Transferable Multi-level Attention Neural Network for Accurate Prediction of Quantum Chemistry Properties via Multi-task Learning

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Abstract

The development of efficient models for predicting specific properties through machine learning is of great importance for the innovation of chemistry and material science. However, predicting global electronic structure properties like frontier molecular orbital HOMO and LUMO energy levels and their HOMO-LUMO gaps from the small-sized molecule data to larger molecules remains a challenge. Here we develop a multi-level attention neural network, named DeepMoleNet, to enable chemical interpretable insights being fused into multi-task learning through (1) weighting contributions from various atoms and (2) taking the atom-centered symmetry functions (ACSFs) as the teacher descriptor. The efficient prediction of 12 properties including dipole moment, HOMO, and Gibbs free energy within chemical accuracy is achieved by using multiple benchmarks, both at the equilibrium and non-equilibrium geometries, including up to 110,000 records of data in QM9, 400,000 records in MD17 and 280,000 records in ANI-1ccx for random split evaluation. The good transferability for predicting larger molecules outside the training set is demonstrated in both equilibrium QM9 and Alchemy datasets at density functional theory (DFT) level. Additional tests on non-equilibrium molecular conformations
from DFT-based MD17 dataset and ANI-1ccx dataset with coupled cluster accuracy as well as the public test sets of singlet fission molecules, biomolecules, long oligomers, and protein with up to 140 atoms show reasonable predictions for thermodynamics and electronic structure properties. The proposed multi-level attention neural network is applicable to high-throughput screening of numerous chemical species in both equilibrium and non-equilibrium molecular spaces to accelerate rational designs of drug-like molecules, material candidates, and chemical reactions.

1. Introduction

Chemistry is indispensable in human daily life as well as the research and development of clothing, drugs, and materials, etc. Nowadays, the powerful quantum chemical calculations in combination with big databases and artificial intelligence are changing the painstaking “try and error” works to rational discovery of novel molecules and materials with the desired properties.\(^1\text{-}^8\) Computational cost of quantum chemistry calculation increases rapidly as the sizes of systems increase. To facilitate the discovery process, quantum chemistry calculations based on density functional theory (DFT) have been widely used in various chemical systems with the computational scaling of \(O(N_b^3)\), where \(N_b\) is the number of basis sets. The gold standard CCSD(T)/CBS is even more expensive with the cost of \(O(N_b^7)\). Development of lower and even linear scaling methods has aroused great interest in the past decade.\(^9\text{-}^{16}\) In spite of these advances in the quantum chemical methods, the quick prediction of various electronic structure properties is highly desired in high-throughput searching of large chemical spaces with all possible combinations of functional groups, towards the material or drug design.\(^4\text{-}^{17},^{18}\)

Many machine learning methods have been introduced in quantum chemical study for the rapid predictions of atomic forces, molecular energy, and electronic structure properties, which are of great importance for the construction of accurate force fields and complicated potential energy surfaces as well as rational design of various materials and drug-like candidates.\(^1,^{19\text{-}37}\) If the data-driven model is trained properly, it could remarkably reduce computational costs but with similar accuracy to quantum chemical calculation. High-throughput computational screening hence
becomes possible, outputting the properties of millions of compounds with broad applications in a fast and accurate way. For example, functions of novel optical materials and electronic devices are correlated with the descriptors of dipole moment, polarizability, energy levels of the highest occupied molecular orbital (HOMO) and the lowest unoccupied molecular orbital (LUMO), and so on.\(^{38, 39}\) Drug-like molecules can be screened with cohesive Gibbs free energy and other descriptors.\(^8\)

The term of electronic spatial extent, \(<R^2>\), may be useful for the design of batteries.\(^{40}\) Heat capacity is an important descriptor in predicting the properties of ionic liquids and many thermal materials.\(^{41, 42}\) However, it is still a big challenge to accurately predict some electronic structure properties such as the HOMO and LUMO energy levels, and furthermore, the HOMO-LUMO energy gap, of much larger molecules that beyond the training set with good transferability.\(^{43-45}\)

To improve the prediction accuracy of HOMO and LUMO energy levels, we propose in this paper an efficient multi-level attention neural network, named as DeepMoleNet, for the molecular systems. Our target is to establish an implicit relationship between the molecular structural information and 12 electronic structure properties including dipole moment, polarizability, HOMO, LUMO, HOMO-LUMO gap, zero point vibration energy (ZPVE), electronic spatial extent \(<R^2>\), internal energy at zero and room temperature \((U_0, U)\), enthalpy \((H)\), free energy \((G)\), and heat capacity \((C_v)\). It may be illustrative to draw an analogy between the language learning and property prediction from quantum chemical datasets. In the machine learning and translation of a sentence in a certain kind of language, the meaning of the translated word is associated with specific words and phrases in the context in neural machine translation. For a certain molecule in chemistry realm, every atom node is affected by its chemical environment adjacent to it with different weights, like that an oxygen atom always behaves differently in tetrahydrofuran and tetrahydro-2H-pyran; nitrogen atom in pyridine versus pyrimidine. The weights to address the different impacts from ‘environment’ atoms on the ‘center’ atom are called ‘attention’. In the present work, we apply multi-level attention in every message passing step which would gradually capture the influence of different atomic nodes at each ‘time step’, T,
and the attention weight varies as the node representation changes in a dynamic manner. Such a multi-level attention strategy differs significantly from some other attention algorithms, in which attention is used once at the defined step. To help the neural network learn a richer representation from atomic structural information, we use the multi-task learning (MTL), combining different known information of input data sample as the related auxiliary tasks to improve the predictive power of our main targets. In addition to the pursuit of 12 quantum chemistry properties as the learning targets, the atom-centered symmetry functions (ACSFs) descriptor is selected as the auxiliary prediction targets. ACSFs have been widely applied as the input in many chemistry applications such as predictions of organic reactions, phase transition, surface catalysis, etc. In the proposed DeepMoleNet model, we obtain the final node feature after T step multi-level attention cycle to predict the auxiliary ACSFs task. In other words, chemistry knowledge underlying the ACSFs descriptor is combined with multi-task learning to improve the generalizability and transferability of the deep learning model. As shown in Figure 1, we train with small molecules in QM959 (70,000 record with the number of atoms in the molecule, \( N_{\text{atom}} < 19 \)) and Alchemy44 (72,000 with \( N_{\text{atom}} < 22 \)) datasets, respectively, to predict the 12 quantum chemistry properties of larger molecules with up to \( N_{\text{atom}} = 29 \) in QM9 and \( N_{\text{atom}} = 38 \) in Alchemy. The present DeepMoleNet model could simultaneously predict 12 properties with better performance, accuracy, and robustness compared with other single-target training models. We even get better results by using the early epoch of multi-targets model as model initialization to train the single-target training model. Furthermore, this multi-task model is transferable to larger molecules with broad application scopes of drug-like molecules, peptides, macrocyclic molecules, oligomers, protein, and singlet fission molecules than those \( N_{\text{atom}} \leq 29 \) used for training in the QM9 dataset. This indicates the potential of our model in predicting the electronic structure properties of complex quantum chemical systems with satisfactory generalizability and transferability.
Figure 1. Illustration of the multi-level attention neural network, DeepMoleNet, which was subjected to extensive tests on quantum chemistry datasets at both equilibrium and non-equilibrium conformations as well as multiple public test sets with different applications. Each dataset is split into 2 different groups, the small sized training group (for examples, QM9: $N_{\text{atom}} \leq 18$; Alchemy: $N_{\text{atom}} \leq 22$) and the large test group (for QM9: $19 \leq N_{\text{atom}} \leq 29$; Alchemy: $23 \leq N_{\text{atom}} \leq 38$). Multi-level attention is applied to the message passing phase, and then after $T$ steps, the auxiliary task is achieved. Finally, after the readout phase, the main tasks are done to test the transferability of the model.

