Silylated Initiators for the Efficient Preparation of Borane-End-Functionalized Polymers via ATRP

Yang Qin, Chengzhong Cui, and Frieder Jäkle*

Department of Chemistry, Rutgers University Newark, 73 Warren Street, Newark, New Jersey 07102 Received July 26, 2006; Revised Manuscript Received December 13, 2006

ABSTRACT: Borane-end-functionalized polymers have been prepared through (i) atom transfer radical polymerization (ATRP) of styrene using a silylated initiator, 1-bromo-1-(4-trimethylsilylphenyl)ethane (PEB-SiMe₃), (ii) exchange of the SiMe₃ end groups with BBr₃, and (iii) modification of the Lewis acidic boron centers with nucleophiles. Ditelechelic α, ω -borane-functionalized polymers were obtained through atom transfer radical coupling (ATRC) of the silane-end-functionalized polymers followed by exchange with BBr₃ and subsequent reaction with nucleophiles. Self-assembly of the borane-functionalized mono- and ditelechelic polymers into supramolecular structures by means of reversible noncovalent and covalent interactions including Lewis acid—base complexation and boroxine formation has been explored.

Introduction

Boron-containing polymers continue to attract much attention due to their potential use in catalysis, sensor applications, and material science.¹ Research in these areas has been mainly focused on the incorporation of boron moieties into the main chain and the attachment as side groups of polymeric structures. The preparation of polymers containing borane moieties as end groups, on the other hand, is intriguing since boron-end-functionalized polymers may serve as precursors to various other telechelic polymers using well-established functional group conversion techniques and can also be advantageously used as building blocks for the formation of block or graft copolymer architectures.²

The methods that are most commonly applied for the preparation of boron-end-functionalized polymers involve (i) the use of borane initiators, (ii) the addition of borane chain transfer agents in olefin polymerization, and (iii) the functional group conversion of suitable preformed telechelic polymers with a borane source. (i) Living cationic polymerizations are frequently initiated with boron halides and thus represent a facile method for the incorporation of boron moieties at the termini of polymer chains.^{3–5} For instance, Faust and co-workers have studied the living cationic polymerization of isobutylene and α-methylstyrene initiated with BCl₃. The resulting telechelic polymers bear a Cl atom at one chain end and a BCl2 group at the other, which then can be further transformed to other borane functionalities such as B(OMe)2 and B(OH)2. Another methodology that takes advantage of boron initiators is the polyhomologation of organoboranes, which has been pioneered by Shea and co-workers.⁶⁻⁸ The living polymerization of an ylide such as dimethylsulfoxonium methylide is catalyzed/initiated by trialkylboranes, resulting in well-defined organoboron star polymers that subsequently are converted into linear polymethylenes through oxidative B-C bond cleavage. (ii) In a different approach, the incorporation of borane moieties at the polymer chain ends by using borane chain transfer agents in metallocene-mediated olefin polymerizations has been demonstrated by Chung and co-workers. 9,10 With 9-BBN and HB(Mes)₂ (9-BBN = 9-borabicyclononane; Mes = 2,4,6-trimethylphenyl)as chain transfer agents in the polymerization of ethylene, the

respective borane-terminated polyethylenes were obtained, while the bulkier borane $HB(Trip)_2$ (Trip = 2,4,6-triisopropylphenyl) did not get incorporated due to increased steric hindrance of the Trip groups. The terminal borane groups were further transformed to hydroxyl groups by oxidative cleavage with H₂O₂ under basic conditions. Interestingly, the 9-BBN borane end groups can also react with limited amount of oxygen to form peroxide (B-O-O-C) species, and the resulting polymers were used in turn as macroinitiators for the free radical chain extension with other monomers such as methyl methacrylate.9 (iii) Finally, functional group conversion by hydroboration of vinyl-terminated polymers also gives straightforward access to borane-end-functionalized polymers. For instance, Katayama and co-workers showed that hydroboration of vinvl-terminated poly(norbornene)s with 9-BBN led to borane-terminated poly(norbornene)s which were later transformed to hydroxylterminated polymers. 11 Similarly, hydroboration of vinylterminated polypropylene was used by Chung and co-workers for the synthesis of maleic anhydride-modified polypropylene. 12 Moreover, Chuio and co-workers studied the hydroboration polymerization of α,ω -vinyl-terminated polymers with bifunctional boranes for the synthesis of organoboron segmented block polymers. 13 In recent work, Ohno and co-workers introduced 9-BBN moieties at both ends of poly(propylene oxide) (PPO) through either hydroboration or dehydrocoupling reactions.¹⁴ The borane end groups served as anion traps, and high ion conductivity and transference were observed for the boraneterminated PPO.

Herein we report a new synthetic route to borane-end-functionalized polymers, in which the use of functional initiators in a controlled polymerization protocol is combined with facile and high yielding polymer modification procedures. We have previously described a straightforward methodology for the incorporation of boron moieties into the side chains of well-defined polystyrenes through highly selective exchange of trimethylsilyl substituents with boron halides. ^{15,16} We have also demonstrated that the halide substituents on boron can easily be exchanged, and the properties of these polymers can thus readily be fine-tuned. We show here that controlled free radical polymerization of olefins with silyl-functionalized initiators can serve as an excellent tool for the preparation of well-defined polymers, in which either one or both chain ends are function-

^{*} Corresponding author. E-mail: fjaekle@rutgers.edu.

Scheme 1. Synthesis of Dibromoboryl-End-Functionalized Polymers by Atom Transfer Radical Polymerization and Subsequent Silicon—Boron Exchange

alized with boranes. We further describe how these borane moieties reversibly bind to nucleophiles and can be advantageously used in supramolecular polymer chemistry. In this context, the self-assembly of polymers through donor—acceptor interaction and reversible boroxine ring formation^{17–19} is discussed.

