Semiconductor nanoparticle-based hydrogels prepared via self-initiated polymerization under sunlight, even visible light

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Since ancient times, people have used photosynthesized wood, bamboo, and cotton as building and clothing materials. The advantages of photo polymerization include the mild and easy process. However, the direct use of available sunlight for the preparation of materials is still a challenge due to its rather dilute intensity. Here, we show that semiconductor nanoparticles can be used for initiating monomer polymerization under sunlight and for cross-linking to form nanocomposite hydrogels with the aid of clay nanosheets. Hydrogels are an emerging multifunctional platform because they can be easily prepared using solar energy, retain semiconductor nanoparticle properties after immobilization, exhibit excellent mechanical strength (maximum compressive strength of 4.153 MPa and tensile strength 1.535 MPa) and high elasticity (maximum elongation of 2784%), and enable recyclable photodegradation of pollutants. This work suggests that functional nanoparticles can be immobilized in hydrogels for their collective application after combining their mechanical and physiochemical properties.

Hydrogels, which have a three-dimensional porous network, have attracted significant attention due to their use as biomaterials and model matrices for biological studies\textsuperscript{1–3}. Most hydrogels, whether covalently or non-covalently cross-linked, are mechanically weak and exhibit poor elasticity and a lack of physiochemical functions\textsuperscript{4}. Pioneering double network hydrogels can achieve certain mechanical requirements using a typical two-step polymerization\textsuperscript{5,6}. A nanocomposite (NC) hydrogel is another original material that can achieve ultra-high mechanical properties due to the multiple non-covalent effects between clay nanosheets (Clay-NS) and polyacrylamide chains\textsuperscript{7–10}. Recently, researchers have developed a supramolecular hydrogel by mixing Clay-NS with dendritic polymers using a multiple salt bridge\textsuperscript{11}. However, the preparation of these tough hydrogels requires complicated molecular designs or preparation processes and does not permit the combination of the strengths and functions of the individual components. Until now, semiconductor nanoparticles (NPs) could only be immobilized in weak polymer hydrogels by physical mixing or by vinyl-modified covalent cross-linking\textsuperscript{12–14}.

Here, we chose a sunlight-initiated approach to prepare multifunctional NC hydrogels using semiconductor NPs as the initiator and backbone; this approach combines the mechanical properties and physiochemical performances of the materials into a single platform. Our system consists of four components: water, watersoluble semiconductor nanoparticles (ZnO, TiO\textsubscript{2}, Fe\textsubscript{3}O\textsubscript{4}, SnO\textsubscript{2}, ZrO\textsubscript{2}, CdSe, or CdTe), N,N-dimethylacrylamide (DMAA), and Clay-NS. Semiconductor NPs, including TiO\textsubscript{2}, ZnO, and CdTe, have been shown to undergo photo-initiated polymerization during heterogeneous polymer synthesis\textsuperscript{15–18}. These inorganic initiators have rarely been used to prepare polymer gels under light. The DMAA monomer is a water-soluble acrylamide derivative, and the corresponding PDMAA exhibits stability against hydrolytic effects and temperature changes. We recently observed that sunlight irradiation can increase the viscosity of a mixture of semiconductor NPs and DMAA due to polymerization. The insufficient number of cross-linking points on the NP surfaces determine the final viscous situation other than three-dimensional hydrogel. This phenomenon suggests the possibility of preparing semiconductor NP-based hydrogels under sunlight by introducing additional cross-linking points. Based on the literature, we introduced Clay-NS, a layered structured mineral with a high specific surface\textsuperscript{19}, to the mixture as a second building block.
Results

