levels of the system in the absence **of** an applied magnetic field and *n* and *m* label particular degenerate states within a given energy level $\epsilon_{\rm F}$. For the U(BH₃CH₃)₄ system Γ designates a particular *SLJ* state while *n* and *m* designate the J_z components of that state. Equation 1 was derived for a rigid system, and the subcript *i* refers to the direction of the magnetic field in the molecular coordinate system *j.* The solution shift is found by calculating the shift along the three principal axes of the molecular coordinate system and then averaging. S_i is the total electron spin operator, (s_i) is the single electron spin operator for electron *j*, L_i is the total orbital angular momentum operator, $(l_i)_j$ is the orbital angular momentum operator for electron *j*, β_e and β_N are the Bohr and nuclear magnetons, respectively, g_e is the free-electron g value (2.0023), g_N is the

- **(1)** Shinomoto, R.; Gamp, E.; Edelstein, N. M.; Templeton, D. H.; Zalkin, A. *Inorg. Chem.* **1983,** *22,* 2351.
- (2) Bernstein, E. R.; Keiderling, T. A. *J. Chem. Phys.* **1973,** *59,* 2'105. (3) (a) Rajnak, **K.;** Gamp, E.; Shinomoto, R.; Edelstein, N. *J. Chem. Phys.* **1984,80,** 5942. (b) Rajnak, R.; Banks, R. H.; Gamp, E.; Edelstein, N.
- *Ibid.* **1984,** *80,* 5951. (c) Gamp, E.; Edelstein, N. *Ibid.* **1984.80,** 5963.
- (4) See e.g. in: *NMR ofParamagnetic Molecules;* LaMar, G. N., Hor-rocks, W. D., Jr., Holm, R. H. Academic: New York, 1973. *(5)* von Ammon, R.; Kanellakopulos, B.; Fischer, R. D. *Chem. Phys. Lett.*
- **1968**, 2, 513. *(6)* Folcher, G.; Marquet-Ellis, H.; Rigny, P.; Soulie, E.; Goodman, G. *J. Inorg. Nucl. Chem.* **1976, 38,** 147.
- (7) Folcher, G.; Lambard, J.; Marquet-Ellis, H. *Nouv. J. Chim.* **1980**, *4*,
- *IJI.* (8) Kiener, C.; Folcher, G.; Langlet, G.; Rigny, P.; Virlet, **J.** *Now. J. Chim.* **1979,** *3,* 99.
- (9) Edelstein, N.; LaMar, G. N.; Mares, F.; Streitwieser, A., Jr. *Chem. Phys. Lett.* **1971, 8,** 399.
- (10) Kurland, R. J.; McGarvey, B. R. *J. Magn. Reson.* **1970,** *2,* 286.

^{&#}x27;Lawrence Berkeley Laboratory and University of California.

^{*}Present address: EMPA, CH-8600 Dubendorf, Switzerland.

Present address: Department of Chemistry, University of Southern California, **Los** Angeles, CA 90089.

nuclear g value, $(r_i)_j$ is the i component of the \vec{r}_i vector between the nucleus and the *j*th electron, and $\delta(r_j)$ is Dirac's δ function. T is temperature in kelvin, and *k* is Boltzmann's constant.

If we approximate the $|\Gamma_m\rangle$ functions by purely f-orbital functions, eq 1 reduces to the familiar dipolar shift equations,¹⁰ which predict the average shift to be proportional to the magnetic susceptibility anisotropy. The molecular unit of $U(BH_3CH_3)_4$, dissolved in toluene, has T_d symmetry and has no such anisotropy. Therefore, the dipolar shift (pseudocontact shift) should be zero and the observed paramagnetic shift is solely due to mechanisms that transfer electron spin into the ligand orbitals. In this paper we shall refer to this shift as the delocalized-spin shift. Most of the literature calls this shift the contact (Fermi) shift, but this is incorrect and misleading when dealing with the paramagnetic shifts of lanthanide and actinide complexes, as it implies that the shift comes only from electron spin delocalized into ligand **s** orbitals. While this is generally (but not always) true for transition-metal complexes (for which the term was invented), the following discussion will show that it is not the case for systems in which the orbital angular momentum **is** not quenched.

Two spin-transfer mechanisms have been proposed to explain NMR, ESR, and ENDOR results from lanthanide and actinide systems. Lewis et al.¹¹ and Reuben and Fiat¹² concluded from ¹⁷O NMR shift studies on hydrated lanthanide ions that the shifts were primarily Fermi contact in origin with a negative contact term. Lewis et al. attributed the negative term to a polarization mechanism in which 4f electrons polarized the bonding electrons in a bond formed by the ligand atom donating electrons from its 2s and 2p orbitals into the empty 6s orbital of the lanthanide ion. Watson and Freeman¹³ proposed that a negative spin density in the outer **5s** and 5p electrons of the lanthanide ion is produced by a polarization interaction with the unpaired **4f** electrons and that this negative spin density is transferred to adjacent ligand atoms by covalent and overlap interactions between these **5s** and 5p orbitals and the **2s** and 2p orbitals of the ligand.

The exact distinction between overlap and covalent transfer of spin has been discussed by McGarvey.¹⁴ Basically this polarization mechanism transfers spin from the outer metal **s** and p orbitals to the ligand orbitals and this spin results from a polarization mechanism that favors spin of sign opposite to that in the inner f shell. It was further assumed that effects of spin in ligand p orbitals would average out in solution (ignoring the coupling of electron spin and electron orbital angular momentum), leaving only the isotropic contribution of electron spin in ligand **s** orbitals. Most of the chemical literature on NMR shifts in lanthanide and actinide complexes has assumed the above explanation to be correct and has, therefore, referred to all isotropic shifts (obtained after subtraction of the dipolar shift) as "contact shifts".

