1. Introduction

Simon [1] concluded from the fact that a finite entropic difference is retained between the vitreous state and the crystalline state of a pure substance in the limit $T \to 0$ ($T$: absolute temperature) that one is not dealing with an internal equilibrium state in the case of a glass but rather with a "frozen" non-equilibrium state. Within the definitions of the thermodynamics of irreversible processes, the vitreous state, therefore, has to be classified as an arrested equilibrium state [2, 3].

In the following, we will explicitly assign a (macroscopic) internal variable $\xi$ to the molecular internal degree of freedom frozen (arrested) in the vitreous state. When cooling a liquid at constant pressure $p$ at a constant and not too high rate $T = \beta < 0$, one schematically obtains the following picture, for example, for the volume $V$ (Figure 1): To begin with, one should take into account that every process which occurs at a finite rate $T \neq 0$ is an irreversible process connected with a finite entropy production. In this case, the volume follows a non-equilibrium curve $V(T, \xi)$ in which the temperature $T$ and the internal variable $\xi$ vary independently of each other. However, if one cools, proceeding from a temperature $T_0$ at a not too high rate $\beta = \text{const.}$, a volume is measured at the beginning of the process which practically follows the curve $V(T, \xi_0(T))$ of the internal equilibrium of the liquid. The development of this curve is uniquely determined by the temperature. At a temperature $T < T_0$, however the volume curve starts to considerably deviate from the equilibrium curve. Below $T_0$, the internal variable $\xi$ clearly deviates from its equilibrium value $\xi_0(T)$, and the volume curve $V(T, \xi)$ is no longer uniquely determined by the temperature. At a temperature $T_g$, the internal molecular degree of freedom then freezes with $\xi = 0$. Subsequently, the volume follows the volume curve $V(T, \xi = \text{const.})$ of the arrested equilibrium, whose development depends solely on the temperature, but whose level, on the other hand, is determined by the non-equilibrium value $\xi(T_g)$ of the internal variable frozen in at $T_g$. The glass transition region, i.e., the position of the temperature interval $(T_g', T_g)$, as well as the non-equilibrium value $\xi$, which freezes in at $T_g$, depends on the cooling rate $T = \beta < 0$. According to the statements pointed out

![Fig. 1. Volume V as a function of the temperature T when cooling a non-crystallizing melt with a not too high constant cooling rate T under constant pressure p. ξ: internal variable; ξ0(T): its equilibrium value; T0: initial temperature; Tg: temperature at which first deviations from the internal equilibrium become noticeable; Tg: so-called glass temperature; Tg': freezing temperature; Tg'': Vogel temperature (see Sect. 5).](image)

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in Sect. 5, however, the internal degree of freedom does not freeze completely at \( T'_g \). Very small changes in \( \zeta \) are still found below \( T'_g \). A complete arrest of the internal degree of freedom with \( \zeta = 0 \) is only observed normally at a considerably lower temperature \( T''_g \). The position of \( T'_g \) decreases with the cooling rate \( \beta \). In the limit \( \beta \to 0 \), i.e., in the case of a fictitious, pure equilibrium process, \( T'_g \) finally reaches the value \( T''_g \).

If one proceeds from the linearized mechanical equations of state, one obtains at constant pressure \( p \) for the internal equilibrium the straight line

\[
V = V_e[1 - \alpha_e(T - T_e)], \quad T \geq T_e
\]

and for the arrested equilibrium the straight line

\[
V = V_e[1 - \alpha_{e\infty}(T - T_e)], \quad T \leq T'_e
\]

\( \alpha_e \) and \( \alpha_{e\infty} \) are the coefficients of thermal expansion of the internal and the arrested equilibria (see Sect. 2). The two intersect at \( T'_e \), the so-called glass temperature. In this approximation (the so-called “simple freezing-in model”; Rehage [3]), the glass transition region is reduced to a single temperature \( T_e \). The glass transition becomes a discontinuous process. The value \( \zeta_g \) of the internal variable, which freezes at \( T'_e \), is necessarily the equilibrium value \( \zeta_e = \zeta_e(T_e) \) in this model. The coefficient of thermal expansion suffers a discontinuous jump at \( T'_e \):

\[
\Delta^e \alpha = \alpha_e - \alpha_{e\infty}.
\]

Physically, \( T'_e \) and \( \zeta_e(T_e) \) are, of course, purely fictitious quantities. Nevertheless, \( T'_e \) is well-defined. The temperatures \( T_F \) and \( T'_e \), which are characteristic for the real glass transition, on the other hand, can only be determined rather vaguely. This is due to the fact that, as already mentioned, the volume curve measured at a finite cooling rate is a non-equilibrium curve which develops continuously in the whole range \( (T'_e, T_0) \).

2. The Fundamental Relations of the Thermodynamics of Irreversible Processes*

In the following, we will consider a homogeneous, fluid single-component system with constant mass. According to the assumptions of the thermodynamics of irreversible processes, the Gibbs formalism of equilibrium thermodynamics also holds in non-equilibrium for such systems. The Gibbs potentials, however, must be complemented by the so-called internal variables, which represent the additional degrees of freedom in non-equilibrium. In the following, we will only consider a single internal variable \( \zeta \). In the G-representation, Gibbs’ fundamental equation of the system is then

\[
g = g(T, p, \zeta) = h(T, p, \zeta) - TS(T, p, \zeta).
\]

It connects the specific free enthalpy \( g \) with the mutually independent variables \( T, p, \zeta \) of the system.

\[
s = -\left(\frac{\partial g}{\partial T}\right)_{p, \zeta} \quad \text{and} \quad h = g - T\left(\frac{\partial g}{\partial T}\right)_{p, \zeta}
\]

are the specific entropy and the specific enthalpy of the system.

In non-equilibrium, the independent variables \( T(t), p(t), \zeta(t) \) are explicit functions of time \( t \). If the temperature and the pressure of the system are also identical with the temperature and the pressure of the surroundings of the system in non-equilibrium, the changes with time \( \dot{T} = dT/dt \) and \( \dot{p} = dp/dt \) are determined by the manner of external perturbation of the system. One can assume a simple dynamic law in the form

\[
\dot{\zeta} = La, \quad L \geq 0
\]

for the change of the internal variable with time. \( a(T, p, \zeta) \) is the so-called affinity, which can be thought of as a driving force for the internal variable. The affinity is given by

\[
a = -\left(\frac{\partial g}{\partial \zeta}\right)_{T, p} = T\sigma_{Tp} - \eta_{Tp},
\]

whereby

\[
\sigma_{Tp} = \left(\frac{\partial s}{\partial \zeta}\right)_{T, p} \quad \text{and} \quad \eta_{Tp} = \left(\frac{\partial h}{\partial \zeta}\right)_{T, p}
\]

denote the partial specific entropy and the partial specific enthalpy relative to the relevant internal degree of freedom.

