ML4Chem: A Machine Learning Package for Chemistry and Materials Science

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ML4Chem is an open-source machine learning library for chemistry and materials science. It provides an extendable platform to develop and deploy machine learning models and pipelines and is targeted to the non-expert and expert users. ML4Chem follows user-experience design and offers the needed tools to go from data preparation to inference. Here we introduce its atomistic module for the implementation, deployment, and reproducibility of atom-centered models. This module is composed of six core building blocks: data, featurization, models, model optimization, inference, and visualization. We present their functionality and ease of use with demonstrations utilizing neural networks and kernel ridge regression algorithms.

I. INTRODUCTION

In the last decade, machine learning (ML) has undergone fast development due to large amounts of available data and advancements in computational hardware e.g. faster and cheaper central processing units (CPU), graphics processing units (GPU), and more recently the introduction of tensor processing units (TPU). Algorithmic improvements on how to compute the gradient in weight space of feedforward neural networks with respect to a loss function[1] reduced the computational time of training deep neural networks significantly. As a result companies like Google, and Facebook, introduced the most useful deep learning platforms available right now: TensorFlow[2], and Pytorch[3]. These frameworks positively impacted and advanced ML research because they helped with democratizing and simplified access to ML technologies to a larger audience.

In the field of physical chemistry and materials sciences, ML models are being standardized and applied to solve tasks such as the acceleration of atomistic simulations[4][5], prediction of the electronic Hamiltonian with generative models[6] [7], extraction of continuous latent representations for the generation of molecules[8], and even the prediction of the scent of small organic molecules[9]. It also is becoming the norm to release software solutions as support to validate results of publications that apply ML models, and alleviate the “reproducibility crisis in artificial intelligence and machine learning”[10][11]. Nevertheless, this obliquely fragments the software ecosystem because each software implementation a) requires specific data structures and b) would likely experience a lack of continuous support. There already are packages that democratize ML in chemistry. For example, DeepChem[12] has played a critical role in providing users a helpful platform of ML algorithms and featurizers for drug discovery, quantum chemistry, material sciences, and biology. More recently ChemML has been introduced as a machine learning and informatics program suite for the analysis, mining, and modeling of chemical and materials data[13]. What differentiates ML4Chem is that it focuses on easing the implementation of new functionality, extraction of intermediate quantities, interfacing with

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external programs, and exportation of any of its modules’ outputs. Also, **ML4Chem** is in its infancy bringing up the possibility to shape its future directions based on current users’ needs and ML paradigms.

Here we introduce the **atomistic** module where ML algorithms learn underlying relationships between molecules and properties treating atoms as central objects. They exploit the principle of locality in Physics: *a global quantity is defined as a sum that runs over many localized contributions*. These localized contributions usually account for interactions of an atom and its nearest-neighbor atoms (many-body interactions). Atomistic models are very useful and have been successfully applied for the acceleration of molecular dynamics simulations[17–19], identification of phase transitions in materials[20], determination of energy and atomic forces with high accuracy [21 22], the search of saddle-points [23], and the prediction of atomic charges [24, 25].

This publication is organized as follows: in section II, we will discuss the design and architecture of **ML4Chem**’s **atomistic** module. Each of its core blocks is introduced in Section III and we will demonstrate the code’s capabilities through a series of demonstration examples in Section IV. Finally, conclusions and perspectives are drawn.

### II. ATOMISTIC MODULE: DESIGN AND ARCHITECTURE

**ML4Chem** and its modules are written in Python in an object-oriented programming paradigm and are built on top of popular open-source projects to avoid duplication of efforts. In this regard, all deep learning computations are implemented with Pytorch [3]. Mathematical and linear algebra operations are executed by Numpy [26] or Scipy [27, 28] that are widely used and recognized for this purpose. Parallelism is achieved with a flexible library for parallel computing called Dask [29]. Dask enables computational scaling-up from a laptop to High-Performance Computing (HPC) clusters effortlessly and offers a web dashboard to real-time monitoring. This is particularly valuable because it provides a good estimation to users about the status of calculations, and helps at profiling computations. Good documentation is another important aspect, as the lack of it can harm usability. **ML4Chem**’s source code is documented using Numpy Python docstrings and rendered in HTML and PDF format. Also, we provide diverse information ranging from installation, theory, usage of modules, and examples.

**ML4Chem**’s modules are developed following user-experience (UX) design practices to deliver usability, accessibility, and desirability. For example, in **ML4Chem** the names of modules, classes, and functions tend to be idiomatic and easy to remember semantically. Getting the latent space from an autoencoder is performed by calling `autoencoder.get_latent_space(X)`, or the computation of atomistic features is done with a `.calculate()` class method that is provided in all featurizers under the `atomistic.features` module *e.g.* `features.calculate(X, purpose="inference")`. All modules are designed to have the same structure, enabling users to become familiar and gain intuition on their usage quickly. The library can be used in interactive Python environments such as iPython, Jupyter or JupyterLab notebooks. Or if desired, as scripts that are invoked by the Python interpreter.

Figure 1 shows a schematic representation of the machine learning workflow that drives the design philosophy used to develop the **atomistic** module in **ML4Chem**:

1. Atoms positions are mapped into atomistic feature vectors with the `atomistic.features` module and chemical symbols are used as labels.

2. ML models are instantiated utilizing atomistic feature vectors as input, and depending on the nature of the task
(supervised or unsupervised learning), targets might or might not be known.

3. Model's parameters are trained either by minimizing/maximizing a loss function or solving systems of equations. In the former case, ML4Chem provides loss and optim modules with sets of predefined loss functions and optimizers to train supervised and unsupervised atomistic models.

4. The resulting atomistic model outputs and predictions are expected to be scalar or vector quantities.

FIG. 1: Design approach of the atomistic module of ML4Chem.

As shown in Figure 2, the architecture of the atomistic module is composed of 6 building blocks. They correspond to the methods and tools required to deploy atom-centered simulations from input featurization to inference and visualization, according to our design philosophy. Modules inside the code blocks need to comply with being derived classes from base classes using Python mixins. This guarantees new classes are reusing the code base and inheriting already defined structures implicitly from the base classes to operate seamlessly with other ML4Chem components. This practice is encouraged by providing a base.py file within each module level that contains required base classes, and is enforced in the continuous integration (CI) system.

In the following sections, each of these blocks is discussed with particular attention on what can be achieved with them, their implementation, and code snippets on their usage. When relevant, we will discuss the theory and mathematics behind them as well.

