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Copper depletion-induced tumor cuproptosis[†]

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Copper homeostasis is crucial for cells, especially for rapidly proliferating cancerous cells. Copper imbalance-induced cell death (*i.e.*, cuproptosis) has emerged as a new strategy for tumor therapy. While copper accumulation-induced cuproptosis has been extensively investigated and its underlying mechanism recently elaborated, copper depletion-induced cuproptosis remains largely unexplored. Herein, we demonstrated copper depletion-induced tumor cuproptosis through the development of a smart copper-depleting nanodrug (*i.e.*, ZnS nanoparticles), leveraging a cation exchange reaction between ZnS and copper ions. This cation exchange reaction is driven by the large difference in solubility product constants (K_{sp}) between ZnS and CuS. Our ZnS nanoparticles demonstrated a potent copper-depleting ability, which induced tumor cuproptosis both *in vitro* and *in vivo*. We proposed a copper-depleting mechanism primarily linked to the dysfunction of cellular copper-contained enzymes, contrasting with the mechanism of copper accumulation-induced cuproptosis. Furthermore, by modifying the ZnS nanoparticle with a polydopamine shell and a glucose transporter 1 DNAzyme (GD), we developed a multifunctional copper nanoconsumer with strong tumor growth and metastatic inhibition activity, enhancing copper depletion-promoted tumor therapy.

Introduction

Copper (Cu), an essential trace element for all organisms, participates in many biological processes, including mitochondrial respiration, redox homeostasis, and kinase signaling.^{1–3} Notably, intracellular copper levels must be maintained in a strict range. A slight deviation from its physiological concentration can disrupt a series of cellular processes, causing diseases such as Menkes syndrome (due to copper deficiency), Wilson's disease (due to copper accumulation), and even cancer.^{4–7} Earlier studies reported that tumor cells have a higher copper content to maintain rapid proliferation,^{8,9} but this

content must also be controlled within an optimal range. Therefore, modulating the copper level in cancer cells has emerged as a promising strategy for tumor therapy *via* either copper overaccumulation or depletion.^{10–12}

To date, an enormous amount of effort has been devoted to copper accumulation-induced cuproptosis.^{13–18} Notably, copper ionophores such as elesclomol and disulfuram have been used for exogenous copper delivery. Recently, the mechanism of copper accumulation-induced cuproptosis has been elucidated, characterized by the aggregation of lipoylated proteins and decrease of Fe–S cluster proteins, which then led to cell death.¹⁹ However, copper accumulation inevitably requires supplementary copper ions, which has limited the wide adoption of copper accumulation-induced cuproptosis.

In contrast, copper depletion-mediated cuproptosis provides a promising strategy to tackle the aforementioned challenge by sequestering cellular bioavailable copper ions with copper chelators.¹² Among these copper chelators, several have advanced to the preclinical/clinical phase for cancer therapy, including trientine, ATN-224, and tetrathiomolybdate (TTM).^{20–22} Encouragingly, TTM has already reached clinical phase II for breast cancer and has shown inhibition of tumor growth.^{12,23} However, large dosages of these molecular chelators are needed to effectively sequester cellular copper, which not only cause side effects but also ultimately induce drug resistance. While nanoparticulate and polymeric systems have been developed for copper depletion, the majority still rely on molecular chelators.^{24–26} Therefore, designing new types of

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copper depletion independent of molecular chelators is greatly needed.

Inspired by the solubility product (K_{sp}) concept, we envisioned that mesoporous zinc sulfide (ZnS) nanospheres could be used for copper depletion through a cation exchange reaction between ZnS and copper ions, leveraging the large difference in K_{sp} of ZnS and CuS (K_{sp} of ZnS is 1.6×10^{-24} and CuS is 6.3×10^{-36} , respectively) (Table S1†).^{27–38} Herein, we synthesized ZnS nanoparticles for copper depletion-induced cupropotosis, which is distinct from previously reported chelators. Additionally, we proposed the copper depletion mechanism. Through further functionalization of ZnS, both primary and metastatic tumors were significantly inhibited based on copper depletion.

Results and discussion

Synthesis and copper depleting ability of ZnS

Based on an organic–inorganic interaction mode, mesoporous ZnS nanospheres were synthesized by a ligand-assisted metal coordination strategy.^{39–41} Typically, using zinc nitrate as a metal source, thioacetamide (TAA) as a sulfur source, citric acid (CA) and hexamethylene tetramine (HMTA) as ligands, and cetyltrimethylammonium bromide (CTAB) as a template, these agents reacted together to produce ZnS nanospheres (Fig. 1a). The morphology of ZnS was determined through transmission electron microscopy (TEM), showing a mesoporous-spherical structure with a size of about 97 ± 4 nm (Fig. 1b). The hydrodynamic size and dispersity of ZnS were measured by dynamic light scattering (DLS), showing a size of about 164 nm and good dispersity (polydispersity index, PDI = 0.107) (Fig. 1c).

