Heterogeneous alkane dehydrogenation catalysts investigated via a surface organometallic chemistry approach

Scott R. Docherty, Lukas Rochlitz, Pierre-Adrien Payard and Christophe Copéret*

The selective conversion of light alkanes (C₂–C₆ saturated hydrocarbons) to the corresponding alkene is an appealing strategy for the petrochemical industry in view of the availability of these feedstocks, in particular with the emergence of Shale gas. Here, we present a review of model dehydrogenation catalysts of light alkanes prepared via surface organometallic chemistry (SOMC). A specific focus of this review is the use of molecular strategies for the deconvolution of complex heterogeneous materials that are proficient in enabling dehydrogenation reactions. The challenges associated with the proposed reactions are highlighted, as well as overriding themes that can be ascertained from the systematic study of these challenging reactions using model SOMC catalysts.

1. Introduction

In recent years, the shift away from traditional Naptha-based cracking processes, due to the appearance of abundant source of light alkanes with the emergence of Shale gas, has resulted in a dramatic reduction in the production of short chain alkenes. In parallel, the abundance of light alkanes, coupled with an increasing global demand of light alkenes, has pushed the use of alternative technologies for the production of light alkenes, such as the direction conversion of alkanes to alkenes via dehydrogenation (Scheme 1a). This reaction is however highly endothermic (standard enthalpy: +124.3 kJ mol⁻¹), hence requiring high operating temperatures. For instance, equilibrium conversion (Xₑ) for propane is only circa 30% for working pressures of 1 bar(a) at 550 °C. Such harsh conditions results in increased side reactions such as (thermal) cracking and coke formation; these conditions also result in increased sintering of supported nanoparticles, leading to limited catalyst lifetimes. To overcome coke formation and catalyst deactivation, constant regeneration through oxidative de-coking and re-reduction cycles is required in industrial settings.

These aforementioned constraints highlight the challenge to develop selective and stable catalysts for the non-oxidative...
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Alkane dehydrogenation

(a) Dehydrogenation of light alkanes (thermodynamic constraints, equilibrium conversion of propane at 1 bar and 550 °C)
(b) General structure of CATOFIN\textsuperscript{CrO}_{x}/Al\textsubscript{2}O\textsubscript{3}\right) type catalysts 
(c) General structure of platinum-based catalysts for the dehydrogenation of light alkanes (PtM/MO\textsubscript{x}/Al\textsubscript{2}O\textsubscript{3}).

Scheme 1 (a) Dehydrogenation of light alkanes (thermodynamic constraints, equilibrium conversion of propane at 1 bar and 550 °C); (b) general structure of CATOFIN\textsuperscript{CrO}_{x}/Al\textsubscript{2}O\textsubscript{3}\right) type catalysts; and (c) general structure of platinum-based catalysts for the dehydrogenation of light alkanes (PtM/MO\textsubscript{x}/Al\textsubscript{2}O\textsubscript{3}).

dehydrogenation of light alkanes. Several processes have however been developed and implemented in industry for on-purpose production of short chain alkenes. Today three catalytic systems are currently employed commercially: the established ABB Lummus CATOFIN (CrO\textsubscript{x}/Al\textsubscript{2}O\textsubscript{3}) (Scheme 1b) and UOP OLEFLEX (Pt/Sn/Al\textsubscript{2}O\textsubscript{3}) processes (Scheme 1c),\textsuperscript{3,8} as well as the most recent process developed by Dow based on Pt/Ga/Al\textsubscript{2}O\textsubscript{3}.\textsuperscript{9} One may note that the analogous oxidative dehydrogenation processes have been more difficult to implement due to the high exothermicity of the reaction, and the difficulty to obtain high alkene selectivity at high conversion.

However, recent findings may lead to further development in this field.\textsuperscript{10,11}

Coming back to the non-oxidative alkane dehydrogenation process, while these aforementioned catalysts have been widely studied, the complex nature of the catalytic systems leaves many unanswered questions regarding the nature of the active site, strategies for the suppression of side reactions (cracking, coking), and how catalyst structure can influence selectivity in alkane dehydrogenation processes.

Thus, the development of realistic model systems is highly desired, with the goal of unravelling the nature of the active sites, the mechanism of this reaction, and to better understand deactivation processes. The increased demand of these processes and the emergence of strategies for the synthesis of well-defined sites and the control of interfaces between metal particles on metal oxide supports has recently triggered interest in this field of research. In this review, we limit our discussion to the development of molecular-level understanding of light alkane dehydrogenation using model catalysts synthesized via Surface Organometallic Chemistry (SOMC).\textsuperscript{12} SOMC, a precise molecular approach which exploits the formation of covalent bonds between molecular species and supports (through grafting), has emerged as a powerful technique to understand heterogeneous catalysis when used in conjunction with \textit{in situ} or \textit{operando} spectroscopy combined with computational approaches. This approach is particularly powerful to interrogate the structure of active sites and reaction mechanism (\textit{vide infra}).\textsuperscript{13–15} The review is divided into three main sections: (i) a section describing the SOMC methodology, and two additional sections related to the use of this methodology for the development of alkane dehydrogenation catalysts based on (ii) isolated metal sites on oxide supports that parallel the industrial Cr-based catalysts, and (iii) supported nanoparticles on doped oxide supports as the second important class of catalysts. This review will end with a view on the possible role of SOMC in this field and future research directions.

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His current research is dedicated to experimental and computational studies of reaction mechanisms targeting to overcome the limitations encountered in catalytic systems.

Christophe Copéret (CCH) obtained a PhD in 1996 with E.-i. Negishi (Purdue University) and was a postdoctoral fellow with K. B. Sharpless (Scripps Research Institute). CCH joined CNRS in 1998, became Research director in 2008 and moved to ETH Zürich in 2010, as Chair of Surface and Interfacial Chemistry. His research interest lies at the interface between molecular, surface and material chemistries. The objective of his laboratory is to bring a molecular-level understanding of complex inorganic systems such as heterogeneous catalysts, hence the development of Surface organometallic chemistry methodology as well as spectroscopic techniques such as solid-state NMR.
2. SOMC methodology

Surface organometallic chemistry has emerged as an effective tool for the generation of materials with well-defined surface sites.13–20 Through the design of molecular precursors, such as thermolytic molecular precursors (TMP),21,22 and rigorous characterization at every stage of synthesis, one is now able to selectively generate uniform supported, isolated species (so-called single sites) and supported metallic nanoparticles with a narrow particle size distribution, where the average particle size is largely independent of the dopant(s) and support – enabling direct comparison of physicochemical properties and catalytic performance.14 The precise molecular control conferred by this approach enables explicit study of the interplay between structure and activity, complementing, and deconvoluting, experimental observations from more complex materials synthesized using traditional synthetic approaches.

