

Anne Aimable¹
 Nathalie Jongen¹
 Andrea Testino¹
 Marcel Donnet¹
 Jacques Lemaître¹
 Heinrich Hofmann¹
 Paul Bowen¹

¹Ecole Polytechnique Fédérale de Lausanne (EPFL), Swiss Federal Institute of Technology, Materials Institute, Powder Technology Laboratory (LTP), Lausanne, Switzerland.

Research Article

Precipitation of Nanosized and Nanostructured Powders: Process Intensification and Scale-Out Using a Segmented Flow Tubular Reactor (SFTR)

The successful scale-out and process intensification using a segmented flow tubular reactor (SFTR) for ultrafine CaCO₃, BaTiO₃, and nanosized ZnO from optimized minibatch (20 mL) conditions is presented. The capacity of the SFTR in process intensification was demonstrated by producing ~5 kg batches of BaTiO₃ powders with excellent batch-to-batch reproducibility. The SFTR scale-out or numbering-up capacity was demonstrated for a nanostructured CaCO₃ in 500 g batches by scaling-out from one to six segmented flow tubular reactors run in parallel (scale-out/-up ratio of 5000 compared to lab batch experiments). The SFTR was then used to demonstrate its potential for nanosized ZnO powders producing 50 g lots of these nanopowders in a continuous process, a scale-out/-up ratio of 250 compared to lab batch experiments without any loss of powder quality. The SFTR allows a precise control of precipitation conditions, leading to an excellent reproducibility in powder characteristics, and shows great promise as a simple production process of powders and advanced nanomaterials with highly controlled properties.

Keywords: Nanoparticles, Plug flow, Powders, Precipitation, Segmented flow tubular reactor

Received: July 30, 2010; *revised:* August 26, 2010; *accepted:* October 19, 2010

DOI: 10.1002/ceat.201000324

1 Introduction

Nanomaterials and nanotechnology attract an extraordinary amount of interest, as shown by the increasing amount of publications and patents published on the subject every year and the continuous growth of funding: in the United States, the federal funding for nanotechnology has increased from approximately \$464 million in 2001 to nearly \$1.5 billion in 2009 and was more than \$5 billion worldwide [1]. The unique properties of these nanomaterials bring them into a key position for a large spectrum of devices and systems [2]. But after 20 years of exciting new discoveries, researchers face the real challenge: to convert science-based findings and research laboratory methods into applied technologies and processes suitable for use in a commercial environment. This is more and

more emphasized in current reviews on nanotechnology [3, 4], as, e.g., by Ashby et al. [5]: “Making effective design application use of new scientific findings that seemingly appear every day within the nanomaterial field is not as easy as might first appear”, or by Pitkethly [6]: “There are still many challenges for nanomaterials companies to overcome before the potential is fully realized. These include: how to produce materials in volume commercially at viable prices...//...consistency and reliability in volume production.”

Projects funded by the European Union also show this necessity in developing new solutions for nanopowder production in a volume that could lead to real applications. A specific call NMP-2008-2.1-2 *Processing and Upscaling of Nanostructured Materials* was dedicated to this issue in the seventh Research Program FP7 in 2008 *Nanosciences, Nanotechnologies, Materials and New Production Technologies*.

Over recent years a new type of tubular plug flow reactor – the segmented flow tubular reactor (SFTR) – has been conceived and used in powder synthesis [7]. This type of tubular or segmented flow reactors has shown excellent performances in liquid-liquid reactions as microreactors with segmented flow in capillaries or microchannels [8–10]. The microfluidics

Correspondence: Dr. A. Aimable (anne.aimable@epfl.ch), Ecole Polytechnique Fédérale de Lausanne (EPFL), Swiss Federal Institute of Technology, Materials Institute, Powder Technology Laboratory (LTP), Station 12, 1015 Lausanne, Switzerland.

of such reactors has been studied in detail [11, 12] showing the advantages of the increased mass and heat transfer between the two liquid phases, which enhance conversion and selectivity for phase transfer-catalyzed reactions [13]. These microreactors have also improved performance efficiency in liquid-liquid extraction [14] and have great potential in synthetic biology [15]. When compared with batch reactors, they are of great interest for both liquid-liquid reactions and powder production targets because they can be operated in a continuous mode and ensure controlled hydrodynamics, residence times, and reaction conditions.

For powder synthesis the main tool used has been the SFTR which has proved its versatility and its robustness through the preparation of different products: CaCO_3 [16], BaTiO_3 [17], copper, calcium and nickel-manganese oxalates [18, 19] with a high powder quality (chemical and phase composition, particle size, and shape) and reproducibility [20].

The use of droplet-based microfluidics proved to be a viable method for nanosized powders [21] but has not been fully exploited with respect to the scale-out/-up and process intensification potential which reactors such as the SFTR possess. This paper presents examples of both scale-out from a single-tube to a six-tube reactor for nanostructured CaCO_3 and the process intensification showing its capabilities to produce kg quantities of high-quality BaTiO_3 powders. The main subject will focus on the synthesis of nonagglomerated redispersible nanoparticles of ZnO (< 100 nm) by using the SFTR.