Given the disparity in K_{sp} values between ZnS and CuS, we envisioned that ZnS could exchange with Cu^{2+} and produce CuS (Fig. S1†). To test this, we explored the cation-exchanging ability of ZnS by adding Cu^{2+} (CuCl_2 as the donor of Cu^{2+}) to the ZnS solution. Gradually increasing amounts of Cu^{2+} were dropwise added into ZnS solution (20 μg per mL ZnS). An obvious color change of the reaction solution from ivory-white (ZnS, inset photo of Fig. 1c) to yellowish green (Cu-exchanged ZnS, inset photo of Fig. 1e) was observed. The products were harvested and analyzed for copper and zinc contents using inductively coupled plasma-optical emission spectrometry (ICP-OES). The results revealed that the copper stack proportion (compared with Zn) increased in a Cu^{2+} concentration-dependent manner and reached 98% when 30 μM Cu^{2+} was added (final concentration, in 20 μg per mL ZnS solution) (Fig. S2†), thus verifying the copper-exchanging ability of ZnS. When higher concentrations of Cu^{2+} were supplemented, a plateau of copper proportion was observed. This was probably due to the saturation point beyond which Zn could not be exchanged with the remaining Cu^{2+} . Then, we characterized the Cu-exchanged ZnS product (final concentration, 30 μM Cu^{2+} exchanging with 20 μg per mL ZnS). It exhibited similar morphology and size to ZnS in the TEM image (Fig. 1d) and good dispersity as determined by DLS (Fig. 1e). These results indicated the occurrence of cation exchange.

Subsequently, X-ray photoelectron spectroscopy (XPS) was performed to determine the composition of ZnS and Cu-exchanged ZnS. The Cu 2p peak emerged while the intensity of the Zn 2p peak decreased (using the peak of S 2p as an internal reference) in the Cu-exchanged ZnS (Fig. 1f). The powder X-ray diffraction (XRD) analysis indicated the crystalline phase of Cu-exchanged ZnS with peaks consistent with those of CuS (PDF card no.: 01-078-0876) (Fig. 1g), confirming the formation of CuS through a cation exchange reaction. High-resolution TEM (HRTEM) and energy-dispersive X-ray spectroscopy (EDS) elemental mapping revealed a homogeneous distribution of Cu and S elements in the produced CuS, with a smaller portion of Zn content. This result was consistent with the earlier observation that cation exchange has a saturation point that prevents further Cu^{2+} exchange (Fig. S2†). Semiquantitative EDS analysis was also performed, showing the weight content of Cu (4.99%) and Zn (1.33%) in the produced CuS, compared with 5.62% Zn in ZnS (Fig. S3†). These findings were mutually consistent, confirming the copper-exchanging ability of ZnS and the formation of CuS. Therefore, we hypothesized that ZnS could be used as a powerful copper-depleting agent for copper depletion therapy.

Fabrication of a “copper nanoconsumer”

To facilitate targeted copper depletion, a “copper nanoconsumer” was fabricated by functionalizing the surface of ZnS (Fig. 2a). A thin shell of polydopamine (PDA), a bio-inspired acid-degradable material widely applied for surface coating,^{42,43} was polymerized on ZnS for core protection, obtaining ZnS@PDA (termed ZP). It has been reported that cellular copper depletion could induce mitochondrial dysfunction, shifting cancer cell metabolism more to glycolysis.^{7,8,44} Therefore, we designed a Zn^{2+} dependent glucose transporter 1 (Glut 1) mRNA-cleaving DNAzyme (*i.e.*, GD) for selective glycolytic inhibition.^{45–47} Finally, GD was conjugated on ZP *via* a metal ion assisted DNA adsorption strategy,^{48,49} ZnS@PDA/GD (termed ZPD) as the “copper nanoconsumer” was obtained.

The monodispersed core–shell structure of ZP was demonstrated by TEM imaging (Fig. 2b), showing a thickness of the PDA shell about 3 nm. The successful conjugation of GD was confirmed by fluorescence quenching analysis and gel electrophoresis (Fig. S4 and S5†). The GD loading efficiency in ZPD was shown to reach 79.34% by fluorescence measurement (Fig. 2c). ZPD was characterized by TEM imaging (Fig. 2d), size distribution (Fig. S6†), and elemental mapping analysis (Fig. S7†), showing a similar morphology to ZP with a small change in the hydrodynamic diameter and zeta potential (Fig. 2e). These results demonstrated the successful synthesis of ZPD.

Subsequently, the properties of ZPD were explored. First, the degradation of PDA was monitored by treating ZPD (Cy3-labeled GD) in different pH buffers (20 mM phosphate buffer, pH = 7.4, 6.8, and 5.8). When ZPD was treated with pH 5.8 buffer (termed as treated-ZPD), the highest GD release efficiency was achieved, as indicated by fluorescence recovery (Fig. 2f). Degradation of the PDA shell was also observed by TEM imaging (Fig. S8†). Both results verified PDA degradation under acidic conditions,



