Effect of pressure on the number of dynamically correlated molecules when approaching the glass transition

Christiane Alba-Simionesco and Cécile Dalle-Ferrier
Laboratoire Léon Brillouin, UMR 12, CEA-CNRS, 91191 Saclay, France

Gilles Tarjus
LPTMC, CNRS-UMR 7600, Université Pierre et Marie Curie, 4 Pl. Jussieu, 75252 Paris cédex 05, France

We characterize the heterogeneous character of the dynamics of liquids approaching the glass transition through an experimental determination of the number of dynamically correlated molecules \( N_{\text{corr}} \) as obtained from dynamical susceptibilities. To do so, we have obtained a new set of dielectric spectroscopy data for liquid dibutyl-phthalate on a fine and extended temperature and pressure grid, and we have used it in conjunction with high-pressure data from the literature. We have been able to evaluate the contributions to \( N_{\text{corr}} \) that are due to fluctuations associated with density and with temperature separately, thereby improving the estimate of \( N_{\text{corr}} \). We find that \( N_{\text{corr}} \) increases along the glass transition line, and more generally along any isochronic line, as pressure increases (up to 1 GPa), a result which is at odds with recent reports and theoretical predictions.

I. INTRODUCTION

A tempting explanation of the dramatic slowing down of the dynamics observed in liquids approaching their glass transition is the existence of a growing underlying length scale. The relevance of such a length, characterizing the cooperative nature of the structural relaxation, was already suggested by Adam and Gibbs in the sixties [1]. The search for length scales went in two directions [2]: on the one hand, studies of non-trivial spatial correlations in the structure, associated with some hidden order parameter or in the form of point-to-set correlations describable through the influence of amorphous boundary conditions [3] (both being essentially inaccessible in laboratory experiments); on the other hand, based on the mounting evidence in experiments and simulations of the increasingly heterogeneous character of the dynamics [4–6], investigations of a “dynamic” length describing the spatial extent of the dynamic heterogeneities. However, direct experimental measurement of the latter in molecular liquids is hard to obtain. In consequence, the corresponding results are scarce [4–8], moreover providing no information on the evolution with temperature.

A few years ago, on the basis of theoretical arguments, a new method for estimating the typical length scale of the dynamical heterogeneities and access its temperature separately, thereby improving the estimate of \( N_{\text{corr}} \). From the multi-point dynamical susceptibilities, one can extract an estimate of the “number of dynamically correlated molecules” [9, 11], which is the focus of the present work.

Specifically, the four-point dynamical susceptibility

\[
\chi_4(t) = N \langle \delta C(0,t)^2 \rangle,
\]

where \( N \) is the number of molecules in the system and \( \delta C(0,t) = \delta O(0)\delta O(t) - F(t) \) characterizes deviations from the average dynamics, quantifies the overall amount of spatial correlations between spontaneous fluctuations taking place between times \( 0 \) and \( t \). This quantity is still, unfortunately, very difficult to directly measure in experiments, especially in the case of molecular liquids. Interestingly, however, it is related through fluctuation-dissipation relations [9, 12] to other dynamical susceptibilities associated with fluctuations that are induced by an external control parameter \( x \) (such as temperature, pressure or, for colloids, volume fraction): these are three-point susceptibilities, defined as

\[
\chi_3(t) = \frac{\partial F(t)}{\partial x},
\]

which describe the response of the two-time correlation function \( F(t) \) to a change in an external parameter. Such quantities are much easier to experimentally access than \( \chi_4(t) \), as the dependence of \( F(t) \) on temperature, pressure, or any external field, may be measured by several well-developed methods such as dielectric or photon correlation spectroscopy for example [9]. It was moreover shown by computer simulation studies [12] that a good estimate for \( \chi_4(t) \) is obtained from the \( \chi_3(t) \)’s in the viscoelastic regime. (Note that another 3-point dynamical susceptibility has also been directly measured by means of an investigation of the nonlinear dielectric response [13].)

