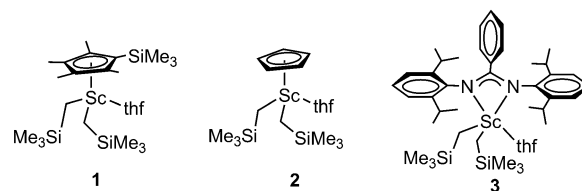


Chain-Shuttling Polymerization at Two Different Scandium Sites: Regio- and Stereospecific “One-Pot” Block Copolymerization of Styrene, Isoprene, and Butadiene**

Li Pan, Kunyu Zhang, Masayoshi Nishiura, and Zhaomin Hou*

The development of new generations of polymerization catalysts and technology for efficient, selective synthesis of novel polymer materials having well-controlled microstructures and desired properties has been a long-standing research subject of both academic and industrial scientists. In this endeavor, exploring the potential of untapped elements is an important strategy. Recently, the chain-shuttling copolymerization of two different monomers by two Group 4 metal catalysts that show different monomer selectivities has emerged as a powerful technology for the preparation of olefin block copolymers with different segments.^[1] In contrast, the potential of rare-earth catalysts in such chain-shuttling copolymerization has remained unexplored to date.^[2,3] Moreover, the success of chain-shuttling copolymerization reported to date has been limited mostly to the copolymerization of ethylene and propylene (or 1-hexene), while the chain-shuttling copolymerization of styrene and a conjugated diene such as isoprene or butadiene has not been reported.

We report herein the first example of chain-shuttling copolymerization of styrene, isoprene, and butadiene by two different scandium catalysts. In the presence of scandium catalyst **1** (Scheme 1), which shows high activity and high selectivity for the syndiotactic polymerization of styrene but low activity and low selectivity for isoprene,^[4a-c] and scandium catalyst **2**, which shows high activity and high *cis*-1,4-selectivity for the polymerization of isoprene^[4c] but low activity and low selectivity for styrene, together with a chain-shuttling agent, the copolymerization of styrene and isoprene has been achieved regio- and stereoselectively at each monomer, affording for the first time block copolymers with perfect syndiotactic polystyrene (sPS) and highly regulated *cis*-1,4-polyisoprene (PIP) blocks. In a similar manner, the terpolymerization of styrene, isoprene, and butadiene has also been achieved for the first time to give terpolymers containing perfect sPS, highly regulated *cis*-1,4-PIP and highly



Scheme 1. Scandium catalysts of different activities and selectivities.

1: High activity and high syndiospecific selectivity for styrene polymerization but low activity and low selectivity for isoprene polymerization. **2:** High activity and high *cis*-1,4-selectivity for isoprene or butadiene polymerization but low activity and low selectivity for styrene polymerization. **3:** High activity and high 3,4-selectivity for isoprene polymerization but low activity for styrene polymerization.

regulated *cis*-1,4-PBD blocks. By replacement of the catalyst **2** with a catalyst **3**,^[4d] which showed high 3,4-selectivity for the polymerization of isoprene in the chain-shuttling polymerization of styrene and isoprene, novel styrene–isoprene block copolymers with sPS and 3,4-PIP blocks have been obtained for the first time.

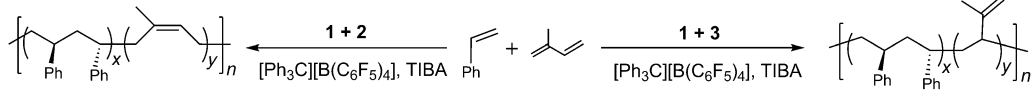
Some representative results of the polymerization and copolymerization of styrene and isoprene by three different Sc catalysts **1**, **2**, and **3** (Scheme 1) under various conditions are summarized in Table 1. As reported, the half-sandwich scandium dialkyl complex **1** bearing the sterically demanding C₅Me₅SiMe₃ ligand in combination with [Ph₃C][B(C₆F₅)₄] showed high activity and high syndiospecific selectivity (*rrrr* > 99%) for the polymerization of styrene (Table 1, run 1),^[4a] but low activity and poor regio- and stereoselectivity for the polymerization of isoprene (Table 1, run 4).^[4b,c] The smaller C₅H₅-bound Sc analogue **2** exhibited much lower activity and no stereoselectivity for the polymerization of styrene (Table 1, run 2) but showed high activity and high *cis*-1,4-selectivity (97%) for the polymerization of isoprene (Table 1, run 5) under the same conditions.^[4c] In the presence of both **1** and **2** without a chain-shuttling agent, the copolymerization of styrene and isoprene gave a polymer mixture of *cis*-1,4-PIP homopolymer and a styrene–isoprene copolymer containing mainly sPS blocks with mixed 1,4-/3,4-PIP units, as confirmed by solvent extraction (Table 1, run 7). In sharp contrast, when five equivalents (per Sc center) of *i*Bu₃Al (TIBA) were added to the above reaction system, the copolymerization of styrene and isoprene took place selectively and rapidly to give the block copolymer containing perfect sPS (*rrrr* > 99%) and highly regulated *cis*-1,4-PIP (97% selectivity) blocks with high molecular weight (*M*_n = 109.4 kg mol^{−1}) and narrow molecular-weight distribution (*M*_w/*M*_n = 1.43, Table 1, run 8). When the amount of TIBA was increased from five equiv-