Preparation and characterization. A dispersion of semiconductor NPs, DMAA, and Clay-NS (Figure 1a) was used as a precursor. Of these materials, the semiconductor NPs can initiate the polymerization under sunlight to form inorganic-PDMAA brushes (Figure 1b) and a non-covalent NC hydrogel network via the further cross-linking of the brushes (Figure 1c)⁷,⁸,²⁰,²¹. As described in the Methods section, a typical ZnO nanoparticle-based nanocomposite hydrogel (ZnO-NC gel) was prepared from an aqueous solution (Figure 1d) with 0.2% ZnO, 5.0% DMAA, and 5.0% Clay-NS (Supplementary movie). The final ZnO-NC gel was translucent and tough after approximately 1 h of sunlight irradiation (Figures 1e and 1f and Supplementary Figure S1). As shown in Table 1 and in Supplementary Figure S2, simple sunlight irradiation yielded elastic and functional NC hydrogels with various nanoparticles, with the exception of ZrO₂ and Fe₂O₃ nanoparticles. We used a xenon lamp with an AM 1.5 G filter to mimic sunlight (similar spectrum composition and equivalent intensity). All functional nanoparticles, with the exception of ZrO₂ and Fe₂O₃, can be used in similar preparations under artificial sunlight. For quantum dots (CdSe and CdTe) with a low bandgap, an elastic hydrogel was prepared via visible light irradiation with a wavelength over 420 nm (Supplementary Figure S3), which can be produced by cutting off the UV light of artificial sunlight. A morphology analysis confirms the fabrication of a functional NC hydrogel using three components. SEM analysis of a supercritically treated sample (Figure 1g) reveals irregular pores with sizes ranging from several tens to several hundreds of nanometres and highly dispersed brushes 100–300 nm in size. The porous aerogel was tough rather than fragile, in contrast to traditional inorganic aerogels. Bio-cutting TEM analysis (Figure 1h) reveals that aggregated NPs, with sizes of approximately several tens of nanometres, and individual Clay-NS particles, which have the same size as in solution, were dispersed in the gel (Supplementary Figure S4).

In our functional hydrogels, the photoinitiator semiconductor NPs were similar to organic initiators²²–²⁴, which enable spatial/temporal control and a mild curing process relative to the traditional preparation of most hydrogels²²–³⁴. The important advantages of this method include the use of inexhaustible sunlight as a substitute for an ultraviolet generator. The mild light irradiation initiated by successful semiconductor NPs can lead to the effective and complete polymerizations of DMAA. Other acrylamide derivatives, such as N-isopropylacrylamide, can also be polymerized via the irradiation effect. Time-dependent¹H NMR analysis of the TiO₂ precursors exhibits an approximately 99% DMAA conversion rate after 1 h of irradiation under sunlight and under artificial light (Figure 2a and Supplementary Figures S5 and S6). The clear gelation of these precursors during irradiation occurs concurrently with a rapid increase

Figure 1 | Proposed mechanism, photographs, and SEM and TEM images of the functional ZnO-NC hydrogel. (a–c) Proposed mechanism of the functional NC hydrogels under sunlight. The semiconductor NPs, monomer, and Clay-NS are homogeneously dispersed in water (a). When exposed to sunlight, polymerization is initiated by the semiconductor NPs (b). Next, the NPs and Clay-NS are cross-linked to develop a 3D network (c). (d–f) Optical images of the hydrogelation process. A mixture solution of ZnO nanoparticles, DMAA, and Clay-NS (d). Hydrogelation after 1 h of irradiation (e). The elastic ZnO-NC hydrogel (f). (g) SEM image of the ZnO-NC hydrogel after supercritical drying. (h) TEM image of the ZnO-NC hydrogel after a bio-cutting treatment.
in monomer conversion after 15 min (Figure 2a), which accompanies by a sharp increase in the decay time and a decrease in the correlation coefficient and the dynamic light scattering (DLS) spectrum (Figure 2b). Comparison of the small angle X-ray scattering (SAXS) spectra at various irradiation times reveals the formation of a regular pore structure (Supplementary Figure S7). If a light source with a higher intensity is used, the gelation could be completed more rapidly; this phenomenon was investigated using an in situ rheological measurement with strong ultraviolet light (Supplementary Figure S8). For a visible light-irradiated CdTe system, a greater than 95% DMAA conversion rate can be achieved within 1.5 h (Supplementary Figure S9). Other semiconductor NPs, such as ZnO, SnO2, CdS, and CdSe, also permit polymerization irradiation under various light sources. For the exceptions, the ZrO2 and Fe2O3 nanoparticles, the NMR data indicate a 5% DMAA conversion rate after the same irradiation time (Supplementary Figure S10). The large bandgap of ZrO2 (4.95 eV) is most likely the main barrier to electron-hole excitation. Although Fe2O3 nanoparticles can initiate photo-polymerization in organic solvents, the free radicals are easily inhibited by the reduction of Fe3+ and water molecules in an aqueous solution. For use in our self-initiated system, a semiconductor NP should have a moderate bandgap, which is easy to be excited by sunlight to form active radicals.