This approach has been exploited for the synthesis of isolated M$^{n+}$ sites dispersed on oxide supports. Through the selective reaction of homoleptic metal siloxides, hydrocarbyls, alkoxides and aluminates with hydroxyl groups found on the surface of oxides (henceforth described as grafting), it has been possible to generate isolated surface species which can be subsequently thermally treated, under reduced pressure, inert or even reactive conditions (e.g. synthetic air), to generate well-defined surface sites of known oxidation state, geometry and nuclearity (Scheme 2a).13,19 The materials generated can be considered surface-doped oxide materials, with effective metal utilization ($M_{\text{surface}} = 100\%$). This approach has been widely exploited for an array of transition metals (Ti, Zr, Hf, Nb, Ta, Cr, Mo, W, Mn, Fe, Co, Ni) as well as Cu, Zn, main group elements (Al, Ga, etc.) and lanthanides, where the strong M–O bonds allow grafting and retention of highly dispersed metal sites. In some cases, like Ni and Cu, oxidative treatment is needed to avoid reduction of the metal sites and to retain isolated metal sites.23,24

Secondly, SOMC can also be used for the synthesis of supported metal nanoparticles. Following the grafting of late-transition metal precursors (siloxides, hydrocarbyls, alkoxides,
amides etc.) through the surface hydroxyl groups of oxide supports, that enables the generation of materials with a uniform initial distribution of metal across the support, the treatment under hydrogen (typically 1 bar(a), in flow) at elevated temperatures yields small, uniform nanoparticles, free of organic ligands/capping agents (Scheme 2b). This approach has been exploited for metals which possess greater ‘noble’ character (Ir, Rh, Pt, Cu, Ag, and Au), though the lower M–O bond strength reduces the thermodynamic impetus for grafting, and renders the range of relevant precursors more limited.25–29

These two approaches can also be combined, exploiting the stability of doped oxide materials generated by SOMC (Scheme 2a), and further functionalizing these materials through the grafting of a second molecular precursor, through remaining/regenerated surface hydroxyls.14 Upon treatment under H₂, one of two outcomes occurs: (i) the reduction of the second grafted species to form monometallic nanoparticles, of uniform size, atop silica containing isolated metal ions (Scheme 2c); or (ii) the reduction of the second grafted species to form metallic nanoparticles, of uniform size, atop oxide supports containing isolated metal ions, with gradual reduction and intercalation of the second metal to the nanoparticle, forming an alloy (Scheme 2c).30–32 The use of this approach, in the absence of alloying enables the disambiguation of bulk/surface effects often described in heterogeneous catalysis, as well as explicit study of particle–support interfaces. Where alloying occurs, it is possible to study, in the absence of large volumes of spectator species (typically found in the bulk of pure oxides), the role of alloying in determining the reactivity for reactions catalysed by supported nanoparticles.

In addition, SOMC has also been shown to generate so-called single atom alloys as well as bimetallic bulk alloyed materials. In this approach, supported metal (M) nanoparticles, prepared via classical or SOMC approach, are reacted, typically under H₂, with organometallic compounds or molecular hydrides (E = Sn, Si, Ge, Pb, As).33–38 This approach involves (Scheme 2d) the selective reaction of surface hydrides present on the metallic nanoparticles under H₂ with the organometallic reagent to generate usually organometallic fragment covalently bound to the remaining/regenerated surface hydroxyls.14 Upon treatment in H₂, the second grafted species to form monometallic nanoparticles, of uniform size, atop silica containing isolated metal ions (Scheme 2c); or (ii) the reduction of the second grafted species to form metallic nanoparticles, of uniform size, atop oxide supports containing isolated metal ions, with gradual reduction and intercalation of the second metal to the nanoparticle, forming an alloy (Scheme 2c).30–32 The use of this approach, in the absence of alloying enables the disambiguation of bulk/surface effects often described in heterogeneous catalysis, as well as explicit study of particle–support interfaces. Where alloying occurs, it is possible to study, in the absence of large volumes of spectator species (typically found in the bulk of pure oxides), the role of alloying in determining the reactivity for reactions catalysed by supported nanoparticles.

3 Modelling supported metal oxides via SOMC

The CATOFIN process is based on chromium oxide dispersed on alumina, Cr₂O₃/Al₂O₃, that falls into a broader class of alkane dehydrogenation catalysts composed of transition-metal or main group metal oxides dispersed on redox-inert supports. This section will first start by examining SOMC models developed to understand the Cr-based CATOFIN catalysts, before looking at alternative systems based on other supported metal oxides.

3.1 Models for supported CrO₃

Regarding Cr-based catalysts, the identity of the active sites remains unknown, in part due to the range of Cr oxidation states detected in Cr₂O₃/Al₂O₃ under reaction conditions.40–48 In addition, the presence of mononuclear and polynuclear sites has been discussed. Both Cr⁴⁺ and Cr⁶⁺ have been proposed to be catalytically active for alkane dehydrogenation with a large consensus pointing to Cr⁶⁺ being the most relevant active species.45–56 Therefore, there has been a strong interest in generating isolated Cr sites of known oxidation state and nuclearity at the surface of tailored oxide supports, to help the identification of active sites in supported metal oxides. One approach is the combined SOMC/TMP approach (see previous section), which provides access to dispersed low coordinated metal sites on oxide supports with controlled oxidation state and nuclearity.13,19 For instance, SOMC/TMP was used to generate a series of isolated Cr⁶⁺ sites at the surface SiO₂, SiO₂–Al₂O₃, and Al₂O₃. More specifically, these materials were obtained by grafting Cr⁶⁺ molecular precursors followed by a subsequent thermal treatment under vacuum: Cr(OSi(OtBu)₃)₃(THF)₂ on dehydroxylated SiO₂-700,57,58 and Al₂O₃-700 as well as Cr(Al(OiPr)₃)₃,
on dehydroxylated Al\textsubscript{2}O\textsubscript{3}-700 (Scheme 3).\textsuperscript{43} This approach enables the formation of Cr\textsuperscript{III} isolated mononuclear sites on silica, silica-alumina and alumina.

