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Combined hydrothermal and mechanochemical control of structural modifications of zirconium dioxide for catalytic applications

In this work the influence of hydrothermal treatment and subsequent mechanochemical treatment on the porous and crystalline structure of precipitated zirconium dioxide was studied. Such new synthetic protocol enabled to obtain purephase materials with the stable defected structure for catalytic applications as revealed by as indicated by the results obtained using XRD and UV-vis spectroscopy.

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# Combined hydrothermal and mechanochemical control of structural modifications of zirconium dioxide for catalytic applications†

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The influence of hydrothermal treatment (HTT) and subsequent mechanochemical treatment (MChT, milling) on the porous and crystalline structure of precipitated zirconium dioxide was studied. It has been established that HTT at 300 °C promotes the transformation of amorphous ZrO2 into a pure monoclinic phase, as well as the formation of a uniform mesoporous structure which has higher thermal stability. Soft dry milling (300 rpm, 0.5-1 h) of hydrothermally modified monoclinic  $ZrO_2$  causes the introduction of defects into its structure without a noticeable change in the phase composition. The presence of defects is confirmed directly using UV-vis spectra and indirectly by the manifestation of photocatalytic activity of milled samples under visible irradiation. Importantly, it is found that after calcination of milled samples at 500 °C a high fraction of defects remains preserved which opens up the potential of using zirconium dioxide modified in this way as a catalyst or catalytic support with added specific properties offered by defects.

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## Introduction

Zirconium dioxide (ZrO<sub>2</sub>) is a versatile material thanks to the tunability of its physicochemical characteristics. ZrO2 has been receiving increasing attention both as a heterogeneous catalyst and catalyst support since it has adjustable phase composition, redox and acid-base properties, and defects, notably oxygen vacancies, which affect its own catalytic performance and/or of the catalytic moieties dispersed on it.1-5 In particular, ZrO2 has been used in a wide spectrum of industrially relevant processes such as water-gas shift, selective reduction of NO2, or biodiesel production. ZrO<sub>2</sub> combined with active metals or metal oxides dispersed on its surface is used in hydrogenation<sup>6,7</sup> or ketonization<sup>7</sup> reactions, as well as in CO<sub>2</sub> and CH<sub>4</sub> conversion<sup>1,8</sup> via CO<sub>2</sub> methanation or dry methane reforming (DMR).9 Its role is still not fully understood, but theory calculations indicate that oxygen vacancies significantly lower the adsorption energy of

One of the most important properties is the phase composition of ZrO2 along its porous vs. crystalline structure, both highly stable in a wide range of applied temperatures. Coking resistance and hydrothermal stability are other major challenges for catalytic materials. 1,4,9 Bulk zirconia is hardly reducible, as demonstrated by the high oxygen vacancy formation energies observed by EPR;14 however, reducibility can be altered by lowering ZrO<sub>2</sub> particle size. Under-coordination and nanostructuring can strongly affect the properties of ZrO2 making non reducible bulk zirconia reducible at the nanoscale, impacting its role as an oxide support as well. 1,4,9 Thus, tailored synthesis protocols offer ways of tuning the structure, morphology and chemical nature of ZrO2 for catalytic applications. 12,15-21

The influence of ZrO<sub>2</sub> precipitation conditions on its porosity and specific surface area value is relatively well described in the literature.17,22-24 However, the disadvantages of precipitated ZrO<sub>2</sub> are the significant content of micropores and an underdeveloped mesoporous structure as well as an amorphous state.25,26 Also, the rapid sintering of micropores and a dramatic decrease in the specific surface area with increasing temperature have been observed.27-30 Post-synthetic modification is necessary to transform amorphous ZrO2 into a crystalline state. Calcination in air, which is most often used for this purpose, results in the formation of an unstable tetragonal phase at low temperature. The complete transformation of tetragonal ZrO<sub>2</sub> into a monoclinic phase does not occur even at 500 °C31,32 and is

CO<sub>2</sub> and promote its dissociation<sup>10,11</sup> with the added benefit of suppressing coke formation. 12,13

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also accompanied by a significant decrease in the specific surface area.<sup>33</sup> It appears that it is more advantageous to use hydrothermal treatment (HTT) as a post-synthetic modification of precipitated ZrO<sub>2</sub> by either processing the wet gel or the dried xerogel.<sup>34</sup> Hydrothermal processes are distinguished as a technique for the synthesis and modification of the structure of porous oxides and nanomaterials<sup>31–34</sup> and excel as simple and cost-effective alternative methods.<sup>35–44</sup> A specific feature of wet gels HTT is the possibility of obtaining materials that simultaneously have high specific surface area and accessible pore size as was shown for zirconium oxide.<sup>26,45</sup> Such modified materials are moreover resistant to sintering,<sup>46</sup> as a co-precipitated composition, *e.g.*, CeO<sub>2</sub>–ZrO<sub>2</sub>.<sup>47,48</sup>

Ball milling is considered another environmentally friendly and effective method used for the preparation of oxide materials, including catalysts<sup>46,49–51</sup> with an added important feature that nanostructured milled oxides contain various defects (e.g., oxygen vacancies). The latter can act as active sites in different catalytic processes,<sup>52–55</sup> e.g., in test photocatalytic reactions of safranin degradation.<sup>27</sup> It should be noted that studies on ZrO<sub>2</sub> milling are not very numerous, do not have a systematic character and are not focused on obtaining catalysts.<sup>56–60</sup> As a rule, they are devoted to the milling of monoclinic ZrO<sub>2</sub> with its transformation into unstable cubic and tetragonal phases. The effect of milling of ZrO<sub>2</sub> on its porous and surface structure has not been studied, to the best of our knowledge.

The purpose of this work is to fill the gap in the synthesis of ZrO<sub>2</sub> structures with controllable and thermostable porosity,

crystallinity, specific surface area, surface chemistry, redox properties and defects by a unique combination of hydrothermal and mechanochemical routes and explore the properties of this new class of materials.

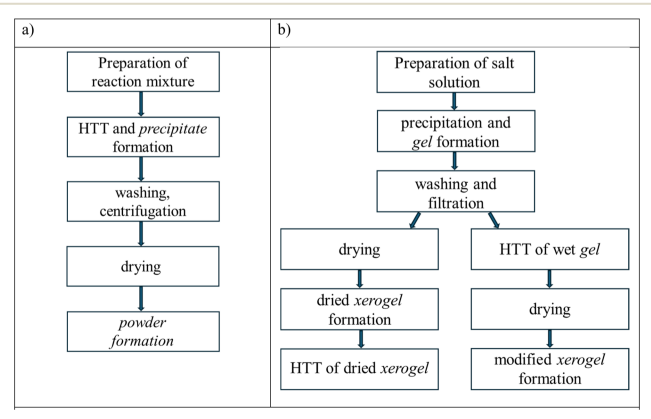
## 2. Experimental

#### 2.1 Precipitation

An 11.5 M ammonium hydroxide solution (28%, Sigma-Aldrich, analytical grade) was gradually added to the 0.35 M solution of zirconium(IV) oxynitrate dihydrate (99%, Sigma-Aldrich, analytical grade) under intensive stirring (500 rpm) while monitoring the pH value. During this process, a white gelatinous precipitate was formed. The pH values at the end of the precipitation were chosen to be at the edges of the interval favorable for the formation of the monoclinic phase, 9.5 and 7.0, respectively. The resulting gel was aged in the mother solution for 24 hours at room temperature. Afterward, it was separated by decantation and then washed on a filter to the neutral pH value with distilled water and compacted to a water content of approximately 85% resulting in a wet gel. Part of the obtained wet gel was dried for 48 hours at room temperature to form a dried xerogel.

