Properties of Bulk In-Pt Intermetallic Compounds in Methanol Steam Reforming

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Heterogeneous catalysts are often complex materials containing different compounds. While this can lead to highly beneficial interfaces, it is difficult to identify the role of single components. In methanol steam reforming (MSR), the interplay between intermetallic compounds, supporting oxides and redox reactions leads to highly active and CO₂-selective materials. Herein, the intrinsic catalytic properties of unsupported In₃Pt, In₃Pt, and In₅Pt as model systems for Pt/In₃O₃-based catalytic materials in MSR are addressed. In₃Pt was identified as the essential compound responsible for the reported excellent CO₂-selectivity of 99.5% at 300 °C in supported systems, showing a CO₂-selectivity above 99% even at 400 °C. Additionally, the partial oxidation of In₅Pt revealed that too much In₂O₃ is detrimental for the catalytic properties. The study highlights the crucial role of intermetallic In-Pt compounds in Pt/In₃O₃ materials with excellent CO₂-selectivity.

Introduction

Intermetallic compounds are an interesting and promising class of materials for a broad range of catalytic reactions.[1–3] The altered electronic structure and geometric effects of these materials result in changed and often beneficial catalytic properties compared to their parent metals.[4] Among the most intensively studied reactions with intermetallic compounds as catalytic materials is methanol steam reforming [MSR, Equation (1)].[5–14] One of the major concerns in this reaction is suppressing CO formation via methanol decomposition [MD, Eq. (2)] or the reverse water gas shift reaction [rWGSR, Eq. (3)]. A high CO₂-selectivity enables the direct use of the product stream in a proton-exchange membrane fuel cell, while even a few ppm of CO inhibit the PEM catalyst.[15]

\[
\text{CH}_3\text{OH} + \text{H}_2\text{O} \rightleftharpoons 3 \text{H}_2 + \text{CO}_2 \quad (1)
\]

\[
\text{CH}_2\text{O} = 2 \text{H}_2 + \text{CO} \quad (2)
\]

\[
\text{CO}_2 + \text{H}_2 = \text{H}_2\text{O} + \text{CO} \quad (3)
\]

Among the different intermetallic catalytic materials for MSR, ZnPd is the most intensively investigated one.[19,10,16–23] Its high CO₂-selectivity is ascribed to the formation of ZnO patches on ZnPd particles, while a clean ZnPd surface was shown to be unselective towards CO₂.[16,17] The discrimination between the catalytic properties of ZnPD and a ZnPd/ZnO interface was achieved by determining the intrinsic catalytic properties of unsupported bulk ZnPd.[16]

The formation of oxide layers on the intermetallic particles was also identified for GaPd,[27] and the In-Pd system.[13,24–26] Especially for In₃O₃-containing materials, the role of partially reduced species or oxygen vacancies was additionally investigated in the hydrogenation of CO₂ to methanol,[27,28] the reverse reaction of MSR. This emphasizes the complex nature of catalytic materials consisting of oxide-supported intermetallic compounds.

The In–Pt system, despite being known as a promising class of catalytic materials since the early 21st century,[20] showing excellent CO₂-selectivity, was subject of only a few further studies in MSR.[29–32] Investigations on the surface structure of a Pt/In₃O₃/AI₂O₃ material concluded that the active surface consists of metallic platinum and partly reduced In₂O₃[32] and no intermetallic In–Pt compound was considered, despite the earlier work.[30] In our recent study on Pt/In₃O₃ aerogels however, a reactive equilibrium of In₃Pt and In₅Pt₃ with In₂O₃ was identified in the active and selective state, resulting in a highly complex mixture of three different compounds in the active sites.[33]

