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Recent progress on phase- and morphology-controlled solution combustion synthesis envisages mass fabrications of nanomaterials with more specified phases and morphologies.

Nanomaterials via solution combustion synthesis: a step nearer to controllability

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Solution combustion synthesis (SCS) is a worldwide adopted technique to synthesize nanomaterials, especially for oxides, because of its simplicity, energy and time-effectiveness, and low cost. The general difficulty encountered in SCS is the controllability over phases and morphologies for the products, which arises from the inherent rapid and uncontrollable combustion procedure. In this regard, the present work 10 is devoted to review the recent progress on phase- and morphology-controlled SCS in detail. Besides the various metal oxides, SCS is now applicable to fabricate nanomaterials of metal phosphates, metal silicates, metal borates, metal sulfides, metals, and even alloys, through careful selections of solution compositions. Oxides with regular morphologies of belts, triangles, tubes, wires, and rods can be synthesized by SCS, in the presence of certain templates, or through a self-assembly procedure. The 15 recent progress made on the synthesis of porous materials via SCS is summarized. SCS is also capable of growing metal oxide thin films at low temperatures, enabling the fabrication of low-cost and highperformance electronics on flexible plastic substrates.

1. Introduction

Nanomaterials have sparked worldwide interest because of their 20 unique physical and chemical properties arising from the small dimensions when compared with those of their bulk counterparts. In industry, oxide powders are generally fabricated by hightemperature solid-state reactions, where the desired products are achieved simply by maintaining the solid mixture of reactants at a 25 high temperature. 1,2 Such a ceramic route is energy and time consuming, and the products are usually inhomogeneous, coarse, impure, and away from stoichiometry because of a big diffusion barrier resulted from the lack of molecular-level mixing. Combustion synthesis (CS) is just the choice to achieve a rapid 30 and energy-efficient synthesis. A typical CS procedure utilizes a self-sustained exothermic reaction among well-mixed reactants to achieve the rapid and economical synthesis of particulate products. Up to 2008, CS has been adopted to fabricate more than 1000 kinds of oxide powders³ over more than 65 countries.⁴

Based on the physical nature of the reactants and also reaction mediums, CS can be classified as self-propagating high temperature synthesis (SHS) and solution combustion synthesis (SCS). 4-6 Various materials of oxides, carbides, nitrides, borides, and aluminides have been obtained by SHS, some of which have 40 been commercialized sucessfully. 7-9 In industry, ceramic-lined steel pipes have also been prepared by a centrifugal SHS.¹⁰ The reactants in conventional SHS are all in solid state, still suffering from an inhomogeneous mixing. The coarse initial powders (10-100 µm in size) and high reaction temperatures (>2000 K) result 45 in products with low surface areas and coarse sizes. 4,5 Additional treatments such as mechanical activation, 11 intensive milling, 12 or chemical dispersion^{4,13} are required after SHS to reduce the particle size, which increase the preparation steps and cost. On the contrary, SCS begins from an aqueous solution, which 50 guarantees a molecular-level mixing of reactants. As a result, SCS serves as a promising route for rapid and direct synthesis of mass nanomaterials.

2. Solution combustion synthesis (SCS)

In 1988, Patil firstly reported the synthesis of Al₂O₃ using 55 Al(NO₃)₃ • 9H₂O and urea. ¹⁴ A typical SCS relies on an exothermic reaction between an oxidizer, typically metal nitrates, like Al(NO₃)₃ • 9H₂O, and an organic fuel such as urea. The driving force for lattice formations derives from the internal chemical energy. The temperature input for SCS is not high, 60 which is required just to trigger the combustion reaction, rather than to provide a continuous energy input to form lattices as the case in conventional material fabrication routes. Various watersoluble organics containing large quantities of C and H are utilized as fuels, which facilitates the liberation of heat by 65 combustion. Certain fuels, such as glycine, also form complexes with metal ions to improve the mixing level of the reactants.^{5,15} For example, citric acid bonds with metal ions to form stable chelate complex compounds; amino acids, such as glycine, bonds with metal ions to build very stable chelate complex compounds. 70 The formation of complex helps to achieve a more homogeneous mixing and hence avoids segregations, which is benificial for producing complex and multicomponent oxides.¹⁶

Most of the solution combustion procedures involve a transition from solution to a colloidal sol and then to a gel, which 75 are thus also termed as gel (or sol-gel) combustion. Based on the precursors utilized, the sol-gel method can be categorized into three types: hydrolysis-condensation of metal-alkoxides, the gelation route via concentration of aqueous solutions involving metal-chelates, and the Pechini method.¹⁷ Amough 80 them, the Pechini technique includes a combined process of metal complex formation and in situ polymerization of organics and is versatile for fabricating metal oxides with homogeneous multicomponents. 17 A typical Pechini procedure involves the introduction of metal salts or alkoxides into an 85 ethylene glycol solution that contains citric acid. The formation

of citric complexes leads to a homogeneous mixing and avoids the separation of components at the later stage. The polycondensation of citric acid and ethylene glycol results in the polymer gel formation. A subsequent heating of the gel leads to 5 the removal of the polymer matrix and the formation of the products with a good homogeneity. If metal nitrates are used, instead of the metal salts or alkoxides, as precursors, and fuels are used as complexing agents, the Pechini sol-gel routes would involve a combustion reaction during the final heating process.