Results and Discussion

Synthesis of Silane-End-Functionalized Polymers. Silaneterminated polystyrene (PS-SiMe₃) was obtained by atom transfer radical polymerization (ATRP)20,21 using the silanefunctionalized initiator 1-bromo-1-(4-trimethylsilylphenyl)ethane (PEB-SiMe₃, Scheme 1),²² which is an analogue of the commonly used ATRP initiator 1-bromo-1-phenylethane (PEB).²⁰ A typical ATRP protocol was used with CuBr/N, N', N', N'', N''pentamethyldiethylenetriamine (PMDETA) as the catalyst system in anisole (50% w/w) at 110 °C and ratios of St:PEB-SiMe₃: CuBr:PMDETA of 50:1:1:1 and 40:1:1:1, respectively. The polymerization was stopped at about 40-50% conversion to ensure preservation of the bromine end groups. According to gel permeation chromatography (GPC) and GPC with in-line multiangle laser light scattering detector (GPC-MALLS) analysis, the polymers show narrow molecular weight distributions (PDI < 1.2), indicative of a controlled polymerization process. The polymers were used in the preparation of monotelechelic organoboron polymers as described below.

The bromine end groups of the silane-functionalized polymer PS-SiMe₃ can be further manipulated through atom transfer radical coupling (ATRC)²³ to form the ditelechelic α,ω -silanefunctionalized polymer, Me₃Si-PS-SiMe₃ (Scheme 1). The sample of PS-SiMe₃ ($M_n = 3358$, GPC-RI) used for ATRC was prepared in bulk, and the polymerization was stopped at an early stage (~30% conversion) to ensure full coverage of the bromine end groups. As previously described by Matyjaszewski and coworkers for the coupling of bromine-terminated PS,²³ CuBr/ PMDETA and activated Cu powder²⁴ were used as the catalyst system. The Cu powder was added in order to reduce the amount of CuBr₂, which acts as a deactivator during the coupling process. A slightly modified procedure with a ratio of PS-SiMe₃: CuBr:PMDETA:Cu of 1:2:10:5 at 90 °C was applied in order to maximize the coupling efficiency. The coupling reaction was monitored by GPC analysis, as shown in Figure 1.

According to the GPC results, the coupling reaction reaches about 90% after 30 min and 94% after 90 min. The extent of coupling was calculated from the equation $\chi_c = 2(1 - M_{n,0}/M_n)$, where χ_c is the extent of coupling, $M_{n,0}$ is the initial molecular weight, and M_n is the molecular weight after

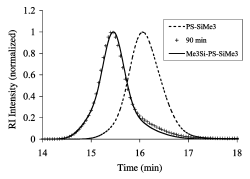


Figure 1. Gel permeation chromatography (GPC) traces for the atom transfer radical coupling (ATRC) process of PS-SiMe₃ (THF, 1 mL/min, refractive index detector; ---, PS-SiMe₃ precursor; +, crude product after 90 min reaction time; —, isolated product after precipitation).

coupling reaction.²³ Further reaction was not observed with extended reaction time based on GPC analyses. As shown in Figure 1, only a small low molecular weight tail can be seen in the trace after 90 min, indicating an efficient coupling reaction. The coupling product Me₃Si-PS-SiMe₃ was purified by precipitation from THF into methanol/hexanes (50/50). Polymer chains with lower molecular weight did not as effectively precipitate under these conditions and were decanted with the supernatant. The low molecular weight fraction significantly reduces in the trace of Me₃Si-PS-SiMe₃ after precipitation and the extent of coupling was calculated to be 96%. Accordingly, the multiplet resonances at ca. 4.4 and 4.5 ppm for the proton of the -CPh(H)Br end group of PS-SiMe3 are not observed in the ¹H NMR spectrum of the coupled product Me₃Si-PS-SiMe₃ (see Figure S1 in the Supporting Information). Moreover, a significantly higher glass transition temperature was observed for Me₃Si-PS-SiMe₃ (T_g (onset) = 95 °C) relative to PS-SiMe₃ $(T_{\rm g}({\rm onset})=80~{\rm ^{\circ}C})$, consistent with the increased molecular weight of Me₃Si-PS-SiMe₃ ($M_n = 6439$ for Me₃Si-PS-SiMe₃ vs $M_n = 3358$ for PS-SiMe₃).

Functionalization of PS-SiMe₃ and Me₃Si-PS-SiMe₃. The trimethylsilyl groups in PS-SiMe₃ and Me₃Si-PS-SiMe₃ were exchanged for BBr2 groups through reaction with a slight excess of BBr₃ in CH₂Cl₂ over a period of 12-24 h (Scheme 1). The dibromoborylated polymer, PS-BBr2, was isolated by precipitation into cold hexanes followed by freeze-drying from benzene. The ¹¹B NMR spectrum of the product shows a broad peak at 56 ppm, which is in the range of typical dibromoarylboranes. In the ¹H NMR spectrum, the resonance at 0.3 ppm due to the trimethylsilyl groups completely disappeared, and a new resonance in the aromatic region at 8.04 ppm evolved, which corresponds to the protons at the ortho positions of the phenyl end group (see Figure 5).²⁵ The multiplet structure of this signal reflects the atactic nature of the PS chain. The molecular weight derived from ¹H NMR end-group analysis matched very well with that obtained from GPC analysis for the starting material, indicating that the boron-silicon exchange reaction proceeded smoothly. Ditelechelic α, ω -borane-functionalized polymers are obtained in a similar procedure by reaction of Me₃Si-PS-SiMe₃ with BBr₃ (Scheme 1). The NMR spectra of Br₂B-PS-BBr₂ are identical to those of PS-BBr2, except for the lack of signals for the Br terminus observed for monotelechelic PS-BBr₂ (see Figure S2 in the Supporting Information).

