Research Article

A Simple Synthesis of Higher Refractive Index Polymeric Nanocomposite Containing the Pendant ZnS Nanocrystals Capping Different Amount of Mercaptoethanol

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In this paper, three kinds of ZnS NPs capping different amount of mercaptoethanol (ME) were synthesized, and the effect of capping amount on ZnS NPs was studied, indicating that the capping amount of ME on ZnS surface decreases, while ZnS size increases as the decrease of added capping agent in synthesis process, and ZnS NP capping middle amount of ME (~27.7 wt%) shows a highest apparent refractive index (RI) value. ZnS NPs were composited into polymeric matrices by a simple “one-step” thermocuring method characterized by adding a small amount of functional monomer of glycidyl methacrylate (GMA) and solvent of N,N-dimethylformamide (DMF). The reaction mechanism, studied by $^1$H NMR spectra, indicates that the epoxy of GMA monomer can be easily opened and chemically grafted on ME-capped ZnS surface under the catalysis of DMF, and then copolymerized with other monomer. By the route, ZnS NPs can be composited into pure poly(N,N-dimethylacrylamide)-type (DMA-type), DMA-type copolymer, and DMA-free matrices to fabricate transparent films, and its RI value can be improved by either optimizing capping amount on ZnS surface or increasing ZnS content in the nanocomposites. The RI value of resulting dried nanocomposites can be improved to 1.764 by compositing 80 wt% ZnS capping middle amount of ME.

1. Introduction

High refractive index (RI) optical materials have extensive applications in lenses, prism, waveguides, and light-emitting diodes (LEDs) [1–3], and great progress has been made to synthesize high RI materials. An effective way is grafting high RI organic molecules with aromatic [4], fused ring [5] groups, or heteroatoms [6] in the optical materials. The other promising strategy is compositing high RI inorganic nanoparticles (NPs) into polymer matrices, because many kinds of high RI inorganic NPs, e.g., quantum dots (QDs), have been successfully synthesized, and its size less than 10 nm can ensure transparency of resulting nanocomposite materials. Two main approaches have been developed to composite inorganic nanoparticles in polymeric matrices including either in situ formation of inorganic NPs in the polymeric matrix or bulk polymerization of organic monomer solution in the present of premade nanoparticles. The latter approach is usually proved to be more effective, because it can provide full synthetic control over both the nanoparticles and the polymeric matrices.

It is crucial to maintain high particle dispersion homogeneity in nanocomposites for optical application. However, nanoparticles tend to aggregation in polymeric matrices, resulting in transmittance loss due to light scattering. Two methods can be used to reduce particle aggregation either bonded covalently nanoparticles on the polymeric skeleton or embedded physically nanoparticles in special polymeric matrices that interact with the nanoparticles. Usually, the former method can ensure a more homogeneous intermixing
between organic and inorganic components, because stronger interaction from covalent bond can effectively block nanoparticle aggregation.

ZnS NPs have high RI (2.36 at 620 nm) and low absorption coefficient (400-1400 nm), and it can be composited into polymeric matrices to prepare high RI nano-composite film. Moreover, ZnS can also be used as a higher band-gapped shell material around QDs to improve the fluorescence efficiency and stability of nanocrystals due to decreased surface defects [7]. Nowadays, many kinds of quantum dots such as CdS [8], CdSe [9], CuInS₂ [10], and ZnSe [11] have been wrapped ZnS shell material. Therefore, it is highly expected to develop novel approach to composite ZnS NPs in all kinds of polymeric matrices. Poly(N,N-dimethylacrylamide) is the most commonly used polymeric matrices embedding physically ZnS NPs [12]. We have also bound typical semiconductor nanocrystals such as photoluminescence (PL) properties of ZnS NPs were composited into polymeric matrices to fabricate high RI transparent ZnS-polymer nanocomposite films. However, a large amount of capping agent on ZnS surface (about 37 wt%) limits further RI improvement of resulting nanocomposite films.

