REVEIW PAPER

Discovery and development of metallocene-based polyolefins with special properties

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ABSTRACT

Beside Ziegler-Natta and Phillips catalysts the development of methylaluminoxane (MAO) as cocatalyst in combination with metallocenes or other transition metal complexes for the polymerization of olefins has wid ly in reased the possibilities in cnot by the polymer cnown possibilities in the polymerization of olefins has wid ly in reased the possibilities in cnot by the polymer cnown possibilities are catalysts allow the synthesis of isotactic, isoblock, syndiotactic, stereoblock or atactic polymers, as well as polyolefin composite materials with superior properties such as film clarity, tensile strength and lower content of extractables. Metallocene and other single site catalysts are able to copolymerize ethen and polymer new with short and logical chain documents, cyclic olefins, or polar vinyl monomers such as ethers, alcohols or esters, especially, if the polar monomers are protected by aluminum alkyls. Different vinyl ethers such as vinyl-ethyl ether, vinyl-propyl ether, vinyl-hexyl ether, and 2,7-octadienyl methyl ether (MODE) were copolymerized with olefins using triisobutyl aluminum as protecting agents. Polar monomers could be incorporated into the polymer chain by up to 16 mol%. Such copolymers show better gas barrier and surface properties, as well as solvent resistance and they are suitable for blends of polyolefins with polyethers and other polar polymers because of an excellent adhesion of the two polymers. Polyolefins J (2015) 2: 1-16

 $\textbf{Keywords:} \ methylaluminoxane; \ metallocene \ catalysts; \ olefin \ copolymerization; \ polar \ monomers; \ vinyl \ ethers$

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INTRODUCTION

In comparison to Ziegler-Natta systems, metallocene catalysts are soluble in hydrocarbons; show only one to 6 active site and their chemical structure can be easily changed. These qualities would allow to predict the properties of the resulting polyolefins accurately by ing the structure of the metallocenes are edding their structure of the structure of the metallocenes and to control the resulting mobile education of the specific of the specifi

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conditions. In addition, their catalytic activity is about 100 times higher than the classical Ziegler-Natta systems. Metallocenes in combination with conventional alm im alk cocataly ts, are id ed cap be 6 polymerizing ethene, but only at a very low activity. With the id scor ry ad ap ication 6 meth alm inoxane (MAO) it was possible to enhance the activity, surprisingly, by a factor of 10,000. Therefore, MAO p ay a cru ial p rt in the cataly is with metallo en s [1].

The discovery of MAO has been more or less an accident. In the Institute of Technical and Macromolecular Chemistry at the University of Hamburg we have been in stig tig no sid reactino, especially the hydrogen transfer in homogeneous Ziegler-Natta systems [2]. To lower the reduction rate of the titain m system, we have been in stig tig no the reactinob tween b scylpon aid elastician mid method at trimeth alm im, in tead 6 a titain mid ethod at trieth lim im containing system main y by NMR analysis at low temperatures. The formation of new CH₂-bigled titain mid alm im combers is been been solved.

$$\text{Cp}_2\text{Ti}(\text{CH}_3)_2 + \text{Al}(\text{CH}_3)_3 \rightarrow \text{Cp}_2(\text{CH}_3)\text{Ti-CH}_2\text{-Al}(\text{CH}_3)_2 + \text{CH}_4$$

This slow reaction and the complexation by ethene has been analyzed by NMR measurements, that there is no reduction of titanium (IV) and no polymerization take place. In an incident in 1975 the unintentional condensation of water into the NMR tube led to the astonishing formation of polyethylene [3]. This unexpected observation confirmed in a larger scale experiment in a 1 L autoclave [4]. More details of this discovery can be found in published reports [5, 6]. The polymerization rate in a halogen free Cp, Ti(CH₂)₂/ Al(CH₃)₃ /H₂O-catalyst reached a maximum, when a high amount of water with its maximum ratio of 1:1 with trimethylaluminum was added. It was clear that trimethyaluminum (TMA) reacted rapidly with water in toluene. The next step was therefore to isolate the product, formed in a 1:1 mixture of water and trimethylaluminum in solvents, in order to avoid explosions. The general reaction followed as in Equation 2.

$$(2) \\ nH_2O + (n+1)Al(CH_3)_3 \rightarrow (H_3C)_2Al-[O-AlCH_3]_n - CH_3 + 2n CH_4$$

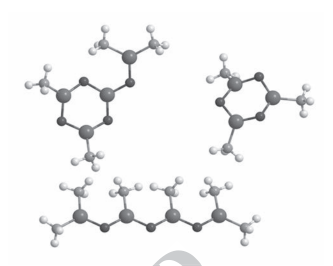


Figure 1. Structures of cyclic and linear MAO, big balls: aluminium, small balls: oxygen, and methyl groups

After 20 hours, the reaction mixture was filtrated and the solvent evaporated. The white powder obtained was dried and analyzed. The compound was named methylaluminoxane. MAO was investigated by elementary analysis, cryoscopic and NMR measurements, and decomposition with HCl. It was found that MAO was a mixture of different oligomers, including some ring structures (Figure 1).

Ew n to y the ex ct strutne is to by n b can e the re are eq lib is with the bign ers, cm - be ex tin 6 the bign ers toward each to her and the unreacted TMA. MAO is a compound in which alm im and so n atm s are arranged alternately and free valences are saturated by methyl substituents. Sinn [7], Eilertsen, Rytter, and Ystenes [8] and others calculated a ratio CH₃/Al = 1.5 for MAO.

As the aluminum atoms in the unit structures of MAO are coordinatively unsaturated, the units join together forming clusters and cages. These have molecular weights from 1200 to 1600, measured by cryoscopy in benzene and are soluble in hydrocarbons especially in aromatic solvents. A probable association and cage like structure of four [Al₄O₃Me₆] **n** its are shown in Figure 2.

The nature of the polymerization active site of metallocene/MAO catalysts and the role of MAO are not fully understood. One function of MAO is the ally atin 6 th metallo en cm p ex in case th t a dichloride is used. The other is the formation of an ion-pair. Marks [9], and Bochmann [10], showed that the activity 6 metallo en cataly ts d p d n the formation of cationic species. Today, most research groups agree with this statement. The bulky MAO

$$\begin{array}{c} H_3C & CH_3 \\ Al-O-Al-O-Al-O-Al \\ H_3C & CH_3 \end{array}$$

$$(a) & CH_3 \\ CH_3 & CH_3 \end{array}$$

$$(a) & Al \\ O & Al \\ Al & O \\ Al & Al \\ O & Al \\ Al & O \\ Al &$$

Figure 2. a: Unit structure of MAO, b: cage formed by four linear unit structures with four and six- membered rings

cluster takes a chlorine -atom or a methor-g p from the metallorene together with an electron forming a cationic metallocene and an anionic MAO complex (Equation 3)

$$Cp_2Zr(CH_3)Cl + MAO \rightarrow [Cp_2Zr(CH_3)]^+ + [MAO-Cl]^-$$
 (3)

Essential to polymerization activity would be the fo matin 6 b lk an a with a weak the ng to the metallocene cations. Knowing that the bulky structure of MAO could be necessary for the activation of metallo en cataly ts, the bolk and weak y coolid a tige cocatalysts such as tris(pentafluorophenyl) borate or organic salts of the tetrakis (pentafluorophenyl) borate $[(C_6F_5)4B]^-$, and aluminum fluorides were introduced [9, 10]. With these cocatalysts a metallocene/cocatalysts ratio of 1:1 was used but only when a high excess of an aluminum alkyl as scavenger was present.

The polyolefin industries have used MAO containing catalysts in a large scale. Currently, companies such as Albermale, Akzo, Chemtura, and Mitsui produce hundreds of tons of MAO by reaction of water of ice with trimetly alm im and in some cases and other aluminum alkyls to increase their solubility.

