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Polymer-Derived Ultra-High Temperature Ceramics (UHTCs) and Related Materials

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Ultra-high temperature ceramics (UHTCs) represent an emerging class of materials capable of providing mechanical stability and heat dissipation upon operation in extreme environments, e.g., extreme heat fluxes, chemically reactive plasma conditions. In the last few decades, remarkable research efforts and progress were done concerning the physical properties of UHTCs as well as their processing. Moreover, there are vivid research activities related to developing synthetic access pathways to UHTCs and related materials with high purity, tunable composition, nano-scaled morphology, or improved sinterability. Among them, synthesis methods considering preceramic polymers as suitable precursors to UHTCs have received increased attention in the last few years. As these synthesis techniques allow the processing of UHTCs from the liquid phase, they are highly interesting, e.g., for the fabrication of ultra-high temperature ceramic composites (UHT CMCs), additive manufacturing of UHTCs, etc. In the present review, UHTCs are in particular discussed within the context of their physical properties as well as energetics. Moreover, various synthesis methods using preceramic polymers to access UHTCs and related materials (i.e., (nano)composites thereof with silica former phases) are summarized and critically evaluated.

1. Introduction

1.1. Definitions, Terms, and Limitations

There are two ways to define ultra-high temperature ceramics (UHTCs). Firstly, a strict definition relates to their melting

points and requires that UHTCs melt at temperatures beyond 3000 °C (sometimes 3000 K).[1,2] These materials are relevant mainly for applications which operate at extremely high temperatures (e.g., $T \gg$ 2000 °C) and for short times (typically in the order of minutes, sometimes few hours). Within this context (and due to reasons discussed in some detail below), borides, carbides and nitrides of group 4 transition metals are considered to be typical UHTCs, as they offer the fundamental advantage not to melt at these tremendously high temperatures. However, they suffer from poor environmental compatibility (e.g., poor oxidation resistance), so they usually resist UHTs only for a very short exposure time.[3] Secondly, an alternative definition for UHTCs relates to materials which can be used at temperatures higher than those addressed by typical high-temperature ceramics (such as silicon nitride or silicon carbide).[1,2] Thus, materials capable of long-term operation at temperatures beyond 1600 $^{\circ}$ -

C (this temperature is generally accepted as the long-term operation limit for Si_3N_4 or SiC) are considered. Their melting points do not necessarily have to exceed 3000 °C (however, the higher the temperature of melting ($T_{\rm m}$), the better); whereas their operation time should be significantly longer than those of the materials mentioned within the context of the first

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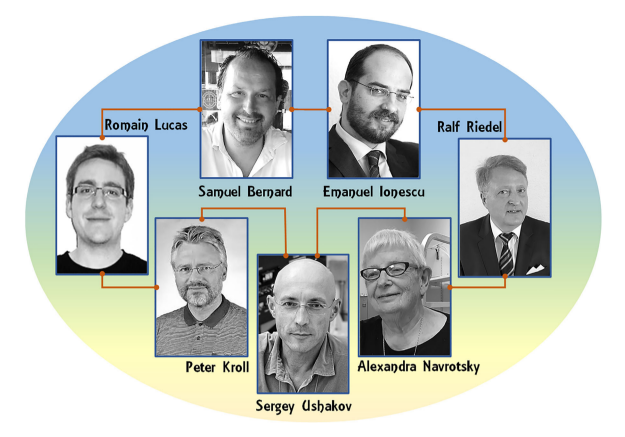
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DOI: 10.1002/adem.201900269



definition. Thus, UHTCs are either 1) materials with $T_{\rm m}$ higher than 3000 °C and which can resist T \gg 2000 °C and short exposure times to hostile environments or 2) materials which can withstand T > 1600 °C and long-term exposure to environmental loading.

When operated at high temperatures (HT) or ultra-high temperatures (UHT) and in extreme environments, structural parts and components are exposed to additional and more severe design constraints as compared to those used in service at or near ambient temperature. [1,4,5] A fundamental property which is required in order to be able to operate materials at (ultra)high temperatures relates to their melting point. Naturally, (ultra)high melting points of the chosen materials are necessary, as usually the maximum operation temperature of a material is round half of its melting point with some exceptions, such as Ni-base superalloys, which can be operated at temperatures up to 90% of their melting points. [6-9] Moreover, the environmental degradation of the materials becomes critical at high and ultra-high temperatures and consequently materials capable to withstand thermal decomposition as well as oxidation and corrosion should be taken into consideration. Hence, a basic requirement to be fulfilled relates to thermal stability and environmental (chemical) inertness of the materials of choice. Operation of materials at high and ultra-high temperatures is also related to other factors such as thermally induced stresses and time dependent inelastic strain (creep), which may significantly contribute to catastrophic failure of the components during operation. Depending on the failure consequences (i.e., economic vs. safety-related aspects),

those requirements are very strict and materials/components design might suffer here from an extremely high level of conservatism.

Generally, high tolerance to thermal and mechanical stress is desirable in order to avoid catastrophic failure of the materials during operation. Thus, high compliance to thermal stress can be provided by using highly thermal shock resistant materials. Additionally, suitably optimized ductility of the materials or, in the case of brittle (e.g., ceramic based) components, improved fracture toughness is desirable in order to provide high tolerance to mechanical stresses. Finally, no phase transformation of the materials in the range from ambient to operation temperature should occur, as such processes usually are accompanied by significant volume changes which consequently are likely to generate tremendous stresses and induce catastrophic failure. In the case of amorphous materials, increased crystallization resistance may be beneficial for avoiding the generation of mechanical stress upon operation at (ultra)high temperatures.

Intense efforts have been made in the last decades to accommodate those requirements, which are often strongly conflicting. Thus, several types and classes of (ultra)high-temperature materials have been proposed and developed over the years, such as highly refractory metals, intermetallics/alloys, metal matrix composites (MMCs) as well as UHTCs or ceramic matrix composites (CMCs).

When trying to identify suitable materials for operation at UHTs and in extreme environments, one major criterion to be taken into account is their melting point. There are more than 300 materials which exhibit a melting temperature above

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2000 °C, including highly refractory metals (such as Hf, Nb, Ta, W, Re, Ir) and numerous ceramics (oxides, carbides, nitrides, borides). However, only few materials are known to exhibit melting points beyond 3000 °C, among them elements (W, Re, C), as well as binaries with high covalent bonding character (e.g., BN, ThO₂, TiC, ZrC, ZrB₂, HfC, HfN, HfB₂, NbC, TaC, TaN, TaB₂). [1,10] They are denoted as ultra-high temperature ceramics (UHTCs). Beside their ultra-high melting points, UHTC materials should withstand non-steady-state interactions with combustion environments and friction with the atmosphere (as for re-entry vehicles). These extreme conditions strongly limit the number of suitable materials among those presented above.

Thus, materials which are left after considering the mentioned criteria are some group 4 carbides, nitrides and diborides (especially that of zirconium and hafnium).[11] This class of materials was first investigated between the 1950s and 1970s, mainly in former USSR and USA and has received increased attention within the last 10–15 years, due to the efforts in developing hypersonic flight vehicles. [4,12] It has been shown. however, that bulk single-phase UHTCs exhibit rather poor oxidation resistance and damage tolerance under extreme environments. Within this context, the development of UHTCs moved toward composite formulations with improved performance in harsh oxidative environments. Thus, oxidation resistant (mostly particulate) phases have been added to UHTCs, such as SiC or silicides (TaSi₂, MoSi₂, etc.).^[1,2] Especially ZrB₂/ SiC and HfB₂/SiC (with ca. 20 vol% SiC) composites have been reported to exhibit promising environmental behavior at UHTs, as they form a protective borosilicate glass coating.[13,14] Moreover, intense efforts were made to further improve the environmental stability of SiC-modified diboride-based UHTCs by increasing the viscosity of the liquid silica-based layer, improving the immiscibility of multi-phasic liquid layers formed during oxidation process.[15]

1.2. Thermodynamic Considerations and Solid-State Synthetic Approaches for UHTCs

1.2.1. Borides

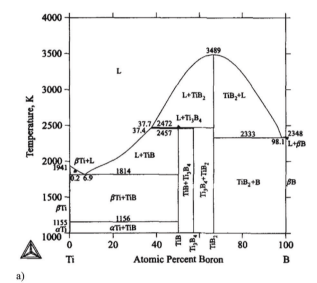
The excellent thermal stability as well as high hardness of group 4 transition metal diborides rely on their crystalline structure. They exist in a primitive hexagonal crystal structure (AlB₂-type, space group P6/mmm) with two-dimensional graphene-like boron layers alternating with hexagonally close-packed metal layers. Thus, the extremely robust B–B and M–B bonds in MB₂ (M = Zr, Hf) are naturally responsible for their ultra-high refractoriness and hardness (Table 1). Accordingly, ZrB₂ and HfB₂ exhibit extremely high melting temperatures (3245 and 3380 °C, respectively) and the highest hardness values among their periods (23 and 28 GPa, respectively). [13,16,17]

In contrast to rocksalt nitrides and carbides, diborides have little variation in stoichiomentry and are essentially line compounds (see **Figure 1**a as for the Ti–B system^[18]). Diborides of Ti, Zr, and Hf are the highest melting borides; however, in the Ta–B system (Figure 1b),^[19,20] the monoboride TaB has a marginally higher melting temperature than that of the

Table 1. Some physical and structural properties of ZrB₂ and HfB₂ (adapted from ref. [13]).

	Density [g cm ⁻³]	Melting point [°C]	CTE [ppm K ⁻¹]	Thermal conductivity [W/(m K)]	Hardness [GPa]	Young's modulus [GPa]
ZrB_2	6.2	3245	5.9	60	23	489
HfB_2	11.2	3380	6.3	104	28	480

corresponding diboride TaB₂ (3363 \pm 15 K for TaB vs. 3310 \pm 20 for TaB₂). TaB is not found in NaCl-type structure, but possesses an orthorombic CrB-type (*Cmcm*) structure. ^[21] The monoboride of Zr was reported ^[22] in NaCl structure, but it is only stable in a narrow temperature range below 1500 K.



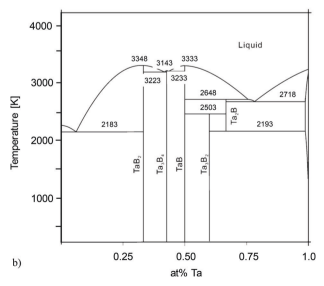


Figure 1. Phase diagrams of a) Ti-B (http://www.calphad.com/titanium-boron.html) and b) Ta-B systems. Reproduced with permission.^[20] Copyright, Elsevier.