The polymers PS-BBr₂ and Br₂B-PS-BBr₂ are highly reactive toward air and moisture and therefore most conveniently used in situ for further reaction with nucleophiles. Scheme 2 describes the functionalization of PS-BBr₂ with different nucleophiles. The functionalization of Br₂B-PS-BBr₂ follows the same methodology, and the resulting ditelechelic borane polymers

Scheme 2. Functionalization of PS-BBr2 and Br2B-PS-BBr2

show similar features in the NMR spectra as their monofunctionalized counterparts.

As shown in Scheme 2, the polymeric Lewis acid PS-BTh₂ was obtained by treatment of PS-BBr₂ with 2-(trimethylstannyl)thiophene under inert atomsphere. The product was isolated in high yield (81%) by precipitation into hexanes and dried under high vacuum. The ¹¹B NMR spectrum of PS-BTh₂ shows a broad peak at 50 ppm, which is in the range of typical triarylboranes and similar to that of the bis(thienyl)boryl homopolymer ($\delta = 47$) and the related molecular species (4-t-BuPh)BTh₂ ($\delta = 55$). ^{15b} According to GPC analysis of PS-BTh₂ in THF, the molecular weight ($M_n = 3517$) and polydispersity (PDI = 1.09) are similar to those of the starting material PS-SiMe₃ ($M_n = 3688$, PDI = 1.10), indicating that no crosslinking or degradation of the polymer occurred during the boron-silicon and boron-tin exchange reactions. The polymer slowly degrades under ambient conditions and was therefore handled under an inert atmosphere. A hydrolytically stable boronic acid-end-functionalized polymer, PS-B(OH)₂, was obtained simply by hydrolysis of PS-BBr₂ (Scheme 2). The formation of a boroxine linkage and self-assembly of the polymers through the boronic acid end groups will be discussed in the next section.

Self-Assembly of Polymers through the Borane End **Groups.** Supramolecular chemistry has become a basic concept and an expanding area of today's chemical research since Lehn, Pederson, and Cram received the Nobel prize in 1987.²⁶⁻²⁸ Supramolecular polymers, assembled from traditional polymers through noncovalent interactions including complementary hydrogen bonding and metal-ligand complexation, are especially intriguing for the combination of novel structures and features of supramolecular species and the well-known polymeric properties.^{29,30} These noncovalent interactions allow for establishment of a stimuli-responsive equilibrium between the supramolecular structures and the polymeric building blocks. We studied the possibility of using the borane-end-functionalized and ditelechelic polymers as building blocks toward supramolecular polymer structures through (i) the reversible formation of Lewis acid-base complexes³¹ and (ii) the reversible construction of boroxine rings.

(i) Assembly through Lewis Acid-Base Complexes. We have shown previously that for bis(thienyl)borane side-group functionalized polystyrene the boron centers are highly Lewis acidic, and it is therefore possible to form isolable complexes with pyridine bases.³² Indeed, the pyridine complex of PS-BTh₂,

Scheme 3. Formation of Polymeric Lewis Acid-Base Complexes

PS-BTh₂•Py, was readily obtained by stirring PS-BTh₂ with excess pyridine in CH2Cl2 followed by removal of volatile materials under vacuum (Scheme 3). A sharp signal at 0 ppm in the ¹¹B NMR spectrum confirmed the tetracoordinated nature of the boron centers. Moreover, integration of the ¹H NMR spectrum is in agreement with one pyridine molecule bound to each polymer chain.33

Treatment of the ditelechelic polymer Th₂B-PS-BTh₂ with 1 equiv of 4,4'-dipyridyl (bipy) provides an opportunity for the self-assembly of longer polymer chains [Th₂B-PS-BTh₂•bipy]_n through donor-acceptor interactions (Scheme 3).33 Under relatively concentrated conditions (15.5 mM boryl groups and 7.7 mM 4,4'-dipyridyl in CDCl₃), the ¹¹B NMR spectrum shows a single peak at 0 ppm consistent with pyridine-coordinated boron centers. The ¹H NMR spectra of PS-BTh₂•Py and [Th₂B- $PS-BTh_2 \cdot bipy|_n$ show very similar patterns and further indicate complexation of the Lewis acidic boron centers (Figure 3). Integration confirms the presence of one 4.4'-dipyridyl molecule per polystyrene chain. DSC studies revealed a glass transition temperature of $T_g = 105$ °C for $[Th_2B-PS-BTh_2 \cdot bipy]_n$, which is similar to that of high-MW PS and significantly higher than that of the low-MW starting material $Th_2B-PS-BTh_2$ ($T_g =$ 88 °C) (Figure 2), thus confirming assembly of Th₂B-PS-BTh₂ through 4,4'-dipyridyl into a polymer with considerably higher molecular weight.

To further investigate the formation of the extended structure in solution, we monitored the titration of Th2B-PS-BTh2 with 4,4'-dipyridyl in anhydrous CDCl₃ using ¹H NMR spectroscopy. The original concentration of Th₂B-PS-BTh₂ was ca. 15.5 mM boryl groups, and upon addition of 4,4'-dipyridyl only one set of averaged signals was observed. The chemical shifts

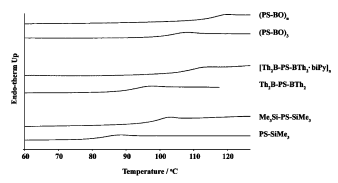


Figure 2. Differential scanning calorimetry (DSC) plots of the endfunctionalized polymers and their self-assembled aggregates (onset, second heating curve, 20 °C/min).

