Intramolecular Force Field of Some Tetrahedral Thio and Seleno Anions *

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The orbital valence force field (OVFF) and the Urey-Bradley force field (UBFF) have been employed to evaluate the force constants of thio and seleno anions of VA, VB, VIB and VIIB group atoms of the periodic table. These constants were utilized to examine the relative strength of chemical bonds in isoelectronic sequences. Further, for these anions, the variation of the ratio L_{12}/L_{21} with the mass coupling term $(T=G_{12}/\sqrt{|G|})$ has been studied. This appears to be an excellent support to the extended L-matrix approximation **.

Recently, Müller et al. 1,2 have reported the infrared and Raman spectra of tetrathio and tetraseleno anions of Vanadium, Niobium and Tantalum and the tetrathio anion of Phosphorus. They have studied the force constants of the above anions 1 along with those of $\mathrm{MoS_4}^{2^-}$, $\mathrm{WS_4}^{2^-}$, $\mathrm{ReS_4}^-$, $\mathrm{MoSe_4}^{2^-}$ and $\mathrm{WSe_4}^{2^-}$ on the basis of tetrahedral symmetry employing the L-matrix approximation 3 , the P.E.D. method 4 and Fadini's approximation 5 . In the present communication, OVFF^{6-8} and UBFF^{9} have been applied to describe the nature of the force field. The variation of the ratio L_{12}/L_{21} with the mass coupling term 10 $(T=G_{12}/V|G|)$ has also been examined.

Table 1. Vibrational frequencies ^{1, 2} (in cm⁻¹) for some tetrahedral thio and seleno anions.

Ion	$v_1(a_1)$	r,(e)	$v_3(f_2)$	$v_4(f_2)$
	71(41)	2(0)	3 (72)	4 (72)
PS ₄ ³⁻	421	201	547	266
VS_{4}^{3}	375	178	460	178
NbS_4^{3-}	408	163	421	163
TaS_4^{3-}	424	170	399	170
MoS_4^{2-} (Cs – Salt)	460	195	480	195
WS_4^{2-} (Cs-Salt)	485	185	465	185
ReS ₄	501	200	486	200
VSe_4^{3-}	232	121	365	121
NeSe ₄ ³⁻	239	100	316	100
TaSe ₄ 3-	249	103	277	103
$MoSe_2^{2-}$ (NH ₄ -Salt)	255	120	340	120
$WSe_4^{\bar{2}-}$ (Cs-Salt)	281	107	309	107

The fundamental vibrations of $XY_4(T_d)$ type systems are distributed as $a_1+e+2\,f_2$. Wilson's ¹¹ FG matrix method has been used to evaluate the

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force constants. The expressions for the F and G matrices were taken from Krebs and Müller 6 .

Table 2. OVFF and the UBFF constants (in mdyn/Å) for some tetrahedral thio and seleno anions.

Ions	$k_1(K)$	$k_a'(3H)$	A(F/2)	$B/R\left(F^{\prime}\right)$
PS_4^{3-}	1.832	0.475	0.168	-0.018
	(2.03) *	(0.45)*	(0.16)*	(-0.02)*
VS_4^{3-}	2.011	0.537	0.081	-0.099
	(2.020)	(0.471)	(0.079)	(0.032)
$\mathrm{NbS_4^{3^-}}$	2.064	0.271	0.135	-0.039
	(2.089)	(0.246)	(0.132)	(0.007)
TaS_4^{3-}	2.108	0.235	0.161	-0.011
1	(2.142)	(0.216)	(0.157)	(-0.017)
$\mathrm{MoS_4^{2^-}}$	2.713	0.464	0.160	-0.067
	(2.750)	(0.420)	(0.155)	(0.010)
WS_4^{2-}	2.931	0.282	0.189	0.013
	(2.975)	(0.258)	(0.184)	(-0.019)
$\mathrm{ReS_4}^-$	3.232	0.405	0.189	-0.027
	(3.294)	(0.372)	(0.181)	(-0.021)
VSe_4^{3-}	1.819	0.675	0.076	-0.146
	(1.856)	(0.579)	(0.081)	(0.064)
4	2.068	0.399	0.074	-0.081
	(2.065)	(0.348)	(0.074)	(0.032)
4	2.051	0.333	0.104	-0.048
	(2.104)	(0.315)	(0.098)	(0.016)
$\mathrm{MoSe_4^{2^-}}$	2.402	0.639	0.078	-0.124
	(2.373)	(0.546)	(0.082)	(0.039)
$\mathrm{WSe_4^{2^-}}$	2.616	0.324	0.132	-0.055
	(2.638)	(0.291)	(0.130)	(0.016)

