Chemical Science

EDGE ARTICLE

Check for updates

Cite this: Chem. Sci., 2023, 14, 12739

All publication charges for this article have been paid for by the Royal Society of Chemistry

Received 6th September 2023 Accepted 2nd October 2023

DOI: 10.1039/d3sc04711c

rsc.li/chemical-science

Introduction

The non-oxidative propane dehydrogenation (PDH) reaction has become an intense field of research in both academia and industry, due to a steadily increasing demand for propene and the emergence of shale gas as a major chemical feedstock, resulting in a gap between offer and demand.¹ In order to address the supply of propene, industrial processes catalysed by K-CrO_x/Al₂O₃ (Lummus Catofin process, 1986), or Pt-based bimetallic systems such as K-Pt-Sn/Al₂O₃ (UOP Oleflex process, 1990),² Pt-Zn-Sn/CaO-Al₂O₃ (Star process, 1999)³ and K-Pt-Ga/Si-Al₂O₃ (FDCh process, 2016)³ have been developed for this highly energy intensive process ($\Delta H_{298K}^{0} = 124.3$ kJ mol⁻¹). These Pt-based systems are rather complex due to the use of various promoters and additives with

Implications of Ga promotion and metal—oxide interface from tailored PtGa propane dehydrogenation catalysts supported on carbon†

Enzo Brack, ^[]^a Milivoj Plodinec, ^[]^{ab} Marc-Georg Willinger^{‡b} and Christophe Copéret ^[]^{*a}

Propane Dehydrogenation is a key technology, where Pt-based catalysts have widely been investigated in industry and academia, with development exploring the use of promoters (Sn, Zn, Ga, etc.) and additives (Na, K, Ca, Si, etc.) towards improved catalytic performances. Recent studies have focused on the role of Ga promotion: while computations suggest that Ga plays a key role in enhancing catalytic selectivity and stability of PtGa catalysts through Pt-site isolation as well as morphological changes, experimental evidence are lacking because of the use of oxide supports that prevent more detailed investigation. Here, we develop a methodology to generate Pt and PtGa nanoparticles with tailored interfaces on carbon supports by combining surface organometallic chemistry (SOMC) and specific thermolytic molecular precursors containing or not siloxide ligands. This approach enables the preparation of supported nanoparticles, exhibiting or not an oxide interface, suitable for state-ofthe art electron microscopy and XANES characterization. We show that the introduction of Ga enables the formation of homogenously alloyed, amorphous PtGa nanoparticles, in sharp contrast to highly crystalline monometallic Pt nanoparticles. Furthermore, the presence of an oxide interface is shown to stabilize the formation of small particles, at the expense of propene selectivity loss (formation of cracking side-products, methane/ethene), explaining the use of additives such as Na, K and Ca in industrial catalysts.

> the possible formation of alloys and interfacial sites, which have been proposed to be driving the catalytic performance. In academic laboratories, numerous supported Pt-based bi- and multi-metallic model systems have been investigated with the goal to understand the effect of promoters, focusing primarily on Sn,² Zn,⁴ Ga,⁴ B,^{5,6} and In.⁷ However, most catalysts are quite ill-defined, partially due to the preparation methods, which often rely on (incipient) wetness impregnation – a technique, involving complex dissolution/precipitation processes and leading to the presence of numerous spectator species, rendering molecular-level understanding of promotional effects very challenging even when using advanced spectroscopic techniques or microscopy.

> In recent years, surface organometallic chemistry (SOMC) has emerged as a versatile tool to prepare silica-supported mono- and bi-metallic catalysts with tailored interfaces and compositions. This approach has been applied to study promoter effects in CO₂ hydrogenation and propane dehydrogenation reactions.⁸⁻¹⁰ In that context, silica-supported PtGa,¹¹ PtZn,¹² as well as PtMn¹³ nanoparticles have been developed and shown to be particularly active and selective in PDH. In these Ptbased systems, the major role of promoters has been linked to the formation of alloyed nanoparticles, which are responsible for improved propene selectivity and increased stability against



View Article Online

View Journal | View Issue

^aDepartment of Chemistry and Applied Biosciences ETH Zurich, Vladimir Prelog Weg 2/10, CH-8093 Zurich, Switzerland. E-mail: ccoperet@ethz.ch

^bScientific Center for Optical and Electron Microscopy (ScopeM) ETH Zurich, Otto-Stern-Weg 3, CH-8093 Zurich, Switzerland

[†] Electronic supplementary information (ESI) available. See DOI: https://doi.org/10.1039/d3sc04711c

[‡] Present address: Department of Chemistry TU Munich, Arcisstraße 21, D-80333 Munich, Germany.

Chemical Science

coking. Computational studies have inferred that the addition of Ga leads to Pt-site isolation and increases surface atom mobility, while the interaction with the oxide interface helps in stabilizing the PtGa nanoparticles against sintering.¹⁴ However, assessing these hypotheses and developing structure–activity relationships have been hindered due to experimental limitations, in particular charging effects in electron microscopy. Herein, we thus report the preparation and investigation of carbon-supported Pt and PtGa based catalysts, that unlock the use of state-of-the-art (scanning) transmission electron microscopy ((S)TEM), in order to probe the morphology of the supported nanoparticles and to evaluate the influence of the interface.

