A Solid State Approach to Synthesis Advanced Nanomaterials for Thermal Spray Applications

Behrooz Movahedi
Department of Nanotechnology Engineering,
Faculty of Advanced Sciences and Technologies, University of Isfahan,
Iran

1. Introduction

Preparation of feedstock powders is the first step for synthesis of thermal spray coatings. A number of techniques that are capable of producing these materials include gas/water atomization, mechanical alloying/milling, thermo-chemical method, spray drying, agglomeration and sintering, plasma fusion, and sol–gel processing techniques (Berndt, 1992). Solid state synthesis is attributed to the chemical reaction and alloying performed at temperature which reactants are solid. As a result, the kinetics of solid state reactions are limited by the rate at which reactant species are able to diffuse across phase boundaries and through intervening product layers. Hence, the conventional solid state technique invariably require the use of high processing temperatures to ensure that diffusion rate is maintained at a high level (Schmalzried, 1995; Stein et al., 1993). More recently, the mechanical milling process has attracted considerable interest, primarily as a result of its potential to generate nanocrystalline and other non-equilibrium structures in large quantities at low temperature. This process is considered as a means to mechanically induced solid state reaction that occur in feedstock powder mixture during collision in the grinding media (Suryanarayana, 2001).

The aim of this chapter is to describe the fundamental, mechanisms and recent developments of solid state approach to synthesis thermal spray advanced materials and related coatings.

2. Solid state synthesis of thermal spray powders

Mechanical milling as a solid state synthesis usually performed using ball milling equipments that generally divided to “low energy” and “high energy” category based on the value of induced the mechanical energy to the powder mixture. The ball milling equipments used for mechanical grinding or mixing are low energy such as Horizontal mill (Tumbler). The speed of the low energy rod or ball mill is quite critical with regards to the efficiency of the process (Fig. 1). It is necessary for the balls (or rods) to drop from the top of the mill onto the feedstock material that is being ground (Fig. 1b). If the mill speed is too fast then the media will not fall at all due to centrifugal forces or it will fall directly onto the media near the bottom of the mill (Fig. 1c). At low speeds the media does not drop at all, whereas at the optimum speed the media continuously "cascades" onto the feedstock material that is being crushed (Brendt, 1992).
Fig. 1. Low energy mechanical milling variables that control particle grinding and efficiency: (a) low speed, (b) optimum speed, and (c) high speed (Berndt, 1992).

In mechanical milling processes that utilize to change the chemical composition of precursors, the high energy ball milling equipment is generally used. This phenomenon can be performed in various types of high energy ball mills, including attrition, planetary, and vibratory mills that are schematically shown in Fig. 2. In an attrition mill, the rotating impeller causes relative movement between balls and powders. In a planetary ball mill, a rotating disc and vials revolve in opposite directions in order of several hundred rpm. In a vibratory mill that is also known as a shaker mill, the vessel is set in 1D or 3D vertical oscillatory motion. Spex 8000 is a commercial type of 3D vibratory mills. (Suryanarayana, 2001, 2004). Of the above types of mills, only the attritor mill has the highest capacity of powder charge. Accordingly, the attritor milling is employed to synthesize thermal spray feedstock powders used for the fabrication of nanostructured coatings.

Fig. 2. Various types of high energy ball mills: (a) attrition mill, (b) vibratory mill, and (c) planetary mill (Suryanarayana, 2004).

High-energy mechanical milling is a low-cost process for the production of nanostructured powders and applicable to a variety of advanced materials (Koch, 1997). In such a process the elemental blended powders are continuously welded and fractured to achieve alloying at the atomic level (Suryanarayana, 2004). By milling ceramic hard phase and metallic binder, refined composite powders are obtained (He et al, 1998, 2002). By varying the milling conditions, different sizes of the hard phase can be adjusted. Two different terms are most
Mechanical alloying describes the process when mixtures of powders (of different metals or alloys/compounds) are milled together. Thus, if powders of pure metals A and B are milled together to produce a solid solution (either equilibrium or supersaturated), intermetallic, or amorphous phase, the process is referred to as MA. Material transfer is involved in this process to obtain a homogeneous alloy. When powders of uniform (often stoichiometric) composition, such as pure metals, intermetallics, or pre-alloyed powders, are milled in a high-energy ball mill, and material transfer is not required for homogenization, the process has been termed mechanical milling (MM).

Mechanical alloying is a complex process involving optimization of a number of process variables to achieve the desired product phase, microstructure, and/or properties. For a given composition of the powder, some of the important variables that have an important effect on the final constitution of the milled powder are as: type of mill, milling container, milling energy/speed, milling time, size distribution of grinding medium, ball-to-powder weight ratio (BPR), extent of vial filling, milling atmosphere, process control agent (PCA), and temperature of milling. For more information about the milling parameters and the effect of these variables on the final product, refer to the book entitled “Mechanical Alloying and Milling” was written by C. Suryanarayana (Suryanarayana, 2004).

Powder particle size and morphology of the feedstock powder influence the melting conditions during spraying and therefore determine coating and microstructure formation. To obtain particles with dissimilar morphologies, powders can be produced by different processing routes including high-energy milling, spray drying and sintering, as depicted by Eigen et al; in Fig. 3. Route A includes milling under agglomerating conditions. If sufficient agglomeration is obtained directly in the milling process, further powder refinement only requires sieving to cut off larger size fractions. In an effort to minimize milling time, milling can therefore, also be stopped at an earlier state followed by a classifying process to yield particle sizes in the sprayable range, thus saving time and energy (route B). In a third route C, high-energy ball milling may be combined with spray drying and sintering, leading to spherical and more open powder morphologies with a finer and more homogeneous phase distribution as compared to conventional feedstock powders (Eigen et al; 2003, 2005).

