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# **openforcefield Documentation**

*Release 0.4.0*

**Open Force Field Consortium**

**Jun 11, 2019**



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A modern, extensible library for molecular mechanics force field science from the Open Force Field Initiative



## USER GUIDE

### 1.1 Installation

#### 1.1.1 Installing via *conda*

The simplest way to install the Open Forcefield Toolkit is via the [conda](#) package manager. Packages are provided on the [omnia Anaconda Cloud channel](#) for Linux, OS X, and Win platforms. The [openforcefield Anaconda Cloud](#) page has useful instructions and [download statistics](#).

If you are using the [anaconda](#) scientific Python distribution, you already have the conda package manager installed. If not, the quickest way to get started is to install the [miniconda](#) distribution, a lightweight minimal installation of Anaconda Python.

On linux, you can install the Python 3 version into \$HOME/miniconda3 with (on bash systems):

```
$ wget https://repo.continuum.io/miniconda/Miniconda3-latest-Linux-x86_64.sh  
$ bash ./Miniconda3-latest-Linux-x86_64.sh -b -p $HOME/miniconda3  
$ source ~/miniconda3/etc/profile.d/conda.sh  
$ conda activate base
```

On osx, you want to use the osx binary

```
$ curl https://repo.continuum.io/miniconda/Miniconda3-latest-MacOSX-x86_64.sh -o  
$ bash ./Miniconda3-latest-MacOSX-x86_64.sh -b -p $HOME/miniconda3  
$ source ~/miniconda3/etc/profile.d/conda.sh  
$ conda activate base
```

You may want to add the new source `~/miniconda3/etc/profile.d/conda.sh` line to your `~/.bashrc` file to ensure Anaconda Python can enabled in subsequent terminal sessions. `conda activate base` will need to be run in each subsequent terminal session to return to the environment where the toolkit will be installed.

Note that `openforcefield` will be installed into this local Python installation, so that you will not need to worry about disrupting existing Python installations.

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**Note:** Installation via the `conda` package manager is the preferred method since all dependencies are automatically fetched and installed for you.

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### 1.1.2 Required dependencies

The openforcefield toolkit makes use of the [Omnia](#) and [Conda Forge](#) free and open source community package repositories:

```
$ conda config --add channels omnia --add channels conda-forge  
$ conda update --all
```

This only needs to be done once.

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**Note:** If automation is required, provide the `--yes` argument to `conda update` and `conda install` commands. More information on the `conda` command-line API can be found in the [conda](#) online documentation.

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### Release build

You can install the latest stable release build of openforcefield via the `conda` package with

```
$ conda config --add channels omnia --add channels conda-forge  
$ conda install openforcefield
```

This version is recommended for all users not actively developing new forcefield parameterization algorithms.

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**Note:** The `conda` package manager will install dependencies from binary packages automatically, including difficult-to-install packages such as OpenMM, numpy, and scipy. This is really the easiest way to get started.

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### Upgrading your installation

To update an earlier `conda` installation of openforcefield to the latest release version, you can use `conda update`:

```
$ conda update openforcefield
```

### Optional dependencies

This toolkit can optionally make use of the [OpenEye toolkit](#) if the user has a license key installed. Academic laboratories intending to release results into the public domain can [obtain a free license key](#), while other users (including academics) intending to use the software for purposes of generating protected intellectual property must [pay to obtain a license](#).

To install the OpenEye toolkits (provided you have a valid license file):

```
$ conda install --yes -c openeye openeye-toolkits
```

No essential openforcefield release capabilities *require* the OpenEye toolkit, but the Open Force Field developers make use of it in parameterizing new open source force fields. It is known that there are certain differences in toolkit behavior between RDKit and OpenEye when reading a small fraction of molecules, and we encourage you to report any unexpected behavior that may be caused by toolkit differences to our [issue tracker](#).

## 1.2 Release History

Releases follow the `major.minor.micro` scheme recommended by [PEP440](#), where

- major increments denote a change that may break API compatibility with previous major releases
- minor increments add features but do not break API compatibility
- micro increments represent bugfix releases or improvements in documentation

### 1.2.1 0.4.0 - Performance optimizations and support for SMIRNOFF 0.3 specification

This update contains performance enhancements that significantly reduce the time to create OpenMM systems for topologies containing many molecules via `ForceField.create_openmm_system`.

This update also introduces the [SMIRNOFF 0.3 specification](#). The spec update is the result of discussions about how to handle the evolution of data and parameter types as further functional forms are added to the SMIRNOFF spec.

We provide methods to convert SMIRNOFF 0.1 and 0.2 forcefields written with the XML serialization (`.offxml`) to the SMIRNOFF 0.3 specification. These methods are called automatically when loading a serialized SMIRNOFF data representation written in the 0.1 or 0.2 specification. This functionality allows the toolkit to continue to read files containing SMIRNOFF 0.2 spec force fields, and also implements backwards-compatibility for SMIRNOFF 0.1 spec force fields.

**Warning:** The SMIRNOFF 0.1 spec did not contain fields for several energy-determining parameters that are exposed in later SMIRNOFF specs. Thus, when reading SMIRNOFF 0.1 spec data, the toolkit must make assumptions about the values that should be added for the newly-required fields. The values that are added include 1-2, 1-3 and 1-5 scaling factors, cutoffs, and long-range treatments for nonbonded interactions. Each assumption is printed as a warning during the conversion process. Please carefully review the warning messages to ensure that the conversion is providing your desired behavior.

#### SMIRNOFF 0.3 specification updates

- The SMIRNOFF 0.3 spec introduces versioning for each individual parameter section, allowing asynchronous updates to the features of each parameter class. The top-level SMIRNOFF tag, containing information like `aromaticity_model`, `Author`, and `Date`, still has a version (currently 0.3). But, to allow for independent development of individual parameter types, each section (such as Bonds, Angles, etc) now has its own version as well (currently all 0.3).
- All units are now stored in expressions with their corresponding values. For example, distances are now stored as `1.526*angstrom`, instead of storing the unit separately in the section header.

- The current allowed value of the potential field for ProperTorsions and ImproperTorsions tags is no longer charmm, but is rather  $k*(1+\cos(\text{periodicity}*\theta-\phi))$ . It was pointed out to us that CHARMM-style torsions deviate from this formula when the periodicity of a torsion term is 0, and we do not intend to reproduce that behavior.
- SMIRNOFF spec documentation has been updated with tables of keywords and their defaults for each parameter section and parameter type. These tables will track the allowed keywords and default behavior as updated versions of individual parameter sections are released.

## Performance improvements and bugfixes

- PR #329: Performance improvements when creating systems for topologies with many atoms.
- PR #347: Fixes bug in charge assignment that occurs when charges are read from file, and reference and charge molecules have different atom orderings.

## New features

- PR #311: Several new experimental functions.
  - Adds `convert_0_2_smirnoff_to_0_3`, which takes a SMIRNOFF 0.2-spec data dict, and updates it to 0.3. This function is called automatically when creating a ForceField from a SMIRNOFF 0.2 spec OFFXML file.
  - Adds `convert_0_1_smirnoff_to_0_2`, which takes a SMIRNOFF 0.1-spec data dict, and updates it to 0.2. This function is called automatically when creating a ForceField from a SMIRNOFF 0.1 spec OFFXML file.
  - NOTE: The format of the “SMIRNOFF data dict” above is likely to change significantly in the future. Users that require a stable serialized ForceField object should use the output of `ForceField.to_string('XML')` instead.
  - Adds `ParameterHandler` and `ParameterType.add_cosmetic_attribute` and `delete_cosmetic_attribute` functions. Once created, cosmetic attributes can be accessed and modified as attributes of the underlying object (eg. `ParameterType.my_cosmetic_attrib = 'blue'`) These functions are experimental, and we are interested in feedback on how cosmetic attribute handling could be improved. (See Issue #338) Note that if a new cosmetic attribute is added to an object without using these functions, it will not be recognized by the toolkit and will not be written out during serialization.
  - Values for the top-level Author and Date tags are now kept during SMIRNOFF data I/O. If multiple data sources containing these fields are read, the values are concatenated using “AND” as a separator.

## API-breaking changes

- `ForceField.to_string` and `ForceField.to_file` have had the default value of their `discard_cosmetic_attributes` kwarg set to False.
- `ParameterHandler` and `ParameterType` constructors now expect the `version` kwarg (per the SMIRNOFF spec change above) This requirement can be skipped by providing the kwarg `skip_version_check=True`
- `ParameterHandler` and `ParameterType` functions no longer handle `x_unit` attributes in SMIRNOFF data (per the SMIRNOFF spec change above).

- The scripts in utilities/convert\_frosst are now deprecated. This functionality is important for provenance and will be migrated to the openforcefield/smirkoff99Frosst repository in the coming weeks.
- `ParameterType ._SMIRNOFF_ATTRIBS` is now `ParameterType ._REQUIRED_SPEC_ATTRIBS`, to better parallel the structure of the ParameterHandler class.
- `ParameterType ._OPTIONAL_ATTRIBS` is now `ParameterType ._OPTIONAL_SPEC_ATTRIBS`, to better parallel the structure of the ParameterHandler class.
- Added class-level dictionaries `ParameterHandler ._DEFAULT_SPEC_ATTRIBS` and `ParameterType ._DEFAULT_SPEC_ATTRIBS`.

## 1.2.2 0.3.0 - API Improvements

Several improvements and changes to public API.

### New features

- PR #292: Implement `Topology.to_openmm` and remove `ToolkitRegistry.toolkit_is_available`
- PR #322: Install directories for the lookup of OFFXML files through the entry point group `openforcefield.smirkoff_forcefield_directory`. The ForceField class doesn't search in the `data/forcefield/` folder anymore (now renamed `data/test_forcefields/`), but only in `data/`.

### API-breaking Changes

- PR #278: Standardize variable/method names
- PR #291: Remove `ForceField.load/to_smirkoff_data`, add `ForceField.to_file/string` and `ParameterHandler.add_parameters`. Change behavior of `ForceField.register_X_handler` functions.

### Bugfixes

- PR #327: Fix units in tip3p.offxml (note that this file is still not loadable by current toolkit)
- PR #325: Fix solvent box for provided test system to resolve periodic clashes.
- PR #325: Add informative message containing Hill formula when a molecule can't be matched in `Topology.from_openmm`.
- PR #325: Provide warning or error message as appropriate when a molecule is missing stereochemistry.
- PR #316: Fix formatting issues in GBSA section of SMIRNOFF spec
- PR #308: Cache molecule SMILES to improve system creation speed
- PR #306: Allow single-atom molecules with all zero coordinates to be converted to OE/RDK mols
- PR #313: Fix issue where constraints are applied twice to constrained bonds

## 1.2.3 0.2.2 - Bugfix release

This release modifies an example to show how to parameterize a solvated system, cleans up backend code, and makes several improvements to the README.

## Bugfixes

- PR #279: Cleanup of unused code/warnings in main package `__init__`
- PR #259: Update T4 Lysozyme + toluene example to show how to set up solvated systems
- PR #256 and PR #274: Add functionality to ensure that links in READMEs resolve successfully

### 1.2.4 0.2.1 - Bugfix release

This release features various documentation fixes, minor bugfixes, and code cleanup.

## Bugfixes

- PR #267: Add neglected <ToolkitAM1BCC> documentation to the SMIRNOFF 0.2 spec
- PR #258: General cleanup and removal of unused/inaccessible code.
- PR #244: Improvements and typo fixes for BRD4:inhibitor benchmark

### 1.2.5 0.2.0 - Initial RDKit support

This version of the toolkit introduces many new features on the way to a 1.0.0 release.

## New features

- Major overhaul, resulting in the creation of the [SMIRNOFF 0.2 specification](#) and its XML representation
- Updated API and infrastructure for reference SMIRNOFF ForceField implementation
- Implementation of modular ParameterHandler classes which process the topology to add all necessary forces to the system.
- Implementation of modular ParameterIOHandler classes for reading/writing different serialized SMIRNOFF forcefield representations
- Introduction of Molecule and Topology classes for representing molecules and biomolecular systems
- New ToolkitWrapper interface to RDKit, OpenEye, and AmberTools toolkits, managed by ToolkitRegistry
- API improvements to more closely follow [PEP8](#) guidelines
- Improved documentation and examples

### 1.2.6 0.1.0

This is an early preview release of the toolkit that matches the functionality described in the preprint describing the SMIRNOFF v0.1 force field format: [\[DOI\]](#).

## New features

This release features additional documentation, code comments, and support for automated testing.

## Bugfixes

### Treatment of improper torsions

A significant (though currently unused) problem in handling of improper torsions was corrected. Previously, non-planar impropers did not behave correctly, as six-fold impropers have two potential chiralities. To remedy this, SMIRNOFF impropers are now implemented as three-fold impropers with consistent chirality. However, current force fields in the SMIRNOFF format had no non-planar impropers, so this change is mainly aimed at future work.

## 1.3 The SMIRks Native Open Force Field (SMIRNOFF) specification

SMIRNOFF is a specification for encoding molecular mechanics force fields from the [Open Force Field Initiative](#) based on direct chemical perception using the broadly-supported [SMARTS](#) language, utilizing atom tagging extensions from [SMIRKS](#).

### 1.3.1 Authors and acknowledgments

The SMIRNOFF specification was designed by the [Open Force Field Initiative](#).

Primary contributors include:

- Caitlin C. Bannan (University of California, Irvine) <bannanc@uci.edu>
- Christopher I. Bayly (OpenEye Software) <bayly@eyesopen.com>
- John D. Chodera (Memorial Sloan Kettering Cancer Center) <john.chodera@choderlab.org>
- David L. Mobley (University of California, Irvine) <dmobley@uci.edu>

SMIRNOFF and its reference implementation in the `openforcefield` toolkit was heavily inspired by the `ForceField` class from the [OpenMM](#) molecular simulation package, and its associated [XML format](#), developed by Peter K. Eastman (Stanford University).

### 1.3.2 Representations and encodings

A force field in the SMIRNOFF format can be encoded in multiple representations. Currently, only an [XML](#) representation is supported by the reference implementation of the [openforcefield toolkit](#).

#### XML representation

A SMIRNOFF force field can be described in an [XML](#) representation, which provides a human- and machine-readable form for encoding the parameter set and its typing rules. This document focuses on describing the XML representation of the force field.

- By convention, XML-encoded SMIRNOFF force fields use an `.offxml` extension if written to a file to prevent confusion with other file formats.
- In XML, numeric quantities appear as strings, like "1" or "2.3".
- Integers should always be written without a decimal point, such as "1", "9".
- Non-integral numbers, such as parameter values, should be written with a decimal point, such as "1.23", "2.".

- In XML, certain special characters that occur in valid SMARTS/SMIRKS patterns (such as ampersand &) must be specially encoded. See [this list of XML and HTML character entity references](#) for more details.

### Future representations: JSON, MessagePack, YAML, and TOML

We are considering supporting [JSON](#), [MessagePack](#), [YAML](#), and [TOML](#) representations as well.

### 1.3.3 Reference implementation

A reference implementation of the SMIRNOFF XML specification is provided in the [openforcefield toolkit](#).

### 1.3.4 Support for molecular simulation packages

The reference implementation currently generates parameterized molecular mechanics systems for the GPU-accelerated [OpenMM](#) molecular simulation toolkit. Parameterized systems can subsequently be converted for use in other popular molecular dynamics simulation packages (including [AMBER](#), [CHARMM](#), [NAMD](#), [Desmond](#), and [LAMMPS](#)) via [ParmEd](#) and [InterMol](#). See [Converting SMIRNOFF parameterized systems to other simulation packages](#) for more details.

### 1.3.5 Basic structure

A reference implementation of a SMIRNOFF force field parser that can process XML representations (denoted by `.offxml` file extensions) can be found in the `ForceField` class of the `openforcefield.typing.engines.smirnoff` module.

Below, we describe the main structure of such an XML representation.

#### The enclosing `<SMIRNOFF>` tag

A SMIRNOFF forcefield XML specification always is enclosed in a `<SMIRNOFF>` tag, with certain required attributes provided. The required and permitted attributes defined in the `<SMIRNOFF>` are recorded in the `version` attribute, which describes the top-level attributes that are expected or permitted to be defined.

```
<SMIRNOFF version="0.3" aromaticity_model="OEAroModel_MDL">
...
</SMIRNOFF>
```

#### Versioning

The SMIRNOFF force field format supports versioning via the `version` attribute to the root `<SMIRNOFF>` tag, e.g.:

```
<SMIRNOFF version="0.3" aromaticity_model="OEAroModel_MDL">
...
</SMIRNOFF>
```

The version format is  $x.y$ , where  $x$  denotes the major version and  $y$  denotes the minor version. SMIRNOFF versions are guaranteed to be backward-compatible within the *same major version number series*, but it is possible major version increments will break backwards-compatibility.

SMIRNOFF tag version	Required attributes	Optional attributes
0.1	aromaticity_model	Date, Author
0.2	aromaticity_model	Date, Author
0.3	aromaticity_model	Date, Author

The SMIRNOFF tag versions describe the required and allowed force field-wide settings. The list of keywords is as follows:

### Aromaticity model

The `aromaticity_model` specifies the aromaticity model used for chemical perception (here, `OEAroModel_MDL`).

Currently, the only supported model is `OEAroModel_MDL`, which is implemented in both the RDKit and the OpenEye Toolkit.

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**Note:** Add link to complete open specification of `OEAroModel_MDL` aromaticity model.

---

### Metadata

Typically, date and author information is included:

```
<Date>2016-05-25</Date>
<Author>J. D. Chodera (MSKCC) charge increment tests</Author>
```

The `<Date>` tag should conform to ISO 8601 date formatting guidelines, such as `2018-07-14` or `2018-07-14T08:50:48+00:00` (UTC time).

### Parameter generators

Within the `<SMIRNOFF>` tag, top-level tags encode parameters for a force field based on a SMARTS/SMIRKS-based specification describing the chemical environment the parameters are to be applied to. The file has tags corresponding to OpenMM force terms (Bonds, Angles, ProperTorsions, etc., as discussed in more detail below); these specify functional form and other information for individual force terms.

```
<Angles version="0.3" potential="harmonic">
  ...
</Angles>
```

which introduces the following `Angle` child elements which will use a harmonic potential.

### Specifying parameters

Under each of these force terms, there are tags for individual parameter lines such as these:

```
<Angles version="0.3" potential="harmonic">
  <Angle smirks="[a,A:1]-[#6X4:2]-[a,A:3]" angle="109.50*degree" k="100.0*kilocalorie_per_mole/
  ↪radian**2"/>
  <Angle smirks="#1:1]-[#6X4:2]-#1:3]" angle="109.50*degree" k="70.0*kilocalorie_per_mole/radian**2"/
  ↪>
</Angles>
```

The first of these specifies the `smirks` attribute as `[a,A:1]-[#6X4:2]-[a,A:3]`, specifying a SMIRKS pattern that matches three connected atoms specifying an angle. This particular SMIRKS pattern matches a tetravalent carbon at the center with single bonds to two atoms of any type. This pattern is essentially a [SMARTS](#) string with numerical atom tags commonly used in [SMIRKS](#) to identify atoms in chemically unique environments—these can be thought of as tagged regular expressions for identifying chemical environments, and atoms within those environments. Here, `[a,A]` denotes any atom—either aromatic (`a`) or aliphatic (`A`), while `[#6X4]` denotes a carbon by element number (#6) that with four substituents (`X4`). The symbol `-` joining these groups denotes a single bond. The strings `:1`, `:2`, and `:2` label these atoms as indices 1, 2, and 3, with 2 being the central atom. Equilibrium angles are provided as the `angle` attribute, along with force constants as the `k` attribute (with corresponding units included in the expression).

---

**Note:** The reference implementation of the SMIRNOFF specification implemented in the Open Force Field toolkit will, by default, raise an exception if an unexpected attribute is encountered. The toolkit can be configured to accept non-spec keywords, but these are considered “cosmetic” and will not be evaluated. For example, providing an `<Angle>` tag that also specifies a second force constant `k2` will result in an exception, unless the user specifies that “cosmetic” attributes should be accepted by the parser.

---

### SMIRNOFF parameter specification is hierarchical

Parameters that appear later in a SMIRNOFF specification override those which come earlier if they match the same pattern. This can be seen in the example above, where the first line provides a generic angle parameter for any tetravalent carbon (single bond) angle, and the second line overrides this for the specific case of a hydrogen-(tetravalent carbon)-hydrogen angle. This hierarchical structure means that a typical parameter file will tend to have generic parameters early in the section for each force type, with more specialized parameters assigned later.

### Multiple SMIRNOFF representations can be processed in sequence

Multiple SMIRNOFF data sources (e.g. multiple OFFXML files) can be loaded by the `openforcefield` `ForceField` in sequence. If these files each contain unique top-level tags (such as `<Bonds>`, `<Angles>`, etc.), the resulting forcefield will be independent of the order in which the files are loaded. If, however, the same tag occurs in multiple files, the contents of the tags are merged, with the tags read later taking precedence over the parameters read earlier, provided the top-level tags have compatible attributes. The resulting force field will therefore depend on the order in which parameters are read.

This behavior is intended for limited use in appending very specific parameters, such as parameters specifying solvent models, to override standard parameters.

## 1.3.6 Units

To minimize the potential for [unit conversion errors](#), SMIRNOFF forcefields explicitly specify units in a form readable to both humans and computers for all unit-bearing quantities. Allowed values for units are given in `simtk.unit` (though in the future this may change to the more widely-used Python `pint` library). For example,

for the angle (equilibrium angle) and k (force constant) parameters in the <Angle> example block above, both attributes are specified as a mathematical expression

```
<Angle smirks="#1:1-[#6X4:2]-[#1:3]" angle="109.50*degree" k="70.0*kilocalorie_per_mole/radian**2"/>
```

For more information, see the [standard OpenMM unit system](#).

### 1.3.7 SMIRNOFF independently applies parameters to each class of potential energy terms

The SMIRNOFF uses direct chemical perception to assign parameters for potential energy terms independently for each term. Rather than first applying atom typing rules and then looking up combinations of the resulting atom types for each force term, the rules for directly applying parameters to atoms is compartmentalized in separate sections. The file consists of multiple top-level tags defining individual components of the potential energy (in addition to charge models or modifiers), with each section specifying the typing rules used to assign parameters for that potential term:

```
<Bonds version="0.3" potential="harmonic">
  <Bond smirks="#6X4:1-[#6X4:2]" length="1.526*angstrom" k="620.0*kilocalories_per_mole/angstrom**2"/>
  <Bond smirks="#6X4:1-[#1:2]" length="1.090*angstrom" k="680.0*kilocalories_per_mole/angstrom**2"/>
  ...
</Bonds>

<Angles version="0.3" potential="harmonic">
  <Angle smirks="[a,A:1]-[#6X4:2]-[a,A:3]" angle="109.50*degree" k="100.0*kilocalories_per_mole/radian**2"/>
  <Angle smirks="#1:1-[#6X4:2]-[#1:3]" angle="109.50*degree" k="70.0*kilocalories_per_mole/radian**2"/>
  ...
</Angles>
```

Each top-level tag specifying a class of potential energy terms has an attribute potential for specifying the functional form for the interaction. Common defaults are defined, but the goal is to eventually allow these to be overridden by alternative choices or even algebraic expressions in the future, once more molecular simulation packages support general expressions. We distinguish between functional forms available in all common molecular simulation packages (specified by keywords) and support for general functional forms available in a few packages (especially OpenMM, which supports a flexible set of custom forces defined by algebraic expressions) with an **EXPERIMENTAL** label.

Many of the specific forces are implemented as discussed in the [OpenMM Documentation](#); see especially [Section 19 on Standard Forces](#) for mathematical descriptions of these functional forms. Some top-level tags provide attributes that modify the functional form used to be consistent with packages such as AMBER or CHARMM.

### 1.3.8 Partial charge and electrostatics models

SMIRNOFF supports several approaches to specifying electrostatic models. Currently, only classical fixed point charge models are supported, but future extensions to the specification will support point multipoles, point polarizable dipoles, Drude oscillators, charge equilibration methods, and so on.

## <LibraryCharges>: Library charges for polymeric residues and special solvent models

**Warning:** This functionality is not yet implemented and will appear in a future version of the toolkit. Please see [Issue 25 on the Open Force Field Toolkit issue tracker](<https://github.com/openforcefield/openforcefield/issues/25>).

A mechanism is provided for specifying library charges that can be applied to molecules or residues that match provided templates. Library charges are applied first, and atoms for which library charges are applied will be excluded from alternative charging schemes listed below.

For example, to assign partial charges for a non-terminal ALA residue from the AMBER ff14SB parameter set:

```
<LibraryCharges version="0.3">
  <!-- match a non-terminal alanine residue with AMBER ff14SB partial charges-->
  <LibraryCharge name="ALA" smirks="[NX3:1]([#1:2])([#6])[#6H1:3]([#1:4])([#6:5]([#1:6])([#1:7])[
  ↪[#1:8])[#6:9](=[#8:10])[#7]" charge1="-0.4157*elementary_charge" charge2="0.2719*elementary_charge"_
  ↪charge3="0.0337*elementary_charge" charge4="0.0823*elementary_charge" charge5="-0.1825*elementary_
  ↪charge" charge6="0.0603*elementary_charge" charge7="0.0603*elementary_charge" charge8="0.
  ↪0603*elementary_charge" charge9="0.5973*elementary_charge" charge10="-0.5679*elementary_charge">
  ...
</LibraryCharges>
```

In this case, a SMIRKS string defining the residue tags each atom that should receive a partial charge, with the charges specified by attributes charge1, charge2, etc. The name attribute is optional. Note that, for a given template, chemically equivalent atoms should be assigned the same charge to avoid undefined behavior. If the template matches multiple non-overlapping sets of atoms, all such matches will be assigned the provided charges. If multiple templates match the same set of atoms, the last template specified will be used.

Solvent models or excipients can also have partial charges specified via the <LibraryCharges> tag. For example, to ensure water molecules are assigned partial charges for TIP3P water, we can specify a library charge entry:

```
<LibraryCharges version="0.3">
  <!-- TIP3P water oxygen with charge override -->
  <LibraryCharge name="TIP3P" smirks="[#1:1]-[#8X2H2+0:2]-[#1:3]" charge1="+0.417*elementary_charge"_
  ↪charge2="-0.834*elementary_charge" charge3="+0.417*elementary_charge"/>
</LibraryCharges>
```

## <ChargeIncrementModel>: Small molecule and fragment charges

**Warning:** This functionality is not yet implemented and will appear in a future version of the toolkit. This area of the SMIRNOFF spec is under further consideration. Please see [Issue 208 on the Open Force Field Toolkit issue tracker](<https://github.com/openforcefield/openforcefield/issues/208>).

In keeping with the AMBER force field philosophy, especially as implemented in small molecule force fields such as [GAFF](#), [GAFF2](#), and [parm@Frosst](#), partial charges for small molecules are usually assigned using a quantum chemical method (usually a semiempirical method such as [AM1](#)) and a [partial charge determination scheme](#) (such as [CM2](#) or [RESP](#)), then subsequently corrected via charge increment rules, as in the highly successful [AM1-BCC](#) approach.

Here is an example:

```

<ChargeIncrementModel version="0.3" number_of_conformers="10" quantum_chemical_method="AM1" partial_charge_method="CM2">
    <!-- A fractional charge can be moved along a single bond -->
    <ChargeIncrement smirks="#[6X4:1]-[#6X3a:2]" chargeincrement1="-0.0073*elementary_charge" />
    <ChargeIncrement smirks="#[6X4:1]-[#6X3a:2]-[#7]" chargeincrement1="+0.0943*elementary_charge" />
    <ChargeIncrement smirks="-#0.0943*elementary_charge" />
    <ChargeIncrement smirks="#[6X4:1]-[#8:2]" chargeincrement1="-0.0718*elementary_charge" />
    <ChargeIncrement smirks="+0.0718*elementary_charge" />
    <!-- Alternatively, fractional charges can be redistributed among any number of bonded atoms -->
    <ChargeIncrement smirks="N:1(H:2)(H:3)" chargeincrement1="+0.02*elementary_charge" chargeincrement2="-0.01*elementary_charge" chargeincrement3="-0.01*elementary_charge" />
</ChargeIncrementModel>

```

The sum of formal charges for the molecule or fragment will be used to determine the total charge the molecule or fragment will possess.

`<ChargeIncrementModel>` provides several optional attributes to control its behavior:

- The `number_of_conformers` attribute (default: "10") is used to specify how many conformers will be generated for the molecule (or capped fragment) prior to charging.
- The `quantum_chemical_method` attribute (default: "AM1") is used to specify the quantum chemical method applied to the molecule or capped fragment.
- The `partial_charge_method` attribute (default: "CM2") is used to specify how uncorrected partial charges are to be generated from the quantum chemical wavefunction. Later additions will add restrained electrostatic potential fitting (RESP) capabilities.

The `<ChargeIncrement>` tags specify how the quantum chemical derived charges are to be corrected to produce the final charges. The `chargeincrement#` attributes specify how much the charge on the associated tagged atom index (replacing #) should be modified. The sum of charge increments should equal zero.

Note that atoms for which library charges have already been applied are excluded from charging via `<ChargeIncrementModel>`.

Future additions will provide options for intelligently fragmenting large molecules and biopolymers, as well as a capping attribute to specify how fragments with dangling bonds are to be capped to allow these groups to be charged.

#### `<ToolkitAM1BCC>`: Temporary support for toolkit-based AM1-BCC partial charges

**Warning:** Until `<ChargeIncrementModel>` is implemented, support for the `<ToolkitAM1BCC>` tag has been enabled in the toolkit. This tag is not permanent and may be phased out in future versions of the spec.

This tag calculates partial charges using the default settings of the highest-priority cheminformatics toolkit that can perform [AM1-BCC charge assignment](#). Currently, if the OpenEye toolkit is licensed and available, this will use QuacPac configured to generate charges using [AM1-BCC ELF10](#) for each unique molecule in the topology. Otherwise [RDKit](#) will be used for initial conformer generation and the [AmberTools antechamber/sqm software](#) will be used for charge calculation.

If this tag is specified for a force field, conformer generation will be performed regardless of whether conformations of the input molecule were provided. If RDKit/AmberTools are used as the toolkit backend for this calculation, only the first conformer is used for AM1-BCC calculation.

The charges generated by this tag may differ depending on which toolkits are available.

Note that atoms for which prespecified or library charges have already been applied are excluded from charging via <ToolkitAM1BCC>.

### Prespecified charges (reference implementation only)

In our reference implementation of SMIRNOFF in the openforcefield toolkit, we also provide a method for specifying user-defined partial charges during system creation. This functionality is accessed by using the `charge_from_molecules` optional argument during system creation, such as in `ForceField.create_openmm_system(topology, charge_from_molecules=molecule_list)`. When this optional keyword is provided, all matching molecules will have their charges set by the entries in `molecule_list`. This method is provided solely for convenience in developing and exploring alternative charging schemes; actual force field releases for distribution will use one of the other mechanisms specified above.

## 1.3.9 Parameter sections

A SMIRNOFF force field consists of one or more force field term definition sections. For the most part, these sections independently define how a specific component of the potential energy function for a molecular system is supposed to be computed (such as bond stretch energies, or Lennard-Jones interactions), as well as how parameters are to be assigned for this particular term. Each parameter section contains a version, which encodes the behavior of the section, as well as the required and optional attributes the top-level tag and SMIRKS-based parameters. This decoupling of how parameters are assigned for each term provides a great deal of flexibility in composing new force fields while allowing a minimal number of parameters to be used to achieve accurate modeling of intramolecular forces.

Below, we describe the specification for each force field term definition using the XML representation of a SMIRNOFF force field.

As an example of a complete SMIRNOFF force field specification, see the prototype [SMIRNOFF99Frosst offxml](#).

---

**Note:** Not all parameter sections *must* be specified in a SMIRNOFF force field. A wide variety of force field terms are provided in the specification, but a particular force field only needs to define a subset of those terms.

---

#### <vdW>

van der Waals force parameters, which include repulsive forces arising from Pauli exclusion and attractive forces arising from dispersion, are specified via the <vdW> tag with sub-tags for individual Atom entries, such as:

```
<vdW version="0.3" potential="Lennard-Jones-12-6" combining_rules="Lorentz-Berthelot" scale12="0.0"_
  ↪scale13="0.0" scale14="0.5" scale15="1.0" switch_width="8.0*angstrom" cutoff="9.0*angstrom" long_
  ↪range_dispersion="isotropic">
  <Atom smirks="#1:1" sigma="1.4870*angstrom" epsilon="0.0157*kilocalories_per_mole"/>
  <Atom smirks="#1:1-[#6]" sigma="1.4870*angstrom" epsilon="0.0157*kilocalories_per_mole"/>
  ...
</vdW>
```

For standard Lennard-Jones 12-6 potentials (specified via `potential="Lennard-Jones-12-6"`), the `epsilon` parameter denotes the well depth, while the size property can be specified either via providing the `sigma`

attribute, such as `sigma="1.3*angstrom"`, or via the `r_0/2` (`rmin/2`) values used in AMBER force fields (here denoted `rmin_half` as in the example above). The two are related by  $r_0 = 2^{(1/6)} * \text{sigma}$  and conversion is done internally in ForceField into the `sigma` values used in OpenMM.

Attributes in the `<vdW>` tag specify the scaling terms applied to the energies of 1-2 (`scale12`, default: 0), 1-3 (`scale13`, default: 0), 1-4 (`scale14`, default: 0.5), and 1-5 (`scale15`, default: 1.0) interactions, as well as the distance at which a switching function is applied (`switch_width`, default: `"1.0*angstrom"`), the cutoff (`cutoff`, default: `"9.0*angstroms"`), and long-range dispersion treatment scheme (`long_range_dispersion`, default: `"isotropic"`).

The potential attribute (default: "none") specifies the potential energy function to use. Currently, only `potential="Lennard-Jones-12-6"` is supported:

$$U(r) = 4\epsilon \left( \frac{\sigma}{r} \right)^{12} - \left( \frac{\sigma}{r} \right)^6$$

The `combining_rules` attribute (default: "none") currently only supports "Lorentz-Berthelot", which specifies the geometric mean of `epsilon` and arithmetic mean of `sigma`. Support for other Lennard-Jones mixing schemes will be added later: Waldman-Hagler, Fender-Halsey, Kong, Tang-Toennies, Pena, Hudson-McCoubrey, Sikora.

Later revisions will add support for additional potential types (e.g., Buckingham-exp-6), as well as the ability to support arbitrary algebraic functional forms using a scheme such as

```
<vdW version="0.3" potential="4*epsilon*((sigma/r)^12-(sigma/r)^6)" scale12="0.0" scale13="0.0" scale14="0.5" scale15="1" switch_width="8.0*angstrom" cutoff="9.0*angstrom" long_range_dispersion="isotropic">
  <CombiningRules>
    <CombiningRule parameter="sigma" function="(sigma1+sigma2)/2"/>
    <CombiningRule parameter="epsilon" function="sqrt(epsilon1*epsilon2)"/>
  </CombiningRules>
  <Atom smirks="#1:1" sigma="1.4870*angstrom" epsilon="0.0157*kilocalories_per_mole"/>
  <Atom smirks="#1:1-[#6]" sigma="1.4870*angstrom" epsilon="0.0157*kilocalories_per_mole"/>
  ...
</vdW>
```

If the `<CombiningRules>` tag is provided, it overrides the `combining_rules` attribute.

Later revisions will also provide support for special interactions using the `<AtomPair>` tag:

```
<vdW version="0.3" potential="Lennard-Jones-12-6" combining_rules="Lorentz-Berthelot" scale12="0.0" scale13="0.0" scale14="0.5" scale15="1">
  <AtomPair smirks1="#1:1" smirks2="#6:2" sigma="1.4870*angstrom" epsilon="0.0157*kilocalories_per_mole"/>
  ...
</vdW>
```

vdW section tag version	Tag attributes and default values	Required parameter attributes	Optional parameter attributes
0.3	<code>potential="Lennard-Jones-12-6", combining_rules="Lorentz-Berthelot", scale12="0", scale13="0", scale14="0.5", scale15="1.0", cutoff="9.0*angstrom", switch_width="1.0*angstrom", method="cutoff"</code>	<code>smirks, epsilon, (sigma OR rmin_half)</code>	<code>id, parent_id</code>

## <Electrostatics>

Electrostatic interactions are specified via the <Electrostatics> tag.

```
<Electrostatics version="0.3" method="PME" scale12="0.0" scale13="0.0" scale14="0.833333" scale15="1.0"/>
```

The method attribute specifies the manner in which electrostatic interactions are to be computed:

- PME - particle mesh Ewald should be used (DEFAULT); can only apply to periodic systems
- reaction-field - reaction-field electrostatics should be used; can only apply to periodic systems
- Coulomb - direct Coulomb interactions (with no reaction-field attenuation) should be used

The interaction scaling parameters applied to atoms connected by a few bonds are

- scale12 (default: 0) specifies the scaling applied to 1-2 bonds
- scale13 (default: 0) specifies the scaling applied to 1-3 bonds
- scale14 (default: 0.833333) specifies the scaling applied to 1-4 bonds
- scale15 (default: 1.0) specifies the scaling applied to 1-5 bonds

Currently, no child tags are used because the charge model is specified via different means (currently library charges or BCCs).

For methods where the cutoff is not simply an implementation detail but determines the potential energy of the system (reaction-field and Coulomb), the cutoff distance must also be specified, and a switch\_width if a switching function is to be used.

Electrostatics section tag version	Tag attributes and default values	Required parameter attributes	Optional parameter attributes
0.3	scale12="0", scale13="0", scale14="0.833333", scale15="1.0", cutoff="9.0*angstrom", switch_width="0*angstrom", method="PME"	N/A	N/A

## <Bonds>

Bond parameters are specified via a <Bonds>...</Bonds> block, with individual <Bond> tags containing attributes specifying the equilibrium bond length (length) and force constant (k) values for specific bonds. For example:

```
<Bonds version="0.3" potential="harmonic">
  <Bond smirks="#6X4:1]-[#6X4:2]" length="1.526*angstrom" k="620.0*kilocalories_per_mole/angstrom**2"/>
  <Bond smirks="#6X4:1]-[#1:2]" length="1.090*angstrom" k="680.0*kilocalories_per_mole/angstrom**2"/>
  ...
</Bonds>
```

Currently, only potential="harmonic" is supported, where we utilize the standard harmonic functional form:

$$U(r) = (k/2)*(r-length)^2$$

Later revisions will add support for additional potential types and the ability to support arbitrary algebraic functional forms. If the potential attribute is omitted, it defaults to harmonic.

Note that AMBER and CHARMM define a modified functional form, such that  $U(r) = k*(r-length)^2$ , so that force constants would need to be multiplied by two in order to be used in the SMIRNOFF format.

Constrained bonds are handled by a separate <Constraints> tag, which can either specify constraint distances or draw them from equilibrium distances specified in <Bonds>.

### Fractional bond orders (EXPERIMENTAL)

**Warning:** This functionality is not yet implemented and will appear in a future version of the toolkit.

Fractional bond orders can be used to allow interpolation of bond parameters. For example, these parameters:

```
<Bonds version="0.3" potential="harmonic">
  <Bond smirks="[#6X3:1]-[#6X3:2]" k="820.0*kilocalories_per_mole/angstrom**2" length="1.45*angstrom"/>
  <Bond smirks="[#6X3:1]:[#6X3:2]" k="938.0*kilocalories_per_mole/angstrom**2" length="1.40*angstrom"/>
  <Bond smirks="[#6X3:1]=[#6X3:2]" k="1098.0*kilocalories_per_mole/angstrom**2" length="1.35*angstrom"/>
  ...
  ...
```

can be replaced by a single parameter line by first invoking the fractional\_bondorder\_method attribute to specify a method for computing the fractional bond order and fractional\_bondorder\_interpolation for specifying the procedure for interpolating parameters between specified integral bond orders:

```
<Bonds version="0.3" potential="harmonic" fractional_bondorder_method="Wiberg" fractional_bondorder_
  interpolation="linear">
  <Bond smirks="[#6X3:1]!#[#6X3:2]" k_bondorder1="820.0*kilocalories_per_mole/angstrom**2" k_
  bondorder2="1098*kilocalories_per_mole/angstrom**2" length_bondorder1="1.45*angstrom" length_
  bondorder2="1.35*angstrom"/>
  ...
  ...
```

This allows specification of force constants and lengths for bond orders 1 and 2, and then interpolation between those based on the partial bond order.

- fractional\_bondorder\_method defaults to none, but the Wiberg method is supported.
- fractional\_bondorder\_interpolation defaults to linear, which is the only supported scheme for now.

Bonds section tag version	Tag attributes and default values	Required parameter attributes	Optional parameter attributes
0.3	potential="harmonic", fractional_bondorder_method="Wiberg", fractional_bondorder_interpolation="linear"	smirks, length, k	id, parent_id

### <Angles>

Angle parameters are specified via an <Angles>...</Angles> block, with individual <Angle> tags containing attributes specifying the equilibrium angle (angle) and force constant (k), as in this example:

```
<Angles version="0.3" potential="harmonic">
  <Angle smirks="[a,A:1]-[#6X4:2]-[a,A:3]" angle="109.50*degree" k="100.0*kilocalories_per_mole/
  ↪radian**2"/>
  <Angle smirks="#1:1]-[#6X4:2]-[#1:3]" angle="109.50*degree" k="70.0*kilocalories_per_mole/radian**2
  ↪"/>
  ...
</Angles>
```

Currently, only `potential="harmonic"` is supported, where we utilize the standard harmonic functional form:

$$U(r) = (k/2) * (\theta - \theta_0)^2$$

Later revisions will add support for additional potential types and the ability to support arbitrary algebraic functional forms. If the `potential` attribute is omitted, it defaults to harmonic.

**Note that AMBER and CHARMM define a modified functional form**, such that  $U(r) = k * (\theta - \theta_0)^2$ , so that force constants would need to be multiplied by two in order to be used in the SMIRNOFF format.

Angles section tag version	Tag attributes and default values	Required parameter attributes	Optional parameter attributes
0.3	<code>potential="harmonic"</code>	<code>smirks, angle, k</code>	<code>id, parent_id</code>

### <ProperTorsions>

Proper torsions are specified via a `<ProperTorsions>...</ProperTorsions>` block, with individual `<Proper>` tags containing attributes specifying the periodicity (`periodicity#`), phase (`phase#`), and barrier height (`k#`).

```
<ProperTorsions version="0.3" potential="k*(1+cos(periodicity*theta-phase))">
  <Proper smirks="[a,A:1]-[#6X4:2]-[#6X4:3]-[a,A:4]" idivf1="9" periodicity1="3" phase1="0.0*degree"
  ↪k1="1.40*kilocalories_per_mole"/>
  <Proper smirks="#6X4:1]-[#6X4:2]-[#8X2:3]-[#6X4:4]" idivf1="1" periodicity1="3" phase1="0.0*degree"
  ↪k1="0.383*kilocalories_per_mole" idivf2="1" periodicity2="2" phase2="180.0*degree" k2="0.
  ↪1*kilocalories_per_mole"/>
  ...
</ProperTorsions>
```

Here, child Proper tags specify at least `k1`, `phase1`, and `periodicity1` attributes for the corresponding parameters of the first force term applied to this torsion. However, additional values are allowed in the form `k#, phase#, and periodicity#`, where all # values must be consecutive (e.g., it is impermissible to specify `k1` and `k3` values without a `k2` value) but # can go as high as necessary.

