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Supraparticles on beads for supported catalytically active liquid metal solutions - the SCALMS suprabead concept†

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A novel GaPt-based supported catalytically active liquid metal solution (SCALMS) material is developed by exploiting the suprabead concept: Supraparticles, i.e. micrometer-sized particles composed of nanoparticles assembled by spray-drying, are bonded to millimeter-sized beads. The suprabeads combine macroscale size with catalytic properties of nanoscale GaPt particles entrapped in their silica framework.

At least one catalytic step is required in the fabrication of more than 85% of all chemical products in our daily lives. Therefore, it is evident that catalysis plays a major role in successfully

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- † Electronic supplementary information (ESI) available: Experimental section, spray-drying scheme, dynamic light scattering measurements, transmission electron microscopy, nano-CT projection, laser diffraction measurements, ICP results, SEM studies, nitrogen sorption measurements, reactor flow scheme, propane dehydrogenation experiments, temperature programmed oxidation measurements, ESI Video 1: HRES PC nano-CT tilt series (supraparticles as prepared), ESI Video 2: virtual reconstructed slices of HRES PC nano-CT tilt series, ESI Video 3: 3D visualization of HRES PC nano-CT tilt series. See DOI: https://doi.org/10. 1039/d3mh01020a
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New concepts

In this work, it is proposed for the first time to lift the size of micrometerscaled particles by supporting them on a millimeter-sized bead to create advanced catalyst-support entities, called suprabeads. This approach increases the size of the units while maintaining the unique properties of the micron-sized particles. While this idea is familiar for supported nanoparticles, attaching µm-particles to a support material requires completely dissimilar, novel synthetic procedures because of the different size regimes and thus, dissimilar physicochemical forces involved in binding the individual species. This concept extends the applicability of µm-scale particles in catalysis and many other fields. Here, it was implemented to synthesize a catalyst-support unit in which catalytically active µm-sized supraparticles are supported on mm-beads. The supraparticles, which represent a relatively recent and unique class of multifunctional materials, consist of assembled SiO2 and Ga (or GaPt) nanoparticles. Additionally, the possibility to spray-dry gallium dispersions to create supraparticles with enclosed gallium particles or droplets, depending on the temperature of the environment, was shown for the first time. This offers a new approach for generating catalytically active materials.

implementing the sustainable use of resources and preserving of our environment. Catalysis helps reduce energy demands in the production of fuels and chemicals. It supports the conversion of renewable energy resources, is crucial for energy storage, and enables the purification of air as well as water.^{2,3} Most catalytic processes are heterogeneous, meaning that the reaction in the gas or liquid phase occurs at the surface of a catalyst possessing a different aggregate state.1 Consequently, there has been and continues to be a continuously ever-growing demand for new catalyst materials for a sustainable future. For this, creative concepts that go beyond traditional strategies are highly needed. 4,5 Heterogeneous catalysts often consist of a catalytically active material such as metallic nanoparticles supported by a high specific surface area material like porous alumina or silica.^{6,7} While the support material has no catalytic properties, it can ensure retrievability of the catalyst material from a reaction medium or prevent coalescence (and deactivation) of catalytically

active nanoparticles. Thus, in the design of high-performance catalysts, both the active material and the support must interact optimally.

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In the last 15 years, the concept of considering nanoparticles as building blocks to create more complex particulate units has gained increasing attention.8 Such entities can be termed supraparticles9 and may represent a smart method for the creation of heterogeneous catalysts. Compared to nanoparticles and macroscale materials, supraparticles exhibit unique features: first, they preserve nanoparticulate properties, such as nano-derived catalytic activity, while elevating particle sizes to the (lower) microscale. 10,11 Second, they often provide additional unexpected functionalities beyond the sum of properties of their constituent building blocks, such as an emergent internal porosity of the material¹²⁻¹⁴ or a well-controllable 3-dimensional spatial distribution of catalytic spots within a supportive surrounding. 15-17 This ultimately permits the development of catalytic materials that exceed the limits of current ones in terms of activity, selectivity, and stability.3,18 Therefore, their high potential for catalytic applications is beyond all doubt. 15,19-21

Despite the great potential of supraparticles, their applicability for catalysis in fixed-bed reactors is still limited because of their still too-small microscale sizes. In these reactors, particle (or pellet) sizes of at least 0.5 to 2 mm are required to avoid massive pressure drops as well as particle losses.²² While there have been a some attempts to create millimetersized supraparticles using nanoparticles as building blocks, 23-29 these suffer from several drawbacks such as poor upscalability, involvement of complex surfaces as synthesis platforms, or challenging template fabrication and removal steps, as well as limitations in the materials that can be used.