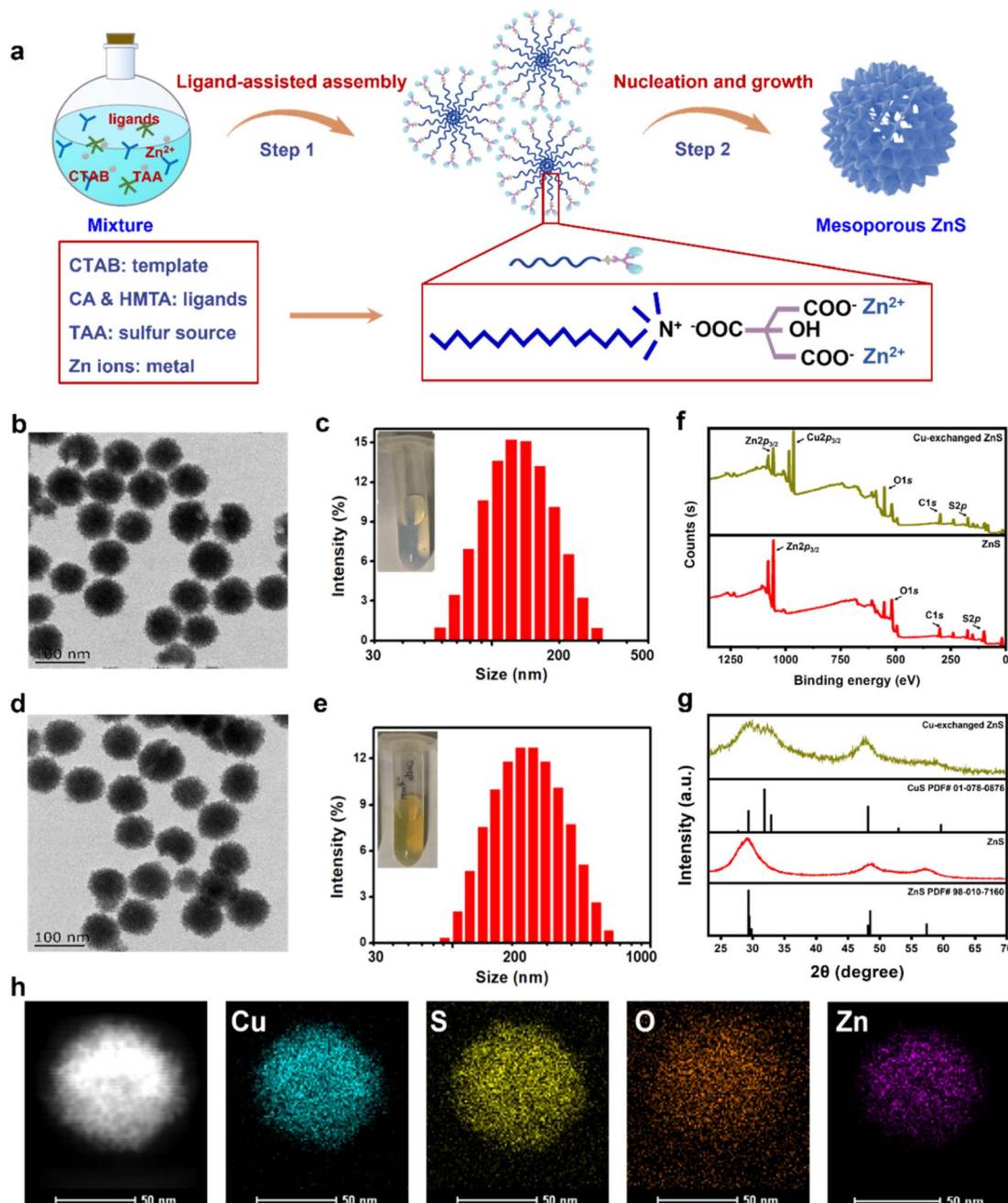


Fig. 1 Characterization and copper depleting ability of ZnS. (a) Schematic of mesoporous ZnS nanosphere synthesis. (b) TEM image of ZnS. (c) Size distribution of ZnS. Inset: digital photo of ZnS solution. (d) and (e) TEM image and size distribution of Cu-exchanged ZnS by TEM and DLS. Inset: digital photo of ZnS after Cu depletion. (f) XPS analysis of ZnS and Cu-exchanged ZnS. (g) XRD patterns of ZnS and Cu-exchanged ZnS. (h) Element distribution mapping of ZnS after Cu depletion by HRTEM.

similar to the acidic microenvironment in tumors. Second, after PDA degradation, the “naked” ZnS could mediate copper depletion and CuS could be produced *in situ*. With the photothermal effect of CuS, an apparent temperature increase was noticed under laser irradiation (808 nm laser, 2 W cm⁻² at different time points) in the treated-ZPD group by adding Cu²⁺, while the ZPD group showed a weak temperature rise due to the protection of the PDA shell (Fig. 2g and h). These results demonstrated that after PDA degradation, ZnS-mediated cation exchange could occur in the presence of Cu²⁺ and produce

photothermal active CuS. Third, the released Zn²⁺ during the copper-cation exchange reaction could specifically activate the cleaving ability of GD DNAzyme (Fig. 2i). As shown in Fig. 2j, both GD + substrate with the treated-ZPD + Cu²⁺ group and GD + substrate with the Zn²⁺ group could successfully cleave the substrate, whereas no cleaved substrate was noticed in the only GD + substrate group. Our findings demonstrated the occurrence of ZnS-mediated copper depletion and GD activation by the released Zn²⁺ ions.