In the specific case of silica, CO adsorption monitored by FT-IR, in combination with DFT, evidenced the presence of two types of Cr\textsuperscript{II}I sites: tri-coordinated sites containing three Si–O groups coordinated to a Cr\textsuperscript{III} center as well as tetra-coordinated sites featuring an additional siloxane bridge coordinated to the metal center (Scheme 4).\textsuperscript{58} The oxidation state of Cr after treatment was confirmed to be Cr\textsuperscript{III} by EPR and XANES. Such Cr\textsuperscript{III} sites are able to catalyse PDH, albeit with low activity (Table 1).

In contrast, Cr\textsuperscript{III} sites dispersed on alumina, independently of the molecular precursors (Table 1), show significantly higher activity in the dehydrogenation of propane. Their activity even exceeds that of catalysts prepared by classical impregnation methods (Table 1, entry 4), probably because of the increased concentration of active sites in the former. On the other hand, these single sites catalysts are significantly less selective, presumably due to the presence of Brønsted sites (e.g. Si–OH or Al–OH interacting with adjacent Lewis sites), favouring cracking. Further surface chemistry can be foreseen to tune the acidity of the surface towards the development of highly active and selective PDH catalysts.

To further understand the reactivity of these sites, the hydrogenation of propene, that has the advantage to take place at much lower temperatures than the reverse PDH reaction, was studied in detail. Kinetic investigation on Cr\textsuperscript{III} single-site catalysts show low activation energies (0–40 kJ mol\textsuperscript{-1}) while activation entropies are very negative (−145 to −194 J K\textsuperscript{-1} mol\textsuperscript{-1}). Under reaction pressure (2 bar(g)), estimation of reaction order revealed first order kinetics law with respect to H\textsubscript{2} for all catalysts and first or zero order rate law with respect to propene for catalysts Cr\textsuperscript{III}@SiO\textsubscript{2}–Al\textsubscript{2}O\textsubscript{3} and CrO\textsubscript{x}/Al\textsubscript{2}O\textsubscript{3} and for catalysts Cr\textsuperscript{III}/Al\textsubscript{2}O\textsubscript{3} and Cr\textsuperscript{III}@Al\textsubscript{2}O\textsubscript{3}, respectively. At lower pressures, the rate law always depends on the concentration of propene, following a Langmuir-like behaviour. This observation is likely due to a strong adsorption of propene on the sites (site-saturation). Furthermore, ParaHydrogen-Induced Polarization (PHIP),\textsuperscript{59–62} was used to probe the mechanistic pathway for hydrogenation – and thereby the microscopic reverse reaction (dehydrogenation) – on Cr\textsuperscript{III} supported single sites. PHIP shows that hydrogenation

Scheme 3

(a) Isolated Cr\textsuperscript{III} sites prepared using an SOMC/TMP approach (a) synthesis of Cr\textsuperscript{III} single sites on SiO\textsubscript{2} partially dehydroxylated at 700 °C; and (b) SiO\textsubscript{2}–Al\textsubscript{2}O\textsubscript{3} and Al\textsubscript{2}O\textsubscript{3}.

Scheme 4

(a) Proposed non-redox mechanism for the dehydrogenation of light alkanes over M\textsuperscript{III} isolated sites (M = Cr & E = Si or Al; or M = Ga & E = Si); and (b) structure of model tri- and tetra-coordinated Cr\textsuperscript{III} sites on Al\textsubscript{2}O\textsubscript{3}. Reproduced from ref. 43 with permission from American Chemical Society, copyright 2016.
likely takes place via pairwise heterolytic $H_2$ activation, as evidenced by the partial induced polarization transfer, in agreement with earlier proposed mechanism. Modelling the Cr$^{III}$ sites dispersed on alumina by periodic DFT calculations further support the proposed mechanism, which involves the non-redox heterolytic cleavage of a terminal C–H bond on a Cr–O bond (Scheme 4a) leading to the formation of Cr-alkyl and OH group followed by hydrogen transfer, olefin de-coordination and desorption of $H_2$ to regenerate the initial Cr–O reactive sites. This mechanism is overall more favoured on tetra- than tri-coordinated Cr$^{III}$ sites (Scheme 4b), and is favoured on alumina compared to silica, likely because of the greater polarization of the Cr–O bonds in the more ionic alumina support.

### 3.2 Supported Ga$^{III}$ and related systems

In parallel to Cr$^{III}$-based systems employed industrially, main group oxides dispersed on oxide supports have also been shown to selectively dehydrogenate alkanes. In fact, the Cyclar$^{63}$ process (BP/UOP) that is used industrially to produce BTX aromatics is based on a Ga-exchanged (ZSM-5) zeolite, where isolated Ga$^{III}$ sites and Brønsted acid sites are thought to dehydrogenate propane and aromatize the olefins, respectively. A wide variety of Ga- as well as In-exchanged zeolites show similar reactivity. Intriguingly, it has been observed that when the Ga is introduced into the zeolite using an organometallic precursor (e.g. GaMe$_3$), the resulting material is selective for the dehydrogenation of propane to propene, likely because GaMe$_3$ reacts with the Brønsted acid sites – quenching the Brønsted acidity, and preventing subsequent reaction steps. Furthermore, numerous Ga-containing systems, that are not supported on hierarchical materials, have been shown to be proficient in light alkane dehydrogenation. For example, numerous Ga$_2$O$_3$-based systems have been reported to selectively catalyse propane dehydrogenation, though these systems typically suffer from rapid deactivation due to Ga$_2$O$_3$ reduction.

To suppress reduction of Ga$^{III}$ in reaction conditions, isolated Ga$^{III}$ sites dispersed on silica, a non-reducible support, have been developed and are highly active, selective and stable PDH catalysts. These Ga$^{III}$ isolated sites are prepared by the SOMC/TMP approach, similarly the Cr$^{III}$ sites discussed above (Scheme 5a). Here, [Ga(OSi(OBu)$_3$)](THF) is grafted on silica (Aerosil$^{64}$) partially dehydroxylated at 700 °C, followed by a thermal treatment under reduced pressure (10$^{-3}$ Pa, 500 °C) that removes the organic ligands leaving Ga$^{III}$ isolated sites.