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[**] This work was supported by a Grant-in-Aid for Scientific Research (S) (No. 21225004) from Japan Society for the Promotion of Science.

Supporting information for this article is available on the WWW under <http://dx.doi.org/10.1002/ange.201104011>.

Table 1: Chain-shuttling copolymerization of styrene and isoprene by two different scandium catalysts.^[a]

														
Run	Cat.	St/IP ^[b]	TIBA/ [Sc] ^[c]	t [min]	Conv. [%]	M _n ^[d] [kg mol ⁻¹]	M _w /M _n ^[d]	sPS [%]	PIP [%]	Microstructure of PIP ^[e]			T _m ^[f] [°C]	T _g ^[f] [°C]
										<i>cis</i> -1,4-	<i>trans</i> -1,4-	3,4-		
1	1	500/0	0	1	100	154.5	1.24	> 99	—	—	—	—	272	95
2	2	500/0	0	60	59	5.8	1.34	— ^[g]	—	—	—	—	—	90
3	3	500/0	0	60	< 1	—	—	—	—	—	—	—	—	—
4	1	0/500	0	60	27	24.5	1.31	—	100	5	54	41	—	−38
5	2	0/500	0	5	100	86.7	2.15	—	100	97	0	3	—	−61
6	3	0/500	0	5	100	63.3	1.71	—	100	1	< 1	98	—	40
7	1 + 2	250/250	0	20	35, ^[h] 92 ^[i]	39.8 ^[j] 82.4 ^[k]	1.40 1.90	92	8	4	50	46	261	—
8	1 + 2	250/250	5	10	100	109.4	1.43	50	50	97	0	3	272	−59
9	1 + 2	250/250	10	10	100	71.5	1.52	50	50	98	0	2	270	−59
10	1 + 2	250/250	20	10	100	46.2	1.79	50	50	98	0	2	268	−60
11	1 + 2	500/500	20	60	100	96.7	1.75	50	50	98	0	2	267	−59
12	1 + 2	1000/1000	20	120	100	194.3	1.67	50	50	98	0	2	268	−59
13	1 + 2	500/250	10	10	100	126.1	1.78	67	33	97	0	3	270	−60
14	1 + 2	250/500	10	10	100	87.2	1.54	34	66	97	0	3	267	−60
15	1 + 2	125/125	20	10	100	27.8	1.66	50	50	98	0	2	268	−59
16		2nd 125/125 ^[l]	20	30	100	40.7	1.78	50	50	98	0	2	268	−59
17		3rd 125/125 ^[l]	20	30	100	52.6	1.89	50	50	98	0	2	268	−59
18		4th 125/125 ^[l]	20	60	100	65.2	1.94	50	50	98	0	2	268	−59
19	1 + 3	250/250	0	60	100	70.4 ^[j] 75.1 ^[k]	1.60 1.41	77 21 ^[m]	23 79	16 9	32 8	52 83	254 —	6 26
20	1 + 3	250/250	5	10	100	67.2	1.79	50	50	10	2	88	263	22
21	1 + 3	250/250	10	10	100	26.5	1.68	50	50	8	2	90	262	35

[a] Conditions: catalyst feed: [1] = [2] = [3] = 20 μmol, [Ph₃C][B(C₆F₅)₄]/[Sc] = 1, toluene V = 40 mL, 25 °C. [b] Molar ratio of monomer to total catalyst feed [Sc]. [c] Molar ratio of tri(isobutyl)aluminum (TIBA) to total catalyst used. [d] Determined by GPC against PS standards in *o*-dichlorobenzene at 140 °C or in THF at 40 °C. [e] Molar ratio determined by ¹³C NMR spectroscopy in 1,1,2,2-tetrachloroethane. [f] Measured by differential scanning calorimetry. [g] Atactic polystyrene. [h] Conversion of styrene. [i] Conversion of isoprene. [j] THF-insoluble portion. [k] THF-soluble portion. [l] Sequential addition of 125 equiv styrene and 125 equiv of isoprene to run 15. [m] *a*PS.

