Composite Materials

Ceramic Matrix Composite Manufacturing Techniques and Applications Areas
Ceramic-fiber–ceramic-matrix composites provide improved strength and fracture toughness compared with conventional ceramics. Fiber reinforcements improve the toughness of the ceramic matrix in several ways.

First, a crack moving through the matrix encounters a fiber; if the bonding between the matrix and the fiber is poor, the crack is forced to propagate around the fiber in order to continue the fracture process.

In addition, poor bonding allows the fiber to begin to pull out of the matrix.

Both processes consume energy, thereby increasing fracture toughness.

Finally, as a crack in the matrix begins, unbroken fibers may bridge the crack, providing a compressive stress that helps keep the crack from opening.
Ceramic Matrix Composites

Two failure modes in ceramic-ceramic composites: (a) Extensive pull-out of SiC fibers in a glass matrix provides good composite toughness. (From Metals Handbook, American Society for Metals, Vol. 9, 9th Ed., 1985.) (b) Bridging of some fibers across a crack enhances the toughness of a ceramic-matrix composite. (From Journal of Metals, May 1991.)
Ceramic Matrix Composites

- The matrix is relatively hard and brittle

- Fiber reinforcement must have high tensile strength to arrest crack growth

- Fiber reinforcement must be free to pull out as a crack extends, so the reinforcement-matrix bond must be relatively weak
Ceramic Matrix Composites

Unlike polymer and metal matrix composites, poor bonding-rather than good bonding-is required. Consequently, control of the interface structure is crucial.

In a glass-ceramic reinforced with SiC fibers, an interface layer containing carbon and SiC is produced that makes debonding of the fiber from the matrix easy.

If, however, the composite is heated to a high temperature, the interface is oxidized; the oxide occupies a large volume, exerts a clamping force on the fiber, and prevents easy pull-out. Fracture toughness is then decreased.
Applications of CMCs

- Automotive industry.
- Heat exchangers
- Aerospace and military applications.
- Bearings in missiles.
- Other applications include wear parts, such as seals, nozzles, pads, liners, grinding wheels, brakes, etc. For instance, carbon fiber reinforced carbon composites are being used in aircraft brakes.
- They are also used in dies and tool bits, medical implants and land-based power and transport engines.
Applications of CMCs

Carbon-Ceramic Matrix Rotors

Applications of CMCs

The Porsche Carrera GT's carbon-ceramic (SiC) composite disc brake

Ceramic Matrix Composite Turbine Blade
CMCs are excellent candidates for replacing the nickel-based superalloys currently used in exhaust nozzle parts, primarily due to their capacity to withstand the high temperatures and severe operational environment for much longer periods of time with minimal changes in structural behavior.

In examining the feasibility of using the A500 seals on the divergent section of the exhaust nozzles, AFRL researchers are addressing a number of key Air Force issues--one of which involves the performance comparison of CMC parts in flight and during engine ground testing. SPS has developed a novel CMC that uses carbon fibers in a sequentially layered carbide matrix produced via chemical vapor infiltration. Because this resultant matrix is self-sealing, it helps protect the carbon fibers from oxidation. The fibers are woven in a multidimensional, ply-to-ply angle interlock pattern to reduce the chance of delamination.

Applications of CMCs

SiC/SiC for Nuclear Applications

- SiC/SiC composites are considered as advanced structural materials in fission and fusion energy systems
  - Heat resistance
  - Intrinsic safety features

- VHTR / NGNP
  - Control rods
  - Control rod supports
  - HX

- GFR
  - Core structures
  - Fuel pins
  - HX

- Fusion
  - Blanket structures
  - Blanket channel liners

Japanese new programs, INERI, GNEP, GIF, BA and many other programs include SiC/SiC applications in LWR, SFR, VHTR, GFR and Fusion
Applications of CMCs

Braided and unidirectional S-2 Glass and carbon fibers are used to produce forks with different stiffness.

http://matse101.mse.uiuc.edu/
Particulate-, platelet-, whisker- and short-fiber-reinforced ceramic-matrix composites can be fabricated via the usual ceramic processes used for multiphase ceramics.

Whisker and short fiber reinforced composites can be processed by hot pressing. For continuous-fiber reinforced composites the first step is building up a 3D architecture from the fibers by textile production techniques (weaving, stitching, knitting and braiding).