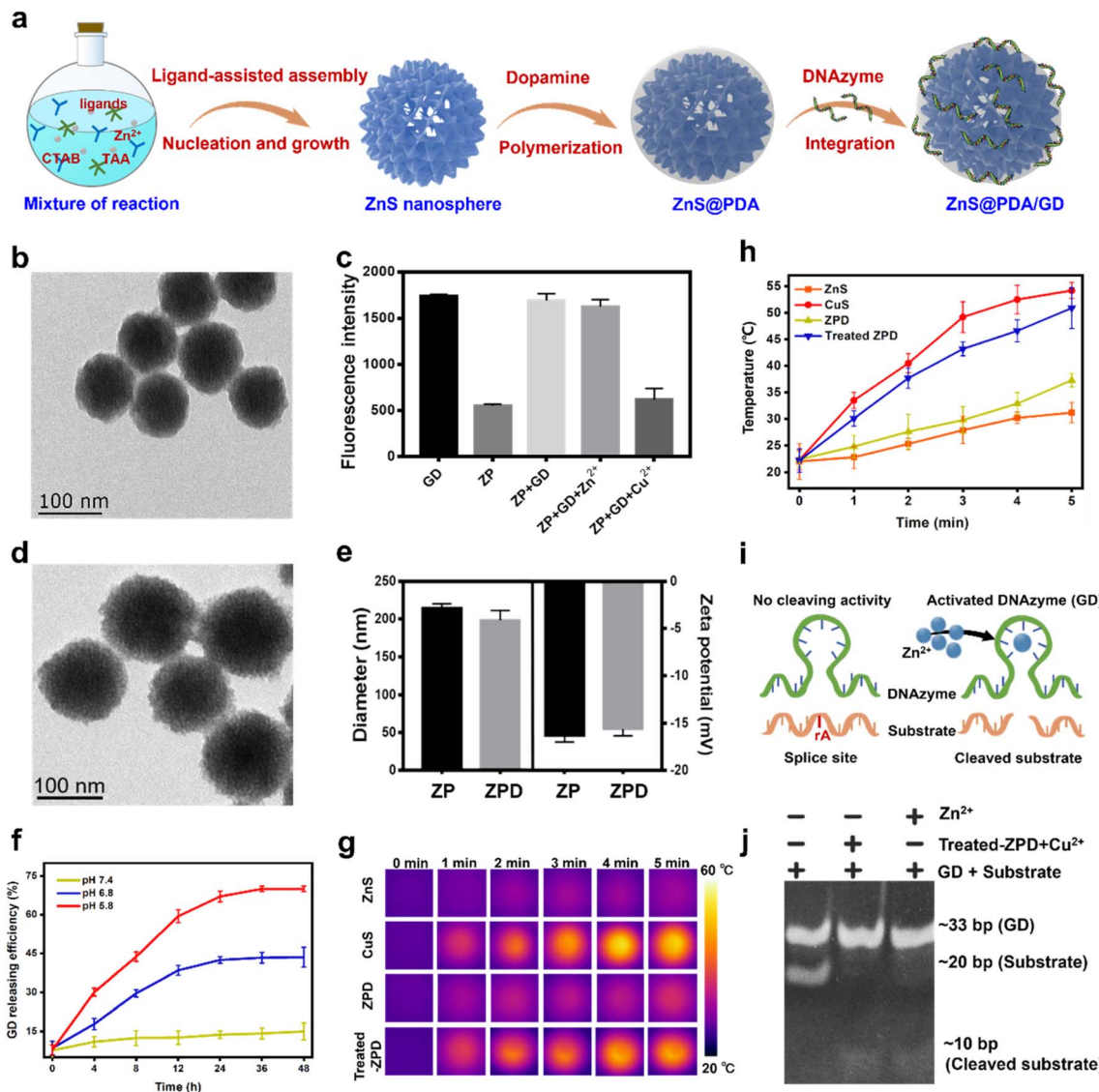


Fig. 2 Synthesis and characterization of “copper nanoconsumer” ZPD. (a) Scheme of “copper nanoconsumer” synthesis. (b) TEM image of ZP. (c) Metal ion-mediated adsorption between ZP and GD, probed by fluorescence quenching analysis. (d) TEM image of ZPD. (e) Comparison of diameters and zeta potentials between ZP and ZPD by dynamic light scattering. (f) GD release profile at different pH values. (g) Photothermal effect analysis by monitoring the temperature change under laser irradiation, 808 nm, 2 W cm^{-2} for 5 min. (h) Photothermal effect analysis by monitoring the temperature change under laser irradiation, 808 nm, 2 W cm^{-2} for 5 min. (i) Schematic description of DNAzyme-mediated mRNA cleavage. (j) Gel electrophoresis of GD cleaving activity. Each error bar represents the standard deviation of four independent measurements.

Copper depletion by ZPD at the cellular level

Copper is usually associated with cancer development, and cancerous cells typically accumulate more copper to support rapid proliferation.^{11,50–52} To confirm this phenomenon, we measured cellular Cu content with and without Cu supplementation in a culture medium. As shown in Fig. 3a, in 4T1 tumor cells, copper content increased 3.6-fold with copper addition (about 5.4 fg per cell) compared to the one without supplementation (about 1.5 fg per cell), whereas there was no obvious change in normal L02 cells after Cu supplementation for 24 h. Prolonging the incubation time with copper to 48 h and 72 h showed similar trends, with increased copper uptake observed in 4T1 cells after copper addition (Fig. S9†).

Meanwhile, copper content in other cancerous cells was also detected, and the same phenomenon was observed, with increased copper content after Cu supplementation (Fig. S10†).

Given the higher copper accumulation within cancer cells and the copper-depleting property of ZPD *in vitro*, we explored its performance at the cellular level. ZPD exhibited dose-dependent cytotoxicity for cancerous cells, but had no obvious influence on normal cells (Fig. 3b and S11†). We inferred that the cytotoxicity towards cancerous cells was associated with ZPD-mediated copper depletion. To validate the occurrence of cellular copper depletion, high-angle annular dark-field scanning transmission electron microscopy (HAADF-STEM) imaging and elemental mapping were performed. Greater amounts of