III. CORE MODULES

A. Data

ML is focused on finding underlying patterns and relationships based on data examples. The format of data, a central part of ML, varies depending on the sources, and the ML algorithm adopted for solving a particular task. ML4Chem provides a data class that creates an object with a data structure that facilitates interoperability with any of its available atomistic modules.

Our atomistic ML algorithms require molecules in the form of Atoms objects as implemented in the Atomic Simulation Environment (ASE). One of the reasons behind choosing ASE is its stability and that it supports more than 50 file formats in its ase.io.read module e.g. XYZ, NWChem, GPAW, and Gaussian. For instance, an XYZ file can
be parsed and converted to an `Atoms` object in the following way: `molecule = ase.io.read("file.xyz")`. `Atoms` objects hold molecular information such as Cartesian coordinates, atom types, chemical symbols, molecular charge, cell type, energy, and atomic forces. Molecules in ASE format can be stored to disk as a list of molecules (`Atoms`) using ASE’s `Trajectory` module. Besides the formats supported by the `ase.io.read` module, we also can parse the Chemical JSON (CJSON) format\[31\] and the ANI-1 data set\[32\] using the `data.parser` module. Support of input formats such as those available in pymatgen\[33\] and MolSSI’s QCSchema are planned for future releases.

The `Data` class uses a list of molecules, or in other words a list `Atoms` objects, to generate a unique sha1 hash to label each molecule and store their respective pairs input/targets. Targets refer to the expected output of ML models and in atomistic simulations they may correspond to total energy, atomic forces, dipole moments, etc. Duplicated data points are automatically removed during the hashing procedure to avoid poor performance and numerical instability. In Listing 1 the `Data` class is instantiated by passing an ASE trajectory file with name `dataset.trajectories` containing some molecules for the purpose of "training" an atomistic ML algorithm. After hashing the molecules present in the trajectory file, pairs of input/targets examples are yielded by invoking the `.get_data()` class method and assigned to the `training_set` and `targets` variables.

Finally, once the data is loaded in memory and arranged by the `Data` class, it can be mapped into features by the `atomistic.features` module of `ML4Chem`. The content of this object, an ordered dictionary, can be easily exported as a pandas dataframe using the `.to_pandas()` class method and saved or serialized for subsequent use.
ML features are defined as a set of measurable unique characteristics or properties of an observable. They are fundamental to any ML algorithm because they represent what models “see”. Feature engineering is the process of applying domain knowledge to generate sets of numerical features that make ML algorithms work and learn meaningful representations from data. In physics constrained domains, like atomistic ML models, feature extraction is also elusive because features require to fulfill a series of properties that are commonly expected from physical systems such as rotational and translational invariance (equivariance). Featurization is a challenging time-consuming process and the feature engineering cycle encompasses their creation, selection according to importance, and validation for the task of interest.

Features can be classified in i) human-engineered features where assumptions about data are made by humans to assign properties to observables or ii) machine-engineered features where the ML algorithms discover meaningful representation during the training procedure. The `atomistic.features` module of ML4Chem supports the former case with Gaussian type features, and the latter with a `LatentFeatures` class (introduced in the following subsections).

To avoid duplication of efforts, atomistic features such as Coulomb matrix[34], smooth overlap of atomic positions (SOAP)[35], and many-body tensor representation (MBTR)[36] are supported through DScribe which is a software package for ML that provides popular feature transformations (“descriptors”) for atomistic materials simulations. This accelerates the application of ML for atomistic property prediction by providing a user-friendly, off-the-shelf descriptor implementations[37]. Our own implementation of the Coulomb matrix feature vectors is available in the `atomistic.features` module of ML4Chem, and serves as an example of how easily our package can be extended.

1. Gaussian Features

In 2007, Behler and Parrinello[7] introduced Gaussian feature vectors, also referred to as “symmetry functions” (SF), for the representation of high-dimensional potential energy surfaces with artificial neural networks. These features overcome limitations related to the image-centered models and are built for each atom in a molecule or extended system. They fingerprint the relevant chemical environment of atoms in molecules, and their computation only requires chemical symbols and atomic positions.

To delimit the effective range of interactions within the domain of a central atom, a cutoff function \( f_c \) is introduced:
\[ f_c(r) = \begin{cases} 
0.5(1 + \cos(\pi \frac{r}{R_c})), & \text{if } r \leq R_c, \\
0, & \text{if } r \geq R_c. 
\end{cases} \tag{1} \]

Where \( R_c \) is the cutoff radius (in unit length), and \( r \) is the inter-atomic distance between atoms \( i \) and \( j \). The cutoff function, with Cosine shape as shown in Eq. (1) vanishes for inter-atomic separations larger than \( R_c \) and takes finite values below the cutoff radius. These cutoff functions aim to avoid abrupt changes in the magnitudes of the features near the boundary by smoothly damping them.

There are two sets of interactions to consider when building Gaussian features: i) the radial (two-body term) and ii) angular (three-body terms) SFs. The radial SFs account for all possible interactions between a central atom \( i \) and its nearest neighbors atoms \( j \). It is defined by Eq. (2):

\[ G^2_i = \sum_{j=1}^{N_{\text{atom}}} e^{-\eta(R_{ij} - R_s)^2/R_c^2} f_c(R_{ij}), \tag{2} \]

where \( R_{ij} \) is the Euclidean distance between central atom \( i \) and neighbor atom \( j \), \( R_s \) defines the center of the Gaussian, and \( \eta \) is related to its width. Each pairwise contribution to the feature in the sum is normalized by the square of the cutoff radius \( R_c^2 \) as proposed in Ref. \[38\]. In practice, one builds a high-dimensional feature vector by choosing different \( \eta \) values.