Towards this goal, we have developed a SOMC methodology on carbon¹⁵ using tailored molecular precursors to control the composition and the interface of the supported nanoparticles (Fig. 1A). Two different classes of organometallic precursors amidinate/alkyl vs. siloxide - were used as a strategy to prepare a set of Pt and PtGa based catalysts with a tailored interface on carbon. On the one hand, molecular precursors such as [GaMes₃]¹⁶ and platinum allyl-(N-N'-diisopropyl)acetamidinate ([PtAMD])17 are used for the preparation of carbon-supported Pt and PtGa nanoparticles, while thermolytic molecular siloxide precursors¹⁰ $[Ga(OSi(O^tBu_3)_3)_3(THF)]^{18}$ and $[Pt(OSi(O^tBu_3)_3)_2(COD)]^{19}$ are used to prepare the corresponding supported Pt and PtGa nanoparticles with an oxide interface (Fig. 1B). This approach based on molecular precursors containing or not a source of Si thus enables the study of the implications of Ga promotion and metal-oxide interfaces on PtGa PDH catalysts.

Results and discussion

Surface organometallic chemistry (SOMC) approach on carbon

As a first step to enable grafting, we introduce oxygenfunctionalities on carbon via a treatment of a mesoporous carbon support under oxidizing conditions, as previously reported.^{15,20} The treatment of mesoporous carbon in refluxing nitric acid for 12 hours leads to the incorporation of oxygen functional groups as evidenced from the O 1s XPS spectrum of the material after the treatment, revealing the presence of C=O, C-OH functional groups (Fig. S1 and S2[†]). The amount of protic functional groups (R-OH) is *ca.* 1.2 OH nm^{-2} (0.17 $mmol_{OH}$ g^{-1}), as determined by titration with $[Mg(Bn)_2(THF)_2]$; a value significantly lower than found for the corresponding carbon black support (0.41 mmol g^{-1} , 3.4 OH nm $^{-2}$).¹⁵ The difference in OH density most likely results from the highly graphitic structure of mesoporous carbon, rendering it more robust against oxidative conditions. Upon nitric acid treatment, the specific surface area of the carbon support slightly increases from 68 to 83 m² g⁻¹, as evidenced from BET analysis (Fig. S3[†]), likely as a result of etching.

Next, the functionalized carbon is used as a platform for SOMC, using two specific classes of molecular Pt and Ga precursors, with the aim to prepare two sets of carbonsupported Pt and PtGa PDH catalysts (Fig. 1B). First of all, isolated surface metal sites are obtained *via* grafting of a Ga or Pt precursor, followed by a thermal treatment under reduced pressure (Fig. 1B). Depending on the tailored precursor (siloxide



Fig. 1 (A) Classical SOMC/TMP approach on oxide supports for the preparation of supported nanoparticles. (B) Strategy for SOMC/TMP on carbon for the generation of carbon-supported nanoparticles with tailored interfaces and methodology for the preparation of carbon-supported (bi)metallic nanoparticles with or without a silica interface, depending on the (non) thermolytic nature of the tailored precursor.

or alkyl/amidinate), two sets (with and without a silica interface) of dispersed metal sites are prepared and denoted $Pt_x(_Si_z)/C_{grafted}$, where "x" and "z" indicate the measured weight loadings of the corresponding element from elemental analysis (EA) and the subscript "grafted" designates the material after grafting and thermal treatment under reduced pressure. Note, that the formation of isolated metal sites after the grafting step is confirmed via high-resolution high-angle annular dark-field (HAADF)-STEM imaging (Pt_{0.68}_Si_{0.1}/C_{grafted}) (Fig. S13[†]). From these highly dispersed metal sites in $Pt_x(Si_z)/C_{grafted}$, supported nanoparticles are formed via exposure to H₂ at 600 °C resulting in two monometallic materials: Pt_{0.68}_Si_{0.1}/C_{H2} and Pt_{0.60}/C_{H2} (Fig. 1B). The corresponding bimetallic materials are prepared and coined $Pt_xGa_v(_Si_z)/C_{H_z}$, where "y" additionally denotes the obtained Ga weight loading from EA. In order to prepare the bimetallic $Pt_xGa_v(Si_z)/C_{H_z}$ materials, dispersed Ga surface sites are first generated, followed by the introduction of Pt with a matching tailored precursor (siloxide or amidinate/alkyl) and finally a H₂ treatment to give $Pt_{0.26}Ga_{0.10}/C_{H_2}$ and $Pt_{0.89}Ga_{1.00}Si_{1.64}/C_{H_2}$. In order to ensure a more accurate comparison of the catalytic performances of the two sets of materials, an additional low-loading $Pt_{0.15}Ga_{0.10}Si_{0.31}/C_{H_2}$ material is prepared *via* substoichiometric grafting. All metal- and silicon-loadings of the catalysts correspond to the EA results of the materials after H₂ treatment and are summarized in Table 1.

