Novel zwitterionic vectors: Multi-functional delivery systems for therapeutic genes and drugs

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Abstract
Zwitterions consist of equal molar cationic and anionic moieties and thus exhibit overall electroneutrality. Zwitterionic materials include phosphorylcholine, sulfo betaine, carboxy betaine, zwitterionic amino acids/peptides, and other mixed-charged zwitterions that could form dense and stable hydration shells through the strong ion–dipole interaction among water molecules and zwitterions. As a result of their remarkable hydration capability and low interfacial energy, zwitterionic materials have become ideal choices for designing therapeutic vectors to prevent undesired biosorption especially nonspecific biomacromolecules during circulation, which was termed antifouling capability. And along with their great biocompatibility, low cytotoxicity, negligible immunogenicity, systematic stability and long circulation time, zwitterionic materials have been widely utilized for the delivery of drugs and therapeutic genes. In this review, we first summarized the possible antifouling mechanism of zwitterions briefly, and separately introduced the features and advantages of each type of zwitterionic materials. Then we highlighted their applications in stimuli-responsive “intelligent” drug delivery systems as well as tumor-targeting carriers and stressed the multifunctional role they played in therapeutic gene delivery.

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1. Introduction

Over the past decades, large quantities of novel materials have been widely investigated and developed for drug and gene delivery to maintain their structural stability, alleviate cytotoxicity towards normal tissues, improve the pharmacokinetic profile and promote therapeutic efficacy [1]. However, nonspecific biomolecular and microorganism adherence is still a challenge obstructing their application in disease diagnose and treatment. Therefore, it is significant to find a method which can not only meet the requirements of nanodrug vehicles, but also resist undesired bioadsorption effectively. The ability to avoid bioadsorption is termed antifouling property. Contributing to the character that can increase water solution and resist undesired adherence of biomacromolecules, poly(ethylene glycol) (PEG) has become the most commonly used biocompatible antifouling material to prevent the interactions with blood components, and prolong the in vivo circulation time [2–4]. However, it was reported that PEGylations would lose their protein repulsive capability at temperatures above 35 °C [5]. And PEG has been found susceptible to oxidation in the presence of oxygen or transition metal ions, as its non-ionic structure damage, resulting in the loss of function in most biochemically relevant solutions [6]. Aside from that, PEG was proved to be difficult to metabolize naturally, and repeated injection of PEGylated preparations would reduce the bioactivity of loaded biomolecules due to the antibodies produced by immune system [7]. Additionally, the gene entrapment efficiency of PEGylated cationic liposomes would decrease due to the decline of positive charges [8–10]. Hence, it is necessary to find alternative materials that can not only resist bioadsorption during blood circulation but can also keep stable in different environment, be biodegraded via metabolism and have negligible immunogenicity other than PEG.

Recent years, zwitterionic materials have attracted much attention in the field of biomaterials for their superhydrophilicity, biocompatibility, antifouling property and some other unique advantages [11]. Characterized by high dipole moment and highly charged groups, zwitterionic materials simultaneously possess an equimolar number of cationic and anionic moieties, maintaining overall electroneutrality and high hydrophilicity [12,13]. Unlike PEG binding water molecules by hydrogen-bond interaction, zwitterions could form stronger hydration shells through ion–dipole interaction with denser and tighter adsorbed water [14], which finally leads to the ultralow of nonspecific protein adsorption, bacterial adhesion, and biofilm formation, making zwitterions preferable substitutes for PEG [15].

In addition to their marvelous antifouling property, zwitterionic materials can also enhance biocompatibility, reduce immune response, prolong the circulation time as well as promote cellular uptake of traditional chemical drugs and therapeutic genes. And zwitterionic modifications could even endow various special functions to common drug carriers, such as stimuli-responsive and tumor targeting abilities. In the following content, we will describe the features of diverse zwitterionic materials, introduce their unique advantages and summarize their applications in gene and drug delivery systems.

2. The antifouling mechanism of zwitterions

Biofouling is the spontaneous but unexpected, non-specific adsorption of biomolecules, cells or bacteria on the surface of materials [16]. In the blood, serum proteins are the main force of biosorption, which always leads to the phagocytosis and metabolism of foreign substances by reticuloendothelial systems (RES) [17]. For nanomedicines, it could greatly shorten their in vivo circulation time and results in unsatisfied therapy effect [18]. This kind of protein adhesion happens through electrostatic and hydrophobic interaction with charged segments and non-hydrophilic pockets in proteins [19]. Therefore, various approaches have been investigated to improve the antifouling properties, and the basic principle to design antifouling materials is to avoid electrostatic and hydrophobic interaction with biomolecules, which usually requires the overall neutral surface charge, remarkable hydrophilicity, hydrogen bond acceptors but no hydrogen bond donors [20].

For a long time, PEGylation of nanocarriers has been considered to be the standard anti-adsorption strategy for the hydration layer formed by hydrogen bonding [21,22]. However, this type of interaction is not strong enough that even at a high PEG grafting density, it is difficult for materials with simple PEG layers to eliminate protein adsorption completely [23–25]. Unlike PEG, zwitterions could form denser and more stable hydration shells through a totally different way. On the one hand, since zwitterions have a large amount of cationic and anionic groups, their ion–dipole interaction with water molecules brings them super hydrophilicity and high hydration capability [26]. This type of coulombic force is more powerful compared with hydrogen bonding, so that zwitterionic materials could form stronger water shells than that of PEGylations to prevent proteins form adhering [19]. On the other hand, in saline solution such as the physiological environment, the charged groups on zwitterionic polymers could attract counterions in solution to counteract the internal electrostatic force of the zwitterionic brushes, and change their conformation from shrinking to stretching relatively [27–29]. As the result of that, the hydration ability becomes more significant and brings about an enhanced antifouling property. Moreover, since the positively and negatively charged groups of zwitterions are equimolar, the overall electroneutrality further reduces the chances of non-specific electrostatic bonding. For better understanding, the schematic diagram of hydration layer formation is showed in Fig. 1.

3. Classification of zwitterionic materials

According to whether the cationic and anionic groups are on the exact same unit of zwitterions, these materials can be roughly classified as betaine-like zwitterions and mixed-charge zwitterionic materials (Fig. 2). Most of zwitterionic polymers belong to the
former, they always take quaternary ammonium as cations to constitute phosphorylcholine (PC), sulfobetaine (SB) and carboxybetaine (CB) with phosphonates ($PO_3^{2-}$), sulfonates ($SO_3^{2-}$) and carboxylates ($COO^-$) respectively. Except for zwitterionic polymers containing both charged segments on the same side chains, there are some mixed-charge materials containing balanced positive and negative charged moieties in different monomer units or binding to the same media (e.g. mesoporous silica nanoparticles) to maintain the overall electrical neutrality. These “spurious” zwitterionic materials own similar antifouling properties because of their semblable structures. Besides, since amino acids are a category of natural zwitterions, researchers have managed to introduce amino acids or peptides to develop zwitterionic carriers, which also exhibited great nonspecific absorption resistance and some unique properties.

3.1. Betaine-like zwitterions

3.1.1. Phosphorylcholine

PC is a zwitterionic polar group of phospholipids which play an important role in cell membranes. Therefore, PC is promising for biomimetic materials, and PC-containing nanostructures have been proved to enter living cells through fusogenic interaction with the plasma membranes [30,31]. Ever since the famous PC-type monomer 2-methacyrloyloxyethyl phosphorylcholine (MPC) was synthesized by Ishihara et al. in 1990, MPC has been utilized to build a variety of zwitterionic polymers for its great biocompatibility and strong hydration caused by electrostatic interactions [32]. About 14 years after that, Ishihara and his team reported the first PC-modified drug delivery system (DDS), poly(2-MPC-co-n-butyl methacrylate) (PMB), which was designed to carry the poor hydrophilic paclitaxel (PTX) [33]. The zwitterionic MPC served as hydrophilic segments, which could make the copolymers self-assemble into micelles in aqueous solution together with hydrophobic parts. Thus, the water solubility of PTX was enhanced, and its stability in the blood was also improved taking advantages of the antifouling property of zwitterionic MPC [12,34]. However, it is not easy to synthesize and handle PC-based polymers in general since their monomers, such as MPC, are moisture sensitive, and studies found out that their phosphoester groups had a tendency to be hydrolyzed, which would shorten their in vivo circulation time and badly obstruct application for drug or gene delivery [35,36].

