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Open-celled silicon carbide foams with high porosity from boron-modified polycarbosilanes

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\textbf{A B S T R A C T}

Open-Celled silicon carbide (SiC) foams were prepared from a mixture of a boron-modified polycarbosilane as a preceramic polymer and poly(methylnorprene) (PMMA) microbeads as sacrificial agents. The process consists in the cross-linking of the liquid allyhydridopolycarbosilane (AHPCS, SiC precursor) with borane dimethylsulphide (BDMS, boron source) to form a solid boron-modified polycarbosilane with an adjusted cross-linking degree. The latter is mixed with PMMA microbeads (25 μm) in a 20:80 ratio and the mixture is warm-pressed at 120 °C forming consolidated green bodies to be pyrolyzed at 1000 °C under argon and to deliver open-celled SiC foams with an interconnected porosity of 73.4 \textpm 2%. These foams combine a low density with a compressive strength of 3.49 ± 0.56 MPa and a thermal and mechanical stability under argon up to 1300 °C.

1. Introduction

Porous solids possessing a large amount of internal surface area are suitable for performing specific functions for a large variety of applications related to separation techniques, i.e., especially for energy and environment-related fields, that require fast and efficient transport in, out, or within a high porosity. Their performance and function are governed by the class of materials they belong to, i.e., polymers, metals, ceramics, the chemical composition of the considered porous scaffold material, and the porosity characteristics (pore shape, average pore size and size distribution, pore interconnectivity and tortuosity).

An appropriate classification of the porosity is based on the investigations made by Dubinin [1] and adopted by Everett [2] then Sing et al. [3]. It consists in classifying pores as macropores ( > 50 nm, sintered ceramics, foams, ...), mesopores in the range of 2-50 nm (mesoporous silica and carbon, ...) and micropores < 2 nm (zeolite, activated carbon, metal organic framework, membranes, ...) [4]. In such systems, (large) macropores are ideal for providing easy access to internal sites, while micro- or mesopores increase the surface area and therefore improve selectivity [5,6].

The development of porosity in ceramics extends the functionality of the latter toward applications where a defined permeability for various fluids is required, especially at high temperature [7]. A typical example is the Diesel Particulate Filter (DPF) usually made of silicon carbide (SiC), mainly used for the safe and effective reduction of Diesel engine exhaust gases and soot [8].

The porosity characteristics, thus the production of highly porous ceramic components such as SiC, are all determined by the selected processing strategies. In particular, designing porosity from nanoscale to microscale is a unique attribute of the chemical material technologies. Among these technologies, the chemical approaches based on tailored preceramic polymers, such as the polymer-derived ceramics (PDCs) route, offer a precise control over chemical compositions, microstructure and porosity characteristics at different length scales for ceramics.

The PDCs method is based on the synthesis of a preceramic polymer as a precursor [9–14]. Its main purpose is to produce ceramic components, often as non-oxide, through the thermo-chemical conversion of...
preceramic polymers via shaping (to form the desired component), curing (to further crosslink the polymer), pyrolysis (to transform the cured polymer into an amorphous ceramic) and optionally annealing (to crystallize the amorphous ceramic). Interestingly, because of the possibilities of controlling the cross-linking degree, the type of bonds linking monomeric units and the nature of functional groups in preceramic polymers, polymer shaping (before or during curing) can be carried out to produce thin parts or complex architectures such as fibers, coatings, or dense monoliths [15–17]. If the PDCs route is coupled with a porous network design method, at the polymer level, a control of the porosity can be achieved at various length scales [18–22]. Based on the advantages provided by the PDCs route, the processing of open-celled SiC foams has been exploited in the present paper.

An approach to prepare open-celled foams consists in using sacrificial agents serving as placeholder substances to create open pore networks in the final ceramics [21–26]. Such agents mainly include organic materials that decompose without leaving residues during heating, such as poly(methyl methacrylate) (PMMA) beads, and that can be directly mixed with the SiC precursors. The selection of the latter is important: because uniaxial warm-pressing is a typical forming method to shape mixtures of preceramic polymers and PMMA and deliver highly porous non-oxide ceramics in monolithic form after pyrolysis [22,23], an ideal open-celled SiC foam precursor would display a relatively high crosslink degree to be processed into pellets by warm-pressing while plastic deformation then precursor particle joining occur, leading to insulable and consolidated green bodies. Subsequent pyrolysis should retain the shape of the material (i.e., controlled volume shrinkage) and lead to open-celled PDC foams (if the PMMA beads mixed with the ceramic precursor before warm-pressing are embedded in an amount large enough to create an interconnecting network), through the conversion of the precursor into ceramics, while PMMA is removed to leave voids, i.e., macro pores.