For convenience, and optional attribute specifies a torsion multiplicity by which the barrier height should be divided (`idivf#`). The default behavior of this attribute can be controlled by the top-level attribute `default_idivf` (default: "auto") for `<ProperTorsions>`, which can be an integer (such as "1") controlling the value of `idivf` if not specified or "auto" if the barrier height should be divided by the number of torsions impinging on the central bond. For example:

```
<ProperTorsions version="0.3" potential="k*(1+cos(periodicity*theta-phase))" default_idivf="auto">
  <Proper smirks="[a,A:1]-[#6X4:2]-[#6X4:3]-[a,A:4]" periodicity1="3" phase1="0.0*degree" k1="1.
  ↪40*kilocalories_per_mole"/>
  ...
</ProperTorsions>
```

Currently, only `potential="k*(1+cos(periodicity*theta-phase))"` is supported, where we utilize the functional form:

```
U = \sum_{i=1}^N k_i * (1 + cos(periodicity_i * phi - phase_i))
```

**Note:** AMBER defines a modified functional form, such that  $U = \sum_{i=1}^N (k_i/2) * (1 + \cos(\text{periodicity}_i * \phi - \text{phase}_i))$ , so that barrier heights would need to be divided by two in order to be used in the SMIRNOFF format.

If the potential attribute is omitted, it defaults to `k*(1+cos(periodicity*theta-phase))`.

ProperTorsions section tag version	Tag attributes and default values	Required parameter attributes	Optional parameter attributes
0.3	<code>potential="k*(1+cos(periodicity*theta-phase))", k, phase, default_idivf="auto"</code>	<code>periodicity</code>	<code>idivf, id, parent_id</code>

### <ImproperTorsions>

Improper torsions are specified via an `<ImproperTorsions>...</ImproperTorsions>` block, with individual `<Improper>` tags containing attributes that specify the same properties as `<ProperTorsions>`:

```
<ImproperTorsions version="0.3" potential="k*(1+cos(periodicity*theta-phase))">
  <Improper smirks="[:1]~[#6X3:2](=[#7X2,#7X3+1:3])~[#7:4]" k1="10.5*kilocalories_per_mole"
    <periodicity1="2" phase1="180.*degree"/>
  ...
</ImproperTorsions>
```

Currently, only `potential="charmm"` is supported, where we utilize the functional form:

```
U = \sum_{i=1}^N k_i * (1 + cos(periodicity_i * phi - phase_i))
```

**Note:** AMBER defines a modified functional form, such that  $U = \sum_{i=1}^N (k_i/2) * (1 + \cos(\text{periodicity}_i * \phi - \text{phase}_i))$ , so that barrier heights would need to be divided by two in order to be used in the SMIRNOFF format.

If the potential attribute is omitted, it defaults to `charmm`.

The improper torsion energy is computed as the average over all three impropers (all with the same handedness) in a `trefoil`. This avoids the dependence on arbitrary atom orderings that occur in more traditional typing engines such as those used in AMBER. The second atom in an improper (in the example above, the trivalent carbon) is the central atom in the trefoil.

ImproperTorsions section tag version	Tag attributes and default values	Required parameter attributes	Optional parameter attributes
0.3	<code>potential="k*(1+cos(periodicity*theta-phase))", k, phase, default_idivf="auto"</code>	<code>periodicity</code>	<code>idivf, id, parent_id</code>

### <GBSA>

**Warning:** This functionality is not yet implemented and will appear in a future version of the toolkit.

Generalized-Born surface area (GBSA) implicit solvent parameters are optionally specified via a `<GBSA>...` `</GBSA>` using `<Atom>` tags with GBSA model specific attributes:

```
<GBSA version="0.3" gb_model="OBC1" solvent_dielectric="78.5" solute_dielectric="1" sa_model="ACE"_
  ↪surface_area_penalty="5.4*calories/mole/angstroms**2" solvent_radius="1.4*angstroms">
  <Atom smirks="#1:1" radius="0.12*angstrom" scale="0.85"/>
  <Atom smirks="#1:1~#[#6]" radius="0.13*angstrom" scale="0.85"/>
  <Atom smirks="#1:1~#[#8]" radius="0.08*angstrom" scale="0.85"/>
  <Atom smirks="#1:1~#[#16]" radius="0.08*angstrom" scale="0.85"/>
  <Atom smirks="#6:1" radius="0.22*angstrom" scale="0.72"/>
  <Atom smirks="#7:1" radius="0.155*angstrom" scale="0.79"/>
  <Atom smirks="#8:1" radius="0.15*angstrom" scale="0.85"/>
  <Atom smirks="#9:1" radius="0.15*angstrom" scale="0.88"/>
  <Atom smirks="#14:1" radius="0.21*angstrom" scale="0.8"/>
  <Atom smirks="#15:1" radius="0.185*angstrom" scale="0.86"/>
  <Atom smirks="#16:1" radius="0.18*angstrom" scale="0.96"/>
  <Atom smirks="#17:1" radius="0.17*angstrom" scale="0.8"/>
</GBSA>
```

## Supported Generalized Born (GB) models

In the `<GBSA>` tag, `gb_model` selects which GB model is used. Currently, this can be selected from a subset of the GBSA models available in OpenMM:

- HCT : [Hawkins-Cramer-Truhlar](#) (corresponding to `igb=1` in AMBER): requires parameters [`radius`, `scale`]
- OBC1 : [Onufriev-Bashford-Case](#) using the GB(OBC)I parameters (corresponding to `igb=2` in AMBER): requires parameters [`radius`, `scale`]
- OBC2 : [Onufriev-Bashford-Case](#) using the GB(OBC)II parameters (corresponding to `igb=5` in AMBER): requires parameters [`radius`, `scale`]

If the `gb_model` attribute is omitted, it defaults to OBC1.

The attributes `solvent_dielectric` and `solute_dielectric` specify solvent and solute dielectric constants used by the GB model. In this example, `radius` and `scale` are per-particle parameters of the OBC1 GB model supported by OpenMM. Units are for these per-particle parameters (such as `radius_units`) specified in the `<GBSA>` tag.

## Surface area (SA) penalty model

The `sa_model` attribute specifies the solvent-accessible surface area model (“SA” part of GBSA) if one should be included; if omitted, no SA term is included.

Currently, only the [analytical continuum electrostatics \(ACE\)](#) model, designated ACE, can be specified, but there are plans to add more models in the future, such as the Gaussian solvation energy component of [EEF1](#). If `sa_model` is not specified, it defaults to ACE.

The ACE model permits two additional parameters to be specified:

- The `surface_area_penalty` attribute specifies the surface area penalty for the ACE model. (Default: 5.4 `calories/mole/angstroms**2`)
- The `solvent_radius` attribute specifies the solvent radius. (Default: 1.4 `angstroms`)

GBSA sec- tion tag version	Tag attributes and default values	Required parameter attributes	Optional parameter attributes
0.3	gb_model="OBC1", solvent_dielectric="78.5", solute_dielectric="1", sa_model="ACE", surface_area_penalty="5.4*calories/mole/angstrom**2", solvent_radius="1.4*angstrom"	smirks, radius, scale	id, parent_id

### <Constraints>

Bond length or angle constraints can be specified through a <Constraints> block, which can constrain bonds to their equilibrium lengths or specify an interatomic constraint distance. Two atoms must be tagged in the smirks attribute of each <Constraint> record.

To constrain the separation between two atoms to their equilibrium bond length, it is critical that a <Bonds> record be specified for those atoms:

```
<Constraints version="0.3" >
  <!-- constrain all bonds to hydrogen to their equilibrium bond length -->
  <Constraint smirks="#[1:1]-[*:2]" />
</Constraints>
```

Note that the two atoms must be bonded in the specified Topology for the equilibrium bond length to be used.

To specify the constraint distance, or constrain two atoms that are not directly bonded (such as the hydrogens in rigid water models), specify the distance attribute (and optional distance\_unit attribute for the <Constraints> tag):

```
<Constraints version="0.3">
  <!-- constrain water O-H bond to equilibrium bond length (overrides earlier constraint) -->
  <Constraint smirks="#[1:1]-[#8X2H2:2]-[#1]" distance="0.9572*angstrom"/>
  <!-- constrain water H...H, calculating equilibrium length from H-O-H equilibrium angle and H-O_
  ↪equilibrium bond lengths -->
  <Constraint smirks="#[1:1]-[#8X2H2]-[#1:2]" distance="1.8532*angstrom"/>
</Constraints>
```

Typical molecular simulation practice is to constrain all bonds to hydrogen to their equilibrium bond lengths and enforce rigid TIP3P geometry on water molecules:

```
<Constraints version="0.3" >
  <!-- constrain all bonds to hydrogen to their equilibrium bond length -->
  <Constraint smirks="#[1:1]-[*:2]" />
  <!-- TIP3P rigid water -->
  <Constraint smirks="#[1:1]-[#8X2H2:2]-[#1]" distance="0.9572*angstrom"/>
  <Constraint smirks="#[1:1]-[#8X2H2]-[#1:2]" distance="1.8532*angstrom"/>
</Constraints>
```

Constraint section tag version	Required tag attributes and de- fault values	Required parameter attributes	Optional parameter attributes
0.3		smirks	distance

### 1.3.10 Advanced features

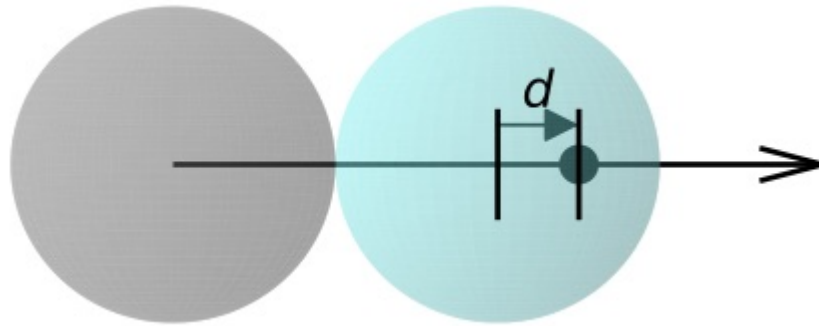
Standard usage is expected to rely primarily on the features documented above and potentially new features. However, some advanced features will also be supported.

#### <VirtualSites>: Virtual sites for off-atom charges

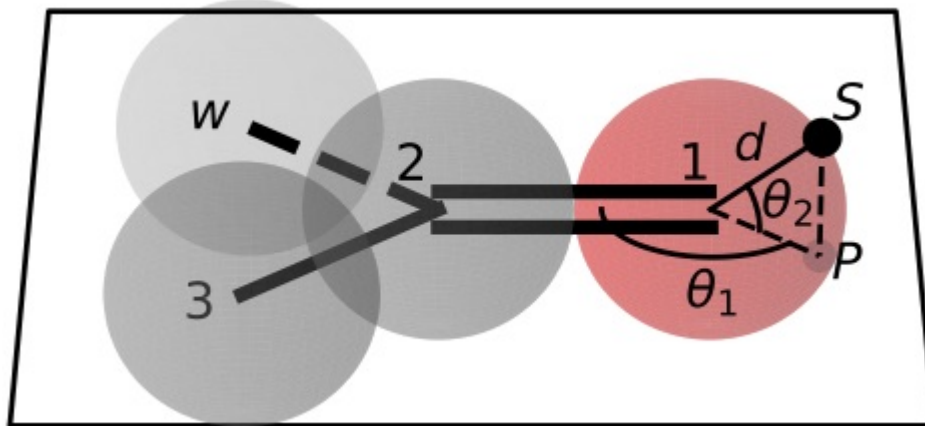
**Warning:** This functionality is not yet implemented and will appear in a future version of the toolkit

We will implement experimental support for placement of off-atom (off-center) charges in a variety of contexts which may be chemically important in order to allow easy exploration of when these will be warranted. We will support the following different types or geometries of off-center charges (as diagrammed below):

- **BondCharge:** This supports placement of a virtual site S along a vector between two specified atoms, e.g. to allow for a sigma hole for halogens or similar contexts. With positive values of the distance, the virtual site lies outside the first indexed atom (green in this image).

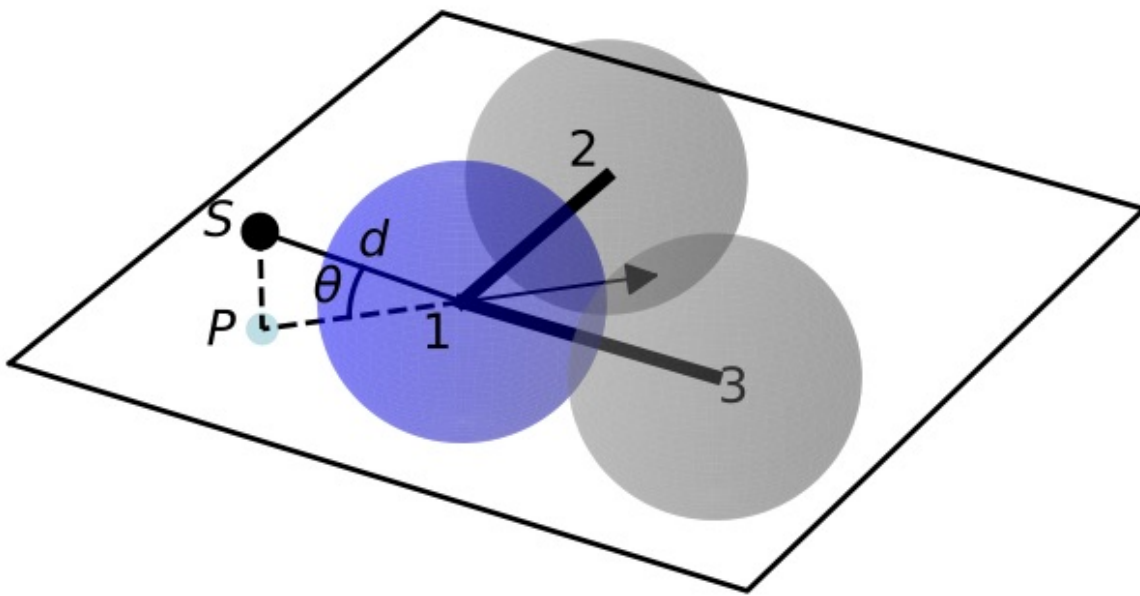


- **MonovalentLonePair:** This is originally intended for situations like a carbonyl, and allows placement of a virtual site S at a specified distance d, inPlaneAngle (theta 1 in the diagram), and outOfPlaneAngle (theta 2 in the diagram) relative to a central atom and two connected atoms.

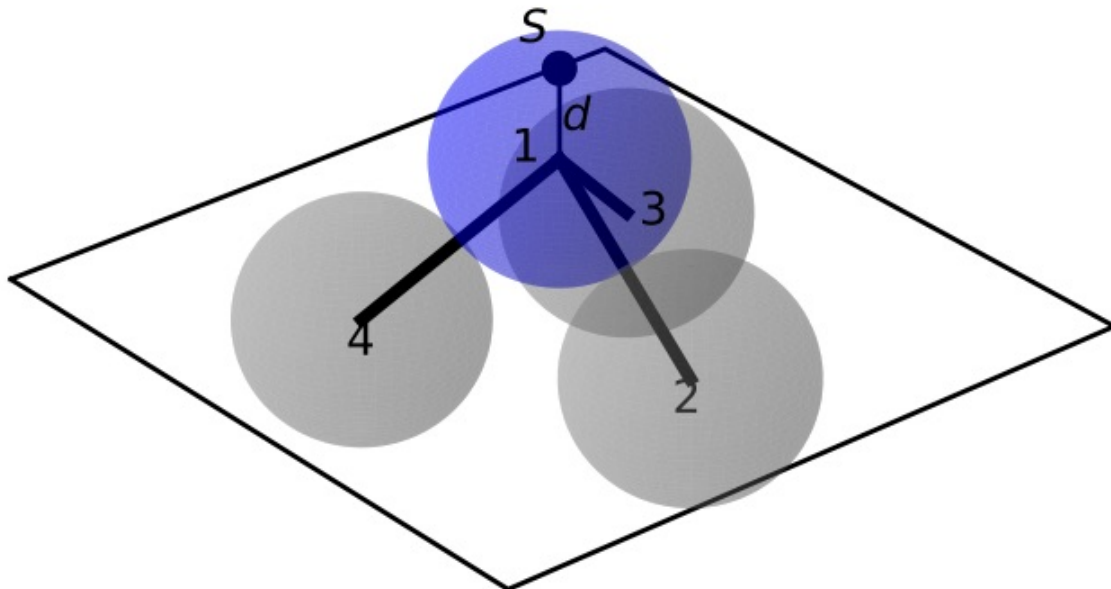


- **DivalentLonePair:** This is suitable for cases like four-point and five-point water models as well as pyrimidine; a charge site S lies a specified distance d from the central atom among three atoms (blue) along the bisector of the angle between the atoms (if outOfPlaneAngle is zero) or out of the plane by

the specified angle (if `outOfPlaneAngle` is nonzero) with its projection along the bisector. For positive values fo the distance  $d$  the virtual site lies outside the 2-1-3 angle and for negative values it lies inside.



- `TrivalentLonePair`: This is suitable for planar or tetrahedral nitrogen lone pairs; a charge site  $S$  lies above the central atom (e.g. nitrogen, blue) a distance  $d$  along the vector perpendicular to the plane of the three connected atoms (2,3,4). With positive values of  $d$  the site lies above the nitrogen and with negative values it lies below the nitrogen.



Each virtual site receives charge which is transferred from the desired atoms specified in the SMIRKS pattern via a `chargeincrement#` parameter, e.g., if `chargeincrement1=+0.1*elementary_charge` then the virtual site will receive a charge of -0.1 and the atom labeled 1 will have its charge adjusted upwards by +0.1. N may index any indexed atom. Increments which are left unspecified default to zero. Additionally, each virtual site can bear Lennard-Jones parameters, specified by `sigma` and `epsilon` or `rmin_half` and `epsilon`.

If unspecified these also default to zero.

In the SMIRNOFF format, these are encoded as:

```
<VirtualSites version="0.3">
    <!-- sigma hole for halogens: "distance" denotes distance along the 2->1 bond vector, measured from
    atom 2 -->
    <!-- Specify that 0.2 charge from atom 1 and 0.1 charge units from atom 2 are to be moved to the
    virtual site, and a small Lennard-Jones site is to be added (sigma = 0.1*angstroms, epsilon=0.05*kcal/
    mol) -->
        <VirtualSite type="BondCharge" smirks="[Cl:1]-[C:2]" distance="0.30*angstrom" chargeincrement1="+0.
        2*elementary_charge" chargeincrement2="+0.1*elementary_charge" sigma="0.1*angstrom" epsilon="0.
        05*kilocalories_per_mole" />
            <!-- Charge increments can extend out to as many atoms as are labeled, e.g. with a third atom: -->
            <VirtualSite type="BondCharge" smirks="[Cl:1]-[C:2]~[*:3]" distance="0.30*angstrom"_
            chargeincrement1="+0.1*elementary_charge" chargeincrement2="+0.1*elementary_charge" chargeincrement3=
            "+0.05*elementary_charge" sigma="0.1*angstrom" epsilon="0.05*kilocalories_per_mole" />
                <!-- monovalent lone pairs: carbonyl -->
                <!-- X denotes the charge site, and P denotes the projection of the charge site into the plane of 1
                and 2. -->
                <!-- inPlaneAngle is angle point P makes with 1 and 2, i.e. P-1-2 -->
                <!-- outOfPlaneAngle is angle charge site (X) makes out of the plane of 2-1-3 (and P) measured from
                1 -->
                    <!-- Since unspecified here, sigma and epsilon for the virtual site default to zero -->
                    <VirtualSite type="MonovalentLonePair" smirks="[0:1]=[C:2]-[*:3]" distance="0.30*angstrom"_
                    outOfPlaneAngle="0*degree" inPlaneAngle="120*degree" chargeincrement1="+0.2*elementary_charge" />
                        <!-- divalent lone pair: pyrimidine, TIP4P, TIP5P -->
                        <!-- The atoms 2-1-3 define the X-Y plane, with Z perpendicular. If outOfPlaneAngle is 0, the
                        charge site is a specified distance along the in-plane vector which bisects the angle left by taking
                        360 degrees minus angle(2,1,3). If outOfPlaneAngle is nonzero, the charge sites lie out of the plane
                        by the specified angle (at the specified distance) and their in-plane projection lines along the angle
                        's bisector. -->
                        <VirtualSite type="DivalentLonePair" smirks="[*:2]~[#7X2:1]~[*:3]" distance="0.30*angstrom"_
                        outOfPlaneAngle="0.0*degree" chargeincrement1="+0.1*elementary_charge" />
                            <!-- trivalent nitrogen lone pair -->
                            <!-- charge sites lie above and below the nitrogen at specified distances from the nitrogen, along
                            the vector perpendicular to the plane of (2,3,4) that passes through the nitrogen. If the nitrogen is
                            co-planar with the connected atom, charge sites are simply above and below the plane-->
                                <!-- Positive and negative values refer to above or below the nitrogen as measured relative to the
                                plane of (2,3,4), i.e. below the nitrogen means nearer the 2,3,4 plane unless they are co-planar -->
                                    <VirtualSite type="TrivalentLonePair" smirks="[*:2]~[#7X3:1](~[*:4])~[*:3]" distance="0.30*angstrom"
                                    chargeincrement1="+0.1*elementary_charge"/>
                                    <VirtualSite type="TrivalentLonePair" smirks="[*:2]~[#7X3:1](~[*:4])~[*:3]" distance="-0.30*angstrom"
                                    chargeincrement1="+0.1*elementary_charge"/>
</VirtualSites>
```

## Aromaticity models

Before conducting SMIRKS substructure searches, molecules are prepared using one of the supported aromaticity models, which must be specified with the `aromaticity_model` attribute. The only aromaticity model currently widely supported (by both the [OpenEye toolkit](#) and [RDKit](#)) is the `OEArModel_MDL` model.

## Additional plans for future development

See the [openforcefield GitHub issue tracker](#) to propose changes to this specification, or read through proposed changes currently being discussed.

### 1.3.11 The openforcefield reference implementation

A Python reference implementation of a parameterization engine implementing the SMIRNOFF force field specification can be found [online](#). This implementation can use either the free-for-academics (but commercially supported) [OpenEye toolkit](#) or the free and open source [RDKit cheminformatics toolkit](#). See the [installation instructions](#) for information on how to install this implementation and its dependencies.

#### Examples

A relatively extensive set of examples is made available on the [reference implementation repository](#) under `examples/`.

#### Parameterizing a system

Consider parameterizing a simple system containing a the drug imatinib.

```
# Create a molecule from a mol2 file
from openforcefield.topology import Molecule
molecule = Molecule.from_file('imatinib.mol2')

# Create a Topology specifying the system to be parameterized containing just the molecule
topology = molecule.to_topology()

# Load the smirnoff99Frosst forcefield
from openforcefield.typing.engines import smirnoff
forcefield = smirnoff.ForceField('test_forcefields/smirkoff99Frosst.offxml')

# Create an OpenMM System from the topology
system = forcefield.create_openmm_system(topology)
```

See `examples/SMIRNOFF_simulation/` for an extension of this example illustrating to simulate this molecule in the gas phase.

The topology object provided to `create_openmm_system()` can contain any number of molecules of different types, including biopolymers, ions, buffer molecules, or solvent molecules. The openforcefield toolkit provides a number of convenient methods for importing or constructing topologies given PDB files, Sybyl mol2 files, SDF files, SMILES strings, and IUPAC names; see the [toolkit documentation](#) for more information. Notably, this topology object differs from those found in [OpenMM](#) or [MDTraj](#) in that it contains information on the *chemical identity* of the molecules constituting the system, rather than this atomic elements and covalent connectivity; this additional chemical information is required for the [direct chemical perception](#) features of SMIRNOFF typing.

#### Using SMIRNOFF small molecule forcefields with traditional biopolymer force fields

While SMIRNOFF format force fields can cover a wide range of biological systems, our initial focus is on general small molecule force fields, meaning that users may have considerable interest in combining SMIRNOFF small molecule parameters to systems in combination with traditional biopolymer parameters from conventional force fields, such as the AMBER family of protein/nucleic acid force fields. Thus, we provide an example of setting up a mixed protein-ligand system in `examples/mixedFF_structure`, where an AMBER family force field is used for a protein and smirnoff99Frosst for a small molecule.

### The optional `id` and `parent_id` attributes and other XML attributes

In general, additional optional XML attributes can be specified and will be ignored by ForceField unless they are specifically handled by the parser (and specified in this document).

One attribute we have found helpful in parameter file development is the `id` attribute for a specific parameter line, and we *recommend* that SMIRNOFF force fields utilize this as effectively a parameter serial number, such as in:

```
<Bond smirks="[#6X3:1]-[#6X3:2]" id="b5" k="820.0*kilocalorie_per_mole/angstrom**2" length="1.  
→45*angstrom"/>
```

Some functionality in ForceField, such as `ForceField.label_molecules`, looks for the `id` attribute. Without this attribute, there is no way to uniquely identify a specific parameter line in the XML file without referring to it by its smirks string, and since some smirks strings can become long and relatively unwieldy (especially for torsions) this provides a more human- and search-friendly way of referring to specific sets of parameters.

The `parent_id` attribute is also frequently used to denote parameters from which the current parameter is derived in some manner.

### A remark about parameter availability

ForceField will currently raise an exception if any parameters are missing where expected for your system—i.e. if a bond is assigned no parameters, an exception will be raised. However, use of generic parameters (i.e. `[*:1]~[*:2]` for a bond) in your `.offxml` will result in parameters being assigned everywhere, bypassing this exception. We recommend generics be used sparingly unless it is your intention to provide true universal generic parameters.

## 1.3.12 Version history

### 0.3

This is a backwards-incompatible update to the SMIRNOFF 0.2 draft specification. However, the Open Force Field Toolkit version accompanying this update is capable of converting 0.1 spec SMIRNOFF data to 0.2 spec, and subsequently 0.2 spec to 0.3 spec. The 0.1-to-0.2 spec conversion makes a number of assumptions about settings such as long-range nonbonded handling. Warnings are printed about each assumption that is made during this spec conversion. No mechanism to convert backwards in spec is provided.

Key changes in this version of the spec are:

- Section headers now contain individual versions, instead of relying on the `<SMIRNOFF>`-level tag.
- Section headers no longer contain `X_unit` attributes.
- All physical quantities are now written as expressions containing the appropriate units.
- The default potential for `<ProperTorsions>` and `<ImproperTorsions>` was changed from `charmm` to `k*(1+cos(periodicity*theta-phase))`, as CHARMM interprets torsion terms with periodicity 0 as having a quadratic potential, while the Open Force Field Toolkit would interpret a zero periodicity literally.

### 0.2

This is a backwards-incompatible overhaul of the SMIRNOFF 0.1 draft specification along with ForceField implementation refactor:

- Aromaticity model now defaults to `OEAroModel_MDL`, and aromaticity model names drop OpenEye-specific prefixes
- Top-level tags are now required to specify units for any unit-bearing quantities to avoid the potential for mistakes from implied units.
- Potential energy component definitions were renamed to be more general:
  - `<NonbondedForce>` was renamed to `<vdW>`
  - `<HarmonicBondForce>` was renamed to `<Bonds>`
  - `<HarmonicAngleForce>` was renamed to `<Angles>`
  - `<BondChargeCorrections>` was renamed to `<ChargeIncrementModel>` and generalized to accommodate an arbitrary number of tagged atoms
  - `<GBSAForce>` was renamed to `<GBSA>`
- `<PeriodicTorsionForce>` was split into `<ProperTorsions>` and `<ImproperTorsions>`
- `<vdW>` now specifies 1-2, 1-3, 1-4, and 1-5 scaling factors via `scale12` (default: 0), `scale13` (default: 0), `scale14` (default: 0.5), and `scale15` (default 1.0) attributes. It also specifies the long-range vdW method to use, currently supporting cutoff (default) and PME. Coulomb scaling parameters have been removed from `StericsForce`.
- Added the `<Electrostatics>` tag to separately specify 1-2, 1-3, 1-4, and 1-5 scaling factors for electrostatics, as well as the method used to compute electrostatics (PME, reaction-field, Coulomb) since this has a huge effect on the energetics of the system.
- Made it clear that `<Constraint>` entries do not have to be between bonded atoms.
- `<VirtualSites>` has been added, and the specification of charge increments harmonized with `<ChargeIncrementModel>`
- The `potential` attribute was added to most forces to allow flexibility in extending forces to additional functional forms (or algebraic expressions) in the future. `potential` defaults to the current recommended scheme if omitted.
- `<GBSA>` now has defaults specified for `gb_method` and `sa_method`
- Changes to how fractional bond orders are handled:
  - Use of fractional bond order is now specified at the force tag level, rather than the root level
  - The fractional bond order method is specified via the `fractional_bondorder_method` attribute
  - The fractional bond order interpolation scheme is specified via the `fractional_bondorder_interpolation`
- Section heading names were cleaned up.
- Example was updated to reflect use of the new `openforcefield.topology.Topology` class
- Eliminated “Requirements” section, since it specified requirements for the software, rather than described an aspect of the SMIRNOFF specification
- Fractional bond orders are described in `<Bonds>`, since they currently only apply to this term.

## 0.1

Initial draft specification.

## 1.4 Examples using SMIRNOFF with the toolkit

The following examples are available in [the openforcefield toolkit repository](#):

### 1.4.1 Index of provided examples

- `SMIRNOFF_simulation` - simulation of a molecule in the gas phase with the SMIRNOFF forcefield format
- `forcefield_modification` - modify forcefield parameters and evaluate how system energy changes
- `using_smirnoff_in_amber_or_gromacs` - convert a System generated with the Open Forcefield Toolkit, which can be simulated natively with OpenMM, into AMBER prmtop/inpcrd and GROMACS top/gro input files through the ParmEd library.
- `inspect_assigned_parameters` - check which parameters are used in which molecules and generate parameter usage statistics.
- `using_smirnoff_with_amber_protein_forcefield` - use SMIRNOFF parameters for small molecules in combination with more conventional force fields for proteins and other components of your system (using ParmEd to combine parameterized structures)

## 1.5 Developing for the toolkit

### 1.5.1 Style guide

Development for the openforcefield toolkit conforms to the recommendations given by the [Software Development Best Practices for Computational Chemistry](#) guide.

The naming conventions of classes, functions, and variables follows [PEP8](#), consistently with the best practices guide. The naming conventions used in this library not covered by PEP8 are: - Use `file_path`, `file_name`, and `file_stem` to indicate path/to/stem.extension, stem.extension, and stem respectively, consistently with the variables in the standard `pathlib` library. - Use `n_x` to abbreviate “number of X” (e.g. `n_atoms`, `n_molecules`).

### 1.5.2 Contributing

We always welcome [GitHub pull requests](#). For bug fixes, major feature additions, or refactoring, please raise an issue on the [GitHub issue tracker](#) first to ensure the design will be amenable to current developer plans.

### 1.5.3 How can I become a developer?

If you would like to contribute, please post an issue on the [GitHub issue tracker](#) describing the contribution you would like to make to start a discussion.

## 1.6 Frequently asked questions (FAQ)

### 1.6.1 Input files for applying SMIRNOFF parameters

SMIRNOFF force fields use direct chemical perception meaning that, unlike many molecular mechanics (MM) force fields, they apply parameters based on substructure searches acting directly on molecules. This creates unique opportunities and allows them to encode a great deal of chemistry quite simply, but it also means that the *starting point* for parameter assignment must be well-defined chemically, giving not just the elements and connectivity for all of the atoms of all of the components of your system, but also providing the formal charges and bond orders.

Specifically, to apply SMIRNOFF to a system, you must either:

1. Provide Open Force Field Toolkit Molecule objects corresponding to the components of your system, or
2. Provide an OpenMM Topology which includes bond orders and thus can be converted to molecules corresponding to the components of your system

Without this information, our direct chemical perception cannot be applied to your molecule, as it requires the chemical identity of the molecules in your system – that is, bond order and formal charge as well as atoms and connectivity. Unless you provide the full chemical identity in this sense, we must attempt to guess or infer the chemical identity of your molecules, which is a recipe for trouble. Different molecules can have the same chemical graph but differ in bond order and formal charge, or different resonance structures may be treated rather differently by some force fields (e.g. c1cc(ccc1c2cc[nH+]cc2)[O-] vs C1=CC(C=CC1=C2C=CNC=C2)=O, where the central bond is rotatable in one resonance structure but not in the other) even though they have identical formal charge and connectivity (chemical graph). A force field which uses the chemical identity of molecules to assign parameters needs to know the exact chemical identity of the molecule you are intending to parameterize.

### 1.6.2 Can I use an AMBER (or GROMACS) topology/coordinate file as a starting point for applying a SMIRNOFF force field?

In a word, “no”.

Parameter files used by typical molecular dynamics simulation packages do not currently encode enough information to identify the molecules chemically present, or at least not without drawing inferences. For example, one could take a structure file and infer bond orders based on bond lengths, or attempt to infer bond orders from force constants in a parameter file. Such inference work is outside the scope of SMIRNOFF.

If you have such an inference problem, we recommend that you use pre-existing cheminformatics tools available elsewhere (such as via the OpenEye toolkits, such as the `OEPceiveBondOrders` functionality offered there) to solve this problem and identify your molecules before beginning your work with SMIRNOFF.

### 1.6.3 What about starting from a PDB file?

PDB files do not in general provide the chemical identity of small molecules contained therein, and thus do not provide suitable starting points for applying SMIRNOFF to small molecules. This is especially problematic for PDB files from X-ray crystallography which typically do not include proteins, making the problem even worse. For our purposes here, however, we assume you begin with the coordinates of all atoms present and the full topology of your system.

Given a PDB file of a hypothetical biomolecular system of interest containing a small molecule, there are several routes available to you for treating the small molecule present:

- Use a cheminformatics toolkit (see above) to infer bond orders

- Identify your ligand from a database; e.g. if it is in the Protein Data Bank (PDB), it will be present in the [Ligand Expo](#) meaning that it has a database entry and code you can use to look up its putative chemical identity
- Identify your ligand by name or SMILES string (or similar) from the literature or your collaborators

#### 1.6.4 What do you recommend as a starting point?

For application of SMIRNOFF force fields, we recommend that you begin your work with formats which provide the chemical identity of your small molecule (including formal charge and bond order). This means we recommend one of the following or equivalent:

- A .mol2 file or files for the molecules comprising your system, with correct bond orders and formal charges. (Note: Do NOT generate this from a simulation package or tool which does not have access to bond order information; you may end up with a .mol2 file, but the bond orders will be incorrect)
- Isomeric SMILES strings for the components of your system
- InCHI strings for the components of your system
- Chemical Identity Registry numbers for the components of your system
- IUPAC names for the components of your system

Essentially, anything which provides the full identity of what you want to simulate (including stereochemistry) should work, though it may require more or less work to get it into an acceptable format.

## API DOCUMENTATION

### 2.1 Molecular topology representations

This module provides pure-Python classes for representing molecules and molecular systems. These classes offer several advantages over corresponding Topology objects in [OpenMM](#) and [MDTraj](#), including offering serialization to a variety of standard formats (including [XML](#), [JSON](#), [YAML](#), [BSON](#), [TOML](#), and [MessagePack](#)).

#### 2.1.1 Primary objects

<code>FrozenMolecule</code>	Immutable chemical representation of a molecule, such as a small molecule or biopolymer.
<code>Molecule</code>	Mutable chemical representation of a molecule, such as a small molecule or biopolymer.
<code>Topology</code>	A Topology is a chemical representation of a system containing one or more molecules appearing in a specified order.

#### `openforcefield.topology.FrozenMolecule`

```
class openforcefield.topology.FrozenMolecule(other=None, file_format=None,
                                             toolkit_registry=<openforcefield.utils.toolkits.ToolkitRegistry object>, allow_undefined_stereo=False)
```

Immutable chemical representation of a molecule, such as a small molecule or biopolymer.

#### Examples

Create a molecule from a sdf file

```
>>> from openforcefield.utils import get_data_file_path
>>> sdf_filepath = get_data_file_path('molecules/ethanol.sdf')
>>> molecule = FrozenMolecule.from_file(sdf_filepath)
```

Convert to OpenEye OEMol object

```
>>> oemol = molecule.to_openeye()
```

Create a molecule from an OpenEye molecule

```
>>> molecule = FrozenMolecule.from_openeye(oemol)
```

Convert to RDKit Mol object

```
>>> rdmol = molecule.to_rdkit()
```

Create a molecule from an RDKit molecule

```
>>> molecule = FrozenMolecule.from_rdkit(rdmol)
```

Create a molecule from IUPAC name (requires the OpenEye toolkit)

```
>>> molecule = FrozenMolecule.from_iupac('imatinib')
```

Create a molecule from SMILES

```
>>> molecule = FrozenMolecule.from_smiles('Cc1ccccc1')
```

**Warning:** This API is experimental and subject to change.

## Attributes

`angles` Get an iterator over all i-j-k angles.

`atoms` Iterate over all Atom objects.

`bonds` Iterate over all Bond objects.

`conformers` Iterate over all conformers in this molecule.

`impropers` Iterate over all proper torsions in the molecule

`n_angles` int: number of angles in the Molecule.

`n_atoms` The number of Atom objects.

`n_bonds` The number of Bond objects.

`n_conformers` Iterate over all Atom objects.

`n_impropers` int: number of improper torsions in the Molecule.

`n_particles` The number of Particle objects, which corresponds to how many positions must be used.

`n_proper` int: number of proper torsions in the Molecule.

`n_virtual_sites` The number of VirtualSite objects.

`name` The name (or title) of the molecule

`partial_charges` Returns the partial charges (if present) on the molecule

`particles` Iterate over all Particle objects.

`proper` Iterate over all proper torsions in the molecule

`properties` The properties dictionary of the molecule

`torsions` Get an iterator over all i-j-k-l torsions.

`total_charge` Return the total charge on the molecule

`virtual_sites` Iterate over all VirtualSite objects.

## Methods

<code>chemical_environment_matches(self[, ...])</code>	<code>query[, ...]</code>	Retrieve all matches for a given chemical environment query.
<code>compute_partial_charges(self[, toolkit_registry])</code>		<b>Warning! Not Implemented!</b> Calculate partial atomic charges for this molecule using an underlying toolkit
<code>compute_partial_charges_am1bcc(self[, ...])</code>		Calculate partial atomic charges for this molecule using AM1-BCC run by an underlying toolkit
<code>compute_wiberg_bond_orders(self[, ...])</code>		Calculate wiberg bond orders for this molecule using an underlying toolkit
<code>from_bson(serialized)</code>		Instantiate an object from a BSON serialized representation.
<code>from_dict(molecule_dict)</code>		Create a new Molecule from a dictionary representation
<code>from_file(file_path[, file_format, ...])</code>		Create one or more molecules from a file
<code>from_iupac(iupac_name, \*\*kwargs)</code>		Generate a molecule from IUPAC or common name
<code>from_json(serialized)</code>		Instantiate an object from a JSON serialized representation.
<code>from_messagepack(serialized)</code>		Instantiate an object from a MessagePack serialized representation.
<code>from_openeye(oemol[, low_undefined_stereo])</code>	<code>al-</code>	Create a Molecule from an OpenEye molecule.
<code>from_pickle(serialized)</code>		Instantiate an object from a pickle serialized representation.
<code>from_rdkit(rdmol[, allow_undefined_stereo])</code>		Create a Molecule from an RDKit molecule.
<code>from_smiles(smiles[, ...])</code>		Construct a Molecule from a SMILES representation
<code>from_toml(serialized)</code>		Instantiate an object from a TOML serialized representation.
<code>from_topology(topology)</code>		Return a Molecule representation of an open-forcefield Topology containing a single Molecule object.
<code>from_xml(serialized)</code>		Instantiate an object from an XML serialized representation.
<code>from_yaml(serialized)</code>		Instantiate from a YAML serialized representation.
<code>generate_conformers(self[, ...])</code>		Generate conformers for this molecule using an underlying toolkit
<code>get_bond_between(self, i, j)</code>		Returns the bond between two atoms
<code>get_fractional_bond_orders(self[, ...])</code>	<code>method,</code> <code>[...])</code>	Get fractional bond orders.
<code>is_isomorphic(self, other[, ...])</code>		Determines whether the molecules are isomorphic by comparing their graphs.
<code>to_bson(self)</code>		Return a BSON serialized representation.
<code>to_dict(self)</code>		Return a dictionary representation of the molecule.

Continued on next page

Table 2 – continued from previous page

<code>to_file(self, file_path, file_format[, ...])</code>	Write the current molecule to a file or file-like object
<code>to_iupac(self)</code>	Generate IUPAC name from Molecule
<code>to_json(self[, indent])</code>	Return a JSON serialized representation.
<code>to_messagepack(self)</code>	Return a MessagePack representation.
<code>to_networkx(self)</code>	Generate a NetworkX undirected graph from the Molecule.
<code>to_openeye(self[, aromaticity_model])</code>	Create an OpenEye molecule
<code>to_pickle(self)</code>	Return a pickle serialized representation.
<code>to_rdkit(self[, aromaticity_model])</code>	Create an RDKit molecule
<code>to_smiles(self[, toolkit_registry])</code>	Return a canonical isomeric SMILES representation of the current molecule
<code>to_toml(self)</code>	Return a TOML serialized representation.
<code>to_topology(self)</code>	Return an openforcefield Topology representation containing one copy of this molecule
<code>to_xml(self[, indent])</code>	Return an XML representation.
<code>to_yaml(self)</code>	Return a YAML serialized representation.

`__init__(self, other=None, file_format=None, toolkit_registry=<openforcefield.utils.toolkits.ToolkitRegistry object at 0x7fba569d3128>, allow_undefined_stereo=False)`  
Create a new FrozenMolecule object

### Parameters

**other** [optional, default=None] If specified, attempt to construct a copy of the Molecule from the specified object. This can be any one of the following:

- a `Molecule` object
- a file that can be used to construct a `Molecule` object
- an `openeye.oecchem.OEMol`
- an `rdkit.Chem.rdcchem.Mol`
- a serialized `Molecule` object

**file\_format** [str, optional, default=None] If providing a file-like object, you must specify the format of the data. If providing a file, the file format will attempt to be guessed from the suffix.

**toolkit\_registry** [a `ToolkitRegistry` or `ToolkitWrapper` object, optional, default=GLOBAL\_TOOLKIT\_REGISTRY] `ToolkitRegistry` or `ToolkitWrapper` to use for I/O operations

**allow\_undefined\_stereo** [bool, default=False] If loaded from a file and `False`, raises an exception if undefined stereochemistry is detected during the molecule's construction.

