Intrinsic aging and effective viscosity in the slow dynamics of a soft glass with tunable elasticity

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We investigate by rheology and light scattering the influence of the elastic modulus, $G_0$, on the slow dynamics and the aging of a soft glass. We show that the slow dynamics and the aging can be entirely described by the evolution of an effective viscosity, $\eta_{\text{eff}}$, defined as the characteristic time measured in a stress relaxation experiment times $G_0$. At all time, $\eta_{\text{eff}}$ is found to be independent of $G_0$, of elastic perturbations, and of the rate at which the sample is quenched in the glassy phase. We propose a simple model that links $\eta_{\text{eff}}$ to the internal stress built up at the fluid-to-solid transition.

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A large variety of disordered soft solids exhibit fast dynamics that are associated with overdamped elastic modes activated by thermal energy. As a consequence, the fast dynamics of systems such as polyelectrolyte or colloidal gels, concentrated emulsions, and surfactant or amphiphilic copolymer phases is related to the elastic modulus, $G_0$, of the material. In addition, most of these systems exhibit a slow and non-stationary dynamics, whose origin is still under debate. One emerging idea is that the slow dynamics be due to the relaxation of internal stress built up at the fast transition from a fluid state to a solid state. In this scenario, the internal stress would result from a deformation of the local structure with respect to the ideal, relaxed configuration and thus would be proportional to the elasticity of the system. Accordingly, one may expect the slow dynamics to be intimately connected to the elastic properties of the material. However, experiments that directly test these ideas are still lacking, due to the difficulty of quenching a system into a glassy phase without perturbing the internal stress distribution. Indeed, for most soft glasses the fluid-to-solid transition is obtained upon cessation of a large preshear, which certainly influences the initial configuration of internal stress.

In this Letter, we investigate the slow dynamics of a disordered soft material, for which $G_0$ can be varied over more than one decade without changing significantly the structure, and whose dynamics can be initialized without imposing any shear. The sample is formed by a compact arrangement of soft and polydisperse elastic spheres, where the transition from a fluid to a solid state can be controlled by varying the temperature. In a previous paper, we have used multispeckle dynamic light scattering (MDLS) and linear rheology to show that this system exhibits slow dynamics, due to the rearrangement of the spheres, whose characteristic time increases as a power law of sample age, $t_w$. Here, we address explicitly the question of how the elasticity of the system influences the slow dynamics. We show that $\tau_R$, the characteristic relaxation time of the slow dynamics as measured in a stress relaxation experiment, depends only on the age of the sample and its elastic modulus. Remarkably, at all ages $\tau_R$ is found to be inversely proportional to $G_0$, thus suggesting that the slow dynamics and the aging of all samples can be described by the time evolution of a single parameter, an effective viscosity defined as $\eta_{\text{eff}} = \tau_R G_0$. We moreover show that the evolution of $\eta_{\text{eff}}$ is independent of perturbations of the elastic modulus and of the rate at which the sample is quenched in the glassy phase. We propose a simple model that relates the age-dependent effective viscosity to the relaxation of the internal stress.

The samples are surfactant lamellar phases constituted of a regular unidimensional stacking of bilayers that spontaneously roll up, resulting in a dense packing of multilamellar vesicles (MLVs) (due to their polydispersity and softness, the volume fraction of the MLVs is one). Bilayers are composed of a mixture of cetylpyridinium chloride (CpCl) and octanol (Oct) (weight ratio CpCl/Oct = 0.95) and diluted in brine ([NaCl] = 0.2M) at a weight fraction $\phi$, which ranges from 12% to 16%. The bilayers are decorated by an amphiphilic copolymer, Symperonics F68 by Serva ((EO)$_{76}$ – (PO)$_{29}$ – (EO)$_{76}$, where EO is ethylene oxide and PO is propylene oxide). The copolymer-to-bilayer weight ratio $\alpha$ ranges between 0.2 and 1.6. Upon copolymer addition, a transition from a flat lamellar phase to a MLV phase occurs. MLVs are polydisperse with a maximum size of a few microns. The MLV phase behaves mechanically as a gel with a frequency-independent storage modulus, $G'$, about one order of magnitude larger than the loss modulus, $G''$. Experimentally, we take $G_0 \approx G'(\nu = 1\text{Hz})$. A simple model relates $G_0$ to the MLV size distribution and the repulsive interaction between the bilayers. Both parameters depend on i) the interlamellar distance, set by $\phi$; ii) the amount of copolymer adsorbed to the bilayer, $\alpha$; iii) the hydrophobicity of the PO block of the copolymer, controlled by the temperature, $T$. Therefore, the elastic modulus can be varied experimentally by chang-
ing three independent parameters, $\phi$, $\alpha$, and $T$.

