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Zirconocene-Catalyzed Stereoselective Cyclocopolymerization of 2-Methyl-1,5-Hexadiene with Propylene

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Abstract: The copolymerization of 2-methyl-1,5-hexadiene (MHD) with propylene has been studied with different single-site group 4 metal catalysts. Systems based on ansa-zirconocene precursors such as rac-{Me2Si(2-Me-4-Ph-Ind)}ZrCl2 (1) and C1- or C5-symmetric ansa-{CpCR2Flu}ZrCl2 (2 and 3, respectively), once activated by MAO, are highly active (20–600 kg pol./g cat. h-1 at 60–70 °C) and yield copolymers in which MHD is cyclopolymerized as methylene-(1-methyl)-1,3-cyclopentane (MMCP) units. 13C NMR studies revealed, depending on the symmetry of the precatalyst used, either highly isotactic (1, 2) or syndiotactic (3) polypropylene (PP) backbones, with isolated MMCP units. Fully trans-diastereoselective cyclopolymerization of MHD was observed with 1/MAO, while a mixture of trans and cis MMCP rings was observed with 2 and 3/MAO. The amount of MMCP units in PP (0.2–1.6 mol-%) can be controlled by the amount of MHD in the feed. In contrast, the constrained

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geometry catalyst system based on \{C_5Me_4SiMe_2NtBu\}TiCl_2 (4) and MAO showed a much lower productivity (ca. 3 kg_{pol}·g_{cat}^{-1}h^{-1} at 60 °C) and yields a regioirregular, atactic copolymer in which MHD is simply vinyl-inserted in quite moderate amount (0.2 mol-%).
Introduction

Homopolymerization of short, yet non-conjugated diolefins such as 1,5-hexadiene leads to macromolecular structures that include in the polymer main-chain cyclic units interfaced by methylene groups, such as methylene-1,3-cyclopentane (MCP). Initially investigated with heterogeneous Ziegler-Natta catalyst systems, formation of such cyclopolyyolefins from 1,5-hexadiene was later extended with homogeneous group 4 (half) metallocenes and a variety of other molecular catalysts (rare earths: Sc, Y, Sm; V; Pd). The cyclopolymerization was shown to be a two-step process. The enantioface selectivity of olefin insertion determines the tacticity of the polymer, and the diastereoselectivity of the cyclisation step determines whether cis or trans rings are formed. Although the stereoselectivity of the first 1,2 insertion was controlled by the enantiomorphic site of the catalyst (if any), the diastereoselectivity of the second insertion leading to the cyclization was attributed to the bulkiness of the ligand rather than to the symmetry of the catalyst precursor. Hence, for instance, catalyst systems based on Cp\(_2\)ZrCl\(_2\), 2a,c \{EBTHI\}ZrCl\(_2\) (\{EBTHI\} = ethylenebis(tetrahydroindenyl))\(_{2b}\) and half-sandwich scandium complexes\(_{3c}\) yielded mainly trans cycles, while Cp\(^*\)\(_2\)ZrCl\(_2\) yielded mainly cis cycles.\(_{2a,c}\)

