High Strain Rate Behaviour of Nano-quasicrystalline Al93Fe3Cr2Ti² Alloy and Composites

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Abstract

We demonstrate the outstanding dynamic strength of nano-quasicrystalline $\text{Al}_3\text{Fe}_3\text{Cr}_2\text{Ti}_2$ at.% alloy and composites. Unlike most crystalline Al alloys, this alloy exhibits substantial strain rate sensitivity and retains ductility at high strain rates. This opens new pathways for use in safety-critical materials requiring impact resistance.

Keywords: Aluminium alloys, Quasicrystals, Dynamic testing, Hopkinson bar, Metal matrix composites

Main

Quasicrystals were first observed in rapidly solidified Al-Mn-(Fe/Cr) alloys by Daniel Shechtman (*1*), who received the 2011 Nobel prize for chemistry for the discovery of this new state of matter (*2*). Since their discovery, quasicrystals have been observed in a variety of systems including laboratory-made binary (*3*), ternary (*4*), quaternary alloys (*5*), soft matter (*6*) and naturally occurring minerals (*7*, *8*). Quasicrystals have shown extremely versatile functional properties due to the ordered, non-translational nature of their quasi-periodic lattice (*9*), making them suitable for an incredibly wide range of applications such as superconductivity (*10*), photonics (*11*–*13*), coatings (due to a low-friction coefficient along aperiodic axes) (*14*), as well as outstanding mechanical strength under quasi-static loading conditions (*15*) and thermal stability up to 75% of the alloys homologous melting temperature (*16*). Studies have analysed the active deformation mechanisms under quasi-static loading that result in excellent mechanical response at both ambient temperature (*17*) and high temperatures (*18*), while no work has been performed to date on the response of these materials at high strain rates.

We studied the dynamic mechanical performance of a quaternary quasicrystalline Al93Fe3Cr₂Ti₂ at.% alloy (hereafter denoted QC) and the composites obtained by mixing 10 and 20 vol.% pure Al fibres into a matrix of QC alloy (referred to as QC-10 and QC-20 respectively). Rapid solidification is required for quasicrystals production in the selected alloy (atomisation has been previously shown to produce a nano-quasicrystalline structure (*17*), while spray forming produced a microstructure made of other nano-scale intermetallic phases (*19*)). The alloy atomisation conditions, consolidation methods and influence of processing parameters are available in a previous publication (*17*). X-ray diffraction (XRD) was performed for the identification of bulk phases, using a Philips 1810 θ–2θ diffractometer with Cu-Kα radiation, (λ =1.5406Å). Diffractograms were collected over 20 angles from 20 to 100° (scattering vector Q between $1.4 - 4.0 \text{ Å}^{-1}$) with a tube voltage of 35 kV, current of 50 µA, scanning step size 0.02°. Samples were prepared by grinding extruded bars into powders to remove texture effects and ease phase identification. Diffractogram peaks were indexed using a combination of data sheets (*20*) and published papers (*16*). Scanning Electron Microscopy (SEM) was performed on a JEOL 6500F microscope. A working distance of 15 mm was used, 300 pA and a voltage of 30 kV. Samples were prepared by cutting longitudinal sections of the extruded bars, grinding and polishing them using increasingly finer grades of diamond and finally 0.04 m colloidal silica. Transmission Electron Microscopy (TEM) was performed on a Philips CM20 W-filament microscope. Samples were prepared with a Gatan dimple grinder and Gatan Precision Ion Polishing System. Convergent beam diffraction (CBD) was performed using the smallest spot sizes, which when converged would allow diffraction patterns to be taken from microstructural features as small as ~40 nm. Manual measurements of particle sizes were performed on bright-field images for maximum accuracy over ~120 measurements, which were then plotted as a graph and fitted to a Gaussian distribution. Results are quoted as mean value \pm full-width-half-maximum (FWHM) of the distribution. Energy dispersive X-ray spectroscopy (EDX) was used to measure chemical compositions of quasicrystals and their surrounding matrix with an Oxford Instruments EDX detector. Values quoted are mean \pm standard deviation obtained over 10 measurements in randomly distributed locations on the 3mm TEM disc.

