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# Sulfonated porous organic polymer supported Ziegler—Natta catalysts for the synthesis of ultrahigh molecular weight polyethylene

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Porous organic polymers (POPs) are attracting attention for their easy functionalization and potential as catalyst supports in olefin polymerization. In this study, sulfonated POP (s-POP) supported Ziegler–Natta catalysts were used for ethylene polymerization, producing ultra-high molecular weight polyethylene, with  $M_{\rm v}$  reaching up to  $6.83\times10^6$  g mol $^{-1}$ . The maximum  $M_{\rm v}$  of polyethylene was achieved by Cat-3 with DIBP as the internal donor, albeit with a partial loss of catalytic activity. Polymerization conditions also play a pivotal role in determining the molecular weight of polyethylene. Hydrogen, being the most efficient chain transfer agent, can decrease the molecular weight to  $9.68\times10^4$  g mol $^{-1}$  at higher hydrogen concentrations ([H $_2$ ]:[C $_2$ H $_4$ ] = 0.83), and the s-POP-supported ethylene polymerization catalysts were observed to exhibit high sensitivity to hydrogen response. The effects of polymerization temperature, [Al]:[Ti] molar ratio, and ethylene pressure on ethylene polymerization were thoroughly investigated.

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# 1. Introduction

Ultra-high molecular weight polyethylene (UHMWPE) is a specialized type of semi-crystalline thermoplastic engineering plastic known for its good chemical resistance, high impact resistance, high wear resistance, durability and biocompatibility. It plays an important role in fields such as national defense, military, marine engineering, ropes and textiles, and sports equipment. Additionally, UHMWPE has a Young's modulus of approximately 0.5–0.8 GPa, similar to human bones, with low friction coefficient and good biocompatibility, making it suitable for the production of artificial knee joints, artificial hip joints, artificial heart valves, and more.<sup>1-6</sup>

The performance of UHMWPE is closely related to its molecular weight, and place high demands on the catalyst system. Single-site catalysts, including metallocene catalysts, Fhenoxy imine-based catalysts, and post-transition metal catalysts, among others, can be utilized for the production of UHMWPE.<sup>7-15</sup> Through the design and synthesis of ligands, precise control over polymer structure can be achieved, resulting in high molecular weight narrow distribution UHMWPE, disentangled UHMWPE, branched UHMWPE, and other UHMWPE variants with specific structural properties. However, the industrial application of single-site catalyst systems still faces challenges such as reaction conditions, catalyst cost, and

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activity. Currently, most commercial UHMWPE is manufactured using conventional Ziegler–Natta (Z–N) catalytic systems employing TiCl<sub>4</sub>/MgCl<sub>2</sub>.<sup>16</sup> In the industry, Z–N catalysts continue to be the predominant catalyst system utilized for olefin polymerization. Ongoing research on Z–N catalyst systems primarily focuses on innovation and refinement catalyst supports. In comparison with traditional inorganic supports like MgCl<sub>2</sub> and SiO<sub>2</sub>, polymer supports can be synthesized using versatile strategies, allowing for easy modification with substituents and controlled pore structures.<sup>17,18</sup> And polystyrene or polysiloxane based supports have been proved to effectively prepare Z–N catalysts and catalyze ethylene/propylene polymerization.<sup>19-24</sup>

Porous organic polymers (POPs) composed mainly of C, H and O atoms are garnering increasing attention due to their easier functionalization to accommodate active cites, while maintaining minimal impact on the produced polymer properties.25-28 Based on our prior research, POPs based metallocene catalysts have demonstrated outstanding capabilities in ethylene homo-polymerization and ethylene/α-olefin copolymerization. Moreover, we found that the 4-hydroxyethylmethacrylate (HEMA) functionalized and sulfonated POPs could be modified using a Grignard reagent to synthesize Z-N catalysts. Despite the slightly lower activity of the resulting Z-N catalysts compared to the MgCl<sub>2</sub>-based catalysts, the prepared polypropylene displayed notable characteristics with broad molecular weight distribution (MWD > 11) and high stereoselectivity.29,30 The functional comonomer not only influences the properties of POPs, but also affects the microchemical environment of immobilized active centers. Thus, polymers

with specific structure and property can be prepared through the integrated design and preparation of functional POPs.