3. Advanced materials and related thermal spray coatings

In thermal spraying technology, molten or semi-molten powders are deposited onto a substrate to produce a coating. The microstructure and properties of the materials depend on the thermal and momentum characteristics of the impinging particulate (Pawlowski, 2008), which are determined by both the spraying methodology and the type of feedstock materials employed. Various coatings are deposited on the surface of a substrate to either provide or improve the performance of materials in industrial applications. Nanostructured and advanced materials are characterized by a microstructural length scale in the 1–100 nm. More than 50 vol.% of atoms are associated with grain boundaries or interfacial boundaries when the grain is small enough. Thus, a significant amount of interfacial component between neighboring atoms associated with grain boundaries contributes to the physical and mechanical properties of nanostructured materials. Using nanostructured or advanced feedstock powders, thermal spraying has allowed researchers to generate coatings having
higher hardness, strength, and corrosion resistance than the conventional counterparts (He et al; 2002).

![Diagram of process routes for the production of thermal spray feedstock powders](https://www.intechopen.com)

Fig. 3. Schematic of process routes for the production of thermal spray feedstock powders (Eigen et al; 2003).

### 3.1 Intermetallic compounds

Self-bonding materials are widely used in the thermal spray industry. Because the coating to substrate bond is often the weakest link in thermal spray coating systems, the ability of self-bonding materials to give a tenacious, reliable bond to the substrate greatly enhances the adhesion and therefore the performance of entire coating systems. While a number of these materials exist, the most popular and widely used is the nickel–aluminium powders (Pawlowski, 2008; Steenkiste et al; 2002). The reason for the good adhesion of these powders on non-pretreated surfaces (preheated only at 80–100°C to eliminate condensation), is the nascent heat evolved by the exothermic reaction of Ni and Al which form nickel–aluminide during spraying (Devi et al; 1997). Different results are available concerning the completeness of the aluminide reaction during the spraying, and the type of aluminide formed. Phase composition in coatings sprayed using Ni–Al powders of different compositions with vacuum plasma spray (VPS) and air plasma spray (APS) methods were studied by Sampath et al. (Sampath et al; 1990). They described that the VPS coatings have a microstructure that results from the reaction between Ni and Al, and APS coatings contain the Ni and Al oxides from air or contain Al₂O₃ together with a solid solution of Al in Ni (α-Ni). It might be concluded that the spraying atmosphere (especially its oxygen content) has a major effect on final microstructure. The chemical reactions resulting directly in the coating phase composition take place during the particle flight in a flame (Chung et al; 2002; Hearley et al; 2000).

Movahedi et al; synthesised a thermal spray Ni–10 wt%Al powder consists of an aggregate containing the two components consisting of an alternative nickel and aluminium layers by
low energy ball mills, such as tumbler, whereby the components are milled together for extended periods to form homogeneous powders (Movahedi et al; 2005a, 2009). Such a layered structure can be readily produced by mechanical alloying of elemental powders (Eigen et al; 2003). A rigid bonding between the particles is caused by a cold welding when mechanical energy is applied to different kinds of powders (Chen et al; 1999). Such binderless composite powders, which can be thermally sprayed to form coatings on various substrates, are suitable for using in a thermal spray process (Maric et al; 1996).

Fig. 4. shows XRD patterns of Ni–10 wt%Al powders after different low energy milling times. X-ray diffraction patterns include only elemental Ni and Al peaks without any identification of oxides or intermetallic phases. In contrast, some researchers (Enayati et al; 2004) reported that ball milling of Ni75Al25 powder mixture in a high-energy ball mill (i.e., planetary), led to the formation of a Ni(Al) solid solution that transformed to nickel aluminide on further milling. The extent of plastic deformation of powder particles, the local increase of temperature, and also the increase in the density of lattice defects in low energy ball mills and therefore the mass transport by diffusion are smaller compared to those in high-energy ball mills, making it impossible to obtain nickel aluminide phase (Boldyrev & Tkacova, 2000; El-Eskandarany, 2001).

Fig. 4. X-ray diffraction patterns of Ni–10 wt%Al composite powders as received and after different milling times (Movahedi et al; 2009).

It is noted that Ni and Al XRD peaks have a lower intensity and higher width than those of initial powders due to the refinement of the crystallite size as well as an increase in the non-uniform internal strain of Ni and Al crystal lattices. Fig. 5. shows cross-sectional images of powder particles after different milling times. At the early stage of ball milling (10h), the Al particles were flattened and cold welding to the Ni particles. After 20h of low energy mechanical milling, the Ni particles also deformed plastically, and a typical lamellar structure consisting of pure Al and Ni layers with a layer thickness of ~10µm was formed (white areas are Ni and black areas are Al). On continuous ball milling, the layered structure refined so that after 35h of milling time the average layer thickness was ~5µm.
Fig. 5. Cross-sectional SEM images of Ni–10 wt-%Al composite powders after different milling times (Movahedi et al; 2005a, 2009).

Fig. 6 plots the average particle size of the powders on milling time. As milling time increased to 20h, the average Ni–10 wt-%Al particle size increased and reached to the maximum value of 300µm. For milling times longer than 20h, the average particle size decreased and finally approached a constant value of 5µm after 100 h of milling time. In addition, the powder particle size becomes more uniform by increasing the milling time.

Fig. 6. Average particle size of Ni–10 wt%Al composite powders versus milling time (Movahedi et al; 2009).

The changes of powder particle size during ball milling are caused by fracture and cold welding of Ni and Al powder particles during milling. The smaller particles grow while the larger ones fracture. A rigid bonding between the particles is caused by a welding when mechanical energy is applied to different kinds of powders (Chen et al; 1999). During milling the ingredients of the powder, mixtures are reduced in size and brought into intimate contact by flattening and crushing the particles, welding them together, and repeating the process again and again. The resultant powders essentially consist of a homogeneous and uniform distribution of the initial component within the powder particles (Maric et al; 1996). The particle size distribution of the powders has a major effect on quality and morphology of thermal spray coatings. It is generally accepted that the optimum particle size for thermal spray powders is within the range 35–100µm. These powders are large enough to be easily fed from simple hoppers but small enough for efficient melting to
occur. Under a given set of plasma conditions, there is likely to be an optimum size of particle that will be melted in the plasma, transported with sufficient momentum, and subsequently deposited. Smaller particles will lose velocity or may be vaporised with subsequent loss of spraying efficiency, whereas large particles will be incompletely melted and will produce a coating with large pore and low strength. In addition, the particle size significantly affects particle temperature and speed during flight, which subsequently influences the properties of the coating (He & Schoenung, 2002). Movahedi et al; reported that the Ni-10 wt%Al powders made by low energy ball milling have a microstructure consisting of alternative Ni and Al layers and include high defect density such as dislocations and vacancies. Therefore, the exothermic reaction between Ni and Al layers during flight in plasma spray process, readily occurs which subsequently enhances the bonding strength.