ZnO has already many applications, such as gas sensors or varistors [22]. Optical properties are also widely exploited as a dye (zinc white) or UV-light emitter [23]. By decreasing the size, ZnO can be used as quantum dots [24]. The synthesis of ZnO nanoparticles (< 100 nm) with a homogeneous morphology and a narrow particle size distribution was investigated in a previous study using a small batch reactor (reactor volume 40 cm^3 , powder yield 0.2 g) [25]. The size and morphology of the precipitated ZnO can be tuned with the addition of polymers or by modifying the pH. By conducting the synthesis under basic conditions, the morphology was modified towards a flower-like shape with an aspect ratio of the primary particles increasing with the pH. Under slightly acidic conditions, the anionic additives poly(acrylic acid) (PAA 2000, Dispex A40) and sodium dodecyl sulfate (SDS) reduced the final particle size and degree of agglomeration by an assumed preferential adsorption on the (0001) positive surface. Nonagglomerated equiaxed nanoparticles (around 100 nm) with a narrow size distribution (span < 1) could be obtained with a very good reproducibility. These features make these materials very interesting for further applications in electronic devices and should allow a higher density after sintering.

In order to investigate more such properties and conduct ceramic processing and sintering tests, it is necessary to have batches of powders of minimum size, at least 50 g, whereas a typical laboratory batch process only produced 0.2 g per batch. The present paper describes the transfer of ZnO synthesis from the minibatch to a continuous production using the SFTR with a significant scale-up ratio of 250. The powder characteristics obtained with a simple batch volume or with the SFTR is compared and the robustness of the SFTR is investigated. The effect of the micromixer used in the process on the material

properties is also studied, as it is well-known that the mixing conditions can determine the resulting product properties in precipitation systems, especially influencing the particle size [26–28]. A key factor for nanopowder production is its colloidal stability and capacity to be dried and redispersed without loss of powder quality. There are many nanopowders available in small quantities and some can be stabilized in suspension but suffer significant nonreversible agglomeration on drying. The cost of transporting and storing suspensions even at 30 % weight solids fraction is significantly higher than transporting a dried powder – the key is the ability to redisperse the powder after drying. This will also be demonstrated for the ZnO powder investigated in the current study.

2 Materials and Methods

2.1 The Segmented Flow Tubular Reactor: SFTR

The SFTR has been developed to overcome the classical problems of powder production scale-up from batch processes, which are mainly linked with mixing, homogeneity, and heat transfer. A schematic view of the SFTR is given in Fig. 1a). The SFTR is composed of three distinct parts: a micromixer which ensures that the coreactants are efficiently mixed, a segmenter, and a tubular reactor, placed in a thermostatic bath. The reactants are introduced through the micromixer where the initial supersaturation is created. The reacting mixture is then segmented by an immiscible solvent: nitrogen gas in the case of BaTiO_3 precipitation, dodecane in the case of CaCO_3 , and ZnO precipitation. Small suspension volumes ($\sim 0.2 \text{ cm}^3$) are thus created, producing microdroplets or microreactors, where precipitation and growth takes place. These small volumes ensure a high homogeneity inside each droplet, all circulating through the tube with an identical *history* (e.g., residence time and heat exchange). Finally, fouling at the wall surface is avoided due to the immiscible fluid by forming a thin film at the surface [12] and thus allowing long continuous process times (> 12 h). The residence time is determined by the pump flow speeds, the tube length, and reaction kinetics and was around 10 min for ZnO (tube diameter: 4 mm, tube length: 20 m). This is a steady-state process in which each step, mixing and ageing, is well separated, leading to a better control and reproducibility. If needed, after the tubular reactor a heat-exchanger can be placed to cool down the reacting mixture to room temperature and stop the reaction. Then a separation unit or decanter allows the separation of the aqueous suspension and the immiscible fluid. The powder in the aqueous phase is collected, while dodecane is recycled in the process for environmental and cost considerations.

2.2 Micromixers

In precipitation systems, the role of the mixing is preponderant and can affect to a great extent the final material, its phase purity, morphology, size, and particle size distribution. Two micromixer designs were investigated in this study: The Y-configuration (Fig. 1b)) consists of two impinging sheets mixing

