



# New Challenges for the Pressure Evolution of the Glass Temperature

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The ways of portrayal of the pressure evolution of the glass temperature ( $T_g$ ) beyond the dominated Simon–Glatzel-like pattern are discussed. This includes the possible common description of  $T_g(P)$  dependences in systems described by  $dT_g/dP > 0$  and  $dT_g/dP < 0$ . The latter can be associated with the maximum of  $T_g(P)$  curve hidden in the negative pressures domain. The issue of volume and density changes along the vitrification curve is also discussed. Finally, the universal pattern of vitrification associated with the crossover from the low density (isotropic stretching) to the high density (isotropic compression) systems is proposed. Hypothetically, it may obey any glass former, from molecular liquids to colloids.

Keywords: glass transition, high pressures, negative pressures, melting, universality, dynamics, glass-forming ability

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Liquids on cooling solidify in the ordered crystalline state when passing the melting temperature  $(T_m)$ . However, the fluidity can be also preserved below melting, down to the glass temperature  $T_g \ll T_m$ , where the solidification from the metastable ultraviscous/ultraslowing liquid to the solid amorphous glass state occurs (Donth, 2000; Rzoska et al., 2010; Berthier and Ediger, 2016). There are also numerous semi-crystalline systems where the vitrification is related to the solidification of one or few elements of symmetry: as examples can serve orientationally disordered crystals (plastic crystals) (Drozd-Rzoska et al., 2006a,b) or liquid crystals (Drozd-Rzoska, 2006, 2009). For many systems, passing  $T_m$  without crystallization is associated with an extreme temperature quench (Donth, 2000). However, there are also numerous glass formers where entering the metastable ultraviscous/ultraslowing domain is possible at any practical experimental cooling rate (Donth, 2000; Rzoska et al., 2010; Berthier and Ediger, 2016). Turnbull (Turnbull, 1969; Angell, 2008) formulated the broadly used empirical Glass-Forming Ability (GFA) rule distinguishing poor  $(T_g/T_m < 2/3)$  and good glass formers  $(T_g/T_m > 2/3)$  and linking  $T_g$  and  $T_m$ . Notwithstanding, there is a notable difference between melting and vitrification: melting is related to the "sudden and almost non-signaled" fusion on cooling whereas the glass transition is hallmarked by far previtreous super-Arrhenius (SA) changes of viscosity  $\eta(T)$ , primary relaxation time  $\tau(T)$ , or other related dynamic properties (Avramov and Milchev, 1988; Donth, 2000; Rzoska et al., 2010; Berthier and Ediger, 2016). This opens the possibility of estimating the glass temperature from the analysis of previtreous effects well above  $T_g$ : as the general reference values  $\eta(T_g) = 10^{13}$  Poise for viscosity or  $\tau(T_g) = 100$  s for the primary (alpha, structural) relaxation time are assumed, since they correlate with the thermodynamic estimation (heat capacity or density scan) of Tg related to 10 K/min cooling rate (Donth, 2000; Rzoska et al., 2010). Although the ultimate form of  $\tau(T,P)$  or  $\eta(T, P)$  portrayal in previtreous ultraviscous/ultraslowing liquids on approaching  $T_g$  remains puzzling (Martinez-Garcia et al., 2013, 2014, 2015), most often the Vogel-Fulcher-Tammann (VFT) relation is used

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(Tammann, 1903; Vogel, 1921; Fulcher, 1925; Donth, 2000; Rzoska et al., 2010; Martinez-Garcia et al., 2013; Berthier and Ediger, 2016):

$$\tau(T) = \tau_0 \exp\left(\frac{D_T T_0}{T - T_0}\right), \qquad P = const \tag{1}$$

where  $\tau_0 = 10^{-14\pm 2}$  is the prefactor,  $T_0 < T_g$  is the VFT singular temperature, and  $D_T$  denotes the fragility strength coefficient linked to fragility metric  $m = \lfloor d \log_{10} \tau/d(T_g/T) \rfloor_{T \to T_g}$  via the empirical dependence  $D_T = 590/(m + \log_{10} \tau_0/\log_{10}(T_g))$  (Böhmer et al., 1993), in which  $\tau_0 = 10^{-14}$  s is assumed.

The pressure counterpart of the VFT equation was first proposed for the analysis of viscosity changes in glycerol by Johari and Whalley (1972) and later for the primary relaxation time in dibutyl phthalate (Paluch et al., 1996):

$$\eta = \eta_0^P \exp\left(\frac{A}{P_0 - P}\right)$$
 and  $\tau = \tau_p^P \exp\left(\frac{A}{P_0 - P}\right)$ , (2)

where: T = const,  $\eta_0^P$  and  $\tau_o^P$  denote prefactors, the amplitude A = const and  $P_0 > P_g$  is the "VFT-like" singular pressure.

However, Eq. 2 can reliably portray experimental data only for "strong" (weakly non-Arrhenius) glass formers, assuming that measurements terminates at  $P_{max} \ll P_0$ . In Paluch et al. (1998), the relation able to portray the previtreous dynamics for an arbitrary glass former and range of pressure was proposed:

$$\tau(P) = \tau_0^P \exp\left(\frac{A(P)}{P_0 - P}\right) = \tau_0^P \exp\left(\frac{D_P P}{P_0 - P}\right).$$
(3)

In this relation the amplitude is pressure dependent  $A = A(P) = D_P P$ , and the pressure fragility strength coefficient  $D_P$  was introduced. It is notable that for the basic VFT Eq. 1 the prefactor is "approximately universal," i.e.,  $\tau_0 \approx 10^{-14\pm 2}$  s, whereas for Eqs 2 and 3, it ranges between  $\tau_0^P \approx 10s$  and  $\tau_0^P \approx 10^{-14}s$  (Drozd-Rzoska and Rzoska, 2006; Drozd-Rzoska et al., 2008). Such enormous discrepancy results from the location of the  $\tau(P)$  isotherm selected for  $\tau(P)$  or  $\eta(P)$  tests in the respect to  $T_g(P)$  curve. This can be illustrated *via* the "general" SA equation:

$$\tau(T,P) = \tau_0^P \exp\left(\frac{PV_a(P)}{RT}\right) = \tau_0 \exp\left(\frac{E_a(T)}{RT}\right)$$
$$\exp\left(\frac{PV_a(P)}{RT}\right) = \tau_0 \exp\left(\frac{E_a(T) + PV_a(P)}{RT}\right). \quad (4)$$

