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Citation: Wakeda, Masato, Junji Saida, Ju Li, and Shigenobu Ogata. "Controlled Rejuvenation of Amorphous Metals with Thermal Processing." Scientific Reports 5 (May 26, 2015): 10545.

As Published: http://dx.doi.org/10.1038/srep10545

Publisher: Nature Publishing Group

Persistent URL: <http://hdl.handle.net/1721.1/98444>

Version: Final published version: final published article, as it appeared in a journal, conference proceedings, or other formally published context

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SCIENTIFIC REPERTS

Received: 30 November 2014 accepted: 17 April 2015 Published: 26 May 2015

Controlled Rejuvenation of OPENAmorphous Metals with Thermal Processing

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Rejuvenation is the configurational excitation of amorphous materials and is one of the more promising approaches for improving the deformability of amorphous metals that usually exhibit macroscopic brittle fracture modes. Here, we propose a method to control the level of rejuvenation through systematic thermal processing and clarify the crucial feasibility conditions by means of molecular dynamics simulations of annealing and quenching. We also experimentally demonstrate rejuvenation level control in Zr₅₅Al₁₀Ni₅Cu₃₀ bulk metallic glass. Our local heat-treatment recipe (rising **temperature above 1.1***T***g, followed by a temperature quench rate exceeding the previous) opens avenue to modifying the glass properties after it has been cast and processed into near component shape, where a higher local cooling rate may be afforded by for example transient laser heating, adding spatial control and great flexibility to the processing.**

Amorphous metals¹⁻³ have excellent properties such as high corrosion resistance⁴, high strength^{[5](#page-7-2)}, and large elastic elongation⁶. At room temperature, however, plastic deformation localizes into narrow band region, and sudden fracture occurs exhibiting almost no macroscopic plastic deformation under uniaxial tensile and compressive loadings^{$7-9$}. This brittle nature is the most serious flaw of amorphous metals for use as structural materials. Therefore, improvement, and moreover tuning of the deformability has been the largest challenge of the decades in the field of amorphous metals¹⁰⁻¹³. Rejuvenation¹⁴⁻¹⁷ is the structural excitation of amorphous solids accompanied by an increase in the enthalpy and free volume and is thus the inverse of aging. Since rejuvenation has to change the elastic and plastic deformation behavior¹⁷ of amorphous metals, it is a new but promising approach for tuning the deformability of amorphous metals. It has recently been reported that shot-peening¹⁶ and sever plastic deformation^{[17](#page-8-1)} can mechanically induce the rejuvenation. On the other hand, structural excitation driven by a thermal loading process^{18–19} also has potential application for practical usage, because the thermal loading process realizes a local rejuvenation control, which is more difficult by mechanical processing. However, the feasibility rejuvenation driven by thermal annealing and quenching is still controversial because of few experimental evidence¹⁸ and a lack of clear knowledge of the nonequilibrium glass properties.

In this work, we theoretically propose a recipe to control the level of rejuvenation through systematic thermal processing and clarify the crucial feasibility conditions by means of molecular dynamics (MD) simulations and figure out a rejuvenation map with respect to annealing temperature and quenching rate. Our rejuvenation map provides temperature-cooling rate conditions to achieve desired level of rejuvenation while avoiding crystallization and/or gross shape change. Based on the rejuvenation map we also experimentally demonstrate the realization of rejuvenation level control in $Zr_{55}Al_{10}Ni_{5}Cu_{30}$ bulk

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Figure 1. Schematic illustration of the initial melt-quenching and subsequent thermal loading process. The system was initially quenched at a constant cooling rate of \widetilde{R}_q (B \rightarrow D). The instantaneous potential energy at temperature T in the initial-quenching process is represented as $\widetilde{E}_q(T)$. The as-quenched model was then subjected to thermal loading consisting of heating (D→E), isothermal annealing (E→F), and quenching (F→G) at various annealing temperatures T_a and constant cooling rates R_q^i . The instantaneous potential energy at temperature *T* in the second quenching process (F \rightarrow G) is represented as $E_a^i(T)$.

metallic glass, which evidences our rejuvenation map and recipe do work well even for typical experimental cooling rates. Moreover, the internal structural features and mechanical properties of rejuvenated amorphous metals are revealed.

