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Experimental Characterization of Milling, Compaction and Sintering of Nanocrystalline FC-0205 Copper-Steel Powder

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7 Abstract

8 The effect of ball milling on the compaction and sintering of nanocrystalline copper steel

9 powder (FC-0205) was evaluated within this work. The as- received micron-sized FC-0205

¹⁰ copper steel powder were subjected to High Energy Ball Milling (HEBM) in an argon

atmosphere at different milling times of 0, 16, 20 and 24 hours to obtain nanocrystalline

¹² structures. Unmilled, 8 and 16 hour milled powder specimen were compacted using uniaxial

¹³ die compression at pressures ranging from 274 MPa to 775 MPa to obtain a relative density

¹⁴ range of 74

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16 Index terms— nanocrystalline powder, ball milling, uniaxial die compaction, sintering, characterization.

17 1 INTRODUCTION

anocrystalline materials have experienced significant growth in recent years due to their wide range of applications and impressive physical and mechanical properties they exhibit. Since nanocrystalline materials possess a large number of grain boundaries and are dependent on grain size and distribution, they are superior when compared to conventional coarse-grain materials [1]. Properties include enhanced diffusivity, improved ductility, lower thermal conductivity and increased strength [2]. A nanocrystalline powder is characterized by a structural length or grain size on the order of 100 nm (0.1 µm) or less [3][4]. They can consist of single crystals of less than 100 nm or of conventionally sized particles composed of a nanocrystalline grain structure.

Materials that have grain sizes in the nanosize range tend to have higher mechanical properties, such as increased strength and hardness as compared to their coarser grained counterparts [5]. Nanosized grains prompt dislocation build up at grain boundaries, hence, making dislocation movement into adjacent grains more difficult. The higher the applied stress needed to move dislocations, the higher the yield strength of the material. The correlation between improved strength characteristics with decreased grain size is captured by the Hall-Petch (HP) relation [6] in equation (1),?? ?? = ?? 0 + ?? ??? (1)

Where ?? ?? is the yield strength, ?? 0 is the lattice friction stress to move individual dislocations, ?? ?? is a constant (HP slope), and d is the grain size in diameter. The HP relation implies that yield strength increases with decreasing grain size; however, there is a limit to which the material can be decreased to avoid weakening [7][8]. Experiments on nanocrystalline materials have shown that if the grains reached a critical grain size, typically around 10 nm, an inverse Hall-Petch effect is observed [9][10][11][12][13][14]. As the grain size decreases to approximately 10 nm, grain boundaries tend to slide past one another, which results in a reduction of the material's strength.

There are three main techniques to obtain nanocrystalline structures during powder production which influence the size, shape, microstructure, chemistry, and cost of a material [15]. One method involves ball-milling micronsized powders to obtain the nanosize powder. Another method uses gas or water atomization from which powder is formed from molten metal using a spray of the droplet, however, this method has a lower particle size limitation of ~1-5 µm [17]. The third method is inert gas condensation which results in no impurities but produces a small output capacity **??**15, 17, and 18]. Among these techniques, ball milling has proven to be the most cost-effective in the production of nanosized powder [15,17] and is the production method used in most current research. High Energy Ball Milling (HEBM) is a type of mechanical milling (MM) technique that is utilized in powder metallurgy
 (PM) consisting of continuous cold welding, fracturing, and re-welding of powder particles [17].

In 1966, John Benjamin and his colleagues experimented by milling nickel and aluminum powders in an 47 oxidizing atmosphere to form nickel super alloys hardened by oxide dispersion. The experiment showed that 48 repeated fracturing and cold welding during milling exposed new oxide layers on the surface of the particles, 49 which resulted in metal alloying [15][16][17][18][19]. This technique became the genesis of the milling and alloying 50 technique and has since then been extremely useful in many industrial applications. Akhtar et al. [17] studied the 51 mechanical milling of iron and copper powder from micrometer size to nanosize particles using HEBM. Within 52 the experiment, mechanical energy was generated through the collision of 0.5-inch hard steel milling balls and 53 the milling vial container in a controlled argon atmosphere. The energy produced caused deformation of the steel 54 powder into nanosize particles. The final particle size, shape, and size distribution were dependent on material 55 property, milling energy/ frequency, ball size, and the quantity of ball and powder [15]. 56 Akhtar and Fecht [2,20] observed that milling time played a significant role in obtaining the nanocrystalline 57

aluminum powder. Brian and Poquillon [15,22] observed that the energy produced during milling enhanced 58 the interaction between steel powder particles, resulting in plastic deformation, fracturing, and cold welding. 59 60 Suryanarayana et al. [21] found that excessive cold welding of particles greater than 100 nm should be avoided 61 by ensuring a stable point between fracturing and cold welding. Similarly, Joao et al. [19] suggested the addition 62 of 1-2wt% of stearic acid which acts as a Process Control Agent (PCA) to avert immoderate cold welding during 63 milling of powder particles. Particle Size, morphology, and agglomeration all contribute to the overall density and mechanical strength of a green compact. The particle size distribution during compaction also determines 64 the level of porosity and homogeneity of the green compact. 65

Poquillon et al. and German [22,23] reported on the dependence of particle size on the tensile stress of green compacts made from spherical copper powders. Their research discovered that the tensile stress increased as the particle size of the copper powder decreased. Also, the smaller the particle size of the specimen, the higher the fatigue strength due to variation in the stress concentration factor with particle size [24][25]. Mechanical milling has become widely used because of its simplicity and most importantly, cost efficiency compared to other techniques [17]. However, mechanical milling still has its shortcomings that are yet to be addressed by material

regimeers and scientists [19][20][21].

