RSC Advances



PAPER

View Article Online
View Journal | View Issue



Cite this: RSC Adv., 2025, 15, 5801

Unusual phenomena of ethylene homopolymerization catalyzed by α -diimine nickel emerging in a microflow platform†

Zeen Zhang, Tianjiao Li, Yong Dong and Yangcheng Lu 🗅 *

This study focuses on ethylene homopolymerization catalyzed by α -diimine catalysts within a designed microfluidic system. Distinct from conventional batch reactors, this microfluidic configuration effectively eradicates the reaction environment heterogeneity, thereby guaranteeing homogeneous reaction conditions and decoupling the mass transfer from the reaction process. This unique setup enables precise modulation of ethylene concentration, facilitating the isolation of variables to scrutinize the effects of temperature and pressure on the polymerization. By varying residence times, it's revealed that the reaction rate is ethylene concentration-independent. Variable separation studies between concentration and temperature show an inverse relationship between temperature and branching degree (BD), contrasting traditional batch reactions due to ethylene concentration changes. Pressure studies indicate that higher pressure leads to lower BD. Different product structures' steric hindrances influence the reaction rate; greater hindrance slows it down. These novel findings suggest microreactors offer new perspectives on late transition metal-catalyzed ethylene homopolymerization, potentially enabling breakthroughs in mass production.

Received 23rd January 2025 Accepted 16th February 2025

DOI: 10.1039/d5ra00557d

rsc.li/rsc-advances

1. Introduction

Polyolefin materials are crucial in modern life and industry, widely used in packaging, automotive, *etc.* Polyethylene, the most-produced polyolefin, has chemical stability, good mechanics, and low cost. Its synthesis *via* ethylene homopolymerization is a key area in polymer chemistry.

The development of ethylene polymerization catalysts has been remarkable. Early Ziegler–Natta¹ catalysts initiated industrial polyethylene production but had limited structure regulation. Then early-transition metal metallocene complexes in combination with MAO and non-metallocene early-transition metal complexes were proved useful in the synthesis of polyolefin.²-⁴ Subsequently, late-transition metal catalysts exhibited a range of unique advantages.⁵-7 Ever since Brookhart's pioneering research on α -diimine catalyst,⁴ extensive studies have been carried out on this catalyst system over the past several decades.⁵-10 Owing to its distinctive chain walking mechanism, this particular kind of catalysts is capable of synthesizing polyolefins possessing branched chain architectures as well as diverse polymer topologies.¹1,12

State Key Laboratory of Chemical Engineering, Department of Chemical Engineering, Tsinghua University, Beijing 100084, China. E-mail: luyc@mail.tsinghua.edu.cn

However, traditional ethylene homopolymerization often employs batch reactors, which exhibit significant deficiencies in mass and heat transfer efficiency. During the exothermic reaction, slow heat transfer can lead to uneven local temperatures, creating temperature gradients that degrade reaction uniformity and affect the consistency of polymer properties. Simultaneously, limited mass transfer hinders adequate contact between reactants and catalysts, not only reducing reaction rates but also increasing the likelihood of side reactions due to excessively high local concentrations, making it challenging to meet the stringent requirements for high quality and performance in modern polyolefin production.

The advent of microreactor technology offers innovative ideas and effective solutions to these challenges. Microreactors are characterized by their tiny channel dimensions (typically in the micrometer range), which bestow them with a significantly high specific surface area, resulting in an order-of-magnitude improvement in mass and heat transfer efficiency compared to traditional batch reactors. In microreactors, reactants can achieve rapid mixing and uniform distribution, effectively avoiding local concentration and temperature differences, thus significantly enhancing reaction selectivity and yield. 13,14 Moreover, microreactors allow for precise control of key parameters such as reaction temperature, pressure, and residence time, providing unprecedented accuracy and flexibility for optimizing reaction conditions.15 In radical polymerization reactions, they can precisely control radical concentrations and optimize the molecular weight distribution of the polymers.