In addition to the radial SFs (two-body term), it is possible to include triplet many-body interactions within the cutoff radius \( R_c \) using the following equation:

\[ G^3_i = 2^{1-\zeta} \sum_{j,k \neq i} (1 + \lambda \cos\theta_{ijk})^\zeta e^{-\eta(R_{ij}^2 + R_{ik}^2 + R_{jk}^2)/R_c^2} f_c(R_{ij}) f_c(R_{ik}) f_c(R_{jk}). \tag{3} \]

This part of the feature vector is built from considering the Cosine between all possible \( \theta_{ijk} \) angles of a central atom \( i \) and a pair of neighbors \( j \) and \( k \). There exists a variant of \( G^3_i \) that includes three-body interactions of triplets forming 180° inside the cutoff sphere but having an inter-atomic separation larger than \( R_c \). These SFs account for long-range interactions as described by Behler in Ref. \[39\]:

\[ G^4_i = 2^{1-\zeta} \sum_{j,k \neq i} (1 + \lambda \cos\theta_{ijk})^\zeta e^{-\eta(R_{ij}^2 + R_{ik}^2)/R_c^2} f_c(R_{ij}) f_c(R_{ik}). \tag{4} \]

In ML4Chem, Gaussian features can be built with the atomistc.features module as shown in Listing 2. The Gaussian() class is instantiated with the desired cutoff radius (units are Å) to define the neighbor atoms, the type of angular symmetry functions (either \( G^3_i \) or \( G^4_i \)), and we normalized dividing them by \( R_c^2 \). It also is possible to pass the svm keyword argument to calculate features for SVM algorithms. Note that we need to pass the data_handler and training_set objects created by the data class (see Listing 1). It is important to preprocess, scale and normalize features. In this way, models learn meaningful and noiseless underlying representations. ML4Chem uses scikit-learn\[40\] for the preprocessing of atomistic features. This can be activated by passing the preprocessor keyword argument. Currently, the supported preprocessors in ML4Chem for atomistic features are: MinMaxScaler, StandardScaler, and Normalizer. Finally, the .calculate() method calculates features.
from ml4chem.atomistic.features import Gaussian

features = Gaussian(
    cutoff=6.5,
    normalized=True,
    preprocessor="MaxMinScaler",
    save_preprocessor="features.scaler",
    angular_type="G3",
)

X = features.calculate(
    training_set, purpose="training", data=data_handler, svm=False
)

Listing 2: Computing Gaussian features with the `atomistic.features` module of ML4Chem.

2. Latent Features

In deep learning, latent features are non-directly observed variables inferred by causality[41]. These features fall under the machine-engineered classification and are determined by the ML algorithm itself during training without human intervention. A clear example would correspond to the informational bottleneck inferred when training autoencoders (AE, see Section III C 2). This informational bottleneck encodes hidden information by making a dimensionality reduction that can reconstruct the input space (directly observed variables). In physics constrained ML models, latent spaces might correspond to chemical physics aspects of atoms in molecules depending on how the model’s parameters are penalized and optimized. Their advantage over human-engineered features is that they facilitate the flexibility of the models, and when extracted with posterior inference they generalize well. Nevertheless, latent features tend to be difficult to interpret and posterior inference relies on Bayesian variational methods that are challenging to train[42]. The `atomistic.features` module in ML4Chem provides a class to ease latent feature extraction to train any of the available atomistic ML algorithms. It uses AE algorithms, like the ones described in Sections III C 2 and III C 3 and convert raw features into latent variables by forward-propagating them through the encoder part of the AE architecture. Listing 3 shows an example of the `LatentFeatures` featurization module where two keyword arguments are passed to instantiate this class:

- A tuple called `args` contains the name of the type of raw features and a dictionary with their respective parameters to be computed and subsequently converted into latent variables with a trained AE.

- We also assign to a variable `encoder` a dictionary with keys "model", and "params" containing the paths on disk to load the model and its parameters.

After instantiation, latent variables are computed with the `.calculate()` class method as it was done with the `Gaussian` class, and are returned in the right structure to be used as input to train other ML algorithms. Note that they can also be converted to a Pandas data frame.
```python
from ml4chem.atomistic.features import LatentFeatures

ae_path = "my_autoencoder/"

# Arguments to build raw features
normalized = True
preprocessor = ("MinMaxScaler", {"feature_range": (-1, 1)})
args = {
    "Gaussian",
    {
        "preprocessor": preprocessor,
        "cutoff": 6.5,
        "normalized": normalized,
        "save_preprocessor": "iso.scaler",
        "overwrite": False,
    },
}

# Dictionary to load trained autoencoder
encoder = {
    "model": ae_path + "vae.ml4c",
    "params": ae_path + "vae.params"}

features = LatentFeatures(encoder=encoder, features=args)
latent = features.calculate(
    inputs,
    purpose="training",
    data=data_handler,
    svm=False
)

df = features.to_pandas()
```

Listing 3: Extraction of latent features using the `atomistic.features` module of ML4Chem.

C. Models

This section describes atomistic ML regression algorithms in ML4Chem under the `atomistic.models` module. At this point, it is important to differentiate ML algorithms from ML models. ML algorithms refer to all procedures and steps carried out to solve a determined ML task while models are well-defined results of algorithms. Another important difference is that ML models are fed inputs to infer or predict some output. Atomistic ML algorithms exploit the physical phenomenon of “locality” where atoms are fundamental entities and whose ML features can be extracted by measuring interactions between each atom and its nearest-neighbor atoms. Predictions $P$ are therefore calculated as the sum of many individual contributions as shown in Eq. 5:

$$P = \sum_{i=1}^{n} p_i(F_i(R_i)),$$  \hspace{1cm} (5)
where a local contribution $p_i$ is a functional of a feature mapping function $F_i$ that takes as arguments atom positions $R_i$, and chemical symbols. There are two possible flavors of these algorithms: \textit{i}) a sub regression model for each chemical symbol in the data set exists or \textit{ii}) a unique regression model is used for all chemical element types.