Powder contamination is a issue in mechanical milling. The milling condition, milling time, surface formation,
 particle size and milling environment all determine the level of contamination of the new powder particles.

Possible approaches to reducing contamination are to create a milled environment free of impurities, by ensuring the powder particles are free of impurities, reducing the mill time, and by using a milling vial and ball of similar material to the powdered particles.

Several research engineers and scientists have studied the quantification and modeling of high energy ball milling, but limited success has been achieved. Since the quantification and modeling has not been adequately scoped, production at the industrial-scale is difficult because predicting, controlling and optimization of the process are underdeveloped. Ball milling or mechanical milling is the least expensive method to produce large quantities of nanocrystalline powders [17].

The potential application of nanoscale materials for use as novel structural or functional engineering materials 83 largely depends on the consolidation of nanopowders into bulk nanoscale solids. A central issue in adding 84 nanopowder to engineered parts is the consolidation of the powder into sintered parts with full density and 85 nanograined microstructures ??3-4, 9-11, 26]. Nanocrystalline powders are much more difficult to process and 86 handle than powders in the micron size range. Changes in properties are caused by an increase in surface atoms 87 of nanoscale powders as the particle diameter decreases. Production and processing of nanopowders are difficult 88 due to small particle size and high specific surface areas. Nanoscale powders tend to agglomerate, which requires 89 additional processing to de-agglomerate or suspend the powders. The nanopowder consolidation and densification 90 process strongly depend on the state of powder agglomeration. 91

Consolidation of powders into various shapes are a direct result of cold or hot-pressing techniques. Hot pressing is a technique whereby compaction and sintering of metal powder are done at the same time, however, [27] it is time-consuming. Cold pressing involves the application of pressure on metal powders in a closed shaped die to form a green compact. Vagnon et al. [28] reported that compacts formed via hot pressing had higher densities and improved homogeneity. Powder pressing is usually done at ambient room temperature and the compacted sample is called a green compact [15].

The compressibility of green compacts is extensively dependent on the compaction pressure [29]. The pressure 98 reduces the volume and porosity of the compact powder and increases the density [15]. The amount of pressure 99 needed for compaction of green compacts largely depend on powder characteristics, additives and desired density 100 [22]. Pranav et al. [27] found that higher pressures were required to compress powders with coarser particles 101 in comparison to powders that contained fine particles of the same material. Poquillon et al. and Sang et al. 102 [22,30] investigated the morphology of iron powder and its impact on compaction behavior. They compacted two 103 different iron powders, one with spherical grains and the other with spongy grains at different pressures ranging 104 from 100 to 350 MPa. Their experimental result showed that the iron spherical powder compacts produced lower 105 density compared to the spongy powder compacts at the same compaction pressure. 106

107 Camila and Lirio [31] expanded further on the impact of morphology on the compressibility of steel powders.

They observed that during compaction, particles with flattened morphologies had greater deformation capacity 108 on compacts due to their high specific surface areas, which increased the internal friction between particles, 109 ultimately, reducing the compressibility of the compacts. During compaction, particles tend to rearrange thereby 110 reducing spaces and gaps, which leads to an increase in density. The density increases due to the more organized 111 112 packing of the particles that create contacts points between them are referred to as the initial stage of compaction [32]. Cold welding occurs during further application of pressure because forces increase between the contact points 113 of the particles. Plastic deformation occurs in the second stage of compaction. Material deformation takes place 114 at this stage due to the stress between the contact points of the powder particles. The plastic flow of particles 115 increases the stress between powder particle contact points which causes deformation of the material. Density 116 increase is much slower in the second stage because frictional force is higher which hinders the easy movement 117 of material. Conversely density in initial stage increases rapidly because of lower resistance of frictional force. 118 Work hardening of the powder occurs in the third stage because of increase in compaction pressure [22]. 119

Lubricants are necessary during compaction to reduce frictional forces that hinder the free movement of powder particles. Lubricants help to form a uniform distribution of particles during compaction [33]. However, immoderate lubrication can store in interparticle pores thereby preventing a uniform distribution and proper compaction of powder. The green compact is then heated at lower temperatures to remove the lubricants so that it has no effect on the density and strength of the sample [15].

The final stage in the press-sinter approach is to sinter the green compact at a temperature below its melting point. Sintering of green compacts is essential in that the loosely bounded particles are taken from their green state to higher densified compact parts with enhanced thermal and mechanical properties through metallurgical bonding. There are three processes for sintering; solid-state sintering, liquid-phase sintering and activated sintering [34]. Liquid-sintering of alloy metal powders can be accomplished in one of the two ways; by sintering at temperatures above the melting point of one of the constituent powders or by sintering at temperatures below the melting point of the base powder according to a Fe-Cu phase diagram [34,35].

Compact shrinkage occurs during sintering due to diffusion thereby reducing space and porosity within the sample. Vagnon et al. [28] studied the effect of sintering on swelling and shrinkage of steel powder. They observed shrinkage within the steel powders at lower temperatures during the removal of the lubrication. However, at higher sintering temperatures, the copper diffused into the iron thereby causing the compact to swell and reduce in shrinkage. Also, during sintering, recrystallization of atoms, grain growth, and phase change were all noticed [36].