Fig. 7. shows the XRD pattern of plasma spray coating. It can be seen that plasma spray coating include NiAl intermetallic phase along with α-Ni solid solution. Significant feature of powders produced by mechanical milling is that a lamellar structure consisting of pure Al and Ni layers forms during milling process. This structure provides a large Ni/Al interfaces for exothermic reaction to occur during flight in plasma spraying. The incomplete Ni and Al reaction is due to the rapid heat losses of powder particles during thermal spraying (Movahedi et al; 2005b, 2009).

![Fig. 7. X-ray diffraction patterns of Ni-10 wt%Al plasma spray coatings (Movahedi et al; 2009).](www.intechopen.com)

Fig. 8. shows the cross-sectional microstructure images of the plasma spray coatings. The shape of the pores suggests that they were formed due to the expansion of trapped air when the impacting particles were still molten. Cavities and pores are formed between the individual particles in the sprayed coatings because inter-particle diffusion is limited and particle flow is hindered during the splat cooling process (Movahedi et al; 2009).

Ni-Al intermetallics are being recognized as high temperature structural materials because of their excellent oxidation resistance, high thermal conductivity, low density, and high melting point (Stoloff et al; 2000). Research on nickel aluminides has been expanded during the last 30 years, not only as bulk materials, but also as coatings. Studies have indicated that nickel aluminide alloys have significant potential in wear applications as wear properties of
carbon steel parts can be significantly improved by applying nickel aluminide coating (Goldenstein et al; 2004; Houben et al; 1973; Knotek et al; 1973; Rickerby et al; 1991). The possible applications of Ni-Al include: gas turbine engine, rotor blades, and stator vanes, however, low ductility at ambient temperature is the major limitation of this material. A number of attempts have been made to overcome this drawback such as nanocrystallization of structure which may transform nominally brittle compound into the ductile material (Morsi, 2001). One of the synthesis methods of nanocrystalline Ni-Al intermetallic compounds is high energy mechanical alloying. Two different mechanisms, via a rapid explosive reaction or through a gradual diffusion, were found for Ni-Al formation during MA (Atzmon, 1990; Enayati et al; 2008; Mashreghi et al; 2009). The powder prepared by MA can be deposited on surfaces of engineering parts using different thermal spraying techniques including plasma spray and, high velocity oxy fuel (HVOF) process. There are limited investigations in the literature concerning thermally sprayed coatings of nickel aluminides. Hearley et al. used inert gas atomized and reaction sintered Ni-30 wt.%Al powders to prepare NiAl intermetallic coatings by HVOF thermal spraying. They reported that a spherical inert gas atomized powder with narrow particle size range between 15 and 45 µm produced coatings of better quality. Their results also showed that both the fuel and oxygen flow rates influence the coating deposition characteristics and properties (Hearley et al; 1999, 2000).

Fig. 8. Cross-sectional SEM images of Ni–10 wt.%Al plasma spray coating at different magnifications (Movahedi et al; 2005b, 2009).

Fig. 9. shows the XRD patterns of Ni50Al50 powder mixture as-received and after 60, 90, and 120 min of high energy milling times. The XRD patterns of the as-received Ni50Al50 powder showed diffraction peaks of the crystalline Ni and Al. Increasing milling time to 60 min led to the disappearance of the Ni and Al peaks, while NiAl peaks began to appear. Complete transformation of elemental Ni and Al powder mixture to the NiAl intermetallic phase appeared to occur after 90 min of high energy ball milling. This result shows that the reaction between Al and Ni is promoted by the extensive Ni/Al interface areas as well as the short circuit diffusion paths provided by the large number of defects such as dislocations and grain boundaries introduced during high energy ball milling (Enayati et al; 2011). Hu et al. reported that complete transformation of Ni +Al to NiAl compound during MA occurred after 240h which is much longer than MA time obtained in the work of Enayati, et al. This discrepancy can be due to the different mill machines used.
Fig. 9. The development of NiAl intermetallic compound from Ni50Al50 powder mixture during mechanical alloying (Enayati et al; 2011).

The XRD patterns of the as-milled powder for 90 min and as-deposited coatings prepared with two sets of spray parameters (Table 1) are presented in Fig. 10. Besides NiAl main peaks several additional small, broad peaks can be observed on the XRD patterns of the coatings. These broad peaks were identified as Ni and Al$_2$O$_3$ phases. Enayati, et al. suggested that oxidizing of Al and subsequent separation of Ni occurred as NiAl particles are subjected to the high temperature (typically 3000 °C) during HVOF spraying. The intensity of Ni and Al$_2$O$_3$ peaks are higher in coating II as the fuel/oxygen ratio increases. It means that a higher fuel/oxygen ratio results a higher flame temperature and therefore more oxidation during HVOF processing.

<table>
<thead>
<tr>
<th>HVOF Parameters</th>
<th>Condition</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>I</td>
</tr>
<tr>
<td>Oxygen flow rate (l/min)</td>
<td>830</td>
</tr>
<tr>
<td>Fuel flow rate (ml/min)</td>
<td>210</td>
</tr>
<tr>
<td>Fuel/oxygen volume ratio</td>
<td>0.025</td>
</tr>
<tr>
<td>Spray distance (mm)</td>
<td>360</td>
</tr>
<tr>
<td>Powder rate (g/min)</td>
<td>80</td>
</tr>
<tr>
<td>Number of passes</td>
<td>3</td>
</tr>
</tbody>
</table>

Table 1. HVOF Spraying parameters for NiAl coatings (Enayati et al; 2011).

