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# e3fp Documentation

*Release 1.2.1*

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**Feb 27, 2020**



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# CHAPTER ONE

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## CONTENTS

### 1.1 Overview of E3FP

#### 1.1.1 Introduction

The Extended 3-Dimensional FingerPrint (E3FP)<sup>1</sup> is a 3D molecular fingerprinting method inspired by Extended Connectivity FingerPrints (ECFP)<sup>2</sup>, integrating tightly with the [RDKit](#). It is developed by the [Keiser Lab at UCSF](#) and maintained primarily by [Seth Axen](#).

For a thorough description of E3FP, please consult the original paper<sup>1</sup> and [paper repository](#) or [Usage and Examples](#). Documentation is hosted by [ReadTheDocs](#).

#### 1.1.2 Contributing

Development occurs on [GitHub](#). Contributions, feature requests, and bug reports are greatly appreciated. Please consult the [issue tracker](#).

#### 1.1.3 License

E3FP is released under the [GNU Lesser General Public License version 3.0 \(LGPLv3\)](#).

Briefly, this means E3FP can be used in any manner without modification, with proper attribution. However, if the source code is modified for an application, this modified source must also be released under GPLv3 so that the community may benefit.

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<sup>1</sup> Axen SD, Huang XP, Caceres EL, Gendelev L, Roth BL, Keiser MJ. A Simple Representation Of Three-Dimensional Molecular Structure. *J. Med. Chem.* **60** (17): 7393–7409 (2017). doi: [10.1021/acs.jmedchem.7b00696](https://doi.org/10.1021/acs.jmedchem.7b00696) bioRxiv 136705

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<sup>2</sup> Rogers D & Hahn M. Extended-connectivity fingerprints. *J. Chem. Inf. Model.* **50**: 742-54 (2010). doi: [10.1021/ci100050t](https://doi.org/10.1021/ci100050t)

## 1.1.4 Citing E3FP

To cite E3FP, please reference the original paper<sup>1</sup>.

## References

# 1.2 Setup and Installation

## 1.2.1 Dependencies

E3FP is compatible with Python 2.7.x and 3.x. It additionally has the following dependencies:

### Required

- NumPy
- SciPy
- RDKit
- mmh3
- python\_utilities

### Optional

The following packages are required for the specified features:

- parallelization:
  - mpi4py
  - futures
- molecular standardisation:
  - standardiser
- protonation states:
  - cxcalc
- storing conformer energies:
  - h5py

## 1.2.2 Installation

The following installation approaches are listed in order of recommendation. All but the first of these approaches requires a prior installation of RDKit.

## Option 1: Install with Conda

E3FP is on the [Anaconda distribution](#). Conda is a cross-platform package manager. This approach is highly recommended as it installs *all* required packages.

```
$ conda create -c keiserlab -c rdkit -c sdaxen --name e3fp_env e3fp
```

**Warning:** Due to a [potential bug](#) in some versions of conda, rdkit may not import on some systems. If this is the case, simply upgrade to the latest version of conda before creating the above environment:

```
conda update conda
```

To get the latest version of E3FP, follow [Option 3: Clone the Repository](#).

## Option 2: Install with Pip

1. Install with

```
$ pip install e3fp
```

2. To install the optional Python dependencies, run

```
$ pip install mpi4py futures standardiser h5py
```

## Option 3: Clone the Repository

0. Install any of the optional dependencies above.

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**Note:**

The easiest way to install the dependencies is with conda using Option 1 above. If e3fp is also installed with conda, you will need to uninstall it before continuing below with

```
conda uninstall e3fp
```

---

1. Download this repository to your machine.

- Clone this repository to your machine with

```
$ git clone https://github.com/keiserlab/e3fp.git
```

- OR download an archive by navigating to the repository and clicking “Download ZIP”. Extract the archive.

2. Install with

```
$ cd e3fp
$ python setup.py build_ext --inplace
$ python setup.py install
```

### 1.2.3 Testing

After installation, it is recommended to run all tests with nose,

```
$ pip install nose
$ nosetests e3fp
```

## 1.3 Usage and Examples

To facilitate flexible use of the E3FP package, we provide multiple interfaces for performing the same tasks. We have organized these below in the order in which we expect them to be most of use to the average user.

### 1.3.1 Configuration

E3FP configurational parameters are stored in the widely used [INI](#) file format. These may be passed to *Command Line Interface* programs or parsed to Python dicts for *Pipeline Methods* or other lower-level functions.

#### Loading Default Parameters

The below example shows all default parameters, accessed via the `e3fp.config` module.

Listing 1: defaults.cfg

```
[preprocessing]
standardise = False
protonate = False

[conformer_generation]
num_conf = -1
first = -1
pool_multiplier = 1
rmsd_cutoff = 0.5
max_energy_diff = None
forcefield = uff
out_dir = conformers
compress = 2
seed = -1

; Optimized parameters used in
; Axen et al. 2017
[fingerprinting]
bits = 1024
level = 5
first = 3
radius_multiplier = 1.718
stereo = True
counts = False
include_disconnected = True
rdkit_invariants = False
remove_duplicate_substructs = True
exclude_floating = True
```

`configparser` is used internally to parse and store these config parameters.

```
>>> from e3fp.config.params import default_params
>>> default_params
<ConfigParser.SafeConfigParser instance at 0x...>
>>> print(default_params.sections())
['preprocessing', 'conformer_generation', 'fingerprinting']
>>> default_params.items('fingerprinting')
[('bits', '1024'), ('level', '5'), ('first', '3'), ('radius_multiplier', '1.718'), (
    'stereo', 'True'), ('counts', 'False'), ('include_disconnected', 'True'), ('rdkit_
    invariants', 'False'), ('merge_duplicate_substructs', 'True'), ('exclude_floating',
    'True')]
```

## Parsing User-Provided Parameters

A user may provide a custom config file.

Listing 2: new\_params.cfg

```
[conformer_generation]
first = 10

[fingerprinting]
bits = 4096
first = 10
```

```
>>> from e3fp.config.params import read_params
>>> config = read_params("source/examples/data/new_params.cfg")
>>> config.items('fingerprinting')
[('bits', '4096'), ('first', '10')]
```

When passing these parameters to any downstream methods, default options will be used except where these options are specified.

## Converting Parameters to Argument Dicts

To pass the parameters to Python methods for fingerprinting and conformer generation, we need to convert them to Python dicts.

```
>>> from e3fp.pipeline import params_to_dicts
>>> confgen_params, fprint_params = params_to_dicts(config)
>>> fprint_params
{'bits': 4096, 'first': 10}
```

## 1.3.2 Command Line Interface

Command line interfaces (CLI) are provided for the two most common tasks: conformer generation and fingerprinting. At the moment, using the CLI requires *downloading the E3FP source*.

In the below examples, we assume the E3FP repository is located at \$E3FP\_REPO.

## Conformer Generation CLI

To see all available options, run

```
$ python $E3FP_REPO/e3fp/conformer/generate.py --help
usage: Generate conformers from mol2 or SMILES [-h] [-m MOL2 [MOL2 ...]]
                                                [-s SMILES [SMILES ...]]
                                                [--standardise STANDARDISE]
                                                [-n NUM_CONF] [--first FIRST]
                                                [--pool_multiplier POOL_MULTIPLIER]
                                                [-r RMSD_CUTOFF]
                                                [-e MAX_ENERGY_DIFF]
                                                [-f {uff,mmff94,mmff94s}]
                                                [--seed SEED] [-o OUT_DIR]
                                                [-C {0,1,2,None}] [-O]
                                                [--values_file VALUES_FILE]
                                                [--prioritize]
                                                [--params PARAMS] [-l LOG]
                                                [-p NUM_PROC]
                                                [--parallel_mode {mpi,processes,
→threads,serial}]
                                                [-v]

optional arguments:
-h, --help            show this help message and exit
-m MOL2 [MOL2 ...], --mol2 MOL2 [MOL2 ...]
                      Path to mol2 file(s), each with one molecule.
                      (default: None)
-s SMILES [SMILES ...], --smiles SMILES [SMILES ...]
                      Path to file(s) with SMILES and name. (space-
                      separated) (default: None)
--standardise STANDARDISE
                      Clean molecules before generating conformers by
                      standardisation. (default: False)
-n NUM_CONF, --num_conf NUM_CONF
                      Set single number of conformers to use. -1 results in
                      auto choosing. (default: -1)
--first FIRST
                      Set maximum number of first conformers to accept.
                      Conformer generation is unaffected, except it may
                      terminate early when this number of conformers is
                      reached. (default: -1)
--pool_multiplier POOL_MULTIPLIER
                      Factor to multiply `num_conf` by to generate
                      conformers. Results are then pruned to `num_conf`.
                      (default: 1)
-r RMSD_CUTOFF, --rmsd_cutoff RMSD_CUTOFF
                      Choose RMSD cutoff between conformers (default: 0.5)
-e MAX_ENERGY_DIFF, --max_energy_diff MAX_ENERGY_DIFF
                      Maximum energy difference between lowest energy
                      conformer and any accepted conformer. (default: None)
-f {uff,mmff94,mmff94s}, --forcefield {uff,mmff94,mmff94s}
                      Choose forcefield for minimization. (default: uff)
--seed SEED
                      Random seed for conformer generation. (default: -1)
-o OUT_DIR, --out_dir OUT_DIR
                      Directory to save conformers. (default: conformers)
-C {0,1,2,None}, --compress {0,1,2,None}
                      Compression to use for SDF files. None and 0 default
                      to uncompressed ".sdf". 1 and 2 result in gzipped and
```

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-O, --overwrite	bzipped SDF files, respectively. (default: 2)
--values_file VALUES_FILE	Overwrite existing conformer files. (default: False)
--prioritize	Save RMSDs and energies to specified hdf5 file. (default: None)
--params PARAMS	Prioritize likely fast molecules first. (default: False)
-l LOG, --log LOG	INI formatted file with parameters. If provided, all parameters controlling conformer generation are ignored. (default: None)
-p NUM_PROC, --num_proc NUM_PROC	Generate logfile. (default: None)
--parallel_mode {mpi,processes,threads,serial}	Set number of processors to use. (default: None)
-v, --verbose	Set number of processors to use. (default: None)
	Run with extra verbosity. (default: False)

We will generate conformers for the molecule whose SMILES string is defined in `caffeine.smi`.

Listing 3: `caffeine.smi`

```
CN1C=NC2=C1C(=O)N(C(=O)N2C)C caffeine
```

The below example generates at most 3 conformers for this molecule.

```
$ python $E3FP_REPO/e3fp/conformer/generate.py -s caffeine.smi --num_conf 3 -o ./
2017-07-17 00:11:05,743|WARNING|Only 1 processes available. 'mpi' mode not available.
2017-07-17 00:11:05,748|INFO|num_proc is not specified. 'processes' mode will use all ↴
→8 processes
2017-07-17 00:11:05,748|INFO|Parallelizer initialized with mode 'processes' and 8 ↴
→processors.
2017-07-17 00:11:05,748|INFO|Input type: Detected SMILES file(s)
2017-07-17 00:11:05,748|INFO|Input file number: 1
2017-07-17 00:11:05,748|INFO|Parallel Type: processes
2017-07-17 00:11:05,748|INFO|Out Directory: ../
2017-07-17 00:11:05,749|INFO|Overwrite Existing Files: False
2017-07-17 00:11:05,749|INFO|Target Conformer Number: 3
2017-07-17 00:11:05,749|INFO|First Conformers Number: all
2017-07-17 00:11:05,749|INFO|Pool Multiplier: 1
2017-07-17 00:11:05,749|INFO|RMSD Cutoff: 0.5
2017-07-17 00:11:05,749|INFO|Maximum Energy Difference: None
2017-07-17 00:11:05,749|INFO|Forcefield: UFF
2017-07-17 00:11:05,749|INFO|Starting.
2017-07-17 00:11:05,779|INFO|Generating conformers for caffeine.
2017-07-17 00:11:05,823|INFO|Generated 1 conformers for caffeine.
2017-07-17 00:11:05,829|INFO|Saved conformers for caffeine to ./caffeine.sdf.bz2.
```

The result is a multi-conformer SDF file called `caffeine.sdf.bz2` in the current directory.

## Fingerprinting CLI

To see all available options, run

```
$ python $E3FP_REPO/e3fp/fingerprint/generate.py --help
usage: Generate E3FP fingerprints from SDF files. [-h] [-b BITS]
                                                [--first FIRST] [-m LEVEL]
                                                [-r RADIUS_MULTIPLIER]
                                                [--stereo STEREO]
                                                [--counts COUNTS]
                                                [--params PARAMS]
                                                [-o OUT_DIR_BASE]
                                                [--out_ext {.fp.pkl,.fp.gz,.fp.bz2}]
                                                [-d DB_FILE] [--all_iters]
                                                [-O] [-l LOG] [-p NUM_PROC]
                                                [--parallel_mode {mpi,processes,
→threads,serial}]
                                                [-v]
sdf_files [sdf_files ...]

positional arguments:
  sdf_files            Path to SDF file(s), each with one molecule and
                      multiple conformers.

optional arguments:
  -h, --help           show this help message and exit
  -b BITS, --bits BITS Set number of bits for final folded fingerprint. If -1
                      or None, unfolded (2^32-bit) fingerprints are
                      generated. (default: 4294967296)
  --first FIRST        Set maximum number of first conformers for which to
                      generate fingerprints. (default: 3)
  -m LEVEL, --level LEVEL, --max_iterations LEVEL
                      Maximum number of iterations for fingerprint
                      generation. If -1, fingerprinting is run until
                      termination, and `all_iters` is set to False.
                      (default: 5)
  -r RADIUS_MULTIPLIER, --radius_multiplier RADIUS_MULTIPLIER, --shell_radius RADIUS_
→MULTIPLIER          Distance to increment shell radius at around each
                      atom, starting at 0.0. (default: 1.718)
  --stereo STEREO     Differentiate by stereochemistry. (default: True)
  --counts COUNTS     Store counts-based E3FC instead of default bit-based.
                      (default: False)
  --params PARAMS     INI formatted file with parameters. If provided, all
                      parameters controlling conformer generation are
                      ignored. (default: None)
  -o OUT_DIR_BASE, --out_dir_base OUT_DIR_BASE
                      Basename for output directory to save fingerprints.
                      Iteration number is appended to basename. (default:
                      None)
  --out_ext {.fp.pkl,.fp.gz,.fp.bz2}
                      Extension for fingerprint pickles. (default: .fp.bz2)
  -d DB_FILE, --db_file DB_FILE
                      Output file containing FingerprintDatabase object
                      (default: fingerprints.fpz)
  --all_iters          Save fingerprints from all iterations to file(s).
                      (default: False)
  -O, --overwrite      Overwrite existing file(s). (default: False)
```

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```
-l LOG, --log LOG      Log filename. (default: None)
-p NUM_PROC, --num_proc NUM_PROC
                      Set number of processors to use. (default: None)
--parallel_mode {mpi,processes,threads,serial}
                      Set parallelization mode to use. (default: None)
-v, --verbose          Run with extra verbosity. (default: False)
```

To continue the above example, we will fingerprint our caffeine conformers.

