

MILLING PARAMETER OPTIMIZATION FOR THE REFINEMENT OF NiO/Al MIXTURES AND THE SYNTHESIS OF Ni/Al₂O₃ NANOCOMPOSITES

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Abstract

This study examined the effects of milling parameters on the development of Ni/Al₂O₃ nanocomposites and the refinement of NiO and Al powders. Ball milling of certain mixtures was followed by sintering at 800 and 1100 °C for 2 h. The X-ray diffraction results of the dry-milled powders indicated that increasing the ball-to-powder weight ratio from 20:1 to 42:1 resulted in finer particles, which enabled the synthesis of Ni/Al₂O₃ nanocomposites by milling at 200 rpm for 1.5 h. Extending the milling duration at lower rotational speeds yielded powders with nanoscale particle sizes. However, as shown by scanning electron microscopy and energy dispersion spectroscopy, a nanocomposite with metallic matrix was formed by the mechanochemical reaction, and the crystallite size was estimated using the Williamson–Hall plot. Furthermore, we used differential scanning calorimetry diagrams to analyze the effects of milling on the temperatures of phase transformation and/or reduction reactions. The tribological performance of the developed nickel metal matrix composite was investigated using a ball-on-disc tribometer under various loading conditions. Indeed, the friction coefficient increases with the applied forces and decreases with milling. Comprehensive examinations of the worn surfaces were carried out using a scanning electron microscope and a 3D optical profiler.

Keywords: Mechanical alloying; Ni/Al₂O₃ nanocomposite; Sintering; Aluminothermic reduction; Friction coefficient; Wear mechanism

1. Introduction

The mechanical alloying (MA) technique is widely employed to synthesise new nanocomposite materials for a variety of applications. Mechanochemical reactions are defined as chemical reactions (exchange reaction, reduction/oxidation, decomposition of compounds, or phase transformation) occurring during the milling process. Among these reactions, the aluminothermic reaction (thermite reaction) occurs between a metal oxide (NiO, CuO, ZnO, TiO₂, Fe₃O₄, ZrO₂, etc.) and aluminium [1–8]. However, two distinct reaction kinetics can emerge depending on the applied milling parameters: first, a gradual transformation; second, self-propagating high-temperature synthesis (SHS) [9] or combustion synthesis. In principle, the SHS requires a critical period before the ignition is

triggered inside the vials during milling. After the gradual increase in the temperature of the vials in the first stage of milling, the temperature increases rapidly indicating the ignition of the reaction [10]. In recent years, several researchers have employed various methods to utilize the NiO/Al system for developing Ni matrix composites incorporating Al₂O₃ particles, which are promising options for applications requiring enhanced wear and corrosion resistance [11–12]. However, Vrel et al. [13,14] investigated the reaction between NiO and Al via SHS by adding various concentrations of alumina and adjusting the stoichiometric ratio of the reagents to minimize the amount of heat released during the thermite reaction, which successfully minimized the sputtering and destruction of the sample. Troncy et al. [15,16] employed pre-oxidised Ni and Al to fabricate self-regenerating coatings of NiAl/Al₂O₃ via heat

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treatment. In addition, high-energy ball milling has been applied to create similar composite materials composed of Al, NiO, and Ni particles. Previous research reported that a critical period of 110 min is required before the ignition of the mixture inside the vials. After 10 h of milling, only the product phases were detected, and upon extending the milling duration to 60 h, refined grain sizes of NiAl and Al₂O₃, i.e., 8-nm particles, were observed [17]. Moreover, NiO/Al powders were used with other materials to prepare composites with various reinforcements [18–21]. Matteazzi et al [22] employed two types of mill, specifically a vibratory ball-mill and a planetary ball mill, in their study of various M_xO_y and Al systems. The α-Al₂O₃-Ni composite was successfully synthesized within a milling time of 1.5 hours, using a ball-to-powder ratio (BPR) of 42:1. Nevertheless, the literature available on this system is relatively sparse and provides only limited information [22,27]. Li [23] and Udhayabanu et al. [24] used a ball-to-powder weight ratio (BPR) of 10:1 along with various process control agents to develop Ni/Al₂O₃ nanocomposites from NiO and Al powders, obtaining Ni and amorphous Al₂O₃ after a long milling time of 10 and 5 h, respectively. They concluded that mechanical activation poses a gradual effect on the reduction of NiO by Al. Mameri et al [25] studied the synthesis of Ni/Al₂O₃ composite by powder a metallurgy process (milling and pressure-assisted sintering) and also investigated the tribological characteristics of these composites as part of their study.

According to the literature review, all existing research focusing on the NiO/Al system applied mechanical alloying with various milling. However, no study has attempted to optimize the milling conditions, whether to refine only the particle size of the reactants or to produce Ni/Al₂O₃ nanocomposites. Furthermore, this gap extends to the realm of tribology as well. There has been no investigation into the tribological behavior of Ni/Al₂O₃ composites obtained through the conventional powder metallurgy process, which involves milling, compacting, and sintering. Therefore, the fundamental objective of this study is to display the effect of various milling parameters, e.g., BPR, milling duration, and milling speed on the refinement of NiO/Al powders. Moreover, we attempted to achieve the smallest ignition period required to produce Ni/Al₂O₃ nanocomposite without using any reducing or protective agent under air. We performed sintering tests to validate the results obtained under the adopted milling conditions. Furthermore, we delved into the tribological attributes of the obtained composites and analysed how milling and applied loads affect the friction coefficient and the characteristics of the worn surfaces.

2. Experimental

Aluminium (Al) powder and nickel oxide (NiO) with an average particle size of 74 μm and <44 μm were purchased from Sigma-Aldrich and Acros-Organics, respectively. The powders were weighed with a stoichiometric composition corresponding to the following equation:



The mechanical alloying of the powders was conducted using a planetary micromilling Pluverisette 7 premium line FRITISH. Stainless steel vials with a capacity of 80 mL and balls of diameter Ø = 10 mm were used. Two ball-to-powder weight ratio (BPR) of 20:1 and 42:1, were used with a powder mixture of about 2g and 1g, respectively. Notably, the mechanical alloying was conducted under air and without adding a process-controlling agent (PCA). The milling conditions applied in this study are summarised in the following table.