2. Method

Shown in Figure 2 is an illustrative flowchart for the implementation of the proposed DeepMoleNet model, which has three steps, i.e., input, message passing, and readout. In this section, we will introduce details of the learning method, databases, and error analysis of our predicted results.
Figure 2. Implementation steps of DeepMoleNet. Representation of each node is generated at first through multi-linear perceptron, and subsequently, the attention coefficient is initialized to make Hadamard product with each node. Then the node hidden states are aggregated with edge hidden states. Hidden states of LSTM are used for multi-level attention generation. After $T$ steps, ACSFs are predicted by each node representation. Finally, each node hidden state is used in set2set and atomwise neural network to predict the 12 quantum properties.

2.1 DeepMoleNet architecture

**Input.** In our DeepMoleNet model, each molecule (labeled as $M$) in QM9 and Alchemy datasets is input as a set of atom types (also called nodes), $a_i$, and atom pairs (edge types), $b_{ij}$, as shown in Figure 2. The selected features of nodes ($a_i$) and edges ($b_{ij}$) inputs are given in Table 1. For nodes, we use atom types, chirality, atomic number, hybridization, etc., which were generated by using RDkit code\textsuperscript{60}. Gasteiger partial charge, as embedded in RDKit, was used for calculating atomic charges. The selected edges include descriptors of bond type, same ring, topological path length, shortest path bonds, graph distance, extended distance, geometric distance, in which the position information is implicitly encoded in atom pair features through Gaussian expansion\textsuperscript{61} in equation (1).
\[ F_{\text{Gaussian expansion}} = e^{-\frac{(r-r_0)^2}{\sigma^2}} \] (1)

where \( r \) is the distance between 2 nodes. The Gaussian functions are centered at 20 locations with the peak center parameters, \( r_0 \), which are linearly placed between 0 and 4. We set the peak width parameter \( \sigma = 0.5 \).

In this work, we aim to design a neural network algorithm to map the relation between molecule, \( M (a_i, b_{ij}) \), and their 12 quantum chemical properties, \( P \), through \( f: \{M_{ij}\} \rightarrow P_{i=1}^{12} \). It is noticed that extended-connectivity fingerprints (ECFP4) were applied as inputs for quantum chemical property predictions.

In Table 1, we also list the input features of other machine learning methods for quantum chemical applications. The ACSFs were widely used in the construction of high-dimensional neural network potential-energy surfaces. The kernel-based machine learning of molecular properties is realized by transforming fingerprints and representations non-linearly with kernel functions. \[25, 34, 62-66 \] There are also some descriptors, such as the smooth overlap of atomic positions (SOAP) method, the bag of bonds approach and Fourier series of atomic radial distribution functions. \[67 \] However, deep learning \[70 \] can directly learn from low-level molecular structure information (e.g., atom types and bond types), and then gradually extract high-level representation through deep multiple neural network layers to predict targets. \[43, 61, \]

Since we can generally achieve acceptable performance in deep learning by ‘focusing’ on our target tasks, the information of ACSFs could be excluded from the input, instead we set ACSFs as one of the targets in the present DeepMoleNet model. Such ACSFs information could come from the training signals of the related tasks in the multi-task learning (MTL), which has been successfully used in many fields ranging from natural language processing and speech recognition, computer vision, and drug discovery. \[49-51 \] MTL is able to improve generalization by parallel training tasks with the shared representations between the related tasks. \[87 \]
Table 1. Comparisons between various neural network methods

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**Message passing neural networks with multi-level attention.** After the input section, DeepMoleNet will subsequently run message passing and readout processes. Message passing neural networks (MPNN)\textsuperscript{73} have been used to learn features from molecular graphs through abstracting the commonalities between several existing
neural models for graph structured data. It is useful to solve molecular systems on
molecular graphs $M$ with node and edge features. Several kinds of graph neural
networks have been developed with the emphasis laid on the dynamically updating
the node features in response to its neighboring node features and edge information.

When performing message passing, every node is connected to all its neighbors. In
addition, the edges between two nodes act as a modulator to transmit these messages.

We can exemplify the message passing by energy prediction of a molecule, which
consists of several atoms, and there are various interactions among these atoms. If the
two atoms are too far away from each other, their atomic interaction may be very
weak and the edges in the graph would reduce messages; but if two atoms are close to
each other, the edges in the graph would enlarge the messages due to the significant
atom-pair interaction.

We can give a mathematic expression for the message passing phase, in which
node features $x_v$ are fed into a node network to obtain a hidden feature of dimension $d$,
and edge features $e_{vw}$ are fed into an edge network to obtain a $d \times d$ matrix, respectively.
Then hidden states of each node are aggregated with message function $M_t$ to
generate the node hidden message, $m_{v}^{t+1}$, where $N(v)$ denotes the neighbors of node
$v$, as shown in equation (2).

\[
m_{v}^{t+1} = \sum_{w \in N(v)} M_t(h_v^t, h_w^t, e_{vw})
\]

Here, $M_t$ is defined as follows.

\[
M_t(h_v^t, h_w^t, e_{vw}) = A(e_{vw})h_w
\]

where $A(e_{vw})$ is the edge network.

The message function is followed by the update function. The update process can
be described by analogy of molecular system. The chemically active atoms would
strongly affect the surrounding atom; in return, the adjacent active atoms are more
likely to surround an active atom.

In the present work, we extend the MPNN framework with the multi-level
‘attention mechanism’ in the update function of the message passing stage to capture
the long ranged interaction information in quantum chemistry calculation data. Attention is similar to the way of observing an object. Our visual system tends to pay attention to some parts of the image selectively, while ignoring other irrelevant information. Therefore, attention mechanisms assign different weights to different parts of the inputs, allowing the extraction of more critical and important information with modest computational costs. In such a way, attention enables the model to distinguish different input information and explain what the model has learned, so as to give an interpretable view for the ‘black box’ deep learning systems.

The update process is presented as follows.

\[ X_v^{t+1}, q_v^{t+1} = U_t(h_v^t, m_v^{t+1}) \] (4)

When there is no attention mechanism applied, \( X_v^{t+1} \), the output of \( U \), goes back to \( h_v^{t+1} \) in the update function of the traditional MPNN\textsuperscript{73}. The \( q_v^{t+1} \) term in equation (4) is the hidden state of time-dependent update function, similar to that used in Long Short Term Memory Network (LSTM)\textsuperscript{88}. LSTM is used to update the node state for further aggregation in an order free manner. As shown in equation (5), the sigmoid activation function is used to obtain the attention score, \( Z \), which indicates the importance of this atom in the molecule.

\[ Z = \sigma(MLP(q_v^{t+1}) \cdot X_v^{t+1}) \] (5)

Then, \( h_v^{t+1} \) is the weighted information of \( X_v^{t+1} \) with the broadcast of the attention score \( Z \), as shown in equation (6).

\[ h_v^{t+1} = X_v^{t+1} \odot Z \] (6)

where \( \odot \) is the Hadamard operator.

We utilize information extracted from message passing phase and the LSTM hidden state to generate attention vector through multiplying this vector by the coefficient generated by the attention mechanism, and feed it to the LSTM block. Therefore, the whole model could be viewed as a model with the embedded pooling layer MPNN.
and attention mechanism.

Node features, $x_v$, and edge features, $e_{vw}$, run with both message function and update function for $T$ time steps in this message passing phase. After the multi-level attentional node message aggregation, the final node representation at $T$ step is used to predict the ACSFs and the outputs of readout function are used for the main target prediction of 12 quantum chemistry properties.

**Read Out.** In the present work, two readout functions, Set-to-set (Set2Set) and atom-wise summation neural networks, are used to simultaneously predict the 12 targets in a neural network. In the readout phase, a graph level representation is calculated with defined readout function $R$ according to the following equation.

$$\hat{y} = R(\{h^T_v, \; |v \in M\}).$$  \hfill (7)

where, $h^T_v$ means the final node representation of message passing in the $T$ step, $v$ is the node of the molecular graph, $M$. The selection of two kinds of readout functions is judged by the intensive or extensive nature of the 12 predicted molecular properties. Among them, HOMO, LUMO, HOMO-LUMO gap, dipole moment, and heat capacity belong to the intensive properties, for which the Set2Set is used in the multi-task training progress to make predictions. In contrast, the energy-related properties, $G$, $H$, $U$, and $U_0$, are extensive properties, whose prediction is completed by using the atom-wise summation neural networks. The remaining three kinds of properties, i.e., polarizability, $<R^2>$, and ZPVE are predicted by using the Set2Set readout for convergence consideration.

