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Cite this: *Inorg. Chem. Front.*, 2020, 7, 3961

## Syndio- and *cis*-1,4 dually selective copolymerization of polar fluorostyrene and butadiene using rare-earth metal catalysts†

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Synthesizing functional butadiene–styrene rubber through coordination polymerization is a theoretical challenge for polymer science, since functional monomers usually deactivate to the applied catalyst. Herein, we report the coordination copolymerization of polar *para*-fluorostyrene (*p*FS) and butadiene (BD) using pyridyl–methylene–fluorenyl supported complexes  $[(\text{Py}-\text{CH}_2-\text{Flu})\text{Ln}(\text{CH}_2\text{SiMe}_3)_2(\text{THF})_n$  ( $\text{Ln} = \text{Sc}$  (**1a**),  $n = 0$ ;  $\text{Ln} = \text{Lu}$  (**1b**),  $n = 1$ )] and pyridyl–cyclopentadienyl supported complexes  $[(\text{Py}-\text{Cp})\text{Ln}(\eta^3-\text{C}_3\text{H}_5)_2$  ( $\text{Ln} = \text{Sc}$  (**2a**),  $\text{Lu}$  (**2b**))]. Strikingly, complexes **2a** and **2b** exhibited dually  $>99\%$  syndio- and  $>95\%$  *cis*-1,4 regio- selectivities and showed obvious characteristics of living polymerization. The insertion of *p*FS can be facilely tuned in the full range of 0–100% by changing the *p*FS-to-BD feed ratio. Diblock P(*p*FS-BD) copolymers were isolated by concurrent addition of monomers and the kinetics study of the copolymerization reaction revealed that BD had the privilege to coordinate to the active metal center. Interestingly, the polymerizations of BD and *p*FS via pulse loading of BD afforded multi-block copolymers of a novel type of fluoro styrene–butadiene rubber with high thermal stability ( $T_d = 368$  °C). The microstructures of resultant copolymers were confirmed by <sup>1</sup>H and <sup>13</sup>C NMR measurements and different phase morphologies of the di- and multi-block polymers were displayed through atomic force microscopy (AFM).

Received 18th June 2020,  
Accepted 21st August 2020

DOI: 10.1039/d0qi00719f  
[rsc.li/frontiers-inorganic](http://rsc.li/frontiers-inorganic)

## Introduction

Synthetic rubbers have played an irreplaceable role in our modern society since we are living in a world “on wheels”.<sup>1</sup> On the other hand these materials are difficult to reuse and recycle and it is nearly impossible for them to be bio-degraded, and so they are the main source of “black pollution”.<sup>2,3</sup> For the sustainable development of tyre and rubber industries, fabricating “green tyres” has been the challenging project in the past three decades.<sup>2</sup> One of the strategies is to synthesize functional rubbers to improve their compatibility with polar fillers, anticipated to endow tyres with excellent wear resistance (prolonged use time) and low rolling resistance without compromising the wet-skid resistance.<sup>4,5</sup> Styrene–butadiene rubber (SBR) is the most widely used synthetic rubber in passenger cars, to fabricate green tyres, by mixing SBR with functionalized fillers.<sup>6–9</sup> End-functionalizing SBR prepared by anionic

solution polymerization with a halogen group is a convenient method to achieve this target; however, obviously, the number of the incorporated functional groups is rather limited.<sup>10,11</sup> Copolymerization with polar group modified monomers is an alternative, anticipated to control the amount and distribution of polar groups in the main chain.<sup>12–16</sup> Radical copolymerization of styrene with nitrile-functionalized butadiene,<sup>17</sup> 2,3-bis(4-ethoxy-4-oxobutyl)-1,3-butadiene<sup>18</sup> or *N,N*-diethyl-2-methylene-3-butenamide (DEA),<sup>19</sup> respectively, provides functionalized poly(styrene–butadiene) rubbers, where the *cis*-1,4 selectivity of the functional butadiene is rather low and its incorporation also needs to improve. Coordination copolymerization is the most powerful method to adjust the regio- and stereoregularity, when the active species are prone to be deactivated by functional groups.<sup>20–22</sup> We reported for the first time the coordination homo- and copolymerizations of polar styrene monomers including methoxystyrene (MOS) and halogen substituted styrenes with the use of side-armed half-sandwich rare-earth metal complexes, with styrene exhibiting high activity and stereoselectivity.<sup>23–25</sup> Thereafter copolymers composed of polar styrenes/ethylene and anisylpropylenes/ethylene were obtained.<sup>26–28</sup> With respect to polar dienes, copolymerization of 2-(2-methylidenebut-3-enyl)furan (MBEF) with isoprene without a masking reagent was achieved by using the bis