[†] Electronic supplementary information (ESI) available: ¹H-NMR spectra of ligand and polymer and additional experimental details and methods. See DOI: https://doi.org/10.1039/d5ra00557d

In the study of ethylene homopolymerization catalyzed by α diimine nickel catalysts, although this catalyst has shown unique potential, the interplay between mass transfer and reaction processes makes it difficult to gain clear insights into the reaction laws, hindering a deeper understanding of this catalytic system and its further application development. To address this issue, we innovatively placed the ethylene homopolymerization catalyzed by α-diimine nickel catalysts in a microreactor platform. Leveraging the characteristics of microreactors, the polymerization process proceeded effectively and smoothly, mostly conforming to common reaction characters. Surprisingly, we noticed some unusual but confirmative phenomena in a certain conditional window, such as, elevated pressure resulting in decreased reaction rate, reflecting the complexity of reaction system. We tried to interpret these phenomena at microscale, and indicated its significance for polymerization process control and design toward product functionalization or high efficiency.

2. Results and discussion

2.1 Reaction performances in a microflow platform

In the microflow system, the amount of ethylene added can be accurately controlled by regulating the flow rate of ethylene gas. Then, the mass of the product is obtained through the weighing method, and further the monomer conversion of ethylene can be calculated, which is difficult to obtain in traditional batch reactions. In this work, toluene was selected as the solvent. As a preliminary experiment, the dissolution of ethylene in toluene was observed in microflow system at first. It was found that ethylene could be sheared into small bubbles in the T-shaped micromixer and then dissolved within a few seconds. This implies that, compared with the traditional batch reactors, the dissolution of ethylene in the microfluidic system is significantly intensified, and it can be ensured that there is no presence of ethylene gas in the reaction tubes, allowing the reaction to proceed in a homogeneous state. Therefore, the influence of the mass transfer process during which ethylene gas dissolves in toluene can be eliminated, and thus the intrinsic kinetics of the reaction may be obtained.

Firstly, the reliability of the polymerization platform was verified through repeated experiments. The experimental conditions are shown in Table 1. The ethylene conversion rates of the three sets of experiments were 46.59%, 45.24% and 43.99%, respectively, with the relative error being less than 3%. Meanwhile, the branching degrees (BD) of polyethylene obtained from the three sets of experiments were 128, 126 and 125, respectively, with the relative error being less than 2%. Judging from the results of both the raw material conversion and the

Table 1 Experimental conditions for repeatability verification experiments

P/MPa	$T/^{\circ}\mathbf{C}$	$c_{\text{ethylene,0}}/\%$ wt	$c_{\rm cat}/\%_{\rm oo}$ wt	$c_{\mathrm{MAO}}/\%$ wt	t/min
0.8	10	0.75	0.14	0.13	10

product structure, the platform has good experimental repeatability and the experimental data are reliable. GPC were also employed to test the molecular weight and its distribution of the polymer. Under the condition (Table 1), the weight-average molecular weight ($M_{\rm W}$) of the polymer was measured to be 7.3 × 10⁴, the number-average molecular weight ($M_{\rm n}$) was 4.3 × 10⁴, and the molecular weight distribution (PD) was 1.7. Compared with the data in the literature, the molecular weight and its distribution are reasonable. Since the focus of this paper is not on the discussion of molecular weight, the exploration of the laws regarding molecular weight will not be carried out in the context.

By changing the reaction time at the same combination of temperature, pressure, ethylene concentration, catalyst concentration and cocatalyst concentration, the time profile of polyethylene yield and ethylene conversion were obtained, as shown in Fig. 1. Fig. 1 also shows the results of linear fitting of X-t. The fitting parameter of correlation coefficient $R^2 = 0.99$ and the fitted straight line passes through the origin. The fitting results are in line with the characteristics of zero-order kinetics. That is, the reaction rate is independent of the ethylene concentration.

Since in traditional batch reactions, the ethylene concentration varies with changes in pressure and temperature, the conclusions regarding the variation of the branching degree (BD) with pressure and temperature discussed in most of the literature are often the result of the coupling of multiple factors, without controlling variables. Moreover, polymerization is a highly exothermic reaction. In batch polymerization, problems such as hot spots often occur, making it rather difficult to control the reaction conditions. By taking advantage of the merits of microreactors, such as the ability to adjust the initial concentration of the system and their good heat transfer performance, the above problems can be solved.

