In-flight Synthesis of Nanosized ZrC Particles from Various Precursors in RF Thermal Plasma

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Abstract

Synthesis of zirconium carbide (ZrC) powder was investigated applying a non-conventional atmospheric radiofrequency (RF) thermal plasma process. In one case, zirconium dioxide (ZrO$_2$) was reacted with solid carbon or with methane with varying molar ratio. In the other, zirconium-propoxide (NZP), containing both constituents, was thermally decomposed in the Ar plasma. Temperature-dependent thermodynamic analysis was performed in the 500-5500 K temperature range to estimate the formation of possible equilibrium products for each reaction stoichiometry. Broad temperature range exists for the stability of solid ZrC for each explored reaction system. In accordance with this prediction, X-ray diffraction studies detected the ZrC as the major phase in all the prepared powders. The yield of particular runs ranged from 39 % to 98 %. Practically, full conversion was typical for the case of NZP precursor, however only partial conversion could be detected in ZrO$_2$ reactions. The average particle size of the powders falls between 10 nm and 100 nm depending on the type of the reaction systems (either calculated from the specific surface area or derived from broadening the XRD reflections). The transmission electron micrographs indicated mostly globular shape of the nanosize particles. Quantitative analysis of the surface of the powders by X-ray photoelectron spectroscopy revealed the presence of oxygen and carbon. Evaluating the spectra of the powders prepared from NZP, and taking in the account its spherical shape, a ZrC core covered by a very thin ($\approx$1.0 nm) ZrO$_2$ layer may be accounted for the measured oxygen and a thicker carbonaceous layer.

Keywords

nanopowder synthesis, RF thermal plasma, ceramic nanopowders, zirconium carbide

1 Introduction

Zirconium carbide (ZrC) is a unique ceramic material belonging to the class of ultra-high temperature ceramics (UHTC) and is an important non-oxide ceramic material with attractive physical and chemical properties, such as high hardness (28.7 GPa), high strength, high melting point (3445 °C), high chemical inertness and good thermal shock resistance. Based on these outstanding properties, ZrC is widely used in different ultra-high temperature applications, as cutting tools in the mechanical, as a coating in the nuclear industry and as reinforcing particles in wear-resistant materials and electrical devices. Furthermore, it is known as one of the main kinds of far infrared ceramics [1–4]. ZrC, however, shows very poor sinterability, hindering its extensive applications. The latter can be improved by applying smaller particles.

Over the last decades, nanosized grain materials have gained increasing interests due to the absence of extended defects in these particles [5]. Nanosized ZrC can decrease the sintering temperature and facilitate densification processes due to the larger surface area and higher reactivity of the smaller particles [6]. Moreover, it was shown that sintered ceramics from ultrafine grain size powder may also improve their mechanical properties significantly [7–8].
The conventional synthesis of metal carbides generally involves the reduction of metal oxides in a carbon-rich environment. The carbothermal reduction method generally requires relatively high reaction temperatures, such as 1600–2200 °C and long reaction time, e.g., several hours in the case of ZrC. In addition, these solid-state reactions result in powders with relatively large particle sizes. Hence in the case of many applications, additional milling is required, which tends to be expensive and may lead to contamination of the powder [9–10]. Several other methods have also been applied for ZrC synthesis, such as self-propagating high-temperature synthesis (SHS), pyrolysis and high energy ball milling [11–14], however, these synthesis routes yield again only submicron-sized particles. Meanwhile, only a few methods have been reported on preparing ZrC nanopowders. These are mainly solution-based processes, such as sol-gel or liquid polymer precursor methods [15–19].

Plasma processes are a special group of gas-phase reactions. A large number of publications, including a set of our previous papers, pointed out that due to its unique properties, thermal plasma processes offer an appealing route for preparing nanosized oxide, nitride, carbide and boride ceramic nanopowders [20–26]. Thermal plasmas possess extremely high temperatures (up to 15,000 K in the flame), large plasma volume, high enthalpy and high concentrations of chemically reactive species, enhancing the reactivity of the precursors injected in the plasma [27–29]. Besides, the typically fast cooling rate (10⁴ K s⁻¹) inhibits grain growth of the small molecular clusters thus promotes the formation of nanopowder. Plasma synthesis involves a one-step process, in which reactions take place within milliseconds, thus the production of nanoparticles can be well controlled with operating time and applied specific power, providing possibility for mass production of nanoparticles with high rate. In addition to above listed advantages, the thermal plasma synthesis is flexible in terms of precursor material selection [30].