## 2.2 Modification procedures

The obtained  $ZrO_2$  in the form of a wet gel and a dried xerogel was subsequently subjected to hydrothermal treatment (HTT) at 260–320 °C and equilibrium vapor pressure for 5–7 hours. HTT



Scheme 1 Preparation of ZrO<sub>2</sub> using procedures (a) previously described and (b) used in this work.

Table 1 List of samples with their method of preparation. The symbols H, M and T denote hydrothermal treatment, milling and thermal treatment, X and G - xerogel and gel, and the numbers after them - the conditions of these treatments in terms of time and temperature that were used for the designation

Sample designation	Method of preparation	
N1-9.5	Initial pH 9.5	
N2-9.5_HX	HTT xerogel 260 °C 5 h	
N3-9.5_HXM_300	HTT xerogel 260 °C + MChT 300 rpm 0.5h BPR 10	
N4-9.5_HXM_450	HTT xerogel 260 °C + MChT 450 rpm 0.5h BPR 10	
N5-9.5_HX	HTT xerogel 320 °C 5 h	
N6-9.5_HXM_300	HTT xerogel 320 °C 5 h + MChT 300 rpm 0.5h BPR 10	
N7-9.5_HG	HTT gel 260 °C 5 h	
N8-9.5_HGM_300	HTT gel 260 °C 5 h + MChT 300 rpm 0.5h BPR 10	
N9-7	Initial pH 7	
N10-7_HX	HTT xerogel 260 °C 5 h	
N11-7_HXM_300_1h_10	HTT xerogel 260 °C 5 h + MChT 300 rpm 1h BPR 10	
N12-7_HXM_300_0.5h_13	HTT xerogel 260 °C 5 h + MChT 300 rpm 0.5 h BPR 13	
N13-7_HXM_300_1h_13	HTT xerogel 260 °C 5 h + MChT 300 rpm 1h BPR 13	
N14-7_T	Initial pH 7 +TT 500 °C	
N15-7_HG	HTT gel 300 °C 5 h	
N16-7_HG_T	HTT gel 300 °C 5 h + TT 500 °C	
N17-7_H7hG	HTT gel 300 °C 7 h	
N18-7_H7hG_T	HTT gel 300 °C 7 h + TT 500 °C	
N19-7_H7hGM_500_0.5_25	HTT gel 300 °C 7 h + MChT 500 rpm 0.5h BPR 25	
N20-7_H7hGM_500_0.5_25_T	HTT gel 300 °C 7 h + MChT 500 rpm 0.5h BPR 25 + TT 500 °C	
N21-7_ H7hGM_400_0.5_10	HTT gel 300 °C 7h + MChT 400 rpm 0.5 h BPR 10	

was carried out in Teflon-lined steel autoclaves with a volume of 45 mL. In both cases, the wet gel and dried xerogel were loaded in a quartz tube that was placed in an autoclave. 15 mL of water was added to the bottom of the autoclave in the processing of the dried xerogel so that HTT was carried out in the vapor phase.34 Hydrothermally treated samples were dried at 100 °C for 6 hours. The difference between the HTT option proposed in this paper and the previously described treatments 40-45,47-49 can be seen in Scheme 1 presented below.

In the first case (Scheme 1a), HTT of the initial reaction mixture is used, resulting in the formation of powders. Their particles can be porous but in general, they are loosely dispersed materials.<sup>61</sup> In the second case (Scheme 1b), washed wet gels or the dried xerogel can be subjected to HTT, the products of which are consolidated or porous materials.61

Hydrothermally treated dried xerogels were subjected to mechanochemical treatment (MChT) in air (dry milling) at 300-500 rpm for 30-60 min. MChT was performed using a planetary ball mill Pulverisette-7, premium line (Fritsch Gmbh) with a vessel of zirconium dioxide. 134 Zirconium dioxide balls with a diameter of 5 mm and a total mass of 91.5 g were used as working bodies. The use of a zirconia ball-milling assembly was chosen to reduce the introduction of contaminants into the product.<sup>62</sup> The ratio of the mass of the balls to the mass of the sample (ball-to-powder, BPR) was 10, 13, and 25. The prepared samples, their designations and synthesis conditions are listed in Table 1.

#### Physicochemical characterization

The porous structure of ZrO2 samples was characterized using the adsorption-structural method. The isotherms of nitrogen

adsorption-desorption were obtained using an automatic gas adsorption analyzer ASAP 2405N (Micromeritics Instrument Corp.) after outgassing the samples at 150 °C for 20 h. The specific surface area S, volume of mesopores  $V_{\rm me}$ , and volume of micropores  $V_{mi}$  were calculated from these isotherms using the Brunauer-Emmett-Teller (BET), Barrett-Joyner-Halenda (BJH), and t-methods, respectively. Sorption pore volume V was determined at a relative pressure of nitrogen  $p/p_0$  close to 1. The curves of pore size distribution (PSD) were then plotted using the desorption branches of isotherms.

X-ray phase analysis and structural and microstructural characterization of the samples were performed by using a DRON-3M diffractometer (CuK $\alpha$  radiation,  $\lambda = 0.154$  nm), fullprofile Rietveld refinement was performed by using WinCSD software<sup>63</sup> to obtain lattice parameters of the monoclinic ZrO<sub>2</sub> structure, and microstructural parameters of the samples (average grain size D and microstrains  $\langle \varepsilon \rangle$ ) were derived by full profile Rietveld refinement. The Garvie-Nicholson method allows determination of the molar content of the monoclinic phase  $X_{\rm m}$  according to the following equation:<sup>64</sup>

$$X_{\rm m} = [I_{\rm m}(111) + I_{\rm m}(-111)]/[I_{\rm m}(111) + I_{\rm m}(-111) + I_{\rm t}(101)],$$

where  $I_{\rm m}(111)$  and  $I_{\rm m}(-111)$  denote the intensity of peaks of the monoclinic phase at 28.2 and 31.3°, respectively, and  $I_t(101)$ denotes the intensity of the peak of the tetragonal phase at 30.4°.

The volumetric fractions of the monoclinic phase  $V_{\rm m}$  were calculated using the so-called Toraya method based on only some X-ray diffraction (XRD) peak intensity values:56,57

$$V_{\rm m} = 1.311 X_{\rm m} / (1 + 0.311 X_{\rm m})$$

Accordingly, the content of the tetragonal phase  $X_t(V_t)$  was calculated as follows:

$$X_{\mathsf{t}}(V_{\mathsf{t}}) = 1 - X_{\mathsf{m}}(V_{\mathsf{m}})$$

The size of the crystallites ( $D_t$  and  $D_m$ ) was calculated according to Scherrer's formula using the most intense reflexes of each phase.

The curves of differential thermal analysis (DTA) and thermogravimetry (TG) were recorded using the Derivatograph-C apparatus (F. Paulik, J. Paulik, L. Erdey) in the temperature range 20-1000 °C at a heating rate of 10 °C min<sup>-1</sup>. The initial mass of the sample was about 200 mg.

The UV-visible diffuse reflectance spectra were recorded with a UV-vis/DRS Lambda 35 spectrometer (PerkinElmer Instruments) in the range of 200-600 nm. MgO was the reference sample. The absorption edge  $\lambda$  was determined based on the spectra plotted in the Kubelka-Munk equation coordination. The bandgap  $E_{\rm g}$  was determined using a Tauc plot.<sup>58</sup>

The morphology of the ZrO2 powders was characterized with a Hitachi S4800 scanning electron microscope (SEM) equipped with a Nanotrace electron diffraction EDX detector (Thermo Electron).