For fiber- and whisker-reinforced composites, the reaction between the reinforcement and the matrix must be minimized, i.e. The processing temperature should not be too high.
CMCs

PREFORM ARCHITECTURES

Biaxial Weave
Triaxial Weave
Knit
Multiaxial Multilayer Warp Knit
3-D Cylindrical Construction
3-D Braiding
3-D Orthogonal Fabric
Angle-Interlock Construction

Illustrations—Scientific American
Processing of CMCs

PARTICULATE COMPOSITES

Ceramic composites containing particulates are similar to multiphase ceramics, and these materials can be fabricated through traditional ceramic processes.

WHISKER COMPOSITES

The green density of the mixture of whiskers and matrix powders is generally low due to a high aspect ratio of whiskers. Pressure sintering, such as hot pressing, is therefore used for densification. Another problem is due to the agglomeration of whiskers, and careful mixing of whiskers with matrix forming powders is required to avoid inducing serious damage to the whiskers. A typical procedure for this is to disperse the whiskers in liquid followed with successive filtration to remove agglomerated whiskers. The whiskers are then prepared for composite fabrication by blending with powders of the matrix material.

Chapter 15.1, Ceramic-Matrix Composites, AKIRA OKADAS, Handbook of Advanced Ceramics, 2003, Elsevier Inc., Somiya et al. (Eds.),
The mechanical behavior of continuous fiber composites is very different than that of other brittle ceramics. In tensile loading, a change in the linear stress-strain relation occurs after matrix cracking, and sliding pullout contributes to the load bearing ability afterward.

Moreover, shear failure and compression failure are often observed in flexural tests, resulting from delamination due to shear stresses and fiber buckling due to compression.

Such a failure mode is obviously derived from the weak interface between the fiber and matrix. Therefore, the presence of lubricant carbon and boron nitride at the interfaces is preferred. Although excess carbon on the surface of SiC fibers acts as a lubricant, the carbon layer may oxidize in air. Boron nitride coating on fibers has been carried out to maintain weak interfaces at high temperatures.
Continuous fibers such as carbon fibers and SiC fibers can be used to reinforce a glass matrix.

The strengthening mechanism is similar to that in resin matrix composites, and the fibers carry most of the load due to their much higher Young's modulus compared to the matrix.

Of all the continuous fiber composites, glass matrix composites are particularly dense as they are produced by impregnation of a glass melt.

The mechanical properties are characterized by high strength and large fracture energy. The large fracture energy is explained in terms of intensive pull out of fibers from the matrix glass. This indicates that the fiber to matrix bond is poor due to the presence of lubricant carbon layers on the surfaces of both graphite and silicon carbide fibers. In contrast glass matrix composites using oxide fibers exhibit low flexural strength due to the high bonding strength between fiber and glass matrix.
Carbon fiber glass matrix composites were intensively studied in the 1970s and SiC fiber composites in the 1980s.

In both cases the fabrication method is essentially the same.

Fiber tows or fabrics are first immersed in a glass powder suspension, and the powder-containing fiber sheets are stacked ready to be hot-pressed into laminates.
The development of carbon/carbon composites began in 1958, and they have been applied to the hot parts of missiles and the Space Shuttle, such as nose caps and leading edges.

Carbon/carbon composites can withstand temperatures higher than 3000°C in a vacuum and in an inert atmosphere, without losing strength as the operating temperature is increased. However, they oxidize and sublime when in an oxygen atmosphere at 600°C.

Silicon carbide coatings are therefore coated with a layer of glass to protect them in high temperature applications. When the part is cooled down from the coating temperature, microcracks develop in the silicon carbide layer, resulting from thermal expansion mismatch between carbon and silicon carbide. These cracks might cause oxidation of the substance if exposed to the air, but are immediately impregnated with the overcoated glass.
Carbon/carbon composites have successfully replaced metallic brake discs in racing cars and aircraft because of their lightweight.

Civilian aircraft, such as the Concorde supersonic jet and Boeing 767 use carbon/carbon composite brakes.

In comparison to steel brakes, a 40% weight saving is achieved using carbon/carbon composites due to their large heat capacity (2.5 times that of steel) and high strength (twice that of steel) at elevated temperatures.