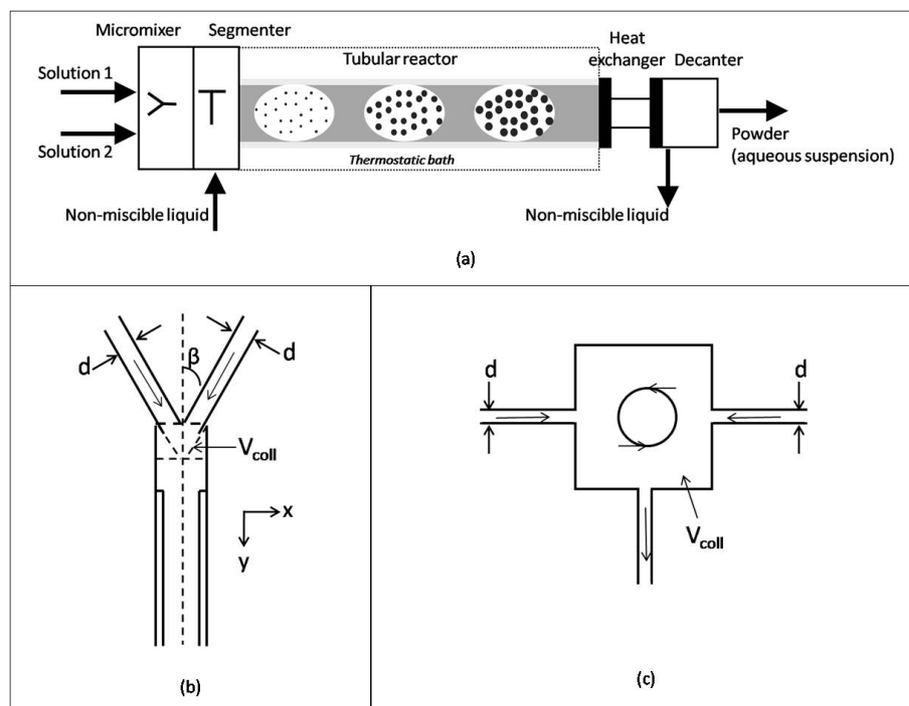


Figure 1. Schematic views: (a) SFTR, (b) Y-micromixer, (c) Roughton micromixer.

at the impingement zone V_{coll} with an angle of 30° which ensures an inelastic collision [29]; the Roughton configuration [30] is a T-mixer where the feed flows are displaced from a direct collision geometry to create an internal vortex in the mixing chamber V_{coll} (Fig. 1c)). For these two designs, the micromixing efficiency can be approximated and compared by calculating the Kolmogorov length [28, 31]. The specific energy dissipation rate ε is determined by the geometry of the impingement zone and the speed of the fluid.

$$\varepsilon = \frac{2P}{m} \quad (1)$$

Factor 2 results from the consideration of the two single sheets, which are assumed to be equal. P is the power lost during the collision, which is simply the kinetic energy divided by the time in the direction of the x -axis, and m is the mass over which P is lost (mass of liquid in V_{coll}). The Kolmogorov length is given by Eq. (2):

$$\lambda = \left(\frac{\nu^3}{\varepsilon}\right)^{1/4} \quad (2)$$

where λ is the Kolmogorov length, ν is the kinematic viscosity, and ε is the energy dissipation rate.

One can then estimate a mixing time, defined as follows:

$$t = \frac{(\lambda/2)^2}{D} \quad (3)$$

where λ is the Kolmogorov length and D is the species diffusivity. This last value can be calculated as:

$$D = \frac{k_B T}{6\pi\eta r} \quad (4)$$

where k_B is the Boltzmann constant, T is the temperature, η is the dynamic viscosity of the precipitation medium, and r is the ionic radius of the considered species. Taking these equations and the geometry into account, the Kolmogorov length and an approximate mixing time in the two micromixers were calculated [28, 31] and both geometries were used in the precipitation of ZnO.

2.3 Powder Characterization

The phase identification of the precipitates was made with powder X-ray diffraction (XRD, Philips X'Pert diffractometer, Cu- K_α radiation). XRD peak broadening was used to determine the size of the primary crystallite using the Scherrer equation (Eq. (5)). The instrumental broadening was determined

using alumina with a large crystal size ($> 1 \mu\text{m}$).

$$d_{\text{XRD}} = \frac{K\lambda_X}{\beta_{\text{Xp}} \cos(\theta)} \quad (5)$$

where K is equal to 0.9, λ_X is the X-ray wavelength, β_{Xp} is the integral breadth of the material, calculated using $\beta_{\text{Xp}} = \sqrt{(\beta^2 - \beta_{\text{alumina}}^2)}$, with the integral breadth β defining the ratio of the area and height of the diffraction peak.

The powder morphologies were analyzed by scanning electron microscopy (SEM, Philips XL 30 FEG microscope). The SEM samples were prepared by dispersing the powder in ethanol. One drop of the suspension was then deposited on an aluminum support and dried in air. TEM micrographs were obtained from a Philips CM 200 TEM operating at 200 kV and room temperature.

Brunauer-Emmett-Teller (BET)-specific surface areas S_{BET} (m^2g^{-1}) were estimated from N_2 adsorption isotherms (Micromeritics Gemini 2375). The size of the primary particles, d_{BET} (nm) [32], was calculated by assuming spherical monodisperse particles (Eq. (6)), where ρ is the density of the material.

$$d_{\text{BET}} = \frac{6000}{S_{\text{BET}} \rho} \quad (6)$$

Before BET measurements the samples were dried at 200°C in flowing nitrogen for 1 h.

