Highly Effective Propane Dehydrogenation Using Ga–Rh Supported Catalytically Active Liquid Metal Solutions

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ABSTRACT: Our contribution demonstrates that rhodium, an element that has barely been reported as an active metal for selective dehydrogenation of alkanes becomes a very active, selective, and robust dehydrogenation catalyst when exposed to propane in the form of single atoms at the interface of a solid-supported, highly dynamic liquid Ga–Rh mixture. We demonstrate that the transition to a fully liquid supported alloy droplet at Ga/Rh ratios above 80, results in a drastic increase in catalyst activity with high propylene selectivity. The combining results from catalytic studies, X-ray photoelectron spectroscopy, IR-spectroscopy under reaction conditions, microscopy, and density-functional theory calculations, we obtained a comprehensive microscopy picture of the working principle of the Ga–Rh supported catalytically active liquid metal solution.

KEYWORDS: catalysis, dehydrogenation, alkane, alkene, gallium, rhodium

INTRODUCTION

Following our general interest in novel supported liquid catalysts for high temperature applications, we have very recently proposed supported catalytically active liquid metal solutions (SCALMS) as a very promising class of heterogeneous catalysts. Published SCALMS systems are composed of catalytically active liquid alloy droplets (Ga,Pd) or Ga,Pt on a porous support. In contrast to conventional supported liquid phase catalysis, the catalytic reaction in SCALMS occurs only at the highly dynamic liquid metal/gas interface, as the liquid metal does not provide any relevant reactant solubility. The first account of SCALMS systems in catalysis described Ga-rich Ga,Pd mixtures (Ga,Pd ratio > 10) on porous glass and revealed high activity and high robustness of these systems against coking in n-butane dehydrogenation at 450 °C. The liquid nature of the supported alloy droplet under the reaction conditions was confirmed through a combination of X-ray diffraction, scanning electron microscopy (SEM), X-ray photoelectron spectroscopy (XPS), and ab initio dynamics calculations.

Shortly after our initial SCALMS work was published, Upahm et al. reported a bulk liquid metallic solution of nickel in bismuth, which could be used up to temperatures of 1100 °C for methane pyrolysis to hydrogen and elemental carbon. These authors showed that the activity and stability of the bulk liquid metallic catalyst were higher than that of its solid counterparts. While a Ni wire lost 50% of its activity in less than 1 h, the bulk liquid metallic catalyst could be used up to 1200 h, without any significant loss of performance. With their findings, these authors confirmed general statements in earlier works by Ogino, claiming that the intrinsic dynamics of liquid metal surfaces would help to avoid the loss of activity in bulk liquid metal catalysis. Severe corrosion issues, however, arise when operating bulk liquid metals and their alloys in contact with metallic reaction equipment, making this sort of bulk liquid metal catalysis challenging from an application point of view.

So far, only a small number of examples of rhodium-based alkane dehydrogenation catalysts have been reported. The excellent and very comprehensive review on alkane dehydro-
Rh phase diagram,11 these compositions lead to the formation of a atomic Ga/Rh ratio of 25 and 34. According to the Ga aqueous suspensions of the metallic Ga/AlO was achieved by partial galvanic replacement of Ga. Briefly, mixed with di-

When the reaction temperature was reached, a out in a tubular quartz reactor. The reactor was heated to the °

in order to obtain the materials with the intended di-

These highly diluted compositions result in supported, fully liquid droplets of the Ga–Rh alloy at a temperature of 550 °C.11

We demonstrate that the addition of Ga to the alumina-supported Rh catalyst leads to an enhanced catalytic performance. Interestingly, a strong additional boost in Rh-based activity can be found at the transition to the fully liquid Rh–Ga system. We performed XPS, IR-spectroscopy, and microscopy studies in combination with density-functional calculations to reveal how catalysis at the liquid interface of Rh–Ga SCALMS systems proceeds at the microscopic level.

