Deep Learning for Automated Classification and Characterization of Amorphous Materials

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It is difficult to quantify structure-property relationships and to identify structural features of complex materials. The characterization of amorphous materials is especially challenging because their lack of long-range order makes it difficult to define structural metrics. In this work, we apply deep learning algorithms to accurately classify amorphous materials and characterize their structural features. Specifically, we show that convolutional neural networks and message passing neural networks can classify two-dimensional liquids and liquid-cooled glasses from molecular dynamics simulations with greater than 0.98 AUC, with no a priori assumptions about local particle relationships, even when the liquids and glasses are prepared at the same inherent structure energy. Furthermore, we demonstrate that message passing neural networks surpass convolutional neural networks in this context in both accuracy and interpretability. We extract a clear interpretation of how message passing neural networks evaluate liquid and glass structures by using a self-attention mechanism. Using this interpretation, we derive three novel structural metrics that accurately characterize glass formation. The methods presented here provide us with a procedure to identify important structural features in materials that could be missed by standard techniques and give us a unique insight into how these neural networks process data.

I. INTRODUCTION

Classifying material structures and predicting their properties are important tasks in materials science. The behavior of materials often depends strongly on their underlying structure, and understanding these structure-property relationships relies on accurately describing the structural features of a material. However, quantifying structure-property relationships and identifying structural features in complex materials are difficult tasks.

A variety of standard techniques have been developed to analyze material structures. Some of the most common techniques include the Steinhardt bond order parameters,¹ Bond Angle Analysis (BAA),² and Common Neighbor Analysis (CNA),³ which are useful for detecting order-disorder transitions and differentiating between crystal structures in ordered samples.⁴ Radial distribution functions, which measure a spatial average of interparticle distances, are also widely used for analyzing different materials and phases.⁵

However, these standard techniques have limitations, especially for analyzing weakly crystalline or amorphous materials. As discussed in Reinhardt et al.,⁶ the Steinhardt bond order parameters can be stymied by thermal fluctuations or ambiguous crystal structures. BAA relies on a small set of crystalline reference structures that may not be present in amorphous samples. CNA is more flexible than BAA, but it cannot provide accurate information about particles that do not exhibit known symmetries, making analysis of irregular structures challenging. Radial distribution functions rely on spatial averaging, which interferes with the ability to characterize complex anisotropic structures.

To overcome some of these limitations, recent studies have focused on developing machine learning algorithms to automate material structure characterization. Many of these studies have concentrated on using supervised machine learning for crystal structure identification and have shown improvements over standard techniques. Geiger and Dellago and Dietz et al.⁷ for example, classify crystalline structures in polymorphic and mixed phase systems more accurately by using a set of local structure functions that are fed into a neural network. Ziletti et al.⁸ use convolutional neural networks to automate the process of classifying crystal symmetries, even in the presence of substantial defects, using images of computationally-generated diffraction patterns. Similar supervised learning methods have been used to analyze a variety of other ordered systems, including graphene.⁹

Other studies have concentrated on crystal structure identification using unsupervised learning techniques, such as dimensionality-reduction algorithms. These unsupervised methods allow for the discovery of previously unidentified local structural features, a task that is often out of reach for standard methods and many supervised learning algorithms. For example, Reinhardt et al.¹⁰ use diffusion maps and cluster-
ing to identify new local structures in colloidal crystals, and Spellings and Glotzer use spherical harmonic functions in conjunction with Gaussian mixture models to automatically identify different crystalline arrangements without relying on a set of reference structures. Unsupervised learning has also been used to study systems ranging from Lennard-Jones crystals to proteins.13,18

Although these machine learning approaches have shown substantial improvements over standard techniques when applied to a variety of systems, many of them still rely on the development of complex, hand-crafted descriptors of local particle environments. The results of machine learning analyses can be quite sensitive to the definitions of these descriptors.14

Moreover, most of these studies have focused on characterization of crystalline or semi-crystalline structures, with little attention given to analysis of completely amorphous systems such as glassy materials.

The characterization of glassy systems is especially challenging because their lack of long-range order makes it difficult to define structural metrics. Nevertheless, several recent studies have been able to manually identify structural metrics for glassy systems. For example, Hu et al.20 defined a metric, the average degree of local five-fold symmetry, that can differentiate between configurations of metallic liquids and glasses and that has a quantitative relationship with dynamical properties of these liquid-cooled glasses generated by molecular dynamics simulations. By representing particle configurations of liquids and glasses as graphs, we are able to distinguish between them with high accuracy. Moreover, by using a technique called self-attention, we are able to extract an interpretation of how message passing neural networks evaluate liquid and glass configurations. This not only provides us with a general method for identifying important structural features in amorphous materials, but it also gives us a unique insight into how these neural networks process data. This result provides clear proof of concept that message passing neural networks could be used in more complex and demanding classification and characterization tasks that stymie standard techniques.

II. METHODS

A. Simulation Details

Glasses are kinetically arrested states of matter which are generally prepared by cooling a liquid to temperatures below the glass transition, \( T_g \), of the corresponding bulk material.21 When cooling is sufficiently rapid, the system avoids crystallization and instead solidifies into a glass, an amorphous state which has an atomic structure similar to that of a liquid but with the mechanical properties of a solid. The specific properties of these liquid-cooled glasses depend on the rate at which they are cooled, as lower cooling rates lead to materials that lie deeper in the underlying potential energy landscape.

To simulate two-dimensional liquids and liquid-cooled glasses, we used the Kob-Andersen model, which consists of a binary mixture of spheres whose glass-forming behavior in the bulk has been studied extensively.22 This binary mixture is comprised of 65% type A and 35% type B particles which have unit mass and which interact according to the pairwise Lennard-Jones potential,

\[
V_{ij}(r) = \begin{cases} 
4\varepsilon_{ij} \left[ \left( \frac{\sigma_{ij}}{r} \right)^{12} - \left( \frac{\sigma_{ij}}{r} \right)^{6} \right] & \text{if } r \leq r_{\text{cut,ij}} \\
0 & \text{if } r > r_{\text{cut,ij}} 
\end{cases}, \quad i, j \in \{A, B\},
\]

In this work, we use deep learning to accurately classify amorphous materials and to derive new metrics that characterize their structures. We use two-dimensional liquids and liquid-cooled glasses generated by molecular dynamics simulations as archetypal examples of amorphous materials. Our classification algorithms make no a priori assumptions about local relationships between particles and are not dependent on complex hand-crafted descriptors that define local particle environments.