Therefore, we propose the concept of attaching supraparticles to beads using a binder material to create suprabeads that maintain the supraparticle features while providing high micron to millimeter sizes. The idea is being implemented in the development of supraparticle-based supported catalytically active liquid metal solutions (SCALMS).30 This recently proposed new class of heterogeneous catalysts consists of small amounts of catalytically active species (such as Pd, Pt, Rh, or Ni) dissolved in an inactive low-melting-point metal (such as Ga) supported by a porous inert material (such as SiO2, Al2O3 or SiC). $^{6,31-34}$ At elevated temperatures (T > 250 °C), these alloys become fully liquid and thus the catalytically active metal is embedded in the highly dynamic matrix metal. The catalytic reaction occurs only at the liquid metal-gas interface (unlike conventional supported liquid phase catalysis). The liquid metal provides no reactant solubility and the active metal appears at the liquid surface as an active single atom during catalysis.34 This high surface mobility is probably one of the main reasons for the high selectivity and excellent coking resistance of SCALMS in high-temperature dehydrogenation reactions, as shown for butane, propane, and methylcyclohexane dehydrogenation. 30,33,35,36

Recent publications have highlighted the high potential of SCALMS while the deposition of gallium on porous supports remains an open challenge (if chemical Ga deposition via

impregnation and decomposition of air-sensitive and expensive (Et₃N)GaH₃ is to be avoided) due to the high surface tension of gallium droplets and thus their poor wetting behavior and scarce intrusion into the pores of commonly used substrates. 37 This limitation can be overcome by using the building block approach to create supraparticles. It allows us to pursue gallium particle encapsulation by a porous support matrix (Fig. 1(a) and (b)): single support nanoparticles act as a scaffold in which the gallium is trapped during supraparticle synthesis via spray-drying, which is a forced assembly of nanoparticles within a droplet (Fig. S1, ESI†). In the next step, the catalytically active platinum species is deposited by galvanic displacement into the gallium contained in the supraparticles. This results in catalytically active supraparticles consisting of GaPt and SiO2. For the applicability of these supraparticle-based SCALMS in propane dehydrogenation reactions, the subsequent generation of suprabeads using these supraparticles is essential. The suprabead envisaged as a SCALMS material (Fig. 1(c) and (d)) consists of an inert millimeter-sized (aluminum oxide) bead covered with supraparticles. Their attachment to the bead is ensured by an (alumina-based) binder material. The beads and binder are mixed and the so-coated beads are rolled in a bed of supraparticles.

Following the detailed procedure described in the Experimental Section (ESI†), the concept of suprabead fabrication is successfully implemented (initially without Pt) using supraparticles consisting of commercial SiO2 and Ga nanoparticles synthesized via an ultrasonication procedure^{37–39} (dynamic light scattering measurements and transmission electron microscopy of the nanoparticles are provided in ESI† Fig. S2). As shown in a scanning electron microscopy (SEM) study (Fig. 2(a)) as well as by nano X-ray computed tomography (nano-CT) (Fig. S3 and ESI† Videos 1-3), gallium particle encapsulation by a porous support matrix was successfully achieved during spray-drying.

