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Effects of cooling rate on structural relaxation in amorphous drugs: elastically collective nonlinear langevin equation theory and machine learning study

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Theoretical approaches are formulated to investigate the molecular mobility under various cooling rates of amorphous drugs. We describe the structural relaxation of a tagged molecule as a coupled process of cage-scale dynamics and collective molecular rearrangement beyond the first coordination shell. The coupling between local and non-local dynamics behaves distinctly in different substances. Theoretical calculations for the structural relaxation time, glass transition temperature, and dynamic fragility are carried out over twenty-two amorphous drugs and polymers. Numerical results have a quantitatively good accordance with experimental data and the extracted physical quantities using the Vogel–Fulcher–Tammann fit function and machine learning. The machine learning method reveals the linear relation between the glass transition temperature and the melting point, which is a key factor for pharmaceutical solubility. Our predictive approaches are reliable tools for developing drug formulations.

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1. Introduction

Amorphous drugs have attracted much attention^{1–5} owing to their large solubility and enhanced bioavailability compared to the crystalline counterparts. The disordered structure of amorphous pharmaceutical products is formed by the rapid cooling of the molten material. The molecular mobility of an amorphous material is characterized by structural (alpha) relaxation time, τ_α . Since the structural relaxation process originates from liquid structure reorganization, τ_α is temperature-dependent and significantly slowed down at low temperatures. Below the glass transition temperature T_g , which is defined by $\tau_\alpha(T_g) = 100$ s, the drug stays in a disordered state for a long time, which is larger than the experimental observation time scale. However, the pharmaceutical can possibly be recrystallized during manufacturing or storage processes.^{1,2} It turns out that the physical stability of many amorphous systems is relatively poor.

Comprehensive understanding of glassy states and the molecular mobility of amorphous drugs is crucial to formulate pharmaceutical products having desired properties^{1,2} and understand the fundamentals of glassy state physics.

The relaxation processes of amorphous materials can be experimentally measured using broadband dielectric spectroscopy (BDS) and differential scanning calorimetry (DSC).^{1,2} BDS technique determines the structural relaxation time corresponding to thermal variation. DSC can measure the glass transition temperature and analyze phase separation in experimental samples at different cooling rates. The relevant time-scale of molecular motions measured by BDS spans from picosecond above melting temperature to hundreds seconds in vicinity of the glass transition temperature. This technique can be used to investigate both structural (primary or long-time) and transient (secondary or short-time) relaxation processes.

Recently, the Elastically Collective Nonlinear Langevin Equation (ECNLE) theory has been developed to understand structural relaxation time of amorphous systems,^{6–11} in which an amorphous material is modeled as a fluid of molecular particles. The ECNLE theory considers a single molecular motion affected by the nearest-neighbor interactions and cooperative motions of molecules outside a particle cage formed by the neighboring molecules.^{6–10} This physical picture leads to a local barrier of a dynamic free energy and a collective barrier for each molecule caused by its nearest-neighbor interactions and effects of cooperative molecular rearrangements, respectively. These barriers are density-dependent and mutually correlated.

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Plugging the two barriers calculated using the ECNLE theory into Kramer's theory gives the structural relaxation time.^{6–11} To find the temperature dependence of alpha relaxation time, one proposed an analytical conversion (thermal mapping) from averaged particle density to temperature based on experimental dimensionless compressibility data associated with equation of state.^{6–10} The theoretical calculations have also provided quantitative good predictions for the glass transition temperature and dynamic fragility of colloidal systems,^{8,9} supercooled liquids,^{7,9} and polymer melts.^{6,10} However, the experimental equation-of-state data needed for the original thermal mapping has rarely measured in amorphous drugs. Thus, we have recently proposed another density-to-temperature conversion using the thermal expansion and experimental glass transition temperature values.¹¹ Our approach has successfully described temperature-dependent molecular dynamics in one- and multi-component amorphous drugs.

Most ECNLE calculations have assumed a universal correlation between local and collective barrier for all substances when inserting into Kramer's theory. The assumption simplifies roles of chemical and biological complexities on the glass transition. Consequently, the quantitative agreement between theoretical calculations and experiments is imperfect. Moreover, since the dynamic fragility of a material depends strongly on the form of $\tau_\alpha(T)$, a small theory-experiment deviation in $\tau_\alpha(T)$ leads to an inaccurate prediction of the fragility.

In this work, we introduce an adjustable parameter characterizing for a non-universal local-collective correlation in pharmaceuticals. The new version of the ECNLE theory accurately and simultaneously predicts the glass transition temperature and dynamic fragility of amorphous drugs. We employ machine learning technique to reveal a undiscovered relation between melting point and glass transition. We also predict the glass transition temperature based on BDS data. Our ECNLE numerical results are quantitatively compared to experimental data and machine learning calculations.