### Examples

Create an empty molecule:

```
>>> empty_molecule = FrozenMolecule()
```

Create a molecule from a file that can be used to construct a molecule, using either a filename or file-like object:

```
>>> from openforcefield.utils import get_data_file_path
>>> sdf_filepath = get_data_file_path('molecules/ethanol.sdf')
>>> molecule = FrozenMolecule(sdf_filepath)
>>> molecule = FrozenMolecule(open(sdf_filepath, 'r'), file_format='sdf')
```

```
>>> import gzip
>>> mol2_gz_filepath = get_data_file_path('molecules/toluene.mol2.gz')
>>> molecule = FrozenMolecule(gzip.GzipFile(mol2_gz_filepath, 'r'), file_format='mol2')
```

Create a molecule from another molecule:

```
>>> molecule_copy = FrozenMolecule(molecule)
```

Convert to OpenEye OEMol object

```
>>> oemol = molecule.to_openeye()
```

Create a molecule from an OpenEye molecule:

```
>>> molecule = FrozenMolecule(oemol)
```

Convert to RDKit Mol object

```
>>> rdmol = molecule.to_rdkit()
```

Create a molecule from an RDKit molecule:

```
>>> molecule = FrozenMolecule(rdmol)
```

Create a molecule from a serialized molecule object:

```
>>> serialized_molecule = molecule.__getstate__()
>>> molecule_copy = Molecule(serialized_molecule)
```

## Methods

<code>__init__(self[, other, file_format, ...])</code>	Create a new FrozenMolecule object
<code>chemical_environment_matches(self, query[, ...])</code>	Retrieve all matches for a given chemical environment query.
<code>compute_partial_charges(self[, toolkit_registry])</code>	<b>Warning! Not Implemented!</b> Calculate partial atomic charges for this molecule using an underlying toolkit
<code>compute_partial_charges_am1bcc(self[, ...])</code>	Calculate partial atomic charges for this molecule using AM1-BCC run by an underlying toolkit
<code>compute_wiberg_bond_orders(self[, ...])</code>	Calculate wiberg bond orders for this molecule using an underlying toolkit
<code>from_bson(serialized)</code>	Instantiate an object from a BSON serialized representation.
<code>from_dict(molecule_dict)</code>	Create a new Molecule from a dictionary representation
<code>from_file(file_path[, file_format, ...])</code>	Create one or more molecules from a file

Continued on next page

Table 3 – continued from previous page

<code>from_iupac(iupac_name, \*\*kwargs)</code>	Generate a molecule from IUPAC or common name
<code>from_json(serialized)</code>	Instantiate an object from a JSON serialized representation.
<code>from_messagepack(serialized)</code>	Instantiate an object from a MessagePack serialized representation.
<code>from_openeye(oemol[, low_undefined_stereo])</code>	Create a Molecule from an OpenEye molecule.
<code>from_pickle(serialized)</code>	Instantiate an object from a pickle serialized representation.
<code>from_rdkit(rdmol[, allow_undefined_stereo])</code>	Create a Molecule from an RDKit molecule.
<code>from_smiles(smiles[, ...])</code>	Construct a Molecule from a SMILES representation
<code>from_toml(serialized)</code>	Instantiate an object from a TOML serialized representation.
<code>from_topology(topology)</code>	Return a Molecule representation of an openforcefield Topology containing a single Molecule object.
<code>from_xml(serialized)</code>	Instantiate an object from an XML serialized representation.
<code>from_yaml(serialized)</code>	Instantiate from a YAML serialized representation.
<code>generate_conformers(self[, ...])</code>	Generate conformers for this molecule using an underlying toolkit
<code>get_bond_between(self, i, j)</code>	Returns the bond between two atoms
<code>get_fractional_bond_orders(self[, method, ...])</code>	Get fractional bond orders.
<code>is_isomorphic(self, other[, ...])</code>	Determines whether the molecules are isomorphic by comparing their graphs.
<code>to bson(self)</code>	Return a BSON serialized representation.
<code>to_dict(self)</code>	Return a dictionary representation of the molecule.
<code>to_file(self, file_path, file_format[, ...])</code>	Write the current molecule to a file or file-like object
<code>to_iupac(self)</code>	Generate IUPAC name from Molecule
<code>to_json(self[, indent])</code>	Return a JSON serialized representation.
<code>to_messagepack(self)</code>	Return a MessagePack representation.
<code>to_networkx(self)</code>	Generate a NetworkX undirected graph from the Molecule.
<code>to_openeye(self[, aromaticity_model])</code>	Create an OpenEye molecule
<code>to_pickle(self)</code>	Return a pickle serialized representation.
<code>to_rdkit(self[, aromaticity_model])</code>	Create an RDKit molecule
<code>to_smiles(self[, toolkit_registry])</code>	Return a canonical isomeric SMILES representation of the current molecule
<code>to_toml(self)</code>	Return a TOML serialized representation.
<code>to_topology(self)</code>	Return an openforcefield Topology representation containing one copy of this molecule
<code>to_xml(self[, indent])</code>	Return an XML representation.
<code>to_yaml(self)</code>	Return a YAML serialized representation.

## Attributes

angles	Get an iterator over all i-j-k angles.
atoms	Iterate over all Atom objects.
bonds	Iterate over all Bond objects.
conformers	Iterate over all conformers in this molecule.
impropers	Iterate over all proper torsions in the molecule
n_angles	int: number of angles in the Molecule.
n_atoms	The number of Atom objects.
n_bonds	The number of Bond objects.
n_conformers	Iterate over all Atom objects.
n_impropers	int: number of improper torsions in the Molecule.
n_particles	The number of Particle objects, which corresponds to how many positions must be used.
n_propers	int: number of proper torsions in the Molecule.
n_virtual_sites	The number of VirtualSite objects.
name	The name (or title) of the molecule
partial_charges	Returns the partial charges (if present) on the molecule
particles	Iterate over all Particle objects.
propers	Iterate over all proper torsions in the molecule
properties	The properties dictionary of the molecule
torsions	Get an iterator over all i-j-k-l torsions.
total_charge	Return the total charge on the molecule
virtual_sites	Iterate over all VirtualSite objects.

**to\_dict(self)**

Return a dictionary representation of the molecule.

**Returns**

**molecule\_dict** [OrderedDict] A dictionary representation of the molecule.

**classmethod from\_dict(molecule\_dict)**

Create a new Molecule from a dictionary representation

**Parameters**

**molecule\_dict** [OrderedDict] A dictionary representation of the molecule.

**Returns**

**molecule** [Molecule] A Molecule created from the dictionary representation

**to\_smiles(self, toolkit\_registry=<openforcefield.utils.toolkits.ToolkitRegistry object at 0x7fb4569d3128>)**

Return a canonical isomeric SMILES representation of the current molecule

---

**Note:** RDKit and OpenEye versions will not necessarily return the same representation.

---

**Parameters**

**toolkit\_registry** [openforcefield.utils.toolkits.ToolRegistry or openforcefield.utils.toolkits.ToolkitWrapper, optional, default=None] ToolkitRegistry

or ToolkitWrapper to use for SMILES conversion

#### Returns

**smiles** [str] Canonical isomeric explicit-hydrogen SMILES

#### Examples

```
>>> from openforcefield.utils import get_data_file_path
>>> sdf_filepath = get_data_file_path('molecules/ethanol.sdf')
>>> molecule = Molecule(sdf_filepath)
>>> smiles = molecule.to_smiles()
```

**static from\_smiles**(smiles, hydrogens\_are\_explicit=False, toolkit\_registry=<openforcefield.utils.toolkits.ToolkitRegistry object at 0x7fba569d3128>)

Construct a Molecule from a SMILES representation

#### Parameters

**smiles** [str] The SMILES representation of the molecule.

**hydrogens\_are\_explicit** [bool, default = False] If False, the cheminformatics toolkit will perform hydrogen addition

**toolkit\_registry** [openforcefield.utils.toolkits.ToolRegistry or openforcefield.utils.toolkits.ToolkitWrapper, optional, default=None] ToolkitRegistry or ToolkitWrapper to use for SMILES-to-molecule conversion

#### Returns

**molecule** [openforcefield.topology.Molecule]

#### Examples

```
>>> molecule = Molecule.from_smiles('Cc1ccccc1')
```

**is\_isomorphic**(self, other, compare\_atom\_stereochemistry=True, compare\_bond\_stereochemistry=True)

Determines whether the molecules are isomorphic by comparing their graphs.

#### Parameters

**other** [an openforcefield.topology.molecule.FrozenMolecule] The molecule to test for isomorphism.

**compare\_atom\_stereochemistry** [bool, optional] If False, atoms' stereochemistry is ignored for the purpose of determining equality. Default is True.

**compare\_bond\_stereochemistry** [bool, optional] If False, bonds' stereochemistry is ignored for the purpose of determining equality. Default is True.

#### Returns

**molecules\_are\_isomorphic** [bool]

**generate\_conformers**(self, toolkit\_registry=<openforcefield.utils.toolkits.ToolkitRegistry object at 0x7fba569d3128>, n\_conformers=10, clear\_existing=True)

Generate conformers for this molecule using an underlying toolkit

#### Parameters

**toolkit\_registry** [openforcefield.utils.toolkits.ToolRegistry or openforcefield.utils.toolkits.ToolkitWrapper, optional, default=None] ToolkitRegistry or ToolkitWrapper to use for SMILES-to-molecule conversion

**n\_conformers** [int, default=1] The maximum number of conformers to produce

**clear\_existing** [bool, default=True] Whether to overwrite existing conformers for the molecule

#### Raises

**InvalidToolkitError** If an invalid object is passed as the toolkit\_registry parameter

#### Examples

```
>>> molecule = Molecule.from_smiles('CCCCC')
>>> molecule.generate_conformers()
```

**compute\_partial\_charges\_am1bcc(self, toolkit\_registry=<openforcefield.utils.toolkits.ToolkitRegistry object at 0x7fba569d3128>)**

Calculate partial atomic charges for this molecule using AM1-BCC run by an underlying toolkit

#### Parameters

**toolkit\_registry** [openforcefield.utils.toolkits.ToolRegistry or openforcefield.utils.toolkits.ToolkitWrapper, optional, default=None] ToolkitRegistry or ToolkitWrapper to use for the calculation

#### Raises

**InvalidToolkitError** If an invalid object is passed as the toolkit\_registry parameter

#### Examples

```
>>> molecule = Molecule.from_smiles('CCCCC')
>>> molecule.generate_conformers()
>>> molecule.compute_partial_charges_am1bcc()
```

**compute\_partial\_charges(self, toolkit\_registry=<openforcefield.utils.toolkits.ToolkitRegistry object at 0x7fba569d3128>)**

**Warning! Not Implemented!** Calculate partial atomic charges for this molecule using an underlying toolkit

#### Parameters

**quantum\_chemical\_method** [string, default='AM1-BCC'] The quantum chemical method to use for partial charge calculation.

**partial\_charge\_method** [string, default='None'] The partial charge calculation method to use for partial charge calculation.

**toolkit\_registry** [openforcefield.utils.toolkits.ToolRegistry or openforcefield.utils.toolkits.ToolkitWrapper, optional, default=None] ToolkitRegistry or ToolkitWrapper to use for SMILES-to-molecule conversion

#### Raises

**InvalidToolkitError** If an invalid object is passed as the toolkit\_registry parameter

## Examples

```
>>> molecule = Molecule.from_smiles('CCCCC')
>>> molecule.generate_conformers()
```

**compute\_wiberg\_bond\_orders**(*self*, *charge\_model*=None, *toolkit\_registry*=<*openforcefield.utils.toolkits.ToolkitRegistry* object at 0x7fba569d3128>)

Calculate wiberg bond orders for this molecule using an underlying toolkit

### Parameters

**toolkit\_registry** [*openforcefield.utils.toolkits.ToolRegistry* or *openforcefield.utils.toolkits.ToolkitWrapper*, optional, default=None] *ToolkitRegistry* or *ToolkitWrapper* to use for SMILES-to-molecule conversion

**charge\_model** [string, optional] The charge model to use for partial charge calculation

### Raises

**InvalidToolkitError** If an invalid object is passed as the *toolkit\_registry* parameter

## Examples

```
>>> molecule = Molecule.from_smiles('CCCCC')
>>> molecule.generate_conformers()
>>> molecule.compute_wiberg_bond_orders()
```

**to\_networkx**(*self*)

Generate a NetworkX undirected graph from the Molecule.

Nodes are Atoms labeled with particle indices and atomic elements (via the *element* node attribute). Edges denote chemical bonds between Atoms. Virtual sites are not included, since they lack a concept of chemical connectivity.

### Returns

**graph** [networkx.Graph] The resulting graph, with nodes (atoms) labeled with atom indices, elements, stereochemistry and aromaticity flags and bonds with two atom indices, bond order, stereochemistry, and aromaticity flags

## Examples

Retrieve the bond graph for imatinib (OpenEye toolkit required)

```
>>> molecule = Molecule.from_iupac('imatinib')
>>> nxgraph = molecule.to_networkx()
```

**property partial\_charges**

Returns the partial charges (if present) on the molecule

### Returns

**partial\_charges** [a simtk.unit.Quantity - wrapped numpy array [1 x n\_atoms]] The partial charges on this Molecule's atoms.

**property n\_particles**

The number of Particle objects, which corresponds to how many positions must be used.

**property n\_atoms**

The number of Atom objects.

**property n\_virtual\_sites**

The number of VirtualSite objects.

**property n\_bonds**

The number of Bond objects.

**property n\_angles**

int: number of angles in the Molecule.

**property n\_proper\_torsions**

int: number of proper torsions in the Molecule.

**property n\_improper\_torsions**

int: number of improper torsions in the Molecule.

**property particles**

Iterate over all Particle objects.

**property atoms**

Iterate over all Atom objects.

**property conformers**

Iterate over all conformers in this molecule.

**property n\_conformers**

Iterate over all Atom objects.

**property virtual\_sites**

Iterate over all VirtualSite objects.

**property bonds**

Iterate over all Bond objects.

**property angles**

Get an iterator over all i-j-k angles.

**property torsions**

Get an iterator over all i-j-k-l torsions. Note that i-j-k-i torsions (cycles) are excluded.

**Returns**

**torsions** [iterable of 4-Atom tuples]

**property proper\_torsions**

Iterate over all proper torsions in the molecule

**property improper\_torsions**

Iterate over all improper torsions in the molecule

**property total\_charge**

Return the total charge on the molecule

**property name**

The name (or title) of the molecule

**property properties**

The properties dictionary of the molecule

**chemical\_environment\_matches(self, query, toolkit\_registry=<openforcefield.utils.toolkits.ToolkitRegistry object at 0x7fba569d3128>)**

Retrieve all matches for a given chemical environment query.

## Parameters

**query** [str or ChemicalEnvironment] SMARTS string (with one or more tagged atoms) or ChemicalEnvironment query Query will internally be resolved to SMIRKS using query.asSMIRKS() if it has an .asSMIRKS method.

**toolkit\_registry** [openforcefield.utils.toolkits.ToolRegistry or openforcefield.utils.toolkits.ToolkitWrapper, optional, default=GLOBAL\_TOOLKIT\_REGISTRY] ToolkitRegistry or ToolkitWrapper to use for chemical environment matches

## Returns

**matches** [list of atom index tuples] A list of tuples, containing the indices of the matching atoms.

## Examples

Retrieve all the carbon-carbon bond matches in a molecule

```
>>> molecule = Molecule.from_iupac('imatinib')
>>> matches = molecule.chemical_environment_matches('[#6X3:1]~[#6X3:2]')
```

**classmethod from\_iupac(iupac\_name, \*\*kwargs)**

Generate a molecule from IUPAC or common name

## Parameters

**iupac\_name** [str] IUPAC name of molecule to be generated

**allow\_undefined\_stereo** [bool, default=False] If false, raises an exception if molecule contains undefined stereochemistry.

## Returns

**molecule** [Molecule] The resulting molecule with position

---

**Note:** This method requires the OpenEye toolkit to be installed. ..

---

## Examples

Create a molecule from a common name

```
>>> molecule = Molecule.from_iupac('4-[(4-methylpiperazin-1-yl)methyl]-N-(4-methyl-3-[(4-  
→(pyridin-3-yl)pyrimidin-2-yl]amino}phenyl)benzamide')
```

Create a molecule from a common name

```
>>> molecule = Molecule.from_iupac('imatinib')
```

**to\_iupac(self)**

Generate IUPAC name from Molecule

## Returns

**iupac\_name** [str] IUPAC name of the molecule

---

**Note:** This method requires the OpenEye toolkit to be installed. ..

---

## Examples

```
>>> from openforcefield.utils import get_data_file_path
>>> sdf_filepath = get_data_file_path('molecules/ethanol.sdf')
>>> molecule = Molecule(sdf_filepath)
>>> iupac_name = molecule.to_iupac()
```

### **static from\_topology(topology)**

Return a Molecule representation of an openforcefield Topology containing a single Molecule object.

#### Parameters

**topology** [openforcefield.topology.Topology] The `Topology` object containing a single `Molecule` object. Note that OpenMM and MDTraj Topology objects are not supported.

#### Returns

**molecule** [openforcefield.topology.Molecule] The Molecule object in the topology

#### Raises

`ValueError` If the topology does not contain exactly one molecule.

## Examples

Create a molecule from a Topology object that contains exactly one molecule

```
>>> molecule = Molecule.from_topology(topology) # doctest: +SKIP
```

### **to\_topology(self)**

Return an openforcefield Topology representation containing one copy of this molecule

#### Returns

**topology** [openforcefield.topology.Topology] A Topology representation of this molecule

## Examples

```
>>> molecule = Molecule.from_iupac('imatinib')
>>> topology = molecule.to_topology()
```

### **static from\_file(file\_path, file\_format=None, toolkit\_registry=<openforcefield.utils.toolkits.ToolkitRegistry object at 0x7fba569d3128>, allow\_undefined\_stereo=False)**

Create one or more molecules from a file

#### Parameters

**file\_path** [str or file-like object] The path to the file or file-like object to stream one or more molecules from.

**file\_format** [str, optional, default=None] Format specifier, usually file suffix (eg. ‘MOL2’, ‘SMI’) Note that not all toolkits support all formats. Check ToolkitWrapper.toolkit\_file\_read\_formats for your loaded toolkits for details.

**toolkit\_registry** [openforcefield.utils.toolkits.ToolRegistry or openforcefield.utils.toolkits.ToolkitWrapper,]

**optional, default=GLOBAL\_TOOLKIT\_REGISTRY** ToolkitRegistry or ToolkitWrapper to use for file loading. If a Toolkit is passed, only the highest-precedence toolkit is used

**allow\_undefined\_stereo** [bool, default=False] If false, raises an exception if oemol contains undefined stereochemistry.

### Returns

**molecules** [Molecule or list of Molecules] If there is a single molecule in the file, a Molecule is returned; otherwise, a list of Molecule objects is returned.

### Examples

```
>>> from openforcefield.tests.utils import get_monomer_mol2_file_path
>>> mol2_file_path = get_monomer_mol2_file_path('cyclohexane')
>>> molecule = Molecule.from_file(mol2_file_path)
```

**to\_file(self, file\_path, file\_format, toolkit\_registry=<openforcefield.utils.toolkits.ToolkitRegistry object at 0x7fba569d3128>)**

Write the current molecule to a file or file-like object

### Parameters

**file\_path** [str or file-like object] A file-like object or the path to the file to be written.

**file\_format** [str] Format specifier, one of ['MOL2', 'MOL2H', 'SDF', 'PDB', 'SMI', 'CAN', 'TDT'] Note that not all toolkits support all formats

**toolkit\_registry** [openforcefield.utils.toolkits.ToolRegistry or openforcefield.utils.toolkits.ToolkitWrapper,]

**optional, default=GLOBAL\_TOOLKIT\_REGISTRY** ToolkitRegistry or ToolkitWrapper to use for file writing. If a Toolkit is passed, only the highest-precedence toolkit is used

### Raises

**ValueError** If the requested file\_format is not supported by one of the installed cheminformatics toolkits

### Examples

```
>>> molecule = Molecule.from_iupac('imatinib')
>>> molecule.to_file('imatinib.mol2', file_format='mol2') # doctest: +SKIP
>>> molecule.to_file('imatinib.sdf', file_format='sdf') # doctest: +SKIP
>>> molecule.to_file('imatinib.pdb', file_format='pdb') # doctest: +SKIP
```

**static from\_rdkit(rdmol, allow\_undefined\_stereo=False)**

Create a Molecule from an RDKit molecule.

Requires the RDKit to be installed.

**Parameters**

**rdmol** [rkit.RDMol] An RDKit molecule

**allow\_undefined\_stereo** [bool, default=False] If false, raises an exception if oemol contains undefined stereochemistry.

**Returns**

**molecule** [openforcefield.Molecule] An openforcefield molecule

**Examples**

Create a molecule from an RDKit molecule

```
>>> from rdkit import Chem
>>> from openforcefield.tests.utils import get_data_file_path
>>> rdmol = Chem.MolFromMolFile(get_data_file_path('systems/monomers/ethanol.sdf'))
>>> molecule = Molecule.from_rdkit(rdmol)
```

**to\_rdkit(self, aromaticity\_model='OEArroModel\_MDL')**

Create an RDKit molecule

Requires the RDKit to be installed.

**Parameters**

**aromaticity\_model** [str, optional, default=DEFAULT\_AROMATICITY\_MODEL] The aromaticity model to use

**Returns**

**rdmol** [rkit.RDMol] An RDKit molecule

**Examples**

Convert a molecule to RDKit

```
>>> from openforcefield.utils import get_data_file_path
>>> sdf_filepath = get_data_file_path('molecules/ethanol.sdf')
>>> molecule = Molecule(sdf_filepath)
>>> rdmol = molecule.to_rdkit()
```

**static from\_openeye(oemol, allow\_undefined\_stereo=False)**

Create a Molecule from an OpenEye molecule.

Requires the OpenEye toolkit to be installed.

**Parameters**

**oemol** [openeye.oechem.OEMol] An OpenEye molecule

**allow\_undefined\_stereo** [bool, default=False] If false, raises an exception if oemol contains undefined stereochemistry.

**Returns**

**molecule** [openforcefield.topology.Molecule] An openforcefield molecule

## Examples

Create a Molecule from an OpenEye OEMol

```
>>> from openeye import oechem
>>> from openforcefield.tests.utils import get_data_file_path
>>> ifs = oechem.oemolistream(get_data_file_path('systems/monomers/ethanol.mol2'))
>>> oemols = list(ifs.GetOEGraphMols())
>>> molecule = Molecule.from_openeye(oemols[0])
```

**to\_openeye(self, aromaticity\_model='OEArroModel\_MDL')**

Create an OpenEye molecule

Requires the OpenEye toolkit to be installed.

### Parameters

**aromaticity\_model** [str, optional, default=DEFAULT\_AROMATICITY\_MODEL] The aromaticity model to use

### Returns

**oemol** [openeye.oechem.OEMol] An OpenEye molecule

## Examples

Create an OpenEye molecule from a Molecule

```
>>> molecule = Molecule.from_smiles('CC')
>>> oemol = molecule.to_openeye()
```

**get\_fractional\_bond\_orders(self, method='Wiberg', toolkit\_registry=<openforcefield.utils.toolkits.ToolkitRegistry object at 0x7fba569d3128>)**

Get fractional bond orders.

**method** [str, optional, default='Wiberg'] The name of the charge method to use. Options are: \* 'Wiberg' : Wiberg bond order

**toolkit\_registry** [openforcefield.utils.toolkits ToolkitRegistry] The toolkit registry to use for molecule operations

## Examples

Get fractional Wiberg bond orders

```
>>> molecule = Molecule.from_iupac('imatinib')
>>> molecule.generate_conformers()
>>> fractional_bond_orders = molecule.get_fractional_bond_orders(method='Wiberg')
```

**classmethod from\_bson(serialized)**

Instantiate an object from a BSON serialized representation.

Specification: <http://bsonspec.org/>

### Parameters

**serialized** [bytes] A BSON serialized representation of the object

### Returns

**instance** [cls] An instantiated object

**classmethod from\_json(*serialized*)**  
Instantiate an object from a JSON serialized representation.  
Specification: <https://www.json.org/>

**Parameters**  
**serialized** [str] A JSON serialized representation of the object

**Returns**  
**instance** [cls] An instantiated object

**classmethod from\_messagepack(*serialized*)**  
Instantiate an object from a MessagePack serialized representation.  
Specification: <https://msgpack.org/index.html>

**Parameters**  
**serialized** [bytes] A MessagePack-encoded bytes serialized representation

**Returns**  
**instance** [cls] Instantiated object.

**classmethod from\_pickle(*serialized*)**  
Instantiate an object from a pickle serialized representation.

**Warning:** This is not recommended for safe, stable storage since the pickle specification may change between Python versions.

**Parameters**  
**serialized** [str] A pickled representation of the object

**Returns**  
**instance** [cls] An instantiated object

**classmethod from\_toml(*serialized*)**  
Instantiate an object from a TOML serialized representation.  
Specification: <https://github.com/toml-lang/toml>

**Parameters**  
**serialized** [str] A TOML serialized representation of the object

**Returns**  
**instance** [cls] An instantiated object

**classmethod from\_xml(*serialized*)**  
Instantiate an object from an XML serialized representation.  
Specification: <https://www.w3.org/XML/>

**Parameters**  
**serialized** [bytes] An XML serialized representation

**Returns**

**instance** [cls] Instantiated object.

**classmethod from\_yaml(serialized)**

Instantiate from a YAML serialized representation.

Specification: <http://yaml.org/>

#### Parameters

**serialized** [str] A YAML serialized representation of the object

#### Returns

**instance** [cls] Instantiated object

**to\_bson(self)**

Return a BSON serialized representation.

Specification: <http://bsonspec.org/>

#### Returns

**serialized** [bytes] A BSON serialized representation of the object

**to\_json(self, indent=None)**

Return a JSON serialized representation.

Specification: <https://www.json.org/>

#### Parameters

**indent** [int, optional, default=None] If not None, will pretty-print with specified number of spaces for indentation

#### Returns

**serialized** [str] A JSON serialized representation of the object

**to\_messagepack(self)**

Return a MessagePack representation.

Specification: <https://msgpack.org/index.html>

#### Returns

**serialized** [bytes] A MessagePack-encoded bytes serialized representation of the object

**to\_pickle(self)**

Return a pickle serialized representation.

**Warning:** This is not recommended for safe, stable storage since the pickle specification may change between Python versions.

#### Returns

**serialized** [str] A pickled representation of the object

**to\_toml(self)**

Return a TOML serialized representation.

Specification: <https://github.com/toml-lang/toml>

#### Returns

**serialized** [str] A TOML serialized representation of the object

**to\_xml(self, indent=2)**  
Return an XML representation.  
Specification: <https://www.w3.org/XML/>

**Parameters**

**indent** [int, optional, default=2] If not None, will pretty-print with specified number of spaces for indentation

**Returns**

**serialized** [bytes] A MessagePack-encoded bytes serialized representation.

**to\_yaml(self)**  
Return a YAML serialized representation.  
Specification: <http://yaml.org/>

**Parameters**

**serialized** [str] A YAML serialized representation of the object

**get\_bond\_between(self, i, j)**  
Returns the bond between two atoms

**Parameters**

**i, j** [int or Atom] Atoms or atom indices to check

**Returns**

**bond** [Bond] The bond between i and j.

## openforcefield.topology.Molecule

**class** openforcefield.topology.Molecule(\*args, \*\*kwargs)  
Mutable chemical representation of a molecule, such as a small molecule or biopolymer.

### Examples

Create a molecule from an sdf file

```
>>> from openforcefield.utils import get_data_file_path
>>> sdf_filepath = get_data_file_path('molecules/ethanol.sdf')
>>> molecule = Molecule(sdf_filepath)
```

Convert to OpenEye OEMol object

```
>>> oemol = molecule.to_openeye()
```

Create a molecule from an OpenEye molecule

```
>>> molecule = Molecule.from_openeye(oemol)
```

Convert to RDKit Mol object

```
>>> rdmol = molecule.to_rdkit()
```

Create a molecule from an RDKit molecule

```
>>> molecule = Molecule.from_rdkit(rdmol)
```

Create a molecule from IUPAC name (requires the OpenEye toolkit)

```
>>> molecule = Molecule.from_iupac('imatinib')
```

Create a molecule from SMILES

```
>>> molecule = Molecule.from_smiles('Cc1ccccc1')
```

**Warning:** This API is experimental and subject to change.

## Attributes

`angles` Get an iterator over all i-j-k angles.

`atoms` Iterate over all Atom objects.

`bonds` Iterate over all Bond objects.

`conformers` Iterate over all conformers in this molecule.

`impropers` Iterate over all proper torsions in the molecule

`n_angles` int: number of angles in the Molecule.

`n_atoms` The number of Atom objects.

`n_bonds` The number of Bond objects.

`n_conformers` Iterate over all Atom objects.

`n_impropers` int: number of improper torsions in the Molecule.

`n_particles` The number of Particle objects, which corresponds to how many positions must be used.

`n_proper_torsions` int: number of proper torsions in the Molecule.

`n_virtual_sites` The number of VirtualSite objects.

`name` The name (or title) of the molecule

`partial_charges` Returns the partial charges (if present) on the molecule

`particles` Iterate over all Particle objects.

`proper_torsions` Iterate over all proper torsions in the molecule

`properties` The properties dictionary of the molecule

`torsions` Get an iterator over all i-j-k-l torsions.

`total_charge` Return the total charge on the molecule

`virtual_sites` Iterate over all VirtualSite objects.

## Methods

<code>add_atom(self, atomic_number, formal_charge, ...)</code>	Add an atom
<code>add_bond(self, atom1, atom2, bond_order, ...)</code>	Add a bond between two specified atom indices
<code>add_bond_charge_virtual_site(self, atoms, ...)</code>	Create a bond charge-type virtual site, in which the location of the charge is specified by the position of two atoms.
<code>add_conformer(self, coordinates)</code>	# TODO: Should this not be public? Adds a conformer of the molecule
<code>add_divalent_lone_pair_virtual_site(self, ...)</code>	Create a divalent lone pair-type virtual site, in which the location of the charge is specified by the position of three atoms.
<code>add_monovalent_lone_pair_virtual_site(self, ...)</code>	Create a bond charge-type virtual site, in which the location of the charge is specified by the position of three atoms.
<code>add_trivalent_lone_pair_virtual_site(self, ...)</code>	Create a trivalent lone pair-type virtual site, in which the location of the charge is specified by the position of four atoms.
<code>chemical_environment_matches(self, query[, ...])</code>	Retrieve all matches for a given chemical environment query.
<code>compute_partial_charges(self[, toolkit_registry])</code>	<b>Warning! Not Implemented!</b> Calculate partial atomic charges for this molecule using an underlying toolkit
<code>compute_partial_charges_am1bcc(self[, ...])</code>	Calculate partial atomic charges for this molecule using AM1-BCC run by an underlying toolkit
<code>compute_wiberg_bond_orders(self[, ...])</code>	Calculate wiberg bond orders for this molecule using an underlying toolkit
<code>from bson(serialized)</code>	Instantiate an object from a BSON serialized representation.
<code>from_dict(molecule_dict)</code>	Create a new Molecule from a dictionary representation
<code>from_file(file_path[, file_format, ...])</code>	Create one or more molecules from a file
<code>from_iupac(iupac_name, \*\*kwargs)</code>	Generate a molecule from IUPAC or common name
<code>from_json(serialized)</code>	Instantiate an object from a JSON serialized representation.
<code>from_messagepack(serialized)</code>	Instantiate an object from a MessagePack serialized representation.
<code>from_openeye(oemol[, low_undefined_stereo])</code>	Create a Molecule from an OpenEye molecule.
<code>from_pickle(serialized)</code>	Instantiate an object from a pickle serialized representation.
<code>from_rdkit(rdmol[, allow_undefined_stereo])</code>	Create a Molecule from an RDKit molecule.
<code>from_smiles(smiles[, ...])</code>	Construct a Molecule from a SMILES representation
<code>from_toml(serialized)</code>	Instantiate an object from a TOML serialized representation.
<code>from_topology(topology)</code>	Return a Molecule representation of an open-forcefield Topology containing a single Molecule object.

Continued on next page

Table 5 – continued from previous page

<code>from_xml(serialized)</code>	Instantiate an object from an XML serialized representation.
<code>from_yaml(serialized)</code>	Instantiate from a YAML serialized representation.
<code>generate_conformers(self[, ...])</code>	Generate conformers for this molecule using an underlying toolkit
<code>get_bond_between(self, i, j)</code>	Returns the bond between two atoms
<code>get_fractional_bond_orders(self[, method, ...])</code>	Get fractional bond orders.
<code>is_isomorphic(self, other[, ...])</code>	Determines whether the molecules are isomorphic by comparing their graphs.
<code>to_bson(self)</code>	Return a BSON serialized representation.
<code>to_dict(self)</code>	Return a dictionary representation of the molecule.
<code>to_file(self, file_path, file_format[, ...])</code>	Write the current molecule to a file or file-like object
<code>to_iupac(self)</code>	Generate IUPAC name from Molecule
<code>to_json(self[, indent])</code>	Return a JSON serialized representation.
<code>to_messagepack(self)</code>	Return a MessagePack representation.
<code>to_networkx(self)</code>	Generate a NetworkX undirected graph from the Molecule.
<code>to_openeye(self[, aromaticity_model])</code>	Create an OpenEye molecule
<code>to_pickle(self)</code>	Return a pickle serialized representation.
<code>to_rdkit(self[, aromaticity_model])</code>	Create an RDKit molecule
<code>to_smiles(self[, toolkit_registry])</code>	Return a canonical isomeric SMILES representation of the current molecule
<code>to_toml(self)</code>	Return a TOML serialized representation.
<code>to_topology(self)</code>	Return an openforcefield Topology representation containing one copy of this molecule
<code>to_xml(self[, indent])</code>	Return an XML representation.
<code>to_yaml(self)</code>	Return a YAML serialized representation.

`__init__(self, *args, **kwargs)`  
Create a new Molecule object

#### Parameters

`other` [optional, default=None] If specified, attempt to construct a copy of the Molecule from the specified object. This can be any one of the following:

- a `Molecule` object
- a file that can be used to construct a `Molecule` object
- an `openeye.oechem.OEMol`
- an `rdkit.Chem.rdcchem.Mol`
- a serialized `Molecule` object

#### Examples

Create an empty molecule:

```
>>> empty_molecule = Molecule()
```

Create a molecule from a file that can be used to construct a molecule, using either a filename or file-like object:

```
>>> from openforcefield.utils import get_data_file_path
>>> sdf_filepath = get_data_file_path('molecules/ethanol.sdf')
>>> molecule = Molecule(sdf_filepath)
>>> molecule = Molecule(open(sdf_filepath, 'r')), file_format='sdf')
```

```
>>> import gzip
>>> mol2_gz_filepath = get_data_file_path('molecules/toluene.mol2.gz')
>>> molecule = Molecule(gzip.GzipFile(mol2_gz_filepath, 'r')), file_format='mol2')
```

Create a molecule from another molecule:

```
>>> molecule_copy = Molecule(molecule)
```

Convert to OpenEye OEMol object

```
>>> oemol = molecule.to_openeye()
```

Create a molecule from an OpenEye molecule:

```
>>> molecule = Molecule(oemol)
```

Convert to RDKit Mol object

```
>>> rdmol = molecule.to_rdkit()
```

Create a molecule from an RDKit molecule:

```
>>> molecule = Molecule(rdmol)
```

Create a molecule from a serialized molecule object:

```
>>> serialized_molecule = molecule.__getstate__()
>>> molecule_copy = Molecule(serialized_molecule)
```

## Methods

<code>__init__(self, *args, **kwargs)</code>	Create a new Molecule object
<code>add_atom(self, atomic_number, formal_charge, ...)</code>	Add an atom
<code>add_bond(self, atom1, atom2, bond_order, ...)</code>	Add a bond between two specified atom indices
<code>add_bond_charge_virtual_site(self, atoms, ...)</code>	Create a bond charge-type virtual site, in which the location of the charge is specified by the position of two atoms.
<code>add_conformer(self, coordinates)</code>	# TODO: Should this not be public? Adds a conformer of the molecule
<code>add_divalent_lone_pair_virtual_site(self, ...)</code>	Create a divalent lone pair-type virtual site, in which the location of the charge is specified by the position of three atoms.

Continued on next page

Table 6 – continued from previous page

<code>add_monovalent_lone_pair_virtual_site(self, ...)</code>	Create a bond charge-type virtual site, in which the location of the charge is specified by the position of three atoms.
<code>add_trivalent_lone_pair_virtual_site(self, ...)</code>	Create a trivalent lone pair-type virtual site, in which the location of the charge is specified by the position of four atoms.
<code>chemical_environment_matches(self, query[, ...])</code>	Retrieve all matches for a given chemical environment query.
<code>compute_partial_charges(self[, toolkit_registry])</code>	<b>Warning! Not Implemented!</b> Calculate partial atomic charges for this molecule using an underlying toolkit
<code>compute_partial_charges_am1bcc(self[, ...])</code>	Calculate partial atomic charges for this molecule using AM1-BCC run by an underlying toolkit
<code>compute_wiberg_bond_orders(self[, ...])</code>	Calculate wiberg bond orders for this molecule using an underlying toolkit
<code>from_bson(serialized)</code>	Instantiate an object from a BSON serialized representation.
<code>from_dict(molecule_dict)</code>	Create a new Molecule from a dictionary representation
<code>from_file(file_path[, file_format, ...])</code>	Create one or more molecules from a file
<code>from_iupac(iupac_name, \*\*kwargs)</code>	Generate a molecule from IUPAC or common name
<code>from_json(serialized)</code>	Instantiate an object from a JSON serialized representation.
<code>from_messagepack(serialized)</code>	Instantiate an object from a MessagePack serialized representation.
<code>from_openeye(oemol[, low_undefined_stereo])</code>	Create a Molecule from an OpenEye molecule.
<code>from_pickle(serialized)</code>	Instantiate an object from a pickle serialized representation.
<code>from_rdkit(rdmol[, allow_undefined_stereo])</code>	Create a Molecule from an RDKit molecule.
<code>from_smiles(smiles[, ...])</code>	Construct a Molecule from a SMILES representation
<code>from_toml(serialized)</code>	Instantiate an object from a TOML serialized representation.
<code>from_topology(topology)</code>	Return a Molecule representation of an open-forcefield Topology containing a single Molecule object.
<code>from_xml(serialized)</code>	Instantiate an object from an XML serialized representation.
<code>from_yaml(serialized)</code>	Instantiate from a YAML serialized representation.
<code>generate_conformers(self[, ...])</code>	Generate conformers for this molecule using an underlying toolkit
<code>get_bond_between(self, i, j)</code>	Returns the bond between two atoms
<code>get_fractional_bond_orders(self[, method, ...])</code>	Get fractional bond orders.
<code>is_isomorphic(self, other[, ...])</code>	Determines whether the molecules are isomorphic by comparing their graphs.
<code>to_bson(self)</code>	Return a BSON serialized representation.

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Table 6 – continued from previous page

<code>to_dict(self)</code>	Return a dictionary representation of the molecule.
<code>to_file(self, file_path, file_format[, ...])</code>	Write the current molecule to a file or file-like object
<code>to_iupac(self)</code>	Generate IUPAC name from Molecule
<code>to_json(self[, indent])</code>	Return a JSON serialized representation.
<code>to_messagepack(self)</code>	Return a MessagePack representation.
<code>to_networkx(self)</code>	Generate a NetworkX undirected graph from the Molecule.
<code>to_openeye(self[, aromaticity_model])</code>	Create an OpenEye molecule
<code>to_pickle(self)</code>	Return a pickle serialized representation.
<code>to_rdkit(self[, aromaticity_model])</code>	Create an RDKit molecule
<code>to_smiles(self[, toolkit_registry])</code>	Return a canonical isomeric SMILES representation of the current molecule
<code>to_toml(self)</code>	Return a TOML serialized representation.
<code>to_topology(self)</code>	Return an openforcefield Topology representation containing one copy of this molecule
<code>to_xml(self[, indent])</code>	Return an XML representation.
<code>to_yaml(self)</code>	Return a YAML serialized representation.

## Attributes

<code>angles</code>	Get an iterator over all i-j-k angles.
<code>atoms</code>	Iterate over all Atom objects.
<code>bonds</code>	Iterate over all Bond objects.
<code>conformers</code>	Iterate over all conformers in this molecule.
<code>impropers</code>	Iterate over all proper torsions in the molecule
<code>n_angles</code>	int: number of angles in the Molecule.
<code>n_atoms</code>	The number of Atom objects.
<code>n_bonds</code>	The number of Bond objects.
<code>n_conformers</code>	Iterate over all Atom objects.
<code>n_impropers</code>	int: number of improper torsions in the Molecule.
<code>n_particles</code>	The number of Particle objects, which corresponds to how many positions must be used.
<code>n_propers</code>	int: number of proper torsions in the Molecule.
<code>n_virtual_sites</code>	The number of VirtualSite objects.
<code>name</code>	The name (or title) of the molecule
<code>partial_charges</code>	Returns the partial charges (if present) on the molecule
<code>particles</code>	Iterate over all Particle objects.
<code>propers</code>	Iterate over all proper torsions in the molecule
<code>properties</code>	The properties dictionary of the molecule
<code>torsions</code>	Get an iterator over all i-j-k-l torsions.
<code>total_charge</code>	Return the total charge on the molecule
<code>virtual_sites</code>	Iterate over all VirtualSite objects.

### property angles

Get an iterator over all i-j-k angles.

### property atoms

Iterate over all Atom objects.

**property bonds**

Iterate over all Bond objects.

**chemical\_environment\_matches(self, query, toolkit\_registry=<openforcefield.utils.toolkits.ToolkitRegistry object at 0x7fba569d3128>)**

Retrieve all matches for a given chemical environment query.

**Parameters**

**query** [str or ChemicalEnvironment] SMARTS string (with one or more tagged atoms) or ChemicalEnvironment query Query will internally be resolved to SMIRKS using query.asSMIRKS() if it has an .asSMIRKS method.

**toolkit\_registry** [openforcefield.utils.toolkits.ToolRegistry or openforcefield.utils.toolkits.ToolkitWrapper, optional, default=GLOBAL\_TOOLKIT\_REGISTRY] ToolkitRegistry or ToolkitWrapper to use for chemical environment matches

**Returns**

**matches** [list of atom index tuples] A list of tuples, containing the indices of the matching atoms.

**Examples**

Retrieve all the carbon-carbon bond matches in a molecule

```
>>> molecule = Molecule.from_iupac('imatitinib')
>>> matches = molecule.chemical_environment_matches('[#6X3:1]~[#6X3:2]')
```

**compute\_partial\_charges(self, toolkit\_registry=<openforcefield.utils.toolkits.ToolkitRegistry object at 0x7fba569d3128>)**

**Warning! Not Implemented!** Calculate partial atomic charges for this molecule using an underlying toolkit

**Parameters**

**quantum\_chemical\_method** [string, default='AM1-BCC'] The quantum chemical method to use for partial charge calculation.