To study the lattice vibrations of ZrO2, diffuse reflectance measurements were conducted in the Far-Infrared (Far-IR) region (30-600 cm<sup>-1</sup>), at the Canadian Light Source (CLS) farinfrared beamline using a Bruker IFS125HR spectrometer equipped with a T222 Mylar beamsplitter. The infrared beam was directed out of the sample compartment into a Pike Instruments diffuse IR accessory and then to a OMC superconducting niobium Transition-Edge Sensor (TES) bolometer. A synchrotron was used as a source because of its high flux of photons below 200 cm<sup>-1</sup>. The ZrO<sub>2</sub> samples were packed into ceramic sample cups and placed into the environmental chamber of the DiffusIR accessory. The chamber with an internal volume of 63 cm<sup>3</sup> was equipped with an Si window. Argon gas was continuously flowing through the chamber at a rate of 200 mL min<sup>-1</sup>, regulated by a mass flow controller (Brooks SLA5850). All spectra were collected with 4 cm<sup>-1</sup> resolution and averaged over 5 sets of measurements each averaging over 2048 scans. Data processing was done using the OPUS software package to obtain the reflectance spectra, which were produced by dividing the ZrO<sub>2</sub> spectra by the spectra of polyethylene powder. The spectra of polyethylene powder were collected directly before and after the collection of ZrO2 spectra and using the same conditions and served as a background.

Selected samples with a pure monoclinic phase were tested in the photocatalytic degradation of safranin T (ST), a dye which is commonly used as a model compound due to its high stability.65 Along with the UV-vis spectra, the reactivity is considered an indirect measure for assessing the formation of defects in the structure of the zirconium dioxide at different stages of its preparation (hydrothermal treatment, milling, and post-calcination). Testing of pure monoclinic ZrO2 samples under visible irradiation was carried out in an aqueous medium using  $1 \times 10^{-5}$  mol dm<sup>-3</sup> solution of ST<sup>59</sup> with a Philips LED

cool daylight (100 W) lamp as an irradiation source. A Shimadzu spectrophotometer UV-2450 was used for the analysis of changes in the concentration of dye during the photocatalytic degradation (band at 520 nm). The optimal dose of catalysts, 1 g dm<sup>-3</sup>, was used as it was established in other studies.<sup>59</sup> Prior to irradiation, the ZrO<sub>2</sub> dispersed in the ST-containing solution was stirred until an adsorption-desorption equilibrium was established for 1 h; conditions were applied as those in the testing of other oxide photocatalysts.27,65

#### 3. Results and discussion

## Effect of pH during precipitation

## 3.1.1 Structural analysis of the samples prepared at pH 9.5.

X-ray diffraction was used to determine the crystal structure of the materials. The initial sample precipitated at pH 9.5 is amorphous (sample N1-9.5, diffraction pattern not shown). For the xerogel sample after HTT (N2-9.5\_HX), the reflections of the tetragonal phase are very intense (Fig. 1). The content of the particular phase determined based on the diffractograms showed that the volumetric percentage of monoclinic ZrO<sub>2</sub> is 38% for this modified xerogel, see Table 2 (the indicated content refers to the crystalline part of zirconium dioxide). As mentioned above, the transition from the tetragonal to the monoclinic phase occurs already at low temperatures in the case of HTT reaction mixtures of zirconium salt with a precipitant. This is due to the mineralizing effect of anions in the case of direct hydrothermal synthesis of zirconium dioxide. 35,41 On the other hand, these anions can interfere with the preparation of pure ZrO<sub>2</sub>.

Simultaneously, dry milling of the ZrO2 samples, which had been hydrothermally processed as xerogels, gave a very promising result in the formation of the monoclinic phase. For the series of samples with a pH of 9.5 the initial material HTT promotes the formation of crystalline ZrO2 in all cases, which is represented by a mixture of tetragonal and monoclinic phases;

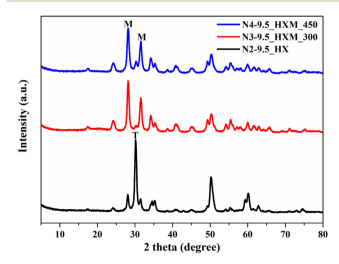


Fig. 1 XRD patterns for ZrO<sub>2</sub>, precipitated at pH 9.5, after HTT and post-milling at 300 and 450 rpm (T - tetragonal phase, peak from plane [101]; M - monoclinic phase, peaks from planes [111] and [-111]).

Table 2 Volume content of the monoclinic and tetragonal phases and their crystallite size, calculated using the Scherrer formula

-
$D_{\rm m}$ , nm
_
14.2
13.8
12.7

however, small content of the amorphous material cannot be excluded. Significant transformation of the tetragonal phase into the monoclinic one (Fig. 1) reaching 93% was achieved already with soft and short ball milling of the xerogel in sample N3-9.5\_HXM\_300. An increase in the intensity of milling (sample N4-9.5\_HXM\_450) does not promote further transformation (Table 2). A similar effect of reflexes' intensification can be observed for the gel and its milled counterpart (ESI materials, Fig. S1 and Table S1,† samples N7-9.5\_HG and N8-9.5\_HGM\_300). At the same time, increasing the temperature of the HTT to 320 °C does not give an additional effect (ESI materials, Table S1,† samples N5-9.5\_HX and N6-9.5\_HXM\_300).

It should be noted that in earlier studies, the milling of the tetragonal phase was scarcely studied. 60,66,67 It was suggested that this transformation (tetragonal into monoclinic) becomes possible in the presence of water vapor.68,69 Indeed, hydrothermal samples, which are subjected to subsequent milling, contain a lot of adsorbed water, as is shown below based on thermogravimetric measurements. Also, there is a certain critical crystallite size (~46 nm) above which the transformation of the tetragonal phase into the monoclinic phase may occur.70 However, this mechanism refers to the transformation that occurs during calcination and it is hardly applicable for milling. In this work, the effect is rather attributed to the presence of water and ammonia impurities, which is further discussed in Subsection 3.5.

The size of the crystallites was calculated according to Scherrer's formula using the half-width (FWHM) of the most intense peaks of both phases ( $D_t$  and  $D_m$ ). As can be seen, the crystallites of both monoclinic and tetragonal ZrO2 are reduced after milling (Table 2).

To gather further information about changes in the ZrO<sub>2</sub> structure after milling, FIR spectra were acquired for samples N2-9.5\_HX, N3-9.5\_HXM\_300, and N4-9.5\_HXM\_450, which have different contents of monoclinic phase and defects. Generally, in a crystalline structure, molecules are arranged in a repeating, three-dimensional pattern, leading to well-defined and sharp peaks in the IR spectrum. Commercial ZrO2 was used as a reference material. In general, spectra confirm the results of XRD analysis in terms of the tetragonal and monoclinic mixture of phases. Also, the signal in the lattice vibration range which is characteristic of the monoclinic phase becomes more pronounced with milling (Fig. 2).