The high complexity of supported intermetallic materials often hinders the assignment of the catalytic properties to distinct intermetallic compounds. Due to these limitations of the supported materials, the compounds In₃Pt₃, space group Im3m, a = 9.4359 Å,[34] In₃Pt, space group Fm3m, a = 6.365 Å,[35]
and In₃Pt₂, space group P3m1, \( a = 4.53 \text{ Å}, c = 5.51 \text{ Å}\) are synthesized as model systems, similar to the approach on ZnPd. This potentially allows separating the catalytic properties of the intermetallic compounds from the intermetallic/oxide interface if no oxidation occurs. Thus, it enables identifying the intermetallic compound being responsible for the high CO₂-selectivity in the supported systems in an ideal scenario or at least reduces the complexity, enabling a better differentiation of different active components. The materials were characterized concerning their phase composition by X-ray diffraction (XRD), their elemental composition by inductively coupled plasma with optical emission spectroscopy (ICP-OES) and their thermal behavior under MSR conditions by operando thermogravimetry coupled with mass spectrometry (TG/MS). To further correlate the obtained catalytic properties to the surface state, operando X-ray photoelectron spectroscopy (XPS) was conducted.

**Experimental**

**Material Preparation**

For the preparation of the bulk intermetallic In–Pt compounds In₃Pt₄, In₅Pt and In₃Pt₂, Pt-foil (ChemPur, 99.99%) was cut and weighed in a glovebox (MBraun, O₂ and H₂O < 0.1 ppm). Afterwards indium granules were cut and weighed to achieve the targeted concentration of 70 at-%, 66.6 at-% and 60 at-% indium, respectively. The total sample mass was around 500 mg for each compound. The metals were then transferred into a quartz glass ampoule and evacuated to a pressure below 2.0 × 10⁻⁶ mbar. Subsequently, the ampoules were re-filled with Ar (AirLiquide, 99.999 %) to 0.5 bar and sealed off. Afterwards the samples were molten in a furnace at 1200 °C for one day and quenched in water. The obtained ingots were annealed at 800 °C for 60 days to obtain the target phases.

**Characterization**

Elemental analysis was conducted via inductively-coupled plasma/optical emission spectroscopy (ICP-OES, Varian Vista RL). The samples were dissolved in freshly prepared aqua regia (hydrochloric acid, 37 wt-%, nitric acid, 68 wt-%, 3:1 ratio, VWR chemicals AnalaR) and diluted to 5 vol-% acid with deionized water. The prepared samples were measured in triplicate.

Phase analysis was conducted via powder X-ray diffraction (XRD, Enraf Nonius FR590) with monochromatic X-rays (Cu Kα, λ = 1.54060 Å, Ge (111) monochromator) on a zero-background Si single-crystal sample holder in Bragg-Brentano geometry. The samples were crushed in an agate mortar until the metallic luster was not visible anymore. The obtained powder was re-annelled in a evacuated and sealed quartz glass ampoule containing 0.5 bar Ar for 1 h at 800 °C and subsequently quenched in ice-water prior to the XRD measurements to release the stress from crushing. Some samples were prepared with grease to enable preparation of the sample holder, resulting in a increased background at low 2θ values. Operando thermogravimetric experiments coupled with mass spectrometry (TG/MS, Netzsch STA 449 F3 Jupiter, Pfeiffer Omnistar) were conducted with 150–200 mg of crushed and sieved samples with a diameter < 20 µm inside an Al₂O₃ crucible. Prior to the measurements the powder was reduced in situ at 400 °C with 5% H₂/He (AirLiquide, 99.999%, 40 mL/min) for 1 h. The samples were heated to 160 °C under 40 mL/min He-flow with 5 K/min. After 30 minutes of equilibration, 40 mL/min of 10 vol-% methanol-water-vapor mixture (1:1 atomic ratio, 0.194 g/l liquid flowrate, Fisher Scientific, HPLC grade) in helium were injected into the apparatus and the samples were heated to 500 °C with a heating rate of 1 K/min followed by an isothermal segment of 1 h. Ion currents for fragments of H₂, CO and CO₂ were recorded utilizing m/z = 2, m/z = 28 and m/z = 44, respectively. TG and MS curves were background-corrected by subtraction of a blank measurement under identical conditions.