```
$ python $E3FP_REPO/e3fp/fingerprint/generate.py caffeine.sdf.bz2 --bits 1024
2017-07-17 00:12:33,797|WARNING|Only 1 processes available. 'mpi' mode not available.
2017-07-17 00:12:33,801|INFO|num_proc is not specified. 'processes' mode will use all ↵
→8 processes
2017-07-17 00:12:33,801|INFO|Parallelizer initialized with mode 'processes' and 8 ↵
→processors.
2017-07-17 00:12:33,801|INFO|Initializing E3FP generation.
2017-07-17 00:12:33,801|INFO|Getting SDF files
2017-07-17 00:12:33,801|INFO|SDF File Number: 1
2017-07-17 00:12:33,802|INFO|Database File: fingerprints.fpz
2017-07-17 00:12:33,802|INFO|Max First Conformers: 3
2017-07-17 00:12:33,802|INFO|Bits: 1024
2017-07-17 00:12:33,802|INFO|Level/Max Iterations: 5
2017-07-17 00:12:33,802|INFO|Shell Radius Multiplier: 1.718
2017-07-17 00:12:33,802|INFO|Stereo Mode: True
2017-07-17 00:12:33,802|INFO|Connected-only mode: on
2017-07-17 00:12:33,802|INFO|Invariant type: Daylight
2017-07-17 00:12:33,802|INFO|Parallel Mode: processes
2017-07-17 00:12:33,802|INFO|Starting
2017-07-17 00:12:33,829|INFO|Generating fingerprints for caffeine.
2017-07-17 00:12:33,935|INFO|Generated 1 fingerprints for caffeine.
2017-07-17 00:12:34,011|INFO|Saved FingerprintDatabase with fingerprints to ↵
→fingerprints.fpz
```

The result is a file `fingerprints.fpz` containing a *FingerprintDatabase*. To use such a database, consult *Fingerprint Storage*.

### 1.3.3 Pipeline Methods

E3FP can be easily plugged into an existing pipeline using the methods in the `e3fp.pipeline` module. Each of these methods wraps functionality in other modules for generating various outputs from inputs and specified options.

---

**Note:** As fingerprinting many molecules is embarrassingly parallel, we highly recommend employing a parallelization strategy. We use our own `python_utilities` package.

First we must choose configuration options. See *Configuration* for detailed instructions. Here we will use defaults for all but a few options.

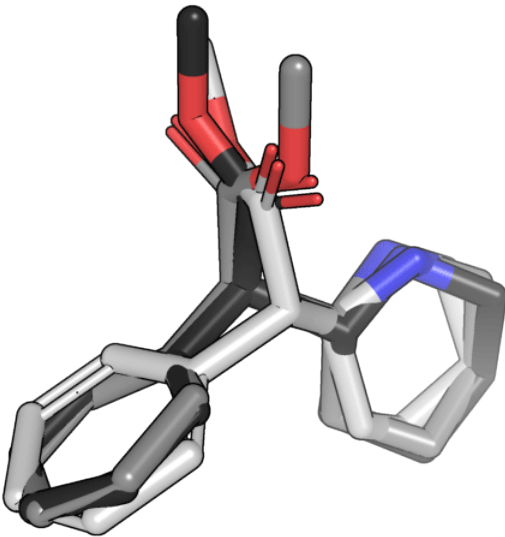
```
>>> fprint_params = {'bits': 4096, 'radius_multiplier': 1.5, 'rdkit_invariants': True}
>>> confgen_params = {'max_energy_diff': 20.0, 'first': 3}
>>> smiles = "COC(=O)C(C1CCCCN1)C2=CC=CC=C2"
```

## Generating Conformers from SMILES

The following code snippet generates a multi-conformer molecule:

```
>>> from e3fp.pipeline import confs_from_smiles
>>> mol = confs_from_smiles(smiles, "ritalin", confgen_params=confgen_params)
>>> mol.GetNumConformers()
3
```

This produces the following conformers:



## Generating Fingerprints from Conformers

```
>>> from e3fp.pipeline import fprints_from_mol
>>> fprints = fprints_from_mol(mol, fprint_params=fprint_params)
>>> len(fprints)
3
>>> fprints[0]
Fingerprint(indices=array([188, 224, ..., 3775, 4053]), level=5, bits=4096, ↴
    ↪name=ritalin_0)
>>> fprints[1]
Fingerprint(indices=array([125, 188, ..., 3693, 4053]), level=5, bits=4096, ↴
    ↪name=ritalin_1)
>>> fprints[2]
Fingerprint(indices=array([188, 206, ..., 3743, 4053]), level=5, bits=4096, ↴
    ↪name=ritalin_2)
```

## Generating Fingerprints from SMILES

```
>>> from e3fp.pipeline import fprints_from_smiles
>>> fprints = fprints_from_smiles(smiles, "ritalin", confgen_params=confgen_params,
    ↪fprint_params=fprint_params)
>>> fprints[0]
Fingerprint(indices=array([188, 224, ..., 3775, 4053]), level=5, bits=4096,
    ↪name=ritalin_0)
```

## Parallel Fingerprinting

The following script demonstrates use of `python_utilities` for fingerprinting all SDF files in a directory in parallel. This essentially is the same as the *Fingerprinting CLI*, albeit with a less convenient interface.

```
>>> from glob import glob
>>> from python_utilities.parallel import Parallelizer
>>> from e3fp.conformer.util import smiles_to_dict
>>> smiles_dict = smiles_to_dict(smiles_file)
>>> print(smiles_dict)
{'CHEMBL1643866': 'CCCC[C@H]1CN(O)C(=O)C(=O)[C@@H](NCC(=O)C(C)C)C(C)C', ...}
>>> len(smiles_dict)
10
>>> smiles_iter = ((smiles, name) for name, smiles in smiles_dict.items())
>>> kwargs = {"confgen_params": confgen_params, "fprint_params": fprint_params}
>>> parallelizer = Parallelizer(parallel_mode="processes")
>>> fprints_list = parallelizer.run(fprints_from_smiles, smiles_iter, kwargs=kwargs)
>>> len(fprints_list)
10
```

For all pipeline methods, please see the `e3fp.pipeline` module API.

### 1.3.4 Using Fingerprints

While molecular fingerprints are widely used, few packages provide simple interfaces for working with them and interfacing with machine learning packages. E3FP provides a number of general utility classes and methods for doing precisely this.

#### Fingerprints

The simplest interface for molecular fingerprints are through three classes in `e3fp.fingerprint.fprint`:

***Fingerprint*** a fingerprint with “on” bits

***CountFingerprint*** a fingerprint with counts for each “on” bit

***FloatFingerprint*** a fingerprint with float values for each “on” bit, generated for example by averaging conformer fingerprints.

In addition to storing “on” indices and, for the latter two, corresponding values, they store fingerprint properties, such as name, level, and any arbitrary property. They also provide simple interfaces for fingerprint comparison, some basic processing, and comparison.

---

**Note:** Many of these operations are more efficient when operating on a *FingerprintDatabase*. See *Fingerprint Storage* for more information.

---

In the below examples, we will focus on *Fingerprint* and *CountFingerprint*. First, we execute the necessary imports.

```
>>> from e3fp.fingerprint.fprint import Fingerprint, CountFingerprint
>>> import numpy as np
```

**See also:**

*Fingerprint Storage*, *Fingerprint Comparison*

### Creation and Conversion

Here we create a bit-fingerprint with random “on” indices.

```
>>> bits = 2**32
>>> indices = np.sort(np.random.randint(0, bits, 30))
>>> indices
array([ 243580376,  305097549, ..., 3975407269, 4138900056])
>>> fp1 = Fingerprint(indices, bits=bits, level=0)
>>> fp1
Fingerprint(indices=array([243580376, ..., 4138900056]), level=0, bits=4294967296,
             name=None)
```

This fingerprint is extremely sparse

```
>>> fp1.bit_count
30
>>> fp1.density
6.984919309616089e-09
```

We can therefore “fold” the fingerprint through a series of bitwise “OR” operations on halves of the sparse vector until it is of a specified length, with minimal collision of bits.

```
>>> fp_folded = fp1.fold(1024)
>>> fp_folded
Fingerprint(indices=array([9, 70, ..., 845, 849]), level=0, bits=1024, name=None)
>>> fp_folded.bit_count
29
>>> fp_folded.density
0.0283203125
```

A *CountFingerprint* may be created by also providing a dictionary matching indices with nonzero counts to the counts.

```
>>> indices2 = np.sort(np.random.randint(0, bits, 60))
>>> counts = dict(zip(indices2, np.random.randint(1, 10, indices2.size)))
>>> counts
{80701568: 8, 580757632: 7, ..., 800291326: 5, 4057322111: 7}
>>> cfp1 = CountFingerprint(counts=counts, bits=bits, level=0)
>>> cfp1
CountFingerprint(counts={80701568: 8, 580757632: 7, ..., 3342157822: 2, 4057322111: 7},
                  level=0, bits=4294967296, name=None)
```

Unlike folding a bit fingerprint, by default, folding a count fingerprint performs a “SUM” operation on colliding counts.

```
>>> cfp1.bit_count
60
>>> cfp_folded = cfp1.fold(1024)
>>> cfp_folded
CountFingerprint(counts={128: 15, 257: 4, ..., 1022: 2, 639: 7}, level=0, bits=1024, ↵
˓→name=None)
>>> cfp_folded.bit_count
57
```

It is trivial to interconvert the fingerprints.

```
>>> cfp_folded2 = CountFingerprint.from_fingerprint(fp_folded)
>>> cfp_folded2
CountFingerprint(counts={9: 1, 87: 1, ..., 629: 1, 763: 1}, level=0, bits=1024, ↵
˓→name=None)
>>> cfp_folded2.indices[:5]
array([ 9, 70, 72, 87, 174])
>>> fp_folded.indices[:5]
array([ 9, 70, 72, 87, 174])
```

RDKit Morgan fingerprints (analogous to ECFP) may easily be converted to a *Fingerprint*.

```
>>> from rdkit import Chem
>>> from rdkit.Chem import AllChem
>>> mol = Chem.MolFromSmiles('Cc1ccccc1')
>>> mfp = AllChem.GetMorganFingerprintAsBitVect(mol, 2)
>>> mfp
<rdkit.DataStructs.cDataStructs.ExplicitBitVect object at 0x...>
>>> Fingerprint.from_rdkit(mfp)
Fingerprint(indices=array([389, 1055, ..., 1873, 1920]), level=-1, bits=2048, ↵
˓→name=None)
```

Likewise, *Fingerprint* can be easily converted to a NumPy ndarray or SciPy sparse matrix.

```
>>> fp_folded.to_vector()
<1x1024 sparse matrix of type '<type 'numpy.bool_''>'>
...with 29 stored elements in Compressed Sparse Row format>
>>> fp_folded.to_vector(sparse=False)
array([False, False, False, ..., False, False, False], dtype=bool)
>>> np.where(fp_folded.to_vector(sparse=False))[0]
array([ 9, 70, 72, 87, ...])
>>> cfp_folded.to_vector(sparse=False)
array([0, 0, 0, ..., 0, 2, 0], dtype=uint16)
>>> cfp_folded.to_vector(sparse=False).sum()
252
```

## Algebra

Basic algebraic functions may be performed on fingerprints. If either fingerprint is a bit fingerprint, all algebraic functions are bit-wise. The following bit-wise operations are supported:

### Equality

```
>>> fp1 = Fingerprint([0, 1, 6, 8, 12], bits=16)
>>> fp2 = Fingerprint([1, 2, 4, 8, 11, 12], bits=16)
>>> fp1 == fp2
False
>>> fp1_copy = Fingerprint.from_fingerprint(fp1)
>>> fp1 == fp1_copy
True
>>> fp1_copy.level = 5
>>> fp1 == fp1_copy
False
```

### Union/OR

```
>>> fp1 + fp2
Fingerprint(indices=array([0, 1, 2, 4, 6, 8, 11, 12]), level=-1, bits=16, ↵
˓→name=None)
>>> fp1 | fp2
Fingerprint(indices=array([0, 1, 2, 4, 6, 8, 11, 12]), level=-1, bits=16, ↵
˓→name=None)
```

### Intersection/AND

```
>>> fp1 & fp2
Fingerprint(indices=array([1, 8, 12]), level=-1, bits=16, name=None)
```

### Difference/AND NOT

```
>>> fp1 - fp2
Fingerprint(indices=array([0, 6]), level=-1, bits=16, name=None)
>>> fp2 - fp1
Fingerprint(indices=array([2, 4, 11]), level=-1, bits=16, name=None)
```

### XOR

```
>>> fp1 ^ fp2
Fingerprint(indices=array([0, 2, 4, 6, 11]), level=-1, bits=16, name=None)
```

With count or float fingerprints, bit-wise operations are still possible, but algebraic operations are applied to counts.

```
>>> fp1 = CountFingerprint(counts={0: 3, 1: 2, 5: 1, 9: 3}, bits=16)
>>> fp2 = CountFingerprint(counts={1: 2, 5: 2, 7: 3, 10: 7}, bits=16)
>>> fp1 + fp2
CountFingerprint(counts={0: 3, 1: 4, 5: 3, 7: 3, 9: 3, 10: 7}, level=-1, bits=16, ↵
˓→name=None)
>>> fp1 - fp2
CountFingerprint(counts={0: 3, 1: 0, 5: -1, 7: -3, 9: 3, 10: -7}, level=-1, bits=16, ↵
˓→name=None)
>>> fp1 * 3
CountFingerprint(counts={0: 9, 1: 6, 5: 3, 9: 9}, level=-1, bits=16, name=None)
>>> fp1 / 2
FloatFingerprint(counts={0: 1.5, 1: 1.0, 5: 0.5, 9: 1.5}, level=-1, bits=16, ↵
˓→name=None)
```

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Finally, fingerprints may be batch added and averaged, producing either a count or float fingerprint when sensible.

```
>>> from e3fp.fingerprint.fprint import add, mean
>>> fps = [Fingerprint(np.random.randint(0, 32, 8), bits=32) for i in range(100)]
>>> add(fps)
CountFingerprint(counts={0: 23, 1: 23, ..., 30: 20, 31: 14}, level=-1, bits=32,
    ↪name=None)
>>> mean(fps)
FloatFingerprint(counts={0: 0.23, 1: 0.23, ..., 30: 0.2, 31: 0.14}, level=-1, bits=32,
    ↪ name=None)
```

## Fingerprint Storage

The most efficient way to store and interact with fingerprints is through the `e3fp.fingerprint.db.FingerprintDatabase` class. This class wraps a matrix with sparse rows (`scipy.sparse.csr_matrix`), where each row is a fingerprint. This enables rapid I/O of the database while also minimizing the memory footprint. Accessing the underlying sparse representation with the `.FingerprintDatabase.array` attribute is convenient for machine learning purposes, while the database class itself provides several useful functions.

---

**Note:** We strongly recommend upgrading to at least SciPy v1.0.0 when working with large fingerprint databases, as old versions are much slower and have several bugs for database loading.

---

## Database I/O and Indexing

See the full `e3fp.fingerprint.db.FingerprintDatabase` documentation for a description of basic database usage, attributes, and methods. Below, several additional use cases are documented.

## Batch Database Operations

Due to the sparse representation of the underlying data structure, an un-folded database, a database with unfolded fingerprints does not use significantly more disk space than a database with folded fingerprints. However, it is usually necessary to fold fingerprints for machine learning tasks. The `FingerprintDatabase` does this very quickly.