In the present study, 4-vinylbenzenesulfonic acid sodium salt and sodium allylsulfonate were selected as functional groups for synthesizing s-POPs, which reacted with methylmagnesium chloride to prepare Z-N catalysts. The findings from ethylene polymerization experiments revealed that the s-POP-based Z-N catalysts exhibited strong ethylene polymerization capability, yielding a polymer with a viscosity average molecular weight  $(M_y)$  as high as  $6.83 \times 10^6$  g mol<sup>-1</sup>, indicating their considerable potential for UHMWPE production.

# Materials and methods

#### 2.1 Materials

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Divinylbenzene (80% mixtures of isomers, DVB), 2,2'-azo-bisisobutyronitrile (≥98%, AIBN), 4-vinylbenzenesulfonic acid sodium salt (≥90%), sodium allylsulfonate (≥90%), diisobutyl phthalate (DIBP, ID-1), decahydronaphthalene and methylmagnesium chloride (3 M in tetrahydrofuran solution) were all purchased from Shanghai Aladdin Biochemical Technology Co., Ltd. (Shanghai, China). DVB was pretreated before use according to our previous work.<sup>26,28</sup> Poloxamer 407 (F<sub>127</sub>, BASF), titanium tetrachloride (Tianjing Yongda Chemical Reagent Co. Ltd), 3-methyl-5-tert-butyl-1,2-phenylene dibenzoate (Tianjin Scaxchem Limited Company, ID-2) and ethanol (≥99.5%, Sinopharm Chemical Reagent Co., Ltd.) were used as received. Deionized water, ethylene, nitrogen, toluene, hexane and triethylaluminium (TEAL, 10% in hexane solution) were kindly donated by PetroChina Lanzhou Petrochemical Company.

## 2.2 Sulfonated porous organic polymer and catalysts preparation

Sulfonated porous organic polymers were synthesized through dispersion polymerization as previously reported, utilizing 4vinylbenzenesulfonic acid sodium salt (A) and sodium allylsulfonate (B) as functional comonomers.29 Some of the resulting s-POPs underwent further acidification, while all s-POPs were subjected to vacuum treatment at 80 °C for 8 hours to prepare the catalysts. Under a nitrogen atmosphere, 20 mL of CH<sub>3</sub>MgCl (3 mol L<sup>-1</sup> in THF) was added into a toluene solution containing s-POPs (5 g), stirred at 35 °C for 3 hours. The reaction mixture was then filtered, washed with toluene, and 50 mL of fresh toluene was added. Subsequently, 50 mL of TiCl<sub>4</sub> was added dropwise, and the slurry was heated to 80 °C. Upon reaching the desired temperature, 0.75 g of internal donor (Cat-3 and Cat-4) was added and stirring for 3 hours. Finally, the slurry was filtered, washed sequentially with three portions of toluene and hexane, and dried under vacuum at 80 °C for 1 hour.

## 2.3 Ethylene polymerization in slurry process

Ethylene polymerization was conducted in a nitrogen-replaced 2 L stainless steel reactor. Initially, 300 g of hexane and 5-20 mL of aluminum triethyl (TEAL, 10% in hexane) were introduced into the reactor, stirred for approximately 10 minutes at room temperature to remove the impurities. Following this, a moderate number of catalysts was added, along with an additional 400 g of hexane. Ethylene was continuously supplied under pressure of 0.4-1.1 MPa, while maintaining a temperature of 60-80 °C. Subsequently, the reactor was cooled to room temperature, excess gas vented, and the polyethylene collected through filtration and drying.

