Atomic picture of elastic deformation in a metallic glass

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The tensile behavior of a Ni₆₀Nb₄₀ metallic glass (MG) has been studied by using ab initio density functional theory (DFT) calculation with a large cell containing 1024 atoms (614 Ni and 410 Nb). We provide insight into how a super elastic limit can be achieved in a MG. Spatially inhomogeneous responses of single atoms and also major polyhedra are found to change greatly with increasing external stress when the strain is over 2%, causing the intrinsically viscoelastic behavior. We uncover the origin of the observed super elastic strain limit under tension (including linear and viscoelastic strains) in small-sized MG samples, mainly caused by inhomogeneous distribution of excess volumes in the form of newly formed subatomic cavities.

In general, the relationship between the stress and strain of materials follows the classical Hooke’s law when the stress is in the elastic regime¹. After the stress is released, the strain recovers to zero linearly without any hysteresis. The stress-strain relationship may deviate from the linear mode when creep², stress-induced phase transition³ or yielding⁴ etc., happens. Metallic glasses (MGs), a new class of engineering materials, have nearly 2% elastic strain limit, which is much higher than that of their crystalline counterparts. Moreover, it has been shown in recent experiments that this value can be even higher, for instance, about 3%, 4.4%, and 6.6% respectively for nanometer-sized MG samples cut from a bulk Zr-Ti-Co-Be MG⁵, a ribbon Cu-Zr MG⁶, and a 50 nm-film Ni-Nb MG⁷. Although much work has been done on elasticity of traditional glassy materials⁸⁻¹⁰, the above results still triggered much interest in the elastic behavior of MGs. The local heterogeneity in elastic properties within the nanoscale was observed in bulk MG samples¹¹. It was also speculated that MGs at room temperature may consist of tightly bonded atomic clusters and loosely bonded free volume regions¹². During the elastic deformation, the tightly bonded regions were assumed to undergo pure elastic response whereas the loosely bonded regions to behave like supercooled liquids, thus showing a viscoelastic feature on the whole. Using x-ray diffraction and bond orientational anisotropy analysis, Dmowski et al. demonstrated that 3/4 volume fraction of MGs deforms elastically and the rest one quarter belongs to the inelastic portion¹². More recently, Ding et al. proposed a solidity/liquidity index, based on bond orientational anisotropy to assess the varying degree of local structures in MGs upon shear stress and the heterogeneous elasticity¹³⁻¹⁵ by classical molecular dynamics (MD). However, the atomistic mechanism underlying the superior elastic properties of MGs is still not fully understood. How to separate the solid-like regions from the liquid-like regions on the atomic scale? What difference in elastic responses between them? What is the major factor behind to control these? Such questions still remain and thus it is imperative to investigate how the glassy structures respond to the stress on the atomic level in MGs in the elastic regime. In this paper, we apply the ab initio density functional theory (DFT) to calculate a Ni₆₀Nb₄₀ MG model containing 1024 atoms (an extremely large number for DFT calculations) upon tension and trace the structural evolution of atoms with increasing tensile strain from 0 to 7%. It is found that not only single atoms but also major polyhedra have inhomogeneous responses to external tensile stress when the strain is over 2%. These heterogeneities suggest that the elasticity of MGs becomes viscoelastic by forming new cavities that evolve with external tensile stress, resulting in the observed super elastic strain limit in small-sized MG samples.

Fig. 1a illustrates the comparison in structure factor S(q) and pair distribution function G(r) between the simulation data calculated from the fully relaxed configuration under zero stress and the experimental data extracted from high energy x-ray diffraction (100 keV) on the BW5, Hamburg, using a ribbon sample prepared.
by melting spinning. More experimental details are given in Supplementary Materials (SM). Due to time scale restriction, the simulation-produced sample cannot reach the atomic relaxation level in the ribbon sample. However, the general structure features of the real-world MG can be captured (e.g., peak position and shape), suggesting that the simulation data could be still reliable. Fig. 1b shows the stress vs. strain curve of Ni-Nb MG under loading and unloading processes. In the loading process, when the macro-strain is less than 2%, the stress increases almost linearly with the strain. However, when the strain is further increased to 7%, the stress-strain relationship deviates from the linear mode, exhibiting an obvious stress release, which is close to our experimental value 6.6%7.

Figure 1 | (a) Comparison in $S(q)$ and $G(r)$ (inset) extracted from XRD measurement (solid line) and VASP simulation (open circle) for the Ni$_{60}$Nb$_{40}$ MG, respectively, showing good agreement between them. (b) Stress-strain curve for a Ni$_{60}$Nb$_{40}$ MG upon loading and unloading produced by ab initio DFT simulations.