The particle size distribution (PSD) was collected using a centrifugal method (CPS, Disc Centrifuge Model DC 24000, or XDC, X-Ray Disc Centrifuge Brookhaven Instruments). For this measurement, powders were suspended in a 0.1 wt-%

Table 1. Experimental data and geometric values used for the calculation of the micromixing times in the two micromixers for the ZnO precipitation system.

Y-Micromixer		Roughton Micromixer			
Internal diameter	d [m]	$2 \cdot 10^{-3}$	Internal diameter	d [m]	$3 \cdot 10^{-4}$
Flow rate	Q [$\text{m}^3 \text{s}^{-1}$]	$5 \cdot 10^{-6}$	Flow rate	Q [$\text{m}^3 \text{s}^{-1}$]	$1.67 \cdot 10^{-7}$
β	angle [°]	30	Flow speed	v [m s^{-1}]	2.36
Flow speed	v_x [m s^{-1}]	0.79	Volume of collision	V_{coll}	$2 \cdot 10^{-9}$
Dissipation rate	ε [$\text{m}^2 \text{s}^{-3}$]	109	Dissipation rate	ε [$\text{m}^2 \text{s}^{-3}$]	466
Kolmogorov length	λ [μm]	9.8	Kolmogorov length	λ [μm]	6.8
Mixing time	t [ms]	17	Mixing time	t [ms]	10

solution of PAA (pH ~ 10 , concentration 0.01 to 0.25 wt-%), and then treated 15 min with an ultrasonic horn. Median volume particle diameters (d_{v50}) and either the distribution standard deviation or span $(d_{v90} - d_{v10})/d_{v50}$ were used to characterize the widths of the distributions [32].

Thermogravimetric curves (TGA, Mettler TGA/DSC/TMA analyzer) were collected in the range from room temperature to 800 °C under flowing air at a heating rate of 10 °C \cdot min $^{-1}$.

3 Barium Titanate Precipitation: Process Intensification

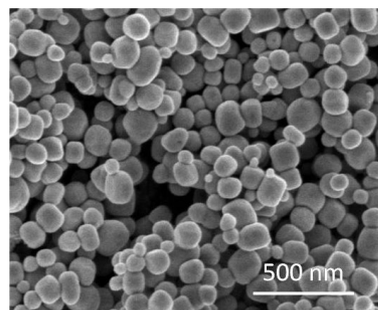
Barium titanate precipitation was carried out using barium chloride and titanium tetrachloride in one reactant stream and sodium hydroxide in excess as the second stream. The experimental details and particle growth mechanism have been well described elsewhere where the reactant concentrations and temperatures were initially investigated in small-batch reactors (50 g) and then transferred to the SFTR [25, 33]. The SFTR micromixer was of the Y-type. Nitrogen gas was used as a segmenting fluid and also protects against carbonate formation. The reaction was carried out at 95 °C and the residence time in the reactor was 10 min.

Regular ultrafine particles with an almost cubic morphology were obtained (see SEM picture in Fig. 2). The tetragonal phase was directly obtained and proved to be very pure with no traces of barium carbonate often found in even the best commercial powders. This powder presents a specific surface area of $S_{\text{BET}} = 10.5 \text{ m}^2 \text{ g}^{-1}$. During early investigations small batches of up to 50 g were produced in short production runs (1 or 2 h). To illustrate how the SFTR can intensify a process, increased batch sizes without loss of powder quality batches of 3 to 7 kg were produced by running the SFTR continuously for up to 30 h. The ultrafine nature and high-quality powder is illustrated with a high degree of reproducibility, as shown by the comparison in PSDs measured on seven different

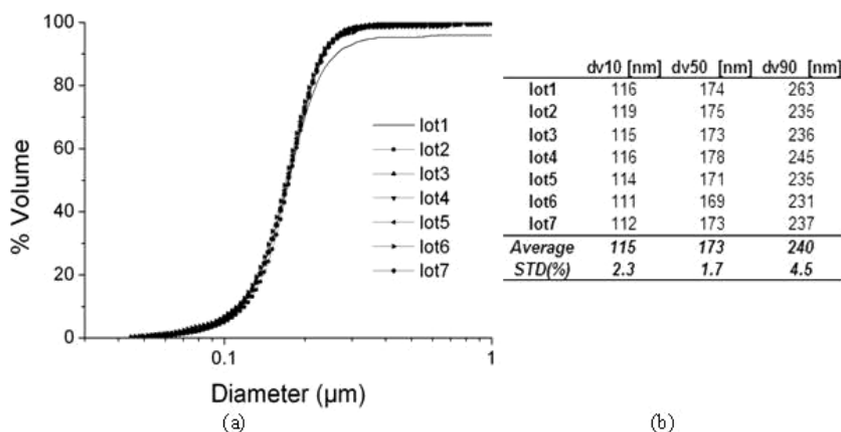
synthesis batches (Fig. 3a) where the PSDs are almost indistinguishable. The d_{v50} measured on seven different lots was 173 nm, with a very low standard deviation of 1.7%, illustrating the capacity of the SFTR in process intensification.