The pioneered commercially available polycarbosilane (PCS), nominally [SiH(CH)=CH2]n, obtained upon thermolysis of polysilanes (PS) [27,28], is a solid preceramic polymer displaying a relatively high ratio of C to Si (2:1) and a complex structure which appears to contain largely carbosilane linkages. PCS is melt-spinnable and led in the middle of the seventies to the achievement of the 1st generation of SiC fibers called Nicalon™ [27,28]. However, the preparation of other geometries than fibers is difficult because of the low ceramic yield of PCS and the strong shrinkage occurring during its pyrolytic conversion. Moreover, the compositional purity of the SiC ceramic produced by this process is limited by the lack of cross-linking moieties in the precursor, which requires oxidative curing to generate a non fusible polymeric body. As a second commercially available PCS precursor, allyldihydridopolycarbosilane (AHPCS), nominally [Si(CH2=CH2)=CH2]n HCH32,nSiH2CH3,a [29] is produced by the Starfire Systems company [30]. It is a liquid and can produce SiC with an almost stoichiometric C:Si ratio. It has been previously investigated to produce porous SiC through its combination with PE [26] and PMMA [31] microbeads. The latter reported the mixing of AHPCs and PMMA microbeads in a slurry to be casted and then pyrolysed at 900 °C, forming foams. As a proof of concept, a thick disc-shaped porous ceramic body was successfully produced using 68 w% of PMMA beads (90 μm diameter), after pyrolysis at 900°C. No detailed information was provided about the morphological, mechanical and thermal characteristics of the materials. The major issue of AHPCS is its relatively low degree of crosslinking and the lack of low temperature cross-linkable functional groups, both involving the releases low molecular weight species such as oligomers in the early stages of the pyrolysis. This inherently reduces the ceramic yield while increasing the volume shrinkage, making the process difficult to scale up. Furthermore, using liquid AHPCs and solid sacrificial agents, it is really challenging to obtain highly porous components (porosity above 70 vol%) with a high degree of pore interconnectivity.

For further processing AHPCS into open-celled SiC foams, a solution would consist in firstly cross-linking the liquid AHPCS, allowing its transformation into rigid solid-state materials that could be then homogeneously mixed with PMMA microbeads. Thus, the latter will exhibit a higher ceramic yield to form open-celled SiC foams with a reduced volume shrinkage. This strategy has been demonstrated to prepare dense SiC objects [32]. However, this requires relatively high cross-linking temperatures (T° approaching 250 °C) and uniaxial warm-pressing should be performed at a significant higher temperature (T° > 250 °C). Furthermore, it is difficult to precisely control the cross-linking evolution via the temperature (and the time at the set temperature), and the resulting polymers can be extensively crosslinked and therefore no longer processible.

To form a solid-state SiC precursors with a tuned plasticity at relatively low temperature, we modified AHPCS with boron dimethylsulphide to synthesize a boron-modified AHPCS with a tailored cross-linking degree according to a strategy reported recently by our group on various types of commercially available preceramic polymers [33–37]. Thus, this strategy allowed us to produce open-celled SiC foams containing free carbon and a low boron content by warm-pressing and pyrolysis. Carbon and boron additives are often used as sintering aids during the densification of SiC, based on the fact that carbon inhibits the in situ effective mass transport while boron favors diffusion and hence the ceramic densification [38]. Based on this strategy, we report, in the first part of the paper, the design of open-celled SiC foams by mixing the boron-modified AHPCS with PMMA microbeads of 25 μm in diameter. As-synthesized materials were characterized in detail by elemental analyses, SEM, mercury porosimetry and mechanical tests. To identify the potential of using these materials in harsh environments, their behavior at high temperature was investigated via HT-TGA, SEM, mercury porosimetry and BET. Then, aging as well as microfiltration tests were performed to simulate the conditions of a primary loop of coolant in a Pressurized Water Reactor (PWR). The schematic representation of the synthetic process is shown in Fig. 1.