Thanks to these high-order dynamical susceptibilities, direct experimental evidence of the increase of the number of dynamically correlated molecules \( N_{\text{corr}}(T) \) while approaching the glass transition by cooling was obtained for several molecular liquids and polymers [11, 13–17]. The growth of \( N_{\text{corr}}(T) \) was found to be moderate compared to the dramatic change in relax-
ation time in the same temperature range, with $N_{corr}$ reaching a value of the order of $10^2$ at the glass transition temperature $T_g$ at atmospheric pressure. No clear conclusions could be drawn concerning the influence of the chemical nature of the system on the variation of $N_{corr}$. In particular, no correlation was observed between the growth of $N_{corr}(T)$ and the fragility of the system, i.e. the deviation of the temperature dependence of relaxation time from an Arrhenius behavior.

When cooling a liquid at constant pressure, its density increases. It is therefore useful to disentangle the effect coming from the densification of the system from that due to the decrease of thermal energy. This provides clearer insight into the physics of the slowing down of relaxation [18]. In addition, when considering estimates of the four-point dynamical susceptibility $\chi_4(t)$, it may prove important because a better description of the associated fluctuations in the $NPT$ ensemble is provided by summing the separate contributions of the density-induced and the temperature-induced fluctuations. However, except for partial results at atmospheric pressure [11], this latter procedure was never attempted. Previous pressure studies along several isobars considered the crudest bound on $\chi_4(t)$ only [16,17].

In this work, we consider the effect of density and temperature on the number of dynamically correlated molecules in a fragile glass-forming liquid, dibutylphthalate. To do so, we have measured the (linear) dielectric susceptibility along both isobars and isothersms, covering thermodynamic points on a fine grid in temperature and pressure, and combined these results with existing high-pressure data [19,20]. We have calculated the response of the linear dielectric susceptibility to temperature along isobars and to density along isothersms to provide estimates of the variation of the number of dynamically correlated molecules. To go beyond this and, as explained above, improve the estimate of $N_{corr}$ by considering separately the contributions associated with density and temperature at any state point, we have used the density-temperature scaling already shown to describe relaxation data in supercooled liquids in a large domain of temperature and density [21–25]. With the help of this scaling description, one can more easily compute the various dynamic quantities entering in the expression of $N_{corr}$. We have taken special care to minimize and to evaluate the systematic errors that come with the calculation of $N_{corr}$. Our main finding is that $N_{corr}$ significantly increases (by a factor of almost 3) along the glass transition line as one increases the pressure up to 1 GPa (the density changes by 25%). The same trend is found for other isochronic lines (i.e. lines of equal relaxation time). Contrary to previous reports [16,17] and to theoretical predictions [26,27], the extent of the spatial correlations in the dynamics is therefore not uniquely determined by the relaxation time.

![FIG. 1. Imaginary part of the dielectric spectrum of liquid DPB measured at $T = 216$ K from atmospheric pressure (leftmost curve) to $P = 250$ MPa (rightmost curve) obtained by both compression and decompression. The average pressure step between 50 and 250 MPa is about 10 MPa.](image)

**II. EXPERIMENTAL DETAILS**

**A. Sample and experimental setup**

The dibutylphthalate (DBP) sample (99% purity) was acquired from Sigma Aldrich and used as purchased. The experimental pressure dielectric setup for the dielectric measurements was described in a previous work [28]. The main characteristic of this setup is that it ensures a completely isotropic compression by the use of an external pressure fluid. For the present study, we measured a new set of data with very small pressure or temperature steps (see Fig. 1 for an illustration). This data set is fully compatible with that reported in Ref. [28] but is more complete in order to perform the detailed analysis involved in estimating $N_{corr}$. The reproducibility was always checked very carefully, especially under pressure where the experiments were carried out alternatively by compression and decompression for comparison.

