Rejuvenation of metallic glasses by non-affine thermal strain

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When a spatially uniform temperature change is imposed on a solid with more than one phase, or on a polycrystal of a single, non-cubic phase (showing anisotropic expansion-contraction), the resulting thermal strain is inhomogeneous (non-affine). Thermal cycling induces internal stresses, leading to structural and property changes that are usually deleterious. Glasses are the solids that form on cooling a liquid if crystallization is avoidedthey might be considered the ultimate, uniform solids, without the microstructural features and defects associated with polycrystals. Here we explore the effects of cryogenic thermal cycling on glasses, specifically metallic glasses. We show that, contrary to the null effect expected from uniformity, thermal cycling induces rejuvenation, reaching less relaxed states of higher energy. We interpret these findings in the context that the dynamics in liquids become heterogeneous on cooling towards the glass transition¹, and that there may be consequent heterogeneities in the resulting glasses. For example, the vibrational dynamics of glassy silica at long wavelengths are those of an elastic continuum, but at wavelengths less than approximately three nanometres the vibrational dynamics are similar to those of a polycrystal with anisotropic grains². Thermal cycling of metallic glasses is easily applied, and gives improvements in compressive plasticity. The fact that such effects can be achieved is attributed to intrinsic non-uniformity of the glass structure, giving a non-uniform coefficient of thermal expansion. While metallic glasses may be particularly suitable for thermal cycling, the non-affine nature of strains in glasses in general deserves further study, whether they are induced by applied stresses or by temperature change.

In glasses there are no equivalent atoms, and elastic deformation must be non-affine. For metallic glasses this has been explored through atomistic simulations^{3,4} and diffraction-based measurements^{5,6} of the pair distribution function (see, for example, the review in ref. 7). Compared to their crystalline counterparts, metallic glasses have lower elastic moduli, associated with local atomic rearrangements occurring even well within the nominally elastic regime⁸. While these rearrangements (shears) contribute to apparently elastic strain, they are not individually reversible⁷, and there is net structural change. In atomistic simulations, shear events are localized in \sim 1 nm soft spots with low-frequency vibrational modes⁴, and can be associated with the range of density and stiffness of nearest-neighbour clusters (that is, the shell around a central atom)^{3,4}.

Measurements of metallic glass properties do not have such fine spatial resolution, but non-uniformity is still clearly detected. Strongly and weakly bonded regions have been postulated to account for \sim 5 nm zones of accelerated crystallization⁹. The initial yield load (in nano-indentation) shows a dispersion of values associated with structural heterogeneity¹⁰. Mapping of heterogeneity by atomic-force microscopy, over areas much larger than in simulation, shows variations of

energy dissipation 11 and of elastic modulus ($\sim\!30\%)^{12}$ with correlation lengths of 2.5–20 nm.

In metallic glasses under elastic loading, non-affine strains lead to local structural change; this led us to question whether thermal strains show analogous effects. In elastic deformation, the effects of non-affine strain and modulus variation are strongest for shear. For thermal expansion/contraction of a glass, however, the macroscopic strain is hydrostatic. In simulations of amorphous iron, the local shear modulus G shows a relative standard deviation of 27%; the local bulk modulus *B* shows a smaller, but still substantial, value of 18%¹³. This variation is likely to be greater for the range of bonding found in a multicomponent system¹⁴. As the coefficient of thermal expansion (CTE) is—other factors being constant—inversely proportional to B (ref. 15), we expect that it also shows significant variation. Noting that neighbouring regions of different CTE must satisfy elastic compatibility, atomic-level shears must develop on temperature change, and given the existence of soft spots, local structural change may occur. On annealing, metallic glasses undergo thermally activated structural relaxation to states of higher density and lower enthalpy. To avoid such effects, thermal cycling in the present work is only from near room temperature (293-343 K) to lower temperatures, always far below the glass-transition temperature T_g near which thermal relaxation occurs (Fig. 1) 16 .

As a metallic glass is heated towards $T_{\rm gr}$ there is exothermic structural relaxation (Fig. 2a). The relaxation spectrum (heat release rate as a function of temperature) can be detected by differential scanning calorimetry (DSC). While the shape of the spectrum may be of interest¹⁷, we focus here only on the overall heat of relaxation ($\Delta H_{\rm rel}$, shaded areas in Fig. 2b). Changes in $\Delta H_{\rm rel}$ have been used to characterize the effects of thermal and mechanical treatments on metallic glasses; for example, heavy plastic deformation (shot-peening¹⁷) can induce ageing (that is, relaxation to a lower-energy state) or rejuvenation. For lanthanum (La)-based metallic glasses, cycling between room temperature (293 K) and liquid-nitrogen temperature (77 K) leads to increases in $\Delta H_{\rm rel}$, that is, rejuvenation (Fig. 2b). For melt-spun ribbons, $\Delta H_{\rm rel}$ peaks at ~10 cycles; for bulk glass, $\Delta H_{\rm rel}$ is smaller, peaking at ~15 cycles; in each case, the maximum increase in $\Delta H_{\rm rel}$ is ~50% relative to the as-cast sample.

We next addressed whether such changes are due to cycling, or to the time spent at 77 K. Ribbons were held at 77 K for 4 h, much longer than the sum of the times spent at 77 K in cycling treatments, but this 'anneal' did not give a detectable change in $\Delta H_{\rm rel}$ (see Methods and Extended Data Fig. 1).