XPS investigations were conducted at the ISISS beamline at BESSY II. Details of the experimental setup are described in reference. For the sample preparation, 150–200 mg of the crushed material were pressed to pills with 8 mm diameter in air with a pressure of 4 tons and 1 min holding time. After generating the pills, the samples were reduced in 5% H₂/He at 350 °C for 1 h with a flow of 40 mL/min at atmospheric pressure. Further handling and storage of the samples was done under argon atmosphere. Operando measurements were conducted at 0.5 mbar of a 1:1 mixture of water and methanol vapor at 400 °C. Ion currents for fragments of H₂, CO and CO₂ were recorded by mass spectrometry. A heating rate of 10 K/min was applied during heating and samples were equilibrated for 15 minutes prior to the XPS measurements.

**Catalytic Testing**

Catalytic investigations were conducted in a plug-flow reactor (PID Eng&Tech Microactivity Reference) with a micro-GC (Varian CP 4900, 10 m back flushed M5A column, 20 m back flushed M5A column and a 10 m PPU column) for the simultaneous analysis of H₂, CO and CO₂. For the catalytic testing, the samples were crushed and sieved. 150 mg of the sieve fraction of 20–32 µm, with a geometric surface area of roughly 0.02 m²/g, were mixed with 200 mg catalytically inactive graphite powder (ChemPur, < 100 µm, 99.9%). The prepared samples were placed on a quartz glass fleece inside of the reactor tube (SiO₂-coated stainless steel, inner diameter of 7.9 mm). A carrier gas flow of 10% He/N₂ at 15 mL/min, pre-heated to 120 °C, was mixed with an equimolar water/methanol vapor (0.01 mL/min H₂O, 0.0225 mL/min CH₃OH, Fisher scientific, HPLC grade). After the reactor all vapors were condensed in a cooling trap at 4°C and the gas flow further dried with a Naion membrane with a N₂-counterflow of 100 mL/min. Activity and selectivity were calculated according to Equations (4) and (5).

\[
a = \frac{n_1}{n_2} \cdot \frac{m_1}{m_2}\text{, with a \( m \) of } H_2 \text{ and Pt in Equation (4)}
\]

\[
S_{CO} = \frac{c_{CO}}{c_{CO} + c_{CO_2} + c_{H_2}}\text{, with } S_{CO} \text{ in Equation (5)}
\]

The chosen activity calculation allows comparison of the samples regarding the atomic efficiency of platinum. It also enables comparison of different materials without the need of the surface area. For the determination of the apparent activation energy \( E_a \), the natural logarithm of the conversion \( X \) was plotted against the reciprocal temperature.
Results and Discussion

XRD analysis of the samples shows that the three intended samples In₃Pt₂, In₂Pt and In₇Pt₃ were obtained as single-phase intermetallic compounds (Figure 1). No additional reflections were observed by XRD and all low intensity reflections originating from the ordering of the respective structure types are identified. Elemental analysis by ICP-OES confirmed the target elemental composition of the samples of 60 at-%, 66.67 at-% and 70 at-% indium with 60(1) at-%, 66(1) at-% and 70(1) at-%, respectively. Consequentially, three distinct intermetallic compounds were obtained, which have different structural and electronic properties.

To identify a suitable temperature range for the catalytic tests and potential oxidation of the investigated compounds, operando TG/MS was conducted with a 1:1 methanol-water mixture to simulate the catalytic conditions. As the catalytic tests aim to reveal the intrinsic catalytic properties of the individual intermetallic compounds, decomposition of them has to be avoided by choosing a suitable temperature regime at which the compounds are stable, if possible. Operando TG/MS from 160–500 °C with a heating rate of 1 K/min revealed no mass changes in the whole temperature range for In₃Pt₂ and In₂Pt. Applying such a low heating rate ensures to observe thermodynamically controlled material changes. In contrast, a continuous mass increase is observed for In₇Pt₃, starting as early as 200 °C (Figure 2). After 1 h at 500 °C, the mass gain equals 0.09(1) wt-%, which corresponds to an oxidation of 5% of the In₇Pt₃ into In₂Pt and In₂O₃ according to Equation (6) (see also Figure 3).

