

Platelet-Based Nanoparticles with Stimuli-Responsive for Anti-Tumor Therapy

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Abstract: In addition to hemostasis and coagulation, years of studies have proved that platelets are involved in the whole process of tumor progression, including tumor invasion, intravasation, extravasation, and so on. It means that this property of platelets can be used in anti-tumor therapy. However, traditional platelet-based antitumor drugs often cause autologous platelet damage due to lack of targeting, resulting in serious side effects. Therefore, the researchers designed a variety of anti-tumor drug delivery systems based on platelets by targeting platelets or platelet membrane coating. The drug delivery systems have special response modes, which is crucial in the design of nanoparticles. These modes enhance the targeting and improve the anti-tumor effect. Here, we present a review of recent discoveries in the field of the crosstalk between platelets and tumors and the progress of platelet-based anti-tumor nanoparticles.

Keywords: platelet, tumor, targeted platelet, platelet membranes, stimuli-responsive drug delivery system, nanoparticle

Introduction

After the discovery of platelets in 1865, these anucleated cells once regarded as metabolic fragments of megakaryocytes, have progressively emerged as a compelling subject of research.¹ Extensive investigations over the years have yielded greater insights into the functions of platelets, particularly their interaction with tumor development.² The interaction between tumors and platelets serves as a prerequisite for hematogenous metastasis of tumors.³ Upon entering the bloodstream, tumor cells are the first to recruit and activate platelets, which subsequently facilitate the completion of the invasion-metastasis cascade by supporting the tumor cells.^{3,4} The study of tumor-platelet interactions plays a pivotal role in the development of platelet-associated anti-tumor drugs, while identifying the key targets between these entities offers valuable insights for anti-tumor therapy.^{5,6} However, common platelet inhibitors such as aspirin, ticlopidine, clopidogrel and prasugrel can attack platelets indifferently in vivo, causing autologous platelet damage and bleeding.⁷

Obviously, the defect of traditional platelet inhibitors is related to their lack of targeting. To this end, researchers designed nano-drug delivery systems. Drugs are carried in the shell assembled by excipients and modified by targeted peptides, so that the nano-drug delivery system can be actively or passively delivered to the target organ or target tissue. The nano-drug delivery system uses targeting and strong permeability retention effect to enrich the drug in the target organ / tissue, which not only enhances the anti-tumor effect, but also reduces the systemic side effects of the drug.⁸ Inspired by this, some teams designed platelet-based drug delivery systems using the natural targeting of platelets to tumors such as (i) targeted platelet

nanoparticles and (ii) platelet membrane-coated anti-tumor nanoparticles. Nevertheless, Shortcomings such as uncontrolled release of the drug make this antineoplastic drug do not achieve the desired effect.⁹

The development of stimuli-responsive nano-drug delivery system can solve the problem of random drug release in the complex background of human body. On the basis of the original nano-drug delivery system, the research teams add related peptides, photothermal agents, sound sensitizers and other substances to the system, or use materials that can be degraded by oxides, acids and other substances to make the shell of the system. Then the drug delivery system was stimulated by a variety of special conditions in vivo and in vitro (in vivo factors: enzyme, ROS, pH, H₂O₂, glutathione (GSH), etc., and in vitro factors: temperature, light and ultrasound (US), etc.) to unload the drug in specific release sites such as tumors.

Interaction Between Tumor and Platelets

The interaction between tumor and platelet in vivo has been widely studied, and a large number of studies have found that platelets play a key role in the process of tumor invasion, intravasation and extravasation. Firstly, platelets are activated by platelet activation mediators such as CD40, thrombin and thromboxane A₂ (TXA₂) secreted by tumor cells or stromal cells. Activated platelets secrete transforming growth factor-beta (TGF- β), platelet derived growth factor-(PDGF) and other factors to assist the epithelial-mesenchymal transition (EMT) of tumor cells and promote tumor entry into the blood. Subsequently, platelets will pass through the physical barrier, inhibit NK cells, inhibit apoptosis and other processes to protect the tumor and make it metastasize smoothly. Platelets play an important role in the cascade of tumor invasion and metastasis, so it is of great significance to study this interaction and related targets for the research and development of anti-tumor drugs.

Tumor Activates Platelets

Tumors are often regarded as ‘non-healing wounds’, which play a crucial role in platelet activation. Tumors activate platelets through diverse mechanisms, with thrombin continuing to be the most effective platelet activator in the platelet activation process. Tissue factor (TF) is expressed on the cell membranes of some cancer cells, triggering the plasma coagulation cascade, resulting in thrombotic reactions and thrombin generation, which is activated primarily through protease-activated receptors (PARs).^{10–12} Among the PARs (PAR1, PAR2, PAR3, and PAR4), platelet activation is mainly associated with PAR1 and PAR4, while PAR-2 is not expressed on platelets, and PAR-3 acts solely as a cofactor for PAR-4.^{13,14} Tumor cells can also secrete soluble mediators, including adenosine diphosphate (ADP) and TXA₂, to activate platelets.^{15,16} Furthermore, activated platelets release ADP and enhance their own activation through autocrine and paracrine mechanisms.¹⁷ ADP acts as an agonist for two purinergic G protein-coupled receptors (P2Y₁, P2Y₁₂) in platelets, contributing to platelet activation and aggregation.¹⁸ TXA₂, through its receptor TXR, also influences platelet activation and aggregation.¹⁹ When interacting with CD40 factors produced by tumors, platelets secrete the CD40 receptor (CD40L), a vital source of CD40L in the blood, resulting in broad platelet activation.^{20,21}

In recent years, CLEC-2 has been shown to be a platelet receptor for podoplanin proteins on tumor cells,²² and podoplanin proteins, which are widely expressed in tumor cells, can activate platelets through the immunoreceptor tyrosine-based activation motif (ITAM) and downstream kinase, thereby cross-linking platelet CLEC-2.⁶ Additionally, receptors for advanced glycosylation end product, including S100, amyloid β , and high mobility group frame 1 (HMGB1), can also stimulate local platelet activation, with HMGB1 predominantly binding to Toll-like receptors on the surface of tumor cells.²³ P-selectin on platelets can also induce platelet activation by interaction with the P-selectin receptor (PSGL-1) on tumor cells.²⁴ Furthermore, CD97, an adhesive G protein-coupled receptor expressed by cancer cells, was discovered by Ward et al to activate platelets.²⁵ Other tumor patients lack the enzyme that cleaves von Willebrand factor (vWf), leading to platelet activation by a highly polymerized form of vWf in the circulation.²⁶ Platelets contain matrix metalloproteinases (MMPs), which, when triggered by malignancies, can activate platelets as well.^{27,28} Glycoprotein VI can direct ITAM signaling to activate platelets, thereby controlling several processes including platelet adhesion, aggregation, and procoagulant activity.²⁹ To enhance the effects of platelet activation, Tumor cells also stimulate platelet aggregation (TCIPA).³⁰ Depending on the type of stimulation, aggregated platelets through TCIPA can secrete proangiogenic and tumor-promoting factors that are essential for tumor migration and growth.^{31,32}

