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THE STUDY OF THE INTERACTION OF INTENSE PICOSECOND LIGHT PULSE WITH MATERIALS (1) MEASUREMENT OF PICOSECOND PULSE WIDTH USING TWO-PHOTON CONDUCTIVITY IN GAAS. (2) THREE PHOTON CONDUCTIVITY IN CDS

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Maryland University College Park, Maryland

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TECHNICAL RESEARCH REPORT

# THE STUDY OF THE INTERACTION OF INTENSE PICOSECOND LIGHT PULSE WITH MATERIALS

A QUARTERLY TECHNICAL REPORT

SUBMITTED TO

THE U.S. ARMY RESEARCH OFFICE

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Two-photon conductivity effect in GaAs has been utilized to map the					
second order intensity correlation function of the picosecond laser					
pulse for the first time. Reproducible data were obtained. The					
This is limited by the thickness of the					
sample. In principle this method can be extended to use very thin					
sample to obtained subpicosecond time resolution.					
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Mr. S. Jayaraman and Mr. V. Bhanthumnavin

"The Study of the Interaction of Intense Picosecond Light Pulses with Materials"

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#### THE STUDY OF THE INTERACTION OF INTENSE PICOSECOND LIGHT PULSE WITH MATERIALS

- I. Measurement of picosecond pulse width using two-photon conductivity in GaAs.
- 2. Three photon conductivity in CdS.

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## I. <u>Measurement of picosecond pulse width using two photon conductivity in</u> GaAs:

The two photon conductivity in GaAs (Cr-doped high resistivity type) was investigated using a Nd: glass mode locked laser and was reported in the earlier publication<sup>(1)</sup>. The photo conductivity versus laser intensity in a log-log scale is shown in Fig. 1. At lower light intensities the photo conductivity shows a square law dependence on intensity and changes to a linear and sublimear dependence because of the thickness of the material and stimulated intraband absorption. The square law dependence of the photo conductivity on intensity could in principle be utilized to map the 2nd order correlation function of the intensity of the laser pulse. This will give us a measure of the pulse width.

The experimental set up used in the preset experiment is shown in Fig. 2. ANd: glass laser was mode-locked using Kodak 9860 dye in dichloroethane solution. The cavity length was adjusted to give a mode locked pulse train width 4 nanoseconds period. The mode locked pulse train was partially reflected by a plane glass beam splitter onto an ITT photodiode the output of which was monitored on a 519 oscilloscope. Another beam splitter reflects part of a beam on a reference GaAs crystal through ND filters. The transmitted beam was attenuated by ND filters and then split into two equal perpendicular components by a 50% - 50% dielectric beam splitter. The two beams were then made to collide on a GaAs crystal using two 99% dielectric reflectors. Both the reference and signal GaAs crystals were cleaved from a Cr doped high resistivity GaAs wafer of thickness. 35 mm. Indium solder was alloyed to the end faces of the crystals and ohmic contact was thus established. Both ends of the crystals were connected in

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series with  $1200\Omega$  resistor through a 22.5 volts battery. The voltage developed across the resistance was monitored in a dual beam oscilloscope.

The mode locked train of pulses was monitored on a 519 oscilloscope. Almost 80% of the shots gave a single neat mode locked pulse train probably due to the contact dye cell used in the experiment. First, the slope two region of the two GaAs samples were confirmed. The ND-filters were adjusted to keep the GaAs samples well inside the slope two region.

The photoconductivity  $\Delta G$  was calculated from the voltage " $\vartheta$ " across the resistor R (1200 $\Omega$ ).

$$\Delta G = \frac{\theta}{(v-\theta)} \times \frac{1}{R}$$

where V = 22.5 volts

 $\mathbf{R} = 1200 \ \Omega$ 

In the slope two region,  $\theta$  was of the order of .02 volts.

Since 3 < < V,  $\Delta G$  is proportional to V and so the voltage  $\theta$  measured on the oscilloscope can be taken as a measure of the two photon conductivity (TPC). The crystal GaAs #1 monitors the photoconductivity produced by the overlap of the pulse with itself while the crystal GaAs #2 monitors the p. c. due to a single passage of the short pulses, thereby providing the usual reference signal. The TPC pattern is scanned by moving the crystal #1 along the direction  $M_1$  to  $M_2$  and plotting the ratio of the pulse height from Sample #2 as a function of distance. The result is shown in the figure 3. Ratios obtained have been normalized to 1 in the wings to conform with units defined in Ref (2) where T P C yield due to a single pass was assigned a value of 1/2. Only data points for which a single neat mode locked pulse train was monitored on the 519 scope

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were plotted on the graph. Each data point was an average of 8 to 10 shots.

The contrast ratio for two photon conductivity is the same as for the TPF experiment. To effect a comparison, the two photon fluoresence as obtained by Dugnay et al (3, 4) for a modelocked and a free running laser is also drawn on the same figure 3.

