Growth And Characterization Of Semi-Organic Nlo Material: L-Valine Potassium Nitrate And L-Valine Lithium Nitrate

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Abstract: Single crystals of semi organic non-linear L-Valine potassium nitrate and L-Valine lithium nitrate grown by slow evaporation method using water as a solvent. The L-Valine phase was confirmed by single crystal powder X-ray diffraction analysis. Presence of various functional groups of L-Valine was characterized by Fourier transform infra-red spectrum (FT-IR) and non-linear optical property is examined by Kurtz powder technique. The optical behavior was analysed by Ultra violet –vis spectrum and found that the crystal is transparent in the region between the 200-1100nm. Hence it may be very much useful for the second harmonic generation (SHG) applications. The crystal was thermally stable up to 215°C (VPN) and 235°C (VLN) as determined by DSC-TGA studies and mechanical stabilities of crystal have been confirmed by Vicker's microhardness study. Dielectric constant was measured with various frequencies as a function of temperature.

Keywords: L-Valine, Second harmonic generation, Micro hardness, dielectric studies, thermal analysis

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I. Introduction

Second order non-linear optical (SONLO) materials have recently attracted much attention because of their potential applications in emerging optoelectronics technologies [1,2]. Materials with large second order optical non-linearity, short transparency cutoff wavelengths, and stable physico-thermal performance are needed to realize many of these applications. The search for new frequency conversion materials over the past decade has concentrated primarily on organics. It has been demonstrated that organic crystals can have very large non-linear susceptibilities compared with inorganic crystals, but they used in low optical transparency, poor mechanical properties, low laser damage threshold, and the unable to produce and process large crystals [3, 4]. Purely inorganic nonlinear optical (NLO) materials typically have excellent mechanical and thermal properties with relatively modest optical nonlinearities because of the lack of extended π -electron delocalization. In semi-organics, polarizable organic molecules are stochiometrically bound within an organic host [5]. In recent years, the NLO properties of semi-organic complex products has attracted great interest because of these metal-organic complexes

Here, L-valine is a branched chain amino acid, which has both a primary amino group and a primary carboxylate group. The carboxlate acid group donates its proton to the amino group. So in solid state, amino acid exists as zwitterions, which create hydrogen bonds, in the form of $N-H^+-O-C$, which are very strong bonds. Hydrogen bonds have also been used in the possible generation of non Centro-symmetric structures, which is a prerequisite for an effective SHG crystal.

This paper describes the synthesis of crystal structure of L-valine potassium nitrate, L-Valine Lithium nitrate. The grown crystals were characterized by powder XRD,FTIR,optical transmission measurement, DSC–TGA, dielectric measurement, microhardness measurement and Kurtz and Perry powder SHG test was performed to confirm the second order nonlinearity of the grown crystal.

II. Experimental

Synthesis and crystal growth

L-valine, potassium nitrate and lithium nitrate was received from Sisco Research Laboratories PVT. Ltd (India). This is a the long recrystallization processes and available raw material is used one after purification. L-valine and potassium nitrate is taken in a particular molar ratio and added 20 ml of double distilled water and stirring till dissolved the mixture. After dissolving the above mixture is transferred in to 100 ml beaker. The mixing solution kept in slow evaporation in beakers covered with aluminum foil sheet at room temperature. The L-Valine and lithium nitrate followed by the same procedure. Two grown crystal are colourless, good transparent crystals has obtained by 3 to 4 weeks

Chemical reaction

$$C_5H_{11}NO_2 + KNO_3$$
-----> $C_5H_{11}NO_2$. KNO_3
 $C_5H_{11}NO_2 + Li (NO_3)_2$ -----> $C_5H_{11}NO_2$. $Li(NO_3)_3$

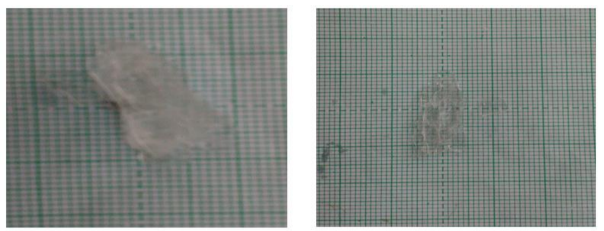


Fig.1 As grown the crystals of VPN and VLN

III. 3. Results And Discussions

3. 1. Powder X-ray diffraction

The grown single crystal of L-Valinepotassium nitrate and L-Valine Lithium nitrate has been subjected to powder X-ray diffraction. Powder form of the above mentioned crystal is taken for the analysis using XPERT PRO diffract meter. The indexed powder x-ray diffraction pattern of the grown crystal is presented in fig 2&3. The lattice parameters for L-Valine potassium nitrate obtained from the data of powder XRD pattern using UNITCELL software package are a = 9.788 Å, b = 6.532 Å and c = 12.00372 ÅCellvol=436.11ų. The lattice parameters for L-Valine Lithium nitrate obtained from the data of powder XRD pattern using UNITCELL software package are and a = 9.890 Å, b = 6.788 Å and c = 12.00222ÅCell vol= 438 ų are found to be in good agreement with the literature

