ABSTRACT

This work investigates whether the unique low thermal expansion property of Invar (64Fe–36Ni) is retained after processing using the additive manufacturing process selective laser melting (SLM). Using this process, near-full-density components (99.96%) were formed by melting thin (20 µm) layers of powdered Invar (15–45 µm particle size). The mechanical properties of SLM Invar were comparable to that of cold-drawn Invar\(^\text{36}\); however, the thermal coefficient of expansion was observed to be a lower value and negative up until 100 °C. This negative value was attributed to residual stress in the as-deposited parts. The low thermal expansion property of Invar was still maintained when processed using a non-conventional layer-based additive manufacturing technique.
components. The method of layered fabrication, combined with the high precision of laser melting, allows for a greatly expanded design freedom with minimal feedstock waste. It is increasingly being used in high value markets for the production of various aerospace, automotive and medical components.

This work builds on the work by Qiu et al. [3] who investigated the microstructure and properties of selective laser-melted Invar36®. Qiu reported that the as-processed microstructure comprised of columnar vertically orientated γ grains, interspersed by nanocrystalline α-phase. The as-processed material displayed anisotropic tensile behaviour, with specimens built in the horizontal orientation displaying superior yield strength and UTS to those built in the vertical; but elongation was superior in the vertical. This behaviour has been noted in other investigations for nickel-base alloys [4–6] processed by metal powder bed fusion. In addition, it was reported that SLM-fabricated Invar36® displayed the same low thermal expansion properties up to 300 °C as conventionally manufactured Invar36®. It was reported that there was a marginal reduction in thermal expansion after heat treatment when tested in the vertical orientation.

In this work, more focus is placed on the effect of the SLM process on the thermal expansion of Invar and similar alloys, and what influence the process characteristics, if any, have on the magnetovolume phenomenon. Analysis of microstructure and mechanical properties is also conducted for reference.

**Experimental methodology**

SLM processing parameters, such as laser energy density and beam velocity, were optimised for the fabrication of fully dense parts with minimal defects, and the physical properties of the resultant fabricated Invar components were investigated. Samples were fabricated on a Renishaw SLM125 using a modulated 200 W ytterbium fibre laser with a 125 × 125 × 125 mm build volume. The Invar powder feedstock (sourced from LPW Technology, UK) was manufactured using gas atomisation to attain a consistent spherical morphology and was sized to 15–45 μm with a composition of 63.7 wt% Fe, 35.8 wt% Ni, 0.47 wt% Mn and 0.032 wt% C.

In previous optimisation trials for Fe–Ni based alloys, it was established that a layer thickness of 20 μm and hatch spacing of 0.09 mm were optimal for achieving maximum density [7] and thus would be set as fixed parameters for the remainder of this investigation. The scan strategy used was a raster pattern with a 67° rotation after each layer, known as a ‘meander’ strategy; it is designed to maximise interlayer density and reduce thermal stress build-up within the parts. Parameter optimisation in this study focussed on laser power (LP), point distance (PD) and exposure time (ET). The laser scan motion on the Renishaw SLM125 is a ‘point-to-point’ traverse, where the laser path is comprised of a series of single exposure points (of time ET) separated by a point distance (PD), as opposed to a continuous laser exposure which moves at a ‘beam velocity’ along the full length of the scan path. PD and ET can be combined, with the addition of idle time, to give an apparent scan velocity. Using Design of Experiment (DOE) software Minitab®, an experimental plan was devised to create samples from a range of SLM parameters consisting of minimum and maximum values of 180–200 W, 90–150 μs and 50–90 μm for LP, ET and PD, respectively. The central point of 190 W, 120 μs and 70 μm was repeated three times for validation purposes. The density results from the initial experiment were input into Minitab®, and a second full-density solution parameter set was calculated.

