Production and Characterization of Al-Si Coatings Fabricated by Mechanical Alloying Method on Inconel 625 Superalloy Substrates

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Abstract: Inconel superalloys are used substantially in high-temperature environments. However, these alloys suffer from corrosion and wear. Attempts to overcome these drawbacks involve coating the metal with different techniques and materials. In this study, a new method with increasing potential was utilized. Using the mechanical alloying process in a planetary ball mill vial, alloying and the Al-Si coatings were concurrently achieved on Inconel 625 substrates. Different process control agent (PCA) ratios, milling ball diameters, and milling times were used to improve coating properties. Macro and microstructure, morphology, microhardness, and roughness values of samples were evaluated and compared. Additionally, crystallographic and cross-sectional properties were investigated in order to optimize the processing conditions. The results indicated that increasing the diameter of the grinding ball enhanced the hardness and thickness of these coatings and increased the roughness values. Longer processing time also enhanced the thickness with mechanical values. However, under these conditions, coating homogeneity decreased, and incompatible regions were formed on the coatings. PCA content brought a refined grain structure, hence showed better mechanical properties. On the other hand, processing time should be increased to get a denser and thicker protective layer against the operational conditions.

Keywords: mechanical alloying; Inconel 625; coating; characterization; Al-Si alloy

1. Introduction

Due to their excellent high-temperature creep and mechanical properties, superalloys are widely used in aerospace, especially in gas turbine engines and energy production systems exposed to extremely high temperatures. However, exposure to long-term oxidant and corrosive environments calls attention to another important property of these materials, surface oxidation [1]. The corrosion properties of these materials could be improved by adding sufficient amounts of alloying elements like chromium, aluminum, or silicon to form \( \text{Cr}_2\text{O}_3 \), \( \text{Al}_2\text{O}_3 \), and \( \text{SiO}_2 \) on the surface, respectively [2]. The best way to achieve a protective surface is by applying a coating of protective layer(s). High-temperature coatings, i.e., diffusion [3,4], overlay [5], and thermal barrier coatings (TBC) [6], are mainly preferred for providing these materials with advanced protection for decades.

High-temperature coatings produced on nickel-based superalloys can be classified into three different material combinations. These are aluminides (M-Al), modified (with precious metals) aluminides (M, Pt)Al or (M, Pd)Al [7], and MCrAlX alloys (Ni, Co or Fe as M), and (Y, Zr or Si as X). Additionally, there are TBC systems containing a ceramic top layer, e.g., \( \text{ZrO}_2 \), together with the coatings mentioned above [8,9]. Moreover, Al-Si coatings produced on these high-temperature alloys were reported to improve the high-temperature oxidation resistance of these alloys [10,11].

Silicon addition was found as an effective way to improve the oxidation resistance of aluminide coatings. Low oxygen pressure fusing [10], powder pack cementation [12], hot
dipping [13], slurry method [14], and hot stamping [15] methods were studied to form Al-Si coatings on different substrate materials to improve oxidation resistance. The formation of an additional oxide phase (SiO$_2$) further benefits the protection of the nickel-based superalloy substrate, by delaying the formation of harmful phases (NiO and $\gamma^\prime$-Ni$_3$Al) [16]. In addition, Fu et al. (2014) reported that both coating elements react with nickel and form Ni$_2$Si and Ni$_3$Al, and the addition of Si promotes the formation of higher Al content NiAl phase (which is favorable to form protective Al$_2$O$_3$) from Ni$_3$Al [11]. Zang et al. also reported that adding silicon helps to form a dense Al$_2$O$_3$ layer, and sets a barrier to oxygen diffusion [13]. Mechanical alloying (MA), also known as mechanical milling, is a solid-state powder mixing and powdered alloy production [17] technique. This technique produces homogeneous equilibrium or non-equilibrium alloy phases using mixtures of different powders, including at least one ductile metal, in an appropriate ball milling device. The particles collide with each other with the help of milling balls. The impact and sliding friction energy make the powders cold weld with severe deformation. After this severe deformation, particles fracture, and fine, in situ alloyed structures are obtained [18–20]. As a coating process, MA has recently become an alternative and attractive operation for producing in situ obtained alloy layers directly by using a powder mixture on bulk substrates to improve properties like mechanical, wear, oxidation, thermal shock, and thermal conductivity [21–28].