Thermodynamic quantities which refer to an internal equilibrium state are indicated with “e”. In the internal equilibrium, the affinity goes to zero. Thus, an internal equilibrium state is characterized by

\[
a_e = 0 \quad \text{or} \quad T\sigma_{Tp}^e = \eta_{Tp}^e.
\]

This equation fixes one of the variables \( T, p, \zeta \) as a function of the other, for example

\[
\zeta = \zeta_e(T, p).
\]

Although the internal variable remains variable in internal equilibrium, it is no longer an independent variable.

* For example, compare [4–6] or [26].
In addition to (6a),

\[(da)_{e} = 0 \] (6b)

must be fulfilled in the internal equilibrium. For the change in the affinity with the independent variables, one gets:

\[da = \left(\frac{\partial a}{\partial T}\right)_{p, \zeta} dT + \left(\frac{\partial a}{\partial p}\right)_{T, \zeta} dp + \left(\frac{\partial a}{\partial \zeta}\right)_{T, p} d\zeta. \] (7)

According to (2) and (4), one obtains in the internal equilibrium:

\[\left(\frac{\partial a}{\partial T}\right)_{p, \zeta} = -\left(\frac{\partial^{2} g}{\partial T \partial \zeta}\right)_{e} = -\left(\frac{\partial^{2} g}{\partial \zeta \partial T}\right)_{e} = \sigma_{T}^{e}, \]

\[\left(\frac{\partial a}{\partial \zeta}\right)_{T, p} = -\left(\frac{\partial^{2} g}{\partial \zeta^{2}}\right)_{T, p} \equiv -\gamma_{T}^{e}. \]

Hence, according to (6b), the change of the internal variable with the temperature at constant pressure in the internal equilibrium is given by

\[\left(\frac{d\zeta}{dT}\right)_{p} = \frac{\sigma_{T}^{e}}{\gamma_{T}^{e}}. \] (8)

In addition, we have in a stable or metastable internal equilibrium

\[\gamma_{T}^{e} > 0. \] (9)

For the change in the entropy with respect to the independent variables, one obtains

\[ds = \left(\frac{\partial s}{\partial T}\right)_{p, \zeta} dT + \left(\frac{\partial s}{\partial p}\right)_{T, \zeta} dp + \left(\frac{\partial s}{\partial \zeta}\right)_{T, p} d\zeta. \] (10)

For the specific heat capacity at constant pressure, this leads to

\[c_{p} = T \left(\frac{\partial s}{\partial T}\right)_{p, \zeta} = T \left(\frac{\partial s}{\partial T}\right)_{p, \zeta} + T \sigma_{T}^{e} \left(\frac{d\zeta}{dT}\right)_{p}. \] (11)

Here,

\[c_{p, \zeta} = T \left(\frac{\partial s}{\partial T}\right)_{p, \zeta} \] (12)

is the specific heat capacity of the arrested equilibrium (\(\zeta = \text{const.}\)). This heat capacity is measured when the change in temperature occurs so rapidly that the internal degree of freedom is not capable of following this change.

\[\left(\frac{\zeta}{T} \right) \text{ or if the internal degree of freedom is frozen} \]

\[\left(\zeta = 0 \right) \text{ because of} \; L \to 0; \text{ see below}. \]

In the internal equilibrium, one measures, according to (8), the heat capacity

\[c_{p}^{e} = c_{p, \zeta} + \Delta c_{p} \] (13a)

with

\[\Delta c_{p} = \frac{T \sigma_{T}^{e} \zeta}{\gamma_{T}^{e}} \geq 0. \] (13b)

\[\Delta c_{p} \] is the contribution of the internal degree of freedom to the heat capacity in the internal equilibrium. Outside the equilibrium, i.e., in non-equilibrium, \(T \) and \( \zeta \) are mutually independent but time-dependent variables. With \( p = \text{const.} \), one must then set

\[\left(\frac{d\zeta}{dT}\right)_{p} = \frac{\zeta}{T}. \] (14)

Under conditions of non-equilibrium, one measures, according to (11), the heat capacity

\[c_{p} = c_{p, \zeta} + \Delta c_{p} \] (15a)

with

\[\Delta c_{p} = T \sigma_{T}^{e} \frac{\zeta}{T}. \] (15b)

\[\Delta c_{p} \] is the contribution of the internal degree of freedom to the heat capacity under conditions of non-equilibrium.

Corresponding expressions can be derived for the coefficient of thermal expansion \( \alpha \) and for the isothermal compressibility \( \kappa_{T} \). One obtains as the coefficient of thermal expansion:

\[\alpha = \frac{1}{v} \left(\frac{\partial v}{\partial T}\right)_{p} \]

\[= \frac{1}{v} \left(\frac{\partial v}{\partial T}\right)_{p, \zeta} + \frac{1}{v} \varphi_{T} \left(\frac{d\zeta}{dT}\right)_{p}. \] (16)

and as the compressibility:

\[\kappa_{T} = -\frac{1}{v} \left(\frac{\partial v}{\partial p}\right)_{T} \]

\[= -\frac{1}{v} \left(\frac{\partial v}{\partial p}\right)_{T, \zeta} - \frac{1}{v} \varphi_{T} \left(\frac{d\zeta}{dp}\right)_{T}. \] (17)

\[\varphi_{T} \equiv \left(\frac{\partial v}{\partial \zeta}\right)_{T, p} \] (18)
is the partial specific volume with respect to the internal degree of freedom, \( v \) the specific volume of the system. For the change of the internal variable as a function of pressure at constant temperature and in the internal equilibrium, one obtains, corresponding to (8),

\[
\left( \frac{d\xi}{dp} \right)_T \equiv -\frac{\varphi_{\xi_T}^e}{\gamma_{\xi_T}^e}.
\]  

(19)

\( \alpha_{\xi} \equiv \frac{1}{v} \left( \frac{d\varphi}{dT} \right)_{p,\xi} \) and \( \kappa_{\xi_T} \equiv -\frac{1}{v} \left( \frac{d\varphi}{dp} \right)_{T,\xi} \)

are the response functions which are measured in the arrested equilibrium. According to (8) and (16), the contribution of the internal degree of freedom to the coefficient of thermal expansion \( \alpha_{\xi_T} \) under conditions of internal equilibrium is

\[
\Delta^e \alpha = \frac{1}{v} \varphi_{\xi_T}^e \gamma_{\xi_T}^e.
\]

(20)

According to (17) and (19), the contribution of the internal degree of freedom to the compressibility \( \kappa_{\xi_T}^e \) of the internal equilibrium is

\[
\Delta^e \kappa_T = \frac{1}{v} \left( \varphi_{\xi_T}^e \right)^2 \geq 0.
\]

(21)

As can easily be seen, the differences (13b), (20), and (21) fulfill the Davies relation

\[
\Delta^c \Delta^e \geq T \nu (\Delta^e \alpha)^2.
\]

(22a)