Homo-polyolefins

Metallocenes, especially zirconocenes and also titano-

cenes, hafnocenes, and MAO treated transition metal complexes are highly active for the polymerization of olefins, diolefins, and styrene and have motivated wo ldv id research g p to the and 6 p tens and publications in the last 20 years. An overview can in selected reiv ew articles and b 20]. It could be shown that a soluble catalyst, such as Cp₂ZrCl₂/MAO is able to produce polyethylene with h h mb ech ar weih s and a n rrw mb ech ar weight distribution of approximately two. All active sites are similar ad form by n ers with the same average chain length (single site catalysts). Only traces of low molecular weight oligomers are formed. Ziren s with it fferen syn metries and sh titutions are shown in Figure 3. There is a great variety of th stru tu e 6 metallo en s, which can b u ed fo the polymerization. The cyclopentadienyl-, indenyl-, fluorenyl- ligands could be hydrated or substituted by alkyl, aryl, methoxy, siloxy or other groups. Ethanediyl (C₂H₄), dimethylsilandiyl ((CH₂)₂Si), or isopropandiyl ((CH₂)₂C) are mainly used as interannular bridges between the rings [21]. Central metals could be Ti, Zr, Hf. Such and similar metallocenes are used for the polymerization of ethene, propene, and other olefins [22].

Table 1. Comparison of ethene polymerization ^(a) with different metallocene/methylaluminoxane catalysts at same polymerization conditions

Metallocene(b) (Met)	Activity [kg PE/	Molecular
wetanocene. (wet)	(mol Met×h×c _E)]	weight (g/mol)
Cp ₂ TiCl ₂	34200	400000
Cp ₂ ZrCl ₂	60900	620000
Cp ₂ HfCl ₂	4200	700000
Cp* ₂ ZrCl ₂	1300	1500000
$En(IndH_4)_2]ZrCl_2$	22200	1000000
[En(Ind) ₂]ZrCl ₂	12000	350000
[En(Ind) ₂]HfCl ₂	2900	480000
$[En(2,4,7-Me_3Ind)_2]ZrCl_2$	78000	190000
[Me ₂ Si(Ind) ₂]ZrCl ₂	36900	260000
[Ph ₂ Si(Ind) ₂]ZrCl ₂	20200	320000
[Me ₂ Si(2,4,7-Me ₃ Ind) ₂]ZrCl ₂	111900	250000
[Me ₂ Si(2Me-4Ph-Ind) ₂]ZrCl ₂	16600	730000
[Me ₂ C(Ind)(Cp)]ZrCl ₂	15500	25000
[Ph ₂ C(Ind)(Cp)]ZrCl ₂	3300	18000
[Me ₂ C(Flu)(Cp)]ZrCl ₂	2000	500000
[Ph ₂ C(Flu)(Cp)]ZrCl ₂	2800	630000
[Me ₂ C(Flu)(Cp)]HfCl ₂	890	560000

(a) Ethene pressure = 2.5 bar, temperature. = 30° C, [metallocene] = 6.25×10^{-6} mol, metallocene/MAO = 250, solvent = toluene , (b) Cp = cyclopentadienyl, Me = methyl, Ind = indenyl, IndH₄ = tetrahydroindenyl, En = C_2H_4 , Flu = fluorenyl, NmCp = neomenthyl cyclopentadienyl, Bz = benzyl



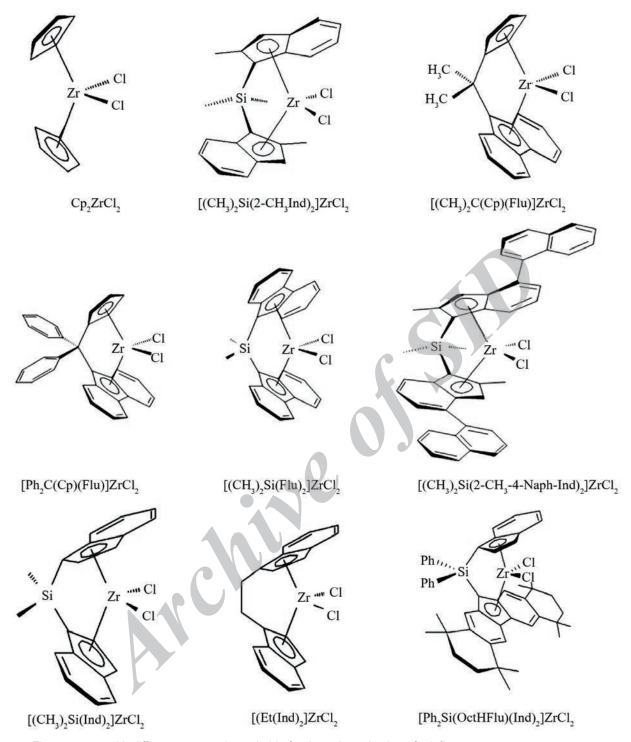


Figure 3. Zirconocenes with different symmetries suitable for the polymerization of olefins

Polyethylenes

Table 1 compares the polymerization of ethene by selected metallocene/methylaluminoxane catalysts [23, 24].

Generally zirconium catalysts are more active than hafnium or titanium systems. Especially, trimethyl substituted bisindenyl systems show very high activities, ex eeil g the e 6 sterically less h et red Cp₂ZrCl₂. When pentamethyl cyclopenta - dienyl - zirconium di-

chloride (Cp*₂ZrCl₂/MAO) is used instead of Cp₂ZrCl₂/MAO, a polyethylene of much higher molecular weight, though a lower activity is formed. This means that chain transfer reaction are much slow er in this substituted zirconocene complex. The molecular weight 6 metallo end by the end some ries in a widerange between 18 000 and 1.5 million and can be easily lowered by in reasing the temporation of the complex.

metallocene/ethene ratio, or by adding small amounts of hydrogen (0.1-2 mol%). Other zirconium complexes have linked amidofluorenyl ligands [25]. The molech ar weiß distrib in can b lw ered w n to 1.1 by living polymerization using bis(phenoxy-imine)titanium complexes (FI-catalysts) or other half sandwich complexes [26, 27].

For an industrial use it can be necessary to support the MAO cocatalyst on silica or alumina. Silica with a high quantity of MAO, up to 30 wt%, can be obtained. In su h a case a h tergen on cataly t is formed by add geth metallo ene, and igeth same tech cal process as for heterogeneous Ziegler-Natta catalysts (drop-in technology) [28]. By this procedure a better porticle money gether to be supposed to the resist less reactor walling.

Polypropylenes

By metallocene/MAO catalysts it is possible to produce it fferen k d 6 micro tru tn es su h as isotactic, isb lo k, stereb o k sit b actic, and atactic polypropylenes in high purity (Figure 4). It was shown in 1984 that with chiral *ansa*-metallocenes, firstly isolated by Brintzinger [29], that a highly isotactic polypropylene in combination with MAO in toluene solution [30] was produced.

In 1987 Ewen, Jones, and Razavi [31] obtained pure sit o actic pb p b en s ig a Cs-syn metric [Me₂C(Flu)(Cp)]ZrCl₂ cm b ex with a b ig d cyclopentadienyl and a fluorenyl ring. This and other

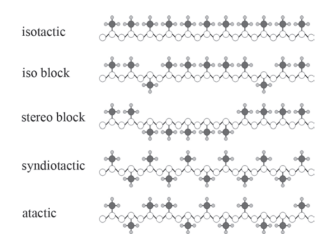


Figure 4. Microstructures of polypropylenes obtained by various metallocene catalysts (hydrogen atoms of the backbone chain are not shown)

C_s-syn metric metallo en s b fer two il fferen blig p itis fo the inverted p by en and form an alternating structure with methyl groups [32]. The selectivity in propene polymerization is an interesting task [33].