Using a combination of Ga K-edge EXAFS, wavelet transform analysis (WT) of the EXAFS and analysis of Ga K-edge XANES, the thermally treated material (Ga@SiO$_2$) was shown to be composed of isolated tetra-coordinate Ga sites dispersed on...
the surface of silica. The material shows a high activity per Ga, which is attributed to effective metal utilization associated with SOMC. In addition, a high selectivity (93%) is observed (Table 2), which is likely related to the absence of Bronsted acid sites that are often invoked in cracking and dehydroaromatization reactions. Recent computational studies demonstrate that GaIII-catalyzed propane dehydrogenation most likely proceeds through an analogous mechanism to that described for CrIII (Scheme 4a), and highlight that the presence of strain and specific configuration are critical for highly active isolated GaIII sites. While isolated GaIII sites are thought to be formed via impregnation method using (Ga(NO₃)₃·nH₂O/citric acid/Davisil 646), their activity and selectivity in PDH are significantly lower (Table 2) than the materials prepared via SOMC, indicating that the nature of the material must be different. This may be attributed to the complexity of the material prepared by impregnation in water, where dissolution/precipitation can readily take place, thus leading to more complex materials.

Using an approach akin to the aforementioned work with GaMe₃ on zeolites, the SOMC approach has also been used with silica and alumina support materials, using the homoleptic precursor tri-isobutylgallane – Ga(iBu)₃ (Scheme 5b). A dimeric surface species is proposed to be formed upon grafting of Ga(iBu)₃ on silica partially dehydroxylated at 700 °C (Ga(iBu)₃/SiO2-700m Degussa Aerosil 200), as evidenced by W-EXAFS and NMR. Conversely, monomeric Ga sites are formed on Al₂O₃ partially dehydroxylated at 500 °C (Ga(iBu)₃/Al₂O₃-500γ-alumina) as evidenced by EXAFS and NMR. The alumina supported catalyst is significantly more active in PDH (550 °C, 20% propane in Ar) than the silica-supported material (Table 2); the difference of reactivity is attributed to the greater reactivity towards C–H bonds of Ga–O in bonds in [−Ga–O−Al−] vs. [−Ga–O–Si−] systems, which parallels what is found for Cr (vide supra). Note however that Al₂O₃ also displays a considerable PDH activity and selectivity (ca. 4% conversion, ≈40% total activity after 20 hours on-stream). In this case, the high background activity of alumina precludes more direct comparison of site requirements (such as nuclearity).

3.3 1st row transition metal single-site PDH catalysts

While supported vanadium oxides are key catalysts for the oxidative dehydrogenation of propane, they have also been studied in non-oxidative PDH discussed in this review. These catalysts are thought to require partial reduction of V₂O₅ to VO₂ sites. Pre-treatment by H₂ has been shown to influence the activity, and both VIII and V⁴V isolated sites have been proposed to catalyse this reaction.

To gain insight into the active structures for the vanadium catalysed dehydrogenation of propane, silica-supported V⁸V and V⁴V single sites have thus been prepared via SOMC (Scheme 6) using the well-defined V⁸V and V⁴V molecular precursors V⁸(VMes)(THF) and V⁴(V)(OMes)₃. Characterization by FT-IR, UV-vis, solid-state ¹H and ⁵¹V NMR and XAS shows that these precursors can generate the corresponding mono [(≡SiO)V⁴(V)(OMes)₂] or bis-grafted [(≡SiO)V⁸(V)(OMes)] sites depending on the temperature at which the silica support is dehydroxylated (Scheme 6). Activity for these catalysts is reported in Table 3. Following an induction period, [(≡SiO)V⁴(V)(OMes)₂] shows high PDH activity (Table 3) compared to V₂O₅/SiO₂ prepared by impregnation, likely due to the absence of inactive polynuclear VO₄ species. [(≡SiO)V⁸(V)(OMes)₂(THF)] shows comparable TOF to [(≡SiO)V⁴(V)(OMes)₂], albeit with a significantly reduced induction period, suggesting that V⁸V species are the active species (Table 3). Pre-treatment of [≡(≡Si)VO⁻(OMes)] under H₂ allows the total suppression of this induction period. This has been attributed to the formation of surface hydride species. Furthermore, a more recent study indicates that a V⁸V species is likely formed upon hydrogen treatment of [(≡SiO)V⁵(V)(OMes)₂] at elevated temperatures (250–500 °C), suggesting that in spite of the different precursors used, the active site structures and ‘activated’ materials are very similar. While various plausible pathways have been proposed, initiation likely starts with the C–H activation of propane to form mesitylene and a propyl vanadium species, followed by β-H elimination and formation of a propylene π-complex. An alternative pathway invokes the cleavage of a V–O bond in a fashion akin to that described for Cr⁷ VII (vide supra, Scheme 4a). However, to obtain further insight into the mechanism additional in situ or operando studies would be beneficial. Studies using PHIP show that pairwise replacement of two hydrogens from propane with the ones originating from para-H₂ is not an efficient process over [(≡SiO)V⁵(V)(OMes)₂], further highlighting the ambiguity in interpretation of the propane dehydrogenation mechanism in reaction conditions. Detailed mechanistic studies of the microscopic reverse reaction (i.e. propene hydrogenation) show that by switching between hydrogenation and dehydrogenation conditions, the hydrogenation capacity of the catalyst

Table 2  Catalytic properties of Ga-based catalysts for propane dehydrogenation discussed in Section 3.2