Copolymerization of 1,5-hexadiene with ethylene\(^6\) and propylene\(^2g,j,4,7\) mediated by group 4 metallocenes and related single-site catalysts proceeds via intramolecular insertion of the latter diolefin and eventually leads to polyolefins incorporating 5-membered rings.\(^8\) The copolymerization of higher diolefins such as 1,6-heptadiene,\(^9\) 1,7-octadiene and 1,9-decadiene\(^10\) with ethylene and propylene was also studied with different types of catalysts. Both 1,2 insertion and eventual cyclization were observed for 1,6-heptadiene and 1,7-octadiene,\(^9,10\) but only 1,2 insertion occurred for 1,9-decadiene,\(^10\) showing that the critical step for cyclization of the monomer is the favorable formation of an intermediate where the
pendant vinyl-end is coordinated to the metal center. Cycloinsertion of diallylsilane in copolymerization with propylene was also reported, forming 6-membered rings. Examples of catalysts enabling the cyclopolymerization of vinyl-vinylidene monomers are much more limited. Kesti and Waymouth have shown that the homopolymerization of 2-methyl-1,5-hexadiene (MHD) catalyzed by Cp*₂ZrMe₂ combined with various activators (MAO, B(C₆F₅)₃, [PhNHMe₂]⁺[B(C₆F₅)₄]⁻) produces a polymer with a repeating methylene-(1-methyl)-1,3-cyclopentane (MMCP) unit (Scheme 1). The regio- and diastereoselective cyclopolymerization of the monomer results from the incorporation of the vinyl-end of the monomer, followed by the coordination and 1,2-insertion of the vinylidene group. The absence of β-hydrogen in the propagating species prevents β-H elimination termination reactions and the addition of another MHD molecule is favored over the β-methyl elimination. The stereochemistry of the MMCP ring was tentatively assigned as trans on the basis of NMR of oligomers and comparison with poly(methylnorbornene), a model polymer that contains only cis methylcyclopentane rings; it was further consolidated by conformational calculations. On the other hand, the copolymerization of ethylene with 5,7-dimethyl-1,6-octadiene (5,7-DMO), a one-carbon higher homologue of MHD, with a variety of catalytic systems proceeded only via insertion of the vinyl moiety, leaving the vinylidene moiety unreacted. Noteworthy, the possibility to fine control the stereoselectivity of these cyclopolymerizations of vinyl-vinylidene monomers, with catalysts having drastically different stereocontrol abilities, has not been investigated yet in details.
Scheme 1. Homopolymerization of MHD with repeating methylene-(1-methyl)-1,3-cyclopentane (MMCP) motifs.\textsuperscript{12}

Herein, we report on the first attempts to copolymerize propylene with MHD. A series of molecular catalyst precursors, including \textit{ansa}-zirconocenes with quite different stereocontrol (iso- vs. syndioselective) abilities and a “constrained-geometry” hemi-titanocene, has been investigated. Our aim was to study their influence on the catalytic performance (productivity, comonomer incorporation) of the copolymerization process and on the microstructure of the cyclopolymer produced.

**Results and Discussion**

The copolymerization of MHD with propylene has been investigated with a series of \textit{ansa}-zirconocene precatalysts (Figure 1), activated by MAO. These precatalysts were selected due to their known propensity to produce high molecular weight isotactic (1,\textsuperscript{15} 2,\textsuperscript{16}) and syndiotactic (3\textsuperscript{17}) polypropylenes. In addition, we have investigated the essentially non-stereoselective “constrained geometry” hemi-titanocene precatalyst 4, because of its well-known great ability to incorporate \textit{\alpha}-olefins.\textsuperscript{18} Representative results are reported in Table 1. The \textit{ansa}-zirconocenes and hemi-titanocene behaved quite differently and, for the sake of clarity, these two types of catalysts are discussed separately hereafter.
**Figure 1.** Structures of the *ansa*-zirconocenes and constrained geometry hemi-titanocene precatalysts used in this study for the cyclo/copolymerization of MHD with propylene.

**Ansa-Zirconocene Catalysts**

**General Trends.** All three *ansa*-zirconocene catalyst systems investigated afforded selectively P(P-co-MHD) copolymers, as indicated by the unimodal and relatively narrow molecular weight distributions determined by GPC analysis (see the Supporting Information). This is also supported by detailed $^1$H and $^{13}$C NMR analyses (*vide infra*) which evidenced complete cyclopolymerization of MHD with propylene, eventually providing isolated MMCP rings within the PP chain; no pendant vinylidene group was observed in any case (Scheme 2). These first observations suggest that: (*i*) after insertion of a vinyl MHD group, insertion of the MHD vinylidene group is favored over insertion of a propylene molecule; (*ii*) insertion of a propylene molecule after cycloinsertion of a MHD molecule is favored over possible β-Me elimination from a [M]–MMCP intermediate.
Scheme 2. Cyclopolymerization of MHD with propylene promoted by ansa-zirconocene catalysts.