Percentage density was optically calculated using micrographs of sectioned samples by taking area fraction measurements of the binary images on Image J software. Samples were mounted in conducting resin, ground and polished using SiC pads in grades from P800-P2500 and diamond suspension polish from 3 to 1 μm, to reveal polished vertical and horizontal sections. Two sections were prepared of each sample, with three micrographs taken for each section—allowing for calculation of mean values with standard error. This method is preferred over displacement techniques as it reveals the quantity and morphology of porosity that may exist within a sample. Samples for microstructural analysis were etched using a 2% Nital solution (98 ml of IMS with 2 ml of nitric acid). Tensile rounds were machined from cuboids fabricated in the x–y orientation, see Fig. 1, to ASTM A370 standard sizing. Tensile components built in x–y (built with the gauge length in the x–y plane, rather than z plane) have been found to display slightly higher tensile strength than those built in z [8]. Therefore, the x–y orientation represents a maximum tensile strength for the as-deposited
It is noted that the same investigation [8] found that elongation was greater in tensile components built in the z plane; therefore, x–y plane components represent a minimum for elongation. Ultrasonic analysis was conducted on as-deposited samples to determine Young’s modulus, independently of the tensile results, and was performed on an Olympus EPOCH 600 Ultrasonic Flaw Detector. Young’s modulus is calculated from a measured longitudinal wave sound velocity and known material density (8.105 g/cm³ as measured by helium gas pycnometry on a Micromeritics AccuPyc 1340 Pycnometer) and Poisson’s ratio (0.3) [2]. The samples were fabricated as 8 mm cubes, with the parallel surfaces ground to ensure sufficient contact, and measurements were taken four times per sample for two samples. Thermal expansion analysis was conducted on a Perkin-Elmer Diamond TMA; the CTE measurements were taken on the same 8 mm cubes from the ultrasonic measurements. Testing was conducted for both x–y and z orientations, see Fig. 1, as per ASTM standard E831 using a heating rate of 5 K per min, with three cycles per sample. The sensitivity of the device is 0.02 μm.

**Results and analysis**

Power, exposure time and point distance are all controlling variables of laser energy, either per unit time or unit length. Plotting density against one of the three is nonsensical as it neglects the influence of the other two. As such, LP, ET and PD have been dimensionally reduced into a single parameter of 1D line energy density. This is an appropriate use of energy density since layer thickness and hatch spacing are fixed; thus, any variation in density is purely a consequence of the absorbed energy. If the energy density is considered over a set length, it can be described as Energy per point × Points per unit length or $P \times ET \times 1/PD$, and this becomes:

$$\frac{Pt_{exp}}{x_{PD}} = \frac{Q}{l}$$

where $P$ is the power, $t_{exp}$ is the exposure time, $x_{PD}$ is the point distance, $Q$ is the energy, and $l$ is the unit length. Note this relationship was first used in a previous publication by Harrison et al. [7].

Figure 2 indicates a strong relationship between part density and 1D line energy density, with a global maximum located between 0.35 and 0.40 J/mm. The highest observed density was 99.96% with a 1D energy density of 0.33 J/mm, of which the scan parameters were LP 190 W, ET 120 μs and PD 70 μm. There were 10 samples which achieved densities ≥99.5%.

Energy densities which fail to produce samples with density of ≥99.5% have been split into two regions according to the type of porosity that was observed (shown in Figs. 2, 3). Lack of fusion occurs when the energy density is not sufficient to generate full melting of the powdered layer, leading to pockets of unmelted particles and, in extreme cases, delamination between previously processed layers. In cases of high energy densities, the generated surface temperature can exceed the evaporation temperature of the alloy, leading to vapourisation recoil which ejects particles and molten material from the heat-affected zone resulting in large irregular voids. Gas may also be trapped within these voids leading to large spherical pores.

In Fig. 4, it is observed that the grains have grown epitaxially and transcend the layers, with vertical grain length being in the order of 200–300 μm. The grain structure is columnar and orientated in the build z-direction and is in agreement with that reported in other investigations [9–13], including Qiu et al. [3]. The meander scan strategy used in this investigation has resulted in more grain intersections than if processed using a raster pattern without rotation, which would have allowed for more continual epitaxial growth. Chen et al. [14] describe how crystallographic orientation selection is influenced by the scan direction-dependent heat flux and that the
epitaxial growth is dependent on the angle between the grain growth and (scan direction-dependent) heat flux with high angles likely to result in stoppage due to the grain impinging on another and low angles more readily continuing to the track surface. The meander strategy likely increases the chance high angles being formed due to the rotation of scan direction after each layer. The result is a less ordered grain structure, made up of large columnar grains interspersed with small more irregular grains. This structure was also observed in an investigation by Harrison et al. [7] where the same meander scan strategy was used on a nickel-base superalloy. The dark arcs observed in Fig. 4 are the melt traces, or base of the individual melt pools, which have been highlighted by the etching process indicating minor solute segregation.
Table 1 displays the results of the tensile testing of SLM Invar samples in comparison with the literature values for cold-rolled and annealed Invar36\textsuperscript{®}. Both UTS and offset yield strength of the SLM-fabricated Invar exceed that of the annealed, which is consistent with other findings for solid solution alloys \cite{7, 15, 16} and is more comparable to cold-drawn Invar36\textsuperscript{®}. The mechanical properties are also consistent with those reported by Qiu et al. \cite{3} for the same conditions. Elongation is marginally reduced for SLM Invar over annealed; however, the reduction of area is greater, implying fewer internal defects.