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Over the last decade, multiple groups studied the thermodynamic properties of ZrB_2 and HfB_2 . Lawson et al. analyzed in detail the bonding situation in ZrB2 and HfB2[23] and concluded that covalent bonding provides cohesion between boron atoms, while metallic bonding persists within the metal layer. A substantial charge transfer from the metal layer to the boron layer yields to strong ionic bonding between layers and is responsible for the substantial refractory character. Zhang et al. recently reported the remarkable structural stability of ZrB₂.^[25] Even when subjected to extreme pressures, the hexagonal structure remains the only competitive modification of the diboride. Further Zr-B compounds have been investigated as well. Without doubt, ZrB in the rocksalt structure is mechanically stable. [26] However, computed lattice parameters do not agree with experimental data within the usual tolerances of firstprinciples calculations. Further mechanically and dynamically stable structures are Zr₃B₂, Zr₃B₄, Zr₂B₃, ZrB₂, and ZrB₁₂. [27] Only ZrB₂ appears to be thermochemically stable, and results for ZrB₁₂ are still undecided. [28] However, a mononitride ZrB appears as the least competitive phase, and the rocksalt type of ZrB is less favorable than an orthorhombic modification. This, furthermore, suggests that the observed rocksalt-type may either be defect-rich or even stabilized by carbon. Noteworthy in this context is the work of Lee et al., who identified insertion of boron into octahedral sites of the hcp Zr structure as energetically most favorable.[29]

The thermochemistry of Hf–B compounds is very similar to those in the Zr–B system. HfB_2 dominates the phase diagram, with orthorhombic HfB and HfB_{12} competitive but not favored by enthalpy. $^{[28,30]}$

The Ta–B system shows more diversity than borides of Zr and Hf (see Figure 1b). With exception of the well-known AlB_2 -type of TaB₂, all appearing structures are quite complex thanks to prevalent Ta–Ta bonding. Boron-rich phases exhibit fragments of the TaB₂ structure, e.g., a one-dimensional strip of condensed B₆-rings, interrupted by additional Ta atom. Tantalum-rich structures display motifs of dense packings with B atoms filling interstitial sites, however, with significant distortions. Calculations indicate that Ta₃B₂, TaB, Ta₅B₆, Ta₃B₄, Ta₂B₃, and TaB₂ are thermochemically stable. [²⁸]

Diboride-based UHTCs can be synthesized by different approaches. Most of them rely on reduction processes of the oxides (zirconia, hafnia, boria). Carbon and boron are the most used reducing agents. The carbothermal process has been the mostly used technique to produce diborides and involves the reaction of a mixture consisting of metal oxide (i.e., $\rm ZrO_2$ or $\rm HfO_2$) and $\rm B_2O_3$ with carbon. As these processes are strongly endothermic, temperatures of synthesis typically around 2000 °C are needed. Usually, boria is added in excess in order to promote the formation of the diborides and to inhibit carbide formation. Common impurities in carbothermally synthesized diboride powders are carbon, boria, and metal carbides. $^{[11,13]}$

The reaction of the group 4 transition metal oxides with boron (borothermic reaction) as well as aluminothermic processes (MO₂ and boria reacting with Al) are also strongly endothermic and are performed at temperatures exceeding $1400-1500\,^{\circ}\text{C}.^{[11]}$ The reduction of the metal oxides with boron carbide (B₄C) was shown to become thermodynamically

favorable at temperatures as low as $1200\,^{\circ}\text{C}$. An additional advantage of the reduction process with B₄C relies on the reduced amounts of carbon and oxide impurities in the synthesized diboride powders. [31,32]

Group 4 transition metal diborides (as well as other transition metal borides) can also be synthesized from elements. Depending on the purity of the starting materials, diborides of high purity are accessible upon this process. However, due to the poor oxidation resistance of the group 4 transition metals, these processes have to be performed in inert atmosphere in order to avoid the formation of the corresponding metal oxides. As the reactions of Zr and Hf with B are highly exothermic, [111] self-propagating high-temperature synthesis (SHS) techniques have been developed to prepare diboride powders from elements. [33,34]

Table 2 summarizes best available values for melting temperatures, standard entropies, and enthalpies of formation at 298 K for selected borides, carbides, and nitrides of Ti, Zr, Hf, and Ta. The data for corresponding oxides are included for comparison. The data referenced to NIST-JANAF and Glushko's thermochemical tables are largely based on the evaluation of measurements performed before 1970. Formation enthalpies for TiC, TaB₂, and TiB₂ are given after more recent calorimetry measurements performed in Kleppa's group. [35,36] The uncertainties are given when provided in the original reference.

Synthetic approaches using reactive processes have also been applied to the preparation of diboride-silicon carbide composite materials. Thus, hot-pressing of a mixture of metallic zirconium, carbon, and silicon tetraboride, SiB₄, led to monolithic ZrB₂/SiC composites. A small amount of ZrC impurity has been also detected. $^{[50]}$

Reactive hot-pressing of a powder mixture containing zirconium, boron carbide B_4C , and silicon was also used to prepare ZrB_2/SiC monoliths with high densities at a temperature as low as $1900\,^{\circ}C.^{[51]}$

1.2.2. Carbides and Nitrides

Generally, transition metal carbides, nitrides, and carbonitrides represent a unique class of materials with outstanding structural

Table 3. Selected physical and structural properties at room temperature of stoichiometric group 4 transition metal carbides and nitrides (i.e., composition near to 50 at% metal).

	ZrC	HfC	ZrN	HfN
Density [g cm ⁻³]	6.5	12.3	7.3	13.8
Melting point [°C]	3420	3930	3000	3330
CTE [ppm K ⁻¹]	7.5	6.1	7.8	8.5
Th. conductivity [W/(m K)]	24.6	25.1	11	11
Hardness [GPa]	25	20	15	18
Young's modulus [GPa]	350	420	460	380
Electrical resistivity [$\mu\Omega$ cm]	75	67	24	27
Curie temperature, T_c [K]	_	_	10.47	6.92

Data taken from ref. [52].

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properties, such as very high melting points, exceptional high hardness, or high-temperature ablation resistance. [52] Furthermore, they exhibit metallic character with respect to electrical and thermal conductivities, fascinating surface properties, and interesting catalytic behavior. In particular, group 4 transition metal carbides and nitrides have been shown to exist over broad ranges of composition with an upper limit to C:M or N:M ratio close to unity. They exhibit tremendous thermal stability (e.g., group 4 transition metal nitrides can melt without suffering decomposition, unlike groups 5 or 6 transition metal nitrides), which has been shown to rely on their free energy of formation. [52]

Group 4 carbides and nitrides belong to the class of interstitial compounds, having as characteristics simple metallic lattices (face-centered cubic, fcc) and carbon or nitrogen atoms located in their octahedral interstitial sites. Their bonding is characterized by a mixture of metallic, ionic, and covalent components. With respect to thermal or electronic transport, carbides and nitrides of group 4 transition metals exhibit metallic behavior. As the carbides show lower density of state (DOS) at the Fermi level than the corresponding nitrides, the electrical and thermal conductivity as well as the $T_{\rm c}$ values of the group 4 carbides are lower than those of the analogous nitrides. [52]

Transition metal carbides are known for their extremely high melting points (Table 3). For instance, ZrC shows a congruent melting point of $3420\,^{\circ}$ C, whereas HfC melts congruently at $3930\,^{\circ}$ C. The material predicted to have the highest known melting point ($4215\,^{\circ}$ C) is a solid solution of HfC and TaC in a molar ratio 4:1, with a composition of Ta_4 HfC₅ (compare to the melting point of TaC, $3985\,^{\circ}$ C). Also, transition metal nitrides have extremely high melting points, often above $3000\,^{\circ}$ C (TiN: $3030\,^{\circ}$ C; ZrN $3000\,^{\circ}$ C; HfN $3330\,^{\circ}$ C).

Table 2. Standard entropies, enthalpies of formation, and melting temperatures for selected borides, carbides; and nitrides of Ti, Zr, Hf, and Ta.

<u> </u>	V110C tl 1=J2	60 u 1-1 v-1	T 10.61
Compound	$\Delta H^{\circ} f_{298.15} [kJ mol^{-1}]$	$S^{\circ}_{298.15} [J \text{ mol}^{-1} \text{ K}^{-1}]$	T _m [°C]
TiC	$-186\pm18^{[35]}$	$24.7 \pm 0.2^{[37]}$	$3067 \pm 25^{[37]}$
TiB ₂	$-328\pm10^{[36]}$	$28.9 \pm 0.4^{[37]}$	$3225 \pm 25^{[38,39]}$
TiN	$-338 \pm 4^{[40]}$	$30.3 \pm 0.2^{\rm [40]}$	$2945 \pm 30^{[41]}$
TiO ₂	$-944 \pm 1^{[37]}$	$50.3 \pm 0.2^{[37]}$	$1870\pm15^{[37]}$
ZrC	$-207 \pm 3^{[37]}$	$33.3 \pm 0.1^{[37]}$	$3572 \pm 30^{[42]}$
ZrB_2	$-323 \pm 7^{[40]}$	$35.9 \pm 0.1^{[40]}$	3244 ^[43]
ZrN	$-342^{[44]}$	$38.9 \pm 0.2^{[37]}$	$2955 \pm 30^{[41]}$
ZrO_2	$-1101\pm1^{[37]}$	$50.4 \pm 0.4^{[37]}$	2710 ^[45]
HfC	$-208 \pm 8^{[37]}$	$40.1 \pm 0.2^{[37]}$	$3982 \pm 30^{[42]}$
HfB ₂	$-329 \pm 9^{[46]}$	42.6 ^[37]	$3378 \pm 20^{[22]}$
HfN	$-374 \pm 2^{[37]}$	$45\pm1^{[37]}$	$3330 \pm 50^{[41]}$
HfO ₂	$-1118\pm 2^{[37]}$	$59.3 \pm 0.6^{[37]}$	2800 ^[45]
TaC	$-142 \pm 3^{[37]}$	$42.3 \pm 0.2^{[37]}$	$4003 \pm 30^{[42]}$
TaB ₂	$-160 \pm 6^{[47]}$	$47 \pm \mathbf{4^{[37]}}$	$3037 \pm 20^{[48]}$
TaN	$-252 \pm 2^{[49]}$	$41.8 \pm 1.2^{[49]}$	$\dagger 2930 \pm 30^{[41]}$
TaO _{2.5}	$-1024\pm1^{[37]}$	$71 \pm 1^{[37]}$	$1872 \pm 10^{[37]}$

Group 4 transition metal carbides can be prepared from elements as well. As the reactions are strongly exothermic, SHS techniques can be used to synthesize carbide powders. Transition metal nitrides can be produced similarly. [53] Carbides can also be synthesized by reduction of oxides with carbon at high temperatures. Thus, zirconia or hafnia powders can be reacted with carbon at temperatures as low as 1600 °C to prepare fine and ultrafine carbide powders. [54–58] Due to their high sinter activity (and to some extent to small oxygen impurities), they can be densified at relatively low temperatures. The carbothermal process was combined with the SPS technique to prepare

The corresponding bulk nitrides can be prepared from group 4 transition metals or metal hydrides via heat treatment in nitrogen or ammonia atmosphere. Carbo-nitridation of metal oxides at high temperatures also furnishes metal nitrides (which contain oxygen and carbon impurities).^[52]

monolithic zirconium and hafnium carbides.^[59]

Except for TaN, all compounds listed in Table 2 are reported to melt congruently. TaN decomposes to Ta_2N which has a slightly higher melting temperature (3232 K). It is not uncommon to find discrepancies of a hundred degrees or more in melting temperatures between different reports, especially for borides and nitrides. Melting temperatures of nitrides are reported after measurements by Ettmayer et al., [41] which are the average from experiments performed at 1–80 atm nitrogen pressure.

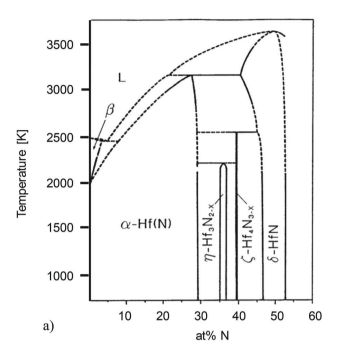
Ti, Zr, Hf, and Ta carbides and nitrides are found in NaCl (rocksalt) structure, in whichs the metal forms a cubic close-packed lattice and C or N fill some or all octahedral sites comprising from 35 to 50 at%. These compounds possess high electrical conductivities. They are sometimes referred to as δ -phase^[60] or interstitial phases^[61] referring to intercalation of *fcc* metal lattice with C or N.

