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SiC, from Amorphous to Nanosized Materials, 
the Exemple of SiC Fibres Issued of 
Polymer Precursors

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1. Introduction

Silicon carbide materials are interesting because of their high thermal stability, high thermal conductivity, extreme hardness and good electrical properties. Some of these properties are directly related to the SiC structure that alternates the Si and C atom layers. SiC have applications as single crystal wafer (wide-gap semiconductor for high power device), as air, spacecraft and nuclear plant thermostructural composites because it is possible to prepare the materials in various forms (fibre, matrix and composite) from organic liquid or vapour precursors. Thus it is possible to adjust the crystallinity, from the amorphous to the crystalline state, including intermediate nanocrystalline state. The later state guaranties the optimal mechanical properties. Non-destructive Raman microspectrometry intrinsically probes the matter at the subnanoscale and offers a "bottom-up" approach that is especially efficient for the analysis of ill-crystallised and nanostructured materials. The great advantage of Raman spectroscopy is first to make possible the recording of series of spectrum by automatic mapping. However, because the Raman intensity depends on the polarisability tensor derivative, the scattered intensity - and the sensitivity - varies order of magnitude with the nature of the chemical bond. Comparison with transmission electron microscopy (TEM) is in general required to discriminate if the phonon coherence length is determined by the domain or the grain size. Both techniques are efficient to identify the different polytypes as well as the nature and degree of the disorder.

2. About the chemistry

The properties of the materials are determined by their chemical bonds and the arrangement of the latter. The uniqueness of silicon carbide properties arises from the nature of the bond. Si-C bond as Si-O and Si-N ones are among the strongest chemical bonds in the matter (Bond dissociation energy > 350 KJ/mole). Consequently, this gives a very good chemical and thermal stability as well very high melting temperature, high mechanical properties and hardness. Silicon atoms form with C, N and O atoms the same tetrahedral coordination that allows very intimate mixture (oxycarbides, oxynitrides, etc.). Furthermore, a hydrogen atom can establish a bond with these oxygen, nitrogen and carbon atoms. This offers a large versatility of preparation routes, including polymerisation process. Consequently, because
silicon and carbon atoms form easily X-C and X-H bonds, a large variety of liquid or
gaseous hybrid/organic precursors are available (Mc Diarmid, 1961; Fritz et al., 1965; Corriu &
Lanneau, 1970; Lucke et al., 1997) and the control of their reactivity, viscosity/diffusivity
dilution, etc. gives rise to a variety of preparation routes leading to monoliths, films, fibres,
composites, crystal,..., from the amorphous state to the single crystal, including
nanomaterials (Gough & Kern, 1967; Parlier & Colomban, 1996; Naslain, 2004).
The mix of atomic orbital led to an intermediate electric behaviour between that of carbon (a
semi-metal) and silica (an insulator), the electronic gap is intermediate (2-3 eV) and the
material is a semiconductor. Since the diffusion coefficient of oxygen atom in silica is very
low (due to the very covalent character of the Si-O bond), the oxidation is blocked in a very
large temperature range (except in water-rich atmosphere when volatile species are formed)
that gives a good stability to Si(X)n compounds in air. The latter properties are searched for
thermomechanical applications: the nanosized state combines the advantage of amorphous
(homogeneity, lack of interface/phases) and crystalline (high density of bonds per volume
unit) phases. Very high thermal conductivity contributes well to the unique properties of
silicon carbide. Since nothing is perfect, the building of the SiC with a single type of bond
makes that orientation disorder is possible and gives rise to two types of Si-C bilayer
stacking. Their combination leads to series of structures: the polytypes and it is difficult to
prepare materials without coexistence of the different phases. This behaviour is observed for
all structures which are built up by corner-sharing tetrahedron. They can be regarded as
consisting of stacked bi-layers.
The availability of volatile/liquid silanes led to the preparation of SiC materials in various
forms (McDiarmid, 1961; Fritz et al., 1965; Gough & Kern, 1967; Lear & Khan, 1970; Corriu &
Lanneau, 1970; Kumar & Litt, 1988; Schmidt et al., 1991): films/coatings (Rynders et al.,
1991), fibres (a SiC coating is deposited on a carbon fibre core (Hurwitz et al., 1991; Tanaka
et al., 1995; idem, 1996; Popovska et al., 1997), monoliths and fibre-reinforced composites
(Yajima et al., 1980; Tanaka & Kurachi, 1988; Bouillon et al., 1991 a & b; Monthioux et al.,
1990; idem, 1991; Delplancke et al., 1991; Lee & Yano, 2004). The growth rate was slow and
for the preparation of thick monoliths or composites weeks are required with sometimes an
intermediate machining of the parts in order to avoid the closure of the porosity that occurs
at the sample surface, avoiding the completion of the core densification. This technology
was used to prepare important parts of civil and military launch vehicles, missiles as well as
aircraft engines (Parlier & Colomban, 1996; Mouchon & Colomban, 1996; Naslain, 2004) and
extended to the preparation of other Si-based materials such as nitrides (Durham et al.,
1991).

