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Bioresponsive drug delivery systems for the treatment of inflammatory diseases

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ABSTRACT

Inflammation is intimately related to the pathogenesis of numerous acute and chronic diseases like cardiovascular disease, inflammatory bowel disease, rheumatoid arthritis, and neurodegenerative diseases. Therefore anti-inflammatory therapy is a very promising strategy for the prevention and treatment of these inflammatory diseases. To overcome the shortcomings of existing anti-inflammatory agents and their traditional formulations, such as nonspecific tissue distribution and uncontrolled drug release, bioresponsive drug delivery systems have received much attention in recent years. In this review, we first provide a brief introduction of the pathogenesis of inflammation, with an emphasis on representative inflammatory cells and mediators in inflammatory microenvironments that serve as pathological fundamentals for rational design of bioresponsive carriers. Then we discuss different materials and delivery systems responsive to inflammation-associated biochemical signals, such as pH, reactive oxygen species, and specific enzymes. Also, applications of various bioresponsive drug delivery systems in the treatment of typical acute and chronic inflammatory diseases are described. Finally, crucial challenges in the future development and clinical translation of bioresponsive anti-inflammatory drug delivery systems are highlighted.

1. Introduction

Inflammation is an essential immune response to harmful stimuli, such as pathogens, damaged cells, chemical irritants, physical trauma, burns, radiation, and frostbite [1]. Whereas inflammation is essential for repair, remodeling, and even renewal of different tissues under a variety of noxious conditions [2], inflammatory responses need to be spatiotemporally regulated to maintain tissue or systemic homeostasis. Uncontrolled or unresolved inflammation is intimately linked to the initiation and progression of numerous acute and chronic diseases, such as acute liver/lung injury, acute bronchitis, asthma, inflammatory bowel disease (IBD), rheumatoid arthritis (RA), atherosclerosis, and neurodegenerative diseases [3–6]. Also, inflammation is a key feature of metabolic disorders such as obesity and diabetes [7,8]. Consequently, anti-inflammation is a highly promising strategy for the prevention and treatment of acute and chronic inflammatory diseases as well as diseases associated with inflammation [5,9,10].

Historically, small molecule agents, such as glucocorticoids, non-steroidal anti-inflammatory drugs, and antioxidants have been extensively used for the treatment of different inflammatory diseases [11–13]. Despite their effectiveness to varied degrees, these drugs show undesirable side effects such as osteoporosis [14], aseptic joint necrosis [15], gastrointestinal bleeding [16,17], liver/kidney injury [18], and cardiovascular risk [19]. By regulating molecular and cellular events closely associated with inflammatory responses, monoclonal antibodies to various pro-inflammatory cytokines and chemokines or other biomacromolecules have been clinically used or pre-clinically examined for therapy of RA [20], IBD [21,22], atherosclerotic disease [23,24], and asthma [25]. However, clinical applications of these biological agents have been considerably impaired by their limitations and adverse effects, such as high cost [26], a primary or secondary non-response [27], and the increased risk of severe infections and malignancies [28–30]. To a large degree, systemic distribution in non-target tissues and cells is mainly responsible for undesirable effects, regardless
of small molecule drugs and biological therapies. Also, site-specific delivery and controlled release in inflammatory tissues/cells is critical to some newly discovered therapies, such as specialized pro-resolving mediators that have been found highly effective in the treatment of inflammatory disorders, owing to their potent activity of resolving inflammation [10,31,32].

To circumvent disadvantages of anti-inflammatory therapies based on traditional formulations, different advanced drug delivery systems have been investigated over the past decades. In this aspect, dendrimers, lipid nanoparticles, polymeric nanoparticles, micelles, nanocapsules, nanoeomulsions, nanofibers, microparticles, hydrogels, and multi-compartmental delivery systems were examined for delivery of either molecular therapies or therapeutic cells for the treatment of inflammatory diseases [33–38]. In addition to the totally synthesized vehicles across multiple length scales, some biomimetic particles such as exosomes, cell membrane-based nanovehicles, and cellular carriers have been developed for delivery of anti-inflammatory agents [39–42]. The majority of these delivery systems were employed for treating inflammation in specific tissues by local delivery of various therapeutics [33–35,43–46]. In other cases, nanoparticles have been extensively investigated for site-specific delivery of therapeutic cargos to inflammatory sites after systemic administration [34,37,39,41,42,47–53], since inflamed tissues generally show enhanced vascular permeability [54,55]. In addition, surface engineering with either molecular moieties or cell-derived membranes have been conducted for different nanovehicles to increase their targeting capability [41,47–52,56], in view of the fact that specific biomarkers are overexpressed in inflammatory tissues and/or cells. Furthermore, macroscale biomaterials, such as hydrogels were examined for delivery of different cells to achieve cell therapy or tissue repair at inflammatory sites [57–59].

Recently, increasing evidence has demonstrated the advantages of bioresponsive delivery systems in the treatment of a diverse array of diseases, such as cancers, metabolic syndrome, and autoimmune/inflammatory diseases [33,40,60–66]. By more precisely regulating the cargo release behaviors at the sites of interest, bioresponsive carriers are able to considerably potentiate efficacies of different therapeutics and simultaneously reduce their side effects. In particular, biomaterials sensitive to various biochemical signals like abnormally changed pH, reactive oxygen species (ROS), and enzymes in diseased sites have been studied as stimuli-responsive delivery systems for targeted treatment of inflammatory diseases. Herein we provide a state-of-the-art review on bioresponsive drug delivery systems for the management of inflammatory diseases.

2. Inflammation and the inflammatory microenvironment

Inflammation is a biological self-response to endogenous and/or exogenous biophysicochemical stimuli, which is characterized by orchestration of a diverse array of molecular mediators and inflammatory cells (Fig. 1). According to the duration of inflammation, it can be classified into acute and chronic inflammatory responses. Acute inflammation, with the cardinal clinical signs of redness, heat, swelling, and pain [67], frequently lasts for several hours to days. Responding to invading pathogens, tissue injuries, or other noxious stimuli, acute inflammation is generally accompanied by rapid and dramatic accumulation of fluid, chemokines, cytokines, and inflammatory cells (particularly neutrophils) in the involved tissues/organs [68,69]. The acute inflammatory response is usually self-limiting, which can be rapidly resolved to restore tissue homeostasis. However, failure to remove a pathological insult or inability to realize resolution will lead to persistent tissue injury and a sustained inflammatory response for months to years, termed as chronic inflammation. Monocytes/macrophages are the primary contributors to the chronic inflammatory process by releasing a large number of inflammatory cytokines, which are intimately related to the pathogenesis of numerous chronic diseases. Although the initiation and progression of inflammation largely depends on types of stimuli and degrees of the body response, its common pathological and physiological features are intriguing targets for diagnosis and therapy of numerous inflammatory diseases. In the following section, we briefly introduce typical pathophysiological characteristics of the inflammatory microenvironment, which are informative for rational design of bioresponsive materials and drug delivery systems for the management of inflammation-associated diseases.

2.1. Increased vascular permeability and infiltration of inflammatory cells

The most significant characteristic in the early stage of inflammation is the altered anatomy and function of the microvasculature, resulting in vasodilation, increased vascular permeability, and tissue leukocytosis [55]. The immediate responses are dominated by plasma- and cell-derived vasoactive mediators, such as fibrin, kinins, complement components (like C3a and C5a), histamine, serotonin, platelet-activating factor, and prostaglandins. These mediators can bind specific receptors on vascular endothelial cells, causing their contraction and gap formation, thereby leading to increased vascular permeability and extravasation of intravascular fluids within 30 minutes after injury. Meanwhile, these inflammatory mediators activate endothelial cells to increase the expression of different cell adhesion molecules, largely classified into four molecular families, i.e., selectins, cadherins, integrins, and immunoglobulins. These adhesion molecules expressed on endothelial cells then mediate extravasation of leukocytes from the bloodstream into the injured tissues via a precisely controlled process with sequential tethering, rolling, firm adhesion, and transmigration.

Different leukocyte subpopulations are involved in inflammatory reactions, mainly including neutrophils, eosinophils, basophils, lymphocytes, monocytes/macrophages, and mast cells. Particularly, as the most abundant leukocytes in the circulation (50–70%), neutrophils are the major cellular participant in acute inflammatory response, by carrying out their diverse functions, such as phagocytosis and digestion of pathogens, production of ROS, formation of neutrophil extracellular traps, as well as activation and regulation of innate and adaptive immunity [70,71]. In addition, neutrophils actively participate in chronic inflammatory conditions like chronic obstructive pulmonary disease [72], arthritis [73], IBD [74], and atherosclerosis [75]. Consequently, neutrophils have emerged as an exciting target for both imaging and therapeutic intervention of diseases associated with inflammation [76–79]. Depending on the degree and persistence of injury as well as activation of different mediators, monocytes can migrate into the inflamed sites and differentiate into macrophages. The tissue-resident macrophages, together with the infiltrated macrophages and their cytokines play a key role in host defense and maintaining tissue homeostasis in the acute phase response of inflammation. Furthermore, sustained and uncontrolled macrophage responses in affected tissues are central to prolonged inflammatory and immune reactions, generally leading to chronic inflammation that is associated with a variety of infectious and noninfectious diseases, such as chronic hepatitis and atherosclerosis [4,5,80]. Macrophages have therefore been investigated as promising targets for imaging and treatment of various chronic inflammatory diseases [5,52,81]. As for mast cells, basophils, and eosinophils, they are important players in allergy, hypersensitivity, asthmatic responses, and parasite-associated inflammatory reactions [82–84]. Accordingly, these cells are interesting therapeutic targets for the related inflammatory diseases, despite their relatively low numbers in the circulation and local tissues. In addition to functioning as therapeutic targets, these different effector cells involved in inflammation can engulf different materials, followed by migration into inflammatory sites, making them interesting “Trojan Horses” for targeted delivery of different molecular and particulate therapeutics to inflamed sites [56,85,86].
2.2. Acidosis at inflammatory sites

Local acidification has been confirmed at inflamed sites of various diseases associated with acute and chronic inflammation, such as the peritoneum of peritonitis [87], injured arteries [88], ischemic sites [89,90], atherosclerotic plaques [91], asthmatic airways [92], rheumatic or gouty joints [93], and cerebrospinal fluid in the brain with meningitis [94]. Compared to healthy tissues, the inflamed areas generally display lower pH values. For instance, extracellular pH in the location of myocardial ischemia is approximately 6.5-6.0 [90], while pH as low as 4.2 was reported in the fracture microenvironment [95]. For patients with acute asthma, their exhaled vapor condensate showed pH 5.2, in contrast to pH 7.7 in control healthy subjects [96]. Normal synovial fluid exhibits pH 7.4-7.8, while synovial fluid in arthritic joints decreased to 6.6-7.2 [93,97,98], which is related to the intensity and state of inflammation and disease activity.