Carbon/carbon composites are produced by pyrolyzing an organic matrix or by CVI. CVI of carbon from hydrocarbon gas is normally accomplished at 1100°C, and pyrolytic carbon is obtained. Carbonized organic composites are typically produced from graphite fabrics pre-impregnated with phenolic resin. The fabrics are laid up as a laminate and cured. They are then pyrolyzed to form a matrix of glassy carbon around the graphite fibers. As a result of repeated impregnations of resins and pyrolyzations, densification is achieved.
Since the oxidation resistance of SiC is much better than that of carbon, SiC/SiC composites have been developed for aerospace application such as propulsion and high velocity systems.

Similar to carbon/carbon composites, the SiC/SiC continuous fiber composites consist of a fiber architecture made of silicon carbide fibers in a matrix of silicon carbide.

The matrix is usually produced by CVI or preceramic polymer impregnation and pyrolysis.
The major advantage is that oxide materials have no oxidation problems. Development of all-oxide composites has been a major goal of recent research. Such composites have an interface configuration which allows a crack to propagate along the interface after matrix cracking. There are several microstructural design strategies. The first is to use fugitive layers, the second is to use stable oxide interfaces with suitably low fracture toughness, and the third is to use a porous matrix because the porous interlayers act as crack deflection paths.
Recently, $\text{Al}_2\text{O}_3$-$Y_3\text{Al}_5\text{O}_{12}$ eutectic composites have been attracting considerable attention.

Firstly, the creep rates of $\text{Al}_2\text{O}_3$-$Y_3\text{Al}_5\text{O}_{12}$ eutectic composites are considerably lower due to the excellent creep resistance of $Y_3\text{Al}_5\text{O}_{12}$ single crystals, and meet the design guidelines for use in gas turbines.

Furthermore, these composites have greater fracture toughnesses than single crystals, and maintain their high flexural strengths up to 1700°C.
The choice of manufacturing method is determined by the

- reinforcement and component geometry,
- the complexity of shape and
- the production volume of the component.
To put a rigidized or densified matrix in place, the precursor of the matrix has to be positioned within the mass of reinforcement. This can be done by a number of methods:

(i) Powder dispersion
(ii) Liquid precursors
(iii) Gaseous infiltration
Processing of CMCs

Powder Dispersion

Impregnating the reinforcement with a suspension of matrix precursor in powder form, either by passing the reinforcement through a slurry or by pressure impregnation of a preform, or by electrophoretic infiltration.

The powder dispersion method is the most widely used where a simple approach using substances of known composition or characteristics is required. Most types of matrix can be positioned using this method. Normally it requires that the fiber or whisker architecture is opened up so that powder particles can completely surround each reinforcing element, and that there is sufficient powder entrained for the densified matrix to fill the space around the reinforcement to an adequate degree, usually completely. Matrix precursors entrained in this way, even if sinter-active, tend to be reluctant to sinter to full density because of the restraint posed by the nonshrinking reinforcement structure, and hot-pressing, usually uniaxial, is required to close voids between reinforcing elements. This process does not apply to complex shapes. Generally plate shapes with planar reinforcement are easiest to produce because the reinforcement is not greatly distorted and thus it is not particularly versatile for producing components with complex architecture.
Processing of CMCs

Particulate reinforcements are compacted to as much as 75 vol% by filtration of solid loaded slurry and subsequent drying.
Processing of CMCs

Liquid precursors

Impregnating the reinforcement preform with a liquid organic or organometallic or inorganic substance, typically a polymer or a sol, which on heating rigidizes by curing or gelling, and then decomposes to leave a ceramic matrix.

A more reliable way of penetrating the reinforcement architecture is to use a liquid precursor. This can be in the form of an aqueous sol (e.g., boehmite, AIOOH, later to form alumina), or an organic or organometallic polymer which decomposes on heating to form a ceramic.

The disadvantage of this method is that it is not very efficient because the volume occupied by the resulting solid is much less than that of the impregnating liquid. Several stages of repeated impregnation and decomposition may be needed to obtain an impervious final product, which even so still contains interreinforcement closed porosity.

The advantage of the method is that complex shapes can be made based on the reinforcement preform shape, but the disadvantage is the cost in terms of the number of processing steps involved.
Preceramic polymer can fill a preform with liquid polymers, either molten or in solution, which are then pyrolized to make a ceramic matrix.

