Poly-3-thienylboronic acid: a chemosensitive derivative of polythiophene

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Abstract
Poly-3-thienylboronic acid was synthesized by electrochemical polymerization from 3-thienylboronic acid dissolved in the mixture of boron trifluoride diethyl etherate and acetonitrile. Cyclic voltammetry during electropolymerization shows oxidative and reductive peaks growing in each next cycle. An investigation by scanning electron microscopy displayed the polymer layer like a highly flexible film of 110 nm thick with grains of 60–120 nm in size. Strong negative solvatochromic effect was observed. Optical spectra of poly-3-thienylboronic acid at different potentials and pH were studied. Potential cycling leads to a well reversible electrochromic effect. At pH 7.4, the increase of potential leads to the decrease in the absorption band at 480 nm and to the rise in the absorption band at 810 nm with an isosbestic point at 585 nm. Spectroelectrochemical behavior of poly-3-thienylboronic acid and polythiophene was compared. Binding of sorbitol at fixed electrode potential leads to an increase in the absorbance in the shortwave band and to the decrease in the longwave band; the effect depends on the electrode potential and pH. Perspectives of application of poly-3-thienylboronic acid as new chemosensitive material are discussed.

Keywords Chemosensitive conducting polymers · Polythiophene · Poly-3-thienylboronic acid · Spectroelectrochemistry · Sorbitol sensing

Introduction
Extraordinary properties of conducting polymers (CP) [1–3] resulted in wide applications in different fields of science and industry: electroluminescent and electrochromic devices [4], membranes and ion exchangers [5, 6], materials for energy technologies [7, 8] chemical sensors [9] and biosensors [10], artificial muscles [11], electrocatalysis [12]. A large number of CP such as polyacetylene, poly(3,4-ethylenedioxythiophene) (PEDOT), polyaniline, polypyrrrole, polythiophene (PTh), and their derivatives have acquired a widespread recognition due to their inherent conductivity as well as tunable electrochemical, acid-base, and optical properties. Among these CP, the polymers with PTh backbone belong to the most explored ones, both theoretically and experimentally [13, 14]. This class of CP attracts close attention because of the high chemical and long-time stability, high electrical conductivity, and ease of structural modification. Numerous applications of PTh and its derivatives in chemical sensing were suggested [15, 16]. Based on PTh nitrogen dioxide sensors with electrochemically controllable affinity allowed one to get a fast sensor recovery [17]. PTh was also used as a conductometric indicator for redox-titration in organic media [18], thus providing electrochemical transducing in non-conducting media. However, PTh itself is not chemosensitive, and in all these works, only redox properties of this polymer were used. An application of PTh in selective chemical sensors requires incorporation of a specific receptor. There are different ways of the conjugation of receptor moieties with CP [19], while the strongest coupling between a binding event and physical properties of CP is achieved if the receptor is incorporated directly into or very close to the conducting path of the polymer. Also, a chemosensitive material should possess no additional moieties which can lead to any interference. But in our attempt to apply electrochemically synthesized polyphenylboronic acid for conductometric sensing of fructose (this unpublished work was performed in 2003 together with Q.
Hao), the effects were so small that we were not able to distinguish between an effect of the target analyte and possible pH effect of acidic impurities: despite a strong buffer, it was impossible to exclude pH effect in the range of 0.01–0.001 which would be enough to get the same effect as the analyte. In this sense, a sensor based on PTh has important advantages because of the absence of basic/ acidic groups in its polymer backbone which lead to strong pH dependence of the physical properties of polyaniline and its derivatives. Therefore, preparation of PTh with receptors incorporated close to its conducting path is a perspective way for the development of new chemosensitive materials.

One of the most used artificial receptors is the boronic acid moiety [20, 21]. Boronic acid–functionalized materials overall and boronic acid substituted polymers, in particular, have been widely implemented in chemical sensing of biological analytes containing diol moiety [22, 23], for instance, carbohydrates and α-hydroxy acids. Boronic acid forms reversible covalent complexes with compounds containing a diol group. The reaction occurs in aqueous solutions; the boronic acid is transformed into the corresponding boronate ester. The perspective to use boronic acid as the receptor for diol containing compounds was the motivation to incorporate this group into various conducting polymers. However, only recently the polythiopheneboron derivatives, namely polythiophene substituted by boronic acid, have been prepared. To our knowledge, there is only one publication discussing the physical properties of poly-3-thienylboronic acid (PThBA) [24]; the polymer was synthesized using oxidation by potassium dichromate. The electrochemical synthesis of PThBA was used in [25–27]. Chen et al. [25] applied this polymer to manufacture an electrochromic device [25]. The works [26, 27] were focused on the development of sensors for hydrogen peroxide and glucose correspondingly, but the authors either did not prove the existence of the polymer film [26] or did not described film properties [27]. The receptor properties of the boronic group of PThBA have never been exploited so far.