In some cases, for the preparation of different high-temperature transition metal carbides, oxide precursors were used in the plasma synthesis method [30, 31]. Recently, using of organometallic precursors, containing both the metal and carbon constituents are preferred [32].

Plasma synthesis of nanosized ZrC powder was also carried out using hydrocarbon as carbon and zirconium(IV) chloride (ZrCl₄) as zirconium source [33]. The disadvantage of applying ZrCl₄ precursor includes the formation of corrosive HCl as a reaction by-product.

In this study, we investigated the preparation of ZrC nanopowder in a radiofrequency (RF) thermal plasma using liquid (NZP) or solid (ZrO₂) as precursors. These two cases involve different evaporation and synthesis routes. In case of NZP, both elements of the ZrC are present in the precursor. Methane or graphite powder was used as carbon sources in the case of ZrO₂. Involvement of ZrO₂ in this study was motivated by consideration of a perspective mass production of ZrC, because a cost-effective process can be based primarily on the readily available ZrO₂.

2 Experimental

The experiments were carried out in an inductively coupled RF thermal plasma system using a five turn coil torch (TEKNA PL-35, TEKNA Ltd., Canada) connected to a radio frequency (RF) generator (LEPEL, USA) with a maximum power of 30 kW at 4–5 MHz. The torch was mounted vertically on top of the reactor connected simultaneously to a cyclone and a vacuum pump. A detailed description of the system and schematic presentation is illustrated in our previous publications [34–36]. Zirconium(IV)-propoxide solution (NZP: Zr(OCΗ₂CH₂CH₃)₄, 70 wt% in 1-propanol, from Sigma-Aldrich) was applied as the zirconium and carbon source, the liquid was delivered by a peristaltic pump (Masterflex) to the atomizer probe placed in the hottest zone of the plasma flame through a water-cooled probe by a powder feeder (PRAXAIR). When graphite powder (KS4, Timcal Co., mean particle size 2.8 μm) was used as carbon sources, it was mixed with the ZrO₂ and fed together. The plasma (central) gas was a mixture of Ar and He, where the role of helium is to increase heat conductivity and the enthalpy of the plasma [20], while Ar was used as sheath gas as well. The precursors were fed with a carrier gas of Ar, or in one case N₂. The purity of these gases (Linde Gáz Magyarország Ltd.) was 99.996 %. In the other set of ZrO₂ experiments, methane (99.5 %, Messer Hungarianz Ltd.) was introduced in the plasma reactor mixed with Ar in the sheath gas. The power was set to 25 kW for each run. The selection of working parameters (Table 1) was based on our previous experiments at the synthesis of different (oxide, boride) nanoparticles using the same equipment [23, 37–39]. In this study the following input ratios were explored: ZrO₂ + 3C, ZrO₂ + 5C, ZrO₂ + 2CH₄, ZrO₂ + 3CH₄, NZP and NZP + 15N₂.

Solid product for the analysis was collected from the inner surface of the reactor wall.
Table 1 Experimental condition for the synthesis of ZrC nanoparticles

| Typical operating parameters in RF thermal plasma |
|--------------------------------------------------|
| Sheath gas and flow rate                         | Ar 30 dm$^3$ min$^{-1}$ |
| Plasma gas and flow rate                         | Ar–He (15:10) 25 dm$^3$ min$^{-1}$ |
| Carrier gas and flow rate                        | Ar or N$_2$ 8 dm$^3$ min$^{-1}$ |
| RF power                                         | 25 kW |
| Reactor pressure                                 | 80 kPa |

X-ray diffraction (XRD) patterns were obtained by a Philips PW 3710 instrument equipped with a PW 1050 Bragg–Brentano parafocusing goniometer, using monochromatized Cu Kα radiation. The XRD scans were digitally recorded with steps of 0.041 in the 2 theta range from 4° to 80°.

The quantitative phase composition was evaluated using a full profile fitting method with corrections for preferred orientation and micro absorption [40]. The specific surface area was measured by nitrogen adsorption at 77 K and evaluated based on BET isotherms (Autosorb 1C, Quantachrome, seven point isotherm). The average particle size of synthesized powders was calculated by the following relation: $SSA = \frac{6}{\rho d}$, where $\rho$ is the absolute particle density (cm$^3$g$^{-1}$), $d$ is the diameter (μm) of the spherical particles and SSA is the measured specific surface area (m$^2$g$^{-1}$).