Fig. 11. shows the cross-sectional SEM images of the coatings at several magnifications. The coating exhibits a typical splat-like and layered morphology due to the deposition and resolidification of molten or semi-molten droplets. The light and dark gray layers are Ni-rich and Al-rich phases, respectively, which are consistent with the work of Movahedi, et al. (Movahedi et al; 2005B, 2009). Enayati, et al. suggested that an improved in uniformity of microstructure was observed for coating II due to the higher fuel flow rate and flame temperature (Enayati et al; 2011).
A dense Ni/Al alloy coating was deposited by cold spraying using a mechanically alloyed powder was reported by Zhang, et al (Zhang et al; 2008). Fig. 12. shows typical SEM images of cross-sectional microstructure of cold-sprayed Ni/Al alloy coating using milled Ni/Al alloy powder. It was observed that the coating exhibited a dense microstructure and some apparent thick layers with a white contrast appeared on the coating microstructure.
XRD patterns of the as-sprayed coating and feedstock powder are shown in Fig. 13. Only the peaks of nickel and aluminum were identified in XRD pattern of the ball-milled powder. It is clear that the XRD pattern of the cold spray coating is almost the same as that of the milled powder. This fact indicates that the coating and feedstock exhibited the same phase structure and no oxide was identified in the powder and the coating by XRD. In cold spraying, the particle deposition takes place in a solid state. Consequently, the lamellar structure of the milled powder will be completely retained in the coating, giving a unique effect on the microstructure and properties of the cold-sprayed coating. According to EDS analysis of the coating, was mentioned by Zhang, et al. the thicker layer in a white contrast was a Ni-rich phase and the fine lamella was a Ni-Al solid solution with high-Al content.

The XRD patterns of annealed Ni/Al cold spray coatings are shown in Fig. 14. After annealing treatment at 500°C for 3h, Ni and Al peaks completely disappeared and peaks corresponding to Ni$_2$Al$_3$ and NiAl appeared (Fig. 14a). With annealing temperature rising to 600°C, it can be found from Fig. 14(b) that NiAl became the main phase, only minor Ni$_2$Al$_3$ exists. As the temperature was raised to 850°C, the diffraction peaks of Ni$_2$Al$_3$ disappeared.
completely, and only diffraction peaks of NiAl phase were present in the XRD pattern, as shown in Fig. 14(c). This fact indicates that the annealing at temperature of higher than 850°C completely converts Ni/Al alloy to NiAl intermetallic compound. As the temperature reached 1050°C, no additional reaction was detected, and the NiAl phase was present in the coating (Fig. 14d) (Zhang et al; 2008).

Fig. 14. XRD patterns of the Ni/Al cold spray coatings annealed at different temperatures (Zhang et al; 2008).

### 3.2 Intermetallic-ceramic and nanocomposites

As mentioned earlier, intermetallic compounds are an important class of materials because of a combination of their high tensile strength, low density, good wear, and creep resistance. These properties have led to the identification of several potential usages including structural applications and protective coatings (Sauthoff, 1995). Two major problems that restrict the application of intermetallic compounds are poor low-temperature ductility and inadequate high temperature creep resistance. These limitations can be overcome by introducing ceramic particles as reinforcements (Morris, 1998). Originally, reinforcement phase can be introduced in the matrix by two routs namely ex-situ addition of reinforcement particles and in-situ formation of reinforcement phase via a displacement reaction which both phase (Intermetallic and ceramic) are formed during ball milling. A rigid bond between the particles is created by cold welding when mechanical energy is applied to powder particles (Chen et al; 1999). Incorporation of hard second phases into an intermetallic composite (IC) matrix is a strategy for effective high-temperature strengthening, creating an intermetallic matrix composite (IMC). Recently, a great deal of work has been done on intermetallic matrix composites (IMCs). Various continuous or discontinuous ceramic reinforcements such as SiC, Al₂O₃, TiB₂, and TiC were explored to obtain increased high temperature strength and better creep resistance, together with adequate ductility and toughness (Inoue et al; 2000). Among these reinforcements, SiC fibers were commercialized for use in IMCs. The SiC reinforcements were added into different nickel aluminate matrices by reaction synthesis, mechanically alloying, and sintering (powder metallurgy) (Lee et al; 2001; Zhang et al; 2004) to improve oxidation and mechanical properties and the workability.
of the matrices. Hashemi et al. reported the synthesis of nickel aluminide matrix composite coating reinforced by SiC particulates that was fabricated by the plasma spraying of Ni-Al-SiC powder prepared by low energy ball milling. The cross-sectional SEM image of the powder particles after 15h of milling time is presented in Fig. 15. As it can be shown the SiC particles were incorporated into the Ni/Al powder particles.

![Cross-sectional SEM image of Ni-Al-SiC powder particles](image)

Fig. 15. Cross-sectional SEM images of Ni-Al-SiC powder particles produced by low energy ball milling (Hashemi et al; 2009).

The formation of Ni-Al intermetallic compounds required a long time and a high temperature for the diffusion of Al and Ni. In thermal spray processes, the temperature is high enough for diffusion, but the exposure time of powders to plasma flame is too short. After powder deposition on the substrate, the high-temperature diffusion and therefore the Ni-Al reaction is stopped (Houben et al; 1973; Sampath et al; 2004). Hashemi et al, suggested that by increasing the plasma spray distance the diffusion time increased, while by increasing current density the plasma flame temperature increased. These two parameters determine the content of intermetallic compound in the coatings. They also mentioned that by increasing the current intensity from 600 to 700 A, the relative amount of Ni decreased while that of Ni₂Al₃ phase increased, whereas by increasing the current intensity from 700 to 800 A, a reverse trend was observed. On the other hand, the increased current density will increase the velocity of the powder flow rate, thus declining the dwelling time of the powder particles and further decreasing the amount of Ni₂Al₃. In this approach, powder particles experience lower heat, which eliminates the diffusion of Ni and Al and therefore the development of Ni-Al compounds. The cross-sectional microstructure of as-sprayed coating in optimum spray condition is shown in Fig. 16. (Hashemi et al; 2009).