Baker¹⁵ has shown that although the polarization mechanism would explain ¹⁹F ENDOR results for Eu^{2+} and Gd^{3+} (f^7 configuration) in CaF₂, it could not account for the results obtained for Tm^2 or Yb^{3+} (f¹³) configuration) in which the isotropic component of the hyperfine interaction was positive rather than negative. He showed that the results for Tm^{2+} and Yb^{3+} could be explained by a direct covalent transfer of the 2s and 2p electrons of the fluoride ion into the 4f shell. McGarvey^{14,16} extended the covalent calculations to Yb^{3+} in octahedral sites and in distorted cubic sites.

The **I9F** NMR shift at room temperature was measured for crystals of CdF₂¹⁷ and CaF₂¹⁸ containing Yb³⁺, and the isotropic shift was found to be upfield in the same direction found for ¹⁷O NMR shifts¹¹ in solution, which had been interpreted to mean a negative isotropic hyperfine interaction. Thus ENDOR and NMR studies in the same system appeared to arrive at contradictory results. McGarvey¹⁹ has shown that the same covalent mechanism used to explain the hyperfine interaction measured by ENDOR could also explain the upfield shift measured by NMR by using eq **1-5.** It was found that the largest term came from the second part of eq 1 and the third term in eq **5.** In other words, the sign of the NMR shift in this case was not determined by the Fermi contact term (which in this case did predict a downfield shift) but rather

- (12) Reuben, J.; Fiat, D. *J. Chem. Phys.* **1969,** *51,* 4909.
- (13) Watson, R. E.; Freeman, A. J. *Phys. Rev.* **1967,** *156,* 251.
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- (14) McGarvey, B. R. J. Chem. Phys. 1976, 65, 955.

(15) Baker, J. M. J. Phys. C 1968, 1, 1670.

(16) McGarvey, B. R. J. Chem. Phys. 1979, 70, 4971.

(17) Mustafa, M. R.; Jones, W. E.; McGarvey, B. R.; Greenblatt, M.; Bank
- **(18)** Booth, R. J.; Mustafa, M. R.; McGarvey, B. R. *Phys. Rev. E Condem. Mutter* **1978,** *17,* 4150.
- (19) McGarvey, B. R. *J. Chem. Phys.* **1976,** *65,* 962.

by the interaction between the nuclear spin and the unquenched orbital angular momentum of electrons in ligand p orbitals. Further, the shift was not determined by a hyperfine matrix element for any ground or, excited state (that could be measured by ESR or ENDOR) but rather by a hyperfine matrix element between different crystal field states of the system. ¹⁹F NMR shift studies on a series of elpasolites²⁰ have shown that the polarization mechanism is dominant in the first half of the lanthanide series but both the covalent and polarization mechanisms must be invoked to explain results for the second half of the series.

About the only satisfactory way to do a theoretical calculation for the polarization mechanism is to do an unrestricted Hartree-Fock calculation on the complete complex to obtain the $|\Gamma_m\rangle$ functions. Baker¹⁵ has suggested that the effect should be proportional to the spin in the f orbitals and therefore one could estimate the matrix elements $\langle \Gamma' m | A_{N_i} / g_N \beta_N | \Gamma n \rangle$ in eq 1 from the spin matrix elements $\langle \Gamma' m | S_i | \Gamma n \rangle$, which are readily calculated from the *SLJJ₁* functions. That is

$$
\langle \Gamma' m | A_{N_i} / g_N \beta_N | \Gamma n \rangle = (K / g_N \beta_N) \langle \Gamma' m | S_i | \Gamma n \rangle \tag{6}
$$

with the proportionality constant *K* being independent of the *SLJ* quantum numbers. The constant *K* has a negatve value for the polarization mechanism and units of energy. Baker found, in the case of **Eu2+** and Gd^{3+} in CaF_2 , that different *K* values were required for the directions parallel and perpendicular to the vector connecting the fluoride ion and the rare-earth ion. He found $K_{\parallel} = -3.60$ MHz and $K_{\perp} = -0.93$ MHz for Gd³⁺ and $K_{\parallel} = -3.49$ MHz and $K_{\perp} = -1.60$ MHz for Eu²⁺.

For the isotropic shift we can assume an average value for *K* and calculate the shift for the *z* direction. Putting eq 6 into eq 1, we obtain

$$
\Delta H/H = (K/g_{\rm N}\beta_{\rm N})\langle S_z\rangle/H \tag{7}
$$

where $\langle S_z \rangle$ (the average total electronic spin) is found from the equation

$$
\langle S_z \rangle / H = -(\beta_e / kTq) \sum_{\Gamma n, \Gamma m} e^{-\epsilon_{\Gamma}/kT} \langle \Gamma n | L_z + g_e S_z | \Gamma m \rangle \langle \Gamma m | S_z | \Gamma n \rangle -
$$

$$
(\beta_e / q) \sum_{\Gamma n, \Gamma' m} Q_{\Gamma \Gamma'} \langle \Gamma n | L_z + g_e S_z | \Gamma' m \rangle \langle \Gamma' m | S_z | \Gamma n \rangle
$$
 (8)

Equation **7** is the usual equation put forward as the contact shift equation⁴ except the parameter *K* is replaced by *A*, which is assumed to be the isotropic hyperfine constant.