In the 80s Japanese groups developed liquids with appropriate viscosity (polycarbosilanes:
PCS) as precursors for SiC fibre (Okamura et al., 1985). The difficulty concerned the
production of a form of polycarbosilane which would have the "good" viscosity to be spun
and then converted in small-diameter ceramic filaments. The small diameter (~10 µm) is
required to permit the bending of the fibre during the weaving process and hence to prepare
a variety of textile fabrics: satin, taffeta, etc. (Mouchon & Colomban, 1995; idem, 1996). It
was found that very high molecular weight was not necessarily best and polymers with
molecular weights of around 1500 were used for commercial production. Then the progress
allowed preparing more complex compositions (Si/TiC, etc. (Stewart et al., 1991; Hasegawa
et al., 1991; Burns et al., 1992; Hurwitz et al. 1993)) under more complex states (Tanaka
et al., 1988; Langguth, 1995). The viscosity was adapted for an easy infiltration of the precursor.
in the porous performs (Greil, 1995; Parlier & Colomban, 1996; Lucke et al., 1997; Greil, 2000) and the technology was extended to multicomponent oxide precursors with the development of sol-gel routes (Colomban, 1989; Colomban & Wey, 1997; Colomban, 2005). A big problem was the achievement of the right stoichiometry: silicon excess was very detrimental because of the relatively low temperature melting of silicon; on the other hand, carbon excess, not detrimental from the thermomechanical point of view, led to a great reactivity to air and to high electrical conductivity (Chauvet et al., 1992a; idem, 1992b; Mouchon & Colomban, 1996). However, the polymer precursors are linear macromolecules (1D) but SiC synthesis requires a 3D crosslinking. This was first achieved by Si-O-Si bridges but the oxygen atoms react with carbon excess at ~1200-1300°C (Karlin & Colomban, 1997; idem, 1998). The use of gamma irradiation for the crosslinking allows approaching better the stoichiometry and the thermal stability of SiC materials issued from polymeric precursor was increased (Ishikawa et al., 1991; Yang et al., 1991; Torecki et al., 1991; idem, 1992; Ishikawa, 1995; Ishikawa et al., 1998). Residual hydrogen atoms evolve above 1000°C whereas the free carbon created by the destruction of the methyl groups reacted with the Si-O group with the evolution of CO gas and forming new Si-C bonds above 1300°C, enhancing the crystallisation process.

Alternative route was developed by US Sacks’ group (Toreki et al., 1992; Sacks et al., 1998): they used high-molecular-weight PCS. The infusible PCS was prepared by pressure pyrolysis of polydimethylsilane and the fibres formed by the dry spinning of concentrated PCS-based polymer solutions which were then pyrolysed in an inert atmosphere at 1000 to 1200°C.

The room temperature ultimate tensile strength ranges between 2.5 and 3.5 GPa as a function of the fibre grade with Young’s modulus ranging between ~200 and 400 GPa, very close to the best properties achieved for bulk ceramics, films or crystals: ~430 GPa (Biswas et al., 2001; Dirras et al., 2004; Chung & Han, 2008). The axial room temperature thermal conductivity ranges between ~2-3 (oxygen rich, amorphous SiC) up to 50-60 W/mK for crystalline nearly stoichiometric SiC (Simon & Bunsell, 1984). The electrical conductivity from ~0.01 to 10 S.cm\(^{-1}\) is highly related to the carbon content (Chauvet et al., 1992 a & b; Mouchon & Colomban, 1996). The reduction of the oxygen content in the fibres made by irradiating PCS produced fibres ~35% stiffer than the first generation fibres. The 1400°C creep rates are slightly improved (Sha et al., 2004). Actually the thermal stability in air remains poor for the first and 2\(^{\text{nd}}\) generation SiC fibres, ca. ~1100°C. The third generation stoichiometric fibres exhibit properties much closer to those of bulk SiC but the thermal stability in air remains much lower than that of oxide fibers (Mouchon & Colomban, 1995; Colomban, 1997; Baxter et al., 2000, Ruggles-Wrenn & Kutsal, 2010; Ruggles-Wrenn & Whiting, 2011).

3. Polytypes

Polytypes are observed in a variety of covalent compounds where rotation along the covalent bond does not require a lot of energy. This often gives rise to more or less lamellar compounds with stacking faults along one direction: carbides (Feldmann et al., 1968; Salvador & Sherman, 1991; Choyke & Pensl, 1997), sulfides (CdS, ZnS, TiS\(_2\), ... (Kaflawi et al., 1969; Schneide & Kirby, 1972; Tronc et al., 1975; Moret & Huber, 1976; Lincot et al., 1997; Agrosi et al., 2009; Alvarez-Garcia et al., 2009; Chi et al., 2011), nitrides (Komatsu et al., 2010) but also some oxides (e.g. BaTiO\(_3\) perovskites (Wu et al., 2006; idem, 2009)) and even diamond (Fayette et al., 1995; Bhargava et al., 1995).
As sketched in Figure 1, SiC structures consist of alternate layers of Si and C atoms forming a bi-layer. These bi-layers are stacked together to form face-centre cubic unit-cell (cubic stacking = ABC-ABC-ABC-, the so-called zinc-blende type cell, to be abbreviated c-SiC) or closed-packed hexagonal system (hexagonal stacking = AB-AB-AB-, the so-called wurzite cell, to be abbreviated h-SiC). Two consecutive layers form a bilayer which is named “h” (h for hexagonal) if it is deduced from the one below by a simple translation. If not, when an additional 180° rotation (around the Si-C bond linking the bilayers) is necessary to get the superposition, the bilayer is named “k” (for “kubic”). The “k” stacking is the reference of β-SiC cubic symmetry, only. The infinite combination of h/c stacking sequences led to hundreds of different polytypes (Feldman et al., 1968; Choyke & Pensl, 1997). Very similar structures are known for many compounds. Formation of polytypes arises because the energy required to change from one type to the other is very low. Consequently, different structures can be formed during the synthesis, simultaneously, especially for layer materials (CdS, SiC, TiS₂, MoS₂, BN, AlN, talc, micas, illites, perovskites, see references above) including MBE superlattices (Yano et al., 1995). Polytypes structure consists of close packed planes stacked in a sequence which corresponds neither to the face-centered cubic system nor the close-packed hexagonal system but to complex sequences associating both cubic and hexagonal stackings, ones such as = -ABABCABAB-, or −ABCAABABA-, or -ABABCABBA-, etc.).

