

A unified phonon dynamical study of europium sulphide (EuS)

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Abstract

A systematic unified theoretical analysis for Europium Sulphide (EuS) have been investigated by a lattice dynamical model which includes the effect of three-body interaction (TBI) in the framework of second neighbour rigid shell model (SNTRSM) and second neighbor rigid ion model (SNTRIM). The significance of these two approaches thus obtained, have been applied to study the elastic constant of first and second order, Cauchy discrepancy, pressure derivatives, anharmonic properties and high pressure phase transition study of Europium sulphide (EuS) by the supplication of TRSM (three-body force shell model) and TRIM (three-body force rigid ion model). It is concluded that our theoretical results predicted by SNTRSM on phonon dynamics and derivable properties will be very much close to their measured data. These are probably the first reports of its kind and they will certainly help the experimental workers to analyze their data in future.

Keywords: phonon dispersion curves, rigid shell model, rigid ion model, debye temperature

Introduction

The electronic structure of Europium sulphide (EuS) which is a family of Europium Chalcogenides (EuO, EuS, EuSe, EuTe) crystallize in f.c.c. NaCl structure and are also called as rare earth europium chalcogenides. Unlike other rare earth compounds europium chalcogenides generally, show non-mixed valance character. The study of lattice dynamical behavior of these chalcogenides is incomplete even today due to insufficient experimental data for phonon dispersion curves and a very few attention has been paid to it.

Although europium chalcogenides have a large application as magnetic semiconductors, yet no serious attention has paid so far. Only few information about the optical frequencies [3-5], elastic properties [2] and magnetic [29, 30] have been presented. Complete experimental data on phonon dispersion is not available for these compounds except for EuSe, for which limited information about phonon frequency has been reported by Silberstein *et al* [1]. Zeyher and Kress have applied a phenomenological model (OSM) [14] to discuss the complete phonon dispersion curves (PDCs) and combined density of states (CDS). Furthermore, Osaka *et al.* [11] have investigated the phonon frequencies only for EuSe using Breathing shell model (BSM). For better result Mischenko and Kikoin [19] have modified Zayher and Kress overlap shell model (OSM) to predict the phonon dispersions curves of EuO and EuS. These authors modified the dynamical matrix by incorporating the charge density deformation effects. But on the basis of overlap achievements, their results are far away from success because none has considered the many body interactions (the first important term is three body interactions) for these compounds. Due to the unfilled 4f shells, the radii of rare earth ion changes and therefore overlapping of the chalcogen ions, also changes. The Europium chalcogenides show deviations from the Cauchy relation $C_{12}=C_{44}$. The BSM used by Onsaka *et al.* [11] and Sakale *et al.* [25] of PDC

only explains the acoustic branches well. Therefore it is evident that OSM and BSM fail to explain the optical branches of PDC of these crystals. It has been found that three body interactions explain well the optical branches and Cauchy discrepancy both simultaneously and successfully to almost all the ionic and semiconducting crystals [17]. In addition, these compounds exhibited strong optical phonon anomalies throughout the Brillouin zone and peculiar acoustic phonon propagation along $[qqq]$ directions. These facts suggest that some types of many-body interactions are responsible for the elastic and phonon anomalies in these compounds. This has motivated the present author to the basic need of two phenomenological lattice dynamical models. The aim of this paper is to test the applicability and utility of second neighbor three-body force rigid shell model (SNTRSM) and second neighbor three-body force rigid ion model (SNTRIM) for the satisfactory description of phonon dispersion relations and other phonon properties of these compounds.

