Hydrogenolysis of Polyethylene and Polypropylene into Propane over Cobalt-Based Catalysts

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ABSTRACT: The development of technologies to recycle polyethylene (PE) and polypropylene (PP), globally the two most produced polymers, is critical to increase plastic circularity. Here, we show that 5 wt % cobalt supported on ZSM-5 zeolite catalyzes the solvent-free hydrogenolysis of PE and PP into propane with weight-based selectivity in the gas phase over 80 wt % after 20 h at 523 K and 40 bar H₂. This catalyst significantly reduces the formation of undesired CH₄ (≤5 wt %), a product which is favored when using bulk cobalt oxide or cobalt nanoparticles supported on other carriers (selectivity ≤95 wt %). The superior performance of Co/ZSM-5 is attributed to the stabilization of dispersed oxidic cobalt nanoparticles by the zeolite support, preventing further reduction to metallic species that appear to catalyze CH₄ generation. While ZSM-5 is also active for propane formation at 523 K, the presence of Co promotes stability and selectivity. After optimizing the metal loading, it was demonstrated that 10 wt % Co/ZSM-5 can selectively catalyze the hydrogenolysis of low-density PE (LDPE), mixtures of LDPE and PP, as well as postconsumer PE, showcasing the effectiveness of this technology to upcycle realistic plastic waste. Cobalt supported on zeolites FAU, MOR, and BEA were also effective catalysts for C₇−C₉ hydrocarbon formation and revealed that the framework topology provides a handle to tune gas-phase selectivity.

KEYWORDS: polyolefin waste, chemical recycling, cobalt, ZSM-5, propane, plastic circularity, selective hydrogenolysis

Plastic waste represents one of the most pressing problems of modern society, as over 5000 million tonnes have been landfilled or leaked into the environment since the 1950s.¹−⁴ Polyethylene (PE) and polypropylene (PP), the two most produced plastics worldwide, account for over half of this waste by weight.⁵−⁶ While mechanical recycling can process only a fraction of plastic waste, chemical recycling could convert it into processable products that can be reintegrated into a circular polymer economy.⁷−⁹ However, a key challenge lies in activating the strong and inert C–C bonds of the PE and PP backbones.

Among the different strategies reported to this end, hydrogenolysis,⁹−¹⁸ which utilizes H₂ to cleave the polymer into shorter alkanes, is a promising approach to deconstruct PE and PP under moderate conditions. Notably, platinum- and ruthenium-based catalysts have been shown to convert model polymers and postconsumer polyolefin plastics into liquid alkanes (C₇−C₂₀).¹⁹−²⁴ To date, these catalytic systems often require high loadings of precious metals (up to 6 wt %) and typically produce a wide array of products ranging from CH₄ and other light alkanes to liquid paraffins (C₇−C₂₄) and solid waxes (C₂₅+), which stems from the internal C–C bond cleavage mechanism favored in these systems.²⁰−²⁴ The development of catalysts that are not based on precious metals and that promote narrower product distributions would enable more efficient technologies to valorize plastic waste. This would require catalysts adept at selectively cleaving the C–C bond positions in the polyolefin chain that are nonterminal, thus minimizing generation of undesired CH₄, but near the chain end, forming a limited range of hydrocarbons, such as ethane, propane, and butane. These could be readily converted into the corresponding olefins and H₂ by commercial, scalable dehydrogenation technologies,²⁹−³¹ thereby regenerating the monomers and closing both the carbon and hydrogen loop in PE and PP production (Figure 1).

To identify a catalyst with the aforementioned properties and reactivity, the solvent-free hydrogenolysis of n-tetracosane (n-C₂₄H₄₀), an oligomer of PE, was first investigated in a batch reactor, as discussed in detail in the Supporting Information.