This acidosis feature (pH < 7.35) has been considered relevant to increased energy and oxygen consumption by infiltrated inflammatory cells, which shifts metabolism from aerobic to anaerobic styles in the inflamed tissues, leading to oxygen-independent glycolysis and increased lactic acid production (Fig. 2A) [99]. In the case of infection, bacterial metabolites like lactate may also contribute to acidification in the interstitial spaces [94,100]. In addition to metabolic acidosis, respiratory acidosis in inflammatory lung diseases (particularly asthma and chronic obstructive pulmonary disease) can be resulted from accidently inhaled acids such as those in fogs and air pollution [92]. The long-term exposure to acids is an important risk factor in the pathogenesis of asthma and bronchitis, which in turn leads to amplified acidosis due to altered airway pH homeostasis. By serving as an endogenous danger signal and regulating the synthesis and release of inflammatory mediators, extracellular acidosis itself can alter biological functions of different inflammatory and immune cells [99,101,102]. Whereas pH values in different inflammatory tissues were determined and compared to the corresponding normal tissues, few studies have been performed to quantify the degree of acidosis during the progression of specific inflammatory diseases. Nevertheless, some available findings revealed a first pH decrease upon the initiation of inflammation, followed by approximately sustained low pH values once inflammation persists [94]. In addition, acidosis can be alleviated by drug therapy [96]. Therefore targeting pH homeostasis might be a new therapeutic strategy to curb different inflammatory diseases [92,99,103]. Meanwhile, acidosis can be used as a very useful endogenous cue for designing delivery systems with acid-triggerable release behaviors for different therapeutics.

It should be noted that, besides these pathological conditions, varied degrees of acidic environments exist in specific organs, tissues, and cells of the human body under normal physiological conditions. For instance, the pH value along the gastrointestinal tract has a well-controlled gradient, ranging from 1.0-3.5 in the stomach to 5.5-7.5 in the intestine [104]. The normal pH range in the vagina is 3.8-4.5. In addition, subcellular compartments show different acidic microenvironments, with pH 5.0-6.5 in early and late endosomes, pH 4.5-5.0 in lysosomes, and...
ROS are chemically reactive species containing oxygen. Typical ROS include hydrogen peroxide (H$_2$O$_2$), superoxide anion radical (•O$_2^−$), hydroxyl radical (•OH), singlet oxygen (¹O$_2$), and hypochlorous acid (HOCl). Physiological levels of ROS play an important role in preserving and controlling cellular functions, such as cell differentiation, proliferation, migration, and survival. In addition, ROS are crucial bactericidal molecules for the host defense against various pathogens. However, excessive ROS, generally produced in cellular organelles such as mitochondria, peroxisomes, and endoplasmic reticulum of activated neutrophils and macrophages under inflammatory conditions (Fig. 2B), can cause oxidation of DNA, proteins, and lipids, thereby causing cell death by necrosis and/or apoptosis [108].

The ROS-mediated oxidative damage is closely related to the pathogenesis of numerous diseases associated with acute and chronic inflammation, such as acute ischemic stroke, acute liver/lung injury, acute bronchitis, acute gastroenteritis, neurodegenerative diseases, cardiovascular diseases, asthma, diabetes, IBD, and RA. In patients with IBD or Helicobacter pylori infections, their mucosal levels of ROS are 10-100 folds higher than those of the healthy population [109,110]. The injured artery of rats showed an average ROS level approximately 10 times higher than that of the normal controls [111]. In some brain diseases, the extracellular H$_2$O$_2$ concentration increased to 100 μM during ischemia and reperfusion, which caused modification of neuronal proteins and structural components as well as loss of cognitive function, eventually resulting in the occurrence of Alzheimer’s and Parkinson’s disease [112]. In inflammatory lung diseases, respiratory lining cells experienced 20-fold higher H$_2$O$_2$ levels than those in the healthy tissues [113]. Studies in animal models of acute lung injury, alcoholic liver injury, and colitis demonstrated that the ROS level generally exhibits a parabolic profile during the development of different inflammatory diseases, with the peak time dominating by the disease types and endogenous or exogenous inflammatory stimuli [77–79]. Consequently, the significantly increased oxidative stress in inflammatory sites is another endogenous biochemical trigger for targeted therapy of inflammatory diseases.

2.4. Overexpressed enzymes and other features under inflammatory conditions

In inflammatory diseases, a diverse array of enzymes will be up-regulated. During acute inflammation, different enzymes required for degradation of pathogens or foreign materials are generated in cytoplasmic granules of polymorphonuclear leukocytes, particularly neutrophils. For instance, lysozyme, myeloperoxidase (MPO), collagenase, elastase, and gelatinase can be released by degranulation of neutrophils [114,115], while acid hydrolases, elastases, cathepsins, and matrix metalloproteases (MMPs) will be secreted by activated monocyes/macrophages in both acute and chronic inflammatory responses [116]. The increased expression of some enzymes such as cyclooxygenase-2 directly mediates inflammation [117], and therefore being used as important therapeutic targets for the management of inflammatory disorders. In the case of infection, bacteria produce some specific enzymes like urease or azoreductase [118]. Regardless of their different sources, the inflammation-associated enzymes frequently display a notable high expression at the initial stage of inflammatory diseases [77–79,119–121], which then decrease to a certain degree and maintain abnormally high levels during the whole process of inflammation.
and hard materials [137]. For POE-based delivery systems, such as wafers and microspheres, their erosion and drug release periods may be facilely regulated from a few days to many months. Initially, different forms of POEs were fabricated to deliver small molecule drugs for the treatment of post-surgical pain, osteoarthritis, and ophthalmic disease [138], which were used later as carriers for peptides, proteins, and nucleic acids [137,139,140]. Also, block copolymers based on POEs and poly(ethylene glycol) (PEG) were synthesized and used as nanoparticles or micelles for targeted drug delivery [141].

Whereas POEs show desirable pH-sensitivity and good
Materials with hydrazone linkages have been widely used in pharmaceutical sciences due to their excellent pH-sensitivity as well as synthetic flexibility and diversity [133,185,186]. Similar to other acid-labile linkages, hydrazones are generally employed to synthesize both prodrugs and delivery carriers to achieve programmed drug release in response to the intracellular or extracellular acidic microenvironment. As a decent example, Kataoka’s group synthesized an amphiphilic block copolymer in which doxorubicin was conjugated onto the polysaccharide block via hydrazone [187]. Polymeric micelles assembled by this pH-sensitive copolymer showed notable pH-dependent release profiles. By the similar strategy, other small molecule drugs, such as dexamethasone, prednisolone, isoniazid, and all-trans retinal, were conjugated onto hydrophilic polymers or inorganic nanoparticles via the hydrazone bond to construct pH-sensitive drug-polymer conjugates or produg nanoparticles with on-demand therapeutic release under different inflammatory conditions [188–193]. As compared to pH-sensitive linkers of ortho esters, acetals, or ketals, the hydrazone bond is more useful to develop acid-triggerable produg systems.

On the other hand, pH-sensitive hydrazone linkages have been utilized to synthesize different pharmaceutical materials, enabling the development of diverse smart delivery systems, such as liposomes, micelles, nanoparticles, and hydrogels [194–197]. Of note, PEG conjugates with hydrazone linkers are frequently used to fabricate long-circulating nanocarriers with acid-responsive sheddable coatings [198], which are particularly intriguing for intracellular delivery of nucleic acids. Nevertheless, these delivery platforms generally show limited control over pH-sensitive drug release, due to the presence of passive diffusion, as compared to hydrazone-derived produg nanoparticles and other vehicles with acid-labile matrices.

Materials bearing acetics or ketals

Acetals/ketals are frequently used as protective groups for aldehydes and ketones in organic synthesis [149]. In drug delivery fields, both acetics and ketals were initially used as acid-cleavable agents to develop prodrugs derived from either small molecule drugs or proteins [150,151]. Studies by Fréchet’s group suggested that acetals can function as pH-sensitive linkages to fabricate diverse drug delivery systems, such as polymer-drug conjugates, micelles, nanoparticles, and microgels [152–154]. A similar approach was adopted by other researchers to develop pH-sensitive polymeric nanogels or hydrogels [155,156]. Of note, the hydrolysis rate of resulting systems at different pH values can be easily tailored by modulating the types of alcohol and carbonyl (aldehyde and ketone). In other studies, acetals/ketals moieties were embedded in the main chain of hydrophobic polymers to synthesize fully acid-degradable biocompatible materials that can be further processed into either nanoparticles or microparticles for site-specific delivery of different therapeutic agents [157–162].

Fréchet et al. also reported facile synthesis of acid-responsive materials by acetalation of dextran using 2-methoxypropene or other functional molecules [163–165]. Thus obtained acetalated dextran polymers have been examined for encapsulation and pH-triggerable release of small molecules, nucleotides, and proteins in the forms of nanoparticles or microparticles [164,166–169]. Our group demonstrated that kinetically controlled acetalation can be carried out for cyclic oligosaccharides, such as α-CD, β-CD, and their polymers, thereby affording a series of acid-labile biocompatible materials [170,171]. By regulating acetalation time, the pH-sensitive hydrolysis rate of resulting CD materials may be easily modulated. Notably, systemic in vivo studies suggested that acetalated CD materials exhibited good safety profile after administration by subcutaneous, intramuscular, intraperitoneal, or intravenous (i.v.) injection in mice, rats, and rabbits [170–173]. In other cases, acetals were introduced as acid-labile linkages in hyperbranched polymers and dendrimers to construct different delivery systems [174–177]. Also, linear and cyclic acetics can be incorporated in the side chains of polymers to develop acid-sensitive polymers or polymer-drug conjugates [178–180]. In addition to their excellent acid-sensitivity and facile synthesis procedures, most acetal- and/or ketal-derived pH-responsive materials do not generate acidic byproducts after hydrolysis, thereby avoiding significant proinflammatory effects [171,181,182]. By contrast, frequently used polyesters, such as PLA and poly(lactide-co-glycolide) (PLGA), can induce local inflammation in different tissues [183,184]. For future translation, however, in vivo safety of different types of acetal/ketal-derived materials and delivery systems need to be intensively examined in various animal models.

Materials with hydrazone linkages

Hydrazone linkages have been used in pharmaceutical sciences to design prodrugs with hydrazone linkages. For example, ortho esters were used to prepare pH-responsive lipid conjugates for liposome formulations [142]. Pendent cyclic ortho esters were introduced in hydrophilic polymers to prepare acid-labile carrier materials for pH-triggerable drug delivery [143,144]. In addition, ortho esters have been incorporated in the polymer main chains or used as cross-linkers to construct pH-responsive nanovehicles [145,146]. By chemical functionalization of β-cyclodextrin (β-CD) with ortho esters, pH-sensitive CD derivatives were obtained, which were fabricated to pH-sensitive supramolecular nanoparticles and polysuperoxotaxanes for drug delivery [147,148]. Nevertheless, for these ortho-ester-containing materials, their in vivo safety profiles remain to be addressed by comprehensive in vivo studies.