We explore the application of two different types of deep learning algorithms: convolutional neural networks and message passing neural networks. Convolutional neural networks have been used in a variety of material classification tasks,31 including the classification of ordered and disordered configurations from simulations of the Ising model,32 but to the best of our knowledge they have never before been used to classify different amorphous material structures from molecular dynamics simulations. By rendering particle configurations of liquids and glasses as two-dimensional images, we are able to distinguish between them with high accuracy using convolutional neural networks. However, there are several limitations that accompany the use of convolutional neural networks, including the potential introduction of artifacts via image rendering and limited interpretability.

We overcome these issues by using message passing neural networks, which operate directly on the Cartesian coordinates of particles by representing a configuration of particles as a graph. Message passing neural networks have been used previously to predict properties of molecules, such as toxicity and solubility,33 and to predict properties of crystals, such as formation energy and shear modulus.22 However, to the best of our knowledge, message passing neural networks have never before been used to classify different amorphous material structures from molecular dynamics simulations. By representing particle configurations of liquids and glasses as graphs, we are able to distinguish between them with high accuracy. Moreover, by using a technique called self-attention, we are able to extract an interpretation of how message passing neural networks evaluate liquid and glass configurations. Using this interpretation, we derive three novel structural metrics that characterize glass formation and that can differentiate between liquid and glass configurations without the use of machine learning. This not only provides us with a general method for identifying important structural features in amorphous materials, but it also gives us a unique insight into how these neural networks process data. This result provides clear proof of concept that message passing neural networks could be used in more complex and demanding classification and characterization tasks that stymie standard techniques.
where $r$ is the distance between a pair of particles, $\varepsilon_{ij}$ characterizes the depth of the potential, and $\sigma_{ij}$ characterizes the finite distance below $r_{cut,ij}$ at which the potential is zero. Specific parameter values for different values of $i$ and $j$, given in Lennard-Jones units, are shown in Table I.

We performed 20,000 independent simulations of this model using LAMMPS. Each simulation contained a fixed total of 4,320 particles in a simulation box of length 60 in the $x$- and $y$-directions. The inherent structure energy per particle of the system using the FIRE minimization algorithm is the potential energy brought to its local minimum.

Each of these simulations models a liquid that cools and solidifies into a glass below $T_g$. For 10,000 of the simulations we used $t_{cool} = 2 \times 10^7$ cooling steps, and for the other 10,000 we used $t_{cool} = 2 \times 10^5$ cooling steps. We calculated the average glass transition temperature, $T_g$, for each cooling rate by identifying two linear regimes in each of the average inherent structure energy curves, one corresponding to the supercooled liquid regime and the other to the glass regime. We then fit the data in these regimes and calculated the intersection point, as shown in Figure 1(a), which gives $T_g \approx 0.37$ for $t_{cool} = 2 \times 10^7$ and $T_g \approx 0.39$ for $t_{cool} = 2 \times 10^5$. The glass configurations generated using $t_{cool} = 2 \times 10^7$ are at a lower average inherent structure energy than those generated using $t_{cool} = 2 \times 10^5$ because $t_{cool} = 2 \times 10^5$ corresponds to a lower cooling rate.

### B. Datasets

Our goal was to train machine learning algorithms to perform a binary classification task: identify a particle configuration as a liquid or a glass. To that end, we used the particle configurations generated by the simulations described in §II A to construct six datasets.

Datasets 1 through 5 are each composed of 10,000 glass configurations and 10,000 liquid configurations, all taken from the simulations with $t_{cool} = 2 \times 10^7$. The glass configurations in each dataset are all at a temperature of 0.05. In order of increasing dataset number, the liquid configurations are at temperatures 1.99, 1.76, 0.96, 0.55, and 0.44, as enumerated in Figure 1(b).

In datasets 1 through 5, the liquid configurations are at higher inherent structure energies than the glass configurations. In order to compare liquid and glass configurations at the same average inherent structure energy, we constructed dataset 6, labeled in Figure 1(a). This dataset has 10,000 glass configurations at a temperature of 0.05 taken from simulations with $t_{cool} = 2 \times 10^5$ and 10,000 liquid configurations at a temperature of 0.55 taken from simulations with $t_{cool} = 2 \times 10^7$.

Each dataset serves as a test for a machine learning algorithm’s ability to distinguish between amorphous material structures, and they are numbered roughly in order of increasing difficulty. Dataset 1 represents a relatively easier test, because the liquid and glass configurations in this dataset have a large difference in average inherent structure energy and distinctly different local structures, as exhibited by their average radial distribution functions (see Figure 1 in the Supplementary Information). Datasets 2 through 5 represent increasingly difficult tests as the differences in average inherent structure energy decrease and the radial distribution functions converge. Dataset 6 represents one of the most difficult tests, because the liquid and glass configurations are, on average, at the same inherent structure energy, as shown in Figure 1(a), and have very similar average radial distribution functions (see Figure 1 in the Supplementary Information).

### C. Convolutional Neural Networks

In this section we give a brief overview of convolutional neural networks (CNNs) and a description of the architecture and training routine for CNNs used in this work.

Neural networks have been proven capable of approximating a wide set of functions. Convolutional neural networks, which use convolutional layers in addition to the fully connected layers that characterize a basic feed forward network, have been shown to excel at computer vision tasks ranging from assessing cancer risk in radiology scans to galaxy morphology classification in telescope images. As explained in 11 CNNs have also been used to classify a variety of materials, including crystal structures and Ising model configurations.

In a convolutional layer, a set of matrices, or kernels, is convolved with an input matrix to produce a set of output matrices, or feature maps. Notably, convolutions are equivariant to translation, a property that underlies the effectiveness of CNNs in detecting features such as edges and shapes in different locations of an image, while also significantly reducing the number of parameters compared to a basic fully connected feedforward network.

