Activation volume in superpressed glass-formers

Aleksandra Drozd-Rzoska

In pressurized glass-forming systems, the apparent (changeable) activation volume \( V_a(P) \) is the key property governing the pre提速的 behavior of the structural relaxation time (\( \tau \)) or viscosity (\( \eta \)), following the Super-Barus behavior: \( \tau(P) \propto \exp(V_a(P)/RT) \), \( T = \text{const} \). It is usually assumed that \( V_a(P) = V^a(P) \), where \( V^a(P) = RT \ln \eta(P)/dP \). This report shows that \( V_a(P) \propto V^a(P) \) for \( P \rightarrow P_g \), where \( P_g \) denotes the glass pressure, and the magnitude \( V^a(P) \) is coupled to the pressure steepness index (the apparent fragility). \( V^a(P) \) and \( V_a(P) \) coincides only for the basic Barus dynamics, where \( V_a(P) = V_a = \text{const} \) in the given pressure domain, or for \( P \rightarrow 0 \). The simple and non-biased way of determining \( V_a(P) \) and the relation for its parameterization are proposed. The derived relation resembles Murnaghan - O’Connel equation, applied in deep Earth studies. It also offers a possibility of estimating the pressure and volume at the absolute stability limit. The application of the methodology is shown for diisobutyl phthalate (Diip, low-molecular-weight liquid), isooctyloxycyanobiphenyl (8°OCB, liquid crystal) and bisphenol A/epichlorohydrin (EPON 828, epoxy resin), respectively.

Previtrous changes of the structural (primary, alpha) relaxation time (\( \tau \)), viscosity (\( \eta \)), electric conductivity (\( \sigma \)), heat conductivity (\( \kappa \)), diffusion (\( d \)) or chemical reactions rates (\( k \)) in systems ranging from low-molecular-weight liquids and polymers to liquid crystals and plastic crystals are the key manifestation of the hypothetical universal dynamics emerging on approaching the glass transition (\( T_g(P) \)). Similar patterns are observed both for the temperature and pressure path. The temperature path is associated with the Super-Arrhenius (SA) dynamics, and it is governed by changes of the apparent activation energy \( E_a(T) \), which strongly increases on approaching the glass transition temperature \( T_g \). The non-biased way of determining \( E_a(T) \) and its properties are discussed in refs 12-14 and recalled in Supplementary Info.

This report focuses on the still puzzling case of the (high) pressure-induced glass transition. For compressed glass-formers, general features of the pre提速的 dynamics are described by the Super-Barus (SB) equation:

\[
\tau(P) = \tau_0 \exp \left( \frac{PV_a(P)}{RT} \right), \quad \eta(P) = \eta_0 \exp \left( \frac{PV_a(P)}{RT} \right)
\]

where \( T = \text{const} \) and \( P < P_g \); \( V_a(P) \) denotes the apparent activation volume, which changes on compressing. Generally, the name ‘activation volume’ is reserved for the basic Barus equation with \( V_a(P) = V_a = \text{const} \) in the given domain of pressures.

Prefactors \( \tau_0 \) and \( \eta_0 \) in Eq. (1) refer to \( P = 0 \), but within the experimental error they can be approximated by atmospheric pressure values, i.e., \( \tau_0 = \tau(0) \approx \tau(P = 0.1 \text{MPa}) \) for the tested isotherm \( T \). For high-pressure studies the experimental errors are \( \Delta P \approx \pm 0.2 \text{MPa} \) (moderate pressures) and \( \Delta P \approx \pm 1 \text{MPa} \) (GPa domain). The shift of pressure by 0.1 MPa does not yield detectable changes of dielectric relaxation time.

Similar SB dependences describe pressure changes of all physical properties recalled above: pressure dependencies of \( \tau(P) \) and \( \eta(P) \) are parallel (Eq. (1))\(^{3-4} \), but for the remaining dynamic properties the translational - orientational decoupling have to be taken into account. For instance, for DC electric conductivity:

\[
\sigma^{-1}(P) = \sigma_0^{-1} \left( \frac{PV_a(P)}{RT} \right)^{1/S} \quad \text{(2)}
\]

where \( S < 1 \) is the decoupling exponent associated with the fractional Debye-Stokes-Einstein (f-DSE) dependence \( \sigma(P)|\tau(P)|^{1/S} = C = \text{const}^{10,16} \).