Set-to-set (Set2Set) framework has been used in enn-s2s\textsuperscript{73} model, which produces graph-level embedded representation with global attention, instead of summing up the final node states. The atom-wise summation neural networks are applied with atom-wise layers for each node separately to the node hidden representations $h_i$ with shared weights $W^l$ and biases $b^l$ on layer $l$ as follows.

$$h_v^{T,l+1} = W^l h_v^{T,l} + b^l$$  \hfill (8)
where the $h^T_{\nu}^{l}$ is the final T step output of the message passing phase at layer $l$.

The atom-wise neural network produces an output on each node representation for each node, and then predicts the target properties by simply summing up or averaging the final node states. In the DeepMoleNet, the message passing steps is set as 6 by trial and error. The learning rate is $1 \times 10^{-5}$. Adam is used as the optimization model, and the batch size is 32. Our model is run in 400 epochs, in which 20 epochs are used for warm-up. More details are shown in Table S1.

In addition, DeepMoleNet is featured as the combination of feature engineering and the deep learning approach within the framework of multi-task learning (MTL) theory. Feature engineering could gain knowledge from dataset, so we introduce an auxiliary task to predict ACSFs at the end of the message passing phase. Outputs of the readout function are used for the main targets of 12 properties prediction with the predicting loss, $L$, defined as follows.

$$L = L_{\text{main targets}} + \lambda L_{\text{ACSFs}}$$

(9),

where $L_{\text{main targets}}$ represents the loss of 12 predicted molecular properties (main targets), as shown in equation (10). $L_{\text{ACSFs}}$ is the loss of the predicted ACSFs values, shown in equation (10). Here, $\lambda$ is the loss weight of $L_{\text{ACSFs}}$. It is set to be 16 by trial and error.

$$L_{\text{main targets}} = \frac{\sum_{p}^{8 \text{ properties in Set2Set}} \sum_{i}^{all} |p^\text{DeepMoleNet}_i - p^\text{DFT}_i|}{N_{\text{molecules}}}$$

$$+ \rho \frac{\sum_{p}^{4 \text{ properties in Atom-wise NN}} \sum_{i}^{all} |p^\text{DeepMoleNet}_i - p^\text{DFT}_i|}{N_{\text{molecules}}}$$

(10),

where the first term is set for the readout 8 properties, including dipole, polarizability, HOMO, LUMO, HOMO-LUMO gap, <R2>, ZPVE, and heat capacity. For those properties using Set2Set, they are normalized, and for extensive properties, the weight is 0.5. In the second term of equation (10), $\rho=0.5$, which is the weight for all 4
energy-related properties, $U_0$, $U$, $H$, and $G$, in Atom-wise NN.

$$L_{ACSFS} = \frac{\sum_{i=1}^{N_{\text{molecules}}} |P_i^{\text{DeepMoleNet}} - P_i^{\text{ACSFs}}|}{N_{\text{molecules}}}$$ (11)

During the message passing phase, the hidden state of node is aggregated by the influence of its neighbors. However, the spatial information is lost in this way of topological information aggregation. There are many efforts devoted to MTL. One way is to combine Coulomb matrix with neural network \textsuperscript{90}. Another is using atom-centered environment vectors (AEVs) as input like AIMNet\textsuperscript{91}. Instead of using AEVs as input attributes in the deep learning approach like AIMNet\textsuperscript{91}, DeepMoleNet sets the ACSFs descriptors as the predicting targets at the end of message passing phase to help every node get the spatial information fused with the learned representations. For spatially sensitive properties such as molecular dipole moment, the auxiliary ACSFs prediction tasks may be helpful to fuse spatial information with the representation learned in the data-driven way. In subsection 3.4, we will further elaborate the performance of using ACSFs as both descriptor and prediction task.

To summarize, our model has features of (1) introducing more detailed description of atom and atom pair information in the input section; (2) using multi-level attention for node message aggregation; (3) employing the auxiliary target, ACSFs, to help the graph convolution model obtain better learned representation; and (4) adopting both set2set readout function and atomwise neural network in simultaneously predicting the 12 quantum properties, which are the main targets of this work.

2.2 Comparison with other methods

It is useful to make comparison between various deep learning methods for molecular property predictions in Table 1. In this work, the inclusion of the complete edge feature vector (bond type, spatial distance) and treating hydrogen atoms as explicit nodes in the graph are found to be crucial to get good predictions for a number of targets in multi-task learning. The enn-s2s\textsuperscript{73} selected less descriptors of node and bond
features in message passing. The difference between the Set2Set\textsuperscript{73} and the multi-level attention we used lies in the node information aggregation. We made attention on local chemical environment of each atom to extract the final node representation, while Set2Set\textsuperscript{73} further made attention with the final node representation to produce the final global graph representation. In addition, SchNet\textsuperscript{77} used element embedding for node feature and radial basis functions to incorporate the continuously filter convolutional layers in message passing and atom-wise layers for final prediction. MGCN\textsuperscript{74} developed the multi-level interaction (atom-wise, atom-pair, triple-wise atom interaction, etc.) in message passing; and DimeNet\textsuperscript{43} used directional message of angle to pass information of the surrounding atoms with Fourier-bessel basis functions. AIMNet\textsuperscript{91} used node embedding from both atomic coordinates and atomic numbers to form the atom-centered environment vectors, then obtained atomic feature vectors (AFVs) in the iterative message passing for downstream tasks (Table 1).

2.3 Datasets

In the application of DeepMoleNet, we employed multiple public datasets including QM9\textsuperscript{59}, Alchemy\textsuperscript{44}, MD17\textsuperscript{92}, ANI-1ccx\textsuperscript{93},\textsuperscript{94} quantum chemistry computation results, as shown in Table 2. Some typical molecules with potential applications in drug and material design were also studied using DeepMoleNet.

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<tr>
<td>QM9\textsuperscript{59}</td>
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<td>133,885/129,428</td>
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<td>MAE, std\textsubscript{MAE}</td>
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</tr>
<tr>
<td><strong>DFT or CCSD data @ sampled conformations</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>MD17\textsuperscript{92}</td>
<td>PBE+vdW-TS</td>
<td>1</td>
<td>3,611,115/560,000</td>
<td>Random</td>
<td>MAE</td>
</tr>
<tr>
<td>ANI-1ccx</td>
<td>CCSD(T)/CBS</td>
<td>1</td>
<td>489,571/332,196</td>
<td>Random</td>
<td>MAE, RMSE</td>
</tr>
<tr>
<td><strong>Transferability test sets</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Drug-like molecules\textsuperscript{95}</td>
<td>All the test molecules</td>
<td>2</td>
<td>24/17</td>
<td></td>
<td>MAE, Δ\textsubscript{MAE}</td>
</tr>
<tr>
<td>MPCONF196\textsuperscript{96}</td>
<td>were re-optimized</td>
<td>2</td>
<td>192/11</td>
<td></td>
<td>MAE, Δ\textsubscript{MAE}</td>
</tr>
<tr>
<td>Singlet fission molecules\textsuperscript{37}</td>
<td>with</td>
<td>2</td>
<td>262/9</td>
<td></td>
<td>MAE, Δ\textsubscript{MAE}</td>
</tr>
<tr>
<td>oligomers\textsuperscript{37}</td>
<td>B3LYP/6-31G(2df,p)</td>
<td>2</td>
<td>12</td>
<td></td>
<td>MAE, Δ\textsubscript{MAE}</td>
</tr>
<tr>
<td>protein\textsuperscript{37}</td>
<td></td>
<td>2</td>
<td>2/1</td>
<td></td>
<td>MAE, Δ\textsubscript{MAE}</td>
</tr>
</tbody>
</table>

\textsuperscript{a} The amount of data in the public dataset is given before the slash, and the number of data used in this work is shown after the slash.
2.3.1 Chemical equilibrium datasets

**QM9.** The QM9$^{59}$ dataset contains about 130,000 organic molecules composed of 9 heavy atoms including C, O, N, and F, etc. within the GDB-17 database. Various molecular properties were calculated at the theoretical level of DFT/B3LYP/6-31G(2df,p). The collected molecules in QM9 cover a wide range including (hetero-) alkane, amide, amine, alcohol, epoxy, ether, ester, chloride, aliphatic, and aromatic groups. In this work, the 110,000 QM9 data are randomly selected as the training set, 10,000 data for validation, and the rest for test set, with the molecular size, \(N_{\text{atom}}\), varying from 3 to 29 atoms (Figure S1).