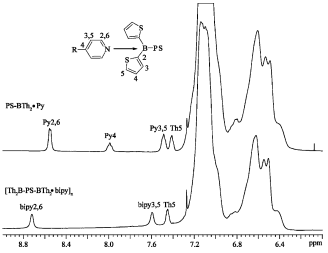


Figure 3. ¹H NMR spectra of PS-BTh-Py (R=H) and [Th₂B-PS-BTh₂·bipy]_n (R = pyridyl) in CDCl₃ (ca. 15.5 mM functional group).

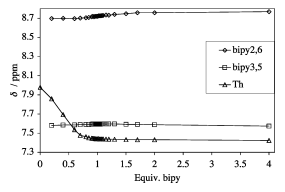


Figure 4. ¹H NMR titration of Th₂B-PS-BTh₂ with 4,4′-dipyridyl in CDCl₃ (ca. 15.5 mM boryl group).

changed gradually, as shown in Figure 4. The signal for the thiophene protons Th-H5 shifted from the initial value of 7.98 to 7.44 ppm upon addition of 1 equiv of bipy per polymer chain and leveled off at higher bipy concentration, indicating that the boryl groups were essentially fully coordinated in the presence of 1 equiv of bipy. The signal for bipy-H3,5 did not experience a significant change, and the bipy-H2,6 resonance remained at ca. 8.70 ppm up to equimolar bipy to polymer ratio. However, the chemical shift gradually increased to 8.75 ppm when up to 2 equiv of bipy was added. Averaged ¹H NMR signals for dicoordinated and monocoordinated bipy groups are apparently observed in this regime. The data presented thus further support the formation of [ThB-PS-BTh•bipy]_n in solution at a ratio of bipy to organoboron polymer of 1:1.

(ii) Assembly through Boroxine Linkages. Boronic acids³⁴ and polymer-supported boronic acids have found applications as scaffolds for the solid-phase synthesis of carbohydrates,³⁵ sensors and stimuli responsive polymers including the sensing of sugars,^{36–38} and separation media in biochemical applications.^{39,40} Boroxines, the cyclic anhydrides of boronic acids formed by elimination of three molecules of water, have found commercial use as flame retardants,⁴¹ dopants that enhance lithium ion transference in polymer electrolyte,⁴² and boronic acid alternative in Suzuki–Miyaura coupling reactions.⁴³

PS-BBr₂ was treated with water to give PS-B(OH)₂, which was then subjected to azeotropic distillation in benzene. Upon elimination of water, (PS-BO)₃, a triarm star with boroxine core is expected to form. The ¹H NMR signals of the two protons at the ortho positions of the phenyl ring attached to boron are used to monitor the process. As shown in Figure 5, the signal of PS-B(OH)₂ at 7.7 ppm disappeared upon azeotropic distillation and a new peak formed at 8.1 ppm. These chemical shifts are very similar to those reported in the literature for phenylboroxine and phenylboronic acid derivatives, 18 respectively, suggesting the transformation into a boroxine-centered star polymer. Moreover, DSC analysis of the product (PS-BO)₃ showed a significantly higher glass transition temperature ($T_{\rm g} = 101$ °C) over that of the starting material PS-SiMe₃ ($T_g = 80$ °C) (Figure 2), which indicates a significant increase of molecular weight and thus provides further evidence for the assembly of PS-B(OH)₂ through boroxine linkages. Importantly, the boroxine can readily be transformed back to boronic acid functionalities by hydrolysis in the presence of a small amount of D₂O as shown by ¹H NMR spectroscopy (Figure 5). Thus, the formation of the star polymer is a reversible process.

When the ditelechelic polymer Br₂B-PS-BBr₂ was similarly subjected to hydrolysis, the bis(boronic acid) polymer (HO)₂B-PS-B(OH)₂ was obtained (Scheme 2). Upon azeotropic distillation of a benzene solution of (HO)₂B-PS-B(OH)₂ a white precipitate formed, which was isolated by removal of benzene under high vacuum. The resulting white solid was found to be insoluble in anhydrous acetone.⁴⁴ Presumably, instead of a triarm star-like structure as in the case of (PS-BO)3, a network-like structure with boroxine cross-links was formed, as illustrated in Figure 6.45 We performed DSC and TGA measurements to further confirm the formation of the boroxine network and to investigate its properties. Indeed, a significantly higher glass transition temperature ($T_{\rm g} = 112$ °C) was observed compared to the starting material Me₃Si-PS-SiMe₃ ($T_g = 95$ °C), consistent with a more rigid structural framework (Figure 2). In the TGA measurements, both Me₃Si-PS-SiMe₃ and the boroxine network showed a similar decomposition onset of ca. 400 °C. No residue was found for Me₃Si-PS-SiMe₃, and only 3% weight retained in the case of (PS-BO)_n, presumably due to residual boron oxide.

When a drop of D_2O was added to a suspension of $(PS-BO)_n$ in dry acetone- d_6 , a clear solution was obtained. The 1H NMR spectrum was similar to that of $PS-B(OH)_2$ in acetone- d_6/D_2O (Figure 5), indicating the breakup of the boroxine network and re-formation of the boronic acid-functionalized polymer, $(HO)_2B-PS-B(OH)_2$. Thus, the formation of boroxine cross-links is reversible, and the polymer properties can be modified by the presence or absence of water.