EA shows that the bimetallic $Pt_xGa_ySi_z/C_{H_2}$ catalysts prepared using the Ga and Pt siloxide TMPs do indeed contain Si (Table 1), while the other materials do not contain any detectable Si, thus confirming the presence/absence of silicone. For $Pt_{0.68}$ _Si_{0.1}/C_{H₂}, the amount of Si is below detection limit (≤ 0.1 w%), due to the low amount of siloxide precursor decomposed in the preparation (grafting via protonolysis of the ligand, Fig. 1B). For the monometallic $Pt_x(Si_z)/C_{H_2}$ catalysts – Pt_{0.68}(_Si_{0.1})/C_{H2} and Pt_{0.60}/C_{H2} - very similar Pt weight loadings of 0.68 w% (0.25 Pt nm⁻²) and 0.60 w% (0.22 Pt nm⁻²) are obtained. On the other hand, the Pt and Ga weight loadings of the prepared bimetallic $Pt_xGa_y(_Si_z)/C_{H_2}$ catalysts vary to a greater extent. The $Pt_{0.89}Ga_{1.00}$ _Si_{1.64}/C_{H₂} and $Pt_{0.26}Ga_{0.10}$ /C_{H₂} catalysts differ significantly in terms of Ga weight loadings, with 1.00 w% Ga (1.04 nm^{-2}) vs. 0.10 w% Ga (0.10 nm^{-2}) , depending on the Ga molecular precursor used. The two catalysts also vary in terms of Pt content with 0.89 w% (0.33 Pt nm^{-2}) and 0.26 w% (0.10 Pt nm⁻²), respectively. The higher Pt loading of $Pt_{0.89}Ga_{1.00}$ _Si_{1.64}/C_{H₂} can be explained by the formation of an *in* situ formed silicate interface from the thermolysis of the Ga siloxide TMP ([Ga(OSi(O^tBu₃)₃)₃(THF)]), whose functional

groups can further be exploited for the grafting of the Pt precursor in the second grafting step. This is in line with a higher Pt loading for $Pt_{0.89}Ga_{1.00}Si_{1.64}/C_{H_2}$ (0.89 w% Pt, 0.33 nm⁻²) compared to the monometallic $Pt_{0.68}$ Si_{0.1}/C_{H₂} catalyst (0.68 w% Pt, 0.25 Pt nm⁻²) (Fig. 1B and Table 1). In the absence of silicon, the Pt content of the bimetallic $Pt_{0.26}Ga_{0.10}/C_{H_2}$ (0.26 w% Pt, 0.10 Pt nm⁻²) catalyst is lower with respect to the monometallic Pt_{0.60}/C_{H2} (0.60 w% Pt, Pt 0.22 nm $^{-2}$), as a result of the partial consumption of functional groups from the grafting of the Ga alkyl precursor ([Ga(Mes)₃]) and the absence of oxide formation. In order to assess the effect of the metal-oxide interface on the catalytic performance, a low-loading Pt_{0.15}Ga_{0.10}_Si_{0.31}/C_{H2} catalyst exhibiting a silicate interface with similar loadings (0.10 Ga nm⁻², 0.06 Pt nm⁻²) compared to the purely carbon-supported Pt_{0.26}Ga_{0.10}/CH₂ was prepared via substoichiometric grafting, with respect to the titrated protic R-OH functional groups.

Catalytic propane dehydrogenation (PDH)

We next evaluate the performance of the set of prepared catalysts - $Pt_{0.68}Si_{0.1}/C_{H_2}$, $Pt_{0.60}/C_{H_2}$, $Pt_{0.89}Ga_{1.00}Si_{1.64}/C_{H_3}$, $Pt_{0.26}Ga_{0.10}/C_{H_2}$ and $Pt_{0.15}Ga_{0.10}Si_{0.31}/C_{H_2}$ - in PDH under flow conditions (50 mL min⁻¹, 550 °C, $1:4 C_3H_8:Ar$) as shown in Fig. 2 and summarized in Table 2. As a consequence of their different Pt loading, the catalysts vary in terms of activity. Thus, the amount of catalyst is adjusted for each run, in order to be in a similar conversion range, while still being far from equilibrium conversion levels (\sim 40% conversion at 550 °C). From the catalytic data, it can be seen, that the bimetallic $Pt_xGa_{\nu}(Si_z)/C_{H_a}$ catalysts, display relatively good activity (8-3% conversion) and stability over the course of the reaction (>35 hours on stream), with a much-increased stability compared to the monometallic $Pt_x(Si_z)/C_{H_z}$ catalysts, as analogously observed for the silicasupported counterparts (Fig. 2).11 After reaching a stable initial state after 1 hour on stream, the deactivation behavior of the bimetallic catalysts (Pt_{0.89}Ga_{1.00}Si_{1.64}/C_H, $k_{\rm D} = 0.013$ h⁻¹; $Pt_{0.26}Ga_{0.10}/C_{H}$, $k_D = 0.015 h^{-1}$ slightly deviate from one another while both still displaying a slower deactivation compared to the silica-supported counterpart $PtGa/SiO_2$ ($k_D =$ 0.04 h^{-1}).¹¹ The low-loading Pt_{0.15}Ga_{0.10}Si_{0.31}/C_{H₂} showed a deactivation constant of $k_{\rm D} = 0.015 \text{ h}^{-1}$, thus lower compared to the one of $Pt_{0.89}Ga_{1.00}Si_{1.64}/C_{H_2}$.

