INORGANIC CHEMISTRY







FRONTIERS

RESEARCH ARTICLE

View Article Online
View Journal | View Issue



Cite this: *Inorg. Chem. Front.*, 2024, **11**, 4439

GSH/pH dual-activated POM@MOF for tumor cell-specific synergistic photothermal and chemodynamic therapy†

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Photothermal therapy (PTT) is emerging as a promising therapeutic approach for tumor treatment. However, the low therapeutic effect of photothermal agents due to their non-specificity and the monotherapy approach remains challenging for further applications. Herein, tumor microenvironment (TME) specifically responsive POM@MOF composites with sequential activation properties were constructed by encapsulating POMs (PMo₁₂, short for H_3 PMo₁₂O₄₀) into metal organic framework (MIL-101) carriers for synergistic PTT/chemodynamic therapy (CDT) treatment. PMo₁₂@MIL-101 releases PMo₁₂ and Fe³⁺ under acidic TME conditions, which are subsequently converted into reduced PMo₁₂ and Fe²⁺ by GSH in the TME. PMo₁₂ in its reduced state exhibits excellent photothermal conversion efficiency (47.03%) under near-infrared laser irradiation, which boosts the CDT effects by accelerating the reaction between H_2O_2 and Fe^{2+} and producing more *OH radicals. The resulting PTT/CDT synergistic therapeutic effects are highly dependent on the pH value and GSH concentration in the TME, and exhibit a strong inhibition of HepG2 cell proliferation compared to monotherapy.

Received 30th April 2024, Accepted 11th June 2024 DOI: 10.1039/d4qi01079e rsc.li/frontiers-inorganic

Introduction

Photothermal therapy (PTT) is an emerging cancer treatment that uses a photothermal agent (PTA) to convert near-infrared (NIR) light into heat for tumor ablation. Due to the minimal invasiveness and precise spatial specificity, PTT has been recognized as an effective strategy for cancer treatment and has attracted extensive attention in recent years.^{2,3} Many PTAs, including inorganic nanomaterials (e.g., black phosphorus nanosheets and Au nanorods), 4,5 organic molecules (e.g., ICG), 6-9 and polymeric materials (e.g., polydopamine), 10,11 have been developed for cancer treatments. However, the practical application of PTAs in cancer treatment remains challenging due to two main reasons: (a) most PTAs are non-specific and randomly distributed throughout the body, requiring manual intervention to achieve targeted therapy, which demands a high level of manipulation; (b) PTT alone is insufficient in eradicating tumors due to the inhomogeneous heat

Polyoxometalates (POMs) are a class of nanoscale inorganic polymetallic oxygen clusters that have a wide range of applications in catalysis, assembly, and biomedicine. 14-17 Recently, POMs have shown great potential in biomedical research fields due to their high biocompatibility, versatile structures, adjustable compositions, low cost and easy availability. 18,19 Particularly, the reduction of Mo-based POMs to molybdenum blue produces strong absorption in the NIR region, giving Mo-based POMs great potential as PTAs. 20-22 In addition, POMs have exhibited acid-activated self-assembly properties that enhance the retention by self-assembling into aggregates once they enter the acidic tumor microenvironment (TME). 23-25 These characteristics endow POMs with a tumor cell-specific activation property as PTAs. Nevertheless, the highly negatively charged surface of POMs makes it difficult to cross cell membranes, 26-28 and thus carriers are often used to enhance the cellular uptake of POMs.

Metal organic frameworks (MOFs) are considered as a promising class of drug carriers due to their well-defined structure, tunable pore size, pH-responsiveness, and biodegradability.^{29,30} A variety of POM@MOF materials have been reported in recent years, and the corresponding preparation approaches are well-established.^{17,31} POMs residing in

distribution throughout the tumor tissue, resulting in inevitable tumor recurrence and metastasis. Therefore, it is crucial to develop tumor cell-specific PTAs and to integrate them with other therapeutic strategies.