The supraparticles obtained (hereafter referred to as Ga-SPs) possess an almost spherical shape and sizes of around 4 to 10 μm (confirmed by laser diffraction measurements (LDM) ESI† Fig. S4). It should be noted that the observed morphology of Ga-SPs is common for spray-dried assemblies: During droplet evaporation, the contained nanoparticles initially form a dense shell at the air/liquid interface. Further shrinkage of the droplet leads to a collapse of the spherical template structure and to the formation of the observed doughnut-like or spherelike morphology with one or more craters. 40,41 The gallium nanoparticles are statistically distributed within the supraparticle with a SiO_2 to Ga weight ratio of ~ 3.5 (calculated based on the Ga weight content of the sample as determined by inductively coupled plasma atomic emission spectroscopy (ICP-AES) ESI† Table S1). This statistical mixing of binary nanoparticle dispersion within a spray-dried supraparticle has already been extensively studied using models supported by experimental results. 42 The obtained Ga-SP-loaded suprabeads (hereafter referred to as Ga-SBeads) possess a spherical shape (Fig. 2(b)). They consist of a 0.73 mm alumina core, which is covered by an approximately 12 µm thick layer of supraparticles Nanoparticles Supraparticle Suprabead **SCALMS Building blocks** Catalysis Cross-section Cross-section Schematic $C_3H_6 + H_2$ um mm Pt deposition in Ga Binder Support SiO₂ particles particles via galvanic Ga-matrix Ga particles Pt-atoms displacement Al₂O₃ core

Fig. 1 Scheme showing the forced assembly of a mixed dispersion of gallium and silica nanoparticles (a) to supraparticles (b) via spray-drying. The supraparticles thus consist of gallium nanoparticles, which contain platinum (after its deposition via galvanic displacement) and are statistically distributed within a matrix consisting of silica nanoparticles. (Note that the perfect sphericity of the displayed schematic supraparticle shown does not exactly reflect its real morphology, but has been chosen for the sake of simplicity.) The supraparticles are subsequently attached to an aluminum oxide core using a binder, resulting in suprabeads (c). The suprabeads display a SCALMS material for catalytic propane dehydrogenation (d).

Suprabead fabrication

(without any specific ordered arrangement). The Ga-SBeads contain 11 weight-% Ga-SPs (see ESI† Table S2). The supraparticles are bonded to the core by an alumina binder, thus providing a uniform distribution of the supraparticle components (Ga and SiO₂) on its surface. Suprabeads providing complete coverage of the core with undamaged supraparticles and binder layer can only be obtained if the ratio of binder, core, and supraparticles is optimal, heat treatment during the fabrication is carefully conducted, and the supraparticles provide sufficient mechanical stability (examples for failed fabrication attempts are shown in ESI† Fig. S5).

Spray-drying

The platinum deposition, essential for the SCALMS preparation, is realized on Ga-SPs by a galvanic displacement protocol, 43 and does not affect the structural properties of the supraparticles. In this context, it has to be mentioned that the synthesis of first Ga droplets and a later galvanic displacement reaction have been chosen to keep the SCALMS fabrication as close as possible to the established protocol.³⁷ The size, morphology, and pore characteristics of the obtained sample (hereafter referred to as Ga₁₃₀Pt-SP), remain comparable to those of Ga-SP (as shown by LDM ESI† Fig. S2, SEM ESI† Fig. S6, and nitrogen sorption measurements (NSM) ESI† Fig. S7). The same holds for the prepared Ga₁₃₀Pt-SP loaded suprabeads (Ga₁₃₀PtSBeads, SEM, and photograph ESI† Fig. S8). The envisaged approximate molar gallium-to-platinum ratio of 130 is achieved in Ga₁₃₀Pt-SBead (ESI† Table S1). This ensures that the Ga₁₃₀Pt alloy is liquid under the reaction conditions.44 Comparing the pore characteristics of the supraparticles with the suprabeads (using NSM ESI† Fig. S9, and Table S3), the specific surface area $S_{
m BET}$ and the total pore volume $V_{
m pore}$ are both significantly reduced after suprabead fabrication due to the heavy macroporous/non-porous core material (a detailed explanation of the pore system characteristics of SPs and SBeads can be found in the ESI†). The mesopore network of the SPs in SBeads can be slightly infiltrated by binder material (resulting in slightly lowered mean pore diameters) while the general accessibility of these pores is still ensured.