Sintering parameters [29,37] affect the strength, density and grain size of the of the sintered compact. These parameters include sintering temperature, material composition, powder morphology, time and atmosphere. [17,28]. Nanocrystalline powders densify at sintering temperatures significantly lower than conventional powders. Densification typically occurs at 0.2 to 0.4 times the melting temperature (T m) as compared to 0.5 to 0.8 T m for conventional powders [38].

Several researchers have studied the effects of sintering time and temperatures on the mechanical properties 143 of metallic powders. Akhtar et al. [2] studied the effects of sintering on steel powders and reported a 50% 144 increase in grain size from the initial powder postcompaction, sintering, and annealing at elevated temperatures. 145 Pranav et al. [27] investigated the sintering mechanisms of molybdenum powder and reported that less sintering 146 work was required to achieve the same densification at higher compaction pressures than at lower compaction 147 pressures. Narasimhan [34] stated that metallic powder type and desired geometry determine both sintering time 148 and temperature. He performed an experiment on several alloying elements and concluded that Fe-B-C has the 149 potential to undergo higher sintering temperature than Fe-Cu-C due to the low solubility of boron in iron as 150 compared to copper. Therefore, Fe-B-C can be used in the production of hard steels. 151

Sintering atmosphere is also a key factor in the sintering process of metal powders [29,37]. The sintering 152 atmosphere should be controlled to prevent existing oxides and control carburization and decarburization. 153 [37]. Several types of sintered atmospheres are used in powder metallurgy; however, the method in which 154 the atmosphere is controlled during sintering is consequential in relation to the strength of the green compact. 155 The sintering atmosphere is usually controlled with hydrogen, argon, helium, nitrogen, or under vacuum [27,34]. 156 Previous studies have shown that sintering in a hydrogen environment results in better mechanical properties than 157 under vacuum [37]. Depending on the sintering time and temperature, powder particles start to bond together 158 by forming necks throughout the diffusion process. These necks grow at the points of particle contact and as 159 sintering time increases the neck size ratio increases which help to increase the mechanical strength [37]. At 160 elevated temperatures and increased time, grain growth was noticeable, and pores became spherical and isolated 161 [36]. Vagnon et al. [28] postulated two methods to reduce sintering temperature to minimize the growth of 162 grains. They suggested a small amount of elements such as Ni, Pt, Pd, and Co be added to the base powder to 163 activate the sintering process thereby reducing the sintering temperature and time. Nevertheless, the addition 164 of some of these elements with the base powder can reduce the ductility of the sintered sample thereby affecting 165 the mechanical properties. The researchers [37] also stated that nanosize particles require higher compaction 166 pressures at lower sintering times and temperature for grain growth reduction. 167

This experimental study establishes a correlation between milling time, particle size, temperature and grain growth through HEBM on copper steel powder. The aim of this work was to characterize the structure-property relations of ball-milled nanocrystalline FC-0205 copper steel powder with Ancorsteel 1000B as the base Fe powder; analyze the morphology of powder particles from different milling times and how that particle size affects the density and mechanical properties of the green compact; and lastly, analyze the effects of ball milling on powder consolidation into dense compacts using uniaxial die compression and sintering. The FC-0205 powder system was chosen due to its widespread use in the powder metallurgy industry for automotive applications. The study utilizes a press-sintering technique using uniaxial die compaction followed by sintering.

176 **2** II.

¹⁷⁷ **3** Experimental Procedures a) Milling

As-received copper steel powder FC-0205 mixed with 0.6% zinc stearate lubricant, also known as Acrawax from 178 Hoeganaes was used for this study. Table 1 summarizes the nominal chemical composition of the as-received iron 179 powder. The prefix (F) designates an iron-based material and the second prefix (C) is copper, which is known as 180 the primary alloying element. The first two digits (02) represent the percentage of the primary alloying element 181 which is 2% copper. Carbon content is represented by the third and fourth digits whereby 05 designates the 182 amount of carbon in a range between 0.3%-0.6%. The base iron powder for FC-0205 is an Ancorsteel 1000B 183 alloy mix commonly used in the powder metallurgy industry. The chemical composition of Ancorsteel 1000B is 184 provided in Table 2. The as-received FC-0205 powder was milled by Union Process at intervals of 0.8,16, 20 and 185 24 hours. The milling process was carried out in a Model SD-1 Laboratory Attritor with 6.35 mm (0.25 inch) 186 stainless steel grinding media. A ball-to-powder ratio of 16/1 was used, and the milling speed was set to 350 RPM 187 in a sealed argon environment to lessen the contamination of oxygen and humidity. The stearic acid lubricant 188 was added to the milling process to reduce friction, minimize die wear, aid in the part ejection, and avoid cold 189 welding of the materials. The material properties of the final components are unaffected by the lubricant since 190 the lubricant is burned off before sintering. The shaped die used for compaction and sintering was made of Vasco 191 Max C-350 and had an inner diameter of 32.63 mm (1.3 inches), and an outer diameter of 101.33mm (4 inches) as 192 shown in Figure 1. A cylindrical split hollow die with a diameter of 12.67 mm (0.5 inches) was made of the same 193 material for easy removal of the sample after compaction. An Instron 5882 mechanical test frame with 100 kN 194 load cell was used to exert a compressive force on the shaped die at a constant compression rate of 10 mm/min 195 in a controlled environment. The steel powder was filled into the shaped die, and the upper punch compressed 196 the powder at various heights to achieve different densities. Density measurements were taken before sintering. 197