The morphology of synthesized powders was characterized by scanning electron microscopy (SEM, Zeiss type DSM 982 GEMINI with a field emission tungsten cathode) equipped with energy dispersive X-ray analysis (EDX) and transmission electron microscopy (TEM, Morgagni 268D). Raman spectroscopy measurements were performed using a Horiba Jobin-Yvon LabRAM-type micro spectrometer equipped with an Nd-YAG laser source, 40 mW laser power supplied with an Olympus BX-40 optical microscope. The measurements were carried out with 532 nm excitation wavelength, at low temperature (−20 °C) using a Linkam THMS600 temperature-controlled microscope stage. X-ray photoelectron spectra were recorded on a Kratos XSAM 800 spectrometer operating in fixed analyzer transmission mode, using Mg Kα$_{1,2}$ (1253.6 eV) excitation. The pressure of the analysis chamber was lower than 1 × 10$^{-7}$ Pa. Photoelectron lines of the main constituent elements, i.e., the Zr3d, C1s and O1s, were recorded by 0.1 eV steps. Spectra were referenced to the energy of the Zr3d$_{5/2}$ line of the carbide type zirconium, set at 179.4 ± 0.1 eV binding energy (B.E.). Data were processed by the Kratos Vision 2 program, while the quantitative analysis, based on peak area intensities, and model calculations were performed by the XPS MultiQuant program [41, 42], using experimentally determined photo-ionization cross-section data of Evans et al. [43] and asymmetry parameters of Reilman et al. [44]. Surface composition was calculated by the “infinitely thick homogeneous sample” and layer thickness by the Layers-on-Sphere models. Chemical states of the constituent elements were identified and assigned by reference data-bases and publications [45, 46].

3 Results

3.1 Thermodynamic calculations

We calculated the thermodynamic equilibrium concentrations of the reactant and products using FACTSAGE [47] thermodynamic software based on the Gibbs free energy minimization method. The calculations were performed for atmospheric pressure in the 500–5500 K temperature range for the following systems:

1. ZrO$_2$ + 3C + Ar (in excess),
2. ZrO$_2$ + 2CH$_4$ + Ar (in excess),
3. ZrO$_2$ + 3CH$_4$ + Ar (in excess),
4. Zr(OCH$_2$CH$_2$)$_2$ + Ar (in excess),
5. Zr(OCH$_2$CH$_2$)$_2$ + 15N$_2$ + Ar (in excess).

Since Ar is not taking part in the plasma reactions, it will not be indicated in further descriptions.

During the evaluation of the results, it must be kept in mind that due to the short residence time of the species in the plasma, thermodynamic equilibrium might not be achieved. The results, however, allow to estimate the potential type and concentration of products with the assumption that the precursors evaporate and decompose completely (atomize) in the high-temperature region of the plasma and reaction occurs in the gas phase. Temperature distribution and reagent trajectories (particles, molecules, atoms, radicals, etc.) in the reactor are usually based on, somehow simplified, mathematical simulations [48, 49]. Another hypothesis is that the reaction product formed first can be considered as the final one as further reactions are kinetically hindered at lower temperature regions because of the short dwell time.

Fig. 1 (a) shows the equilibrium composition as the function of temperature for the case of ZrO$_2$ and C precursor with 1:3 molar ratios. The main species in the gas phase at 3000–5500 K temperature range are atomic Zr, C, O, small carbon clusters (C$_n$, C$_x$) and oxygen-containing molecules such as ZrO. Over 3000 K temperature ZrO$_2$ is found practically in a reduced vapor phase, only Zr, C and CO are present in a considerable amount thus between the Zr and C reaction is very likely to occur in the gas phase. The first condensed phase is ZrC that forms below 3500 K, followed by carbon and ZrO$_2$ below 2000 K.
Thermodynamic equilibrium conditions of ZrO$_2$ and 2CH$_4$ system (Fig. 1 (b)) are quite similar to the former one regarding the prevailing species above 2000 K. Due to methane addition atomic H and H$_2$ molecules also appears at temperatures above 1000 K. Zr-bearing gas-phase species found at the high temperatures (above 3000 K) are atomic Zr, and in a small extent ZrO and ZrH molecules. The first condensed state again is ZrC (below 3500 K), while ZrO$_2$ and carbon are developed only below 2000 K.