^{*} Reference 2.

(The values in parentheses indicate the UBFF constants $K,\,3\,H,\,F/2$ and F'.)

It is evident from Table 2 that the OVFF constants k_1 , k_{α}' and A are comparable with the corresponding UBFF constants \vec{K} , 3H, and F/2. It is worth noting that for the same metal, the metalsulphur bond is stronger than the metal-selenium bond. This conclusion, supported also by the electronegativity differences, shows that as the mass of the peripheral atom increases, the force constant decreases and the bond gets weakened. The variation of the stretching force constant (k_1) in thio anions of different metals indicates that as the mass of the central atom increases the relative strength of the chemical bonds increases. This is also true in the case of seleno anions of different metals. In the iso- $\begin{array}{lll} {\rm electronic\ groups\ viz.\ NbS_4{}^{3^-},\ MoS_4{}^{2^-},\ TaS_4{}^{3^-},} \\ {\rm WS_4{}^{2^-},\ ReS_4{}^-,\ NbSe_4{}^{3^-},\ MoSe_4{}^{2^-},\ and\ TaSe_4{}^{3^-},} \end{array}$ WSe₄²⁻, it is observed that the stretching force constant increases with the increase of the oxidation state. It is, therefore, inferred that the relative



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^{**} Z. Naturforsch. 27 a, 129 [1972].

Notizen 827

strength of chemical bonds increases with the decrease of the formal charge. The interaction constant, A, shows the same trend and supports the above conclusion which indicates that the interaction constant is greater for smaller non-bonded distances. The angle bending force constant k_a decreases with increasing mass of the central atom in the same group only. The present calculation leads to a negative value of B/R for all the ions. It follows that the OVFF model, making use of a single interaction term in B/R, cannot represent satisfactorily the interaction between non-bonded atom pairs. Several other investigators 6, 7, 12, 13 have also reported the negative B/R in the case of ions. Another set of OVFF constants has also been computed assuming the relation A = 6.5 (B/R), but the constants obtained do not lead the satisfactory reproduction of the observed frequencies. This part is not reported here. Krebs and Müller 6 also found the failure of

0.271

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0.141

0.024

0.020

0.015

0.015

0.011

0.016

0.013

0.009

0.010

0.010

0.018

0.015

0.014

0.013

0.018

0.015

0.007

0.012

0.012

0.015

0.013

0.013

0.005

0.014

0.014

0.009

0.008

0.008

0.012

0.011

0.014

0.008

0.008

0.006

-0.359

-0.247

-0.211

-0.223

-0.151

-0.114

-0.089

-0.043

-0.049

-0.148

-0.108

-0.117

-0.085

-0.047

-0.052

-0.087

-0.049

-0.055

-0.283

-0.265

-0.274

-0.181

-0.170

-0.175

-0.095

-0.098

-0.187

-0.168

-0.176

-0.115

-0.094

-0.090

-0.118

0.108

0.479

0.326

0.129

0.068

0.312

-0.349

0.346

0.290

-0.610

-0.506

0.197

0.318

0.165

-0.219

-0.298

-0.293

-0.297

-0.299

0.252

0.223

-0.006

0.123

0.223

0.062

0.002

0.202

0.214

-0.033

0.062

0.203

-0.214

-0.215

-0.188

-0.190

No.