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† Electronic supplementary information (ESI) available. See DOI: [10.1039/d0qi00719f](https://doi.org/10.1039/d0qi00719f)

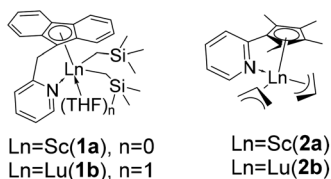


Chart 1 Structures of rare-earth metal catalysts 1 and 2.

(phosphino)carbazolideyttrium precursor.<sup>29</sup> The amino-functionalized styrene and butadiene were mediated to copolymerize; however, a solvent fractionation experiment was needed to obtain pure copolymers.<sup>30</sup> Guo *et al.* recently realized the copolymerization of 4-(*N,N*-diphenylamino)styrene and isoprene to produce functionalized poly(isoprene-styrene).<sup>31</sup>

Despite remarkable characteristics such as low surface energy and superior chemical resistance of fluoropolymers<sup>32</sup> as well as the excellent properties of thermoplastic syndiotactic poly(*p*-fluorostyrene) and elastic *cis*-1,4-PBD,<sup>33</sup> incorporating all these factors into one macromolecular chain, such as fluorinated butadiene-styrene copolymers, with highly regulated microstructures has remained a promising but challenging subject for academic and industrial fields.

Herein, we report the coordination copolymerization of *p*-fluorostyrene (*p*FS) and butadiene (BD) by using rare-earth metal precursors bearing various sterics and electronics (Chart 1). When the sterically less hindered and Lewis acidic rare-earth metal precursors are used, copolymerization of *p*FS and BD occurs rather successfully, in particular in the living mode. The incorporation of polar *p*FS can be adjusted facilely by adjusting the feed-ratio, and its distribution can be di-block and multi-block according to the loading mode. The resultant P(*p*FS-BD) copolymers have >99% syndioregularity for P(*p*FS) sequences and >95% *cis*-1,4-tacticity for PBD sequences, which show a much higher thermal stability ( $T_d = 368\text{ }^\circ\text{C}$ ) than that of commercial SBR.

## Experimental section

### General experimental procedures

All experiments were performed under a dry and oxygen-free nitrogen atmosphere using standard high vacuum Schlenk techniques or in a glove box. All solvents were purified *via* a solvent purification system. *p*-Fluorostyrene was purchased from Aldrich and dried by  $\text{CaH}_2$  under stirring for 48 hours and distilled before use. Catalysts 1 and 2 were synthesized according to the literature.<sup>34,35</sup>  $^1\text{H}$ ,  $^{13}\text{C}$  NMR spectra were recorded on a Bruker AV400 (FT, 400 MHz for  $^1\text{H}$ ; 100 MHz for  $^{13}\text{C}$ ) or AV500 (FT, 500 MHz for  $^1\text{H}$ ; 125 MHz for  $^{13}\text{C}$ ) at 25 °C and  $\text{CDCl}_3$  was used as the solvent with  $\text{CHCl}_3$  as the internal standard (7.26 ppm in  $^1\text{H}$  NMR and  $77.16 \pm 0.06$  ppm in  $^{13}\text{C}$  NMR). The molecular weights ( $M_n$ ) and molecular weight distributions of the polymers ( $M_w/M_n$ ) were measured by TOSOH HLC-8220 GPC at 40 °C using THF as the eluent (the flow rate was 0.35 mL min<sup>-1</sup>) against polystyrene standards. Differential

scanning calorimetry analyses were carried out on a Q100 DSC from TA Instruments under a nitrogen atmosphere at heating and cooling rates of 10 °C min<sup>-1</sup>. Thermo-gravimetric analysis (TGA) was carried out on the TA Instruments SDT Q600 under a nitrogen atmosphere to characterize the thermal stability. AFM was used to study the surface topography of the spin-coated film. Images were obtained using a SPI3800N AFM (Seiko Instruments Inc., Japan). The cantilevers were operated slightly below their resonance frequency of around 20–150 kHz. Image acquisition was performed under ambient conditions. AFM was used in the tapping mode to reduce tip-induced surface degradation and sample damages. Imaging was conducted in the height and phase modes. The sample was prepared by dipping a silicon wafer into a tetrahydrofuran solution of a polymer sample and then placing it on a flat poly(tetrafluoroethylene) plate to allow the solvent to gradually evaporate. Next, the wafer was heated in a vacuum oven at 40 °C to eradicate the solvent before test.