Interaction Between Platelets and Immunocytes

It is worth noting that platelets also interact with immunocytes, which is also extremely important in tumor progression. In previous studies, it has been found that platelets are involved in regulating the function of immunocytes such as natural killer (NK) cells, dendritic cells, neutrophils, macrophages and lymphocytes.³ In tumor progression, platelets play an important role in extracellular neutrophil trapping (NET) and NK cell inhibition.³³

NET is a unique immune response of neutrophils to neutrophil elastin and DNA excretion substances in pathogens. The DNA network in NET can also be used as a chemokine to attract and capture circulating tumor cells (CTC), but this capture is not harmful, NET will enhance the proliferation and invasion of CTC.³⁴ In recent years, it has been found that NET can also enhance the permeability of local blood vessels and facilitate the extravasation of tumor cells.³⁵ In addition, substances such as elastin released by NET can also break down laminin and provide help for tumor invasion.³⁶ NET also captures platelets and activates platelets to form thrombus, which in turn promotes the formation of NET and amplifies this effect.³⁷ In tumors, activated platelets immediately bind to neutrophils through P-selectin, and induce Net production through thrombin, TLR4 and HMGB1 produced by platelets, thus enhancing platelet activation and aggregation, thus promoting tumor progression.^{38,39}

NK cells are deadly to CTC, but platelets can help CTC survive the killing of NK cells. Platelets can transfer histocompatibility class I molecules (MHC-1) to the surface of tumor cells, shielding tumor cells from immune recognition and diminishing NK cell function, and platelet-derived TGF- β can also reduce NK cell activity.⁴⁰ Platelet-derived RGS18 can also protect CTC from NK-mediated immune surveillance by participating in immune checkpoint HLA-E: CD94-NKG2A.⁴¹ In addition, platelets can induce the exfoliation of NKG2D³³ and NKG2DL⁴² ligands to enhance the immune escape of CTC to NK cells.

Platelets Assist in Tumor Invasion

Tumor invasion is associated with tumor EMT, and crucial soluble mediators such as platelet-derived growth factors and matrix metalloproteinases, which are released by activated platelets are essential for the induction of EMT.¹⁹ EMT confers mesenchymal cell competence to epithelial cells, significantly increases cell motility, contributes to tumor cell invasiveness and facilitates tumor invasion by degrading the extracellular matrix (ECM) of metastasis.^{43,44}

EMT is a biological process through which epithelial cells undergo a specific program to acquire a mesenchymal phenotype. EMT is a biological process that is regulated by genes such as the SNAIL family, the TWIST family, and the zinc finger E-box binding homology box (ZEB) family. These genes, in turn, are influenced by signaling pathways such as TGF- β , Smad, P38, Wnt, NF- κ B, and PI3K-AKT. The expression of epithelial genes such as E-cadherin, Occludins, claudins, and ZO1 is suppressed, while the expression of mesenchymal phenotype-related genes such as N-cadherin, Vimentin, and Fibronectin is enhanced. Additionally, increased degradation of the extracellular matrix by MMPs is a significant feature of EMT.^{45,46}

In addition to gene regulation, platelets are also involved in the regulation of EMT-related signaling pathways to promote the invasive-metastatic cascade response of tumor cells. For instance, platelet-secreted TGF β 1 can activate EMT through the TGF β /Smad pathway. Platelets can also activate the NF- κ B pathway upon contact with tumors and thus enhance TGF β 1 secretion from tumor cells, promoting EMT activation.⁴⁷ Platelets in breast cancer can also activate EMT by inducing the NF- κ B pathway, conferring an aggressive phenotype.⁴⁸ Moreover, a study on cholangiocarcinoma found that platelet-released PDGF activates EMT via the P38/MAPK pathway.⁴⁹ In hepatocellular and breast cancers, platelets can also promote EMT through PAR activation of TWIST.⁵⁰

Tumor cells that complete EMT are also vulnerable to anoikis, a programmed cell death that occurs when cells detach from the extracellular matrix.⁵¹ Resistance to anoikis is a crucial prerequisite for tumor cells to metastasize. Platelets frequently assist tumor cells in resisting evading by producing autotoxin (ATX) during tumor growth, mainly through the conversion of lysophosphatidylcholine to lysophosphatidic acid (LPA), followed by LPA binding to LPA receptor (LPAR-1) on tumor cells via RhoA-G α 12/ 13-YAP-1 signaling pathway promoting resistance to anoikis and facilitating tumor progression.⁵²

Platelets-Based Tumor Intravasation

Tumor cells that have undergone EMT and survived anoikis can infiltrate. Tumor cell invasion into lymphatic systems, blood vessels, and adjacent tissues occur in various ways. This article primarily focuses on the processes that occur when tumor cells enter the bloodstream. Tumor endosmosis and extravasation require the destruction of the endothelium connection to pass through endothelial cells, known as transendothelial cell migration (TEM).^{53,54} Due to the distinct ways in which tumor cells migrate through endothelial cells, endosmosis and extravasation differ. Tumor cells invade by creating intercellular connections with the body's blood vessels and producing local blood vessels during intravasation. This process is regulated by angiogenesis-related factors such as vascular endothelial growth factor (VEGF), angiopoietin, and PDGF.^{55,56} Ward et al discovered that expressed in cancer cells, CD97 adhesion GPCR may activate platelets and cause LPA release from platelets, promoting tumor cell migration across the endothelium.²⁵ Endothelial cells activated by thrombin have a circular appearance and lack adhesion connections, which can encourage tumor cell movement across the endothelium.⁵⁷ Furthermore, vascular diameter, specific proteases such as uPAR and MMPs can also influence tumor invasion into blood arteries.^{55,58}

The tumor invades blood arteries and transforms into CTC, which enter the circulatory system and prepare for metastasis. However, most CTC is destroyed during this process by blood flow shear stress and NK cells. As a result, CTC must work with platelets to avoid numerous hazards in the circulatory system.

To begin, CTC must conserve enough platelets to create tumor cell immune thrombi by TCIPA. The mechanism of CTC-mediated TCIPA is primarily associated with various platelet activation pathways, including PAR binding to TF, ADP and P2Y₁₂ binding, TXA₂ and TXR binding, CD40L and CD40 binding, podoplanin and CLEC-2 binding, HMGB-1 and TLR4 binding and so on. Excessive vWF binding to platelet glycoprotein GPIb IX–V, as well as enhanced synthesis of PAI-1 and MMPs and accelerated platelet aggregation.³ This implies that CTCs stimulate and recruit platelets, enabling them to promptly respond to multiple life-threatening challenges in the circulatory system.