[7, 8] (compare also [4]). It should be stressed that the Davies relation holds for every arbitrary internal equilibrium state \( Z[T, p, \xi_{\xi_T}(T, p)] \). Hence, it does not have anything to do with the formally homologous Ehrenfest relation, which is fulfilled on the line \( T_{\xi}(p_{\xi}) \) of a second-order transition. In (22), the differences \( \Delta^e \) refer to the response functions, which are measured during a quasi-static perturbation \([\xi = \xi_{\xi_T}(T, p); \text{internal equilibrium}]\) and during a sufficiently fast perturbation \((\xi_{\xi} = \text{const.}; \text{arrested equilibrium})\). If several internal degrees of freedom are simultaneously macroscopically relevant, (22a) is replaced by the Davies inequality [9]

\[
\Delta^c \Delta^e \geq T \nu (\Delta^e \alpha)^2.
\]

(22b)

The contributions of the internal degree of freedom to the coefficient of thermal expansion or to the compressibility under conditions of non-equilibrium are given by

\[
\Delta \alpha = \frac{1}{v} \varphi_{\xi_T} \frac{\zeta_{\xi_T}}{\tilde{T}_p} / \tilde{T}_p \quad \text{or}
\]

\[
\Delta \kappa_T = -\frac{1}{v} \varphi_{\xi_T} \frac{\zeta_T}{\tilde{T}_p} / \tilde{T}_p.
\]

(23)

One must distinguish here if the process is run at constant pressure or at constant temperature. The differences (15b) and (23) do not fulfill the Davies relation (22). Since the coefficient of thermal expansion, defined by (16), must be measured at constant pressure, it would be more correct to write \( \alpha_{\xi} \) instead of \( \alpha \) and in (23) \( \Delta \alpha_{\xi} \) instead of \( \Delta \alpha \). According to the equations of equilibrium thermodynamics, we can then also define an isothermal coefficient of expansion

\[
\alpha_T = -\frac{1}{v} \left( \frac{d\sigma}{dp} \right)_T.
\]

\[
\Delta \alpha_T = -\frac{1}{v} \varphi_{\xi_T} \frac{\zeta_T}{\tilde{T}_p} / \tilde{T}_p.
\]

In non-equilibrium, the contribution of the internal degree of freedom to this coefficient is

\[
\Delta \alpha_{\xi_T} = -\frac{1}{v} \varphi_{\xi_T} \frac{\zeta_{\xi_T}}{\tilde{T}_p} / \tilde{T}_p.
\]

Obviously, we now have in place of (22a)

\[
\Delta \alpha_{\xi_T} \Delta \kappa_T = T \nu (\Delta \alpha_{\xi_T})^2.
\]

However, in non-equilibrium, in general, \( \Delta \alpha_{\xi} \neq \Delta \alpha_T \).

It should be noted that the product \( L \gamma_{\xi_T}^p \) has the dimension of reciprocal time if the internal variable is dimensionless (e.g., a concentration).

\[
\tau_{\xi_T}^P(T, p) = 1/L \gamma_{\xi_T}^P
\]

(24)

proves to be the so-called Debye relaxation time of the system. Analogously, one can introduce the relaxation time

\[
\tau_{\xi_T}^P(T, p, \xi) = 1/L \gamma_{\xi_T}^P
\]

(25)

with

\[
\gamma_{\xi_T}^P = \left( \frac{\partial^2 \xi}{\partial \xi_T^2} \right)_{T, p}.
\]

(26)

In contrast to the Debye relaxation time, the relaxation time (25) also depends on the internal variable. In order to determine the internal variable as a function of time, one obtains, if the pressure \( p \) and the coupling factor \( L \)
are constant, from (3) and (7) with (4) and (25) the non-linear differential equation:

\[ \ddot{\zeta} + \frac{1}{\tau_{Tp}} \dot{\zeta} = -L \left( \frac{\partial^2 g}{\partial T \partial \zeta} \right) \dot{T}. \]  

(27a)

If \( \eta_{Tp} \) and \( \sigma_{Tp} \) are not dependent on the temperature [e.g., as in the case of the Eqs. (38) and (39) in Sect. 3], we have, as in the equilibrium,

\[ -\frac{\partial^2 g}{\partial T \partial \zeta} = \sigma_{Tp}. \]  

(27b)

Equation \( \dot{\zeta} = 0 \) holds for the frozen (arrested) states. According to (3), however, \( \dot{\zeta} \) does not disappear because of \( a = 0 \) (internal equilibrium), but because of \( L \rightarrow 0 \). Hence, when describing a freezing process, e.g. the glass transition, \( L \) must necessarily be regarded as a variable, i.e., as a function \( L(T, p, \zeta) \) of the mutually independent variables of the system. In this case, the differential equation (27) is formally maintained if the relaxation time \( \tau_{Tp} \) is replaced by the effective relaxation time

\[ \frac{1}{\tau_{eff}} = \frac{1}{\tau_{Tp}} \left( \frac{\partial \ln L}{\partial T} \right)_{p, \zeta} \dot{T} - \left( \frac{\partial \ln L}{\partial \zeta} \right)_{T,p} \dot{\zeta}. \]  

(28)

Thus, the effective relaxation time \( \tau_{eff} \) of a freezing process under constant pressure depends not only on the independent variables \( T, \zeta \), but also on their rates \( \dot{T}, \dot{\zeta} \).

3. A Gibbs Fundamental Equation for the Description of the Glass Transition

A fundamental problem when describing the glass transition within the framework of phenomenological thermodynamics is the formulation of a suitable Gibbs fundamental equation of the melt. Here, we proceed from the simple model of the Eyring liquid [10–12]. Eyring regards liquids as a mixture of molecules and vacancies. What freezes during the glass transition of a melt, is without doubt the diffusive translational motion of the molecules. The concentration of the vacancies or the volume fraction of the vacancies (the relative free volume) of the Eyring liquid can serve as a measure of the intensity of the diffusive translational motion. In this case, the diffusion process should not be understood as a hopping of the molecules into holes of the same size. In a relatively compact melt, the diffusion process is mainly a cooperative process, for which a considerably smaller vacancy size is sufficient [13]. We thus proceed from a mixture whose species differ widely in their volumes.

According to Flory [14, 15], one can write for the free enthalpy \( G \) of a mixture of molecules with two different volumes:

\[ G = G_0 + RT \left[ N_1 \ln \frac{N_1}{N_1 + \rho N_2} + N_2 \ln \frac{\rho N_2}{N_1 + \rho N_2} \right] + G_E. \]  

(29)

\[ \rho \equiv v_2/v_1 > 1 \]  

(30)

is the ratio of the molar volumes of the two mixing partners, \( N_1 \) and \( N_2 \) their mole numbers, and \( R \) the gas constant. With respect to the Eyring liquid, \( N_1 \) designates the mole number of the vacancies* and \( N_2 \) the mole number of the material particles (molecules of a low-molecular-weight substance or mobile units of a flexible polymer). The standard term \( G_0 \) is solely determined by the material particles, as the vacancies in the pure state do not possess a chemical potential \( \mu \):

\[ G_0 = \mu_1^0 N_1, \mu_1^0 = 0. \]  

(31)

For the deduction of an expression for the excess free enthalpy \( G_E \), we proceed from a simple mixture as defined by Guggenheim [15]:

\[ G_E = zN \frac{N_1 N_2}{N_1 + N_2}. \]  