### Homopolymerization of *p*FS

A typical procedure for polymerizing *p*FS is as follows (Table 2, entry 1): under a nitrogen atmosphere and at 20 °C, a toluene (0.5 mL) solution of  $[\text{Ph}_3\text{C}][\text{B}(\text{C}_6\text{F}_5)_4]$  (4.8 mg, 5  $\mu\text{mol}$ ) and a toluene solution (0.5 mL) of complex 2a (1.9 mg, 5  $\mu\text{mol}$ ) were added to a 10 mL flask. Then purified *p*FS (0.122 g, 1.0 mmol) was added under stirring to initiate the polymerization. The reaction was terminated after 20 min by the addition of a small amount of acidic methanol and the mixture was poured into methanol (50 mL). The precipitated polymer was collected by filtration and dried under vacuum at 40 °C to a constant weight (0.122 g, >99%). All other polymerization data were obtained by following the same procedure but with different *p*FS or catalyst feed amounts.

### Copolymerization of *p*FS and BD

A typical procedure for butadiene and *p*FS copolymerization by the concurrent addition of both monomers is as follows (Table 1, entry 5): a toluene (1 mL) solution of  $[\text{Ph}_3\text{C}][\text{B}(\text{C}_6\text{F}_5)_4]$  (4.8 mg, 5  $\mu\text{mol}$ ) and a toluene solution (1 mL) of complex 2a (1.9 mg, 5  $\mu\text{mol}$ ) were added to a 10 mL flask under a nitrogen atmosphere and the mixture was kept under stirring for a few minutes. Then a mixture of *p*FS (0.30 g, 2.5 mmol) and butadiene (0.135 g, 2.5 mmol, 19%<sub>wt</sub> in toluene) was added. The reaction was terminated after 10 min by the addition of a small amount of acidic methanol and the mixture was poured into methanol (50 mL). The precipitated copolymer was collected by filtration and dried under vacuum at 40 °C to a constant weight (0.294 g, 66%). All other polymerization data were obtained following the same procedure but with different *p*FS-to-BD feed ratios or catalyst.

### Synthesis of multi-block P(*p*FS-BD)

A typical procedure for synthesizing a multi-block P(*p*FS-BD) is as follows: a toluene (0.5 mL) solution of  $[\text{Ph}_3\text{C}][\text{B}(\text{C}_6\text{F}_5)_4]$  (4.8 mg, 5  $\mu\text{mol}$ ) and a toluene solution (0.5 mL) of complex 2a (1.9 mg, 5  $\mu\text{mol}$ ) were added to a 10 mL flask under a nitro-

Table 1 Copolymerization of *para*-fluorostyrene and butadiene<sup>a</sup>

Entry	Cat	<i>p</i> FS fed (mol %)	Time/min	Conv. (%)	<i>p</i> FS <sup>b</sup> (mol%)	<i>rrrr</i> <sup>b</sup> (%)	Selectivity <sup>b</sup> (%)		<i>M<sub>n</sub></i> <sup>c</sup> × 10 <sup>-4</sup>	<i>M<sub>w</sub></i> / <i>M<sub>n</sub></i> <sup>c</sup>
							<i>cis</i> -1,4 : <i>trans</i> -1,4 : 1,2			
1 <sup>d</sup>	<b>1a</b>	50	10	60.4	n. d	n. d	n. d	n. d	n. d	n. d
2 <sup>d</sup>	<b>1b</b>	50	60	32.8	34.2	n. d	n. d	16.7/1.5	1.1/1.8	
3	<b>2a</b>	90	10	90.7	89.1	>99	95 : 3 : 2	25.7	1.3	
4	<b>2a</b>	70	10	76.0	63.8	>99	95 : 3 : 2	21.7	1.2	
5	<b>2a</b>	50	10	66.1	38.7	>99	96 : 3 : 1	15.8	1.1	
6	<b>2a</b>	30	10	47.4	4.3	>99	97 : 2 : 1	15.5	1.1	
7	<b>2a</b>	10	10	79.3	3.6	>99	97 : 2 : 1	13.8	1.1	
8	<b>2b</b>	90	60	84.6	89.6	>99	96 : 2 : 2	24.1	1.3	
9	<b>2b</b>	70	60	61.2	56.8	>99	96 : 2 : 2	13.4	1.5	
10	<b>2b</b>	50	60	41.1	16.9	>99	96 : 2 : 2	12.4	1.2	
11	<b>2b</b>	30	60	49.5	6.8	>99	96 : 2 : 2	15.2	1.2	
12	<b>2b</b>	10	60	80.0	3.5	>99	96 : 2 : 2	17.0	1.2	

