Optical Properties of Poly (Vinyl Chloride-co-Vinyl Acetate-co-2-Hydroxypropyl Acrylate) / (Acrylonitrile-Butadiene-Styrene) Blends

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ABSTRACT

The morphology and optical properties of PVVH, ABS and its polyblend samples have been studied using different techniques such as atomic force microscopy, AFM, Fourier transform-Infrared, FTIR, Raman spectroscopy and UV-Vis spectroscopy. AFM images showed that roughness of polyblend samples is increased as PVVH content is decreased in the polyblend samples. FTIR spectroscopy has been investigated for PVVH, ABS and its polyblend samples in the range from 4000 to 500 cm⁻¹. The characteristic absorption bands of PVVH and ABS are detected. On the other hand, the main absorption bands of pure materilas have been affected in their positions and intensities in the FTIR spectra of polyblend samples. Raman Spectroscopy is carrid out for all the samples in the range 3650-50 cm⁻¹. Analysis of UV spectroscopy showed that absorption edge and indirect optical energy gap are decreased with decreasing PVVH content in polyblend samples.

Keywords: PVVH, ABS, Polyblend, Miscibility, AFM, Raman shift.

INTRODUCTION

The polymer blends have been attracted much attention in the area of research and development in polymer science in the past three decades, because of their

potential usage in many applications, such as, electro-optical devices, pipes, sensor technology, shielding, polymer electrolytes and membranes [1-6]. One of the commercial advantages is that polymer blends provide a method to produce new materials with new property profiles, which reduces development costs. Polymer blending is an interesting method for investigating the phase separation and polymer interactions at microsopic and mesoscopic levels.

ABS terpolymer is widely used in engineering thermoplastics containing rubber. It is characterized by two phase systems with a glassy poly (styrene acrylonitrile), (SAN) copolymer and a rubbery polybutadiene (PB). PVVH is an amorphous terpolymer and has a sensitive chemical structure to variation of temperature. The properties of polymer blends can be improved by varying composition ratio and conditions of the processing. Both dipole-dipole interaction and hydrogen bonding are responsible for polymer-polymer compatibility and miscibility [7-8].

The investigation of both optical properties and electrical properties of polyblend materials is very important as well as the study of their thermal and mechanical properties. The present article is devoted for investigating the morphology and optical properties of PVVH/ABS polymer blend.

Experimental Work

Poly (vinyl chloride-co-vinyl acetate-co-2-hydroxypropyl acrylate), PVVH, is supplied fom Aldrich Chemicals. Acrylonitrile-butadiene-styrene, ABS, is supplied from Poly Sci., USA. The films of samples are prepared using casting method. Tetrahydrofuran (THF) is used a common solvent for PVVH and ABS. The solvent was eleminated in an oven, at temparture of ~ 323 K for two days.

Atomic force microscope (AFM) is investigated using Bruker Dimension Icon system with Scan Asys mode (tapping). Fourier- Transform Infrared Spectroscopy, FT-IR, is carried out in the range from 4000 to 500 cm⁻¹ using FT-IR spectroscopy, Omnic. Raman Spectroscopy is carrid out using Senterra II, Bruker in the range 3650-50 cm⁻¹. Ultra violte–Visible (UV-Vis) spectroscopy is carried out in the range from 200 to 800 nm by UV 5200 Spectrophotometer, at Physics Department, College of Science and Humanities, Prince Sattam bin Abdulazziz University.

RESULTS and DISCUSSION

Atomic Force Microscope (AFM)

The AFM amplitude and three-dimensional (3D) height images of PVVH, ABS and its polyblend samples are shown in Fig. 1. Large numbers of hills and valleys are observed in the 3D height images. AFM images in Fig. 1 exhibited surface with some agglomerates and low amplitude of roughness.

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Figure 1: AFM images of a) PVVH, b) 70wt% PVVH, c) 50 wt% PVVH, d) 30 wt% PVVH and e) ABS.

The values of R_{ms} , R_a and surface area of the samples are estimated and summarized in Table 1. Generally, the values of both R_{ms} and R_a will be similar to each other if the sample surface is flat and does not contain any deviations from the mean surface level. On the other hand, values of R_{ms} will be greater than R_a if the surface of the sample is very rough and contains significant numbers of holes and large bumps [9]. From the R_{ms} and R_a values of these blends in Table 1, one can conclude that the surfaces of the polyblend samples are very rough and large bumps and holes are found in appreciable numbers.

Material	Average surface roughness R _a (nm)	RMS surface roughness R _q (nm)	Average height R _{max} (nm)	Surface area (µm ²)
PVVH	0.57	0.74	12.1	1.04
70 wt% PVVH/ 30 wt% ABS	0.92	1.90	34.5	26.3
50 wt% PVVH/ 50 wt% ABS	2.97	4.99	49.1	1.07
30 wt% PVVH/ 70 wt% ABS	4.41	6.23	83.3	1.03
ABS	1.75	2.53	30.7	1.06

Table 1: The values of R_a, R_q, R_{max} and surface area of all samples.