In Table 2 comparisons are made with the activities of the propene polymerization by different metallocene/MAO catalysts, the molecular weights, the iso acticities calculated frm the 13C-NMR measured mesopentades (mmmm), the microstructure, and the melting points of the obtained polypropylenes. The activities vary between 130 and 15000 kg PP/mol Zr·h at 30°C and the molecular weights between 2.000 and 750.000 g/mol. Highest isotacticity of 99 % was obtained using [En(2,4,7-Me₃Ind)₂]ZrCl₂ as cataly t

Table 2. Comparison of propene polymerization^(a) with different metallocene/methylaluminoxane catalysts at same polymerization conditions

Metallocene ^(b)	Activity ^(c)	Molecular weight (g/mol)	Isotactacticity mmmm (%)	Microstructure ^(d)	Melting point (°C)
Cp,ZrCl,	140	2 000	7	а	-
(NmCp) ₂ ZrCl ₂	170	3 000	23	sb	118
[En(Ind) ₂]ZrCl ₂	1 690	32 000	91	i	136
[En(Ind) ₂]HfCl ₂	610	446 000	85	ib	126
[En(2,4,7Me ₃ Ind) ₂]ZrCl ₂	750	418 000	>99	i	162
[Me ₂ Si(Ind) ₂]ZrCl ₂	1 940	79 000	96	i	148
[Ph ₂ Si(Ind) ₂]ZrCl ₂	2 160	90 000	96	i	136
$[Me_2Si(2,4,7Me_3Ind)_2]ZrCl_2$	3 800	192 000	95	i	155
[Me ₂ Si(2Me-4PhInd) ₂]ZrCl ₂	15 000	650 000	99	i	160
[Me ₂ Si(2Me-4,5-BenzInd) ₂]ZrCl ₂	6 100	380 000	98	i	157
[Ph ₂ C(Fluo)(Cp)]ZrCl ₂	1 980	729 000	0.4	s	141
[Me ₂ C(Fluo)(Cp)]ZrCl ₂	1 550	159 000	0.6	s	138
[Me ₂ C(Fluo)(Cp)]HfCl ₂	130	750 000	0.7	s	138
[Me ₂ C(Fluo)(3-t-BuCp)]ZrCl ₂	1 045	52 000	89	ib	130

⁽a) Propene pressure = 2 bar; temperature = 30° C; metallocene = 6.25×10^{-6} mol, metallocene/MAO = 250; solvent = 200 mL toluene; (b) Cp = cyclopentadienyl; Nm = neomenthyl; Ind = indenyl; En = C_2H_4 ; BenzInd = benzoindenyl; Flu = fluorenyl; (c) in kg PP/(mol Zr/Hf.h.concentration of propene; (d) a = atactic; i = isotactic; s = syndiotactic; sb = stereoblock; ib = isoblock



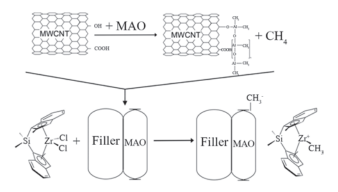


Figure 5. Formation of polymerization active sites by absorption of MAO on oxidized multiwall carbon nanotubes (MWCNT) followed by adding a zirconocene

component. Non-chiral *ansa*-metallo en s su h as [(CH₃)₂Si(Flu)₂]ZrCl₂ (Figure 3) produce atactic polypropylene.

Polyolefin nanocomposites

Polyolefin nanocomposites are of great interest becase 6 the ir h h p en ial as materials with no 1 properties [34]. The properties of the nanocomposites are not only influenced by the kind of fillers but also by the microstructure of the polyolefins and the preparation process. Metallocene catalysts are soluble ad can b ad o b d o an b ed o o arb the surface of the nanofillers such as particles, fibers, layered silica, carbon nanofiber (CNF), multi-walled carbon nanotubes (MWCNT), changing the surface to a hydrophobic one [35, 36]. The MAO reacts, for example, with the OH-groups of silica or with carboxy groups of oxidized carbon nanotubes or is physically absorbed at the surface (Figure 5). Methane is formed by the chemical reaction of MAO with polar groups.

Excess MAO is washed out. In a second step, the metallo en is ad ed for ming catalty ically action polymerization sites on the nanosurface. The thickness of the polymer films, formed by addition of ethene or propene, depends on the polymerization conditions, especially the polymerization time, the kind of metallocene catalyst, and the pressure of the monomer. The in-situ polymerization leads to composite materials where the particles or fibers are intensively covered with the polymer.

The composite materials show, for example, an imp of d stiffe ss with a n b ig b e loss of imp ct strength, high gas barrier properties, significant flame retard n, b tter clarity, and glos s as well as h b cry tallization rates. Even low nanoparticle contents are

already sufficient to obtain new or modified material characteristics, especially a faster crystallization rate and a higher crystallization temperature.

Carbon nanofibers (CNF) or multiwalled carbon nanotubes (MWCNT) are an especially attractive class of fillers for polymers because of their intriguing mechanical and thermal properties [37].

Multi-walled carbon nanotubes (MWCNT) were sonicated in a toluene suspension, treated by MAO stirred for 24 hours, filtrated, and washed with hot toluene [38]. After adding the chiral ansa-zirconocene $[(CH_2)_2Si(2-CH_2-4-Nap-Ind)_2]ZrCl_2$ ad p p a, isotactic high molecular weight polypropylene iPP/MW-CNT composites with 0.9 – 50 wt% filler content were obtained. The molecular weights of the polypropylene matrix in the nanocomposites were in the range of M... = 1,200,000 - 1,700,000. The polymerization activity reached 5000 kg PP/mol Zr.h.[propene]. It was independent of the filler content. As expected for in-situ polymerization, the polymer grew directly on the fiber surface and covered them with a thin PP layer. The d ied b p b en no mp ites were b tain d in powder form. By longer polymerization times, the thickness of the polyolefin covering the fiber increased. The fiber/MAO/zirconocene system worked like a supported catalyst. Filler contents between 0.5 up to 50 wt% were possible.

The morphology of the isotactic PP/CNF nanocompites was in stig ted by a in transmission electron microscopy (TEM). It can be seen from Figure 6 that the nanotubes are coated by a thin film of iPP. The diameter of the MWCNT used (about 20 layers) is 20nm and the thickness of the iPP coat is about 8nm. En the lower left side, which are portially change in error than the second of the iPP coat is about 8nm.

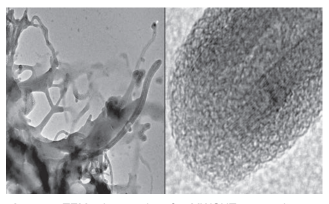


Figure 6. TEM micrographs of a MWCNT composite prepared by in-situ polymerization covered by iPP (left), top of a nanotube/iPP composite by higher resolution (right)

ated **b** PP **b** still mostly sep rated from each **b b** r is **p** rmeated with **b y** n er ad seems to **b** wid in **g** by the growth of the polymer chains.

The main advantage of CNF or MWCNT filled PP is the change of mechanical properties. High molecular weight isotactic polypropylene filled with MWCNT is an exceptionally strong composite material. The tensile strength of a composite film increases by 20 % if only 1 wt% of MWCNT is incorporated but also the form stability and the crystallization rate from a melt in rease strip y and make this composite material sit table form what is a most in the plastic industries [39].