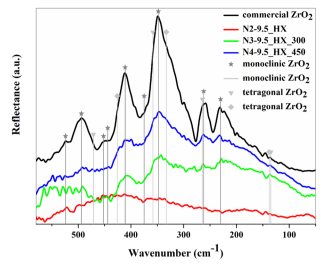


Fig. 2 Far IR spectra of samples N2-9.5\_HX, N3-9.5\_HXM\_300, N4-9.5\_HXM\_450 and commercial ZrO2 (reference). The spectra were obtained at 20 °C with an Si window, and polyethylene was used as a background. Marked  $\star$  are the bands of monoclinic ZrO<sub>2</sub> (ref. 71) and ▼ denote bands of tetragonal ZrO<sub>2</sub>,72 and ◆ bands of tetragonal

The spectrum of the reference (commercial monoclinic ZrO<sub>2</sub>) shows all 9 peaks characteristic of the ZrO2 structure observed by Maczka et al.,73 El Boutaybi et al.74 and previously by Hirata et al.75 with doublets at 494 cm<sup>-1</sup> and 524 cm<sup>-1</sup>. The doublets at 452 and 444 cm<sup>-1</sup> and the shoulder at 375 cm<sup>-1</sup> are less defined for commercial ZrO2 compared to the ones presented in the cited work<sup>73</sup> where highly crystalline ZrO<sub>2</sub>, calcined at 1200 °C, was studied. The band at 348 cm<sup>-1</sup> is the most prominent feature of monoclinic ZrO2. The spectrum of the reference sample also contains intense bands at 411, 262 and 230 cm<sup>-1</sup>.

For the selected synthesized samples with different tetragonal/monoclinic phase ratios, starting with N2-9.5\_HX (the as-prepared hydrothermal material), the features of the spectrum are difficult to distinguish. One possible cause of the loss of spectral details may be a strong scattering on this sample with fine crystallite size. According to the XRD results, the crystal structure of sample N2-9.5\_HX is characterized as a mixture of tetragonal and monoclinic phases, reaching about 62% of the former and the rest of the latter (Table 2). The spectrum of this hydrothermal sample is poorly resolved, possibly due to the presence of an amorphous component, as well as adsorbed water and surface hydroxyl groups. Also, the more random distribution of molecular vibrations, leading to a less sharp and often flat IR spectrum,76 may indicate the lack of a well-defined lattice.

With the milling of N2-9.5\_HX, the effect of tetragonal into monoclinic transformation is visible already after treatment at 300 rpm for 30 min (N3-9.5\_HXM\_300) and even more intense bands appear for sample N4-9.5\_HXM\_450 (appearance of features originating from translations and vibration of the ZrO<sub>2</sub> molecules). The spectra of these milled samples, containing about 90% of the monoclinic phase, are similar to each other and the bands characteristic for monoclinic ZrO2 are present at

406, 347, 262, and 230 cm<sup>-1</sup>. The shape of the spectra is strongly reminiscent of that of the commercial sample, suggesting that milling of the N2-9.5\_HX hydrothermal sample leads to a more ordered crystal structure with a monoclinic phase.

3.1.2 Structural analysis of the samples prepared at pH 7. It was noted that the monoclinic phase of ZrO2 is stable in the high-temperature range (200-900 °C), which is favorable for industrial catalytic processes.4 The usage of a tetragonal phase, which is unstable under these conditions, or a mixture of these two phases is undesirable because the phase shift from tetragonal to monoclinic is accompanied by an increase in volume by 3-5%, which may cause cracking of ZrO2 granules.27,60 Since modification of ZrO2 obtained at pH 9.5 did not lead to the formation of a pure monoclinic phase even after HTT at 320 °C, an attempt to reach this by lowering the precipitation pH was made. In terms of pH influence, it was established in previous studies that monoclinic zirconia is formed from precursor precipitated in the 6.5 to 10.5 pH range while the tetragonal precursor is obtained for the 3 to 4 or 13 to 14 pH range. 65,77,78 Precipitation from nitrate solutions also contributes to the formation of the monoclinic phase during post-synthetic modification.79

For further investigations, pH 7 was chosen for ZrO<sub>2</sub> precipitation since its lowering should promote the formation of monoclinic ZrO<sub>2</sub>,<sup>21,31,32,34</sup> albeit with a lower specific surface area.<sup>17,23,25,31</sup> HTT of the xerogel, precipitated at pH 7 and 260 °C (N10-7\_HX) results in the formation of a sample with a monoclinic phase content reaching 96% but the diffractogram of this sample contains very broad peaks (Fig. 3). This indicates that an imperfect crystal structure with a crystallite size of about 4.2 nm is formed. Subsequent milling of this hydrothermal sample at 300 rpm (N11-7\_HXM\_300\_1h\_10, N12-7\_HXM\_300\_0.5h\_13, and N13-7\_HXM\_300\_1h\_13) practically does not change the diffraction patterns of the resulting samples (Fig. 3). Finally, the elevation of HTT to 300 °C (N15-7\_HG) allows for obtaining a pure monoclinic phase with an improved structure and crystallite size of about 12.2 nm.

The negligible presence of an amorphous component in the hydrothermal samples can be confirmed by comparing the results of thermogravimetric analysis (DTA-TG) for the initial sample and the sample after HTT at 300 °C (Fig. S2a and b†). It is known that crystallization of the amorphous precipitated ZrO<sub>2</sub> is accompanied by an intense exo-effect on the DTA curve at 420–450 °C.  $^{26}$  Indeed, the acquired DTA curve of the initial sample (sample N9-7) contains a sharp exo-effect at 436 °C which corresponds to the crystallization of amorphous ZrO<sub>2</sub> (Fig. S2a†). In comparison, this effect is absent on the DTA curve recorded for the sample after HTT at 300 °C (N15-7\_HG, Fig. S2b†).

To completely transform the tetragonal phase into a monoclinic phase, HTT at 300 °C was increased to 7 hours (N17-7 H7hG). The diffraction pattern of sample N17-7 H7hG contains only peaks attributed to the monoclinic phase (Fig. 4). The milling of this hydrothermal sample at 400 rpm and BPR = 10 does not cause a change in the phase composition and the diffraction pattern of the milled sample N21-7\_H7hGM\_400\_0.5\_10 shows only reflexes of monoclinic ZrO2. Besides, the slight background in its diffraction pattern and calculations indicate the absence of a noticeable amount of amorphous phase. Also interestingly, even harsh milling (500 rpm, BPR 25) of this sample does not lead to broadening of the peaks in the diffraction pattern (N19-7 H7hGM 500 0.5 25 in Fig. 4). This indicates that the crystallite size does not change for this series of samples. The diffractogram shows a lowintensity peak at  $2\Theta = 30.18^{\circ}$ , indicating the presence of a cubic rather than a tetragonal phase. This can be concluded based on the absence of the peak at 43.2° and two peaks in the range of 73-75° by which tetragonal and cubic phases can be distinguished.80 Also, the monoclinic phase can be transformed into a cubic phase during milling, as reported in earlier studies.60,62,66,72,81

It is also important to compare the thermal behavior of the X-ray amorphous initial sample (N9-7) with the samples after HTT

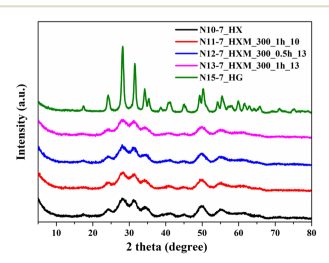


Fig. 3 XRD patterns for ZrO<sub>2</sub>, precipitated at pH 7, after HTT additional MChT under different conditions.

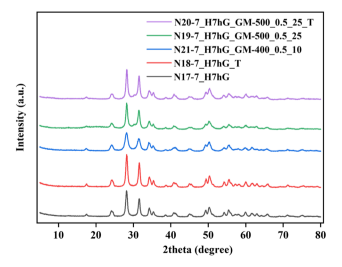


Fig. 4  $\,$  XRD patterns for ZrO<sub>2</sub>, precipitated at pH 7, after HTT, additional milling and calcination.

