Incorporation of Acetoacetanilide Crystals in Host PMMA Polymer Matrix and Characterizations of the Hybrid Composite

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ABSTRACT

Blending of polymer with organic/inorganic material has given a new direction for developing novel materials. This is an easy and inexpensive method of modifying various properties of the individual materials and composites. The aim of the present work is an attempt to incorporate crystals in polymer host and investigate the effect, on optical properties of the of polymer-crystal combine. In this paper a novel technique of incorporating inorganic/organic crystal in a host polymer matrix is reported. Crystals of acetoacetanilide (AA) are grown in the host Poly Methyl Methacrelate (PMMA) by simple evaporation technique. The scanning electron micrograph (SEM) studies reveal the growth of Acetoacetanilide crystals of varying sizes and shapes (flakes) in prepared samples. The results are confirmed by various spectroscopic characterization studies such as X-ray diffraction (XRD), Fourier Transform Infrared Spectroscopy (FT-IR) and the UV-Vis spectroscopy. The investigations carried out, show a change in optical absorption band edge which is attributed due to change in band gap because of crystal sizes.

Key words: Polymer composite, Optical properties, PMMA-acetoacetanilide composite, Nano composites, Spectroscopic techniques.

1. INTRODUCTION

Inorganic/organic hybrid materials have attracted increasing interests due to their unique optical and electronic properties [1-6]. Organic crystals have a narrow transparency and exhibit low thermal and mechanical resistance. It is established that the polymers are excellent host
materials. When the nano-particles are embedded or encapsulated in polymer, the polymer acts as surface capping agent. In addition, the particle size is controlled well within the desired regime. For application in opto-electronics and electronics, the control of particle size and their uniform distribution within the polymer is the key to technology based on the nano-particles in polymers. With these advantages of organic crystal and polymer, a new approach, the growth of organic crystals in polymer matrix has been contemplated. This approach is intended to combine the properties of both classes of material that is the organic acetoacetanilide and the polymer PMMA. The authors herein describe the technique of incorporating acetoacetanilide crystals in PMMA matrix and to study the effect on the optical properties of acetanilide crystals due to the presence of the host PMMA. Various spectroscopic and morphological techniques are useful in characterizing the nano-composites. In the present paper the preparation, characterisation of the nano-composites and the effect on optical properties of acetoacetanilide crystals due to the presence of polymer matrix have been reported. The results are demonstrated by X-ray diffraction (XRD), Fourier Transform Infrared Spectroscopy (FT-IR) and the UV-Vis spectroscopy and scanning electron microscopy (SEM) studies.

2. EXPERIMENTAL

2.1 General

Research grade Acetoacetanilide and PMMA from Merck are used to prepare the composites. Using JEOL JD8p Visual XRD, (with automation software:VISX122D), with cu Kα of \( \lambda = 1.5405 \) Å (30KV) as a source, the Xrd data has been recorded. The FT-IR spectroscopy analysis has been carried out using Nicolet AVATAR 330 FT-IR instrument in the wave number range from 400cm\(^{-1}\) to 4000cm\(^{-1}\). Shimadzu UV-Vis –NIR Spectrophotometer model 3101 PC has been employed for UV-Vis measurements. JEOL JSM- 5800 LV scanning electron microscope (SEM) has been used to obtain the microimages of the crystals-polymer composites.

2.2 Preparation of Acetoacetanilide-PMMA Composites

The research grade materials of acetoacetanilide and PMMA have been procured from Merk and Co. To prepare the PMMA solution 10gm of PMMA is dissolved in 20 ml of double distilled water. A homogeneous transparent solution of PMMA is obtained after the solution is stirred continuously at room temperature for 6-8 hours. Then 5 ml of PMMA solution is taken in a beaker. A saturated solution of acetoacetanilide in chloroform is prepared and filtered. 5 ml of the filtered solution is mixed with the polymer solution in the beaker and stirred to obtain a uniform and homogeneous mixture. The blend is obtained by casting the prepared mixture into a glass petri dish and left for slow evaporation. The resulting thin sheet of acetoacetanilide and PMMA combined obtained (PM-AA), peels off the petri dish when completely dried. Two
samples of the PM-AA blend have been prepared by changing the concentration of acetoacetanilide (10 ml for the first sample and 15 ml for the second sample).