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Among the different bulk metal oxides screened, cobalt oxide (CoO) was found to be highly active, fully converting the paraffin substrate into almost exclusively CH$_4$ (Table S1). Given these results, CoO was tested for the batch solvent-free hydrogenolysis of PE and PP with mass average molecular weights ($M_w$) of 4000 Da and 12 000 Da, respectively (Figures S1 and S2). After 20 h at 523 K and 40 bar of H$_2$ both polymers could be fully converted into CH$_4$ over 100 mg of CoO. The dominant products across temperatures, reaction times, and catalyst loadings tested over CoO were solids or gases, with gaseous products composed exclusively of CH$_4$. Under all conditions investigated, only trace liquid products were formed (Figures S1 and S2). These observations support the hypothesis that Co$_3$O$_4$ favors a terminal C–C bond cleavage mechanism.

Support interactions have been shown to strongly affect the activity and selectivity of C–C bond cleavage over cobalt-based systems in different reactions. Thus, we dispersed 5 wt % cobalt over various supports, i.e., amorphous silica–alumina (SIRAL), silica (SiO$_2$), ceria (CeO$_2$), zirconia (ZrO$_2$), titania (TiO$_2$; anatase phase), and ZSM-5 zeolite (Si/Al = 11.5) via incipient wetness impregnation. The catalysts were characterized by temperature-programmed reduction with H$_2$ (H$_2$-TPR), exhibiting at least two peaks at ca. 533 and 603 K, similar to bulk Co$_3$O$_4$ (Figure S3), which are characteristic of Co$^{3+}$ and Co$^{2+}$ reduction, respectively. Notable exceptions to this trend were Co/SIRAL and Co/ZSM-5, both showing a single peak at 533 and 568 K, respectively. The catalysts were then tested in the batch solvent-free hydrogenolysis of model PE ($M_w = 4000$ Da) using an equivalent mass of cobalt across the experiments (Figure 2), where $W_M$ represents the mass of Co metal. The catalytic activity was measured by quantifying the H$_2$ conversion, $X_{H_2}$, which provides an estimate of the numbers of C–C bonds cleaved. After 20 h at 523 K and 40 bar H$_2$ differences were observed in the catalyst performance. In summary, the catalytic activity increased as Co/SIRAL < Co/SiO$_2$ < Co/CeO$_2$ < Co/ZrO$_2$ < Co/TiO$_2$ ≈ Co/ZSM-5 (Figure 2a). The reactivity hierarchy was well reflected by the phase distribution of products, with Co/SIRAL, Co/SiO$_2$, Co/CeO$_2$, Co/ZrO$_2$, and Co/TiO$_2$ predominantly resulting in solids (85–90%), while Co/ZSM-5 mainly yielded gaseous products (87%). Still, the H$_2$ conversion over Co/ZSM-5 was significantly lower than that obtained over CoO under equivalent conditions (30% vs 60%).

The weight-based selectivity distributions in the gas phase (eq 3 in the SI) were compared at $X_{H_2} \approx 5\%$ (Figure 2b), obtained by varying the reaction time. This allows for a more direct performance comparison at an approximately equal number of C–C bonds cleaved. Co/ZrO$_2$ exhibited a gas-phase composition virtually comparable to that of bulk Co$_3$O$_4$, Co/TiO$_2$, and especially Co/SIRAL (Figure S4). However, the total gas yield was limited to approximately 15%, and by increasing the H$_2$ conversion, the amount of CH$_4$ rose quickly at the expense of other alkanes (Figure 2b). Interestingly, at reduced H$_2$ conversions ($\leq$1%) as obtained after 20 h, a considerably higher fraction of C$_2$H$_6$ and other light alkanes (60–90 wt %) were detected over Co/SiO$_2$ and especially Co/SIRAL (Figure S4). However, the total gas yield was limited to approximately 15%, and by increasing the H$_2$ conversion, the amount of CH$_4$ rose quickly at the expense of other alkanes (Figure 2b). Notably, Co/ZSM-5 was the only system that did not produce CH$_4$ and instead favored the generation of C$_2$H$_6$ (77%), C$_3$H$_8$ (12%), and C$_4$H$_{10}$ (8%) at $X_{H_2} \approx 5\%$ (Figure S5). Under these conditions, almost 30% of the products comprised liquid hydrocarbons (primarily C$_3$)
with an iso/normal paraffin ratio of ca. 5:1, while higher alkanes were significantly less abundant (Figure S5). The liquid distribution observed for Co/ZSM-5 strongly differs from that obtained over archetypical Ru/C that peaks at around C\textsubscript{4}\textsubscript{5}H\textsubscript{11} \textsuperscript{20}. On the contrary, it qualitatively resembled that obtained over other cobalt-based catalysts, i.e., Co/SiO\textsubscript{2}, Co/Co\textsubscript{2}O\textsubscript{4}, and Co/SIRAL (Figure S6), although these systems generated considerably less liquids (<5 wt %). These results also significantly differ from reports of the hydropyrolysis and hydrocracking of PE over cobalt-based catalysts including CoAl\textsubscript{2}O\textsubscript{4} and Co/MOR at temperatures ranging from 623 to 823 K, which produce gasoline-range hydrocarbons.\textsuperscript{39,30}