Fourier Transform- Infrared (FTIR)

FTIR spectroscopy is a highly effective method to examine the polymer-polymer interactions. This method is used extensively to investigate the mechanism of interpolymer miscibility qualitatively and quantitatively through the hydrogen bonding [10-12]. FTIR spectrum of pure PVVH is characterized by bands at 2968 and 2916 cm⁻¹ are related to C- H stretches, 1733 cm⁻¹ is attributed to C=O, 1441 cm^{-1} is attributed to CH₃ rocking, 1385, 1329, 1251, 1094, and 1049 cm⁻¹ are attributed to C-O stretch coupled with C-C vibration, 966 cm^{-1} is attributed to skeletal vibrations and 691 and 615 cm^{-1} are attributed to C-Cl stretches and skeletal vibrations, as shown in Fig. 2a [13-15]. FTIR spectrum of ABS is characterized by many absorption peaks, such as, 2234 cm⁻¹ and 1452 cm⁻¹, as shown in Fig. 2e. These bands are attributed to nitrile group in ABS, i.e., C≡N and double bonds of butadiene blocks. The absorption at 1677 and 1495 cm⁻¹ are related to styrene double bonds in ABS terpolymer. The absorption band at 2851 cm⁻¹ is attributed to the vibration of aliphatic and aromatic C-H bond. Also, the bands at 3022 cm⁻¹ and 2926 cm⁻¹ are related to aromatic C-H bonds and the vibration of aliphatic [16]. The bands at 700 and 755 cm⁻¹ are related to C-H def (out of phase) and at 964 cm⁻¹ is related to C=C-H *def* (aliphatic). On the other hand, it is observed that FTIR spectra of PVVH/ABS polyblend samples exhibit all the characteristic absorption spectral bands of PVVH and ABS with some changes in both their intensities and positions, such as 2234 cm⁻¹, 1670 cm⁻¹ and 1452 cm⁻¹,

as displayed in Fig. 2(b-d). This behavior is confirming to the high miscibility of the polyblend due to the formation of hydrogen bonding between the functional groups of both polymers [6].





Figure 2: FTIR spectrum of PVVH, ABS and samples of polyblends.

Figure 3: Ratio of I(615 cm⁻¹)/I(692 cm⁻¹) versus PVVH wt %.

The intensity ratio of both 638 cm⁻¹ and 692 cm⁻¹ absorption peaks $[I(638cm^{-1})/I(691cm^{-1})]$ is highly sensitive for any change in the molecular conformation and structure of the crystal [17]. It is observed that, this ratio of PVVH/ABS polyblend samples is decreased as PVVH content decresed, as shown in Fig. 3. Also, it is observed the intensity of many absorption bands such as 1733 cm⁻¹, 1251 cm⁻¹ and 966 cm⁻¹ has been decreased as the content of PVVH is decreased in the polyblend samples, as shown in Fig. 4.



Figure 4: The intensity of the bands 1733 cm⁻¹ and 1251 cm⁻¹ versus PVVH content.

Raman Spectroscopy

The advantage of Raman spectroscopy technique is its high sensitivity to nonpolar bonds and its ability to give qualitative and quantitative information on the macromolecules in various configurational and phase states, involving the description of different crystalline modifications and amorphous regions having different states of order [18]. Fig. 5a displays Raman spectrum of pure PVVH. It is observed that, Raman spectrum of PVVH is characterized by seven obvious bands located at 639 cm⁻¹, 695 cm⁻¹, 860 cm⁻¹, 1105 cm⁻¹, 1381 cm⁻¹, 1870 cm⁻¹, 2920 cm⁻¹ and 3283 cm⁻¹, respectively. The band at 2920 cm⁻¹ is attributed to C-H stretching. The band at 1381 cm⁻¹ is related to the twist or bending of CH₂ and CH₃ bonds. The region of 1000–1150 cm⁻¹ involves C-C stretching vibrations and this region is very sensitive to the conformational changes, i.e., the properties of the bands in this region has been modified, as shown in Fig. 5c for the PVVH/ABS polyblend samples [17]. The bands in the region $639-690 \text{ cm}^{-1}$ are related to the C-Cl stretching. Raman spectrum of ABS is characterized by three bands detected at 1604 cm⁻¹, 1666 cm⁻¹ and 2238 cm⁻¹, as shown in Fig. 5b. These bands are attributed to styrene, butadiene and acrylonitrile phases, respectively.



Figure 5: Raman shift of a) PVVH, b) ABS and c) polyblend amples.