A temperature jump from $T = 4^\circ C$ to $T \geq 10^\circ C$ induces a fast transition from a fluid to a solid state and initializes the dynamics of the system (we take $t_w = 0$ as the time where the sample becomes a solid, as indicated by $G'(1\text{Hz}) \geq G'(1\text{Hz})$). The slow dynamics is subsequently probed by rheology measurements. The stress relaxation that follows a step strain imposed in the linear regime exhibits a very slow stretched exponential decay, $\sigma(\tau) = \sigma_0 \exp[-(\tau/\tau_R)^p]$, with $p$ in the range 0.2–0.4, depending on sample composition \cite{16}. In fig. 1 we show $\tau_R$, extracted from the stretched exponential fit of $\sigma(\tau)$, as a function of the elastic modulus $G_0$ for samples with the same age ($t_w \simeq 24000 \text{ sec}$) and surfactant weight fraction ($\phi = 16\%$), but different $G_0$. The elastic modulus was varied either by changing copolymer content $\alpha$ at a fixed temperature ($0.2 \leq \alpha \leq 1.6, T = 20^\circ C$, circles), or by varying $T$ at fixed $\alpha$ ($11^\circ C \leq T \leq 30^\circ C$, $\alpha = 0.8$, stars). Strikingly, both sets of data collapse onto a master curve, for which $\tau_R$ monotonically decreases as $G_0$ increases, thus indicating that the slow dynamics is faster for harder systems, independently of the detailed sample composition. A power law fit to the data yields an exponent $-1.1 \pm 0.1$, suggesting that the slow dynamics may be described by introducing an effective viscosity defined by $\eta_{eff} = \tau_R G_0$, whose value is independent of the composition and of the elasticity of the material over more than one order of magnitude in $G_0$.

To test the robustness of the concept of effective viscosity, we follow by linear rheology the aging of samples of various composition, and thus different elasticity (both $\phi$ and $\alpha$ were varied leading to $60 \leq G_0 \leq 700 \text{ Pa}$). Each sample is quenched in the solid phase and aged without any perturbation during a time $t_w$ before being submitted to a step strain allowing the measurement of $\tau_R$, as described previously. For all samples, we find that $\tau_R \sim t_w^{\alpha}$ with an average exponent $\alpha = 0.87 \pm 0.09$, consistently with our previous experiments \cite{16} and similarly to what observed for other soft glassy materials \cite{1, 2, 11}. Remarkably, all data collapse onto a single master curve when plotting the age dependence of the effective viscosity $\eta_{eff} = \tau_R G_0$, as shown in fig. 2. This scaling demonstrates that not only does the effective viscosity characterize the slow dynamics at a given age, but it also accounts for the aging of samples with different elasticity in a unified way.