Materials with other acid-labile linkages

To prepare pH-sensitive delivery systems, β-thiopropionate, phosphoramidate, vinyl ether, boronate, and silyl ether were also used as acid-sensitive linkages. The acid-labile β-thiopropionate linkage was employed by Kataoka and colleagues to synthesize conjugates of PEGylated antisense oligonucleotides or siRNA, which were further assembled into polycion complex micelles for intracellular gene silencing [199]. Similarly, oligonucleotides can be covalently conjugated with PEG via an acid-cleavable phosphoramidate linkage, and the obtained conjugate is able to complex with a cationic peptide to form polyelectrolyte complex micelles for efficient intracellular delivery [200]. Likewise, phosphoramidate linkers were used to synthesize nanoformulated polymeric produgs for antiretroviral therapy [201,202]. In other cases, PEGylated dioleoylphosphatidylethanolamine (DOPE) conjugate with an acid-labile vinyl ether linker was used to prepare liposomes or nanoassemblies to achieve pH-triggerable drug release or gene delivery [203,204]. Also, boronate and silyl ether were used as cross-linkers to synthesize acid-sensitive biomaterials in the forms of nanoparticles, macroscale particles, suture, or stents [205,206]. For these delivery vehicles, however, only a few studies have been conducted to investigate their pH-responsive drug release performance.

Different from the above mentioned pH-sensitive materials based on acid-labile linkages, Gao’s group designed and developed a library of tunable, ultrasensitive pH-activatable micelles self-assembled by ionizable block copolymers, in which tertiary amines serve as precisely controlled hydrophobic substituents in the ionizable hydrophobic blocks [207,208]. This type of micellar nanoparticles exhibited fast and ultrasensitive response to changes over the entire physiological range of pH, depending on the types of tertiary amine substituents. In addition to functioning as ultra-pH-sensitive fluorescent nanoprobe, these nanoparticles can be used for developing pH-activatable nanotherapies and nanovaccines [209,210].

ROS-responsive materials and delivery systems

In view of overproduced ROS at different diseased sites, considerable advances have been achieved in recent years in the development of
Table 2
ROS-responsive materials for on-demand drug delivery [355-359].

| Linkages         | Chemical structures of typical materials | Formulation forms | Release mechanisms                                                                 | Evaluation models          | References        |
|------------------|-------------------------------------------|-------------------|------------------------------------------------------------------------------------|-----------------------------|-------------------|
| Boronic ester    | ![Chemical structure of Ox-DEx](image)     | Nanoparticles and microparticles | Passive diffusion and ROS-triggered carrier hydrolysis | In vitro in CD8+T and PC-12 cells; in vivo in mice/rats | [62, 213, 214, 355] |
|                  | ![Chemical structure of Ox-bCD](image)    | Nanoparticles     | Passive diffusion and ROS-responsive hydrolysis of nanoparticles                     | In vitro in RAW264.7, MOVAS, B16F10, and MDA-MB-231 cells; in vivo in mice/rats | [111, 182, 214, 215, 298, 319, 320, 325, 329] |
|                  | ![Chemical structure of TPCD](image)      | Nanoparticles and polyplexes | Passive diffusion, ROS-triggered material hydrolysis, and dissociation of polyelectrolyte complexes | In vitro in A549 cells and neutrophils; in vivo in mice | [218, 219] |
|                  | ![Chemical structure of 4CBE](image)      | Micelles and nanoparticles, | Passive diffusion and ROS-triggered hydrolysis of delivery vehicles | In vitro in RAW264.7 cells; in vivo in mice | [223-229] |
| Polyoxalate      | ![Chemical structure of HPOX](image)      | Nanoparticles and microparticles | Passive diffusion and ROS-responsive hydrolysis of nanoparticles | In vitro in RAW264.7 cells; in vivo in mice | [223-229] |
| Thioketal        | ![Chemical structure of PPADT](image)     | Polyplexes, nanoparticles, microspheres, and fibrous patches | Passive diffusion, ROS triggered hydrolysis, and/or dissociation of polyelectrolyte complexes | In vitro in lymphocytes as well as MDA-MB-231, PC, HeLa, KB, and RAW264.7 cells; in vivo in mice/rats | [231-235, 239] |
| Thioether        | ![Chemical structure of PPS](image)       | Micelles, nanoparticles, microspheres, and hydrogels | Passive diffusion and ROS-triggered disassembly of delivery vehicles | In vitro in RAW264.7, dendritic, CD8+T, and LR-3T3s cells; in vivo in mice/rats | [242-250, 356-358] |
| Selenium/tellurium | ![Chemical structure of PEG-PSSe-PEG](image) | Micelles | Passive diffusion and ROS-triggered disassembly of micelles | In vitro in L-02 and HepG2 cells | [251-253] |
|                  | ![Chemical structure of PEG-PUTE-PSSe-PEG](image) | Micelles | Passive diffusion and ROS-triggered disassembly of micelles | In vitro in L-02 and HepG2 cells | [251-253] |

ROS-responsive materials and delivery systems for on-demand drug release and targeted therapy (Table 2). In this section, we introduce typical ROS-responsive materials and delivery platforms.

3.2.1. Boronic ester-containing materials

In organic synthesis, boronic acids and their derivatives are frequently used as good protective groups for diols. Since the linkages between boronic esters and the conjugated molecules can be easily cleaved under oxidative conditions, they have been extensively employed for the development of ROS-responsive fluorescent probes, prodrugs, and pharmaceutical materials [62, 211–214]. An early study by Fréchet and coworkers demonstrated successful synthesis of ROS-responsive materials by covalently conjugating an oxidation-labile unit of 4-(hydroxymethyl)phenylboronic acid pinacol ester (PBAP) onto...
dextran, and the functionalized polymer can be further produced into microparticles for ROS-triggerable therapeutic delivery [213]. Our studies showed that this functionalization strategy can be conducted for cyclic oligosaccharides (such as β-CD) [214,215], thereby offering ROS-responsive carrier materials. Notably, H$_2$O$_2$-sensitivity of resulting materials is regulatable by changing the content of PBAP or the linker type between PBAP and β-CD [215]. At a comparable content of PBAP, while materials containing either carbonate or carbamate linkages showed rapid hydrolysis in the presence of H$_2$O$_2$, the ester bond-based materials displayed considerably low H$_2$O$_2$-sensitivity. Regardless of their chemical structures, all these materials are able to be processed into nanoparticles with tunable sizes by a nanoprecipitation/self-assembly method. Furthermore, both in vitro and in vivo evaluations by our group substantiated that this type of ROS-responsive CD materials exhibited good safety profiles [214,215]. Also, phenylboronic acid or its ester was employed to functionalize hydrophilic polymers to achieve ROS-triggerable gene delivery [216,217].

In addition to functionalization of hydrophilic materials, phenylboronic acid pinacol ester was incorporated into the main or side chains of polymers to synthesize ROS-responsive materials for delivery of small molecule therapeutics and efficient gene delivery [218–220]. In other cases, pH/ROS dual-responsive polymers were synthesized by covalently incorporating phenylboronic acid pinacol ester into ortho ester-containing block copolymers [221]. Our recent studies suggested that pH/ROS dual-responsive nanoparticles can be easily constructed by physically combining acid-labile acetalated β-CD and PBAP-conjugated β-CD materials [88]. Of note, for ROS-responsive materials based on the boronic ester strategy, they can be easily obtained by relatively facile and effective synthesis methods, which is a considerable advantage for large-scale production and future translation. Nevertheless, hydrolysis of all the aforementioned boronic ester-containing materials can generate boronic acids that exhibit biological activities to a certain degree, therefore their possible side effects and/or synergistic effects should be taken into account during the treatment of specific diseases.

### 3.2.2. Polyoxalate materials

Polyoxalate-based nanoparticles were initially examined for imaging of H$_2$O$_2$, owing to the nanomolar H$_2$O$_2$-sensitivity of polyoxalate materials and their specificity to H$_2$O$_2$ over other ROS types [222]. Later Lee et al. synthesized polyoxalate materials for H$_2$O$_2$-responsive therapeutic delivery [223]. As a representative example, they synthesized a bioactive polyoxalate material (i.e., HPOX) by a condensation reaction of oxalyl chloride, 1,4-cyclohexamethanol, and p-hydroxybenzyl alcohol (HBA) [224]. In the presence of ROS, HPOX can be triggerably hydrolyzed into three safe compounds, i.e., cyclohexanediol, HBA, and CO$_2$. Of note, HBA is a major bioactive component of Gastrodia elata, a widely used herbal agent for the treatment of inflammation and oxidative stress-related diseases, such as ischemic injury and coronary heart diseases. HPOX can be produced to nanoparticles by an oil-in-water solvent evaporation method, affording nanotherapies with H$_2$O$_2$-activatable hydrolysis and drug release properties as well as antioxidant activity [225,226]. According to the similar procedures, nanoparticles or microparticles derived from a vinyl alcohol-containing copolyoxalate PVAX were developed as H$_2$O$_2$-responsive, biodegradable, and bioactive therapeutic agents for the treatment of acute oxidative injury-related diseases [227–229]. Despite the facile method for synthesis of this type of ROS-responsive materials, it is worth noting that oxalate should be linked with aryl mieties to afford materials with desirable H$_2$O$_2$-sensitivity, while only a very limited number of monomers with good biocompatibility are commercially available.

#### 3.2.3. Thioketal-containing materials

Due to their excellent stability in both acidic and basic conditions, thioketals are also popularly used protecting groups for carbonyl in organic synthesis, which can be efficiently cleaved by ROS [230]. Taking advantage of this unique ROS-labile property, Murthy et al. synthesized a ROS-responsive polymer, i.e., poly(1,4-phenylenecac-tone dimethylene thioketal) (PPADT), by condensation polymerization of 1,4-benzendimethanethiol and 2,2-dimethoxypropane [231]. In the presence of superoxide, PPADT was hydrolyzed into small molecules. It was also found that PPADT nanoparticles were able to triggerably release the loaded fluorescent molecules in activated macrophages. Inspired by this study, different thioketal-containing polymers have been designed and synthesized as ROS-responsive materials for diverse delivery applications. For example, a thioketal polymer was synthesized by step-growth polymerization of (4,4′-bis(mercaptomethyly)biphenyl and 2,2-dimethoxypropane [232]. ROS-responsive microspheres were further prepared using this polymer for inflammation-triggered local delivery of tacrolimus, a powerful immunosuppressant. Gao et al. synthesized ROS-responsive and biodegradable polyurethane by reaction of poly(ε-caprolactone) diol, 1,6-hexamethylene diisocyanate, and a thioketal-containing chain extender, which was fabricated into fibrous patches for therapeutic delivery [233]. Wang’s group reported a thioketal linkage-bridged diblock copolymer of PEG and PLA for on-demand drug delivery [234].

