NORMAL COORDINATE TREATMENT OF LANTHANIDE HEXAHALIDE ANIONS $(L_n X_2^{3-})^{\frac{1}{4}}$

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A. INTRODUCTION

Hexahalide complexes of the lanthanide metals have been prepared $^{1-4}$ in the form $(RH)_3 LnX_6$, where RH represents the pyridinium, triphenyl-phosphonium or tetrabutyl-ammonium cation. The lanthanide hexahalide anion, LnX_6^{3-} , is six-coordinate and believed to occupy a local site of near-octahedral symmetry in the crystalline state $^{1-4}$. These six-coordinate complexes are particularly interesting since lanthanide metals tend to go to higher coordination numbers (e.g. 8, 9, 10 or 12) in other complexes. In a previous paper the far-infrared and Raman skeletal vibrations of the LnX_6^{3-} anion have been determined. These vibrational data for the $(pyH)_3 LnX_6$ complexes have been used to perform normal coordinate treatments by assuming five different potential fields. The force constants determined indicate several behavioral trends occurring within the series. Ferraro and coworkers recently presented a complete survey of metal hexahalide systems as studied using these five force fields, indicating that no theoretical work or force constants were previously available for any lanthanide hexahalide units with the exception of $CeCl_6^{2-}$.

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TABLE 1	
Symmetrized G matrix elements for the octahedral $\operatorname{LnX}_6^{3-}$ unit	it

Symmetry species	G matrix element a
410	$G_{i1} = \mu_{\chi}$
	$G_{Z2} = \mu_X$
4 _{1g} 5g 51u	$G_{33} = \mu_x (1 + 2\mu_{Ln}/\mu_x)$
- -	$G_{34} = -4\mu_{Ln}$
	$G_{44} = 2\mu_{X}(1 + 4\mu_{Ln}/\mu_{X})$
72g	$G_{55} = 4\mu_{\chi}$
	$G_{66} = 2\mu_{x}$

 $a \mu_n = \text{reciprocal mass of } n$.

B. CALCULATIONS

Computer calculations in this study were performed by an IBM 360-195 or a Sigma V computer. Computer drawings were carried out using a Cal Comp 780 incremental plotter.

C. NORMAL COORDINATE TREATMENT OF LnX₆³-

Normal coordinate treatment of the LnX_6^3 anion (where Ln = Nd, Eu, Gd, Dy, Er, Yb and X = Cl, Br) was performed by assuming local octahedral symmetry. All the force fields used here⁵⁻¹² have been described in the literature. They are the General Valence Force Field (GVFF), Urey—Bradley Force Field (UBFF), Modified Urey—Bradley Force Field (MUBFF), Orbital Valence Force Field (OVFF) and Modified Orbital Valence Force Field (MOVFF). The elements of the F and G matrices are listed in Tables 1 and 2.

The GVFF contains five force constants. These are f_r , f_{α} , f_{rr} , $f_{\alpha\alpha}$ and $f_{r\alpha}$, where f_r and f_{α} are the stretching and bending terms respectively. The remainder are interaction force constants where f_{rr} is the interaction term for two adjacent bonds, $f_{\alpha\alpha}$ for two angles sharing a common bond and in the same plane, and $f_{r\alpha}$ for an angle and a bond included in the angle. The UBFF contains four force constants, where K and H are the stretching and bending force constants and F and F' are repulsion constants between non-bonded atoms. The MUBFF has five independently variable force constants. Three of these are the standard UBFF constants K, H and F. F' is allowed to equal $-\frac{1}{10}F$ as calculated when it is assumed that the forces between non-bonded atoms are proportional to $1/r^9$, where r is the distance between non-bonded atoms. Two additional GVFF-type interaction constants are added. These are the constants h and h, where h is the interaction constant between two angles sharing a common bond and at right angles to each other, and h is the interaction constant for two bonds trans to each other. The OVFF has four force constants -K, h, h and h. These are analogous to UBFF constants except that h is the

