Fabrication of redispersible silica nanoparticles by a facile one-step one-pot approach

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Abstract: In this study, special redispersible silica nanoparticles (SNP) was synthesized by a facile one-step one-pot approach. The synthesized amorphous SNP had a spherical and uniform particle morphology with a diameter of 30 nm and exhibited high monodispersity. The characteristics of the particles were investigated by Fourier transform infrared spectroscopy and thermogravimetric analysis. The results revealed that there were a large number of methyl groups grafted on the surface of the SNP, and the organic materials content in the as-synthesized SNP was approximately 10%. After a simple ultrasonic process, the SNP solid powder showed excellent stable and transparent redispersion in various organic solvents. In addition, the results revealed that the different polarities of the solvents had a significant effect on the visible light transparency of the SNP redispersion. SNP showed the best redispersion performance in N-methyl-pyrrolidone, exhibiting a transmittance of more than 80% at 600 nm at a high concentration of 25 wt%. The synthesis process developed in this study is simple, inexpensive, and can be used in the industrial application of SNP.

1 Introduction

In recent decades, nanotechnology has attracted significant scientific interest owing to the promising potential applications of nanoparticles and nanotechnology in various fields. [1-2] Silica nanoparticles (SNP) such as precipitated silica, fumed silica, and colloidal silica are inorganic chemical materials with an average particle size of less than 100 nm and are widely used in various fields including the fields of functional coatings, catalysis, nanocomposites, adsorption, biomedicine, etc. [3-7] SNP is usually added to solvents, polymers, and other matrices as special fillers in the preparation processes of functional coatings or nanocomposites. However, the agglomeration of SNP in the matrix due to its small particle size and high surface energy limits its further application. The agglomeration of SNP can be attributed to its poor dispersion property in the matrix, which in turn leads to the degradation of the physical and chemical properties of the matrix, such as a reduction in the transparency and strength of resin. In summary, the particle agglomeration caused by the poor dispersion ability of SNP significantly reduces the functional effect of its application. Consequently, several surface modification steps have been introduced to improve the dispersibility of synthesized SNP [8-10]. However, the high cost of these surface modification steps and their inability to fundamentally improve the dispersion of SNP in the application matrix has limited the effective application of these methods. Generally, the modification of SNP with a suitable modifier and modification process improves the dispersibility of SNP in specific solvents or polymers. However, agglomeration occurs when SNP is redispersed in another solvent or polymer matrix. In addition, due to the extremely high surface activity of SNP, synthesized or modified SNP are prone to secondary agglomeration during the drying process, which reduces the redispersibilities of dried SNP in solvents or polymers. [11-12] SNP with good redispersibility can be packaged as solid powders to avoid storage and transportation safety problems caused by packaging the SNP in the form of dispersion in solvent. SNP solid powders with good redispersibility can be directly redispersed in different solvents or polymer matrices for practical applications, thus making them more flexible and easy to use. However, studies on the redispersibility of SNP and the preparation of redispersible SNP are rare.

In this study, we report a facile approach to fabricate redispersible SNP by a one-step one-pot method without additional surface modification treatment. The dried SNP solid powder showed excellent redispersible properties and was easily redispersed in various organic solvents to obtain highly stable and transparent SNP redispersion.

2 Experiment

2.1. Materials
Tetraethoxysilane (TEOS, AR), ethanol (ETOH, AR),
tetrahydrofuran (THF, AR), ethyl acetate (EA, AR),
N-methyl-pyrrolidone (NMP, AR), and ammonium
hydroxide (NH₄OH, 25–28%, AR) were purchased from
Damao Chemical Reagent Factory (Tianjin, China).
Dimethoxydimethylsilane (DDS, AR) was purchased
from Hangzhou Jessica Chemical Co., Ltd (Hangzhou,
China). The commercial SNP powder was purchased
from Wacker Chemie Co. Ltd (Germany). All chemicals
were used as received without any further purification.