Polymer impregnation and thermal decomposition are repeated several times.

Resins, such as phenol and pitch, are used for producing carbon matrix composites, and organosilicon compounds, such as polycarbosilane and polyvinylsilane, are used for the impregnation of silicon carbide.

Sols have also been used for infiltrating preforms to produce oxide matrices, such as alumina, silica, zirconia and mullite.

Following infiltration, they are gelled by drying or by adjusting the temperature, and the matrix is formed after heat treatment.
Processing of CMCs

Liquid infiltration

It is similar to the resin transfer molding process which is used for the processing of polymer matrix composites and squeeze casting used for metal matrix composites. The three major issues to be considered in the liquid infiltration process are:

a) Chemical reactivity: The process is done at an elevated temperature and at high temperatures; the reinforcement and the matrix may react resulting in unnecessary reaction which deteriorates the bonding between the reinforcement phase and the matrix phase.

b) Infiltrant viscosity: The viscosity of ceramic suspensions are high which can result in the inability of the ceramic infiltrant to infiltrate into the fibrous ceramic preform.

c) Wettability of the reinforcement: The infiltrant may not be able to wet the reinforcement resulting in improper bonding which further leads to the failure of the product at the interface of reinforcement and the matrix.

Applications of Liquid infiltrated CMCs

Multiple-mission spacecrafts necessitate reusable heatshields and heatexposed elements. Ceramic Matrix Composite (CMC) are the most suitable structural heatdurable materials for these functions, carbon fibre reinforced silicon carbide (C/ SiC) in particular.

For producing these composites, the carbon fiber mats are imprignated with the Fast-Sol-Gel, a resin based on rapid hydrolysis and polymerization of a mixture of (Me)_xSi(O-Me)_{4-x} monomers. After a gradual heat-pressure process under inert atmosphere the green composites are converted into C-SiC composites. Schematic reaction: (SiRO_{3/2})_n \rightarrow SiC + CO_2 + H_2O

http://www.aac-research.at/products/products_AAC_Service_CeramicMatrix_en.html
Applications of Liquid infiltrated CMCs

In addition such materials require an Oxidation Protection System (OPS) to prevent oxidative damage to the carbon fibres during re-entry. These OPS should reliably protect the C/SiC structures at temperatures up to 1600°C and must remain crack-free over the whole temperature range from approx. 450 to 1600°C. This system consists of the Fast-Sol-Gel resin and ceramic filler like n-Al₂O₃ or n-ZrO₂.


SEM view of C/SiC composite with two layers of OPS.

http://www.aac-research.at/products/products_AAC_Service_CeramicMatrix_en.html
In the **liquid phase routes**, the fibres first coated with an interphase (e.g. by I-CVI) are embedded in a liquid precursor of the matrix.

In the **reactive melt infiltration (RMI) processes**, a fibre preform is impregnated by capillary forces with a liquid which reacts either with a solid phase used to consolidate the fibre preform (SiC-Si matrices formed through liquid silicon infiltration of a carbon-consolidated preform) or with the atmosphere (Al₂O₃-Al matrices formed through liquid aluminium infiltration and chemical reaction with an oxidizing atmosphere).

Among other **advantages**, the RMI-processes are fast and **can be applied to thick preforms**. They also yield materials of **low residual porosities** and **high thermal conductivities**.
Processing of CMCs

In the *polymer impregnation and pyrolysis (PIP) processes*, the fibres are embedded in a polymeric precursor of the matrix, such as a thermosetting resin or a pitch for carbon or a polycarbosilane for SiC, and the green composite is then pyrolyzed.

Such processes are *relatively flexible* since the composition of the precursor can be tailored.

Conversely, a shrinking of the matrix occurs during the pyrolysis step owing to the evolution of gaseous species. As a result, several PIP-sequences have to be applied in order to achieve a low enough residual porosity, which is time and labour consuming. Shrinkage can be limited by loading the liquid precursor with suitable fine powder, i.e. by using a slurry. Finally, the residual porosity can also significantly be reduced through a hot pressing step, an alternative that supposes that the matrix displays enough plasticity not to damage the fibres. This liquid impregnation/hot pressing technique is well suited to the fabrication of glass-ceramic matrix composites.
Carbon-carbon composites are made by forming a polyacrylonitrile or carbon fiber fabric into a mold, then impregnating the fabric with an organic resin, such as a phenolic.

