
Chainer Chemistry Documentation

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Chainer Chemistry is a collection of tools to train and run neural networks for tasks in biology and chemistry using Chainer .

- State-of-the-art deep learning neural network models (especially graph convolutions) for chemical molecules (NFP, GGNN, Weave, SchNet etc.)
- Preprocessors of molecules tailored for these models
- Parsers for several standard file formats (CSV, SDF etc.)
- Loaders for several well-known datasets (QM9, Tox21 etc.)

Introductory to deep learning for molecules and Chainer Chemistry is also available [here \(SlideShare\)](#).

1.1 Installation

1.1.1 Dependency

Following packages are required to install Chainer Chemistry and are automatically installed when you install the library by *pip* command.

- `chainer`
- `pandas`
- `scikit-learn`
- `tqdm`

Also, it uses following library, which you need to manually install.

- `rdkit`

See the [official document](#) for installation. If you have setup `anaconda`, you may install `rdkit` by following command:

```
$ conda install -c rdkit rdkit
```

1.1.2 Install via pip

It can be installed by pip command:

```
$ pip install chainer-chemistry
```

1.1.3 Install from source

The tarball of the source tree is available via `pip download chainer-chemistry`. You can use `setup.py` to install Chainer Chemistry from the tarball:

```
$ tar xzf chainer-chemistry-x.x.x.tar.gz
$ cd chainer-chemistry-x.x.x
$ python setup.py install
```

Install from the latest source from the master branch:

```
$ git clone https://github.com/pfnet-research/chainer-chemistry.git
$ pip install -e chainer-chemistry
```

1.1.4 Run example training code

The [official repository](#) provides examples of training several graph convolution networks. The code can be obtained by cloning the repository:

```
$ git clone https://github.com/pfnet-research/chainer-chemistry.git
```

The following code is how to train Neural Fingerprint (NFP) with the Tox21 dataset on CPU:

```
$ cd chainer-chemistry/examples/tox21
$ python train_tox21.py --method=nfp --gpu=-1 # set --gpu=0 if you have GPU
```

1.2 Tutorial

1.2.1 Abstract

In this tutorial, we predict Highest Occupied Molecular Orbital (HOMO) level of the molecules in [QM9 dataset](#) [1][2] by [Neural Finger Print \(NFP\)](#) [3][4]. We concentrate on explaining usage of Chainer Chemistry briefly and do not look over the detail of NFP implementation.

1.2.2 Tested Environment

- Chainer Chemistry $\geq 0.0.1$ (See [Installation](#))
- Chainer $\geq 2.0.2$
- CUDA ≥ 8.0 , CuPy $\geq 1.0.3$ (Required only when using GPU)
 - For CUDA 9.0, CuPy $\geq 2.0.0$ is required
- sklearn $\geq 0.17.1$ (Only for preprocessing)

1.2.3 QM9 Dataset

QM9 is a publicly available dataset of small organic molecule structures and their simulated properties for data driven researches of material property prediction and chemical space exploration. It contains 133,885 stable small organic molecules made up of CHONF. The available properties are geometric, energetic, electronic, and thermodynamic ones.

In this tutorial, we predict HOMO level in the properties. Physically, we need quantum chemical calculations to compute HOMO level. From mathematical viewpoint it requires a solution of an internal eigenvalue problem for a Hamiltonian matrix. It is a big challenge to predict HOMO level accurately by a neural network, because the network should approximate both calculating the Hamiltonian matrix and solving the internal eigenvalue problem.

1.2.4 HOMO prediction by NFP

At first you should clone the library repository from [GitHub](#). There is a Python script `examples/qm9/train_qm9.py` in the repository. It executes a whole training procedure, that is, downloads QM9 dataset, preprocess it, define an NFP model and run training on them.

Execute the following commands on a machine satisfying the tested environment in environment.

```
~$ git clone git@github.com:pfnet-research/chainer-chemistry.git
~$ cd chainer-chemistry/examples/qm9/
```

Hereafter all shell commands should be executed in this directory.

If you are a beginner for Chainer, [Chainer hands-on](#) will greatly help you. Especially the explanation of inclusion relationship of Chainer classes in Sec. 4 in [Chap. 2](#) is helpful when you read the sample script.

Next the dataset preparation part and the model definition part in `train_qm9.py` are explained. If you are not interested in them, skip [Dataset Preparation](#) and [Model Definition](#), and jump to [Run](#).

Dataset Preparation

Chainer Chemistry accepts the same dataset type with Chainer, such as `chainer.datasets.SubDataset`. In this section we learn how to download QM9 dataset and use it as a Chainer dataset.

The following Python script downloads and saves the dataset in `.npz` format.

```
#!/usr/bin/env python
from chainer_chemistry import datasets as D
from chainer_chemistry.dataset.preprocessors import preprocess_method_dict
from chainer_chemistry.datasets import NumpyTupleDataset

preprocessor = preprocess_method_dict['nfp']()
dataset = D.get_qm9(preprocessor, labels='homo')
cache_dir = 'input/nfp_homo/'
os.makedirs(cache_dir)
NumpyTupleDataset.save(cache_dir + 'data.npz', dataset)
```

The last two lines save the dataset to `input/nfp_homo/data.npz` and we need not to download the dataset next time.

The following Python script read the dataset from the saved `.npz` file and split the data points into training and validation sets.

```
#!/usr/bin/env python
from chainer.datasets import split_dataset_random
from chainer_chemistry import datasets as D
from chainer_chemistry.dataset.preprocessors import preprocess_method_dict
from chainer_chemistry.datasets import NumpyTupleDataset

cache_dir = 'input/nfp_homo/'
dataset = NumpyTupleDataset.load(cache_dir + 'data.npz')
train_data_ratio = 0.7
train_data_size = int(len(dataset) * train_data_ratio)
train, val = split_dataset_random(dataset, train_data_size, 777)
print('train dataset size:', len(train))
print('validation dataset size:', len(val))
```

The function `split_dataset_random()` returns a tuple of two `chainer.datasets.SubDataset` objects (training and validation set). Now you have prepared training and validation data points and you can construct `chainer.iterator.Iterator` objects, needed for updaters in Chainer.

Model Definition

In Chainer, a neural network model is defined as a `chainer.Chain` object.

Graph convolutional networks such as NFP are generally connection of graph convolution layers and multi perceptron layers. Therefore it is convenient to define a class which inherits `chainer.Chain` and compose two `chainer.Chain` objects corresponding to the two kind of layers.

Execute the following Python script and check you can define such a class. NFP and MLP are already defined `chainer.Chain` classes.

```
#!/usr/bin/env python
import chainer
from chainer_chemistry.models import MLP, NFP

class GraphConvPredictor(chainer.Chain):

    def __init__(self, graph_conv, mlp):
        super(GraphConvPredictor, self).__init__()
        with self.init_scope():
            self.graph_conv = graph_conv
            self.mlp = mlp

    def __call__(self, atoms, adjs):
        x = self.graph_conv(atoms, adjs)
        x = self.mlp(x)
        return x

n_unit = 16
conv_layers = 4
model = GraphConvPredictor(NFP(n_unit, n_unit, conv_layers),
                           MLP(n_unit, 1))
```

Run

You have defined the dataset and the NFP model on Chainer. There are no other procedures specific to Chainer Chemistry. Hereafter you should just follow the usual procedures in Chainer to execute training.

The sample script `examples/qm9/train_qm9.py` contains all the procedures and you can execute training just by invoking the script. The following command starts training for 20 epochs and reports loss and accuracy during training. They are reported for each of `main` (dataset for training) and `validation` (dataset for validation).

The `--gpu 0` option is to utilize a GPU with device id = 0. If you do not have a GPU, set `--gpu -1` or just drop `--gpu 0` to use CPU for all the calculation. In most cases, calculation with GPU is much faster than that only with CPU.

```
~/chainer-chemistry/examples/qm9$ python train_qm9.py --method nfp --label homo --gpu 0
↪ 0 # If GPU is unavailable, set --gpu -1

Train NFP model...
epoch      main/loss    main/accuracy  validation/main/loss  validation/main/accuracy
↪ elapsed_time
1          0.746135    0.0336724      0.680088             0.0322597
↪ 58.4605
2          0.642823    0.0311715      0.622942             0.0307055
↪ 113.748
(...)
19         0.540646    0.0277585      0.532406             0.0276445
↪ 1052.41
20         0.537062    0.0276631      0.551695             0.0277499
↪ 1107.29
```

After finished, you will find log file in `result/` directory.

Evaluation

In the loss and accuracy report, we are mainly interested in `validation/main/accuracy`. Although it decreases during training, the `accuracy` field is actually mean absolute error. The unit is Hartree. Therefore the last line means validation mean absolute error is 0.0277499 Hartree. See `scaled_abs_error()` function in `train_qm9.py` for the detailed definition of mean absolute error.

You can also train other type models like GGNN, SchNet or WeaveNet, and other target values like LUMO, dipole moment and internal energy, just by changing `--model` and `--label` options, respectively. See output of `python train_qm9.py --help`.

1.2.5 Using your own dataset

You can use your own dataset in Chainer Chemistry. [example/own_dataset](#) shows an example.

1.2.6 Reference

- [1] L. Ruddigkeit, R. van Deursen, L. C. Blum, J.-L. Reymond, Enumeration of 166 billion organic small molecules in the chemical universe database GDB-17, *J. Chem. Inf. Model.* 52, 2864–2875, 2012.
- [2] R. Ramakrishnan, P. O. Dral, M. Rupp, O. A. von Lilienfeld, Quantum chemistry structures and properties of 134 kilo molecules, *Scientific Data* 1, 140022, 2014.
- [3] Duvenaud, D. K., Maclaurin, D., Iparraguirre, J., Bombarell, R., Hirzel, T., Aspuru-Guzik, A., & Adams, R. P. (2015). Convolutional networks on graphs for learning molecular fingerprints. In *Advances in neural information processing systems* (pp. 2224–2232).
- [4] Gilmer, J., Schoenholz, S. S., Riley, P. F., Vinyals, O., & Dahl, G. E. (2017). Neural message passing for quantum chemistry. arXiv preprint arXiv:1704.01212.

1.3 Contribution guide

We welcome any type of contribution that helps to improve and promote Chainer Chemistry. Typical contribution includes:

- Send pull requests (PRs) to the [repository](#) (We recommend developers making PRs to read the [Development policy](#) before starting to implement).
- Report bugs or problems as [issues](#).
- Send questions to developer community sites like [Stackoverflow](#) or Chainer Slack ([en](#), [jp](#)).
- Write a blog post about Chainer Chemistry or its use case.

1.4 Development policy

In this section, we describe the development policy that the core developers follow. Developers who are thinking to send PRs to the repository are encouraged to read the following sections before starting implementation.

1.4.1 Versioning policy

Basically, we follow the [semantic versioning v2.0.0](#). In Chainer Chemistry, *public APIs* in the sense of semantic versioning are ones in [the document](#).

We follow these rules about versioning during the major version zero in addition to ones described in the the semantic versioning:

- We do not plan any scheduled releases.
- We do not plan any pre releases.
- We release the minor version when the core development team agrees. Typically, we do so when (1) sufficient number of features are added since the last minor release (2) the latest release cannot run the example code in the master branch of the repository (3) critical bugs are found. But we are not restricted to them.
- If we find critical bugs, we should release a patch version or a minor version that fixes them. The core development team will determine which version to release.

We do not have a concrete plan about versioning strategy after v1.0.0.

1.4.2 Compatibility policy

As an immediate consequence of the semantic versioning, we may break compatibility of public APIs including addition, deletion, and changes in their semantics anytime in the major version zero. Since APIs of Chainer Chemistry are still immature and unstable, we expect introduction of new features can sometime involve compatibility break. If we are faced with a dilemma between cost for backward compatibility and benefit of new features, we are likely to give up the former because we want to place importance on introducing new features as soon as possible. Of course, we care backward compatibility whenever it is easy and low-cost.

Like [ChainerCV](#), Chainer Chemistry provides several off-the-shelf deep learning models (e.g. Neural Finger Print) whose papers are available in such as arXiv or conferences related to machine learning. Although, most of published papers reports evaluation results of the models with publicly available datasets, we do *NOT* guarantee the reproducibility of experiments in the papers.

At some point, coding examples in the master branch of the official repository may not work even with the latest release. In that case, users are recommended to either use the example code of the latest release or update the library code to the master branch.

As of v0.3.0, we have introduced *BaseForwardModel*, which provides methods for serializing itself to and loading from a file. As these methods internally use `pickle`, portability of the class depends on that of pickling. Especially, serialized instances of *BaseForwardModel* made with older Chainer Chemistry may not be loaded with newer one, partly because we may change their internal structures for refactoring, performance improvement, and so on. See the document of *BaseForwardModel* and their subclasses (e.g. *Classifier*, *Regressor*).

1.4.3 Branch strategy

The official repository of Chainer Chemistry is <https://github.com/pfnet-research/chainer-chemistry>. We use the *master* branch of the repository for development. Therefore, developer who makes PRs should send them to the master branch.

During major version zero, we do not maintain any released versions. When a bug is found, changes for the bug should be merged to the next version (either minor or patch). If the bug is critical, we will release the next version as soon as possible.

1.4.4 Coding guideline

We basically adopt *PEP8* <<https://www.python.org/dev/peps/pep-0008/>> as a style guide. You can check it with *flake8*, which we can install by:

```
$ pip install flake8
```

and run with `flake8` command.

In addition to PEP8, we use upper camel case (e.g. `FooBar`) for class names and snake case (e.g. `foo_bar`) for function, method, variable and package names. Although we recommend developers to follow these rules as well, they are not mandatory.

For documents, we follow the [Google Python Style Guide](#) and compile it with [Napoleon](#), which is an extension of [Sphinx](#).