#### 2.4 Characterization

The N2 adsorption/desorption isotherms of s-POPs were examined using a Quantachrome Nova 2000e N<sub>2</sub> sorption instrument (Boynton Beach, FL, USA) at 77.3 K. Prior to analysis, the particles were degassed at 120 °C overnight. X-ray diffraction (XRD) measurements of the catalysts were performed on a Bruker D8 ADVANCE using Cu K $\alpha$  radiation ( $\lambda = 0.154$  nm) with  $2\theta$  scanning angle from  $10^{\circ}$  to  $65^{\circ}$ , and samples were protected from contact with air by a polyethylene film. X-ray photoelectron spectroscopy (XPS) analysis utilized an ESCA Lab250 spectrometer from Thermo Fisher Scientific (USA) with Al Kα radiation at 1486.6 eV (500 μm spot-size). The FT-IR of s-POPs and s-POP supported Z-N catalysts were examined on the NEXUS 670 FTIR. Ti and Mg content analysis of the catalysts were carried out on a VISTAICP-MPX (VARIAN, Palo Alto, CA, USA). The viscosity molecular weight  $(M_{\nu})$  of the polymer was measured in decahydronaphthalene at 135 °C following ASTM D-4020. The  $M_{\nu}$  was calculated according to the formula (1).

$$M_{\nu} = 5.37 \times 10^4 [\eta]^{1.37} \tag{1}$$

#### 3. Results and discussion

#### Preparation of sulfonated porous organic polymers

In this study, s-POPs were synthesized via a dispersion polymerization strategy with two functional comonomers. POP-SO<sub>3</sub>Na-A and POP-SO<sub>3</sub>H-A used 4-vinylbenzenesulfonic acid sodium salt (A) as functional comonomer, POP-SO3Na-B and POP-SO<sub>3</sub>H-B used sodium allylsulfonate (B) as functional comonomer, respectively. POP-SO<sub>3</sub>H-A and POP-SO<sub>3</sub>H-B were acidified to convert -SO<sub>3</sub>Na groups to -SO<sub>3</sub>H. Conversely, nonacidified s-POPs (POP-SO<sub>3</sub>Na-A and POP-SO<sub>3</sub>Na-B) retained their original composition. The specific surface area (SSA) and total pore volume (TPV) of obtained s-POPs were assessed by nitrogen sorption analysis. As presented in Table 1, s-POPs exhibited good porosity with the SSA exceeding 300 m<sup>2</sup> g<sup>-1</sup>. In comparison, POP-SO3H-B, employing functional comonomer B, obtained a higher SSA of 471 m<sup>2</sup> g<sup>-1</sup> with a TPV value of 0.31 cm<sup>3</sup> g<sup>-1</sup> compared to POP-SO<sub>3</sub>H-A. This difference could be

Table 1 Porosimetry analysis results of POP-SO<sub>3</sub>H-A and POP-SO<sub>3</sub>H-B

	SSA $(m^2 g^{-1})$	TPV (cm $^3$ g $^{-1}$ )	
POP-SO <sub>3</sub> H-A	385	0.22	
POP-SO <sub>3</sub> H-B	471	0.31	

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attributed to the different thermodynamic compatibility among comonomers, solvent systems and the prepared s-POPs.<sup>27</sup>

#### 3.2 Catalysts preparation

In this work, Mg-Cl moiety was anchored onto the s-POPs via CH<sub>3</sub>MgCl mediated reaction, allowing the modified s-POPs to immobilize TiCl4 and form Ti/Mg@s-POP Z-N catalysts. For Cat-2 and Cat-6, TiCl4 was immobilized onto the Mg-Cl moiety generated from the -SO<sub>3</sub>Na on the surface of non-acidified POP-SO<sub>3</sub>Na-A and POP-SO<sub>3</sub>Na-B. To investigate the effect of internal donor on the catalyst's polymerization performance and the prepared polymers, DIBP (ID-1) and 3-methyl-5-tert-butyl-1,2phenylene dibenzoate (ID-2) were employed as internal donor in Cat-3 and Cat-4, respectively.

The contents of magnesium and titanium loaded in s-POPs were determined by ICP analysis after calcination and acidolysis. As detailed in Table 2, the magnesium contents in the Ti/ Mg@s-POP catalysts were controlled in a relatively narrow range of 3.4-4.2%, which was significantly lower than that of MgCl<sub>2</sub> supported catalysts.31 Moreover, the titanium contents were all controlled below 4%. Catalysts (Cat-2 and Cat-6) derived from unacidified POP-SO<sub>3</sub>Na-A and POP-SO<sub>3</sub>Na-B exhibited lower magnesium contents compared to catalysts prepared from acidified supports. This indicates that the unacidified s-POPs have a weaker interaction with the Grignard reagent, possibly resulting in the physical adsorption of the Grignard reagent onto the s-POP structure. However, the titanium exhibited opposite trend, catalysts derived from unacidified supports showed higher titanium loading content. TiCl<sub>4</sub> probably adsorbed onto the s-POPs via polarized adsorption of -SO<sub>3</sub>Na groups.