```
>>> from e3fp.fingerprint.db import FingerprintDatabase
>>> from e3fp.fingerprint.fprint import Fingerprint
>>> import numpy as np
>>> db = FingerprintDatabase(fp_type=Fingerprint, name="TestDB")
>>> print(db)
FingerprintDatabase[name: TestDB, fp_type: Fingerprint, level: -1, bits: None, fp_
    ↪num: 0]
>>> on_inds = [np.random.uniform(0, 2**32, size=30) for i in range(5)]
>>> fps = [Fingerprint(x, bits=2**32) for x in on_inds]
>>> db.add_fingerprints(fps)
>>> print(db)
FingerprintDatabase[name: TestDB, fp_type: Fingerprint, level: -1, bits: 4294967296,
    ↪fp_num: 5]
>>> db.get_density()
6.984919309616089e-09
```

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```
>>> fold_db = db.fold(1024)
>>> print(fold_db)
FingerprintDatabase[name: TestDB, fp_type: Fingerprint, level: -1, bits: 1024, fp_
↪num: 5]
>>> fold_db.get_density()
0.0287109375
```

A database can be converted to a different fingerprint type:

```
>>> from e3fp.fingerprint.fprint import CountFingerprint
>>> count_db = db.as_type(CountFingerprint)
>>> print(count_db)
FingerprintDatabase[name: TestDB, fp_type: CountFingerprint, level: -1, bits:_
↪4294967296, fp_num: 5]
>>> count_db[0]
CountFingerprint(counts={2977004690: 1, ..., 3041471738: 1}, level=-1,_
↪bits=4294967296, name=None)
```

The `e3fp.fingerprint.db.concat` method allows efficient joining of multiple databases.

```
>>> from e3fp.fingerprint.db import concat
>>> dbs = []
>>> for i in range(10):
...     db = FingerprintDatabase(fp_type=Fingerprint)
...     on_inds = [np.random.uniform(0, 1024, size=30) for j in range(5)]
...     fps = [Fingerprint(x, bits=2**32, name="Mol{}".format(i)) for x in on_inds]
...     db.add_fingerprints(fps)
...     dbs.append(db)
>>> dbs[0][0]
Fingerprint(indices=array([94, 97, ..., 988, 994]), level=-1, bits=4294967296,_
↪name=Mol0)
>>> print(dbs[0])
FingerprintDatabase[name: None, fp_type: Fingerprint, level: -1, bits: 4294967296, fp_
↪num: 5]
>>> merge_db = concat(dbs)
>>> print(merge_db)
FingerprintDatabase[name: None, fp_type: Fingerprint, level: -1, bits: 4294967296, fp_
↪num: 50]
```

## Database Comparison

Two databases may be compared using various metrics in `e3fp.fingerprint.metrics`. Additionally, all fingerprints in a database may be compared to each other simply by only providing a single database. See [Fingerprint Comparison](#) for more details.

## Performing Machine Learning on the Database

The underlying sparse matrix may be passed directly to machine learning tools in any package that is compatible with SciPy sparse matrices, such as `scikit-learn`.

```
>>> from sklearn.naive_bayes import BernoulliNB
>>> clf = BernoulliNB()
>>> clf.fit(db.array, ypred)
BernoulliNB(alpha=1.0, binarize=0.0, class_prior=None, fit_prior=True)
>>> clf.predict(db2.array)
...
```

## Fingerprint Comparison

The `e3fp.fingerprint.metrics` sub-package provides several useful methods for batch comparison of fingerprints in various representations.

### Fingerprint Metrics

These metrics operate directly on pairs of `Fingerprint` and `FingerprintDatabase` objects or on a combination of each. If only a single variable is specified, self-comparison is performed. The implemented methods are common functions for fingerprint similarity in the literature.

### Array Metrics

To efficiently compare fingerprint databases above, we provide comparison metrics that can operate directly on the internal sparse matrix representation without the need to “densify it”. We describe these here, as they have several additional features.

The array metrics implemented in `e3fp.fingerprint.metrics.array_metrics` are implemented such that they may take any combination of dense and sparse inputs. Additionally, they are designed to function as `scikit-learn-compatible kernels` for machine learning tasks. For example, one might perform an analysis using a support vector machine (SVM) and Tanimoto kernel.

```
>>> from sklearn.svm import SVC
>>> from e3fp.fingerprint.metrics.array_metrics import tanimoto
>>> clf = SVC(kernel=tanimoto)
>>> clf.fit(X, y)
...
>>> clf.predict(test)
...
```

Most common fingerprint comparison metrics only apply to binary fingerprints. We include several that operate equally well on count- and float-based fingerprints. For example, to our knowledge, we provide the only open source implementation of Soergel similarity, the analog to the Tanimoto coefficient for non-binary fingerprints that can efficiently operate on sparse inputs.

```
>>> from e3fp.fingerprint.metrics.array_metrics import soergel
>>> clf = SVC(kernel=soergel)
>>> clf.fit(X, y)
...
>>> clf.predict(test)
...
```

## 1.4 Developer Notes

We welcome contributions to E3FP! These notes are designed to help developers contribute code

### 1.4.1 Authoring Code

#### Code Formatting

E3FP's code should be *readable*. To ensure this, we rigorously follow the [PEP8](#) style conventions and [PEP257](#) docstring conventions, which maximize readability of the code and ease of future development. You may check your code for conformation to these conventions with the [pycodestyle](#) and [pydocstyle](#) utilities, respectively. Where the code is necessarily complicated, inline comments should reorient the reader.

#### Utility Methods and Classes

Three sets of utility methods and classes are provided: [`e3fp.util`](#), [`e3fp.conformer.util`](#), and [`e3fp.fingerprint.util`](#). These provide general and often-used functionality in their corresponding packages. Additionally, they provide E3FP-specific errors and exceptions.

#### Warnings and Errors

By default, warnings in Python are silent. We therefore provide a warning base class [`e3fp.util.E3FPWarning`](#) that is not silent by default. We provide several general warnings:

[`E3FPDeprecationWarning`](#) warns when a deprecated method is called or class is instantiated.

**See also:**

[`Deprecation`](#)

[`E3FPEfficiencyWarning`](#) warns when a method, module version, or combination of parameters is known to be inefficient.

---

**Note:** If possible, the warning message should advise on a more efficient approach.

---

E3FP-specific errors should inherit [`e3fp.util.E3FPError`](#) base class. Several fingerprinting-specific errors are defined in [`e3fp.fingerprint.util`](#).

#### Deprecation

Whenever changing the interface or behavior of a user-facing method or class, it is proper to deprecate it for at least one release, so that the users have time to update their scripts accordingly. A deprecated method should providing an [`e3fp.util.E3FPDeprecationWarning`](#), notifying the user in which release to expect the method or class to be removed, and updating the documentation accordingly. This functionality is automated with the [`e3fp.util.deprecated`](#) decorator, as shown in this example:

```
>>> import sys
>>> sys.stderr = sys.stdout
>>> from e3fp.util import deprecated
>>> @deprecated("1.1", remove_version="1.3", msg="Function no longer needed.")
... def deprecated_method():
```

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```

...
    """A method to demonstrate method deprecation."""
...
    pass
>>> deprecated_method()
...: E3FPDeprecationWarning: Function `my_function` was deprecated in 1.1 and will be
...: removed in 1.3. Function no longer needed.

```

In the api documentation, the method will appear as:

**deprecated\_method()**

---

**Note:** Deprecated in e3fp 1.1. *deprecated\_method* will be removed in e3fp 1.3. Function no longer needed.

---

A method to demonstrate method deprecation.

---

**Note:** If no *remove\_version* is specified, then the remove version defaults to the next release after deprecation. For example, if the method was deprecated in 1.1, it is by default marked for removal in 1.2.

---

## Contributing Code

Before contributing code to E3FP, it is advisable for major modifications to submit an issue to the [issue tracker](#) to enable other developers to contribute to the design of the code and to reduce the amount of work necessary to conform the code to E3FP's standards. After writing the code, create a [pull request](#). This is best even if you have push access to the E3FP repo, as it enables the test suite to be run on the new code prior to merging it with the remaining code base.

## Writing Tests

The standard in E3FP is to commit a test for new functionality simultaneously with the new functionality or within the same pull request. While this slows development, it prevents building a large backlog of untested methods and classes.

These should ideally be unit tests, though for some complicated functionalities, such as fingerprinting, integration tests are also necessary. For these complicated functions, specific units may still be tested using [unittest.mock](#). For example, `unittest.mock.patch()` may be used to force a high level method to produce a specific output. For examples, see the [fingerprinting tests](#).

## Continuous Integration

E3FP uses [Travis CI](#) for continuous integration. This ensures that each commit and pull request passes all tests on a variety of systems and for all supported versions of Python. Additionally, Travis CI updates code coverage on [Coveralls](#) and tests all usage examples in the documentation using [doctest](#).

## 1.4.2 Documentation

In general, it is best to document the rationale and basic usage of a module, class, or method in its docstring instead of in a separate documentation file. See, for example, the docstring for `e3fp.fingerprint.db.FingerprintDatabase`. We use a variety of tools to ensure that our documentation is always up-to-date. The official documentation is hosted on [ReadtheDocs](#) and is automatically generated when new code is committed to the repository.

### Documenting Code

E3FP uses NumPy's docstring conventions for all docstrings. These are parsed by Sphinx using Napoleon. All usage examples must be fully functional, as these are tested using `doctest`.

The purpose of a docstring is to explain the purpose of a class/method, any relevant implementation details, its parameters, its attributes, its outputs, and its usage. The goal is clarity. For self-evident methods with descriptive variables, a simple one-line summary is all that is needed. For complicated use cases, often involving other methods/classes, it is better to document the usage elsewhere in the documentation.

### Documentation Usage

Coming soon.

## 1.4.3 Releasing Code

## 1.5 e3fp API

### 1.5.1 e3fp package

#### Subpackages

#### e3fp.config package

#### Submodules

#### e3fp.config.params module

Get E3FP default parameters and read parameters from files.

Author: Seth Axen E-mail: [seth.axen@gmail.com](mailto:seth.axen@gmail.com)

`get_default_value(*args, **kwargs)`

`get_value(params, section_name, param_name, dtype=<class 'str'>, auto=False, fallback=None)`

Get value from params with fallback.

#### Parameters

- **params** (*SafeConfigParser*) – Parameters
- **section\_name** (*str*) – Name of section in *params*
- **param\_name** (*str*) – Name of parameter in *section*
- **dtype** (*type, optional*) – Type to return data as.

- **auto** (*bool, optional*) – Auto-discover type of value. If provided, *dtype* is ignored.
- **fallback** (*any, optional*) – Value to return if getting value fails.

**Returns** `value` – Value of parameter or *fallback*.

**Return type** `any`

**params\_to\_sections\_dict** (*params, auto=True*)

Get dict of sections dicts in *params*, with optional type discovery.

**Parameters**

- **params** (*str or SafeConfigParser*) – Params to read
- **auto** (*bool, optional*) – Auto typing of parameter values.

**Returns** `dict`

**Return type** dict matching sections to parameters to values.

**read\_params** (*params=None, fill\_defaults=False*)

Get combination of provided parameters and default parameters.

**Parameters**

- **params** (*str or SafeConfigParser, optional*) – User provided parameters as an INI file or `SafeConfigParser`. Any parameters provided will replace default parameters.
- **fill\_defaults** (*bool, optional*) – Fill values that aren't provided with package defaults, if *params* is file.

**Returns** `all_params` – Combination of default and user-provided parameters.

**Return type** `SafeConfigParser`

**update\_params** (*params\_dict, params=None, section\_name=None, fill\_defaults=False*)

Set `SafeConfigParser` values from a sections dict.

Sections dict key must be parameter sections, and value must be dict matching parameter name to value. If existing `SafeConfigParser` is provided, parameter values are updated.

**Parameters**

- **params\_dict** (*dict*) – If *section\_name* is provided, dict must match parameter names to values. If *section\_name* is not provided, dict key(s) must be parameter sections, and value(s) must be parameter dict.
- **params** (*SafeConfigParser, optional*) – Existing parameters.
- **section\_name** (*str, optional*) – Name of section to which to add parameters in *params\_dict*
- **fill\_defaults** (*bool, optional*) – Fill values that aren't provided with package defaults, if *params* is file.

**write\_params** (*params, params\_file='params.cfg'*)

Write params to file.

**Parameters**

- **params** (*SafeConfigParser*) – Params
- **params\_file** (*str*) – Params file

## Module contents

### e3fp.conformer package

#### Submodules

##### e3fp.conformer.generate module

Generate conformers from SMILES or MOL2 files.

Author: Seth Axen E-mail: [seth.axen@gmail.com](mailto:seth.axen@gmail.com)