1.4.5 Testing guideline

Chainer Chemistry uses `pytest` as a unit-test framework. All unit tests are located in `tests/` directory. We can run tests with normal usage of `pytest`. For example, the following command runs all unit tests:

```
$ pytest tests
```

Some unit tests require GPUs, which are annotated with `@pytest.mark.gpu`. Therefore, you can skip them with `-m` option:

```
$ pytest -m "not gpu" tests
```

If a developer who writes a unit test that uses GPUs, you must annotate it with `@pytest.mark.gpu`.

Similarly, some unit tests take long time to complete. We annotated them with `@pytest.mark.slow` and can skip them with `-m` option:

```
$ pytest -m "not slow" tests
```

Any unit test that uses GPUs must be annotated with `@pytest.mark.slow`.

We can skip both GPU and slow tests with the following command:

```
$ pytest -m "not (gpu or slow)" tests
```

1.4.6 Terminology

In the context of machine learning, especially chemoinformatics, we use several terms such as feature, feature vectors, descriptor and so on to indicate representation of inputs. To avoid disambiguity and align naming convention within the library code, we use these terms in the following way:

- *Feature* is a representation of a sample of interest (typically molecules in Chainer Chemistry).
- *Label* is a target value of we want to predict.
- *Input feature* is a representation of a sample from which we want to predict the target value.

For example, consider a supervised learning task whose dataset consisting of input-output pairs $((x_1, y_1), \dots, (x_N, y_N))$, where N is the number of samples. In Chainer Chemistry `x_i` and `y_i` are called input feature and label, respectively and a pair of (x_i, y_i) is feature for each i .

1.4.7 Relation to Chainer

Chainer is a deep learning framework written in Python that features dynamic computational graph construction (the “define-by-run” paradigm) for flexible and intuitive model development. As the name indicates, Chainer Chemistry is an extension library of Chainer built on top of it. The core development team members of Chainer and that of Chainer Chemistry work together tightly.

1.5 API Reference

1.5.1 Dataset

Converters

<code>chainer_chemistry.dataset.converters.concat_mols</code>	Concatenates a list of molecules into array(s).
---	---

`chainer_chemistry.dataset.converters.concat_mols`

`chainer_chemistry.dataset.converters.concat_mols` (*batch*, *device=None*, *padding=0*)

Concatenates a list of molecules into array(s).

This function converts an “array of tuples” into a “tuple of arrays”. Specifically, given a list of examples each of which consists of a list of elements, this function first makes an array by taking the element in the same position from each example and concatenates them along the newly-inserted first axis (called *batch dimension*) into one array. It repeats this for all positions and returns the resulting arrays.

The output type depends on the type of examples in *batch*. For instance, consider each example consists of two arrays (x, y) . Then, this function concatenates x ’s into one array, and y ’s into another array, and returns a tuple of these two arrays. Another example: consider each example is a dictionary of two entries whose keys are `'x'` and `'y'`, respectively, and values are arrays. Then, this function concatenates x ’s into one array, and y

's into another array, and returns a dictionary with two entries `x` and `y` whose values are the concatenated arrays.

When the arrays to concatenate have different shapes, the behavior depends on the `padding` value. If `padding` is `None`, it raises an error. Otherwise, it builds an array of the minimum shape that the contents of all arrays can be substituted to. The padding value is then used to the extra elements of the resulting arrays.

The current implementation is identical to `concat_examples()` of Chainer, except the default value of the `padding` option is changed to 0.

Example

```
>>> import numpy
>>> from chainer_chemistry.dataset.converters import concat_mols
>>> x0 = numpy.array([1, 2])
>>> x1 = numpy.array([4, 5, 6])
>>> dataset = [x0, x1]
>>> results = concat_mols(dataset)
>>> print(results)
[[1 2 0]
 [4 5 6]]
```

See also:

`chainer.dataset.concat_examples()`

Parameters

- **batch** (*list*) – A list of examples. This is typically given by a dataset iterator.
- **device** (*int*) – Device ID to which each array is sent. Negative value indicates the host memory (CPU). If it is omitted, all arrays are left in the original device.
- **padding** – Scalar value for extra elements. If this is `None` (default), an error is raised on shape mismatch. Otherwise, an array of minimum dimensionalities that can accommodate all arrays is created, and elements outside of the examples are padded by this value.

Returns The type depends on the type of each example in the batch.

Return type Array, a tuple of arrays, or a dictionary of arrays

Indexers

<code>chainer_chemistry.dataset.indexer.BaseIndexer</code>	Base class for Indexer
<code>chainer_chemistry.dataset.indexer.BaseFeatureIndexer</code>	Base class for FeatureIndexer
<code>chainer_chemistry.dataset.indexers.NumpyTupleDatasetFeatureIndexer</code>	FeatureIndexer for NumpyTupleDataset

chainer_chemistry.dataset.indexer.BaseIndexer

class `chainer_chemistry.dataset.indexer.BaseIndexer`

Base class for Indexer

`__init__()`

Initialize self. See `help(type(self))` for accurate signature.

chainer_chemistry.dataset.indexer.BaseFeatureIndexer

class chainer_chemistry.dataset.indexer.**BaseFeatureIndexer** (*dataset*)

Base class for FeatureIndexer

FeatureIndexer can be accessed by 2-dimensional indices, axis=0 is used for dataset index and axis=1 is used for feature index. For example, let *features* be the instance of *BaseFeatureIndexer*, then *features[i, j]* returns *i*-th dataset of *j*-th feature.

features[ind] works same with *features[ind, :]*

Note that the returned value will be numpy array, even though the dataset is initialized with other format (e.g. list).

__init__ (*dataset*)

Initialize self. See help(type(self)) for accurate signature.

Methods

__init__ (<i>dataset</i>)	Initialize self.
check_type_feature_index (<i>j</i>)	
create_feature_index_list (<i>feature_index</i>)	
extract_feature (<i>i, j</i>)	Extracts <i>i</i> -th data's <i>j</i> -th feature
extract_feature_by_slice (<i>slice_index, j</i>)	Extracts <i>slice_index</i> -th data's <i>j</i> -th feature.
features_length ()	Returns length of features
postprocess (<i>item</i>)	
preprocess (<i>item</i>)	

Attributes

dataset_length
shape

chainer_chemistry.dataset.indexers.NumpyTupleDatasetFeatureIndexer

class chainer_chemistry.dataset.indexers.**NumpyTupleDatasetFeatureIndexer** (*dataset*)

FeatureIndexer for NumpyTupleDataset

Parameters **dataset** (*NumpyTupleDataset*) – dataset instance

__init__ (*dataset*)

Initialize self. See help(type(self)) for accurate signature.

Methods

__init__ (<i>dataset</i>)	Initialize self.
check_type_feature_index (<i>j</i>)	
create_feature_index_list (<i>feature_index</i>)	
extract_feature (<i>i, j</i>)	Extracts <i>i</i> -th data's <i>j</i> -th feature
extract_feature_by_slice (<i>slice_index, j</i>)	Extracts <i>slice_index</i> -th data's <i>j</i> -th feature.
features_length ()	Returns length of features

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<code>postprocess(item)</code>
<code>preprocess(item)</code>

Attributes

<code>dataset_length</code>
<code>shape</code>

Parsers

<code>chainer_chemistry.dataset.parsers.BaseParser</code>	
<code>chainer_chemistry.dataset.parsers.CSVFileParser</code>	csv file parser
<code>chainer_chemistry.dataset.parsers.SDFFileParser</code>	sdf file parser
<code>chainer_chemistry.dataset.parsers.DataFrameParser</code>	data frame parser
<code>chainer_chemistry.dataset.parsers.SmilesParser</code>	smiles parser

chainer_chemistry.dataset.parsers.BaseParser

class `chainer_chemistry.dataset.parsers.BaseParser`

`__init__()`
Initialize self. See `help(type(self))` for accurate signature.

Methods

<code>__init__()</code>	Initialize self.
-------------------------	------------------

chainer_chemistry.dataset.parsers.CSVFileParser

class `chainer_chemistry.dataset.parsers.CSVFileParser` (*preprocessor*, *labels=None*,
smiles_col='smiles', *post-*
process_label=None, *post-*
process_fn=None, *log-*
ger=None)

csv file parser

This FileParser parses .csv file. It should contain column which contain SMILES as input, and label column which is the target to predict.

Parameters

- **preprocessor** (`BasePreprocessor`) – preprocessor instance
- **labels** (*str* or *list*) – labels column

- **smiles_col** (*str*) – smiles column
- **postprocess_label** (*Callable*) – post processing function if necessary
- **postprocess_fn** (*Callable*) – post processing function if necessary
- **logger** –

__init__ (*preprocessor, labels=None, smiles_col='smiles', postprocess_label=None, postprocess_fn=None, logger=None*)
Initialize self. See help(type(self)) for accurate signature.

Methods

__init__ (<i>preprocessor[, labels, smiles_col, ...]</i>)	Initialize self.
extract_total_num (<i>filepath</i>)	Extracts total number of data which can be parsed
parse (<i>filepath[, return_smiles, ...]</i>)	parse csv file using <i>preprocessor</i>

chainer_chemistry.dataset.parsers.SDFFileParser

class chainer_chemistry.dataset.parsers.**SDFFileParser** (*preprocessor, labels=None, postprocess_label=None, postprocess_fn=None, logger=None*)

sdf file parser

Parameters

- **preprocessor** (*BasePreprocessor*) – preprocessor instance
- **labels** (*str or list*) – labels column
- **postprocess_label** (*Callable*) – post processing function if necessary
- **postprocess_fn** (*Callable*) – post processing function if necessary
- **logger** –

__init__ (*preprocessor, labels=None, postprocess_label=None, postprocess_fn=None, logger=None*)
Initialize self. See help(type(self)) for accurate signature.

Methods

__init__ (<i>preprocessor[, labels, ...]</i>)	Initialize self.
extract_total_num (<i>filepath</i>)	Extracts total number of data which can be parsed
parse (<i>filepath[, return_smiles, ...]</i>)	parse sdf file using <i>preprocessor</i>

chainer_chemistry.dataset.parsers.DataFrameParser

class chainer_chemistry.dataset.parsers.**DataFrameParser** (*preprocessor, labels=None, smiles_col='smiles', postprocess_label=None, postprocess_fn=None, logger=None*)

data frame parser

This FileParser parses pandas dataframe. It should contain column which contain SMILES as input, and label column which is the target to predict.

Parameters

- **preprocessor** (*BasePreprocessor*) – preprocessor instance
- **labels** (*str or list or None*) – labels column
- **smiles_col** (*str*) – smiles column
- **postprocess_label** (*Callable*) – post processing function if necessary
- **postprocess_fn** (*Callable*) – post processing function if necessary
- **logger** –

__init__ (*preprocessor, labels=None, smiles_col='smiles', postprocess_label=None, postprocess_fn=None, logger=None*)

Initialize self. See help(type(self)) for accurate signature.

Methods

__init__ (<i>preprocessor[, labels, smiles_col, ...]</i>)	Initialize self.
extract_total_num (<i>df</i>)	Extracts total number of data which can be parsed
parse (<i>df[, return_smiles, target_index, ...]</i>)	parse DataFrame using <i>preprocessor</i>

chainer_chemistry.dataset.parsers.SmilesParser

class chainer_chemistry.dataset.parsers.**SmilesParser** (*preprocessor, postprocess_label=None, postprocess_fn=None, logger=None*)

smiles parser

It parses *smiles_list*, which is a list of string of smiles.

Parameters

- **preprocessor** (*BasePreprocessor*) – preprocessor instance
- **postprocess_label** (*Callable*) – post processing function if necessary
- **postprocess_fn** (*Callable*) – post processing function if necessary
- **logger** –

__init__ (*preprocessor, postprocess_label=None, postprocess_fn=None, logger=None*)

Initialize self. See help(type(self)) for accurate signature.

Methods

__init__ (<i>preprocessor[, postprocess_label, ...]</i>)	Initialize self.
extract_total_num (<i>smiles_list</i>)	Extracts total number of data which can be parsed
parse (<i>smiles_list[, return_smiles, ...]</i>)	parse <i>smiles_list</i> using <i>preprocessor</i>

Preprocessors

Base preprocessors

<code>chainer_chemistry.dataset.preprocessors.BasePreprocessor</code>	Base class for preprocessor
<code>chainer_chemistry.dataset.preprocessors.MolPreprocessor</code>	preprocessor class specified for rdkit mol instance

chainer_chemistry.dataset.preprocessors.BasePreprocessor

class chainer_chemistry.dataset.preprocessors.**BasePreprocessor**

Base class for preprocessor

`__init__()`

Initialize self. See help(type(self)) for accurate signature.

Methods

<code>__init__()</code>	Initialize self.
<code>process(filepath)</code>	

chainer_chemistry.dataset.preprocessors.MolPreprocessor

class chainer_chemistry.dataset.preprocessors.**MolPreprocessor** (*add_Hs=False, kekulize=False*)

preprocessor class specified for rdkit mol instance

Parameters

- **add_Hs** (*bool*) – If True, implicit Hs are added.
- **kekulize** (*bool*) – If True, Kekulizes the molecule.

`__init__(add_Hs=False, kekulize=False)`

Initialize self. See help(type(self)) for accurate signature.