Polyoxometalates (POMs) are a class of panoscale inorganic

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† Electronic supplementary information (ESI) available. See DOI: https://doi.org/10.1039/d4qi01079e

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the MOF cavity avoid the ineffective encapsulation caused by the surface loading, and maintain a high stability of the POM structure.32 Notably, MOFs have adjustable inorganic metal centers, which endow them with intrinsic Fenton catalytic activity as a chemodynamic therapy (CDT) agent by tuning the metal center such as Fe and Cu. 33,34 As such, the encapsulation of POMs into MOFs will not only enhance cellular uptake efficacy by improving the surface charges of POMs, but also endow them with tumor cell-specifically activated PTT/CDT combined therapeutic effects.

In this work, we report a TME-responsive POM@MOF (PMo₁₂@MIL-101, denoted as P@M) therapeutic agent with sequential activation properties for synergistic PTT/CDT therapy (Fig. 1). By confining PMo₁₂ with activated photothermal conversion properties to MIL-101, the low cellular uptake efficacy of PMo₁₂ that is caused by its highly negative surface charge can be avoided. The MIL-101 structure collapses in the acidic and reducing TME, favoring the activated release of PMo₁₂ in tumor cells. The released PMo₁₂ is reduced and aggregated under endogenous GSH and acidic conditions, resulting in electron transfer and a subsequent photothermal effect that triggers thermal ablation of tumor cells under NIR laser irradiation (808 nm). Concurrently, the released Fe³⁺ is reduced to Fe2+ by GSH, which further causes massive generation of 'OH from overexpressed H2O2 in the TME via the Fenton reaction. In addition, local hyperthermia promotes the Fenton reaction and enhances the effect of CDT, and thus the synergistic effect of combined PTT/CDT therapy shows excellent efficacy in suppressing tumor cell proliferation.

Results and discussion

As shown in Fig. 2A, P@M was synthesized via the one-pot solvothermal method.³⁵ Scanning electron microscopy (SEM) images showed that P@M resembled MIL-101 in the monodisperse octahedral morphology, with a size of approximately 800 nm (Fig. 2B and S1†), suggesting that the morphology of MIL-101 remained unchanged upon encapsulation of PMo₁₂. No obvious small particles were observed on the surface of the octahedral structure, which indicated that PMo₁₂ was orderly assembled with MIL-101 rather than being self-assembled and then composited with MIL-101. Elemental mapping results displayed that the Mo, Fe, and C elements were uniformly distributed over the entire P@M (Fig. 2C). In addition, the X-ray powder diffraction (XRD) pattern (Fig. S2†) of P@M closely resembled that of MIL-101,36 providing additional evidence that the framework of MIL-101 was not affected by PMo₁₂. Note that no obvious XRD patterns attributable to PMo₁₂ were detected in P@M, which further demonstrated that PMo12 clusters were well dispersed without aggregation. The content of PMo₁₂ in P@M was calculated to be 26.8% from the ICP results (Table S1†). Moreover, as shown in Fig. 2D, PMo₁₂ alone exhibited a negative zeta potential because of the highly

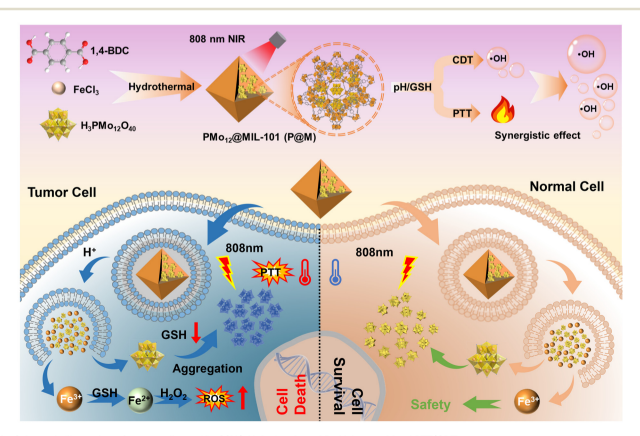


Fig. 1 Schematic illustration of tumor cell-specific GSH/pH dual activated P@M for combined PTT/CDT treatment.



