Cooling rate, heating rate, and aging effects in glassy water

Nicolas Giovambattista, ¹ H. Eugene Stanley, ¹ and Francesco Sciortino²

¹Center for Polymer Studies and Department of Physics, Boston University, Boston, Massachusetts 02215, USA

²Dipartimento di Fisica, Istituto Nazionale per la Fisica della Materia, and INFM Center for Statistical Mechanics and Complexity, Università di Roma La Sapienza, Piazzale Aldo Moro 2, I-00185 Roma, Italy

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We report a molecular dynamics simulation study of the properties of the potential energy landscape sampled by a system of water molecules during the process of generating a glass by cooling, and during the process of regenerating the equilibrium liquid by heating the glass. We study the dependence of these processes on the cooling/heating rates as well as on the role of aging (the time elapsed in the glass state). We compare the properties of the potential energy landscape sampled during these processes with the corresponding properties sampled in the liquid equilibrium state to elucidate under which conditions glass configurations can be associated with equilibrium liquid configurations.

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One recent activity in the physics of supercooled liquids and glasses [1,2] is the search for the conditions under which a glass can be considered a liquid whose structural properties have been "frozen" during the preparation process. If the glass can be connected to a liquid state, then a thermodynamic description of the glass state can be developed [3–6]. Many routes can bring a system to an arrested disordered state [7], such as vapor deposition, pressure induced amorphization, hyperquenching, or standard cooling. Only the last two provide a continuous path from the liquid to the glass state and hence are the best candidates for studying the connection between glass and liquid configurations.

Since water can be glassified by cooling using hyperquenching techniques (i.e., with rates of the order of 10^5 K/s [8]), understanding the connection between the liquid state and glasses generated with different cooling rates is important. When the hyperquenched glass of water is properly annealed at T=130 K [8], a reproducible weak endothermic transition is observed which has been associated with the calorimetric glass transition temperature T_g . For water, the standard cooling rate is not sufficiently fast to overcome crystallization, so active debate [9–11] concerns how to relate the T_g of hyperquenched water to the unmeasurable T_g of the slowly cooled glass.

In this work we use molecular dynamics (MD) simulations to address the relation between liquid and glass configurations. We reproduce the same procedure followed experimentally to generate glasses. We use two cooling rates q_c differing by almost three orders of magnitude. We also study the heating rate and aging effects in the glass. We work in the framework of the potential energy landscape (PEL) approach, in which the 6N-dimensional configurational space—defined by the 3N center of mass coordinates and by the 3N Euler angles—is partitioned into a set of basins, each associated with a different local minimum of the PEL [12,13]. We focus, in particular, on the depth of the local minimum closest to the system point, and on the local curvatures of the PEL around the local minimum. We discover that with slow cooling rates, the glass retains a configuration very similar to a configuration sampled by the liquid at higher T, and hence all structural properties of the glass can be related to the structural properties of the liquid. In this case, a fictive temperature can be defined, the temperature at which the glass configuration is sampled by the liquid. In the case of a fast cooling rate, aging phenomena are very active already during the cooling process. Dynamics moves the configuration in regions of the PEL which are never explored in equilibrium [14]. Further aging at low temperature increases the differences between the glass and the liquid. When this is the case, the glass does not possess a structure corresponding to the equilibrium liquid at any temperature and hence it is not possible to associate a unique fictive T with the glass configuration.

We perform MD simulations for a system of N=216 molecules at fixed density, $\rho=1$ g/cm³, interacting with the extended simple-point-charge (SPC/E) potential [15], with periodic boundary conditions. Interactions are cut off at a distance of $r=2.5\sigma$ (σ parametrizes the Lennard-Jones part of the SPC/E potential) and reaction field corrections are added to account for the long-range interactions. Quantities are averaged over 32 independent trajectories. We perform three types of MD calculations: (i) cooling scans at constant rate, starting from equilibrium liquid configurations at T =300 K, (ii) heating scans at constant rate (from glass configurations at ≈5 K), and (iii) aging runs at constant aging temperature (at $T_{\rm age}$ =100 K and $T_{\rm age}$ =180 K, where significant aging effects are observed). The two cooling rates are q_c =-3×10¹⁰ K/s and q_c =-10¹³ K/s, and the two heating $q_h = +3 \times 10^{10} \text{ K/s}$ and rates K/s. We will denote by fast-quenched glass the glass obtained with the fast cooling rate and by slow-quench glass the glass obtained with the slow cooling rate. An averaged slow scan requires a simulation lasting 320 ns, close to the maximum possible by our method. This limitation prevents us from studying larger systems [17]. The location of the system on the PEL is studied by performing numerical minimizations (conjugate gradient algorithm) along the runs to estimate the closest local minimum configuration or inherent structure (IS), its energy e_{IS} , and the set of 6N eigenvalues ω_i^2 of the Hessian matrix [19]. The local curvatures around the minimum define in the harmonic approximation the mul-

