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Nano/ultrafine grained immiscible Fe-Cu alloy with ultrahigh strength produced by selective laser melting

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ABSTRACT

A nano/ultrafine grained immiscible Fe-Cu alloy was produced using selective laser melting (SLM) with an average grain size of 250 nm and many 50–100 nm, the finest ever achieved using additive manufacturing (AM). The substantial grain refinement was attributed to liquid separation, monotectic reaction and solid-state phase transformations upon cyclic heating. Cu particles of ∼5 nm in size and ∼10 nm in spacing were contained in some grains, resulting in dispersion strengthening which, together with grain boundary strengthening, led to a significant increase in the yield strength from ∼400 MPa in an SLM-fabricated Fe to ∼900 MPa in Fe-Cu.

IMPACT STATEMENT

Nano/ultrafine Fe grains containing Cu particles of 5 nm in size and 10 nm in spacing were achieved by SLM of Fe-Cu, resulting in an ultrahigh strength material.

1. Introduction

Alloys with a positive enthalpy of mixing (ΔH_{mix}) are immiscible, exhibiting little solubility between solute and solvent. Systems with large positive ΔH_{mix} show immiscibility in both solid and liquid states, possessing a liquid miscibility gap [1]. Although systems with less positive ΔH_{mix} may be miscible in the liquid state [1], a metastable liquid miscibility gap is observed if sufficiently undercooled upon rapid cooling [2–4]. In the miscibility gap, the liquid decomposes to two with the minority forming droplets in a matrix of the majority, with the droplets subsequently solidifying into either layers or particles depending on the cooling rate (T) [5,6]. Such alloys have potential for applications, including bearing materials (e.g. Cu- and Al-based alloys [7–9]) and high strength alloys (e.g. Fe-Cu containing nano-Cu precipitates [3,10]).

The fabrication of immiscible alloys is challenging, though, because of Stokes sedimentation and Marangoni motion causing macro-scale segregation [5,11,12]. Efforts have been made to overcome this difficulty using various methods including atomisation [13], melt spinning [14,15], spray deposition [16,17] and laser and electron beam cladding [3,18]. They have rapid solidification in common, hindering macro-segregation and creating more uniform dispersion of particles. However, these methods only produce coatings, thin sheets or powders requiring post-processing.

The full potential can be exploited using AM, delivering cooling rates of 10^3–10^8°C/s and near-net-shape. Moreover, cyclic heating in AM would allow solute atoms trapped in the matrix during rapid solidification to precipitate as finely spaced nano-particles [10,19,20], leading to increased strength. A recent study [21] showed that a dense Fe-Cu immiscible alloy can be produced using AM, although microstructural evolution and mechanical properties were not investigated.

A secondary phase in immiscible alloys can facilitate grain refinement in AM. Although equiaxed Fe-grains form due hypothetically to α → γ → α transformation
grains in most AM-fabricated cubic metals are columnar owing to epitaxial growth (e.g. Al [23,24], and β-Ti [25,26]). Much attention has been paid to columnar-to-equiaxed transformation and grain refinement in AM-fabricated metals by adding high melting-point secondary particles [24,27], vibrating the melt pool by ultrasound [28] and triggering eutectic [29] or peritectic [30] reactions. However, immiscible alloys have not been produced using AM to utilise liquid separation and monotectic reactions to refine grains. Hypothetically the secondary phase in a previously deposited layer could act as nucleation sites for grains in the newly added layer, preventing epitaxial growth and creating refined equiaxed grains. A good heat-conducting secondary phase (e.g. Cu in Fe-Cu [18]) can also increase undercooling and enhance grain refinement [12,31]. Additionally, a secondary liquid may wet the solidifying grains [32] and separate them from the primary liquid, causing grain refinement.

Here, we used SLM to process an immiscible Fe-Cu alloy. Nano/ultrafine Fe-grains containing nano-spaced, nano-Cu precipitates were achieved. Grain boundary and dispersion strengthening mechanisms substantially increased the yield strength.

2. Materials and methods

Pure Fe and a mechanically mixed powder of gas atomised Fe (99.8%) and 20 vol.% Cu (99.7%) (both 15–45 μm) were used in Renishaw AM250 to produce rods of ∅10×8 mm using stripe scanning strategy on 304SS substrate at room temperature in an atmosphere set at 100 ppm oxygen. Other parameters are listed in Table 1.