Despite its efficient cyclopolymerization, MHD has a noticeable, slightly detrimental influence on the catalyst productivity/activity. As compared to simple propylene homopolymerizations, the addition of MHD decreased by ca. 25–50% the productivity of the ansa-zirconocene catalyst systems (compare entries 1 vs. 2, 4 vs. 5–8, 9 vs. 10–11).\textsuperscript{19} As illustrated with system 2/MAO,\textsuperscript{20} the amount of MHD incorporated in the copolymers appears to be proportional to the concentration of monomer initially loaded (entries 4–8, Figure 2). Hence, from ca. 0.34 mol-% up to 1.3–1.6 mol-% of MHD (that is from ca. 11 up to 52 insertions for 10,000 carbons) are incorporated by 2/MAO at [MHD]\textsubscript{0} in the range 12–80 mmol.L\textsuperscript{-1} (entries 4–8). In addition, the MHD incorporation rate remains stable during the polymerization course (as long as the MHD concentration does not vary much), as noted with the 3/MAO system after 10 and 20 min (entries 10 and 11). Both isoselective 2 and syndioselective 3 ansa-\{CpCR\textsubscript{2}Flu\}-zirconocene-based systems afforded comparable MHD incorporation rates (1.6 and 1.2 mol-%, respectively, at [MHD]\textsubscript{0} = 80 mmol.L\textsuperscript{-1}; entries 7,8 and 10,11). Apparently, Spaleck’s bis(indenyl)-zirconocene 1 showed, apart from a much higher activity than \{CpCR\textsubscript{2}Flu\}-zirconocene 2, a similar ability to incorporate MHD, with ca. 0.37 mol-%\textsuperscript{21} vs. 0.34 mol-% incorporation rates at [MHD]\textsubscript{0} = 12.0 mmol.L\textsuperscript{-1}, respectively (entries 2 and 5).
Figure 2. Plot of MHD incorporated in PP as a function of initial loading, with catalyst system 2/MAO (Table 1, entries 4, 5, 6 and 8).

For comparison purposes, the copolymerization of higher α-olefins comparable in size with MHD but having no additional polymerizable function, namely 8-trimethylsilyl-1-octene (8-TMSO) and 7-methyl-1,6-octadiene (1,6-MOD), has been carried out using the 5/MAO system (Figure 3), which has similar reactivity and properties as 2/MAO\textsuperscript{16} (Table 1, entries 14–16). The amounts of incorporated 8-TMSO and 1,6-MOD, which are easily determined using the TMS or internal vinylidene groups as NMR probes (see the Supporting Information), are quite comparable to those observed for MHD under similar conditions (ca. 0.3 mol-%). This confirms that it is the rate of the terminal vinyl insertion that determines the ability of cyclic incorporation of MHD.
Figure 3. Structures of precatalyst 5 and higher \( \alpha \)-olefins 8-TMSO and 1,6-MOD.

The cyclopolymerization of MHD affects the molecular features of the PP produced. The P(P-co-MHD) copolymers featured systematically lower molecular weights than the corresponding homoPP (entries 1–2, 4–8 and 9–11). This decrease of the molecular weight follows the same trend as the decrease of the activity/productivity. The incorporation of a monomer more difficult to insert than propylene (\textit{i.e.}, the vinylidene moiety of MHD) results in a kind of “dormant” species and likely leads to a higher probability for this latter species to undergo termination or chain transfer to aluminum reactions. This might also account for the larger dispersity (yet still unimodal) observed with the highly active catalyst 1/MAO (entry 2). On the other hand, copolymers produced by 2/MAO and 3/MAO (entries 5–8 and 10,11) show a relatively narrow dispersity, comparable with that of homoPPs produced by the same catalyst (entries 4 and 9, respectively).

As anticipated, the incorporation of little MHD as MMCP rings induced a small decrease in the stereoregularity compared to the corresponding homoPP, that is ca. 2–3\% (at the \( m^f \) level) for 2 (entries 4–8) to ca. 4\% for 1 (entries 1–3). Syndioselective catalyst 3 followed the same trend with a drop of 3\% at the \( r^d \) level (entries 9–11). The presence of MMCP rings in the polymer backbone also decreased the crystallinity of the copolymers, as evidenced by a pronounced decrease of the melting temperature up to 11 \(^\circ\)C in the case of the copolymer produced with 1/MAO (entries 1–3). For less stereoregular copolymers produced
with 2/MAO or 3/MAO, a difference of only 1–2 °C was noted in the \( T_m \) values (entries 4–8 and 9–11).