UBFF ¹⁶	MUBFF ^{16,24}	OVFF ²⁵	MOVEF ⁵	GVFF
$F_{11} = K + 4F$	F11 = K + 4F + F	F11 = K+4F	$F_{11} = K + 4F + k$	$F_{11} = f_p + 5.33 f_{PP}$
$F_{12} = K + F + 3F'$	$F_{22} = K + 0.7F + k$	$F_{22} = K + F + 3F'$	$F_{22} = K + F + 3F' + k$	$F_{22} = f_F - 0.667 f_{FT}$
$F_{33} = K + 2F + 2F'$	$F_{33} = K + 1.8F - k$	$F_{33} = K + 2F + 2F'$	$F_{33} = K + 2F + 2F' - k$	$F_{33} = f_{\nu} - 1.33 f_{\tau}$
F34 = F + F'	$F_{34} = 0.9 F$	F34 = F + F'	F34 = F+F'	F34 = 2/1a
$F_{44} = H + F/2 - 3/2F'$	$F_{44} = H + 0.65 F + 2h$	$F_{44} = D/2 + F/2 - 3/2F'$	$F_{44} = D/2 + F/2 - 3/2F'$	$F_{44} = f_{\alpha} + 2f_{\alpha\alpha}$
$F_{55} = H + 1/2F - 1/2F'$	$F_{55} = H + 0.55F$	$F_{55} = D/4 + F/2 - F'/2$	$F_{55} = D/4 + F/2 - F'/2$	$F_{55} = f_{\alpha} - 2f_{\alpha\alpha}$
Fec = H + 1/2F + 1/2F'	$F_{66} = H + 0.45 F - 2h$	$F_{66} = D/2 + F/2 + F'/2$	$F_{66} = D/2 + F/2 + F'/2$	$F_{66} = f_{\alpha} - 2f_{\alpha\alpha}$

Observed and calculated frequencies and force constants for several lanthanide hexalialide, ${\rm Ln}\chi_6^3$, complexes a

TABLE 3

		Treamen	Treduction (cm-1)					2			6			
							•	anna.	# CO 21	nts (mo	rotte constants (mayne/A)		Average percentage deviation	centage
Hexahalide b Ln χ_6^3		(81 W) 14	$ u_2(E_g) $	1,3 (F _{LL})	ν4 (F ₁₁₁)	vs(F2g)	ν ₆ (F ₂₁₄)	* ~	ند م <u>ـ</u> د	H(D)	h(F') ∫αα	* ~ 5	(64,54,14)	(4, 1, 1, 5, 1, 6)
NdCl3~	Observed	ł	204	226	121	113	(80)			1		-		
•	UDFF	241	193	242	116	121	1	0.69	0.13		-0.01		5.6	6.0
	MUBFF	246	207	226	116	120	11	0.71	0.11		0.00	0.11	4.1	. 4
	OVFF		194	242	119	116	22	0.71	0.11		0.0	;	6.2	~ - -
	MOVFF		208	226	120	115	80	0.72	0.10		-0.01	0.11	1.6	8.0
	GVFF		204	226	121	113	80	0.91	0.08	90.0	0.01	90.0	0.0	0.0
Ndbr.3	Observed		(123)	167	8	92	(54)							
,	UBFF		119	172	88	83	25	0.57	0.13	0.01	-0.01		2.3	5,3
	MUBFF	153	123	167	68	81	52	0.58	0.12	0.01	0.001	0.05	0.3	5.2
	OVFF		119	174	6	79	54	0.59	0.12	0.03	-0.01		3.5	2.5
	MOVFF		125	167	93	11	54	0.60	0.10	90.0	-0.02	0.08	1.2	1:1
	GVFF		123	167	94	91	54	92.0	0.07	0.03	0.01	0.04	0.0	0.1
EuCl3-	Observed	_	(216)	226	144	111	(80)							
,	UBFF	243	196	246	125	124	75	0.74	0.12	0.01	-0.02		7.7	10.4
	MUDEF		220	226	126	124	75	0.78	60.0	0.03	0.01	0.16	1.3	10.2
	OVFF		197	249	133	118	80	0.79	60.0	0.07	-0.02		9.3	4.6
	MOVFF		221	228	136	115	80	0.85	90.0	0.10	-0.02	0.19	2.0	3.4
	GVFF C		216	235	128	112	5	1.02	0.01	0.11	0.02	0.12	1.4	4,3
EuBr?	Observed		(126)	164	6	81	(51)							
•	UBFF	156	119	171	33	98	Se	0.57	0.14	0.01	0.01		3.1	4,5
	MUDEF	156	126	164	93	86	55	0.59	0.12	0.02	0.001	0.0	0.1	4.3
	OVFF 1	154	119	172	8	83	57	0.58	0.13	0.03	-0.01		3.9	2.1
	MOVFF	155	126	164	6	81	57	0.60	0.11	90.0	-0.02	0.0	0.3	0.3
	GVFF		126	164	76	81	57	0.79	0.07	0.09	0.01	0.05	0.0	0.1

