Bacteria responsive polyoxometalates nanocluster strategy to regulate biofilm microenvironments for enhanced synergetic antibiofilm activity and wound healing

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Abstract

Background: Nowadays, biofilms that are generated as a result of antibiotic abuse cause serious threats to global public health. Such films are the primary factor that contributes to the failure of antimicrobial treatment. This is due to the fact that the films prevent antibiotic infiltration, escape from innate immune attacks by phagocytes and consequently generate bacterial resistance. Therefore, exploiting novel antibacterial agents or strategies is extremely urgent.

Methods: Herein, we report a rational construction of a novel biofilm microenvironment (BME)-responsive antibacterial platform that is based on tungsten (W)-polyoxometalate clusters (POMs) to achieve efficient bactericidal effects.

Results: On one hand, the acidity and reducibility of a BME could lead to the self-assembly of POMs to produce large aggregates, which favor biofilm accumulation and enhance photothermal conversion under near-infrared (NIR) light irradiation. On the other hand, reduced POM aggregates with BME-induced photothermal-enhanced efficiency also exhibit surprisingly high peroxidase-like activity in the catalysis of bacterial endogenous hydrogen peroxide (H2O2) to produce abundant reactive oxygen species (ROS). This enhances biofilm elimination and favors antibacterial effects. Most importantly, reduced POMs exhibit the optimal peroxidase-like activity in an acidic BME.

Conclusion: Therefore, in addition to providing a prospective antibacterial agent, intelligent acid/reductive dual-responsive POMs will establish a new representative paradigm for the areas of healthcare with minimal side effects.

Key words: Biofilm, polyoxometalate, acid/reductive-responsive, peroxidase-like activity, photothermal antibacterial effect

Introduction

Bacterial infection diseases remain a major public health concerns to worldwide because of the high morbidity and mortality. [1-5]. In particular, the efficiency of bacteria that are disposed of via conventional antibiotics has been threatened by the developing emergence of antibiotic resistance. This is mainly because of structure changes and genetic mutations of planktonic bacteria and the formation of
biofilms [6, 7]. Biofilms are self-generated protective communities on an extracellular matrix that are composed of healthy bacterial cells. Biofilms irreversibly adhere to the surface of a sample or organ and can be embedded in an extracellular polymeric substance (EPS) matrix, which is mainly composed of extracellular bacterial DNA, exopolysaccharides, proteins and enzymes [7, 8]. As the major component of biofilms, an EPS protects pathogenic bacteria from attack by extraneous antibiotics. This is attributed to lowered biofilm penetration, which consequently induces bacterial resistance to antibacterial agents and causes antibiotic treatments to fail [9-13]. Additionally, because the encapsulation of EPS generates highly acidic niches (pH values close to 4.5), a biofilm microenvironment (BME) that is infected with bacteria is characterized as hypoxic and more acidic than the microenvironment of healthy tissues, and this leads to failure to eradicate infectious disease and to the recurrence of infection [7, 14, 15]. Thus, novel anti-biofilm strategies that target bacterial-infected microenvironments are important and have attracted significant interest. Specially, they have the potential to open new treatment paradigms for infections.

Recently, numerous strategies that take advantage of a unique bacterial-infected microenvironment as an important regulated target to facilitate the destruction of biofilm structure and to degrade the EPS have attracted widespread attention [16-18]. These strategies include chemical-mediated approaches using antibiotics, antibacterial peptides, quaternary ammonium compounds, and peptide-polysaccharides [19, 20], nonchemotherapeutic means such as nanotechnology-based strategies that involve the unique antibacterial nanostructures themselves and external physical stimuli such as light, electricity, magnetism and heat-activated alternative therapies [21-37]. Among these therapies, nanoparticle-based antibacterial agents have recently demonstrated their potential in the suppression of biofilm formation. This is achieved via reducing bacterial adhesion and viability, as well as regulating the unique bacterial-infected microenvironment as a method of inducing transformation. These new strategies can enhance antibacterial properties without causing side-effects and are able to overcome drug-resistance mechanisms with a high, sustained locally administrated dosage. Therefore, a promising strategy that is based on a combination of antibacterial agents is highly desirable for combating bacterial infectious disease. This is probably beneficial for regulating bacterial-infected microenvironments, overcoming biofilm recalcitrance, reducing antibiotic resistance and increasing anti-biofilm activity.

Among several antibacterial agents, hydrogen peroxide (H₂O₂), which is an endogenously active molecule, plays a critical role in physiological and pathophysiological processes such as wound disinfection. However, the concentration of H₂O₂ that is traditionally used is in the range of 166 mM to 1 M (volume ratio: 0.5–3 %) [38-40], and this prevents wound healing and damages adjacent normal organs during bacterial sterilization. Currently, various novel types of peroxidase-like nanomaterials that have high peroxidase-mimic activity, such as carbon-based nanomaterials [41-44], noble metals [45-51], iron-based nanomaterials [52-55], transitional metal dichalcogenides [56, 57], and polyoxometalates [58], have been known to have obvious antibacterial effects. They can disrupt the integrity of bacterial cell membranes and cell wall permeability. In particular, they restrain bacterial adhesion and biofilm formation. Unfortunately, these peroxidase-like nanomaterials that have high peroxidase-mimic activity still have disadvantageous, such as their intrinsic cytotoxicity, and because they generate oxidative stress at low concentrations of H₂O₂ and increase antimicrobial resistance under single antibacterial methods. In this regard, the development of new biocompatible and intelligent peroxidase-like nanostructures with BME stimuli-responsive character and the performance of the combination of multiple antibacterial substances are highly desirable for enhancing antibacterial efficiency and for achieving better antibacterial outcomes with potential therapeutic effect and decreased inevitable side-effects.