#### 1. Network Architecture

The CNN that we developed, shown in Figure 2, has two convolutional layers. The first has 6 kernels with dimensions $10 \times 10 \times 3$ along with a bias vector and ReLU activation, with

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**TABLE I:** Lennard-Jones parameters for simulations of liquids and liquid-cooled glasses. Note that $\varepsilon_{ij} = \varepsilon_{ji}$, $\sigma_{ij} = \sigma_{ji}$, and $r_{cut,ij} = r_{cut,ji}$.

<table>
<thead>
<tr>
<th></th>
<th>$\varepsilon$</th>
<th>$\sigma$</th>
<th>$r_{cut}$</th>
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</thead>
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<tr>
<td>$\varepsilon_{AA}$</td>
<td>1.00</td>
<td>1.00</td>
<td>$r_{cut,AA} = 2.50$</td>
</tr>
<tr>
<td>$\varepsilon_{AB}$</td>
<td>1.50</td>
<td>0.80</td>
<td>$r_{cut,AB} = 2.00$</td>
</tr>
<tr>
<td>$\varepsilon_{BB}$</td>
<td>0.50</td>
<td>0.88</td>
<td>$r_{cut,BB} = 2.20$</td>
</tr>
</tbody>
</table>

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The inherent structure energy of a configuration, used to quantify a configuration’s stability, is the potential energy brought to its local minimum. Each simulation contained a fixed total of 4,320 particles in a simulation box of length 60 in the $x$- and $y$-directions. The inherent structure energy per particle of the system using the FIRE minimization algorithm is the potential energy brought to its local minimum. Each simulation was performed in the canonical NVT ensemble with periodic boundary conditions in the $x$- and $y$-directions, and the temperature was reduced linearly from an initial temperature of 2.0 to a final temperature of 0.05 in Lennard-Jones units using a Nose-Hoover thermostat for $t_{cool}$ simulation steps. During each simulation we recorded the inherent structure energy per particle of the system using the FIRE minimization algorithm. The inherent structure energy curves, one corresponding to the supercooled liquid regime and the other to the glass regime, were fit in these regimes and calculated the intersection point, as shown in Figure 1(a), which gives $T_g \approx 0.37$ for $t_{cool} = 2 \times 10^7$ and $T_g \approx 0.39$ for $t_{cool} = 2 \times 10^5$. The glass configurations generated using $t_{cool} = 2 \times 10^7$ are at a lower average inherent structure energy than those generated using $t_{cool} = 2 \times 10^5$ because $t_{cool} = 2 \times 10^5$ corresponds to a lower cooling rate.
Figure 1: (a) Average inherent structure energies per particle as a function of temperature for 10,000 configurations with $t_{cool} = 2 \times 10^5$ and 10,000 configurations with $t_{cool} = 2 \times 10^7$. $T_g$ for each curve is calculated as the intersection of linear approximations to the supercooled liquid regime and the glass regime, as shown by the dotted black lines. We consider configurations at temperatures above $T_g$ to be liquids and configurations below $T_g$ to be glasses. The magenta and blue points indicate the temperatures at which we select glass and liquid configurations for dataset 6, respectively. Standard deviations of energy values, not shown here, are on the order of $10^{-3}$.

(b) Average inherent structure energy per particle as a function of temperature for the 10,000 configurations with $t_{cool} = 2 \times 10^7$. The magenta point indicates the temperature at which we select glass configurations for datasets 1 through 5. The blue points indicate the temperatures at which we select liquid configurations, labeled by dataset number.

Figure 2: Convolutional neural network (CNN) architecture used in this work. Two convolutional layers are followed by a fully connected layer and an output layer. In order to be analyzed by a CNN, particle configurations are rendered as images.

no zero padding and a stride of 1. The second has 16 kernels with dimensions $5 \times 5 \times 6$ along with a bias vector and ReLU activation, also with no zero padding and a stride of 1. The 16 feature maps that are output by these layers, which have dimensions $237 \times 237$, are then flattened and fed into a fully connected layer with 80 neurons and a bias vector, followed by ReLU activation. Dropout, a regularization method that has been shown to reduce overfitting, is applied to this fully connected layer followed by a final output layer with two neurons. We used TensorFlow to build and train these models and all of our code is available in our GitHub repository.

D. Message Passing Neural Networks

In this section we give a brief overview of message passing neural networks (MPNNs) and a description of the architecture, training routine, and interpretation scheme for MPNNs used in this work.

As explained in §I, MPNNs have been used to predict properties of molecules and crystals by representing them as graphs, with atoms corresponding to nodes and bonds corresponding to edges. To the best of our knowledge, however,
MPNNs have never before been used to classify and analyze large ensembles of particles or amorphous materials such as glasses.

As described in the message passing framework established by Gilmer et al., MPNNs operate on undirected graphs \( G \) with node features \( x_v \) and edge features \( e_{vw} \). The MPNN processes these graphs in two phases: a message passing phase and a readout phase. In the message passing phase, the MPNN builds a representation of the input graph, and in the readout phase, the MPNN uses this representation to predict properties of interest.

The message passing phase runs for \( T \) steps. During each step \( t \), hidden states \( h_v^t \) and messages \( m_{vw}^{t+1} \) at each node \( v \) in the graph are updated using message function \( M_t \) and vertex update function \( U_t \) according to

\[
\begin{align*}
    m_{vw}^{t+1} & = \sum_{w\in N(v)} M_t(h^t_v, h^t_w, e_{vw}) \\
    h^t_v & = U_t(h^t_v, m_{vw}^{t+1}),
\end{align*}
\]

where \( N(v) \) are the neighbors of \( v \) in graph \( G \) and \( h^0_v \) is a function of the initial node features \( x_v \). The readout phase uses some function \( R \) to make a property prediction of interest based on the final hidden states according to

\[
\hat{y} = R\left(\{h^T_v \mid v \in G\}\right).
\]

### 1. Network Architecture

Here we describe the particular variant of MPNN that we use in this work, as described in Yang et al. and proposed in Dai et al. This variant, called Directed MPNN (DMPNN), uses messages associated with directed edges rather than messages associated with nodes. Our motivation in using this particular architecture is that having messages passed along edges instead of nodes in the graph is more conducive to extracting an interpretation of the network using self-attention, as described in \[11, 13\].