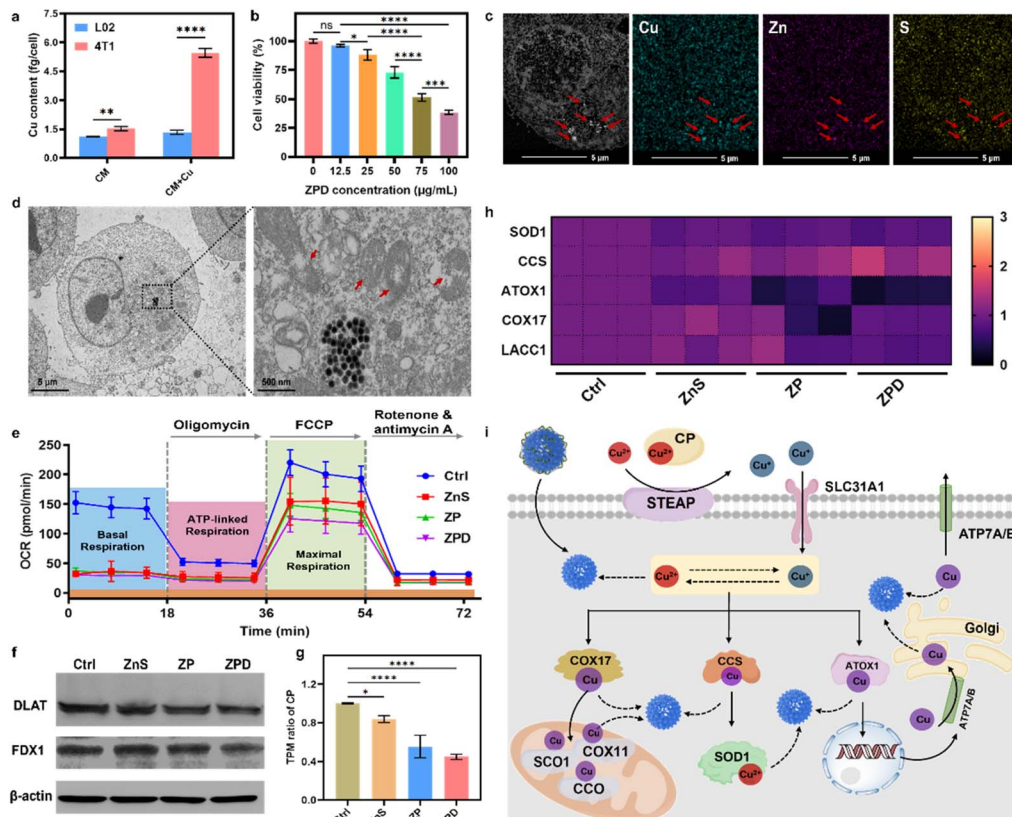


Fig. 3 Cellular copper depletion by “copper nanoconsumer” ZPD and mechanistic analysis. (a) Cellular copper content with/without copper supplement in a culture medium. (b) Cell viability analysis of ZPD at different concentrations in 4T1 cells. (c) HAADF-STEM and elemental mapping of ZPD-treated 4T1 cells. Red arrows indicate the position of endocytosed nanoparticles. (d) Bio-TEM analysis of ZPD-treated 4T1 cells. Red arrows indicate the damage of mitochondria. (e) Analysis of mitochondrial respiration by using the oxygen consumption rate (OCR). (f) Western blotting analysis of DLAT expression. (g) Transcripts per million (TPM) of ceruloplasmin (CP) in different treatments. (h) Heat map of TPM in copper-associated metalloenzymes including SOD1 (Cu/Zn superoxide dismutase), CCS (copper chaperone for superoxide dismutase), ATOX1 (antioxidant 1 copper chaperone), COX17 (cytochrome c oxidase assembly protein 17), and LACC1 (laccase (multicopper oxidoreductase) domain containing 1). (i) Possible steps of copper-depletion/copper-sequestration in cells. SOD1, Cu/Zn superoxide dismutase; CCS, copper chaperone for superoxide dismutase; ATOX1, antioxidant 1 copper chaperone; COX11, cytochrome c oxidase assembly protein 11; COX17, cytochrome c oxidase assembly protein 17; STEAP, belongs to SLC31A1 solute carrier family 31 member 1, as metalloreductases; CCO, cytochrome c oxidase; SCO1, synthesis of cytochrome c oxidase 1; ATP7A/B, for a copper transporter. Solid lines: established pathways; dashed lines: proposed pathways without known mechanisms. Each error bar in (a) and (b) represents the standard deviation of four independent measurements, in (e) represents the standard deviation of twenty independent measurements, and in (g) represents the standard deviation of three independent measurements. Ordinary one-way ANOVA was used for discrepancy analysis (* $P < 0.05$, ** $P < 0.01$, *** $P < 0.001$, and **** $P < 0.0001$).

Cu were observed in locations of endocytosed-nanoparticles, demonstrating Cu-ZnS exchange-mediated cellular copper depletion (Fig. 3c). In contrast, the distribution of Cu was uniform in untreated cells (Fig. S12†), further implying ZPD-mediated cellular copper depletion. Cellular copper depletion is associated with mitochondrial dysfunction.^{7,8,44} Bio-TEM shed light on this process. As shown in Fig. 3d, ZPD was successfully endocytosed by 4T1 cells and mitochondrial damage was clearly observed. Because of mitochondrial damage, the oxygen consumption rate (OCR), as an indicator of mitochondrial respiration, was monitored using a Seahorse XFe96 analyzer. 4T1 cells were co-incubated with ZnS, ZP, and ZPD. As shown in Fig. 3e, basal respiration and maximal respiration were reduced in each treated group compared to the untreated one, corroborating the copper depletion phenomenon.

Mechanism of copper depletion by ZPD

Having demonstrated cellular copper depletion by ZnS, we further explored its mechanism. The mechanism of copper overaccumulation-mediated cuproptosis is characterized by the aggregation of lipoylated protein and decrease in Fe-S cluster proteins.¹⁹ Therefore, we analyzed the expression of DLAT (dihydrolipoamide S-acetyltransferase, as one indicator of lipoylated protein) and FDX1 (ferredoxins, as a family of Fe-S cluster proteins) by western blotting (Fig. 3f). There was no obvious discrepancy in the expression of these two proteins, which was obviously changed in copper-accumulated cuproptosis with DLAT aggregation and FDX1 decrease. We conjectured that the mechanism of copper-depleting cuproptosis should be different from the overaccumulation mechanism. To get further insights into the copper depletion mechanism, cellular transcriptomic analysis was carried out. ZnS, ZP, and

ZPD showed different transcripts per million (*e.g.*, TPM) compared to the control group (Fig. S13†). In contrast, TPM of DLAT and Fe-S clusters showed no significant change compared to the control group (Fig. S14†). These results further indicated distinct mechanism of copper depletion from overaccumulation.