■ RESULTS AND DISCUSSION

Synthesis of Ga–Rh-SCALMS Materials. The Ga–Rh SCALMS materials were prepared using the procedure recently described for Ga/Pd systems.1 Analogously, the deposition of Rh was achieved by partial galvanic replacement of Ga. Briefly, aqueous suspensions of the metallic Ga/AlO material were mixed with different amounts of stock solution of RhCl₃·3H₂O, in order to obtain the materials with the intended different Rh–Ga ratios. After filtration, the products were dried at 130 °C and used in the catalytic experiment without any further treatment (for details see Supporting Information).

PDH Experiments. The PDH experiments were carried out in a tubular quartz reactor. The reactor was heated to the desired reaction temperature under a flow of an inert gas. When the reaction temperature was reached, a flow of propane was added to the He flow, resulting in a catalyst bed contact time of 0.7 s and a gas hour space velocity (GHSV) of 4900 h⁻¹. The effluent gas flow was analyzed with an online-GC (see Supporting Information for details).

Figure 1 shows the comparison of Rh on alumina with the Ga–Rh alloys of different compositions (as prepared in this work) at 550 °C. Rh on alumina displays under these conditions an initial productivity of 15 gpropylene gRh⁻¹ h⁻¹ that remains stable over 15 h time-on-stream. The conversion with the Rh/AlO₃ catalyst is around 4% with propylene selectivity rising from an initial 12% level to close to 90% after 15 h time-on-stream. Note, that the empty reactor shows a conversion of around 0.3% with a propylene selectivity of 35% at 550 °C (propane flow: 8.9 mL min⁻¹, He flow: 89 mL min⁻¹, see Supporting Information for further details).

As the baseline experiment, we also evaluated the catalytic activity of Ga on AlO₃ at 550 °C in PDH. It was found that the initial conversion was 4% with a propylene selectivity of around 95%. Note, that gallium oxide has recently been described as an active catalyst for PDH.12,13 We cannot exclude that some amount of gallium oxide is present in the precious metal-free, Ga on the AlO₃ material in the form of a passivation layer of the Ga droplets adsorbed on the support, and that this amount caused the observed catalytic activity. However, it is known from the literature that in the presence of noble metals under the reductive conditions of PDH experiments, gallium oxide species are reduced to elemental Ga.14,15 Indeed, we find clear evidence from our XPS and IR spectroscopy experiments that gallium oxide species are reduced to elemental Ga in the presence of Rh (see below and Supporting Information). All Ga–Rh alloys under investigation showed significantly higher initial conversions, with ~20% (Ga₃₄Rh), and ~29% (Ga₁₂₅Rh).

FIGURE 1

Figure 1. Conversion (top) and Rh-based productivity (bottom) of different Ga–Rh-SCALMS in PDH at 550 °C and 1.2 bar. Molar ratios of Ga–Rh are 0 (open circles), 34 (downward-pointing triangles), 89 (upward-pointing triangles), and 125 (diamonds). Reaction conditions: 1.2 g catalyst (Rh/Al₂O₃: 0.19 wt % Rh; Ga₃₄Rh: 0.26% Rh, 5.90% Ga; Ga₈₂Rh: 0.11% Rh, 5.9% Ga; Ga₁₂₅Rh: 0.07% Rh, 5.94% Ga), He flow 89 mL min⁻¹, C₃H₈ flow 8.9 mL min⁻¹, GHSV 4900 h⁻¹.

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Comparing the catalytic performance of the Ga–Rh alloys with maximum and minimum compositions ratios (125 vs 34) at 550 °C, a remarkable difference in productivity is observed.

This change in alloy composition resulted in a productivity increase by a factor of 4–5. The maximum initial activities were 263 gpropylene gRh h⁻¹ versus 49 gpropylene gRh h⁻¹ for Ga₁₂₅Rh and Ga₃₄₄Rh, respectively, and the productivities after 15 h time-on-stream were 127 gpropylene gRh h⁻¹ versus 31 gpropylene gRh h⁻¹, respectively. Selectivities were very similar in all experiments with Ga–Rh alloys: typically around 92% for conversions between 10 and 20% (see Supporting Information for details). All SCALMS material showed slight deactivation over time. No apparent changes in the catalyst morphology could be seen by means of SEM imaging, for example droplet coalescence with consequent loss of the active surface (see Supporting Information, Figure S3). However, deactivation due to morphological changes cannot be completely excluded.