<table>
<thead>
<tr>
<th>Material</th>
<th>Feed</th>
<th>Time (h)</th>
<th>T (°C)</th>
<th>Conv. (%)</th>
<th>Select. (%)</th>
<th>Productivity ²</th>
<th>wt% Ga</th>
<th>Ref.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ga@SiO₂</td>
<td>3.3% propane in Ar (2 bar)</td>
<td>0.1</td>
<td>550</td>
<td>9.3</td>
<td>94</td>
<td>13</td>
<td>1.6</td>
<td>76</td>
</tr>
<tr>
<td></td>
<td></td>
<td>20</td>
<td>6.5</td>
<td>93</td>
<td>9</td>
<td></td>
<td></td>
<td></td>
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<tr>
<td>Ga/SiO₂</td>
<td>2.3% propane in Ar (1.4 bar)</td>
<td>1</td>
<td>550</td>
<td>9.8</td>
<td>97</td>
<td>0.9</td>
<td>2.64</td>
<td>79</td>
</tr>
<tr>
<td></td>
<td></td>
<td>1</td>
<td>550</td>
<td>8</td>
<td>90</td>
<td>4.7</td>
<td>3.34</td>
<td>80</td>
</tr>
<tr>
<td>Ga[iBu₃]/SiO₂-700</td>
<td>20% propane in Ar (1 bar)</td>
<td>1</td>
<td>550</td>
<td>24</td>
<td>—</td>
<td>13</td>
<td>1.97</td>
<td>80</td>
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<tr>
<td></td>
<td></td>
<td>25</td>
<td>2.5</td>
<td>72</td>
<td>2.3³</td>
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<tr>
<td>Ga[iBu₃]/Al₂O₃-500</td>
<td>20% propane in Ar (1 bar)</td>
<td>1</td>
<td>550</td>
<td>8</td>
<td>90</td>
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<td>90</td>
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</table>

² 20 h on-stream. ³ Not disclosed.
is reduced, suggesting that the hydrogenation is enabled by a site that is not stable in relevant conditions – precluding mechanistic insights being obtained using this strategy.89

3.4 Late-transition metal PDH catalysts based on single-sites

Later 3d transition-metals dispersed on oxide supports such as Fe and Co have also shown PDH activity.6,94–96 For example, silica-supported FeII single sites have been prepared by grafting bis(2,4-dimethylpentadienyl)iron(II) on silica (Scheme 7a). As evidenced NMR, FTIR and analysis of washings, the grafting takes place by reaction of one Si–OH group with bis(2,4-dimethylpentadienyl)iron(II) leading to the release of 1 equiv. of 2,4-dimethylpentadiene (Scheme 7). Reduction of this material under H2 at 400 °C leads to the formation of small iron oxide clusters that can be re-dispersed by thermal treatment at 650 °C. The associated FeII isolated sites show PDH activity (Table 3), while the corresponding iron nanoparticles show poor selectivity (coke formation is the dominant reaction), and iron oxide nanoparticles supported on silica show significantly lower activity and selectivity for PDH, suggesting that FeII isolated sites are catalytically competent in PDH.97

Using the SOMC/TMP approach, mononuclear silica-supported CoII PDH catalysts have been prepared starting from the dinuclear CoII precursor [Co2(OSi(OtBu))3]2 (Scheme 7b).98 EXAFS data, consistent with the presence of a Co–O path of degeneracy 4 (1.94 Å), attributed to anionic oxygen directly bound to Co centre and adjacent siloxane oxygens interacting in a dative fashion, was used to establish the proposed surface structure. Worth of note, the activity toward PDH of such

Table 3 Catalytic properties of 1st row transition metal based catalysts described in Sections 3.3 and 3.4

<table>
<thead>
<tr>
<th>Material Feed</th>
<th>Time (h)</th>
<th>T (°C)</th>
<th>Conv. (%)</th>
<th>Select. (%)</th>
<th>Productivity (gprod g M⁻¹ h⁻¹ wt% M)</th>
<th>Ref.</th>
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<tbody>
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<td>[(SiO)2VIV(O)Mes2] 20% propane/Ar</td>
<td>0.1</td>
<td>500</td>
<td>17</td>
<td>83</td>
<td>—</td>
<td>V: 1.40 83</td>
</tr>
<tr>
<td>[(SiO)2VIV(O)Mes] 20% propane/Ar</td>
<td>0.1</td>
<td>500</td>
<td>17</td>
<td>90</td>
<td>—</td>
<td>V: 1.64 83</td>
</tr>
<tr>
<td>[(SiO)2VIVMes] 2.3% propene/Ar</td>
<td>0.3</td>
<td>550</td>
<td>8</td>
<td>94</td>
<td>0.19</td>
<td>V: 2.8 89</td>
</tr>
<tr>
<td>FeII@SiO2</td>
<td>3% propane/Ar</td>
<td>0</td>
<td>650</td>
<td>4.1</td>
<td>69a</td>
<td>0.83</td>
</tr>
<tr>
<td>FeII/SiO2200</td>
<td>3% propane/Ar</td>
<td>0</td>
<td>650</td>
<td>27</td>
<td>14</td>
<td>32.3</td>
</tr>
<tr>
<td>FeII/SiO2</td>
<td>3% propane/Ar</td>
<td>0</td>
<td>650</td>
<td>32</td>
<td>27</td>
<td>—</td>
</tr>
<tr>
<td>CoII@SiO2 (SOMC)</td>
<td>20% propane/Ar</td>
<td>0.1</td>
<td>550</td>
<td>9.5</td>
<td>73</td>
<td>—</td>
</tr>
<tr>
<td>CoII/SiO2 (IWI)</td>
<td>3% propane/Ar</td>
<td>0</td>
<td>550</td>
<td>6.5</td>
<td>92</td>
<td>8.9</td>
</tr>
<tr>
<td>CoII/SiO2 (IWI)</td>
<td>3% propane/Ar</td>
<td>0</td>
<td>550</td>
<td>4</td>
<td>89</td>
<td>3.7</td>
</tr>
</tbody>
</table>

a The reported corrected selectivity is 99% upon subtraction of cracking products – the observed selectivity is below 70%. Calculated based on metal loading in the material prior to reduction in H2. b Corrected selectivity by subtracting the amount of thermal cracking attributed to silica.

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Scheme 6 Preparation of silica supported VV and VV aryl complexes for PDH, (a) SOMC approach for the preparation of monopodal and bipodal supported VV single sites employed in vanadium catalysed propane dehydrogenation; and (b) preparation of VV single sites for propane dehydrogenation prepared using an SOMC approach.
SOMC-TMP prepared CoII single sites is significantly higher than the one of single sites prepared via impregnation approach (Table 3). However, the stability of this catalyst is poor, due to the formation of nanoparticles under reaction conditions. A mechanism similar to the one proposed for CrIII (vide supra, Scheme 4a) has been calculated on model clusters and is consistent with kinetic and PHIP experiments on SOMC prepared single sites in the reverse hydrogenation reaction. Both experiment and calculations indicate that PDH likely involves the heterolytic C–H activation of propane, similarly to what has been proposed for CrIII.

