Polyethylene Hydrogenolysis at Mild Conditions over Ruthenium on Tungstated Zirconia

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ABSTRACT:
Plastics waste has become a major environmental threat, with polyethylene being one of the most produced and hardest to recycle plastics. Hydrogenolysis is potentially the most viable catalytic technology for recycling. Ruthenium (Ru) is one of the most active hydrogenolysis catalysts but yields too much methane. Here we introduce ruthenium supported on tungstated zirconia (Ru-WZr) for hydrogenolysis of low-density polyethylene (LDPE). We show that the Ru-WZr catalysts suppress methane formation and produce a product distribution in the diesel and wax/lubricant base-oil range unattainable by Ru-Zr and other Ru-supported catalysts. Importantly, the enhanced performance is showcased for real-world, single-use LDPE consumables. Reactivity studies combined with characterization and density functional theory calculations reveal that highly dispersed (WO\textsubscript{x})\textsuperscript{n} clusters store H as surface hydroxyls by spillover. We correlate this hydrogen storage mechanism with hydrogenation and desorption of long alkyl intermediates that would otherwise undergo further C−C scission to produce methane.

KEYWORDS: plastic waste, hydrogenolysis, low-density polyethylene, ruthenium, tungstated zirconia, hydrogen spillover

INTRODUCTION

Over 12 billion tons of plastic waste are projected to end in landfills or leak into the environment by 2050.\textsuperscript{1} Roughly 60\% of the global plastics are polyolefins, such as polyethylene and polypropylene, whose recycling rates are currently too low.\textsuperscript{2−4} The chemical composition and structure giving rise to the materials’ desired properties, such as their stability, is the very reason their recycling is so challenging. Mechanical recycling and incineration are somewhat limited in scope for these plastics. Chemical recycling and specifically catalytic depolymerization is potentially attractive to combating polyolefin plastic waste.\textsuperscript{5−12} Among approaches, catalytic cracking is facile, but it requires high reaction temperatures (typically above 673 K), and is prone to low-value gas products.\textsuperscript{5,11,13−16} Olefin-mediated metathesis\textsuperscript{17} has recently been applied to polyethylene decomposition using a large volume of pentane and long residence times.\textsuperscript{5} Bifunctional hydrocracking has been proven feasible at moderate temperatures and H\textsubscript{2} pressures.\textsuperscript{5,12,18−22} High branching imparts desirable properties to fuels and lubricants. However, it requires suitable catalysts to achieve appropriate metal−acid balance in strength and site density;\textsuperscript{12} the rate-limiting acid functionality often impedes high activities at lower temperatures.

Metal-catalyzed hydrogenolysis has recently drawn significant attention due to its decent reactivity and selectivity at low temperatures. Prior work has focused on discovering hydrogenolysis catalysts to produce fuels for proof-of-con-cept.\textsuperscript{6,8,23−26} For example, the hydrogenolysis of polyethylene over an ordered mesoporous shell/active site/core, Pt/SiO\textsubscript{2} catalyst was showcased to demonstrate a processive mechanism.\textsuperscript{23} Ruthenium-supported catalysts, e.g., Ru/C,\textsuperscript{24,26,28} Ru/zeolites,\textsuperscript{29} and Ru/ CeO\textsubscript{2},\textsuperscript{23} possess higher activity than Pt\textsuperscript{25} and are potentially promising materials. Unfortunately, Ru generates significant methane, which is an undesirable, low-value product. Besides, most studies involve either a large catalyst-to-polymer ratio, long processing times, or solvents. Ultralow molecular weight plastics are commonly employed as substrates to enable high conversions at low temperatures.\textsuperscript{23,24} While exciting initial results have been obtained, there is a clear need for more active and selective catalysts to enable practical implementation.

Here we demonstrate that ruthenium on tungstated zirconia (Ru-WZr) is highly active for the hydrogenolysis of low-density polyethylene (LDPE, molecular weight, M\textsubscript{w}, ∼76 kDa) at mild conditions (523 K and 50 bar H\textsubscript{2} for 2 h). The deconstruction rate is fast, although a direct comparison to previous contributions is impossible due to different
experimental conditions, substrates (especially regarding the large variability in molecular weight), and catalyst supports. Compared to Ru supported on bare zirconia (Ru-Zr), tungstated silica (Ru-WSi), HY zeolite (Ru-HY), and mesoporous [Al]MCM-41 (Ru-[Al]MCM-41) as benchmarks, the Ru-WZr catalyst significantly suppresses methane; it produces a heavier and more valuable carbon product distribution for fuel and wax/lubricant base-oil. We hypothesize that excessive methane is primarily made via cascade surface reactions when hydrogenation and desorption of long alkanes are rate-limiting. Extensive catalyst characterization and density functional theory (DFT) calculations demonstrate highly dispersed (WO\textsubscript{x})\textsubscript{n} clusters that increase the H storage capacity in surface hydroxyls by spillover from Ru to WO\textsubscript{x}. We provide evidence for a strong correlation of the excess hydrogen storage and the hydrogenation of intermediates, possibly by reverse spillover, interfacial sites, and/or catalyst−polymer interactions and desorption from the catalyst to avoid significant cracking leading to small hydrocarbons and methane. The implications for the rational design of polyolefin hydrogenolysis catalysts are discussed.

**RESULTS**

**LDPE Hydrogenolysis on Ru-WZr Catalysts**

We first demonstrate the promotional role of WO\textsubscript{x} in Ru-WZr catalysts on the mild hydrogenolysis of low-density polyethylene (LDPE, M\textsubscript{w} \sim 76\,\text{kDa}). The operating conditions are rather mild, namely, a relatively low temperature of 523\,K (250\,°C), ensuring depolymerization proceeds in the melt phase, a short processing time of 2\,h, and a low amount of catalyst (polymer to catalyst mass ratio of 40).

