



Cite this: *React. Chem. Eng.*, 2025, 10, 2238

Received 29th May 2025,
 Accepted 17th August 2025

DOI: 10.1039/d5re00239g

rsc.li/reaction-engineering

Short reaction times for hydrogenolysis of polyolefins by overcoming mass transfer limitations

E. van Daatselaar, ^a A. G. J. van der Ham,^a S. R. A. Kersten^a and M. P. Ruiz ^{*ab}

The recycling of polyolefins is gaining attention as society transitions toward a more circular economy. Pyrolysis is a promising method; however, its product distribution can be unpredictable. Moreover, the resulting compounds often require additional hydrogenation if they are to be used as feedstock for naphtha crackers. An alternative approach is hydrogenolysis, in which polyolefins are depolymerised into shorter, fully saturated alkanes using a heterogeneous catalyst under a hydrogen atmosphere. Literature indicates that the hydrogenolysis of polyolefins appears to be a slow process, requiring reaction times up to 96 hours to achieve a significant yield of useful products, such as naphtha or fuels. In this work, it is shown that these long reaction times are resolved when physical mass transport limitations are overcome: in 40 minutes, full conversion of low-density polyethylene to gas and liquid products is reached. Introducing a hollow-shaft mechanical stirrer instead of no or limited stirring significantly increases the gas contact area and mass transfer coefficient to the polymer melt, resulting in a decrease in mass transport limitations and thus an increase in overall reactivity. Monitoring the (hydrogen) pressure over time generates more insight into the reaction kinetics, as at a similar hydrogen consumption level, the product distribution changes if the system is stirred instead of kept stagnant. The authors would like to emphasise the importance of these findings regarding the influence of hydrogen mass transfer through the melt, as this could also result in novel catalysts possibly performing even better than currently reported, making hydrogenolysis a more viable option for the chemical recycling of polyolefins.

In recent years, an increasing number of scientific articles have been published researching the application of hydrogenolysis as a novel chemical recycling technique for the conversion of

polyolefins, such as polyethylene and polypropylene, to smaller alkanes used as fuels or as feedstock for the existing plastic production process.^{1–9} Published work focuses mainly on the development of highly active catalysts and the research on reaction mechanisms, to be able to obtain full conversion and, preferably, high yields in the liquid and wax range. This mixture with compounds in the range of pentane (C₅H₁₂) to eicosane (C₂₀H₄₂), is the desired product, as the lower range is similar to the naphtha feedstock for hydrocrackers, while the higher range equals fuel in the gasoline or diesel range. However, current reported reaction times to reach this product distribution are substantial, ranging from 3 to 96 hours.^{2,3,7,10} Reaction times must be significantly reduced to develop an economically feasible chemical recycling process of polyolefin waste into fuels using hydrogenolysis.

Current literature often neglects the influence of, amongst others, feedstock, and reactor kinetics and configuration.^{11–13} As the scission of the polyolefins depends on the presence of hydrogen at the catalyst surface in the liquid phase,¹⁰ we prove in this work that mass transfer of hydrogen is of high importance to be able to quickly hydrogenolyse polyolefins.

In published work, hydrogenolysis of polyolefins is often performed with a monofunctional ruthenium on carbon (Ru/C) catalyst, either as the main focus or as a benchmark experiment. In Table 1, an overview of several studies that apply similar reaction conditions, such as temperature and initial hydrogen pressure, is given. The reactor volumes are comparable, ranging from 25 to 190 mL.^{2,5} The main difference is the level of stirring: either not at all, with a magnetic bar, or a mechanical impeller. As can be seen in Table 1, to achieve full conversion of the polymer to gases and liquids using Ru/C, reaction times of up to 24 hours are required. To enable a consistent comparison of the literature data, the required hydrogen consumption in moles was calculated from the reported product distributions, extracted from either figures or tables. Complete saturation of all products was assumed. Per mole of converted polyolefin, this corresponds to a hydrogen usage of $x - 1$ moles of H₂, where

^a Sustainable Process Technology, University of Twente, PO Box 217, 7500 AE Enschede, The Netherlands. E-mail: m.p.ruizramiro@utwente.nl