3.1.2. Sulfobetaine

Betaine is a type of natural zwitterionic molecule widely existing in many plants. As a derivate of it, SB has the characteristics of common zwitterions and exhibits extremely remarkable
By means of differential scanning calorimetry, researchers found that there are 8 mol H₂O bound to 1 mol SB group, which is far more than a single unit of PEG [37,38]. For that reason, SB has attracted growing attention with its good biocompatibility and resistance to nonspecific protein adsorption. Recent years, several studies have attempted to introduce SB groups to delivery systems, in which the modified carriers exhibited effective drug loading, high cellular uptake rate, tiny cytotoxicity and faint immune response. For instance, researchers developed a starch-based polymer SB-ST-OC (SSO) with hydrophobic octane groups (OC) and zwitterionic SB, and the results proved that SSO micelles had less macrophage activation potential compared with ST-OC and naked DOX, through detecting the secretion of TNF-α and IL-6 on the macrophage phagocytosis of micelles [39]. And another important reason why SB-coupled materials, especially sulfobetaine methacrylate (SBMA), become so attractive should be the lower synthetic cost compared with PC-modified ones. Since the strong charge-charge or dipole–dipole interactions among the zwitterionic sulfobetaine moieties, SB-based polymers usually exhibit a sharp phase transition at the upper critical solution temperature (UCST) in aqueous solution [40]. That means their solubility would be influenced by the polarities and ionic concentration of solvent [41–44], and by adjusting the solvent conditions, the UCST could be modulated to a desired temperature to build thermo-responsive polymers [45].

Fig. 2. Models and structures of zwitterionic materials. (A) Chemical structures of the common zwitterionic groups. (B) Zwitterionic poly(amino acids) and polypeptide. (C) Mixed-charge zwitterionic polymers that have balanced cationic and anionic groups in different monomer units, and pseudo-zwitterionic materials with equimolar negative and positive charge binding to the same medium.

3.1.3. Carboxybetaine

CB could be one of the most popular zwitterions in the last few years. Similar to SB, CB is also a derivative of the betaine that is ultralow-fouling characterized. Although SB and CB are alike in structures and are both good at resisting undesired adherence, the difference between their anionic groups (COO⁻ for CB and SO₃ for SB) makes these two zwitterions have diverse properties in hydration, ionic interaction, and self-association [12]. As mentioned above, the solubility of PSB depends on the specific conditions of solvents, while polycarboxybetaine (PCB) has been proved to have remarkable hydration properties in a wide range of media [12]. And scientists found that the cationic and anionic groups of CB segments formed few inter-/intra-associations, so that CB-grafted polymers hardly exist thermal or salt responses caused by the self-associations among zwitterionic moieties, whereas SB polymers do [41]. More importantly, the carboxylate anion groups of PCB are able to be functionalized for the immobilization of amine-containing biomolecules including some antibodies, targeting ligands and fluorescent dyes, allowing PCB and its derivatives multifunctional materials [46].

3.2. Mixed-charge zwitterionic materials

3.2.1. Zwitterionic amino acids and peptides

Amino acids (AAs) are natural zwitterions that have both positive charged amino groups and negative charged carboxyl groups on the α-carbon atoms. AAs possess the common advantages zwitterionic materials have, so it is valid to conjugate them as terminal groups with conventional polymers that are weak in undesirable biosorption resistance or stability [47]. Some basic AAs including lysine (Lys), histidine (His) and arginine (Arg) have one more amine group, while some AAs such as aspartic acid (Asp) and glutamic acid (Glu) have one more carboxyl group. What makes it special is that these types of AAs have redundant functional groups, so they could form peptide bonds with α-carboxyl/amino groups to get mixed-charge zwitterionic peptides, or perform the dehydration condensation reaction with their redundant carboxyl/amino groups to get betaine-like zwitterionic side chains with terminal
AAs (Fig. 2B). Since the zwitterionic polypeptides (ZIPPs) are another kind of biomimetic materials that are similar to the surface proteins on cellular membranes, they generally exhibit outstanding biodegradability, biocompatibility and in vivo stability [48]. Even more, the amino terminals and carboxyl ends make it possible to introduce functional biomolecules just like CB-based materials, so zwitterionic AAs and polypeptides are great choices for drug/gene delivery requiring abundant bio-functions such as tumor targeting.

3.2.2. Other mixed-charge zwitterionic materials

In addition to the mixed-charge zwitterionic peptides, zwitterionic polymers could also be synthesized with different components bearing equivalent opposite charges to maintain the overall electroneutrality (Fig. 2C) [49]. For example, taking zwitterionic amino acid decorated amphiphilic DDSs as models, poly(aminobutyl methacrylate)-co-poly(methacrylic acid)-co-poly(n-butyl methacrylate) (CPMA) was developed for the delivery of doxorubicin (DOX) [50]. Although the cationic amino groups and anionic carboxyl groups of this copolymer were distributed in the different side chains, they still owned excellent antifouling property, biocompatibility as well as biostability. What’s more, trough linking side chains, they still owned excellent antifouling property, biocompatibility. But that materials are applied to carry therapeutic agents for their ultra-tumor targeting.

4. Applications of zwitterionic materials in drug delivery systems

As mentioned at the beginning, more and more zwitterionic materials are applied to carry therapeutic agents for their ultratargeting property and wonderful biocompatibility. But that alone is far from enough to meet clinical demands, since competent carriers should also have great loading volumes, promote cellular uptake, and control the drug release as expected. Therefore, sorts of “intelligent” carriers have been engineered to undergo corresponding changes in material properties when exposed to different biological stimuli [53], so that they are able to protect fragile cargos until they reach the targets and unload them effectively at active sites. The zwitterionic drug delivery systems involved in this review are shown in Table 1.

4.1. pH-responsive zwitterionic DDSs

The inherent differences of pH value among different tissues, organs and cellular compartments make pH an attractive factor to control drug delivery according to diverse internal environment [54–56]. It’s well known that the tumor microenvironments (pH 6.4–6.8) are always more acid than that in bloodstream and normal tissues (pH 7.4) [57], taking use of it could promote the accumulation of drugs around tumor cells to reduce side effects. And since the pH decreases from pH 7.4 to pH 5.5–6.0 in endosomes and even below pH 5.0 in lysosomes, it could act as a trigger for drug release by lowering the pH during endocytosis [58]. Therefore, scientists

| Category | Polyampholytes | Working morphology | Loaded drugs | Special properties | Ref. |
|----------|----------------|-------------------|--------------|--------------------|-----|
| PC       | PMB            | micelles          | PTX          | -                  | [33]|
| PMBC-PDPA| micelles/vesicles | PTX/DOX       | pH-responsive | [61,63]           |     |
| NaHiso-PCc | NPs            | quercetin         | DOX          | pH-responsive      | [73]|
| PMBC-PFLA| vesicles       | DOX/DOX HCl      | pH-responsive | [85]              |     |
| PMBC-Bink-CPLA | micelles | PTX          | pH-responsive | [86]              |     |
| PLL-PMBC(DOX) | micelles | DOX          | pH-responsive | [87]              |     |
| PMBC-PCL | micelles       | DOX              | -            | [104]             |     |
| PMBC-6sPCL| micelles      | PTX              | -            | [105]             |     |
| PMBC-ss-PCL| micelles    | DOX              | redox-responsive | [106]           |     |
| PMBC-TMZ | micelles       | TMZ              | redox-responsive | [111]            |     |
| FA-MPC-DPA | micelles    | tamoxifen, PTX   | pH-responsive tumor-targeting | [136] |     |
| FAMEG-MPC-PCL | micelles | DOX              | tumor-targeting | [140]            |     |
| SB       | SSO            | micelles         | DOX          | -                  | [39]|
| PSBMA-PDEA-PCL | micelles | CUR              | pH-responsive | [68]             |     |
| PSBMA    | nanogels       | DOX              | redox-responsive | [110]          |     |
| PCL-ss-PDASEB | micelles | DOX              | pH- and redox-responsive | [119]        |     |
| PSBMA-PDPA| micelles      | CUR              | pH- and thermo-responsive | [127]         |     |
| CB       | PCBMA-ss-PCL-ss-PCBMA | micelles | DOX          | redox-responsive | [108]|
| PCBMA-PMAEL | micelles    | DOX               | redox-responsive | [111]          |     |
| PCBMA(DC-C) | NPs           | DOX              | pH- and redox-responsive | [120]         |     |
| PCBMA-DMAEMA@ZnO | NPs | DOX              | pH-responsive | [122]            |     |
| cRGD-PCSSL | NPs           | DOX              | redox-responsive tumor-targeting | [145]       |     |
| cRGD-PCSSD | NPs           | DOX              | pH-, redox-responsive tumor-targeting | [152]       |     |