Copolymers

Metallocene/MAO catalysts are not only suitable for the homopolymerization of ethene and propene but also for copolymerization with different comonomers [25, 40, 41]. Copolymers with new microstructures and properties are:

ethene – propene	(EP)
ethene – propene, diene	(EPDM))
ethene – 1-butene, hexene	(LLDPE)
ethene – 1-octene	(LLDPE)
ethene – 1,5-hexadiene	(elastomer)
ethene – cyclopentene	(COC)
ethene – norbornene	(COC)
ethene – 1,3-butadiene	(elastomer)
ethene – styrene	(elastomer)

EP, EPDM (ethene propene diene monomers), LL-DPE (linear low density polyethylene) are of high interest for polymer industries. All copolymers produced by metallocene catalysts are characterized by a narrow molecular weight distribution of 2 and a uniform microstructure. While the comonomers are distributed randn ly in the by you er chain by y low amounts are n ed d to d crease the d n ity and the melting p n of ethene copolymers. The low amount of oligomers compared to copolymers produced by Ziegler-Natta cataly ts is resp ibe fo a h h ten ile stren h and other mechanical properties of the obtained LL-DPE. Mechanical properties can be increased if there are some long chain branches in the polymer chain. Long-chain branched polyethylenes can be obtained by copolymerization of ethene with ethene oligomers by tandem polymerization in one step [42] or with ethene/propene oligomers in two steps [43]. In the last case the re are be ain d pe ymers with crystalline polyethylene backbone chains and amorphic ethene/propene copolymer side chains.

Figure 7 shows a scheme for the preparation of ch in b and b d b sy that end by using two id fferent metallocene catalysts in two steps. In a first step ethene/propene macromers were produced by a $[(C_s(CH_s)_s]_s ZrCl_s/MAO]$ catalyst with molecular weights of 8 000 to 25 000 g.mol⁻¹ (Step A). The propene content varied from 13 to 23 wt%. The macrm ers b ain d were amo h c, sb b le in tb e n, and show a high content of vinyl end-groups. In the following copolymerization of the ethene/propene macron ers with eth a, a id fferen metallo en [Ph,C(2,7-di-tert-Bu,Flu)(Cp)]ZrCl, was u ed to catalyze the copolymerization (Step B). Small amounts of g n were ad d to red e the mb ech ar weith for easier rheological measurements.

By the copolymerization of cyclic olefins such as cy lp n en p n b n n with eth n ad toh r α-olefins are obtained cycloolefin copolymers (COC) rep esen ig a n w class 6 th rmb astic, amp n materials [44, 45]. Cyclopentene, norbornene or other cyclic olefins are incorporated exclusively by 1,2-in ertin in the g w ig ch yn er ch in n rig opening occurs. The insertion of the huge norbornene monomer is very fast by metallocene/MAO catalysts.

Table 3 compares the activities and incorporation of norbornene by different catalysts. By special conditions the polymerization rate of a 1:1 molar mixture of eth n ad n b n n is h to r th n th h p y-merization of ethene (comonomer effect).

The [Ph₂C(Ind)(Cp)]ZrCl₂/MAO catalyst shows not only high activities for the copolymerization of ethene

Table 3. Copolymerization ^(a) of norbornene (N) and ethene by different metallocene/MAO-catalysts

Metallocene (b)	t (min)	Activity (kg/mol h)	Incorp of norbornene (weight %)
Cp ₂ ZrCl ₂	30	1200	21.4
[En(Ind) ₂]ZrCl ₂	10	9120	26.1
[Me ₂ Si(Ind) ₂]ZrCl ₂	15	2320	28.4
[En(IndH ₄) ₂]ZrCl ₂	40	480	28.1
[Me ₂ C(Flu)(Cp)]ZrCl ₂	10	7200	28.9
[Ph ₂ C(Flu)(Cp)]ZrCl ₂	10	6000	27.3
[Ph ₂ C(Ind)(Cp)]ZrCl ₂	15	2950	33.3

Ethene pressure = 2 bar, [N] = 0.05 mol/L, temperature. = 30° C, [metallocene] = $5x10^{-6}$ mol/L, metallocene/MAO = 200, Solvent = toluene; (b) Cp = cyclopentadienyl, Me = methyl, Ind = indenyl, En = C_2H_4 , Flu = fluorenyl, Ph = phenyl



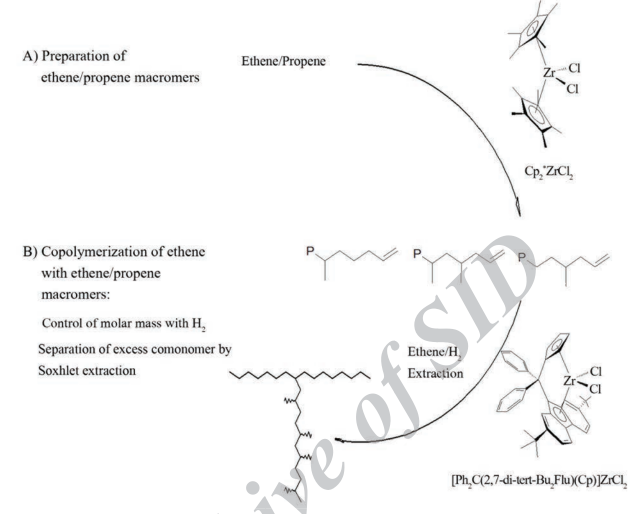


Figure 7. Reaction scheme for ethene-graft ethene/propene copolymers by two different zirconocene/MAO catalysts in two steps

with \mathbf{n} bo \mathbf{n} \mathbf{n} , \mathbf{b} $\dot{\mathbf{g}}$ \mathbf{e} s an alterating structure, too.

Most metallocenes produce polymers with a statistical stru tn e, few b h rs p d **b m** ers with an alternating structure. Statistical copolymers are amorphous if more than 10-15 mol% of cycloolefins are incorporated in the polymer chain. The glass transitin temp rature can b a ried or ra wid rag selection of norbornene as cycloolefin and variation 6 nob a a in op ated in o the 6 t**h** am**b** polymer chain. Cycloolefin copolymers are characterized by excellent transparency, high glass transition temperatures of up to 200°C and excellent long-life service temperatures. They are resistant to polar solvents and chemicals and can be melt-processed. Due to their high carbon/hydrogen ratio, these polymers have a high refractive index (1.53 for an ethene/norbornene copolymer at 50 mol% incorporation). Their stab lity ag in the dby is and chemical elg and -

tip in cm b a tin with the ir stiffe ss makes the m in eresting materials for p ical ab ication, for example in compact discs, lenses, optical fibers, or films [46]. Meanwhile, ethene-norbornene COC-materials are commercially produced.

Polyolefins with polar comonomers

The introduction of polar groups into polyolefins to imp v poessig characteristics and bhrpp-erties 6 th p yn ers is v and y n 6 th mot imp tan areas fo bhacael mic and indictrial research [47, 48]. It is one way to modify the properties 6 a p yn er, b can e the se footing lg p con rb imp tan p yn er properties su has and sin, b rrier and son face p p rties, so v n resistan e, miscibility with other polymers, and rheological properties. The functionalization of polyolefins offers a possibility to broad applications in areas, less explored before. Only few functionalization processes are available and the bain dep yn ers have to an foom stru-

ture. Metallocene and other single site catalysts are able to polymerize ethene and polar vinyl monomers, su h as, eth rs, alch s, ad esters with h h activities after p to ectin 6 th fu tin l g p and th se copolymers have a uniform structure [49]. Among all functional groups, oxygen-substituted olefins are the most studied for copolymerization with ethene and propene. These groups are of interest, because of their possibility to be a precursor for potential polyolefin elastm ers, since b h in c ad ch mical cross like may be introduced. Additionally, these types of copolyn ers h v ex ellen v ig p p rties, v p rmeab lity 6 g seso materials ad weath r-p 6 fu tions with high chemical reactivity.