Herein, a series of polymerization experiments in which the ethylene temperature was varied, but the pressure held at 0.8 MPa, is summarized in Table 2 and Fig. 2. The temperature range for the experiment was selected from 0 °C to 20 °C. Within this temperature range, during the reaction time of 10

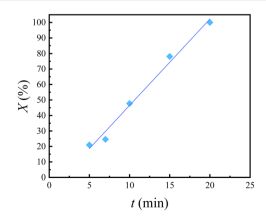


Fig. 1 Conversion of $C_2H_4\ \textit{versus}$ of time refer to entries 1 to 5 in Table 2.

Table 2 Summary of reaction condition and results

Entry	P/MPa	T/°C	$c_{\rm ethylene,0}$ /% wt	$c_{\rm cat}/^{\circ}_{\circ oo}$ wt	$c_{\mathrm{MAO}}/\%$ wt	t/min	Conversion/%	BD
1	0.8	10	0.75	0.14	0.13	5	19.89	125
2	0.8	10	0.75	0.14	0.13	7	23.34	130
3	0.8	10	0.75	0.14	0.13	10	45.24	126
4	0.8	10	0.75	0.14	0.13	15	74.06	124
5	0.8	10	0.75	0.14	0.13	20	100	115
6	0.8	0	0.75	0.14	0.13	10	31.91	150
7	0.8	5	0.75	0.14	0.13	10	42.24	132
8	0.8	15	0.75	0.14	0.13	10	52.27	113
9	0.8	20	0.75	0.14	0.13	10	67.16	100
10	0.5	10	0.75	0.14	0.13	10	85.39	138
11	1	10	0.75	0.14	0.13	10	9.42	121
12	0.5	10	0.5	0.14	0.13	10	60.94	247
13	0.8	10	0.5	0.14	0.13	10	76.85	189
14	1	10	0.5	0.14	0.13	10	83.67	157

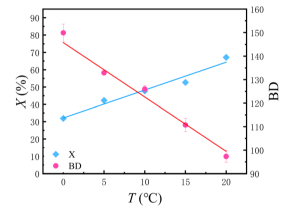


Fig. 2 Conversion of C_2H_4 and BD *versus* temperature refer to entries 3, 6–9 in Table 2.

minutes, the activity of the catalyst can be maintained relatively well, which means there will be no significant decrease in the reaction rate or abnormal variations in BD due to catalyst deactivation.

It can be seen from the Fig. 2 that under the same reaction time, the ethylene conversion increases linearly with the increase of reaction temperature. In other words, in the case of ensuring that the catalyst does not lose activity, increasing the temperature is conducive to accelerating the reaction rate.

As for the BD, it can be seen from the Fig. 2 that BD shows a linear decreasing trend with the increase in temperature. This is contrary to the conclusions obtained in many traditional batch reactors. Derek P. Gates $et\ al.^{16}$ once made speculations on the mechanism of ethylene polymerization catalyzed by α -diimine nickel catalysts: basically, the chain walking takes place in the cationic alkyl complex (B). As the ethylene concentration rises, the lifetime of this complex is reduced, and consequently the degree of chain walking is also decreased. Our experiment provided further evidence for this mechanism conjecture. As the temperature increases, the solubility of ethylene in toluene decreases. For traditional batch polymerization, the concentration of ethylene in the solution is exactly the solubility of

ethylene in toluene under the corresponding temperature and pressure. That is to say, with the rising temperature, the concentration of ethylene drops, which is the main reason why the BD increases with the increase in temperature during batch polymerization. However, in the flow platform, the concentration of ethylene is controlled by a gas mass flowmeter, which can ensure that the ethylene concentration remains equal under different temperature conditions. Therefore, the experimental data can reflect the real impact of temperature. As the temperature rises, the movement rate of molecules accelerates and the rate of capturing ethylene increases, resulting in the decrease of BD (Fig. 3).

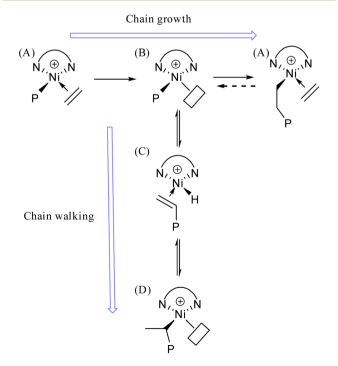


Fig. 3 Proposed mechanism of propagation for the preparation of branched polyethylene.

2.2 Effect of system pressure

A series of polymerization experiments in which the system pressure was varied from 0.5 MPa to 1 MPa, but the temperature held at 10 °C, is summarized in Table 2 and Fig. 4.

