Photoactivatable nanogenerators of reactive species for cancer therapy

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A B S T R A C T

In recent years, reactive species-based cancer therapies have attracted tremendous attention due to their simplicity, controllability, and effectiveness. Herein, we overviewed the state-of-art advance for photo-controlled generation of highly reactive radical species with nanomaterials for cancer therapy. First, we summarized the most widely explored reactive species, such as singlet oxygen, superoxide radical anion (O₂•−), nitric oxide (•NO), carbon monoxide, alkyl radicals, and their corresponding secondary reactive species generated by interaction with other biological molecules. Then, we discussed the generating mechanisms of these highly reactive species stimulated by light irradiation, followed by their anticancer effect, and the synergetic principles with other therapeutic modalities. This review might unveil the advantages of reactive species-based therapeutic methodology and encourage the pre-clinical exploration of reactive species-mediated cancer treatments.

1. Introduction

Reactive species are necessary regulatory substances in various physiological functions [1, 2]. As the most abundant components in organisms, the common nonmetal reactive species are generally based on oxygen, nitrogen, carbon, sulfur, etc [3–5]. Reactive species include a collective term of non-radical molecular forms (hydrogen peroxide (H₂O₂), CO, etc.) and the radical forms. Radicals refer to those atoms, ions, or molecules that contain unpaired electrons in the outermost electron orbitals, such as alkyl radical [6], nitric oxide (•NO) [7–8], peroxynitrite anion (ONOO−) [9], hydroxyl radical (•OH) [10], chlorine radical (•Cl) [11] and superoxide radical anion (O₂•−) [12]. These reactive species generally possess a short lifetime due to their high reactivity [13, 14]. Low concentrations of reactive species exhibit beneficial effects in immune response and maintenance of redox homeostasis and normal metabolisms of the cells [15–17]. In contrast, abnormal levels of reactive species can also lead to oxidative stress and even cell death by irreversibly damaging biofunctional macromolecules (nucleic acid, lipids, and proteins) [18, 19]. The human body has an antioxidant defense system, comprising some small-molecule antioxidants (e.g. Vitamin C and glutathione) and antioxidant enzymes (superoxide dismutases, catalase, glutathione peroxidases, peroxiredoxins, etc.) [20–22]. These antioxidants can abolish reactive species-induced oxidative stress to maintain the redox balance of the biological system [23].

Accumulated evidence has indicated that cancer cells possessed a high level of reactive species [24]. To keep the redox balance, most cancer cells have to generate more endogenous antioxidants [25]. The flexible alteration ability of redox status reduced the therapeutic effect of chemotherapy and radiotherapy on malignant tumor cells. Interestingly, some experimental results suggested that reactive species could exhibit higher damaging effects to cancer cells than normal cells [26]. The preferential inhibition outcome might be attributed to the direct suppression of reactive species to endogenous antioxidants in cancer cells. Despite reactive species-based nanomedicine have intrinsic advantages, reactive species-based cancer therapy is restricted by several limitations: (1) oxygen dependence limited the therapeutic effects to hypoxic tumors; (2) the short absorption wavelength of photactivatable nanogenerators kept traditional reactive species-based nanoparticle platform from eliminating deep-seated tumors; (3) undesirable side effects derived from the nonspecific distribution of nanogenerators remain an issue. Regarding these concerns, researchers have been devoted to designing and creating therapeutic platforms derived from...
Abbreviations

| Abbreviation | Definition |
|--------------|------------|
| NPs          | Nanoparticles |
| PTT          | Photothermal therapy |
| NMOFs        | Nanoscale metal-organic frameworks |
| ACQ          | Aggregation caused quenching |
| AIE          | Aggregation-induced emission |
| FRET         | Förster resonance energy transfer |
| NIR          | Near-infrared |
| CONs         | Covalent organic nanosheets |
| EPR          | Enhanced permeability and retention |
| DOX          | Doxorubicin |
| UCNPs        | Upconversion nanoparticles |
| TPZ          | Tirapazamine |
| OVA          | Ovalbumin |
| AuNCs        | Au nanocages |
| PDT          | Photodynamic therapy |
| PSs          | Photosensitizers |
| POP          | Porous organic polymer |
| SPNs         | Semiconducting polymer nanoparticles |
| CA-IX        | Carbonic anhydrase IX |
| P-gp         | P-glycoprotein |
| PTX          | Paclitaxel |
| iNOS         | NO synthase |
| DTX          | Docetaxel |
| BSA          | Bovine serum albumin |
| Hb           | Hemoglobin |
| PCM          | Phase change material |
| AIPH         | 2,2'-azobis[2-(2-imidazolin-2-yl)propane]dihydrochloride |

Scheme 1. Schematic illustration of the photoactivatable nanogenerators of reactive species including reactive oxygen, reactive nitrogen, and reactive carbon species for cancer therapy.
reactive species-based monotherapy or the featured synergistic therapy with other therapeutic strategies.

The formation of reactive species in cells can originate from internal and external stimuli. The internal stimuli are specifically derived from some metabolic processes [27–32], or pathological conditions [33]. The external triggers are mostly starting from a range of physicochemical or biochemical reactions induced by reactive species generators. Growing bodies of the experimental data indicated that reactive species generators activated by light irradiation are excellent therapeutic candidates due to their noninvasiveness and spatiotemporally controllable profile in biomedical applications [34]. In this review, we first summarized the recent advances of photo-activatable nanogenerators for reactive species including reactive oxygen species, reactive nitrogen species, and reactive carbon species (Scheme 1). We then discussed the principles for the design of photoactivatable nanoplatforms and their application for cancer therapy. We finally provided our perspectives for improving the therapeutic effect and promoting the clinical translation of the photo-activatable nanoparticles.

2. Reactive oxygen species-based antitumor therapy

Reactive oxygen species (ROS) is composed of reactive radicals or non-radical molecules evolved from oxygen atoms. It is a collective term including singlet oxygen (\(1^1O_2\)), superoxide radical anion (\(O_2^-\)), hydroxyl radical (\(\cdot OH\)), hydrogen peroxide (\(H_2O_2\)) [35]. ROS can destroy tumor cells through multifactorial mechanisms, including oxidizing the biological macromolecules, destroying angiogenesis to cut off the supply of nutrients to tumor cells [36,37]. Typically, five therapeutic strategies including radiation therapy [38–41], chemodynamic therapy [42–46], sonodynamic therapy [47–50], bioreductive chemotherapeutics (e.g., tirapazamine, TPZ) [51–54], and photodynamic therapy (PDT) [55–62] can generate ROS. Among all these approaches, PDT-based cancer therapies have been extensively exploited.

PDT can be generally classified into oxygen-independent type-I and oxygen-dependent type-II mechanisms. The type-II PDT procedure can transform the photon energy to the photosensitizers (PSs) by exciting the PSs from the ground state to the singlet state. The excited singlet state PSs can further be converted to an excited triplet state through the intersystem crossing process. The excited triplet state PSs eventually transfer the photo energy to the molecular oxygen for generating singlet oxygen (\(1^1O_2\)). In contrast, the type-I PDT can generate superoxide radical anions and hydrogen peroxide without the involvement of molecular oxygen (Fig. 1). The occurrence ratio of the two types of PDT is determined by the types of PSs and the concentration of the environment oxygen [63]. Furthermore, PDT can induce both apoptotic and necrotic death of the tumor cells. Notably, the dying tumor cells crack the plasma membrane and spill out the cytosolic constituents for provoking the inflammatory response, which can activate the protective immune response and suppress tumor growth via recruiting the tumor-infiltrating cytotoxic T lymphocytes [64]. In this section, we mainly discuss the advance of PDT-triggered reactive species generation with the nanogenerators.