**Alchemy.** Alchemy$^{44}$ includes 202,579 molecules with a maximum of 14 heavy atoms (including C, O, N, F, S, Cl, etc.) and \(N_{\text{atom}} = 11 \sim 38\) sampled from GDB MedChem dataset. All the 12 properties were obtained with Python-based Simulations of Chemistry Framework (PySCF)$^{97}$. All geometries were calculated with the density fitting approximation for electron repulsion integrals using the B3LYP/6-31G(2df,p). Unlike the QM9 dataset, the auxiliary basis cc-pVDZ-jkfit was used in density fitting to build the Coulomb matrix and the HF exchange matrix. The meta-Lowdin population analysis was employed to obtain the atomic charges. It should be mentioned that some of the quantum chemistry results obtained from Alchemy and QM9 datasets are somewhat different from each other$^{44}$ due to the different biased generation routines used in these two datasets despite the same computation level of B3LYP/6-31G(2df,p) was used. We find that both Alchemy and QM9 datasets give nearly identical results for Gibbs free energy with the pearson correlation coefficient of 1.0. However, the pearson correlation coefficient is just about 0.63 for the predicted dipole moments and 0.85 for HOMO results in Alchemy with the QM9-trained DeepMoleNet model (Table S2).

2.3.2 Non-equilibrium conformation datasets

**MD17.** The MD17$^{92}$ contains eight organic molecules with up to 21 atoms composed of heavy atoms like C, N, O, F. For each molecule, ab initio molecular dynamics simulation was performed to obtain the energy and forces using PBE +
vdW-TS electronic structure method. At each time step, the energy and forces together with its coordination were recorded. Here, we focus on the relative conformational energies of a certain molecule in potential energy surface (PES).

ANI-1ccx. ANN-1ccx is a high-quality and diverse data set which contains 500,000 molecules in both wide chemical and conformer spaces in the benchmark calculations at CCSD(T)/CBS level for isomerization energies, reaction energies, molecular torsion profiles, and energies and forces at non-equilibrium geometries with molecule size $N_{\text{atom}}$ ranging from 2 to 54. The ANI-1ccx dataset is intelligently selected 10% sub-sample of the ANI-1x dataset, in which the molecular conformations were derived from 57,000 distinct molecular configurations containing the C, H, N and O elements computed with DFT through an active learning algorithm with four kinds of sampling methods, namely, molecular dynamics simulations, normal mode analysis, dimer sampling, and torsion sampling.

2.3.3 Other test sets

Some typical molecules, which are of great importance in drug or material design, were selected as external data set for the transferability test (Figure S2) based on the model trained with 110,000 QM9 data.

Drug-like molecules data set. There are 24 drug molecules like aspirin, abacavir, vitamins, and drug-like molecules in this data set. All of them are natural products with pharmacological activity. They all contain rings with rotatable bonds and certain amount of hydrogen bond donors and acceptors in the spatial structure. The $N_{\text{atom}}$ of these molecules ranges from 14 to 73. In this work, 17 molecules, including vitamin B3, vitamin C, aspirin, amphetamine, vitamin B5, abiraterone, cocaine, amitriptyline, testosterone, vitamin A1, vitamin B12, progesterone, abacavir, glucosepane, cholic acid, vitamin D3, and vitamin D2 were randomly selected for the test.

MPCONF196. This dataset with the data size of 196 selects 13 acyclic and cyclic model peptides and several macrocyclic compounds, with 15 or 16 conformers (in both high- and low-energy regions) for each compound, whose conformation
energies were computed by DFT(-D3) and CCSD(T)/CBS methods. The selected macrocycles are also collected in the Cambridge Structural Database (CSD) and are further denoted by their CSD codes. In this work, 5 peptides (FGG, GGF, WG, WGG, and GFA) and 6 macrocyclic molecules (with CSD codes of POXTRD, CAMVES, COHVAW, CHPSAR, Gpd_A, and Gpd_B) were randomly selected from the MPCONF196 dataset for the transferability test of the proposed DeepMoleNet method in prediction of Gibbs free energy and HOMO properties.

Singlet fission molecules data set\textsuperscript{95}. This dataset includes 262 singlet fission molecule candidates. The dataset was screened out through a procedure of exploiting quantum chemical calculations of excitation energies, which were calibrated against experimental data. The candidate molecules were stored into different chemical families, enabling the design of further singlet fission materials using the hits as lead compounds for further exploration. For simplified, the selected 9 test molecules are also named by their CSD codes, AFZPYM, OXTPTZ, BATWUO, FEFLEK, EVAWON, LULLOT04, DIFQEP, TOSYAD, and QELNOK, in our transferability test of the Gibbs free energy and some other properties.

Oligomers data set\textsuperscript{37}. The originally reported data set selects 5 classes of oligomers, i.e., polyethylene (PE, n=28), polyacetylene (PA, n=15), polylactic acid (PLA, n=10), and alanine peptide (ala, n=10), quaternary ammonium polysulphon (bQAPS, n=3). Here, we adopt 4 kinds of oligomers with different polymerization degrees, including polyethylene (PE, n=3, 7, 14), polyacetylene (PA, n=6, 8, 15), polylactic acid (PLA, 3, 5, 10), and alanine peptide (ala, n=2, 4, 10), as our test sets.

2.4 Error analysis of predicted results

DeepMoleNet features regression tasks among 12 different electronic properties at DFT level and energy prediction at CCSD(T)/CBS calculation level. Similar to other publications, MAE and RMSE metrics were calculated for evaluating the prediction error. To further reflect the average error compared to the standard deviation of each target, we report the standardized MAE, std.\textsubscript{MAE}, of different properties. For each target, the std.\textsubscript{MAE} is defined as follows.
\[ \text{std}_{\text{MAE}} = \frac{1}{N} \sum_{i=1}^{N} \frac{|\text{Predict}_{m}^i - \text{DFT}_{m}^i|}{\sigma_m} \] (12),

where \( \text{Predict}_{m}^i \) and \( \text{DFT}_{m}^i \) are the machine learning predicted and actual DFT values for the property \( m \) (e.g., the dipole moment) of the \( i \)-th molecule, respectively. The term \( \sigma_m \) is the standard deviation from DFT result of property \( m \). \( N \) is the number of data for the test set. In some cases, for example, to reflect the relative deviation of HOMO prediction, we also calculate the relative MAE error, \( \Delta_{\text{MAE}} \), which comes from the division of MAE by the real DFT value, as shown below.

\[ \Delta_{\text{MAE}} = \left| \frac{\text{Predict}_{m}^i - \text{DFT}_{m}^i}{\text{DFT}_{m}^i} \right| \] (13).

3. Results and Discussion

3.1. Learning QM9/Alchemy molecular properties

In the first step, the 110,000 QM9 data are randomly selected as the training set, 10,000 data for validation and the rest for test set, with the molecular size, \( N_{\text{atom}} \), varying from 3 to 29 atoms (Figure S1). The transferability test is then done on the randomly split 60,000 molecules in Alchemy, which contains the computational data of the 12 electronic structure properties (dipole moment, polarizability, HOMO, LUMO, HOMO-LUMO gap, ZPVE, \( \langle R^2 \rangle \), \( U_0 \), \( U \), \( H \), \( G \), and \( C_v \)) at the same B3LYP/6-31G(2df,p) level but with different calculation workflow from those data in QM9. As mentioned above, it is not practical to use QM9 trained model to directly predict the corresponding Alchemy properties. Instead, we used the Pearson correlation coefficient to test the relative trends between the 12 properties for Alchemy molecules predicted by the model trained with QM9 and the real values reported in Alchemy data set (Table S2).