**Mechanical properties.** As shown in the supporting movie, our semiconductor NP-NC gels exhibits high toughness and elasticity. Table 2 lists the detailed mechanical data for the TiO2-NC hydrogels with various semiconductor NP and Clay-NS proportions. A three-component hydrogel could be formed with a proper concentration range of NPs (0.015% to 1% for TiO2, ZnO, SnO2, CdSe, and CdTe), DMAA (5% to 10%), and Clay-NS (5% to 20%). As shown in Figure 3a, the TiO2-NC hydrogels with 0.2% TiO2 (5% DMAA and 5% Clay-NS) exhibit a very large elongation of close to 27 times the original length. The minimum proportion of TiO2 for complete

![Figure 2](https://www.nature.com/scientificreports/)

**Figure 2 | Gelation process of the TiO2-NC hydrogels.** (a) The conversion of the monomer by TiO2 under a xenon lamp with an AM 1.5 G filter. The precursor consisted of 3% DMAA, 3% Clay-NS, 94% D2O, and 0.1% TiO2. (b) DLS analysis of the sample during irradiation. The precursor solution containing 3% DMAA, 3% clay, 94% H2O, and 0.1% TiO2 was irradiated for different times. (c) The EPR spectrum of the precursor consisting of TiO2 and DMAA under UV irradiation. (d) The molecular weight distribution of the PDMAA polymers formed from the aqueous mixture of TiO2 and DMAA after 1 h of sunlight irradiation.

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| NP        | Absorption edge (nm) | Band energy (eV) | Solution colour | Sun lighta | Xenon lampb | Visible lightb |
|-----------|-----------------------|------------------|-----------------|------------|-------------|----------------|
| Fe2O3     | 919.1                 | 1.35             | Red brown       | No         | No          | No             |
| CdSe      | 570.4                 | 2.17             | Yellow          | Gel        | Gel         | Gel            |
| CdTe      | 559.8                 | 2.22             | Yellow          | Gel        | Gel         | Gel            |
| ZnO       | 434.2                 | 2.85             | Translucent     | Gel        | Gel         | No             |
| TiO2      | 414.8                 | 2.99             | White           | Gel        | Gel         | No             |
| SnO2      | 384.9                 | 3.22             | Translucent     | Gel        | Gel         | No             |
| ZrO2      | 250.3                 | 4.95             | White           | No         | No          | No             |

aThe average intensities of sunlight and of the xenon lamp with an AM 1.5 G filter at 365 nm are both approximately 2.0 mw cm-2.
bThe visible region of the xenon lamp was obtained by cutting off the light at λ < 420 nm.
conversion is 0.05%, which yields an approximate elongation of 2740%. Figure 3b shows that the relative elongation decreases slightly to 2720%, 2784%, and 2688%, when the proportion of TiO2 increases to 0.1%, 0.2%, and 0.4%, respectively. The additional cross-linking density from TiO2 nanoparticles should correspond to a decrease in the elongation of our hydrogels along with an increase in the proportion of TiO2. As shown in Supplementary Figure S12, the ZnO-based hydrogel also exhibits decreasing elongation as the proportion of ZnO was increased. The final optimized concentration of TiO2 in our system is 0.2%, taking into account the elongation, tensile modulus, and broken strength (Figure 3b). Synthetic PDMAA with a broad distribution provides the high elasticity of our NC gel system, which is the result of the space linkage between the dispersed inorganic cross-linkers.