Simulation Procedures

For all of the MD simulations described here, we use the Lennard-Jones (L-J) potentials²⁰ developed for the Cu-Zr amorphous alloy and a 30,000-atom simulation box with three-dimensional (3D) periodic boundary conditions. We consider alloys with atomic compositions of $Cu_{50}Zr_{50}$, $Cu_{30}Zr_{70}$, and $Cu_{57}Zr_{43}$ to ensure composition-independent generality (see Supplementary Information in regard to the latter two). The MD time step was set to 1 fs, and we employed the *NPT* ensemble for which the temperature and pressure are controlled by the Nose-Hoover²¹ and Parrinello-Rahman methods²², respectively. An amorphous model was constructed via a melt-quenching process as follows [\(Fig. 1](#page-2-0)). We first arranged the atoms randomly in the simulation box and melted the structure by keeping the model at a high temperature of 3000K for 20ps [\(Fig. 1;](#page-2-0) process A→B). The melted model was then quenched at a constant cooling rate of $\widetilde{R}_q(=1 \times 10^{12} \text{ K/s})$ from 3000 K to 0 K (process B→D). This quenched product is referred to as the "as-quenched model" hereafter, and we use a tilde to represent the initial quenching process. By monitorinng the specific volume change during the rapid quenching process with a cooling rate of \widetilde{R}_{q} , the glass transition temperature T_{g} was determined to be approximately 900K from the kink in volume-temperature curve. Since T_g generally depends on the cooling rate²³, we also determined a set of temperatures T_g^i for five different cooling rates \tilde{R}_g^i ($i=1-5$): $\tilde{R}_q^1=1\times 10^{11}$ K/s, $\tilde{R}_q^2=3\times 10^{11}$ K/s, $\widetilde{R}_{q}^{3} = 1 \times 10^{12}$ K/s, $\widetilde{R}_{q}^{4} = 3 \times 10^{12}$ K/s, and $\widetilde{R}_{q}^{5} = 1 \times 10^{13}$ K/s (see Supplementary Information, Fig. S1).

We conducted successive thermal loading simulations of the as-quenched model as follows. First, the as-quenched model was reheated at a heating rate of 1×10^{13} K/s up to a temperature T_a (process D→E) and then annealed for a time period t_a (=2000 ps) under constant T_a and zero external pressure conditions (process E→F). Isothermal annealing processes with t_a =500 and 1000 ps were also conducted and revealed that the annealing time t_a had a small effect on our simulation results (see Supplemental Information, Fig. S2). We assumed various T_a values ranging from 0.44 T_g to 1.67 T_g to examine the effect of the annealing temperature on the level of thermal rejuvenation. After isothermal annealing at T_a for t_a , the model was quenched again at five different cooling rates, R^i_q (= \widetilde{R}^i_q) (*i* = 1 – 5), from T_a to 0K (process F→G). This final product is referred to as the "annealed model" hereafter.

Results and Discussion

Rejuvenation map. [Figure 2\(a\)](#page-3-0) shows the generated excess potential energy $E_a^i(0)$ due to the thermal loading process from state D to G for different T_a and R_q^i values. If any aging occurs during the thermal loading process, then the excess potential energy value should fall below zero. On the other hand, if thermal rejuvenation occurs, then the excess potential energy value should remain positive. Thus, the excess potential energy can represent the level of rejuvenation or aging depending on the sign. As we see in Fig. 2(a), significant aging is observed whenever the cooling rate is less than \widetilde{R}_q , regardless of the

