Chemical Science



PERSPECTIVE

View Article Online



Cite this: Chem. Sci., 2024, 15, 9915

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Received 31st March 2024 Accepted 5th June 2024

DOI: 10.1039/d4sc02129k

rsc.li/chemical-science

Current progress in the regulation of endogenous molecules for enhanced chemodynamic therapy

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Chemodynamic therapy (CDT) is a potential cancer treatment strategy, which relies on Fenton chemistry to transform hydrogen peroxide (H₂O₂) into highly cytotoxic reactive oxygen species (ROS) for tumor growth suppression. Although overproduced H₂O₂ in cancerous tissues makes CDT a feasible and specific tumor therapeutic modality, the treatment outcomes of traditional chemodynamic agents still fall short of expectations. Reprogramming cellular metabolism is one of the hallmarks of tumors, which not only supports unrestricted proliferative demands in cancer cells, but also mediates the resistance of tumor cells against many antitumor modalities. Recent discoveries have revealed that various cellular metabolites including H₂O₂, iron, lactate, glutathione, and lipids have distinct effects on CDT efficiency. In this perspective, we intend to provide a comprehensive summary of how different endogenous molecules impact Fenton chemistry for a deep understanding of mechanisms underlying endogenous regulation-enhanced CDT. Moreover, we point out the current challenges and offer our outlook on the future research directions in this field. We anticipate that exploring CDT through manipulating metabolism will yield significant advancements in tumor treatment.

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Introduction

Cancer has been a severe threat to human life.1 Currently, the main cancer treatments typically involve surgery, radiotherapy, and chemotherapy, yet they offer limited clinical benefits and are usually associated with serious side effects.2 Therefore, there is an urgent need to explore effective and tumor-specific treatment modalities aiming to completely eradicate tumors, prevent tumor recurrence and metastasis, and improve prognosis.3,4 Importantly, cancer cells show elevated reactive oxygen species (ROS) generation compared to non-cancer cells, owing to



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alterations in genetic and energy metabolism patterns.⁵ In contrast, excessive ROS can cause irreversible oxidative cell damage, ultimately resulting in cell death.6 Considering the vulnerability of cancer cells to oxidative stress, several ROS-based therapies have emerged, including photodynamic therapy (PDT),8 radiotherapy (RT),9 sonodynamic therapy (SDT),10 chemodynamic therapy (CDT),11,12 immunotherapy, etc.13

CDT, an emerging efficient ROS-mediated therapeutic modality, was first given by Shi and coworkers in 2016.14 It catalyzes the conversion of intracellular hydrogen peroxide (H₂O₂) through intratumoral Fenton or Fenton-like reactions, generating a cytotoxic hydroxyl radical ('OH), which is among the most harmful ROS.15-17 This ROS generation is directly activated by endogenous chemical energy conversion without the need for exogenous stimulations (e.g., laser, ultrasound, or X-rays), thus adequately overcoming the significant limitation of PDT, RT, and SDT, namely low tissue penetration depth.¹⁸⁻²⁰ Moreover, Fenton chemistry triggers the generation of intracellular cytotoxic radicals that are only specifically activated at tumor sites, enabling improved therapeutic effects and fewer side effects of CDT.²¹⁻²³ Consequently, substantial efforts have been made to engineer various Fenton reaction catalysts with amplified Fenton reaction performance for effectively treating tumors.24-27

Despite extensive attempts, the therapeutic outcome of CDT relying solely on Fenton reaction catalysts remains unsatisfactory.28 The main reason is that tumor cells can construct a robust defense system that allows them to escape from ROSinduced damage by metabolic adaption, inevitably diminishing the antitumor efficacy of CDT.29-31 Mounting evidence indicated that in specific biochemical contexts, aberrant metabolism of tumor cells can cause differential susceptibility of cells to CDT.32,33 A detailed understanding of how endogenous metabolic pathways in tumor cells influence CDT processes is of great therapeutic interest. Herein, we first summarize the mechanisms by which H₂O₂, iron, lactate, glutathione (GSH), and lipid metabolism impact Fenton chemistry-mediated CDT. Then, the current state-of-the-art designs of CDT nanomedicines integrating metabolic regulation abilities for enhanced tumor therapeutic efficiency will be critically discussed. Last but not least, we discuss the representative challenges and future directions for developing CDT



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agents. We expect the emergent metabolic reprogramming strategy to push forward the great process of research in the CDT field, which will benefit the clinical treatment of cancers.

Pathways controlling CDT

Considering that various endogenous molecules' metabolic processes interfere with the efficiency of Fenton-type reactions that are involved in CDT, researchers are beginning to focus on identifying the metabolic pathways that can impact therapeutic efficacies of CDT. The classifications of currently identified metabolic pathways in tumors, including H2O2, iron, lactate, GSH, and lipid metabolism, are summarized in Fig. 1.