Tensile and compressive quasi-static tests were performed at a strain rate of 10^{-4} s⁻¹ at room temperature. Tensile quasi-static tests were performed by Westmoreland plc using an Instron servo-hydraulic mechanical testing machine and measuring strain by means of an extensometer. Compressive quasi-static tests were performed in-house using an Instron servo-hydraulic testing machine in displacement control. In compression cylindrical specimens were used, of diameter 2 mm and height 2 mm. The number of tests was limited by the volume of material available but at least 2-4 repeats of each test were performed. Quasi-static tests were highly repeatable, with scatter less than 1%. Dynamic tests were performed using a split-Hopkinson bar. Test pieces were marked using evenly spaced markers and the tests were filmed by means of a high-speed camera. Strain was measured through digital image correlation (using the marks as references). Between 2-4 repeats of each test were performed and representative graphs were selected and displayed. Dog-bone shaped threaded round-section samples of ASTM standard size E8/E 8M sub-size 4 (diameter 4 mm, gauge length 16 mm) were used in tension. Dynamic equilibrium in the tensile tests was achieved only in a small number of experiments, and only just prior to the specimen failure; for this reason the measured tensile stress versus strain curves are not presented, but only the measured tensile strength is reported; the average strain rates indicated are obtained by averaging the measured strain rate between the point of dynamic equilibrium and the time at which peak stress is achieved. Specimens of diameter 3 mm x 6 mm were used in compression. During dynamic compression tests, dynamic equilibrium was achieved towards the end of the elastic response.

Detailed microstructural characterisation was performed. Phase identification within samples of the QC alloy and QC-10 and QC-20 composites was performed by X-ray diffraction (XRD), displayed in Figure 1. The α-Al peaks corresponding to the (111), (200), (220) and (311) were indexed (*20*) alongside the icosahedral phase, with reflections visible at angles (16) $2\theta \sim 23^{\circ}$, 41°, 43°, 62°, 74° (scattering vectors Q=1.6, 2.7, 2.8, 3.6 and 3.9 Å⁻¹), indicating that the overall microstructure was constituted solely of those two phases (or that if other phases were present, their volume fraction did not exceed the XRD detection limit of $~4\%$) after extrusion. Before extrusion, an additional peak of the metastable θ -Al₁₃(Cr, Fe)₂₋₄ was detected. This phase has a distorted monoclinic crystal structure, produced by rapid solidification. Due to overlaps only one reflection is observed, therefore unambiguous determination of this phase is difficult. However, its presence is expected in small volume fractions with its strongest peak ($\overline{820}$) overlapping with the FCC-Al (200) peak, the second strongest reflections the $(\overline{4}24)$ and (331) overlapping with two icosahedral peaks. The $(1\bar{1}31)$ reflection in the position observed here, so the highest intensity peak which does not overlap with other reflections. Following extrusion the reflection from the θ-Al₁₃(Fe,Cr)₂₋₄ phase can no longer be distinguished. Peak intensity comparisons were performed on the α -Al peaks, comparing them to an untextured powder sample, indicating that the thermo-mechanical production of all samples through extrusion induced a ~60% texture along the (220) direction (preferred grain orientation), which is itself an unusual finding for an aluminium alloy. This was measured by peak-intensity comparison, using the textures bar and untextured powder produced by grinding the bar.

Figure 1: X-ray diffractogram of the QC, QC-10 and QC20 samples confirming the presence of icosahedral phase, α-Al and the same phases were present in the atomised powders before extrusion, with the exception of θ *which decomposed during extrusion.*

Figure 2 (a-c) shows SEM micrographs of longitudinal sections of the extruded QC, QC-10 and QC-20 bars, revealing a microstructure made of deformed well-bonded powder particles, with no evidence of residual porosity remaining after extrusion. Some randomly distributed pure Al contamination was detected in the QC sample, which likely originates from the standard industrial practice of cleaning the atomiser with pure Al in between production of different alloys. This contamination was quantified at ~0.35 vol.% by synchrotron X-ray tomography (*17*), which confirmed SEM-based observations. The extruded microstructure of the QC-10 and QC-20 composite bars consists of a quasicrystalline alloy matrix with 10-20 vol% pure Al fibres. Fibres are made from extruded $\langle 25 \rangle$ um diameter pure Al powder particles, which during extrusion were deformed into fibres $< 200 \mu m$ in length. The sample longitudinal section and cross section show a random distribution of fibres within the composites, with no observed powder clustering. Grain size and texture was previously measured by EBSD (17) giving a mean planar grain diameter of d_s = 0.97 ± 0.3 µm (mean \pm standard deviation).