2.1. Photactivatable nanoparticles for Type-II PDT of cancer

Given the successful application of hematoporphyrin for PDT, extensive efforts have been devoted to designing various PSs and exploring their biomedical application. Up to date, both inorganic and organic nanomaterials have been exploited for PDT. The inorganic nanomaterials display superior colloidal stability in the physiological milieu [65–70]. The organic PSs harbor the special merits of good biocompatibility and biodegradability. Various organic nanomaterial-based PSs, such as porphyrin [71–75], boron dipyrromethene (BODIPY) [76–78], cyanine dyes [79–82], aggregation-induced emission (AIE) molecules [83–90], have attracted considerable attention.

The hydrophobic properties lead to the aggregation tendency of most PSs with \(\pi-\pi\) stacking structures, which heavily restricted the biomedical application of common PSs molecules. Thanks to the molecular assembly strategies and the discovery of various carriers, these insoluble molecules could be prepared as multifunctional nanotherapeutic nanoplatforms. For instance, Yan et al. constructed a novel type of nano-photosensitise via coordination reaction between metal-binding peptides, \(\text{Zn}^{2+}\) ions, and PS with multiple weak interactions using a multi-component self-assembly strategy (Fig. 2a) [91]. The obtained self-assembled nano-phototherapeutics exhibited unique advantages including uniform size distribution, high PS loading efficiency, robust colloidal stability in physiological solution, and prolonged blood circulation time in blood. Unsurprisingly, the self-assembled nano-phototherapeutic displayed quenched photoactivity due to aggregation of PS in physiological conditions. In contrast, their photoactivity was restored inside the tumor cells by responding to acidic pH and elevated glutathione (GSH) expression, which endowed the nanoplatform with tumor-specific and highly efficient PDT of cancer. The self-assembly strategy could efficiently solve the solubility problem of PSs molecules with conjugated structures by forming nanoparticles.

To acquire a favorable therapeutic effect, the nanoplatform should facilitate PSs drugs dwell in the tumor sites. Strategies that empower the nanoplatform with high accumulation and long-term retention at the tumor sites is worthy of being investigated. The solid tumor is of an acidic microenvironment due to glycolysis of the tumor cells. To take the advantage of the acidic tumor microenvironment, Pu et al. prepared a

![Fig. 1. The mechanisms of type-I and type-II PDT of cancer (R represents the substrate participating in the type-I photochemical reactions, ISC: intersystem crossing).](image-url)
Fig. 2. (a) A schematic diagram of preparing supramolecular metallo-nanodrugs by the coordination of peptide, PSs, and Zn$^{2+}$. Reprinted with permission [91]. Copyright 2018, American Chemical Society. (b) The transformation of peptide-porphyrin nanoparticles to peptide-porphyrin nanofibers activated by acid and the PDT application after intravenous injection. Reprinted with permission [92]. Copyright 2020, Wiley-VCH. (c) The formation of FHP photoactive nanostructure for NIRF imaging-guided PDT. Reprinted with permission [93]. Copyright 2020, Wiley-VCH. (d) The molecule structure of RDM-BDP for mitochondria targeting PDT. Reprinted with permission [94]. Copyright 2018, American Chemical Society.
kind of self-assembled acid-activated NPs with the peptide-porphyrin conjugates (Fig. 2b) [92]. The obtained peptide-porphyrin conjugate NPs could transform the spheroidal morphology into fibril nanostructure by forming molecular hydrogen bonds under the acidic tumor microenvironment, thereby improving $^{1}O_{2}$ generation efficiency and extending tumor accumulation of the nanoparticles. The peptide-porphyrin combined with morphology transformation function displayed excellent PDT performance. In this work, the acid-mediated morphology transformation strategy provided novel ways for elevating the accumulation of nanoparticles in the tumor regions.

To enhance tumor accumulation of the nanomaterials in the deep tissues, the active targeting ligand can be used. Recently, Yin et al. prepared a perylene monoimide-based nanocluster with a size of about 100 nm for fluorescence imaging-guided PDT of deep tumor (Fig. 2c) [93]. In their nanostructure system, the introduction of folate enabled the targeting capability to the tumor sites. The enzyme-triggered hydrolysis reduced the size of the nanostructure, eliciting the formation of ultrasmall NPs with a size of about 10 nm, which promised the successful penetration of NPs into the deep tumor sites. After the hydrolysis-induced disassembly, the fluorescence emission of the nanocluster was recovered, and their $^{1}O_{2}$ generation efficiency was also elevated for PDT. Featuring the enzyme-triggered size changes and the modified folate ligand in the structure, the nanoclusters achieved a deep tumor inhibition effect.

To boost the therapeutic index, more strategies elevating the tumor accumulation amount of theranostic materials should be explored. Studies have found that some materials exhibited the nature tumor accumulation tendency due to the presence of delocalized positive charge in their molecular structures. Utilizing the special characteristics, Peng et al. designed a phototheranostic material-based RDM-BDP molecule with the mitochondria-targeting property (Fig. 2d). The smart structure with delocalized positive charges endowed RDM-BDP with the structure-inherent targeting function for enhanced tumor-targeting potency. To further amplify the phototherapeutic outcome, they

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**Fig. 3.** (a) Synthetic procedure of Hf-DBP NMOM and the $^{1}O_{2}$ generation under light illumination. Reprinted with permission [98]. Copyright 2014, American Chemical Society. (b) The structure of DBC-UiO and the photosensitization under light irradiation. Reprinted with permission [99]. Copyright 2015, American Chemical Society. (c) The formation schematic of UNM. Reprinted with permission [102]. Copyright 2017, American Chemical Society. (d) Structure schematic of HUC-PEG. Reprinted with permission [103]. Copyright 2020, Elsevier. (e) A schematic diagram of preparing TPE-PHO nanostructure and the substitution activated phototherapy mechanism. Reprinted with permission [101]. Copyright 2020, American Chemical Society.
introduced the Förster resonance energy transfer (FRET) mechanism into molecular design. The FRET effect could optimize the photochemical property of PSs by expanding their absorption regions, thus increasing their photon utilization efficiency, and promoting \( \text{O}_2 \) generation. Assisted by the structure-inherent targeting function and FRET mechanism, the RDM-BDP-based nanoplatform exhibited notable photocytotoxicity to cancer cells and remarkable tumor inhibition effect in vivo upon 660 nm laser irradiation [94].

Apart from the above assembly strategies or direct dissolution by the organic solution to solve the hydrophobicity-induced aggregation problem of common PSs, the carriers have been intensively explored. The traditional polymer-carriers cannot completely avoid the aggregation state of hydrophobic molecules, which will partly counteract the photochemical property of phototherapeutic materials. Thanks to the development of materials chemistry, researchers demonstrated that nanoscale metal-organic frameworks (NMOFs) loaded with PSs molecules exhibited great advantage and potential. NMOFs as the vectors of porphyrin derivatives have been extensively employed for cancer therapy [95]. NMOFs with the periodic and ordered structures separate the porphyrin molecules from each other in the framework, thereby avoiding the aggregation caused quenching (ACQ) effect of PSs [96,97]. Moreover, NMOFs are rich in porous structures, which provide passable diffusion channels for the generated reactive species and enhance the antitumor performance of PDT. For instance, Lin et al. fabricated a porphyrin-based NMOF (i.e., DBP-Uio) with plate morphology (Fig. 3a). PDT by DBP-Uio exhibited outstanding antitumor performance toward the drug-resistant tumors under 640 nm light irradiation with a dose of 90 J cm\(^{-2}\). The therapeutic efficacy of DBP-Uio could be attributed to the enhanced \( \text{O}_2 \) generation efficiency induced by the heavy atom effect of Hf and ROS diffusion inside the MOF structures [98].