MA is very dependent on many process parameters as well as the properties of the powders used in the blend [29]. So far, only parameters like substrate hardness and the type of process device have been examined [30,31] for coatings produced by mechanical alloying. In this present study, the effect of grinding ball size, process control agent (PCA) content, and milling time were investigated as coating process parameters. These are some of the first parameters that come to mind when mechanical alloying is the subject [32,33]. In addition, an Al-Si powder mixture was used to produce a dense, homogeneous, strongly adhering, and potentially protective intermetallic alloy coating on Inconel 625 superalloy surface by using a planetary ball mill.

2. Materials and Methods

2.1. Materials

Inconel 625 superalloy discs with 2.5 mm thickness were cut from a rod with 12 mm diameter and were used as a substrate material for coating studies. Stearic acid (SA) was used as the process control agent (PCA), to obtain the balance between cold welding and fracturing of the particles while mechanical alloying, and enhance the process efficiency [33]. High purity Al powder (Sigma, St. Louis, MO, USA: 7429-90-5) with 60 $\mu$m max. particle size and Si powder (Sigma: 7440-21-3) with 45 $\mu$m max. powder size were used as mechanical alloying input, with a weight ratio of 85/15 respectively to maintain a hypereutectic alloy structure (see Table 1).

| Material   | Ni  | Cr | Mo | Al  | Si  | Others |
|------------|-----|----|----|-----|-----|--------|
| Inconel 625| 60±2| 21 | 9  | 0.4 max | 0.5 max | bal. |
| Al Powder  | -   | -  | -  | 99.9 | -   | bal.  |
| Si Powder  | -   | -  | -  | -   | 99  | bal.  |

2.2. In Situ Mechanical Alloying and Coating Production

For mechanical milling operations, a planetary ball mill device (Retsch PM 100, Hahn, Germany) with a standard 500 mL hardened steel grinding jar and 200 g of grinding balls (Retsch, Hahn, Germany; 58 to 63 HRC hardness) were used. Grinding balls with different diameters (5 and 10) were selected in order to understand the effects of ball size on the homogeneity of the coating layer. Powder to ball ratio was specified as 1/10 (20 g of metal
powder), different PCA ratios, and milling times were selected to examine the effect of these parameters (Table 2). The milling processes were performed in ambient conditions with a milling speed of 250 rpm. In order to avoid the rise in temperature during the coating process, 5 min of coating application was followed by 5 min of cooling of the vial (see Figure 1).

Table 2. Specifications of mechanically alloyed samples.

| Sample No | Ball Diameter (mm) | PCA (SA) wt.% | Milling Time (Hour) |
|-----------|--------------------|---------------|---------------------|
| 1         | 10                 | 0             | 6                   |
| 2         | 10                 | 0.25          | 6                   |
| 3         | 10                 | 0.5           | 6                   |
| 4         | 5                  | 0             | 6                   |
| 5         | 5                  | 0.25          | 6                   |
| 6         | 5                  | 0.5           | 6                   |
| 7         | 5                  | 0.25          | 12                  |
| 8         | 5                  | 0.5           | 18                  |
| 9         | 5                  | 1             | 18                  |

Following each coating process, in order to prevent contamination, the grinding vial and balls were cleaned by using sodium hydroxide solution for aluminum cleaning. Aluminum reacts with sodium hydroxide to produce aluminum hydroxide and becomes easily removable from the system. For further cleaning, quartz sand was used in the vial with grinding balls, and milling was applied for 20 min. Finally, methanol was used to get a perfectly cleaned surface on the vial and balls.
2.3. Characterization

After production of the coatings, remaining mechanically alloyed powders were examined to understand their alloying performance. The particles were prepared using a standard specimen preparation procedure by mounting in two-component epoxy resin, grinding from 80 to 1200 grit sandpaper, and polishing with 3 μm diamond paste. Their dimensional, microstructural, and mechanical properties were identified using a stereomicroscope (ZEISS SteREO Discovery, AG, Jena, Germany), an optical microscope (Nikon Eclipse ME 600, Tokyo, Japan), and a micro-hardness tester (Shimadzu HMV-2, Tokyo, Japan).