Because surfaces respond more rapidly than interiors, sample temperature is not spatially uniform during thermal cycling; surfaces must be in tension during cooling and in compression during heating. In nano-indentation, stress cycling can harden metallic glasses¹⁸, raising the question of whether the ΔH_{rel} changes (Fig. 2b) are due to the

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Figure 1 | **Thermal cycling of metallic glasses. a**, At high fractions of the glass-transition temperature, T_{g} to as low as 0.6 T_{g} , metallic glasses can undergo α and β relaxations, giving peaks in loss modulus as a function of T/T_{g} as shown by the curve¹⁶. **b**, The thermal cycling in the present work explores lower temperatures. Samples are cycled from near room temperature, 293–343 K (RT), to liquid-nitrogen temperature (77 K). The ranges of T/T_{g} reflect both range of temperature and the range of T_{g} for the metallic glasses, from 474 K for La₅₅Ni₁₀Al₃₅ (at.%) up to 700 K for Cu₄₆Zr₄₆Al₇Gd₁. **c**-**e**, Schematic depictions of the degree of heterogeneity in a metallic glass in the as-cast state (**c**), and after increasing numbers of thermal cycles (**d**, **e**). The population and intensity of soft spots (dark), with lower elastic stiffness and higher CTE, increases with cycling. The scale of these heterogeneities is not given by the present results (so no scale bars are given), but mapping of elastic properties^{11,12} suggests characteristic lengths of <10 nm.

temperature cycling itself, or to the associated stresses. As-cast rods of $La_{55}Ni_{10}Al_{35}$ bulk metallic glass (BMG) were subjected to room temperature–77 K cycles at the same time as thin discs cut from the rods. As detailed in Methods, the changes in ΔH_{rel} are independent of sample size, showing that stresses generated during cycling have a negligible effect (Extended Data Fig. 2). Thermal calculations (Extended Data Table 1) suggest that the stresses themselves are negligible, consistent with this conclusion.

Many aspects of thermal cycling could be tailored: upper and lower temperatures, holding times at these temperatures, rates of cooling and heating, number of cycles. We report preliminary studies in Methods; our focus here, however, is on property changes induced by thermal cycling.

In instrumented indentation (nano-indentation) of metallic glasses^{10,18}, initial yielding is indicated by a sharp 'pop-in' (increase in indentation depth *h*) on the load-*h* curve (Extended Data Fig. 3), corresponding to shear-banding onset. Cumulative distributions of initial yield pressure P_y (Fig. 3a) for La₅₅Ni₂₀Al₂₅ glass ribbon in three states permit the effects of a longer hold at 77 K to be compared with cycling. The median value of P_y , 2.98 GPa in the as-cast sample, decreases by 3% after 10-min hold at 77 K, and by a further 17% after ten room temperature–77 K (1 min) cycles. After the cycling treatments the relative width (1st to 9th decile) of the P_y distribution (\pm 15%) is greater than in the as-cast sample (\pm 7%), suggesting that cycling induces greater heterogeneity.

The corresponding distributions of hardness H (Fig. 3b), determined from h at full load, show similar trends. The median H of the as-cast sample, 2.65 GPa, is 1% lower after 10 min at 77 K, and a further 4% lower after ten room temperature–77 K cycles. For a given sample, the H distribution is narrower than that for P_y ; for example, after cycling, the relative width is only $\pm 3\%$ (one-fifth that of P_y). Evidently H, reflecting conditions for continuance of flow, is much



Figure 2 | Differential scanning calorimetry (DSC) of melt-spun ribbons of La₅₅Ni₂₀Al₂₅ and bulk rods of La₅₅Ni₁₀Al₃₅ metallic glasses. a, DSC traces on heating show an exotherm before the glass transition at $T_{\rm g}$ (here for ribbon). b, By subtraction of successive traces of specific heat $C_{\rm p}$ (see Methods), the heat of relaxation, $\Delta H_{\rm rel}$, associated with the exotherm can be determined (shaded areas) and is increased after room temperature–77 K thermal cycles. In both **a** and **b**, vertical bars give units and scales for the *y* axes. **c**, For ribbons, $\Delta H_{\rm rel}$ peaks at ~10 room temperature–77 K cycles; for discs (250–500 µm thick) cut from the bulk rod, $\Delta H_{\rm rel}$ is lower and peaks at ~15 cycles; the data points are for individual measurements and the changes lie outside the error range of ±30 J mol⁻¹ (standard deviation estimated as in Methods).

less sensitive to local heterogeneities than is P_y . The Young modulus, determined on unloading, shows similar behaviour to H (Extended Data Fig. 4).

The distribution curves of P_y and H confirm changes related to the number of cycles rather than the holding time at 77 K. The marked reduction in P_y induced by cycling is accompanied by a reduction in initial yield increments (Δh , Extended Data Fig. 4a).

Metallic glasses show exceptionally high yield stress and yield strain¹⁹, and can be very tough²⁰, but there is a clear need to improve their plasticity. Annealing reduces their plasticity, often inducing brittleness. Figures 2 and 3 show property changes opposite to those expected from annealing, suggesting that thermal cycling gives rejuvenation. The effects on macroscopic plasticity are explored for two BMG compositions tested in uniaxial compression (Fig. 4). The plastic strain of 1.5-mm-diameter samples of Zr₆₂Cu₂₄Fe₅Al₉ is improved (from 4.9% to 7.6%) by successive thermal cycles (Fig. 4a). Improvements in plasticity, smaller in absolute terms (for reasons given in Methods), but larger relatively, are seen in larger-diameter samples of the same glass (Fig. 4b), which also show a reduction in microhardness (by 4%, inset), similar to that for the median hardness of the La-based glass in Fig. 3b. Cu₄₆Zr₄₆Al₇Gd₁ BMG $(1.5 \times 1.5 \times 3.0 \text{ mm}^3 \text{ cuboid})$ shows an increase in plastic strain from 1.4% in the as-cast state to 5.1% after ten room temperature-77 K cycles (Fig. 4c). A sample fully relaxed by annealing is brittle and is not successfully rejuvenated by cycling, but a strong rejuvenation effect is possible after partial thermal relaxation, when thermal cycling gives a plastic strain similarly improved to that of as-cast samples.