Lewis et al.¹¹ have pointed out that since the lowest energy states belong to the same *J* manifold (assuming no *J* mixing by the crystal field), the operators in eq 8 can be replaced by equivalent *J* operators. That is, $L_z + g_e S_z = g_J J_z$ and $S_z = (g_J - 1) J_z$, where *g* is the Lande *g* factor. In this case the sign of the shift is determined by the sign of $g_J(1)$ $-g_J$)K. For ions with less than seven f electrons g_J is less than 1 and therefore a negagive *K* will give a negative or downfield shift. Ions with more than seven f electrons have g_J greater than 1, and therefore eq 8 will predict a positive or upfield shift.

If the crystal field states have energies of the same magnitude as *kT* or larger than kT , eq 7 and 8 will not predict a simple T^{-1} (Curie law) behavior for $\Delta H/H$. It will approach such behavior only for temperatures in which $kT \gg \epsilon_{\rm F}$ of the crystal field states. The behavior at very low temperatures depends on the nature of the ground state. If the groundstate matrix elements $\langle 0m|S_z|0n\rangle$ are not zero, the temperature dependence becomes T^{-1} at low temperatures and $\Delta H/H$ becomes very large at low temperatures. If $\langle 0m|\dot{S}_z|0n\rangle = 0$ for the ground state but corresponding matrix elements between the ground state and excited state are not zero, *AH/H* approaches a constant value at low temperatures. This is the situation for $U(BH_3CH_3)_4$. Finally if all matrix elements involving the ground state are zero, $\Delta H/H$ approaches zero at very low temperatures.

In actinide compounds, especially at the beginning of the series, spin-orbit coupling and crystal field energies are of the same magnitude and extensive f mixing takes place so that calculations within the ground-state multiplet alone can no longer explain finer effects. It has been shown, also, that orbital reduction factors are necessary to explain experimental results satisfactorily.^{3,9,21} For this reason an empirical reduction factor k_z has been introduced in which the operator L_z has been replaced by $k_z L_z$ in any calculation. The wave functions $|\Gamma_n\rangle$ are linear combinations within the familiar $|\gamma SLJJ_z\rangle$ basis.²² The computational effort grows exceedingly if all free ion terms of the **f"** configuration are included in the calculation. We give, therefore, easily programmable equations for all necessary matrix elements in the Appendix.

The problem is more complex for the covalent mechanism, and no simple rules can be formulated that will predict the sign of the shift. In

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- (20) Reuveni, A.; McGarvey, B. R. J. Magn. Reson. 1979, 34, 181.
(21) Gamp, E.; Edelstein, N.; Khan Malek, C.; Hubert, S.; Genet, M. J.
Chem. Phys. 1983, 79, 2023.
- (22) Wybourne, 8. *G. Spectroscopic Properties* of *Rare Earths;* Wiley: New York, 1965.

^(1 1) Lewis, W. B.; Jackson, J. **A.;** Lemons, J. F.; Taube, H. *J. Chem. Phys.* **1962,** *36,* 694.

view is along an **S4** axis.

Figure 2. T^{-1} dependence of the paramagnetic shifts in $U(BH_3CH_3)_4$ referred to $Th(BH_3CH_3)_4$. All shifts are to lower field and given in ppm. The solid lines were calculated from eq 7and 8 with $k_z = 0.85$, $\Delta_1 = 215$ cm-I, *K* values given in Table **11,** and crystal field energies and eigenvectors of the optical analysis.^{3a} The dashed line for B is a combined polarization plus covalent shift theoretical curve.

most of the literature it has been argued that a covalent transfer leads to a positive Fermi contact term and therefore to a shift opposite in sign to that predicted by the polarization mechanism. This argument was based on two false assumptions: (I) the electrons in ligand p orbitals will not contribute to isotropic solution shifts because this is a dipolar interaction that averages to zero and (2) the s electrons that interact through the Fermi contact term give an isotropic and positive interaction. While both assumptions are correct for systems in which the orbital angular momentum is quenched to first order, they are not true when angular momentum is not quenched (as in the case for all rare-earth ions except the **8S** state of the **f'** configuration). The p electrons contribute to the shift through the second and third terms in *eq* 5. For spin-only systems, the second term gives a traceless shift matrix that will average to zero in solution and the third term is zero. For complexes with orbital angular momentum the third term is neither zero nor traceless and therefore contributes to the isotropic shift. Even the second term gives a nontraceless shift matrix in some ground states. Thus the first assumption is incorrect and it is wrong to call the isotropic shift a contact shift in such systems.

The second assumption is sometimes also wrong. McGarvey^{14,16} has

Table I. Lowest Calculated Energy Levels (cm⁻¹) from the Optical Analysis of $U(BD_4)_4/Hf(BD_4)_4$ and Corresponding Eigenvectors^{3a}

state	$E_{\rm calcd}$	eigenvector ^a
Е		94 ${}^{3}H_{4}$ + 3 ${}^{1}G_{4}$
T_1	215 ^b	$76 \text{ }^{3}\text{H}_{4} + 11 \text{ }^{3}\text{H}_{5} + 7 \text{ }^{1}\text{G}_{4}$
T ₂	531	$78 \text{ }^{3}H_{4} + 11 \text{ }^{1}G_{4} + 6 \text{ }^{3}F_{3}$
A_1	2036	63 ${}^{3}H_{4}$ + 27 ${}^{1}G_{4}$
Е	2750	$57 \text{ } ^3\text{F}_2 + 31 \text{ } ^3\text{H}_4 + 9 \text{ } ^1\text{D}_2$
T,	3562	58 ${}^3\text{F}_2$ + 20 ${}^3\text{H}_3$ + 10 ${}^1\text{D}_2$

include 90% of the state. $\frac{b}{b}$ From analysis of the susceptibility. "Percent of *SU* state. Enough components are given, at least 2, to

shown that the Fermi contact term leads to a positive and isotropic hyperfine interaction for the Γ_7 ground state of Yb³⁺ in cubic symmetry in CaF₂, but the ¹⁹F hyperfine parameter for the Γ_6 ground state of Yb³⁺ in the octahedral symmetry of KMgF, is highly anisotropic and gives a negative value for the average hyperfine interaction. The calculaion of the NMR shift¹⁹ for ¹⁹F of Yb³⁺ in CaF₂ gave an anisotropic shift for the Fermi contact contribution even though the ground state itself had an isotropic Fermi term. Thus the second assumption can also be wrong.