To implement new deep learning atomistic algorithms in \textit{ML4Chem}, developers have to derive their classes by inheriting the structure from the \texttt{DeepLearningModel} base class shipped in the \texttt{base.py} file and shown in Listing 4:

```python
from abc import ABC, abstractmethod
import torch

class DeepLearningModel(ABC, torch.nn.Module):
    @abstractmethod
    def name(cls):
        """Return name of the class""
        return cls.NAME

    @abstractmethod
    def __init__(self, **kwargs):
        """Arguments needed to instantiate the model""
        pass

    @abstractmethod
    def prepare_model(self, **kwargs):
        """Prepare model for training or inference""
        pass

    @abstractmethod
    def forward(self, X):
        """Forward propagation pass""
        pass

Listing 4: Abstract base class for the implementation of new atomistic deep learning models under \texttt{atomistic.models} module of \textit{ML4Chem}.

Deep learning classes require a \texttt{name()} method that returns the name of the model, list of keyword arguments to instantiate the model using the reserved \texttt{__init__()} constructor, a \texttt{prepare_model()} method where parameters are initialized and the algorithm is prepared for the purposes of training or inference. Finally, a \texttt{forward()} method is responsible to perform a forward pass and return predictions. Support vector machine algorithms, require all these methods except for the \texttt{forward()} function. The fulfillment of this structure enables inter-operability within \textit{ML4Chem}.

1. \textit{Supervised Learning}

Supervised learning refers to the ML task of determining a complex function that maps inputs into outputs from labeled pairs of input/target examples. Its application assumes there exists a good understanding of the structure and existent classes of the data. It is worth noting that the interpretation of these models is usually easier than the
ones obtained from unsupervised learning ML tasks.

In atomistic ML algorithms, the inputs correspond to attributes of molecules such as atom positions, atom types, total charge, band-gap, etc. All these can be directly used as features, or mapped applying domain knowledge rules. To train the models, targets can be any scalar or vector quantity associated with a molecule like total energy, dipole moment, or atomic forces. Supervised algorithms tend to require large amounts of data, and featurization tends to be biased because it is human-engineered. Thus, models are usually prone to perform poorly beyond training set regimes. Also, the data sets have to be designed with enough diversity to capture meaningful underlying structures from input data. Active learning protocols[43,44] are known to help in assuring this diversity. After the training procedure, and assessment of the predictive power of models by cross-validation[45] or other model validation techniques, their parameters can be stored to perform inference in unknown data.

In the following sections, we discuss the type of neural network architectures and support vector machine algorithms supported in the atomistic.models module of ML4Chem.

a. Neural Networks

Neural Networks (NN) are algorithms inspired by how the human brain works. Their building blocks are constituted by hidden layers with interconnected neurons. NN can approximate any function with arbitrary accuracy and their outputs can be represented by the following equation:

\[
y = WX + b,
\]

where \( X \) are the inputs, \( W \) are learnable parameters, \( b \) are the biases and \( y \) are outputs. Eq. 6 is nothing but a linear function and without any extra modification, it can only fit data having a linear response. In the context of deep learning, the product \( WX \) is a nested function[46] composed of \( l \) hidden layers that returns either a vector or a scalar:

\[
y = Wx + b = f_4(f_3(f_2(f_1(x)))).
\]

From the equation above, \( l = 4 \) means the model has four layers that output a vector. The nested function \( f_l(z) \) is defined as:

\[
f_l(z) = a_l(W_lz + b_l),
\]

where we have introduced an activation function denoted by \( a_l \). The effect of the activation function on the outputs \( z \) of the neurons is to add non-linear response making the NN suitable to approximate more complex non-linear functions. Some of the most common activation functions are \( tanh(z) \), \( sigmoid(z) \), and \( relu(z) \):

\[
tanh(z) = \frac{(e^z - e^{-z})}{(e^z + e^{-z})}
\]

\[
sigmoid(z) = \frac{1}{1 + e^{-z}}
\]
When forward-propagating information through NN, the type of activation function applied to the last layer determines whether the task is regression (linear or non-linear activation function), or classification (logistic activation function)\(^\text{1}\). When more than two hidden layers are stacked between the input and output layers, NN are called \textit{deep neural networks}.\(^\text{1}\)

In Figure 3 we show a neural network with one input layer, two hidden layers, and an output layer. In this multi-layer perceptron, we assume all outputs of a layer are connected to the inputs of succeeding layers.

\[
\text{relu}(z) = \begin{cases} 
0 & \text{for } z \leq 0 \\
z & \text{for } z \geq 0 
\end{cases}
\]  

(11)

Our atomistic neural network implementation follows the structure described by Behler and Parrinello\(^\text{7}\), where each set of chemical elements in the data set has its own NN algorithm. For instance, if the data set contains C, H, N, and O atoms, then there will be four different NN. Hidden layers are \textit{fully connected} and the same activation function is applied to each of them. Different activation functions to each hidden layer and application of convolutions are features planned for future releases. However, various NN with different activation functions can be merged and trained simultaneously with the \texttt{ModelMerger} module described in Section III C 3. NN algorithms are trained by forward-propagating the atomistic feature vectors of molecules through their respective NN. The outputs of the models are atomic contributions. If targets are global quantities, such as the total energy, then the local atomic energy contributions of a molecule can be summed up and used to evaluate the loss function against corresponding targets. In this way, when backward-propagating the models, the optimization process will account for the global quantity. The usage of NN in the \texttt{atomistic.models} module is illustrated in Listing 5. A \texttt{NeuralNetwork} class with two hidden layers of 10 nodes each is instantiated, and a ReLU activation function is applied to all hidden layers except for the output.
layer. The model is subsequently prepared for training with atomistic feature vectors holding 4 dimensions per atom. Note that we also feed the data_handler object created with the Data class.