Figure 2: (a) backscattered low magnification overview of the QC bar, (b) the QC-10 bar, (c) the QC-20 bar, (d) bright field TEM micrograph and CBDP showing the characteristic 5-fold symmetry of the icosahedral phase.

TEM micrographs of the icosahedral quasicrystalline particles and their respective 5-fold symmetry convergent beam diffraction pattern (CBDP) are shown in Figure 2d. A bimodal distribution of well-dispersed icosahedral quasicrystalline phases was identified with mean spherical radii $r_1=43 \pm 6$ nm and $r_2=129 \pm 6$ nm. The volume fraction of quasicrystals in the FCC-Al matrix was 0.41, slightly lower than the 0.45 value measured by Inoue and Kimura, who produced the same alloy by melt spinning instead of atomisation (*21*). Quasicrystal composition measured by EDX was $Al_{87.8}Fe_{4.6}Cr_{4.2}Ti_{3.4}$ at.%. The composition of the α -Al matrix between quasicrystalline particles was Al_{98.1}Fe_{0.3}Cr_{0.3}Ti_{1.3} at.%. All matrix values are higher than the equilibrium solid solubility of each element in an α -Al matrix, particularly titanium (*22*), a likely result of the rapid solidification production route.

Figure 3: compressive true stress-strain curves of the QC alloy, QC-10 and QC-20 composites under quasi-static $(1x10³ s⁻¹)$ and dynamic $(3x10³ s⁻¹)$ loading conditions.

Figure 3 shows compressive true stress-strain curves of the as-extruded QC alloy and QC-10 and QC-20 composites at strain rates ranging from 10^{-4} s⁻¹ to 2840 s⁻¹, with a summary of all tests performed shown in Table 1. In compression, the QC alloy strength was 0.74 ± 0.10 GPa at slow strain rate (10^{-4} s^{-1}) and 0.90 ± 0.25 GPa at fast strain rates $(-3x10^{3} \text{ s}^{-1})$ ¹). Strength of the QC alloy is strain-rate dependent, increasing by ~20% from quasi-static to dynamic loading conditions, though the dynamic ductility, of order 5-6%, was reduced compared to the quasi-static case (10-20%). Localised plastic deformation initiated along shear bands inclined at approximately 45° from the direction of loading, corresponding to a plateau in the true stress-strain curves, shown in Figure 3. The shear bands width progressively increased to 2-3 mm and quickly evolved into a macroscopic crack, causing ultimate failure of the specimen (see insert of Figure 3). This behaviour was observed both during quasi-static and dynamic tests, though the width of shear bands was narrower after dynamic tests.

Test #	Type (T/C)	Material	Strain rate, /s	Strength, MPa
1	C	QC	0.0001	740
$\overline{2}$	C	$QC-10$	0.0001	690
3	C	$QC-20$	0.0001	626
4	T	QC	0.0001	660
5	T	$QC-10$	0.0001	605
6	T	$QC-20$	0.0001	530
7	C	QC	1770	926
8	C	QC	2840	878
9	T	QC	1490	943
10	T	QC	2720	998
11	C	$QC-10$	1620	856
12	T	$QC-10$	1530	822
13	C	$QC-20$	1750	689
14	T	$OC-20$	1740	754

Table 1: Summary of all tests performed in this study and relevant measurements.