The dielectric spectra were fitted by Havriliak-Negami expressions [29]:

$$\varepsilon(\omega) = \varepsilon_\infty + \frac{\Delta \varepsilon}{1 + (i\omega\tau)^{\alpha}} \gamma^\gamma$$

with $\Delta \varepsilon$, $\tau$, $\alpha$, and $\gamma$ temperature and pressure dependent adjustable parameters. This provided a convenient and faithful parametrization of all data, allowing for instance to conveniently extract the relaxation function through Fourier transformation and to take numerical derivatives with respect to the control parameters via the dependence of the adjustable parameters.
B. Thermodynamic data

Information on the thermodynamics is needed for the computation of the number of dynamically correlated molecules and it must be considered with care in order to minimize the sources of uncertainty. The PVT data was obtained from the literature and included very high pressure results [19, 30, 31]. (This has already been described in Ref. 28.) The density and temperature dependent isothermal compressibility $\kappa_T(T, \rho)$ and the thermal expansion coefficient $\alpha_P(T, \rho)$ were estimated from this data set.

Some results for the temperature dependence of the heat capacity $C_p$ of DBP are available in the literature [32, 33], but only at atmospheric pressure. We estimated from thermodynamic arguments and comparisons to other molecular liquids that the pressure variation of $C_p$ is negligible for DBP. This is in part due to the fact that the absolute value of $C_p$ is very high for DBP (about 440 $\text{JK}^{-1}\text{mol}^{-1}$ at $T_g$ and atmospheric pressure [32]). The variation for the constant volume heat capacity $C_V$ can in turn be estimated from the relation $C_V = C_p - T\alpha_P V / \kappa_T$ and is also found to be small (typically less than 10%). The jump of $C_p$ at $T_g$ is known at atmospheric pressure [32] and is about 150 $\text{JK}^{-1}\text{mol}^{-1}$. Unfortunately, no data are available for the jump of $C_V$ for these characteristic temperatures. For other molecular glass-forming liquids, one observes that the heat capacity $C_p$ of the liquid at $T_g$ slightly increases with pressure: a few % for 3-methylpentane and 1-propanol up to 200 MPa [34], a few % for m-fluoroaniline up to 400 MPa [35], and 10 – 12% for toluene up to 400 MPa [36]. However, the $C_p$ of the glass at $T_g$ has been found to also increase in 3-methylpentane and 1-propanol where it has been measured [33], so that the heat capacity jump $\Delta C_{P, g}(P)$ barely changes with pressure in these systems.

III. NUMBER OF DYNAMICALLY CORRELATED MOLECULES $N_{corr}$ ALONG ISOBARS AND ISOTHERMS

We are interested in evaluating the four-point dynamical susceptibility $\chi_4$ and the associated number of dynamically correlated molecules in the NPT ensemble. From considerations about fluctuation-dissipation relations [9, 12], two expressions may be used. The first one, into account the contributions associated with temperature (at constant volume) and density (at constant temperature) separately:

$$
\chi_4^{NPT}(t) = \frac{k_B T^2}{c_V} [\chi_4^{NVT}(t)]^2 + \rho^2 k_B T \kappa_T |\chi_4^{NPT}(t)|^2 + \chi_4^{NVE}(t),
$$

where $c_V$ is the isochoric heat capacity per molecule, relates $\chi_4^{NPT}(t)$ to the three-point-dynamical susceptibility $\chi_4^{NPT}(t) = \frac{\partial^3 \phi(t)}{\partial t^3}$, with a strictly positive residual term describing the fluctuations in the NPH ensemble (where $H$ is the enthalpy). The second expression takes

$$(3)$$

where $\chi_4^{NVT}(t)$ is the isothermal heat capacity per molecule, $\chi_4^{NVE}(t)$ is the (strictly positive) four-point-dynamical susceptibility in the NVE ensemble. For a more detailed discussion concerning these different quantities, see Refs. [11] and [12].