4 Calcium Carbonate Precipitation: Scale-Up/-Out Demonstration

Calcium carbonate precipitation has been studied intensively in our group for more than ten years [16, 20, 34–37]. The precipitation of calcite is

**Figure 2.** SEM picture of BaTiO₃ ultrafine particles obtained using the SFTR.

controlled by the addition of seeds stabilized by PAA. These seeds are added to a potassium carbonate solution (0.02 mol L $^{-1}$) as one stream, mixed with a calcium nitrate solution (0.02 mol L $^{-1}$) as another stream, using a Y-type micromixer. The above route has been optimized with a 20 mL batch reactor [35] allowing the production of high-surface-area nanostructured powders. Dodecane was used as the segmenting fluid. The reaction was carried out at ambient temperature and pressure, and the residence time was 10 min. To illustrate the stability of product quality as a function of reactor opera-

**Figure 3.** (a) Particle size distributions of seven different lots of BaTiO₃ powders obtained using the SFTR, measured by XDC, (b) percentile volume diameters measured, average, and standard deviations.

tion time, the SFTR was run continuously for 25 h. To illustrate the scale-out concept, a multiple-tube configuration with six tubes running in parallel was carried out for a 15 h period, which needed 100 L of reactants.

Typical calcium carbonate particles are shown in Fig. 4. They have a slightly elongated shape but were pure calcite within the detection limits of X-ray powder diffraction [34]. The powders have SSAs around $35 \text{ m}^2 \text{ g}^{-1}$, and the rough surface illustrates their nanostructured nature ($d_{\text{XRD}} = 67 \text{ nm}$). The PSDs for the minibatch, the single-tube SFTR as well as the multi-tube SFTR precipitates are shown in Fig. 5a) and are very similar. The span and d_{v50} for each type of powder is shown in Fig. 5b). This scale-up of 5000 going from the minibatch reactor (20 cm^3) to the 100 L of the multitube run without any loss of powder quality is remarkable and shows the power of the segmented flow reactor approach for the production and scale-out/up of high-quality powders.

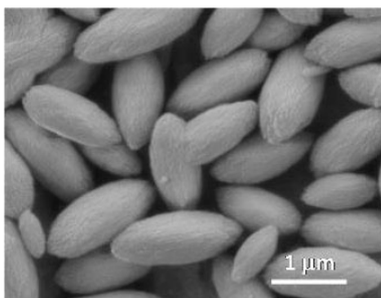


Figure 4. SEM picture of CaCO_3 nanostructured particles obtained using the SFTR.

5 Zinc Oxide Precipitation: Continuous Synthesis of Well-Dispersed Nanoparticles <100 nm

5.1 ZnO Precipitation Procedure

In this study, the synthesis of ZnO using the continuous SFTR process is presented for the first time. The precipitation of

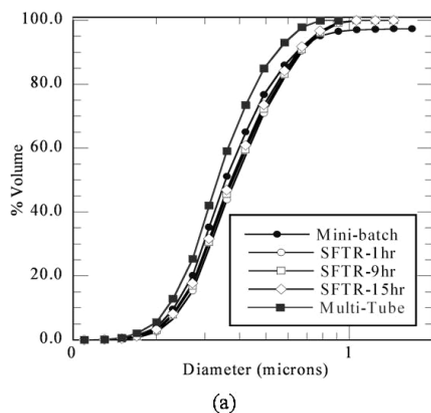
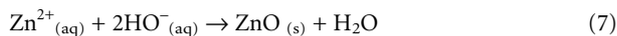


Figure 5. (a) Particle size distributions of different runs of CaCO_3 synthesis, (b) comparison of span and d_{v50} for different runs of CaCO_3 synthesis.

	Span	d_{v50} [μm]
Mini-batch (20 mL)	1.02	0.36
SFTR-25hr (25 L)	1.09	0.37
Multi-Tube (100 L)	1.06	0.39

ZnO results from the mixing of zinc nitrate and sodium hydroxide aqueous solutions (Eq. (7)).



The precipitation conditions were optimized in a previous study using a minibatch reactor [38]. The 0.10 M Zn^{2+} reaction solution was prepared by dissolving $\text{Zn}(\text{NO}_3)_2 \cdot 6\text{H}_2\text{O}$ in ultrapure water. The 0.11 M NaOH reaction solution was prepared by diluting a titrated solution of 1 M NaOH in ultrapure water. PAA (M_w 2000) at 0.05 wt-\% was added to the NaOH reaction solution for a better control of the precipitated powder characteristics. After reaction, the suspended powder was washed four times with ultrapure water, before being filtered and dried for 24 h at 70°C .

ZnO precipitation was conducted in a 400 cm^3 vessel, called batch reactor B, and in a 40 cm^3 vessel, called minibatch reactor MB, both placed in a thermostatic water bath at 90°C . The same experiments were also performed using the SFTR, called S (see Tab. 2). Dodecane was used as the segmenting fluid. The reaction was carried out at 90°C and the residence time was 10 min. All synthesis conditions are summarized in Tab. 2.

Table 2. Description of the process used (micromixer and reactor) for the different precipitated ZnO samples.