Microstructures were characterised using SEM (FEI Quanta and Teneo) and TEM (FEI Tecnai F20). TEM samples were cut by in-situ lift out FIB (FEI Nova 200 Nanolab). EBSD was performed using 17.5 kV, 9.5 nA and step sizes of 10–20 nm. Compression tests were conducted on specimens of ∼∅5×8 mm at a strain rate of 10⁻³ s⁻¹.

3. Results

Figure 1(a) shows equiaxed grains of ∼1 μm in the SLM-Fe, finer but comparable to the average grain size (D) of ∼5–12 μm observed by others [22,33–35]. Figure 1(b–i) shows the SLM Fe-Cu. In each melt pool, clusters of Fe and Cu would form, driven by Marangoni convection, a layered structure of Cu-rich regions (Fe-Cu alloy) and Fe-rich ones (nearly pure Fe), the latter being much thicker thanks to much higher Fe concentration.

**Table 1.** Critical printing parameters.

| Stripe size | Layer thickness | Point distance | Exposure time | Power | Hatch spacing | Laser spot size | Rotation angle between layers |
|-------------|----------------|----------------|---------------|-------|---------------|-----------------|-----------------------------|
| 5 mm        | 50 μm          | 60 μm          | 90 μs         | 200 W | 110 μm        | 66 μm           | 67°                         |

**Figure 1.** (a) SEM on the as-SLM pure Fe, showing grains of ∼1 μm. (b–i) SEM on the as-SLM Fe-Cu: (b) Cu-rich layers created by the Marangoni convection (dashed lines) and Cu fibres stretching out along the solidification direction, (c) closeup of the area framed in (b), showing fibres and their fragmentation to spherical Cu particles (selectively arrowed), (d) a layer similar to those dashed lined in (b), showing irregular and spherical Cu (bright) and dark Fe matrix, (e) fine Fe grains bounded by Cu fibres (arrowed), (f) Fe grains surrounded by Cu (arrowed), (g) higher resolution, showing finer Fe grains compared to those in (a) and (h,i) closeups of the boxed areas in (g) revealing (h) larger grains away from Cu fibres compared to (i) finer ones between them.
Upon solidification, liquid separation would occur in the Cu-rich area and Cu fibres form perpendicular to the layer, periodically separated by the Fe-rich layers. Such a structure is observed everywhere in all the melt pools. Closeups (Figure 1(c)) of the frame in Figure 1(b) reveals fragmentation of fibres into spherical particles (arrowed). The pattern of the Cu-rich phase (bright) in the Fe-rich matrix (dark) is in Figure 1(d), comparable to the microstructures from Fe-Cu liquid separation [3,18,36]. Further, Figure 1(e,f) shows nano/ultrafine Fe-grains between Cu fibres (arrowed). Figure 1(g–i) shows that the grains in Fe-Cu were significantly finer, mainly of the order of 100 nm. The grains confined by Cu fibres (Figure 1(i)) were much finer (< 200 nm) than those further away (D ~ 400 nm, Figure 1(h)).

EBSD in the Cu-rich region is shown in Figure 2. Figure 2(a,b) shows IPF-Y (Y is the build direction) maps for α-Fe and Figure 2(c,d) are the corresponding band contrast maps. The pole figures in Figure 2(e) revealed weak texture, unlike in other bcc-metals (e.g. β-Ti [25,26]) strongly textured along the build direction. Figure 2(f) shows the grain size distribution, revealing D of 250 nm with many in the nano-range of 50–100 nm and very few of the order of 1 μm (the median is ~ 140 nm). The aspect ratio distribution in Figure 2(g) shows mostly equiaxed grains. EBSD, therefore, confirms the formation of equiaxed grains, as previously reported [22, 33–35] and observed here (Figure 1(a)), and much smaller grains (by one order of magnitude) in Fe-Cu. It should be noted that the Cu fibres are too thin (< 30 nm) to detect by EBSD in Figure 2.

Figure 3(a,b) shows DF for Cu and BF TEM (Fe grains selectively delineated) with a closeup of the framed area on the right, revealing mostly equiaxed (some with aspect
Figure 3. (a–h) TEM on the as-SLM Fe-Cu: (a) DF for Cu, showing Cu fibres and equiaxed Fe grains surrounded by Cu and (b) corresponding BF (Fe grains are selectively delineated) with a closeup from the framed area on the right, (c,d) BF and DF for Cu from a cross section of Cu fibres, respectively, showing a network of Cu in Fe grain boundaries, and (e) STEM-EDS elemental map from one of the hexagons in (c) with green representing Cu and red Fe, (f,g) DF for Cu showing fine Cu particles of ∼ 5–10 nm, (h) HAADF-STEM showing continuous Cu in Fe grain boundaries with a high density of nano Cu particles in its vicinity. (i–l) TEM on the as-SLM Fe-Cu after compression, showing (i) dislocations at the Fe/Cu-fibre interface, (j,k) dislocations pinned by Cu particles (selectively arrowed) and (l) formation of dislocation loops around Cu precipitates (selectively arrowed).