Fig. 1. Schematic diagrams of the (a) hexagonal, (b) cubic, (c,d) polytypes modifications and of the stacking fault disorder (e). SiC structures alternate layers of Si and C atoms to form a SiC bi-layer, AB or AC (e).

4. From amorphous to crystalline materials

The precursor route led to a rather progressive transformation of a more or less 1D organised framework to a 3D amorphous one and subsequent thermal treatments control
the crystallization. The first problem to solve (Table 1) was the way to establish the bridge between the polymeric (Si-C)\textsubscript{n} chains: i) the first route (NLM\textsuperscript{TM} Nippon Carbon fibre (Ishikawa, 1995)) is the thermal oxidation (Si-O-Si bridge) at relatively low temperature (~200°C), the resulting SiO\textsubscript{2} content decreases from ~25 to ~10 wt% with improvements), ii) the second one is the electronic irradiation that allows forming Si-C bridges but leads to a carbon excess (C/Si ~1.4 in Hi-Nicalon\textsuperscript{TM} Nippon Carbon fibre (Berger et al; 1995; idem, 1999); alternatively the grafting of Ti or Zr alkoxide (Ti or Zr addition) leads to rather similar material but the fibres could be made with smaller diameter (UBE Industries Tyranno\textsuperscript{TM} LOX-M, ZE and TE grade fibres (Berger et al., 1997; idem, 1999); iii) the optimization of the organic precursor and associated thermal treatments gives stoichiometric SiC fibre (SA3\textsuperscript{TM} Ube Industries, Sylramic\textsuperscript{TM} Dow Corning Corp. Fibres and Hi-Nicalon\textsuperscript{TM} Type S (Lipowitz et al., 1995; Ishikawa et al., 1998; Berger et al., 1999; Bunsell & Piant, 2006). The high temperature of the manufacture process leads to much larger grain sizes.

| Generation | 1\textsuperscript{st} | 2\textsuperscript{nd} | 3\textsuperscript{rd} |
|------------|-----------------|-----------------|-----------------|
| Producer   | Nippon Carbon   | Nippon Carbon   | Ube Industries  |
| Grade      | NLM Nicalon     | Hi-Nicalon      | ZE,TE SA3       |
| Reticulation | Si-O bond     | Electron irradiation | Electron irradiation |
| Grain size/ nm | ~<2           | 5-10            | 5-10            |
| Si/C stoichiometry | 1.3           | <1.3            | <1.3            | ~1             | ~1             | ~1             |
| Diameter/ µm (+/- 3) | 15            | 12              | 11              | 7.5            | 10             | 12             |

Table 1. Small diameter SiC fibre generations.

The first generations fibre microstructures consist of an amorphous ternary phase made of SiO\textsubscript{x}C\textsubscript{y} tetrahedra (Porte & Sartre, 1989) with \(x+y = 4\), with ~1.4-1.7 nm SiC crystallites and ~5% of randomly oriented free carbon aggregates, 1 nm in size (Nicalon\textsuperscript{TM} 200 grade, \(x=1.15\)). Carbon (002) lattice fringe images showed small stacks of two fringes of around 0.7 nm in size suggesting that the basic structural unit (BSU) was a face-to-face association of aromatic rings, called dicoronenes, in which the hydrogene-to-carbon atomic ratio is 0.5. Accordingly, a porosity level of 2\% was present (Le Coustumer et al., 1995 a & b). Other studies proposed that the intergranular phase should be written as SiO\textsubscript{x}C\textsubscript{1-x/2}, which suggests that the composition varies continuously from SiC to SiO\textsubscript{2} as the oxygen traces varied (Bodet et al., 1995). The removal of oxygen from the cross-linking process resulted in a stoichiometry closer to Si/C = 1 and an increase in size of the β-SiC grains which were in the range of 5 to 10 nm in commercial fibres. The TEM images show well ordered SiC
surrounded by highly disorderd/amorphous SiC interphase and free carbon grains (Monthioux et al., 1990; idem, 1991; Havel, 2004; Havel et al., 2007).

5. How to identify the polytypes, the stacking disorder and the relative proportion of each polytypes?

The challenge for the nanotechnologies, which is to achieve perfect control on nanoscale related properties, requires correlating the production conditions to the resulting nanostructure.

Transmission electron microscopy (darkfield and high resolution images, electronic diffraction, etc. (see e.g. Mirguet et al., 2009; Sciau et al., 2009)) is the most efficient technique to determine the grain size, the defaults (disorder, superstructures, amorphous interface, voids, etc.) but the technique is destructive, time-consuming and may modify the sample structure. Moreover the representativity of the samples is always poor.