We have used the same methods¹⁹ used to calculate the ¹⁹F NMR shift of Yb^{3+} in CaF₂ to estimate the covalent contribution to the paramagnetic shift in $U(BH_3CH_3)_4$. To keep the calculation simple, we have assumed a pure 'H, state for the 5f2 configuration with no *J* mixing. Rajnak et al.³ gives the ground state for $U(BH_4)_4$ as E with the excited states in ascending energy as being T_1 , T_2 , and A_1 . We have also left out the A_i state from the calculation as it is much higher in energy than the T_1 and T_2 states and has no matrix elements connecting it to the E ground state. The calculation was done only for ligand atoms along a \bar{C}_3 axis of the tetrahedron and is therefore applicable only to the B and C atoms of $U(BH_3CH_3)_4$. The resulting equation is

$$
\Delta H/H = [g_{J}\beta_{e}/27kTqg_{N}\beta_{N}][e^{-\Delta_{1}/kT}[1.13939a_{\sigma}^{2}A_{2s} + (-5.24848b_{\sigma}^{2} - 2.01818b_{\sigma}^{2} + 15.45405b_{\sigma}b_{\tau})A_{2p}] +e^{-\Delta_{2}/kT}[5.88745a_{\sigma}^{2}A_{2s} + (1.25108b_{\sigma}^{2} + 19.84416b_{\sigma}^{2} +62.43548b_{\sigma}b_{\sigma})A_{2p}] + Q_{01}[24.84040a_{\sigma}^{2}A_{2s} + (-0.50101b_{\sigma}^{2} -29.44242b_{\sigma}^{2} + 208.16209b_{\sigma}b_{\sigma})A_{2p}] +Q_{02}[6.21299a_{\sigma}^{2}A_{2s} + (-9.13593b_{\sigma}^{2} - 9.53824b_{\sigma}^{2} +111.31923b_{\sigma}b_{\sigma})A_{2p}] - Q_{12}[2.90505a_{\sigma}^{2}A_{2s} + (16.61010b_{\sigma}^{2} +41.50909b_{\sigma}^{2} - 64.50323b_{\sigma}b_{\sigma})A_{2p}] \qquad (9)
$$

where

$$
Q_{01} = (1 - e^{-\Delta_1/k} T) k T / \Delta_1
$$
 (10)

$$
Q_{02} = (1 - e^{-\Delta_2/k} T) k T / \Delta_2
$$
 (11)

$$
Q_{12} = (e^{-\Delta_1/kT} - e^{-\Delta_2/kT})kT/(\Delta_1 - \Delta_2)
$$
 (12)

$$
q = 2 + 3e^{-\Delta_1/k} + 3e^{-\Delta_2/k} + e^{-\Delta_3/k} \tag{13}
$$

$$
\Delta_1 = \epsilon(T_1) - \epsilon(E) \tag{14}
$$

$$
\Delta_2 = \epsilon(T_2) - \epsilon(E) \tag{15}
$$

$$
\Delta_3 = \epsilon(A_1) - \epsilon(E) \tag{16}
$$

$$
A_{2s} = (8\pi/3)(g_e \beta_e g_N \beta_N) |2s(0)|^2
$$
 (17)

$$
A_{2p} = \frac{2}{5} (g_e \beta_e g_N \beta_N) \langle r^{-3} \rangle_{2p} \tag{18}
$$

The a_{σ} term is the molecular orbital coefficient for the ligand 2s orbital mixing with the 5f₀ metal orbital, b_{σ} is for mixing of 2p_{σ} with 5f₀, and b_{τ} is for mixing of 2p_{± 1} with 5f_{±1}, Baker¹⁵ and McGarvey¹⁴ found for F⁻ that $b_{\sigma} \approx b_{\tau}$ as was found earlier for transition-metal fluorides.²³ If we assume this to be the case for B and C in $U(BH_3CH_3)_4$, the above equation predicts an upfield shift for the energies reported by Rajnak) et al.

⁽²³⁾ Hall, T. **P.** P.; Hayes, W.; Stevenson, R. W. H.; Wilkens, J. J. *Chem. Phys.* **1963,** *38,* **1977; 1963,** *39,* **35.**

Figure 3. Temperature dependence of the total average spin $\langle S_z \rangle /H$ from eq 8: (a) $k_z = 1.00$, $\Delta_1 = 215$ cm⁻¹, complete set of eigenvalues and eigenvectors from ref 3a; (b) $k_z = 1.00$, $\Delta_1 = 370$ cm⁻¹, complete set of eigenvalues and eigenvectors from ref 3a; (c) $k_z = 0.85$, $\Delta_1 = 215$ cm⁻¹, complete set of eigenvalues and eigenvectors from ref 3a; (d) $k_z = 1.00$, $\Delta_1 = 215$ cm⁻¹, pure ³H₄ functions only; (e) $k_z = 1.00$, $\Delta_1 = 370$ cm⁻¹, pure ${}^{3}H_4$ functions only.