**partial\_charge\_method** [string, default='None'] The partial charge calculation method to use for partial charge calculation.

**toolkit\_registry** [openforcefield.utils.toolkits.ToolRegistry or openforcefield.utils.toolkits.ToolkitWrapper, optional, default=None] ToolkitRegistry or ToolkitWrapper to use for SMILES-to-molecule conversion

**Raises**

**InvalidToolkitError** If an invalid object is passed as the toolkit\_registry parameter

**Examples**

```
>>> molecule = Molecule.from_smiles('CCCCC')
>>> molecule.generate_conformers()
```

---

**compute\_partial\_charges\_am1bcc(self, toolkit\_registry=<openforcefield.utils.toolkits.ToolkitRegistry object at 0x7fba569d3128>)**

Calculate partial atomic charges for this molecule using AM1-BCC run by an underlying toolkit

#### Parameters

**toolkit\_registry** [openforcefield.utils.toolkits.ToolRegistry or openforcefield.utils.toolkits.ToolkitWrapper, optional, default=None] ToolkitRegistry or ToolkitWrapper to use for the calculation

#### Raises

**InvalidToolkitError** If an invalid object is passed as the toolkit\_registry parameter

#### Examples

```
>>> molecule = Molecule.from_smiles('CCCCC')
>>> molecule.generate_conformers()
>>> molecule.compute_partial_charges_am1bcc()
```

**compute\_wiberg\_bond\_orders(self, charge\_model=None, toolkit\_registry=<openforcefield.utils.toolkits.ToolkitRegistry object at 0x7fba569d3128>)**

Calculate wiberg bond orders for this molecule using an underlying toolkit

#### Parameters

**toolkit\_registry** [openforcefield.utils.toolkits.ToolRegistry or openforcefield.utils.toolkits.ToolkitWrapper, optional, default=None] ToolkitRegistry or ToolkitWrapper to use for SMILES-to-molecule conversion

**charge\_model** [string, optional] The charge model to use for partial charge calculation

#### Raises

**InvalidToolkitError** If an invalid object is passed as the toolkit\_registry parameter

#### Examples

```
>>> molecule = Molecule.from_smiles('CCCCC')
>>> molecule.generate_conformers()
>>> molecule.compute_wiberg_bond_orders()
```

#### property conformers

Iterate over all conformers in this molecule.

#### classmethod from\_bson(serialized)

Instantiate an object from a BSON serialized representation.

Specification: <http://bsonspec.org/>

#### Parameters

**serialized** [bytes] A BSON serialized representation of the object

#### Returns

**instance** [cls] An instantiated object

#### classmethod from\_dict(molecule\_dict)

Create a new Molecule from a dictionary representation

## Parameters

**molecule\_dict** [OrderedDict] A dictionary representation of the molecule.

## Returns

**molecule** [Molecule] A Molecule created from the dictionary representation

**static from\_file(file\_path, file\_format=None, toolkit\_registry=<openforcefield.utils.toolkits.ToolkitRegistry object at 0x7fba569d3128>, allow\_undefined\_stereo=False)**

Create one or more molecules from a file

## Parameters

**file\_path** [str or file-like object] The path to the file or file-like object to stream one or more molecules from.

**file\_format** [str, optional, default=None] Format specifier, usually file suffix (eg. ‘MOL2’, ‘SMI’) Note that not all toolkits support all formats. Check ToolkitWrapper.toolkit\_file\_read\_formats for your loaded toolkits for details.

**toolkit\_registry** [openforcefield.utils.toolkits.ToolRegistry or openforcefield.utils.toolkits.ToolkitWrapper,]

**optional, default=GLOBAL\_TOOLKIT\_REGISTRY** ToolkitRegistry or ToolkitWrapper to use for file loading. If a Toolkit is passed, only the highest-precedence toolkit is used

**allow\_undefined\_stereo** [bool, default=False] If false, raises an exception if oemol contains undefined stereochemistry.

## Returns

**molecules** [Molecule or list of Molecules] If there is a single molecule in the file, a Molecule is returned; otherwise, a list of Molecule objects is returned.

## Examples

```
>>> from openforcefield.tests.utils import get_monomer_mol2_file_path
>>> mol2_file_path = get_monomer_mol2_file_path('cyclohexane')
>>> molecule = Molecule.from_file(mol2_file_path)
```

**classmethod from\_iupac(iupac\_name, \*\*kwargs)**  
Generate a molecule from IUPAC or common name

## Parameters

**iupac\_name** [str] IUPAC name of molecule to be generated

**allow\_undefined\_stereo** [bool, default=False] If false, raises an exception if molecule contains undefined stereochemistry.

## Returns

**molecule** [Molecule] The resulting molecule with position

---

**Note:** This method requires the OpenEye toolkit to be installed. ..

---

## Examples

Create a molecule from a common name

```
>>> molecule = Molecule.from_iupac('4-[(4-methylpiperazin-1-yl)methyl]-N-(4-methyl-3-[(4-pyridin-3-yl)pyrimidin-2-yl]amino}phenyl)benzamide')
```

Create a molecule from a common name

```
>>> molecule = Molecule.from_iupac('imatinib')
```

### `classmethod from_json(serialized)`

Instantiate an object from a JSON serialized representation.

Specification: <https://www.json.org/>

#### Parameters

`serialized` [str] A JSON serialized representation of the object

#### Returns

`instance` [cls] An instantiated object

### `classmethod from_messagepack(serialized)`

Instantiate an object from a MessagePack serialized representation.

Specification: <https://msgpack.org/index.html>

#### Parameters

`serialized` [bytes] A MessagePack-encoded bytes serialized representation

#### Returns

`instance` [cls] Instantiated object.

### `static from_openeye(oemol, allow_undefined_stereo=False)`

Create a Molecule from an OpenEye molecule.

Requires the OpenEye toolkit to be installed.

#### Parameters

`oemol` [openeye.oecore.OEMol] An OpenEye molecule

`allow_undefined_stereo` [bool, default=False] If false, raises an exception if oemol contains undefined stereochemistry.

#### Returns

`molecule` [openforcefield.topology.Molecule] An openforcefield molecule

## Examples

Create a Molecule from an OpenEye OEMol

```
>>> from openeye import oechem
>>> from openforcefield.tests.utils import get_data_file_path
>>> ifs = oechem.oemolistream(get_data_file_path('systems/monomers/ethanol.mol2'))
>>> oemols = list(ifs.GetOEGraphMols())
>>> molecule = Molecule.from_openeye(oemols[0])
```

```
classmethod from_pickle(serialized)
```

Instantiate an object from a pickle serialized representation.

**Warning:** This is not recommended for safe, stable storage since the pickle specification may change between Python versions.

#### Parameters

**serialized** [str] A pickled representation of the object

#### Returns

**instance** [cls] An instantiated object

```
static from_rdkit(rdmol, allow_undefined_stereo=False)
```

Create a Molecule from an RDKit molecule.

Requires the RDKit to be installed.

#### Parameters

**rdmol** [rkit.RDMol] An RDKit molecule

**allow\_undefined\_stereo** [bool, default=False] If false, raises an exception if oemol contains undefined stereochemistry.

#### Returns

**molecule** [openforcefield.Molecule] An openforcefield molecule

## Examples

Create a molecule from an RDKit molecule

```
>>> from rdkit import Chem
>>> from openforcefield.tests.utils import get_data_file_path
>>> rdmol = Chem.MolFromMolFile(get_data_file_path('systems/monomers/ethanol.sdf'))
>>> molecule = Molecule.from_rdkit(rdmol)
```

```
static from_smiles(smiles, hydrogens_are_explicit=False, toolkit_registry=<openforcefield.utils.toolkits.ToolkitRegistry object at 0x7fba569d3128>)
```

Construct a Molecule from a SMILES representation

#### Parameters

**smiles** [str] The SMILES representation of the molecule.

**hydrogens\_are\_explicit** [bool, default = False] If False, the cheminformatics toolkit will perform hydrogen addition

**toolkit\_registry** [openforcefield.utils.toolkits.ToolRegistry or openforcefield.utils.toolkits.ToolkitWrapper, optional, default=None] ToolkitRegistry or ToolkitWrapper to use for SMILES-to-molecule conversion

#### Returns

**molecule** [openforcefield.topology.Molecule]

## Examples

```
>>> molecule = Molecule.from_smiles('Cc1ccccc1')
```

**classmethod from\_toml(serialized)**

Instantiate an object from a TOML serialized representation.

Specification: <https://github.com/toml-lang/toml>

**Parameters**

**serialized** [str] A TOML serialized representation of the object

**Returns**

**instance** [cls] An instantiated object

**static from\_topology(topology)**

Return a Molecule representation of an openforcefield Topology containing a single Molecule object.

**Parameters**

**topology** [openforcefield.topology.Topology] The `Topology` object containing a single `Molecule` object. Note that OpenMM and MDTraj Topology objects are not supported.

**Returns**

**molecule** [openforcefield.topology.Molecule] The Molecule object in the topology

**Raises**

**ValueError** If the topology does not contain exactly one molecule.

## Examples

Create a molecule from a Topology object that contains exactly one molecule

```
>>> molecule = Molecule.from_topology(topology) # doctest: +SKIP
```

**classmethod from\_xml(serialized)**

Instantiate an object from an XML serialized representation.

Specification: <https://www.w3.org/XML/>

**Parameters**

**serialized** [bytes] An XML serialized representation

**Returns**

**instance** [cls] Instantiated object.

**classmethod from\_yaml(serialized)**

Instantiate from a YAML serialized representation.

Specification: <http://yaml.org/>

**Parameters**

**serialized** [str] A YAML serialized representation of the object

**Returns**

**instance** [cls] Instantiated object

**generate\_conformers**(*self*, *toolkit\_registry*=<*openforcefield.utils.toolkits.ToolkitRegistry* object at 0x7fba569d3128>, *n\_conformers*=10, *clear\_existing*=True)  
Generate conformers for this molecule using an underlying toolkit

#### Parameters

**toolkit\_registry** [*openforcefield.utils.toolkits.ToolRegistry* or *openforcefield.utils.toolkits.ToolkitWrapper*, optional, default=None] *ToolkitRegistry* or *ToolkitWrapper* to use for SMILES-to-molecule conversion

**n\_conformers** [int, default=1] The maximum number of conformers to produce

**clear\_existing** [bool, default=True] Whether to overwrite existing conformers for the molecule

#### Raises

**InvalidToolkitError** If an invalid object is passed as the *toolkit\_registry* parameter

### Examples

```
>>> molecule = Molecule.from_smiles('CCCCC')
>>> molecule.generate_conformers()
```

**get\_bond\_between**(*self*, *i*, *j*)

Returns the bond between two atoms

#### Parameters

**i, j** [int or Atom] Atoms or atom indices to check

#### Returns

**bond** [Bond] The bond between i and j.

**get\_fractional\_bond\_orders**(*self*, *method*='Wiberg', *toolkit\_registry*=<*openforcefield.utils.toolkits.ToolkitRegistry* object at 0x7fba569d3128>)

Get fractional bond orders.

**method** [str, optional, default='Wiberg'] The name of the charge method to use. Options are: \* 'Wiberg' : Wiberg bond order

**toolkit\_registry** [*openforcefield.utils.toolkits.ToolkitRegistry*] The toolkit registry to use for molecule operations

### Examples

Get fractional Wiberg bond orders

```
>>> molecule = Molecule.from_iupac('imatinib')
>>> molecule.generate_conformers()
>>> fractional_bond_orders = molecule.get_fractional_bond_orders(method='Wiberg')
```

**property impropers**

Iterate over all proper torsions in the molecule

**is\_isomorphic**(*self*, *other*, *compare\_atom\_stereochemistry*=True, *compare\_bond\_stereochemistry*=True)

Determines whether the molecules are isomorphic by comparing their graphs.

## Parameters

**other** [an openforcefield.topology.molecule.FrozenMolecule] The molecule to test for isomorphism.

**compare\_atom\_stereochemistry** [bool, optional] If False, atoms' stereochemistry is ignored for the purpose of determining equality. Default is True.

**compare\_bond\_stereochemistry** [bool, optional] If False, bonds' stereochemistry is ignored for the purpose of determining equality. Default is True.

## Returns

**molecules\_are\_isomorphic** [bool]

**property n\_angles**

int: number of angles in the Molecule.

**property n\_atoms**

The number of Atom objects.

**property n\_bonds**

The number of Bond objects.

**property n\_conformers**

Iterate over all Atom objects.

**property n\_impropers**

int: number of improper torsions in the Molecule.

**property n\_particles**

The number of Particle objects, which corresponds to how many positions must be used.

**property n\_propers**

int: number of proper torsions in the Molecule.

**property n\_virtual\_sites**

The number of VirtualSite objects.

**property name**

The name (or title) of the molecule

**property partial\_charges**

Returns the partial charges (if present) on the molecule

## Returns

**partial\_charges** [a simtk.unit.Quantity - wrapped numpy array [1 x n\_atoms]] The partial charges on this Molecule's atoms.

**property particles**

Iterate over all Particle objects.

**property propers**

Iterate over all proper torsions in the molecule

**property properties**

The properties dictionary of the molecule

**to\_bson(self)**

Return a BSON serialized representation.

Specification: <http://bsonspec.org/>

## Returns

**serialized** [bytes] A BSON serialized representation of the object

**to\_dict(self)**

Return a dictionary representation of the molecule.

**Returns**

**molecule\_dict** [OrderedDict] A dictionary representation of the molecule.

**to\_file(self, file\_path, file\_format, toolkit\_registry=<openforcefield.utils.toolkits.ToolkitRegistry object at 0x7fba569d3128>)**

Write the current molecule to a file or file-like object

**Parameters**

**file\_path** [str or file-like object] A file-like object or the path to the file to be written.

**file\_format** [str] Format specifier, one of ['MOL2', 'MOL2H', 'SDF', 'PDB', 'SMI', 'CAN', 'TDT'] Note that not all toolkits support all formats

**toolkit\_registry** [openforcefield.utils.toolkits.ToolRegistry or openforcefield.utils.toolkits.ToolkitWrapper,]

**optional, default=GLOBAL\_TOOLKIT\_REGISTRY** ToolkitRegistry or ToolkitWrapper to use for file writing. If a Toolkit is passed, only the highest-precedence toolkit is used

**Raises**

**ValueError** If the requested file\_format is not supported by one of the installed cheminformatics toolkits

**Examples**

```
>>> molecule = Molecule.from_iupac('imatinib')
>>> molecule.to_file('imatinib.mol2', file_format='mol2') # doctest: +SKIP
>>> molecule.to_file('imatinib.sdf', file_format='sdf') # doctest: +SKIP
>>> molecule.to_file('imatinib.pdb', file_format='pdb') # doctest: +SKIP
```

**to\_iupac(self)**

Generate IUPAC name from Molecule

**Returns**

**iupac\_name** [str] IUPAC name of the molecule

---

**Note:** This method requires the OpenEye toolkit to be installed. ..

---

**Examples**

```
>>> from openforcefield.utils import get_data_file_path
>>> sdf_filepath = get_data_file_path('molecules/ethanol.sdf')
>>> molecule = Molecule(sdf_filepath)
>>> iupac_name = molecule.to_iupac()
```

**to\_json(self, indent=None)**

Return a JSON serialized representation.

Specification: <https://www.json.org/>

#### Parameters

**indent** [int, optional, default=None] If not None, will pretty-print with specified number of spaces for indentation

#### Returns

**serialized** [str] A JSON serialized representation of the object

#### `to_messagepack(self)`

Return a MessagePack representation.

Specification: <https://msgpack.org/index.html>

#### Returns

**serialized** [bytes] A MessagePack-encoded bytes serialized representation of the object

#### `to_networkx(self)`

Generate a NetworkX undirected graph from the Molecule.

Nodes are Atoms labeled with particle indices and atomic elements (via the `element` node attribute). Edges denote chemical bonds between Atoms. Virtual sites are not included, since they lack a concept of chemical connectivity.

#### Returns

**graph** [networkx.Graph] The resulting graph, with nodes (atoms) labeled with atom indices, elements, stereochemistry and aromaticity flags and bonds with two atom indices, bond order, stereochemistry, and aromaticity flags

### Examples

Retrieve the bond graph for imatinib (OpenEye toolkit required)

```
>>> molecule = Molecule.from_iupac('imatinib')
>>> nxgraph = molecule.to_networkx()
```

#### `to_openeye(self, aromaticity_model='OEArroModel_MDL')`

Create an OpenEye molecule

Requires the OpenEye toolkit to be installed.

#### Parameters

**aromaticity\_model** [str, optional, default=DEFAULT\_AROMATICITY\_MODEL] The aromaticity model to use

#### Returns

**oemol** [openeye.oecore.OEMol] An OpenEye molecule

### Examples

Create an OpenEye molecule from a Molecule

```
>>> molecule = Molecule.from_smiles('CC')
>>> oemol = molecule.to_openeye()
```

**to\_pickle(self)**

Return a pickle serialized representation.

**Warning:** This is not recommended for safe, stable storage since the pickle specification may change between Python versions.

**Returns**

**serialized** [str] A pickled representation of the object

**to\_rdkit(self, aromaticity\_model='OEAroModel\_MDL')**

Create an RDKit molecule

Requires the RDKit to be installed.

**Parameters**

**aromaticity\_model** [str, optional, default=DEFAULT\_AROMATICITY\_MODEL] The aromaticity model to use

**Returns**

**rdmol** [rkit.RDMol] An RDKit molecule

**Examples**

Convert a molecule to RDKit

```
>>> from openforcefield.utils import get_data_file_path
>>> sdf_filepath = get_data_file_path('molecules/ethanol.sdf')
>>> molecule = Molecule(sdf_filepath)
>>> rdmol = molecule.to_rdkit()
```

**to\_smiles(self, toolkit\_registry=<openforcefield.utils.toolkits.ToolkitRegistry object at 0x7fb4569d3128>)**

Return a canonical isomeric SMILES representation of the current molecule

---

**Note:** RDKit and OpenEye versions will not necessarily return the same representation.

---

**Parameters**

**toolkit\_registry** [openforcefield.utils.toolkits.ToolRegistry or openforcefield.utils.toolkits.ToolkitWrapper, optional, default=None] ToolkitRegistry or ToolkitWrapper to use for SMILES conversion

**Returns**

**smiles** [str] Canonical isomeric explicit-hydrogen SMILES

**Examples**

```
>>> from openforcefield.utils import get_data_file_path
>>> sdf_filepath = get_data_file_path('molecules/ethanol.sdf')
>>> molecule = Molecule(sdf_filepath)
>>> smiles = molecule.to_smiles()
```

**to\_toml(self)**

Return a TOML serialized representation.

Specification: <https://github.com/toml-lang/toml>

**Returns**

**serialized** [str] A TOML serialized representation of the object

**to\_topology(self)**

Return an openforcefield Topology representation containing one copy of this molecule

**Returns**

**topology** [openforcefield.topology.Topology] A Topology representation of this molecule

**Examples**

```
>>> molecule = Molecule.from_iupac('imatinib')
>>> topology = molecule.to_topology()
```

**to\_xml(self, indent=2)**

Return an XML representation.

Specification: <https://www.w3.org/XML/>

**Parameters**

**indent** [int, optional, default=2] If not None, will pretty-print with specified number of spaces for indentation

**Returns**

**serialized** [bytes] A MessagePack-encoded bytes serialized representation.

**to\_yaml(self)**

Return a YAML serialized representation.

Specification: <http://yaml.org/>

**Returns**

**serialized** [str] A YAML serialized representation of the object

**property torsions**

Get an iterator over all i-j-k-l torsions. Note that i-j-k-i torsions (cycles) are excluded.

**Returns**

**torsions** [iterable of 4-Atom tuples]

**property total\_charge**

Return the total charge on the molecule

**property virtual\_sites**

Iterate over all VirtualSite objects.

**add\_atom**(*self*, *atomic\_number*, *formal\_charge*, *is\_aromatic*, *stereochemistry=None*, *name=None*)

Add an atom

#### Parameters

**atomic\_number** [int] Atomic number of the atom

**formal\_charge** [int] Formal charge of the atom

**is\_aromatic** [bool] If True, atom is aromatic; if False, not aromatic

**stereochemistry** [str, optional, default=None] Either ‘R’ or ‘S’ for specified stereochemistry, or None if stereochemistry is irrelevant

**name** [str, optional, default=None] An optional name for the atom

#### Returns

**index** [int] The index of the atom in the molecule

### Examples

Define a methane molecule

```
>>> molecule = Molecule()
>>> molecule.name = 'methane'
>>> C = molecule.add_atom(6, 0, False)
>>> H1 = molecule.add_atom(1, 0, False)
>>> H2 = molecule.add_atom(1, 0, False)
>>> H3 = molecule.add_atom(1, 0, False)
>>> H4 = molecule.add_atom(1, 0, False)
>>> bond_idx = molecule.add_bond(C, H1, False, 1)
>>> bond_idx = molecule.add_bond(C, H2, False, 1)
>>> bond_idx = molecule.add_bond(C, H3, False, 1)
>>> bond_idx = molecule.add_bond(C, H4, False, 1)
```

**add\_bond\_charge\_virtual\_site**(*self*, *atoms*, *distance*, *charge\_increments=None*, *weights=None*, *epsilon=None*, *sigma=None*, *rmin\_half=None*, *name=”*)

Create a bond charge-type virtual site, in which the location of the charge is specified by the position of two atoms. This supports placement of a virtual site S along a vector between two specified atoms, e.g. to allow for a sigma hole for halogens or similar contexts. With positive values of the distance, the virtual site lies outside the first indexed atom. Parameters ——  
*atoms* : list of openforcefield.topology.molecule.Atom objects or ints of shape [N]

The atoms defining the virtual site’s position or their indices

*distance* : float

**weights** [list of floats of shape [N] or None, optional, default=None] *weights*[index] is the weight of particles[index] contributing to the position of the virtual site. Default is None

**charge\_increments** [list of floats of shape [N], optional, default=None] The amount of charge to remove from the VirtualSite’s atoms and put in the VirtualSite. Indexing in this list should match the ordering in the atoms list. Default is None.

**epsilon** [float] Epsilon term for VdW properties of virtual site. Default is None.

**sigma** [float, default=None] Sigma term for VdW properties of virtual site. Default is None.

**rmin\_half** [float] Rmin\_half term for VdW properties of virtual site. Default is None.

**name** [string or None, default=”] The name of this virtual site. Default is “.

**Returns**

**index** [int] The index of the newly-added virtual site in the molecule

**add\_monovalent\_lone\_pair\_virtual\_site(self, atoms, distance, out\_of\_plane\_angle, in\_plane\_angle, \*\*kwargs)**

Create a bond charge-type virtual site, in which the location of the charge is specified by the position of three atoms.

TODO : Do “weights” have any meaning here?

**Parameters**

**atoms** [list of three openforcefield.topology.molecule.Atom objects or ints] The three atoms defining the virtual site’s position or their molecule atom indices

**distance** [float]

**out\_of\_plane\_angle** [float]

**in\_plane\_angle** [float]

**epsilon** [float] Epsilon term for VdW properties of virtual site. Default is None.

**sigma** [float, default=None] Sigma term for VdW properties of virtual site. Default is None.

**rmin\_half** [float] Rmin\_half term for VdW properties of virtual site. Default is None.

**name** [string or None, default=”] The name of this virtual site. Default is “.

**Returns**

**index** [int] The index of the newly-added virtual site in the molecule

**add\_divalent\_lone\_pair\_virtual\_site(self, atoms, distance, out\_of\_plane\_angle, in\_plane\_angle, \*\*kwargs)**

Create a divalent lone pair-type virtual site, in which the location of the charge is specified by the position of three atoms.

TODO : Do “weights” have any meaning here?

**Parameters**

**atoms** [list of 3 openforcefield.topology.molecule.Atom objects or ints] The three atoms defining the virtual site’s position or their molecule atom indices

**distance** [float]

**out\_of\_plane\_angle** [float]

**in\_plane\_angle** [float]

**epsilon** [float] Epsilon term for VdW properties of virtual site. Default is None.

**sigma** [float, default=None] Sigma term for VdW properties of virtual site. Default is None.

**rmin\_half** [float] Rmin\_half term for VdW properties of virtual site. Default is None.

**name** [string or None, default=”] The name of this virtual site. Default is “.

**Returns**

**index** [int] The index of the newly-added virtual site in the molecule

```
add_trivalent_lone_pair_virtual_site(self,      atoms,      distance,      out_of_plane_angle,
                                         in_plane_angle, **kwargs)
```

Create a trivalent lone pair-type virtual site, in which the location of the charge is specified by the position of four atoms.

TODO : Do “weights” have any meaning here?

#### Parameters

**atoms** [list of 4 openforcefield.topology.molecule.Atom objects or ints] The three atoms defining the virtual site’s position or their molecule atom indices

**distance** [float]

**out\_of\_plane\_angle** [float]

**in\_plane\_angle** [float]

**epsilon** [float] Epsilon term for VdW properties of virtual site. Default is None.

**sigma** [float, default=None] Sigma term for VdW properties of virtual site. Default is None.

**rmin\_half** [float] Rmin\_half term for VdW properties of virtual site. Default is None.

**name** [string or None, default=""] The name of this virtual site. Default is “”.

#### Returns

**index** [int] The index of the newly-added virtual site in the molecule

```
add_bond(self, atom1, atom2, bond_order, is_aromatic, stereochemistry=None, fractional_bond_order=None)
```

Add a bond between two specified atom indices

#### Parameters

**atom1** [int or openforcefield.topology.molecule.Atom] Index of first atom

**atom2** [int or openforcefield.topology.molecule.Atom] Index of second atom

**bond\_order** [int] Integral bond order of Kekulized form

**is\_aromatic** [bool] True if this bond is aromatic, False otherwise

**stereochemistry** [str, optional, default=None] Either ‘E’ or ‘Z’ for specified stereochemistry, or None if stereochemistry is irrelevant

**fractional\_bond\_order** [float, optional, default=None] The fractional (eg. Wiberg) bond order

#### Returns

**index: int** Index of the bond in this molecule

```
add_conformer(self, coordinates)
```

# TODO: Should this not be public? Adds a conformer of the molecule

#### Parameters

**coordinates: simtk.unit.Quantity(np.array) with shape (n\_atoms, 3)**

Coordinates of the new conformer, with the first dimension of the array corresponding to the atom index in the Molecule’s indexing system.

#### Returns

---

**index: int** Index of the conformer in the Molecule

## openforcefield.topology.Topology

**class** openforcefield.topology.Topology(*other=None*)

A Topology is a chemical representation of a system containing one or more molecules appearing in a specified order.

**Warning:** This API is experimental and subject to change.

## Examples

Import some utilities

```
>>> from simtk.openmm import app
>>> from openforcefield.tests.utils import get_data_file_path, get_pckmol_pdb_file_path
>>> pdb_filepath = get_pckmol_pdb_file_path('cyclohexane_ethanol_0.4_0.6')
>>> monomer_names = ('cyclohexane', 'ethanol')
```

Create a Topology object from a PDB file and sdf files defining the molecular contents

```
>>> from openforcefield.topology import Molecule, Topology
>>> pdbfile = app.PDBFile(pdb_filepath)
>>> sdf_filepaths = [get_data_file_path(f'systems/monomers/{name}.sdf') for name in monomer_names]
>>> unique_molecules = [Molecule.from_file(sdf_filepath) for sdf_filepath in sdf_filepaths]
>>> topology = Topology.from_openmm(pdbfile.topology, unique_molecules=unique_molecules)
```

Create a Topology object from a PDB file and IUPAC names of the molecular contents

```
>>> pdbfile = app.PDBFile(pdb_filepath)
>>> unique_molecules = [Molecule.from_iupac(name) for name in monomer_names]
>>> topology = Topology.from_openmm(pdbfile.topology, unique_molecules=unique_molecules)
```

Create an empty Topology object and add a few copies of a single benzene molecule

```
>>> topology = Topology()
>>> molecule = Molecule.from_iupac('benzene')
>>> molecule_topology_indices = [topology.add_molecule(molecule) for index in range(10)]
```

## Attributes

**angles** Iterable of Tuple[TopologyAtom]: iterator over the angles in this Topology.

**aromaticity\_model** Get the aromaticity model applied to all molecules in the topology.

**box\_vectors** Return the box vectors of the topology, if specified

**charge\_model** Get the partial charge model applied to all molecules in the topology.

**constrained\_atom\_pairs** Returns the constrained atom pairs of the Topology

**fractional\_bond\_order\_model** Get the fractional bond order model for the Topology.

**impropers** Iterable of Tuple[TopologyAtom]: iterator over the improper torsions in this Topology.

**n\_angles** int: number of angles in this Topology.

`n_impropers` int: number of improper torsions in this Topology.  
`n_proper`s int: number of proper torsions in this Topology.  
`n_reference_molecules` Returns the number of reference (unique) molecules in in this Topology.  
`n_topology_atoms` Returns the number of topology atoms in in this Topology.  
`n_topology_bonds` Returns the number of TopologyBonds in in this Topology.  
`n_topology_molecules` Returns the number of topology molecules in in this Topology.  
`n_topology_particles` Returns the number of topology particles (TopologyAtoms and TopologyVirtualSites) in in this Topology.  
`n_topology_virtual_sites` Returns the number of TopologyVirtualSites in in this Topology.  
`proper`s Iterable of Tuple[TopologyAtom]: iterator over the proper torsions in this Topology.  
`reference_molecules` Get an iterator of reference molecules in this Topology.  
`topology_atoms` Returns an iterator over the atoms in this Topology.  
`topology_bonds` Returns an iterator over the bonds in this Topology  
`topology_molecules` Returns an iterator over all the TopologyMolecules in this Topology  
`topology_particles` Returns an iterator over the particles (TopologyAtoms and TopologyVirtualSites) in this Topology.  
`topology_virtual_sites` Get an iterator over the virtual sites in this Topology

## Methods

<code>add_constraint(self, iatom, jatom[, distance])</code>	Mark a pair of atoms as constrained.
<code>add_molecule(self, molecule[, ...])</code>	Add a Molecule to the Topology.
<code>add_particle(self, particle)</code>	Add a Particle to the Topology.
<code>assert_bonded(self, atom1, atom2)</code>	Raise an exception if the specified atoms are not bonded in the topology.
<code>atom(self, atom_topology_index)</code>	Get the TopologyAtom at a given Topology atom index.
<code>bond(self, bond_topology_index)</code>	Get the TopologyBond at a given Topology bond index.
<code>chemical_environment_matches(self, query[, ...])</code>	Retrieve all matches for a given chemical environment query.
<code>from_bson(serialized)</code>	Instantiate an object from a BSON serialized representation.
<code>from_dict(d)</code>	Static constructor from dictionary representation.
<code>from_json(serialized)</code>	Instantiate an object from a JSON serialized representation.
<code>from_mdtraj(mdtraj_topology[, unique_molecules])</code>	Construct an openforcefield Topology object from an MDTraj Topology object.
<code>from_messagepack(serialized)</code>	Instantiate an object from a MessagePack serialized representation.

Continued on next page

Table 8 – continued from previous page

<code>from_molecules(molecules)</code>	Create a new Topology object containing one copy of each of the specified molecule(s).
<code>from_openmm(openmm_topology[, unique_molecules])</code>	Construct an openforcefield Topology object from an OpenMM Topology object.
<code>from_parmed(parmed_structure[, unique_molecules])</code>	
<b>Warning:</b>	
	This functionality will be implemented in a future toolkit release.
<code>from_pickle(serialized)</code>	Instantiate an object from a pickle serialized representation.
<code>from_toml(serialized)</code>	Instantiate an object from a TOML serialized representation.
<code>from_xml(serialized)</code>	Instantiate an object from an XML serialized representation.
<code>from_yaml(serialized)</code>	Instantiate from a YAML serialized representation.
<code>get_bond_between(self, i, j)</code>	Returns the bond between two atoms
<code>get_fractional_bond_order(self, iatom, jatom)</code>	Retrieve the fractional bond order for a bond.
<code>is_bonded(self, i, j)</code>	Returns True if the two atoms are bonded
<code>is_constrained(self, iatom, jatom)</code>	Check if a pair of atoms are marked as constrained.
<code>to_bson(self)</code>	Return a BSON serialized representation.
<code>to_dict(self)</code>	Convert to dictionary representation.
<code>to_json(self[, indent])</code>	Return a JSON serialized representation.
<code>to_messagepack(self)</code>	Return a MessagePack representation.
<code>to_openmm(self)</code>	Create an OpenMM Topology object.

Continued on next page

Table 8 – continued from previous page

`to_parmed(self)`

**Warning:**  
 This functionality will be implemented in a future toolkit release.

<code>to_pickle(self)</code>	Return a pickle serialized representation.
<code>to_toml(self)</code>	Return a TOML serialized representation.
<code>to_xml(self[, indent])</code>	Return an XML representation.
<code>to_yaml(self)</code>	Return a YAML serialized representation.
<code>virtual_site(self, vsite_topology_index)</code>	Get the TopologyAtom at a given Topology atom index.

`__init__(self, other=None)`

Create a new Topology.

#### Parameters

**other** [optional, default=None] If specified, attempt to construct a copy of the Topology from the specified object. This might be a Topology object, or a file that can be used to construct a Topology object or serialized Topology object.

#### Methods

<code>__init__(self[, other])</code>	Create a new Topology.
<code>add_constraint(self, iatom, jatom[, distance])</code>	Mark a pair of atoms as constrained.
<code>add_molecule(self, molecule[, ...])</code>	Add a Molecule to the Topology.
<code>add_particle(self, particle)</code>	Add a Particle to the Topology.
<code>assert_bonded(self, atom1, atom2)</code>	Raise an exception if the specified atoms are not bonded in the topology.
<code>atom(self, atom_topology_index)</code>	Get the TopologyAtom at a given Topology atom index.
<code>bond(self, bond_topology_index)</code>	Get the TopologyBond at a given Topology bond index.

Continued on next page

Table 9 – continued from previous page

<code>chemical_environment_matches(self, query[, ...])</code>	Retrieve all matches for a given chemical environment query.
<code>from_bson(serialized)</code>	Instantiate an object from a BSON serialized representation.
<code>from_dict(d)</code>	Static constructor from dictionary representation.
<code>from_json(serialized)</code>	Instantiate an object from a JSON serialized representation.
<code>from_mdtraj(mdtraj_topology[, unique_molecules])</code>	Construct an openforcefield Topology object from an MDTraj Topology object.
<code>from_messagepack(serialized)</code>	Instantiate an object from a MessagePack serialized representation.
<code>from_molecules(molecules)</code>	Create a new Topology object containing one copy of each of the specified molecule(s).
<code>from_openmm(openmm_topology[, unique_molecules])</code>	Construct an openforcefield Topology object from an OpenMM Topology object.
<code>from_parmed(parmed_structure[, unique_molecules])</code>	

**Warning:**  
 This functionality will be implemented in a future toolkit release.

<code>from_pickle(serialized)</code>	Instantiate an object from a pickle serialized representation.
<code>from_toml(serialized)</code>	Instantiate an object from a TOML serialized representation.
<code>from_xml(serialized)</code>	Instantiate an object from an XML serialized representation.
<code>from_yaml(serialized)</code>	Instantiate from a YAML serialized representation.
<code>get_bond_between(self, i, j)</code>	Returns the bond between two atoms
<code>get_fractional_bond_order(self, iatom, jatom)</code>	Retrieve the fractional bond order for a bond.
<code>is_bonded(self, i, j)</code>	Returns True if the two atoms are bonded
<code>is_constrained(self, iatom, jatom)</code>	Check if a pair of atoms are marked as constrained.
<code>to_bson(self)</code>	Return a BSON serialized representation.

Continued on next page

Table 9 – continued from previous page

<code>to_dict(self)</code>	Convert to dictionary representation.
<code>to_json(self[, indent])</code>	Return a JSON serialized representation.
<code>to_messagepack(self)</code>	Return a MessagePack representation.
<code>to_openmm(self)</code>	Create an OpenMM Topology object.
<code>to_parmed(self)</code>	

**Warning:**

This functionality will be implemented in a future toolkit release.

<code>to_pickle(self)</code>	Return a pickle serialized representation.
<code>to_toml(self)</code>	Return a TOML serialized representation.
<code>to_xml(self[, indent])</code>	Return an XML representation.
<code>to_yaml(self)</code>	Return a YAML serialized representation.
<code>virtual_site(self, vsite_topology_index)</code>	Get the TopologyAtom at a given Topology atom index.

**Attributes**

<code>angles</code>	Iterable of Tuple[TopologyAtom]: iterator over the angles in this Topology.
<code>aromaticity_model</code>	Get the aromaticity model applied to all molecules in the topology.
<code>box_vectors</code>	Return the box vectors of the topology, if specified Returns —— <code>box_vectors</code> : simtk.unit.Quantity wrapped numpy array The unit-wrapped box vectors of this topology
<code>charge_model</code>	Get the partial charge model applied to all molecules in the topology.
<code>constrained_atom_pairs</code>	Returns the constrained atom pairs of the Topology
<code>fractional_bond_order_model</code>	Get the fractional bond order model for the Topology.
<code>impropers</code>	Iterable of Tuple[TopologyAtom]: iterator over the improper torsions in this Topology.

Continued on next page

Table 10 – continued from previous page

<code>n_angles</code>	int: number of angles in this Topology.
<code>n_impropers</code>	int: number of improper torsions in this Topology.
<code>n_proper</code>	int: number of proper torsions in this Topology.
<code>n_reference_molecules</code>	Returns the number of reference (unique) molecules in in this Topology.
<code>n_topology_atoms</code>	Returns the number of topology atoms in in this Topology.
<code>n_topology_bonds</code>	Returns the number of TopologyBonds in in this Topology.
<code>n_topology_molecules</code>	Returns the number of topology molecules in in this Topology.
<code>n_topology_particles</code>	Returns the number of topology particles (TopologyAtoms and TopologyVirtualSites) in in this Topology.
<code>n_topology_virtual_sites</code>	Returns the number of TopologyVirtualSites in in this Topology.
<code>proper</code>	Iterable of Tuple[TopologyAtom]: iterator over the proper torsions in this Topology.
<code>reference_molecules</code>	Get an iterator of reference molecules in this Topology.
<code>topology_atoms</code>	Returns an iterator over the atoms in this Topology.
<code>topology_bonds</code>	Returns an iterator over the bonds in this Topology
<code>topology_molecules</code>	Returns an iterator over all the TopologyMolecules in this Topology
<code>topology_particles</code>	Returns an iterator over the particles (TopologyAtoms and TopologyVirtualSites) in this Topology.
<code>topology_virtual_sites</code>	Get an iterator over the virtual sites in this Topology

**property `reference_molecules`**

Get an iterator of reference molecules in this Topology.

**Returns****iterable of `openforcefield.topology.Molecule`****classmethod `from_molecules(molecules)`**

Create a new Topology object containing one copy of each of the specified molecule(s).

**Parameters****molecules** [Molecule or iterable of Molecules] One or more molecules to be added to the Topology**Returns****topology** [Topology] The Topology created from the specified molecule(s)**assert\_bonded(*self*, *atom1*, *atom2*)**

Raise an exception if the specified atoms are not bonded in the topology.

**Parameters**

**atom1, atom2** [openforcefield.topology.Atom or int] The atoms or atom topology indices to check to ensure they are bonded

**property aromaticity\_model**

Get the aromaticity model applied to all molecules in the topology.

**Returns**

**aromaticity\_model** [str] Aromaticity model in use.

**property box\_vectors**

Return the box vectors of the topology, if specified Returns ——— box\_vectors : simtk.unit.Quantity wrapped numpy array

The unit-wrapped box vectors of this topology

**property charge\_model**

Get the partial charge model applied to all molecules in the topology.

**Returns**

**charge\_model** [str] Charge model used for all molecules in the Topology.

**property constrained\_atom\_pairs**

Returns the constrained atom pairs of the Topology

**Returns**

**constrained\_atom\_pairs** [dict] dictionary of the form d[(atom1\_topology\_index, atom2\_topology\_index)] = distance (float)

**property fractional\_bond\_order\_model**

Get the fractional bond order model for the Topology.

**Returns**

**fractional\_bond\_order\_model** [str] Fractional bond order model in use.

**property n\_reference\_molecules**

Returns the number of reference (unique) molecules in in this Topology.

**Returns**

**n\_reference\_molecules** [int]

**property n\_topology\_molecules**

Returns the number of topology molecules in in this Topology.

**Returns**

**n\_topology\_molecules** [int]

**property topology\_molecules**

Returns an iterator over all the TopologyMolecules in this Topology

**Returns**

**topology\_molecules** [Iterable of TopologyMolecule]

**property n\_topology\_atoms**

Returns the number of topology atoms in in this Topology.

**Returns**

**n\_topology\_atoms** [int]

**property topology\_atoms**

Returns an iterator over the atoms in this Topology. These will be in ascending order of topology index (Note that this is not necessarily the same as the reference molecule index)

**Returns**

**topology\_atoms** [Iterable of TopologyAtom]

**property n\_topology\_bonds**

Returns the number of TopologyBonds in in this Topology.

**Returns**

**n\_bonds** [int]

**property topology\_bonds**

Returns an iterator over the bonds in this Topology

**Returns**

**topology\_bonds** [Iterable of TopologyBond]

**property n\_topology\_particles**

Returns the number of topology particles (TopologyAtoms and TopologyVirtualSites) in in this Topology.

**Returns**

**n\_topology\_particles** [int]

**property topology\_particles**

Returns an iterator over the particles (TopologyAtoms and TopologyVirtualSites) in this Topology. The TopologyAtoms will be in order of ascending Topology index (Note that this may differ from the order of atoms in the reference molecule index).

**Returns**

**topology\_particles** [Iterable of TopologyAtom and TopologyVirtualSite]

**property n\_topology\_virtual\_sites**

Returns the number of TopologyVirtualSites in in this Topology.

**Returns**

**n\_virtual\_sites** [iterable of TopologyVirtualSites]

**property topology\_virtual\_sites**

Get an iterator over the virtual sites in this Topology

**Returns**

**topology\_virtual\_sites** [Iterable of TopologyVirtualSite]

**property n\_angles**

int: number of angles in this Topology.

**property angles**

Iterable of Tuple[TopologyAtom]: iterator over the angles in this Topology.

**property n\_proper\_torsions**

int: number of proper torsions in this Topology.

**property proper\_torsions**

Iterable of Tuple[TopologyAtom]: iterator over the proper torsions in this Topology.

**property n\_impropers**

int: number of improper torsions in this Topology.

**property impropers**

Iterable of Tuple[TopologyAtom]: iterator over the improper torsions in this Topology.