**Microstructural NMR Analysis.** The microstructure of the P(P-co-MHD) copolymers produced with the three ansa-zirconocene catalysts was analyzed by \(^{13}\)C\{\(^1\)H\} NMR spectroscopy. It was compared with that of the corresponding PP homopolymers in order to distinguish the resonances specific to the MMCP motifs from those which correspond to intrinsic branching and stereo/regio-defects in the PP backbone (not relevant to this study). In the NMR spectra of isotactic and syndiotactic polymers (Figures 4 and 5, respectively), the absence of a resonance at \( \delta \) ca. 34.5 ppm, which is observed when the cyclic MMCP units are adjacent to one another as in PMHD homopolymers [for the C4 methylene group],\(^{12}\) indicates that all of the MMCP units are separated by propylene units in the copolymers; this was not unexpected when considering the low MHD incorporation rates (< 2 mol-%).

Depending on the ansa-zirconocene catalyst used, one or two sets of resonances assigned to the MMCP motifs incorporated in the PP backbone were identified. The two sets of signals for the MMCP motifs produced by syndioselective catalyst 3/MAO (identified with markers + (major set) and \( \times \) (minor set)) are different from the signals identified for the corresponding motifs formed by isoselective catalysts 1/MAO and 2/MAO (identified with markers * (major) and ° (minor))\(^{22}\) (Table 2). As described hereafter, each of these four sets of signals corresponds to a specific MMCP stereoisomer (trans/cis) formed upon vinyldiene insertion within either an isotactic or a syndiotactic PP backbone (Scheme 3, microstructures trans-isol/cis-iso and trans-syndiol/cis-syndio). A \(^{13}\)C-\(^{13}\)C INADEQUATE NMR experiment was performed on a copolymer produced by 2/MAO and containing 0.5 mol-% of MHD (Table 1, entry 6), enabling a complete assignment of the major set of signals observed for the MMCP motifs (Figure 6, Table 2; the minor set was of too low intensity to observe cross-
peaks); only the resonance of the 1-methyl MMCP (δ ca. 28.5 ppm)\(^{12}\) could not be unambiguously identified, as it overlaps with the CH signals of the iPP backbone.

(insert Figures 4 and 5 here)

Figure 6. \(^{13}\)C-\(^{13}\)C INADEQUATE NMR spectrum (1,2,4-trichlorobenzene/C\(_6\)D\(_6\) (5:1), 125 MHz, 135 °C) of a P(P-co-MHD) copolymer produced by 2/MAO (Table 1, entry 6).
Table 2. Assignment of individual sets of $^{13}$C NMR resonances (in ppm) for cyclic MMCP motifs in isotactic and syndiotactic PP backbones. The right structure shows experimental chemical shifts for the major set (*) of resonances in an iPP backbone and (in parentheses) the corresponding calculated chemical shifts (ACD Lab).

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<th>$^\circ$ (isotactic, minor set; cis-iso)</th>
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<th>$^\times$ (syndiotactic, minor set; cis-syndio)</th>
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<td>21.6</td>
<td>21.2</td>
<td>21.2</td>
</tr>
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</table>

[Diagram showing isospecific and syndiospecific insertion with chemical shifts indicated for major and minor sets.]
**Scheme 3.** Cyclopolymerization mechanism of MHD with propylene by isoselective (left) and syndioselective (right) *ansa*-zirconocene catalysts, leading to the formation of two *trans/cis* MMCP stereoisomers in iPP and sPP backbones, respectively.

In fact, the use of stereoselective catalysts for the copolymerization of MHD and propylene reduces the number of possible stereoisomers. With the {CpCR$_2$Flu}-type isoselective 2/MAO and syndioselective 3/MAO catalysts, two sets of resonances (° (major) o (minor) and + (major) x (minor), respectively) were clearly observed in the $^{13}$C NMR spectra. On the other hand, the incorporation of MHD by isoselective bis(indenyl) catalyst 1/MAO leads to only one observable set (°). Based on earlier observations by Waymouth *et al.*, Naga *et al.*, and computations by Cavallo *et al.*, this latter set can be confidently assigned to the *trans*-iso isomer; similarly, the major and minor sets derived from 2 and 3/MAO can be assigned to the *trans*-iso and *cis*-iso, and *trans*-syndio and *cis*-syndio isomers, respectively. This shows that vinylidene insertion is more stereoselective with 1 than with 2 and 3. This is line with the higher stereoselectivity of this former catalyst system toward propylene homopolymerization, as compared to the latter {CpCR$_2$Flu}-type catalysts (97.7% [$m^4$] for 1/MAO vs. 94.1% [$m^4$] for 2/MAO; Table 1, entries 1 and 4, respectively).