Theory

The general formulation of present Lattice Dynamical model is given by

1. Three-body force rigid shell model (SNTRSM)
2. Three-body force rigid ion model (SNTRIM)

The general formulation of SNTRSM can be derived from the crystal potential whose relevant expression per unit cell is given by

$$\Phi = \Phi^C + \Phi^R + \Phi^{TBI} \dots \dots \dots (1)$$

Where the first two terms represent, respectively, long range Coulomb and three body interactions (TBI) energies. The next term is the form of SR overlap repulsive energy extended to the next nearest neighbor ions. where first term

Φ^C is Coulomb interaction potential which is long-range in nature, second term Φ^R is short range overlap repulsion potential operative up to the second neighbors and third term Φ^{TBI} is three body interaction potential. The secular determinant $D(q)$, is the (6x6) dynamical matrix which is given by:

$$D(q) = (R' + Z_m CZ_m) - (T + S_m CY_m)(S + K + Y_m CY_m)^{-1}(T^* + Y_m CZ_m) \quad \text{----- (2)}$$

The Number of adjustable parameters has largely been reduced by considering the short range interaction to act only through the shells. This assumption leads to $R=T=S$. The expressions derived for elastic constants corresponding to SNTRSM have been obtained as:-

$$\frac{4r_0^4}{e^2} C_{11} = [-5.112Z_m^2 + A_{12} + \frac{1}{2}(A_{11} + A_{22}) + \frac{1}{2}(B_{11} + B_{22}) + 9.3204\xi^{12}] \quad \text{---- (3)}$$

$$\frac{4r_0^4}{e^2} C_{12} = [0.226Z_m^2 - B_{12} + \frac{1}{4}(A_{11} + A_{22}) - \frac{5}{4}(B_{11} + B_{22}) + 9.3204\xi^{12}] \quad \text{----- (4)}$$

$$\frac{4r_0^4}{e^2} C_{44} = [2.556Z_m^2 + B_{12} + \frac{1}{4}(A_{11} + A_{22}) + \frac{3}{4}(B_{11} + B_{22})] \quad \text{----- (5)}$$

In View of the equilibrium condition $[(d\Phi/dr)_0]$ We obtain.

$$B_{11} + B_{22} + B_{22} = -1.165Z_m^2 \quad \text{----- (6)}$$

Where

$$Z_m^2 = Z^2 \left(1 + \frac{12}{Z} f_0\right) \quad \text{and} \quad \xi^{12} = Zr_0 f_0$$

The term f_0 is a function dependent on the overlap integrals of the election wave-funtions and the subscript zero indicates the equilibrium value. By solving the secular equation along $[q00]$ direction and subjection the short and long-range coupling coefficients to the long-wavelength limit $q \rightarrow 0$, two distinct optical vibration frequencies are obtained as:

$$(\mu\omega_L^2)_{q=0} = R_0 \frac{(Z'e)^2}{Vf_r} \cdot \frac{8\pi}{3} (Z_m^2 + 6\xi^{12}) \quad \text{----- (7)}$$

$$(\mu\omega_r^2)_{q=0} = R_0 \frac{(Z'e)^2}{Vf_r} \cdot \frac{4\pi}{3} Z_m^2 \quad \text{----- (8)}$$

Since, in these compounds, $\omega_L = \omega_r$ at Γ - point, therefore, Eqs (7) and (8) lead to the expression:

$$\frac{Z_m^2 + 6\xi^{12}}{\xi^{12}} = -\frac{f_L}{2f_r} \quad \text{----- (9)}$$

Where the abbreviations stand for

$$R_0 = R_0 - e^2 \left(\frac{d_1^2}{\alpha_1} + \frac{d_2^2}{\alpha_2} \right); Z' = Z_m + d_1 - d_2$$

$$f_L = 1 + \left(\frac{\alpha_1 + \alpha_2}{v} \right) \cdot \frac{8\pi}{3} (Z_m^2 + 6\xi^{12})$$

$$f_r = 1 - \left(\frac{\alpha_1 + \alpha_2}{v} \right) \cdot \frac{4\pi}{3} Z_m^2$$

By solving the dynamical matrix along $[0.5, 0.5, 0.5]$ directions at L-Point modified expressions for $\omega_{L_o}(L)$, $\omega_{T_o}(L)$, $\omega_{L_A}(L)$, and $\omega_{T_A}(L)$, are as follows.