2. Experimental procedure

For all the synthesis reactions, a ventilated hood equipped with a vacuum/argon line allowed to manipulate the chemical products under argon atmosphere. Argon (≥ 99.995%) was purified by passing through a column of phosphorus pentoxide (Sigma-Aldrich, Saint Quentin, France), sacipent™ (Millipore S.A.S, Molsheim, France) and copper oxide-based catalysts (Sigma-Aldrich, Saint Quentin, France). The glassware was stored in an oven at 120 °C overnight before being assembled and pumped under vacuum for 30 min and then filled with argon. In order to keep inert conditions after the synthesis reactions, the manipulation of the chemical products was made inside argon-filled glove boxes (MBraun MB200B; O2 and H2O concentrations kept at ≤ 0.1 ppm and Jacomex, Campus-type; O2 and H2O concentrations kept at ≤ 0.1 ppm and ≤ 0.8 ppm, respectively). Allyldihydridopolycarbosilane (AHPCS labeled SMP-10, Starfire Systems® Incorporation, New York, USA), with a density of 0.998 g·cm−3, was used as-received. Toluene (99.85%, Extra Dry over Molecular Sieve, AcroSeal(R)) was obtained from Acros Organics. Borane dimethylsulphide BH3 Si(CH3)2 (BDMS complex solution 2.0 M in toluene) was obtained from Sigma Aldrich. Poly(methylmetacrylate) (PMMA) microbeads with a diameter of 25 μm were provided by Atotina Italia, Milan, Italy.

2.1. Material preparation

Open-celled SiC foams were produced from a boron-modified allyldihydridopolycarbosilane labelled AHPCSB30 (30 being the Si:B ratio fixed during the polymer synthesis) which was synthesized by reaction between AHPCS and BDMS according to a previously published report [39]. AHPCSB30 was ground in an agate mortar, then it was mixed with 25 μm PMMA microbeads in AHPCSB30:PMMA weight ratios.
ranging from 10:90 to 30:70. The mixture was homogenized in a mortar and the mixture was then pressed at 74 MPa, in a steel die with diameter 13 mm with a heating mantle, from RT to 120 °C, using an automatic uniaxial press (Atlas Auto model, Specac) set-up in a nitrogen-filled glove-box. We also investigated a higher steel die diameter (25.6 mm) for further scaling-up. The homogeneous mixture was introduced in the steel die, put under load and heated to the desired temperature of warm-pressing, then the temperature was held for 0.5 h. The samples were slowly de-loaded after cooling down to RT (within 1 h). Typical dimensions of the cylindrical green bodies were 13 mm × 1.6 mm (Please note that compressive strength measurements required a sample thickness of 4 mm; thus, we used a higher quantity of mixture for warm-pressing to perform mechanical measurements). Green samples were placed in alumina boats, and then transferred under argon flow into a silica tube inserted in a horizontal furnace (Carbolite BGHA12/450B). The tube was pumped under vacuum (1.10⁻¹ mbar) and refilled with argon (99.995%) at atmospheric pressure. Subsequently, the samples were subjected to a cycle of ramping of 0.5 °C.min⁻¹ to 450 °C (dwell time of 2 h) then to 1000 °C (dwell time of 2 h) to form mSiBC30_10 samples. A constant flow of argon (120 mL.min⁻¹) was passed through the tube during the pyrolysis cycle. In order to investigate the high temperature behavior of the mSiBC30_10 samples, annealing treatments were performed above 1000 °C in a graphite furnace (Gero Model HTK 8). The furnace was pumped under vacuum (1.10⁻¹ mbar), refilled with argon and maintained under a constant flow of argon (200 mL.min⁻¹) during the whole heat treatment. The program consisted of a 5 °C.min⁻¹ heating ramp up to the maximum temperature fixed in the range 1000–1500 °C, dwelling at this temperature for 2 h, and cooling down to RT at 5 °C.min⁻¹. The samples were labelled as mSiBC30_X, X being the two first digits of the temperature at which the open-celled SiC foams were heated, e.g., X = 13 for 1300 °C.

2.2. Material characterization

AHPCS830 was fully characterized in our previous report [33]. High-temperature thermogravimetric analysis (HT-TGA, Setaram Setsys 2400 CS evolution equipment) of the foams prepared at 1000 °C was performed in flowing argon from 25 to 1500 °C at a heating rate of 5 °C.min⁻¹ using tungsten crucibles. X-ray diffraction was carried out from 20 to 80° with a step size of 0.0167° using a Philips PW 3040/60 X’Pert PRO X-ray diffraction system operating at 30 mA and 40 kV to identify the crystalline phase assemblage of the samples. Samples were also investigated by SEM (Hitachi S4800) coupled with EDX to assess the morphology of the samples while measuring the Si, C and O content in materials. ICP-OES (Perkin Elmer Optima 7000 DV) was performed at the IUT Montpellier-Sète (France), N₂ adsorption-desorption isotherms were carried out using a Model Sorptomatic 1900 analyser (Fisons); prior to the analysis samples were degassed overnight at 150 °C. The Brunauer–Emmett–Teller (BET) method was used to calculate the specific surface area of samples. A Micromeritics Autopore IV 9500 mercury porosimeter was used to investigate the pore size distribution of the samples. Compression tests were carried out on espe-cially designed cylindrical samples with 10 mm diameter and 4 mm height using a universal testing machine (Instron 1121 UTM, Instron Danvers, MA) at a cross head speed of 1 mm/min; at least three samples were tested for each data point, and values are reported as mean ± standard deviation. The aging tests of foams including mSiBC30_10 and mSiBC30_13 samples were performed in six micro-autoclaves made of Inconel 600 arranged set-up in an oven at 330 °C under 140 bar and containing demineralized water with 2.24 ppm (mg.L⁻¹) of lithium and 590 ppm of boron at a pH of 6.8 to reproduce the working conditions of a primary loop of coolant [39] (Fig. 2).