¹⁹⁸ 4 c) Dilatometer Tests

A vertical push rod dilatometer (Anter Corporation Unitherm Model 1161) was used for the dilatometry analysis 199 to examine the density and phase transformations of the unmilled samples during sintering. FC-0205 compacts 200 were heated to 1150 °C (2102 °F) in a nitrogen atmosphere at different heating rates (2C/min, 5C/min, and 201 10C/min). The samples were held for 30 minutes followed by an uncontrolled cool down from the sintering 202 cycle. Dilatometry measurements were captured when the push rod attached to a transducer displaced with 203 the shrinkage or expansion of the compact. The signals obtained from the motion transducer and thermocouple 204 were correlated by a computer which provided the dimensional change of the sample as a function of time and 205 temperature. 206

²⁰⁷ 5 d) Sintering

A Carbolite 1600°C (2912 °F) tube furnace was used to sinter the compacts in a controlled environment of 75% hydrogen and 25% nitrogen to avoid oxidation and contaminants. The bulk specimens were de-lubed at 400 °C (752 °F) to get rid of the stearic acid in order not to hinder the overall density and strength. Bulk samples from each milling hour were sintered at 900 °C (1652 °F) and 1120 °C (2048 °F) which is close to the critical sintering temperature of FC-0205 powder [31]. The temperature was set to ramp at 15 °/min and dwell for 30 minutes before it was slowly cooled to room temperature. The compaction and sintering schedule minimize grain growth and maximize densification during the process.

²¹⁵ 6 e) Microstructure Analysis and Sample Preparation

Struers LaboPress hot mounting technique was carried out on each sample for easy handling of the sample during 216 grinding and polishing. Each surface of the mounted samples was subjected to grinding and polishing for proper 217 218 analysis of the microstructure using SEM, optical microscopy (OM) and image analysis. The sintered samples 219 were etched with 2% nitric acid and 78% ethanol to remove a uniform thin layer of the surface off for analysis of 220 the grain size and grain growth. Zeiss Evo Supra 40 Field Emission Gun Scanning Electron Microscope FEG-SEM 221 was used to analyze the morphology, particle distribution, grain size and agglomeration of milled and compacted 222 FC-0205. Rikagu Ultima III x-ray diffraction system was used to analyze the average grain size of milled FC-0205. A thin layer of FC-0205 iron powder was evenly spread on a glass sample holder. This procedure was carried out 223 on each powder sample of different milling time. The $2?^{\circ}$ (FWHM) range was set from 10° to 90° at $1^{\circ}/\text{min}$ to 224 cover the essential part of the powder pattern. ImageJ analysis was performed on images obtained from optical 225 microscopy to analyze the extent of grain growth in the FC-0205 samples. 226

²²⁷ 7 f) Hardness Test

Rockwell hardness (HRA) measurements using Leco Rockwell Hardness Tester were carried out on bulk specimens obtained through the compaction and sintering of FC-0205 steel powders. The indenter had a diamond cone shape for indenting. The indenter was forced into the material under a total load of 60 kgf. At least four different readings were taken from each surface of the sample, and the arithmetic average was obtained.

232 **8 III.**

²³³ 9 Milling, XRD, And Microscopy Results

²³⁴ 10 a) Ball Milling of Nanocrystalline Copper Steel Powders

Table 3 and Figure 2 shows analysis of FC-0205 particle size using particle size analyzer. A linear function relationship is obtained between milling time and average particle size of the copper steel powder. The linear function relationship is experimentally expressed in equation (2).

238 11 D=kt+71.57 (2)

Where D is the average particle size in (μm) , t is the milling time in (hour) and k is a constant. The correlation coefficient of equation (??) is 0.998.

It was observed that as milling time of powder increased the average particle size decreased which agrees with 241 previous studies [22,2] (see Table 3). As milling time increased from 0 to 16 hour, the rate of fragmentation of 242 particles was high leading to a high reduction in particle size from 72.37 (µm) to 25.64 (µm). After further milling 243 from 16 to 24 hours, it was noticed that fragmentation of particles decreased which led to a reduction in particle 244 size from $25.64 \,(\mu m)$ to $3.7 \,(\mu m)$. The reduction in particle size is due to an increase in milling energy which 245 causes an increase in the contact area between the grains. The particle size analysis experiments also offered 246 information on the distribution of particle diameters in each sample. Figure 3 shows the particle size distribution 247 plots for each powder analysis. For each successive increase in milling time, the peak of the distribution is seen 248 to gradually skew from a right-sided distribution at 0-hour to a significantly left-sided distribution after 24 hours 249 of milling with the peaks becoming narrower. The particle size distribution for the 8, 16, and 20-hour milled 250 samples shows a mono-modal peak type compared to 0 (unmilled) and 24 milling hour which show a bi-modal 251 252 peak type. The reason for the bimodal peak could be as a result of higher particle size and coarser morphology of the 0 (unmilled) powder and high agglomeration obtained in the 24-hour milled powder. It was observed that 253 at maximum peak for each milling distribution, the corresponding particle size decreased with increased milling 254 time. For the unmilled powder, the average particle size is 72 (µm). In comparison, for the 24-hour milled 255 256 powder, the average particle size is less than 5 (μ m).