Capping agents are well recognized as a key factor in synthesis and manipulation of quantum dot nanocrystals [13], in which capping agent maintain a dynamic bonding with the surface of nanocrystals at high reaction temperature above 250°C, ensuring a sufficient coverage of capping agent on a nanocrystal at a given moment resulting in a good solution stability [14]. Many kinds of capping agents have been bound on the surface of quantum dots, and research shows that capping agent plays also integral role in the function of the nanocrystal, such as photoluminescence (PL) properties of semiconductor nanocrystals [15]. However, to our knowledge, the effect of capping amount on nanocrystal structure and its RI value is rarely reported.

In this paper, three kinds of ZnS NPs capping different amount of ME were synthesized, and the effects of capping amount on the size, crystal structure, and refractive index of ZnS NPs were studied for the first time. Furthermore, ZnS NPs were composited into polymeric matrices to fabricate high RI transparent nanocomposites by a novel, simple, and universal route. The effect of capping amount on the RI value of the resulting nanocomposites was also studied emphatically. The paper involves the design and optimization of the nanoparticle with different amount capping agent and polymeric monomer as well as a suitable polymerization process. The approach has been proved to be effective in the fabrication of both DMA-type and DMA-free nanocomposites with higher RI value by compositing ZnS NPs with middle capping amount of ME.

2. Experimental Section

2.1. Materials. Zinc acetate dehydrate (Zn(Ac)_2·2H₂O), thiourea, and dibutyltin dilaurate (DBTDL) were purchased from Damao Chemicals (Tianjin, China); mercaptoethanol (ME) was purchased from Aladdin Chemical Reagent Co., Ltd. (Shanghai, China); methyl methacrylate (MMA), N, N-dimethylacrylamide (DMA), 2-isocyanatoethyl methacrylate, 2-hydroxyethyl methacrylate (HEMA), glycidyl methacrylate (GMA), and azobisisobutyronitrile (AIBN) were purchased from SA Chemical Technology Co., Ltd. (Shanghai, China).

2.2. Synthesis of ME-Capped and Polymerizable-Group-Capped ZnS NPs

2.2.1. ME-Capped ZnS NPs. Three kinds of ZnS nanoparticles, with different capping amount of ME, were synthesized according to a modified literature method [16]. Firstly, Zn(Ac)_2·2H₂O (5.5 g, 0.025 mol), thiourea (1.375 g, 0.018 mol), and different amount of ME (1.46 g, 2.18 g, 2.92 g) were, respectively, dissolved in 150 mL N,N-dimethylformamide (DMF) in a 250-mL three-necked round-bottom flask to obtain clear solution. Then, the solution was refluxed 10 h at 160°C under continuous magnetic stirrer and nitrogen protection. The resultant solution was concentrated to 20 mL using rotary evaporation under vacuum at 30°C, and then precipitated in excess ethanol to obtain crude white ME-capped ZnS NPs. The crude ZnS NPs was dispersed in 20 mL DMF, and then reprecipitated in excess ethanol again. Finally, the white precipitation was collected and washed thoroughly with methanol before dried in vacuum at 30°C. Three kinds of ZnS NPs with varying amount of ME from low to high are named as ZnS₁, ZnS₂, and ZnS₃, respectively.

2.2.2. Polymerizable-Group-Capped ZnS NPs. Polymerizable-group-capped ZnS was synthesized according to our previous route [17, 18]. Briefly, ME-capped ZnS NPs (3 g) and one drop of DBTDL were dispersed in 75 mL DMF, in which DMF solution of 2-isocyanatoethyl methacrylate (0.3 g in 15 mL DMF) was added drop by drop in half an hour, and then, the solution was kept at 30°C for 2.5 h under continuous stirring and ultrasound. Finally, the solution was concentrated to 10 mL by vacuum distillation, and then precipitated and washed with excess methanol. Three kinds of polymerizable-group-capped ZnS NPs with different amount of capping agent, named, respectively, as polymerizable-group-capped ZnS₁, polymerizable-group-capped ZnS₂, and polymerizable-group-capped ZnS₃, were dried in vacuum at 30°C.