In parallel, the microfiltration capacity of mSiBC30_10 samples was tested for 220 h by using a closed system under 90 bar up to 275 °C, which was based on a filter cascade as illustrated in Fig. 2. The filtration system, as a dead-end filtration process, contained differential pressure detectors at each of the three consecutive filters with different porosity ranges [40,41]. The mSiBC30_10 samples were considered as the filter 1 with the largest porosity. The foam was inserted in a PTFE ring for sealing before being mounted between two stainless pieces. The system was then set-up in the chamber of the filter 1. The two other filters with lower porosity were commercially available. Such a system allowed for the in-situ measurement of the pressure upstream and downstream from the open-celled SiC foams, during filtration at a flow rate of 40 mL.min⁻¹ of demineralized water containing ethanolamine (3 ppm) and ammonia (1 ppm) as well as magnetite particles (around 5 μm in diameter and 10 g of particles for 50 mL of demineralized water), simulating the corrosion products coming from structural materials used in the primary loop of coolant.

Fig. 1. Overall synthetic path employed to generate open-celled SiC foams from boron-modified AHPCS. High magnification SEM image of the mSiBC30_10 samples as inset.
3. Results and discussion

3.1. Design and characterization of open-celled SiC foams

The modification of AHPCS by BDMS allows adjusting the degree of cross-linking of the derived polymer, i.e., AHPCS830 for solid-state processing. Thus, AHPCS830 is a solid compound with a ceramic yield as high as 91% [33] that could be mixed with 25 µm PMMA microbeads as a pore former to form homogeneously mixed powders with controlled AHPCS830:PMMA weight ratios. Powders could be then warm-pressed and pyrolyzed to generate foams. The 20:80 ratio was selected as the most appropriate AHPCS830:PMMA weight ratio, based on the fact that the corresponding samples did not collapse upon pyrolysis (as observed with samples prepared with 10:90 and 15:85 wt ratios) while being mechanically stable and presenting a high level of interconnected porosity after pyrolysis (samples prepared with the 30:70 wt ratio did not display a significant level of interconnectivity). The pyrolysis at 1000°C produced the mSiBC30_10 samples with an empirical formula of Si2.8B0.6C1.8O1.8. As-obtained samples are X-ray amorphous, based on the presence of diffuse SiC diffraction peaks in their powder X-Ray Diffraction (XRD) pattern, and contain free carbon, as indicated by the chemical composition and the Raman spectrum through the appearance of the D and G bands of free carbon (See Fig. 1S in ESI). As shown in Fig. 1, as-obtained materials reproduce the monolithic shape of the warm-pressed samples without the presence of any defects or cracks, whereas a volume shrinkage equivalent to 52.6% occurs (very close to bulk analogs). As shown in the high-magnification SEM image (as inset in Fig. 1), the mSiBC30_10 samples display a typical combination of spherical voids with a size controlled by the diameter of the PMMA beads. These macropores, homogeneously distributed within the bulk, contain a large amount of cell windows, allowing high interconnectivity among adjacent cells to the foams. In order to better qualify and quantify the macroporosity and pore interconnectivity of mSiBC30_10 samples, mercury intrusion porosimetry was carried out (Fig. 3).