4 Modelling alkane dehydrogenation catalysts using supported metal nanoparticles and alloys prepared via SOMC

Early studies have shown that PtM (M = Pd, Rh, Ir, Sn) nanoparticles can catalyze cyclohexane dehydrogenation as well as other important hydrocarbon conversion reactions. The addition of a second metal component has been shown to result in enhanced selectivity and stability. The superior catalytic performances of bimetallic catalysts in alkane dehydrogenation, when compared to their monometallic counterparts, has raised a considerable interest in bimetallic systems, and has led to the development of industrial systems for the dehydrogenation of light alkanes. First commercialized in the 1990s, the UOP Oleflex process consists of alloyed Pt–Sn nanoparticles supported on alumina, and is used for the selective dehydrogenation of propane. In addition to the licensed Pt–Sn systems, there are an array of related systems that have garnered attention in both academia and industry. This broad class of catalysts, of general structure PtM/MOx, have been widely studied as model systems for the Oleflex catalyst.

In spite of the complexity of these systems and the presence of multiple metal species, SOMC has also been used to study supported metal nanoparticles as models for the Oleflex system. Building upon pioneering work on the Pt–Sn system by Yermakov in the late 1970’s, SOMC research in the 1980’s and 1990’s focused on modifying supported metal nanoparticles and generating alloys for alkane dehydrogenation through the selective grafting of organometallic compounds on the surface hydrides present at the surface of platinum particles. In the specific case of Pt–Sn, this approach could be successfully carried out by reacting organotin species selectively with supported Pt nanoparticles at ambient temperature, followed by a high temperature treatment under H2 (Scheme 2d). The first step leads to the formation of R3Sn–Pt moieties at the nanoparticle surface, which, upon treatment under H2 at elevated temperatures, lose their alkyl ligands and form a Pt–Sn alloy by migration of the Sn inside of the particle. This approach was used more recently, to prepare a trimetallic Pt–Sn–Li material active in the dehydrogenation of 2,3-dimethylbutane (Scheme 8a). The treatment of alumina with BuLi and Pt(acac)2, followed by a calcination under air at 400 °C and a reduction at 550 °C under H2 yielded a material containing dispersed Li+ aluminate sites (quenching of Brønsted acidity for the purpose of suppressing side reactions) and small Pt particles. Subsequent reaction of this material with Bu4Sn, in the presence of H2 (550 mbar), followed by a treatment under H2 at 550 °C did not lead to a change in particle size. The addition of both Li and Sn by SOMC leads to an increased productivity and stability compared to the monometallic Pt catalyst (Table 4). Addition of Sn leads to an increased selectivity, and is manifested in H2 chemisorption measurements. The beneficial role of Li has been attributed to decreasing the acidity of the Al2O3 support and...
increasing Pt dispersion, leading to an overall improved catalytic performance.

Alternatively, the consecutive grafting of organometallic precursors via the OH groups of metal oxide supports was implemented for the synthesis of bimetallic materials (Scheme 8b). For instance, PtSn supported on \( \theta \)-Al\(_2\)O\(_3\) have been recently prepared via SOMC through the consecutive grafting of \([\text{Pt(COD)Me}_2]\) and \(\text{HSnPh}_3\) onto the partially dehydroxylated support via Pt–Me and Sn–H cleavage as evidenced by GC analysis (with the respective formation of \(\text{CH}_4\) and \(\text{H}_2\)).

Powder XRD measurements only show contributions of the \(\theta\)-Al\(_2\)O\(_3\) structure which indicates no significant amounts of crystalline PtSn, Pt or Sn domains in the material prior to a reduction step under \(\text{H}_2\) at 150 °C that yields supported sub-nanometric (0.8 nm), highly dispersed and size-homogeneous Pt–Sn particles as evidenced by HAADF-STEM. CO adsorption IR indicates formation of a Pt–Sn alloy as the bridge-type CO vibrational band (\(\nu_{\text{CO}} < 2000 \text{ cm}^{-1}\)) is absent in these spectra. XPS analysis indicates the coexistence of Sn\(^{IV}\), Sn\(^{II}\) and Sn\(^0\) species with increasing volumes of oxidized species with increased Sn/Pt ratio. While Sn\(^0\) enhances the selectivity due to alloy formation, the presence of Sn\(^{IV}\)/Sn\(^{IV}\) is proposed to prevent sintering, hence the formation of small particles. In this study, a PtSn material containing a ratio of 3 : 1 Sn : Pt shows highest PDH performance with 40% conversion over 72 h at >99.5% selectivity for propene – stable in a WHSV range from 11.8 h\(^{-1}\) to 3300 h\(^{-1}\) at 550 °C (Table 4).

Besides Sn, Ga has garnered attention in alkane dehydrogenation processes, and has recently been implemented industrially, with the Fluidized Catalytic Dehydrogenation process introduced by Dow Chemical in 2017.

To gain insight into this family of catalysts, which contain both Pt and Ga, a PtGa bimetallic material was prepared via SOMC by grafting of \([\text{Pt(COD)(OSi(OtBu)}_3)\text{]}_2\) onto the

