Predicting Dynamic Heterogeneity in Glass-Forming Liquids by Physics-Inspired Machine Learning

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We introduce GlassMLP, a machine learning framework using physics-inspired structural input to predict the long-time dynamics in deeply supercooled liquids. We apply this deep neural network to atomistic models in 2D and 3D. Its performance is better than the state of the art while being more parsimonious in terms of training data and fitting parameters. GlassMLP quantitatively predicts four-point dynamic correlations and the geometry of dynamic heterogeneity. Transferability across system sizes allows us to efficiently probe the temperature evolution of spatial dynamic correlations, revealing a profound change with temperature in the geometry of rearranging regions.

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Glasses are formed by the continuous solidification of supercooled liquids under cooling, while maintaining an amorphous microstructure [1]. Understanding glass formation and the phenomenon of the glass transition has been the focus of an intense research activity [2,3].

An important feature of supercooled liquids is the growth of spatial heterogeneity characterizing the relaxation dynamics, where some regions actively rearrange while others appear completely frozen [4]. Recently, an important effort was devoted to understanding the connection between dynamic heterogeneity and structural properties [5–7]. Several structural parameters were shown to correlate with the dynamics, including density, potential energy [8,9], and locally favored structures [10–12], but also more complicated quantities such as soft modes [13], local yield stress [14], and Franz-Parisi potential [15]. The search intensified with the emergence of machine learning (ML) allowing the detection of correlations from unsupervised [16–19] or supervised [20–26] learning. The explored methodologies range from simple linear regression and support vector machines using a set of handcrafted structural descriptors [20] to graph neural networks (GNN) with tens of thousands of adjustable parameters [22,26]. Despite this versatility, none of the proposed networks can so far predict dynamic heterogeneities and related multipoint correlation functions that quantitatively agree with the actual dynamics.

Here, we bridge this major gap by leveraging and combining previous ML approaches. We construct a physics-inspired deep neural network that uses established structural order parameters as input to predict long-time dynamics in deeply supercooled liquids. The proposed methodology, which surpasses the state of the art, allows us to very efficiently obtain quantitative predictions about heterogeneous dynamics and hence to gather novel physical insights about their temperature evolution.

We simulate a Lennard-Jones nonadditive Kob-Andersen mixture in 3D (KA, [27]) for comparison with earlier work [22] and a 2D ternary mixture (KA2D) where lower temperatures can be accessed. We focus on KA2D since its interactions were adapted to efficiently prevent crystallization [28] and enable the use of the swap Monte Carlo (SWAP) algorithm [29,30]. Equilibrium configurations are created with \( N = 1290 \) particles \( (M_{\text{type}} = 3, N_1 = 600, N_2 = 330, N_3 = 360) \) and box length \( L = 32.896 \) using periodic boundary conditions and reduced units. We use SWAP to equilibrate the system and create a statistical ensemble. The average over equilibrium configurations is denoted \( \langle \cdots \rangle \). For each configuration, \( N_R = 20 \) replicas are created by drawing initial velocities from the Maxwell distribution to analyze the isoconfigurational ensemble [13,31] in which one averages over velocities at fixed initial configuration. We then simulate the dynamics using molecular dynamics (MD) and calculate for each particle \( i \) the isoconfigurational average of the bond-breaking (BB) correlation function \( C_{BB}(t) = \langle n_t^{BB}/n_0^{BB} \rangle_{\text{iso}} \), which following Refs. [13,31] we call “propensity”; \( C_{BB}^i(t) \) describes the number \( n_t^i \) of nearest neighbors particle \( i \) still has after a time \( t \) relative to its \( n_0^i \) initial number of neighbors [32].

From the averaged propensity \( \tilde{C}_B(t) = \langle 1/N_i \sum_{i \in N_i} C_{BB}^i(t) \rangle \), we extract a structural relaxation time, \( \tau_{BB}^i \), defined as \( \langle \tilde{C}_B(t = \tau_{BB}^i) \rangle = 0.5 \). We report results for type 1 but verified that all findings are independent of particle type.