On the other hand, the observed deactivation could be attributed to the formation of coke under reaction conditions, that is, at 550 °C. Indeed, XPS of spent catalysts showed an increase of the C 1s component in all samples when compared to the as-prepared materials (see Supporting Information, Figure S11). In addition, the presence of coke in SCALMS after PDH was confirmed using Raman spectroscopy (see Supporting Information, Figure S10). However, the moderate fluorescence of Al₂O₃ due to the applied laser wavelength prohibited the clear identification of coke in spent SCALMS after PDH at a temperature ≤ 500 °C, suggesting a minor formation of carbon on these catalysts under these milder conditions. Remarkably and in strong contrast to the tested SCALMS, the formation of coke was clearly observable for the Ga-free Rh catalyst after PDH, already at the lowest temperature tested of 450 °C (i.e., indicating the deposition of a sufficient amount of coke to quench the fluorescence of Al₂O₃). In other words, although the formation of coke could not be completely suppressed, the SCALMS materials displayed an increased resistance against coking compared to the pure Rh catalyst used as the benchmark. In fact, the formation of a detectable amount of carbon on SCALMS could be only observed under the most severe conditions applied, that is, running the PDH at 550 °C. Accordingly, we suggest that the observed conversion with the Ga-free Rh catalyst is just the residual activity after substantial coking deactivation. While the SCALMS systems, conversely, show strongly enhanced performance over a long time on stream operations, as the poisoning by coke formation is significantly reduced (Figure 1).

**XPS Investigations.** The XPS measurements of the supported alloy samples, before and after catalytic operation, clearly indicate the formation of metallic Ga and Rh species [Ga, Rh, or intermetallic compounds (IMCs)] during the reaction (see Supporting Information for details). Moreover, to understand the surface behavior/composition of Rh–Ga alloy droplets, we also studied model alloy samples with compositions comparable to those screened in the catalytic tests, as a function of temperature under ultrahigh vacuum conditions.

In Figure 2, the temperature-dependent composition, as obtained from the quantitative analysis of the XP spectra, is shown for the three different mixtures Ga₁₆₆₆Rh, Ga₂₃₄₄Rh, and Ga₄₉₄₄Rh (0.6, 1.2, and 2.0 at. % Rh in Ga); for exemplary XP spectra collected from a sample of composition Ga₂₃₄₄Rh see the Supporting Information, Figure S13. The measured composition is representative for the composition of the liquid phase because of the rather high information depth in lab-based XPS (93 Å for Ga 3d, 77 Å for Rh 3d in Ga). The temperature-dependent change of Rh intensity is ascribed to the formation or melting of Rh-rich IMC, which form in the bulk of the Ga–Rh droplet or at the droplet interface with alumina (see also microscopy results below) at lower temperatures. These solid IMCs phases, present in the bulk of the liquid have macroscopic dimensions and are not probed by the surface-sensitive XPS measurement of the droplet. A similar behavior and analysis was recently reported for the Ga–Pd system at low Pd concentrations. For compositions of Ga₃₆₆₆Rh and Ga₂₃₄₄Rh (0.6 and 1.2 at. % Rh in Ga), the respective plateaus at 480 and 530 °C indicate that the liquidus temperature (where all IMC are dissolved, and the XPS data thus give the nominal bulk concentration) is reached, as seen in Figure 2. Below the liquidus temperature, the composition of the liquid phase is given by the concentration on the liquidus line. Here, we observe a qualitative agreement with the bulk phase diagram of Rh–Ga, and the bulk phase diagrams of comparable systems (Pt–Ga, Pd–Ga). For Ga₃₆₆₆Rh, the measurement at the highest temperature also reached the nominal bulk composition.