Herein, we report a representative acid/reductive-responsive inorganic paradigm as a result of tungsten (W)-based POM clusters (GdW₁₀O₃₆ NCs) for the regulation of BMEs to achieve efficient bactericidal effects and wound healing. Owing to the unique self-adaptive electronic structure and chemical versatility of POMs, such as polyanionic feature, rigid framework, monodispersed size, versatile properties, and biofunctionality, this representative W-based POM can self-assemble into larger aggregates in bacteria under the activation by the BME via protonation-triggered hydrogen bonding formation. This promotes effective adhesion to bacteria that have negatively charged cell surfaces within biofilms. Meanwhile, a reductive BME can “regenerate” the strong NIR absorption of POM through the chemical conversion of GSH-promoted W (VI) to W (V), which significantly increases their photothermal conversion performance. Most importantly, the obtained reduced POM aggregates in the BME have self-adaptive enhanced photothermal conversion, they catalyze endogenous hydrogen peroxide (H₂O₂) to generate...
abundant hydroxide radicals (•OH) via intrinsic peroxidase-like catalytic activity. This increases the obliteration of biofilm and cause bacteria to be more vulnerable to damage; they also can strengthen the antibacterial effect via the combination of NIR laser-induced hyperthermia with a low endogenous concentration of H2O2. Therefore, we believe that this work will advance the development of these acid/reductive-responsive POMs in the field of enzymatic catalysis and areas of healthcare.

**Methods**

**Materials**

All of the reagents and solvents were used as received without any modification. Sodium tungstate dehydrate (Na2WO4·2H2O), gadolinium chloride hexahydrate (GdCl3·6H2O), glutathione (GSH), and L-ascorbic acid (AA) were obtained from Alfa Aesar Reagent Company (China, Shanghai). Tetramethyl benzidine (TMB), o-phenylenediamine (OPD), 2,2’-azino-bis(3-ethylbenzothiazoline-6-sulfonic acid) diammonium salt (ABTS), 3, 3’-diaminobenzidine (DAB), horseradish peroxidase (HRP), propidium iodide (PI) and calcein (AM) were all obtained from Aladdin Company (China, Shanghai). P-Phthalic acid (TA) was obtained from the Chengdu area of the industrial development zone Xindu Mulan. Crystal violet staining solution was obtained from Scientific Phygene. All of the other chemical agents were used as received without any modification.

**Preparation of GdW10O36 nanoclusters before and after reduction**

Na9[Gd(W5O18)2]·×H2O nanoclusters (GdW10O36 NCs) were fabricated via a facile recrystallization method that has previously been described.[59, 60] Briefly, 25 mmol of Na2WO4·2H2O was dispersed in 20 mL of ultrapure water under continuous stirring at room temperature to form a homogeneous solution. A solution of acetic acid was then added dropwise until the pH was in the range of 7.40–7.50. Subsequently, 1.25 mmol/mL GdCl3·6H2O (477.95 g mol⁻¹, 2.0 mL) aqueous solution was added dropwise to this system under continuous stirring and heated to 85 °C until a transparent solution formed. Finally, crude crystals were isolated from solution when the mixtures were cooled to room temperature. To obtain the reduced GdW10O36 nanoclusters (reduced GdW10O36 NCs), another solution of glutathione with a different concentration or a solution of glycol was injected dropwise into the system, which was irradiated with a hand-held ultraviolet lamp. After irradiation for 12 h, water and ethanol were added three times to precipitate the reduced clusters, which were then dried in a freeze dryer to obtain reduced cluster powders.

**Peroxidase-like catalytic activity of reduced GdW10O36 nanoclusters**

TMB (1 mM) and OPD (1.85 mM) were chosen to evaluate the capacity of reduced GdW10O36 NCs (33 μg/mL) dispersed in tris buffer solution containing Na2HPO4·12H2O (0.2 M) and citric acid (0.1 M) to convert H2O2 (33 mM) into •OH radicals in the absence or presence of 808 nm NIR laser irradiation for 30 min. UV-Vis spectrophotometer (UV-Vis, UV-6100, MAPADA) was used to monitor the absorbance change of the characteristic peak at 652 nm or 450 nm. In addition, TA was used as a probe to react with the generated •OH to form TAOH, which is highly fluorescent. Fluorescence spectroscopy (HORIBA Dual-FL) was then used to record fluorescence spectra of samples after different treatments. Also the color change of peroxidase-like reduced GdW10O36 NCs was also photographed after 30 min of co-incubation at room temperature.