As shown in Fig. 4, two sets of experiments with different ethylene concentrations simultaneously demonstrated that as the pressure of the reaction system increases, the BD decreases. Regarding the conversion, at lower ethylene concentrations, the conversion increases with an increase in pressure. However, at higher ethylene concentrations, the conversion shows an opposite trend.

As for the changes in BD, with the elevation of pressure, the interaction among molecules in the liquid is enhanced and the distance between molecules becomes smaller, which makes it easier for the catalyst to capture ethylene. As a result, the chain growth rate is accelerated relative to the chain walking rate, thus leading to the reduction of BD.

As for the changes in the conversion, that is, the reaction rate, under different pressures, the two sets of experiments showed completely opposite trends. Under a higher ethylene concentration, the reaction rate decreases as the pressure increases; while under a lower ethylene concentration, the reaction rate increases with the increase in pressure. We believe that just like the steric hindrance effect of the catalyst ligand, the steric hindrance caused by the change in product structure can also significantly affect the reaction rate. When the steric hindrance is relatively large, it becomes difficult to capture ethylene and the reaction rate slows down.

Under the conditions of lower pressure and lower ethylene concentration, the degree of branching of polyethylene increases, and when the pressure is relatively low, the BD of the polymer increases sharply. According to the free volume theory, the volume of a polymer material consists of two parts: one part is the volume occupied by the molecules themselves, which is called the occupied volume; the other part is the voids between molecules, which is called the free volume. Branched polymers have a larger occupied volume and a smaller free volume compared with linear polymers, resulting in an increase in the steric hindrance of polyethylene and an enhanced shielding effect on the metal active center. When the pressure is relatively

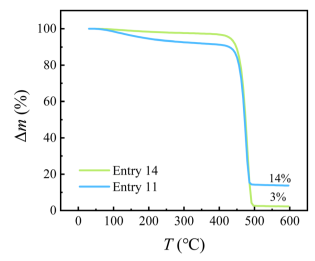


Fig. 5 The TGA results refer to entries 11 and 14 in Table 2.

high and the concentration is also high, the polymer tends to form a straight chain. Under the action of the polymer shape and pressure, physical cross-linking occurs in the polymer, which further leads to an increase in steric hindrance and a decrease in the ethylene capture rate.

The cross-linking of polyethylene can significantly increase its thermogravimetric char yield. After the cross-linking treatment of polyethylene materials, their molecular structures become more complex and stable, which enables more carbon atoms to be retained to form char during high-temperature decomposition. Therefore, cross-linked polyethylene is superior to non-cross-linked polyethylene in terms of thermal stability, and its thermogravimetric char yield is significantly increased. By comparing the TGA results of ethylene concentrations of 0.50% wt and 0.75% wt respectively at 1 MPa under the same reaction temperature, catalyst and cocatalyst concentrations, it was found that the char yield was only 3% when the concentration was 0.50% wt, while it was as high as 14% when the concentration was 0.75% wt. This indicates that the polyethylene produced by polymerization at an ethylene concentration of 0.75% wt has more cross-linking, which is consistent with the reasons we speculated previously (Fig. 5).

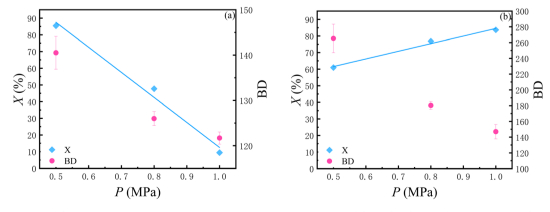


Fig. 4 (a) Conversion of C_2H_4 and BD *versus* pressure with the initial concentration of C_2H_4 0.75% wt refer to entries 3, 10, 11 in Table 2. (b) Conversion of C_2H_4 and BD *versus* pressure with the initial concentration of C_2H_4 0.5% wt refer to entries 12–14 in Table 2.

. . . .

Paper

3. Conclusions

In summary, a microfluidic system was designed to accomplish the ethylene homopolymerization catalyzed by α -diimine catalysts. It enabled to eliminate the impacts brought about by the heterogeneity of the reaction environment in conventional batch reactors, and achieve uniform reaction conditions. It is also capable of separating the mass transfer process from the reaction process, which makes it possible to study the intrinsic kinetics. Moreover, the concentration of the fed ethylene can be precisely controlled to separate variables and explore the influences of factors such as temperature and pressure.