Subsequently, platelets that aggregate around tumor cells adhere to them through the action of adhesion molecules including integrin α Ib β 3, P-selectin, CD97, and GPIb-IX-V/GPIIb-IIIa.^{59,60} These proteins are also involved in platelet activation and the development of TCIPA.⁶¹ Platelets adhering to the surface of tumor cells support the survival of CTCs in the circulatory system, shielding them from blood flow shear, tumor necrosis factor- α (TNF- α), and cell death induced by NK cells.⁶²

Tumor Extravasation via Platelets

CTCs that survive in the circulation eventually come to a gradual standstill as peripheral capillary blood flow slows down and adheres to endothelial cells in the presence of P-selectin and integrins, among others, and undergo TEM, similar to the process of endo-osmosis.⁵² In addition to actively releasing MMPs to break down the ECM to promote extravasation, platelets also release PDGF, 12(S)-hydroxyeicosatetraenoic acid (12-HETE), and LPA to, in turn, stimulate the production of MMPs by CTCs,⁶³ with 12-HETE also promoting contraction of endothelial cells to promote extravasation.⁶⁴ ATP binds to the P2Y₂ receptor on vascular endothelial cells, leading to the disruption of tight junctions and increasing endothelial permeability.⁶⁵ ATP released from activated platelets has also recently been found to promotes endothelial cell contraction to increase permeability and enhance the expression of endothelial adhesion molecules, which in turn promotes extravasation.⁶⁶ However, in some cases, CTCs are not required for TEM, and stagnant cancer cells may proliferate within the vascular lumen, leading to the growth of large intraluminal tumor colonies that eventually rupture the nearby endothelial wall, allowing direct access to the tissue parenchyma.⁶⁷ Figure 1 provides an illustrative overview of this interaction.

Platelet-Based Anti-Tumor Nanoparticles

Nanoparticles, as emerging anti-tumor tools in recent years, are designed to transport anti-tumor drugs using unique drug delivery systems, addressing challenges such as limited specificity, high toxicity, and drug resistance encountered by conventional chemotherapeutic drugs like Tegretol and Doxorubicin. Platelets have garnered significant attention in anti-tumor nanoparticles research due to their innate tumor-targeting properties. Anti-tumor nanoparticles with stimulus-responsive drug delivery system designed using platelets have higher specificity, and when combined with radiotherapy and ultrasound, these drugs can be concentrated near the tumor site, minimizing damage to healthy cells. Currently, platelet-based anti-tumor nanoparticles

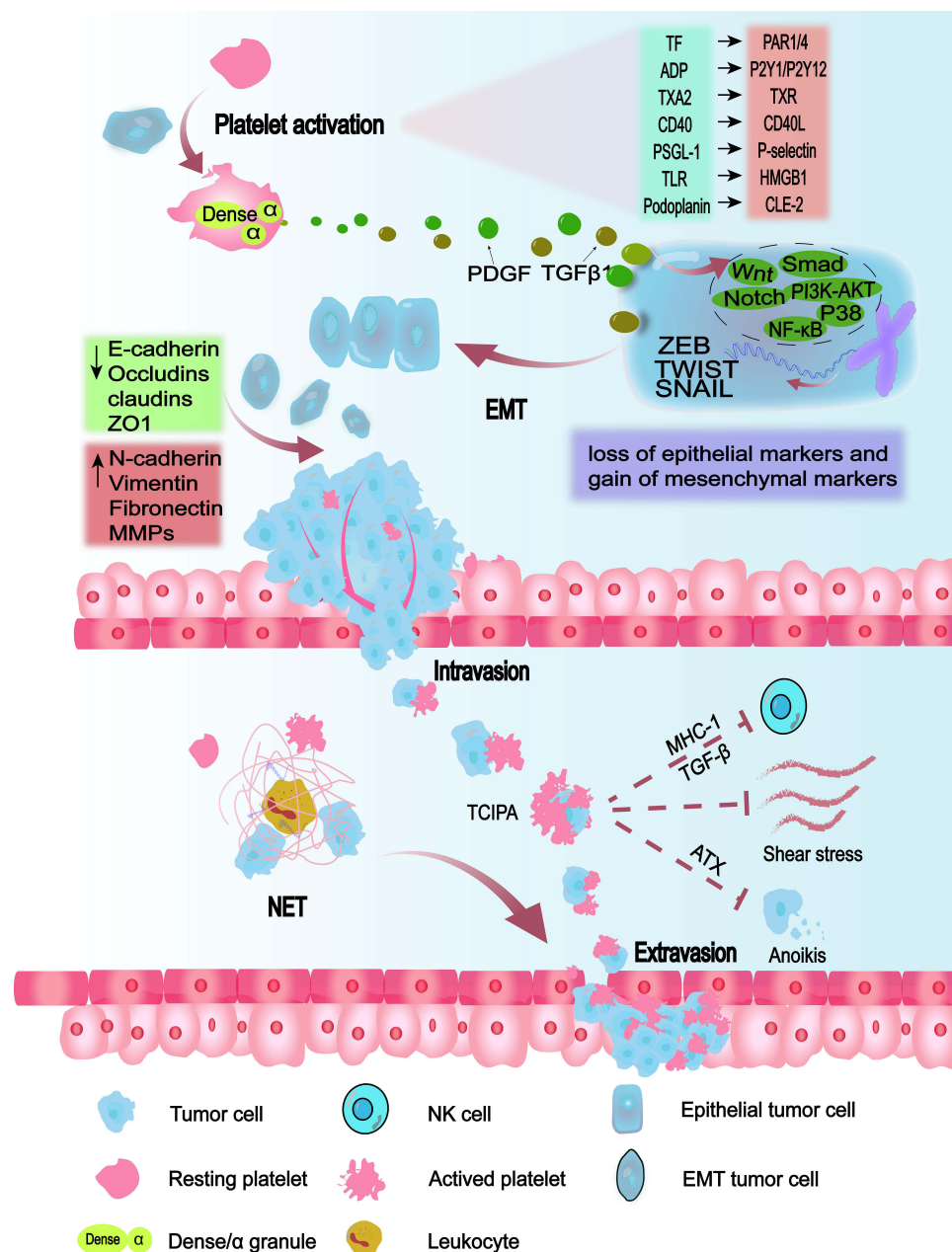


Figure 1 Platelets participate in the cascade of tumor invasion and metastasis. Tumor activated platelets: (1) tumor cell Tf binds platelet PAR1/4; (2) tumor cell ADP binds platelet P2Y2/P2Y12; (3) tumor cell TXA2 binds platelet TXR; (4) tumor cell CD40 binds platelet CD40L; (5) tumor cell PSGL-1 binds platelet P-selectin; (6) tumor cell TLR binds platelet HMGB1. (7) tumor cell podoplanin binds to platelet CLEC-2. PDGF and TGF-β released by activated platelets regulates Snail, Twist, ZeB and other genes and participates in tumor EMT through Wnt, Notch, NF-κB, SmAD, P38, PI3K-AKT and other pathways. During this process, the expression of E-cadherin, Occludins, Claudins and ZO1 was down-regulated, while the expression of N-cadherin, Vimentin, MMPs and fibronectin was enhanced. Then the tumor infiltrates into the blood circulation with the help of platelets. Tumor recruits activate platelets through TCIPA to form a physical barrier on the surface of tumor cells to protect the tumor from blood flow shear. On the basis of physical barrier, platelets also use MHC-I and transforming growth factor-β to protect tumor cells from recognition and killing of NK cells, and release ATX to participate in the resistance of tumor cells to apoptosis. At the same time, platelets help neutrophils form NET. NET can promote the extravasation of tumor cells. Eventually, tumor cells will complete the metastasis with the help of platelets.