3. RESULTS AND DISCUSSIONS

3.1 XRD Analysis

The X-ray diffraction scan is a useful tool to examine the influence of PMMA contents on the crystalline structure of acetoacetanilide. From the Xrd data recorded, the Xrd spectrograph for pure PMMA, acetoacetanilide and the composites acetoactanilde-PMMA (PM-AA1 and PM-AA2) is shown in the Figure1.

From the fig. 1 it is clear that for the pristine acetoacetanilide (AA) the diffraction peaks observed at $8^\circ (2\theta)$ is due to crystalline structure of acetoacetanilide. Pure PMMA exhibits an amorphous feature which is characterized by a hump at 16 degrees ($2\theta$) with no sharp peaks. From the diffraction scans of the samples it is found that

- Amorphous hump is prominent in the sample of PM-AA1

- A decrease in the relative intensity of peaks appearing at 8 degrees is observed in the samples of PM-AA1 and PM-AA2. This is attributed to the complexation of the PMMA-acetoacetanilide blend.

![XRD spectrograph patterns of the composites](image-url)

Fig. 1 XRD spectrograph patterns of the composites
The peaks in Xray diffraction patterns of the PM-AA combined are well in agreement with that of acetoacetanilide. More peaks are observed in the prepared composite PM-AA2 than the pristine acetoacetanilide. This may be due to the presence of crystals of bigger sizes and orientation. In the XRD pattern of PM-AA1, broadening of peaks is seen. This is attributed to the presence of the crystallites of smaller sizes. Large intensity peaks and the increase in the number of peaks observed in the sample of PMAA-2 are correlated to a high degree of orientation.

3.2 FT-IR Studies

The FT-IR spectrographs of PMMA-AA composites are depicted in Fig. 2.

In the spectra of the PM-AA blend characteristic absorption bands have been identified and assigned, by comparison with the pristine acetoacetanilide and PMMA. These bands are listed in the table 1. The details of functional groups of PMMA are a sharp intense peak at 1731 cm\(^{-1}\) appeared due to the presence of ester carbonyl group stretching vibration. The broad peak ranging from 1260-1000 cm\(^{-1}\) is owing to the bending of C-O stretching vibration. The stretching vibration is shown by the presence of a broad peak between 3100-2900 cm\(^{-1}\). In spectra of acetoacetanilide the absorption peak appearing at 1650 cm\(^{-1}\) is due to NH bending and the CH bending at 1425 cm\(^{-1}\). The IR spectra of the PM-AA composites (Sample 1 and Sample 2) show the major peaks observed in the individual PMMA and acetoacetanilide, with a small shift and change in the intensity of the observed peaks. This confirms the presence of acetoacetanilide crystals in PMMA matrix [9-10].
Table 1. Characteristic absorption peaks of PMMA, AA and the composite PMMA-AA

|                  | PMMA                  | Acetoacetanilide          | Composite PM-AA1 and PM-AA2 |
|------------------|-----------------------|---------------------------|-----------------------------|
| Peak position cm$^{-1}$ | Bond Assignment | Peak Position cm$^{-1}$ | Bond Assignment | Peak Position cm$^{-1}$ | Bond Assignment |
| 752              | CH                    | 3300                      | OH Stretching              | 3252                      | OH Stretching   |
| 987              | Bending (C-O)         | 2950                      | CH aliphatic str           | 2948                      | CH aliphatic str |
|                  |                       | 1650                      | NH bending                 | 1597                      | NH Bending      |
| 1200             | (C-O)Stretching       | 1425                      | CH bending                 | 1200                      | (C-O) Str       |
| 1723             | Stretching C=O        |                            |                            | 1723                      | Str C=O         |

3.3 UV-Vis Spectral Analysis

The UV-Vis spectral data recorded in the wavelength range from 200-3000 nm for the pure and PM-AA composites samples are shown in the Fig. 3.