Institute of High Pressure Physics Polish Academy of Sciences, ul. Sokolowska, 29/37.01-142, Warsaw, Poland. Correspondence and requests for materials should be addressed to A.D.-R. (email: arzoska@unipress.waw.pl)
The first discussion regarding \( \eta(P) \) or \( \tau(P) \) behavior in compressed liquids can be associated with the relation \( \eta(P) \propto \exp(aP) \) proposed by Barus at the end of the 19th century when studying viscosity of natural oils\(^{15} \). Whalley\(^{16,17}\) and Williams\(^{18} \) applied such description for pressurized polymers and dielectric relaxation time, introducing the activation volume \( V_a \), what led to Eq. (1) with \( V_a = \text{const}. \) The Barus (B) or Barus-Williams dependence can be considered as the pressure counterpart of the basic, temperature-related Arrhenius equation which was originally introduced as \( k(T) = k_0 \exp(E/RT) \), where \( E_a \) stands for the activation energy and \( k \) is the reaction rate coefficient\(^{20} \). Generally, for ultraviscous/ultraslow glass-forming systems one should expect the Super-Barus (SB) behavior (Eqs (1) and (2)), where dynamic properties are governed by changes of the pressure-dependent activation volume: the apparent activation volume.

According to the above discussion one concludes that \( V_a(P) \) governs the dynamics of ultraviscous/ultraslow systems, and its determination and understanding is the key to the ultimate insight into the glass transition problem\(^{6–9} \), the behavior of soft matter under pressure\(^{7–21} \), high-pressure chemistry and biochemistry\(^{22} \), innovative material engineering\(^{23} \), high-pressure biotechnology\(^{24} \), geophysics, and deep Earth science\(^{25} \).

Usually, for the previtreous domain the apparent activation volume \( V_a(P) \) is calculated from \( \tau(P) \) or \( \eta(P) \) experimental data via\(^{30–35} \):

\[
V_a(P) \rightarrow V^*(P) = RT \left[ \frac{d \ln \tau(P)}{dP} \right]
\]

under the assumption that \( V_a(P) = V^*(P) \) and for \( 0.1 \, MPa < P \leq P_g \).

The analysis exploring the (apparent) activation volume determined via Eq. (3) is the key point of numerous research reports. In the framework of the transition state theory, the activation volume describes the difference between volumes occupied by a molecule in activated and non-activated states\(^{22} \). It is the essential parameter characterizing the sensitivity of the structural relaxation time or other dynamic properties to pressure changes\(^{7,8,21} \). It estimates the local volume required for a given dynamical process (in the case of \( \tau \) denotes molecular rearrangements)\(^{26–35} \). Hong \textit{et al.}\(^{36,37} \) indicated that the activation volume correlates with the length scale of dynamical heterogeneities \( \xi \), which are considered as one of the essential sources of the previtreous 'universality' of dynamic previtreous properties. Tests in hydrogen-bonded molecular liquids showed the case-sensitivity of the activation volume, determined via Eq. (3) to subtle features of molecular structures\(^{38–40} \). Worth stressing is also the broadly used link between the activation volume at the glass transition, and the fragility: \( \Delta V = m_p \times 2.303R \) \( (dT_g/dP)^{\eta \rightarrow 0} \), where the fragility \( m_p = \left[ \frac{d \ln \tau(T)}{d(T_g/T)} \right]_{T=T_g} \) is one of key 'universality' metrics for the glass transition phenomenon\(^{3,5,9,10,34,44} \). The analysis via Eq. (3) was also used for showing different activation volumes determined by dielectric and light scattering spectroscopies\(^{34,35} \). Reasonings based on such analysis can yield important checkpoints for glass transition models\(^{44–50} \). The activation volume is also significant for the thermodynamic scaling linking \( \tau(T,P,V) \) experimental data\(^{8,34,35} \). There are also reports where \( V^*(P) \) is recalled as the (apparent) activation volume, but the link to the steepness index \( m_\eta(P) \) is indicated\(^{56,57} \). For the validation of Eq. (3) reports by Whalley\(^{17,18} \) and Williams\(^{19} \) are most often cited\(^{8,30–34,38–42,48–54} \). However, these reports did not consider the Super-Barus dynamics with the pressure depending apparent activation volume but the basic Barus behavior with the constant activation volume. This issue is worth stressing, since the SB Eq. (1) directly yields the differential equation:

\[
RT \frac{d \ln \tau(P)}{dP} = V_a(P) + P \frac{dV_a(P)}{dP}
\]

Comparing Eqs (3) and (4) one obtains that generally: \( V_a(P) = V^*(P) \). The second term in Eq. (4) disappears only for two 'special' cases: (i) for \( P = 0 \), or (ii) for \( V_a(P) = V_a = \text{const} \), i.e., for the basic Barus behavior in the given pressure domain.