Through the study of polymerization reactions with different residence times, it was found that the reaction rate is independent of the ethylene concentration. Through the study of the separation of variables between concentration and temperature, it was discovered that an increase in temperature would lead to a decrease in the branching degree (BD), which is contrary to the conclusion of traditional batch reactions. The difference come from that the traditional batch reactions were deviated from the change in ethylene concentration meanwhile. Through the study of the separation of variables between concentration and pressure, it was found that the higher the pressure, the smaller the branching degree. Different product structures will bring about different steric hindrances, which in turn affect the reaction rate. The greater the steric hindrance, the stronger the shielding of the active catalytic sites and the slower the reaction rate. These unusual phenomena gave us a reminder that new platform like microreactor might provide new insights into ethylene homopolymerization catalyzed by late transition metal and potential breakthroughs in mass production.

4. Experimental sections

4.1 Materials

Ethylene (C_2H_4 , $\geq 99.9\%$, Air Liquide (Tianjin) Co., Ltd), methylaluminoxane (MAO) (10% wt solution in toluene, Rhawn), acenaphthenequinone ($C_{12}H_6O_2$, 98%, Macklin), 2,6-diisopropylaniline ($C_{12}H_{19}N$, 95%, Macklin), acetonitrile (C_2H_3N , $\geq 99.9\%$, extra dry, water ≤ 30 ppm, J&K), acetic acid ($C_2H_4O_2$, 99.7%, Aladdin), hexane (C_6H_{14} , 97.5%, extra dry, water ≤ 50 ppm, J&K), acetone (C_3H_6O , 99.5%, Tongguang) and nickel(II) bromide ethylene glycol dimethyl ether complex ((DME)NiBr₂, 97%, Adamas) were used directly without further purification. Dichloromethane ($C_4C_2C_1$, 99.9%, extra dry with molecular sieves water ≤ 50 ppm, J&K) and toluene ($C_6H_5C_4$, $\geq 99.9\%$, Tongguang) were purged with nitrogen and then dehydrated by adding molecular sieves with a volume fraction of 10%. Deuterated chloroform (CDC_1 , (C_1 , 99.8%)) were purchased from Adamas and was used directly as received.

4.2 Equipment and procedures

4.2.1 Synthesis of ligand ([*N***-(2,6-diisopropylphenel)imino**] **acenaphthene**). This ligand was prepared according to a method reported by Asselt, Elsevier *et al.*¹⁷

Acenaphthenequinone (1.35 g, 7.4 mmol) was heated under reflux in 65 mL of acetonitrile at 80 °C for 30 minutes. Subsequently, 12 mL of acetic acid was added, and heating continued until the anthraquinone was completely dissolved. To this hot solution, 3 mL (16 mmol) of 2,6-diisopropylaniline was directly added, and the mixture was further heated under reflux for 1.5 hours. The solution was then allowed to cool to room temperature, and the solid was filtered to obtain a yellow product, which was washed with hexane and air-dried.

4.2.2 Synthesis of catalyst (bis[N,N'-(2,6-diisopropylphenyl) imino]acenaphthene nickel(n) dibromide). The catalyst was prepared according to the method reported by Brookhart $et\ al.^6$ 500 mg (1.62 mmol) of (DME)NiBr $_2$ and 850 mg (1.70 mmol) of ligand were mixed under an argon atmosphere. To this mixture, 30 mL of dichloromethane was added, resulting in an orange solution. The mixture was stirred for 20 hours, yielding a reddish-brown solution. The liquid was then added to n-hexane for anti-solvent precipitation, producing a reddish-brown solid. The solid was washed with n-hexane and dried under vacuum.