During Hg porosity experiments, we observed a continuous increase in mercury uptake with the decrease in pore diameter within one intrusion step (i.e., increase of Hg intrusion pressure), as illustrated through the evolution of the cumulative pore volume versus pore diameter. This indicates a sharpening of the distribution of the cell windows in the samples. As a consequence, the log differential intrusion versus diameter, demonstrating the rate-limited filling of the pores by mercury under pressure, is represented by one signal with a shoulder on the left part from 5 to 7 µm allowing to range the size of the windows from 2 to 25 µm. The mSiBC30_10 samples exhibit a total porosity of 73.4 vol% and an intrusion volume as high as 1.47 cm³. g⁻¹. They display a bulk density as low as 0.51 g.cm⁻³ and a skeletal density of 1.92 g.cm⁻³. We confirmed such data by increasing the diameter of the foams to 20 mm (See Fig. 2S in ESI): A total porosity of 77.2 vol% and an intrusion volume as high as 1.66 cm³. g⁻¹ have been determined. They display a bulk density as low as 0.47 g.cm⁻³ and a skeletal density of 2.04 g.cm⁻³. According to the relatively high porosity, an average compressive strength of 3.49 ± 0.56 MPa was measured. Although it is challenging to speculate on these results, we can consider reports using a similar strategy to prepare foams. Based on an equivalent precursor (polysilazane):PMMA ratio (i.e., 20:80), C. Vakifahmetoglu et al. prepared Si(O)CN foams with an average compressive strength of 5.6 ± 0.7 MPa for a higher porosity (77.24 vol%) [23]. SiOC microcellular produced from a silicone resin using the same amount and size of PMMA microbeads had a compression strength after pyrolysis, of about 5.4 ± 0.9 MPa [42]. In contrast, using a physical blowing agent, the same authors produced oxygen-free SiCN foams with a porosity of 70.68 vol% and a flexural strength of 2.89 ± 1.04 MPa. Authors applied the model proposed by Gibson and Ashby [43] to discuss on the compressive strength of their samples, for which the validity of the model is appropriate (relative density lower than 0.3).
Thus, we can consider mSiBC30_10 samples as open-celled SiC foams and this is most probably the reason of their lower compressive strength compared to those measured for Si (O)CN foams using an equivalent precursor:PMMA ratio (i.e., 20:80).

### 3.2. High temperature behavior of open-celled SiC foams

The high temperature behavior of mSiBC30_10 samples was first investigated through high temperature TG experiments up to 1500 °C in an argon atmosphere (Fig. 4). As a comparison, the TG curve of the bulk analog labelled SiBC30_10, prepared by direct pyrolysis of AHPCS830 in an argon atmosphere at 1000 °C was reported [33]. Then, complementary characterization experiments of samples pyrolyzed at different temperatures in the temperature range 1000–1500 °C were carried out to follow the evolution of the porosity characteristic and the mechanical behavior of open-celled SiC foams at high temperature.

As expected, the mSiBC30_10 samples do not undergo any weight loss after heat-treatment to 1000 °C; temperature at which they were prepared. Then, a two-step weight loss occurs: a first and very small weight loss (0.25%) was measured in the temperature range 1000–1300 °C and a second one, more significant (3.75%), was then identified from 1300 to 1500 °C. Thus, a total weight loss of 4% was measured at 1500 °C for mSiBC30_10 samples. It should be pointed out that the shrinkage upon annealing to 1500 °C is difficult to quantify because its evolution is very poor. As a comparison, the SiBC30_10 samples display a single weight loss from 1120 °C to 1500 °C. A total weight loss as low as 0.55% has been measured at 1500 °C for the SiBC30_10 samples.

The differences in terms of high temperature thermal behavior between both samples are related to both the architecture and composition of the materials. First, it should be pointed out that the mSiBC30_10 samples contain a certain amount of oxygen (5.6 wt%) whereas the bulk analogs exhibit a very low oxygen content (0.3 wt%). The boron and carbon contents in the macroporous samples are 0.9 and 35.6 wt%, respectively. These values, especially for boron, are relatively close to those found in the bulk analog [33]. The higher carbon content in the porous samples most probably comes from the presence of carbon residue from decomposition of PMMA. The oxygen intake could be attributed to the combined effects of (a) the use of PMMA (its decomposition during the pyrolysis involving release of oxygen-based species that could react with the system which is not yet fully converted into a ceramic), (b) relatively high sensitivity of the preceramic polymer to moisture (during warm pressing, although performed in a glove-box, transfer into the furnace, handling for characterization) [23], and (c) the presence of an open porosity that makes the sample more sensitive to oxidation. In addition, we cannot exclude in both samples the presence of residual hydrogen that is removed above 1000 °C during the transformation of AHPCS830 into SiC [33].

Secondly, the mSiBC30_10 samples possess an open and accessible network structure (because of interconnected pores), as demonstrated by SEM images (Fig. 1), that clearly facilitates the release of gaseous species, especially compared to “closed” systems as represented by the bulk analog, i.e., SiBC30_10 samples. Thus, the release of residual hydrogen is most probably associated with the first weight loss identified between 1000 and 1300 °C for the mSiBC30_10 samples, and shifted to higher temperature, i.e., above 1120 °C, for SiBC30_10 samples. The weight loss observed in mSiBC30_10 samples when the temperature is raised beyond about 1300 °C could be attributed to the evolution of oxygen-containing gaseous species such as CO and SiO [44,45].