**Microscopic Investigations.** To further elucidate the structure, composition, and phase stability of the Ga–Rh systems under investigation, microscopy investigations on Ga–Rh model nanoalloys were carried out with transmission electron microscopy (TEM), either static at room temperature (RT) or in situ during heating using a furnace TEM holder. Together with the XPS analysis, these investigations are particularly valuable as the phase diagram of the Ga–Rh system is not safely established in the low Rh concentration range. Moreover, phase stability and melting points may be altered for bimetallic nanoparticles by nanoscale effects. TEM studies were performed on a series of model GaₓRh₁₋ₓ nanoalloys (with x = 16−160, i.e., Rh 0.6−6 at. %) supported on thin SiO₂ membranes. Details on the preparation of the alloy nanoparticles by physical vapor deposition are provided in the Supporting Information.

We first describe the structural properties at RT. A typical two-phase structure of crystalline (c-) precipitates in amorphous (a-) nanoparticles is found throughout all samples, and exemplary results from the sample with x = 36 (Rh = 2.7 at. %) are summarized in Figure 3a–c. By combining EDXS mapping, quantification, and careful analysis of the sharp diffraction rings in electron diffraction patterns, the crystal phase is identified as Ga₁₆₆₆Rh₁₋ₓ or Gaₓ₂₃₄₄Rh₁₋ₓ and their derivatives (see Supporting Information for details). The crystallites possess a laminate shape and show a preferential orientation.
which is likely due to the presence of the SiO2 membrane below. Stacking disorder can often be seen in the laminites (cf. Figure 3c white arrows). The Ga-rich particle matrix shows an amorphous contrast in the high-resolution TEM (HRTEM) image, which agrees with the broad diffuse rings in the electron diffraction patterns (see Supporting Information, Figure S15). With increasing Rh concentration from 0 to 6 at. % a systematic evolution of the particle size and shape toward a larger size with a more irregular shape is found (see Supporting Information, Figure S16).

From the sparse literature on Ga–Rh systems, the (equilibrium) melting temperature at the low Rh concentrations studied in the present work is expected to be in the range of 200–600 °C. To elaborate the phase stability of the c-GaRh/a-Ga–Rh nanoparticles, electron diffraction has been performed in situ while heating the samples in the TEM. Figure 3d,e show key frames from a series of in situ electron diffraction patterns taken from the samples with $x = 36$ (Rh = 2.7 at. %) and $x = 36$ (Rh = 2.7 at. %). Temperatures are indicated at the top-right corner of each pattern. See the text for details, complete time series (movie) of in situ SAED are provided in Supporting Information.

Density-Functional Calculations Elucidating the Mode of Action of the Ga–Rh SCALMS Systems in PDH. To gain further insight into the nature of the catalytic phase, its surface composition and the active site for the catalytic transformation, we carried out ab initio molecular dynamics (AIMD) simulations based on density-functional theory (DFT) with periodic boundary conditions. An estimation of the uncertainty of the AIMD simulations can be found in Supporting Information.

In detail, we studied the composition of the surface using a slab model with a tetragonal simulation box containing 180 atoms with a Ga–Rh ratio of 89 (178 Ga, 2 Rh atoms) and Rh initially located at the surface (one Rh at each side of the slab). The time evolution of the z-position of Rh atoms in the slab is shown in Figure 4a (green curve). It is obvious that already after a few picoseconds Rh disappears from the surface. This is in line with a simulation with the Ga–Rh ratio of nine (162 Ga, 18 Rh atoms) that shows a Rh surface depletion (see Supporting Information, Figure S21). However, in order to be catalytically active, the rare Rh surface atoms have to stay at the surface if a reactant or reaction intermediate is adsorbed on them. To probe this, we ran AIMD simulations starting with

