

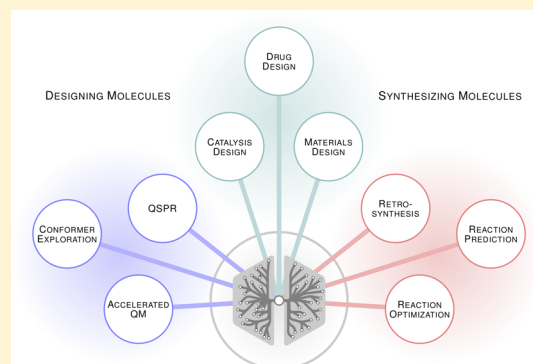
# Deep Learning in Chemistry

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**ABSTRACT:** Machine learning enables computers to address problems by learning from data. Deep learning is a type of machine learning that uses a hierarchical recombination of features to extract pertinent information and then learn the patterns represented in the data. Over the last eight years, its abilities have increasingly been applied to a wide variety of chemical challenges, from improving computational chemistry to drug and materials design and even synthesis planning. This review aims to explain the concepts of deep learning to chemists from any background and follows this with an overview of the diverse applications demonstrated in the literature. We hope that this will empower the broader chemical community to engage with this burgeoning field and foster the growing movement of deep learning accelerated chemistry.

**KEYWORDS:** Machine learning, Representation learning, Deep learning, Computational chemistry, Drug design, Materials design, Synthesis planning, Open sourcing, Quantum mechanical calculations, Cheminformatics



## INTRODUCTION

Deep learning has emerged as a dominant force within machine learning over the last 10 years through a series of demonstrations of its frequently superhuman predictive power.<sup>1–7</sup> These initial demonstrations have fostered a desire among researchers to harness its abilities to address challenges in a diverse range of areas. Chemistry stands as one of these areas, with a variety of immensely complex problems such as retrosynthesis, reaction optimization, and drug design. Historically, these have presented fierce opposition to computational approaches based on hand-coded heuristics and rules, with these approaches being met with skepticism by chemists.<sup>8–11</sup> There are strong analogies between these problems and those which deep learning has come to dominate, such as computer vision and natural language processing.<sup>12</sup> As a result of this, chemistry has seen a steady increase in the deployment of these technologies, with many demonstrating significant improvements in predictive accuracy and ability to replicate human decision making.<sup>13–24</sup>

With the prevalence deep learning is likely to achieve within chemistry, it is important that chemical researchers not familiar with the minutiae of deep learning become comfortable with how these techniques function. There have been a number of reviews covering subfields of deep learning in chemistry. Goh et al.'s<sup>14</sup> review serves as an excellent overview for theoretical chemists and has accessible explanations of the core deep learning concepts. While not strictly a review, Wu et al.'s<sup>13</sup> paper on MoleculeNet provides an extensive summary of the available descriptors and data sets as well as model comparisons. In addition to this, there are a number of broader reviews covering machine learning for drug design,<sup>25,26</sup> synthesis planning,<sup>11</sup> materials science,<sup>27</sup> quantum mechanical calculations,<sup>28</sup> and cheminformatics.<sup>29</sup> This paper seeks to adopt a central stance on

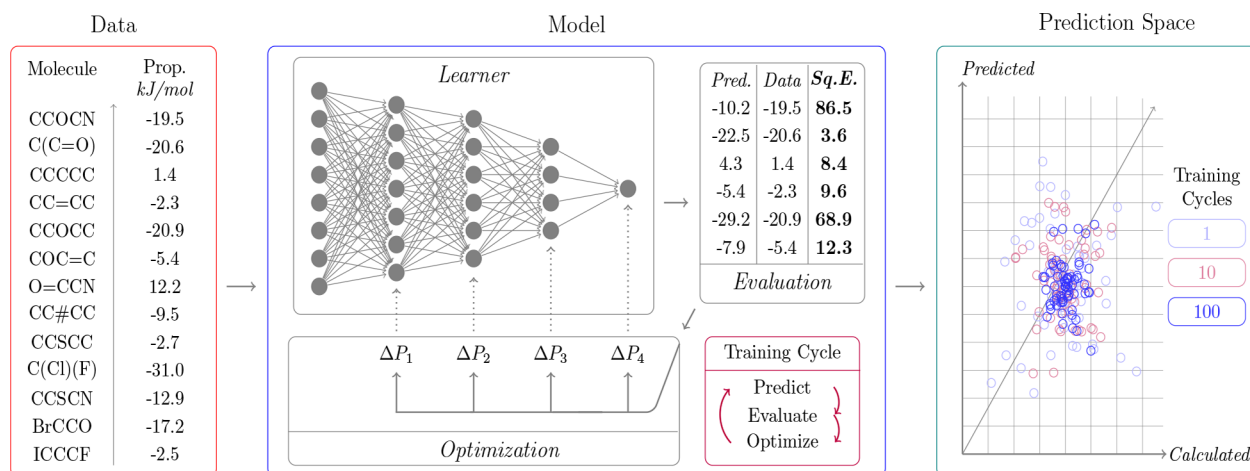
deep learning in chemistry, explaining the core ideas in the broadest possible sense, without emphasis on mathematical detail, and with reference to chemical applications. This understanding will then be used to provide a broad overview of the influences deep learning has so far had across applied and theoretical chemistry.

## THE BIG PICTURE

Machine learning is an extremely broad subfield of artificial intelligence that aims to solve the problem of computers learning from data. Representation learning is a subset of machine learning in which computational models learn internal representations of objects that inform the decisions or predictions that they make. Finally, deep learning is a subset of representation learning in which multiple layers of internal representations, initially of simple shapes such as edges, are combined to form increasingly complex objects, like faces.<sup>30</sup> Chemistry stands as an exemplar of this phenomenon, with the behavior of molecules determined not simply by atoms but their immediate grouping into functional groups, followed by interactions between these groups at increasing ranges. Ostensibly, this makes chemistry an ideal candidate for these methods. Unfortunately, molecules also supply a set of challenging problems including sampling sufficiently diverse molecules and their accompanying conformational space, effectively representing molecules, and obtaining suitably large data sets.

Understanding how these problems are being addressed requires an introduction to the methods of deep learning.

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**Figure 1.** The Big Picture of Deep Learning. The learner shown in this image is a deep feedforward network; however, this same procedure applies to a plethora of learners. The  $\Delta P$  term indicates the change to the parameters in each network layer after the input layer. The data in this image is fictitious and thus labeled simply as property.

Machine learning, and thus deep learning, at its core contains three components: the data (and its associated representation), the model that will learn to interpret the data, and a prediction space from which we draw utility. The model in deep learning (as well as other methodologies) represents an optimization cycle of three subcomponents: the learner, evaluation, and optimization. These ideas are summarized in Figure 1. Understanding chemical deep learning requires familiarity with each of these ideas and the unique challenges chemistry presents in each. The first section of this review seeks to disambiguate these topics, beginning with an exploration of data and how molecules are represented. This leads into a discussion of three of the dominant model architectures in chemical deep learning. The prediction space is then examined to explain how chemical problems must be phrased in order to make them amenable to deep learning. This section concludes with a brief overview of terms that are frequently referenced in the literature.