^b Circular Chemical Engineering, Maastricht University, 6167 RD Geleen, The Netherlands



Table 1 Overview of publications performing hydrogenolysis of polyolefins over a monofunctional ruthenium catalyst. The hydrogen consumption rates are calculated based on reported data, assumptions can be found in section S1

Feed (MW)	Catalyst ^a	Type of stirring	Stirring rate [rpm]	Ratio F : C	Mass feed [g]	Temp. [°C]	Initial H ₂ press. (STP) [bar]	Reaction time [h]	Gas yield (C1–C4) [wt%]	Liquid yield (C5–C45) [wt%]	Solid yield (C45+) [wt%]	H ₂ cons. rate (calc. & norm.) [mol _{H₂} g _{Ru} ^{−1} s ^{−1}]	Source
LDPE (4 kDa)	Ru/C	No stirring	—	10 : 1	1.0	250	30	18	100	0	0	1.9×10^{-4}	1
LDPE (4 kDa)	Ru/C	Magnetic	600	25 : 1	1.4	200	22	16	57	43	0	2.8×10^{-4}	2
LDPE (4 kDa)	Ru/C	Magnetic	600	4 : 1	0.1	225	30	16	100	0	0	7.6×10^{-5}	2
i-PP (250 kDa)	Ru/C	Magnetic, 0.7 mL stir bar	300	20 : 1	2.0	250	30	16	97	3	0	3.8×10^{-4}	3
LDPE (~100 kDa)	Ru/CeO ₂ ^b	Magnetic	400	4 : 1	2.0	250	20	24	13	87	0	3.0×10^{-4}	4
LDPE (4 kDa)	Ru/C	Magnetic, glass coated stir bar	450	34 : 1	3.4	240	35	6	23	39	38	2.7×10^{-4}	5
HDPE (200 kDa)	Ru/TiO ₂	Mechanical, solid shaft	750	20 : 1	0.5	225	20	4	60 ^c	14 ^c	26	4.8×10^{-4d}	6
LDPE (4 kDa)	Ru/C	Mechanical, hollow shaft	1300	9.3 : 1	3.5	250	40	0.67	22	78	0	1.4×10^{-3}	This work

^a 5 wt% Ru, unless noted otherwise. ^b 0.2 wt% Ru. ^c Gas range reported as sum of C1–C5, liquid range starts at C6. ^d Based on reported reaction rate.

$x - 1$ equals the total number of moles of alkane products formed. This approach ensures closure of both the carbon and hydrogen balances. The required amount of moles H₂ to obtain the given product distributions is normalised over the amount of ruthenium and averaged over the reaction time (mol_{H₂} g_{Ru}^{−1} s^{−1}). The assumptions for these calculations can be found in the SI, section S1.

This work reevaluates the same reaction system of LDPE (MW 4 kDa) and 5 wt% Ru/C, while applying a novel reactor configuration. As hydrogen is required to hydrogenolyse the polyolefins, a higher concentration of hydrogen in the polymer melt near the catalyst surface should enhance the reaction rate. For that reason, a hollow shaft stirrer is applied, which physically introduces hydrogen bubbles in the liquid phase, increasing the contact area and thus the mass transfer of hydrogen from the gas phase to the liquid phase (Fig. 1 and S5). During operation of this batch-wise system, the pressure and temperature are logged over the entire reaction time, resulting in information about the reactivity of the chemical system. More details on the experimental method, reactor set-up and components, all designed and built in-house, can be found in the SI (section S2).

As the reaction progresses, the measured pressure in the reactor is a combination of the hydrogen and product partial pressures in the gas phase, *i.e.* $P_{\text{tot}} = P_{\text{H}_2, \text{G}} + P_{\text{C}_x, \text{G}}$. A pressure change is thus a direct result of both consumption of H₂ and the formation of gaseous hydrocarbons (C_x). The concentration of H₂ in the liquid is proportional to the concentration (and thus partial pressure) of hydrogen in the gas phase *via* the solubility ($C_{\text{H}_2, \text{L}} = m \times C_{\text{H}_2, \text{G}}$), the volumetric mass transfer coefficient $k_L a$ and consumption during reaction. Due to the mixing shown in Fig. 1 and S5, the value for $k_L a$ is high. Measurements and calculations in section S3