Worth recalling is the difference between the free volume \( V_f \) as the volume not occupied by molecules and the activation volume \( V_a \) as the volume required for the given process, for instance, the molecular rearrangement or reorientation. Then, one can expect \( V_f \approx V_a \)\(^{8,9} \).

Consequently, the question arises for the (proper) estimation of the (apparent) activation volume in the previtreous domain. This report proposes the solution to this problem and discusses the meaning and behavior of both \( V^*(P) \) and \( V_a(P) \) for \( P < P_g \). The discussion is supported by the analysis of the \( \tau(P) \) experimental data for glass-forming low molecular weight liquid \textit{disobutyl phthalate} (DIB, \( T_g(0.1 \, MPa) = 196.8K \)), epoxy resin \textit{bisphenol A/epichlorohydrin} (EPON 828, \( T_g(0.1 \, MPa) = 253.9K \)) and liquid crystalline \textit{isooctylxyanobiphenyl} (8\(^{0} \)OCB, \( T_g(0.1 \, MPa) = 220.7K \)). The latter vitrifies in the isotropic liquid phase, and the possible nematic phase is hidden below the glass transition. In given studies, pressures up to \( P \approx 1.2 \, GPa \) were reached, i.e. for the domain hardly available in high resolutions tests carried out so far\(^{3,10} \). Experimental details are described in the Methods section.

Results and Discussion

Figure 1 shows the pressure evolution of the structural relaxation time for selected isotherms for three qualitatively different glass formers \( 8^{0} \)OCB, DIB and EPON 828, in the pressure range 0.1 \( \, MPa < P < P_g \). They served for estimating both \( V^*(P) \) and \( V_a(P) \).

When discussing the physical meaning of \( V^*(P) \) one can recall the definition of the pressure-related steepness index (the normalized rate of changes of the relaxation time, viscosity, ….) in the previtreous domain\(^{8} \), which leads to the relation:
τ \tau = m_P \log P \left( \frac{P}{P_g} \right) \text{, } T = \text{const} \tag{5}

It terminates at the pressure-related fragility \( m_T(P) = m_T(P_g) = \left[ \log_{10} \tau(P)/d(P/P_g) \right]_{P=P_g} \), which is the key
metric for glass-forming ultraviscous/ultraslow systems. Then, the pressure-dependent (isothermic) coefficient
\( m_T(P) \) for \( P < P_g \) can be called the apparent fragility. Linking Eqs (3) and (5) one obtains the relation showing that
\( V'(P) \propto m_T(P) \):

\[ V'(P) = \left( \frac{RT \log_{10} \left( \frac{P}{P_g} \right)}{P_g} \right) \times m_T(P), \quad T = \text{const} \tag{6} \]

Considering further the ratio of fragilities along \( T_g(P) \) curve/line: \( m_T^{\prime}(T)/m_T(P) = \left[ \log_{10} \tau(T)/d(T_g'/T) \right]_{P=P_g} \)
\left[ \log_{10} \tau(P)/d(P/P_g) \right] = d(P/P_g)/d(T_g'/T) and linking this with Eq. (6) the following relations are obtained:

\[ m_T^{\prime}(T) = m_T(P) \left( \frac{dT_g'}{dT_g} \right) \left( \frac{P}{P_g} \right)^{-1} \tag{7a} \]

and

\[ m_T^{\prime}(T) = V'(P) \left( \frac{dT_g'}{dT_g} \right) \left( \frac{P}{P_g} \right)^{-1} \tag{7b} \]

Equation (7b), originally derived in ref.\(^{40}\), is broadly used for calculating isobaric fragilities \( m = m_T(T_g) \) for
different isobars, based on the knowledge of the \( \sim \)activation volume \( V'(P) \) calculated via Eq. (3) and the pressure
shift of \( T_g \). \(^{8,26-34,38-42,47-54} \) Notwithstanding, Eq. (7a) is fundamentally more correct than Eq. (7b), since \( V'(P) \)
should not be recalled as the apparent activation volume.