primarily consist of platelet-targeted drugs and platelet membrane-encapsulated anti-tumor nanoparticles designed to specifically target tumors.

Tumor-Based Platelet Targeted Nanoparticles

Platelets are commonly implicated in tumor growth through the interaction of their membrane receptors with tumor cells. Platelet-targeted medicines that frequently exhibit positive imaging and anti-tumor effects have targeted these receptors. In order to inhibit platelet activation, Zhou's team developed nanoparticles made of albumin that can coat the surface of platelets with

ADP, collagen (a GPVI agonist), and a thrombin activation site. Utilizing this nano platform can enhance the effectiveness of anti-tumor immunotherapy by promoting increased permeability of tumor blood vessels and T-cell intravasation.⁶⁸

Nevertheless, systemic administration of anti-platelet drugs often harms autologous platelets and disrupts functions, including coagulation. Nanoparticles with targeted delivery capabilities circumvent damage to autologous platelets and enable localized dosage at the tumor site for improved anti-tumor effects (Table 1). Initially, researchers altered nanoparticles by incorporating tumor-targeting proteins, such as the CREKA (Cys-Arg-Glu-Lys-Ala) peptide,⁵⁹ TM33 peptide,⁶⁰ fucoidan segment,⁶¹ and P-selectin-targeting peptide (PSN),⁶² and coupled them with platelet inhibitors to concentrate them near tumor cells. Subsequently, they employed specific excitation techniques crucial for the anti-tumor effects to facilitate drug release. Currently, two distinct types of excitation techniques exist: internal stimuli, such as an acidic pH, redox reactions, or tumor microenvironment (TME) enzymes, and external stimuli, including laser and ultrasound.

Excessive acidity results from the accumulation of lactate and hypoxia in the TME due to the increased metabolic rate of tumor cells.⁷⁹ Building upon this knowledge, researchers have successfully achieved drug release by incorporating ionizable groups of the drug into nanomaterials. When the nanoparticles is exposed to an acidic environment, the binding sites between the drug and the nanomaterial become ionized, resulting in drug release. In one study, Zhang's team prepared an acid-responsive nanoparticles, CREKA-lipo-T. The platelet inhibitor ticagrelor was encapsulated in liposomes and conjugated with the tumor-targeting pentapeptide CREKA, facilitating the enrichment of the nanoparticles in the vicinity of the tumor. In a 4T1 mouse model, CREKA effectively targeted the fibronectin-fibronectin complex in the tumor vessel wall. CREKA-Lipo-T actively bound to tumor vessel microthrombi, leading to local release of ticagrelor, which interacts with the P2Y12 receptor, through an acid response. This mechanism successfully inhibited tumor-associated platelet function and blocked tumor lung metastasis, without inducing significant side effects such as bleeding complications in the experiment.⁷⁷ Subsequent research will heavily concentrate on acid-responsive nanoparticles as it provides promising approaches for the delivery and release of anti-tumor medications.

Besides its highly acidic nature, the tumor microenvironment exhibits elevated levels of glutathione (GSH) and hydrogen peroxide (H₂O₂). These components reduce the susceptibility of tumor cells to oxidative stress and position GSH/H₂O₂ as potential catalysts in the advancement of nanoparticles. The existing design strategy for GSH relies primarily on exploiting the GSH present in tumor cells to cleave disulfide bonds and facilitate drug release from nanoparticles. For example, Xu et al modified albumin by incorporating terminal sulfhydryl groups (-SH) and subsequently s-nitrosylated the albumin to generate

Table 1 Tumor-Based Platelet-Targeted Nanoparticles

Nanoparticles	Nanoplatform	Content	Response way	Reference
Fu-UK/ICG@SiAu NRs	Mesoporous silica-coated gold nanorods	Indocyanine green/urokinase-type plasminogen activator	Laser	[69]
BDCCN	FeIII -doped C3 N4	Chlorogenic acid	pH/H ₂ O ₂	[70]
PLP-D-R	Copolymer poly(ether imide)-poly(lactic-co-glycolic acid)2	Doxorubicin/ antiplatelet antibody R300	MMP2	[71]
BSA-IRLA@RVs	Scaffold of bovine serum albumin	L-Arginine/ photosensitizer IR783	Laser	[72]
IGBN	Copper-based MOF	Cisplatin-arginine (Pt-Arg) prodrug	pH/GSH	[73]
CyBA/PFM NPs	Amphiphilic phenyl boronated fucoidan/ maltodextrin polymers	Fucoidan segment	H ₂ O ₂	[74]
Ptx@AlbSNO	S-nitroso albumin (AlbSNO)	Paclitaxel/NO	GSH	[75]
DPC@ICD-Gd-Tic	Liposomal nanoparticle	Ticagrelor/ photosensitizer IR780	GSH	[76]
CREKA-Lipo-T	Liposomal nanoparticles	Ticagrelor	pH	[77]
TM33-GON/TNA	Gelatin/oleic acid nanoparticle	Tanshinone IIA	MMP2	[78]

nitrosylated human serum albumin (AlbSNO). They employed AlbSNO as a shell to construct nanoparticles that encapsulated paclitaxel (Ptx), resulting in the formation of Ptx@AlbSNO complexes. Specifically, AlbSNO can induce targeted release of nitric oxide (NO) upon reductive stimulation by the high GSH concentration found in the 4T1 and CT26 tumors. NO effectively inhibits platelet activation and the secretion of TGF- β , which is crucial in alleviating the immunosuppressive tumor micro-environment. In turn, disruption of the AlbSNO shell leads to the release of paclitaxel, which in turn develops anti-tumor effects.⁷⁵ Hydrogen peroxide (H₂O₂), another vital reducing agent in the TME, plays a significant role in the elimination of reactive oxygen species (ROS) and platelet activation processes. Wang et al developed an H₂O₂-responsive NIR probe self-assembly-triggered nanosystem for targeted platelet nanoformulations CyBA/PFM. The CyBA/PFM NPs actively bind thrombi by associating with highly expressed P-selectin, and the rocket algae etiolated glycan segment within the vector also facilitates specific delivery of therapeutic polymers and probes to platelet-rich arterial thrombi. In a mouse model of FeCl₃-induced arterial thrombosis, the phenyl borate group captures the electron-donating oxygen atom. Subsequently, H₂O₂ prompts the release of the phenyl boride group, resulting in nanoparticle dissociation and the generation of fluorescent CyOH. During this process, H₂O₂ is significantly consumed to reduce platelet activation and achieve inhibition of platelet function.⁷⁴