Table 1 Summary of stimuli-responsive zwitterionic DDSs.

| Category | Poly(AAs) and polyptides | Working morphology | Loaded drugs | Special properties | Ref. |
|----------|--------------------------|-------------------|--------------|--------------------|-----|
| HHG2C18-L | liposomes | coumarin6/teosirinum | pH-responsive | [75,77]           |     |
| Lys-val-N, N-bis(acryloyl) dianmohexane | micelles | DOX          | pH-responsive | [76]             |     |
| M-HHG2C18-L | NPs            | Erlotinib/DOX     | pH-responsive | [79]             |     |
| PALHy-hyd-DOX | micelles | DOX              | pH-responsive | [92]             |     |
| Lysine methacrylate | micelles | CPT          | redox-responsive | [116]          |     |
| cRGD-PMIX | vesicles | DOX              | pH-responsive tumor-targeting | [146]       |     |
| CPMA | micelles | DOX              | pH-responsive | [50]             |     |
| MSN-HA-SiO2-TSA/DMA | NPs         | DOX              | pH-, redox-and HAase-responsive | [51]          |     |
| Z-MSN | NPs         | DOX              | pH-responsive | [52]             |     |
| BCM-41 | NPs         | LEVO            | pH-responsive | [81]             |     |
| dHAD | micelles | DOX              | pH-responsive tumor-targeting | [155]       |     |
have developed a series of pH-responsive zwitterionic carriers to protect drugs from non-specific adsorption in bloodstream, prevent cargo premature leakage during circulation and help drugs escape from endosomes/lysosomes rapidly with their morphological or chemical changes at target sites [59].

To design pH-responsive carriers, one of the most common strategies is using charge shifting polymers that would change in conformation or solubility according to the pH value of the surrounding environments. Armes and his partners synthesized a series of zwitterionic amphiphatic block copolymers based on poly(2-methacryloyloxyethyl phosphorylcholine) (PMPC) and the pH-sensitive poly(2-diisopropylamino)ethyl methacrylate) (PDPA) through atom transfer radical polymerization (ATRP) method [60–62]. The PDPA blocks would be protonated and hence hydrophilic under dilute acidic environment but become deprotonated when adjusting the solution to physiological pH value. This would lead to the self-assembly of PMPC-PDPA micelles in bloodstream, and loaded the water insoluble cargoes by the hydrophobic PDPA core [61]. When the micelles were endocytosed into cells, PDPA blocks would be protonated and cannot maintain the micelle structure, which was followed by quick drug release in the cytoplasm. By adjusting the polymerization degree of PMPC and PDPA, the drug release kinetics could be regulated, as well as their working morphology (micelles, vesicles, etc.) [61,63]. In fact, Armes and his co-workers also attempted to use PMPC-PDPA as novel non-viral vectors for gene delivery [64]. Through the electrostatic interactions between the cationic groups and negatively charged nucleic acids, genes are condensed by these zwitterionic polymers. And the MPC moieties could provide a “stealth” shell to protect the loaded pDNA from promiscuous absorption, and the DPA segments endowed the vectors pH-responsive capability [64]. It has been verified that about 90% of the loaded DNA could be released from the PMPC-PDPA, with pH switching from physiological level (pH 7.4) to endocytic (pH 6.0) level [65,66]. The PMPC-PDPA copolymer was also used to transfect siRNA to knockdown MDM2 gene in p53 mutant cells for cancer inhibition, and the animal experiment showed a significant effect in cell cycle arrest, apoptosis and growth inhibition [67].

Similarly, by replacing the DPA block with other tertiary amine containing monomers, numerous pH-sensitive zwitterionic polymers have been developed. For example, an amphiphilic triblock copolymer, poly(ε-caprolactone)-poly(diethylaminoethyl methacrylate)-poly(sulfobetaine methacrylate) (PSBMA-PDEA-PCL) has been synthesized [68]. The PDEA block worked as a pH-responsive switch, owing to its amphoteric nature by protonation-deprotonation brought from unsaturated nitrogen of tertiary amine [68,69]. However, PDEA is so sensitive to environmental pH that might influence stability of the core, and the cytotoxicity caused by its charge property also disturbed its application [70]. To solve this problem, PSBMA was combined with PDEA to maintain stable micelles and limit its toxicity, in the meanwhile to serve as a hydrophilic protective layer to resistant nonspecific biosorption. In addition, the unsaturated nitrogen of the imidazole group in histidine would be protonated to become hydrophilic, and the size of histidine containing self-assembled nanoparticles (NPs) will significantly increase due to the breakdown of the hydrophobic/hydrophilic balance [71]. Wu et al. respectively linked pH-responsive N-acetyl-L-histidine (NAcHis) and membrane-like PC groups to chitosan, a typical polysaccharide that can be disintegrated in vivo [72], to prepare NAcHis-PCCs conjugate [73]. The PC shell provided excellent antifouling property and biocompatibility to avoid adverse biological response. And since the pKa of the imidazole group in histidine is around 6.5, NAcHis would be protonated to be hydrophilic and destabilized the nanocarriers which eventually led to cargo release when the milieu pH < 6.5. By using Quercetin (QUE) as a model drug, the scientists found an acid-triggered behavior of NAcHis-PCCs that the release efficiency of QUE under acidic conditions (pH 5.5) was enhanced compared to physiological conditions (pH 7.4), which consisted well with the pH-responsive size changing behavior measured by dynamic light scattering (DLS).

Since some amino acids have extra amino/carboxy groups that can act as charge-reversing switch, researchers attempted to add AAs into zwitterionic polymers for pH-sensitive drug delivery [74,75]. For example, a pH-sensitive charge-reversal zwitterionic copolymer containing basic lysine, poly(Lys-alt-N,N-bis(acryloyl) diaminohexane) was synthesized through the Michael addition polymerization between N,N-bis(acryloyl) diaminohexane and lysine [76]. This amino acid-based copolymer could self-assemble into nanomicelles and show a slightly negative charge in blood but exhibit a positive charge in acid environments. Because of the high protonation of tertiary amino groups in the backbones and zwitterions of side chains in nanomicelles under the acidic condition, the hydrodynamic radius of the nanomicelles would increase thus to facilitate the encapsulated DOX diffusion at pH 5.0. Similarly, Takeoka and co-workers developed two amino acid-based lipids with different head groups, in which 1,5-dihexadecyl N-glutamyl-L-glutamate (L1) and 1,5-dihexadecyl N-N-diglutamyl-lysyl-L-glutamate (L2) were served as pH-responsive components to form charge-converse DOX-encapsulating liposomes [74]. The L2 moiety had two glutamic residues while the L1 had only one, both of the L1- and L2-liposomes exhibited good pH-responsibility as well as improved membrane fusogenic potential, but the DOX-loading efficiency of L1-liposomes was lower than the other ones for the distinction of head groups. Hence L2-liposomes was chosen to be intravenously injected to model rats, displaying comparable blood persistence and biodistribution to conventional phosphatidylcholine-based liposomes, but a higher toxicity to Hela cells.