Figure 1 compares the product distributions over Ru-Zr, Ru-15WZr, and Ru-25WZr catalysts (hereafter, the number in front of W indicates the weight percent of the added WO\textsubscript{x}). Overall, the catalysts are significantly active. The overall nonsolid (liquids plus gases) yield is relatively high given the mild conditions (Figure 1a: 75%, 73%, and 55%, respectively) and is similar among catalysts, Ru-25WZr is less active but more selective to heavy liquids. Figure 1a−d shows that WO\textsubscript{x} has a profound effect on product distribution at similar conversions. It effectively decreases the gas yield from 21 to 7% and the undesired methane yield from 16% to 4.6%. It consequently enhances the selectivity to liquids as follows: gasoline from 25% to 28%, jet fuel from 33% to 42%, diesel fuel from 47% to 66%, and wax/lubricant base-oils from 22% to 35%. The results obtained over Ru-15WZr were benchmarked by good performing catalysts, Ru−Ce and Ru−C, reported previously\textsuperscript{23,24} (Figure S1 and the discussion in the Supporting Information). Effectively, the Ru-WZr catalysts shift the distribution to heavier, more valuable, soluble products (the peak of the distribution shifts from C\textsubscript{11} to C\textsubscript{15}). We elaborate with extensive characterizations that the apparent reaction rate of the Ru-WZr catalysts is affected by the interplay of reduced Ru site density and an increased turnover frequency and is comparable to that on Ru-Zr. The key difference between catalysts (Figure 1) is in selectivity.

Differential scanning calorimetry (DSC) of the solid residue (Figure 1e) and pristine LDPE (melting temperature, T\textsubscript{m} \sim 385\,K) shows that the leftover solid over Ru-Zr possesses a low-temperature broad feature centered at 342\,K, a shoulder at 358\,K, and a high-temperature sharp feature centered at 373\,K. The results obtained over Ru-15WZr were benchmarked by good performing catalysts, Ru−Ce and Ru−C, reported previously\textsuperscript{31,32} (Figure S1 and the discussion in the Supporting Information). Effectively, the Ru-WZr catalysts shift the distribution to heavier, more valuable, soluble products (the peak of the distribution shifts from C\textsubscript{11} to C\textsubscript{15}). We elaborate with extensive characterizations that the apparent reaction rate of the Ru-WZr catalysts is affected by the interplay of reduced Ru site density and an increased turnover frequency and is comparable to that on Ru-Zr. The key difference between catalysts (Figure 1) is in selectivity.

Figure 1. Effect of WO\textsubscript{x} incorporation in Ru-WZr catalysts on LDPE hydrogenolysis. Yields and selectivities for LDPE hydrogenolysis over Ru-Zr (red), Ru-15WZr (blue), and Ru-25WZr (green) catalysts. a, Carbon balance (gray dots), yields of soluble (solid bars), and gaseous (faded bars) products. b, Selectivities by fuel range: gasoline, C\textsubscript{5}−C\textsubscript{12}; jet fuel, C\textsubscript{8}−C\textsubscript{16}; diesel, C\textsubscript{9}−C\textsubscript{22}; and waxes/lubricant base-oils, C\textsubscript{20}−C\textsubscript{35}. c and d, Detailed carbon distributions of the nonsolid products. e, Differential scanning calorimetry curves of the solid product residues referenced to pristine LDPE (gray curve). Auxiliary lines at 342, 358, 373, and 385\,K. Reactions performed at 523\,K and 50\,bar H\textsubscript{2} for 2\,h with 2\,g of LDPE and 50 mg of catalyst.
nonsolid is heavier (Figure 1c,d), indicating a heavier carbon distribution of the products. Reduction in the melting temperature may also result from varying branching degrees;34 we rule out this possibility due to forming mainly normal alkanes with low branching (Table S1). Excitingly, Figure 1 demonstrates that WO\textsubscript{x} modification of the Ru-WZr catalysts can suppress methane formation and shift the carbon distribution to larger, value-added alkanes, such as diesel and waxes/lubricant base-oils. WO\textsubscript{x} modification overcomes an essential barrier of the otherwise excellent Ru catalyst reported in other works and does so at short times and low temperatures.

Excitingly, hydrogenolysis of real-world, single-use plastic consumables, such as LDPE bottles, cling wraps, and pipettes (Figure 2e), over the Ru-15WZr catalyst gave near quantitative conversion (Figure 2a), with selective product distributions (Figure 2b−d) similar to that of the pristine LDPE shown in Figure 1.

Structural Characterization of Ru-WZr Catalysts

High magnification BF-TEM, HAADF-STEM, and EDS elemental mapping of the Ru-15WZr (Figure 3) and Ru-Zr (Figure S2a) catalysts shows that the Ru-Zr has a characteristic mesoporous structure of around 10 nm diameter (Figure S2a) and ∼1−5 nm Ru nanoparticles (Figure S2c,d) consistent with BET (Table S2) and H\textsubscript{2} titration data (Table S3). In situ EXAFS results (Table S4) show that the Ru−Ru coordination numbers of the reduced Ru-Zr and Ru-15WZr catalysts are 8.3 and 6.6, respectively, which indicate that the Ru particle size decreases on average only slightly with the WZr support.35