Results and Discussion

The T_d molecular symmetry of the U(BH₃CH₃)₄ unit is shown in Figure 1. Each ligand is bound to the metal center via three bridging hydrogens. The boron atoms lie at the tetrahedral corners, and the methyl groups attached to them point out of the complex along the threefold axes of the molecular T_d unit. This leads to only four magnetically nonequivalent nuclei: bridging hydrogens (H_b) , terminal hydrogens (H_t) , boron, and carbon. Their paramagnetic shifts (referenced to the isostructural, but diamagnetic Th $(BH_3CH_3)_4$ ¹) are shown in Figure 2 as a function of T^{-1} .

No splittings from spin-spin coupling $(^1H-^{11}B$ or $^1H-^{13}C$) have been observed due to the rather large line widths. There is no indication of low-symmetry components in solution, e.g. solvent adducts (which have a markedly different NMR spectrum)²⁴ or dimerization. The shifts can therefore be considered as having no dipolar (pseudocontact) shift. All shifts are to low fields, and this indicates that the polarization mechanism is dominant in each case. The $T⁻¹$ dependences shows significant curvatures and nonzero extrapolated intercepts for $T^{-1} = 0$, indicating that a first-order treatment and the assumption that the splitting of the free ion ground term (${}^{3}H_{4}$) by the crystal field is smaller than kT are not correct in the present case.

This is confirmed by the optical and magnetic analyses, which have shown that it is possible to explain the optical spectrum of $U(BD_4)_4/Hf(BD_4)_4$ and the susceptibility of $U(BH_3CH_3)_4$ with the same set of parameters only if the full basis set of the $f²$ configuration is used and an orbital reduction factor included.^{3a} Table **I** shows the lowest levels and the main components of the corresponding eigenvectors in the *ISWJ,)* basis from the analysis. The lowest T_1 state has been calculated to be at 370 cm⁻¹ in the optical analysis of $U(BD_4)_4$ diluted in $Hf(BD_4)_4$. However, the analysis of the susceptibility of $U(BH_3CH_3)_4$ is especially sensitive to the position of the thermally populated lower levels and allowed a more reliable determination of the T_1 energy at 215 cm⁻¹. The discrepancy is also plausible in view of the slightly different optical spectra in solution of $U(BH_3CH_3)_4$ and $U(BH_4)_4$, where shifts of up to 250 cm-l occur. The introduction of an isotropic orbital reduction factor $k = 0.85$ has also been necessary to explain the susceptibility and account for the reduction of about 10% of the free ion spin-orbit coupling parameter ζ in the complex.

Since the polarization mechanism appears to be dominant, we will first analyze the results in terms of eq 7 and 8. $\langle S_z \rangle / H$ was

Table II. Best-Fit Values of Hyperfine Parameter *K*

	H_b	H,	в	C		
	$k_2 = 1.00$, $\Delta_1 = 215$ cm ⁻¹ , Complete Calculation					
K, MHz -4.069 -0.4245 -0.788 -0.451						
std dev, ppm 0.44 0.21 9.55				2.06		
$k_2 = 0.85$, $\Delta_1 = 215$ cm ⁻¹ , Complete Calculation						
K, MHz $-5.640 -0.588 -1.093 -0.625$						
std dev, ppm 0.43 0.18 9.76 2.21						
	$k_z = 1.00$, $\Delta_1 = 370$ cm ⁻¹ , Complete Calculation					
K, MHz $-4.526 -0.472 -0.856 -0.498$						
std dev, ppm 6.97 0.91 5.84 1.13						
$k_7 = 1.00$, $\Delta_1 = 215$ cm ⁻¹ , ³ H ₄ Functions Only						
K, MHz $-3.520 -0.367 -0.673 -0.389$						
std dev, ppm 3.40 0.54 7.38 0.52						
$k_7 = 1.00$, $\Delta_1 = 370$ cm ⁻¹ , ³ H ₄ Functions Only						
K, MHz $-3.915 -0.408 -0.732 -0.430$						
std dev, ppm 9.83 1.21 3.88				2.56		

calculated from all energies and complete eigenvectors as given in ref 3a up to 16000 cm⁻¹ with $k_z = 1.00$ and with both 370 and 215 cm⁻¹ for the lowest T_1 state. The calculation was repeated for $k_z = 0.85$ and the T₁ state at 215 cm⁻¹. For comparison purposes, the calculation was also done by assuming a pure ${}^{3}H_4$ state with no *J* mixing and ignoring other excited states. In this case *k,* was taken to be unity. The results are plotted in Figure 3 vs. T^{-1} . The non-Curie behavior is quite apparent as well as the pronounced dependence on the energy chosen for the lowest T_i state. The effect of changing k_z is mainly to change the magnitude of $\langle S_z \rangle / H$ rather than the functional dependence on temperature. Using just the pure ${}^{3}H_{4}$ functions gives larger values of $\langle S_z \rangle / H$, but the temperature dependence is similar to that of the complete calculation.

For each $\langle S_z \rangle / H$ function a best fit between theory and experiment was obtained by varying the value of the parameter K in eq 7. The best-fit values of *K* and the standard deviation for each case are given in Table 11. Excellent fits were obtained for both the bridging protons (H_b) and terminal protons (H_t) for 215 cm⁻¹ by using the full basis set andd $k_z = 0.85$ or 1.00. The solid lines in Figure 2 are plots of fitted curves for $k_z = 0.85$. The fit for the ¹³C shifts is less satisfactory and for the ¹¹B shifts is very poor.