In simulation studies on model glass-formers [12], $\chi_4^{NPH}$ and $\chi_4^{NVE}$ were found to be small compared to the other terms in the viscous liquid regime at low temperature, becoming even negligible at the lowest temperatures. This was also indirectly confirmed by a comparison between results obtained from $(k_B T^2/c_V)|\chi_4^{NPT}(t)|^2$ and from the direct experimental measurement of a nonlinear dielectric susceptibility $\chi_4^{NVT}(t)$. It should nonetheless be stressed that, since $\chi_4^{NVE} \leq \chi_4^{NPH}$, considering the separate contributions associated with temperature and density, $(k_B T^2/c_V)|\chi_4^{NVT}(t)|^2$ provides a better approximation to the dynamical four-point susceptibility than $(k_B T^2/c_V)|\chi_4^{NPT}(t)|^2$.

In this work we focus on the numbers of dynamically correlated molecules $N_{corr}$ which may be defined as the maximum over time of the relevant time-dependent susceptibilities (or their absolute value if negative). This maximum takes place for a time of the order of the average relaxation time of the system $\tau_a$. One therefore considers $N_{corr} = \max_{t} [\chi_4^{NPT}(t)]$, $N_{corr, T} = \sqrt{k_B / c_V} T \max_{t} |\chi_4^{NVT}(t)|$, $N_{corr, T} = \sqrt{k_B / c_V} T \max_{t} |\chi_4^{NVT}(t)|$, and $N_{corr, A} = \sqrt{k_B / c_V} T \max_{t} |\chi_4^{NPT}(t)|$. From Eqs. (2) and (3), $N_{corr}^{NPT}$ may then be estimated either from

$$
N_{corr, A} \geq (N_{corr, T})^2
$$

or from

$$
N_{corr, A} \geq (N_{corr, T})^2 + (N_{corr, A})^2,
$$

with the latter expression providing a better approximation than the former one.

We have first evaluated the four-point dynamical susceptibility along isobars through the simpler formula in Eq. (2). To do so we have computed $\chi_4^{NPT}(t)$ from a numerical derivative of the Fourier transform of the Havriliak-Negami fits of the experimental dielectric data, as explained in detail in Ref. 11. We display the resulting estimate for $N_{corr, A}^{NPT}$ versus temperature $T$ on different isobars in Fig. 3(a). The results are similar to those obtained at atmospheric pressure in previous papers [11, 14, 15] and in Ref. 16; the number of
dynamically correlated molecules is found to increase when the temperature is decreased along any isobar, indicating growing spatial correlations in the dynamics as one approaches the glass transition.

We have also considered the contribution coming from the density-triggered fluctuations along isotherms, \( \text{i.e.} N_{\text{corr}}(\rho) \). To compute this term, we have numerically differentiated with respect to density the Havriliak-Negami fits of the dielectric spectra. The results are shown in Fig. 2 (b).

We observe that \( N_{\text{corr}}(\rho) \) increases with increasing density along any isotherm. The other contribution involving \( \chi_T^{\text{NVT}}(t) = \left( \frac{\partial F(t)}{\partial P} \right) / T \) is virtually impossible to get directly from the dielectric data with a good accuracy, due to the fact that data are not collected along isochores. A different method will then be considered in the following section. Note finally that the span of most curves in Figs. 2 (a) and (b) is quite restricted. In spite of the large number of state points studied and the fine \( T, P \) grid covered, the range of relaxation times that are directly accessible experimentally on given isobars and isotherms remains limited, especially at low temperature and high pressure.

IV. \( N_{\text{corr}} \) THROUGH THE SCALING DESCRIPTION OF THE RELAXATION TIME

We have already stressed that considering separately the fluctuation contributions due to density and temperature provides a crisper estimate of the number of dynamically correlated molecules \( N_{\text{corr}}(\rho) \). However, to compute the various terms required \([\text{see Eqs. (3) and (4)}]\), an alternative procedure to the direct differentiation of the experimental spectra must be employed. We have therefore relied on the, by now, well established scaling description of the density and temperature dependences of the relaxation time.