4.2.3 Stop-flow method. Fig. 6 shows the schematic diagram of the microflow system for stop-flow method. Ethylene gas was introduced into the system through a mass flow controller (MFC, FlowMethod Measure & Control Systems Co. Ltd; flow rate, 0-50 sccm, operating differential pressure, 0.5-1 MPa; maximum allowable pressure, 5.0 MPa). A syringe pump (Chemyx Fusion6000 with 50 mL syringe; linear force, 500 lbs; flow rate, 0.19 μL min⁻¹ to 120 mL min⁻¹) was used to deliver toluene. After pre-cooling to the specified temperature, the twophase reactants were mixed in a T-shape micromixer (i.d. = 1.2mm), where ethylene can be sheared into small bubbles and dissolved quickly. To ensure sufficient dissolution process, a delay microtube (L = 3.0 m; i.d. = 1.0 mm) was connected to the T-shape micromixer. A toluene solution of catalyst and cocatalyst MAO was also added to the system via a syringe pump and mixed with toluene dissolved with ethylene. The reaction begins when the two solutions are mixed. When the flow is stable, close the valves at both ends of the reaction coil and start timing. The reaction was terminated automatically due to the deactivation of the catalyst in contact with air when opening the back valve to receive the sample at preset time. A bypass line was set up to keep the liquid flowing, preventing clogging of the line. The outlets of the reaction tubes were connected to a stainless buffer vessel as surplus solution collector. A back pressure regulator (pressure range, 0-3 MPa, Beijing Xiongchuan Technology Co., Ltd) was also connected to the buffer vessel to control the system pressure. The reaction mixture was collected and dropped into excessive acetone. After centrifugation, the products could be completely isolated from the mixture and was vacuum dried at 80 °C for 24 h.

To compare batch polymerization and flow polymerization, we conducted comparative experiments under the same conditions. The experimental conditions and results are shown in Table 3. For batch polymerization, the experimental method we adopted was as follows: ethylene was continuously introduced into an autoclave filled with a toluene solution containing the catalyst and cocatalyst (which had been pre-pressurized

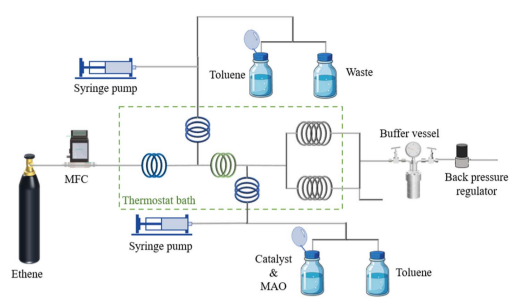


Fig. 6 Micro-flow system.

Table 3 The comparison of batch and flow reactor

Mode	P/MPa	t/min	T/°C	$c_{\rm cat}/^{\circ}_{\sim o}$ wt	$c_{\rm MAO}/\%$ wt	$m_{ m ethylene}/{ m g}$	$m_{\rm poly}/{\rm g}$	Conversion/%
Batch Flow	0.1	1.2	20	0.14	0.13	0.00514 0.3396	0.00235 0.00204	45.70 0.59

with nitrogen to the specified pressure), and the mixture was continuously stirred. The time when ethylene started to be introduced was regarded as the starting time of the reaction. When the reaction ended, the ethylene feed was shut off and the vent valve was opened to terminate the reaction. The amount of ethylene introduced was the sum of the ethylene dissolved in toluene (estimated by the solubility) and the ethylene in the headspace (estimated by the ideal gas equation of state). Judging from the data, the mass transfer process indeed has a significant impact on the rate of the entire polymerization reaction. Microreactors, however, can separate mass transfer from the reaction, enabling the exploration of the intrinsic reaction laws. Meanwhile, by taking advantage of their enhanced mass transfer, microreactors can improve the apparent reaction rate.

4.3 Characterization

4.3.1 NMR spectroscopy. The sample for proton nuclear magnetic resonance (1 H-NMR) was prepared by dissolving 2–3 mg polymer into 0.5 mL CDCl $_3$ at 65 °C. All NMR spectra were recorded on Bruker, AVANCE III HD400 at room temperature. Chemical shifts were referenced to residual CDCl $_3$ as an internal standard and given in ppm.

Branching degree (BD) is represented by the number of methyl groups per thousand carbon atoms ($M_{\rm e}/1000$ C). The $M_{\rm e}/1000$ C values for different polyethylene were calculated based on the ratio of the integral area of the peak attributed to the

hydrogen atoms in the methyl groups to the integral area of the peaks attributed to all hydrogen atoms in the ¹H-NMR spectra. The calculation formula¹⁸ shows as follows.

$$M_{\rm e}/1000~{\rm C} = \frac{A_1}{A_1 + A_2} \times \frac{2}{3} \times 1000$$

Among them, A_1 represents the integral area of the peak assigned to the hydrogen in the methyl group in the ¹H-NMR, and A_2 represents the integral area of the peak assigned to the hydrogen in the methylene and methine groups.

