Rhodium-Mediated Stereoselective Polymerization of “Carbenes”

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Abstract: Unprecedented rhodium-catalyzed stereoselective polymerization of “carbenes” from ethyl diazoacetate (EDA) to give high molecular mass poly(ethyl 2-ylidene-acetate) is described. The mononuclear, neutral [(N,O-ligand)M(ED)2] (M = Rh, Ir) catalytic precursors for this reaction are characterized by (among others) single-crystal X-ray diffraction. These species mediate formation of a new type of polymers from EDA: carbon-chain polymers functionalized with a polar substituent at each carbon of the polymer backbone. The polymers are obtained as white powders with surprisingly sharp NMR resonances. Solution and solid state NMR data for these new polymers reveal a highly stereoregular polymer, with a high degree of crystallinity. The polymer is likely syndiotactic. Material properties are very different from those of atactic poly(diethyl fumarate) polymer obtained by radical polymerization of diethyl fumarate. Other diazoacetates are also polymerized. Further studies are underway to reveal possible applications of these new materials.

Introduction

The thermal decomposition of diazomethane (reported around 1900) is the first known route to polymethylene.1 Neither this explosive process, however, nor the milder catalytic versions2 ever found large scale practical applications,3 and it was more than half a century later that transition metal mediated ethene polymerization gave easy access to linear polyethylene.4,5 Unlike the inherently unstable diazoalkanes, diazocarbonyl compounds such as diazoacetates (N2CHCO2R) are reasonably stable, safe (even in large scale/industrial synthesis),6 easy to prepare, and extensively used as carbene precursors in organic synthesis. The most frequently observed reactions include carbene dimerization to form olefins, carbene transfer to olefins in cyclopropanation reactions, and carbene insertion into O–H, N–H, and C–H bonds. The reactions are often mediated by transition metals such as platinum, copper, rhodium acetates, and ruthenium catalysts, which have been reported to give efficient and, in many cases, stereoselective conversion of the carbenido to more or less sophisticated products.7 In this framework, though, only three reports describe the polymerization (oligomerization) of α-carbonyl stabilized “carbenes” from diazocarbonyl compounds. This concerns Cu- and Rh-mediated formation of low molecular mass polymers (oligomers) from alkyl diazoacetates and related diazocarbonyls (number average degree of polymerization up to ca. 100).8,9 These materials are notable because they carry a polar functionality (a) Clack, J. D.; Shah, A. S.; Peterson, J. C. Thermochim. Acta 2002, 392–392, 177–186. (b) Clack, J. D.; Shah, A. S.; Peterson, J. C.; Patelis, L.; Kersten, R. J. A.; Heemskerk, A. H. Thermochim. Acta 2002, 386, 65–72. (d) Clack, J. D.; Heise, J. D.; Shah, A. S.; Peterson, J. C.; Chou, S. K.; Levine, J.; Karakas, A. M.; Ma, Y.; Ng, K.-Y.; Patelis, L.; Springer, J. R.; Stano, D. R.; Wettach, R. R.; Dutra, G. A. Org. Process Res. Dev. 2004, 8, 176–185.


The variations in the M-
This effect is more pronounced for the one of the two vacant sites somewhat from its environment.

Rh-Mediated Polymerization of “Carbenes”

Results and Discussion

Synthesis and Characterization of the Catalysts. The [M\(^{III}\)(L-prolinate)(cod)] complexes 1a (M = Rh) and 1b (M = Ir) (cod = ZZ-L,5-cyclooctadiene) were prepared by reaction of the [(cod)M\(^{II}\)(Cl)] \(_2\) precursors with in situ deprotonated L-proline in methanol at room temperature (Scheme 1). The benzyl analogues [M\(^{III}\)(N-benzyl-L-prolinate)(cod)] (2a, M = Rh; 2b, M = Ir) and the nonchiral picolinate and quinolinolate Rh analogues 3 and 4 were prepared via similar routes. The structures of 1–3 were determined by single-crystal X-ray diffraction (Figure 1; Table 1). The reported structure of 4 is very similar.\(^{11}\)

Complexes 1–4 all adopt a square-planar geometry. The picolinate ligand in 3 is planar and causes little steric hindrance to the two vacant sites of the Rh center. The aliphatic prolinate ring of 1a on the other hand is bent, thus shielding one of the two vacant sites somewhat from its environment. This effect is more pronounced for the N-benzyl-L-prolinate analogue 2a. The Ir species 2b is isostructural to its Rh analogue 2a. The M–ligand bond lengths of 1–4 are quite comparable. The variations in the M–N bond lengths in the order N\(_{Ir}\) < NHR\(_2\) < NR\(_3\) are as expected for sp\(^2\) and secondary and tertiary sp\(^3\) hybridized nitrogen atoms.