Cross-sections of coatings were prepared with the same specimen preparation route. To protect the coating from any harm, grinding was applied parallel to the interface. The specimens were investigated using SEM (JEOL JSM 6060, Tokyo, Japan) and EDS to observe the substrate-coating interface, adhesion, and homogeneity of the alloyed coating. The overall hardness values of coatings were measured by using a 1-kg load (HV₁). Average coating thicknesses were calculated randomly from all over the cross-section profiles of coatings (both from narrower and thicker parts), and the average of the 20 measurements by using an optical microscope. Cross-sectional hardness profiles were prepared by using five hardness values, and these values were obtained by using a microhardness tester (Shimadzu HMV-2) with 50 g load (HV₀.₀₅). Following the application of background fitting, the average surface roughness of coatings (Ra) was determined using a surface profilometer (Ambios Technology XP-2, Santa Cruz, CA, USA). Furthermore, crystalline phase structures of the coatings and powder mixture were determined by using Rigaku Ultima 3 device (Rigaku, Tokyo, Japan) with Cu—Kα radiation (λ-Kα₁ = 1.54 Å), the device was set to 40 kV and 30 mA power, and scanning was done between 10° and 80° range with 2°/min scan rate and 0.02°/step. Crystallite sizes were calculated by using Sherrer’s equation [34].

\[
d = \frac{k \times \lambda}{\beta \times \cos \theta}
\]

where \(d\) is the crystallite size, \(\lambda\) is the X-ray wavelength, \(\beta\) is the width of the peak (full width at half maximum (FWHM)), \(\theta\) is the Bragg angle, and \(K\) is the Scherrer constant (\(K\) was taken as 0.94).

3. Results and Discussion

3.1. Macroscopic Examination

The coatings and powder produced on Inconel 625 surfaces by mechanical alloying were first examined macroscopically. Optical and stereographic images of the specimens can be seen in Figures 1 and 2, respectively. The grain sizes for each coating are large because of the amount of PCA used in the process. In order to enable powders to agglomerate and create a dense layer on the surface, PCA amount in the process was kept low in PCA-containing mixtures. For each coating parameter, the surfaces of samples show a continuous layer on the substrates. Because as-produced alloys have large grain sizes, the surfaces of coatings result in high roughness values as well. This macroscopical examination also shows that the resultant particles are larger and the surface structures are rougher for the materials processed with 10-mm grinding balls (the first, second, and third samples) than those processed with 5-mm grinding balls.
Figure 2. Macroscopic appearances of produced alloy powders and coatings.

Stereographic examination of the specimens shows further evidence of the surface structure of the coatings. Uniform coating consolidation was achieved by producing a dense layer on the substrate and having ductile particles performing plastic deformation. With the addition of PCA (left to right in Figure 3a), the surface grain structure becomes smaller. However, this addition also changes the geometrical shape of these surfaces. The irregular shape of particles makes the surface hard to cover. The coating process will fail if there are gaps, cracks, or other interfacial voids [35]. Moreover, one-dimensional powders (spheres) will be more controllable and will have higher accuracy in transforming the particles into the cold-welded coating [36]. Additionally, with the increasing PCA content, the surface morphology of coatings shows some crack-like separation on particle junctures (Figure 3b—coating 6).

3.2. Characterization of Alloy Particles

As the powder mixture starts to form an alloy, it either produces an agglomerate or forms a coating layer on substrates. Thus, the resultant particles and surface layer produced during the alloying and coating processes are very similar in their structure. Produced particles were examined on an optical microscope to clearly understand the formation process of the coating (Figure 4). The lamellar structure of the ductile Al constituent surrounds the brittle Si particles.