```
generate_conformers(input_mol, name=None, standardise=False, num_conf=-1,
                      pool_multiplier=1, rmsd_cutoff=0.5, max_energy_diff=None, forcefield='uff',
                      seed=-1, out_file=None, out_dir='conformers', save=False, compress=2,
                      overwrite=False)
```

Generate and save conformers for molecules.

#### Parameters

- **input\_mol** (*RDKit Mol*) – Mol with a single conformer from which to generate conformers.
- **name** (*str, optional*) – Name of molecule.
- **standardise** (*bool, optional*) – Standardise mol before generating conformers.
- **num\_conf** (*int, optional*) – If int, this is the target number of conformations. If -1, number of conformations is automatically chosen based on number of rotatable bonds.
- **first** (*int, optional*) – Number of first conformers to return. Does not impact conformer generator process, except may terminate conformer generation early when this many of conformers have been accepted.
- **pool\_multiplier** (*int, optional*) – Factor to multiply by *num\_conf*. The resulting number of conformations will be generated, then pruned to *num\_conf*.
- **rmsd\_cutoff** (*float, optional*) – RMSD threshold above which to accept two conformations as different
- **max\_energy\_diff** (*float, optional*) – Maximum energy difference between lowest energy conformer and any accepted conformer.
- **forcefield** (*{'uff', 'mmff94', 'mmff94s'}*, *optional*) – Forcefield to use for minimization of conformers.
- **seed** (*int, optional*) – Random seed for conformer generation. If -1, the random number generator is unseeded.
- **out\_file** (*str, optional*) – Filename to save output, if *save* is True. If None, filename will be *name.sdf*, optionally with a compressed extension.
- **out\_dir** (*str, optional*) – Directory where output files will be saved if *save* is True.
- **save** (*bool, optional*) – Save conformers to *out\_file* in *out\_dir*.
- **compress** (*int, optional*) – Compression of SDF files. None: auto. Mode is chosen based on extension, defaulting to SDF. 0: File is not compressed. 1: File is gzipped (.gz) 2: File is bzipped (.bz2)
- **overwrite** (*bool, optional*) – Overwrite output files if they already exist.

#### Returns

- *bool* – If something went wrong, only return False. Otherwise return below.
- *tuple* – Tuple with molecule name, number of rotatable bonds, numpy array of indices of final conformations, numpy array of energies of all conformations generated, and 2D numpy array of pairwise RMSDs between final conformations.

**run** (*mol2=None*, *smiles=None*, *standardise=False*, *num\_conf=-1*, *first=-1*, *pool\_multiplier=1*, *rmsd\_cutoff=0.5*, *max\_energy\_diff=None*, *forcefield='uff'*, *seed=-1*, *params=None*, *prioritize=False*, *out\_dir='conformers'*, *compress='2'*, *overwrite=False*, *values\_file=None*, *log=None*, *num\_proc=None*, *parallel\_mode=None*, *verbose=False*)  
Run conformer generation.

**values\_to\_hdf5** (*hdf5\_buffer*, *values*)  
Write specific values to *hdf5\_buffer*.

#### Parameters

- **hdf5\_buffer** (*HDF5Buffer*) – HDF5 buffer to write to.
- **values** (*tuple*) – Tuple of values to be written to buffer. Values and types should be *name* (str), *num\_rotatable\_bonds* (int), *target\_conformers* (int), *indices* (numpy long array), *energies* (numpy float array), and *rmsd* (numpy float array). *rmsd* should be a square array with size along each dimension equal to length of *indices*.

**Returns** True if success, False if not.

**Return type** *bool*

## e3fp.conformer.generator module

Conformer generation.

Author: Seth Axen E-mail: [seth.axen@gmail.com](mailto:seth.axen@gmail.com)

**class ConformerGenerator** (*num\_conf=-1*, *first=-1*, *rmsd\_cutoff=0.5*, *max\_energy\_diff=-1.0*, *forcefield='uff'*, *pool\_multiplier=1*, *seed=-1*, *get\_values=False*, *sparse\_rmsd=True*, *store\_energies=True*)

Bases: *object*

Generate conformers using RDKit.

1. Generate a pool of conformers.
2. Minimize conformers.
3. Filter conformers using an RMSD threshold and optional minimum energy difference.

Note that pruning is done \_after\_ minimization, which differs from the protocol described in the references.

## References

- [#working-with-3d-molecules](http://rdkit.org/docs/GettingStartedInPython.html)
- <http://pubs.acs.org/doi/full/10.1021/ci2004658>
- [https://github.com/skearnes/rdkit-utils/blob/master/rdkit\\_utils/conformers.py](https://github.com/skearnes/rdkit-utils/blob/master/rdkit_utils/conformers.py)

**embed\_molecule** (*mol*)

Generate conformers, possibly with pruning.

**Parameters** *mol* (*RDKit Mol*) – Molecule.

**filter\_conformers** (*mol*)

Filter conformers which do not meet an RMSD threshold.

**Parameters** *mol* (*RDKit Mol*) – Molecule.

**Returns**

- A new *RDKit Mol* containing the chosen conformers, sorted by
- increasing energy.

**generate\_conformers** (*mol*)

Generate conformers for a molecule.

**Parameters** *mol* (*RDKit Mol*) – Molecule.

**Returns** **RDKit Mol**

**Return type** copy of the input molecule with embedded conformers

**get\_conformer\_energies** (*mol*)

Calculate conformer energies.

**Parameters** *mol* (*RDKit Mol*) – Molecule.

**Returns** **energies** – Minimized conformer energies.

**Return type** array\_like

**get\_molecule\_force\_field** (*mol*, *conf\_id=None*, *\*\*kwargs*)

Get a force field for a molecule.

**Parameters**

- **mol** (*RDKit Mol*) – Molecule.
- **conf\_id** (*int, optional*) – ID of the conformer to associate with the force field.
- **\*\*kwargs** (*dict, optional*) – Keyword arguments for force field constructor.

**static get\_num\_conformers** (*mol*)

Return ideal number of conformers from rotatable bond number in model.

**Parameters** *mol* (*Mol*) – RDKit *Mol* object for molecule

**Yields** **num\_conf** (*int*) – Target number of conformers to accept

**minimize\_conformers** (*mol*)

Minimize molecule conformers.

**Parameters** *mol* (*RDKit Mol*) – Molecule.

**static reverse\_enumerate** (*iterable*)

Enumerate, but with the last result first but still numbered last.

**Parameters** *iterable* (*some 1-D iterable*)

**Returns** Reverse of *enumerate* function

**Return type** iterable

## e3fp.conformer.protonation module

Functions for generating protonation states of molecules.

Author: Seth Axen E-mail: [seth.axen@gmail.com](mailto:seth.axen@gmail.com)

```
smiles_dict_to_proto_smiles_dict (in_smiles_dict, max_states=3, pka=7.4, dist_cutoff=20.0,
add_missing=False, parallelizer=None, chunk_size=100)
```

Generate dict of SMILES for protonated states from SMILES dict.

```
smiles_list_to_proto_smiles_list (in_smiles_list, max_states=3, pka=7.4, dist_cutoff=20.0)
```

Generate list of SMILES for protonated states from single SMILES.

```
smiles_to_proto_smiles (smiles, mol_name, max_states=3, pka=7.4, dist_cutoff=20.0)
```

Generate list of SMILES for protonated states from single SMILES.

This is very inefficient in batch.

## e3fp.conformer.util module

Utilities for handling SMILES strings and RDKit mols and conformers.

Author: Seth Axen E-mail: [seth.axen@gmail.com](mailto:seth.axen@gmail.com)

```
class MolItemName (mol_name=None, proto_state_num=None, conf_num=None, proto_delim='-', conf_delim='_')
```

Bases: `object`

Class for parsing mol item names and converting to various formats.

```
property conf_name
```

```
copy()
```

```
classmethod from_str (mol_item_name, mol_item_regex=re.compile('(?P<mol_name>.+?)(?::-(?P<proto_state_num>\d+))?(?:_(?P<conf_num>\d+))?$'), mol_item_fields=('mol_name', 'proto_state_num', 'conf_num'), **kwargs)
```

```
classmethod from_tuple (fields_tuple)
```

```
property mol_item_name
```

```
static mol_item_name_to_dict (mol_item_name, mol_item_regex=re.compile('(?P<mol_name>.+?)(?::-(?P<proto_state_num>\d+))?(?:_(?P<conf_num>\d+))?$'), mol_item_fields=('mol_name', 'proto_state_num', 'conf_num'))
```