The comparison of Eqs 3 and 4 yields  $E_a(T) = RD_T/(1T_0 - 1/T)$  and  $V_a(P) = TD_P R/(P_0 - P)$  for "VFT-type estimations" of the activation energy and activation volume, respectively. Notwithstanding, the general and model-free forms of  $E_a(T)$  and  $V_a(P)$  dependencies are still not known. The solution of the problem of the poorly defined prefactor  $\tau_0^P$  in Eqs 2 and 3 was proposed in Drozd-Rzoska and Rzoska (2006) and Drozd-Rzoska et al. (2008) by introducing the equation:

$$\tau(P) = \tau_0^P \exp\left(\frac{D'_P \left(P - P_{S_P}\right)}{P_0 - P}\right) = \tau_0 \exp\left(\frac{D'_P \Delta P}{P_0 - P}\right).$$
(5)

This dependence takes into account that the liquid state terminates at the absolute stability limit pressure (spinodal  $P_{Sp}$ ), in negative pressures domain. The ultimate description needs both positive (isotropic compression, hydrostatic pressures, P > 0) and negative pressures (isotropic stretching, P < 0) domains (Angell and Quing, 1989, Imre et al., 2002). For Eq. 5, the prefactor is approximately the same,  $\tau_0^P = \tau (P_{Sp}) \approx 10^{-12}$ s, for any isotherm. When comparing Eqs 3 and 5 worth noting is that the latter can penetrate negative pressures domain but the fragility strength coefficients changes:  $D_P/D'_P = P_0/(P_0 - P_{Sp})$  (Drozd-Rzoska and Rzoska, 2006; Drozd-Rzoska et al., 2008).

The characterization of  $T_g(P)$  dependence has a notable impact on the behavior under atmospheric pressure, being included *via* the coefficient  $dT_g(P)/dP$  in numerous relations (Drozd-Rzoska et al., 2007a,b; Rzoska et al., 2010; Donth, 2000; Rzoska and Mazur, 2007; Floudas et al., 2011). The reliable knowledge of  $T_g(P)$  description seems to be essential for silicate glasses, in which practically important features are created due to the high pressure—high temperature annealing with induced "exotic" features preserved after decompressing. They are, for instance: (i) the notable increase of density, (ii) the increase of hardness, and (iii) the anty-cracking ability (Smedskjaer et al., 2014; Januchta et al., 2016, 2017; Svenson et al., 2017). Still puzzling is the description of  $T_g(P)$  behavior in systems where  $dT_g/dP < 0$  (Donth, 2000; Drozd-Rzoska et al., 2007a,b, 2008).

All above show that the reliable and effective portrayal of the pressure evolution of the glass temperature can constitute one of milestones in dealing with the glass transition. This report presents the resume of this issue, supplemented by some extensions beyond the current state-of-the art.

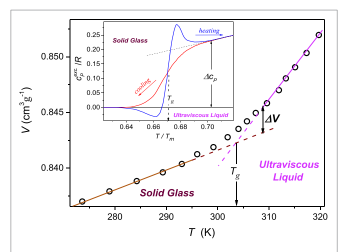
## PARAMETERIZATION OF THE PRESSURE EVOLUTION OF MELTING AND GLASS TEMPERATURES

There are several relations for describing the pressure evolution of melting temperature: the most popular is the Simon–Glatzel (SG) equation due to its simple form and the limited number of fitted parameters (Simon and Glatzel, 1929; Skripov and Faizulin, 2006):

$$T_m(P) = T_0 \left(1 + \frac{P}{a}\right)^{1/b},$$
 (6)

where  $T_0$ , a, and b are adjustable parameters.

It can be derived from the Clausius–Clapeyron (C–C) equation  $dT/dP = T\Delta V/\Delta H = \Delta V/\Delta S$ , where  $\Delta V$ ,  $\Delta H$ , and  $\Delta S$  are for the volume, enthalpy, and entropy changes at the transition, assuming  $(dT/dP)_{fusion} = a + bP$  (Skripov and Faizulin, 2006). This relation is used for the description of melting, where the "sudden and sharp" change of volume or density ( $\Delta V$ ,  $\Delta \rho$ ) and entropy takes place. However, the C–C equation can be linked to any fusion phenomenon, provided it is associated with detectable changes in entropy and volume/density. This occurs also for the glass transition temperature, although the transformation is "stretched" in temperature or pressure and occurs between the disordered (ultravisous) liquid and the disordered solid (glass), as shown in **Figure 1**.



**FIGURE 1** | The temperature dependence (P = 0.1 MPa) of the proper volume  $V = 1/\rho$ ,  $\rho$  denotes density, for polyvinyl acetate in the ultraviscous and solid amorphous phases. Dashed lines show extrapolations of the experimental behavior remote from the "stretched" glass transition domain  $T_g$ . The apparent discontinuity of the volume can be estimated as  $\Delta V' = 0.0021 \, cm^3 g^{-1}$  and  $\Delta V'' = 0.0030 \, cm^3 g^{-1}$  (double arrows in the plot). The inset, based on data from McKinney and Goldstein (1974), Roland and Casalini (2003), and Tropin et al. (2012) is for the excess of the specific heat  $\Delta c_{\rho}(T) = c_{\rho}^{solid}(T) = a + bT$  described the behavior well below. The resulting discontinuity  $\Delta c_{\rho}(T)/R = 0.23$ . Data in this figure are for 10 K/min. cooling/heating rate.

As mentioned above the "reasonable" metric of the glass transition is the isochronal or isoviscous condition  $\tau(T_g, P_g) = 100 \text{ s}$ or  $\eta(T_g, P_g) = 10^{13}$  Poise (Donth, 2000). Generally, such condition is absent along the melting curve within the P-T plane (Skripov and Faizulin, 2006). However, the isochronal condition for  $T_m(P)$  is clearly fulfilled if melting is associated with only one element of symmetry, as for the isotropic-nematic transition in liquid crystals (Roland et al., 2008). Heuristic similarities between melting and vitrification can be strengthen recalling the empirical link between  $T_g$  and  $T_m$ , used as the indicator of the GFA:  $T_g/T_m > 2/3$  (near-spherical molecules) and  $T_g/T_m > 1/2$  (elongated molecules) (Turnbull, 1969; Donth, 2000; Angell, 2008). Consequently, one can expect that the pressure dependence of  $T_m$ can be paralleled by  $T_g(P)$  evolution. Regarding the vitrification, S. Peter Andersson and Ove Andersson (AA) introduced the SGtype relation for describing the pressure evolution of the glass temperature in poly(propylene) glycol (Andersson and Andersson, 1998):

$$T_g(P) = k_1 \left( 1 + \frac{k_2}{k_3} P \right)^{1/k_2}$$
(7)

where  $k_1$ ,  $k_2$ , and  $k_3$  are empirical, adjustable parameters.