**The Data.** Learning cannot happen without data, and in the case of supervised learning, this data must be labeled. These labels indicate the ground truth associated with the data point, such as associating a label of “truck” with an image of a truck. In a chemical sense, the data can be a representation of a molecule with its free energy of solvation labeled or any other property. This creates one of the first big challenges of deep learning: how can enough data be obtained? The most dominant demonstrations of deep learning’s potential are in fields where data is abundant, typically where millions, if not billions, of data points can be obtained through distributed collection via social networks or even more broadly, the Internet.<sup>1,31</sup> In the case of science, the requisite volume of data only exists in certain applications. In chemistry, all levels of data are present, with extensive data available for successful reactions or ground state energies, a moderate amount of data for specific properties such as ionization energies, through to relatively small databases for properties such as free energies of solvation.<sup>32–34</sup> As a result of this need for data, chemical deep learning has formed a strong link with computational chemistry due to the latter’s capacity to generate huge volumes of data significantly faster than it could be obtained in a laboratory.<sup>33,35</sup> This presents challenges, however, due to the poorer accuracy of these calculations relative to experimentally obtained results. Lab-derived data sets are available and are the gold standard, but aside from reaction

databases, the number of data points they contain is not usually on the same order of magnitude.<sup>36</sup>

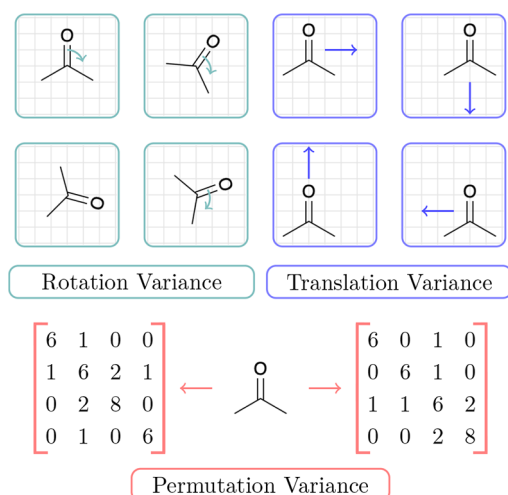
Additionally, effective assessment of deep learning models requires that the data undergoes subsequent splitting. Assessing a model on the data it was trained on leads to significant overfitting in which the model learns to reproduce that specific set of data but not the trends underlying it. To stop this “memorization” of data, it is common to test the models on data that they have not yet seen. This is typically done by dividing the data into three separate sets: training, validation, and test sets. The training set (typically 60%–80% of the data) is given to the network in its entirety, and its labels are used to adjust the network’s parameters in supervised learning. The validation set (typically 10%–20% of the data) is used to ensure that the model is not overfitting by providing a constant estimate of its performance on unseen examples. In addition to this, when training multiple models, validation data is used to identify the best performing model. Finally, the third data set, the test set, is used as the final performance evaluation of the chosen model on the remainder of the withheld data. In order to remove any bias in the partitioning of the data into these sets, *k*-fold cross validation is used, in which the data partitioning process is randomized *k* times.<sup>37</sup> Any model is highly dependent on the way in which the data is represented. Due to this, deep learning has a strong interest in the long-standing cheminformatics problem of how best to represent chemical structures for a computer.

There are three key invariances that must be captured, two of which are intuitively captured by the human visual processing. Formally, these are as follows:

- **Permutation invariance:** the representation must be unaltered by a change in the specified order of the atoms.
- **Translational invariance:** the representation must not be changed by a translation in space.
- **Rotational invariance:** the representation must be unchanged by a rotation operation.

Familiar examples of these variances are shown in Figure 2. An additional requirement for some models is a fixed size input. This is typically achieved by padding the representation with zeros for smaller molecules.

A molecular graph is a set of vertices (atoms) that are connected by edges (bonds). This can be expressed in matrix



**Figure 2.** Three key variances in common molecular descriptors that must be overcome for deep learning. The top two invariance grids show acetone undergoing rotation and translation in a fixed reference grid. Permutation invariance shows two equivalent acetone representations as atom connectivity matrices introduced by Spialter.<sup>38</sup> The atom connectivity matrix has nuclear charges listed along the diagonal, with off diagonal elements representing bonds, with associated bond order, between the diagonally located atoms that they link.

form, with an example shown in Figure 2. Originally, deep learning models utilized extended connectivity fingerprints (ECFP). These involve assigning an integer identifier to each atom and updating it to include information from neighboring atoms by expanding a circular radius that analyzes the atoms contained within. Within this circle, the atoms are sorted to achieve permutation invariance, and by compressing spatial information into integer identifiers, the two spatial invariances were also satisfied. Each of these integer identifiers are passed through a hashing function to produce a number, which, when combined with modulo arithmetic, allowed a particular index within a fixed vector to be switched to a one.<sup>39</sup> This vector has a fixed size, achieves the three invariances, but contains only zeroes and ones and is thus referred to as a bit vector. This is the basic methodology that inspired the molecular graph-based models that are described below. The idea of gathering information about an atom's local environment while preserving their invariances was retained, but critically, they encode the molecular information in a real valued vector allowing for significantly richer information to be embedded.

The Simplified Molecular Input Line Entry System (SMILES) is a classic cheminformatics representation that uses a set of ordered rules and specialized syntax to encode three-dimensional chemical structures as strings of text.<sup>40,41</sup> An additional procedure can be applied on top of this to create permutation invariance, a process known as canonicalization. The other frequently used text-based identifier, the international chemical identifier (InChI), is not regularly used in deep learning due to multiple studies finding that its more complex, numeric formulations lead to deterioration in predictive performance.<sup>42,43</sup> A reaction variant of SMILES, which contains specialized grammar to describe chemical transformations, is also frequently used in machine learning for models that operate on reaction data sets.<sup>44,45</sup>

Molecular graph inputs currently dominate due to their ability to extract higher-level features and the increase in predictive performance that comes with this. It must be noted that there are

additional representations such as point clouds<sup>46</sup> and Coulomb matrices<sup>47</sup> that are also used. Finally, regardless of representation, molecules must be entered into data sets in order to be transformed into a model input. To digitize the enormous number of structures in the literary corpus, deep learning has been used to automate the digitization of these structures.<sup>48</sup>

**The Model.** In any given deep learning framework, the model is the component that transforms the data into a prediction, classification, or action. The model relies on an interplay between its learner, evaluation, and optimization. The learner contains a set of parameters which define how each input point is converted into an output. This prediction is then quantitatively compared to the desired output via an evaluation or cost function. Finally, optimization alters the parameters of the model to decrease the difference between the predicted and the desired output for each data point. This cycle of the model making predictions, which are then evaluated and finally used to optimize the model's parameters, is bundled into a single training cycle. These ideas are summarized visually in Figure 1.