Distinguishing Ru from (WO\textsubscript{x})\textsubscript{n} in Ru-15WZr based on the TEM contrast is challenging due to similar atomic numbers. However, the surface roughness of Ru-15WZr (Figure 3a,b) and Ru-Zr (Figure S2a,b) indicates that the highly dispersed ∼1 nm clusters in the former are most likely (WO\textsubscript{x})\textsubscript{n}39 HAADF-STEM images (Figure 3e,f), XRD (Figure S3a), and STEM-EDS mapping (Figure 3g) further confirm the high dispersion of Ru nanoparticles and (WO\textsubscript{x})\textsubscript{n} clusters. Intriguingly, a Ru particle depicted in Figure 3c is partially covered with an overlayer, identified as a WO\textsubscript{x} cluster, using the lattice parameter36−38 (Figure 3d). This suggests that the (WO\textsubscript{x})\textsubscript{n} species are mobile40 and may redisperse onto the metal of Ru during impregnation/reduction. Ambient temperature Raman spectra (Figure S3b) confirmed that the (WO\textsubscript{x})\textsubscript{n} species are highly amorphous and lack WO\textsubscript{3} crystalline structure in the presence of Ru. Furthermore, the HAADF-STEM data and EDS Ru and W maps shown in Figure 3g indicate that the two elements show substantial spatial overlap. These partial overlayers significantly decrease the Ru site density when increasing WO\textsubscript{x} loading (Table S3). However, the LDPE reactivity on Ru-WZr catalysts does not decrease as much (Figure 1a), suggesting that WO\textsubscript{x} enhances the Ru turnover rate.

The 15WZr sample has a WO\textsubscript{x} surface density (∼6.6 W/nm\textsuperscript{2}, Table S2) corresponding close to a monolayer coverage.41,42 WO\textsubscript{3} crystallites are nearly absent in the XRD or Raman spectra (Figure S4a,b). Unlike ZrO\textsubscript{2} with a tetragonal or monoclinic structure, ZrO\textsubscript{2} in 15WZr is predominantly tetragonal (Figure S4a). Characteristic Raman bands in ZrO\textsubscript{2} decrease significantly with increasing W loading (Figure S4b), suggesting a high WO\textsubscript{x} coverage. The 25WZr sample has similar structural features except for WO\textsubscript{3} crystallites seen in XRD and Raman spectra (Figure S4a,b) due to a higher loading (∼10.7 W/nm\textsuperscript{2}, Table S2; more than one monolayer coverage). The 15WZr sample exhibits Bronsted acidity based on pyridine adsorption (Figure S4c), and Ru does not
significantly affect the acidity of the 15WZr support (Figure S4d). However, the Ru-15WZr shows negligible isomerization activity (Table S1), suggesting a lack of metal−acid bifunctional hydrocracking−hydroisomerization chemistry. Compared to the bifunctional Pt-WZr catalyst we reported recently, which promotes a high degree of branching,19,43 Ru is much more active for deep dehydrogenation and subsequent C−C scission due to stronger metal−carbon bond strength.44,45 This prevents Ru from producing olefins that undergo hydrocracking and isomerization over the Brønsted acid sites.46 Two additional Ru catalysts on highly acidic HY (Si/Al = 30) zeolite and mesoporous [Al]MCM-41 (Si/Al = 25) were prepared to understand the role of acidity. Moderate branching (Table S5) was observed, likely due to the stronger Brønsted acid strength47 and much greater site density of these supports.19 However, these catalysts are not as active (56% and 68% conversion over Ru-HY and Ru-[Al]MCM-41, respectively) and produce much higher methane yields (36% and 52%, respectively) compared to Ru-15WZr under identical conditions. These findings suggest that acidity is not as critical a factor to methane suppression over the Ru-WZr catalysts.

Proposed Reaction Mechanism over Ru-WZr Catalysts

The time evolution for LDPE hydrogenolysis over Ru-15WZr is depicted in Figure 4. The data exhibits characteristic sequential reactions converting longer alkanes to shorter ones, evidenced by bell-shaped trajectories of C20+, C17−C19, and C13−C16 (longer molecules reach maxima at shorter times), with a monotonic growth of C8−C12, C5−C7, and C1−C4. The nonsolid products progressively shift toward lower molecular weights (Figure 4c,d). Methane becomes dominant (20% yield) only at long reaction times after severe over-reaction due to the sequential nature of the reaction. Noticeably, the nonsolid yields experience a 10-fold increase from 7% to 73% between 1 to 2 h (Figure 4a). In this time
interval, further experiments (Figure 4b) reveal a light-off in nonsolid yields between 1.5 and 1.75 h (from 12% to 32%). TEM images of postreaction (2 h) catalyst (Figure S5) show no noticeable morphological or size change in Ru and (WO₄)ₙ clusters compared to the fresh sample (Figure 3a−d). No sudden changes in the solid DSC curves occur at this time interval (Figure S6), i.e., the melt of the LDPE is steadily depolymerized. This indicates the hydrogenolysis product distribution swiftly shifts to lower carbon numbers in the early stages of the reaction and becomes GC detectable in the C₁−C₃₅ carbon range after 1.5 h (Figure 4b). At 1 h, the yield of C₁−C₃₅ alkanes is only 7% (of which methane is 3%); during this time, the LDPE (Mₖ ~ 76 kDa) is substantially decomposed to the ~C₃₀₀ range (estimated from the Mₖ of ~4 kDa LDPE standard).

Notably, methane is produced in disproportionally large fractions early on (Figure 4b), and its yield increases relatively linearly with time (inset in Figure 4b), indicating its parallel production with the hydrogenolysis of long alkanes. Continuous flow hydrogenolysis experiments using n-hexane as a surrogate are consistent with this hypothesis (Figure S7, see discussion in Supporting Information), whereby methane forms via a cascade of multiple C−C bond scissions (iv−vii) before hydrogenation and desorption (vi). Since these intermediates are bound to the Ru surface with newly formed terminal carbons (ii), further C−C bond breaking favors demethylation to make methane (iv−vii). Ru catalysts favor terminal C−C bond breaking even in the primary C−C scission (i−iii). The proposed cascade mechanism (iv−vii) must be invoked in the decomposition to rationalize the disproportionally high methane yields in both LDPE and model hexane reactions consistent with recent contributions. Catalytic performance at various hydrogen pressures supports this mechanism (Figures S8−S10, see detailed discussion in Supporting Information). Activities and selectivities (lower methane yield and heavier product distributions) increase over Ru-Zr and Ru-15WZr catalysts with increasing H₂ pressure. A higher H₂ pressure promotes the rate-limiting hydrogenation of intermediates followed by their desorption (iii), thereby enhancing the apparent rate and suppressing methane formation (iv−vii).