Table 1. The milling conditions applied to the mixtures of NiO/Al powders

| case | BPR | Speed (rpm) | Time (h) |
|------|-------|-----------------------|-------------------|
| A | 20:01 | 100, 150, 200 and 250 | 1 |
| B | 42 :1 | 100, 150, 200 and 250 | 1 |
| C | 42 :1 | 100 | 1, 2, 3, 4, 5, 8. |
| D | 42 :1 | 100, 150 and 200 | 1.5 |

Prior to initiation, the mill was set up to halt for 15 min after every 30 min of operation; 2 g of the milled powders were uniaxially pressed at room temperature via a cylindrical tungsten carbide die (D = 13 mm) under a pressure of 510 MPa. Subsequently, the compacted samples were subjected to sintering using a tubular electric furnace in a neutral atmosphere provided by a vacuum pump at 800 and 1100 °C. In addition, differential scanning calorimetry (DSC) was performed for the green mixture, and an amount of the milled powders was heated at a rate of 25 °C/min in an argon atmosphere using a Netzsch STA 449 F3 Jupiter® to determine the thermal stability of the various samples. The particle morphology evolution of the milled powders and sintered samples was analysed using a Philips XL30 and FEI Quanta-250 scanning electron microscopes (SEM) equipped with an energy dispersive X-ray spectrometer (EDS). The structural evolution and phase formation occurring during mechanical alloying was determined using high-resolution powder X-ray diffraction (HRPXRD) performed at the ID 31 beamline (Ei = 31 KeV, corresponding to λ = 0.4 Å) of the European Synchrotron Radiation Facility (ESRF) in Grenoble and X-ray diffraction (XRD, X-Pert Panalytical)



using Cu-K α radiation for the sintered samples. Furthermore, the Williamson–Hall plot (W–H) method was used to calculate the crystallite size and lattice strain of the milled samples [26]. The tribological characteristics were evaluated by employing a ball-on-disc tribometer (CSM tribometer, manufactured by CSM Instruments Inc., located in Peseux, Switzerland) with alumina balls (diameter=6mm) as a counter-face material. The measurement of the volume loss on the worn surface was acquired through the utilization of CT 100 a 3D optical profilometer.

3. Results and Discussion

3.1. Milling process

3.1.1. X-Ray Diffraction

Effect of the ball to powder weight ratio (BPR)

The XRD plots of the specimens milled for 1 h as a function of milling speed (100, 150, 200, and 250 rpm) with a BPR of 20:1 representing Case A and a BPR of 42:1, representing Case B in Table 1 are depicted in Figures 1 and 2. As observed, the as-received powder mixture was characterized by sharp intense peaks, reflecting the higher crystallinity of NiO and Al powders. However, as the milling speed increased up to 200 rpm, the NiO and Al peak intensities notably diminished with large broadening of the peaks, which was potentially caused by the gradual reduction in the crystallite size of the reactants coupled with the increased lattice strain during milling. Furthermore, milling with a rotation speed of 250 rpm generated Ni and Al₂O₃ particles, indicating that the aluminothermic reaction between NiO and Al occurred during the mechanical alloying process (Eq. 1) by a self-propagating reaction as noted

earlier [23]. However, as observed from Figures 1 and 2, the peaks ascribed to the NiO phase are still present even when the thermite reaction has taken place, suggesting oxidation of a small amount of nickel after reaction and during cooling, due to the processing conditions in air. Notably, as the BPR increased to 42:1 the peak attributed to the nickel particles broadened, indicating that alumina became sharper. However, the peaks became broader than those of Ni and Al₂O₃ obtained in the first case with a BPR of 20:1.

The crystallite size of the green mixture particles was 491 nm for NiO and 810 nm for Al. As such, milling at a speed of 200 rpm and BPR of 20:1 produced NiO and Al particles with diameters of 79.7 and 325 nm, respectively. However, the grain sizes of the NiO and Al particles obtained with a BPR of 42:1 were 64.5 and 212 nm, respectively. Moreover, the grain sizes of Ni and Al₂O₃ of case B were 87 and 78 nm with a lattice strain of 0.171 and 0.241%, respectively (Table 2). As the particle size of the reinforcement (Al₂O₃) and metallic matrix (Ni) were less than 100 nm, the results confirmed the formation of a nanocomposites [27]. Milling cannot reduce Al particles to an extremely fine crystallite size owing to its ductility [17]. As fresh surfaces were generated by the reduction of the crystallite size via repeated cold welding and fracturing, the reactions that require elevated temperatures were rendered feasible at low temperatures [28]. During the phase transformation, the temperature of the vials increased, which can be attributed to the highly exothermic reaction between NiO and Al powder, exceeding 3156 K [29-30]. Prior to milling, the mixed powders were pale green in color, which subsequently varied to two distinct colours: first, a dark-green color for samples with no thermite reaction during mechanical alloying (100,

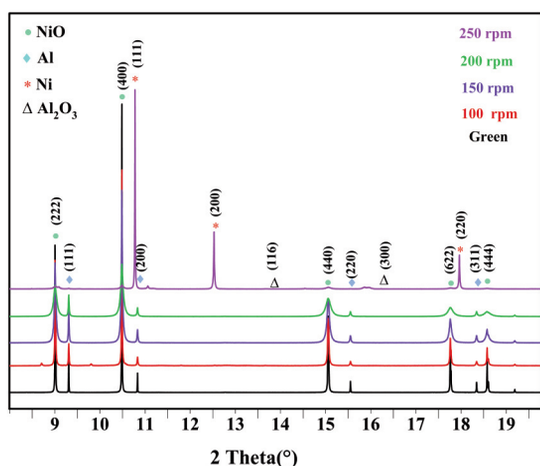


Figure 1. XRD plots of pure and milled NiO/Al powder mixture with a BPR of 20:1 milled for 1 hour as a function of milling speed 100, 150, 200 and 250 rpm

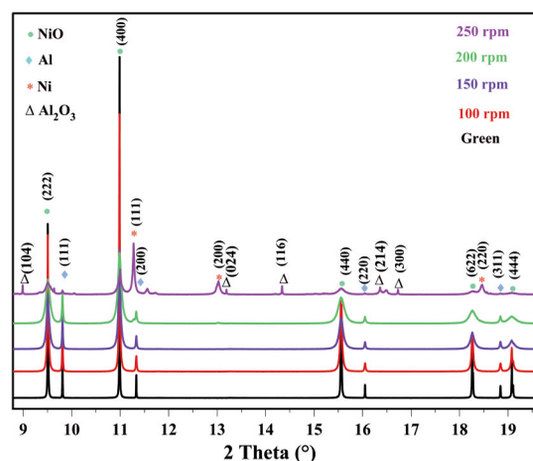


Figure 2. XRD plots of specimens milled for 1 hour as a function of milling speed (100, 150, 200 and 250 rpm) with a BPR of 42:1

Table 2. Mean crystallite size of NiO, Al, Ni and Al₂O₃ after milling for 1h as a function of milling speed for a BPR of 20:1 and 42:1

| BPR | Speed (rpm) | D NiO (nm) | D Al (nm) |
|------|-------------|------------------------------|--|
| 20:1 | 100 | 497 | 800 |
| | 150 | 113 | 366 |
| | 200 | 79.7 | 325 |
| | 250 | D _{Ni} (nm) 564 | D _{Al₂O₃} (nm) 144 |
| 42:1 | 100 | 206 | 759 |
| | 150 | 83 | 316 |
| | 200 | 64.5 | 212 |
| | 250 | D _{Ni} (nm) 87.8 | D _{Al₂O₃} (nm) 78 |

150, and 200 rpm); second, a black color related to the reduction of the NiO by Al into a Ni/Al₂O₃ nanocomposite (250 rpm), which are consistent with the previously reported observations [31].