The tensile performances (elongation, modulus, and strength) all weakly depend on the DMAA content when the concentration is greater than 5%. A maximum elongation of 2784% occurs for the TiO2-NC gel containing 7% DMAA (0.2% TiO2 and 5% Clay-NS). We select 5% DMAA as the optimized concentration to decrease the organic component and the final cost. The tensile performance (elongation, modulus, and strength) all strongly depend on the Clay-NS content (Figure 3c and Table 2). Therefore, we attempt to use the viscous precursors to fabricate a hydrogel with up to 20% Clay-NS, which should have excellent mechanical properties due to the high inorganic concentration. The maximum tensile modulus of the functional NC gel with 20% Clay-NS (0.2% TiO2 and 10% DMAA) is 2.404 MPa. The elongation values decrease from 2688% to 1086% as the proportion of Clay-NS increase from 5–20% due to the increased cross-linking density. A maximum broken strength of 1.535 MPa is obtained for the functional NC gel with 15% Clay-NS.

For the compressive test, we found that all of the hydrogels can tolerate a compression greater than 95% and can exhibit partially rubber-like properties. The compressive performances of the hydrogels decrease slightly with increasing DMAA concentration (0.2% TiO2 and 5% Clay-NS) (Table 2). The compressive modulus and strength both strongly depend on the Clay-NS content. As shown in Figure 3d, the compressive modulus and strength at a 95% strain increase with increasing Clay-NS amount. The TiO2-NC hydrogel

| Table 2 | The mechanical properties of the TiO2-NC gels with various proportions |
| No. | Composition TiO2/DMAA/Clay-NS (%) | Tensile Modulus (MPa) | Tensile Strength (MPa) | Extensibility (%) | Comp. Modulus (MPa) | Comp. Strength (MPa) |
|------|---------------------------------|----------------------|-----------------------|------------------|---------------------|---------------------|
| 1    | 0.2/5/5                         | 0.0082 ± 0.0004      | 0.2575 ± 0.0143       | 2688 ± 73        | 0.0074 ± 0.0004     | 0.386 ± 0.0111      |
| 2    | 0.2/7/5                         | 0.0081 ± 0.0004      | 0.2525 ± 0.0125       | 2784 ± 58        | 0.0068 ± 0.0004     | 0.363 ± 0.0091      |
| 3    | 0.2/10/5                        | 0.0083 ± 0.0003      | 0.2595 ± 0.0108       | 2767 ± 80        | 0.0063 ± 0.0004     | 0.335 ± 0.0088      |
| 4    | 0.2/10/10                       | 0.167 ± 0.026        | 1.130 ± 0.074         | 2280 ± 103       | 0.177 ± 0.0111      | 2.103 ± 0.116       |
| 5    | 0.2/10/15                       | 0.507 ± 0.032        | 1.535 ± 0.089         | 1506 ± 78        | 0.610 ± 0.026       | 3.356 ± 0.143       |
| 6    | 0.2/10/20                       | 2.404 ± 0.150        | 1.075 ± 0.063         | 1086 ± 86        | 1.958 ± 0.080       | 4.153 ± 0.121       |

*Comp. = Compressive.
*Comp. strength at a maximum strain of 95%.

Figure 3 | Mechanical properties of the TiO2-NC hydrogels with different proportions. (a) Images of the TiO2-NC hydrogel during the tensile testing process. (b) Tensile properties of the TiO2-NC hydrogels containing 5% DMAA, 5% clay, and different proportions of TiO2. (c) Tensile properties of the TiO2-NC hydrogels containing 0.2% TiO2 and different Clay-NS concentrations. (d) Compressive properties of the TiO2-NC hydrogels with different clay concentrations.
(0.2% TiO$_2$, 5% DMAA, and 20% Clay-NS) achieves a maximum of 1.958 MPa for the compressive modulus and 4.153 MPa for the compressive strength. Other semiconductor NP-NC gels, such as the ZnO system, also exhibited excellent compressive performances similar to that of the TiO$_2$ gels (Supplementary Figures S11 and S12). Another interesting mechanical property is the very rapid recovery of rheological strength that our sample exhibits after a large-amplitude oscillatory breakdown; this property is known as self-healing$^{35,36}$ (Supplementary Figure S13). This recovery is fully reproducible for three strain cycles ($\gamma$ changes as 1%-100%-1%-100%-1%-100%-1%).