$$2 \text{In}_7\text{Pt}_3(\text{s}) + 3 \text{H}_2\text{O}(\text{g}) \rightarrow 6 \text{In}_2\text{Pt}(\text{s}) + \text{In}_2\text{O}_3(\text{s}) + 3 \text{H}_2(\text{g})$$

(6)

The evolution of the m/z = 2 signal, indicating hydrogen formation from MSR, shows an onset of 320 °C for In₃Pt₂ and deactivation is observed from 367 °C onwards. Since no mass changes were observed for In₃Pt₂, the deactivation is either caused by sintering of surface irregularities and/or healing of
surface defects or deposited (carbonaceous) species, which were also detected by C1s XPS. In₃Pt and In₇Pt₃ show almost identical behavior in the evolution of the m/z = 2 signal. The onset temperatures are 366 °C and 365 °C, respectively. Since In₃Pt₃ is oxidized under reaction conditions this indicates that the resulting species exhibit similar catalytic properties and temperature stability as In₃Pt. According to these findings, In₃Pt and In₇Pt₃ are less prone to deactivation at elevated temperatures than In₇Pt₃. Stabilization can be caused by the formation of surface oxides, which act as sintering inhibitors or stabilize surface defects. Despite the observed differences in onset temperature and deactivation behavior, catalytic testing is ideally conducted in the same temperature range to compare activity, selectivity and long-term stability. Thus, a maximum temperature of 400 °C was chosen for the catalytic tests of the three materials as compromise between limited deactivation of In₇Pt₃ and expected observability of catalytic activity of In₃Pt and In₇Pt₃.

Phase analysis by XRD of the samples after operando TG/MS measurements confirms the stability of In₇Pt₃ and In₃Pt (Figure 3). For these compounds, no additional phases were detected. In agreement with the mass increase in the case of In₇Pt₃, In₇O₃ and In₃Pt were identified here as additional phases. According to these findings, the catalytic properties of In₃Pt are expected to be greatly influenced by the formation of In₃Pt and In₇Pt while In₇Pt₃ and In₃Pt₃ are expected to exhibit their intrinsic catalytic properties.

Catalytic tests on the crushed materials were conducted from 200 to 400 °C with a heating rate of 5 K/min and 1 h holding time in 50 °C steps (Figure 4). After the initial heating, the samples were cooled down to 225 °C and heated to 400 °C again with the same heating protocol. By this, stable catalytic properties at different temperatures were achieved, allowing to identify temperature-induced differences. The low temperature regime of 200–300 °C was chosen to be investigated for potential low-temperature activity or activation after the initial heating, despite the higher onset temperature observed in the operando TG/MS measurements, since the samples have a much higher interaction with the gas flow in the flow reactor as in the TG/MS device. After the dynamic temperature profile, a 20 h isothermal segment at 400 °C is utilized to address the catalytic stability. In₇Pt₃ exhibits detectable catalytic activity from 350 °C onwards upon heating and reaches its maximum activity of 68 mol(H₂)/mol(Pt) h⁻¹ at 400 °C with a CO₂-selectivity of 90%. In the isothermal segment at 400 °C, a strong deactivation to 1 mol(H₂)/mol(Pt) h⁻¹ is observed during 20 h while the corresponding CO₂-selectivity is 94%. For In₇Pt₃, catalytic activity is observed from 400 °C onwards with a maximum activity of 30 mol(H₂)/mol(Pt) h⁻¹ and a CO₂-selectivity of 99.8%. In the subsequent isothermal segment, the activity drops to 4 mol(H₂)/mol(Pt) h⁻¹ and the CO₂-selectivity decreases slightly to 99.2%. In₃Pt₃ (together with In₇O₃) shows a maximum activity of 6 mol(H₂)/mol(Pt) h⁻¹ at 400 °C with a selectivity of 99.2%. In the isothermal segment these decrease to 2 mol(H₂)/mol(Pt) h⁻¹ and 97.5%, respectively. All materials exhibit a higher activity at low temperature after the initial heating up, indicating changes under catalytic operation, which might be formation of surface oxides (undetectable by TG/MS in the case of In₇Pt₃ and In₃Pt) or removal of carbonaceous deposits from atmospheric hydrocarbons.