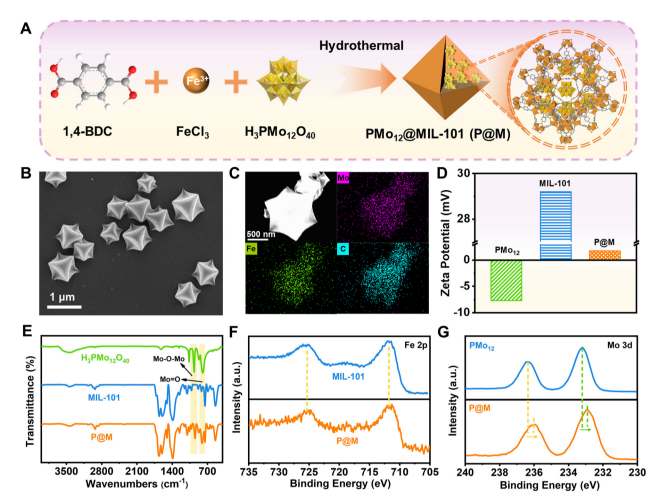


Fig. 2 Preparation and characterization of P@M. (A) Schematic illustration of the synthetic route for P@M. (B) SEM image of P@M. (C) Elemental mappings of Mo, Fe, and C signals. (D) Zeta potential of PMo₁₂, MIL-101 and P@M. (E) FT-IR spectra of PMo₁₂, MIL-101 and P@M. High-resolution XPS spectra of (F) Fe 2p and (G) Mo 3d for PMo₁₂ and P@M.

negatively charged surface of the clusters. In contrast, MIL-101 showed a positive zeta potential.³⁷ The as-prepared P@M exhibited a slightly positive zeta potential, suggesting that PMo₁₂ was confined within the MIL-101 cages.

The Fourier transform infrared (FT-IR) spectrum of P@M showed the characteristic bands of both PMo₁₂ and MIL-101 (Fig. 2E). The bands at about 542, 749, 1390 and 1598 cm⁻¹ were attributed to $\nu(\text{Fe-O})$, $\nu(\text{C-H})$, $\nu_s(\text{COO-})$ and $\nu_{as}(\text{COO-})$ of MIL-101, respectively.³⁸ The four bands at ~1064, 967, 870 and 794 cm $^{-1}$ corresponded to the $\nu(\text{P-O})$ of the PO $_4$ tetrahedron and $\nu(\text{Mo=O})$, $\nu(\text{Mo-O}_{b}\text{-Mo})$ and $\nu(\text{Mo-O}_{e}\text{-Mo})$ of the PMo₁₂ cluster, respectively.^{39,40} This result indicated that the structure of PMo₁₂ was maintained during the encapsulation process. Additionally, the composition and elemental valence of P@M were investigated by X-ray photoelectron spectroscopy (XPS) measurements. For both MIL-101 and P@M, the highresolution spectrum of Fe 2p exhibited two broad peaks at 711.74 and 725.35 eV, corresponding to Fe^{3+} $2p_{3/2}$ and Fe^{3+} 2p_{1/2}, respectively (Fig. 2F).⁴¹ The Mo 3d peaks for PMo₁₂ located at ~233.2 and 236.3 eV were assigned to Mo⁶⁺ 3d_{5/2}

and Mo⁶⁺ 3d_{3/2}, respectively (Fig. 2G).⁴² Compared with the individual PMo₁₂, the two Mo 3d peaks of P@M shifted slightly toward lower binding energies at ~232.9 and 236.0 eV, which was consistent with those reported in the literature.³⁵ The results indicated a charge transfer between the MOF and the encapsulated POM, which was attributed to the host-guest interaction.