The batch hydrogenolysis of n-tetracosane was also investigated over Co/ZSM-5. After 2 h, around 70% of C\textsubscript{24}H\textsubscript{50} and 10% of H\textsubscript{2} were converted into an almost equal mass mixture of gases, liquids, and solids, resembling the product distribution obtained after 5 h in PE hydrogenolysis at similar (8%) H\textsubscript{2} conversion (Figures S5 and S7). Notably, the gas phase was predominantly composed of propane (76%), and the distribution of liquids also resembled that obtained after 5 h in PE hydrogenolysis, demonstrating the replicability of our results in a model system.

To probe catalyst stability, we performed a recycling experiment of PE (M\textsubscript{w} = 4000) hydrogenolysis over Co/ZSM-5. The spent catalyst from hydrogenolysis (20 h, 523 K, 40 bar H\textsubscript{2}) was extracted, washed with cyclohexane, dried at 393 K for 12 h, and then reloaded in the Parr reactor with fresh polymer and run under the same reaction conditions. The recycled catalyst exhibited minimal change in activity, maintaining propane as the major gaseous product with only a slight decrease in X\textsubscript{h\textsubscript{2}} (Figure S8). To further demonstrate the stability of the Co/ZSM-5 system, we also ran vapor-phase hydrogenolysis of n-dodecane in continuous flow, which maintained stable activity and selectivity for at least 24 h on stream (Figure S9a,b). We estimated this system to have a turnover number (TON) of at least 2000 under these conditions. This value fits the criteria of Kozuch and Martin, which state that an industrially relevant catalyst should have a TON \( \geq 1000 \).\textsuperscript{30}

By progressively varying the reaction time from 5 to 80 h, the H\textsubscript{2} conversion over Co/ZSM-5 increased from ca. 7% to 42% along with the gas fraction (up to ca. 94%) (Figure 2c and Figure S5). Additionally, CH\textsubscript{4} weight-based selectivity in the gas phase rose with reaction time from 1% to 12% at the expense of C\textsubscript{2}H\textsubscript{6} production, suggesting that CH\textsubscript{4} formation is kinetically limited. The cobalt-based catalysts were also tested in the batch solvent-free hydrogenolysis of model PP (M\textsubscript{w} = 12 000 Da) (Figure S10). A comparable activity hierarchy to that in PE hydrogenolysis was observed when deconstructing PP: Co/SIRAL < Co/SiO\textsubscript{2} < Co/Co\textsubscript{2}O\textsubscript{4} < Co/ZrO\textsubscript{2} < Co/TiO\textsubscript{2} \ll Co/ZSM-5. Contrastingly, the H\textsubscript{2} conversions were lower, and the fraction of solids was higher when compared to those obtained for PE hydrogenolysis over the same systems, in agreement with the previous results on Co\textsubscript{3}O\textsubscript{4} (Figure S10). However, a similar product distribution to that in PE hydrogenolysis was observed, with Co/ZSM-5 favoring C\textsubscript{2}H\textsubscript{6} formation (ca. 80%), while other systems preferably yielded CH\textsubscript{4}.