Theoretical understanding of how glassy dynamics of amorphous materials varies with the cooling rate is limited. Previous theoretical studies^{12–14} have given phenomenological/qualitative descriptions for the T_g change as a function of cooling rate. Based on the theoretical development of this paper, we propose, for the first time, an analysis to provide quantitative determinations for this phenomenon and insightful discussions for experiments.

II. Structural relaxation time of amorphous drugs

Amorphous drugs/materials are theoretically described as a fluid of disconnected spherical rigid particles, which interact with each other *via* hard-sphere interaction. The particle diameter is d and the number density of particles is ρ . A particle is assumingly acted by three forces including (1) the caging force caused by the surrounding fluid, $-\partial F_{\text{dyn}}(r)/\partial r$, (2) the thermal white noise, δf , (3) the friction force, $-\zeta_s(\partial r/\partial t)$, here ζ_s is a short-time friction constant and $r \equiv r(t)$ is the displacement

of the particle. The key quantity is $F_{\text{dyn}}(r)$ known as the dynamic free energy of the tagged particle due to its nearest neighbors.^{6,7,15,16} Based on an overdamped equation-of-motion for the scalar displacement of a tagged particle, we have

$$-\zeta_s \frac{\partial r}{\partial t} - \frac{\partial F_{\text{dyn}}(r)}{\partial r} + \delta f = 0. \quad (1)$$

The thermal noise force satisfies $\langle \delta f(0) \delta f(t) \rangle = 2k_B T \zeta_s \delta(t)$, where k_B is the Boltzmann constant and T is temperature. According to ECNLE theory, the dynamic free energy is^{6,7,15,16}

$$\frac{F_{\text{dyn}}(r)}{k_B T} = \int_0^\infty dq \frac{q^2 d^3 [S(q) - 1]^2}{12\pi\Phi[1 + S(q)]} \exp\left[-\frac{q^2 r^2 (S(q) + 1)}{6S(q)}\right] - 3 \ln \frac{r}{d}, \quad (2)$$

where $\Phi = \rho\pi d^3/6$ is the volume fraction, q is the wavevector, and the static structure factor $S(q)$ is calculated using Percus-Yevick (PY) integral equation theory¹⁷ for a hard-sphere fluid. The PY theory expresses $S(q)$ *via* the direct correlation function $C(q) = [S(q) - 1]/\rho S(q)$. While the Fourier transform of $C(q)$ or the real-space direct correlation function is¹⁷

$$C(r) = -\frac{(1 + 2\Phi)^2}{(1 - \Phi)^4} + \frac{6\Phi(1 + \Phi/2)^2}{(1 - \Phi)^4} \frac{r}{d} - \frac{\Phi(1 + 2\Phi)^2}{2(1 - \Phi)^4} \left(\frac{r}{d}\right)^3 \quad \text{for } r \leq d \quad (3)$$

$$C(r) = 0 \text{ for } r > d. \quad (4)$$

Recall that the ECNLE theory ignores effects of rotational motions and only consider translational motions, which are angularly-averaged. The first term of eqn (2), which depends strongly on the fluid structure and density, is the dynamic trapping potential and favors the particle localization. While the second term independent of the system structure represents the ideal fluid state.

Fig. 1 shows an illustration of the dynamic free energy as a function of r and key physical quantities in the local caging constraint. In dilute suspension ($\Phi \leq 0.43$),^{15–17} $F_{\text{dyn}}(r)$ monotonically decreases with an increase of r and particles move without constraint. When $\Phi > 0.43$, the tagged particle is dynamically arrested within a particle cage formed by its neighbors and one observes an emergence of a free-energy barrier. The particle cage radius, r_{cage} , as depicted in Fig. 1 is determined as the first minimum position in the radial distribution function, $g(r)$. Since $S(q)$ and $g(r)$ are a Fourier-transform pair, one has $g(r) - 1 = \frac{1}{2\pi^2 \rho r} \int_0^\infty [S(q) - 1] q \sin(qr) dq$. Thus, the radius of the particle cage is about $1.3 - 1.5d$. When the dynamic free energy reaches the local minimum and maximum, one obtains the localization length (r_L) and the barrier position (r_B). The energy difference between these two positions is the local hopping barrier $F_B = F_{\text{dyn}}(r_B) - F_{\text{dyn}}(r_L)$. The jump distance from the localized position to the barrier position is defined as $\Delta r = r_B - r_L$. $K_0 = |\partial^2 F_{\text{dyn}}(r)/\partial r^2|_{r=r_L}$ and $K_B = |\partial^2 F_{\text{dyn}}(r)/\partial r^2|_{r=r_B}$