P@M exhibited a slightly positive zeta potential in phosphate buffer solution (PBS) at pH 7.4 since PMo₁₂ was encapsulated in MIL-101 cages (Fig. 3A and B). Intriguingly, the surface potential of P@M became negative in PBS at pH = 5.5, indicating the collapse of the MIL-101 structure and the release of negatively charged POM clusters under acidic conditions. The pH-sensitive property of P@M will make it responsive to the acidic tumor microenvironment. Similar to the individual PMo₁₂ clusters (Fig. S3†), P@M showed a strong absorbance in the near-infrared-I (NIR-I) window upon reduction by glutathione (GSH) (Fig. 3C). The absorbance intensity of P@M in its reduced form (denoted as rP@M) increased with an increase in the concentration of GSH. The XPS results (Fig. 3D)

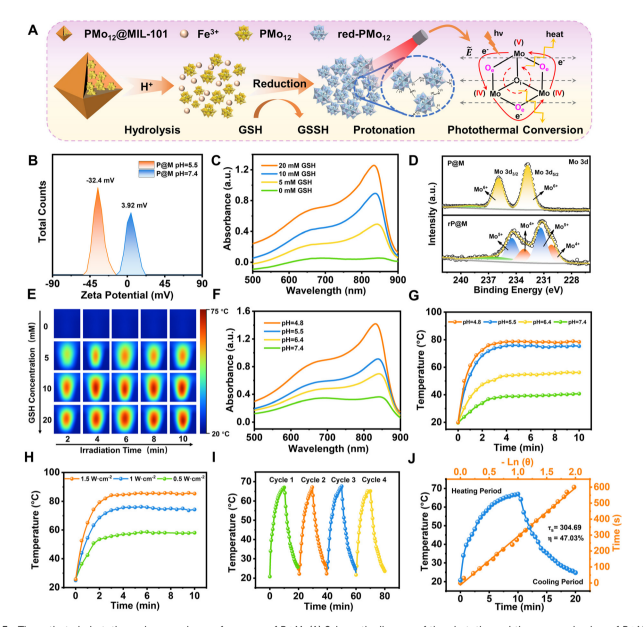


Fig. 3 The activated photothermal conversion performance of P@M. (A) Schematic diagram of the photothermal therapy mechanism of P@M. (B) Zeta potential of P@M in PBS buffer at different pH values. (C) UV-vis spectra of 1 mg mL⁻¹ P@M at different GSH concentrations in pH 5.5 buffer. (D) XPS profiles of Mo 3d for P@M and rP@M. (E) Visualized IR thermograms of 1 mg mL⁻¹ P@M solutions at different GSH concentrations (0-20 mM) irradiated with an 808 nm laser (1.0 W cm⁻²) for 10 min. (F) UV-vis spectra of 1 mg mL⁻¹ P@M at different pH values in the presence of 10 mM GSH. (G) Temperature changes of 1 mg mL⁻¹ P@M solution at different pH values during 10 min of irradiation at 808 nm (1.0 W cm⁻²) when the GSH concentration is 10 mM. (H) Temperature profiles for 1 mg mL $^{-1}$ P@M under 808 nm laser irradiation at different power densities when the pH value is 5.5 and GSH concentration is 10 mM. (I) Thermal stability of 1 mg mL $^{-1}$ P@M under 808 nm laser irradiation in the presence of 10 mM GSH. (J) Heating and cooling curves of P@M and fitted linear data obtained from the cooling process.

indicated that GSH reduced the Mo in the POM from the oxidation state of Mo(vi) to mixed valence states of Mo(v) and Mo (iv). As illustrated in Fig. 3A, the inter-valence charge transfer between Mo(v) and Mo(v) via the oxo-bridge endowed the reduced P@M with NIR photothermal conversion behaviour. 20,43 As shown in Fig. S4,† a prominent photothermal conversion phenomenon was observed for P@M upon treatment with GSH in PBS (pH = 5.5). The photothermal behavior was found to correlate with the reducibility of Mo over a

certain range of GSH concentrations. Visualized IR thermograms showed that the highest solution temperature of P@M could reach about 76 °C when 10 mM GSH was applied to reduce Mo (Fig. 3E and S4†).