Complete miscibility can be found among Ti, Zr, Hf, Ta carbides and nitrides at high tempeatures (ca. 2000 °C). The stability of these solid solutions and exact temperatures and shapes of miscibility gaps depend on vacancies concentrations. [62] From the analysis of phase diagrams, the congurently melting compositions for rock salt carbides and nitrides deviate from ideal stoichiometry with 44–48 at% of nonmetal.

Yu et al. investigated several structures in the Ta–C system and showed the elasticity of TaC_{1-x} decreases with loss of carbon in the structure, which is related to a decrease in covalent bonding between Ta and C paralleled by an increase in metallic bonding. ^[63] This is also the underlying motif of the perfectly ordered ζ -like structure of Ta_3N_2 with $[\text{M}_6-\text{C}_4/\text{V}_2-\text{M}_6-\text{C}_4/\text{V}_2]==M_3\text{C}_2$.

An interesting topic are mixtures of metal carbides forming UHTCs. Not only is the enthalpy of mixing relevant to stability of phases, but also a potential mixing entropy among the metal cations. Vorotilo et al. investigated solid-solutions of TaC and ZrC in the rocksalt structure with full occupation of cation and anion positions. [64] Computed lattice parameters and bulk moduli of solid solutions follow a rule of mixture and agree with experimental data. The enthalpy of mixing ΔH of intermediate compositions $Ta_x Zr_{1-x}C$ is negative for all x with a maximum ΔH = -0.030 eV atom⁻¹ at x = 0.625. Experimental work carried out in parallel did not indicate decomposition of solid solution, suggesting complete solubility without miscibility gap in the $Ta_x Zr_{1-x}C$ phase diagram.

The Hf–N system (see phase diagram in **Figure 2**a^[65]) displays several compounds favored by enthalpy. Two recent comprehensive studies present structures and thermochemistry in this system using DFT calculations. ^[66,67] Besides HfN, structures of compositions Hf₄N₃, Hf₃N₂, Hf₂N, Hf₃N, Hf₆N, and, probably,



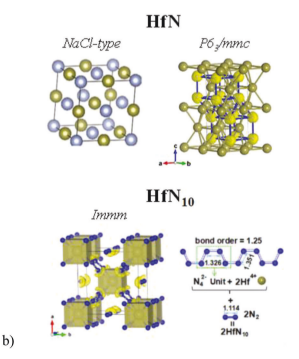


Figure 2. a) Phase diagram of the Hf–N system (Reproduced with permission. [65] Copyright, Springer); b) crystalline structures of HfN (NaCl-type and $P6_3/mmc$) and of HfN $_{10}$ (Immm). For Immm-HfN $_{10}$, infinite polymeric nitrogen chains and $N\equiv N$ bondings are shown (Reproduced with permission. [67] Copyright, American Physical Society).

Hf₈N appear thermochemically stable towards the metal-rich side of the phase diagram. All structures can be described as being related to densest packings of metal atoms with N filling octahedral interstitials. Surprisingly, a new polymorph of HfN with hexagonal structure ($P6_3/mmc$; Figure 2b) was discovered to be the most favorable modification of HfN. [67] Careful work indicated that the rocksalt type of HfN (Figure 2b), previously assumed to be dominant, only becomes thermodynamically stable above 670 °C. Metal cations in hexagonal P63/mmc-HfN fall onto the sites of the La structure type, a simple intergrowth of hcp and ccp with repetition every four layers, also termed double hexagonal closest packing (*dhcp*). Anions fill all octahedral sites, giving the hexagonal HfN structure the flavor of NiAs-type and rocksalt type joined together. Toward the nitrogen-rich side of the phase diagram structures with composition Hf₁₁N₁₂, Hf₇N₈ and Hf₄N₅ appear, all described as cation-deficient rock-salt structures. Besides the high-pressure phase Hf₃N₄ with Th₃P₄-type structure, ^[60] the most nitrogen-rich compound of Hf thermochemically stable at ambient pressure is Hf₃N₄, for which a new monoclinic structure has been proposed. [67] At pressures beyond 23 GPa, an Immm-HfN₁₀ phase was predicted via DFT calculations and exhibits Hf atoms eightfold coordinated by nitrogen and consists of infinite polymeric nitrogen chains (N-N bond has a bond order of 1.25) and encapsulated N≡N molecular units (Figure 2b).

Calculations of the Ta-N system do not yield as many thermochemically stable structures as the Hf-N system. Ta₃N₅, which can be synthesized, appears together with Ta2N3 as nitrogen rich phases in the phase diagram at low temperatures. It is well known, however, that upon annealing to 1000 °C, Ta₃N₅ decomposes through a series of phases of intermediate composition into TaN. The phase diagram (**Figure 3a** $^{[68,69]}$) is then dominated by ϵ -TaN $^{[70-72]}$ with a hexagonal structure related to the intermetallic CoSb (Figure 3b). Ta atoms form dense packed layers which are tightly stacked similar to a hexagonal closest packing. With the shorter stacking distance, Ta-Ta bonds persist between next-nearest layers. Anions are then located within the plane of every other Ta layer and do not occupy interstitial sites (Figure 3b). In comparison to rocksalttype TaN, ε-TaN has a major energy advantage, but lacks a similar defect chemistry. The next stable phase then is Ta₂N, which structure is best described as a hexagonal closest packing of Ta with N filling octahedral sites in every other layer (Figure 3b). Although the metal arrangements in both ϵ -TaN and Ta2N are described in similar terms, intermediate structures with anion deficiency do not appear with competitive enthalpy of formation.[66]

It is worthy to mention that first-principles calculations of thermochemistry for nitride UHTCs face a tough challenge. Reaction enthalpies are computed at 0 Kelvin. Entropy differences between reactants and products are often neglected, and enthalpy differences are assumed to remain constant over the range of temperatures. Even if those differences are taken into consideration — and there are some fine works that go the extra mile^[67] — such an approach is impossible for nitrogen. N is a gaseous molecule at ambient conditions, quite unlike elemental B and C, which are solid-state compounds. Being a gas, both enthalpy and entropy of nitrogen change significantly larger with increasing temperature than those of the solids, and the

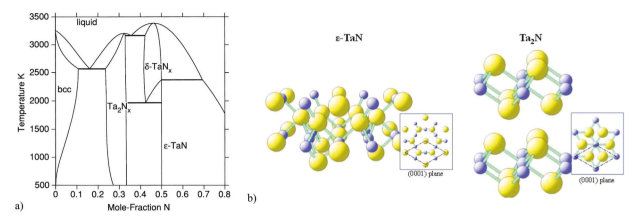


Figure 3. a) Phase diagram of the Ta–N system^[68,69] (Reproduced with permission.^[69] Copyright, Springer); b) crystalline structures of ε-TaN (*P6/mmm*) and Ta₂N ($P\bar{3}ml$): N and Ta: blue and yellow spheres, respectively; dashed lines in the insets indicate the cell unit (Reproduced with permission.^[72] Copyright, American Physical Society).

computational arsenal to assess heat capacity and vibrational entropy loses its strength. Consequently, one cannot hope for fortuitous error cancellation in reaction enthalpies between products and reactants. This sets N apart from B and C, and makes incorporation of the chemical potential of nitrogen at elevated temperatures indispensable for predicting nitride phases.^[73–75]

Standard enthalpies of formation are the least negative for carbides and most negative for oxides. This does not reflect the bond strength in compounds, but is mainly related to the difference in the reference states for nonmetal atoms, which are N_2 , O_2 , and graphite. Formation enthalpies show that carbides, nitrides, and borides are expected to be unstable to oxidation. For Zr, Hf, and Ta, melting temperatures decrease in the order of carbides – borides – nitrides – oxides, spanning a range of 1–2000 degrees. Titanium diboride is an exception from this trend, with its melting temperature (3498 K) exceeding that of TiC (3340 K).

No experimental measurements of fusion enthalpies for Ti, Zr, Hf, and Ta carbides, borides and nitrides have been reported. First principles computations are increasingly used to evaluate thermodynamic properties of these compounds: thermal expansion and heat capacities were computed for ZrC and ZrN up to 3000–K^[76] and the effect of carbon vacancies on miscibility gap in TiC–ZrC carbides was analyzed, ^[62] and melting temperatures were studied in the Hf–C–N system. ^[77]

Since carbides, borides, and nitrides of transition metals are good electrical conductors, most of the measurements of their melting temperatures were performed by direct bulk heating of the samples with electric current using an experimental arrangement known as Pirany furnace. In this method, first used by Pirany and Alterthum in 1923,^[78] the melting occurs in the bottom of the blind hole drilled in the test specimen. The hole also serves as black body cavity to aim pyrometer for the temperature measurements, eliminating the uncertainty from sample emissivity. This arrangement was used by Rudy et al.^[79] on carbides and borides and by Ettmayer^[41] to study nitrides. For metals which can be drawn as a wire, the Joule heating evolved

into exploding wire technique, which now also yields enthalpy as a function of temperature in the solid and liquid state. [80]

For almost 40 years no new studies on melting temperatures of carbides, borides, and nitrides by Joule heating method have been performed; however, the renewed interest in refractory carbides for UHTCs resulted in a new surge of experimental efforts. Pulsed laser melting technique, originally developed for refractory oxides, was recently applied to study the melting of Ta, Hf, and Zr carbides. [42,81] In this method, melting is induced by millisecond laser pulses, the surface temperature is recorded with a fast spectropyrometer and liquidus and solidus temperatures are derived from thermal arrests on cooling. Pulsed laser measurements of HfC melting point performed by Barraza et al. [81] in Germany (3959 \pm 84 °C) and by Sheindlin et al. [42] in Russia (3982 \pm 30 °C) are in reasonable agreement with each other and with earlier Pirany furnace measurements by Rudy et al. [79] in California (3928 \pm 40 °C). However, there is a large discrepancy for reported TaC melting point from pulsed laser measurements: $(3768 \pm 8 \,^{\circ}\text{C})^{[81]}$ versus $(4003 \pm 30 \,^{\circ}\text{C})^{[42]}$ The higher value reported by Sheindlin et al.[42] (and included in Table 2) is also consistent with earlier results from Pirany furnace (3983 \pm 15 °C). [79] Recently, a pulsed Joule heating technique was also used on 250 µm thick carbide samples, [82,83] to analyze melting temperature, heat capacity and fusion enthalpy for Zr carbide and nitride. [84,85]

The prize for highest melting solid was given to $Ta_{0.8}Hf_{0.2}C$ in 1930 by Agte and Alterthum^[86] (\approx 60 °C higher than TaC), in 1967 Rudy et al.^[79] gave it to TaC (3983 °C). Both teams were using Pirany furnace. Barraza et al.^[81] gave it to HfC from laser pulse melting, and recent work using electric pulse method returns it back to $Ta_{0.8}Hf_{0.2}C$ (4027 °C).^[83] However, computational work predict that experimentalists are looking in the wrong system and the highest melting solid is a Hf–C–N alloy^[77] (Figure 4).

There is much to be done in refining experimental measurements and synthesis, and characterization techniques and, for practical applications, oxidation behavior and properties of oxycarbides are more critical than absolute values of melting temperatures for pure compounds.

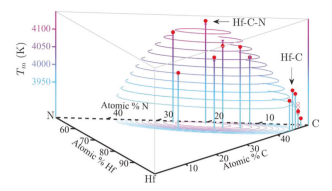


Figure 4. Melting temperatures of materials compositions in the Hf–C–N and Hf–C (red filled circles) as well as Hf–Ta–C (empty circles) systems (Reproduced with permission.^[77] Copyright, American Physical Society).