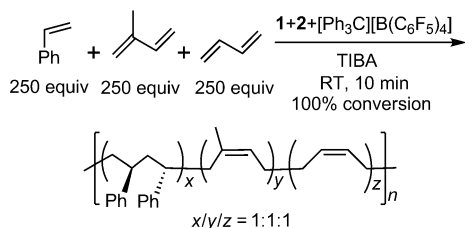
alents to 10 and 20 equivalents, the molecular weight of the resulting copolymers decreased significantly, but the molecular-weight distribution remained rather narrow and the syndiospecific selectivity for styrene (> 99%) and *cis*-1,4-selectivity for isoprene (98%) remained constantly high (Table 1, runs 9 and 10). In the presence of 20 equivalents TIBA, the increase of the monomer/catalyst ratio resulted in an almost linear increase of the molecular weight of the copolymer products, while the molecular-weight distribution and the selectivity for either styrene or isoprene remained almost unchanged (Table 1, runs 10–12). The styrene/isoprene content in the resulting copolymers could be easily changed simply by changing the styrene/isoprene feed ratio (Table 1, runs 13 and 14). Sequential additions (four steps) of a 125/125 mixture of styrene and isoprene to the chain-shuttling copolymerization catalyst system afforded the copolymer products with increased molecular weights (*M_n* from 27.8 to 65.2 kg mol⁻¹) and almost constant molecular-weight distributions (*M_w*/*M_n* = 1.66–1.94) as well as consistently high syndiospecific selectivity for styrene (> 99%) and *cis*-1,4-selectivity for isoprene (98%, Table 1, runs 15–18), suggesting that the chain-shuttling reactions in the present polymerization system are highly reversible and efficient. The resulting styrene–isoprene copolymers showed melting points around 270 °C originating from the *s*PS blocks^[4a,b,5] and glass

transition temperatures (*T_g*) at approximately −60 °C originating from the *cis*-1,4-PIP blocks^[4d,6] (Table 1, runs 8–18).

To further confirm the composition of the copolymer products obtained in chain-shuttling copolymerization, solvent (THF) extraction of the crude polymer product obtained in Table 1, run 9 was carried out. The THF-soluble portion (32 wt %) showed a *s*PS content of 18 mol % and a *cis*-1,4-PIP content of 82 mol % with *M_n* = 68.2 kg mol⁻¹ and *M_w*/*M_n* = 1.45. The THF-insoluble portion (68 wt %) showed *s*PS content of 72 mol %, *cis*-1,4-PIP content of 28 mol %, *M_n* = 74.2 kg mol⁻¹, and *M_w*/*M_n* = 1.61. These results suggest that this crude polymer product contains predominantly *s*PS–*cis*-1,4-PIP block copolymers rather than a mixture of homopolymers.

The present chain-shuttling copolymerization of styrene and isoprene constitutes the first efficient protocol for the preparation of styrene–isoprene copolymers having perfect *s*PS and highly regulated *cis*-1,4-PIP blocks. *s*PS is a well-known semicrystalline polymer which possesses many useful properties, such as high modulus of elasticity and excellent resistance to heat and chemicals,^[5] while *cis*-1,4-PIP is a major component of natural rubber and can serve as an excellent elastomer.^[6] Although styrene–isoprene copolymers with both *s*PS and highly regulated *cis*-1,4-PIP blocks are of much interest, such highly stereoregulated copolymers are difficult to prepare and have not been reported to date.^[4b]

The scandium amidinate complex **3**, similar to its known yttrium analogue,^[4d] showed high activity for the 3,4-polymerization of isoprene (selectivity 98%, Table 1, run 6) but very poor activity for the polymerization of styrene (Table 1, run 3). In the presence of **1** and **3** without a chain-shuttling agent, the copolymerization of styrene and isoprene afforded a crude copolymer product containing PS and PIP sequences. Solvent extraction experiments revealed that the THF-soluble portion (44 wt%) contained blocks rich in 3,4-PIP (79 mol%, 83% 3,4-selectivity) and atactic PS units (21 mol%), while the THF-insoluble portion (56 wt%) possessed *s*PS blocks (77 mol%) and mixed 1,4-/3,4-PIP units (23 mol%; Table 1, run 19). In contrast, in the presence