In addition, like other Mo-based POMs that are reported in the literature, the individual PMo₁₂ was reduced and selfassembled into larger-sized aggregates with a concomitant increase in the number of Mo at mixed-valence states under acidic conditions, which promoted reversible multistep electron exchange and enhanced the electronic relaxation polarization under an applied electric field (laser), leading to an increased NIR absorbance (Fig. S5†).44,45 As such, for P@M, the acidic solution conditions can not only destruct the MIL-101 framework and release the encapsulated PMo₁₂, but also promote the self-assembly of reduced PMo₁₂ and improve the NIR photothermal conversion. As shown in Fig. 3F, the absorption of P@M in the NIR-I window was significantly enhanced with decreasing pH in the presence of GSH (10 mM), confirming the low-pH-induced aggregation property of PMo₁₂. Correspondingly, the temperature of the P@M system with a concentration of 1000 µg mL⁻¹ significantly increased with decreasing pH under laser irradiation at 808 nm (1 W cm⁻²), and the temperature could be elevated to more than 70 °C within 2 min at pH 5.5 (Fig. 3G and S6†). It should be noted that P@M showed no obvious temperature change under 808 nm laser irradiation in the absence of GSH under neutral pH conditions, reflecting that P@M had a pH/ GSH dual-activated photothermal conversion Moreover, the photothermal behavior of P@M was irradiation power and concentration dependent (Fig. 3H and S7-S9†). P@M maintained good stability in the 4-cycle laser on/off test and its photothermal conversion efficiency was calculated to be 47.03% (Fig. 3I and J), which was comparable to the performance of recently reported inorganic photothermal materials (Fig. S10†). These experimental results demonstrated that P@M has tumor microenvironment-activated photothermal conversion properties, and is a high-performance photothermal material that can be used for photothermal therapy.

Intriguingly, in addition to PMo₁₂, the destruction of P@M also led to the release of free Fe ions. As illustrated in Fig. 4A, the released Fe ions in their reduced state (Fe²⁺) were speculated to catalyse a Fenton-like reaction, thus functioning as a chemodynamic therapy (CDT) agent. To verify the potential of P@M as a CDT agent, the change in the Fe valence state after treatment with GSH was firstly investigated by XPS. As shown in Fig. 4B, after GSH treatment, the broad Fe³⁺ peaks in P@M, originally located at 712.4 and 725.9 eV, were significantly shifted to lower binding energies in red-P@M, appearing at 709.3 (Fe²⁺ $2p_{3/2}$) and 722.5 eV (Fe²⁺ $2p_{1/2}$), respectively, which indicated that the released Fe³⁺ could be reduced to Fe²⁺ by GSH. In addition, the reduction of Fe3+ in P@M was assessed by a chromogenic reaction using o-phenanthroline, which reacted with Fe2+ to form a complex with an absorbance centered around 510 nm. As shown in Fig. 4C, in a slightly acidic (pH = 5.5) environment, the absorbance of the Fe-o-phenanthroline complex gradually increased with increasing GSH concentration. In contrast, no obvious Fe2+ was detected without GSH, revealing that GSH induced the reduction of Fe^{3+} to Fe^{2+} . Besides, an obvious increase in the absorbance of the Fe-ophenanthroline complex was observed with a decline in the pH of the P@M solution (Fig. S11†), which further indicated the pH-dependent Fe release of P@M.