TABLE 3 (continued)

	<u>.</u>	Frequen	Frequencies (cm ⁻¹)					Force	Force constants (mdyne/A)	nts (md	yne/A)		Average percentage deviation	entage
Hoxahalide ^b LnX ₆		(4 1g)	v ₂ (E _g)	v3(F1u)	"4(F _{1,"})	v3(F1u) v4(F1u) v5(F2g) v6(F2u)	"6(F2u)	* *	12. LE	H(D)	h(F') fax	عري	(v ₁ , ν ₂ , ν ₃)	(V, 'US, U6)
GdCi3-	Observed	258	(202)	227	135	120	(85)	0.70	0.14]	-0.02		5.4	6.3
	MUBEE		206	228	127	130		0.73	0,12	0,02	0.003	0.09	1.9	6,2
	OVFF		194	243	131	125	85	0.73	0.12		-0.02		0'9	2.3
	MOVFF	247	207	227	134	123	8	0.74	0.11		-0.02	0.1	2.2	1.4
	GVFF	258	202	227	135	120	82	0.91	0.09	_	0.01	9.0 B	0.0	0.0
GdBr ³ .	Observed	157	(122)	163	90	85	(61)							
•	UBFF		120	167	88	88	8	0.54	0.15	0,0	-0.01		1,6	2.2
	MUBFF		122	163	68	83	9	0.57	0.13	0.02	-0.005	9. 2	6.3	1.8
	OVFF		120	166	68	85	9	0.54	0.15	0.03	-0.01		1.2	0.5
	MOVFF		122	163	8	82	61	0.54	0.15	0.03	-0.01	0.03	0.0	0.0
	GVFF		122	163	8	98	61	0.75	0.08	0.09	0.003	0.03	0.0	9,4
DyCl3.	Observed		(198)	230	139	118	(81)							
B	UBFF		193	242	128	129	11	0.71	0.14	0.006	-0.02		4.0	7.0
	MUBFF		203	231	128	129	11	0.70	0.13	0.02	0.006	0.07	2.2	7.2
	OVER		194	243	134	123	18	0.75	0.11	0.05	-0.02		5.1	3.7
	MOVFF		204	230	137	122	8	0.76	0.0	0.07	-0.03	0:0	2.8	1.9
	GVFF	257	198	230	139	116	82	0.88	60.0	0.10	0.01	0.02	0.0	6.0
DyBr ³ -	Observed		(124)	161	\$	90	(64)							
	UBFF		120	166	35	93	63	0.54	0.16	0.05	-0.01		2.2	1.5
	MUBFF		124	191	93	6	83	0.57	0.14	0.02	0.005	9.0	0.2	1.2
	OVEF		120	165	93	8	æ	0.53	0.17	0.03	-0.006		2.2	0 .4
	MOVFF		123	191	25	8	\$	0.53	0.16	0.04	-0.01	0.0	9.0	0.5
	GVFF	159	124	191	94	8	Ē	0.78	0.08	0.10	0.002	0.04	0:0	2.1