Endogenous H₂O₂, as an important substrate of the Fenton reaction, can be produced from various cellular metabolic processes in cancer cells. For instance, electrons leaked from the endoplasmic reticulum stress or mitochondrial respiratory chain can be captured by molecular oxygen (O2) to form a superoxide ('O2"), which is further disproportionated into H₂O₂ by superoxide dismutase (SOD).³⁴ In addition, the production of H₂O₂ can also be initiated by activated nicotinamide adenine dinucleotide phosphate (NADPH) oxidase (NOX) or nicotinamide adenine dinucleotide (phosphate): quinone oxidoreductase1 (NQO1).35 And then, the ferrous ion (Fe2+) possesses more powerful 'OH-producing ability compared to other Fenton-reactive metal ions, enabling the strategy of regulating iron metabolism attractive for achieving efficient CDT.^{36,37} First, the extracellular transferrin (Tf)-ferric ion (Fe³⁺) (Tf-Fe³⁺) complex is carried into cells through cell surface Tf receptor protein 1 (Tfr1).38 In the endosome, ferrireductaseactive prostate 3 (STEAP3) mediates the transformation of Fe³⁺ into Fe²⁺ and divalent metal transporter 1 (DMT1) transports Fe²⁺ to a cytosolic labile iron pool (LIP).³⁹ Excess Fe²⁺ is excreted out of cells via ferroportin or is oxidized to Fe³⁺ by the ferritin heavy chain (FHC) with ferroxidase activity to initiate subsequent Fe sequestration in ferritin, which may be limiting the Fe-driven Fenton reaction,40 while iron deficiency can trigger nuclear receptor coactivator 4 (NCOA4)-dependent autophagic ferritin degradation, namely ferritinophagy, to release free iron on demand.41 Lactate metabolism regulation can interfere with the construction of the acidic tumor microenvironment (TME), which provides a feasible strategy to meet the strong acidity requirements of an effective Fenton reaction.42,43 Glucose is imported into cells by the glucose transporter (GLUT) on the cell membrane and subsequently transformed into pyruvate in the cytoplasm. Then lactate dehydrogenase (LDH) mediates the conversion between pyruvate and lactate, and the monocarboxylic acid transporter (MCT) mediates lactic acid efflux, inducing tumor acidification.44 The cellular antioxidative system plays an indispensable role in maintaining the equilibrium of a reductive environment against excessive ROS.45 Such a redox homeostasis can alleviate oxidative stress which can attack key biomacromolecules and cause irreversible cellular damage.46 As the most abundant lowmolecular-weight thiol-containing tripeptide synthesized in the cell cytosol, GSH has long been considered the representative reducing agent for constantly neutralizing cellular oxidative

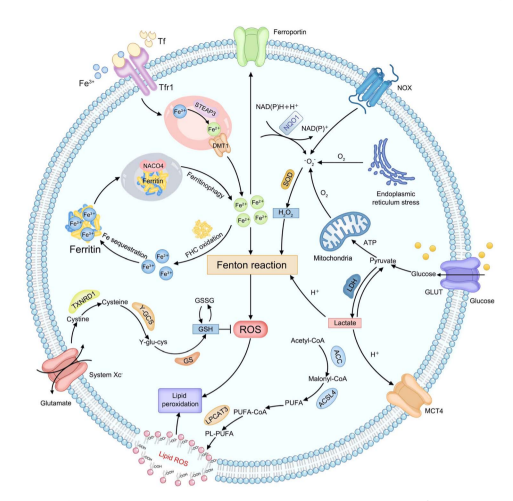


Fig. 1 Schematic illustration of endogenous metabolic pathways regulating Fenton chemistry-mediated CDT in tumor cells.

damage and detoxifying lipid ROS.47 The high GSH content in tumor cells can weaken the effect of CDT by directly scavenging ROS.48 For GSH synthesis, extracellular cystine is internalized by cells via the cystine/glutamate antiporter (system X_c^-) and reduced into cysteine by thioredoxin reductase 1 (TXNRD1).49 Then cysteine is conjugated with glutamate in the presence of γ glutamylcysteine synthetase (γ -GCS) to form γ -glutamylcysteine (γ -glu-cys). Finally, GSH synthetase (GS) catalyzes γ -glutamylcysteine and glycine to produce GSH. Lipid metabolism is also closely related to the effect of CDT.50 Typically, acetyl-CoA is first carboxylated to form malonyl-CoA by acetyl-CoA carboxylase (ACC). Free polyunsaturated fatty acid (PUFA) generated from malonyl-CoA can be esterified by activated acyl-CoA synthetase long-chain family member 4 (ACSL4) and is inserted into the cell membrane with the assistance of lysophosphatidylcholine acyltransferase 3 (LPCAT3), eventually improving sensitivity of cells to CDT.51 The above metabolic processes of endogenous molecules provide numerous practical targets for CDT treatment. Next, the mechanism of CDT regulation by each metabolic pathway will be elucidated and discussed in detail, using representative examples in the corresponding section, with the aim of providing references for designing more efficient chemodynamic agents.