2. Ceramic Synthesis from Preceramic Polymers (the PDC Route) – A Brief Introduction

Single-source preceramic precursors provide a facile and inexpensive access to various ceramic systems with tailored phase compositions, microstructures and property profiles. Among them, polymeric single-source precursors are of special relevance, as they can be used for the preparation of ceramic bulk powders, dense monoliths, macro, meso and micro porous bodies, thin films and coatings, miniaturized (microstructured) components, fibers, etc. Furthermore, polymeric single-source precursors allow preparation of unique ceramic compositions and microstructures, which are not accessible via other preparative techniques. The molecular structure and chemistry of the single-source precursors greatly influence the composition, microstructure, and phase distribution of the final ceramic product, providing the advantage to control and design ceramic materials with various macroscopic and physical properties.^[87] In this context, a crucial role is played by the synthesis of suitable precursors with tailored composition and molecular architecture.[87-89]

Polymer-based single-source precursors for the preparation of ceramic nanocomposites were taken into account in the 1980s, [90,91] as researchers started to look for appropriate synthesis techniques for nanoscale non-oxidic ceramics. However, already in the late 1950s and 1960s studies related to the conversion of molecular precursors into non-oxidic ceramics were reported^[92–94]; furthermore, in the 1970s, polymeric precursors such as polycarbosilanes (PCS) or polysilazanes (PSZ) were used to prepare small-diameter SiC and SiC/Si₃N₄ ceramic fibers. [95,96] These preparative efforts culminated in the commercialization of the Yajima production process of continuous SiC ceramic fibers. [96,97] These SiC fibers were shown to exhibit excellent high temperature stability and promising mechanical properties. However, there are some limitations concerning their use at high temperatures, as they possess limited oxidation resistance and poor mechanical properties above 1300 °C. [98,99] In this context, numerous studies have been performed with respect to the modification of the SiCbased fibers with metals in order to improve their properties. Mainly, PCS was modified with metal alkoxides or acetyl

acetonates (metal = Ti, Zr, Al, etc.).[100-102] Depending on the metal precursor used for the chemical modification, different reaction pathways of PCS with the metal precursor have been reported. Thus, titanium, zirconium, or aluminum alkoxides react with PCS upon alkane evolution and formation of Si-O-M units^[101,102]; whereas metal acetonates were assumed to react upon Si-M bond formation and release of acetyl acetone. [101] However, in both cases the chemical interaction between PCS and the metal precursors has not been clarified unambiguously. For instance, the modification of polycarbosilane with metal alkoxides was supported by spectroscopic data (such as FTIR or NMR spectroscopy) only in some cases^[102]; while in other cases no direct evidence for the formation of the Si-O-M units was obtained and the alkoxide was assumed to be physically incorporated within the PCS-based matrix (i.e., the formation of a blend consisting of alkoxide and PCS). [103] Also, in the case of the reaction of PCS with metal acetylacetonates, no direct evidence of the formation of Si-M units was provided (this was claimed only on the basis of the decrease of the amount of Si-H bonds in PCS; which is, however, not a strong argument). [101] Recently, a case study on the chemical modification of an allylsubstituted hydrido polycarbosilane with vanadium acetylacetonate indicated that the PCS reacts with metal acetylacetonate complexes upon formation of Si-O-M units, as also reported for the reactions of PCS with metal alkoxides. $^{[104]}$

Thermal treatment of the prepared single-source precursors was shown to lead in a first step to amorphous single-phase ceramics, which subsequently undergo phase separation and crystallization processes to furnish bi- or multi-phase nanocomposites (Figure 5). The conversion of the single-source precursors into ceramics usually consists of two steps: 1) crosslinking processes at low temperatures (100-400 °C) leading to infusible organic/inorganic networks; and 2) ceramization at temperatures up to 1000-1400 °C. Whereas mainly amorphous ceramics are obtained upon pyrolysis, subsequent annealing at high temperatures leads to (poly)crystalline materials. During cross-linking, the single-source precursors are converted into organic/inorganic materials. This process prevents the loss of low molecular weight components of the precursor as well as their fragmentation during ceramization, and thus increases the ceramic yield. Furthermore, the conversion of the single-source precursors into infusible materials prevents their melting during ceramization, while retaining the shape upon pyrolysis. The ceramization process of the cross-linked precursors consists of the thermolysis and evolution of their organic groups at 600-1000 °C. In this temperature range, the conversion of the preceramic materials into amorphous covalent ceramics (usually being single-phasic) occurs. The multifaceted mechanisms involved in the pyrolysis of preceramic polymers are not well understood yet. This is mainly due to the complex structure of the highly cross-linked polymeric materials and the resulting amorphous ceramics. The reactions which occur during pyrolysis can be investigated by solid-state NMR, FTIR, and Raman spectroscopy, as well as via thermogravimetric analysis (TGA) coupled with evolved gas analysis (EGA) (i.e., in situ mass spectrometry and FTIR spectroscopy). [89]

Synthesis and phase formation of silicon-based ceramic nanocomposites were studied extensively in the last decade. [87] Pyrolysis of suitable alkoxysilanes, Si(OR)₄, or polysiloxanes,

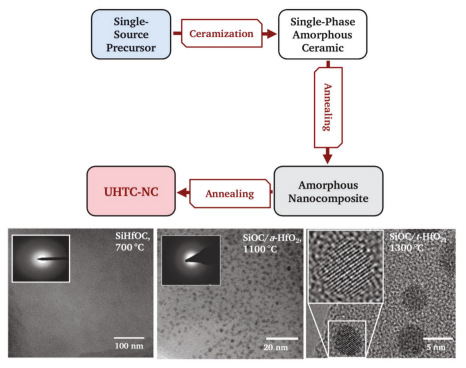


Figure 5. Typical preparative approach of polymer-derived ceramic nanocomposites starting from single-source precursors. [87–89]

[Si(R)₂-O]n, chemically modified with metal alkoxides was shown to give $MO_x/SiOC$ -based PDC-NCs, as reported for M = Al, Ti, Zr, Hf, etc. For M = Zr and Hf a single-phase SiMOC ceramic is obtained upon pyrolysis at rather low temperatures (ca. 700 °C), while at higher temperatures amorphous MO2 nanoparticles precipitate (800–1100 °C). Upon increasing the annealing temperature to 1300°C, MO2 nanoparticles crystallize, forming microstructures comprised of tetragonal zirconia/hafnia particles finely dispersed within an amorphous SiOC matrix. [87,105-107] When using other metals to modify the SiOC ceramic matrix, different and apparently contradictory results have been observed. Thus, in the mentioned case study, the phase composition of different SiMOC systems (M = Sn, Fe, Mn, V, and Lu) was investigated. [108] Depending on the metal, different ceramic phases formed. For M = Mn and Lu, MO_x/SiOC ceramic nanocomposites were formed, whereas other compositions revealed the formation of M/SiOC (M = Sn), $MSi_x/SiOC$ (M = Fe) or $MC_x/SiOC$ (M = V) upon pyrolysis^[87,108] (**Figure 6**). The strong effect of the precursor composition on the phase evolution upon ceramization reflects the reducing conditions during pyrolysis and annealing. Thus, the thermodynamic stability of the metal oxides generated during pyrolysis, particularly with respect to the system C-O, plays a crucial role. Based on thermodynamic data of the respective oxides, the phase composition of SiMOC/SiMCNO ceramics upon annealing at high temperatures can be predicted for different metals (Figure 2). The prediction agrees with the experimental results from this study and those reported in the literature for both SiMOC and SiMCNO ceramic nanocomposites. However, in addition to the stability of the oxides with respect to their reduction in carburizing conditions, some other aspects must be taken into account for predicting the phase composition of SiMOC/SiMCNO

nanocomposites, such as thermodynamic stabilization through conversion into silicates (for MO_x being stable with respect to carbothermal conversion into M) or into silicides or carbides (for MO_x not being stable against carbothermal reduction).

3. Synthesis of UHTCs and (Nano)Composites Thereof from Preceramic Polymers

3.1. Synthesis of Pure UHTC Phases from Polymeric Precursors

The synthesis of polymeric precursors used to access UHTC phases can be divided into two distinct categories: 1) methods using chemical modifications of metal alkoxides with, e.g., (poly) alcohols, (poly)acetates or other compounds such as acetylacetone, salicylic acid, etc.; and 2) methods using chemical modification of organometallic compounds such as metal-locene-type compounds or other metal complexes with alkyl, alkenyl, alkynyl, or aryl ligands.

Thorne et al. studied the reaction of Ti isopropoxide with various organic di-acetates leading to polymeric compounds, so-called "polymeric titanates" (**Figure 7**), which were converted into titanium carbide upon thermal treatment in Ar atmosphere at temperatures of $1000-1600\,^{\circ}\mathrm{C}$. All prepared materials contained specific amounts of excess carbon. Interestingly, despite the presence of excess carbon, the obtained TiC_x titanium carbide materials were sub-stoichiometric, with x=0.4-0.6.

Using the same principle, metal alkoxides may be used to prepare polymeric metallates upon reactions with di-

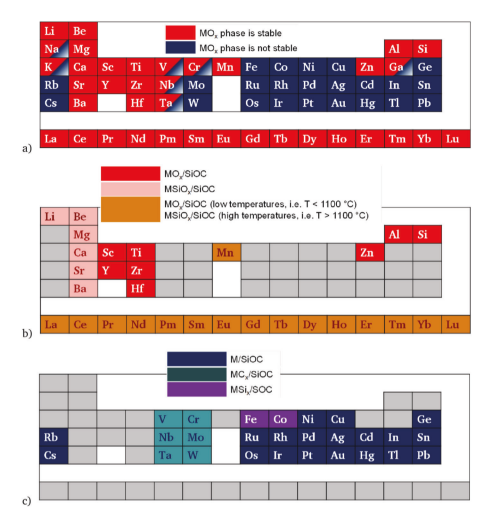


Figure 6. Thermodynamic control in Si–M–O–C-based ceramic nanocomposites: a) oxides of the red marked elements are stable against carbothermal reduction; whereas those of the blue marked elements get reduced in carburizing conditions; b) metals which upon incorporation into SiOC lead directly to silicate formation (pink); the orange marked elements induce firstly the precipitation of MO_x (at $T < 1100\,^{\circ}C$), followed by the formation of silicates at higher temperatures; modification of SiOC with the red marked elements leads to $MO_x/SiOC$ nanocomposites; c) the blue marked metals lead to metallic nanoparticles within the SiOC matrix; the modification of SiOC with the purple marked metals (e.g., Fe, Co) generates metal silicide phases; groups 5 and 6 transition metals (green marked) lead to carbide crystallization within SiOC matrix. [108]

alcohols,^[110–113] acetylacetone, or combinations thereof^[114–120] (**Figure 8**). Whereas in some cases the prepared polymers were structurally characterized to some extent, many other publications do not consider the molecular structure of the synthesized polymeric metallates and report rather on their conversion into metal carbide compounds upon pyrolysis at elevated temperatures (1000–1600 °C) in inert gas atmosphere.

The conversion of the polymeric metallates into metal carbides is considered to occur via the intermediate formation of metal oxide and excess carbon at temperatures ranging from 800 to $1100-1300\,^{\circ}\text{C}$; at higher temperatures, the metal oxide phase reacts with carbon and converts into metal carbide. Depending on the molecular structure of the polymeric metallates (i.e., nature and amount of organic substituents), various amounts of excess carbon can be generated in the resulting C/MC_x nanocomposites.