In SCS, the phase, morphology, particle size and surface area of products can be altered to certain extents by adjusting the fuel. the fuel/oxidizer ratio, and the pH value of the solution. 15,16 A higher combustion temperature during SCS usually results in improved crystallinity, larger grain size, more serious 15 agglomeration, and lower specific surface area for the products. The propellant chemistry theory is usually used to calculate the stoichiometry of the fuel and oxidizer. 18,19 Taking the combustion reaction between zinc nitrate and urea (CH₄N₂O) as an example, the elemental valency of Zn, C, H, N, and O is +2, +4, +1, 0, and 20 -2, respectively. Accordingly, the oxidizing valency of zinc nitrate and the reducing valency of urea are -10 and +6, respectively. The equilibrium combustion reaction can be described as.

$$Zn(NO_3)_2 + \frac{5}{3}\varphi CH_4N_2O + 5(\varphi - 1)O_2 - \cdots \rightarrow ZnO + \frac{5\varphi + 3}{3}N_2(g) + \frac{5}{3}\varphi CO_2(g) + \frac{10}{3}\varphi H_2O(g)$$

where, $\varphi=1$ means a stoichiometric state, in which atmospheric oxygen is not needed for a complete oxidation of the fuel. $\varphi > 1$ and φ <1 indicates a fuel-rich and fuel-lean condition, respectively. The fuel initiates the combustion reaction with metal nitrates at low ignition temperatures.⁵ When $\varphi=1$, the 30 combustion is intense and complete, which releases lots of heat. φ >1 or φ <1 means one of the reactants is excess in the combustion system. Trace impurities, such as carbon and carbonates, would remain in the product, especially under a fuelrich condition. The phase of products is also determined by the 35 fuel/oxidizer ratio, because the fuel and metal nitrate acts as reducer and oxidizer, respectively. For example, an appropriate fuel/oxidizer ratio in a fuel-rich condition results in the formation of metal rather than metal oxides, as will be discussed latter in Section 3.2. A very high fuel/oxidizer ratio in a fuel-rich 40 condition would result in the formation of amorphous metal complexes, which decompose to form porous metal oxides during the subsequent calcinations (as discussed latter in Section 4.2). Fig. 1 shows a typical combustion procedure for the synthesis of ZnO using urea as a fuel. The combustion reaction sustains after 45 stopping the supply of external energy input, which demonstrates clearly a self-propagating nature of SCS.²⁰

The advantages of SCS can be summarized as follows^{15,16,21}: (1) time and energy efficient; (2) the equipment is simple and the raw materials are usually less expensive; (3) a molecular level 50 mixing for the raw materials; and (4) the composition of products can be tuned. Thanks to such merits, SCS has been adopted to fabricate various materials for many applications of catalysis, 22-30 luminescent materials,³¹⁻³⁶ fuel cells,³⁷⁻³⁹ energy conversion,⁴⁰⁻⁴² and energy storage. 43-46 Complicated Garnets are also widely 55 synthesized by SCS. 47-50 For example, Huczko et al. synthesized cerium-doped yttrium aluminum garnet by SCS. 47 Recently, SCS

is also applied to synthesize thin films of metal oxides. 51-55 Fabrications of low-cost and high-performance electronics at low temperatures are thus possible with the help of SCS.51-55 A 60 detailed description of SCS and its applications is out of the region of this review and the interested readers can refer to other reviews^{3-6,21,22} and books.^{15,16} For example, a book edited by Lackner et al.16 described in detail the characteristics and mechanisms of SCS. The present review focus on the recent 65 progress on SCS to fabricate nanomaterials with more controllable phase and morphology.

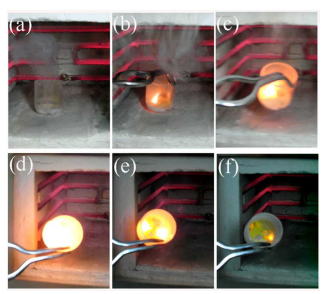


Fig. 1 The typical combustion procedure in SCS: (a) after complete dehydration but before combustion, (b) ignition, (c-f) the combustion 70 process after stopping energy input by taking the reactor out of the oven. 20 Adopted from Ref. 20, Copyright 2013, with permission from Elsevier.

3. Phase control in SCS

Most of binary, ternary or even more complicated oxides have been achieved by SCS; on the contrary, reports on nanomaterials 75 other than oxides are relatively rare. Encouragingly, it is reported that many phosphates can be obtained by SCS followed by a subsequent heat treatment, in the presence of additional $(NH_4)_2HPO_4$ or $NH_4H_2PO_4$. $^{56-61}$ Li_2SrSiO_4 and Li_2FeSiO_4/C^{63} are also prepared by SCS and a subsequent heat treatment, in 80 which SiO₂ serves as a silica source. Wang et al. synthesized copper-doped Li₂B₄O₇ by SCS and a subsequent heat treatment, using H₃BO₃ as a boron source. 64 Recently, metal sulfides, pure metals, and alloys have been achieved by SCS via carefully selecting fuels, changing the ratio of fuel to oxidizer, and 85 controlling the reaction atmosphere.