The D-MPNN works as follows, as shown in Figure 3. It operates on hidden states \( h^t_{vw} \) and messages \( m_{vw}^t \) at each edge connecting nodes \( v \) and \( w \). The edges in graph \( G \) are directed, so that for any two connected nodes \( v \) and \( w \), there is an edge from \( v \) to \( w \) and an edge from \( w \) to \( v \). In this way, the messages are also directed: \( m_{vw}^t \) and \( m_{vw}^{t+1} \) are distinct from \( m_{wv}^t \) and \( m_{wv}^{t+1} \).

In the message passing phase, the hidden states and messages are updated according to

\[
\begin{align*}
    m_{vw}^{t+1} & = \sum_{k\in \{N(v)\} \setminus w} M_t(x_v, x_k, h^t_{kv}) \\
    h^t_{vw} & = U_t(h^t_{vw}, m_{vw}^{t+1}).
\end{align*}
\]

Note that the message \( m_{vw}^{t+1} \) is not a function of the reverse message \( m_{wv}^t \) from the previous step. Prior to the first step of message passing, edge hidden states are initialized according to

\[
h^0_{vw} = \alpha(W_t \text{cat}(x_v, e_{vw})),
\]

where \( W_t \in \mathbb{R}^{h \times h} \) is a learned matrix, \( \text{cat}(x_v, e_{vw}) \in \mathbb{R}^h \) is the concatenation of the node features \( x_v \) for node \( v \) and the edge features \( e_{vw} \) for edge \( vw \), and \( \alpha \) is the ReLU activation function. The message passing functions \( M_t \) are given by

\[
M_t(x_v, x_w, h^t_{vw}) = h^t_{vw},
\]

and the edge update functions are given by a neural network,

\[
U_t(h^t_{vw}, m_{vw}^t) = \alpha(h^t_{vw} + W_m m_{vw}^{t+1}),
\]

where \( W_m \in \mathbb{R}^{h \times h} \) is a learned matrix with hidden size \( h \). Dropout is then applied. The presence of \( h^0_{vw} \) in the above equation provides a skip connection to the original feature vector for that edge. After \( T - 1 \) steps of this message passing, a node representation of the graph is constructed by summing inbound edge features in a final message passing step according to

\[
\begin{align*}
    m_v & = \sum_{k\in N(v)} h^{T-1}_{kv} \\
    h_v & = \alpha(W_a \text{cat}(x_v, m_v)),
\end{align*}
\]

where \( W_a \in \mathbb{R}^{h \times h} \) is a learned matrix.

The readout phase is the same as for general MPNNs. For the readout function \( R \), node hidden states are summed to obtain a feature vector for the graph

\[
h = \sum_{v \in G} h_v.
\]

Property predictions are then generated according to

\[
\hat{y} = f(h),
\]

where \( f \) is a feed forward neural network with ReLU activation and dropout at each layer.

---

**Figure 3:** Representative images of glass and liquid particle configurations from dataset 1 prepared for a CNN. Orange dots are type A particles and blue dots are Type B particles.
To prepare a dataset for a D-MPNN we extract a matrix of scaled particle coordinates $x, y \in [0, 1]$ and particle types (A or B) for each configuration. In order to be processed by a D-MPNN, the raw coordinate data for a batch of configurations is transformed into a graph, which happens on-the-fly for each batch that is loaded for training.

To do this, we first implement a step for each configuration in a batch that effectively increases the size of the dataset. We specify a hyperparameter called the window length, $w_{\text{win}} \in (0, 1)$, whose square is the area of a square window that we randomly select from each particle configuration. We select this window by drawing two random numbers uniformly from the interval $[0, 1 - w_{\text{win}}]$, called $x_{\text{rand}}$ and $y_{\text{rand}}$, and then select particles whose $x, y$ coordinates satisfy $x_{\text{rand}} \leq x \leq x_{\text{rand}} + w_{\text{win}}$ and $y_{\text{rand}} \leq y \leq y_{\text{rand}} + w_{\text{win}}$. For example, if $w_{\text{win}} = 0.5$, we select a subset of particles in a random square window corresponding to about 25% of the original set of particles. This process effectively augments the size of the dataset.

We then transform this batch of modified configurations into a graph. Each individual configuration is transformed into a graph by connecting each particle to its $k$ nearest neighbors (graph construction). Then, after the edge hidden states are initialized via a fully connected layer, message passing along the edges is carried out, as in Eqns. 3 - 7 (message passing phase). Finally, property predictions are generated via a fully connected neural network, as in Eqns. 8 and 9 (readout phase).

### Figure 4: Directed message passing neural network (D-MPNN) architecture used in this work. First, particle configurations are transformed into a graph representation by connecting each particle to its $k$ nearest neighbors (graph construction). Then, after the edge hidden states are initialized via a fully connected layer, message passing along the edges is carried out, as in Eqns. 3 - 7 (message passing phase). Finally, property predictions are generated via a fully connected neural network, as in Eqns. 8 and 9 (readout phase).
formation to \( \mathbf{h} \):
\[
\sigma = \text{softmax} \left( \alpha \left( \mathbf{h}^T \mathbf{W}_{\text{attn}} \right) \mathbf{v}_{\text{attn}} \right),
\]
(10)
where \( \alpha \) is the ReLU transformation, \( \mathbf{W}_{\text{attn}} \in \mathbb{R}^{h \times h} \) is a learned parameter matrix, and \( \mathbf{v}_{\text{attn}} \in \mathbb{R}^{h} \) is a learned parameter vector. The weights \( \sigma \) are then dotted with the initial row vectors to yield a new set of hidden edge states:
\[
\mathbf{h}' = \sigma \mathbf{h}.
\]
(11)
Finally, the row vectors of \( \mathbf{h}' \) replace the corresponding values of the edge features from Eqn. \( \text{[3]} \) and are applied as such to the next round of message passing. The weights \( \sigma \) represent the "attention" that the network is giving to each edge in the graph. As discussed in §III, we then used the \textit{networks} package in Python to quantify attributes of the graph structure of these self-attention weights for glass and liquid particle configurations.