Conclusions

We have developed a modular synthetic methodology for the preparation of borane-end-functionalized polymers through ATRP of styrene using a silylated initiator, followed by exchange with BBr₃ and modification through treatment with nucleophiles. Our methodology is highly versatile since various

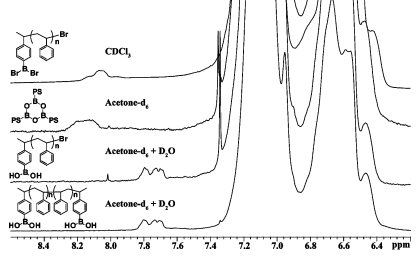


Figure 5. Reversible interconversion of PS-B(OH)₂/(PS-BO)₃ and (HO)₂B-PS-B(OH)₂/(PS-BO)_n (insoluble in acetone) as monitored by ¹H NMR spectroscopy.

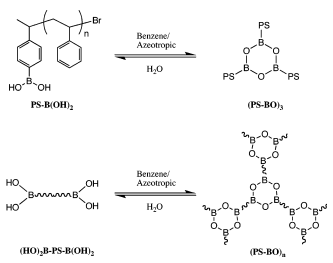


Figure 6. Schematic representation of the formation of the triarm star structure of (PS-BO)3 with a boroxine core and the boroxine crosslinked polystyrene (PS-BO) $_n$ (the structural representation is idealized as a few non-cross-linked -PS-B(OH)₂ and -PS-B(OH)-PS- moieties likely also exist).

substituents can be introduced to the borane end groups, which can serve as precursors to other end-functionalities and be used for the construction of polymer assemblies. In pursuit of these goals, we have shown that the bis(thienyl)boryl- and boronic acid-end-functionalized polymers can be reversibly converted to supramolecular polymer structures through Lewis acid-base interactions and boroxine linkages, respectively. Thus, new methodologies for the reversible assembly of supramolecular polymer structures in addition to traditional hydrogen bonding and metal-ligand interactions are provided. Especially the reversible assembly of polymers through strong covalent B-O bonds in an equilibrium that is easily shifted by addition or removal of water, and can be influenced by the presence of Lewis bases,⁴⁷ is expected to provide new opportunities in the field.

Experimental Section

Materials and General Methods. The compounds 1-bromo-4ethylbenzene, Mg (turnings), Me₃SiCl, N-bromosuccinimide, BBr₃, CuBr, CuBr₂, Cu (electrical dust), pyridine, 4,4'-dipyridyl (bipy), and anisole were purchased from Acros. N,N',N',N'',N''-Pentamethyldiethylenetriamine (PMDETA) was purchased from Aldrich. BBr₃ was distilled under vacuum, and Me₃SiCl, PMDETA, anisole, and pyridine were distilled from CaH2 prior to use; CuBr was purified according to a literature procedure.⁴⁸ Cu powder was activated according to a literature procedure²⁴ and used immediately. The synthesis of 2-trimethylstannylthiophene was reported previously. 16a All other chemicals were used as received without further purification. Reactions and manipulations involving reactive boron species were carried out under an atmosphere of prepurified nitrogen using either Schlenk techniques or an inert-atmosphere glovebox (Innovative Technologies). Ether solvents were distilled from Na/ benzophenone prior to use. Hydrocarbon and chlorinated solvents were purified using a solvent purification system (Innovative Technologies; alumina/copper columns for hydrocarbon solvents), and the chlorinated solvents were subsequently distilled from CaH2 and degassed via several freeze-pump-thaw cycles. The 499.893 MHz ¹H and 125.681 MHz ¹³C NMR spectra were recorded on a Varian INOVA 500 MHz spectrometer. The 399.844 MHz ¹H and $100.541~\mathrm{MHz}$ $^{13}\mathrm{C}$ NMR spectra were recorded on a Varian VXR-S 400 MHz spectrometer. The 160.4 MHz ¹¹B NMR spectra were recorded on the Varian INOVA 500 spectrometer equipped with a boron-free probe using boron-free quartz NMR tubes. All solution ¹H and ¹³C NMR spectra were referenced internally to the solvent peaks. 11B NMR spectra were referenced externally to BF3. Et2O $(\delta = 0)$ in C₆D₆. GPC analyses were performed in THF (1 mL/ min) using a Waters Breeze system equipped with a 717plus autosampler, a 1525 binary HPLC pump, a 2487 dual λ absorbance detector, and a 2414 refractive index detector. Two Styragel columns (Polymer Laboratories; 5 µm Mix-C), which were kept in a column heater at 35 °C, were used for separation. The columns were calibrated with polystyrene standards (Polymer Laboratories). Multiangle laser light scattering (MALLS) experiments were performed at 690 nm (30 mW linear polarized GaAs laser) using a Wyatt Dawn EOS instrument in-line with the GPC; differential refractive indices dn/dc were calculated from in-line GPC-MALLS mode with Wyatt Astra software assuming 100% mass recovery. DSC measurements were performed on a Perkin-Elmer differential scanning calorimeter Pyris 1 system with ca. 10 mg of the polymer and at a scan rate of 20 °C/min. The results reported correspond to the onset from the second heating cycle. TGA measurements were performed on a Perkin-Elmer Pyris 1 thermogravimetric analyzer at a scan rate of 20 °C/min and up to 800 °C.

Caution! BBr₃ is toxic and highly corrosive and should be handled appropriately with great care. Fluorinated grease was used for ground glass joints in all reactions involving boron tribromide.