3.1 Synthesis of metal sulfides

Tukhtaev et al. used thiourea or thiosemicarbazide as a fuel, which also complexed with metal ions, to prepare various metal sulfides by SCS in an inert atmosphere. 65 The particle size and morphology of the products are reported to be dependent on the atmosphere pressure. 65 CdS with crystallite size of 10~15 nm was also obtained by a microwave-assisted combustion synthesis using thiourea as a fuel.66 Amutha et al. also obtained CdS nanoparticles by the microwave-assisted combustion synthesis

using cadmium thiocyanate complex as a single source precursor.⁶⁷ The key to achieve metal sulfides instead of the corresponding metal oxides is the selection of appropriate fuel, such as thiourea, which contains abundant sulfur. However, the 5 fabrication of other metal sulfides beside CdS by SCS in an air atmosphere is rarely reported. Further work is needed to clarify the exact formation mechanism of CdS during SCS in an air atmosphere.

3.2 Synthesis of metals and alloys

10 Fuels in SCS act as reducing agents; therefore, it is possible to further reduce the metal oxides produced in SCS by the residual fuel. Rao et al. obtained metallic Cu, Ni, and CuNi alloys by SCS using N-tertiarybutoxy-carbonylpiperazine (C₉H₁₈N₂O₂) as a fuel.⁶⁸ Later, Jung et al. directly synthesized nickel particles 15 through a microwave-induced SCS with a fuel-rich composition in an air atmosphere without a further reducing calcination process.⁶⁹ Erri et al. chose an appropriate fuel/oxidizer ratio to control the combustion velocity and thus a nickel foam was obtained. 70 The fuel-rich condition was demanded to provide a 20 reducing atmosphere for the metallic Ni formation. 70

Jiang et al. also used SCS to prepare various metal and alloy nanoparticles of Ni, Co, Cu, Ag, Bi, Ni_{0.5}Co_{0.5} in an inert atmosphere. 71 H2, H2O, CH4, NO, CO2, NH3, and NO2 species were detected near the combustion temperature. Among them, H₂ 25 and CH₄ have the reducing ability to obtain metals from the corresponding metal oxides. An appropriate fuel/oxidizer ratio is critical for the synthesis of metals by SCS. A low fuel/oxidizer ratio restricts the reduction reaction; however, when the ratio is too high, the combustion temperature decreases, which also 30 inhibits the reduction of metal oxides.⁷¹

Mukasyan's group investigated the reaction pathways of metal nanopowders in SCS both in experiment and thermodynamic analysis, taking nickel⁷² and copper⁷³ as a model, respectively. The formation of nickel oxide appeared at the early stage in the 35 reaction front, followed by a reduction to nickel in the post combustion zone. The reducing environment was achieved in fuel-rich conditions.⁷² Similar reaction pathways were also observed in a copper nitrate-glycine system.⁷³ Recently, they further used several in situ techniques and electron microscopy to 40 study the mechanism in combustion reaction of a nickel nitrateglycine system.⁷⁴ The results further confirmed that metal oxides formed at the early stage by decomposition of metal nitrate, which were then reduced to the corresponding metals by the reducing gases released in the combustion process in fuel-rich 45 conditions. 74 In the nickel nitrate-glycine system, the source of energy for self-sustained reaction was the exothermic reaction between N₂O and NH₃ species from the decomposition of nickel nitrate and glycine. 74 The excess NH3 gas produced in fuel-rich conditions reduces nickel oxide to nickel in the reaction front.⁷⁴ 50 The mechanism of combustion synthesis of Ni in nickel nitrate-glycine system is summaried in Fig. 2. It is not favorable to obtain metallic iron in the iron nitrate-glycine system because iron(II) oxide can not be reduced to iron by NH3 (or H2) even at 1500 °C, as suggested by a thermodynamic calculation. 74 Pure 55 iron was recently prepared by combustion of an iron nitrate-citric acid system in a nitrogen atmosphere, using the mixture of ethanol and deionized water as a solvent. 75 Interestingly, Fe₃C was also obtained by further increasing the amount of citric acid

(fuel).⁷⁵ It is also found that more completed reduction of Ni₁₋ 60 Co, alloys can be achieved by adding appropriate ethanol, which thus improves the saturation magnetization.⁷⁶ The additive of a suitable amount of ethanol was supposed to increase the reduction ability of the sol-gel combustion process. 75,76 The metals and alloys obtained by SCS, including Fe, Co, Ni, Cu, Ag, 65 Bi, Ni-Co alloys, Ni-Cu alloys, and Ni-Cu-Fe alloys, are summarized in Table 1.

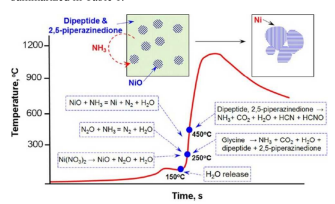


Fig. 2 A schematic illustration showing the mechanism of Ni formation during self-sustained combustion reactions of nickel nitrate-glycine Adopted from Ref. 74, Copyright 2013, with permission from American Chemical Society.