*s*PS > 99%, *cis*-1,4-PIP > 97 mol%, *cis*-1,4-PBD > 97 mol%

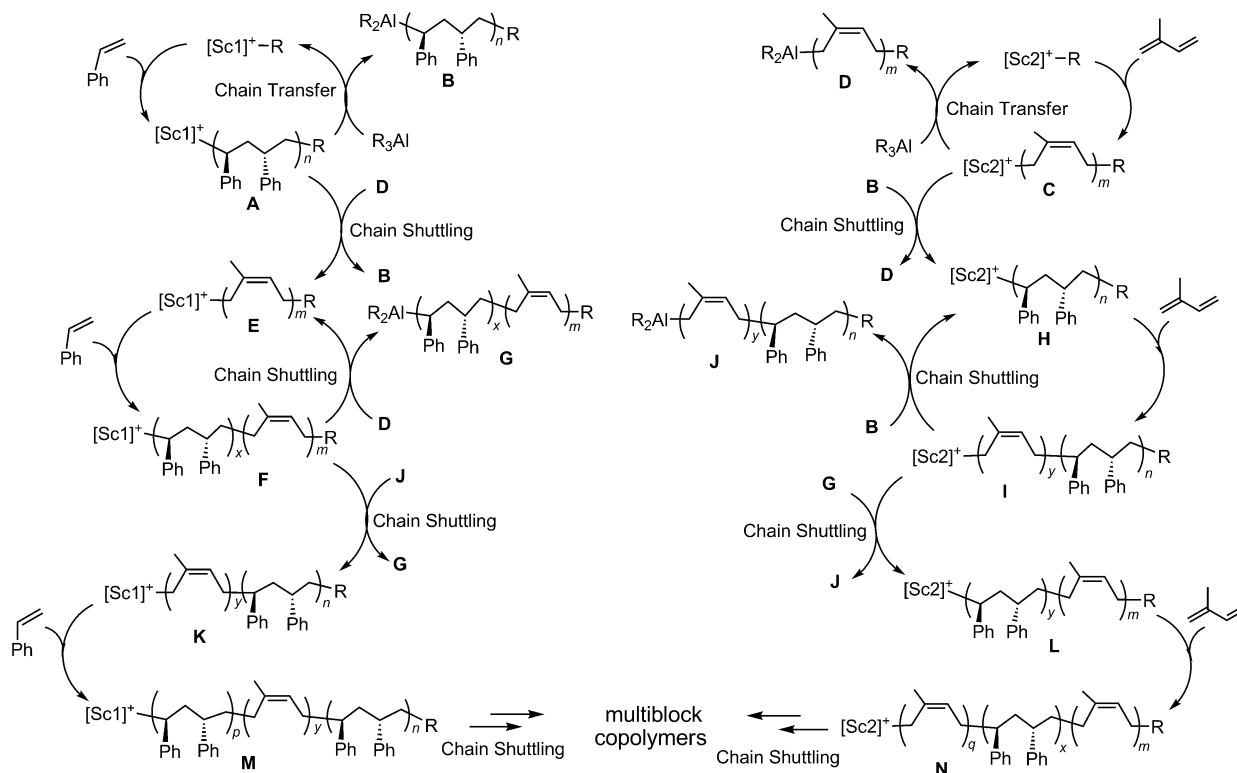
5 equiv TIBA: $M_w = 166.8 \text{ kg mol}^{-1}$, $M_w/M_n = 1.38$, $T_m = 267^\circ\text{C}$, $T_g = -77^\circ\text{C}$
10 equiv TIBA: $M_w = 94.4 \text{ kg mol}^{-1}$, $M_w/M_n = 1.50$, $T_m = 266^\circ\text{C}$, $T_g = -76^\circ\text{C}$

Scheme 2. Scandium-catalyzed regio- and stereospecific terpolymerization of styrene, isoprene, and butadiene in the presence of a chain-shuttling agent (TIBA).

of TIBA, the copolymerization of styrene and isoprene took place selectively and more rapidly to give the corresponding block copolymers containing perfect *s*PS (*rrrr* > 99%) and 3,4-PIP (ca. 90%) blocks (Table 1, runs 20 and 21). The resulting copolymers showed T_g around room temperature (22–35 °C) originating from the 3,4-PIP blocks,^[4d,7] in addition to melting points at approximately 263 °C originating from the *s*PS blocks (Table 1, runs 20 and 21). This is the first example of styrene–isoprene copolymerization with high syndiospecific selectivity for styrene and high 3,4-selectivity for isoprene.