The AA equation has become the key tool for describing  $T_g(P)$  experimental data till nowadays (Roland et al., 2005; Drozd-Rzoska et al., 2007a; Rzoska and Mazur, 2007; Rzoska et al., 2010; Floudas et al., 2011). This success was notably strengthen by its derivation within the Avramov–Milchev (AM) phenomenological model for the vitrification (Avramov and Milchev, 1988; Roland and Casalini, 2003; Hu et al., 2017):

$$T_g(P) = \varepsilon T_0 \left( 1 + \frac{P}{\Pi} \right)^{\beta/\alpha}, \tag{8}$$

TABLE 1   Systems in which	the application	of pressure	decreases	the	glass
temperature ( $dT_g/dP < 0$ ).					

Glass former $dT_r/dP$ Reference				
dT <sub>g</sub> /dP (K/GPa)	Reference			
-8.5	Kanno et al. (1981)			
-5	Williams and Angell (1977)			
-52	Giovambattista et al. (2012)			
-8.4	Bagdassarov et al. (2004)			
-45	Bagdassarov et al. (2004)			
-57	Deb et al. (2001)			
-30	Ramesh (2014)			
-78	Ramesh et al. (2016)			
-41.5	Trybuła and Stankowski (1998)			
	-8.5 -5 -52 -8.4 -45 -57 -30 -78			

For the dominant group of glass formers (molecular liquids, polymers, etc.):  $dT_g/dP > 0$  (Donth, 2000; Roland et al., 2005; Floudas et al., 2011).

where the coefficient  $\varepsilon = [30\log_{10}(e)/(\log_{10}(\tau(T_g)) - \log_{10}\tau_0)]^{1/\alpha}$ .

Notable is some discrepancy between Eqs 7 and 8 because the coefficient  $\varepsilon \gg 1$ . Worth recalling is also the criticism (Martinez-Garcia et al., 2013, 2014) regarding the basic AM model (Avramov and Milchev, 1988) output relation  $\tau(T) = \tau_0 \exp(A/T^D)$  or  $\eta(T) = \eta_0 \exp(A/T^D)$ , for P = const.

It is worth stressing that for SG Eq. 6 and AA Eqs 7 and 8 always  $dT_{g,m}/dP > 0$ , i.e.,  $T_m(P)$  and  $T_g(P)$  have to increase permanently with rising pressure. However, there are also systems  $dT_{g,m}/dP < 0$ , although the experimental evidence for such glass formers is still limited: some of them are collected in **Table 1**.

It seems that such behavior may occur only for some strongly bonded glass formers. Notwithstanding, taking into account the clear evidence of systems with  $T_m(P)$  maximum (Kechin, 1995, 2001; Tonkov and Ponyatovsky, 2004), the similar behavior can be expected for  $T_g(P)$  curves. It is notable, that already a century ago it was indicated that the reversal melting  $dT_m/dP > 0 \rightarrow (T_m^{max}, P_m^{max}) \rightarrow dT_m/dP < 0$  can be the general phenomenon (Tammann, 1903), although it can be hidden in the negative pressures domain or its emergence can be stopped by a phase transition. The description of the reversal melting phenomenon was first clearly proposed by Rein and Demus (RD) (Demus and Pelzl, 1988; Rein and Demus, 1992) and subsequently by Kechin (K) (Kechin, 1995, 2001):

$$T_m(P) = T_0 \left(1 + \frac{P}{a}\right)^{1/b} \exp(-a_1 P) = R(P) \times D(P),$$
 (9)

where a, b and  $a_1$  are adjustable parameters. R(P) denotes the SG-type "rising" term and D(P) is for the "damping term."

In subsequent decades Eq. 9, most often recalled as the "Kechin equation," became the key tool for describing experimental data associated with melting curve maximum (Drozd-Rzoska, 2005; Skripov and Faizulin, 2006; Drozd-Rzoska et al., 2007a; Rzoska and Mazur, 2007; Rzoska et al., 2010). Regarding the meaning of parameters in Eqs 6–9 one can generalize the reasoning of Burakovsky et al. (Burakowsky et al., 2000; Burakovsky et al., 2003), who considered the volume-related compression factor (modulus):  $\eta' = \Delta V_0 / \Delta V = (V(\pi) - V(P_0))/(V(P) - V(P_0))$ 

and linked it to the bulk (compressibility) modulus *via*  $B = -\Delta V(d(\Delta P)/d(\Delta V)) = \eta' d(\Delta P)/d\eta'$ , with the pressure dependence given as  $B(P) = B_0 + B'_0P + \dots$  and  $\Delta P = P - P_0$ :

$$\eta'(P) = \left(1 + \frac{B'_0}{B_0}P\right)^{1/B'_0} \to T_m(P) = T_0 * (\eta')^{-1}$$
$$= T_0 \left(1 + \frac{P}{B_0./B'_0}\right)^{1/B'_0}, \tag{10}$$

where the index " $_0$ " is related to the reference point ( $T_0$ ,  $P_0$ ).

Hence, taking the atmospheric pressure as the reference one can indicate the following meaning of parameters in Eqs 6-9  $a = B_0/B'_o = \pi$  and for the power exponent  $b = B'_0$ . For SG and AA Eqs 6 and 7, as well as K&RD Eq. 8, the reference has to be taken as  $T_0 = T_{g,m}(P_0 = 0) \approx T_{g,m}(P_0 = 0.1MPa)$ . Other selections of  $T_0$  yields non-optimal and effective values of fitted coefficients. In Skripov and Faizulin (2006) as the general reference the triple point was proposed: and the  $T_0 = T_{triple}$ and  $P \rightarrow \Delta P = P - P_{triple}$  in the SG Eq. 6. Such reference cannot be implemented for the glass transition. Drozd-Rzoska (Drozd-Rzoska, 2005; Drozd-Rzoska et al., 2007a, 2008) proposed as the reference arbitrary values  $(T_0, P_0)$  along melting or vitrification curves, assuming  $\Delta P = P - P_0$ . Subsequently, considering the Clausius-Clapeyron equation along the melting or vitrification curve  $(\Delta H/\Delta V)_{T_{g,m},P_{g,m}} = (b\Pi + b\Delta P)/(1 - c(b\Pi + b\Delta P))$  the following relation was derived (Drozd-Rzoska, 2005):

$$T_{g,m}(P) = T_0 \left( 1 + \frac{P - P_{Sp}}{\pi + P_0} \right) \times \exp\left(-\frac{P - P_0}{c}\right)$$
$$= T_0 \left( 1 + \frac{\Delta P}{\Pi} \right)^{1/b} \times \exp\left(-\frac{\Delta P}{c}\right), \quad (11)$$

where  $\Delta P = P - P_0$ ,  $-\pi$  is the extrapolated negative pressure value for which  $T_{g,m}(P \rightarrow -\pi) \rightarrow 0$ : it correlates with the onset of  $T_{Sp}(P_{Sp})$  absolute stability limit curve in negative pressures domain; *c* is the damping pressure coefficient.