Raman spectrometry is a very interesting technique to study nanomaterials since it investigates the matter at a sub-nanometer scale, i.e. the scale of the chemical bonds. The automatic mapping (best spatial resolution ~0.5 to 1 µm² as a function of objective aperture and laser wavelength) allows a very representative view of the sample surface. Each Raman peak corresponds to a specific vibration (bending, stretching, librational, rotational and lattice modes) of a given chemical bond, and provides information (even on heterogeneous materials, e.g. composites) such as the phase nature and symmetry, distribution, residual stress,... (Colomban, 2002; Gouadec & Colomban, 2007). Since the Raman scattering efficiency depends on the polarisability of the electronic cloud, it can be very sensitive to light elements involved in covalent bonds (C, H, N, B, O, ...), which is a valuable advantage, when compared to X-ray/electron-based techniques (EDS, micro-probe,...). In the case of coloured materials if the exciting laser energy is close to that of absorbing electronic levels, resonance Raman scattering occurs and the technique becomes a surface analysis in the range of ~20 to 100 nm in-depth penetration (also depending on the wavelength, (Gouadec & Colomban, 2007)). Then, the selection of a given wavelength allows probing specific layers. The main advantages compared to infrared spectrometry are that the laser in a Raman equipment can be focused down to ~0.5-1 µm², allowing for imaging specific areas (Gouadec et al, 2001; Colomban, 2003; idem, 2005) and that Raman peaks are narrower that IR bands (Gouadec & Colomban, 2007 and references herein).

Fig. 2a shows the representative electronic diffraction pattern ([2-1-10] axis) of a SA3™ fibre thermally treated at 1600°C in inert atmosphere. Most of the Bragg spots correspond to 6H SiC (hexagonal P6mmc space group), i.e. to the most simple polytype (Fig. 1). The diffuse scattering along the horizontal axe ([01-11], arises from the stacking disorder of the SiC bilayer units. On the contrary, the disorder signature is weaker on the X-ray diffraction pattern (small polytype peak at d = 0.266 pm, Fig. 2b). However Bragg diffraction highlights the most crystalline part and sweeps the information on low crystalline (e.g. carbon) second phases. Fig. 3 shows the corresponding Raman spectra. For 1st and even 2nd generation fibres the Raman spectrum is dominated by the carbon doublet that overlaps the SiC Raman fingerprint. Specific thermal and chemical treatments are necessary to eliminate most of the carbon second phases and thus to have access to the Raman signal of the SiC phases (Havel & Colomban, 2005).
Fig. 2. a) Representative electron diffraction pattern recorded on SA3\textsuperscript{TM} (Ube Industries Ltd, see Table 1) fibre thermally treated at 1600°C under inert atmosphere (Courtesy, L. Mazerolles); b) X-ray diffraction pattern recorded on powdered SA3\textsuperscript{TM} fibre (the immersion in molten NaNO\textsubscript{3} do not modify the pattern, (Havel & Colomban, 2005)).

Fig. 3. Representative spectra of the as-produced fibres (a) and after different thermal/chemical treatments in order to highlight the SiC fingerprint (b).
Fig. 4. Variations of a) the ~1320 cm\(^{-1}\) Raman peak area (A\(_{1320}\)) and b) its wavenumber shift across the diameter of a NLM\(^{TM}\) fibre polished section, as-received (dot) and after a chemical attack (triangle) eliminating the carbon phase; a comparison of the variation of the “carbon rate” (Raman peaks surfaces ratio A\(_{1598}\)/A\(_{795}\)(C/SiC)) along the diameter of SA3\(^{TM}\) (c) and Sylramic\(^{TM}\) fibres section (d) (\(\lambda = 632\) nm, P= 0.5 mW, t= 60s).

Raman peaks attribution of the disordered carbons present in SiC fibres has been previously discussed (Karlin & Colomban, 1997; idem, 1998; Gouadec et al., 1998). Pure diamond (sp\(^3\) C-C bonds) and graphite (in plane sp\(^2\) C=C bond) have sharp stretching mode peaks at 1331 and 1581 cm\(^{-1}\) respectively. The two main bands of amorphous carbons are then assigned to diamond-like (D band for diamond and disorder) and graphite-like (G band for graphite) entities. Because diamond Raman scattering cross-section is much lower than that of graphite (~10\(^{-2}\)), a weak C\(_{sp3}\)-C\(_{sp3}\) stretching mode is expected. Actually, given the small size of carbon moieties and the strong light absorption of black carbons the contribution of the chemical bonds located near their surface will be enlarged (resonance Raman, the Raman wavenumbers shift with used laser wavelength, see in (Gouadec & Colomban, 2007)). The D band corresponds to vibration modes involving C\(_{sp3}\)-C\(_{sp2}\)/sp\(^3\) bonds also called sp\(^{2,3}\). This band presents a strong resonant character, evidenced by a high dependence of the intensity and position on wavelength. Additional components below 1300 cm\(^{-1}\) arise from hydrogenated carbons and those intermediate between D and G bands have been assigned to oxidised and special carbon phases (Karlin & Colomban, 1997; idem, 1998; Colomban et al., 2002). The wavenumber of the sp\(^3\) carbon bond (D peak) measures the aromaticity degree (aromaticity is a function of the “strength and extension size” of the \(\pi\) electronic clouds and thus also function of the crystal order) and hence is directly related to the electric properties of the material (Mouchon & Colomban, 1996). This value depends directly on the thermal treatment temperature history and hence is also related to the mechanical properties, see details in (Gouadec & Colomban, 2001; Colomban, 2003).

The plot of the carbon fingerprint parameters recorded across the fibre section diameter (on fracture) shows the very anisotropic carbon distribution (Fig. 4). Chemical treatments eliminate the carbon in the analysed SiC volume and hence allow a better study of the SiC phases (Havel & Colomban, 2005).
The Raman spectrum of well crystallised SiC phases is observed between 600 and 1000 cm\(^{-1}\) (Feldman et al., 1968; Nakashima et al., 1986; idem, 1987; idem, 2000; Nakashima & Hangyo, 1991; Nakashima & Harima, 1997; Okimura et al., 1987; Tomita et al., 2000; Hundhausen et al., 2008). The main Raman peaks centred at 795 and 966 cm\(^{-1}\) correspond to the transverse (TO) and longitudinal (LO) optic modes respectively of the (polar) cubic 3C phase, also called \(\beta\) SiC. Any other definite stacking sequence is called \(\alpha\)-SiC and displays either hexagonal or rhombohedral lattice symmetry. Polytypes in the \(\alpha\)-SiC structure induce the formation of satellite peaks around 766 cm\(^{-1}\) and of additional features between the TO and LO modes (Figs 5 & 6). However, the TO mode is twice degenerated; while TO\(_1\) is centred at 796 cm\(^{-1}\), TO\(_2\) is a function of the “h” layers concentration in the structure. A linear variation of 0.296 cm\(^{-1}\)/% has been demonstrated (Salvador & Sherman, 1991; Feldman et al., 1968).