Methods

<code>__init__([add_Hs, kekulize])</code>	Initialize self.
<code>get_input_features(mol)</code>	get molecule's feature representation, descriptor.
<code>get_label(mol[, label_names])</code>	Extracts label information from a molecule.
<code>prepare_smiles_and_mol(mol)</code>	Prepare <i>smiles</i> and <i>mol</i> used in following preprocessing.
<code>process(filepath)</code>	

Concrete preprocessors

<code>chainer_chemistry. dataset.preprocessors. AtomicNumberPreprocessor</code>	Atomic number Preprocessor
<code>chainer_chemistry.dataset. preprocessors.ECFPPreprocessor</code>	
<code>chainer_chemistry.dataset. preprocessors.GGNNPreprocessor</code>	GGNN Preprocessor
<code>chainer_chemistry.dataset. preprocessors.NFPPreprocessor</code>	NFP Preprocessor
<code>chainer_chemistry.dataset. preprocessors.SchNetPreprocessor</code>	SchNet Preprocessor
<code>chainer_chemistry.dataset. preprocessors.WeaveNetPreprocessor</code>	WeaveNet must have fixed-size atom list for now, zero_padding option
<code>chainer_chemistry.dataset. preprocessors.RelGATPreprocessor</code>	RelGAT Preprocessor
<code>chainer_chemistry.dataset. preprocessors.RelGCNPreprocessor</code>	RelGCN Preprocessor
<code>chainer_chemistry.dataset. preprocessors.RSGCNPreprocessor</code>	RSGCN Preprocessor

chainer_chemistry.dataset.preprocessors.AtomicNumberPreprocessor

```
class chainer_chemistry.dataset.preprocessors.AtomicNumberPreprocessor (max_atoms=-1,  
out_size=-1)
```

Atomic number Preprocessor

Parameters

- **max_atoms** (*int*) – Max number of atoms for each molecule, if the number of atoms is more than this value, this data is simply ignored. Setting negative value indicates no limit for max atoms.
- **out_size** (*int*) – It specifies the size of array returned by `get_input_features`. If the number of atoms in the molecule is less than this value, the returned arrays is padded to have fixed size. Setting negative value indicates do not pad returned array.

__init__ (*max_atoms=-1, out_size=-1*)

Initialize self. See `help(type(self))` for accurate signature.

Methods

<code>__init__([max_atoms, out_size])</code>	Initialize self.
<code>get_input_features(mol)</code>	get input features
<code>get_label(mol[, label_names])</code>	Extracts label information from a molecule.
<code>prepare_smiles_and_mol(mol)</code>	Prepare <i>smiles</i> and <i>mol</i> used in following preprocessing.
<code>process(filepath)</code>	

chainer_chemistry.dataset.preprocessors.ECFPPreprocessor

class chainer_chemistry.dataset.preprocessors.**ECFPPreprocessor** (*radius=2*)

__init__ (*radius=2*)
Initialize self. See help(type(self)) for accurate signature.

Methods

__init__ ([<i>radius</i>])	Initialize self.
get_input_features (<i>mol</i>)	get molecule's feature representation, descriptor.
get_label (<i>mol</i> [, <i>label_names</i>])	Extracts label information from a molecule.
prepare_smiles_and_mol (<i>mol</i>)	Prepare <i>smiles</i> and <i>mol</i> used in following preprocessing.
process (<i>filepath</i>)	

chainer_chemistry.dataset.preprocessors.GGNNPreprocessor

class chainer_chemistry.dataset.preprocessors.**GGNNPreprocessor** (*max_atoms=-1*,
out_size=-1,
add_Hs=False,
kekulize=False)

GGNN Preprocessor

Parameters

- **max_atoms** (*int*) – Max number of atoms for each molecule, if the number of atoms is more than this value, this data is simply ignored. Setting negative value indicates no limit for max atoms.
- **out_size** (*int*) – It specifies the size of array returned by *get_input_features*. If the number of atoms in the molecule is less than this value, the returned arrays is padded to have fixed size. Setting negative value indicates do not pad returned array.
- **add_Hs** (*bool*) – If True, implicit Hs are added.
- **kekulize** (*bool*) – If True, Kekulizes the molecule.

__init__ (*max_atoms=-1*, *out_size=-1*, *add_Hs=False*, *kekulize=False*)
Initialize self. See help(type(self)) for accurate signature.

Methods

__init__ ([<i>max_atoms</i> , <i>out_size</i> , <i>add_Hs</i> , <i>kekulize</i>])	Initialize self.
get_input_features (<i>mol</i>)	get input features
get_label (<i>mol</i> [, <i>label_names</i>])	Extracts label information from a molecule.
prepare_smiles_and_mol (<i>mol</i>)	Prepare <i>smiles</i> and <i>mol</i> used in following preprocessing.
process (<i>filepath</i>)	

chainer_chemistry.dataset.preprocessors.NFPPreprocessor

```
class chainer_chemistry.dataset.preprocessors.NFPPreprocessor (max_atoms=-1,
                                                             out_size=-1,
                                                             add_Hs=False,
                                                             kekulize=False)
```

NFPPreprocessor

Parameters

- **max_atoms** (*int*) – Max number of atoms for each molecule, if the number of atoms is more than this value, this data is simply ignored. Setting negative value indicates no limit for max atoms.
- **out_size** (*int*) – It specifies the size of array returned by *get_input_features*. If the number of atoms in the molecule is less than this value, the returned arrays is padded to have fixed size. Setting negative value indicates do not pad returned array.
- **add_Hs** (*bool*) – If True, implicit Hs are added.
- **kekulize** (*bool*) – If True, Kekulizes the molecule.

```
__init__ (max_atoms=-1, out_size=-1, add_Hs=False, kekulize=False)
```

Initialize self. See help(type(self)) for accurate signature.

Methods

<code>__init__</code> (<i>max_atoms</i> , <i>out_size</i> , <i>add_Hs</i> , <i>kekulize</i>)	Initialize self.
<code>get_input_features</code> (<i>mol</i>)	get input features
<code>get_label</code> (<i>mol</i> , <i>label_names</i>)	Extracts label information from a molecule.
<code>prepare_smiles_and_mol</code> (<i>mol</i>)	Prepare <i>smiles</i> and <i>mol</i> used in following preprocessing.
<code>process</code> (<i>filepath</i>)	

chainer_chemistry.dataset.preprocessors.SchNetPreprocessor

```
class chainer_chemistry.dataset.preprocessors.SchNetPreprocessor (max_atoms=-1,
                                                                    out_size=-1,
                                                                    add_Hs=False,
                                                                    kekulize=False)
```

SchNetPreprocessor

Parameters

- **max_atoms** (*int*) – Max number of atoms for each molecule, if the number of atoms is more than this value, this data is simply ignored. Setting negative value indicates no limit for max atoms.
- **out_size** (*int*) – It specifies the size of array returned by *get_input_features*. If the number of atoms in the molecule is less than this value, the returned arrays is padded to have fixed size. Setting negative value indicates do not pad returned array.
- **add_Hs** (*bool*) – If True, implicit Hs are added.
- **kekulize** (*bool*) – If True, Kekulizes the molecule.

`__init__` (*max_atoms=-1, out_size=-1, add_Hs=False, kekulize=False*)
 Initialize self. See help(type(self)) for accurate signature.

Methods

<code>__init__</code> (<i>max_atoms, out_size, add_Hs, kekulize</i>)	Initialize self.
<code>get_input_features</code> (<i>mol</i>)	get input features
<code>get_label</code> (<i>mol, label_names</i>)	Extracts label information from a molecule.
<code>prepare_smiles_and_mol</code> (<i>mol</i>)	Prepare <i>smiles</i> and <i>mol</i> used in following preprocessing.
<code>process</code> (<i>filepath</i>)	

chainer_chemistry.dataset.preprocessors.WeaveNetPreprocessor

class `chainer_chemistry.dataset.preprocessors.WeaveNetPreprocessor` (*max_atoms=20, add_Hs=True, use_fixed_atom_feature=False, atom_list=None, include_unknown_atom=False, kekulize=False*)

WeaveNet must have fixed-size atom list for now, zero_padding option is always set to True.

Parameters

- **max_atoms** (*int*) – Max number of atoms for each molecule, if the number of atoms is more than this value, this data is simply ignored. Setting negative value indicates no limit for max atoms.
- **add_Hs** (*bool*) – If True, implicit Hs are added.
- **use_fixed_atom_feature** (*bool*) – If True, atom feature is extracted used in original paper. If it is False, atomic number is used instead.
- **atom_list** (*list*) – list of atoms to extract feature. If None, default *ATOM* is used as *atom_list*
- **include_unknown_atom** (*bool*) – If False, when the *mol* includes atom which is not in *atom_list*, it will raise *MolFeatureExtractionError*. If True, even the atom is not in *atom_list*, *atom_type* is set as “unknown” atom.
- **kekulize** (*bool*) – If True, Kekulizes the molecule.

`__init__` (*max_atoms=20, add_Hs=True, use_fixed_atom_feature=False, atom_list=None, include_unknown_atom=False, kekulize=False*)
 Initialize self. See help(type(self)) for accurate signature.

Methods

<code>__init__</code> (<i>max_atoms, add_Hs, ...</i>)	Initialize self.
<code>get_input_features</code> (<i>mol</i>)	get input features for WeaveNet

Continued on next page

Table 22 – continued from previous page

<code>get_label(mol[, label_names])</code>	Extracts label information from a molecule.
<code>prepare_smiles_and_mol(mol)</code>	Prepare <i>smiles</i> and <i>mol</i> used in following preprocessing.
<code>process(filepath)</code>	

chainer_chemistry.dataset.preprocessors.RelGATPreprocessor

```
class chainer_chemistry.dataset.preprocessors.RelGATPreprocessor (max_atoms=-1,
                                                                out_size=-1,
                                                                add_Hs=False)
```

RelGAT Preprocessor

Parameters

- **max_atoms** (*int*) – Max number of atoms for each molecule, if the number of atoms is more than this value, this data is simply ignored. Setting negative value indicates no limit for max atoms.
- **out_size** (*int*) – It specifies the size of array returned by *get_input_features*. If the number of atoms in the molecule is less than this value, the returned arrays is padded to have fixed size. Setting negative value indicates do not pad returned array.

__init__ (*max_atoms=-1, out_size=-1, add_Hs=False*)
Initialize self. See help(type(self)) for accurate signature.

Methods

<code>__init__([max_atoms, out_size, add_Hs])</code>	Initialize self.
<code>get_input_features(mol)</code>	get input features
<code>get_label(mol[, label_names])</code>	Extracts label information from a molecule.
<code>prepare_smiles_and_mol(mol)</code>	Prepare <i>smiles</i> and <i>mol</i> used in following preprocessing.
<code>process(filepath)</code>	

chainer_chemistry.dataset.preprocessors.RelGCNPreprocessor

```
class chainer_chemistry.dataset.preprocessors.RelGCNPreprocessor (max_atoms=-1,
                                                                out_size=-1,
                                                                add_Hs=False,
                                                                kekulize=False)
```

RelGCN Preprocessor

Parameters

- **max_atoms** (*int*) – Max number of atoms for each molecule, if the number of atoms is more than this value, this data is simply ignored. Setting negative value indicates no limit for max atoms.
- **out_size** (*int*) – It specifies the size of array returned by *get_input_features*. If the number of atoms in the molecule is less than this value, the returned arrays is padded to have fixed size. Setting negative value indicates do not pad returned array.

- **add_Hs** (*bool*) – If True, implicit Hs are added.
- **kekulize** (*bool*) – If True, Kekulizes the molecule.

__init__ (*max_atoms=-1, out_size=-1, add_Hs=False, kekulize=False*)
Initialize self. See help(type(self)) for accurate signature.

Methods

__init__ (<i>max_atoms, out_size, add_Hs, kekulize</i>)	Initialize self.
get_input_features (<i>mol</i>)	get input features
get_label (<i>mol[, label_names]</i>)	Extracts label information from a molecule.
prepare_smiles_and_mol (<i>mol</i>)	Prepare <i>smiles</i> and <i>mol</i> used in following preprocessing.
process (<i>filepath</i>)	

chainer_chemistry.dataset.preprocessors.RSGCNPreprocessor

class chainer_chemistry.dataset.preprocessors.**RSGCNPreprocessor** (*max_atoms=-1, out_size=-1, add_Hs=False, kekulize=False*)

RSGCN Preprocessor

Parameters

- **max_atoms** (*int*) – Max number of atoms for each molecule, if the number of atoms is more than this value, this data is simply ignored. Setting negative value indicates no limit for max atoms.
- **out_size** (*int*) – It specifies the size of array returned by *get_input_features*. If the number of atoms in the molecule is less than this value, the returned arrays is padded to have fixed size. Setting negative value indicates do not pad returned array.
- **add_Hs** (*bool*) – If True, implicit Hs are added.
- **kekulize** (*bool*) – If True, Kekulizes the molecule.

__init__ (*max_atoms=-1, out_size=-1, add_Hs=False, kekulize=False*)
Initialize self. See help(type(self)) for accurate signature.

Methods

__init__ (<i>max_atoms, out_size, add_Hs, kekulize</i>)	Initialize self.
get_input_features (<i>mol</i>)	get input features
get_label (<i>mol[, label_names]</i>)	Extracts label information from a molecule.
prepare_smiles_and_mol (<i>mol</i>)	Prepare <i>smiles</i> and <i>mol</i> used in following preprocessing.
process (<i>filepath</i>)	

Utilities

<code>chainer_chemistry. dataset.preprocessors. MolFeatureExtractionError</code>	
<code>chainer_chemistry.dataset. preprocessors.type_check_num_atoms</code>	Check number of atoms in <i>mol</i> does not exceed <i>num_max_atoms</i>
<code>chainer_chemistry. dataset.preprocessors. construct_atomic_number_array</code>	Returns atomic numbers of atoms consisting a molecule.
<code>chainer_chemistry.dataset. preprocessors.construct_adj_matrix</code>	Returns the adjacent matrix of the given molecule.

chainer_chemistry.dataset.preprocessors.MolFeatureExtractionError

exception `chainer_chemistry.dataset.preprocessors.MolFeatureExtractionError`

chainer_chemistry.dataset.preprocessors.type_check_num_atoms

`chainer_chemistry.dataset.preprocessors.type_check_num_atoms` (*mol*,
num_max_atoms=-1)

Check number of atoms in *mol* does not exceed *num_max_atoms*

If number of atoms in *mol* exceeds the number *num_max_atoms*, it will raise *MolFeatureExtractionError* exception.