The good fit between *eq* **7** and the experimental 'H shifts with the same eigenvectors and energies that have been used to fit the optical spectra of $U(BH_4)_4$ and magnetic susceptibility of $U(B H_3CH_3$)₄ is strong support for the polarization mechanism being the only important spin-transfer mechanism for both ${}^{1}H$ nuclei. The very poor fit for $11B$ means either that another mechanism for spin transfer is also important for boron or that the chemical shift for ^{11}B is not the same in both $U(BH_3CH_3)_4$ and Th(B- H_3CH_3)₄ since an improved fit could be attained by not forcing the data to extrapolate to zero when *T* becomes infinite.

The polarization mechanism predicts the same functional dependence on temperature for all nuclei with only the K parameter being different. If this were true, we could write the following
equation for each of our four shifts
 $(\Delta H/H)_j = K_j f(T) + A_j$ (19) equation for each of our four shifts

$$
(\Delta H/H)_j = K_j f(T) + A_j \tag{19}
$$

where the subscript *j* denotes a particular nuclear shift, $f(T)$ is the common temperature dependence function, and *A,* is the chemical shift in the absence of the paramagnetic shift. All *AJ's* are 0 if our assumption about $U(BH_3CH_3)_4$ and $Th(BH_3CH_3)_4$ having the same chemical shift is valid. Plotting shift 1 vs. shift 2 would yield a straight line whose equation is

$$
(\Delta H/H)_1 = (K_1/K_2)(\Delta H/H)_2 + [A_1 - (K_1/K_2)A] \tag{20}
$$

In Figure 4 the H_t , C, and B shifts are plotted vs. the H_b shifts. For C and B the appropriate H_b shifts were interpolated or extrapolated from the H_b vs. T^{-1} plot. The data were fitted by a least-squares procedure against a straight line. The fitted slopes and intercepts are given in Table 111, and the fitted straight lines

Figure 4. $\Delta H/H$ for C, B, and H_t plotted as a function of $\Delta H/H$ for H_b. The solid lines are least-squares-fitted lines whose slopes and intercepts are given in Table **111.**

are shown in Figure 4 as solid lines. In each case the fit is good, but it is difficult to detect curvature over the small temperature interval of the measurements and it is of the intercept that is more informative. The intercept near zero for the H_t vs. H_b plot is a clear indication that the functional temperature dependence is nearly the same for both ¹H shifts. It is difficult to attribute the large intercepts for both ^{11}B and ^{13}C shifts as due to differences in chemical shifts, and therefore, we take these large shifts as evidence that an additional spin-transfer mechanism is present for these two nuclei that has a different functional dependence on temperature.

To see if inclusion of the covalent mechanism for spin transfer might better explain the temperature behavior of the ¹¹B shift, we have fitted the experimental shifts to a sum of eq 7 and eq 9 assuming $b_{\sigma} = b_{\tau} = b$. Since the covalent equations assumed pure ${}^{3}H_4$ functions, we have used the equivalent calculations for the polarization shift. The fitted equation is plotted in Figure 2 as a dashed curve. The standard deviation is 0.62 ppm. The fitted parameters were $K = -2.76 \text{ MHz}, a_{\sigma}^2 A_{2s} \approx 0$, and $b^2 A_{2p} = 1.26$ MHz. With $A_{2p} = 53.1 \text{ MHz}^{25}$ for ¹¹B, the value of the molecular orbital coefficient is $b^2 = 0.024$, which would mean a 2% covalent transfer to the boron p orbitals or an 8% reduction in the f orbitals. This is a reasonable number considering the 15% reduction in *k,.* It would appear reasonable to ascribe the different temperature dependence in 11 B and 13 C to the presence of a covalent contribution (of sign opposite to that for the polarization shift) to the paramagnetic shift.

The fitted values of K and $b²$ given above cannot be taken as reliable due to the assumption in the calculation of pure ${}^{3}H_{4}$ functions. The data in Table I1 show that these functions gave poorer agreement for the 'H shifts. It is clear, however, that the actual K values for both ^{11}B and ^{13}C are larger than the fitted values of Table II. Also the actual value of b^2 is probably of the same order of magnitude as found above.

It is of interest to compare our values of *K* to those reported by Baker¹⁵ for ¹⁹F-Gd³⁺ and $-Eu^{2+}$. Scaled to ¹H, his average values would be -1.93 MHz for Gd³⁺ and -2.23 MHz for Eu²⁺. Our values for the bridging proton ere only twice as large in magnitude. Scaled down to ¹¹B, the *K* values become –0.62 MHz for Gd^{3+} and -0.81 MHz for Eu^{2+} . Our computations show that *K* in our system is about 3-4 times larger in magnitude.

The contact shifts of H_b of approximately -150 ppm at room temperature are the largest reported to date in U(IV) compounds except for $U(BH_4)_4$, where they are approximately the same.⁷ Reports of ¹¹B shifts are rare. In (C_5H_5) , UBH₄, the delocalization shift contribution for $11B$ has been estimaed to be -240 ppm and that of the bridging protons to be -61 ppm.²⁶ It is not possible

Table 111. Slope and Intercept for Shifts Plotted vs. Bridging Proton Shifts

nucleus	slope	intercept, ppm	correln factor
в	0.205 ± 0.008	68.2 ± 1.4	0.9994
C	0.360 ± 0.010	14.8 ± 3.7	0.9988
H.	0.113 ± 0.006	-1.6 ± 0.1	0.9999

to explain this inversion in magnitudes without some information about the ground and excited states of the molecule. A different ground state will have a pronounced effect on both $\langle S_z \rangle / H$ and on the covalent shift. A different ground state could have a much smaller covalency contribution for ¹¹B, making the measured shift more negative than is found in $U(BH_3CH_3)_4$. Comparison with other similar σ -bonding ligands is difficult due to lack of experimental results. In $(C_5H_3)_3UBH_3CH_2CH_3$ the ¹H delocalization shifts are -55.3 ± 8 ppm for the bridging protons and -7.3 ± 1.2 ppm for the methylene protons.²⁷ Both of these shifts are about one-third, in magnitude, of the shifts found here for corresponding protons in $U(BH_3CH_3)_4$. The bridging proton shift is, however, nearly the same as that found for the related $(C_5H_5)_3UBH_4$, indicating that both have the same ground state. A shift of -90 ppm has been reported⁶ for ¹³C in U(NCS)₈⁴⁻, which is not much different from the -70 ppm reported here.