The aging behavior of many glassy systems is deeply affected by a temperature \cite{17} or a mechanical \cite{12, 13} perturbation, leading to surprising effects such as the memory effect and rejuvenation. The question naturally arises whether the effective viscosity introduced here for the MLV phase is sensitive to a perturbation of the elastic modulus during the aging. To address this issue, we study the time evolution of the dynamics of a sample quenched from $T = 4$ to $T = 26^\circ C$ and afterwards submitted to a square wave temperature perturbation, from 26 down to 20 and back to $26^\circ C$, resulting in a variation of the elastic modulus of 20%. We follow the aging by MDLS: the multispeckle technique allows time-resolved information on the dynamics to be obtained, thus probing both the aging behavior and the instantaneous response of the soft glass configuration to the change in temperature. Figure 3 shows $g_2(t_w, t_w + t) - 1$, the two-time intensity correlation function measured at a scattering vector $q = 13.1 \mu m^{-1}$ before, during, and after the temperature perturbation. Simultaneously to the temperature jump, the correlation function drops abruptly to 0 (circles in fig. 3), thus revealing that the temperature perturbation not only modifies $G_0$, but it also significantly affects the local glass configuration. In fact, the complete loss of correlation is indicative of rearrangements on a length scale larger than $2\pi / q \simeq 0.5 \mu m$, a significant fraction of the MLV size \cite{13}. The characteristic time of the decay of $g_2 - 1$, $\tau_{DLS}$ \cite{20}, is plotted in fig. 3.
as a function of age, for the same sample as in a). Before
the perturbation, $\tau_{DLS}$ exhibits a power law growth with
sample age, as observed in previous work \[12\]. When the
sample is cooled at $T = 20^\circ \text{C}$, $\tau_{DLS}$ suddenly increases
by more than a factor of two; however, note that the
growth of $\tau_{DLS}$ follows a trend similar to that before the
temperature jump. Values of $\tau_{DLS}$ higher at $T = 20^\circ \text{C}$
than at $T = 26^\circ \text{C}$ are in qualitative agreement with the
trend observed for the relaxation time measured by rheo-
logy, since the lower $T$ the smaller $G_0$ and hence the
larger $\tau_R$. However, we note that the variation of $\tau_{DLS}$
is much larger than that of $\tau_R$ (for the former is 130%,
while for the latter is 20%). Surprisingly, when $T$ is
increased back to its initial value, $\tau_{DLS}$ drops abruptly and recovers the power law evolution it would have had
if the temperature was never changed. Similar results
have been obtained for a positive square wave tempera-
ture perturbation. Light scattering data thus indicate
that, in spite of the significant change in the local con-
figuration and in the relaxation time produced by the $T$
jump, the evolution of the dynamics is not affected by
the perturbation but rather follows an “intrinsic aging”
behavior.

It is tempting to associate this intrinsic aging to a
steady growth of $\eta_{eff}$ similar to that reported in fig. 2
for the undisturbed soft glasses. Unfortunately, time-
resolved stress relaxation experiments that would directly
confirm a continuous increase of $\eta_{eff}$ for the samples sub-
mitted to a temperature perturbation are not available.
Instead, we note that if the evolution of the dynam-
ics does not depend on the elastic history of the sam-
ple, as suggested by the MDLS experiments, the rate
at which the sample is quenched from the fluid to the
solid phase should not modify the aging. In fact, a
slow and continuous increase of $T$ may be viewed as
a series of small $T$ increments. In fig. 4, we compare
the characteristic time measured by rheology for samples
brought from 4 to 20$^\circ \text{C}$ either rapidly (“fast quench”,
heating rate $r \approx 0.64^\circ \text{C}/\text{min}$) or slowly (“slow quench”,
$r = 0.03^\circ \text{C}/\text{min}$). If $t_w$ is defined as the time elapsed af-
fter reaching the final temperature (main plot), a marked
difference exists between the aging of samples quenched
at different rates, as observed for spin glasses \[17\]. By
contrast, the data for both fast and slow quench rates
collapse onto a single curve (inset of fig. 4) when defining
the age as the time elapsed since the materials is a solid
(defined by $G' \geq G''$), as expected if the effective viscos-
ity does not depend on the elasticity of the sample or its
elastic history, but only on the time spent in the glassy
phase.

To further support the intrinsic aging scenario and
to gain insight on the physical mechanism responsible
for the growth of $\eta_{eff}$, we follow the time evolution of
$G' (\nu = 1 \text{ Hz})$ for a sample submitted to a tempera-
ture perturbation similar to that of the MDLS experiments.
At constant temperature, $G'$ slowly but continuously de-
creases with time, as shown in fig. 8. This behavior sug-
gests that the storage modulus can be written as the sum
of two contributions: the elastic modulus at equilibrium,
$G_{eq}$, and the internal stress, built up at the fast transition
from the fluid to the solid phase and whose slow relax-
ation is responsible for the aging dynamics. When the
temperature is varied from 23 to 20$^\circ \text{C}$, the storage mod-