The MMPs enzyme family plays a crucial role in the tumor progression during the invasion metastasis cascade of malignancies. MMP2 is overexpressed selectively on the plasma membranes of tumor endothelial cells and tumor stromal cells. Therefore, Li's team utilized this property to develop a nanocomposite consisting of a core made of a block copolymer, poly (ether imide)-poly(lactic-co-glycolic acid)₂ (PEI-(PLGA)₂), loaded with the antiplatelet antibody R300 and the chemotherapy drug doxorubicin (Dox). The core was coated with an MMP2 cleavable peptide, lecithin, and polyethylene glycol phospholipid to form the shell. R300, a monoclonal antibody targeting the GPIIb α subunit of the platelet vascular hematopoietic factor receptor, effectively depletes platelets by inducing microaggregation (3–5 platelets). The resulting macroaggregates are rapidly cleared through phagocytosis without activating the coagulation system. Upon intravenous injection into MCF7 tumor-bearing nude mice, the polyethylene glycolated phospholipids in the shell layer of the nanoparticles inhibit nonspecific aggregation, prolonging their half-life. Once they reach the tumor microenvironment, tumor-associated MMP2 triggers a burst release of the drug by cleaving the MMP2-cleavable peptide on the shell layer.⁷¹ Enzyme-responsive drugs pose greater challenges to the nanoshell than the two response modalities mentioned earlier. The selection of enzymes and the modification of the nanoshell require more stringent criteria. Further research is necessary to establish their potential as clinical candidates.

In addition to internal pathways, external stimuli play a crucial role in stimulating the desired effects. As one of the routine treatment modalities in cancer therapy, conventional radiation therapy is less effective on deep-seated tumors due to the diminishing energy of light in the body, and the abundance of GSH and H₂O₂ in tumor tissue can also weaken the harm of ROS produced by radiation. To address these shortcomings of radiotherapy, researchers have combined it with the development of nanoparticles, which use specially designed nanoparticles to deliver metal particles or photothermal agents with excellent photothermal conversion near the tumor to enhance the effect of radiotherapy in the presence of radiation for anti-tumor purposes. For instance, Chang's group successfully developed a nanocomposite called Fu-Uk/ICG@SiAu NRs. This nanocomposite demonstrated high efficiency and safety in targeting tumor thrombi, exhibiting photothermal/enzymatic thrombolytic effects.⁶⁹ Ma et al developed a BSA-IRLA@RVs complex that targets platelets through the encapsulation of protein nanoparticles within red blood cell membranes. Using the constructed cRGD immune membrane structure, the nanoparticles could actively target platelets and cancer cells, respectively, through overexpression of integrins α Ib β 3 and α v β 3, among others. The drug is loaded with biological nitric oxide synthase, l-arginine (LA), and the photosensitizer IR783, with the LA molecule acting as a nitric oxide precursor. In the U14 tumor-bearing mice model, IR783 utilizes oxygen under NIR stimulation to generate reactive oxygen species (ROS). This process leads to the oxidation of the guanidine group of LA, releasing nitric oxide (NO). Consequently, NO specifically inhibits tumor-associated platelet activation, resulting in weakened endothelial junctions and erythrocyte intravasation, along with a significant increase in drug concentration within the tumor tissue.⁷²

Platelets Delivery Anti-Tumor Drugs

Researchers have designed a variety of drug delivery systems with platelets. At present, the main ideas are as follows: 1. platelet membrane-coated nanoparticles. 2. living platelets-based nanoparticles. 3. platelets with drug or antibody coupled on its surface. 4. platelet EV based nanoparticles (Table 2).

Table 2 Platelet-Based Nanoparticles

Nanoparticles	Nanoplatform	Content	Response Way	Reference
Platelet membrane-coated nanoparticles				
PM-NP/PTX	Dextran nanoparticles	Paclitaxel	Thrombin	[80]
PMNPs	Poly(lactic-co-glycolic) acid	Doxorubicin /indocyanine green	PTT	[81]
PSCI	Mesoporous silica nanoparticles	Cinnamaldehyde / sonosensitizer IR780	US	[82]
PINPs@PM	Poly(d, l-lactide-co-glycolide)	Indocyanine green	PTT	[83]
PLTM-DOX@MPDA	Mesoporous polydopamine nanoparticles	Doxorubicin	PTT	[84]
rVAR2-PM/PLGA-ss-HA	Biodegradable PLGA conjugate nanoparticles	Docetaxel	GSH	[85]
PLTM-CS-pPLGA/Bu NPs	Chitosan oligosaccharide (CS)-poly(lactic-co-glycolic acid) (PLGA) copolymer	Bufalin	pH	[86]
BMSNR@PM	Chitosan oligosaccharide (CS)-poly(lactic-co-glycolic acid) (PLGA) copolymer	Bismuth nanorod	PTT	[87]
PMS	Mesoporous Fe single-atom nanozyme	-	PTT	[88]
H-L-D-Z@PM NPs	Zeolitic imidazolate framework-8 (ZIF8)	Lactate oxidase / doxorubicin / hemoglobin	pH	[89]
PG@HGNs	Hollow gold nanoparticles	-	pH	[90]
PB/PM/HRP/APT	Prussian blue nanoparticles	-	PTT	[91]
PCLP-CUR	Chitosan-modified liposome	Curcumin	pH	[92]
HMMD@PG	Hollow mesoporous copper sulfide	MnO ₂ / doxorubicin	PTT	[93]
PMBNs	Biomimetic mesoporous organosilicon	Type I AIE photosensitizer TBP-2	GSH	[94]
IR780@PLGA/DOX NPs	Poly (lactic-co-glycolic acid) (PLGA)	Sonosensitizer IR780/dobiroxacin	pH	[95]
PCDD NPs	The third-generation poly-L-lysine dendrimer	Chlor-in e6/ docetaxel	ROS	[96]
P-MOF-siRNA	Metal-organic framework	si-RNA	pH	[97]
Living platelets-based nanoparticles				
PDNGs	Nanogels	Doxorubicin	Tumor	[98]
PLT@PDA-DOX	Polydopamine	Dobiroxacin	PTT	[99]
P-P-IO	-	aPDL1 / IONPs	Endothelium	[100]
IO-PG-DOX-Ce6	Iron Oxide (IO) / polyglycerol	Chlor-in e6	laser	[101]
PDA@Dox NPs	Polydopamine	Fluorescent photothermal agent IR820 / doxorubicin	PTT/pH	[102]
α MM-PEG@PLTs	MnO ₂ -coated porphyrinic metal-organic framework	α -methyl-DL-tryptophan	US/GSH	[103]

(Continued)

Table 2 (Continued).

