A STUDY OF THERMOPHYSICAL AND COHESIVE PROPERTIES OF $LiCl_{1-x}I_x$ MIXED ALKALI HALIDE

A.K. DIXIT

Professor of Physics, Govt. P.G. College, Satna (M.P.), India

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I have studied the thermophysical and cohesive properties such as Gruneisen parameter, molecular force constant, Restrahl frequency, Debye temperature compressibility and Anderson-Gruneisen parameter etc of mixed LiCl_{1-x}l_x mixed alkali halide under the banner of three body force potential model .Three body force potential (TBFP) model contains long-range forces (coloumbian and three body interactions) as well as short range forces (vander-Waal's and overlap repulsive forces) [10-15] . I have selected mixed alkali halides for the discussion because mixed crystals are very useful and known as solid solutions. Mixed alkali halides have received considerable attention of investigators [1-9] due to their scientific and technological importance. The observations show that the micro hardness of mixed alkali halides is larger than the pure crystals. Some results are very close to their experimental value which shows the superiority of this model over other models.

Introduction

Donic solids are model crystal for the analysis of lattice properties. In the same sprit we have used our three body force potential model (TBFP) for the calculation of lattice and thermo-physical properties of mixed alkali halides $\operatorname{LiCl}_{1-x}I_x$. This potential model consists long-range (coloumbian and three body interaction) forces as well as short range forces (overlap repulsive force operating upto next nearest neighbour ions and vander-Waal's interaction) [25-30]. This shows that the inclusion of three body interaction effect makes the present model suitable for the study of thermophysical properties of mixed alkali halides. Most of the theoretical and experimental workers [10-36] have explained the elastic and thermophysical properties of the ionic solids.

Theory and method of Calculations:

From the early history of science and its evolution attempts have been made to explain the behavior of the ionic solids under some boundary conditions through some potential models. We have applied the TBP model for the calculation of thermophysical properties of mixed alkali halides. Three body potential model contains long-range forces as well as short-range forces. Coloumbian force associated with three body force is long-range force while vander-Waal's and Hafemeister – Flygare are short-range forces. The values of input data for different concentrations (x) have been evaluated by applying Vegard's law for second order elastic constants, lattice constants, ionic radii and vander-Waal's constants of host and dopant materials as-

$$C_{ij}(\text{mix}) = x c_{ij}(A) + (1 - x) c_{ij}(B)$$

 $r_{ij}(\text{mix}) = x r_{ij}(A) + (1 - x) r_{ij}(B)$
 $\rho(\text{mix}) = x \rho(A) + (1 - x) \rho(B)$
 $C(\text{mix}) = x C(A) + (1 - x) C(B)$
 $D(\text{mix}) = x D(A) + (1 - x) D(B)$

The constituent of mixed crystals are held together by the harmonic elastic forces with no internal stress within the crystal. The values of input data are given in table 1.1. Thermophysical properties included in the discussion are Gruneisen parameter (γ), molecular force constant (f) Debye Temperature (θ_D), Restrahl frequency (ν_D), ratio of volume thermal expansion coeffcient to specific heat at constant volume ($\alpha_v/C\nu$), Compressibility (β), Anderson-Gruneisen parameter (δ_T). The relevent expressions have been derived from three body potential model [25-36].

U (r) =
$$-\alpha_m z e^2/r [z + 12 f(r)] + 6b\beta_+ - \exp[(r_1 + r_2 - r)/\rho]$$

+ $6b\beta_+ \exp[(2r_1 - 1.4142r)/\rho]$
+ $6b\beta_- \exp[(2r_2 - 1.4142r)/\rho] - C/r^6 - D/r^8$

(i) Gruneisen parameter

$$\gamma = -r_0/6 [U'''(r)/U''(r)] r = ro$$

(ii) Molecular force constant

$$f = 1/3 [U']_{SR}(r) + 2/r U_{SR}(r)]_{r=ro}$$

(iii) Restrahl frequency

$$υ$$
o = $1/2π\sqrt{f/μ}$

where μ is the reduced mass of the crystal.