Conclusions

The observed contact shifts of the bridging hydrogens are the largest reported to date in U(1V) compounds. The temperature dependence and sign of the 'H shifts (both bridging and terminal) agree very well with what is predicted by a polarization model in which spin is transferred by an overlap between outer 6s and 6p electrons of uranium with ligand orbitals. This spin has been negatively polarized to that of the inner 5f electrons by an interaction between the 5f and 6s, 6p electrons. The agreement between the experimental and theoretical temperature dependence is best when the eigenvectors and energies for the entire $f²$ set used are those that best fit the optical and magnetic data for the system.

The negative shift for both ${}^{13}C$ and ${}^{11}B$ indicate the polarization contribution is the largest, but the sizable divergences between prediced and experimental dependencies is strong evidence for a large contribution from a covalent mechanism directly involving the 5f orbitals of the uranium atom.

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Appendix

It is obvious that for a calculation utilizing the entire $fⁿ$ baasis set and not just one term of the free ion computer programs must be used. In the following we give formulas that are easily programmable and allow separation of the *L* and *S* parts so that orbital reduction factors can be introduced.

The wave function Ψ in the matrix element $\langle \Psi | 0 | \Psi \rangle$ consists of a number of components

$$
|\Psi_i\rangle = \sum_i c_j' |\gamma_j S_j L_j J_j J_z\rangle \tag{A1}
$$

and the problem reduces to the calculation of matrix elements of the form^{22,28}

$$
\langle \gamma SLJJ_z|k_zL_z + g_e S_z | \gamma' S' L' J' J_z' \rangle \tag{A2}
$$

(where $g_e = 2.0023$), which can be expressed as

$$
\langle \gamma SLJJ_z | k_z L_z + g_e S_z | \gamma' S' L' J' J_z' \rangle =
$$

$$
(-1)^{J-J_z} \begin{pmatrix} J & 1 & J' \\ -J_z & 0 & J_z \end{pmatrix} \langle \gamma SLJ | k_z L_z + g_e S_z | \gamma' S' L' J' \rangle
$$
 (A3)

(27) Marks, T. **J.; Kolb, J.** R. *J. Am. Chem. Sot.* **1975,** *97,* **27.**

⁽²⁵⁾ Morton, J. R.; Rowlands, J. R.; Whiffen, O. H. Natl. Phys. Lab. (U.K.), Rep. 1962, BPR13. Morton, J. R. Chem. Rev. 1964, 64, 453.

⁽²⁶⁾ Fischer, R. D. *Organometallics of the f Elements*; Marks, T. J., Fischer, R. D., Eds.; D. Reidel: Dordrecht, The Netherlands, **1979 p 337,** and references therein.

⁽²⁸⁾ Judd, **B.** R. *Operator Techniques in Atomic Spectroscopy;* McGraw-Hill: **New** York, 1963.

where the quantity in parentheses is a *3j* symbol. The above *3j* symbol is not zero only if $J_z = J'_z$ and $|J - J'| \ge 1$. The reduced matrix element can now be separated into an orbital and a spin part

$$
\langle \gamma SLJ | k_z L_z + g_e S_z | \gamma' S' L' J' \rangle =
$$

\n
$$
k_z(-1)^{S+L+J'+1} \begin{cases} J & 1 & J' \\ L' & S & L \end{cases} \{ (2J+1)(2J'+1) L(L+1)(2L+1) \}^{1/2} +
$$

\n
$$
g_e(-1)^{S+L'+J+1} \begin{cases} J & 1 & J' \\ S' & L & S \end{cases} \{ (2J+1)(2J'+1)S(S+1)(2S+1) \}^{1/2}
$$
\n(A4)

where the quantities in braces are *6j* symbols. It can easily be shown that the reduced matrix element is not zero only for γ = γ' , $S = S'$, and $L = L'$. Equation A4 can also be used to calculate matrix elements of S_z alone by simply setting k_z to zero and dividing the result by **g,.** The necessary *3j* and *6j* symbols are obtained from easily programmable equations.29

(29) Rotenberg, M.; Bivins, R.; Metropolis, N.; Wooten, J. K., Jr. *The 3j and 6j Symbols;* MIT Press: Cambridge, MA, 1959.

Contribution from the Laboratorio di Chimica Nucleare and Centro di Strutturistica Diffrattometrica, Dipartimento di Chimica, Universita di Ferrara, 44100 Ferrara, Italy, and Department of Chemistry, University of Cincinnati, Cincinnati, Ohio 45221

Synthesis and Structure of an Amino Sugar-Schiff Base Complex of Technetium(V) Containing Salicylddehyde in an Unusual Coordination Mode

Adriano Duatti,*+ Andrea Marchi,' Luciano Magon,' **Edward** Deutsch,*t Valerio Bertolasi,§ and Gastone Gilli[§]