In this work, we report electrochemical polymerization of thienny-3-boronic acid (Fig. 1) and optimization of this process. Electrochemical synthesis in the mixture of boron trifluoride diethyl etherate (BFEE) and acetonitrile allowed us to get a highly flexible polymer film with a thickness of ~100 nm. We describe the structural, electrochemical, and spectroelectrochemical properties of this polymer and demonstrate its applicability for chemosensing of sugars.

**Experimental**

Prior to electropolymerization, 150-nm-thick gold electrodes on a glass wafer were cleaned by acetone (Chemsolute), ethanol (Merck), water, and piranha solution (a 1:3 (v/v) mixture of 30% H₂O₂/concentrated H₂SO₄). **Caution: this mixture reacts violently with most organic materials and must be handled with extreme care.** Then, the electrodes were rinsed thoroughly with deionized water additionally purified by the EGLA-classic system. Indium tin oxide (ITO)-coated glass slides (Sigma-Aldrich) were cleaned with ethanol and rinsed with deionized water.

Synthesis of PThBA and PTh films was performed in 50-mM solution of the corresponding monomer (Sigma-Aldrich) either in pure acetonitrile (Honeywell) containing 100 mM of tetrabutylammonium hexafluorophosphate (NBu₄PF₆) (Sigma-Aldrich) or in the mixture of 90% BFEE (Sigma-Aldrich) and 10% acetonitrile (v/v) containing 50 mM 2,6-di-tert-butylpyridine (DTBP) (Sigma-Aldrich) by potential cycling from −0.2 V to +1.8 V vs. Ag/AgCl electrode (Metrohm) filled by 2 M LiCl (Merck) in ethanol (Merck) with the same solution in the salt bridge. The proton scavenger DTBP was added to avoid a hydrogenating of BFEE during polymerization in consequence of its high sensitivity to moisture leading to side reactions. An application of the BFEE/ acetonitrile mixture for electropolymerization was optimized and exploited earlier for electropolymerization of PTh [28].

Cyclic voltammetry was performed using General Purpose Electrochemical System Autolab PGSTAT-12 with Nova 2.1.4 software. A three-electrode configuration was used. Platinum wire (ChemPur) of 0.5 mm in diameter served as the counter electrode in spectroelectrochemical measurements, while a planar gold electrode with an area of ~1.5 cm² was used for the same purpose in cyclic voltammetry. Measurements in buffer solution (pH 7.4) consisting of 10 mM sodium phosphate (Merck) and 140 mM sodium chloride (Roth) were performed using Ag/AgCl reference electrode without a salt bridge. Polymer films deposited on ITO-coated glass slides were rinsed with acetonitrile, dried nitrogen flow, and then applied for spectroelectrochemical measurements.

Scanning electron microscopy (SEM) was performed at Zeiss Microscope Evo® MA 15. Spectral and spectroelectrochemical measurements were done using spectrophotometer Evolution 200 Series from Thermo Scientific. Nikon Eclipse LV 100 microscope equipped with CCD-camera was applied for optical microscopy. The experiments were performed at room temperature. Other experimental details are described in [28].

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**Fig. 1 Suggested structure of poly-3-thienylboronic acid**
Results and discussion

At the beginning of the work, we tried to make electrochemical deposition of PThBA from the solution of acetonitrile with 100 mM NBu4PF6, but the film adhesion to the electrode surface was poor, and the film growth was inhibited by relatively low conductivity of the formed material. The obtained cyclic voltammograms were similar to those reported in [27] (except the current polarity which was inverted in that work). Much better results were obtained for electrochemical polymerization of PThBA from BFEE/acetonitrile solution [29, 30]. The cyclic voltammetric curves during electropolymerization display an increase in the cathodic and anodic peaks in each next cycle (Fig. 2); such behavior is typical for the deposition of highly conducting materials. Both oxidation and reduction peaks were shifted symmetrically to anodic and cathodic directions correspondingly.