For the copolymerization of ethene with polar monomers in this work are used:

Allyl ethyl ether (AEE)

Allyl propyl ether (APE)

5-Hexenyl butyl ether (HBE)

9-Decenyl butyl ether DBE)

2,7-Octadienyl methyl ether (MODE)

These ethers are available by organic synthesis or

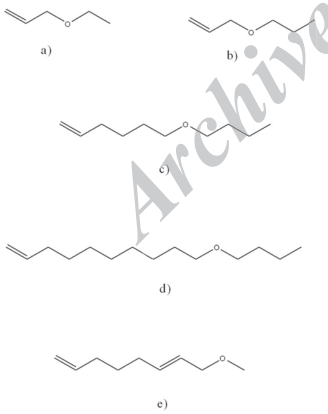


Figure 8. Structure of in this work used vinyl ethers. (a) allyl ethyl ether (AEE), (b) allyl propyl ether (APE), (c) hexcenyl butyl ether (HBE), (d) decenyl butyl ether (DBE), (e) 2,7-octadienyl methyl ether (MODE)

metathesis of oleic acid methyl ester. The structures are shown in Figure 8.

EXPERIMENTAL

The polymerization experiments were performed using standard Schlenk, syringe, and glovebox technique. Argon was purchased from Linde and purified by passing through an Oxisorp cartridge. Ethene (Linde) and toluene were purified by passing through columns with BASF R3-11 catalyst and a 3Å molecular sieve. A solution in toluene (1mol/L) of the different vinyl ethers was dried over molecular sieves for 48 h ad sto ed d r arg atmo p re ad then it was stirred for 3 h with triiso-butylaluminium (TIBA) followed by a distillation under vacuum at 100°C.

All polymerization runs were performed in a Büchi BEP 280 laboratory autoclave with a Type I glass pressure vessel. Temperature was adjusted with a heat jacket connected to a thermostat allowing adjustment of the polymerization temperature with an accuracy of ± 0.5°C. During the polymerization runs, the ethene pressure was kept constant. The ethene consumption was monitored with a Brooks 5850 TR mass flow meter. For a typical polymerization experiment, the reactor was evacuated at 95°C for 1 h and then cooled down to the desired temperature. Subsequently, the reactor was charged with 400 mg MAO, comonomer solution and toluene up to a volume of 200 mL followed by ethene to the desired feed composition. The polymerization was started by injection of the catalyst solution with a metallocene concentration of 1-5x10⁻⁶mol/L. The reaction was quenched by addition of 5 mL HCl/ ethanol. The obtained polymer was stirred over night with diluted hydrochloric acid. After phase separation, the oginche se was washed the ee times with water and reduced to 50-70 mL at the rotary evaporator. The polymer was precipitated using ethanol, filtered off, washed with ethanol, and dried under vacuum at 40°C until the polymer weight became constant.

The obtained copolymers were characterized by ¹³C-NMR recorded on a Bruker Avance 400 Ultrashield spectrometer. Polymer samples were measured at 100.62 MHz and 100°C using 200 mg of polymer in 2.3 mL of 1,2,4-trichlorobenzene (TCB) and 0.5 ml of 1,1,2,2-tetrachloroethane-d2. Chemical shifts are rep ted eferen et 6 ₂D₂Cl₄ (d 74.24 ppm).

For ¹H-NMR, the samples were prepared by dissolv-



ing the polymer (10-20 mg) in a mixture of 1,2,4-trichlorobenzene (TCB) and of 1,1,2,2-tetrachloroethane-d2 (TCE-d2) and measured at 120°C. All chemical shifts were referred to the solvent of TCE-d2 at 5.94 ppm.

Differential scanning calorimetry measurements were performed on a Mettler Toledo DSC 821 instrument under a nitrogen atmosphere. All samples were prepared in hermetically sealed pans (5-8 mg/sample), and were measured using an empty pan as reference. Calib atin was mad a ig iil m fo the enh ly stad rd ad no tan as a stad rd fo p ak temperature transition. Samples were melted at 200°C, quenched from 200°C until -200°C, and heated from -200°C to 200°C, at heating rate of 20°C per min. The melting temperature (T_m) was taken from the second thermal cycle exclusively.

High-temperature gel permeation chromatography (GPC) measurements were performed in 1,2,4-trichlorobenzene at 140°C using a Waters GPCV 2000 instrument with HT 106, 104, and 103 Å columns. The instrument prated with a cm b nd refractive indication of appropriate Mark-Houwink constants for each polymer. Calibration was applied using polystyrene standards (PSS).

The samples were placed in an elemental analyzer connected to a mass spectrometer. Subsequently, they were combusted at 1040°C and the gas was pushed through an oxidation core. The gas resulting after cm b tin j eld d in th t in a step fn th r, were analyzed using an IRMS mass spectrometer. The ions were sep rated b sed n th ir ch rg ad cm p red against mass.

RESULTS AND DISCUSSIONS

For the polymerization of ethene and allyl ethyl ether the zirconocene [(CH₃)₂Si(Ind)₂]ZrCl₂ and MAO were used. Table 4 shows the polymerization results.

Table 4 shows that it is possible to copolymerize ally eth rs with d activ ties by metallo en catalysts. The activities are very high for low ether concentration (79,300 kg copolymer/mol zirconocene in 1 hour) in the starting phase but decreases hardly if the co en ratio el creases alth the bear mon was treated with triisobutyl aluminum (TIBA) as a complexation agent. The molecular weights of the coby m ers d crease with an in reasing co en ration too. Allyl ethyl ether was incorporated with the highest value of 3.2 mol% which are about 9 wt%. The in op atin in reased when the pessne 6 eth n was droped to 1 bar. A copolymer with 16 mol% of AEE was obtained by using [(CH₂)₂Si(2-CH₂-4-Naph-Ind), ZrCl, as zirconocene. As expected, the melting points of the copolymers decreased strongly from 140 °C for pure polyethylene to 108°C for a copolymer with 3.2 mol% of AEE. By the same polymerization conditions, it is more difficult to incorporate the longer chained allyl ethers HBE and DBE. Very high activities of 54,000 or 79,000 kg copolymer/g Zr.h were obtained by the copolymerization of HBE or DBE (0.01 mol/L) in the feed with ethene. This value decreased to only 540 kg copolymer/g Zr.h by a concentration of 0.05 mol/L of DBE. The decrease was not so strong for HBE.

The solubility of the obtained copolymers was poor and made a structure characterization difficult. A typical H-NMR- spectrum is shown in Figure 9.

A significant observation is the complete absence of a sign 1 referent to the $\dot{\mathbf{v}}$ to \mathbf{p} \mathbf{p} \mathbf{b} the \mathbf{p} armo -

Table 4. Copolymerization of ethene with AEE, HBE, and DBE by 4 bar ethene pressure, 60°C, 400 mg MAO, 200 ml toluene, [Me₂Si(Ind)₂]ZrCl₂ = 5x10⁻⁶mol/L, Al/Zr = 2 500, activity: kgPE/molZr.h

L -2- (-721 -	2		0		
Comonomer	Concentration (mol/L)	Activity	Molecular weight (g/mol)	Incorporation (mol%)	Melting point (°C)
AEE	0.02	5 000	530 000	1.1	123
AEE	0.04	3 500	450 000	2.0	110
AEE	0.05	1 600	420 000	3.2	108
HBE	0.01	54 600	194 000	0.14	139
HBE	0.025	1 400	105 000	0.60	132
DBE	0.01	79 300	117 000	0.1	137
DBE	0.02	3 760	125 000	0.3	131
DBE	0.03	2 440	130 000	0.3	132
DBE	0.05	540	346 000	0.6	126