The generated Fe2+ can react with H2O2 and produce ROS through a Fenton reaction (Fig. 4A), 46,47 which will oxidize 3,3',5,5'-tetramethylbenzidine (TMB) and lead to a color reaction from colorless to blue. As shown in Fig. 4D and S12,† the absorbance of oxTMB at 650 nm was enhanced with an

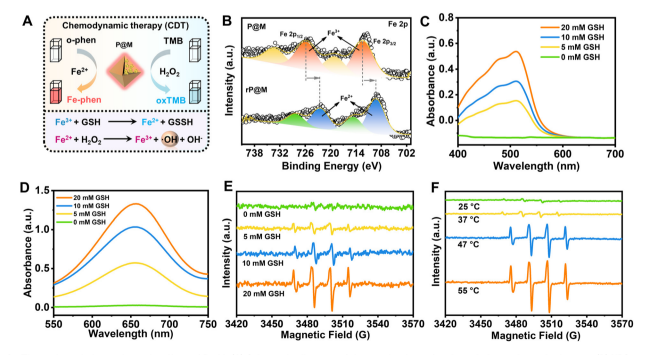


Fig. 4 The activated chemodynamic effect of P@M. (A) Schematic diagram of the mechanism of activated chemodynamic therapy. (B) XPS profiles of Fe 2p for P@M and rP@M. (C) UV-vis spectra of P@M (in PBS, pH 5.5) with o-phenanthroline at different GSH concentrations. (D) UV-vis spectra of P@M (in PBS, pH 5.5) with TMB in the presence of 100 μM H₂O₂ at different GSH concentrations. (E) ESR spectra of DMPO/ OH for P@M at different GSH concentrations. (F) ESR spectra of DMPO/OH for P@M at different temperatures.

increase in GSH concentration or decline in the pH value. The GSH concentration/pH-dependent nature of TMB-related color reaction was consistent with the above-mentioned chromogenic reaction from the complexation of Fe2+ and o-phenanthroline, which further confirmed the CDT potential of P@M via a Fenton reaction. To identify the generated ROS species, an electron spin resonance (ESR) measurement was performed using 5,5-dimethyl-1-pyrroline N-oxide (DMPO) as a trapping agent. As shown in Fig. 4E, no signal was detected for P@M without GSH. In contrast, the typical 1:2:2:1 quartet pattern was observed for P@M that reacted with H₂O₂ in the presence of GSH, indicating the generation of the 'OH species.

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A strengthened ERP signal of 'OH was detected with the increase in the concentration of GSH, which proved that the enhanced amount of Fe2+ would promote the generation of 'OH through a Fenton reaction. The above-mentioned results suggested that P@M had obvious tumor microenvironmentactivated Fenton activity and could function as an excellent CDT agent. Interestingly, the production of 'OH showed a temperature-dependent behavior. As shown in Fig. 4F, there was a significant increase in 'OH production when the temperature rose from 25 to 55 °C, indicating that high temperatures would promote Fenton-like reaction. Notably, the ROS production at body temperature (37 °C) was much lower than that when heated, which ensured the safety of P@M for normal cells or tissues. The results suggested that P@M might have a great synergistic effect by exerting combined PTT and CDT functions.

Encouraged by the prominent photothermal effect and efficient Fenton reaction activity of P@M, its potential therapeutic effect on cancer cells was further investigated. Fig. 5A shows that after a 24 h-incubation with P@M, distinct dark