Nanoparticles	Nanoplatform	Content	Response Way	Reference
Platelets with drug or antibody coupled on its surface				
PDNGs	Nanogels	Doxorubicin	Tumor	[98]
P-P-IO	-	aPDLI/ IONPs	Endothelium	[100]
CAR-T-P aPDLI@gel	Hyaluronic acid hydrogel	CAR-T cells/ P-aPDLI	Thrombin	[104]
GALA-GNGs-P	Protein nanogels	Pore-forming peptide/Granzyme B	pH	[105]
Platelet EV based nanoparticles				
RGD-NPVs@MNPs/DOX	-	Melanin nanoparticles/ doxorubicin	PTT	[106]

Platelet Membrane-Coated Nanoparticles

Platelet membrane-coated nanoparticles overcome the limitations of conventional coated non-scaling polymers like polyethylene glycol (PEG) and dextran, enhancing targeting capabilities and extending the half-life. Conventional anti-tumor therapy faces challenges in accessing the tumor cytoplasm, whereas platelet membrane-coated drugs utilize signaling molecules on the membrane surface to target tumors and facilitate platelet phagocytosis, significantly improving drug delivery. Platelet membrane-coated drugs are more intricate than platelet-targeted drugs, with the preparation and coating of platelet membranes being crucial. Platelets are initially isolated from the blood through differential or gradient centrifugation. Subsequently, they undergo repeated freeze-thaw or platelet lysis using hypotonic media. Afterward, centrifugation is performed to obtain platelet membrane fragments. These isolated platelet membranes are then mixed with inner core nanoparticles and coated onto the nanoparticles using ultrasound, co-extrusion, or microfluidic electroporation. The resulting nanocomposites are characterized using various techniques, including transmission electron microscopy, dynamic light scattering, sodium dodecyl sulfate-polyacrylamide gel electrophoresis, and Western blotting analysis.^{84,85,97}

Researchers have developed Platelet membrane-coated anti-tumor nanoparticles that target specific tumor features and activate multiple pathways, thereby enhancing safety. Zhuang et al encapsulated siRNA within acid-responsive porous metal-organic framework (MOF) nanoparticles. These nanoparticles were then coated with platelet membranes to create P-MOF-siRNA complexes. The P-MOF-siRNA complex specifically targeted breast cancer cells through the presence of platelet membranes on its surface. Upon endocytosis by the target cells, the acidic environment triggered the dissociation of the MOF scaffold, resulting in drug release and subsequent anti-tumor effects. In conclusion, this method presents a promising approach for developing clinical anti-tumor drugs through the preparation of acid-responsive platelet membrane-coated nanoparticles. Moreover, the highly reproducible P-MOF-siRNA complex offers versatility for exploring alternative therapeutic options by modifying the loaded drug.⁹⁷

To target the role of GSH in tumor tissue, Ning's team loaded the type I AIE photosensitizer TBP-2 into mesoporous organo-silica nanoparticles. Subsequently, these nanoparticles were coated with platelet membranes, resulting in the formation of PMBN complexes. The P-selectin present on the platelet membrane specifically targets CD44 in cancer stem cells (CSCs), leading to the phagocytosis of CSCs. Subsequently, intracellular GSH triggers the degradation of the disulfide bonds in the organo-mesoporous silica nanoparticles, resulting in the release of TBP-2. Ultimately, the release of TBP-2, when combined with external light, exerts an anti-tumor effect. In a murine breast cancer tumor model, the results demonstrated the successful targeting of CSCs by PMBN, leading to a decrease in the number of CSCs at the tumor site and reduced recurrence following treatment.⁹⁴

Huang et al employed mesoporous silica nanoparticles (MSN) as carriers, loaded with the oxidative stress amplifier cinnamaldehyde (CA) and the acoustically sensitive agent IR780. These nanoparticles were further coated with platelet membranes. Upon ultrasound exposure, CA was released from the MSN, resulting in GSH depletion and subsequent ROS increase. This disruption of the cell membrane led to drug release, ultimately exerting anti-tumor effects in the 4T1 cancer-bearing mouse model. Experiments have demonstrated the effective inhibition of tumor growth by this drug. CA-induced GSH depletion hinders the ability of tumors to counteract ultrasound-induced oxidative stress. In summary, the combined approach of ultrasound treatment with anti-tumor drugs offers improved biosafety and efficacy, with significant potential for clinical application in combination with traditional anticancer therapies.⁸²

The results of radiotherapy in targeting platelet nanoparticles are equally favorable in platelet-coated nanoparticles. Qi's team developed a mesoporous Fe-SA enzyme (PMS) coated with platelet membranes, utilizing photothermal therapy (PTT) to induce mitochondrial damage in breast cancer. This approach enables secondary near-infrared (NIR-II) driven PTT at mild temperatures. The developed PMS exhibited favorable NIR-II photothermal properties, high peroxidase (POD) activity, and excellent tumor-targeting capabilities. Additionally, it demonstrated the ability to carry protein drugs. Upon targeting and subsequent phagocytosis by the tumor, the SAzyme, possessing high POD activity, catalyzes the generation of -OH from endogenous H₂O₂ within the tumor, resulting in mitochondrial damage. In turn, mitochondria encode genes that contribute to the development of heat resistance in eukaryotic cells. This suggests that damaging mitochondria can reduce the heat resistance of tumor cells, enabling their eradication under mild PTT conditions. These findings are supported by data from *in vivo* experiments.⁸⁸

Living Platelets-Based Nanoparticles

In addition to using platelet membranes to cover nanoparticles, researchers also use living platelets as a drug delivery system to treat diseases. At present, drugs such as doxorubicin (DOX) are used for platelet loading, which is mainly loaded into platelets through passive osmosis of drugs.¹⁰⁷ In the study of Xia,¹⁰⁸ they prepared Au-Hb@PLT complex. In tumor-bearing mice, the engineered platelets will activate and release gold nanoparticles under radiation, greatly enhancing the effect of radiotherapy. This laser-responsive living platelet complex shows a good effect in tumor therapy, and the BNPD-Ce6@Plt constructed by Xu et al has also proved this in the glioblastoma mouse model.¹⁰⁹

