

# TensorAlloy

**TensorAlloy** is a TensorFlow based machine learning framework for metal alloys. **TensorAlloy** builds direct computation graph from atomic positions to total energy. Thus, atomic forces and the virial stress tensor can be derived by the **AutoGrad** module of TensorFlow directly.

## 1. Requirements

- Python>=3.7.0
- TensorFlow==1.13.1
- scikit-learn
- scipy
- numpy
- ase>=3.18.0
- matplotlib>=3.0.0
- toml==0.10.0
- joblib
- wheel

Anaconda3 can install above packages without pain. However, the performance of conda-provided tensorflow is not that good.

Natively compiled TensorFlow, with all CPU features (SSE, AVX, etc.) enabled, is strongly recommended.

**Note:** [prefix] indicates the top-level directory where the program is unzipped.

## 2. Installation

This package can be installed with `pip` or `python`:

```
cd tensoralloy
pip install .
```

or

```
cd tensoralloy
python setup.py install
```

A command-line interface `tensoralloy` will also be installed.

## 3. Training

### 3.1 Usage

[TOML](#) is the configuration file format used by TensorAlloy. All the necessary keys are included in the two examples. Default keys and values can be found in [default.toml](#).

To run a training experiment, the easiest way is using the command-line program `tensoralloy`. Here are some key commands:

- `tensoralloy build database [extxyz]`: build a database from an `extxyz` file.
- `tensoralloy run [input.toml]`: run an experiment from a `toml` input file.

- `tensoralloy print [logfile]`: read the evaluation results from a logfile and print these results to a csv file.
- `tensoralloy --help`: print the help messages.

## 3.2 Output

After the training, a binary `pb` file (the trained model) will be exported to the `model_dir` specified in the input toml file. This exported `pb` file can be used by the ASE-style [TensorAlloyCalculator](#).

We also provide three pre-trained models:

- Ni: energy, force
- Mo: energy, force, stress
- Ni-Mo: energy, force, stress

## 3.3 Examples

We also provide two training examples:

### 1. QM7

- This example starts from the raw `extxyz` file. The first step is building the SQLITE3 database.
- Angular symmetry functions are enabled.

### 2. SNAP/Ni-Mo

- The binary alloy dataset.
- Only radial symmetry functions are used by default.

For each example, a `run.sh` is provided. The evaluation results at each evaluation step can be found in the `summary.csv` in the directory `train`. This csv file is generated by the command `tensoralloy print train/logfile`.

## 4. Prediction

The ASE-style calculator, [TensorAlloyCalculator](#), shall be used to make predictions.

```
#!/coding=utf-8
""" Simple usage. """
from tensoralloy.calculator import TensorAlloyCalculator
from ase.build import bulk
from ase.units import GPa

calc = TensorAlloyCalculator("NiMo.pb")
atoms = bulk("Ni", cubic=True)
atoms.calc = calc
print(atoms.get_total_energy())
print(atoms.get_forces())
print(atoms.get_stress() / GPa)
```

## 5. Input

In this section we will introduce the options and values of the input toml file.

### 5.1 Root

- `precision`: medium (float32) or high (float64). Default is medium.

- `seed`: the global seed. Default is 611.

## 5.2 Dataset

- `dataset.sqlite3`: a string, the [ASE Sqlite3 Database](#) to use.
- `dataset.name`: a string, the name of this experiment.
- `dataset.rc`: a float, the cutoff radius.
- `dataset.tfrecords_dir`: a string, the directory to save/load tfrecords files.
- `dataset.test_size`: an integer, the number of examples for evaluation.
- `dataset.serial`: a boolean. Default is `false`. Typically this should be `false` so that structures in the `sqlite3` database can be written to tfrecords in parallel. For some old Linux systems or debugging, this can be set to `true`.

## 5.3 NN

### 5.3.1 General options

- `nn.activation`: a string, the activation function to use. Options are: `elu`, `leaky_relu`, `softplus`, `softsign`, `tanh`, `sigmoid`.
- `nn.minimize`: a list of string, the properties to minimize. Some examples: `["energy", "forces"]`, `["energy", "stress"]`
- `nn.export`: a list of string, the properties that the exported model should predict. This option has the same format with `nn.minimize`.