**chemical\_environment\_matches(self, query, toolkit\_registry=**  
*aromaticity\_model='MDL', object at 0x7fba569d3128>)*

Retrieve all matches for a given chemical environment query.

TODO: \* Do we want to generalize this to other kinds of queries too, like mdtraj DSL, pymol selections, atom index slices, etc?

We could just call it topology.matches(query)

**Parameters**

**query** [str or ChemicalEnvironment] SMARTS string (with one or more tagged atoms) or ChemicalEnvironment query Query will internally be resolved to SMARTS using query.as\_smarts() if it has an .as\_smarts method.

**aromaticity\_model** [str] Override the default aromaticity model for this topology and use the specified aromaticity model instead. Allowed values: ['MDL']

**Returns**

**matches** [list of Topology.\_ChemicalEnvironmentMatch] A list of tuples, containing the topology indices of the matching atoms.

**to\_dict(self)**

Convert to dictionary representation.

**classmethod from\_dict(d)**

Static constructor from dictionary representation.

**classmethod from\_openmm(openmm\_topology, unique\_molecules=None)**

Construct an openforcefield Topology object from an OpenMM Topology object.

**Parameters**

**openmm\_topology** [simtk.openmm.app.Topology] An OpenMM Topology object

**unique\_molecules** [iterable of objects that can be used to construct unique Molecule objects] All unique molecules must be provided, in any order, though multiple copies of each molecule are allowed. The atomic elements and bond connectivity will be used to match the reference molecules to molecule graphs appearing in the OpenMM Topology. If bond orders are present in the OpenMM Topology, these will be used in matching as well. If all bonds have bond orders assigned in mdtraj\_topology, these bond orders will be used to attempt to construct the list of unique Molecules if the unique\_molecules argument is omitted.

**Returns**

**topology** [openforcefield.topology.Topology] An openforcefield Topology object

**to\_openmm(self)**

Create an OpenMM Topology object.

The OpenMM Topology object will have one residue per topology molecule. Currently, the number of chains depends on how many copies of the same molecule are in the Topology. Molecules with

more than 5 copies are all assigned to a single chain, otherwise one chain is created for each molecule. This behavior may change in the future.

#### Parameters

**openmm\_topology** [simtk.openmm.app.Topology] An OpenMM Topology object

**static from\_mdtraj(mdtraj\_topology, unique\_molecules=None)**

Construct an openforcefield Topology object from an MDTraj Topology object.

#### Parameters

**mdtraj\_topology** [mdtraj.Topology] An MDTraj Topology object

**unique\_molecules** [iterable of objects that can be used to construct unique Molecule objects] All unique molecules must be provided, in any order, though multiple copies of each molecule are allowed. The atomic elements and bond connectivity will be used to match the reference molecules to molecule graphs appearing in the MDTraj Topology. If bond orders are present in the MDTraj Topology, these will be used in matching as well. If all bonds have bond orders assigned in `mdtraj_topology`, these bond orders will be used to attempt to construct the list of unique Molecules if the `unique_molecules` argument is omitted.

#### Returns

**topology** [openforcefield.Topology] An openforcefield Topology object

**static from\_parmed(parmed\_structure, unique\_molecules=None)**

**Warning:** This functionality will be implemented in a future toolkit release.

Construct an openforcefield Topology object from a ParmEd Structure object.

#### Parameters

**parmed\_structure** [parmed.Structure] A ParmEd structure object

**unique\_molecules** [iterable of objects that can be used to construct unique Molecule objects] All unique molecules must be provided, in any order, though multiple copies of each molecule are allowed. The atomic elements and bond connectivity will be used to match the reference molecules to molecule graphs appearing in the structure's topology object. If bond orders are present in the structure's topology object, these will be used in matching as well. If all bonds have bond orders assigned in the structure's topology object, these bond orders will be used to attempt to construct the list of unique Molecules if the `unique_molecules` argument is omitted.

#### Returns

**topology** [openforcefield.Topology] An openforcefield Topology object

**to\_parmed(self)**

**Warning:** This functionality will be implemented in a future toolkit release.

Create a ParmEd Structure object.

**Returns**

**parmed\_structure** [parmed.Structure] A ParmEd Structure object

**get\_bond\_between(self, i, j)**

Returns the bond between two atoms

**Parameters**

**i, j** [int or TopologyAtom] Atoms or atom indices to check

**Returns**

**bond** [TopologyBond] The bond between i and j.

**is\_bonded(self, i, j)**

Returns True if the two atoms are bonded

**Parameters**

**i, j** [int or TopologyAtom] Atoms or atom indices to check

**Returns**

**is\_bonded** [bool] True if atoms are bonded, False otherwise.

**atom(self, atom\_topology\_index)**

Get the TopologyAtom at a given Topology atom index.

**Parameters**

**atom\_topology\_index** [int] The index of the TopologyAtom in this Topology

**Returns**

An openforcefield.topology.TopologyAtom

**virtual\_site(self, vsite\_topology\_index)**

Get the TopologyAtom at a given Topology atom index.

**Parameters**

**vsite\_topology\_index** [int] The index of the TopologyVirtualSite in this Topology

**Returns**

An openforcefield.topology.TopologyVirtualSite

**bond(self, bond\_topology\_index)**

Get the TopologyBond at a given Topology bond index.

**Parameters**

**bond\_topology\_index** [int] The index of the TopologyBond in this Topology

**Returns**

An openforcefield.topology.TopologyBond

**add\_particle(self, particle)**

Add a Particle to the Topology.

**Parameters**

**particle** [Particle] The Particle to be added. The Topology will take ownership of the Particle.

**add\_molecule(self, molecule, local\_topology\_to\_reference\_index=None)**

Add a Molecule to the Topology.

**Parameters**

**molecule** [Molecule] The Molecule to be added.

**local\_topology\_to\_reference\_index: dict, optional, default = None** Dictionary of {TopologyMolecule\_atom\_index : Molecule\_atom\_index} for the Topology-Molecule that will be built

**Returns**

**index** [int] The index of this molecule in the topology

**classmethod from\_bson(serialized)**

Instantiate an object from a BSON serialized representation.

Specification: <http://bsonspec.org/>

**Parameters**

**serialized** [bytes] A BSON serialized representation of the object

**Returns**

**instance** [cls] An instantiated object

**classmethod from\_json(serialized)**

Instantiate an object from a JSON serialized representation.

Specification: <https://www.json.org/>

**Parameters**

**serialized** [str] A JSON serialized representation of the object

**Returns**

**instance** [cls] An instantiated object

**classmethod from\_messagepack(serialized)**

Instantiate an object from a MessagePack serialized representation.

Specification: <https://msgpack.org/index.html>

**Parameters**

**serialized** [bytes] A MessagePack-encoded bytes serialized representation

**Returns**

**instance** [cls] Instantiated object.

**classmethod from\_pickle(serialized)**

Instantiate an object from a pickle serialized representation.

**Warning:** This is not recommended for safe, stable storage since the pickle specification may change between Python versions.

**Parameters**

**serialized** [str] A pickled representation of the object

**Returns**

**instance** [cls] An instantiated object

**classmethod from\_toml(serialized)**  
Instantiate an object from a TOML serialized representation.

Specification: <https://github.com/toml-lang/toml>

**Parameters**

**serialized** [str] A TOML serialized representation of the object

**Returns**

**instance** [cls] An instantiated object

**classmethod from\_xml(serialized)**  
Instantiate an object from an XML serialized representation.

Specification: <https://www.w3.org/XML/>

**Parameters**

**serialized** [bytes] An XML serialized representation

**Returns**

**instance** [cls] Instantiated object.

**classmethod from\_yaml(serialized)**  
Instantiate from a YAML serialized representation.

Specification: <http://yaml.org/>

**Parameters**

**serialized** [str] A YAML serialized representation of the object

**Returns**

**instance** [cls] Instantiated object

**to bson(self)**  
Return a BSON serialized representation.

Specification: <http://bsonspec.org/>

**Returns**

**serialized** [bytes] A BSON serialized representation of the object

**to json(self, indent=None)**  
Return a JSON serialized representation.

Specification: <https://www.json.org/>

**Parameters**

**indent** [int, optional, default=None] If not None, will pretty-print with specified number of spaces for indentation

**Returns**

**serialized** [str] A JSON serialized representation of the object

**to\_messagepack(self)**  
Return a MessagePack representation.

Specification: <https://msgpack.org/index.html>

**Returns**

**serialized** [bytes] A MessagePack-encoded bytes serialized representation of the object

**to\_pickle(self)**

Return a pickle serialized representation.

**Warning:** This is not recommended for safe, stable storage since the pickle specification may change between Python versions.

**Returns**

**serialized** [str] A pickled representation of the object

**to\_toml(self)**

Return a TOML serialized representation.

Specification: <https://github.com/toml-lang/toml>

**Returns**

**serialized** [str] A TOML serialized representation of the object

**to\_xml(self, indent=2)**

Return an XML representation.

Specification: <https://www.w3.org/XML/>

**Parameters**

**indent** [int, optional, default=2] If not None, will pretty-print with specified number of spaces for indentation

**Returns**

**serialized** [bytes] A MessagePack-encoded bytes serialized representation.

**to\_yaml(self)**

Return a YAML serialized representation.

Specification: <http://yaml.org/>

**Returns**

**serialized** [str] A YAML serialized representation of the object

**add\_constraint(self, iatom, jatom, distance=True)**

Mark a pair of atoms as constrained.

Constraints between atoms that are not bonded (e.g., rigid waters) are permissible.

**Parameters**

**iatom, jatom** [Atom] Atoms to mark as constrained These atoms may be bonded or not in the Topology

**distance** [simtk.unit.Quantity, optional, default=True] Constraint distance True if distance has yet to be determined False if constraint is to be removed

**is\_constrained(self, iatom, jatom)**

Check if a pair of atoms are marked as constrained.

**Parameters**

**iatom, jatom** [int] Indices of atoms to mark as constrained.

**Returns**

**distance** [simtk.unit.Quantity or bool] True if constrained but constraints have not yet been applied Distance if constraint has already been added to System

**get\_fractional\_bond\_order(self, iatom, jatom)**  
Retrieve the fractional bond order for a bond.

An Exception is raised if it cannot be determined.

**Parameters**

**iatom, jatom** [Atom] Atoms for which a fractional bond order is to be retrieved.

**Returns**

**order** [float] Fractional bond order between the two specified atoms.

## 2.1.2 Secondary objects

Particle	Base class for all particles in a molecule.
Atom	A particle representing a chemical atom.
Bond	Chemical bond representation.
VirtualSite	A particle representing a virtual site whose position is defined in terms of Atom positions.

### openforcefield.topology.Particle

**class openforcefield.topology.Particle**  
Base class for all particles in a molecule.

A particle object could be an Atom or a VirtualSite.

**Warning:** This API is experimental and subject to change.

**Attributes**

**molecule** The Molecule this atom is part of.

**molecule\_particle\_index** Returns the index of this particle in its molecule

**name** The name of the particle

**Methods**

<b>from_bson(serialized)</b>	Instantiate an object from a BSON serialized representation.
<b>from_dict(d)</b>	Static constructor from dictionary representation.
<b>from_json(serialized)</b>	Instantiate an object from a JSON serialized representation.
<b>from_messagepack(serialized)</b>	Instantiate an object from a MessagePack serialized representation.

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<code>from_pickle(serialized)</code>	Instantiate an object from a pickle serialized representation.
<code>from_toml(serialized)</code>	Instantiate an object from a TOML serialized representation.
<code>from_xml(serialized)</code>	Instantiate an object from an XML serialized representation.
<code>from_yaml(serialized)</code>	Instantiate from a YAML serialized representation.
<code>to_bson(self)</code>	Return a BSON serialized representation.
<code>to_dict(self)</code>	Convert to dictionary representation.
<code>to_json(self[, indent])</code>	Return a JSON serialized representation.
<code>to_messagepack(self)</code>	Return a MessagePack representation.
<code>to_pickle(self)</code>	Return a pickle serialized representation.
<code>to_toml(self)</code>	Return a TOML serialized representation.
<code>to_xml(self[, indent])</code>	Return an XML representation.
<code>to_yaml(self)</code>	Return a YAML serialized representation.

`__init__(self, /, *args, **kwargs)`

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

## Methods

<code>__init__(self, /, *args, **kwargs)</code>	Initialize self.
<code>from_bson(serialized)</code>	Instantiate an object from a BSON serialized representation.
<code>from_dict(d)</code>	Static constructor from dictionary representation.
<code>from_json(serialized)</code>	Instantiate an object from a JSON serialized representation.
<code>from_messagepack(serialized)</code>	Instantiate an object from a MessagePack serialized representation.
<code>from_pickle(serialized)</code>	Instantiate an object from a pickle serialized representation.
<code>from_toml(serialized)</code>	Instantiate an object from a TOML serialized representation.
<code>from_xml(serialized)</code>	Instantiate an object from an XML serialized representation.
<code>from_yaml(serialized)</code>	Instantiate from a YAML serialized representation.
<code>to_bson(self)</code>	Return a BSON serialized representation.
<code>to_dict(self)</code>	Convert to dictionary representation.
<code>to_json(self[, indent])</code>	Return a JSON serialized representation.
<code>to_messagepack(self)</code>	Return a MessagePack representation.
<code>to_pickle(self)</code>	Return a pickle serialized representation.
<code>to_toml(self)</code>	Return a TOML serialized representation.
<code>to_xml(self[, indent])</code>	Return an XML representation.
<code>to_yaml(self)</code>	Return a YAML serialized representation.

## Attributes

<code>molecule</code>	The Molecule this atom is part of.
<code>molecule_particle_index</code>	Returns the index of this particle in its molecule
<code>name</code>	The name of the particle

### `property molecule`

The Molecule this atom is part of.

### `property molecule_particle_index`

Returns the index of this particle in its molecule

### `property name`

The name of the particle

### `to_dict(self)`

Convert to dictionary representation.

### `classmethod from_dict(d)`

Static constructor from dictionary representation.

### `classmethod from_bson(serialized)`

Instantiate an object from a BSON serialized representation.

Specification: <http://bsonspec.org/>

#### Parameters

`serialized` [bytes] A BSON serialized representation of the object

#### Returns

`instance` [cls] An instantiated object

### `classmethod from_json(serialized)`

Instantiate an object from a JSON serialized representation.

Specification: <https://www.json.org/>

#### Parameters

`serialized` [str] A JSON serialized representation of the object

#### Returns

`instance` [cls] An instantiated object

### `classmethod from_messagepack(serialized)`

Instantiate an object from a MessagePack serialized representation.

Specification: <https://msgpack.org/index.html>

#### Parameters

`serialized` [bytes] A MessagePack-encoded bytes serialized representation

#### Returns

`instance` [cls] Instantiated object.

### `classmethod from_pickle(serialized)`

Instantiate an object from a pickle serialized representation.

**Warning:** This is not recommended for safe, stable storage since the pickle specification may change between Python versions.

#### Parameters

**serialized** [str] A pickled representation of the object

#### Returns

**instance** [cls] An instantiated object

#### **classmethod from\_toml(serialized)**

Instantiate an object from a TOML serialized representation.

Specification: <https://github.com/toml-lang/toml>

#### Parameters

**serialized** [str] A TOML serialized representation of the object

#### Returns

**instance** [cls] An instantiated object

#### **classmethod from\_xml(serialized)**

Instantiate an object from an XML serialized representation.

Specification: <https://www.w3.org/XML/>

#### Parameters

**serialized** [bytes] An XML serialized representation

#### Returns

**instance** [cls] Instantiated object.

#### **classmethod from\_yaml(serialized)**

Instantiate from a YAML serialized representation.

Specification: <http://yaml.org/>

#### Parameters

**serialized** [str] A YAML serialized representation of the object

#### Returns

**instance** [cls] Instantiated object

#### **to\_bson(self)**

Return a BSON serialized representation.

Specification: <http://bsonspec.org/>

#### Returns

**serialized** [bytes] A BSON serialized representation of the object

#### **to\_json(self, indent=None)**

Return a JSON serialized representation.

Specification: <https://www.json.org/>

#### Parameters

**indent** [int, optional, default=None] If not None, will pretty-print with specified number of spaces for indentation

#### Returns

**serialized** [str] A JSON serialized representation of the object

#### **to\_messagepack(self)**

Return a MessagePack representation.

Specification: <https://msgpack.org/index.html>

#### Returns

**serialized** [bytes] A MessagePack-encoded bytes serialized representation of the object

#### **to\_pickle(self)**

Return a pickle serialized representation.

**Warning:** This is not recommended for safe, stable storage since the pickle specification may change between Python versions.

#### Returns

**serialized** [str] A pickled representation of the object

#### **to\_toml(self)**

Return a TOML serialized representation.

Specification: <https://github.com/toml-lang/toml>

#### Returns

**serialized** [str] A TOML serialized representation of the object

#### **to\_xml(self, indent=2)**

Return an XML representation.

Specification: <https://www.w3.org/XML/>

#### Parameters

**indent** [int, optional, default=2] If not None, will pretty-print with specified number of spaces for indentation

#### Returns

**serialized** [bytes] A MessagePack-encoded bytes serialized representation.

#### **to\_yaml(self)**

Return a YAML serialized representation.

Specification: <http://yaml.org/>

#### Returns

**serialized** [str] A YAML serialized representation of the object

## openforcefield.topology.Atom

```
class openforcefield.topology.Atom(atomic_number, formal_charge, is_aromatic, name=None,
                                    molecule=None, stereochemistry=None)
```

A particle representing a chemical atom.

Note that non-chemical virtual sites are represented by the `VirtualSite` object.

**Warning:** This API is experimental and subject to change.

### Attributes

- `atomic_number` The integer atomic number of the atom.
- `bonded_atoms` The list of `Atom` objects this atom is involved in bonds with
- `bonds` The list of `Bond` objects this atom is involved in.
- `element` The element name
- `formal_charge` The atom's formal charge
- `is_aromatic` The atom's `is_aromatic` flag
- `mass` The standard atomic weight (abundance-weighted isotopic mass) of the atomic site.
- `molecule` The `Molecule` this atom is part of.
- `molecule_atom_index` The index of this Atom within the the list of atoms in Molecules.
- `molecule_particle_index` The index of this Atom within the the list of particles in the parent Molecule.
- `name` The name of this atom, if any
- `partial_charge` The partial charge of the atom, if any.
- `stereochemistry` The atom's stereochemistry (if defined, otherwise None)
- `virtual_sites` The list of `VirtualSite` objects this atom is involved in.

### Methods

<code>add_bond(self, bond)</code>	Adds a bond that this atom is involved in ..
<code>add_virtual_site(self, vsite)</code>	Adds a bond that this atom is involved in ..
<code>from_bson(serialized)</code>	Instantiate an object from a BSON serialized representation.
<code>from_dict(atom_dict)</code>	Create an Atom from a dict representation.
<code>from_json(serialized)</code>	Instantiate an object from a JSON serialized representation.
<code>from_messagepack(serialized)</code>	Instantiate an object from a MessagePack serialized representation.
<code>from_pickle(serialized)</code>	Instantiate an object from a pickle serialized representation.
<code>from_toml(serialized)</code>	Instantiate an object from a TOML serialized representation.

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<code>from_xml(serialized)</code>	Instantiate an object from an XML serialized representation.
<code>from_yaml(serialized)</code>	Instantiate from a YAML serialized representation.
<code>is_bonded_to(self, atom2)</code>	Determine whether this atom is bound to another atom
<code>to_bson(self)</code>	Return a BSON serialized representation.
<code>to_dict(self)</code>	Return a dict representation of the atom.
<code>to_json(self[, indent])</code>	Return a JSON serialized representation.
<code>to_messagepack(self)</code>	Return a MessagePack representation.
<code>to_pickle(self)</code>	Return a pickle serialized representation.
<code>to_toml(self)</code>	Return a TOML serialized representation.
<code>to_xml(self[, indent])</code>	Return an XML representation.
<code>to_yaml(self)</code>	Return a YAML serialized representation.

`__init__(self, atomic_number, formal_charge, is_aromatic, name=None, molecule=None, stereochemistry=None)`

Create an immutable Atom object.

Object is serializable and immutable.

#### Parameters

`atomic_number` [int] Atomic number of the atom

`formal_charge` [int] Formal charge of the atom

`is_aromatic` [bool] If True, atom is aromatic; if False, not aromatic

`stereochemistry` [str, optional, default=None] Either ‘R’ or ‘S’ for specified stereochemistry, or None for ambiguous stereochemistry

`name` [str, optional, default=None] An optional name to be associated with the atom

#### Examples

Create a non-aromatic carbon atom

```
>>> atom = Atom(6, 0, False)
```

Create a chiral carbon atom

```
>>> atom = Atom(6, 0, False, stereochemistry='R', name='CT')
```

#### Methods

<code>__init__(self, atomic_number, formal_charge, ...)</code>	Create an immutable Atom object.
<code>add_bond(self, bond)</code>	Adds a bond that this atom is involved in ..
<code>add_virtual_site(self, vsite)</code>	Adds a bond that this atom is involved in ..
<code>from_bson(serialized)</code>	Instantiate an object from a BSON serialized representation.
<code>from_dict(atom_dict)</code>	Create an Atom from a dict representation.

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Table 16 – continued from previous page

<code>from_json(serialized)</code>	Instantiate an object from a JSON serialized representation.
<code>from_messagepack(serialized)</code>	Instantiate an object from a MessagePack serialized representation.
<code>from_pickle(serialized)</code>	Instantiate an object from a pickle serialized representation.
<code>from_toml(serialized)</code>	Instantiate an object from a TOML serialized representation.
<code>from_xml(serialized)</code>	Instantiate an object from an XML serialized representation.
<code>from_yaml(serialized)</code>	Instantiate from a YAML serialized representation.
<code>is_bonded_to(self, atom2)</code>	Determine whether this atom is bound to another atom
<code>to_bson(self)</code>	Return a BSON serialized representation.
<code>to_dict(self)</code>	Return a dict representation of the atom.
<code>to_json(self[, indent])</code>	Return a JSON serialized representation.
<code>to_messagepack(self)</code>	Return a MessagePack representation.
<code>to_pickle(self)</code>	Return a pickle serialized representation.
<code>to_toml(self)</code>	Return a TOML serialized representation.
<code>to_xml(self[, indent])</code>	Return an XML representation.
<code>to_yaml(self)</code>	Return a YAML serialized representation.

## Attributes

<code>atomic_number</code>	The integer atomic number of the atom.
<code>bonded_atoms</code>	The list of Atom objects this atom is involved in bonds with
<code>bonds</code>	The list of Bond objects this atom is involved in.
<code>element</code>	The element name
<code>formal_charge</code>	The atom's formal charge
<code>is_aromatic</code>	The atom's is_aromatic flag
<code>mass</code>	The standard atomic weight (abundance-weighted isotopic mass) of the atomic site.
<code>molecule</code>	The Molecule this atom is part of.
<code>molecule_atom_index</code>	The index of this Atom within the the list of atoms in Molecules.
<code>molecule_particle_index</code>	The index of this Atom within the the list of particles in the parent Molecule.
<code>name</code>	The name of this atom, if any
<code>partial_charge</code>	The partial charge of the atom, if any.
<code>stereochemistry</code>	The atom's stereochemistry (if defined, otherwise None)
<code>virtual_sites</code>	The list of VirtualSite objects this atom is involved in.

`add_bond(self, bond)`

Adds a bond that this atom is involved in .. todo :: Is this how we want to keep records?

### Parameters

`bond: an openforcefield.topology.molecule.Bond` A bond involving this atom

```
add_virtual_site(self, vsite)
    Adds a bond that this atom is involved in .. todo :: Is this how we want to keep records?

Parameters
    bond: an openforcefield.topology.molecule.Bond A bond involving this atom

to_dict(self)
    Return a dict representation of the atom.

classmethod from_dict(atom_dict)
    Create an Atom from a dict representation.

property formal_charge
    The atom's formal charge

property partial_charge
    The partial charge of the atom, if any.

Returns
    float or None

property is_aromatic
    The atom's is_aromatic flag

property stereochemistry
    The atom's stereochemistry (if defined, otherwise None)

property element
    The element name

property atomic_number
    The integer atomic number of the atom.

property mass
    The standard atomic weight (abundance-weighted isotopic mass) of the atomic site.

    TODO (from jeff): Are there atoms that have different chemical properties based on their isotopes?

property name
    The name of this atom, if any

property bonds
    The list of Bond objects this atom is involved in.

property bonded_atoms
    The list of Atom objects this atom is involved in bonds with

is_bonded_to(self, atom2)
    Determine whether this atom is bound to another atom

Parameters
    atom2: openforcefield.topology.molecule.Atom a different atom in the same molecule

Returns
    bool Whether this atom is bound to atom2

property virtual_sites
    The list of VirtualSite objects this atom is involved in.
```

**property molecule\_atom\_index**

The index of this Atom within the the list of atoms in Molecules. Note that this can be different from molecule\_particle\_index.

**property molecule\_particle\_index**

The index of this Atom within the the list of particles in the parent Molecule. Note that this can be different from molecule\_atom\_index.

**classmethod from\_bson(*serialized*)**

Instantiate an object from a BSON serialized representation.

Specification: <http://bsonspec.org/>

**Parameters**

**serialized** [bytes] A BSON serialized representation of the object

**Returns**

**instance** [cls] An instantiated object

**classmethod from\_json(*serialized*)**

Instantiate an object from a JSON serialized representation.

Specification: <https://www.json.org/>

**Parameters**

**serialized** [str] A JSON serialized representation of the object

**Returns**

**instance** [cls] An instantiated object

**classmethod from\_messagepack(*serialized*)**

Instantiate an object from a MessagePack serialized representation.

Specification: <https://msgpack.org/index.html>

**Parameters**

**serialized** [bytes] A MessagePack-encoded bytes serialized representation

**Returns**

**instance** [cls] Instantiated object.

**classmethod from\_pickle(*serialized*)**

Instantiate an object from a pickle serialized representation.

**Warning:** This is not recommended for safe, stable storage since the pickle specification may change between Python versions.

**Parameters**

**serialized** [str] A pickled representation of the object

**Returns**

**instance** [cls] An instantiated object

**classmethod from\_toml(*serialized*)**

Instantiate an object from a TOML serialized representation.

Specification: <https://github.com/toml-lang/toml>

**Parameters**

**serialized** [str] A TOML serialized representation of the object

**Returns**

**instance** [cls] An instantiated object

**classmethod from\_xml(serialized)**

Instantiate an object from an XML serialized representation.

Specification: <https://www.w3.org/XML/>

**Parameters**

**serialized** [bytes] An XML serialized representation

**Returns**

**instance** [cls] Instantiated object.

**classmethod from\_yaml(serialized)**

Instantiate from a YAML serialized representation.

Specification: <http://yaml.org/>

**Parameters**

**serialized** [str] A YAML serialized representation of the object

**Returns**

**instance** [cls] Instantiated object

**property molecule**

The Molecule this atom is part of.

**to bson(self)**

Return a BSON serialized representation.

Specification: <http://bsonspec.org/>

**Returns**

**serialized** [bytes] A BSON serialized representation of the object

**to json(self, indent=None)**

Return a JSON serialized representation.

Specification: <https://www.json.org/>

**Parameters**

**indent** [int, optional, default=None] If not None, will pretty-print with specified number of spaces for indentation

**Returns**

**serialized** [str] A JSON serialized representation of the object

**to messagepack(self)**

Return a MessagePack representation.

Specification: <https://msgpack.org/index.html>

**Returns**

**serialized** [bytes] A MessagePack-encoded bytes serialized representation of the object

**to\_pickle(self)**

Return a pickle serialized representation.

**Warning:** This is not recommended for safe, stable storage since the pickle specification may change between Python versions.

**Returns**

**serialized** [str] A pickled representation of the object

**to\_toml(self)**

Return a TOML serialized representation.

Specification: <https://github.com/toml-lang/toml>

**Returns**

**serialized** [str] A TOML serialized representation of the object

**to\_xml(self, indent=2)**

Return an XML representation.

Specification: <https://www.w3.org/XML/>

**Parameters**

**indent** [int, optional, default=2] If not None, will pretty-print with specified number of spaces for indentation

**Returns**

**serialized** [bytes] A MessagePack-encoded bytes serialized representation.

**to\_yaml(self)**

Return a YAML serialized representation.

Specification: <http://yaml.org/>

**Returns**

**serialized** [str] A YAML serialized representation of the object

## openforcefield.topology.Bond

```
class openforcefield.topology.Bond(atom1, atom2, bond_order, is_aromatic, fractional_bond_order=None, stereochemistry=None)
```

Chemical bond representation.

**Warning:** This API is experimental and subject to change.

**Attributes**

**atom1, atom2** [openforcefield.topology.Atom] Atoms involved in the bond

**bondtype** [int] Discrete bond type representation for the Open Forcefield aromaticity model TODO: Do we want to pin ourselves to a single standard aromaticity model?

**type** [str] String based bond type  
**order** [int] Integral bond order  
**fractional\_bond\_order** [float, optional] Fractional bond order, or None.  
.. warning :: This API is experimental and subject to change.

## Methods

<code>from_bson(serialized)</code>	Instantiate an object from a BSON serialized representation.
<code>from_dict(molecule, d)</code>	Create a Bond from a dict representation.
<code>from_json(serialized)</code>	Instantiate an object from a JSON serialized representation.
<code>from_messagepack(serialized)</code>	Instantiate an object from a MessagePack serialized representation.
<code>from_pickle(serialized)</code>	Instantiate an object from a pickle serialized representation.
<code>from_toml(serialized)</code>	Instantiate an object from a TOML serialized representation.
<code>from_xml(serialized)</code>	Instantiate an object from an XML serialized representation.
<code>from_yaml(serialized)</code>	Instantiate from a YAML serialized representation.
<code>to_bson(self)</code>	Return a BSON serialized representation.
<code>to_dict(self)</code>	Return a dict representation of the bond.
<code>to_json(self[, indent])</code>	Return a JSON serialized representation.
<code>to_messagepack(self)</code>	Return a MessagePack representation.
<code>to_pickle(self)</code>	Return a pickle serialized representation.
<code>to_toml(self)</code>	Return a TOML serialized representation.
<code>to_xml(self[, indent])</code>	Return an XML representation.
<code>to_yaml(self)</code>	Return a YAML serialized representation.

`__init__(self, atom1, atom2, bond_order, is_aromatic, fractional_bond_order=None, stereochemistry=None)`

Create a new chemical bond.

## Methods

<code>__init__(self, atom1, atom2, bond_order, ...)</code>	Create a new chemical bond.
<code>from_bson(serialized)</code>	Instantiate an object from a BSON serialized representation.
<code>from_dict(molecule, d)</code>	Create a Bond from a dict representation.
<code>from_json(serialized)</code>	Instantiate an object from a JSON serialized representation.
<code>from_messagepack(serialized)</code>	Instantiate an object from a MessagePack serialized representation.
<code>from_pickle(serialized)</code>	Instantiate an object from a pickle serialized representation.

Continued on next page

Table 19 – continued from previous page

<code>from_toml(serialized)</code>	Instantiate an object from a TOML serialized representation.
<code>from_xml(serialized)</code>	Instantiate an object from an XML serialized representation.
<code>from_yaml(serialized)</code>	Instantiate from a YAML serialized representation.
<code>to_bson(self)</code>	Return a BSON serialized representation.
<code>to_dict(self)</code>	Return a dict representation of the bond.
<code>to_json(self[, indent])</code>	Return a JSON serialized representation.
<code>to_messagepack(self)</code>	Return a MessagePack representation.
<code>to_pickle(self)</code>	Return a pickle serialized representation.
<code>to_toml(self)</code>	Return a TOML serialized representation.
<code>to_xml(self[, indent])</code>	Return an XML representation.
<code>to_yaml(self)</code>	Return a YAML serialized representation.

## Attributes

<code>atom1</code>	
<code>atom1_index</code>	
<code>atom2</code>	
<code>atom2_index</code>	
<code>atoms</code>	
<code>bond_order</code>	
<code>fractional_bond_order</code>	
<code>is_aromatic</code>	
<code>molecule</code>	
<code>molecule_bond_index</code>	The index of this Bond within the the list of bonds in Molecules.
<code>stereochemistry</code>	

### `to_dict(self)`

Return a dict representation of the bond.

### `classmethod from_dict(molecule, d)`

Create a Bond from a dict representation.

### `property molecule_bond_index`

The index of this Bond within the the list of bonds in Molecules.

### `classmethod from_bson(serialized)`

Instantiate an object from a BSON serialized representation.

Specification: <http://bsonspec.org/>

## Parameters

`serialized` [bytes] A BSON serialized representation of the object

## Returns

`instance` [cls] An instantiated object

### `classmethod from_json(serialized)`

Instantiate an object from a JSON serialized representation.

Specification: <https://www.json.org/>

**Parameters**

**serialized** [str] A JSON serialized representation of the object

**Returns**

**instance** [cls] An instantiated object

**classmethod from\_messagepack(serialized)**

Instantiate an object from a MessagePack serialized representation.

Specification: <https://msgpack.org/index.html>

**Parameters**

**serialized** [bytes] A MessagePack-encoded bytes serialized representation

**Returns**

**instance** [cls] Instantiated object.

**classmethod from\_pickle(serialized)**

Instantiate an object from a pickle serialized representation.

**Warning:** This is not recommended for safe, stable storage since the pickle specification may change between Python versions.

**Parameters**

**serialized** [str] A pickled representation of the object

**Returns**

**instance** [cls] An instantiated object

**classmethod from\_toml(serialized)**

Instantiate an object from a TOML serialized representation.

Specification: <https://github.com/toml-lang/toml>

**Parameters**

**serialized** [str] A TOML serialized representation of the object

**Returns**

**instance** [cls] An instantiated object

**classmethod from\_xml(serialized)**

Instantiate an object from an XML serialized representation.

Specification: <https://www.w3.org/XML/>

**Parameters**

**serialized** [bytes] An XML serialized representation

**Returns**

**instance** [cls] Instantiated object.

**classmethod from\_yaml(serialized)**

Instantiate from a YAML serialized representation.

Specification: <http://yaml.org/>

**Parameters**

**serialized** [str] A YAML serialized representation of the object

**Returns**

**instance** [cls] Instantiated object

**to\_bson(self)**

Return a BSON serialized representation.

Specification: <http://bsonspec.org/>

**Returns**

**serialized** [bytes] A BSON serialized representation of the object

**to\_json(self, indent=None)**

Return a JSON serialized representation.

Specification: <https://www.json.org/>

**Parameters**

**indent** [int, optional, default=None] If not None, will pretty-print with specified number of spaces for indentation

**Returns**

**serialized** [str] A JSON serialized representation of the object

**to\_messagepack(self)**

Return a MessagePack representation.

Specification: <https://msgpack.org/index.html>

**Returns**

**serialized** [bytes] A MessagePack-encoded bytes serialized representation of the object

**to\_pickle(self)**

Return a pickle serialized representation.

**Warning:** This is not recommended for safe, stable storage since the pickle specification may change between Python versions.

**Returns**

**serialized** [str] A pickled representation of the object

**to\_toml(self)**

Return a TOML serialized representation.

Specification: <https://github.com/toml-lang/toml>

**Returns**

**serialized** [str] A TOML serialized representation of the object

**to\_xml(self, indent=2)**

Return an XML representation.

Specification: <https://www.w3.org/XML/>

## Parameters

**indent** [int, optional, default=2] If not None, will pretty-print with specified number of spaces for indentation

## Returns

**serialized** [bytes] A MessagePack-encoded bytes serialized representation.

**to\_yaml(self)**

Return a YAML serialized representation.

Specification: <http://yaml.org/>

## Returns

**serialized** [str] A YAML serialized representation of the object

## openforcefield.topology.VirtualSite

**class** openforcefield.topology.VirtualSite(atoms, charge\_increments=None, epsilon=None, sigma=None, rmin\_half=None, name=None)

A particle representing a virtual site whose position is defined in terms of Atom positions.

Note that chemical atoms are represented by the Atom.

**Warning:** This API is experimental and subject to change.

## Attributes

**atoms** Atoms on whose position this VirtualSite depends.

**charge\_increments** Charges taken from this VirtualSite's atoms and given to the VirtualSite

**epsilon** The VdW epsilon term of this VirtualSite

**molecule** The Molecule this atom is part of.

**molecule\_particle\_index** The index of this VirtualSite within the the list of particles in the parent Molecule.

**molecule\_virtual\_site\_index** The index of this VirtualSite within the list of virtual sites within Molecule Note that this can be different from particle\_index.

**name** The name of this VirtualSite

**rmin\_half** The VdW rmin\_half term of this VirtualSite

**sigma** The VdW sigma term of this VirtualSite

**type** The type of this VirtualSite (returns the class name as string)

## Methods

---

**from\_bson(serialized)** Instantiate an object from a BSON serialized representation.

---

**from\_dict(vsite\_dict)** Create a virtual site from a dict representation.

Continued on next page

Table 21 – continued from previous page

<code>from_json(serialized)</code>	Instantiate an object from a JSON serialized representation.
<code>from_messagepack(serialized)</code>	Instantiate an object from a MessagePack serialized representation.
<code>from_pickle(serialized)</code>	Instantiate an object from a pickle serialized representation.
<code>from_toml(serialized)</code>	Instantiate an object from a TOML serialized representation.
<code>from_xml(serialized)</code>	Instantiate an object from an XML serialized representation.
<code>from_yaml(serialized)</code>	Instantiate from a YAML serialized representation.
<code>to_bson(self)</code>	Return a BSON serialized representation.
<code>to_dict(self)</code>	Return a dict representation of the virtual site.
<code>to_json(self[, indent])</code>	Return a JSON serialized representation.
<code>to_messagepack(self)</code>	Return a MessagePack representation.
<code>to_pickle(self)</code>	Return a pickle serialized representation.
<code>to_toml(self)</code>	Return a TOML serialized representation.
<code>to_xml(self[, indent])</code>	Return an XML representation.
<code>to_yaml(self)</code>	Return a YAML serialized representation.

`__init__(self, atoms, charge_increments=None, epsilon=None, sigma=None, rmin_half=None, name=None)`

Base class for VirtualSites

#### Parameters

**atoms** [list of Atom of shape [N]] atoms[index] is the corresponding Atom for weights[index]

**charge\_increments** [list of floats of shape [N], optional, default=None] The amount of charge to remove from the VirtualSite's atoms and put in the VirtualSite. Indexing in this list should match the ordering in the atoms list. Default is None.

**sigma** [float, default=None] Sigma term for VdW properties of virtual site. Default is None.

**epsilon** [float] Epsilon term for VdW properties of virtual site. Default is None.

**rmin\_half** [float] Rmin\_half term for VdW properties of virtual site. Default is None.

**name** [string or None, default=None] The name of this virtual site. Default is None.

**virtual\_site\_type** [str] Virtual site type.

**name** [str or None, default=None] The name of this virtual site. Default is None

#### Methods

<code>__init__(self, atoms[, charge_increments, ...])</code>	Base class for VirtualSites
<code>from_bson(serialized)</code>	Instantiate an object from a BSON serialized representation.
<code>from_dict(vsite_dict)</code>	Create a virtual site from a dict representation.

Continued on next page

Table 22 – continued from previous page

<code>from_json(serialized)</code>	Instantiate an object from a JSON serialized representation.
<code>from_messagepack(serialized)</code>	Instantiate an object from a MessagePack serialized representation.
<code>from_pickle(serialized)</code>	Instantiate an object from a pickle serialized representation.
<code>from_toml(serialized)</code>	Instantiate an object from a TOML serialized representation.
<code>from_xml(serialized)</code>	Instantiate an object from an XML serialized representation.
<code>from_yaml(serialized)</code>	Instantiate from a YAML serialized representation.
<code>to_bson(self)</code>	Return a BSON serialized representation.
<code>to_dict(self)</code>	Return a dict representation of the virtual site.
<code>to_json(self[, indent])</code>	Return a JSON serialized representation.
<code>to_messagepack(self)</code>	Return a MessagePack representation.
<code>to_pickle(self)</code>	Return a pickle serialized representation.
<code>to_toml(self)</code>	Return a TOML serialized representation.
<code>to_xml(self[, indent])</code>	Return an XML representation.
<code>to_yaml(self)</code>	Return a YAML serialized representation.

## Attributes

<code>atoms</code>	Atoms on whose position this VirtualSite depends.
<code>charge_increments</code>	Charges taken from this VirtualSite's atoms and given to the VirtualSite
<code>epsilon</code>	The VdW epsilon term of this VirtualSite
<code>molecule</code>	The Molecule this atom is part of.
<code>molecule_particle_index</code>	The index of this VirtualSite within the the list of particles in the parent Molecule.
<code>molecule_virtual_site_index</code>	The index of this VirtualSite within the list of virtual sites within Molecule Note that this can be different from particle_index.
<code>name</code>	The name of this VirtualSite
<code>rmin_half</code>	The VdW rmin_half term of this VirtualSite
<code>sigma</code>	The VdW sigma term of this VirtualSite
<code>type</code>	The type of this VirtualSite (returns the class name as string)

### `to_dict(self)`

Return a dict representation of the virtual site.

### `classmethod from_dict(vsite_dict)`

Create a virtual site from a dict representation.

### `property molecule_virtual_site_index`

The index of this VirtualSite within the list of virtual sites within Molecule Note that this can be different from particle\_index.

### `property molecule_particle_index`

The index of this VirtualSite within the the list of particles in the parent Molecule. Note that this

can be different from molecule\_virtual\_site\_index.

**property atoms**

Atoms on whose position this VirtualSite depends.

**property charge\_increments**

Charges taken from this VirtualSite's atoms and given to the VirtualSite

**property epsilon**

The VdW epsilon term of this VirtualSite

**property sigma**

The VdW sigma term of this VirtualSite

**property rmin\_half**

The VdW rmin\_half term of this VirtualSite

**property name**

The name of this VirtualSite

**property type**

The type of this VirtualSite (returns the class name as string)

**classmethod from\_bson(serialized)**

Instantiate an object from a BSON serialized representation.

Specification: <http://bsonspec.org/>

**Parameters**

**serialized** [bytes] A BSON serialized representation of the object

**Returns**

**instance** [cls] An instantiated object

**classmethod from\_json(serialized)**

Instantiate an object from a JSON serialized representation.

Specification: <https://www.json.org/>

**Parameters**

**serialized** [str] A JSON serialized representation of the object

**Returns**

**instance** [cls] An instantiated object

**classmethod from\_messagepack(serialized)**

Instantiate an object from a MessagePack serialized representation.

Specification: <https://msgpack.org/index.html>

**Parameters**

**serialized** [bytes] A MessagePack-encoded bytes serialized representation

**Returns**

**instance** [cls] Instantiated object.

**classmethod from\_pickle(serialized)**

Instantiate an object from a pickle serialized representation.

**Warning:** This is not recommended for safe, stable storage since the pickle specification may change between Python versions.

**Parameters**

**serialized** [str] A pickled representation of the object

**Returns**

**instance** [cls] An instantiated object

**classmethod from\_toml(serialized)**

Instantiate an object from a TOML serialized representation.