Effect of milling duration

Based on the first two cases, we conclude that increasing the BPR to 42:1 yields finer particles. Therefore, pure and milled NiO/Al powders blended for various milling periods (1, 2, 3, 4, 5, and 8 h) with a BPR of 42:1 and rotation speed of 100 rpm is visualized in Figure 3, which corresponds to Case C in Table 1. As observed, no additional peaks were detected corresponding to other new or intermediate phases, and both NiO and Al peaks continued to exist. However, milling up to 8 h with a small rotation speed did not alter the composition of the product. The broadness of the NiO and Al peaks increased

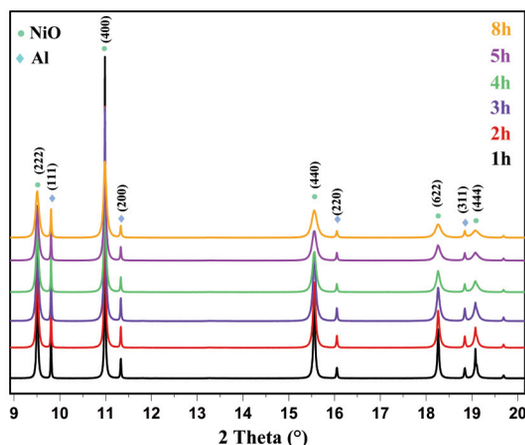


Figure 3. XRD plots of pure and milled NiO/Al powders blended for different periods of milling time 1, 2, 3, 4, 5 and 8h with a BPR of 42:1 and a rotation speed of 100 rpm

significantly with the milling duration. Moreover, the prolonged milling duration aided the development of homogenous powders at a nanometre scale, which can be attributed to the equilibrium achieved between the mechanisms of fracturing and cold welding [1]. Furthermore, the ball milling process can impact the crystallite size by inducing dislocations during the formation of sub-grains [32].

The crystallite size and lattice strain evolution are presented in Figure 4 as a function of milling period determined using the W–H plot method. According to the data acquired from the NiO peaks, the mean particle size after milling for 1 h was observed to be approximately 206 nm. Upon milling for 3 h, the crystallite size decreased considerably to 104 nm, and after milling for 8 h, the particle size further decreased to 76.2 nm. However, as per the W-H relation, the lattice strain should be inversely proportional to the crystallite size, which is not exactly the case in our analysis. It is proportional up to 3 h, after that the crystallite size is stabilized while the lattice strain is increased. A possible explanation for this phenomenon is the increase of stacking fault energy during milling, which impede the formation of new crystallite and increase the strains due to impacts during milling [33,34].

Effect of milling speed

The XRD patterns of pure and milled powders for 1.5 h using a BPR of 42:1 for varying milling speeds of 100, 150, and 200 rpm are illustrated in Figure 5. In Case D, the Ni/Al₂O₃ nanocomposite was formed only upon applying a milling speed of 200 rpm. This result is consistent with the literature on the same system [22,27]. Milling with a speed of 100 and 150 rpm only activated the powder mixture and refined the reactant particles. The mechanical activation of the powders during milling can be related to the repeated collision applied to the particles between the balls and vial walls or between the balls [35]. The objective of

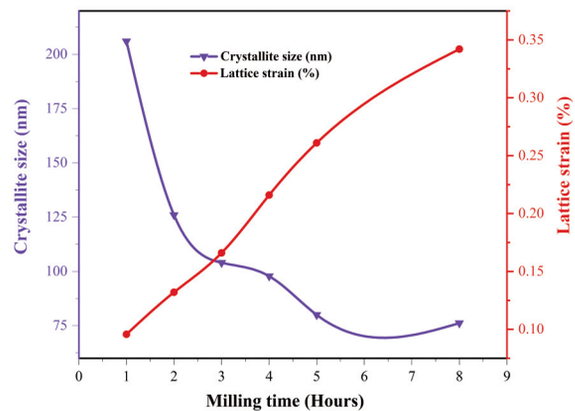


Figure 4. The crystallite size and lattice strain evolution as a function of milling time

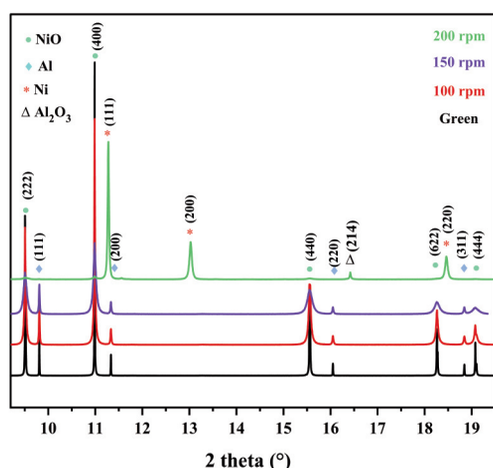


Figure 5. XRD patterns of pure and milled powders during 1.5 hour using a BPR of 42:1 for different milling speed 100, 150 and 200 rpm

searching for optimal conditions to enhance the refinement of NiO/Al powders is to generate mechanically activated and finely refined powders. The milling time is a key factor in the milling process and is closely related to the Ball-to-Powder Ratio (BPR) [27]. A larger BPR results in higher milling intensity and a shorter time required for reaction completion, and vice versa. The selection of BPR in this study was determined based on relevant literature [22,27]. Prolonged milling time increases the risk of contamination and the formation of undesirable phases [27]. It also necessitates the use of a diluent [22], and excessively high speeds can strongly accelerate the reaction without achieving the desired refinement of the co-milled powders. These refined powders will then be introduced into the Ti-C system to assess their impact on the ensuing TiC synthesis reaction, as part of an upcoming study. Notably, in all Cases (A, B, C, and D), the intensity of the peaks ascribed to Al was substantially less than that of NiO owing to the low volume fraction of Al in the primary mixture (80.6 wt.% of NiO with 19.4 wt.% of Al). The

disappearance of the Al_2O_3 peaks in this sample was possibly caused by the low volume fraction of the new phase formed during milling, which is related to the extremely fine particles obtained. Moreover, as NiO is extremely brittle in nature, it reduces to ultrafine sizes after milling. The crystallite sizes of the Ni and Al_2O_3 phases determined using the W–H plot method were 85 and 45 nm, respectively. As reported by Suryanarayana, the evolution of the grain size depends on increasing the milling energy and BPR at low temperatures [1].