4. Morphology control in SCS

Controlling nuclei formation and the subsequent crystal growth is crucial for the synthesis of high-quality crystals with certain 75 morphologies. The rapid combustion reaction and less controllable combustion process make it difficult to control the morphology of products in SCS. The typical morphology of products by SCS is agglomerations of irregular nanoparticles.

4.1 Products with regular shapes

80 Introducing appropriate templates in SCS results in special nanostructures. For example, hydroxylapatite nanotubes have been obtained by introducing a porous anodic aluminum oxide (AAO) template in SCS.⁷⁹ After the removal of the AAO template, uniform hydroxylapatite nanotubes with a fiber-brush 85 like shape were obtained, as shown in Fig. 3a-c. The length and diameter of the nanotubes is ca. 60 µm and 100 nm, respectively. The TEM image (Fig. 3c) clearly shows the hollow structure. Just like other template-assisted synthesis methods, the preparation and removal of templates make the synthesis procedure complex 90 and expensive.

Interestingly, one-dimensional nanostructures can sometimes be obtained by annealing the preformed nanoparticles derived form SCS. 80-82 For example, Ding et al. obtained CaIn₂O₄ nanorods (Fig. 3f) by SCS combined with a high-temperature 95 post-annealing. 80 Firstly, CaIn₂O₄ nanoparticles were formed by SCS (Fig. 3d). Then, the as-combusted CaIn₂O₄ nanoparticles self-assembled into nanocapsules and subsequently formed the regular rods (Fig. 3e, f). The formation of rods was believed to be related to the asymmetric crystal structure of CaIn2O4 and the 100 instantaneous high temperature involved in the combustion.⁸⁰ V₂O₅ rods were also synthesized by SCS combined with a subsequent annealing at 550 °C for 24 h.81 Chen et al. also obtained Sn_{1-x}RE_xO_{2-x/2} (RE=Y, La, Gd and Nd) nanorods by

annealing the as-synthesized products from a microwave-induced KCl-assisted SCS.⁸² The diameter and length of the nanorods is 8-12 nm and 100-200 nm, respectively. The preformed nanoparticles by the KCl-assisted SCS transfer to nanorods in the 5 molten salt during annealing.⁸²

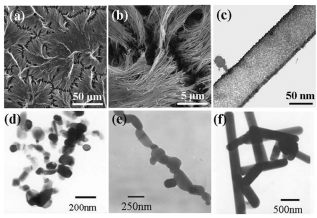


Fig. 3 SEM (a, b) and TEM (c) images of hydroxyapatite nanotubes.⁷⁹ TEM images of CaIn₂O₄: (d) as-combusted, (e) post-annealed at 1,373 K for 2 h and (f) post-annealed at 1,373 K for 12 h.80 (a)-(c) adopted from 10 Ref. 79, Copyright 2008, with permission from Elsevier. (d)-(f) adopted from Ref. 80, Copyright 2009, with permission from Springer.

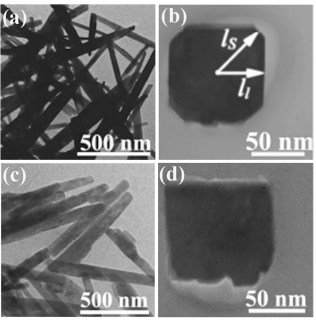


Fig. 4 TEM images of (a, b) Al₄B₂O₉ nanowires synthesized at 900 °C for 2 h and (c, d) $Al_{18}B_4O_{33}$ Nanowires synthesized at 1050 °C for 2 h.84 15 Adopted from Ref. 84, Copyright 2007, with permission from American Chemical Society.

More interestingly, calcium phosphate nanowhiskers were directly prepared by a microwave-assisted auto ignition molten salt synthesis without the subsequent annealing.⁸³ NaNO₃ was 20 used as fluxing agent and urea served as the fuel for combustion. The combustion process raises the extent of superheating in the molten-salt bath. Embryonic particles of calcium phosphates are formed through the nucleation and growth procedure. Calcium phosphate nanowhisker growth occurs by a sequential 25 "dissolution-crystallization-whisker growth" process as the

molten ionic bath cools down.83 Tao et al. achieved singlecrystalline Al₄B₂O₉ (Fig. 4a, b) and Al₁₈B₄O₃₃ nanowires (Fig. 4c, d) by the one-step high-temperature calcination at 900 °C and 1050 °C, respectively, in which the combustion were maintained 30 for ca. 6 min and final products were obtain after 2 h annealing.⁸⁴ The cross-sectional observation reveals that the Al₄B₂O₉ nanowires, which grew along the [100] direction, exhibited four large facets with lower surface energy and four small facets with high surface energy (Fig. 4b). For the Al₁₈B₄O₃₃ nanowires, 35 which grew along the [001] direction, the small facets with high surface area disappeared, resulting in a nearly squre cross section (Fig. 4d). The anisotropical growth of both nanowires was believed to result from the requirement to minimize the surface energy for a free particle. The single crystalline nature for the 40 nanowires with high purity seems to be the key to achieve the nanowires during the SCS procedure.