Parameters

- **mol** (*Mol*) –
- **num_max_atoms** (*int*) – If negative value is set, not check number of atoms.

chainer_chemistry.dataset.preprocessors.construct_atomic_number_array

`chainer_chemistry.dataset.preprocessors.construct_atomic_number_array` (*mol*,
out_size=-1)

Returns atomic numbers of atoms consisting a molecule.

Parameters

- **mol** (*rdkit.Chem.Mol*) – Input molecule.
- **out_size** (*int*) – The size of returned array. If this option is negative, it does not take any effect. Otherwise, it must be larger than the number of atoms in the input molecules. In that case, the tail of the array is padded with zeros.

Returns

an array consisting of atomic numbers of atoms in the molecule.

Return type `numpy.ndarray`

chainer_chemistry.dataset.preprocessors.construct_adj_matrix

`chainer_chemistry.dataset.preprocessors.construct_adj_matrix(mol, out_size=-1, self_connection=True)`

Returns the adjacent matrix of the given molecule.

This function returns the adjacent matrix of the given molecule. Contrary to the specification of `rdkit.Chem.rdmolops.GetAdjacencyMatrix()`, The diagonal entries of the returned matrix are all-one.

Parameters

- **mol** (`rdkit.Chem.Mol`) – Input molecule.
- **out_size** (`int`) – The size of the returned matrix. If this option is negative, it does not take any effect. Otherwise, it must be larger than the number of atoms in the input molecules. In that case, the adjacent matrix is expanded and zeros are padded to right columns and bottom rows.
- **self_connection** (`bool`) – Add self connection or not. If True, diagonal element of adjacency matrix is filled with 1.

Returns

The adjacent matrix of the input molecule. It is 2-dimensional array with shape (atoms1, atoms2), where atoms1 & atoms2 represent from and to of the edge respectively. If `out_size` is non-negative, the returned its size is equal to that value. Otherwise, it is equal to the number of atoms in the the molecule.

Return type `adj_array` (`numpy.ndarray`)

Splitters

<code>chainer_chemistry.dataset.splitters.RandomSplitter</code>	Class for doing random data splits.
<code>chainer_chemistry.dataset.splitters.StratifiedSplitter</code>	Class for doing stratified data splits.
<code>chainer_chemistry.dataset.splitters.ScaffoldSplitter</code>	Class for doing data splits by chemical scaffold.

chainer_chemistry.dataset.splitters.RandomSplitter

class `chainer_chemistry.dataset.splitters.RandomSplitter`

Class for doing random data splits.

__init__()

Initialize self. See `help(type(self))` for accurate signature.

Methods

<code>k_fold_split(dataset, k)</code>	
<code>train_valid_split(dataset[, frac_train, ...])</code>	Generate indices to split data into train and valid set.
<code>train_valid_test_split(dataset[, ...])</code>	Generate indices to split data into train, valid and test set.

chainer_chemistry.dataset.splitters.StratifiedSplitter**class** chainer_chemistry.dataset.splitters.StratifiedSplitter

Class for doing stratified data splits.

__init__()

Initialize self. See help(type(self)) for accurate signature.

Methods

k_fold_split(dataset, k)

train_valid_split(dataset[, labels, ...]) Split dataset into train and valid set.

train_valid_test_split(dataset[, labels, ...]) Split dataset into train, valid and test set.

chainer_chemistry.dataset.splitters.ScaffoldSplitter**class** chainer_chemistry.dataset.splitters.ScaffoldSplitter

Class for doing data splits by chemical scaffold.

Referred Deepchem for the implementation, <https://git.io/fXzF4>**__init__**()

Initialize self. See help(type(self)) for accurate signature.

Methods

k_fold_split(dataset, k)

train_valid_split(dataset, smiles_list[, ...]) Split dataset into train and valid set.

train_valid_test_split(dataset, smiles_list) Split dataset into train, valid and test set.

1.5.2 Datasets**Dataset implementations**

*chainer_chemistry.datasets.
NumpyTupleDataset*

Dataset of a tuple of datasets.

chainer_chemistry.datasets.NumpyTupleDataset**class** chainer_chemistry.datasets.NumpyTupleDataset (*datasets)

Dataset of a tuple of datasets.

It combines multiple datasets into one dataset. Each example is represented by a tuple whose *i*-th item corresponds to the *i*-th dataset. And each *i*-th dataset is expected to be an instance of numpy.ndarray.**Parameters datasets** – Underlying datasets. The *i*-th one is used for the *i*-th item of each example. All datasets must have the same length.**__init__**(*datasets)

Initialize self. See `help(type(self))` for accurate signature.

Methods

<code>__init__(*datasets)</code>	Initialize self.
<code>get_datasets()</code>	
<code>load(filepath)</code>	
<code>save(filepath, numpy_tuple_dataset)</code>	save the dataset to filepath in npz format

Attributes

<code>features</code>	Extract features according to the specified index.
-----------------------	--

Dataset loaders

<code>chainer_chemistry.datasets.tox21.get_tox21</code>	Downloads, caches and preprocesses Tox21 dataset.
<code>chainer_chemistry.datasets.qm9.get_qm9</code>	Downloads, caches and preprocesses QM9 dataset.
<code>chainer_chemistry.datasets.molnet.get_molnet_dataset</code>	Downloads, caches and preprocess MoleculeNet dataset.
<code>chainer_chemistry.datasets.molnet.get_molnet_dataframe</code>	Downloads, caches and get the dataframe of MoleculeNet dataset.

chainer_chemistry.datasets.tox21.get_tox21

`chainer_chemistry.datasets.tox21.get_tox21` (*preprocessor=None*, *labels=None*, *return_smiles=False*, *train_target_index=None*, *val_target_index=None*, *test_target_index=None*)

Downloads, caches and preprocesses Tox21 dataset.

Parameters

- **preprocessor** (`BasePreprocessor`) – Preprocessor. This should be chosen based on the network to be trained. If it is `None`, default `AtomicNumberPreprocessor` is used.
- **labels** (*str or list*) – List of target labels.
- **return_smiles** (*bool*) – If set to `True`, smiles array is also returned.
- **train_target_index** (*list or None*) – target index list to partially extract train dataset. If `None` (default), all examples are parsed.
- **val_target_index** (*list or None*) – target index list to partially extract val dataset. If `None` (default), all examples are parsed.
- **test_target_index** (*list or None*) – target index list to partially extract test dataset. If `None` (default), all examples are parsed.

Returns The 3-tuple consisting of train, validation and test datasets, respectively. Each dataset is composed of *features*, which depends on *preprocess_method*.

chainer_chemistry.datasets.qm9.get_qm9

```
chainer_chemistry.datasets.qm9.get_qm9(preprocessor=None, labels=None, re-
                                         turn_smiles=False, target_index=None)
```

Downloads, caches and preprocesses QM9 dataset.

Parameters

- **preprocessor** (*BasePreprocessor*) – Preprocessor. This should be chosen based on the network to be trained. If it is *None*, default *AtomicNumberPreprocessor* is used.
- **labels** (*str* or *list*) – List of target labels.
- **return_smiles** (*bool*) – If set to *True*, smiles array is also returned.
- **target_index** (*list* or *None*) – target index list to partially extract dataset. If *None* (default), all examples are parsed.

Returns dataset, which is composed of *features*, which depends on *preprocess_method*.

chainer_chemistry.datasets.molnet.get_molnet_dataset

```
chainer_chemistry.datasets.molnet.get_molnet_dataset(dataset_name, pre-
                                                       processor=None, la-
                                                       bels=None, split=None,
                                                       frac_train=0.8, frac_valid=0.1,
                                                       frac_test=0.1, seed=777,
                                                       return_smiles=False,
                                                       return_pdb_id=False,
                                                       target_index=None,
                                                       task_index=0, **kwargs)
```

Downloads, caches and preprocess MoleculeNet dataset.

Parameters

- **dataset_name** (*str*) – MoleculeNet dataset name. If you want to know the detail of MoleculeNet, please refer to [official site](#) If you would like to know what dataset_name is available for chainer_chemistry, please refer to *molnet_config.py*.
- **preprocessor** (*BasePreprocessor*) – Preprocessor. It should be chosen based on the network to be trained. If it is *None*, default *AtomicNumberPreprocessor* is used.
- **labels** (*str* or *list*) – List of target labels.
- **split** (*str* or *BaseSplitter* or *None*) – How to split dataset into train, validation and test. If *None*, this functions use the splitter that is recommended by MoleculeNet. Additionally You can use an instance of *BaseSplitter* or choose it from ‘random’, ‘stratified’ and ‘scaffold’.
- **return_smiles** (*bool*) – If set to *True*, smiles array is also returned.
- **return_pdb_id** (*bool*) – If set to *True*, PDB ID array is also returned. This argument is only used when you select ‘pdbbind_smiles’.
- **target_index** (*list* or *None*) – target index list to partially extract dataset. If *None* (default), all examples are parsed.
- **task_index** (*int*) – Target task index in dataset for stratification. (Stratified Splitter only)

Returns (dict): Dictionary that contains dataset that is already split into train, valid and test dataset and 1-d numpy array with dtype=object(string) which is a vector of smiles for each example or *None*.

chainer_chemistry.datasets.molnet.get_molnet_dataframe

chainer_chemistry.datasets.molnet.get_molnet_dataframe(*dataset_name*, *pdbbind_subset=None*)

Downloads, caches and get the dataframe of MoleculeNet dataset.

Parameters

- **dataset_name** (*str*) – MoleculeNet dataset name. If you want to know the detail of MoleculeNet, please refer to [official site](#) If you would like to know what dataset_name is available for chainer_chemistry, please refer to *molnet_config.py*.
- **pdcbind_subset** (*str*) – PDBbind dataset subset name. If you want to know the detail of subset, please refer to *official site* <http://www.pdcbind.org.cn/download/pdcbind_2017_intro.pdf>

Returns (pandas.DataFrame or tuple): DataFrame of dataset without any preprocessing. When the files of dataset are separated, this function returns multiple DataFrame.

1.5.3 Functions

Function implementations

<code>chainer_chemistry.functions.matmul</code>	Computes the matrix multiplication of two arrays.
<code>chainer_chemistry.functions.mean_squared_error</code>	Mean squared error function.
<code>chainer_chemistry.functions.mean_absolute_error</code>	Mean absolute error function.
<code>chainer_chemistry.functions.r2_score</code>	Computes R ² (coefficient of determination) regression score function.

chainer_chemistry.functions.matmul

chainer_chemistry.functions.matmul(*a*, *b*, *transa=False*, *transb=False*)

Computes the matrix multiplication of two arrays.

Parameters

- **a** (*Variable*) – The left operand of the matrix multiplication. If *a* and *b* are both 1-D arrays, *matmul* returns a dot product of vector *a* and vector *b*. If 2-D arrays, *matmul* returns matrix product of *a* and *b*. If arrays' dimension is larger than 2, they are treated as a stack of matrices residing in the last two indexes. *matmul* returns a stack of each two arrays. *a* and *b* must have the same dimension.
- **b** (*Variable*) – The right operand of the matrix multiplication. Its array is treated as a matrix in the same way as *a*'s array.
- **transa** (*bool*) – If *True*, each matrices in *a* will be transposed. If *a.ndim == 1*, do nothing.
- **transb** (*bool*) – If *True*, each matrices in *b* will be transposed. If *b.ndim == 1*, do nothing.

Returns The result of the matrix multiplication.

Return type Variable

Example

```
>>> a = np.array([[1, 0], [0, 1]], 'f')
>>> b = np.array([[4, 1], [2, 2]], 'f')
>>> F.matmul(a, b).data
array([[ 4.,  1.],
       [ 2.,  2.]], dtype=float32)
```

chainer_chemistry.functions.mean_squared_error

chainer_chemistry.functions.**mean_squared_error**(x0, x1, ignore_nan=False)

Mean squared error function.

This function computes mean squared error between two variables. The mean is taken over the minibatch. Note that the error is not scaled by 1/2.

Parameters

- **x0** (Variable or numpy.ndarray or cupy.ndarray) – Input variable.
- **x1** (Variable or numpy.ndarray or cupy.ndarray) – Input variable.
- **ignore_nan** (bool) – If *True*, this function compute mean squared error ignoring NaNs. The arithmetic mean is the sum of the non-NaN elements along the axis divided by the number of whole elements.

Returns A variable holding an array representing the mean squared error of two inputs.

Return type Variable

chainer_chemistry.functions.mean_absolute_error

chainer_chemistry.functions.**mean_absolute_error**(x0, x1, ignore_nan=False)

Mean absolute error function.

This function computes mean absolute error between two variables. The mean is taken over the minibatch.

Parameters

- **x0** (Variable or numpy.ndarray or cupy.ndarray) – Input variable.
- **x1** (Variable or numpy.ndarray or cupy.ndarray) – Input variable.
- **ignore_nan** (bool) – If *True*, this function compute mean absolute error ignoring NaNs. The arithmetic mean is the sum of the non-NaN elements along the axis divided by the number of whole elements.

Returns A variable holding an array representing the mean absolute error of two inputs.

Return type Variable

chainer_chemistry.functions.r2_score

`chainer_chemistry.functions.r2_score(pred, true, sample_weight=None, multioutput='uniform_average', ignore_nan=False)`

Computes R^2 (coefficient of determination) regression score function. :param pred: Variable holding a vector, matrix or tensor of

estimated target values.

Parameters

- **true** (*Variable*) – Variable holding a vector, matrix or tensor of correct target values.
- **sample_weight** – This argument is for compatibility with scikit-learn’s implementation of `r2_score`. Current implementation admits `None` only.
- **multioutput** (*string*) – [‘uniform_average’, ‘raw_values’]. If ‘uniform_average’, this function returns an average of R^2 score of multiple output. If ‘raw_average’, this function return a set of R^2 score of multiple output.

Returns A Variable holding a scalar array of the R^2 score if ‘multioutput’ is ‘uniform_average’ or a vector of R^2 scores if ‘multioutput’ is ‘raw_values’.