<sup>a</sup> Conditions: [Sc] 5 μmol, [pFS] + [BD]/[Sc]/[Ph<sub>3</sub>C][B(C<sub>6</sub>F<sub>5</sub>)<sub>4</sub>] = 1000 : 1 : 1 (mol/mol/mol), toluene 2 mL, *T* = 20 °C. <sup>b</sup> Measured by <sup>1</sup>H NMR and <sup>13</sup>C NMR in CDCl<sub>3</sub> at 25 °C. <sup>c</sup> Determined by GPC in THF at 40 °C against polystyrene standard. <sup>d</sup> [Sc]/[Al<sup>i</sup>Bu] = 1 : 10.

gen atmosphere. The mixture was stirred at room temperature for a few minutes. As soon as purified *p*FS (0.3 g, 2.5 mmol) was added into the above system, butadiene (0.26 g, 4.8 mmol, 13%<sub>wt</sub> in toluene) was added dropwise over 10 min. Then the reaction was terminated by the addition of a small amount of acidic methanol followed by pouring the mixture into methanol (50 mL) to precipitate the copolymer product. The copolymer product was collected by filtration and dried under vacuum at 40 °C to a constant weight (0.47 g, 84%).

#### Kinetics studies for *p*FS and BD copolymerization

To a 50 mL flask were added a toluene solution (5 mL) containing [Ph<sub>3</sub>C][B(C<sub>6</sub>F<sub>5</sub>)<sub>4</sub>] (24 mg, 25 μmol) and complex **2a** (9.5 mg, 25 μmol), under a nitrogen atmosphere at 20 °C. After vigorous stirring for a few minutes, a mixture of *p*FS (0.61 g, 5.0 mmol) and butadiene (0.27 g, 5.0 mmol, 17%<sub>wt</sub> in toluene) was added to the above system, which was evenly divided into five portions, and the reaction was terminated by adding acidic methanol at the set time (44s, 61s, 75s, 92s, 122s), respectively. Each copolymer was collected by filtration and was dried under vacuum at 40 °C and weighed for calculating the conversions of BD and *p*FS. The kinetics study for the copolymerization of *p*FS and BD by complex **2b** was performed following the same procedure as described above.

## Results and discussion

First, the copolymerization of *p*-fluorostyrene (*p*FS) and butadiene (BD) in toluene at room temperature was investigated by using the catalytic systems of **1**+[Ph<sub>3</sub>C][B(C<sub>6</sub>F<sub>5</sub>)<sub>4</sub>]/[Al<sup>i</sup>Bu], which are highly active towards the homopolymerization of *p*FS and the copolymerization with styrene and ethylene, respectively.<sup>25,26</sup> The polymerization catalyzed by **1a** (Flu<sub>cent</sub>-Sc-N (96.26°)) occurred fast but gave a copolymer insoluble in many solvents even under a high temperature of 160 °C. We proposed that cross-linking took place, since the scandium active species is Lewis acidic and can initiate polymerization of the dangling C=C bonds of the 1,2-regulated PBD segments.

When switched to catalyst **1b** (Flu<sub>cent</sub>-Lu-N (92.16°)), the polymerization activity turned out to be very slow. The GPC curve of the isolated polymer shows multi-peaks, suggesting poor controllability of catalyst **1b** during polymerization (Table 1, entries 1 and 2). Catalytic systems based on complexes **2a** and **2b** with smaller bite angles (Cp<sub>cent</sub>-Sc-N (86.6°) and Cp<sub>cent</sub>-Lu-N (84.1°)) possess dual *cis*-1,4 regio- and perfect stereo-selectivities for the polymerization of butadiene and styrene, respectively; however, whether they are tolerant to polar fluorine substituted monomers is still unknown.<sup>36</sup> Thus the homopolymerization of *p*FS using complex **2a** or **2b** upon activation with [Ph<sub>3</sub>C][B(C<sub>6</sub>F<sub>5</sub>)<sub>4</sub>] was first attempted. Complex **2a** showed a high activity to consume *p*FS completely in 20 min and provided perfect syndiotactic P(*p*FS) (*rrrr* > 99%, Fig. 1, entry 1). Changing the *p*FS-to-scandium molar ratio from 200 : 1 to 400 : 1, the molecular weight (*M<sub>n</sub>*) of the resultant poly(*p*FS) increased from 5.2 × 10<sup>4</sup> to 10.3 × 10<sup>4</sup> while the molecular weight distribution remained constant and narrow (Table 2 entries 1 and 2). In addition, the *M<sub>n</sub>* of P(*p*FS) increased linearly in proportion to the monomer conversion ([*p*FS]/[Complex **2a**]/[Ph<sub>3</sub>C][B(C<sub>6</sub>F<sub>5</sub>)<sub>4</sub>]) = 1000/1/1) as shown in Fig. 2, suggesting a living polymerization mode. On the other