3.1.2. SEM Microstructure

The SEM micrographs of the unmilled and milled samples for 1 h at 200 rpm are depicted in Figure 6 for BPR 20:1 and 42:1. As observed, the unmilled powder mixture exhibited a particle morphology composed of irregular particles of Al surrounded by smaller grains of NiO. The size of the Al and NiO particles diminished significantly upon milling. The inhomogeneous distribution of these particles produced an irregular morphology. Moreover, the fine particles were agglomerated by the repeated fragmentation and cold welding of the powders due to the absence of a process control agent (PCA). The PCA in the mixture ensured that the particle surfaces do not face any barrier between themselves or against the vials [1]. Moreover, the SEM images of the samples displaying the presence of Ni and Al_2O_3 phases in their XRD spectra because of the mechanochemical reaction at the end of the milling process are presented in Figure 7. Two phases, bright and dark, corresponding to the Ni and Al_2O_3 phases, respectively, were observed with small irregular particles and a few agglomerates, but with finer product particles. Thereafter, a specimen milled with a BPR of 42:1 and 250 rpm for 1 h (Figure 7(d)) revealed another morphology. The refinement of the product grains was evident, indicating the predominant fracture in the cold-welding process

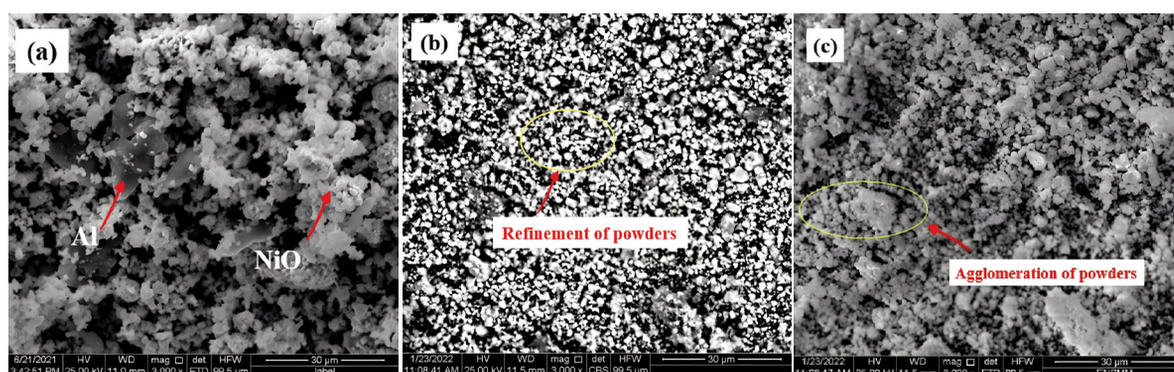


Figure 6. SEM micrographs of unmilled and milled samples with a rotation speed of 200 rpm for 1 hour (a) unmilled (b) BPR 20:1 (c) BPR 42:1



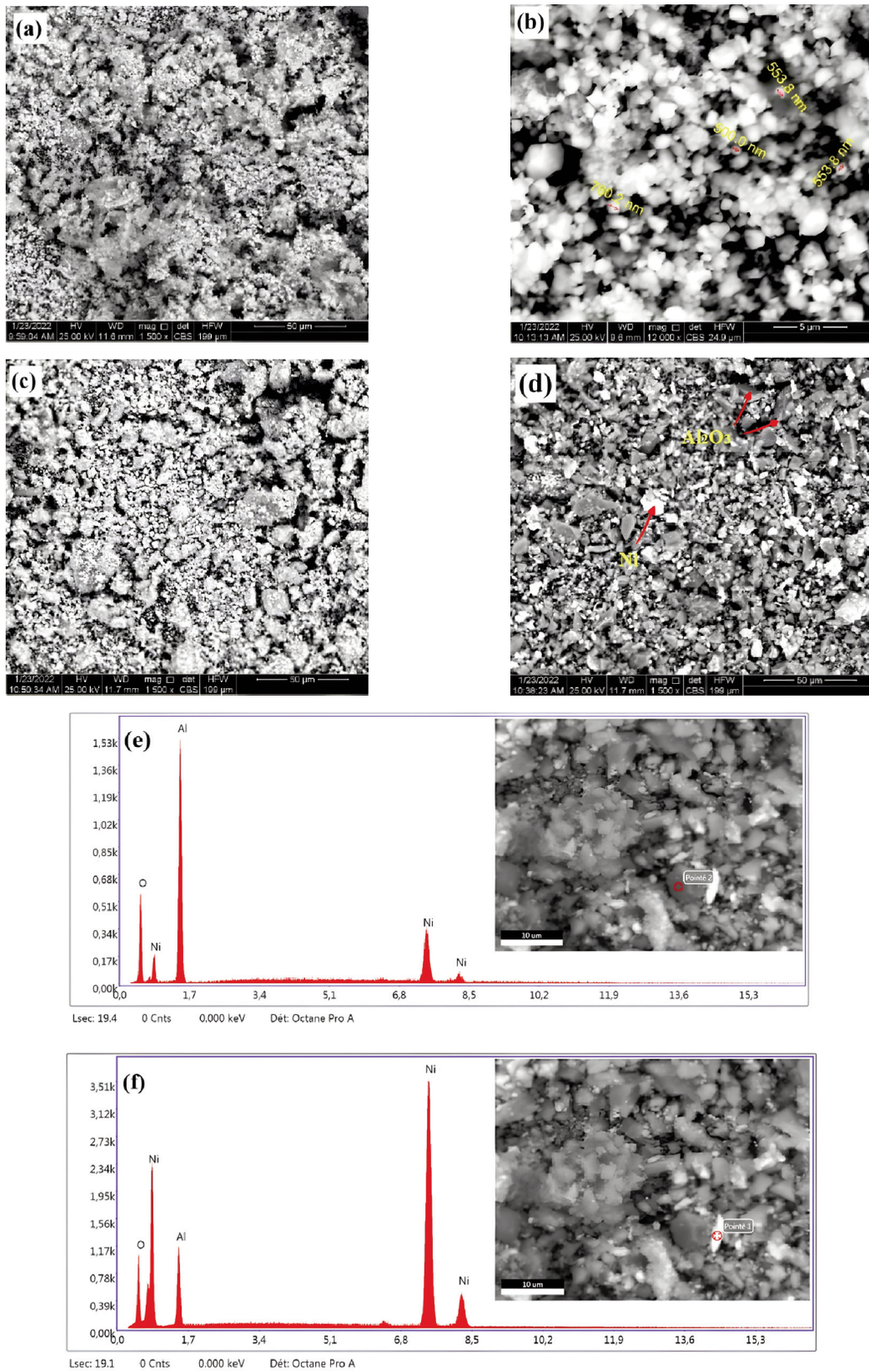


Figure 7. SEM micrographs and EDS plots of samples (a, b) milled with BPR of 20:1 and 250 rpm for 1 hour, (c) BPR of 42:1 and 200 rpm for 1.5 hour; (d) BPR of 42:1 and 250 rpm for 1 hour; (e, f) EDS of sample d

following the reduction reaction. This result is consistent with previous reports on similar systems [36,37] as confirmed by the EDS analyses (Figure 7(e-f)). For a BPR of 20:1 and speed of 250 rpm for 1 h (Figure 7(a)), the average size of the Ni particles ranged from 500 to 700 nm (Figure 7(b)), which confirms the previous reports of Ni particle size calculated using the W-H plot method. Moreover, the same observations are recorded in Figure 7(c).