In addition to external stimulation to activate engineering platelets, live platelets are also activated by tumor cells, which greatly reduces the difficulty of design. In the study of Li et al,¹¹⁰ living platelets loaded with DOX were activated by Lewis lung cancer cells and unloaded drugs, the results showed that the therapeutic effect was significantly higher than that of DOX alone, and there was no serious toxicity related to DOX in mice. In addition, some similar engineered platelets have been tested in many different tumors, and the results have shown positive significance (Table 2). However, it is worth noting that these engineered platelets can be activated by tumor cells, thrombosis, etc., which means that they may also be activated by other substances, especially in the complex background of the human blood system. Moreover, the preparation and storage of platelets have great restrictions on this kind of engineered platelets. Platelets stored at room temperature often have problems such as bacterial contamination, activation and degranulation, while platelets stored at low temperature have the problem of functional change. These defects limit the participation of living platelets in the construction of these engineered platelets, which need to be overcome in order to be widely used.¹¹¹

Platelets with Drug or Antibody Coupled on Its Surface

As mentioned earlier, there are a large number of surface markers on the surface of platelets, which means a new research idea. By coupling antibodies or drugs with platelet membranes, the researchers improve the anti-tumor ability of drugs through the natural targeting between platelets and tumors. Wang et al¹¹² coupled programmed death ligand 1 to the platelet surface. This engineered platelet performed well in the mouse model of primary melanoma or triple negative breast cancer, which significantly increased the overall productivity of mice. This idea was also verified in Li's experiment, which used hydrogel to locally consume tumor-related macrophages, which enhanced the coupling of anti-programmed cell death protein 1 antibody to platelets and had a good inhibitory effect on tumor recurrence.¹¹³ However, it is worth noting that the coupling of platelets with these substances may lead to excessive concealment of platelet surface markers, thus weakening the targeting ability of platelets to tumor cells and affecting the anti-tumor effect.

Platelet EV Based Nanoparticles

Extracellular vesicles (EV), as a kind of cell information granules, are substances involved in a variety of functions such as coagulation, immunity and tumor progression in the human body.¹¹⁴ As the most abundant source of EV, platelet microparticle (PMP) is an important element in the process of tumor invasion and metastasis. Because PMP comes from platelets, it retains the characteristics of platelets to some extent, including surface markers such as CD41, CD42, P-selectin and functional substances such as platelet activating factor, angiogenic factor and chemokines.^{115,116} The tumor-promoting function of PMP is similar to that of platelets has been confirmed. Therefore, PMP is a potential substance for the development of antineoplastic drugs.¹¹⁷

Some studies have proved that EV, as a new nano-scale platform, can be used to carry drugs and treat related diseases. In jing's experiment, they built a nanomedicine RGD-NPVs@MNPs/DOX developed by encapsulating melanin nanoparticles (MNPs) and doxorubicin (DOX) inside RGD peptide (c (RGDyC))-modified nanoscale platelet vesicles (RGD-NPVs). With the help of platelet vesicles, nanoparticles evaded immune clearance and targeted tumor cells, and finally activated and unloaded drugs under laser irradiation. The results showed that the nanoparticles enhanced the effect of radiotherapy and effectively killed tumor cells.¹⁰⁶

Unfortunately, the research on anti-tumor drugs carried by PMP is still lack of focus, and the research of PMP in the development of anti-tumor drugs needs to be further explored.

Nanoparticles with Multiple Excitation Modes

In addition, nanoparticles combining multiple excitation modalities have also shown better results in experiments. These nanoparticles often provide two or more excitation modes, which can significantly increase the payload of the drug and unload the drug through external stimulation and the internal physical or chemical characteristics of the tumor. So as to improve the drug release efficiency and anti-tumor effect. However, due to the complexity of the tumor environment, a variety of excitation modes may antagonize each other and affect the drug release. For example, the anoxic environment caused by laser excitation will significantly reduce the drug release.¹¹⁸ Therefore, the nanoparticles bound by a variety of stimuli must be combined in an appropriate way.

For platelet-targeting nanoparticles, Yue et al conducted a study in which they utilized C3N4 nanosheets (CN) doped with the two-dimensional nanomaterial FeIII. These nanosheets were loaded with both chlorogenic acid (CA) and DOX to synthesize a pH/H₂O₂-responsive nanocomplex. As a result, the drugs exhibit more potent anti-tumor effects in the U14 tumor-bearing mice model.⁷⁰ Wang's team also prepared an 'ion/gas' bioactive nanogenerator (called IGBN) comprising a copper-based MOF and a loaded cisplatin-arginine (Pt-Arg) prodrug, which is a pH/GSH-responsive complex. The acidic pH in the tumor microenvironment breaks down the MOF, releasing Pt-Arg. Subsequently, GSH dissociates Pt-Arg to release active cisplatin and arginine (Arg). Arginine significantly enhances H₂O₂ production within the tumor and undergoes oxidation facilitated by H₂O₂ self-supply, resulting in the production of a self-enhancing cascade of NO gas. Ultimately, NO effectively inhibits platelet activation within the tumor in a 4T1 mammary adenocarcinoma model.⁷³

For platelet-membrane-coated nanoparticles, Zhang's team developed a photothermal/acid-responsive PDA@Dox NPs complex, which exhibited outstanding therapeutic effects and effectively prevented tumor recurrence in a breast cancer mouse model.¹⁰² Zhou developed an ultrasound and GSH-responsive MM-PEG@PLTs complex. The results demonstrated that ultrasound and GSH effectively triggered the controlled release of the contents, leading to tumor amino acid starvation, mTOR inhibition, and iron death. These effects significantly enhanced the anti-tumor response in breast cancer.¹⁰³ Figure 2 illustrates the response modes of the mentioned above drugs.

Limitation and Perspective

These nanoparticles use the targeted effect of platelets on the tumor to aggregate at the tumor site and achieve site-specific drug release through specific stimulation. This strategy enhances drug targeting while minimizing systemic toxicity. However, their shortcomings are also obvious.

First of all, there are still great challenges in large-scale experiment and promotion of human-derived platelets because of ethical and other problems. At present, the storage of platelets also poses a challenge to the experiment. Platelets stored at room temperature often face the risk of pollution, which leads to a short shelf life, and even the latest