Synthesis of 1-Ethyl-4-trimethylsilylbenzene.²² To a suspension of magnesium (7.0 g, 0.29 mol) in 300 mL of THF was added slowly 20 mL of 1-bromo-4-ethylbenzene (26.9 g, 0.145 mol). The reaction mixture was subsequently heated to reflux under nitrogen for 2 h and cooled down to 0 °C. Chlorotrimethylsilane (25 mL, 21.4 g, 0.197 mol) was added slowly, and the reaction mixture was stirred at room temperature overnight. After standard aqueous workup, the product 1-ethyl-4-trimethylsilylbenzene was distilled (31 °C, 10^{-2} Torr) and collected as a colorless liquid (22.8 g, 88.3%). ¹H NMR (399.851 MHz, C₆D₆): δ = 7.45 (d, ³J = 8 Hz, 2H, Ph-H2,6), 7.12 (d, ³J = 8 Hz, 2H, Ph-H3,5), 2.47 (q, ³J = 7.6 Hz, 2H, CH₂CH₃), 1.10 (t, ³J = 7.6 Hz, 3H, CH₂CH₃), 0.23 (s, 9H, SiMe₃).

Synthesis of 1-Bromo-1-(4-trimethylsilylphenyl)ethane (PEB-SiMe₃).²² 1-Ethyl-4-trimethylsilylbenzene (22.5 g, 0.126 mol) was dissolved in 300 mL of CCl₄, and benzoyl peroxide (0.2 g) and N-bromosuccinimide (22.3 g, 0.126 mol) were added in one portion. The reaction mixture was heated to reflux for 12 h while being irradiated by a 450 W Hanovia medium-pressure Hg broadband UV lamp through standard Pyrex glassware. After cooling, the solid succinimide was removed by filtration, and the filtrate was washed with saturated aqueous NaHSO₃. The organic layer was dried over Na₂SO₄ and concentrated. The remaining oil was distilled (56 °C, 10^{-2} Torr) to give the product as a colorless liquid (22.7 g, 70.0%). ¹H NMR (499.893 MHz, C₆D₆): $\delta = 7.34$ (d, ³J = 8 Hz, 2H, Ph-H2,6), 7.21 (d, ${}^{3}J = 8$ Hz, 2H, Ph-H3,5), 4.81 (q, ${}^{3}J = 7.0$ Hz, 1H, CHBr), 1.71 (d, ${}^{3}J = 7.0$ Hz, 1H, CHC H_3), 0.17 (s, 9H, SiMe₃). ¹³C NMR (125.698 MHz, C_6D_6): $\delta = 144.1$ (Ph-C1), 140.6 (Ph-C4), 133.9 (Ph-C2,6), 126.5 (Ph-C3,5), 49.5 (CHBr), 26.8 $(CHCH_3)$, -1.2 (SiMe₃).

Solution Polymerization of Styrene with PEB-SiMe₃: Synthesis of PS-SiMe₃. Styrene (57 g, 0.547 mol), 1-bromo-1-(4trimethylsilylphenyl)ethane (2.80 g, 10.9 mmol), CuBr (1.57 g, 10.9 mmol), and PMDETA (1.90 g, 10.9 mmol) were dissolved in 57 g of anisole, and the mixture was degassed by three freezepump-thaw cycles. The reaction mixture was then heated to 110 °C under nitrogen for 90 min and precipitated into methanol (1 L). The precipitate was redissolved in THF and passed through a short plug of neutral alumina gel. Precipitation into methanol gave a white solid, which was collected by filtration and dried under vacuum at 60 °C overnight (24 g, 42%). ¹H NMR (499.893 MHz, CDCl₃): δ = 7.4-6.3 (all aromatic H's), 4.6-4.3 (m, CHBr), 2.6-1.2(polymer backbone), 1.05 (m, 3H, CHCH₃), 0.26 (9H, SiMe₃). ¹³C NMR (125.698 MHz, C_6D_6): $\delta = 145.7$ (br, styrene Ph-C1), 133.7 (SiPh-C2,6), 126.8 (SiPh-C3,5), 126.1 (br, styrene Ph-C's), 47-40 (polymer backbone), -0.95 (SiMe₃). GPC-RI: $M_n = 4038$, M_w = 4469, PDI = 1.11. GPC-MALLS (THF): M_n = 4256, M_w = 4653, PDI = 1.09, dn/dc (25 °C in THF) = 0.179 mL/g. DSC (onset): $T_g = 80$ °C; TGA (onset): $T_{dec} = 409$ °C, 100% weight loss. Another sample was obtained under similar conditions with similar molecular weight. GPC-RI: $M_{\rm n}=3688,\ M_{\rm w}=4051,$ PDI = 1.10. Both samples were used to prepare monotelechelic organoboron polymers as described below.

Bulk Polymerization of Styrene with PEB-SiMe₃: Synthesis of PS-SiMe₃(b). 1-Bromo-1-(4-trimethylsilylphenyl)ethane (1.76 g, 6.85 mmol), CuBr (0.98 g, 6.85 mmol), CuBr₂ (0.077 g, 0.34 mmol), and PMDETA (1.25 g, 7.2 mmol) were dissolved in styrene (71.4 g, 0.69 mol); the mixture was purged with N₂ for 0.5 h and then immersed into an oil bath preset at 80 °C. After 1 h the polymer was recovered by precipitation into methanol (\sim 1 L), redissolved in THF, and passed through a short plug of neutral alumina gel. Precipitation into methanol gave a white solid, which was collected by filtration and dried under vacuum at 60 °C overnight (19.6 g, 28%). GPC-RI: $M_n = 3358$, $M_w = 3919$, PDI = 1.17. The sample was used to prepare the ditelechelic polymer Me₃Si-PS-SiMe₃ via ATRC.