Fig. 5. Representative Raman spectra recorded for NLM™ Nicalon fibres thermally and chemically treated (a,b). Detail on the disorder-activated acoustic modes observed for ZE™ fibre (c) and for very amorphous SiC zone are shown.

The main effect of the disorder is the break of the symmetry rules that excludes the Raman activity of the vibrational, optical and acoustical, modes (phonons) of the whole Brillouin
zone: only zone centre modes give rise to a Raman activity. Because the wavenumber of these modes shift with wavevector value, they give broad asymmetric bands. Fig. 6 illustrates the apparition of satellite peaks because the step-by-step Brillouin Zone folding associated to the formation of polytypes. On the contrary, stacking disorder lead to a projection of the vibrational density of state on the vertical energy axis and broad asymmetric bands are observed.

Fig. 6. a) Sketch of the folding of the original phonon Brillouin zone in the stretching LO/TO mode region along the stacking axis of the reference cubic symmetry by factor 2 (2H polytype), 4 (4H) and 6 (6H). b) Satellite peak wavenumbers for series of polytypes (after Nakashima & Harima, 1997).

The comparison of the Figures 2a (Diffraction & diffuse scattering) and 2b (Raman scattering) points out the very different sensitivity of these two methods. Fig. 4 compares the Raman spectra of the different generation SiC fibres, with carbon excess ranging from ~20 wt% (1st generation) to less than 1 wt% (3rd generation). A small wavenumber shift may be associated to the change of the exciting wavelength. Another important point is that for coloured materials, the interaction between laser light and matter must be very strong and hence the light absorption. This may have detrimental effect (local heating – and thermal induced wavenumber shift – (Colomban, 2002), oxidation and phase transition (Gouadec et al., 2001) in the lack of attention but this also controls the penetration depth of the laser light: the penetration can be limited to a few (tenths of) nanometers (Gouadec & Colomban, 2007).

Figs 5 to 9 give examples of the variety of Raman signatures observed on SiC materials issued of the organic precursor routes.

The narrow peaks pattern of crystalline polytypes is obvious and assignments are univocal with the comprehensive work of Nakashima (Nakashima et al., 1986; idem, 1987; Nakashima & Harima, 1997), see Fig. 6. The most stringent new features are the very broad bands observed at ~730 and 870 cm⁻¹ and the structured pattern below 600 cm⁻¹. The first feature corresponds to the amorphous silicon carbide and the second one to the acoustic modes rendered active because of the very poor crystallinity of the fibre.
Fig. 7. a) Raman spectra recorded every 2µm along a line from the centre of a SCS-6 Textron™ fibre (L= 532nm, 1mW, 120s/spectrum); b) representative spectra of the pure SiC (III) zone; the different components have been fitted with Gaussian or Lorentzian lines: the broad 740 and 894 cm\(^{-1}\) bands correspond to amorphous SiC, the 767 cm\(^{-1}\) to 6H-SiC and the 795 cm\(^{-1}\) band to 3C-SiC polytypes.

The apparition of disordered activated acoustic phonon in the Raman spectrum is not surprising in compounds with large stacking disorder (Chi et al., 2011). Additional multiphonon features are not excluded. However, many Raman studies of such materials have been made using exciting laser line leading to a resonance spectrum, simpler, in which the contribution of the disordered activated modes is low or even not detected.

Very similar features are observed for SiC materials prepared by Chemical Vapour Infiltration. The Raman spectra of the SiC coating deposited on a small diameter (~7µm) carbon fibre core to obtain the SCS-6 Textron™ fibre, a ~120 µm thick fibre used to reinforce metal matrix consist in features where the acoustic phonon intensity becomes stronger than the optical ones. Furthermore the latter group is dominated by the broad bands of the amorphous SiC.

Because of the different laser line absorption, Rayleigh confocal imaging allows to have very interesting image of the heterogeneous material (Colomban & Havel, 2002; Colomban, 2003; Havel & Colomban, 2003; idem, 2004; idem, 2005; idem, 2006). Fig. 8 shows representative spectra recorded on the deposit obtained around the fibres of a textile perform. In order to
optimise the thermomechanical properties of the composite a first coating of the SiC fibre with BN has been made. The spectra show the 3C (narrow peak at 799 and 968 cm\(^{-1}\)), 6H (786 cm\(^{-1}\)), 8H or 15R (768 cm\(^{-1}\)) as well the broad and strong contribution of amorphous SiC (optical modes at 750 & 900 cm\(^{-1}\) and acoustic modes at 450 cm\(^{-1}\) with shoulder at 380 and 530 cm\(^{-1}\)). Traces of carbon (1350-1595 cm\(^{-1}\) doublet) are also observed. We assign the broad Gaussian peaks at \(~ 700 \) cm\(^{-1}\) and \(~ 882 \) cm\(^{-1}\) to the amorphous SiC. Indeed, the position of the band at ca 882 cm\(^{-1}\) is exactly between the two optical modes at a wavenumber of \((796+969) / 2 = 882.5 \) cm\(^{-1}\). Dkaki et al. (Dkaki et al., 2001) already assigned the band at ca. 740 cm\(^{-1}\) to the amorphous SiC phase.