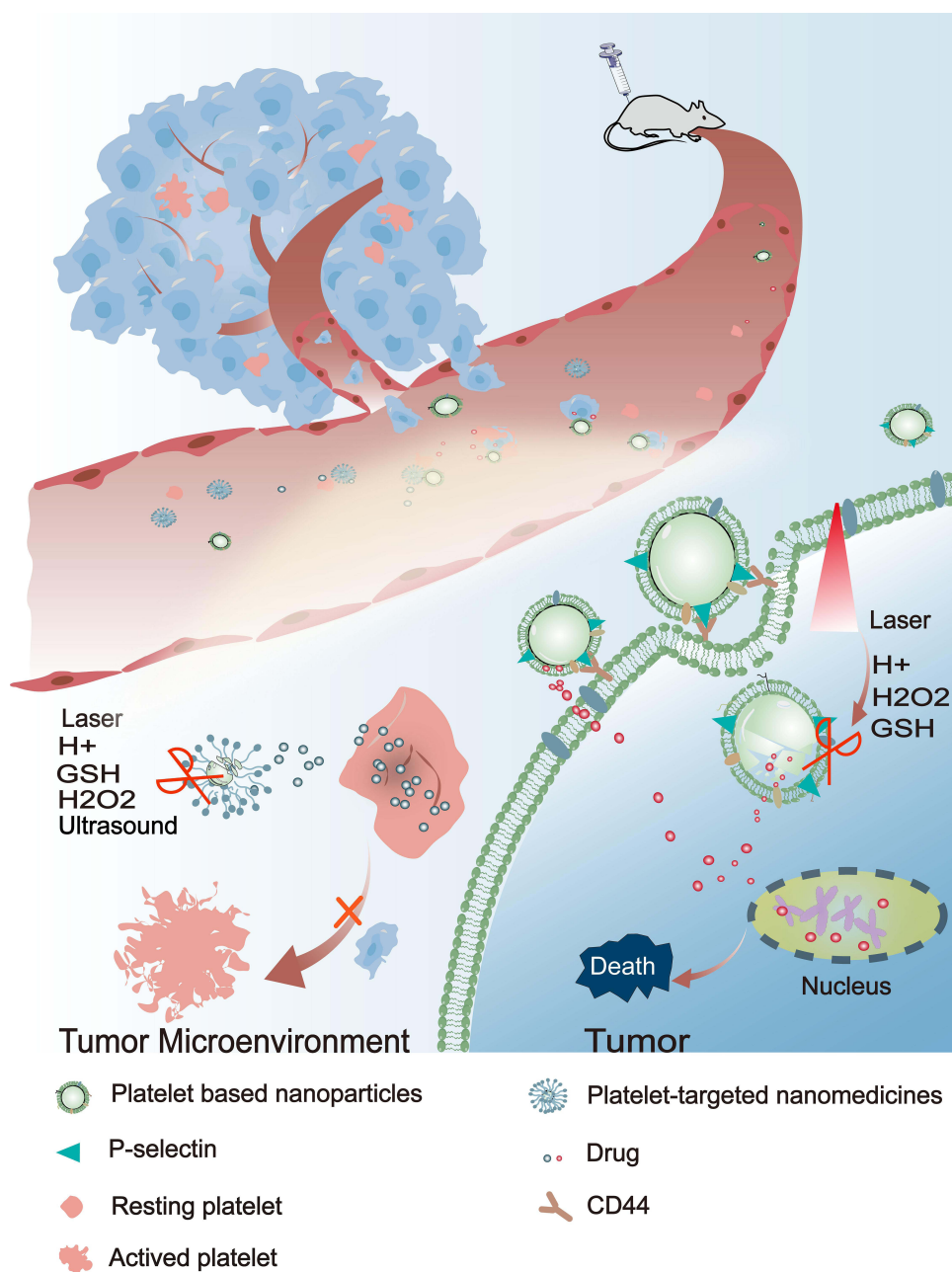


Figure 2 Nanoparticles enter the blood circulation and actively target tumor-related platelets or tumor cells. Platelet targeting nanoparticles target platelets with their surface modifiers and release loaded drugs under the conditions of GSH, H⁺, hydrogen peroxide, ultrasound and laser, and inhibit platelet activation by covering platelet surface markers. Platelet-based nanoparticles mainly use P-selectin on platelet membrane to recognize CD44 targeting tumor on the surface of tumor cells, and enter tumor cells through tumor endocytosis, and then release drugs under the conditions of GSH, H⁺, hydrogen peroxide, laser and so on. In addition, platelet-based nanoparticles can also be directly activated by tumor cells and release drugs.

pathogen technology only delays the shelf life to about 7 days.¹¹⁹ At the same time, as a highly dynamic cell, platelets usually communicate with their environment (for example, tumor educated platelets), and the obtained platelets often have certain donor characteristics.¹²⁰ Therefore, whether the engineered platelets prepared by non-autologous platelets have other effects on tumors is a question that needs to be verified.

Not only that, the current research on these nanoparticles is still based on mouse experiments, the lack of clinical trials means a lot of unknown possibilities because of differences between mice and human body. It requires more clinical trials to verify whether these specially designed nanoparticles can work smoothly in the human body. In the existing experiments, these kinds of nanoparticles need sufficient membrane coverage, which will lead to the recognition

and removal of nanoparticles by the immune system, and excessive clearance may lead to systemic damage due to the premature release of drugs. This puts forward higher requirements for the preparation of nanoparticles. In addition, the design of these drugs is often complex and does not improve efficacy, posing challenges for drug production and storage. Further optimization is necessary to promote their clinical translation. Although the targeting ability of the drug has been improved, experimental evidence shows that it accumulates in organs such as liver and kidney, indicating possible hepatotoxicity. Last but not least, the strategies of radiotherapy and chemotherapy are different in different stages of cancer, so further experiments are needed to determine the appropriate doses of drugs at different stages of cancer to ensure the effectiveness and safety of these drugs.

At present, artificial synthetic platelets have been paid attention to, and great progress has been made in simulated platelet adhesion. These analogues can prolong storage life and minimize side effects, synthetic platelets are more controllable, and the problems of platelet supply are also solved. The use of synthetic platelet analogues to carry drugs may be a way out for future development.¹¹¹

Conclusion

Platelets play an important role in tumor progression, and anti-tumor drugs inspired by platelets have become an important direction of cancer drug development in the future. In this review, researchers have worked hard to design a variety of platelet-based nanoparticles, and these strategies have proved their ability in experiments, and have good performance on the natural targeting, effectiveness and safety of tumor cells. With the development of intelligent delivery, personalized treatment for cancer will become possible, and these nanoparticles will greatly enhance the specificity of anti-tumor therapy. Accordingly, these nanoparticles can also be modified for the diagnosis of tumor cells. These nanoparticles show great potential in anti-tumor and will play a great role in clinical treatment in the future, but due to the lack of clinical trials to further prove their performance, more efforts are needed for their application in the future.

Data Sharing Statement

Data sharing not applicable to this article as no datasets were generated or analyzed during the current study.

Author Contributions

All authors made a significant contribution to the work reported, whether that is in the conception, study design, execution, acquisition of data, analysis and interpretation, or in all these areas; took part in drafting, revising or critically reviewing the article; gave final approval of the version to be published; have agreed on the journal to which the article has been submitted; and agree to be accountable for all aspects of the work.

Funding

This research was funded by the Sichuan Provincial Cadre Health Research Project (Chuan Gan Yan 2022-802); the Sichuan Science and Technology Program, China (2021JDRC0152, 2022YFS0006, 23NSFSC3334), Sichuan Province Medical Association Research Project (S21021).

Disclosure

The authors declare no competing interests in this work.

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