2.3. Preparation of Transparent ZnS/Polymer Nanocomposite Films

2.3.1. Pure DMA-Type Nanocomposites Films. ME-capped or polymerizable-group-capped ZnS, with desired weight percent of the final dried nanocomposites, was dispersed in an equal-mass mixture of monomer of DMA and solvent of DMF by ultrasound, in which functional monomer of GMA (10 wt% of added ME-capped ZnS) was added and mixed up by ultrasound. Then, the initiator of AIBN (0.5 wt% of total monomer weight) was added in above solutions and dissolved by ultrasound to obtain a uniform transparent polymerization solution. Finally, the polymerization solution was sealed in the cavity of polypropylene plate molds separated by 0.16-mm thickness polypropylene frame, and then, the molds were placed in oven. The oven was heated up from room temperature to 80°C and maintained the temperature for 12 h to obtain transparent pure DMA-type polymeric nanocomposite films. The films were taken
off from the molds and removed the solvent of DMF under vacuum at 70°C for 4 hours.

2.3.2. DMA-Type Copolymeric Nanocomposite Films. ME-capped ZnS was dispersed in a mixture of monomer of DMA and solvent of DMF (1:1, m/m) by ultrasound, in which functional monomer of GMA (10 wt% of ME-capped ZnS) was uniformly added by ultrasound. Then, the other monomer of HEMA and initiator of AIBN was orderly added and mixed by ultrasound. Finally, DMA-type copolymeric nanocomposite films were prepared by a similar thermocuring process to pure DMA-type nanocomposites films.

2.3.3. DMA-Free Polymeric Nanocomposites. ME-capped ZnS was dispersed in solvent of DMF by ultrasound, in which functional monomer of GMA (10 wt% of added ME-capped ZnS) was added and mixed up by ultrasound. The resulting solution was maintained in oven at 50°C for 3h, and then, the monomer of MMA and initiator of AIBN were orderly added in and mixed up by ultrasound. Finally, DMA-free polymeric nanocomposites were fabricated by a similar thermocuring process to pure DMA-type nanocomposites films.

2.4. Characterization of the Nanocomposite Films Compositing ZnS NPs. 1H NMR spectra were recorded on a 400 MHz instrument (Bruker AC200) using DMSO-d6 as solvent. X-ray diffraction (XRD) spectra were recorded on a Bruker AXS D8 X-ray diffractometer with a Cu Kα (λ = 1.5406 Å) source, operated at 40 kV and 20 mA. The 2θ scanning range was set from 5° to 90° with a step size of 0.02° and at a scanning speed of 1 degree/min. Transmission electron microscopy (TEM) of ZnS NPs was observed with a JEM-2100 transmission electron microscope (Japan). Thermogravimetric curves of ZnS NPs were recorded using a SDT-Q600 TGA instrument in a temperature range from ambient temperature to 600°C at a heating rate of 5°C/min under nitrogen protection. The water content in the hydrated nanocomposite hydrogels was calculated by the ratio of weight difference between swollen and dried hydrogels to the weight of swollen hydrogels. The transmittance spectra of nanocomposite hydrogel films were recorded at the wavelength range from 200 nm to 800 nm using distilled water as a reference solution. The RI values of dried ZnS/polymer nanocomposite films were determined by a L116 ellipsometer (gaertner, USA), equipped with a He-Ne laser (wavelength 632.8 nm). The RI of swollen nanocomposite hydrogels was measured on a WYA Abbe refractometer at 589.3 nm. Mechanical properties of swollen nanocomposite hydrogel films were determined using a Series IX Automated Materials Testing System (Instron Corporation), with a crosshead speed of 20 mm/min at room temperature and a relative humidity of 50%.