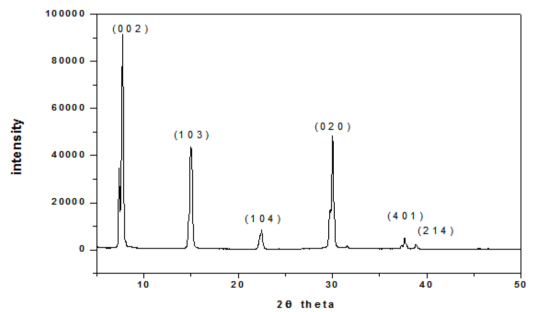


Fig. 2 XRD patteren of L-Valine with Potassium nitrate

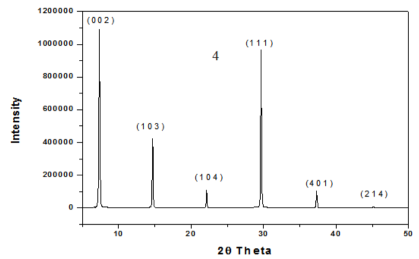


Fig. 3 XRD patteren L-Valine with Lithium nitrate

Table 1 for	XRD	Values of	VPN and	VLN	crystals

ND values of virt and virt crystals				
Parameters(VPN)	Present study	Reported velues		
A	9.788	9.9714		
В	6.532	6.2930		
С	12.000372	12.6480		
V	436.11	434.22		
System	Monoclinic	Monoclinic		
Space group		P2		
Parameters(VLN)	Present study	Reported velues		
A	9.890	9.9714		
В	6.788	6.2930		
С	12.002223	12.6480		
V	438.00	434.22		
System	Monoclinic	Monoclinic		
Space group		P2		

3.2. Optical transmission spectra

A transmission spectrum is very important for any NLO materials, because a nonlinear optical material can be of any practical use if it has a wide transparency window. In the present study, we have recorded the UV-Vis NIR transmission spectrum in the range of 200nm-1100nm is shown in fig 4&5and the instrument used in the analysis is LAMBDA-35 UV-Vis spectrophotometer. From the spectrum, it is seen that the crystal has a lower cut-offwavelength of 272nm.and 280 nm(VPN). The spectrum further indicates that the crystal has a wide optical window from 272nm to 1100nm and 280nm to 1100 nm (VLN). The crystal is transparent in the visible and infrared spectral regions. Optical transmittance of about 100% is observed for 1.5mm plates of L-Valinepotassium nitrate and L-Valine Lithium nitratecrystals is sufficiently good for SHG.

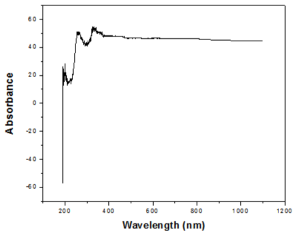


Fig. 4 UV spectrum for L-Valine with potassium nitrate

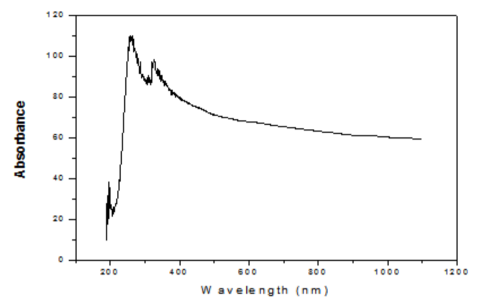


Fig. 5 UV spectrum for L-Valine with lithium nitrate

3.3. FTIR spectral analysis

The FTIR spectrum of VPN and VLN crystals were recorded in the range 400-4000cm-1 employing a Perkin-Elmer spectrometer by KBr pellet method to study the metal organic coordination. Fig.6&7 shows the recorded FTIR spectrum of the grown crystal of VPN and VLN. The vibrational frequency of various functional groups of VPN and VLN and the tentative frequency assignment are presented in table.