**Bacterial culture and dose-dependent antibacterial ability of reduced GdW10O36 nanoclusters**

E.coli (ATCC 25922) and S.aureus (ATCC 29213) bacteria were planted in a Luria–Bertani (LB) agar plate and then moved to 10 mL of LB with shaking for 8 h at 37 °C. When the logarithmic phase (A600 = 0.4) was achieved, Bacteria were collected via centrifugation at a speed of 8000 rpm for 1 min. The bacteria that collected at the bottom were washed and resuspended in phosphate buffered solution (PBS, 10 mM, pH = 7.4), which was attenuated to ~ 10⁶ colony-forming units (CFU·mL⁻¹) for subsequent experiments. To evaluate the dose-dependent antibacterial ability of reduced GdW10O36 NCs against E.coli and S.aureus bacteria, a series of reduced GdW10O36 NCs at different concentrations was mixed with 500 μL of the as-prepared bacteria solution for 30 min. Finally, bacterial strains with a concentration of 10⁴ CFU·mL⁻¹ were cultured in solid medium for 24 h using the spread plate method to count the number of the bacterial colonies.

**In vitro antibacterial effect of reduced GdW10O36 NCs with peroxidase-like catalytic activity and photothermal therapy**

To investigate the antibacterial property of peroxidase-like reduced GdW10O36 NCs with respect to E.coli and S.aureus bacteria, the following treatments were used on eight separate groups: (I) Control, (II) Reduced GdW10O36, (III) H2O2 (IV) Reduced GdW10O36 + H2O2, (V) Control + NIR, (VI)...
Reduced GdW10O36 + NIR, (VII) H2O2 + NIR, and (VIII) Reduced GdW10O36 + H2O2 + NIR. The final concentration of bacteria was 1.0 × 10^6 CFU mL\(^{-1}\), that of reduced GdW10O36 NCs was 200 μg mL\(^{-1}\), and that of H2O2 was 200 μM. After 10 min of co-incubation, groups V, VI, VII, and VIII were irradiated with an 808 nm NIR laser at a power density of 1.0 W cm\(^{-2}\) for 15 min. Then, an infrared thermal imager (E40, FLIR) was used to monitor the temperature. When co-incubation is conducted for 30 minutes, 100 μL of bacterial suspension in all of the groups was transferred to an agar culture plate for continuous cultivation for 12 h at 37 °C. All of the experiments were repeated in parallel three times.

**Fluorescent staining and SEM morphology observation of E. coli and S. aureus materials**

Propidium iodide (PI) (Aladdin Company, China, Shanghai) was chosen to evaluate the dead fluorescent staining of bacteria. In the assay, E. coli and S. aureus bacteria were used in different experimental groups and were incubated with fluorescing propidium iodide (30 μM) at 25 °C (avoiding light) for 15 min. The stained bacterial suspension (5 μL) was dropped onto a microslide that was then covered a coverslip, and an inverted fluorescence microscope (Carl-Zeiss) was used to visualize the sample. For further visual observations of the specific morphology of the bacteria after different treatments, E. coli and S. aureus bacteria were obtained via centrifugation at 8,000 rpm for 1 min and fixed with 4.0 % glutaraldehyde solution overnight at 4 °C. After three repeated rinsings with PBS, a graded series of ethanol solutions (30, 50, 70, 80, 90, and 100 %) was used to dehydrate bacterial cells for 10 min. Finally, the obtained specimens were set on a silicon glide, sprayed with gold, and imaged under an ultrahigh-resolution field emission SEM (S-4700, Hitachi, Japan).

**Bacterial biofilm formation and antibacterial effects of reduced GdW10O36 NCs on bacterial biofilm**

First, E. coli and S. aureus bacteria were washed three times with PBS (0.01 M) to discard the floating samples, and then 0.9 % saline solution was used to suspend the above-mentioned bacteria to an OD\(_{600}\) value up to 0.1. The diluted bacterial suspension (100 μL) was then co-incubated with 900 μL of LB or TSB medium for E. coli and S. aureus bacteria in a 96 well-plates. The samples were then continuously cultured and for 48 h at 37 °C. After that, fresh medium was substituted every 12 h. To evaluate the combined antibacterial effect of reduced GdW10O36 on bacterial biofilm, separate experiment groups were used: (I) PBS, (II) Reduced GdW10O36, (III) H2O2, (IV) Reduced GdW10O36 + H2O2, (V) PBS + NIR, (VI) Reduced GdW10O36 + NIR, (VII) H2O2 + NIR, and...
(VIII) Reduced GdW\textsubscript{10}O\textsubscript{36} + H\textsubscript{2}O\textsubscript{2} + NIR. After different disposes, bacterial biofilms were set aside in a shaker at 37 °C for 12 h. Then, 0.9 % physiological saline was used to rinse the biofilm three times to remove the samples and dead bacteria. Next, 500 μL of anhydrous alcohol was immediately used to fix the biofilm in each group, which was further cultured at 4 °C for 15 min. Thereafter, 300 μL of 1 % crystal violet staining solution was added to each well for 30 min of biofilm staining. Then, 0.01 M PBS was used to wash the biofilm in each well again, and the biofilms were then dried at room temperature. Last, each well was blended with anhydrous ethanol (500 μL) and co-incubated for 30 min. For the biofilm in each group, a multi-function microplate reader (Thermo Scientific, Varioskan LUK) was used to monitor the OD\textsubscript{590} value, and a digital camera was used to record the color.

**Morphology analysis of biofilms using confocal laser scanning microscopy (CLSM)**

For visually observations of biofilms using CLSM, biofilms were disposed with different methods. They were then cultured with Calcein-AM for 20 min and rinsed with 0.9 % NaCl to replace the discarded bacteriologic medium. An A1R-si Meta Inverted CLSM was then used to collect photographs with a size of 212.1 μm×212.1 μm and a framework size of 805×805. All of the samples were imaged with the same experimental facilities and analyzed using Zen 2011 software (Carl Zeiss). Each sample was independently measured in parallel three times.