Recently, it was shown experimentally that changes of the pressure-related apparent fragility can exhibit a
universal previtreous behavior\(^{57} \):

\[ m_T(P) = \frac{A}{P^* - P} \tag{8} \]

where \( T = \text{const} \), the amplitude \( A = \text{const} \), and \( P^* \) is for the extrapolated singular pressure. Regarding pressures:
\( P < P_g \) and \( P^* > P_g \).

Following Eqs (6) and (8) one can conclude that: \( 1/V'(P) \propto 1/m_T(P) \propto P^* - P \). Such behavior is confirmed in
the insets in Figs 2–4. One of the basic features of the previtreous domain is the appearance of two dynamical
domains, i.e., regions with different SA or SB behavior remote and close to the glass transition, respectively\(^{58,59} \).

Roland\(^{60}\) showed the pressure-temperature invariance of the dynamic crossover
time-scale for a set of glass-forming liquids \( \tau(T_g, P_g) \sim 10^{-7.5 \pm 1} \) s. Until recently, the detection of \( P_g \) was associated

Figure 1. The isothermal pressure evolution of the structural relaxation time in previtreous domains of
tested glass formers recalled in the Figure. Arrows show terminals of the reliable fitting using the simple
approximation via Eq. (10a).
Figure 2. The pressure evolution of the normalized apparent activation volume $V_a(P)/RT$ (Eq. 9) and the reciprocal of $V^T(P)/RT \propto m_T(P)$ (Eqs 3 and 8; the inset) in superpressed disobutylphthlate. In the inset, the manifestation of the dynamical crossover pressure $P_B$ is indicated. Results are for $T = 317$ K isotherm. The thin dashed green line is for the Kornilov et al. moderate pressures approximation (Eq. (10b)). The thick blue curve is related to Eq. (13). The vertical arrow shows the glass transition. In the inset, the dynamic crossover pressure is indicated.

Figure 3. The pressure evolution of the normalized apparent activation volume $V_a(P)/RT$ (Eq. (9)) and the reciprocal of $V^T(P)/RT \propto m_T(P)$ (Eqs (3) and (8); the inset) in superpressed epoxy resin EPON 828. In the inset, the manifestation of the dynamical crossover pressure $P_B$ is indicated. Results are for $T = 303$ K isotherm. The thin dashed green line is for the Kornilov et al. moderate pressures approximation (Eq. 10b). The thick blue curve is related to Eq. (13). The vertical arrow shows the glass transition. The dynamic crossover pressure is also indicated.

with $\phi_T(P) = (d \log \tau(P)/dP)^{-1/2}$ vs. $P$ plot, which is parallel to $(m_T(P))^{-1/2}$ vs. $P$ presentation. Such analysis assumes a priori the validity of the pressure counterpart of the Vogel-Fulcher-Tammann (VFT) relation for describing $\tau(P)$ experimental data. Recently, an alternative and a model-free way for detecting the dynamic crossover via $1/m_T(P)$ vs. $P$ analysis was indicated. This report shows that the pressure evolution of $1/V^T(P)$ follows the pattern noted for $1/m_T(P)$; this is shown in insets in Figs 2 and 3. The lack of $P_B$ for $8^\circ$OCB in the inset in Fig. 4 results from the limited tested pressure range, between $P_g(T) = 0.56$ GPa and $P = 0.1$ MPa. Consequently, in $8^\circ$OCB measurements were carried out only in the high-pressure dynamical domain ($P_g > P > P_B$).

The above discussion shows the direct link between the magnitude considered so far as the apparent activation volume $V^T(P)$ and the apparent fragility $m_T(P)$. To determine the ‘real’ apparent activation volume $V_a(P)$ the
protocol developed for the apparent activation energy \( E_a(T) \) can be applied. The latter is given in refs\(^{12–14} \) and the Supplementary Info. For the pressure path, this means the numerical solution of the differential Eq. (4) for the given set of \( \tau(P) \) experimental data. However, for the SB Eq. (1) a more straightforward way of determining \( V_a(P) \) is possible since the prefactor in SB Eq. (1) is known:

\[
\tau_0 P = \tau(T) P = 0 \approx \tau(T) P = 0.1 \text{ MPa}.
\]

Consequently, the apparent activation volume can be calculated directly from the SB Eq. (1), via the simple rearrangement:

\[
V_a(P) = \frac{RT}{P} \ln \left( \frac{\tau(P)}{\tau_0} \right) \quad T = \text{const}
\]  

(9)

Results of such analysis present in main panels of Figs 2–4.

