Review Article

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Phenylboronic acid-decorated polymeric nanomaterials for advanced bio-application

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Abstract: The paradigm of using phenylboronic acid-decorated polymeric nanomaterials for advanced bio-application has been well established over the past decade. Phenylboronic acid and its derivatives are known to form reversible complexes with polyols, including sugar, diol and diphenol. This unique chemistry of phenylboronic acid has given many chances to be exploited for diagnostic and therapeutic applications. This review highlights the recent advances in fabrication of phenylboronic acid-decorated polymeric nanomaterials, especially focus on the interactions with glucose and sialic acid. Applications of these phenylboronic acid-decorated nanomaterials in drug delivery systems and biosensors are discussed.

Keywords: Phenylboronic acid; Polymeric nanomaterials; Polyols; Drug delivery

1 Introduction

Phenylboronic acid (PBA) and its derivatives are known to reversibly form cyclic boronic esters with polyol compounds [1, 2], hence they could recognize carbohydrates, such as glucose in blood and sialic acid (SA) on cancer cells. Carbohydrates are the foundation of life and play irreplaceable roles in a variety of cellular events, including proliferation, recognition, intercellular communication and immunoregulation. When disease occurs, the type and expression amount of carbohydrates would change. The unique chemistry of PBA with polyol compounds, especially carbohydrates, provides a way to design PBA-based polymeric nanomaterials. As polymeric nanomaterials could greatly enhance molecular interactions by multivalent effects, they have attracted much attention in biomedical applications [3–5], such as drug delivery systems and biosensors for diagnosis and therapeutics of diseases [6].

The most typical application of PBA and its derivatives is focused on glucose-responsive materials to recognize blood glucose and delivery insulin for the treatment of diabetes mellitus [7–10]. However, PBA-based glucose-responsive materials could only work in alkaline media but not at physiological pH, due to the high $pK_a$ value ($pK_a > 8.0$) of PBA moiety [11–14]. Numerous investigations have been focused on the design and synthesis of PBA-based polymers to decrease $pK_a$ of PBA moiety and increase the glucose-sensitivity as well as effectively release insulin at physiological pH [15–18]. Interestingly, PBA and its derivatives can stably bind with SA in acid conditions, such as tumor acid microenvironment, resulting in another application of PBA-based polymeric nanomaterials to construct anticancer drugs delivery systems [19–21].

Herein, we review the recent advances in fabrication of PBA-decorated polymeric nanomaterials based on the specific interactions with carbohydrates, and highlight the biomedical applications in drug delivery systems and biosensors.

2 PBA-based glucose-sensitive polymeric nanomaterials

Insulin-dependent diabetes mellitus characterized by accumulating glucose concentrations in blood has been a major global health epidemic [22, 23], leading to a highly need to design glucose-responsive materials for self-regulated insulin release responding to the variation of blood glucose concentrations. PBA-based glucose-responsive polymeric materials have been well-investigated because they are versatile enough to design different formulations and more stable than protein-based systems (glucose oxidase and concanavalin A).
The interaction between PBA and glucose is reversible equilibrium, which lays the foundation for the glucose-responsiveness of PBA-based materials. PBA-based glucose-sensitive nanomaterials could be fabricated by different approaches, such as self-assembly of amphiphilic copolymers, polymerizing into nano/microgels, and conjugating with inorganic hybrid nanoparticles, etc. [25–31].

As most of the PBA-based materials reported demonstrate maximum glucose-sensitivity at pH 9 ~ 10, significant efforts have been focused on decreasing their $pK_a$ values and enhancing the sensitivity to glucose at physiological conditions. Sumerlin group prepared macromolecular stars with boronic ester linkages to self-assemble into nanoparticles [25]. The boronic esters were formed between 3-acrylamidophenyl boronic acid (APBA) and monofunctional diols, and the formation-dissociation process was proved to be repeatable over multiple cycles (Figure 1A). Later, they synthesized another boronic acid-based polymeric aggregations with glucose-sensitivity at pH = 7.4, indicating promising prospects for controlled insulin delivery systems under physiological conditions (Figure 1B) [26]. Kim group prepared two kinds of insulin-loaded polymersomes from polyboroxole-based block copolymers, which could release insulin responding to the glucose-triggered disassembly under physiological conditions (see Figure 2) [28, 29]. Differently, Yao et al. designed a novel nanocarrier with phenylboronate ester as a leaving group in the hydrophobic block, and the nanocarrier successfully realized glucose-responsiveness and insulin release at neutral pH (Figure 3) [27].