Here, however, we substitute with

\[ zN_i \rightarrow o_i N_i \]  

the coordination number \( z \) by the contact surfaces \( o_i \) per mole of the mixing partners [16]. It is obvious that the interaction between partners of different sizes and shapes can be described much better by the surfaces \( o_i \) than by the mere number \( z \) of the nearest neighbours. Hence, the excess free enthalpy should amount to

\[ G_E = \Delta w \frac{N_1 o_1 N_2}{N_1 + o_2 N_2}. \]  

In

\[ \Delta w = w_{12} - \frac{1}{2} (w_{11} + w_{22}) \]

The mole number and the number of vacancies are fictitious quantities, since the vacancies have no mass and, continuously smeared, do not have an individuality. According to (34) and (37), on the other hand, a real physical significance is attributed to the relative free volume \( \varphi \), introduced instead of \( N_1 \) further down, and to the internal variable \( \zeta \).
describes the binding energy between two adjacent material particles and \( w_{12} = 0 \) the binding energy between two vacancies. \( w_{12} \) can be interpreted as the binding energy between two material particles separated by vacancies.

We approximatively assume that the excess free enthalpy \( G_E \) neither depends on the temperature nor the pressure. The temperature independence means that we confine ourselves to a regular mixture with

\[
G_E = H_E, \quad S_E = 0.
\]

The pressure independence allows an equation of the molar volumes \( v_i \) with the partial molar volumes \( \bar{v}_i \). In place of the mole numbers \( N_i \), we can then easily introduce the volume fractions

\[
\varphi_1 = \frac{\bar{v}_1 N_1}{v_1 N_1 + \bar{v}_2 N_2} = \frac{N_1}{N_1 + \rho N_2}; \quad \varphi_2 = 1 - \varphi_1
\]

as variables of composition. In the following, we will use the volume fraction \( \varphi = \varphi_1 \) of the vacancies (the relative free volume) as the independent variable of the composition. Furthermore, we will refer all the extensive quantities to the mole number \( N_2 \) of the material particles and once again denote these molar quantities with small letters

\[
g/N_2 = g, \quad v/N_2 = v, \quad c_p/N_2 = c_p etc.
\]

(29-32) then yield as the Gibbs fundamental equation of the melt interspersed with vacancies:

\[
g = g_0 + RT \left[ \frac{\rho \varphi}{1 - \varphi} \ln \varphi + \ln (1 - \varphi) \right] + g_E \quad (33a)
\]

with

\[
g_0 = \mu_e^0(T, \rho)
\]

(33b)

and

\[
g_E = h_E = h_c \frac{\rho \varphi}{1 - \varphi + \lambda \varphi}.
\]

(33c)

\[
h_c = o_1 \Delta w
\]

(33d)

is the energy required to generate one mole of vacancies in the vacancy-free melt.

\[
\lambda = \frac{o_1}{o_2}, \quad i.e., \quad \frac{o_1}{o_2} = \lambda \frac{v_1}{v_2},
\]

(33e)

is a geometric factor which not only considers the different sizes but also the different shapes of the mixing partners. As dimensionless internal variable, we will finally introduce the ratio \( \zeta = N_1/N_2 \) of the mole number of the vacancies to the mole number of the material particles. The internal variable, which is often rather vaguely referred to as the “ordering parameter” in the literature (e.g., compare [3], [8], [17]), thus acquires a concrete physical meaning. With the relative free volume \( \varphi \), we find the interrelation

\[
\zeta = \frac{\rho \varphi}{1 - \varphi} \quad ; \quad d\zeta = \frac{\rho}{(1 - \varphi)^2} d\varphi.
\]

(34)

According to (2) and (33), the entropy and enthalpy of the system are given by

\[
s = -\left( \frac{\partial g_0}{\partial T} \right)_p - R \left[ \frac{\partial \varphi}{1 - \varphi} \ln \varphi + \ln (1 - \varphi) \right], \quad (35)
\]

\[
h = h_0 + h_E \quad \text{with} \quad h_0 = g_0 - T \left( \frac{\partial g_0}{\partial T} \right)_p. \quad (36)
\]

One obtains for the volume

\[
v = v_1 \zeta + v_2 = \frac{v_2}{1 - \varphi}.
\]

(37)

The vacancies are not of importance in the response functions \( c_p, \alpha, \kappa \) of the arrested equilibrium. According to (12) and (35), one obtains, for example, for the heat capacity of the arrested equilibrium

\[
c_p, \zeta = -T \left( \frac{\partial^2 g_0}{\partial T^2} \right)_p
\]

Equations (33) and (34) yield for the quantities (5), (18), and (26), which determine the dynamics of the system

\[
\eta_{Tp} = h_c \left( \frac{1 - \varphi}{1 - \varphi + \lambda \varphi} \right)^2, \quad (38)
\]

\[
\sigma_{Tp} = -R \left[ \ln \varphi + \left(1 - \frac{1}{\rho}\right)(1 - \varphi) \right], \quad (39)
\]

\[
\varphi_{Tp} = \frac{v_1}{\rho} \frac{1 - \varphi}{\varphi},
\]

(40)

and

\[
\gamma_{Tp} = \left( \frac{\partial \eta_{Tp}}{\partial \zeta} \right)_{T, p} - T \left( \frac{\partial \sigma_{Tp}}{\partial \zeta} \right)_{T, p}, \quad (41a)
\]

with

\[
\left( \frac{\partial \eta_{Tp}}{\partial \zeta} \right)_{T, p} = -2 h_c \lambda \left( \frac{\eta_{Tp}}{h_c} \right)^{3/2}, \quad (41b)
\]

\[
\left( \frac{\partial \sigma_{Tp}}{\partial \zeta} \right)_{T, p}
\]
In consideration of the equilibrium condition (6a), one can also write for the equilibrium value of $\gamma_{TP}$:

$$\gamma_{TP}^e = RT \frac{1 - \varphi_e}{\rho}$$

$$\times \left[ \frac{2 \lambda}{1 - \varphi_e + \lambda \varphi_e} \left[ \ln \varphi_e + \left( 1 - \frac{1}{\rho} \right) (1 - \varphi_e) \right] 
+ \frac{1 - \varphi_e}{\varphi_e} \left( 1 - \varphi_e + \frac{\varphi_e}{\rho} \right) \right].$$

As already mentioned, $\tau_{\gamma_{TP}}^e = 1/L_{\gamma_{TP}}^e$ is the Debye relaxation time of the system.

In the internal equilibrium, the liquid interspersed with vacancies (33) is stable up to $T \to 0$. However, the liquid is not capable of holding an arbitrary number of vacancies. With increasing temperature, a critical point $(T_{\text{max}}; \varphi_{e_{\text{max}}})$ is reached, after which the system becomes unstable with $\gamma_{TP}^e < 0$.

4. The Glass Transition in the “Simple Freezing-in Model”

In the “simple freezing-in model” (Sect. 1), the glass transition region is reduced to the so-called glass temperature $T_g$. The glass transition appears as a discontinuous process in which an equilibrium value $\zeta_{e}(T)$ of the internal variable of the liquid state is abruptly subjected to the arrest $\zeta_e = \text{const}$. As a result, the response functions undergo a precipitous change in the amounts given by the equilibrium differences (13b), (20), and (21).