3. Results and Discussion

3.1. Synthesis of ZnS Nanoparticles with Different Amount of Capping Agent. Three kinds of ZnS NPs capping different amount of ME were synthesized by changing the added amount of ME in the process of synthesis. When the mass ratio between capping agent of ME and zinc acetate decreased to 1:5.5, white precipitate insoluble in DMF emerged. Increasing the ratio to 1:3.77 for the synthesis of ZnS capping low amount of ME (ZnS1), Tyndall scattering was observed after 1 hour of reaction at 160°C, and then, the reaction solution turns white, indicating that the ZnS particle size increases rapidly. As the reaction goes on, the solution slowly turns clear again. Surface capping agent plays a decisive role in the internal structure of nanocrystals [19]. To observe the effect of capping amount on ZnS nanocrystals, high-resolution transmission electron microscopy (HRTEM) images of ZnS1-3 were performed, and the results were shown in Figures 1(a)–1(c). It can be seen that there is no obvious difference of ZnS crystalline structure with different capping amount of ME that all show distinct lattice fringes with a distance about 0.275 nm. This is important for ZnS NPs to maintain its attractive high RI properties [20]. HRTEM photography indicates also that ZnS capping middle amount of ME (ZnS2) shows a wider size distribution, and its nanocrystal shows an appearance from aggregative growth [21] as shown in Figure 1(b).

As shown in Figure 1(d), the X-ray diffraction (XRD) patterns confirm further the crystal structure and crystallinity of ME-capped ZnS NPs capping different amount of ME. Three broad diffraction peaks at 28.7°, 47.6°, and 56.4° were observed, indicating the (111), (220), and (311) planes of ZnS with zinc blend structure [22]. The size of ZnS1 and ZnS2 is 3.25 nm, and the size of ZnS3 is 2.95 nm calculated by Scherrer’s formula (d = kλ/β cos θ) based on the width at half maximum of peak (111) plane, indicating that increasing the capping amount helps to decrease the size of ZnS nanocrystals.

The capping amount of ME on ZnS NPs was determined by thermogravimetric analysis, and the curves were shown in Figure 1(e). The weight loss of ZnS1-3 is, respectively, 4.31%, 3.88%, and 1.43% at the temperature from 100 to 200°C that may be ascribed to residual solvent. Thiols like mercaptoethanol have the property of getting tightly adsorbed on the surface of the clusters, which can be decomposed at about 200°C [23]. Therefore, the rapid weight loss from 200°C to 325°C can be ascribed to the loss of tightly absorbed ME molecule on ZnS surface, and the weight loss of ZnS1-3 is, respectively, 23.59%, 27.7%, and 33.84% at the temperature range. Additionally, the decompose temperature of ME from ZnS surface decreases as the increment of added ME during ME-capped ZnS synthesis, indicating a more tight absorb ability of ME on the surface of ZnS NPs capping a smaller amount of ME. The content of pure ZnS in ME-capped ZnS1-3 nanoparticles is, respectively, 69.78%, 66.99%, and 62.63%, indicating that a large amount of capping agent exist still on the ZnS surface although greatly decreased the amount of added capping agent in the process of synthesis.

All three kinds of ZnS can be dispersed in DMF to obtain clear solution. The apparent RI of the DMF solution of ME-capped ZnS was shown in Figure 1(f). It can be seen that ZnS with 33.01 wt% capping amount has a higher apparent RI value than ZnS with higher or lower capping amount of ME at the same ME-capped ZnS content in DMF. This may be ascribed to its inhomogeneous particle size distribution.
of ZnS$_2$ NP, in which bigger size has a greater contribution to RI value.