Fig. 1 (c) illustrates the thermodynamic equilibrium composition for NZP. There are some significant differences comparing to former systems. First, NZP has much lower evaporation and decomposition temperature than ZrO$_2$ thus once introduced in the high-temperature zone of the plasma complete evaporation and decomposition can be assumed. Furthermore, molar ratios of O, C and H to Zr are considerably higher than in the previous systems. In other words, carbon is provided in a great excess that facilitates the formation of ZrC. The prevailing species are quite the same as compared to former described systems, but their ratio reflects the increased molar ratio of C and O to Zr in the NZP molecule. Accordingly, H$_2$ molecules are present at a higher concentration, while small hydrocarbons such as C$_2$H$_2$, C$_2$H and CH also appear in the gas phase at a temperature already above 2000 K. Carbon is available in the form of atomic carbon and small carbon clusters (C$_2$, C$_3$) in a wide temperature range. The majority of the oxygen is irreversibly bound to carbon as CO. When CO is formed in this system, it behaves essentially as an inactive gas, and these carbon and oxygen are unavailable for further reactions. The condensed phase below 3000 K comprises ZrC and carbon as well, while ZrO$_2$ is formed below 2000 K.

The addition of nitrogen to NZP (Fig. 1 (d)) diversifies the equilibrium compositions in the studied temperature range. Greatly increases the number of species with nitrogen-containing molecules such as N, CN, (CN)$_2$, HCN, NH, NO above 2500 K and NH$_3$ at temperature below 2000 K. Even a small amount of ZrN (g) is present in the gas phase at high-temperature range (4500–5500 K). Nevertheless, it does not appear among the condensed phases. Similar to former systems, the first condensed phase is ZrC with a considerable amount of solid carbon, too.
Results of the thermodynamic calculations (Fig. 1 (a–d)) show that solid ZrC could be formed over a wide temperature range from 1700 to 3500 K. In each case ZrC is the first solid phase formed at or below 3500 K. However, for ZrO$_2$ + 2CH$_4$ and ZrO$_2$ + 3C systems, the optimum temperature range for ZrC formation is narrower (2100–2800 K) than in case of the NZP system (1900–3600 K). It can be assumed that in each investigated system at higher temperatures, ZrC is the thermodynamically favored state. However, when the temperature of the system falls below 2000 K ZrO$_2$ becomes thermodynamically favored. ZrC must be formed and condensed before the temperature falls below the aforementioned temperature.

The results of thermodynamic equilibrium calculations have shown that the complete reduction of the NZP or ZrO$_2$ to ZrC can be achieved in the Ar, N$_2$-Ar, and methane–Ar mixture.

Thermodynamic computations certainly provide information about equilibrium states, but they did not take into consideration the kinetic limitations. The extremely high temperatures, steep gradients and the rapid cooling create unusual process conditions which, in some circumstances, can give rise to significant deviations from the calculated equilibrium [50, 51]. In the plasma synthesis process, the typical residence times of particles in the plasma are in the order of several hundred ms [48], consequently, the time for condensed phase reactants evaporation and gas-phase chemical reactions is very short. Due to this short residence time of the reactants in the plasma flame, many chemical reactions may not reach the equilibrium state. Nevertheless, these investigations may give hints of the thermal prehistory of the precursors in the plasma.

### 3.2 Powder characterization

In Table 2, the semi-quantitative composition, based on the XRD peak intensities [52], the average crystallite size of ZrC determined from the broadening of XRD peaks and particle sizes calculated from the specific surface area values of the prepared powders are presented.

The mean sizes of ZrC particles were calculated from the values of the specific surface areas (SSA) of the product powders, assuming dense spherical shape (Table 2). The SSA varied in the particular Runs within the range of 10–100 m$^2$g$^{-1}$. The mean size was estimated to fall between 11 nm and 79 nm.

#### 3.2.1 Crystallographic phase relations

X-ray diffraction analysis was performed on the plasma synthesized samples to investigate the phase composition and the crystallite size.

In Fig. 2 the typical X-ray diffraction patterns of some selected plasma products synthesized from different precursors (Runs 1, 3, 5, 6) are illustrated.