Our model can predict 12 molecular properties with satisfactory accuracy, as shown in Table 3 (for single-task) and Tables S3 (multi-task). Most of MAEs of our simultaneous predictions of 12 properties relative to DFT results are slightly lower than those of other multi-target models trained on MoleculeNet \(^\overset{84}{\text{84}}\) and multi-target model of DimeNet (Table S3). Among all the 12 properties, the present model works very well for the energy-related properties, \( U_0 \), \( U \), \( H \), and \( G \), with MAEs of 7.7 meV, 7.8 meV, 7.8 meV, and 8.6 meV, respectively. The performance of the prediction of
electronic structure properties, such as HOMO (MAE: 23.9 meV), LUMO (MAE: 22.7 meV), and HOMO-LUMO gap (MAE: 33.2 meV), is also satisfactory. For each target, we also calculated the mean standardized MAE, std. MAE. As shown in Table S4 and Table S5, the calculated std. MAE values of the present model are about 1%, approaching the chemical accuracy. It is still a big challenge to achieve the balanced and accurate predictions for all those 12 targets simultaneously. In fact, the MAEs of our multi-target model (Table S3) is even smaller than those of other single-target models for the predictions of dipole, HOMO, HOMO-LUMO gap, $U_0$, $U$, $H$, and $G$ (Table 3), which is perhaps the first report for the multi-task model to have such a performance to the best of our knowledge. The single-target prediction accuracy of 12 targets using our DeepMoleNet model is the best with small MAEs of HOMO (21.8 meV), LUMO (18.5 meV), and HOMO-LUMO gap (32.1 meV). To illustrate the statistic performance, the error bars of dipole moment, HOMO energy level, and free energy in single-target prediction and those 12 properties in multi-target task on QM9 are calculated in Table S6 and Table S7, respectively.

Table 3. MAEs of DeepMoleNet single-target training and other single-target training methods using QM9 dataset.

<table>
<thead>
<tr>
<th>Property</th>
<th>Unit</th>
<th>SchNet</th>
<th>enn-s2s</th>
<th>MEGNET</th>
<th>Cormorant</th>
<th>MGCN</th>
<th>DimeNet</th>
<th>DeepMoleNet</th>
</tr>
</thead>
<tbody>
<tr>
<td>$\mu$ D</td>
<td>0.033</td>
<td>0.03</td>
<td>0.05</td>
<td>0.13</td>
<td>0.056</td>
<td>0.0286</td>
<td>0.0178±0.0000</td>
<td></td>
</tr>
<tr>
<td>$\alpha$ $a_0$</td>
<td>0.235</td>
<td>0.318</td>
<td>0.081</td>
<td>0.092</td>
<td>0.03</td>
<td>0.0469</td>
<td>0.0475</td>
<td></td>
</tr>
<tr>
<td>$\epsilon_{\text{HOMO}}$ meV</td>
<td>41</td>
<td>43</td>
<td>43</td>
<td>36</td>
<td>42.1</td>
<td>27.8</td>
<td>21.9±0.1</td>
<td></td>
</tr>
<tr>
<td>$\epsilon_{\text{LUMO}}$ meV</td>
<td>34</td>
<td>37</td>
<td>44</td>
<td>36</td>
<td>57.4</td>
<td>19.7</td>
<td>18.5</td>
<td></td>
</tr>
<tr>
<td>$\Delta_e$ meV</td>
<td>63</td>
<td>69</td>
<td>66</td>
<td>60</td>
<td>64.2</td>
<td>34.8 a</td>
<td>32.1</td>
<td></td>
</tr>
<tr>
<td>$&lt;R^2&gt;$ $a_0^2$</td>
<td>0.073</td>
<td>1.8</td>
<td>0.302</td>
<td>0.673</td>
<td>0.11</td>
<td>0.331</td>
<td>0.115</td>
<td></td>
</tr>
<tr>
<td>ZPVE meV</td>
<td>1.7</td>
<td>1.5</td>
<td>1.43</td>
<td>1.43</td>
<td>1.12</td>
<td>1.29</td>
<td>1.22</td>
<td></td>
</tr>
<tr>
<td>$U_0$ meV</td>
<td>14</td>
<td>19</td>
<td>12</td>
<td>28</td>
<td>12.9</td>
<td>8.02</td>
<td>6.1</td>
<td></td>
</tr>
<tr>
<td>$U$ meV</td>
<td>19</td>
<td>19</td>
<td>13</td>
<td>21</td>
<td>14.4</td>
<td>7.89</td>
<td>6.1</td>
<td></td>
</tr>
<tr>
<td>$H$ meV</td>
<td>14</td>
<td>17</td>
<td>12</td>
<td>21</td>
<td>14.6</td>
<td>8.11</td>
<td>6.1</td>
<td></td>
</tr>
<tr>
<td>$G$ meV</td>
<td>14</td>
<td>19</td>
<td>12</td>
<td>20</td>
<td>16.2</td>
<td>8.98</td>
<td>7.1±0.0</td>
<td></td>
</tr>
<tr>
<td>$C_v$ cal/mol K</td>
<td>0.033</td>
<td>0.040</td>
<td>0.029</td>
<td>0.031</td>
<td>0.038</td>
<td>0.025</td>
<td>0.0241</td>
<td></td>
</tr>
</tbody>
</table>

The $\Delta_e$ is predicted simply by taking $\epsilon_{\text{LUMO}} - \epsilon_{\text{HOMO}}$ in DimeNet. 43
We apply the multi-target DeepMoleNet model, which was trained on 110,000 QM9 data, to predict the 12 properties of the molecules in the whole QM9 dataset. The prediction MAEs of the energy-related properties, $U_0$, $U$, $H$, and $G$, for all 130,000 molecules are just about 2.5-4.0 meV (0.058-0.092 kcal mol$^{-1}$). However, predictions of HOMO, LUMO, and HOMO-LUMO gap are relatively more difficult with MAEs of about 6.3-9.3 meV (Figure S3).

The difficulty in predicting HOMO and LUMO energy levels probably lies in the complexity of modulation of frontier molecular orbitals and their energy levels by different molecular topology and substitution groups. As shown in Figure S4 and Table S8, there is no evident correlation between the substituent groups and the frontier orbitals for the substituted furans at $\alpha$-position (also called 2-position in this work). According to our chemical intuition, the electron-donating or accepting ability of the substituted groups may be closely related to the upshift or down shift of the HOMO and LUMO energy levels. For this purpose, we calculated the group charge, $\rho_{\text{sub}}$, of substituent, X= H, NH$_2$, OCH$_3$, CH$_3$, OCHO by summation of the atomic charges (Table S9) in the substituted group. However, different population analysis methods, Hirshfeld$^{98}$ and Mulliken$^{99-101}$ population, give different group charges, as shown in Figure S4. Accordingly, these two different the population analysis methods exhibit different relationship between the group charges and the frontier molecular orbital energy levels. Therefore, the factors that affecting the HOMO and LUMO energy levels are rather complicated, leading to the difficulty in the accurate prediction of frontier molecular orbitals. More examples, with or without the molecular rings, are given in Table 4. It can be seen that our method is able to accurately predict molecular properties for various systems with different topology and molecular sizes (Table 4 and Table S10).

We further tested model predictivity with model trained on 110,000 data in QM9 to make predictions on molecules in Alchemy. As mentioned above, we used the Pearson correlation coefficient to make an indirect comparison between our predicted results on the randomly sampled 60,000 molecules and the original values given in Alchemy, as shown in Table S2. Good correlations were obtained for most of the
predicted molecular properties, except for the dipole moment and polarizability properties.