Well below T_g , plastic flow in metallic glasses is localized into thin shear bands. Macroscopic plasticity is limited by premature failure on



Figure 3 Cumulative distributions of (a) initial yield pressure P_y and (b) hardness *H*. La₅₅Ni₂₀Al₂₅ metallic glass ribbon is tested in instrumented indentation, to a maximum load of 40 mN, in three states: as-cast, after a 10-min hold at 77 K, and after a further ten room temperature–77 K cycles each with 1 min hold. The cycling reduces P_y and *H*, and widens the distribution of their values. The widening is taken to represent increasing heterogeneity of the glass structure (Fig. 1*c*–*e*).

one or a few dominant shear bands. The plasticity can be improved by proliferation of shear bands, making the overall flow more uniform²¹. The shear-band spacing of ~10 µm in the as-cast sample (Fig. 4d) is reduced to ~2.5 µm in the thermally cycled sample, suggesting greater ease of shear-band initiation, consistent with reductions in P_y (Fig. 3a) and in initial yield displacements (Extended Data Fig. 4a). Resonant ultrasound spectroscopy (RUS) was used to measure the elastic moduli of Cu₄₆Zr₄₆Al₇Gd₁ BMG, which show no detectable change on thermal cycling (Extended Data Fig. 5).

Across a range of metallic glass compositions, melt-spun and bulk, thermal cycling induces changes in properties. As the cycling is down to 77 K, these changes can be compared with the property improvements (hardness, wear resistance) obtained by deep cryogenic treatment (DCT) of steels²². DCT, however, works through phase changes (transformation of residual austenite, precipitation of fine carbides) on holding at low temperature. In the present case, there is no discernible phase change (X-ray diffraction confirms that the samples remain fully



Figure 4 | Improved plasticity after thermal cycling. a, 1.5-mm-diameter rod samples of $Zr_{62}Cu_{24}Fe_5Al_9$ BMG (yield stress 1.68 \pm 0.05 GPa) show an increase in plastic strain under uniaxial compression when treated with 338 K to 77 K thermal cycles. b, The compressive plasticity of this glass is strongly dependent on sample dimension, but for all the diameters tested, thermal cycling leads to an increase in plastic strain, accompanied (inset) by a reduction in microhardness (in kg mm⁻²; each data point is the average of 20 measurements). c, $1.5 \times 1.5 \times 3.0$ mm³ cuboids of Cu₄₆Zr₄₆Al₇Gd₁ BMG (yield stress 1.61 \pm 0.04 GPa), both as-cast and annealed (1.0 h at 400 $^{\circ}$ C) to partial relaxation, show an increase in plastic strain under uniaxial compression after ten room temperature-77 K cycles; for full relaxation (1.5 h at 400 °C), the sample remains brittle. d, For the Cu₄₆Zr₄₆Al₇Gd₁ BMG samples, scanning electron microscopy shows that the population density of shear bands near the dominant shear band is lower in an as-cast sample (top) than in a similar sample tested after ten room temperature-77 K cycles (bottom). In a and c, horizontal bars indicate the scale and units for the x axes; quoted yield stress gives a guide to the y axes.

glassy after cycling; Extended Data Fig. 6), and the effects are more analogous to those induced in single-phase polycrystalline non-cubic metals, for example, thermal-cycling growth in uranium²³ and increased dislocation density in zirconium²⁴. The widespread use of DCT²⁵ does, however, suggest that cycling to cryogenic temperatures may be a practicable process.

The changes in $\Delta H_{\rm rel}$ (Fig. 2c) show rejuvenation followed by ageing, opposing trends known in processes such as irradiation or plastic deformation, which introduce both damage (raising the energy of the system) and mobility (enabling relaxation to lower energy). Shotpeening increases ΔH_{rel} in an annealed BMG, but for an as-cast BMG even this intense plastic deformation reduces ΔH_{rel} (ref. 17). Thermal cycling increases ΔH_{rel} even for a rapidly quenched metallic glass. The maximum increase in $\Delta H_{\rm rel}$ (Fig. 2c), 340 J mol⁻¹, is (correcting for differing DSC heating rates) $\sim 60\%$ of the increase in $\Delta H_{\rm rel}$ in a Zr-based BMG induced by one rotation in high-pressure torsion²⁶, and reductions in hardness (Figs 3b and 4b and ref. 26) are, pro rata, similar. The increases in $\Delta H_{\rm rel}$ (Fig. 2c) are very similar to those seen in elastostatic loading²⁷; it is remarkable that both these apparently mild processes give effects comparable with heavy plastic deformation, suggesting that non-affine strains in the nominally elastic regime are efficient in generating structural damage (disordering).