The morphology of films obtained from acetonitrile and BFEE/acetonitrile solution was also very different. The polymer films synthesized in acetonitrile were very heterogeneous and looked like a broken solid layer. No lateral polymer growth between gold strips on the electrode was observed (Fig. 3a). In BFEE/acetonitrile solution, the polymer growth was very fast, and a well visible lateral growth filling the gap between the strip electrodes was observed (Fig. 3b). A similar difference was observed earlier for the electropolymerization of PTh [28]. Therefore, the second electrolyte was used for electrochemical polymerization in further experiments. Being deposited at such conditions onto ITO glass slides, the polymer formed a smooth scarlet film (SI, movie 1) with a characteristic metallic shine. Already during electropolymerization, a strong electrochromic effect was observed—the color of the film was changed between scarlet and blue (SI, movie 1). The polymer film on the electrodes was stable in acetonitrile, but in aqueous solutions, the adhesion was much weaker. This can be related to the hydration of the boronic acid moiety.

Further structural investigation of PThBA films polymerized in the second electrolyte was performed using scanning electron microscopy (Fig. 4). The thickness of the film formed after 20 cycles of cyclic voltammetry was determined by SEM measurements as 110 nm. Very close value gives a calculation based on the integration of cyclic voltammetry measured in organic electrolyte after electropolimerization (Fig. 5) and using Faraday low in the assumption that the polymer density is the same as for PTh (1.06 g/mL [31]). It means that almost the whole amount of the synthesized polymer is adsorbed on the electrode surface. Figure 4a shows the polymer film detached from the ITO. One can see high flexibility and elasticity of this film: its curvature in the field of wrinkles is comparable with the film thickness. At higher magnification, one can observe a formation of some thicker round spots, probably as a result of secondary nucleation; similar structures on the smooth polymer layer were observed for electropolymerization of N-methylaniline [32]. The film surface at higher magnification looks to be formed by densely packed grains of the size ~ 60–140 nm.

The electrochemical activity of PThBA in the aqueous electrolyte is very low; no clear oxidation or reduction peaks were observed in cyclic voltammograms. The same effect was described earlier for PTh [28]. In the case of PTh, the low electrochemical activity in aqueous electrolytes was explained by low penetration of hydrophilic ions from aqueous solution into the highly hydrophobic film of polythiophene [28]. This can be formally described as an increase of the oxidation potential of polythiophene in an aqueous environment but the potential value required for the polymer oxidation cannot be reached because of limited polymer stability. Due to the presence of boronic acid moieties, PThBA should be less hydrophobic than PTh, but probably this hydrophilicity still is not enough to provide a good excess for inorganic ions from aqueous solution.

The electrochemical activity of PThBA in acetonitrile is well measurable and similar to that of PTh (Fig. 5). The values of the redox potential obtained as the middle point between the peaks are 0.78 V for PTh and 0.82 V for PThBA. The values of the peak current are very similar. For both polymers, there is complete reversibility of oxidation and reduction. The gap between oxidative and reduction peaks is some higher for PThBA than for PTh; this may indicate additional hindrances for the penetration of hydrophobic ions from the electrolyte into the polymer or for the electron transfer between polymer molecules.

The polymer films of ~ 100-nm thickness on ITO were used for optical spectroscopy and spectroelectrochemical measurements. The spectra of PThBA in acetonitrile show a well-pronounced absorption band at 420–570 nm and a wide weak adsorption band around ~ 800 nm (Fig. 6a) being similar
to the spectra of PTh. Assuming that the density of PThBA is close to that of PTh, we get the value of extinction recalculated to the molar concentration of monomers $5.9 \times 10^3 \text{ L cm}^{-1} \text{ mol}^{-1}$. In aqueous solutions, the spectra have a similar shape but with an essential shift in the wavelengths of the absorption peaks (SI Fig. S1). Incorporation of boronic acid moiety into PTh leads to the small hypsochromic shift in the main absorption band which is increasing from $\sim 5$ nm until $\sim 20$ nm with polymer oxidation. The spectra are qualitatively similar to those reported in [25]. Both PTh and PThBA show a strong negative solvatochromic effect, but for PTh, this effect is much larger than for PThBA. The difference in the absorption maxima in aqueous solution and in acetonitrile for PTh in the reduced state is $\sim 260$ nm, while for PThBA this effect is only $\sim 110$ nm. This may be explained by much lower energy of the ground state of PThBA in aqueous solvent due to the strong hydrophilic contribution of the boronic acid moiety (Fig. 6b).

Spectroelectrochemistry of PTh and PThBA also displays an essential difference between these polymers. Practically no influence of the electrode potential on the spectra of PTh was observed in aqueous solutions (Fig. 6b), this corresponds to the data [32]. In a contrast, an increase of the electrode potential leads to remarkable changes in the optical spectra of PThBA (Fig. 6b): the shortwave peak is decreasing while the longwave peak is increasing with an isosbestic point at $\sim 585$ nm.
In acetonitrile, the spectroelectrochemistry of PTh and PThBA is more similar (Fig. 6a, c, d). In this case, for both polymers, an increase of the electrode potential leads to a decrease in the main adsorption band and to the increase in the adsorption in the longwave adsorption band (600–900 nm) with a less pronounced isosbestic point at 570–590 nm. Both in aqueous solution and in acetonitrile the change in the transmission at 450–500 nm was 28–30% for the potential change between \(-0.4\) V and 1.0 V, about the same value was obtained in the electrochromic device for the potential change for 3 V in acetonitrile based gel electrolyte [25].