TABLE 3 (continued)

والمراسسة والمقدمة المراسم والمراسم	Fieq	Frequencies (cm ⁻¹)	_			 	Force	constan	nts (md	Force constants (mdync/A)		Average percentage deviation	entage
Hexahalide b	, ,,(4 ₁ g)	1g) "2(Eg)	$v_3(F_{1M})$	"4(F14)	vs(F _{2g}) v ₆ (F _{2u})	, v ₆ (F ₂ u)	* **	7.	H(D)	h(F') Saa	75 75	(v ₁ ,p ₂ ,v ₃)	(4, v _S , v ₆)
EC13-	Observed 260 UBFF 247 MUBFF 248 OVFF 238 MOVFF 244 GVFF 260	(200) 193 205 195 207 207	229 244 230 229 229	143 130 130 137 143	118 130 130 124 123	(81) 77 77 81 80	0.77 0.79 0.79 0.90	0.14 0.11 0.09 0.09	0.03 0.06 0.08 0.08	0.007 0.007 0.03 0.03	0.08 0.11 0.02	4.7 2.4 5.9 3.1 0.0	8.4 8.1 3.3 2.3 0.9
Вты г.	Observed 161 UBFF 160 MUBFF 160 OVFF 161 MOVFF 161	(121) 121 121 121 121 121	166 166 166 166 166 166	\$ 22 9 9 9 9 2 22 24 24 24 24 24 24 24 24 24 24 24 24	8 8 8 8 8 8 8 8 8 8 8 8 8 8 8 8 8 8 8	<u>6</u> 88222	0.55 0.57 0.54 0.54 0.75	0.16 0.19 0.17 0.17 0.09	0.02 0.01 0.03 0.03 0.11	-0.007 -0.005 -0.007 -0.005	0.02 0.004 0.02	0.5 0.3 0.1 0.0	1.9 1.6 0.1 0.1
₹bc1,	Observed 263 UBFF 245 MUBFF 250 OVFF 234 MOVFF 245 GVFF 263	(207) 197 213 1198 215	216 243 227 246 226 226	139 123 123 131 135	111 124 124 118 116	(6) 4 4 6 8 8 6 6 6 6 6 6 6 6 6 6 6 6 6 6 6	0.75 0.76 0.80 0.84 0.96	0.13 0.08 0.06 0.09	0.01 0.02 0.07 0.09	-0.02 0.006 -0.02 -0.03	0.16 0.05	6.5 2.7 8.0 3.7 0.0	9.8 8.9 9.2 9.2 9.2
YbBrg	Observed 160 UBFF 157 MUBFF 156 OVFF 151 MOVFF 152 GVFF 160	(119) 118 121 118 122 122	161 163 163 167 161 161	96 93 94 96	78 84 84 82 81 78	(55) 53 53 55 55	0.56 0.56 0.58 0.59 0.73	0.15 0.14 0.12 0.11 0.09	0.000 0.005 0.03 0.05 0.19	-0.02 0.002 -0.02 -0.02 0.06	0.03 0.06 0.17	1.9 1.2 3.4 0.0	5.9 5.8 2.9 0.1.

 g Observed frequencies listed in parentheses were derived from combination bands or calculations, b ALI observed frequencies are those for the solid pyridinium complex, $(pyH)_{3}LnX_{6}$. c Oscillating, best fit given.

contribution to the potential energy of $\Delta\beta$. Here $\Delta\beta$ is the angle deformation of the bond M-X away from its idealized hybrid orbital as opposed to $\Delta\alpha$ (of UBFF), which is the angle deformation of XMX. The MOVFF has five force constants. Four are basic OVFF constants (i.e. K, D, F and F') and the fifth is the GVFF-type interaction, k, as defined for MUBFF. The derivation and more detailed discussion of these force fields can be found elsewhere S^{-12} .