```python
from ml4chem.atomistic.models import NeuralNetwork

# Parameters
n = 10  # Hidden layers
activation = "relu"  # Activation function
input_dimension = 4  # Input Dimension

# Model instantiation and preparation
nn = NeuralNetwork(hiddenlayers=(n, n), activation=activation)
nn.prepare_model(
    input_dimension,
    data=data_handler,
    purpose="training"
)
```

Listing 5: Usage example script of the atomistic neural network class in ML4Chem.

### 6. Support Vector Machines

A Support Vector Machine (SVM) is a discriminative classifier that for some given labeled examples, it outputs an optimal hyperplane which categorizes new examples. For atomistic ML models, Kernel Ridge Regression (KRR) is a very useful algorithm based on linear ridge regression that aims to minimize a squared error loss function with a $l_2$-norm regularization term:

\[
\arg\min_{\beta \in \mathbb{R}^D} \sum_{i=1}^{N} (\beta x_i - y_i)^2 + \lambda \|\beta\|^2_2,
\]

where $\beta$ are regression coefficients, $x_i$ is the $i^{th}$ model input, $y_i$ is its corresponding $i^{th}$ target, and $\lambda \geq 0$ is a hyperparameter representing the penalty during the minimization of the loss function. Although the loss function in Eq. 12 is only suitable for linear regression, it can be applied to non-linear problems using the kernel trick (KT)\cite{17}. The KT allows model inputs (atom features in this framework) to be mapped into other feature spaces through the use of kernel functions (KF). The importance of KF lies in that they are computed directly in raw input space (atomistic feature vectors are not explicitly modified). KF are non-negative real-valued functions between two vectors $k(x, y)$. According to Mercer’s theorem, matrices built from KFs are squared definite-positive covariant kernel matrices\cite{18}. Thus, they provide some numerical advantages because they can be factorized and generally provide well-defined solutions. The KRR algorithm of the atomistic module of ML4Chem supports radial basis kernel functions (RBF), squared exponential kernel, linear, and Laplacian kernels in both their isotropic and anisotropic\cite{19} variants as seen on Eqs. 13 and 14 respectively:

\[
k(x, y)^{iso}_{rbf} = \exp \left( -\frac{||x - y||_2^2}{2\sigma^2} \right),
\]

\[ (13) \]
\[ k(x, y)_{aniso} = \exp \left( -\sum_{i=1}^{D} \frac{(x_i - y_i)^2}{2\sigma_i^2} \right), \quad (14) \]

where \( x \) and \( y \) are input and reference atom feature vectors respectively, \( \sigma \) is the Gaussian width, and \( D \) is the number of dimensions of the feature vectors. These kernels are bivariate functions that compute the similarity based on distances between two vectors in a normalized vector space. The output of these KF is within the interval \([0, 1]\).

Note that anisotropic variants of kernels allow the assignment of specific variances for each dimension of the feature vectors without the need to modifying them explicitly.

Our implementation is inspired by Ref. [50] and uses the atomistic feature vectors to build the kernel matrix shown in Eqs. [15]. Kernel matrices are positive definite which make their Cholesky decomposition to provide a unique solution to the regression coefficients. This is a particular advantage of these models since their solution is deterministic. By deterministic we mean a model that for a given input always reproduces the same regression coefficients, and consequently the same output. That is not the case in models such as neural networks that are randomly initialized leading to different local minima that are valid solutions.

\[
\begin{bmatrix}
    k_{11}^E & k_{12}^E & k_{13}^E & \ldots & k_{1N}^E \\
    k_{21}^E & k_{22}^E & k_{23}^E & \ldots & k_{2N}^E \\
    \vdots     & \vdots     & \vdots     & \ddots & \vdots     \\
    k_{N1}^E & k_{N2}^E & k_{N3}^E & \ldots & k_{NN}^E \\
\end{bmatrix}
\begin{bmatrix}
    \beta_1 \\
    \beta_2 \\
    \vdots \\
    \beta_N \\
\end{bmatrix}
= 
\begin{bmatrix}
    E_1 \\
    E_2 \\
    \vdots \\
    E_N \\
\end{bmatrix} 
\quad (15)
\]

Solving the system of equations in Eq. [15] requires atomic quantities that might not be available in quantum mechanics such as atomic energies. In those cases, one has to rely on atomic decomposition Ansatz as described by Bartók [35, 51, 52]. Instead of training using a loss function, the regression coefficients \( \beta \) in Eq. [15] are determined by forward and backward substitution after matrix factorization using the Cholesky decomposition method.

After training, a prediction is carried out by

\[
f(x) = \sum_{i=1}^{n} \beta_i k(x_i, y_j). \quad (16)\]

The atomistic module of ML4Chem can perform kernel ridge regression as shown in Listing [6]. The KernelRidge class is instantiated with the kernel function to be used, the isotropic variance \( \sigma \) and penalization values. Then, the model is prepared by passing the feature and reference features to build the covariance matrix. We also support Gaussian process regression in which case the listing has to be modified to import the GaussianProcess class instead. When doing so, the predictions will also return a tuple with the associated predictive uncertainty.
```python
from ml4chem.atomistic.models import KernelRidge

# Parameters
sigma = 1.0  # Sigma kernel value
target = "rbf"  # Kernel type
lamda = 1e-5  # Penalization

# Model instantiation and preparation
krr = KernelRidge(sigma=sigma, kernel=target, lamda=lamda)
krr.prepare_model(
    feature_space,
    reference_features,
    data=data_handler)
```

Listing 6: Usage example script of the atomistic Kernel ridge regression class in ML4Chem.

2. *Unsupervised Learning*

Probably the best example of an unsupervised learning algorithm corresponds to autoencoders (AE). AE are a type of artificial neural network architecture composed by an encoder and a decoder that can learn data representations without human intervention. As shown in Figure 4, an AE forward propagates the input through the encoder architecture to reach an informational bottleneck where latent features, denoted with $h_d$, are extracted. The bottleneck layer is usually of lower dimensionality compared to the input. Afterward, these latent features are used by the decoder to reconstruct the inputs. This reconstruction task might seem uninteresting but depending on the type of AE architecture, and what is being reconstructed, the model can learn low-dimensional representations of the input space, denoise data, or even how to generate new examples.

An autoencoder with fully connected hidden layers tends to only “memorize” how to reconstruct the input space or denoise data. Their power can be enhanced by penalizing the latent space with a loss function\[53\], attachment of external task\[11\], or just by forcing sparse activations\[54, 55\].

Our implementation of autoencoders supports two architectures that can be set with the *one_for_all* boolean keyword argument. If set to true, a single autoencoder is used for all types of atoms in the training data, otherwise, each set of atom type will have its autoencoder as in the Behler-Parrinello scheme. In Listing 7 an *AutoEncoder()* class in the *atomistic.models* module of ML4Chem is instantiated with an encoder/decoder architecture where input’s dimensions are lowered from 40 to 4 dimensions. In this case, the hyperbolic tangent activation function is applied to all layers with the *activation* keyword argument, and when the *purpose* is not set then training is assumed. Currently, only the same activation function for all layers is supported but this functionality can be extended in future releases. Note that the *prepare_model()* class method sets the dimensionality of the output to be the same as the input.

In 2013 Kingma and Welling\[56\] proposed a modification to autoencoder architectures employing a variational Bayes method that infers a posterior probability. Instead of encoding a latent vector, as seen in Figure 5 the variational autoencoder (VAE) encodes a non-differentiable latent normal distribution with unit variance. To make the model trainable, a latent vector is randomly sampled from the distribution and used by the decoder to reconstruct the input
from ml4chem.atomistic.models import AutoEncoder

hiddenlayers = {"encoder": (20, 10, 4), "decoder": (4, 10, 20)}
activation = "tanh"
purpose = "training"
input_dimension = 40

ae = AutoEncoder(
    hiddenlayers=hiddenlayers,
    activation=activation,
    one_for_all=False
)

ae.prepare_model(input_dimension, input_dimension, data=data_handler)

Listing 7: Training example script of the atomistic autoencoder class in ML4Chem.

space. This is known as the reparameterization trick and allows the calculation of the gradient of the loss function with respect to the parameters of this architecture[56]. VAEs are known to produce blurry reconstructions[57, 58] of the inputs, but more meaningful data representations, unlike vanilla AEs. That is because the VAEs are forced to minimize reconstruction errors from feature vectors sampled from the latent distribution. VAEs are classified as generative models because they learn smooth conditional distributions of the input space $X$ for given evidence and are
even able to generate new examples. VAEs can be invoked in ML4Chem by importing `VAE()` instead of the `AutoEncoder()` class in Listing 7.

In our VAE implementation, we also support the `one_for_all` keyword argument, and provide three variants when passing the `variant` string keyword argument:

1. "multivariate": the decoder outputs a distribution with mean and unit variance, and the model is trained by minimizing the negative of the log-likelihood plus the Kullback–Leibler divergence. This is useful when outputs are continuous variables. Expected feature range $[-\infty, \infty]$.

2. "bernoulli": the sigmoid activation function is applied to the decoder’s outputs and models are also minimized with the negative of the log-likelihood plus the Kullback–Leibler divergence. Features must be in a range $[0, 1]$.

3. "dcgan": hyperbolic tangent activation function is applied to the decoder’s outputs, and the model is trained by minimizing the negative of the log-likelihood plus the Kullback–Leibler divergence. Useful for feature ranges $[-1, 1]$.

![Variational Autoencoder architecture](image)

**FIG. 5: Variational Autoencoder architecture.**

### 3. Semi-Supervised and Hybrid Learning

In previous sections, we have discussed the supervised and unsupervised learning algorithms. Supervised learning requires large amounts of data to be labeled in input/target pairs which usually requires a very costly process carried
out by an ML engineer or a data scientist. On the other hand, unsupervised learning does not require labeled data but its applicability is very limited and dependent on the problem that is tried to be solved. To overcome these limitations, semi-supervised learning tasks consist of building algorithms that can work with both labeled and unlabeled data. Mixing both tasks is challenging as several loss functions are used to then backward propagate the ensemble model.

The `atomistic.models` module allows access to semi-supervised learning with the `ModelMerger()` class. In Listing 8 we use the neural network and autoencoder models already instantiated in Listings 5 and 7 and add them to a list called `models`. We import the required loss functions from the `models.loss` module and assigned them to a list named `losses`. Similarly, the targets for each model are added to a list with name `targets`, and for the inputs, note that the neural network model takes as input the latent space from the autoencoder.

During training, there are two important options that can be passed to the `ModelMerger` class:

- A boolean "independent_loss" keyword argument to set whether or not the loss functions are merged. If set to true, models are aware of each other.

- A "loss_weights" keyword argument with a list to set how much the loss of model(i) contributes to the total loss function.

More about training procedures is elaborated in Section III D.