Sample	Micromixer	Reactor	Productivity
B	none	Batch	400 cm^3 2 g/batch
B-Y	Y	Batch	400 cm^3 2 g/batch
B-V	Roughton (Vortex)	Batch	400 cm^3 2 g/batch
MB-Y	Y	Minibatch	40 cm^3 0.2 g/batch
MB-V	Roughton	Minibatch	40 cm^3 0.2 g/batch
S-V	Roughton	SFTR	0.2 cm^3 4 g/h

5.2 Influence of the Process

Whatever the process (reactor and micromixer), pure and crystalline ZnO nanoparticles were obtained by precipitating zinc nitrate and sodium hydroxide in the presence of PAA at 90°C , as shown with the diffractogram of the sample S-V in Fig. 6. This diffractogram matches the ZnO pattern of wurtzite (ICDD 075-0576). Crystallite sizes were calculated using the [100] reflection and are given in Tab. 3. All powders present nanodomains in the same range, varying between 24 and 30 nm.

The TG curve of S-V is presented in Fig. 7. The first weight loss between 30 and 200°C was attributed to the removal of physically adsorbed water. The second weight loss between 200 and 400°C is the consequence of two phenomena: the decomposition of chemically bound hydroxyl groups and

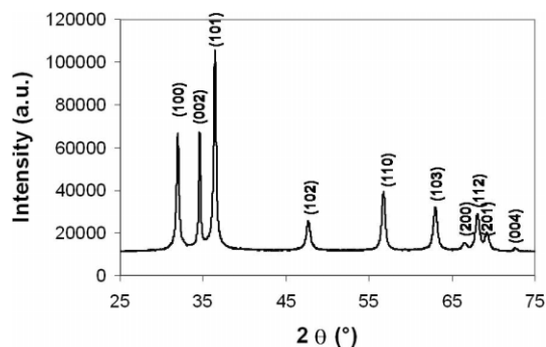


Figure 6. XRD pattern of ZnO synthesized with the SFTR (sample S-V).

the removal of the organic compound PAA that was adsorbed at the surface. The total weight loss measured for the different samples is in the same range, between 12.7 and 14.5%, as shown in Tab. 3.

There is, however, a significant influence of both micromixer and type of reactor on the specific surface area and on the PSD. This is demonstrated by the three characteristic sizes of the PSD (d_{v10} , d_{v50} , d_{v90}) given in Tab. 3 and the cumulative distributions compared in Fig. 8.

The span is calculated from Eq. (8), giving an indication of the width of the PSD:

$$\text{span} = \frac{d_{v90} - d_{v10}}{d_{v50}} \quad (8)$$

If no micromixer is used (sample B), the resulting particles from the batch reactor have a span of 1.8 with sizes ranging from 100 to 700 nm and a specific surface area of $45 \text{ m}^2 \text{ g}^{-1}$. The lack of control on the mixing and thus the initial supersaturation presumably leads to such heterogeneity in the final powder.

Using a micromixer with the batch reactor (Y- or Roughton configuration) decreases the particle size and a span below 1.0 is attained. With the Y-micromixer, the d_{v50} of the particles is around 100 nm (B-Y: 109 nm), whereas with the Roughton micromixer the d_{v50} of the particles is below 100 nm (B-V: 82 nm). The Roughton configuration allows a better mixing as shown by the calculation of the Kolmogorov length and the

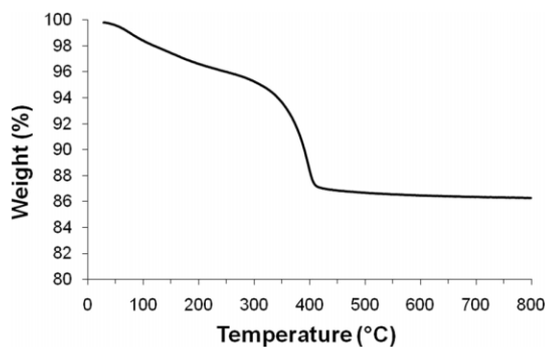


Figure 7. TG curve obtained for ZnO synthesized with the SFTR (sample S-V). Heating rate $10^\circ\text{C min}^{-1}$ in flowing air (30 mL min^{-1}).

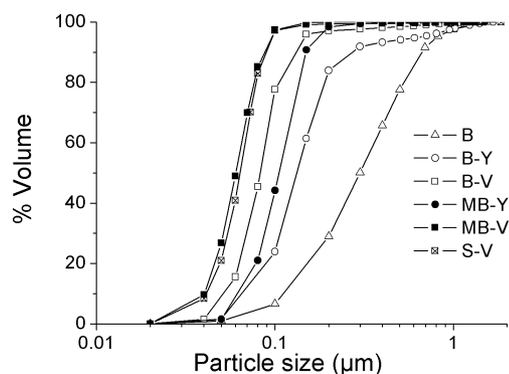


Figure 8. Particle size distributions of ZnO powders measured by centrifugal sedimentation (CPS).

estimated mixing time (Tab. 2: 17 ms with the Y-configuration and 10 ms with the Roughton configuration). Consequently, with a faster mixing, it can be assumed that the local supersaturation will be higher, leading to smaller particle sizes. These trends with the two mixer geometries were also observed for the minibatch and SFTR reactors.