Figure 2. Rejuvenation map. (**a)** Change in the potential energy caused by the thermal loading process (D→G), where $E^i_a(0)$ represents the potential energy of the annealed model with a cooling rate of R^i_q . The potential energy of the as-quenched model with a cooling rate \tilde{R}_q^i is shown by the dashed lines. The grey rectangles represent 1.3 T_g^i , where T_g^i is the estimated glass transition temperature in the melt-quenching
process with a cooling rate $\tilde{R}_q^i (= R_q^i)$. (b) Change in the potential energy calculated by MD and the c the enthalpy obtained from the experiment. For the cooling rate after annealing, R_q^i is normalized by \widetilde{R}_q in the MD simulations, and $R_{\text{Exp.}}$ is normalized by $R^*_{\text{Exp.}}$ for the experimental data.

annealing temperature T_a , whereas thermal rejuvenation is observed in the opposite case. Thus, a higher cooling rate after isothermal annealing than that of the initial quenching process is a *necessary* condition for thermal rejuvenation, but this alone is insufficient. Thermal rejuvenation is only realized when the annealing temperature is above a certain critical temperature $T_c \approx 1.1 T_g$. Thus, the two conditions, $R_0^i > \tilde{R}_a$ and $T_a > T_c$, are crucial conditions for feasible thermal rejuvenation. We also found that a higher R_1^i and a higher T_a result in a higher level of rejuvenation under these conditions. [Figure 2\(a\)](#page-3-0) can then be used as a rejuvenation map to identify conditions for controlling the rejuvenation level. It should be noted that T_c is not an intrinsic material property but depends on both \widetilde{R}_q and R^i_q as follows: 1) Since a higher $\widetilde{R}_{\rm q}$ leads to a thermodynamically unstable as-quenched state, further aging proceeds easily during later thermal loading, and the T_c value is shifted to a higher temperature. 2) Since a higher R_q^i does not allow a more thermodynamically stable state to be found during the final quenching process, limited aging proceeds, and the T_c value is shifted to a lower temperature ([Fig. 2\(a\)](#page-3-0)).

The levels of aging and rejuvenation are relative quantities and depend on the choice of reference amorphous state. Thus, if we choose a perfectly rejuvenated amorphous state as the reference amorphous state, which is supposedly constructed through melt quenching at an infinitely fast cooling rate, then further rejuvenation will not occur. In reality, however, it is impossible to realize such a perfectly rejuvenated state; even though a near-perfectly rejuvenated state can be realized, we would not be able to observe it because of the extremely short lifetime of the highly excited state. For this reason, the discussions above are valid for actual glass systems in general, in which both aging and rejuvenation can be realized.

It is worth noting that evaluating the level of aging and rejuvenation based on the volume change (i.e., mass density change) yields qualitatively similar results to those in [Fig. 2\(a\)](#page-3-0) (see Supplementary Information, Fig. S3).

Based on the above findings, we demonstrate thermal rejuvenation experimentally in a $Zr_{55}Al_{10}Ni_5Cu_{30}$ bulk glassy alloy ($T_g = 684$ K at a heating rate of 0.33 K/s). The as-quenched $Zr_{55}Al_{10}Ni_5Cu_{30}$ alloy rod was sliced into 0.5-mm-thick disc samples that were initially annealed at 685K for 120s at heating and cooling rates of 0.17K/s, denoted as R_{Exp}^* . The fully relaxed samples were then annealed again at 735K (=1.07T_g) for 120s at a heating rate of 0.33K/s and then quenched at cooling rates of $R_{\text{Exp}} = 0.33$, 0.83, 1.7, 3.2, or 4.4K/s (Ref. [24](#page-8-8)). The excess enthalpy generated during the second annealing process ∆*H* is calculated as $\Delta H = \int_{RT}^{725} C_{p,s} - C_{p,q} dT$ $\frac{p_{723K}}{p_{13}}$ $C_{\rm p,s}-C_{\rm p,q}dT$, where RT is the room temperature. $C_{\rm p,s}$ and $C_{\rm p,q}$ are the specific heats of the sample reheated to 723K and the as-secondary annealed state, respectively. The specific heat measurement was conducted with a heating and cooling rates of 0.33 K/s using DSC. Figure 2(b) shows the change in the potential energy calculated by the MD simulation compared to the change in the enthalpy obtained from the experiment. The cooling rates after annealing employed in the MD simulation and experiment are normalized by the typical cooling rates, \widetilde{R}_q (=1 × 10¹² K/s) and R^*_{Exp} . (=0.17K/s), respectively. Note that due to the different alloy chemistries, the MD and the experimental