Return type Variable

Note: This function is non-differentiable.

1.5.4 Iterators

Iterator Implementations

<code>chainer_chemistry.iterators.BalancedSerialIterator</code>	Dataset iterator that serially reads the examples with balancing label.
<code>chainer_chemistry.iterators.IndexIterator</code>	Index iterator

chainer_chemistry.iterators.BalancedSerialIterator

```
class chainer_chemistry.iterators.BalancedSerialIterator(dataset, batch_size,  
                                                         labels, repeat=True,  
                                                         shuffle=True,  
                                                         batch_balancing=False,  
                                                         ignore_labels=None,  
                                                         logger=<Logger  
chainer_chemistry.iterators.balanced_serial_iterator>  
                                                         (WARNING)>)
```

Dataset iterator that serially reads the examples with balancing label.

Parameters

- **dataset** – Dataset to iterate.
- **batch_size** (*int*) – Number of examples within each minibatch.

- **labels** (*list* or *numpy.ndarray*) – 1d array which specifies label feature of *dataset*. Its size must be same as the length of *dataset*.
- **repeat** (*bool*) – If `True`, it infinitely loops over the dataset. Otherwise, it stops iteration at the end of the first epoch.
- **shuffle** (*bool*) – If `True`, the order of examples is shuffled at the beginning of each epoch. Otherwise, the order is permanently same as that of *dataset*.
- **batch_balancing** (*bool*) – If `True`, examples are sampled in the way that each label examples are roughly evenly sampled in each minibatch. Otherwise, the iterator only guarantees that total numbers of examples are same among label features.
- **ignore_labels** (*int* or *list* or *None*) – Labels to be ignored. If not `None`, the example whose label is in *ignore_labels* are not sampled by this iterator.

`__init__(dataset, batch_size, labels, repeat=True, shuffle=True, batch_balancing=False, ignore_labels=None, logger=<Logger chainer_chemistry.iterators.balanced_serial_iterator (WARNING)>)`
 Initialize self. See help(type(self)) for accurate signature.

Methods

<code>__init__(dataset, batch_size, labels[, ...])</code>	Initialize self.
<code>finalize()</code>	Finalizes the iterator and possibly releases the resources.
<code>next()</code>	Returns the next batch.
<code>reset()</code>	
<code>serialize(serializer)</code>	Serializes the internal state of the iterator.
<code>show_label_stats()</code>	

Attributes

<code>epoch_detail</code>
<code>previous_epoch_detail</code>

chainer_chemistry.iterators.IndexIterator

class `chainer_chemistry.iterators.IndexIterator` (*index_list*, *shuffle=True*, *num=0*)
 Index iterator

`IndexIterator` is used internally in `BalancedSerialIterator`, as each label's index iterator

Parameters

- **index_list** (*list*) – list of int which represents indices.
- **shuffle** (*bool*) – shuffle flag. If `True`, indices specified by *index_list* will be randomly shuffled.
- **num** (*int*) – number of indices to be extracted when `__next__` is called.

`__init__(index_list, shuffle=True, num=0)`
 Initialize self. See help(type(self)) for accurate signature.

Methods

<code>__init__(index_list[, shuffle, num])</code>	Initialize self.
<code>finalize()</code>	Finalizes the iterator and possibly releases the resources.
<code>get_next_indices(num)</code>	get next indices
<code>next()</code>	Python2 alternative of <code>__next__</code> .
<code>serialize(serializer)</code>	Serializes the internal state of the iterator.
<code>update_current_index_list()</code>	

1.5.5 Links

Link implementations

<code>chainer_chemistry.links. EmbedAtomID(out_size)</code>	Embedding specialized to atoms.
<code>chainer_chemistry.links. GraphLinear(in_size)</code>	Graph Linear layer.
<code>chainer_chemistry.links. GraphBatchNormalization([...])</code>	Graph Batch Normalization layer.

Scaler implementations

<code>chainer_chemistry.links. StandardScaler()</code>	
--	--

Update implementations

<code>chainer_chemistry.links. GGNNUpdate([...])</code>	GGNN submodule for update part.
<code>chainer_chemistry.links.NFPUpdate(...[, ...])</code>	NFP submodule for update part.
<code>chainer_chemistry.links. RelGATUpdate(...[, ...])</code>	RelGAT submodule for update part.
<code>chainer_chemistry.links. RelGUNUpdate(...[, ...])</code>	RelGUN submodule for update part.
<code>chainer_chemistry.links. RSGCNUpdate(...)</code>	RSGCN submodule for message and update part.
<code>chainer_chemistry.links. SchNetUpdate([...])</code>	

Readout implementations

<code>chainer_chemistry.links. GeneralReadout([...])</code>	General submodule for readout part.
<code>chainer_chemistry.links. GGNNReadout(out_dim)</code>	GGNN submodule for readout part.

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<code>chainer_chemistry.links.NFPReadout(...)</code>	NFP submodule for readout part.
<code>chainer_chemistry.links.SchNetReadout([...])</code>	SchNet submodule for readout part.

1.5.6 Models

Model implementations

<code>chainer_chemistry.models.NFP</code>	Neural Finger Print (NFP)
<code>chainer_chemistry.models.GGNN</code>	Gated Graph Neural Networks (GGNN)
<code>chainer_chemistry.models.MLP</code>	Basic implementation for MLP
<code>chainer_chemistry.models.SchNet</code>	See Kristof et al, SchNet: A continuous-filter convolutional neural network for modeling
<code>chainer_chemistry.models.WeaveNet</code>	WeaveNet implementation
<code>chainer_chemistry.models.RelGAT</code>	Relational Graph Attention Networks (GAT)
<code>chainer_chemistry.models.RelGCN</code>	Relational GCN (RelGCN)
<code>chainer_chemistry.models.RSGCN</code>	Renormalized Spectral Graph Convolutional Network (RSGCN)

chainer_chemistry.models.NFP

class `chainer_chemistry.models.NFP` (*out_dim*, *hidden_dim*=16, *n_layers*=4, *max_degree*=6, *n_atom_types*=117, *concat_hidden*=False)

Neural Finger Print (NFP)

See: David K Duvenaud, Dougal Maclaurin, Jorge Iparraguirre, Rafael Bombarell, Timothy Hirzel, Alan Aspuru-Guzik, and Ryan P Adams (2015). Convolutional networks on graphs for learning molecular fingerprints. *Advances in Neural Information Processing Systems (NIPS)* 28,

Parameters

- **out_dim** (*int*) – dimension of output feature vector
- **hidden_dim** (*int*) – dimension of feature vector associated to each atom
- **n_layers** (*int*) – number of layers
- **max_degree** (*int*) – max degree of atoms when molecules are regarded as graphs
- **n_atom_types** (*int*) – number of types of atoms
- **concat_hidden** (*bool*) – If set to True, readout is executed in each layer and the result is concatenated

__init__ (*out_dim*, *hidden_dim*=16, *n_layers*=4, *max_degree*=6, *n_atom_types*=117, *concat_hidden*=False)

Initialize self. See help(type(self)) for accurate signature.

Methods

__init__ (<i>out_dim</i> [, <i>hidden_dim</i> , <i>n_layers</i> , ...])	Initialize self.
---	------------------

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Table 45 – continued from previous page

<code>add_hook(hook[, name])</code>	Registers a link hook.
<code>add_link(name, link)</code>	Registers a child link to this chain.
<code>add_param(name[, shape, dtype, initializer])</code>	Registers a parameter to the link.
<code>add_persistent(name, value)</code>	Registers a persistent value to the link.
<code>addgrads(link)</code>	Accumulates gradient values from given link.
<code>children()</code>	Returns a generator of all child links.
<code>cleargrads()</code>	Clears all gradient arrays.
<code>copy([model])</code>	Copies the link hierarchy to new one.
<code>copyparams(link[, copy_persistent])</code>	Copies all parameters from given link.
<code>count_params()</code>	Counts the total number of parameters.
<code>delete_hook(name)</code>	Unregisters the link hook.
<code>disable_update()</code>	Disables update rules of all parameters under the link hierarchy.
<code>enable_update()</code>	Enables update rules of all parameters under the link hierarchy.
<code>init_scope()</code>	Creates an initialization scope.
<code>links([skipself])</code>	Returns a generator of all links under the hierarchy.
<code>namedlinks([skipself])</code>	Returns a generator of all (path, link) pairs under the hierarchy.
<code>namedparams([include_uninit])</code>	Returns a generator of all (path, param) pairs under the hierarchy.
<code>params([include_uninit])</code>	Returns a generator of all parameters under the link hierarchy.
<code>register_persistent(name)</code>	Registers an attribute of a given name as a persistent value.
<code>repeat(n_repeat[, mode])</code>	Repeats this link multiple times to make a <code>Sequential</code> .
<code>serialize(serializer)</code>	Serializes the link object.
<code>to_cpu()</code>	Copies parameter variables and persistent values to CPU.
<code>to_gpu([device])</code>	Copies parameter variables and persistent values to GPU.
<code>to_intel64()</code>	Copies parameter variables and persistent values to CPU.
<code>zerograds()</code>	Initializes all gradient arrays by zero.

Attributes

<code>local_link_hooks</code>	Ordered dictionary of registered link hooks.
<code>update_enabled</code>	True if at least one parameter has an update rule enabled.
<code>within_init_scope</code>	True if the current code is inside of an initialization scope.
<code>xp</code>	Array module for this link.

chainer_chemistry.models.GGNN

```
class chainer_chemistry.models.GGNN(out_dim,          hidden_dim=16,          n_layers=4,
                                   n_atom_types=117,      concat_hidden=False,
                                   weight_lying=True,    activation=<function identity>,
                                   num_edge_type=4)
```

Gated Graph Neural Networks (GGNN)

See: Li, Y., Tarlow, D., Brockschmidt, M., & Zemel, R. (2015). Gated graph sequence neural networks. [arXiv:1511.05493](https://arxiv.org/abs/1511.05493)

Parameters

- **out_dim** (*int*) – dimension of output feature vector
- **hidden_dim** (*int*) – dimension of feature vector associated to each atom
- **n_layers** (*int*) – number of layers
- **n_atom_types** (*int*) – number of types of atoms
- **concat_hidden** (*bool*) – If set to True, readout is executed in each layer and the result is concatenated
- **weight_lying** (*bool*) – enable weight_lying or not
- **activation** (*Function or FunctionNode*) – activate function
- **num_edge_type** (*int*) – number of edge type. Defaults to 4 for single, double, triple and aromatic bond.

__init__ (out_dim, hidden_dim=16, n_layers=4, n_atom_types=117, concat_hidden=False, weight_lying=True, activation=<function identity>, num_edge_type=4)
Initialize self. See help(type(self)) for accurate signature.

Methods

__init__ (out_dim[, hidden_dim, n_layers, ...])	Initialize self.
add_hook (hook[, name])	Registers a link hook.
add_link (name, link)	Registers a child link to this chain.
add_param (name[, shape, dtype, initializer])	Registers a parameter to the link.
add_persistent (name, value)	Registers a persistent value to the link.
addgrads (link)	Accumulates gradient values from given link.
children ()	Returns a generator of all child links.
cleargrads ()	Clears all gradient arrays.
copy ([model])	Copies the link hierarchy to new one.
copyparams (link[, copy_persistent])	Copies all parameters from given link.
count_params ()	Counts the total number of parameters.
delete_hook (name)	Unregisters the link hook.
disable_update ()	Disables update rules of all parameters under the link hierarchy.
enable_update ()	Enables update rules of all parameters under the link hierarchy.
init_scope ()	Creates an initialization scope.
links ([skipself])	Returns a generator of all links under the hierarchy.

Continued on next page

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<code>namedlinks([skipself])</code>	Returns a generator of all (path, link) pairs under the hierarchy.
<code>namedparams([include_uninit])</code>	Returns a generator of all (path, param) pairs under the hierarchy.
<code>params([include_uninit])</code>	Returns a generator of all parameters under the link hierarchy.
<code>register_persistent(name)</code>	Registers an attribute of a given name as a persistent value.
<code>repeat(n_repeat[, mode])</code>	Repeats this link multiple times to make a <code>Sequential</code> .
<code>reset_state()</code>	
<code>serialize(serializer)</code>	Serializes the link object.
<code>to_cpu()</code>	Copies parameter variables and persistent values to CPU.
<code>to_gpu([device])</code>	Copies parameter variables and persistent values to GPU.
<code>to_intel64()</code>	Copies parameter variables and persistent values to CPU.
<code>zerograds()</code>	Initializes all gradient arrays by zero.

Attributes

<code>local_link_hooks</code>	Ordered dictionary of registered link hooks.
<code>update_enabled</code>	True if at least one parameter has an update rule enabled.
<code>within_init_scope</code>	True if the current code is inside of an initialization scope.
<code>xp</code>	Array module for this link.

chainer_chemistry.models.MLP

```
class chainer_chemistry.models.MLP(out_dim, hidden_dim=16, n_layers=2, activation=<function relu>)
```

Basic implementation for MLP

Parameters

- **out_dim** (*int*) – dimension of output feature vector
- **hidden_dim** (*int*) – dimension of feature vector associated to each atom
- **n_layers** (*int*) – number of layers
- **activation** (*chainer.functions*) – activation function

```
__init__(out_dim, hidden_dim=16, n_layers=2, activation=<function relu>)
```

Initialize self. See `help(type(self))` for accurate signature.

Methods

<code>__init__(out_dim[, hidden_dim, n_layers, ...])</code>	Initialize self.
<code>add_hook(hook[, name])</code>	Registers a link hook.