²⁵⁷ 12 b) Scanning Electron Microscopy (SEM) of Copper Steel ²⁵⁸ Powders

Scanning electron microscopy of the copper steel powder was performed to compare the morphology, agglomer-259 ation and particle size at different milling time and to determine the effect of particle size on consolidation. The 260 SEM micrographs of the 0, 16, 20 and 24-hour milled copper steel powder at 5000X magnification is shown in 261 Figure ??. It was observed that as milling time increased from 0 to 24-hour, the particle size became less coarse 262 but was still in the micron meter range. However, higher agglomeration and morphology change of the steel 263 powder was noticed as milling time increased. The morphology of the particle size for 0-hour (unmilled) powder 264 265 is spherical as shown in Figure ??. As the milling time increased the particle size became flat and spongy with little agglomeration as shown in Figures ?? (b) and 4(c). The morphology change of the particles is due to an 266 increase in milling energy which causes the contact area between grains to increase, as well as, an increase in 267 the amount of coalesced particles, and a reduction in the plastic deformation of particles. The flat and spongy 268 structure obtained due to increase milling time will be difficult to compact because of the high surface area which 269 increases internal friction between particles. Hence, higher compaction pressure will be required to avoid high 270 porosity and obtain high density. Increased agglomeration and a changes in morphology are also apparent in 271 the 24-hour milled powder, as shown in Figure ?? (d). Increase in milling time causes more fracture of particles 272 which results in further deformation and fragmentation of the steel particles. where L is average grain size in 273 (nm), t is milling time in (hour) 274

A linear reduction in grain size is obtained as milling time increases from 0 (unmilled) to 24-hour milled powder as shown in Figure ??. The X-ray diffraction peak patterns with angle 2?° values in the range from 10° to 100° for FC-0205 copper steel powders as shown in Figure ??. Figure ?? (a) shows a sharp diffraction line known as the full widths at half maximum (FWHM), which occurs in the micrometer range as shown in Table 4. However, as milling time increased, the diffraction line widths broadened, as shown in Figures ?? (b, c, and d) and grain decreased to the nanosize range as shown in Figure ??. The change noticed as a result of milling time is due to grain size refinement and increase in atomic level strains.

The results of the X-ray diffraction experiment, used to characterize the correlation between the grain sizes and milling time, are shown in Table 4 and

²⁸⁴ 13 IV. Compaction And Sintering Results

Figure ?? shows the variation in relative density of 0 (unmilled), 8 and 16-hour green compacts of FC-0205 285 286 powder as a function of compaction pressure. As pressure increased from 300 MPa, the 0-hour (unmilled) green compacts had higher relative density than 8-hour green compacts. Similarly, the 8-hour green compacts show 287 a higher relative density when compared to the 16hour green compacts. Thus, increased milling time of the 288 powders resulted in decreased the density of the green compacts at a given pressure. The observation agrees 289 with previous studies on copper steel powder [39]. It was observed that between 300 MPa to 400 MPa, 0-hour 290 (unmilled) compacts show a higher relative density than 8-hour 16-hour compacts. Also, an insignificant difference 291 in relative density between 0-hour and 8-hour green compacts is noticed. However as milling time increased to 292 16 hours, a significant drop in relative density is observed. The spongy structure that resulted from increased 293 milling, hindered compaction. Therefore, higher compaction pressures are required to increase the density of the 294 295 milled powder compacts.

Where D is relative density, P is compaction pressure and K is a proportionality constant. The correlation 296 297 coefficient of the equations is 0.99, 0.97 and 0.98 respectively. The resulting dimensional change from the 298 dilatometer experiments for the unmilled FC-0205 compacts heated at 10 C/min, along with the corresponding temperature cycle is shown in Figure 10. The FC-0205 compact presents dimensional change after sintering 299 due to the difference in alloy composition and powder size. As shown in Figure 10, the copper steel compacts 300 experienced a thermal expansion of 0.7% at 700 °C (1292 °F). At this point, the iron ferrite, having a bcc 301 structure, started to transform to austenite an FCC structure with some evidence of sample contraction. As 302 temperature increased, carbon diffused into the austenite region. The copper in the sample began to melt at 303 1083 °C (1981 °F) which caused expansion of the compact. Near 1120 °C (2048 °F) all the copper melted and 304 305 penetrated the austenite structure. During the 30 minutes hold time the sample started to contract again. As 306 temperature decreased during uncontrolled cooling, copper precipitated from iron and austenite iron transformed into ferrite and pearlite (a mixture of ? ferrite and cementite Fe 3 C). After sintering the copper regions were no 307 308 longer in the form of discrete particles, and the pearlite structure dominated the ferrite structure. The sintering behavior of the iron-coppercarbon system is very different compared to that of other PM metals. Figure 11 309 shows the typical shrinkage behavior during heating for nickel, 316L stainless steel, and bronze as compared to 310 the expansion of the FC-0205 material during heating. The FC-0205 sample expanded to 1.5% at the maximum 311 heating temperature 1150 °C (2102 °F). As the FC-0205 sample cooled, the material contracted, resulting in an 312 overall dimensional change (shrinkage) of only 0.2% in the sample. FC-0205 copper steel powder samples were 313 314 compressed to various heights to achieve different densities. Subsequently, the bulk samples were sintered at 315 900 °C (1652 °F) and 1120 °C (2048 °F) in a controlled nitrogen and hydrogen atmosphere to analyze density 316 change with an increase in temperature. An insignificant difference in relative density Was obtained between green compacts and bulk compacts sintered at 900 °C for 0 (unmilled) powder as shown in Figure 12. However, 317 318 as milling time increased to 8 hours, a slight increase in relative density was noted between green compacts and compacts sintered at 900 °C (1652 °F) and 1120 °C (2048 °F) as shown in Figure 13. Figure 14 shows a 319 significant increase in relative density with increased temperature for the 16-hour milled compacts. The increase 320 in relative density is due to finer particles dissolving or diffusing into the base metal thereby creating a bonded 321 final compact. Shrinkage during sintering can also be a factor due to a reduction in space and pores in the 322 compact. The microstructure in the 0-hour and 8-hour compacts sintered at 900 °C (1652 °F) show a better 323 324 compaction and sintering of iron particles as shown in Figure 15 (b & d). Large pores are noticed in the 0-hour 325 (unmilled) green compact as shown in Figure 15 (a) due to low compaction pressure. The large pores can affect the strength of the compact and act as a point of crack initiation which can lead to pre-mature fracture. High 326 compaction pressure and sintering temperature can help control large porosity. Figure 15 (a) also shows elements 327 and particles that are distinctly separated from each other, i.e., heterogeneous due to the unmilled powder of 328 the green compact. The heterogeneous separation of particles allows the fine particles to fill in the inter-space 329 between large particles, hence, increasing the powder packing and density of the green compact as obtained in 330 the compressibility curve. The microstructure in 0 and 8-hour milled compacts sintered at 900 °C (1652 °F) show 331 sinter bonding of powder particles and also the formation of pearlite as seen in Figure 15 (b and d). The shape, 332 space and amount of pearlite depend on carbon diffusion and processing conditions. As temperature increases 333 to 900 °C (1652 °F), iron ferrite transforms to austenite. As the sintering temperatures increase from 900 °C to 334 335 1120 °C (2048 °F), the diffusion of carbon into austenite takes place, along with the formation of pearlite in 0 and 8hour sintered compacts as shown in Figure 15 (c and e). 336