We calculate the atomistic local strain by comparing the changes in atomic positions between the present and a reference state. The local shear strain invariant $\eta^{\text{diss}}$ of each atom in the sample is calculated using equation$^{16}$: $\eta^{\text{diss}} = \sqrt{\frac{1}{2} \text{Tr}(\eta - \eta_{\text{ref}})^2}$, where $\eta$ and $\eta_{\text{ref}}$ are local Lagrangian strain and local hydrostatic strain for that atom, respectively. To calculate the $\eta^{\text{diss}}$ of each atom, we use a reference configuration that has 1% strain lower. The values of $\eta^{\text{diss}}$ can reveal whether the local shear strain distribution is homogeneous in the elastic regime for all atoms. Fig. 2a shows the distribution of von Mises strain for the total atoms with increasing tensile strain. When the tensile strain increases to 2%, the distribution of von Mises strain for all atoms remains localized, indicating that most atoms have the same local shear strain around 0.008. It is fully consistent with the 2% elastic strain limit for MG bulk samples. After that, the distribution of von Mises strain becomes delocalized, spanning a wide range from 0.002 to 0.015 when the tensile strain reaches about 7%. These results point the fact that the atoms in the sample have significantly diverse responses to the external tensile stress when the tensile strain is larger than 2% and the sample mainly undergoes viscoelastic deformation.

It is commonly believed that metallic glasses consist of different types of polyhedra, which have a wide distribution$^{17}$. In the dense random hard sphere model, the hard spheres closely pack together and form solid-like regions. Naturally, the volumes with disconnected cavities make up the void regions. The deformation behavior of MGs was demonstrated to have close relations with their inside free volume or open volume changes$^{18,19}$. By using the algorithm proposed by Sastry$^{20,21}$ (more details seen in SM and Fig. S1), the open volume regions in Ni$_{60}$Nb$_{40}$ alloy can be evaluated by setting suitable exclusion radii (about 1.39 times Schmidt radii) for Ni and Nb atoms, respectively. In doing so, only those relatively large cavities are enumerated and the tiny ones, e.g. tetrahedral and octahedral cavities, are excluded. Thus, in the so-called “solid” regions, no cavities are involved. By this means, the atoms can be grouped as locating in the “solid” and cavity regions. As shown in Fig. 2b, with increasing tensile strain, the distribution of von Mises strain of atoms either in the “solid” or cavity regions becomes broader, indicating the increased local heterogeneities with external strains. However, the
main peak of the von Mises strain for the atoms in the cavity regions moves to the large strain values, while for those in the “solid” regions shifts to the low ones. This result demonstrates that most atoms are involved in the inelastic deformation when the strain is larger than 2%, whereas the atoms in the cavity regions have slightly larger local strain than those in the “solid” regions. This is consistent with local atomic displacements of atoms changing with tensile strain in both regions, as shown in Fig. S2.

Figs. 3a–c demonstrate that the total number of atoms near the cavities increases with increasing tensile strain, especially for the strain from 3% to 7%. The fraction of atoms near the cavities is estimated to be about 32.7% at zero strain, about 35.1% at 3% strain and then rapidly increases to 41.0% at 7% strain. Compared to 25% atoms undergoing inelastic deformation estimated in Ref. 13, our results demonstrate that this value could be even larger in the small-sized MG samples. It is also found that the tensile stress applied favors the formation of new voids after 3% strain. The larger the tensile strain, the more voids are formed in response. During deformation, the formation of new voids is a dynamical process associated with the annihilation of existing ones, and is found to be randomly but inhomogeneously distributed. Some of them are inclined to form near the existing ones and accumulate to form relatively big cavities when the strain is larger than 3%. Fig. 4a shows the total volume change of the simulation cell and Fig. 4b exhibits the cavity volume changes during loading and unloading. Compared with the stress vs. strain curves in Fig. 1b, the total volume change exhibits almost the similar variation trend. However, some difference still exists. The total volume change deviating from the linear mode is not so significant when the strain is from 2 to 4%. Upon unloading, the total volume can be nearly recovered while the rest strain still remains about 1.6%. In contrast, the cavity volume change is more consistent with the stress-strain curves in Fig. 1b. During the loading process, the cavity volume increases linearly in the pure elastic strain range. It increases much faster than predicted by the linear relationship and can be fitted by an exponential function when the deformation is in the inelastic regime (2% to 7% strain). Due to the creation of more cavities, viscoelasticity becomes dominant. Upon unloading, cavity volume and stress (in Fig. 1b) almost linearly decrease with strain, but the residual cavity volume in the sample is always larger than the value at the same strain during loading. Fig. 4c exhibits a trend that the cavity volume increase results from both the increases of cavity number and size with increasing tensile strain. Our results obtained here indicate that the viscoelasticity of MGs under tension is mainly governed by fast proliferation and slow annihilation of excess volumes in the form of subatomic cavities at high stresses.