Although continued cycling leads to reversal in the ΔH_{rel} increase (Fig. 2c), the microhardness reduction appears to saturate (Fig. 4b inset) while the plasticity continues to improve (Fig. 4a, b); the onset

of yield (Fig. 3a) is facilitated more than general yielding (Fig. 3b) and the effect on high-frequency (MHz in RUS) macroscopic elastic moduli is negligible (Extended Data Fig. 5). These contrasts for different properties lead us to suggest that thermal cycling introduces heterogeneities (soft spots)⁴ that are particularly effective in initiating flow and improving plasticity, with lesser effects on the average structure of the metallic glass. That the effects (at least for ΔH_{reb} , Fig. 2c) are greater in melt-spun ribbon than in BMG of similar composition suggests that pre-existing heterogeneity (greater in a less-relaxed metallic glass) helps cycling to introduce more heterogeneity; this is expected if, as we speculate, the effect is due to non-affine thermal strains arising from the heterogeneity itself. The present results, however, do not identify any characteristic length scale for the heterogeneity.

Rejuvenation of metallic glasses, with improved plasticity, has been achieved by elastostatic loading^{27,28}, ion irradiation²⁹ and plastic deformation²¹. In comparison with these methods, thermal cycling is attractive: it is non-destructive, not involving shape change; unlike an applied elastostatic stress, it is isotropic and cannot introduce anisotropy; it can be applied to any sample (thin film, ribbon or bulk) and repeated as necessary; it changes the whole sample (not just the surface as, for example, in shot-peening or ion irradiation, or only inside shear bands); and it is controllable, can be performed *in situ*, induces no macroscopic residual stresses, involves strains far below the elastic limit, and induces no macroscopic plastic flow.

Online Content Methods, along with any additional Extended Data display items and Source Data, are available in the online version of the paper; references unique to these sections appear only in the online paper.

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Author Contributions Preparation of metallic glasses was by S.V.K. and Z.L. The thermal-cycling treatments were performed by Y.H.S., S.V.K., A.R.B. and A.C.; calorimetry by Y.H.S., S.V.K., Z.L., A.C. and A.R.B.; nano-indentation by S.N. and A.R.B.; compression and microhardness tests by S.V.K., Y.H.S., A.R.B. and A.C.; Y.-R.Y. and Y.H.S.; and resonant ultrasound spectroscopy by Y.H.S., M.A.C., A.R.B. and A.C. Direction of the work was by D.V.L.-L. (Sendai), H.Y.B. and W.H.W. (Beijing), and A.L.G (Cambridge). A.L.G. led the project and the writing of the pare. All authors contributed to interpretation and presentation of the results.

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METHODS

Metallic glasses in this study. Sample preparation and property measurements took place in three laboratories. Kinetic phenomena in glasses are best compared on a homologous temperature scale, using the glass-transition temperature T_{g} as the normalizing parameter. The selected glasses allow studies over a range of T/T_g : for example, room temperature (293 K) covers the range of $T/T_g = 0.42-0.62$ for these glasses. (The values of Tg were determined using differential scanning calorimetry at standard heating rates of 20 and 40 K min⁻¹.) The four selected glasses are: (1) $Cu_{46}Zr_{46}Al_7Gd_1$ rod, 3 mm diameter ($T_g = 700$ K (ref. 30)); (2) $La_{55}Ni_{20}Al_{25}$ (at.%) melt-spun ribbon 40 μ m thick ($T_g = 475$ K); (3) La₅₅Ni₁₀Al₃₅ rod 3 mm diameter ($T_g = 474$ K); and (4) $Zr_{62}Cu_{24}Fe_5Al_9$ rods, 1.5, 2.0 or 2.5 mm diameter $(T_g = 658 \text{ K})$. Master alloys were prepared by arc-melting 3–4N pure elements. Glassy ribbons, 40 µm thick, were produced by melt-spinning, under argon atmosphere, on to a single copper wheel at 4,000 r.p.m. Bulk metallic glass (BMG) rods, with diameter 1.5-3.0 mm and length up to 80 mm, were cast, under Ti-gettered argon atmosphere, into water-cooled Cu-moulds, using induction melting and suction casting.

Thermal cycling. The La-based and $Cu_{46}Zr_{46}Al_7Gd_1$ samples were inserted into liquid nitrogen for 1 min, then heated by a hair-dryer set at room temperature, and held for 1 min; samples were treated with up to 25 such cycles. The $Zr_{62}Cu_{24}Fe_5Al_9$ samples were inserted into liquid nitrogen for 1 min, then into ethanol at 333–343 K for 1 min; samples were treated with up to 60 such cycles.

The paper reports only results from cycling between near room temperature and liquid-nitrogen temperature (77 K). Keeping the upper temperature fixed (near room temperature), it is expected that there would be an optimum lower temperature for maximum effect of thermal cycling (at given cooling and heating rates, and hold times at the upper and lower temperatures): for higher temperatures closer to room temperature, there would be less driving force for structural change; for lower temperatures further from room temperature, the rate of structural change near the lower temperature would be lower.

In addition to cycling down to 77 K, melt-spun ribbons of La₅₅Ni₂₀Al₂₅ glass were cycled down to dry-ice (CO₂) temperature (195 K) and to liquid-helium temperature (4.2 K) to explore the role of the lower temperature. The state of the samples was assessed using DSC (with heating rates from 20 to 40 K min⁻¹). For a given type of sample, the heat of relaxation $\Delta H_{\rm rel}$ is lower for higher heating rate; it is therefore appropriate to normalize the $\Delta H_{\rm rel}$ values relative to that for the as-cast glass measured at the same DSC heating rate. Measurements were made for samples: (1) held at 195 K for 100 min, for which the ratio of $\Delta H_{\rm rel}$ to that of the as-cast glass is 1.23; (2) treated with ten cycles of 1 min at 195 K (ratio = 1.02); (4) treated with ten cycles of 10 min at 195 K (ratio = 1.37). Given the low number of samples tested, there is significant uncertainty in these relative values (±15%); nevertheless, it seems that, within the limited range of these tests, cycling down to 77 K produces the strongest effect.