Polymerization of Carbenes from Diazoacetates. Doyle-type dinuclear tetracarboxylate-bridged Rh\(^{III}\)–Rh\(^{III}\) species have been extensively studied as catalysts for a variety of carbene transfer reactions using diazoacetates as carbene precursors.\(^{12}\)

| Table 1. Selected Bond Lengths (Å) and Angles (deg) Observed for Complexes 1–3 |
|-----------------|-----------------|-----------------|-----------------|
| param           | 1a (M = Rh)     | 2a (M = Rh)     | 2b (M = Ir)     | 3 (M = Rh)      |
| O1-M-O1         | 2.042(2)        | 2.0593(11)      | 2.0548(15)      | 2.0597(13)      |
| M-N             | 2.109(2)        | 2.1558(14)      | 2.152(2)        | 2.0857(16)      |
| M-C1            | 2.095(3)        | 2.0986(15)      | 2.092(2)        | 2.1028(19)      |
| M-C2            | 2.088(3)        | 2.1155(15)      | 2.108(2)        | 2.084(2)        |
| M-C5            | 2.124(3)        | 2.1049(17)      | 2.099(2)        | 2.1370(19)      |
| M-C6            | 2.115(3)        | 2.1432(17)      | 2.127(3)        | 2.127(2)        |
| O1-M-N          | 81.88(8)        | 79.61(5)        | 79.40(7)        | 80.00(6)        |
| O1-M-C1         | 163.61(18)      | 155.49(6)       | 155.74(8)       | 164.13(7)       |
| N-M-C1          | 96.58(12)       | 98.84(7)        | 100.00(9)       | 100.45(7)       |

Scheme 1. Synthesis of Catalysts 1a,b and the Structure of 2–4

Figure 1. X-ray structures of 1a, 2a, and 3.

(12) See for example: Davies, H. M. L.; Beckwith, R. E. J. Chem. Res. 2003,
103, 2861 and references therein.
(13) Krumper, J. R.; Gerisch, M.; Suh, J. M.; Bergman, R. G.; Tilley, T. D. J.

J. AM. CHEM. SOC. VOL. 128, NO. 30, 2006 9747
mass polymers were obtained (observed with PEA as a white powder. The highest yields in polymer were from the reaction mixture by quenching with methanol, yielding on the reaction conditions (Table 2). The polymer was separated gave PEA in isolated yields between 10% and 50% depending At 20°C calibrated with matrix-assisted laser light scattering (MALLS). Solvents (e.g., THF, acetonitrile, or toluene) yields were lower, using the catalysts (PEA, Scheme 2). Formation of PEA was further investigated fumarate according to NMR. The remaining material (EDA, this yields at most 40% diethyl maleate and diethyl dimerization yielding diethyl maleate and diethyl fumarate in a 1:2 ratio. However, while 1a mediates full conversion of EDA, this yields at most 40% diethyl maleate and diethyl fumarate according to NMR. The remaining material (~60%) proved to be the new poly(ethyl 2-yldiene) polymer (PEA, Scheme 2). Formation of PEA was further investigated using the catalysts 1–4.

Treating EDA with a catalytic amount of 1 (2 mol %) gave PEA in isolated yields between 10% and 50% depending on the reaction conditions (Table 2). The polymer was separated from the reaction mixture by quenching with methanol, yielding PEA as a white powder. The highest yields in polymer were observed with 1a in chloroform or dichloromethane. In other solvents (e.g., THF, acetone, or toluene) yields were lower, possibly due to partial catalyst precipitation.