By use of the combination of **1** and **2** with TIBA as a chain-shuttling agent, the terpolymerization of styrene, isoprene, and butadiene has also been achieved to give rapidly and selectively novel block copolymers containing *s*PS (*rrrr* > 99%), *cis*-1,4-PIP (selectivity > 97%), and *cis*-1,4-PBD (selectivity > 97%) blocks for the first time (Scheme 2). Such highly stereoregulated *s*PS-*cis*-1,4-PIP-*cis*-1,4-PBD copolymers should be of particular interest in comparison with the nonstereoregular styrene–isoprene–butadiene polymers (SIBR) prepared by conventional anionic polymerization.^[8,9]

A possible scenario for the copolymerization of styrene and isoprene by the catalyst combination **1** + **2** in the presence of TIBA is shown in Scheme 3. Chain growth of styrene at the active site $[\text{Sc1}]^+-\text{R}$ generated from **1** and $[\text{Ph}_3\text{C}][\text{B}(\text{C}_6\text{F}_5)_4]$ should give a scandium active species bearing a *s*PS block such as **A**,^[4a,b] which after transmetalation with TIBA (R_3Al ,



Scheme 3. A possible chain-shuttling mechanism for the formation of multiblock *s*PS-*cis*-1,4-PIP copolymers by the combination of catalysts **1** and **2**. $[\text{Sc1}]$ denotes an active species generated from **1**, which shows high activity and syndiospecific selectivity for styrene polymerization. $[\text{Sc2}]$ denotes an active species generated from **2**, which shows high activity and *cis*-1,4-selectivity for isoprene polymerization. The polymer species containing aluminum end groups, such as **B**, **D**, **G**, and **J**, could undergo reversible transmetalation with scandium active species, thus leading to efficient chain shuttling.

chain transfer) would afford sPS having an R_2Al -functionalized end group (**B**). Analogously, the R_2Al -functionalized *cis*-1,4-PIP species **D** could be generated by IP chain propagation at the $[Sc2]^+$ site formed from **2** and $[Ph_3C][B(C_6F_5)_4]^{[4c]}$ and subsequent transmetalation between the resulting $[Sc2]$ -PIP species **C** and R_3Al . If transmetalation between the $[Sc1]$ -sPS species **A** and the Al -PIP species **D** or between the $[Sc2]$ -*cis*-1,4-PIP species **C** and the Al -sPS species **B** takes place, the $[Sc1]$ -*cis*-1,4-PIP species **E** or the $[Sc2]$ -sPS species **H** would be formed, respectively (chain shuttling). Continuous styrene insertion into **E** could give the $[Sc1]$ -sPS-*cis*-1,4-PIP species **F**, which on transmetalation with the Al -PIP species **D** would afford the Al -functionalized sPS-*cis*-1,4-PIP diblock copolymer **G**. Analogously, the R_2Al -*cis*-1,4-PIP-sPS diblock copolymer **J** could be formed by isoprene insertion into **H** followed by transmetalation between the resultant $[Sc2]$ -*cis*-1,4-PIP-sPS species **I** and the Al -sPS species **B**. The repeated chain-shuttling polymerization should give multi-block styrene-isoprene copolymers with stereoregulated sPS and *cis*-1,4-PIP blocks (Scheme 3). The chain-shuttling terpolymerization of styrene, isoprene, and butadiene by **1** + **2** to give copolymers containing sPS, *cis*-1,4-PIP, and *cis*-1,4-PBD blocks and the copolymerization of styrene and isoprene by **1** + **3** to yield copolymers with sPS and 3,4-PIP blocks could take place in an analogous fashion.

In summary, by use of two scandium catalysts such as **1** and **2** or **1** and **3**, which show different monomer selectivity and stereoselectivity, with TIBA as a chain-shuttling agent, the regio- and stereospecific copolymerization of styrene, isoprene, and butadiene has been achieved for the first time to afford a novel family of copolymer materials composed of highly stereoregulated sPS, *cis*-1,4-PIP, and *cis*-1,4-PBD blocks or copolymers containing sPS and 3,4-PIP blocks. This work demonstrates for the first time that rare-earth-catalyzed chain-shuttling copolymerization of styrene and conjugated dienes can constitute a unique protocol for efficient synthesis of novel polymer materials that are difficult to obtain by other means. Further studies on the chain-shuttling copolymerization of other monomers by rare-earth-based catalysts are in progress.

Received: June 11, 2011

Revised: August 15, 2011

Published online: October 25, 2011

Keywords: block copolymers · chain shuttling · copolymerization · scandium · syndiotactic polystyrene

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