Ceruloplasmin (*e.g.*, CP), the predominant carrier of copper, has been regarded as an indicator of copper depletion.^{12,53,54} TPM of CP in “copper nanoconsumers” was significantly lower than in the control group (Fig. 3g), suggesting a cellular deficiency of available copper. Copper is an essential cofactor for important antioxidant enzymes, such as Cu/Zn superoxide dismutase (SOD1), copper chaperone for superoxide dismutase (CCS), antioxidant 1 copper chaperone (ATOX1), cytochrome c oxidase assembly protein 17 (COX17), and LACC1 (laccase (multicopper oxidoreductase) domain containing 1).^{2,3,12} Once copper is depleted in cells, the function of these copper-containing enzymes would likely be impaired. TPM of these metalloenzymes was investigated (Fig. 3g). Four enzymes (*i.e.*, SOD1, ATOX1, COX17, and LACC1) exhibited an obvious decrease in TPM, especially in the ZPD group. Because of lowered TPM of antioxidant enzymes, cellular ROS increased (Fig. S15†), consequently deactivating SOD1 and decreasing the GSH level (Fig. S16†), particularly in the ZPD group. These phenomena were similar to the reported performance of copper-chelator DC_AC5.⁵⁵

We further carried out gene function classification by GO (gene ontology) enrichment analysis for ZPD (Fig. S17 and S18†). To our surprise, the ZPD treatment was strongly associated with a series of cellular stress responses, including not only “detoxification of inorganic compounds” and “stress response to metal ions” but also “detoxification of copper ions” and “stress response to copper ions” (highlighted in the red squares). We speculated that these observations were associated with copper depletion. Taken together, it is reasonable to state that ZPD could deplete and sequestrate copper ions, leading to the dysfunction of copper-containing metalloenzymes and disruption of cellular homeostasis, ultimately promoting cell death. We illustrated the possible steps of the ZPD depleting/sequestrating cellular copper in Fig. 3h.

Copper depletion induces glycolytic inhibition and enables the photothermal effect

As demonstrated above, the copper depletion by ZPD disrupted mitochondrial respiration and consequently shifted cells more to glycolysis. Moreover, the GD conjugated onto ZPD could be specifically activated by cation exchange-released Zn²⁺, which would downregulate the expression of Glut 1 and further induce glycolytic inhibition (Fig. 4a). To demonstrate this hypothesis, Glut 1 expression was monitored by confocal imaging. As shown in Fig. 4b, fluorescence intensity of Glut 1 was notably downregulated in the ZPD group, which was also identified by Glut

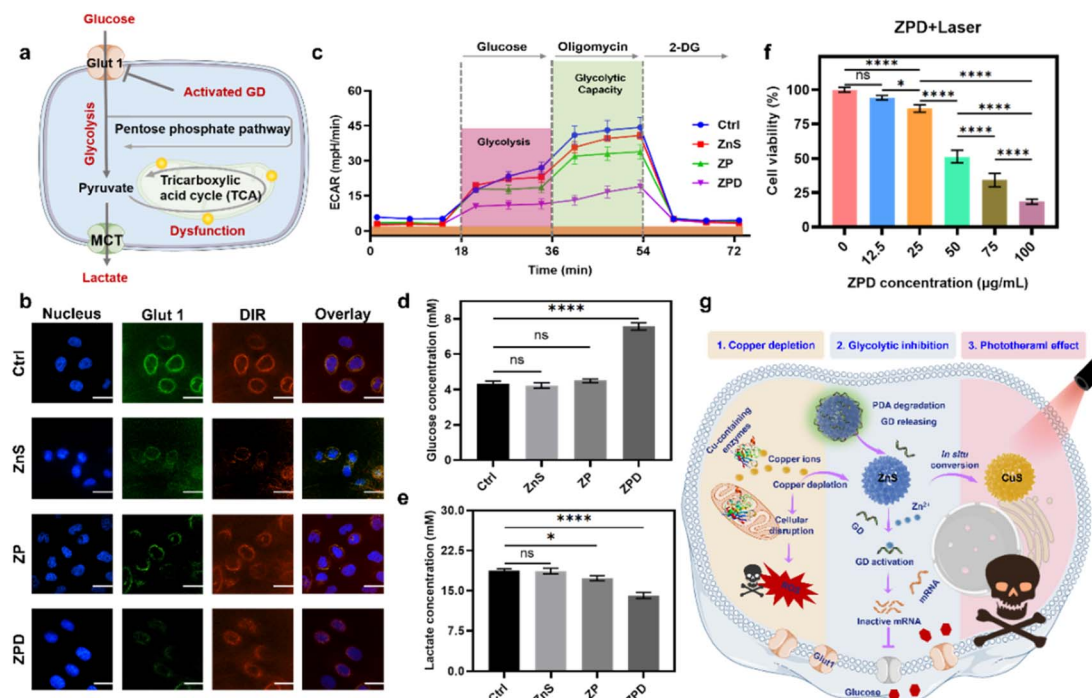


Fig. 4 Copper depletion induces glycolytic inhibition and enables the photothermal effect. (a) Schematic of glycolytic inhibition. (b) Analysis of Glut 1 expression after indicated treatments by confocal imaging. Scale bar: 5 μ m. (c) Analysis of glycolytic capacity by using the extracellular acidification rate (ECAR). (d) and (e) Glucose and lactate content in the culture supernatant after different treatments. (f) Cell viability after treatment with different concentrations of ZPD with laser irradiation (808 nm, 2 W cm^{-2} for 5 min). (g) Schematic of “copper nanoconsumer” ZPD’s multifunctionalities. Each error bar in (c) represents the standard deviation of twenty independent measurements and in (d)–(f) represents the standard deviation of four independent measurements. Ordinary one-way ANOVA was used for discrepancy analysis (* $P < 0.05$, ** $P < 0.01$, *** $P < 0.001$, and **** $P < 0.0001$).