The part is pyrolyzed to convert the phenolic resin to carbon. The composite, which is still soft and porous, is impregnated and pyrolyzed several more times, continually increasing the density, strength, and stiffness.

Finally, the part is coated with silicon carbide to protect the carbon-carbon composite from oxidation.

Carbon-carbon composites have been used as nose cones and leading edges of high-performance aerospace vehicles such as the space shuttle, and as brake discs on racing cars and commercial jet aircraft.
Carbon-carbon (C—C) composites are used for extraordinary temperature resistance in aerospace applications.

Carbon-carbon composites can operate at temperatures of up to 3000°C and, in fact, are stronger at high temperatures than at low temperatures.

A comparison of the specific strength of various carbon-carbon composites with that of other high-temperature materials relative to temperature.
Gaseous infiltration

Using a reactive gas mixture which deposits a ceramic material within a preform of the reinforcement, commonly known as chemical vapor deposition (CVD) or chemical vapor infiltration (CVI), and typically performed at high temperature; all of which have particular advantages or disadvantages for different types of matrix material.

For those matrix materials which can be produced by reaction between gases, such as carbon and silicon carbide, a gas phase route can substitute for the liquid impregnation route. This tends to be a slow process, because if deposition is allowed to occur too quickly, it mostly occurs at the external surface, blocking penetration.

Developments in the technique have concentrated on ensuring that deposition occurs internally to the component, by forcing reactive gas flow through the walls of the component (CVI), using temperature gradients, and on speeding the process up, e.g., by using microwave heating which also has the advantage of improving deposition internally to the preform. However, as with liquid infiltration, full densification of the matrix never occurs as access to remaining pores becomes blocked, but the method can be used for complex shapes.
In gas phase routes, i.e. the so-called chemical vapor infiltration (CVI) processes, the reinforcements (usually as a multidirectional preform) is densified by the matrix deposited from a gaseous precursor, e.g. an hydrocarbon for carbon or a mixture of methyltrichlorosilane and hydrogen for silicon carbide.

There are several versions of the CVI-process.

It is now well established that a fibre coating, referred to as the interphase, has to be deposited on the fibre prior to the infiltration of the matrix in order to control the fibre-matrix (FM) bonding and the mechanical behavior of the composite.

The main role of the interphase is to deflect the microcracks which form in the matrix under loading and hence to protect the fibre form notch effect (mechanical fuse function).
Chemical vapor impregnation (CVI) is a method of infiltrating fiber architectures with matrix particles via the vapor phase.

Although this is a similar process to chemical vapor deposition (CVD) in terms of gas reaction, deposition conditions in CVI are chosen for in-depth deposition rather than coating the surface of the substrate.

Several techniques have been developed to introduce the reaction gases into the fiber architecture, such as temperature gradient, pressure gradient and pulse CVI.
Processing of CMCs

CHEMICAL VAPOR IMPREGNATION

CVI equipment for the temperature gradient technique

Chapter 15.1, Ceramic-Matrix Composites, AKIRA OKADAS, Handbook of Advanced Ceramics, 2003, Elsevier Inc., Somiya et al. (Eds.),
When fabricating ceramic composites, the reaction between the reinforcement and matrix must be minimized. Consequently, a suitable low processing temperature should be selected to minimize degradation of the reinforcing phases.

Use of continuous fibers can modify the mechanical properties if suitable fiber orientations are chosen. A well-designed fiber architecture is first prepared, and the matrix material introduced into the voids of the structure. Infiltration into the fiber architecture is often performed using chemical vapor infiltration or pre-ceramic polymer infiltration-pyrolysis techniques. These techniques have the advantage of requiring low temperatures in comparison with sintering processes.

Textile production techniques, such as weaving, stitching, knitting and braiding are used for producing three-dimensional composite structures, and the stacked sheets of fabrics are used for manufacturing panel structures.

Hot pressing is used for fabricating glass matrix composites due to the relatively low processing temperatures, and this is also used for the fabrication of whisker composites.
Uniaxial and two-dimensional fiber composites with glass matrices can be produced by hot pressing.