In recent years, Zhang group synthesized a series of PBA-based amphiphilic copolymers by radical polymerization and living-controlled polymerization to deliver insulin at physiological conditions [32–37]. By introducing carbohydrate moieties into copolymers, $pK_a$ values of PBA moieties were decreased and glucose sensitivity was enhanced at physiological conditions, which ensured the delivery effectiveness and self-regulated release of insulin. To explore the effect of polymer structures on self-assembled aggregates and glucose-sensitivity, a series of amphiphilic block and random PBA-based copolymers with similar monomer compositions were synthesized by reversible addition–fragmentation chain transfer (RAFT) polymerization [38]. These copolymers could easily self-assemble into spherically shaped nanoparticles in aqueous solution and two types of nanoparticles facilitated insulin delivery at physiological pH, despite insulin release rate of random copolymer was slightly quicker than that of the block ones (Figure 4).

Besides PBA-based amphiphilic copolymers self-assembling into nanoparticles, the chemically cross-linked PBA-decorated nanogels (microgels) also attract much attention because they can meet different needs, such as prompt drug delivery, oral administration, and intravenous injection. Typical PBA-based glucose-responsive nanogels were prepared through modification of poly(N-isopropylacrylamide-co-acrylic acid) [P(NIPAM-co-AA)] with 3-amino-phenylboronic acid (Figure 5) [39–41]. The introduction of PBA groups significantly decreased volume phase transition temperature of microgels, yet a pH value higher than 7.4 was needed to release loaded-insulin in response to glucose.

Wang et al. reported multifunctional microgels based on PNIPAM, N-acryloyl-3-aminophenylboronic acid and two fluorescent molecules [42]. The FRET efficiencies and shrink-swelling properties of microgels could be tuned via thermo-induced collapse or glucose-induced swelling at proper pH and temperature (Figure 6). Tang et al. synthesized a specific glucose-sensitive P(NIPAM-AAPBA) microgel that displayed specific glucose sensitivity at physiological pH [43]. Due to the interference from fructose, galactose, and lactate was negligible, the author considered the contraction-type specific glucose-sensitive microgels had potential applications in self-regulated insulin delivery and glucose sensing (Figure 7). Zhao et al. prepared a novel nanogel with high glucose sensitivity
Figure 2: (A) Self-assembly of PEG-b-PBOx polymersome and its disassembly responding to monosaccharides. Reprinted with permission from [28]. Copyright 2012, American Chemical Society. (B) Schematic illustration for self-assembly and glucose-responsive disassembly of polymersomes. Reprinted with permission from [29]. Copyright 2012, American Chemical Society

Figure 3: Schematic illustration for sugar-responsive behavior of nanoassembly. Reproduced with permission from [27]. Copyright 2011, The Royal Society of Chemistry
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Figure 4: Schematic illustration for the self-assembly and insulin release behaviors of block and random copolymeric nanoparticles. Adapted with permission from [38]. Copyright 2015, American Chemical Society

Figure 5: Synthesis of glucose-sensitive P(NIPAM-PBA) microgels with pendant 3-amino-phenylboronic acid. Adapted with permission from [41]. Copyright 2006, American Chemical Society

Figure 6: Synthesis and modulation of FRET efficiencies for microgels by temperature variations and adding glucose. Reprinted with permission from [42]. Copyright 2011, American Chemical Society

through one-pot thiol-ene copolymerization of pentaerythritol tetra-(3-mercaptopropionate), poly(ethylene glycol) diacrylate, methoxyl poly(ethylene glycol) acrylate and N-acryloyl-3-aminophenylboronic acid [44]. The release of loaded-insulin was triggered and modulated at pH 7.4 responding to different glucose concentrations (Figure 8).