For small or moderate pressures one obtains the SG or AA-type equation(Drozd-Rzoska, 2005; Drozd-Rzoska et al., 2007a, 2008):

$$T_{g,m}(P) \approx T_0 \left( 1 + \frac{P - P_0}{\pi + P_0} \right) = T_0 \left( 1 + \frac{\Delta P}{\Pi} \right)^{1/b}$$
(12)

Equation 11 is able to portray systems with the maximum of melting or vitrification curve, even if they are hidden in the negative pressures domain. It can be also applied for systems were  $dT_{g,m}/dP < 0$ . Equation 12 can describe experimental data if  $dT_{g,m}(P)/dP > 0$  and the set of data is well below the maximum of  $T_{g,m}(P)$  curve. Both relations can be implemented in the negative pressures domain. Applying results of Burakowsky et al. (2000), one obtains:  $b = B'_0$  and  $B_0/B'_0 = P_0 + \pi$  and then  $B_0 = B'_0P_0 + B'_0\pi$ . The latter equation is in agreement with the empirical relation for the pressure evolution of the bulk modulus recalled above (Murnaghan, 1944). It is notable that both the basic AA Eq. 7 and the extended Eq. 12 are able to portray experimental data in the negative pressures domain. For Eq. 7 such portrayal was successfully applied in Adrjanowicz et al. (2015). However, for Eq. 7, one has to assume  $T_0(P=0)$  as the reference. Moreover, the direct substitution of negative pressures, is not possible if hallmarks of the reversal vitrification ( $T_g(P)$  maximum) appears: Demus–Rein–Kechin (Eq. 9) is not able to portray experimental data if substituting P < 0, contrary to Drozd–Rzoska et al. (Eq.11).

There are few other approaches considering  $T_m(P)$  evolution which start from the C–C or related Lindemann relations (Skripov and Faizulin, 2006). They are briefly presented below, with indications of their applicability for the glass formation. Schlosser et al. (1989) starting from the Lindemann relation  $T_m = CV^{2/3}\Theta_D$ (*C* is a constant,  $\Theta_D$  is the Debye reduced temperature) (Lindemann, 1910; Skripov and Faizulin, 2006) and the definition of the Grüneisen parameter as  $\gamma = (\partial \Theta_D / \partial V)_T = - \partial \ln \Theta_D / \partial \ln V$ (Grüneisen, 1912) obtained the relation focusing on the volume dependence of the melting temperature. Generalizing this dependence for the arbitrary fusion process one obtains:

$$T_{g,m}(V) = T_0 \left(\frac{V}{V_0}\right)^{2/3} \exp\left(2\gamma_0 \frac{V - V_0}{V_0}\right) = T_0 X^2 \exp\left(2\gamma_0 \frac{\Delta V}{V_0}\right).$$
(13)

where the index " $_0$ " is for the zero-pressure (~atmospheric pressure) reference.

Assuming for the  $X^2 \approx 1 - 2\Delta V/3V_0 \approx \exp(-2\Delta V/3V_0)$  following relation was derived (originally for melting):

$$T_{g,m}(V) = T_0 \exp\left(\frac{-2\Delta V}{3V_0}\right) \exp\left(2\gamma_0 \frac{\Delta V}{V_0}\right)$$
(14)

One may expect that it is able to portray systems described both by  $dT_{g,m}/dP > 0$  and  $dT_{g,m}/dP < 0$ . For small/moderate pressures Eq. 14 can be reduced to the Kraut–Kennedy relation (Kraut and Kennedy, 1966; Schlosser et al., 1989), originally developed for melting:

$$T_{g,m} \approx T_0 \left[ 1 + 2 \left( B_0 - 1/3 \right) \frac{\Delta V}{V_0} \right] = T_0 \left( 1 + C \Delta V / V_0 \right).$$
 (15)

It can be converted to the density related dependence along melting or vitrification curves:

$$T_{g,m} \approx T_0 \left( 1 + C \frac{\rho_0 - \rho}{\rho} \right) = T_0 \left( 1 + C \frac{\Delta \rho}{\rho} \right).$$
 (16)

Linking Eqs 12 and 15 one obtains the relation for pressureinduced volume changes along melting or vitrification curve:

$$\left(\frac{\Delta V}{V_0}\right)_{g,m} = \frac{(1 + \Delta P/\Pi)^{1/b} - 1}{C}.$$
 (17)

This relation is in fair agreement with the Murnaghan equation, broadly used is earth sciences (Murnaghan, 1944; Poirier, 2000; Skripov and Faizulin, 2006). Recalling the dependence  $\Delta V/V_0 = \ln(1 + \beta P)/\alpha$ , where  $\alpha = B' + 1$  and  $\beta = \alpha/B = (B'_0 + 1)/B$  Eq. 15 can be converted to the SG- or AA-type equation (Schlosser et al., 1989):

$$T_{g,m}(P) \approx T_0 (1 + \beta P)^{2(B-1/3)/\alpha}$$
 (18)

It this relation the SG exponent  $b = (B'_0 + 1)/(2(B_0 - 1/3))$ , i.e., it differs from Burakovsky (Burakovsky et al., 2003) predictions.

Kumari and Dass (Kumari and Dass, 1988; Dass, 1995) also applied the framework of the Lindemann criterion (Lindemann, 1910) and workout the relation originally focused on the pressure evolution of the melting temperature, focusing on alkali metals:

$$\ln\left(\frac{T_{m,g}}{T_0}\right) = -2\alpha P + \left[2\left(C + \frac{\alpha}{\beta}\right)\ln\left(1 + \beta P\right)\right], \quad (19)$$

where  $\alpha = (\gamma'/B')P_0$ ,  $T_0$ ,  $\beta = (B'/B)_{P_0,T_0}$ ,  $C = [(\gamma - 1/3)/B']_{P_0,T_0}$ ,  $\gamma, \gamma, '$  and *B*, *B*' stands for the Grüneisen parameter, bulk modulus and their first derivatives.