Solutions of secular equations were performed using a program of Yeranos and Foss¹³. A least-squares analysis was employed fitting calculated frequencies to observed experimental frequencies to give a "best" set of force constants. Table 3 gives the results of these calculations. Observed and calculated frequencies, as well as converged force constants are given for each force field. The percentage deviation of the calculated from the observed frequencies are given as an average for v_1 , v_2 and v_3 (essentially stretching) and v_4 , v_5 and v_6 (essentially bending). The General Valence Force Field values are given as a reference although an artificial problem where the number of knowns is equivalent to the number of unknowns has been created. This is a result of the calculation fixing $v_6 = v_5/\sqrt{2}$ which is implicit in GVFF assumptions. The result was a "perfect fit" solution for eleven of the twelve complexes for which GVFF calculations were performed. However, no convergence could be achieved for EuCl $_6^3$, and the refinement problem oscillated between two sets of force constants, neither of which gave a "perfect fit", Although several methods of damping the refinement process were tried for EuCl $_6^3$, no convergence could be achieved.

In this paper we assume that the force field giving the lowest average percentage deviation (i.e. best fit of calculated to observed frequencies) is the "best" potential energy force field. For all twelve of the LnX_6^{3-} units studied here, the MOVFF gives the "best" overall fit for both stretching and bending vibrations. MUBFF also gives a "good" fit for the stretching frequencies. However, MUBFF is noticeably bad for those vibrations which are essentially bending $(\nu_4, \nu_5$ and ν_6). This is manifest in the higher average percentage deviation for ν_4 , ν_5 and ν_6 for MUBFF compared with MOVFF. Previous studies comparing these force fields for octahedral complexes indicate that MOVFF gives best fits with observed frequencies for most hexachlorides and hexabromides⁵.

D. FORCE CONSTANTS

It is important to consider the significance of several individual force constants. The metal-halogen stretching force constant (f_r) or K) takes the largest value and is most variable among the force constants studied here. Further, this stretching force constant readily lends itself to direct comparison with such properties of the $\text{Ln}X_6^{3-}$ unit as bond strength, bond length, oxidation number and bond order. This metal-halogen stretching force constant is 0.72-0.79 mdyne/Å for the $\text{Ln}Cl_6^{3-}$ complexes and 0.54-0.59 mdyne/Å for the $\text{Ln}Br_6^{3-}$ complexes. The lower value for the bromide complexes was expected as

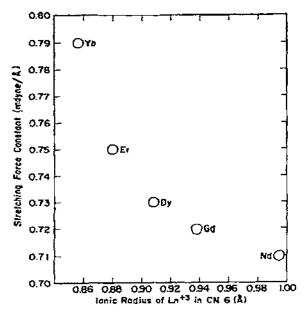


Fig. 1. Variation of the stretching force constant with the ionic radius of the metal for LnC_6^{3-} .

the observed fundamental halide shift can be accounted for only partially by the mass change⁴. In general, the metal-halogen stretching force constant, f_{LnX} , decreases in going from fluorine to iodine⁵. Our results are consistent with these observations. This order may be expected since the lanthanide metals act as Chatt-Ahrland Group A metals², so that stronger bonds are formed with lighter halogens (F > Cl > Bt > I).

Some variation of the metal—halogen stretching force constant occurs for different metals within the lanthanide series. As the ionic radius of the metal ion decreases by lanthanide contraction (moving towards higher atomic number) the force constant increases indicating a stronger and shorter metal—halogen bond. Figures 1 and 2 show the variation of the stretching force constant with the ionic radius of the Ln^{3+} metal. The force constant (K) plotted is the average of the refined force constant for UBFF, MUBFF, OVFF and MOVFF. The GVFF stretching force constant (f_r) was not included for the reasons discussed earlier. The ionic radii are those given for Ln^{3+} ions by Douglas and McDaniel¹⁴. Although Ln^{3+} ionic radii, as found in the literature, vary considerably in absolute value the trend observed here is consistent for several different reported radii. The value for $EuCl_6^{3-}$ was not included because it deviated widely and inexplicably from the other values. Figure 1 shows a consistent dependence of the stretching force constant on the ionic radius for the $LnCl_6^{3-}$ complexes. The $LnBr_6^{3-}$ series appears somewhat less consistent (Fig. 2). The bromides seem to be divided into two behavioral groups — one containing Nd and Eu, and one containing the heavier metals such as Gd, Dy, Er and Yb.