From the temperature-dependent measurements, apparent activation energies of $E_a = 107(24)$ kJ/mol, $E_a = 112(32)$ kJ/mol and $E_a = 55(6)$ kJ/mol were determined by Arrhenius plots (Figure S1 in the Supporting Information) for In₇Pt₃, In₃Pt and In₇Pt₃, respectively. The limited accuracy of the obtained values for In₇Pt₃ and In₃Pt is due to the ongoing deactivation after the initial reaching of 400 °C. Obtained values for In₇Pt₃ and In₃Pt are in the same region as determined for supported In₃Pt on In₇O₃ ($E_a = 119(2)$ kJ/mol)[31,33]. In summary, In₃Pt exhibits the highest activity and CO₂-selectivity after 20 h at 400 °C, making it superior to In₇Pt₃ and In₇Pt₃. Compared to supported Pt/In₇O₃ at 300 °C with a maximum CO₂-selectivity of 99.5%[31,33], bulk In₃Pt keeps the same CO₂-selectivity even at 400 °C, which is above the WGSIR equilibrium of 98.7%, calculated according to Reference [38]. It can be concluded that the decomposition of In₇Pt₃ into In₇Pt and In₇O₃ leads to a less active and selective state in the form of an In₇O₃-enriched material. However, for In₇Pt and In₇Pt₃, surface sensitive analysis of sample composition is mandatory as decomposition of the near-surface region cannot be identified by XRD.

XPS measurements were conducted to ascribe the differences in the observed catalytic properties to the corresponding surface composition, with a focus on the formation of surface oxides. The as-prepared In₃Pt sample reveals an asymmetric signal in the Pt4f region with a binding energy of roughly 71.8 eV (Figure 5). The signal shape is in agreement with previous studies on In–Pt materials[32] while the binding energy is slightly higher than for Pt-richer intermetallic compounds.[39] The obtained total signal in the as-prepared state cannot be fitted with the chosen parameters for the Pt4f core-level alone. This deviation of the fitted signal to the experimental spectrum was only observed for the most surface-sensitive measurement under UHV conditions (see Figure S2 for comparison) and might be related to a high In-concentration at the surface, leading to

![Figure 4. Catalytic MSR tests (H₂O:MeOH = 1:1) on In₃Pt₃, In₇Pt and In₇Pt₃ under dynamic temperature from 200–400 °C. Activity is given in strong colors and selectivity in pale colors.](image-url)
imposing of the Pt4f signal on the In4p signal,\(^{32}\) The In3d signal was deconvoluted into a signal at low binding energy for the intermetallic compound, 444.2 eV, which is slightly higher than for previously reported intermetallic compounds containing indium,\(^ {41,42}\) and a signal for oxidic indium with a binding energy of roughly 444.8–445.0 eV, which is in the range of In\(_2\)O\(_3\) and In(OH)\(_3\).\(^ {42,43}\) Thus, in the as-prepared state, small amounts of oxidized indium species are present on the surface of In\(_2\)Pt. Similar results were obtained during the operando measurement. No changes in the Pt4f signal except for the removal of the underlying In4p signal are observed. In the In3d signal, a small shift to lower binding energy for the oxidic species is detectable, possibly resulting from the formation of partially reduced species under reaction conditions. In comparison to the UHV measurements, the relative amount of oxidic indium is decreasing, which confirms the stability of In\(_2\)Pt under MSR conditions, as no continuous oxidation of the bulk occurred. Since investigations on supported In\(_2\)Pt/In\(_2\)O\(_3\) showed that oxidic indium is actively participating in MSR,\(^ {33}\) the catalytic properties cannot be assigned to In\(_2\)Pt alone but are the result of In\(_2\)Pt, small amounts of oxidic indium and possibly a more Pt-rich (inter)metallic surface species, since In\(_2\)Pt is described as a line compound without significant homogeneity range.\(^ {44}\)