$$m_1 \omega_{L_A}^2(L) = R_0 + \frac{e^2}{V} (2A_{11} + B_{11} - \frac{e^2 d_1^2}{\alpha_1} + \left(\frac{e^2}{V} \right) C_{1L} (Z_m + d_1)^2 \left[1 + \left(\frac{\alpha_1}{V} \right) C_{1L} \right]^{-1} \quad \text{----- (10)}$$

$$m_2 \omega_{L_o}^2(L) = R_0 + \frac{e^2}{V} (2A_{22} + B_{22} - \frac{e^2 d_1^2}{\alpha_1} + \left(\frac{e^2}{V} \right) C_{1L} (Z_m + d_2)^2 \left[1 + \left(\frac{\alpha_2}{V} \right) C_{1L} \right]^{-1} \quad \text{----- (11)}$$

$$m_2 \omega_{T_o}^2(L) = R_0 + \frac{e^2}{2V} (2A_{22} + B_{22} - \frac{e^2 d_2^2}{\alpha_2} + \left(\frac{e^2}{V} \right) C_{1T} (Z_m + d_2)^2 \left[1 + \left(\frac{\alpha_2}{V} \right) C_{1T} \right]^{-1} \quad \text{----- (12)}$$

$$m_1 \omega_{T_A}^2(L) = R_0 + \left(\frac{e^2}{2V} \right) (A_{11} + 5B_{11} - \frac{e^2 d_1^2}{\alpha_1} + \frac{e^2}{V} C_{1T} (Z_m + d_1)^2 \left[1 + \left(\frac{\alpha_1}{V} \right) C_{1T} \right]^{-1} \quad \text{----- (13)}$$

Where

$$C'_{1L} = - \left[(C_{1xx} + 2C_{1xy}) + (V_{1xx} + 2V_{1xy}) Z_m^{-2} Zr_0 f_0' \right] 0.5, 0.5, 0.5$$

$$C'_{1T} = - \left[(C_{1xx} + C_{1xy}) + (V_{1xx} + 2V_{1xy}) Z_m^{-2} Zr_0 f_0' \right] 0.5, 0.5, 0.5$$

Where $(C_{1xx} + C_{1xy})$ and $(V_{1xx} + V_{1xy})$ are Coulomb and three - body coupling coefficients evaluated at L-point. polarizability is negligibly small and the negative ion polarizability of nitride ion is almost zero. Therefore, it has been considered to utilize the second neighbour three-body force rigid ion model (SNTRIM) for further calculations of phonon frequencies.

In an attempt to solve the expressions for SNTRIM, all the Eqs (1-6) will remain the same, only the difference is in the expressions from Eqs. (7-13), which can be written as follows:

$$(\mu\omega_L^2)_{q=0} = R_0 + \frac{8\pi e^2}{3V}(Z_m^2 + 6\xi^2) \quad \text{----- (14)}$$

$$(\mu\omega_T^2)_{q=0} = R_0 - \frac{4\pi e^2}{3V}(Z_m^2) \quad \text{----- (15)}$$

Since, in these compounds, $\omega_L = \omega_T$ at Γ - point, therefore, Eqs (14) and (15) lead to the expression:

$$Z_m^2 = -4Zr_0f_0' \quad \text{----- (16)}$$

Again, by solving the dynamical matrix along [0., 0.5, 0.5] directions at L-point, the modified expressions for $\omega_{Lo}(L)$, $\omega_{To}(L)$, $\omega_{LA}(L)$, and $\omega_{TA}(L)$ are derived as follows:

$$\omega_{Lo}(L) = R_0 + \frac{e^2}{v}(2A_{22} + B_{22}) + \frac{e^2}{v}C'_{1T}Z_m^2 \quad \text{---- (17)}$$

$$m_2\omega_{To}^2(L) = R_0 + \frac{e^2}{v}(A_{22} + 5B_{22}) + \frac{e^2}{v}C'_{1T}Z_m^2 \quad \text{--- (18)}$$

$$m_1\omega_{LA}^2(L) = R_0 + \frac{e^2}{v}(2A_{11} + B_{11}) + \frac{e^2}{v}C'_{1L}Z_m^2 \quad \text{--- (19)}$$

$$m_1\omega_{TA}^2(L) = R_0 + \frac{e^2}{v}(A_{11} + 5B_{11}) + \frac{e^2}{v}C'_{1T}Z_m^2 \quad \text{--- (20)}$$