```python
def train...
```

Listing 8: Example of a hybrid model in ML4Chem combining an autoencoder with a neural network.
D. Model Optimization

Deep learning is a challenging optimization problem that employs the gradient descent algorithm. Partial derivatives of the loss function with respect to the model’s parameter space are required to update ML model’s parameters in the direction of steepest descent as defined by the negative of the gradient. This computation experiences the vanishing or exploding gradient problems. In the former case, the partial derivatives of the loss function with respect to the parameters of the model become so small that the optimizers cannot adjust weights to minimize or maximize the loss function. In the latter case, partial derivatives are so large that optimizers oscillate around a minimum or maximum in the loss function space and convergence is never reached. These problems remained elusive for decades but were resolved by gradient clipping\[60\], regularization\[46\], and usage of activation functions that only saturate in one direction e.g. the ReLU activation function\[61\]. The atomistic module of ML4Chem supports most of the optimizers available in Pytorch such as ADAM\[62\], stochastic gradient descend\[63\], and Adagrad\[64\].

We train models exploiting the data parallelism paradigm as illustrated in Figure 6. In this scheme, data is partitioned in mini-batches that are set with the batch_size keyword argument. Mini-batches with the input data and targets are scattered through workers in a local or remote distributed cluster. The ML algorithm also is replicated on each of these workers, and parallel forward propagation is carried out. After the forward pass, we proceed with the parallel evaluation of the loss function for each mini-batch. In this step, the Pytorch automatic differentiation package autograd computes the gradient of the loss functions with respect to the weights of the model for each mini-batch and performs parallel backward-propagation passes. Finally, the gradients are reduced in CPU-1 and we call a step in the optimizer to update weights according to this gradient. This process is repeated until the number of epochs is
exhausted or some convergence criterion is reached. The parallelism scheme described above can also be visualized with the Dask dashboard as shown in Figure 7.

Each algorithm in the `atomistic.models` module provides a `train` class to optimize its parameters. Each model has its particularities but, in general, the `train` class requires at least input, outputs, batch size, optimizer, and loss function (if needed). The `train` class is responsible for taking all this information and carry out all necessary steps to train a model including calling steps in the optimizer. The optimizers are set with the `atomistic.models.optim` module that wraps Pytorch optimizer objects from a tuple composed by the name of the optimizer, and a dictionary with its required parameters e.g.: `optimizer = ("adam", {"lr": float, "weight_decay": float})`. When ML models are trained, their parameters can be stored to disk and subsequently, be used for prediction.

![Dask dashboard tool tracking computations in real time for the case where ML4Chem is optimizing a neural network over 16 processes.](image)

**FIG. 7:** Dask dashboard tool tracking computations in real time for the case where ML4Chem is optimizing a neural network over 16 processes.

### E. Inference and Visualization

Inference is the step in an ML pipeline where a trained ML model is used to infer/predict over unseen samples. This procedure involves a similar forward pass as the carried out during training for the prediction of targets.

Probably the biggest difference with a model in training mode is that during inference no backward-propagation is carried out. Therefore, the inference is a very fast computation that requires matrix multiplication and summation. In ML4Chem, atomistic models can be saved and loaded as seen in Listing 9.

We use the `.save()` method of the `Atomistic` class to save a neural network and gaussian features passing as arguments their objects. The `label` argument is used to save them to disk with name “nn”. A trajectory file is loaded in memory that contains molecules stored as `Atoms()` ASE objects. Then, we assign the model to the `calc` variable with the `load()` function and the following arguments: i) path to optimized parameters stored in a file named `nn.ml4c`, ii) path to the `nn.params` file that contains parameters in JSON format to recreate the model and features, and iii) the preprocessor to scale the features from `nn.scaler`. 
```python
from ml4chem.atomistic import Atomistic
from ase.io import Trajectory