2.2. Synthesis of the redispersible SNP

The redispersible SNP was synthesized by a one-step
one-pot method based on a modified Stöber protocol.[13]
To explain in brief, ETOH (150 g), NH₄OH (10 g),
TEOS (10 g), and DDS (2.88 g) were placed in a
three-neck round-bottom flask, after which the mixture
was stirred vigorously at 38 °C for 6 h. After the reaction,
the redispersible SNP dispersion in ETOH was obtained.
After drying the dispersion in an oven at 60 °C for 4 h,
the redispersible SNP solid powder was obtained.

2.3. Characterization

X-ray powder diffraction (XRD) analysis of the
as-synthesized SNP was conducted (DX-2700BH,
Haoyuan). The chemical structure of the as-synthesized
SNP was examined using a Fourier transform infrared
(FTIR) spectrometer (TENSOR II, Bruker). Transmission
electron microscopy (TEM, F30, FEI Tecnai) was
conducted to analyze the morphology of the
as-synthesized SNP. The thermal stability of the
as-synthesized SNP was determined through
thermogravimetric analysis (TGA, STA 449 F5, Netzsch).
The optical transparency of the SNP dispersions was
examined by UV–visible (UV–vis) spectroscopy
(UV-1800, Shimadzu).

3 Results and discussion

3.1. Characterization of the as-synthesized SNP

The XRD pattern of the as-synthesized SNP is shown in
Figure 1. The XRD diffraction pattern reveals that the
synthesized SNP possessed an amorphous-like structure.
TEM was carried out to investigate the morphology and
the size of the SNP, and the TEM images are shown in
Figure 2. The high- and low-magnification TEM images
reveal that the well-dispersed SNP had a spherical and
uniform particle morphology with a diameter of
approximately 30 nm.

3.2. Characterization of the as-synthesized SNP

Figure 3 shows the FTIR spectra of the
as-synthesized SNP. As seen, an absorption band
appeared at approximately 1095 cm⁻¹, which could be
attributed to the vibrations of Si-O-Si bonds.[14]
Additional absorption peaks were observed at ~2865
cm⁻¹ and ~1460 cm⁻¹, which could be attributed to the
presence of -CH₃ group.[15] The absorption peaks
observed at ~860 cm⁻¹ corresponded to the characteristic
vibrations of the Si-CH₃ bonds.[16] The observed
characteristic bands of the -CH₃ and Si-CH₃ groups
indicated the successful synthesis of the SNP grafted with
a large number of methyl groups. Figure 4 shows the TG
curve of the as-synthesized SNP. The TG curve revealed
that the organic matter content of the SNP was about 10
wt%, and was completely decomposed at approximately 600 °C.

Figure 3. FTIR spectra of the as-synthesized SNP.

Figure 4. TGA curve of the as-synthesized SNP.

3.2. Redispersible properties of the as-synthesized SNP

The obtained SNP was stably dispersed in ETOH, and a light-blue transparent dispersion was observed, as shown in Figure 5 (a), indicating that the SNP dispersion in ETOH exhibited excellent transparency. After drying the SNP dispersion in ETOH in an oven at 60 °C for 4 h, a white SNP solid powder was obtained, as shown in Figure 5 (b). To investigate the redispersibility of the as-synthesized SNP, 1 wt% SNP content of different SNP redispersion was prepared by mixing 0.1 g of the dried SNP solid powder with 9.9 g of various solvents, after which ultrasonication was performed for 1 min to redisperse the SNP in the solvent. For the redispersibility investigation, we selected a variety of solvents that are widely used in industries including alcohol solvents (ETOH), ether solvents (THF), ester solvents (EA), and ketone solvents (NMP). As shown in Figure 6(a), after the simple ultrasonic dispersion process, transparent SNP redispersion was obtained in all solvents, indicating that the as-synthesized SNP exhibited excellent redispersibilities in various solvents. For comparison, the redispersibility of a commercial SNP powder in ETOH with a concentration of 1 wt% was investigated. As shown in the photo of the suspension in Figure 6(b), the transparency of the suspension was very low, indicating that the poor redispersibility of the commercial SNP.