Figure 3. Static (a–c) and in situ (d,e) TEM investigations of the structure, chemical composition, and temperature-dependent phase properties of Ga–Rh nanoparticles. (a) BF-TEM image of the particles prepared on the thin SiO2 membrane. The particles contain crystalline precipitates as revealed by dark diffraction contrast. (b) Composed Ga and Rh map extracted from a scanning TEM–EDX dataset confirming an increased Rh concentration (~12–15 at. %) in the areas showing precipitates. (c) HRTEM image of a single Ga–Rh particle, the dashed circles indicate the crystalline Ga–Rh phase. Key frames of electron diffraction patterns taken during an in situ heating experiment from the samples with (d) $x = 36$ (Rh = 2.7 at. %) and (e) $x = 36$ (Rh = 2.7 at. %). Temperatures are indicated at the top-right corner of each pattern. See the text for details, complete time series (movie) of in situ SAED are provided in Supporting Information.
Rh plus an attached adsorbate located at the surface. For all tested adsorbates, namely, one or two CO molecules (Figure 4), a propyl residue with one additional hydrogen atom, propylene, and a hydrogen molecule (see Supporting Information, Figure S22) the adsorbed species can retain Rh at the surface until it desorbs. After desorption of the adsorbed species, the Rh atoms immediately diffuse away from the surface. As will be shown later, DFT calculations suggest that at the end of the dehydrogenation, propylene is adsorbed to Ga atoms, whereas H₂ is bound to the catalytically active Rh atom. Once H₂ is desorbed, Rh immediately moves from the surface into the Ga matrix. The fact that adsorbates can hold dynamically appearing Rh atoms at the surface may lead to a certain enrichment of catalytically active centers during catalysis.

To further elucidate the mechanism of PDH with our Ga−Rh SCALMS catalyst, we constructed a solid model system consisting of a face-centered cubic (fcc)-Ga(100) surface with one Rh atom located in a slightly distorted fourfold surface hollow site to represent the isolated catalytically active center. We choose this system for two reasons: (1) the partial radial distribution function between Rh and Ga was calculated from the MD simulation for the slab model used in the MD simulations as described before. Here, the maximum of the peak corresponding to the first coordination sphere of Rh is located at a distance of 2.54 Å away from Rh. Once H₂ is desorbed, Rh immediately moves from the surface into the Ga matrix. The fact that adsorbates can hold dynamically appearing Rh atoms at the surface may lead to a certain enrichment of catalytically active centers during catalysis.

In addition, we calculated Bader charges for the liquid system used in the MD as well as the solid model system with the result that Rh in both cases is negatively charged (−0.83 e as average in the MD simulation, −0.82 e in the solid model). DFT geometry optimizations were performed in combination with the climbing image-nudged elastic band (cNEB) approach and the dimer method19,20 to determine reaction barriers between possible reaction intermediates (see Supporting Information for details). Additionally, we carried out calculations for free energies and barriers. The corresponding reaction profile is shown in the Supporting Information and agrees very well with the energies and barriers without free energy corrections. The corresponding reaction diagram for some potential pathways is shown in Figure 4c. Note that this is the first step to get insight into the basic reaction mechanism of the dehydrogenation at the atomic level using a specific slab model in conjunction with the cNEB technique to determine the transition-state energies. Further details of the reaction mechanism could be obtained in future work for example by advanced, computationally very demanding MD methods. In the initial step, the first C−H bond of the weakly bound propane (Figure 4c (a), E_ads ≈ 0.46 eV) breaks with an activation barrier of 1.42 eV leaving both H and the residual 2-propyl species bound to the surface Rh atom (b). The subsequent diffusion of hydrogen to an adjacent Ga atom (c) is calculated to have a larger energy barrier (0.66 eV) than the diffusion of the 2-propyl residue (d, 0.32 eV) because of the large affinity of hydrogen to Rh. The resulting geometry (d) is 0.76 eV more stable than the structure (c). To complete the dehydrogenation starting from (d), a rotation of 2-propyl is