Yet, this difference in isoselectivity between catalysts might not be important enough to justify the absence of the minor MMCP stereoisomer in the copolymer produced by 1/MAO. In line with previous reports, we suspect that the different geometries between, on one hand, the {CpCR$_2$Flu}-type catalysts based on 2 and 3 and, on the other hand, the bis(indenyl) catalyst 1 might also account for the higher stereoselectivity of the process. The key factor in the formation of those stereoisomers is which enantioface of vinylidene is coordinated to the metal center (Scheme 3). In this process, the distance between the free coordination sites and the steric hindrance between the bridging carbons of MHD and the
substituents of the ligand certainly are determining factors in the formation of either one or the other intermediate.

**Hemi-Titanocene “Constrained Geometry” Catalyst**

The “constrained geometry” 4/MAO catalyst system behaved quite differently from the *ansa*-zirconocene ones (Table 1, entries 12, 13). First, under comparable conditions, the catalytic productivity was lower by one or two orders of magnitude. More, although a copolymer with a relatively narrow, unimodal distribution was indeed produced, no cyclization of MHD was observed (Scheme 4). The $^{13}$C NMR spectra of copolymers produced with catalyst 4/MAO proved much more complicated than those of polymers prepared with stereoselective *ansa*-zirconocene catalysts (Figure 7). Despite the complexity of the spectra, arising as anticipated from the presence of many regio-errors (2,1-insertions, as evidenced by signals in the $\delta$ 15–17 ppm region)$^{23}$ and the absence of stereocontrol,$^{18}$ informative observations could be made: i) CH$_2$ signals at $\delta$ 48–50 ppm, characteristic of MMCP rings (*vide supra*) were absent; ii) instead, signals at $\delta$ 110.0 and 146.1 ppm, assignable to the pendant vinylidene group of vinyl-inserted MHD, were observed.$^{24}$ Simple vinyl-insertion of MHD was corroborated by the observation in the $^1$H NMR spectra of signals at $\delta$ 4.69 and 4.67 ppm for the pendant vinylidene group (see Figure S2 in the Supporting Information)$^{24}$
Scheme 4. Vinyl-copolymerization of MHD with propylene promoted by hemi-titanocene “constrained geometry” catalyst 4/MAO.

(insert Figure 7 here)

The amount of MHD inserted was estimated by $^{13}$C NMR to be ca. 0.2 mol-%, that is ca. 5 fold less than with ansa-zirconocenes under similar conditions (compare entries 13 vs. 7/8 and 10/11). Overall, these data show that the 4/MAO system polymerizes much more easily vinyl than vinylidene groups. This was somehow unexpected considering that constrained geometry titanium catalysts have been reported to promote the copolymerization of isoalkenes with olefins (ethylene), yet with bimetallic catalysts leading to incorporation rates significantly enhanced as compared to those achieved with monometallic ones.25

Summary

The copolymerization of MHD with propylene by isoselective and syndioselective ansa-zirconocene catalysts leads to the production of copolymers in which MHD is selectively incorporated as methylene-(1-methyl)-1,3-cyclopentane (MMCP) units via consecutive insertion of the vinyl and vinylidene bonds. The MMCP rings feature specific $^{13}$C NMR resonances, depending on their trans/cis configuration and on the isotactic/syndiotactic PP backbone, and some of them could be assigned by a $^{13}$C-$^{13}$C INADEQUATE NMR experiment. Fully trans-diastereoselective cyclopolymerization of MHD was observed with the bis{indenyl} catalyst 1/MAO, while a mixture of trans (major) and cis (minor) MMCP rings was observed with the {CpCR$_2$Flu}-type catalysts 2 and 3/MAO. This behavior is attributed to the intrinsic higher stereoselectivity of the former catalyst and geometric differences between the catalysts. The copolymers produced with 2 and 3/MAO, which
contains up to 1.6 mol-% of MHD/MMCP, have melting temperatures close to those of the corresponding i/sPP homopolymers. In contrast, incorporation of 0.4–0.6 mol-% of MHD in iPP by 1/MAO resulted in a much more pronounced drop of the melting temperature.