# Save a model
Atomistic.save(nn, features=gaussian, path='', label='nn')

# Load a model
molecules = Trajectory("test.traj")
calc = Atomistic.load(model="nn.ml4c", params="nn.params", preprocessor="nn.scaler")

for molecule in molecules:
    energy = calc.get_potential_energy(molecule)
```

Listing 9: Example of save, and load trained neural network model in ML4Chem.

(a) Parity plot.

(b) Loss function progress.

(c) 2D latent space (seaborn).

(d) 3D latent space (plotly).

FIG. 8: Visualization tools offered in ML4Chem.
Another important part of ML pipelines has to do with visualization. We provide a visualization module built on top of matplotlib, seaborn and plotly. In Figure 8 we show a parity plot between ML4Chem predictions and true values, the progress of the loss function in real-time, a 2D visualization of atomistic latent space obtained using a VAE, and its 3D interactive visualization with plotly. All these plots are offered as standalone functions. We also provide a command-line tool called ml4chem that allows quick access to these visualizations. For example, the latent space visualization shown in Figure 8c can be plotted from a stored features file using the command line ml4chem --plot tsne --file latent_space.db.

IV. DEMONSTRATIONS OF THE ML4CHEM FRAMEWORK

In this section, we present some demonstrations of running ML pipelines with ML4Chem. An atomistic neural network is trained with the ANI-1 data set, and a kernel ridge regression algorithm is trained with the QM7 data set.

A. Data Sets

The ANI-1 data set is publicly available at https://github.com/isayev/ANI1_dataset. It is provided in HDF5 format, and can be used with the atomistic module of ML4Chem when converted to ASE and passed to the Data class. In Listing 10 we show how to load ANI-1 files to memory with pyanitools and assign them to a ani_data variable. Next, this list is passed as an argument to the ML4Chem’s ani_to_ase() parser function that converts HDF5 ANI-1 data sets to ASE trajectory files. After the conversion is done, the trajectory file is saved to disk as ani.traj.

```
import ase
import pyanitools as pya
from ml4chem.data.parser import ani_to_ase
from ml4chem.data.utilis import split_data

# Load ANI-1 hdf5 files to a list
files = ["ani_gdb_s01.h5", "ani_gdb_s02.h5","ani_gdb_s03.h5"]
ani_data = [pya.anidataloader(f) for f in files]

# Pass list of hdf5 files to ML4Chem ani_to_ase() parser function.
ani_dataset = ani_to_ase(ani_data,
data_keys="energies",
trajfile="ani.traj")

# Load trajectory file and split data in 80% training and 20% test set
traj = ase.io.Trajectory("ani.traj")
split_data(traj, test_set=20, randomize=True)
```

Listing 10: Conversion of ANI-1 data set to the atomic simulation environment format (ASE).
The QM7\cite{67, 68} is available at \url{http://quantum-machine.org/datasets} as a Matlab file. According to the website, “The data set is composed of three multidimensional arrays X (7165 x 23 x 23), T (7165) and P (5 x 1433) representing the inputs (Coulomb matrices), the labels (atomization energies) and the splits for cross-validation, respectively. The data set also contains two additional multidimensional arrays Z (7165) and R (7165 x 3) representing the atomic charge and the Cartesian coordinate of each atom in the molecules”. Its conversion into an ASE trajectory file to be used in atomistic module of ML4Chem is trivial as illustrated in Listing 11.

```python
import scipy.io as sio
import ase
from ml4chem.data.utils import split_data

traj = ase.io.Trajectory("qm7.traj", mode="w")
dataset = sio.loadmat("qm7.mat")

# Cartesian coordinates
cartesian = dataset["R"]

# Atomic charge (Z), atomization energy (T)
Z = dataset["Z"]
T = dataset["T"]

for i, molecule in enumerate(Z):
    numbers, positions = [], []
    for j, z in enumerate(molecule):
        if z != 0:
            numbers.append(z)
            pos = tuple(cartesian[i][j])
            pos = [c * ase.units.Bohr for c in pos]
            positions.append(pos)
    atoms = ase.Atoms(numbers=numbers, positions=positions)
    traj.write(atoms, energy=float(T[0][i]))