This relation can describe systems notably diverging from the SG pattern, including the crossover  $dT_{g,m}/dP > 0 \rightarrow dT_{g,m}/dP < 0$ . It can be also converted to the form coincided with Rein and Demus and Kechin Eq. 8:

$$T_{m,g} = T_0 (1 + \beta P)^{2C + 2\alpha/\beta} \exp\left(-2\alpha P\right)$$
(20)

The coefficient  $\alpha = \gamma'/B'$ , what makes it possible to define the "damping pressure" parameter in DR Eq. 11:  $c = B'/2\gamma'$ . Equation 20 can be reduced to the SG or AA forms assuming  $\alpha = 0$  (Dass, 1995), i.e.,  $\gamma(P) = const$  in the given range of pressures:

$$T_{m,g}(P) = T_0 (1 + \beta P)^{2C}.$$
 (21)

It is also notable that Eq. 19 makes it possible to estimate the location of the maximum of  $T_{g,m}(P)$  curves as  $P_{g,m}^{\max} = (\gamma - 1/3)/\gamma'$ . Taking into account the form of the exponent *C* worth recalling is Lindemann–Gilvary law (Gilvarry, 1966)  $dT_m/dP = T_m[2(\gamma - 1/2)/B]$ , what indicates the pressure dependence of the power exponent in the SG-type Eq. 21. Schlosser et al. Equation 13 and Kumari–Dass Eq. 19 and can be extended to the negative pressures domain when introducing the reference related to the absolute stability limit in the negative pressures domain:  $P \rightarrow \Delta P = P - P_{Sp}, V \rightarrow \Delta V = V - V_{Sp}, \rho \rightarrow \rho - \rho_{Sp}$ .

The formal base of above relations, including the Andersson–Andersson equation, are the extended C–C relation or/and the Grüneissen parameter definition. Their implementations are related to different pressure dependences of the volume and the modules in neighboring phases. The latter give rise to the nonlinear dependence of the enthalpy. For melting at the welldefined temperature such behavior is easily detectable in neighboring phases. For the glass transition there are "stretched" gradual changes of mentioned properties over the transition region between coexisting ultraviscous/ultraviscous and solid states. Notwithstanding, also for the glass transition one can define the equivalents of "jumps" for  $\Delta V$  and  $\Delta S$ , or equivalently  $\Delta H$ (Figure 1).

### THE ANALYSIS OF EXPERIMENTAL DATA

When considering the parameterization of  $T_g(P)$  or  $T_m(P)$  experimental data, some basic problems emerges:

(i) Does the selected equation is proper for portraying the given set of data?

- (ii) What is the pressure range of applicability of the description?
- (iii) Is it possible to estimate optimal values of parameters, avoiding the uncertainty associated with the number of parameter and the nonlinear fitting?

To address these questions, in Drozd-Rzoska (2005), Drozd-Rzoska and Rzoska (2006), and Drozd-Rzoska et al. (2007a), the preliminary derivative-based and distortions-sensitive analysis of  $T_m(P)$  and  $T_g(P)$  experimental data was proposed:  $T_g(P) \Rightarrow [d(\ln T_{g,m})/dP]^{-1}$ . For SG/AA or DR Eqs 6, 7, and 12, one obtains the linear behavior of transformed experimental data (Drozd-Rzoska, 2005; Drozd-Rzoska et al., 2007a):

$$(d \ln T_{g,m}/dP)^{-1} = ba + bP$$
 and  
 $(d \ln T_{g,m}/dP)^{-1} = b\pi + bP.$  (22)

It is visible that the description via DR and SG/AA relations overlaps and both can be extended into the negative pressures domain. However, such possibility for the AA and SG relation may be casual since it does not takes place for Rein and Demus and Kechin Eq. 9, for Kumari and Dass Eq. 19 or for pressure counterparts of the VFT relation (Eqs 2 and 3).

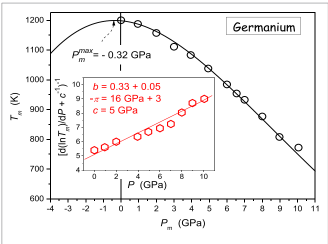
Regarding the "general" DR Eq. 11, the following transformation of experimental data was proposed to test the domain of its validity (Drozd-Rzoska et al., 2007a, 2008):

$$\left[d(\ln T_m)/dP + c^{-1}\right]^{-1} = A + BP$$
(23)

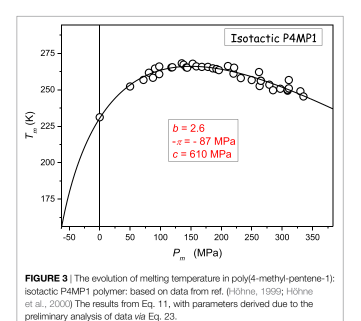
For the optimal selection of the damping pressure coefficient c one obtains the linear behavior of transformed experimental data and the linear regression fit yields optimal values of  $\pi$ , b, and c coefficients. Subsequently, they can be substituted to Eq. 11, avoiding the nonlinear fitting.

Concluding, Eqs 22 and 23 define the way of the preliminary transformation and analysis of experimental  $T_{g,m}(P)$  via the plot  $d \ln T_{g,m}/dP$  vs. P, which indicates the domain of the domain of validity of the given description and optimal values of parameters. The derivative-based and distortions-sensitive preliminary analysis can reveal even "weakly emergent" hallmarks of approaching  $dT_{g,m}/dP > 0 \Leftrightarrow dT_{g,m}/dP < 0$  crossover, hardly "eye-detectable." Below, practical applications of above reasoning are discussed. First, they are focused on melting of germanium  $(dT_m/dP < 0)$ (Vaidya et al., 1969; Porowski et al., 2015) and subsequently for the "soft" material, P4MP1 polymer, with  $T_m(P)$  maximum (Höhne, 1999; Höhne et al., 2000). It is worth stressing that for the vast majority of systems tested so far  $dT_m/dP > 0$  (Kechin, 1995, 2001; Skripov and Faizulin, 2006) and there is much lesser number of systems where  $dT_m/dP < 0$  (see Table 1). Figure 2 presents such data for germanium, which can be well portrayed by DR Eq. 11, with parameters obtained from the pre-analysis of experimental data via Eq. 23, as shown in the inset. Notable, is the possible maximum of  $T_m(P)$  curve hidden in the negative pressures domain at  $P_{\rm max} \approx -0.32 \, {\rm GPa}.$ 