E. Hyperparameter Optimization and Cross Validation

There are seven hyperparameters in our D-MPNN: window length \( l_{\text{win}} \), number of nearest neighbors \( k \), number of message passing steps \( T \), dropout probability \( p \), hidden size \( h \), number of feed forward layers \( f \), and batch size \( n_b \). There are two hyperparameters in our CNN: dropout probability and batch size. To discover optimal values for these hyperparameters, we used a Bayesian optimization scheme called a Tree-structured Parzen Estimator (TPE), implemented in the \textit{hyperopt} package. As described in Bergstra \textit{et al.}, TPE is a type of sequential model-based global optimization algorithm which discovers optimal hyperparameters by modeling the loss function with a surrogate probability model and making increasingly well-informed guesses for a specified number of iterations. We found that batch sizes greater than five for D-MPNNs sometimes exceeded GPU memory, so we fixed \( n_b = 5 \) for D-MPNNs.

<table>
<thead>
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<th>Hyperparameter</th>
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</tr>
</thead>
<tbody>
<tr>
<td># nearest neighbors (( k ))</td>
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<td>5</td>
<td>1</td>
</tr>
<tr>
<td>window length (( l_{\text{win}} ))</td>
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<td>hidden size (( h ))</td>
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<tr>
<td># message passing steps (( T ))</td>
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<td>6</td>
<td>1</td>
</tr>
<tr>
<td>dropout probability (( p ))</td>
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<td>0.5</td>
<td>0.05</td>
</tr>
<tr>
<td># feed forward layers (( f ))</td>
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<td>3</td>
<td>1</td>
</tr>
<tr>
<td>batch size (( n_b ))</td>
<td>10</td>
<td>100</td>
<td>10</td>
</tr>
</tbody>
</table>

TABLE II: Ranges of values used for hyperparameter optimization. To discover optimal values in these ranges, we used a Bayesian optimization scheme called a Tree-structured Parzen Estimator (TPE). TPE discovers optimal hyperparameters by modeling the loss function with a surrogate probability model and making increasingly well-informed guesses for a specified number of iterations. We found that batch sizes greater than five for D-MPNNs sometimes exceeded GPU memory, so we fixed \( n_b = 5 \) for D-MPNNs.

III. RESULTS AND DISCUSSION

A. CNNs

The CNNs classified liquid and glass configurations in datasets 1 through 6 with greater than 0.98 AUC, as shown in Figure 6. All of the models generalized successfully to other datasets, but the models trained...
on the more difficult tasks (i.e. datasets 5 and 6) performed best across the board.

A central question that arises is how the CNNs are making classification decisions. Since a CNN can accurately identify liquid and glass configurations prepared at the same inherent structure energy (dataset 6), these algorithms are not simply computing energy as a means for classification. We tried several methods for interpreting the CNNs, including visualizing feature maps and examining whether specific geometrical structures exist. These methods have been used in previous work to facilitate interpretation of neural networks. However, none of these methods were successful at providing an interpretation. Further work is needed to answer this question.

Besides the challenge of interpreting the CNNs, there were several other issues. The CNNs do not incorporate rotational invariance, a symmetry that is present in our data, and they require each input image to have the same size, limiting their flexibility. Moreover, rendering particle configurations as images introduces an artificial radius to these particles which is not present in the underlying system. These issues motivated our study of message passing neural networks. As discussed above, we found that message passing neural networks overcome all of these challenges.

### B. D-MPNNs

D-MPNNs classified liquid and glass configurations in datasets 1 through 6 with greater than 0.98 AUC. As shown in Figure 5, the D-MPNNs performed at least as well as the CNNs on all six datasets and had a higher average AUC than the CNNs on datasets 4 through 6 (see Figure 15 in the Supplementary Information for a complementary accuracy report). These results show that, like CNNs, D-MPNNs are capable of classifying amorphous materials with no a priori information about particle interactions and no hand-crafted descriptors of local particle environments, even when the material structures are only subtly different. It is true that we created the particle graphs by connecting nearest neighbors, but we do not restrict the D-MPNN to focus on any specific local environments in the graph (different values for number of nearest neighbors (k) are explored during optimization).

Similar to the CNNs, the D-MPNNs trained on one dataset were able to make accurate classifications on configurations from other datasets, as exhibited in Figure 15(b). Again, the models trained on the more difficult tasks perform best across the board. All of the D-MPNN models, on average, generalize better than the CNN models.

Besides their superior performance, the D-MPNNs have several distinct advantages over the CNNs. Because information from neighboring particles in the graph is summed, the D-MPNNs are invariant to permutations and rotations of the graph, taking advantage of a natural symmetry in the system. The D-MPNNs also operate directly on the particle coordinates without introducing any unnecessary artifacts that might appear in an image representation. In addition, they are flexible and can process different graph sizes (number of particles). We took one of the optimal D-MPNNs trained on dataset 1, which was trained using \( l_{\text{win}} = 0.2 \) (approximately 173 particles per graph), and successfully performed inference on both smaller and larger graphs from dataset 1, as shown in Figure 7.

Perhaps the most significant advantage of D-MPNNs is that, when imbued with the self-attention mechanism described in §II D 3, they are able to produce a clear interpretation of how they are making classification decisions. We illustrated this by training a D-MPNN on one of the outer train sets of dataset 1 with the self-attention mechanism in place and using an optimal set of hyperparameters. The model achieved an AUC of 0.995 on the corresponding outer test set. We then visualized the attention weights from configurations in the outer test set. Representative visualizations for glass and liquid configurations are shown in Figures 8(a) and 8(b), respectively.

There are clear differences in the graph structure of self-attention weights in these visualizations. The high attention weight edges form one large connected graph in the glass but multiple smaller disjoint graphs in the liquid. Several type B (blue) particles are connected to each other with high attention edges in the liquid, but none are connected in the glass. There are a larger number of type A (orange) particles that are isolated from high attention weight graphs in the liquid compared to the glass. Isolated type A particles and pairs of type B particles connected with high attention edges are highlighted in Figures 8(c) and 8(d). Also, all of the high attention edges in both liquid and glass are connected to at least one type B particle (there appear to be no isolated type B particles).