A. Density-temperature scaling law for relaxation time

The relaxation times \( \tau_\alpha \) obtained from the Havriliak-Negami fits of the dielectric spectra \( \tau_\alpha \) is equivalent to \( \tau \) in Eq. (1) are plotted versus pressure at different temperatures in the inset of Fig. 3 (a). The main panel of this figure shows that the same points, combined with literature data \([19, 20]\), can be collapsed onto a master curve when plotted versus a scaling parameter \( \epsilon(\rho)/T \): the function \( \epsilon(\rho) \) is obtained from the best collapse of the whole data set and has the physical meaning (up to an overall constant factor) of a bare or noncooperative activation energy characterizing the dynamics at high temperature \([21]\). The fact that all the \( \tau_\alpha 's \) can be plotted on a master curve,

\[
\log \left( \frac{\tau_\alpha(\rho, T)}{\tau_0} \right) = \mathcal{F}(\frac{\epsilon(\rho)}{T}) ,
\]

with \( \mathcal{F} \) a species-dependent scaling function, has been already proposed and verified for a great variety of glass-forming liquids and polymers \([22–24]\). In many cases, the function \( \epsilon(\rho) \) can be approximated by a power law, \( \rho^x \), \([22–24]\). Often, however, the density range under study is too limited to allow a discrimination between different functional forms for \( \epsilon(\rho) \). Interestingly, in the case of DBP, data are available up to 1 GPa, and it was found in Ref. \([28]\) that a simple power law does not provide a good collapse of the data over the whole density range and should be replaced by a more general function, \textit{e.g.} a polynomial function. This is illustrated in Fig. 3 (b). In such a case, the exponent \( x \) can be gener-
alized to a density dependent parameter

\[ x(\rho) = \frac{\partial \log e(\rho)}{\partial \log(\rho)}, \quad (7) \]

which varies between 1.5 and 4 for DBP over the range of density covered. As discussed elsewhere[37] (and recently confirmed in the case of “strongly correlating” liquids [38]), there are no physical reasons for why the density-temperature scaling law of the relaxation time should always be expressible in terms of a simple power law. In this respect, the study presented in this paper is more general than previous ones [16][17] that have only considered liquids and ranges of pressure for which the exponent \( x \) can roughly be taken as constant. We will see that this feature has some important consequences on the variation of the number of dynamically correlated molecules.

**B. Density and temperature contributions to \( N^{NPT}_{\text{corr,}A} \)**

The density-temperature scaling of the relaxation time provides a continuous parametrization of the latter over the whole domain of density and temperature and more specifically allows us to compute derivatives with respect to the temperature at constant density, an otherwise horrendous task experimentally. To further facilitate the computation of the various contributions to \( N^{NPT}_{\text{corr,}A} \), we have simplified the form of the relaxation function \( F(t) \) by fitting the result of the Havriliak-Negami parametrization to a stretched exponential, \( F(t) = \exp[-(t/\tau_a)^\beta] \), with \( \beta \) a state dependent parameter. This enables us to rewrite \( N^{NPT}_{\text{corr,T}} \), \( N^{NPT}_{\text{corr,ρ}} \), and \( N^{NPT}_{\text{corr,T}} \) as

\[ N^{NPT}_{\text{corr,T}} = \sqrt{\frac{k_B}{c_V}} \beta e^{-1} \left[ \frac{e(\rho)}{T} F'(\frac{e(\rho)}{T}) \right], \quad (8) \]

\[ N^{NPT}_{\text{corr,ρ}} = \sqrt{p k_B T \kappa T} \beta e^{-1} x(\rho) \left[ \frac{e(\rho)}{T} F'(\frac{e(\rho)}{T}) \right], \quad (9) \]