We hypothesize that the promotional effect of WO₄ (Figure 1) may share the same origin as the hydrogen pressure. Indeed, the benefit of high H₂ pressure diminishes with increasing WO₄ loading (Figure S10). At sufficiently high H₂ pressure, such as 130 bar, the hydrogenation/desorption (iii) step no longer limits the overall rate. The product distributions on Ru-Zr and Ru-15WZr become similar at comparable conversions (Figure S8g,h) with rates proportional to their Ru site density (123 and 76 μmol/g, respectively, Table S3). This suggests...
that the Ru-catalyzed C–C bond activation, which is ratelimiting at high pressures, is not modified by WO$_x$ incorporation. This is consistent with the n-hexane model reaction results (Figure S7c), which show an identical TOF per C–C bond scission over Ru-Zr and the two Ru-WZr catalysts (see detailed discussion in Supporting Information). Our structural characterizations show little difference in particle size of Ru-Zr and Ru-15WZr catalysts (see EXAFS results in Table S4), and XPS data also provide no evidence of electronic modification of Ru by WO$_x$. These findings altogether indicate that the contribution of Ru is not impacted by the WO$_x$ incorporation. The benefit of Ru-WZr catalysts in H$_2$ deficient environments may arise from additional hydrogen, polymer–surface interaction, and/or interfacial sites created on these catalysts. The data suggests that the (WO$_x$)$_n$ clusters may supply extra H via reverse-spillover to facilitate the rate-limiting hydrogenation/desorption (iii) on the Ru surface or the interface or modify the polymer–catalyst interactions, enhancing both activity and selectivity and effectively changing the mechanism from Scheme 1c to Scheme 1b. We elaborate on this point next.

TPR-MS and H$_2$-TPD-MS measurements provide further insights into the enhanced performance of Ru-15WZr. TPR-MS results (Figure 5a) show that, at low temperatures (300–600 K), Ru-Zr consumes roughly 0.6 mmol H$_2$/g, primarily due to the reduction of surface Ru oxides and Ru-15WZr consumes an extra of 0.3 mmol H$_2$/g. This additional hydrogen likely remains on the catalyst at least until 600 K because the formation of H$_2$O below 600 K on both Ru-Zr and Ru-15WZr is quite similar (Figure S11). The reduction of WO$_x$ on 15WZr consumes $\sim$1.9 mmol H$_2$/g. It occurs at a much higher temperature of $\sim$900 K than Ru-15WZr. On the latter, the WO$_x$ reduction peak is much broader and starts at much lower temperatures of $\sim$600 K. The lower reduction temperature of WO$_x$ on Ru-15WZr stems from Ru’s role. These results indicate a possible hydrogen spillover, where Ru dissociates H$_2$ and H diffuses over to the (WO$_x$)$_n$ clusters. This extra H on (WO$_x$)$_n$ can be readily available for hydrogenation reactions on metallic Ru by the reverse spillover mechanism.

TPD-MS results (Figure 5b) show that H$_2$ desorbed from Ru-15WZr (0.16 mmol/g) is 3-fold that from Ru-Zr (0.05 mmol/g).
mmol/g), with a higher H binding energy, based on the desorption temperatures that extend beyond those of Ru-Zr and up to 800 K. DFT calculations (Table S6) of a W4O12 cluster on Ru (0001), chosen as a model structure, confirms that the surface H on W−OH groups is bound significantly more strongly (E(binding) = −1.43, −1.29, and −1.05 eV) than on the Ru sites (E(binding) = −0.93, −0.82, and −0.44 eV). In contrast, a Ru-20WSi catalyst does not possess synergistic features in TPR and TPD measurements (Figure S12); at the same time, it lacks highly dispersed (WOx)n clusters (XRD and Raman, Figure S13) and does not suppress methane formation (Figure S14).

Air-free XPS spectra in the W 4f + Zr 4p region (Figure 5c) and Zr 3d region (Figure 5e) were collected after pretreating Ru-15WZr at 523 K and 1 bar in a 10% H2 flow. Deconvoluted and benchmarked spectra against the air-equilibrated ones are shown in Figure 5d,f. Both the air-exposed W and Zr oxides are at the highest valence, evidenced by doublets for W 4f at 38.0 and 36.0 eV, Zr 4p at 32.5 and 30.9 eV (Figure 5d), and Zr 3d at 185.0 and 182.7 eV (Figure 5f). Air-free Ru-15WZr (Figure 5c) shows reduction bands associated with both W5+ cations (36.4 and 34.4 eV) and Zr suboxides (30.7 and 29.1 eV for Zr 4p and 183.4 and 181.2 eV for Zr 3d). Deconvolution of the W4+ center is impossible due to overlap with the Zr 4p and is omitted from the analysis. Since (WOx)n clusters are predominantly ∼1 nm in size (within photoelectron escape depth), we estimate that roughly 30% of W 6+ centers are reduced to 5+ (or lower) oxidation state based on the deconvolution area ratios.