Horlock, et al. synthesised a reactive powder having a nominal composition of 50wt.%Ni(Cr)-40wt.%Ti-10wt.%C with the planar-type ball milling. They reported that the reaction to produce TiC is initiated within individual powder particles during HVOF spraying, leading to the formation of a coating containing TiC particles within a nanocomposite Ni-rich matrix. The TiC particles were found in the coating (Fig. 17) on the order of 50 to 200 nm in size. The Low-magnification SEM image of a cross section through an HVOF-sprayed coating shows the characteristic layered morphology with little porosity as well as the nanoscale TiC grains (arrowed), which are embedded in a light contrast metallic matrix.
Fig. 16. Cross-sectional SEM images of plasma spray Ni-Al-SiC coating at different magnification (Hashemi, 2009).

Fig. 17. (a) Low-magnification (b) high-magnification BSE image of a region from (a) of a cross section through an HVOF-sprayed Ni(Cr)-TiC coating (Horlock et al; 2005).

In Horlock’s work a pre-alloyed Ni(Cr) powder was used to produce a metallic matrix with the potential to resist corrosion and high-temperature oxidation, while TiC is a ceramic with high hardness and chemical stability. They also proposed that the final powder microstructure contains Ti, Ni(Cr), and C all embedded in the same particle, there is a decreased possibility of fresh Ti surfaces being in contact with free C. Consequently, the possibility of Ti oxidation of TiC is greatly diminished during spraying. The XRD pattern from the as-deposited coating, shown in Fig. 18. has major peaks that can be identified as a Ni-rich solid solution phase and TiC as well as the smaller peaks corresponding to NiTi, TiO₂, and NiTiO₃ (Horlock et al; 2004).

TiC-Ni based nanocomposite powders for thermal spraying were produced by high-energy attrition and vibration mills in scale up were studied by Eigen, et al. X-ray diffraction analysis shows (Fig. 19) that the crystallite sizes of both hard phase and binder are refined with increasing milling time reaching a steady state after about 20 h.

Correspondingly, the mixing and refining of the phases can be observed by SEM (Fig. 20). At early stage of milling (Fig. 20a), the ductile binder particles are deformed to plate-like particles while hard particles are initially pressed into the surface of the binder platelets and
cover them. In later milling stages, densification takes place, i.e. hard phase particles are fully embedded into the matrix and the matrix is completely cold-welded (Fig. 20b). In the final stage in Fig. 20c, microstructures are characterized by carbide phase dimensions ranging from less than 20 nm (see Fig. 19) to about 500 nm (Eigen et al; 2003).

The development of a nanocomposite thermal spray powder is also studied by He, et al. as schematically summarized in Fig. 21. In a nanocomposite powder, such as Cr$_3$C$_2$-NiCr and WC-Co, there are hard and brittle carbide particles and a tough metal binder constituent. Hard and brittle carbide particles are fractured into sharp fragments and embedded into the metal binder. The metal binder, with lower hardness, is subjected to enhanced milling from
both the balls and hard carbide particles. As milling time increases, carbide fragments are continually embedded into the metal binder. The metal binder and the polycrystalline composite, experience continuous overlapping, cold welding, and fracturing. Finally, a polycrystalline nanocomposite powder system, in which round nanoscale carbide particles are uniformly distributed in a metal binder, is formed. As an example, Fig. 22 shows such a Cr$_3$C$_2$-25 (Ni20Cr) polycrystalline nanocomposite powder. Clearly shown that the large proportions of carbides, in the form of round particles, are uniformly distributing themselves in the NiCr solid solution. It is possible to use mechanical milling to synthesize other nanocomposite powder systems with a hard particle and tough binder duplex structure; examples of such systems are WC-NiCr, TiC-NiCr, TiC-Ti, and SiC-Al. The microstructures of conventional and nanostructured Cr$_3$C$_2$-25 (Ni20Cr) coatings, examined using SEM, are shown in Fig. 23. A uniform and dense microstructure is observed in the nanostructured coatings, compared to the conventional Cr$_3$C$_2$-25 (Ni20Cr) coating that is observed to have an inhomogeneous microstructure (He et al; 2002).

Fig. 20. Refinement of phase distribution after: (a) 2 h; (b) 10 h; and (c) 40 h (BPR 23:1) in the vibration mill; and (d) after 20 h in the attrition mill (BPR 20:1). Black arrows in (a) indicate fragments of one broken hard phase particle (Eigen et al; 2003).
Fig. 21. Schematic diagram of milling mechanism for duplex structure powder: (a) initial stage; (b) NiCr matrix overlaps and deforms, Cr$_3$C$_2$ fractures and embed into NiCr; (c) binders deform, fracture, and weld, carbide fracture further; and (d) nanocomposite powder (He et al; 2002).

Fig. 22. Cr$_3$C$_2$–NiCr powder milled for 20 h: (a) Bright field image; and (b) dark field image. White particles are Cr$_3$C$_2$, and dark matrix is NiCr (He et al; 2002).
3.3 Amorphous-nanocrystalline materials

Amorphous metallic alloys have been of interest not only for fundamental studies, but also for potential applications for over 40 years. Amorphous structures have been made in many alloy systems and show a variety of unique properties compared to their crystalline counterparts. These properties are associated with the amorphous atomic structure and include high yield strength, large elastic limit, high corrosion resistance, good wear resistance, and low elastic modulus (Greer et al; 2002; Schuh et al; 2007). Amorphization by high energy mechanical alloying of elemental powders occurs by an inter-diffusion reaction at relatively low temperature along constituent interfaces. The formation of amorphous phase by MA process depends on the energy provided by the milling machine and thermodynamic properties of the alloy system. The thermodynamic and kinetic principles for amorphization through solid state synthesis or mechanical alloying are discussed by Schwarz and Johnson (Schwarz & Johnson, 1883). They identified two rules for the formation of amorphous alloy by MA in an A–B binary system: (1) A large negative heat of mixing, $\Delta H_{\text{mix}}$, between the elemental constituents. (2) A large asymmetry in the diffusion coefficients of the constituents. An amorphous phase is kinetically obtained only if the amorphization reaction is much faster than that for the crystalline phases. It is also believed that during mechanical milling of a homogeneous crystalline alloy, the internal energy of lattice increases due to the introduction of crystal defects. When the free energy of the crystalline structure exceeds the free energy of the amorphous phases, crystalline structure can transform to an amorphous phase (Suryanarayana, 2001, 2004).