A Ti-based BMG cycled up to 400 times between 77 K and 423 K (to simulate conditions on the surface of a spacecraft in low earth orbit) showed slight relaxation rather than rejuvenation³¹. This suggests that there is a significant annealing effect during the holds (25 min each) at the upper temperature, which itself is higher than in our work (and corresponds to $T/T_g = 0.68$, inside the range associated with β relaxation, see Fig. 1a in main text).

Stresses associated with thermal cycling. As noted in the paper, it is important to understand whether the effects of thermal cycling are due to the temperature changes themselves or to stresses arising from non-uniform temperature in the samples during cooling and heating. Whether the temperature is significantly nonuniform can be assessed using the dimensionless Biot number,

$$\mathrm{Bi} = \frac{hL}{\kappa} \tag{1}$$

where *h* is the heat-transfer coefficient between the sample and the surrounding medium, *L* is a characteristic sample dimension (usually taken to be the ratio of sample volume to surface area), and κ is the thermal conductivity of the sample material. For a flat plate, *L* is the half-thickness; for a cylinder the half-radius. If Bi < 0.1, the temperature differences within the sample are negligible compared to the step in temperature between the sample and its surroundings (Newtonian cooling). Conversely, if Bi > 1, there are significant temperature gradients in the sample. We examine the conditions for samples immersed in liquid nitrogen.

Cooling in liquid nitrogen has been widely studied, particularly in connection with cryopreservation³². The 'French straw' sample containers used are cylinders of 2.8 mm diameter, and so are very similar in dimension to the cast BMG rods in the current work. On first immersion, there is film boiling, in which a film of vapour limits the heat transfer; in this regime³² h = 148 W m⁻² s⁻¹. As the sample cools, there is a transition to the nucleate-boiling regime³² in which h = 1,355 W m⁻² s⁻¹. The transition point has been measured for a variety of metallic samples

of geometry comparable to the samples in the present work³²; it occurs at temperatures ranging from 150 to 100 K. Thus a room-temperature sample immersed in liquid nitrogen spends around 75% of the temperature range down to 77 K with the relatively poor heat transfer associated with the film-boiling regime.

We used materials parameters determined for Vitreloy 1; Zr-based BMGs of this kind are among those for which thermal-cycling treatments might be put into practice. For Vitreloy 1, the thermal conductivity decreases with decreasing temperature³³ and starts to level off below room temperature. At 300 K, $\kappa = 4.59 \text{ W K}^{-1} \text{ m}^{-1}$; we use a rough estimate of $\kappa = 4 \text{ W K}^{-1} \text{ m}^{-1}$ for the temperature range in the present work. Extended Data Table 1 gives the estimates of Bi for the various sample geometries of interest.

We can see that for nearly all of these cases Bi < 0.1, and therefore temperature gradients (and thermal stresses) in the samples are negligible. Only for the largest-diameter cylinders in the final (nucleate boiling) stages of cooling is there any significant deviation from Newtonian cooling. These calculations support the experimental work (Extended Data Fig. 2 and associated discussion, see below) suggesting that sample size is not a significant factor in the thermal-cycling effect, and reinforce the conclusion that the effect is due to the temperature changes themselves and not due to any stresses arising from non-uniform temperature profiles in samples during cooling and heating.

It is of interest to consider the possible extent of thermal stresses associated with non-uniform sample temperature. Taking the value³⁴ determined for the Zr-based BMG Vitreloy 106, the linear coefficient of thermal expansion is $\alpha = 8.7 \times 10^{-6} \text{ K}^{-1}$. For a room-temperature sample immersed in liquid nitrogen, the maximum possible temperature difference during cooling is between 293 K (room temperature, possible at the sample centre) and 77 K (possible at the sample surface). For this temperature difference, the linear thermal strain is 0.19%, which would give internal stresses of the order of 10% of the macroscopic yield stress. In practice, as highlighted by the low values for the Biot number in Extended Data Table 1, any thermal stresses would be much lower than this value. Rejuvenation of metallic glasses by elastostatic loading²⁸ typically involves much higher stresses (up to 95% of the yield stress) for much longer times (tens of hours, rather than a few minutes). Again, it seems that stress effects in the current thermal cycling should be negligible.

Differential scanning calorimetry. This was performed (as in Fig. 2) using a Q2000 DSC (TA Instruments). Samples were heated at 20 K min⁻¹ from room temperature to 503 K (into the supercooled liquid state), held for 2 min, then cooled to room temperature at 20 K min⁻¹. A second cycle using the same procedure was used as the baseline for subtraction from the first cycle. The heat of relaxation $\Delta H_{\rm rel}$ is calculated from the area between the two curves, from the onset of relaxation to the glass transition³⁵. Additional data were acquired using a Perkin Elmer DSC 8500 at a heating rate of 40 K min⁻¹.

Extended Data Figure 1a shows examples of DSC traces used to assess the reproducibility of measurements of the heat of relaxation $\Delta H_{\rm rel}$ (using the base-line-subtraction method outlined above). Six samples of as-cast ribbons were tested, giving an average value of $\Delta H_{\rm rel} = 740 \,\text{J mol}^{-1}$, with a standard deviation of 30 J mol⁻¹ (that is, ±4%).