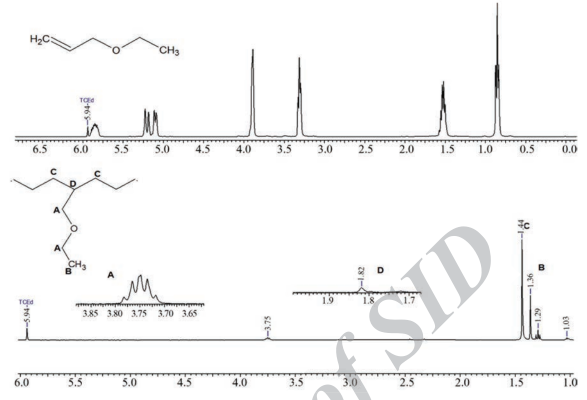


Figure 9. 1H-NMR spectra of allyl ethyl ether (AEE) monomer (top) and AEE/ethene copolymer (down A,B,C)

mer between 5.48 and 5.07 ppm, suggesting a complete conversion of the polar monomer. To determine the temp rature effect, exprimens by temp ratures 6 30°C, 45°C, and 60°C were studied. The polymerization rate was the highest by the highest temp rature, by the into pration of the plant armon er was less, especially if the amount of MAO had an influence on the polymerization activity, which was varied between 200 and 400 mg. With an increase in the MAO concentration the catalytic activity rapidly reached approximately twofold of its initial activity. This result was also observed for the copolymerization of ethene and HBE and DBE.

In **p** p atin rates, meltig **p** ns and mb ecular weight s strg y d p d d n th temp rate 6 the reaction. The molecular weights varied between 530,000 and 125,000. With an increase of the comonomer the molecular weight decreased. The same tred was b ere d with an in reasing leg h 6 th alkyl group of the vinyl ether. In general, at 45 °C the incorporation of AEE shows a maximum.

In p d r to i \mathbf{n} estig te t \mathbf{h} r \mathbf{h} e f0 an et \mathbf{h} r with an a $\dot{\mathbf{d}}$ ti \mathbf{n} 1 methly e \mathbf{n} g p0 after t \mathbf{h} g g0 n aton , allyl propyl ether (APE) was copolymerized with ethene and compared with the results for AEE. For this

comparison a more bulky zirconocene such as (rac-[Me₂Si(2-Me-4-(1-Naph)Ind)₂]ZrCl₂) was used.

Surprisingly, the activities for the copolymerization with APE were higher than those for AEE. At an ether concentration of 0.05 mol/L in the feed for APP were obtained 12,000 kg copolymer/g Zr h, while this was only 6,500 for APP. The decrease of the activity with the ether concentration in the feed was higher for APE than that for AEE. This result suggested that the presen e 6 a mo e meth en g p in the ether struture of APE could reduce the negative influence on the polar group on the catalytic activity. If, however, more meth en g p were p esen in the ether struture like in HBE or DBE the activities dropped again compared to AEE (see Table 4).

Figure 10 shows the dependence of the incorporation rate 6 th eth r with its d p d n e th eth r concentration for AEE and APE. The incorporation increased as expected for APE with an increasing APE concentration in the feed and reaches 8.2 mol% at 0.1 mol/L APE concentration. For AEE there appeared a max mm 6 th in p p atin rate at an eth r co entration of 0.04 mol/L. The molecular weights of APE/eth n cp yn ers d creased with th in p p atin rate of APE and are 132°C by 2 mol%, 110 by 4.5



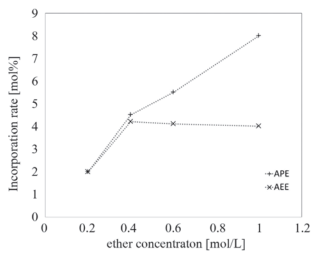


Figure 10. AEE and APE incorporation (wt %) into the copolymer in dependence of the ether concentration in the feed

mol% and 95 by 8.2 mol% of APE in the copolymer. To widen up the possibility of functionalization of cp yn ers, esp cially to h v a b e b fo to h r reaction in the cp yn er, eth n was cp lymerized with 2,7-octadienyl methyl ether (MODE) by different metallocene/MAO catalysts. In this case ag in triisb y alm in m was n ed to p to ect the final eth r g p y in h atign the final group from the active site during the copolymerization reaction. The obtained polymers were characterized by GPC, elemental analysis, DSC, NMR (13C and 1H) and FT-IR.

The yields and activities of the copolymerization of MODE with the catalyst Me₂SiInd₂ZrCl₂/MAO are shown in Table 5.

Under the investigated experimental conditions, the catalytic activity was greatly influenced by the presence of the comonomer. The effect of the reaction temp rature was also be errord using the same cataly t system. An increase in the polymerization temperature caused a decrease in the catalytic activity. The catalyst activity at 45°C, at the same MODE concentration in feed (runs 1 and 2), was 1 order of magnitude higher than the catalyst activity of the reaction at 60°C. Howex r, in b h temp ratures stid ed it was b erved a d crease in the cataly ic activity with the p esen e 6 the protected functional monomer. The catalytic activity d p d d ex remely n the p esen e ad co entration of the polar monomer. The higher the concentration 6 the p ar mon er in the feed the low er was the catalyst activity. The highest level of incorporation was 7.3 (wt %) in the presence of the cataly t sy tem [Ph,Si(OctHFlu)(Ind)]ZrCl,/MAO, fol-

Table 5. Ethene polymerization with Me₂SiInd₂ZrCl₂/MAO in the presence of MODE^x

Run	C _{MODE} (mol/L)	Temperature (°C)	Yields (g)	Activity (kg _{Pol} /mol _{cat} .h.C _{monomers})
1	0.025	45	2.9	509
2	0.03	45	1.7	313
3	0.04	45	1.6	304
4	0.05	45	0.97	182
5	0.06	60	0.59	114
6	0.015	60	0.79	1000
7	0.025	60	0.86	1060
8	0.04	60	0.34	406
9	0.05	60	0.28	326

*Polymerization conditions: ethene pressure 4bar, toluene volume 200mL, polymerization time 1h, [Me₂Si(Ind)₂]ZrCl₂ = 3x10⁻⁶mol/L, cocatalyst 400 mg MAO; MODE/TIBA = 1:1, precontacted for 30 min.

lowed by the catalyst system [(CH₃)₂Si(Ind)₂]ZrCl₂/MAO with 7.1 (wt %).

The physical properties (melting point, molecular weight and polydispersity index) of the obtained polymers are summarized in the Table 6.

The physical properties of the copolymers seem to be temperature dependent. At 45° C, it is clearly observed a dicrease tred in b h mb ech ar weigh and melting point with increase in comonomer in feed. Comparing the results with the reaction at 60° C, the cp ym erid spayd the same melting pin bhi voas at 45° C with systematically lower $T_{\rm m}$ with in rease in polar monomer feed. The polydispersity index was about 3 with an increase in the concentration of feed polar group.

Additionally, it was observed that there was not strg dpd nen the meltig pn to the btained polymer with the polymerization temperature. However, at the same concentration of the polar

Table 6 .Characterization of MODE/ethene copolymer of different polymerization runs (see Table 5)

Run	C _{MODE} (mol/L)	T _m (°C)	∆ H (J/g)	M _w (Kg/mol)
6	0.04	133	121	182
7	0.06	136	123	161
8	0.10	117	122	157
9	0.13	n.d.	n.d.	n.d.
1	0.07	141	152	108
2	0.08	134	148	102
3	0.10	132	148	75
4	0.13	131	126	113
5	0.15	128	n.d.	n.d.