All of the force fields considered here, except GVFF, include Urey—Bradley-type repulsions of non-bonded atoms. Steele 15 has pointed out that these interactions occur

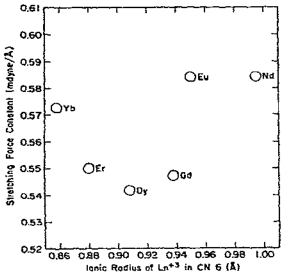


Fig. 2. Variation of the stretching force constant with the ionic radius of the metal for LnBr3.

predominantly in systems where steric effects between non-bonded nuclei are significant. We would expect that the lanthanide bromides would demonstrate more steric interactions than the chlorides since bromine is larger than chlorine. In fact, our calculations show larger repulsive force constants, F, for bromides than chlorides. For example, for $GdCl_6^3$, $F_{CI-CI} = 0.118$ mdyne/Å and $F_{Br-Br} = 0.145$ mdyne/Å. Further, as the ionic radius of the metal decreases (drawing the halogens closer together), the repulsive force constant, F, tends to increase for the bromides. For example, NdBr₆³, $F_{Br-Br} = 0.118$ m/dyne/A and for ErBr₆³, $F_{Br-Br} = 0.165$ mdyne/Å. It is rather interesting to note that an opposite trend is seen for the chlorides. YbBr3 ~ does not behave consistently. It is, perhaps, worthy of comment that the bromides show smaller average percentage deviations than the chlorides for all the force fields which include Urey-Bradley-type interactions (UBFF, MUBFF, OVFF, MOVFF). This tends to verify Steele's belief that these force fields are most applicable to systems involving large steric interactions¹⁵. A second explanation for the consistently low percentage deviation for LnBr₆ calculations (as compared with LnCl3 may be that the bromides more closely approach the assumed octahedral symmetry. On the basis of electronic spectra, Ryan2 draws similar conslusions that the LnX62 anion approaches octahedral symmetry in the order I > Br > Cl where $LnCl_a^3$ is the most distorted.

The F' repulsion constant appears in the UBFF, OVFF and MOVFF. In the MUBFF, F' was set equal to $-\frac{1}{10}$ F, assuming van der Waal type interactions between non-bonded atoms and covalently bonded molecules. Although the Ln-X bonds of the lanthanide hexahalide anions are somewhat ionic in character, the validity of the $F' = -\frac{1}{10} F$ approximation for LnX₆³ complexes becomes evident when F' is allowed to vary inde-

pendently of F (that is in UBFF, OVFF and MOVFF); the least-squares refined F' value always converged to very near $-\frac{1}{10}F$ (see Table 3).

Although the stretching interaction force constant $k(f'_{rr})$ of MUBFF and MOVFF is small, it is obviously highly significant. Both MOVFF and MUBFF give excellent fits for the stretching vibrations. This "good" fit indicates the importance of the interaction between non-adjacent (trans) metal—halogen bond stretchings. That is, the metal—halogen stretching vibration has a significant influence on the bond trans to it. Such an observation has been expressed previously in terms of the "trans effect" 17.

The force constant h of MUBFF is very small for all the molecules studied here. This force constant represents the interaction of adjacent angles sharing a common bond but not in the same plane. The smallness of this force constant and the poor fit for bending fundamentals (large percentage deviation for ν_4 , ν_5 , ν_6) in MUBFF lead us to conclude that $h(f'_{\alpha\alpha})$ is not an important interaction to be included in the potential energy description of the $\text{Ln}X_0^{3-}$ system.