3.1.3. Differential Scanning Calorimetry analysis DSC

To further explain and understand the kinetics of the thermite reaction, a range of samples that correlated with the previous results of the XRD analysis were treated with DSC. The DSC profile of the unmilled NiO/Al powder mixture heated at a rate of $25\text{ }^{\circ}\text{C min}^{-1}$ is plotted in Figure 8. The first endothermic peak was observed at $665\text{ }^{\circ}\text{C}$, corresponding to the melting temperature of Al, whereas the exothermic peak detected at $1015\text{ }^{\circ}\text{C}$ represents the NiO reduction according to Equation (1). This result confirms that the reaction between NiO and Al occurs at a high temperature after the melting of Al, which is consistent with the findings of Udhayabanu [24] and Padhan [31]. The DSC curves of the mechanically activated samples were identical to those of the unmilled sample, varying only in terms of the onset temperature. The sample milled with a BPR of 20:1 and rotation speed of 250 rpm for 1 h is similar in shape and nature to the sample milled with a BPR of 42:1 and speed of 200 rpm for 1.5 h. However, the two exothermic peaks observed at 454 and $591\text{ }^{\circ}\text{C}$ are likely to be associated with the partial reduction of NiO. Notably, a small endothermic peak was observed at $636\text{ }^{\circ}\text{C}$, which was certainly related to the melting temperature of Al. The ultimate exothermic peak observed at $910\text{ }^{\circ}\text{C}$ was related to the exothermic reaction between the unreacted NiO and melted Al, which resulted in the formation of the Ni/Al₂O₃ nanocomposite. The DSC curve of the sample milled with a BPR of 42:1 at 250 rpm for 1 h exhibited a distinct pattern, displaying a faint exothermic peak at $982\text{ }^{\circ}\text{C}$ that represents the complete reduction of NiO. The absence of the Al endothermic peak indicated that the entire Al was oxidized in the mixture. More importantly, the benefit of mechanical alloying on powders where a thermite reaction can occur results in a lower ignition temperature for the combustion synthesis [4]. No endothermic peak corresponding to the melting of NiO was observed in any of the DSC curves due to its exceptionally high melting temperature of $1984\text{ }^{\circ}\text{C}$, which is above the temperature range covered by the DSC analysis. These results are consistent with the XRD patterns of the sintered samples.

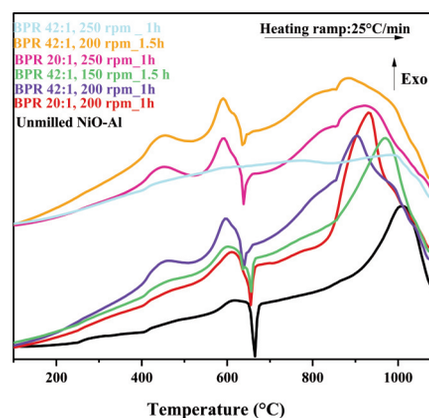


Figure 8. DSC plots comparing the unmilled NiO/Al mixture to the milled samples

3.2. Sintering process

3.2.1. XRD analysis

The sintering process was applied to samples, wherein the final product was not formed using MA. Depending on the starting conditions applied to the NiO/Al system, the reduction process might occur during milling or sintering [39]. The milled samples sintered at $800\text{ }^{\circ}\text{C}$ for 2 h were analyzed using XRD, and the results are depicted in Figure 9. As observed, new phases of Ni and Al₂O₃ were formed during sintering, and the NiO peaks remained. This can be attributed to the low sintering temperature that is insufficient to ensure a complete reduction ($800\text{ }^{\circ}\text{C} < 904$ and $930\text{ }^{\circ}\text{C}$) according to the present DSC results. Moreover, the quantity of NiO present in the pellet was not completely reduced. However, we observed a marginal reduction in the intensity of the peaks as the peak corresponding to the sample milled with a BPR of 42:1 broadened wider than that milled with a BPR of 20:1. The XRD profile of the milled sample sintered at $1100\text{ }^{\circ}\text{C}$ for 2 h is shown in Figure 10. Moreover, the sharp intense peaks were recorded for both Ni and Al₂O₃ phases; the presence of NiO in the mixture after sintering likely resulted from the oxygen present within the tube during the sintering process. Additionally, it may be associated with the oxidation of nickel during the extraction of the mixture following the milling process. As the analytical section presided over molten Al₂O₃, alumina peaks were more frequent than nickel peaks. A similar tendency was observed in the X-ray study performed by Jones et al. [40].

3.2.2. SEM microstructure

The morphology of the new product phases in the samples sintered at $800\text{ }^{\circ}\text{C}$ for 2 h is depicted in Figure 11. As observed in Figure 11(a) and (c), the product particles were arranged in a homogenous

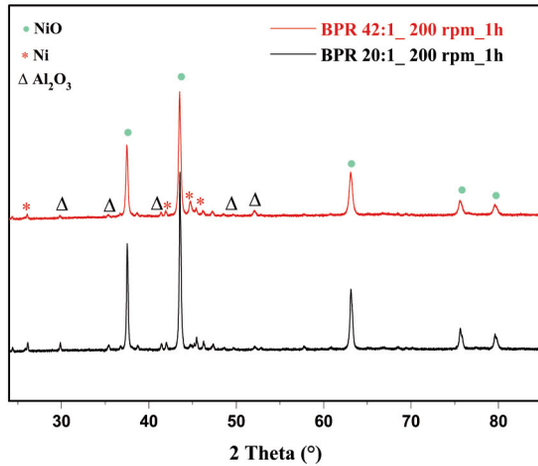


Figure 9. XRD patterns of milled and sintered samples at 800°C for 2 hours

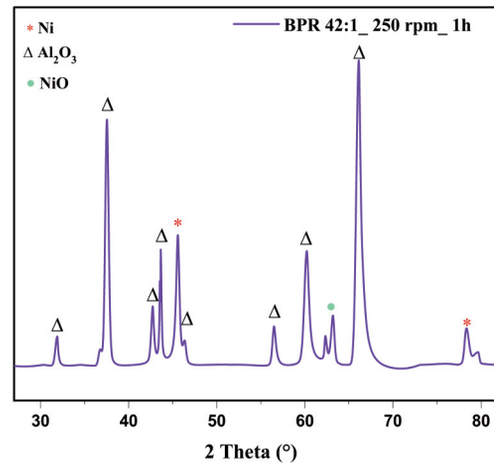


Figure 10. XRD profile of milled and sintered sample at 1100°C for 2 hours

distribution with no agglomeration. The number of voids in the composite is reflected in the dark zones; dark grey particles denote Al_2O_3 , whereas white particles represent the Ni phase. The EDS analysis of the two sintered samples revealed no contamination during milling and/or sintering; only the reactants and product elements were detected. The composition of the area in Figure 11(b) was nearly identical to that in (d). Despite selecting an extremely small area of analysis, all the selected areas did not exhibit a single phase, depicting fine grains and particles of the composite. In the sintering process, the thermal