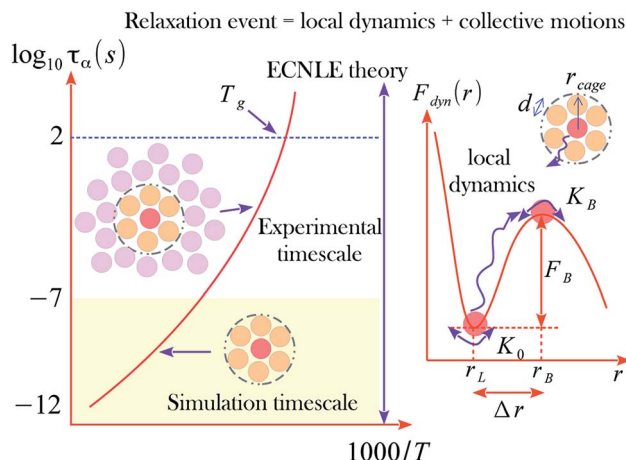


Fig. 1 A general form of temperature dependence of structural relaxation time described by a coupled process of local cage-scale dynamics and collective motions. The dynamic free energy profile of a tagged particle indicates key length scales and a barrier in the local dynamics.

are absolute curvatures at the localization length and barrier position.

The rearrangement of particles in the first shell causes a small expansion on the surface of the particle cage and generates a harmonic displacement field $u(r)$ in surrounding medium *via* collective motions of other particles. In bulk systems, one can obtain analytical form of the distortion field by Lifshitz's continuum mechanics analysis¹⁸

$$\left(K + \frac{G}{3}\right) \nabla(\nabla \cdot \mathbf{u}) + G \nabla^2 \mathbf{u} = 0, \quad (5)$$

where K and G are the bulk and shear modulus, respectively. In ref. 7 and 8, the cage expansion amplitude Δr_{eff} is found to be

$$\Delta r_{\text{eff}} = \frac{3}{r_{\text{cage}}^3} \left[\frac{r_{\text{cage}}^2 \Delta r^2}{32} - \frac{r_{\text{cage}} \Delta r^3}{192} + \frac{\Delta r^4}{3072} \right]. \quad (6)$$

Solving eqn (5) with the boundary condition at the cage surface gives

$$u(r) = \frac{\Delta r_{\text{eff}} r_{\text{cage}}^2}{r^2}, \quad r \geq r_{\text{cage}}. \quad (7)$$

The spatial harmonic displacement energy of a particle at separation distance r from the center of its arrested cage is $K_0 u^2(r)/2$. Since the local time-averaged density is $\rho g(r)$, the number of particles found at a distance between r and $r + dr$ is $\rho g(r) 4\pi r^2 dr$. One can integrate the elastic energies of particles outside the cage to quantify effects of cooperative motions. The collective elastic barrier, F_e , is

$$F_e = 4\pi\rho \int_{r_{\text{cage}}}^{\infty} dr r^2 g(r) K_0 \frac{u^2(r)}{2}. \quad (8)$$

For $r \geq r_{\text{cage}}$, one can approximate $g(r) \approx 1$. The collective motions of molecules play a more important role than the local dynamics in the glass transition at high densities or low temperatures^{6–8} as depicted in Fig. 1.

The activated relaxation is governed by both local and non-local processes. If a universal correlation between a local and non-local process is assumed, the total barrier is simply $F_{\text{total}} = F_B + F_e$. The alpha relaxation time for a particle to diffuse from its particle cage is quantified by Kramer's theory:

$$\frac{\tau_\alpha}{\tau_s} = 1 + \frac{2\pi}{\sqrt{K_0 K_B}} \frac{k_B T}{d^2} e^{F_{\text{total}}/k_B T}, \quad (9)$$

where τ_s is a short relaxation time scale. The mathematical form of τ_s is given by^{7,8}

$$\tau_s = g^2(d) \tau_E \left[1 + \frac{1}{36\pi\Phi} \int_0^\infty dq \frac{q^2 (S(q) - 1)^2}{S(q) + b(q)} \right], \quad (10)$$

where τ_E is the Enskog time scale, $b(q) = 1/[1 - j_0(q) + 2j_2(q)]$, and $j_n(x)$ indicates the spherical Bessel function of order n . Based on many previous studies of thermal liquids, polymers and amorphous drugs,^{6–8,11} one can assume $\tau_E = 10^{-13}$ s.

To convert our hard-sphere calculations into the temperature dependence of the structural relaxation time, we proposed¹¹ a thermal mapping, which is based on the thermal expansion process during temperature variation, to convert from a volume fraction of hard-sphere fluid to temperature of experimental material. The mapping is

$$T \approx T_0 - \frac{\Phi - \Phi_0}{\beta\Phi_0}. \quad (11)$$

where β is the volume thermal expansion coefficient, Φ_0 and T_0 are the characteristic volume fraction and temperature, respectively. Since a typical value for linear thermal expansion coefficient of many glass-forming liquids is $2\text{--}5 \times 10^{-4} \text{ K}^{-1}$,^{19,20} the volume thermal expansion coefficient β is approximately $6\text{--}15 \times 10^{-4} \text{ K}^{-1}$. From a recent work,¹¹ we estimated $\Phi_0 = 0.50$ and $\beta\Phi_0 = 6 \times 10^{-4} \text{ K}^{-1}$.