Molecular masses of the new PEA polymers were determined from size exclusion chromatography (SEC) measurements calibrated with matrix-assisted laser light scattering (MALLS). At 20°C, with all four rhodium-based catalysts high molecular mass polymers were obtained (M_w typically in the range 120–155 kDa) within 14 h. Not unexpectedly, shortening the reaction time leads to formation of lower weight polymers in lower yields (Table 2, entry 4). Decreasing the reaction temperature and increasing the reaction time leads to an increase of the polymer molecular mass (M_w up to 190 kDa for polymerization with 1a in chloroform at ~20°C).

The polydispersities of the polymers obtained with the Rh catalysts are all slightly higher than 2.0. The polymerization experiments were run with full conversion of EDA, which contributes to somewhat broader weight distributions (Table 2, entries 3 and 4). Therefore, we consider the obtained polydispersity to be in agreement with a nonliving polymerization process at a single center active species.

Quite remarkably, 1b (the iridium analogue of 1a) does not produce any PEA polymer from EDA. Instead, 1b catalyzes quantitative carbene dimerization to yield a mixture of diethyl fumarate and diethyl maleate (in a ratio of 1:9). Iridium complex 2b on the other hand, containing a weaker coordinating benzyl-functionalized N-donor, does produce PEA in low yields (4%). Both the molecular weight (M_w = 12 kDa) and the polydispersity (M_w/M_n = 1.6) of the polymer obtained with 2b are much lower than those of the polymers obtained with the Rh species.

A variety of diazocarbonyl “monomers” are readily obtained via cheap and easy routes. Addition of nucleophiles to dioxenones, followed by diazo transfer to the resulting β-keto esters and base-assisted elimination of acetic acid, provides a convenient route to such species (Scheme 3).7,14 Preliminary results reveal that treatment of thus obtained crude 3-butenyl-diazoacetate (Nu = OCH2CH2CH=CH2) and 3-butynylidiazooacetate (Nu = OCH2CH2C≡CH) with 1a yields the corresponding polymers, albeit in low yields (2–8%). Copolymers from 3-butynylidiazooacetate and EDA (1:10) are obtained in higher yields (32%). Polymerization of n-butyldiazooacetate (Nu = OBu) gives a yield of ~20%. Clearly, the scope of the catalytic process introduced herein is not limited to EDA, and we are currently investigating other diazocarbonyl compounds as carbene precursors in Rh-mediated polymerization reactions.

Characterization of Stereoregular PEA. Quite surprisingly, the new high molecular weight PEA polymers obtained with catalysts 1–4 reveal sharp resonances in solution 1H and 13C NMR spectra indicative for stereoregular polymers. The low molecular weight (M_w ~ 1.2 kDa) PEA polymers (oligomers) obtained with Pd-based catalysts reported by Ibarra and coworkers are viscous oils which show very broad NMR resonances.9a Apart from the possibility that these polymers are atactic, the broadening of the NMR signals of these low molecular weight polymers could also be caused by chain-end effects. For none of the PEA samples obtained with the Rh and Ir catalysts 1–4 we were able to detect polymer chain-end signals or indirect chain-end effects with NMR. To investigate tacticity effects, we prepared atactic PEA by radical polymerization (initiated by AIBN) of diethyl fumarate.9a,10 These atactic PEA samples have a larger molecular weight (M_w = 6.1 kDa) but still reveal very broad NMR resonances. NMR spectra of the new stereoregular PEA polymers obtained with catalysts 1–4 are clearly different from those of atactic PEA (Figure 2).

Even at 600 MHz for 1H NMR and 150 MHz for 13C NMR, the main chain methine signal of stereoregular PEA appears as a highly symmetrical singlet, with a half-height width of 11 and 20 Hz, respectively. A weak shoulder at higher frequency

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Scheme 2. Stereoregular Polymerization of Ethyl Diazoacetate