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<code>add_link(name, link)</code>	Registers a child link to this chain.
<code>add_param(name[, shape, dtype, initializer])</code>	Registers a parameter to the link.
<code>add_persistent(name, value)</code>	Registers a persistent value to the link.
<code>addgrads(link)</code>	Accumulates gradient values from given link.
<code>children()</code>	Returns a generator of all child links.
<code>cleargrads()</code>	Clears all gradient arrays.
<code>copy([model])</code>	Copies the link hierarchy to new one.
<code>copyparams(link[, copy_persistent])</code>	Copies all parameters from given link.
<code>count_params()</code>	Counts the total number of parameters.
<code>delete_hook(name)</code>	Unregisters the link hook.
<code>disable_update()</code>	Disables update rules of all parameters under the link hierarchy.
<code>enable_update()</code>	Enables update rules of all parameters under the link hierarchy.
<code>init_scope()</code>	Creates an initialization scope.
<code>links([skipself])</code>	Returns a generator of all links under the hierarchy.
<code>namedlinks([skipself])</code>	Returns a generator of all (path, link) pairs under the hierarchy.
<code>namedparams([include_uninit])</code>	Returns a generator of all (path, param) pairs under the hierarchy.
<code>params([include_uninit])</code>	Returns a generator of all parameters under the link hierarchy.
<code>register_persistent(name)</code>	Registers an attribute of a given name as a persistent value.
<code>repeat(n_repeat[, mode])</code>	Repeats this link multiple times to make a <code>Sequential</code> .
<code>serialize(serializer)</code>	Serializes the link object.
<code>to_cpu()</code>	Copies parameter variables and persistent values to CPU.
<code>to_gpu([device])</code>	Copies parameter variables and persistent values to GPU.
<code>to_intel64()</code>	Copies parameter variables and persistent values to CPU.
<code>zerograds()</code>	Initializes all gradient arrays by zero.

Attributes

<code>local_link_hooks</code>	Ordered dictionary of registered link hooks.
<code>update_enabled</code>	True if at least one parameter has an update rule enabled.
<code>within_init_scope</code>	True if the current code is inside of an initialization scope.
<code>xp</code>	Array module for this link.

chainer_chemistry.models.SchNet

```
class chainer_chemistry.models.SchNet(out_dim=1, hidden_dim=64, n_layers=3, read-
out_hidden_dim=32, n_atom_types=117, con-
cat_hidden=False)
```

See Kristof et al, SchNet: A continuous-filter convolutional neural network for modeling quantum inter-

actions. [arXiv:1706.08566](#)

Parameters

- **out_dim** (*int*) – dimension of output feature vector
- **hidden_dim** (*int*) – dimension of feature vector associated to each atom
- **n_layers** (*int*) – number of layers
- **readout_hidden_dim** (*int*) – dimension of feature vector associated to each molecule
- **n_atom_types** (*int*) – number of types of atoms
- **concat_hidden** (*bool*) – If set to True, readout is executed in each layer and the result is concatenated

__init__ (*out_dim=1, hidden_dim=64, n_layers=3, readout_hidden_dim=32, n_atom_types=117, concat_hidden=False*)
Initialize self. See help(type(self)) for accurate signature.

Methods

__init__ ([out_dim, hidden_dim, n_layers, ...])	Initialize self.
add_hook (hook[, name])	Registers a link hook.
add_link (name, link)	Registers a child link to this chain.
add_param (name[, shape, dtype, initializer])	Registers a parameter to the link.
add_persistent (name, value)	Registers a persistent value to the link.
addgrads (link)	Accumulates gradient values from given link.
children ()	Returns a generator of all child links.
cleargrads ()	Clears all gradient arrays.
copy ([mode])	Copies the link hierarchy to new one.
copyparams (link[, copy_persistent])	Copies all parameters from given link.
count_params ()	Counts the total number of parameters.
delete_hook (name)	Unregisters the link hook.
disable_update ()	Disables update rules of all parameters under the link hierarchy.
enable_update ()	Enables update rules of all parameters under the link hierarchy.
init_scope ()	Creates an initialization scope.
links ([skipself])	Returns a generator of all links under the hierarchy.
namedlinks ([skipself])	Returns a generator of all (path, link) pairs under the hierarchy.
namedparams ([include_uninit])	Returns a generator of all (path, param) pairs under the hierarchy.
params ([include_uninit])	Returns a generator of all parameters under the link hierarchy.
register_persistent (name)	Registers an attribute of a given name as a persistent value.
repeat (n_repeat[, mode])	Repeats this link multiple times to make a Sequential.
serialize (serializer)	Serializes the link object.
to_cpu ()	Copies parameter variables and persistent values to CPU.

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<code>to_gpu([device])</code>	Copies parameter variables and persistent values to GPU.
<code>to_intel64()</code>	Copies parameter variables and persistent values to CPU.
<code>zerograds()</code>	Initializes all gradient arrays by zero.

Attributes

<code>local_link_hooks</code>	Ordered dictionary of registered link hooks.
<code>update_enabled</code>	True if at least one parameter has an update rule enabled.
<code>within_init_scope</code>	True if the current code is inside of an initialization scope.
<code>xp</code>	Array module for this link.

chainer_chemistry.models.WeaveNet

class `chainer_chemistry.models.WeaveNet` (*weave_channels=None*, *hidden_dim=16*, *n_atom=20*, *n_sub_layer=1*, *n_atom_types=117*, *readout_mode='sum'*)

WeaveNet implementation

Parameters

- **`weave_channels`** (*list*) – list of int, output dimension for each weave module
- **`hidden_dim`** (*int*) – hidden dim
- **`n_atom`** (*int*) – number of atom of input array
- **`n_sub_layer`** (*int*) – number of layer for each *AtomToPair*, *PairToAtom* layer
- **`n_atom_types`** (*int*) – number of atom id
- **`readout_mode`** (*str*) – ‘sum’ or ‘max’ or ‘summax’

`__init__` (*weave_channels=None*, *hidden_dim=16*, *n_atom=20*, *n_sub_layer=1*, *n_atom_types=117*, *readout_mode='sum'*)
Initialize self. See `help(type(self))` for accurate signature.

Methods

<code>__init__([weave_channels, hidden_dim, ...])</code>	Initialize self.
<code>add_hook(hook[, name])</code>	Registers a link hook.
<code>add_link(name, link)</code>	Registers a child link to this chain.
<code>add_param(name[, shape, dtype, initializer])</code>	Registers a parameter to the link.
<code>add_persistent(name, value)</code>	Registers a persistent value to the link.
<code>addgrads(link)</code>	Accumulates gradient values from given link.
<code>children()</code>	Returns a generator of all child links.
<code>cleargrads()</code>	Clears all gradient arrays.
<code>copy([mode])</code>	Copies the link hierarchy to new one.
<code>copyparams(link[, copy_persistent])</code>	Copies all parameters from given link.
<code>count_params()</code>	Counts the total number of parameters.

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Table 53 – continued from previous page

<code>delete_hook(name)</code>	Unregisters the link hook.
<code>disable_update()</code>	Disables update rules of all parameters under the link hierarchy.
<code>enable_update()</code>	Enables update rules of all parameters under the link hierarchy.
<code>init_scope()</code>	Creates an initialization scope.
<code>links([skipself])</code>	Returns a generator of all links under the hierarchy.
<code>namedlinks([skipself])</code>	Returns a generator of all (path, link) pairs under the hierarchy.
<code>namedparams([include_uninit])</code>	Returns a generator of all (path, param) pairs under the hierarchy.
<code>params([include_uninit])</code>	Returns a generator of all parameters under the link hierarchy.
<code>register_persistent(name)</code>	Registers an attribute of a given name as a persistent value.
<code>repeat(n_repeat[, mode])</code>	Repeats this link multiple times to make a <code>Sequential</code> .
<code>serialize(serializer)</code>	Serializes the link object.
<code>to_cpu()</code>	Copies parameter variables and persistent values to CPU.
<code>to_gpu([device])</code>	Copies parameter variables and persistent values to GPU.
<code>to_intel64()</code>	Copies parameter variables and persistent values to CPU.
<code>zerograds()</code>	Initializes all gradient arrays by zero.

Attributes

<code>local_link_hooks</code>	Ordered dictionary of registered link hooks.
<code>update_enabled</code>	True if at least one parameter has an update rule enabled.
<code>within_init_scope</code>	True if the current code is inside of an initialization scope.
<code>xp</code>	Array module for this link.

chainer_chemistry.models.RelGAT

```
class chainer_chemistry.models.RelGAT(out_dim, hidden_dim=16, n_heads=3, negative_slope=0.2, n_edge_types=4, n_layers=4, dropout_ratio=-1.0, activation=<function identity>, n_atom_types=117, softmax_mode='across', concat_hidden=False, concat_heads=False, weight_tying=False)
```

Relational Graph Attention Networks (GAT)

See: Veličković, Petar, et al. (2017). Graph Attention Networks. *arXiv:1701.10903* <<https://arxiv.org/abs/1710.10903>> Dan <<https://openreview.net/forum?id=Bklzkh0qFm>>

Parameters

- `out_dim` (*int*) – dimension of output feature vector

- **hidden_dim** (*int*) – dimension of feature vector associated to each atom
- **n_layers** (*int*) – number of layers
- **n_atom_types** (*int*) – number of types of atoms
- **n_heads** (*int*) – number of multi-head-attentions.
- **n_edge_types** (*int*) – number of edge types.
- **dropout_ratio** (*float*) – dropout ratio of the normalized attention coefficients
- **negative_slope** (*float*) – LeakyRELU angle of the negative slope
- **softmax_mode** (*str*) – take the softmax over the logits ‘across’ or ‘within’ relation. If you would like to know the detail discussion, please refer Relational GAT paper.
- **concat_hidden** (*bool*) – If set to True, readout is executed in each layer and the result is concatenated
- **concat_heads** (*bool*) – Whether to concat or average multi-head attentions
- **weight_tying** (*bool*) – enable weight_tying or not

__init__ (*out_dim*, *hidden_dim*=16, *n_heads*=3, *negative_slope*=0.2, *n_edge_types*=4, *n_layers*=4, *dropout_ratio*=-1.0, *activation*=<function identity>, *n_atom_types*=117, *softmax_mode*=‘across’, *concat_hidden*=False, *concat_heads*=False, *weight_tying*=False)
Initialize self. See help(type(self)) for accurate signature.

Methods

__init__ (<i>out_dim</i> [, <i>hidden_dim</i> , <i>n_heads</i> , ...])	Initialize self.
add_hook (<i>hook</i> [, <i>name</i>])	Registers a link hook.
add_link (<i>name</i> , <i>link</i>)	Registers a child link to this chain.
add_param (<i>name</i> [, <i>shape</i> , <i>dtype</i> , <i>initializer</i>])	Registers a parameter to the link.
add_persistent (<i>name</i> , <i>value</i>)	Registers a persistent value to the link.
addgrads (<i>link</i>)	Accumulates gradient values from given link.
children ()	Returns a generator of all child links.
cleargrads ()	Clears all gradient arrays.
copy ([<i>mode</i>])	Copies the link hierarchy to new one.
copyparams (<i>link</i> [, <i>copy_persistent</i>])	Copies all parameters from given link.
count_params ()	Counts the total number of parameters.
delete_hook (<i>name</i>)	Unregisters the link hook.
disable_update ()	Disables update rules of all parameters under the link hierarchy.
enable_update ()	Enables update rules of all parameters under the link hierarchy.
init_scope ()	Creates an initialization scope.
links ([<i>skipself</i>])	Returns a generator of all links under the hierarchy.
namedlinks ([<i>skipself</i>])	Returns a generator of all (path, link) pairs under the hierarchy.
namedparams ([<i>include_uninit</i>])	Returns a generator of all (path, param) pairs under the hierarchy.
params ([<i>include_uninit</i>])	Returns a generator of all parameters under the link hierarchy.

Continued on next page

Table 55 – continued from previous page

<code>register_persistent(name)</code>	Registers an attribute of a given name as a persistent value.
<code>repeat(n_repeat[, mode])</code>	Repeats this link multiple times to make a <code>Sequential</code> .
<code>serialize(serializer)</code>	Serializes the link object.
<code>to_cpu()</code>	Copies parameter variables and persistent values to CPU.
<code>to_gpu([device])</code>	Copies parameter variables and persistent values to GPU.
<code>to_intel64()</code>	Copies parameter variables and persistent values to CPU.
<code>zerograds()</code>	Initializes all gradient arrays by zero.

Attributes

<code>local_link_hooks</code>	Ordered dictionary of registered link hooks.
<code>update_enabled</code>	True if at least one parameter has an update rule enabled.
<code>within_init_scope</code>	True if the current code is inside of an initialization scope.
<code>xp</code>	Array module for this link.

chainer_chemistry.models.RelGCN

```
class chainer_chemistry.models.RelGCN(out_channels=64, num_edge_type=4,
                                     ch_list=None, n_atom_types=117, input_type='int',
                                     scale_adj=False)
```

Relational GCN (RelGCN)

See: Michael Schlichtkrull+, Modeling Relational Data with Graph Convolutional Networks. March 2017. [arXiv:1703.06103](https://arxiv.org/abs/1703.06103) <<https://arxiv.org/abs/1703.06103>>

Parameters

- **out_channels** (*int*) – dimension of output feature vector
- **num_edge_type** (*int*) – number of types of edge
- **ch_list** (*list*) – channels of each update layer
- **n_atom_types** (*int*) – number of types of atoms
- **input_type** (*str*) – type of input vector
- **scale_adj** (*bool*) – If `True`, then this network normalizes adjacency matrix

```
__init__(out_channels=64, num_edge_type=4, ch_list=None, n_atom_types=117, input_type='int',
         scale_adj=False)
```

Initialize self. See `help(type(self))` for accurate signature.