If the thermal conversion of the polymeric metallates is performed in reactive atmospheres, other compounds are accessible. For instance, Kurokawa et al. converted polymeric metallate precursors in ammonia atmosphere into metal nitrides MN_x (M = Ti, Nb, Ta). [122]

In a case study, the thermal conversion of a polymeric precursor consisting of polymeric zirconate (polyzirconoxanesal^[117]) modified with boric acid was shown to lead to a mixture of zirconia, boria, and segregated carbon at $1000-1200\,^{\circ}\text{C}$, which finally converted into phase-pure ZrB_2 at temperatures above $1400\,^{\circ}\text{C}^{[123]}$ (Figure 9).

A second preparative tool to produce UHTC phases from polymeric precursors is based on organometallic precursors. [124–126] Already in the 80s, titanocene derivatives $CpTiCl_3$, Cp_2TiCl_2 , $Cp_2Ti(R)Cl$, and Cp_2TiR_2 (Cp=cyclopentadienyl; R=alkyl, aryl) were used to prepare titanium carbide by gas-phase approaches (MO CVD) or thermal decomposition (see ref. [124] and references therein). Moreover, various sandwich compounds of group 4 transition metals $(R-Cp)_2MR'_2$ (R=H, CH_3 , $Si(CH_3)_3$; R'=Cl, alkyl,

$$Ti(O^{f}Pr)_{4}$$

$$+ \frac{H_{3}C}{C} - O - \frac{H_{2}}{C} - \frac{H_{2}}{R} - C - O - \frac{H_{2}}{C} - O - \frac{H_{2}$$

Figure 7. Schematic description of the synthesis of TiC from polymeric titanates. $^{[109]}$

aryl, NH₂, SH, OCN, SCN, etc. $^{[124]}$) were prepared and converted via thermal treatment in inert gas atmosphere into the respective metal carbides. It was shown that the decomposition of the molecular organometallic compounds occurs via formation of polymeric (macromolecular) intermediates, as shown in **Figure 10**. $^{[124]}$

Recently, the synthesis of Hf-containing polymeric precursors based on organometallic compounds and their high-temperature conversion into HfC/C composites was

reported. [125] Typically, Cp_2HfCl_2 was reacted with bifunctional dilithio diynes (e.g., 1,4-dilithio-1,3-butadiyne, or 1,4-bis-(lithioethynyl)-benzene) to generate the corresponding polymeric precursors. [125] Additionally, a dialkenyl-substituted hafnocene was polymerized upon thermal treatment in the presence of dicumyl peroxide as radical polymerization catalysts (Figure 11). [125]

Also, di-vinyl substituted zirconocene was reacted with borane to deliver a polymeric precursor which was thermally converted at $1000-1500\,^{\circ}\text{C}$ into a ZrC/ZrB2 nanocomposite. [126]

3.2. Synthesis of (Nano)Composites Consisting of UHTCs and Silica Formers

The relatively poor oxidation stability of pure UHTCs phases (especially at moderate temperatures, i.e., $500-600\,^{\circ}\mathrm{C}$ and $1200-1500\,^{\circ}\mathrm{C}$) pushed forward efforts to find different methods to improve their oxidation resistance. Thus, different approaches such as solid solutions, ternary diboride compositions, or the use of secondary phases were studied. Currently the most promising approach involves the use of additive phases (e.g., silicon carbide^[127]), which can alter the composition and behavior of the oxide scale.

The beneficial effect of SiC addition on the oxidation behavior of diborides has been explained to rely on the formation of a borosilicate glass layer on the oxidized ceramics. [15] For instance, extensive studies on the oxidation behavior of SiC-containing diboride materials indicate the formation of a silica layer at the beginning of the oxidation

$$(a) \qquad \qquad (RO)_{x_2} M \bigcirc R \qquad (RO)_{x_2} M \bigcirc R \qquad (RO)_{x_3} M \bigcirc R \qquad (RO)_{x_4} M \bigcirc R \qquad (RO)_{x_4} M \bigcirc R \qquad (RO)_{x_5} M \bigcirc R \qquad (RO$$

Figure 8. Synthetic access to polymeric metallates; a) based on metal alkoxides and di-alcohols $(R = alkyl; R' = C_x H_{2x})^{[113,121]}$; (b) based on metal alkoxides and acetylacetone^[118] as well as on metal salts and a combination of acetylacetone and a di-alcohol.^[117]

Figure 9. a) Synthesis of ZrB_2 upon high thermal conversion of $B(OH)_3$ -modified polyzirconoxanesal in argon atmosphere. [123]

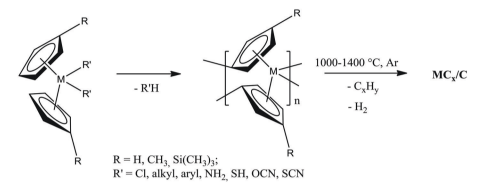
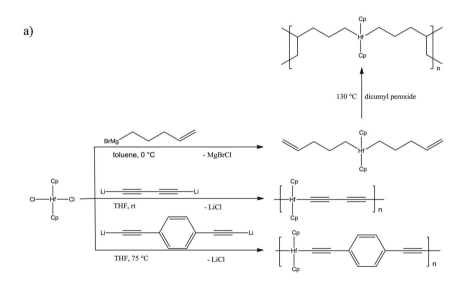


Figure 10. Use of sandwich metal complexes (M = Ti, Zr, Hf) as precursors for the synthesis of MC_x/C nanocomposites. Their thermal decomposition was shown to occur via formation of polymeric intermediates. [124]



b)

$$Cp_2ZrCl_2 + 2 H_2C = CHMgCl \xrightarrow{toluene, 0 \ ^{\circ}C} \xrightarrow{l} H_2C = CH_2 \xrightarrow{t} CH_2 \xrightarrow{l} H_3B \cdot S(CH_3)_2 \xrightarrow{l} H_3C \xrightarrow{l}$$

Figure 11. a) Synthesis of non-oxidic preceramic polymers for HfC starting from $Cp_2HfCl_2^{[125]}$; (b) synthesis of ZrC/ZrB_2 using a preceramic polymer synthesized from a divinyl-substituted zirconocene and borane. [126]

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process.^[127] Accordingly, the further oxidation is expected to be limited by the inward oxygen transport through the formed silica scale. Additionally, boria is formed as an oxidation product of the diborides and contributes to a significant decrease of the viscosity of the silica scale. Consequently, the oxygen transport through the borosilicate glassy scale is increased and at the same time boria partially vaporizes from the surface of the glass scale. A multilayer structure has been proposed for the oxidized diborides, which show a porous oxide layer (zirconia, hafnia) below the borosilicate glass layer. At high oxidation temperatures (i.e., beyond 1500 °C) an additional layer located between the porous oxide layer and the parent materials was analyzed. It consists of SiC-depleted diboride and was stated to relate to an active oxidation process of silicon carbide. Accordingly, the silicon carbide converts into gaseous silicon monoxide, SiO, which is transported through the porous oxide layer towards the surface, where it reacts with oxygen to form silica, SiO₂. [15,128–130]

In addition to silicon carbide as a silica former, silicon nitride, $^{[131,132]}$ molybdenum disilicide, $^{[133,134]}$ or other transition metal silicides $^{[135,136]}$ have been used. These systems exhibit similar oxidation behavior to that of MB₂/SiC composites.

3.2.1. UHTC (Nano) Composites with Low Content of Silica Former (i.e., 10–vol%)

Similar to the preparative methods presented in Section 4.1 for UHTC phases, the access to UHTC-based (nano)composites with typically 10–20 vol% silica former phase (e.g., SiC) was reported in the last decade using the same synthesis tools. Thus, various publications from the last decade report the synthesis of MC/SiC (M = Zr, Hf, Ta) using polymeric precursors based on polymeric metallates either blended or modified with poly(carbo) silanes. $^{[118,137-139]}$

For instance, a polymeric zirconate (polyzirconoxanesal^[117]) was mixed with boric acid and poly(methylsilylene-ethynylene)^[140] and was converted via thermal treatment in argon atmosphere at 1000–1400 °C into $ZrB_2/ZrC/SiC/C$ composites (Figure 12).^[141]

Similarly, ZrC/SiC, HfC/SiC as well as TaC/SiC were prepared from corresponding polymeric precursors based on polymeric metallates blended with, e.g., poly-(allylhydridocarbosilane). $^{[118,137-139]}$ More recently, a case study reported on the synthesis of HfC/SiC from precursors based on polymeric metallates modified with TEOS as a source for SiC. $^{[142]}$

A different approach was reported by Amorós et al., based on poly(dimethylsilane), PDMS, or poly-(methylhydridocarbosilane), PCS, which were mixed with Cp_2MCl_2 . The polymeric precursors, resulted from the thermal treatment under reflux condition, were then thermally converted into MC/SiC/C in argon atmosphere at temperatures in the range from 900 to $1400\,^{\circ}$ C. This synthesis methods allows adjusting the M:Si ratio (i.e., from 1:1 to 1:10) as well as the amount of free carbon (between ca. 15 and 50 wt%) in the prepared nanocomposites. As expected, the use of PCS led to an improved ceramic yield after pyrolysis, which relies on the presence of cross-linking active Si–H bonds in PCS. However, for both PDMS and PCS, rather low ceramic yields (10–25 wt %) after pyrolysis at 900 °C were reported (Figure 13). [143]

3.2.2. UHTC (Nano) Composites with High Content of Silica Former (i.e., 80-vol%)

In this part, we consider (nano)composite materials in which the UHTC phase is preferentially distributed in a silica former matrix such as Si_3N_4 , SiC, and more complex systems like Si–B–C–N. A general strategy is to modify precursors of the silica former phase at molecular scale by metalorganic compounds. This allows the generation of composites or nanocomposites made of (at least) one more phase in addition to the SiC, Si_3N_4 , or Si–C–B–N phase. Such materials often exhibit dramatically enhanced mechanical properties and/or structure stability. [87,144,145] Additional routes consist of 1) mixing polymers with metal fillers, [146] 2) blending two precursors of the desired composition, [147,148] 3) copolymerizing two monomeric precursors to form co-polymers. [149]

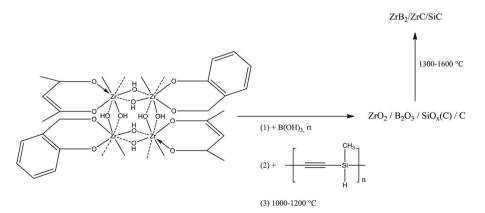


Figure 12. Synthesis of ZrB₂/ZrC/SiC from a polymer blend consisting of B(OH)₃-modified polyzirconoxanesal and poly(methylsilylene-ethynylene) upon high-temperature thermal treatment in argon atmosphere.^[141]

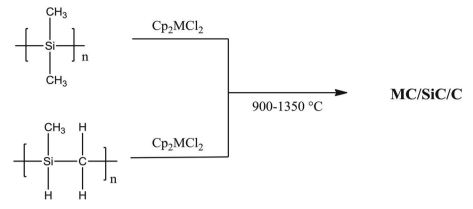


Figure 13. Synthesis of MC/SiC/C nanocomposites (M = Ti, Zr, Hf) from polydimethylsilane or polymethylhydridocarbosilane modified with metallocene dichlorides (Cp = cyclopentadienyl). [143]