conditions applied to compacted powder particles promote grain growth, allowing them to connect to each other across of grain boundaries. Increasing the sintering temperature promotes increased grain growth, resulting in a reduction of the trapped air between particles and subsequently enhancing the density of the samples [33]. The microstructure of the specimen milled with a BPR of 42:1, speed of 250 rpm for 1 h, and sintered at 1100 °C for 2 h is depicted in Figure 12. After sintering, the recovered sample was melted in a ceramic container, where it nearly separated into two parts—nickel and alumina. The

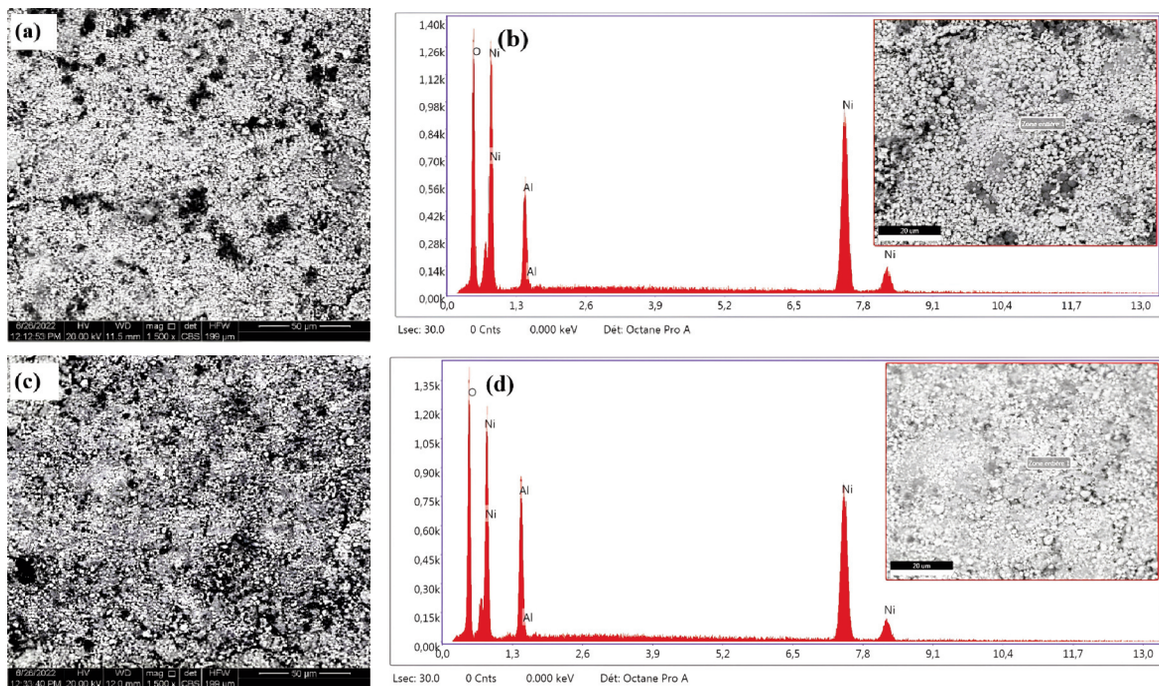


Figure 11. SEM images and EDS plots of sintered samples at 800°C for 2 hours of (a, b) BPR 20:1 and 200 rpm for 1 hour; (c, d) BPR 42:1 and 200 rpm for 1 hour