Cheng et al. used SCS followed by a subsequent hightemperature calcination to synthesize Eu3+ incorporated ZnO belts⁸⁵ (Fig. 5a, b) and SrAl_xO_y: Eu²⁺, Dy³⁺ belts (Fig.5c, d).⁸⁶ 45 The introduction of N(CH₂CH₂OH)₃ (triethanolamine, TEA) is believed to be the key to achieve the final specific morphology, because it is a multifuctional reagent serving simultaneously as a chelating agent to form metal-TEA complexes, a surfactant to control the shape of precursors, and also a a fuel that is capable of 50 reducing the reaction rate of the combustion synthesis. 85,86 Mapa et al. synthesized N doped ZnO micrometer-sized triangular particles (Fig. 5e) by a combustion reaction between urea and zinc nitrate with a molar ratio of ca. 1.87 The size is relatively less uniform and is large in micro-scale rather than in nano-scale. The 55 authors argued that, when the urea/Zn(NO₃)₂ molar ratio increased to beyond 3, an increasing exothermicity was achieved for the combustion reaction. As a result, no specific morphology can be observed for the resultant particles.⁸⁷ The trangular particles are believed to result from the stress-free and well-60 grown crystals, which is expected from slow crystallization. The exact mechanism for the formation of the trangular-shaped particles by SCS, which is indeed a rapid preparation method, is therefore surprising; however, Mapa's study suggests that, for polycrystalline oxides, a moderate combustion favors the 65 formation of particles with specific morphologies.

Recently, copper oxide (CuO) nanoflowers, as shown in Fig. 5f, were prepared by SCS, where glycine was used as a fuel.⁸⁸ contain CuO nanoflowers well-crystallized nanopetals/sheets. The flower-like structure was believed to come 70 into being through the local sintering of the preformed nanoparticles during the high temperature combustion.

Although some progresses have been made, most of regular morphologies reported to date are in micro-scale or need a subsequent annealing. For the AAO-assisted SCS to obtain one-75 dimensional (1D) nanostructures, the penetration of the precursor into the templates is the key point. ⁷⁹ For crystals with asymmetric structures, the subsequent annealing is capable of inducing selfassembly of nanoparticles preformed during the SCS procedure, especially for those in the molten salts, to achieve 1D 80 nanostructures. 80-82 It still remains a challenge to directly obtain regular and uniform nanostructures by SCS. The exact mechanism for the formation of the unique nanostructure during SCS is still unclear because of the rapid and unknown

combustion process. For asymmetric crystals with high purity, it is possible to achieve directly 1D structures because of the driving force for minimizing the total surface energy.⁸⁴ For 3D structures of flower-like particles, an appropriate combustion 5 environment favoring the local sintering of the preformed nanoparticles is of importance. 88 Further work is needed to clarify the exact formation mechanism of the various morphologies. We believe a deep understanding on the nucleation and crystal growth procedure characteristic of SCS would help clarifying the 10 formation of the specific nanostructure during SCS. In situ electron microscopic observation would be also helpful to understand the mechanism of morphology evolution during SCS.

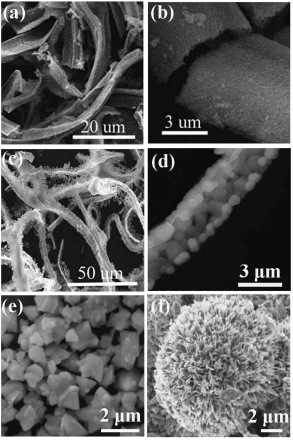


Fig. 5 SEM images of the ZnO post-annealing at 600 °C (a, b), 85 SrAl_xO_v 15 post-annealing at 1300 °C (c, d), 86 triangular ZnO (e), 87 nanoflowers (f).88 (e) adopted from Ref. 87, Copyright 2009, with permission from American Chemical Society. (f) adopted from Ref. 88, Copyright 2013, with permission from Elsevier.

4.2 Porous materials

20 Porous materials have promising applications in the various fields of gas sensors, catalysis, separation, energy storage and energy conversion. 89-91 Most methods for synthesis of porous materials demand the assistance of templates: hard templates (mesoporous silica, silica sphere, or polystyrene sphere) and soft templates 25 (surfactants or block copolymers). 92,93 Because of the advantages of rapid and energy efficient, it is interesting to adopt SCS to synthesize porous materials, which is simple, template-free and low cost. Sometimes porous structures can be obtained as a result of the gas evolution during the combustion reaction; however, the 30 pores are usually inhomogeneous and most of pore size is in tens

of micrometers. It is of importance to obtain relatively homogeneous porous materials with pore sizes ranging from nanometers to several micrometers, which is favorable for gas or liquid diffusion and provides a relatively large surface area.