With (39), (40), and (42), (20) now leads to

$$\Delta^e \alpha = - \frac{1}{T} \frac{\varphi_e}{2 \lambda \varphi_e} + \frac{1 - \varphi_e}{1 - \varphi_e + \lambda \varphi_e} \ln \varphi_e + \left( 1 - \frac{1}{\rho} \right) (1 - \varphi_e).$$

According to (13b) and (20), we have further

$$\Delta^e c_p = vT \Delta^e \alpha \frac{\sigma_{TP}^e}{\varphi_{e_{TP}}^e},$$

i.e., with (39), (40), and (43)

$$\Delta^e c_p = \frac{R \rho \varphi_e}{1 - \varphi_e}$$

$\ln \varphi_e + \left( 1 - \frac{1}{\rho} \right) (1 - \varphi_e)$

$$+ \frac{2 \lambda \varphi_e}{1 - \varphi_e + \lambda \varphi_e} \left[ \ln \varphi_e + \left( 1 - \frac{1}{\rho} \right) (1 - \varphi_e) \right].$$

These equations hold for every arbitrary internal equilibrium state of the system (33), i.e., with the assumption $\zeta_e = \zeta_e(T_g)$ also at the glass temperature $T_g$. If $\Delta^e \alpha(T_g)$, $\Delta^e c_p(T_g)$, $\rho$, and $T_g$ are known, the relative free volume $\varphi_e(T_g)$ and the interaction parameter $\lambda$ can be determined from (43) and (44).

Using equivalent equations (under the precondition $\varphi_e \ll 1$ and omission of $1/\rho$ as compared to 1 in Eq. (39)), Kanig [16] determined the volume fraction $\varphi_{g_{\text{max}}} = \varphi_e(T_g)$ and the interaction parameter $\lambda$ for eleven different organic polymers with experimental data for $\Delta^e \alpha$, $\Delta^e c_p$, and $T_g$. For these polymers, he found on average

$$\langle \varphi_{g_{\text{max}}} \rangle = 0.0235 \pm 0.0050,$$

$$\langle \lambda \rangle = 3.15 \pm 0.35.$$
Hence, in the “simple freezing-in model” the Davies relation in the form (22a) or also in the form (22b) appears as a characteristic of the glass transition and, as such, is also designated as the Prigogine-Defay relation (e.g., compare [2], [22–24]). One should notice here that the Davies relation can, strictly speaking, never become valid at a finite cooling rate, since we are dealing with an irreversible process subject to a non-vanishing entropy production in the whole interval \((T'_g, T_0)\) (Figure 9). The Davies relation approximately holds for the states \(Z(T, p, \xi)\) in the interval \((T_F, T_0)\), as in this region the response functions still approximately follow the equilibrium curves with \(\xi = \xi_c(T, p)\). Below \(T_F\), however, the internal degree of freedom provides the “dynamic” contributions (15b) and (23) to the response functions, which do not comply with the Davies relation. These contributions continuously diminish from \(T_F\) until they approximately disappear at \(T'_g\) and finally completely disappear at \(T''_g\) (Figs. 7 and 8). With regard to the glass transition, the Davies or Prigogine-Defay relation can only attain significance upon quasi-static cooling with \(T = \beta \rightarrow 0\). According to the statements in the next section, this is the case at the temperature \(T''_g = T_w\), which is usually some ten degrees below the glass temperature.

5. The Glass Transition as an Irreversible Process

In the thermodynamics of irreversible processes, the system temperature \(T\) is defined by

\[
T(s, v, \xi) = \left( \frac{\partial u}{\partial s} \right)_{v, \xi}
\]

\((u:\) specific internal energy of the system). It reflects all the internal degrees of freedom of the system which are in internal equilibrium. The macroscopically relevant internal degree of freedom \(\xi\), on the other hand, must be regarded as being arrested with \(\xi = \text{const.}\). If \(\xi\) is assigned to the diffusive translational motion of the molecules of a liquid in non-equilibrium, one must, therefore, expect a considerable difference between the system temperature and that temperature \(T^*\) at which the heat exchange of the system with its surroundings takes place. According to Meixner [25], the differentiation between a thermostatic temperature \(T\) and a thermodynamic temperature \(T^*\) is even generally imperative in the thermodynamics of irreversible processes. This completely corresponds to the differentiation between the hydrostatic and the hydrodynamic pressures in hydromechanics. In the following, the differentiation between \(T\) and \(T^*\) finally proves to be essential for a correct formulation of the boundary conditions. \(T = T^*\) naturally holds in the internal as well as in the arrested equilibrium.

In non-equilibrium, the change in entropy \(s\) with time is separated into two additive parts:

\[
\dot{s} = \frac{d_a s}{dt} + \frac{d_i s}{dt}.
\]

\(d_a s/dt\) describes the amount of entropy exchanged by the system with its surroundings per time and mass unit. \(d_i s/dt \geq 0\) is the entropy produced in the interior of the system per time and mass unit [26]. If one distinguishes between the static temperature \(T\) and the dynamic temperature \(T^*\), one obtains

\[
\frac{d_a s}{dt} = \frac{1}{T^*} \dot{q}.
\]

\(\dot{q}\) is the heat exchanged by the system with its surroundings at the temperature \(T^*\) per time and mass unit. If we neglect the possible difference between the hydrostatic and the hydrodynamic pressures and assume \(p = \text{const.}\), we have

\[
\dot{q} = \dot{h},
\]

whereby (36) with (5) and (12) leads to

\[
\dot{h} = c_p, \dot{T} + \eta_{fp} \dot{\xi}.
\]

According to (10) with (5) and (12), the change in entropy with time is given by

\[
\dot{s} = \frac{1}{T} c_p, \dot{T} + \sigma_{fp} \dot{\xi}.
\]

Hence, according to (47), (48), and (49), the entropy produced in the interior of the system must be

\[
\frac{d_i s}{dt} = c_p, \left( 1 - \frac{1}{T^*} \right) \dot{T} + \left( \sigma_{fp} - \frac{1}{T^*} \eta_{fp} \right) \dot{\xi}.
\]

In the thermodynamics of irreversible processes, the product \(T(d_i s/dt)\) now always appears in a bilinear form of the effective “fluxes” and “forces” [26]. Therefore, we can conclude from (50) that, in the case of a differentiation between the static and the dynamic temperatures, Eq. (3) will be replaced by the dynamic laws of the form

\[
\dot{T} = L_{11} c_p, T^* - T \\
+ L_{12} T \left( \sigma_{fp} - \frac{1}{T^*} \eta_{fp} \right),
\]
\[ \dot{\zeta} = L_{21} c_p \zeta \frac{T^* - T}{T^*} + L_{22} T \left( \sigma_{T_p} - \frac{1}{T^*} \eta_{T_p} \right). \] (51b)

The matrix of the coefficients \( L_{ik} \) is positive definite because of \( \frac{d \sigma}{d T} > 0 \). If we ignore with \( L_{12} = L_{21} = 0 \) the possible interference effects, we obtain the simple dynamic laws

\[ \dot{T} = L_{11} c_p \zeta \frac{T^* - T}{T^*}, \] (52)

\[ \dot{\zeta} = L_{22} \frac{T}{T^*} (T^* \sigma_{T_p} - \eta_{T_p}). \] (53)

With a constant cooling or heating rate, we also have the differential equation

\[ T^* = \beta = \text{const.} \] (54)

For the internal variable \( \zeta \), there is, as before, a simple dynamic law of the form (3):

\[ \dot{\zeta} = L_{22} a^*. \] (55)

However, the affinity \( a \) must now be replaced by the "dynamic" affinity

\[ a^* = \frac{T}{T^*} (T^* \sigma_{T_p} - \eta_{T_p}). \] (56)

Between \( a \) and \( a^* \), we find the interrelation

\[ a^* = a + \eta_{T_p} \frac{T^* - T}{T^*}. \] (57)

According to Tool [27] (see also [8]), one can further introduce a fictive temperature

\[ T_f = \frac{\eta_{T_p}}{\sigma_{T_p}}. \] (58)

This is the temperature which the system in a non-equilibrium state \( Z(T, \zeta) \) would have if it were in internal equilibrium [see Eq. (6)]. With the fictive temperature (58), one can also write instead of (4) and (56)

\[ a = \sigma_{T_p}(T - T_f) \] (59)

and

\[ a^* = T \sigma_{T_p}(T^* - T_f)/T^*. \] (60)

The functional dependence of the coefficients \( L_{ik} = L_{ik}(T, p, \zeta) \) cannot be derived in the macroscopic-phenomenological theory. In this respect, one rather depends on ad-hoc formulations. \( L_{22} \) has the dimension fluidity/volume. With respect to the description of the glass transition, it is reasonable to assume a proportionality

\[ L_{22} \sim \frac{1}{\eta} \]

(\( \eta \): viscosity of the melt). In order to describe the temperature dependence of the viscosity, we proceed from the so-called Vogel-Fulcher-Tammann-Hesse equation [28, 29]

\[ \ln \eta = \ln \eta^0 + \frac{c}{T - T_m} \]

and correspondingly set

\[ L_{22}(t) = \]

\[ L_{22}(0) \exp \left[ \frac{c}{T^* - T_m} - \frac{1}{T^* - T_m} \right]. \] (61)

Here and in the following, the subscript "0" refers to the initial state. \( T_m \) is the Vogel temperature. At the same time, \( T_m \) and \( c \) are the invariants of the so-called Williams-Landel-Ferry equation [18, 28, 29]. With (61), (53) leads to \( \zeta \to 0 \) if \( \eta \to \infty \). Hence, the internal degree of freedom freezes completely at the Vogel temperature. \( T_m \) is identical with the temperature denoted with \( T'' \) in Figure 1.

The boundary conditions should particularly be considered when determining the coefficient \( L_{11} \). Proceeding from an internal equilibrium state \( Z(T_0', \zeta_0(T_0')) \), we cool the melt under constant pressure at the constant rate \( T^* = \beta \). In the initial state, we have \( T_0 = T_i = T_0' \) and \( \dot{T}_0 = \dot{T}_i = 0 \). Approaching the Vogel temperature, the internal degree of freedom is arrested with a non-equilibrium value \( a^* = a = 0 \). Hence, \( T^* = T \neq T_i \) and \( T^*_f = \beta = T_f = 0 \) must hold in the limit \( T^* \to T_m \). Moreover, according to (8) and (14),

\[ \lim_{T^* \to T_m} \frac{\dot{\zeta}}{T} = \left( \frac{d^2 \zeta}{dT^2} \right)_p \]

must hold if the transition from the internal equilibrium to the non-equilibrium is supposed to proceed continuously in \( T \) and \( \zeta \) at the beginning of the process. With (8), (52), (55), and (60), this leads to

\[ \lim_{T^* \to T_m} \frac{L_{22} T \sigma_{T_p} (T^* - T_f)}{L_{11} c_p \zeta (T^* - T)} = \frac{\sigma_{T_p}^0}{\gamma_{T_p}^0} \]

and because of

\[ \lim_{T^* \to T_m} \frac{T^* - T_f}{T^* - T} = 1. \]
to

$$L_{11}(0) = \frac{T_0}{c_{p,\zeta}} L_{22}(0) \gamma_{T_p}^0 = \frac{T_0}{c_{p,\zeta}} \frac{T_0}{\tau_{T_p}}.$$ (62)

The boundary conditions for $T^* \to T_\infty$ are satisfied if one first generalizes (62) as

$$L_{11} = \frac{T}{c_{p,\zeta} \tau_{T_p}}$$

and replaces the relaxation time $\tau_{T_p}$ by the effective relaxation time (28). According to (28) and (61), one obtains for $\beta < 0$:

$$\frac{1}{\tau_{eff}} = \frac{1}{\tau_{T_p}} \frac{c \beta}{(T^*-T_\infty)^2} + \frac{1}{\tau_{T_p}} \frac{c |\beta|}{(T^*-T_\infty)^2}.$$ (63)

In order to maintain the limiting value (62), we should then set

$$L_{11} = \frac{T}{c_{p,\zeta}} \left[ L_{22} \gamma_{T_p} + c |\beta| \right] \left( \frac{1}{(T^*-T_\infty)^2} - \frac{1}{(T_0^*-T_\infty)^2} \right).$$ (63)

Equation (63) can also be used for $\beta > 0$, since $L_{11} > 0$ is always valid regardless of the sign of $\beta$.

The differential equations (52–54) with the phenomenological coefficients (61) and (63), and the thermodynamic coefficients (38–41) are thus the determining equations for the glass transition. Numerical solutions of these equations are represented in the following figures. For the constants of the equations, we will choose values which approximately apply for polystyrene:

$$h_c = 10 \text{ kJ/mol}, \quad \rho = 10, \quad \lambda = 3,$$ (64a)

according to [16],

$$T_\infty = 340 \text{ K}, \quad c = 400 \text{ K},$$ (64b)

according to [28], and with $T_0 = T_0^* = 400 \text{ K}$:

$$L_{22}(0) = 0.0001 \text{ mol/J sec}.$$ (64c)

The heat capacity $c_{p,\zeta}$ of the arrested equilibrium only plays a role in Figs. 6 and 9. There, we assume

$$c_{p,\zeta} = 0.75 (T - T_0) + c_{p,\zeta}(T_0) \quad \text{with} \quad c_{p,\zeta}(T_0) = 150 \text{ J/mol K}.$$ (64d)

Strictly speaking, the quantities (64a) are naturally not constant. The values assumed with (64a) have the effect that the melt in internal equilibrium already loses its stability towards higher temperatures at

$$T_\infty^e = 456 \text{ K}, \quad \phi_e^\infty = 0.125.$$

However, this should at least not qualitatively influence the glass transition to be described.

In Fig. 2, the relative free volume $\phi$ is represented as a function of the temperature $T^*$ for different cooling rates $T^* = \beta < 0$. It becomes obvious that at normal cooling rates the internal degree of freedom freezes practically at approximately 30 K above the Vogel temperature $T_\infty = 340 \text{ K}$. The frozen value of $\phi$ is higher the faster the cooling. It is much higher than the value $\phi_e(T_\infty) = 0.01679$, which would be reached upon quasi-

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Fig. 2. Relative free volume $\phi$ as a function of the dynamic temperature $T^*$ when cooling at different constant rates $T^* = \beta$ according to (52–54) with (61), (63), (64) and (34), (38), (39). a) $\beta = -5 \text{ K/min}$; b) $\beta = -1 \text{ K/min}$; c) $\beta = -0.2 \text{ K/min}$. According to (6), (38), (39), the equilibrium value of the relative free volume is at the initial temperature $T_0^* = 400 \text{ K}$: $\phi_e = 0.038017$, at $T^* = 370 \text{ K}$: $\phi_e = 0.025343$, and at the Vogel temperature $T_\infty'' = T_\infty = 340 \text{ K}$: $\phi_e = 0.016791$.

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Fig. 3. Relative free volume $\phi$ as a function of the dynamic temperature $T^*$ when cooling at the rate $\beta = -1 \text{ K/min}$ down to $T^* = 350 \text{ K}$ and subsequently heating at the rate $\beta = +1 \text{ K/min}$.
static cooling ($\beta \to 0$) at the Vogel temperature. If the melt is cooled down to the freezing range at the rate $\beta < 0$ and then immediately heated up again at the same rate $|\beta|$, the heat capacity goes through a hysteresis cycle (Figure 3). The cause for this can be found in the affinity $a^*$, which with $\zeta < 0$ necessarily assumes negative values when cooled. If one heats up again, $a^*$ at first stays negative. $\zeta$ and $\phi$ decrease further until $a^*$ becomes positive. Upon heating, $a^*$ must necessarily approach the equilibrium value $a^* = 0$ from the positive side. Therefore, $a^*$ first intersects the equilibrium axis and passes through a maximum within the positive range in order to finally reach the equilibrium value $a^* = 0$ “from above” (Figure 4).

According to (52), if $\dot{T}^*$ and $\dot{T}$ have the same sign, the static temperature $T$ is always above the dynamic temperature $T^*$ when cooling and always below it when heating. The temperature of the system retards with respect to the temperature of the surroundings. The maximum distance $|T^* - T|$ is smaller, the smaller the cooling or heating rate. In the limit $\beta \to 0$ we, of course, have $T^* - T = 0$. Upon cooling, because of the proximity to the internal equilibrium, $T$ first follows the fictive temperature $T_f$ [Eq. (59) with $a = 0$] and then turns towards the dynamic temperature (arrested equilibrium with $T = T^* = T_f$) (Figure 5). The fictive temperature $T_f$ freezes in with $\phi$ during the glass transition. Upon heating, $T_f$, like $\phi$, passes through a hysteresis cycle (Figure 5). The behaviour of the static temperature $T$ is, of course, completely a result of our formulation (63). Except for “pathological” cases, however, a different behaviour does not seem to be possible – at least qualitatively.

In Fig. 6, the heat capacity $c_p$, according to (15), is represented for different cooling rates. The faster the cooling, the faster the curves separate from the equilibrium curve, and the faster the melt freezes. The width of the freezing range increases with the cooling rate. The bar on the right of the figure indicates the jump $\Delta c_p$, which the heat capacity would suffer upon quasi-static cooling ($\beta \to 0$) at the Vogel temperature $T_v = 340$ K. Only this quantity satisfies the Davies relation (22). It is considerably smaller than the step in the heat capacity which one finds in the freezing range about 30 K above the Vogel temperature in the case of normal cooling rates.

If the melt is cooled down to the freezing range at the rate $\beta < 0$ and then heated up again at the same rate $|\beta|$, the heat capacity $\Delta c_p$, according to (15b), first passes through a minimum and then through a maximum during the heating process (Figure 7). This effect is also entirely due to the affinity. According to (15b), we have

$$\Delta c_p = L_{22} \frac{\alpha^*}{T} \sigma_{T_p} \frac{T}{\dot{T}}.$$
Fig. 6. Heat capacity per mole of the material particles $c_p$, according to (15), as a function of the dynamic temperature $T^*$ when cooling at the rates a) $\beta = -0.2 \text{ K/min}$; b) $\beta = -1 \text{ K/min}$; c) $\beta = -5 \text{ K/min}$. $\Delta c_p(T^*)$: jump in the heat capacity which, according to (13), occurs upon quasi-static cooling at the Vogel temperature $T_v = 340 \text{ K}$.

Fig. 7. Contribution $\Delta c_p$ of the vacancies to the heat capacity, according to (15b), when cooling from $400 \text{ K}$ down to $350 \text{ K}$ at the rate $\beta = -1 \text{ K/min}$ and subsequently heating at the rates a) $\beta = 0.2 \text{ K/min}$; b) $\beta = 1 \text{ K/min}$; c) $\beta = 5 \text{ K/min}$.

Fig. 8. Heat capacity $\Delta c_p$, according to (15b), when cooling down to $T^* = 350 \text{ K}$ at the rate $\beta = -1 \text{ K/min}$, “annealing” at $T^* = 350 \text{ K}$, and subsequently heating at the rate $\beta = +1 \text{ K/min}$. The initial values of the free volume $\varphi$ upon heating were a) $\varphi = 0.0268907$ (the value which was attained upon cooling at $350 \text{ K}$), b) $\varphi = 0.0240000$, c) $\varphi = \varphi_e = 0.0193056$ (the equilibrium value of the melt at $350 \text{ K}$).

Fig. 9. Entropy production $d_s/dt$, according to (50), when cooling down to $T^* = 350 \text{ K}$ at the rate $\beta = -1 \text{ K/min}$ and subsequently heating at the rate $\beta = +1 \text{ K/min}$. Although the free volume seems to be fully frozen below $365 \text{ K}$ (see Fig. 3), there is still a considerable entropy production in this range. When cooling, the entropy production only disappears after reaching the Vogel temperature (at $340 \text{ K}$). The position of the maxima of the entropy production indicates rather precisely the temperature at which the freezing of the free volume seems to be complete or alternatively the thawing of the free volume starts (compare Fig. 9 with Figure 3).

The initial values of the free volume $\varphi$ upon heating were a) $\varphi = 0.0268907$ (the value which was attained upon cooling at $350 \text{ K}$), b) $\varphi = 0.0240000$, c) $\varphi = \varphi_e = 0.0193056$ (the equilibrium value of the melt at $350 \text{ K}$).