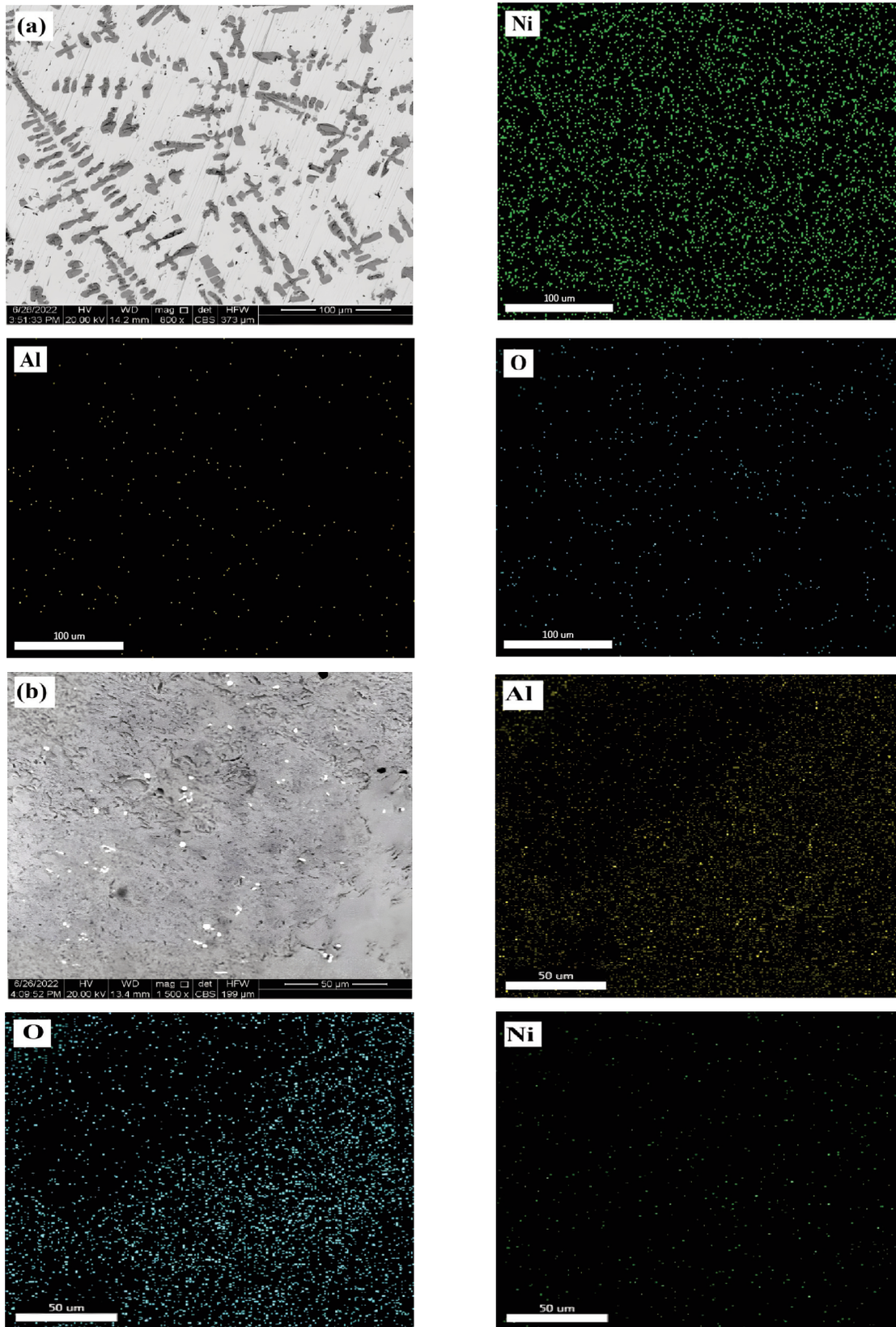


Figure 12. SEM images and EDS mapping of sintered sample at 1100°C for 2 hours (a) Nickel phase and (b) Alumina phase

mapping analysis in the Ni phase (Figure 12(a)) exhibited the presence of oxygen and Al, but with low concentrations. Similar observations were recorded for mapping the alumina phase, and the amount of nickel was considerably small (Figure 12(b)).

3.3. Tribological Properties

3.3.1. Effect of milling and the applied load on the friction coefficient

Milling parameters have a significant and consequent influence on both the microstructural properties of the acquired powder mixtures and the resulting sintered composites, as a result, they have a significant influence on their respective tribological properties. Two samples were examined: (a) an unmilled sample sintered at 1100 °C and (b) a sample milled with a BPR of 42:1 at 200 rpm for 1 hour and subsequently sintered at 1100 °C. The tests were conducted under standard environmental conditions, subjecting the specimens to vertical forces of 5 N and 10 N, with a sliding distance of 50 m and a sliding speed of 25 mm/s. Figure 13 depicts the development of the friction coefficient for both samples. It is noticeable that as the applied load increases, the friction coefficient increases in both samples. Increasing the applied load from 5 to 10 N increases the friction coefficient of the unmilled sample from 0.48 to 0.66, respectively. Moreover, the friction coefficients of milled sample reached 0.44 and 0.60 when applying loads of 5 and 10 N, respectively. An increase in the friction coefficient may occur as a result of the removal of surface asperities, which leads to the formation of a third body. These particles from the third body, located at the contact surface, contribute to elevated value of friction [40-42]. It is noteworthy that the coefficient of friction reduces with milling. These findings show the impact of milling on the friction coefficient. The decreased

coefficient of friction observed in milled sample can be attributed to the occurrence of work hardening resulting from the repeated fracture and welding caused by milling. As a result, the Young's modulus increased, leading to a reduction in the friction coefficient [43].

3.3.2. Wear mechanism

Figure 14 reveals the worn surface of the two sintered specimens (a) unmilled (10N) and (b) milled with a BPR of 42:1 at 200 rpm for 1 hour (5 and 10 N). As observed, Figure 14 (a) the worn surface displays clear signs of abrasive wear, featuring the emergence of several microcracks. Due to the existence of microcracks, particles can be readily dislodged from the matrix, resulting in the creation of a rough wear track. The primary factor contributing to the significant weight loss is the detachment of particles. As depicted in Figure 14b (SEM), the wear tracks on the specimens subjected to a 10 N load are characterized by their depth and narrowness, in contrast to those exposed to 5 N loads. A minimal quantity of debris was detected around the wear track. Moreover, the worn surfaces of sample (b) reveals the presence of plastic deformation and more cracks. An extensive examination of changes in the 3D surface morphology of both sintered samples with an applied load of 10N was provided in figures 14 (c and d). The mean wear track depth measured -18 μm for milled and sintered specimens compared to -5 μm for the unmilled samples. The irregularities observed at the base of the wear tracks, particularly as shown in Figure 14(d), suggest that abrasive wear is also a common occurrence during contact. This is likely attributed to three-body abrasion mechanisms resulting from the fragmentation of oxide particles and wear debris [44]. The findings above suggest that the tribological characteristics are influenced by

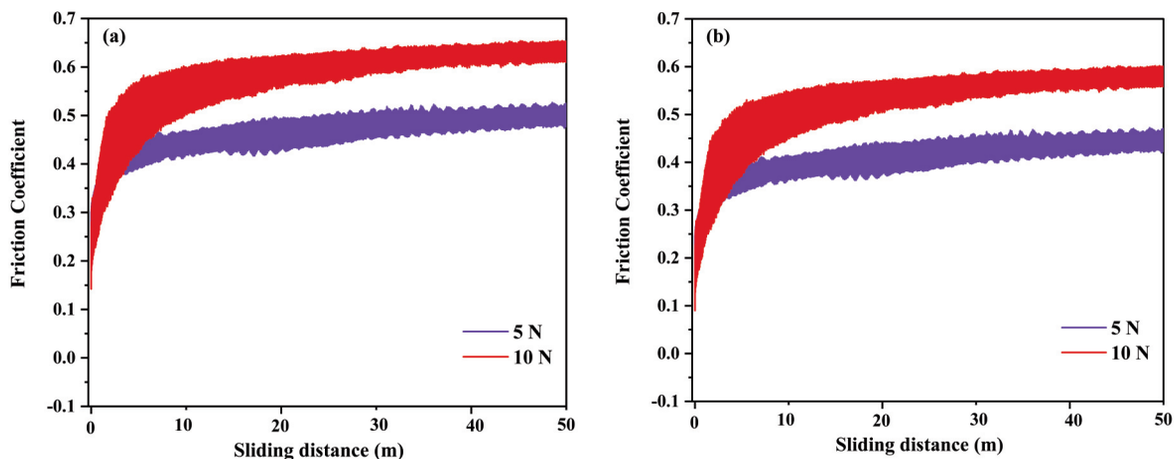


Figure 13. Impact of applied forces (5 and 10 N) on the friction coefficient of sintered specimens, (a) unmilled sample and (b) a sample milled with a BPR of 42:1 at 200 rpm for 1 hour and subsequently sintered at 1100 °C