We focus on three different temperatures: (i) slightly below...
the onset temperature \(T = 0.4, \tau_{\alpha}^{BB} = 1.7 \times 10^3\), (ii) slightly above the mode-coupling temperature \(T = 0.3, \tau_{\alpha}^{BB} = 3.4 \times 10^4\), and (iii) slightly below the mode-coupling temperature \(T = 0.23, \tau_{\alpha}^{BB} = 4.0 \times 10^6\). More details are given in the Supplemental Material [33].

The first step in the ML approach is to select physics-inspired inputs: a number \(M_S\) of structural descriptors constructed for each particle \(i\) from \(K\) different observables. Inspired by the handcrafted features in Refs. [24,25] we also calculate coarse-grained averages of these descriptors on different length scales \(L\). The first descriptor is the coarse-grained local density, \(\bar{\rho}_{L,\beta} = \sum_{j \in N_i^\beta} e^{-R_{ij}/L}\), where the sum runs over all \(N_i^\beta\) particles of type \(\beta\) within distance \(R_{ij} = |R_i - R_j| < 20\) of particle \(i\). Particle positions are evaluated in the inherent structures \(R_i\). Similar in philosophy to Ref. [23] we additionally choose three different physics-inspired descriptors: the coarse-grained potential energy, \(E_i^{L,\beta} = \sum_{j \in N_i^\beta} E_i e^{-R_{ij}/L}/\bar{\rho}_{L,\beta}\), extracted from the pair potential \(E_i = \sum_{j \neq i} V(R_{ij})/2\), the local Voronoi perimeter \(P_i^{L,\beta} = \sum_{j \in N_i^\beta} P_i e^{-R_{ij}/L}/\bar{\rho}_{L,\beta}\), using the perimeter \(p_i\) of the Voronoi cell around particle \(i\), extracted using the software Voro++ [39], and finally local variance of potential energy, \(\overline{\Delta E}_{L,\beta} = \sum_{j \in N_i^\beta} (E_i - \bar{E}_i)^2 e^{-R_{ij}/L}/\bar{\rho}_{L,\beta}\).

As coarse-graining lengths we choose \(M_{CG} = 16\) values \(L = \{0.0, 0.5, ..., 7.5\}\). In addition to coarse graining the descriptors separately for each of the \(M_{type}\) types we also calculate the coarse-grained average by iterating over all particles independently of type. In total, this procedure therefore produces a set of \(M_S = KM_{CG}(M_{type} + 1) = 256\) descriptors. To simplify the learning, each descriptor is shifted and rescaled to have zero mean and unit variance over the training set.

We then apply a supervised ML procedure to train a multilayer perceptron (MLP) to give a prediction \(\chi'_{MLP}\) [40] for the propensity of particle \(i\). Between the input and output layers, we introduce three hidden layers with 2, 10, and 10 nodes, respectively, as sketched in Fig. 1. In total, our model has around 650 fitting parameters, about 100 times less than the GNN proposed in Ref. [22], and slightly fewer than the networks used in Refs. [24,25] due to a significant reduction in the number of structural descriptors \(M_S\). The intermediate layer with only two nodes is a bottleneck layer. Its introduction is crucial to prevent overfitting of the training data and represents a major difference from the MLP suggested in Ref. [25] where unsatisfying results were reported. We name our deep neural network “GlassMLP”. We use \(N_S = 300\) initial structures, which are equally divided into training, validation, and test sets. During learning, we compute for each configuration as loss function the mean absolute error between true and predicted labels [22,24,25]. In the loss we also include terms that penalize deviations from the true variance and spatial correlations of the propensities. Both quantities are evaluated by averaging over all particles in the configuration for which the loss function is evaluated. For the training we apply stochastic gradient descent with an Adam optimizer [41]. The hyperparameters used for training are the same for all times and temperatures. The training of GlassMLP on one state point requires less than five minutes on a Laptop GPU (NVIDIA T600 Laptop).