Methods

<code><u>__init__</u></code> ([out_channels, num_edge_type, ...])	Initialize self.
---	------------------

Continued on next page

Table 57 – continued from previous page

<code>add_hook(hook[, name])</code>	Registers a link hook.
<code>add_link(name, link)</code>	Registers a child link to this chain.
<code>add_param(name[, shape, dtype, initializer])</code>	Registers a parameter to the link.
<code>add_persistent(name, value)</code>	Registers a persistent value to the link.
<code>addgrads(link)</code>	Accumulates gradient values from given link.
<code>children()</code>	Returns a generator of all child links.
<code>cleargrads()</code>	Clears all gradient arrays.
<code>copy([model])</code>	Copies the link hierarchy to new one.
<code>copyparams(link[, copy_persistent])</code>	Copies all parameters from given link.
<code>count_params()</code>	Counts the total number of parameters.
<code>delete_hook(name)</code>	Unregisters the link hook.
<code>disable_update()</code>	Disables update rules of all parameters under the link hierarchy.
<code>enable_update()</code>	Enables update rules of all parameters under the link hierarchy.
<code>init_scope()</code>	Creates an initialization scope.
<code>links([skipself])</code>	Returns a generator of all links under the hierarchy.
<code>namedlinks([skipself])</code>	Returns a generator of all (path, link) pairs under the hierarchy.
<code>namedparams([include_uninit])</code>	Returns a generator of all (path, param) pairs under the hierarchy.
<code>params([include_uninit])</code>	Returns a generator of all parameters under the link hierarchy.
<code>register_persistent(name)</code>	Registers an attribute of a given name as a persistent value.
<code>repeat(n_repeat[, mode])</code>	Repeats this link multiple times to make a <code>Sequential</code> .
<code>serialize(serializer)</code>	Serializes the link object.
<code>to_cpu()</code>	Copies parameter variables and persistent values to CPU.
<code>to_gpu([device])</code>	Copies parameter variables and persistent values to GPU.
<code>to_intel64()</code>	Copies parameter variables and persistent values to CPU.
<code>zerograds()</code>	Initializes all gradient arrays by zero.

Attributes

<code>local_link_hooks</code>	Ordered dictionary of registered link hooks.
<code>update_enabled</code>	True if at least one parameter has an update rule enabled.
<code>within_init_scope</code>	True if the current code is inside of an initialization scope.
<code>xp</code>	Array module for this link.

chainer_chemistry.models.RSGCN

```
class chainer_chemistry.models.RSGCN(out_dim,          hidden_dim=32,          n_layers=4,
                                       n_atom_types=117, use_batch_norm=False, read-
                                       out=None, dropout_ratio=0.5)
```

Renormalized Spectral Graph Convolutional Network (RSGCN)

See: Thomas N. Kipf and Max Welling, Semi-Supervised Classification with Graph Convolutional Networks. September 2016. [arXiv:1609.02907](https://arxiv.org/abs/1609.02907)

The name of this model “Renormalized Spectral Graph Convolutional Network (RSGCN)” is named by us rather than the authors of the paper above. The authors call this model just “Graph Convolution Network (GCN)”, but we think that “GCN” is bit too general and may cause namespace issue. That is why we did not name this model as GCN.

Parameters

- **out_dim** (*int*) – dimension of output feature vector
- **hidden_dim** (*int*) – dimension of feature vector associated to each atom
- **n_atom_types** (*int*) – number of types of atoms
- **n_layers** (*int*) – number of layers
- **use_batch_norm** (*bool*) – If True, batch normalization is applied after graph convolution.
- **readout** (*Callable*) – readout function. If None, *GeneralReadout(mode='sum')* is used. To the best of our knowledge, the paper of RSGCN model does not give any suggestion on readout.
- **dropout_ratio** (*float*) – ratio used in dropout function. If 0 or negative value is set, dropout function is skipped.

```
__init__(out_dim, hidden_dim=32, n_layers=4, n_atom_types=117, use_batch_norm=False, read-
         out=None, dropout_ratio=0.5)
```

Initialize self. See `help(type(self))` for accurate signature.

Methods

<code>__init__(out_dim[, hidden_dim, n_layers, ...])</code>	Initialize self.
<code>add_hook(hook[, name])</code>	Registers a link hook.
<code>add_link(name, link)</code>	Registers a child link to this chain.
<code>add_param(name[, shape, dtype, initializer])</code>	Registers a parameter to the link.
<code>add_persistent(name, value)</code>	Registers a persistent value to the link.
<code>addgrads(link)</code>	Accumulates gradient values from given link.
<code>children()</code>	Returns a generator of all child links.
<code>cleargrads()</code>	Clears all gradient arrays.
<code>copy([model])</code>	Copies the link hierarchy to new one.
<code>copyparams(link[, copy_persistent])</code>	Copies all parameters from given link.
<code>count_params()</code>	Counts the total number of parameters.
<code>delete_hook(name)</code>	Unregisters the link hook.
<code>disable_update()</code>	Disables update rules of all parameters under the link hierarchy.
<code>enable_update()</code>	Enables update rules of all parameters under the link hierarchy.

Continued on next page

Table 59 – continued from previous page

<code>init_scope()</code>	Creates an initialization scope.
<code>links([skipself])</code>	Returns a generator of all links under the hierarchy.
<code>namedlinks([skipself])</code>	Returns a generator of all (path, link) pairs under the hierarchy.
<code>namedparams([include_uninit])</code>	Returns a generator of all (path, param) pairs under the hierarchy.
<code>params([include_uninit])</code>	Returns a generator of all parameters under the link hierarchy.
<code>register_persistent(name)</code>	Registers an attribute of a given name as a persistent value.
<code>repeat(n_repeat[, mode])</code>	Repeats this link multiple times to make a <code>Sequential</code> .
<code>serialize(serializer)</code>	Serializes the link object.
<code>to_cpu()</code>	Copies parameter variables and persistent values to CPU.
<code>to_gpu([device])</code>	Copies parameter variables and persistent values to GPU.
<code>to_intel64()</code>	Copies parameter variables and persistent values to CPU.
<code>zerograds()</code>	Initializes all gradient arrays by zero.

Attributes

<code>local_link_hooks</code>	Ordered dictionary of registered link hooks.
<code>update_enabled</code>	True if at least one parameter has an update rule enabled.
<code>within_init_scope</code>	True if the current code is inside of an initialization scope.
<code>xp</code>	Array module for this link.

Wrapper models

<code>chainer_chemistry.models.BaseForwardModel</code>	A base model which supports forward functionality.
<code>chainer_chemistry.models.Classifier</code>	A simple classifier model.
<code>chainer_chemistry.models.Regressor</code>	A simple regressor model.

chainer_chemistry.models.BaseForwardModel

class `chainer_chemistry.models.BaseForwardModel`

A base model which supports forward functionality.

It also supports *device* id management and pickle save/load functionality.

Parameters `device` (*int*) – GPU device id of this model to be used. -1 indicates to use in CPU.

`_dev_id`

Model's current device id

Type `int`

`__init__()`

Initialize self. See `help(type(self))` for accurate signature.

Methods

<code>__init__()</code>	Initialize self.
<code>add_hook(hook[, name])</code>	Registers a link hook.
<code>add_link(name, link)</code>	Registers a child link to this chain.
<code>add_param(name[, shape, dtype, initializer])</code>	Registers a parameter to the link.
<code>add_persistent(name, value)</code>	Registers a persistent value to the link.
<code>addgrads(link)</code>	Accumulates gradient values from given link.
<code>children()</code>	Returns a generator of all child links.
<code>cleargrads()</code>	Clears all gradient arrays.
<code>copy([model])</code>	Copies the link hierarchy to new one.
<code>copyparams(link[, copy_persistent])</code>	Copies all parameters from given link.
<code>count_params()</code>	Counts the total number of parameters.
<code>delete_hook(name)</code>	Unregisters the link hook.
<code>disable_update()</code>	Disables update rules of all parameters under the link hierarchy.
<code>enable_update()</code>	Enables update rules of all parameters under the link hierarchy.
<code>get_device()</code>	
<code>init_scope()</code>	Creates an initialization scope.
<code>initialize([device])</code>	Initialization of the model.
<code>links([skipself])</code>	Returns a generator of all links under the hierarchy.
<code>load_pickle(filepath[, device])</code>	Load the model from <i>filepath</i> of pickle file, and send to <i>device</i>
<code>namedlinks([skipself])</code>	Returns a generator of all (path, link) pairs under the hierarchy.
<code>namedparams([include_uninit])</code>	Returns a generator of all (path, param) pairs under the hierarchy.
<code>params([include_uninit])</code>	Returns a generator of all parameters under the link hierarchy.
<code>register_persistent(name)</code>	Registers an attribute of a given name as a persistent value.
<code>repeat(n_repeat[, mode])</code>	Repeats this link multiple times to make a <code>Sequential</code> .
<code>save_pickle(filepath[, protocol])</code>	Save the model to <i>filepath</i> as a pickle file
<code>serialize(serializer)</code>	Serializes the link object.
<code>to_cpu()</code>	Copies parameter variables and persistent values to CPU.
<code>to_gpu([device])</code>	Copies parameter variables and persistent values to GPU.
<code>to_intel64()</code>	Copies parameter variables and persistent values to CPU.
<code>update_device([device])</code>	
<code>zerogradsg()</code>	Initializes all gradient arrays by zero.

Attributes

<code>local_link_hooks</code>	Ordered dictionary of registered link hooks.
<code>update_enabled</code>	True if at least one parameter has an update rule enabled.
<code>within_init_scope</code>	True if the current code is inside of an initialization scope.
<code>xp</code>	Array module for this link.

chainer_chemistry.models.Classifier

```
class chainer_chemistry.models.Classifier(predictor, lossfun=<function softmax_cross_entropy>, accfun=None, metrics_fun=<function accuracy>, label_key=-1, device=-1)
```

A simple classifier model.

This is an example of chain that wraps another chain. It computes the loss and accuracy based on a given input/label pair.

Parameters

- **predictor** (*Link*) – Predictor network.
- **lossfun** (*function*) – Loss function.
- **accfun** (*function*) – DEPRECATED. Please use *metrics_fun* instead.
- **metrics_fun** (*function or dict or None*) – Function that computes metrics.
- **label_key** (*int or str*) – Key to specify label variable from arguments. When it is *int*, a variable in positional arguments is used. And when it is *str*, a variable in keyword arguments is used.
- **device** (*int*) – GPU device id of this Classifier to be used. -1 indicates to use in CPU.

predictor

Predictor network.

Type *Link*

lossfun

Loss function.

Type *function*

accfun

DEPRECATED. Please use *metrics_fun* instead.

Type *function*

y

Prediction for the last minibatch.

Type *Variable*

loss

Loss value for the last minibatch.

Type *Variable*

metrics

Metrics computed in last minibatch

Type *dict*

compute_metrics

If True, compute metrics on the forward computation. The default value is True.

Type `bool`

Note: The differences between original *Classifier* class in chainer and chainer chemistry are as follows. 1. *predict* and *predict_proba* methods are supported. 2. *device* can be managed internally by the *Classifier*. 3. *accfun* is deprecated, *metrics_fun* is used instead. 4. *metrics_fun* can be *dict* which specifies the metrics name as key

and function as value.

Note: This link uses `chainer.softmax_cross_entropy()` with default arguments as a loss function (specified by `lossfun`), if users do not explicitly change it. In particular, the loss function does not support double backpropagation. If you need second or higher order differentiation, you need to turn it on with `enable_double_backprop=True`:

```
>>> import chainer.functions as F
>>> import chainer.links as L
>>>
>>> def lossfun(x, t):
...     return F.softmax_cross_entropy(
...         x, t, enable_double_backprop=True)
>>>
>>> predictor = L.Linear(10)
>>> model = L.Classifier(predictor, lossfun=lossfun)
```

__init__(*predictor*, *lossfun*=<function softmax_cross_entropy>, *accfun*=None, *metrics_fun*=<function accuracy>, *label_key*=-1, *device*=-1)
Initialize self. See `help(type(self))` for accurate signature.

Methods

<code>__init__(predictor[, lossfun, accfun, ...])</code>	Initialize self.
<code>add_hook(hook[, name])</code>	Registers a link hook.
<code>add_link(name, link)</code>	Registers a child link to this chain.
<code>add_param(name[, shape, dtype, initializer])</code>	Registers a parameter to the link.
<code>add_persistent(name, value)</code>	Registers a persistent value to the link.
<code>addgrads(link)</code>	Accumulates gradient values from given link.
<code>children()</code>	Returns a generator of all child links.
<code>cleargrads()</code>	Clears all gradient arrays.
<code>copy([mode])</code>	Copies the link hierarchy to new one.
<code>copyparams(link[, copy_persistent])</code>	Copies all parameters from given link.
<code>count_params()</code>	Counts the total number of parameters.
<code>delete_hook(name)</code>	Unregisters the link hook.
<code>disable_update()</code>	Disables update rules of all parameters under the link hierarchy.
<code>enable_update()</code>	Enables update rules of all parameters under the link hierarchy.
<code>get_device()</code>	

Continued on next page

Table 64 – continued from previous page

<code>init_scope()</code>	Creates an initialization scope.
<code>initialize([device])</code>	Initialization of the model.
<code>links([skipself])</code>	Returns a generator of all links under the hierarchy.
<code>load_pickle(filepath[, device])</code>	Load the model from <i>filepath</i> of pickle file, and send to <i>device</i>
<code>namedlinks([skipself])</code>	Returns a generator of all (path, link) pairs under the hierarchy.
<code>namedparams([include_uninit])</code>	Returns a generator of all (path, param) pairs under the hierarchy.
<code>params([include_uninit])</code>	Returns a generator of all parameters under the link hierarchy.
<code>predict(data[, batchsize, converter, ...])</code>	Predict label of each category by taking .
<code>predict_proba(data[, batchsize, converter, ...])</code>	Calculate probability of each category.
<code>register_persistent(name)</code>	Registers an attribute of a given name as a persistent value.
<code>repeat(n_repeat[, mode])</code>	Repeats this link multiple times to make a <code>Sequential</code> .
<code>save_pickle(filepath[, protocol])</code>	Save the model to <i>filepath</i> as a pickle file
<code>serialize(serializer)</code>	Serializes the link object.
<code>to_cpu()</code>	Copies parameter variables and persistent values to CPU.
<code>to_gpu([device])</code>	Copies parameter variables and persistent values to GPU.
<code>to_intel64()</code>	Copies parameter variables and persistent values to CPU.
<code>update_device([device])</code>	
<code>zerograds()</code>	Initializes all gradient arrays by zero.