3.2. Pure DMA-Type Copolymeric Nanocomposite Hydrogels from Polymerizable-Group-Capped ZnS. ZnS NPs have been embedded physically into DMA-based polymeric matrices. Herein, we bond covalently ZnS NPs on the polymeric matrices. To achieve this, polymerizable-group-capped ZnS NPs capping different amount of ME were synthesized by our previous method [17, 18]. However, polymerizable-group-capped ZnS capping low ME amount can not be dispersed in DMF and the monomer of DMA. The copolymeric hydrogels from polymerizable-group-capped ZnS with middle capping amount of ME (ZnS$_2$) shows obvious Tyndall scattering.
effect when added ZnS$_2$ is more than 30 wt\% as shown in Figure 2(a), and its transmittance decreases greatly with increased ZnS$_3$ as shown in Figure 2(b). However, The copolymeric hydrogels from polymerizable-group-capped ZnS capping high ME content (ZnS$_2$) are transparent at hydrated and dried state. Its transmittance slightly decreases with the increment of ZnS$_3$ content especially in the wavelength between 300 nm and 400 nm as shown in Figure 2(c). The transmittance loss is mainly ascribed to the light scattering of the particles in nanocomposites. The relation between light intensity and nanoparticle size abides by Rayleigh’s law as follows:

$$I = I_0 \exp \left[ -\frac{3\phi l R^4}{4\lambda^4} \left( \frac{n_{\text{NP}}}{n_p} - 1 \right) \right],$$

where $I_0$ and $I$ are the intensity of the incident and scattered light, $\phi$ is the volume fraction of the particles, $L$ means the thickness of the nanocomposite film, and $\lambda$ is the wavelength of the incident light. $n_{\text{NP}}$ and $n_p$ are the refractive index of nanoparticles and the polymer matrix in the nanocomposite films, respectively. The nanocomposite films should be highly transparent when the nanoparticle size is less than 40 nm. The size of our ZnS$_{1-3}$ is about 3 nm, far less than 40 nm, and the thickness is about 160 $\mu$m. The RI values of polymeric matrices of PDMA and ZnS nanocrystal are about 1.5 and 2.5, respectively. The volume fraction of the ZnS nanoparticles is less than 20%. It can be calculated that the loss of light intensity from particle scattering should be less than 1% according to the Rayleigh’s law. Therefore, it can be concluded that ZnS$_3$, nanoparticle aggregates in the nanocomposite films; although, ZnS$_2$ particles are also pendant on polymer chain by copolymerization [24].

RI value of hydrated hydrogels is highly correlated with its water content [25] and has been widely used to estimate its water loss in contact lens application [26]. For swollen hydrogel, its RI value is usually low because of the low RI value of polymeric matrices and water in the hydrogel. As shown in Figure 2(d), the RI value increases with improved ZnS content, which can be ascribed to decreased water content (Figure 2(d)) and improved high RI ZnS NPs content. In the hydrogels with high water content, polymer backbone has usually little influence on its RI value. Therefore, its RI value can be calculated according to their water content [27]. The hydrogel by copolymerization between polymerizable-group-capped ZnS has a higher RI value compared to reported hydrogels with a similar water content, e.g., the RI value of hydrogel with 30 wt\% ZnS (ESR: about 70\%, RI: about 1.39) is higher than that of Neficon A hydrogel (ESR: 69\%, RI: 1.38); the RI value of hydrogel with 50 wt\% ZnS NPs (ESR: about 47\%, RI: 1.46) is higher than that of polymacon hydrogel (water content: 38.6\%, RI: 1.43). All these indicate that ZnS NPs in the nanocomposite hydrogels play a more important role on increasing RI value than decreased water content in the hydrogels. Moreover, it is unexpected that the RI value of the hydrogels compositing ZnS$_3$ is higher than that of the hydrogel compositing a same content of ZnS$_2$; although, the ZnS$_3$ shows a higher refractive index than ZnS$_3$ at the same mass concentration in DMF as shown in Figure 1(g). This can be explained by its more serious microphase separation of ZnS$_3$ compositing hydrogels, which results in a higher water content as shown in Figure 2(d).

All hydrogels with different ZnS content shows high elastic state as shown in Figures 2(e) and 2(f), and their Young’s modulus increasing as improved ZnS content, indicating polymerizable-group-capped ZnS act as cross linker in the hydrogels. Moreover, the hydrogels with ZnS$_2$ shows better mechanical properties, e.g., higher tensile strength and Young’s modulus, longer elongation at break.