In other studies, the thioketal linkage was introduced into cationic polymers for ROS-responsive gene transfection [235,236]. Also, different therapeutic molecules can be covalently conjugated onto polymers via the thioketal linker to formulate prodrug nanotherapies [237–239]. In addition, by simultaneously incorporating thioketal and disulfide linkages in the backbone of polymers, biodegradable nanopolymers dually responsive to H$_2$O$_2$ and glutathione (GSH) were developed for programmable drug release [240]. It should be emphasized that for thioketal-containing materials and delivery systems, good ROS-sensitivity is only achieved at high levels of H$_2$O$_2$ or in the presence of other strong oxidizing agents.

#### 3.2.4. Materials with thioether linkages and other chalcogen-containing polymers

While the above mentioned ROS-responsive materials are hydrolyzed to realize on-demand drug release, another types of ROS-sensitive materials are based on the hydrophobic to hydrophilic change after oxidation of specific groups. Organic sulfides are frequently used to synthesize ROS-responsive materials and delivery systems, since they can undergo conversion from hydrophobic sulfides to more hydrophilic sulfoxides or sulfones under oxidative conditions. Early in 2004, Hubbell’s group developed oxidation-responsive polymeric vesicles by self-assembly of triblock copolymers consisting of PEG and poly(propylene sulfide) (PPS), which can be easily synthesized by one-pot ring-opening polymerization of episulfide [241]. In the presence of H$_2$O$_2$, the assembled vesicles showed oxidative destabilization, resulting from the increased hydrophilicity due to the oxidation of sulfide moieties in the hydrophobic PPS block into sulfoxides/sulfones.

On the basis of this finding, different types of PPS-derived materials were synthesized to prepare ROS-responsive delivery vehicles. Duvall’s group developed ROS-responsive micelles for triggered drug release, using a copolymer of PPS and poly(N,N-dimethylacrylamide) [242]. The micelles demonstrated efficient drug release in response to H$_2$O$_2$ or endogenous oxidants generated in lipopolysaccharide (LPS)-induced RAW264.7 macrophages. By the living emulsion polymerization of episulfides, cross-linked polysulfide nanoparticles with oxidation-sensitivity were produced [243]. Similarly, Allen et al. fabricated PPS nanoparticles by ring-opening emulsion polymerization using activated pentaerythritol tetraethoxierster as a four-armed star initiator, thereby affording ROS-responsive nanoplatorms capable of loading hydrophobic molecules [244]. In addition, microspheres were directly prepared by the well-established oil-in-water emulsion solvent evaporation method using PPS [245], while hydrogels can be fabricated using PPS-containing copolymers [246]. Nanoparticles, polymericosomes, microparticles, and hydrogels based on PPS-derived biomaterials have been...
extensively investigated as ROS-responsive delivery vehicles for small molecule drugs, proteins, and therapeutic cells, in which therapeutics can be packaged by either physical entrapment or covalent conjugation [245–250].

Other chalcogen-containing polymers have also been examined as bioresponsive materials for drug delivery. In a pioneer study by Zhang and Xu et al., an amphiphilic copolymer (PEG-PUSeSe-PEG) with one block of diselenide-containing polyurethane and two PEG blocks was synthesized [251], which can assemble into micelles in aqueous solutions. Upon stimulation by either oxidants or reductants, PEG-PUSeSe-PEG micelles were disassembled, resulting from a structural dissociation of the diselenide bonds. Following studies suggested that micelles assembled from monoselenide-containing amphiphilic block copolymers also showed oxidation-responsive dissociation and drug release behaviors [252], mainly due to the structural transition of hydrophobic selenide groups into hydrophilic selenoxide or selenone groups. Subsequently, different selenium-containing polymers with diverse structures were synthesized and explored as ROS-responsive materials for biomedical applications [253,254]. Based on the similar design principle, Xu’s group also synthesized tellurium-containing compounds or polymers to construct ROS-responsive assemblies [255]. Nevertheless, similar to thiopektal-derived responsive systems, chalcogen-containing polymers show good sensitivity only at high ROS levels.

### 3.3. Enzyme-responsive materials and delivery systems

Overexpressed enzymes under diseased conditions have long been employed as biological signals for the design and engineering of different bioresponsive materials [256–258]. The responsive mechanisms mainly involve the reduction/oxidation of substrates and the formation/cleavage of chemical bonds in the presence of enzymes. Many enzymes, such as proteases, phosphatases, kinases, and oxidoreductases have been concerned in the development of stimuli-responsive biomaterials. Herein we mainly describe biomaterials sensitive to enzymes overexpressed under inflammatory conditions (Table 3).

Extracellular enzymes like neutrophil elastase, cathepsins, and MMPs are the most frequently studied enzymatic cues in inflammation for designing responsive materials and delivery systems. Aimetti et al. developed a novel enzyme-cleavable hydrogel delivery system based on a neutrophil elastase-sensitive peptide linker (Ala-Ala-Pro-Val), its degradability was demonstrated in the presence of corresponding elastase in vitro [259]. Homma et al. found that Phe-Phe chain can serve as a cathepsin-cleavable peptide linkage to synthesize a polymeric prodrug of methotrexate-conjugated hyaluronic acid, showing effective in vivo efficacy in rats with collagen-induced arthritis [260]. By introducing MMP-cleavable peptide sequences (such as GPOGIAGQ, GPOGIWQ, and GGRMSPMV) into hydrogels as crosslinkers, MMP-sensitive hydrogels based on either synthetic or natural polymers can be synthesized [261–263]. In most recent studies, the peptide PLGLAG linker was used to synthesize MMP-2/9-responsive prodrug nanotherapies [264], nanovehicles [265], and hydrogels [266].

Also, biomaterials and delivery systems responsive to intracellular enzymes, such as furin, protein kinase Ca (PKCα), and caspases have been synthesized. A study by Tang and Gu et al. demonstrated that carriers containing furin-cleavable peptide RVRRSK crosslinkers can be degraded gradually to release cargo protein along their cellular uptake pathway [267]. The same group developed a graphene-based co-delivery system, in which a cell membrane targeting anticancer protein was covalently linked with a furin-cleavable peptide to achieve sequential delivery of different therapeutics [268]. In other studies, polymers containing PKCα-specific peptide substrates were designed and prepared for targeted gene delivery [269], while caspase-3-cleavable polymeric nanocarriers were constructed using a caspase-3 substrate-based peptide linker (such as α-CD and β-CD), MPO-responsive materials can be

| Table 3 | Enzyme-responsive systems for drug delivery. | Sensitive sequences or moieties | Formulation forms | Release mechanisms | Evaluation models | References |
| --- | --- | --- | --- | --- | --- | --- |
| MMPs | GGRMSPMV, GPRQGIAGQ, GPRQGIWQ, GPGQGQ | Nanoparticles and hydrogels | Passive diffusion and enzyme-triggered hydrolysis of delivery | In vitro in B16-F10 cells; in vivo in mice, rats, and pigs | [260] | |
| Neutrophil elastase | KQ(ROX)AAPVRGGGK(QXL) | Hydrogels | Passive diffusion and enzyme-triggered hydrogel hydrolysis | In vitro in HeLa and A549 cells; in vivo in mice | [267,268] | [78,271] |
| Cathepsin D | Phe-Phe | Prodrugs | Enzyme-triggered breakage of peptide linkages | In vitro in HFLS cells; in vivo in mice | [260] | [259] |
| Furin | RVRRSK | Nanocapsules | Passive diffusion and enzyme-triggered hydrolysis of nanocapsules | In vitro in B16 melanoma cells; in vivo in mice | [261–263] | [251] |
| Protein kinase Ca | HFKKQGSFAKKK-NH<sub>2</sub> | Nanoparticles | Passive diffusion and enzyme-triggered hydrolysis of nanoparticles | In vitro in Hela cells; in vivo in mice | [264] | [252] |
| Caspase-3 | AGVA | Nanoparticles | Passive diffusion and enzyme-triggered hydrolysis of nanoparticles | In vitro in HeLa cells; in vivo in Hela cells | [265] | [253] |
| MPO | Luminol | Nanoparticles | ROS- and enzyme-triggered hydrolysis of nanoparticles | In vitro in hydrogel matrices; in vivo in mice | [266] | [254] |
easily synthesized for imaging and therapy of inflammatory diseases [78,79,271]. Whereas enzyme-responsive delivery systems exhibit extremely higher specificity than those responsive to pH or ROS, sensitive peptides are generally required for their preparation, thereby resulting to complicated procedures and high cost in most cases.

3.4. Other bioresponsive materials and delivery systems

Besides the above mentioned biomaterials, other bioresponsive materials have also been explored for anti-inflammatory applications. In this aspect, different redox-sensitive materials and delivery systems have been developed for the treatment of different inflammatory diseases. For example, Xiao et al. synthesized a reducible cationic polymer, by Michael addition between cysteamine bisacrylamide (CBA) and branched polyethyleneimine (PEI, Mw = 800 Da), which was further conjugated with mannosylated PEG. The final material, named as PPM, can complex with tumor necrosis factor (TNF)-α siRNA to obtain redox-sensitive nanoparticles of 211-275 nm for colitis therapy [272]. Kim et al. designed redox-sensitive poly(oligo-L-arginine) (rsPOLA) that can target nanoparticles with endothelial nitric oxide synthase (eNOS) DNA for targeted therapy of atherosclerosis [273]. In the presence of a reducing agent β-mercaptoethanol (β-ME), eNOS/rsPOLA nanoparticles showed considerably increased diameter, resulting from reduced condensation interaction. Recently, Liu and coworkers developed an amphiphilic inulin polymer of 4-aminothiophenol-conjugated carboxylmethylinulin (i.e., ATP-CMI), which could assemble into redox-sensitive nanoparticles of about 210.18 nm [274]. In the presence of GSH, ATP-CMI nanoparticles loaded with budesonide showed notable increased drug release, due to breaking of disulfide bonds and subsequent disassembly of ATP-CMI chains. To further improve the responsive efficiency, a pH/redox-dual responsive polymeric nanoliposome system was designed for targeted delivery of a copper-liganded bioactive complex to the myocardium, resulting in notably improved cardiac function and reduced the infarct size [275].

4. Treatment of inflammatory diseases by bioresponsive drug delivery systems

To overcome limitations of traditional formulations of different anti-inflammatory drugs, bioresponsive drug delivery systems have received much attention for the treatment of inflammatory diseases in recent years. Herein we describe applications of different bioresponsive systems in the management of typical diseases associated with acute or chronic inflammation.