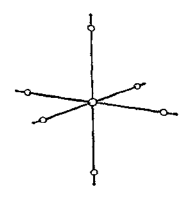
E. POTENTIAL ENERGY DISTRIBUTION

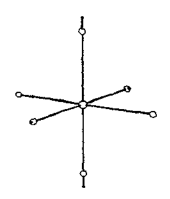
The potential energy distribution coefficient, PED, is defined by

$$PED = \Lambda^{-1} (JZ) \Phi$$

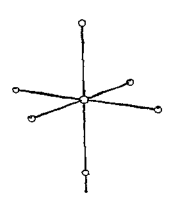
where JZ is the Jacobian matrix expressing the variation of the eigen-value with the force constants, Φ is the column force constant matrix, and Λ^{-1} is the reciprocal diagonal matrix of the eigenvalues¹⁹. The PED has been calculated over the force constants for each of the force fields and LnX_6^3 units. The results are not listed for the sake of brevity but are available from the authors on request.

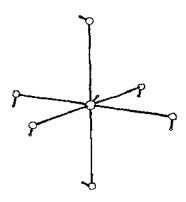
From the PED it is evident that v_1 and v_2 are pure stretching modes since these two vibrations include no contribution from the bending constants. The opposite is true for v_5 and v_6 which can be considered pure bending modes. Although v_4 is primarily a bending vibration and v_3 is largely stretching, a slight amount of coupling between these two modes is seen for some LnX_6^{3-} groups. However, the contribution is usually less then 5% and can be considered nearly insignificant. The importance of the $k(f_{rr}^i)$ constant, added to the MUBFF and MOVFF, is evident in the PED only for stretching vibrations. The $h(f_{\alpha\alpha}^i)$ force constant of MUBFF is insignificant for many of the complex ions, although it does indicate some participation in the v_4 and v_6 vibrations of some heavier metal chlorides. The PED of the repulsive force constant, F, is rather interesting. As might be expected, the PED of F tends to be greater for bending modes than for stretching modes. Further, F participation is, in general, larger for the bromides than for the chlorides and increases as the radius of the central metal decreases.



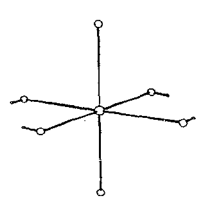


FREQ. 1 (A1G)

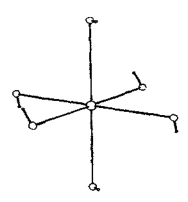




FREQ 3A (F1U)



FREG. 4A (F1U)



FREQ. 6A (F2U)

FREQ. 5A (F20)
Fig. 3. Normal vibrations of Nd X_6^3 . Symmetry species labels appear, as computer plotted, without subscripts, i.e. F1U is equivalent to F_{1M} .

F. CARTESIAN COORDINATE DISPLACEMENTS AND PLOTTING OF NORMAL MODES

The solution of the secular equation provides eigenvalues, PED information and force constants. However, it is rather difficult to visualize the actual vibration from these data. We can plot the molecular motion by transforming the normal mode of vibration into Cartesian coordinate space. The transformation matrix from the normal to Cartesian coordinates can easily be obtained using Schachtschneider's programs¹⁸. The calculations have been completed using both GVFF and UBFF with no significant changes in the results. The results were plotted using the crystallographer's ORTEP program¹⁹ as described by LaBonville and Williams²⁰. Figure 3 shows the resulting normal modes of vibrations. These displacements have been multiplied six times in order to make small displacements observable.

The importance of plotting out the actual normal vibrations for any system becomes apparent when the drawings of Fig. 3 are compared with any theoretical representation of normal vibrations for octahedral symmetry. Several standard texts give the theoretically expected vibration for MX₆ octahedral units²¹⁻²³. Although these are probably correct for the general case, some variations always occur in practice. In this study, the ν_1 , ν_2 , ν_3 and ν_5 vibrations agree with the standard descriptions of the normal modes. However, ν_4 and ν_5 vary slightly but significantly.

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