Scheme 3. Preparation of Diazocarbonyl Compounds from Dioxenones

**Table 2. Polymerization of EDA with Catalysts 1–4**

<table>
<thead>
<tr>
<th>entry</th>
<th>catalyst</th>
<th>T, °C</th>
<th>solvent</th>
<th>reaction time</th>
<th>yield, %</th>
<th>M_w, kDa</th>
<th>M_w/M_n</th>
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<tr>
<td>1</td>
<td>1a</td>
<td>−20</td>
<td>CHCl₃</td>
<td>7 days</td>
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<td>2</td>
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<td>35</td>
<td>169</td>
<td>3.8</td>
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<tr>
<td>3</td>
<td>1a</td>
<td>20</td>
<td>CHCl₃</td>
<td>14 h</td>
<td>45</td>
<td>133</td>
<td>2.4</td>
</tr>
<tr>
<td>4</td>
<td>1a</td>
<td>20</td>
<td>CHCl₃</td>
<td>30 min</td>
<td>5</td>
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</tr>
<tr>
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<td>14 h</td>
<td>30</td>
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<tr>
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<td>CH₂Cl₂</td>
<td>14 h</td>
<td>40</td>
<td>143</td>
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<td>1a</td>
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<td>THF</td>
<td>14 h</td>
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<td>14 h</td>
<td>25</td>
<td>120</td>
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<td>20</td>
<td>CHCl₃</td>
<td>14 h</td>
<td>25</td>
<td>154</td>
<td>2.7</td>
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<tr>
<td>11</td>
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<td>14 h</td>
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<tr>
<td>12</td>
<td>2b (M = Ir)</td>
<td>20</td>
<td>CHCl₃</td>
<td>14 h</td>
<td>4</td>
<td>12</td>
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<tr>
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<td>14 h</td>
<td>15</td>
<td>147</td>
<td>2.9</td>
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</tbody>
</table>

* Conditions: 0.037 mmol of catalysts; 1.84 mmol of EDA, 6 mL of solvent. After precipitation with MeOH.

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of the $^1$H NMR signal is possibly traceable to stereo defects (in all cases well below 10%, as estimated by peak deconvolution).

Solid-state NMR is consistent with the above conclusion and provides more information. In the CP/MAS NMR spectrum \(^{13}\)C (Figure 3), the methine signal reveals two sharp resonances ($\delta$ = 43.73 and 45.30 ppm) in an approximate 1:1 ratio, partly overlapped by a weaker and broader peak at higher frequency ($\delta$ = 47.8 ppm). The former can be attributed to a crystalline phase, while the latter represents an amorphous fraction. Fourier transformation with a resolution enhancing window function revealed a similar splitting for the carbonyl peak (1:1 sharp doublet at $\delta$ = 171.0 and 171.8 ppm) and more complicated fine structures for the methylene ($\delta$ = 59.6–63.7 ppm) and methyl ($\delta$ = 12.8–14.6 ppm) peaks (Supporting Information, Figure S6 and S7).

By deconvolution of the methine resonance in a quantitative single pulse $^{13}$C MAS NMR spectrum (recycle time, 60 s), the relative abundance of the crystalline phase was estimated to be slightly above 70%. In a single pulse $^{13}$C MAS NMR spectrum run with very short recycle time (1 s), so as to minimize the contribution of more rigid, ordered structures, which generally tend to have a longer $T_1$ relaxation time, \(^{16}\) the methylene and methyl resonances turned out to be clearly stronger than the methine and carbonyl ones, which suggests some conformational disorder of the ethyl residue even in the crystalline domains. Powder X-ray diffraction spectra of stereoregular PEA reveal strong but relatively broad peaks (Supporting Information, Figure S9). This seems in good agreement with a high degree of crystallinity but with substantial disorder of the ethyl residues.

Stereoregular PEA can in principle be isotactic or syndiotactic. Preliminary molecular mechanics (MM2) calculations on model compounds indicated that low-energy conformations with a two-backbone-bond conformational repeat are found for both polymers in the TG (trans-gauche) and GT domains; this is consistent with a shift to lower frequency of the methine peak for the ordered phase relative to the disordered one in the CP/MAS NMR spectrum. The $\text{H}^\text{--C}^\text{--C}^\text{--O}$ torsions are all close to anti (anti $\approx 160^\circ$) in the isotactic polymer, whereas their sequence in the syndiotactic polymer is ...syn, syn, anti, anti, syn, syn... (syn $\approx 30^\circ$, anti $\approx 170^\circ$), with the syn, anti or anti, syn arrangements flanking the main chain T bonds. The SCF-MO calculated NMR spectra are compatible with the experimental one for both structures; therefore additional information is needed for a definitive assignment of the configuration. We recently prepared carbonyl $^{13}$C-labeled samples of the new stereoregular PEA polymers, and we are currently investigating their configuration by 2-dimensional carbon–carbon correlation experiments (DOQSY). \(^{17}\) Analyzing the results of these experiments is not straightforward and requires elaborate data-fitting. \(^{18}\) These results will be published in a subsequent paper.