Figure 5 displays the linear expansion curve of SLM Invar for a temperature range of 30–300 °C, as well as those for stress-relieved Invar samples which will be discussed in “Observed reduction of thermal expansion coefficient” section. The values have been normalised against the room temperature dimension of each sample for fair comparison. Table 2 gives the mean coefficient of thermal expansion \( \alpha_{\text{CTE}} \) for five temperature ranges, as well as comparison values for annealed Invar36\textsuperscript{®}.

It is noted from Fig. 5 that the normal linear thermal expansion begins at approximately 240 °C for both orientations; this is taken as the temperature at which the magnetovolume phenomenon is fully overcome by atomic vibrations. However, this does not indicate a conflicting Curie point to that of the literature, and it is merely the temperature at which the phenomenon’s effect becomes negligible.

Temperature ranges for \( \alpha_{\text{CTE}} \) measurements shown in Table 2 were chosen for the best comparison with the literature values. The final temperature range represents room to Curie temperature.

**Observed reduction of thermal expansion coefficient**

In comparison with Invar36\textsuperscript{®}, as-deposited SLM Invar displays significantly lower CTE values for all temperature ranges and negative CTE for 30–100 °C. The initial consideration for this variance was
differences in chemical composition; however, the reported chemical compositions of LPW-Invar and Invar36°/[2] are similar [2]. In addition, the CTE of an alloy is only sensitive to changes in concentration of ferromagnetic elements Fe, Ni and Co, but no quantitative relationships for this sensitivity exist. Given that the elemental constituents of the LPW-Invar do not vary from the standardised composition—i.e. additions of Co—and that any deviations from the 64Fe–36Ni form would reduce the Invar effect rather than boost it [1], this is again not considered to be a significant factor in the observed reduction of thermal expansion.

It was also considered that the unique microstructure of as-deposited SLM samples may have an effect on the thermal expansion. The thermal expansion of a crystal must possess the symmetry of the crystal [19], and therefore, cubic crystals, like the fcc of Invar, will exhibit isotropic thermal expansion in all crystallographic directions. It is therefore expected that, without external influence, a grain of Invar dendrites will expand homogenously and \( \alpha_{\text{CTE}} \) will be the same regardless of its shape or size. If grain morphology did have an effect, dramatic variation between x-y and z measured \( \alpha_{\text{CTE}} \) would be observed because of the high aspect ratio grains and columnar dendrites. Although a disparity has been observed, it is only marginal and therefore unlikely to be as a result of grain structure.

An investigation by Wang et al. [20] demonstrated that residual stress anisotropy can lead to effective

| Table 2 | Mean coefficient of thermal expansion \( \alpha_{\text{CTE}} \) for SLM-fabricated Invar, Invar 36°/[2], and stress-relieved SLM-fabricated Invar |
| Temp (°C) | 30–100 | 30–150 | 30–200 | 30–260 | 30–279 |
| SLM z | −0.355 | 0.164 | 0.889 | 2.77 | 3.56 |
| SLM x–y | −0.471 | 0.098 | 0.767 | 2.58 | 3.34 |
| Invar 36° | 1.6° | 2.0 | – | 4.1 | – |
| SLM stress-relieved z | 1.00 | 1.34 | 1.85 | 2.96 | 3.68 |
| SLM stress-relieved x–y | 0.70 | 0.99 | 1.53 | 2.74 | 3.36 |

*a For temperature range 25–93 °C

Units are \( (10^{-6}/ {\degree}C) \); uncertainty of measurement is ±0.01 \( 10^{-6}/ {\degree}C \)

| Table 3 | Mean coefficient of thermal expansion \( \alpha_{\text{CTE}} \) for SLM-fabricated stainless steel 316l (316l), Hastelloy X (Hast X) and literature values of the same alloys fabricated from conventional techniques [22, 23] |
| Temp (°C) | 30–100 | 30–200 | 30–300 | 30–400 | 30–500 |
| SLM 316l z | 17.27 | 17.48 | 17.76 | 17.93 | 18.17 |
| SLM 316l x–y | 16.63 | 17.21 | 17.51 | 17.64 | 17.57 |
| Cold-rolled 316l | 16 | 16.5 | 17 | 17 | 18 |
| SLM Hast X z | 12.87 | 13.57 | 13.98 | 14.28 | 14.55 |
| SLM Hast X x–y | 12.59 | 13.52 | 14.02 | 14.35 | 14.65 |
| Sheet Hast X | 13 | 13.5 | 13.8 | 14.5 | 14.5 |

Units are \( (10^{-6}/ {\degree}C) \)
CTE anisotropy. This could not only be the cause of the disparity but also potentially explain the observed negative, and or low, expansion of the SLM-fabricated samples.