and

\[ N^{NPT}_{\text{corr,}A} = \sqrt{\frac{k_B}{c_p}} \beta e^{-1} \left[ 1 + x(\rho) T \kappa \right] \left[ \frac{e(\rho)}{T} F'(\frac{e(\rho)}{T}) \right], \quad (10) \]

where \( \kappa \) is the thermal expansion coefficient, \( \beta \) is the stretching parameter, \( e^{-1} \) is the inverse of Euler’s mathematical constant, which should not be confused with the activation energy \( e(\rho) \). \( x(\rho) \) is defined in eq. \( (7) \), and \( F' \) is the first derivative of the scaling function in Eq. \( (6) \). To derive the above equations, we have used the fact that the maximum over time of the various susceptibilities calculated from a stretched exponential description occurs for \( t = \tau_a \).

![Graph](image)

**FIG. 3.** -a Logarithm (base 10) of the (dielectric) relaxation time \( \tau_a \) (in units of \( \tau_0 = 1 \text{sec} \)) of DBP measured at different pressures and temperatures versus the scaling parameter \( e(\rho)/T \). The latter is adjusted to obtain the best collapse of all data points. The blue symbols denote our new set of data, the black ones our previous data from Ref. [28], and the violet ones are for literature data from Refs. [20] and [29]. Inset: All our data points plotted versus pressure. -b Bare activation energy \( e(\rho) \) obtained from the collapse of relaxation time data (dotted line). For DPB the function cannot be simply described by a power-law dependence \( \rho^x \): the full line represents the best fit with a constant \( x = 2.5 \).

Finally, \( N^{NPT}_{\text{corr,}A} \) is estimated from Eq. \( (11) \) as

\[ N^{NPT}_{\text{corr,}A} \simeq (N^{NVT}_{\text{corr,T}})^2 (1 + R), \quad (11) \]

where \( N^{NVT}_{\text{corr,T}} \) is given in Eq. \( (6) \) and \( R \) is defined as \( (11) \)

\[ R = \rho T \kappa c_V x(\rho)^2. \quad (12) \]

\( R \) is therefore a measure of the relative importance of the density-induced contribution to the number of dynamically correlated molecules at constant pressure.

Before discussing the results obtained from Eq. \( (11) \), it is worth assessing various sources of uncertainty in the calculation. We have thus compared the values of \( N^{NPT}_{\text{corr,}A} \)
and of $N_{\text{corr},T}^{\text{NPT}}$ computed from Eqs. (9) and (10), respectively, to their counterparts obtained through a numerical derivative of the Havriliak-Negami parametrization of the dielectric data (see previous section). We find that the difference between the two sets are at most of the order of 10% [which amounts to at most 20% in the square values entering in $N_{\text{corr},A}^{\text{NPT}}$: see Eqs. (4,5)], with no systematic trend with density or temperature.

Additional errors come with the account of the thermodynamic input. First, there can be a change of an overall factor for the absolute value of the $N_{\text{corr}}$'s according to whether one considers the total fluctuations of energy, enthalpy or density (as rigorously enters in the lower bound for $N_{\text{corr},A}^{\text{NPT}}$) or only that part of the fluctuations which is associated with the structural relaxation and can be ascribed to the values in excess to the glass ones (as physically motivated) [11]. For instance, as described in section II-B, $\Delta C_p$ is about 1/3 of $C_p$ for DBP at the glass transition, which, when replacing $C_p$ by $\Delta C_p$ in the estimate of the number of dynamically correlated molecules, introduces an overall multiplying factor of 3. However, we are not interested in absolute magnitudes but in relative variations with $T$, $\rho$, or $P$. Concerning the latter, the situation is much more favorable, as one can estimate that the relative change in the various thermodynamic factors is never more than 10% over the whole range of state points under study. To summarize this discussion, we conservatively conclude that a variation of the estimate of $N_{\text{corr},A}^{\text{NPT}}$ that is beyond 20 – 30% is physically significant. Smaller ones need to be interpreted with caution.