We also tested this model, which was trained using \( l_{\text{win}} = 0.2 \) (approximately 173 particles per graph), on larger configurations generated with \( l_{\text{win}} = 0.3 \) (approximately 389 particles per graph). The model successfully generalized to these larger configurations, achieving an AUC of 0.957, and yielded similar self-attention features, as shown in Figure 16 in the Supplementary Information.

We quantified these features by computing the average number of disjoint graphs, average number of high attention edges in both liquid and glass configurations. These calculations can be used as a means of interpretation.
Figure 6: These plots show how models trained on one dataset perform on all other datasets. In each of the plots in this figure, the number in row \( i \), column \( j \) is the average classification AUC on dataset \( j \) of the optimal models trained on dataset \( i \). For \( i = j \) we report the average three-fold nested cross validation AUC. For \( i \neq j \), we report average AUC using an outer test subset of dataset \( j \). The optimal models, three CNNs and three D-MPNNs for each dataset, were generated from the three-fold nested cross validation procedure described in §II E. (a) shows average CNN AUCs and (b) shows average D-MPNN AUCs.

Figure 7: D-MPNN AUC on an outer test set of dataset 1 with different values of \( l_{\text{win}} \) (number of particles). This model was trained on dataset 1 using only \( l_{\text{win}} = 0.2 \) (approximately 173 particles), which is highlighted in purple.

Average values of the three metrics are plotted as a function of temperature for configurations from simulations with \( t_{\text{cool}} = 2 \times 10^7 \) in Figure 10. All three of these structural metrics exhibit a similar dependence on temperature. They decrease at rapid rates above \( T_g \), but immediately below \( T_g \), these rates change and the metrics decrease more slowly. We fit each metric to a linear regression model of the form

\[
y_i = \alpha_0 + \alpha_1 T_i + \alpha_2 \mathbb{I} + \alpha_3 T_i \mathbb{I} + \epsilon_i
\]  

using ordinary least squares, where \( T \) is temperature, \( \mathbb{I} \) is an indicator variable with a value of 0 below \( T_g \) and a value of 1 above \( T_g \), and \( \epsilon \) is a random error term. The regressions returned positive-valued interaction coefficients \( \alpha_3 \) with p-values less than 0.05, confirming that the difference in slopes above and below \( T_g \) is statistically significant for all three metrics. Figure 17 shows these metrics plotted separately as a
Figure 8: Self-attention visualizations, with the attention weights computed on the $T - 1$th step of message passing. All connected particles in the graph are joined with a green line whose width is proportional to the magnitude of the attention weight. Some of the edges have such small attention weights that these lines are just barely visible. Note that each connected pair of particles actually has two edges, because the graph is directed. Here, we visualize the edge with the higher weight. (a) Glass configuration generated with $l_{win} = 0.2$. (b) Liquid configuration generated with $l_{win} = 0.2$. (c) and (d) show the same attention weight visualizations as (a) and (b), respectively, but with isolated type A particles highlighted with red circles and pairs of type B particles connected by high attention edges highlighted with blue ellipses.

Function of temperature along with predictions from the linear regression models. Similar to the five-fold symmetric metric in Hu et al., which was derived by hand, these metrics have a temperature dependence that describes the structural evolution of a liquid during glass formation.

We were able to derive novel structural metrics for two-dimensional liquid and glass configurations directly from the self-attention features generated by a D-MPNN. The neural network generated these features based on raw particle coordinates alone and did not rely upon complex local descriptors, extensive spatial averaging, or a set of reference structures. This provides clear proof of concept that D-MPNNs are an effective tool not only for classifying amorphous materials but also for identifying structural features in complex systems.

IV. CONCLUSION

In this work, we showed that CNNs and D-MPNNs are both effective tools for amorphous materials classification, as they can classify two-dimensional liquids and liquid-cooled glasses with greater than 0.98 AUC. We also demonstrated ways in which D-MPNNs are superior to CNNs in this context, including their ability to operate on raw particle data without introducing artifacts, to achieve better classification.
performance, to process configurations with different numbers of particles, and to provide a clear and quantifiable interpretation of the classification process. Using the interpretation that we extracted from D-MPNNs with a self-attention mechanism, we derived three novel structural metrics that characterize glass formation.

Moving forward, we believe that D-MPNNs could be applied to more difficult classification tasks that elude standard techniques. Specifically, we are interested in applying D-MPNNs to analyze other types of glassy materials, such as vapor-deposited glasses. A natural next step also includes using D-MPNNs to classify three-dimensional liquids and glasses, which could provide new insight into the local structure of these materials. In our work we performed graph-level predictions, but it is also possible to use D-MPNNs to perform node-level predictions, which could be used to extract even more granular information about local structures around specific particles or to identify defect sites in materials. And finally, further steps can be taken to improve the interpretability of attention mechanisms in these networks, including incorporating node attention in addition to edge attention and using multi-headed attention or other attention pooling techniques. In our work, we manually analyzed self-attention visualizations, but as the field of neural network interpretation advances, future work on developing tools that improve and automate this process would make deep learning analysis of amorphous material structures even more effective.

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In each of the 20,000 simulations that we performed, the same initial configuration of particles is used as a starting point. This configuration is minimized using the FIRE minimization algorithm, which brings the configuration to its local potential energy minimum. To do this we used an energy stopping tolerance of $1 \times 10^{-10}$, a force stopping tolerance of $1 \times 10^{-5}$, a maximum of 1,000 iterations of the minimizer, and a maximum of 1,000 force and energy evaluations. To ensure independence of the simulations, the particles in this configuration are then given initial velocities drawn from a Gaussian distribution with mean zero and a total linear momentum of zero and a temperature of 2.0. The seeds used to initialize the random number generator for each simulation are generated using the Bash $RANDOM$ function, which returns a pseudorandom integer in the range 0 to 32,767.

These specific random displacement values were determined by visual inspection and trial and error.

Appendix A: Simulation Details

Unscaled coordinates refer to the raw coordinates output by a simulation. Scaled coordinates refer to the raw coordinates normalized so that they lie within the unit interval [0, 1].

For example, see https://distill.pub/2017/feature-visualization/.