The parameter T_0 depends on molar mass and particle size. Our numerical calculations indicate the structural relaxation time $\tau_\alpha \approx 100$ s at $\Phi \approx 0.611$. Thus, one can approximately obtain $T_0 = T_g - (0.611 - \Phi_0)/\beta\Phi_0$. The experimental value of T_g can be found in many literatures. Based on the calculation, we investigated the temperature dependence of the structural relaxation time of many amorphous drugs and their mixtures (binary and ternary composites).¹¹ The theoretical calculations without any adjustable parameters quantitatively agree with various experimental data over 14 decades in time. While simulations can determine relaxation times only over first 3–6 decades and do not access experimental timescale as illustrated in Fig. 1.

Fig. 2 shows $\log_{10} \tau_\alpha$ of five representative pure amorphous drugs^{21–24} as a function of $1000/T$ calculated using eqn (9)–(11). Overall, the theoretical curves are relatively close to the experimental counterpart, except for calculations of vitamin A. A deviation of experimental data of the vitamin-A drug from theoretical calculations is expected. This is because the theory ignores many chemical and structural complexities such as



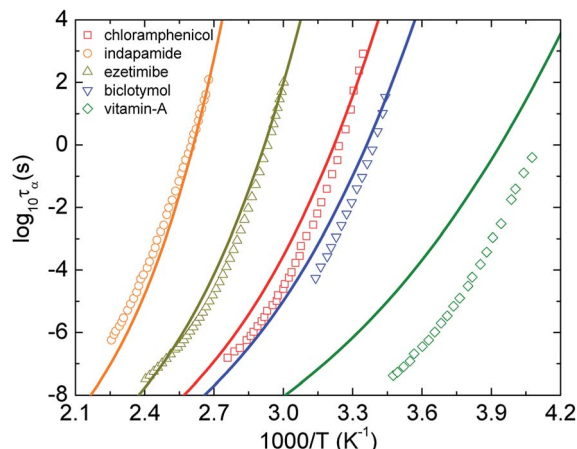


Fig. 2 The temperature dependence of structural relaxation time of chloramphenicol,²¹ indapamide,²² ezetimibe,²² bicalutamide,²³ and vitamin-A acetate.²⁴ Open points are experimental data in literatures and solid curves correspond to our ECNLE calculations.

hydrogen-bonding, network formers, and flexible molecular docking. Moreover, the approach seems to provide less quantitatively accurate predictions for the dynamic fragility of amorphous materials.

III. Nonuniversal coupling of local and cooperative dynamics

The dynamic fragility is calculated by

$$m = \left. \frac{\partial \log_{10}(\tau_s)}{\partial (T_g/T)} \right|_{T=T_g} \quad (12)$$

One adopts the physical quantity to classify into two main categories: “strong” or “fragile” for glass-forming materials. For $m \leq 30$, the glass formers are strong. The glass formers having $m \geq 100$ are fragile. The remaining materials are called intermediate glass-forming materials.

The dynamic fragility is very sensitive to the slope of $\tau_s(T)$ at $T = T_g$. Thus, a good agreement between theory and experiment in τ_s versus T does not mean that another consistency in calculations of m occurs. In ref. 10, authors introduced an adjustable parameter a_c to scale the collective elastic barrier as $F_e \rightarrow a_c^2 F_e$. The parameter a_c captures chemical and biological complexities, conformational configuration, and chain connectivity in different thermal liquids and polymers. The parameter assesses the relative importance of the collective elastic distortion by assuming a non-universal coupling of the cage-scale hopping and collective rearrangements of fluid particles. The previous work¹⁰ obtained contemporaneously the quantitative accordance between theoretical ECNLE calculations and experimental values of both dynamic fragility and glass transition temperature for 17 polymers.