3.3. “One-Step” Route for Pure DMA-Type Nanocomposite Hydrogels. In order to composite ZnS NPs capping low amount of ME (ZnS$_1$) into transparent nanocomposite film, a novel and simple “one-step” route was developed. In the route, ZnS NPs capping different amount of ME were directly dispersed in a mixture of DMF and DMA, and then, functional monomer of GMA and initiator of AIBN were added and dispersed by ultrasound after every addition. The polymerizable solution can be polymerized by a simple thermal curing process. We suppose the reaction mechanism as illustrated in Figure 3(a). Radical polymerization undergoes a relative slow chain initiation process [28], in which GMA is linked on ZnS surface under the catalysis of solvent DMF to obtain polymerizable ZnS that then copolymerized with monomer of DMA. To confirm further the reaction mechanism, the polymerization solution without initiator of AIBN was placed in 50° C oven for 3 h, and then, ZnS NPs were precipitated in excess ethanol, and its chemical structure was confirmed by $^1$H NMR spectra. As shown in Figure 3(b), obvious carbon-carbon double bond peaks, locating at $\delta =$ 6.11 and 5.57 ppm, were observed, confirming the grafting reaction mechanism of GMA on ME-capped ZnS surface in DMF solution.

It can be seen that the nanocomposite film compositing ZnS NPs capping low amount of ME (ZnS$_1$) shows even a higher transmittance than that of the nanocomposite compositing ZnS NPs capping high amount of ME (ZnS$_2$) as shown in Figures 4(a) and 4(c). Moreover, it is interesting that the nanocomposite films with 60 wt\% and 70 wt\% ME-ZnS show higher transmittance than that of the nanocomposites with 40 wt\% and 50 wt\% ME-ZnS. The mechanical properties of “one-step” nanocomposites were shown in Figures 4(d)–4(f). Obviously, the tensile strength and Young’s modulus are higher than that of the copolymeric DMA-type nanocomposites from polymerizable-group-capped ZnS (Figures 2(e) and 2(f)), indicating higher crosslink density of “one-step” nanocomposites ascribe to the chemical link of two GMA molecules due to epoxy group of a GMA molecule reaction with the hydroxyl group of the other GMA molecule that is formed from GMA link onto ME-ZnS surface.

High RI ZnS/PDMA hydrogels have been reported [12], and its RI can be regulated from 1.58 to 1.70 in the dry states. Obviously, dry ZnS/PDMA hydrogels fabricated by “one-step” route can achieve a higher RI value above 1.73 by compositing more amount ZnS NPs. Moreover, dry “one-
Figure 2: Pure DMA-type nanocomposite films from copolymerization between polymerizable-group-capped ZnS and the monomer of DMA. Photograph of hydrated nanocomposite hydrogels immersed in distilled water (a). Transmittance of the hydrated nanocomposite hydrogels compositing ZnS₂ (b) and ZnS₃ (c). Water content and refractive index of hydrated nanocomposite hydrogels (d). Stress-strain curve of hydrated nanocomposite hydrogels compositing ZnS₂ (e) and ZnS₃ (f).
step” ZnS$_2$ nanocomposite film shows a higher RI value about 1.764 than ZnS$_1$ and ZnS$_3$ nanocomposite films (about 1.73) as shown in Figure 4(g), indicating a contrary regulation of pure ZnS and capping agent on the RI value. Higher RI films (about 1.645) with low ZnS content (less than 50 wt%) can also be achieved by compositing ZnS NPs in high RI polystyrene. However, styrene monomer only can be used as comonomer of DMA monomer due to phase separation. There is no obvious difference among the water content of three kinds of hydrated nanocomposite with a similar ZnS content, and the typical relation between water content/refractive index and free ME-ZnS content was shown in Figure 4(h). It seems that “one-step” nanocomposite hydrogels show lower water content than copolymeric hydrogels from polymerizable-group-capped ZnS (Figure 2(d)), resulting in a higher refractive index. This can be also ascribed to its higher crosslink density.