**Experimental section**

**General.** All manipulations were performed under a purified argon atmosphere using standard Schlenk techniques or in a glovebox. Solvents were distilled Na/K alloy (toluene, hexane, pentane) under argon, degassed thoroughly and stored under argon prior to use. *Ansa-*metalallocene rac-\{Me₂Si(2-Me-4-Ph-Ind)ZrCl₂ (1) was generously provided by Total Raffinage Chimie. *Ansa-*metalallocenes 2, 16, 3, 26 and 5, 16 were synthesized as described in the literature. Hemi-titanocene \{C₅Me₄SiMe₂NtBu\}TiCl₂ (4) was purchased from Boulder Scientific. Methyl-1,5-hexadiene (MHD), 8-trimethylsilyl-1-octene (8-TMSO) and 7-methyl-1,6-octadiene (1,6-MOD) were purchased from Aldrich, and kept over activated molecular sieves. Propylene (99.99%) was purchased from Air Liquide and used as received. MAO (30 wt-% solution in toluene, Albermale; contains ca. 10 wt-% of free AlMe₃) was used as received.

\[^{13}\text{C}\{^{1}\text{H}\} \text{NMR analyses of polymer samples were run in the research center of Total Raffinage-Chimie in Feluy (Belgium) on a Avance 500 Bruker spectrometer equipped with a cryoprobe using the following conditions: solutions of ca. 200–700 mg of polymer in Cl₂CDCDCl₂ or in a 1,2,4-trichlorobenzene/C₆D₆ (5:1) mixture at 135 °C in 10 mm tubes, inverse gated experiment, pulse angle = 90°, delay = 30 s, acquisition time = 1.25 s, number of scans = 500–1,000.} \[^{13}\text{C} \text{chemical shifts are reported in ppm using Me₃SiOSiMe₃ (δ 2.03 ppm; equivalent to δ 0.00 ppm for SiMe₄) as internal standard. The MHD incorporation rate was determined as MHD (mol-%) = [average of integral values of MMCP signals] / [integral value for CH of PP + average of integral values of MMCP signals] × 100. The}\]^{13}\text{C}_{-}\text{^{13}\text{C}}
INADEQUATE spectrum was recorded using 700 mg of polymer dissolved in a 10 mm tube in a 1,2,4-trichlorobenzene/C₆D₆ (5:1) mixture at 135 °C, with a 50 ppm spectral width centered at δ 34 ppm, using 240 increments containing each 384 scans, and a delay of 4 s.

DSC measurements were performed on a SETARAM Instrumentation DSC 131 differential scanning calorimeter at a heating rate of 10 °C.min⁻¹; first and second runs were recorded after cooling to 30 °C; the melting temperatures reported in tables correspond to the second run. GPC analyses were carried out in 1,2,4-trichlorobenzene at 135 °C in the research center of Total Raffinage-Chimie in Feluy (Belgium), using polystyrene standards for universal calibration.

**Propylene/2-methyl-1,5-hexadiene copolymerization procedure.** Copolymerization experiments were performed in a 300 mL high-pressure glass reactor equipped with a mechanical stirrer (Pelton turbine) and externally heated with a double mantle with a thermostated circulating water bath. The reactor was charged with toluene (80 to 150 mL), the appropriate amount of 2-methyl-1,5-hexadiene (MHD) and MAO (1.5 mL of a 30 wt-% solution in toluene), and propylene (5 bar) was introduced. The reactor was thermally equilibrated at the desired temperature for 30 min. Propylene pressure was decreased to 1 bar, and a solution of the catalyst precursor in toluene (ca. 2 mL) was added by syringe. The propylene pressure was immediately increased to 5 bar (kept constant with a back regulator) and the solution was stirred for the desired time. The temperature inside the reactor was monitored using a thermocouple. The polymerization was stopped by venting the vessel and quenching with a 10 wt-% solution of aqueous HCl in methanol (ca. 3 mL). The polymer was precipitated in methanol (ca. 200 mL) and 35 wt-% aqueous HCl (ca. 1 mL) was added to dissolve possible catalyst residues. The polymer was collected by filtration, washed with methanol (ca. 200 mL), and dried under vacuum overnight. **Homopolymerization of**
propylene (Table 1, entries 1, 4, 9, 12, 14) and copolymerization of propylene with other model \(\alpha\)-olefins (Table 1, entries 15 and 16) were conducted in a similar way.