Figure 3. Schematic illustration of the energy/volume change during the initial melt-quenching (A→D) and subsequent thermal loading processes (D→G) under conditions of $T_a \gg T_c$ and $R_q^i > \tilde{R}_q$ (top left); $T_a \gg T_c$ and $R_q^i < \widetilde{R}_q$ (bottom left); $T_a > T_c$ and $R_q^i > \widetilde{R}_q$ (top right); and $T_a < T_c$ and $R_q^i > \widetilde{R}_q$ (bottom right). The equilibrium liquid is represented as a solid black line. In the top right figure, the solid red curve represents the energy/volume change under $t_a > t_c$ conditions, while the dotted red curve represents that under $t_a < t_c$ conditions. In the top and bottom left figures, the vertical dotted lines represent the fictive temperature T_f , at which the extrapolated glass line intersects the equilibrium liquid line. The aging decreases and rejuvenation increases T_f by changing the energy/volume versus temperature curve of the glass.

curves in [Fig. 2\(b\)](#page-3-0) are not directly comparable; yet the basic trend and the order of magnitude roughly match. The enthalpy obtained from the experiment increases with increasing cooling rate after annealing, and the positive enthalpy of the plots for $R_{\text{Exp}} = 0.83$, 1.7, 3.2, and 4.4K/s indicate that thermal rejuvenation was achieved. This is clear evidence of the feasibility of thermal rejuvenation via thermal loading, provided the necessary conditions are satisfied. The rough matching between MD and experimental curves in [Fig. 2\(b\)](#page-3-0) implies that the level of rejuvenation is dominated by a cooling rate ratio between initial melt-quenching and final quenching, rather than the absolute magnitude of the cooling rate.

[Figure 3](#page-4-0) shows a schematic of the energy/volume change during the initial melt-quenching $(A\rightarrow D)$ and subsequent thermal loading processes (D→G). (1) If the annealing temperature T_a is much higher than T_c ($T_a \gg T_c$), then an amorphous metal may quickly "relax" to an equilibrium liquid state within the heating stage (green curve in [Fig. 3,](#page-4-0) top and bottom left) before the annealing process starts. As the equilibrium liquid is cooled (F→G), sooner or later the liquid deviates from the equilibrium liquid state and eventually the structure is frozen into a glass. A cooling rate R_q^i higher than \widetilde{R}_q induces an earlier deviation from the equilibrium liquid and leads to a rejuvenated glass with a higher energy/volume (red curve in [Fig. 3](#page-4-0), top left), while a cooling rate lower than \widetilde{R}_q leads to an aged glass (blue curve in [Fig. 3](#page-4-0), bottom left).

Figure 4. The change in the topological order induced by the thermal loading process. (**a**) Change in the fraction of the icosahedral atomic cluster through thermal loading with different final cooling rates R_q^i . The as-quenched model has an icosahedral cluster fraction of 0.081. (**b**) Change in the average cluster coefficient of the icosahedral MRO \overline{X} during the thermal loading process (D→G) with different final cooling rates R_q^i . The \overline{X} value of the as-quenched model is 0.327. The four figures to the right of the graph represent the MRO network structure^{[27](#page-8-11)} of the annealed models. The final cooling rate and annealing temperature are shown at the bottom left of each figure.