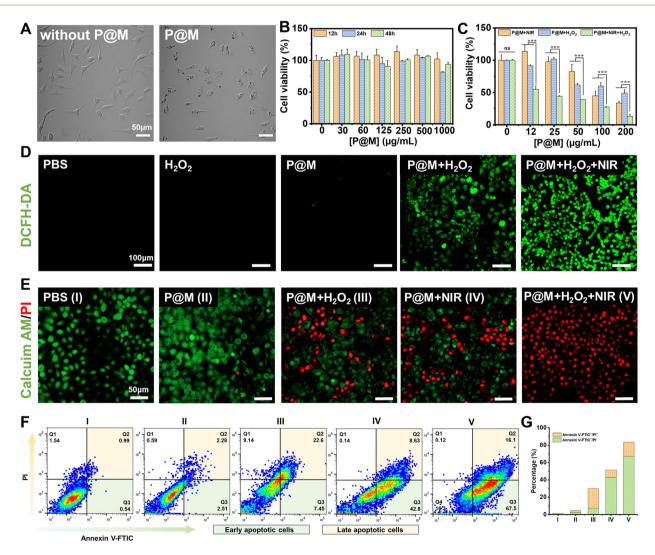


Fig. 5 (A) Cellular uptake of P@M in HepG2 cells. (B) HepG2 cell viabilities after the treatment with different concentrations of P@M for 12, 24 and 48 h. (C) Cytotoxicity assays of HepG2 cells with PTT, CDT and PTT/CDT treatments. Statistical analysis was performed via two-way ANOVA with Bonferroni's multiple comparisons test. ***P < 0.001; data are presented as means \pm SD (n = 3). (D) CLSM images of intracellular ROS levels with different treatments. (E) Calcein AM/PI double staining of HepG2 cells with different treatments. (F) Flow cytometry analysis of the apoptosis of HepG2 cells subjected to different treatments and (G) corresponding quantitative analysis of cell apoptosis percentages.

granules were observed in the cells compared to untreated HepG2 cells, indicating the cellular uptake of P@M. As shown in Fig. 5B, no significant decline in cell viability was observed after treating HepG2 cells with $0-500~\mu g~mL^{-1}$ P@M for 12, 24, and 48 h. Notably, cell viability remained above 90% even after 48 h of co-culturing with high concentrations of P@M up to 1000 μg mL⁻¹, which indicated that P@M had negligible cytotoxicity against the HepG2 cell line. H2O2 alone has no obvious cytotoxicity to HepG2 cells (Fig. S13†).

To assess the therapeutic efficacy of P@M as a PTT and CDT agent, HepG2 cells were exposed to 808 nm laser irradiation (1 W cm⁻², 5 min, NIR), H₂O₂ and NIR/H₂O₂ combination in the presence of P@M (Fig. 5C). Cell survival in all the three groups exhibited a predominant dose-dependent trend with P@M. At a concentration of 200 µg mL⁻¹, the cell viability was 34.3% for the P@M + NIR group, 49.8% for the $P@M + H_2O_2$ group, and 14.8% for the $P@M + NIR + H_2O_2$ group, respectively. These results demonstrated the potential of P@M for PTT and CDT, as well as its synergistic effect when used in a PTT/CDT-combination therapy.

To investigate P@M-induced intracellular ROS generation, HepG2 cells were stained with 2',7'-dichlorodihydrofluorescein diacetate (DCFH-DA), a commercially available fluorescent probe, under various treatment conditions (Fig. 5D). Compared to the control group treated with PBS, neither the H₂O₂ nor the P@M treated group showed an increase in ROS levels. In contrast, the P@M + H2O2 group showed a remarkable green fluorescence, indicative of an intracellular Fenton reaction with ROS generation. Intriguingly, a significant elevation in ROS levels was observed in the P@M + NIR + H₂O₂ group compared to that of the P@M + H₂O₂ group, which demonstrated that hyperthermia may boost the CDT process.

Calcein-AM/PI double staining was further performed for the direct visualization of cell viability (Fig. 5E). Negligible cell death was observed in both the PBS control and P@M (200 µg mL^{-1}) treatment groups. Notably, CDT (P@M + H₂O₂) and PTT (P@M + NIR) treatments resulted in a substantial increase in

cell death, especially in the PTT and CDT co-treated group (P@M + H₂O₂ + NIR), where nearly all cells in the irradiated region were killed. An annexin V-FITC and PI double staining assay was carried out to investigate the ability of P@M to induce cell apoptosis (Fig. 5F). The PTT + CDT combination treatment group (P@M + H₂O₂ + NIR) exhibited the highest efficiency in inducing cell apoptosis, with an apoptotic ratio (sum of Q_2 and Q_3) of 83.6%, evidently higher than the ratios observed in the P@M (4.79%), P@M + H₂O₂ (30.05%) and P@M + NIR (51.43%) groups under identical conditions (Fig. 5G). These results provided compelling evidence that the combination therapy of P@M with PTT and CDT activities was effective in triggering apoptosis compared to therapy with PTT and CDT alone, suggesting a synergistic role for P@M in cancer therapy.