2.1 Elevation of H₂O₂ contents

The H₂O₂ produced by cell metabolism can be converted into lethal 'OH by Fenton/Fenton-like reactions.52 Despite higher levels of H₂O₂ in cancer cells versus non-cancer cells, the endogenously produced H2O2 is still insufficient to assure a satisfactory tumor-killing effect.53 Developing pertinent strategies aimed at promoting the accumulation of intracellular H_2O_2 is still the primary focus of research on CDT. For example, Yang et al. constructed H₂O₂-responsive supramolecular polymers (PCSNs) by self-assembly of β-cyclodextrin-ferrocene (CD-Fc) conjugates with platinum(w) complex modification to overcome this shortcoming (Fig. 2A).54 The hydrophobic Fc in PCSNs was oxidized into water-soluble ferrocenium (Fc⁺) upon exposure to elevated H2O2 in the tumor. The disruption of hostguest interaction in the nanostructure resulted in dissociation behavior. Meanwhile, the released platinum(w) prodrug could be reduced into cisplatin(II) to activate NOX-mediated H2O2 replenishment, which could further promote the Fc-initiated Fenton reaction. In vivo experiments confirmed that such a positive feedback loop enabling self-boosting ROS generation dramatically inhibited tumor growth. Wang and co-workers developed amphiphilic polypeptide self-assembling nanomedicine (PtkDOX-NM) for the encapsulation of Fe³⁺ and β-

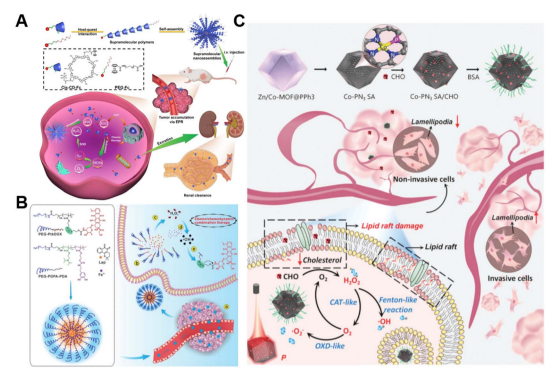


Fig. 2 (A) The main synthesis and therapeutic process of PCSNs. Reproduced with permission from ref. 54. Copyright 2021, Wiley-VCH GmbH. (B) Schemes displaying the construction process of PtkDOX-NM and its therapeutic mechanisms. Reproduced with permission from ref. 55. Copyright 2019, Wiley-VCH GmbH. (C) Illustrative diagram of the antitumor mechanism of Co-PN₃ SA/CHO. Reproduced with permission from ref. 59. Copyright 2023, Wiley-VCH GmbH.

lapachone (Lap), an o-naphthoquinone that can generate H₂O₂ via futile redox cycles under the catalysis of overexpressed NQO1 (Fig. 2B).55 PtkDOX-NM was formed by co-assembly of two amphiphilic polymers: poly(ethylene glycol) (PEG)-blockpoly diisopropylaminoethyl methacrylate block-poly dopamine (PEG-PDPA-PDA) with pH-responsiveness and PEG-block-poly cinnamaldehyde (CA)-Fc polyprodrug (PEG-PCASFc) with ROSresponsiveness, thus achieving dual responses to pH and ROS in the TME. Next, the hydrophobic-to-hydrophilic transition of PDPA responding to the acidity of endo/lysosomes in tumor cells resulted in nanostructure disassembly and Lap release. Lap-induced H₂O₂ generation could enhance 'OH production through the Fe³⁺-catalyzed Fenton reaction, causing the rapid release of the DOX prodrug and the further collapse of nanostructure cleavage of the ROS-responsive thioacetal linker. Their study demonstrated that CDT combined with H2O2 replenishment realizes efficient tumor inhibition with improved survival in mice.

Additionally, natural enzymes or artificial nanoenzymes have been widely applied to catalyze the transformation of intracellular substances such as glucose,56,57 lactate,58 and cholesterol into H₂O₂, which can provide an adequate substrate to accelerate the Fenton reaction. In a recent study by Lin's group, multienzyme-like Co-PN3 SA/CHO nanoagents were designed and constructed by utilizing a well-dispersed phosphorus (P)doped cobalt single atom nanozyme loaded with cholesterol oxidase (CHO) to achieve efficient tumor catalytic therapy for tumor metastasis inhibition (Fig. 2C).59 The as-prepared Co-PN₃

SA/CHO could concurrently mimic catalase (CAT), oxidase (OXD), and peroxidase (POD) to produce 'O2 and highly cytotoxic 'OH for cancer cell killing. Moreover, CHO effectively catalyzed the oxidization of intratumoral cholesterol to generate H₂O₂ and cholestenone, promoting the Co-PN₃ SA-mediated Fenton reaction. Simultaneously, the CHO-catalytic depletion of cholesterol resulted in the disruption of lipid raft integrity and the inhibition of invasive lamellipodium formation, suppressing tumor progression and metastasis. This work provides a promising strategy to improve therapeutic efficiency through the combination of CHO-mediated cholesterol consumption and Co-PN3 SA-induced 'OH generation. Notably, bacteria engineered with overexpression of respiratory chain enzyme II (NDH-2 enzyme) have also been utilized to promote intracellular H₂O₂ content elevation. For example, Fan et al. designed an integrative bioreactor by assembling magnetic Fe₃O₄ nanoparticles on the surface of NDH-2 enzyme-functionalized nonpathogenic bacterium Escherichia coli MG1655.60 During Escherichia coli MG1655 respiration, NDH-2 enzyme accepted electrons from NADH and subsequently transferred electrons to O₂ to generate H₂O₂. The Fe₃O₄ nanoparticles converted the sustainably synthesized H2O2 to produce 'OH through the Fenton-like reaction, achieving H2O2 self-supplying CDT.