#### 3.3 Ethylene polymerization results

The ethylene polymerization was conducted using the six prepared heterogeneous catalysts in conjunction with TEAL, and the polymerization results are presented in Table 2. The Ti/ Mg@s-POP catalytic system was observed to produce polyethylene with  $M_{\nu}$  exceeding 2 million g mol<sup>-1</sup>, making it suitable for producing UHMWPE. Specifically, Cat-1, originating from acidified POP-SO<sub>3</sub>H-A, exhibited high activity of 4966 g PE per g cat per h, resulting in polyethylene with a  $M_v$  of 2.35  $\times$ 10<sup>6</sup> g mol<sup>-1</sup>. In contrast, Cat-2, prepared from unacidified s-POP, showed lower activity of 2677 g PE per g cat per h. Similarly, Cat-5, derived from acidified POP-SO<sub>3</sub>H-B without an internal donor, yielded polyethylene with a  $M_{\nu}$  of 4.28  $\times$  10<sup>6</sup> g mol<sup>-1</sup>, demonstrating the highest polymerization activity of 7356 g PE per g cat per h. However, the unacidified POP-SO<sub>3</sub>Na-B supported catalyst Cat-6 exhibited lower activity but yield polyethylene with a higher molecular weight. This disparity could be attributed to the weaker reaction of CH3MgCl with the unacidified -SO<sub>3</sub>Na groups, leading to most of the CH<sub>3</sub>MgCl and TiCl4 being physically adsorbed onto the s-POP. Consequently, only a small portions of effective Mg-Ti active centers were loaded onto the s-POP and took effect during the ethylene polymerization. Moreover, Cat-5, prepared from POP-SO<sub>3</sub>H-B, exhibited higher polymerization activity and produced polyethylene with a higher molecular weight, suggesting that the choice of functional comonomer impacted the active sites of the prepared catalysts. Compared to traditional MgCl<sub>2</sub> or MgCl<sub>2</sub> (ethoxide type) supported Z-N catalysts, the decrease in activity observed in s-POP based Z-N catalysts for ethylene polymerization is not as pronounced as the decrease observed for propylene polymerization.32

Regarding the results of Cat-3 and Cat-4, catalysts prepared with internal donors exhibited lower activity but higher molecular weight. Cat-3, employing DIBP as the internal donor, produced polyethylene with the highest  $M_{\rm v}$  of 6.83  $\times$  10<sup>6</sup> g mol<sup>-1</sup>, almost three times that of Cat-1 without internal donor. Cat-4 with ID-2 as the internal donor produced polyethylene with a similar  $M_{\nu}$  to Cat-1 but showed lower catalytic activity. Internal donors can bind to the Mg-Ti active centers formed during the preparation process and affect the molecular weight. This may be cause the disruption of active centers responsible for producing low molecular weight polymer by the internal donor, or the reduction in chain transfer to TEAL.16,33

#### 3.4 FTIR, XPS and XRD analysis of catalysts

The FTIR spectra of Ti/Mg@s-POP catalysts are depicted in the Fig. 1. A prominent, wide absorption band near 3200-3700 cm<sup>-1</sup> indicates the presence of -OH stretching vibrations, likely due to water adsorbed in Ti/Mg@s-POP catalysts. Peaks around 2920 cm<sup>-1</sup> correspond to C-H stretching vibrations, aligning with the chemical structure of the supports used. Notably, a sharp absorption peak at approximately  $1600 \, \, \text{cm}^{-1}$ can be ascribed to C=C stretching vibrations in aromatic rings, particularly associated with divinylbenzene. Additionally, bands near 1170 cm<sup>-1</sup> and 1030 cm<sup>-1</sup> are indicative of asymmetric and