# Load trajectory file and split data in 80% training and 20% test set
traj = ase.io.Trajectory("qm7.traj")
split_data(traj)
```

Listing 11: Conversion of QM7 data set to the atomic simulation environment format (ASE).

Each data set was randomized and split in 80% as training set and 20% as test set using the `split_data()` function.

B. Neural Network with Gaussian Features

In this demonstration we trained an atomistic machine learning potential using the `NeuralNetwork` class with Gaussian type features and the ANI-1 data set. For this example, we will use the high-level `Potentials` class.
from ase.io import read
from ml4chem.utils import logger
from ml4chem.atomistic.features import Gaussian
from ml4chem.atomistic.models import NeuralNetwork
from ml4chem.atomistic import Potentials
from dask.distributed import Client, LocalCluster

def run():
    # Use 500 molecules for a total of 3606 feature vectors
    n_molecules, batch_size = 500, 30
    logger("nn.log")

    training = read("../training_images.traj", index="0:{:1}.format(n_molecules))

    # Instantiate the Potentials class
    calc = Potentials(
        features=Gaussian(
            batch_size=batch_size, cutoff=6.5, normalized=True, save_preprocessor="model.scaler"
        ),
        model=NeuralNetwork(
            hiddenlayers=(10, 10), activation="tanh"
        ),
        label="nn_training",
    )

    # Optimizer options and convergence criterion
    convergence = {"energy": 5e-2}
    lr = 1.0e-2
    weight_decay = 1e-5
    optimizer = ("adam", {"lr": lr, "weight_decay": weight_decay})

    # Train the algorithm
    calc.train(
        training_set=training, convergence=convergence, optimizer=optimizer, batch_size=batch_size
    )

if __name__ == "__main__":
    cluster = LocalCluster(n_workers=16, threads_per_worker=1)
    client = Client(cluster)
    run()

Listing 12: Training a neural network algorithm in ML4Chem using the Potentials class.

In Listing 12 the Potentials class can be instantiated with any of the atomistic.features and atomistic.models objects of ML4Chem, and automates all of the necessary steps to train an atomistic ML potential.

The architecture of the NN is of two hidden layers of 10 nodes each, a hyperbolic tangent activation function,
and the ADAM optimizer. The convergence criterion is set to be a root-mean-squared error of 0.05 Hartree. The optimization converged at about 5000 epochs.

In Figure 9 we show a parity plot with results obtained by predicting over 1000 unknown molecules of the test set. The NN model can do predictions with high accuracy, achieving a MAE 0.04 Hartree.

![Parity Plot](image)

**FIG. 9:** Parity plot of neural network atomistic model predictions over 1000 molecules in the ANI-1 data set. Units are in Hartree.

It is worth noting that data points in the ANI-1 dataset are obtained through normal model sampling (NMS) around the equilibrium geometry. Therefore, Gaussian features of atoms are very close to each other in the feature space. To prove this hypothesis we carried out a principal component analysis (PCA) dimensionality reduction of the Gaussian features of the training and test sets. In Figure 10 we can see how the training and test data points are close in the lower PCA dimensional space making it easy for the model to correctly predict them.

C. Kernel Ridge Regression with Coulomb Matrix Features

In this demonstration, we train a kernel ridge regression (KRR) model for the task of predicting atomization energies with the QM7 data set. Instead of the Potentials class, we will use a modular approach to show the flexibility of ML4Chem. We proceed, as shown in Section III A, to the instantiation of the Data() class with the qm7.traj trajectory
file for the purpose of "training". The training set and targets are obtained with the .get_data() class method to interoperate with ML4Chem (see Listing 13).

We use the rows of the Coulomb matrix features [69] as atomistic features vectors. The training set is featurized by instantiating the CoulombMatrix class implemented using the DScribe package and calling the .calculate() method. In the next step, the KernelRidge class is instantiated with keyword arguments to set training data, batch size, kernel function (radial basis function, or RBF), and a sigma value. The model is prepared with features and reference space. In the last step, we call the .train() method to fit the KRR algorithm passing as arguments the training set, targets, and Data objects and the model’s parameters are saved to disk.

Now, we can proceed to load this model and predict unknown data points (see Listing 14). The results are shown in a parity plot in Figure 11 with the root-mean-squared (RMSE) and mean squared (MAE) error metrics. This model showed an MAE of 14.2 kcal/mol, compared to 9.9 kcal/mol when using the same RBF kernel on the Coulomb matrix sorted eigenspectrum in Ref. [34]. This significant difference is expected because, for this demonstration, we determined sigma to be the average of the euclidean distance of the Coulomb matrix atomic feature vectors. This is not an exhaustive way of determining this hyperparameter, and in practice, one has to rely on k-fold cross-validation to find the best value that fits the data of interest. Also, the number of training set data points in our demonstration is smaller than the used in Ref [34].
from ml4chem.data.handler import Data
from ase.io import read
from ml4chem.utils import logger
from ml4chem.atomistic.features import CoulombMatrix
from ml4chem.atomistic.models import KernelRidge
from ml4chem.atomistic import Atomistic
from dask.distributed import Client, LocalCluster

def run():
    # Use 500 molecules for a total of 7727 feature vectors
    n_molecules, batch_size = 500, 10

    # Start a logger object to write to file
    logger("experiments.log")

    # Read training data
    training = read("training_images.traj", index="0:{}:1".format(n_molecules))

    # Prepare Data object
    data = Data(training, purpose="training")
    training, targets = data.get_data(purpose="training")

    # Featurization using Coulomb Matrix
    n_atoms_max = max(data.atoms_per_image)
    cm = CoulombMatrix(n_atoms_max=n_atoms_max, batch_size=batch_size)
    features, reference_space = cm.calculate(training, data=data, svm=True, purpose="training")

    # Instantiate model, prepare and train. sigma is set to average Euclidean
    # distance of feature vectors
    krr = KernelRidge(
        trainingimages="training_images.traj", batch_size=batch_size, kernel="rbf",
        sigma=26.808046418478668
    )
    krr.prepare_model(features, reference_space, data=data, purpose="training")
    krr.train(training, targets, data=data)

    # Save model to disk
    Atomistic.save(krr, features=cm, path="krr/", label="publication")

if __name__ == "__main__":
    cluster = LocalCluster(n_workers=15, threads_per_worker=1)
    client = Client(cluster)
    run()

Listing 13: Training a kernel ridge regression algorithm in ML4Chem.
from ase.io import read
from ml4chem.utils import logger
from ml4chem.atomistic import Atomistic
from dask.distributed import Client, LocalCluster
from ml4chem.data.visualization import parity
import pandas as pd

def run():
    # Use 1000 molecules from the test set
    n_molecules = 1000

    # Start a logger object to write to file
    logger("inference.log")

    # Read test data
    calc = Atomistic.load(model="krr/publication.ml4c", params="krr/publication.params")

    # Set the reference space
    calc.reference_space = "features.db"

    # Compute energies
    energies = []
    trues = []

    for index, atoms in enumerate(test):
        energy = calc.get_potential_energy(atoms)
        true = atoms.get_potential_energy()
        print(true, energy)
        trues.append(true)
        energies.append(energy)

    df = pd.DataFrame(
        {"ML4Chem Energies": energies, "True Values": trues}
    )
    df.to_pickle("inference_results.pkl")
    parity(
        energies, trues, scores=True, filename="parity_inference.png"
    )

if __name__ == "__main__":
    cluster = LocalCluster(n_workers=15, threads_per_worker=1)
    client = Client(cluster)
    run()

Listing 14: Inference using trained kernel ridge regression ML4Chem parameters.
V. CONCLUSIONS

We presented the atomistic module of ML4Chem, an open-source software package aiming to ease the deployment and implementation of ML models in chemistry and materials science. Its structure is designed with strict modularity and defined in such a way that each of its parts can be used as standalone programs. ML4Chem provides all needed methods and tools for an ML discovery pipeline – that is to go from raw data to inference and visualization. We showed with code snippets and demonstration cases what can be achieved with the core atomistic modules, and the intended intuitiveness derived from the use of UX design rules. These attributes make ML4Chem a potential platform for the implementation of new models, targeted to both non-experts and expert users.

For future development directions, we plan to extend the support of other input data formats of the atomistic module beyond the atomic simulation environment (ASE); introduce a geometric module in ML4Chem to work with geometric deep learning; addition of convolutional neural networks for both the atomistic and geometric modules; transfer learning; and implementation of active learning protocols.
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