Since little water forms from WOx reduction up to 600 K, based on the TPR-MS on Ru-15WZr (Figure S11), we envision that hydrogen spillover leads to Hδ+ interacting with the Lewis basic oxygen atoms (W=O or W−O−W) while injecting electrons into the W6+ centers to lower their oxidation state (Scheme 1b), similar to frustrated Lewis pairs.53,54 Since the degree of hydrogen incorporation is in part

Figure 5. Role of hydrogen on Ru-15WZr catalyst. a, TPR-MS (negative features, m/z = 2). b, H2-TPD-MS (positive features, m/z = 2) measurements on 15WZr (black), Ru-Zr (red), and Ru-15WZr (blue). c−f, XPS spectra of (c, e) air-free and (d, f) air-exposed Ru-15WZr catalyst in the (c, d) W 4f + Zr 4p region and (e, f) Zr 3d region. Contributions from W6+, W5+, Zr4+, and partially reduced ZrO2 were deconvoluted and are illustrated in purple, light purple, red, and light red, respectively. g, XPS spectra of air-free (black) and air-exposed (red) Ru-15WZr in the region of C 1s and Ru 3d for Ru metal, RuO2, and/or RuOx·xH2O. All catalysts for XPS were pretreated in 10% H2/He at 523 K for 1 h with the air-exposed samples being equilibrated at air room temperature and the air-free samples being immediately transferred (at room temperature) for measurements without exposing to air (P2<sub>O2</sub> < 1000 ppm) using a vacuum transfer vessel. h and i, Difference IR spectra of Ru-15WZr at (pink) 323 K, (blue) 373 K, (green) 423 K, and (red) 523 K collected upon exposure to steady-state 10% H2 flow carried by 60 sccm He. h, Stretching vibration regime of W=O and W−O−W groups. i, Bending vibration regime of O−H groups. Difference spectra were obtained by subtracting the absorbance spectra collected prior to H2 exposure from the spectra post-H2 exposure at each temperature. Horizontal dashed lines represent estimated baselines. j, Simulated bending vibration modes of hydroxyl on a W4O12 cluster, supported on Ru(0001).
determined by the ability to delocalize the injected electrons into the conjugated base, i.e., the (WO)₆₈ clusters, it is not surprising then that the hydrogen storage capacity is associated with the ~1 nm (WO)₆₈ clusters on Ru-15WZr rather than bulk crystallites on Ru-20WSi. The slight reduction of the ZrO₂ surface may also result from hydrogen spillover. Since Ru and WO₆ species are highly dispersed on ZrO₂ (Figure 3), the partially reduced ZrO₂ sites may be implicated by bridging H₆ species and electrons between Ru and (WO)₆₈ islands (Scheme 1b). Such an effect is impossible on nonsemiconducting supports, such as SiO₂ (Scheme S1).

XPS bands associated with Ru 3d (Figure S5g) are complicated due to C 1s overlaps and interference between the Ru metal and hydrated/dehydrated oxides. By subtracting the two Ru-15WZr spectra with and without air exposure, contributions of the metallic Ru 3d₅/₂ band (~280 eV) become evident in the air-free sample; yet, Ru is likely to remain slightly oxidized in the air-free sample, judging by the bandwidth and according to the literature on Ru/TiO₂.

We performed in situ transmission FTIR measurements on Ru-15WZr catalyst under relevant reaction conditions (323–523 K and 1 bar in 10% H₂/He flow) and obtained difference absorbance spectra by subtracting the collected spectra before and after steady-state H₂ exposure at each temperature (see Figure S15 for spectra overview and peak assignments). All samples were pretreated in H₂ and purged in He at 623 K to ensure water formation does not interfere with the results acquired at lower temperatures.

Upon H₂ exposure, the difference IR features from 323 to 523 K in the W/O stretching regime (Figure S5h) are similar to a decrease in the dehydrated ν(W=O) mode (1012 cm⁻¹) and an increase in the hydrated ν(W=O) mode (972 cm⁻¹). Since pretreatments rule out water formation, we assign this band shift to W=O hydrogen bonding with the proximal H₆ species and/or OH groups (i.e., W=O···H and/or W=O···H–O–W). The relative intensity of hydrated (+)/dehydrated (−) ν(W=O) bands decreases at higher temperatures, suggesting consumption of the W=O bonds at high temperatures and probable transformation into W–OH species with formal O–H bonds. The broad spectral envelope at 918 and 826 cm⁻¹ can have contributions from both ν(W=O–Zr) and ν(W–O–W) modes, as in the multiwavelength Raman approach on transitional metal oxides. The decrease in the difference absorbance (Figure S5h) is possibly due to electron delocalization on WO₆ clusters induced by charge separation of spillover hydrogen. Similar trends in the overtone regime at ~2000 cm⁻¹ further corroborate the above changes (Figure S16). The same methodology on a benchmark 15WZr sample shows 1–2 orders of magnitude weaker changes in difference spectra (Figure S17) when the intensities of absolute absorbance spectra are comparable (Figure S18a,b). Moreover, the relative band areas of hydrated (+)/dehydrated (−) W=O features do not change significantly at elevated temperatures, suggesting that no W=O/W–OH transformations occur without hydrogen spillover from Ru. Difference features associated with ν(W=O–Zr) and ν(W–O–W) modes are absent, suggesting no charge separation on the (WO)₆₈ clusters without Ru. The W/O vibrational changes in response to H₂ exposure on both Ru-15WZr and 15WZr catalysts were subsequently confirmed by in situ Raman measurements (Figure S18c–f) performed under the same conditions.