Specification: <https://github.com/toml-lang/toml>

**Parameters**

**serialized** [str] A TOML serialized representation of the object

**Returns**

**instance** [cls] An instantiated object

**classmethod from\_xml(serialized)**

Instantiate an object from an XML serialized representation.

Specification: <https://www.w3.org/XML/>

**Parameters**

**serialized** [bytes] An XML serialized representation

**Returns**

**instance** [cls] Instantiated object.

**classmethod from\_yaml(serialized)**

Instantiate from a YAML serialized representation.

Specification: <http://yaml.org/>

**Parameters**

**serialized** [str] A YAML serialized representation of the object

**Returns**

**instance** [cls] Instantiated object

**property molecule**

The Molecule this atom is part of.

**to\_bson(self)**

Return a BSON serialized representation.

Specification: <http://bsonspec.org/>

**Returns**

**serialized** [bytes] A BSON serialized representation of the object

**to\_json(self, indent=None)**

Return a JSON serialized representation.

Specification: <https://www.json.org/>

**Parameters**

**indent** [int, optional, default=None] If not None, will pretty-print with specified number of spaces for indentation

**Returns**

**serialized** [str] A JSON serialized representation of the object

**to\_messagepack(self)**

Return a MessagePack representation.

Specification: <https://msgpack.org/index.html>

**Returns**

**serialized** [bytes] A MessagePack-encoded bytes serialized representation of the object

**to\_pickle(self)**

Return a pickle serialized representation.

**Warning:** This is not recommended for safe, stable storage since the pickle specification may change between Python versions.

**Returns**

**serialized** [str] A pickled representation of the object

**to\_toml(self)**

Return a TOML serialized representation.

Specification: <https://github.com/toml-lang/toml>

**Returns**

**serialized** [str] A TOML serialized representation of the object

**to\_xml(self, indent=2)**

Return an XML representation.

Specification: <https://www.w3.org/XML/>

**Parameters**

**indent** [int, optional, default=2] If not None, will pretty-print with specified number of spaces for indentation

**Returns**

**serialized** [bytes] A MessagePack-encoded bytes serialized representation.

**to\_yaml(self)**

Return a YAML serialized representation.

Specification: <http://yaml.org/>

**Returns**

**serialized** [str] A YAML serialized representation of the object

## 2.2 Forcefield typing tools

### 2.2.1 Chemical environments

Tools for representing and operating on chemical environments

**Warning:** This class is largely redundant with the same one in the Chemper package, and will likely be removed.

ChemicalEnvironment	Chemical environment abstract base class that matches an atom, bond, angle, etc.
---------------------	--

#### openforcefield.typing.chemistry.ChemicalEnvironment

```
class openforcefield.typing.chemistry.ChemicalEnvironment(smirks=None, label=None, replacements=None, toolkit='openeye')
```

Chemical environment abstract base class that matches an atom, bond, angle, etc.

**Warning:** This class is largely redundant with the same one in the Chemper package, and will likely be removed.

#### Methods

<code>Atom([ORtypes, ANDtypes, index, ring])</code>	Atom representation, which may have some ORtypes and ANDtypes properties.
<code>Bond([ORtypes, ANDtypes])</code>	Bond representation, which may have ORtype and ANDtype descriptors.
<code>addAtom(self, bondToAtom[, bondORtypes, ...])</code>	Add an atom to the specified target atom.
<code>asSMIRKS(self[, smarts])</code>	Returns a SMIRKS representation of the chemical environment
<code>getAlphaAtoms(self)</code>	Returns a list of atoms alpha to any indexed atom
<code>getAlphaBonds(self)</code>	Returns a list of Bond objects that connect
<code>getAtoms(self)</code>	
	<b>Returns</b>
<code>getBetaAtoms(self)</code>	Returns a list of atoms beta to any indexed atom
<code>getBetaBonds(self)</code>	Returns a list of Bond objects that connect
<code>getBond(self, atom1, atom2)</code>	Get bond between two atoms
<code>getBondOrder(self, atom)</code>	Returns minimum bond order around a given atom 0 if atom has no neighbors aromatic bonds count as 1.5 any bond counts as 1.0
<code>getBonds(self[, atom])</code>	

#### Parameters

---

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Table 25 – continued from previous page

<code>getComponentList(self, component_type[, ...])</code>	Returns a list of atoms or bonds matching the descriptor
<code>getIndexedAtoms(self)</code>	returns the list of Atom objects with an index
<code>getIndexedBonds(self)</code>	Returns a list of Bond objects that connect two indexed atoms
<code>getNeighbors(self, atom)</code>	Returns atoms that are bound to the given atom in the form of a list of Atom objects
<code>getType(self)</code>	Uses number of indexed atoms and bond connectivity to determine the type of chemical environment
<code>getUnindexedAtoms(self)</code>	returns a list of Atom objects that are not indexed
<code>getUnindexedBonds(self)</code>	Returns a list of Bond objects that connect
<code>getValence(self, atom)</code>	Returns the valence (number of neighboring atoms) around the given atom
<code>isAlpha(self, component)</code>	Takes an atom or bond and returns True if it is alpha to an indexed atom
<code>isBeta(self, component)</code>	Takes an atom or bond and returns True if it is beta to an indexed atom
<code>isIndexed(self, component)</code>	returns True if the atom or bond is indexed
<code>isUnindexed(self, component)</code>	returns True if the atom or bond is not indexed
<code>isValid(self[, smirks])</code>	Returns if the environment is valid, that is if it creates a parseable SMIRKS string.
<code>removeAtom(self, atom[, onlyEmpty])</code>	Remove the specified atom from the chemical environment.
<code>selectAtom(self[, descriptor])</code>	Select a random atom fitting the descriptor.
<code>selectBond(self[, descriptor])</code>	Select a random bond fitting the descriptor.
<code>validate(smirks[, ensure_valence_type, toolkit])</code>	Validate the provided SMIRKS string is valid, and if requested, tags atoms appropriate to the specified valence type.

`__init__(self, smirks=None, label=None, replacements=None, toolkit='openeye')`

Initialize a chemical environment abstract base class.

**smirks = string, optional** if smirks is not None, a chemical environment is built from the provided SMIRKS string**label = anything, optional** intended to be used to label this chemical environment could be a string, int, or float, or anything**replacements = list of lists, optional, [substitution, smarts] form for parsing SMIRKS**

## Methods

<code>__init__(self[, smirks, label, ...])</code>	Initialize a chemical environment abstract base class.
<code>addAtom(self, bondToAtom[, bondORtypes, ...])</code>	Add an atom to the specified target atom.
<code>asSMIRKS(self[, smarts])</code>	Returns a SMIRKS representation of the chemical environment
<code>getAlphaAtoms(self)</code>	Returns a list of atoms alpha to any indexed atom

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Table 26 – continued from previous page

<code>getAlphaBonds(self)</code>	Returns a list of Bond objects that connect
<b>Returns</b>	
<code>getBetaAtoms(self)</code>	Returns a list of atoms beta to any indexed atom
<code>getBetaBonds(self)</code>	Returns a list of Bond objects that connect
<code>getBond(self, atom1, atom2)</code>	Get bond between two atoms
<code>getBondOrder(self, atom)</code>	Returns minimum bond order around a given atom 0 if atom has no neighbors aromatic bonds count as 1.5 any bond counts as 1.0
<code>getBonds(self[, atom])</code>	
<b>Parameters</b>	
<code>getComponentList(self, component_type[, ...])</code>	Returns a list of atoms or bonds matching the descriptor
<code>getIndexedAtoms(self)</code>	returns the list of Atom objects with an index
<code>getIndexedBonds(self)</code>	Returns a list of Bond objects that connect two indexed atoms
<code>getNeighbors(self, atom)</code>	Returns atoms that are bound to the given atom in the form of a list of Atom objects
<code>getType(self)</code>	Uses number of indexed atoms and bond connectivity to determine the type of chemical environment
<code>getUnindexedAtoms(self)</code>	returns a list of Atom objects that are not indexed
<code>getUnindexedBonds(self)</code>	Returns a list of Bond objects that connect
<code>getValence(self, atom)</code>	Returns the valence (number of neighboring atoms) around the given atom
<code>isAlpha(self, component)</code>	Takes an atom or bond and returns True if it is alpha to an indexed atom
<code>isBeta(self, component)</code>	Takes an atom or bond and returns True if it is beta to an indexed atom
<code>isIndexed(self, component)</code>	returns True if the atom or bond is indexed
<code>isUnindexed(self, component)</code>	returns True if the atom or bond is not indexed
<code>isValid(self[, smirks])</code>	Returns if the environment is valid, that is if it creates a parseable SMIRKS string.
<code>removeAtom(self, atom[, onlyEmpty])</code>	Remove the specified atom from the chemical environment.
<code>selectAtom(self[, descriptor])</code>	Select a random atom fitting the descriptor.
<code>selectBond(self[, descriptor])</code>	Select a random bond fitting the descriptor.
<code>validate(smirks[, ensure_valence_type, toolkit])</code>	Validate the provided SMIRKS string is valid, and if requested, tags atoms appropriate to the specified valence type.

**class Atom(ORtypes=None, ANDtypes=None, index=None, ring=None)**

Atom representation, which may have some ORtypes and ANDtypes properties.

#### Attributes

**ORtypes** [list of tuples in the form (base, [list of decorators])] where bases and decorators are both strings The descriptor types that will be combined with logical OR

**ANDtypes** [list of string] The descriptor types that will be combined with logical

## AND

## Methods

<code>addANDtype(self, ANDtype)</code>	Adds ANDtype to the set for this atom.
<code>addORType(self, ORbase, ORdecorators)</code>	Adds ORtype to the set for this atom.
<code>asSMARTS(self)</code>	Return the atom representation as SMARTS.
<code>asSMIRKS(self)</code>	Return the atom representation as SMIRKS.
<code>getANDtypes(self)</code>	returns a copy of the list of ANDtypes for this atom
<code>getORTypes(self)</code>	returns a copy of the dictionary of ORtypes for this atom
<code>setANDtypes(self, newANDtypes)</code>	sets new ANDtypes for this atom
<code>setORTypes(self, newORTypes)</code>	sets new ORtypes for this atom

`asSMARTS(self)`

Return the atom representation as SMARTS.

## Returns

`smarts` [str]**The SMARTS string for this atom**`asSMIRKS(self)`

Return the atom representation as SMIRKS.

## Returns

`smirks` [str]**The SMIRKS string for this atom**`addORType(self, ORbase, ORdecorators)`

Adds ORtype to the set for this atom.

## Parameters

`ORbase: string, such as '#6'``ORdecorators: list of strings, such as ['X4','+0']``addANDtype(self, ANDtype)`

Adds ANDtype to the set for this atom.

## Parameters

`ANDtype: string` added to the list of ANDtypes for this atom`getORTypes(self)`

returns a copy of the dictionary of ORtypes for this atom

`setORTypes(self, newORTypes)`

sets new ORtypes for this atom

## Parameters

`newORTypes: list of tuples in the form (base, [ORdecorators])` for example:  
('#6', ['X4','H0','+0']) -> '#6X4H0+0'`getANDtypes(self)`

returns a copy of the list of ANDtypes for this atom

`setANDtypes(self, newANDtypes)`

sets new ANDtypes for this atom

## Parameters

`newANDtypes: list of strings` strings that will be AND'd together in a SMARTS

**class Bond(ORtypes=None, ANDtypes=None)**  
Bond representation, which may have ORtype and ANDtype descriptors.

#### Attributes

**ORTypes** [list of tuples of ORbases and ORdecorators] in form (base: [list of decorators]) The ORtype types that will be combined with logical OR

**ANDtypes** [list of string] The ANDtypes that will be combined with logical AND

#### Methods

<code>addANDtype(self, ANDtype)</code>	Adds ANDtype to the set for this atom.
<code>addORTYPE(self, ORbase, ORdecorators)</code>	Adds ORtype to the set for this atom.
<code>asSMARTS(self)</code>	Return the atom representation as SMARTS.
<code>asSMIRKS(self)</code>	

#### Returns

<code>getANDtypes(self)</code>	returns a copy of the list of ANDtypes for this atom
<code>getORTYPES(self)</code>	returns a copy of the dictionary of ORtypes for this atom
<code>getOrder(self)</code>	Returns a float for the order of this bond for multiple ORtypes or ~ it returns the minimum possible order the intended application is for checking valence around a given atom
<code>setANDtypes(self, newANDtypes)</code>	sets new ANDtypes for this atom
<code>setORTYPES(self, newORTYPES)</code>	sets new ORtypes for this atom

#### `asSMARTS(self)`

Return the atom representation as SMARTS.

#### Returns

**smarts** [str] The SMARTS string for just this atom

#### `asSMIRKS(self)`

#### Returns

**smarts** [str] The SMIRKS string for just this bond

#### `getOrder(self)`

Returns a float for the order of this bond for multiple ORtypes or ~ it returns the minimum possible order the intended application is for checking valence around a given atom

#### `addANDtype(self, ANDtype)`

Adds ANDtype to the set for this atom.

#### Parameters

**ANDtype: string** added to the list of ANDtypes for this atom

#### `addORTYPE(self, ORbase, ORdecorators)`

Adds ORtype to the set for this atom.

#### Parameters

**ORbase: string, such as '#6'**

**ORdecorators: list of strings, such as ['X4','+0']**

#### `getANDtypes(self)`

returns a copy of the list of ANDtypes for this atom

**getORtypes(self)**

returns a copy of the dictionary of ORtypes for this atom

**setANDtypes(self, newANDtypes)**

sets new ANDtypes for this atom

**Parameters**

**newANDtypes:** list of strings strings that will be AND'd together in a SMARTS

**setORtypes(self, newORtypes)**

sets new ORtypes for this atom

**Parameters**

**newORtypes:** list of tuples in the form (base, [ORdecorators]) for example:  
(' #'6', ['X4','H0','+0']) -> '#6X4H0+0'

**static validate(smirks, ensure\_valence\_type=None, toolkit='openeye')**

Validate the provided SMIRKS string is valid, and if requested, tags atoms appropriate to the specified valence type.

**Parameters**

**smirks** [str] The SMIRKS expression to validate

**ensure\_valence\_type** [str, optional, default=None] If specified, ensure the tagged atoms are appropriate to the specified valence type

This method will raise a :class:`SMIRKSParsingError` if the provided SMIRKS string is not valid.

**isValid(self, smirks=None)**

Returns if the environment is valid, that is if it creates a parseable SMIRKS string.

**asSMIRKS(self, smarts=False)**

Returns a SMIRKS representation of the chemical environment

**Parameters**

**smarts:** optional, boolean if True, returns a SMARTS instead of SMIRKS without index labels

**selectAtom(self, descriptor=None)**

Select a random atom fitting the descriptor.

**Parameters**

**descriptor:** optional, None None - returns any atom with equal probability int - will return an atom with that index 'Indexed' - returns a random indexed atom 'Unindexed' - returns a random unindexed atom 'Alpha' - returns a random alpha atom 'Beta' - returns a random beta atom

**Returns**

a single Atom object fitting the description

or None if no such atom exists

**getComponentList(self, component\_type, descriptor=None)**

Returns a list of atoms or bonds matching the descriptor

**Parameters**

**component\_type:** string: 'atom' or 'bond'

**descriptor:** string, optional 'all', 'Indexed', 'Unindexed', 'Alpha', 'Beta'

**selectBond(*self*, *descriptor*=*None*)**

Select a random bond fitting the descriptor.

**Parameters**

**descriptor: optional, None** None - returns any bond with equal probability int - will return an bond with that index 'Indexed' - returns a random indexed bond 'Unindexed' - returns a random unindexed bond 'Alpha' - returns a random alpha bond 'Beta' - returns a random beta bond

**Returns**

a single Bond object fitting the description

or None if no such atom exists

**addAtom(*self*, *bondToAtom*, *bondORtypes*=*None*, *bondANDtypes*=*None*, *newORtypes*=*None*, *newANDtypes*=*None*, *newAtomIndex*=*None*, *newAtomRing*=*None*, *beyondBeta*=*False*)**

Add an atom to the specified target atom.

**Parameters**

**bondToAtom: atom object, required** atom the new atom will be bound to

**bondORtypes: list of tuples, optional** strings that will be used for the ORtypes for the new bond

**bondANDtypes: list of strings, optional** strings that will be used for the ANDtypes for the new bond

**newORtypes: list of strings, optional** strings that will be used for the ORtypes for the new atom

**newANDtypes: list of strings, optional** strings that will be used for the ANDtypes for the new atom

**newAtomIndex: int, optional** integer label that could be used to index the atom in a SMIRKS string

**beyondBeta: boolean, optional** if True, allows bonding beyond beta position

**Returns**

**newAtom: atom object for the newly created atom**

**removeAtom(*self*, *atom*, *onlyEmpty*=*False*)**

Remove the specified atom from the chemical environment. if the atom is not indexed for the SMIRKS string or used to connect two other atoms.

**Parameters**

**atom: atom object, required** atom to be removed if it meets the conditions.

**onlyEmpty: boolean, optional** True only an atom with no ANDtypes and 1 ORtype can be removed

**Returns**

**Boolean True: atom was removed, False: atom was not removed**

**getAtoms(*self*)****Returns**

**list of atoms in the environment**

**getBonds(*self*, *atom*=*None*)**

**Parameters**

**atom:** Atom object, optional, returns bonds connected to atom  
returns all bonds in fragment if atom is None

**Returns**

a complete list of bonds in the fragment

**getBond(self, atom1, atom2)**

Get bond between two atoms

**Parameters**

**atom1 and atom2:** atom objects

**Returns**

bond object between the atoms or None if no bond there

**getIndexedAtoms(self)**

returns the list of Atom objects with an index

**getUnindexedAtoms(self)**

returns a list of Atom objects that are not indexed

**getAlphaAtoms(self)**

Returns a list of atoms alpha to any indexed atom that are not also indexed

**getBetaAtoms(self)**

Returns a list of atoms beta to any indexed atom that are not alpha or indexed atoms

**getIndexedBonds(self)**

Returns a list of Bond objects that connect two indexed atoms

**getUnindexedBonds(self)**

Returns a list of Bond objects that connect an indexed atom to an unindexed atom or two unindexed atoms

**getAlphaBonds(self)**

Returns a list of Bond objects that connect an indexed atom to alpha atoms

**getBetaBonds(self)**

Returns a list of Bond objects that connect alpha atoms to beta atoms

**isAlpha(self, component)**

Takes an atom or bond and returns True if it is alpha to an indexed atom

**isUnindexed(self, component)**

returns True if the atom or bond is not indexed

**isIndexed(self, component)**

returns True if the atom or bond is indexed

**isBeta(self, component)**

Takes an atom or bond and returns True if it is beta to an indexed atom

**getType(self)**

Uses number of indexed atoms and bond connectivity to determine the type of chemical environment

**Returns**

**chemical environment type:** ‘Atom’, ‘Bond’, ‘Angle’, ‘ProperTorsion’, ‘ImproperTorsion’ None if number of indexed atoms is 0 or > 4

**getNeighbors(self, atom)**

Returns atoms that are bound to the given atom in the form of a list of Atom objects

**getValence(self, atom)**

Returns the valence (number of neighboring atoms) around the given atom

**getBondOrder(self, atom)**

Returns minimum bond order around a given atom 0 if atom has no neighbors aromatic bonds count as 1.5 any bond counts as 1.0

## 2.2.2 Forcefield typing engines

Engines for applying parameters to chemical systems

### The SMIRks-Native Open Force Field (SMIRNOFF)

A reference implementation of the SMIRNOFF specification for parameterizing biomolecular systems

#### ForceField

The ForceField class is a primary part of the top-level toolkit API. ForceField objects are initialized from SMIRNOFF data sources (e.g. an OFFXML file). For a basic example of system creation using a ForceField, see examples/SMIRNOFF\_simulation.

---

`ForceField`

A factory that assigns SMIRNOFF parameters to a molecular system

---

`openforcefield.typing.engines.smirnoff.forcefield.ForceField`

```
class openforcefield.typing.engines.smirnoff.forcefield.ForceField(*sources, parameter_handler_classes=None, parameter_io_handler_classes=None, disable_version_check=False, allow_cosmetic_attributes=False)
```

A factory that assigns SMIRNOFF parameters to a molecular system

`ForceField` is a factory that constructs an OpenMM `simtk.openmm.System` object from a `openforcefield.topology.Topology` object defining a (bio)molecular system containing one or more molecules.

When a `ForceField` object is created from one or more specified SMIRNOFF serialized representations, all ParameterHandler subclasses currently imported are identified and registered to handle different sections of the SMIRNOFF force field definition file(s).

All ParameterIOHandler subclasses currently imported are identified and registered to handle different serialization formats (such as XML).

The force field definition is processed by these handlers to populate the ForceField object model data structures that can easily be manipulated via the API:

Processing a Topology object defining a chemical system will then call all :class`ParameterHandler` objects in an order guaranteed to satisfy the declared processing order constraints of each :class`ParameterHandler`.

## Examples

Create a new ForceField containing the smirnoff99Frosst parameter set:

```
>>> from openforcefield.typing.engines.smirnoff import ForceField
>>> forcefield = ForceField('test_forcefields/smirnoff99Frosst.offxml')
```

Create an OpenMM system from a `openforcefield.topology.Topology` object:

```
>>> from openforcefield.topology import Molecule, Topology
>>> ethanol = Molecule.from_smiles('CCO')
>>> topology = Topology.from_molecules(molecules=[ethanol])
>>> system = forcefield.create_openmm_system(topology)
```

Modify the long-range electrostatics method:

```
>>> forcefield.get_parameter_handler('Electrostatics').method = 'PME'
```

Inspect the first few vdW parameters:

```
>>> low_precedence_parameters = forcefield.get_parameter_handler('vdW').parameters[0:3]
```

Retrieve the vdW parameters by SMIRKS string and manipulate it:

```
>>> parameter = forcefield.get_parameter_handler('vdW').parameters['[#1:1]-[#7:1]
>>> parameter.rmin_half += 0.1 * unit.angstroms
>>> parameter.epsilon *= 1.02
```

Make a child vdW type more specific (checking modified SMIRKS for validity):

```
>>> forcefield.get_parameter_handler('vdW').parameters[-1].smirks += ' ~[#53]'
```

**Warning:** While we check whether the modified SMIRKS is still valid and has the appropriate valence type, we currently don't check whether the typing remains hierarchical, which could result in some types no longer being assignable because more general types now come *below* them and preferentially match.

Delete a parameter:

```
>>> del forcefield.get_parameter_handler('vdW').parameters['[#1:1]-[#6X4]']
```

Insert a parameter at a specific point in the parameter tree:

```
>>> from openforcefield.typing.engines.smirnoff import vdWHandler
>>> new_parameter = vdWHandler.vdWType(smirks='[*:1]', epsilon=0.0157*unit.kilocalories_per_mole,_
>>> rmin_half=0.6000*unit.angstroms)
>>> forcefield.get_parameter_handler('vdW').parameters.insert(0, new_parameter)
```

**Warning:** We currently don't check whether removing a parameter could accidentally remove the root type, so it's possible to no longer type all molecules this way.

## Attributes

**parameters** [dict of str] parameters[tagname] is the instantiated ParameterHandler class that handles parameters associated with the force tagname. This is the primary means of retrieving and modifying parameters, such as parameters['vdW'][0].  
 $\sigma \approx 1.1$

**parameter\_object\_handlers** [dict of str] Registered list of ParameterHandler classes that will handle different forcefield tags to create the parameter object model. parameter\_object\_handlers[tagname] is the ParameterHandler that will be instantiated to process the force field definition section tagname. ParameterHandler classes are registered when the ForceField object is created, but can be manipulated afterwards.

**parameter\_io\_handlers** [dict of str] Registered list of ParameterIOHandler classes that will handle serializing/deserializing the parameter object model to string or file representations, such as XML. parameter\_io\_handlers[iotype] is the ParameterHandler that will be instantiated to process the serialization scheme iotype. ParameterIOHandler classes are registered when the ForceField object is created, but can be manipulated afterwards.

## Methods

<code>create_openmm_system(self, topology, \*\*kwargs)</code>		Create an OpenMM System representing the interactions for the specified Topology with the current force field
<code>create_parmed_structure(self, topology, ...)</code>		Create a ParmEd Structure object representing the interactions for the specified Topology with the current force field
<code>get_parameter_handler(self, tagname[, ...])</code>		Retrieve the parameter handlers associated with the provided tagname.
<code>get_parameter_io_handler(self, io_format)</code>		Retrieve the parameter handlers associated with the provided tagname.
<code>label_molecules(self, topology)</code>		Return labels for a list of molecules corresponding to parameters from this force field.
<code>parse_smirnoff_from_source(self, source)</code>		Reads a SMIRNOFF data structure from a source, which can be one of many types.
<code>parse_sources(self, sources[, ...])</code>		Parse a SMIRNOFF force field definition.
<code>register_parameter_handler(self, ...)</code>		Register a new ParameterHandler for a specific tag, making it available for lookup in the ForceField.
<code>register_parameter_io_handler(self, ...)</code>		Register a new ParameterIOHandler, making it available for lookup in the ForceField.
<code>to_file(self, filename[, io_format, ...])</code>		Write this Forcefield and all its associated parameters to a string in a given format which complies with the SMIRNOFF spec.

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Table 30 – continued from previous page

<code>to_string(self[, io_format, ...])</code>	Write this Forcefield and all its associated parameters to a string in a given format which complies with the SMIRNOFF spec.
--	--

`__init__(self, *sources, parameter_handler_classes=None, parameter_io_handler_classes=None, disable_version_check=False, allow_cosmetic_attributes=False)`  
Create a new `ForceField` object from one or more SMIRNOFF parameter definition files.

### Parameters

**sources** [string or file-like object or open file handle or URL (or iterable of these)]

A list of files defining the SMIRNOFF force field to be loaded. Currently, only the [SMIRNOFF XML format](#) is supported. Each entry may be an absolute file path, a path relative to the current working directory, a path relative to this module's data subdirectory (for built in force fields), or an open file-like object with a `read()` method from which the forcefield XML data can be loaded. If multiple files are specified, any top-level tags that are repeated will be merged if they are compatible, with files appearing later in the sequence resulting in parameters that have higher precedence. Support for multiple files is primarily intended to allow solvent parameters to be specified by listing them last in the sequence.

**parameter\_handler\_classes** [iterable of ParameterHandler classes, optional, default=None] If not None, the specified set of ParameterHandler classes will be instantiated to create the parameter object model. By default, all imported subclasses of ParameterHandler are automatically registered.

**parameter\_io\_handler\_classes** [iterable of ParameterIOHandler classes] If not None, the specified set of ParameterIOHandler classes will be used to parse/generate serialized parameter sets. By default, all imported subclasses of ParameterIOHandler are automatically registered.

**disable\_version\_check** [bool, optional, default=False] If True, will disable checks against the current highest supported forcefield version. This option is primarily intended for forcefield development.

**allow\_cosmetic\_attributes** [bool, optional. Default = False] Whether to retain non-spec kwargs from data sources.

### Examples

Load one SMIRNOFF parameter set in XML format (searching the package data directory by default, which includes some standard parameter sets):

```
>>> forcefield = ForceField('test_forcefields/smirkoff99Frosst.offxml')
```

Load multiple SMIRNOFF parameter sets:

```
forcefield = ForceField('test_forcefields/smirkoff99Frosst.offxml', 'test_forcefields/tip3p.offxml')
```

Load a parameter set from a string:

```
>>> offxml = '<SMIRNOFF version="0.2" aromaticity_model="OEArroModel_MDL"/>'  
>>> forcefield = ForceField(offxml)
```

## Methods

<code>__init__(self, \*sources[, ...])</code>	Create a new ForceField object from one or more SMIRNOFF parameter definition files.
<code>create_openmm_system(self, topology, \*\*kwargs)</code>	Create an OpenMM System representing the interactions for the specified Topology with the current force field
<code>create_parmed_structure(self, topology, ...)</code>	Create a ParmEd Structure object representing the interactions for the specified Topology with the current force field
<code>get_parameter_handler(self, tagname[, ...])</code>	Retrieve the parameter handlers associated with the provided tagname.
<code>get_parameter_io_handler(self, io_format)</code>	Retrieve the parameter handlers associated with the provided tagname.
<code>label_molecules(self, topology)</code>	Return labels for a list of molecules corresponding to parameters from this force field.
<code>parse_smirnoff_from_source(self, source)</code>	Reads a SMIRNOFF data structure from a source, which can be one of many types.
<code>parse_sources(self, sources[, ...])</code>	Parse a SMIRNOFF force field definition.
<code>register_parameter_handler(self, ...)</code>	Register a new ParameterHandler for a specific tag, making it available for lookup in the ForceField.
<code>register_parameter_io_handler(self, ...)</code>	Register a new ParameterIOHandler, making it available for lookup in the ForceField.
<code>to_file(self, filename[, io_format, ...])</code>	Write this Forcefield and all its associated parameters to a string in a given format which complies with the SMIRNOFF spec.
<code>to_string(self[, io_format, ...])</code>	Write this Forcefield and all its associated parameters to a string in a given format which complies with the SMIRNOFF spec.

## Attributes

<code>author</code>	Returns the author data for this ForceField object.
<code>date</code>	Returns the date data for this ForceField object.

### `property author`

Returns the author data for this ForceField object. If not defined in any loaded files, this will be None.

#### Returns

`author` [str] The author data for this forcefield.

### `property date`

Returns the date data for this ForceField object. If not defined in any loaded files, this will be None.

#### Returns

`date` [str] The date data for this forcefield.

### `register_parameter_handler(self, parameter_handler)`

Register a new ParameterHandler for a specific tag, making it available for lookup in the ForceField.

**Warning:** This API is experimental and subject to change.

#### Parameters

**parameter\_handler** [A ParameterHandler object] The ParameterHandler to register.  
The TAGNAME attribute of this object will be used as the key for registration.

**register\_parameter\_io\_handler(self, parameter\_io\_handler)**

Register a new ParameterIOHandler, making it available for lookup in the ForceField.

**Warning:** This API is experimental and subject to change.

#### Parameters

**parameter\_io\_handler** [A ParameterIOHandler object] The ParameterIOHandler to register. The FORMAT attribute of this object will be used to associate it to a file format/suffix.

**get\_parameter\_handler(self, tagname, handler\_kwargs=None, allow\_cosmetic\_attributes=False)**

Retrieve the parameter handlers associated with the provided tagname.

If the parameter handler has not yet been instantiated, it will be created and returned. If a parameter handler object already exists, it will be checked for compatibility and an Exception raised if it is incompatible with the provided kwargs. If compatible, the existing ParameterHandler will be returned.

#### Parameters

**tagname** [str] The name of the parameter to be handled.

**handler\_kwargs** [dict, optional. Default = None] Dict to be passed to the handler for construction or checking compatibility. If this is None and no existing ParameterHandler exists for the desired tag, a handler will be initialized with all default values. If this is None and a handler for the desired tag exists, the existing ParameterHandler will be returned.

**allow\_cosmetic\_attributes** [bool, optional. Default = False] Whether to permit non-spec kwargs in smirnoff\_data.

#### Returns

**handler** [An openforcefield.engines.typing.smirnoff.ParameterHandler]

#### Raises

**KeyError** if there is no ParameterHandler for the given tagname

**get\_parameter\_io\_handler(self, io\_format)**

Retrieve the parameter handlers associated with the provided tagname. If the parameter IO handler has not yet been instantiated, it will be created.

#### Parameters

**io\_format** [str] The name of the io format to be handled.

**Returns**

**io\_handler** [An openforcefield.engines.typing.smirnoff.ParameterIOHandler]

**Raises**

**KeyError** if there is no ParameterIOHandler for the given tagname

**parse\_sources(self, sources, allow\_cosmetic\_attributes=True)**

Parse a SMIRNOFF force field definition.

**Parameters**

**sources** [string or file-like object or open file handle or URL (or iterable of these)]

A list of files defining the SMIRNOFF force field to be loaded. Currently, only [the SMIRNOFF XML format](#) is supported. Each entry may be an absolute file path, a path relative to the current working directory, a path relative to this module's data subdirectory (for built in force fields), or an open file-like object with a `read()` method from which the forcefield XML data can be loaded. If multiple files are specified, any top-level tags that are repeated will be merged if they are compatible, with files appearing later in the sequence resulting in parameters that have higher precedence. Support for multiple files is primarily intended to allow solvent parameters to be specified by listing them last in the sequence.

**allow\_cosmetic\_attributes** [bool, optional. Default = False] Whether to permit non-spec kwargs present in the source.

**.. notes ::**

- New SMIRNOFF sections are handled independently, as if they were specified in the same file.
- If a SMIRNOFF section that has already been read appears again, its definitions are appended to the end of the previously-read definitions if the sections are configured with compatible attributes; otherwise, an `IncompatibleTagException` is raised.

**parse\_smirnoff\_from\_source(self, source)**

Reads a SMIRNOFF data structure from a source, which can be one of many types.

**Parameters**

**source** [str or bytes] sources : string or file-like object or open file handle or URL

(or iterable of these) A list of files defining the SMIRNOFF force field to be loaded. Currently, only [the SMIRNOFF XML format](#) is supported. Each entry may be an absolute file path, a path relative to the current working directory, a path relative to this module's data subdirectory (for built in force fields), or an open file-like object with a `read()` method from which the forcefield XML data can be loaded.

**Returns**

**smirnoff\_data** [OrderedDict] A representation of a SMIRNOFF-format data structure. Begins at top-level 'SMIRNOFF' key.

**to\_string(self, io\_format='XML', discard\_cosmetic\_attributes=False)**

Write this Forcefield and all its associated parameters to a string in a given format which complies with the SMIRNOFF spec.

**Parameters**

**io\_format** [str or ParameterIOHandler, optional. Default='XML'] The serialization format to write to

**discard\_cosmetic\_attributes** [bool, default=False] Whether to discard any non-spec attributes stored in the ForceField.

#### Returns

**forcefield\_string** [str] The string representation of the serialized forcefield

**to\_file(self, filename, io\_format=None, discard\_cosmetic\_attributes=False)**

Write this Forcefield and all its associated parameters to a string in a given format which complies with the SMIRNOFF spec.

#### Parameters

**filename** [str] The filename to write to

**io\_format** [str or ParameterIOHandler, optional. Default=None] The serialization format to write out. If None, will attempt to be inferred from the filename.

**discard\_cosmetic\_attributes** [bool, default=False] Whether to discard any non-spec attributes stored in the ForceField.

#### Returns

**forcefield\_string** [str] The string representation of the serialized forcefield

**create\_openmm\_system(self, topology, \*\*kwargs)**

Create an OpenMM System representing the interactions for the specified Topology with the current force field

#### Parameters

**topology** [openforcefield.topology.Topology] The Topology corresponding to the system to be parameterized

**charge\_from\_molecules** [List[openforcefield.molecule.Molecule], optional] If specified, partial charges will be taken from the given molecules instead of being determined by the force field.

#### Returns

**system** [simtk.openmm.System] The newly created OpenMM System corresponding to the specified topology

**create\_parmed\_structure(self, topology, positions, \*\*kwargs)**

Create a ParmEd Structure object representing the interactions for the specified Topology with the current force field

This method creates a [ParmEd](#) Structure object containing a topology, positions, and parameters.

#### Parameters

**topology** [openforcefield.topology.Topology] The Topology corresponding to the System object to be created.

**positions** [simtk.unit.Quantity of dimension (natoms,3) with units compatible with angstroms] The positions corresponding to the System object to be created

#### Returns

**structure** [parmed.Structure] The newly created parmed.Structure object

**label\_molecules(self, topology)**

Return labels for a list of molecules corresponding to parameters from this force field. For each molecule, a dictionary of force types is returned, and for each force type, each force term is provided with the atoms involved, the parameter id assigned, and the corresponding SMIRKS.

## Parameters

**topology** [openforcefield.topology.Topology] A Topology object containing one or more unique molecules to be labeled

## Returns

**molecule\_labels** [list] List of labels for unique molecules. Each entry in the list corresponds to one unique molecule in the Topology and is a dictionary keyed by force type, i.e., `molecule_labels[0]['HarmonicBondForce']` gives details for the harmonic bond parameters for the first molecule. Each element is a list of the form: [ ( [ atom1, ..., atomN], parameter\_id, SMIRKS), ... ].

## Parameter Type

ParameterType objects are representations of individual SMIRKS-based SMIRNOFF parameters. These are usually initialized during ForceField creation, and can be inspected and modified by users via the Python API. For more information, see examples/forcefield\_modification.

ParameterType	Base class for SMIRNOFF parameter types.
BondHandler.BondType	A SMIRNOFF bond type
AngleHandler.AngleType	A SMIRNOFF angle type.
ProperTorsionHandler.ProperTorsionType	A SMIRNOFF torsion type for proper torsions.
ImproperTorsionHandler.ImproperTorsionType	A SMIRNOFF torsion type for improper torsions.
vdWHandler.vdWType	A SMIRNOFF vdWForce type.

## openforcefield.typing.engines.smirnoff.parameters.ParameterType

```
class openforcefield.typing.engines.smirnoff.parameters.ParameterType(smirks=None, allow_cosmetic_attributes=False, **kwargs)
```

Base class for SMIRNOFF parameter types.

**Warning:** This API is experimental and subject to change.

### Attributes

`smirks`

### Methods

<code>add_cosmetic_attribute(self, attr_name, ...)</code>	Add a cosmetic attribute to this ParameterType object.
<code>delete_cosmetic_attribute(self, attr_name)</code>	Delete a cosmetic attribute from this ParameterType object.
<code>to_dict(self[, discard_cosmetic_attributes])</code>	Convert this ParameterType object to dict.

```
__init__(self, smirks=None, allow_cosmetic_attributes=False, **kwargs)
```

Create a ParameterType

## Parameters

**smirks** [str] The SMIRKS match for the provided parameter type.

**allow\_cosmetic\_attributes** [bool optional. Default = False] Whether to permit non-spec kwargs (“cosmetic attributes”). If True, non-spec kwargs will be stored as an attribute of this parameter which can be accessed and written out. Otherwise an exception will be raised.

## Methods

<code>__init__(self[, smirks, ...])</code>	Create a ParameterType
<code>add_cosmetic_attribute(self, attr_name, ...)</code>	Add a cosmetic attribute to this ParameterType object.
<code>delete_cosmetic_attribute(self, attr_name)</code>	Delete a cosmetic attribute from this ParameterType object.
<code>to_dict(self[, discard_cosmetic_attributes])</code>	Convert this ParameterType object to dict.

## Attributes

### smirks

**to\_dict(self, discard\_cosmetic\_attributes=False)**

Convert this ParameterType object to dict. A unit-bearing attribute ('X') will be converted to two dict entries, one ('X') containing the unitless value, and another ('X\_unit') containing a string representation of its unit.

#### Parameters

**discard\_cosmetic\_attributes** [bool, optional. Default = False] Whether to discard non-spec attributes of this ParameterType

#### Returns

**smirnoff\_dict** [dict] The SMIRNOFF-compliant dict representation of this ParameterType object.

**output\_units** [dict[str: simtk.unit.Unit]] A mapping from each simtk.unit.Quantity-valued ParameterType attribute to the unit it was converted to during serialization.

**add\_cosmetic\_attribute(self, attr\_name, attr\_value)**

Add a cosmetic attribute to this ParameterType object. This attribute will not have a functional effect on the object in the Open Force Field toolkit, but can be written out during output.

#### Parameters

**attr\_name** [str] Name of the attribute to define for this ParameterType object.

**attr\_value** [str] The value of the attribute to define for this ParameterType object.

**delete\_cosmetic\_attribute(self, attr\_name)**

Delete a cosmetic attribute from this ParameterType object.

#### Parameters

**attr\_name** [str] Name of the attribute to delete.

**openforcefield.typing.engines.smirnoff.parameters.BondHandler.BondType**

**class** openforcefield.typing.engines.smirnoff.parameters.BondHandler.BondType(\*\*kwargs)  
A SMIRNOFF bond type

**Warning:** This API is experimental and subject to change.

**Attributes****smirks****Methods**

<code>add_cosmetic_attribute(self, attr_name, ...)</code>	Add a cosmetic attribute to this ParameterType object.
<code>delete_cosmetic_attribute(self, attr_name)</code>	Delete a cosmetic attribute from this ParameterType object.
<code>to_dict(self[, discard_cosmetic_attributes])</code>	Convert this ParameterType object to dict.

**\_\_init\_\_(self, \*\*kwargs)**

Create a ParameterType

**Parameters**

**smirks** [str] The SMIRKS match for the provided parameter type.

**allow\_cosmetic\_attributes** [bool optional. Default = False] Whether to permit non-spec kwargs (“cosmetic attributes”). If True, non-spec kwargs will be stored as an attribute of this parameter which can be accessed and written out. Otherwise an exception will be raised.

**Methods**

<code>__init__(self, \*\*kwargs)</code>	Create a ParameterType
<code>add_cosmetic_attribute(self, attr_name, ...)</code>	Add a cosmetic attribute to this ParameterType object.
<code>delete_cosmetic_attribute(self, attr_name)</code>	Delete a cosmetic attribute from this ParameterType object.
<code>to_dict(self[, discard_cosmetic_attributes])</code>	Convert this ParameterType object to dict.

**Attributes**

---

**smirks**

---

**add\_cosmetic\_attribute(self, attr\_name, attr\_value)**

Add a cosmetic attribute to this ParameterType object. This attribute will not have a functional effect on the object in the Open Force Field toolkit, but can be written out during output.

**Parameters**

**attr\_name** [str] Name of the attribute to define for this ParameterType object.

**attr\_value** [str] The value of the attribute to define for this ParameterType object.

**delete\_cosmetic\_attribute(self, attr\_name)**  
Delete a cosmetic attribute from this ParameterType object.

**Parameters**

**attr\_name** [str] Name of the attribute to delete.

**to\_dict(self, discard\_cosmetic\_attributes=False)**  
Convert this ParameterType object to dict. A unit-bearing attribute ('X') will be converted to two dict entries, one (['X']) containing the unitless value, and another (['X\_unit']) containing a string representation of its unit.

**Parameters**

**discard\_cosmetic\_attributes** [bool, optional. Default = False] Whether to discard non-spec attributes of this ParameterType

**Returns**

**smirnoff\_dict** [dict] The SMIRNOFF-compliant dict representation of this ParameterType object.

**output\_units** [dict[str: simtk.unit.Unit]] A mapping from each simtk.unit.Quantity-valued ParameterType attribute to the unit it was converted to during serialization.

## openforcefield.typing.engines.smirnoff.parameters.AngleHandler.AngleType

**class** openforcefield.typing.engines.smirnoff.parameters.AngleHandler.AngleType(\*\*kwargs)  
A SMIRNOFF angle type.

**Warning:** This API is experimental and subject to change.

### Attributes

**smirks**

### Methods

<code>add_cosmetic_attribute(self, attr_name, ...)</code>	Add a cosmetic attribute to this ParameterType object.
<code>delete_cosmetic_attribute(self, attr_name)</code>	Delete a cosmetic attribute from this ParameterType object.
<code>to_dict(self[, discard_cosmetic_attributes])</code>	Convert this ParameterType object to dict.