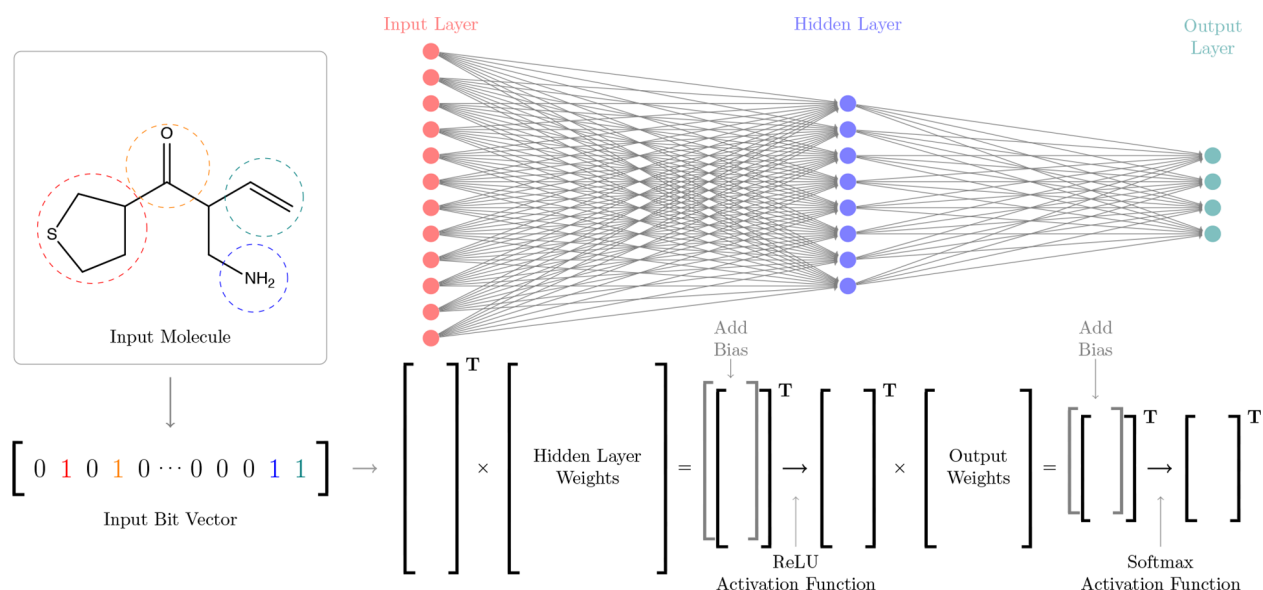
Deep learning is named for the computational depth of its learner; i.e., how many sequential layers of calculations are required. The learner is thus the defining feature of deep learning methods, with an intimate link being formed with the field of connectionism. Connectionism is focused on the development of artificial neural networks (ANNs) and their many variants. These learners are neurologically inspired systems of interconnected virtual neurons (an example network is shown as the learner in Figure 1). Due to their prominence in deep learning methods, the remainder of the model discussion focuses on variants of ANNs. A mathematical discussion is not the intent of this review; however, much of this discussion is inspired by *Deep Learning* by Goodfellow et al.<sup>30</sup> which contains an extensive and rigorous treatment of deep learning methods.

Despite the enormous diversity in the learner architecture, the evaluation and optimization procedures are dominated by a few methods. In the case of neural networks, the evaluation step is typically a simple function that assesses the learner's performance across batches, or all, of the data; two common examples are the root mean squared deviation (RMSD) or the cross-entropy cost function. The optimization typically employed for neural networks is the powerful backpropagation algorithm.<sup>49</sup> This method propagates the gradients backward from the outputs through to the inputs and using the information contained within these alters the parameters of each noninput node in a manner that lowers to deviation between the predicted and true values.<sup>49</sup> To highlight what makes the learner networks so different, three of the dominant architectures are now discussed.

**Deep Neural Network (DNN).** A deep neural network (DNN) is the prototypical deep learning architecture. DNNs contains three separate layer types, input, hidden, and output. Each layer is composed of a set of neurons, and in fully connected systems, each hidden layer neuron connects to all neurons in the previous and following layers. The "wiring" of the network (how many layers there are and how they are connected), as well as what function each neuron performs, is typically referred to as the network's topology, and the performance of the network is highly dependent on the chosen topology.

Each neuron in the input layer receives a single, real number from each data point and is thus represented as a fixed size vector. DNNs were frequently used with ECFP representations, in which a one indicates the presence of a particular





**Figure 3.** Matrix view of a typical neural network forward pass. The input molecule was chosen at random, and the bit vector is a simple structural representation that can roughly be viewed as ones indicating the presence of certain substructural feature, and zeros representing the absence. The bold T's above the vectors indicate that the transpose is used in the multiplication in order to make the operation defined.

substructural feature, which may or may not correspond to a recognizable function group, and a zero its absence.<sup>39</sup>

The neurons within the hidden and output layers have two types of trainable parameters. Every incoming connection has a scalar weight associated with it that is expressed within a matrix, and then, each neuron has its own scalar term called a bias, collected into a vector for each layer. The forward data pass is computed by multiplying the input vector with the weight matrix to produce an output vector. The bias is then added to this output vector, and it is then passed through an activation function. This function is critical as it allows the network to model nonlinear phenomena. One of the simplest and most widely used activation functions is the rectified linear unit (ReLU),<sup>50</sup> which simply maps any nonpositive number to zero and returns any positive number unchanged. This vector now becomes the input for the next layer of the network, and the process continues until the output layer is reached.

The output layer is typically either a single real number, indicating that the network is built for regression (i.e., for predicting a property such as the enthalpy of combustion), or a vector that contains the likelihood of the input being classified as certain objects, thus a classification network. In the case of classification tasks, the softmax activation function is commonly used; it converts a vector of real numbers into a probability distribution where the sum of all terms is one, and all terms are between zero and one. This allows the network to produce a distribution over the classes, indicating which is most likely. The utilization of matrix operations allows these models to leverage graphical processing units (GPUs) to massively accelerate the computation.<sup>51</sup> A summary of this matrix multiplication process is given in Figure 3.

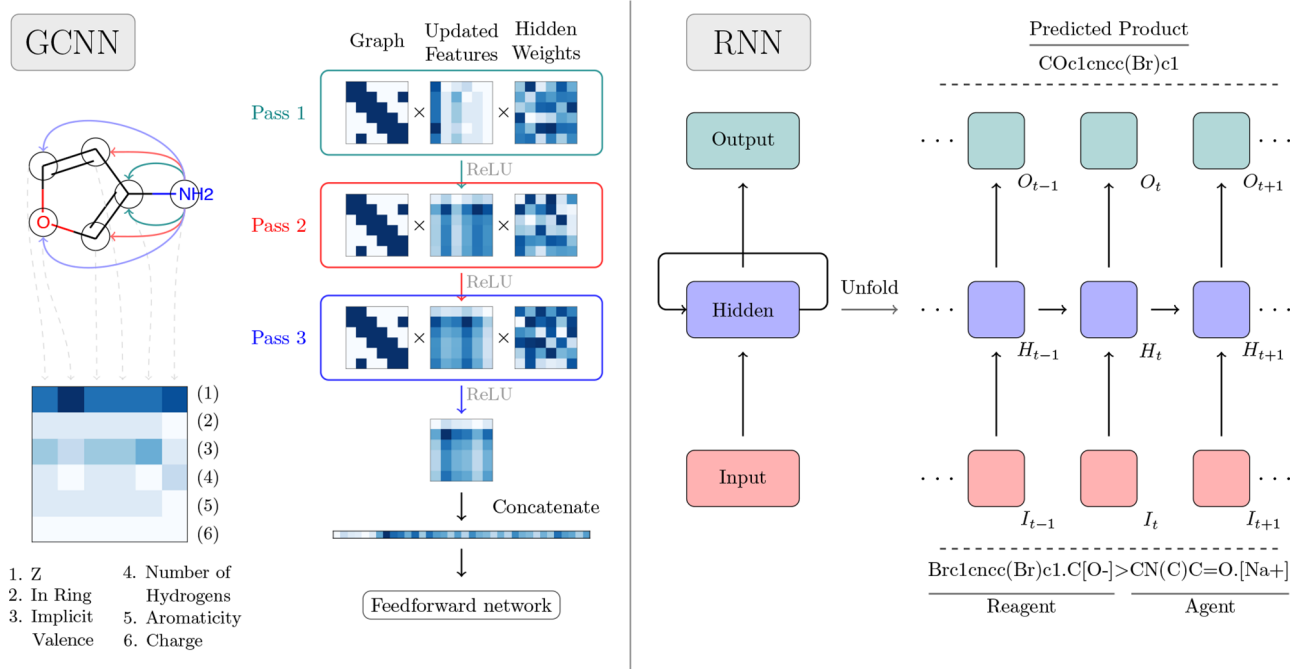
Learning in these networks involves the backpropagation algorithm, which applies the multivariate chain rule from calculus to efficiently calculate the gradients of each trainable parameter in the network, and then uses these to alter the parameters in a way that lowers the cost function. DNNs have been effective at addressing chemical problems. However, other deep learning architectures that evolved in two of AI's largest