Fe-based amorphous alloys and thermal spray coatings are perhaps the most important system for possible applications because of the low cost of iron, and the relatively high strength and hardness of Fe-based amorphous alloys (Chen et al; 2006; Sunol et al; 2001). Extensive research has been carried out on the mechanical alloying of Fe-based amorphous alloys including binary Fe–B, Fe–Cr, and Fe–Zr (Schuh et al; 2007), ternary Fe–Zr–B (Suryanarayana, 2001, 2004) and Fe–Si–B (Chen et al; 2006) and multi-component Fe–Ni–Si(P)–B (Chen et al; 2005), Fe–Al–P–C–B (Minic et al; 2009) and Fe–Nb–Cu–Si–B (Inoue et al; 2000) alloys. Movahedi et al, reported the MA amorphization of 70Fe–15Cr–4Mo–5P–1C–
1Si–4B (wt.%) elemental powder which includes four types of elements: late transition metal (Fe), early transition metals (Cr, Mo), metalloids (B, P, Si), and graphite. The atomic sizes of these elements are in the order of Mo (0.139 nm) > Si (0.132 nm) > P (0.128 nm) > Cr (0.127 nm) > Fe (0.126 nm) > B (0.098 nm) > C (0.091 nm). This composition yields new atomic pairs of Fe–(Cr, Mo), (Cr, Mo)–(B, P, Si) and Fe–(B, P, Si) with various negative heats of mixing. These properties suggest that Fe–Cr–Mo–B–P–C–Si composition has a high glass forming ability (GFA) and appropriate thermal stability (Movahedi et al; 2010a & b).

Fig. 24. shows the XRD patterns of powder mixture as a function of high energy milling (i.e., Retch PM100) times. As-received powder mixture shows sharp crystalline peaks of elemental Fe, Cr, Mo, B, C and Si. Red phosphorus is absent on XRD pattern because of its amorphous nature. As milling progresses, the XRD peaks of the elemental constituents are broadened with a corresponding decrease in their intensities. These effects are caused by a continuous decrease effective crystallite size and an increase of the atomic level strain, as a result of the induced-plastic deformation during MA (Filho et al; 2000; Zhang, 4004). After 15h of milling time, the Cr, Mo, Si, B and C peaks vanished. This may be due to the dissolution of these elements into Fe matrix and/or to their ultra fine crystallite size. There is no significant reaction between the elemental powders at this stage of milling as no new phase was detected. On continued milling a broad peak was developed on the XRD pattern, owing to the formation of an amorphous phase. Meanwhile the crystalline peaks of Fe remain distinct. This structure transforms to an amorphous phase on further milling till 80h of milling time (Movahedi et al; 2010a).

Fig. 24. XRD patterns of 70Fe–15Cr–4Mo–5P–1C–1Si–4B powders as-received and after different milling times (Movahedi et al; 2010a).

Movahedi et al. suggested that the transformation of supersaturated solid solution of Fe to amorphous structure during milling is believed to occur as a result of the internal energy increase of the crystalline structure due to the creation of a high density of lattice defects as well as the dissolution of a great amount of solute atoms with different sizes in Fe lattice. When the free energy of the crystalline solid solution exceeds the free energy of the amorphous state, the crystalline structure thermodynamically becomes unstable and
transform to the amorphous structure. This behavior can be accounted for by considering the interaction parameters, proposed by Inoue et al. (Inoue et al; 2000) which describe the difference in bonding energy between the atomic pairs Fe–P–C and Fe–B–C in the ternary Fe–A–C system. The effect of B addition is presumably to generate attractive bonding among the constituent elements. In the Fe–A–C system with a negative interaction parameter, the enthalpy of mixing is also negative. In such a case, the formation of the Fe–A–C solution decreases the free energy of the system by lowering the system mixing enthalpy. These interaction parameters characterize the effect of a third element on amorphization reaction of Fe–C binary alloys processed by mechanical alloying (Inoue et al; 2000; Olofinjana et al; 2007; Suryanarayana, 2001).

Fig. 25. shows cross-sectional SEM images of the powder particles after 2, 4, 10, 15, 30 and 80h of milling. During the early stage of milling, the powder particles are flattened by compressive forces due to the collision of the balls. Thus, micro-forging action deforms the powder particles plastically leading to work hardening and fracturing. The creation of new surfaces enable the particle to weld together and thus to produce a typical lamellar structure consisting of pure elements (Movahedi et al; 2009). On continued ball milling, the layered structure is progressively refined (Fig. 25d–f). At longer milling times the structure became featureless on SEM as a result of development of Fe-base solid solution and subsequent amorphous phase.

![Fig. 25. cross-sectional SEM images of 70Fe–15Cr–4Mo–5P–1C–1Si–4B powders after different milling times (Movahedi et al; 2010a).](image)

High resolution transition electron microscopy (HRTEM) and selected-area diffraction pattern (SADP) of powders milled for 15h confirmed the formation of a nanocrystalline structures (Fig. 26a). After 40 h of milling time amorphous and nanocrystalline phases co-existed in the milled powders. Fig. 26b shows that most amorphous phase is developed at the edge of powder particles indicating that the amorphization reaction starts at edge of
particles and progress into the internal regions as MA proceeds. The SADP in Fig. 26b shows some diffraction spots within the amorphous diffuse ring. Furthermore the fast Fourier transform (FFT) images showed a broad diffuse ring at the edge and crystalline diffraction spots at the center of particles (arrows in Fig. 26b) suggesting that the sample contains both amorphous and nanocrystalline phases. Fig. 26c is the HRTEM image and SADP of mechanically alloyed powder after 80h of milling time, showing a fully amorphous microstructure (Movahedi et al; 2010a, 2011).