Consistent difference absorbance for the δ(OH) modes on Ru-15WZr is depicted in Figure S1. The corresponding ν(OH) modes exhibit very broad features at ~3000 cm⁻¹ due to H-bonding and are shown in Figure S16b. In the δ(OH) regime, the band at ~1610 cm⁻¹ resembles the deformation vibrations of water hydroxyls. In the absence of molecular water (Figure S19), we ascribe this feature to water-like surface species that release molecular H₂ instead of H₂O upon heating or evacuation. The difference features at ~1414 and 1200 cm⁻¹ were previously assigned to δ(OH) modes of W–OH interaction with surface oxygen atoms to a different degree. The band at 1610 cm⁻¹ diminishes at elevated temperatures due to thermal instability. The 1414 cm⁻¹ feature remains stable and is ultimately reduced at 523 K. At high temperatures, a third, quite unusual δ(OH) band emerges at ~1220 cm⁻¹. DFT calculations of a W₂O₁₂ cluster on Ru (0001), as a model structure (Figure S5j), reveal that this unique δ(OH) mode may be assigned to W–O–H groups adjacent to a W=O group (Movie 1; details in Supporting Information). This bending vibration is absent in W₂O₁₂ without Ru (Movie 2). We observed the ~1220 cm⁻¹ band grows with increasing temperature, whereas the 1600 and 1414 cm⁻¹ bands decrease in unison (Figure S1). This is consistent with the decrease in the relative band areas of hydrated (+)/dehydrated (−) ν(W=O) features (Figure S5h), which suggest the transformation from W=O to W–O–H bonds. Experiments on 15WZr or Ru-Zr (Figure S19) hardly give any such difference features, highlighting the Ru-WO₆ synergy in Ru-WZr. The δ(OH) bands at ~1610, 1414, and 1220 cm⁻¹ were definitively assigned by replicating experiments using molecular deuterium (Figure S20). The observed δ(OD) wavenumbers agree well with those derived from the corresponding δ(OH) modes calculated using reduced mass. Relaxation of the H₂-equilibrated Ru-15WZr catalyst (Figure S21) shows that all the different features associated with ν(W/O) and δ(OH) modes nearly reversed to the original states when H₂ was removed from the feed stream at 523 K. This suggests that the extra H stored in hydroxyls is highly mobile, can reverse spillover back to the Ru surface, and may be subject to recombinative desorption of molecular H₂ (TPD, Figure S5b).

Collectively, multiple characterization approaches and DFT calculations corroborate the presence of an extra amount of highly mobile and active H species on the Ru-WZr catalysts. These H species can undergo reversible spillover between Ru and (WO)₆ clusters, aiding hydrogenation and desorption (iii, Scheme 1b), enhancing activity and selectivity in LDPE hydrogonyalysis under H₂ deficient conditions. The correlation of H storage with selectivity is unambiguous. However, direct proof of the implication of the reverse spillover mechanism is not possible at this moment. We hypothesize that new interfacial sites created by OH and modified polymer–catalyst interactions could also possibly be at play. Further work is needed to dissect the precise mechanism.

**Discussion**

We demonstrated that Ru-WZr catalysts are highly active and selective for LDPE hydrogonyalysis at mild conditions (low temperatures, short times, a low catalyst to polymer mass ratio). Importantly, this performance extends to real-world plastic consumables, such as LDPE bottles, cling wraps, and pipettes, where reaction times could be shorter and conversions higher. Excitingly, they significantly suppress low-value methane formation and produce higher molecular weight fuels and wax/lubricant base-oils compared to Ru/C.
and Ru/zeolites. They are also superior to Ru-Zr and Ru-WSi catalysts, demonstrating the WO₃ and ZrO₂ support’s profound role. The WO₃ addition overcomes Ru’s high hydrogenolysis activity while overcoming its extensive chain cracking to methane. Characterization and DFT calculations confirm that the highly dispersed (WO₃)ₙ clusters store H in surface hydroxyls by the spillover of H from the Ru active phase that dissociates molecular H₂. WO₃ increases the catalyst hydrogen storage capacity essential for hydrogenating long-chain alkyl intermediates through reverse spillover to Ru or by creating new interfacial sites and/or modifying the catalyst—polymer interactions, leading to their desorption before they get overcracked. This novel mechanism of hydrogen storage overcomes significant compression costs associated with high H₂ pressures and opens up possibilities for materials design to further engineer plastics recycling.

## MATERIALS AND METHODS

### Preparation of Ru-WZ Catalysts

Ruthenium supported on tungstated zirconia (Ru-WZr) was synthesized using the successive incipient wetness impregnation method. The tungstated zirconia (WZr) support was prepared by impregnation of zirconium oxyhydroxide (ZrO₂(OH)₂·nH₂O) with a solution of ammonium metatungstate hydrate ((NH₄)₆H₂W₁₂O₄0·xH₂O, Aldrich, 99.99%) and stirring. The mixture was then transferred into a 50 mL stainless-steel Parr reactor with a 1/4-in. ID quartz tube placed vertically in a tube furnace. The catalyst bed (typically 100 mg) was diluted 100 times using inert ZrO₂ supports and held in place by quartz wool plugs and coarse quartz pellets loaded on both ends of the catalyst bed. The reactor was custom-made to insert a thermocouple, protected by a quartz protection sleeve, at the center of the catalyst bed. Before the kinetic measurements, the catalyst was pretreated at 523 K (10 K/min ramping rate) for 1 h in 10% H₂ in 100% He. n-Hexane (1–2%) was injected into a He carrier flow using a syringe pump (New Era Pump System) and mixed with H₂ flow before entering the reactor.

### Product Analysis

Products from the probe reaction of n-hexane hydrogenolysis were analyzed online using an FID gas chromatography equipped with an Agilent HP-Plot GC column. For the LDPE hydrogenolysis reactions, gaseous products, typically C1–C6, from the Parr reactor headspace were sampled at 283 K using a 1 L Tedlar bag and analyzed using the aforementioned GC. Quantification was done using standard calibration mixtures. Hydrocarbons, typically C4–C35, were extracted from the liquid—solid residues using a 20 mL of dichloromethane (CH₂Cl₂, Fisher Scientific, ACS grade) solution with 20 mg of n-octacosane (external standard, n-C28, TCI chemicals, >98%). Since the reaction produces a small amount of n-C28, the signal of internal standard is calculated from the raw n-C28 product signal using an average of n-C27 and n-C29. The soluble products were analyzed using GC-MS (Agilent, DB-1 column) for identification and GC-FID (Agilent HP-1 column) for quantification. A representative raw GC trace is shown in Figure S23. Calibration of GC signals was conducted using normal alkanes standard mixtures, and alkane isomers were assumed to have the same calibration factor as the corresponding normal alkanes. The catalyst and remaining solids were filtered, dried in air for 24 h, and weighed prior to further characterization.