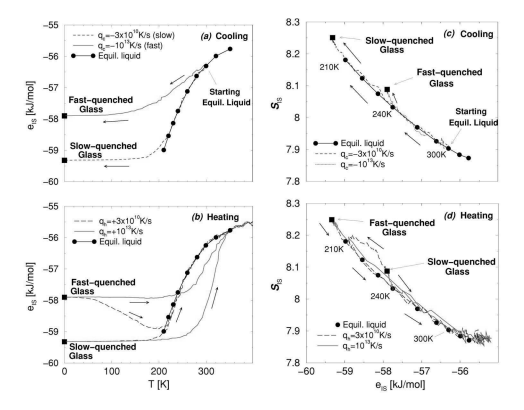


FIG. 1. (a) Cooling of equilibrium liquid configurations at T=300 K to $T\approx 0 \text{ K}$ with a fast $(q_c=-10^{13} \text{ K/s})$ and slow $(q_c=-3 \times 10^{10} \text{ K/s})$ cooling rate to generate fast-quenched and slow-quenched glasses, respectively. While during fast cooling the system is out of equilibrium already at T=290 K, slow cooling allows the system to sample the equilibrium IS as low as T=220 K. (b) Heating of the fast-quenched and slow-quenched glasses with a fast $(q_h=10^{13} \text{ K/s})$ and slow $(q_h=3\times 10^{10} \text{ K/s})$ heating rate. In the slow heating scan, the fast-quenched glass shows relaxation to deeper IS, a feature missing in the slow-quenched glass case. Fast heating shifts plots to higher T and avoids relaxation in the fast-quenched glass. (c) Basin shape function S_{IS} during fast and slow cooling. For the slow cooling rate the system is able to follow the equilibrium $S_{IS}(e_{IS})$, but for the fast cooling rate the system ends up outside the equilibrium curve. (d) $S_{IS}(e_{IS})$ during heating of the fast-quenched and slow-quenched glasses. During heating, the slow-quenched glass follows the $S_{IS}(e_{IS})$ path for equilibrium liquids. In the case of the fast-quenched glass the system approaches the equilibrium $S_{IS}(e_{IS})$ only at large times.

tidimensional parabolic shape of each PEL basin. As a global indicator of shape we use the shape function defined as

$$S_{IS} \equiv \left[\sum_{i=1}^{6N-3} \ln(\hbar \omega_i / A_0) \right] / N, \tag{1}$$

where $A_0=1$ kJ/mol. At density $\rho=1$ g/cm³ the temperature dependence of e_{IS} and S_{IS} for the SPC/E model has been previously calculated [16].

Focusing on landscape properties has the advantage that vibrational features are suppressed, and hence the remaining temperature dependence reflects structural changes. The landscape approach is particularly useful for comparing properties of different configurations. Indeed, at constant volume, in equilibrium, all landscape properties (i.e., e_{IS} , S_{IS}) are functions only of T. By eliminating T, we can establish a direct relation between S_{IS} and e_{IS} , i.e., between properties of the landscape sampled in equilibrium. A set of configurations with the same e_{IS} can be considered representative of equilibrium configurations if and only if S_{IS} satisfies the equilibrium $S_{IS}(e_{IS})$ landscape relation. We compare the relation between shape and depth during cooling scans, heating

scans, and aging with the equilibrium relation. This allows us to test if the region of configuration space explored during the scans is identical to the region explored by the equilibrium liquid at some higher temperature.

Figure 1(a) shows $e_{IS}(T)$ for the fast and the slow cooling rates, together with equilibrium values from Ref. [16]. The equilibrium data show that in the liquid state, the system explores deeper and deeper basins on lowering T and that at each T, basins with a well-defined energy depth are populated. The fact that different depths are associated with different T provides a mechanism for associating a configuration (whose e_{IS} value is known) to a fictive temperature T_f , as suggested in Ref. [20]. In the fast cooling case the liquid loses equilibrium at a very high temperature (around 290 K), i.e., when the system is still sampling basins of high energy. In the slowest cooling case, the system loses equilibrium around 220 K. For the fast cooling case, the range of T in which e_{IS} changes is significantly larger, and more energy (from the T where equilibrium is lost to the final T) is released during the cooling scan.