UV-Vis Spectroscopy

The electronic structure of polymeric materials can be studied by investigation its optical absorption. The absorption is occurred when the photon has an enough amount of energy to excite the electrons from lowest energy to highest energy levels. Based on Davis–Mott model, the wave function is localized and the transition probability depends mainly on the overlapping of wave functions [19]. In the amorphous material, the localized states extend from the conduction and valence bands to the energy gap band. Hence, these localized states can affect the electron transition in polymeric materials [20].

UV spectra of PVVH, ABS and its polyblends have been investigated in the range of wavelength from 200 to 850 nm, as shown in Fig. 6a. The investigation of optical

absorption gives more valuable information regarding the band structure of polymeric materials. The absorption coefficient of all samples is calculated using the following equation [21]

$$\alpha(\nu) = 2.303 \frac{A}{d}$$

Where, A and d are defined as the absorbance and thickness of the sample, respectively. The valuable information of both the optical energy band gap (E_g) and electronic band structure can be obtained using absorption coefficient (α). Fig. 6b displays the variation of optical absorption coefficient (α) against photon energy (h υ) for all samples. The values of absorption edge are calculated by extrapolating the linear portion of α to zero absorption value and summarized in Table 1. On the other hand, the optical energy band gap (E_g) values for pure and polyblend samples are calculated using the following formula [22]

$$\alpha = B \left(\frac{h \upsilon - E_g}{h \upsilon} \right)^n / h \upsilon$$

Where, *B* is a constant and depends mainly on probability of the transition and localized state width inside the band gap, hv is the photon energy, *n* is an index depends on the transition nature and is attributed to the density of states distribution. For the allowed direct and indirect transitions, *n* will equal to 1/2 and 2, whereas, for forbidden direct and indirect transition *n* equals to 3/2 and 3, respectively. Fig. 6c shows $(\alpha hv)^2$ against hv for all samples under investigation. The allowed indirect optical energy gap values are estimated from the intercept of the extrapolated linear portion of $(\alpha hv)^2$ on the photon energy axis, as shown in Fig. 6c, and summarized in Table 1.



Figure 6: a) Absorbance, b) Absorbance coefficient (α) and c) $(\alpha h \upsilon)^2$ versus h υ for pure and polyblend samples.

It is observed that with decreasing PVVH content the optical band gap of PVVH/ABS polyblend samples are decreased. This behavior can be interpreted based on the increasing in the number of unsaturated defects will increase the final

states density in the band structure resulting in low optical band gap energy or may attributed to structural rearrangement [23].

The physical processes that primarily control the behavior of the gap states in the amorphous polymeric materials are structural disorder which responsible for structural defects and tail states in deep states. The lattice configuration fluctuation in polymeric materials will distort the fundamental edge resulting in Urbach tail [22]. Fig. 7 displays the variation of $ln \alpha$ against ho for all sapples to get some information about the band tails. The values of Urbach energy (E_U) are calculated using the slope of linear relationship of ($ln \alpha$) versus ($h\nu$) using the following equation



Figure 7: $\ln \alpha$ against photon energy hu.

where α_0 and E_U are defined as a constant and Urbach energy, respectively and is often explained as the tail of localized states width in the band gap. The exponential tail is appeared because amorphous and/or disordered materials produce extended localized states in the bandgap.

Table 1: The v	alues of Absor	ption Edge,	Optical Energy	' Gap
a	nd Urbach ene	rgy of all sat	mples.	

Material	Absorption Edge (eV)	Optical Energy Gap (eV)	Urbach enegy (eV)
PVVH	4.88	5.12	0.60
70 wt% PVVH/ 30 wt% ABS	3.68	4.17	0.98
50 wt% PVVH/ 50 wt% ABS	2.97	3.94	0.69
30 wt% PVVH/ 70 wt% ABS	3.27	3.59	0.53
ABS	3.12	3.91	0.66

CONCLUSION

AFM images of pure and polyblend samples revealed that the roughness of sample surface is increased as PVVH content is decreased in the polyblend samples. FTIR spectroscopy has been investigated for PVVH, ABS and its polyblend samples. It is found that FTIR of PVVH is characterized by main absorption bands at 2968 cm⁻¹, 1733 cm⁻¹, 1441 cm⁻¹ 691 cm⁻¹ and 615 cm⁻¹, while the absorption bands characterizing acrylonitrile, butadiene and styrene of ABS are obtained at 2234 cm⁻¹, 1452 cm⁻¹, 1677 cm⁻¹ and 1495 cm⁻¹. On the other hand, the main absorption bands of pure materilas have been affected in their positions and intensities due to the blending, as observed in both FTIR and Raman spectra, confirming the high miscibility between the polymers. Analysis of UV spectroscopy showed that absorption edge, indirect optical energy gap and Urbach energy are decreased with decreasing PVVH content in polyblend samples. This behavior is attributed to structural rearrangement.

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