3.4. “One-Step” Route for DMA-Type Copolymeric Nanocomposite Hydrogels. To observe the applicability of “one-step” route for transparent nanocomposites, ZnS NPs capping different amount of ME were also incorporated into poly(HEMA-DMA) based copolymer matrices (DMA-type copolymer nanocomposite hydrogels). We found that the functional monomer of GMA plays an important role for its transmittance, e.g., the DMA-type copolymeric nanocomposite hydrogel, without adding functional monomer of GMA, exhibits serious irregular nanoparticle aggregation. This can be confirmed by its TME photography and its opaque appearance as an inserted image as shown in Figure 5(a). However, “one-step” route greatly overcomes microphase separation; although, only a small quantity of functional monomer was added. The uniform dispersibility of ZnS particles can be confirmed by their transparent appearance and TEM photography (Figure 5(b)–5(d)). Our previous study has shown that it is difficult to disperse more than 10 wt% ME-ZnS NPs in poly(HEMA-DMA) copolymeric matrices to prepare transparent nanocomposites [17]. Therefore, it can be concluded that the reaction between GMA and ME-capped ZnS is rapid enough that major ME-capped ZnS is conversed into polymerizable ZnS in the chain initiation process, ensuring almost all ZnS can be pendant on polymer chain in the next rapid polymerization process.

The transmittance of copolymer nanocomposite hydrogels was shown in Figures 6(a)–6(c). It can be seen that all the copolymer nanocomposite hydrogels show even a higher transmittance than pure DMA-type “one-step” nanocomposite hydrogels. This may be ascribed to its higher crosslink density. In the polymerizable solution composed of ME-capped ZnS NPs, the functional monomer of GMA, polymerizable monomer of HEMA and DMA, at least three kinds of reaction from opening epoxy of GMA monomer can be predicted: grafting reaction of GMA on ME-capped ZnS surface; linkage reaction between GMA and HEMA; chemical reaction between GMA and the products of the above two kinds of reactions. The latter two reactions will produce short chain
Figure 4: Continued.
crosslinker, resulting in increased crosslink density in the nanocomposite films.

All hydrogels with different ZnS content show also high-elastic state as shown in Figures 6(d)–6(f); although, the elongation at break is lower than that of pure DMA-type nanocomposite hydrogels from polymerizable-group-capped ZnS (Figures 2(e) and 2(f)). Young’s modulus increasing as improved ZnS content, and the hydrogel with high ZnS content is obviously higher than that of poly(HEMA-DMA) nanocomposite hydrogels from polymerization between polymerizable-group-capped ZnS and the monomer of DMA and HEMA as our previous report [17]. This further confirms its higher crosslink density as mentioned above. The hydrogels with ZnS show higher tensile strength and Young’s modulus, longer elongation at break than the hydrogels with ZnS1 or ZnS2.

The RI value and water content of the nanocomposite hydrogels after equilibrium in distilled water were shown in

![Figure 4](image-url)

**Figure 4:** Pure DMA-type “one-step” nanocomposite hydrogels. Transmittance of the hydrated hydrogels compositing ZnS1 (a), ZnS2 (b) and ZnS3 (c). Stress-strain curves of hydrated nanocomposite hydrogel compositing ZnS1 (d), ZnS2 (e) and ZnS3 (f). RI value of dried pure DMA-type nanocomposite films (g). Water content and RI value of hydrated nanocomposite hydrogels (h).

![Figure 5](image-url)

**Figure 5:** TEM photography of DMA-type copolymeric nanocomposite hydrogels and corresponding appearance as insert image. Hydrogels physically embedding 50 wt% ZnS capping high amount of ME (ZnS3) without addition of functional monomer of GMA (a). Hydrogels compositing 50 wt% ZnS1 (b), 50 wt% ZnS2 (c), 50 wt% ZnS3 (d) by “one-step” route characterized by adding functional monomer of GMA.
Figure 6: Continued.
Figure 6(g). There is no obvious difference was observed for the water content of three kinds of hydrogels. The incorporation of ZnS NPs in the hydrogel increases its refractive index, for example, the RI value reaches 1.52 when ZnS content is 60 wt%. Moreover, similar to the RI value of pure DMA-type nanocomposite hydrogels, the nanocomposite film with ZnS2 possesses also higher RI value than the nanocomposite with ZnS1 and ZnS3.