As noted in the paper, it is important to know whether the effects of thermal cycling between near room temperature and 77 K are due just to the time spent at 77 K. Samples of La₅₅Ni₂₀Al₂₅ glassy ribbons were held at 77 K for 4 h (a time much longer than the total spent at 77 K in the course of ten room temperature–77 K cycles). Within experimental error (Extended Data Fig. 1b), such holds at 77 K appear to have no effect on ΔH_{rel} . Of course, a hold at 77 K represents a single cycle. As shown by the nano-indentation results in Fig. 3 in the main paper, even one cycle can have a measurable effect on properties. In the present case of ΔH_{rel} , however, any change as a result of one cycle is within the experimental uncertainty.

As noted in the main paper and discussed above in the Methods, it is necessary to examine whether the effects of thermal cycling are dependent on sample size. The non-uniform temperature in a larger sample could give rise to stresses and the cycling of those stresses could have effects. As-cast rods (3 mm diameter) of La₅₅Ni₁₀Al₃₅ BMG were subjected to room temperature–77 K cycles at the same time as thin (250–500 µm thick) discs pre-cut from the rods. After cycling, discs were cut from the centres of the bulk rods, so that samples of the same glass and same DSC-sample size could be compared after thermal cycling in two different geometries (thin disc and bulk). Extended Data Figure 2 shows that the changes in $\Delta H_{\rm rel}$ are essentially the same for the samples thermally cycled in disc and in bulk form, suggesting that the stresses generated during cycling have a negligible effect. The two sets of data are combined in Fig. 2c.

Instrumented indentation. This was performed using an XP Nanoindenter (MTS Systems Corp.) at room temperature under load control up to a maximum load (F_{max}) of 40 mN. A diamond spherical indenter of tip radius 8.074 µm was used, its area function calibrated using fused silica and sapphire standards. Loading and unloading rates were 0.5 mN s⁻¹; the drift rate was maintained below

0.07 nm s⁻¹. Samples were loaded to $F_{\rm max}$ then unloaded to $0.05F_{\rm max}$ and maintained at that load for 60 s for measurement of thermal drift. The cumulative distribution curves (Fig. 3) include 39–48 data points. Initial yield events (pop-ins) were identified directly from the *F*-*h* curve and the associated spike in indenter tip velocity, or from the first deviation from the Hertzian elastic solution fitted to the *F*-*h* curve. Hardness was determined by the standard Oliver and Pharr method³⁶, on unloading at $F_{\rm max}$.

Tests were conducted on melt-spun ribbons of $La_{55}Ni_{20}Al_{25}$ glass under the conditions noted above. Typical load-displacement curves (Extended Data Fig. 3) show pop-ins (that is, the transition from purely elastic to elastic-plastic deformation) with a moderate sharpness that is not affected by thermal cycling. The pop-ins are characterized by the values of initial yield load F_y and initial yield displacement Δh . Initial yield pressure P_y is calculated from:

$$P_{\rm y} = \frac{F_{\rm y}}{\pi a_{\rm y}^2} \tag{2}$$

where a_y is the contact radius at initial yield. The reduced or indentation modulus E_r is determined by the standard method^{36,37}, on unloading from F_{max} . E_r is also determined by using the Hertzian elastic solution to fit the shape of the load-displacement curve up to the first pop-in. E_r is given by³⁸:

$$\frac{1}{E_{\rm r}} = \frac{(1-v_{\rm i}^2)}{E_{\rm i}} + \frac{(1-v^2)}{E}$$
(3)

where *E* and *v* are, respectively, the Young modulus and Poisson ratio of the glass, while the corresponding values³⁶ for the diamond indenter tip are taken to be $E_i = 1,141$ GPa and $v_i = 0.07$. Using equation (3), values of *E* were obtained.

Tests were conducted on ribbon samples in three states: (1) as-cast; (2) after one cycle from room temperature to a 10-min hold 77 K and back to room temperature; and (3) after a further ten cycles to 77 K each with 1 min hold. The effect of a longer hold at 77 K can thus be compared with cycling. All metallic-glass samples, as-cast or treated, are found to have some variation in properties over length scales of millimetres. For this reason it is important to compare data taken from a given region of the sample surface. In the present work, indentations were made in the same region (within 200 μ m) of the given sample surface in the three successive states. Extended Data Table 2 collects data on the instrumented-indentation tests in the present work. Cumulative distributions, such as those shown in Fig. 3, are characterized by the median value and by the 1st and 9th deciles.

Extended Data Figure 4a shows the distribution of values of initial yield pressure P_y and initial yield displacement Δh for the three states of the La₅₅Ni₂₀Al₂₅ glass ribbons. On cycling, both P_y and Δh are reduced. There is a correlation, as expected, between higher values of P_y and higher values of Δh . The spread in values of Δh is particularly wide. After ten room temperature–77 K cycles, there is a marked reduction in the incidence of larger values: the initial yield events are smaller, reflecting less catastrophic shear banding.

The Young's modulus of the glass *E* determined by unloading from full load (Extended Data Fig. 4b) shows similar effects to those found for hardness *H* (Fig. 3b). The median value of E_{unload} in the as-cast sample decreases by 0.6% after 10 min hold at 77 K (compared to a 0.9% decrease in *H*), and by a further 7% after ten room temperature–77 K (1 min) cycles (compared to 4% for *H*). After the cycling treatments the relative width (1st to 9th decile) of the *E* distribution (\pm 2.4%, compared to \pm 3.1% for *H*) is greater than in the as-cast sample (\pm 1.4%, compared to \pm 1.3% for *H*).