Table 4. The randomly selected test sets of QM9 dataset and their absolute errors of the predicted properties with the proposed deep learning method

| absolute error | CCC(=N)OC(=O)CC | CCC(=O)OCC1CC1 | COC(=O)C1(NC1C | OC1CCCC1OC=N | CC1COC2C=CCC12 |
|---------------|------------------|------------------|------------------|------------------|
| µ(D)          | 0.03             | 0.02             | 0.03             | 0.02             | 0.01             |
| α(au^-3)      | 0.07             | 0.00             | 0.06             | 0.02             | 0.00             |
| ε_HOMO(meV)   | 22.8             | 4.9              | 4.5              | 13.5             | 2.2              |
| ε_LUMO(meV)   | 6.2              | 14.5             | 29.6             | 16.8             | 15.8             |
| Δ(ε)(meV)     | 29.0             | 9.5              | 22.4             | 30.0             | 13.7             |
| U (meV)       | 2.0              | 4.9              | 3.0              | 8.8              | 18.5             |
| G (meV)       | 0.0              | 3.9              | 2.9              | 12.7             | 17.5             |
| C_v(cal/mol K)| 0.02             | 0.01             | 0.02             | 0.01             | 0.01             |

3.2. Learning non-equilibrium conformation energies

In most cases, especially when the molecules aggregate in the condensed phase or take place chemical reactions at room or even higher temperature, molecules are not always staying in their lowest energy states with the equilibrium geometry. As shown in Table 5, we adopt eight organic molecules, i.e., aspirin, benzene, ethanol, malonaldehyde, naphthalene, salicylic acid, toluene, and uracil, with up to N_atom = 21 collected in the MD17\textsuperscript{92} dataset to test the predicting ability of DeepMoleNet toward conformational energies in the non-equilibrium molecular configuration space. For each molecule, a training, validation, and test split scheme is set as 50,000, 10,000, and 10,000 molecules, respectively. A satisfactory prediction was achieved by using our DeepMoleNet model with MAE values of less than 0.08 kcal/mol for the selected molecules in MD17\textsuperscript{71, 92} dataset. The MAE values of other models using the same training numbers of 50,000 points are also listed in Table 5. It should be mentioned that different models were trained with different training set numbers as well as different approaches to get the energies. The SchNet\textsuperscript{77} was trained on energies, and
both energies and forces with 50k for training and 1k for validation, respectively. Both DTNN\textsuperscript{78} and DeepMoleNet were only trained with energies on 50,000 points with 1k for DTNN as validation set and 10k for DeepMoleNet, respectively. Some models were trained only on a small number of configurations. For example, sGDML\textsuperscript{71,92} was trained with 1,000 point (validation was part of the training set) on forces. But some other model like the DeepMD\textsuperscript{102} used a much larger training set of 95,000 points (validation was part of the training set) for the training on both energies and forces. In other words, one can get low error with 1k points used for sGDML which is comparable to errors obtained with other models trained on 50k+ points.

We choose one molecule, aspirin, of the MD17 collected molecules, to draw the PES of the AIMD sampled conformers, which were characterized by two flexible dihedral angles, $\phi$, and $\varphi$, shown in Figure 3. The AIMD potential energy surface is well reproduced by DeepMoleNet predictions, indicating the applicability of deep learning method to predict reaction barriers of chemical reactions. We carried out three independent calculations with the MAE values of 0.662, 0.665, 0.661, respectively. The MAE with the standard error of aspirin is thus presented as 0.07 ± 0.00 (Table 5). However, DeepMoleNet was trained only with energies, without force. Thus, the present model is not able to perform dynamics simulations. Further work is desired to extend the DeepMoleNet to the force predictions of non-equilibrium configurations.

<table>
<thead>
<tr>
<th>Test species (N\textsubscript{atom})</th>
<th>sGDML\textsuperscript{71,92,103} trained with force</th>
<th>SchNet\textsuperscript{77} trained with force and energy</th>
<th>DeepMD\textsuperscript{102} trained with force and energy</th>
<th>SchNet\textsuperscript{77} trained with energy</th>
<th>DTNN\textsuperscript{78} trained with energy</th>
<th>DeepMoleNet trained with energy</th>
</tr>
</thead>
<tbody>
<tr>
<td>Training size/validation size</td>
<td>1k</td>
<td>50k/1k</td>
<td>95k</td>
<td>50k/1k</td>
<td>50k/1k</td>
<td>50k/10k</td>
</tr>
<tr>
<td>Aspirin (21)</td>
<td>0.19</td>
<td>0.12</td>
<td>0.20</td>
<td>0.25</td>
<td>-</td>
<td>0.07 ± 0.00</td>
</tr>
<tr>
<td>Benzene (12)</td>
<td>0.10</td>
<td>0.07</td>
<td>0.07</td>
<td>0.08</td>
<td>0.04</td>
<td>0.02</td>
</tr>
<tr>
<td>Ethanol (9)</td>
<td>0.07</td>
<td>0.05</td>
<td>0.06</td>
<td>0.07</td>
<td>-</td>
<td>0.02</td>
</tr>
<tr>
<td>Malonaldehyde (9)</td>
<td>0.10</td>
<td>0.08</td>
<td>0.09</td>
<td>0.13</td>
<td>0.19</td>
<td>0.03</td>
</tr>
<tr>
<td>Naphthalene (18)</td>
<td>0.12</td>
<td>0.10</td>
<td>0.10</td>
<td>0.20</td>
<td>-</td>
<td>0.08</td>
</tr>
<tr>
<td>Salicylic Acid (15)</td>
<td>0.12</td>
<td>0.11</td>
<td>0.11</td>
<td>0.25</td>
<td>0.41</td>
<td>0.07</td>
</tr>
<tr>
<td>Tolunene (15)</td>
<td>0.10</td>
<td>0.09</td>
<td>0.09</td>
<td>0.16</td>
<td>0.18</td>
<td>0.05</td>
</tr>
<tr>
<td>Uracil (12)</td>
<td>0.11</td>
<td>0.10</td>
<td>0.09</td>
<td>0.14</td>
<td>-</td>
<td>0.06</td>
</tr>
</tbody>
</table>
Figure 3. (a) AIMD calculated versus (b) DeepMoleNet predicted potential energy surfaces of aspirin, whose conformations are characterized by two flexible dihedral angles, $\phi$, and $\varphi$, shown in inset.

To further test DeepMoleNet at a more accurate theoretical level like CCDS(T)/CBS, the molecules in ANN-1ccx$^{93,94}$ with molecular size, $N_{\text{atom}}$, ranging from 2 to 54 were also studied, with the prediction errors shown in Table 6. DeepMoleNet achieves comparable accuracy in comparison with ANI-1ccx results. Being limited by the computational costs and some really challenging conformations for DeepMoleNet model, we only use 280,000 data (about half of the whole ANI-1ccx dataset) to train our model. The well and poorly predicted systems by our DeepMoleNet model are presented in Figure S5, from which one can find that the MAE values of the most of the acyclic or aromatic cyclic systems are close to 0.0 kcal/mol but the nitrogen-containing five or seven membered ring (5-MR, 7-MR) heterocyclic molecules are not predicted well. Improving the predictive performance of molecules with complex chemical environments will be an important direction in our future works.

Table 6. Prediction errors, in units of kcal/mol, on ANI-1ccx dataset with different ANI models (ANI-1ccx model and ANI-1ccx-R model) and our DeepMoleNet model.
<table>
<thead>
<tr>
<th></th>
<th>ANI-1ccx$^{94,a}$</th>
<th>ANI-1ccx-R$^{94,b}$</th>
<th>DeepMoleNet$^c$</th>
</tr>
</thead>
<tbody>
<tr>
<td>MAE</td>
<td>1.8</td>
<td>2.3</td>
<td>2.3</td>
</tr>
<tr>
<td>RMSE</td>
<td>2.6</td>
<td>3.3</td>
<td>3.3</td>
</tr>
</tbody>
</table>

$a$ training based on ANI-1x 5 M data points and 500,000 ANI-1ccx data points through transfer learning and only considers conformations from 8x model ensemble.