Synthesis of Me₃Si-PS-SiMe₃. PS-SiMe₃(b) ($M_n = 3358$ by GPC-RI, 3.0 g, 0.76 mmol end group units), CuBr (0.11 g, 0.76 mmol), PMDETA (0.66 g, 3.8 mmol), Cu metal (0.19 g, 3.0 mmol), and 20 g of anisole were charged into a Schlenk flask; the mixture was degassed by three freeze—pump—thaw cycles and then immersed into an oil bath preset at 90 °C. The reaction was monitored by GPC analysis and stopped after 3 h by cooling down to room temperature and passing through a short plug of neutral alumina gel. The solution was then concentrated and precipitated into methanol. The solid obtained was further purified by precipitation from THF into methanol/hexanes (50/50) to remove uncoupled low molecular weight material and finally into methanol and dried

at 60 °C under high vacuum overnight (1.8 g, 60%). GPC-RI: $M_{\rm n}=6439,\,M_{\rm w}=7532,\,{\rm PDI}=1.17.\,{\rm DSC}$ (onset): $T_{\rm g}=95$ °C; TGA (onset): $T_{\rm dec}=405$ °C, 100% weight loss.

Synthesis of PS-BTh₂. To a solution of PS-SiMe₃ ($M_n = 3688$ by GPC-RI, 4.0 g, 1.14 mmol) in ∼40 mL of CH₂Cl₂ was added slowly a solution of BBr₃ (0.32 g, 1.26 mmol) in \sim 10 mL of CH₂-Cl₂. The reaction mixture was kept stirring overnight, and 2-trimethylstannylthiophene (0.78 g, 3.16 mmol) in ~10 mL of CH₂Cl₂ was added slowly. The mixture was stirred for another 12 h and concentrated to about 5 mL before precipitation into cold hexanes (\sim 500 mL, -30 °C). The hexanes were decanted, and the precipitate was dried under vacuum to give 3.24 g of white powdery material (81%). ¹H NMR (399.844 MHz, CDCl₃): $\delta = 7.99$ (Th-H5), 7.83 (Th-H3), 7.70 (BPh-H2,6), 7.40 (Th-H4), 7.3-6.3 (all other aromatic H's), 2.4-1.2 (polymer backbone), 1.12 (CHCH₃). ¹³C NMR (125.698 MHz, CDCl₃): $\delta = 145.9$ (br, styrene Ph-C1), 142.6 (Th-C3/5), 137.5 (BPh-C2,6), 137.2 (Th-C3/5), 129.1 (Th-C4), 128.1, 125.9 (m, styrene Ph-C's), n.o. (BPh-C1), 47-40 (polymer backbone). 11B NMR (160.380 MHz, CDCl₃): $\delta = 50 \ (w_{1/2} = 3800 \ \text{Hz}). \ \text{GPC-RI:} \ M_{\text{n}} = 3517, \ M_{\text{w}} = 3828,$ PDI = 1.09.

Synthesis of Th₂B-PS-BTh₂. To a solution of Me₃Si-PS-SiMe₃ ($M_n = 6439$ by GPC-RI, 0.30 g, 0.093 mmol) in \sim 10 mL of CH₂-Cl₂ was added slowly a solution of BBr₃ (45 mg, 0.18 mmol) in \sim 10 mL of CH₂Cl₂. The reaction mixture was kept stirring overnight, and 0.14 g of 2-trimethylstannylthiophene (0.56 mmol) in \sim 5 mL of CH₂Cl₂ was added slowly. The mixture was stirred for another 12 h and concentrated to about 1 mL before precipitation into cold hexanes (\sim 100 mL, -30 °C). The precipitate was isolated by filtration and dried under vacuum to give 0.25 g of a white powdery material (83%). ¹H NMR (499.893 MHz, CDCl₃): $\delta = 7.97$ (Th-H5), 7.80 (Th-H3), 7.63 (BPh-H2,6), 7.38 (Th-H4), 7.3–6.2 (all other aromatic H's), 2.4–1.2 (polymer backbone). GPC-RI: $M_n = 6270$, $M_w = 7311$, PDI = 1.17. DSC (onset): $T_g = 88$ °C; TGA (onset): $T_{dec} = 400$ °C, 97% weight loss.

Complexation with Pyridine: Synthesis of PS-BTh₂·Py. PS-BTh₂ ($M_n = 3517$ by GPC-RI, 0.50 g, 0.143 mmol) was dissolved in 10 mL of CH₂Cl₂, and pyridine (13.6 mg, 0.171 mmol) was added. The mixture was stirred for 1 h, and all volatile material was removed under high vacuum. PS-BTh₂·Py was recovered quantitatively as a white solid. 1 H NMR (399.844 MHz, CDCl₃, ca. 10 mM boryl groups): $\delta = 8.55$ (Py-H2,6), 7.80 (Py-H4), 7.49 (Py-H3,5), 7.42 (Th-H5), 7.3–6.3 (all other aromatic H's), 2.5–1.2 (polymer backbone), 1.07 (CHC H_3). 13 C NMR (100.541 MHz, CDCl₃): $\delta = 154.4$ (Th-C2), 147.4 (Py-C2,6), 145.8 (styrene Ph-C1), 140.8 (Py-C4), 133.3 (BPh-C2,6), 132.4 (Th-C3), 127.5 (Th-C5), 125.7 (Th-C4), 125.1 (Py-C3,5), 128.1, 126.2 (all other aromatic C's), n.o. (BPh-C1), 47–40 (polymer backbone). 11 B NMR (160.380 MHz, CDCl₃): $\delta = 0$ ($w_{1/2} = 800$ Hz). GPC-RI: $M_n = 2379$, $M_w = 2905$, PDI = 1.22.