Finally, the potential in vivo toxicity of P@M was evaluated. Normal 8-week-old male Balb/c mice were used in the in vivo experiments. All animal experiments comply with the Animal Ethical and Welfare Committee of Beijing Institute of Technology. The hematological and histological data were collected and analyzed after intravenous injection with P@M (dose: 20 mg kg^{-1}) into the tails of healthy mice. As shown in Fig. S14,† in routine blood tests, there was no statistically significant difference between the P@M administration group and the control group in the measurement results of white blood cells (WBC), red blood cells (RBC), hemoglobin (HGB), hematocrit (HCT), mean corpuscular volume (MCV), mean corpuscular hemoglobin (MCH) and mean platelet volume (MPV). Moreover, renal function indicators such as blood urea nitrogen (BUN), uric acid (UA) and creatinine (CRE) were comparable to those of the control group. Meanwhile, the corresponding histological changes in the major organs of mice, including the heart, liver, spleen, lungs, and kidneys, harvested after administration were analyzed by typical H&E staining (Fig. 6). The results showed no obvious signs of damage to the normal anatomical structure of the organs. These findings demonstrated the high biocompatibility and biosafety of P@M.

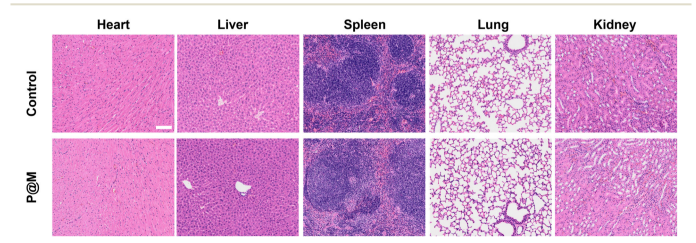


Fig. 6 Histological examinations of major organs of mice after intravenous injection with P@M (scale bar: 100 μm).

Conclusions

In conclusion, GSH/pH dual-activated P@M that specifically responds to the TME was successfully developed for synergistic PTT/CDT therapy. Fe³⁺ and PMo₁₂ clusters were released from P@M when exposed to relatively acidic conditions, and were subsequently reduced by GSH in the TME. The generated Fe²⁺ showed Fenton-like activity to catalyze the formation of highly toxic 'OH from H2O2, while the reduced PMo12 clusters displayed an excellent photothermal conversion efficiency as high as 47.03% under NIR irradiation. Both ESR and intracellular experimental results demonstrated a photothermal-enhanced CDT effect due to the increasing production of ROS at elevated temperature or upon laser treatment. The inhibitory effect of P@M on HepG2 cells further confirmed the efficacy of combined PTT/CDT treatment. These findings suggest potential for the design of tumor cell specifically activated PTT/CDT combined therapy based on POM-nanoplatforms.

Data availability

The data supporting this article have been included as part of the ESI. \dagger

Author contributions

The manuscript was written through contributions of all authors. All authors have given approval to the final version of the manuscript.

Conflicts of interest

The authors declare no conflict of interest.

Acknowledgements

This research was supported by the Beijing Natural Science Foundation (2242015 and L233031), National Natural Science Foundation of China (22007004, 22178019, 22208013, 22288102, and 22207010) and the Fundamental Research Funds for the Central Universities (buctrc202010, XK1802-6, XK1803-05, and XK1902). All the experimental animals were humanely treated in accordance with the guidelines for animal care, and were approved by the Experimental Animal Welfare Ethics Committee of Beijing Institute of Technology.

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