(2) If T_a is slightly above T_c ($T_a > T_c$), then the annealing time t_a becomes a key factor in addition to the cooling rate R_q^i because the relaxation time to the equilibrium liquid is considerable compared with the heating and annealing times. Thus, during isothermal annealing the energy/volume increases to approach that of the equilibrium liquid. We actually observed the energy increase during the isothermal annealing process under $T_a > T_c$ conditions (see Supplementary Information, Fig. S4). As shown in Fig. 3 (top right), a longer annealing time leads to a larger energy/volume increase during isothermal annealing and eventually to a rejuvenated glass after cooling (solid red curve), while a shorter annealing time may lead to an aged glass (dotted red curve). Therefore, there is a critical minimum annealing time for rejuvenation t_c ; rejuvenation is realized only if $t_a > t_c$. (3) If T_a is below T_c ($T_a < T_c$), then the annealing-cooling process always leads to aged glass as shown in [Fig. 3](#page-4-0) (bottom right). In [Fig. 3](#page-4-0) (top and bottom left), the fictive temperature T_f (Ref. [25\)](#page-8-9) is also indicated, at which the extrapolated glass line intersects the equilibrium liquid line. In a similar manner to T_g , glass produced at a higher cooling rate from a high temperature liquid has a higher fictive temperature and vice versa²⁶.

Change in the topological order by rejuvenation. We investigated the change in the topological order induced by the thermal loading process. The internal structures of amorphous metals, such as the short-range order (SRO) and medium-range order (MRO) have been attracted much attention, because they are correlated with the intrinsic properties of amorphous metals. [Figure 4\(a\)](#page-5-0) shows the change in the fraction of icosahedral SRO, which has an energetically stable, high packing structure and is known to be a core amorphous metal structure. The figure shows clearly that aging increases and rejuvenation decreases icosahedral SRO.

[Figure 4\(b\)](#page-5-0) shows the topological MRO composed of icosahedra. To characterize the MRO network structure, we employ the MRO clustering coefficient parameter \overline{X} , which was employed in our previous work²⁷. For the center atom of the icosahedron α , the clustering coefficient parameter X^{α} can be defined as, $X^{\alpha} = 2L^{\alpha}/\{n_{B}^{\alpha}(n_{B}^{\alpha} - 1)\}\$, where n_{B}^{α} is the icosahedral bond number, and L^{α} is the number of triangular geometries. As \overline{X} is the average clustering coefficient parameter over all icosahedra in an amorphous model, a large \overline{X} value implies a highly developed MRO network structure. As shown in [Fig. 4\(b\)](#page-5-0), \overline{X} increases during the thermal loading process (D→G) when the annealing temperature T_a is below 1.1 T_g or the cooling rate is below \widetilde{R}_q . In contrast, although some fluctuation is observed, \overline{X} decreases when T_a is above 1.1 T_g and the cooling rate is above \widetilde{R}_q .

Together with [Fig. 2\(a\)](#page-3-0) and [Fig. 4,](#page-5-0) we conclude that aging increases and rejuvenation decreases the topological SRO and MRO, which are typical characteristics of glass structures. The clear correspondence between the potential energy profile in [Fig. 2\(a\)](#page-3-0) (and the volume profile in Supplementary Information,

Figure 5. The change in the mechanical properties induced by the thermal loading process. (**a**) Change in the elastic stiffness coefficients C11⁵_a and C44⁵_a during thermal loading (process D→G) at a final cooling rate of R_q^5 and various annealing temperatures T_a . The elastic stiffness coefficients of the as-quenched model with a cooling rate \widetilde{R}_q (state G) are $\widetilde{C}11_q = 171.9$ GPa and $\widetilde{C}44_q = 57.5$ GPa; these are adopted as reference values. The elastic stiffness coefficient of the as-quenched model with a cooling rate \tilde{R}_q^5 are shown by the dashed lines. (**b**) The shear stress change during the shear deformation tests for four models; two asquenched models (state D) with different cooling rates, \widetilde{R}_q and \widetilde{R}_q^5 (> \widetilde{R}_q), and two annealed models (state G) with different annealing temperatures, T_g (aged model) and 1.67 T_g (rejuvenated model) with the same cooling rate of after annealing, $R_0^5 (= \widetilde{R}_q^5)$. Inset figures represent snapshots of the two annealed models colored by local Mises strain at the same $\gamma = 0.4$ engineering shear strain.