The Fe-Cu exhibited a compressive yield strength (CYS) of ∼ 900 MPa, more than double that of ∼ 400 MPa for SLM-Fe. The ultimate compressive strength reached ∼ 1200 MPa at fracture strain ($\epsilon_f$) of ∼ 10%. Table 2 compares CYS of Fe-Cu with that of Fe produced using different methods. The CYS of Fe is ∼ 200–420 MPa remarkably smaller than that of Fe-Cu by up to ∼ 80%. $\epsilon_f$ of ∼ 10% was quite good considering the high strength.

4. Discussion

Two important observations can be made. First, a nano/ultrafine grained structure was produced using SLM, the finest obtained using AM. Second, thanks to the significant grain refinement and dispersion of nano-Cu particles, ultrahigh strength Fe-Cu with CYS of 900 MPa...
Figure 4. Schematic illustration of the microstructural evolution in Fe-Cu during SLM: (a) A mixture of Fe and Cu powders in a melt pool before melting, (b) the Marangoni convection leading to (c) formation of a layered structure of liquids, including Fe + Cu liquid (L0) in a near pure Fe liquid (L), (d) liquid separation in L0, creating Fe-rich (L1) and Cu-rich (L2) liquids, (e) formation of a fibrous Cu structure among αM upon solidification, showing a network (A-A section) originated from the monotectic reaction L1 → L2 + γ-Fe, followed by solidification of L2 into Cu-rich fibres and transformation of γ into αM, and (f) αM transformation to α and precipitation of nano-Cu particles upon αM → γ → α transformation, similar to the mechanism suggested for the formation of nano-grains in steels during cyclic heating through γ → αM → γ [39]. Another possibility, as assumed in an SLM-fabricated Fe [22], is that α solidifies directly from liquid without first forming δ and γ due to rapid cooling, and the subsequent reheating leads to the columnar-α → γ → equiaxed-α transformation. However, although the high ˙T might prevent L → δ, the suppression of L → γ is unlikely since γ is stable in a wide temperature range (912–1400°C). This is supported by a study [40] on a splat-quenched Fe showing the sequence of L → γ → αM at ˙T of ∼10^5°C/s comparable to ˙T in SLM. Since there are different variants of αM formed in the prior γ, it is possible that the epitaxial growth and columnar grain structure in SLM of other cubic metals [23–26] would not occur in Fe.

While equiaxed α-grains can similarly form through γ → αM → γ → α in SLM of Fe-Cu, important differences are noted. The primary γ in Fe-Cu is confined by Cu fibres (bottom in Figure 4(e)), leading to finer αM upon solidification. The subsequent heating cycles would convert αM to equiaxed α (Figure 4(f)) whose size is largely dependent on the spacing between Cu fibres in the range of 50–300 nm, much finer than in SLM-pure Fe. In contrast, grains were coarser in the ultrafine range in the region without restriction from Cu fibres (Figure 1(g–i), top in 4f). Moreover, Cu fibres and particles in a previously deposited layer might become nucleation sites for Fe solidification, facilitating grain refinement. An
increase in thermal conductivity by adding Cu, particularly when fibrous Cu forms [18], can introduce higher \( T \) and undercooling, further enhancing grain refinement [12,31]. Further, a gradient of Cu concentration from its rejection by the solidification front may result in constitutional undercooling, contributing to grain refinement. In Fe-Cu, \( \gamma \)-Fe undergoes the eutectoid reaction \( \gamma \)-Fe\(\rightarrow\)\(\alpha\)-Fe+\(\epsilon\)-Cu [19]. Further, under equilibrium the solubility limit of Cu in \( \gamma \)-Fe at elevated temperatures is \( \sim 14 \text{ wt.\%} \), rising to \( \sim 35 \text{ wt.\%} \) upon rapid cooling [3], but it reduces to almost zero at room temperature, leading to \( \epsilon \)-Cu precipitation. Therefore, during cyclic heating/cooling, \( \gamma \)-Fe\(\rightarrow\)\(\alpha\)-Fe transformation is accompanied by the formation of nano-sized (\( \sim 5 \text{ nm} \)) Cu particles (Figures 3,4(f)), via either the eutectoid reaction or precipitation. The concentration of these nano-Cu particles is higher near the continuous Cu at the grain boundary (Figure 3(f,h) and A-A section in Figure 4(e)). The network is formed as Cu is rejected from L1 during the monotectic reaction, leading to increasing Cu contents in solidifying \( \gamma \)-Fe towards the grain boundary and a high density of Cu particles there.