4.1. Therapy of acute inflammatory diseases

4.1.1. Acute cardiovascular diseases

Acute cardiovascular diseases (CVDs), such as acute heart failure, acute myocardial infarction (MI), and acute ischemic stroke, remain one of the leading causes of global death. These fatal diseases are always characterized by acute inflammatory response and elevated oxidative stress that need to be rapidly eliminated in clinical treatment [276,277]. Recently, there has been increasing interest in advanced drug delivery systems for the treatment of acute CVDs [46,49,66,278–281]. Among them, bioresponsive materials and systems show considerable advantages in site-specific delivery and responsive release of different therapeutic agents at diseased tissues [282,283].

Taking advantage of the mild acidic microenvironment of ischemic myocardium, Davis et al. synthesized pH-sensitive poly(cyclohexane-1,4-diy acetone dimethylene ketal) (PCADK), which was further formulated into microparticles (~20 μm) containing a p38 mitogen-activated protein kinase (MAPK) inhibitor SB239063 [181]. Treatment by direct cardiac injection of drug-loaded microparticles (defined as PK-p38) significantly improved cardiac dysfunction following MI, resulting from sustained and controlled release of therapeutics within the damaged myocardium. Notably, PK-p38, showed considerably higher efficacy than free drug and PLGA-based microparticles. Moreover, degradation of PCADK generates neutral and excretable compounds, different from frequently used polyesters such as PLGA that produces proinflammatory acidic byproducts after hydrolysis. Consequently, PCADK-derived delivery systems are very promising for treating MI and other inflammatory diseases. In a following study, the same group demonstrated that acid-sensitive polyketal nanoparticles (~500 nm) loaded with NOX2-NADPH oxidase siRNA can significantly attenuate NOX2 expression and recover cardiac function by inhibiting oxidative stress, after intramyocardial injection in mice with MI [162]. In other studies, pH-responsive microparticles based on acetalated dextran were examined for tunable release of a cardioprotective growth factor to the heart by local delivery [284], and peptide-functionalized acetalated dextran-derivated nanoparticles (100-200 nm) were used for M2-like macrophage-mediated targeted delivery of small hydrophobic compounds to the infarcted heart for MI therapy [285].

In view of overproduced ROS in ischemic myocardium, ROS-responsive delivery systems were investigated for the treatment of acute MI. For example, ROS-responsive nanoparticles self-assembled by a diblock copolymer PEG-b-PPS were used to encapsulate a natural product ginsenoside Rg3. In a rat model of myocardial ischemia-reperfusion injury, intramyocardial treatment with Rg3-loaded PEG-b-PPS nanoparticles significantly decreased the plasma levels of TNF-α, interleukin (IL)-1β, IL-6, and C-reactive protein (CRP), therefore resulting in notably improved cardiac function and reduced the infarct size [247]. In addition, fibrous patches based on polyurethane containing thioetalalinkageswereusedasaROS-responsive delivery platform of methylprednisalone, showing desirable effects with respect to free radical scavenging, rebuilding structures, and improving functions of infarcted myocardium after implantation in MI rats [233]. Also, captopril-loaded mesoporous silica nanoparticles of ~120 nm with H2O2-triggerable release performance were examined for therapy of heart failure in a zebrafish model [297].

Also, abnormally increased enzymes can serve as effective endogenous biological stimuli to realize enhanced accumulation and/or triggerable release of different therapeutics for the treatment of acute CVDs. Christian’s group designed enzyme-responsive nanoparticles for targeted delivery and prolonged retention of therapeutic agents in the heart tissue after MI (Fig. 3) [288]. The mentioned nanoparticles with the average diameter of 20 nm were assembled by a brush peptide-polymer amphiphile (PPA) with a polynorbornene backbone bearing peptide sequences that can be specifically recognized by MMP-2 and MMP-9, while both MMP-2 and MMP-9 are endogenously expressed inflammation-related enzymes post MI. After i.v. injection, the enzyme-responsive nanoparticles efficiently accumulated in the infarct tissue through the leaky vasculature post-MI. Subsequently, up-regulated MMPs at the infarct site induced a morphological transition from discrete nanoparticles into network-like scaffolds, resulting in dramatically increased retention at the injured site for up to 28 days after injection. In another study by Gianschi and Christian et al., they engineered peptide proglulators that are water-soluble cyclic proglulators showing free flow in solution until proteolytically activated by MMP-2/
9 and elastase [289]. After local injection at the site of MI in rats, enzymatic cleavage of cyclic progelators generated linear self-assembling peptides, which rapidly assembled into hydrogels. Consequently, these peptides are promising structurally dynamic biomaterials for minimally invasive therapeutic delivery to the heart after MI. Nevertheless, no further experiments have been conducted using appropriate candidate drugs to demonstrate in vivo efficacies of these enzyme-responsive delivery systems in MI models. Most recently, Dai and Zhao et al. developed a GSH-modified collagen hydrogel (Gel-GSH) for MMP-2/9-responsive on-demand delivery of bFGF [266]. To this end, bFGF was fused with glutathione-S-transferase (GST) and MMP-2/9 cleavable peptide PLGLAG (TIMP), giving rise to GST-TIMP-bFGF that was loaded into Gel-GSH via a bond between GST and GSH (Fig. 4). After intramyocardial injection in rats post MI, substrate peptides were hydrolyzed by MMP-2/9, leading to bFGF release in the pathological microenvironment, thereby promoting vascularization and ameliorating myocardium remodeling. Similarly, MMP-responsive and injectable hydrogel was prepared for sequestration and triggerable release of MMP-2 siRNA for MI treatment in rats [195], which led to significantly improved functions, such as increased ejection fraction, stroke volume, and cardiac output, as compared to controls.

4.1.2. Acute lung injury

Acute inflammation is crucially related to many life-threatening diseases in the respiratory system, such as acute lung injury (ALI) and acute pulmonary infection [290]. ALI is generally resulted from direct and indirect injury to the lungs, showing the most severe manifestation of acute respiratory distress syndrome, which affects approximately 1 million people worldwide annually [291]. Acute pulmonary infection caused by coronaviruses can lead to much more serious global health threats, like novel coronavirus disease 2019 (COVID-2019), severe acute respiratory syndrome, and Middle Eastern respiratory syndrome [292]. Over the past decades, extensive studies have demonstrated the effectiveness and advantages of advanced drug delivery systems for the treatment of pulmonary diseases [293–296].

As well-documented, oxidative stress, resulting from overproduced ROS, is intimately associated with progression of ALI [297]. Therefore ROS can function as either therapeutic targets or biological stimuli for ALI therapy. Our group recently designed and developed a ROS-responsive and ROS-eliminating material (defined as TPCD) by simultaneously conjugating Tempol (Tpl) and PBAP onto β-CD (Fig. 5A-B) [298]. For synthesized TPCD bearing 2 Tpl and 5 PBAP units, it can be hydrolyzed into water-soluble products in the presence of H2O2, a typical ROS. TPCD is able to effectively scavenge superoxide anion, H2O2, radical, and hypochlorite (Fig. 5C-F), thereby serving as a broad-spectrum ROS-eliminating material. Using a nanoprecipitation/self-assembly method in the existence of lecithin and 1,2-distearoyl-sn-glycero-3-phosphoethanolamine-N-[methoxy(polyethylene glycol)-2000, TPCD was produced into nanoparticles with the mean diameter of 109 ± 2 nm and ζ-potential of -16 ± 0.1 mV. TPCD nanoparticles displayed
desirable H₂O₂-responsive hydrolysis performance (Fig. 5G). By efficiently scavenging different types of ROS, TPCD nanoparticles protected RAW264.7 macrophages from H₂O₂-induced apoptosis. After i.v. injection, TPCD nanoparticles could accumulate in injured lungs in mice with LPS-induced ALI by passive targeting. Correspondingly, i.v. treatment with TPCD nanoparticles efficaciously mitigated symptoms relevant to ALI, as implicated by the significantly decreased lung wet/dry weight ratio, notably reduced proinflammatory cytokines (TNF-α and IL-1β) and oxidative mediators (H₂O₂ and MPO) in bronchoalveolar lavage fluid, remarkably decreased infiltration of neutrophils in pulmonary tissues, and improved lung tissue structure (Fig. 5H). These findings suggested that TPCD nanoparticles are promising for the treatment of pulmonary diseases related to acute inflammation and oxidative stress.

Considering acidosis and increased expression of lipase and alkaline phosphatase in bacterial-induced ALI [299], [300], Wang et al. developed pH-responsive nanoparticles for target therapy of ALI [301,302]. To this end, infectious microenvironment-targeting and bioresponsive nanoparticles were assembled using a pH and enzymes multiple-responsive amphiphilic block copolymer consisting of biotinylated PEG-b-poly(β-amino ester)-b-PEG grafted with PEGylated lipid, which were subjected to surface modification with intercellular adhesion molecule-1 (ICAM-1) antibody [302]. The assembled nanoparticles of 120 nm in diameter were loaded with an antibiotic and an anti-inflammatory agent, giving rise to a multifunctional nanotherapy that can target infectious microenvironments and release drug molecules upon triggering by local infectious cues, as demonstrated in an acute lung bacterial infection model in mice. In vivo therapeutic studies in a sepsis mouse model revealed that i.v. treatment with the bioresponsive nanotherapy simultaneously eliminated bacteria and alleviated the host inflammatory response, as indicated by the significantly decreased levels of infiltrating leukocytes and proinflammatory cytokines (such as TNF-α, IL-1β, and IL-6) in infected lungs, and therefore it can be further developed as a mechanism-based nanotherapy for infectious diseases.

In a most recent study, the same group prepared a nanotherapy assembled by an amphiphilic copolymer of poly(β-amino esters) and PEG, loading an anti-inflammatory agent TPCA-1 in the pH-sensitive core [301]. Additional decoration using anti-ICAM-1 antibody was carried out for this TPCA-1 nanotherapy to enhance lung targeting capability. In mice with ALI, i.v. injection of the TPCA-1-loaded targeting nanotherapy (~100 nm) demonstrated efficient targeting and acid-triggered drug release in inflamed lungs, thereby inhibiting lung
inflammation and injury.

