Reprinted from the Proceedings of the National Academy of Sciences Vol. 61, No. 3, pp. 812–818. November, 1968.

THE EFFECT OF PRESSURE ON THE OXIDATION STATE OF IRON, II. OXYGEN LIGANDS*

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FEB 24 1969

Communicated August 12, 1968

In a series of papers from this laboratory¹⁻⁴ we have discussed the effect of pressure on the oxidation state of iron, on the isomer shift (s-electron density at the nucleus), and on the quadrupole splitting (splitting of the nuclear level of spin 3/2 due to interaction with an electric field gradient at the nucleus) for a variety of iron compounds. Since high-spin ferrous ions exhibit a relatively low electron density at the nucleus (isomer shift $\sim 1.2-1.4$ mm/sec relative to metallic iron) and large quadrupole splitting (2–3 mm/sec), whereas high-spin ferric compounds show a distinctly lower isomer shift (0.3–0.5 mm/sec) and relatively small quadrupole splitting (0.3–0.6 mm/sec), it is easy to identify one phase in the presence of the other.

The observations can be summarized as follows:5

(1) With increasing pressure, Fe(III) ion reduces to the Fe(II) state (reversibly) in a wide variety of compounds including halides, phosphate, sulfate, citrate, ferricyanide, etc. The conversion as a function of pressure is of the form:

$$K = AP^B, \tag{1}$$

where $K = C_{II}/C_{III}$, with C_{II} and C_{III} the nominal concentrations of Fe^{II} and Fe^{III} measured from peak areas, and A and B are constants. From thermodynamic arguments^{1, 5} one can write:

$$\frac{\partial \ln C_{\rm II}}{\partial \ln P} = \frac{P(V^{\rm III} - V^{\rm II})}{RT} C_{\rm III}.$$
(2)

The fractional increase in conversion per fractional increase in pressure is thus proportional to the concentration of sites available for conversion: the proportionality coefficient is the work to convert a site measured in thermal units, and is independent of pressure. For typical ionic materials $B \sim 0.5$. The reaction is endothermic with a heat of reaction which increases with temperature and is independent of pressure for compounds like the halides.

The electron transfer is from ligand nonbonding to metal antibonding orbital $(t_{2u} \rightarrow t_{2g} \text{ in octahedral symmetry})$. The reduction of the energy difference between these two sets of orbitals with increasing pressure is associated with a spreading of the 3*d* metal orbitals observed optically and probably also with a relative increase in pi bonding that stabilizes the t_{2g} orbitals.^{1, 5}

(2) With increasing pressure, the isomer shift decreases (the electron density at the nucleus increases) for most compounds. This can be discussed in terms of decreased shielding of the 3s-electrons by the 3d, due to the spreading of the 3d orbitals mentioned above, and of deformation of the s orbitals.⁵

Vol. 61, 1968 CHEMISTRY: FUNG, LEWIS, AND DRICKAMER

(3) For high-spin ferric ions there is generally a significant increase in quadrupole splitting with pressure. Since the ground state is an s state, the splitting is due to lack of cubic symmetry in the surroundings, and the electric field gradient is very sensitive to compression of the lattice. The much larger quadrupole splitting of high-spin ferrous ions, primarily caused by the aspherical 3d shell, tends to increase or decrease by modest amounts with increasing pressure.

In this paper we present high-pressure Mössbauer resonance data for ferric acetyl acetonate $[Fe(C_5H_7O_2)_3]$, basic ferric acetate $[Fe_3(CH_3COO)_6 (OH)_2CH_3-COOH_2O]$, ferric oxalate $[Fe(C_2O_4)_3 \cdot 5H_2O]$, and strontium trisoxalatoferrate (III) $[Sr_3(Fe(C_2O_4)_3)_2 \cdot 2H_2O]$. The compounds were synthesized from iron enriched to 85 per cent in Fe⁵⁷. The basic acetate and the acetyl acetonate were made as described in reference 1; the ferric oxalate was made by the procedure of Weinland and Rein,⁶ and the strontium salt was synthesized by the technique utilized by Gallagher.⁷ The oxalates were handled in the dark or by red light to prevent photoreduction. All compounds gave the correct chemical analyses and exclusively ferric Mössbauer spectra at one atmosphere.

All four compounds have in common the fact that the ligands are bidentate and the iron is surrounded by six oxygen atoms (or ions) in essentially octahedral symmetry.⁷⁻¹⁰ Early data on the acetate and acetyl acetonate indicated some conversion of Fe(III) to Fe(II) with increasing pressure at room temperature,¹ but the results were not quantitative. The high-pressure Mössbauer techniques have been described elsewhere.¹¹

We shall be primarily concerned with the conversion of Fe(III) to Fe(II) as a function of pressure and temperature. The experimental results are exhibited in Figures 1 and 2. As can be seen, they follow the relationship of equation (1), but $B \neq 0.5$ and is a distinct function of temperature, in contrast to the behavior of compounds previously discussed. Values of A and B are listed in Table 1 and heats of formation in Table 2.