Fig. 4 shows the variation with temperature and density of the number of correlated molecules $N_{\text{corr},A}^{\text{NPT}}$ in DBP, as obtained from Eq. (11) in conjunction with Eqs. (6,7,8,12). $N_{\text{corr},A}^{\text{NPT}}$ is found to increase when the temperature decreases or when density increases, as anticipated. The number of dynamically correlated molecules obtained through this method is furthermore always slightly larger than those calculated through the other, less efficient, bound in Eq. (4) or (10).

V. $N_{\text{corr},A}$ ALONG THE GLASS TRANSITION LINE AND OTHER ISOCHRONES

From the 3-dimensional plot displayed in Fig. 4, one can follow the evolution of the number of dynamically correlated molecules along any chosen path in the $(T,\rho)$ plane. An interesting choice of paths is provided by the isochronic, i.e. equal relaxation time, lines. The most commonly considered among such lines is the glass transition line, where $T_g$ is defined by a given value of the relaxation time, say $\tau_g = 100\,\text{sec}$. It was reported in Ref. [16] for 4 glass-forming liquids and polymers that the number of dynamically correlated molecules is uniquely determined by the relaxation time. If this were true in general, $N_{\text{corr},A}^{\text{NPT}}$ should be constant along isochronic lines. As shown in Fig. 5(a), this is however not valid in the case of DBP. We find in particular that $N_{\text{corr},A}^{\text{NPT}}$ increases with pressure (and temperature) along the $T_g$ line, at least up to the highest pressure available (the evolution appears to saturate or reach a maximum for the highest pressures or temperatures considered). The same systematic trend is observed for all isochrones, although the increase in $N_{\text{corr},A}^{\text{NPT}}$ becomes smaller as the relaxation time decreases [see Fig. 5(b)]. We stress that the pressure range covered in the present study on DBP is significantly larger than in Ref. [16]: here, we consider $P$ from the atmospheric value up to 1 GPa (with a change in density of almost 25%) whereas $P$ varies from the atmospheric value to a few tens to a few hundreds MPa in [16].

Two remarks are worth making in connection with the above result. First, the stretching parameter $\beta$ of the relaxation functions was shown to be constant along any
isochnic line, and in particular along the $T_g$ line, in several glass-forming liquids [25, 28]. The same observation applies here to liquid DBP. Secondly, the isochoric fragility $m_\rho$, defined as $m_\rho = \left( \frac{\partial \log (\tau_\alpha/\tau_0)}{\partial (1/T)} \right)_\rho$, is also found to be constant along any isochoric line, as a result of the density-temperature scaling of the relaxation time [21, 37]. The important consequence is that neither the stretching of the relaxation nor the dynamic fragility are correlated with the increase in the number of dynamically correlated molecules observed along the glass transition line and other isochromes.

As the stretching parameter $\beta$ and the isochoric fragility $m_\rho$ are essentially constant, whereas $c_V$ varies by 10% along the glass transition line, it follows from Eqs. [11] and [6] that the variation of $N_{NPT \text{corr,4}}$ with pressure should be mostly due to that of the parameter $R$. This is indeed what is observed, as illustrated in Fig. 6: $1+R$ increases by a factor of almost 3 (and $R$ by a factor of almost 4), which roughly corresponds to the increase in $N_{NPT \text{corr,4}}$.

The physical significance of this increase in $R$ is that the fluctuation effects triggered by density become increasingly dominant over those associated with temperature as pressure and temperature increase. We stress that this should not be confused with the relative importance of density effects (versus temperature effects) in the slowing down of the relaxation. While $R$ concerns the fluctuations around the average dynamics, the latter deals with the average dynamics (see the Introduction) and, in the scaling description of the relaxation time, it is measured by the factor $T_\alpha P(T, \rho)x(\rho)$ [37]. For DBP along the $T_g(P)$ line, $T_\alpha P$ is equal to 0.12 at atmospheric pressure and decreases by about 25% at high pressure, whereas, as already quoted, $x$ varies from 2.5 to 4. As a result, the factor $T_\alpha P x$ changes at the glass transition from 0.30 to 0.36 over the same pressure range. At $T_g(P)$, the influence of density on the average dynamics at constant pressure is thus significantly less than that of.
temperature and increases by only 20% under pressure, a result very different from that found for $R$ in connection with the dynamical heterogeneities.