**Supporting Information Available.** \(^1\)H NMR spectrum of a P(P-co-MHB) copolymer produced with “constrained geometry” catalyst 4/MAO; \(^1\)H NMR spectra of copolymers of propylene with 8-TMSO and 1,6-MOD; representative GPC and DSC traces of copolymers.

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**Notes and References**

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5 For a review on stereo-controlled synthesis of polyolefins with cycloalkane groups by late transition metal catalysts, see: D. Takeuchi, Polymer J., 2012, 44, 919.


Interest in polyolefins incorporating cyclic groups is a result of the improved transparency and high glass transition temperatures of such materials. See O. Henschke, F. Köller and M. Arnold, *Macromol. Rapid Commun.*, 1997, **18**, 617.


19 All reactions were typically conducted at a consigned temperature of 60 °C. However, with highly active systems such as 1, 2 and 3/MAO, and despite the low precatalyst loading and vigorous cooling system used, it often turned out difficult to evacuate the exothermicity of the polymerization; the temperature in the reactor hence often reached 70–75 °C (see Table 1). This increase of the reaction temperature accounts for some different productivities of a given system.

20 This study of MHD incorporation rate as a function of MHD loading could not be easily performed with 1/MAO, as this is system is so active that it exhausts rapidly MHD from the reaction medium.

21 Note yet that, as total conversion of MHD was achieved for this run with 1, it is likely that the amount of incorporated comonomer would have been higher if the polymerization was stopped before running out of MHD. However, due to technical constraints with this highly active system, this aspect was no further investigated.
The intensity ratio between the major (*) and minor (°) sets of resonances was determined to be 73:27 in P(P-co-MHD) copolymers produced by 2 (Table 1, entries 5 and 6).

Most of the signals in the region δ 30-45 ppm are also assignable to regio/stereoerrors in the polypropylene backbone.

Regular vinylidene groups in PP are observed at δ 111.6 and 144.5 ppm in $^{13}$C NMR and at δ 4.64 and 4.70 ppm in $^1$H NMR. The 0.06 ppm difference between the latter $^1$H signals is significantly larger than the 0.02 ppm difference observed in the P(P-co-MHD) copolymers produced by 4/MAO; see: L. Resconi, F. Piemontesi, I. Camurati, O. Sudmeijer, I. E. Nifant’ev, P. V. Ivvchenko and L .G. Kuz’mina, J. Am. Chem. Soc., 1998, 120, 2308.


Table 1. Copolymerization of propylene with MHD and model α-olefins by catalytic systems 1–5/MAO