Consider first the acetate and acetyl acetonate. The work $P\Delta V$ to form Fe(II) from Fe(III) is considerably larger at a given temperature and pressure for these complexes than for the simpler ionic crystals. The volume change $V^{III} - V^{II}$ amounts to several cubic centimeters at 10 kilobars (kb). One can understand that for these bidentate ligands it would be more difficult to distribute the strain involved in the charge transfer and formation of ion radicals than it would be for FeCl₃ or FeBr₃. The smaller value of B (and of ΔV) at higher temperature of the strain the charge transfer and formation of the strain temperature of the strain temperature of the strain for the smaller value of the strain temperature of t

$a D for R - AI^{-}.$		
Temperature (°K)	A	В
295	$1.24 imes10^{-5}$	2.23
375	$0.96 imes 10^{-2}$	1.013
378	$0.22 imes10^{-6}$	3.05
418	$2.21 imes10^{-2}$	0.983
295	0.041	0.51
335	0.029	0.833
383	0.043	1.146
295	0.115	0.301
383	0.058	0.844
	Temperature (°K) 295 375 378 418 295 335 383 295 383	Temperature (°K) A 295 1.24×10^{-5} 375 0.96×10^{-2} 378 0.22×10^{-6} 418 2.21×10^{-2} 295 0.041 335 0.029 383 0.043 295 0.115 383 0.058

TABLE 1. Constants A and B for $K = AP^{B}$.

Material (kb)						
Acetyl acetonate	325°K		375°K			
60	0.15		0.25			
100	0.105		0.15			
150	0.065		0.085			
200	0.032		0.037			
Basic acetate		398°K				
75		0.93				
100		0.60				
150		0.44				
Oxalate	315°K		360°K			
25	0.19		0.34			
70	0.23		0.39			
100	0.26		· 0.42			
150	0.29					
Strontium ferric oxalate		333°K				
20		0.11				
50		0.15				
100		0.20				
200		0.24				

tures can probably be associated with thermal relaxation of the strains. The heat of reaction decreases markedly with increasing pressure.

On the other hand, the oxalates exhibit a value of $B(P\Delta V/RT)$ and a heat of formation of the Fe(II) site which increases with pressure, i.e., the work to perform the reduction is greater at high than at low temperature. It is known that oxalates tend to reduce photochemically,^{12, 13} and when heated in the dark, they undergo a series of reactions; first reduction, then oxidation.¹⁴ In both cases CO₂ is evolved and the process is irreversible.

As indicated earlier, the high-pressure reduction is reversible and involves a lowering of energy of metal antibonding levels with respect to ligand nonbonding p-orbitals, so it is not necessary to postulate formation of CO₂ in the lattice. Nevertheless, the radical formed in the process quite possibly has a volume in the lattice that is rather temperature-dependent. It is of interest to note that the slope B, and therefore the volume difference $(V^{III} - V^{II})$, is noticeably smaller for the strontium salt than for the ferric oxalate, although the number and gross symmetry of the ligands is apparently the same for both compounds. This illustrates the important effect of the ability to redistribute the strain to next-nearest neighbors and beyond on the conversion equilibrium.

While the main interest in this study was the conversion of Fe(III) to Fe(III), isomer shift and quadrupole splitting data were obtained both for the Fe(III)and for the Fe(II) formed at high pressure. These results appear in Tables 3 and 4. Isomer shift data are shown only at room temperature, for it was necessary to change the source location slightly with increasing pressure, which caused a slight change in its temperature. Peak locations established with changing amounts of Fe(III) and Fe(II) present are necessarily less accurate than those obtained in nonreacting systems, but one can comment on two trends of interest whose magnitude are well beyond any experimental error.

814



FIG. 1.—Ln K vs. ln P—acetate and acetyl acetonate.



FIG. 2.—Ln K vs. ln P—ferric oxalate and strontium ferric oxalate.

CHEMISTRY: FUNG, LEWIS, AND DRICKAMER PROC. N. A. S.