Using ZrO$_2$ precursor (Run 1–4), the product powder is mainly a mixture of carbon, monoclinic (m-ZrO$_2$) and tetragonal (t-ZrO$_2$) zirconia, while reflections of lower intensity also indicate the presence of ZrC. Peak corresponding to graphitic carbon can be detected regardless graphite or methane was used in the reactions as carbon source. The presence of zirconia in the product implies that a considerable amount of the zirconia particles just travel through the plasma flame without evaporation or getting reacted with the carbon. The zirconia precursor is composed of both monoclinic and tetragonal crystalline phases in nearly equal ratio. The intensity of m-ZrO$_2$ and t-ZrO$_2$, however, slightly decreases with the increase of carbon. In the case of methane reduction agent, no m-ZrO$_2$ peaks reflections were observed.

It was found in several earlier papers [9, 53] that synthesis performed in oxygen-rich environment oxygen easily lead to the formation of oxycarbide (ZrO$_2$C$_x$) phases, in which oxygen substitutes some part of carbon atoms in the ZrC lattice. According to the XRD diffractograms, however neither separate reflections nor expansion/distortion of the ZrC reflections could be observed, which would correspond to the formation of oxycarbide face in our experiments.

To reduce the residual ZrO$_2$ content in the product, in Runs 2 and 4 excess carbon was applied. According to thermodynamic calculations, higher molar ratio of the

| Run | Precursors molar ratio | Carrier gas | Phase composition of the product (wt%) by XRD | ZrC crystallite size (nm) from XRD | dBET (nm) |
|-----|------------------------|-------------|---------------------------------------------|----------------------------------|----------|
| 1   | ZrO$_2$ + 3C           | Ar          | ZrC 41, ZrO$_2$ 53                          | 6                                | 37       |
| 2   | ZrO$_2$ + 5C           | Ar          | ZrC 39, ZrO$_2$ 48                          | 13                               | 29       |
| 3   | ZrO$_2$ + 3CH$_4$      | Ar          | ZrC 48, ZrO$_2$ 49                          | 3                                | 25       |
| 4   | ZrO$_2$ + 4CH$_4$      | Ar          | ZrC 51, ZrO$_2$ 42                          | 7                                | 23       |
| 5   | Zr(OCH$_2$CH$_2$CH$_3$)$_2$ | Ar  | ZrC 98, ZrO$_2$ 2                           | -                               | 17       |
| 6   | Zr(OCH$_2$CH$_2$CH$_3$)$_2$ | N2      | ZrC 95, ZrO$_2$ 5                            | -                               | 19       |
carbon among the reactants increases the ZrC content of the product as well. However, it was not the case in our experiments. The addition of excess carbon did not promote the ZrC formation versus ZrO$_2$ reduction rather only increase the solid carbon phase formation. Although thermodynamic calculations showed that reduction of ZrO$_2$ to ZrC can be achieved in ZrO$_2$ + 3C or ZrO$_2$ + 2CH$_4$ mixtures, XRD analysis indicates that only a part of the starting oxide was converted to ZrC. This observation can be explained by the short residence times of ZrO$_2$ particles in the plasma, being not enough for complete evaporation and decomposition for an in-flight reaction.

In Runs 5–6 using a NZP precursor, ZrC becomes the predominant phase. The sharp lines and absence of impurity peaks in the XRD pattern indicate the formation of the crystalline ZrC phase (Fig. 2). It has been reported in the literature that non-stoichiometric ZrC$_{1-x}$ could be formed in the Zr–C binary system [54, 55]. In our case, the calculated lattice parameters for ZrC powders are slightly higher (4.699 Å) than the standard value 4.693 Å (PDF#350784), however, in all experiments (Run 1–Run 6) the lattice parameter displays the same value. All the intense peaks correspond to face-center cubic (fcc) ZrC in agreement with the reference data indicating high structural purity of ZrC.

The addition of nitrogen in the plasma resulted in the same phase composition: ZrC remained the dominant phase in the product, however its amount was slightly decreased and a small increase of the amount of ZrO$_2$ was observed in Run 6 as compared to Run 5. The difference could be attributed to higher oxygen contamination of the plasma gases.