To determine the response of local atomic environment to stress, the Voronoi tessellation method22 is adopted with a cutoff of 3.6 Å, corresponding to the first minimum in pair correlation function (PCF) curves. At the same time, different types of atomic clusters are traced to see how they deform upon tension. Figs. 5a–b illustrate the fraction changes of Voronoi polyhedra centered by Ni in both “solid” and cavity regions with increasing strain. The evolution of polyhedra centered by Nb atoms can be seen in Fig. S3. It is found that the fractions of major polyhedra in the “solid” regions on the whole do not change so drastically as opposed to those near the cavities. The top three types of polyhedra centered by Ni atoms in the “solid” regions
are successively $<0,2,8,2>$, $<0,2,8,1>$ and $<0,0,12,0>$, as shown in Fig. 5a, each of which has a population larger than 10%. With increasing tensile strain, the fraction of $<0,2,8,2>$ in the solid regions tends to decrease while the fraction of $<0,0,12,0>$ remains almost unchanged. These can also be reflected from the large distortions for $<0,2,8,2>$ polyhedra and less distortions of perfect $<0,0,12,0>$ icosahedra induced by stress, as shown in Fig. 5c, consistent with the reports in Refs. 23,24. Interestingly, the fractions of $<0,2,8,1>$ and $<0,3,6,4>$ centered by Ni atoms near the cavity region in Fig. 5b significantly increase, while those of $<0,2,8,2>$ and $<0,0,12,0>$ are reduced accordingly in comparison to the "solid" regions (see Fig. 5a). These results demonstrate a structural heterogeneity in atomic structures in MG from "solid" to cavity regions. An opposite evolution trend is found in the fractions of $<0,2,8,2>$ and $<0,3,6,4>$ polyhedra, as reflected from the trajectory of atoms analysis in Fig. 5d, in which two cluster types $<0,2,8,1>$ and $<0,3,6,4>$ coexist at zero strain. At 6% stain, the structure of polyhedron $<0,2,8,1>$ does not change much, while the $<0,3,6,4>$ evolves into the $<0,2,8,2>$ via the expansion and breaking of A–B bonds, and simultaneous contraction and formation of C–D bonds. As expected, bond orientational anisotropy is observed in the elastic regime for Ni60Nb40 MG.

Figure 5 | Fractions of major Voronoi polyhedra centered by Ni atoms (a) in the “solid” regions and (b) near the cavities change with tensile strain. (c) Structure changes of icosahedron $<0,0,12,0>$ and polyhedron $<0,2,8,2>$. (d) Two clusters of $<0,2,8,1>$ and $<0,3,6,4>$ under zero strain and strain of 6%, in which bond orientational anisotropy is observed in the elastic regime for Ni60Nb40 MG.

above 2% strain is mainly controlled by the generation, evolution and annihilation of stress-induced defects, e.g., subatomic cavities. It is found that the elastic strain limit of a Cu50Zr50 MG in compression is smaller than that in tension as shown in Fig. S5a. When the stress is over 0.8 GPa, the viscoelasticity becomes easy to start under tension. Fig. S5b–c indicates that the compression decreases the total volume and also subatomic cavities, thus reducing the viscoelasticity accordingly compared to the tension. In Refs. 27 the viscoelasticity of MGs was supposed to correlate with the amount of irreversible shear transformation zones (STZs). However, the irreversible STZs most likely are the sources of viscoplastic deformation28 rather than viscoelasticity of MGs. In contrast, our results demonstrate that the subatomic cavity change could give a general picture for describing the elasticity of MGs.