**Fluorescence and photocatalytic properties.** One advantage of our systems is the remnant physicochemical properties of the semiconductor NPs after in situ immobilization. We can observe that the NPs within a tough hydrogel retain their intrinsic properties. As shown in Figure 4a, the CdTe-NC hydrogel (CdTe/DMAA/Clay-NS of 0.015%/3%/3%) has absorption and emission peaks that are analogous to those of a 0.015% quantum dot solution of CdTe. The fluorescence intensity of the CdTe-NC hydrogel is slightly higher than that of the original CdTe solution at an identical concentration due to light activation during processing$^{37}$. Hydrogels with different fluorescence intensities can be easily prepared using variously sized CdTe dots (Figures 4b–d).

Semiconductor NPs can act as photocatalysts even within a functional hydrogel. The TiO$_2$-NC hydrogel (TiO$_2$/DMAA/Clay-NS of 0.4%/5%/5%) is selected to collectively treat dyes using the adsorption-photodegradation-readsorption cycle$^{38,39}$. As drawn in Figure 4e, the TiO$_2$-NC hydrogel can enrich a large amount of methylene blue to form a dark blue hydrogel via the electrostatic effect. The first maximum absorption of methylene blue is 59.5 mg for 1 g of dried hydrogel (Figure 4f). The dark blue hydrogel in 5-fold excess of water can change to light blue after 2 h of UV (middle-pressure mercury lamp, 100 mW cm$^{-2}$) degradation via the loaded TiO$_2$. The colourless products after photodegradation easily diffuse in water. Notably, the reused TiO$_2$-NC hydrogel can readorb the methylene blue and be re-treated by the immobilized photocatalyst. The adsorbed amount of methylene blue after two recycles can reach 86.3% of the fresh hydrogel. As a control, the traditional NC hydrogel of Clay-NS and DMAA$^8$ was studied; this hydrogel could adsorb an equivalent amount of methylene blue but could not be obviously degraded under the same UV irradiation conditions (Supplementary Figure S14). Another pollutant, acid fuchsin can also be captured and degraded with the TiO$_2$-NC hydrogel (Supplementary Figure S15).

**Discussion**

With our aqueous precursors, semiconductor NPs produce the same free radical mechanism as organic initiators$^{20,21}$ due to water inhibition of the chain propagated ions. The OH radical (produced by the photogenerated hole and electron) and the hole are the two initiating species that form a DMAA radical and further propagating radicals. The competitive OH radical is the main species in our precursor with a high pH (approximately 9.0) in the absence of deliberate degassing treatment$^{15}$. The existence of a hydroxyl radical in the TiO$_2$ and CdTe base solutions is confirmed by electron paramagnetic resonance (EPR) spectroscopy after DMPO adduction; the spectroscopy data reveal a characteristic DMPO-OH radical spectrum with a 1:2:2:1

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**Figure 4** | Fluorescence and photocatalytic properties of the semiconductor NP-NC hydrogels. (a) The UV-Vis absorbance and PL spectra of the CdTe-NC hydrogel and the CdTe quantum dot solution with the same concentration of CdTe. (b–d) Images of three different CdTe-NC hydrogels under an ultraviolet lamp. The average diameters of the CdTe quantum dots are estimated to be 2.8 nm (b, green), 3.3 nm (c, yellow), and 4.0 nm (d, red). (e) The recyclable application of the TiO$_2$-NC hydrogel for the treatment of polluted water and images of the TiO$_2$-NC hydrogels at various stages are shown (the scale is 1 cm). (f) The amount of MB adsorbed by the TiO$_2$-NC gel in the three reusing cycles.
ratio (Supplementary Figures S16a and 16c). The hole initiator can directly react with non-covalent adsorbed DMAA to form a surface monomer radical15–18. The OH radicals then diffuse from the NPs and initiate polymerization near the NP surfaces in the polymers that only have hydroxyl end groups (Supplementary Figure S17a). The OH radicals then diffuse from the NPs and initiate polymerization near the NP surfaces in the polymers that only have hydroxyl end groups (Supplementary Figure S17b).