research areas, computer vision and natural language processing, have largely superseded them.

**Graph Convolutional Neural Network (GCNN).** Computer vision is the field of research that aims to use computers to see in a manner similar to humans. Convolutional neural networks (CNNs) are networks specialized for interacting with grid-like data, such as a 2D image. As molecules are typically not represented as 2D grids, chemists have focused on a variant of this approach: the graph convolutional neural network (GCNN) on molecular graphs.

Molecular graphs confer key advantages: they bypass the conformational challenge of using 3D representations while maintaining invariance to rotation and translation due to their pairwise definition. A wide variety of molecular graph implementations have been developed in recent years,<sup>18,22,23,52–55</sup> and the MoleculeNet paper by Wu et al.<sup>13</sup> offers a concise conceptual comparison of six major variants. To facilitate the following explanation, the framework of neural message passing networks put forth by Gilmer et al.<sup>53</sup> is used.

Neural message passing networks are a chemically motivated system to understand and compare these GCNN systems. Fundamentally, this approach utilizes a convolutional layer, simply a matrix of scalar weights, to exchange information between atoms or bonds within a molecule and produce a fixed length, real-valued vector that embeds the molecular information. To begin, they generate or compute a feature vector for each atom within the molecule; this can contain information such as how many hydrogens are attached to the atom, its hybridization, whether or not it is aromatic or in a ring, etc. These feature vectors are then collected into a matrix. Additionally, they generate a graph topology matrix that specifies the connectivity of the graph, similar to Figure 2 although often without bond order or atomic number along the diagonal. In a forward convolutional pass, these three matrices are multiplied together. This allows information to be exchanged between the feature vectors of each atom with its immediate neighbors, in accordance with the connectivity specified by the topology matrix. This updates each atom's feature vector to include information about its local environment. This updated



**Figure 4.** Illustration of GCNN and RNN architectures for chemical applications. Colored arrows stemming from the amine group indicate the information transfer from the nitrogen to other heavy atoms, with the color corresponding to the convolutional pass. Light gray arrows indicate each atom's feature vector in the matrix. Importantly, properties such as atomic number (Z) are often encoded using one hot vectors, which are binary, but for spatial efficiency, the integer is used in its place. The RNN model shows a simplified “many to many” recurrent network, with the text above and below the dashed lines indicating a stylized reaction prediction system inspired by the work of Schwaller et al.<sup>61</sup> This system takes in reagent and agent SMILES and predicts the variable length product string; however, the LSTM architecture they used is significantly more complex than the one shown above.

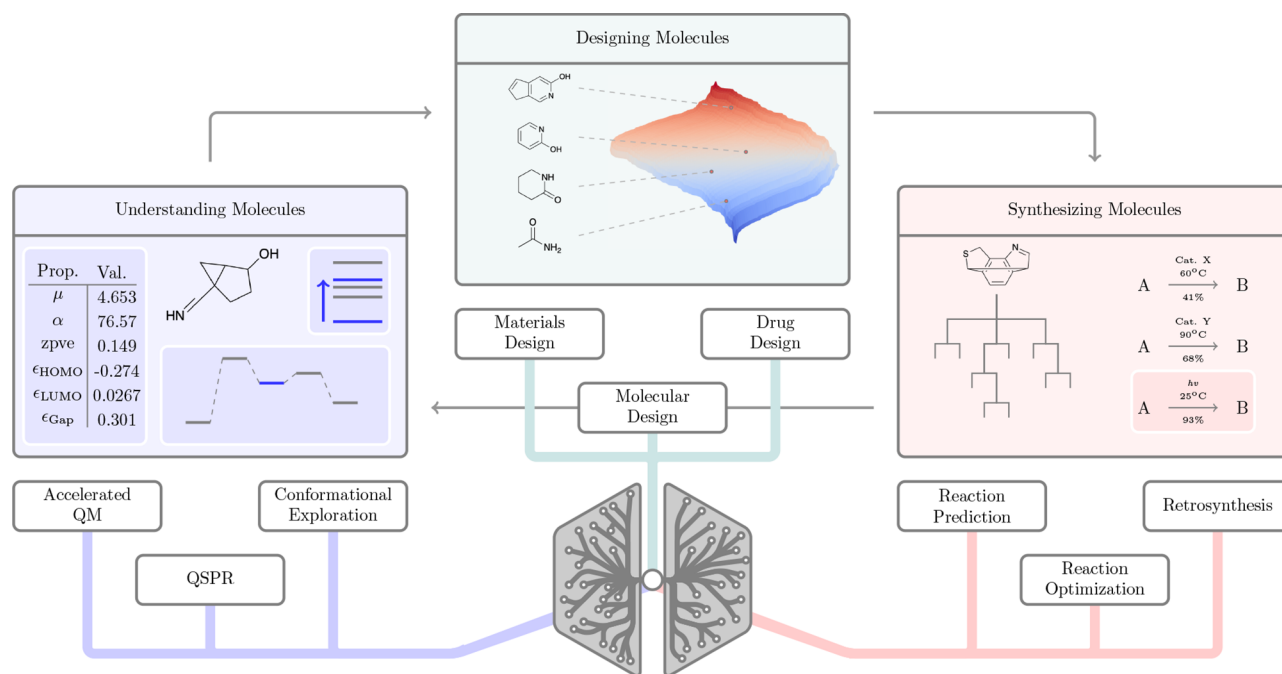
feature vector matrix is then passed through an activation function (i.e., ReLU) and can then be iteratively updated by using it as the feature matrix in another convolutional pass. This propagates information throughout the molecule. Finally, these atom feature vectors are either summed or concatenated to give a unique, learned representation of the molecule as a real valued vector (Figure 4). Alternative approaches to generating this learned representation have been put forth, such as using traditional computer vision CNNs on 2D grid images<sup>56</sup> of molecules. However, molecular graphs remain the dominant paradigm.