![Fig. 3 UV-Vis Absorption spectra of Pure AA and composite PM-AA Samples](image)

The strong absorption peaks are evident from the UV absorption spectra of the samples shown in Fig. 3. A sharp absorption peak is exhibited by acetoacetanilide at 360 nm. The addition of
acetoacetanilide in the host polymer environment causes the growth of crystals that is realized in very strong absorption peaks in the PM-AA1 sample at 240 nm and a peak at 310 nm in the case of the PM-AA2 sample. This suggests the blue shift in the absorption edge of acetoacetanilide in the polymer matrix (PM-AA1 and PM-AA2) with respect to bulk acetoacetanilide. The observed blue shift indicates decrease in particle size.

Energy band of materials is related to absorption coefficient $\alpha$ by the Tauc relation

$$(\alpha h\nu) = A (h\nu - E_g)^n.$$ 

Where $A$ is a constant, $h\nu$ is the photon energy, $E_g$ is the band gap and $n$ is an index which assumes values $\frac{1}{2}$, $\frac{3}{2}$, 2 or 3 depending on the nature of electronic transition responsible for the absorption. By taking the least index $n=1/2$, $(\alpha h\nu)^2$ and $h\nu$ in eV, the plot at room temperature shows a linear behaviour of the absorption edge. The extrapolation of the linear part of the curve on X axis gives the energy gap. The plot made for the samples (PM-AA1, PM-AA2) is shown in the Fig. 4.

![Fig. 4 $(\alpha h\nu)^2$ and hv plot for the samples of PM-AA](image)

The optical energy gap for the samples of PM-AA1 and PM-AA2 is increased to 3.75 eV and 4.95 eV respectively compared to bulk AA, which is 3.25 eV [11-12]. The absorption peaks and the calculated energy gap from the absorption peaks (Fig. 3) and the energy gap from $(\alpha h\nu)^2$ and hv plot for the bulk AA and composites of PM-AA1 and PM-AA2 are tabulated in the table. 2 shown below.
Table 2. Absorption peaks and Energy Gap of acetoacetanilide and the composites of PM-AA

| Samples   | Absorption peaks [λ vs Absorbance] | Calculated energy gap $E_g$ in eV | Energy gap in eV $[(\alpha h\nu)^{2} vs E_g$ plot] |
|-----------|------------------------------------|-----------------------------------|-----------------------------------------------|
| AA        | 360 nm                              | 3.45                              | 3.25                                          |
| PM-AA1    | 240 nm                              | 5.175                             | 4.95                                          |
| PM-AA2    | 310 nm                              | 4                                 | 3.75                                          |

3.4 SEM Analysis

The morphology of PM-AA blends has been studied using SEM. The SEM image of PM-AA sample is shown in Figure 5.

![Figure 5 SEM micrograph images of PM-AA1 (a) and PM-AA2 (b) composites](image-url)
The SEM images of PM-AA1 and PM-AA2 samples are shown in the Figure 5. The images show plate type (flakes) crystals with high crystal density. This is attributed to the agglomeration of smaller crystals [13] due to high surface to volume ratio of the crystals.

3.5 SHG Measurement

The second harmonic generation (SHG) measurement using the 1.06 µm pulsed radiation of a Nd:YAG laser, as a fundamental wave has been carried out. It is demonstrated that PMMA and acetoacetaenilide are both non-linear and has second harmonic conversion efficiency. The blends of PM-AA1 and PM-AA2 samples do not show any second order conversion. That is the second harmonic conversion efficiency is absolutely absent [14] for the composites of PM-AA.

4. CONCLUSIONS

The XRD analysis reveals that the PMMA and acetoacetaenilide composite has been formed. The studies suggest the effectiveness of crystal size control, by temperature and evaporation restriction and volume of the solutions used. Characteristic absorption bands from FTIR spectrum of composites PM-AA are identified and assigned by comparison with the values found for PMMA and acetoacetanilide individual materials. The presence of absorption peaks of individual materials in the PM-AA composites confirms the incorporation of acetoacetanilide in the PMMA host polymer. The change in the UV-visible spectrum is due to complex formation which can be reflected in the form of change (decrease in PM-AA samples) in the optical energy gap. Morphology of PM-AA shows crystalline domains uniformly shaped. The blend PM-AA does not show any second harmonic conversion though acetoacetanilide has SHG property.

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