The main feature of PThBA is the presence of a receptor group for the binding of diol moieties. Here, we demonstrate optical changes due to the binding of sorbitol (Fig. 7). With the goal to get the maximal effect of this analyte on spectral properties, the sorbitol concentration of 500 mM was used. The binding constant of sorbitol to 3-thienylboronic acid at pH 7.4 is \(\sim 300\) mol\(^{-1}\) (SI Fig. S3). Assuming the same value for the polymerized form of this compound (PThBA), we get that at 500 mM sorbitol concentration over 98% of the receptor moiety will be occupied. A formation of the PThBA-sorbitol conjugate leads to an increase in optical absorbance in the range 350–550 nm and to the decrease in the absorbance at longer wavelengths. The effect depends on the electrode potential (Fig. 7, SI Fig. S2). For the absorbance value measured near maximum (455 nm), the maximal value of the relative absorbance change at pH 7.4 is reached at the electrode potential of approximately +0.8 V (Fig. 7b). It is known that the binding constant of the boronic acid moiety with sugars is increasing at alkaline pH [33]. We have observed that pH increase from 7.4 to 8.0 leads to about twice a higher relative absorbance change of PThBA due to sorbitol binding (Fig. 7b). Also, in this case, the maximal effect at pH 8.0 was observed for the oxidized state of the polymer. The magnitude of the observed effect is about the same as for polyphenylboronic acid [34] but the polymer backbone without basic/acidic groups promises a better selectivity.

Finally, the properties of PThBA are close to PTh but the presence of a chemosensitive group in PThBA makes its perspective for applications in chemical sensors. Conjugation of the receptor group with redox-active polymer allows us to consider this new material to be promising for electrochemical chemotransistors [17].

**Fig. 5** Cyclic voltammetry of films of poly-3-thienylboronic acid (red) and polythiophene (blue) in acetonitrile containing 100 mM NBu\(_4\)PF\(_6\). Electrode: ITO, macroscopic electrode area 1.3 cm\(^2\). Scan rate: 100 mV/s

**Fig. 6** Optical spectra of polythiophene (solid line) and poly-3-thienylboronic acid (dotted line) at potentials –0.4, –0.2, 0, +0.2, +0.4, +0.6, +0.8, and +1.0 V (potential increase is indicated by arrows) in acetonitrile containing 0.1 M NBu\(_4\)PF\(_6\) and in aqueous buffer 10 mM sodium phosphate, 140 mM NaCl, pH 7.4 (b) and potential influence on intensity of both adsorption bands for polythiophene (c) and poly-3-thienylboronic (d) for the spectra in (a). The potentials are indicated relative quasi-reference silver electrode (a, c, d) or Ag/AgCl (140 mM NaCl) electrode (b)
35] or other sensors with electrochemically controlled affinity [36, 37]. The stability of the polymer films in an aqueous environment can be improved by using corresponding anchor layers, for example, by deposition of cationic polymers.

Conclusion

We have described preparation of a new chemosensitive material: thin and highly flexible films of poly-3-thienylboronic acid. The material can be prepared by electrochemical synthesis in situ; therefore, it is well compatible with methods of surface design and preparation of sensor arrays or materials libraries based on electrically addressed polymerization [38]. Electrochemical and spectral properties of this material are close to PTh, but the incorporated boronic acid moiety provides its chemical sensitivity to diol containing compounds. The ability to form stable films allows one to consider applications in support-free conditions, for example, for the formation of chemosensitive electromechanical actuators. Development of such systems and an analysis of its electrical conductivity and a preparation of conductometric chemosensors and electrochemical chernotransistors are the topics of our future work on this new material.

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Fig. 7 Influence of sorbitol on optical spectra of poly-3-thienylboronic acid: the spectra in the presence (green) and in the absence (blue) of 500 mM sorbitol at electrode potential + 0.8 V measured in aqueous buffer 10 mM sodium phosphate, 140 mM NaCl, pH 7.4 (a) and influence of the electrode potential and pH on the relative absorbance at 455 nm (b). The electrode potential is indicated vs. Ag/AgCl (140 mM NaCl), pH 7.4 (squares) or 8.0 (circles)
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