The Young's modulus determined from the Hertzian fit of the F-h curve up to the first pop-in (Extended Data Fig. 4c) has a median value \sim 5% lower than that determined by unloading. The median value of $E_{\rm Hertz}$ in the as-cast sample decreases by 2.4% after 10-min hold at 77 K, and by a further 8.8% after ten room temperature-77 K (1 min) cycles. The relative width of the E_{Hertz} distribution in the as-cast sample ($\pm 1.6\%$) is similar to that for E_{unload} , but this width increases more (to $\pm 7.3\%$) after the cycling treatments. This suggests that the elastic properties up to the first pop-in are more sensitive to local variations in the sample. Compression tests. These were made on Instron 5581 and Hounsfield 25 kN machines, loading at a strain rate of 5×10^{-4} s⁻¹. The Zr₆₂Cu₂₄Fe₅Al₉ samples were cylindrical rods of 2:1 height:diameter, with diameter of 1.5, 2.0 or 2.5 mm. The $Cu_{46}Zr_{46}Al_7Gd_1$ samples were $1.5 \times 1.5 \times 3.0 \text{ mm}^3$ cuboids cut from the original cast rod of 3 mm diameter. All samples were polished to a mirror finish. The gradients of the elastic portions of the stress-strain curves in Fig. 4a are affected by the machine compliance and underestimate the Young's modulus. The curves are used only to extract values for the plastic strain as shown. The curves in Fig. 4c are almost fully corrected for machine compliance effects.

The curves in Fig. 4a and Fig. 4c show engineering stress and engineering strain. The strain values are calculated from the displacement of the testing machine and not from strain gauges on the samples. The gradients of the elastic portions of the curves are therefore affected by the machine compliance and significantly underestimate the Young's modulus of the materials. The curves are used only to extract values for the plastic strain before catastrophic failure. Figure 4b collects data for rod samples of $Zr_{62}Cu_{24}Fe_5Al_9$ BMG of three diameters: 1.5 mm, 2.0 mm and 2.5 mm, and corresponding heights 3.0 mm, 4.0 mm and 5.0 mm. The stress-train curves for 1.5 mm samples are shown in Fig. 4a; those for 2.0 mm and 2.5 mm samples in Extended Data Fig. 7.

As has been much discussed^{21,39}, larger samples are more susceptible to catastrophic failure through shear localization. Such size effects are extrinsic, and relate only to the mechanical tests themselves. In addition, there is an intrinsic effect: larger samples have a lower cooling rate during casting and so are more relaxed and tend to be more brittle in the as-cast state.

Resonant ultrasound spectroscopy. RUS measurements were conducted as in earlier work^{40,41}. Measurements were at room temperature on $1.5 \times 1.5 \times 3.0$ mm³ cuboids cut from the as-cast 3-mm-diameter rod of Cu₄₆Zr₄₆Al₇Gd₁ BMG. For each sample, a resonance spectrum was collected in the range 0.05–2.0 MHz with 65,000 data points. To detect changes in shear and bulk modulus, 10,000 data points were collected in each case in the ranges 0.900–0.910 MHz and 0.978–0.982 MHz, respectively. Within experimental error (less than ±0.1% for the shear modulus), room temperature–77 K thermal cycling has no effect on the elastic moduli of the glass (Extended Data Fig. 5). This suggests that the reductions in Young's modulus (Extended Data Fig. 4b, c) and correspondingly hardness (Fig. 3b) seen under quasi-static conditions are largely attributable to anelastic (that is, time-dependent elastic) strains.

Dynamic mechanical analysis. This technique (not discussed in the paper), permitting measurement of the loss and storage moduli of a material, was applied (using methods reported previously⁴²) to ribbons of La₅₅Ni₂₀Al₂₅ glass (Extended Data Fig. 8) to explore the nature of the rejuvenation effect. The loss modulus shows a sharp maximum near the glass transition, associated with α relaxation. After ten room temperature–77 K cycles, the maximum is lowered by 2.3 K. This earlier onset of softening on heating is consistent with a rejuvenated state. The broad maximum at lower temperature (centred around 115 °C, Extended Data Fig. 8) shows β relaxation. The β relaxation in metallic glasses has been linked to mechanical properties^{42,43}. In view of the improved plasticity seen after thermal cycling (Fig. 4 and Extended Data Fig. 7), it is surprising that the effect of thermal cycling on the β relaxation is so slight (inset in Extended Data Fig. 8), especially compared to the clear effect on the α relaxation.

X-ray diffraction. There are reports that under cyclic mechanical loading at room temperature, in the elastic regime, a Zr-based BMG can undergo some crystallization⁴⁴. Deep cryogenic treatment of steels certainly induces phase changes^{22,25}. In the present work, X-ray diffraction (Bragg–Brentano geometry, Bruker D8 instrument, Cu K α radiation) was used to check that the samples remained fully glassy throughout, and especially after thermal cycling. Representative diffractograms for ribbon and bulk samples are shown in Extended Data Fig. 6. We found no evidence for any crystallization induced by the thermal-cycling treatments. The property changes suggest that the cycling does induce changes are too subtle to be detectable with the simple X-ray methods used in the present work.

Other methods. Microhardness was measured on an Akashi MVK-HVL hardness testing machine, with 20 measurements for each data point. Scanning electron microscopy (secondary-electron imaging) was performed using a JEOL 5200 instrument.