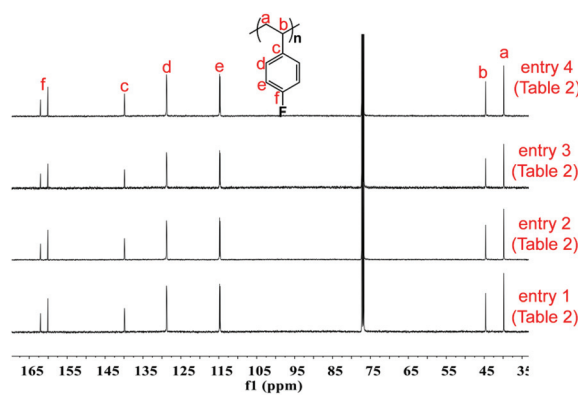


Table 2 Polymerization of *para*-fluorostyrene<sup>a</sup>

Entry	Cat	<i>p</i> FS/Cat	Time (min)	Conv. (%)	<i>rrrr</i> <sup>b</sup> (%)	$M_n \times 10^{-4}$	$M_w/M_n$	$T_m$ (°C)
1	2a	200	20	>99	>99	5.2	1.7	323
2	2a	400	20	>99	>99	10.3	2.3	323
3	2b	200	60	51	>99	3.7	1.6	323
4	2b	400	60	50	>99	5.7	2.0	323

<sup>a</sup> Conditions: [Sc] 5 μmol, [Sc]/[Ph<sub>3</sub>C][B(C<sub>6</sub>F<sub>5</sub>)<sub>4</sub>] = 1:1 (mol/mol), Toluene 1 mL, *T* = 20 °C. <sup>b</sup> Measured by <sup>1</sup>H NMR and <sup>13</sup>C NMR in CDCl<sub>3</sub> at 25 °C. <sup>c</sup> Determined by GPC in THF at 40 °C against polystyrene standard. <sup>d</sup> Determined by DSC.

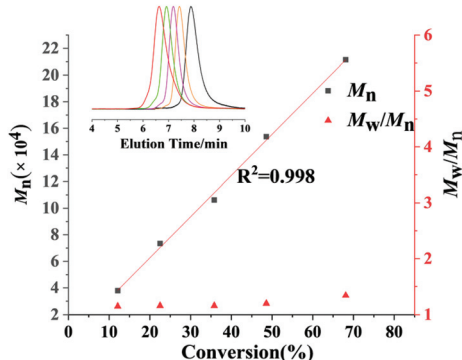


Fig. 2 Linear growth of molecular weight ( $M_n$ ) as a function of *p*FS conversion. Inset: GPC traces as detected from the retention time.

hand, complex **2b** also showed a perfect syndioselectivity albeit in a low activity, probably owing to the less acidic lutetium metal center.

The excellent catalysis of **2**/[Ph<sub>3</sub>C][B(C<sub>6</sub>F<sub>5</sub>)<sub>4</sub>] for highly selective polymerization of *p*FS prompted us to explore its copolymerization behavior with BD. The copolymerization with **2a** was performed in toluene at 20 °C for 10 min by the concurrent addition of the two monomers under the *p*FS molar fraction varying from 10 to 90 mol%. As shown in Table 1, high conversions were obtained in the full range of monomer feed ratios, indicating distinguished high activities. The <sup>1</sup>H NMR spectrum of the resultant polymer (Fig. 3 (top)) (Table 2, entry 4) shows that the peaks assigned to phenyl protons in P(*p*FS) units appear at  $\delta$  = 6.29–6.89 ppm, whilst the peaks arising from olefinic protons in the 1,4-tactic PBD segments appear at  $\delta$  = 5.30–5.46 ppm. According to the integration intensity ratio of the phenyl and olefinic protons, it was found that the *p*FS molar fraction in the copolymers varies corresponding to the loaded one, reaching up to 89.1 mol% when the *p*FS feed ratio was 90 mol%. Catalyst **2b** showed a parallel catalytic behavior to **2a** albeit with lower activity, which is much better than its performance for the homopolymerization of *p*FS.

