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Sonogenerated metal-hydrogen sponges for reactive hard templating

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As featured in:
Sonogenerated metal-hydrogen sponges for reactive hard templating†

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We present sonogenerated magnesium-hydrogen sponges for effective reactive hard templating. Formation of differently organized nanomaterials is possible by variation of sonochemical parameters and solution composition: Fe₂O₃ nanorods or composite dendritic Fe₂O₃/Fe₃O₄ nanostructures.

Synthesis of nanostructured materials by using a reactive hard template (RHT) approach was first introduced in 2007 by Thomas et al.1 with great advantage of omitting high temperatures or harsh reagents for removal of a template. Herein we combine both RHT and the sonochemical approaches to synthesise hierarchically structured materials and propose a novel method of template-assisted synthesis of nanostructures by using sonochemistry (sono-RHT, Fig. 1). The sono-RHT concept, which combines the advantages of hard templating with in situ decomposition, i.e., removal of a template, is a time saving and step-reducing process.

Synthesis and nanostructuring of solids by using sonochemical methods attract fundamental and technological interest due to the unique potential of locally generated high temperature (up to 5000 K) and high pressure (several hundreds of bars) processes provided by intensive ultrasonication.2 The propagation of the acoustic waves in liquid generates a cavitation field consisting of a large number of interacting microbubbles.2 These microbubbles can be considered as microreactors containing free radicals and excited species that can be used for chemical synthesis and nanostructuring of solids.2c Symmetric and asymmetric microbubble collapses can provide constant delivery of chemicals to the reaction zone.2d Recently, we demonstrated that inorganic materials could adopt a variety of different morphologies upon sonication of aqueous suspensions of solid particles. For instance, depending on sonication time, amorphous or crystalline silicon particles and quantum dots can be formed from silicon slurries.3a Sonochemical modification of Zn particles3b leads to the formation of nanocomposite particles where ZnO nanorods are attached to the metallic Zn core. Sonochemically modified Al and Mg3c exhibit a mesoporous inner structure and a significantly increased surface area (up to 300 m² g⁻¹). The mechanism of ultrasound-assisted modification of solid particles in water relies on the interplay of the sonomechanical effect – fragmentation of particles accompanied by the increase of the surface area and the sonochemical effect – the interfacial Red/Ox reactions and etching of a metal surface triggered by free radicals formed during water sonolysis. The contribution of these two sono-effects depends on physical (melting point, hardness, and crystallinity) and chemical (electrode potentials) properties of the metals. Furthermore, the sonochemical

Fig. 1 (a) Temperature programmed desorption of hydrogen from the sonogenerated metal-hydrogen sponge for reactive hard templating (sono-RHT); arrows show a ramping rate increase from 5 K min⁻¹ to 20 K min⁻¹ with the 5 K min⁻¹ step. Insets show SEM and TEM images of sono-RHT. (b) Normalized per number of Mg-atoms desorption energies of hydrogen of different thickness slabs within the DFT level.

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oxidation of “reactive” metals (Al and Mg) is followed by the production of a highly concentrated reducing agent, e.g. hydrogen, on the metal surface.

The “reactive” Al and Mg are not only oxidized during ultrasonic treatment in water, but they also serve as effective donors of electrons that result in H2 production. Sonochemical modification of Al and Mg leads to the simultaneous formation of mesoporous metal frameworks and generation of H2 in the pores. Magnesium as one of the most reactive metals was used as a model reactive metal for the proof of the principles of sono-RHT. The images of Mg used as a sono-RHT are shown in Fig. 1a (insets). Sonication of aqueous suspensions of Mg leads to the formation of highly active magnesium hydrides of different compositions. The temperature programmed desorption of hydrogen (Fig. 1a) indicates the presence of hydrogen in the sonicated Mg. Moreover, several adsorbed hydrogen clusters can be suggested since there are at least two peaks visible in the spectrum.

The calculated energy of hydrogen desorption is ca. 65.7 kJ and ca. 389.3 kJ. The magnesium hydrides rapidly decompose under extreme conditions during sonication in water with the production of H2, and thus, magnesium is a promising material that can be used for the sono-RHT synthesis. The concentration of the residual hydrogen in the sonicated and dried Mg powder was calculated ca. 1.13 μl mg⁻¹.