Motivated by the idea in ref. 10, in our calculations, we adjust parameters T_0 and a_c to achieve the best fit to the experimental

temperature dependence of structural relaxation times. Fig. 3 shows experimental data and our theoretical calculations for $\tau_s(T)$ of 22 pure amorphous materials. We carry out the same procedure as calculations in Fig. 2 except for now $F_{\text{total}} = F_B + a_c^2 F_e$. Our numerical results agree quantitatively well with a wide range of experimental data. Remarkably, the activated events of carvedilol, celecoxib, chloramphenicol, and polystyrene below T_g , where τ_s ranges from 100 s to 10^4 s, are well-described using the ECNLE theory. Many previous works^{26,27,43,44} have observed a distinctive deviation, so-called a dynamic structural decoupling, in the relaxation process at low temperatures. For example, in Fig. 3a, the growth of $\tau_s(T)$ of carvedilol drug below T_g is abruptly deviated from what it is supposed to be. Currently, there is poor theoretical understanding for the interesting but challenging feature. In the framework of the ECNLE theory, the decoupling could be related to a temperature dependence of thermal expansion coefficient β in our thermal mapping.

From calculations in Fig. 3, we can estimate the glass transition temperature and dynamic fragility. The local-nonlocal coupling parameter a_c , the characteristic temperature T_0 , the

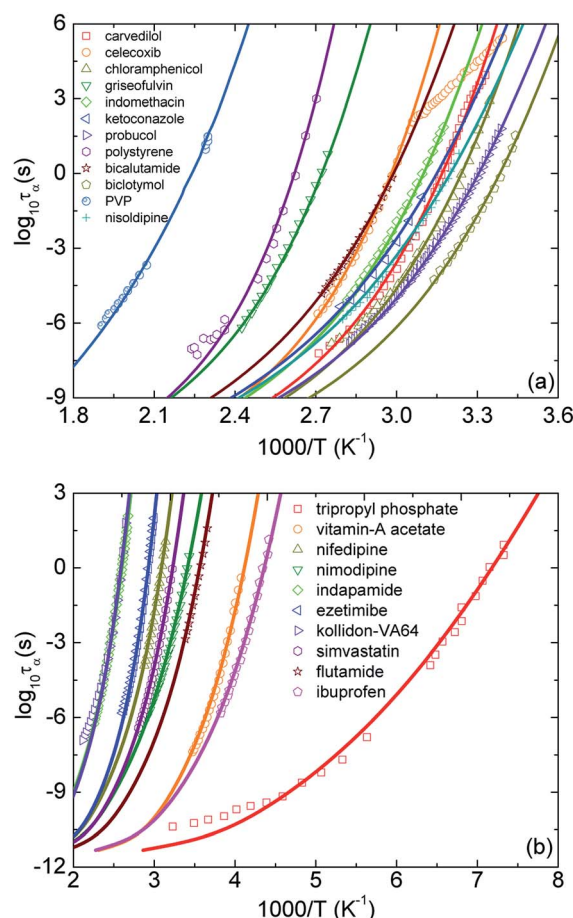


Fig. 3 The temperature dependence of structural relaxation time of twenty-two various amorphous drugs and polymers listed in Table 1.^{11,18–39} Open points are experimental data in literatures and solid curves correspond to our ECNLE calculations. PVP is an abbreviation of polyvinyl pyrrolidone K30.

melting temperature T_m , and theoretical and experimental values for T_g and m of the studied materials are listed in Table 1. Clearly, the theoretical T_g is in perfect accordance with the experimental counterpart. The different accuracy of our calculations for the fragility is somehow expected and reflects a complicated $T_g - m$ correlation.

IV. Machine-learning based analysis of glassy dynamics

There are two main methods to obtain T_g from experiments. First, one can use the DSC measurement for samples to find the value of T_g . Second, experimentalists have widely used the Vogel–Fulcher–Tammann (VFT) equation to fit data measured by the BDS, which is inaccessible to the deeply supercooled state or ($\tau_\alpha \leq 1$ s). From this fitting function, they can obtain the glass transition temperature at $\tau_\alpha = 100$ s *via* extrapolation. However, the fitting depends strongly on a focused regime of data.⁴⁵ The selection process possibly causes large over-prediction or underprediction of T_g . This is possibility why many works have reported different T_g values for a given material.

Here we introduce, for the first time, another approach based on machine learning techniques. After obtaining a predictive model by applying the support vector regression (SVR) in the Scikit-learn Python library⁵⁴ to experimental data of the temperature dependence of τ_α at high temperatures, we can predict new relaxation times at lower temperatures than the coolest temperature in the training dataset. The SVR with the radial basis function (RBF) kernel has two controlled parameters including a regularization parameter C_{RBF} and a RBF kernel parameter γ . While γ is considered as the inverse of the

standard deviation of the RBF kernel, C_{RBF} determines the penalty of large slack variables. In our calculations, we chose $\gamma = 0.1$ and obtained equal performance when $C_{\text{RBF}} \geq 100$. As shown in Fig. 4, $\tau_\alpha(T)$ versus $1000/T$ predicted by the SVR with $C_{\text{RBF}} = 10^3$ grows smoothly and are close to experimental data for some representative amorphous drugs. The number of BDS data points for simvastatin, ketoconazole, bicalutamide, griseofulvin, vitamin-A acetate, and nisoldipine used for training are 24, 8, 17, 17, 22, and 28, respectively. The SVR calculations give T_g s for simvastatin, ketoconazole, bicalutamide,