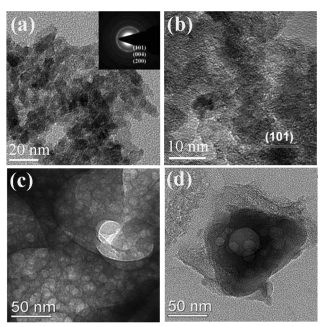
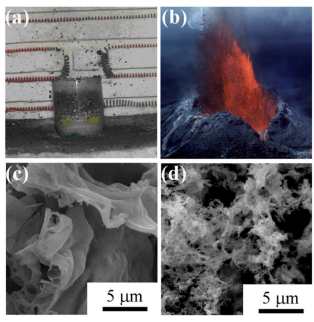


Fig. 6 (a) TEM image and (b) HRTEM image of TiO2 with wormhole mesopores. 98 (c, d) TEM images of porous Li₃V₂(PO₄)₃/C. 61 (c), (d) adopted from Ref. 61, Copyright 2012, with permission from Elsevier.

Phase and surface area of products in SCS are greatly affected 40 by fuels and fuel/oxidizer ratio. Anatase TiO2 with high spedific surface area and disordered mesopores (Fig. 6a, b) was successfully synthesized through SCS by using many kinds of fuels, such as glycine, ⁹⁴⁻⁹⁷ urea, ⁹⁸⁻¹⁰¹ hexamethylenetetramine, ⁹⁵ and oxalyldihydrazide. ⁹⁵ Zhang et al. obtained porous 45 Li₃V₂(PO₄)₃/C (as shown in Fig. 6c, d) with a surface area of 25.7 m² g⁻¹ by annealing sucrose and combustion-driven amorphous precursor at 800 °C for 8 h in a N₂ atmosphere, and this material exhibited excellent electrochemical properties. 61 Mesoporous MgO with a surface area of 107 m²/g was obtained by SCS, 50 which exhibited excellent fluoride removal capacity. 102 Mesoporous alumina with high surface area and narrow pore size distribution was also prepared by SCS in a fuel (urea)-lean condition. The surface area decreased with increasing amounts of urea. 103 However, the as-synthesized alumina was amorphous and 55 the surface area greatly decreased upon the subsequent crystallization procedure. 103 It is still difficult to extend this procedure for synthesis of other porous oxides with relatively uniform pores because of the unknown and uncontrollable combustion process. 16

Chen et al. is the first to introduce salts in SCS, which greatly increases the surface area of products in SCS. 104 The role of salts is believed to be two folds: decreasing the reaction temperature because the salt additives absorb reaction heat; salt precipitated in situ coats the newly formed nanoparticles, preventing them from 65 sintering and agglomerating. 104-107 The salts are inexpensive and can be easily removed from the products by water washing. For the stoichiometric combustion reaction between nickel nitrate hexahydrate and citric acid, NaF is found to be more effective

than NaCl for improving the surface area. 108 More interestingly, the additive of appropriate amounts of NaF alters the combustion mode, which results in the appearance of the so-called eruption combustion. In the eruption combustion process, lots of powders 5 were lifted off and naturally fall down around the container, leaving highly fluffy products with a quite big volume, as shown in Fig. 7a. This phenomenon is very similar to a volcano eruption in nature (Fig. 7b). The eruption combustion in nickel nitratecitric acid system resulted in smaller particles, improved 10 dispersity, and remarkably increased specific surface area, as shown in Fig. 7c, d. 108 Moreover, it was shown that introduction of high surface area carriers in SCS also helps to obtain high surface area oxide-based supported catalysts. 109



15 Fig. 7 Optical photographs of eruption combustion process (a) and volcanic eruption in nature (b). The optical photograph showing a natural volcanic eruption is from the Internet. SEM images of Ni/NiO powders derived with conventional combustion (c) and eruption combustion (d). 108 Adopted from Ref. 108, Copyright 2011, with permission from American 20 Chemical Society.

Fuel-rich conditions in SCS usually result in residual organics in the products. Carefully selecting fuels and adopting a high fuel/oxidizer ratio can obtain an organic-rich and amorphous precursor, which further decomposes to form porous metal oxide 25 by the subsequent calcination because of gas evolutions. By using this method, homogeneously macroporous ZnO can be achieved, as shown in Fig. 8a, b. 110 The synthesis route can also be extended to the preparation of other porous oxides. For example, porous Co₃O₄ (Fig. 8c, d) can be also obtained by this method. 111 30 The decomposition temperature of the Co-based precursor is much lower than that of the Zn-based precursor. 110,111 The low decomposition temperature of the Co-based precursor is benificial for the formation of the mesoporous Co₃O₄ network (Fig. 8c). Increasing the calcination temperature results in a larger 35 particle size and larger pore size of the porous Co₃O₄, as shown in Fig. 8d, due to the enhanced grain growth. 111

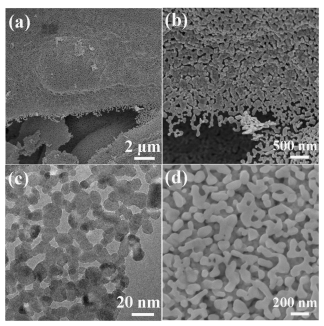


Fig.8 (a, b) SEM images of porous ZnO network. 110 (c) TEM image of mesoporous Co₃O₄ calcined at 355 °C. 111 (d) SEM image of macroporous 40 Co₃O₄ calcined at 550 °C. 111 (a), (b) adopted from Ref. 110, Copyright 2012, with permission from AIP Publishing.