![SiC fibre reinforced-SiC matrix composite](image)

Fig. 8. Optical photomicrograph (a) and Rayleigh image (b) of a SiC (BN coated) fibre reinforced-SiC matrix composite. Examples of SiC spectra are given in c). Polytypes are evidenced by 786 (4H) and 768 (6H) cm\(^{-1}\) TO modes. The fingerprints of 3C (799 cm\(^{-1}\)) and amorphous (900 cm\(^{-1}\) broad band) SiC are also present.

When classically used, a Raman spectrometer is built to avoid the elastic (Rayleigh) scattering which is much more intense (\( \times 10^6 \)) than the inelastic one (Raman) and masks it. However, the Rayleigh signal contains useful information (volume of interaction and dielectric constant) that can be recorded in only few seconds, giving rise to topological and/or chemical maps (a high resolution Raman image requires tenths of hours!). The combination of Rayleigh image and Raman scattering is very interesting to study indentation figures (Colomban & Havel, 2002). Rayleigh scattering gives image of the topology mixed with information on the chemical composition through the variation of the optical index. Fig. 9 presents the Rayleigh image of the Vickers indented zone of the mixed SiC+C region (zone II) of a SCS-6 polished section (see Fig. 7). The automatic XY mapping has been performed with an objective with an Z axis extension of the focus volume sufficiently large to be bigger than the indentation depth. Thus, a 3D view is obtained.
up-deformation of the fibre matter close to the edges resulting from the pyramidal shape of the Vickers indenter is obvious. The residual stress is calculated using the experimental relationship previously established under pressure (Salvador & Sherman, 1991; Olego et al., 1982). The amorphization is obvious at the center of the indented area with the relative increase of the intensity of the 760-923 cm\(^{-1}\) doublet and the decrease of the TO/LO doublet; note, the up-shift of the TO mode from 796 to 807 cm\(^{-1}\). Similar information can be extracted from the D carbon band using the relationship established by Gouadec & Colomban, 2001.

| Peak  | Out of the indented area | At the tip position |
|-------|-------------------------|---------------------|
|       | \(\bar{\nu}\) (cm\(^{-1}\)) | P (GPa)            | \(\bar{\nu}\) (cm\(^{-1}\)) | P (GPa) |
| TO    | 796 ± 2                 | 0                  | 807 ± 6                   | 3 ± 2   |
| LO    | 969 ± 2                 | 0                  | 969 ± 4                   | 3 ± 2   |
| D     | 1351 ± 3                | 0                  | 1369 ± 4                  | 3 ± 1   |

Table 2. Comparison between the TO/LO peak wavenumbers measured at the tip and out of the 50 g Vickers indented area on SCS-6 Textron\(\text{TM}\) fibre, mixed SiC-C zone II (see Fig. 7a).

Fig. 9. (a,b) Rayleigh images of the Vickers indented area on the mixed SiC+C II region of a SCS-6 Textron\(\text{TM}\) fibre (100x100 spectra, 3s/Spectrum, 10\(^{-6}\) mW, \(\lambda = 532\) nm); (c,c’) representative spectra (step: 0.1\(\mu\)m) recorded at the core (c’) and the periphery (c) of the indented area; the fitting of the different component allows calculating the residual hydrostatic pressure (see Table 2).
6. Microstructure and defects

Fig. 10 shows representative high resolution Transmission Electron Microscopy (TEM) images recorded on thermally treated NLM 202 Nicalon™ and SA3™ fibres (Table 1). Structural studies of SiC nanocrystals were carried out on fragments of fibres deposited on a copper grid after crushing in an agate mortar (Havel, 2004; Havel et al., 2007). In SA3™ fibre the carbon phase appears to be well organized, graphitic, according to the narrow doublet of the Raman spectra (Fig. 3a). The interplane spacing is 0.33 nm. The stacking sequence of SiC bilayers is clear in Fig. 10c & d, because the contrast jump relative to the Bragg peak shifts. The domain sizes along the stacking direction might rich 3-4 nm. Figure 10c shows a typical HRTEM image of a nanocrystal with a size of 15 nm along its longest axis consisting of two regions corresponding to the α and β phases. The stacking faults, which are clearly seen on the micrograph, show no periodicity along the c axis of the hexagonal structure. Stacking faults can be considered as a perturbation of the β-SiC 3C stacking sequence so that the α phase can be seen as a sequence of β-SiC domains of various sizes ranging from 0.2 to 5 nm. The progressive transition between crystalline domains explains the variety of Raman fingerprint. There is a good agreement between Raman and TEM data.

7. Quantitative extraction of the (micro)structural information present in the Raman spectrum

For the decomposition of the SiC Raman peaks we used the spatial correlation model (SCM), which was established by Richter et al. (Richter et al., 1981), and by Nemanich et al.
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(Nemanich et al., 1981) and then popularised Parayantal and Pollack (Parayantal & Pollack, 1984). A comprehensive description for non-specialist has been given in our previous work (Gouadec & Colomban, 2007). It can be briefly explained as follows. In "large" crystals, phonons propagate "to infinity" and because of the momentum selection rule the first order Raman spectrum only consists of "q=0" phonon modes, i.e. the centre of the Brillouin Zone (Fig. 6). However, since crystalline perfection is destroyed by impurities or lattice disorder, including at the surface where atoms environment is singular, the phonon function of polycrystals is spatially confined. This results in an exploration of the wavevectors space and subsequent wavenumber shifts and band broadening. Another effect is the possible activation of "symmetry forbidden" modes. This is linked to the Brillouin zone folding as illustrated in Fig. 6. In the 6H polytype structure, the zone is folded three times at the Γ centre point and the reduced wave vectors that can be observed are at q = 0, 0.33, 0.67 and 1 (Feldman et al., 1968; Nakashima et al., 1987; Nakashima & Harima, 1997). The Raman line broadening can be described by the (linear) dependence of its half width upon the inverse grain size, as reported previously for many nanocrystalline materials including CeO$_2$ (Kosacki et al., 2002), BN (Nemanich et al., 1981), Si (Richter et al., 1981), etc.