Our semiconductor NP-based NC gels, which are produced by easy sunlight-initiated polymerization and a non-covalent cross-linking effect, have not only mechanical properties that are tunable by changing the various components but also no toxic residue in comparison with organic initiators. The self-immobilized NPs within the functional NC gels exhibit inherent physicochemical properties, such as absorption and fluorescence, that are comparable to those of their original solution. The semiconductor NP-NC hydrogels can be used as a recyclable platform for the collective treatment of industrial pollutants. By coupling other applications of NPs, functional NC gels could be designed as an effective and recyclable water-rich platform for photohydrolysis and solar cells. Our semiconductor NP-based hydrogel addresses one disadvantage of hydrogels fabricated by complicated methods, that is, excellent mechanical performances but a lack of functionality.

Methods
Preparation. In a typical hydrogelation, 0.12 mL of a ZnO aqueous suspension (10%), DMAA monomer (300 mg, 0.31 mL), Clay-NS (300 mg), and water were mixed with vigorous stirring to form a 6.0 g aqueous suspension within 5 minutes (Figure 1d). The last precursor can form a ZnO-NC gel with semitransparent performance and tough elasticity via irradiation under sunlight (1 h), with an average intensity of 2.0 mW cm⁻² at 365 nm (Figure 2d). The propagating radical signal has the largest intensity at 4 min and then decreases with time (Supplementary Figure S16b). The semiconductor NPs should be immobilized within the functional hydrogels because the non-covalent cross-linking points prevent easy leakage during incubation. A mechanism for the preparation of our NC gels under sunlight irradiation is proposed as follows. First, more active holes react with the non-covalent adsorbed DMAA to form a surface monomer radical, which initiates the polymerization directly at the unmovable NP surfaces. The corresponding polymers exhibit both double bond and hydroxyl end groups (Supplementary Figure S17a). The OH radicals then diffuse from the NPs and initiate polymerization near the NP surfaces in the polymers that only have hydroxyl end groups (Supplementary Figure S17b).

Photoluminescence (PL) spectra were obtained on a UV-Vis spectrometer (Hitachi F-7000), respectively.