**Animal study and wound healing process**

First, female BALB/c mice (6-8 weeks, 18-23 g) were acquired from Beijing Vital River Laboratory Animal Technology Co., Ltd. and separated into six groups: (I) no treatment, (II) H\textsubscript{2}O\textsubscript{2}, (III) reduced GdW\textsubscript{10}O\textsubscript{36} NCs, (IV) reduced GdW\textsubscript{10}O\textsubscript{36} NCs + NIR, (V) H\textsubscript{2}O\textsubscript{2} + reduced GdW\textsubscript{10}O\textsubscript{36} NCs, and (VI) H\textsubscript{2}O\textsubscript{2} + reduced GdW\textsubscript{10}O\textsubscript{36} NCs + NIR with three mice in each group. Then, an E.coli (ATCC 6538) bacterial suspension (1×10\textsuperscript{6} CFU mL\textsuperscript{-1}) was infected four times on each mouse’s back to produce subcutaneous abscesses with a wound area of ~78 mm\textsuperscript{2} after anesthesia. Twelve hours later, 5 μL of reduced GdW\textsubscript{10}O\textsubscript{36} NCs (100 μg/mL) and H\textsubscript{2}O\textsubscript{2} solutions (100 μM) were added dropwise onto the subcutaneous abscess at each infected site in each corresponding group. The mouse in groups IV and VI was then further irradiated under an 808 nm NIR laser at a power density of 1.0 W cm\textsuperscript{-2} for 15 min; meanwhile, a thermal infrared imager was used to monitor temperature changes. Toward the end of all of the treatments, the wound site of each mouse was photographed every other day, and the wound tissues were collected and fastened with 4% paraformaldehyde solution for histological analysis with Hematoxylin and eosin (H&E) staining, Masson’s trichrome staining and Giemsa staining.

**Results and Discussions**

To demonstrate the feasibility of our assumption, we first fabricated W-based polyoxometalate clusters (GdW\textsubscript{10}O\textsubscript{36} NCs) according to a previously described method (Figure 1A) [59, 60]. The GdW\textsubscript{10}O\textsubscript{36} NCs were composed of cations (Na\textsuperscript{+} and NH\textsubscript{4}\textsuperscript{+}) and Keggin-type macroanionic units. The successful fabrication was confirmed by Fourier transform infrared (FT-IR) spectroscopy and Raman spectroscopy (Figure S1). GdW\textsubscript{10}O\textsubscript{36} NCs had a monodispersed spherical morphology and an ultrasmall diameter of about 1–3 nm that exhibited high hydrophilicity and dispersity at a pH of 7.4 (Figure 1B). Interestingly, as observed from transmission electron microscopy (TEM) (Figure 1C, D), these clusters exhibited distinct acidity-induced self-assembly with pH values that ranged from 7.4 to 4.0 when these clusters were acidified. They were homogeneously dispersed and had a diameter of ~30 nm in a slightly acidic environment (pH 6.4), and they further aggregated into a larger structured morphology when the pH was further decreased to 4.0. In particular, the acidity-activated self-assembly characteristic of GdW\textsubscript{10}O\textsubscript{36} NCs was further confirmed by dynamic light scattering measurements (Figure 1E). The acidity-induced self-assembly of GdW\textsubscript{10}O\textsubscript{36} NCs was ascribed to the protonation of POM macroanions through pH-induced hydrogen bond formation, which was beneficial for regulating the BME to achieve an enhanced antibacterial effect. Subsequently, to further investigate the redox-activated properties of the synthesized GdW\textsubscript{10}O\textsubscript{36} NCs, glutathione (GSH) and ethylene glycol (EtOH), which are reducing agents, were used as patterns for incubating with GdW\textsubscript{10}O\textsubscript{36} NCs at different concentrations. As seen in Figure 1F, G and S2, the pure solution of GdW\textsubscript{10}O\textsubscript{36} NCs was colorless and had no absorption in the near infrared (NIR) region. However, the color of the solution gradually changed to blue after reduction with GSH and EtOH, This color change was simultaneously accompanied by a corresponding strong characteristic absorption in the NIR region, and this gradually became stronger with an increase in the GSH, EtOH, or GdW\textsubscript{10}O\textsubscript{36} NCs concentration. Consistent with this, the relative residual amount of AA and GSH gradually decreased with GdW\textsubscript{10}O\textsubscript{36} NCs oxidation, as shown in Figure S3, and this, further confirms the superior redox-activated properties of GdW\textsubscript{10}O\textsubscript{36} NCs. Beyond that,
to obtain fundamental insight into the redox-activated behavior of these POM clusters, X-ray photoelectron spectroscopy (XPS) was recorded. The XPS results reveal the oxidation state of W in GdW_{10}O_{36} NCs before and after the reduction process. As seen in Figure 1H-J, GdW_{10}O_{36} NCs were reduced by GSH to form reduced GdW_{10}O_{36} NCs, and this confirms that the reduced GdW_{10}O_{36} NCs were successfully synthesized. This phenomenon is the results of multiple reversible steps of electron exchange between delocalized electron density and occupiable cation site of W (V), which are present because of the increased reduction of GdW_{10}O_{36} NCs [61, 62]. This enhances the electron relaxation polarization and strengthens NIR absorption. Therefore, all of these results indicate that the GdW_{10}O_{36} NCs have admirable acidic/reductive dual-responsive capability that enables the efficient regulation of the acidic BME and promotes oxidation of endogenous GSH, which play a critical role in the bacterial antioxidant defense system.