Figure 4. Results from DFT calculations. (a) z-position (perpendicular to the surface) of Rh atoms in the liquid slab model (see main text and Supporting Information) as function of time. The Rh atoms were initially located at the two surfaces of the slab, one at the top and one at the bottom surface. The green lines correspond to a simulation without any adsorbate attached to the Rh atoms at the surface. The red and the blue lines correspond to a starting configuration for the simulations, where one and two CO molecules are adsorbed at each of the Rh atoms at the surface. On the right the Ga-density along the z-direction of the slab is shown for orientation. Horizontal lines indicate the first maximum of the Ga-density for the two surfaces. (b) Simulated vibrational power spectra for Rh(CO)₂ (upper part) and RhCO (lower part); snapshots of one corresponding geometry from the MD simulations are additionally depicted. (c) Reaction diagram for the dehydrogenation of propane on fcc-Ga(100) with one Rh ad-atom. Ga is depicted in blue, Rh in light red, C in black, and H in white. Reaction intermediates are indicated by blue lines and transition states by red lines.
The surface and uses its kinetic energy for the phase with high kinetic energy hits the reactive Rh center at follows: in the recombine and desorb as H_2. After the H_2 desorption, our matrix and can desorb. Hydrogen at the Rh center can barriers, which can easily be overcome at elevated temperatures. At the end of the reaction, propene di geometries, it is obvious that the more hydrogen is bound to Ga (g) requires a lower activation energy (0.29 eV) than hydrogen diffusion (f) (0.52 eV). Considering the final geometries, it is obvious that the more hydrogen is bound to Rh, the more energetically favorable is the structure. This suggests (i) that at the end of the dehydrogenation reaction, propylene is bound to Ga atoms and can diffuse away from the active Rh and desorb in the gas phase, while the resulting two H atoms bound to Rh recombine and desorb as H_2. From our mechanismic study, we conclude that the reaction proceeds as follows: in the first step, propane that comes from the gas phase with high kinetic energy hits the reactive Rh center at the surface and uses its kinetic energy for the first dehydrogenation step. The other reaction steps require smaller barriers, which can easily be overcome at elevated temperatures. At the end of the reaction, propane diffuses onto the Ga matrix and can desorb. Hydrogen at the Rh center can recombine and desorb as H_2. After the H_2 desorption, our AIMD simulations show that the Rh atom immediately moves away into the Ga matrix. To further investigate the origin of the catalytic activity of SCALMS, we calculated Bader charges for the liquid SCALMS model as well as the Rh(111) surface. We found that the Rh in the liquid Ga slab has an average charge of −0.83 e in the MD simulation, while in the Rh(111), the surface Rh atoms are obviously metallic Rh^0 species. We conclude that Ga plays the role of an electron donor for the more electronegative Rh. The negatively charged Rh species, in contrast to Rh^0, is a strongly active dehydrogenation catalyst. This also is one explanation why the Rh/AlO_3 catalyst shows a significantly lower conversion than SCALMS catalysts. Furthermore, this emphasizes the important role of the Ga matrix for the catalytic process.

Besides this, Ga also modifies the adsorption properties of the reactants. One of the main issues in dehydrogenation catalysis, for example for industrially used Pt/AlO_3 catalysts, is coking of the surface because the desired products like propene bind to the active sites too strongly and therefore desorption is hindered and further dehydrogenation or cracking reactions can occur, which lead to undesired side products and coking of the surface. This can be prevented by lowering the adsorption strength of the desired products, which in this case is achieved by the Ga matrix around the catalytically active center. Our calculated mechanism also shows that it is thermodynamically more stable for propene to diffuse away from the reactive Rh center to the Ga matrix, while hydrogen stays at Rh. Propene can then easily diffuse away from the Ga matrix. Again, this underlines the importance of the Ga matrix in the SCALMS system.

**Infrared Spectroscopy.** Finally, we performed in situ infrared spectroscopy to gain further information on the state of Rh in the catalytically active SCALMS. We probed the noble metal sites at the Ga−Rh interface with adsorbed CO and investigated Ga−Rh alloys of various compositions (Ga_{25}Rh, Ga_{48}Rh, and Ga_{125}Rh on Al_{2}O_{3}) monometallic Ga/Al_{2}O_{3} and Rh/Al_{2}O_{3} as a reference. Both experiments and DFT calculations showed that elemental Ga is inactive toward CO adsorption, while experimental data showed the presence of hydrides. In contrast, shows rich carbonyl chemistry, especially when supported on oxides. The two main features observed on such systems are CO bound to metallic Rh in an on-top fashion (Rh^0CO) and the atomically dispersed dicarbonyl Rh^1(CO)_2 formed via the oxidative disruption of metallic Rh particles. Most bands on the Rh and Rh−
Ga samples can be assigned based on previous studies (see assignments summarized in Table 1).