4.1.3. Acute kidney injury

Acute kidney injury (AKI), the most common cause of kidney dysfunction, remains a critical health issue with high morbidity and
mortality worldwide [303]. AKI is mainly caused by sepsis, ischemia-reperfusion, and/or nephrotoxicity, and it is characterized by rapid progression of inflammation and oxidative stress. Different drug delivery systems have recently been explored for AKI therapy in murine models, including drug conjugates [304], micelles [305], nanoparticles [306], microspheres [307], and injectable hydrogels [308]. Of note, Ji and Du et al. designed pH-responsive and AKI-kidney targeting nanoparticles by electrostatic forces between anionic hyaluronic acid and cationic chitosan for effective delivery of SS-31 (a mitochondrial targeting peptide with potent antioxidant activity) to AKI. The developed SS-31 nanoparticles showed the mean diameter of 53 ± 0.17 nm, with ζ-potential of -19.6 ± 0.7 mV. In low pH environments, SS-31-containing nanoparticles will be broken due to dissociation of polyelectrolyte complexes, thereby allowing SS-31 release and subsequent mitochondrial targeting to achieve therapeutic effects [309]. While pH-sensitivity of this SS-31 nanotherapy was demonstrated by in vitro experiments, in vivo fluorescence imaging confirmed desirable accumulation of Cy5-labeled nanotherapy in kidneys of AKI mice after i.v. administration. Consistently, the pH-sensitive SS-31 nanotherapy effectively inhibited oxidative stress, protected mitochondrial structure, attenuated inflammatory response, and reduced apoptosis and necrosis of tubular cells after i.v. injection in AKI mice.

In view of overproduced ROS at the AKI site, the newly engineered ROS-responsive and broad-spectrum ROS-scavenging TPCD nanoparticles were employed for the treatment of acute renal injury [298]. Preliminary studies in mice with acetaminophen-induced AKI showed desirable efficacy after i.v. administration, as indicated by notably attenuated hydrenephrosis and significantly decreased serum levels of creatinine (CREA) and urea (UREA). In a most recent study, a ROS-responsive drug delivery system was developed using an amphiphilic copolymer of PEG and PLGA covalently linked via the thioketal bond [310], in which mitochondrial targeting ceria nanoparticles and an anti-inflammatory drug atorvastatin were simultaneously loaded, giving rise to a nanotherapy with the average diameter of 25.5 ± 5.8 nm and ζ-potential of -0.551 ± 0.114 mV. After i.v. injection, fluorescence imaging demonstrated accumulation of the engineered nanotherapy in injured kidneys of mice with sepsis-induced AKI. In vivo studies revealed that this ROS-responsive and ROS-scavenging nanotherapy significantly decreased the levels of malondialdehyde (MDA), TNF-α, and IL-6, thereby notably reducing oxidative stress and inflammatory response. Also, i.v. treatment with this nanotherapy effectively protected the mitochondrial structure and inhibited tubular cell apoptosis and tubular necrosis in AKI mice. According to other pathologically increased biomolecules (such as MMP-2), enzyme-responsive polymeric prodrugs were synthesized for targeted therapy of AKI [311]. For different bioresponsive nanotherapies examined thus far, whereas these studies demonstrated their effectiveness in AKI treatment, the kidney targeting efficiency need to be considerably improved in future studies by delicate modulation on the particle size, shape, surface charge, and density of targeting moieties, to avoid their undesirable distribution in other non-target organs (such as the liver). For example, nanoparticles of ~75 ± 25 nm have demonstrated efficient targeting to the mesangium of the kidney [312]. Also, metabolism, renal excretion, and other safety issues should be elucidated for most nanocarriers by in-depth evaluations.

4.1.4. Acute liver injury

Similar to the above mentioned diseases associated with acute inflammation, the pathophysiological microenvironment in the liver has been deployed to treat acute liver injury, a main cause of diverse liver diseases, such as hepatic fibrosis, cirrhosis, and hepatocellular carcinoma [313]. In this context, pH-sensitive nanoparticles were formulated using ketalized maltodextrin (KMD) to achieve triggerable therapeutic delivery to the injured hepatic tissue resulting from inflammation [314]. KMD was synthesized by covalent conjugation of acid-cleavable hydrophobic groups onto maltodextrin through carbonate bonds. Using silymarin as a candidate drug, in vitro pH-dependent release profiles were found for KMD nanoparticles. Cell culture experiments revealed desirable antioxidant and anti-inflammatory activity of silymarin-loaded KMD nanoparticles. Correspondingly, this pH-responsive nanotherapy effectively suppressed acute inflammation and protected liver from acetaminophen-induced hepatic injury. Also, acid-responsive nanoparticles based on acetalated dextran were simultaneously loaded with a therapeutic compound and gold nanoparticles for targeted therapy and computer tomography imaging of acute liver failure in mice [315].

In addition, ROS-responsive delivery systems were investigated for the treatment of acute liver injury. A study by Lee and coworkers showed that H₂O₂-responsive polymeric prodrug PVAX microparticles containing manganese-porphyrin (a mimic of superoxide dismutase) effectively protected the liver from acetaminophen-induced injury in mice [228]. Zhou and Dong et al. prepared ROS-responsive nanoparticles by self-assembly of PEG-b-PPS, which were used for delivery of melatonin, an antioxidant molecule [316]. In vitro release of melatonin from PEG-b-PPS nanoparticles was considerably accelerated by H₂O₂. In a mouse model of sepsis-induced acute liver injury, the formulated ROS-responsive nanotherapy showed significantly potentiated efficacy compared to free drug after intraperitoneal injection, with respect to attenuating oxidative stress and inflammatory response as well as inhibiting liver injury. Our recent finding revealed a higher accumulation of i.v. delivered nanopoulos in the injured liver, compared to that of the healthy liver in normal mice [298]. Further, treatment of mice with acetaminophen-induced acute liver injury by ROS-responsive and ROS-eliminating TPCD nanoparticles at 1 mg/kg significantly reduced the levels of serum alanine aminotransferase (ALT) and aspartate aminotransferase (AST), in comparison to the model group treated with saline. Moreover, TPCD nanoparticles effectively suppressed neutrophil infiltration, significantly inhibited the expression of proinflammatory cytokines (TNF-α and IL-1β), and notably reduced ROS levels in hepatic tissues. Additionally, the histological abnormalities of the liver were remarkably improved after TPCD NP therapy (Fig. 5). Given the critical role of proinflammatory cytokines in the development of acute liver failure, Yin’s group developed ROS-responsive and macrophage-targeting polyplexes to site-specifically deliver TNF-α siRNA (siTNF-α) to mannose receptor-overexpressed hepatic macrophages [317]. The polyplexes were prepared using low molecular weight PEI cross-linked with a ROS-responsive diselenide bond, which were further coated with carboxylated mannan. After i.v. administration, thus engineered siTNF-α nanotherapy showed effective active targeting to mannose receptor-overexpressed macrophages in the liver and significantly attenuated hepatic inflammation in mice with LPS/D-galactosamine-induced acute liver failure, by notably down-regulating the levels of hepatic and systemic TNF-α.

Collectively, the available findings strongly suggested that ROS-responsive drug delivery systems are particularly promising for the treatment of liver diseases related to acute injury. Notably, liver targeting can be easily achieved by most nanoparticles due to the passive tissue accumulation mediated by the mononuclear phagocyte system. Nevertheless, precision targeting of specific cell types in the liver is still challenging to realize more significant therapeutic benefits and minimize side effects.

4.2. Treatment of chronic inflammatory diseases

Chronic inflammatory diseases, such as IBD, RA, and atherosclerosis, are major causes of death worldwide among older adults. Long-term treatment with traditional anti-inflammatory agents is generally associated with notable side effects. Increasing evidence has demonstrated multiple advantages of bioresponsive drug delivery systems in the management of chronic inflammatory diseases. This section highlights applications of bioresponsive delivery strategies for the treatment of typical diseases related to chronic inflammation.
4.2.1. Inflammatory bowel disease (IBD)

IBD, including ulcerative colitis and Crohn’s disease, is a generic term for a group of chronic relapsing gastrointestinal diseases. It has been extensively demonstrated that overproduced ROS play a critical role in the pathogenesis of IBD [318], and therefore there is an increasing interest in developing ROS-responsive drug delivery systems for IBD therapy. Using a ROS-responsive polymer PAPDT, Murthy et al. developed nanoparticles loaded with siRNA against TNF-α (termed as TKNs, with diameter of ~600 nm) [231]. TKNs released siRNA in response to abnormally high levels of ROS at the site of intestinal inflammation. Oral administration of TKNs significantly reduced the TNF-α mRNA level in the colon of mice with dextran sodium sulfate (DSS)-induced ulcerative colitis, thereby effectively protecting mice from colitis. Of note, TKNs showed significantly higher in vivo activity than TNF-α siRNA-containing nanoparticles derived from PLGA or a cationic lipid 1,2-dioleoyl-3-trimethylammonium-propane (DOTAP), with respect to inhibiting TNF-α mRNA in inflamed intestinal tissues. Our group developed a ROS-responsive nanotherapy mimicking superoxide dismutase/catalase [319]. This nanotherapy is consisted of an oxidation-responsive β-CD material (OxbCD) and a free radical scavenger superoxide dismutase/catalase (SOD/CAT). This composition showed significant inhibition of TNF-α expression, which is associated with many types of ROS, such as superoxide anion, hydroxyl radical, and H2O2. Interestingly, Tpl/OxbCD nanoparticles were stable at pH 1.2, while responsive release profiles were observed under colonic pH conditions, in a ROS-responsive manner. After oral administration, OxbCD nanoparticles showed considerable accumulation in the inflamed colons of mice bearing DSS-induced colitis, with significantly higher targeting efficiency than that of control PLGA nanoparticles. In mice with DSS or 2,4,6-trinitrobenzene sulfonic acid (TNBS)-induced acute colitis as well as DSS-induced chronic colitis, oral delivery of Tpl/OxbCD nanoparticles notably alleviated symptoms relevant to colitis, as exemplified by significantly decreased levels of CD98-positive intestinal epithelial cells and typical inflammatory factors, including TNF-α, IL-1β, and interferon (IFN-γ). In all these mouse models, Tpl/OxbCD nanoparticles were more effective than Tpl/PLGA nanoparticles. In a separate study, we found that this type of H2O2-responsive and ROS-eliminating nanoparticles can potentiote in vivo antitumor activity of a ROS-responsive camptothecin-11 nanotherapy in mice with chronic colitis-associated colon cancer, by normalizing the proinflammatory microenvironment [320]. More recently, our studies showed that OxbCD nanoparticles are able to package a proresolving annexin A1-mimetic peptide Ac2-26, offering a ROS-responsive peptide nanotherapy (defined as AON) (Fig. 6A-E) [321]. AON effectively protected Ac2-26 from degradation in the gastrointestinal environment (Fig. 6F-G). Moreover, site-specific delivery and triggerable release of Ac2-26 at the inflamed colon were achieved after oral delivery of AON (Fig. 6H). In mice bearing DSS-induced acute or chronic colitis, oral treatment with AON significantly decreased the levels of proinflammatory cytokines (TNF-α, IL-1β, and IFN-γ) and oxidative stress-related molecular mediators (MDA, H2O2, and MPO) in the colonic tissues, notably reduced disease activity index, and considerably improved colonic microstructure, thereby achieving desirable therapeutic outcomes (Fig. 61-K).