### Table 4  Catalytic properties of the materials described in Section 3.2

<table>
<thead>
<tr>
<th>Material (particle size, nm)</th>
<th>Feed</th>
<th>Time (h)</th>
<th>T (°C)</th>
<th>Conv. (%)</th>
<th>Select.(^a) (%)</th>
<th>Productivity(^a) (\text{g prod/g Pt h})</th>
<th>wt% M</th>
<th>Ref.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pt/SiO(_2) (2.2 ± 0.8)</td>
<td>20% propane/argon</td>
<td>0.1</td>
<td>550</td>
<td>2.5</td>
<td>75</td>
<td>14.3</td>
<td>3.96 Pt</td>
<td>112</td>
</tr>
<tr>
<td>Pt–Sn–Li/Al(_2)O(_3) [n.d.]</td>
<td>35% 2,3-dimethylbutane/H(_2)</td>
<td>0.1</td>
<td>500</td>
<td>40</td>
<td>77</td>
<td>55</td>
<td>2.0 Pt, 1.32 Sn, 2.8 Li</td>
<td>119</td>
</tr>
<tr>
<td>Pt–Sn/Al(_2)O(_3) (0.8 ± 0.2)</td>
<td>10% propane/10% H(_2)/Helium</td>
<td>0.1</td>
<td>550</td>
<td>43</td>
<td>99</td>
<td>860</td>
<td>0.5 Pt, 1.5 Sn</td>
<td>120</td>
</tr>
<tr>
<td>Pt–Ga/SiO(_2) (1.0 ± 0.2)</td>
<td>20% propane/argon</td>
<td>0.3</td>
<td>550</td>
<td>32</td>
<td>&gt;99.8</td>
<td>766</td>
<td>4.82 Pt, 1.48 Ga</td>
<td>111</td>
</tr>
<tr>
<td>Pt–Zn/SiO(_2) (0.8 ± 0.2)</td>
<td>20% propane/argon</td>
<td>0.3</td>
<td>550</td>
<td>30</td>
<td>98</td>
<td>793</td>
<td>3.05 Pt, 1.54 Zn</td>
<td>112</td>
</tr>
<tr>
<td>Pt–Zn/SiO(_2) (ALD) (1.2 ± 0.3)</td>
<td>5% (n)-butane/argon</td>
<td>0.1</td>
<td>500</td>
<td>79</td>
<td>92.5</td>
<td>165</td>
<td>0.6 Pt, 9.9 Zn</td>
<td>121</td>
</tr>
<tr>
<td>Pt–B/SiO(_2) (2.6 ± 0.6)</td>
<td>1% (n)-butane/argon</td>
<td>0.1</td>
<td>550</td>
<td>1.6</td>
<td>70</td>
<td>8.7</td>
<td>0.021 Pt, 0.60 B</td>
<td>122</td>
</tr>
</tbody>
</table>

\(^a\) Only desired products are included. \(C_6\) olefins in the case of 2,3-dimethylbutane, propene in the case of propane, butenes in the case of \(n\)-butane.
silica-supported Ga\textsuperscript{III} single-sites discussed above (Ga@SiO\textsubscript{2}, Section 3.2) followed by a thermal treatment under H\textsubscript{2} (Scheme 9a).\textsuperscript{111} This approach builds on the presence of isolated Ga\textsuperscript{III} sites in close proximity of –OH groups (ca. 0.6 Ga and 1.3 –OH nm\textsuperscript{-2}) that enables introduction of a controlled quantity of Pt through grafting onto the –OH groups. After H\textsubscript{2} treatment, highly dispersed and size-homogeneous 1 nm particles are generated, even at a high Pt loading of 4.37 wt%.\textsuperscript{111} Detailed CO adsorption IR, XPS and XAS studies have shown that a Pt–Ga alloy is formed along a 0.1 to 0.5 fraction of the overall Ga remaining as Ga\textsuperscript{III} on the support surface. EXAFS fitting indicates that bulk Ga\textsubscript{2}O\textsubscript{3} domains are not present. The material shows high initial productivity (Table 4) and selectivity (>99%) in the PDH reaction at 550 °C. The productivity of this catalyst, per g, exceeds that of the Ga\textsuperscript{III} single-site PDH catalysts, described in Section 3.2 by ca. two orders of magnitude. In addition to the high selectivity that is observed, this material also shows an unusually high long-term stability over 20 h during which the average structure of the catalyst does not change, according to \textit{in situ} XAS. Finally, this material has been shown to be partially regenerated upon consecutive O\textsubscript{2}/H\textsubscript{2} treatments, with a recovery of 75% of the initial activity with a high propene selectivity (>99%) retained (Table 4).

In an analogous approach to the synthesis of bimetallic Pt–Ga materials prepared \textit{via} SOMC described above, the corresponding silica-supported Pt–Zn bimetallic system was prepared, starting from isolated Zn\textsuperscript{II} sites supported on silica.\textsuperscript{112} This approach yielded silica-supported subnanometric Pt–Zn clusters (0.8 nm). CO adsorption IR, XPS and XAS studies confirm the formation of a Pt–Zn alloy with an approximate 1 : 1 ratio of Zn : Pt in the alloyed particles while an overall 3 : 2 ratio of Zn: Pt is evidenced by EA. This indicates that 33% of Zn\textsuperscript{II} remains on the support surface, in line with observations from XPS and XANES linear combination fits (LCF) indicating the presence of 20–45% of Zn\textsuperscript{II} (Scheme 9b). The material shows excellent performance in the PDH reaction with a higher initial productivity (Table 4) than most other Pt–Zn based systems. Propene selectivity (>98%) drops slightly after 30 h on stream despite the relatively high stability over 30 h when compared to other Pt-based propane dehydrogenation catalysts (Table 4).

The use of one-cycle atomic layer deposition (ALD) has also recently been applied to prepare Pt–Zn nanoclusters on a SiO\textsubscript{2} support, that are shown to be active in the dehydrogenation of n-butane to butenes and 1,3-butadiene.\textsuperscript{122} Consecutive ALD of Zn(C\textsubscript{3}H\textsubscript{5})\textsubscript{2} followed by [(MeCp)PtMe\textsubscript{3}] yields a material containing 10 wt% Zn and 0.5 wt% Pt. After reduction in an H\textsubscript{2} stream at 550 °C for 2 h, the material consists of 1.2 nm alloyed PtZn particles as shown by HAADF-STEM, XAS and XPS and CO adsorption studies. The material exhibits high selectivity to butenes (>90%, of which <20% 1,3-butadiene) at high conversion levels at 550 °C. (Table 4) A long-term stability test over 7 days shows a 60% deactivation of the catalyst while the 1,3-butadiene selectivity increases from 12% to 28%.