To improve the efficacy of PDT, Lin et al. further built a chlorin-based NMOF (DBC-Uio) with elevated photochemical reactivity and efficiently eliminated the colorectal tumors (Fig. 3b) [99]. The designed DBC-Uio possessed an elevated extinction coefficient in the lowest-energy Q band due to the successful reduction of porphyrin to chlorin [100,101], which endowed the amplified photon-harvesting capability for improved singlet oxygen generation efficiency. The ionic DBC-Uio with optimized photophysical properties exhibited superior PDT efficacy.

Similar to NMOF, porous organic polymers (POP) integrating porphyrin were also exploited for PDT of cancer. For instance, Xie et al. used the core-shell strategy to prepare a nanocomposite (UNM) by combining a metal-organic framework (UiO-66) with a porphyrin-based POP (Fig. 3c) [102]. Upon 450 nm light irradiation with a LED lamp, the porphyrin-based POP produced sufficient \( \text{O}_2 \) to kill the tumor cells. Moreover, they demonstrated that chlorin-based POP displayed amplified \( \text{O}_2 \) generation ability upon 671 nm laser irradiation (Fig. 3d) [103]. These porous materials-based phototheranostics remarkably inhibited tumor growth.

ACQ phenomenon of the conventional PSs is a major obstacle for hydrophobic nanomaterial-based PDT. In past years, AIE molecules-based nanomaterials have been investigated as alternative phototherapeutic. The AIE molecules can emit fluorescence in an aggregate state and generate cytotoxic \( \text{O}_2 \) under laser illumination [87]. Recently, Tang et al. designed a pyridinium-functionalized tetrathienylethylene AIE molecule (TPE-PHO) via host-guest interactions with water-soluble calixarene. The prepared complex structure formed the nanoplatform with well-defined nanoscale assembly (Fig. 3e) [101]. In normal conditions, the phototoxicity of TPE-PHO was quenched for avoiding dark toxicity during blood circulation. In contrast, the TPE-PHO molecules were competitively replaced by 4,4′-benzidine dihydrochloride from the cavity of calixarene and the photoactivity of TPE-PHO molecules was restored, causing the generation of \( \text{O}_2 \) upon white light illumination for PDT at the tumor sites. Moreover, the supramolecular assembly strategy in this work not only facilitates the nanometer form of PSs molecules but also provides the feasible methods for the application of materials with high dark cytotoxicity. Aiming at the hydrophobility problem of phototherapeutic agents, the AIE molecules exhibited great advantage by converting enemies into friends. However, the short absorption wavelength and high dark cytotoxicity presented a formidable challenge to their further in vivo biomedical application.

For oxygen-dependent type II PDT, the inherent tumor hypoxia limitation must be faced and conquered for superior phototherapeutic efficacy. To overcome the hypoxia drawbacks, four pathways have been studied in detail: (1) the direct method is supplying oxygen to the tumor sites utilizing the perfuorocarbon molecules or \( \text{MnO}_2 \); (2) the employment of hypoxia-activated drugs (such as tirapazamine) can counteract the hypoxia drawback; (3) the way using the decomposition of \( \text{H}_2\text{O}_2 \) for supplying continuous \( \text{O}_2 \) mediated by some metal-induced catalytic re-actions have demonstrated their effectiveness in phototherapy of hypoxic tumors; (4) the introduction of inhibitors towards hypoxia inducing factors can efficiently relieve the hypoxia stress.

To demonstrate this, Chen et al. recently developed an iron-porphyrin MOF-based oxygen-supplying nanoplatform for PDT of cancer. Upon NIR laser irradiation, the nanoparticles were loaded with AuNPs, the latter can catalyze the glucose to generate \( \text{H}_2\text{O}_2 \) via glucose oxidase-like catalytic activity for starvation therapy. Subsequently, the produced \( \text{H}_2\text{O}_2 \) was decomposed to generate oxygen via photocatalysis reaction of Fe-porphyrin constructed MOF (Fig. 4a) [104]. Meanwhile, the porphyrin-based PSs could facilitate the generation of cytotoxic \( \text{O}_2 \) excited by the visible light emitted from the UCNPs. This novel system not only provided a continuous source of oxygen for PDT but also used the up-conversion mechanism to convert near-infrared light (980 nm) to visible light, which can solve the tissue-penetration problem of short-wavelength light.

Similarly, Liu et al. used the oxidation etching strategy to prepare holey palladium nanosheets (namely H-Pd NSs) for overcoming the hypoxic tumor microenvironment and improving PDT performance (Fig. 4b) [105]. The palladium nanosheet without etching treatment was employed as a PTT agent. However, the etching decoration endowed the palladium nanosheets with the functionality of catalyzing the decomposition of hydrogen peroxide to generate \( \text{O}_2 \) (Fig. 4c). The generated \( \text{O}_2 \) was transferred into \( \text{O}_2 \) photocatalyzed by H–Pd NSs upon 808 nm laser irradiation. This oxygen-supplying advantage endowed the nanosystems with similar photoinduced killing efficacy to 4T1 cells under hypoxia and normoxic conditions (Fig. 4d). The combinatory effect of photothermal and PDT, the H–Pd NSs displayed satisfying therapeutic efficacy in vitro and in vivo, which have achieved a tumor inhibition rate of 99.7%.

In addition to the delicate nanoplatform to elevate the oxygen concentration at tumor sites, the direct inhibition of hypoxia-related cascade reactions for enhanced therapeutic effect has been investigated. To relieve the side effects of PDT-induced hypoxic tumor microenvironment, Pu et al. further designed a smart nanoplatform integrating a semi-conducting polymer-based PS and an antagonist of carbonic anhydrase IX (CA-IX) (Fig. 5a) [106]. The CA-IX inhibitor suppressed proliferation and metastasis of the tumor cells by dysregulating the balance of both intracellular and extracellular pH (Fig. 5b). The integration of CA-IX inhibitor enhanced the targeting ability of the whole platform to CA-IX positive breast cancer cells. Upon 680 nm laser irradiation, the semiconducting polymer PS could efficiently produce \( \text{O}_2 \) for PDT. By combining the synergistic effect of PDT and anti-tumor metastasis mechanism, a remarkable therapeutic effect could be obtained upon phototherapeutic.

The synergistic therapy could integrate the collective merits of employed multiple treatments and exhibited enhanced therapeutic effect. The introduction of hypoxia-responsive drug systems can efficiently inhibit the tumor cells in hypoxic conditions. For instance, Pu et al. conjugated a chemotherapy drug (IPM-Br) to the backbone of semi-conducting polymer nanoparticles (SPNs) via a hypoxia-responsive linker (Fig. 5c) [107]. Upon 808 nm light illumination, the SPNs consumed \( \text{O}_2 \) to generate \( \text{O}_2 \) for PDT. The PDT intensified hypoxia.
would cleave the hypoxia-sensitive linker and release the IPM-Br drug to crosslink the DNA macromolecules, thereby leading to cellular apoptosis (Fig. 5d). The synergistic PDT and chemotherapy exhibited 4.3-times higher tumor inhibition effectiveness than its chemotherapy process.