In equation (1), the SCM describes the crystalline quality by introducing a parameter L$_0$, the coherence length, which is the average extension of the material homogeneity region. Noting q the wave vector expressed in units of π/a (a being the lattice unit-cell parameter) and Γ$_0$ the half width of Raman peaks for the ordered reference structure, the intensity I(ν) at the wavenumber ν is then given by equation (2). (Richter et al., 1981; Nemanich et al., 1981; Gouadec & Colomban, 2007).

The exponential function represents a Gaussian spatial correlation and ν(q) is the mode dispersion function, which can be deduced from neutron scattering measurements or from calculations often based on a rigid-model structure (Parayantal & Pollak, 1984; Weber et al., 1993; Kosacki et al., 2002).

\[ I(\nu) = I_0 \times \int_{q=0}^{q=1} e^{-\frac{k_B T \pi (q-\nu_0)^2}{16 \pi^2 \nu^2}} \times \frac{dq}{[\nu - \nu(q)]^2 + \left(\frac{\Gamma_0}{2}\right)^2} \]  (1)

While the one dimensional disorder (in the stacking direction) leads to the polytypes formation, a complete disorder induces the total folding of the Brillouin zone and the apparition of a very broad Raman signal (density of state spectrum, e.g. Fig. 9c). The phonon confinement is observed for small grains in a well crystallized state. The dispersion curve can be modelled with the Eq. 2-4 (Parayantal & Pollak, 1984). Our 6H reference corresponds to coefficients A and B of respectively $3.18 \times 10^5$ and $1.38 \times 10^{10}$ for TO and $4.72 \times 10^5$ and $8.52 \times 10^{10}$ for LO modes (Havel & Colomban, 2004).

\[ \nu(q) = \sqrt{A + \sqrt{A^2 - 8 \pi (1 - \cos(\pi q))}} \quad 0 \leq q \leq 1 \]  (2)

with

\[ A = \frac{1}{2} \times \nu_0^2 \]  (3)
and

\[ B = \frac{1}{2} \nu (\nu - 2) \left( \nu (\nu - 2) \right) \]

The SCM has been used to determine the size and structure of SiC nanocrystals extracted from annealed SiC fibre. The Raman spectra of the NLM fibres annealed 1h and 10h are shown in Fig 5. The SiC Raman signature, is composed of the 2 optical TO and LO modes. A satellite at 768 cm\(^{-1}\) indicates the presence of the 6H-SiC polytype (Fig. 6). The most interesting parameter in this SiC signature is the strong asymmetry of the LO peak at ~ 969 cm\(^{-1}\) (see also Fig. 7). The TO peak is much less asymmetric and centred at 796 cm\(^{-1}\). The elementary peaks obtained from the decomposition of the experimental spectrum are shown in Fig. 5 and the adjustment parameters (position, \(q_0\) and \(L_0\)) are summarized in Table 3.

| Peak | Parameter | 1h       | 10h      | 1h + corroded |
|-----|-----------|----------|----------|--------------|
| TO  | \(\nu\) (cm\(^{-1}\)) | 796.8    | 795.6    | 794.2 ± 2    |
|     | \(q_0\)   | 0.22 ± 0.02 | 0.26 ± 0.01 | 0.30 ± 0.01 |
|     | \(L_0\) (nm) | 5.6 ± 1.5 | 6.5 ± 1.2 | 5.6 ± 0.6 |
| LO  | \(\nu\) (cm\(^{-1}\)) | 961.7    | 969.5    | 954.5        |
|     | \(q_0\)   | 0.18 ± 0.03 | 0.00 ± 0.07 | 0.26 ± 0.01 |
|     | \(L_0\) (nm) | 2.9 ± 0.4 | 3.8 ± 0.9 | 5.2 ± 0.5 |

Table 3. Peak fitting parameters of the TO and LO peaks of SiC calculated from the Raman spectra of the NLM fibres annealed 1h and 10h at 1600°C and annealed 1h then corroded 100h in NaNO\(_3\) (Havel & Colomban, 2005).

For the fibre annealed for 1h, the \(L_0\) parameters of both TO and LO peaks show a confinement dimension in the range of 2.5 to 7 nm, in good agreement with the TEM image. After 10h annealing, the TO and LO peaks become sharper and more intense, indicating an increase in the size of the nanocrystals. This is confirmed by the \(L_0\) parameter, which gives a confinement dimension slightly higher, between 3 and 8 nm, according to the polytype domain size (Fig. 10).

8. Raman imaging

Raman imaging is very powerful, especially for heterogeneous materials but its rise is limited because of a lack of real control on the \(x, y, z\) spatial resolution (changing the diameter of confocal hole allows however some possibility) and of the huge recording time required (the spectrometer has often to be used during night time). However, a precise study of the laser shape, can improve the control on the resolution and since the CCD detectors are more and more sensitive, Raman images will now require more reasonable acquisition time (hours!). Note, that once the image is recorded, the set of spectra (also called hyperspectrum) has to be
analysed, which is much more time-consuming than the acquisition itself. This is why automatic decomposition software must be developed (Havel et al., 2004; Gouadec et al., 2011).