Co/ZSM-5, Co/SiO\textsubscript{2}, Co/SIRAL, and the corresponding bare supports, were further characterized by temperature-programmed desorption of NH\textsubscript{3} (Figure S11). While Co/SiO\textsubscript{2} is not acidic, Co/ZSM-5 and Co/SIRAL showed two similar main peaks for NH\textsubscript{3} desorption (423 K, 778 K vs 393 K, 723 K), indicating qualitatively similar acidic strength, although differing in total quantity of sites (0.87 mmol NH\textsubscript{3} g\textsubscript{cat} \textsuperscript{-1} vs 0.04 μmol NH\textsubscript{3} g\textsubscript{cat} \textsuperscript{-1}, respectively). No differences were observed between the NH\textsubscript{3}-TPD profiles of the metal-containing catalysts and the bare supports, with the exception of a 100 K upward shift of the high-temperature desorption peak in Co/ZSM-5 and of a new peak at 723 K over Co/
SIRAL, which might be related to the ability of cobalt to coordinate with NH₃.⁴¹ These results, in combination with the analysis of iso/normal alkane distribution in the liquid phase during PE hydrogenolysis, provide important mechanistic insights. Specifically, over nonacidic catalysts, such as Co/SiO₂, normal C₄₅ alkanes were the only liquid products detected in PE hydrogenolysis. In contrast, systems with acid supports like SIRAL and ZSM-5 generated a distribution of iso- and n-alkanes in the liquid phase (Figure S6). After 5 h of PE hydrogenolysis, Co/ZSM-5 generated primarily C₅₃ iso-alkanes, indicating the occurrence of isomerization. Accordingly, tertiary carbon could be formed even in the case of PE,⁴² which would explain the similar product distribution observed for PP. These insights point to a possible bifunctional hydrocracking mechanism, also known as ideal hydrocracking,⁴³ where both metal-based phases and acid sites are involved.

Co/ZSM-5 was further analyzed by transmission electron microscopy (TEM) prior to and after PE hydrogenolysis. The average particle diameters before and after reaction were 1.50 ± 0.32 and 1.66 ± 0.40 nm, respectively, demonstrating minimal change in particle size (Figure 3a,b and Figures S12 and S13). The minor decrease in activity exhibited by the recycled catalyst (Figure S8) could be due to the slight sintering of nanoparticles after the reaction. The choice of the support has been shown to significantly influence the oxidation state of cobalt, allowing the stabilization of either metallic or partially reduced phases in H₂-rich atmospheres.⁴⁴ It has been shown that metal confinement within zeolites can stabilize metastable phases,⁴⁵–⁴⁷ which could offer a possible explanation for the pronounced stabilization effect with ZSM-5. Interestingly, Have et al. demonstrated that H₂ activation routes differ depending on the cobalt oxidation state in CO₂-based Fischer–Tropsch synthesis. In particular, they showed that while H₂ is adsorbed dissociatively over metallic cobalt, cobalt oxide follows a H₂-assisted pathway.⁴⁸ In addition to the potential influence of acidity as previously discussed, the observed shift from main terminal to close-to-terminal C=C bond cleavage in polyolefin hydrogenolysis over Co/ZSM-5 might stem from the preferential stabilization of oxidized cobalt phases by the zeolite that are different from those observed for bulk Co₃O₄ or other supports under reductive conditions.