Table 2 Preparation and ethylene polymerization results of s-POP supported Z-N catalysts<sup>a</sup>

Catalyst	Support	Internal donor(ID)	Mg (wt%)	Ti (wt%)	Activity (g PE per g cat per h)	$M_{\scriptscriptstyle  m V}  imes 10^{-4}~{ m g~mol}^{-1}$
Cat-1	POP-SO <sub>3</sub> H-A	No ID	4.1	2.8	4966	235
Cat-2	POP-SO <sub>3</sub> Na-A	No ID	3.4	3.2	2677	302
Cat-3	POP-SO <sub>3</sub> H-A	ID-1	3.8	2.7	2397	683
Cat-4	POP-SO <sub>3</sub> H-A	ID-2	4.0	2.0	2208	265
Cat-5	POP-SO <sub>3</sub> H-B	No ID	4.2	2.8	7356	428
Cat-6	POP-SO <sub>3</sub> Na-B	No ID	3.7	3.6	3347	459

<sup>&</sup>lt;sup>a</sup> Polymerization conditions: Cat = 50-60 mg; TEAL = 10 mL; temperature = 70 °C; pressure = 0.6 MPa; time = 60 min.

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2920 cm<sup>-1</sup> 1600 cm<sup>-1</sup> 1030 cm<sup>-1</sup> 1600 cm<sup>-1</sup> 1700 c

Fig. 1 FTIR spectra of Ti/Mg@s-POP catalysts.

symmetric S=O stretching, respectively. These spectral features exhibit variations across different catalyst samples, suggesting alterations in the sulfonic groups' environment, potentially due to interactions with metal centers or other structural changes during the catalyst synthesis on s-POP.<sup>34,35</sup> For samples Cat-3 and Cat-4, additional absorption bands are observed near 1700 cm<sup>-1</sup> and 1300 cm<sup>-1</sup>, corresponding to C=O and C-O stretching vibrations, respectively. These are attributed to the inclusion of ID-1 and ID-2 during the catalyst preparation. Furthermore, a band centered at approximately 580 cm<sup>-1</sup> is identified as C-S stretching vibration.

To investigate the influence of s-POP on the localized microchemical environments of magnesium and titanium sites,

X-ray photoelectron spectroscopy (XPS) analysis was conducted. The obtained spectra, shown in Fig. 2, provide detailed insights into the binding energies (BE) of S, O, Ti, Mg and Cl atoms. As shown in the Fig. 2(a), compared to the binding energy (BE) of the S 2p orbital in pure s-POP (POP-SO<sub>3</sub>H-A), the Ti/Mg@s-POP catalysts display a slight increase in the BE of the S 2p, indicating an electron density decrease around the sulfur atoms in the s-POP. Additionally, as illustrated in the Fig. 2(b), the broadened shape of the O 1s spectra in the s-POP supported catalysts indicates the presence of oxygen in at least three distinct states, contrasting with the narrow peak observed in standalone s-POP at a binding energy around 531.5 eV. The varied peaks within the O 1s range can be attributed to different functional groups: S=O/C=O bonds appearing in the 533.8-532.5 eV range, S-O/C-O bonds around 532.5-531.0 eV, and Ti-O or Mg-O bonds manifesting in the 531.0-529.5 eV range.33 This diversity in oxygen states reflects the complex interactions and bonding environments in the s-POP supported catalysts. Cat-3 and Cat-4, contain similar BE curves around the range of 533.8-532.5 eV, it could be ascribed to the addition of internal donor, which contribute C=O bonds to the prepared catalysts. In contrast, Cat-1 and Cat-5 exhibit distinct metal-oxygen peak profile, reflecting variations in their chemical composition and structure.