Fig. S3) and the SRO and MRO in [Fig. 4](#page-5-0) indicates that aging and rejuvenation change the macroscopic structural properties of amorphous metals, such as the free volume, by changing the internal topological order.

Change in the mechanical properties by rejuvenation. To demonstrate the possibility of controlling the mechanical properties via aging and rejuvenation, the change in elastic stiffness due to thermal loading (process D→G) was investigated. The elastic stiffness coefficients of the as-quenched model with a cooling rate \widetilde{R}_q (state D in [Fig. 1](#page-2-0)), $\widetilde{C}11_q$ and $\widetilde{C}44_q$, were calculated from the second spatial derivative of the total potential energy $\widetilde{E}_q(0)$; these $\widetilde{C}11_q$ and $\widetilde{C}44_q$ values are taken as the reference. The elastic stiffness coefficients of the models annealed at T_a and with a final cooling rate R_q^5 (state G) were also calculated, and the change in the elastic stiffness coefficients due to thermal loading (process D→ G), C11⁵ and C44⁵_a, are shown Fig. 5(a). Both C11⁵_a and C44⁵_a increase in the case of aging, while they decrease in the case of rejuvenation. Moreover, both values saturate in the high T_a region and take values of $\tilde{C}11_q^5$ and $\tilde{C}44_q^5$, respectively, which are the elastic stiffness coefficients of the as-quenched model with a cooling rate of \tilde{R}_q^5 (= R_q^5). The clear correspondence between the potential energy profile (*i* = 5) in [Fig. 2](#page-3-0) and the elastic stiffness coefficient profile in [Fig. 5\(a\)](#page-6-0) shows the possibility of controlling the elastic properties of amorphous metals via thermally induced rejuvenation and aging. The correlation between SRO in [Fig. 4\(a\)](#page-5-0) and elastic stiffness in [Fig. 5\(a\)](#page-6-0) in this study agrees with that found in our previous studies $27, 28$ $27, 28$, where the elastic constant increases with increasing icosahedral SRO.

Plastic deformation behavior of the "aged model" and "rejuvenated model" was also investigated by simple shear MD deformation tests, in which the models were applied incremental affine shear strain at 0K. At each strain step, the atomic structure and simulation box were relaxed using the conjugate gradient method. [Figure 5\(b\)](#page-6-0) shows the shear stress change during the shear deformation tests for four models; two as-quenched models (state D) with different cooling rates, \tilde{R}_q and $\tilde{R}_q^5(\geq \tilde{R}_q)$, and two annealed models (state G) with different annealing temperatures, $T_{\rm g}$ (aged model) and 1.67 $T_{\rm g}$ (rejuvenated model) with the same cooling rate of after annealing, $R_9^5 (= \tilde{R}_9^5)$. All the model shows a maximum shear stress at around $\gamma = 0.1$ engineering shear strain, and the peak shear stress increases by the thermal loading process D→G with the annealing at $T_a = T_g$ and decreases with the annealing at $T_a = 1.67 T_g$. The two figures of atomic structure in [Fig. 5\(b\)](#page-6-0) represent snapshots of the annealed models at $\gamma = 0.4$ engineering shear strain, in which atoms are colored by Mises strain²⁹. The aged model shows an inhomogeneous deformation, in which Mises strain is localized into band region parallel to shear direction, while the rejuvenated model shows more homogeneous deformation. The clear difference in deformation mode

between the aged and rejuvenated models demonstrates that the aging and rejuvenation can change the plastic behavior of amorphous metals. It is well known that deformation at \approx 2% uniaxial strain of well-aged amorphous metals at room temperature is severely localized into narrow band region, so called shear banding, and the shear banding severely limits the macroscopic deformability of amorphous metals because of the local meltin[g30–33](#page-8-14). The homogeneous deformation in the rejuvenated model suggests that the rejuvenation conduces better macroscopic deformability to amorphous metals. Tensile ductility of amorphous metal induced by mechanical rejuvenation has been reported in a recent experimental study³⁴.

We also experimentally observed the change in the mechanical properties via rejuvenation through a micro Vickers hardness test (see Supplementary Information, Fig. S5).