To test this hypothesis, we conducted ex situ H₂-TPR as well as X-ray absorption spectroscopy (XAS) at the Co K-edge of Co/ZSM-5 and Co₃O₄ after PE hydrogenolysis and compared it to the corresponding data in fresh form (Figure 3c,d and Figure S14). Analysis of the X-ray absorption near-edge structure (XANES) revealed that, while fresh Co₃O₄ featured the expected mixed Co²⁺ and Co³⁺ oxidation states, the white line and edge position of the Co/ZSM-5 spectrum resemble those of highly dispersed Co²⁺ species (Figure S14).⁴⁹,⁵⁰ Notably, the H₂-TPR profile of the spent Co/ZSM-5 showed new peaks at 603 and 828 K (Figure 3c), indicating the presence of species that are less reducible. These data are consistent with the XANES profiles of spent Co/ZSM-5 showing the preservation of Co²⁺ species, as well as the FT-EXAFS analysis that showed unchanged local coordination environments after reaction, with the first peak centered at 1.61 Å that can be assigned to Co–O by EXAFS analysis (Figures S14–S16 and Table S2). Still, XAS cannot clearly differentiate between Co–O bonds of oxides and Co–C bonds of carbidic or oxycarbides, which might form under the reaction conditions studied here. During continuous-flow n-dodecane hydrogenolysis, a 4 h induction period was observed where conversion increased from ca. 30% to 46%, and CH₄ production leveled down from 12% to 2% (Figure S9b). These results suggest that the state of the catalyst is evolving during hydrogenolysis, possibly forming carbide-like species that can be stabilized in MFI frameworks under hydrocarbon-rich reductive environments and that have been shown in Fischer–Tropsch synthesis to be active and to reduce CH₄ formation.⁵¹,⁵² Furthermore, differences in intensity and broadening of the pre-edge XANES region were observed between the spectra of fresh and used Co/ZSM-5 (Figure S17), suggesting changes in the coordination geometry,⁵³ which could be related to catalyst evolution. Contrarily, there was extensive reduction of the CoO₂ during PE hydrogenolysis, as observed by a >100 K downward shift in the maxima of the H₂-TPR profile and an order of magnitude lower H₂ consumption (Figure 3c) as well as by XAS, which detected almost exclusively metallic cobalt species with the main peak centered at 2.15 Å (not phase corrected) ascribed to Co–Co pairs (Figure 3d and Figures S14–S18). Finally, TEM and XAS were also conducted on Co/SiO₂ prior to and after PE hydrogenolysis. These analyses revealed that this catalyst was composed of large clusters (d∞ ≈ 17 nm) of CoO₂ that reduced to CoO-like species upon reaction (Figures S15–S17, S19, and S20). These results suggest that the stabilization of highly dispersed oxidic cobalt nanoparticles by the zeolite prevents complete reduction to metallic species and/or formation of large clusters, which may be related to the observed promotion of C₅H₈ and suppression of CH₄. These results exemplify the complexity of this reaction network, which calls for dedicated and systematic full studies that can properly deconvolute acid- from metal-based bond activation, as well as mass transport and confinement effects, among others.

Thermogravimetric analysis (TGA) of Co/ZSM-5 and Co/SiO₂ was conducted prior to and after PE hydrogenolysis to probe the presence of coke, since the latter is not directly observable by TEM and XAS. No differences were observed between the TGA curves of fresh and used Co/ZSM-5, indicating the absence of any carbonaceous residues (Figure S21a). Conversely, almost 50% weight loss was observed in the profile of spent Co/SiO₂ (Figure S21b), which was attributed mainly to the catalytic being coated by solid long chain alkanes rather than coke, as they were the main product obtained over this system (Figure 2a).