Also, Zhang et al. synthesized a pH-responsive liposome based on zwitterionic oligopeptides that were able to convert the electrical property according to different pH in tumor matrix and intracellular compartments [75,77]. Besides common phosphatidylcholine and cholesterol in normal liposomes, this oligopeptide-based zwitterionic lipid (HHG2C18_L) employed glutamine acids (G), histidine (H) and pH-cleavable hexahydrobenzoic amide groups as its hydrophobic block, while stearyl alkane chains (−2C18H37) as the hydrophobic block. After the HHG2C18_L liposome arriving at tumor sites (pH 6.5) and convert surface charges from negative to positive ones, the cellular uptake could be enhanced via electrostatic interaction. And the imidazole groups of histidine promoted proton influx to endosomes/lysosomes, resulting in their bursting. As pH got lower, hexahydrobenzoic amide would be hydrolyzed, which brought about even stronger positive surface charge and escaped from endo/lysosomes to release the renal cancer-against drug temsirolimus eventually (Fig. 3). Furthermore, with the constant further studies on combination chemotherapy, “drug cocktail” has become an optimal therapeutic strategy for cancer treatment. Studies demonstrated that pretreatment of lung carcinoma cells using erlotinib could significantly synergize with DOX [78], but this combination needed a specific time lag between the drugs to maximize the effect, which asked an adaptive DDS to match the pharmacokinetic parameters for each drug and control their release sequence precisely. To fit for the “drug cocktail” method, Zhang’s group constructed a charge reversal erlotinib/DOX co-delivery nanoparticle (M-HHG2C18_L) by using their pH-sensitive lipid bilayer HHG2C18_L as a coat to cover amino-functionalized mesoporous silica nanoparticles (MSNs) [79]. They entrapped DOX in MSN-NH2 followed by coverage with erlotinib-carried lipid bilayer, so that the exterior EGF inhibitor would release faster than DOX. This pH-sensitive charge conver-
tion co-delivery system not only possessed all the properties of HHG2C18-L, but also achieved staggered sequential drug release without any damage to other major organs, showing great potential for cancer therapy.

In addition, a great number of mixed-charged zwitterionic materials are designed based on MSNs, which have good biocompatibility, tunable pore sizes, modifiable surface and remarkable drug loading capacity [80]. To improve drug loading efficiency and cellular uptake, a zwitterionic MSN (Z-MSN) was prepared by introducing a composition of carboxylic groups and quaternary amino groups as gatekeepers responding to tumor microenvironment [52]. The surface charge of Z-MSN would switch from negative to positive with cleavage of the acid-sensitive amide linkages, enhancing effective uptake of DOX into tumor tissues, leading to much higher antitumor effect compared with free DOX. And a Spanish team designed a mixed-charge zwitterionic MSN (MCM-41) whose external surface was decorated with short chains of aminopropyl silane triol and trihydroxy-silyl-propyl-methyl phosphonate mixed-charge brushes [81]. Levofloxacin (LEVO) was loaded into the pores of MCM-41, which displayed a remarkable antimicrobial activity against pathogenic E. coli and S. aureus in a low-fouling and reduced macrophage-uptake method.

Another strategy to design pH-responsive zwitterionic carriers is introducing acid-labile linkages (such as hydrazine, acetal and imine) that could be cleaved at acid environments but stable at neutral pH [55,82,83]. Liu et al. at first used a common aliphatic biodegradable polyester poly(D,L-lactide) (PLA) as hydrophobic segments to synthesize diblock copolymer PMPC-PLA by ATRP of MPC monomer with PLA-Br as the initiator [84]. PMPC-PLA vesicles were prepared via a dialysis method and DOX/DOX-HCl were loaded into the vesicle membrane and interior respectively. The results showed significantly faster drug release at pH 5.0 compared to pH 7.4, which might attribute to the hydrolysis of PMPC-PLA, resulting in morphology transformation from vesicle to micelle thus to extrude the drugs (Fig. 4) [85]. Later, they introduced pH-sensitive benzoyl imine linkage (Blink) in PMPC-PLA as a bridge to get a new diblock copolymeric micelle for PTX delivery. And the PLA block was ended by cinnamyl alcohol (CPLA) whose benzene rings could interact with PTX via π-π stacking to improve the loading content [86]. The in vitro study showed over 90% of PTX were released from the PMPC-Blink-CPLA micelles in 48 h at pH 5.5, which was ascribed to the cleavage of their acid-labile switch Blink.

Except for being used as block linkages, diverse polymer-drug conjugates (also called prodrugs) have been developed through connecting drugs and carriers with some suitable chemical bonds that can not only maintain the micelles structure, but also hold drugs firmly with neglectable drug leakage during blood circulation than traditional micelles [83]. In order to enable the prodrug response to environmental pH value, Wang et al. developed zwitterionic prodrug PMPC-PLL(DOX) using poly(L-lysine) as the functional segment to connect DOX via imine bonds. The PMPC-PLL (DOX) prodrg could self-assemble into core-shell structural stable micelles under physiological environment, but when
transferred to the acidic environment (pH 5.5), their imine bonds would be cut off to reverse surface charge so that contributed to the endosomal escape and accelerated drug release [87].

Similarly, quantities of prodrugs have been synthesized with the help of hydrazone bonds which are easy to be cut off in acidic environments too [88–91]. Wu et al. conjugated zwitterionic L-Lys and hydrazine grafted poly(aspartamide) (PALHy) by successive amination reactions of poly(succinimide) with L-lysine and hydrazine hydrate following hydrolysis [92]. The DOX molecules were conjugated to the copolymer chemically via hydrazone bonds to form PALHy–hyd-DOX conjugates. Performing as a hydrophobic moiety, DOX could trigger self-assembly of the nanoparticle in an aqueous solution, which resulted in hydrophobic interactions as well as π–π stacking to encapsulate more dissociative DOX molecules subsequently.

Taking advantages of the acid-labile characteristics of acetal segments, Wang et al. fabricated a pH-sensitive SB-modified dendrimer (Gn–SB) for antitumor drug delivery [93]. Dendrimers are regarded as a novel kind of promising micelle in the field of DDSs owning to the unique merits of uniform architectures, controllable sizes, abundant terminal groups, huge internal cavities, surface functionality and low intrinsic viscosity in solutions [94]. They selected polyacetal as the acid-labile segment for its ability to be hydrolyzed in an acidic environment with non-acidic metabolites, which makes it biodegradable [95,96]. And zwitterionic SB was conjugated to the tails of polyacetal to promote endocytosis by tumor cells for its high affinity to negatively charged cytomembranes in faintly acid conditions [60].

4.2. Redox-sensitive drug carriers

In addition to taking advantage of the pH difference, the usual stimulating factors include redox potential, temperature, light, specific enzymes and magnetic field [97–100]. Due to the overexpression of glutathione (GSH) in the intracellular components of tumor cells (10 mM) which could be much higher than that of the extracellular fluid and blood (10 μM), it is possible to achieve faster tumor intracellular drug release triggered by GSH [101,102]. And since disulfides could be easily reduced to sulfhydryl group under the reduction circumstance, disulfide linkages were widely utilized to build redox-sensitive therapeutic delivery systems [103].

At the outset, amphiphilic diblock polymers based on zwitterionic PMPC and biodegradable poly-ε-caprolactone (PMPC–PCL) with different polymerization degree were developed as substitutes for PEG–PCL. Both of them could load hydrophobic drugs efficiently, but PMPC–PCL micelles exhibited longer circulation time and higher accumulation rate by tumor cells attributing to the strong hydration and bioinspired effect of PMPC [104,105]. However, the peripheral dense shells prevent the enyzmolysis of PCL core, which is not conducive to drug release. Thereafter, scientists introduced disulfide bond as a bridge between PMPC and PCL moieties so that the new block polymer (PMPC–ss–PCL) could be broken up and unload cargoes rapidly in the presence of high GSH concentration in tumor cells. Cai et al. found an obvious size increase of the PMPC–ss–PCL micelles in presence of 10 mM dithiothreitol (DTT, another reducing agent to mimic intracellular tumor microenvironment) but no significant size change for 16 h in absence of DTT [106]. This phenomenon might happen because the disulfide bonds were cut off by DTT, so the PMPC shell pulled off from the micellar surface leading to micelle aggregation [107]. In this research, DOX–loaded PMPC–ss–PCL micelles showed faster drug release under reducing conditions in vitro, and much better anti-tumor effect than non-sensitive ones and positive DOX–HCl. Analogously, a reduction-sensitive triblock zwitterionic polymer PCBMA–ss–PCL–ss–PCBMA was fabricated by connecting hydrophilic PCBMA and lipophilic PCL groups with disulfide bonds [108]. Compared to a diblock polymer, this triblock one contains more disulfides and zwitterionic moieties in one unit, which may bring stronger anti-biosorption and redox–responsive properties.