2.2 Disruption of iron homeostasis

The conversion of intracellular H2O2 into 'OH via Fenton chemistry is dependent on Fenton catalysts.⁶¹ Fe²⁺ and its derivatives could serve as excellent Fenton catalysts due to their

high catalytic activity in 'OH generation.⁶² Generally, tumor cells require more iron ingestion compared to non-tumor cells to maintain rapid proliferation, while coordinating the regulation of iron metabolism to avoid excess iron-induced toxicity.63 The regulation of intracellular iron homeostasis encompassing iron storage/release and import/export has been regarded as a promising approach for reinforcing CDT. For instance, Bu's group prepared a mesoporous silicon nanocarrier loaded with amorphous elemental Fe⁰ and 2-amino-1,2,4-triazole (AT, a catalase inhibitor) (DMON@Fe⁰/AT) to integrate redox regulation and iron metabolism disruption for enduring CDT (Fig. 3A).64 Once DMON@Fe⁰/AT was endocytosed by tumor cells, acidic pH-triggered Fe2+ and AT release from this nanocarrier occurred. The released AT suppressed the catalase activity to promote H₂O₂ accumulation and the exposed DMONmediated GSH depletion, thus enhancing the Fe²⁺-driven Fenton reaction. The elevated intracellular 'OH induced mitochondrial dysfunction, leading to the downregulation of cell membrane protein ferroportin 1 expression and the further disruption of intracellular iron homeostasis. In short, the asprepared intelligent biodegradable DMON@Fe⁰/AT demonstrates a new method for effective CDT treatment by blocking the iron exporter and regulating redox homeostasis.

Our group recently proposed a Fe⁰-small interfering RNA (siRNA) composite system (Fe⁰-siRNA) to realize FHC downregulation-potentiated CDT (Fig. 3B).65 When the Fe⁰siRNA nanoparticles (Fe⁰-siRNA NPs) were internalized into tumor cells and located in mildly acidic lysosomes, they were disassembled in the presence of O2 and released Fe2+ and siRNA. The acidity amplification resulting from O₂ depletion could promote further degradation of Fe⁰-siRNA NPs. The Fe²⁺ from Fe⁰-siRNA NPs not only activated the conversion of intracellular H₂O₂ into highly toxic 'OH for CDT, but also facilitated endosome escape of FHC siRNA through 'OH-triggered lipid peroxide (LPO) of endo/lysosomal membranes. More importantly, FHC silencing with siRNA suppressed the transformation of Fe2+ into Fe3+ with less Fenton acitivity and the sequestration of Fe in ferritin, allowing the accumulation of higher reactive Fe²⁺, as well as increasing the CDT efficiency. In another study, Du et al. decorated a thiamine pyrophosphate (TPP)-etched open cavity metal-organic framework (MOF) with chitosan oligosaccharide (COS) through electrostatic interaction to develop a nanoplatform (termed COS@MOF) enabling hydrogen sulfide (H2S)-activated CDT and ferroptosis for colon cancer treatment (Fig. 3C).66 On the one hand, the upregulated H₂S could boost the transformation of Fe³⁺ into Fe²⁺ to improve

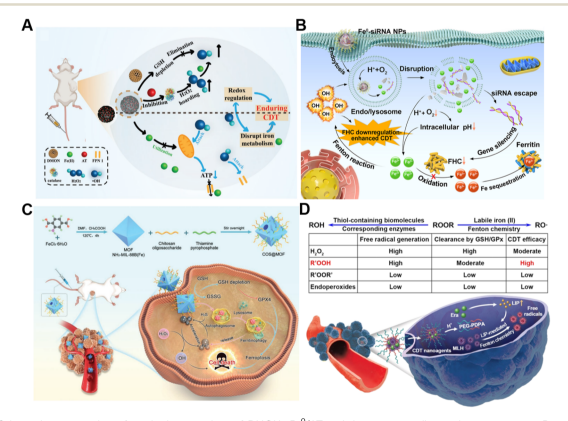


Fig. 3 (A) Schematic presentation of synthetic procedure of DMON@Fe⁰/AT and the corresponding anticancer process. Reproduced with permission from ref. 64. Copyright 2021, Wiley-VCH GmbH. (B) Schematic description of the sequential release of Fe²⁺ and siRNA from Fe⁰-siRNA NPs responding to tumor acidity for enhancement of CDT. Reproduced with permission from ref. 65. Copyright 2023, Wiley-VCH GmbH. (C) Illustration of an engineered MOF with amplifying ferritinophagy to cause iron ion overload for improved antitumor therapy. Reproduced with permission from ref. 66. Copyright 2023, Wiley-VCH GmbH. (D) The scheme of factors affecting endogenous labile iron-based CDT outcome of ROOR and the utilization of MLH-loading nanomedicines for CDT. Reproduced with permission from ref. 68. Copyright 2020, American Chemical Society.