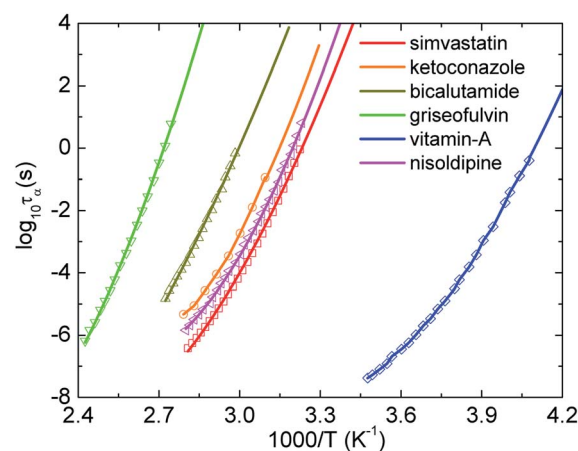


Fig. 4 The temperature dependence of structural relaxation time of simvastatin¹¹, ketoconazole,³⁰ bicalutamide,³³ griseofulvin²⁸, vitamin-A acetate,²⁴ and nisoldipine.³⁵ Open points are experimental data in literatures and solid curves correspond to machine learning-based calculations.

Table 1 System parameters and theoretical and experimental results

Materials	$T_g(\text{th})$ (K)	$T_g(\text{expt})$ (K)	$m(\text{th})$	$m(\text{expt})$	$T_m(\text{expt})$ (K)	a_c	T_0 (K)
Carvedilol ²⁵	308	310 (ref. 25)	91.5		387.5 (ref. 38)	2.1	450
Celecoxib ^{26,27}	328	328 (ref. 26)	97.8	110 (ref. 26)	432 (ref. 38)	2.1	470
Chloramphenicol ²¹	301.1	301 (ref. 21)	89	116 (ref. 21)	423.5 (ref. 38)	2.1	443
Griseofulvin ²⁸	358	359 (ref. 28)	88.7	84.6 (ref. 28)	489 (ref. 39)	1.2	533
Indomethacin ²⁹	314.1	315 (ref. 29)	86.4	77, 67, 64 (ref. 29)	432 (ref. 39)	1.5	476
Ketoconazole ³⁰	308	316.3 (ref. 30)	71.37		419 (ref. 38)	1.0	493
Probuco ³¹	294	294.7 (ref. 31)	79.4	85 (ref. 31)	398 (ref. 38)	1.5	456
Polystyrene ³²	373	373 (ref. 10)	105.53	116, 143, 97, 121 (ref. 10)	513 (ref. 40)	1.7	528
Bicalutamide ³³	325.3	325.4 (ref. 33)	78.03	84 (ref. 33)	464 (ref. 39)	1.1	505
Biclotymol ²³	288	288 (ref. 23)	85.42	85 (ref. 23)		2.1	430
Polyvinylpyrrolidone K30 (ref. 26)	431.2	431 (ref. 26)	58.47	70 (ref. 26)	523 (ref. 41)	0.3	672
Tripropyl phosphate ³⁴	132.1	134 (ref. 34)	40.3		194 (ref. 38)	2.4	266
Vitamin-A acetate ²⁴	236.3	244.3 (ref. 24)	76.6	83 (ref. 24)	330 (ref. 42)	3.4	349
Nisoldipine ³⁵	303	305 (ref. 35)	70.2	81 (ref. 35)	425 (ref. 42)	1.0	488
Nifedipine ³⁵	315	315 (ref. 35)	74.9	84 (ref. 35)	444 (ref. 39)	1.1	494
Nimodipine ³⁵	284.1	285 (ref. 35)	62	82 (ref. 35)	398 (ref. 38)	0.9	474
Indapamide ²²	373.4	373.5 (ref. 22)	75.3	73 (ref. 22)	433 (ref. 38)	0.7	578
Ezetimibe ²²	333.3	333.1 (ref. 22)	91.1	93 (ref. 22)	436 (ref. 38)	1.5	495
Kollidon VA64 (ref. 11)	376.2	378 (ref. 11)	79			0.8	573
Simvastatin ¹¹	301	303 (ref. 11)	74.8	73	411 (ref. 38)	1.3	471
Flutamide ³⁶	272	271 (ref. 36)	71.6		385 (ref. 38)	1.4	438
Ibuprofen ³⁷	222	225 (ref. 37)	68	87 (ref. 37)	346 (ref. 39)	2.4	356



griseofulvin, vitamin-A acetate, and nisoldipine are 300.2, 308.64, 323, 358.2, 237.8, and 304 K, respectively. These values quantitatively agree experimental data and our ECNLE results in Table 1. This is a new reliable approach for investigating molecular dynamics of amorphous materials near T_g , particularly when the structural decoupling appears.^{26,27,33,43,44} The linear regression cannot be used in this protocol since it enforces a Arrhenius nature on any experimental data and keeps it unchanged in the predictive process.

