Strong configurational dependence of elastic properties for a binary model metallic glass

Gang Duan, a) Mary Laura Lind, Marios D. Demetriou, and William L. Johnson
W.M. Keck Laboratory of Engineering Materials, California Institute of Technology, Mail Code 138-78, Pasadena, California 91125

William A. Goddard III
Materials and Process Simulation Center, California Institute of Technology, Mail Code 139-74, Pasadena, California 91125

Tahir Çağin
Department of Chemical Engineering, Texas A&M University, College Station, Texas 77845

Konrad Samwer
I. Physics Institute, University of Göttingen, 37077 Göttingen, Germany

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In this work, the strong dependence of elastic properties on configurational changes in a Cu–Zr binary metallic glass assessed by molecular dynamics simulations is reported. By directly evaluating the temperature dependence and configurational potential energy dependence of elastic constants, the shear modulus dependence on the specific configurational inherent state of metallic glasses is shown to be much stronger than the dependence on Debye-Grueneisen thermal expansion. © 2006 American Institute of Physics. [DOI: 10.1063/1.2360203]

Bulk metallic glasses (BMGs) have acquired considerable attention from scientific and technological view points in the last two decades.1–9 BMGs have very high yield strength, at least double that of ordinary commercially used crystalline materials, and high elastic strain limit, roughly 2% in tension or compression due to their disordered atomic structure.4,14–16 Upon yielding BMGs tend to form highly localized shear bands, where one observes large local plastic strains.10–12 To describe the yielding in metallic glasses a cooperative shear model13 has been developed for the glassy state based on potential energy landscape (PEL)/inherent state (IS) theory.14–16 A scaling relationship among the shear flow barrier, a universal critical yield strain, and the isoconfigurational shear modulus $G$ was constructed. The model reveals that for a fixed glass configuration the barrier height for shear flow is proportional to the isoconfigurational shear modulus $G$, which makes the elastic properties of great importance to fully understand the mechanical behaviors of BMGs. A wide variety of experimental work has been performed to measure the elastic properties of BMGs.4,17,18 Molecular dynamics (MD) simulations have been extensively utilized to calculate the elastic stiffness coefficients in crystals. In this work we present the Debye-Grueneisen thermal expansion effect as well as the strong configurational inherent state dependence of the elastic properties for a model binary metallic glass.

In order to obtain numerical results for the elastic stiffness coefficients, we perform MD simulations using an interatomic many-body Rosato-Guillope-Legrand-type potential model developed for Cu–Zr binary alloy system.21 The original simulation cell contains $N=2000$ atoms, arranged in a random bcc structure with periodic boundary conditions. At the beginning, the system was heated to 2400 K and the structure of the liquid phase was allowed to equilibrate. The system was then cooled to 50 K under zero pressure using a range of quenching rates from 1 to 10 K/ps (1 ps=$10^{-12}$ s) to generate glass configurations. This yielded the reference shapes and size matrices, $h_0$, in Parrinello-Rahman formalism22 to calculate elastic stiffness coefficients by constant temperature and constant volume ($nvt$) simulations. The glass transition temperature occurs at 700 K under the cooling rate of 5 K/ps, and there is a slight difference among the glass transition temperatures when varying the cooling rates.

Upon determining elastic stiffness coefficients, $nvt$ simulations for each state point were carried out. After equilibration of 20 000 steps (20 ps with a time step of 1 fs; 1 fs=$10^{-15}$ s), we calculated the elastic stiffness coefficients at different temperatures and at different potential energies after collecting statistics over 100 000 steps for the convergence of the fluctuation terms. The elastic stiffness coefficients were evaluated using the following statistical fluctuation formula:23

$$C_{ijkl}^T = \frac{\Omega_0}{k_B T} \left( \langle P_{ij} P_{kl} \rangle - \langle P_{ij} \rangle \langle P_{kl} \rangle \right) + \frac{2Nk_BT (\delta_{ij}\delta_{jk} + \delta_{ij}\delta_{ik})}{\Omega_0} + \langle \chi_{ijkl} \rangle,$$

where $\langle \rangle$ denotes the averaging over time and $\Omega_0=\det h_0$ is the reference volume for the model system. The first term represents the contribution from the fluctuation of the microscopic stress tensor, the second term represents the kinetic energy contribution, and the third term is the Born term. The thermal relaxation effect on the elastic stiffness constants was simulated to guarantee that the calculations performed in this work reached the steady states. For instance, at the highest temperature of 500 K the shear moduli for the samples...
RESULTS

obtained by the cooling rate of 10 K/ps are, respectively, 9.5, 8.0, and 8.2 GPa when collecting statistics by 50,000 steps (50 ps), 80,000 steps (80 ps), and 100,000 steps (100 ps) for the convergence of the fluctuation terms.