Fenton reaction efficiency. On the other hand, utilization of autophagy inducer COS enabled the degradation of ferritin to promote the release of iron from ferritin and Fe2+ replenishment, further enhancing CDT and ROS-mediated ferroptosis. In another example, Li et al. developed a nanodrug (T10@cLAV) by the integration of transferrin-homing peptide T10 and crosslinked lipoic acid vesicles for Fe2+ self-supplying tumor therapy.67 The T10@cLAV bound to the overexpressed Tf receptor on cancer cells (Tf@T10@cLAV) and was subsequently imported into endosomes/lysosomes. The acidic environment of endosomes/lysosomes could trigger the degradation of Tf, thus inducing the sustained release of iron ions and activating the Fe-driven Fenton reaction. More importantly, cLAV was disrupted with the assistance of GSH and TXNRD1 to release dihydrolipoic acid (DHLA) for DHLA-mediated reduction of Fe³⁺ to Fe²⁺, resulting in an increase in Fe²⁺ levels to further enhance nanocatalytic therapy. Completely unlike traditional treatment strategies involving the introduction of exogenous Fenton-type metal ions, our group first utilized the intracellular labile LIP to provide the continuous catalyst for catalytic generation of toxic free radicals through Fenton-type chemistry (Fig. 3D).68 We developed a pH-sensitive polymer for the co-delivery of the model ROOH molecule and LIP-increasing agent, which can sequentially respond to pH and labile iron in the TME to realize enhanced CDT.

2.3 Interference of lactate metabolism

In addition to the Fenton catalyst activity, the reaction conditions, particularly acidity, also influence the reaction kinetics of the Fenton reaction.⁶⁹ It has been confirmed that acidic pH

favors the Fenton reaction.70 Lactate is considered a metabolic byproduct of glycolysis in tumor cells, leading to elevated tumor acidity.71 Consequently, lactate metabolism regulation is an alternative strategy for enhancing CDT.72 Shi's group reported a pH-responsive and self-augmented nanoplatform (defined as FePt@FeO_x@TAM-PEG) fabricated by incorporating a core-shell FePt@FeO_r nanocatalyst and pH-responsive tamoxifen (TAM) drug in a poly(styrene-co-maleic anhydride) (PSMA) polymer polymeric matrix and further modifying the surface with PEG (Fig. 4A).⁷³ Once internalized by cancer cells, the nanoplatform disintegrated due to acidic pH-triggered hydrophobic-hydrophilic transitions of TAM and released the FePt@FeOr nanocatalyst. The FePt@FeOx nanocatalyst catalyzed the inherent H₂O₂ decomposition to produce lethal 'OH to damage cancer cells. Meanwhile, the liberated TAM could suppress the mitochondrial complex I, thus increasing the intracellular lactate level as well as elevating environmental acidity. This simultaneously amplified 'OH production and accelerated disassembly of the FePt@FeOx@TAM-PEG nanostructure, inducing more therapeutic cargo release. As a result, this intelligent nanoplatform overcomes the limitation of weak acidic conditions in tumors by introducing metabolism regulators to upregulate lactate content, thereby achieving the enhanced antitumor effect of CDT. Another related study involving the transport modulation of lactate to remodel the reaction environment is worth noting as well. Wang et al. prepared a calcium phosphate-based biomineralized multifunctional nanosystem with the co-delivery of a DOX-Fe²⁺ complex and MCT4-inhibiting siRNA (siMCT4) (CaP-DOX@Fe²⁺-siMCT4-PEG-HA) for interfering lactate efflux and enhancing anti-tumor CDT (Fig. 4B).74 The nanosystem was specifically taken up by tumor cells with the aid of hyaluronic

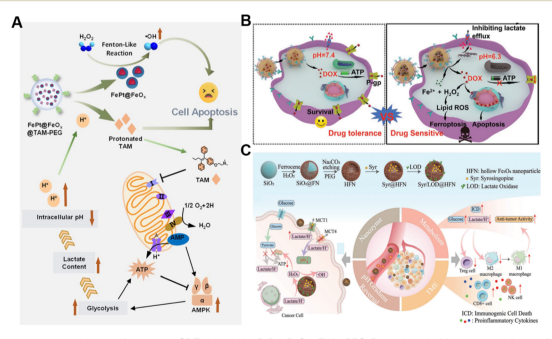


Fig. 4 (A) The underlying mechanism of enhanced CDT induced by $FePt@FeO_x@TAM-PEG$. Reproduced with permission from ref. 73. Copyright 2021, Wiley-VCH GmbH. (B) Schematic diagram of CaP-DOX@ Fe^{2+} -siMCT4-PEG-HA enabling TME remodeling for improved CDT. Reproduced with permission from ref. 74. Copyright 2021, Wiley-VCH GmbH. (C) Illustration of the construction and application of the Syr/LOD@HFN nanoplatform. Reproduced with permission from ref. 76. Copyright 2023, Wiley-VCH GmbH.