In vivo efficacy of AON was further confirmed in IL-10-deficient mice that can spontaneously develop colitis. In all cases, AON showed more efficacious effects than a control PLGA nanotherapy. In addition to notably attenuated local inflammation and oxidative stress, AON treatment promoted intestinal mucosal wound healing, reshaped the gut microbiota, and enhanced short-chain fatty acid production. Importantly, AON displayed good safety profile after oral delivery at a high dose of 5 g/kg in mice. Consequently, AON is a promising ROS-responsive nanotherapy for precision therapy of IBD and other inflammatory diseases.

Therapeutic advantages of bioresponsive delivery systems in IBD therapy have also been demonstrated by other studies. Siepmann et al. fabricated microbiota-sensitive 5-aminosalicylic acid pellets [322]. The resulting pellets notably mitigated manifestations relevant to TNBS-induced colitis in rats and significantly decreased inflammatory markers. Also, multi-responsive approaches were examined to deliver therapeutics for IBD management, attempting to overcome the limitations of single signal-dependent strategies. Xiao’s group designed curcumin-loaded nanoparticles derived from natural silk fibroin with surface decoration using chondroitin sulfate [323]. The generated nanoparticles displayed good bioreponsiveness upon triggering by acidity, GSH, and ROS, in activated macrophages. After oral administration or i.v. injection, this nanotherapy notably mitigated the symptoms of colitis, maintained the gut microbiota, and enhanced the survival rate of colitis mice. In addition, mannosylated bioreducible nanoparticles, pH/enzyme dual-responsive nanogels, and pH/ROS-responsive nano-in-micro composites were investigated as delivery vehicles of different drugs for targeted treatment of IBD [272,324,325].

4.2.2. Inflammatory vascular diseases

Sustained vascular inflammation has been implicated in many CVDs that remain the leading cause of global morbidity and mortality. Excessive ROS production has been found in injured arteries of many vascular diseases, such as restenosis, atherosclerosis, and aneurysm. According to local acidosis and elevated levels of ROS at the injured sites of arteries, our group developed pH- or ROS-responsive nanotherapies of 186.5 ± 2 or 253.5 ± 3.9 nm in diameter, by loading rapamycin (RAP) into nanoparticles based on acetalated β-CD or PBA conjugated β-CD, respectively (Fig. 7A) [111,182]. The obtained RAP nanoparticles displayed either pH- or ROS-triggerable drug release behaviors. Compared to free RAP and a nonresponesive PLGA nanotherapy, both responsive nanotherapies exhibited significantly higher anti-migration and antiproliferative effects in rat vascular smooth muscle cells (VSMCs). After i.v. injection, the related nanoparticles accumulated at the injured carotid arteries of rats with balloon angioplasty injury (Fig. 7B), which is a typical animal model of arterial restenosis. Moreover, i.v. treatment with pH- or ROS-responsive nanotherapy more significantly inhibited neointimal hyperplasia, compared to free RAP and a PLGA nanotherapy (Fig. 7C). In a most recent study, we demonstrated nanoparticles with tunable dual-sensitivity can be easily prepared by combining pH- or ROS-responsive β-CD materials [88]. In rat VSMCs, thus engineered pH/ROS dual-responsive RAP nanotherapies showed potentiated anti-migration, anti-proliferation, and cell cycle regulation capabilities, when compared with corresponding single-responsive RAP nanotherapies. Also, in vivo studies revealed enhanced therapeutic benefits of the pH/ROS dual-responsive nanotherapy, in comparison to the pH- or ROS-responsive RAP nanotherapy. Further, surface decoration of the dual-responsive nanotherapy via a type IV collagen-targeting peptide (KLWLPKGGGC) notably potentiated its in vivo anti-restenosis efficacy, resulting from the increased accumulation in injured carotid arteries by active targeting.

Atherosclerosis, a chronic inflammatory disorder characterized by the formation of atheromatous plaques in the intima of arteries, may cause severe diseases such as MI and stroke. In a previous study, we found that the above mentioned pH- or ROS-responsive RAP nanotherapy derived from chemically functionalized β-CD materials remarkably decreased the macrophage count and MMP-9 level in aortas, and therefore delayed progression of atherosclerosis and significantly enhanced stability of plaques in apolipoprotein E-deficient (ApoE−/−) mice [182], compared to a non-responsive PLGA nanotherapy (Fig. 7D-F). This enhanced anti-atherosclerotic activity of RAP by responsive nanocarriers should be mainly due to triggerable and effective release of drug molecules in atheromatous plaques, following targeting by the lymphatic translocation mediated via inflammatory infiltration of neutrophils and monocytes/macrophages that was demonstrated by ex vivo imaging in combination with flow cytometric analysis. Moreover, these responsive nanocarriers displayed good safety after long-term...
administration in mice. In a following study, we explored anti-atherosclerotic activity of our previously developed ROS-responsive and broad-spectrum ROS-eliminating TPCD nanoparticles [326], in view of deleterious effects of overproduced ROS on atherosclerosis. In vitro experiments revealed that TPCD nanoparticles notably suppressed foam cell formation in both macrophages and VSMCs. After i.v. administration, TPCD nanoparticles could accumulate in atherosclerotic lesions of ApoE-/- mice, mainly resulting from passive targeting via the permeable endothelium and translocation by inflammatory cells. Further in vivo therapeutic studies indicated that i.v. treatment with TPCD nanoparticles attenuated the development of atherosclerosis and stabilized plaques, as implicated by thicker fibrous caps, smaller necrotic cores, and lower macrophage counts, compared to control antioxidant drugs against atherosclerosis. The desirable in vivo efficacies of TPCD nanoparticles were largely contributed by effectively decreasing the levels of TNF-α, IL-1β, and H2O2 in the serum and aorta, which in turn led to attenuated systemic/local inflammation and oxidative stress. Since TPCD nanoparticles showed good biocompatibility in different animal models [298,326], they can be further intensively studied as a safe and effective nanotherapy for both acute and chronic inflammatory diseases. Also, ROS-responsive nanoparticles containing atorvastatin were functionalized with macrophage cell membrane [327], leading to nanotherapies with further increased in vivo anti-atherosclerotic efficacies. Most recently, we engineered a pH-responsive anti-miRNA-33 nanotherapy (defined as AAM NP), by packaging an antisense oligonucleotide against microRNA-33 (anti-miR33) into nanoparticles derived from an acetalated α-CD material and PEI (Mw = 1800 Da) for targeted therapy of atherosclerosis [328], considering lesions acidosis. AAM NP was further decorated with a peptide ligand cRGDfk targeting integrin to develop an active targeting nanotherapy RAAM NP. After i.v. delivery, both nanotherapies can be accumulated in the plaques and efficiently internalized by endothelial cells and macrophages, with RAAM NP showing considerably enhanced targeting capability. Treatment with both anti-miR33 nanotherapies significantly attenuated atherosclerosis in ApoE-/- mice and notably reduced vulnerable plaques by promoting reverse cholesterol transport and regulating adaptive immunity via modulating macrophage polarization and regulatory T cell differentiation. In other studies, redox-sensitive gene delivery nanoplatforms were examined for reducing atherosclerotic inflammation [273], by targeted delivery of eNOS plasmid to endothelial cells. These studies unambiguously demonstrated that pH-, redox-, or ROS-responsive nanoparticles are highly effective delivery vehicles for precision therapy of atherosclerosis by site-specific release of different therapeutic agents.

In an attempt to develop targeting therapies for abdominal aortic aneurysm (AAA), a leading cause of mortality and morbidity in the
elderly, we recently examined the effectiveness of ROS-responsive RAP nanotherapies (Fig. 8) [329]. To this end, nonresponsive or ROS-responsive RAP nanotherapies were prepared by a nanoprecipitation/self-assembly method, using PLGA or OxbCD as carrier materials, respectively. While both nanotherapies remarkably inhibited calcium and inorganic phosphorus-induced calcification and cell death as well as decreased H2O2-induced ROS generation and apoptosis in VSMCs, much better results were observed for the ROS-responsive nanotherapy (OR NP). After i.v. administration in rats with CaCl2-induced AAAs, OxbCD nanoparticles displayed significantly higher accumulation in aneurysmal aortas than PLGA nanoparticles. By decreasing expressions of MMPs (MMP-2 and MMP-9) and proinflammatory cytokines (TNF-α, IL-1β, and IFN-γ), inhibiting aortic calcification and macrophage infiltration, and lowering ROS production, OR NP afforded more desirable in vivo efficacies than PLGA-based nanotherapy. Subsequently, OR NP was decorated with a cyclopeptide ligand cRGDfK that can target neovessels via binding with αvβ3 integrin (Fig. 8A). Both in vitro and in vivo studies revealed notably enhanced anti-aneurysmal efficacy of the active targeting and ROS-responsive nanotherapy ROR NP (Fig. 8B-E). Finally, functionalization with macrophage cell membrane was conducted to further increase targeting capability of ROR NP. The engineered biomimetic and bioreponsive nanotherapy more significantly prevented the development of AAAs, as indicated by the notably attenuated expansion of the aortic diameter as well as remarkably decreased calcification and elastin degradation, resulting from improved aneurysmal targeting after macrophage membrane cloaking.

In other studies, ROS-responsive as well as pH- and/or thrombin-responsive nanotherapies were examined for targeted treatment of thrombus formation [330], peripheral arterial disease [226], and ischemic stroke [331–334], offering promising therapeutic effects in different animal models. Collectively, studies by others and our group demonstrated that bioresponsive nanotherapies possess great potential for precision therapy of different inflammatory vascular diseases.