```
property mol_name
```

```
property proto_name
```

```
to_conf_name (conf_num=None, conf_delim='_')
```

```
to_mol_name (as_proto=False)
```

```
to_proto_name (proto_state_num=None, proto_delim='-')
```

```
to_str()
```

```
to_tuple()
```

```
class MolItemTuple (mol_name, proto_state_num, conf_num)
```

Bases: `tuple`

**property conf\_num**  
Alias for field number 2

**property mol\_name**  
Alias for field number 0

**property proto\_state\_num**  
Alias for field number 1

**add\_conformer\_energies\_to\_mol** (*mol, energies*)  
Add conformer energies as mol property.

See discussion at <https://sourceforge.net/p/rdkit/mailman/message/27547551/>

**dict\_to\_smiles** (*smiles\_file, smiles\_dict*)  
Write SMILES dict to file.

**get\_conformer\_energies\_from\_mol** (*mol*)  
Get conformer energies from mol.

**iter\_to\_smiles** (*smiles\_file, smiles\_iter*)  
Write iterator of (mol\_name, SMILES) to file.

**mol2\_generator** (\**filenames*)  
Parse name from mol2 filename and return generator.

**Parameters** **files** (*iterable object*) – List of mol2 files, where filename should be molecule name followed by “.mol2”

**Yields** *tuple* – *tuple* of the format (file, name).

**mol\_from\_mol2** (*mol2\_file, name=None, standardise=False*)  
Read a mol2 file into an RDKit PropertyMol.

#### Parameters

- **mol2\_file** (*str*) – path to a mol2 file
- **name** (*str; optional*) – Name of molecule. If not provided, uses file basename as name
- **standardise** (*bool*) – Clean mol through standardisation

**Returns** RDKit PropertyMol

**Return type** Molecule.

**mol\_from\_sdf** (*sdf\_file, conf\_num=None, standardise=False*)  
Read SDF file into an RDKit Mol object.

#### Parameters

- **sdf\_file** (*str*) – Path to an SDF file
- **conf\_num** (*int or None, optional*) – Maximum number of conformers to read from file. Defaults to all.
- **standardise** (*bool (default False)*) – Clean mol through standardisation

**Returns** RDKit Mol

**Return type** Mol object with each molecule in SDF file as a conformer

**mol\_from\_smiles** (*smiles, name, standardise=False*)  
Generate a n RDKit PropertyMol from SMILES string.

#### Parameters

- **smile** (*str*) – SMILES string
- **name** (*str*) – Name of molecule
- **standardise** (*bool*) – Clean Mol through standardisation

**Returns** RDKit PropertyMol

**Return type** Molecule.

**mol\_to\_sdf** (*mol*, *out\_file*, *conf\_num=None*)

Write RDKit Mol objects to an SDF file.

#### Parameters

- **mol** (RDKit Mol) – A molecule containing 1 or more conformations to write to file.
- **out\_file** (*str*) – Path to save SDF file.
- **conf\_num** (*int or None, optional*) – Maximum number of conformers to save to file. Defaults to all.

**mol\_to\_standardised\_mol** (*mol*, *name=None*)

Standardise mol(s).

**smiles\_generator** (\**filenames*)

Parse SMILES file(s) and yield (name, smile).

**Parameters** **files** (*iterable object*) – List of files containing smiles. File must contain one smile per line, followed by a space and then the molecule name.

**Yields** *tuple* – *tuple* of the format (smile, name).

**smiles\_to\_dict** (*smiles\_file*, *unique=False*, *has\_header=False*)

Read SMILES file to dict.

## Module contents

### e3fp.fingerprint package

#### Subpackages

### e3fp.fingerprint.metrics package

#### Submodules

### e3fp.fingerprint.metrics.array\_metrics module

Fingerprint array comparison metrics.

Each is fully compatible with both dense and sparse inputs.

Author: Seth Axen E-mail: [seth.axen@gmail.com](mailto:seth.axen@gmail.com)

**cosine** (*X*, *Y=None*, *assume\_binary=False*)

Compute the Cosine similarities between *X* and *Y*.

#### Parameters

- **X** (*array\_like or sparse matrix*) – with shape (*n\_fprints\_X*, *n\_bits*).

- **Y** (*array\_like or sparse matrix, optional*) – with shape  $(n\_fprints\_Y, n\_bits)$ .
- **assume\_binary** (*bool, optional*) – Assume data is binary (results in efficiency boost). If data is not binary, the result will be incorrect.

**Returns cosine**

**Return type** array of shape  $(n\_fprints\_X, n\_fprints\_Y)$

**See also:**

`dice()`, `soergel()`, `tanimoto()`

**dice** ( $X, Y=None$ )

Compute the Dice coefficients between  $X$  and  $Y$ .

Data must be binary. This is not checked.

**Parameters**

- **X** (*array\_like or sparse matrix*) – with shape  $(n\_fprints\_X, n\_bits)$ .
- **Y** (*array\_like or sparse matrix, optional*) – with shape  $(n\_fprints\_Y, n\_bits)$ .

**Returns dice**

**Return type** array of shape  $(n\_fprints\_X, n\_fprints\_Y)$

**See also:**

`cosine()`, `soergel()`, `tanimoto()`, `pearson()`

**pearson** ( $X, Y=None$ )

Compute the Pearson correlation between  $X$  and  $Y$ .

**Parameters**

- **X** (*array\_like or sparse matrix*) – with shape  $(n\_fprints\_X, n\_bits)$ .
- **Y** (*array\_like or sparse matrix, optional*) – with shape  $(n\_fprints\_Y, n\_bits)$ .

**Returns pearson**

**Return type** array of shape  $(n\_fprints\_X, n\_fprints\_Y)$

**See also:**

`soergel()` Soergel similarity for non-binary data

`cosine()`, `dice()`, `tanimoto()`

**soergel** ( $X, Y=None$ )

Compute the Soergel similarities between  $X$  and  $Y$ .

Soergel similarity is the complement of Soergel distance and can be thought of as the analog of the Tanimoto coefficient for count/float-based data. For binary data, it is equivalent to the Tanimoto coefficient.

**Parameters**

- **X** (*array\_like or sparse matrix*) – with shape  $(n\_fprints\_X, n\_bits)$ .
- **Y** (*array\_like or sparse matrix, optional*) – with shape  $(n\_fprints\_Y, n\_bits)$ .

**Returns soergel**

**Return type** array of shape  $(n\_fprints\_X, n\_fprints\_Y)$

**See also:**

`tanimoto()` A fast version of this function for binary data.

`pearson()` Pearson correlation, also appropriate for non-binary data.

`cosine()`, `dice()`

`tanimoto(X, Y=None)`

Compute the Tanimoto coefficients between  $X$  and  $Y$ .

Data must be binary. This is not checked.

#### Parameters

- $\mathbf{X}$  (*array\_like or sparse matrix*) – with shape  $(n\_fprints_X, n\_bits)$ .
- $\mathbf{Y}$  (*array\_like or sparse matrix, optional*) – with shape  $(n\_fprints_Y, n\_bits)$ .

#### Returns `tanimoto`

**Return type** array of shape  $(n\_fprints_X, n\_fprints_Y)$

#### See also:

`soergel()` Analog to Tanimoto for non-binary data.

`cosine()`, `dice()`, `pearson()`

## e3fp.fingerprint.metrics.fprint\_metrics module

Fingerprint comparison metrics.

Author: Seth Axen E-mail: [seth.axen@gmail.com](mailto:seth.axen@gmail.com)

`cosine(fp1, fp2)`

Calculate cosine similarity between fingerprints.

#### Parameters

- `fp1` (*Fingerprint*) – Fingerprint 1
- `fp2` (*Fingerprint*) – Fingerprint 2

#### Returns `float`

**Return type** Cosine similarity.

`dice(fp1, fp2)`

Calculate Dice coefficient between fingerprints.

#### Parameters

- `fp1` (*Fingerprint*) – Fingerprint 1
- `fp2` (*Fingerprint*) – Fingerprint 2

#### Returns `float`

**Return type** Dice coefficient.

`distance(fp1, fp2)`

Calculate Euclidean distance between fingerprints.

#### Parameters

- `fp1` (*Fingerprint*) – Fingerprint 1

- **fp2** (*Fingerprint*) – Fingerprint 2

**Returns float**

**Return type** Euclidian distance.

**hamming** (*fp1*, *fp2*)

Calculate Hamming distance between fingerprints.

**Parameters**

- **fp1** (*Fingerprint*) – Fingerprint 1
- **fp2** (*Fingerprint*) – Fingerprint 2

**Returns float**

**Return type** Hamming distance.

**pearson** (*fp1*, *fp2*)

Calculate Pearson correlation between fingerprints.

**Parameters**

- **fp1** (*Fingerprint*) – Fingerprint 1
- **fp2** (*Fingerprint*) – Fingerprint 2

**Returns float**

**Return type** Pearson correlation.

**soergel** (*fp1*, *fp2*)

Calculate Soergel similarity between fingerprints.

Soergel similarity is the complement of Soergel distance and can be thought of as the analog of the Tanimoto coefficient for count/float-based fingerprints. For *Fingerprint*, it is equivalent to the Tanimoto coefficient.

**Parameters**

- **fp1** (*Fingerprint*) – Fingerprint 1
- **fp2** (*Fingerprint*) – Fingerprint 2

**Returns**

- **float** (*Soergel similarity*.)
- *Reference*
- \_\_\_\_\_

**tanimoto** (*fp1*, *fp2*)

Calculate Tanimoto coefficient between fingerprints.

**Parameters**

- **fp1** (*Fingerprint*) – Fingerprint 1
- **fp2** (*Fingerprint*) – Fingerprint 2

**Returns float**

**Return type** Tanimoto coefficient.

## Module contents

Efficient comparison metrics for fingerprints and their databases.

Author: Seth Axen E-mail: [seth.axen@gmail.com](mailto:seth.axen@gmail.com)

### **cosine** (*A, B=None*)

Compute cosine similarities between fingerprints.

Fingerprints must have same number of bits. If only one fingerprint/database is provided, it is compared to self.

**Parameters** **A, B** (*Fingerprint or FingerprintDatabase*) – Fingerprint(s) to be compared

**Returns** cosine

**Return type** float or ndarray [shape (num\_fps\_A, num\_fps\_B)]

**See also:**

[dice\(\)](#), [pearson\(\)](#), [soergel\(\)](#), [tanimoto\(\)](#)

### **dice** (*A, B=None*)

Compute Dice coefficients between fingerprints.

Fingerprints must have same number of bits. If not bit-fingerprints, arrays will be cast to binary. If only one fingerprint/database is provided, it is compared to self.

**Parameters** **A, B** (*Fingerprint or FingerprintDatabase*) – Fingerprint(s) to be compared

**Returns** dice

**Return type** float or ndarray [shape (num\_fps\_A, num\_fps\_B)]

**See also:**

[cosine\(\)](#), [pearson\(\)](#), [soergel\(\)](#), [tanimoto\(\)](#)

### **pearson** (*A, B=None*)

Compute Pearson correlation between fingerprints.

Fingerprints must have same number of bits. If only one fingerprint/database is provided, it is compared to self.

**Parameters** **A, B** (*Fingerprint or FingerprintDatabase*) – Fingerprint(s) to be compared

**Returns** pearson

**Return type** float or ndarray [shape (num\_fps\_A, num\_fps\_B)]

**See also:**

[cosine\(\)](#), [dice\(\)](#), [soergel\(\)](#), [tanimoto\(\)](#)

### **soergel** (*A, B=None*)

Compute Soergel similarities between fingerprints.

Soergel similarity is the complement of the Soergel distance and is analogous to the Tanimoto coefficient for count/float fingerprints. For binary data, it is equivalent to [tanimoto](#).

**Parameters** **A, B** (*Fingerprint or FingerprintDatabase*) – Fingerprint(s) to be compared

**Returns** soergel

**Return type** float or ndarray [shape (num\_fps\_A, num\_fps\_B)]

**See also:**

[cosine\(\)](#), [dice\(\)](#), [pearson\(\)](#), [tanimoto\(\)](#)

**tanimoto** (*A, B=None*)

Compute Tanimoto coefficients between fingerprints.

Fingerprints must have same number of bits. If not bit-fingerprints, arrays will be cast to binary. For non-binary data, use `soergel`. If only one fingerprint/database is provided, it is compared to self.

**Parameters** **A, B** (*Fingerprint or FingerprintDatabase*) – Fingerprint(s) to be compared

**Returns** **tanimoto** – Pairwise tanimoto(s) between fingerprint(s) in *A* and *B*.

**Return type** float or ndarray [shape (num\_fps\_A, num\_fps\_B)]

**See also:**

`cosine()`, `dice()`, `pearson()`, `soergel()`

## Submodules

### e3fp.fingerprint.array\_ops module

Various array operations.

Author: Seth Axen E-mail: [seth.axen@gmail.com](mailto:seth.axen@gmail.com)

**as\_unit** (*v, axis=1*)

Return array of unit vectors parallel to vectors in *v*.

**Parameters**

- **v** (ndarray of float)
- **axis** (int, optional) – Axis along which to normalize length.

**Returns** ndarray of float – magnitude along *axis*.

**Return type** Unit vector of *v*, i.e. *v* divided by its

**calculate\_angles** (*vec\_arr, ref, ref\_norm=None*)

Calculate angles between vectors in *vec\_arr* and *ref* vector.

If *ref\_norm* is not provided, angle ranges between 0 and pi. If it is provided, angle ranges between 0 and 2pi. Note that if *ref\_norm* is orthogonal to *vec\_arr* and *ref*, then the angle is rotation around the axis, but if a non-orthogonal axis is provided, this may not be the case.

**Parameters**

- **vec\_arr** (Nx3 array of float) – Array of N 3D vectors.
- **ref** (1x3 array of float) – Reference vector
- **ref\_norm** (1x3 array of float) – Normal vector.

**Returns** Array of N angles

**Return type** 1-D array

**make\_distance\_matrix** (*coords*)

Build pairwise distance matrix from coordinates.

**Parameters** **coords** (ndarray of float) – an Mx3 array of cartesian coordinates.

**Returns** ndarray of float

**Return type** square symmetrical distance matrix

**make\_rotation\_matrix**(*v0, v1*)

Create 3x3 matrix of rotation from *v0* onto *v1*.

Should be used by dot(R, v0.T).T.

**Parameters**

- **v0** (*1x3 array of float*) – Initial vector before alignment.
- **v1** (*1x3 array of float*) – Vector to which to align *v0*.

**make\_transform\_matrix**(*center, y=None, z=None*)

Make 4x4 homogenous transformation matrix.

Given Nx4 array A where A[:, 4] = 1., the transform matrix M should be used with dot(M, A.T).T. Order of operations is 1. translation, 2. align y x z plane to yz-plane 3. align y to y-axis.

**Parameters**

- **center** (*1x3 array of float*) – Coordinate that should be centered after transformation.
- **y** (*None or 1x3 array of float*) – Vector that should lie on the y-axis after transformation
- **z** (*None or 1x3 array of float*) – Vector that after transformation should lie on yz-plane in direction of z-axis.

**Returns** 4x4 homogenous transformation matrix.

**Return type** 4x4 array of float

**pad\_array**(*a, n=1.0, axis=1*)

Return *a* with row of *n* appended to *axis*.

**Parameters**

- **a** (*ndarray*) – Array to pad
- **n** (*float or int, optional*) – Value to pad *a* with
- **axis** (*int, optional*) – Axis of *a* to pad with *n*.

**Returns** Padded array.

**Return type** ndarray

**project\_to\_plane**(*vec\_arr, norm*)

Project array of vectors to plane with normal *norm*.

**Parameters**

- **vec\_arr** (*Nx3 array*) – Array of N 3D vectors.
- **norm** (*1x3 array*) – Normal vector to plane.

**Returns** Array of vectors projected onto plane.

**Return type** Nx3 array

**quaternion\_to\_transform\_matrix**(*quaternion, translation=array([0., 0., 0.])*)

Convert quaternion to homogenous 4x4 transform matrix.

**Parameters**

- **quaternion** (*4x1 array of float*) – Quaternion describing rotation after translation.
- **translation** (*3x1 array of float, optional*) – Translation to be performed before rotation.

**rotate\_angles**(*angles, amount*)

Rotate angles by *amount*, keeping in 0 to 2pi range.

### Parameters

- **angles** (*1-D array of float*) – Angles in radians
- **amount** (*float*) – Amount to rotate angles by

### Returns 1-D array of float

**Return type** Rotated angles

## `transform_array` (*transform\_matrix, a*)

Pad an array with 1s, transform, and return with original dimensions.

### Parameters

- **transform\_matrix** (*4x4 array of float*) – 4x4 homogenous transformation matrix
- **a** (*Nx3 array of float*) – Array of 3-D coordinates.

### Returns Nx3 array of float

**Return type** Transformed array

## `transform_matrix_to_quaternion` (*transform\_matrix, dtype=<class 'numpy.float64'>*)

Convert homogenous 4x4 transform matrix to quaternion.

### Parameters

- **transform\_matrix** (*4x4 array of float*) – Homogenous transformation matrix.
- **dtype** (*numpy dtype, optional*) – Datatype for returned quaternion.

## `unpad_array` (*a, axis=1*)

Return *a* with row removed along *axis*.

### Parameters

- **a** (*ndarray*) – Array from which to remove row
- **axis** (*int, optional*) – Axis from which to remove row

### Returns Unpadded array.

**Return type** ndarray

## e3fp.fingerprint.db module

Database for accessing and serializing fingerprints.

Author: Seth Axen E-mail: [seth.axen@gmail.com](mailto:seth.axen@gmail.com)

**class FingerprintDatabase** (*fp\_type=<class 'e3fp.fingerprint.fpint.Fingerprint'>, name=None*)  
Bases: object

Efficiently build, access, compare, and save fingerprints.

Fingerprints must have the same values of *bits* and *level*. Additionally, all fingerprints will be cast to the type of fingerprint passed to the database upon instantiation.

### Parameters

- **fp\_type** (*type, optional*) – Type of fingerprint (Fingerprint, CountFingerprint, FloatFingerprint).
- **level** (*int, optional*) – Level, or number of iterations used during fingerprinting.

- **name** (*str, optional*) – Name of database.

## Variables

- **array** (*scipy.sparse.csr\_matrix*) – Sparse matrix with dimensions N x M, where M is *bits*, and M is *fp\_num*.
- **bits** (*int*) – Number of bits (length) of fingerprints.
- **fp\_names** (*list of str*) – Names of fingerprints.
- **fp\_names\_to\_indices** (*dict*) – Map from fingerprint name to row indices of *array*.
- **fp\_num** (*int*) – Number of fingerprints in database.
- **fp\_type** (*type*) – Type of fingerprint (Fingerprint, CountFingerprint, FloatFingerprint)
- **level** (*int*) – Level, or number of iterations used during fingerprinting.
- **name** (*str*) – Name of database
- **props** (*dict*) – Dict with keys specifying names of fingerprint properties and values corresponding to array of values.

## Notes

Since most fingerprints are very sparse length-wise, *FingerprintDatabase* is implemented as a wrapper around a *scipy.sparse.csr\_matrix* for efficient memory usage. This provides easy access to underlying data for tight integration with NumPy/SciPy and machine learning packages while simultaneously providing several fingerprint-specific features.

### See also:

`e3fp.fingerprint.fprint.Fingerprint` A fingerprint that stores indices of “on” bits

## Examples

```
>>> from e3fp.fingerprint.db import FingerprintDatabase
>>> from e3fp.fingerprint.fprint import Fingerprint
>>> import numpy as np
>>> np.random.seed(2)
>>> db = FingerprintDatabase(fp_type=Fingerprint, name="TestDB")
>>> print(db)
FingerprintDatabase[name: TestDB, fp_type: Fingerprint, level: -1, bits: None, fp_
->num: 0]
>>> bvs = (np.random.uniform(size=(3, 1024)) > .9).astype(bool)
>>> fps = [Fingerprint.from_vector(bvs[i, :], name="fp" + str(i))
...     for i in range(bvs.shape[0])]
>>> db.add_fingerprints(fps)
>>> print(db)
FingerprintDatabase[name: TestDB, fp_type: Fingerprint, level: -1, bits: 1024, fp_
->num: 3]
```

The contained fingerprints may be accessed by index or name.

```
>>> db[0]
Fingerprint(indices=array([40, ..., 1012]), level=-1, bits=1024, name=fp0)
>>> db['fp2']
[Fingerprint(indices=array([0, ..., 1013]), level=-1, bits=1024, name=fp2)]
```

Alternatively, the underlying `scipy.sparse.csr_matrix` may be accessed.

```
>>> db.array
<3x1024 sparse matrix of type '<... 'numpy.bool_>'>
...with 327 stored elements in Compressed Sparse Row format>
>>> db.array.toarray()
array([[False, False, False, ..., False, False, False],
       [False, False, False, ..., False, False, False],
       [ True, False, False, ..., False, False, False]])
```

Fingerprint properties may be stored in the database.

```
>>> db.set_prop("prop", np.arange(3))
```

The database can be efficiently stored and loaded.

```
>>> db.savez("/tmp/test_db.fpz")
>>> db = FingerprintDatabase.load("/tmp/test_db.fpz")
>>> print(db)
FingerprintDatabase[name: TestDB, fp_type: Fingerprint, level: -1, bits: 1024, fp_
˓→num: 3]
```

Various comparison metrics in `e3fp.fingerprint.metrics` can operate efficiently directly on databases

```
>>> from e3fp.fingerprint.metrics import tanimoto, dice, cosine
>>> tanimoto(db, db)
array([[1., 0.0591133, 0.04245283],
       [0.0591133, 1., 0.0531401],
       [0.04245283, 0.0531401, 1.]])
>>> dice(db, db)
array([[1., 0.11162791, 0.08144796],
       [0.11162791, 1., 0.10091743],
       [0.08144796, 0.10091743, 1.]])
>>> cosine(db, db)
array([[1., 0.11163878, 0.08145547],
       [0.11163878, 1., 0.10095568],
       [0.08145547, 0.10095568, 1.]])
```

### `add_fingerprints(fprints)`

Add fingerprints to database.

**Parameters** `fprints` (*iterable of Fingerprint*) – Fingerprints to add to database

### `as_type(fp_type, copy=False)`

Get database with fingerprint type `fp_type`.

**Parameters**

- `fp_type` (*type*) – Type of fingerprint (Fingerprint, CountFingerprint, FloatFingerprint)
- `copy` (*bool, optional*) – Force copy of database. If False, if database is already of requested type, no copy is made.

**Returns** Database coerced to fingerprint type of `fp_type`.

**Return type** *FingerprintDatabase*

**property bits**

**fold**(*bits*, *fp\_type=None*, *name=None*)

Get copy of database folded to specified bit length.

**Parameters**

- **bits** (*int*) – Number of bits to which to fold database.
- **fp\_type** (*type or None, optional*) – Type of fingerprint (Fingerprint, CountFingerprint, FloatFingerprint). Defaults to same type.
- **name** (*str, optional*) – Name of database

**Returns** Database folded to specified length.

**Return type** *FingerprintDatabase*

**Raises BitsValueError** – If *bits* is greater than the length of the database or database cannot be evenly folded to length *bits*.

**property fp\_num**

**classmethod from\_array**(*array*, *fp\_names*, *fp\_type=None*, *level=-1*, *name=None*, *props={}*)

Instantiate from array.

**Parameters**

- **array** (*numpy.ndarray or scipy.sparse.csr\_matrix*) – Sparse matrix with dimensions *N* x *M*, where *M* is the number of bits in the fingerprints.
- **fp\_names** (*list of str*) – *N* names of fingerprints in *array*.
- **fp\_type** (*type, optional*) – Type of fingerprint (Fingerprint, CountFingerprint, FloatFingerprint).
- **level** (*int, optional*) – Level, or number of iterations used during fingerprinting.
- **name** (*str or None, optional*) – Name of database.
- **props** (*dict, optional*) – Dict with keys specifying names of fingerprint properties and values corresponding to length *N* array of values.

**Returns** Database containing fingerprints in *array*.

**Return type** *FingerprintDatabase*

**get\_density**(*index=None*)

Get percentage of fingerprints with ‘on’ bit at position.

**Parameters index** (*int or None, optional*) – Index to bit for which to return positional density.  
If None, density for whole database is returned.

**Returns** Density of ‘on’ position in database

**Return type** *float*

**get\_prop**(*key*)

Get property.

**Raises KeyError** – If *key* not in *props*.

**get\_subset**(*fp\_names*, *name=None*)

Get database with subset of fingerprints.

**Parameters**

- **fp\_names** (*list of str*) – List of fingerprint names to include in new db.
- **name** (*str, optional*) – Name of database

**classmethod load** (*fn*)

Load database from file.

The extension is used to determine how database was serialized (`save` vs `savez`).

**Parameters** **fn** (*str*) – Filename

**Returns** Database

**Return type** *FingerprintDatabase*

**save** (\*\**kwargs*)

---

**Note:** Deprecated in e3fp 1.2. `save` will be removed in e3fp 1.3. Use `savez` instead.

---

Save database to file.

**fn** [*str, optional*] Filename or basename if extension does not include ‘.fps’

**savetxt** (*fn, with\_names=True*)

Save bitstring representation to text file.

Only implemented for *fp\_type* of *Fingerprint*. This should not be attempted for large numbers of bits.

**Parameters**

- **fn** (*str or filehandle*) – Out file. Extension is automatically parsed to determine whether compression is used.
- **with\_names** (*bool, optional*) – Include name of fingerprint in same row after bitstring.

**Raises**

- **E3FPInvalidFingerprintError** – If *fp\_type* is not *Fingerprint*.
- **E3FPEfficiencyWarning** – If *bits* is over  $2^{14} = 16384$ .

**savez** (*fn='fingerprints.fpz'*)

Save database to file.

Database is serialized using `numpy.savez_compressed`.

**Parameters** **fn** (*str, optional*) – Filename or basename if extension is not ‘.fpz’

**set\_prop** (*key, vals, check\_length=True*)

Set values of property for fingerprints.

**Parameters**

- **key** (*str*) – Name of property
- **vals** (*array\_like*) – Values of property.
- **check\_length** (*bool, optional*) – Check to ensure number of properties match number of fingerprints already in database. This should only be set to False for temporary iterative updating.

**update\_names\_map** (*new\_names=None, offset=0*)

Update map of fingerprint names to row indices of *self.array*.

**Parameters**

- **new\_names** (*iterable of str, optional*) – Names to add to map. If None, map is completely rebuilt.
- **offset** (*int, optional*) – Number of rows before new rows.

**update\_props** (*props\_dict, append=False, check\_length=True*)

Set multiple properties at once.

#### Parameters

- **props\_dict** (*dict*) – Dict of properties. Values must be array-like of length *fp\_num*.
- **append** (*bool, optional*) – Append values to those already in database. By default, properties are overwritten if already present.
- **check\_length** (*bool, optional*) – Check to ensure number of properties match number of fingerprints already in database. This should only be set to False for temporary iterative updating.

**concat** (*dbs*)

Efficiently concatenate *FingerprintDatabase* objects.

The databases must be of the same type with the same number of bits, level, and property names.

**Parameters** **dbs** (*iterable of FingerprintDatabase*) – Fingerprint databases

**Returns** Database with all fingerprints from provided databases.

**Return type** *FingerprintDatabase*

**See also:**

*FingerprintDatabase()*

## Examples