This microstructure in Fe-Cu enhanced strength thanks to dispersion strengthening by the nano-Cu particles and boundary strengthening by grain refinement. Based on the Orowan mechanism, the stress (\( \sigma_{\text{or}} \)) for a dislocation to bow between particles is [41]

\[
\sigma_{\text{or}} = M \cdot \tau_{\text{or}} = M \cdot \frac{0.84 \cdot G \cdot b}{l} \quad (1)
\]

where \( \tau_{\text{or}} \) denotes the shear stress, \( M \) Taylor factor (\( \sim 3 \) in polycrystalline cubic metals [41]), \( G \) shear modulus (79 GPa for Fe [33]), \( b \) Burgers vector (0.25 nm for Fe [33]), and \( l \) mean distance between particles. \( l \) is calculated by [42]

\[
l = \left( \frac{6 f}{\pi} \right)^{-\frac{1}{2}} \cdot d \quad (2)
\]

where \( f \) is the dispersed volume fraction, measured to be \( \sim 0.09 \) in the region with a high density of nano-Cu particles, and \( d \) diameter of the dispersoid (\( \sim 5 \text{ nm} \)), giving rise to \( l \approx 9 \text{ nm} \). Using eq.1, \( \sigma_{\text{or}} \) is 5.5 GPa. This high strength, however, applies only to the area near the Cu network with a high concentration of nano-Cu particles. In other words, this represents the strongest part of the grain. The number of Cu particles reduces quickly away from the boundary, becoming nearly Cu-free at the centre. The volume fraction (\( f' \)) of the region containing a high density of Cu nanoparticles (i.e. with \( \sigma_{or} = 5.5 \text{ GPa} \)) is estimated to be \( < \sim 0.2 \) from TEM (same area as in Figure 3(d)). If the rest of the grain is assumed to have the strength of pure-Fe (i.e. \( \sigma_F = \sim 400 \text{ MPa} \)), the overall strength in this Cu fibre-containing region (\( \sigma_{Fe-Cu} \) corresponding to the bottom part in Figure 4(f)) can be roughly estimated to be \( < \sim 1420 \text{ MPa} \) using the rule of mixture

\[
\sigma_{H-P} = 130 (\text{MPa}) + \frac{310 (\text{MPa} \cdot \mu \text{m}^2) + 130 (\text{MPa})}{\sqrt{d(\mu \text{m})}} \quad (3)
\]

where \( d \) is the grain size. Taking \( d = 0.4 \mu \text{m} \) (Figure 1(h)), \( \sigma_{H-P} = 620 \text{ MPa} \) is obtained. The fraction of the Cu-free region is estimated to be \( < \sim 20\% \) (Figure 1(b)). The total strength combining \( \sigma_{H-P} \) and \( \sigma_{Fe-Cu} \), again assuming the rule of mixture, would be (620 x 0.2)+(1420 x 0.8) = 1260 MPa, higher than the observed 900 MPa owing to the simplistic assumptions and roughly estimated volume fractions used. It nonetheless demonstrates the significant contributions from dispersion of nano-Cu particles and nano/ultrafine grain sizes to the much-enhanced strength in Fe-Cu. If the Cu-rich area with uniformly distributed Cu nanoparticles can be obtained in the entire alloy, yield strength of \( > 1.5-2 \text{ GPa} \) is achievable.

In summary, the SLM-fabricated Fe-Cu exhibited CYS more than double that of Fe thanks to the dispersion of 5 nm-Cu precipitates and ultrafine/nano grains. The formation of such a unique microstructure in the immiscible Fe-Cu system was attributable to rapid solidification and cyclic heating in SLM, preventing macro-segregation encountered in conventional processing and creating the nanostructure. The outcome demonstrates great potential for producing ultrahigh strength immiscible alloys using AM.

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**Disclosure statement**

No potential conflict of interest was reported by the author(s).

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