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Extended Data Figure 1 | DSC traces for ribbons of La₅₅Ni₂₀Al₂₅ glass (heated at 20 K min⁻¹). a, These example traces for as-cast samples show the exotherm, just below the glass-transition temperature, from which the heat of relaxation $\Delta H_{\rm rel}$ is determined. b, The effect of holding at 77 K: the traces show that the heat of relaxation $\Delta H_{\rm rel}$ given by the exotherm just below the glass-transition temperature, is, within the experimental error of 4%, the same for an as-cast ribbon, and for a sample held for 4 h at 77 K.



Extended Data Figure 2 | Differential scanning calorimetry (DSC) of La₅₅Ni₁₀Al₃₅ bulk metallic glass. The heat of relaxation $\Delta H_{\rm rel}$ is compared for discs (250–500 µm thick) pre-cut from the bulk rod and then treated with room temperature–77 K cycles, and for discs post-cut from treated rod samples. There is no difference between these cases within the error of \pm 30 J mol⁻¹.



Extended Data Figure 3 | **Load-displacement curves for ribbons of La**₅₅Ni₂₀Al₂₅ metallic glass tested up to a maximum load F_{max} of 40 mN. The initial yield load F_y is indicated on the curves for the glass in three states: as-cast, after a 10-min hold at 77 K, and after a further ten room temperature–77 K cycles each with 1 min hold.



Extended Data Figure 4 | Instrumented indentation of La₅₅Ni₂₀Al₂₅ metallic glass ribbon. a, Initial yielding, characterized by initial yield pressure P_y and initial yield displacement Δh . Indentations are made for three states of the sample: as-cast, after a 10-min hold at 77 K, and after a further ten room temperature-77 K cycles each with 1-min hold. b, c, Distributions of the Young's modulus, using data from the same indentations of the sample in three states as in Fig. 3 and in a. Values of the Young's modulus in the glass *E* are determined (b) by the standard Oliver and Pharr³⁶ method on unloading from F_{max} and (c) from Hertzian fitting to the *F*-*h* curve up to the first pop-in.



Extended Data Figure 5 | Elastic moduli derived from resonant ultrasound spectroscopy for $Cu_{46}Zr_{46}Al_7Gd_1$ BMG treated with room temperature-77 K thermal cycles. Shear modulus (left axis); bulk modulus (right axis). Error bars, root mean square errors in the fitted frequencies.



Extended Data Figure 6 | X-ray diffraction traces for Zr-based metallic glasses subjected to 338–77 K thermal cycles. a, For melt-spun ribbons of $Zr_{61.1}Cu_{26.3}Fe_{2.1}Al_{10.5}$; b, for rods of $Zr_{62.2}Cu_{23.9}Fe_{4.8}Al_{9.1}$. No clear changes are induced by the cycling treatments.



Extended Data Figure 7 | Compressive stress–strain curves for rod samples of $Zr_{62}Cu_{24}Fe_5Al_9$ BMG. a, For rods of 2 mm diameter; b, for rods of 2.5 mm diameter. In each case, increasing numbers of 338 K to 77 K thermal cycles cause the plastic strain to increase.



Extended Data Figure 8 | Dynamic mechanical analysis of ribbons of La₅₅Ni₂₀Al₂₅ metallic glass. The heating rate is 3 K min⁻¹. The general form of the curve matches that shown in Fig. 1a, where an example was chosen of a metallic glass showing β relaxation at a particularly low value of T/T_g . For La₅₅Ni₂₀Al₂₅ the β relaxation is centred at $T/T_g \approx 0.8$.

Extended Data Table 1 | Values of the Biot number Bi for immersion in liquid nitrogen of the various sample geometries in the present work

Sample geometry	L (μm)	Bi, film boiling	Bi, nucleate boiling	
melt-spun ribbon, 40 μ m thick	20	7.4×10 ⁻⁴	6.8×10 ⁻³	
disc, 250–500 μm thick	125–250	(4.6-9.2)×10 ⁻³	(4.2-8.5)×10 ⁻²	
cylinder, 1.5 mm diam.	375	1.4×10 ⁻²	0.13	
cylinder, 2.0 mm diam.	500	1.8×10 ⁻²	0.17	
cylinder, 2.5 mm diam.	625	2.3×10 ⁻²	0.21	
cylinder, 3.0 mm diam.	750	2.8×10 ⁻²	0.26	

The film-boiling regime applies until the sample surface temperature decreases to between 150 K and 100 K, and the nucleate-boiling regime applies thereafter.



Extended Data Table 2 | Summary of instrumented-indentation tests

State	as-cast			10 min LN			10 min & 10 cycles LN		
Property	10%	median	90%	10%	median	90%	10%	median	90%
No. of indents		48			39			47	
F_{y} (mN) ± 0.1	10.6	14.3	16.0	10.6	13.5	14.3	5.8	9.3	11.5
∆ <i>h</i> (nm) ± 0.2	5.1	8.2	11.4	3.6	6.8	10.7	3.2	4.5	6.5
$P_{\rm y}$ (GPa) ± 0.015	2.69	2.98	3.12	2.61	2.89	2.96	1.97	2.38	2.66
$h_{\max}(nm) \pm 1$	442	445	450	444	449	453	458	469	481
H (GPa) ± 0.003	2.607	2.647	2.678	2.576	2.622	2.672	2.450	2.518	2.607
E_{unload} (GPa) ± 0.04	43.52	44.29	44.79	43.47	44.04	44.47	39.65	40.93	41.59
E_{Hertz} (GPa) ± 0.04	41.95	42.65	43.33	40.83	41.64	42.22	34.84	37.90	40.35

Cumulative distributions, such as those shown in Fig. 3, are characterized by the median value and by the first and ninth deciles.