Gel permeation chromatography (GPC) analyses of these crude P(*p*FS-BD) copolymers without extraction show that the molecular weights are high ( $M_n$  = 12.4–25.7  $\times 10^4$ ) and the molecular weight distributions are unimodal and narrow (PDI

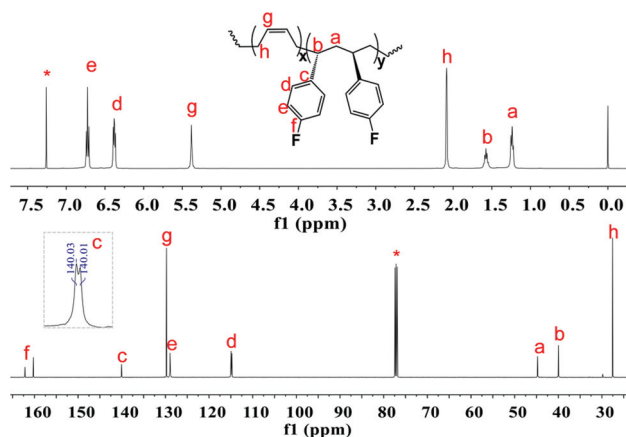


Fig. 3 <sup>1</sup>H and <sup>13</sup>C NMR spectra of poly(*p*FS-BD) (Table 1, entry 4) (500 MHz for <sup>1</sup>H NMR and 125 MHz for <sup>13</sup>C NMR, CDCl<sub>3</sub>, 25 °C).

= 1.1–1.5), suggesting that copolymers were obtained successfully instead of a mixture of homopolymers, which is consistent with the single-sited catalytic behavior and obvious characteristics of the living copolymerization fashion.

The microstructures of these copolymers were defined by <sup>13</sup>C NMR spectroscopy analyses. The representative spectrum of a poly(*p*FS-BD) product with a moderate *p*FS content of 63.8 mol% is shown in Fig. 3 (bottom) (Table 2, entry 4). The *ipso*-carbon C<sub>c</sub> giving a doublet at  $\delta$  = 140.02 ppm with a coupling constant <sup>4</sup>J<sub>C-F</sub> = 2.5 Hz indicates the syndiotactic (*rrrr* > 99%) P(*p*FS) block. The strong singlet at  $\delta$  = 27.42 ppm indicates a high degree of *cis*-1,4-PBD unit (>95%). However, the resonances arising from the carbon–carbon linkages of *p*FS-BD joints are not observed. The <sup>13</sup>C NMR spectra of the copolymers with different *p*FS contents show similar topologies. Meanwhile the *cis*-1,4 tacticity of the PBD segments remains unchanged (95–97%) in all copolymers, which is not affected by the *p*FS content. All these characterizations suggest that the copolymers obtained by using **2a**/[Ph<sub>3</sub>C][B(C<sub>6</sub>F<sub>5</sub>)<sub>4</sub>] might have a diblock sequence. To confirm this deduction, a kinetics study of the copolymerization reaction in toluene at 20 °C by the concurrent addition of both monomers was carried out. As shown in Fig. 4 (top), when the polymerization catalyzed by **2a**/[Ph<sub>3</sub>C][B(C<sub>6</sub>F<sub>5</sub>)<sub>4</sub>] was carried out for 1 min, the isolated product was almost pure PBD corresponding to the almost complete conversion of BD (>99%), whereas no *p*FS incorporation was found. With the polymerization going on and consumption of BD monomer, the *p*FS fraction in the copolymer increased dramatically. A similar phenomenon was observed using the **2b**/[Ph<sub>3</sub>C][B(C<sub>6</sub>F<sub>5</sub>)<sub>4</sub>] system (Fig. 4 (bottom)). These results indicate that in the presence of *p*FS, BD has the priority when coordinating to the active metal center and propagating into polymer. Until BD is almost fully consumed, *p*FS starts to insert into Sc-PBD active species to generate the diblock structure without homopolymer contaminants.