Hydrazine has a high energy N-N bond and lone electron pairs, which can coordinate with metal ions to form metal complexes.¹⁵ Such metal hydrazine complexes can serve as low-temperature 45 precursors to oxide materials. 112-114 The energetic complexes usually decompose in an exothermal way along with combustion at low temperatures. Moreover, the raw materials for the complexes are water-soluble. Thus, the synthesis route starting from metal hydrazine complexes is very similar to the SCS 50 procedure and can be classified as SCS. Recently we used metal nitrate, metal acetate, hydrazine hydrate, and glycine to synthesize homogeneous macroporous NiO/Ni115 and porous ZnO. 116 The morphology of macroporous NiO/Ni can be changed by adjusting the complex composition. For example, 55 macroporous NiO/Ni with three-dimensional (3D) and twodimensional (2D) architectures can be achieved, as shown in Fig. 9a-d. The content of NiO in the macroporous NiO/Ni can be controlled by a subsequent calcination, with the macroporous structure preserved. 117 The obtained ZnO exhibits a coral-like 60 macro-/mesoporous structure (Fig. 9e, f). 116 Interestingly, this kind of metal hydrazine complexes can transform to macro-/meso-/ microporous nanofoam structures under focused electron beam irradiation during TEM operation. For example, porous ZnO nanofoams with ultrathin walls (Fig. 10a-c) are obtained via 65 focused electron beam irradiation of the zinc hydrazine complex, whose preparation process is the same to that in the reference. 116 As shown in the HRTEM image (Fig. 10c), the measured lattice spacing of 0.25 nm and 0.26 nm represents the (101) and (002) atomic plane of wurtzite ZnO, respectively. The SAED pattern 70 (Fig. 10d) further demonstrates that the product is polycrystalline wurtzite ZnO. This difference of mophorlogy (comparing Fig. 10a-c with Fig. 9e, f) may be attributed to the different heating rates and atmospheric pressures. The much higher heating rate of the focused electron beam and lower atmospheric pressure during 75 TEM operation result in the formation of porous nanofoams with

ultrathin walls. This one-step route to porous structures is simple and rapid; however, hydrazine is toxic, and the reaction is relatively violent and hence careful operation and safety protection are required.

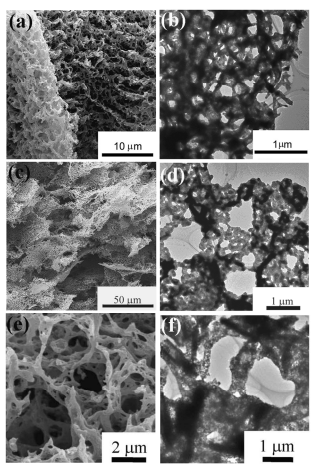


Fig. 9 SEM image (a) and TEM image (b) of 3D macroporous NiO/Ni. SEM image (c) and TEM mage (d) of 2D macroporous NiO/Ni. 115 SEM image (e) and TEM image (f) of 3D porous ZnO. 116 (a)-(d) adopted from Ref. 115, Copyright 2013, with permission from Elsevier.

10 4.3 Thin films

SCS is more commonly used to produce materials in form of powders. Many precursors employed in SCS have an uncanny resemblance to corresponding compounds utilized in sol-gel routes.²¹ Thus, it is possible to obtain metal oxide thin films by 15 SCS. Recently, Marks et al. adopt SCS as a new general route to fabricate diverse metal oxide thin films of In₂O₃, Zn-Sn-O, In-Zn-O, and ITO.51 Taking the synthesis of ZnO films for example, 297.5 mg Zn(NO₃)₂ • 6H₂O and 100.1 mg urea were dissolved in 5 ml 2-methoxyethanol and aged for 72 h. Then, the films were 20 fabricated by spin coating and annealed at the desired temperature (150-400 °C) for 30 min in air. 51 Fig. 11a-c shows the SEM images of SCS-derived In₂O₃ thin films with controlled thickness.⁵¹ Thin films with a thickness of 30 nm or less exhibit smooth surfaces and negligible porosity; on the contrary, those 25 with a thickness over 70 nm exhibit voids and evidence of porosity because of gas release during SCS. Growth of thicker and denser films thus demands multiple depositions.

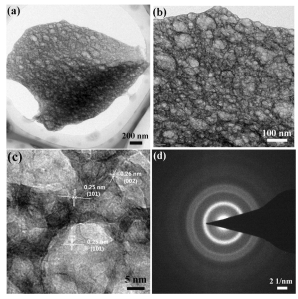


Fig. 10 TEM images (a, b), HRTEM image (c), and (d) of porous ZnO 30 nanofoams derived from focused electron beam irradiation of the zinc hydrazine complex under TEM operation.