In the TPC measurement, the points more or less follows the TPF curve for a mode-locked laser not that of a free-running or Q-switched laser as can be seen in Fig. 3. However, we get a contrast ratio of 1.75 only. This is because of the poor resolution of the crystal. GaAs used in this particular experiment is of thickness 0.35 mm and this correspondes to a resolution of  $\sim 4 \text{ p}$  sec.  $(\frac{t}{c} \times n, n = 3.4, c = 3 \times 10^{-10} \text{ cm/sec})$ . If we sample and integrate the Dugnay's curve over 4 p secs at various points of the curve, we would expect the TPC or TPF curve with ~1.8 contrast ratio.

Thus we show that the two photon conductivity can be used to me asure the width of the picosecond pulses. Thin film crystals of GaAs or some other thin photoconductors (e. g Cd  $S_x - Se_{1 - x}$ ) are suggested for future experiments to measure picosecond pulses. The crystals like Cd  $S_x - Se_1$  have lower two photon absorption cross section and hence will insure a better square law region over a wide range of intensities. This elimiates the narrow square law region of GaAs. To improve the resolution, it is required to perform the experiment with thin film crystals. In conclusion, we state that we measured the auto correlation of the pulse intensity using two photon conductivity in GaAs which yielded the same shape of the curve as got by Dugnay but with a reduced contrast ration of 1.75 because of limited resolution of the detector.

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#### II. Three photon conductivity in Cd S: -

The availability of powerful sources of optical radiation had made it possible to perform a variety of experiments involving many-photon transitions in the optical range of wavelengths. The two-photon absorption process had been observed in cadmium sulfide (5), in an investigation of the recombination radiation excited with light from a q-switched ruby laser. In a different investi-, the two-photon absorption coefficient was measured. B. M. Ashkinadye gation reported an investigation of two-photon conductivity in Cd S at room et al temperature, excited with giant pulses from a ruby laser. However, the three photon process in Cd S with the use of Nd: glass laser was experimentally observed detected the luminescence emitted by only recently. B. M. Ashkinadge et al by Cd S at 77° K excited by neodymium laser. Cd S has a forbidded band width of of 2.57 ev at 77° K, the light of a Nd: glass laser (1.17 ev) should produce the three quantum absorption. They detected the luminescence recombination • radiation) at 5200 A and found that the luminescent intensity depended on the excitation intensity as  $I_{lum} = I_{excitation}^{3.4}$ . They observed the luminescence when the excitation intensity was varied between 20 MW/cm<sup>2</sup> and 100 MW/cm<sup>2</sup>. Since they were using a q-switched laser, they have to focus it to get higher intensities. With the availability of mode locked Nd:glass lasers, it is possible to produce up to a few gega watts  $/_{cm}^2$  without focussing. Further because of the high peak power, the absorption process can be easily observed in the case of picosecond pulse excitation. Arsene et al (9) used the three photon absorption process in Cd S to estimate indirectly the picosecond pulse width from a measurement of the decay of luminescence along the length of the crystal. We investigated the three-photon conductivity in Cd S at room temperature using a mode locked

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Nd: glass laser. The photo conductive cell has a dark resistance of well over a meg ohm. Since the Cd S cell is sensitive to ordinary light, we enclosed the cell in a box and cut off the light from the flash lamp and other sources using optical filters which cut off light other than 1.06 micron laser source. The cell was connected to a 22.5 volts battery through a 127  $\Omega$  resistor. The voltage across the resistance was monitored and measured on a dual beam oscilloscope along with the laser pulse. The mode locking of the pulse from the laser was monitored on a 519 scope. The change in conductivity  $\Delta G$  is estimated from the voltage measured ( $\theta$ ) across the resistance R (127  $\Omega$ ).

$$\Delta G = \frac{\theta}{(V - \theta) R} , \quad V = 22.5 \text{ volts.}$$

The intensity of the excitation intensity was varied using calibrated neutral density filters.

The photo conductivity  $\Delta G$  in millimhos versus relative laser intensity is shown in Fig. 4. A least square fit was made and the slope was found to be 2.9. This indicated a power law of ~ I<sup>3</sup>, characteristics of a three photon process. The generation of non-equilibrium carriers can be due to absorption of non-phase matched second harmonic generation in Cd S. Since the second harmonic is of 5300 Å, it will be absorbed only as a result two photon process. Hence, photo conductivity due to such a process is of fourth order and will be very weak compared to a three photon process. Therefore the observed photo conductivity is due to three photon absorption in Cd S.