Reactor testing

Consequently, the material characterizations already indicate the presence and accessibility of GaPt within the millimetersized suprabeads. To demonstrate their usability as SCALMS catalysts, propane dehydrogenation reactions are carried out at 550 °C using a continuous gas-flow reactor. The detailed experimental procedure is described in the Experimental Section of the ESI† and a flow scheme of the reactor is shown in ESI† Fig. S10. For these catalytic experiments, three suprabead reproductions are delivered and two reference suprabead samples are used,

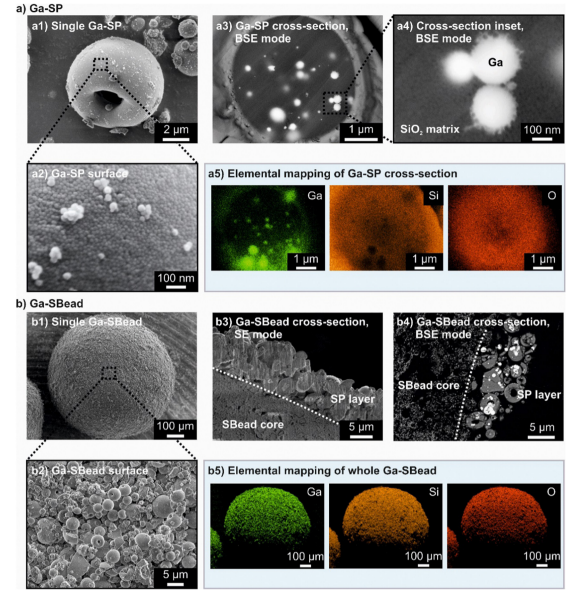


Fig. 2 SEM images of Ga-SP (a) and Ga-SBead (b), including micrographs showing their surface (a1, a2, b1, b2 – all secondary electron (SE) contrast), their cross-sections (a3 – backscattered electron (BSE) contrast, a4 - BSE, b3 – FIB cross-section SE, b4 – cross-section polish BSE) as well as elemental mappings of a Ga-SP cross-section (a5) and a whole Ga-SBead (b5).

namely Ga-SBeads and Pt-SBeads. The Pt-SBeads were prepared by wet impregnation of the SiO₂ supraparticles using a platinum precursor and subsequent reductive treatment to convert Pt(4+) into Pt(0) nanoparticles (which is an established route for the preparation of common heterogeneous Pt catalysts). This was followed by the fabrication of the suprabeads (referred to as Pt-SBeads). It should be noted that the gallium content of Ga-SBeads is comparable to that of Ga₁₃₀Pt-SBeads and the Pt content in the Pt-SBeads is even twice as much compared to the amount of Pt in the Ga₁₃₀Pt-SBeads (see ICP-AES, ESI† Table S1). The structural characterizations of the reference samples via NSM and SEM (shown in Fig. 2, as well as ESI† Fig. S7, Table S3, and Fig. S11) show similar morphologies and comparable pore characteristics.

The aim is to report for the first time the catalytic performance of GaPt SCALMS on suprabeads (Ga130Pt-SBead-1, denoting the first of three reproductions of the Ga₁₃₀Pt-SBead synthesis) in comparison with the reference materials containing only one metal species (Ga-SBead and Pt-SBead). To confirm the absence of blind activity, the beads, and binder material without the metals were also tested under the same reaction conditions (ESI† Fig. S12). As shown in Fig. 3, Ga₁₃₀Pt-SBead-1 provides superior catalytic performance compared to the Pt-SBead and Ga-SBead materials. Ga-SBeads provide an initial conversion of 1.1% followed by a steady decline in activity over the remaining 15 h time-on-stream (TOS) to a final conversion value of 0.5%. Due to the oxophilic nature of gallium, the presence of a thin passivating Ga₂O₃ layer has been