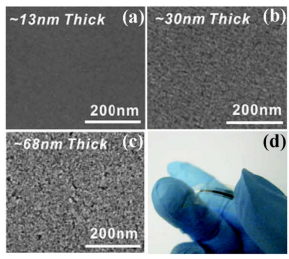


Fig. 11 (a-c) SEM images of In₂O₃ films on n++ Si with controlled thickness. (d) Optical image of a flexible combustion-processed In2O3 35 device. 51 Adopted from Ref. 51, Copyright 2011, with permission from Macmillan Publishers Ltd.

The driving force of oxide lattice formations in SCS is from the internal chemical energy; therefore, the required temperature is just for reaction initiation rather than continuous energy input. 40 These unique advantages endow SCS the ability to grow metal oxide films at temperatures as low as 200 °C.51 It is thus convenient to achieve optically transparent transistors on flexible plastic substrates (Fig. 11d). 51,52 In a word, SCS enables the fabrication of low-cost and high-performance electronics at low 45 temperatures. 51-55

5. Conclusions and perspectives

Solution combustion synthesis has been adopted worldwide to synthesize numerous nanomaterials, especially simple and complex oxides, because of its simplicity, energy- and time-50 effectiveness, and low cost. The recent progress on controlled synthesis in SCS is described in detail in this review. It is a big challenge to obtain products with well defined phases and morphologies by SCS, which is partially attributed to the rapid and uncontrollable combustion procedure.

- Besides the various metal oxides, it is now possible to synthesize metal sulfides, metals, and alloys, through careful selections of fuel and appropriate fuel/oxidant ratio. The metals or alloys are believed to result from the further reduction of the pre-formed corresponding metal oxides by the reducing 10 atmosphere, which comes from either the excessive reducing fuel or the gas evolution during the combustion in fuel-rich conditions. The synthesis of other species, such as metal nitrides, metal carbides, and metal borides for a wider application still remains a challenge.
- Oxides with regular morphologies of belts, triangles, tubes, wires, and rods can be synthesized by SCS, in the presence of certain templates, or through a self-assembly procedure. Utilizing the driving force for minimizing the total surface energy, SCS is capable of fabricating directly 1D structures when the target 20 products are of asymmetric crystals. To achieve 3D structures of flower-like particles in one-step by SCS, an appropriate combustion environment should be maintained to favor the local sintering of the preformed nanoparticles to the resultants with specific morpholgies.
- More commonly, products in SCS are aggregates of nanoparticles, with relatively low specific surface area. The saltassisted SCS favors the synthesis of oxides with small particles and thus high specific surface area. The additive of NaF in certain SCS system induced a novel eruption combustion mode, which 30 further improves the dispersion and specific surface area of the produced oxides. Organic-rich precursor can be obtained by carefully selecting fuels and adopting a high fuel/oxidizer ratio, which further decomposes to form porous metal oxide by the subsequent calcination because of gas evolutions. Decomposition 35 of metal hydrazine complexes is capable of producing oxides with high porosity. By adjusting the complex intensity, the architecture of the products can also be controlled. SCS is also developed to fabricate metal oxide thin films at low temperatures, enabling the fabrication of low-cost and high-performance 40 electronics on flexible plastic substrates.

With the efforts of researchers worldwide, remarkable progresses have been made towards a more controllable SCS. However, the exact mechanism of SCS still remains unclear and it is a big challenge to achieve SCS products with well-defined 45 phase structure and pre-designed morphology. Further work is needed to clarify the phase and microstructure evolution in the combustion process and to effectively control the combustion process.

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Notes and references

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Table 1 Various metals and alloys derived from SCS.

Metals or alloys	Fuel	Other starting reactants	Fuel to oxidizer molar ratio	Atmosphere	Solvent	рН	Heating temperature / °C	Pre-dried temperature / °C	Ref.
-carbonylpiperazine	nitrates							,	
Ni	glycine	$Ni(NO_3)_2 \cdot 6H_2O$	2.44	air	water	\	microwave	\	69
Ni, Cu, Co, Ni-Cu, and Ni-Co	glycine	Hydrated metal	1.4 for Ni, 2.8 for	air	water	\	-	\	70
		nitrates	Cu, and 1.7 for						
			$Ni_{0.9}Cu_{0.1}$						
Co, Ni, Cu, Ag, Bi, and Ni-Co	citric acid	Hydrated metal	0.8-1.2 for Ni	N_2	water	7	300	95	71
		nitrates							,
Ni	glycine	$Ni(NO_3)_2 \cdot 6H_2O$	1.4	air	water	\	-	-	72
Cu and Ni-Cu	glycine	Hydrated metal	3.3 for Cu and 1.9	air	water	\	-	-	73
		nitrates	for Cu-Ni						1
Fe	citric acid	Fe(NO ₃) ₃ • 9H ₂ O	1.25	N_2	water+	6	300, 400, 500,	95	75
					ethanol		600, 700		
Ni-Co	citric acid	Hydrated metal	1.0	N_2	water+	7	300, 400, 500,	90	76
		nitrates			ethanol		600, 700		
Cu, Ag, and Ni	glycine	Hydrated metal	2.0 for Ni	air	water	\	250	\	77
		nitrates							
Ni-Cu-Fe	glycine	Metal nitrates	1.9	air	water	\	-	-	78

[\] means without treatment

⁻ means that it was not described in the reference.