The latter is expected to readily react with free carbon to form stoichiometric SiC by carbothermal reaction at higher temperature, involving CO. To confirm the formation of SiC after annealing to 1500 °C, we observed the evolution of the porous structure of the samples by SEM/EDS after annealing to 1000 °C (See Fig. 1 as inset), 1300 °C and 1500 °C (Fig. 4 as inset). Whereas the SEM images of mSiBC30_15 samples, i.e., mSiBC30_10 samples annealed at 1300 °C (Fig. 4 as inset), do not show any difference in terms of pore structure and distribution compared to the mSiBC30_10 samples, we observed the formation of SiC (as confirmed by EDS) whiskers in the mSiBC30_15 samples, i.e., mSiBC30_10 samples annealed at 1500 °C (Fig. 4 as inset), highlighting the occurrence of vapor phase reactions at high temperature, most probably in the temperature range 1300–1500 °C. In such porous samples, the diffusion paths for CO (resulting from the reaction between silicon oxide gases and free carbon) are shorter and hence, formation of stoichiometric SiC is kinetically favored compared to bulk samples in which no SiC whiskers were identified after annealing to 1500 °C.

In order to understand the effect of the high temperature treatment of open-celled SiC foams on their porosity characteristics, the evolution of the pore architecture of the mSiBC30_10 samples between 1000 and 1500 °C was firstly assessed at the mesoscopic length scale by nitrogen gas adsorption-desorption measurements at 77K (Fig. 5). Thus, mSiBC30_X (X = 12 (1200 °C), 13 (1300 °C) and 15 (1500 °C)) samples were obtained by annealing treatment of the mSiBC30_10 samples at 1200, 1300 and 1500 °C (a dwelling time of 2 h was applied at the final temperature).

Based on IUPAC classification [3,46], the adsorption–desorption isotherms of the mSiBC30_10 samples has a type-II shape, characterized by a progressive increase of the adsorbed quantity of nitrogen with the P/P0 increase: such an isotherm is in general produced from non-porous or macroporous samples. As a confirmation of the macroporous nature of this sample, a specific BET surface area as low as 1.7 m²·g⁻¹ was measured, whereas a total pore volume (for pores smaller than 79.61 nm) of only 0.0017 cm³·g⁻¹ was determined from the amount of nitrogen adsorbed at P/P0 = 0.97. The horizontal isotherm characteristic of the mSiBC30_12 samples indicates a saturation of the adsorbent. Such isotherms are in general obtained with adsorbents containing micropores. This is confirmed with the relatively fast increase of the adsorbed volume at very low P/P0, indicating the development of microporosity in this sample. This can be related to the release of dihydrogen from the mSiBC30_10 samples from 1000 to 1200 °C, involving the first weight loss identified in the TG curve depicted in Fig. 4. As a consequence, an increase of the specific BET surface area to 6.7 m²·g⁻¹ was measured, whereas a total pore volume (for pores smaller than 79.6115 nm) of 0.0040 cm³·g⁻¹ was determined from the amount of nitrogen adsorbed at P/P0 = 0.97 in the mSiBC30_12 samples. The
specific BET surface area still increased in the mSiBC30_13 samples to
reach a value of 58.44 m²·g⁻¹, because of the contribution of micro-
pores (fast increase of the adsorbed volume at very low P/Po). To il-
lustrate this, a pore size of 2.93 nm was computed at the mesoscopic
length scale in the mSiBC30_13 samples, whereas it was 20 nm in the
mSiBC30_12 samples. The total pore volume (for pores smaller than
79.6115), determined from the amount of nitrogen adsorbed at P/
Po = 0.97, increased to 0.029 cm³·g⁻¹ confirming the formation of a
hierarchical porosity within the sample. The profile of the isotherms
of the mSiBC30_15 samples shows a H3-type hysteresis from P/Po = 0.4
to P/Po = 1, which is characteristic of a low degree of pore curvature
and non-rigidity of the aggregate structure. Although the BET surface
area decreased to 16.9 m²·g⁻¹, the total pore volume (for pores smaller
than 79.6115 nm), determined from the amount of nitrogen adsorbed at
P/Po = 0.97, still increased to 0.042 cm³·g⁻¹ most probably because of
the increased content of macropores to the detriment of mesopores. We
suggested that this was due to the coalescence of micropores and me-
sopores into macropores. Therefore, the evolution of the pore archi-
tecture of the mSiBC30_10 samples between 1000 and 1500 °C, at the
mesoscopic length scale, confirms the TG experiments concerning the
release of gases from the samples that help to develop the micro-
and mesoporosity in open-celled SiC foams at high temperature. Thus, open-
celled SiC foams with hierarchical porosity were produced at 1300 and
1500 °C. In contrast, the macroporous volume, measured by Hg por-
osimetry, did not change drastically upon annealing to 1500 °C; it varies
from 73.4 vol% (mSiBC30_10 samples) to 70.5 vol% (mSiBC30_15
samples). The mSiBC30_13 samples possess a porosity of 71.6 vol%. In
the same time, the skeletal density varies from 1.92 g·cm⁻³
(mSiBC30_10 samples) to 2.06 g·cm⁻³ (mSiBC30_13 samples) before
decreasing to 1.54 g·cm⁻³ (mSiBC30_15 samples). The mechanical
properties of the mSiBC30_10 samples remain stable even after at
1300 °C, i.e., mSiBC30_13 samples, which possess a compression
strength of 3.37 ± 1.86 MPa. It decreases to 2.26 ± 0.49 MPa for the
mSiBC30_15 samples that tend to confirm the degradation of open-
celled SiC foams upon annealing above 1300 °C.