This suggests, that the difference in deactivation behavior between $Pt_{0.89}Ga_{1.00}$ _Si_{1.64}/C_{H₂} and $Pt_{0.26}Ga_{0.10}/C_{H_2}$ could be related to the higher Pt to Ga ratio (0.3 *vs.* 0.5) of the $Pt_{0.26}Ga_{0.10}/C_{H_2}$ catalyst, rather than an effect caused by the presence of an oxide interface, as recently suggested in PtGa based dehydrogenation of *n*-heptane.²¹ Overall, when comparing the initial propene

Table 1 Summary of Elemental Analysis (EA) results of the prepared catalysts											
Sample	Pt [w%]	Ga [w%]	Si [w%]	$Pt [nm^{-2}]$	$Ga \left[nm^{-2} \right]$	Si [nm ⁻²]					
Pt _{0.68} _Si _{0.1} /C _{H₂}	0.68	n.d.	$0.1 \le$	0.25	n.d.	$0.3 \leq$					
$Pt_{0.60}/C_{H_2}$	0.60	n.d.	$0.1 \leq$	0.22	n.d.	$0.3 \leq$					
Pt _{0.89} Ga _{1.00} _Si _{1.64} /C _{H₂}	0.89	1.00	1.64	0.33	1.04	4.2					
$Pt_{0.15}Ga_{0.10}Si_{0.31}/C_{H_2}$	0.15	0.10	0.31	0.06	0.10	0.8					
Pt _{0.26} Ga _{0.10} /C _{H₂}	0.26	0.10	$0.1 \le$	0.10	0.10	$0.3 \leq$					



Fig. 2 (Left) conversion vs. Time on Stream (TOS) and (right) product selectivities after 20 hours on stream of $Pt_{0.68}$ - $Si_{0.10}/C_{H_{2'}}$ $Pt_{0.60}/C_{H_{2'}}$ $Pt_{0.89}Ga_{1.00}$ - $Si_{1.64}/C_{H_2}$, $Pt_{0.26}Ga_{0.10}/C_{H_2}$ and $Pt_{0.15}Ga_{0.10}$ - $Si_{0.31}/C_{H_2}$ in PDH under flow conditions (50 mL min⁻¹, 1: 4 C₃H₈: Ar, 550 °C).

productivity of the silicate decorated Pt0.15Ga0.10_Si0.31/CH2 $(904 g_{C_3H_c}/(g_{Pt} h); 4192 mol_{C_3H_c}/(mol_{Pt} h))$ to the purely carbon-supported $Pt_{0.26}Ga_{0.10}/C_{H_2}$ (486 $g_{C_2H_2}/(g_{Pt} h)$; 2254 mol_{C.H.}/(mol_{Pt} h)) catalyst, an enhanced initial propene productivity is observed, likely as a result of the presence of an oxide interface (Table 2 and Fig. S4[†]). From comparison of the product selectivities after 20 hours on stream, a lower selectivity towards propene along with an increased formation of cracking products (Fig. 2), ethene and methane, is observed for Pt_{0.26}Ga_{0.10}/C_{H₂} compared to the low-loading Pt_{0.15}Ga_{0.10}Si_{0.31}/C_{H₂} (where an oxide interface is present, Fig. S7†). The shift in selectivity can be attributed to the presence of a metal-oxide interface, since upon increasing the Pt to Ga ratio, the formation of cracking products is enhanced, as evidenced from the comparison of the product selectivities of Pt_{0.89}Ga_{1.00}_Si_{1.64}/C_{H2} and Pt_{0.15}Ga_{0.10}Si_{0.31}/C_{H2} and Pt_{0.68}Si_{0.1}/C_{H₂} (Fig. 2). Similarly, comparison of the product selectivities of the monometallic $Pt_{0.68}$ _Si_{0.1}/C_{H₂} and $Pt_{0.60}$ /C_{H₂} catalysts, consistently shows a shift in selectivity towards ethene and methane upon presence of an oxide interface (Fig. 2 and S5[†]). Although typically regarded as an innocent support, these findings demonstrate, that the introduction of a silicate interface enhances the propene productivity, at the expense of promoted cracking activity, as observed from a selectivity shift towards cracking products. This shift in selectivity is likely related to the introduction of acidity from the deposited silicate interface, possibly explaining the use of Na, K and Ca in industrial catalyst formulations,^{2,3} presumably as a strategy to remove the acidic properties of the oxide support. Additionally, as previously reported, Ga is known to form isolated single-sites on SiO₂.¹⁸ Therefore, the migration of Ga during reaction conditions from Pt to the silicate interface, might additionally favor a decrease in propene selectivity of $Pt_{0.15}Ga_{0.10}$ -Si_{0.31}/C_{H₂} compared to $Pt_{0.26}Ga_{0.10}/C_{H_2}$.