The fibers are immersed in a slurry of matrix particles and then dried.

The fiber/matrix powder preform is then cut into suitable dimensions, and stacked for hot pressing in order to produce dense composites.

The fabricated composites are usually in the shape of flat disks or rectangular plates. Densification of whisker composites is usually performed by hot pressing.

Careful control is required both to avoid damage to the whiskers and to achieve a homogeneous dispersion of whiskers.
Processing of CMCs

HOT PRESSING

**Hot isostatic pressing**

- Pressure
- Pressurized gas (argon)
- Pressed part
- Steel can container
- Heated chamber

**Hot pressing**

- Die fill stage
  - Upper punch
- Pressed part
- Lower punch
- Hot die

**Part ejection**

www.substech.com
NOVEL TECHNIQUES

Melt infiltration into fibrous preforms combined with oxidation of the metal matrix can produce ceramic matrix composites.

Since this type of process was developed by Lanxide Corporation, it is called the Lanxide process.

For example, a Nicalon SiC fiber and alumina matrix composite was produced as follows: stacked fibrics of Nicalon fiber were coated by CVD. The major purposes of the coating were to protect the fiber from aluminum alloys during the matrix growth process and to provide a weak fiber-matrix interface. The fiber architectures were then placed on molten aluminum to allow the matrix to grow in the fiber preform by direct oxidation of the aluminum alloy.
Eutectic consolidation using a single crystal production technique can also be used in the production of two-phase ceramic composites.

Aligned composite structures can be produced by unidirectional solidification of binary eutectics. The BaFe$_{12}$O$_{19}$-BaFe$_2$O$_4$ eutectic has been, for example, produced from powder mixtures of Fe$_2$O$_3$ and BaCO$_3$. The melt was consolidated by moving the specimen gradually in the furnace, in a similar way to the Bridgman technique.

Using a similar technique, a variety of eutectic systems were investigated. The microstructure revealed that fibrous structures are developed along the longitudinal direction, and the flexural strength is greater when the crack propagates across the fibrous structure.
## Summary of Processing Routes

<table>
<thead>
<tr>
<th>Processing Route</th>
<th>Matrices</th>
</tr>
</thead>
<tbody>
<tr>
<td>Chemical vapour infiltration</td>
<td>Carbides, nitride, carbon, oxides, borides</td>
</tr>
<tr>
<td>Viscous phase hot pressing (2D preforms)</td>
<td>Glasses, ceramic-glasses</td>
</tr>
<tr>
<td>Sol-gel, slurry routes (2D, 3D preforms)</td>
<td>Oxides, carbides, nitrides</td>
</tr>
<tr>
<td>Polymer precursor route (3D preforms)</td>
<td>SiC, SiₙNₘ, SiₓCᵧNₗ</td>
</tr>
<tr>
<td>Liquid metal infiltration</td>
<td>Si → SiC</td>
</tr>
<tr>
<td>Gas-metal reaction</td>
<td>Oxide (Al, nitrides (Al, Zn, Ti))</td>
</tr>
<tr>
<td>Solid-state hot pressing</td>
<td>SiC, Si₃N₄</td>
</tr>
<tr>
<td>Prepreg curing and pyrolysis</td>
<td>SiC, Si₃N₄</td>
</tr>
<tr>
<td>Hot pressing (2D preforms)</td>
<td>Oxides</td>
</tr>
</tbody>
</table>
# Processing of CMCs