Almost identical results were obtained for In\(_3\)Pt\(_2\) (Figure 6). As for In\(_2\)Pt, a slightly asymmetric signal was obtained for the Pt4f core level. In addition, the In3d signal revealed minor surface oxidation in the as-prepared state, which does not increase upon exposure to reaction conditions. The presence of oxidic indium species can also be seen in the more bulk-sensitive measurements with a kinetic energy of 1080 eV (Figure S2). Thus, it can be concluded that the observed catalytic properties of In\(_3\)Pt\(_2\) are resulting from oxidic indium species and a Pt-enriched surface species on top of bulk In\(_3\)Pt\(_2\), and cannot be assigned to In\(_3\)Pt\(_2\) alone, analogous to In\(_2\)Pt.

Since both In\(_2\)Pt and In\(_3\)Pt\(_2\) show a slight surface oxidation in the as-prepared state and under reaction conditions, it has to be concluded that the excellent CO\(_2\)-selectivity of In\(_2\)Pt cannot be assigned to the formation of the surface oxides alone but depends strongly on the intermetallic compound. Besides the surface oxides, In\(_2\)Pt is mandatory to obtain an excellent CO\(_2\)-selectivity, thus making In\(_2\)Pt superior to In\(_3\)Pt\(_2\). However, using In\(_3\)Pt\(_2\) as precursor for In\(_2\)Pt and In\(_2\)O\(_3\) does not to lead to the excellent catalytic properties of bulk In\(_2\)Pt with slight surface oxidation. From this, it can be concluded that the amount of In\(_2\)O\(_3\) has to be limited on the In\(_2\)Pt surface to obtain excellent catalytic properties. The pronounced differences between in the catalytic properties of In\(_2\)Pt and In\(_3\)Pt\(_2\) clearly show that the presence of In\(_3\)Pt\(_2\) is an essential criterium for high CO\(_2\)-selectivity in catalytic In–Pt materials. In\(_3\)Pt\(_2\) as oxidation product of In\(_2\)Pt, is most likely also relevant for the high CO\(_2\)-selectivity but the obtained data clearly shows that In\(_2\)O\(_3\) and In\(_3\)Pt\(_2\) are not responsible for the excellent catalytic properties alone.
Conclusions

Three In-rich intermetallic compounds, In$_3$Pt$_2$, In$_2$Pt and In$_7$Pt$_3$, were synthesized as bulk materials and investigated regarding their catalytic properties and structural stability in methanol steam reforming. By operando TG/MS and XPS investigations, it was shown that In$_3$Pt$_2$ and In$_2$Pt are stable under reaction conditions and only exhibit slight surface oxidation, whereas In$_7$Pt$_3$ decomposes into In$_2$O$_3$ and In$_2$Pt. Upon linking these findings with supported In$_2$Pt/In$_2$O$_3$, the excellent CO$_2$-selectivity of more than 99\% of In$_2$Pt, which is significantly outperforming In$_3$Pt$_2$ and In$_7$Pt$_3$, can be ascribed to the presence of In$_2$Pt and a small amount of oxidic indium. Large amounts of In$_2$O$_3$ are detrimental to the activity and selectivity of In$_2$Pt, as observed for the strong decomposition in the case of In$_7$Pt$_3$. This study reveals that In$_2$Pt, in combination with small amounts of In$_2$O$_3$ and In$_3$Pt$_2$ as decomposition products, is responsible for the high CO$_2$-selectivity of In–Pt materials in MSR and confirms the high capability of intermetallic bulk materials to understand the intrinsic roles of different compounds in heterogeneous catalysts.

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Conflict of Interest

The authors declare no conflict of interest.

Data Availability Statement

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

Keywords: methanol steam reforming · intermetallic compounds · heterogeneous catalysis · renewable hydrogen · operando measurements