Synthesizing amorphous and/or nanocrystalline layers on metal substrates can be utilized to improve surface performance such as wear and corrosion resistance (Kim et al; 2007). Greer et al. reported that amorphous alloys can have very good resistance to sliding and abrasive wear and the coatings can have low friction coefficient (Greer et al; 2002). Thermal spraying process is one of the techniques to deposit amorphous coatings on surfaces, where the amorphous structure is retained due to the sufficiently rapid cooling that inhibits long-range diffusion and crystallization. On impact with the substrate, droplet spreading occurs to give lamellar morphologies with cooling rates of $10^7$-$10^8$ K/s (Wu et al; 2006). A number of researchers have investigated the use of air plasma spraying (APS), low pressure plasma
spraying (LPPS) and vacuum plasma spraying (VPS) to deposit alloys, which are capable of solidifying as metallic glasses. Kishitake et al. reported that mixed amorphous and crystalline structures are obtained in APS and LPPS coatings (Kishitake et al; 1996). In recent years, there has been an increasing interest in the use of HVOF for depositing protective coatings. The general methodology involves designing alloys that have low critical cooling rates for glass formation. Alloys with high glass forming ability (GFA) would be favorable for forming fully amorphous phase coating by HVOF process.

The amorphous coatings, while exhibiting interesting properties, can be heated up above their crystallization temperature to initiate devitrification and yield amorphous-nanocrystalline mixture. Since the driving force for the crystallization is extremely high and the diffusion rate in the solid state, at the crystallization temperature, is very low, an extremely high nucleation frequency results. There is limited time for growth before impingement between neighbouring crystallites occurs, resulting in the formation of nanoscale microstructures. Some researchers (Branagan et al; 2001; Kishitake et al; 1996; Otsubo et al; 2000), mentioned that the formation of Fe-base amorphous coatings by LPPS, high-energy plasma spraying (HPS), and HVOF processes with using atomized feedstock powder. In recent years, as mentioned earlier, Movahedi, et al. first synthesised the amorphous mechanical alloying feedstock powder in a new composition 70Fe-15Cr-5Mo-5P-4B-1C-1Si (wt.%) with high GFA and then developed them to amorphous-nanocrystalline coatings with HVOF and plasma spray process (Movahedi, 2010c, 2011).

Fig. 27. illustrates the XRD patterns of mechanically alloyed Fe-Cr-Mo-P-B-C-Si feedstock powder and the as-sprayed HVOF coatings. The presence of halo on XRD pattern confirms that the feedstock MA powder used for HVOF spraying has an amorphous structure. The XRD pattern of HVOF-G1 coating also shows a halo characteristic indicating that this coating has an amorphous structure similar to feedstock MA powder. However, in HVOF-G2 there is an emergent crystalline peak on the top of the amorphous hub suggesting that this coating is a mixture of amorphous and crystalline phases. Structure of HVOF-G3 coating mainly consists of crystalline phases such as $\alpha$-Fe, $\text{Fe}_2\text{C}_6$, and $\text{Fe}_5\text{C}_2$.

![XRD patterns of mechanically alloyed feedstock powder and Fe-Cr-Mo-P-B-C-Si HVOF coatings](http://www.intechopen.com)
Movahedi, et al. inferred from the diffraction patterns that a full range of amorphous to fully crystalline microstructures can be obtained by adjusting of HVOF parameters especially fuel/oxygen ratio (Table 2). They suggested that the difference in the glassy fraction is related to the mechanism of the amount of cooling rate and remelting of individual particles in HVOF flame at various fuel/oxygen ratios. By increasing fuel/oxygen ratio, the flame temperature becomes too high and the velocity of the powder particles is much higher. Thus, the powder particles were completely remelted in the HVOF flame and then were rapidly solidified and quenched on the cold substrate forming an amorphous structure. By decreasing the fuel/oxygen ratio the particles did not remelt to a significant extent as well as the velocity of them becomes lower, thus the conditions appear to crystallize the amorphous feedstock powder in flame which explains the higher percentage of crystalline phase in the HVOF-G3 (Movahedi, 2010c, 2011; Shin et al; 2007).

Kishitake et al. reported that for Fe-17Cr-38Mo-4C gas atomized powder, amorphous coatings are obtained by the APS while a mixture of amorphous and crystalline phases are formed by HVOF (Kishitake et al; 1996). They suggested that this difference may result from the difference of the cooling rate between the APS and HVOF processes. It is regarded that the composition chosen by Movahedi, et al. satisfies the three empirical rules (Pang et al; 2002) for the stabilization of the super-cooled liquid during HVOF and plasma spraying, leading to highly dense random packed atomic configurations, higher viscosity, and lower atomic diffusivity (Kobayashi et al; 2008) which is primarily attributed to the high GFA.

<table>
<thead>
<tr>
<th>Parameters</th>
<th>Microstructure</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Amorphous (HVOF-G1)</td>
</tr>
<tr>
<td>Oxygen gas flow rate (SLPM)</td>
<td>833</td>
</tr>
<tr>
<td>Fuel (Kerosene) flow rate (SLPM)</td>
<td>0.37</td>
</tr>
<tr>
<td>Fuel/Oxygen (Vol%)</td>
<td>0.044</td>
</tr>
<tr>
<td>Powder feed rate (g/min)</td>
<td>35</td>
</tr>
<tr>
<td>Spray distance (mm)</td>
<td>300</td>
</tr>
<tr>
<td>Scanning Velocity (mm/s)</td>
<td>50</td>
</tr>
<tr>
<td>Deposit thickness (µm)</td>
<td>300</td>
</tr>
<tr>
<td>Nozzle length (mm)</td>
<td>100</td>
</tr>
<tr>
<td>Compress air cooling</td>
<td>yes</td>
</tr>
</tbody>
</table>

Table 2. HVOF spraying parameters for advanced structures (Movahedi et al; 2010c, 2011).