FIG. 3: (a) Intensity correlation functions taken at age $t_w = 
7000 \text{ sec (squares), 17000 \text{ sec (circles), 35000 \text{ sec (up triangles)}
and 175000 \text{ sec (down triangles)} for a sample (with $\phi = 16\%$,
and $\alpha = 0.8$) submitted to the temperature history shown in
(b). (b) (left axis) Age dependence of the characteristic ti me
extracted from the correlation functions (the dashed line i s a
guide for the eye). (right axis) The solid line shows the tem-
perature of the sample and the symbols indicate the times at
which the correlation functions shown in (a) are measured.
(c) Age dependence of the storage modulus measured at fre-
quency $\nu = 1 \text{ Hz}$, for a sample submitted to the temperature history shown as a solid line. The dashed line is a guide for the eye.
FIG. 4: Characteristic relaxation time measured by rheology for a sample (with $\phi = 16\%$ and $\alpha = 0.8$), (circles) rapidly, (stars) slowly, brought from $T = 4^\circ$C to $T = 20^\circ$C. In the main plot the age $t_w$ of the sample is defined as usual as the time elapsed since $T$ has reached its final value, while in the inset $t_w'$ is defined as the time elapsed since the sample is solid.

ulus drops by 10%, a change in remarkable agreement with the difference of $G'$ for samples directly quenched to these two temperatures. In analogy with the results of the MDLS experiments, when $T$ is brought back to its initial value $G'$, resumes the same time evolution it would have had without any temperature perturbation. These results provide additional evidence that the aging of the MLV phase is described by the evolution of an effective viscosity that only depends on $t_w$.

In order to rationalize our experimental findings, we present a simple model that links the effective viscosity to the internal stress, $\sigma_{int}$. In this picture $\sigma_{int}$ stems from the elastic deformation of the MLVs with respect to a spherical shape, due to the random growth of closely packed MLVs at the fluid-to-solid transition. With time, the elastic deformation is slowly reduced through structural relaxations that are at the origin of the slow dynamics, leading to the aging behavior. We consider that the stress relaxation measured by rheology is due to rearrangements of regions of size $L$ (presumably containing several MLVs) that are submitted to a driving force, due to $\sigma_{int}$, and a viscous drag force. The driving force and the viscous drag scale respectively as $\sigma_{int}L^2$ and $\eta_{micro}LV$. Here, $\eta_{micro}$ is a microscopic viscosity related to the solvent viscosity and the MLVs’ volume fraction, and $V$ is the local velocity of the rearranging zone. By balancing the driving force and the viscous drag, one obtains $\sigma_{int} \sim \eta_{micro}V/L$. Because both the elastic modulus and $\sigma_{int}$ have as a common microscopic origin the elastic energy due to the deformation of the MLVs, one expects $\sigma_{int}$ to be, at all time, proportional to the equilibrium elastic modulus of the material: $\sigma_{int}(t_w) = G_{eq}/\xi(t_w)$, with $\xi(t_w)$ a dimensionless proportionality factor whose growth with $t_w$ describes the aging. Writing as suggested previously $G_0 = G_{eq}+\sigma_{int}$, one gets $\sigma_{int} = G_0/(1+\xi) \sim \eta_{micro}V/L$. Experimentally, the characteristic time of the stress relaxation is related to the time needed for a region of size $L$ to move over a distance equal to its size, $\tau_R \sim L/V$. From $\eta_{eff} = G_0\tau_R$, one finally obtains $\eta_{eff}(t_w) \sim \eta_{micro}(1+\xi(t_w))$. In agreement with experiments, this scaling shows that the effective viscosity is independent of the elastic modulus of the material and that the relaxation of the internal stress leads to an increase of $\eta_{eff}$ with sample age. The intrinsic aging reported above suggests that neither an isolated temperature jump (fig. 3) nor a series of very small $T$ increments (fig. 4) are able to relax significantly the MLVs deformation, although they change the local configuration, as indicated by the drop of $g_2\sim 1$. As a consequence, they leave essentially unchanged $\xi$ and hence $\eta_{eff}$. Increasing the amplitude or the number of $T$ jumps may eventually suppress the intrinsic aging: experiments are in progress to test this conjecture. Finally, we note that the elasticity of the MLV phase is formally identical to that of concentrated emulsions [1]: our findings should therefore be directly generalizable to the latter. Whether the same conclusions may be drawn also for other concentrated soft systems, e.g. deformable colloidal particles, remains an open issue.

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