The learned representation in vector form is referred to as a representation in latent space and is then used as the input for a traditional fully connected DNN to finally make the classification or prediction. This process of learning its own molecular representation is the cause of it being in the broader class of representation learning methods. Backpropagation is once again used to train these networks by propagating gradients backward and determining how to change the convolution matrix weights and the parameters in the DNN.

**Recurrent Neural Networks (RNNs).** Recurrent neural networks (RNNs), introduced by Hopfield<sup>57</sup> in 1982, are specialized for dealing with sequences of arbitrary length. This makes them ideally suited to handling textual representation of chemical information, such as SMILES.<sup>40</sup> The critical difference is that in the previous architectures each data input is distinct, while in an RNN each input will influence the next one. An illustrative example is viewing any particular input, such as a SMILES string, as time series data. The presence of a carbon atom at one moment in time influences what the next character is likely to be. This is expressed in the architecture by feeding the

output of the hidden layer for that carbon into the hidden layer of the next atom. As a more complex illustration, this process can be used to model reactions by utilizing the SMILES reaction strings to encode the information and train the network to predict the product (Figure 4). The feeding of one hidden state into the next gives the system a recursive relationship within the hidden layer, but it can be viewed as directional by “unfolding” the network to form of an unfolded, acyclic network graph. By doing this, it maintains a history of all previous inputs, and they influence its prediction at a later time. The network can then be trained using a recursive form of backpropagation.<sup>58</sup> This is the simplest RNN, but more sophisticated and powerful variants such as neural Turing machines<sup>59</sup> and long–short-term memory networks (LSTM)<sup>60</sup> that incorporate memory into the network are the current leaders. This ability to use previous information has led to their dominance in sequence-based tasks such as machine translation, as previous words define the context and thus what the next word is likely to be.

**The Prediction Space.** The prediction space is the set of all possible outputs for the network. More intuitively, it can be thought of as the utility of the network or the question that the network can produce an output for. As discussed above, supervised learning requires labeled data that allows the model to iteratively improve its predictive performance. This model relies on a quantitative error assessment by the evaluation component, and thus, each deep learning problem needs to be framed in such a way that it can be quantitatively evaluated. This creates a significant challenge in chemistry, as questions such as “what is the best synthetic route?” require systematic analysis to produce a question that can be numerically evaluated and thus produce quantitatively labeled data. In the broader context of



**Figure 5.** Deep learning influence on the idealized chemical workflow. Illustrative examples of each task are shown in the dialogue boxed with arrows indicating the closed cycle that is contained within the framework. The property values in the blue panel were obtained from the QM9 data set for a randomly chosen molecule.<sup>33</sup>

artificial intelligence, this means that these systems are weak AI, capable of solving only a single, extremely narrow task and not capable of meaningfully answering even slight deviations from the question it was trained on.

**Commonly Used Terms.** Before concluding this section, a brief explanation of commonly used ideas and terms is provided. Each term is linked to seminal papers and, where appropriate, accompanied by an example of its application in chemistry.

- **Transfer Learning:** Transfer learning involves using a network that has been trained on a related task and then tweaking its parameters to adapt to a new task, often with less data.<sup>62</sup> It has been used to adapt a model trained on DFT to a smaller database of higher fidelity calculations by Smith et al.<sup>63</sup>
- **Multitask learning:** This involves training a model on multiple prediction tasks at the same time to decrease the likelihood of overfitting.<sup>64</sup> It has been used to increase predictive performance for toxicity or bioactivity prediction.<sup>65</sup>
- **One Shot Learning:** A technique used to overcome applications with extremely limited data that uses networks to compress inputs into a continuous latent space and then compares the representation in this space to a larger, trained latent space.<sup>66</sup> It has been used in chemistry for low-data drug discovery.<sup>67</sup>
- **Autoencoders:** A network architecture used to compress an input to a real valued vector, commonly referred to as the latent space. A decoder network then takes this vector as its input and tries to reproduce the original input data.<sup>68</sup> It has been used to design molecules by training the latent space to reflect a particular property and then navigating it.<sup>43</sup>
- **Generative Adversarial Networks (GANs):** GANs utilize two networks in a competitive scheme. One network has to generate data, and another has to

determine if a particular data point is a fake generated by the network or a real one from the data set. By competing with one another, the generating network learns to create high quality imitations of the data set.<sup>69</sup> It has been utilized for the inverse molecular design problem.<sup>70</sup>

- **Data Augmentation:** This involves expanding a data set by creating new training examples through reasonable manipulations of the data. One of the simplest demonstrations of this is rotating images in a data set but maintaining the same label in a way that is obvious to humans; i.e., a car is still a car at different angles.<sup>71</sup> This has been used with SMILES to enumerate the different potential orderings and increase the predictive performance.<sup>72</sup>
- **Reinforcement Learning:** This is when the model learns iteratively through trial and error by making its cost function measure its progress toward a particular goal.<sup>73</sup> It has been used to train a model to optimize reactions.<sup>74</sup>
- **Supervised Learning:** Supervised learning involves giving the model a labeled data set, effectively telling it what it needs to learn. While this is currently the dominant learning paradigm in machine learning, it is not representative of how humans tend to learn.
- **Unsupervised Learning:** Unsupervised learning is learning in which the model is not told what to reproduce and instead tries to separate the data into its underlying clusters. Algorithms such as *k*-mean clustering fall into this category, and it is much closer to how humans learn.<sup>75</sup>

## DEEP LEARNING APPLICATIONS

This section reviews the multiple areas of chemistry that deep learning has thus far impacted, presenting examples in each that highlight particular achievements. To create a logical narrative, this discussion follows an idealized chemical workflow. To build

a molecule with a particular property would first require developing methods to accurately correlate any given structure to the property. These can then be used to intelligently design a molecule that maximizes the desired property. The final step is to design an efficient synthesis from readily available starting materials (Figure 5). This creates a closed feedback cycle in which the synthesized molecule can be experimentally analyzed, and this information can then improve the models that link molecules to properties. Deep learning has influenced every stage of this workflow, beginning with understanding molecules.

**Accelerated Computational Models.** Computational modeling in chemistry seeks to use physics-based calculations to determine the properties and behavior of a given molecular system. There are two distinct ways which deep learning can be used within this space. The first is to integrate the deep learning method with physics style approaches to alleviate computational bottlenecks. The second is directly predicting properties from molecular structures, thereby bypassing physical laws altogether.

Integrating deep learning methodologies with physics-based approaches involves training the network to predict a key component of the overall calculation. These include using the deep learning model to predict potential energy surfaces,<sup>52,76,77</sup> force fields,<sup>78</sup> add corrections to ab initio calculations,<sup>79</sup> and to bypass expensive stages in both density functional and wave functional methods.<sup>80,81</sup> There is an excellent review and tutorial on using neural networks for the prediction of potential energy surfaces by Behler.<sup>82,83</sup> Many of these methods adapt a method introduced by Behler and Parrinello<sup>84</sup> in 2007 that determines the energy of the system by summing the energetic contribution of each atom. This method transforms the Cartesian coordinates of a molecule using radial symmetry functions, using this to then describe the atom's local environment. This transformed representation is then passed through a neural network that predicts the contribution of this atom to the total energy. This general method of using functions to capture an atom's local environment, then predicting its energy through a network and finally summing these contributions, has been refined in a variety of ways. Notable work in the field includes that of Schütt et al.<sup>24</sup> which produced size extensive predictions with an average error of 1 kcal/mol, and the work of Smith et al.<sup>52</sup> which produced errors below 1 kcal/mol and generalization to larger molecules. Schütt et al.'s<sup>85</sup> work has been further refined and developed into an open source software package (SchNetPack) that can be used to predict properties.