Here, we limit the discussion to samples with low Rh loading. Figure 5 shows the DRIFT spectra obtained from Ga$_{88}$Rh$_1$/Al$_2$O$_3$ and monometallic Ga$^{−}$Rh samples. The spectra acquired from Ga$_{25}$Rh$_1$/Al$_2$O$_3$ and Ga$_{125}$Rh$_1$/Al$_2$O$_3$ are provided in the Supporting Information (Figure S26) together with a detailed discussion of all DRIFT experiments.

For the Ga$_{88}^{−}$Rh$_1$/Al$_2$O$_3$ sample, the bands at 2024 and 1993 cm$^{-1}$ obtained at 450 °C are of particular interest. We applied AIMD slab calculations (see Supporting Information and Figure 4b) to elucidate the nature of these absorptions. Both RhCO and Rh(CO)$_2$ remain stable at the surface during the complete simulation (up to 45 ps). Additionally, vibrational power spectra (Figure 4b) were calculated. We found that the calculated spectra are in excellent agreement with the experimental spectrum for the Ga$_{88}^{−}$Rh$_1$ sample above 300 °C. They show a feature at 1993 cm$^{-1}$ for RhCO and two bands at 2002 and 1978 cm$^{-1}$ for the symmetric and asymmetric CO-stretching frequencies of Rh(CO)$_2$, respectively. Based on this data, we identify two possible explanations for the experimental observations. On the one hand, the signal at 2024 cm$^{-1}$ may be assigned to hydrides bound on metallic Ga (Ga$^{−}$H peak found at 2021 cm$^{-1}$ in the case of the monometallic Ga/Al$_2$O$_3$ sample, compare also Table 1), while the peak at 1993 cm$^{-1}$ originates from a CO stretching vibration of a RhCO species. On the other hand, the DFT results also are in agreement with the formation of a Rh(CO)$_2$ species. In the latter case, the two experimental peaks would be ascribed to the symmetric and asymmetric CO-stretching modes. From the experimental and theoretical data, both interpretations appear reasonable, as the peak positions of Rh(CO)$_2$ (symmetric CO stretch) and Ga$^{−}$H bands both lie around 2025 cm$^{-1}$ and the features may overlap.

The effect of site isolation on the CO-stretching frequency is evaluated by performing frequency calculations in harmonic approximation for CO at a Rh top-site on a perfect Rh(111) surface at different coverages. The results as shown in Table 1 are in line with previously calculated values$^{32,33}$ which show that the frequency is strongly coverage dependent. Compared to the experimental data, the DFT values are in excellent agreement at a high coverage, while being ∼20 cm$^{-1}$ too low at lower coverage. Therefore, we conclude that the main contribution to the redshift in the vibrational spectra of the Rh−Ga mixtures originates from the isolation of the Rh atoms.

The spectra obtained from the Ga$_{88}^{−}$Rh$_1$ and Ga$_{125}^{−}$Rh$_1$ (see Supporting Information, Figure S26) samples recorded at 400 and 450 °C show only two peaks, which are assigned to Ga$^{−}$H species and/or Rh(CO)$_1^{−}$ in liquid Ga. There is no evidence for adsorbed CO on Rh atoms interacting with the support or for the formation of larger Rh clusters. Thus, the thermal evolution of the DRIFT spectra provides further evidence for the presence of a liquid Ga phase with isolated Rh atoms exposed at the surface. Figure 6 illustrates our conclusions from infrared spectroscopy and DFT calculation for the different Ga$^{−}$Rh samples under investigation.