Except for being used as linkages to build block polymers, disulfide bonds could also act as cross-linkers to form reduction-responsive cross-linking structures which have emerged as a feasible strategy to minimize premature drug release and facilitate unloading within tumor cells [109]. For instance, a biodegradable zwitterionic PSBMA nanogel was synthesized by one-step reflux precipitation copolymerization of SBMA and N,N'-bis-(acryloyl) cysteamine (BAC) [110]. DOX was held firmly by PSBMA nanogel in physiological conditions with only 7% leakage for 24 h, but 85% release in 8 h under a high GSH environment by the cleavage of the disulfide bonds in BAC. After intravenous injection in human hypopharyngeal carcinoma-bearing mouse models, the PSBMA nanogel showed excellent properties including high tumor accumulation, excellent biodegradation, long circulation, and sufficient drug release. And to enhance the stability of zwitterionic amphilic copolymers, Chen’s team synthesized a random copolymer P(CMBA-co-MAEL) by using zwitterionic carboxybetaine methacrylate as hydrophilic part and disulfide containing 2-(methacryloyloxy) ethyl lipoate as hydrophobic segment [111]. They first used Tris(2-carboxyethyl) phosphate to reduce the disulfides from MAEL, then stirred the mixture to form shell–cross-linked micelles via O2-mediated oxidation. DOX was loaded into the carriers before the micelles were cross-linked, and it proved that the PCBMA moieties provided excellent hemocompatibility and remarkable resistance to biosorption, with selective release of DOX in reducing environments owing to the cross-linked shell.

Besides, various polymeric prodrugs linked by disulfide bonds have already been tested for chemotherapeutics [112,113], and scientists also managed to connect therapeutic drugs to zwitterionic polymers by disulfide linkers directly to gift zwitterionic prodrugs the reduction-responsive properties. Temozolomide (TMZ) is a first-line drug for glioblastoma, but it is hydrolytic instability and is impeded by enzyme-mediated chemoresistance after crossing the blood brain barrier [114]. To improve its stability in aqueous solution, Emrick et al. synthesized a battery of PMPC–TMZ conjugates that showed enhanced drug stability as well as much longer half-life times [115]. Afterward, they expanded this delivery platform by introducing disulfide linkers to trigger TMZ release efficiently under high GSH intracellular tumor condition. And another broad-spectrum antitumor drug camptothecin was conjugated to the lysine residue side chains of zwitterionic poly(lysine methacrylate) via GSH-sensitive disulfide bonds to form a reduction sensitive polymer-drug conjugate [116]. In the meanwhile, the authors linked anisamide, a sigma-1 receptor to this polymer as a guide to target cancerous endothelial cells, but the ligands utility in tumor targeting is now controversial [117].

In fact, scientists soon came up with the great idea of preparing “smart” amphoteric delivery systems that can respond to more than just one stimulus to help therapeutics work more effectively [118]. The first pH and redox dual-sensitive zwitterionic DDS was synthesized by linking pH-responsive PDEA and PCL via redox-sensitive disulfide bonds, and the SB segment was added to gain the zwitterionic copolymer PCL–ss–PDEASB [119]. This amphiphilic copolymer could self-assemble into micelles and have long circulation in blood as well as effective internalization by cells owing to the SB shell. After endocytosis, the micelles could not only release the encapsulated DOX which was triggered by the protonation of PDEA groups in acid condition, but also unload the drugs quickly with the reduction of disulfide bonds in respond to intracellular high GSH concentration. Hence, the drug-loaded PCL–ss–PDEASB micelles were able to release lipophilic antitumor drugs fast under acidic and reductive conditions simultaneously to kill tumor cells.
efficiently, and the modification with SB made it possible even to eliminate the cytotoxicity of drug carriers with PEDA (Fig. 5).

Through a different approach, another typical pH- and redox-sensitive DDS was designed to cover peptide dendrimer-modified carbon dots by PCBMA through the interaction between the terminal guanidine on arginine dendrimers and carboxyl groups on PCBMA [120]. In this work, PCBMA not only provided “stealth” delivery in blood, but also achieved electric reverse after enriching at tumor sites. The zwitterionic coat would thus be taken-off from the PCBMA(DC-C) to expose the abundant peripheral guanidine groups with positive charges, which were beneficial to enhance endocytosis across the negative charged cell membranes. Eventually, the highly reductive and acid condition in tumor cells would cut-off the disulfide bonds between peptide dendrimer and carbon dots, desorbing DOX/CHCl to accelerate the nucleus-specific drug release for cancer therapy. And PCBMA has also been utilized to decorate quantum dots to improve their properties for drug delivery. Just like ZnO quantum dots which can be decomposed into zinc ions completely in the aqueous solution with a pH of 5.0 thus become toxic to tumor cells [121,122]. But their water insolubility and ease of agglomeration require antifouling modifications, so PCBMA-poly(2-(dimethylamino) ethyl methacrylate (PDMAEMA)) PCBMA-PDMAEMA was employed as a pH-sensitive coat to compensate for their shortcomings [122].

4.3. Thermo-responsive zwitterionic vehicles

Thermo-responsiveness is one of the most widely used stimulus that can lead to reversible phase transition with just a slight change of temperature, so that thermo-sensitive DDSs have attracted much attention to control drug release [123]. As mentioned in the classification chapter, SB-based polymers such as PSBMA exhibit a sharp phase transition below their UCST in aqueous solution owing to the strong electrostatic interactions among their SB moieties [124]. That means the combination of zwitterionic SB and other suitable non-ionic segments might achieve thermo-responsive carriers which could transit from soluble to insoluble through adjusting the environmental temperature.

Through free radical polymerization, Chen and co-workers prepared dually responsive copolymers consisting of thermo-responsive SBMA and pH-sensitive DPA copolymer for drug oral delivery (P(SBMA-co-DPA)) [125]. In this work, SBMA was selected owing to its strong electrostatic attraction between their ammonium cation and sulfo-anion, which made the zwitterionic copolymer self-aggregate into NPs in aqueous media within their UCSTs [124], and thus to load the drug curcumin (CUR) [126]. When the temperature increased above the UCST, collapsed associations of polymer chains would transit to the unimer state to facilitate drug release. Interestingly, the authors found the UCSTs of P

Fig. 5. The forming process and intracellular drug release of the PCL-ss-PDEASB/DOX system. With the shield of zwitterionic SB shell, these micelles could avoid undesired absorption, and internalized by tumor cells effectively. After endocytosis, the loaded DOX would be release rapidly as the result of PDEA protonation in response to low pH and disulfide bonds cleavage triggered by cytosol high GSH concentration.
(SBMA-co-DPA) varied with different solution pH and exhibited a pH-dependently thermal-responsive behavior, that might be caused by the protonation/deprotonation of DPA segments. When the environmental pH value was below the pKa of DPA moieties, the protonated amino groups brought positive charges to the copolymer which might introduce strong electrostatic repulsion to diminish the attractive interaction among SB moieties. That means the more content of DPA segments, the lower UCSTs of P(SBMA-co-DPA) would be achieved at the same pH value. So they tried different molar ratios of SBMA and DPA and results showed P(SBMA90-co-DPA10) could be well-dispersed at extremely acidic condition like stomach (pH 1.2–3.0), but at neutral pH such as intestine (pH 6.5–8.0) the NPs would be solvated as unimer and unloaded CUR since their UCST became lower than body temperature [127]. Thus, these pH-sensitive and thermo-responsive zwitterionic copolymers PSBMA-PDPA are promising to be applied as oral delivery drug carrier.

4.4. Tumor targeting zwitterionic carriers

Achieving high uptake rate by tumor cells is still a challenge demanding prompt solution for the development of antitumor drugs’ delivery which could improve the therapeutic efficiency and limit side effects as possible. To enhance the specific accumulation of drugs in tumor sites, various tumor target-specific moieties are conjugated to the surface of carriers, such as folic acids (FA) [128,129], mucopolysaccharide [130], peptides [131–133], and antibodies [134]. Exceptionally, these target-specific ligands have also been widely introduced to zwitterionic vehicles so that they can be recognized by the corresponding receptors that are overexpressed on the surface of specific tumor cells.

Since FA shows extremely high affinity for the folate receptor (FR) that is known to be overexpressed on the membrane of malignant cells include ovary, cervical and nasopharyngeal carcinomas [135], FA-decorated carriers could promote cellular uptake for FR-positive cancer cells. Therefore, Armes’ team conjugated FA to the primary amine terminus of the PMPC-PDPA to endow this pH-sensitive diblock copolymer tumor targeting property [136]. They first prepared PMPC-PDPA through ATRP using an Fmoc-protected initiator and treated the purified Fmoc-MPC-DPA with they first prepared PMPC-PDPA through ATRP using an Fmoc-protected initiator and treated the purified Fmoc-MPC-DPA with and FAMEG [141]. Thus, the content of FA groups in this copolymer (FAMEG) was first synthesized by a two-step synthetic strategy of FA-modified poly(ethylene glycol) methacrylate macromonomer (FAMEG) was first synthesized by a two-step synthetic strategy [140]. The copolymer FAMEG-MPC-PCL was then fabricated through free radical polymerization of PCL macromonomer, MPC, and FAMEG [141]. Thus, the content of FA groups in this copolymer could be expediently adjusted via changing the molar ratio of monomers.