Figure 1(b) shows e_{IS} for the fast and slow heating rates starting both from the fast-quenched glass and the slow-

quenched glass. For the fast heating rate, to reach equilibrium T must reach a value larger than the T at which the same system lost equilibrium on cooling [Fig. 1(a)]. In the slow heating case, a different phenomenon appears for the fast-quenched glass, the system starts to explore PEL basins with lower and lower e_{IS} as it relaxes toward equilibrium, significantly changing the location of the system in configuration space.

Figures 1(c) and 1(d) show the $S_{IS}(e_{IS})$ relation for the scans reported in Figs. 1(a) and 1(b), respectively. One sees that in the slow cooling case, the cooling scan proceeds via a sequence of equilibrium configurations [since the $S_{IS}(e_{IS})$ relation coincides with the equilibrium one], i.e., basins with same depth explored during the slow scan and those in equilibrium have also the same shape. However, fast cooling generates a glass (the black square) which we see lies outside the equilibrium curve of Fig. 1(c). In this case, basins explored during fast cooling that have the same depth as those found in equilibrium have larger curvatures. During the slow heating process $S_{IS}(e_{IS})$ moves even further from the equilibrium line [Fig. 1(d)].

We now focus on the trajectory of the fast-quenched glass during aging at two different aging temperatures $T_{\rm age}$ = 100 K and $T_{\rm age}$ =180 K. For both values of $T_{\rm age}$ we find that e_{IS} is a monotonically decreasing function of aging time $t_{\rm age}$ [Fig. 2(a)], suggesting that the system changes basin by finding specific directions on the PEL which involve low energy barriers. For long $t_{\rm age}$, $e_{IS} \approx \ln(t_{\rm age})$, as found in atomic liquids [20,18]. During aging, two different behaviors are again observed for the PEL $S_{IS}(e_{IS})$ relation [Fig. 2(b)]. In the high- $T_{\rm age}$ scan, the system approaches equilibrium configurations, while for low $T_{\rm age}$ the opposite is seen. Note [Fig. 2(b)] that aging at low $T_{\rm age}$ moves the system even further from the equilibrium $S_{IS}(e_{IS})$ curve.

Next we show (Fig. 3) heating scans for different aging times after isothermal aging at $T_{\rm age}$ =100 K. Data for $T_{\rm age}$ = 180 K (not shown) are qualitatively similar. The IS energy of the starting configuration is lower the longer the time spent at T_{age} = 100 K before beginning to heat. We see [Fig. 3(c)] that e_{IS} is monotonic on heating. Figure 3(b) shows the results for the fast heating rate. In this case, T increases so fast that the system does not have time to find deeper IS. Therefore, there are no aging effects and e_{IS} grows monotonically with T. From Figs. 3(a) and 3(b), we see that for the fast heating rate, equilibrium is reached at a much higher temperature compared to the case of the slow heating rate, in full agreement with the no-aging (t_{age} =0) case, as observed in Fig. 1(b). The aging effects on $S_{IS}(e_{IS})$ are shown in Figs. 3(c) (for slow) and 3(d) (for fast) heating rates, confirming that in the slow heating case the fast-quenched glass keeps exploring new basins never seen in equilibrium before meeting the equilibrium curve.

In summary, we have shown that the fast-quenched glass significantly differs from the slow-quenched glass. To the extent that this model captures key features of real water, our work provides support to the idea that, if crystallization would not interrupt the cooling process in water, the

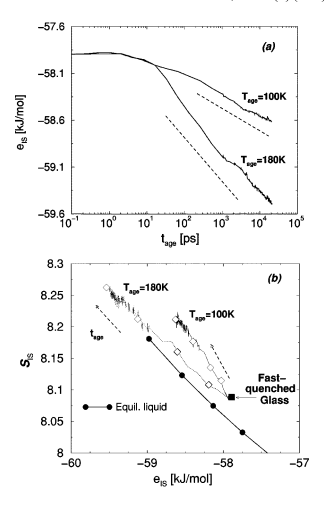


FIG. 2. (a) Time dependence of e_{IS} and (b) relation between basin shape S_{IS} and basin depth during aging of the fast-quenched glass at two temperatures $T_{\rm age} = 100$ K and $T_{\rm age} = 180$ K. For both cases, for long $t_{\rm age}$, $e_{IS} \approx \ln(t_{\rm age})$, (dashed lines). Note that during aging at low $T_{\rm age}$, the system explores basins for which the relation between shape and depth does not correspond to the equilibrium $S_{IS}(e_{IS})$ relation. At large $T_{\rm age}$, aging moves the system closer to equilibrium. At each $T_{\rm age}$, open diamonds correspond to (from right to left) $t_{\rm age} = 20$ ps, $t_{\rm age} = 20$ ps, $t_{\rm age} = 2$ ns, and $t_{\rm age} = 20$ ns.