Compared to their crystalline counterparts, MGs may exhibit high yield strength approaching the theoretical limits (e.g., ~$E/10$)29 due to the lack of dislocations in the material. In bulk MG samples, except the excess free volume, the probability for the occurrence of casting defects increases, which may act as stress concentration sources30. With increasing tensile stress (beyond the experimentally observed yield strength), the material usually fails in a catastrophic fashion along a major shear band. This most likely occurs at the casting defects, corresponding to a strain value usually less than 2%. This value is consistent with the linear elastic strain limit obtained in our simulations. By contrast, in nanometer-sized MG samples the chance having casting defects becomes slim. Most of the newly-formed cavities during tension are rather randomly distributed, which restrain the formation of major shear band by dispersing locally concentrated...
stresses, and thus lead the deformation into the viscoelastic regime. Consequently, nanometer-sized MGs can exhibit higher values in elastic strain limit, as observed in Refs. 5–7. Consequently, nanometer-sized MGs can exhibit higher values in elastic strain limit, as observed in Refs. 5–7. It is often restricted by the availability of precise potentials in the classical MD (see Supplementary Fig. S5) or by the limited atom numbers in the ab initio MD. In principle, DFT is a parameter-free accurate method that has been demonstrated to yield comparable results with the experiments. We build a cubic box containing 1024 atoms (614 Ni and 410 Nb) to simulate the heating and cooling processes by LAMMPS48 using an embedded-atom-method (EAM) potential of Ni-Nb with periodic boundary conditions in three directions. The sample is heated to 2400 K, and then quenched to 300 K at a cooling rate of 10° K/s with NPT ensemble. The obtained configuration is further fully relaxed using Vienna ab initio Simulation Package (VASP)49 with periodic boundary conditions but without any constrictions in cell shape and atomic positions. The box size obtained is a = 24.362 Å, b = 24.541 Å, c = 24.096 Å, ʋ = 89.3°, ʋ = 89.8° and γ = 90.03° with the final density of about 8.571 g/cm³, in good agreement with the experimental data of about 8.5 ± 0.5 g/cm³. The elastic coefficients and mechanical properties of this fully relaxed model are calculated using a strain vs. stress approach50 to obtain the bulk mechanical properties (bulk modulus K = 192.7 GPa, shear modulus G = 43.6 GPa). The module’s elastic strain is E = 121.7 GPa, averaged Poisson’s v = 0.395) including directionally dependent Poisson’s ratio based on the Voigt-Reuss-Hill (VRH) approximation, which are comparable with the experimental data of K = 174.9 GPa, G = 48.2 GPa, γ = 120 GPa and γ = 0.37 reported for a Ni34Nb66 MG. The use of directionally dependent Poisson’s ratio in uniaxial tension simulation is crucial. The model is then subjected to successive extension in the x direction with an interval of 1% elongation. At each step, the model is fully relaxed using DFT and the cell dimensions, the other directions and the volume is adjusted according to the directionally dependent Poisson’s ratio. The stress and the atomic coordinates of the model at each step are then recorded. The total time used on this project is over 2 million processor hours on Hopper machine at National Energy Research Scientific Computing Center (NERSC).

Methods

Numerical simulation, e.g., molecular dynamics (MD), is a powerful tool that can track the trajectory of atoms in MGs upon deformation51,52,53,54. However, it is often restricted by the availability of precise potentials in the classical MD (see Supplementary Fig. S5) or by the limited atom numbers in the ab initio MD. In principle, DFT is a parameter-free accurate method that has been demonstrated to yield comparable results with the experiments. We build a cubic box containing 1024 atoms (614 Ni and 410 Nb) to simulate the heating and cooling processes by LAMMPS48 using an embedded-atom-method (EAM) potential of Ni-Nb with periodic boundary conditions in three directions. The sample is heated to 2400 K, and then quenched to 300 K at a cooling rate of 10° K/s with NPT ensemble. The obtained configuration is further fully relaxed using Vienna ab initio Simulation Package (VASP)49 with periodic boundary conditions but without any constrictions in cell shape and atomic positions. The box size obtained is a = 24.362 Å, b = 24.541 Å, c = 24.096 Å, ʋ = 89.3°, ʋ = 89.8° and γ = 90.03° with the final density of about 8.571 g/cm³, in good agreement with the experimental data of about 8.5 ± 0.5 g/cm³. The elastic coefficients and mechanical properties of this fully relaxed model are calculated using a strain vs. stress approach50 to obtain the bulk mechanical properties (bulk modulus K = 192.7 GPa, shear modulus G = 43.6 GPa). The module’s elastic strain is E = 121.7 GPa, averaged Poisson’s v = 0.395) including directionally dependent Poisson’s ratio based on the Voigt-Reuss-Hill (VRH) approximation, which are comparable with the experimental data of K = 174.9 GPa, G = 48.2 GPa, γ = 120 GPa and γ = 0.37 reported for a Ni34Nb66 MG. The use of directionally dependent Poisson’s ratio in uniaxial tension simulation is crucial. The model is then subjected to successive extension in the x direction with an interval of 1% elongation. At each step, the model is fully relaxed using DFT and the cell dimensions, the other directions and the volume is adjusted according to the directionally dependent Poisson’s ratio. The stress and the atomic coordinates of the model at each step are then recorded. The total time used on this project is over 2 million processor hours on Hopper machine at National Energy Research Scientific Computing Center (NERSC).

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Additional information

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