³³⁷ Copper completely diffuses into austenite as temperature increases from 900 °C (1652 °F) to 1120 °C (2048 ³³⁸ °F) for 0, 8 and 16 hour compacts. The diffusion of copper into iron austenite created some pores as shown in ³³⁹ Figure 15 (c & e). Similarly, increase in sintering temperature of the compact causes phase change and grain ³⁴⁰ redistribution. As temperature increases from 900 °C (1652 °F) to 1120 °C (2048 °F) for 0 and 8 hour compacts, ³⁴¹ phase changed occurred from solid state phase to liquid state phase with the formation of high dominant pearlite ³⁴² in the microstructure as shown in Figure 15 (c & e). In the process of slow cooling, copper precipitate with the ³⁴³ formation of pearlite and ferrite (cementite Fe 3 C) as shown in Figure 15 (c & e). V.

³⁴⁴ 14 Microstructure Measurement

The microstructural grains of FC-0205 sintered compacts were measured using ImageJ analysis free software 345 according to ASTM standard E 112. The objective was to ascertain the extent of growth concerning sintering 346 temperature. The sintered microstructure obtained from OM shows the variation of grain size distribution of 347 copper steel compact in Figure 16 and also the image analysis grain size data in Table 5. As temperature 348 increases to 900 °C (1652 °F), the formation of big and small microstructural non-uniform grain structures was 349 noticed as seen in Figure 15 (b and d). Further increase in sintering temperature to 1120 °C (2048 °F) resulted in 350 smaller grains coalescing to form a more uniform grain structure with pores due to copper diffusion into austenite 351 iron as shown in Figure 15 (c and e). The increase in temperature resulted in grain size increase from 13.45 352 (um) to 13.76 (um) for 0-hour (unmilled) sintered compacts, similarly from 13.49 (um) to 13.77 (um) for 8-hour 353 sintered compacts as obtained in Table 5. As temperature increased, grain growth is obtained as seen in Figure 354 16. Unmilled and 8-hour sintered compacts indicated little increase in grain size and density as temperature 355 increased from 900 °C (1652 °F) to 1120 °C (2048 °F). However, as temperature increased from 900 °C (1652 °F) 356 to 1120 °C (2048 °F) in 16-hour sintered compacts, a significant increase in grain size and density is evident due 357 to morphology and size of the powder. 358

359 15 a) Hardness Test

Table 6 shows the mechanical properties obtained for FC-0205 copper steel powder using Rockwell hardness test 360 at a maximum load of 60 kg. Four separate tests were performed on each sample, and the average was obtained for 361 proper analysis of the strength. Each sample was polished to avoid spatial variability in hardness. It was noticed 362 that Rockwell hardness for each milling hour increased as temperature increased, demonstrating the impact of 363 sintering temperature on compact strength. However, Rockwell hardness for the 0-hour (unmilled) and 8-hour 364 compacts was relatively close, but a significant difference was observed in the 16-hour compacts as temperature 365 increased. Increase in density increased hardness of the sintered compacts as shown in Figure 17 and Table 366 7. Also, in This experimental study establishes a correlation between milling time, particle size, temperature 367 and grain growth through HEBM on copper steel powder. Results from this study agrees with previous studies 368 which show that increased milling time of powder particles decreased the average particle size as well as grain 369 size due to high rate of particle fragmentation. Although at higher milling time, fragmentation rate was low 370 resulting in low reduction in particle size. The particle size reduction rate was as a result of higher milling 371 energy which increases the contact areas between grains, therefore, welding of particles is obtained. Particle 372 size analysis showed distribution peaks that was getting narrower and also skewing from right to left due to 373 increased milling time of the copper-steel powder. The increase in milling time caused change in morphology 374 and higher agglomeration of the powder particles. The change in morphology noticed with increased milling 375 can hinder compaction because of high surface area that increases internal friction between particles. Therefore, 376 higher compaction pressure will be required to achieve a good dense compact. A linear reduction in grain size 377 was obtained as milling time increased. As pressure was increased in green compacts, density also increased. It 378 was noticed that at the same pressure range, lower milling hour compacts obtained higher density. Therefore 379 higher compaction pressure is required for higher milling hour to avoid a high porous compact. Contraction of 380 copper steel compact was noticed at 700 °C (1292 °F) as the microstructure transformed from iron ferrite to iron 381 austenite. Further increase in temperature to 1083 °C (1981 °F) resulted in copper melting into iron austenite 382 which led to expansion of the compacts. It was observed that increase in milling time between 0 (unmilled) 383 384 and 8 hour compacts showed a slight increase in relative density as temperature increased to 1120 °C (2048 °F). 385 A significant increase in relative density was obtained for 16 hour compacts at the same temperature. Hence, 386 higher milling time and higher temperatures resulted in increased density of the compacts. increases the density 387 of the compacts at the same temperature increase. However, grain growth was significant in the 16-hour milled compacts as temperature was increased during sintering as compared to the 0 (unmilled) and 8 hour compacts 388 sintered at the same temperature. As compared to the 0-hour and 8-hour milled and sintered samples, the 16hour 389 milled, sintered compacts The finer sized powders require higher compaction pressure to increase presintering 390 $1 \ 2 \ 3$ density and reduced sintering temperature to prevent grain growth. 391