**a) Thermal Analysis.** Stereoregular PEA as obtained by precipitation from the reaction mixture with MeOH is largely amorphous. However, slow evaporation of the solvent from a solution of stereoregular PEA in chloroform results in a “crystallized” brittle film of the polymer. These “amorphous” and crystallized samples of stereoregular PEA have different properties.

Thermogravimetric measurements (TGA) reveal that both the amorphous and crystallized PEA samples decompose above 300 °C, but DSC data for these samples at lower temperatures are clearly different from each other. The amorphous samples showed a reversible first-order phase transition at 100 °C (75 °C on heating/cooling (Supporting Information, Figure S11). In the first heating/cooling cycle these transitions are very weak but become stronger ($\Delta h = 21$ J g$^{-1}$) in a second cycle after annealing the polymer for a few seconds at 240 °C.

Samples which were first crystallized from CHC1$_3$ reveal quite pronounced ($\Delta h = 24$ J/g) and sharp DSC transitions from the beginning, without the need to anneal the polymer (see Figure...
The transitions of the crystallized samples also occur at higher temperatures (150 °C/110 °C) compared to those of the amorphous samples (100 °C/75 °C). Since the polymer is a flexible rubber above 155 °C, whereas it is hard and brittle below 70 °C, the transitions at 150 °C/110 °C for crystallized PEA and 100 °C/75 °C for amorphous PEA could be interpreted as the softening and glass-transition temperatures, in accord with reported glass–rubber transitions of atactic PEA (between 65 and 80 °C). The fact that crystallized samples of stereoregular PEA reveal transitions at higher temperatures compared to the amorphous samples and the fact that these transitions appear as real peaks in the DSC could point to a substantial influence of the glass–rubber transitions by the crystalline domains of this material. However, at this point we cannot exclude an alternative interpretation of the data, in which the DSC peaks correspond to crystalline–liquid crystalline transitions instead of glass–rubber transitions. We are currently investigating this possibility.

The reported softening (78–80 °C) and glass transition temperatures (65 °C) of atactic PEA (obtained by radical polymerization of diethyl maleate with AIBN) are somewhat lower than the above transitions of our amorphous stereoregular PEA. For bulky polymers, containing only one repeating side group/two carbons of the polymer backbone, the highest glass transition temperatures are usually observed for the atactic and syndiotactic polymers. For poly[methyl(methacrylate)] (PMMA) for example, quite large differences in glass transition temperatures are reported for the atactic (∼90 kDa, $T_g = 105 \degree C$), isotactic (30 kDa, $T_g = 56 \degree C$), and syndiotactic (60 kDa, $T_g = 140 \degree C$) polymers. Van der Waals forces between polymeric chains in amorphous domains are stronger for the less symmetrical atactic and syndiotactic polymers as compared to isotactic polymers. If one reasons in analogy (and assumes that the transitions belong to glass–rubber transitions), stereoregular PEA might be syndiotactic because even the amorphous samples reveal transitions at higher temperatures than atactic PEA. However, since we have no reference material for glass–rubber transitions of atactic, isotactic, and syndiotactic carbon-chain polymers which are substituted at each carbon of the polymer backbone (nor information excluding the alternative interpretation of the transitions as crystalline–liquid crystalline transitions), these data are only an indication for the possible configuration of stereoregular PEA.

Under a polarization microscope, the crystallized PEA samples show birefringence. Above the softening temperature, also the amorphous PEA samples start to show birefringence but only after physical contact. We take the birefringence as a further indication for the semicrystalline behavior of stereoregular PEA (or perhaps thermotropic liquid-crystalline behavior in the rubber state).