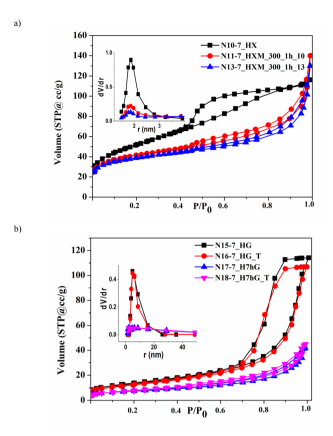


Fig. 5 Nitrogen adsorption-desorption isotherms and PSD curves (insets) for samples of ZrO2, precipitated at pH 7, after HTT of the xerogel (a) and gel (b).

and milling. It can be seen that the initial sample after calcination at 500 °C (N14-7\_T) contains a predominant monoclinic phase and about 12% of a tetragonal phase (Fig. 5). Calcination of the hydrothermally treated sample (N15\_7\_HG) at 500 °C results in conservation of the pure monoclinic phase (N16-7\_HG\_T). The phase composition of the milled sample does not change after its post-calcination at 500 °C. Moreover, the position of the main peak of the monoclinic phase also does not change in all these diffractograms and the values of FWHM for this peak are within 0.55-0.57°.

Increasing the time of HTT from 5 to 7 h (N17-7\_H7hG) allowed a material to be obtained that contains only the monoclinic phase. Therefore, for a more precise analysis of the crystal structure of the pure phase samples presented in Fig. 4, lattice parameters and microstructural parameters (average grain size D and microstrains  $\langle \varepsilon \rangle$ ) were derived by full profile Rietveld refinement, Table 3. It should be noted that the lattice parameters of the sample after HTT at 300 °C are very close to the tabulated data for monoclinic ZrO<sub>2</sub> (JCPDS, no. 65-1025).<sup>71</sup>

A comparison of the results presented in Table 3 shows that the most detectable decrease in all lattice parameters, monoclinic angle  $\beta$ , and unit cell volume V (by 0.1–0.4%) is observed after milling (samples N19-7\_H7hGM\_500\_0.5\_25 and N20-7\_H7hGM\_500\_0.5\_25\_T). Also, an increase in the concentration of the microstrain value  $\langle \varepsilon \rangle$  by 52% occurs, similarly as reported in the work<sup>62</sup> in which  $\langle \varepsilon \rangle$  is approximately 0.00555 after milling commercial monoclinic ZrO2 for 30 min at 500 rpm using a stainless-steel assembly. This indicates the introduction of defects into the crystal structure, which takes place during milling. It is also important that post-calcination of the milled sample (N19-7\_H7hGM\_500\_0.5\_25) at 500 °C (as in ref. 60) - sample N20-7\_H7hGM\_500\_0.5\_25\_T, leads to minimal changes in lattice parameters (less than 0.1%), the microstrain values, and average grain size D. The results provide evidence that the introduced defects are quite stable at this temperature.

#### 3.2 Porous structure of the samples prepared at pH 9.5 and 7

It is known that zirconium dioxide, precipitated from aqueous solutions, is a micro-mesoporous material. 26,27 The parameters of the porous structure for all prepared samples are presented in Table 4. It should be noted that only the as-precipitated samples (N1-9.5 and N9-7) contain micropores, the volume of which is given in brackets in Table 4: 0.04 and 0.03 cm<sup>3</sup> g<sup>-1</sup>, respectively. This is 40-50% of the total pore volume (column 3 of Table 4).

HTT of both as-precipitated samples promotes the formation of a developed and uniform mesoporous structure: the volume of mesopores increases to 0.17-0.23 cm<sup>3</sup> g<sup>-1</sup> and their size is 7.9-9.7 nm, as can be seen for samples N2-9.5\_HX, N5-9.5\_HX, N7-9.5 HG, N15-7 HG, and N17-7 H7hG (Table 4). For all

Table 3 Lattice and microstructural parameters (average grain size D and microstrains  $\langle \epsilon \rangle$ ) for samples containing a pure monoclinic phase

Samples	Lattice parameters		Microstructural parameters
N17-7_H7hG	a = 5.1484(7)  Å	b = 5.2096(8)  Å	$\langle arepsilon  angle = 0.00208$
	c = 5.3228(8)  Å $V = 140.83(6) \text{ Å}^3$	$\beta = 99.437(7)^{\text{ o}}$	D=26.9  nm
N18-7_H7hG_T	v = 140.83(6)  A a = 5.1488(6)  Å	b = 5.1969(8)  Å	$\langle arepsilon  angle = 0.00228$
	c = 5.3193(7)  Å	$\beta = 99.418(6)^{\text{ o}}$	D = 26.5  nm
N19-7 H7hGM 500 0.5 25	$V = 140.41(6) \text{ Å}^3$ a = 5.143(2)  Å	b = 5.203(2)  Å	$\langle arepsilon  angle = 0.00318$
1V15 /_11/1IGH1_000_0.0_20	c = 5.315(2)  Å	$\beta = 99.03(1)^{\circ}$	D = 32.7  nm
	$V = 140.5(2) \text{ Å}^3$	1 - 10=(0) 3	/ \
N20-7_H7hGM_500_0.5_25_T	a = 5.146(1)  Å c = 5.319(2)  Å	b = 5.197(2)  Å $\beta = 99.122(9) ^{\text{o}}$	$\langle arepsilon  angle = 0.00325$ $D = 30.1 \ \mathrm{nm}$
	$V = 140.5(1) \text{ Å}^3$	,(*)	

Table 4 Parameters of the porous structure of prepared samples

Samples	$S,  \mathrm{m^2  g^{-1}}$	<i>V</i> , cm <sup>3</sup> g <sup>-1</sup>	$d_{ m me}$ , nm
N1-9.5	166	0.06(0.04)	3.5
N2-9.5_HX	61	0.20	7.9
N3-9.5_HXM_300	66	0.16	6.5
N4-9.5_HXM_450	67	0.15	3.9
N5-9.5_HX	66	0.23	9.5
N6-9.5_HXM_300	74	0.19	6.5
N7-9.5_HG	87	0.175	6.6
N8-9.5_HGM_300	100	0.165	3.7
N9-7	130	0.07(0.03)	2.8
N10-7_HX	184	0.18	3.7
N11-7_HXM_300_1h_10	149	0.215	3.7
N12-7_HXM_300_0.5h_13	144	0.205	3.7
N13-7_HXM_300_1h_13	139	0.20	3.7
N14-7_T	27	0.07	6.5
N15-7_HG	60	0.17	9.7
N16-7_HG_T	61	0.24	9.7
N17-7_H7hG	48	0.18	9.7
N18-7_H7hG_T	44	0.17	9.7
N19-7_H7hGM_500_0.5_25	28	0.07	3.8; 13.0
N20-7_H7hGM_500_0.5_25_T	28	0.07	3.8; 13.0

modified samples, the total pore volume V coincides with the volume of mesopores  $V_{\rm me}$ . This coincidence, as well as the calculations of microporosity using the t-method, indicates the absence of micropores in all modified samples.

One can see that the isotherms recorded for hydrothermal samples of both series contain pronounced capillary-condensation hysteresis which is characteristic of mesoporous materials. This is related to the samples precipitated at pH 9.5 (Fig. S3†). Similar isotherms were obtained for samples precipitated at pH 7 (Fig. 5) which have larger sizes of mesopores – 9.7 nm (Table 4, N15-7\_HG and N17-7\_H7hG).