An aqueous solution of the reduced GdW_{10}O_{36} NCs had a navy blue color and strong absorption in the NIR region. Thus, the photothermal effects of samples were carefully investigated. The results indicate that the photothermal effect of reduced GdW_{10}O_{36} NCs depended on laser power intensity and GSH concentration. With an increase in the NIR laser power density, the temperature of this photothermal material increased (Figure 2A). Furthermore, after 10 min of irradiation under an NIR laser, GdW_{10}O_{36} NCs that were mixed with different concentrations of GSH had a high temperature (Figure 2B-D).

Figure 1. Synthesis and characterization of oxidized and reduced GdW_{10}O_{36} NCs. (A) Schematic diagram of synthesis. (B-D) TEM images of nanoclusters at pH = 7.4, 6.4 and 4.0. (E) DLS measurements of GdW_{10}O_{36} NCs with continuous acidification from pH = 7.4 to 4.0. UV-vis spectra and corresponding photographs of GdW_{10}O_{36} clusters before and after reduction in the presence of various concentrations of GSH (0, 10, 15, and 20 mM) on the 2nd (F) and 5th day (G). (H) XPS spectra of GdW_{10}O_{36} NCs. (I-J) XPS spectra of W 4f in GdW_{10}O_{36} NCs before and after reduction.
concentrations of GSH showed a higher increase in temperature. Moreover, GdW\textsubscript{10}O\textsubscript{36} NCs that were co-incubated with 150 mM GSH at a pH of 7.4 even presented a remarkable temperature change of 36 °C (Figure 2B). Interestingly, we found that the photothermal effect of reduced GdW\textsubscript{10}O\textsubscript{36} NCs was also pH-dependent. This corresponds to low pH-induced self-assembly of GdW\textsubscript{10}O\textsubscript{36} NCs and proportionally increases NIR absorbance (Figure 2C, D). The results showed that a more efficient increase in temperature can be monitored under BME-mimicked conditions under irradiation of an 808-nm laser, which facilitate to be a photothermal agent for photothermal antibacterial application. All of these results indicate that the obtained GdW\textsubscript{10}O\textsubscript{36} NCs have the potential to be used for pH/GSH dual-responsive photothermal therapy.

Furthermore, one of the unique features of our GdW\textsubscript{10}O\textsubscript{36} NCs system is that, except for intelligent acidic/reductive dual-responsive enhanced photothermal efficacy under BME-mimicked conditions; the GdW\textsubscript{10}O\textsubscript{36} NCs after reduction could act simultaneously as a robust and effective intrinsic peroxidase mimic for catalyzing bacterial endogenous H\textsubscript{2}O\textsubscript{2} to achieve enhanced biofilm elimination and favorable antibacterial effects. In the following, we employed terephthalic acid (TA) and 3,3,5,5-tetramethylbenzidine (TMB) as peroxidase substrates for evaluating the peroxidase-like activity of reduced GdW\textsubscript{10}O\textsubscript{36} NCs or the commonly used enzyme HRP.

The results indicate that reduced GdW\textsubscript{10}O\textsubscript{36} NCs and HRP can both efficiently catalyze •OH radical generation to form a strong fluorescence intensity with TA or a characteristic absorption peak at 652 nm with the TMB substrate (Figure 3); with 33 mM of H\textsubscript{2}O\textsubscript{2} The system was blue (Figure S4). Simultaneously, like other peroxidase mimics, the catalytic activity of reduced GdW\textsubscript{10}O\textsubscript{36} NCs, was dependent on pH, temperature, H\textsubscript{2}O\textsubscript{2} and concentrations; these characteristics are similar to those of HRP. Notably, the high activity of reduced GdW\textsubscript{10}O\textsubscript{36} NCs and HRP was optimal at a pH of 5.5, which is approximately the acidic BEM (Figure 3A). Hence, the enzyme mimics that have high catalytic activity in the important pH range of 3.5-5.5 (i.e., the pH of an acidic BME) will be promising candidates for use in biological and antibacterial research. Additionally, we discovered that compared to HRP, reduced GdW\textsubscript{10}O\textsubscript{36} NCs can withstand higher temperature and higher H\textsubscript{2}O\textsubscript{2} concentration to achieve maximum values of peroxidase activity. Nevertheless, the peroxidase-like activity was suppressed if the temperature were further elevated or if the H\textsubscript{2}O\textsubscript{2} concentration were further increased, as indicated by the enzymic catalytic reaction (Figure 3B-D). Thus, all of the above results demonstrate that reduced GdW\textsubscript{10}O\textsubscript{36} NCs can serve as peroxidase mimics that have high activities and stability for enhancing biofilm removal and for facilitating antibacterial effects.