western blotting analysis (Fig. S19†). Then, glycolytic function was measured by using the extracellular acidification rate (ECAR) using a Seahorse analyzer. As shown in Fig. 4c, the ZPD group exhibited the lowest glycolysis and glycolytic capacity due to GD introduction, while ZnS and ZP groups showed slightly lower levels than the control group. To further validate glycolytic inhibition, glucose and lactate contents in the culture supernatant were monitored. Glucose content in the ZPD-treated culture supernatant was higher than in other groups

(Fig. 4d), indicating less glucose uptake. Lactate, a product of glycolysis, was decreased in the treated culture supernatant (Fig. 4e), indicating glycolytic inhibition, especially in the ZPD-treated group. These results were consistent and demonstrated that ZPD could induce glycolytic inhibition.

Besides Zn^{2+} release after copper depletion, the photo-thermally active CuS was formed. Therefore, the photothermal effect was evaluated by irradiating ZPD-treated cells with an 808 nm laser at a power of 2 W cm^{-2} for 5 min. Greater cell growth

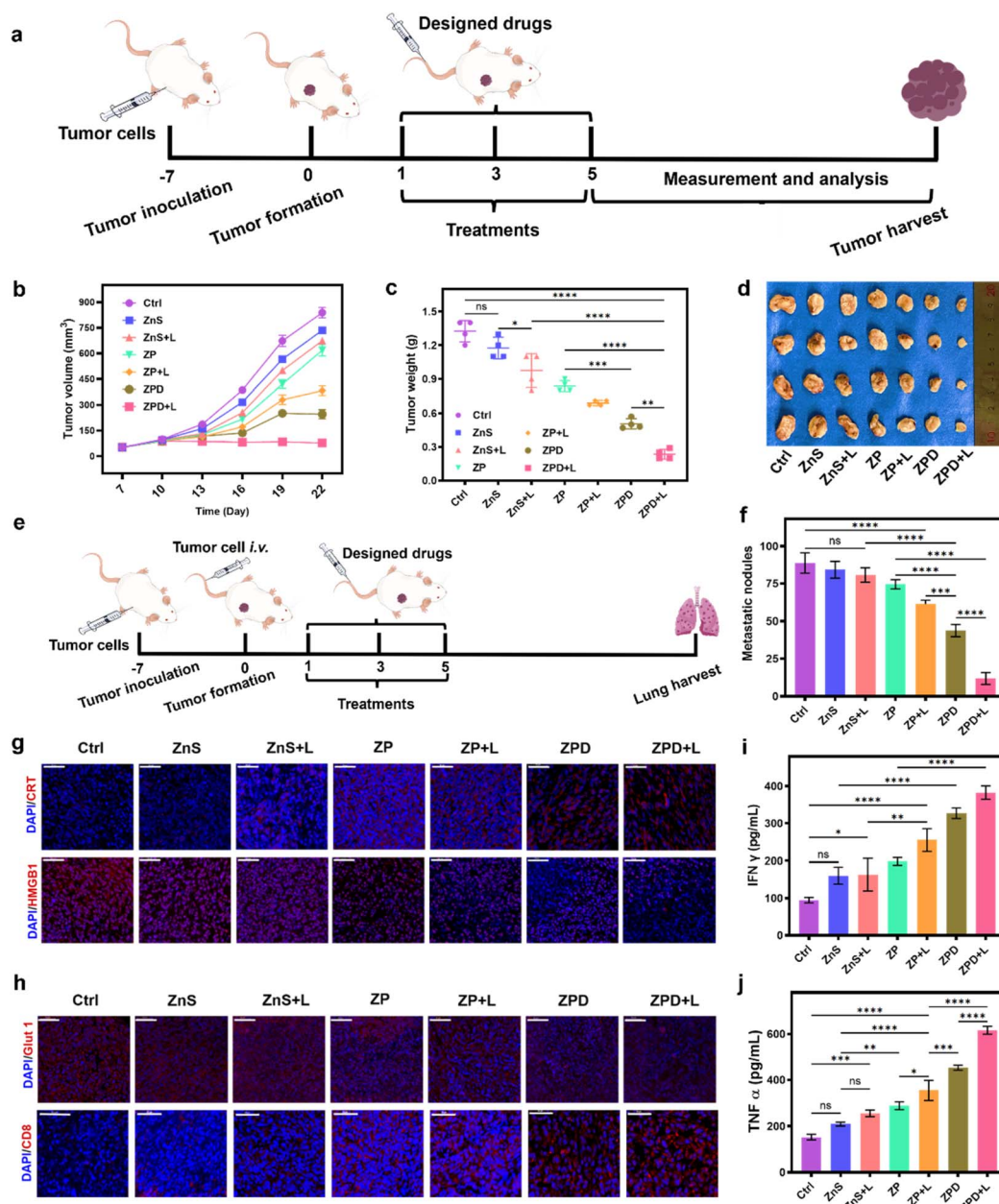


Fig. 5 Anti-primary tumor and anti-metastatic ability analysis. (a) Schematic of anti-primary tumor model construction and treatment. (b) Tumor volume changes over time. (c) Tumor weight analysis by groups. (d) Tumor images by groups. (e) Schematic of constructing the lung metastasis mouse model and treatment. (f) Statistical analysis of metastatic nodules in groups. (g) Immunofluorescence analysis of ICD molecules. (h) Immunofluorescence analysis of Glut 1 expression and CD8⁺ T cell infiltration. Cytokine detection of IFN- γ (i) and TNF- α (j). All designed drugs with intravenous injection at a dosage of 5 mg per kg per mouse. A 808 nm laser with a power of 2 W cm^{-2} is used for 5 min for laser irradiation. Each error bar represents the standard deviation of four independent measurements. Ordinary one-way ANOVA was used for discrepancy analysis (* $P < 0.05$, ** $P < 0.01$, *** $P < 0.001$, and **** $P < 0.0001$). Scale bar: 50 μm .



inhibition was achieved in the laser irradiation groups (Fig. 4f) compared to those without laser irradiation (Fig. 3b). Taken together, we concluded that the “copper nanoconsumer” ZPD could play multifunctional roles in copper depletion, glycolytic inhibition, and the photothermal effect (Fig. 4g), all of which would be significant for tumor synergetic therapy.

Anti-primary and metastatic tumor ability of ZPD

After the cellular studies, we subsequently evaluated the anti-tumor ability of ZPD *in vivo*. A subcutaneous 4T1 breast tumor model was constructed for different treatments (Fig. 5a). First, the biocompatibility was investigated by monitoring mouse weight changes (Fig. S20†) and histopathological staining of major organs (Fig. S21†). These results suggested no obvious side effects, demonstrating the good biosafety of nanoparticles, including ZPD. Furthermore, the biodistribution of Cu, Zn, and S with and without ZPD treatment was analyzed. As shown in Fig. S22,† the levels of Zn and S increased in the ZPD-treated group, especially in tumor areas, suggesting that ZPD could accumulate in tumor areas. Although liver demonstrated the highest content of Zn and S, this was rational due to function of liver and consistent with other nanomaterials’ biodistribution.^{56,57}