The specific surface area of the sample as-precipitated at pH 9.5 (N1-9.5) is higher than that of the as-precipitated sample at pH 7 (N9-7) which is consistent with the results of earlier studies. 17,23,25 Its HTT leads to a reduction in specific surface area: all hydrothermal samples modified at 260-320 °C have a specific surface area within 60-87 m<sup>2</sup> g<sup>-1</sup> (N2-9.5\_HX, N5-9.5\_HX, and N7-9.5\_HG). In contrast, the HTT of the sample, precipitated at pH 7, in the form of a dried xerogel at 260 °C (N10-7\_HX), causes an increase in specific surface area to 184  $m^2$   $g^{-1}$ . This is obviously due to the fact that a fine crystalline structure is formed under these conditions, as was shown above according to the XRD data. As a result, the coalescence of these small crystallites leads to the formation of a finely porous structure with a much smaller size of mesopores – 3.7 nm – and, accordingly, a higher specific surface area. On the other hand, an increase in HTT temperature to 300 °C leads to the reduction of specific surface area to 60 m<sup>2</sup> g<sup>-1</sup> (N15-7\_HG).

Milling at low and medium intensity (300–450 rpm, BPR 10–13) causes partial destruction and transformation of the porous structure of all hydrothermally modified samples due to changes in particle morphology and pore shape (N3-9.5\_HXM\_300, N4-9.5\_HXM\_450, N6-9.5\_HXM\_300, N8-9.5\_HGM 300, N11-7\_HXM 300 1h 10, N12-

7\_HXM\_300\_0.5h\_13, and N13-7\_HXM\_300\_1h\_13). samples precipitated at pH 9.5, the specific surface area increases slightly after milling and the volume of mesopores, on the contrary, decreases (Table 4, N3-9.5\_HXM\_300, N4-9.5 HXM 450, N6-9.5 HXM 300, and N8-9.5 HGM 300). Milling of the hydrothermal sample with high specific surface area (precipitation at pH 7 after HTT of the xerogel at 260 °C) is accompanied by its reduction (samples 7\_HXM\_300\_1h\_10, N12-7\_HXM\_300\_0.5h\_13, and N13-7 HXM 300 1h 13). The volume of mesopores for this sample also slightly decreases after milling. But the diameter of the mesopores  $d_{\mathrm{me}}$  remains unchanged. The change in the shape of nitrogen adsorption-desorption isotherms for milled samples of both series confirms this (Fig. S3<sup>†</sup> and 5). This can also be seen from the pore volume distribution (PSD) curves by size shown in the inset of Fig. 5a.

Increasing the intensity of milling up to 500 rpm at BPR 25 results in more significant destruction of porosity and a corresponding decrease in specific surface area, volume, and size of mesopores (N19-7\_H7hGM\_500\_0.5\_25, Table 4 and Fig. 5b). The PSD curve for this sample contains a second diffuse maximum centered at around 13 nm, which is visible in Fig. S4.† Mesopores with a size greater than 5 nm make a significant contribution to the total pore volume of the milled samples N11-7\_HXM\_300\_1h\_10 and N13-7\_HXM\_300\_1h\_13, while for the hydrothermal sample N10-7\_HX this contribution is minimal. Consequently, the bi-porous structure is formed under milling. The presence of pores with a larger size – 5–50 nm – should contribute to the acceleration of the diffusion of reagents and products in catalytic processes.

It should be noted that the results obtained – multidirectional changes in porosity parameters – coincide with the general trends that are observed during dry milling of porous oxides with different values of specific surface area. 50,76,82

As mentioned above, the porous structure of hydrothermal samples has higher thermal stability, i.e. they are sintered to a lesser extent with increasing temperature. This is mainly because they have a larger pore size and a more uniform pore structure. Thus, if the specific surface area of the as-precipitated sample N9-7 after post-calcination at 500 °C decreases almost 5 times and is 27 m<sup>2</sup> g<sup>-1</sup> (N14-7\_T), then it does not change for hydrothermal sample N15-7\_HG and has a value of 61 m<sup>2</sup> g<sup>-1</sup> (N16-7\_HG\_T). Moreover, the volume of mesopores in this hydrothermal sample is significantly higher after postcomparison calcination. of samples N19-7 H7hGM 500 0.5 25 and N20-7 H7hGM 500 0.5 25 T also indicates that the sample, successively modified by HTT and milling, does not change the porous structure after postcalcination. The described results are well illustrated in Fig. 5b, which shows corresponding isotherms and PSD curves. As can be seen, the pairs of isotherms and PSD curves for samples before and after post-calcination practically coincide.

The monoclinic  $ZrO_2$  phase and the developed uniform mesoporous structure are optimal from the point of view of their application in catalysis. In fact, mesoporous catalysts are the most demanded for many processes because they provide a balance between a good diffusion rate of reactants and

a sufficiently high accessible surface, as well as resistance to sintering and coking through the confinement effect. 12,15-21

#### Morphology

Scanning electron microscopy was used to shed light on the material's morphology undergoing various treatments. Examples of micrographs of hydrothermal, milled and post-calcined samples were acquired for pure phase samples obtained at pH 7 and are shown in Fig. 6. The most pronounced effect can be observed for milling of the sample N10-7\_HX (Fig. 6a and b), which results in the diminution of the grains and formation of a loose layer of smaller particles on the top of the bigger ones (N11-7\_HXM\_300\_1h\_10, Fig. 6c and d). The following calcination leads to the sintering of those smaller particles yet does not influence the overall size of the grains (N20-7\_H7hGM\_500\_0.5\_125\_T, Fig. 6e and f).

SEM images of the selected pure monoclinic ZrO2 samples precipitated at pH 7 (Fig. 7a and b) indicate that milling affects their morphology much more than calcination. On the micrometer scale, the diminution of the grains is well visible, when comparing the hydrothermally treated gel (N17-7\_H7hG) and its

milled counterpart (N19-7\_H7hGM\_500\_0.5\_25), see Fig. 7a and e. On the other hand, the changes observable on the nanometer scale show that an initial fine structure of the material is being crushed into agglomerates when milled at 500 rpm (Fig. 7b and f). The post-calcination does not seem to affect the material grains, see the comparison of samples N17-7\_H7hG with N18-7\_H7hG\_T (Fig. 7a and c) and N19-7\_H7hGM\_500\_0.5\_25 with N20-7 H7hGM 500 0.5 25 T (Fig. 7e and g).

#### 3.4 Optical properties and photocatalytic activity

In general, zirconium dioxide is a semiconductor with a band gap of 5.0-5.8 eV.36,43,83 Its spectra contain the main band in the UV region, namely at 220–250 nm, which is attributed to the  $O^{2-}$ → Zr<sup>4+</sup> transition. UV-vis spectra of the samples obtained at pH 7 gathered in Fig. 8 show that a well-crystallized sample after HTT of a gel at 300 °C (N17-7\_H7hG) has a band with a maximum of around 227 nm and an absorption edge at 241 nm (Fig. 8a). The absorption edge is determined as the point of the intersection of the tangent with the abscissa axis (solid black line). This corresponds to band gap  $E_g = 5.10$  eV (Tauc plot in Fig. 8b, solid black line).