The reactor volume also has an influence on the final PSD. By using the same micromixer (Y or Roughton), the d_{v50} of the powder is higher and the specific surface area is lower in a large-volume batch reactor. The particles B-Y and B-V also

Table 3. Results of S_{BET} , d_{BET} , d_{XRD} , particle size distribution (PSD), and weight loss measured by TGA of ZnO powders obtained through different processes.

Sample	S_{BET} [$\text{m}^2 \text{ g}^{-1}$]	d_{BET} [nm]	d_{XRD} [nm]	PSD			Span ($d_{v90} - d_{v10}$)/ d_{v50}	Weight loss at 800°C [%]
				d_{v10} [nm]	d_{v50} [nm]	d_{v90} [nm]		
B	45	24	27	119	307	674	1.8	13.7
B-Y	57	19	30	67	109	178	1.0	12.7
B-V	64	17	28	56	82	116	0.7	13.7
MB-Y	75	14	24	65	104	147	0.8	14.5
MB-V	78	14	23	40	60	85	0.7	
S-V	68	16	24	42	64	86	0.7	13.3

look bigger and less homogeneous in size in the SEM pictures (Fig. 5). So, by increasing the reacting volume from only 40 to 400 cm³, the homogeneity of the powder and the mean particle size were significantly changed. This is typical of scale-up in processes, which are mainly attributed to poor homogeneity inside the reactor volume, leading to concentration variations, accumulation zones, and also to heat transfer discrepancies, which can affect the precipitation kinetics (nucleation and growth) and hence the final powder quality.

With a smaller volume of reaction a better homogeneity is ensured in the minibatch and the SFTR reactors and a narrower PSD is obtained. SEM and TEM micrographs in Fig. 9 give a qualitative view of the nanometric sizes of these particles and their narrow size distributions, confirmed with the PSD measurement by the CPS centrifugal method (Fig. 4). These results show that the particles are polycrystalline with primary particles from the XRD line broadening being around 23 to 30 nm and the redispersible particles being 60 to 100 nm. Finally, one can observe in Fig. 8 that the PSDs of MB-V and S-V are almost superimposed. This illustrates the possibility to optimize the precipitation parameters at the lab scale (a few mg/batch) and transfer directly these parameters to the SFTR

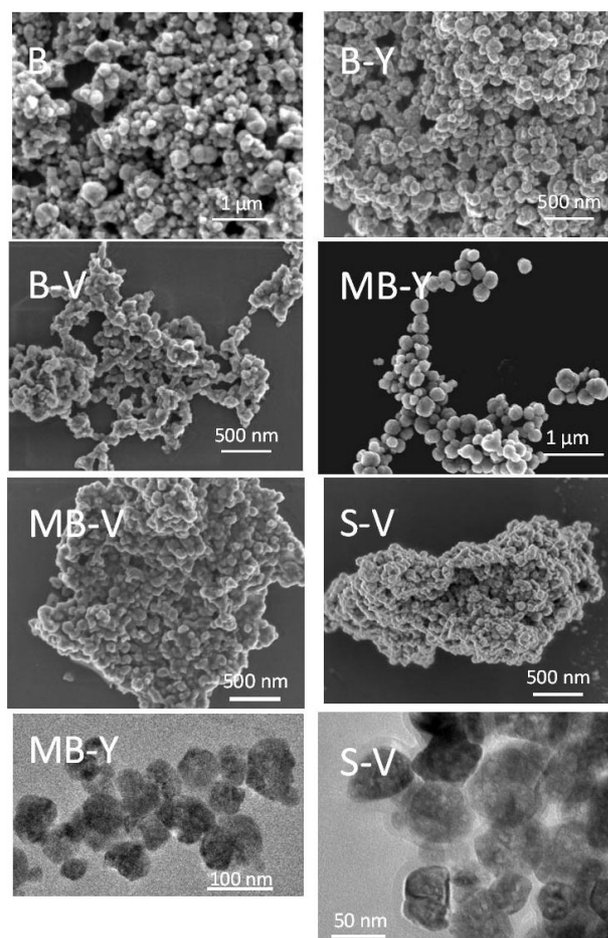


Figure 9. SEM and TEM (MB-Y and S-V) micrographs of ZnO powders obtained from different processes.

for a pilot production scale (g to kg/h, depending on the number of tubes working in parallel and the solubility of the product powder), maintaining the high-quality characteristics of the final powder.