We estimated the three photon absorption coefficient from the measured photo conductivity. For this, we had to know the peak intensity of the pulse. We measured the total energy of the mode locked pulse using a calorimeter. Under similar conditions, the two photon flourescence was photographed using Rhodamine 6G. This gave a value of  $\sim 3$  psecs without measuring contrast

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ratio. 1.4 ge ga watts/cm<sup>2</sup> of peak intensity gave a value of  $\Delta G = 0.8$  milli mhos. The three photon conductivity  $\Delta G_3$  can be easily written for a transient process using Jick Yee's formaula<sup>(11)</sup> as

$$\Delta G_3 = u \tau \frac{I_0}{3\hbar w} \left[ 1 - \frac{1}{(1 + 2K_3 I_0^2 L)^{1/2}} \right]$$

where

$$\alpha = q - \frac{\mu}{c} (\mu + \mu)$$

q = electronic charge

 $\frac{a}{c} = 2$  geometric factor

$$\mu_e + \mu_h = \text{mobility} = 200 \text{ cm}^2 |\text{volt-sec inCds}|$$

$$\mathbf{T}\mathbf{w} = 1.17 \, \mathrm{ev}$$

 $K_{30}^{2}$  = three photon absorption coefficient (cm<sup>-1</sup>)

The thickness of the crystal was measured with a microscope and was found to be approximately 0.2 mms. From the expressions for  $\Delta G_3$ ,  $K_3 I_0^2$  was estimated to be 0.167 am <sup>-1</sup>. For I  $\approx 1.4$  gega watts/cm<sup>2</sup>,  $K_3$  was found to be 0.043 cm<sup>3</sup>/Gw<sup>2</sup>. In a recent paper Jick Yee<sup>(12)</sup> calculated the three photon absorption coefficient using Hartree Fock approximation with a three valence bands model for Cd S and his value of  $K_3$  was 0.25 cm<sup>3</sup>/Gw<sup>2</sup>. Aykinadze et al <sup>(13)</sup> reported a value of 2.5 am<sup>3</sup>/Gw<sup>2</sup> for  $K_3$ . Their experiment was done with a q~switched laser pulse and their intensity dependence was 3.4. Aresenev etal<sup>(9)</sup> estimated using mode locked pulses and they got a value of 0.02 cm<sup>3</sup>/Gw<sup>2</sup>. Our experimental arrangement was similar to Aersenev's and is in the order of magnitude agreement with Jick Yee's theoretical value and Aresenev's experimental value gives one more evidence to the three photon generation process in Cd S.

B. M. Ashkinadrye et al  ${}^{(8)}$  observed a power law of I <sup>3.4</sup> in their o experimental investigation of the recombination radiation of the 5211 A band from Cd S at 77° K after three photon absorption of Q-switch Nd: glass radiation. Since they conducted the experiment at 77° K, they observed a large number of excitons. The excitons formed another recombination channel. They explained a power 'aw of I<sup>2.6</sup> in the case of luminescence of Cd S with two photons excitation <sup>(13)</sup> by assuming two recombination channels and the pumping of carriers from one recombination channel to another. This was supported by their observation of two recombination times exhibited by the decay of the photo current. The power law of I<sup>3.4</sup> could be similarly explained. However, in the case of three photon excitation at room temperature, the photo conductivity exhibited

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a power law of  $I^{3.0 \pm \cdot 2}$ . Since we operated the crystal at room temperature, there are very little excitons formed and so only one recombination channel is present as evidenced by the single recombination time (long time constant of 20 µ secs) in the photo conductivity decay even at the highest light intensitier. No fast recombination was observed corresponding to the deexcitation of the excitons. Further excitons don't contribute to photo conductivity. B. M. Ashkinadye et al <sup>(7)</sup> experimented on two photon conductivity in Cd S at room temperature excited with giant pulses from a ruby laser and observed a power law of  $I^{2.0}$ . Exploring these results, the slope of  $3.0 \pm .2$  in the log-log plot of  $\Delta G_3$  versus I could be justified.

In conclusion, we observed the three photon conductivity in CdS at room temperature using mode locked Nd: glass laser pulses and the three photon conductivity depended on excitation intensity as  $I^{3.0 \pm .2}$ . An order of magnitude estimate of the three photon absorption coefficient was found to be in fair agreement with the theoretically calculated values. The power law of  $I^3$  could be utilized to measure third order intensity correlations of the picosecond pulses. Third order processes like this could be observed easily with the use of picosecond pulses because of higher peak intensity. When Q-switched pulses of the same envelope density as that of the mode locked pulse train were used to excite the CdS crystal, no observable signal was detected. This indicates the advantages of using picosecond pulses in investigating multi-photon processes in semi-conductors.

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#### Figure Caption

- Fig. 1 Two photon conductivity change in GaAs vs laser intensity with mode-locked pulse excitation.
- Fig. 2 Experimental set-up for measurement of picosecond light pulses by using two-photon conductivity effect.
- Fig. 3 Experimental data of second order intensity correlation curve as measured by two photon conductivity effect.
- Fig. 4 Three photon conductivity change in polycrystalline CdS sample vs laser intensity.









Fig. 3