Electronic structure and morphology of the $Pt_x(_Si_z)/C$ and $Pt_xGa_y(_Si_z)/C$ catalysts

In order to gain further insights on the electronic structure, the morphology, the composition and reasons behind catalytic performances, the set of catalysts is next studied *via* (*in situ*) XANES and high-resolution STEM measurements. *Ex situ* XANES of the silicate decorated materials after grafting $(Pt_{0.68}^{II}_Si_{0.1}/C_{grafted} \text{ and } Pt_{0.89}^{II}Ga_{1.00}^{III}_Si_{1.64}/C_{grafted})$ as well as after treatment under a flow of H₂ (Pt_{0.68}^{II}_Si_{0.1}/C_{H_2} and Pt_{0.89}^{II}Ga_{1.00}^{III}_Si_{1.64}/C_{H_2}) (Fig. S8–S10†) are found to be consistent with previously reported findings on the silica-supported counterpart PtGa/SiO₂.¹¹ Prior to reduction, the Pt L₃-Edge (Fig. S8†) and Ga K-edge (Fig. S10†) energies of

Sample	Time [h]	Conversion [%]	Selectivity [%]	$\begin{array}{l} \text{WHSV} \\ [g_{\text{C}_3\text{H}_8} / (g_{\text{Pt}} \ h)] \end{array}$	Initial productivity $[g_{C_3H_6}/(g_{Pt} h)]$	Carbon Balance	$k_{ m D} [{ m h}^{-1}]$
$Pt_{0.68}Si_{0.1}/C_{H_2}$	0.03	8	89	2536	181	>99%	0.07
	35	1	65				
$Pt_{0.60}/C_{H_2}$	0.03	6	81	1434	68	>99%	0.05
	35	1	70				
$Pt_{0.89}Ga_{1.00}_Si_{1.64}/C_{H_2}$	1	8	98	2315	179	>98%	0.013
	35	4	95				
$Pt_{0.26}Ga_{0.10}/C_{\rm H_2}$	1	6	99	3467	486	>99%	0.015
	35	3	94				
PtGa/SiO ₂ (ref. 11)	0.3	32	>99	2243	721	>95%	0.04
	20	17	>99				
$Pt_{0.15}Ga_{0.10}_Si_{0.31}/C_{H_2}$	1	9	95	7202	903	>99%	0.015
	35	5	90				

Table 2Summary of catalytic propane dehydrogenation tests under flow conditions (50 mL min⁻¹, 1: 4 C_3H_8 : Ar, 550 °C)

Pt_{0.68}^{II}_Si_{0.1}/C_{grafted} and Pt_{0.89}^{II}Ga_{1.00}^{III}_Si_{1.64}/C_{grafted} are identical with their silica-supported counterpart at 11 566 eV and 10 374 eV, respectively, confirming the preservation of the oxidation states (Pt^{II}, Ga^{III}) of the molecular precursors upon grafting and formation of dispersed sites. The same is observed for the catalysts in the absence of an oxide interface. After reduction, the white line intensity decreased along with a shift of the edge energy from 11 566 eV to 11 565 eV, consistent with a reduction of Pt from Pt^{+II} to Pt⁰. The decrease in white line intensity and energy shift at the Pt L₃-Edge are attributed to an electron transfer from a metallic phase of Ga to the Pt 5d band, leading to a shift in the central transition of the Pt 5d valence band with respect to the Fermi energy level, as observed for the silicasupported systems, thus indicating alloying of the two metals within the nanoparticles.11 The same can be observed from in situ XANES measurements at the Pt L₃-Edge of the grafted Pt containing catalysts under a flow of H₂ (Fig. 3, S11 and S12[†]). Notably, the reduction of Pt occurs at lower temperatures in the case of the bimetallic PtGa catalysts compared to the monometallic Pt ones. From in situ XANES measurements at the Ga K-Edge of the bimetallic $Pt_{0.89}{}^{II}Ga_{1.00}{}^{III}_Si_{1.64}/C_{grafted}$ and $Pt_{0.26}{}^{II}Ga_{0.10}{}^{III}/C_{grafted}$ catalysts under a flow of H₂ (Fig. 3A), the reduction of Ga is observed from a shift of the edge energy to lower energy. Combined with the information gained from in situ XANES measurements at the Pt L₃-Edge, it appears that the Pt is reduced at a lower temperature, while Ga being gradually reduced and intercalated into the Pt nanoparticles to form alloyed $Pt_xGa_v(Si_z)/C_{H_2}$ nanoparticles, as previously proposed