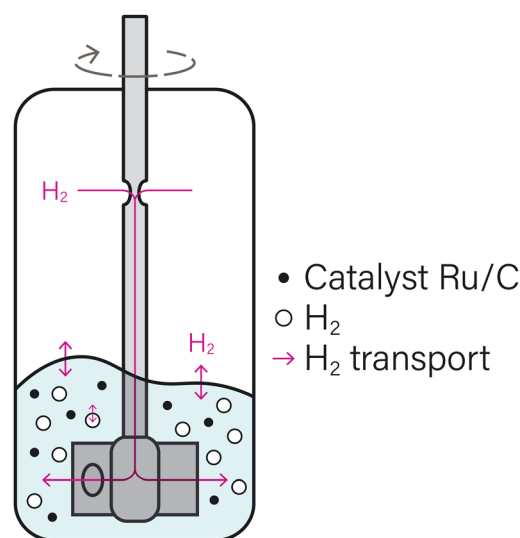


Fig. 1 Schematic overview of the hollow-shaft stirred autoclave. Due to rotation, hydrogen gas is transported through the shaft into the polymer melt, increasing the gas transport area through bubble formation.

Reaction Chemistry & Engineering

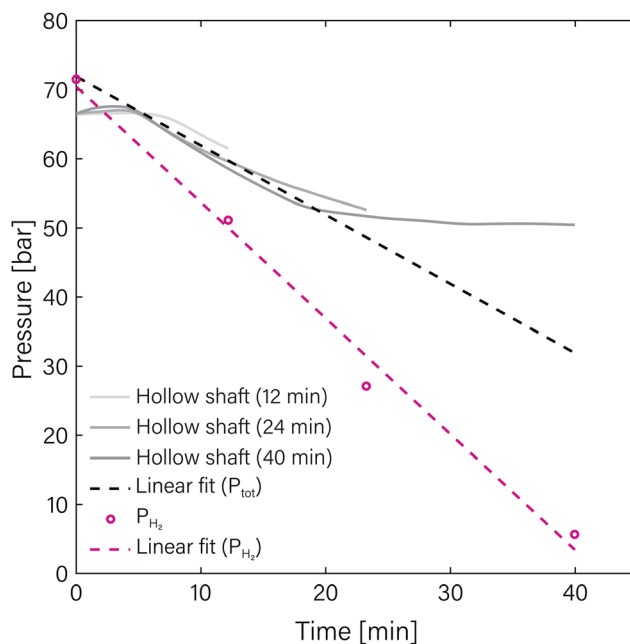


Fig. 3 Pressure over time for hollow-shaft stirred experiments for different reaction times (grey). Pink dots indicate final partial hydrogen pressure. Linear fits are shown (dashed) to extract rates. Experimental settings: 3.5 g LDPE, Ru/C, feed-to-catalyst 9.3 to 1, 40 bar H₂ (at RT), initial peak temperature 260 + 5 °C. after 15 min 250 + 5 °C.

Performing batch reactions with LDPE (4 kDa) and 5 wt% Ru/C at a feed-to-catalyst ratio 9.3 to 1, 250 °C and an initial pressure 40 bar H₂ (at RT) for 40 minutes at reaction temperature, while changing the stirring regime, returns pressure profiles as shown in Fig. 2. This raw data shows a difference in reaction rate, as no stirring results in nearly no pressure decrease and thus almost no polymer conversion. Stirring with a solid shaft at different speeds to mimic recent work using magnetic or mechanical stirrers does increase the conversion rate minimally, as the slope steepens slightly. However, applying the hollow shaft stirrer shows a significant pressure drop, indicating much more conversion within the same time range.