(iv) Debye Temperature.

$$(\Theta_D) = h \upsilon o/k$$

(v) Ratio of volume thermal expression coefficient to specific heat at constant volume

$$\alpha_{v}/C_{v} = -U^{"}(r_{0})/2r_{0}U^{"}(r_{0})^{2}$$

- (vi) Compressibility $\beta = 18r_0/U''$ (r_0)
- (vii) The Anderson-Gruneisen Parameter

$$\delta_T = -\beta r_0^3 / 27 V U'''(r_0)$$

The above relations have been used to calculate the thermophysical properties of mixed halides. The model parameters are given in table 1.1 and calculated thermophysical properties of mixed alkali halide are given in table 1.2.

Table 1.1. Model parameters for mixed alkali halide $LiCl_{1-x}I_x$ at different compositions

Composition	ρ (10 ⁻⁸ cm)	b	$f(r_0)$	$af^{\prime}(r_0)$	С	$(10^{-76}\mathrm{erg}\times\mathrm{cm}^8)$		
(x)		$(10^{-12} ergs)$			$(10^{-60} \operatorname{erg} \times \operatorname{cm}^6)$			
0.0 (LiCl)	0.399	0.408	0.0038	- 0.048	77	78		
0.2	0.407	0.399	0.00386	- 0.044	314.6	179.2		
0.4	0.416	0.391	0.00392	- 0.040	552.2	280.4		
0.6	0.425	0.382	0.00398	- 0.036	789.8	381.6		

0.8	0.433	0.374	0.00404 -	- 0.032	1027.4	482.8
1.0 (LiI)	0.442	0.365	0.0041 -	- 0.028	1265	584

Table 1.2. Thermophysical & cohesive properties of mixed halide $LiCl_{1-x}I_x$ at different compositions

	compositions											
Cohesive Property	X=0 $X=0$		0.2 X = 0.4		X = 0.6		X = 0.8		X = 1.0			
	Calc	Exp.	Calc	Exp.	Calc	Exp.	Calc	Exp.	Calc	Exp.	Calc	Exp.
Gruneisen Prarmeter γ	1.67	1.78 ^C	1.7047	_	1.738	_	1.772	_	1.806	_	1.84	1.9^{25}
Molecular force constant $f (10^4 \text{ dyn/cm})$	6.72	Ι	6.382	-	6.044	_	5.706	_	5.638	-	5.03	3.32^{28}
Restrahl frequency υο (<i>T</i> Hertz)	13.92	Ι	13.356	-	12.792	_	12.23	_	11.66	-	11.1	_
Debye Temperature (Θ_D) (k)	658.28	-	628.22	-	598.17	_	568.11	_	538.08	_	508	_
Ratio of volume thermal expression coefficient to specific heat at cons tant volume α_v/C_v (10 ¹⁰ ergs unit)	4.66	ı	4.78	ı	4.896		5.014	ı	5.132	_	5.25	ı
Compressibility (β) (10 ⁻¹² dyn/cm)	2.91	3.36 ^K	3.256	_	3.602	-	3.948	_	4.294	_	4.64	5.83 ^K
The Anderson-Gruneisen Prarmeter δ_T	3.25	-	3.338	_	3.426	_	3.514	_	3.602	_	3.69	_

Result and discussion

look at the table 1.2, it is clear that presently estimated end point values of Gruneisen parameter and Anderson-Gruneisen parameter are very close to their experimental values. The end point values of Debye temperature show a good agreement with their available experimental values (21-29). Some results of ours are still higher than the experimental values. The reason behind is to change in bond length in ionic solids. The observations show that the micro hardness of mixed alkali halides is larger than the pure crystals. Some results are very close to their experimental value which shows the superiority of this model over other models. On the basis of over all achievements the present TBP model can be regarded to be adequately suitable for the prediction of thermo physical properties of mixed alkali halides.

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