$b$ training with ANI-1ccx 500,000 data points and only considers conformations from 8x model ensemble. The data were taken from literature.$^{94}$

$c$ training with ANI-1ccx 280,000 data points

### 3.3. Transferability test to the larger sized molecules

It is very useful to use small data samples for training to predict the data of larger sized molecules. We know that the DFT computation time increases with the scaling of $O(N_b^3)$, while the increase in prediction time for the machine learning model is modest. The scaling of DeepMoleNet is $O(N_{atom}^{-2})$. The QM9 data set only contains molecules with the size ranging from $3 \leq N_{atom} \leq 29$, so the Alchemy dataset which contains relatively larger molecules (with up to 38 atoms) was introduced for further transferability test. A series of tests were conducted with small molecules in the training dataset while big molecules for tests on these two datasets, respectively. In the first step, we sampled all molecules with $N_{atom} \leq 18$ (about 72,000 data) for training. Then we made tests on molecules with larger $N_{atom}$, as shown in Figure 4 and Figure S6. The MAE is satisfactory with the training on 18-atoms molecules and testing on 29-atom ones, although the error slightly accumulates with the size increasing. Due to the data sparsity, we could use all 72,000 data for training. Accuracy could be improved if the training data size is big enough. Even though, it is still a big challenge for realizing good transferability.

In Alchemy dataset, molecules with $N_{atom}$ less than 23 (instead of $N_{atom} \leq 18$ in QM9) were used for training to make sure that same percentage of about 50% training samples in each dataset is used as QM9 to make fair comparison of DeepMoleNet. As shown in Figure 5, a good transferability is still held on prediction of free energy $G$. 

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HOMO, LUMO, and HOMO-LUMO gap. Predictions of all 12 properties could be found in Table S11 and Table S12. It should be mentioned that the chemical space of the medium sized molecules in Alchemy is not as rich as QM9 so the prediction errors are larger than those test for QM9.

Figure 4. Data distribution of dipole moment, HOMO, and Gibbs free energy, $G$, with training set of 72,367 molecules, validation set of 8,000 molecules, and test data of 49,061 molecules in QM9 on transferability test.
Figure 5. Transferability on QM9 trained with $N_{\text{atom}} < 19$ and Alchemy with $N_{\text{atom}} < 23$ to predict larger molecules in each dataset, respectively.

In the next step, we carried out more transferability tests on many other molecules including drug molecules, peptides and macrocycles, oligomers, protein, and singlet fission molecules, using deep learning model trained with 110,000 samples of $N_{\text{atom}} \leq 29$ in QM9, as shown in Figure 6. All the selected test molecules were optimized under the B3LYP/6-31G(2df,p) level. Among those test molecules, the largest one, chignolin protein has 140 atoms, much larger than the biggest trained molecule which only consists of 29 atoms in the training set. Good correlations are indicated for all the studied chemical systems between the calculated DFT Gibbs free energies and the DeepMoleNet predicted results.
The prediction of HOMO remains a big challenge for the test molecules.
Among them, some natural products and PLA oligomers have relatively small prediction errors within 5%, as exemplified in Figure 7. In fact, the relationship between MAE and $N_{\text{atom}}$ is not evident for the prediction of electronic structure property such as the HOMO energy level. However, for some $\pi$-conjugated systems, such as, semiconducting PA oligomers ($n=15$), vitamin B12, and singlet fission molecule, TOSYAD, the prediction power of DeepMoleNet is significantly decreased (Figure S8). The difficulty in the prediction of the electron delocalized systems containing the naphthalene or anthracene ring and long $\pi$-conjugated chain in PA may be caused by the sparse in the chemical space of delocalized systems in the training set. On the other hand, DeepMoleNet uses the one-hot encoding to represent the molecular path length. In this case, if the molecules have larger sized ring structure than the trained molecules, these systems cannot be accurately described even in the input.

![Figure 7. Transferability test on HOMO energy levels of some well predicted molecules based on the model trained with 110,000 QM9 data ($N_{\text{atom}} \leq 29$)](image)

3.4. Factors that influence the performance of deep learning model
Steps of message passing. Multi-level attention mechanism has been widely used in many areas including urban computing, computer vision, and natural language processing. In this work, the T steps of message passing local attention were performed after each aggregation of different node feature levels, leading to the importance variance after each step. At the different T steps, the node is affected by the neighbors, as exemplified by the α- and β-substituted furans with T=4 and T=6, respectively, in Figure 8. The attention tends to give large weights (labelled in red color) to non-hydrogen heavy atoms at the beginning, and then gradually diffuses to the surrounding H atoms. The weights of the seemingly ‘less-important’ H atoms gradually increase, and finally climb to 1. These coefficients are learned by the neural network itself. It can be found that the setting of T=6 is sufficient to get satisfactory prediction results.
Figure 8. Illustration of multi-level attention mechanism in different message passing phase. The color indicates the relative importance. The red color corresponds to the importance of 1 and the blue color indicates the importance of 0, respectively.
ACSFs as the auxiliary teaching descriptor. For the i-th atom, ACSFs are composed of three types of radial functions, $G_{i}^{1, z_1}$, $G_{i}^{2, z_1}$, $G_{i}^{3, z_1}$, as shown in equation (14), equation (15), equation (16), and two types of angular functions, $G_{i}^{4, z_1, z_2}$ and $G_{i}^{5, z_1, z_2}$, in equation (17) and equation (18), respectively. ACSFs of the i-th atom are presented as follows, where the summation for j runs over all atoms with atomic number $z_1$ in two-body term, and for the three-body term, summations for j and k run over all atoms with atomic number $z_1$ and $z_2$, respectively.

$$G_{i}^{1, z_1} = \sum_{j} f_c(R_{ij})$$  \hspace{1cm} (14)

$$G_{i}^{2, z_1} = \sum_{j} e^{-\eta(R_{ij}-R_s)^2} f_c(R_{ij})$$  \hspace{1cm} (15)

$$G_{i}^{3, z_1} = \sum_{j} \cos(\kappa R_{ij}) f_c(R_{ij})$$  \hspace{1cm} (16)

$$G_{i}^{4, z_1, z_2} = 2^{1-\zeta} \sum_{j\neq i} \sum_{k\neq i} (1 + \lambda \cos \theta_{ijk})^{\zeta} e^{-\eta(R_{ij}^2+R_{ik}^2+R_{jk}^2)} f_c(R_{ij}) f_c(R_{ik}) f_c(R_{jk})$$  \hspace{1cm} (17)

$$G_{i}^{5, z_1, z_2} = 2^{1-\zeta} \sum_{j\neq i} \sum_{k\neq i} (1 + \lambda \cos \theta_{ijk})^{\zeta} e^{-\eta(R_{ij}^2+R_{ik}^2)} f_c(R_{ij}) f_c(R_{ik})$$  \hspace{1cm} (18)

In the above equations, the $R_{ij}$ is inter-atom distance between atoms i and j. In the three-body angular function, the angle $\theta_{ijk}$ between the three atoms (with the i-th atom in the center) is calculated by

$$\theta_{ijk} = \cos \frac{R_{ik} R_{jk}}{R_{ij} R_{jk}}$$  \hspace{1cm} (19)

The inter-atom distance related term, $f_c$, is the cutoff symmetry function, which is defined as,

$$f_c(R_{ij}) = \begin{cases} \frac{1}{2} \left[ \cos \left( \frac{R_{ij}}{R_c} \right) + 1 \right] & \text{if } R_{ij} \leq R_c \\ 0 & \text{if } R_{ij} > R_c \end{cases}$$  \hspace{1cm} (20)

where $R_c$ is a cutoff radius, which is set to be 6.0 here.

In the functions of $G_{i}^{2, z_1}$, $G_{i}^{3, z_1}$, $G_{i}^{4, z_1, z_2}$ and $G_{i}^{5, z_1, z_2}$, $\eta$, $R_s$, $\kappa$, $\zeta$, and $\lambda$ are user-defined parameters. We applied $G_{i}^{2, z_1}$ and $G_{i}^{4, z_1, z_2}$ functions in the following calculations. Two-body $G_{i}^{2, z_1}$ values are calculated by using different $(\eta, R_s)$
parameters, (1,1), (1,2), and (1,3). Three-body $G_i^{4,z_1,z_2}$ is obtained by triplet $(\eta, \zeta, \lambda)$ parameter sets of (1,1,1), (1,2,1), (1,1,-1), and (1,2,-1).