Figure 2. Photothermal effects of reduced GdW\textsubscript{10}O\textsubscript{36} NCs in vitro. (A) Temperature evaluation of reduced GdW\textsubscript{10}O\textsubscript{36} NCs (5.3 ppm Gd) as a function of irradiation time (0-10 min) using an 808 nm laser at different power densities (0.2, 0.4, 0.6, 0.8, and 1.0 W/cm\textsuperscript{2}). (B) Temperature evaluation of reduced GdW\textsubscript{10}O\textsubscript{36} NCs that were incubated with different concentrations of GSH (50, 100, and 150 mM) as a functional of irradiation time (0-10 min) using an 808 nm laser at a power density of 1.0 W/cm\textsuperscript{2}. (C) Acidity-dependent photothermal effects (C) and the corresponding IR images (D) of reduced GdW\textsubscript{10}O\textsubscript{36} NCs (5.3 ppm Gd) under the irradiation of an 808 nm NIR laser at a power density of 1.0 W cm\textsuperscript{-2} for 10 min.
To further understand the fundamental catalytic mechanism of reduced GdW_{10}O_{36} NCs, we tested whether the reduced GdW_{10}O_{36} NCs can promote the decomposition of H_{2}O_{2} into hydroxyl radicals (•OH) under the irradiation of an 808 nm laser to improve antibacterial activities for synergistic wound therapy. Thus, we used 3,3,5,5-tetramethylbenzidine (TMB), OPD and TA as substrates to record the production of hydroxyl radicals (•OH) that originate from the effective photothermal conversion of the reduced GdW_{10}O_{36} NCs to promote the low concentration of H_{2}O_{2} species decomposition. As expected, the fluorescence intensity is significantly enhanced in the presence of reduced GdW_{10}O_{36} NCs compared the fluorescence intensity of the pure H_{2}O_{2} species. This indicates that reduced GdW_{10}O_{36} NCs powerful catalysts for the production of •OH from H_{2}O_{2} because of their optimal peroxidase-like catalytic activity. More interestingly, when reduced GdW_{10}O_{36} NCs were treated successively with H_{2}O_{2} for 10 min and irradiation from an 808 nm laser for 30 min, an apparent increase in fluorescence intensity was monitored in this system. In contrast, only a relative weak fluorescence signal was obtained in the absence of either H_{2}O_{2} or irradiation from an 808 nm laser (Figure 4A-C). This phenomenon was also shown in the enzyme-catalyzed reactions (Figure 4D). A clearer explanation of the aforementioned results is described below for the brief enhancement mechanism of reduced GdW_{10}O_{36} NCs that are irradiated with an 808 nm laser to produce •OH radicals (Figure 4E). Therefore, these results effectively confirm that reduced GdW_{10}O_{36} NCs significantly enhance the formation of •OH species via the combination of their photothermal effect and peroxidase-like catalytic activity.

To demonstrate the antibacterial effect of the reduced GdW_{10}O_{36} NCs-based biocatalytic process under pathological conditions, a peroxidase-mediated antibacterial assessment against E. coli or and S. aureus bacteria was carried out using the standard colony counting method to explore the feasibility of reduced GdW_{10}O_{36} NCs as peroxidase mimics and photothermal agents to enhance the generation of reactive oxygen species. After incubation with large concentrations of reduced GdW_{10}O_{36} NCs for 24 h, no obvious reduced bacteria population and cells were observed in E. coli and S. aureus bacteria as well as 4T1.
cells, even at a dose as high as 1 mg/mL (Figures S5 and S6). This indicates that reduced GdW_{10}O_{36} NCs with superior biocompatibility and low cytotoxicity can meet the requirements for biomedical applications. Next, we explored the antibacterial effects of these NCs as simultaneous peroxidase and photothermal agents. As seen in Figure 5A, B, significantly reduced numbers of colonies were observed after bacteria were incubated with reduced GdW_{10}O_{36} NCs in the presence of H_{2}O_{2} for 10 min and then irradiated with an 808 nm NIR laser for 15 min. Specifically, the bacteria proliferation inhibition percentages reached 80% for *E. coli* bacteria. In contrast, in the absence of either H_{2}O_{2} or irradiation under an 808 nm laser, the reduced GdW_{10}O_{36} NCs group had relatively inferior antibacterial behavior. Meanwhile, because of the low concentration of H_{2}O_{2} (200 μM) and reduced GdW_{10}O_{36} concentration (200 μg/mL), the single H_{2}O_{2} group and reduced GdW_{10}O_{36} NCs group both exhibited negligible antibacterial effects towards *E. coli*. The survival percentages were 85.4% and 64.1%, respectively, which indicates that peroxidase catalysis alone cannot achieve ideal antibacterial effects. Correspondingly, similar experimental results were also confirmed in *S. aureus* bacteria (Figure 5C, D). Consequently, all of the results further substantiate that reduced GdW_{10}O_{36} NCs have a superior capacity to be function as peroxidase-like enzymes and photothermal agents with an effective combined antibacterial application.

Figure 4. Enzyme-like properties of reduced GdW_{10}O_{36} NCs. (A, B) Absorbance spectra and visual color changes of TMB (1 mM) and OPD (1.85 mM) in different reaction systems: (1) H_{2}O+TMB, (2) 808 nm+TMB, (3) Reduced GdW_{10}O_{36} NCs+TMB, (4) Reduced GdW_{10}O_{36} NCs+808 nm+TMB, (5) H_{2}O_{2}+TMB, (6) Reduced GdW_{10}O_{36} NCs+H_{2}O_{2}+TMB, and (7) Reduced GdW_{10}O_{36} NCs+808 nm+H_{2}O_{2}+TMB. (C) Fluorescence spectra of terephthalic acid for hydroxyl radical detection at the maximum emission wavelength of 435 nm. (D) Different color reactions and the mechanism scheme of reduced GdW_{10}O_{36} NCs catalysis in the presence of H_{2}O_{2} and various peroxidase substrates represented by AH. (E) Scheme of the catalytic mechanism of reduced GdW_{10}O_{36} NCs combined with irradiation from an 808 nm laser in the presence of H_{2}O_{2}.