Figure 5. Photos of (a) as-synthesized SNP dispersion in ethanol (ETOH), and (b) SNP solid powder after drying.

Figure 6. Photos of (a) as-synthesized SNP redispersion in different solvents, and (b) purchased SNP redispersion in ETOH.
Figure 7 shows the UV–vis transmittance spectra of the as-synthesized SNP redispersion in different solvents and that of the purchased SNP redispersion in ETOH. The concentration of SNP in all the redispersion was 1 wt%. The light transmittance of the different redispersion were tested, and the results revealed that the SNP redispersion in NMP showed the highest transmittance with the light transmittance at 600 nm exceeding 90 %, whereas, the transparency of the commercial SNP redispersion in ETOH was significantly lower. In addition, there were obvious differences in the transparencies of the SNP redispersion in different solvents, indicating that the SNP has different redispersibilities in different solvents. Previous studies have reported that nanoparticles aggregates in a dispersion medium significantly increase the intensity of scattered light as described by Rayleigh’s law, which would in turn induce a considerable decrease in the transparency of the dispersion. Owing to the different polarities of different solvents, which affects the aggregation state of SNP in the solvent, the degrees of light scattering of the dispersion differ, which in turn affected the transparency of the redispersion. Generally, the redispersion obtained by dispersing the as-synthesized SNP solid powder in different solvents have high visible light transparency, which is consistent with the results in Figure 6, and the light transmittance of all the redispersion at 600 nm exceeded 50 %. In contrast, the light transmittance of the purchased SNP redispersed in ETOH was significantly low, and that of the suspension at 600 nm was less than 30 %, indicating the low redispersibility of the purchased SNP in ETOH. It is worth noting that the redispersion of SNP obtained by redispersing the SNP solid powder in different solvents showed excellent stabilities. After storage at room temperature for 7 days, no delamination or precipitation occurred in the transparent SNP redispersion.

Figure 7. UV–vis transmittance spectra of as-synthesized SNP redispersion in (a) ETOH; (b) THF; (c) EA; (d) NMP, and (e) the purchased SNP redispersion in ETOH.

The results of the above analysis indicated that the as-synthesized SNP showed the best redispersibility in NMP. Therefore, we investigated the redispersibilities of different concentration of the as-synthesized SNP in NMP using a simple ultrasonic method. The photos of the redispersion are shown in the Figure 8. As seen, the redispersion with SNP concentrations of 1 wt% and 10 wt% exhibited excellent transparencies, whereas the redispersion with 25 wt% SNP concentration exhibited a slightly poorer transparency. The UV–vis transmittance spectra of the as-synthesized SNP redispersion in NMP at different concentrations is shown in Figure 9. The transmittance of the redispersion with 1 wt%, 10 wt%, and 25 wt% SNP content at 600 nm was 91.6 %, 88.1 %, and 83.1 %, whereas the transmittance of pure NMP at 600 nm was 92.3 %. This result indicated that the transmittance of the redispersion decreased gradually with an increase in the SNP content, however, they were all higher than 80 %, and high transparencies were maintained.

Figure 8. Photos of the as-synthesized SNP redispersed in NMP at different concentrations.

Figure 9. UV–vis transmittance spectra of the as-synthesized SNP redispersion in NMP at different concentrations.

4 Conclusion

In this study, we reported the synthesis of SNP by a simple one-step one-pot method. The obtained SNP with a diameter of approximately 30 nm exhibited a stable dispersion in ETOH. By drying the dispersion in a traditional oven, a redispersible and user-friendly SNP solid powder was obtained. The dried SNP solid powder exhibited stable redispersion in various organic solvents including ETOH, THF, EA, and NMP. In addition, the SNP powder exhibited excellent redispersion in NMP even at a high concentration of 25 wt%. The preparation method proposed in this study can provide technical insights for the development of new SNP-based functional coatings and nanocomposites.
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