To investigate this further, as-deposited SLM samples were stress-relieved by heat treatment. The samples were held at 850 °C for 1 h under an argon atmosphere to prevent oxidation. After the heat treatment, the thermal expansion analysis was conducted again for both x–y and z orientations. The stress relief resulted in an overall increase in thermal expansion coefficient for the samples shown in Table 2; however, the disparity between the test orientations increased in magnitude; see also Fig. 5 and Table 2. This implies that the residual stress was contributory to the exceptionally low CTE values in the as-deposited samples, but was not responsible for the anisotropy.

This behaviour is not in agreement with Qiu et al. who reported very similar $\alpha_{\text{CTE}}$ values for as-deposited and heat-treated material. However, Qiu employed a different heat treatment regime of multiple exposures with reducing temperature and increasing dwell time, and critically, the initial stress relieving cycle held the maximum temperature (830 °C) for half of the duration (0.5 h) compared to this work (1 h). The lower dwell time may have not been sufficient to fully stress-relieve the material, and therefore, the atomic movement was still being restricted.

To investigate whether reduction of thermal expansion occurred in other alloy systems when processed by SLM, as-deposited samples of stainless steel 316l (316l) and Hastelloy X were analysed. Thermal expansion analysis was performed for both x–y and z orientations, as with Invar; see Fig. 6. The values were then compared to those values for conventionally manufactured equivalents; see Table 3. Unlike Invar, the 316l and Hastelloy X samples did not display significantly different $\alpha_{\text{CTE}}$ than that reported for conventional manufacturing techniques. Although not quantified, it is expected that both alloys will be under significant residual stress in the as-deposited state [21] and likely that this residual stress will be greater than that for Invar, given their comparative yield strengths. The implication is therefore that although stress-relieved Invar does display a higher CTE than that of the as-deposited material, the residual stress does not affect the thermally driven expansion of the lattice but rather the magnetovolume contraction. Observations of the thermal expansion curves in Fig. 5 support this, where, beyond the range of the Invar effect’s influence ($T > 280 ^\circ\text{C}$), the as-deposited and stress-relieved Invar expand at the same rate.

A consistent disparity between the test orientations was observed for 316l but not Hastelloy X. In Hastelloy X, the z orientation displays the larger $\alpha_{\text{CTE}}$ values up to 300 °C, after which point the x–y orientation measurements become greater. Most notable though is that the difference in $\alpha_{\text{CTE}}$ between orientations for all three alloys does not vary greatly from $0.5 \times 10^{-6}/\text{°C}$. The implication is therefore that the disparity is a process-related phenomenon, and is not specific to Invar.

The only other feature which varies specifically with orientation is defect population. Given the fine columnar grain structure and fabrication through layering of powder, it is plausible that defects such as micropores and microcracks will be aligned with either scan direction or grain orientation. Even if the expansion of the individual defect is isotropic, a preferential concentration of defects in a particular orientation will result in bulk anisotropy. This would also explain the small absolute magnitude of disparity and why it remains consistent with increasing temperature and not affected by the magnetovolume Invar effect.

**Conclusions and summary**

Full-density (>99.5% dense) components of Invar were fabricated after optimisation of laser scan parameters. The optimised parameters were then used to build tensile and thermal expansion test components. SLM-fabricated Invar displayed tensile properties comparable to that cold-drawn, and superior to annealed, Invar36®.

As-deposited SLM-fabricated Invar was shown to have a lower CTE than conventionally manufactured Invar36®. In addition to this, a small but consistent anisotropy in the CTE was observed between the x–y and z test orientations for the as-deposited SLM Invar samples. After considering the effects of microstructure and residual stress, the anisotropy was attributed to heterogeneous distribution of defects within the as-deposited components.

Residual stress was found to be a significant contributor to the reduced CTE of SLM-fabricated Invar.
However, comparisons of thermal expansion behaviour between other Fe–Ni alloy systems, and stress-relieved Invar samples, implied that the influence of process-induced residual stress was not universal. Rather, it affected the magnetovolume contraction (Invar effect) and not the thermally induced expansion of the lattice.