Fig. 2(d) illustrates the Ti 2p spectra of the catalysts, characterized by a doublet resulting from spin-orbit coupling of the 2p electrons, corresponding to Ti  $2p_{3/2}$  and Ti  $2p_{1/2}$ 

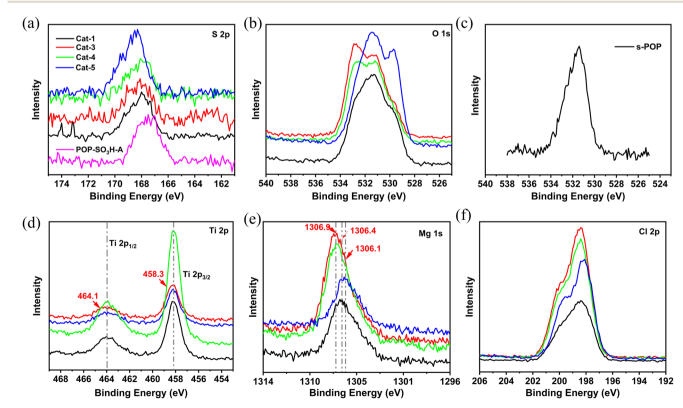


Fig. 2 (a) S 2p XPS spectra of s-POP and s-POP supported catalysts; (b) O 1s XPS spectra of s-POP supported catalysts; (c) O 1s XPS spectra of s-POP; (d) Ti 2p XPS spectra of s-POP supported catalysts; (e) Mg 1s XPS spectra of s-POP supported catalysts; (f) Cl 2p XPS spectra of s-POP supported catalysts.

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Table 3 XPS analysis data of s-POP supported catalysts

Entry	Ti 2p <sub>1/2</sub>	Ti 2p <sub>3/2</sub>	Mg 1s	Cl 2p <sub>1/2</sub>	Cl 2p <sub>3/2</sub>
Cat-1	464.0	458.2	1306.4	198.2	199.7
Cat-3	464.1	458.3	1306.9	198.3	200.0
Cat-4	464.0	458.2	1306.9	198.3	200.0
Cat-5	463.9	458.1	1306.1	198.1	199.7

photoelectrons.<sup>36</sup> The BE values for all supported catalysts are nearly identical. As shown in Table 3, Cat-3 with DIBP as internal donor shows a marginally higher BE of 458.3 eV and 464.1 eV for Ti 2p<sub>3/2</sub> and Ti 2p<sub>1/2</sub>, respectively, potentially due to the enhanced electron-withdrawing effect by the combination DIBP and s-POP. However, the BE for Ti 2p<sub>3/2</sub> of Cat-3 was lower than that reported in the literature in the case of Ti/DIBP/ MgCl<sub>2</sub>.<sup>37</sup> BE shifts are influenced by both the oxidation state and the charge on atoms adjacent to the excited atom.38 In comparison with Cat-1, Cat-5 displays a slightly lower BE for Ti  $2p_{3/2}$  (458.1 eV) and Ti  $2p_{1/2}$  (463.9 eV). This suggests that POP-SO<sub>3</sub>H-B has weaker electron-withdrawing capability than the s-POP employed sodium p-styrene sulfonate as comonomer. In turn, Cat-3 and Cat-4 obtained increased BE for Mg 1s, attributed to the formed Mg-O by the interaction with internal donor. Additionally, POP-SO<sub>3</sub>H-B exhibits the weakest electron withdrawal from the Mg atom, reflected in the smallest Mg 1s BE at 1306.1 eV. Furthermore, Fig. 2(f) reveals a slight increase in the BE of Cl 2p<sub>3/2</sub> for Cat-3 and Cat-4 to 200.0 eV. This increase is likely due to the electron-withdrawing effect from oxygen atoms in the internal donors, indicating a complex interaction between these components within the catalyst structure.

Fig. 3 illustrate the powder X-ray diffraction (XRD) patterns for Ti/Mg@s-POP catalysts alongside anhydrous MgCl<sub>2</sub>. The anhydrous MgCl<sub>2</sub> exhibits distinct and sharp diffraction peaks at  $2\theta$  values of 15°, 30°, 35° and 50°, corresponding to the (003), (012), (104), and (110) crystallographic planes of the  $\alpha$ -MgCl<sub>2</sub> phase, respectively. The XRD patterns of the Ti/Mg@s-POP catalysts mirror the MgCl<sub>2</sub> pattern to some extent, suggesting the presence of MgCl<sub>2</sub> crystals on the s-POP support. However,