Where R_0 and C'_{1T} have already been defined.

Computations and Results

The input data along with their relevant references and calculated model parameters from SMTRSM and SNTRIM for EuS are given in Table-1. A comparative results on phonon dispersions curves from the two models have been shown in Figure-1. These results have also been compared with the observed data of Kress *et al* [1], for visual comparison.

Table 1- Input data and model parameters for EuS

C_{ij} (in 10^{12} dyne cm^{-2}), v (in THz), r_0 (in 10^{-8} cm) and α_i (in 10^{-24} cm^3)

Table 1

Input Data		Model Parameters		
Properties	Values	Parameters	TRSM	TRIM
C ₁₁	13.1	Z _m ²	0.6745	0.6522
C ₁₂	1.10	rof ₀	-0.3005	-0.3005
C ₄₄	2.73	A	8.5110	8.396
v _{Lo} (Γ)	8.01	B	-0.7858	-0.7598
v _{To} (Γ)	5.35	d ₁	0.01634	--
r ₀	2.984	d ₂	1.4081	--
α_1	1.5	Y ₁	-0.7943	--
α_2	5.5	Y ₂	-2.1497	--

* Value extrapolated from measured PDC

** Reasonable value taken from ionic radii.

Table 2

Raman				IR Raman		
C.D.S. v (Cm ⁻¹)	Experimental	Assignment	Value (cm ⁻¹)	Experimental	Assignment	Value (cm ⁻¹)
53	-	LA-TA(X)	54	-	-	-
90	-	LA-TA(L)	92	-	2TA(Δ)	100
128	-	TO-TA(X)	129	-	-	-
155	-	LA+TA(Δ)	154	-	LA+TA(Δ)	154
184	-	2TA(L)	183	-	TO+LA(Δ)	191
208	-	2LA(Δ)	208	-	2LA(Δ)	208
225	-	LA+TA(l)	225	-	-	-
262	-	TO+TA(X)	262	-	TO+TA(L)	258
290	-	TO+LA(Δ) LO+TA(Δ) LO+TA(Δ)	292	-	LO+TA(Δ)	292
317	-	TO+LA(X)	316	-	TO+LA(Δ)	300
357	-	2TO(Γ)	357	-	LO+TA(L)	337
412	-	LO+TO(L)	412	-	LO+TO(Δ)	429
450	-	2LO(X)	450	-	-	-
483	-	2LO(Δ)	483	-	2LO(Δ)	483
492	-	2LO(L)	491	-	-	-

Discussion and Conclusion

From figure 1, it is clear that the results reported from SNTRSM for EuS. EuS are comparatively more close to ensured data on PDCs. There are certain features in PDC of EuS which deserve special mention. The three body interactions have influenced both LO and TO branches much more than acoustic braches (LA and TA). Another striking feature of the present study is noteworthy from the excellent reproduction of optical and acoustic branches.

The model parameters of TRSM and TRIM have been used to calculate the phonon spectra for allowed 48 non-equivalent wave vector in first Brillouin zone. The frequency along with symmetry directions have been plotted against the wave vectors to obtain the phonon dispersion curves (PDCs) from both the models. These curves are compared with each other and with inelastic neutron scattering technique.