The arginyl-glycyl-aspartic acid (RGD) is an oligopeptide that expresses on many extracellular matrix proteins and plays a significant role in cell direction and cell linkage [142]. RGD could selectively recognize and bind to the $\alpha_v \beta_3$ class of integrins, which are overexpressed during tumor progression and have been regarded as tumor markers [143]. For this reason, different forms of RGD include liner and cyclic structures were widely investigated for tumor targeting [144], but the cyclic ones exhibited better activity attributing to a conformation-less assembly that was able to withstand proteolysis and had higher affinities to interact with integrin receptors [140], so that lots of cyclic-five-member-ring RGD contained pentapeptides (termed cRGD) were utilized to endow drug vehicles tumor-targeting properties. By combining cyclo(Arg-Gly-Asp-D-Phe-Lys) with PCB-ss-PCL, Huang et al. developed a multifunctional zwitterionic block copolymer cRGD-PCSSL [145]. DOX was encapsulated by the biodegradable PCL segments and the zwitterionic PCB shell would resist nontarget-specific absorption after intravenous injection to prolong circulation time. Under the guiding of cRGD ligands, the DOX-loading nanoparticles accumulated around cancer cells and would be endocytosed into cells mediated by $\alpha_v \beta_3$ integrin receptor (Fig. 6). Afterwards, disulfide bonds were cut off for reductively drug release.

On the other hand, since cRGDs are class of pentapeptides, using them to modify zwitterionic polypeptides can achieve unique biomimetic properties as well as tumor targeting function. Given this, Lin and his partners synthesized a cRGD-conjugated polypeptide composed of glutamic acid and lysine to mimic albumin [146]. Because the pair of Glu-Lys has been found as the most abundant AA pairs in albumin, they were chosen for a longer half-life circulation time [147] taking albumin-bound paclitaxel (Abraxane) as a successful precedent [148]. The negative charged DOX was encapsulated in cRGD-poly(L-glutamic acid-co-L-lysine) (cRGD-PMIX) by electrostatic interaction via tuning the ratio of Glu to Lys over 1:1 and the DOX-loaded vesicle was formed through an emulsion solvent evaporation technique. This targeting-functionalized vesicle exhibited pH-sensitive drug release behavior and the existing of trypsin obviously promoted the release of DOX which has been demonstrated to be expressed in kinds of cancers during proliferation, invasion and metastasis [149,150].

In order to limit drug premature leakage during blood circulation, scientists always take advantage of disulfide bonds to build cross-linked shell. Based on this, Huang et al. designed and synthesized a triblock zwitterionic copolymer that was composed of the disulfide supplier polyN-(2-(2-pyridyl disulide) ethyl methacrylamide (PDS), the ultra pH-sensitive PDPA core to hold DOX (151) and zwitterionic PCBMA corona to stabilize the nanoparticles and to prolong half-life [152]. The triblock copolymer (PCBMA-PDS-PDPA, called PCSSD) was tumor-targeting functionalized with RGD, so the RGD-PCSSD shell-cross-linked NPs could be guided to tumor sites with negligible biosorption and less premature drug release during systemic circulation. The pH and redox dual-sensitivity of the RGD-PCSSD copolymer was illustrated in Fig. 7. After receptor-mediated endocytosis, the PDPA segment would be protonated to promote escape from endosomes/lysosomes, and the shell linkers would be cut off in respond to GSH with high concentration to facilitate drug release in an intracellular reductive environment.

Hyaluronic acid (HA) is another popular tumor-targeting ligand that performs strong affinity to CD44 which is overexpressed on the surface of many kinds of cancer cells and is associated with tumor progress, invasion and metastasis [153,154]. Attributed to the carboxyl groups and acetamide groups in HA, Gao et al. prepared a pH-sensitive mix-charged zwitterionic polymer $\mathbf{dHAD}$ by deacetylation of HA and grafting dodecylamine onto it [155]. The polymer exhibited negative charge at pH 7.4 and became positive at pH 6.2 which led to an enhanced uptake rate by tumor cells in acidic environment [156]. And since HA is a natural glycosaminoglycan that could be degraded completely by hyaluronidase (HAase) in cells [157], $\mathbf{dHAD}$ can provide a choice to form biodegradable DDSs while traditional zwitterionic polymers could not.
**Fig. 6.** The tumor targeting cRGD-PCSSL micelles guided by cRGDfc peptide. The functional zwitterionic micelles could accumulate around tumor cells leading by cRGD ligands. Through the interaction between cRGD and \(\alpha_v\beta_3\) integrin receptor, the uptake rate of micelles by tumor cells would be enhanced.

**Fig. 7.** The reduction and pH dual-responsive behavior of the tumor targeting RGD-PCSSD shell-cross-linked nanoparticles. When the pH value was 7.4, the liner RGD-PCSSD copolymers could assemble into NPs using the hydrophobic PDPA core to encapsulate DOX. Treating with ultrasound promoted the formation of disulfide bonds to get compacted cross-linked NPs and hold DOX tightly. When the pH value decreased, the PDPA moieties would become protonated which made the NPs swell and a part of drug leak slowly. And as the concentration of GSH in the environment increased, the cross-linked disulfide bonds would be cut off to disrupt the whole system and release the cargoes rapidly.
It is interesting to note that Jin et al. developed a triple-layered MSN which used HA as a switch to trigger drug release in response to high concentration HAase, rather than for its tumor targeting [51]. Above the first HA-layer, disulfide bond-embedded silica was utilized as the second GSH-sensitive layer, which was covered by a mix-charged zwitterionic shell (Fig. 8B). This zwitterionic surface consisted of trimethoxysilylpropyl-N,N,N-trimethylammonium chloride (TSA) and 2,3-dimethylmaleic anhydride (DMA), and the net charge of MSN-HA-SiO$_2$-TSA/DMA could switch from neutral to positive in the acidic environment surrounding tumor cells. Mix-charged zwitterions located on the surface of MSN could work as gatekeepers to hold drugs and open the “doors” at expected sites (Fig. 8A) [52]. More than that, the HA molecules were labeled by fluorescein isothiocyanate (FITC) which made it possible for real-time drug tracing and selective tumor cell imaging.

5. Zwitterionic materials applied in gene delivery

As the studies continue persistently, gene therapy has become a predominant strategy for the treatment of genomic and non-genomic diseases [158–160]. So far, many therapeutic genes are in clinical trials, and some have even been approved for marketing [161,162]. However, to send the therapeutic genetic molecules to the desired sites and ensure they could transcript and translate into functional proteins successfully, safe and effective carriers are indispensable [163–165]. The available gene delivery vectors at present generally include viral vectors, non-viral cationic polymers and inorganic metal materials [166–168]. Although the viral vectors have showed relatively higher gene delivery efficiencies, their immunogenicity makes repeated administration impossible which seriously obstructs the application in clinic. And the risks of insertional mutation as well as difficulties in mass production also limit their application. While cationic polymers can condense negatively charged gene molecules and facilitate cellular uptake of cell membranes, but positively charged macromolecules have considerable cytotoxicity and prefer to interact with other biomolecules in blood [169].

Due to the versatile capabilities, zwitterionic materials also exhibit great potential for gene delivery. The copolymers contain cationic segments and zwitterionic parts are expected as ideal vectors which inherit the gene-carrying ability of cationic vectors via the interactions with genetic molecules, while the zwitterionic moieties provide a “stealth” surface and stabilize the gene/polymer complexes (polyplexes) sterically [165,170]. The vinyl-based cationic monomers used to build cationic gene vectors always have a linear/branched structure with monomer unit sequences, such as the derivatives from methacrylamine and methacrylate (shown in Fig. 9) [171]. Most often, zwitterionic molecules are conjugated with these derivatives to form novel gene delivery vectors, and some other types take a zwitterionic coat on the surface of traditional non-viral vectors to endow them with antifouling properties.
and enhance their stability at meanwhile. Here, the novel gene vectors involved with zwitterionic materials are summarized and displayed in Table 2.