2.2. Nanoparticles for Type-I PDT

Just as a Chinese proverb says: “Better to divert rather than blocking and better to change rather than diverting”. This motto guides researchers that the hypoxia targeting therapeutic modalities may be better than the direct oxygen-supplying methods, however, the designing of oxygen-independence nanomedicine platforms may possess the optimal therapeutic effect. Thanks to the repaid development of material synthetic chemistry tailed to biomedical application, researchers demonstrated that some photoactivatable nanomaterials could perform type-I PDT in an oxygen-independent manner. These nanomaterials are of great potentials for PDT of hypoxic solid tumors. For example, Huang et al. prepared a novel nanostructure based on the self-assembled phthalocyanine molecules (NanoPcA) (Fig. 6a) [108]. NanoPcA with a unique structure design can avoid the ACQ phenomenon and perform PDT with the type-I mechanism. Compared with organic nanomaterials-based PSs, metal-based PSs also exhibit remarkable photocytotoxicity in the tumor cells. Recently, Ruiz et al. constructed a PS complex by conjugating the cyclometallated Ir(III) to the coumarin molecules (Ir(III)–COUPY) (Fig. 6b) [109]. Ir(III)–COUPY was effectively endocytosed by HeLa cells and exhibited good biocompatibility in dark conditions. Upon blue or green light irradiation, Ir(III)–COUPY induced a similar photocytotoxicity effect to HeLa cells under normoxic or hypoxic conditions. Mechanism studies indicated that Ir (III)–COUPY caused cell death by generating superoxide

Fig. 4. (a) A schematic diagram of the oxygen self-supplying UMOFs@Au NPs for enhanced PDT efficacy in vivo. Reprinted with permission [104]. Copyright 2020, American Chemical Society. (b) The preparation process of holey Pd NSs (H–Pd NSs). Reprinted with permission [105]. Copyright 2020, American Chemical Society. (c) The detection of the decomposition process of H2O2 into O2 by electron spin resonance spectra. Reprinted with permission [105]. Copyright 2020, American Chemical Society. (d) Viability of 4T1 cells treated with different conditions. Reprinted with permission [105]. Copyright 2020, American Chemical Society.

Fig. 5. (a) The preparation of OPNi and (b) the mechanism of OPNi induced synergistic phototherapy. Reprinted with permission [106]. Copyright 2019, Wiley-VCH. (c) The preparation of SPNs photoactive nanomaterials for combined PDT/chemotherapy application. Reprinted with permission [107]. Copyright 2019, Wiley-VCH. (d) The release mechanism of IPM-Br for chemotherapy.
radical anion after visible light irradiation by performing the type I PDT process under hypoxia conditions. The generated superoxide radical anion could be confirmed by the emitting green fluorescence of dihydrodorhamine 123 (a ROS indicator) in a cell-free-media and the increased luminescence of luminol in the test of a cell-based assay. It is presumed that the Ir(III)–COUPY conjugates selectively produced superoxide radical anion via the possible mechanism of suppressing intramolecular charge recombination. This work excavated the increased luminescence of luminol in the test of a cell-based assay. It is understood process under hypoxia conditions. The generated superoxide radical anion under light irradiation in a hypoxic environment through the type-I peroxide radical anion via the possible mechanism of suppressing the Ir(III) complex and coomarin by exploring their cytotoxic \( \cdot \text{O}_2 \) generation mechanism. The special photochemical behavior could be used to treat hypoxia tumor cells. However, the shorter absorption wavelength of Ir(III)–COUPY conjugates limited their further biomedical application in vivo.

The above PS nanoplatorm designing all employed ROS-active motifs as the main photoresponsive group, the ROS-inert motifs could also be favorable photoactivatable agents by forming a special structure. For instance, Deng et al. constructed two kinds of COF nanomaterials (called COF-808 and COF-909) with the photo-inactivatable molecular monomer L-3C (Fig. 6c) and L-3N (Fig. 6d) as building units, respectively [110]. The cross-linked COF-808 and COF-909 generated a large number of superoxide radical anions under 630 nm light irradiation. The generated superoxide radical anions subsequently reacted with the \( \cdot \text{H}_2\text{O} \) molecule to form the hydroxyl species. PDT study in vitro showed that COF-909 killed more than 80% of the tumor cells at a concentration as low as 50 \( \mu \text{g/mL} \). The antitumor studies demonstrated that COF-909 inhibited over 90% of tumor growth compared with that of the untreated control group, indicating the promising potential of COF-909 for type-I PDT. However, the clinical translation of these COF structures was restricted by their poor physiological stability.

Besides the traditional nanoparticle mode-based phototherapy, the direct small organic molecules have also been explored as efficient PDT agents. For instance, Peng et al. developed a superoxide radical anion generator (ENBS-B) based on the benzophenothiazine compounds, which could be activated by NIR light (Fig. 6e) [111]. Upon 660 nm laser irradiation, the obtained ENBS-B could produce sufficient \( \cdot \text{O}_2 \) under light irradiation in a hypoxic environment through the type-I photochemical reaction for PDT (2% \( \text{O}_2 \)). The intracellular uptake amount of ENBS-B was about 87-times higher than the control groups due to the targeting effect of biotin ligands. The generated \( \cdot \text{O}_2 \) species can destroy the intracellular lysosomes and induced tumor cell apoptosis while participating in SOD-mediated reactions to form \( \cdot \text{H}_2\text{O}_2 \) and highly toxic \( \cdot \text{OH} \). ENBS-B specifically targeted the tumor tissues and efficiently inhibited tumor growth by low-dose laser irradiation when administrated by intravenous injection.

To improve the photochemical property of the nanogenerators, Peng et al. further developed another nanophototherapeutic generating superoxide radical anion \( (\cdot \text{O}_2) \) for type-I PDT (Fig. 6f) [12]. The photon-absorption capacity of ENBOS in the NIR region was significantly amplified via the FRET mechanism. ENBOS could be activated by laser irradiation at low photodensity. Remarkably, ENBOS displayed excellent “structure inherent targeting” ability. The signal-to-noise ratio of the tumor and surrounding tissues reach 25.2 when administrated by intravenous injection, indicating the excellent tumor targeting ability of ENBOS. Furthermore, ENBOS effectively eliminated the hypoxic tumor owing to the structural intrinsic localization function, strong photon absorption capacity, and oxygen-independence characteristic.

To alleviate the hypoxia burden of tumor cells and acquire more right to use the existing oxygen in the tumor cell, Peng et al. further designed a superoxide radical anion \( (\cdot \text{O}_2) \) generator (called SORgenTAM), which was able to reverse the hypoxia-induced therapeutic resistance by acting on mitochondrial respiration (Fig. 6g) [112]. Specifically, the tamoxifen molecule in the nanotherapeutic structure affected the normal process of mitochondrial respiration, reduced the hypoxic pressure of tumor cells by preventing \( \text{O}_2 \) consumption in the tumor cells, and simultaneously down-regulated HIF-1 expression, thereby preventing the tumor cells from becoming hypoxic. The photosensitization studies demonstrated that the superoxide radical anion could transform into hydroxyl radical via a cascade of catalytic reactions. Moreover, the antitumor study suggested that SORgenTAM inhibited self-repair of the tumor cells by activating the AMPK metabolism signaling pathway and enhanced the lethal effect to tumor cells. This study suggested that the SORgenTAM phototherapeutic platform could be tailored for treating deep-seated hypoxia tumors.

### 2.3. Nanoparticles for combined Type I and II PDT

Thanks to the richness and diversity of existing materials and the advance of synthetic chemistry, researchers have found that some phototherapeutic nanomaterials could simultaneously perform the \( \text{O}_2 \)-dependent and \( \text{O}_2 \)-independent PDT. For instance, Lin et al. developed a Ti-based NMOF (Ti-TBP) for combining Type-I and Type-II PDT process (Fig. 7a) [113]. Upon 650 nm laser irradiation, the Ti-TBP NMOF produced \( \cdot \text{O}_2 \) and transferred electrons from the excited TBP* species to Ti\(^{4+}\)-based secondary building units, generating TBP* ligands and Ti\(^{4+}\) centers, thereby promoting the generation of superoxide anions hydrogen peroxide and hydroxyl radicals. Thus the bifunctional Ti-TBP NMOF nanophototherapeutics-mediated PDT exhibited a superior anticancer effect by generating four different types of ROS and efficiently inhibited the solid tumor growth (~98%) under 650 nm light irradiation.