slow-quenched glass would be characterized by a different T_g compared to the fast-quenched glass [9–11]. The analysis of simulated configurations has allowed us to clarify the differences between the slow- and the fast-quenched glass. The differences lie not only in the expected difference in e_{IS} but, more significantly, in the fact that the fast-quenched glass during cooling loses contact with the liquid state and starts to explore regions of the PEL which are never explored in equilibrium. The fast-quenched glass does not possess a structure corresponding to the equilibrium liquid structure at any temperature, and hence it is not possible to associate with it a unique fictive temperature. Finally, we have found that aging in the glass phase provides a mechanism for reducing

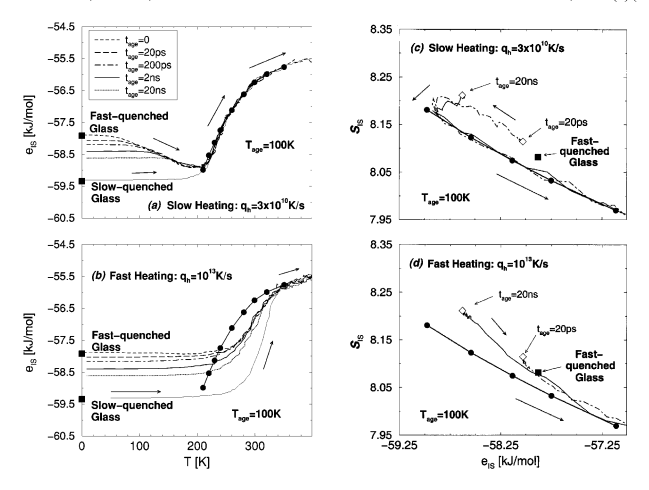


FIG. 3. Inherent structure energy e_{IS} and basin shape function S_{IS} during heating of the fast-quenched glass with (a), (c) slow and (b), (d) fast heating rates after aging for different times t_{age} at a temperature T_{age} =100 K. Results for T=180 K are qualitatively similar. Filled circles correspond to data for equilibrium liquid configurations from Ref. [16]. For comparison, in (a) and (b) we show e_{IS} during the heating scan from the slow-quenched glass. In (c) and (d), we only show S_{IS} in the heating scans after aging for t_{age} =20 ps and t_{age} =20 ns.

differences between the liquid and the glass structures only if the temperature at which the system ages is sufficiently high. Remarkably, at low $T_{\rm age}$, differences between the liquid and the glass structures appear to increase with $t_{\rm age}$.

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^[1] P. G. Debenedetti, *Metastable Liquids* (Princeton University Press, Princeton, 1996).

^[2] P.G. Debenedetti, J. Phys.: Condens. Matter 15, R1669 (2003).

^[3] T. M. Nieuwenhuizen, Phys. Rev. Lett. 80, 5580 (1998); J. Phys. A 31, L201 (1998).

^[4] M. Mézard and G. Parisi, Phys. Rev. Lett. 82, 747 (1999).

^[5] R. J. Speedy, J. Chem. Phys. 114, 9069 (2001).

^[6] S. Mossa et al., Eur. Phys. J. B 30, 351 (2002).

^[7] C. A. Angell, Science **267**, 1924 (1995).

^[8] G. P. Johari, A. Hallbrucker, and E. Mayer, Nature (London) 330, 552 (1987).

^[9] V. Velikov et al., Science 294, 2335 (2001).

^[10] G. P. Johari, J. Chem. Phys. 116, 8067 (2002).

^[11] Y.-Z. Yue and C. A. Angell, Nature (London) **427**, 717 (2004).

^[12] F. H. Stillinger and T. A. Weber, Phys. Rev. A 25, 978 (1982).

^[13] P. G. Debenedetti and F. H. Stillinger, Nature (London) 410, 259 (2001).

^[14] S. Mossa et al., Philos. Mag. B 82, 695 (2002).

^[15] H. J. Berendsen et al., J. Phys. Chem. 91, 6269 (1987).

^[16] F. Sciortino et al., Phys. Rev. Lett. 91, 155701 (2003).

^[17] A previous study for a Lennard-Jones system has convincingly shown that size effects in the supercooled state do not significantly alter statistical properties of the PEL provided that the number of particles is larger than ≈65 [S. Büchner and A. Heuer, Phys. Rev. Lett. **84**, 2168 (2000)].

^[18] L. Angelani et al., Phys. Rev. Lett. 87, 055502 (2001).

^[19] The Hessian matrix is the matrix formed by the second derivatives of the potential energy with respect to the center of mass coordinates and Euler angles of each molecule.

^[20] W. Kob et al., Europhys. Lett. 49, 590 (2000).