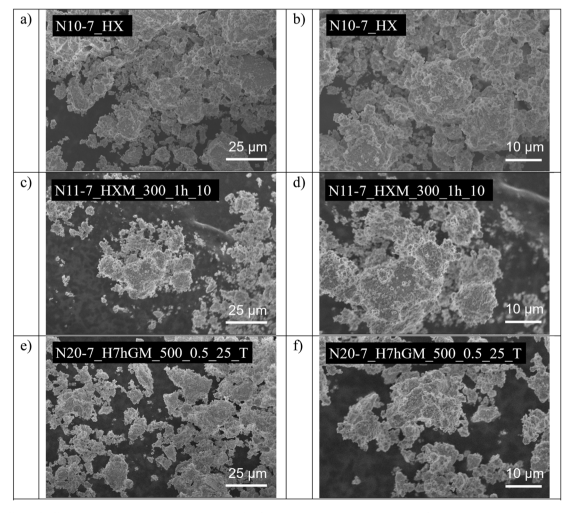


Fig. 6 The effect of different treatments on the morphology of ZrO<sub>2</sub> precipitated at pH 7, samples N10-7\_HX: hydrothermal treatment (a and b), milling (c and d), and post-calcination (e and f).

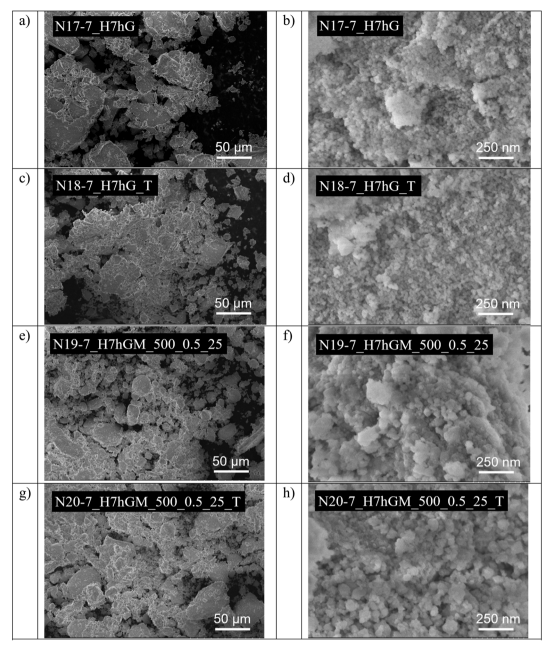


Fig. 7 The effect on the morphology of the hydrothermal sample N17-7\_H7hG (a and b), and its subsequent calcination (c and d), milling (e and f), and post-calcination (g and h).

Milling this sample at 500 7\_H7hGM\_500\_0.5\_25) and post-calcination of the latter (N20-7\_H7hGM\_500\_0.5\_25\_T) results in a shift of the absorption edge toward 247 nm (blue dashed line in Fig. 8a) which corresponds to band gap  $E_{\rm g}$  of about 5.02 eV (blue solid line in Fig. 8b). It is noteworthy that these two spectra contain a socalled Urbach tail (marked with an arrow in Fig. 8a), extending to the visible region and characterizing the set of defect states of milled ZrO2.84 As can be seen from Fig. 8a, this is not observed for the hydrothermal sample N17-7\_H7hG. The presence of an Urbach tail may be caused by the formation of oxygen vacancies in the ZrO2 structure, which is typical for the dry milling of oxides.85,86 As a result of the extended absorption

range, the second band gap at about 2.80 eV is recorded on the Tauc plot for milled and post-calcined samples (Fig. 8b, blue dashed line) which is observed for oxygen vacancy-rich ZrO<sub>2</sub>. 83,87,88 It is important that post-calcination of this milled sample at 500 °C (N20-7\_H7hGM\_500\_0.5\_25\_T) practically does not lead to a change in the spectra, see Fig. 8a and b. This is consistent with the thermal stability of the crystal structure of milled samples as described above. As can be seen, the second band gap is absent on the Tauc plot for the sample before milling – for sample N17-7\_H7hG (Fig. 8b).

Due to its wide band gap, zirconium dioxide is photocatalytically active only under the influence of UV radiation. However, as previously reported in several studies, the

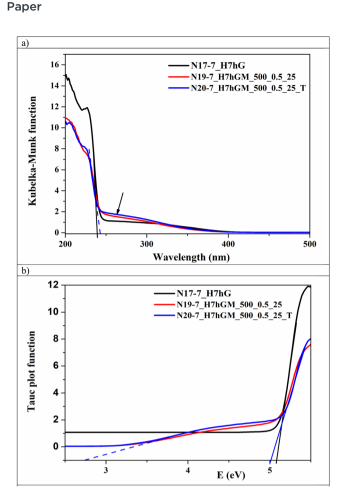


Fig. 8 UV-vis spectra (a) and Tauc plots (b) for several ZrO<sub>2</sub> samples, precipitated at pH 7.

introduction of oxygen vacancies into the structure of ZrO<sub>2</sub> promotes its acquisition of photocatalytic activity during the decomposition of organic pollutants in visible light.26,87,89 Testing the selected samples with the pure monoclinic phase and high specific surface area under visible radiation was carried out in the process of degradation of a stable model dye molecule - safranin T. The initial sample precipitated at pH 7, as well as the same after HTT at 260 and 300 °C, was predicted to be inactive according to our previous experiments,27 while milling of hydrothermal sample N10-7\_HX at 300 rpm and hydrothermal sample N17-7 H7hG at 500 rpm (N11-7\_HXM\_300\_1h\_10 and N19-7\_H7hGM\_500\_0.5\_25, respectively) activates them as photocatalysts in this process. An example of change in the spectra of an aqueous solution of safranin T vs. time of irradiation for a sample N19-7H7hGM 500 0.5 25 is shown in Fig. S5.†

As can be seen, the intensity of the characteristic band at 520 nm steadily decreases with increasing irradiation time. In parallel, a decrease in the intensity of the bands in the region of 240-280 nm is also observed, which indicates the splitting of the chromophore rings in the safranin molecule.65 As a result, the decolorization of the dye solution reaches 75% for the most active sample N11-7\_HXM\_300\_1h\_10 during the applied time window. In addition, the reduction of total organic carbon

(TOC) in the solution as a result of the photocatalytic process reached 62%. The degree of TOC reduction is a very important additional signature, in addition to the decolorization of the solution, since it indicates the degree of mineralization of the dye, i.e. the transformation of the organic pollutant into less harmful inorganic substances,65 potentially offering the applications of these new materials in the remedy of notoriously stable water pollutants.

The resulting temporal changes in the optical density of the band at 520 nm were approximated using a pseudo-first-order kinetic equation. The plotted kinetic curves are shown in Fig. S6.† The results fit well with linear fitting ( $R^2 = 0.96-0.99$ ). As mentioned above, the sample subjected to HTT alone was process. Milled inactive in this samples 7\_HXM\_300\_1h\_10, N19-7\_H7hGM\_500\_0.5\_25), as well as the post-calcined sample (N20-7\_H7hGM\_500\_0.5\_25\_T), demonstrate fairly high activity. This can be explained by the fact of speculative formation of oxygen vacancies in the ZrO<sub>2</sub> structure, which is beneficial for photocatalytic activity, as previously reported.87,89 The observed difference in the activity of tested samples may be caused by their different physicochemical characteristics. 59,90,91 Thus, a slightly higher value of the rate constant of safranin degradation  $k_d$  for a sample milled at 300 rpm (N11-7\_HXM\_300\_1h\_10) compared to that milled at 500 rpm (N19-7\_H7hGM\_500\_0.5\_25) - reaching  $4.6 \times 10^{-5}$  and  $4.2 \times 10^{-5}$  s<sup>-1</sup>, respectively – may be associated with a higher specific surface area of the first sample (Table 4). Simultaneously, the lower  $k_d$  value (3.2 × 10<sup>-5</sup> s<sup>-1</sup>) for the post-calcined sample (N20-7\_H7hGM\_500\_0.5\_25\_T), despite the same values of the specific surface area and band gap, may be due to dehydroxylation of the surface during calcination. This leads to a decrease in the content of active sites and adsorption of safranin on the surface of the post-calcined sample (N20-7\_H7hGM\_500\_0.5\_25\_T) compared to the as-milled one (N19-7 H7hGM 500 0.5 25). Since the degree of the dve's photocatalytic breakdown is directly correlated with safranin adsorption, the post-calcined sample's activity is lowered. 26,59,90