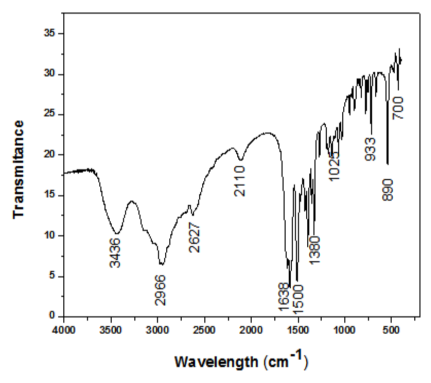


Fig. 6 FT-IR analysis of L-Valine with Potassium nitrate

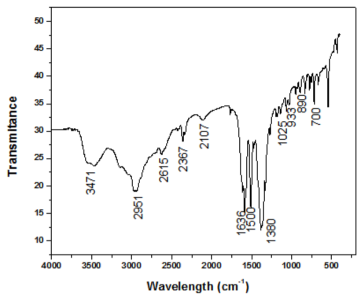


Fig. 7 FT-IR analysis of L-Valine with Lithium nitrate

Table: 2 Comparison IR frequencies of VPN and VLN

IR peak	Assianments	Frequencies		
	Assignments	VPN	VLN	
1	NH ₃ asymmetric stretching	1636	1630	
2	NH ₃ symmetric stretching	1500	1450	
3	CH ₃ symmetric stretching	1380	1360	
4	(CH ₃) ₂ symmetricstretching	1025	1000	
5	C-C symmetric stretching	933	940	
6	C-C-N symmetric stretching	890	870	
7	CH₃ rocking	700	720	

3.4. Second Harmonic Generation Test

The second harmonic generation (SHG) test on the VLN and VPNcrystal was performed by Kurtz powder SHG method [18]. The powdered sample of crystal was illuminated using the fundamental beam of 1064 nm from Q-switched Nd:YAG laser. Pulse energy 4ml/pulse and pulse width of 6 ns and repetition rate of 10Hz were used. The second harmonic signal generated in the crystalline sample was confirmed from the emission of green radiation of wavelength 532 nm collected a monochromator after separating the 1064 nm pump beam with an IR-blocking filter. A photomultiplier tube is used as a detector. It is observed that the measured SHG efficiency of VLN crystal was 0.5 times and VPN was 0.6times that of potassium dihydrogen phosphate (KDP).

Sl. No.	Code / Name of the Sample	Output Energy (milli joule)	Input Energy (joule)
1	L-valine+Lithium nitrate	4.54	0.701
2	L-valine+potassium nitrate	5.08	0.701
3	KDP (Reference)	8.91	0.701

3.5 TGA and DTA analysis

To find the thermal characteristics of LVPN&LVLN, differential analysis (DSC) and thermogravimetric analysis (TGA) were carried out simultaneously in a TA Instruments Q 600 SDT DSC: Simultaneous Thermal Analyzer. The sample was heated at a rate of 10°C/min in protected nitrogen gas flow and 1.25 mg of the sample was taken to carry out the experiment. Fig.8&9shows the thermograms illustrating simultaneously recorded TGA and DTA. From fig 8 DTA curve, it is observed that the material undergoes an irreversible exothermic transition at about 215°C where the decomposition starts, which indicate the material stable up to 215°C. The material is fully decomposed above 600°C. The sharpness of the exothermic peak shows good degree of crystallinity of the grown LVPN crystal. From TGA curve the weight loss curve is observed starts at 145°C and ends at 230°C. This weight loss is due to the liberation of volatile substances. The peak at 230°C indicates a phase change from liquid to vapor state as evidence from the loss of weight in the TGA curve.

From fig 9 DTA curve, it is observed that the material undergoes an irreversible exothermic transition at about 235°C where the decomposition starts, which indicate the material stable up to 235°C. The material is fully decomposed above 600°C. The sharpness of the exothermic peak shows good degree of crystallinity of the grown LVLN crystal. From TGA curve the weight loss curve is observed starts at 145°C and ends at 255°C. This weight loss is due to the liberation of volatile substances. The peak at 255°C indicates a phase change from liquid to vapor state as evidence from the loss of weight in the TGA curve.