To figure out which operation is important in improving the accuracy of molecular property predictions, in Table 7 we made comparison between ‘level A’, i.e., DeepMoleNet with ACSFs as learning input descriptors with both multi-level attention and auxiliary learn ACSFs target (not as input), and those cases with multi-level attention but without the auxiliary ACSFs target (level B) and without both multi-level attention and the auxiliary ACSFs target (level C), respectively. As expected, the knowledge of attention mapping to the atoms in level A and level B could improve the prediction accuracy for HOMO and LUMO energy levels, relative to those obtained at level C without attention.

Specifically, the use of auxiliary target ACSFs in level A is crucial to get good predictions for the dipole moment, HOMO, LUMO, HOMO-LUMO gap, $U_0$, $U$, $H$, and $G$. ACSFs have wide applications in constructing potential energy surfaces (PESs) and properties prediction including catalysis, reactions, phase transition, etc. The main idea of ACSFs is to represent a chemical system's geometry with symmetrized invariant functions as input descriptors. To the best of our knowledge, it is the first attempt to set the ACSFs to be one of the prediction targets in this work. Ablation studies are further carried out to find out which information of ACSFs is helpful for the final node hidden states to learn better representation. We make comparisons among different combination of various test cases, including that only applying two-body radial functions (learn $G_i^{2,z_1}$ without using the angular functions $G_i^{4,z_1,z_2}$), only three-body angular functions (learn $G_i^{4,z_1,z_2}$ without using radial functions $G_i^{2,z_1}$), and both radial and angular functions (learn both $G_i^{2,z_1}$ and $G_i^{4,z_1,z_2}$) as the auxiliary targets, respectively. This ablation study indicates that ACSFs information could not be learned by the data-driven manner of the topological message passing. Instead, it can be obtained by predicting ACSFs. Directly using ACSFs as input in the message passing phase obtains the unsatisfactory prediction results. This suggests that in deep learning, ACSFs may be not necessarily needed to function as the input descriptors.
Instead, when ACSFs are taken as one of the prediction targets, a boost in the model performance is observed in Table 7.

One can also find from Table 7 that using either the radial functions ($G_i^{2,z_1}$) or the angular functions ($G_i^{4,z_1,z_2}$) in ACSFs could achieve better performance than that without using ACSFs. Especially for the prediction of the dipole moments, the employment of radial or angular functions is crucial to greatly improve the accuracy, implying that the structural information is important for predicting properties sensitive to structural distance. Moreover, the introduction of three-body angular functions ($G_i^{4,z_1,z_2}$) is able to give better predictions than that only using the two-body radial functions ($G_i^{2,z_1}$). For the most challenging prediction tasks of the HOMO, LUMO, and HOMO-LUMO gap, the solely use of two-body radial functions is insufficient. Fortunately, the combination of radial and three-body angular functions in level A could greatly improve the prediction accuracy for HOMO, LUMO, and HOMO-LUMO gap. This displays the importance of the many-body interactions in learning better node representations, and hence, properly predicting electronic structure properties. It is still not clear why level A scheme would work. Further exploration is going on in our laboratory. The error bars of the estimated properties in Table 7 were obtained from three independent calculations, which were listed in Table S7.
Table 7. The MAE values of QM9 predictions made by DeepMoleNet multi-target training with both multi-level attention and auxiliary target ACSFs (level A), in comparison between those cases with ACSFS as inputs (level B), without multi-level attention and auxiliary target ACSFs (level C), and without auxiliary target ACSFs, respectively.

<table>
<thead>
<tr>
<th>Property</th>
<th>Unit</th>
<th>Without using attention and ACSFs (level C)</th>
<th>Using attention</th>
<th>Using ACSFs as prediction task (level A)a</th>
</tr>
</thead>
<tbody>
<tr>
<td>µ</td>
<td>D</td>
<td>0.061</td>
<td>0.0438</td>
<td>0.0265</td>
</tr>
<tr>
<td>α</td>
<td>a₀³</td>
<td>0.109</td>
<td>0.0802</td>
<td>0.0731</td>
</tr>
<tr>
<td>E_HOMO</td>
<td>meV</td>
<td>32.1</td>
<td>25.4</td>
<td>26.1</td>
</tr>
<tr>
<td>E_LUMO</td>
<td>meV</td>
<td>31.8</td>
<td>24.7</td>
<td>24.6</td>
</tr>
<tr>
<td>Δe</td>
<td>meV</td>
<td>44.5</td>
<td>35.4</td>
<td>35.7</td>
</tr>
<tr>
<td>&lt;R²&gt;</td>
<td>a₀²</td>
<td>1.86</td>
<td>1.58</td>
<td>0.91</td>
</tr>
<tr>
<td>ZPVE</td>
<td>meV</td>
<td>3.2</td>
<td>4.0</td>
<td>2.5</td>
</tr>
<tr>
<td>U₀</td>
<td>meV</td>
<td>12.8</td>
<td>11.0</td>
<td>8.6</td>
</tr>
<tr>
<td>U</td>
<td>meV</td>
<td>12.9</td>
<td>11.1</td>
<td>8.6</td>
</tr>
<tr>
<td>H</td>
<td>meV</td>
<td>12.9</td>
<td>11.0</td>
<td>8.7</td>
</tr>
<tr>
<td>G</td>
<td>meV</td>
<td>13.4</td>
<td>11.7</td>
<td>9.7</td>
</tr>
<tr>
<td>C_v</td>
<td>cal/mol K</td>
<td>0.0459</td>
<td>0.0376</td>
<td>0.0313</td>
</tr>
</tbody>
</table>

*We investigated three cases, i.e., only applying two-body radial functions (learn 𝐺_i^{2,1} without using the angular functions 𝐺_i^{4,2,2}), only three-body angular functions (learn 𝐺_i^{4,2,1} without using radial functions 𝐺_i^{2,1}), and both radial and angular functions (learn both 𝐺_i^{2,1} and 𝐺_i^{4,2,1}) as the auxiliary targets, respectively.

As mentioned above, the using of ACSFs has an advantage in representing the local neighboring environment of an atom by using a fingerprint, which is composed of the output of several two-body and three-body functions that can be customized to detect specific structural features. Surprisingly, we found that its prediction error is closely sensitive to the number of atoms, N_atom, and the square of the number of atoms, (N_atom)², with the Pearson correlation coefficients of 0.93 and 0.87, respectively (Figure S9). In the process of message passing, every added atom will affect other atoms and increase prediction errors through many-body interactions, which should be considered in our future work.
4. Conclusion

In summary, the DeepMoleNet model provides an efficient way to generate node representation that could feel chemical environments with multi-level attention and auxiliary target ACSFs. Source codes\textsuperscript{107,108} were available on both https://github.com/Frank-LIU-520/DeepMoleNet and http://106.15.196.160:5659 for academic use. The website of DeepMoleNet and login page are illustrated in Figures S10-S14. DeepMoleNet can accurately predict electronic structure properties. Furthermore, the DeepMoleNet model can be generalized to larger sized molecules than the training molecules. It should be mentioned that QM9 is just used as the dataset to test the graph neural network model performance. We will do more calculations in our future work for making comparisons including confidence intervals, tests for significance, etc.

Indeed, the HOMO/LUMO prediction is still a challenge for the large-sized molecules. DeepMoleNet could be improved in some aspects. The present model needs to calculate the ACSFs for each atom. The combination of ACSFs knowledge with deep multi-level attention is crucial to get good accuracy. In addition, choosing the appropriate chemical descriptors as the auxiliary task requires a deep understanding of the problem. It is also a pity that when the information of atomic nodes is aggregated, the surrounding structure cannot be well touched. What is learned is an isolated node embedding information. Further improvement is anticipated to use less sample data with the goal of reaching similar prediction ability. The multi-level attention neural network will become an efficient tool to accelerate rational designs of functional molecules and chemical reactions.

ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at the attachment. Details of data distribution, standard deviation MAE, single-target results, multi-target prediction for all the QM9 dataset, multi-target prediction on different molecules, transferability test results for both QM9 and
Alchemy datasets, and instruction for using the source codes of DeepMoleNet.

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Notes

The authors declare no competing financial interest.

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