(Nano) Composites with $Si_3N_4(C,O)$ Matrix as Silica Former. Riedel et al. described the reaction between TiCl4 and bis(trimethylsilyl)carbodiimide leading to a single source polymeric {[Ti(NCN)₂₁₃NCNSiMe₃} compound as precursor suitable to form a TiC_{0.3}N_{0.7}/Si-C-N nanocomposite. [150] Accordingly, the main process to design UHTC nanocomposites with Si₃N₄ and Si-N(C,O) matrices as silica former is based on the chemical modification of pre-formed preceramic polymers with low molecular weight species such as metalorganic compounds to form so called polymeric single-source precursors (SSP), in which a uniform chemical composition is established at the molecular scale. Then, the SSPs are converted in a first pyrolysis step under ammonia to reduce or even eliminate the carbon-based organic groups present in the polymer while a single-phase amorphous ceramic is generated at 1000 °C. The latter is subsequently heat-treated at higher temperatures under nitrogen to form the UHTC nanophase providing nanocomposites composed of MN nanocrystals (M = Ti, Hf) homogeneously distributed in an amorphous matrix. [151-155] The first report focused on the design of TiN/Si₃N₄ nanocomposites involved the reaction between a commercially available polysilazane, i.e., perhydropolysilazane (PHPS), and a metallorganic compound namely tetrakisdimethylaminotitanium (TDMATi) in a 2.5:1 molar ratio. The reaction formed a perhydropolytitanosilazane with a chemical formula of $[Si_{1.1}Ti_{0.4}C_{2.2}N_{1.1}H_{6.2}]_n$ and a molecular weight of 4200 g mol^{-1} [152] After ammonolysis at $-40\,^{\circ}\text{C}$, the polymer could be warm-pressed at 110 °C under 162 MPa uniaxial pressure. Subsequent pyrolysis of the formed body under ammonia (1000 °C) and nitrogen (1300 °C) produced bulk nanocomposites with 70 wt% yield and comprised of TiN nanocrystallites with an average diameter of 3.1 nm homogeneously embedded in an amorphous Si₃N₄ matrix. The in situ controlled growth of TiN nanocrystals associated with the absence of Si₃N₄ grain growth and the purity of the materials provided Vickers hardness as high as 25.1 GPa. Later, the same authors investigated the chemistry behind the process of preparing such nanocomposites to gain new insight into the reaction mechanisms of these materials and to understand the role of chemistry behind the processability and design of TiN/Si₃N₄ nanocomposites. [154] Based on complementary characterization tools including solid state NMR, elemental

analyses and infrared spectroscopy, two mechanisms have been proposed to govern the reaction between polysilazane and TDMATi. Mechanisms involve N–H and Si–H bonds present in polysilazanes as well as N(CH₃)₂ groups in TDMATi (**Figure 14**).

Detailed solid-state NMR studies in combination with quantitative thermogravimetry provided a comprehensive understanding of the polytitanosilazane-to-nanocomposite transformation process. Accordingly, a predominantly amorphous ceramic network is formed first at 1000°C under ammonia. Subsequent annealing in N2 at 1400 °C produces nanocomposites comprised of homogeneously dispersed TiN nanocrystals embedded in an amorphous matrix (Si₃N₄). The structural evolution above 1000 °C is closely related to 1) the chemistry of polysilazanes (precursors of the Si₃N₄ matrix), 2) the molar Si:Ti ratio fixed during the synthesis of the polytitanosilazane, and 3) the material shape (powder, mesoporous monoliths, or dense materials). [153] It appears that TiN is the first phase to nucleate at 1400 °C in such materials and the TiN crystallite growth mainly depends on the molar Si:Ti ratio fixed in the polytitanosilazane. Interestingly, the onset of the Si₃N₄ crystallization temperature in MN/Si₃N₄ nanocomposites is shifted upward with increase of the volume fraction of TiN in the ceramics. The Si₃N₄ crystallization was significantly reduced even after annealing at 1800 °C for the nanocomposite with the highest Ti content. Similar effects have also been observed by replacing titanium by hafnium to form HfN/Si₃N₄ nanocomposites from Hf-modified PHPS. In the latter, HfN nanocrystals are embedded in an amorphous SiN_x matrix after annealing at 1300 °C whereas full crystallization of both phases is found after annealing at 1600 °C. [155] Thus, it appears that the crystallization behavior of the Si₃N₄ matrix in MN/Si₃N₄ nanocomposites is governed by the chemistry of the polysilazanes and the Si:M ratio fixed at the polysilazane level.

By changing the chemistry of the precursors and using polyhydridomethylsiloxane (PHMS) as a matrix precursor and titanium tetra-n-butoxide as a precursor of the nanophase, nanocomposites made of TiN nanocrystals of approximately \approx 4 nm size homogeneously distributed in an amorphous SiOCN matrix have been generated at 1200 °C. [151] The photocatalytic properties of the nanocomposites with respect to dye degradation have been evaluated. The visible light photocatalytic activity

Figure 14. Reaction between NH and SiH units present in polysilazanes and NCH₃ groups present in TDMATi. [154]

of the synthesized nanocomposite resulted in a degradation efficiency of 64% and the hydroxyl radicals were found to be the active species.

(Nano) Composites with SiC(N) Matrix as Silica Former – MC/ SiC(N) and MN/SiC(N) (Nano) Composites: The design of (nano) composites with SiC(N) matrix as silica former has been well reported in the literature. The first strategy consists of using passive or active fillers mixed in the SiC(N) precursor. Whereas passive fillers such as ZrC powders do not interact with the SiC precursor upon pyrolysis in order to form additional phases in the SiC(N) material after the pyrolysis and annealing treatments, [156,157] active fillers do interact with preceramic polymers. As an illustration, Seyferth et al. investigated this approach to prepare Si-C-W systems using polysilazanes and W powders. [158] In the same study, Ti and Zr were combined with polysilazane and pyrolyzed under Ar at 1500 °C to form mixtures of TiN, ZrN, and SiC. This approach offers easy access to composite materials in a relatively large range of compositions according to the choice of both the metal and the organosilicon precursor. As an illustration, the pyrolysis of polycarbosilane (PCS) mixed with Ti, Zr, and Hf to 1500 °C in a stream of argon delivered MC/SiC composites. [159] Using a carbon-rich SiC precursor, i.e., [Me₂SiC≡C]_n, as a reagent and Ti, Zr, and Hf as active fillers, different MC/SiC compositions have been prepared. $^{\left[159\right]}$ In contrast, the pyrolysis of a Si-rich SiC precursor, i.e., $[(MeSiH)_{\sim 0.4}(MeSi)_{\sim 0.6}]_n$, under argon at 1500 °C led to TiC and Ti₅Si₃ phases. Riedel et al. mixed titanium powders with polysiloxane to form after pyrolysis to 1100°C mixtures of titanium carbide and metal silicide. [160] More recently, a series of studies has been reported on the dispersion of titanium powders in polycarbosilanes with the objective to form the well-known MAX phase Ti₃SiC₂. [161-164] All these works illustrate the simplicity of this polymer approach. However, the use of nanoparticles (diameter <100 nm) as active nanofillers in organosilicon polymers was not reported despite nanostructured materials can provide improved or new properties compared to their microstructured counterparts. In this context, the use of allylhydridopolycarbosilane (AHPCS) as a reagent and titanium (Ti) nanopowders as active fillers (Figure 15) has been described to generate TiC/SiC nanocomposites. [165]

During the pyrolysis under argon, Ti nanopowders reacted with carbon-based gaseous by-products (methane, ethylene, etc.) that evolved from AHPCS during its decomposition and/or with carbon radicals to form TiC as a unique phase after pyrolysis to $800\,^{\circ}$ C. The dominating crystalline TiC phase further crystallized in the temperature range $800-1000\,^{\circ}$ C while marking the Ti $_3$ SiC $_2$

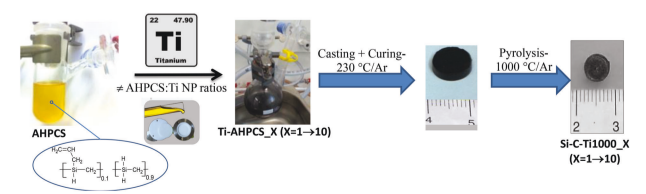


Figure 15. Overall process for preparing dense monolithic Si-C-Ti components. [165]

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and Ti₅Si₃ nucleation. ^[165] This procedure suggests that Ti and/or TiC reacted with the AHPCS-derived matrix to form the Ti₅Si₃ and Ti₃SiC₂ phases. Upon further pyrolysis above 1000 °C, monolithic TiC/SiC composites gradually formed at 1400°C. Most probably, free carbon identified in the samples played a key role in the formation of TiC and SiC from Ti₃SiC₂ and Ti₅Si₃ phases. Based on these studies, it appears that SiC precursors represent appropriate reagents to successfully generate MC/SiC composites via their mixture with metal (M) powders. However, the major disadvantage of this approach is that the processing may result in aggregation of the powders leading to compounds with an inhomogeneous distribution of the phases. Therefore, solution-based methods, in which the SiC and MC (M = Ti, Zr, Hf) precursors are mixed on the molecular scale, as well as single-source precursors (SSP) are preferred although more expensive.

The modification of SiC with Ti, Zr, and Hf has been proposed since the 80s to design (nano)composites with a SiC matrix as a silica former. [12,101–103,166–176] Since that time, different synthetic strategies are described in the literature to form MC/SiC(N) (nano)composites, by using either mixed precursors (or blended polymers), or by inserting the metal in the polymer at the level of the macromolecule (i.e., chemical modification) or directly in the monomer (*single-source* precursors).

Mixing of two polymers respectively with M and Si in their structures represent an easy access to MC/SiC composites. ^[177] Following this route, a polycarbosilane and a polyzirconobutanediol were blended in toluene to be pyrolyzed at 1500 °C under argon. ^[178] The distribution of the ZrC phase as well as the size of the ZrC crystals correlated with the time of the drying method.

Another route to access MC/SiC composites was based on the addition of a crosslinking agent, namely divinylbenzene (DVB), to the blended polymers consisting of polyhafnoxanesal ([-Hf-O-Hf-]_n), and PCS.^[139] HfC was generated at high temperature through a carbothermal reduction of the Hf-O-C phase by the excess carbon in-situ formed from DVB and PCS during pyrolysis. After annealing at 1600 °C under argon, HfC/SiC powder with diameter less than 100 nm and a composition of ca. 13.5 vol% HfC, 64.5 vol% SiC, and 22 vol% excess carbon was formed (**Figure 16**).

A blend composed of a polyzirconoxanesal (Zr source), a phenyl-terminated polysilane (Si source), and bisphenol-A type benzoxazine was reacted to lead to a multi-element precursor, with ring-opening, condensation, and addition polymerizations. This method was used to fabricate ZrC/SiC ceramics, after a suitable pyrolysis of the complex polymeric network up to 1600 °C, under nitrogen atmosphere. The ceramic yields ranged from 40 to 65%, and the

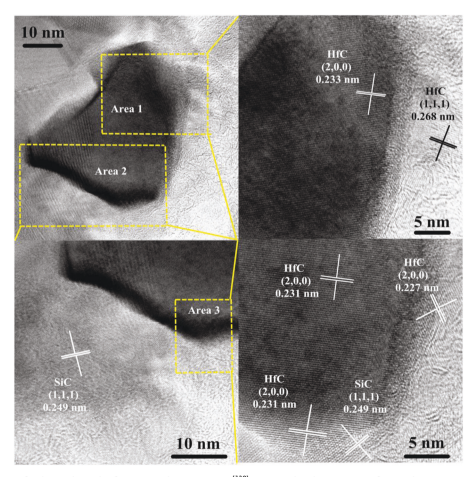


Figure 16. TEM images of polymer-derived HfC/SiC powders at 1600 °C^[139] (reprinted with permission from Wiley, 2018).

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microstructures revealed well-distributed Zr, Si, and C elements in the final material.