(b) MALDI-TOF. The polymers prepared with Rh catalysts 1a, 2a, 3, and 4 invariably give high molecular weight PEA polymers with rather broad molecular weight distributions ($M_w / M_n = 2–3$; see Table 2). As expected, we were unable to measure any reliable MALDI-TOF mass spectra from these PEA samples. However, MALDI-TOF spectra of PEA prepared with Ir catalyst 2b ($M_w = 12$ kDa, $M_w / M_n = 1.6$) were readily obtained, albeit only in the lower mass region of the polymer. As shown in Figure 5, PEA is clearly built from ethyl 2-ylideneacetate ($:CHCOOEt$) repeating units, resulting in a regular 86 Da spaced repeating pattern. Fragments with an equal number degree of polymerization ($n$) give rise to three masses: ($CHCOOEt)_n + 25 + x \times 86$ Da (dominant signals), ($CHCOOEt)_n + 41 + x \times 86$ Da (weaker), and ($CHCOOEt)_n + 56 + x \times 86$ Da (least intense).
Although the masses of the more intense signals correspond with $[\text{Na}^+]_n([\text{H–CHCOOEt})_n])$ (e.g. $m/z = 3467$ Da, with $n = 36$) and $[\text{Na}^+]_n([\text{H–(CHCOOEt)}_n–\text{OH}])$ (e.g. $m/z = 3483$ Da, with $n = 36$), these interpretations do not explain their typical isotope distribution patterns. These patterns are suggestive for the presence of cationic iridium complexes as charge carriers (Figure 6). The signals could well belong to $[\text{Ir}^{III}(\text{OH})_2(\text{cod})(\text{MeOH})]^{+}_n([\text{H–(CHCOOEt)}_n])$ (e.g. $m/z = 3467$ Da, with $n = 36$) and $[\text{Ir}^{III}(\text{OH})_2(\text{cod})(\text{MeOH})]^{+}_n([\text{H–(CHCOOEt)}_n–\text{OH}])$ (e.g. $m/z = 3483$ Da, with $n = 36$), but with different Ir-bound ligand interpretations are possible. Addition of K$^+$ salts does not influence the spectrum. Addition of Ag$^+$ salts does not change the peak positions but does influence their relative intensities, leading to significantly increased intensities of the CHCOOEt$^+_n$ + 41 + $x$ × 86 Da signals (e.g. 3483 Da). Also the isotopic distribution patterns appear to be different in the presence of Ag$^+$. The coincidentally unchanged masses with slightly different isotopic distribution patterns are suggestive for the presence of $[\text{Ag}^+]_n([\text{H–CHCOOEt})_n])$ and $[\text{Ag}^+]_n([\text{H–(CHCOOEt)}_n–\text{OH}])$. Considering the increased intensity of the CHCOOEt$^+_n$ + 41 + $x$ × 86 Da signals in the presence of Ag$^+$, it seems quite safe to conclude that H–(CHCOOEt)$_n$–OH chains are present in the polymer samples. The MS data are also suggestive for the copresence of H–(CHCOOEt)$_n$–H chains, although this picture is somewhat blurred (no clear intensity change upon addition of Ag$^+$ and overlapping signals of Ir-based alternatives).}

At this point we cannot entirely exclude the possibility of organometallic rhodium or iridium complexes being present as polymer chain ends. PEA samples obtained with catalyst 1a reveal the presence of 0.09% Rh, which corresponds to approximately 1 equiv of Rh/polymer chain. Therefore, a final assignment of the polymer end groups (i.e. Rh/Ir based, only H–PEA–OH or a mixture of both H–PEA–OH and H–PEA–H) requires additional data from high-resolution MS spectra (requiring polymer samples with a lower polydispersity) and/or NMR characterization of lower molecular weight polymers.

**Mechanistic Considerations.** At this point we do not understand all details of the formation of PEA. We need a high catalyst loading of 2% to obtain complete conversion of EDA. Half of the starting material converts to diethyl maleate and diethyl fumarate. Since polymers consisting of about 1000 CHCOOEt repeating units are formed, only about 3% of the added catalytic precursor becomes really active as a polymerization catalyst. The exact nature of the active species is therefore unknown, and in part we necessarily rely on (educated) speculations in the mechanistic considerations below.