**Figure 3** presents the unique "soft matter system" where the crossover  $dT_m/dP > 0 \Leftrightarrow dT_m/dP < 0$  takes place at relatively low pressures:  $P_{\text{max}} \approx 150 MPa$ . Recalling the Kumari–Dass model (Dass, 1995; Kumari and Dass, 1988) such small value of  $P_{\text{max}}$ 



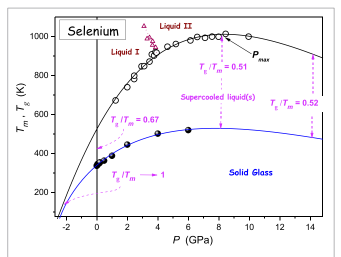
**FIGURE 2** | Pressure dependence of melting temperature of germanium [based on data from ref. Vaidya et al. (1969), Porowski et al. (2015)]. Experimental data are portrayed by DR Eq. 11, with the support of the preliminary derivative-based analysis (Eq. 23) yielding also optimal values of parameters: this is shown in the inset.



may result from the strong pressure dependence of the Grüneissen parameter.

One can expect that different types of  $T_m(P)$  dependences should be paralleled by  $T_g(P)$  behavior, taking into account the form of GFA factor. Unfortunately, the number of experimental data for  $T_g(P)$  is very limited.

**Figure 4** shows the compilation of  $T_g(P)$  and  $T_m(P)$  experimental data available for selenium. It is notable that a single DR Eq. 11 curve can describe the whole set of  $T_m(P)$  data, without a hallmark of passing a liquid I–liquid II (L–L) transition (Imre and Rzoska, 2010). This issue is worth stressing because often  $dT_m/dP$  discontinuity is reported when passing the L–L transition (Imre and Rzoska, 2010). The value of the ratio  $T_g/T_m$  changes from  $T_g/T_m(P \approx 0.1MPa) \approx 2/3 \rightarrow T_g/T_m(P \approx P_{max}) \approx$ 



**FIGURE 4** | The pressure evolution of melting and glass temperature for selenium. The change of  $T_g/T_m$  value is indicated. Solid curves are described by DR Eq. 11: parameters were derived from the preliminary analysis based on Eq. 23. Experimental data were taken from refs. (Deaton and Blum, 1965; Tanaka, 1984; Ford et al., 1988; Katayama et al., 2000; Caprion and Schober, 2002).

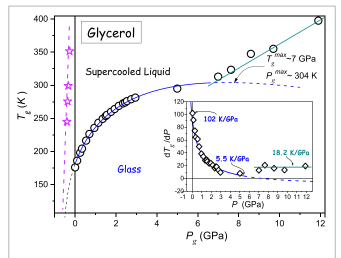
1/2 (Drozd-Rzoska et al., 2007a, 2008). When entering the negative pressures domain the GFA factor  $T_g/T_m \rightarrow 1$ , i.e., the system becomes extremely good glass former.

Glycerol belongs to the group of the most "classical" glassforming ultraviscous liquids (Donth, 2000; Rzoska and Mazur, 2007; Rzoska et al., 2010; Berthier and Ediger, 2016) Figure 5 shows the compilation of data from the authors' broad band dielectric spectroscopy pressure studies and the analysis of the primary relaxation time  $\tau(T, P)$  via Eq. 5 supplemented by earlier  $T_{g}(P)$  estimations (Drozd-Rzoska, 2005; Drozd-Rzoska et al., 2007a). Notable is the emergence of two types of  $T_g(P)$  evolution. The first one leads to the maximum of  $T_g(P)$  curve at  $P_g^{\max} \approx 7GPa$ and it is followed by a hypothetical reversal vitrification associated with  $dT_{q}(dP < 0)$ . However, prior to reaching the maximum, at  $P \approx 6.5 \, GPa$  the "crossover" to the another form of  $T_g(P)$  evolution, described by  $dT_g/dP > 0$  takes place. The dashed curve shows the extrapolation of the solid blue curve, with the indication of a hypothetical "hidden" maximum of  $T_g(P)$  curve. The inset in **Figure 2** shows changes of  $(dT_g/dP)$  coefficient on rising pressure, additionally distinguishing two different types of  $T_g(P)$  evolution.

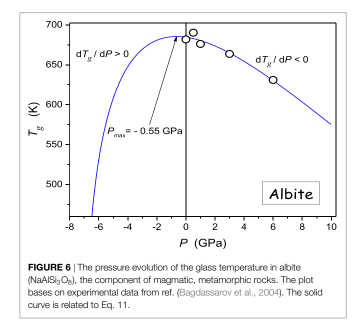
Generally, the experimental evidence of glass formers characterized by  $dT_g/dP < 0$  is very limited (see **Table 1**). Such behavior seems to be characteristic for some strongly bonded systems. **Figure 6** shows results of such studies for albite, a geophysically important material, which can be well portrayed by Eq. 11, revealing the maximum of  $T_g(P)$  curve "hidden" in negative pressures domain.

## UNIVERSAL ASPECTS OF THE PRESSURE EVOLUTION OF THE GLASS TEMPERATURE

The above discussion indicated the possible common phenomenological description of  $T_g(P)$  evolution in glass formers



**FIGURE 5** | The pressure evolution of the glass temperature for glycerol. The solid blue curve, with "dotted" and "dashed" parts is related to DR Eq. 11 and the preliminary analysis *via* Eq. 23. Experimental data are from author's measurements and from Cook et al. (1994), Drozd-Rzoska (2005), Drozd-Rzoska et al. (2007a), Pronin et al. (2010). The dashed line and stars (in magenta) in the negative pressures domain denotes the possible absolute stability limit location: this was determined from the analysis of  $\tau(P)$  experimental data *via* Eq. 5. The inset shows the pressure evolution of  $dT_g/dP$  coefficient.



described by  $dT_g(P)/dP > 0$  and/or  $dT_g(P)/dP < 0$ . The question arises of the more microscopic insight. Voigtmann (2006a) analyzed the vitrification within frames of the square-well (SW) model associated with the relatively simple potential:  $U(r) = \infty$ for distances r < d supplemented with an SW attraction within the range  $\delta$ ,  $U(r) = -U_0$  for  $d < r < d(1 + \delta)$ , and U(r) = 0 beyond was analyzed. The SW approach proved its superior ability for describing colloidal glass formers, which can be thus considered as a kind of archetypical experimental glass-forming model system.