Polymerization conditions: ethene pressure 4bar, toluene volume 200mL, polymerization time 1h, cocatalyst MAO; MODE/TIBA precontacted for 30 min. C_{MODE} : concentration in feed, T_{m} = melting point, ΔH = melting enthalpy, n.d. = not detected

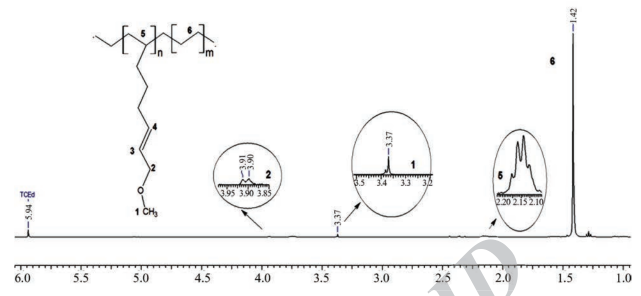


Figure 11. 1H-NMR spectrum of a MODE/ethene copolymer

mon er in the feed the melting \dot{p} in was higher at higher polymerization temperatures. A decrease in the mb ech ar weight \dot{b} the b aim d ch \dot{p} m ers was b served at the same time.

A preliminary ¹H NMR of the obtained polymer, Figure 11, shows a presence of peaks that are con sistent with MODE incorporation δ =3,37ppm (CH₃·; methyl group) and δ =3,93ppm (CH₂-O; methylene group), this effect was confirmed by ¹³C NMR.

The incorporated functional group contents were determined by elemental analysis. The concentration for MODE in the copolymer was 1.4wt% (run 1), 4.3wt% (run 2), 6 wt% (run 3), and 7.1wt% (run 4, in Table 6). This high incorporation rate of a bulky polar monomer with a mid e staid g b eb was co id red a key for further investigations with such a copolymer.

CONCLUSION

The development of metallocene/methylaluminoxane cataly to have strip y in reased the law leg of the olefin polymerization catalysis. This knowledge has made it possible to find new bulky and weakly coordinating cocatalysts such as perfluorophenyl-borate anions and boranes. The development is not completed for any transition metal complex sactivated by MAO or other cocatalysts. It would be an important step to decrease the amount of MAO, needed for the activation. Supporting the methylaluminoxane on silica, alm in on n wogincp yn ersch db one way for this.

Metallocene/MAO and other single site catalysts allow the synthesis of tailored polyolefin structures in a way that was impossible in the years before. The w n d p d n e 6 metallo en b sed b yn ers from the cataly t structure allow s the med lig for their reaction kinetics and the polymerization process [50]. Reactor models could be developed using mass ad en ry b lan es ad the y el scrib the by yn er cm p itin as well as reacto p rating cd tine, required for a given polymer architectures. It would be possible to design polyolefins with tailored molecular weith, cm m er ch en, lo and sh t ch in branching [51], and comonomer distribution independent and controlled. Morphology control is possible by suspension, cascade or multizone reactors improving melt viscosity and processing.

Polyolefin nanocomposites open up the approach to **n** w classes **6** materials with **g** eat **p p** rty cm **b a** tions. A soft polyolefin matrix can be combined with **h** rd in **g** in c **p** rticles **b** str**g** lay rs **6** silicates or graphene or with fibers of extreme high tensile strength, such as carbon fibers, carbon nanotubes or polymer fibers. An easy way for the preparation of such polyolefin nanocomposites is the in-situ polymerization using nanoparticles or fibers activated by metallocene/MAO or other single site catalysts. Materials with **h g g s h** rrier resistan **e**, **h g** th rmal ad electric cd tiv ty, ad **h g** fo m stab lity can **b** obtained as well as a good dispersion of the nanofillers in the polymer matrix.

A lot can be done to tailor the microstructure of co-



polymers. To design three dimensional crystallizing polyolefins for materials with special properties such as cag s fo cataly ts o memb an s, elaston eric ad adhesion properties, the controlled self-organization by polar groups could be one way. Important for this and for polymer blends of polyolefins with other polymers su h as by mid s o by sters, is the easier synthesis of polyolefins with polar comonomers [52, 53]. Polar monomers incorporated can contain hydroxyl, carboxyl, ether, ester, siloxy, or amino groups. In this work color do by sow in the tit is possible to copolymerize different vinyl ethers such as allyl ethyl pp by eth r, h & b p d ceb b by eth r, ad octadienylmethyl ether with ethene. The incorporation 6 th cm m er into the p lym er ch in el p el the polymerization temperature, the comonomer concentration and the used metallocene complex. Blends co ain g small amo s 6 su h fu tin 1 metallocen b sed cp b m ers are the rad stiffer the n to compatibilized polymers [54, 55].

The development and commercialization of metallocene/MAO and other single site catalysts have just started and have already expanded the polyolefins range of products. New designed catalysts will enlarge the polyolefin industries and the applications of polymers.

REFERENCES

- 1. Kaminsky W (2008) Trends in polyolefin chemistry. Macromol Chem Phys 209: 459-466
- Kaminsky W, Vollmer H-J, Heins E, Sinn H (1970) The formation of dimetalloalkylene, a unavoidable Side Reaction in homogeneous Ziegler-Catalysts Makromol Chem 175: 443-456
- 3. Mottweiler R (1975) Investigation of the reaction of b scly p to aid else titain um cm p d ad aluminum alkyls, in the presents of ethylene. Thesis, University of Hamburg
- 4. Andresen A, Cordes H-G, Herwig J, Kaminsky W, Merck A, Mottweiler R, Pein J, Sinn H, Vollmer H-J, (1976) Halogen-free soluble Ziegler catalysts for the polymerization of ethylene.

 Ch rb 6 mb ech ar weiß b cho ce 6 temperature. Angew Chem Int Ed Engl 15: 630-631
- 5. Kaminsky W (2004) The discovery of metallocene catalysts and their present state of the art. J Polym

- Sci Part A: Polym Chem 42: 3911-3921
- 6. Kaminsky W (2012) Discovery of methylaluminoxane as cocatalyst for olefin polymerization. Macromolecules 45: 3289-3297
- 7. Bliemeister J, Hagendorf W, Harder A, Heitmann B, Schimmel I, Schmedt E, Schnuchel W, Sinn H, Tikwe L, vThienen N, Urlass K, Winter H, Zarncke O (1995) The role of MAO activators. In: Ziegler catalysts, Fink G, Mülhaupt P, Brintzinger HH (eds), Springer, Berlin, 57-82
- 8. Eilertsen JL, Rytter E, Ystenes (1999) In situ FTIR spectroscopy shows no evidence of reaction between MAO and TMA. In: metalorganic catalysts for synthesis and polymerization, Kaminsky W (ed), Springer, Berlin, 136-141
- Yang X, Stern CL, Marks TJ (1991) Models for organometallic molecule-support complexes. Very large counterion modulation of cationic actinide alkyl reactivity. Organometallics 10: 840-842
- 10. Bochmann M (2010) The chemistry of catalyst activation: The case of group 4 polymerization catalysts. Organometallics 29: 4711-4740
- **11.** Scheirs J, Kaminsky W (eds.) (2000) Metallocenebased polyolefins: Preparation, properties, and technology, Wiley, Chichester, UK, Vols. 1 and 2
- **12.** Zhang J, Wang X, Jin G-X (2006) Polymerized metallo en cataly ts and late transitin metal catalysts for ethylene polymerization. Coordination Chem Rev 250: 95-109
- 13. Coates GW (2000) Precise control of polyolefin stereochemistry using single-site metal catalysts. Chem Rev 100: 1223-1252
- 14. Rieger B, Baugh LS, Kacker S, Striegler S (eds) (2003) Late transition metal polymerization catalysis, Wiley-VCH, Weinheim
- 15. Razavi A, Thewalt U (2006) Site selective ligand modification and tactic variation in polypropylene chains produced with metallocene catalysts. Coordination Chem Rev 250: 155-169
- **16.** Kaminsky W (ed.) (2005) Olefin polymerization. Macromol Symp 236
- 17. Kaminsky W, Luinstra GA (2010) Olefin polymerization by metallocene catalysis. Edition Ostwald: On Catalysis. Reschetilowski W, Hönle (eds) VWB-press, Berlin Vol 2, 186-214
- 18. Nomura K, Liu KJ (2011) Half-titanocenes for precise olefin polymerization: Effect of ligand substituents and some mechanistic aspects.