These specific random displacement values were determined by visual inspection and trial and error.

is zeroed in each dimension at every timestep. Particle configurations (both scaled and unscaled coordinates) are recorded every 100,000 steps.

After the simulation steps are complete, we measure the system’s inherent structure energy, also at intervals of 100,000 steps, by loading in the scaled configurations output from the simulation. To compute inherent structure energy, we again use the FIRE minimization algorithm as described above, this time with a maximum of 10,000,000 iterations and a maximum of 10,000,000 force and energy calculations.

Figure 11 shows comparisons of the average radial distribution functions of liquid and glass configurations in dataset 1, where the radial distribution function is defined as

$$g_{ij}(r) = \frac{N}{\rho N_i} \sum_{m=1}^{N_i} \sum_{n=1}^{N_j} \langle \delta(r - r_{mn}) \rangle, \quad i, j \in \{A, B\}. \quad (A1)$$

Here, $N$ is the total number of particles, $N_i$ is the total number of type $i$ particles, $N_j$ is the total number of type $j$ particles, $\rho$ is the total density, $\delta$ is the delta functional, $r_{mn}$ is the vector from particle $m$ to particle $n$, and the average is taken over all vectors $r$ with magnitude $r$. We calculated the radial distribution functions in LAMMPS using 500 histogram bins and a cutoff distance of 10$\sigma_{AA}$. As expected, these radial distribution functions exhibit clear structural differences between the liquid and glass configurations, with the glasses having higher and sharper peaks than the liquids.

At $T = 0.55$ the 10,000 configurations from $t_{cool} = 2 \times 10^7$, which are liquids, have a mean inherent structure energy of $-3.86698$ with a standard deviation of 0.00382, while at $T = 0.05$ the 10,000 configurations from $t_{cool} = 2 \times 10^5$, which are glasses, have a mean inherent structure energy of $-3.86744$ with a standard deviation of 0.00288. These sets of configurations, therefore, have approximately the same energy.

We computed average radial distribution functions comparing the configurations from these glasses and liquids, which comprise dataset 6, shown in Figure 12. Note that the structures are much more similar than those shown in Figure 11.

Appendix B: Image Data for CNNs

We utilized CNNs to classify liquid and glass configurations by rendering them as images. For a given configuration, we load its scaled particle coordinates and particle types as a numpy array in Python and used the matplotlib package to save the configuration as a 250 x 250 pixel PNG image with 100 dots per inch. As noted at https://lammps.sandia.gov/doc/dump.html: “Because periodic boundary conditions are enforced only on timesteps when neighbor lists are rebuilt, the coordinates of an atom written to a dump file may be slightly outside the simulation box...atom coords are written in a scaled format (from 0 to 1)...an x value of 0.25 means the atom is at a location 1/4 of the distance from xlo to xhi of the box boundaries.” Upon inspection, most particles have coordinate values between 0 and 1, with only occasional exceptions slightly outside these bounds, e.g. values such as 1.0001. This does not affect our analyses. We
Figure 11: Average (a) A-A, (b) A-B, and (c) B-B radial distribution functions of 10,000 configurations at $T = 1.99$ and 10,000 configurations at $T = 0.05$ with $t_{cool} = 2 \times 10^7$ (dataset 1). Glasses (green) are clearly distinguishable from liquids (blue) because of their higher and sharper radial distribution function peaks.

remove the axes and the frame from the image so that only the particles are rendered in the image, without any additional artifacts. Type A particles are represented as orange dots and type B particles as blue dots using the scatter function, with a dot size of $s = 1$. The image is then slightly cropped in order to remove any unnecessary white space. In some images this slightly truncates particles at the edges of the image, but we found that this does not inhibit the performance of the CNN. Metadata representing these images for a dataset are saved in a JSON file.

During training the generator function loads a batch of training images into a numpy array by reading their file paths from the metadata JSON file. This function converts the liquid and glass labels into a one-hot vector representation, i.e. ([0, 1] for glasses and [1, 0] for liquids).

Appendix C: Hyperparameter Optimization for D-MPNNs

The hyperopt Sequential Model-based Global Optimization (SMBO) algorithms form a probabilistic model that maps hyperparameters to a probability of a score on the loss function, $P(y|x)$. This probabilistic model is called a surrogate. SMBO methods work by choosing the next set of hyperparameters to test on the loss function by selecting hyperparameters that perform best on the surrogate function. 

As
Figure 12: Average (a) A-A, (b) A-B, and (C) B-B radial distribution functions of 10,000 configurations at $T = 0.55$ with $t_{\text{cool}} = 2 \times 10^7$ and 10,000 configurations at $T = 0.05$ with $t_{\text{cool}} = 2 \times 10^5$ (dataset 6). Glasses (green) are distinguishable from liquids (blue) because of higher and sharper radial distribution function peaks, but the differences are much less severe than in Figure 11 as the glasses and liquids here have the same average inherent structure energy.

Each set of hyperparameters is evaluated, the method updates the surrogate probability model in order to make increasingly well-informed guesses. After a specified number of iterations, the method suggests the optimal set of hyperparameters. The specific SMBO method that we use in this work is called a Tree-structured Parzen Estimator (TPE), which is thoroughly described in Bergstra et al. and which we follow closely here. There are different ways of identifying which hyperparameters to select based on the surrogate model in an SMBO method, but one of the most effective is a metric called Expected Improvement, otherwise known as an "exploration-exploitation" criterion. Given a desired threshold value for the objective function, $y^*$, and some set of hyperparameters $x$, the Expected Improvement is given by

$$
\text{EI}_{y^*}(x) = \int_{-\infty}^{y^*} (y^* - y)P(y|x)dy. \tag{C1}
$$

The first factor in the integrand promotes values in regions that are likely to contain objective function minima (exploitation), while the second term promotes regions that have greater uncertainty (exploration). When this integral is positive, it
means that the hyperparameter set $x$ is expected to yield an improvement relative to the threshold value $y^*$. In the TPE algorithm, instead of modeling the surrogate directly as \( p(y|x) \), this method uses Bayes rule, \( p(y|x) = \frac{p(x|y)p(y)}{p(x)} \), to model \( p(x|y) \) and \( p(y) \) instead. \( p(x|y) \) is broken down into \( l(x) \) and \( g(x) \), such that

\[
p(x|y) := \begin{cases} l(x) & y < y^* \\ g(x) & y \geq y^*. \end{cases} \tag{C2}
\]

In other words, we create two different distributions for the hyperparameters: one where the objective function value is less than the threshold, \( l(x) \), and one where the objective function value is greater than the threshold, \( g(x) \). These non-parametric densities are constructed after some number \( K \) of evaluations of the objective function. \( y^* \) is chosen to be slightly greater than the best observed objective function score.