## Some processes for continuous fibre-reinforced CMCs

<table>
<thead>
<tr>
<th>Processing method</th>
<th>Advantages</th>
<th>Disadvantages</th>
<th>Fibre</th>
<th>Matrix</th>
<th>Temperature range* (°C)</th>
</tr>
</thead>
<tbody>
<tr>
<td>I. Slurry infiltration</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>(a) Glass ceramic matrix</td>
<td>• Commercially developed</td>
<td>• Limited max. Temperature due to matrix</td>
<td>• Graphite</td>
<td>• Glass-</td>
<td>• 800-1000</td>
</tr>
<tr>
<td></td>
<td>• Good mechanical properties</td>
<td>• Needs to be hot pressed, expensive</td>
<td>• Nicalon</td>
<td>ceramic</td>
<td>• 800-1000</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Formations of complex shapes is difficult</td>
<td></td>
<td></td>
<td></td>
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<tr>
<td>(b) Ceramic matrix</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1. Sintered matrix</td>
<td>• Potentially inexpensive</td>
<td>• Shrinkage during sintering cracks matrix</td>
<td>• Graphite</td>
<td>Alumina SiC</td>
<td>• 800-1400</td>
</tr>
<tr>
<td></td>
<td>• Could produce complex shapes</td>
<td>• Temperature limit due glassy phase</td>
<td>• Nicalon</td>
<td>Si₃N₄</td>
<td>• 800-1600</td>
</tr>
<tr>
<td>2. Cement bonded matrix</td>
<td>• Inexpensive</td>
<td>• Relatively poor properties to date</td>
<td>• Graphite</td>
<td>Cements</td>
<td>• 800-1500</td>
</tr>
<tr>
<td></td>
<td>• Ability to produce large complex shapes</td>
<td></td>
<td>• Nicalon</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>• Low temperature processing</td>
<td></td>
<td>• “New” fibres</td>
<td></td>
<td></td>
</tr>
<tr>
<td>3. Reaction bonded</td>
<td>• Good mechanical properties</td>
<td>• Has requires hotpressing of Si powder in Si₃N₄ system prior to reaction bonding</td>
<td>• Nicalon</td>
<td>•Si₃N₄</td>
<td>• 800-1500</td>
</tr>
<tr>
<td></td>
<td>• Pressureless densification</td>
<td>• Simple shapes only</td>
<td>• SiC</td>
<td></td>
<td>• 800-1600</td>
</tr>
</tbody>
</table>

*: Temperature limit depends on fibre. Currently all systems are limited to ~1200°C available fibres.

### Processing of CMCs

<table>
<thead>
<tr>
<th>Processing method</th>
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<th>Disadvantages</th>
<th>Fibre</th>
<th>Matrix</th>
<th>Temperature range* (°C)</th>
</tr>
</thead>
</table>
| **II. Sol-gel and polymer processing** | • Good matrix composition control  
• Easy to infiltrate fibres  
• Lower densification temperature | • Low yields  
• Very high shrinkage  
• Would require multiple infiltration/densification steps  
• No promising results reported | • Nicalon | • Non-oxide  
• Alumina  
• Silicates | • 800-1200  
• 800-1400 |
| **III. Melt infiltration** | | | | | |
| a) Ceramic melt | • Potentially inexpensive  
• Should be easy to infiltrate fibres  
• Lower shrinkage on solidification | • High melting temperatures would damage fibres | • Graphite  
• Nicalon  
• “New” fibres | • Glass | • 800-1100  
• 800-1100 |
| b) Metal melt, followed by oxidation | • Potentially inexpensive  
• Cermet type material | • Difficult to control chemistry and produce all ceramic system  
• Difficult to envision in use for large, complex parts for aerospace applications | • Graphite  
• Nicalon | • Alumina | • 800-1200 |

*: Temperature limit depends on fibre. Currently all systems are limited to ~1200°C available fibres.

### Processing of CMCs

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</tr>
</thead>
<tbody>
<tr>
<td><strong>IV. Chemical vapour infiltration</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>a) General approach</td>
<td>• Has been commercially developed • Best mechanical properties • Considerable flexibility in fibres and matrices • High quality matrix, very pure • Little fibre damage • In-situ fibre surface treatment • Ability to fill small pores</td>
<td>• Slow and expensive • Requires iterative process • Never achieved full density • Capital intensive</td>
<td>• “New” fibres • Nicalon</td>
<td>• B₄C • SiC</td>
<td>• 800-1200 • 800-1600</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>• Nextels</td>
<td>• SiC</td>
<td>• 800-1800</td>
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<td></td>
<td></td>
<td>-</td>
<td>• HfC</td>
<td>-</td>
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<td>b) Lanxide</td>
<td>• Ability to produce complex shapes • Properties dominated by ceramic • Very porous grain boundaries • Systems include: AlN/Al, TiN/Ti, ZrN/Zr</td>
<td>• Slow reaction and growth kinetics • Long processing time and high temperature limits chemistry • Wetting and reactions are limitations</td>
<td>• Graphite</td>
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*: Temperature limit depends on fibre. Currently all systems are limited to ~1200°C available fibres.  