INTRODUCTION

The polymer composite material has given the important responsibility of new materials [1]. When it is made a composite and mixed to nano-size materials, polymer composite has applications of one or more dimensional materials and amorphous materials [2]. In the important development of nano-sized materials have a way of increasing the quality of structure and strength of the prepared polymer nanocomposite materials. In this century, nanotechnology is a quality promising field of energy storage and biotechnology systems [3]. In the basic idea of nanotechnology is in controlled prepared nanosize particle and polymer nanocomposites on the size of particles 1 to 100 nm [4]. This nanotechnology effort will help to raise the fields of materials science, physics and integrated chemistry [5]. All type of materials like crystalline and amorphous materials was got the polymer nanocomposite materials from the different type of preparation. These new materials of polymer nanocomposites are helping to increase the quality of societal advantages in the world. So we were interested to prepare the polymer nanocomposites materials in the combination of conducting polymer and metal oxide. Conducting polymers are conjugated chain of organic compounds; it shows good conductivity similar to good conductivity metal because of the presence of large number of charge carrier of extended \( \pi \) electrons [6]. The backbone of the polymer chain has charge mobility, which allows conducting polymer [7]. Their conductivity behaviour is compatible with normal conductors but conducting polymer has large advantages [8], like as being flexibility, light weight and very less amount. These conducting polymers have various applications of mobile phone, LED, batteries, solar cells, many microelectronic devices and sensors.
Among the conducting polymer families, polyaniline (PANI) has been of peculiar characteristics. It is most controllable electrical conductivity, nice redox properties, good conductivity, easy polymerization, chemical stability and low cost monomer. PANI combined with pure inorganic nanoparticles like as MnO₂, SnO₂ and TiO₂ to form PANI/metal oxide nanocomposite to improve electrical, chemical and physical properties. This type of nanocomposite has many advantages, can have good potential applications in latest devices. The combining properties of nanocomposites have improved the optical and dielectric properties [10]. Metal oxide could act as conductive junctions, between the PANI chains that resulted in an increase of the electrical conductivity of the nanocomposites. The among metal oxide nanoparticles used, cerium oxide (CeO₂) nanoparticle are very nice in modern materials. Due to special magnetic, optical and electronic properties. CeO₂ has many applications in capacitors, microelectronics, electrode materials, solar cells and biomedical fields [11]. The synthesis of CeO₂ nanoparticles embedded in PANI using chemical, electrochemical and in situ polymerization methods. In this research article reported that, the PANI/CeO₂ nanocomposites (10 wt %) are synthesized with using of chemical agent of ammonium per sulphate in situ polymerization method. PANI/ CeO₂ nanocomposites material could have nice behaviour between electrical properties of PANI and metal oxide for new devices.

**EXPERIMENTAL**

The two moratilities HCl, ammonium persulphate, prepared CeO₂ nanoparticles and aniline were used to prepare 10 wt % of nanocomposites in situ polymerization technique.

**General procedure:** In a typical preparation PANI/CeO₂ nanocomposites were synthesized in situ polymerization methods as follows: The 4.5 mL of aniline and 70 mL of HCl in 2 M are mixed and stirred at 60 °C. In this prepared solution, 0.9 g (10 wt %) of prepared CeO₂ nanoparticles was added and stirred continuously for 3 h. During the stirring, prepared ammonium persulphate solution was added drop-wise into the solution. The prepared solution was filtered and washed by de-ionized water. The finally green colour precipitate was dried for room temperature for 3 days.

**Detection method:** The powder XRD pattern of the nanocomposites sample was recorded as PAN analytical model diffractometer (Kα = 1.54056 Å at 35 kV, 10 mA). The sample was scanned at the angle of 10° to 80° (2θ). The high resolution transmission electron microscopy (HRTEM) image was carried out at JOEL JEM 2100. The scanning electron microscope (SEM) images were recorded at Hitachi SEM. The optical spectrum of FTIR was carried out using 8400S Shimadzu spectrometer at the range wave number 4000-400 cm⁻¹. Another optical spectrum of UV was recorded in the wavelength range of 190-1100 nm. AC conductivity analysis was carried out using a Zahnner zennium IM meter in the frequency range of 10 μHz to 8 MHz at various temperatures.

**RESULTS AND DISCUSSION**

**X-ray diffraction studies:** Fig. 1 shows that X-ray diffraction pattern of as synthesized nanocomposite (10 wt %) materials. The prepared PANI/CeO₂ nanocomposites has been compared to the result with XRD pattern of the samples of pure CeO₂ and pure PANI. Pure CeO₂ is a nano-crystalline nature [12] and conducting polymer PANI is also nano-crystalline nature materials [13]. From Fig. 1 is clear that peaks show that the prepared nanocomposite materials are crystalline in nature. It was clearly observed that the changes of peak intensities compared to the pure sample of CeO₂ and PANI nanoparticles. This indicates the peaks of XRD pattern and results intercalated morphology of pure sample, the CeO₂ nanoparticles intercalated within the polymer matrix due to the penetration of pure nanoparticles, have in dispersion of the CeO₂ nanoparticles with in the polymer matrix. The particle grain size was calculated using FWHM, the formula is

\[ d = 0.9\lambda/\beta \cos \theta \]

where \( d \) is the crystalline size in nm, \( \lambda \) is the wavelength of X-ray in Å, \( \beta \) is the FWHM and \( \theta \) is the angle of diffraction in degrees. The particle grain size of the nanocomposites are observed and is varied from which compared to the pure materials. Particle grain size of maximum number of nanoparticles is around 16.5 nm.