5.3 Robustness of the SFTR

The SFTR offers the benefit of a continuous process rather than stop-start batch reactors. Moreover, the SFTR allows the production of higher-quality materials due to the segmenting fluid which produces homogeneous microreactors. To demonstrate that the powder quality for nanosized powders could be maintained over a realistic working time, ZnO precipitation was run for 12 h. Some aliquots of the aqueous suspension were collected at different times for the measurement of the particle size distribution by CPS. The results are shown in Fig. 10. The PSD remains very narrow over the considered period of time with an excellent stability ($d_{v50} = 64.6 \pm 2.1$ nm). The particle morphology and phase purity as well as the specific surface area were also constant over the same period (not shown). The SFTR produced 50 g of powder with an excellent control of its properties. Such quantity is a minimum for further investigations on the ceramic processing and sintering tests, which are currently under investigation to determine their applicability for real-ceramic applications.

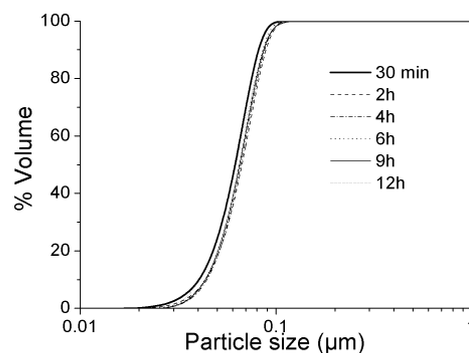


Figure 10. Particle size distributions of ZnO powders during 12 h of continuous precipitation using the SFTR, measured by centrifugal sedimentation (CPS).

It was also interesting to consider the effect of the drying step on the agglomeration of the powder, as most applications prefer the use of dried powders as raw materials. The PSD was measured on an aliquot of the aqueous phase just after collection from the reactor and on resuspended powder previously dried by two different processes: classical drying in an oven at 70 °C during 24 h and freeze-drying. The results are shown in Fig. 11. This figure clearly shows that while freeze-drying allows the redispersion of the powder to give the same PSD as before drying, the classical oven drying leads to the formation of a small percentage of agglomerates (~8 % of the volume). This is a very important aspect for the technological breakthrough of nanosized powders. All too often PSDs are given from TEM images with the degree of agglomeration being qualitatively assessed as low or softly agglomerated. Here, we

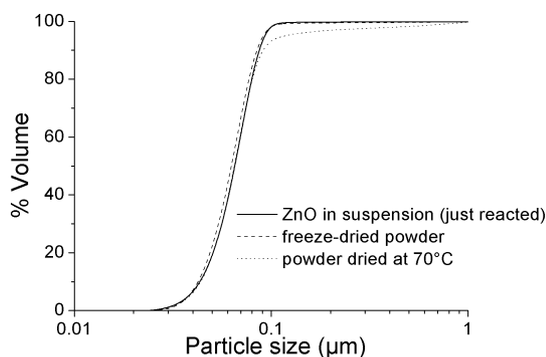


Figure 11. Particle size distributions of ZnO powders synthesized with the SFTR, in suspension after the reaction, after a classical drying of 24 h at 70 °C, and after freeze-drying, measured by centrifugal sedimentation (CPS).

demonstrate the capacity of the SFTR to produce powders which can be redispersed after the drying step, thus allowing for more economical transport.

6 Conclusions

Using a plug flow segmented fluid microreactor, the SFTR, successful scale-out and process intensification for ultrafine CaCO_3 , BaTiO_3 , and nanosized ZnO powders were investigated. The chemistry and operational conditions were optimized using minibatch systems (20–100 mL) and the powder quality could be maintained throughout the scale-up or process intensification steps. This is of key importance in moving from lab scale to pilot and/or production scale, reducing time and cost in the development and testing of new products.

The process intensification was demonstrated by producing 3–7 kg batches of tetragonal BaTiO_3 powders ($d_{v50} = 173 \pm 3$ nm) and with excellent batch-to-batch reproducibility over seven to ten batches. The scale-out or numbering-up capacity of the SFTR was demonstrated for a nanostructured CaCO_3 ($d_{v50} \approx 500$ nm) by scaling-out from one to six tubular reactors run in parallel representing a scale-out/up ratio of 5000 without loss of powder quality. The SFTR's potential for nanosized powders was then demonstrated by producing 50 g of pure and polycrystalline ZnO nanoparticles. This nanopowder ($d_{v50} = 65 \pm 2$ nm) shows a very narrow PSD due to a perfect control of the precipitation conditions by using a Roughton micromixer and the reduced reaction volume after segmentation (~ 0.2 cm³). An excellent reproducibility was demonstrated and in particular the very narrow PSDs were conserved over the whole 12 h run. Dried powders could be perfectly redispersed when using the freeze-drying method.

The ability of the SFTR to scale out the powder production from very low optimization volumes (40 cm³) at the laboratory scale without changes in powder quality has been clearly demonstrated. For the different chemical systems investigated the pilot quantities of powder produced allow their assessment for ceramic or other applications. The SFTR is a powerful tool for the production of powders and nanocrystals with advanced

properties and accompanies the development and optimization of new materials in the growing market of nanotechnologies as well as high added-value powders in general.

Acknowledgements

The authors would like to thank the Swiss cost office, COST Action 539 and the Swiss Federal Office for Education and Science (CTI-TOPNANO 21) for support.

The authors have declared no conflict of interest.

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