To quantify the performance of GlassMLP we compute the Pearson correlation coefficient \(\rho_p = \frac{\text{cov}(C_B, \chi'_{MLP})}{\sqrt{\text{var}(C_B)\text{var}(\chi'_{MLP})}}\), between the true propensities \(C_B\) and the network output, \(\chi'_{MLP}\). Perfect predictions would yield \(\rho_p = 1\) while random ones correspond to \(\rho_p = 0\). As shown in Fig. 2(a), we find that \(\rho_p\) depends nonmonotonically on time and is maximal around \(t \approx \tau_{BB}^{\alpha}/3\). Furthermore, the predictability considerably increases at lower temperatures and reaches values up to \(\rho_p > 0.8\), which is significantly better than previously proposed techniques on KA models [16,17,22,24,25]. A direct comparison to GNNs [22] is presented below for the 3D KA model.

We now go beyond establishing the quality of a correlation and focus on the probability distribution of the propensity. Figure 2(b) shows an excellent agreement between GlassMLP predictions and MD results. Minor discrepancies exist in the tails for small propensities, as the network slightly underestimates variances. Poor results are instead obtained by the ridge regression method suggested in Refs. [24,25], which always outputs nearly Gaussian distributions. This shows that using a nonlinear neural network such as GlassMLP is important to capture the complex shape of the distributions. See the Supplemental Material [33] for further comparison between methods [33].

Because GlassMLP performs excellently at the single-particle level, we now apply it to spatial correlations, thus
promoting GlassMLP as a new tool to probe dynamic heterogeneity \cite{42}. First, we show snapshots of the predicted and calculated propensities for different timescales in Fig. 3(a). The MD results show how marginally rearranged active clusters at small times (white and red) coarsen with time and become both larger and more strongly contrasted to the unrelaxed background (blue) \cite{43}. GlassMLP is able to predict remarkably well the location and the geometry of the relaxing clusters from the sole knowledge of the initial structure.

Spatially heterogeneous dynamics is quantified by the four-point susceptibility $\chi_4(q,t) = N^{-1}_i(\langle \bar{C}_B(t) \rangle - \langle \bar{C}_B(t) \rangle)^2$ shown in Fig. 3(b). Its time dependence is similar to the one of the Pearson correlation, with a maximum at $t \approx \tau_{BB}/3$ that grows upon cooling. This similarity suggests that GlassMLP is particularly powerful in analyzing strongly heterogeneous dynamics. The effect is further enhanced due to the increased structural origin for dynamic heterogeneities at lower temperatures observed in earlier work \cite{42}. Figure 3(b) also highlights that GlassMLP accurately predicts the time and temperature evolution of $\chi_4(t)$. To our knowledge, no ML technique has previously been able to predict $\chi_4(t)$ at a comparable quantitative level. This susceptibility quantifies the average number of correlated particles during structural relaxation \cite{44} and can be accessed experimentally \cite{45,46}.

The evolution of $\chi_4(t)$ results from two factors \cite{47,48}: a growing length scale characterizing the decay of dynamic correlations, and a growing strength of these correlations. We now show that GlassMLP can even disentangle them. Let us define the four-point structure factor, $S_4(q,t) = N^{-1}_i(W(q,t)W(-q,t))$, with $W(q,t) = \sum_{i\in N}\langle \bar{C}_B(t) - \bar{C}_B(t) \rangle \exp[iq \cdot R_i(t)]$. See the Supplemental Material \cite{33} for the analysis of its real space counterpart. The measured $S_4(q,t)$, shown in Fig. 4(a), displays a peak at small $q$ which contains all relevant information about spatial dynamic correlations. For this function the predictions made by GlassMLP are again in excellent agreement with measurements. It is notoriously difficult to quantitatively extract a correlation length scale $\xi$ from $S_4(q,t)$ as one needs systems much larger than $\xi$ \cite{49,50,51,52}. Previous works tackled this challenge by simulating very large systems which becomes a real challenge at low temperatures where long timescales are also needed. GlassMLP fully solves this problem by transferring results from small to large systems. One can train GlassMLP on reasonably small (but not too small) systems and then apply it to very large ($N = 82,560$) equilibrium configurations obtained using SWAP. GlassMLP predicts the propensity field and hence $S_4(q,t)$ for these configurations at essentially no cost because the network is already trained and the slow dynamics of large systems is never simulated. The transferability in system size is possible because the bond-breaking correlation and $S_4(q,t)$ have been shown to be independent of system size for the chosen $N$ values \cite{52,53}.