Multi-block copolymers as a kind of significant macromolecular materials exhibit outstanding properties. To reuse and

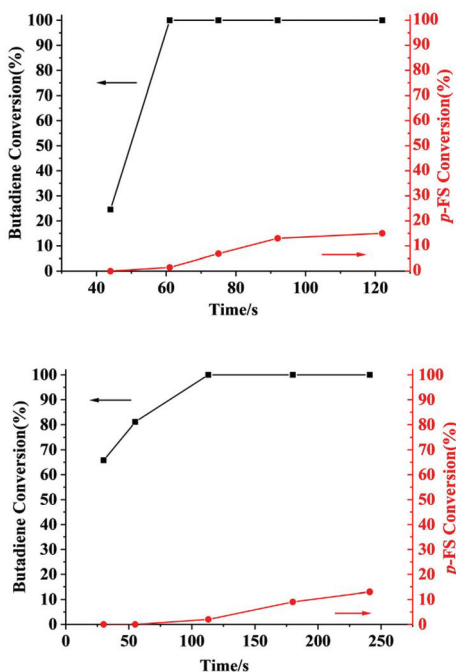


Fig. 4 Monomer conversions versus time with catalytic systems **2a** (top) and **2b** (bottom).

strengthen the interfaces of abandoned PE and iPP, Coates *et al.* produced PE/iPP multi-block polymers by pyridylamido-hafnium catalyst activated with  $B(C_6F_5)_3$ .<sup>37</sup> Miller and Ellison synthesized multi-block poly(ethylene terephthalate)-polyethylene, which enhances the ability to recycle PET/PE mixed waste streams.<sup>38</sup> In other aspects, multi-block sulfonated poly(arylene ether sulfone)s are used in polymer electrolyte membrane fuel cells.<sup>39</sup> According to the living polymerization mode of BD and *p*FS in the catalyst **2a** system, a multiple BD feeding method was adopted to exclude the formation of long BD and *p*FS sequences for preparing multi-block P(*p*FS-BD) copolymers. A typical procedure was carried out in which *p*FS (0.3 g) was added in one portion into the polymerization system catalyzed by **2a**/[Ph<sub>3</sub>C][B(C<sub>6</sub>F<sub>5</sub>)<sub>4</sub>] (5  $\mu$ mol), and then a BD solution (2.0 g, 13%<sub>wt</sub>) was added over 10 min. As soon as the addition of the BD solution was completed, the copolymerization was terminated and a copolymer with 26.7 mol% *p*FS content was isolated (0.47 g,  $M_n = 13.9 \times 10^4$ , PDI = 1.9). Its molecular weight distribution gives a unimodal peak (Fig. S32†), suggesting successful copolymerization. All the resonances in the <sup>1</sup>H NMR spectrum of the multi-block copolymer are broadened as compared to the corresponding resonances in the diblock copolymer, especially at the aromatic region (Fig. 5). In the <sup>13</sup>C spectrum, the resonances of carbons from the joints of the multi-block copolymer are observed (m, n, o, p and q points in Fig. 6). The averaged chain lengths of PBD and P(*p*FS) in the multi-block copolymer obtained by calculating the ratios of the integration intensities of the corresponding repeat units and the joints in the quantitative <sup>13</sup>C NMR spectrum are 161 and 27, respectively (Fig. S21†). Since the highly

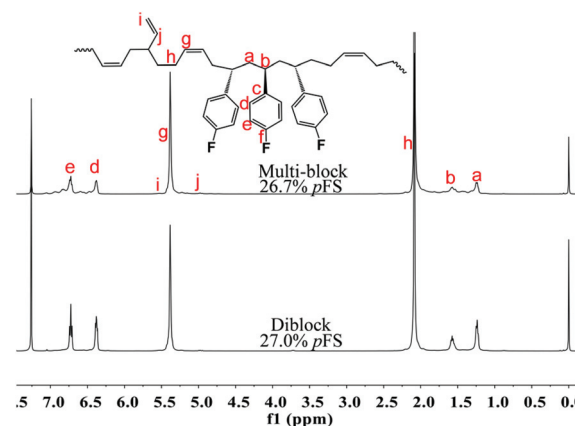


Fig. 5 <sup>1</sup>H NMR spectra of multi-block and diblock poly(*p*FS-BD) (500 MHz, CDCl<sub>3</sub>, 25 °C).