### 5.1. Zwitterion-decorated cationic gene vectors

In the early days, researchers tried to conjugate cationic gene carriers with anionic materials, especially negative charged ligands that are capable of targeting cells to form mix-charged zwitterionic gene carriers. It was reported that conjugating HA to polyethyleneimine (PEI) could reduce the toxicity which was associated with their cationic amine groups, and PEI-HA could promote DNA transfection via the interaction between HA and CD44 on cell surface [172–174]. However, the DNA condense levels were still decreased with the neutralization of cations. To enhance system stability, limit cytotoxicity but not sacrifice loading capacity, scientists turned to zwitterionic materials for inspiration and innovation.

PC-based cationic polymers have been demonstrated as efficient gene delivery vectors. Ever since the first phospholipid polymer, poly(MPC-2-aminoethyl methacrylate (AEMA)) (PMPC-PAEMA) was developed and applied as a plasmid DNA (pDNA) vector, and quantities of MPC containing polymers were developed for gene delivery [175]. In order to alleviate the non-specific intertwining of vectors and gene molecules, Kitagawa et al. synthesized a battery of micelle-like amphiphilic copolymers, consisting of MPC, N-(3-(dimethylamino)propyl) acrylamide (DMAPAA) and stearyl acrylate (SA) [176]. With the protection of DMAPAA-MPC-SA, the expression level of transfected pCMV-Luc (a plasmid encoding luciferase gene) was proved to be three times higher than that carried by liner-type polycations.

As mentioned in the section of drug delivery system, PMPC-PDPA diblock polymers were utilized as zwitterionic pH-sensitive vectors to load pDNA and siRNA molecules [64–67]. The MPC could be copolymerized with any vinyl monomer through a common radical polymerization method [177], by replacing the cationic block DPA with other tertiary amine containing monomers, PMPC-PDEA was copolymerized [64,178,179]. Apart from condensing DNA effectively and stabilizing the colloidal size complexes sterically, all of these MPC-based are probable to incorporate with targeting ligands to prepare specific targeting gene vectors. For instance, FA as a frequent-used ligand has been introduced into PMPC-PDMAEMA and PMPC-

### Table 2

Summary of gene vectors associated with the zwitterions.

| Category          | Polymers                  | Loaded gene | Ref.          |
|-------------------|---------------------------|-------------|---------------|
| Phosphorylcholine | PMPC-PDMAEMA              | pDNA        | [64,180]      |
|                   | PMPC-PDPA                 | pDNA, siRNA | [65–67,181]   |
|                   | PMPC-PAEMA                | pDNA        | [175]         |
|                   | PMPC-PDMAPAA-PSA          | pDNA        | [176]         |
|                   | PMDN-HBs                  | pDNA        | [177]         |
|                   | PMPC-PDEA                 | pDNA        | [178,179]     |
|                   | P/(MPC-co-AEMAm)-b-MPC    | pDNA        | [183]         |
| Sulfobetaine      | PMPD5SAH-PDMAEMA          | pDNA        | [184]         |
|                   | PEI-g-PHEAa-b-MEO2MA      | pDNA        | [185,186]     |
|                   | PEI-g-PHEAa-b-MPD5SAH     | pDNA        | [185,186]     |
|                   | PSBMA-PAA                 | pDNA        | [187]         |
|                   | ZAlS                      | Cas9 mRNA   | [189]         |
|                   |                          | and sgRNA   |               |
|                   | MPDSAH-MEO2MA-MPD5SAH     | pDNA        | [197]         |
| Amino acids       | GEA                       | pDNA        | [188]         |
| Carboxybetaine    | FC20-DSPE                 | siRNA       | [191,192]     |
|                   | CBAA-PAMAMA               | pDNA        | [196]         |
|                   | P(CBMA-EE)                | pDNA        | [198]         |
|                   | PCB-NBE                   | pDNA        | [199]         |
PDPA diblock copolymers, and the in vitro experiments with FA receptor overexpressing cells [180,181]. Besides, Chiba et al. copolymerized the poly(MPC-DMAEMA-p-nitrophenylcarbonylox yethyl methacrylate (NPMA)) (PMDN) as a frame for hepatocyte-specific gene delivery. The NPMA was introduced to immobilize hepatitis B antigen (HBs) so that the PMDN-HBs could transfer therapeutic genes into human hepatocytes specifically [177]. According to the in vivo test, intravenous injection of pDNA/PMDN-HBs did not lead to liver dysfunction or organ weight changes owing to the biocompatibility offered by MPC moieties. More importantly, the specificity of PMDN could be conveniently altered through replacing the liver-targeting HBs with other biorecognition molecules.

Moreover, it was reported that statistical carbohydrate-based copolymers could enhance the expression of loaded genes with low toxicity, but were easier to aggregate in the presence of serum proteins than block copolymers [182]. Therefore, MPC was introduced to form a “block-statistical” copolymer poly(MPC-2-aminoethyl methacrylamide) (PMPC-PAEMAm), exhibiting high gene expression, relatively low-serum protein interaction and low toxicity in hepatocytes and human dermal fibroblasts [183].

Similarly, zwitterionic SB and CB can also perform like PC to improve the cationic gene vectors. As mentioned earlier, the PMPC block could benefit the steric stability of the PDMAEMA/DNA complexes but sacrificed the transfection performance just like the PEGylated cationic vectors. Fortunately, the comb-shaped gene vectors consisted of dextran backbone and diblock copolymeric PCB-PDMAEMA or PSB-PDMAEMA sidechains were demonstrated to work better than PMPC-PDMAEMA, with enhanced DNA transfection as well as high biostability [165]. Liu and co-workers also developed a PSB-cationic methacrylate copolymer poly(N-(3-(met hacylamido)propyl)-N,N-dimethyl-N-(3-sulfo-propyl) ammonium hydroxide (MPDSAH)-b-DMAEMA) [184], and showed a much higher gene transfection efficiency than PMPC-PDMAEMA, which might be caused by the shell charges of MPC block obstructed the cellular internalization, whereas the SB containing MPDSAH remained the improvement of cellular uptake. But the transfection efficiency of MPDSAH-PDMAEMA was still lower than cationic PEI (25 k). Since the high molecule weight PEI (25 k) showed high gene transfection ability but high toxicity at the same time, while the low molecule weight (LMW) PEIs were just the reverse, they attempted to graft poly(MEO-2-

**Fig. 10.** Schematic diagram of PSBMA-PAA/PDMAEMA (or PEI)/DNA core–shell polyplexes. Cationic PDMAEMA (or PEI) first interacted with DNA molecules electrostatically. In physical condition, the diblock polymer PSBMA-PAA served as electroneutral shells to protect the polyplexes from biosorption and fell off around tumor cells to expose the positive charges of the core for enhanced cellular uptake.
Au DENPs display few macrophage cellular uptake to extend their escape from lysosomes via proton sponge effect. And CBAA made terminal amines, PAMAM could condense pDNA strongly and thylated in cancer 1 (HIC1) pDNA [196]. Because of the abundant functionalized Au DENPs as a vector for the delivery of hyperme-
MPDSAH and "B" was PEG analog MEO 2MA with LCST behavior zwitterionic triblock polymers via ATRP, in which "A" represented sizing an double thermo-responsive SB-based "ABA" structured nucleotide molecules, Dai et al. came up with an idea that synthe-
have also participated in therapeutic gene delivery directly.

densation of pDNA and compensate the positive charges of amines to preserve the 3D spherical shape of dendrimers for better con-
denstion of that, they successfully developed a novel photolabile "switch" for active degradation control [199]. Inter-
estingly, in order to maintain the unspecific protein resistance property of PCB, Chen et al. used the polyzwitterionic precursor PCBMEE as a hydrophobic segment to entrap pDNA as well as guard them against the nuclease attack, and the zwitterionic PCBMMA shell was added to improve water-solubility of the core and kept gene form protein in serum [200]. Combining all of the merits, the block polymer PCBMMAE-PCBMA showed high transfection efficiency even to some hard-to-transfect cells like HUVEC, and effective intracellular gene release as well.