## 3.5 Post-treatment and mechanochemical aspects of the preparation

The set of obtained results demonstrates the dependence of the ZrO<sub>2</sub> structure (in a broad sense - porosity, crystallinity, and surface design) on the conditions of its precipitation and postsynthetic modification. It is also important that changes in the main structural characteristics due to modification are interrelated. As-precipitated at pH 9.5 and 7 ZrO<sub>2</sub> samples (N1-9.5 and N9-7) are mostly X-ray amorphous. But the first of them has a higher specific surface area due to the higher content of micropores. The following HTT contributes to the elimination of microporosity and the development of mesoporosity in both precipitated samples, which is a general tendency of hydrothermal processes.<sup>27,35</sup> Differences in the change of the specific surface area may be caused by features of the course of crystallization of both samples under hydrothermal conditions. As mentioned above, crystallization processes during HTT of these samples proceed in different ways. At 260 °C, a mixture of tetragonal and monoclinic phases with a predominance of the first is formed for the sample precipitated at pH 9.5 (N1-9.5), while the HTT product of the sample, precipitated at pH 7 (N9-7), is a pure monoclinic phase (compare Fig. 1 and 3). This difference can be explained in terms of the influence of impurity ions on phase transformation in ZrO2.92 Not only transition metal cations but also alkali metal and ammonium cations have a stabilizing effect on the preservation of the tetragonal phase at low temperatures.93 A sample, precipitated at pH 9.5 (N1-9.5), contains NH<sub>4</sub><sup>+</sup> cations in ion-exchanged positions which are not completely removed by washing. Due to this, the tetragonal phase is stabilized after HTT at 260 °C (N2-9.5\_HX). According to the results of the work,94 the removal of volatile impurities leads to the appearance of shear stresses in the ZrO2 lattice during the desorption of stabilizing species, which can lead to a tetragonal transformation into a monoclinic one. During postmilling of the hydrothermal sample N2-9.5 HX, removal of ammonium impurities is possible due to mechanical action and local temperature increase which takes place under these conditions.52,53 This may be another reason (along with the influence of water) for the rapid transformation of the tetrag-

The sample precipitated at pH 7 (N9-7), *i.e.* near the point of zero charge for ZrO<sub>2</sub>, absorbs fewer ions, that can stabilize the tetragonal phase, and is more easily washed away from them. Therefore, an almost pure fine-crystalline monoclinic phase is formed already at 260 °C during post-HTT of this sample. The latter factor is the reason for the high specific surface area of this hydrothermally modified sample (Table 4, sample N10-7\_HX).

onal phase into a monoclinic phase.

Increasing the HTT temperature and time promotes the complete transformation of the tetragonal phase into a monoclinic phase for both as-precipitated samples, regardless of the conditions of their precipitation. This is also fully consistent with the patterns of hydrothermal transformations in oxides.<sup>35</sup> As a result, a temperature of 300 °C and a duration of 5-7 hours are optimal HTT conditions, if several important requirements are taken into account, as confirmed in this work. They are as follows: (i) a well-crystallized monoclinic phase without a tetragonal phase admixture; (ii) a uniformly developed porous structure; (iii) stability of the phase composition and porous structure with increasing temperature of post-calcination. Relatively soft dry milling applied for this series of samples allowed their properties to be influenced in terms of surface structure as shown with the XRD and UV-vis results. Its advantage lies in providing a controlled introduction of defects into the structure while maintaining the monoclinic phase with minimal change in the porosity, which proves mechanochemistry to be a very useful tool for obtaining highly active catalytic materials. By governing the milling intensity one can achieve a spectrum of materials with tuned properties. It was reported in ref. 87 and 88 that oxygen vacancies are responsible for the photocatalytic activity of defected ZrO<sub>2</sub>. The results obtained on the post-calcined milled samples indicate the stability of the monoclinic phase composition, conserved porous structure, and its defects. This finding is consistent with the conclusion of the authors expressed in studies<sup>83,95</sup> about the stabilization

effect of the defects on monoclinic  $ZrO_2$ . Thus, we hypothesize the formation of defects, most likely oxygen vacancies, based on the observed acquisition of photocatalytic activity of  $ZrO_2$  with vacancies introduced by milling, and these defects can contribute to the stabilization of the monoclinic phase. The presence of oxygen vacancies is indirectly confirmed by the UV-vis spectra (Fig. 8a) and preservation of the photocatalytic activity of these post-calcined samples (Fig. S5†). Based on this, it can be assumed that thermostable  $ZrO_2$ , modified in this way, will be promising for application in other catalytic processes, for example in  $CO_2$  and  $CH_4$  high-temperature conversion.

## 4. Conclusions

The sequential use of hydrothermal treatment at 260-300 °C and milling at 300-450 rpm as post-synthetic modification strategies makes it possible to vary the physicochemical characteristics of precipitated zirconium dioxide and its following catalytic activity. For both ZrO<sub>2</sub> samples - precipitated at pH 9.5 and 7 - elevation of HTT temperature to 300 °C leads to the complete transformation of a mixture of tetragonal/monoclinic phases into pure monoclinic ZrO<sub>2</sub> with varying degrees of crystallinity. The milling of these hydrothermally modified samples introduces defects into their structure, as indicated by the results obtained using XRD and UV-vis spectroscopy. Postmilling of low and medium intensity (300-450 rpm, BPR 10-13) allows ZrO<sub>2</sub> to be obtained with a sufficiently high specific surface area, volume, and pore size, with defects stable also after post-calcination. More intensive milling (500 rpm, BPR 25) causes significant destruction of the porous structure and partial amorphization of ZrO<sub>2</sub>. As a result, milled samples acquire photocatalytic activity under visible irradiation.

Based on the obtained results, new routes for the synthesis of a thermally stable mesoporous monoclinic ZrO<sub>2</sub> support containing stable defects introduced by milling are proposed. This can provide higher material resistance not only against the sintering of the porous structure and indirectly deposited metal particles, but also the reduction of coking during catalytic processes at high temperatures. For *in situ* characterization under real catalytic conditions, ZrO<sub>2</sub> samples with pure phases would facilitate the interpretation of the results and distinguishing particular phase activity. Therefore, the proposed combined synthesis route makes it possible to use the advantages of both hydrothermal and mechanochemical methods in order to prepare pure monoclinic zirconium dioxide with optimal characteristics for application as a catalyst support including in high-temperature catalytic processes.

# Data availability

The data supporting this article have been included as part of the ESI.†

## **Author contributions**

Conceptualization, V. Sydorchuk and J. E. Olszowka; sample preparation, S. Levytska; milling experiments, O. Kiziun; XRD

studies, L. Vasylechko; data acquisition, B. Billinghurst, K. Simkovicova, S. Valtera, and J. E. Olszowka; data curation, K. Simkovicova; visualization, K. Simkovicova and S. Valtera; writing - draft preparation, J. E. Olszowka, V. Sydorchuk, S. Valtera, K. Simkovicova, B. Billinghurst, and S. Vajda. All authors have approved the final version of the manuscript.

## Conflicts of interest

There are no conflicts to declare.

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