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**Fig. 6.** (a) Molecular structure of PcA. (b) Molecular structure of the Ir(III)–COUPY conjugate. (c) The molecular structure of L-3C for constructing COF-809. (d) The molecular structure of ENBS-B. (e) Molecular structure of ENBS-B. (f) Molecular structure of ENBS-B. (g) Molecular structure of SORgenTAM.
To improve the antitumor performance of the nano-phototherapeutic, Lin et al. further prepared a Zr-TBB NMOF by utilizing bacteriochlorophyll as the connecting unit and zirconium as the metal node (Fig. 7b) [98]. Under 740 nm laser irradiation, Zr-TBB NMOF triggered superoxide anions, hydrogen peroxide, hydrogen peroxide, and singlet oxygen generation through the type-I and type-II mechanisms, leading to a favorable PDT effect. The four kinds of reactive oxygen species were verified by aminophenyl fluorescein assay and electron paramagnetic resonance measurement. In vivo studies showed that the Zr-TBB NMOF was of excellent antitumor efficacy in the mouse models of breast and colorectal cancer, with cure rates of 40% and 60%, respectively.

In addition to NMOF, covalent organic porous nanostructures gained extensive attention in recent years due to their framework structures and tunable photochemical properties. For instance, Tian et al. prepared a heterogeneous covalent organic nanosheet (CON) with a donor-acceptor structure (Fig. 7c) [114]. Upon 635 nm laser irradiation, the porphyrin PS molecules in the nanosheet generated O$_2^*$ for type-II PDT. Besides, the heterostructure of CONs provided efficient charge separating efficiency to obtain long-lived electrons and holes. Electrons reduced O$_2$ to form O$_2^*$, and holes oxidized water to produce toxic HO$_2$ to exert the type-I PDT process. The type-I PDT function significantly relieved the hypoxia-induced resistance to type-II PDT. The CONs displayed significant tumor suppression effect by combining type-I, type-II PDT and PTT modality when administrated by intravenous injection. This work explored the photochemical mechanism of porphyrin-based CONs and expanded the biomedical application of indissoluble covalent organic nanomaterials.

3. Reactive nitrogen species-mediated tumor suppression

Reactive nitrogen species (RNS) contains nitric oxide (NO) and a wide range of its metabolites, such as peroxynitrite anion (ONOO$^-$) [23, 115–118]. As a biological regulatory molecule in the human body, NO plays crucial functions in various physiological and pathological processes [7,8,119]. Therefore, NO is closely related to the occurrence and progression of many diseases including cancer [117]. Similar to reactive oxygen species, NO is a double-edged sword, and its biological effects are concentration-dependent [120]. For instance, NO can promote angiogenesis of the solid tumor at a concentration below 0.2 μmol. In contrast, NO caused poisoning effect in the blood at a concentration of NO higher than 1 μmol [121]. When NO is maintained at a certain concentration (>1 μmol), it could inhibit tumor growth and eventually regressed tumor [122–124]. The main antitumor mechanisms of NO are as follows: (1) affecting the energy metabolism of cancer cells, which resulted in tumor suppression; (2) combining with reactive oxygen radicals (such as superoxide radical anion) in the cell to damage the skeleton structure of DNA by generating nitrogen/oxygen-based free radicals; (3) promoting macrophage activation to fight tumors, and inhibiting tumor metastasis by down-regulating the platelet aggregation; (4) inducing and activating the expression of tumor suppressor gene p53 for causing tumor cell apoptosis; and (5) reducing P-glycoprotein (P-gp) expression levels against drug resistance.

As a typical radical, NO is the source of all RNS in the biological systems [125]. NO possess high reactive activity in the presence of other radicals and exhibit oxidation or nitration potential to the local biomolecules. Therefore, precise cancer therapy can be achieved by controlling NO release at the lesion locations. Generally, there are four bonding strategies including R-O-NO, R-S-NO, R-N-NO (R represents alkyl group), and mental-NO for NO delivery [126]. The four bonding or coordination-based NO supplying nanotherapeutic platforms can achieve light irradiation-triggered NO release. The external light irradiation-induced photothermal effect can also induce a controlled release of NO. However, despite various nanomaterials that have been designed for NO-delivery, it remains a challenge to precisely deliver NO to the tumor sites in a controlled release profile.

NO-based monotherapy application is relatively rare due to the concentration-dependent therapeutic effects of NO. However, NO-based combination therapy has been extensively studied. Current NO-based treatment research mainly focuses on four pathways, including high concentration-induced cytotoxicity [127], as a chemotherapetry sensitizer [128], as radiation therapy sensitizer [129], or as phototherapy sensitizer [120,122]. The ever-increasing antitumor research about the combination of NO therapy with various therapeutic strategies demonstrated the great significance of summarizing the related research works. In this section, the mechanism and advantages of synergistic therapy between NO therapy and other therapeutic modalities will be discussed for discovering innovative methods to cure...
cancer.

The drug delivery rate for tumor clinical chemotherapy is low due to the limitation of physiological barriers, which lead to poor therapeutic outcomes and tumor metastasis. Studies have shown that a low dose of NO can amplify the permeability and retention effects (EPR) of the nanoparticles. Moreover, NO has been extensively employed for sensitizing the tumor cells to chemotherapeutics, such as cisplatin, paclitaxel (PTX), and doxorubicin (DOX). For example, Kim et al. designed a polymer micelle formed by nitrobenzene and platinum prodrugs by conjugating with nitrobenzene molecules at both ends of the platinum prodrugs in the axial direction (Fig. 8a) [128]. The strong hydrophobic property of nitrobenzene improved the loading efficiency of Pt(IV) prodrugs when utilizing polymeric micelles as the carrier. The photo-cleavage of the nitro compound transformed the hydrophobic nitrobenzene into a hydrophilic phenol structure, which further increased the release efficiency of platinum (IV) prodrugs. This nanomedicine platform utilizing NO donors for tuning NO release and boosting chemotherapy exhibited effective anticancer effects. However, the tissue penetration of the ultraviolet light is quite limited, which restricted the biomedical application of nitro benzene-based NO donors in vivo.

Similarly, NO can be used to amplify the chemosensitivity of PTX by the bonding method of R-S-NO. Li et al. constructed an antitumor nanoplatform (IPH-NO NPs) by utilizing the NO donor bonded human serum albumin as a drug carrier to support the photosensitizer (IR780) and PTX (Fig. 8b) [130]. IPH-NO NPs slowly released NO in the presence of glutathione (GSH), thus leading to elevated permeability of NPs to the tumor blood vessels and improved drug accumulation at the tumor sites. Moreover, the NIR light irradiation induced the rapid release of NO, which directly killed tumor cells, and inhibiting tumor metastasis. This nanoplatform exhibited improved tumor suppression effect and anti-metastasis effect by the synergistic therapy of NO therapy, photothermal therapy, and chemotherapy.

Also, NO can be used to enlarge the chemosensitivity of DOX by codelivery of NO donor (metal-NO coordination complex) and DOX drugs in one nanoplatform. Shen et al. designed a nanoplatform and proved the positive effect of NO on DOX-based chemotherapy (Fig. 8c) [131]. In this nanoplatform, chitosan was bonded on the surface of upconversion nanoparticles (UCNPs) and then loading DOX and photoactivated NO donor (Ruxin black salt, RBS) to obtain the nanospheres (UCNPs(DOX)@CS-RBS) (Fig. 8c). The UCNPs in the nanospheres effectively converted absorbed NIR photons into visible light, triggering the release of NO. The low pH in the lysosome promoted the release of DOX. They demonstrated that a small dose of NO acted as a P-gp-mediated therapeutic agent against multidrug resistance (MDR), leading to the increased concentration of DOX in the cell. In the meantime, the high dose of NO and DOX synergistically killed cancer cells. The combination of DOX and high doses of NO exhibited an improved tumor proliferation suppression effect. It has been defined that the multidrug resistance had severely limited the therapeutic effect of chemotherapeutic. Therefore, the successful introduction of NO should be of great practical significance.