To summarise, SLM-fabricated Invar displays comparable tensile properties to those fabricated by conventional processes. The SLM process does not affect the low thermal expansion properties of Invar in a negative way; instead, it produces a further reduced thermal expansion coefficient for atmospheric temperatures.

Acknowledgements

The authors would like to thank LPW Technology for their collaboration and material supply within this project. This work was supported by LPW Technology and the EPSRC (Grant Number EP/K503149/1).

Compliance with ethical standards

Conflict of Interest The authors declare that they have no conflict of interest.

Open Access This article is distributed under the terms of the Creative Commons Attribution 4.0 International License (http://creativecommons.org/licenses/by/4.0/), which permits unrestricted use, distribution, and reproduction in any medium, provided you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons license, and indicate if changes were made.

References

[1] Rancourt DG, Dang MZ (1996) Relation between anomalous magnetovolume behavior and magnetic frustration in Invar alloys. Phys Rev B 54(17):12225–12231
[2] Carpenter Invar 36 Alloy: Technical Datasheet. (2014), Carpenter Technology Corporation
[3] Qiu C, Adkins NJE, Attallah MM (2016) Selective laser melting of Invar 36: microstructure and properties. Acta Mater 103:382–395
[4] Kanagarajah P et al (2013) Inconel 939 processed by selective laser melting: effect of microstructure and temperature on the mechanical properties under static and cyclic loading. Mater Sci Eng, A 588:188–195
[5] Bauer T et al (2013) Microstructure and mechanical characterisation of SLM processed Haynes® 230®. J Chem Inf Model 53:1689–1699
[6] Etter T et al (2015) Reduction in mechanical anisotropy through high temperature heat treatment of Hastelloy X processed by selective laser melting (SLM). IOP Conf Series Mater Sci Eng 82(1):012097
[7] Harrison NJ, Todd I, Muntaz K (2015) Reduction of microcracking in nickel superalloys processed by selective laser melting: a fundamental alloy design approach. Acta Mater 94:59–68
[8] Rickenbacher L (2013) High temperature material properties of IN738LC processed by selective laser melting (SLM) technology. Rapid prototyp J 19(4):282–290
[9] Wang Z et al (2012) The microstructure and mechanical properties of deposited-IN718 by selective laser melting. J Alloy Compd 513:518–523
[10] Vilaro T et al (2012) Microstructural and mechanical approaches of the selective laser melting process applied to a nickel-base superalloy. Mater Sci Eng A 534:446–451
[11] Kempen K et al (2011) Microstructure and mechanical properties of selective laser melted 18Ni-300 steel. Phy Procedia 12:255–263
[12] Amato K (2012) Comparison of microstructures and properties for a Ni-base superalloy (Alloy 625) fabricated by electron beam melting. J Mater Sci Res 1(2):3–41
[13] Amato KN et al (2012) Microstructures and mechanical behavior of Inconel 718 fabricated by selective laser melting. Acta Mater 60(5):2229–2239
[14] Chen ZW, Phan MAL, Darvish K (2017) Grain growth during selective laser melting of a Co–Cr–Mo alloy. J Mater Sci 52(12):7415–7427
[15] Wang F (2011) Mechanical property study on rapid additive layer manufacture Hastelloy® X alloy by selective laser melting technology. Int J Adv Manuf Technol 58(5-8):545–551
[16] Savitha U et al (2012) Effect of process parameters on porosity in laser deposited IN625 alloy. Trans Indian Inst Met 65(6):765–770
[17] CES Edupack 2014. Granta Design Limited
[18] Hull FC et al (1987) Effect of composition on thermal expansion of alloys used in power generation. J. Mater Eng 9(1):81–92
[19] Nye, J.F., Physical properties of crystals: their representation by tensors and matrices. 1979, Oxford: Clarendon Press. xiii
[20] Wang XL et al (1999) Influence of residual stress on thermal expansion behaviour. Appl Phys Lett 75(21):3294–3296
[21] Mercelis P, Kruth J-P (2006) Residual stresses in selective laser sintering and selective laser melting. Rapid Prototyp J 12(5):254–265
[22] Inc., H.I. Haynes international Inc—Hastelloy® X alloy datasheet. (1997); Available from: http://www.haynesintl.com/pdf/h3009.pdf
[23] BSI, BS EN 10088-1:2014—Stainless steels. List of stainless steels. 2014, BSI