Recently, the critical fictive temperature concept T_{fc} for plasticity in amorphous metals has been proposed¹³. The critical fictive temperature is a characteristic of amorphous metals: an amorphous metal is ductile in the case of $T_{\text{fc}} < T_f$ and brittle in the case of $T_{\text{fc}} > T_f$. Since rejuvenation increases T_f as schematically shown in [Fig. 3](#page-4-0), it would improve the ductility of amorphous metals through the prevention of catastrophic shear localization behavior such as shear banding. We actually demonstrated that rejuvenation enhances more homogenous deformation in [Fig. 5\(b\),](#page-6-0) which supports a fictive-temperature-based discussion of ductility in amorphous metals.

Conclusion

In summary, we have conducted thermal loading MD simulations of an already-formed amorphous metals subjected to heating, isothermal annealing, and second quenching and have produced a rejuvenation map with respect to the annealing temperature and second quenching rate that allows conditions to be found to control the level of rejuvenation. We found that thermal rejuvenation occurs via a thermal processing of isothermal annealing at temperatures above $1.1T_g$ and subsequent quenching at a cooling rate that is higher than that of the initial quenching process. The level of rejuvenation increases with increasing annealing temperature and quenching rate. We also experimentally demonstrated control of the level of thermal rejuvenation in $Zr_{55}Al₁₀Ni₅Cu₃₀$ bulk glassy alloy. The above results are qualitatively consistent with analytical models of fictive temperature evolution in quenching and heating glass DSC analysis²⁶, but now adding a constructive notion of applying the heating-quenching process *multiple* times to engineer glass properties, with quantitative proofs that the scheme actually works.

The fact that rejuvenation is realized via annealing even slightly above T_g and subsequent quenching is important for modifying the glass properties after the material has been cast and processed into near component shape because amorphous metals are easy to handle at temperatures slightly above T_g because of their high resistance to shape changes and/or crystallization.

One would have the flexibility of applying the thermal rejuvenation very locally, using for example transient laser heating; if the laser-heated spot is very small, the local cooling rate afterwards could be much higher than with the entire glassy block cooling down from melt. In practical thermal loading, the achieved level of rejuvenation will vary across the sample because heterogeneous heat transfer and heat conduction over time and space are expected. If we need to demonstrate the details of the heterogeneity in an actual heat treatment process, then we could employ a coarse-grained heat transfer and temperature distribution analysis such as FEM in combination with our rejuvenation map. This analysis framework could be a useful engineering tool for predicting heterogeneous rejuvenation and engineering the desired rejuvenation distribution while avoiding crystallization and/or gross shape changes with certain boundary conditions. Recently, nanoimprinting of metallic glass³⁵ has been developed using a thermoplastic forming process slightly above T_g , which allows the realization of low-cost fabrication of microand nanodevices. Our study proposes that a thermoplastic forming process consisting of annealing slightly above T_c and then rapid cooling before the glass relaxes into equilibrium liquid would avoid embrittlement by aging in the thermoplastic forming process.

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Acknowledgments

This work was supported by Grants-in-Aid for Challenging Exploratory Research (Nos. 23656418 and 25630013), Scientific Research (A) (No. 23246025), and the Elements Strategy Initiative for Structural Materials (ESISM). J.L. acknowledges support by NSF DMR-1240933 and DMR-1120901.

Author Contributions

M.W. performed the molecular dynamics simulations and J.S. conducted the experiments. J.L. and S.O. assisted with data interpretation. M.W. and S.O. designed, and M.W., J.L. and S.O. wrote the paper. All authors discussed the results and participated in manuscript editing.

Additional Information

Supplementary information accompanies this paper at <http://www.nature.com/srep>

Competing financial interests: The authors declare no competing financial interests.

How to cite this article: Wakeda, M. *et al.* Controlled Rejuvenation of Amorphous Metals with Thermal Processing. *Sci. Rep.* **5**, 10545; doi: 10.1038/srep10545 (2015).

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