In addition to combination with chemotherapy, NO has also been used to optimize the therapeutic effect of PDT on hypoxia tumors. For instance, Dong and Huang et al. synthesized a nano-PS based on the diketopyrrolopyrrole nucleus (DPP-NF) (Fig. 9a) [122]. They introduced a light-responsive NO donor (4-nitropyrole) and pH-sensitive group (dimethylaminophenyl) into the structure of DPP-NF NPs (Fig. 9b). The DPP-NF NPs could be activated under weakly acidic conditions of lysosomes (pH = 4.5–5.0) by forming protonated structure to generate ROS for PDT and enable the photothermal conversion function for PTT under laser irradiation. Moreover, upon 660 nm light illumination, the released NO was able to trigger tumor cell death by inducing DNA damage. The NO therapy could overcome the low efficiency of PDT modality to hypoxic tumors. Both in vitro and in vivo antitumor effects demonstrated that DPP-NF NPs are promising phototherapeutic nanoplatform.

It is well known that PDT could suppress tumor growth by causing irreversible oxidative damages to the lipids, amino acids, and proteins. However, some recent evidence have shown that the concentration of NO synthase (iNOS) was upregulated during the PDT process. The iNOS or NO donors generated NO could suppress the PDT-induced...

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**Fig. 8.** (a) The structure schematic of the photoactive Pt(IV) prodrug. Reprinted with permission [128]. Copyright 2018, Royal Society of Chemistry. (b) Schematic illustration of the IPH-NO for enhanced antitumor therapy. Reprinted with permission [130]. Copyright 2019, Royal Society of Chemistry. (c) The release mechanism of UCNPs(DOX)@CS-RBS nanomaterials activated by light illumination and the acid. Reprinted with permission [131]. Copyright 2017, Royal Society of Chemistry.
photokilling effect to cancer cells via weakening the lipid peroxidation effect [121]. So the nanomaterials designing about combination of NO therapy and PDT need to consider more experimental variables to fully understand the function mechanism, thereby obtaining optimal PDT efficacy.

NO has also been used as an efficient photothermal therapy (PTT) sensitizing agent. Uneven heat distribution often leads to poor accuracy of treatment and side effects of burning normal tissues during the PTT process. Therefore, mild PTT is a considerable alternative and could be explored for antitumor research. Zhao and Gu constructed a nanocomposite based on the bismuth sulfide (Bi$_2$S$_3$) NPs and a NO donor (bis-N-nitroso compound, BNN). Under 808 nm light illumination, BNN released NO facilitated the even distribution of nanocomposite in solid tumors for mild PTT. Moreover, the exploration result indicated that the self-repair function of tumor cells was inhibited, thus achieving an excellent PTT effect [120]. The results suggested that NO could cooperate with PTT modality to suppress the growth of tumor cells. This work provided novel insight for avoiding the side effects of hypothermia therapy.

Radiotherapy is commonly used for cancer treatment in the clinic. However, the therapeutic efficiency of radiotherapy is generally unsatisfactory, due to its low specificity, limited drug resistance, and side effects caused by high doses of radiation. Researchers have found that NO can effectively sensitizer radiotherapy. For instance, Wang et al.
recently synthesized a controlled therapeutic platform by using bovine serum albumin (BSA) as a carrier of Ag2S quantum dots, NO donors (TBN), and ovalbumin (OVA) (Fig. 9c) [129]. The photothermal effect of the nanoplatform upon 808 nm light irradiation precisely triggered the release of NO, which optimized the radiotherapy effects of heavy metal-Ag and immunotherapy effect induced by OVA. The antitumor study in vivo showed that the survival rate of tumor-bearing mice reached 100% after the treatment of this multifunctional nanoplatform. Compared with the control group, the tumor volume of the NO-combining treatment group was the smallest, indicating the radiotherapy sensitization effect of NO.

In addition to the adjuvant function to other therapy modalities, NO can directly improve tumor treatment efficacy at sufficient concentration. For example, Zhang and Feng have doped NO donor (sodium nitroprusside, SNP) into the mesoporous Prussian blue (m-PB) NPs to obtain m-PB-NO NPs, and then loading docetaxel (DTX) to construct the DTX@m-PB-NO platform (Fig. 9d) [127]. By adjusting the intensity and time of NIR light irradiation (808 nm), NO was released in a controlled manner. This system combined NO therapy, PTT, and chemotherapy to effectively inhibit tumor growth and metastasis. Similarly, Fan et al. used F127 to encapsulate semiconducting polymer and s-nitrosothiol group (SNAP) to form a theranostic platform (Fig. 9e) [132]. Upon 808 nm laser irradiation, the encapsulated semiconducting polymer converted light to heat energy for PTT treatment. The generated heat decomposed s-nitrosothiol to produce NO for cooperative treatment of NO and PTT. The combined treatment strategy inhibited the tumor growth up to a 77% percentage. The synergistic therapy by combining NO and PTT modality has great potential due to the favorable antitumor efficacy.

Along with the therapeutic effect of NO, Chan et al. recently synthesized a NO donor (photoNOD) for photacoastic (PA) imaging by using aza-BODIPY as the backbone [133]. This NIR light provided enough energy to break the N-nitroso bonds, which helped BODIPY molecules to generate a strong PA signal under excitation, and the photophysical properties were also affected by the change of chemical structure. This NIR light-controlled property enabled the molecule to be an ideal ratiometric probe. After NIR irradiation, photoNOD-1 and photoNOD-2 released NO and generated aza-BODIPY (rNOD-1 or rNOD-2) (Fig. 9f). The conversion from electron-deficient N-nitroso bond to electron-rich N–H bond caused the absorbance redshift of the molecule. The irradiation of photoNOD and rNOD with two wavelengths of light led to the generation of two corresponding PA signals, and the generated ratio signals could detect the ratio of photoNOD and rNOD and non-invasively monitor the release of NO in living animal tissues. Compared with a fluorescence-based imaging modality, PA imaging of photoNOD offered an non-invasive and high-resolution imaging strategy to detect the NO release process without additional contrast agents (Fig. 9g). This work demonstrated that the NO-based reactive species can not only optimize the therapeutic effect of common cancer therapy but also play a part in the imaging field. The potential of NO for PA imaging will be beneficial for performing tunable and tumor-specific NO delivery with the NO donors-containing nanomaterials.

4. Carbon-containing reactive species for cancer therapy

Carbon-containing reactive species are kinds of carbon-centric derivatives. Herein, we mainly overviewed the advances of CO and alkyl radicals-based carbon-containing reactive species for cancer therapy [4].

4.1. CO-performed cancer therapy

It is well known that the binding affinity of CO for hemoglobin (Hb) is 220-times higher than that of oxygen [134,135]. Therefore, CO can disrupt the oxygen-carrying capacity of Hb in the blood, thus causing poisoning effects. CO can also inhibit the function of mitochondria and induce autophagy of the cells. However, CO is a metabolite of endogenous heme. As an endogenous signaling molecule, CO has a variety of biomedical functions, such as regulating blood pressure, reducing inflammation, sensitizing the tumor cells to chemotherapy, and dysfunction-lining mitochondrial aerobic breathing energy supply [134]. Based on these functionalities, the study of CO as a therapeutic agent has received widespread attention [136,137]. However, to avoid the toxicity of CO, it is essential to deliver CO specifically to the tumor sites and release the payload on demand. Currently, various nanocarriers (such as polymers, ferric oxide, silica nanoparticles, Au nanocages (AuNCs), etc.) have been developed to support the CO donors (Ru carbonyl, Fe carbonyl, and Mn carbonyl). In this section, the potential application and mechanism of CO-based multifunctional nanomaterials for anti-tumor therapy are summarized.