To simultaneously achieve optical detection of glucose, high drug loading capacity, and self-regulated drug delivery, Wu and coauthors reported a series of PBA-based hybrid microgels with glucose-sensitivity to realize self-regulated insulin delivery at physiological pH [16]. The multifunctional hybrid nanogels were made of Ag nanoparticle cores covered with a copolymer gel shell of poly(4-vinylphenylboronic acid-co-2-(dimethylamino)ethylacrylate) [p(VPBA-DMAEA)] [16, 45–47]. The small sized Ag cores (10 ± 3 nm) in microgels provided fluorescence as an optical code. The microgels could not only convert the disruption in homeostasis of glucose level into optical signals to detect the blood glucose levels, but also regulate release of preloaded insulin responding to the glucose level variations (Figure 9A). Later, they reported a fluorescent responsive hybrid nanogel of Zinc oxide@poly[N-isopropylacrylamide (NIPAM)-acrylamide (AAm)-2-aminomethyl-5-fluorophenylboronic acid (FPBA)] [45]. The hybrid nanogels could sensitively and selectively detect glucose via fluorescent signals over the clinically relevant glucose concentrations of 18-540 mg/dl. The nanogels could also regulate loaded-insulin release behaviors in response to different glucose levels and exhibit quicker release rate at higher glucose level (Figure 9B).
Figure 7: The complexation equilibrium between glucose and PBA groups, and the glucose-induced swelling of P(NIPAM-AAPBA) microgels. Reproduced with permission from [43]. Copyright 2014, The Royal Society of Chemistry.

Figure 8: Structures of nanogels and glucose-sensitive behaviors of ARS-loaded nanogels in PBS [44]. Reprinted with permission from [44]. Copyright 2013, Elsevier.
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3 PBA-mediated SA recognition on cancer cells

SA is an anionic monosaccharide that frequently occurs at the termini of glycan chains. Its overexpression on cell membranes implicated in the malignant and metastatic phenotypes of various types of cancers, such as lung, breast, colon, prostate, bladder, and stomach cancer [48, 49]. Thus, advanced techniques to monitor SA expression would have great significance on diagnostics and therapeutics. Interestingly, PBA and its derivatives can recognize SA and form the stable boronic ester with SA not only in the physiological and basic pH but even in acidic medium such as tumor acidic microenvironment [50, 51]. Thus, many researchers have devoted to explore the advanced materials based on PBA-mediated SA-recognition as cancer diagnostic and therapeutic agents [52–58]. Matsumoto et al. prepared a PBA modified monolayer gold electrode to noninvasively detect SA on cell membranes [52]. Deng et al. prepared glucose-bridged and 4-mercaptophenyl boronic acid-decorated silver nanoparticle (Glucose-MPBA@AgNPs) as nanoprobe for highly sensitive recognition of SA expression levels on the surfaces of both normal and cancer cells via surface-enhanced raman scattering spectroscopy (Figure 10) [54].

Among these materials, polymeric nanoparticles occupy the dominant roles owing to their excellent expression in the cancer diagnostic and therapeutic fields [59–63]. Liu et al. prepared PBA and amphiphilic copolymer of maleic anhydride/octadecene coated with semiconductor quantum dots as imaging probes to specifically label SA on living cells [62]. These probes could one-step label and continuously track SA moieties on cell surfaces without any pretreatment of living cells [62]. These probes could one-step label and continuously track SA moieties on cell surfaces without any pretreatment of living cells [62]. These probes could one-step label and continuously track SA moieties on cell surfaces without any pretreatment of living cells [62]. These probes could one-step label and continuously track SA moieties on cell surfaces without any pretreatment of living cells [62]. These probes could one-step label and continuously track SA moieties on cell surfaces without any pretreatment of living cells [62]. These probes could one-step label and continuously track SA moieties on cell surfaces without any pretreatment of living cells [62]. These probes could one-step label and continuously track SA moieties on cell surfaces without any pretreatment of living cells [62]. These probes could one-step label and continuously track SA moieties on cell surfaces without any pretreatment of living cells [62]. These probes could one-step label and continuously track SA moieties on cell surfaces without any pretreatment of living cells [62].