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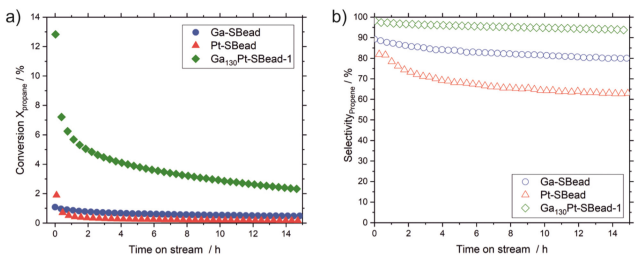


Fig. 3 Conversion (a; filled symbols) and selectivity (b; open symbols) for Ga-SBead (blue circles), Pt-SBead (red triangles), and Ga_{130} Pt-SBead-1 (green diamonds) in propane dehydrogenation. Catalyst mass in bed, 3 g. Pre-treatment conditions: 19.5 mL_N min⁻¹ H_2 , 80.5 mL_N min⁻¹ Ar, 1.2 bar, 550 °C. PDH experiment: 8.9 mL_N min⁻¹ G_3H_8 , 89.9 mL_N min⁻¹ Ar, 550 °C, 1.2 bar, 15 h TOS, GHSV: 1960 h⁻¹.

reported on gallium-based materials (regardless of the preactivation step), 45 together with the presence of Lewis acidic ${\rm Ga^{3^+}}$ sites. 46,47 Both species can activate hydrocarbon C-H bonds under these reaction conditions. 45,48,49

The Pt-SBeads display an initial conversion of 1.89%, which deactivates rapidly to 0.4% and 0.1% after 1 h and 15 h of timeon-stream (TOS) respectively. The initial activity of the Pt-SBead is ascribed to the presence of Pt nanoparticles in the heterogeneous Pt based catalyst prepared by wet impregnation and subsequent reductive treatment. However, under reaction conditions, these Pt nanoparticles are prone to sintering due to weak metal-support interactions.⁴⁹ The process of sintering leads to the loss of active surface and the formation of vicinal sites. The later leads to cracking activity and coke formation which is responsible for the rapid deactivation observed with Pt-SBeads. Interestingly, the synergistic effects of combining Pt with Ga in the GaPt alloy lead to a significant jump in activity. Ga₁₃₀Pt-SBeads show an initial conversion of 12.8%. This represents a 6-fold and 12-fold increase in initial catalytic activity when compared to the Pt-SBeads and Ga-SBeads respectively, under the same reaction conditions. In addition, Ga₁₃₀Pt-SBeads display very high selectivity with an initial selectivity value of 98%. In contrast, Ga-SBeads show an initial selectivity value of 89%, representing a 9% decrease in selectivity compared to the alloy-based catalyst. The activity on Ga-SBeads induced by the presence of Ga₂O₃ is often accompanied by a tendency to crack, 48 which favors the enhanced formation of side products by Ga-SBeads and reduces their selectivity. The Pt-SBead sample shows a far lower selectivity of 82%, which decreases to 70% after 15 h TOS due to the presence of vicinal sites which promote side product formation and subsequent deactivation (note, blind activity shows a selectivity of 45%. ESI† Fig. S12). To further corroborate these results, two reproductions of Ga₁₃₀Pt-SBeads were made and tested for catalytic activity (and provided in ESI† Fig. S13).

The data obtained from structural characterization, metal loading, as determined by ICP-AES, and catalytic performance,