**\_\_init\_\_(self, \*\*kwargs)**  
Create a ParameterType

**Parameters**

**smirks** [str] The SMIRKS match for the provided parameter type.

**allow\_cosmetic\_attributes** [bool optional. Default = False] Whether to permit non-spec kwargs ("cosmetic attributes"). If True, non-spec kwargs will be stored as an

attribute of this parameter which can be accessed and written out. Otherwise an exception will be raised.

## Methods

<code>__init__(self, \*\\*kwargs)</code>	Create a ParameterType
<code>add_cosmetic_attribute(self, attr_name, ...)</code>	Add a cosmetic attribute to this ParameterType object.
<code>delete_cosmetic_attribute(self, attr_name)</code>	Delete a cosmetic attribute from this ParameterType object.
<code>to_dict(self[, discard_cosmetic_attributes])</code>	Convert this ParameterType object to dict.

## Attributes

---

### smirks

---

#### `add_cosmetic_attribute(self, attr_name, attr_value)`

Add a cosmetic attribute to this ParameterType object. This attribute will not have a functional effect on the object in the Open Force Field toolkit, but can be written out during output.

##### Parameters

`attr_name` [str] Name of the attribute to define for this ParameterType object.

`attr_value` [str] The value of the attribute to define for this ParameterType object.

#### `delete_cosmetic_attribute(self, attr_name)`

Delete a cosmetic attribute from this ParameterType object.

##### Parameters

`attr_name` [str] Name of the attribute to delete.

#### `to_dict(self, discard_cosmetic_attributes=False)`

Convert this ParameterType object to dict. A unit-bearing attribute ('X') will be converted to two dict entries, one ([‘X’] containing the unitless value, and another ([‘X\_unit’]) containing a string representation of its unit.

##### Parameters

`discard_cosmetic_attributes` [bool, optional. Default = False] Whether to discard non-spec attributes of this ParameterType

##### Returns

`smirnoff_dict` [dict] The SMIRNOFF-compliant dict representation of this ParameterType object.

`output_units` [dict[str: simtk.unit.Unit]] A mapping from each simtk.unit.Quantity-valued ParameterType attribute to the unit it was converted to during serialization.

## `openforcefield.typing.engines.smirnoff.parameters.ProperTorsionHandler.ProperTorsionType`

`class openforcefield.typing.engines.smirnoff.parameters.ProperTorsionHandler.ProperTorsionType(**kwargs)`  
A SMIRNOFF torsion type for proper torsions.

**Warning:** This API is experimental and subject to change.

## Attributes

**smirks**

## Methods

<code>add_cosmetic_attribute(self, attr_name, ...)</code>	Add a cosmetic attribute to this ParameterType object.
<code>delete_cosmetic_attribute(self, attr_name)</code>	Delete a cosmetic attribute from this ParameterType object.
<code>to_dict(self[, discard_cosmetic_attributes])</code>	Convert this ParameterType object to dict.

**`__init__(self, **kwargs)`**

Create a ParameterType

### Parameters

**smirks** [str] The SMIRKS match for the provided parameter type.

**allow\_cosmetic\_attributes** [bool optional. Default = False] Whether to permit non-spec kwargs (“cosmetic attributes”). If True, non-spec kwargs will be stored as an attribute of this parameter which can be accessed and written out. Otherwise an exception will be raised.

## Methods

<code>__init__(self, \*\*kwargs)</code>	Create a ParameterType
<code>add_cosmetic_attribute(self, attr_name, ...)</code>	Add a cosmetic attribute to this ParameterType object.
<code>delete_cosmetic_attribute(self, attr_name)</code>	Delete a cosmetic attribute from this ParameterType object.
<code>to_dict(self[, discard_cosmetic_attributes])</code>	Convert this ParameterType object to dict.

## Attributes

**smirks**

**`add_cosmetic_attribute(self, attr_name, attr_value)`**

Add a cosmetic attribute to this ParameterType object. This attribute will not have a functional effect on the object in the Open Force Field toolkit, but can be written out during output.

### Parameters

**attr\_name** [str] Name of the attribute to define for this ParameterType object.

**attr\_value** [str] The value of the attribute to define for this ParameterType object.

**`delete_cosmetic_attribute(self, attr_name)`**

Delete a cosmetic attribute from this ParameterType object.

## Parameters

**attr\_name** [str] Name of the attribute to delete.

**to\_dict**(*self*, *discard\_cosmetic\_attributes=False*)

Convert this ParameterType object to dict. A unit-bearing attribute ('X') will be converted to two dict entries, one ('X') containing the unitless value, and another ('X\_unit') containing a string representation of its unit.

## Parameters

**discard\_cosmetic\_attributes** [bool, optional. Default = False] Whether to discard non-spec attributes of this ParameterType

## Returns

**smirnoff\_dict** [dict] The SMIRNOFF-compliant dict representation of this ParameterType object.

**output\_units** [dict[str: simtk.unit.Unit]] A mapping from each simtk.unit.Quantity-valued ParameterType attribute to the unit it was converted to during serialization.

## openforcefield.typing.engines.smirnoff.parameters.ImproperTorsionHandler.ImproperTorsionType

**class** openforcefield.typing.engines.smirnoff.parameters.ImproperTorsionHandler.**ImproperTorsionType**(\*\**kwargs*)  
A SMIRNOFF torsion type for improper torsions.

**Warning:** This API is experimental and subject to change.

## Attributes

**smirks**

## Methods

<code>add_cosmetic_attribute(self, attr_name, ...)</code>	Add a cosmetic attribute to this ParameterType object.
<code>delete_cosmetic_attribute(self, attr_name)</code>	Delete a cosmetic attribute from this ParameterType object.
<code>to_dict(self[, discard_cosmetic_attributes])</code>	Convert this ParameterType object to dict.

**\_\_init\_\_(self, \*\**kwargs*)**

Create a ParameterType

## Parameters

**smirks** [str] The SMIRKS match for the provided parameter type.

**allow\_cosmetic\_attributes** [bool optional. Default = False] Whether to permit non-spec kwargs ("cosmetic attributes"). If True, non-spec kwargs will be stored as an attribute of this parameter which can be accessed and written out. Otherwise an exception will be raised.

## Methods

<code>__init__(self, \*\*kwargs)</code>	Create a ParameterType
<code>add_cosmetic_attribute(self, attr_name, ...)</code>	Add a cosmetic attribute to this ParameterType object.
<code>delete_cosmetic_attribute(self, attr_name)</code>	Delete a cosmetic attribute from this ParameterType object.
<code>to_dict(self[, discard_cosmetic_attributes])</code>	Convert this ParameterType object to dict.

## Attributes

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`smirks`

---

`add_cosmetic_attribute(self, attr_name, attr_value)`

Add a cosmetic attribute to this ParameterType object. This attribute will not have a functional effect on the object in the Open Force Field toolkit, but can be written out during output.

### Parameters

`attr_name` [str] Name of the attribute to define for this ParameterType object.

`attr_value` [str] The value of the attribute to define for this ParameterType object.

`delete_cosmetic_attribute(self, attr_name)`

Delete a cosmetic attribute from this ParameterType object.

### Parameters

`attr_name` [str] Name of the attribute to delete.

`to_dict(self, discard_cosmetic_attributes=False)`

Convert this ParameterType object to dict. A unit-bearing attribute ('X') will be converted to two dict entries, one ('X') containing the unitless value, and another ('X\_unit') containing a string representation of its unit.

### Parameters

`discard_cosmetic_attributes` [bool, optional. Default = False] Whether to discard non-spec attributes of this ParameterType

### Returns

`smirnoff_dict` [dict] The SMIRNOFF-compliant dict representation of this ParameterType object.

`output_units` [dict[str: simtk.unit.Unit]] A mapping from each simtk.unit.Quantity-valued ParameterType attribute to the unit it was converted to during serialization.

[openforcefield.typing.engines.smirnoff.parameters.vdWHandler.vdWType](#)

`class openforcefield.typing.engines.smirnoff.parameters.vdWHandler.vdWType(**kwargs)`  
A SMIRNOFF vdWForce type.

**Warning:** This API is experimental and subject to change.

## Attributes

**smirks**

## Methods

<code>add_cosmetic_attribute(self, attr_name, ...)</code>	Add a cosmetic attribute to this ParameterType object.
<code>delete_cosmetic_attribute(self, attr_name)</code>	Delete a cosmetic attribute from this ParameterType object.
<code>to_dict(self[, discard_cosmetic_attributes])</code>	Convert this ParameterType object to dict.

**`__init__(self, **kwargs)`**

Create a ParameterType

### Parameters

**smirks** [str] The SMIRKS match for the provided parameter type.

**allow\_cosmetic\_attributes** [bool optional. Default = False] Whether to permit non-spec kwargs (“cosmetic attributes”). If True, non-spec kwargs will be stored as an attribute of this parameter which can be accessed and written out. Otherwise an exception will be raised.

## Methods

<code>__init__(self, \*\\*kwargs)</code>	Create a ParameterType
<code>add_cosmetic_attribute(self, attr_name, ...)</code>	Add a cosmetic attribute to this ParameterType object.
<code>delete_cosmetic_attribute(self, attr_name)</code>	Delete a cosmetic attribute from this ParameterType object.
<code>to_dict(self[, discard_cosmetic_attributes])</code>	Convert this ParameterType object to dict.

## Attributes

---

**smirks**

---

**`add_cosmetic_attribute(self, attr_name, attr_value)`**

Add a cosmetic attribute to this ParameterType object. This attribute will not have a functional effect on the object in the Open Force Field toolkit, but can be written out during output.

### Parameters

**attr\_name** [str] Name of the attribute to define for this ParameterType object.

**attr\_value** [str] The value of the attribute to define for this ParameterType object.

**`delete_cosmetic_attribute(self, attr_name)`**

Delete a cosmetic attribute from this ParameterType object.

### Parameters

**attr\_name** [str] Name of the attribute to delete.

**to\_dict(self, discard\_cosmetic\_attributes=False)**

Convert this ParameterType object to dict. A unit-bearing attribute ('X') will be converted to two dict entries, one ([‘X’] containing the unitless value, and another ([‘X\_unit’]) containing a string representation of its unit.

**Parameters**

**discard\_cosmetic\_attributes** [bool, optional. Default = False] Whether to discard non-spec attributes of this ParameterType

**Returns**

**smirnoff\_dict** [dict] The SMIRNOFF-compliant dict representation of this ParameterType object.

**output\_units** [dict[str: simtk.unit.Unit]] A mapping from each simtk.unit.Quantity-valued ParameterType attribute to the unit it was converted to during serialization.

**Parameter Handlers**

Each ForceField primarily consists of several ParameterHandler objects, which each contain the machinery to add one energy component to a system. During system creation, each ParameterHandler registered to a ForceField has its `assign_parameters()` function called..

<code>ParameterList</code>	Parameter list that also supports accessing items by SMARTS string.
<code>ParameterHandler</code>	Base class for parameter handlers.
<code>BondHandler</code>	Handle SMIRNOFF <Bonds> tags
<code>AngleHandler</code>	Handle SMIRNOFF <AngleForce> tags
<code>ProperTorsionHandler</code>	Handle SMIRNOFF <ProperTorsionForce> tags
<code>ImproperTorsionHandler</code>	Handle SMIRNOFF <ImproperTorsionForce> tags
<code>vdWHandler</code>	Handle SMIRNOFF <vdW> tags
<code>ElectrostaticsHandler</code>	Handles SMIRNOFF <Electrostatics> tags.
<code>ToolkitAM1BCCHandler</code>	Handle SMIRNOFF <ToolkitAM1BCC> tags

**openforcefield.typing.engines.smirnoff.parameters.ParameterList**

**class** openforcefield.typing.engines.smirnoff.parameters.`ParameterList`(`input_parameter_list=None`)  
Parameter list that also supports accessing items by SMARTS string.

**Warning:** This API is experimental and subject to change.

**Methods**

<code>append(self, parameter)</code>	Add a ParameterType object to the end of the ParameterList
<code>clear(self, /)</code>	Remove all items from list.
<code>copy(self, /)</code>	Return a shallow copy of the list.
<code>count(self, value, /)</code>	Return number of occurrences of value.

Continued on next page

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<code>extend(self, other)</code>	Add a ParameterList object to the end of the ParameterList
<code>index(self, item)</code>	Get the numerical index of a ParameterType object or SMIRKS in this ParameterList.
<code>insert(self, index, parameter)</code>	Add a ParameterType object as if this were a list
<code>pop(self[, index])</code>	Remove and return item at index (default last).
<code>remove(self, value, /)</code>	Remove first occurrence of value.
<code>reverse(self, /)</code>	Reverse <i>IN PLACE</i> .
<code>sort(self, /, \*[key, reverse])</code>	Stable sort <i>IN PLACE</i> .
<code>to_list(self[, discard_cosmetic_attributes])</code>	Render this ParameterList to a normal list, serializing each ParameterType object in it to dict.

**`__init__(self, input_parameter_list=None)`**

Initialize a new ParameterList, optionally providing a list of ParameterType objects to initially populate it.

**Parameters**

**input\_parameter\_list: list[ParameterType], default=None** A pre-existing list of ParameterType-based objects. If None, this ParameterList will be initialized empty.

**Methods**

<code>__init__(self[, input_parameter_list])</code>	Initialize a new ParameterList, optionally providing a list of ParameterType objects to initially populate it.
<code>append(self, parameter)</code>	Add a ParameterType object to the end of the ParameterList
<code>clear(self, /)</code>	Remove all items from list.
<code>copy(self, /)</code>	Return a shallow copy of the list.
<code>count(self, value, /)</code>	Return number of occurrences of value.
<code>extend(self, other)</code>	Add a ParameterList object to the end of the ParameterList
<code>index(self, item)</code>	Get the numerical index of a ParameterType object or SMIRKS in this ParameterList.
<code>insert(self, index, parameter)</code>	Add a ParameterType object as if this were a list
<code>pop(self[, index])</code>	Remove and return item at index (default last).
<code>remove(self, value, /)</code>	Remove first occurrence of value.
<code>reverse(self, /)</code>	Reverse <i>IN PLACE</i> .
<code>sort(self, /, \*[key, reverse])</code>	Stable sort <i>IN PLACE</i> .
<code>to_list(self[, discard_cosmetic_attributes])</code>	Render this ParameterList to a normal list, serializing each ParameterType object in it to dict.

**`append(self, parameter)`**

Add a ParameterType object to the end of the ParameterList

**Parameters**

**parameter** [a ParameterType object]

**`extend(self, other)`**

Add a ParameterList object to the end of the ParameterList

**Parameters****other** [a ParameterList]**index**(*self*, *item*)

Get the numerical index of a ParameterType object or SMIRKS in this ParameterList. Raises ValueError if the item is not found.

**Parameters****item** [ParameterType object or str] The parameter or SMIRKS to look up in this ParameterList**Returns****index** [int] The index of the found item**insert**(*self*, *index*, *parameter*)

Add a ParameterType object as if this were a list

**Parameters****index** [int] The numerical position to insert the parameter at**parameter** [a ParameterType object] The parameter to insert**to\_list**(*self*, *discard\_cosmetic\_attributes=True*)

Render this ParameterList to a normal list, serializing each ParameterType object in it to dict.

**Parameters****discard\_cosmetic\_attributes** [bool, optional. Default = True] Whether to discard non-spec attributes of each ParameterType object.**Returns****parameter\_list** [List[dict]] A serialized representation of a ParameterList, with each ParameterType it contains converted to dict.**clear**(*self*, /)

Remove all items from list.

**copy**(*self*, /)

Return a shallow copy of the list.

**count**(*self*, *value*, /)

Return number of occurrences of value.

**pop**(*self*, *index=-1*, /)

Remove and return item at index (default last).

Raises IndexError if list is empty or index is out of range.

**remove**(*self*, *value*, /)

Remove first occurrence of value.

Raises ValueError if the value is not present.

**reverse**(*self*, /)Reverse *IN PLACE*.**sort**(*self*, /, \*, *key=None*, *reverse=False*)Stable sort *IN PLACE*.

## openforcefield.typing.engines.smirnoff.parameters.ParameterHandler

```
class openforcefield.typing.engines.smirnoff.parameters.ParameterHandler(allow_cosmetic_attributes=False,  
                                         skip_version_check=False,  
                                         **kwargs)
```

Base class for parameter handlers.

Parameter handlers are configured with some global parameters for a given section. They may also contain a `ParameterList` populated with `ParameterType` objects if they are responsible for assigning SMIRKS-based parameters.

**Warning:** This API is experimental and subject to change.

### Attributes

`known_kwargs` List of kwargs that can be parsed by the function.

`parameters` The ParameterList that holds this ParameterHandler's parameter objects

### Methods

<code>add_cosmetic_attribute(self, attr_name, ...)</code>	Add a cosmetic attribute to this ParameterHandler object.
<code>add_parameter(self, parameter_kwargs)</code>	Add a parameter to the forcefield, ensuring all parameters are valid.
<code>assign_parameters(self, topology, system)</code>	Assign parameters for the given Topology to the specified System object.
<code>check_handler_compatibility(self, handler_kwargs)</code>	Checks if a set of kwargs used to create a ParameterHandler are compatible with this ParameterHandler.
<code>check_parameter_compatibility(self, ...)</code>	Check to make sure that the fields requiring defined units are compatible with the required units for the Parameters handled by this ParameterHandler
<code>delete_cosmetic_attribute(self, attr_name)</code>	Delete a cosmetic attribute from this ParameterHandler object.
<code>find_matches(self, entity)</code>	Find the elements of the topology/molecule matched by a parameter type.
<code>get_parameter(self, parameter_attrs)</code>	Return the parameters in this ParameterHandler that match the parameter_attrs argument
<code>postprocess_system(self, topology, system, ...)</code>	Allow the force to perform a final post-processing pass on the System following parameter assignment, if needed.
<code>to_dict(self[, discard_cosmetic_attributes])</code>	Convert this ParameterHandler to an OrderedDict, compliant with the SMIRNOFF data spec.

`__init__(self, allow_cosmetic_attributes=False, skip_version_check=False, **kwargs)`

Initialize a ParameterHandler, optionally with a list of parameters and other kwargs.

#### Parameters

`allow_cosmetic_attributes` [bool, optional. Default = False] Whether to permit non-spec kwargs. If True, non-spec kwargs will be stored as attributes of this ob-

ject and can be accessed and modified. Otherwise an exception will be raised if a non-spec kwarg is encountered.

**skip\_version\_check: bool, optional. Default = False** If False, the SMIRNOFF section version will not be checked, and the ParameterHandler will be initialized with version set to \_MAX\_SUPPORTED\_SECTION\_VERSION.

**\*\*kwargs** [dict] The dict representation of the SMIRNOFF data source

## Methods

<code>__init__(self[, allow_cosmetic_attributes, ...])</code>	Initialize a ParameterHandler, optionally with a list of parameters and other kwargs.
<code>add_cosmetic_attribute(self, attr_name, ...)</code>	Add a cosmetic attribute to this ParameterHandler object.
<code>add_parameter(self, parameter_kwargs)</code>	Add a parameter to the forcefield, ensuring all parameters are valid.
<code>assign_parameters(self, topology, system)</code>	Assign parameters for the given Topology to the specified System object.
<code>check_handler_compatibility(self, handler_kwargs)</code>	Checks if a set of kwargs used to create a ParameterHandler are compatible with this ParameterHandler.
<code>check_parameter_compatibility(self, ...)</code>	Check to make sure that the fields requiring defined units are compatible with the required units for the Parameters handled by this ParameterHandler
<code>delete_cosmetic_attribute(self, attr_name)</code>	Delete a cosmetic attribute from this ParameterHandler object.
<code>find_matches(self, entity)</code>	Find the elements of the topology/molecule matched by a parameter type.
<code>get_parameter(self, parameterAttrs)</code>	Return the parameters in this ParameterHandler that match the parameterAttrs argument
<code>postprocess_system(self, topology, system, ...)</code>	Allow the force to perform a final post-processing pass on the System following parameter assignment, if needed.
<code>to_dict(self[, discard_cosmetic_attributes])</code>	Convert this ParameterHandler to an OrderedDict, compliant with the SMIRNOFF data spec.

## Attributes

<code>known_kwargs</code>	List of kwargs that can be parsed by the function.
<code>parameters</code>	The ParameterList that holds this ParameterHandler's parameter objects

### `property parameters`

The ParameterList that holds this ParameterHandler's parameter objects

### `add_cosmetic_attribute(self, attr_name, attr_value)`

Add a cosmetic attribute to this ParameterHandler object. This attribute will not have a functional effect on the object in the Open Force Field toolkit, but can be written out during output.

## Parameters

**attr\_name** [str] Name of the attribute to define for this ParameterType object.

**attr\_value** [str] The value of the attribute to define for this ParameterType object.

**delete\_cosmetic\_attribute(self, attr\_name)**

Delete a cosmetic attribute from this ParameterHandler object.

#### Parameters

**attr\_name** [str] Name of the attribute to delete.

**property known\_kwargs**

List of kwargs that can be parsed by the function.

**check\_parameter\_compatibility(self, parameter\_kwargs)**

Check to make sure that the fields requiring defined units are compatible with the required units for the Parameters handled by this ParameterHandler

#### Parameters

**parameter\_kwargs: dict** The dict that will be used to construct the ParameterType

#### Raises

**Raises a ValueError if the parameters are incompatible.**

**check\_handler\_compatibility(self, handler\_kwargs)**

Checks if a set of kwargs used to create a ParameterHandler are compatible with this ParameterHandler. This is called if a second handler is attempted to be initialized for the same tag.

#### Parameters

**handler\_kwargs** [dict] The kwargs that would be used to construct

#### Raises

**IncompatibleParameterError if handler\_kwargs are incompatible with existing parameters.**

**add\_parameter(self, parameter\_kwargs)**

Add a parameter to the forcefield, ensuring all parameters are valid.

#### Parameters

**parameter\_kwargs** [dict] The kwargs to pass to the ParameterHandler.INFOTYPE  
(a ParameterType) constructor

**get\_parameter(self, parameter\_attrs)**

Return the parameters in this ParameterHandler that match the parameter\_attrs argument

#### Parameters

**parameter\_attrs** [dict of {attr: value}] The attrs mapped to desired values (for example {"smirks": "[1]~[#16:2]=,:[#6:3]~[:4]", "id": "t105"} )

#### Returns

**list of ParameterType objects** A list of matching ParameterType objects

**find\_matches(self, entity)**

Find the elements of the topology/molecule matched by a parameter type.

#### Parameters

**entity** [openforcefield.topology.Topology] Topology to search.

#### Returns

**matches** [ValenceDict[Tuple[int], ParameterHandler.Match]]  
 matches[particle\_indices] is the ParameterType object matching the tuple of particle indices in entity.

### `assign_parameters(self, topology, system)`

Assign parameters for the given Topology to the specified System object.

#### Parameters

**topology** [openforcefield.topology.Topology] The Topology for which parameters are to be assigned. Either a new Force will be created or parameters will be appended to an existing Force.

**system** [simtk.openmm.System] The OpenMM System object to add the Force (or append new parameters) to.

### `postprocess_system(self, topology, system, **kwargs)`

Allow the force to perform a final post-processing pass on the System following parameter assignment, if needed.

#### Parameters

**topology** [openforcefield.topology.Topology] The Topology for which parameters are to be assigned. Either a new Force will be created or parameters will be appended to an existing Force.

**system** [simtk.openmm.System] The OpenMM System object to add the Force (or append new parameters) to.

### `to_dict(self, discard_cosmetic_attributes=False)`

Convert this ParameterHandler to an OrderedDict, compliant with the SMIRNOFF data spec.

#### Parameters

**discard\_cosmetic\_attributes** [bool, optional. Default = False.] Whether to discard non-spec parameter and header attributes in this ParameterHandler.

#### Returns

**smirnoff\_data** [OrderedDict] SMIRNOFF-spec compliant representation of this ParameterHandler and its internal ParameterList.

## openforcefield.typing.engines.smirnoff.parameters.BondHandler

### `class openforcefield.typing.engines.smirnoff.parameters.BondHandler(**kwargs)`

Handle SMIRNOFF <Bonds> tags

**Warning:** This API is experimental and subject to change.

#### Attributes

**known\_kwargs** List of kwargs that can be parsed by the function.

**parameters** The ParameterList that holds this ParameterHandler's parameter objects

#### Methods

<code>BondType(**kwargs)</code>	A SMIRNOFF bond type
<code>add_cosmetic_attribute(self, attr_name, ...)</code>	Add a cosmetic attribute to this ParameterHandler object.
<code>add_parameter(self, parameter_kwarg)</code>	Add a parameter to the forcefield, ensuring all parameters are valid.
<code>assign_parameters(self, topology, system)</code>	Assign parameters for the given Topology to the specified System object.
<code>check_handler_compatibility(self, other_handler)</code>	Checks whether this ParameterHandler encodes compatible physics as another ParameterHandler.
<code>check_parameter_compatibility(self, ...)</code>	Check to make sure that the fields requiring defined units are compatible with the required units for the Parameters handled by this ParameterHandler
<code>delete_cosmetic_attribute(self, attr_name)</code>	Delete a cosmetic attribute from this ParameterHandler object.
<code>find_matches(self, entity)</code>	Find the elements of the topology/molecule matched by a parameter type.
<code>get_parameter(self, parameter_attrs)</code>	Return the parameters in this ParameterHandler that match the parameter_attrs argument
<code>postprocess_system(self, topology, system, ...)</code>	Allow the force to perform a final post-processing pass on the System following parameter assignment, if needed.
<code>to_dict(self[, discard_cosmetic_attributes])</code>	Convert this ParameterHandler to an OrderedDict, compliant with the SMIRNOFF data spec.

---

**create\_force**  

**`__init__(self, **kwargs)`**

Initialize a ParameterHandler, optionally with a list of parameters and other kwargs.

**Parameters**

**allow\_cosmetic\_attributes** [bool, optional. Default = False] Whether to permit non-spec kwargs. If True, non-spec kwargs will be stored as attributes of this object and can be accessed and modified. Otherwise an exception will be raised if a non-spec kwarg is encountered.

**skip\_version\_check: bool, optional. Default = False** If False, the SMIRNOFF section version will not be checked, and the ParameterHandler will be initialized with version set to \_MAX\_SUPPORTED\_SECTION\_VERSION.

**\*\*kwargs** [dict] The dict representation of the SMIRNOFF data source

**Methods**

<code>__init__(self, \*\*kwargs)</code>	Initialize a ParameterHandler, optionally with a list of parameters and other kwargs.
<code>add_cosmetic_attribute(self, attr_name, ...)</code>	Add a cosmetic attribute to this ParameterHandler object.

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<code>add_parameter(self, parameter_kwargs)</code>	Add a parameter to the forcefield, ensuring all parameters are valid.
<code>assign_parameters(self, topology, system)</code>	Assign parameters for the given Topology to the specified System object.
<code>check_handler_compatibility(self, other_handler)</code>	Checks whether this ParameterHandler encodes compatible physics as another ParameterHandler.
<code>check_parameter_compatibility(self, ...)</code>	Check to make sure that the fields requiring defined units are compatible with the required units for the Parameters handled by this ParameterHandler
<code>create_force(self, system, topology, \*\*kwargs)</code>	
<code>delete_cosmetic_attribute(self, attr_name)</code>	Delete a cosmetic attribute from this ParameterHandler object.
<code>find_matches(self, entity)</code>	Find the elements of the topology/molecule matched by a parameter type.
<code>get_parameter(self, parameter_attrs)</code>	Return the parameters in this ParameterHandler that match the parameter_attrs argument
<code>postprocess_system(self, topology, system, ...)</code>	Allow the force to perform a final post-processing pass on the System following parameter assignment, if needed.
<code>to_dict(self[, discard_cosmetic_attributes])</code>	Convert this ParameterHandler to an OrderedDict, compliant with the SMIRNOFF data spec.

## Attributes

<code>known_kwargs</code>	List of kwargs that can be parsed by the function.
<code>parameters</code>	The ParameterList that holds this ParameterHandler's parameter objects

```
class BondType(**kwargs)
A SMIRNOFF bond type
```

**Warning:** This API is experimental and subject to change.

## Attributes

`smirks`

## Methods

<code>add_cosmetic_attribute(self, attr_name, ...)</code>	Add a cosmetic attribute to this ParameterType object.
<code>delete_cosmetic_attribute(self, attr_name)</code>	Delete a cosmetic attribute from this ParameterType object.
<code>to_dict(self[, discard_cosmetic_attributes])</code>	Convert this ParameterType object to dict.

**add\_cosmetic\_attribute(self, attr\_name, attr\_value)**

Add a cosmetic attribute to this ParameterType object. This attribute will not have a functional effect on the object in the Open Force Field toolkit, but can be written out during output.

**Parameters****attr\_name** [str] Name of the attribute to define for this ParameterType object.**attr\_value** [str] The value of the attribute to define for this ParameterType object.**delete\_cosmetic\_attribute(self, attr\_name)**

Delete a cosmetic attribute from this ParameterType object.

**Parameters****attr\_name** [str] Name of the attribute to delete.**to\_dict(self, discard\_cosmetic\_attributes=False)**

Convert this ParameterType object to dict. A unit-bearing attribute ('X') will be converted to two dict entries, one ('[X]') containing the unitless value, and another ('[X\_unit]') containing a string representation of its unit.

**Parameters****discard\_cosmetic\_attributes** [bool, optional. Default = False] Whether to discard non-spec attributes of this ParameterType**Returns****smirnoff\_dict** [dict] The SMIRNOFF-compliant dict representation of this ParameterType object.**output\_units** [dict[str: simtk.unit.Unit]] A mapping from each simtk.unit.Quantity-valued ParameterType attribute to the unit it was converted to during serialization.**check\_handler\_compatibility(self, other\_handler)**

Checks whether this ParameterHandler encodes compatible physics as another ParameterHandler. This is called if a second handler is attempted to be initialized for the same tag.

**Parameters****other\_handler** [a ParameterHandler object] The handler to compare to.**Raises****IncompatibleParameterError** if **handler\_kwargs** are incompatible with existing parameters.**add\_cosmetic\_attribute(self, attr\_name, attr\_value)**

Add a cosmetic attribute to this ParameterHandler object. This attribute will not have a functional effect on the object in the Open Force Field toolkit, but can be written out during output.

**Parameters****attr\_name** [str] Name of the attribute to define for this ParameterType object.**attr\_value** [str] The value of the attribute to define for this ParameterType object.**add\_parameter(self, parameter\_kwargs)**

Add a parameter to the forcefield, ensuring all parameters are valid.

**Parameters****parameter\_kwargs** [dict] The kwargs to pass to the ParameterHandler.INFOTYPE (a ParameterType) constructor**assign\_parameters(self, topology, system)**

Assign parameters for the given Topology to the specified System object.

**Parameters**

**topology** [openforcefield.topology.Topology] The Topology for which parameters are to be assigned. Either a new Force will be created or parameters will be appended to an existing Force.

**system** [simtk.openmm.System] The OpenMM System object to add the Force (or append new parameters) to.

#### **check\_parameter\_compatibility(self, parameter\_kwargs)**

Check to make sure that the fields requiring defined units are compatible with the required units for the Parameters handled by this ParameterHandler

##### Parameters

**parameter\_kwargs:** dict The dict that will be used to construct the ParameterType

##### Raises

**Raises a ValueError if the parameters are incompatible.**

#### **delete\_cosmetic\_attribute(self, attr\_name)**

Delete a cosmetic attribute from this ParameterHandler object.

##### Parameters

**attr\_name** [str] Name of the attribute to delete.

#### **find\_matches(self, entity)**

Find the elements of the topology/molecule matched by a parameter type.

##### Parameters

**entity** [openforcefield.topology.Topology] Topology to search.

##### Returns

**matches** [ValenceDict[Tuple[int], ParameterHandler.Match]]  
matches[particle\_indices] is the ParameterType object matching the tuple of particle indices in entity.

#### **get\_parameter(self, parameter\_attrs)**

Return the parameters in this ParameterHandler that match the parameter\_attrs argument

##### Parameters

**parameter\_attrs** [dict of {attr: value}] The attrs mapped to desired values (for example {"smirks": "[\*:1]~[#16:2]=,:[#6:3]~[:4]", "id": "t105"} )

##### Returns

**list of ParameterType objects** A list of matching ParameterType objects

#### **property known\_kwargs**

List of kwargs that can be parsed by the function.

#### **property parameters**

The ParameterList that holds this ParameterHandler's parameter objects

#### **postprocess\_system(self, topology, system, \*\*kwargs)**

Allow the force to perform a final post-processing pass on the System following parameter assignment, if needed.

##### Parameters

**topology** [openforcefield.topology.Topology] The Topology for which parameters are to be assigned. Either a new Force will be created or parameters will be appended to an existing Force.

**system** [simtk.openmm.System] The OpenMM System object to add the Force (or append new parameters) to.

**to\_dict(self, discard\_cosmetic\_attributes=False)**

Convert this ParameterHandler to an OrderedDict, compliant with the SMIRNOFF data spec.

#### Parameters

**discard\_cosmetic\_attributes** [bool, optional. Default = False.] Whether to discard non-spec parameter and header attributes in this ParameterHandler.

#### Returns

**smirnoff\_data** [OrderedDict] SMIRNOFF-spec compliant representation of this ParameterHandler and its internal ParameterList.

## openforcefield.typing.engines.smirnoff.parameters.AngleHandler

**class** openforcefield.typing.engines.smirnoff.parameters.AngleHandler(\*\*kwargs)

Handle SMIRNOFF <AngleForce> tags

**Warning:** This API is experimental and subject to change.

#### Attributes

**known\_kwargs** List of kwargs that can be parsed by the function.

**parameters** The ParameterList that holds this ParameterHandler's parameter objects

#### Methods

<code>AngleType(**kwargs)</code>	A SMIRNOFF angle type.
<code>add_cosmetic_attribute(self, attr_name, ...)</code>	Add a cosmetic attribute to this ParameterHandler object.
<code>add_parameter(self, parameter_kwarg)</code>	Add a parameter to the forcefield, ensuring all parameters are valid.
<code>assign_parameters(self, topology, system)</code>	Assign parameters for the given Topology to the specified System object.
<code>check_handler_compatibility(self, other_handler)</code>	Checks whether this ParameterHandler encodes compatible physics as another ParameterHandler.
<code>check_parameter_compatibility(self, ...)</code>	Check to make sure that the fields requiring defined units are compatible with the required units for the Parameters handled by this ParameterHandler
<code>delete_cosmetic_attribute(self, attr_name)</code>	Delete a cosmetic attribute from this ParameterHandler object.
<code>find_matches(self, entity)</code>	Find the elements of the topology/molecule matched by a parameter type.
<code>get_parameter(self, parameter_attrs)</code>	Return the parameters in this ParameterHandler that match the parameter_attrs argument

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<code>postprocess_system(self, topology, system, ...)</code>	Allow the force to perform a final post-processing pass on the System following parameter assignment, if needed.
<code>to_dict(self[, discard_cosmetic_attributes])</code>	Convert this ParameterHandler to an OrderedDict, compliant with the SMIRNOFF data spec.

`create_force``__init__(self, **kwargs)`

Initialize a ParameterHandler, optionally with a list of parameters and other kwargs.

**Parameters**

**allow\_cosmetic\_attributes** [bool, optional. Default = False] Whether to permit non-spec kwargs. If True, non-spec kwargs will be stored as attributes of this object and can be accessed and modified. Otherwise an exception will be raised if a non-spec kwarg is encountered.

**skip\_version\_check: bool, optional. Default = False** If False, the SMIRNOFF section version will not be checked, and the ParameterHandler will be initialized with version set to \_MAX\_SUPPORTED\_SECTION\_VERSION.

**\*\*kwargs** [dict] The dict representation of the SMIRNOFF data source

**Methods**

<code>__init__(self, \*\*kwargs)</code>	Initialize a ParameterHandler, optionally with a list of parameters and other kwargs.
<code>add_cosmetic_attribute(self, attr_name, ...)</code>	Add a cosmetic attribute to this ParameterHandler object.
<code>add_parameter(self, parameter_kwargs)</code>	Add a parameter to the forcefield, ensuring all parameters are valid.
<code>assign_parameters(self, topology, system)</code>	Assign parameters for the given Topology to the specified System object.
<code>check_handler_compatibility(self, other_handler)</code>	Checks whether this ParameterHandler encodes compatible physics as another ParameterHandler.
<code>check_parameter_compatibility(self, ...)</code>	Check to make sure that the fields requiring defined units are compatible with the required units for the Parameters handled by this ParameterHandler
<code>create_force(self, system, topology, \*\*kwargs)</code>	
<code>delete_cosmetic_attribute(self, attr_name)</code>	Delete a cosmetic attribute from this ParameterHandler object.
<code>find_matches(self, entity)</code>	Find the elements of the topology/molecule matched by a parameter type.
<code>get_parameter(self, parameter_attrs)</code>	Return the parameters in this ParameterHandler that match the parameter_attrs argument

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Table 63 – continued from previous page

<code>postprocess_system(self, topology, system, ...)</code>	Allow the force to perform a final post-processing pass on the System following parameter assignment, if needed.
<code>to_dict(self[, discard_cosmetic_attributes])</code>	Convert this ParameterHandler to an OrderedDict, compliant with the SMIRNOFF data spec.

### Attributes

<code>known_kwargs</code>	List of kwargs that can be parsed by the function.
<code>parameters</code>	The ParameterList that holds this ParameterHandler's parameter objects

`class AngleType(**kwargs)`  
A SMIRNOFF angle type.

**Warning:** This API is experimental and subject to change.

### Attributes

`smirks`

### Methods

<code>add_cosmetic_attribute(self, attr_name, attr_value)</code>	Add a cosmetic attribute to this ParameterType object.
<code>delete_cosmetic_attribute(self, attr_name)</code>	Delete a cosmetic attribute from this ParameterType object.
<code>to_dict(self[, discard_cosmetic_attributes])</code>	Convert this ParameterType object to dict.

**add\_cosmetic\_attribute(self, attr\_name, attr\_value)**

Add a cosmetic attribute to this ParameterType object. This attribute will not have a functional effect on the object in the Open Force Field toolkit, but can be written out during output.

#### Parameters

`attr_name` [str] Name of the attribute to define for this ParameterType object.  
`attr_value` [str] The value of the attribute to define for this ParameterType object.

**delete\_cosmetic\_attribute(self, attr\_name)**

Delete a cosmetic attribute from this ParameterType object.

#### Parameters

`attr_name` [str] Name of the attribute to delete.

**to\_dict(self, discard\_cosmetic\_attributes=False)**

Convert this ParameterType object to dict. A unit-bearing attribute ('X') will be converted to two dict entries, one ('X') containing the unitless value, and another ('X\_unit') containing a string representation of its unit.

#### Parameters

`discard_cosmetic_attributes` [bool, optional. Default = False] Whether to discard non-spec attributes of this ParameterType

#### Returns

`smirnoff_dict` [dict] The SMIRNOFF-compliant dict representation of this ParameterType.

terType object.  
**output\_units** [dict[str: simtk.unit.Unit]] A mapping from each simtk.unit.Quantity-valued ParameterType attribute to the unit it was converted to during serialization.

**check\_handler\_compatibility(self, other\_handler)**

Checks whether this ParameterHandler encodes compatible physics as another ParameterHandler. This is called if a second handler is attempted to be initialized for the same tag.

**Parameters**

**other\_handler** [a ParameterHandler object] The handler to compare to.

**Raises**

**IncompatibleParameterError** if **handler\_kwarg**s are incompatible with existing parameters.

**add\_cosmetic\_attribute(self, attr\_name, attr\_value)**

Add a cosmetic attribute to this ParameterHandler object. This attribute will not have a functional effect on the object in the Open Force Field toolkit, but can be written out during output.

**Parameters**

**attr\_name** [str] Name of the attribute to define for this ParameterType object.

**attr\_value** [str] The value of the attribute to define for this ParameterType object.

**add\_parameter(self, parameter\_kwarg)**

Add a parameter to the forcefield, ensuring all parameters are valid.

**Parameters**

**parameter\_kwarg** [dict] The kwarg to pass to the ParameterHandler.INFOTYPE (a ParameterType) constructor

**assign\_parameters(self, topology, system)**

Assign parameters for the given Topology to the specified System object.

**Parameters**

**topology** [openforcefield.topology.Topology] The Topology for which parameters are to be assigned. Either a new Force will be created or parameters will be appended to an existing Force.

**system** [simtk.openmm.System] The OpenMM System object to add the Force (or append new parameters) to.

**check\_parameter\_compatibility(self, parameter\_kwarg)**

Check to make sure that the fields requiring defined units are compatible with the required units for the Parameters handled by this ParameterHandler

**Parameters**

**parameter\_kwarg: dict** The dict that will be used to construct the ParameterType

**Raises**

**Raises a ValueError if the parameters are incompatible.**

**delete\_cosmetic\_attribute(self, attr\_name)**

Delete a cosmetic attribute from this ParameterHandler object.

**Parameters**

**attr\_name** [str] Name of the attribute to delete.

**find\_matches(self, entity)**

Find the elements of the topology/molecule matched by a parameter type.

**Parameters**

**entity** [openforcefield.topology.Topology] Topology to search.

**Returns**

**matches** [ValenceDict[Tuple[int], ParameterHandler.Match]]

matches[particle\_indices] is the ParameterType object matching the tuple of particle indices in entity.

**get\_parameter(self, parameter\_attrs)**

Return the parameters in this ParameterHandler that match the parameter\_attrs argument

**Parameters**

**parameter\_attrs** [dict of {attr: value}] The attrs mapped to desired values (for example {"smirks": "[1]~[#16:2]=:[#6:3]~[:4]", "id": "t105"} )

**Returns**

**list of ParameterType objects** A list of matching ParameterType objects

**property known\_kwargs**

List of kwargs that can be parsed by the function.

**property parameters**

The ParameterList that holds this ParameterHandler's parameter objects

**postprocess\_system(self, topology, system, \*\*kwargs)**

Allow the force to perform a final post-processing pass on the System following parameter assignment, if needed.

**Parameters**

**topology** [openforcefield.topology.Topology] The Topology for which parameters are to be assigned. Either a new Force will be created or parameters will be appended to an existing Force.

**system** [simtk.openmm.System] The OpenMM System object to add the Force (or append new parameters) to.

**to\_dict(self, discard\_cosmetic\_attributes=False)**

Convert this ParameterHandler to an OrderedDict, compliant with the SMIRNOFF data spec.

**Parameters**

**discard\_cosmetic\_attributes** [bool, optional. Default = False.] Whether to discard non-spec parameter and header attributes in this ParameterHandler.

**Returns**

**smirnoff\_data** [OrderedDict] SMIRNOFF-spec compliant representation of this ParameterHandler and its internal ParameterList.