¹Experimental Characterization of Milling, Compaction and Sintering of Nanocrystalline FC-0205 Copper-Steel Powder

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Figure 2:



Figure 3: Figure 2 :



Figure 4: Figure 3



Figure 5:



Figure 6: Figure 7 . Figure 5 :



Figure 7: Figure 6 : Figure 7 :



Figure 8: Figure 8 : 1 (1 ? 1 (1 ?



Figure 9: Figure 9 :



Figure 10: Figure 10 :



Figure 11: Figure 11 :







Figure 13: Figure 13 :



Figure 14: Figure 14 :



Figure 15: Figure 15 :



Figure 16: Figure 16 :

1

MPIF designation/ Component FC-0205		Fe 97.1 0.5	С	Cu 2	Acrawax 0.6
2	Figure 17: '	Table 1 :			
3	Figure 18: '	Table 2 :			
4	Figure 19: '	Table 3 :			
5	Figure 20: '	Table 4 :			

Figure 21: Table 5 :

Sample	Relative	Count	Total Area	Average	%Area	Mean	Mean
	Density		$(nm\ 2\)$	Size (?m)		(2m 2)	Diameter
	(%)						(?m)
0hr-900C	0.92	284	54660.03	192.47	91.6	142.07	13.45
0hr-1120C	0.95	284	54789.03	192.92	90.82	148.62	13.76
8hr-900C	0.94	235	55422.57	235.84	92.18	142.87	13.49
8hr-1120C	0.95	283	54551.93	192.76	91.2	149	13.77
16hr-900C	0.91	380	54698.83	143.94	90.98	155.97	14.09
16hr-1120C	0.92	211	55683.8	263.9	92.62	173.66	14.87

Figure 22: Table 7 ,

6

 $\mathbf{7}$

Figure 23: Table 6 :

7

Experimental Characterization of Milling, Compaction and Sintering of Nanocrystalline

	Sample	1	2
	0hr 900 31.9		30.8
	Unr 1120 32.2		29.5
	8hr 900 34.4		32.6
12 Year 2018	8hr 1120 35.1 16hr 900 16.1 16hr 1120 22.2	Inc	dentor Diamond cone 34.9 17.9 22.8 Hardness HR
A () Volume XVIII Issue I Version I	Total Load 60kgf Sample Relative Density	0hr	900 0.921 0hr 1120 0.951 8hr 900 0.936 8hr 1120
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neering		T 7	
Global		V.	I.CONCLUSION
	© 2018 Global Journals		

Figure 24: Table 7 :

- [James], Brian W James. Powder Metallurgy Methods and Applications 7. (Powder Metallurgy) 392
- [Nieh et al. ()], T G Nieh, P Luo, W Nellis, D Lesuer, D Benson. Acta Mat 1996. 44 (9) p. . 393
- [Furukawa et al. ()], M Furukawa, Z Horita, M Nemoto, R Z Valiev, T G Langdon. Acta Mater 1996. 44 p. . 394
- [Alymov ()] , M I Alymov , MaltinaE I , StephanovY . Powder Metal Technology and Applications 1998. American 395 Society for Metals. 7 p. 506.
- [Kumar et al. ()], K S Kumar, H V Swygenhoven, S Suresh. Acta Materialia 2003. 51 p. . 397
- [Kumar et al. ()], K S Kumar, H V Swygenhoven, S Suresh. Acta Materialia 2003. 51 p. . 398
- [Tjong and Chen ()], S C Tjong, Haydn Chen. Nanocrystalline Materials and coatings 2004. p. . (R45) 399
- [Carlton and Ferreira ()], C E Carlton, P J Ferreira. Acta Materialia 2007. 55 p. . 400
- [ASM Handbook ()] ASM Handbook, 1998. American Society for Metals. 7 p. 77. 401
- [Poquillon et al. ()] Cold compaction of iron powders relations between powder morphology and mechanical 402 properties, D Poquillon, V Baco-Carles, Ph Tailhades, E Andrieu. 2002. p. . 403
- [Vagnon et al. ()] Deformation of Steel Powder Compacts during Sintering: Correlation between Macroscopic 404 Measurement and in Situ Microtomography Analysis, A Vagnon, O Lame, D Bouvard, Di Micheal, M 405 Bellet , D Kapelski , G . 2006. p. . 406
- [Dieter ()] G Dieter . Mechanical Metallurgy 3 rd Edition, 1986. 407