1. Wang, H., Yang, Z. & Adams, D. J. Controlling peptide-based hydrogelation. Mater. Today 15, 500–507 (2012).
2. Peppas, N. A., Huang, Y., Torres-Lugo, M., Ward, J. H. & Zhang, J. Physicochemical, foundational and structural design of hydrogels in medicine and biology. Annu. Rev. Biomed. Eng. 2, 9–29 (2000).
3. Gong, J. P., Katsuyama, Y., Kurokawa, T. & Osada, Y. Double-network hydrogels with extremely high mechanical strength. Adv. Mater. 15, 1155–1158 (2003).
4. Chen, Y. M., Yang, J. I., Osada, Y. & Gong, J. F. Synthetic hydrogels as scaffolds for nanopatterning endothelium cell behaviors. Chin. J. Polym. Sci. 29, 23–41 (2011).
5. Haraguchi, K. & Takehisa, T. Nanocomposite hydrogel: a unique inorganic–organic network structure with extraordinary mechanical, optical, and swelling/de-swelling properties. Adv. Mater. 14, 1120–1124 (2002).
6. Haraguchi, K., Farkworth, R., Ohbayashi, A. & Takehisa, T. Compositional effects on mechanical properties of nanocomposite hydrogels composed of poly(N,N-dimethylacrylamide) and clay. Macromolecules 36, 5732–5741 (2003).
7. Liu, Y., Zhu, M. F., Liu, X. L., Zhang, W. & Adler, H. J. High content nanocomposite hydrogels with surprising mechanical strength and interesting deswelling kinetics. Polymer 47, 1–5 (2006).
8. Wang, J. F., Lin, L., Cheng, Q. F. & Jiang, L. A strong bio-inspired layered pnp-gramm nanocomposite hydrogel. Angew. Chem. In. Ed. 41, 4676–4680 (2012).
9. Wang, Q. G., Mynar, J. L., Yoshida, M., Lee, E., Lee, M. & Aida, T. High-water-content mouldable hydrogels by mixing clay and a dendiric molecular binder. Nature 463, 339–343 (2010).
10. Satarkar, N. S., Biswal, D. & Hilt, J. Z. Hydrogel nanocomposites: a review of applications as remote controlled biomaterials. Soft Matter 6, 3566–3571 (2010).
11. Wang, C., Flynn, N. T. & Langer, R. Controlled structure and properties of thioresponsive nanoparticle–hydrogel composites. Adv. Mater. 16, 1074–1079 (2004).
12. Zhu, C. H., Hai, Z. B., Cui, C. H., Li, H. H. & Yu, S. H. In situ controlled synthesis of thermosensitive poly(N-isopropylacrylamide)/Au nanocomposite hydrogels by gamma radiation for catalytic application. Small 8, 930–936 (2012).
13. Ni, X. Y., Ye, J. & Dong, C. Kinetics studies of methyl methacrylate photopolymerization initiated by titanium dioxide semiconductor nanoparticles. J. Photochem. Photobio. A 181, 19–27 (2006).
14. Huang, Y. Z., Barber, T., Mills, G. & Morris, M. B. Heterogeneous photopolymerization of methyl methacrylate initiated by small ZnO particles. J. Phys. Chem. B, 12746–12752 (1994).
15. Nakashima, T., Sakashita, M., Nonoguchi, Y. & Kawai, T. Sensitized photopolymerization of anionic liquid-based monomer by using CdTe nanocrystals. Macromolecules 40, 6540–6544 (2007).
16. Stroyuk, A. L. et al. Photoinitiation of butylmethacrylate polymerization by colloidal semiconductor nanoparticles. J. Photochem. Photobio. A 162, 339–351 (2004).
17. Rockwood Additives Ltd., Laponite in personal care products. Laponite technical bulletin L211/01(g) (1990).
18. Perse, B. et al. Gelation mechanism of poly(N-isopropylacrylamide)-clay nanocomposite hydrogels synthesized by photopolymerization. Langmuir 24, 12627–12635 (2008).
19. Haraguchi, K. & Takada, T. Synthesis and characteristics of nanocomposite gels prepared by in situ photopolymerization in an aqueous system. Macromolecules 43, 4294–4299 (2010).
20. Hirst, A. R., Escuder, B., Miravet, J. F. & Smith, D. K. High-tech applications of self-assembling supramolecular nanostructured gel-phase materials: from regenerative medicine to electronic devices. Angew. Chem. Int. Ed. 47, 8002–8018 (2008).
21. Zhang, X. L. et al. Rational Design of a Tetrameric Protein to Enhance Interactions between Self-Assembled Fibers Gives Molecular Hydrogels. Angew. Chem. Int. Ed. 51, 4388–4392 (2012).
22. Kiyonaka, S. et al. Semi-wet peptide/protein array using supramolecular hydrogel. Nature Mater. 3, 58–64 (2004).
23. Gao, Y., Zhao, F., Wang, Q. G., Zhang, Y. & Xu, B. Small peptide nanofibers as the matrix of molecular hydrogels for mimicking enzymes and enhancing the activity of enzymes. Chem. Soc. Rev. 39, 3425–3433 (2010).
24. Ladet, S., David, L. & Domard, A. Multi-membrane hydrogels. Nature 452, 76–78 (2008).
25. Liang, G. L., Ren, H. J. & Rao, J. H. A biocompatible condensation reaction for controlled assembly of nanostructures in living cells. Nature Chem. 2, 54–60 (2010).
26. Sun, J. Y., Zhao, X. H., Mooney, D. J., Vlassak, J. J. & Suo, Z. G. Highly stretchable and tough hydrogels. Nature 489, 133–136 (2012).
27. John, G., Jadhav, S. R., Menon, V. M. & John, V. T. Flexible Optics: Recent Developments in Molecular Gels. Angew. Chem. Int. Ed. 8, 1760–1762 (2012).
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Author contributions
D.Z. and J.H.Y. synthesized semiconductor nanoparticles and analysed the hydrogel properties. S.B. assisted with the photocatalysis analysis. D.Z., J.H.Y., Q.S.W. and Q.G.W. designed the experiments, analysed the data and wrote the paper.

Additional information
Supplementary information accompanies this paper at http://www.nature.com/scientificreports

Competing financial interests: The authors declare no competing financial interests.

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