In Voigtmann (2006a), the possibility of the common description of glass-forming molecular liquids and colloids was shown, using the plot  $\log_{10}P_g^*$  and  $\log_{10}T_g^*$ , where the "natural units," i.e., model normalized glass pressure and temperature were used:  $T_g^* = T_g/T_g^{\text{mod }el}$  and  $P^* = P_g/P_g^{\text{mod }el}$ . In Voigtmann (2006b), the similar plot was tested for the model fluid associated with the Lennard–Jones (LJ)  $V_{LJ} = 4 \in [(r/\sigma)^{-12} - (r/\sigma)^{-6}]$  potential analyzed within the mode-coupling theory approximation. In Voigtmann (2006a),  $T_g(P)$  experimental data for glycerol, dibutyl phthalate, o-terphenyl, and epoxy resin EPON 828 were analyzed  $(dT_q/dP > 0)$ . In Roland and Casalini (2003), only glycerol was discussed, for the clarity of reasoning. This report also focuses on glycerol, but for the notably enhanced range of pressures, basing on data from Figure 5. This is supplemented by experimental data for albite, where  $dT_g/dP < 0$  (Figure 6). In Voigtmann (2006a), the SW model units were used for scaling, namely  $T_g^{\text{mod }el} = T_g^{SW} = U_0/k_B = 826K$  and  $P_g^{\text{mod }el} = P_g^{SW} = U_0/d^3 = 3.09GPa$  and in Voigtmann (2006b), the LJ model units, i.e.,  $T_g^{LJ} = k_B/\in$ 500K and  $P_g^{LJ} = \epsilon/\sigma^3 = 2.5GPa$ : numbers are given for glycerol. In Voigtmann (2006b), the partial agreement between predictions of SW and LJ model was obtained after ad hoc shifting  $T^* \rightarrow 1.5 T^*$ . It is notable that so far experiments in colloids are carried out under atmospheric pressure and obtained phase diagrams are presented using the volume fraction  $(\phi)$ —interaction strength or temperature axes. Such data were model-mapped into the pressure-temperature plane in Voigtmann (2006a). Figure 7 recalls results of Voigtmann (2006a) for: (i) the colloid with the addition of polymer increasing attraction and causing the "reentrant" vitrification (Pham et al., 2002), (ii) glycerol  $(dT_g/dP > 0)$ for experimental data taken from Figure 5, (iii) albite for which  $dT_g/dP < 0$  (Figure 6), and (iv) the SW model predictions for  $\delta = 0.04$  and  $\delta = 0.12$  values of the key parameter, (v) the model using LJ potential with and without the attraction. This is supplemented by results of fitting via DR Eq. 11 for glycerol and albite. One of key findings of Voigtmann (2006a,b) was the "generic steep" anomaly with exactly defined singularity, the same for any molecular glass former:  $T_g^* \rightarrow 0.23$  for SW model units and  $T_g^*$  (anomaly)  $\rightarrow 0.334$  for the LJ model. These led to the conclusion that there are three general regimes of glass formation resulted from  $T_g(P)$  evolution (Voigtmann, 2006a,b):

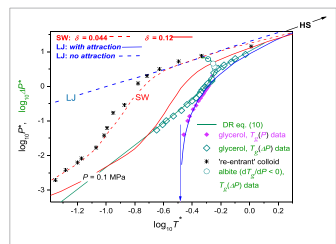
*Regime I*—for  $T_g^* > 1$ : glass formers approach the hard sphere limit. Following Voigtmann (2006a,b) in this domain:  $T_g \propto P_g^{4/5}$ .

*Regime II*—for  $1 > T_g^* > 0.23$  (*or*\_0.334): there is a universal "generic steep" anomaly and this regime is characteristic for molecular glass formers.

*Regime III*—for  $T_g^* \to 0$  the low density and weak interactions domain occurs. It is available for colloidal glass formers and does not accessible for molecular ones.

In Voigtmann (2006a,b), glass-forming systems for which  $dT_g/dP < 0$  were not discussed.

One of the most striking features of Voigtmann (2006a,b) is the "generic steep" anomaly, presumably occurring only for molecular glass formers. However, this unique phenomenon has few surprising features. First, it is very strong and associated with exactly



**FIGURE 7** | The pressure dependence of the glass temperature, summarizing the model discussion (Voigtmann, 2006a,b): SW is for the square-well potential model, LJ—the Lennard–Jones potential model and HS is for the hard spheres model. For details see the text of the given paragraph and refs. (Voigtmann, 2006a). Experimental data for glycerol are taken from **Figure 5**: they are present in the "natural scaled" units. Data for albite are from **Figure 6**. Note that for open green diamonds (glycerol) and open circles (albite) the reference pressure was takes into account:  $P \rightarrow \Delta P = P + \pi$ . Data for the polymer mediated colloid are from Pham et al. (2002) and Voigtmann (2006a). For details see comments in the given paragraph. Note the disappearance of the "generic steep" anomaly (indicated by the vertical arrow) and the ability for describing arbitrary glass former. For scaling model values see the text below.

the same "singular" value of  $T_g^* \approx 0.23$  for arbitrary molecular glass former. Well above the singularity experimental data for all molecular glass formers overlaps. Second, the "generic" anomaly appears in the log–log scale but no hallmarks of such behavior appears in the linear scale for any "native" Tg(P) data (Johari and Whalley, 1972; Andersson and Andersson, 1998; Donth, 2000; Roland et al., 2005; Drozd-Rzoska et al., 2007a, 2008; Rzoska and Mazur, 2007; Floudas et al., 2011). Third, although real high pressure results for colloidal glass formers are still not available, one can easily show that such data also will follow the same "generic steep anomaly" pattern, in disagreement with "re-calculated" data shown in **Figure 7** (stars).

Following all these, one can conclude that the "generic steep" anomaly is the consequence of  $P \rightarrow 0$  (i.e., in practice  $P \rightarrow 0.1 MPa$ ) within the plot applying the log-log scale. This is not a real physical phenomenon. Any fluid can be smoothly crossovered from the hydrostatic pressures region (P > 0) to the isotropically stretched, negative pressures domain (P > 0) (Imre et al., 2002). Experimental evidences clearly show the lack of any hallmarks of passing P = 0, also for supercooled molecular glass formers (Angell and Quing, 1989; Sciortono et al., 1995; Imre et al., 2002). The natural termination of the liquid state is the absolute stability limit spinodal in negative pressures domain, where any liquid "breaks" and the homogeneous cavitation occurs. Taking this as the reference one should consider the "universal plot" based on the scale  $\log_{10}\Delta P^* = \log_{10} \left[ (P + \pi)/P_g^{mod el} \right]$  vs.  $\log_{10}T_g^*$  instead of  $\log_{10}P^*$  vs.  $\log_{10}T_g^*$  plot.