In this approach, the Expected Improvement is given by

\[
EI(x) = \int_{-\infty}^{y^*} (y^* - y) \frac{p(x|y)p(y)}{p(x)} dy,
\]

which can be rearranged as

\[
EI(x) \propto (\gamma + \frac{g(x)}{l(x)}(1-\gamma))^{-1}, \tag{C4}
\]

where \( \gamma = p(y < y^*) \) (no specific \( p(y) \) is necessary). So, the TPE works by drawing sample hyperparameters from \( l(x) \), evaluating them in terms of \( g(x)/l(x) \), and returning the set \( x \) that gives the best expected improvement value.

Appendix D: Attempts to Interpret CNNs

In an attempt to interpret the CNNs, we computed the feature maps produced by the first convolutional layer for several glass and liquid images that were correctly classified as such by the best network trained on dataset 1. Visualizing components of neural networks is gaining traction as a method for interpretation\(^{[15]}\). Visualizations of these feature maps for representative liquid and glass configurations are shown in Figure [13]


We attempted a different method for interpretation based on our prior knowledge about local geometric structure in two-dimensional Kobb-Anderson binary mixtures. As evidenced in Reid et al., the degree of five-fold symmetry in a liquid-cooled glass is much higher than that in liquids at higher temperatures and higher inherent structure energies.\(^{[23]}\) One possibility is that the network has identified this geometric quantity as a means for classifying liquids and glasses and that the kernels in the CNN have been trained to identify local pentagonal arrangements of particles. As in Reid et al., we consider a region of five-fold symmetry to be characterized by a type B (blue) particle surrounded immediately by a pentagon of type A (orange) particles. Perhaps, if the CNN is trained to correlate high concentrations of five-fold symmetry with glassy materials, an artificial image that is saturated with five-fold symmetry patterns will be classified by the network as a glass.

We tested this hypothesis by constructing artificial images imbued with five-fold symmetry as follows. First, we placed 1,512 (35% of 4,320) type B particles in a grid, evenly spaced, with the x- and y-axes ranging from 0 to 1. Each particle was then given a slight random displacement in the x- and y-directions from their initial placements, corresponding to a uniform random number in the range -1/195 to 1/195. We then selected 500 of these type B particles to surround with pentagonal arrangements of type A (orange) particles. To attempt to replicate the five-fold symmetry clustering effect described in Hu et al., we added pentagonal arrangements around type B particles iteratively and selected each subsequent type B particle to be a neighbor of a previously selected particle with a probability of 3/4.\(^{[20]}\) Every time a pentagonal arrangement of type A particles was placed, we arranged the particles to be at a radial distance of 1/78 from the central type B particle and gave the pentagonal arrangement a random angular rotation selected uniformly from the range \(-2\pi\) to \(2\pi\). For any pair of particles within a distance of 1/90 of each other, we removed one of the particles to prevent overlaps. Figure [14](a) shows a representative image of the result.

The remainder of the type A particles are filled in randomly, again avoiding any overlaps, to produce a final result, shown in [14](b). We again used the dataset 1 network to classify several hundred examples of these artificial images. However, all of them were classified as liquids. We also constructed images with other n-fold symmetries and with random configurations of particles, but all were classified as liquids.

Appendix E: CNN and D-MPNN Classification Results

Figure [15] shows accuracy results for CNNs and D-MPNNs.

Appendix F: D-MPNN Self-Attention

Figure [16] shows self-attention visualizations for configurations in an outer test set of dataset 1 with \( l_{win} = 0.3 \). High
Figure 13: First convolutional layer feature map visualizations for representative liquid and glass configurations. The network used here is a model optimized on dataset 1. The glass and liquid images are taken from \( t_{\text{cool}} = 2 \times 10^7 \) data and were correctly classified by the network. The axes on all of these plots correspond to pixel number. (a) is the original glass configuration; (b) is the original liquid configuration; (c) is glass feature map 1; (d) is liquid feature map 1; (e) is glass feature map 2; (f) is liquid feature map 2; (g) is glass feature map 3; (h) is liquid feature map 3; (i) is glass feature map 4; (j) is liquid feature map 4; (k) is glass feature map 5; (l) is liquid feature map 5; (m) is glass feature map 6; (n) is liquid feature map 6.

Attention edges were determined with a hard cutoff; upon inspection, there was a clear trough in the distribution of attention weights that separated those with small, almost negligible magnitudes and those with larger magnitudes. Figure 17 shows the graph-based metrics from Figure 10 plotted individually as a function of temperature.
Figure 14: (a) Initial artificial pentagonal arrangements of type A and type B particles. (b) Final artificial five-fold symmetry configuration.

Figure 15: CNN and D-MPNN average classification accuracy for datasets 1 through 6 with error bars showing standard deviation. These average values were computed using the three-fold nested cross-validation scheme described in §II E.

Figure 16: Self-attention visualizations, with the attention weights computed on the $T-1^{th}$ step of message passing. All connected particles in the graph are joined with a green line whose width is proportional to the magnitude of the attention weight. Note that each connected pair of particles actually has two edges, because the graph is directed. Here, we visualize the edge with the higher weight. (a) Glass configuration with $I_{win} = 0.3$. (b) Liquid configuration with $I_{win} = 0.3$. 
Figure 17: Average number of isolated type A particles (a), edges connecting pairs of type B particles, (b), and number of disjoint graphs (c) in configurations from simulations with $t_{cool} = 2 \times 10^7$ at a variety of temperatures. The dotted orange lines show predictions from the linear regression models for each of these curves, as described in §III. The segments of the linear models above and below $T_g$ are artificially extended to visually highlight the difference in slopes above and below $T_g$. 