Second, the tumor volume and weight were monitored to evaluate therapeutic efficacy (Fig. 5b-d). Without laser irradiation, anti-tumor ability was shown as ZnS < ZP < ZPD. With laser irradiation, anti-tumor ability had a similar trend of ZnS + L < ZP + L < ZPD + L, with the ZPD + L group exhibiting the greatest anti-tumor performance. To demonstrate the broad applicability, the anti-metastatic ability of our nanoparticles was also evaluated (Fig. 5e, f and S23†). The trend of anti-metastatic ability among the designed groups was consistent with the anti-tumor performance. For mechanistic exploration of metastatic inhibition, we assumed that immunity may play a key role. Therefore, molecules of immunogenic cell death (ICD) such as CRT (calreticulin) and HMGB1 (high mobility group protein 1) were detected by immunofluorescence analysis (Fig. 5g and S24†). The ZPD + L group demonstrated the highest CRT exposure and HMGB1 release. Additionally, Glut 1 expression decreased in the ZPD and ZPD + L groups (Fig. 5h and S25†), which also benefited the immunotherapy. Immunofluorescence analysis of CD8⁺ T cell infiltration was also performed (Fig. 5i). The ZPD + L group showed the highest CD8⁺ T cell infiltration. Meanwhile, both IFN- γ (interferon γ) (Fig. 5i) and TNF- α (tumor necrosis factor α) (Fig. 5j), as important cytokines in the peripheral blood of mice, exhibited the highest levels in the ZPD + L group, demonstrating their immunotherapeutic potential. Overall, it was demonstrated that the “copper nanoconsumer” ZPD could successfully deplete cellular copper and therefore promote tumor therapy.

Conclusion

In summary, to address the challenges of copper over-accumulation-mediated tumor cuproptosis, we demonstrated copper depletion-induced cuproptosis using the “copper

nanoconsumer”. Unlike molecular copper chelators, the copper-depleting principle of our design was inspired by the large difference in K_{sp} values between ZnS and CuS, which could promote a cation reaction between ZnS and copper ions and mediate copper depletion. Our “copper nanoconsumer” exhibited good efficacy against both primary and metastatic tumors. Additionally, the mechanism of copper depletion-induced cuproptosis was proposed, which was mainly associated with dysfunction of copper-contained metalloenzymes. Furthermore, we investigated the representative copper-based therapies (Table S2†). Although some copper-based molecular drugs were in preclinical or clinical phases, due to their intrinsic shortcomings, further efforts are needed to advance copper-based tumor therapies.^{11,16,22,25,26,58-64} We envision that copper depletion-induced tumor cuproptosis will provide an alternative strategy for tumor therapy. Furthermore, by exploring the physiochemical characteristics of nanodrugs, more innovative therapeutics can be designed.

Data availability

Data supporting this study are available in the ESI† and further details are available from the authors on reasonable request.

Author contributions

M. Z., H. W. conceived the idea. H. W. supervised the work. M. Z., F. M., J. F. carried out *in vitro* and *in vivo* experiments. Y. Z., T. L. assisted with data analysis. Y. Z. assisted with schematic drawing. M. Z. wrote the original draft and H. W. provided the major revisions. All the authors discussed the results and commented on the manuscript.

Conflicts of interest

The authors declare no conflict of interest.

Acknowledgements

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