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acid (HA)-mediated targeting and degraded in the acidic environment of lysosomes due to the pH-triggered hydrolysis of CaP, resulting in the dissociation of the DOX-Fe2+ complex and the release of Fe²⁺, DOX, and siMCT4. MCT4 silencing by siMCT4 could exacerbate intracellular acidification by blocking lactate efflux, which not only enhanced the Fe²⁺-driven Fenton reaction, but also decreased adenosine triphosphate (ATP) production for inhibition of DOX efflux, thus significantly improving the treatment efficiency. Typically, Wang's group developed an intelligent bioreactor (Sa@FeS) in which ferrous sulfide (FeS) nanoparticles were anchored on the surface of Salmonella typhimurium strain (Sa) via biomineralization.75 The effective tumor penetration of Sa enabled FeS nanoparticles to mediate photothermally enhanced Fenton catalytic reactions in deep tumor tissues upon 1064 nm laser irradiation. At the same time, the H₂S produced by Sa metabolism can facilitate glucose uptake by tumor cells, resulting in the elevation of intracellular lactic acid levels and ultimately boosting FeS-mediated CDT enhancement. Given the limited therapeutic effect of single lactate consumption, simultaneous regulation of multiple metabolic pathways may be a promising way to enhance CDT. Wu et al. proposed a strategy with a dual effect on lactate metabolism based on lactate oxidase

(LOX) and syrosingopine (Syr) co-loaded hollow Fe₃O₄ nanoparticles (denoted as Syr/LOD@HFN), which could combine LOX-mediated lactate consumption and lactate transport chain blockade (Fig. 4C).76 In the tumor TME, LOX could catalyze lactate to generate H₂O₂ in situ while Syr-induced MCT4 downregulation could achieve lactate metabolism blockade, leading to continuous H₂O₂ production and pH reduction for boosting the Fe₃O₄-catalyzed Fenton reaction. Owing to the augmented ROS production and the reversion of the lactate-related immunosuppressive TME, this nanoplatform caused severe immunogenic cell death in tumor cells and promoted antitumor immunity. This work reveals that manipulation of multiple lactate metabolic pathways enables a better therapeutic effect than a single regulation strategy.

2.4 Depletion of GSH

It is well established that GSH is capable of potent ROS scavenging, thus endowing cancer cells with the ability to resist oxidative stress and weaken the effect of CDT. Generally, metals with oxidation states could consume GSH via undergoing redox reactions with GSH.77 Our group first reported the synthesis of

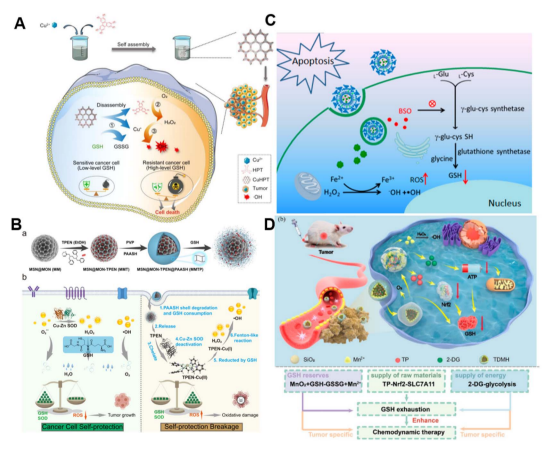


Fig. 5 (A) The synthesis of CuHPT and scheme of its antitumor mechanisms against cancer cells. Reproduced with permission from ref. 79. Copyright 2022, American Chemical Society. (B) Illustration for the fabrication procedure and multifunctional application in vivo of MSN@MON-TPEN@PAASH nanoparticles. Reproduced with permission from ref. 82. Copyright 2019, American Chemical Society. (C) A diagram of BSO/GA-Fe(II)@liposome for enhanced CDT via reducing GSH biosynthesis. Reproduced with permission from ref. 84. Copyright 2021, American Chemical Society. (D) Schematic illustration of the use of the TDMH nanomedicine for multistage GSH depletion-augmented CDT. Reproduced with permission from ref. 85. Copyright 2022, American Chemical Society.

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manganese dioxide (MnO₂)-coated mesoporous silica nanoparticles for precise magnetic resonance imaging (MRI)-guided self-reinforcing CDT. Upon encountering the high level of GSH, MnO₂ NPs were degraded to release Mn²⁺, which not only initiated CDT by Mn²⁺-mediated Fenton-like chemistry but also acted as MRI contrast agents to monitor the CDT process.⁷⁸ downregulated solute carrier family 7 member 11 (SLC7A11) expression and decreased cysteine uptake, and its combination with the glycolysis inhibitor 2-DG could significantly reduce the intracellular ATP level and block the regeneration of GSH synthesis. Owing to the synergistic effect of GSH depletion and GSH synthesis inhibition, this nanoplatform efficiently