1

2

3

4

5

6

7

8

9

10

11

12

 VS_{4}^{3-}

NbS43-

 TaS_4^{3-}

 MoS_{4}^{2-}

 WS_4^{2-}

ReS₄

VSe₄3-

NbSe₄3-

TaSe₄3-

MoSe₄2-

WSe₄2-

the relation $A=6.5\ (B/R)$ in some ions and concluded that the application of a Lennard-Jones potential resulting in the assumption $A=6.5\ (B/R)$ is not reasonable in the case of ions.

Recently Müller et al. ¹⁰ have studied the variation of the ratio L_{12}/L_{21} with the mass coupling term $(T=G_{12}/V|G|)$ for molecules of the type $XY_4(T_d)$, $XY_3(D_{3h})$ and $XY_2(C_{2v})$. They formulated an empirical constraint for the calculation of force constants (extended L-matrix approximation). From their empirical study, they concluded that an average value of the factor $L_{12}/L_{21}=-0.075$ may be taken as an empirical constraint in fixing the force constants of the second order vibrational eigenvalue problem in $XY_4(T_d)$ type systems. They found that the values of the force constants determined from this empirical approach are in good agreement with the exact force field data. We, therefore, examined the dependence of the ratio L_{12}/L_{21}

Table 3. Elements of L- L_{12} L_{12}/L_{21} Method matrix * (a. m. u.) $^{-1/2}$ for L_{11} L_{21} L_{22} some tetrahedral thio and and seleno anions. PS43-0.027 -0.3620.454 -0.0740.271 Fad. OVFF 0.027 -0.3510.342 -0.0770.271

-0.067

-0.081

-0.071

-0.067 -0.073

-0.148

-0.114

-0.101

-0.233

-0.204

-0.119

-0.139

-0.119

-0.150

-0.382

-0.288

-0.080

-0.241

-0.215

-0.053

-0.049

-0.047

-0.028

-0.082

-0.080

-0.076

-0.084

-0.082

-0.064

-0.065

-0.079

-0.070

-0.089

-0.064

UBFF

OVFF UBFF

Fad.

Fad. OVFF

UBFF

OVFF

UBFF

OVFF UBFF

Fad.

Fad. OVFF

UBFF

Fad.

OVFF

UBFF

OVFF

UBFF

UBFF

OVFF

UBFF

OVFF UBFF

Fad

Fad.

OVFF

UBFF

Fad.

Fad. OVFF

Fad.

Fad.

^{*} Calculated by equations given in Reference 3.

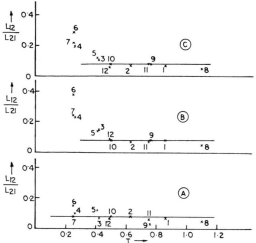


FIG. VARIATION OF $\frac{L_{12}}{L_{21}}$ VS. MASS COUPLING TERM $\left(T = \frac{G_{12}}{\sqrt{|G_{1}|}}\right)$

- $igain \frac{\text{Li2}}{\text{L}_{21}}$ by Fadini's force field contants
- B LIZ BY O.V.F.F. CONTANTS
- $\bigcirc \frac{L_{12}}{L_{21}}$ BY U.B.F.F. CONTANTS

Fig. 1.

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on the mass coupling term, T, in the present case of thio and seleno anions using the force constants calculated by Fadini's method ^{1, 5}, OVFF constants and UBFF constants. It is found from Fig. 1 that the force field data from Fadini's method give an average value of the constraint $L_{12}/L_{21} = -0.08$, which is in good agreement with the value proposed by Müller et al. ¹⁰. The OVFF and UBFF constants also give the same average value of the constraint L_{12}/L_{21} for ions of higher mass coupling terms. However, in the case of ions having lower mass coupling terms, the ratios L_{12}/L_{21} are found to be greater due to the higher F_{12} values from OVFF and UBFF constants. Hence it may be concluded that Fadini's approximation is closer to an exact force field.

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