The process transformed the powder mixture into an alloy, and microstructures of particles showed a typical hypereutectic alloy structure [37,38]. At the same time, plastic flow marks arose on the produced alloy particles and can be seen in higher (500×) magnifications. Particles produced with 10-mm grinding balls are coarser than those with 5-mm balls. Naturally, the particles form alloys faster because of the higher energy input obtained from larger ball sizes; however, the resulting particles show some porosities (Figure 4—no 1). Coarser particles mean coarser coatings, hence rougher surfaces when coated, which is discussed in Section 3.3. While the alloy formation process continues, plastic deformation
occurs on ductile aluminum powders, forming the matrix phase. Brittle silicon powders fracture and decrease in size with increasing time. During MA, Suryanarayana indicated that [18], brittle constituents are encapsulated by ductile constituents. The brittle particle is closely spaced between the interlamellar spacings of elongated ductile structure. With the continuation of the alloying process, the ductile particles undergo work hardening and the lamellar structure is intertwined and the grain structure refines. With increasing MA time, the alloy powder particles get work hardened, the hardness and consequently the brittleness increases [18]. On the other hand, PCA content hinders ductile particles from agglomerating and acts as a surfactant to fracture particles by penetrating into microcracks during the alloying process [39]. However, in order to produce a homogeneous coating on a substrate, particles should cold-weld and remain agglomerated. The sixth sample with 0.5% PCA content shows a very fine grain structure, which is desired. However, this structure also contains very large cracks inside, which could cause problems as a coating.

Figure 3. Stereographic structures of coatings (a) (16× magnification), and (b) showing coatings number 4, and 6 (50× magnification).
The process transformed the powder mixture into an alloy, and microstructures of particles showed a typical hypereutectic alloy structure [37,38]. At the same time, plastic deformation caused the transformation of the structure into a more advanced mechanical property.

In Figure 5, hardness values for different parameters are given. Smaller grain structures show advanced mechanical properties with increasing plastic deformation. However, Figure 4 shows that the increase in the deformation with increasing time causes incompatibility.

**Figure 4.** Optical microstructures of 1st, 4th, 6th, and 9th specimens from top to bottom, with 50× (left), and 500× (right) magnifications.

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ble structures between the center and edges of particles on the ninth sample. The center of alloy particles was formed in the earlier stages of the mechanical milling process. With time, the edges of particles deform further and become harder via deformation hardening. This problem results in cracks and voids in the particles after a processing time longer than 6 h.

![HV1 hardness values for specimens numbered 1, 4, and 9; with different structures.](image)

**Figure 5.** HV$_1$ hardness values for specimens numbered 1, 4, and 9; with different structures.

### 3.3. Coating Characterization

Chemical composition obtained from EDS analysis of mechanically alloyed coating is shown in Figure 6. EDS no 1 shows the general coating composition and contains $\approx 17\%$ silicon content. The silicon amount is a little higher than the powder mixture ratio (15% silicon), just probably because the EDS results could not precisely give the exact amounts. The different phase structures on the image can be identified as the darker matrix phase of $\alpha$-Al matrix containing 10.6% of silicon (approximately eutectic composition, second EDS) and the lighter silicon particles (third EDS). As a result of the MA process, it was concluded that the desired alloy was formed.

Figure 7 shows the XRD phase development of produced alloy coatings with the starting powder mixture. Because of their nature, Al-Si hypereutectic alloys do not form an intermetallic compound after forming an alloy via mechanical alloying; $\alpha$-Al solid solution and Si diffraction peaks remain separate. Still, peak heights and widths changed after MA was applied via increasing silicon content in the $\alpha$-Al matrix. Moreover, increasing time and PCA amount, decreased the crystallinity of MA-alloyed coatings.
Figure 6. SEM image of mechanically alloyed coating and EDS results of marked areas.

Table 3. XRD results of mechanically alloyed coatings.

| Sample | Al (111) Intensity | Property | Powder Mixture | 1 (10 mm 6 h 0% PCA) | 4 (5 mm 6 h 0% PCA) | 6 (5 mm 6 h 0.5% PCA) | 8 (5 mm 18 h 0.5% PCA) |
|--------|--------------------|----------|----------------|----------------------|---------------------|----------------------|-----------------------|
| 1      | 23,804             | 2-θ Value (degrees) | 38.64           | 38.70                | 38.68               | 38.76               | 38.82                 |
| 2      | 1.2342             | d-spacing (Å) | 1.2326          | 1.2331                | 1.2310              | 1.2294              |
| 3      | 0.204              | FWHM (2θ) | 0.335           | 0.271                 | 0.376               | 0.337               |
| 4      | 43.98              | Crystallite size (nm) | 26.26          | 32.46                 | 23.40               | 25.38               |