To find new minimalist correlations or physical insights among the quantities in Table 1, we employ a linear regression model in the scikit-learn library.⁵⁴ This regression algorithm provides the simplest relation to describe a target variable from a set of descriptor variables. By adopting T_g and T_m experimental values of 71 glassy drugs listed in ref. 46 as a training dataset, one obtains $T_m \approx 1.362 T_g$. It is important to note that these drugs are different from amorphous materials in this work. Then, we apply the predicted relation to our twenty substances to evaluate the validity of the model. Numerical results in Fig. 5 indicate that the $T_g - T_m$ correlation works well. This finding suggests the ECNLE theory can be exploited to estimate the melting temperature with a reasonable deviation.

In many prior works,^{47–49} the melting point of amorphous drugs exhibits an essential role in the solubility determination of the drugs. The linear relation between T_g and T_m suggests that one can employ ECNLE theory to evaluate the solubility of amorphous materials and their mixtures.

V. Effects of cooling rate on glassy dynamics

Although it is experimentally well-known that cooling rate has considerable impact on glassy dynamics, but theoretical understanding has remained ambiguous. Thus, in this section, we would propose a simple model to estimate how T_g is varied with different cooling rates.

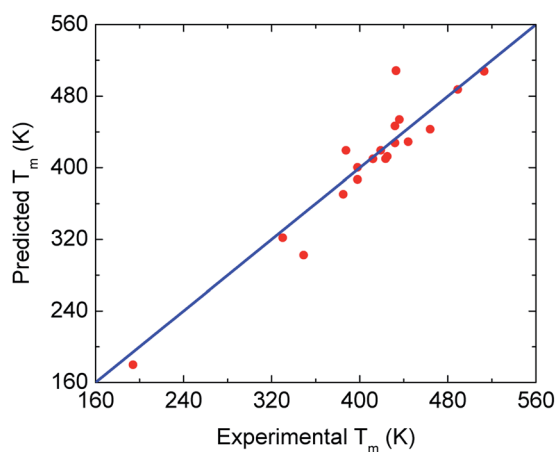


Fig. 5 Cross-plot of predicted and experimental values of the melting temperature for all 20 drugs and polymers from Table 1. The blue line indicates perfect agreement.

According to an assumption introduced by Cooper and Gupta in 1982,¹² the molecular relaxation time is approximately equal to the experimental observation time at T_g . This assumption leads to a new definition of a cooling rate, h ,

$$h = -\frac{dT}{dt} \approx -\frac{dT}{d\tau_\alpha}. \quad (13)$$

The minus sign in eqn (13) represents an inverse variation between mobility and temperature.

Near glass transition temperature, $e^{F_{\text{total}}/k_B T} \gg 1$ and the alpha structural relaxation time in eqn (9) approximately becomes

$$\tau_\alpha(T) \approx \frac{2\pi\tau_s}{\sqrt{K_0 K_B}} \frac{k_B T}{d^2} e^{F_{\text{total}}/k_B T}. \quad (14)$$

Recall that the total barrier here is $F_{\text{total}} = F_B + a_c^2 F_c$ as discussed in Section III. After straightforward transformations, one obtains

$$\frac{1}{\tau_\alpha} \frac{d\tau_\alpha}{dT} = \frac{d}{dT} \ln \left(\frac{2\pi\tau_s}{\sqrt{K_0 K_B}} \frac{k_B T}{d^2} \right) + \frac{d}{dT} \left(\frac{F_{\text{total}}}{k_B T} \right). \quad (15)$$

Since τ_s is order of picoseconds (10^{-12} s), the first term is much smaller than the second term near T_g . Thus,

$$h\tau_\alpha(T_g) \frac{d}{dT} \left(\frac{F_{\text{total}}}{k_B T} \right) \bigg|_{T=T_g} \approx -1. \quad (16)$$

When the temperature dependence of structural relaxation time obeys the Arrhenius behavior, the total barrier F_{total} is a constant and our eqn (16) can be deduced to be the same mathematical form as previous studies^{13,14}

$$h\tau_\alpha(T_g) \frac{F_{\text{total}}}{k_B T_g^2} = 1 \quad \text{or} \quad \frac{h\tau_\alpha(T_g)}{T_g^2} = \text{constant}. \quad (17)$$

Moreover, combining eqn (14) and (16) gives

$$\ln \left(-\frac{d}{dT} \left(\frac{F_{\text{total}}}{k_B T} \right) \bigg|_{T=T_g} \right) + \ln \left(\frac{2\pi\tau_s}{\sqrt{K_0 K_B}} \frac{k_B T_g}{d^2} \right) + \frac{F_{\text{total}}}{k_B T_g} + \ln h = 0. \quad (18)$$

The above equation reveals explicitly a correlation between the cooling rate and the glass transition temperature. Recall that various experimental studies have empirically indicated that $\ln h$ is linearly proportional to $-1/T_g$ in a specific range of T_g . If the relation is universal, it suggests the first two terms in eqn (18) may play a minor role compared others and then our analysis can clearly explain the experimental observation.