Because of their interesting thermal and mechanical properties, SiC composites (SiC fibres + SiC matrix) find numerous applications in the aerospace industry and new ones are expected in fusion ITER plant (Roubin et al., 2005). However, their expensive cost has to be balanced with a long lifetime, which is not yet achieved. To increase their lifetime, we first have to understand their behaviour under chemical and mechanical stresses, and thus, to characterize their nanostructure. In this section, we focus on the SiC fibres, which are analysed across their section. Indeed, this approach allows observing the chemical variations that may exist between the fibre’s core and surface. Fig. 11 shows Raman maps
of the Tyranno SA3™ (Ube Industry) fibre polished sections: a full spectrum is recorded
each 0.5 µm (the hyperspectrum) and after computation, Raman parameters are extracted
and mapped. Figures 11a & b consider the intensity variation of the TO SiC and D carbon
peaks (see also Fig. 4); this later line is assigned to the vibrations of peculiar carbon
moieties, which are thought to be located at the edges of the sp² carbon grains (Fig. 10a).
Fig. 11c (top) shows the wavenumber shift of the later D band. In this particular case, the
wavenumber shift represents the aromaticity of the carbon species. It has been reported
that this parameter also depends on the residual strain as is shown in equation 5
(Gouadec & Colomban, 2001).

\[ \Delta \nu_D = 10 \text{ cm}^{-1} / \text{GPa} \]  

(5)

The radial anisotropy results from the fibre preparation process: the fibre is heated from
outside and the departure of the H and C excess takes place at the fibre surface.
Consequently, because of the thermodynamic rules, the temperature of the fibre surface is
higher than that of the core, that keep C and H excess. After thermal treatment at 1600°C a
better homogeneity is achieved. Obviously, the specific microstructures of the different fibre
grades can be analysed using a “simpler” and faster diameter line-scan (Fig. 4).
The first maps (Figs 10a, b & c-top), representing the distribution of a simple Raman
parameter, may be of limited physical interest. However, it can be translated (through
models) to a property’ map (Colomban, 2003). The resulting image as exemplified in (Fig. 11c-
bottom) gives the distribution of physical parameters; we call it a “Smart image”.
For instance, the size of short-range ordered vibrational units in carbon moieties can be
deduced from the Raman parameters. It is based on the ratio of the intensity, I, of the two
main carbon Raman peaks (I_D/I_G), as first proposed by Tiunstra & Koenig, 1970.

\[ I_D / I_G = C_I / S_g \]  

(6)

with the grain size S_g in nm and the constant C = 44 for 5145.5 nm laser excitation; this
formula works well for relatively large grains (>2 nm). A new model (7) takes into account the
Raman efficiency, d, of the D^{1340} with respect to that of G^{1600}, as well as R, the ratio of atoms on
the surface of each grain with respect to the bulk, e_s the surface thickness and L_g the coherent
length (~ the grain size of Tuinstra and Koenig model). Assuming a spherical shape of all
grains the following equation can be proposed (Colomban et al., 2001).

\[ d \times R = d \times \left[ \left( 1 - \frac{2 \times e_s}{L_g} \right)^{-3} - 1 \right] \]  

(7)

This model has been used to calculate the carbon grains size distribution in SA3™ fibre’s
cross sections. We observe that the intensity ratio is much higher in the core than near the
surface and the carbon grain size appears approximately 2-3 times smaller on the fibre’s core
than on its periphery because the thermal gradient during the process.
The Raman data can be translated through equation 5 to a map of the maximum tolerable
strain (Colomban, 2003). The resulting image (Fig. 12) clearly evidences that the fibre’s
mechanical properties are better (~ 3.5 GPa) in the core than near the surface (~ 2 GPa).
Fig. 11. Raman map of calculated ultimate tensile strength of the SiC zones in a SA3\textsuperscript{TM} fibre section.

“Smart Raman images” in this section bring a lot of interesting information. First, there is a huge difference between the fibre’s core and surface with a radial gradient of physical properties as function of the fibre’ producer and additional treatments. Second, the maximum tolerable strain is observed in the fibre’s core, where the carbon species are the smallest (~ 1.5 nm). The core/skin differences are due to the elaboration process (spinning, sintering steps, etc.).

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We refer to the cited literature for further details.
Silicon Carbide (SiC) and its polytypes, used primarily for grinding and high temperature ceramics, have been a part of human civilization for a long time. The inherent ability of SiC devices to operate with higher efficiency and lower environmental footprint than silicon-based devices at high temperatures and under high voltages pushes SiC on the verge of becoming the material of choice for high power electronics and optoelectronics. What is more important, SiC is emerging to become a template for graphene fabrication, and a material for the next generation of sub-32nm semiconductor devices. It is thus increasingly clear that SiC electronic systems will dominate the new energy and transport technologies of the 21st century. In 21 chapters of the book, special emphasis has been placed on the aspects and developments thereof. To that end, about 70% of the book addresses the theory, crystal growth, defects, surface and interface properties, characterization, and processing issues pertaining to SiC. The remaining 30% of the book covers the electronic device aspects of this material. Overall, this book will be valuable as a reference for SiC researchers for a few years to come. This book prestigiously covers our current understanding of SiC as a semiconductor material in electronics. The primary target for the book includes students, researchers, material and chemical engineers, semiconductor manufacturers and professionals who are interested in silicon carbide and its continuing progression.

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