particles for precise magnetic resonance imaging (MRI)-guided self-reinforcing CDT. Upon encountering the high level of GSH, MnO₂ NPs were degraded to release Mn²⁺, which not only initiated CDT by Mn²⁺-mediated Fenton-like chemistry but also acted as MRI contrast agents to monitor the CDT process.78 Recently, redox-responsive nanosized frameworks coordinated with Cu²⁺ and 2,3,6,7,10,11-hexahydroxytriphenylene (HPT, a catechol ligand), namely CuHPT, were designed to eliminate drug-resistant cancers by disrupting cellular redox homeostasis (Fig. 5A).79 Experimental results indicated that upon being activated by GSH, CuHPT nanoparticles displayed enhanced CDT with remarkable tumor growth inhibition and negligible side effects via a combination of auto-oxidation and Cu+-catalyzed reactions. In addition to introducing oxidizing agents to consume GSH, numerous electrophilic reagents that can directly react with the thiol group in GSH for reduction of GSH content were utilized.80,81 Guo et al. fabricated a hybrid mesoporous silica/organosilicate nanocomposite (MSN@MON) followed by coating the surface with disulfide bond-containing link poly(acrylic acid) and loading with N,N,N',N'-tetrakis(2pyridinyl methyl)-1,2-ethanediamine (TPEN, a metal ion chelator) to obtain MSN@MON-TPEN@PAASH (Fig. 5B).82 Because of GSH-induced disulfide bond cleavage in MSN@MON-TPEN@PAASH, intracellular GSH was partially consumed and the NPs collapsed to release TPEN. The liberated TPEN could chelate with Cu2+ in the Cu-Zn superoxide dismutase, enabling its deactivation and the formation of a TPEN-Cu(II) complex. Then, the TPEN-Cu(II) complex was reduced into TPEN-Cu(1) by remaining GSH, inducing the further decline of antioxidant defense resulting from GSH and ultimately promoting TPEN-Cu(1)-driven Fenton-like reactions for selfreinforced CDT.

Normally, GSH biosynthesis occurs in the cytosol of almost all cells.83 Blocking GSH synthesis using agents has been a good way to decrease GSH levels in cancer cells. Dong and co-workers reported a biocompatible liposomal nanoformulation containing L-buthionine sulfoximine (BSO) and gallic acid-ferrous ion (GA-Fe(II)) complexes to inhibit the synthesis of GSH for combined CDT-RT (Fig. 5C).84 Interestingly, GA boosted the transformation of Fe³⁺ into Fe²⁺, promoting the Fenton reaction to generate highly cytotoxic 'OH. Besides, BSO was capable of limiting the GSH synthesis rate via inhibiting the activity of glutamate-cysteine ligase to reduce GSH levels, thus intensifying the Fenton reaction-enabled cancer cell damage and CDT. Moreover, Huang and co-workers developed a multi-faceted GSH depletion-potentiated CDT system (TP/2-DG@HMnO2@-HA, TDMH), in which 2-deoxy-p-glucose (2-DG) and triptolide (TP) were co-encapsulated in hollow mesoporous MnO₂ (HMnO₂) followed by modifying HA on the nanoparticles for tumor targeting (Fig. 5D).85 When the nanomedicine reached the tumor area, MnO2 participated in a redox reaction with GSH, yielding glutathione disulfide (GSSG) and Mn²⁺ that could drive the Fenton-like reaction with the assistance of bicarbonate to lead to intracellular GSH consumption as well as 'OH production. Meanwhile, the disintegration of the HMnO2 structure occurred, triggering the release of TP and 2-DG. TP

2.5 Regulation of lipid metabolism

strengthened CDT against tumors.

The ROS accumulation induced by Fenton-type reactions can generally cause LPO. PUFAs, a family of membrane lipids possessing two or more carbon-carbon double bonds, are extremely susceptible to ROS-initiated LPO.86,87 Therefore, the CDT susceptibility of cancer cells can be raised by increasing PUFA content in the lipid chain.88 Taking advantage of this strategy, our group constructed a chemodynamic nanoagent (OA@Fe-SAC@EM) capable of modulating PUFA metabolism by loading oleanolic acid (OA) onto a single-atom Fe-anchored hollow carbon nanosphere (Fe-SAC) and further coating with an erythrocyte membrane (EM) for enhanced CDT (Fig. 6A).89 After internalization by tumor cells, Fe-SAC could catalyze intracellular H2O2 to generate 'OH, which not only mediated CDT but also induced the disruption of the surface EM for the controlled release of OA. Notably, OA caused a dramatically elevated expression of ACSL4, thus increasing membrane unsaturation via enriched PUFAs. As a result, this nanoagent exerted enhanced chemodynamic efficiency by LPO amplification. Another relevant study was developed by Liu and coworkers. They successfully prepared the FeCo/Fe-Co dualatom nanozyme (FeCo/Fe-Co DAzyme) possessing fourenzyme-like activities and subsequently loaded lipoxygenase (LOX) and phospholipase A2 (PLA2) into these NPs for the fabrication of a six-enzyme co-expressed nanoplatform (FeCo/ Fe-Co DAzyme/PL) to enhance catalytic therapy and immunotherapy (Fig. 6B).90 FeCo/Fe-Co DAzyme from this nanomedicine exhibited POD-like and OXD-like activities to catalyze both O_2 and H_2O_2 to generate O_2 and OH, as well as exerted glutathione oxidase (GSHOx)-like activity for GSH elimination to reduce 'OH consumption and enhanced ROStriggered LPO. Meanwhile, FeCo/Fe-Co DAzyme also in situ converts H2O2 into O2 that could combine with free arachidonic acid (AA) liberated from PLA2-catalyzed phospholipids, producing AA-OOH through the catalysis of LOX. These two processes synergistically induced an LPO self-amplification effect for effectively initiating immunogenic ferroptosis. The resulting systemic immune activation mediated the reprogramming of ACSL4-associated phospholipids and further strengthened the cascade of immunogenic tumor ferroptosis when combined with AA. Therefore, this strategy targeting lipid metabolism provides new insight into the design of ROSactivated immunotherapy. Additionally, Yang and co-workers co-loaded Fenton catalysts hemin and lipoxidase into a pHresponsive self-assembled CaCO3-encapsulated poly(lactic-coglycolic acid) (PLGA) nanoreactor for cascade catalytic amplification of tumor oxidative stresses.91 Upon entering tumor cells, CaCO₃ could be degraded the acidic