Insets in these Figures show pressure evolutions of \( 1/V(P) \propto 1/mT(P) \) (Eqs (3) and (8)): the inset) in superpressured liquid crystalline 8*OCB (isotropic liquid phase). Results are for \( T = 298 \text{ K} \) isotherm. The thin dashed green line is for the Kornilov et al.\(^{62} \) moderate pressures approximation (Eq. (10b)). The thick blue curve is related to Eq. (13). The vertical arrow indicates the glass transition.

![Figure 4](https://example.com/figure4.png)

Figure 4. The pressure evolution of the normalized apparent activation volume \( V_a(P)/RT \) (Eq. 9) and the reciprocal of \( V(P)/RT \propto m(T) \) (Eqs (3) and (8): the inset) in superpressured liquid crystalline 8*OCB (isotropic liquid phase). Results are for \( T = 298 \text{ K} \) isotherm. The thin dashed green line is for the Kornilov et al.\(^{62} \) moderate pressures approximation (Eq. (10b)). The thick blue curve is related to Eq. (13). The vertical arrow indicates the glass transition.
The main result of the given report is the simple and non-biased way of determining the apparent activation volume. Concluding, the activation volume is the key parameter governing the complex pretransition dynamics for the structural relaxation time, viscosity, diffusion, and reaction rates under high pressure, as indicated in the SB Eqs (1) and (2). In numerous research reports, focusing on the glass transition, this property is considered to be given by $V^*(P) = RT[\ln(\tau(P))/dP]^{9,26-35,38-42,47-55}$, and subsequently used for developing the ‘pressure dimension’ of the material engineering, high pressure preservation of food and biotechnology under pressure.

Table 1. Results of fitting $P^*$ via Eq. (13) for experimental data from Figs 2–4.

| Glass former | $P^*$ (GPa) | $C$ | $\psi$ | $P_{sl}$ (GPa) |
|--------------|-------------|-----|--------|----------------|
| DIIB         | 2.6         | 72.3| 74     | –1.1           |
| EPION 828    | 0.6         | –12.1| 27     | –0.1           |
| 8OOCB        | 1.0         | 5.5 | 28.9   | –0.21          |

where $C = \ln \Psi / \ln \tau(T), T = \text{const.}$

However, in Eq. (12) the apparent activation volume $V_a(P) \to \infty$ for $P \to 0$, and then the anomalous increase occurs for $P \to 0.1 \text{MPa}$. This problem can be avoided when taking into account that for solids, including liquids, the available range of pressures extends from the ‘normal’ positive domain (the isotropic compression) to the negative one, associated with the isotropic stretching. This concept proved its fundamental significance for the general equation of state for water or critical mixtures and blends. Angell and Quing experimentally showed the appearance of negative pressures and passing $P = 0$ without any hallmark, in glass-forming liquids using the centrifugal method. Consequently, the ‘positive’ (isotropic compressing) and ‘negative’ (isotropic stretching) pressure regions can be considered as the common area, terminating at the absolute stability limit (SL) spinodal, hidden under negative pressures. To describe experimental data in both pressure domains, one can consider the transformation $P \to \Delta P = P_P^{\text{sl}}$. Introducing the latter to Eq. (11) the modified dependence appears:

$$V_a(P) = \frac{C}{\Delta P} - \frac{\Psi}{\Delta P} \ln(P^* - P) \approx \frac{C}{\Delta P} - \frac{\Psi}{\Delta P} \left[1 - \left(\frac{P}{P^*}\right)^\psi \right].$$

The above relation describes pressure changes of the apparent activation volume in the whole tested pressure range, up to $P \approx 1.2 \text{GPa}$, as shown in Figs 2–4. It can be also extended into the negative pressures domain. Fitted parameters are given in Table 1. Values $P^*$ were estimated from insets in Figs 2–4 using the condition $1/V^*(P) = 0$; this also includes parameters $C$ and $\Psi$ estimated following ref. 7.

Consequently, the analysis of $V_a(P)$ experimental data via Eq. (13) can be reduced solely to the single fitted parameter ($P_{sl}$). This offers the new route for estimating the absolute stability limit pressure in negative pressures domain, which is considered as one of the most difficult to estimate properties via the experimental determination.