Typical SEM cross section image of HVOF coatings is shown in Fig. 28. As it can be seen, the microstructure of coatings includes very fine lamella structure which is smooth and dense, adhering well with the substrate with no cracking. Moreover, some pores are rarely observed in this microstructure as indicated by arrows can be seen from the images. The big pores located between flattened droplets are mainly caused by the loose packed layer structure or gas porosity phenomenon, while the small pores within the flattened particles originate from the shrinkage porosity (Totemeier, 2005). Obviously, the porosity of the coatings reduces in the order of HVOF-G3, G2, and G1, and it is believed that increasing the fuel/oxygen ratio, increases both the thermal and kinetic energy of the gas flow, so that the majority of the powder particles are better melted and also accelerated to higher velocities.
and deformed extensively on impact to form elongated lamella (Ji, 2005). Some unmelted particles are clearly visible in HVOF-G3 (Fig. 28c) coating because of lower flame temperature (minimum fuel/oxygen ratio).

![Fig. 28. SEM cross-sectional images of Fe-Cr-Mo-P-B-C-Si coatings (a) HVOF-G1, (b) HVOF-G2, and (c) HVOF-G3 (Movahedi et al; 2010c).](image)

HRTEM image has shown in Fig. 29. confirms that HVOF-G1 coating is completely amorphous. As it can be seen in Table 2, this microstructure appears when the fuel/oxygen ratio has a maximum value. As shown in Fig. 30. the HVOF-G2 coating consists of amorphous phase and nanocrystalline grains. The electron diffraction pattern, in Fig. 30(a) was taken with the selected area aperture centered over the amorphous and nanocrystalline region and shows a diffuse amorphous halo with diffraction spots arising from nanocrystalline grains with a size range of 5-30 nm. The HRTEM micrograph and fast Fourier transform (FFT), as shown in Fig. 30(b), confirm the presence of a mixture of nanocrystalline grains within an amorphous matrix. In this case the fuel/oxygen ratio is moderate (HVOF-G2) so this duplex microstructure can be explained by quenching of semimolten particles when impinged to the cold substrate. Therefore, some unmelted particles crystallized inside the HVOF flame to yield nanocrystalline grains which embedded within the amorphous matrix. A nanocrystalline structure with equiaxed nanograins was obtained in case of HVOF-G3 coating (Fig. 30c). In this condition the fuel/oxygen ratio has a minimum value and the HVOF flame temperature is the lowest so the most of the individual powder particles were unmelted and crystallized inside the HVOF flame. Moreover, the cooling rate was sufficiently high to avoid grain coarsening and yielded nanocrystalline structure (Movahedi et al; 2010c, 2011).
Fig. 29. HRTEM micrograph and SADP of amorphous Fe-Cr-Mo-P-B-C-Si HVOF coating (HVOF-G1) (Movahedi et al; 2010c, 2011).

Fig. 30. (a) TEM and (b) HRTEM micrographs, SADP and FFT of amorphous-nanocrystalline Fe-Cr-Mo-P-B-C-Si HVOF coatings (HVOF-G2), and (c) TEM, HRTEM micrographs and SADP of nanocrystalline of HVOF-G3 (Movahedi et al; 2010c, 2011).
Movahedi, et al. reported that the microhardness value of HVOF coatings are all have a high hardness of around 800 to 1200 HV. The value of hardness was similar to that of the bulk Fe-based metallic glass (Kobayashi et al; 2008) but it was higher than that of electroplated chromium and the Ni-based amorphous coatings (Ni et al; 2009). The difference in hardness value of the three groups of HVOF coatings were developed by Movahedi, et al. is attributed mainly to the difference in volume fraction of amorphous and nanocrystalline phases. A fully amorphous coating has lower hardness (830 HV) compared to the duplex amorphous-nanocrystalline coating (950 HV). The fully nanocrystalline coating has the highest microhardness (1230 HV), probably due to precipitation of some carbides such as Fe<sub>23</sub>(C, B)<sub>6</sub> and Fe<sub>5</sub>C<sub>2</sub> during crystallization (Movahedi, 2010c). Some researchers suggested that the hardness of the amorphous Fe-base coating increases after crystallization (Branagan et al; 2005). In contrast Kishitake et al. reported that the duplex microstructure consisting of the both amorphous and nanocrystalline structure exhibits a higher hardness than fully amorphous or nanocrystalline structure (Kishitake et al; 1996). The difference is mainly attributable to the difference of decomposition of amorphous phase during crystallization.

4. Conclusion

This chapter reviews the solid state synthesis and characterization of advanced feedstock powders used in various thermal spray techniques, and processing and characterization of amorphous-nanostructured related coatings. The published results show that mechanical milling can be effectively used to synthesize advanced materials and nanocomposite powders. On the other hand, whether a composite or a single-phase starting powder is involved, mechanical milling leads to the formation of nanocrystalline structure under certain milling conditions.

5. References


Recently, plasma spray has been received a large number of attentions for various type of applications due to the nature of the plasma plume and deposition structure. The plasma gas generated by the arc, consists of free electrons, ionized atoms, some neutral atoms, and undissociated diatomic molecules. The temperature of the core of the plasma jet may exceed up to 30,000 K. Gas velocity in the plasma spray torch can be varied from subsonic to supersonic using converging-diverging nozzles. Heat transfer in the plasma jet is primarily the result of the recombination of the ions and re-association of atoms in diatomic gases on the powder surfaces and absorption of radiation. Taking advantages of the plasma plume atmosphere, plasma spray can be used for surface modification and treatment, especially for activation of polymer surfaces. Addition, plasma spray can be used to deposit nanostructures as well as advanced coating structures for new applications in wear and corrosion resistance. Some state-of-the-art studies of advanced applications of plasma spraying such as nanostructure coatings, surface modifications, biomaterial deposition, and anti wear and corrosion coatings are presented in this book.

How to reference
In order to correctly reference this scholarly work, feel free to copy and paste the following:
