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A brief critique of the Adam-Gibbs entropy model

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ABSTRACT

This paper critically discusses the entropy model proposed by Adam and Gibbs in 1965 for the dramatic temperature dependence of glass-forming liquids' average relaxation time, which is one of the most influential models during the last four decades. We discuss the Adam–Gibbs model's theoretical bases as well as its reported experimental model confirmations; in the process of doing this a number of problems with the model are identified.

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1. Introduction

Any liquid forms a glass when supercooled rapidly enough to avoid crystallization [1–15]. Glass formation is an example of the 'falling-out-of-equilibrium' that takes place for any system the relaxation time of which exceeds laboratory time scales [16]. This phenomenon does not in itself present subtle scientific questions, in our opinion, although there may well be interesting relaxations taking place at T_g affecting details of the glass structure [17]. The ultraviscous liquid in metastable equilibrium above T_g , on the other hand, does present fundamental scientific challenges. The two most important questions relating to the ultraviscous liquid phase preceding glass formation are: (1) What causes the non-exponential relaxations usually observed? (2) What causes the non-Arrhenius temperature dependence of the average (alpha) relaxation time τ ? This paper addresses one of the classical answers to the latter question.

Most viscous liquids require temperature dependence of the activation energy $\Delta E = \Delta E(T)$ if the Arrhenius expression is accepted,

$$\tau(T) = \tau_0 \exp\left(\frac{\Delta E(T)}{k_B T}\right). \tag{1}$$

Molten pure silica and a few other liquids have almost temperature-independent activation energy, but for all other liquids the activation energy increases upon cooling. An unbiased measure of how fast the activation energy increases is the 'temperature index' defined [18] by $I=-d\ln\Delta E/d\ln T\geqslant 0$. The standard measure of the degree of non-Arrhenius behavior is Angell's fragility m defined

by $m=d\log \tau/d(T_g/T)|_{T=T_g}$ [19–21], a quantity that however only refers to liquid properties right at T_g . If the glass transition temperature (by definition) is taken as the temperature where $\tau=100\,\mathrm{s}$ and $\tau_0=10^{-14}\,\mathrm{s}$, Arrhenius behavior corresponds to m=16. In the index terminology Arrhenius behavior corresponds to I=0. Generally, the following relation allows one to calculate the fragility from the index at T_g : $m=16[1+I(T_g)]$ [18].

In the broad research field 'viscous liquids and the glass transition' there is no general agreement about the origin of the non-Arrhenius behavior of viscous liquids. It may well be that no simple, universally valid model or theory exists, but many workers in the field including ourselves prefer to think that such a model exists. This is a reasonable assumption, because ultraviscous liquids approaching the glass transition have physical properties that do not depend on whether the liquid is bonded by covalent bonds, ionic bonds, van der Waals bonds, hydrogen bonds, or metallic bonds [1–15] (we prefer to exclude the often studied polymer glass transition because it is not a liquid–glass transition, but it is noteworthy that this transition has several properties in common with the liquid–glass transition).

Whenever an important scientific problem is unsolved, there is usually not one, but many models allegedly solving the problem. The non-Arrhenius behavior of glass-forming liquids is no exception. Classical phenomenological models relate the relaxation time to other macroscopic liquid properties, like the configurational entropy [22,23], the free volume [24–27], the energy [6,17,28–32], or the high-frequency elastic constants [33–37]. More recently, these were supplemented by models that have more fundamental basis like, e.g., the mode-coupling theory [13,38], the random-first-or-der-transition theory (RFOT) [39,40], energy-landscape based models [41–45], frustration-based approaches [46], the entropic barrier hopping theory [47], kinetically constrained models [48,49], etc.

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This paper deals with one of the most popular classical models, the Adam–Gibbs entropy model [23]. We first briefly review the model and how it was traditionally supported by experiment (Section 2). In Section 3 critiques of the model are presented, relating to both the model's theoretical basis and its experimental validation. Many of these arguments have been made before, but we felt it would be useful to collect them into one paper. Section 4 concludes.

2. The Adam-Gibbs entropy model

2.1. Assumptions and model prediction

According to the Adam–Gibbs model the liquid's relaxation time is controlled by the configurational entropy $S_{\rm c}(T)$. This quantity is defined by subtracting the vibrational entropy, $S_{\rm vib}(T)$, from the entropy S: $S_{\rm c}(T) = S(T) - S_{\rm vib}(T)$. This separation of entropy into two contributions is much in the spirit of the energy landscape paradigm that was subsequently formulated by Goldstein [41] and Stillinger [50], where vibrations around a potential energy minimum (an inherent state) are occasionally interrupted by thermally activated transitions to another minimum.

The Adam-Gibbs model's activation energy obeys

$$\Delta E(T) \propto \frac{1}{S_c(T)}$$
. (2)

This is justified as follows. Any molecular rearrangement is a thermally activated transition that involves all molecules of a 'cooperatively rearranging region.' Such a region is defined as a 'subsystem of the sample which, upon a sufficient fluctuation in energy (or, more correctly, enthalpy), can rearrange into another configuration independently of its environment.' Three crucial ideas/assumptions go into the model: (1) The activation energy is proportional to region volume. This is justified by writing the change in Gibbs free energy upon activation as a chemical potential change $\Delta\mu$ times volume and assuming that 'in a good approximation the dependence of $\Delta\mu$ on temperature and region volume can be neglected. (2) There is a lower limit to the size of a cooperatively rearranging region since it must have at least two configurations 'available to it, one in which the region resides before the transition and another one to which it may move.' (3) The cooperatively rearranging regions are 'independent and equivalent subsystems,' i.e., there are only insignificant interactions of any given region with its surroundings.

2.2. The model's attractive scenario

The Adam-Gibbs model connects two of the most fundamental concepts of physics: Entropy and Time. The model is aesthetically attractive by having this property - the only other quantitative connection of entropy and time that we can think of is that of black hole thermodynamics as theorized by Hawking and others (the fact that entropy cannot decrease for an isolated system is a qualitative entropy-time connection). The entropy model has the further beauty of connecting the observed dramatic slowing down to the Kauzmann paradox and the theory of phase transitions. Recall that the Kauzmann paradox is the observation that the supercooled liquid's excess entropy S_{exc} (the liquid entropy minus the crystal entropy at the same temperature) extrapolates to zero at a temperature T_K not far below T_g [3]. Unless something rather dramatic happens invalidating this extrapolation, the liquid's entropy would fall below the crystal's if the liquid could be equilibrated close to and below T_K . But if – as usually done – the excess entropy is identified with the configurational entropy (a point returned to below),

$$S_{\text{exc}}(T) \cong S_{\text{c}}(T),$$
 (3)

the Adam–Gibbs (AG) model solves the Kauzmann paradox: By Eq. (2) the relaxation time diverges to infinity as the liquid is cooled towards T_K . This means that the liquid cannot equilibrate close to T_K , implying that the glass transition must take place above T_K , no matter how slowly the liquid is cooled.

Based on Eq. (3) the AG model presents a scenario that predicts an underlying phase transition to a state of zero configurational entropy and infinite relaxation time. Thus the model explains the dramatic relaxation-time increase as a consequence of the approach to a phase transition. The predicted slowing down extends over a broader temperature range and is much more dramatic than the usual critical slowing down for second order phase transitions where $\tau \propto |T-T_c|^{-x}$ [51], but the idea is the same. In this way, the paradigm of second order phase transitions comes into play for the glass transition problem.

The above explains the AG model's attraction in general, theoretical terms. Its main attraction, however, is probably the fact that it appears to explain experiments. We shall not detail the evidence for this here, but refer the reader to the several excellent reviews [8,10,11,52]. In many cases the experimental evidence for the AG model relates it to the Vogel–Fulcher–Tammann (VFT) empirical equation for the relaxation time:

$$\tau(T) = \tau_0 \exp\left(\frac{A}{T - T_0}\right). \tag{4}$$

Close to T_K the configurational entropy $S_c(T)$ may be expanded to first order: $S_c(T) \propto T - T_K$, implying that to lowest order the AG model predicts

$$T_K = T_0. (5$$

This prediction has been compared to experiment on many liquids. The general picture reported in numerous papers is that the AG model is obeyed for most, if not all systems studied [52,53]. These include chemically quite different systems with widely differing glass transition temperatures.

3. Critiques of the AG entropy model

3.1. Model assumptions

As mentioned, the three basic assumptions of the AG model are: (1) The activation energy is proportional to the region volume, $\Delta E(T) \propto V_{\rm reg}(T)$. (2) A region must have at least two configurations, i.e., its configurational entropy is larger than $k_B \ln 2$; (3) The 'region assumption' that regions are *independent* and *equivalent* subsystems of the liquid. None of points (1)–(3) are compelling: Molecular rearrangements take place in almost perfect crystals via diffusing vacancies or interstitials, and in a plastic crystal, for instance, one might well have molecular reorientations happening without either Assumption (1) or (2) being obeyed. This is also an example where Assumption (3) breaks down. Even if Assumption (3) holds, however, it is not necessary that a region must have a minimum configurational entropy in order to allow for transitions; also for a low configurational entropy a region would have many states 'available to it' if differing energies are allowed for.

Assumption (1), which is responsible for the non-Arrhenius behavior and the relaxation time divergence as $T \to T_K$, was justified by the suggestion that the chemical potential difference between initial and transition state (barrier) is region-size independent. The question is how well defined a chemical potential difference is for this situation particularly in view of the small region sizes inferred from experiment that makes it difficult to justify ignoring the interactions with the surroundings, see below.

Finally, returning to the region Assumption (3) we note that it can only be justified if regions are very large. As an analogue, note that even for rather large 'regions,' nucleation theory must take into account interactions with the surrounding liquid in order to arrive at realistic predictions. It is not clear why the same should not be done in the Adam–Gibbs theory; indeed, this is done in the more sophisticated RFOT entropy model of Wolynes and coworkers [39,40].

Suppose that we nevertheless accept Assumptions (1)–(3) and go ahead by comparing to experiment. When this is done one typically arrives at regions containing 4–8 molecules [11,54] close to the glass transition. At higher temperatures regions must be even smaller, because it is the increasing regions size upon cooling that is responsible for the non-Arrhenius behavior. The small region sizes of experiment presents a serious challenge to the AG entropy model, because such small regions cannot reasonably be regarded as independent with region-region interactions that may be ignored; every molecule must interact with molecules of other regions as much as with the molecules within its region.

Suppose that one nevertheless accepts the AG idea that the configurational entropy controls the relaxation time's temperature dependence and also accepts Eq. (3) that allows for the entropy model to be tested in experiment. Then, as mentioned, the relaxation time becomes infinite at T_K where the equilibrium state of the liquid has zero configurational entropy - the 'ideal glass' state [55,56]. This state cannot be reached experimentally, of course, but one may still ask what is its nature. A state of zero entropy is unique like a perfect crystal, so one would expect that some simple description of it can be given. Except for the random close packing of hard spheres (the uniqueness of which is questioned), we are not aware of attempts to describe the ideal glassy state in structural terms. This does not rule out that such a description exists, but one would imagine it to be fairly simple (like a quasi-crystal) and thus to have been identified long ago.

3.2. The AG entropy model's experimental validation

Despite the above arguments, suppose that we accept the AG entropy model's prediction Eq. (2). Unfortunately, configurational entropy cannot be measured. For many years this problem was solved by arguing as follows: "The vibrational properties of glass and crystal are very similar, and very similar to the liquid's high-frequency vibrational properties (i.e., on time scales much shorter than those of the alpha (main) relaxation time). Since the crystal-line state has practically zero configurational entropy, the crystal entropy provides a good estimate of the liquid's vibrational entropy. Thus by subtracting crystal entropy from liquid entropy one finds the liquid's configurational entropy (Eq. (3))".

There is now a growing recognition that the above reasoning is problematic [15,57–59]. Dating back to the 1950s, in fact, it was known from sound velocity measurements that the liquid's highfrequency sound velocity is generally much more temperature dependent than that of the crystal or glass phases [5,60,61]. It is easy to understand why this is so if one adopts the simple-minded assumption that the high-frequency sound velocity is a function of density: The thermal expansion coefficient is generally considerably larger in the liquid than in the solid phases (crystal or glass). In this simple approach, the vibrational entropy is a (logarithmic) function of the vibrational force constants that determine the high-frequency sound velocity, so the vibrational entropy is considerably more temperature dependent in the liquid than in the crystal. This severely weakens Eq. (3). An illustration of the problem with Eq. (3) is the fact that it is not generally true that a liquid must have larger entropy than the same temperature crystal: Both in the cases involving so-called inverse melting [62,63] and for the classical hard sphere system, the crystalline phase has larger entropy than the liquid.

Suppose that we nevertheless accept Eq. (3). Then at the Kauzmann temperature T_K there is a second order phase transition to the ideal glassy state – if the liquid has the infinite time needed to equilibrate. But T_K is identified by extrapolation, and one may well question how reliable the extrapolation is. This question arises, in particular, if one accepts that T_g is close to a genuine phase transition as predicted by the AG model. It seems quite possible that the liquid entropy may 'bend over' and stay above the crystalline entropy right down to zero temperature [64–66]. This would imply $T_K = 0$.

Suppose that we nevertheless accept that data conform to Eq. (5) – the intriguing connection of a purely dynamic temperature (T_0) with a purely thermodynamic one (T_K) . Very recently the VFT equation's predicted divergence was questioned in a paper that compiled accurate data for the dielectric relaxation time's temperature dependence for 42 organic liquids [67]. The conclusion was that, while the VFT equation does work well as a mathematically simple representation of data, there is no evidence for any dynamic divergence; in other words, there is no evidence that T_0 exists [68].

Suppose that we nevertheless accept both the extrapolation usually carried out in order to identify T_K and the existence of the VFT T_0 . Then a simple experimental test of the entropy model is the prediction Eq. (5). Numerous papers published the last 30 years have reported confirmation of Eq. (5); indeed this appears to be one of the strongest experimental arguments for the AG entropy model. In 2003, however, Tanaka compiled a large amount of data and concluded that Eq. (5) is disobeyed [69].

4. Concluding remarks

The classical Adam–Gibbs scenario presents several challenges. Thus if entropy is the variable controlling the relaxation time, it seems that more advanced approaches are needed. There are, however, alternatives like the elastic models that date back to the 1940s [15]. According to the shoving model [15,33], one of the elastic models, the activation energy is proportional to the instantaneous shear modulus G_{∞} . This quantity is quite temperature dependent in viscous liquids, in fact precisely enough to explain the non-Arrhenius behavior [70]. Since G_{∞} cannot diverge, there is no underlying phase transition, so the elastic model scenario differs qualitatively from that of the AG entropy model.

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