in a computational study on PtGa/SiO₂.¹⁴ Unfortunately, it is not possible to carry out analysis of the extended X-ray absorption fine structure (EXAFS) due to low weight loadings of the catalysts and consequential poor data quality of the extended fine structure, thus hampering EXAFS fitting. In order to obtain more insights on the alloving of the two metals within the prepared catalysts, they are studied via electron microscopy. The formation of Pt and PtGa nanoparticles in the case of the monometallic $Pt_x(_Si_z)/C_{H_2}$ (Fig. S16 and S21[†]) and bimetallic $Pt_xGa_v(Si_z)/C_H$ catalysts (Fig. 3B and S25[†]) was confirmed from STEM imaging and EDX elemental mapping. Quantitative analysis of the energy dispersive X-ray spectra of $Pt_{0.68}$ _Si_{0.1}/C_{H₂} (Table S1[†]) and $Pt_{0.89}Ga_{1.00}Si_{1.64}/C_{H_2}$ catalysts (Table S3[†]), indicate the presence of Si and confirm the EA results, validating the proposed formation of a silicate interface as a result of the decomposition of the siloxide ligands of the TMPs. The observed silicon most likely deposits in the form of SiO₂ domains in close spatial proximity to the nanoparticles, as indicated by the overlap of Si and O signals in the EDX maps (Fig. S16, S25 and S28[†]). In sharp contrast, no Si was detected (as expected) from the EDX spectra of Pt_{0.60}/C_{H2} (Tables S2 and S5[†]) and Pt_{0.26}Ga_{0.10}/C_{H₂} (Fig. 3B). These findings, strongly support the hypothesis that the observed Si domains in the case of Pt_{0.68}_Si_{0.1}/C_{H2} and Pt_{0.89}Ga_{1.00}_Si_{1.64}/C_{H2} are a result of the thermolytic decomposition of the siloxide containing ligands present in the TMP used in the grafting step(s). Notably, the absence of a silicate interface in Pt_{0.26}Ga_{0.10}/C_{H2} allows for the acquisition of a STEM-EDX map of a single nanoparticle. From



Fig. 3 (A) *In situ* Pt L₃ and Ga K-Edge X-ray Absorption Near-Edge Structure (XANES) spectra are shown for $Pt_{0.89}^{II}Ga_{1.00}^{III}$ _Si_{1.64}/C_{grafted} (left) and $Pt_{0.26}^{III}Ga_{0.10}^{III}/C_{grafted}$ (right), acquired under a flow of 1 barg of H₂ (10 mL min⁻¹) over a temperature range of 25 °C to 600 °C. (B) HAADF-STEM image of a single-nanoparticle, corresponding Energy Dispersive X-ray (EDX) maps of Ga L, Pt M, C K and EDX spectrum of $Pt_{0.26}^{II}Ga_{0.10}^{III}/C_{grafted}$. These maps reveal the spatial correlation between Pt and Ga, indicating homogeneous alloying of the two metals within the nanoparticle. The Cu and Fe signals in the EDX spectrum are associated with the polepiece of the microscope, while the Cu signals are related to the TEM grid.

the spatial superimposition of Ga L and Pt M X-ray peak signals (Fig. 3B) as well as from additional EDX maps (Fig. S25 and S31†), a homogeneously distributed alloy formation of the two metals within the particles is observed. Although, EXAFS fitting is not possible, the spatial correlation of the Ga L and Pt M signals from STEM-EDX mapping, combined with the insights gained from (*in situ*) XANES measurements are consistent with alloying between Pt and Ga. In order to get further insights on the implications of Ga promotion on the morphology and composition of the prepared catalysts, high-resolution STEM measurements are performed next (Fig. 4).

From comparison of the particle size distributions of the prepared catalysts, the formation of slightly smaller monometallic Pt nanoparticles compared to the silica-supported counterpart is observed,²² with a mean size of 1.5(6) nm for Pt_{0.68}_Si_{0.1}/C_{H₂} (Fig. 4A) and 1.5(9) nm for Pt_{0.60}/C_{H₂} (Fig. S19†). In contrast, the bimetallic nanoparticles of Pt_{0.26}Ga_{0.10}/C_{H₂} and Pt_{0.89}Ga_{1.00}_Si_{1.64}/C_{H₂} exhibit a mean particle size of 1.4(8) nm (Fig. S29†) and 1.2(5) nm (Fig. 4B) respectively, thus slightly larger than for its SiO₂₋₇₀₀-supported counterpart, with a mean of 1.0(2) nm.¹¹ These results confirm the formation of uniformly distributed nanoparticles in analogy to the silica-supported catalysts, while indicating the role of the oxide interface in the stabilisation of the nanoparticles during particle growth, as evidenced by the formation of smaller nanoparticles with a narrower size distribution upon increasing the Si content of the interface. Investigation of the spent catalysts show no apparent sintering effects, remaining similarly uniformly distributed supported nanoparticles after exposure to reaction conditions (Fig. S17, S22, S26 and S32†).