The advantages of this approach are that it is more flexible than mapping a structure to a property, and it is more interpretable due to its physical basis. The difficulty is that, as there are typically still physics-based calculations involved, such methods cannot achieve the same speed as those that map purely from a structure to a particular property. It is important to note that there is a large literature base for using kernel ridge regression as the ML method. This approach has achieved excellent results but is not a deep learning method and thus is outside the scope of this review. For an overview of these methods, the reader is referred to von Lilienfeld's excellent review.<sup>86</sup>

**Quantitative Structure Property/Activity Relationships.** The alternative approach to deep learning in computational chemistry is training a direct map from a simple representation of the molecule through to the desired property. This is a diverse field of research that can broadly be captured under the two fields of quantitative structure property relationship (QSPR) and quantitative structure activity relation-

ships (QSAR). Broadly speaking, QSPR seeks to predict properties of molecular systems, such as thermochemistry, while QSAR seeks to predict the activity of that molecule within a broader context, such as toxicity within biological systems. The goal of these methods is to maximize accuracy of prediction, with chemical accuracy for QSPR commonly being set to 1 kcal/mol or approximately 4 kJ/mol.<sup>87,88</sup> The properties that can be predicted are entirely determined by the available training data, and there are many databases available. There are summaries of available databases in both the review by Butler et al.<sup>27</sup> and the MoleculeNet paper by Wu et al.<sup>13</sup> Typically, for properties that can be readily computed, such as ground state energies, ionization energies, or dipole moments, computational data sets are the norm. These are typically computed with a DFT method in order to maximize speed and allow for as much data as possible to be generated. Some of the most commonly utilized are QM9,<sup>33</sup> ANI-1,<sup>35</sup> and the Materials Project.<sup>89</sup> Properties that are difficult or currently impossible to compute accurately, such as toxicity, free energies of solvation, biological activity, or binding affinities, rely instead on experimental data sets that typically contain significantly fewer entries due to the challenge in obtaining them. Frequently used data sets include ChEMBL,<sup>90</sup> PubChem,<sup>91</sup> and FreeSolv.<sup>34</sup>

For this type of problem, DNNs were the most widely used network architecture for the first half of this decade. They have been used to effectively predict electronic properties,<sup>19,87,92</sup> bioactivity,<sup>21,93–95</sup> toxicity,<sup>15,96,97</sup> and reactivity,<sup>92</sup> as well as other physical properties.<sup>98</sup> Multitask networks are also frequently used due to the increase in predictive performance, as well as increased robustness to overfitting.<sup>15,21,93,94</sup> RNNs have been more widely used as the generative networks that produce novel molecules which are discussed later. For predictive purposes, however, they have utilized both graph-type input structures similar to GCNNs to predict aqueous solubility<sup>18</sup> and drug toxicities,<sup>99</sup> as well as the more traditional text-based inputs of SMILES for general property prediction.<sup>100,101</sup>

In almost all cases, however, GCNNs and their many variants have demonstrably better predictive performance than either of the other two classes of methods. Due to the focus on improving network architecture, convolutional models are often tested against a variety of benchmarks. However, there has been a particular push to improve the predictions of electronic properties in order to ease the computational stress imparted by physics-based calculations.<sup>56,102,103</sup> In addition to this, GCNNs have shown dominance in predicting bioactivity,<sup>104</sup> polymer property predictions,<sup>105</sup> and physical properties.<sup>55,106</sup> Work to increase their predictive abilities is ongoing, but errors below 1 kcal/mol are routinely achieved. The accuracy of these methods brings into question the validity of the training data, particularly the accuracy of the labels, as well as potential bias in the data. DFT is known to have large errors,<sup>107,108</sup> while the gold standard methods such as coupled cluster with singles, doubles, and perturbative triples (CCSD(T)) are currently prohibitively expensive for data sets of this size.<sup>109</sup> In order to overcome this deficit, transfer learning has been utilized to fine-tune these networks on smaller data sets of calculations performed at significantly higher levels of theory, such as CCSD(T).<sup>63,110</sup> Additionally, bias in chemical data sets is a well-known problem.<sup>111,112</sup> While there has been recent work to intelligently design them using deep learning,<sup>113</sup> genetic algorithms,<sup>114</sup> or techniques such as query by committee,<sup>115</sup> the large data sets required for chemical deep learning are largely restricted to small



molecules containing only carbon, nitrogen, oxygen, fluorine, and hydrogen. As the coverage of chemical space expands, it is critical that the data sets are intelligently designed to maximize coverage of the rapidly expanding combinatorial space.

The final topic to address is interpretability. Deep learning has a reputation for being a “black box”, as it is almost impossible to understand why the network made the decision that it did.<sup>116</sup> Recent work has attempted to overcome this in chemical deep learning by cleverly designing the architectures to allow for extraction of chemical insights from its decision making. In recent work from Goh et al.,<sup>117</sup> by changing the information available to the network in their descriptor, they were able to infer that the network was learning a different approach to solve different chemical prediction challenges. Schütt et al.,<sup>102</sup> on the other hand, demonstrate not how the network is making decisions but rather that its predictions align with an understanding of chemical ideas such as aromaticity.

**Conformational Exploration.** Regardless of how deep learning influences chemical property mappings, effective exploration of chemical space involves navigating not only the species space but also the conformational space of those species. Conformational screening is an immense challenge in chemistry, as with each new atom, multiple additional local minima appear on the potential energy surface. The aforementioned neural network potentials offer a rapid way to explore the conformational space of a molecule. The leading potential at the moment is the ANI-1 potential that achieved errors below 1 kcal/mol and is trained using off-equilibrium geometries.<sup>52</sup> The data set it was trained on contains approximately 20 million energies of ~57,000 molecules in different stances.<sup>35</sup>

The inverse of conformational screening is to develop a system that can generate equilibrium conformers for a given molecule. This challenge has been undertaken by Gebauer et al.<sup>118</sup> which demonstrated deep learning’s ability to generate equilibrium conformers. This method is an adaption of SchNet architecture developed by Schütt et al.<sup>103</sup> that was able to regenerate molecular geometries with a root mean squared deviation of approximately 0.4 Å. Additionally, a novel, but not as rigorously tested, method was introduced by Thomas et al.<sup>46</sup> in which 3D point clouds were used to regenerate molecule geometries. This work did not place the same emphasis on minima structures but was able to achieve very low errors of approximately 0.15 Å. This field of research is still very young but holds immense potential to minimize the conformational screening bottleneck.