Attributes

<code>accfun</code>	
<code>accuracy</code>	
<code>compute_accuracy</code>	
<code>compute_metrics</code>	
<code>local_link_hooks</code>	Ordered dictionary of registered link hooks.
<code>update_enabled</code>	True if at least one parameter has an update rule enabled.
<code>within_init_scope</code>	True if the current code is inside of an initialization scope.
<code>xp</code>	Array module for this link.

chainer_chemistry.models.Regressor

```
class chainer_chemistry.models.Regressor (predictor,                lossfun=<function
                                         mean_squared_error>,        metrics_fun=None,
                                         label_key=-1, device=-1)
```

A simple regressor model.

This is an example of chain that wraps another chain. It computes the loss and metrics based on a given input/label pair.

Parameters

- **predictor** (*Link*) – Predictor network.
- **lossfun** (*function*) – Loss function.
- **metrics_fun** (*function or dict or None*) – Function that computes metrics.
- **label_key** (*int or str*) – Key to specify label variable from arguments. When it is *int*, a variable in positional arguments is used. And when it is *str*, a variable in keyword arguments is used.
- **device** (*int*) – GPU device id of this Regressor to be used. -1 indicates to use in CPU.

predictor

Predictor network.

Type *Link*

lossfun

Loss function.

Type *function*

y

Prediction for the last minibatch.

Type *Variable*

loss

Loss value for the last minibatch.

Type *Variable*

metrics

Metrics computed in last minibatch

Type *dict*

compute_metrics

If *True*, compute metrics on the forward computation. The default value is *True*.

Type *bool*

__init__ (*predictor, lossfun=<function mean_squared_error>, metrics_fun=None, label_key=-1, device=-1*)

Initialize self. See help(type(self)) for accurate signature.

Methods

<code>__init__(predictor[, lossfun, metrics_fun, ...])</code>	Initialize self.
<code>add_hook(hook[, name])</code>	Registers a link hook.
<code>add_link(name, link)</code>	Registers a child link to this chain.
<code>add_param(name[, shape, dtype, initializer])</code>	Registers a parameter to the link.
<code>add_persistent(name, value)</code>	Registers a persistent value to the link.
<code>addgrads(link)</code>	Accumulates gradient values from given link.
<code>children()</code>	Returns a generator of all child links.
<code>cleargrads()</code>	Clears all gradient arrays.
<code>copy([mode])</code>	Copies the link hierarchy to new one.
<code>copyparams(link[, copy_persistent])</code>	Copies all parameters from given link.
<code>count_params()</code>	Counts the total number of parameters.
<code>delete_hook(name)</code>	Unregisters the link hook.

Continued on next page

Table 66 – continued from previous page

<code>disable_update()</code>	Disables update rules of all parameters under the link hierarchy.
<code>enable_update()</code>	Enables update rules of all parameters under the link hierarchy.
<code>get_device()</code>	
<code>init_scope()</code>	Creates an initialization scope.
<code>initialize([device])</code>	Initialization of the model.
<code>links([skipself])</code>	Returns a generator of all links under the hierarchy.
<code>load_pickle(filepath[, device])</code>	Load the model from <i>filepath</i> of pickle file, and send to <i>device</i>
<code>namedlinks([skipself])</code>	Returns a generator of all (path, link) pairs under the hierarchy.
<code>namedparams([include_uninit])</code>	Returns a generator of all (path, param) pairs under the hierarchy.
<code>params([include_uninit])</code>	Returns a generator of all parameters under the link hierarchy.
<code>predict(data[, batchsize, converter, ...])</code>	Predict label of each category by taking .
<code>register_persistent(name)</code>	Registers an attribute of a given name as a persistent value.
<code>repeat(n_repeat[, mode])</code>	Repeats this link multiple times to make a <code>Sequential</code> .
<code>save_pickle(filepath[, protocol])</code>	Save the model to <i>filepath</i> as a pickle file
<code>serialize(serializer)</code>	Serializes the link object.
<code>to_cpu()</code>	Copies parameter variables and persistent values to CPU.
<code>to_gpu([device])</code>	Copies parameter variables and persistent values to GPU.
<code>to_intel64()</code>	Copies parameter variables and persistent values to CPU.
<code>update_device([device])</code>	
<code>zerograds()</code>	Initializes all gradient arrays by zero.

Attributes

<code>compute_metrics</code>	
<code>local_link_hooks</code>	Ordered dictionary of registered link hooks.
<code>update_enabled</code>	True if at least one parameter has an update rule enabled.
<code>within_init_scope</code>	True if the current code is inside of an initialization scope.
<code>xp</code>	Array module for this link.

1.5.7 Utilities**1.5.8 Training****Extensions**

```
chainer_chemistry.training.
extensions.batch_evaluator.
BatchEvaluator
```

```
chainer_chemistry.training.
extensions.roc_auc_evaluator.
ROCAUCEvaluator
```

Evaluator which calculates ROC AUC score

```
chainer_chemistry.training.
extensions.prc_auc_evaluator.
PRCAUCEvaluator
```

Evaluator which calculates PRC AUC score

chainer_chemistry.training.extensions.batch_evaluator.BatchEvaluator

```
class chainer_chemistry.training.extensions.batch_evaluator.BatchEvaluator (iterator,
                                                                    tar-
                                                                    get,
                                                                    con-
                                                                    verter=<function
                                                                    concat_examples>,
                                                                    de-
                                                                    vice=None,
                                                                    eval_hook=None,
                                                                    eval_func=None,
                                                                    met-
                                                                    rics_fun=None,
                                                                    name=None,
                                                                    log-
                                                                    ger=None)
```

```
__init__(iterator, target, converter=<function concat_examples>, device=None, eval_hook=None,
          eval_func=None, metrics_fun=None, name=None, logger=None)
    Initialize self. See help(type(self)) for accurate signature.
```

Methods

<code>__init__(iterator, target[, converter, ...])</code>	Initialize self.
<code>evaluate()</code>	Evaluates the model and returns a result dictionary.
<code>finalize()</code>	Finalizes the evaluator object.
<code>get_all_iterators()</code>	Returns a dictionary of all iterators.
<code>get_all_targets()</code>	Returns a dictionary of all target links.
<code>get_iterator(name)</code>	Returns the iterator of the given name.
<code>get_target(name)</code>	Returns the target link of the given name.
<code>initialize(trainer)</code>	Initializes up the trainer state.
<code>serialize(serializer)</code>	Serializes the extension state.

Attributes

<code>default_name</code>
<code>name</code>

Continued on next page

Table 70 – continued from previous page

priority
trigger

chainer_chemistry.training.extensions.roc_auc_evaluator.ROCAUCEvaluator

```
class chainer_chemistry.training.extensions.roc_auc_evaluator.ROCAUCEvaluator (iterator,
                                                                              tar-
                                                                              get,
                                                                              con-
                                                                              verter=<function
                                                                              con-
                                                                              cat_examples>,
                                                                              de-
                                                                              vice=None,
                                                                              eval_hook=None,
                                                                              eval_func=None,
                                                                              name=None,
                                                                              pos_labels=1,
                                                                              ig-
                                                                              nore_labels=None,
                                                                              raise_value_error:
                                                                              log-
                                                                              ger=None)
```

Evaluator which calculates ROC AUC score

Note that this Evaluator is only applicable to binary classification task.

Parameters

- **iterator** – Dataset iterator for the dataset to calculate ROC AUC score. It can also be a dictionary of iterators. If this is just an iterator, the iterator is registered by the name 'main'.
- **target** – Link object or a dictionary of links to evaluate. If this is just a link object, the link is registered by the name 'main'.
- **converter** – Converter function to build input arrays and true label. `concat_examples()` is used by default. It is expected to return input arrays of the form $[x_0, \dots, x_n, t]$, where $x_0, \dots, x_n$ are the inputs to the evaluation function and t is the true label.
- **device** – Device to which the training data is sent. Negative value indicates the host memory (CPU).
- **eval_hook** – Function to prepare for each evaluation process. It is called at the beginning of the evaluation. The evaluator extension object is passed at each call.
- **eval_func** – Evaluation function called at each iteration. The target link to evaluate as a callable is used by default.
- **name** (*str*) – name of this extension. When *name* is None, *default_name*='validation' which is defined in super class *Evaluator* is used as extension name. This name affects to the reported key name.
- **pos_labels** (*int* or *list*) – labels of the positive class, other classes are considered as negative.

- **ignore_labels** (*int* or *list* or *None*) – labels to be ignored. *None* is used to not ignore all labels.
- **raise_value_error** (*bool*) – If *False*, *ValueError* caused by *roc_auc_score* calculation is suppressed and ignored with a warning message.
- **logger** –

converter

Converter function.

device

Device to which the training data is sent.

eval_hook

Function to prepare for each evaluation process.

eval_func

Evaluation function called at each iteration.

pos_labels

labels of the positive class

Type *list*

ignore_labels

labels to be ignored.

Type *list*

__init__ (*iterator*, *target*, *converter*=<function *concat_examples*>, *device*=*None*, *eval_hook*=*None*, *eval_func*=*None*, *name*=*None*, *pos_labels*=*1*, *ignore_labels*=*None*, *raise_value_error*=*True*, *logger*=*None*)
Initialize self. See help(type(self)) for accurate signature.

Methods

<code>__init__(iterator, target[, converter, ...])</code>	Initialize self.
<code>evaluate()</code>	Evaluates the model and returns a result dictionary.
<code>finalize()</code>	Finalizes the evaluator object.
<code>get_all_iterators()</code>	Returns a dictionary of all iterators.
<code>get_all_targets()</code>	Returns a dictionary of all target links.
<code>get_iterator(name)</code>	Returns the iterator of the given name.
<code>get_target(name)</code>	Returns the target link of the given name.
<code>initialize(trainer)</code>	Initializes up the trainer state.
<code>roc_auc_score(y_total, t_total)</code>	
<code>serialize(serializer)</code>	Serializes the extension state.

Attributes

<code>default_name</code>
<code>name</code>
<code>priority</code>
<code>trigger</code>

chainer_chemistry.training.extensions.prc_auc_evaluator.PRCAUCEvaluator

```
class chainer_chemistry.training.extensions.prc_auc_evaluator.PRCAUCEvaluator (iterator,
                                                                    tar-
                                                                    get,
                                                                    con-
                                                                    verter=<function
                                                                    con-
                                                                    cat_examples>,
                                                                    de-
                                                                    vice=None,
                                                                    eval_hook=None,
                                                                    eval_func=None,
                                                                    name=None,
                                                                    pos_labels=1,
                                                                    ig-
                                                                    nore_labels=None,
                                                                    raise_value_error=
                                                                    log-
                                                                    ger=None)
```

Evaluator which calculates PRC AUC score

Note that this Evaluator is only applicable to binary classification task.

Parameters

- **iterator** – Dataset iterator for the dataset to calculate PRC AUC score. It can also be a dictionary of iterators. If this is just an iterator, the iterator is registered by the name 'main'.
- **target** – Link object or a dictionary of links to evaluate. If this is just a link object, the link is registered by the name 'main'.
- **converter** – Converter function to build input arrays and true label. `concat_examples()` is used by default. It is expected to return input arrays of the form $[x_0, \dots, x_n, t]$, where $x_0, \dots, x_n$ are the inputs to the evaluation function and t is the true label.
- **device** – Device to which the training data is sent. Negative value indicates the host memory (CPU).
- **eval_hook** – Function to prepare for each evaluation process. It is called at the beginning of the evaluation. The evaluator extension object is passed at each call.
- **eval_func** – Evaluation function called at each iteration. The target link to evaluate as a callable is used by default.
- **name** (*str*) – name of this extension. When *name* is *None*, *default_name*='validation' which is defined in super class *Evaluator* is used as extension name. This name affects to the reported key name.
- **pos_labels** (*int* or *list*) – labels of the positive class, other classes are considered as negative.
- **ignore_labels** (*int* or *list* or *None*) – labels to be ignored. *None* is used to not ignore all labels.
- **raise_value_error** (*bool*) – If *False*, *ValueError* caused by *roc_auc_score* calculation is suppressed and ignored with a warning message.

- **logger** –

converter

Converter function.

device

Device to which the training data is sent.

eval_hook

Function to prepare for each evaluation process.

eval_func

Evaluation function called at each iteration.

pos_labels

labels of the positive class

Type *list*

ignore_labels

labels to be ignored.

Type *list*

__init__(*iterator*, *target*, *converter*=<function *concat_examples*>, *device*=None, *eval_hook*=None, *eval_func*=None, *name*=None, *pos_labels*=1, *ignore_labels*=None, *raise_value_error*=True, *logger*=None)
Initialize self. See help(type(self)) for accurate signature.

Methods

<code>__init__(iterator, target[, converter, ...])</code>	Initialize self.
<code>evaluate()</code>	Evaluates the model and returns a result dictionary.
<code>finalize()</code>	Finalizes the evaluator object.
<code>get_all_iterators()</code>	Returns a dictionary of all iterators.
<code>get_all_targets()</code>	Returns a dictionary of all target links.
<code>get_iterator(name)</code>	Returns the iterator of the given name.
<code>get_target(name)</code>	Returns the target link of the given name.
<code>initialize(trainer)</code>	Initializes up the trainer state.
<code>prc_auc_score(y_total, t_total)</code>	
<code>serialize(serializer)</code>	Serializes the extension state.

Attributes

<code>default_name</code>
<code>name</code>
<code>priority</code>
<code>trigger</code>

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