Based on the results discussed above, it is reasonable to consider
that open-celled SiC foams are relatively stable even after annealing to
1300 °C. Interestingly, we can tune the porosity of the foams at
different length scales through the temperature of annealing: open-
celled SiC foams were produced at 1000 °C whereas foams with a
hierarchical porosity were produced at 1300 °C. In the latter, the
microstructural characterization carried out by XRD revealed the
presence of distinct XRD peaks related to the β-SiC phase at 2θ = 35.6° (111), 60.2° (220),
and 72.1° (311) (mSiBC30_13 samples; See Fig. 2SI in ESI), which were
poorly resolved in the XRD pattern of the mSiBC30_10 samples (See
Fig. 1SIa in ESI). The crystal growth also followed an increasing trend
with processing temperature, according to the calculation of the
average crystallite size by applying the Scherrer equation to the (111)
peak, which showed an increase of the grain size from 0.8 nm for the

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**Fig. 5.** N₂ adsorption-desorption isotherms of the mSiBC30_10 (a), mSiBC30_12 (b), mSiBC30_13 (c) and mSiBC30_15 (d) samples.
mSiBC30_10 samples to 2.3 nm for the mSiBC30_13 samples.

3.3. Ageing and microfiltration tests of open-celled SiC foams

Both mSiBC30_10 and mSiBC30_13 samples were used for investigating the ageing of open-celled SiC foams and their chemical stability in the working conditions of a primary coolant loop in a Pressurized Water Reactor (PWR). In parallel, the microfiltration capacity of foams toward colloids was investigated.

The primary loop of coolant, flowing through the reactor core at very high pressure to remain a liquid, prevents water from boiling within the reactor. It is comprised of a solution made of boric acid and lithium hydroxide to maintain the reactor coolant at an alkaline value. Because the primary loop is made of nickel and zirconium alloys as well as stainless steels, corrosion occurs. The corrosion products coming from the corrosion of the structure materials in contact with the primary coolant are in the form of soluble, colloidal or particulate compounds, which are transported by the primary coolant to the core where they become activated and redistributed out of the core surfaces. The deposits as radionuclides generate dose rate mainly at the bottom of the pressurized reaction vessel and on the surface of the steam generator tubes [47]. To limit the formation of radionuclides and reduce the dose rates, porous ceramics could be employed to separate such radionuclides from the solution. However, porous ceramics should be stable in the conditions imposed by the primary loop. In the present work, ageing tests were performed in a static autoclave under 140 bar at 330 °C in demineralized water containing 2.24 ppm (mg L⁻¹) of lithium and 590 ppm of boron (both measured by ICP), at a pH of 6.8, to reproduce the working conditions of a primary loop of coolant. Tests were carried out on a series of several pieces representative of the mSiBC30_10 samples (See Fig. 2). After the tests, samples labelled mSiBC30_10-at (at for ageing tests) were characterized by XRD, Hg porosimetry, SEM, EDX, and the solution after ageing was analyzed by ICP to measure the Si and B contents.

Samples were weighed before and after the tests according to this procedure: cleaning with demineralized water under ultra-sonication for 5 min, followed by drying in an oven at 100 °C for 24 h then in a desiccator for 12 h. The mSiBC30_10 samples undergo a weight loss of 6.8% after the ageing test, suggesting that the samples degrade under the imposed conditions. The solutions in which the samples were immersed possess a silicon (Si) content of 551 mg L⁻¹ (there was no Si in the solution before tests), whereas the boron (B) content is 728 mg L⁻¹ (590 mg L⁻¹ in the solution before tests) and the pH decreases from 6.8 to 4.3 after the ageing test. Therefore, we could infer that the mSiBC30_10 samples released silica, which is known to be an oxide acid, and boric acid. Although the mSiBC30_10 samples were X-ray amorphous (See Fig. 15la in ESI), it seems that a further “amorphization” process occurred after ageing tests in the mSiBC30_10-at samples, as shown by the broader (111) and (220) XRD peaks of the β-SiC phase (See ESI, Fig. 15la). SEM-EDS investigations showed the presence of spheres (1–2 microns in diameter) at the surface of the mSiBC30_10-at samples (Fig. 6b), which were not present in the mSiBC30_10 samples (Fig. 6a).