The thermodynamic calculation predicts that there is a strong tendency for the system (Runs 5 and 6) to form carbon in addition to ZrC over a wide temperature range. Starting from the NZP precursor, based on these calculations, carbon formation is expected in somewhat larger (Run 5) or lower (Run 6) quantities. However, in XRD patterns of these experiments the carbon phase could not be detected among the crystalline phases. Even when a carbon deposit has not been reported, it may be present in the product in the amorphous state [56]. Plasma processing of various aliphatic alcohols in reducing conditions can results in graphene layers instead of well crystallized carbon structure [57–58].

The average crystallite size of ZrC particles determined by the Scherrer equation based on the broadening of the reflections was found to be between 17 and 37 nm respectively. The highest value, with 37 nm, was found in the reaction of ZrO$_2$ + 3C (Run 1) and the lowest, with 17 nm size, was found in the case of NZP precursor (Run 5).

### 3.2.2 Particle morphology

The TEM and the SEM image in Fig. 3 show the microstructure of the plasma synthesized product powder obtained from the ZrO$_2$ + 2CH$_4$ reaction (Run 3). SEM image (Fig. 3 (a)) suggests a significant heterodispersity in the particle size. Some large spherical particles with a mean particle size of about 10 μm and aggregates of sub-micron and more frequently, of nanometer size particles could be observed.

![Fig. 3 SEM (a) and TEM (b) micrographs of the sample from ZrO$_2$ + 2CH$_4$ precursors](image)

**Fig. 2** X-ray diffraction patterns of plasma products from ZrO$_2$ + 3C, ZrO$_2$ + 2CH$_4$, NZP and NZP + 15N$_2$ precursors, respectively.
EDX analysis revealed that the larger spheres are ZrO$_2$ particles, while the aggregates are composed of zirconium and carbon.

The larger zirconia particles represent the unreacted part of the zirconia precursor. Theoretically, the precursor ZrO$_2$ particles passing through the plasma flame should evaporate. However, part of them neither evaporate nor dissociate due to the insufficient dwell time in the hottest zone of the plasma. The appropriate residence time varies depending on both the size and the actual thermal trajectory of the individual particles. The spherical shapes of the particle suggest, however, that it was also affected, melted or partially evaporated in the plasma. The small size and the relatively smooth surface indicate also significant recrystallization of the precursor ZrO$_2$ particles. ZrC is formed by homogenous nucleation from the gas phase. Due to high cooling rates characteristic for in-flight plasma conditions, the nucleated particles do not have the chance to grow further but tend to deposit on the reactor wall or fly away with the gas stream into the filter unit.

Although the SEM images of the products from the ZrO$_2$ precursor show similar morphology (Fig. 3 (a) and Fig. 4 (a)), in TEM micrographs, we observed differences. The TEM image of the ZrO$_2$ + 3CH$_4$ reaction (Fig. 3 (b)) shows the presence of dark 10–50 nm size grains embedded in a light contrast, presumably graphitic material. The electron diffraction pattern corresponds to crystalline ZrC.

TEM micrograph of the product resulted from the ZrO$_2$ + 3C reaction (Fig. 4 (b)) is in accordance with the XRD results showing three different phases: the darkest crystallites could be assigned to ZrC, the more transparent grains to the ZrO$_2$ phase, while the lightest ones to carbon. This is made on the basis of the different absorption coefficients of electrons increasing with the concentration of the high atomic number element, i.e. to the zirconium content of the phase.

The smallest mean particle size of 11 nm was obtained for the sample prepared from NZP precursor (Run 5). TEM micrographs (Fig. 5) of this sample fully confirm the above cited small size of mainly equiaxed particles. The mean size of ZrC particles were calculated from the values of the specific surface areas (SSA) of the product powders assuming dense spherical shape (Table 2). Although this assumption may not be completely satisfied, the obtained results are in agreement not only with SEM and TEM analysis (Figs. 3–5) but also with the average crystallite size of ZrC particles calculated from XRD patterns.

3.2.3 Raman spectroscopy

Raman spectra were recorded only for samples prepared from NZP precursor (Run 5, Run 6) and shown in Fig. 6. According to previous studies [59], stoichiometric ZrC, due to its perfect NaCl structure, shows no peaks for Raman spectra, meanwhile for nonstoichiometric ZrC$_{1-x}$ phonon optical and acoustic branches associated to the presence of carbon vacancies were observed in the range...
of 200–300 cm\(^{-1}\) and 500–600 cm\(^{-1}\). Peaks corresponding to ZrC\(_{1-x}\) did not occur in our spectra, thus these low-frequency ranges are not depicted in Fig. 6. However, as no peaks corresponding to ZrC\(_{1-x}\) were detected the synthesized ZrC can be considered to be stoichiometric.