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The technetium(V) complex TcOL(sal), where L represents dianionic **N-salicylidene-D-glucosamine** and sal represents monoanionic salicylaldehyde, has been synthesized and characterized. This neutral species precipitates from a concentrated methanol solution after reaction of a 5-fold excess of the glucose derivative with $TcOCl₄$. The presence of chelated salicylaldehyde in the final compound does not appear to result from hydrolysis of the Schiff base ligand L prior to coordination since the reaction of $TcOCl₄$, L, and salicyldldehyde in a 1:1:1 stoichiometric ratio does *nof* yield TcOL(sa1). It thus appears that the formation of TcOL(sa1) is a kinetically controlled process. The crystal structure of the title compound, **(N-salicylidene-D-glucosaminato)(salicyl**aldehydato)oxotechnetium(V), was determined by X-ray analysis using counter data. The technetium atom is in a distorted octahedral coordination environment with the three donor atoms of the Schiff base ligand (Le., the neutral aldimine nitrogen atom, the anionic phenolic oxygen atom of the Schiff base moiety, and the anionic hydroxylic oxygen atom of the glucopyranose ring) occupying the plane normal to the Tc=O linkage. The coordination shell is completed by the binding of a salicylaldehyde ligand through a charged phenolic oxygen atom and a neutral carbonyl oxygen atom in positions cis and trans to the $Tc=O$ linkage, respectively. This unusual coordination mode of salicylaldehyde, in which the neutral atom is trans to the M=O linkage, has not been previously observed in oxo complexes containing similar ligands and presumably results from the formation of this complex by a kinetically controlled process. The length of the Tc-0 bond trans to Tc=O is 2.360 **(9)** A, while the displacement of the Tc atom out of the mean equatorial plane toward the oxo group is **0.422** (1) **A.** This complex crystallizes in the orthorhombic space group $P2_12_12_1$ with $a = 6.533$ (2) \hat{A} , $b = 12.649$ (4) \hat{A} , $c = 23.675$ (7) \hat{A} , and $V = 1956$ (1) \hat{A}^3 with $Z = 4$, for 1445 observed reflections with $I > 2\sigma(I)$.

Introduction

There is great interest in developing ^{99m}Tc radiopharmaceuticals suitable for monitoring the metabolic function of the brain and heart because of the ideal nuclear properties and availability of this isotope.' Glucose is an important substrate for brain and heart metabolism and hence it would be desirable to incorporate this sugar into the design of a 99m Tc radiopharmaceutical. The ultimate goal of this design is an agent in which technetium-99m is linked to a glucose molecule in a manner such that the biodistribution of the metabolically active sugar is not altered or destroyed. In principle, such agents can be prepared by using bifunctional chelating agents (BCA) in which one functional group strongly coordinates to the technetium center (e.g., EDTA, DTPA, thiosemicarbazone, thiols, or dithiols¹⁻¹²) and a second functional group provides the biologically active substrate or binds to it. This **BCA** approach is applied here in the preliminary development of a radiopharmaceutical wherein technetium is bonded to a chelating Schiff base functionality, which in turn is linked to a metabolically active glucose molecule.

A number of Schiff base complexes of Tc(V) and Tc(II1) have been recently reported,¹³⁻²⁰ illustrating the efficiency of this class

- (1) Deutsch, E.; Libson, K.; Jurisson, *S.;* Lindoy, F. L. *Prog. Inorg. Chem.* **1982, 30,** 75-139.
- **(2)** Lanteigne, D.; Hnatovich, D. J. *Int. J. Appl. Radiat. Isof.* **1984,** *35,* **6 1** 1-62 **1.**
- (3) Seifert, S.; Noll, B.; Muenze, R. Int. J. Appl. Radiat. Isot. 1982, 33, 1393-1398.
- **(4)** Noll, B.; Seifert, S.; Muenze, R. *Int. J. Appl. Radiat. Isot.* **1983,** *34,* 581-584.
- **(5)** Burgi, H. B.; Anderegg, G. A,; Blauenstein, P. *Inorg. Chem.* **1981,** *20,* 3829-3834.
- **(6)** Brenner, D.; Davison, A,; Lister-James, J.; Jones, A. L. *Inorg. Chem.* **1984, 23,** 3193-3197.
- **(7)** DePamphilis, B. **V.;** Jones, A. G.; Davison, A. *Inorg. Chem.* **1983,** *22,* 2292-2297.
- (8) Davison, **A.;** Jones, **A.** G.; Orvig, C.; Sohn, M. *Inorg. Chem.* **1981.20,** 1629- 1 632.
- (9) Davison, **A.;** Orvig, C.; Trop, H. **S.;** Sohn, M.; DePamphilis, B. **V.;** Jones, A. G. *Inorg. Chem.* **1980,** *19,* 1988-1992.
- (10) DePamphilis, B. **V.; Jones.** A. G.; Davies, **M. A,;** Davison, A. *J. Am. Chem. SOC.* **1978,** *100,* 5570-5571.
- (11) Smith, **J. E.;** Byrne, **E.** F.; Cotton, F. **A,;** Sekutowsky, J. C. *J. Am. Chem. SOC.* **1978,** *100,* 5571-5572.
- **(1** 2) Pinkerton, T. C.; Dailets, C. P.; Hoch, D. J.; Mikelsons, **M. V.;** Wilson, G. **M.** *J. Chem. Edur.* **1985, 62,** 965-913. (13) Ichimura, **A,;** Heineman, W. R.; Deutsch, E. *Inorg. Chem.* **1985, 24,**
- 2134-21 39.
- (14) Jurisson, *S. S.;* Dancey, K.; McPartlin, **M.;** Tasker, P. A,; Deutsch, E. *Inorg. Chem.* **1984, 23,** 4143-4149.

[†] Laboratorio di Chimica Nucleare, Dipartimento di Chimica, Universită di Ferrara.

^{*}University of Cincinnati.

^fCentro di Strutturistica Diffrattometrica, Dipartimento di Chimica, Universită di Ferrara.