6. Summary and outlook

On account of the super hydration ability arising from the ion–di-
pole interaction among zwitterions and water molecules, zwitte-
rionic materials are gifted with remarkable antifouling property to prevent the adsorption of bacteria, cells and biological macro-
molecules [15]. And the antifouling property, low cytotoxicity, great biocompatibility, negligible immunogenicity, in vivo stability as well as long circulation time, together make the zwitterions a great choice for drug and therapeutic gene delivery. With the development of multi-functional zwitterionic materials, stimuli-
responsive and tumor-targeting ones have been fabricated, which further makes the zwitterions an ideal option for clinical applica-
tion. Via introducing corporate segments that are protonated in acid environment or pH-sensitive chemical bonds, zwitterionic carriers could respond to environmental pH value to enhance cel-
lular uptake and drug release efficiency; via introducing disulfide bonds which are easily reduced by GSH with high concentration, zwitterionic vectors are gifted redox responsiveness to unload cargoes rapidly in cytoplasm of tumor cells; via utilizing zwitterionic copolymers that are composed of SB-based moieties and adjusting to the fit molar ratios, the SB-decorated delivery systems could be thermo-responsive and transfer from soluble to insoluble hence to release the therapeutics. And the participation of targeting mole-
cules endows zwitterionic carriers with cell and tissue specificity.

Besides, when work as vectors for gene transfection, zwitterions could be used to decorate traditional cationic gene vectors to relieve their cytotoxicity and maintain their ability to hold gene molecules as well as their high cellular uptake efficiency. On the other hand, zwitterionic materials could also serve as gene carriers directly in a charge reversal manner.

Although each type of zwitterions displays outstanding antifouling ability and has been utilized to build therapeutic vec-
tors, there are of course some differences among diverse cate-
gories. Since PC is a component of cell membranes, PC-based zwitterions are the earliest to be investigated in this field, and they are also the most ones with relatively thorough study. PC-based biomimetic zwitterionic carriers always show great biocompatibil-
ity, but the high synthetic cost might be a fatal problem that limits their application. As for SB-decorated carriers, the most prominent advantage should be the UCST, which is useful to designed thermo-
responsive zwitterionic polymers. The UCST is influenced by molar radio and concentration of ions in environment, so adjusting these parameters to fit is necessary. Attributing to the abundant carboxy groups, CB-based vectors are easily functionalized with versatile ligands or biomolecules hence to develop “smart” delivery systems according to demands. And zwitterionic poly(amino acids) or polypeptides are another natural biomimetic materials that can be degraded totally by enzymes through metabolism.

In early days, the patents of zwitterionic materials are mainly about detergent, antifouling coatings and polymer compounds with producing methods thereof. But as the expectations for the safety and healthy become higher, more zwitterionic materials have been granted gradually for DDSs and implant materials in last

escape capability of the lipoplexes [191]. Of course, the PCB modi-
fied cationic liposomes could also avoid accelerant blood clearance thus to extend the in vivo circulation time and increase tumor accumulation of therapeutic siRNAs after injection [192]. Recently, PCB-DSPPE micelles were also prepared for oral delivery of insulin which were facilitated to penetrate across the epithelial cell layer owing to the biological properties of carbboxybetaine [193]. This novel zwitterionic platform might improve the life quality of dia-

5.2. Gene carriers based on zwitterionic materials

Apart from being used to modify available cationic gene vectors to afford their antifouling abilities, various zwitterionic materials have also participated in therapeutic gene delivery directly.

Since the charge neutralization of cationic polymers tended to cause precipitate of complex after condensing negative charged nucleotide molecules, Dai et al. came up with an idea that synthe-
sizing an double thermo-responsive SB-based “ABA” structured zwitterionic triblock polymers via ATRP, in which “A” represented MPDSAH and “B” was PEG analog ME0 2MA with LCST behavior [197]. The SB-based MPDSAH acted to condense DNA molecules via their quaternary ammonium cations, while their sulfonic anions were unreacted thus to prevent from the inter-complex coacervation. Since the positive and negative charges of MPDSAH were separated by three methane groups, the spatial barrier was weakened so that DNA could be complexed with MPMDSAH effectively. But with the addition of DNA molecules, the ion pairs in MPDSAH would be disrupted which resulted in the disappearance of their UCST, while the LCST still existed with slightly increasement.

In addition, zwitterionic carbboxybetaine also shows great po-
tential in gene vector designing thanks to their functional car-
boxyl groups. Once the COO− groups of PCBs are esterified, the polymers left cationic thus could condense nucleic acid molecules just like the cationic vectors. After ester hydrolysis and exposure of the cationic COO− groups, the gene molecules would be released efficiently. Esterified with the ethyl ester, Jiang et al. developed the CBMA-ethyl ester polymers PCBMEE with anions hidden by the ester bond partially, so that to render cationic for DNA loading until hydrolysis [198]. They replaced the quaternary ammonium with quaternary (4')- tertiary (3')- and secondary (2')-amine respectively and found a 3:1 mixed copolymer of 3' and 4' CBMA-EE monomers allowed the polyplexes a good size and sur-
face charge to promote transfection and gene expression. On basis of that, they successfully developed a novel photolabile o-nitrobenzyl ester of PCB (PCB-NBE) to endow this platform a

UV-sensitive “switch” for active degradation control [199]. Inter-

stably with cells, Wang et al. modified generation 5 (G5) PAMAM dendrimer with carbboxybetaine acrylamide (CBA), which showed barely changes in radius of hydration [194]. And entrapp-
ing gold nanoparticles into PAMAM dendrimers has been proved to preserve the 3D spherical shape of dendrimers for better con-
densation of pDNA and compensate the positive charges of amines partly to alleviate the cytotoxicity [195]. Therefore, Shi et al. first synthesized CBA and covalently grafted CBA to the surface of G5 PAMAM dendrimers to entrap gold NPs, and finally got CBAA-
functionalyzed Au DENPs as a vector for the delivery of hyperme-
thylated in cancer 1 (HIC1) pDNA [196]. Because of the abundant terminal amines, PAMAM could condense pDNA strongly and escape from lysosomes via proton sponge effect. And CBAA made Au DENPs display few macrophage cellular uptake to extend their circulation time.

PCB, Chen et al. used the polyzwitterionic precursor PCBMAEE as a hydrophobic segment to entrap pDNA as well as guard them against the nuclease attack, and the zwitterionic PCBMA shell was added to improve water-solubility of the core and kept gene form protein in serum [200]. Combining all of the merits, the block polymer PCBMAEE-PCBMA showed high transfection efficiency even to some hard-to-transfect cells like HUVEC, and effective intracellular gene release as well.

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decades. The classic example PMPC-PDA have been granted numbers of patents as carriers for drug or gene worldwide, and the patent for siRNA delivery with long-term circulation is going to be published in 2020 (US20200171169A1). The recent trend in relevant patent filings indicates that more and more interests are focused on the development of multi-functional zwitterionic vectors. But while these novel materials have exhibited great potential in the field of biological medicine, studies either on their structures or characters are just in the primary stage, there is still a long way to go to achieve clinical application. One of the most pressing problems is to simplify the synthesis process and reduce the costs which must be faced in the mass production. Further studies are asked to overcome this crucial challenge in the near future. Meanwhile, with the increasing demands for precise control of drug location and release, the application of multiple stimuli-responsive DDSs is an irresistible trend, which can not only send the drugs to desired destinations with few premature leakage, but also unload them at the exact sites (even at the expected cell-organelle) to limit side effects and ensure the therapeutic effect. In addition, since the quantity of zwitterionic gene vectors at present is obviously less than carriers for drugs, it might suggest a much huger development space and brighter future for versatile zwitterionic materials utilized for therapeutic genes’ delivery. From this prospective, more attention might be paid to enhance the transfection efficiency of therapeutic genes with the help of zwitterionic delivery systems. All in all, in order to achieve clinical application, more investigations are needed to improve zwitterionic materials’ performance in biocompatibility, biodegradability, drug loading capacity, low cytotoxicity, environmental stimuli-sensitivity, specific targeting ability and some other peculiarities in light of requirements.

CRediT authorship contribution statement

**Ling-Yan Zhou:** Data curation, Formal analysis, Writing - review & editing.  
**Yang-Hui Zhu:** Data curation, Writing - original draft, Software.  
**Xiao-Yu Wang:** Investigation, Methodology.  
**Chao Shen:** Methodology.  
**Xia-Wei Wei:** Supervision.  
**Ting Xu:** Conceptualization, Funding acquisition, Project administration, Supervision.  
**Zhi-Yao He:** Conceptualization, Funding acquisition, Methodology, Validation.  

**Declaration of Competing Interest**

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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