In recent years, polymeric nanoparticles with different architectures and formulations as drug delivery carriers based on PBA-SA recognition for cancer therapeutics were investigated by many researchers [55–57, 64]. De-shayes et al. prepared PBA-installed polymeric micelles to target sialylated epitopes overexpressed on cancer cells, and the micelles showed high affinity for SA even under intratumoral pH conditions, suggesting the potential
application of PBA-installed nanocarriers for enhanced tumor targeting (Figure 13) [55]. Zhao et al. developed PBA-terminated polyethylene glycol monostearate (PBA-PEG-C_{18}) and pluronic P_{123} (PEG_{20}-PPG_{70}-PEG_{20}) targeting drug delivery system based on the selective binding with PBA [64]. Different from others’ work, herein PBA located on micelles margin was protected by fructose to avoid nonspecific interaction with normal cells or other components containing cis-diol residues in vivo circulation (Figure 14).

Besides synthetic polymers, PBA could also be decorated on the natural polymers for specific targeting SA on cancer cells [65–67]. Lee et al. prepared nanoparticles based on (3-aminomethylphenyl)boronic acid (AMPB)-functionalized chondroitin sulfate A (CSA)–deoxyxylulose acid (DOCA) to enhance tumor targeting and penetration [65]. Near-infrared fluorescence imaging revealed that CSA–DOCA–AMPB nanoparticles could target and penetrate into tumor based on both CSA–CD44 receptor and boronic acid–SA interactions. Multiple intravenous injections of DOX-loaded CSA–DOCA–AMPB nanoparticles efficiently inhibited the growth of A549 tumor in xenografted mouse models. These boronic acid-rich nanoparticles are promising candidates for cancer therapy and imaging (Figure 15). Jeong et al. fabricated AMPB-installed hyaluronic acid–ceramide (HACE)-based nanoparticles for tumor-targeted delivery [66]. HACE-AMPB/MB nanoparticles improved cellular accumulation efficiency, tumor penetration efficiency, and antitumor efficacy, indicating the PBA-SA interaction played important roles in augmented tumor targeting and penetration (Figure 16).

PBA-based polymeric nanoparticles could also be used to deliver siRNA for cancer therapy [68, 69]. Ji et al. made PBA conjugated onto low molecular weight polyethylenimine (PEI) to generate amphiphilic PBA-grafted PEI (PEI-PBA) [68]. PEI-PBA and siRNA spontaneously self-assembled into PEI-PBA/siRNA nanocomplexes that dramatically increased siRNA uptake up to 70–90% in several cancer cell lines due to the PBA-SA interaction. Moreover, the PEI-PBA nanovector effectively promoted the lysosome escape of siRNA, thereby robustly induced tumor apoptosis and cell cycle arrest. Hence, SA-
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targeted PEI-PBA could be a highly efficient and safe nanovector to improve the efficacy of cancer siRNA therapy (Figure 17). Kim et al. also reported a PBA-based polymer architecture as gene carrier for targeting anti-angiogenic tumor [69]. The nanocarrier was obtained by crosslinking of low-molecular-weight (MW) PEI to form high-MW PEI based on the interaction between PBA and galactose, resulting in strong interaction with anionic DNA. Inside the tumor cells, the linkages of PBA and sugar were disrupted by either acidic endosomal pH or intracellular ATP, resulting in effective release of preloaded gene (Figure 18). Wang et al. developed PBA-mediated chitosan nanoparticles for tumor targeting, and the results showed that PBA-decorated nanoparticles were more easily internalized by tumor cells compared to non-decorated nanoparticles in 2-D and 3-D cell models, indicating a potential application for delivering more drugs into tumor cells due to the active targeting effect of boronic acid groups [67].

4 Biosensor

Due to the binding capacity of PBA and its derivatives towards free sugars or sugar-protein complexes, they have been extensively investigated as the fundamental parts of biosensor systems. The most typical sensor is glucose-sensor for detecting blood glucose concentrations in diabetes [70–75]. However, mostly of sensors were based on gold nanoparticles/surfaces owing to their plasmonic properties, PBA-based polymeric nanopartil-

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**Figure 15:** Schematic illustration of tumor targeting and penetrating action of CSA-DOCA-AMPB/DOX nanoparticles. Reproduced with permission from [65]. Copyright 2018, WILEY-VCH Verlag GmbH & Co. KGaA, Weinheim