Based on the special functionality, Zhang et al. used AuNCs to support CO donor-iron pentacarbonyl (Fe(CO)5) under anaerobic conditions, avoiding leakage and oxidation during the blood circulation. The iron pentacarbonyl on the surface of Au nanocages could turn into iron oxide under aerobic conditions (Fig. 10a) [138]. Upon 808 nm laser irradiation, Au converted light to heat energy and promoted the decomposition of iron pentacarbonyl into CO and iron. The generated CO interfered with the respiration of mitochondrial, induced autophagy, and damaged the lysosome in cancer cells. Besides, the produced iron and the iron oxide degraded under the weak acidity of the lysosome and underwent Fenton reaction for producing cytotoxic hydroxyl radicals, which further destructed the autolysosomes, and improved the anti-cancer performance. This study combined the CO therapy with hydroxyl radicals-induced oxidative damage demonstrated the potential of CO species for antitumor therapy.

Apart from utilizing the influence of CO on the function of mitochondrial for tumor regressing, CO could also be employed to enhance the performance of chemotherapy. To overcome the resistance of chemotherapy, small molecule chemosensitizers are used to make tumor cells more sensitive to chemotherapeutics. Studies indicated that CO could act as a chemotherapeutic sensitizer and also exert the function of anti-inflammatory. For instance, Zhang et al. designed a multifunctional CO nanogenerator based on partially oxidized tin disulfide nanosheets (POS NSs) to inhibit tumor growth and suppress inflammatory response (Fig. 10b) [139]. POS NSs are modified with targeting ligand-functionalized polymers (PEG-cRGD) for improving their tumor specificity. POS NSs can efficiently load DOX to prepare PPOSd due to the large surface area and volume of NS. The resultant PPOSd could catalyze CO2 to generate CO upon 561 nm laser irradiation, and sensitize the tumor cells to DOX treatment. Moreover, the POS NS in PPOSd can induce hyperthermia effect upon laser irradiation for photothermal ablation of the tumor. The produced CO could reduce the inflammatory response caused by PTT. This study effectively explored the anti-inflammatory function of CO and uncovered its action mechanism.

Similarly, Shen et al. developed the MCM@PEG–CO–DOX NPs by firstly oxidizing the ferrocene compound to obtain carbonized magnetic NPs (MC), and then modifying the surface of MC with iron-based MOF (MIL-100) and utilizing MIL-100 as a carrier for supporting the CO donor-Mn(CO)5Br and DOX (Fig. 10e) [140]. Upon 808 nm laser irradiation, MC produced photothermal effects for triggering CO and DOX release. The combination of CO with chemotherapy and PTT elicited an excellent tumor suppression effect. The study demonstrated the potential application of CO for improving chemo- and photothermal therapy of cancer.

CO as the anti-inflammatory agent could precisely avoid the reverse effects (i.e., inflammation) of PTT. To verify the feasibility, Zhang et al. used defective tungsten oxide (WO3) nanosheets (DW NSs) to support carbon dioxide donor-bicarbonate (BC) and then modified their surface with polyethylene glycol (PEG) to make P@DW/BC nanosheets [141]. Upon 808 nm laser irradiation, the DW in P@DW/BC exerted PTT function and converted carbon dioxide (CO2) into CO via photocatalysis process. Combining the photothermal and anti-inflammation effects of
CO, the prepared nanosheets have exhibited excellent antitumor effects with low adverse side effects.

Moreover, recent studies have indicated that CO could accelerate the oxygen consumption of mitochondrial, promoting a hypoxic environment for activating the hypoxia-activatable prodrug. For example, Yin et al. prepared a theranostic nanoplatform by utilizing m-PB NPs as carriers to support the tirapazamine (TPZ) and iron pentacarbonyl (Fe(CO)$_5$, CO donor) (Fig. 10 d) [137]. The modification of polyallylamine hydrochloride (PAH) and polyacrylic acid (PAA) on the surface prolonged the blood circulation of the nanoplatform and promoted its accumulation in tumor sites. After intravenous injection, the photothermal effect of PB under 808 nm light irradiation induced the cleavage of the coordination bonds in Fe(CO)$_5$, thus releasing CO. Subsequently, CO accelerated the aerobic respiration of mitochondria and increased the hypoxic extent of tumor cells. The hypoxic environment further activated the bioreductive chemotherapeutic TPZ, which can generate transient oxidative free radicals during the metabolism of reductase to combine with DNA, leading to DNA damage and subsequent cell death. CO-mediated activation of TPZ significantly enhanced the antitumor efficacy of TPZ-mediated chemotherapy. This pioneering study ingeniously explored the influence of CO on the oxygen consumption of mitochondrial and broadened its biomedical application.

Studies have found that the mechanism of multidrug resistance is related to the P-glycoprotein (P-gp)-mediated drug efflux of the cancer cell, and the function of P-gp highly depends on the binding of ATP [142]. There are various strategies to overcome P-gp-involved MDR: (1) using P-gp inhibitors to reduce nicotinamide adenine dinucleotide (NADPH) and adenosine triphosphate (ATP) to down-regulate P-gp expression. (2) designing anticancer drugs targeting mitochondria and using interfering RNA to silence P-gp expression. Studies have found that CO could reverse the drug resistance of chemotherapy by inactivating mitochondria and suppressing the function of P-gp. For example, Yin et al. reported the strategy of using CO for overcoming the MDR process [143]. They used mesoporous Prussian blue nanoparticles (PB NPs) as a carrier to support iron pentacarbonyl (Fe(CO)$_5$) and DOX. Upon 808 nm laser irradiation, the Fe(CO)$_5$ released CO via photothermal breakdown, resulting in mitochondrial metabolic failure. This process prevented the synthesis of ATP and inhibited ATP-dependent drug outflow, thus reversing drug resistance of MCF-7/ADR tumors. Besides, CO-mediated mitochondrial failure activated the apoptotic protein caspase 3, which further lead to the tumor cell apoptosis.

4.2. Alkyl radical-mediated cancer inhibition

ROS can inhibit the proliferation of the tumor cells by inducing oxidative damage to biomolecules (e.g., lipids, proteins, and DNA) and subcellular organelles (i.e., mitochondria). However, the generation of ROS is highly dependent on the local concentration of oxygen. The intratumoral production of ROS is insufficient since most solid tumors are hypoxic, especially the area away from the blood vessels. Therefore, the development of O$_2$-independent free radical therapy strategies may improve the therapeutic effect by overcoming the hypoxic condition. A feasible method to solve the above problems is delivering free radical-
generating precursors to the tumor sites that can be activated via external stimulation. A recent study showed that 2,2′-azobis[2-(2-imidazolin-2-yl)propane]dihydrochloride (AIPH) as a conventional polymerization initiator can produce alkyl radicals under heat or irradiation activation. The generated free radicals can directly oxidize intracellular components in hypoxia conditions or react with O2 to produce O2·-containing free radicals, such as alkoxy and peroxy groups in a normoxia microenvironment [144]. AIPH may become an effective biomedical candidate for treating hypoxia tumors.