While the initial powder mixture has very intense and narrow aluminum and silicon phase diffractions, the coatings clearly show wider and lower intensity peaks, which is reasonable with mechanical alloying. This change is due to the increase of the amorphousness of the structure [18,40]. Because the number of dislocations, grain boundaries, and...
other faults increases by cold working, the imperfections of crystals increase. Therefore, it
could be concluded that the amorphousness (the amount of disorder) increases too. Fur-
thermore, because the increase in the milling time results in higher deformation, d-spacings
(interplanar distances) of lattice planes shift to the right with an increase in 2θ degrees
(Table 3), which could be explained by the increase of the residual stress. Likewise, the
relation between the fourth and sixth samples shows that increasing stearic acid content
also decreases the crystallinity of the structure. It was observed that the calculated crys-
tallite sizes decreased with the increasing of ball size and PCA. However, no change was
observed in the crystallite dimensions depending on the increase in time.

Table 3. XRD results of mechanically alloyed coatings.

| Sample | Powder Mixture | 1 (10 mm 6 h 0% PCA) | 4 (5 mm 6 h 0% PCA) | 6 (5 mm 6 h 0.5% PCA) | 8 (5 mm 18 h 0.5% PCA) |
|--------|----------------|----------------------|---------------------|----------------------|-----------------------|
|        | Intensity      | 23,804               | 14,464              | 20,692               | 9087                  | 7282                  |
|        | 2-Theta (degrees) | 38.64               | 38.70               | 38.68               | 38.76                 | 38.82                 |
|        | d-spacing (Å)  | 1.2342              | 1.2326              | 1.2331              | 1.2310                | 1.2294                |
|        | FWHM (2θ)      | 0.204               | 0.335               | 0.271               | 0.376                 | 0.337                 |
|        | Crystallite size (nm) | 43.98           | 26.26               | 32.46               | 23.40                 | 25.38                 |

Cross-section SEM examinations (Figures 8 and 9) give some brief information about
the homogeneity and surface morphology of coating thicknesses. Figure 8a represents the
low magnification (33×) coating profile of the ninth specimen. The coatings are continuous
and dense, but rough, showing valleys and ridges. Coating profiles obtained from cross-
sections show that coatings produced with PCA content have higher roughness, with
nonhomogeneous coating thicknesses. On the contrary, coatings without PCA (first and
fourth samples) have a smooth surface.

In Figure 9, coatings prepared with different PCA contents show cracks and voids, as
seen in the third, sixth, seventh, and ninth specimens. Specimens that do not contain PCA
in the MA mixture (the first and fourth specimens) plastically deform and easily bind to
the surface. As mentioned earlier, the presence of ductile particles enables the powders to
merge and plastically deform. On the other hand, PCA acts as a barrier between ductile
particles, making them act as fragile particles [39]. This blockage leads to inferior cold
welding of particles through MA, letting undesired cracks and voids to generate between
them. A horizontal crack originates from the interface, because of the increasing energy
input with the process and increasing hardness of the coating. Plastic deformation and
mechanical interlocking of hard particles become more difficult, and therefore adhesion
properties deteriorates [41].

Another result obtained from these coatings is that 10-mm grinding balls (samples
to three) will have higher coating thicknesses. The larger the grinding ball gets, the
higher the energy produced during the mechanical milling process. The energy makes the
coating process propagate faster, and the layer becomes thicker. Average coating thickness
and Vickers hardness (HV1) values acquired from cross-sections, and the roughness (Ra)
values from surfaces are given in Table 4. As aforementioned, PCA addition increases
the roughness values of coatings. Moreover, with longer milling times, coating thickness
was raised together with surface roughness [42]. This change in surface roughness can be
explained by the effect of ball-substrate-ball collisions during MA, with metal powders
becoming cold welded to the substrate surface [42,43].
sections show that coatings produced with PCA content have higher roughness, with non-homogeneous coating thicknesses. On the contrary, coatings without PCA (first and fourth samples) have a smooth surface.