Fig. 6 shows how the cooling rate influences the glass transition temperature of polystyrene,⁵⁰ PVP,⁵¹ nifedipine,⁵² and indomethacin.⁵¹ Our theoretical calculations are relatively consistent with experimental data. Although both theoretical curves and data points are not perfect straight lines, the rough



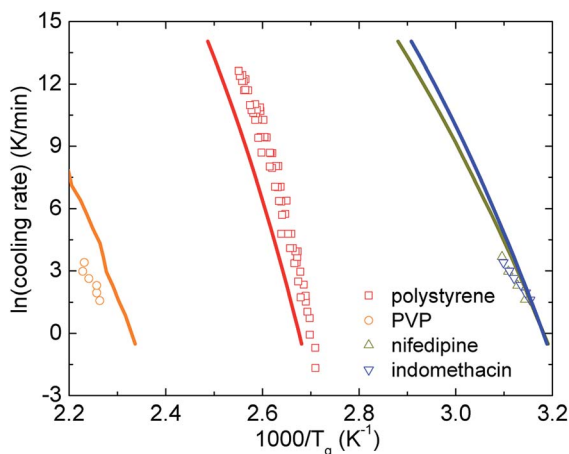


Fig. 6 The logarithm of cooling rate as a function of inverse glass transition temperatures of various amorphous drugs and polymers. Points are experimental data and solid curves correspond to our theoretical calculations using eqn (16).

linearity is possibly observed in a certain range of the glass transition temperature. One can also crudely view these smooth curves as a combination of high- and low- T_g linear branches. Furthermore, as discussed in Section IV, $T_m \approx 1.362T_g$, we expect plotting $\ln h$ versus $1000/T_m$ (not shown here) does not change the variation trends compared to Fig. 6. The finding is in accordance with other experimental works.⁵³

The effects of cooling rate on T_g can be also analyzed using the VFT-type relaxation time, which is

$$\tau_\alpha(T) = \tau_0 \exp\left(\frac{DT_{\text{VFT}}}{T - T_{\text{VFT}}}\right), \quad (19)$$

where τ_0 , D , and T_{VFT} are parameters fitted from experimental data. At $T = T_g$, we have

$$h\tau_\alpha(T_g) = \frac{(T_g - T_{\text{VFT}})^2}{DT_{\text{VFT}}}. \quad (20)$$

Interestingly, both ECNLE theory and VFT-function analysis give us the same form of the nontrivial correlation among the cooling rate, glass transition temperature, and fragility

$$h\tau_\alpha(T_g) = \frac{T_g}{m \ln(10)}. \quad (21)$$

For small $\ln q$ (≤ 0) or low cooling rates, T_g of polystyrene is nearly unchanged, approximately 373 K. Thus, one can utilize the definition of $\tau_\alpha(T_g) = 100$ s to estimate T_g in the range of the cooling rate. It suggests that if τ_α of all substances is measured at the same low cooling rate, $h\tau_\alpha(T_g) \approx 100h$ and T_g is linearly proportional to m . However, since different experiments have been carried out at different cooling rates, it is hard to obtain a $T_g - m$ trivial correlation.

VI. Conclusions

We have shown several theoretical approaches to improve quantitatively accurate predictions of the glass transition temperature and dynamic fragility of twenty-two amorphous drugs and polymers. The temperature dependence of the structural relaxation time is theoretically calculated using the version of ECNLE theory. By introducing an adjustable parameter to describe a non-universal correlation between local and collective molecular dynamics in different materials, our numerical results for the dynamic fragility and T_g measured at various cooling rates show better quantitative agreement with experiments than simply using the universal local-nonlocal coupling. Applying machine-learning calculations to BDS experimental data gives the same T_g values as using the VFT fit function. The finding suggests that machine learning technique can verify the VFT-based results in all BDS studies instead of comparing with DSC experiments. The machine-learning calculation may be more reliable to predict T_g and m when the structural decoupling of relaxation process occurs. Machine learning also reveals the linear relation of $T_m \approx 1.362T_g$. This relation explains why the cooling rate changes the melting point and glass transition temperature in the same manner.

Conflicts of interest

There are no conflicts to declare.

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