Besides the combination of late and post-transition metals a recent study by Teplyakov et al. utilizes the SOMC approach to combine Pt and B supported on SiO\textsubscript{2} for butane dehydrogenation.\textsuperscript{122} The material was synthesized \textit{via} sequential grafting of B(OiPr)\textsubscript{3} and [(MeCp)PtMe\textsubscript{3}] onto partially dehydroxylated SiO\textsubscript{2} with an intermediate calcination step at 400 °C. A treatment under H\textsubscript{2} yielded 2.6 nm monometallic Pt particles supported on B doped SiO\textsubscript{2} as evidenced by HAADF-STEM and HR-TEM. XPS revealed similar trigonal and tetrahedral B sites in Pt–B and B materials as well as mainly Pt\textsuperscript{II}/Pt\textsuperscript{IV} and Pt\textsuperscript{0} in the material before and after H\textsubscript{2} treatment, respectively.\textsuperscript{11}B MQMAS NMR additionally showed significant differences of the B sites in Pt–B and B materials post-catalysis. In the B–Pt material a new feature consistent with a tetrahedral coordination environment appears in the post-catalysis sample which is consistent with calculations that invoke the formation of B–C bonds. Dehydrogenation performance of the B–Pt material shows a two-fold.
increase in activity compared to Pt, a selectivity of around 70% for butenes and minor deactivation over 25 h (Table 4). Redispersion of Pt species can be observed during catalysis as evidenced by post-catalysis HAADF-STEM, being more significant for the B-Pt sample compared to a Pt only material. In addition, Raman and TGA studies show that less coke is formed on the bimetallic material over the course of the reaction.

We have shown that bimetallic systems can be prepared via SOMC and that they show high activity towards propane dehydrogenation. Because it is possible to introduce dopants selectively at the support interface, these systems can be characterized exhaustively to provide molecular-level information about the structure of multi-metallic systems. This opens the possibility for more detailed *operando* and computational studies to address the question of the role of dopants, their effect on catalyst activity and stability as well as the structure and dynamics of active sites. Such insights could then inform the design of better performing catalysts.

5. Conclusions

SOMC, and the combination of SOMC & TMP, have been shown to provide molecular level insight into the structure and nature of active centres for the dehydrogenation of light alkanes for the two classes of dehydrogenation catalysts based on isolated sites or supported Pt nanoparticles. Through the systematic use of tailored molecular precursors, and rigorous characterization of materials at every step of the synthesis, it has been possible to obtain valuable insight into dehydrogenation processes using model catalysts. Furthermore, the ability to systematically change a single component of a system (i.e. identity of dopant), while fixing all other variables other metals, surface area, metal loading etc.) enables direct comparison of otherwise identical materials – something which is extremely challenging using traditional synthetic approaches such as incipient wetness impregnation. This has enabled rigorous study of general classes of catalysts employed in light alkanes dehydrogenation.

Specifically, in the case of isolated metal sites (M**n**+) on non-reducible supports, it has been possible to develop Cr**III**/MO**x** type catalysts, containing isolated metal ions which catalyse the dehydrogenation of light alkanes. This insight indicates that isolated metal ions are proficient in the dehydrogenation of light alkanes, and that there is most likely a non-redox mechanism that enables this reactivity. Broader study of a range of metal ions (Ga**III**, V**IV**+, Fe**II** and Co**II**+) shows that this is a more general approach for the synthesis of oxide-based catalysts for the dehydrogenation of light alkanes. In the case of Ga, it has also been shown that through the dilution of Ga**III** sites, and the formation of strong Ga-O-E (E = Si, Al) linkages, reduction (and thus deactivation) of Ga**III** can be suppressed.

For multi-metallic systems based on Pt-nanoparticles, the SOMC approach has been able to show, by using detailed spectroscopic study, that the presence of an alloyed PtM phase is critical to the stability of Pt-based systems. More generally, it has been possible to demonstrate that alloying, as a general strategy possible with an array of metals (Sn, Ga, Zn), enables the synthesis of more active, selective and stable catalysts for PDH. It has been shown, by using this molecular approach, that alloying serves as an effective strategy for the dilution of Pt centres at the surface of the supported nanoparticles – suppressing cracking and coke formation as well as augmenting selectivity. Interestingly, the materials synthesized using this approach possess a stability which exceeds that of existing catalysts. This can most likely be attributed to the existence of remaining dispersed Lewis acidic sites on the surface of these materials which serve as anchoring sites for the alloyed nanoparticles, hence reducing sintering. The molecular approach employed, and the ability to selectively introduce metal functions in the absence of complex anionic species typical of traditional synthetic approaches enables explicit study of the role of the metal. In future these well-defined model systems could be used to explore the possibility of further optimized ternary systems, similar to those described in the recent literature.

In summary, SOMC provides unprecedented insight into the molecular-level structure of heterogeneous catalysts for light alkane dehydrogenation, informing strategies for the synthesis of improved catalytic systems, and providing a basis for understanding of the subtle interplay between structure and reactivity. As spectroscopic, computational and data-driven high-throughput experimentation approaches improve, we envisage that the insights provided using this approach will serve as a platform for the systematic study of new materials for the dehydrogenation of light alkanes.

Conflicts of interest

There are no conflicts to declare.

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Notes and references

condensate to liquid hydrocarbons through catalytic dehydrogenation and oligomerization, *Processes*, 2016, 6, 139.


29. F. Heroguel, Controlled growth and interfaces of supported metal nanoparticles from late transition metal siloxides, PhD thesis, ETH Zurich, 2014.


32. G. Noh, E. Lam, J. L. Alfke, K. Larmier, K. Searles, P. Wolf and C. Copéret, Selective Hydrogenation of CO2 to CH3OH
Review Article

Chem Soc Rev

on Supported Cu Nanoparticles Promoted by Isolated TiIV Surface Sites on SiO2, ChemSusChem, 2019, 12, 968–972.
87 U. Rodemerk, M. Stoyanova, E. V. Kondratenko and D. Linke, Influence of the kind of VO\(_x\) structures in VO\(_x\)/


100 J. L. Gland, K. Baron and G. A. Somorjai, Low-energy electron diffraction, work function change and mass spectrometric studies of chemisorption and dehydrogenation of cyclohexane, cyclohexene and 1,3-cyclohexadiene on the Pt(111) surface, *J. Catal.*, 1975, 36, 305–312.


120 Z. Xu, R. Xu, Y. Yue, P. Yuan, X. Bao, E. Abou-Hamad, J. M. Basset and H. Zhu, Bimetallic Pt-Sn nanocluster from the hydrogenolysis of a well-defined surface compound consisting of \(\text{[\text{AlO}\_\text{3}](\text{COD})\text{Me}\text{]}\) and \(\text{[\text{AlO}\_\text{3}](\text{SnPh}\_\text{3})\text{]}\) fragments for propane dehydrogenation, *J. Catal.*, 2019, 374, 391–400.