Following refs. (Voigtmann, 2006a,b) the model parameters are related to the LJ potential, which is considered as a realistic interaction model in liquids:  $V_{LJ}(r) = 4U_0[(r/\sigma)^{-12} - (r/\sigma)^{-6}]$ ,

for which the model temperature and pressure  $T^* = k_B T/U_0$ and  $P^* = P\sigma^3/U_0$ . To correlate experimental and model data the "arbitrary" scale shift is also used (see for comparison: Voigtmann, 2006a,b). Following scaling values were assumed:  $P_g^{\text{mod } el} =$ 3.09GPa and  $T_g^{\text{mod } el} = 826K$  for glycerol and  $P_g^{\text{mod } el} = 10.4GPa$ and  $T_g^{\text{mod } el} = 210K$  for albite.

Consequently, the "generic steep" anomaly disappears and  $T_g(P)$  experimental data for molecular glass formers can be mapped also to the low density  $(T^* \rightarrow 0)$  domain. When linking such analysis with Eq. 11 one also obtains the possibility of describing systems characterized by  $dT_g/dP < 0$ , as shown for the extrapolated behavior for glycerol and for albite in **Figure 7**. **Figure 7** also shows that the re-entrant glass-forming colloids mapped from experimental studies under atmospheric pressure to the *P*-*T* plane are related to the case  $dT_g/dP < 0$ .

For glycerol, for very high pressures, the behavior described by  $T_g \propto P_g^{4/5}$  emerges and the evolution approaches the hard sphere limit pattern (Voigtmann, 2006a). One of arguments for the generic importance of the "steepness" anomaly in Voigtmann (2006a,b) was the possibility of it reproduction by the model-fluid with LJ potential containing properly adjusted attraction term. However, for the analysis of  $T_g^*(P_g^*)$  in such model-fluid the absolute stability limit have to be taken into account: after the transformation  $P \rightarrow \Delta P$  the "generic steep anomaly" disappears also for the LJ model fluid.

Concluding, the plot  $\log_{10}\Delta P_g^*$  vs.  $\log_{10}T_g^*$  offers a nice frame for the "universal" presentation and comparison  $T_g(P)$  experimental and model based data. The crossover from  $dT_g/dP > 0 \rightarrow$  $dT_g/dP < 0$  seems to be associated with  $T_g^* \rightarrow 0.6$  and  $T_g^* \rightarrow 3.55$ in such plot. This is the key feature of the intermediate regime II. There are no unique "generic" steep anomalies. Finally, worth indicating is the general difference between  $P_g^*$  vs.  $T_g^*$  data taken from "concentrational" experiment under atmospheric pressure (1) and from the real high-pressure experiment (2) for colloidal glass formers. The case (1) for re-entrant colloidal glass former can be linked to the group of systems where  $dT_g/dP < 0$ . The characterization of the solvent is constant but the number of colloidal particles and distances between them can change when "decreasing pressure" ( $\phi \rightarrow 0$ ). For such system the problem of the absolute stability limit is absent: it is naturally related to  $P_g^* = 0$ and the negative pressures domain does not exist. For the case (2), compressing changes notably not only not only distances between colloidal particles but also properties of the solvent. Changes of density of the solvent (typically ~ 30% for  $P \approx 1$  GPa) are associated with very strong changes in dynamics, particularly near the glass temperature. In this case "rarefication" associated with the isotropic stretching and entering pressures domain can yield even stronger changes for the solvent. Stretching is terminated by the absolute stability limit spinodal in negative pressures domain. All these show that for the case (1) properties of the colloidal glass former are dominated almost exclusively by colloidal particles. In the case (2), at least equally important is the impact of the solvent.

**Figure** 7 indicates the clear link between molecular and colloidal glass formers: they follow the same patter the plot  $\log_{10} \Delta P_g^*$  vs.  $\log_{10} T_g^*$ . Model fluids based on SW and LJ potentials offer the nice frame for getting the fundamental insight into experimental data within such presentation.

## **CONCLUDING REMARKS**

This report presents proposals of few equations for describing the pressure evolution of the glass temperature beyond the dominated SG/AA pattern. They make the description of glass-forming systems where both  $dT_g/dP > 0$  and  $dT_g/dP < 0$  possible. The ways of portrayal were extended also for the evolution of  $T_g(V,\rho)$  and  $P_g(V,\rho)$ . The basic relevance of including into the analysis negative pressures and the preliminary derivative-based and distortions-sensitive analysis has been shown. From results presented the possible general pattern for  $T_g(P)$  evolution for glass-forming systems ranging from low molecular weight liquids, resins, polymer melt, liquid crystals to colloidal fluids emerges.

In the low density region the extended SG-type equation can describe experimental data. On increasing pressures, for intermediate densities, the gradual inclusion of the "damping term" can lead to the reversal (re-entrant,  $dT_g/dP < 0$ ) vitrification. However, for strongly compressed and high density systems the crossover to the second, HS-type, dependence  $T_g(P) \rightarrow P_g^{4/5}$  takes place. The crossover to this second type of vitrification can occur before reaching the maximum of  $T_g(P)$  as for glycerol or well beyond the maximum. For the model-normalized "universal" plot  $\log_{10}\Delta P_g^*$  vs.  $\log_{10}T_g^*$  such general characterization is manifested as the less or more marked S-shape. It is notable that this picture may be valid both for molecular and colloidal glass formers, although for the latter real high-pressure experiments are still required. For the dominated group of systems where  $dT_{g,m}/dP > 0$  most often the SG/AA-type  $(T_{g,m}(P))$ , Kraut–Kennedy type ( $T_{g,m}(V, \rho)$ ) or Murnaghan type ( $P_{g,m}(V, \rho)$ ) dependences are used. The discussion for the latter (Poirier, 2000; Skripov and Faizulin, 2006) indicates that notable distortions

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appears for  $\Delta V/V_0 \rightarrow 1/2$ . Taking into account the compressibility of typical molecular liquids such domain starts for  $P \sim 1.5$  GPa. In the opinion of the authors, equally important can be the distance of the reference point from the possible maximum of  $T_g(P)$ , even if it is "hidden" by a phase transition or crossover to another form of vitrification.

Finally, we would like to stress the significance of the above discussion for the glass transition physics, material engineering and geophysical and planetary studies (Donth, 2000; Poirier, 2000, Berthier and Ediger, 2016; Rodríguez-Tinoco et al., 2016; Svenson et al., 2017).

# **AUTHOR CONTRIBUTIONS**

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