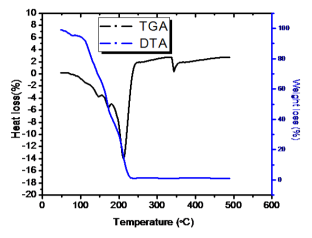


Fig -8 TGA/DTA Curve of L-Valine Potassium nitrate

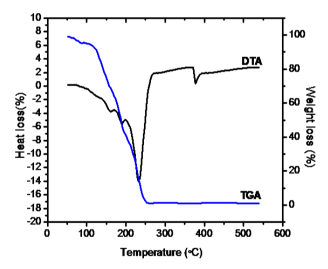


Fig -9 TGA/DTA Curve of L-Valine Lithium nitrate

3.6 Micro Hardness

The mechanical strength of the grown crystal was studied using HMV 2T, Vicker'smicrohardness tester. Microhardness measurement is commonly used to determine the mechanical strength of the material which is related to bond strength and defect structure [11]. The static indentations were made on the surface of crystal by varying the load from 5-100g at room temperature. Vicker'smicrohardness number was determined using H_v =1.8544 $P/d^2kg/mm^2$. The variation of H_v with the applied load P is shown in Fig. 10. In our case, Hv increases with load up to 75g and becomes load independent for P_z 75 g, which can be attributed to the work hardening of the surface and above 75g load significant cracking occurs, which may be due to the release of internal stresses generated with indentation. Finally the maximum value of hardness for LVPN&LVLN crystal at room temperature was found to be $kg72.4~kg/mm^2(LVPN)$ and $81.6~kg/mm^2(LVLN)$ for theload of 75g.

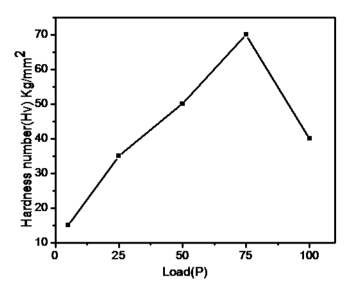


Fig -10 Micro hardness of L-Valine Potassium Nitrate crystal

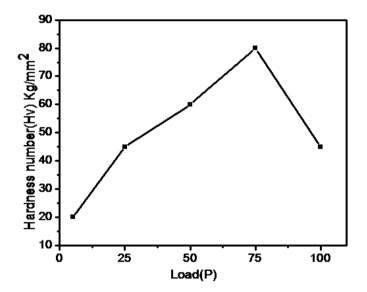


Fig -11 Micro hardness of L-Valine Lithium Nitrate crystal

3.7 Dielectric Constant (E_r)

The dielectric property of LVPN was studied at various temperatures using Agilent A 2484. The dielectric Constant(\mathcal{E}_r) of crystal was found by measuring the capacitance and dielectric loss, which is used to calculate the dielectric constant at various temperatures ranging between room temperature to 150°C for three different frequencies(100Hz, 10KHz and 1MHz). From the figure the dielectric constant increased with increased the temperature⁷. The current investigations showed that dielectric constant was observed maximum at 150°C, since all types of polarization such as electronic, ionic, orientation and space charge polarizations occur at higher temperature. The variation of dielectric constant with temperature at three different frequencies like 100Hz, 10 KHz and 1 MHz is shown in Fig.12&13.

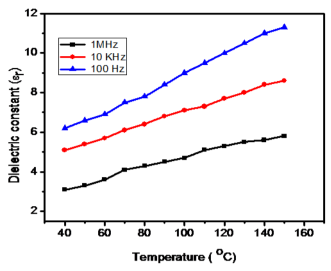


Fig.12 Dielectric studies for L-Valine Pottasium nitrate

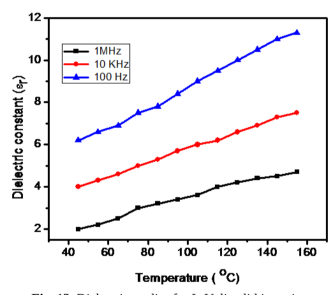


Fig.13 Dielectric studies for L-Valine lithium nitrate

IV. Conclusion

Single crystals of L- valine potassium nitrate and L- valine lithium nitrate were successfullySynthesized by solution growth technique. Its lattice dimensionshave been determined from the powder X-ray diffraction analysis. The various functional groups have been identified from the Fourier transforminfra-red (FT-IR) analysis. The grown crystal has good transmission window in thevisible region between(270 and 280) to 1100 nm, it is suitable for NLO applications. The thermal studies confirm that the crystal structure for LVPN and LVLN is stable up to 215°C& 235°C and indicate its suitability for application in lasers field. Microhardness value was calculated in order to understand the mechanical stability of the grown crystals. From the dielectric studies it is seen that the dielectric constant increased with increased temperature. The powder second harmonic generation efficiency measurement shows the grown VLN and VPN crystal having 0.5 and 0.6 times higher nonlinear optical efficiency than potassium dihydrogen phosphate.

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