In tension, the strength of the QC alloy was 0.66 ± 0.1 GPa at slow strain rate (10^{-4} s^{-1}) and 0.97 ± 0.2 GPa at fast strain rates $(3x10^3 s^{-1})$. Strain-rate dependency was more marked in tension, with a strength increase of 32% from quasistatic to dynamic loading conditions, though the dynamic ductility, of order 5-6%, was substantially reduced compared to the quasi-static case (10-20%). A summary of all collected mechanical data is shown in Table 1. In quasistatic tension, the QC alloy displayed an initial hardening phase followed by brittle fracture along a plane perpendicular to the loading direction, at plastic strains up to ~6%. In dynamic tension, the materials response was brittle and specimens failed by a transverse crack at the end of the elastic response, showing only limited plastic deformation and no visible necking. Repeated dynamic tests revealed wide scatter in the measured strength, of the order ±0.05 GPa. Failure was occasionally initiated by large pores, consequences of imperfect manufacturing. The moduli of the three materials were approximately insensitive to the direction of loading (tensile or compressive), though the compressive strength marginally outperformed the tensile strength independently of the strain rate. As expected from the conventional rule of mixtures, strength decreased linearly with increasing volume fraction of fibres, though the ductility and toughness required at least 20 vol% fibres in order to improve.

Figure 4: Specific strength as a function of the imposed strain rate; the tensile and compressive responses of the Al93Fe3Cr2Ti² *alloy and its composites in comparison with the compressive strength of commercial high-strength AA7075-T6* (*23*)*, AA7108-T6* (*24*) *and Ti6Al4V* (*23*) *alloys for dynamic applications.*

High strain rate tests were performed and the dynamic flow stress in tension measured for the pure alloy bar was ~ 1.00 GPa, while the composites showed 0.82 GPa (10 vol.% Al) and 0.75 GPa (20 vol.% Al). When divided by the density, their specific strength was measured to be 0.35 GPa/gcm^{-3} , 0.28 GPa/gcm^{-3} and 0.26 GPa/gcm^{-3} respectively. Values for commercially available Al alloys tested at similar strain rates were obtained from the literature and used as comparison, as shown in Figure 4. Tests were found on 7000 series Al alloys performed by Chen *et al.* (*25*) at 1000s-1 strain rate. They found specific dynamic flow stresses of 0.10-0.18 GPa/gcm^{-3} . Reyes et al performed tests at $1300s^{-1}$ strain rate and found that 7000-series alloys had a specific dynamic flow stress of 0.14-0.15 GPa/gcm⁻³ (24). Ti6Al4V tested at $3500s^{-1}$ and had a specific flow stress of $0.42 \text{ GPa/gcm}^{-3}$ (23). Cited data is summarised in Figure 4. While commercially available Al-based alloys display negligible strain rate sensitivity, the QC alloy under investigation becomes substantially stronger when tested at high strain rates. The dynamic strength of the alloy and its composites is 100-200% higher than that of commercial Al-alloys, and not far from that of the Ti6Al4V alloy, shown for comparison.

Several mechanisms could be causing the substantial strain rate sensitivity we observed in this aluminium-based alloy, either single-handedly or combined. The Al-Fe-Cr-Ti alloy studied notably has nano-scale grains (<400 nm) therefore the mean free path that each dislocation can cover dislocation motion will be substantially restricted, as will be their ability to cross slip (*17*). This effect will have increased the strength of the material and reduced the ductility, however, strain rate sensitivity is, by nature, time dependent. The introduction and the severity of this timedependency leads us to hypothesise that this effect is likely to be a combination of diffusion and restricted dislocation motion. Some of our previous work has shown that the composition of the α -Al matrix in which the quasicrystalline particles are found contains high levels of Fe, Cr and Ti in solid solution, namely 0.2 at% Fe, 0.3 at% Cr and 1.3 at% Ti (*17*). These values far exceed the nominal equilibrium solid solubility of these elements, which are 0.02 at% Fe, 0.18 at% Cr and 0.032 at% Ti (*22*). The excess of solutes in solid solution has been shown to interfere with dislocation motion by pinning, in one of our other publications (*18*). The pinning effect arises when excess solutes in solid solution segregate in the low-energy position at the dislocation core, creating an energy barrier that needs to be overcome before the dislocation can move. It is possible that as the strain rate is increased, dislocations do not have

time to become un-pinned, leading to the substantial loss in ductility and substantial increases in strength. However, not all the ductility is lost and the outstanding impact strength of the QC alloy and composites open new applications for safety-critical materials for impact resistant engineering components.

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Data Availability

The data that support the findings of this study will be made available upon reasonable request to the authors. A preprint of this publication is available on ArXiv.

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