As the thermodynamic factor $\rho T_{g} c_{e} c_{V}$ varies weakly with pressure along the glass transition line, the increase in $R$ can be assigned to the change in the activation energy parameter $x(\rho)$ [compare with Eq. (12)]. The latter indeed increases from 2.5 to 4 along the $T_{g}(P)$ line, which, once squared, explains the growth of $R$ and of $N_{corr}^{NPT}$. The present study therefore highlights the role of the deviation from simple power-law behavior of the bare activation energy $c(\rho)$ in the variation of the number of dynamically correlated molecules along the glass transition line. As a check, we have repeated for another molecular glass-forming liquid, ortho-terphenyl, the very same procedure as done here. This leads to a value of the number of dynamically correlated molecules that increases along the glass transition line with increasing pressure up to 400 MPa, but only by less than 20%. Accordingly, it was found that the parameter $x(\rho)$ can be taken as essentially constant over the same domain of pressure [21, 24].

Finally, we note that our finding (for DBP up to 1 GPa) of an increase of the number of dynamically correlated molecules, and as a consequence of the associated correlation length, along the $T_{g}(P)$ line is inconsistent with the theoretical predictions made on the basis of the Random First-Order Transition (RFOT) theory [26, 27]. In the RFOT theory, the dynamic length at $T_{g}$ is predicted to be $\zeta = 5.8a$ [26, 27], where $a$ is the diameter of an elementary component of the molecule or the polymer, referred to as a “bead”. This prediction, which only depends on the relative value of the relaxation time compared to a microscopic time, is claimed to be valid for all glass-formers under any thermodynamic conditions. Some uncertainty do exist as to how beads should be defined and operational procedures have been devised to compare different systems [26, 27]. However, when looking at the same liquid under different pressures, the bead should keep unchanged, which sidesteps the fuzziness of its definition. The fact that $N_{corr}^{corr}$ increases along the glass transition line then directly contradicts the claim of the RFOT theory (except if one thinks that the relation between length scale and number of molecules should change with pressure, an a priori groundless assumption). One may of course take the variation by a factor of slightly less than 3 which is observed observed for DBP as “within the noise” of the theoretical predictions and the experimental procedures. Nevertheless, considering the care with which the various sources of uncertainty have been estimated in our calculation, it rather seems as a fact to be seriously taken into account in further theoretical and experimental investigations.

VI. CONCLUSION

In this work, we have characterized the spatial extent of the dynamical heterogeneities in a glass-forming liquid through an experimental determination of the number of dynamically correlated molecules obtained from dynamical susceptibilities. We have focused on the fragile liquid dibutylphthalate (DBP), for which, from our own measurements and from existing ones, a very large domain of temperature and pressure is experimentally covered for both dielectric spectroscopy and thermodynamics. We have calculated different estimates of the number of dynamically correlated molecules $N_{corr}^{NPT}$ including a crisp one making use of the separable contributions due to fluctuations associated with density and with temperature. We have shown that the range of the spatial correlations in the dynamics of DBP grow when approaching the glass transition along any thermodynamic path. In addition, we have found that $N_{corr}^{NPT}$ varies along the glass transition line (by a factor of about 3), and more generally along any isochronic line, as pressure (or equivalently temperature) increases, a result which is at odds with recent reports and theoretical predictions. The spatial extent of the correlations in the dynamics is further found to be uncorrelated with both the (isochoric) fragility of the system and the stretching of the relaxation function. This contradicts the naive view of the dynamics of glass-forming liquids that ties cooperativity, fragility, stretching and growing heterogeneity altogether.

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