High-resolution HAADF-STEM images confirm the formation of crystalline Pt nanoparticles in Pt_{0.68}Si_{0.1}/C_{H₂} (Fig. 4C) and $Pt_{0.60}/C_{H_{c}}$ (Fig. S20[†]). Furthermore, images of a single Pt nanoparticle (Fig. 4C and E) and its corresponding fast-Fourier transform (FFT) (inset in Fig. 4E) reveal the atomic structure of a fcc structured Pt nanoparticle viewed along the [110] zoneaxis. Note, that the graphitic structure of the carbon support is preserved after the nitric acid treatment and SOMC/TMP approach, as evidenced especially from high-resolution annular bright-field STEM (ABF-STEM) images (Fig. S15⁺), showing the preservation of the graphitic layers. In sharp contrast, bimetallic PtGa/C nanoparticles are amorphous (Fig. 4D and F), according to the high-resolution HAADF-STEM measurements which were validated by 4D STEM. While the Pt catalysts consist of highly crystalline nanoparticles (Fig. S33 and S34[†]), only convergence-beam electron diffraction (CBED) disks for the graphitic carbon are observed (Fig. S35 and S36[†]) for the PtGa catalysts. The amorphous nature of the PtGa nanoparticles most likely arises from the intercalation of Ga into Pt nanoparticles during H2 treatment, forming



Fig. 4 Particle size distribution of (A) $Pt_{0.68}$ -Si_{0.1}/ C_{H_2} and (B) $Pt_{0.89}Ga_{1.00}$ -Si_{1.64}/ C_{H_2} . Representative high-resolution HAADF-STEM images of (C) $Pt_{0.68}$ -Si_{0.1}/ C_{H_2} and (D) $Pt_{0.89}Ga_{1.00}$ -Si_{1.64}/ C_{H_2} . Enlarged view of a single (E) Pt and (F) PtGa nanoparticle (obtained from red square regions in (C) and (D)). Insets in (E) and (F) show the fast Fourier transforms of a crystalline Pt nanoparticle viewed along the [110] zone-axis and an amorphous, alloyed PtGa nanoparticle, respectively.

Edge Article

a homogeneous, amorphous, alloy according to *in situ* XANES and STEM-EDX measurements, consistent with the proposed structure based on our earlier computational studies.¹⁴ The homogeneous distribution of Ga and Pt within the PtGa nanoparticles supports the proposed role of Ga promotion *via* stabilizing the nanoparticles against coke formation through Pt-site isolation.^{14,23} Notably, the PtGa nanoparticles with high Ga contents are suspected to be in liquid-like state under reaction conditions according to the PtGa phase diagram and as exploited in supported catalytically active liquid metal solution (SCALMS) propane dehydrogenation catalysts.^{24,25}

Conclusion

The transposition of SOMC from classical oxide supports to carbon has enabled the preparation of tailored catalysts suitable for detailed structural analysis, designed to probe the spatial distribution of elements and the morphology of nanoparticles assessed by electron microscopy. STEM-EDX studies confirmed the preparation of a set of catalysts exhibiting or not a siloxide interface in proximity to PtGa nanoparticles depending on the chosen tailored molecular precursors. In particular, a combination of XANES and STEM-EDX confirm a homogenous alloying of Pt and Ga, and high-resolution HAADF-STEM images show the amorphous nature of PtGa nanoparticles, sharply contrasting the formation of highly crystalline monometallic Pt nanoparticles on carbon prepared according to the same approach. Thus, Ga is shown to form amorphous, alloyed PtGa nanoparticles and thereby to enable Pt-site isolation and superior catalytic performance in PDH, as proposed in earlier computational studies on the silicasupported counterpart.¹⁴ Furthermore, the presence of a SiO₂ interface is shown to enhance the overall propene productivity, but also promotes the formation of cracking products, resulting in loss of propene selectivity. This is consistent with the acidic properties of oxide supports and the use of additional additives such as Na/K/Ca in industrial catalyst formulations.^{2,3} Additionally, while not necessary for the stabilization of the nanoparticles against sintering, the presence of an oxide interface helps to form smaller nanoparticles. Overall, this strategy based on the development of catalysts, composed of nanoparticles with defined composition and tailored interfaces on carbon, offers a promising platform for the investigation of promotional and support effects in thermal catalysis, and can be readily extended to electrochemical applications of multi-metallic nanoparticles dispersed on conductive supports such as carbon. We are currently exploring these avenues.

Data availability

The data supporting the findings of this study is openly accessible on Zenodo at https://doi.org/10.5281/zenodo.8402733.

Author contributions

E. B. and C. C. conceptualized the study and were responsible for the methodology. E. B. and M. P. undertook the

investigations. C. C. was responsible for the funding acquisition. C. C. and M. G. W. were responsible for supervision. E. B. and C. C. wrote the original draft. All authors edited the manuscript.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

C. C., E. B. and M. P. acknowledge NCCR Catalysis (Grant no. SNF 180544), a National Centre of Competence in Research funded by the Swiss National Science Foundation. The Swiss Light Source is acknowledged for beamtime at the SuperXAS beamline (Proposal no. 20222164). Kazutaka Sakamoto, Domenico Gioffrè and Sumant Phadke are thanked for help with XAS measurements. ScopeM, the Scientific center for optical and electron microscopy of ETH Zurich, is acknowledged for providing access to their instruments for the project no. 2658. The Mougel Group at ETH Zurich is thanked for access to the XPS. Dr Seraphine Zhang is acknowledged for the XPS measurements and corresponding data processing. Dr Lukas Rochlitz is thanked for fruitful discussions and help with XAS data processing.