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Figure 5. Photographs and antibacterial activity of reduced GdW$_{10}$O$_{36}$ NCs after different treatments. (A, C) Photographs and (B, D) survival rates of treated E. coli and S. aureus after exposed to (I) Control, (II) Reduced GdW$_{10}$O$_{36}$, (III) H$_2$O$_2$, (IV) Reduced GdW$_{10}$O$_{36}$+ H$_2$O$_2$, (V) Control+NIR, (VI) Reduced GdW$_{10}$O$_{36}$+NIR, (VII) H$_2$O$_2$+NIR, and (VIII) GdW$_{10}$O$_{36}$ + H$_2$O$_2$+NIR. Conditions: 200 µg/mL reduced GdW$_{10}$O$_{36}$, 200 µM H$_2$O$_2$, 1 W/cm$^2$ 808 nm laser and 15 min. P values were based on the Student’s test: *P < 0.05, and **P < 0.01.

Additionally, the biofilm, which is a hybrid of bacterial cells, irreversibly adheres onto the surface of a material or tissue. It is packed in self-generated extracellular polymeric substances (EPS), which act as a barrier in preventing antibiotic permeation into the bacterial cells; consequently, bacterial drug-resistance is produced and antibiotic treatment failure occurs. Therefore, anti-biofilm activity is an important characteristic of antibacterial agents for achieving desirable antibacterial effects. Herein, the crystal violet staining method was used to investigate biofilm formation of reduced GdW$_{10}$O$_{36}$ NCs after different treatments. As seen in Figure 6A, C, the navy blue color remained in the reduced GdW$_{10}$O$_{36}$ NCs, H$_2$O$_2$ and 808-nm laser alone treated groups. A slight decrease in the blue color was observed for the groups that were treated with the combination of reduced GdW$_{10}$O$_{36}$ NCs plus laser or the combination of reduced GdW$_{10}$O$_{36}$ NCs and H$_2$O$_2$. Nevertheless, the blue color significantly faded to colorless after the combined treatment of reduced GdW$_{10}$O$_{36}$ NCs with laser irradiation or with H$_2$O$_2$ treatment. Accordingly, biofilm biomass of E. coli bacteria that were treated with different methods was also quantified using UV-vis spectroscopy (Figure 6B, D). The results indicate that much less biofilm biomass are measured for disposed with reduced GdW$_{10}$O$_{36}$ NCs and laser irradiation in the presence of H$_2$O$_2$ in comparison to the monotherapy groups or the control group.

Furthermore, 3D confocal microscopy was used to analyze the average thickness of biofilms that formed on glasses via different treatments. As seen in Figure 6E&S7, much thinner biofilms formed on reduced GdW$_{10}$O$_{36}$ NCs and H$_2$O$_2$ as well as for the samples that were irradiated with a laser. In contrast, much thicker biofilms were observed on the surfaces of samples in the monotherapy groups or control group. These observations confirm that peroxidase-like catalysis and the photothermal effect of reduced GdW$_{10}$O$_{36}$ NCs more efficiently reduce the thickness of biofilm. This phenomenon is ascribed to the unique electrical structure and acidity of reduced GdW$_{10}$O$_{36}$ NCs for regulating the BME and achieving antibacterial effects. We also used scanning electron microscopy to observe material adsorption on bacterial membranes and to investigate changes in the morphology of treated E. coli and S. aureus. As seen in Figure 7A, C, the negative group exhibited smooth, rod-like and sphere-like shapes, and the surface was intact. In contrast, membranes of bacteria distorted and shrunk to some extent after the sample was irradiated with an 808 nm laser or exposed to a low concentration of H$_2$O$_2$ alone. Moreover, we found that bacteria membranes were seriously destroyed and crashed after the combined photothermal and H$_2$O$_2$ treatment, and this was attributed to the strong production of •OH that was induced by reduced GdW$_{10}$O$_{36}$ NCs and H$_2$O$_2$ under 808 nm laser
exposure. In addition, we used live/dead staining imaging to demonstrate the relative survival of bacteria after different treatments (Figure 7B, D). The results were similar to the previously mentioned bacterial cloning and scanning electron microscopy results, and this further confirms that reduced GdW10O36 NCs have an admirable capacity for efficiently inhibiting bacterial proliferation because of their photothermal and peroxidase-like catalysis.