<table>
<thead>
<tr>
<th>Ent.</th>
<th>Precat.</th>
<th>[Zr] [μmol.L⁻¹]</th>
<th>[Al]/[Zr]</th>
<th>Comon.</th>
<th>nCom. [mmol]</th>
<th>Tpolym [°C]</th>
<th>mpol [g]</th>
<th>Prod. [kgprop.gcat⁻¹.h⁻¹]</th>
<th>Mₚ/Mₜ</th>
<th>Tₗ [°C]</th>
<th>Tcrys [°C]</th>
<th>[mₚ] or</th>
<th>Comonom. incorp. [%]</th>
<th>Comonomer conv. [%]</th>
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<tr>
<td>1</td>
<td>1</td>
<td>2.9</td>
<td>15 510</td>
<td>-</td>
<td>0</td>
<td>60 (72)</td>
<td>21.6</td>
<td>541</td>
<td>41.0</td>
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<td>108</td>
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<tr>
<td>2</td>
<td>1</td>
<td>3.8</td>
<td>12 000</td>
<td>MHD</td>
<td>1.8</td>
<td>60 (75)</td>
<td>22.2</td>
<td>370</td>
<td>13.3</td>
<td>2.7</td>
<td>144</td>
<td>107</td>
<td>93.9</td>
<td>0.37/12</td>
</tr>
<tr>
<td>3</td>
<td>1</td>
<td>3.8</td>
<td>11 900</td>
<td>MHD</td>
<td>1.8</td>
<td>80 (96)</td>
<td>32.4</td>
<td>1 853</td>
<td>nd</td>
<td>nd</td>
<td>145</td>
<td>113</td>
<td>93.6</td>
<td>0.64/21</td>
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<tr>
<td>4</td>
<td>2</td>
<td>8.0</td>
<td>5 000</td>
<td>-</td>
<td>0</td>
<td>60 (72)</td>
<td>15.1</td>
<td>100</td>
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<td>105</td>
<td>94.1</td>
<td>-</td>
</tr>
<tr>
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<td>2</td>
<td>9.3</td>
<td>4 800</td>
<td>MHD</td>
<td>1.8</td>
<td>60 (72)</td>
<td>7.6</td>
<td>42</td>
<td>45.7</td>
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<td>105</td>
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</tr>
<tr>
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<td>MHD</td>
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<td>27</td>
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<td>106</td>
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<tr>
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<td>11.0</td>
<td>2 180</td>
<td>MHD</td>
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<td>21.6</td>
<td>100</td>
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<td>MHD</td>
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<td>60 (65)</td>
<td>2.5</td>
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<td>4</td>
<td>18.0</td>
<td>1 600</td>
<td>-</td>
<td>0</td>
<td>30 (45)</td>
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<td>131.5</td>
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<td>12.0</td>
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<td>1.4</td>
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<td>10.0</td>
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<td>2.5</td>
<td>145</td>
<td>103</td>
<td>91.2</td>
<td>0.30/10</td>
</tr>
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</table>

a Conditions: [Zr or Ti] = 3.8 to 11.0 μmol.L⁻¹, toluene (150 mL), in situ activation with 1,600–15,000 equiv. MAO. Ppropylene = 5 bar (i.e., [propylene] = ca. 1.7 mol.L⁻¹. Tpropylene = 60 °C, time = 10 min. b Reaction time = 20 min. c Reaction time = 30 min. d Values in brackets correspond to the maximal temperature reached in the reactor. e Melting (Tm) and crystallization (Tcrys) temperatures, as determined by DSC. f Determined by 1H (8-TMSO and 1.6-MOD) or 13C (MHD) NMR; see the Experimental section. g Not determined as a small part of this polymer was not soluble in TCB at 135 °C. h Not observed/not determined, gummy materials at room temperature.
Figure 4. Detail of the $^{13}$C-$^1$H NMR spectra (CDCl$_3$, 125 MHz, 135 °C) of a) an iPP homopolymer produced by 1/MAO (Table 1, entry 1). b) a P(P-co-MHD) copolymer produced by 1/MAO (Table 1, entry 2); c) a P(P-co-MHD) copolymer produced by 2/MAO (Table 1, entry 6); d) an iPP homopolymer produced by 2/MAO (Table 1, entry 4). Signals corresponding to MMCP motifs are noted * (major series) and ° (minor series).
Figure 5. Detail of the $^{13}\text{C}\{^1\text{H}\}$ NMR spectra (Cl$_2$CDCl$_2$, 125 MHz, 135 °C) of e) a P(P-co-MHD) copolymer produced by 3/MAO (Table 1, entry 10); signals corresponding to MMCP motifs are noted + (major series) and × (minor series) and f) a sPP homopolymer produced by 3/MAO (Table 1, entry 9).
Figure 7. Details of the $^{13}$C($^1$H) NMR spectrum (1,2,4-trichlorobenzene/$C_6D_6$ (5:1), 125 MHz, 135 °C) of a P(P-co-MHD) copolymer produced by 4/MAO (Table 1, entry 13)
For the Table of content entry

ansa-zirconocene cyclopolymerization

isotactic or syndiotactic P(P-co-MHD)
trans-pure or trans-enriched rings

vinyl copolymerization constrained geometry hemi-titanocene

atactic, regiorregular P(P-co-MHD)