The spectra show intensive bands at ~1350 cm\(^{-1}\), ~1580 cm\(^{-1}\) and ~2700 cm\(^{-1}\), characteristic for carbon. The peak at 1588 cm\(^{-1}\) is conventionally referred as the G mode and assigned to the C–C stretching vibration (E\(_{2g}\)) of graphite, while the peak at 1340 cm\(^{-1}\) is called disorder-induced D-band. The peak observed at 2684 cm\(^{-1}\) originates from a second-order process and is referred as G’ band.

The intensity of the D and G and also the G’ peaks decrease significantly with the addition of N\(_2\), indicating the decrease of residual carbon in the powders. At the same time the decrease of the graphitic peak was more pronounced relative to the D band.

As it was emphasized, till the D to G band ratio of I(D)/I(G) represents the concentration of defects in graphite [1], the number of graphitic or graphene layers can be associated with the FWHM of the G’ band and the I(G’)/I(G) ratio [32]. It was assumed that the number of graphitic or graphene layers decreases when the I(G’)/I(G) ratio increases and the FWHM(G’) decreases [32, 60].

For sample Run 5, the I(D)/I(G) ratio is 0.79, FWHM(G’) is 69 cm\(^{-1}\) while the ratio of I(G’)/I(G) is 0.51. For Run 6 these values are: 1.56, 56 cm\(^{-1}\) and 0.73, respectively. The degree of structural order, which is related to the ratio of I(D)/I(G), is higher in sample of Run 5 as compared to Run 6 when nitrogen was added to the system.

Evaluation of the number of graphene layers based on reported estimation [32, 60], is found to be more than 10 layers for Run 5, while for Run 6 turned to be between 5 and 10 layers, only. These small values are in full agreement with the TEM observations about the presence of transparent carbon layers. The presence of excited N\(_2\) atoms seems to partially inhibit the formation of the well-ordered graphite phase.

### 3.2.4 Surface analysis

The surface characteristics of the as-prepared powder samples were characterized by XPS.

Besides the expected Zr and C, oxygen was also detected. All constituents are present in several chemical states, as illustrated by the decomposed Zr3d and C1s spectra in Fig. 7. Zirconium is found in carbide, Zr3d5/2 179.4 eV, and in oxide, Zr3d3/2 182.8 eV, states. The line positions of the corresponding carbon, C1s 182.0 eV, and oxygen, O1s 531.2 eV, are in agreement with the literature data [45]. The appearance of the ZrO2 minor phase can be attributed to the high affinity between Zr and O [61, 62]. Part of the oxygen is chemically bound to carbon, O1s 533.0 ± 0.2 eV, C1s 286.7 ± 0.2 eV. In the case of sample Run 4, due to the separate carbon particles (as revealed by electron microscopy), differential charging can be observed, causing an extra C1s component (peak labeled C* in Fig. 7). The majority of the
carbon is in elemental state, C1s 284.6 ± 0.2 eV, forming together with the oxidized carbon the "carbonaceous" contamination on the samples.

The surface chemical compositions are shown in Table 3. Although the composition is calculated by the Homogeneous model, these samples are not homogeneous thus these values can be regarded semi-quantitative. A large amount of excess carbon is inevitably demonstrated. The ratios of the constituents of the main components (ZrC, ZrO2) are quite close to their expected values (1:1, 1:2).

For the NZP + Ar and NZP + 15N2 samples, the level of the carbonaceous contamination is significantly lower and no separate carbon particles detected by SEM. Supposing that the oxide covers directly the carbide particles then the carbonaceous layer is formed on the surface of nanoparticles (due to decomposition of the hydrocarbons or propoxide, which are present in the gas phase), the thickness of these layers can be calculated. The results are presented in Table 4. The thickness values are in harmony with the composition data. For NZP + Ar the carbon layer is thicker while the oxide layer is thinner, comparing to NZP + 15N2.

Assuming that the as-calculated carbon layer is actually made up of graphene sheets, and evaluating the number of graphene layers appeared as 12 layers for Run 5, while for Run 6 as 10 layers (calculated with 0.33 nm distances between the layers), which are in excellent agreement with the values derived from Raman spectra.