```
>>> from e3fp.fingerprint.db import FingerprintDatabase, concat
>>> from e3fp.fingerprint.fprint import Fingerprint
>>> import numpy as np
>>> np.random.seed(2)
>>> db1 = FingerprintDatabase(fp_type=Fingerprint, name="TestDB1", level=5)
>>> db2 = FingerprintDatabase(fp_type=Fingerprint, name="TestDB2", level=5)
>>> bvs = (np.random.uniform(size=(6, 1024)) > .9).astype(bool)
>>> fps = [Fingerprint.from_vector(bvs[i, :], name="fp" + str(i), level=5)
...     for i in range(bvs.shape[0])]
>>> db1.add_fingerprints(fps[:3])
>>> db2.add_fingerprints(fps[3:])
>>> print(concat([db1, db2]))
FingerprintDatabase[name: None, fp_type: Fingerprint, level: 5, bits: 1024, fp_
˓→num: 6]
```

## e3fp.fingerprint.fprint module

Classes and methods for chemical fingerprint storage and comparison.

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```
class CountFingerprint (indices=None, counts=None, bits=4294967296, level=-1, name=None,
                        props={}, **kwargs)
```

Bases: [e3fp.fingerprint.Fingerprint](#)

A fingerprint that stores number of occurrences of each index.

### Parameters

- **indices** (*array-like of int, optional*) –  $\log_2(\text{bits})$ -bit indices in a sparse vector, corresponding to positions with counts greater than 0. If not provided, **counts** must be provided.
- **counts** (*dict, optional*) – Dict matching each index in **indices** to number of counts. All counts default to 1 if not provided.
- **bits** (*int, optional*) – Number of bits in bitvector.
- **level** (*int, optional*) – Level of fingerprint, corresponding to fingerprinting iterations.
- **name** (*str, optional*) – Name of fingerprint.
- **props** (*dict, optional*) – Custom properties of fingerprint, consisting of a string keyword and some value.

### Variables

- **bits** (*int*) – Number of bits in bitvector, length of fingerprint.
- **counts** (*dict*) – Dict matching each index in **indices** to number of counts.
- **indices** (*numpy.ndarray of int*) – Indices of fingerprint with counts greater than 0.
- **level** (*int*) – Level of fingerprint, corresponding to fingerprinting iterations.
- **mol** (*RDKit Mol*) – Mol to which fingerprint corresponds (stored in **props**).
- **name** (*str or None*) – Name of fingerprint (stored in **props**).
- **props** (*dict*) – Custom properties of fingerprint, consisting of a string keyword and some value.
- **vector\_dtype** (*numpy.dtype*) – NumPy data type associated with fingerprint values (e.g. bits)

See also:

[\*\*Fingerprint\*\*](#) A fingerprint that stores indices of “on” bits

[\*\*FloatFingerprint\*\*](#) A fingerprint that stores float counts

## Examples

```
>>> import e3fp.fingerprint.fprint as fp
>>> from e3fp.fingerprint.metrics import soergel
>>> import numpy as np
>>> np.random.seed(1)
>>> bits = 1024
>>> indices = np.random.randint(0, bits, 30)
>>> print(indices)
[ 37 235 908 72 767 905 715 645 847 960 144 129 972 583 749 508 390 281
 178 276 254 357 914 468 907 252 490 668 925 398]
>>> counts = dict(zip(indices,
...                     np.random.randint(1, 100, indices.shape[0])))
>>> print(sorted(counts.items()))
[(37, 51), (72, 88), (129, 62), ..., (925, 50), (960, 8), (972, 23)]
>>> f = fp.CountFingerprint(indices, counts=counts, bits=bits, level=0)
>>> f_folded = f.fold(bits=32)
>>> print(sorted(f_folded.counts.items()))
[(0, 8), (1, 62), (5, 113), ..., (29, 50), (30, 14), (31, 95)]
>>> print(f_folded.to_vector(sparse=False, dtype=int))
[ 8   62   0   0 113   61   58   88   97   71 228 111   2   58   10   64   0
 82   0 120   0   0   0   82   0   0   27   50   14   95]
>>> fp.Fingerprint.from_fingerprint(f_folded)
Fingerprint(indices=array([0, 1, ...]), level=0, bits=32, name=None)
>>> indices2 = np.random.randint(0, bits, 30)
>>> counts2 = dict(zip(indices2,
...                     np.random.randint(1, 100, indices.shape[0])))
>>> f_folded2 = fp.CountFingerprint.from_indices(indices2, counts=counts2,
...                                                bits=bits).fold(bits=32)
>>> print(sorted(f_folded2.counts.items()))
[(0, 93), (2, 33), (3, 106), ..., (25, 129), (26, 89), (30, 53)]
>>> print(soergel(f_folded, f_folded2))
0.17492946392...
```

### property counts

#### `fold(*args, **kwargs)`

Fold fingerprint while considering counts.

Optionally, provide a function to reduce colliding counts.

#### Parameters

- **bits** (*int, optional*) – Length of new bitvector, ideally multiple of 2.
- **method** (*{0, 1}*, *optional*) – Method to use for folding.
  - 0** partitioning (array is divided into equal sized arrays of length *bits* which are bitwise combined with *counts\_method*)
  - 1** compression (adjacent bits pairs are combined with *counts\_method* until length is *bits*)
- **linked** (*bool, optional*) – Link folded and unfolded fingerprints for easy referencing. Set to False if intending to save and want to reduce file size.
- **counts\_method** (*function, optional*) – Function for combining counts. Default is summation.

#### Returns `CountFingerprint`

**Return type** Fingerprint of folded vector

```
classmethod from_counts(counts, bits=4294967296, level=-1, **kwargs)
    Initialize from an array of indices.

    Parameters
    • counts (dict) – Dictionary mapping sparse indices to counts.
    • bits (int, optional) – Number of bits in array. Indices will be log2(bits)-bit integers.
    • level (int, optional) – Level of fingerprint, corresponding to fingerprinting iterations.
    • name (str; optional) – Name of fingerprint.
    • props (dict, optional) – Custom properties of fingerprint, consisting of a string keyword and some value.

    Returns fingerprint
    Return type CountFingerprint

classmethod from_fingerprint(fp, **kwargs)
    Initialize by copying existing fingerprint.

    Parameters
    • fp (Fingerprint) – Existing fingerprint.
    • name (str, optional) – Name of fingerprint.
    • props (dict, optional) – Custom properties of fingerprint, consisting of a string keyword and some value.

    Returns fingerprint
    Return type Fingerprint

classmethod from_indices(indices, counts=None, bits=4294967296, level=-1, **kwargs)
    Initialize from an array of indices.

    Parameters
    • indices (array_like of int, optional) – Indices in a sparse bitvector of length bits which correspond to 1.
    • counts (dict, optional) – Dictionary mapping sparse indices to counts.
    • bits (int, optional) – Number of bits in array. Indices will be log2(bits)-bit integers.
    • level (int, optional) – Level of fingerprint, corresponding to fingerprinting iterations.
    • name (str; optional) – Name of fingerprint.
    • props (dict, optional) – Custom properties of fingerprint, consisting of a string keyword and some value.

    Returns fingerprint
    Return type CountFingerprint

get_count (index)
    Return count index in fingerprint.

    Returns int
    Return type Count of index in fingerprint

mean ()
    Return mean of counts.
```

**Returns float**

**Return type** Mean

**reset**(\*args, \*\*kwargs)

Reset all values.

**std()**

Return standard deviation of fingerprint.

**Returns float**

**Return type** Standard deviation

**vector\_dtype**

alias of numpy.uint16

**class Fingerprint**(indices, bits=4294967296, level=-1, name=None, props={}, \*\*kwargs)

Bases: object

A fingerprint that stores indices of “on” bits.

#### Parameters

- **indices** (array\_like of int, optional) – log<sub>2</sub>(bits)-bit indices in a sparse bitvector of bits which correspond to 1.
- **bits** (int, optional) – Number of bits in bitvector.
- **level** (int, optional) – Level of fingerprint, corresponding to fingerprinting iterations.
- **name** (str, optional) – Name of fingerprint.
- **props** (dict, optional) – Custom properties of fingerprint, consisting of a string keyword and some value.

#### Variables

- **bits** (int) – Number of bits in bitvector, length of fingerprint.
- **counts** (dict) – Dict matching each index in indices to number of counts (1 for bits).
- **indices** (numpy.ndarray of int) – Indices of “on” bits
- **level** (int) – Level of fingerprint, corresponding to fingerprinting iterations.
- **mol** (RDKit Mol) – Mol to which fingerprint corresponds (stored in props).
- **name** (str or None) – Name of fingerprint (stored in props).
- **props** (dict) – Custom properties of fingerprint, consisting of a string keyword and some value.
- **vector\_dtype** (numpy.dtype) – NumPy data type associated with fingerprint values (e.g. bits)

#### See also:

[\*\*CountFingerprint\*\*](#) A fingerprint that stores number of occurrences of each index

[\*\*FloatFingerprint\*\*](#) A fingerprint that stores indices of “on” bits

[\*\*e3fp.fingerprint.db.FingerprintDatabase\*\*](#) Efficiently store fingerprints

## Examples

```
>>> import e3fp.fingerprint.fprint as fp
>>> from e3fp.fingerprint.metrics import tanimoto
>>> import numpy as np
>>> np.random.seed(0)
>>> bits = 1024
>>> indices = np.random.randint(0, bits, 30)
>>> print(indices)
[684 559 629 192 835 763 707 359 9 723 277 754 804 599 70 472 600 396
 314 705 486 551 87 174 600 849 677 537 845 72]
>>> f = fp.Fingerprint(indices, bits=bits, level=0)
>>> f_folded = f.fold(bits=32)
>>> print(f_folded.indices)
[ 0  1  3  4  5  6  7  8  9 12 13 14 15 17 18 19 21 23 24 25 26 27]
>>> print(f_folded.to_vector(sparse=False, dtype=int))
[1 1 0 1 1 1 1 1 0 0 1 1 1 0 1 1 0 1 0 1 1 1 1 1 0 0 0 0]
>>> print(f_folded.to_bitstring())
110111111001111011101011110000
>>> print(f_folded.to_rdkit())
<rdkit.DataStructs.cDataStructs.ExplicitBitVect object at 0x...>
>>> f_folded2 = fp.Fingerprint.from_indices(np.random.randint(0, bits, 30),
...                                             bits=bits).fold(bits=32)
>>> print(f_folded2.indices)
[ 0  1  3  5  7  9 10 14 15 16 17 18 19 20 23 24 25 29 30 31]
>>> print(tanimoto(f_folded, f_folded2))
0.5
```

**property bit\_count**

**property bits**

**clear()**

Clear temporary (and possibly large) values.

**property counts**

**property density**

**fold(bits=1024, method=0, linked=True)**

Return fingerprint for bitvector folded to size *bits*.

### Parameters

- **bits** (*int, optional*) – Length of new bitvector, ideally multiple of 2.
- **method** (*{0, 1}*, *optional*) – Method to use for folding.
  - 0** partitioning (array is divided into equal sized arrays of length *bits* which are bitwise combined with OR)
  - 1** compression (adjacent bits pairs are combined with OR until length is *bits*)
- **linked** (*bool, optional*) – Link folded and unfolded fingerprints for easy referencing. Set to False if intending to save and want to reduce file size.

### Returns Fingerprint

**Return type** Fingerprint of folded bitvector

**classmethod from\_bitstring(bitstring, level=-1, \*\*kwargs)**

Initialize from bitstring (e.g. ‘10010011’).

**Parameters**

- **bitstring** (*str*) – String of 1s and 0s.
- **level** (*int, optional*) – Level of fingerprint, corresponding to fingerprinting iterations.
- **name** (*str; optional*) – Name of fingerprint.
- **props** (*dict, optional*) – Custom properties of fingerprint, consisting of a string keyword and some value.

**Returns** `fingerprint`**Return type** `Fingerprint`**classmethod** `from_fingerprint` (*fp, \*\*kwargs*)

Initialize by copying existing fingerprint.

**Parameters** `fp` (*Fingerprint*) – Existing fingerprint.**Returns** `fingerprint`**Return type** `Fingerprint`**classmethod** `from_indices` (*indices, bits=4294967296, level=-1, \*\*kwargs*)

Initialize from an array of indices.

**Parameters**

- **indices** (*array\_like of int*) – Indices in a sparse bitvector of length `bits` which correspond to 1.
- **bits** (*int, optional*) – Number of bits in array. Indices will be  $\log_2(\text{bits})$ -bit integers.
- **level** (*int, optional*) – Level of fingerprint, corresponding to fingerprinting iterations.
- **name** (*str; optional*) – Name of fingerprint.
- **props** (*dict, optional*) – Custom properties of fingerprint, consisting of a string keyword and some value.