Complexation with 4,4'-Dipyridyl (bipy): Synthesis of [Th₂B-PS-BTh₂·bipy]_n. Th₂B-PS-BTh₂ (50 mg, 15.5 μmol) and 4,4'-dipyridyl (1.2 mg, 7.7 μmol) were dissolved in 1 mL of benzene, and [Th₂B-PS-BTh₂·bipy]_n was recovered by freeze-drying. ¹H NMR (399.844 MHz, CDCl₃, ca. 15.5 mM boryl groups): δ = 8.72 (bipy-H2,6), 7.59 (bipy-H3,5), 7.45 (Th-H5), 7.3–6.3 (all other aromatic H's), 2.4–1.2 (polymer backbone). ¹¹B NMR (160.380 MHz, CDCl₃): δ = 1 (w_{1/2} = 1370 Hz). GPC-RI: M_n = 6144, M_w = 7092, PDI = 1.15. DSC (onset): T_g = 105 °C; TGA (onset): T_{dec} = 400 °C, 100% weight loss.

General Procedure of the Titration of Th₂B-PS-BTh₂ with 4,4'-Dipyridyl. Th₂B-PS-BTh₂ (50 mg, ca. 15.5 μ mol boryl groups) was dissolved in 1 mL of CDCl₃ and charged into a NMR tube equipped with a rubber septum. A solution of 4,4'-dipyridyl (ca. 77 mM in CDCl₃) was added in predetermined increments using a microliter syringe. ¹H NMR measurements were performed after each addition of the base solution.

Synthesis of (PS-BO)₃. To a solution of BBr₃ (72 mg, 0.29 mmol) in 20 mL of CH_2Cl_2 was added dropwise a solution of PS-SiMe₃ ($M_n = 4038$ by GPC-RI, 1.0 g, 0.23 mmol end groups) in 10 mL of CH_2Cl_2 , and the mixture was stirred for 12 h. The solution

was then concentrated to \sim 2 mL and precipitated into cold hexanes (−30 °C) to give the intermediate PS-BBr₂ as a gummy material. PS-BBr₂ was freeze-dried from benzene and dried under high vacuum at 50 °C for 12 h. ¹H NMR (399.844 MHz, CDCl₃): δ = 8.04 (BPh-H2,6), 7.4–6.4 (other aromatic H's), 2.6–1.0 (polymer backbone). ¹¹B NMR (160.380 MHz, CDCl₃): $\delta = 56 (w_{1/2} = 1850)$ Hz). PS-BBr₂ was then dissolved in benzene (50 mL), and water (0.5 mL) was added. After azeotropic distillation until no more water separated, the product was isolated by freeze-drying and dried under high vacuum at 60 °C overnight (0.80 g, 80%). ¹H NMR (499.893 MHz, CDCl₃): $\delta = 8.05$ (m, BPh-H2,6), 7.6–6.4 (other aromatic H's), 2.6-1.0 (polymer backbone). ¹H NMR (499.893 MHz, d_6 -acetone/D₂O): $\delta = 7.80-7.71$ (m, BPh-H2,6), 7.4-6.4 (other aromatic H's), 2.5-1.0 (polymer backbone). 11B NMR (160.380 MHz, CDCl₃): $\delta = 22$ ($w_{1/2} = 2840$ Hz). GPC-RI: M_n = 3771, $M_{\rm w}$ = 5219, PDI = 1.38. DSC (onset): $T_{\rm g}$ = 101 °C; TGA (onset): $T_{\text{dec}} = 394 \,^{\circ}\text{C}$, 98.5% weight loss.

Formation of Boroxine Cross-Linked PS: Synthesis of $(PS-BO)_n$. Me₃Si-PS-SiMe₃ ($M_n = 6439$ by GPC-RI, 0.50 g, 0.063 mmol) was dissolved in 10 mL of CH₂Cl₂, and a solution of BBr₃ (63 mg, 0.25 mmol) in 10 mL of CH₂Cl₂ was added dropwise. The reaction mixture was stirred overnight and concentrated to ~ 1 mL. Br₂B-PS-BBr₂ was recovered by precipitation into cold hexanes and redissolved in ~20 mL of benzene. Water (1 mL) was added to the benzene solution, and the mixture was subjected to azeotropic distillation for 6 h until no more water separated and a white precipitate formed. Benzene was then removed under vacuum, and the remaining white solid was dried at 60 °C under high vacuum for 12 h (0.31 g, 62%). ¹H NMR (499.893 MHz, d_5 -pyridine): δ = 8.33 (br, BPh-H2,6), 7.5-6.6 (other aromatic H's), 2.6-1.0(polymer backbone). ¹¹B NMR (160.380 MHz, d_5 -pyridine): $\delta =$ 15. ¹H NMR (499.893 MHz, d_6 -acetone/D₂O): $\delta = 7.80-7.71$ (m, BPh-H2,6), 7.4-6.4 (other aromatic H's), 2.5-1.2 (polymer backbone). DSC (onset): $T_{\rm g} = 112$ °C; TGA (onset): $T_{\rm dec} =$ 395 °C, 97% weight loss.

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Supporting Information Available: ¹H NMR plots of PS-SiMe₃, Me₃Si-PS-SiMe₃, PS-BBr₂, and Br₂B-PS-BBr₂. This material is available free of charge via the Internet at http://pubs.acs.org.

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- (47) Iovine, P. M.; Fletcher, M. N.; Lin, S. Macromolecules 2006, 39, 6324–6326.
- (48) Whitesides, G. M.; Sadowski, J. S.; Lilburn, J. J. Am. Chem. Soc. 1974, 96, 2829–2835.

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