396

- [Edelstein and Cammarata ()] A S Edelstein , R C Cammarata . Nanomaterials: Synthesis, Properties and 408 Applications, 1996. Bristol and Philadelphia. 409
- [Ho Na et al. ()] Effect of Ball-Mill Treatment on Powder Characteristics, Compaction and Sintering Behaviors 410
- of ex-AUC and ex-ADU UO Powder, Sang Ho Na, Si Hyung Kim, Young-Woo Lee, Dong Seong Sohn. 411 2002. р. . 412
- [Pranav et al. ()] Effect of Die Compaction pressure on Densification Behavior of Molybdenum Powders, G 413 Pranav, P Seong-Jin, M Randall. 2007. p. . 414
- [Schaffer ()] Effect of High Energy Milling on the Microstructure and Properties of WC-Ni Composite, Camila 415 Dos Santos Torres & Lirio Schaffer . 2010. p. . 416
- [Montasser Dewidar et al. ()] 'Effect of Processing Parameters and amount of additives on the Mechanical 417 Properties and Wear Resistance of Copper-based Composite'. G T Montasser Dewidar, Mahmoud Abdel-418 Jaber, Hussien Bakreya, Badry. IJMME-IJENS 2013. 10. 419
- [Honda ()] 'Effect of sintering Temperature and Time on Fatigue Strength of Iron Compacts with Different 420 Particle Sizes'. T Honda . J. Jpn. Soc. Powder Powder Metall 1983. 30 p. . 421
- [Ward and Billington ()] 'Effect of Zinc Stearate on Apparent Density, Mixing and Compaction /Ejection of Iron 422 Powder Compacts'. M Ward, & J Billington. Powder Metallurgy 1979. 22. 423
- [Enneti et al. ()] Effects of Lubricant on Green Strength, Compressibility and Ejection of Parts in Die Compaction 424 process, Ravi K Enneti, Adam Lusin, Sumeet Kumar, Randall M German, Sunday V Atre. 2013. p. . 425
- [Haynes ()] 'Fatigue behavior of sintered metal and alloys'. R Haynes . PM 1970. 13 (26) . 426
- [German (1976)] R R M German . The Sintering of 304L Stainless Steel" Metallurgical Transactions, Dec 1976. 427 7 p. . 428
- [Farrokh and Khan ()] Grain Size, Strain Rate, and Temperature Dependence of Flow Stress in Ultr-Fine grained 429 and Nanocrystalline Cu and Al: Synthesis, Experiment, and Constitutive Modeling, Babak Farrokh, Akhtar 430 431 Khan . 2009. p. .
- [Hansen ()] 'Hall-Petch relation and boundary strengthening'. Neils Hansen . Hansen Scripta Materialia 2004. 432 433 51 p.
- [Nieman et al. ()] 'Mechanical behavior of nanocrystalline Cu and Pd'. G W Nieman , J R Weertman , R W 434 Siegel. Journal of Material Research 1991. 6 (5). 435
- [Meyers et al. ()] 'Mechanical properties of nanocrystalline materials'. M Meyers , A Mishra , D Benson . Progress 436 in Materials Science 2006. 51 p. 444. 437
- [Akhtar et al. ()] Mechanical response and modelling of fully compacted nanocrystalline iron and copper, S Akhtar 438 , Hayoyue Khan , Laszlo Zhang , Takacs . 2000. p. 439
- [Akhtar et al. ()] Nanocrystalline Aluminum: Mechanical Behavior at Quasi-Static and high strain rates, and 440 constitutive modeling, S Akhtar, Yeong S Khan, Xu Suh, Hayoyue Chen, Laszlo Zhang, Takacs. 2006. p. 441 442
- [Fecht et al. (1990)] 'Nanocrystalline metals prepared by high-energy ball milling'. H J Fecht, E Hellstern, Z 443 Fu, W L Johnson. Metall. Trans. A Sep. 1990. 21 p. . 444

- [Conrad and Narayan ()] 'On the grain size softening in nanogcrystalline materials'. H Conrad , J Narayan .
 Scripta Mater 2000. 42 p. .
- [Narasimhan ()] Sintering of Powder mixtures and the growth of ferrous powder metallurgy, K Narasimhan .
 2001. p. .
- [GermanR ()] Sintering Theory and Practice, GermanR . 1996. New York: John Wiley and Sons.
- [German] 'Strength Dependence on Porosity for P/M Compacts'. R M German . Int. J. Powder Met. Powder
 Tech
- [Sanders et al. ()] 'Tensile behavior of nanocrystlline Cu the Minerals'. P G Sanders , J A Eastman , J R
 Weertman . Metals and Materials Society 1996. p. .
- 454 [Fogagnolo et al. ()] The Effects of Mechanical Alloying on the Compressibility of Aluminum Matrix Composite
 455 Powder, Joao Fogagnolo, Elisa Ruiz-Navas, Maria Robert, Jose Torralba. 2003. p. .
- [Bonetti et al. ()] The influence of grain size the mechanical properties of nanocrystalline Aluminum, E Bonetti
 , L Pasquini , E Sampaolesi . 1997. 9 p. .
- [Suryanarayana et al. ()] The Science and Technology of Mechanical Alloying, C Suryanarayana, E Ivanov, V
 Boldyrev. 2001. p. .
- [Suryanarayana (2002)] 'The Structure and Properties of Nanocrystalline Materials'. C Suryanarayana . Issues
 and Concerns September 2002. (JOM)