**Returns** `fingerprint`**Return type** `Fingerprint`**classmethod** `from_rdkit` (*rdkit\_fp, \*\*kwargs*)

Initialize from RDKit fingerprint.

If provided fingerprint is of length  $2^{32} - 1$ , assumes real fingerprint is of length  $2^{32}$ .**Parameters**

- **rdkit\_fp** (*RDKit ExplicitBitVect or SparseBitVect*) – Existing RDKit fingerprint.
- **level** (*int, optional*) – Level of fingerprint, corresponding to fingerprinting iterations.
- **name** (*str; optional*) – Name of fingerprint.
- **props** (*dict, optional*) – Custom properties of fingerprint, consisting of a string keyword and some value.

**Returns** `fingerprint`**Return type** `Fingerprint`**classmethod** `from_vector` (*vector, level=-1, \*\*kwargs*)

Initialize from vector.

**Parameters**

- **vector** (*numpy.ndarray or scipy.sparse.csr\_matrix*) – Array of bits/counts/floats
- **level** (*int, optional*) – Level of fingerprint, corresponding to fingerprinting iterations.
- **name** (*str, optional*) – Name of fingerprint.
- **props** (*dict, optional*) – Custom properties of fingerprint, consisting of a string keyword and some value.

**Returns** `fingerprint`

**Return type** `Fingerprint`

**get\_count** (*index*)

Return count index in fingerprint.

Defaults to 1 if index in *self.indices*

**Returns** `int`

**Return type** Count of bit in fingerprint

**get\_folding\_index\_map** ()

Get map of sparse indices to folded indices.

**Returns** `dict`

**Return type** Map of sparse index (keys) to corresponding folded index.

**get\_prop** (*key*)

Get property. If not set, raise KeyError.

**get\_unfolding\_index\_map** ()

Get map of sparse indices to unfolded indices.

**Returns** `dict` – indices.

**Return type** Map of sparse index (keys) to set of corresponding unfolded

**property index\_id\_map**

**property indices**

**property level**

**mean** ()

Return mean, i.e. proportion of “on” bits in fingerprint.

**Returns** `float`

**Return type** Mean

**property mol**

**property name**

**property props**

**reset** ()

Reset all values.

**set\_prop** (*key, val*)

Set property.

**std** ()

Return standard deviation of fingerprint.

**Returns** `float`

**Return type** Standard deviation

**to\_bitstring()**  
Get bitstring as string of 1s and 0s.

**Returns** str

**Return type** bitstring

**to\_bitvector(sparse=True)**  
Get full bitvector.

**Returns** numpy.ndarray or scipy.sparse.csr\_matrix of bool

**Return type** Bitvector

**to\_rdkit()**  
Convert to RDKit fingerprint.

If number of bits exceeds  $2^{31} - 1$ , fingerprint will be folded to length  $2^{31} - 1$  before conversion.

**Returns** rdkit\_fprint – Convert to bitvector used for RDKit fingerprints. If *self.bits* is less than  $10^5$ , ExplicitBitVect is used. Otherwise, SparseBitVect is used.

**Return type** RDKit ExplicitBitVect or SparseBitVect

**to\_vector(sparse=True, dtype=None)**  
Get vector of bits/counts/floats.

**Returns** Vector of bits/counts/floats

**Return type** numpy.ndarray or scipy.sparse.csr\_matrix

**unfold()**  
Return unfolded parent fingerprint for bitvector.

**Returns** Fingerprint – None.

**Return type** Fingerprint of unfolded bitvector. If None, return

**update\_props(props\_dict)**  
Set multiple properties at once.

**vector\_dtype**  
alias of numpy.bool\_

**class FloatFingerprint(indices=None, counts=None, bits=4294967296, level=-1, name=None, props={}, \*\*kwargs)**  
Bases: e3fp.fingerprint.CountFingerprint

A Fingerprint that stores float counts.

Nearly identical to *CountFingerprint*. Mainly a naming convention, but count values are stored as floats.

**See also:**

**Fingerprint** A fingerprint that stores indices of “on” bits

**CountFingerprint** A fingerprint that stores number of occurrences of each index

**property counts**

**vector\_dtype**  
alias of numpy.float64

**add** (*fprints, weights=None*)

Add fingerprints by count to new *CountFingerprint*.

If any of the fingerprints are *FloatFingerprint*, resulting fingerprint is likewise a *FloatFingerprint*. Otherwise, resulting fingerprint is *CountFingerprint*.

**Parameters**

- **fprints** (*iterable of Fingerprint*) – Fingerprints to be added by count.
- **weights** (*iterable of float*) – Weights for weighted sum. Results in *FloatFingerprint* output.

**Returns** Fingerprint with counts as sum of counts in *fprints*.

**Return type** *CountFingerprint* or *FloatFingerprint*

**See also:**

*mean ()*

**coerce\_to\_valid\_dtype** (*dtype*)

Coerce provided NumPy data type to closest fingerprint data type.

If provided *dtype* cannot be read, default corresponding to bit *Fingerprint* is returned.

**Parameters** **dtype** (*numpy.dtype or str*) – Input NumPy data type.

**Returns** Output NumPy data type.

**Return type** *numpy.dtype*

**diff\_counts\_dict** (*fp1, fp2, only\_positive=False*)

Given two fingerprints, returns difference of their counts dict.

**Parameters**

- **fp1, fp2** (*Fingerprint*) – *Fingerprint* objects, *fp2* subtracted from *fp1*.
- **only\_positive** (*bool, optional*) – Return only positive counts, negative being thresholded to 0.

**Returns** **counts\_diff** – Count indices in either *fp1* or *fp2* with value as diff of counts.

**Return type** *dict*

**See also:**

*sum\_counts\_dict ()*

**dtype\_from\_fptype** (*fp\_type*)

Get NumPy data type from fingerprint type.

**Parameters** **fp\_type** (*class or Fingerprint*) – Class of fingerprint

**Returns** NumPy data type

**Return type** *numpy.dtype*

**fptype\_from\_dtype** (*dtype*)

Get corresponding fingerprint type from NumPy data type.

**Parameters** **dtype** (*numpy.dtype or str*) – NumPy data type.

**Returns** **class** – Class of fingerprint

**Return type** {*Fingerprint*, *CountFingerprint*, *FloatFingerprint*}

**load**(*f, update\_structure=True*)Load [Fingerprint](#) object from file.**Parameters**

- **f** (*str or File*) – File name or file-like object to load file from.
- **update\_structure** (*bool, optional*) – Attempt to update the class structure by initializing a new, shiny fingerprint from each fingerprint in the file. Useful for guaranteeing that old, dusty fingerprints are always upgradeable.

**Returns** [Fingerprint](#)**Return type** Pickled fingerprint.**See also:**[loadz\(\)](#), [save\(\)](#)**loadz**(*f, update\_structure=True*)Load [Fingerprint](#) objects from file.**Parameters**

- **f** (*str or File*) – File name or file-like object to load file from.
- **update\_structure** (*bool, optional*) – Attempt to update the class structure by initializing a new, shiny fingerprint from each fingerprint in the file. Useful for guaranteeing that old, dusty fingerprints are always upgradeable. If this doesn't work, falls back to the original saved fingerprint.

**Returns** list of [Fingerprint](#)**Return type** Fingerprints in pickle.**See also:**[load\(\)](#), [savez\(\)](#)**mean**(*fprints, weights=None*)Average fingerprints to generate [FloatFingerprint](#).**Parameters**

- **fprints** (*iterable of Fingerprint*) – Fingerprints to be added by count.
- **weights** (*array\_like of float, optional*) – Weights for weighted mean. Weights are normalized to a sum of 1.

**Returns** [FloatFingerprint](#) – *fprints*.**Return type** Fingerprint with float counts as average of counts in**save**(*f, fp, \*\*kwargs*)Save [Fingerprint](#) object to file.**Parameters**

- **f** (*str or File*) – filename *str* or file-like object to save file to
- **fp** ([Fingerprint](#)) – Fingerprint to save to file
- **protocol** (*{0, 1, 2, None}, optional*) – Pickle protocol to use. If None, highest available protocol is used. This will not affect fingerprint loading.

**Returns** bool**Return type** Success or fail

See also:

`savez()`, `load()`

`savez(f, *fps, **kwargs)`

Save multiple `Fingerprint` objects to file.

#### Parameters

- `f (str or File)` – filename `str` or file-like object to save file to
- `fps (list of Fingerprint)` – List of Fingerprints to save to file
- `protocol ({0, 1, 2, None}, optional)` – Pickle protocol to use. If `None`, highest available protocol is used. This will not affect fingerprint loading.

#### Returns `bool`

**Return type** Success or fail

See also:

`save()`, `loadz()`

`sum_counts_dict(*fps, **kwargs)`

Given fingerprints, return sum of their counts dicts.

If an optional `weights` iterable of the same length as `fps` is provided, the weighted sum is returned.

#### Parameters

- `*fps` – One or more `Fingerprint` objects
- `weights (iterable of float, optional)` – Weights for weighted mean. Weights are normalized to a sum of 1.

**Returns** `dict` – as sum of counts.

**Return type** Dict of non-zero count indices in any of the `fps` with value

See also:

`diff_counts_dict()`

## e3fp.fingerprint.fprinter module

Tools for generating E3FP fingerprints.