**Figure 16:** The tumor targeting and penetration strategy using HACE-AMPB/MB nanoparticles. Reprinted with permission from [66]. Copyright 2017, Elsevier
Figure 17: Synthesis of PEI-PBA conjugates for SA-targeted siRNA delivery. Reprinted with permission from [68]. Copyright 2016, American Chemical Society

Figure 18: Schematic illustration of anti-angiogenic gene delivery mediated by PBA-PEG-Cross PEI vector. Reprinted with permission from [69]. Copyright 2015, Elsevier
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5 Stimuli-responsive drug delivery systems

PBA and its derivatives can combine with cis-diol/diphenol, and the interactions are pH responsive to disassemble. Some PBA derivatives are hydrogen peroxide ($\text{H}_2\text{O}_2$) responsive to degrade. These properties of PBA and its derivatives could be applied in stimuli-responsive drug delivery systems.

5.1 Diols responsiveness

PBA can bind with a variety of cis-diols, such as galactose, mannose, glucose, 1,3-diols, etc. Their interactions are always pH responsive, which could be applied in the stimuli-responsive drug delivery systems [84–86]. Kim et al. formulated a water-soluble nanoconstruct by forming boronic ester between the cis-1,3-diol of andrographolide (AND) and hydrophilically polymerized phenylboronic acid (pPBA) [85]. Due to the pH- and diol-dependent affinity of boronic ester, AND release was regulated by low pH and high ATP concentrations. Besides, pPBA-AND nanoconstruct exhibited an excellent tumor targeting ability both in vitro and in vivo owing to the intrinsic property of PBA moieties (Figure 21).

5.2 Diphenol responsiveness

The pH-sensitive interaction between PBA and diphenol is always used in the stimuli-responsive drug delivery systems [87–89]. Li et al. developed a DOX-loaded fluorescent nanoparticle based on the installed boronic acid-
modified poly(lactic acid)-poly(ethyleneimine)(PLA-PEI) copolymers for intracellular imaging and pH-responsive drug delivery [89]. DOX-loaded nanoparticles showed pH-responsive drug release and effectively suppressed the proliferation of MCF-7 cells. In addition, the fluorescent nanoparticles tracked the process of endosomal escape by real-time imaging (Figure 22).

5.3 H$_2$O$_2$ responsiveness

H$_2$O$_2$ is highly expressed in tumor site, this could be exploited in the design of H$_2$O$_2$-responsive drug delivery systems for cancer therapy. Interestingly, 4-(methylol) phenylboronic acid and its derivatives are H$_2$O$_2$-responsive to induce PBA disintegration and loaded-drug release [56, 90, 91]. Yu et al. recently designed a temperature and H$_2$O$_2$-responsive “nano-valve” of ROSP@MSN by taking advantage of mesoporous silica nanoparticles (MSN) and 4-(4,4,5,5-tetramethyl-1,3,2-dioxaborolan-2-yl)benzyl acrylate modified polymers (ROSP) [91]. ROSP@MSN could achieve cargo loading drug in cold water, and subsequently close the pore by raising temperature to obtain ROSP@MSN@DOX. Upon the stimulus of H$_2$O$_2$, ROSP@MSN@DOX exhibited excellent DOX release behavior under physiological conditions (Figure 23).

6 Conclusion

In this paper, some new advances of PBA-based strategies for diagnostic and therapeutic applications are reviewed. Because of the versatility for different designs and the specific interactions with polyol compounds, PBA and its derivatives have been widely exploited in constructing glucose-responsive, SA-recognized and stimuli-responsive polymeric nanomaterials, such as drug delivery systems and biosensors. A series of progresses have been made in recent years including synthesis of new PBA-based polymers and fabrication of PBA-decorated nanomaterials. These PBA-decorated materials are designed to operate upon the disease microenvironment characters, such as under physiological pH or tumor acid condition. Until now, most drug delivery systems based on PBA-decorated nanoparticles were focused on insulin and anticancer drugs aiming at treatment of diabetes mellitus and cancer, respectively. And most PBA-based biosensors were focused on blood glucose detection. In conclusion, PBA and its derivatives-decorated polymers and nanomaterials have great potential applications in smart drug delivery systems and biosensors for the diagnosis and therapeutics of diseases.

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