ZSM-5-supported cobalt catalysts with different metal loadings (10, 15, and 20 wt %) were further synthesized via incipient wetness impregnation and compared against 5 wt % Co/ZSM-5 in PE hydrogenolysis at equivalent temperature, pressure, time, and cobalt loading (Figure 4a). In particular, while H₂ conversion remained similar over both Co/ZSM-5 and 10-Co/ZSM-5 (~29%), it increased over 15-Co/ZSM-5 (32%) and further over 20-Co/ZSM-5 (38%). While the gas fraction remained roughly constant over the catalysts at ca. 87%, differences were observed in the remaining products. Solids were the main byproducts over Co/ZSM-5 and 10-Co/ZSM-5, while liquids, primarily composed of pentane, were observed over 15-Co/ZSM-5 and 20-Co/ZSM-5. However, the most striking difference was found in the composition of the gaseous components; namely, while Co/ZSM-5 and 10-Co/ZSM-5 maintained high weight-based selectivity in the gas phase toward C₃H₆ (~84%), this value decreased over 15-Co/
ZSM-5 (66%) and especially 20-Co/ZSM-5 (47%) at the conversion and (b) CH$_4$ consumption (Figure S24). The similar morphology and reducibility of the 5 and 10 wt % loaded systems are consistent with their analogous catalytic performance. Conversely, 15-Co/ZSM-5 and particularly 20-Co/ZSM-5 showed large ~30 nm clusters deposited on the external surface of the zeolite together with small nanoparticles dispersed in the microporous structure (Figures S25 and S26), resulting in broad particle size distributions of 4.67 ± 3.30 nm over 15-Co/ZSM-5 and 8.01 ± 5.32 nm over 20-Co/ZSM-5. Their H$_2$-TPR profiles exhibited a main peak at 578 K, similar to those observed over Co/ZSM-5 and 10-Co/ZSM-5 that were attributed to the presence of small nanoparticles stabilized by the zeolite, as well as shoulders at 533 and 603 K likely associated with large clusters of Co$_3$O$_4$ (Figure S24). The area of the combined shoulders represented over 15% and 30% of the total H$_2$ consumption of 15-Co/ZSM-5 and 20-Co/ZSM-5, respectively, which would explain both the increased reactivity and the shift toward CH$_4$ formation. The dependency of the C$_3$H$_8$ and CH$_4$ weight-based selectivity in the gas phase on the average particle size, $d_p$, is depicted in Figure 4b. The generation of C$_3$H$_8$ and CH$_4$ followed a volcano and antivolcano behavior with a maximum at 84% and minimum at 1%, respectively, both centered around 2.22 nm obtained over 10-Co/ZSM-5.

While the ZSM-5 modifies the activity of the cobalt oxide, the zeolite itself is also active for propane formation from PE under these conditions. To ensure that observed trends in Figure 4a,b were not a function of the loading of ZSM-5, which decreases as the cobalt loading increases, the activities over equivalent masses of ZSM-5 for the series were measured for PE hydrogenolysis. The results, shown in Figure 4c and Figure S22b, indicate that ZSM-5 alone has high activity for propane formation, although the product selectivity and trends differ in a number of ways. First, the total quantities of gas and liquid produced, as well selectivity to methane, generally increase with increasing loading of ZSM-5, as opposed to Figure 4b which shows a maximum of propane selectivity at an intermediate cobalt loading. This confirms that the selectivity trends in Figure 4b are not an artifact of the decreasing ZSM-5 content with increasing cobalt loading. Second, the liquid products differ between Co/ZSM-5 and ZSM-5. The liquid products over ZSM-5 are largely aromatic and unsaturated hydrocarbons, as evidenced by GC-MS (Figure S27) and by the yellow color of the products (Figure S28), whereas the liquid products over Co/ZSM-5 also contain saturated hydrocarbons and are translucent in color (Figures S27 and S28). This suggests that the cobalt facilitates H$_2$ activation to saturate hydrocarbons. Third, unlike the reaction over Co/ZSM-5, in which $X_{CH_4}$ increases with increasing gas and liquid produced, the $X_{CH_4}$ over ZSM-5 remains fairly constant. This suggests that there is hydrogen going toward C–C bond cleavage that comes not only from the molecular H$_2$ supplied but also from the substrate itself. The evidence of aromatic products is consistent with the theory that hydrogen transfer reactions from the PE substrate are occurring over ZSM-5.