## ■ MOLECULAR DESIGN

The second stage of the idealized workflow is the problem of molecular design. This problem, sometimes referred to as inverse QSPR, has a history of machine learning applications including Bayesian optimization<sup>119</sup> and genetic algorithms.<sup>120</sup> Recent years have seen the application of generative deep learning models to design molecules. One of the seminal demonstrations of this method is the work of Gómez-Bombarelli et al.<sup>43</sup> which used an autoencoder with a latent space that was optimized by an additional network to reflect a particular property. This “landscape” can then be explored to identify candidate molecules that maximize the property. There are many other approaches that also use autoencoders,<sup>121–124</sup> generative adversarial networks (GANs),<sup>70</sup> or reinforcement learning agents<sup>125–128</sup> to navigate chemical space structured around a particular property. Finally, RNNs have also been used for molecular library generation by an adaptation of their text

generation capabilities.<sup>129</sup> An excellent review of molecular design is provided by Sanchez-Lengeling and Aspuru-Guzik.<sup>130</sup>

General molecular design is seeing a surge of activity; however, there are two special classes of molecules that deserve particular attention: materials and drugs. These are arguably the two most challenging molecule classes to design and optimize but also offer the greatest potential benefits. Therefore, they have motivated significant research efforts with deep learning.

**Materials Design.** Many modern technologies such as batteries, aerospace, and renewable energy rely on advanced materials. Deep learning has only recently begun to influence the field, but there has been a rapid growth in applications in the past few years. The distinction between discrete small molecules and crystalline structures has led to a separate set of convolutional descriptors that seek to capture the crystalline structure. Crystal graph convolutional neural networks (CGCNNs) as introduced by Xie and Grossman<sup>131</sup> show much potential in this field. There has been a push, however, to reconcile the representation systems for these two classes of molecules. SchNet<sup>103</sup> has been demonstrated on both, and MEGNet by Chen et al.<sup>132</sup> has been developed for this specific purpose.

GCNNs, as well as the CGCNN variant, have been used to predict the properties of bulk materials,<sup>133</sup> predict thermoelectric properties,<sup>134</sup> optimize polymer properties,<sup>135</sup> and explore chemical materials space.<sup>136</sup> These applications are still young; however, it has moved beyond predictive models as demonstrated in work by Li et al.<sup>135</sup> in which they successfully used reinforcement learning to train an agent to experimentally control polymer weight distributions and thus the polymer’s properties. Additionally, the exploration of chemical materials space by Xie and Grossman<sup>136</sup> demonstrated the potential of these methods to uncover previously undetected pockets of materials space. Beyond the properties of materials, work has been done to optimize their synthesis parameters<sup>137</sup> and perform defect detection.<sup>138</sup> Finally, a deep learning method that utilizes tensor networks, similarly to Schütt et al.,<sup>24</sup> demonstrated generative design of chiral metamaterials.<sup>139</sup> Most of these applications remain theoretical in nature, and effectively incorporating them with an experimental workflow, such as in the polymer optimization workflow of Li et al.,<sup>135</sup> is a key next step to determine their efficacy.

One key subfield of materials design is catalysis design. Machine learning has seen increased use in catalytic research;<sup>140,141</sup> however, deep learning has seen limited application in this field due to the limited data available, the unique nature of each catalytic process, and the difficulty of representing multimolecular systems. The applications of deep learning within catalyst design largely center around using neural network potentials to model the catalytic system. Recent examples of this include Shakouri et al.’s<sup>142</sup> work to model nitrogen gas on a ruthenium surface and the optimization of platinum clusters by Zhai and Alexandrova.<sup>143</sup> Extending this work beyond using neural network potentials will likely require increased data gathering efforts, as well as the development of newer descriptors to describe interacting, multimolecular systems.

**Drug Design.** Drug design is arguably one of chemistry’s most important applications. Fundamentally, it involves identifying molecules that achieve a particular biological function with maximum efficacy. These can either be obtained from natural sources or built from the ground up. In either case, the goal typically starts with one molecule, or a set of molecules, and the challenge is to optimize its properties to improve



potency, improve specificity, decrease side effects, and decrease production costs. There are a number of reviews on deep learning's impact on this field as it is of great interest to the community.<sup>25,144,145</sup>

The generative models in drug design follow the same trends as general molecular design, with autoencoders,<sup>146</sup> GANs,<sup>147</sup> and reinforcement learning<sup>148</sup> all being used to try and generate potent drug molecules. In addition to these, there are some novel approaches to drug development that include predicting anticancer drug synergy<sup>149</sup> and developing a benchmarking for generative models in drug design.<sup>150</sup> Drug design approaches struggle from limited data, possibly more so than any other field due to the expense of obtaining it. Work by Altae-Tran et al.<sup>67</sup> utilized one shot learning to address this deficiency and make informed predictions about drug candidates with limited data. Finally, while not a molecule-optimizing generative system, work by Segler et al.<sup>151</sup> developed methods to develop focused libraries of drug candidates for screening using RNNs.

## ■ SYNTHESIS PLANNING

Synthesis planning is the final stage in this idealized workflow. It can be simplified into three separate components: Retrosynthesis, in which the product is known, is where it is broken down into a series of simpler starting materials from which it can be made. Reaction prediction, in which reagents are known, is where the dominant product must be determined. Finally, reaction optimization involves taking a reaction with known reagents and products and trying to maximize the yield or efficiency of this process. One important distinction to note here is that reaction optimization and reaction prediction both have well-established computational approaches, kinetic models, and quantum calculations, respectively. Both of these can, however, be expensive and in the case of quantum calculations enormously so.

Computational retrosynthesis, on the other hand, has a long and turbulent history. The original retrosynthesis program was Pensak's and Corey's<sup>8</sup> work on the LHASA software. From this point, there have been a multitude of assistive software packages.<sup>152–154</sup> The beginning of the 21st century saw a loss of interest in this field due to a variety of factors, but it is largely attributed to a widespread belief that computers could not capture the art of synthesis. This field has had a second wind with the advent of deep learning, with the models beginning to challenge the notion of computational inferiority in synthesis planning.<sup>10</sup>

**Retrosynthesis.** The great challenge of retrosynthesis is the exponential scaling of possible moves in synthetic space from any point. This is a property it shares with traditional board games such as Chess or Go. Formally, this can be expressed as a tree search, where the branching factor is how many possible steps you can take from a particular point. The depth is how many steps it takes to reach the desired position. Compared to the aforementioned games, retrosynthesis has a significantly greater branching factor but lower depth.<sup>155</sup> Retrosynthesis may present a far greater challenge due to the immense difficulties in knowing a priori whether a reaction will be successful and produce the desired material, whereas Chess and Go have a perfectly defined set of possible moves. However, these games represent a good starting point to consider the problem, and fortunately, both have succumbed to artificial intelligence approaches. It is not surprising then that one of the dominant displays of retrosynthetic AI was heavily inspired by AlphaGo,

the seminal AI system DeepMind developed to achieve superhuman Go playing ability.<sup>3</sup>

Work by Segler et al.<sup>16</sup> adapted the AlphaGo methodology (Monte Carlo Tree Search with deep neural network policy) to design a state of the art retrosynthetic AI. This system was trained on over 12 million reactions from the Reaxys<sup>156</sup> database and produced human-accepted synthesis routes. Assessing synthesis plans is a thorny challenge, and in order to do this, they performed a double-blind study in which graduate chemists were shown the machine's synthetic plan and the original, literature plan. There was no statistically significant difference in their preferences, thus giving a preliminary indication that its synthetic routes are "human level". It is also possible, however, to argue that the graduate chemists do not yet have the necessary expertise to distinguish the human route. Thus, determining when computers achieve human ability in synthesis planning is a decision that can only be made by the entire field. While this method showed great potential, there are other avenues of research such as the use of RNNs in an encoder/decoder setup to perform retrosynthetic analysis of small molecules.<sup>157</sup>