Calculating the rates for the experiments with no or limited stirring results in similar values for the normalised hydrogen consumption compared to data shown in Table 1,^{1,5,6} namely 1.48×10^{-4} and 2.78×10^{-4} mol_{H₂} g_{Ru}⁻¹ s⁻¹, respectively. These calculations can be found in the SI, section S4. More importantly, compared to the performed non-stirred experiment, the consumption rate of the hollow-shaft stirred system is approximately 10 times faster (Table S5), *i.e.* this finding significantly reduces the required reaction times to obtain liquid yields *via* hydrogenolysis. Analysis of the final product substantiates this statement, as the product is fully converted to alkanes in the range of C1 to C28 within the reaction time of 40 minutes – see Fig. 5.

As previously mentioned, the measured pressure is a combination of hydrogen consumption and gaseous product formation. Therefore, the resulting product distribution was expected to resemble when experiments reach a comparable final pressure. The pressure profiles of a non-stirred and hollow-shaft stirred experiment, conducted for 420 and 24 minutes, respectively, are presented in Fig. 4 together with the earlier shown 40 minute experiment for comparison. An identical final reactor pressure is reached within a $\pm 0.5\%$ margin. However, comparing the resulting product yields in Fig. 5 reveals unexpected results: the enhanced mass transfer influences the product distribution.

As can be seen in Fig. 5, not stirring the reaction mixture does lead to the formation of lower molecular weight alkanes. The methane yield of 15 wt% is notably higher compared to the methane yield of the hollow-shaft stirred reaction mixtures, while the formation of other gases remains

- 3 P. A. Kots, S. Liu, B. C. Vance, C. Wang, J. D. Sheehan and D. G. Vlachos, *ACS Catal.*, 2021, **11**, 8104–8115.
- 4 M. Chu, X. Wang, X. Wang, X. Lou, C. Zhang, M. Cao, L. Wang, Y. Li, S. Liu, T.-K. Sham, Q. Zhang and J. Chen, *Research*, 2023, **6**, 1–10.
- 5 Y. Nakaji, M. Tamura, S. Miyaoka, S. Kumagai, M. Tanji, Y. Nakagawa, T. Yoshioka and K. Tomishige, *Appl. Catal., B*, 2021, **285**, 119805.
- 6 S. D. Jaydev, A. J. Martin, D. Garcia, K. Chikri and J. Perez-Ramirez, *Nat. Chem. Eng.*, 2024, **1**, 565–575.
- 7 G. Celik, R. M. Kennedy, R. A. Hackler, M. Ferrandon, A. Tennakoon, S. Patnaik, A. M. LaPointe, S. C. Ammal, A. Heyden, F. A. Perras, M. Pruski, S. L. Scott, K. R. Poepplmeier, A. D. Sadow and M. Delferro, *ACS Cent. Sci.*, 2019, **5**, 1795–1803.
- 8 S. D. Jaydev, A. J. Martin and J. Perez-Ramirez, *ChemSusChem*, 2021, 1–8.
- 9 Z. Qiu, S. Lin, Z. Chen, A. Chen, Y. Zhou, X. Cao, Y. Wang and B. L. Lin, *Sci. Adv.*, 2023, **9**, 1–10.
- 10 Y. Nakagawa, S. I. Oya, D. Kanno, Y. Nakaji, M. Tamura and K. Tomishige, *ChemSusChem*, 2017, **10**, 189–198.
- 11 P. A. Kots, B. C. Vance and D. G. Vlachos, *React. Chem. Eng.*, 2022, **7**, 41–54.
- 12 R. Helmer and M. Shetty, *Chem Catal.*, 2023, **3**, 1–2.
- 13 M. L. Pennel, Y. Jiang and M. Cargnello, *ACS Sustainable Resour. Manage.*, 2024, **1**, 1047–1052.
- 14 A. Buffo, M. Vanni and D. Marchisio, *Chem. Eng. Sci.*, 2012, **70**, 31–44.
- 15 M. N. Amar, F. M. Alqahtani, H. Djema, K. Ourabah and M. Ghasemi, *J. Taiwan Inst. Chem. Eng.*, 2023, **153**, 105215.
- 16 M.-R. Mohammadi, F. Hadavimoghaddam, M. Pourmahdi, S. Atashrouz, M. T. Munir, A. Hemmati-Sarapardeh, A. H. Mosavi and A. Mohaddespour, *Sci. Rep.*, 2021, **11**, 17911.