See the Supplemental Material \cite{33} for finite-size analysis \cite{33}. This method allows us to obtain for the first time reliable data for $S_4(q,t)$ over an extended range of times, temperatures, and wave vectors; see Fig. 4 \cite{54}. We find that an Ornstein-Zernicke functional form, $S_4 \approx 1/[1 + (q\xi)^2]$ does not describe the numerical data over the entire range of temperature for $q > 0.2$ and a higher-order term is needed. This was proposed theoretically using mode-coupling theory \cite{55} with a quartic term, and in the East model \cite{56} where a fractal exponent $q^{1.58} - 1$ is found. Neither proposal is consistent with our data. Because dynamic heterogeneity appears increasingly contrasted with more compact boundaries at lower temperatures \cite{43}, we introduce a cubic term $q^{3}$ by analogy with Porod’s law describing two-phase systems with sharp interfaces \cite{57}: $S_4(0.2 < q < 0.6, t) = S_4(t)/[1 + (\xi q)^2 + A(\xi q)^3]$. This expression contains the minimal ingredients to describe both the evolution of the characteristic length scale $\xi$.
Isoconfigurational average of particle displacements, by Ref. [22] and similarly define the propensity as the use the same dataset and the pretrained GNNs provided for benchmarking, and to present a fair comparison, we the performance of GlassMLP for a different model. For which the GNN of Ref. [22] was initially applied. The aim is to compare GlassMLP and the GNN and to show

which is consistent with the conclusions in Ref. [62]. The trained networks do not detect any outstanding features, although GlassMLP successfully captures the underlying physics when approaching the glass transition. Improved performance is reached from (i) using prior knowledge about glass transition as inductive bias for neural networks [23]; (ii) including spatial correlations into the loss function; and (iii) adjusting the architecture of the deep neural network to avoid overfitting. Using transferability across system sizes allows one to extract physically meaningful four-point dynamical structure factors and to analyze their physical evolution when approaching the glass transition. Although GlassMLP’s performance is remarkable, the trained networks do not detect any outstanding features, which is consistent with the conclusions in Ref. [62]. The success of GlassMLP therefore demonstrates the importance of combining physics-inspired inputs and deep neural networks able to extract inherent complex and nonlinear features from them, with relative weights that are presumably model dependent.

![Graph of correlation versus temperature](image)

**FIG. 4.** Evolution of length scales and geometry of dynamic heterogeneity in the 2DKA model. (a) Four-point structure factor slightly below the structural relaxation time $t_{\text{sl}}^3/3$ for different temperatures $T$ and system sizes $N$. Dashed lines are fits $S_i (q, t) = \chi_4 (t) / [1 + (\xi q)^2 + A(\xi q)^3]$ as rationalized in the main text. (b) Length scales $\xi$ extracted from nonlinear fits described in the main text. Only points for which the Pearson coefficient $\rho_P > 0.5$ are shown. (c) Rescaled four-point structure factor vs rescaled wave number $q \xi$ for the MLP, $N = 82560$ data. Dashed lines correspond to $[1 + (q \xi)^2]^{-1}$ and dashed-dotted line is $\sim q^{-3}$. Inset shows enlarged data for large $q \xi$. (d) Higher-order prefactor $A$, extracted from fitting $S_i (q, t)$ as described in the main text.

**FIG. 5.** Comparison of two different ML techniques to predict the isoconfigurational average of displacements $R_i (t)$ for the 3D KA model. (a) Pearson correlation coefficient $R$ for different times $t$ at temperature $T = 0.44$. The vertical dotted line marks structural relaxation $t = t_s$ and the dashed-dotted line is the maximal achievable correlation. (b) Susceptibility $\chi_4 (t)$ compared to the ground truth (MD).
The method proposed here could easily be extended to include further descriptors and be applied to other types of systems, including experiments on glass-forming colloidal liquids or granular glasses, where potentially different descriptors can be used. Our findings on spatially correlated dynamics pave the way for a more rigorous analysis of dynamic heterogeneity in deeply supercooled liquids to better understand their physical origin, and the interplay between heterogeneous structure [15] and dynamic facilitation [43] close to the experimental glass transition.

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