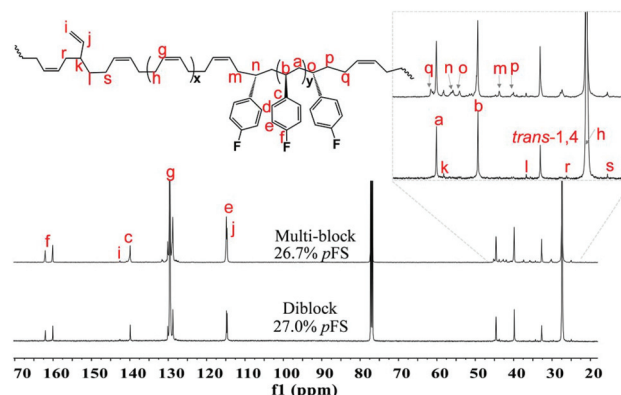


Fig. 6 <sup>13</sup>C NMR spectra of multi-block and diblock poly(*p*FS-BD) (125 MHz, CDCl<sub>3</sub>, 25 °C).

syndiotactic P(*p*FS) is crystalline,<sup>35</sup> the diblock copolymer with a 27.0 mol% P(*p*FS) content (one long sequence) has a microphase separation rather different from the multi-block copolymer with a similar P(*p*FS) content (26.7 mol%, many short sequences). As shown in Fig. 7, the AFM micrograph of the diblock copolymer (left) reveals a larger degree of phase separation than the multi-block one (right). In addition, there is no

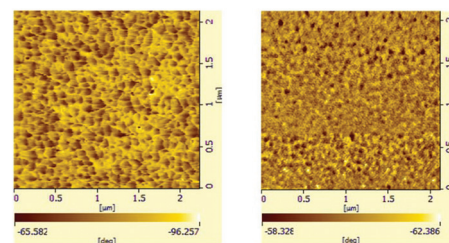


Fig. 7 AFM micrographs of a spin-coated thin film of diblock poly(*p*FS-BD) (left) (Table 2, entry 10, x(*p*FS) = 26.7 mol%) and multi-block poly(*p*FS-BD) (right) (x(*p*FS) = 27.0 mol%) (phase images, scan size: 2.0  $\mu$ m).

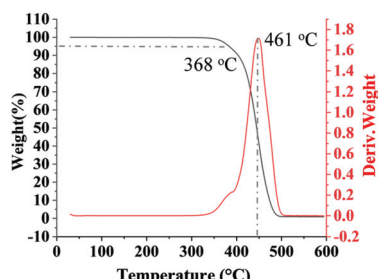


Fig. 8 TG and DTG curves of multi-block poly(*p*FS-BD).

obvious melting point in the multi-block P(*p*FS-BD) as compared with the di-block one ( $T_m = 316$  °C) in the curves of the differential scanning calorimetric analyses (Fig. S34†). The 5% mass loss temperature of the multi-block copolymer is at 368 °C, which is much higher than that of the commercial SBR (Fig. 8),<sup>40</sup> indicating that it is a quite different and more thermally stable material.

## Conclusions

In summary, we have demonstrated the unprecedented highly *cis*-1,4 selective (>95%) and perfectly syndiospecific (>99%) copolymerization of BD and *p*FS using pyridyl functionalized Cp ligated CGC rare-earth metal bis(allyl) complexes, which provide an active rare-earth metal center with an open coordination sphere to avoid low *cis*-1,4 selectivity and cross-linking process. In addition, the excellent tolerance for polar groups of these catalysts permits the insertion of *p*FS facilely tuned in the full range of 0–100% by changing the *p*FS-to-BD ratio. The diblock sequence copolymer is obtained by the concurrent addition of *p*FS and BD, with BD as the privilege, which is consistent with the kinetics study. By the merit of the living copolymerization characteristics, remarkably, a multi-block copolymer is isolated, for the first time, *via* pulse loading of butadiene. An unexpectedly high thermal stability was found for the multi-block copolymer during TGA testing. This work paves a new way to access functional styrene–butadiene copolymers with controllable regio- and stereo-regularities as well as sequence distribution. The mechanical properties of the di- and multi-block copolymers and the comparison of these copolymers with the SBR prepared by the traditional anionic mechanism are under investigation.

## Conflicts of interest

There are no conflicts to declare.

## Acknowledgements

The authors acknowledge the financial support from NSFC (projects No.21634007, 21774118, 21674108 and U1862108),

and Department of Science and Technology of Jilin Province project No. 20190201067JC.

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