To further probe the role of hydrogen transfer reactions, control reactions of PE ($M_w = 4000$) over 5-Co/ZSM-5 and ZSM-5 were performed in a nitrogen atmosphere at 523 K. To gain insights into this phenomenon, we performed TEM and H$_2$-TPR of these systems in fresh form. The micrographs of 10-Co/ZSM-5 evidenced well-dispersed particles with a relatively narrow particle size distribution centered around 2.22 ± 0.46 nm (Figure S23), which agreed well with TEM measurements of similarly synthesized 10-Co/ZSM-5 from the literature. The H$_2$-TPR profile of 10-Co/ZSM-5 showed a main peak at 578 K, comparable to Co/ZSM-5 (5 wt %), with a small shoulder at 533 K, accounting for only 5% of the total H$_2$ consumption (Figure S24). The similar morphology and reducibility of the 5 and 10 wt % loaded systems are consistent with their analogous catalytic performance. Conversely, 15-Co/ZSM-5 and particularly 20-Co/ZSM-5 showed large ~30 nm clusters deposited on the external surface of the zeolite together with small nanoparticles dispersed in the microporous structure (Figures S25 and S26), resulting in broad particle size distributions of 4.67 ± 3.30 nm over 15-Co/ZSM-5 and 8.01 ± 5.32 nm over 20-Co/ZSM-5. Their H$_2$-TPR profiles exhibited a main peak at 578 K, similar to those observed over Co/ZSM-5 and 10-Co/ZSM-5 that were attributed to the presence of small nanoparticles stabilized by the zeolite, as well as shoulders at 533 and 603 K likely associated with large clusters of Co$_3$O$_4$ (Figure S24). The area of the combined shoulders represented over 15% and 30% of the total H$_2$ consumption of 15-Co/ZSM-5 and 20-Co/ZSM-5, respectively, which would explain both the increased reactivity and the shift toward CH$_4$ formation. The dependency of the C$_3$H$_8$ and CH$_4$ weight-based selectivity in the gas phase on the average particle size, $d_p$, is depicted in Figure 4b. The generation of C$_3$H$_8$ and CH$_4$ followed a volcano and antivolcano behavior with a maximum at 84% and minimum at 1%, respectively, both centered around 2.22 nm obtained over 10-Co/ZSM-5.

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To further probe the role of hydrogen transfer reactions, control reactions of PE ($M_w = 4000$) over 5-Co/ZSM-5 and ZSM-5 were performed in a nitrogen atmosphere at 523 K.
In N\textsubscript{2}, both reactions produced gaseous and liquid products, largely composed of light hydrocarbons and a mixture of aromatics. TGA of the ZSM-5 before and after reaction in H\textsubscript{2} (Figure S21c) revealed a small additional weight loss feature, which could be attributed to soft coke, that is not present over used 5-Co/ZSM-5. In a nitrogen atmosphere, both 5-Co/ZSM-5 and ZSM-5 exhibited coking after reaction (Figure S21d,e). The stability of the ZSM-5 was also tested in a flow reactor for the vapor-phase hydrogenolysis of \textit{n}-dodecane at 523 K over 24 h on stream (Figure S9c,d), exhibiting a slight deactivation over time, which could be due to the soft coke formation in the absence of cobalt.

The most selective bifunctional catalyst, 10-Co/ZSM-5, was selected for testing the solvent-free hydrogenolysis of polyolefins representative of plastic waste: low-density PE (LDPE), \textit{PP} (\textit{M}_\text{w} = 12,000 Da), a weight-equivalent mixture of LDPE and PP, as well as a postconsumer PE bottle (post-PE) from VWR International (Figure S), as described in the Supporting Information. High H\textsubscript{2} conversion (\(X_{H_2} \geq 35\%\)) was reached in LDPE and post-PE hydrogenolysis over 10-Co/ZSM-5, leading to gases (ca. 95%). On the contrary, \(X_{H_2}\) decreased to 25% and 18% in the hydrogenolysis of the LDPE+PP mixture and PP, resulting in a higher solid fraction (ca. 30% and 60%, respectively), compared to LDPE alone and post-PE (<3%; Figure 5a), probably due to the lower reactivity of PP. In all cases, liquids were limited to \(<5\%\) wt % and were composed primarily of C\textsubscript{3}, as observed in PE hydrogenolysis over Co/ZSM-5. C\textsubscript{3}H\textsubscript{8} was preferably formed in all tests, reaching ca. 80% weight-based selectivity in the gas phase (Figure 5b). Together with the tests on \(n\text{-C}_{25}H_{50}\) hydrogenolysis, these results showcase that the favored propane formation in the gas phase can be obtained over a wide range of substrates.