Both Mg and MgH2 clusters can be formed. Some of the possible geometries of MgH2 were calculated recently by the density functional theory (DFT) (B97) method. Comparably, to our sono-RHT are hydrogen-enriched MgHx clusters, which were also taken into account. The extra hydrogen atoms are less strongly bound to the hydride structure and can therefore be released at lower temperatures. Moreover, in our case, we formed mesoporous sponge structures in which clusters are difficult to be aggregated. Mesoporous sponge has different nanosized units of MgHx slabs in the composition. We calculated for different thickness for MgH2 crystal (100) slabs the energy of total hydrogen desorption from each single slab by DFT within the hybrid PBE0 method. It was shown that complimentary to our experimental data are slabs with thickness between 2 and 3 nm. The entire dehydrogenation of MgH2 slabs, the sum of desorption energies of all H-layers, is about 65 kJ mol⁻¹ for (110) with z = 3 nm (Fig. 1b). This value is very close to the experimentally obtained value. The ΔH298 of entire dehydrogenation was calculated only for the 1 nm slab (110) due to the high computational effort needed. It is 59.3 kJ mol⁻¹, whereas electronic energy is 62.3 kJ mol⁻¹. The difference of 3 kJ mol⁻¹ is vanishingly small, less than the error of application of different exchange-correlation functionals. In such a way, the first peak of H2 thermodesorption (65.7 kJ mol⁻¹) is most probably the desorption of the whole hydrogen from MgH2 nanoparticles of several nanometer size. This is less than ΔH298 (MgH2) = 64.16 kJ mol⁻¹ due to the unique morphology of MgH2 used for sono-RHT and is promising fact for solid hydrogen storage.

Overall, a general sono-RHT synthetic procedure of nanostructures includes the following steps: (i) sonochemical formation of the mesoporous metal RHT; (ii) ultrasound assisted loading of the RHT with chemical precursors for nanoparticle synthesis; (iii) reduction of precursors due to interaction with the sono-activated metal surface; and (iv) the RHT decomposition due to the electrochemical processes stimulated by ultrasound. Herein, we demonstrate the sono-RHT for the synthesis of hierarchically structured materials.

Magnetic nanoparticles were synthesized by reduction of Fe(II) or Fe(III) ions from the corresponding chloride (Fig. 2) or nitrate (not shown here) salts. The formation of the magnetic ion-based materials in the pores of the sono-RHT (Mg) was followed by simultaneous electrochemical degradation of the RHT triggered by sonication. The morphology and composition of magnetic nanoparticles are dependent on the used precursors. The final magnetic materials are dark brownish powders that are composed of 4 mol% 2 mol% Fe3O4 - 32 mol% Fe2O3 (Fig. S2 right and Table S1, ESI†) or 32 mol% Fe2O3 (Fig. S2 left and Table S1, ESI†). By using the sono-RHT method it is possible to control the parameters during the synthesis such as concentration of the RHT, concentration and type of precursors and solvent as well as duration and intensity of sonication, etc. Certain equilibrium structures can be obtained under particular preparation conditions.

The sono-RHT synthesis was performed by sonication of 1 wt% aqueous suspension of ca. 100 μm magnesium particles. Then aqueous solutions of iron salts (to achieve a concentration of 2 wt% in the resultant solution) were added to the suspension. The structures in Fig. S2 left (ESI†) correspond to the samples that are prepared by using a FeCl2 precursor. The structures in
Fig. S2 right (ESI†) correspond to the samples that are synthesized in the presence of a FeCl₃ precursor.

By using the same concentration of the iron salts, Cl⁻ concentration was higher in the presence of FeCl₃. Since the Cl⁻ anions are known as corrosive agents for Mg we expect that the degradation of the metal sono-RHT will be faster at higher chloride concentrations. Indeed, the degradation of Mg was slower in the presence of nitrate and FeCl₂ in comparison to the degradation of the RHT in the presence of FeCl₃. The rate of Mg degradation strongly affects the morphology and composition of the resultant iron-based materials. In the case of slow decomposition of the sono-RHT decomposition process, Mg²⁺ from the metal framework could be involved in the reduction process together with hydrogen as an additional reduction agent. This results in the formation of the iron-based dendritic porous structure on the surface of the metal surface as evidenced by TEM images (Fig. S2 left, ESI†). Since the sonication was performed in aqueous media, reduced iron nanostructures are rapidly oxidized with the formation of various iron oxides. Thus, a hierarchically structured iron based material consisting of a dendritic porous substance covered by nanoparticles was formed by using the sono-RHT method. The chemical composition of these structures will be discussed below. Shortly, the dendritic porous Fe₅O₄→Fe₃O₄ is covered by Fe₂O₃ nanoparticles.