Computational retrosynthesis is making enormous strides; however, many problems persist. First, planning a retrosynthesis that looks valid and experimentally verifying its predictions are different challenges, and until these methods are rigorously tested, it is unknown whether or not they are useful to chemists. This challenge would likely benefit from a user-friendly software package in order to get chemists' feedback on the computer-generated syntheses. These are beginning to appear with an example being the ASKCOS software developed by the Machine Learning for Pharmaceutical Discovery and Synthesis Consortium.<sup>158</sup>

**Reaction Prediction.** Reaction prediction is the process of taking a set of known reagents and conditions and predicting what products will form; as such, it typically requires greater exploration into uncharted chemical space. Current methods to perform this, such as quantum calculations, are exceedingly expensive and thus limited to smaller molecules. Deep learning methods represent an opportunity to alleviate this computational expense and free up time of trained computational chemists.

Reaction prediction exemplifies the challenge of predicting outliers due to the frequent need to predict outside of the training space. As a result of this, the majority of reaction prediction machine learning methods either integrate the model with a physics-based scheme or apply reaction templates.<sup>159</sup> One of the early works that applied deep learning to reaction prediction involved DNNs with molecular fingerprints to predict what product would form.<sup>44</sup> Additional work has utilized RNN variants,<sup>61</sup> as well as more specialized architectures such as neural machine translation<sup>160,161</sup> and Siamese architectures (which take two identical networks given different inputs and determine the similarity between them).<sup>17</sup> One of the striking challenges for this field is the immense literature bias toward successful reactions. Recently Coley et al.<sup>45</sup> presented a clever approach to overcome this by recognizing that a successful reaction implicitly defines a large number of unsuccessful reactions that can be added to the database. This was performed by identifying high yielding reactions and generating viable alternative products that are thus not formed in high yield. These can then be added to the data set to augment it with negative examples. The current state of the art that also stresses interpretability uses a GCNN to predict reaction outcomes in a manner similar to human intuition.<sup>162</sup>

Due to deep learning's relatively new arrival to reaction prediction, there is a history of traditional machine learning methods for reaction prediction that is reviewed by Coley et al.<sup>11</sup> Current developments are reaching a level that is competitive with humans. With further advancements in predictive ability and transitioning it into user-friendly software, this is likely to become a key addition to the chemical toolset.

**Reaction Optimization.** Reaction optimization involves tuning the conditions of a reaction to increase its efficiency. This is often performed via kinetic models or experimentally through the use of flow chemistry or high throughput combinatorial chemistry. Despite the maturity of these methods, there is scope for a system which can rapidly produce idealized synthetic conditions given a molecule and reaction type. Deep learning has the potential to fill this niche, and research has begun to adapt it to this challenge.

The potential of this approach was demonstrated by Zhou et al.<sup>74</sup> in which an RNN variant learned to optimize the conditions of reactions. Their model used an RNN that learned to evolve the conditions of a reaction toward an optimized state. It was trained on simulated reactions and then outperformed other software-based approaches for multiple experimental reaction setups. It is important to acknowledge here that due to limited availability of data, and the need to flexibly update the model, deep learning methods may not be the best choice here; instead, a method that uses alternative machine learning methodologies such as random forests has been demonstrated to be a potent alternative.<sup>163</sup>

## FUTURE DIRECTIONS

To summarize, deep learning is a subfield of machine learning that uses sequential layers to extract higher level features and use them to learn the patterns present in a data set to predict future behavior. Supervised learning requires large volumes of labeled data and a quantitatively assessable goal or question. With this, a model uses an interplay of a predictive learner, evaluation, and optimization, in the form of a training cycle to iteratively improve its performance until it begins to overfit the training set, at which point training stops and the model is evaluated.

The past decade has seen explosive growth in the application of these methods across chemistry. Through its applications, deep learning shows promise of being a game changer within chemistry. This review has demonstrated that deep learning has and will continue to impact every stage of the idealized chemistry workflow. Realization of its potential will require a concerted effort to address the major challenges deep learning still faces, many of which have been discussed throughout this review. The three main challenges that must be addressed to maximize the potential of this technique within chemistry are (1) obtaining large amounts of high-quality data, (2) developing a standardized framework, and (3) effectively integrating with the broader chemistry community.

The first two challenges will be immensely benefitted by increased collaboration and, in particular, continued open sourcing. The push for open sourcing has increased, and there is strong evidence of it occurring within deep learning particularly through software packages such as DeepChem,<sup>13</sup> TensorMol,<sup>164</sup> ANI,<sup>52</sup> SchNetPack,<sup>85</sup> and chemprop.<sup>165</sup> Addressing the problem of high-quality data also relies on continued advancements in physics-based computational chemistry and the accompanying software packages.<sup>166,167</sup>

The final challenge requires concerted action from specialists and the broader community. Open sourcing software packages is

a step in the right direction, but the chemical community has a long history of resisting assistive software either due to poor usability or unreliable software performance. The latter is demonstrably addressed by these powerful methods, but the former requires conscious development of usable software packages with feedback from the community. These methods are built to empower chemists first and foremost, and that must be a priority as this field matures.

This review hopes to serve as a gateway to this burgeoning field and encourage chemists, regardless of their specialization, to consider how deep learning could be applied to their work. The following are a set of guidelines to assist in the initial application of these methods:

- Python has become the coding language of choice for deep learning and finding someone proficient in it is invaluable.
- Deep learning requires large volumes of data to outperform traditional machine learning methods. Unless transfer learning is an option, a few thousand data points is a minimum.
- To begin with, employ the open source software packages referenced above with default settings to get a baseline.
- From this baseline, adapt the network architecture using techniques presented in the literature referenced in this review to try and improve performance.
- Utilize the wealth of informative online courses and user-friendly software packages<sup>168,169</sup> provided by the deep learning community to aid in learning these techniques.

Deep learning's contributions to chemistry to date demonstrate that it has a bright future within chemistry, but through effective collaboration between specialists and the broader community, it has the potential to offer a revolution.

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## ABBREVIATIONS

ANNs = Artificial Neural Networks

CCSD(T) = Coupled Cluster Singles Doubles with Perturbative Triples

CGCNN = Crystal Graph Convolutional Neural Networks

CNN = Convolutional Neural Network

DFT = Density Functional Theory

DNN = Deep Normal Network

ECFP = Extended Connectivity Fingerprint  
GANs = Generative Adversarial Networks  
GCNN = Graph Convolutional Neural Network  
GPU = Graphical Processing Unit  
InChI = International Chemical Identifier  
LSTM = Long Short-Term Memory  
QSAR = Quantitative Structure Activity Relationship  
QSPR = Quantitative Structure Property Relationship  
RMSD = Root Mean Squared Deviation  
RNN = Recurrent Neural Network  
ReLU = Rectified Linear Unit  
SMILES = Simplified Molecular Input Line Entry System

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