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```
class Fingerprinter(bits=4294967296,      level=5,      radius_multiplier=1.718,      stereo=True,
                      counts=False,     include_disconnected=True,    rdkit_invariants=False,   ex-
                      clude_floating=True, remove_duplicate_substructs=True)
```

Bases: `object`

E3FP fingerprint generator.

#### Parameters

- `bits (int or None, optional)` – Maximum number of bits to which to fold returned fingerprint. Multiple of 2 is strongly recommended.
- `level (int or None, optional)` – Maximum number of iterations for fingerprint generation. If `None` or `-1`, run until no new substructures are identified. Because this could produce a different final level number for each conformer, it is recommended to manually specify a level.

- **radius\_multiplier** (*float, optional*) – Multiple by which to increase shell size. At iteration 0, shell radius is  $0^*\text{radius\_multiplier}$ , at iteration 2, radius is  $2^*\text{radius\_multiplier}$ , etc.
- **counts** (*bool, optional*) – Instead of simple bit-based Fingerprint object, generate CountFingerprint that tracks number of times each bit appears in a fingerprint.
- **stereo** (*bool, optional*) – Differentiate based on stereography. Resulting fingerprints are not comparable to non-stereo fingerprints.
- **remove\_duplicate\_substructs** (*bool, optional*) – If a substructure arises that corresponds to an identifier already in the fingerprint, then the identifier for the duplicate substructure is not added to fingerprint.
- **include\_disconnected** (*bool, optional*) – Include disconnected atoms from hashes and substructure. E3FP's advantage over ECFP relies on disconnected atoms, so the option to turn this off is present only for testing/comparison.
- **rdkit\_invariants** (*bool, optional*) – Use the atom invariants used by RDKit for its Morgan fingerprint.
- **exclude\_floating** (*bool, optional*) – Mask atoms with no bonds (usually floating ions) from the fingerprint. These are often placed arbitrarily and can confound the fingerprint.

## Variables

- **current\_level** (*int*) – The maximum level/iteration to which the fingerprinter has been run on the current conformer.
- **level\_shells** (*dict*) – Dict matching level to set of all shells accepted at that level.

**property current\_level**

**get\_fingerprint\_at\_level** (*level=-1, bits=None, exact=False, atom\_mask={}*)

Get the fingerprint at the specified level.

## Parameters

- **level** (*int or None, optional*) – Level/iteration
- **bits** (*int or None, optional*) – Return fingerprints folded to this number of bits. If unspecified, defaults to bits set when instantiated.
- **exact** (*bool, optional*) – Exact level
- **atom\_mask** (*int or set of int, optional*) – Don't return shells whose substructures contain these atoms.

## Returns Fingerprint

**Return type** Fingerprint at level

**get\_shells\_at\_level** (*level=-1, exact=False, atom\_mask={}*)

Get set of shells at the specified level.

## Parameters

- **level** (*int or None, optional*) – Level/iteration
- **exact** (*bool, optional*) – Exact level
- **atom\_mask** (*int or set of int, optional*) – Don't return shells whose substructures contain these atoms.

## Returns set of Shell

**Return type** Shells at level

**initialize\_conformer**(*conf*)

Retrieve atom coordinates and instantiate shells generator.

**Parameters** *conf* (*RDKit Conformer*) – Conformer to fingerprint

**initialize\_identifiers**()

Set initial identifiers for atoms.

**initialize\_mol**(*mol*)

Set general properties of *mol* that apply to all its conformers.

**Parameters** *mol* (*RDKit Mol*) – Input molecule *Mol* object.

**next**()

Run next iteration of fingerprinting.

**reset**()

Clear all variables associated with the last run.

**reset\_conf**()

Clear only conformer-specific variables.

**reset\_mol**()

Clear all variables associated with the molecule.

**run**(*conf=None*, *mol=None*, *return\_substruct=False*)

Generate fingerprint from provided conformer or mol and conf id.

**Parameters**

- **conf** (*RDKit Conformer or int, optional*) – Input conformer or conformer in *mol*.
- **mol** (*RDKit Mol, optional*) – Input molecule object, with at least one conformer. If *conf* not specified, first conformer is used.
- **return\_substruct** (*bool, optional*) – Return dict mapping substructure to fingerprint indices. Keys are indices, values are list of substructures, represented as a tuple of atom indices where the first index is the central atom and the remaining indices (within the sphere) are sorted.

**substructs\_to\_pdb**(*level=None*, *bits=None*, *out\_dir='substructs'*, *reorient=True*, *exact=False*)

Save all accepted substructs from current level to PDB.

**Parameters**

- **level** (*int or None, optional*) – Level of fingerprinting/number of iterations
- **bits** (*int or None, optional*) – Folding level of identifiers
- **out\_dir** (*str, optional*) – Directory to which to save PDB files.
- **reorient** (*bool, optional*) – Reorient substructure to match stereo quadrants.

**class ShellsGenerator**(*conf*, *atoms*, *radius\_multiplier=0.5*, *include\_disconnected=True*, *atom\_coords=None*, *bound\_atoms\_dict=None*)

Bases: *object*

Generate nested *Shell* objects from molecule upon request.

**back**()

Back up one iteration.

**get\_match\_atoms**(*rad*)

Get atoms within shell at radius *rad*.

**Parameters** *rad* (*float*) – Radius of shell.

**Returns** `dict` – shell

**Return type** Dict matching atom id to set of ids for other atoms within

**get\_shells\_at\_level** (`level`)

Get *dict* of atom shells at specified level/iteration.

If not run to *level*, raises *IndexError*.

**Parameters** `level (int)` – Level/iteration from which to retrieve shells *dict*.

**Returns** `dict`

**Return type** Dict matching atom ids to that atom's `Shell` at that level.

**next** ()

Get next iteration's *dict* of atom shells.

**atom\_tuples\_from\_shell** (`shell, atom_coords, connectivity, stereo`)

Generate sorted atom tuples for neighboring atoms.

**Parameters**

- `shell (Shell)` – Shell for which to build atom tuples
- `atom_coords (dict)` – Dict matching atom ids to coords.
- `connectivity (dict)` – Dict matching atom id pair tuples to their bond order (5 for unbound).
- `stereo (bool)` – Add stereo indicators to tuples

**bound\_atoms\_from\_mol** (`mol, atoms`)

Build *dict* matching atom id to ids of bounded atoms.

Bound atoms not in *atoms* are ignored.

**Parameters**

- `mol (RDKit Mol)` – Input mol
- `atoms (list of int)` – List of atom IDs

**Returns** `dict`

**Return type** Dict matching atom id to set of bound atom ids.

**coords\_from\_atoms** (`atoms, conf`)

Build *dict* matching atom id to coordinates.

**Parameters**

- `atoms (list of int)` – Atom ids
- `conf (RDKit Conformer)` – Conformer from which to fetch coordinates

**Returns** `dict`

**Return type** Dict matching atom id to 1-D array of coordinates.

**get\_first\_unique\_tuple\_inds** (`tuples_list, num_ret, ignore=[], assume_sorted=True`)

Return indices of first *num\_ret* unique tuples in a list.

Only first 2 values of each tuple are considered.

**Parameters**

- `tuples_list (list of tuple)` – List of tuples. Only first two unique values are considered.
- `num_ret (int)` – Maximum number of first unique tuples to return.

- **ignore** (*list, optional*) – Indices for tuples not be considered as unique.
- **assume\_sorted** (*bool, optional*) – If True, assume list is already sorted by tuples.

**Returns** tuple of int – unique tuples in list.

**Return type** List of at most *num\_ret* ints indicating index of

**hash\_int64\_array** (*array, seed=0*)

Hash an int64 array into a 32-bit integer.

#### Parameters

- **array** (*ndarray of int64*) – Numpy array containing integers
- **seed** (*any, optional*) – Seed for MurmurHash3.

**Returns** int

**Return type** 32-bit integer

**identifier\_from\_shell** (*shell, atom\_coords, connectivity, level, stereo*)

Determine new identifier for a shell at a specific level.

#### Parameters

- **shell** (*Shell*) – Shell for which to determine identifier
- **atom\_coords** (*dict*) – Dict matching atom ids to coords.
- **connectivity** (*dict*) – Dict matching atom id pair tuples to their bond order (5 for unbound).
- **level** (*int*) – Level/iteration
- **stereo** (*bool*) – Add stereo indicators

**identifiers\_from\_invariants** (*mol, atoms, rdkit\_invariants=False*)

Initialize ids according to Daylight invariants.

#### Parameters

- **mol** (*RDKit Mol*) – Input molecule
- **atoms** (*list of int*) – IDs for atoms in mol for which to generate identifiers.
- **rdkit\_invariants** (*bool, optional*) – Use the atom invariants used by RDKit for its Morgan fingerprint.

**Returns** ndarray of int64

**Return type** initial identifiers for atoms

**invariants\_from\_atom** (*atom*)

Get seven invariants from atom.

Invariants used are the six Daylight invariants, plus an indicator of whether the atom is in a ring, as detailed in [1].

## References

1. D Rogers, M Hahn. J. Chem. Inf. Model., 2010, 50 (5), pp 742-754 <https://doi.org/10.1021/ci100050t>

**Parameters** `atom` (*RDKit Atom*) – Input atom

**Returns** 1-D array if int64

**Return type** Array of 7 invariants

**pick\_y** (`atom_tuples`, `cent_coords`, `y_precision=0.1`)

Pick a y-coordinate from atom tuples or mean coordinate.

**Parameters**

- `atom_tuples` (*list of tuple*) – Sorted list of atom tuples
- `cent_coords` (*Nx3 array of float*) – Coordinates of atoms with center atom at origin.
- `y_precision` (*str, optional*) – For mean to be chosen for y-coordinate, it must be at least this distance from the origin. Useful when atoms are symmetrical around the center atom where a slight shift in any atom results in a very different y.

**Returns**

- 1x3 array of float or None (*y-coordinate*)
- int or None (*index to y-atom, if y was chosen from the atoms.*)

**pick\_z** (`connectivity`, `identifiers`, `cent_coords`, `y`, `long_angle`, `z_precision=0.01`)

Pick a z-coordinate orthogonal to y.

**Parameters**

- `connectivity` (*dict*) – Dict matching atom id pair tuples to their bond order (5 for unbound).
- `identifiers` (*iterable of int*) – Atom identifiers
- `cent_coords` (*Nx3 array of float*) – Coordinates of atoms with center atom at origin.
- `y` (*1x3 array of float*) – y-coordinate
- `long_angle` (*Nx1 array of float*) – Absolute angle of atoms from orthogonal to y.
- `z_precision` (*str, optional*) – Minimum difference in `long_angle` between two potential z-atoms. Used as a tie breaker to prevent small shift in one atom resulting in very different z.

**Returns** 1x3 array of float or None

**Return type** z-coordinate

**quad\_indicators\_from\_coords** (`cent_coords`, `y`, `y_ind`, `z`, `long_sign`)

Create angle indicators for four quadrants in each hemisphere.

**Parameters**

- `cent_coords` (*Nx3 array of float*) – Array of centered coordinates.
- `y` (*1-D array of float*) – Vector lying along y-axis.
- `y_ind` (*int*) – Index of `cent_coords` corresponding to y.
- `z` (*1-D array of float*) – Vector lying along z-axis
- `long_sign` (*Nx1 array of int*) – Array of signs of vectors in `cent_coords` indicating whether they are above (+1) or below (-1) the xz-plane.

**Returns** Nx1 array of int – indicators are 2, 3, 4, 5 for vectors above the xz-plane and -2, -3, -4, -5 for vectors below the xz-plane.

**Return type** Quadrant indicators. Clockwise from  $z$  around  $y$ ,

#### `rdkit_invariants_from_atom(atom)`

Get the 6 atom invariants RDKit uses for its Morgan fingerprints.

**Parameters** atom (RDKit Atom) – Input atom

**Returns** 1-D array of int64

**Return type** Array of 6 invariants

#### `signed_to_unsigned_int(a, bits=4294967296)`

Convert  $int$  between +/-bits to an int between 0 and  $bits$ .

**Parameters**

- a (int or ndarray of int) – Integer
- bits (int, optional) – Maximum size of int. E.g. 32-bit is  $2^{32}$ .

**Returns** int

**Return type** unsigned integer

#### `stereo_indicators_from_shell(shell, atom_tuples, atom_coords_dict, add_transform_to_shell=True)`

Get list of int indicating location of atoms on unit sphere.

**Parameters**

- shell (Shell) – Shell for which to get stereo indicators.
- atom\_tuples (list of tuple) – List of atom tuples.
- atom\_coords\_dict (dict) – Dict matching atom ids to coords.
- add\_transform\_to\_shell (bool, optional) – Calculate transformation matrix to align coordinates to unit sphere, and add to shell.

**Returns** list of int

**Return type** stereo indicators for atoms in *atom\_tuples*.

## e3fp.fingerprint.generate module

Generate E3FP fingerprints.

Author: Seth Axen E-mail: [seth.axen@gmail.com](mailto:seth.axen@gmail.com)

#### `fprints_dict_from_mol(mol, bits=4294967296, level=5, radius_multiplier=1.718, first=3, counts=False, stereo=True, include_disconnected=True, rdkit_invariants=False, exclude_floating=True, remove_duplicate_substructs=True, out_dir_base=None, out_ext='fp.bz2', save=False, all_iters=False, overwrite=False)`

Build a E3FP fingerprint from a mol with at least one conformer.

**Parameters**

- mol (RDKit Mol) – Input molecule with one or more conformers to be fingerprinted.
- bits (int) – Set number of bits for final folded fingerprint.

- **level** (*int, optional*) – Level/maximum number of iterations of E3FP. If -1 is provided, it runs until termination, and *all\_iters* is set to False.
  - **radius\_multiplier** (*float, optional*) – Radius multiplier for spherical shells.
  - **first** (*int, optional*) – First *N* number of conformers from file to fingerprint. If -1, all are fingerprinted.
  - **counts** (*bool, optional*) – Instead of bit-based fingerprints. Otherwise, generate count-based fingerprints.
  - **stereo** (*bool, optional*) – Incorporate stereochemistry in fingerprint.
  - **remove\_duplicate\_substructs** (*bool, optional*) – If a substructure arises that corresponds to an identifier already in the fingerprint, then the identifier for the duplicate substructure is not added to fingerprint.
  - **include\_disconnected** (*bool, optional*) – Include disconnected atoms when hashing and for stereo calculations. Turn off purely for testing purposes, to make E3FP more like ECFP.
  - **rdkit\_invariants** (*bool, optional*) – Use the atom invariants used by RDKit for its Morgan fingerprint.
  - **exclude\_floating** (*bool, optional*) – Mask atoms with no bonds (usually floating ions) from the fingerprint. These are often placed arbitrarily and can confound the fingerprint.
  - **out\_dir\_base** (*str, optional*) – Basename of out directory to save fingerprints. Iteration number is appended.
  - **out\_ext** (*str, optional*) – Extension on fingerprint pickles, used to determine compression level.
  - **save** (*bool, optional*) – Save fingerprints to directory.
  - **all\_iters** (*bool, optional*) – Save fingerprints from all iterations to file(s).
  - **overwrite** (*bool, optional*) – Overwrite pre-existing file.
- Deleted Parameters**
- \_\_\_\_\_
  - **sdf\_file** (*str*) – SDF file path.

`fprints_dict_from_sdf(sdf_file, **kwargs)`

Build fingerprints dict for conformers encoded in an SDF file.

See [`fprints\_dict\_from\_mol`](#) for description of arguments.

`run(sdf_files, bits=4294967296, first=3, level=5, radius_multiplier=1.718, counts=False, stereo=True, include_disconnected=True, rdkit_invariants=False, exclude_floating=True, remove_duplicate_substructs=True, params=None, out_dir_base=None, out_ext='.fp.bz2', db_file=None, overwrite=False, all_iters=False, log=None, num_proc=None, parallel_mode=None, verbose=False)`  
Generate E3FP fingerprints from SDF files.

## e3fp.fingerprint.structs module

Class for defining 3D atom environments.

Author: Seth Axen E-mail: [seth.axen@gmail.com](mailto:seth.axen@gmail.com)

### **exception FormatError**

Bases: `Exception`

### **class Shell (center\_atom, shells={}, radius=None, last\_shell=None, identifier=None)**

Bases: `object`

A container for other Shells centered on an atom.

Shells represent all atoms explicitly within a container. Atoms are represented by their ids. If atoms are provided instead of shells, they are converted to single-atom shells. A Substruct is generated from a Shell on the fly by recursion through member shells. An optional identifier may be set.

#### **property atoms**

Get all atoms explicitly within the shell.

#### **property center\_atom**

#### **classmethod from\_substruct (substruct)**

Create shell with one shell for each atom in the substruct.

#### **property shells**

#### **property substruct**

Get substruct with all atoms implicitly within the shell.

### **class Substruct (center\_atom=None, atoms={})**

Bases: `object`

A container for atoms optionally centered on an atom.

A Substruct represents all atoms implicitly within a Shell. Two Substructs are equal if they contain the same atoms.

#### **property atoms**

#### **property center\_atom**

#### **classmethod from\_shell (shell)**

### **shell\_to\_pdb (mol, shell, atom\_coords, bound\_atoms\_dict, out\_file=None, reorient=True)**

Append substructure within shell to PDB.

#### **Parameters**

- **mol** (*RDKit Mol*) – Input mol
- **shell** (*Shell*) – A shell
- **atom\_coords** (*dict*) – Dict matching atom id to coordinates.
- **bound\_atoms\_dict** (*dict*) – Dict matching atom id to id of bound atoms.
- **out\_file** (*str or None, optional*) – File to which to append coordinates.
- **reorient** (*bool, optional*) – Use the transformation matrix in the shell to align by the stereo quadrants. If no transformation matrix present, centers the center atom.

#### **Returns list of str**

**Return type** list of PDB file lines, if *out\_file* not specified

## e3fp.fingerprint.util module

Utility methods and class for fingerprinting-related functions.

Author: Seth Axen E-mail: [seth.axen@gmail.com](mailto:seth.axen@gmail.com)

### **exception E3FPBitsValueError**

Bases: `e3fp.util.E3FPError`, `ValueError`

Bits value is invalid.

### **exception E3FPCountsError**

Bases: `e3fp.util.E3FPError`, `ValueError`

Index in counts is invalid.

### **exception E3FPInvalidFingerprintError**

Bases: `e3fp.util.E3FPError`, `TypeError`

Fingerprint is incorrectly formatted.

### **exception E3FPMolError**

Bases: `e3fp.util.E3FPError`, `TypeError`

Mol is of incorrect type.

### **exception E3FPOptionError**

Bases: `e3fp.util.E3FPError`, `ValueError`

Option provided is invalid.

## Module contents

### Submodules

## e3fp.pipeline module

Functions for various pipeline use cases.

Author: Seth Axen E-mail: [seth.axen@gmail.com](mailto:seth.axen@gmail.com)

### **confs\_from\_smiles (smiles, name, confgen\_params={}, save=False)**

Generate conformations of molecule from SMILES string.

### **fprints\_from\_fprints\_dict (fprints\_dict, level=-1)**

Get fingerprint at *level* from dict of level to fingerprint.

### **fprints\_from\_mol (mol, fprint\_params={}, save=False)**

Generate fingerprints for all *first* conformers in mol.

### **fprints\_from\_sdf (sdf\_file, fprint\_params={}, save=False)**

Generate fingerprints from conformers in an SDF file.

### **fprints\_from\_smiles (smiles, name, confgen\_params={}, fprint\_params={}, save=False)**

Generate conformers and fingerprints from a SMILES string.

### **params\_to\_dicts (params)**

Get params dicts for pipeline functions from INI format params file.

### **sdf\_from\_smiles (smiles, name, confgen\_params={}, out\_file=None, out\_ext='.sdf.bz2')**

Generate conformations from SMILES string and save to SDF file.

## e3fp.util module

Utility classes/methods.

Author: Seth Axen E-mail: [seth.axen@gmail.com](mailto:seth.axen@gmail.com)

### **exception E3FPDeprecationWarning**

Bases: `e3fp.util.E3FPWarning, DeprecationWarning`

A warning class for a deprecated method or class.

### **exception E3FPEfficiencyWarning**

Bases: `e3fp.util.E3FPWarning, RuntimeWarning`

A warning class for a potentially inefficient process.

### **exception E3FPError**

Bases: `Exception`

Base class for E3FP-specific errors.

This class is provided for future E3FP-specific functionality.

### **exception E3FPWarning**

Bases: `Warning`

Base E3FP warning class.

Unlike normal warnings, these are by default always set to on.

### **class deprecated(deprecated\_version, remove\_version=None, msg=None)**

Bases: `object`

Decorator to mark a function as deprecated.

Issue a deprecation warning when a function is called, and update the documentation. A deprecation version must be provided.

## Examples

```
>>> from e3fp.util import deprecated
>>> @deprecated("1.1", remove_version="1.3",
...             msg="Function no longer needed")
...     def my_function():
...         pass
```

## Notes

Adapted from <https://wiki.python.org/moin/PythonDecoratorLibrary>

### **deprecate\_function(f)**

Return the decorated function.

### **update\_docstring(obj)**

Add deprecation note to docstring.

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