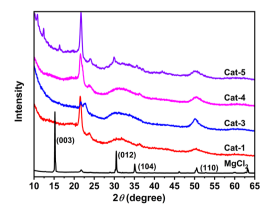


Fig. 3 Powder X-ray diffraction patterns of s-POP based catalysts and MgCl<sub>2</sub>. The peak around  $2\theta=21.7^{\circ}$  is mainly due to the crystal of the polyethylene protection film.

the specific peak for the (003) plane is conspicuously absent, implying that the Cl–Mg–Cl layers within the s-POP based catalysts structure are limited and possibly mainly monolayered. Moreover, the (012) and (104) peaks appear merged into a single broadened peak ( $2\theta$  ranges at 27– $37^{\circ}$ ), combining with the broad reflection at 48– $54^{\circ}$ , indicating the formation of stacking disorder  $\delta$ -MgCl<sub>2</sub> within the s-POP based Z–N catalysts.<sup>39</sup> The diminished intensity of the (110) peak, compared to that of MgCl<sub>2</sub>, likely signifies a reduction in the size of the MgCl<sub>2</sub> crystallites within the catalyst.<sup>40-42</sup>

#### 3.5 Effect of polymerization conditions on the polyethylene

Multiple factors could affect the polymerization activity and the molecular weight of the obtained UHMWPE. The effect of temperature, [Al]: [Ti] molar ratio, hydrogen concentration and ethylene pressure on catalyst activity and the  $M_{\nu}$  of the polyethylene was investigated. For example, while polymerization temperature positively affects activity, it adversely affects polymer molecular weight. As shown in Table 4 and Fig. 4(a), Cat-1 exhibited activity below 3000 g PE per g cat per h at 55 °C, but increased to 6800 g PE per g cat per h at 80 °C, nearly 2.5 times higher than at 55 °C. Within the temperature range of 60-70 °C, polymerization activity remained relatively stable. However, the  $M_{\nu}$  of the resulting polyethylene decreased significantly from  $4.83 \times 10^6$  to  $1.70 \times 10^6$  g mol<sup>-1</sup>, due to an increased chain transfer rate. Alkylaluminum plays a crucial role in olefin polymerization, influencing catalyst behavior and polymer properties as transfer agent. 16,43 In this study, the [Al]: [Ti] molar ratio was consistently maintained between 40 and 294 while keeping the catalyst content constant. An increase in [Al]: [Ti] molar ratio corresponded to higher catalytic activity, ranging from 2964 to 4084 g PE per g cat per h. Moreover, no maximum activity was observed within the studied [Al]: [Ti] range, possibly because both the Ti<sup>3+</sup> and Ti<sup>2+</sup> are active species for ethylene polymerization.44 However, due to chain transfer to TEAL, the viscosity average molecular weight decreased to  $2.05 \times 10^6$  g  $\text{mol}^{-1}$  at a [Al]: [Ti] ratio of 294 (Table 5).

Hydrogen serves as the principal chain transfer agent in the polymerization of all olefins with heterogeneous Ti-based catalysts. <sup>45</sup> As shown in Table 6, the catalyst activity decreased with increasing hydrogen concentration, completely opposite to propylene polymerization. <sup>46,47</sup> Compared to polymerization without hydrogen, the catalyst activity decreased by 2.5 fold

**Table 4** Effect of temperature on polymerization activity of Cat-1 and  $M_{\nu}$ , of the prepared UHMWPE<sup> $\alpha$ </sup>

Entry	PE no.	Temp (°C)	Activity (g PE per g cat per h)	$M_{ m v} imes 10^{-4} \ ({ m g~mol}^{-1})$
1	PE-1	55	2570	102
-				483
2	PE-2	60	4500	342
3	PE-3	65	4740	317
4	PE-4	70	4966	235
5	PE-5	80	6800	170

<sup>&</sup>lt;sup>a</sup> Polymerization conditions: Cat-1 = 50-60 mg; pressure = 0.6 MPa, TEAL = 10 mL, time = 60 min.