### 5.3.2 Loss options

- `nn.loss.energy.weight`: a float, the weight of the energy loss. Default is 1.
- `nn.loss.energy.per_atom_loss`: a boolean. If true, per-atom energy loss will be used. Default is `false`.
- `nn.loss.energy.method`: a string, `"rmse"` or `"logcosh"`.
- `nn.loss.forces.weight`: a float, the weight of the force loss. Default is 1.
- `nn.loss.forces.method`: a string, `"rmse"` or `"logcosh"`.
- `nn.loss.stress.weight`: a float, the weight of the force loss. Default is 1.
- `nn.loss.stress.method`: a string, `"rmse"` or `"logcosh"`.
- `nn.loss.l2.weight`: a float, the weight of the L2 regularization. Default is 1.0. **Note: adaptive optimizers (adam, nadam, etc) are not compatible with L2. If you use any adaptive optimizer, please set this to 0.**
- `nn.loss.l2.decayed`: a boolean. If True, the L2 weight will decay exponentially.
- `nn.loss.l2.decay_rate`: a float controls the decay rate.
- `nn.loss.l2.decay_steps`: an integer.

### 5.3.3 Neural network options

- `nn.atomic.arch`: a string, the architecture of the atomistic neural networks. There two options in this package: `"AtomicCNN"` or `"AtomicResNN"`. The former is the traditional type and later is the modified version proposed by the TensorAlloy paper.

- `nn.atomic.kernel_initializer`: a string, the initialization method.
- `nn.atomic.minmax_scale`: a boolean. If True, the min-max normalization of the input features will be enabled. Default is true.
- `nn.atomic.resnet.fixed_static_energy`: a boolean. If `nn.atomic.arch` is `AtomicResNet` and this is true, the static energy parameters will be fixed.
- `nn.atomic.behler.eta`: a list of float, the eta values for radial symmetry functions.
- `nn.atomic.behler.omega`: a list of float, the omega values for radial symmetry functions.
- `nn.atomic.behler.beta`: a list of float, the beta values for angular symmetry functions.
- `nn.atomic.behler.gamma`: a list of float, the gamma values for angular symmetry functions.
- `nn.atomic.behler.zeta`: a list of float, the zeta values for angular symmetry functions.
- `nn.atomic.behler.angular`: a boolean. If true, angular symmetry functions will be used. Otherwise gamma, zeta and beta will be ignored.

### 5.3.4 Layers

The following section demonstrates how to setup the hidden layers of the atomistic neural network for element C. This block is optional. The default setting is `[64, 32]`.

```
[nn.atomic.layers]
C = [128, 64, 32]
```

## 5.4 Opt

- `opt.method`: a string, the optimizer to used. Options are: adam, nadam, sgd, adadelta, rmsprop. Default is adam.
- `opt.learning_rate`: a float, the initial learning rate. Default is 0.01.
- `opt.decay_function`: a string, the decay function to use. Options are: exponential, inverse\_time, natural\_exp. Default is false which means learning rate decay is disabled.
- `opt.decay_rate`: a float, the decay rate.
- `opt.decay_steps`: an integer.
- `opt.staircase`: a boolean.

## 5.5 Train

- `train.reset_global_step`: a boolean. If True, the global step will be set to 0 by force. If you want to continue a previous training, this should be changed to false.
- `train.batch_size`: an integer, the batch size. Default is 50.
- `train.shuffle`: a boolean. If True, the input dataset will be shuffled. In most cases this should be true.
- `train.model_dir`: a string, the working directory for this experiment.
- `train.train_steps`: an integer, the maximum training steps.
- `train.eval_steps`: an integer, the intervals between two evaluations.
- `train.summary_steps`: an integer, the intervals between two writing summary operations.

- `train.log_steps`: an integer, the intervals between two logging operations.
- `train.profile_steps`: an integer, the intervals between two performance profiling operations. Set this to 0 to disable profiling.
- `train.ckpt.checkpoint_filename`: a string, the previous checkpoint file to read. Default is `false`.
- `train.ckpt.use_ema_variables`: a boolean. If `train.ckpt.checkpoint_filename` is provided, the moving average variables will be loaded if this is true.
- `train.ckpt.restore_all_variables`: a boolean. It only works when `train.ckpt.checkpoint_filename` is provided. If this is true, all variables (including the optimizer-specific variables) will be loaded. If false, only model variables will be used. Typically, if you want to continue a previous training with exactly the same settings but more training steps, set `train.reset_global_step` to false and this option to true. If you want to use the checkpoint values as initial guesses, this option shall be false.

## 6. License

This TensorAlloy program is licensed under GNU Lesser General Public License v3.0. For more information please read [LICENSE](#).