We first analyzed the Debye-Grüneisen thermal expansion effect on the elastic stiffness coefficients for the model glass under the same configuration. The calculations were carried out under zero pressure at different temperatures below the glass transition temperature $T_g$ to ensure no configurational changes happen. Figures 1 and 2 present the temperature dependence of the shear modulus $G$ and the bulk modulus $B$ for the binary Cu$_{46}$Zr$_{54}$ metallic glass prepared at two different cooling rates, 5.0 and 2.5 K/ps, respectively. The shear modulus Debye-Grüneisen slope was measured to be $[dG/dT] = -18 \pm 20$ MPa/K and the bulk modulus Debye-Grüneisen slope $[dB/dT] = -37$ MPa/K. The MD glass samples quenched using different cooling rates have different but fixed configuration states; the linear fits to the low temperature modulus data have essentially the same slope, showing that the low temperature $G$ dependence is intrinsically independent of the configurational states.

FIG. 1. Temperature dependence of shear modulus $G$ for the Cu$_{46}$Zr$_{54}$ metallic glass prepared at two different cooling rates (2.5 and 5 K/ps). The MD glass samples quenched using different cooling rates have different but fixed configuration states; the linear fits to the low temperature modulus data have essentially the same slope, showing that the low temperature $G$ dependence is intrinsically independent of the configurational states.

FIG. 2. Temperature dependence of bulk modulus $B$ for the Cu$_{46}$Zr$_{54}$ metallic glass prepared at two different cooling rates (2.5 and 5 K/ps). The two different configurational states are indistinguishable, demonstrating that $B$ is not as sensitive as $G$ to the configurational inherent state changes.

The configurational inherent state dependence of the elastic constants was further considered. By means of the same MD simulation model we calculated the elastic stiffness coefficients at five temperatures for different configurations (with different potential energies) obtained by quenching the system at different rates from the same liquid state under no pressure. It is found that at five temperatures (300, 350, 400, 450, and 500 K) the slope of shear modulus $G$ with respect to the total potential energy is nearly the same: $\sim -750$ GPa/(eV/atom) (approximately three times larger than the Debye-Grüneisen effect). The volume change at each temperature due to different cooling rates is negligible. It is noted that the volume difference at all five temperatures between the highest cooling rate of 10 K/ps and the lowest cooling rate of 1 K/ps is less than 0.15%, which means that the Debye-Grüneisen thermal expansion effect here is negligibly small.

According to PEL/IS theory, the liquid enthalpy can be separated as the vibrational and configurational contributions, $h_L = h_v + h_c$. The glass transition can be identified with the freezing of configurational inherent states. The vibrational contribution to the enthalpy is $3k_BT/2$. Therefore, to the first approximation, the configurational enthalpy can be directly obtained from the total enthalpy, which is the total potential energy in this work considering the fact that the calculations were made under no pressure. After subtracting the vibrational enthalpy from the total enthalpy, we plotted all the shear modulus data with respect to the configurational potential energy. As indicated in Fig. 3, a dramatically strong configurational dependence of shear modulus $G$ can be observed and the configurational energy slope of $G$ is noticed to be $\sim -720$ GPa/(eV/atom). This direct evaluation of configurational potential energy dependence of shear modulus $G$ for the Cu$_{46}$Zr$_{54}$ metallic glass when varying the cooling rates to obtain different configurations. This direct evaluation of the configurational inherent state dependence of shear modulus shows that $G$ strongly depends on the specific configurational changes of the glassy states, while weakly depends on Debye-Grüneisen thermal expansion effect.

FIG. 3. (Color online) Strong configurational potential energy dependence of shear modulus $G$ for the Cu$_{46}$Zr$_{54}$ metallic glass when varying the cooling rates to obtain different configurations. This direct evaluation of the configurational inherent state dependence of shear modulus shows that $G$ strongly depends on the specific configurational changes of the glassy states, while weakly depends on Debye-Grüneisen thermal expansion effect.
figurational inherent state dependence of shear modulus $G$ from MD simulations shows that $G$ strongly depends on the specific configurational changes of the glassy states while weakly depends on Debye-Grüneisen thermal expansion effect.

Recently Lind et al. experimentally measured the isoconfigurational elastic constants of the Zr$_{46.25}$Ti$_{8.25}$Cu$_{7.5}$Ni$_{10}$-Be$_{27.5}$ (Vit-4) samples using the pulse-echo overlap technique.$^{24}$ The samples were isothermally annealed and quenched near the glass transition temperature. It is found that the shear modulus $G$ has a strong dependence on annealing temperatures and, thus, on the specific configurational potential energy of the equilibrium liquid, although the low temperature dependence of $G$ of the configurationally frozen glasses shows linear temperature dependence. Our simulation results presented in this work are consistent with the experimental progress obtained from Vit-4.

In summary, the temperature and configurational dependences of elastic properties for a binary model metallic glass have been obtained from MD simulations. It is shown that the shear modulus dependence on the specific configurational inherent state of metallic glasses is much stronger than the dependence on Debye-Grüneisen thermal expansion.

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