The yield of product alkanes with i carbons (Cᵢ) was calculated using the following equation:

\[
Y_i = \frac{n_i}{n_{\text{initial}}}
\]

where \(n_i\) is the moles of carbon in Cᵢ and \(n_{\text{initial}}\) is the same in the initial LDPE feedstock. The selectivity of product alkanes with i carbons (Cᵢ) is evaluated from

\[
S_i = \sum_{\text{solid}} Y_i
\]

Here \(\sum_{\text{solid}} Y_i\) corresponds to the yield of nonsolid products—namely, the combined yield of all gaseous and soluble liquid products.

High-resolution differential scanning calorimetry (DSC, Discovery, TA Instruments) measurements were performed to characterize the melting temperature and heat of fusion for pristine LDPE and the solid residues collected after filtration.

### Catalyst Characterization

X-ray diffraction (XRD) patterns of powdered materials were measured using a Bruker D8 diffractometer equipped with a Cu Kα source. The BET surface area and porosity were measured using N₂ as the standard gas.
the adsorbent at 77 K with a Micromeritics ASAP 2020 instrument. Raman spectra were obtained with a LabRAM high-resolution Raman spectrometer (HORIBA) equipped with a He−Cd excitation UV laser with a wavelength of 325 nm and a green line solid-state laser with a wavelength of 532 nm. In situ Raman spectroscopy was performed in an environmentally controlled high-temperature cell (Harrick). Elemental analysis was carried out on a wavelength-dispersive X-ray fluorescence spectrometer (WDXRF, Rigaku, Supermini 200) equipped with a high-powered (200 W) X-ray tube. For X-ray photoelectron spectroscopy (XPS) analysis, powdered samples were pressed onto Cu foils. XPS spectra were collected using a Thermo Fisher K-Alpha Instrument equipped with an Al(K) X-ray source. Samples used in air-free XPS measurements were pretreated in a tubular flow reactor using 10% H2 in a 100 sccm He carrier gas for 2 h at 523 K. The cooled reactor was then sealed and moved into a glovebox (P02 < 1000 ppm), where the samples were transferred into a designated vacuum vessel for XPS measurements. Selected results of compositional and geometrical characterizations described above are summarized in Table S2.

Bright-field transmission electron microscopy (BF-TEM) was performed with a field emission gun transmission electron microscope (JEOL, JEM-2010F), operated at an accelerating voltage of 200 kV. Dark-field scanning transmission electron microscopy was performed with a JEOL NEOARM operating at 200 kV. The samples were diluted in isopropanol and deposited on a lacy carbon film on copper grids provided by Electron Microscopy Sciences. The probe current was 120 pA with a condenser lens aperture of 40 µm and a camera length of 4 cm. Energy-dispersive X-ray spectroscopy was performed with the same parameters, with the exception of a camera length of 2 cm. Two silicon drift detectors (SDD) provided by JEOL were used, and the maps were obtained with DigitalMicrograph, a software provided by Gatan Inc.

Apparent dispersions of Ru were measured on a Micromeritics AutoChem II instrument following a customized procedure of H2 chemisorption reported by Iglesia and coworkers. This method has previously been shown to have an H2/Ru = 1 stoichiometry and is not affected by the reduction history of Ru or SMSI effect. In brief, the sample was pretreated in situ under 10% H2/Ar flow for 1 h at 523 K followed by purging in pure Ar flow for 1 h at 523 K. The sample was cooled under Ar, maintained at 200 K (±10 K) using a liquid N2−ethanol bath, and subsequently treated in a 1% O2/He mixture at 200 K for 15 min followed by purging in Ar flow at 200 K for 15 min. Finally, the sample was heated in Ar flow and maintained at 373 K for 1 h. Ru dispersions were measured using pulse H2 chemisorption at 373 K. Results from all catalysts used in this study are tabulated in Table S3. Conventional chemisorption of either H2 or CO using simple pulse techniques at room temperature is reliable (Table S7). This is in part due to the activation energy of H2 adsorption on Ru at room temperature, especially on catalysts that featured fairly strong metal−support interaction (underestimation of Ru dispersion) and the fast reaction activation of CO on the Ru surface (overestimation of Ru dispersion).

Temperature-programmed reduction (TPR) and temperature-programmed desorption (TPD) of adsorbed hydrogen (H2−TPD) were carried out in combination with mass spectroscopy (MS) analysis (Pfeiffer Vacuum Ominstar GSD 301C quadrupole mass spectrometer) using a continuous flow reactor. For TPR-MS measurements, the sample was purged in He at 523 K for 1 h and cooled down to room temperature in He flow. A 60 sccm, 5% H2 in He flow was stabilized to steady state for at least 1 h. The TPR measurement was then carried out at a ramping rate of 10 K/min from 300 to 1273 K. For H2−TPD analysis, the sample was prerduced in a 60-sccm, 5% H2 in He mixed flow at 523 K for 1 h, cooled down to room temperature, and purged in a 60-sccm He flow for at least 1 h to stabilize the baseline of the H2 MS. The H2−TPD measurement was then carried out using the same ramping procedure as H2−TPD. MS signals were calibrated using a MicroGC (990 MicroGC, Agilent) equipped with an MSSA column. The MS signal for He was used as an internal reference for semiquantification of H2 uptake/desorption.