Pressure	Acetyl Acetonate		Oxa	late	Strontium Oxalate		
(kb)	Fe(III)	Fe(II)	Fe(III)	Fe(II)	Fe(III)	Fe(II)	
0	0.35	-	0.45	-	0.41		
25	0.38	1.28	0.44	1.24	0.40	1.27	
50	0.40	1.20	0.43	1.26	0.39	1.29	
75	0.39	1.16	0.43	1.29	0.385	1.31	
100	0.38	1.14	0.42	1.31	0.38	1.33	
125	0.36	1.12	0.41	1.34	0.375	1.36	
150	0.33	1.11	0.40	1.36	0.37	1.38	
175	0.33	1.10	0.39	1.38	0.365	1.40	

TABLE 3.	Isomer	shift	versus	pressure	at	2950	K.	*
			0.01.0000	10.000000000		1000		×.,

* Mm/sec relative to iron metal at 1 atmosphere.

TABLE 4. Quadrupole splittings (mm/sec).

			(A) Ferric Id	on				
Pressure (kb)	Basic . 378K°	Acetate 418°K	Acetyl A 295°K	cetonate 375°K	Oxa 295°K	late 335°K	Strontium 295°K	m Oxalate 382°K	
0			0.46	0.70	0.59	0.54	0.50	0.72	
25			0.76	1.00	0.82	0.77	0.68	0.88	
50		0.85	0.80	1.10	0.95	0.92	0.78	1.00	
75	0.82	0.90	0.92	1.15	1.02	1.00	0.90	1.10	
100	0.90	0.95	1.03	1.17	1.08	1.07	1.02	1.18	
125	0.96	1.02	1.14	1.20	1.13	1.12	1.12	1.25	
150	1.02	1.06	1.22	1.23	1.20	1.16	1.16	1.32	
175	1.08	1.10	1.28	1.26	1.26	1.20	1.20	1.38	
			(E	3) Ferrous I	on				
Pressure	ressure Basic Acetate		Acetyl Acetonate		-Oxal	Oxalate		Strontium Oxalate	
(kb)	$378^{\circ}\mathrm{K}$	418°K	$295^{\circ}K$	375°K	$295^{\circ}K$	335°K	295°K	383°K	
0								_	
25			2.20	2.10	1.90	1.90	2.00	2.00	
50		1.80	2.15	2.05	2.00	2.00	2.08	2.08	
75	2.20	1.84	2.12	2.00	2.10	2.08	2.14	2.14	
100	2.16	1.86	2.00	2.00	2.18	2.16	2.20	2.20	
125	2.14	1.88	2.00	1.98	2.23	2.21	2.26	2.26	
150	2.12	1.90	2.00	1.96	2.28	2.23	2.30	2.30	
175	2.10	1.90	2.00	1.95	2.30	2.25	2.35	2.35	

In both oxalates the isomer shift of the ferrous ion formed increased with pressure, i.e., the electron density at the iron nucleus decreased. This is in contrast to the behavior of all but a few other systems, and indeed to that of ordinary ferrous oxalate.¹ The bonding must be very strained and changes in occupation of the bonding orbitals with pressure apparently overwhelm the shielding and orbital compression effects.⁵

In the acetyl acetonate, there is a maximum in the isomer shift at about 40–50 kb, as shown in Figure 3. The apparent isomer shift at 102°C is plotted in the same figure to show that it also exhibits the maximum. In this case the changes in orbital occupation dominate in the low-pressure region, whereas the usual orbital distortion effects govern at high pressure. Figure 4 shows the quadrupole splitting for acetyl acetonate as a function of pressure. At room temperature there is a distinct leveling in the region of the isomer shift maximum, then a second rise at higher pressures. It is clear that the change in orbital occupation of the bonding orbitals affects the local asymmetry. At 102°C the change in quadrupole splitting with pressure is too small to determine whether the irregularity is

816



0.4 0.4 100 150 200 PRESSURE (Kbar)

FIG. 4.—Quadrupole splitting vs. pressure—ferric acetyl acetonate (Fe(III)).

present. The increase in quadrupole splitting with increasing temperature in the low-pressure region is unusual. The anomalies in isomer shift and quadrupole splitting are not reflected in the conversion, which shows no irregularities with increasing pressure or temperature. This is not surprising as the conversion involves only nonbonding and antibonding orbitals.

Summary.—The effect of pressure on the oxidation state of ferric ion, in a site of essentially octahedral symmetry with six oxygen ligands, has been measured. Four compounds have been studied: basic ferric acetate, ferric acetyl acetonate, ferric oxalate, and strontium ferric oxalate. For all these compounds the reduction followed the relationship $K = AP^B$ where K is the equilibrium constant C_{II}/C_{III} . For the acetate and acetyl acetonate, B decreased with increasing temperature, whereas both oxalates showed an increase in B with increasing temperature. These observations are discussed in terms of thermodynamic implications. The unusual behavior of the isomer shift and quadrupole splitting of the ferric ion in the acetyl acetonate is briefly discussed.

* This work was supported in part by the U.S. Atomic Energy Commission.

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818