Conclusions

Concluding, the activation volume is the key parameter governing the complex pretransition dynamics for the structural relaxation time, viscosity, diffusion, and reaction rates under high pressure, as indicated in the SB Eqs (1) and (2). In numerous research reports, focusing on the glass transition, this property is considered to be given by $V^*(P) = RT[\ln(\tau(P))/dP]^{9,26-35,38-42,47-55}$, and subsequently used for developing the ‘pressure dimension’ of the glass transition models/theories: the free volume model, Adam-Gibbs model, Cohen-Grest model, and Avramov-Milchev model. This report shows that the activation volume $V_a(P) = V^*(P)$, which magnitudes coincide only for the basic Barus dynamics, i.e. for Eq. (1) with $V_a = \text{const}$. The magnitude $V^*(P)$ is directly linked to the pressure-related apparent fragility $m_a(P)$.

The main result of the given report is the simple and non-biased way of determining the apparent activation volume $V_a(P)$ via Eq. (9) and the proposal for the parameterization of $V_a(P)$ evolution given by Eq. 13. It is worth mentioning that problems with the estimation and meaning of the activation volume seem to be absent for geophysics/deep Earth science where Murnaghan – O’Connel relation is broadly applied:

$$V_a(P) = V_{a}^{ref} \left[1 + \frac{P}{K'_{0}/K'_0} \right]^{1/K'_0}$$

where $K_{0}, K'_0$ denotes 4/9 of the bulk modulus and its first derivative.

Results presented focused on the pressure evolution of the primary relaxation time, but they can also be applied for the viscosity, electric conductivity, diffusion, equilibrium, and reaction rates coefficients, in ultravis-cous/ultraslow systems what indicates the broad range of fundamental and practical applications ranging from the glass transition physics and the solid state physics to ‘extreme’ chemistry, geophysics, petrology, innovative material engineering, high pressure preservation of food and biotechnology under pressure.

Methods

In the last decades, the broadband dielectric spectroscopy (BDS) has become the key tool for studying pretransition behavior, including challenging insights from high-pressure studies. In this report, BDS is used to determine the pressure evolution of the primary (structural, alpha) relaxation time. Studies were carried out using the Novocontrol impedance analyzer, model 2015. BDS studies were carried out between the atmospheric pressure ($P = 0.1 \text{MPa}$) and the glass transition pressure, estimated via the empirical condition $\tau(T_p, P_p) = 100 \text{s}^{-1}$.
structural relaxation times were determined from the peak frequencies of primary relaxation loss curves $\epsilon''(f)$: $\tau = 1/2nf_{\text{peak}}$. Tested samples were placed in the flat-parallel measurement capacitor made from Invar. The gap between plates $d = 0.2 \, \text{mm}$ and their diameter $2r = 16 \, \text{mm}$. Samples were entirely isolated from the pressurized medium (Plexiglas). They were in contact only with Invar, quartz (the spacer between plates) and Teflon. The pressure was transmitted to the sample via the deformation of 50 mm thick Teflon film. The process was supported by the computer–controlled pump, enabling pressure changes and programming with the precision $\Delta P = \pm 0.2 \, \text{MPa}$. The pressure chamber was surrounded by a special jacket associated with the Julabo high–accuracy thermostat with the external circulation and the volume of the thermostated liquid $V = 20 \, \text{L}$. These enabled temperature changes and control with accuracy $\Delta T = \pm 0.02 \, \text{K}$. The temperature was monitored using the thermocouple within the pressure chamber and two platinum mini-resistors placed at the bottom and the top of the chamber. The high-pressure system was designed and produced by UnipressEquipment (Poland). Further experimental details are given in ref.5,7,24. Notable, that the examined range of pressures was extended up to $P = 1.2 \, \text{GPa}$, the still hardly available range in high-resolution BDS pressure studies8,26–54. Experimental results cover time-scales from $\tau(P = 0.1 \, \text{MPa})$ to $\tau(P, T) = 100s$. The latter is commonly applied as the practical empirical estimation of $(T_P, P)^{8,9,72}$.  

Data Availability

The data supporting the findings of this study are available from the author upon reasonable requests.

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A.D.-R. is responsible for all results, reasoning, and conclusions presented in the given report (100%).

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