Nanometer-sized ZrC particles have poor oxidation resistance. However, oxidation resistance can be improved by means of coating [1]. Graphitic carbon-coated nanoparticles are intensively studied worldwide, expecting that the carbon shell in these structures would be able to protect the inner core materials, i.e., it can protect ZrC nanoparticles in oxidative environment (Runs 5 & 6 in the present study) [1].

However, when nanopowder is used as precursor for a sintered body, the presence of free carbon on the surface might prevent sintering. Heat treatment in hydrogen atmosphere would be required to prepare carbon free ZrC phase.

4 Conclusions

- Feasibility of preparation of nanosize ZrC in thermal plasma from ZrO2 reacting with carbon or CH4 and by thermal decomposition of NZP using high-temperature (HT) atmospheric plasma conditions is presented.
- Thermodynamic calculations predict the formation of solid ZrC in certain temperature ranges for all of the selected precursors and their relative proportions. The common temperature range of solid ZrC formation for each explored reaction system may be between approx. 2000 and 3000 K, what is in the cooling tail of the plasma flame. In agreement with the thermodynamic calculations, XRD results confirm the presence of crystalline ZrC in all prepared samples. From here, it became evident that there must be a somewhat similar thermal prehistory of the reaction partners in the conducted experiments.
- Nanometric size powders were produced in each experiment, although with different mean particle size and chemical composition. The highest yield of 98 % was achieved when nanopowders were prepared by the decomposition of NZP, while ZrO2 reacted with C resulted in the lowest yield of 39 %. In the case of ZrO2, the starting oxide could only be partially converted to ZrC at the applied reaction conditions.
- NZP was completely converted to crystalline ZrC, this implies that the four Zr-O bonds are fully dissociated, and the Zr reacted with the carbon-containing species evolved from the propoxide part of the molecules. Only negligible amount of ZrO2 appeared in the resulted powder. The excess carbon from the propoxide chain form solid carbon.
- The particle size of the prepared samples falls in the range of 10–100 nm depending on the type of precursors. The smallest mean particle size, 11 nm, was achieved with the NZP precursor. Important to note that very similar values were obtained for each type of sample, notwithstanding they are calculated from the specific surface area or from the broadening of the X-ray diffraction reflections, and also from the visual observation by TEM. Besides, this fact implies that all of these HT-plasma deposited

| Table 3 | Surface composition (atomic %) of the various samples calculated by the Homogeneous model |
|----------|-----------------------------------------------------|
| Precursor | Zr C | Zr O | O | C | | | |
| NZP + 2CH4 | 0.7 | 0.7 | 0.5 | 1.5 | 89.2 | 5.9 | 1.6 |
| NZP + Ar | 6.5 | 5.8 | 4.3 | 7.0 | 61.7 | 11.0 | 3.8 |
| NZP + 15N2 | 5.5 | 5.6 | 6.7 | 12.8 | 54.1 | 8.6 | 6.7 |

| Table 4 | Thickness (nm) of the carbonaceous and oxide layers on the Zr carbide particles, calculated by the Layers-on-Sphere model from XPS data |
|----------|-----------------------------------------------------|
| Precursor | CHx layer (nm) | ZrO2 layer (nm) |
| NZP + Ar | 4.2 | 1.0 |
| NZP + 15N2 | 3.5 | 1.6 |
samples are rather compact, not having internal pores with extended surface area, and also that the second solid phase, carbon, does not affect those measured particle size values significantly.

- XPS characterization revealed that even the structurally pure ZrC samples, derived from the NZP, contain some amount of oxygen and carbon on their surface. Supposing an oxide and carbonaceous layers on the surface of the spherical ZrC nanoparticles, an ≈1 nm oxide and a ≈4.1 nm thick carbon layer was calculated from Run 5 sample, but somewhat thicker oxide but thinner carbon layer was obtained for Run 6 when nitrogen was added to the process gas.

- Based on the above mentioned results, the process requires optimization with ZrO₂ precursor towards completeness, and with NZP, towards the elimination of the excess solid carbon, e.g. by gasification (into CO₂), if the further application requires it. However, considering some practical aspects of these results, the nanometer range particle size of the samples, synthesized by such HT-plasma technique, is beneficial in the process of hi-tech ZrC ceramics.

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