FTIR transmission spectra were acquired on a Nicolet 8700 spectrometer (Thermo Scientific) equipped with a liquid nitrogen cooled MCT detector and a custom-made Pyrex in situ flow cell sealed with KBr windows on both ends. Powdered samples were pressed into a self-supported disk (~12.5 mg, 1000 PSI) and aligned with the laser path using a quartz holder. FTIR measurements were performed to analyze the hydrogen surface species at temperatures ranging from 323 to 523 K. Moisture was removed from the gas feed by a liquid nitrogen trap. In order to eliminate the interferences from bulk reduction or water formation, all samples were prerduced in situ at 623 K for 1 h in 10% H2, carried by 60 sccm He flow and then purged in pure He at 623 K for 1 h to remove adsorbed hydrogen. Difference spectra were obtained at 323, 373, 423, and 523 K by subtracting each reference spectrum collected in He from the steady-state spectrum taken in H2 in He flow at the same temperature. Experiments with deuterium gas (D2, Cambridge Isotope Laboratories, D-99.8%) with the same methodology were repeated in order to confirm the hydroxyl vibration modes.

FTIR of adsorbed pyridine was also performed to characterize the Brunst ein and Lewis acid sites. The sample was pretreated in He flow at 523 for 1 h and then cooled and maintained at 423 K. The sample was then saturated by pyridine vapor introduced by passing He flow through a bubbler filled with liquid pyridine (Aldrich, 99.8%). Prior to spectrum acquisition, the sample was purged with pure He flow for 30 min to desorb weakly bonded pyridine thermally.

X-ray absorption spectroscopy (XAS) data were collected for the Ru K-edge (22.117 keV) at the QAS 7-8M beamline of NSLS-II at Brookhaven National Laboratory. The spectra of the reference (Ru foil) were measured in transmission mode, while spectra of sample catalysts were recorded in fluorescence mode. Before measurements, the sample was packed in a quartz tube connected to a gas line. Then the catalyst was reduced in 20 mL/min flow of a 50% H2/He mixture at 473 K for 30 min. Spectra were collected after reduction. Spectra were subjected to standard data processing using Athena and Artemis software.

## Modeling

Density functional theory calculations (DFT) were performed by the plane-wave based Vienna ab initio simulation package (VASP), version 5.4.4.68−70 The electron−electron exchange and correlation energies were computed using the Perdew, Burke, and Ernzerhof functionals with D3 dispersion corrections developed by Grimme.71,72 Core electrons were treated with the projector augmented-potential (PAW) pseudopotentials.73 The Ru (0001) surface was simulated using a four-layer 4×4 unit cell. The lattice constant of Ru was calculated to be 2.725 Å, in close agreement with the experimental value of 2.71 Å.74 The supported WO3 (0001) tetramer was constructed by cleaving a 2×1×1 unit cell from the monoclinic bulk WO3, which features lattice parameters of a = 5.358, b = 5.413, and c = 7.722 Å. The lattice parameters are in good agreement with experimental values of a = 5.28, b = 5.16, and c = 7.76 Å.75 The bottom two layers of Ru were held fixed during calculations, whereas the rest of the all atoms were allowed to relax. A vacuum padding between the slabs was set at a height of 16.5 Å to minimize the interaction in the z-direction. In addition, dipole corrections were applied normal to the surface. The Brillouin zone was integrated using a 3×3×1 k-mesh based on the Monkhorst−Pack method,76 with Gaussian smearing of 0.1 eV. For the plane-wave set, a cutoff energy of 400 eV was used. Electronic structure calculations were performed iteratively, and electronic structures were converged when the difference in energy between subsequent steps fell below 10−6 eV. Geometries of adsorbate−metal systems were optimized until the maximum of all forces fell below 0.02 eV/Å. Vibrational modes analysis was carried out using the central finite difference method. In these calculations, atoms were displaced by 0.05 Å in all directions by the VASP default. In the vibrational mode calculations, the adsorbate-hydrogen and the WO3 cluster were displaced.
ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/jacsau.1c00200.

LDPE hydrogenolysis on Ru-WZr catalysts (Figure S1, Table S1); catalyst characterization and comparison of LDPE hydrogenolysis reactivity to zeolite supported Ru (Figures S2–S5, Tables S2–S5); solid residue characterization (Figure S6); steady-state n-hexane hydrogenolysis at 1 bar (Figure S7); effect of hydrogen pressure on LDPE hydrogenolysis (Figures S8–S10); characterization and DFT calculations on hydrogen effect (Figures S11 and S12, Table S6); comparison of catalyst structure and reactivity to a Ru-WSi catalyst (Figures S13 and S14, Scheme S1); computational infrared vibrational modes over Ru-WZr catalyst (Movies 1 and 2); in situ spectroscopic measurements (Figures S15–S21); materials and methods (Figures S22 and S23, Table S7) (PDF)

Raw data and DFT files (ZIP)

Movie S1 showing the vibrational mode of O–H bending in a Ru-(WO₃)₄-H system (MP4)

Movie S2 showing the bending mode unique to the WO₃ cluster on Ru (MP4)

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Author Contributions

C.W. and D.G.V. with the support of P.A.K, W.Z., and S.L. developed the conceptual foundations and led the writing and revision of the original manuscript. T.X. performed DFT calculations and analysis. C.W. led the experimental measurements and data analysis. B.C.V. contributed to the reactivity measurements. K.Y. performed the TPD/TPR sorption experiments. J.F. performed the XPS experiments. P.K. and E.A.S. performed and analyzed the HAADF-STEM and EDS elemental mapping. P.K. performed and analyzed the EXAFS results. G.T. participated in spectroscopic measurements and reviewed the results. W.Z. performed and analyzed BF-TEM. All authors wrote, reviewed, and approved the final manuscript.

Notes

The authors declare no competing financial interest.

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