To achieve tumor-targeted delivery of AIPH, Xia et al. designed a novel nanotherapeutic by integrating AuNC, phase change material (PCM), and AIPH into one single nanoplatform (Fig. 11a) [6]. They used AuNC to load AIPH and wrapped the surface of AuNC with a PCM outer layer. Upon 808 nm laser irradiation, AuNC induced hyperthermia effect for inducing phase transition of PCM and then triggering on-demand AIPH release. As an azo compound, AIPH decomposed rapidly under the stimulation of photothermal-induced heat to generate alkyl radicals. These free radicals directly oxidized cellular components or reacted with oxygen to generate secondary cytotoxic free radicals, including alkoxy and peroxy groups. Even under hypoxic conditions, photothermal-induced heat still induce AIPH to produce free radicals to increase intracellular lipid hydroperoxide and ultimately lead to oxidation-mediated damage and cell apoptosis. In vitro, experimental results showed that the Au-PCM-AIPH in the hypoxia group and the normoxia group showed an approximate inhibition rate on A549 cells at 61.1% and 62.3%, respectively. The experimental data indicated that the tumor suppression effect of Au-PCM-AIPH was independent of the oxygen level. This novel AIPH-based therapeutic strategy expanded the range of reactive species and extended the biomedicine application of reactive species-based nanomaterials.

Lin et al. further prepared a kind of NPs by combining BSA-coated magnetic Fe5C2, AIPH, and PCM (Fig. 11b) [145]. This nanoplatform exhibited a remarkable cancer cell inhibition effect by producing toxic free radicals induced by the photothermal effect. The multifunctional therapeutic nanoplatform also combined magnetic targeting function to remotely control drug delivery. This study combining the alkyl radicals-mediated cancer therapy and the magnetic resonance imaging (MRI) performance could realize the MRI-guided antitumor therapy.

Furthermore, Lin et al. used self-assembly strategies to develop a therapeutic platform based on CuFeSe2 NPs, BSA, and AIPH (Fig. 11c) [146]. Owing to the strong absorption in the NIR-I and NIR-II spectral regions, the CuFeSe2 NPs could activate the release of alkyl radicals and exert a favorable inhibition effect at deep-seated tumor tissues. This work optimized the penetration effect of the photoactivatable nanomaterials and extended the bio-application of reactive species to the deep-seated tumor. Similarly, Qu et al. constructed a thernanostatic platform by combining AIPH, PCM, and the photoabsorbent PEG-MoS2 [147]. The designed system could also induce the generation of free radicals under NIR laser irradiation. Results suggested that the MoS2@AIPH-PCM nanosystem was capable of oxidizing and scavenging GSH inside the tumor cells, thus enhancing the therapeutic effect of alkyl free radicals. This work explored the mechanism of the influence of alkyl radicals on intracellular reducing substances for enhanced antitumor effect.

Organic biomaterials with intrinsic biodegradability have been extensively studied and employed as photothermal inducing agents. For
example, Xie et al. use Pluronic F-127 to encapsulate a conjugate between porphyrin and AIPH to form the TPP-NN NPs (Fig. 11d) [148]. Under 638 nm laser irradiation, the photothermal effect of porphyrin was transferred to AIPH and induced the generation of alkyl radicals. The generated alkyl radicals could be detected by monitoring the absorbance intensity changes of 1,3-diphenylisobenzofuran in DMF and the enhanced fluorescence from 2,7'-dichlorodihydrofluorescein in vitro. In vivo tests indicated that TPP-NN NPs could effectively inhibit tumor growth by producing elevated oxidative stress. These results demonstrated the excellent treatment efficacy of alkyl radicals-based reactive species to cancer cells. The work affirmed that the reactive species-based therapeutic strategies have great potential and are necessary to be further vigorously developed.

5. Conclusions and perspectives

In short, this review first discussed the recent development of ROS-based phototherapeutic nanoplatforms for phototherapy and also summarized the advantages or disadvantages of these various nanoplatforms. The photochemical mechanism and materials optimization approaches are introduced in detail for enhanced therapeutic efficacy to tumor cells. Second, the auxiliary functions of NO to various kinds of therapeutic modalities have been described, which laid the foundation of corresponding synergistic interaction in multifunctional nanomaterials. Third, the in-depth bioapplication mechanism of CO-containing nanoplatforms has been investigated, which holds much promise for the discovery and rational design of precision medicine. Subsequently, the reaction mechanism of photon-triggered generation of alkyl radical species was discussed, which provided innovative direction for biomedical nanomaterials design.

With the rapid development of nanobiotechnology, reactive species-based theranostics has achieved remarkable advances over the past decades [149,150]. Owing to the high controllability and low invasiveness of laser irradiation, photoactivatable nanogenerators have been extensively explored for reactive species-based cancer therapy [123]. Although several reactive species-generating photosensitive drugs have been used clinically, they have not been selected as the first-line treatment due to their intrinsic limitations [35]. The clinically approved reactive species-generators are all oxygen-based photosensitive nanodrugs (5-ALA, Verteporfin, and Porfimer Sodium), and few clinical research data on nitrogen and carbon-based reactive species have been collected. The oxygen-dependence nature of ROS would be doomed to fail to exert ideal therapeutic effects on solid hypoxia tumors. Moreover, the absorption wavelengths of most PSs are below 680 nm. The photo source with a wavelength below 680 nm is difficult to penetrate the deep tumor and activate the PSs distributed inside the tumor tissue [24]. The clinical application of photoactivatable radical species is currently focused on melanoma, lung cancer, and esophageal cancer [96]. For NO treatment strategies, when nitro is used as a ′NO donor, ultraviolet photo source and long-term laser irradiation limit its clinical translation, which required designing more optimized donors to meet the standard of clinical nanomedicine for efficient cancer therapy. For CO treatment, highly tumor-specific delivery of the CO precursor is essential due to strong adhesion between CO and hemoglobin.

Several strategies might be practical to address the above challenges. First, we need to pay more attention to study the therapeutic effects of photoactivatable reactive species, including attempts to treat the in situ tumors and metastatic tumors, as well as injection treatments from mice to larger animals, and systematically evaluating their biological distribution, comprehensive pharmacokinetic pharmacodynamics pathway and other clinical trials, to obtain more useful clinical data. Second, PDI with type I mechanisms should be focused on for treating malignant hypoxic tumors. Oxygen self-supplying drug systems or combining with hypoxia-activated drugs, such as TPZ should be developed. Third, up-conversion materials and two-photon absorption materials could be employed to solve the limitation of light wavelength. Also, the photon absorption capacity can be enhanced by rational molecular chemical designs, and the molecular structure could be smartly designed with high intersystem crossing efficiency from the excited state to the excited triplet state. Furthermore, synergistic therapy methods may endow effective therapeutic effects when the therapeutic outcomes of reactive species based on oxygen, nitrogen, and carbon are ineffective. Notably, due to the huge advantage of immunotherapy in individualized treatment, we must vigorously explore the immunoregulatory potential of reactive species-based nanomedicines [151–154]. Given the safety issues of NO and CO, it is a prerequisite to constructing nano-pharmaceutical systems with a more precisely controlled release profile. Besides, imaging detection strategies based on different reactive species can help qualitatively and quantitatively monitor the reactive species’ biodistribution and corresponding redox state, thus precisely controlling their beneficial treatment and relieving the systemic side effects [155–157].

It is noted that biosafety is a major concern for developing nano-therapeutics. The reactive species-generators comprise a diverse range of photoactivatable nanomaterials or nanoplatforms. The nanoformulations need to be thoroughly optimized [156]. The sense of biosafety and biocompatibility is crucial for developing radical species-based nanomedicine [159–165]. Multiple discipline collaboration, in particular the integration of nanotechnology with medicinal chemistry and pharmaceutical technology is important to promote the clinical translation of the photoactivatable nanomedicine for reactive species-based cancer therapy.

Declaration of competing interest

The authors declare no conflict of interest.

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