If the reduction agent is mostly H₂, which is produced during the course of the fast decomposition of the magnesium framework (at high Cl⁻ concentration when FeCl₃ is used), rod-shaped Fe₃O₄ nanoparticles are formed (Fig. S2 right, ESI†). The results of the magnetization measurements of the iron based nanoparticles and dendritic structures are shown in Fig. 3. All samples show a superparamagnetic behaviour. The material itself is ferromagnetic, thus the observed magnetic moment is much higher than for paramagnets and the sample does not follow the Curie law. However, there is no hysteresis because the particles are small: they change their orientation when the orientation of the external magnetic field is changed. This is observed when the particle size is smaller than the size of the Weiss areas of the ferromagnet/ferri magnet. The temperature dependence was investigated in order to see, if the movement of the particles is blocked below a certain temperature (blocking temperature), which is not the case.

The Mössbauer spectrum (Fig. 3a) of the dendritic particles (Fig. S2 left, ESI†) can be approximated by a superposition of two doublets and two sextets (Table S1, ESI†). The values of quadrupole splitting (QS) to input A of the doublet of the outer shell and core Fe³⁺ decrease in comparison with the values measured for the nanoparticles mostly due to the size effect. (Fig. S2 left, ESI†). It is seen from Fig. 3a that in the case of the dendritic structure there is one maximum in total QS distribution due to an increase in the range of distributions of QS of each doublet, with a decrease of the value of the quadrupole splitting of iron ions from the outer shell of the nanoparticles. The spectrum and distribution of QS for the dendritic structures suggested their porous character which is also seen from electron microscopy (Fig. S2 left, ESI†). The presence of two sextets indicates a magnetic interaction between iron atoms in the dendritic structure. The parameters (Table S1, ESI†) of the sextets (IS) might be assigned to ferromagnetic magnetite Fe₃O₄. One of the sextets with parameters IS = 0.28 ± 0.01 mm s⁻¹ and H_eff = 48.0 ± 0.02 T corresponds to Fe³⁺ ions, which are in tetrahedral positions (4-position). Simultaneously the second sextet with parameters IS = 0.51 ± 0.01 mm s⁻¹ and H_eff = 44.4 ± 0.02 T corresponds to Fe²⁺½ ions (mixture of Fe³⁺ and Fe²⁺), which are in an octahedron coordination (B-positions) in spinel magnetite. The stoichiometric ratio of the first to the second sextets is close to the theoretical value of 2 [A(2)/A(4) = 2.1].

The Mössbauer spectrum (Fig. 3b) of the nanoparticles (Fig. S2 right, ESI†) is a superposition of two doublets with approximately the same values of the isomeric shift (IS = 0.38 ± 0.02 mm s⁻¹), corresponding to iron(III) ions, and different values of quadrupole splitting (QS = 0.50 ± 0.02 mm s⁻¹ and QS = 0.91 ± 0.02 mm s⁻¹). The parameters can be attributed to Fe₂O₃ in the superparamagnetic state. Thus, intermediate Fe⁰ is not observed in the structure of the nanoparticles. However, the quadrupole splitting suggests the asymmetric density of the electric charge from iron ions in the core (QS = 0.91 mm s⁻¹) to the outer shell (QS = 0.50 mm s⁻¹) of the particle and justifies the assumption of an intermediate.

In order to compare the proposed sono-RHT method with a conventional method of synthesis of iron based materials we performed the synthesis with and without sonochemical irradiation of a solution of the Mg RHT and the iron(III) salt. Two types of particles and their magnetic behaviour are shown in Fig. S1 (ESI†). It is seen that particles prepared using a sono-RHT and described in the paper are different from non-magnetic particles prepared when a Mg RHT was not sonochemically irradiated.
In summary, the combination of the sonochemical approach and the RHT concept for a sono-RHT opens great prospects for the formation of a variety of nanomaterials in aqueous solutions. The structures that can be obtained are affected by the preparation conditions. The acoustic cavitation in water and the shape-defining effect of the reactive template material provide the special thermodynamic and kinetic conditions for the synthesis of nanostructures of particular morphology. The results presented here provide guidelines for the expansion of the concept towards a broad variety of chemical systems. As one of the following perspectives of the method we would like to highlight the formation of biocompatible nanostructures. Thus the hybrid inorganic–organic biocompatible structures with polypyrrole can find their application in biosensors, bioactuators, and even nanorobotics.

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Notes and references