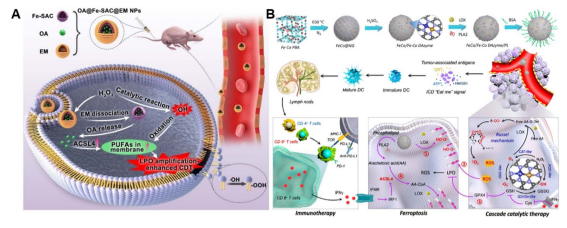


Fig. 6 (A) Schematic diagram of the preparation process of OA@Fe-SAC@EM as a self-reinforcing CDT agent *via* the manipulation of cell membrane unsaturation. Reproduced with permission from ref. 89. Copyright 2023, Wiley-VCH GmbH. (B) Schematic illustration of the design and proposed therapeutic mechanism of FeCo/Fe-Co DAzyme. Reproduced with permission from ref. 90. Copyright 2023, American Chemical Society.

microenvironment to release hemin and lipoxidase. Hemin catalyzed the Fenton reaction to convert H_2O_2 into toxic 'OH for ROS-initiated LPO. Moreover, lipoxidase further oxidized PUFA to generate lipid hydroperoxides and led to massive lipid hydroperoxide accumulation as well as aggravated membrane LPO, eventually causing tumor cell death.

3. Summary and outlook

In this perspective, we focused on the metabolic pathways of endogenous molecules within the TME and discussed the availability of these pathways in modulating Fenton chemistry for antitumor CDT. We briefly outlined the fundamental principles of the regulatory network involving endogenous molecules. We then thoroughly discussed the relevance of Fenton chemistry to $\rm H_2O_2$, iron, lactate, GSH, and lipid metabolism. In these sections, we described recent progress in developing chemodynamic agents that modulate endogenous metabolites to strengthen tumor CDT and tried to attract broad interest of researchers from various fields. While interfering with multiple endogenous metabolic regulations has shown promise in boosting the effectiveness of traditional CDT therapies, addressing the following issues is crucial to advancing the development of CDT.

Firstly, the endogenous metabolic pathways of cancer cells varied greatly due to the significant difference in tumor types, sizes, locations, and stages. Therefore, selecting appropriate experiment models and developing antitumor strategies with enhanced targeting ability are essential to achieve precise therapy. Secondly, the metabolic network of cancer cells is complex, and there can be crosstalk among different metabolic pathways involving various molecules. Inhibiting metabolic pathways of a single endogenous molecule may lead to the development of compensatory metabolic pathways in tumor cells, conferring resistance to the single intervention and compromising therapeutic efficacy. Undoubtedly, gaining

a comprehensive understanding of the interdependencies among different metabolism systems would provide us with new ideas for the design of endogenous regulation-based CDT. Additionally, we have only explored several endogenous metabolic pathways while the abundance of other cellular metabolites has not been exploited for their roles in influencing CDT, where research in this field is much needed. Meanwhile, it is important to consider that most of the metabolic pathways associated with alternative molecules are pervasively present in both normal and cancer cells. Thus, plenty of efforts should be devoted to enhancing the tumorspecificity of these chemodynamic nanomedicines that regulate endogenous metabolites, aiming to mitigate their adverse effects on normal tissues and organs. Despite the existence of certain challenges, growing knowledge about Fenton chemistry and its related endogenous regulators foresees the emergence of novel avenues and significant potential for endogenous regulation-based strategies to enhance CDT in the field of anti-tumor therapy.

Data availability

No primary research results, software or code have been included and no new data were generated or analysed as part of this perspective.

Author contributions

All authors participated in this review work. J. Wang, T. T. Cui, and L. S. Lin conceived the structure of this perspective. J. Wang, Y. N. Liu, and T. T. Cui prepared the draft. H. H. Yang and L. S. Lin reviewed and refined the manuscript.

Conflicts of interest

The authors declare no conflict of interest.

Acknowledgements

This work was supported by the National Key Research and Development Program of China (2020YFA0210800), the National Natural Science Foundation of China (22374026 and 22027805), and the Major Project of Science and Technology of Fujian Province (2020HZ06006).

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