Finally, we examined the effect of different zeolite topologies on the catalytic activity and selectivity for PE depolymerization. Catalysts containing 5 wt % Co were synthesized via incipient wetness impregnation using three additional zeolite supports: FAU (Si/Al = 15), BEA (Si/Al = 12.5), and MOR (Si/Al = 10). Catalyst characterizations for these materials, including XRD, H\textsubscript{2}-TPR, and NH\textsubscript{3}-TPD, are shown in Figure S32, which confirms that the materials are crystalline and that the cobalt oxide domains have relatively similar reducibility. PE reactions were carried out at 523 K, under 40 bar H\textsubscript{2}, over 20 h (Figure 6). As shown in Figure 6a, catalyst activity varies significantly, with \(X_{H_2}\) increasing from Co/MOR < Co/BEA < Co/FAU < Co/ZSM-5. While Co/FAU, Co/BEA, and Co/MOR produced similar quantities of gas, the liquid composition of Co/FAU and Co/BEA was significantly higher (38% and 51%, respectively) compared to 3% and 4% over ZSM-5 and MOR, respectively. The liquid product distributions (comprising C\textsubscript{8}–C\textsubscript{18}) saturated and unsaturated hydro-

**Figure 5.** Mass-based (a) phase distribution of products and molar H\textsubscript{2} conversion and (b) gas composition in the hydrogenolysis of LDPE, PP, LDPE+PP, and post-PE over 10-Co/ZSM-5. Conditions: \(T = 523\) K, \(P = 40\) bar H\textsubscript{2} (35 bar H\textsubscript{2} for post-PE), \(W_M = 50\) mg, \(\tau = 40\) h (50 h for post-PE).

**Figure 6.** Mass-based (a) phase distribution of products and molar H\textsubscript{2} conversion and (b) gas composition in the hydrogenolysis of PE over different zeolite-supported cobalt-based catalysts. Conditions: \(T = 523\) K, \(P = 40\) bar H\textsubscript{2}, \(W_M = 50\) mg, \(\tau = 20\) h.
The catalytic tests were performed in a 25 cm³ stainless steel Parr reactor setup. Briefly, the solid alkanes (weight, \( W = 0.7 \) g)—\( n\)-C\(_{24}\)H\(_{50}\) polyethylene (PE, mass average molecular weight, \( M_w = 4000 \) Da), polypropylene (PP, \( M_w = 12 000 \) Da), low-density PE (LDPE; melt flow index, MFI, 25 g in 10 min at 463 K and 2.16 kg), and postconsumer LDPE (post-LDPE)—were loaded in the reactor together with the catalyst in an appropriate amount to ensure that, across different catalytic tests, the same amount of Co metal was used (mass metal equivalent, \( W_M = 25–100 \) mg). The reactor was then evacuated, flushed, and pressurized (\( P = 40 \) bar) with \( H_2 \). The reactor was then placed in a homemade heater equipped with a Digi-Sense TC9100 temperature controller and a K-type thermocouple to regulate the operating temperature (\( T = 523 \) K), insulated, and run for different reaction times (\( \tau = 5–80 \) h). After the tests, the reactor was quenched in an ice bath. The gaseous alkane products and \( H_2 \) in the headspace were collected in a gas sampling bag and were quantified off-line using a gas chromatograph (GC) coupled to a flame ionization detector (FID) and a thermal conductivity detector. Liquid alkane products (\( C_{12}, C_{13}, C_{14} \)) were extracted from the remaining solid alkanes (\( C_{13}, C_{14} \)) and catalyst using cyclohexane or acetone as solvent. Then, 1,3,5-tri-tertbutyl benzene was added as an external standard, and the suspension was centrifuged to separate liquids and solids. The liquid products were identified using LC coupled to a mass spectrometer and then quantified off-line using a gas chromatograph (GC) coupled to a quadrupole mass spectrometer (QMS).

Catalyst synthesis, reactivity testing, additional figures, and data of further catalytic experiments and catalyst characterization (PDF)
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