3.5. "One-Step" Route for DMA-Free Nanocomposites. The route, "one-step" fabrication of transparent nanocomposites, was further used to prepare DMA-free nanocomposites (PMMA-based nanocomposites). We found the DMA-free nanocomposite is opaque using a similar process with DMA-type nanocomposites. This indicates that a small amount of ME-capped ZnS do not still react with GMA in chain initiation process. Therefore, an additional heat treatment (50°C, 3 h) was performed after the addition of functional monomer of GMA in ZnS DMF solution. The polymerization solution can be cured to obtain transparent nanocomposites at 80°C. This indicates that ZnS NPs can be uniformly dispersed in DMA-free polymeric nanocomposites by a simple adjustment of the "one-step" route. In addition, we also decreased the time of reaction solution in 50°C oven to 1 h and 2 h. However, the resulting nanocomposites were still opaque, and the transmittance increases with extending the time of thermo-treatment. It seems that it is easier to synthesize transparent DMA-type copolymeric nanocomposites than DMA-free nanocomposites, and there is no additional process was set for the grafting reaction. This is mainly ascribed to the monomer of DMA that can play roles as both ligand and solvent effect, resulting effectively disperses and stabilizes ZnS particles. Moreover, it can also be concluded that part ME-capped ZnS was not pendant on polymeric matrices in DMA-type copolymeric nanocomposites, in which it was dispersed and stabilized by DMA component. The PMMA-based nanocomposite film composing ZnS capping 33.01 wt% ME (ZnS2) shows also an obvious higher RI value than that compositing ZnS capping higher and lower amount of ME as shown in Figure 7. This is also consistency with the data of pure DMA-type nanocomposites. Poly(methyl methacrylate) is a key polymer component for nanocomposite filling quantum dots because of its unique properties including excellent transmittance from the near UV that helps more energy can be absorbed by quantum dots under UV irradiation, maintaining the photophysics associated with the photoexcited states of the crystalites in the nanocomposites. Herein, the PMMA-based nanocomposites by the novel "one-step" route also show a good photoluminescence property under UV-irradiation as shown in Figure 7.

4. Conclusions

The amount of capping mercaptoethanol on ZnS surface can be controlled by adjusting the amount of mercaptoethanol added in the synthesis process, and increased capping amount helps to decrease the size of ZnS nanocrystals. The apparent refractive index of ZnS with middle capping amount of ME (27.7 wt%) shows a higher RI value than that of ZnS NPs with lower (23.6 wt%) or higher (33.8 wt%) capping amount of ME, ZnS NPs capping different amount of ME were successfully composited into polymeric matrices by a simple "one-step" thermocuring method characterized by the addition of a small amount of functional monomer of GMA. Epoxy group of GMA monomer can be easily opened and chemically grafted on ME-capped ZnS surface under the catalysis of solvent DMF in the relative slow chain initiation process of the polymerization, and then copolymerized with other monomers. By the route, ZnS
nanocrystals capping different amount of ME can be composited into pure DMA-type, DMA-type copolymer, and DMA-free polymeric matrices to prepare transparent and high refractive index nanocomposite films. The nanocomposite films compositing 80 wt% ZnS nanocrystals capping middle amount of ME shows a higher RI value (1.764) than the nanocomposites composing the same amount of ZnS nanocrystals capping higher or lower amount of ME, indicating that the RI value of resulting nanocomposites can be improved by optimizing the capping amount on ZnS surface. The route is an obvious advantage over reported methods because of not only simple and universal but also higher refractive index and transparent nanocomposite films.

Data Availability
All data are available from the corresponding author by request.

Conflicts of Interest
The authors declare no conflict of interest.

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