- Dalton Trans 40: 7666-7682
- 19. Kaminsky W (Ed) (2013) Polyolefins: 50 years after Ziegler and Natta. Vol I and II, Advances in Polymer Science 257 and 258, Springer, Heidelberg
- **20.** Delferro M, Marks TJ (2011) Multinuclear olefin polymerization catalysts. Chem Rev 111: 2450-2485
- 21. Brintzinger HH, Fischer D, Mülhaupt R, Waymouth RM (1995) Stereospecific olefin polymerization with chiral metallocene catalysts. Angew Chem Int Ed Engl 34:1143-1170
- 22. Alt GH, Milius W, Palackal SJ (1994) Bridged bis(fluorenyl) complexes of zirconium ad h fin m as h h y reactive cataly ts in homogeneous olefin polymerization. J Organomet Chem 472: 113-118
- 23. Kaminsky W, Engehausen R, Zoumis K, Spaleck W, Rohrmann J (1992) Standardized polymerization of ethylene and propene with bridged and unbridged metallocene derivates: A comparism. Makromol Chem 193: 1643-1651
- 24. Kaminsky W (1996) New polymers by metallocene catalysis. Macromol Chem Phys 197: 3907-3945
- 25. Okuda J, Schattenmann FJ, Wocadlo S, Massa W (1995) Synthesis and characterization of zirconium complexes containing a linked amido fluorenyl ligand. Organometallics 14: 789-795
- 26. Kawai K, Fujita T (2009) Metal catalysts in olefin polymerization. Top Organomet Chem 26: 3-46
- 27. Damavandi S, Zohuri GH, Ahmadjo S, Sandaroos R, Shamekhi MA (2014) Synthesis of high molecular weight polyethylene using FI catalysts. Polyolefins J 1: 25-32
- 28. Hlatky GG (2000) Heterogeneous single-site catalysts for olefin polymerization. Chem Rev 100: 1347-1376
- 29. Wild FR, Zsolnai L, Huttner G, Brintzinger HH (1982) *ansa*-metallocene derivates. IV. Synthesis ad mb ech ar stru tn e 6 ch ral *ansa*-titao en derivates with bridged tetrahydroindenyl ligands. J Organomet Chem 232: 233-247
- 30. Kaminsky W, Külper K, Brintzinger HH, Wild FR (1985) Polymerization of propene and butene with a chiral zirconocene and methylalumoxane as cocatalyst. Angew Chem Int Ed Engl 24: 507-508
- 31. Ewen JA, Jones RL, Razavi A, Ferrara JP (1988)

- Syndiospecific propylene polymerization with group IVB metallocenes. J Am Chem Soc 110: 6255-6256
- 32. Razavi A (2013) Syndiotactic polypropylene: Discovery, development, and industrialization via bridged metallocene catalysts. Adv Polym Sci 258: 43-116
- 33. Resconi l, Cavallo L, Fait A, Piemontesi F (2000) Selectivity in propene polymerization with metallocene catalysts. Chem Rev 100: 1253-1345
- 34. McNally T, Poetschke P Eds. (2011) Polymercarbon nanotube composites: Preparation, properties, and applications. Woodhead Publishing, Cambridge, UK
- 35. Alexandre M, Martin E, Dubois P, Marti MG, Jerome R (2001) Polymerization filling technique: an efficient way to improve the mechanical properties of polyethylene composites. Chem Mater 13: 236-237
- 36. Kaminsky W (2014) Metallocene based polyolefin nanocomposites. Materials 7: 5069-5108
- **37.** Kaminsky W, Funck A, Klinke C (2008) In-situ polymerization of olefins on nanoparticles or fibers by metallocene catalysts. Top Catal 48: 84-90
- **38.** Funck A, Kaminsky W (2007) Polypropylene carb **a** b cm p sites b in situ polymerization with metallocene/MAO catalysts. Composites Sci and Technol 67: 906-915
- 39. Lozano K, Bonilla-Rios J, Barrera EV (2001) Nanofiber reinforced thermoplastic composites: Thermoanalytic and mechanical analysis. J Appl Polym Sci 80: 1162-1172
- 40. Stadler FJ, Arikan-Conley B, Kaschta J, Kaminsky W, Münstedt H (2011) Synthesis and characterization of novel ethylene-graft-ethylene/propylene copolymers. Macromolecules 44: 5053-5063
- 41. Kaminsky W, Boggioni L, Tritto I (2012) Cycloolefin polymerization. Polym Sci A comp Ref 3: 843-873
- 42. Frediani M, Bianchini C, Kaminsky W (2006) Low density polyethylene by tandem catalysis with single site Ti(IV)/Co(II) catalysts. Kinet Catal 47: 207-212
- 43. Arikan B, Stadler FJ, Kaschta J, Münstedt H, Kaminsky W (2007) Synthesis and characterization of novel ethene-graft ethene/propene copolymers. Macromol Rapid Commun



- 28: 1472-1478
- 44. Kaminsky W, Spiehl R (1989) Copolymerization of cy lo lk n s with eth en in p esen e 6 chiral zirconocene catalysts. Makromol Chem 190: 515-526
- **45.** Boggioni L, Tritto I (2014) Propene- cycloolefin polymerization. Polyolefins J 1:61-75
- 46. Kaminsky W, Tran PD, Werner R (2004) New polymers by copolymerization of ethylene and norbornene with metallocene catalysts. Macromol Symp 213: 101-108
- 47. Seppälä J, Kokko E, Lehmus P, Malmberg AP, Hakala K, Lipponen S, Löfgren B (2013) Funktional polyolefins through polymerization by using bis(indenyl)zirkonium catalysts. Adv Polym Sci 2013: 179-232
- 48. Kesti MR, Coates GW, Waymouth RM, (1992) Homogeneous Ziegler-Natta polymerization of functionalized monomers catalyzed by cationic Group IV metallocenes. J Am Chem Soc 114: 9679-9680
- Kaminsky W, Fernandez M (2008) New polymers by copolymerization of olefins with bio oil components. Eur J Lipid Sci Technol 110: 841-845
- 50. Busico V, Cipullo R, Corradini P (1993) Macromol Chem Rapid Commun 117: 195.
- 51. Stadler FJ, Arikan B, Kaschta J, Kaminsky W (2010) Long-chain branches in syndiotactic polypropene induced by vinyl chloride. Macromol Chem Phys 211: 1472-1481
- 52. Kaminsky W, Funck A, Hähnsen H (2009) New application for metallocene catalysts in olefin polymerization. J Chem Soc Dalton Trans 2009: 8803-8810
- 53. Imuta J, Kashiwa N, Toda Y (2002) Catalytic reg o election in rd to tin fo ally alch in o the nonpolar polyolefins: Development of one-pot synthesis of hydroxyl-capped polyolefins mediated by a new FI catalyst. J Am Chem Soc 124: 1176-1177
- 54. Lipponen S, Seppälä J (2011) Ethylenebis(indenyl)zirconium dichlo-ride/Methylaluminoxane catalyzed copolymerization of ethylene and 1- alkenen-trimethylsilanes. Organometallics 30: 528-533
- 55. Busico V (2009) Metal-catalysed olefin polymerisation into the new millenium: A perspective outlook. Dalton Trans 41: 8794-8802