The nature of the catalyst has little influence on the obtained stereoregular PEA. The Rh-based catalytic precursors 1a, 2a, 3, and 4 produce roughly the same molecular weight PEA with comparable polydispersities. Only the Ir-based precursor 2b produces somewhat lower molecular weight PEA. There are no obvious differences in the properties between the obtained polymers (including NMR data), suggesting that identical or at least very similar active species are generated. This likely involves dissociation of the NO-ligand from the metal under catalytic conditions. However, the precursors 1a, 2a, 3, and 4 produce PEA in different yields, thus suggesting that the NO-ligands do have an influence on the kinetics of initiation. This might, for example, involve formation of a reactive metal–alkyl by attack of a nitrogen donor (as in the Pd-based systems of Ibara et al.) or oxygen donor of the NO-ligand to a metal-bound carbene. The MALDI-TOF spectra further suggest the presence of –H and –OH end groups. Chain growth might thus start from M–OH and/or M–H species, while termination might involve reductive elimination of M(H)(C$_{\text{chain}}$) fragments, protonation of M–C$_{\text{chain}}$ fragments, and/or oxidation (by e.g. O$_2$).

Formation of a rhodium-bound carbene is commonly believed to be a key step in rhodium-catalyzed carbene transfer reactions.
with diazoacetates. PEA might thus be formed via consecutive (migratory) \( \alpha \)-insertions of metal-bound carbenes into the Rh–C bond of the growing chain (Scheme 4).

(a) Stereocontrol. Since no ancillary ligand effect on the stereoselectivity was observed, we propose that the insertions into the Rh–CH(COOEt)P fragment (P = polymeryl) are sterically controlled by the configuration of the Rh-bound carbon atom of the growing chain (propagation under “chain-end control”). This is illustrated by the Newman projections in Figure 7. Stereoselective propagation under chain-end control has been observed in a number of insertion polymerizations, including the Pd-mediated alternating copolymerization of CO with styrene.

Migratory carbene insertions involving stereogenic Rh–CH(COOEt)P methine carbons in configuration S experience least steric hindrance when the re-face of the rhodium carbene is attacked. This yields a new Rh–CH(COOEt)P methine carbon in configuration R, which in turn preferentially attacks the si-face of a carbene (Figure 7). We thus expect formation of a syndiotactic polymer, having a regular arrangement of alternating R and S configurations concerning the stereogenic carbons of the polymer backbone. This seems in agreement with the DSC data, but a definitive assignment of the polymer configuration still requires additional investigations.

Conclusions

Rh-mediated polymerization of carbenes, generated from ethyl diazoacetate, results in formation of a new highly stereoregular polymer with a high molecular mass: poly(ethyl 2-yldiene-acetate) (PEA). Diazoacetates containing other ester functionalities can also be polymerized. The polymers contain polar functional side groups at each carbon of the carbon-chain backbone.

PEA is obtained as a white amorphous solid but can be crystallized to obtain a material with a high degree of crystallinity (~70%). The stereocontrol in the formation of PEA most likely involves propagation under chain-end control. Therefore, stereoregular PEA is likely syndiotactic, which seems in agreement with the relatively high \( T_g \) of the material.

We expect that these new polymers, having a unique structure, will have special material properties. Further studies, in which we focus of the structure and properties of these new materials, are underway.

Acknowledgment. This work was supported by The Netherlands Organization for Scientific Research (NWO-CW), the Radboud University of Nijmegen, and the University of Amsterdam. We thank J. Opsteen, R. Bovee, and Dr. A. Kros for their assistance with the SEC/MALLS measurements. We thank P. J. Aarnoutse for her assistance with the MALDI-TOF measurements.

Supporting Information Available: Experimental procedures and crystallographic data for 1a, 2a,b, and 3, details on SEC/MALLS, NMR, X-ray diffraction, and DSC/TGA polymer characterization, and MM2 and SCF-MO NMR property calculations. This material is available free of charge via the Internet at http://pubs.acs.org.

JA058722J