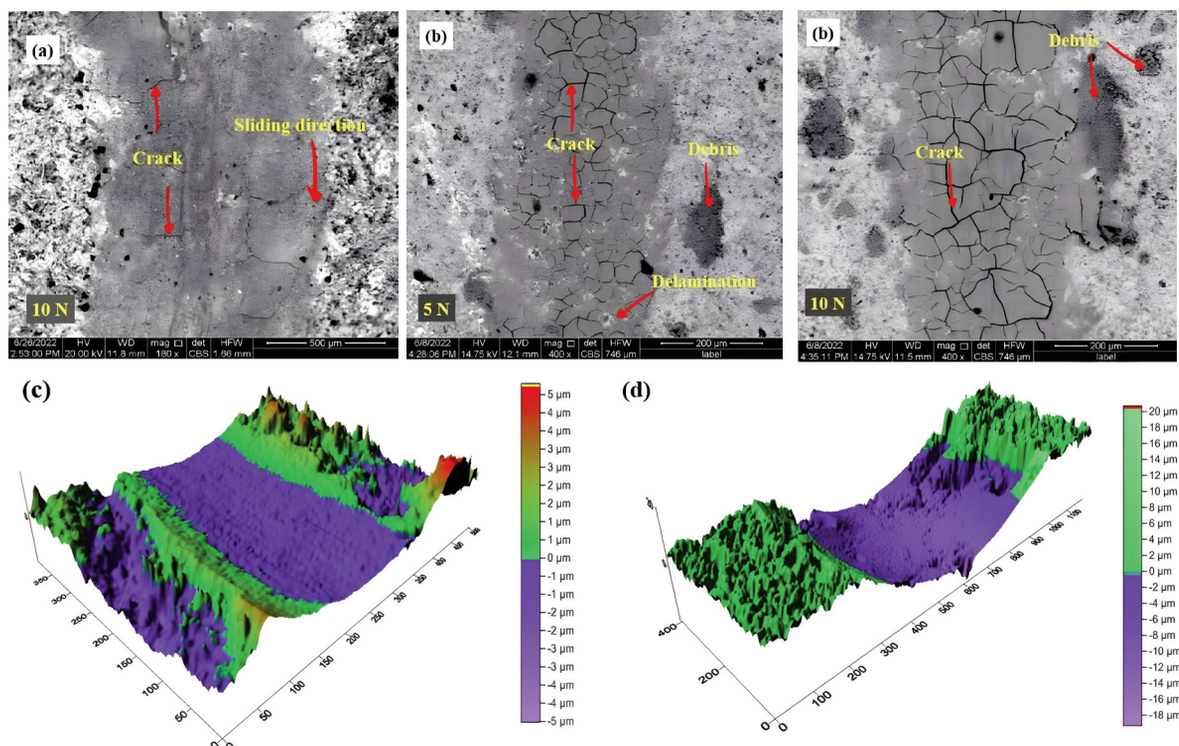


Figure 14. SEM images and 3D morphologies of worm surfaces (a) unmilled (10N), (b) milled with a BPR of 42:1 at 200 rpm for 1 hour (5 and 10 N), (c) 3D of the unmilled sample at 10N and (d) 3D milled sample at 10 N

operational factors such as applied loads and the chemical composition which as mentioned in the first part depends on milling parameters. Milling has a major impact on the grain size and phases formed, which helps explain why the friction coefficient and worm surfaces of samples milled was different from the unmilled.

4. Conclusions

This study successfully investigated the synthesis, characterization and tribological performance of Ni/Al₂O₃ nanocomposites obtained by mechanical alloying. In particular, we examined the effects of milling parameters on the refinement of reactant (NiO and Al) powders and the production of Ni/Al₂O₃ nanocomposite without the addition of PCAs. Sintering process was used to densify the milled powder mixtures. The inferences derived from the present findings are summarised as follows:

Increasing the BPR from 20:1 to 42:1 improved the particle refinement of the reactants and products for each applied speed (100, 150, 200, and 250 rpm). Moreover, homogenous powders at a nanometre-scale were obtained upon exploring the effects of prolonged milling duration with a low milling speed; no composite or intermetallic was detected even after milling for 8 h.

In case of a BPR of 42:1 with a speed of 200 rpm

for 1.5 h, we could generate Ni/Al₂O₃ nanocomposites with particle sizes of 85 nm for Ni particles and 45 nm for alumina.

The findings of SEM and EDS revealed the formation of the metallic matrix nanocomposite via the mechanochemical interaction between NiO and Al and confirmed the crystallite size calculations using the W–H plot.

The DSC data of the milled samples revealed that uncompleted milling reactions might occur during sintering at lower temperatures than those in the unmilled sample.

The XRD patterns of the samples sintered at 800 °C for 2 h exhibited peaks of Ni and alumina phases with traces of unreduced NiO. In particular, the thermite reactions reduced the unreacted NiO and Al during sintering at 1100 °C for 2 h.

The friction coefficient increases with applied forces and decreases with milling.

SEM and 3D morphologies of the worn surface clearly showed that abrasion was the predominant wear process.

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Author contribution

Nor El-Houda Berramdan (PhD student): conceptualization, experimental work, investigation, resources, writing - original draft, writing - review and editing, and visualization. Hafida Boutefnouchet (supervisor): methodology, resources, review and editing. Mosbah Zidani: supervision, visualization and validation. Ridvan Yamanoglu: technical support and help with characterisation. Caroline Curfs: technical support and help with characterisation.

Data availability

Not applicable.

Conflict of interest

The authors declare no competing interests.

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OPTIMIZACIJA PARAMETARA MLEVENJA ZA PREČIŠĆAVANJE MEŠAVINA NiO/Al I SINTEZU NANOKOMPOZITA Ni/Al₂O₃

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Apstrakt

Ova studija je ispitivala efekte parametara mlevenja na razvoj Ni/Al₂O₃ nanokompozita i rafinaciju NiO i Al prahova. Kuglično mlevenje određenih smeša je praćeno sinterovanjem na 800 i 1100 °C tokom 2 h. Rezultati difrakcije rendgenskih zraka suvo mlevenog praha pokazali su da povećanje masenog odnosa kuglica i praha sa 20:1 na 42:1 dovodi do finijih čestica, što je omogućilo sintezu nanokompozita Ni/Al₂O₃ mlevenjem pri 200 o/min. tokom 1,5 h. Produženje trajanja mlevenja pri nižim brzinama rotacije dalo je prahove sa nanoveličinom čestica. Međutim, kao što je pokazano pomoću skenirajuće elektronske mikroskopije i energetske disperzivne spektroskopije, nanokompozit sa metalnom matricom je formiran mehaničko-hemijskom reakcijom, a veličina kristalita je procenjena korišćenjem Vilijamson-Hal dijagrama. Uz to, korišćeni su dijagrami diferencijalne skenirajuće kalorimetrije da bi se analizirali efekti mlevenja na temperature fazne transformacije i/ili reakcije redukcije. Tribološke performanse razvijenog kompozita matrice metala nikla su ispitane korišćenjem tribometra sa kuglicom na disku pod različitim uslovima opterećenja. Zapravo, koeficijent trenja raste sa primenjenim silama i opada sa mlevenjem. Sveobuhvatna ispitivanja istrošenih površina vršena su pomoću skenirajućeg elektronskog mikroskopa i 3D optičkog profilatora.

Ključne reči: Mehaničko legiranje; Ni/Al₂O₃ nanokompozit; Sinterovanje; Aluminotermna redukcija; Koeficijent trenja; Mehanizam habanja