An O content of 57.8 at%, a Si content of 37.6 at% and a C content of 4.6 at% was measured by focusing in the area of spheres by EDX, whereas the part free of spheres (the bulk) had an O content of 12.9 at %, a Si content of 33.1 at% and a C content of 53.9 at%. Therefore, we suggested the formation of silica spheres at the surface of the samples that inherently lead to the “amorphization” of the mSiBC30_10-at samples as shown by XRD. It is important to mention that those spheres were not observed when using Li- and B-free demineralized water during the ageing tests. Therefore, this clearly confirms that silica spheres formed because of the presence of Li and B within the solution simulating the primary loop of coolant. As a result of the formation of silica spheres, Hg porosimetry analysis (See Fig. 2 and Fig. 35i in ESI) showed a distinct shift of the cell windows toward lower sizes in the range 1–6 μm. Interestingly, the degradation did not affect the microfiltration capacity of the foams, since the latter were able to trap magnetite particles present in demineralized water without pressure drop during tests for 220 h. It should be mentioned here that we did not use a solution made of Li and B in order to be able to distinguish magnetite particles (and iron oxide) from silica spheres deposited on the foams by SEM (See Fig. 2). As shown in Fig. 2, a crack appeared along the foams during the removal of the foams from the PTFE ring. However, we can consider that the strength of the foams is sufficiently high to withstand the conditions imposed by the application.

Because of their porosity characteristics and mechanical properties comparable to those of the mSiBC30_10 samples, the stability of the mSiBC30_13 samples was investigated in the same conditions. Interestingly, “amorphization” did not occur in the mSiBC30_13-at samples (See ESI, Fig. 45i), which could be related to the lower presence of silica sphere at the surface of the sample (See Fig. 6c (before ageing tests) and d (after ageing tests)). Furthermore, it appeared that the mSiBC30_13 samples release a significantly lower content of Si (255 mg L⁻¹), whereas the boron content released is similar (773 mg L⁻¹). This could be explained by the higher degree of crystallization of the mSiBC30_13 samples, that improved its chemical stability. Because of the lower release of Si as silica, the pH of the solution was measured to be 5.3. Although the pore junction size was also shifted to lower size from the mSiBC30_13 samples to the mSiBC30_13-at samples (See Fig. 35i in ESI), it had a smaller difference compared to the mSiBC30_10 and mSiBC30_10-at samples. Therefore, we may infer that open-celled SiC foams prepared at 1300 °C display a higher stability in harsh environment compared to those prepared at 1000 °C, and could withstand the harsh conditions imposed by the primary loop of coolant in PWR.

4. Conclusion

Open-Celled silicon carbide (SiC) foams with an interconnected porosity of 73.4 ± 0.5 % were prepared from a mixture of boron-modified polycarbosilane and poly(methymetacrylate) (PMMA) microbeads (25 μm) sacrificial fillers in a 20:80 ratio. The boron-modified polycarbosilane was synthesized through the reaction between allylhydridopolycarbosilane (AHPCS) and borane dimethylsulfide (BODMS) according to a controlled Si:B ratio of 30. It allowed to significantly reduce its processing temperature compared to thermally crosslinked AHPCS. Thus, warm-pressing of the powder mixture was performed at a temperature as low as 120 °C and the green body was then pyrolyzed at 1000 °C under argon. The resulting highly porous components were analyzed in detail at the mesoscopic and macroscopic length scales, and their structural stability as well as compressive strength was investigated. Open-celled SiC foams combined a low density with a compressive strength of 3.49 ± 0.56 MPa and a thermostructural and mechanical stability that can be extended up to 1300 °C, temperature at which a single-step weight loss continuously occurred up to 1500 °C without losing the monolithic structure. It was attributed to the evolution of oxygen-containing gaseous species such as CO and SiO. As a consequence of the release of gaseous species, the BET SSA of foams increased in the temperature range 1000–1500 °C to form open-celled SiC foams with a hierarchical porosity at 1300 and 1500 °C. Ageing and microfiltration tests in the condition of a primary loop of coolant in a Pressurized Water Reactor (PWR) showed that open-celled SiC foams were able to retain particles with diameter around 5 μm without degradation of the performance of filtration, but those prepared at 1300 °C demonstrated a higher chemical stability than open-celled SiC foams prepared at 1000 °C in the tested conditions.

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