Discussion. Merging the information given for the phase behavior of the Ga$^{−}$Rh system,$^{11}$ the solubility data of Rh in Ga,$^{35}$ the analytical data and additional catalytic experiments described here (see Supporting Information for details), it is possible to correlate the physical states of the different materials at varying Ga/Rh ratios with their catalytic properties. In Figure 7, the productivity values obtained when applying different compositions of SCALMS at different temperatures are depicted versus the known phase diagram.

Figure 5. CO-stretching region of the DRIFT spectra obtained from Ga$_{88}$Rh$_1$/Al$_2$O$_3$ and monometallic Ga−Rh samples in an Ar atmosphere after treatment with CO. The temperature in °C and the scaling factors are indicated on the left and right side of each panel, respectively. All spectra were normalized and displayed as difference spectra as described in the Supporting Information.

Figure 6. Surface features observed on the Ga$_x$−Rh$_y$ SCALMS and on the Rh/Al$_2$O$_3$ and Ga/Al$_2$O$_3$ reference samples.
data and the data from our XPS experiments. According to our XPS and TEM data, systems with a Ga-Rh $\leq 33$ are characterized by the presence of solid intermetallic Rh–Ga phases, together with a Ga-rich liquid phase (vide infra) and can be expected to never reach a fully liquid state, at any of the temperatures tested. Conversely, the system with Ga–Rh $> 120$ is expected to be always present in the fully liquid phase under the applied reaction conditions forming a liquid alloy. The systems with $82 < \text{Ga–Rh} \leq 89$ may be defined as borderline cases as the liquidus curve is crossed at a temperature between 450 and 500 °C. Hence, these systems are expected to be fully liquid at 550 °C. Figure 7 shows that the productivities are always significantly higher in the case of fully liquid Ga–Rh SCALMS systems, irrespective of the applied reaction temperature (compare results at 480, 500, and 550 °C). In other words, the catalyst with Ga$_{33.5}$Rh showed the highest performance for the whole set of temperatures, as its fully liquid nature allows for the maximum presence of single Rh atoms at the liquid/gas interface (vide infra). The catalysts with Ga–Rh $\leq 33.5$, conversely, showed for all temperatures the poorest performance, as a significant part of the Rh is always present in solid intermetallic phases, showing lower catalytic activity in these systems.

**CONCLUSIONS**

While hardly any successful example of Rh-catalyzed alkane dehydrogenation catalysis is known, we show in this paper that supported Ga–Rh alloys are a very interesting class of PDH catalysts. Even small additions of Ga to alumina-supported Rh have a positive effect on the catalytic activity and selectivity. A strong boost of the Rh-induced dehydrogenation activity was found under conditions where the supported Ga–Rh alloy is entirely liquid.

Moreover, our paper sheds light on the chemical nature and working principles of Rh-based dehydrogenation catalysis in supported Ga–Rh alloys. From our AIMD simulations, we conclude that the liquid metal interface is highly dynamic whereby Rh appears at the interface only periodically and is trapped there for a longer time if propane is offered from the gas phase. Additionally, our calculations suggest a cooperative mode of action between Ga and Rh during catalysis with the activation of the propane at the Rh single-site atom, followed by transfer of the propyl rest to the Ga surface while recombination of two hydrides to H$_2$ occurs at the Rh single site. Our infrared studies confirm the single atom nature of Rh at the liquid interface.

All our results provide additional evidence that the specific nature of supported liquid Ga alloys offers very beneficial properties for dehydrogenation catalysis. This remarkable fact encourages us to extend this particular material strategy to explore other Ga alloy systems and substrates in our ongoing work.

**ASSOCIATED CONTENT**

1. Supporting Information: The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acscatal.9b02459.

Materials synthesis, reactor, and analytic description, SEM, XPS, and Raman characterization of catalysts, TEM, DFT methods, and DRIFT set-up description (PDF)

Video S1. Electron diffraction series of sample with $x = 36$, temperature 400–530 °C, 20X accelerated (AVI)

Video S2. Electron diffraction series of sample with $x = 105$, temperature 550–RT °C, 20X accelerated (AVI)

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**Notes**

The authors declare no competing financial interest.

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