Figure 8. Low magnification (33×) cross-section coating profile of sample no 9 (a), and 150× profiles of samples no 1, 3, 4, 6, 7, and 9 for detailed comparison (b).
particles, making them act as fragile particles [39]. This blockage leads to inferior cold welding of particles through MA, letting undesired cracks and voids to generate between them. A horizontal crack originates from the interface, because of the increasing energy input with the process and increasing hardness of the coating. Plastic deformation and mechanical interlocking of hard particles become more difficult, and therefore adhesion properties deteriorate [41].

Figure 9. Cross-section SEM images of coatings number 1, 3, 4, 6, 7 and 9 at 1500 × magnification.

Table 4. Average thickness, roughness, and hardness values of coatings.

| Specimen No | Average Thickness (µm) | Roughness (Ra, µm) | Vickers Hardness (HV1) |
|-------------|------------------------|--------------------|-----------------------|
| Substrate   | -                      | -                  | 273 ± 6.7             |
| 1           | 160.5 ± 22.4           | 19.639             | 91.4 ± 17.4           |
| 2           | 190.4 ± 54.1           | 21.095             | 126 ± 21.9            |
| 3           | 256.7 ± 107.1          | 23.881             | 142 ± 24.3            |
| 4           | 122.3 ± 11.4           | 14.264             | 78.8 ± 12.3           |
| 5           | 126.4 ± 27.6           | 19.571             | 107 ± 18.8            |
| 6           | 128.9 ± 46.3           | 22.314             | 134 ± 24.2            |
| 7           | 176.9 ± 43.5           | 22.792             | 160 ± 21.3            |
| 8           | 182.6 ± 73.7           | 23.311             | 166 ± 32.7            |
| 9           | 164.8 ± 42.1           | 22.405             | 178 ± 41.3            |

Micro Vickers hardness (HV0.05) profiles from the cross-sections of coatings are shown in Figure 10. It could be simply understood that materials are harder on the surfaces of coatings, as described earlier on microstructure analysis. The surface of the mechanically alloyed particles gets harder with the continuous impact of the grinding balls. Inner parts of the coatings will be exposed to this impact energy to a lesser degree till the substrate-coating interface. When the profiles are analyzed, the softest coating is the fourth sample, while the ninth sample has the hardest surface. The fourth sample remains the softest on the interface,
but the third sample has become the hardest. The amount of reduction on the hardness profiles of coatings using balls with 10-mm diameter (the first and third specimens) is smaller than coatings produced with 5-mm balls. Higher impact and sliding energy input of larger grinding balls result in a more homogeneous distribution of structure (Figure 4), and therefore hardness profiles of coatings become more stable toward the interface.

Figure 10. Cross-section microhardness profiles of coatings from surface to interface.

4. Conclusions

Mechanically alloyed Al-15Si coatings were successfully applied on the Inconel 625 superalloy substrates using a planetary ball mill. Furthermore, the effects of important process parameters; the diameter of the grinding ball, the milling time, and PCA content were investigated using several characterization techniques. Based on the findings of this study, the following conclusions were drawn:

- Increasing the grinding ball diameter from 5 to 10 mm increases the coating thickness, and provides a homogenized microstructure by introducing a higher energy input on the material’s surface. However, the increased coating thickness in these parameters leads to increased surface roughness values too. Because of the increase, surface roughness values also increase. These values might produce a potential problem on the tribological aspect of potential high temperature, high-speed components.

- Although coating thickness values and mechanical properties improve with the rising milling time, the surface of the coating becomes incompatible with the center or the coating-substrate interface.

- The use of PCA causes cracks and voids in the coatings but enhances the mechanical properties by decreasing grain size. Highest hardness value of 178 ± 41.3 HV1 was attained by using highest amount of PCA (1% stearic acid), and lowest value of 78.8 ± 41.3 HV1 was obtained without using PCA. Therefore, PCA content should be kept at optimal levels to hinder crack formation and for gaining higher hardness values, and milling time needs to be increased when PCA is used.
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