The chemical modification of a selected polysilane or a polycarbosilane containing reactive groups like SiH sites with a metal-containing precursor is probably the most common approach which has been explored to prepare MC/SiC composites and nanocomposites. Starting from polymethylsilane, Wang et al. prepared a series of polyzirconomethylsilanes namely PZMS by adjusting the weight ratio between the Zrbased compound, i.e., $Cp_2Zr(CH = CH_2)_2$ and the polymethylsilane. [180] The pyrolysis of PZMS resulted in a ceramic yield of 78.4 wt%. Crystalline ZrC and SiC phases were well distributed in the composite prepared at 1600 °C with the content of ZrC as high as 54.6% by weight. It was shown that the introduction of ZrC prevented the fast growth of crystalline β -SiC. Based on their work focused on the design of ZrC/SiC composites^[174] and on the pioneering report of Latorre et al., [143] Yu et al. reported the preparation of TiC/SiC composites through the reaction of dichlorobis(cyclopentadienyl)titanium (Cp₂TiCl₂).^[181] The reaction was performed with three Cp2TiCl2:AHPCS ratios, i.e., 1:3, 1:2, and 1:1, followed by a pyrolysis in an argon atmosphere up to 900 °C and subsequent annealing at 1600°C in an argon flow. It was shown that Cp₂TiCl₂ played a double role: 1) The cross-linking of the hybrid precursors was significantly catalyzed by Cp2TiCl2 and Si-Si dehydrocoupling and hydrosilylation effectively occurred during mixing. 2) Cp2TiCl2 reacted with AHPCS through reaction of SiH and TiCl groups forming HCl. These two functions of Cp₂TiCl₂ significantly improved the ceramic yield of the polymer (80.8 wt%) compared to prisitine AHPCS (60.5 wt%). The chemical composition as well as the microstructure could be tailored by the Cp2TiCl2:AHPCS ratios. AHPCS was also used to react with borazine via hydroboration before adding titanium butoxide Ti(OBu)₄ to be cross-linked at 170 °C. [182] By fixing B and excess C, Ti allowed to form crystalline TiB₂ and TiC phases in addition to the crystalline SiC phase through pyrolysis at 900 °C and annealing at 1800 °C under argon. Polycarbosilanes may also react with alkene functional groups present in metalcontaining molecules through hydrosilylation. This approach was developed by Wang et al. using Zr-containing molecules.[183,184] The authors performed functionalization of

zirconocene dichloride ((Cp_2ZrCl_2) by using an organomagnesium reagent (i.e., allyl- ($CH_2=CHCH_2MgCl$) or vinyl-magnesium chloride ($CH_2=CHMgCl$)). Then, the as-obtained diallylbis(cyclopentadienyl)zirconium (DACZ) depicted in **Figure 17** could react with a low molecular weight polycarbosilane to give a soluble crosslinked preceramic polymer labelled PCS-DACZ via free radical polymerization. [184] The PCS-DACZ displayed a relatively high ceramic yield of 80 wt%. ZrC/SiC/C nanocomposites were retrieved after a pyrolysis at 1400 °C under argon atmosphere, displaying an excellent high-temperature resistance up to 2000 °C with well-dispersed ZrC nanoparticles.

This strategy was also investigated with a zirconocene dichloride functionalized with vinyl groups to generate ZrC/SiC ceramics with a high Zr content. [184] The ceramics showed a good resistance to oxidation in air up to 1400 °C. In parallel to the reactivity of alkene groups, other methods were attempted involving N- or O-containing complexes of zirconium. For example, the two-step reaction between allylamine (C₃H₇N), zirconium tetrachloride (ZrCl₄), polymethylsilane (PMS), and CH₂ = CHCH₂MgCl led to a soluble ZrC/SiC precursor. The composite was isolated in a 49% yield at 1000°C, and the material exhibited less than 2 wt% of N in its structure. $^{[185]}$ In the field of polymer-derived ceramics, oxygen containing-zirconium complexes are often used. Reacting with polycarbosilanes, they can lead to ZrC/SiC ceramics after a suitable heat-treatment. In this way, Devasia et al. modified polycarbosilanes with zirconium acetylacetonate (Zr(Acac)4) to synthesize polyzirconocarbosilanes. [186] The authors introduced different Zr.Si ratio and showed that the ceramic yield of the precursors was improved by the introduction of Zr, to reach 76 wt% at 1200 °C, under argon. HRTEM analyses were undertaken on the composites heattreated at 1650 °C, revealing the presence of an oxide-free ZrC/ SiC composite.

In the case of hafnium, an earlier work reported on the synthesis of a precursor suitable to form HfC_xN_{1-x}/SiC nanocomposites.^[187] The procedure involved the reaction between AHPCS, and tetrakis(dimethylamido)hafnium (Hf(N-(CH₃)₂)₄, TDMAH) according to **Figure 18**.

The functionalization of AHPCS was confirmed by IR and NMR spectroscopy. Amorphous SiHfCN-based ceramic powders were then densified using spark plasma sintering (SPS) at

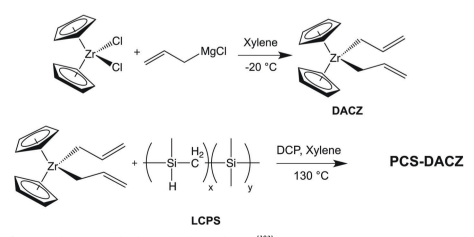


Figure 17. Synthesis of DACZ and reaction with a low molecular weight PCS.[183]



Figure 18. Modification of AHPCS with Hf(NEt₂)₄ leading to two silicon environments in the resulting preceramic polymer (Reproduced with permission^[187]; we acknowledge the Royal Society of Chemistry for the reproduced material).

2200 °C, and led to dense monolithic HfC_xN_{1-x}/SiC nanocomposites, highlighting homogeneously embedded hafnium carbonitride ($HfC_{0.83}N_{0.17}$) grains in a β -SiC matrix, encapsulated by in situ formed carbon layers. These materials were then characterized in detail regarding their dielectric properties, ^[188] oxidation resistance, ^[189-191] and laser ablation behavior. ^[192] HfC_xN_{1-x}/SiC nanocomposites display improved laser ablation resistance compared to pure SiC-based monoliths ^[192] and are especially suited for electromagnetic shielding in harsh environment. ^[188,190] It was reported that the additional incorporation of tantalum (Ta) in the HfC_xN_{1-x} phase leads to an enhanced sintering ability of the oxide scale and to an improved oxidation resistance via a short pre-oxidation step at temperatures sufficiently high to activate the formation of silica. ^[189,192]

The last – but highly important – method to reach multielement polymers with metal from the group 4 elements is the molecular synthesis of *single-source* precursors. This route is very useful especially in order to control the chemical composition of the preceramic precursor, and *in fine* of the ceramic, at the molecular scale. Upon salt metathesis reactions of zirconcene dichloride, methylphenyldichlorosilane, and dilithioacetylene, Cai et al. synthesized a ZrC/SiC nanocomposite precursor according to **Figure 19**.^[193]

Thus, the prepared polymer dissolves well in different solvents, such as THF, toluene, and chloroform etc. The obtained polymer could be converted to nanosized ZrC/C/SiC composites at a relatively low temperature, i.e., 1400 °C with a ceramic yield of over 52 wt%. Starting with oxide compounds, Li et al. detailed the synthesis of a polymer with Si and Zr in its structure. [194] They used the reaction of hydrolysis/condensation between tetraethoxysilane (Si(OC2H5)4 and zirconium oxychloride octahydrate (ZrOCl₂'8H₂O) in the presence of phenol, paraformaldehyde and acetylacetone. The as-obtained precursor yielded ZrC/SiC/C ceramics after pyrolysis up to 1800°C. However, SiO2 and ZrO2 phases were still present in the powders. Focusing on the titanium chemistry, Liu et al. used the incorporation of both Ti and Si in the backbone of a hyperbranched polycarbosilane labelled HPCS derived from Grignard coupling of a mixture made of trichloro(chloromethyl) silane (Cl₃SiCH₂Cl), Cp₂TiCl₂, and dichloromethyl-(chloromethyl)silane ($Cl_2Si(CH_3)CH_2Cl$) followed by LiAlH₄ reduction according to Figure 20. [195]

The thermal decomposition of the final liquid hyperbranched structure denoted as HPTiCS indicated a ceramic yield of 76 wt% at $1200\,^{\circ}$ C (higher by around 31% than that of the ceramic yield measured with the pure hydridopolycarbosilane) and led to TiC/SiC composites.

Figure 19. Synthesis of a Zr and Si containing poly-yne polymer used as a single-source precursor for the synthesis of ZrC/SiC nanocomposites. [193]

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Figure 20. Two-step synthesis of a polymeric single-source precursor containing Ti and Si mixed at molecular scale. [195]

An original approach was recently described, using organolithium reagents instead of organomagnesium compounds. Moreover, the reactivity of the cyclopendatienyl groups from the zirconocene dimethyl ($Cp_2Zr(CH_3)_2$) was investigated this time. Wang et al. synthesized three preceramic polymers from an intermediate dilithiozirconocene complex as a key Zr-Li salt obtained by the reaction of $Cp_2Zr(CH_3)_2$ with tetramethylethylenediamine (TMEDA) and butyl lithium (n-BuLi) according to Figure 21. $^{[196]}$

Di- and trifunctional chlorosilanes were employed to react with the Zr-Li salt. After removal of LiCl by filtration, the respective polymers were obtained (Figure 17). The polymer containing a vinyl group attached to the silicon atoms displayed the highest ceramic yield of 77 wt% under argon atmosphere at $800\,^{\circ}\text{C}$. Annealing experiments from 1000 to 2000 $^{\circ}\text{C}$ showed the crystallization from zirconium silicate (ZrSiO_4) and zirconia (ZrO_2) at lower temperature to ZrC at higher temperature. The prepared micrometer-sized spheres were shown to possess interesting electromagnetic wave absorbing capacity. $^{[196]}$

It is also worth mentioning two other studies using the monomer route: Firstly, bis(cyclopentadienyl)hafnium dichloride (Cp₂HfCl₂) and trans-1,4-dibromo-2-butene were reacted in the presence of magnesium powder in THF to yield a macromolecular HfC precursor. A chain termination with chloromethyltrimethylsilane ((CH₃)₃SiCH₂Cl) is then added to introduce silicon into the structure of the polymer.^[197] This preceramic precursor exhibited a good solubility in common organic solvents, and pyrolysis at 1600 °C resulted in HfC/SiC nanocomposite ceramics. Secondly, a polycarbosilane was reacted with zirconocene at the end of the silicon polymer chain.^[198] Zhang et al. used a zirconocene as a catalyst with

metallic sodium (Na) for an insertion of 1-methylsilene ($H(CH_3)$ $Si = CH_2$) into a preceramic precursor. The reaction product was denoted as polyzirconocenecarbosilane.

Metallic sodium was used to in-situ obtain 1-methylsylene, which polymerizes spontaneously at the Na-surface to PCS with zirconocene at the chain end groups. The thermal decomposition of the polymer with narrow molar-mass dispersity ($M_{\rm w}/M_{\rm n}$) led to ZrC/SiC ceramic composites with a ZrC concentration of about 15 wt%. This method was extended to titanocene and hafnocene, with apparently attractive rheological properties for fiber melt spinning.