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(a) 8000 550 (b) 4200 350 7333 500 4000 325 6667 450 Activity (g PE/g cat) 3800 300 (g PE/g 6000 400 3600 350 275 💍 5333 4667 300 250 ≥ 3400 Activity 4000 250 3200 225 3333 200 3000 200 2667 150 2800 2000 100 70 50 100 150 200 250 300 50 55 60 65 75 85 Temperature (°C) [AI]:[Ti]

(a) Effect of temperature on polymerization results; (b) effect of [Al]: [Ti] molar ratio on polymerization results.

Table 5 Effect of [Al]: [Ti] molar ratio on polymerization activity of Cat-1 and  $M_v$  of the UHMWPE obtained<sup>a</sup>

Entry	PE no.	[Al]:[Ti]	Activity (g PE per g cat per h)	$M_{ m v}  imes 10^{-4} \  m (g\ mol^{-1})$
6	PE-6	40	2964	317
7	PE-7	102	3209	282
8	PE-8	213	3741	237
9	PE-9	294	4080	205

<sup>&</sup>lt;sup>a</sup> Polymerization conditions: Cat-1 = 50-60 mg; temperature = 70 °C; pressure = 0.6 MPa; time = 60 min.

when  $[H_2]$ :  $[C_2H_4] = 0.50$ . Numerous hypotheses have been proposed to explain the reduction in ethylene polymerization activity in the presence of hydrogen, but the mechanism still requires further elucidation. 48,49 Sukulova et al. 50 suggested that this reduction is mainly caused by the decrease in the calculated propagation rate constant. Due to the chain transfer reaction of hydrogen, the molecular weight of the prepared polyethylene sharply decreases. In entry 10, a small amount of hydrogen  $([H_2]: [C_2H_4] = 0.17)$  reduced the  $M_v$  to  $44.5 \times 10^4$  g mol<sup>-1</sup>, nearly 20% of the polyethylene (2.35  $\times$  10<sup>6</sup> g mol<sup>-1</sup>) obtained without hydrogen, and the polyethylene's  $M_v$  was only 9.68  $\times$  $10^4$  g mol<sup>-1</sup> at higher hydrogen concentration ([H<sub>2</sub>]:[C<sub>2</sub>H<sub>4</sub>] = 0.83). It can be concluded that the s-POP-supported ethylene polymerization catalyst exhibit high sensitivity to hydrogen response.

Table 6 Effect of hydrogen and ethylene pressure on polymerization activity of Cat-1 and  $M_v$  of the polyethylene obtained<sup>a</sup>

Entry	PE no.	[C <sub>2</sub> H <sub>4</sub> ] pressure (MPa)		Activity (g PE per g cat per h)	
10	PE-10	0.6	0.17	3672	44.5
11	PE-11	0.6	0.50	2054	13.3
12	PE-12	0.6	0.83	1068	9.68
13	PE-13	1.0	0	15 146	313

<sup>&</sup>lt;sup>a</sup> Polymerization conditions: Cat-1 = 50-60 mg; temperature = 70 °C; TEAL = 10 mL; time = 60 min.

Ethylene polymerization without hydrogen at high pressure was conducted in entry 13. As shown in Table 6, the catalytic activity was two times higher than that of Cat-1 under the pressure of 0.6 MPa. However, in contrast, the molecular weight did not change significantly.

## Conclusion

In conclusion, this study demonstrated the effectiveness of using s-POP supported Ziegler-Natta catalysts for ethylene polymerization. These s-POP-based catalysts successfully produced UHMWPE, with  $M_{\nu}$  reaching up to 6.83  $\times$  10<sup>6</sup> g mol<sup>-1</sup>, particularly with Cat-3 using DIBP as the internal donor. The effects of polymerization temperature, [Al]: [Ti] molar ratio, and ethylene pressure on the polymerization process were thoroughly explored, providing valuable insights for optimizing catalyst performance and polymer characteristics. The s-POP-supported Ziegler-Natta catalysts with the collaborative influence of internal electron donors and supports on the active centers. These functionalized POPs offer a new strategy and a straightforward approach for producing UHMWPE with tailored molecular weights.

# Data availability

All data supporting the findings of this study are included within the manuscript.

# Conflicts of interest

There are no conflicts to declare.

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