Notes and references

- 1 D. Malakoff, Science, 2014, 344, 1464-1467.
- 2 J. J. H. B. Sattler, J. Ruiz-Martinez, E. Santillan-Jimenez and B. M. Weckhuysen, *Chem. Rev.*, 2014, 114, 10613–10653.
- 3 S. Chen, X. Chang, G. Sun, T. Zhang, Y. Xu, Y. Wang, C. Pei and J. Gong, *Chem. Soc. Rev.*, 2021, **50**, 3315–3354.
- 4 S. R. de Miguel, E. L. Jablonski, A. A. Castro and O. A. Scelza, *J. Chem. Technol. Biotechnol.*, 2000, 75, 596–600.
- 5 C. Byron, S. Bai, G. Celik, M. S. Ferrandon, C. Liu, C. Ni, A. Mehdad, M. Delferro, R. F. Lobo and A. V. Teplyakov, *ACS Catal.*, 2020, **10**, 1500–1510.
- 6 M. Aly, E. L. Fornero, A. R. Leon-Garzon, V. V. Galvita and M. Saeys, *ACS Catal.*, 2020, **10**, 5208–5216.
- 7 X. Liu, W.-Z. Lang, L.-L. Long, C.-L. Hu, L.-F. Chu and Y.-J. Guo, *Chem. Eng. J.*, 2014, **247**, 183–192.
- 8 S. R. Docherty, L. Rochlitz, P.-A. Payard and C. Copéret, *Chem. Soc. Rev.*, 2021, **50**, 5806–5822.
- 9 S. R. Docherty and C. Copéret, J. Am. Chem. Soc., 2021, 143, 6767–6780.
- 10 C. Copéret, Acc. Chem. Res., 2019, 52, 1697-1708.
- 11 K. Searles, K. W. Chan, J. A. Mendes Burak, D. Zemlyanov,
 O. Safonova and C. Copéret, *J. Am. Chem. Soc.*, 2018, 140, 11674–11679.
- 12 L. Rochlitz, K. Searles, J. Alfke, D. Zemlyanov, O. V. Safonova and C. Copéret, *Chem. Sci.*, 2020, **11**, 1549–1555.
- L. Rochlitz, Q. Pessemesse, J. W. A. Fischer, D. Klose,
 A. H. Clark, M. Plodinec, G. Jeschke, P.-A. Payard and
 C. Copéret, *J. Am. Chem. Soc.*, 2022, 144, 13384–13393.

- 14 P. A. Payard, L. Rochlitz, K. Searles, L. Foppa, B. Leuthold,
 O. V. Safonova, A. Comas-Vives and C. Copéret, *JACS Au*, 2021, 1, 1445–1458.
- 15 C. K. Mavrokefalos, N. Kaeffer, H.-J. Liu, F. Krumeich and C. Copéret, *ChemCatChem*, 2020, **12**, 305–313.
- 16 O. T. Beachley, Jr., M. R. Churchill, J. C. Pazik and J. W. Ziller, *Organometallics*, 1986, 5, 1814–1817.
- 17 C. Ehinger, X. Zhou, M. Candrian, S. R. Docherty, S. Pollitt and C. Copéret, *JACS Au*, 2023, 3(8), 2314–2322.
- 18 K. Searles, G. Siddiqi, O. V. Safonova and C. Copéret, *Chem. Sci.*, 2017, **8**, 2661–2666.
- 19 D. A. Ruddy, J. Jarupatrakorn, R. M. Rioux, J. T. Miller, M. J. McMurdo, J. L. McBee, K. A. Tupper and T. D. Tilley, *Chem. Mater.*, 2008, **20**, 6517–6527.

- 20 C. Moreno-Castilla, M. V. López-Ramón and F. Carrasco-Marín, *Carbon*, 2000, **38**, 1995–2001.
- 21 O. Sebastian, A. Al-Shaibani, N. Taccardi, U. Sultan, A. Inayat, N. Vogel, M. Haumann and P. Wasserscheid, *Catal. Sci. Technol.*, 2023, **13**, 4435–4450.
- 22 P. Laurent, L. Veyre, C. Thieuleux, S. Donet and C. Copéret, *Dalton Trans.*, 2013, **42**, 238–248.
- 23 Y. Nakaya, J. Hirayama, S. Yamazoe, K.-i. Shimizu and S. Furukawa, *Nat. Commun.*, 2020, **11**, 2838.
- 24 H. Okamoto, J. Phase Equilib. Diffus., 2007, 28, 494.
- N. Raman, M. Wolf, M. Heller, N. Heene-Würl, N. Taccardi, M. Haumann, P. Felfer and P. Wasserscheid, ACS Catal., 2021, 11, 13423–13433.