In 2012, Li et al. reported mixing a polyzirconoxanesal with boric acid and a poly(methylsilylene)ethynylene to produce ZrB₂/ZrC/SiC composites by pyrolysis at 1400 °C under argon. [141] Following this study, He et al. reported the design of ZrB₂/SiC nanocomposites via a precursor route based on a similar simple one-pot method. [199] In this approach, a ZrB₂ precursor and a polycarbosilane (SiC precursor) were mixed in the desired ratio to form a molecular hybrid compound as a ZrB₂/SiC precursor with a targeted ZrB₂-to-SiC volume ratio of 25%. After pyrolysis at 1300 °C, a ceramic yield close to 61 wt% has been measured and t-ZrO₂ and β-SiC phases were identified exclusively by X-ray powder diffraction. Above this temperature, reaction of ZrO2 with B2O3 and C occurred and led to the formation of the ZrB2 phase along with SiC. Thus, after pyrolysis at 1700 °C for 4 h, a nanocomposite made of ZrB₂ nanoparticles embedded within a SiC matrix was generated, although ZrC was identified as an additional phase present in the powders. Later, the same authors performed spark plasma sintering (SPS) of these powders by changing the ZrB2-to-SiC volume ratio from 6.4 to 61.5 vol% and investigated the mechanical properties of the as-prepared pieces. [200] SPS was performed at 1600 °C for

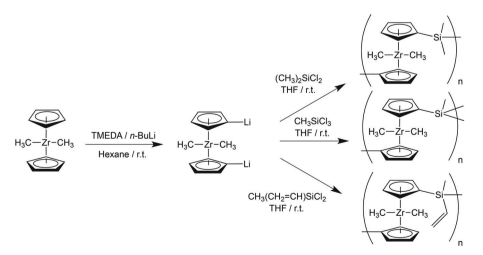


Figure 21. Synthesis of preceramic polymers via the reaction of dilithiozirconocene complex with various chlorosilanes. [196]

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30 min under an argon atmosphere and a uniaxial pressure of 35 MPa. As expected, the Vicker's hardness of the monolithic samples increased with the increase of the density of the materials from 7.42 ± 0.3 GPa (relative density of 87.6%) to 17.54 ± 0.9 GPa (relative density of 99.1%).

(Nano) Composites with Si-B-C-N Matrix as Silica Former. Adding boron to Si-C-N ceramics, i.e., leading to silicon boron carbonitride (Si-B-C-N) ceramics, improves the thermal stability of the amorphous structure of Si-C-N ceramics in terms of crystallization and decomposition.^[201] This phenomenon is related with the strong local covalent bonds associated with the presence of boron in SiBCN. As a consequence, the carbothermal reaction of Si₃N₄ occurring in Si-C-N ceramics is shifted to higher temperature. The use of this amorphous inorganic network appears also to be beneficial for the high temperature creep behavior of these multicomponent ceramics. (Nano)composites with a Si-B-C-N matrix have been prepared following the synthesis process implemented for dense SiC/ HfC_xN_{1-x}-based ultra-high temperature ceramic nanocomposites. [187] A commercially available polysilazane (HTT1800 (now Durazane® 1800), Merck)) was modified with Hf(NEt₂)₄ and BH3 · SMe2 (Figure 22) and subsequently cross-linked and pyrolyzed.[202]

The presence of Hf and B within the molecular structure of the polysilazane leads to low-temperature phase separation processes within the resulting Si-Hf-B-C-N. Depending on the annealing atmosphere, HfC/HfB2/SiC (annealing in argon) and HfN/Si₃N₄/SiBCN (annealing in nitrogen) nanocomposites were obtained. The Si-Hf-B-C-N precursor could be warm-pressed after cross-linking at 250 °C then pyrolyzed and annealed to 1300 °C to form dense pieces. [203] The oxidation behavior of these bulk samples (and powders) was studied at 1200-1400 °C in air. The measurement of the surface-specific weight gain as a function of the isothermal oxidation time showed that both, powder and monolithic samples, exhibited a parabolic oxidation behavior characterized by low parabolic rates. Observations of the oxide scale that formed on the monoliths at different oxidation temperatures revealed that a continuous oxide scale consisting of borosilicate, silica (cristobalite), m- and t-HfO2 formed at temperatures below 1300 °C, limiting the capacity for oxygen transport. At temperatures above 1300 °C, oxide evaporation increased and the oxide scale became discontinuous and consisted of silica, HfSiO₄, and m- and t-HfO₂, and large pores were formed in the bulk. Alternatively, the molecular Si-Hf-B-C-N precursor was cross-linked then pyrolyzed at 900 °C to deliver powders that could be sintered by SPS at 1850 or 1950°C into dense monolithic samples. [204] The high hardness and high elastic modulus of the dense pieces highlighted superior mechanical properties as compared to other polymer-derived ceramics, which rely on their phase compositions and their fine microstructure. The investigation of the oxidation of the prepared dense ceramic nanocomposites at high temperature revealed that the parabolic oxidation rates of Si-Hf-C-N were comparable to those of ultra-high temperature ceramics such as metal-diboride/SiC composites, whereas the parabolic oxidation rates of Si-Hf-B-C-N were several orders of magnitude lower than those reported for UHTCs and even lower than values reported for SiC. Additionally, the Si-Hf-B-C-N precursor could be used to infiltrate carbon fiber preforms (C_f) and form after curing and pyrolysis C_f/Si-Hf-B-C-N composites. [205,206] These ceramic matrix composites (CMCs) were tested regarding hydrothermal corrosion and laser ablation tests. The C_f/Si-Hf-B-C-N composites exhibited enhanced resistance towards hydrothermal corrosion as compared to $C_f/Si-C-N$ due to the improved kinetics upon Hf and B incorporation. Additionally, a tight C_f/matrix interface (which is rather disadvantageous for mechanical behavior) was found to be beneficial for improved corrosion behavior in C_f/Si-Hf-B-C-N composites.^[205] Various ablation mechanisms occurred in three distinct ablation regions, as detected by the study of the evolution of microstructure and phase composition of the CMCs by scanning electron microscopy (SEM) and energy dispersive spectroscopy (EDS)^[206]: 1) in the central region, the erosion of C_f/Si-Hf-B-C-N is controlled by the sublimation of the components of the CMCs; 2) in the transition region, the degradation of C_f/Si-Hf-B-C-N is dominated by the passive

Figure 22. Modification of HTT1800 with Hf(NEt₂)₄ and BH₃·SMe₂ (Reproduced with permission.^[202] Copyright, ACS, 2014)



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oxidation of the Hf-containing components and the active oxidation of the fibers and Si-containing components; 3) in the outer region, the pronounced degradation process of C_f/Si-Hf-B-C-N is the passive oxidation of the Si-containing components. It has been found that the ablation mechanism and the microstructure evolution of CMCs are mainly dependent on the local temperature which directly depends on the distance from the laser spot and the thermal conductivity of the carbon fibers. Similarly, a hafnium chloride compound has been used to modify a liquid polyborosilazane containing borazine rings as bridges between two monomeric silazane units. [207] After pyrolysis at 1100 °C in an argon atmosphere, the final material consisted of HfN nanoparticles embedded in a Si-B-C-N matrix. The HfN nanoparticles distributed uniformly in the Si-B-C-N matrix provided an excellent oxidation resistance: the HfN/Si-B-C-N nanocomposite was shown to be more oxidation-resistive with a weight loss of 2.8 wt% at 1500 °C in air, compared to a weight loss of 5.4 wt% for Si-B-C-N ceramics under the same condition.

The addition of Zr in polymer-derived Si–B–C–N ceramics through the modification of a liquid polyborosilazane with Cp₂ZrCl₂ according to different weight ratios has been also reported. In particular, fibers were prepared from this polymer to be cured with a BCl₃/N₂ mix flow gas, then pyrolyzed and annealed at 1400 °C under Ar. The Zr-modified Si–B–C–N ceramic fibers were amorphous after pyrolysis at 1200 °C. As the temperature was increased to 1400 °C, crystalline phases, namely ZrC_xN_y , ZrB_2 , and Si_3N_4 were identified. Zr was also shown to improve the oxidation resistance of the Si–B–C–N ceramics as reported for the Hf counterpart.

4. Conclusions and Perspectives

The polymer-derived ceramic (PDC) route represents a unique preparative access to UHTCs and their derived (nano)composites containing silica formers. These materials possess adjustable phase compositions and microstructures according to the fact that the chemical composition of the materials is controlled at a molecular scale. Thus, two strategies may be applied to deliver UHTC: 1) methods using chemical modifications of UHT-metal containing alkoxides with, e.g., (poly)alcohols, (poly)acetates, or other compounds such as acetylacetone, salicylic acid, etc.; and 2) methods using chemical modification of organometallic compounds such as metallocene-type compounds or other metal complexes with alkyl, alkenyl, alkynyl, or aryl ligands. In order to make accessible UHTC-containing nanocomposites (with either low or high content of silica former phase), a Sicontaining precursor is in general added to the UHTC precursor forming either precursor blends or, in the case of chemical reactions, single-source polymeric precursors which are further pyrolyzed at elevated temperatures (\gg 1000 °C). It may be mentioned that typically polymeric blends are used to prepare nanocomposites with low volume fractions of silica former phase (i.e., 10-20 vol%); whereas single-source precursors are highly suitable for preparing

nanocomposites with high content of silica former phase (i.e., 80-90 vol%).

There are obvious advantages of the polymeric-precursorbased synthesis methods of UHTCs and their nanocomposites, among them, e.g., nanoscopic particle size, high purity materials, relatively low synthesis temperatures. However, specific issues related to the use of polymeric/organometallic precursors should be also mentioned such as relatively high synthetic efforts (especially for the single-source precursors) or, in some cases, the environmental sensitivity of the polymeric precursors (i.e., for hydrolysis, oxidation processes). The industrial potential of UHTCs and derived (nano)composites is nevertheless evident, despite the PDCs process needs to be further developed in details for industrial and technological applications, e.g., raw materials availability, process scalability, etc. In particular, following aspects should be intensively addressed in the future in order to provide a rational design of UHTCs and their derived (nano)composites with tailored compositions, microstructures and properties: 1) a straightforward synthetic access to single-source precursors with a detailed knowledge on their chemical compositions and molecular architectures; 2) a fundamental understanding of the polymer-to-ceramic conversion process; 3) polymer chemistry - rheology correlation to develop the processability of these polymers and design complex textures and morphologies of materials; 4) joint and tailored experimental and computational studies are necessary in order to assess the properties of the UHTCs and their derived (nano)composites and to understand how the microstructural features of the materials affect their properties; and 5) detailed short and long-term analysis of their high-temperature behavior and mechanical properties (hardness, Young modulus, fracture toughness, etc.). It is anticipated that polymer-derived UHTCs and their derived (nano)composites have the potential to provide specific functionalities as well as improved and/or extended properties to be applied in particular in the societal relevant fields of energy conversion and future aeronautics and reentry systems.

Acknowledgements

The authors team is grateful to Prof. Yanchun Zhou (Aerospace Research Institute of Materials and Processing Technology, Beijing, China) for encouraging to compile the present review on polymer-derived UHTC systems. EI and RR thank the German Science Foundation (DFG, Bonn, Germany) for financial support (project "Micropatterned polymer-derived ceramic catalysts and sensors"). EI furthermore acknowledges the EU COST Action CM1302 (Smart Inorganic Polymers) and the Heisenberg Program of DFG. PK gratefully acknowledges support by the National Science Foundation (CMMI-1634448 and OISE 1743701).

Conflict of Interest

The authors declare no conflict of interest.

Keywords

physical properties, preceramic polymers, refractoriness, synthesis, thermodynamics, UHTC (nano)composites, ultra-high temperature ceramics (UHTCs)

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Received: March 13, 2019 Revised: May 1, 2019 Published online:

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