In a metastable state, it is possible that the free volume decreases even beyond the equilibrium value $\varphi_e(T^*)$ upon annealing, and approaches the equilibrium value $\varphi_e(T^*) < \varphi_e(T^*)$ of the crystalline state. When judged with respect to the potential $g$ of the melt, the states $\varphi(t) < \varphi_e(T^*)$ are non-equilibrium states with a positive affinity. If one heats up from such a state, the maximum of the heat capacity increases further. With $\phi(t) \rightarrow 0$, we finally obtain $\phi^* \rightarrow \infty$ and $\Delta c_p \rightarrow \infty$. The increase in the maximum of the heat capacity with the annealing time,
which is measured when heating a glassy frozen, annealed melt, has repeatedly led to the conclusion in the literature that the glass transition must be a masked phase transition in the sense of equilibrium thermodynamics [21]. But this is definitely not true. In these cases, the maximum of the heat capacity is clearly connected with the maximum of the affinity $a^* \neq 0$, which is characteristic for the non-equilibrium.

The entropy production $d_s/dt$, according to (59), characteristic for the irreversibility of the processes, is represented in Fig. 9 for the case of the process Fig. 3 and Fig. 7b. The entropy production disappears only at the beginning of the process at $T^* = T_0$ (internal equilibrium) and at the end of the process at $T^* = T_m$ (arrested equilibrium). In addition, it becomes very small if one switches from $T^* = -\beta$ to $T^* = +\beta$ in the range $T_m < T^* < T_g$ (in Fig. 9 at $T^* = 350 \, \text{K}$). $T$ passes through zero when switching (see further below), whereas $\zeta$ is very small.

The entropy production reaches a maximum value when cooling at the end of the freezing process and when heating at the beginning of the thawing process (compare Fig. 9 with Figure 3). The level of the maxima increases with |$\beta$|. Figure 9 shows that the glass transition generated at a finite rate $T^*$ is an irreversible process in the whole interval ($T_m$, $T_0$). However, the contribution of the entropy production to the balance (47) of the entropy is relatively small during normal perturbation of the system. The maximum contribution of the entropy production to the balance is $0.95\%$ with $\beta = -5 \, \text{K/min}$, $0.71\%$ with $\beta = -1 \, \text{K/min}$, and $0.55\%$ with $\beta = -0.2 \, \text{K/min}$.

According to (15), one can also write

$$\dot{\zeta} = \frac{\Delta c_p}{T \sigma T_p} \dot{T}. \tag{65}$$

Insertion of this expression into (49) leads to the interrelation

$$\dot{s} = \frac{1}{T} c_p \dot{T}, \tag{66}$$

which completely corresponds to the equilibrium thermodynamics. However, the conclusion

$$\dot{q} = c_p \dot{T} \tag{66}$$

would be wrong, as $\dot{s}$ contains the entropy production which, according to Clausius, corresponds to a non-compensable heat. Instead of (66), (48) and (65) lead to the correct relation

$$\dot{q} = \left( c_p, c^*_p + \frac{\eta TP}{T \sigma T_p} \Delta c_p \right) \dot{T}. \tag{67}$$

Finally, one should point out the following: $\dot{T}$ reacts spontaneously to a discontinuous change from $T^* = -|\beta|$ to $T^* = +|\beta|$ only in the arrested equilibrium. Above $T_m$, $T$ is subject to a retardation with respect to $T^*$ (Figure 5). $T$ then reacts to a discontinuous change of sign from $T^* < 0$ to $T^* > 0$ with a rapid, but still continuous change from $T < 0$ to $T > 0$. Therefore, $T$ necessarily passes through zero during this transition. If $\xi \neq 0$ is valid, $\Delta c_p$, according to (15), passes through a singular infinity due to the sign reversal of $T^*$. $\zeta = 0$ holds if the sign reversal takes place at a temperature $T^*$ within the range $T_m < T^* < T_g$. The temperature interval $\Delta T^*$, in which the singularity of $\Delta c_p$ becomes apparent, is then so small that the singularity can easily be overseen when solving the differential equations (52--54) with finite time steps $\Delta t$ (as in Figs. 7 and 8). In order to avoid the singularity, one must, in addition, postulate that with the sign reversal of the "flux" $T^*$, the sign of the "force" $T^* - T$ also has to be changed. At the instant of the sign reversal of $T^*$, $T$ must then be replaced by $2T^* - T$.

One should point out further that $\varphi$ can also be introduced as an internal variable in place of $\zeta = N_1/N_2$. With $L_{22}(0) = 0.000000925 \, \text{mol/J sec}$ instead of (64c), one obtains practically the same results.

**6. Conclusions**

The thermodynamics of irreversible processes allows an, at least qualitative, understanding of the dynamics of the processes in the glass transition region on the basis of simple dynamic laws and a simple, certainly only approximately valid Gibbs fundamental equation. The glass transition generated with a finite cooling or heating rate is a dissipative process. Such processes require a clear differentiation between equilibrium and non-equilibrium quantities. In the thermodynamics of irreversible processes, this distinction is given *a priori*. The coupling coefficient $L_{22}$ and the affinity $a^*$ appear as determining quantities for the glass transition in the thermodynamics of irreversible processes. We associated the coupling coefficient with the fluidity of the melt according to Vogel, Fulcher, Tamman, and Hesse. Except for the sign, the affinity $a^*$ is essentially given by the partial free enthalpy of the vacancies in the melt. However, it still contains a certain correction which takes into account that in non-equilibrium the system temperature retards with respect to the temperature at which the heat exchange with the surroundings takes place.
It becomes clear that the glass transition is a typical non-linear phenomenon. An explanation of the processes within the linear response theory, especially by the superposition of a spectrum of linear relaxation mechanisms, is, therefore, not very useful. Moreover, relaxation times prove to be auxiliary quantities, whose practical usefulness disappears when the differential equations describing the dynamics of the system are explicitly known. The Davies of Prigogine-Defay relations, which are often considered to be characteristic for the glass transition, do not have any significance in an irreversible process. These relations only become relevant with respect to the glass transition after a fictitious quasi-static cooling at the so-called Vogel temperature, which, according to our results, lies considerably below the temperature \( T_g \) usually designated as the glass temperature.

The fact that, according to our statements, the glass transition occurs during quasi-static cooling only at the Vogel temperature \( T_v \) some ten degrees below \( T_g \), reminds us of the thesis of Gibbs et al. [31, 32], according to which the glass transition is supposed to occur as a second-order transition some ten degrees below \( T_g \) upon quasi-static cooling. As a matter of fact, the glass transition, which we find at \( T_v \) with \( T_s = 0 \), formally exhibits some characteristics of an Ehrenfest second-order transition. However, this is only true as long as we only consider a variation of the temperature. In the case of a pressure variation, the equilibrium value \( \zeta_e(T, p) \), which freezes at \( T_v(p) \), becomes pressure-dependent. This fact alone proves that a freezing process can not be a second-order transition as defined by Ehrenfest [3]. Moreover, the quasi-static glass transition at the Vogel temperature \( T_v \) appears as a limiting case with the dynamic quantities \( \beta \to 0, L_{22} \to 0, or \eta \to \infty \) within the framework of the phenomenological thermodynamics of irreversible processes. In the pure equilibrium theory, dynamic quantities, such as \( \beta, L_{22}, \eta, or a^* \) are completely irrelevant.