Figure 6. Bacterial biofilm elimination effect and images of reduced GdW10O36 NCs after different treatments. (A, C) Bacterial biofilm eradication effect of E. coli and S.aureus bacteria treated with reduced GdW10O36 NCs after after treated with (I) PBS (control), (II) Reduced GdW10, (III) H2O2, (IV) Reduced GdW10 + H2O2 in the presence or absence of NIR. (B, D) OD590 value for evaluating changes in biofilm biomass of E. coli and S.aureus bacteria after different treatments. (E) 3D confocal laser scanning microscopy (CLSM) of biofilms formed from E. coli and S.aureus bacteria treated with reduced GdW10O36 NCs after different treatments. Image sizes: 212.1×212.1 µm. The biofilms were stained with CA. P values were based on the Student’s test: *P < 0.05, and **P < 0.01.
To further investigate the potential that reduced GdW_{10}O_{36} NCs with antibacterial activity have for promoting wound healing in vivo, E. coli bacteria-infected wound-disinfected mice were first designed and then divided into six groups: (I) PBS, (II) H_{2}O_{2}, (III) Reduced GdW_{10}, (IV) Reduced GdW_{10} + NIR, (V) Reduced GdW_{10} + H_{2}O_{2}, and (VI) Reduced GdW_{10} + NIR + H_{2}O_{2}. The concentration of reduced GdW_{10}O_{36} NCs was fixed at 100 μg/mL, and that of H_{2}O_{2} was fixed at 100 μM; these are much less than the concentrations that are commonly used (0.5-3 % and about 166 mM to 1 M) [38-40]. During therapeutic periods, the states of wounds were photographed on days of 4 and 10. As seen in Figure 8A, B, the wounds in the group without treatment and in the group that received reduced GdW_{10}O_{36} NCs or H_{2}O_{2} alone exhibited serious inflammatory reactions with ulceration, erythema, and edema on day 4. Also, the relative wound area was only reduced 81.54±3.09%, 84.48±3.69% and 75.94±3.63%, respectively. In contrast, the group that treated with reduced GdW_{10}O_{36} NCs in the presence of irradiation from an 808 nm laser or with H_{2}O_{2} alone slightly decreased over time, which is contrary to the untreated group. More importantly, the skin of the mice was almost completely recovered on day 10 after treatment with reduced GdW_{10}O_{36} NCs + NIR + H_{2}O_{2} (Figure 8C). Meanwhile, regeneration and differentiation of fibroblasts under the dermis as well as regeneration of the skin around the wound are important processes in wound healing. Therefore, structure changes to the epidermis and formation of collagen fiber during wound healing were further investigated. Hematoxylin and eosin (H&E) and Masson and Giemsa staining of the epidermis of repaired tissues after different treatments were performed on day 10. As seen in Figure 8D, H&E staining indicates that new fibroblasts and newborn blood vessels filled with plenty of red blood cells had begun to move to the wound position from adjacent normal tissue or organ for the reduced GdW_{10}O_{36} + H_{2}O_{2} + NIR group by the end of the experiment. In
contrast, abundant dead cell fragments and a loose disordered incomplete dermal layer are still observed in other groups. Furthermore, Masson’s trichrome staining showed infected necrotic foci, edema, ulceration, and some inflammatory cells were also seen in the control groups. Also, the skin epidermis structure and collagen fiber of the monotherapy group (reduced GdW10O36 + H2O2 and reduced GdW10O36 + NIR) are relatively clear, complete, and regenerated. In contrast, intact established collagen fibers and dermal layer were observed in the reduced GdW10O36 + H2O2 + NIR group, and this implies that the wounds have completely healed. Additionally, Giemsa staining was used to test the amounts of adherent bacteria after 10 days of treatments. The results indicate that the amount of bacteria in the reduced GdW10O36 + H2O2 + NIR group decreased significantly in comparison to the control and monotherapy groups. This confirms the positive effects of reduced GdW10O36 during wound treatment. In brief, this antibacterial platform efficiently inhibits bacterial proliferation, and also promotes differentiation of fibroblasts and regeneration of skin. Therefore, all of these results demonstrated that reduced GdW10O36 NCs have great potential for accelerating wound healing in the E. coli-infected abscesses because of their superior photothermal efficiency and surprisingly high peroxidase-like activity.

Figure 8. Wound-healing studies of reduced GdW10O36 NCs after different treatments in vivo. (A) Relative wound area and (B) weight of mice infected with E.coli after different treatments. (C) Photographs and histologic analyses (D) of E. coli-infected wound treated with (I) PBS (control), (II) H2O2, (III) Reduced GdW10, (IV) Reduced GdW10+NIR, (V) Reduced GdW10+H2O2, and (VI) Reduced GdW10+NIR+H2O2 at days 4 and 10 and their corresponding histologic analyses. P values were based on the Student’s test: *P < 0.05, and **P < 0.01.
Conclusions

In brief, we successfully developed an antimicrobial platform using the intelligent tungsten (W)-polyoxometalate cluster paradigm. This system features pH/GSH dual-response via acidic BME-triggered self-assembly and reductive BME-driven strengthened NIR absorption. Thus, it results in the rapid elimination of biofilm and exhibits remarkable antibacterial activity upon exposure to NIR laser radiation. Additionally, the intrinsic peroxidase-mimic activity of reduced POMs that are generate in a BME can catalyze the decomposition of endogenous H₂O₂ with low concentrations to generate OH. More importantly, in the presence of bacterial endogenous H₂O₂, an enhanced antibacterial effect can be achieved via the strengthened photothermal effect and intrinsic peroxidase-mimic activity of reduced POMs. Therefore, the intelligent pH/GSH dual-responsive tungsten (W)-polyoxometalate cluster resolves the problem of severe biofilm infections and also achieves an unprecedented antibacterial efficacy. In brief, the particular acidic/reductive dual-responsive BME is anticipated to be a powerful tool for illustrating significant guidance on the future design of BME-specific antibacterial nanoplatforms with efficient antibacterial effects.

Supplementary Material

Supplementary figures.
http://www.thno.org/v10p1003I1s1.pdf

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Competing Interests

The authors have declared that no competing interest exists.

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