4.2.3. Rheumatoid arthritis (RA)

RA is an autoimmune and chronic inflammatory disease in the joints that may cause cartilage damage, bone erosion, and loss of joint function. To overcome significant side effects of small molecule anti-inflammatory agents, Wang’s group developed a pH-sensitive drug delivery system by conjugating dexamethasone (Dex) onto a N-(2-hydroxypropyl)methacrylamide copolymer (P-Dex) via an acid-labile

Fig. 7. Inflammation-responsive nanotherapies for targeted treatment of atherosclerosis and restenosis. (A) Design of inflammation-triggerable nanoparticles. (B) Representative ex vivo fluorescence images showing retention of Cy7.5/Ac-bCD NP at the injured site of the carotid artery in rats after intravenous injection. (C) H&E stained histological sections of carotid arteries isolated from rats subjected to various treatments. Scale bars, 200 μm (the upper panel), 50 μm (the lower panel). Reproduced with permission [111]. Copyright 2016, Elsevier. (D) Representative ex vivo fluorescence images illustrating accumulation of Cy5 fluorescent signals in the aorta of ApoE−/− mice at 24 h after intraperitoneal injection of saline, free Cy5, or Cy5/Ac-bCD NP. (E) Representative photographs of en face ORO-stained aortas from each group. (F) Quantitative analysis of the lesion area. **p < 0.01, ***p < 0.001. Reproduced with permission [182]. Copyright 2017, Elsevier. RAP, rapamycin; PLGA, poly(lactide-co-glycolide); Ac-bCD, acetalated β-cyclodextrin; Ox-bCD, oxidation-responsive β-cyclodextrin.
Lydia hydrazone bond, which can release free dexamethasone at acidic pH [189]. This polymeric prodrug showed specific accumulation in inflamed joints of rats with adjuvant-induced arthritis (AIA). Compared to free Dex, i.v. treatment with P-Dex afforded more effective and longer-lasting anti-inflammatory effects, with notably reduced side effects. Therapeutic benefits of P-Dex were also demonstrated in a mouse model of collagen-induced arthritis (CIA) and mice with particle-induced inflammation in joints [190,335]. Based on the similar principle, Sun et al. prepared pH-sensitive micelles assembled by an amphiphilic copolymer derived from a PEG derivative with conjugated prednisolone via the hydrazone linkage (Fig. 9A) [188]. After i.v. injection in CIA mice, prodrug micelles showed significantly higher accumulation in the joints than that of free drug. Correspondingly, pH-sensitive micelles exhibited notably enhanced efficacy and simultaneously reduced myelo- and hepatotoxicity, as compared to free prednisolone. In a most recent study, Chen et al. developed macrophage-targeting pH-sensitive nanoparticles for co-delivery of two anti-inflammatory agents (Fig. 9B) [192]. All-trans-retinal was conjugated onto dextran via the hydrazone bond, followed by conjugation of a targeting unit of galactose. The assembled nanoparticles were physically loaded with triptolide, a potent anti-inflammatory drug isolated from Chinese herb. Both drugs in the assembled nanoparticles displayed acid-triggerable release profiles. In CIA mice, the nanoassembly with galactose decoration showed remarkably higher accumulation in the inflamed joints after i.v. injection, due to combined passive and active targeting effects. Further in vivo studies revealed that encapsulation into pH-responsive and targeting nanoparticles notably potentiated anti-arthritis activity of triptolide, while drug toxicity was considerably reduced. Likewise, folate receptor (FA)-targeting, acid-labile nanoparticles derived from PCADK/lipid/PLGA-PEG or PEGylated amphiphilic hyaluronan nanoparticles with calcium phosphate as a pH-responsive mineral, both containing methotrexate, were investigated for targeted RA treatment in AIA or CIA mice [336,337], respectively, giving rise to anticipated therapeutic benefits.

In addition to acid-labile nanotherapies, ROS-responsive delivery systems were examined for the RA treatment. In view of high ROS levels in diseased tissues of RA, Liu et al. fabricated ROS-responsive micelles of ~153 nm based on a vitamin E-conjugated amphiphilic copolymer derived from PLGA and PEG that were linked via a ROS-responsive diselenide bond [338], in which a candidate anti-RA drug berberine was loaded. In vitro studies based on liquid chromatography-mass spectrometry showed that ROS-responsive micelles increased both cellular and mitochondrial concentrations of berberine in RA fibroblasts nearly ten times higher than those of free berberine. After intraperitoneal injection in AIA rats, the ROS-responsive micellar nanoassembly significantly inhibited the levels of inflammatory cytokines (IL-1 and IL-6) and reduced the bone destruction, with much better efficacy than berberine. In other cases, ROS-responsive and FA-targeting liposomes and nanoparticles were investigated for targeted RA treatment [339,340], and both systems notably reinforced in vivo efficacies of loaded drugs in CIA mice.

Also, enzyme-responsive drug delivery systems have been developed for arthritis therapy. As a representative example, arthritis flare-responsive hydrogel was self-assembled by triglycerol monostearate (TG-18) to encapsulate an anti-arthritis drug triamcinolone acetonide (TG-18) to encapsulate an anti-arthritis drug triamcinolone acetonide [341]. This injectable hydrogel showed on-demand drug release profiles upon triggering by MMP-3 or MMP-9. After intra-articular injection, TG-18 hydrogel underwent disassembly correlating with arthritis severity in a mouse model of inflammatory arthritis. Moreover, local drug delivery with TG-18 hydrogel significantly reduced paw thickness and total clinical scores in mice with arthritis, while free drug did not show notable therapeutic effects, mainly due to rapid clearance from the injection site. Accordingly, bioresponsive systems can be further developed as a promising on-demand delivery strategy to potentiate anti-arthritis activity of different drugs.
4.3. Other inflammation-associated diseases

A growing body of evidence suggests that chronic inflammation and oxidative stress also play an important role in the pathophysiology of neurodegenerative diseases, such as Alzheimer’s disease (AD) and Parkinson’s disease. However, many anti-inflammatory or antioxidant therapies cannot effectively inhibit brain injury due to their limited ability to cross the blood-brain barrier (BBB) and target the disease site. Bioresponsive nanotherapies have been emerging as an effective strategy to overcome the BBB and afford the advantage of site-specific release in the impaired brain. In an elegant study, Jiang’s group synthesized a ROS-responsive amphiphilic copolymer by conjugating phenylboronic ester onto the polylysine block of PEG-polylysine, which was then functionalized with a peptide targeting receptor for advanced glycation end-products [342]. The copolymer-assembled ROS-responsive and targeting micelles were packaged with a model drug curcumin (Fig. 10). After i.v. injection in APPsw/PSEN1dE9 model mice, the micelles efficiently accumulated in the diseased regions by β-amyloid transportation-mediated pathway, significantly decreased the levels of proinflammatory cytokines TNF-α and IL-1β in the brain, and showed neuroprotection and microglia modulation via the synergistic effects of polymers and payloads, thereby decreasing β-amyloid plaque burdens and enhancing cognitive functions.

Chronic inflammation also contributes to common respiratory system diseases, such as asthma, chronic obstructive pulmonary disease, and cystic fibrosis. Both nanoparticles and microparticles are promising delivery vehicles for the treatment of pulmonary diseases [293,295]. Lee et al. explored they previously developed H2O2-responsive and H2O2-scavenging HPOX nanoparticles for the treatment of asthma in a murine model [343]. After intratracheal injection in asthmatic mice induced with ovalbumin, HPOX nanoparticles significantly reduced the infiltration of inflammatory cells (macrophages and neutrophils) and expression of proinflammatory mediators (IL-4 and inducible nitric oxide synthase) as well as decreased airway remodeling. Recently, nanoparticle-embedded microgels responsive to both trypsin and neutrophil elastase were prepared by the thiol-maleimide Michael addition reaction in emulsions, using model polystyrene nanoparticles and enzyme-degradable peptide crosslinkers [344]. After aerosolized delivery, the enzyme-responsive microgels showed desirable pulmonary retention and disease-triggered degradation. Nevertheless, further
therapeutic studies in specific disease models are necessary to demonstrate advantages of this novel multi-stage enzyme-responsive microgel. Collectively, the currently available findings suggest that bioresponsive delivery systems have great potential as novel drug carriers for targeted treatment of neurodegenerative diseases and airway inflammatory diseases, although intensive and in-depth studies are necessary to evaluate their effectiveness, advantages, and benefit-risk ratio in different disease models. In addition, bioresponsive nanotherapies are able to attenuate surgical injury-induced inflammation. As demonstrated by laminectomy models in rats and rabbits [345], the aforementioned TPCD nanoparticles loaded in a poloxamer 407-derived thermosensitive hydrogel displayed notable antioxidative stress and anti-inflammatory activities by significantly reducing the levels of H$_2$O$_2$, MPO, TNF-α, IL-1β, and IL-6, thereby effectively reducing fibrotic tissue formation and preventing postlaminectomy epidural adhesion.

5. Concluding remarks

Over the past decades, considerable progress has been achieved in the development of bioresponsive materials and delivery systems as well as their applications in the treatment of both acute and chronic inflammatory diseases. Particularly, a large number of materials responsive to varied inflammatory microenvironments have been synthesized and evaluated in different cells and/or animal models. By triggerably releasing therapeutic molecules in response to pathologically relevant biochemical signals such as low pH, increased ROS, and overexpressed enzymes, their efficacies can be considerably enhanced, as well demonstrated in preclinical studies based on animal models of acute CVDs, acute liver injury, ALI, AKI, IBD, inflammatory vascular...
diseases, and RA. Despite these achievements, only a very few of bioresponsive drug delivery systems have been intensively investigated in diverse animal models of inflammatory diseases. To the best of our knowledge, currently no translation studies have been conducted for bioresponsive therapies in anti-inflammatory interventions.

Some crucial challenges should be addressed before further clinical translation studies. First, for rational design of bioresponsive drug delivery systems, the optimal biochemical signal need to be selected according to the specific disease types and diseased sites. The degree of local acidosis varies in different inflammatory diseases. Also, some normal tissues and subcellular organelles exhibit acidic environments, such as the stomach and endolysosomes, which may cause off-target drug release from pH-responsive delivery platforms, thereby impairing in vivo efficacies or generating side effects. Likewise, the ROS generation and the major type of ROS may be vary in different inflammatory diseases or at the different stages of the same disease. As for enzyme-sensitive systems, the corresponding enzymes expressed in other non-target tissues also affect their drug release behaviors. Unfortunately, these issues remain elusive, which complicates and limits the rational design and engineering of bioresponsive delivery systems with more precisely controlled release profiles. Whereas multi-responsive systems afford much more sensitivity and specificity, their development frequently involves more complicated materials synthesis and/or formulation procedures. Second, preparation processes of currently available bioresponsive delivery systems remain to be optimized to achieve desirable quality control, such as well-defined material structures, tailorable biophysicochemical properties, and good reproducibility. Cost-effective mass production is also a key issue to simultaneously ensure clinical benefits and low cost, which is especially important for the management of chronic inflammatory diseases. Third, new preclinical animal models that can fully replicate the pathophysiological microenvironment of human inflammatory diseases need to be established. Thus far, most in vivo targeting and therapeutic studies of bioresponsive delivery systems were performed in murine models of inflammatory diseases. The obtained results cannot be directly extrapolated to humans, due to the intrinsic differences between humans and other species. In this context, targeting capability and/or efficacies of currently developed anti-inflammatory delivery systems should be further examined in multiple animal models to afford more convincing results and screen translational formulations. For bioresponsive anti-inflammatory nanotherapies, their accumulation in off-target organs or biomimetic approaches, remain to be systematically examined to maximally enhance targeting efficiency of these nanotherapies at inflammatory sites, thereby notably minimizing their nonspecific distribution and reducing side effects. Last but not least, both acute and chronic toxicity evaluations must be carefully conducted for some promising bioresponsive materials or their delivery systems. The perfect bioresponsive delivery systems should maximize drug efficacies and simultaneously reduce or avoid drug-mediated side effects, thereby providing patients with therapeutic benefits outweighing risks.

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