**SEM and HRTEM studies:** Fig. 2a and 2b show images of SEM and HRTEM. From images it is observed to be spherical with small agglomeration. Fig. 2a and 2b clearly indicate that the pure nanoparticles PANI and CeO₂ penetration on the matrix of polyaniline, was confirmed on the formation of nanocomposites. HRTEM image shows agglomerated cluster of CeO₂, it is reduced that the making nanocomposites [14]. The polymer chains have been occurred nanoparticles in the compositions of PANI/CeO₂ nanocomposites. From the image shows that the nanocomposites particles were nearly spherical shape, it is possible to confirm of various crystallographic lattice plane of different atomic density. These images exhibit that the range of particles from 10 to 20 nm. It will give proof for the confirmation of supra-aggregates as found in the image of HRTEM and supra-aggregates are nearly stable from the nanocomposites suspension [15]. From some images could occurred in very smaller particles size of 5-10 nm, are combined together to grown to irregular shape of nanocomposites within the size of single particles. Since, nanocomposites
exhibit to less surface energy and we found the spherical shape with the size between 10 to 30 from the SEM images.

**Ultraviolet studies:** The inspirational of electron in molecules and atoms to higher energy state, it is results in absorption of UV-visible light. In all atoms undergoes electron rebel coming after absorption of light. In characteristic of chemical structure, the wavelength of light is absorbed; the molecular species was indentified using by absorption spectrum [16]. UV spectra of PANI/CeO$_2$ nanocomposites in 10 wt % shows in Fig. 3a. From figure, nanocomposites material showed two absorption at below 400 nm and one broad absorption above 400 nm. Two absorption bands placed at 324 and 368 nm, other broad bands placed at 858 nm in the range of UV from charge transition of nanocomposites materials. From figure shows, wavelength fall begin above 1000 nm. The first small peak at 324 nm is related to the absorption peak of CeO$_2$ nanoparticle [17], which blue shifted from 341 to 324 nm with decrease due to formation of nanocomposites. The second and third broad absorption peak at 368 and 858 nm are related to the absorption peak of PANI nanoparticles. The UV absorption sharp peak gives information about band gap. The band gap energy have been obtained which extension of straight line plot of $h\gamma$ vs. $(\alpha h\gamma)^2$. Here $h\gamma$ and $h\gamma$ are the normal and usual meaning. $\alpha$ is an absorption coefficient. $\alpha$ is calculated from the relation is $\alpha = 2.303 \log (\text{absorption})/\text{t}$, where $t$ is a grain size of the nanoparticles. The relation between $\alpha$ and $h\gamma$ is $(\alpha h\gamma)^2 = A (h\gamma - E_g)^{2\alpha}$. Where $E_g$ is a band gap and $A$ is a constant. The band gap energy have

**FTIR studies:** The FTIR spectra shows the execution of the formation of CeO$_2$ and remaining organic part on the surface of CeO$_2$ nanoparticles [18]. Fig. 4 shows the FTIR spectra of PANI/CeO$_2$ nanocomposites with 10 wt %. From figure, peaks/bands at 879 cm$^{-1}$ be associated to metal oxygen bond. The peaks at 1477.37, 1242.07, 702.04 and 507.24 cm$^{-1}$ agree with the functional groups of PANI. The complex peaks measured at 1658.66 and 1515.94 cm$^{-1}$ are due to undesirable residues in the nanocomposites. From figure, is measured that there are no peaks linked to residual H$_2$O and hydroxyl group of pure CeO$_2$ nanoparticles. The peaks in 1552.59 cm$^{-1}$ region are assigned to C-C stretching mode for benzenoid ring, 1299.93 cm$^{-1}$ are assigned to C-N stretching mode, 1134.07 cm$^{-1}$ are assigned to C=N stretching of secondary aromatic amine, can be requisite the presence of anion serious of acid atoms which are presume to be occupied to counteract the positive ion on the nitrogen molecule of PANI during polymerization. The peaks of wave number 806.19 cm$^{-1}$ is assigned to aromatic C-H out-of-plane bending [19].
AC conductivity studies: Fig. 5 shows the AC conductivity curve of PANI/CeO$_2$ nanocomposites frequency range from more than 10 $\mu$Hz to less than 8 MHz. Using formula of AC conductivity is

$$\sigma(\omega) = B(T) \omega(T)^{\beta}$$

where $\omega$ is the angular frequency, $\varepsilon_o$ and $\varepsilon_r$ are the permittivity of free space and medium.

The frequency dependence conductivity formula can be written as:

$$\sigma(\omega) = B(T) \omega(T)^{\beta}$$

The characteristics of AC conductivity the nanocomposites divulge a frequency independent conductivity in the very low range frequency area [19]. From figure, the high frequency region conductivity increases as the frequency increases [20]. The AC conductivity varied as per variation of frequency and will dependents on the temperature. Upto around 0.5 MHz, the AC conductivity varied as per variation of frequency and region conductivity increases as the frequency increases [20]. From figure, the high frequency divulge a frequency independent conductivity in the very low range frequency area [19]. From figure, the high frequency divulge a frequency independent conductivity in the very low range frequency area [19].

**Conclusion**

_in situ_ Polymerization method has been used for the synthesis of PANI/CeO$_2$ nanocomposites. The technique is carried out to be efficient for capable to control structural and morphology characteristics. The average grain size of the nanocomposites calculated with help of XRD method and HRTEM image is around 16.5 nm. The collection of nanocomposites samples synthesized is deducted from SEM analysis. The PANI/CeO$_2$ nanocomposites show a three strong UV absorption peak at 324, 368 and 858 nm, the band gap energy is 3.36 eV. The functional group of the nanocomposite materials was analyzed by FTIR spectrum. The frequency dependence AC conductivity was calculated with the help of different temperature region.

**REFERENCES**