Since neutron scattering provide us only a very little data for

symmetry direction, we have studied the specific heat for complete description of frequencies. For this purpose the specific heat has been computed at different temperature using Blackmann's technique [35] and corresponding Debye temperature have been plotted against absolute temperature (T). It may be concluded that SNTRSM provides agreement which is certainly better than those fitted by experimental researchers and SNTRIM, are very much close to the experimental values. Although, qualitatively the agreement achieved from our present model SNTRSM is comparatively better than some of the model values. In addition, some other researchers [18-24] of the field has also tried their best to explain PDCs and other properties of europium chalcogenides but only with moderate success.

Furthermore, in order to increase the merit of this work, we have tested the adequacy of our model by calculating [33] two phonon Raman/IR spectra shown in Table-2 and variation of Debye temperatures and the Combined density of stated (CDS) peaks are shown in figure-2 and figure-3. In order to interpret them the critical point analysis has been used following the method prescribed by Burstein *et al* [36]. It may be concluded that the inclusion of the effect of short range overlap repulsive interaction upto second neighbours in the framework of TRIM and TRSM is important in EuS. The present approach has revealed much better description of the crystal dynamics of the solid than those reported [1] by other models. Furthermore, the inherent shortcomings of the present models are most likely the same as the demerits of models RIM and RSM. It is expected that slight discrepancies still occurring between theory and experiment may be further improved by including the effect of free carrier screening (FCS), Van der Waals interactions (if data are available in future) and by including anharmonicity of vibrations in the present model (SNTRSM).

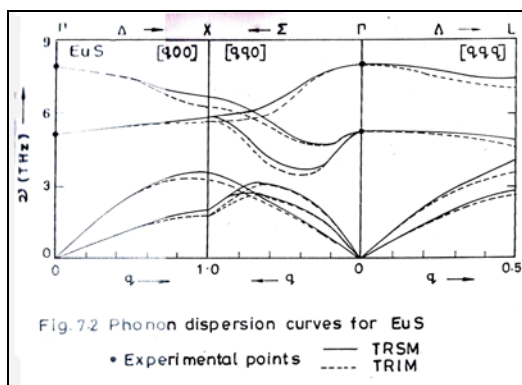


Fig 1

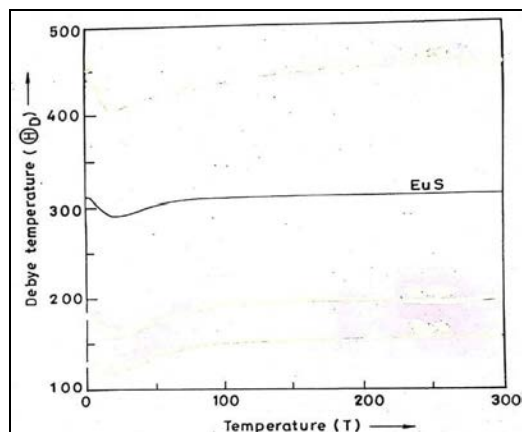


Fig 2: Debye temperature variations of EuS

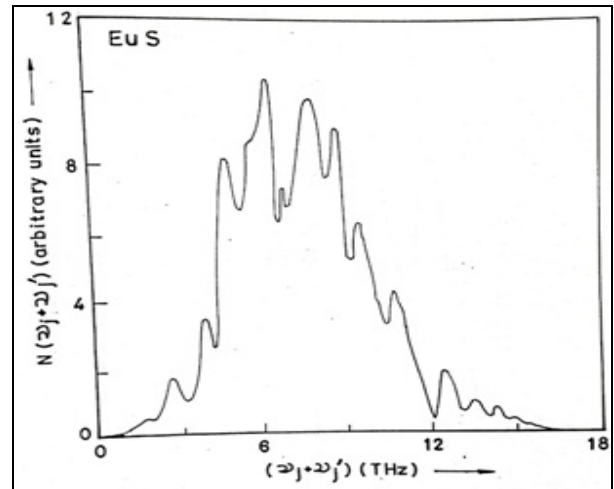


Fig 3: The Combined density states (CDS) peaks of EuS

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