as summarized in ESI† Table S4, strengthen our approach in the preparation of Ga₁₃₀Pt-SBeads and their excellent reproducibility. The nature of the active sites responsible for the activity in Ga₁₃₀Pt-SBeads is believed to be similar to that of already established GaPt SCALMS catalysts, since only its enclosure in the support was modified rather than the GaPt alloy. However, compared to established SCALMS, e.g., Ga45Pt SCALMS, the Ga₁₃₀Pt-SBeads presented here show superior activity (ESI† Fig. S14), which presumably results from the encasement of the GaPt droplets within the silica matrix of the supraparticles. This prevents migration and coalescence of GaPt droplets (both in the catalyst preparation process and the catalytic reaction) and thus, provides a higher catalytically active surface area. The still statistical distribution of Ga within the spent Ga₁₃₀Pt-SBeads (SEM ESI† Fig. S15) indicates the immobilization of the catalytically active phase. In GaPt SCALMS, the small amount of Pt present is alloyed with an excess of Ga. Under reaction conditions, the formed alloy becomes liquid. 50,51 In the presence of a hydrocarbon, the Pt metal is enriched at the liquid metal/gas interface. 30,34 The low solubility of the hydrocarbon ensures that subsequent substrate adsorption, reaction, and product desorption occur at the liquid metal/gas interface.30,34 The absence of vicinal sites due to the isolated and atomically dispersed Pt within the Ga matrix prevents traditional deactivation by sintering. However, other deactivation phenomena may be responsible for the loss of activity seen in Ga₁₃₀Pt-SBeads. Obvious culprits could be changes in catalyst morphology. To investigate this further, analyses of the spent Ga₁₃₀Pt-Sbeads' structure provide no evidence of morphological or porosity changes (SEM ESI† Fig. S15 and NSM ESI† Fig. S16) apart from minor supraparticle-losses. However, subatomic changes in the GaPt alloy under these conditions cannot be completely excluded. In addition, coke analysis was performed on the spent Ga-SBeads, Pt-SBeads, and Ga130Pt-SBeads catalysts by temperature program oxidation (TPO). As shown in ESI† Fig. S17 Ga₁₃₀Pt-SBeads catalysts showed no detectable carbon

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deposits, whereas Ga-SBeads and Pt-SBeads showed a mass loss of 0.04% and 0.1%, respectively. This suggests a different deactivation mechanism than coking. Raman et al. discussed other possible deactivation mechanisms at play, namely the phase separation of the alloy into Ga-rich and Pt-rich phases under reaction conditions.^{37,52} Detailed studies of this phenomenon are currently underway as an extension of this work.

Conclusions

Taking advantage of the suprabead concept, we have succeeded in synthesizing a heterogeneous catalyst unit in which µm-scale catalytically active supraparticles are bound to mm-sized Al₂O₃ beads using an Al₂O₃-based binder. The supraparticles consist of GaPt nanoparticles enclosed by a matrix of SiO2 nanoparticles, which was achieved by spray-drying a mixed dispersion of Ga and SiO2 nanoparticles, followed by Pt deposition. The functionality and high performance of these novel SCALMS were demonstrated in the propane dehydrogenation reaction. The suprabead concept opens up a new world for the creation of supported catalytically active materials and for fully exploiting the high potential of supraparticles in catalysis.

Data availability

Data for this paper, including dynamic light scattering, ICP-AES, laser diffraction, nitrogen sorption, nano CT, propane dehydrogenation runs, photographs, SEM and EDX, TEM, and TPO measurements and analysis are available at Zenodo at https://doi.org/10.5281/zenodo.8193496.

Author contributions

This collaborative research was initiated by K. M., P. W., M. H., and S. W. General research goals were developed by S. W., M. H., T. Z., N. M., and P. G. The supervision of the whole project was done by S. W. and M. H. Performance of the syntheses and characterization were carried out as follows: Ga-nanoparticle dispersion was prepared by N. M., N. T. and N. D., spray-drying was carried out by P. G. and N. D. Suprabead development was conducted by T. Z., T. R. and N. D. Binder synthesis was done by T. Z. Propane dehydrogenation reactions were carried out by N. M. and N. D. Laser-diffraction was measured by T. Z. N₂ sorption measurements and analysis were performed by P. G. Cross-section preparation and analysis of SPs was conducted by P. G. SEM analysis of SPs and SBeads was performed by P. G. and T. R. Cross-section preparation and SEM imaging of SBeads and nano-CT experiments of SPs were performed by S. C. - the corresponding results were interpreted and discussed by S. C., B. A. Z. and E. S. Interpretation and conclusion of the results were done by S. W., M. H., T. Z., N. M., and P. G. Figures were created by T. Z., N. M., and P. G. The original manuscript draft was written by S. W. and N. M. with the contribution of T. Z., and P. G. Reviewing and editing were

performed by T. R., K. M., P. W., and M. H. All authors proofread and commented on the manuscript.

Conflicts of interest

There are no conflicts to declare.

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