4.3.2 TGA. Thermogravimetric measurements were carried out using the TGA (METTLER TOLEDO, Thermal Analysis System TGA/DSC 3+). Samples of approximately 3–5 mg were loaded into an alumina crucible and placed in the TGA furnace. The temperature was ramped from room temperature to 600 °C at a heating rate of 10 °C min⁻¹ under nitrogen atmosphere with a flow rate of 20 mL min⁻¹. During the heating process, the mass change of the sample was continuously monitored and recorded as a function of temperature.

4.3.3 GPC. Number-average molecular weight (M_n) , weight-average molecular weight (M_W) , and molecular weight distribution (PD) of polymers were measured by gel permeation chromatography (GPC). GPC was performed on PL-GPC220 (Agilent) at 150 °C with standard polystyrene as the reference, and 1,2,4-trichlorobenzene was employed as the eluent with a flow rate of 1.0 mL min⁻¹.

Data availability

The data that support the findings of this study are available from the corresponding author upon reasonable request.

Author contributions

Zeen Zhang: writing original draft, conceptualization, methodology, investigation, data curation; Tianjiao Li: investigation, validation; Yong Dong: investigation, data curation; Yangcheng Lu: supervision, conceptualization, methodology, project administration.

Conflicts of interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

Acknowledgements

We appreciate financial support from the National Natural Science Foundation of China (21422603, U1662120, 21978152).

References

- 1 K. Soga and T. Shiono, Prog. Polym. Sci., 1997, 22, 1503-1546.
- 2 A. Andresen, H.-G. Cordes, J. Herwig, W. Kaminsky, A. Merck, R. Mottweiler, J. Pein, H. Sinn and H.-J. Vollmer, *Angew Chem. Int. Ed. Engl.*, 1976, 15, 630–632.
- 3 L. S. Boffa and B. M. Novak, Chem. Rev., 2000, 100, 1479– 1494.

- 4 M. Mitani, T. Nakano and T. Fujita, *Chem.-Eur. J.*, 2003, **9**, 2396–2403.
- 5 E. Drent, R. van Dijk, R. van Ginkel, B. van Oort and R. I. Pugh, *Chem. Commun.*, 2002, 744–745, DOI: 10.1039/B111252J.
- 6 L. K. Johnson, C. M. Killian and M. Brookhart, J. Am. Chem. Soc., 1995, 117, 6414–6415.
- 7 T. R. Younkin, E. F. Connor, J. I. Henderson, S. K. Friedrich, R. H. Grubbs and D. A. Bansleben, *Science*, 2000, 287, 460– 462
- 8 L. K. Johnson, S. Mecking and M. Brookhart, J. Am. Chem. Soc., 1996, 118, 267–268.
- 9 J. M. Kaiser and B. K. Long, *Coord. Chem. Rev.*, 2018, 372, 141–152
- 10 F. Wang and C. Chen, Polym. Chem., 2019, 10, 2354-2369.
- 11 L. H. Shultz, D. J. Tempel and M. Brookhart, *J. Am. Chem. Soc.*, 2001, **123**, 11539–11555.
- 12 L. Guo, S. Dai, X. Sui and C. Chen, *ACS Catal.*, 2016, **6**, 428–441.
- 13 L. Falk and J. M. Commenge, *Chem. Eng. Sci.*, 2010, **65**, 405-411.
- 14 M. N. Kashid and L. Kiwi-Minsker, *Ind. Eng. Chem. Res.*, 2009, 48, 6465–6485.
- 15 S. A. Kazemi Oskooei and D. Sinton, *Lab Chip*, 2010, **10**, 1732–1734.
- 16 D. P. Gates, S. A. Svejda, E. Oñate, C. M. Killian, L. K. Johnson, P. S. White and M. Brookhart, *Macromolecules*, 2000, 33, 2320–2334.
- 17 R. van Asselt, C. J. Elsevier, W. J. J. Smeets, A. L. Spek and R. Benedix, *Recl. Trav. Chim. Pays-Bas*, 1994, **113**, 88–98.
- 18 A. C. Gottfried and M. Brookhart, *Macromolecules*, 2003, 36, 3085–3100.