**openforcefield.typing.engines.smirnoff.parameters.ProperTorsionHandler****class openforcefield.typing.engines.smirnoff.parameters.ProperTorsionHandler(\*\*kwargs)**

Handle SMIRNOFF <ProperTorsionForce> tags

**Warning:** This API is experimental and subject to change.

## Attributes

`known_kwarg` List of kwargs that can be parsed by the function.

`parameters` The ParameterList that holds this ParameterHandler's parameter objects

## Methods

<code>ProperTorsionType(**kwargs)</code>	A SMIRNOFF torsion type for proper torsions.
<code>add_cosmetic_attribute(self, attr_name, ...)</code>	Add a cosmetic attribute to this ParameterHandler object.
<code>add_parameter(self, parameter_kwarg)</code>	Add a parameter to the forcefield, ensuring all parameters are valid.
<code>assign_parameters(self, topology, system)</code>	Assign parameters for the given Topology to the specified System object.
<code>check_handler_compatibility(self, other_handler)</code>	Checks whether this ParameterHandler encodes compatible physics as another ParameterHandler.
<code>check_parameter_compatibility(self, ...)</code>	Check to make sure that the fields requiring defined units are compatible with the required units for the Parameters handled by this ParameterHandler
<code>delete_cosmetic_attribute(self, attr_name)</code>	Delete a cosmetic attribute from this ParameterHandler object.
<code>find_matches(self, entity)</code>	Find the elements of the topology/molecule matched by a parameter type.
<code>get_parameter(self, parameter_attrs)</code>	Return the parameters in this ParameterHandler that match the parameter_attrs argument
<code>postprocess_system(self, topology, system, ...)</code>	Allow the force to perform a final post-processing pass on the System following parameter assignment, if needed.
<code>to_dict(self[, discard_cosmetic_attributes])</code>	Convert this ParameterHandler to an OrderedDict, compliant with the SMIRNOFF data spec.

<code>create_force</code>	
<code>validate_parameters</code>	

`__init__(self, **kwargs)`

Initialize a ParameterHandler, optionally with a list of parameters and other kwargs.

### Parameters

`allow_cosmetic_attributes` [bool, optional. Default = False] Whether to permit non-spec kwargs. If True, non-spec kwargs will be stored as attributes of this object and can be accessed and modified. Otherwise an exception will be raised if a non-spec kwarg is encountered.

`skip_version_check: bool, optional. Default = False` If False, the SMIRNOFF section version will not be checked, and the ParameterHandler will be initialized with version set to `_MAX_SUPPORTED_SECTION_VERSION`.

**\*\*kwargs** [dict] The dict representation of the SMIRNOFF data source

## Methods

<code>__init__(self, **kwargs)</code>	Initialize a ParameterHandler, optionally with a list of parameters and other kwargs.
<code>add_cosmetic_attribute(self, attr_name, ...)</code>	Add a cosmetic attribute to this ParameterHandler object.
<code>add_parameter(self, parameter_kwargs)</code>	Add a parameter to the forcefield, ensuring all parameters are valid.
<code>assign_parameters(self, topology, system)</code>	Assign parameters for the given Topology to the specified System object.
<code>check_handler_compatibility(self, other_handler)</code>	Checks whether this ParameterHandler encodes compatible physics as another ParameterHandler.
<code>check_parameter_compatibility(self, ...)</code>	Check to make sure that the fields requiring defined units are compatible with the required units for the Parameters handled by this ParameterHandler
<code>create_force(self, system, topology, **kwargs)</code>	
<code>delete_cosmetic_attribute(self, attr_name)</code>	Delete a cosmetic attribute from this ParameterHandler object.
<code>find_matches(self, entity)</code>	Find the elements of the topology/molecule matched by a parameter type.
<code>get_parameter(self, parameter_attrs)</code>	Return the parameters in this ParameterHandler that match the parameter_attrs argument
<code>postprocess_system(self, topology, system, ...)</code>	Allow the force to perform a final post-processing pass on the System following parameter assignment, if needed.
<code>to_dict(self[, discard_cosmetic_attributes])</code>	Convert this ParameterHandler to an OrderedDict, compliant with the SMIRNOFF data spec.
<code>validate_parameters(self)</code>	

## Attributes

<code>known_kwargs</code>	List of kwargs that can be parsed by the function.
<code>parameters</code>	The ParameterList that holds this ParameterHandler's parameter objects

**class ProperTorsionType(\*\*kwargs)**  
A SMIRNOFF torsion type for proper torsions.

**Warning:** This API is experimental and subject to change.

## Attributes

`smirks`

## Methods

<code>add_cosmetic_attribute(self, attr_name, attr_value)</code>	Add a cosmetic attribute to this ParameterType object.
<code>delete_cosmetic_attribute(self, attr_name)</code>	Delete a cosmetic attribute from this ParameterType object.
<code>to_dict(self[, discard_cosmetic_attributes])</code>	Convert this ParameterType object to dict.

### `add_cosmetic_attribute(self, attr_name, attr_value)`

Add a cosmetic attribute to this ParameterType object. This attribute will not have a functional effect on the object in the Open Force Field toolkit, but can be written out during output.

#### Parameters

- `attr_name` [str] Name of the attribute to define for this ParameterType object.
- `attr_value` [str] The value of the attribute to define for this ParameterType object.

### `delete_cosmetic_attribute(self, attr_name)`

Delete a cosmetic attribute from this ParameterType object.

#### Parameters

- `attr_name` [str] Name of the attribute to delete.

### `to_dict(self, discard_cosmetic_attributes=False)`

Convert this ParameterType object to dict. A unit-bearing attribute ('X') will be converted to two dict entries, one ([‘X’] containing the unitless value, and another ([‘X\_unit’]) containing a string representation of its unit.

#### Parameters

- `discard_cosmetic_attributes` [bool, optional. Default = False] Whether to discard non-spec attributes of this ParameterType

#### Returns

- `smirnoff_dict` [dict] The SMIRNOFF-compliant dict representation of this ParameterType object.

- `output_units` [dict[str: simtk.unit.Unit]] A mapping from each simtk.unit.Quantity-valued ParameterType attribute to the unit it was converted to during serialization.

### `check_handler_compatibility(self, other_handler)`

Checks whether this ParameterHandler encodes compatible physics as another ParameterHandler. This is called if a second handler is attempted to be initialized for the same tag.

#### Parameters

- `other_handler` [a ParameterHandler object] The handler to compare to.

#### Raises

`IncompatibleParameterError` if `handler_kwarg`s are incompatible with existing parameters.

### `add_cosmetic_attribute(self, attr_name, attr_value)`

Add a cosmetic attribute to this ParameterHandler object. This attribute will not have a functional effect on the object in the Open Force Field toolkit, but can be written out during output.

#### Parameters

- `attr_name` [str] Name of the attribute to define for this ParameterType object.

- `attr_value` [str] The value of the attribute to define for this ParameterType object.

### `add_parameter(self, parameter_kwarg)`

Add a parameter to the forcefield, ensuring all parameters are valid.

**Parameters**

**parameter\_kwarg** [dict] The kwargs to pass to the ParameterHandler.INFOTYPE (a ParameterType) constructor

**assign\_parameters(self, topology, system)**

Assign parameters for the given Topology to the specified System object.

**Parameters**

**topology** [openforcefield.topology.Topology] The Topology for which parameters are to be assigned. Either a new Force will be created or parameters will be appended to an existing Force.

**system** [simtk.openmm.System] The OpenMM System object to add the Force (or append new parameters) to.

**check\_parameter\_compatibility(self, parameter\_kwarg)**

Check to make sure that the fields requiring defined units are compatible with the required units for the Parameters handled by this ParameterHandler

**Parameters**

**parameter\_kwarg**: dict The dict that will be used to construct the ParameterType

**Raises**

Raises a ValueError if the parameters are incompatible.

**delete\_cosmetic\_attribute(self, attr\_name)**

Delete a cosmetic attribute from this ParameterHandler object.

**Parameters**

**attr\_name** [str] Name of the attribute to delete.

**find\_matches(self, entity)**

Find the elements of the topology/molecule matched by a parameter type.

**Parameters**

**entity** [openforcefield.topology.Topology] Topology to search.

**Returns**

**matches** [ValenceDict[Tuple[int], ParameterHandler.\_Match]]  
matches[particle\_indices] is the ParameterType object matching the tuple of particle indices in entity.

**get\_parameter(self, parameter\_attrs)**

Return the parameters in this ParameterHandler that match the parameter\_attrs argument

**Parameters**

**parameter\_attrs** [dict of {attr: value}] The attrs mapped to desired values (for example {"smirks": "[\*:1]~[#16:2]=,:[#6:3]~[:4]", "id": "t105"} )

**Returns**

**list of ParameterType objects** A list of matching ParameterType objects

**property known\_kwarg**

List of kwargs that can be parsed by the function.

**property parameters**

The ParameterList that holds this ParameterHandler's parameter objects

**postprocess\_system**(*self, topology, system, \*\*kwargs*)

Allow the force to perform a final post-processing pass on the System following parameter assignment, if needed.

#### Parameters

**topology** [openforcefield.topology.Topology] The Topology for which parameters are to be assigned. Either a new Force will be created or parameters will be appended to an existing Force.

**system** [simtk.openmm.System] The OpenMM System object to add the Force (or append new parameters) to.

**to\_dict**(*self, discard\_cosmetic\_attributes=False*)

Convert this ParameterHandler to an OrderedDict, compliant with the SMIRNOFF data spec.

#### Parameters

**discard\_cosmetic\_attributes** [bool, optional. Default = False.] Whether to discard non-spec parameter and header attributes in this ParameterHandler.

#### Returns

**smirnoff\_data** [OrderedDict] SMIRNOFF-spec compliant representation of this ParameterHandler and its internal ParameterList.

## openforcefield.typing.engines.smirnoff.parameters.ImproperTorsionHandler

**class** openforcefield.typing.engines.smirnoff.parameters.ImproperTorsionHandler(\*\*kwargs)

Handle SMIRNOFF <ImproperTorsionForce> tags

**Warning:** This API is experimental and subject to change.

#### Attributes

**known\_kwargs** List of kwargs that can be parsed by the function.

**parameters** The ParameterList that holds this ParameterHandler's parameter objects

#### Methods

<code>ImproperTorsionType(**kwargs)</code>	A SMIRNOFF torsion type for improper torsions.
<code>add_cosmetic_attribute(self, attr_name, ...)</code>	Add a cosmetic attribute to this ParameterHandler object.
<code>add_parameter(self, parameter_kwargs)</code>	Add a parameter to the forcefield, ensuring all parameters are valid.
<code>assign_parameters(self, topology, system)</code>	Assign parameters for the given Topology to the specified System object.
<code>check_handler_compatibility(self, other_handler)</code>	Checks whether this ParameterHandler encodes compatible physics as another ParameterHandler.

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<code>check_parameter_compatibility(self, ...)</code>	Check to make sure that the fields requiring defined units are compatible with the required units for the Parameters handled by this ParameterHandler
<code>delete_cosmetic_attribute(self, attr_name)</code>	Delete a cosmetic attribute from this ParameterHandler object.
<code>find_matches(self, entity)</code>	Find the improper torsions in the topology/molecule matched by a parameter type.
<code>get_parameter(self, parameter_attrs)</code>	Return the parameters in this ParameterHandler that match the parameter_attrs argument
<code>postprocess_system(self, topology, system, ...)</code>	Allow the force to perform a final post-processing pass on the System following parameter assignment, if needed.
<code>to_dict(self[, discard_cosmetic_attributes])</code>	Convert this ParameterHandler to an OrderedDict, compliant with the SMIRNOFF data spec.

<code>create_force</code>	
<code>validate_parameters</code>	

`__init__(self, **kwargs)`

Initialize a ParameterHandler, optionally with a list of parameters and other kwargs.

**Parameters**

**allow\_cosmetic\_attributes** [bool, optional. Default = False] Whether to permit non-spec kwargs. If True, non-spec kwargs will be stored as attributes of this object and can be accessed and modified. Otherwise an exception will be raised if a non-spec kwarg is encountered.

**skip\_version\_check: bool, optional. Default = False** If False, the SMIRNOFF section version will not be checked, and the ParameterHandler will be initialized with version set to `_MAX_SUPPORTED_SECTION_VERSION`.

**\*\*kwargs** [dict] The dict representation of the SMIRNOFF data source

**Methods**

<code>__init__(self, \*\*kwargs)</code>	Initialize a ParameterHandler, optionally with a list of parameters and other kwargs.
<code>add_cosmetic_attribute(self, attr_name, ...)</code>	Add a cosmetic attribute to this ParameterHandler object.
<code>add_parameter(self, parameter_kwarg)</code>	Add a parameter to the forcefield, ensuring all parameters are valid.
<code>assign_parameters(self, topology, system)</code>	Assign parameters for the given Topology to the specified System object.
<code>check_handler_compatibility(self, other_handler)</code>	Checks whether this ParameterHandler encodes compatible physics as another ParameterHandler.

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Table 71 – continued from previous page

<code>check_parameter_compatibility(self, ...)</code>	Check to make sure that the fields requiring defined units are compatible with the required units for the Parameters handled by this ParameterHandler	
<code>create_force(self, system, topology, \*\*kwargs)</code>	system,	topology,
<code>delete_cosmetic_attribute(self, attr_name)</code>	Delete a cosmetic attribute from this ParameterHandler object.	
<code>find_matches(self, entity)</code>	Find the improper torsions in the topology/molecule matched by a parameter type.	
<code>get_parameter(self, parameter_attrs)</code>	Return the parameters in this ParameterHandler that match the parameter_attrs argument	
<code>postprocess_system(self, topology, system, ...)</code>	Allow the force to perform a final post-processing pass on the System following parameter assignment, if needed.	
<code>to_dict(self[, discard_cosmetic_attributes])</code>	Convert this ParameterHandler to an OrderedDict, compliant with the SMIRNOFF data spec.	
<code>validate_parameters(self)</code>		

### Attributes

<code>known_kwargs</code>	List of kwargs that can be parsed by the function.
<code>parameters</code>	The ParameterList that holds this ParameterHandler's parameter objects

`class ImproperTorsionType(**kwargs)`  
A SMIRNOFF torsion type for improper torsions.

**Warning:** This API is experimental and subject to change.

### Attributes

`smirks`

### Methods

<code>add_cosmetic_attribute(self, attr_name, attr_value)</code>	Add a cosmetic attribute to this ParameterType object.
<code>delete_cosmetic_attribute(self, attr_name)</code>	Delete a cosmetic attribute from this ParameterType object.
<code>to_dict(self[, discard_cosmetic_attributes])</code>	Convert this ParameterType object to dict.

`add_cosmetic_attribute(self, attr_name, attr_value)`

Add a cosmetic attribute to this ParameterType object. This attribute will not have a functional effect on the object in the Open Force Field toolkit, but can be written out during output.

#### Parameters

`attr_name` [str] Name of the attribute to define for this ParameterType object.

`attr_value` [str] The value of the attribute to define for this ParameterType object.

**delete\_cosmetic\_attribute(self, attr\_name)**

Delete a cosmetic attribute from this ParameterType object.

**Parameters**

**attr\_name** [str] Name of the attribute to delete.

**to\_dict(self, discard\_cosmetic\_attributes=False)**

Convert this ParameterType object to dict. A unit-bearing attribute ('X') will be converted to two dict entries, one ('[X]') containing the unitless value, and another ('[X\_unit]') containing a string representation of its unit.

**Parameters**

**discard\_cosmetic\_attributes** [bool, optional. Default = False] Whether to discard non-spec attributes of this ParameterType

**Returns**

**smirnoff\_dict** [dict] The SMIRNOFF-compliant dict representation of this ParameterType object.

**output\_units** [dict[str: simtk.unit.Unit]] A mapping from each simtk.unit.Quantity-valued ParameterType attribute to the unit it was converted to during serialization.

**check\_handler\_compatibility(self, other\_handler)**

Checks whether this ParameterHandler encodes compatible physics as another ParameterHandler. This is called if a second handler is attempted to be initialized for the same tag.

**Parameters**

**other\_handler** [a ParameterHandler object] The handler to compare to.

**Raises**

**IncompatibleParameterError** if **handler\_kwarg**s are incompatible with existing parameters.

**find\_matches(self, entity)**

Find the improper torsions in the topology/molecule matched by a parameter type.

**Parameters**

**entity** [openforcefield.topology.Topology] Topology to search.

**Returns**

**matches** [ImproperDict[Tuple[int], ParameterHandler.Match]]  
matches[atom\_indices] is the ParameterType object matching the 4-tuple of atom indices in entity.

**add\_cosmetic\_attribute(self, attr\_name, attr\_value)**

Add a cosmetic attribute to this ParameterHandler object. This attribute will not have a functional effect on the object in the Open Force Field toolkit, but can be written out during output.

**Parameters**

**attr\_name** [str] Name of the attribute to define for this ParameterType object.

**attr\_value** [str] The value of the attribute to define for this ParameterType object.

**add\_parameter(self, parameter\_kwarg)**

Add a parameter to the forcefield, ensuring all parameters are valid.

**Parameters**

**parameter\_kwarg** [dict] The kwarg to pass to the ParameterHandler.INFOTYPE (a ParameterType) constructor

**assign\_parameters**(*self, topology, system*)

Assign parameters for the given Topology to the specified System object.

**Parameters**

**topology** [openforcefield.topology.Topology] The Topology for which parameters are to be assigned. Either a new Force will be created or parameters will be appended to an existing Force.

**system** [simtk.openmm.System] The OpenMM System object to add the Force (or append new parameters) to.

**check\_parameter\_compatibility**(*self, parameter\_kwargs*)

Check to make sure that the fields requiring defined units are compatible with the required units for the Parameters handled by this ParameterHandler

**Parameters**

**parameter\_kwargs**: dict The dict that will be used to construct the ParameterType

**Raises**

**Raises a ValueError if the parameters are incompatible.**

**delete\_cosmetic\_attribute**(*self, attr\_name*)

Delete a cosmetic attribute from this ParameterHandler object.

**Parameters**

**attr\_name** [str] Name of the attribute to delete.

**get\_parameter**(*self, parameterAttrs*)

Return the parameters in this ParameterHandler that match the parameterAttrs argument

**Parameters**

**parameterAttrs** [dict of {attr: value}] The attrs mapped to desired values (for example {"smirks": "[1]~[#16:2]=,[#6:3]~[:4]", "id": "t105"} )

**Returns**

**list of ParameterType objects** A list of matching ParameterType objects

**property known\_kwargs**

List of kwargs that can be parsed by the function.

**property parameters**

The ParameterList that holds this ParameterHandler's parameter objects

**postprocess\_system**(*self, topology, system, \*\*kwargs*)

Allow the force to perform a final post-processing pass on the System following parameter assignment, if needed.

**Parameters**

**topology** [openforcefield.topology.Topology] The Topology for which parameters are to be assigned. Either a new Force will be created or parameters will be appended to an existing Force.

**system** [simtk.openmm.System] The OpenMM System object to add the Force (or append new parameters) to.

**to\_dict**(*self, discard\_cosmetic\_attributes=False*)

Convert this ParameterHandler to an OrderedDict, compliant with the SMIRNOFF data spec.

**Parameters**

**discard\_cosmetic\_attributes** [bool, optional. Default = False.] Whether to discard non-spec parameter and header attributes in this ParameterHandler.

#### Returns

**smirnoff\_data** [OrderedDict] SMIRNOFF-spec compliant representation of this ParameterHandler and its internal ParameterList.

## openforcefield.typing.engines.smirnoff.parameters.vdWHandler

**class** openforcefield.typing.engines.smirnoff.parameters.**vdWHandler**(\*\*kwargs)  
Handle SMIRNOFF <vdW> tags

**Warning:** This API is experimental and subject to change.

#### Attributes

**combining\_rules** The combining\_rules used to model van der Waals interactions  
**cutoff** The cutoff used for long-range van der Waals interactions  
**known\_kwargs** List of kwargs that can be parsed by the function.  
**method** The method used to handle long-range van der Waals interactions  
**parameters** The ParameterList that holds this ParameterHandler's parameter objects  
**potential** The potential used to model van der Waals interactions  
**switch\_width** The switching width used for long-range van der Waals interactions

#### Methods

<code>add_cosmetic_attribute(self, attr_name, ...)</code>	Add a cosmetic attribute to this ParameterHandler object.
<code>add_parameter(self, parameter_kwargs)</code>	Add a parameter to the forcefield, ensuring all parameters are valid.
<code>assign_parameters(self, topology, system)</code>	Assign parameters for the given Topology to the specified System object.
<code>check_handler_compatibility(self, other_handler)</code>	Checks whether this ParameterHandler encodes compatible physics as another ParameterHandler.
<code>check_parameter_compatibility(self, ...)</code>	Check to make sure that the fields requiring defined units are compatible with the required units for the Parameters handled by this ParameterHandler
<code>delete_cosmetic_attribute(self, attr_name)</code>	Delete a cosmetic attribute from this ParameterHandler object.
<code>find_matches(self, entity)</code>	Find the elements of the topology/molecule matched by a parameter type.
<code>get_parameter(self, parameter_attrs)</code>	Return the parameters in this ParameterHandler that match the parameterAttrs argument

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Table 74 – continued from previous page

<code>postprocess_system(self, system, topology, ...)</code>	Allow the force to perform a final post-processing pass on the System following parameter assignment, if needed.
<code>to_dict(self[, discard_cosmetic_attributes])</code>	Convert this ParameterHandler to an OrderedDict, compliant with the SMIRNOFF data spec.
<code>vdWType(**kwargs)</code>	A SMIRNOFF vdWForce type.

`create_force``__init__(self, **kwargs)`

Initialize a ParameterHandler, optionally with a list of parameters and other kwargs.

**Parameters**

**allow\_cosmetic\_attributes** [bool, optional. Default = False] Whether to permit non-spec kwargs. If True, non-spec kwargs will be stored as attributes of this object and can be accessed and modified. Otherwise an exception will be raised if a non-spec kwarg is encountered.

**skip\_version\_check: bool, optional. Default = False** If False, the SMIRNOFF section version will not be checked, and the ParameterHandler will be initialized with version set to \_MAX\_SUPPORTED\_SECTION\_VERSION.

**\*\*kwargs** [dict] The dict representation of the SMIRNOFF data source

**Methods**

<code>__init__(self, \*\*kwargs)</code>	Initialize a ParameterHandler, optionally with a list of parameters and other kwargs.
<code>add_cosmetic_attribute(self, attr_name, ...)</code>	Add a cosmetic attribute to this ParameterHandler object.
<code>add_parameter(self, parameter_kwargs)</code>	Add a parameter to the forcefield, ensuring all parameters are valid.
<code>assign_parameters(self, topology, system)</code>	Assign parameters for the given Topology to the specified System object.
<code>check_handler_compatibility(self, other_handler)</code>	Checks whether this ParameterHandler encodes compatible physics as another ParameterHandler.
<code>check_parameter_compatibility(self, ...)</code>	Check to make sure that the fields requiring defined units are compatible with the required units for the Parameters handled by this ParameterHandler
<code>create_force(self, system, topology, \*\*kwargs)</code>	
<code>delete_cosmetic_attribute(self, attr_name)</code>	Delete a cosmetic attribute from this ParameterHandler object.
<code>find_matches(self, entity)</code>	Find the elements of the topology/molecule matched by a parameter type.
<code>get_parameter(self, parameter_attrs)</code>	Return the parameters in this ParameterHandler that match the parameter_attrs argument

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Table 75 – continued from previous page

<code>postprocess_system(self, system, topology, ...)</code>	Allow the force to perform a final post-processing pass on the System following parameter assignment, if needed.
<code>to_dict(self[, discard_cosmetic_attributes])</code>	Convert this ParameterHandler to an OrderedDict, compliant with the SMIRNOFF data spec.

### Attributes

<code>combining_rules</code>	The combining_rules used to model van der Waals interactions
<code>cutoff</code>	The cutoff used for long-range van der Waals interactions
<code>known_kwarg</code> s	List of kwarg that can be parsed by the function.
<code>method</code>	The method used to handle long-range van der Waals interactions
<code>parameters</code>	The ParameterList that holds this ParameterHandler's parameter objects
<code>potential</code>	The potential used to model van der Waals interactions
<code>switch_width</code>	The switching width used for long-range van der Waals interactions

```
class vdWType(**kwargs)
```

A SMIRNOFF vdWForce type.

**Warning:** This API is experimental and subject to change.

### Attributes

`smirks`

### Methods

<code>add_cosmetic_attribute(self, attr_name, ...)</code>	Add a cosmetic attribute to this ParameterType object.
<code>delete_cosmetic_attribute(self, attr_name)</code>	Delete a cosmetic attribute from this ParameterType object.
<code>to_dict(self[, discard_cosmetic_attributes])</code>	Convert this ParameterType object to dict.

```
add_cosmetic_attribute(self, attr_name, attr_value)
```

Add a cosmetic attribute to this ParameterType object. This attribute will not have a functional effect on the object in the Open Force Field toolkit, but can be written out during output.

#### Parameters

`attr_name` [str] Name of the attribute to define for this ParameterType object.

`attr_value` [str] The value of the attribute to define for this ParameterType object.

```
delete_cosmetic_attribute(self, attr_name)
```

Delete a cosmetic attribute from this ParameterType object.

#### Parameters

**attr\_name** [str] Name of the attribute to delete.

**to\_dict(self, discard\_cosmetic\_attributes=False)**

Convert this ParameterType object to dict. A unit-bearing attribute ('X') will be converted to two dict entries, one ([‘X’] containing the unitless value, and another ([‘X\_unit’]) containing a string representation of its unit.

**Parameters**

**discard\_cosmetic\_attributes** [bool, optional. Default = False] Whether to discard non-spec attributes of this ParameterType

**Returns**

**smirnoff\_dict** [dict] The SMIRNOFF-compliant dict representation of this ParameterType object.

**output\_units** [dict[str: simtk.unit.Unit]] A mapping from each simtk.unit.Quantity-valued ParameterType attribute to the unit it was converted to during serialization.

**property potential**

The potential used to model van der Waals interactions

**property combining\_rules**

The combining\_rules used to model van der Waals interactions

**property method**

The method used to handle long-range van der Waals interactions

**property cutoff**

The cutoff used for long-range van der Waals interactions

**property switch\_width**

The switching width used for long-range van der Waals interactions

**check\_handler\_compatibility(self, other\_handler)**

Checks whether this ParameterHandler encodes compatible physics as another ParameterHandler. This is called if a second handler is attempted to be initialized for the same tag.

**Parameters**

**other\_handler** [a ParameterHandler object] The handler to compare to.

**Raises**

**IncompatibleParameterError** if handler\_kwarg are incompatible with existing parameters.

**postprocess\_system(self, system, topology, \*\*kwargs)**

Allow the force to perform a final post-processing pass on the System following parameter assignment, if needed.

**Parameters**

**topology** [openforcefield.topology.Topology] The Topology for which parameters are to be assigned. Either a new Force will be created or parameters will be appended to an existing Force.

**system** [simtk.openmm.System] The OpenMM System object to add the Force (or append new parameters) to.

**add\_cosmetic\_attribute(self, attr\_name, attr\_value)**

Add a cosmetic attribute to this ParameterHandler object. This attribute will not have a functional effect on the object in the Open Force Field toolkit, but can be written out during output.

**Parameters**

**attr\_name** [str] Name of the attribute to define for this ParameterType object.

**attr\_value** [str] The value of the attribute to define for this ParameterType object.

**add\_parameter(self, parameter\_kwarg)**

Add a parameter to the forcefield, ensuring all parameters are valid.

#### Parameters

**parameter\_kwarg** [dict] The kwarg to pass to the ParameterHandler.INFOTYPE (a ParameterType) constructor

**assign\_parameters(self, topology, system)**

Assign parameters for the given Topology to the specified System object.

#### Parameters

**topology** [openforcefield.topology.Topology] The Topology for which parameters are to be assigned. Either a new Force will be created or parameters will be appended to an existing Force.

**system** [simtk.openmm.System] The OpenMM System object to add the Force (or append new parameters) to.

**check\_parameter\_compatibility(self, parameter\_kwarg)**

Check to make sure that the fields requiring defined units are compatible with the required units for the Parameters handled by this ParameterHandler

#### Parameters

**parameter\_kwarg: dict** The dict that will be used to construct the ParameterType

#### Raises

**Raises a ValueError if the parameters are incompatible.**

**delete\_cosmetic\_attribute(self, attr\_name)**

Delete a cosmetic attribute from this ParameterHandler object.

#### Parameters

**attr\_name** [str] Name of the attribute to delete.

**find\_matches(self, entity)**

Find the elements of the topology/molecule matched by a parameter type.

#### Parameters

**entity** [openforcefield.topology.Topology] Topology to search.

#### Returns

**matches** [ValenceDict[Tuple[int], ParameterHandler.Match]]  
matches[particle\_indices] is the ParameterType object matching the tuple of particle indices in entity.

**get\_parameter(self, parameter\_attrs)**

Return the parameters in this ParameterHandler that match the parameter\_attrs argument

#### Parameters

**parameter\_attrs** [dict of {attr: value}] The attrs mapped to desired values (for example {"smirks": "[1]~[#16:2]=,[#6:3]~[:4]", "id": "t105"} )

#### Returns

**list of ParameterType objects** A list of matching ParameterType objects

**property known\_kwargs**

List of kwargs that can be parsed by the function.

**property parameters**

The ParameterList that holds this ParameterHandler's parameter objects

**to\_dict(self, discard\_cosmetic\_attributes=False)**

Convert this ParameterHandler to an OrderedDict, compliant with the SMIRNOFF data spec.

**Parameters**

**discard\_cosmetic\_attributes** [bool, optional. Default = False.] Whether to discard non-spec parameter and header attributes in this ParameterHandler.

**Returns**

**smirnoff\_data** [OrderedDict] SMIRNOFF-spec compliant representation of this ParameterHandler and its internal ParameterList.

**openforcefield.typing.engines.smirnoff.parameters.ElectrostaticsHandler**

**class** openforcefield.typing.engines.smirnoff.parameters.ElectrostaticsHandler(\*\*kwargs)  
Handles SMIRNOFF <Electrostatics> tags.

**Warning:** This API is experimental and subject to change.

**Attributes**

**cutoff** The cutoff used for long-range van der Waals interactions

**known\_kwargs** List of kwargs that can be parsed by the function.

**method** The method used to model long-range electrostatic interactions

**parameters** The ParameterList that holds this ParameterHandler's parameter objects

**switch\_width** The switching width used for long-range electrostatics interactions

**Methods**

<code>add_cosmetic_attribute(self, attr_name, ...)</code>	Add a cosmetic attribute to this ParameterHandler object.
<code>add_parameter(self, parameter_kwargs)</code>	Add a parameter to the forcefield, ensuring all parameters are valid.
<code>assign_parameters(self, topology, system)</code>	Assign parameters for the given Topology to the specified System object.
<code>check_handler_compatibility(self, other_handler)</code>	Checks whether this ParameterHandler encodes compatible physics as another ParameterHandler.
<code>check_parameter_compatibility(self, ...)</code>	Check to make sure that the fields requiring defined units are compatible with the required units for the Parameters handled by this ParameterHandler

Continued on next page

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<code>delete_cosmetic_attribute(self, attr_name)</code>	Delete a cosmetic attribute from this ParameterHandler object.
<code>find_matches(self, entity)</code>	Find the elements of the topology/molecule matched by a parameter type.
<code>get_parameter(self, parameter_attrs)</code>	Return the parameters in this ParameterHandler that match the parameter_attrs argument
<code>postprocess_system(self, topology, system, ...)</code>	Allow the force to perform a final post-processing pass on the System following parameter assignment, if needed.
<code>to_dict(self[, discard_cosmetic_attributes])</code>	Convert this ParameterHandler to an OrderedDict, compliant with the SMIRNOFF data spec.

`create_force``__init__(self, **kwargs)`

Initialize a ParameterHandler, optionally with a list of parameters and other kwargs.

**Parameters**

**allow\_cosmetic\_attributes** [bool, optional. Default = False] Whether to permit non-spec kwargs. If True, non-spec kwargs will be stored as attributes of this object and can be accessed and modified. Otherwise an exception will be raised if a non-spec kwarg is encountered.

**skip\_version\_check: bool, optional. Default = False** If False, the SMIRNOFF section version will not be checked, and the ParameterHandler will be initialized with version set to \_MAX\_SUPPORTED\_SECTION\_VERSION.

**\*\*kwargs** [dict] The dict representation of the SMIRNOFF data source

**Methods**

<code>__init__(self, \*\*kwargs)</code>	Initialize a ParameterHandler, optionally with a list of parameters and other kwargs.
<code>add_cosmetic_attribute(self, attr_name, ...)</code>	Add a cosmetic attribute to this ParameterHandler object.
<code>add_parameter(self, parameter_kwargs)</code>	Add a parameter to the forcefield, ensuring all parameters are valid.
<code>assign_parameters(self, topology, system)</code>	Assign parameters for the given Topology to the specified System object.
<code>check_handler_compatibility(self, other_handler)</code>	Checks whether this ParameterHandler encodes compatible physics as another ParameterHandler.
<code>check_parameter_compatibility(self, ...)</code>	Check to make sure that the fields requiring defined units are compatible with the required units for the Parameters handled by this ParameterHandler
<code>create_force(self, system, topology, \*\*kwargs)</code>	
<code>delete_cosmetic_attribute(self, attr_name)</code>	Delete a cosmetic attribute from this ParameterHandler object.

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Table 79 – continued from previous page

<code>find_matches(self, entity)</code>	Find the elements of the topology/molecule matched by a parameter type.
<code>get_parameter(self, parameter_attrs)</code>	Return the parameters in this ParameterHandler that match the <code>parameter_attrs</code> argument
<code>postprocess_system(self, topology, system, ...)</code>	Allow the force to perform a final post-processing pass on the System following parameter assignment, if needed.
<code>to_dict(self[, discard_cosmetic_attributes])</code>	Convert this ParameterHandler to an OrderedDict, compliant with the SMIRNOFF data spec.

## Attributes

<code>cutoff</code>	The cutoff used for long-range van der Waals interactions
<code>known_kwargs</code>	List of kwargs that can be parsed by the function.
<code>method</code>	The method used to model long-range electrostatic interactions
<code>parameters</code>	The ParameterList that holds this ParameterHandler's parameter objects
<code>switch_width</code>	The switching width used for long-range electrostatics interactions

### `property method`

The method used to model long-range electrostatic interactions

### `property cutoff`

The cutoff used for long-range van der Waals interactions

### `property switch_width`

The switching width used for long-range electrostatics interactions

### `check_handler_compatibility(self, other_handler)`

Checks whether this ParameterHandler encodes compatible physics as another ParameterHandler. This is called if a second handler is attempted to be initialized for the same tag.

#### Parameters

`other_handler` [a ParameterHandler object] The handler to compare to.

#### Raises

`IncompatibleParameterError` if `handler_kwargs` are incompatible with existing parameters.

### `add_cosmetic_attribute(self, attr_name, attr_value)`

Add a cosmetic attribute to this ParameterHandler object. This attribute will not have a functional effect on the object in the Open Force Field toolkit, but can be written out during output.

#### Parameters

`attr_name` [str] Name of the attribute to define for this ParameterType object.

`attr_value` [str] The value of the attribute to define for this ParameterType object.

### `add_parameter(self, parameter_kwargs)`

Add a parameter to the forcefield, ensuring all parameters are valid.

#### Parameters

**parameter\_kwarg** [dict] The kwargs to pass to the ParameterHandler.INFOTYPE (a ParameterType) constructor

**assign\_parameters**(*self*, *topology*, *system*)

Assign parameters for the given Topology to the specified System object.

#### Parameters

**topology** [openforcefield.topology.Topology] The Topology for which parameters are to be assigned. Either a new Force will be created or parameters will be appended to an existing Force.

**system** [simtk.openmm.System] The OpenMM System object to add the Force (or append new parameters) to.

**check\_parameter\_compatibility**(*self*, *parameter\_kwarg*)

Check to make sure that the fields requiring defined units are compatible with the required units for the Parameters handled by this ParameterHandler

#### Parameters

**parameter\_kwarg**: dict The dict that will be used to construct the ParameterType

#### Raises

**Raises a ValueError if the parameters are incompatible.**

**delete\_cosmetic\_attribute**(*self*, *attr\_name*)

Delete a cosmetic attribute from this ParameterHandler object.

#### Parameters

**attr\_name** [str] Name of the attribute to delete.

**find\_matches**(*self*, *entity*)

Find the elements of the topology/molecule matched by a parameter type.

#### Parameters

**entity** [openforcefield.topology.Topology] Topology to search.

#### Returns

**matches** [ValenceDict[Tuple[int], ParameterHandler.\_Match]]  
matches[particle\_indices] is the ParameterType object matching the tuple of particle indices in entity.

**get\_parameter**(*self*, *parameter\_attrs*)

Return the parameters in this ParameterHandler that match the parameter\_attrs argument

#### Parameters

**parameter\_attrs** [dict of {attr: value}] The attrs mapped to desired values (for example {"smirks": "[1]~[#16:2]=,[#6:3]~[:4]", "id": "t105"} )

#### Returns

**list of ParameterType objects** A list of matching ParameterType objects

**property known\_kwarg**

List of kwargs that can be parsed by the function.

**property parameters**

The ParameterList that holds this ParameterHandler's parameter objects

**postprocess\_system**(*self, topology, system, \*\*kwargs*)

Allow the force to perform a final post-processing pass on the System following parameter assignment, if needed.

#### Parameters

**topology** [openforcefield.topology.Topology] The Topology for which parameters are to be assigned. Either a new Force will be created or parameters will be appended to an existing Force.

**system** [simtk.openmm.System] The OpenMM System object to add the Force (or append new parameters) to.

**to\_dict**(*self, discard\_cosmetic\_attributes=False*)

Convert this ParameterHandler to an OrderedDict, compliant with the SMIRNOFF data spec.

#### Parameters

**discard\_cosmetic\_attributes** [bool, optional. Default = False.] Whether to discard non-spec parameter and header attributes in this ParameterHandler.

#### Returns

**smirnoff\_data** [OrderedDict] SMIRNOFF-spec compliant representation of this ParameterHandler and its internal ParameterList.

## openforcefield.typing.engines.smirnoff.parameters.ToolkitAM1BCCHandler

**class** openforcefield.typing.engines.smirnoff.parameters.**ToolkitAM1BCCHandler**(\*\*kwargs)

Handle SMIRNOFF <ToolkitAM1BCC> tags

**Warning:** This API is experimental and subject to change.

#### Attributes

**known\_kwargs** List of kwargs that can be parsed by the function.

**parameters** The ParameterList that holds this ParameterHandler's parameter objects

#### Methods

<code>add_cosmetic_attribute(self, attr_name, ...)</code>	Add a cosmetic attribute to this ParameterHandler object.
<code>add_parameter(self, parameter_kwargs)</code>	Add a parameter to the forcefield, ensuring all parameters are valid.
<code>assign_charge_from_molecules(self, molecule, ...)</code>	Given an input molecule, checks against a list of molecules for an isomorphic match.
<code>assign_parameters(self, topology, system)</code>	Assign parameters for the given Topology to the specified System object.
<code>check_handler_compatibility(self, other_handler)</code>	Checks whether this ParameterHandler encodes compatible physics as another ParameterHandler.

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<code>check_parameter_compatibility(self, ...)</code>	Check to make sure that the fields requiring defined units are compatible with the required units for the Parameters handled by this ParameterHandler
<code>delete_cosmetic_attribute(self, attr_name)</code>	Delete a cosmetic attribute from this ParameterHandler object.
<code>find_matches(self, entity)</code>	Find the elements of the topology/molecule matched by a parameter type.
<code>get_parameter(self, parameter_attrs)</code>	Return the parameters in this ParameterHandler that match the parameter_attrs argument
<code>postprocess_system(self, system, topology, ...)</code>	Allow the force to perform a final post-processing pass on the System following parameter assignment, if needed.
<code>to_dict(self[, discard_cosmetic_attributes])</code>	Convert this ParameterHandler to an OrderedDict, compliant with the SMIRNOFF data spec.

`create_force``__init__(self, **kwargs)`

Initialize a ParameterHandler, optionally with a list of parameters and other kwargs.

**Parameters**

`allow_cosmetic_attributes` [bool, optional. Default = False] Whether to permit non-spec kwargs. If True, non-spec kwargs will be stored as attributes of this object and can be accessed and modified. Otherwise an exception will be raised if a non-spec kwarg is encountered.

`skip_version_check: bool, optional. Default = False` If False, the SMIRNOFF section version will not be checked, and the ParameterHandler will be initialized with version set to `_MAX_SUPPORTED_SECTION_VERSION`.

`**kwargs` [dict] The dict representation of the SMIRNOFF data source

**Methods**

<code>__init__(self, \*\*kwargs)</code>	Initialize a ParameterHandler, optionally with a list of parameters and other kwargs.
<code>add_cosmetic_attribute(self, attr_name, ...)</code>	Add a cosmetic attribute to this ParameterHandler object.
<code>add_parameter(self, parameter_kwarg)</code>	Add a parameter to the forcefield, ensuring all parameters are valid.
<code>assign_charge_from_molecules(self, molecule, ...)</code>	Given an input molecule, checks against a list of molecules for an isomorphic match.
<code>assign_parameters(self, topology, system)</code>	Assign parameters for the given Topology to the specified System object.
<code>check_handler_compatibility(self, other_handler)</code>	Checks whether this ParameterHandler encodes compatible physics as another ParameterHandler.

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Table 82 – continued from previous page

<code>check_parameter_compatibility(self, ...)</code>	Check to make sure that the fields requiring defined units are compatible with the required units for the Parameters handled by this ParameterHandler	
<code>create_force(self, system, topology, \*\*kwargs)</code>		
<code>delete_cosmetic_attribute(self, attr_name)</code>	Delete a cosmetic attribute from this ParameterHandler object.	
<code>find_matches(self, entity)</code>	Find the elements of the topology/molecule matched by a parameter type.	
<code>get_parameter(self, parameter_attrs)</code>	Return the parameters in this ParameterHandler that match the <code>parameter_attrs</code> argument	
<code>postprocess_system(self, system, topology, ...)</code>	Allow the force to perform a final post-processing pass on the System following parameter assignment, if needed.	
<code>to_dict(self[, discard_cosmetic_attributes])</code>	Convert this ParameterHandler to an OrderedDict, compliant with the SMIRNOFF data spec.	

## Attributes

<code>known_kwargs</code>	List of kwargs that can be parsed by the function.
<code>parameters</code>	The ParameterList that holds this ParameterHandler's parameter objects

`check_handler_compatibility(self, other_handler, assume_missing_is_default=True)`

Checks whether this ParameterHandler encodes compatible physics as another ParameterHandler. This is called if a second handler is attempted to be initialized for the same tag.

### Parameters

`other_handler` [a ParameterHandler object] The handler to compare to.

### Raises

`IncompatibleParameterError` if `handler_kwargs` are incompatible with existing parameters.

`assign_charge_from_molecules(self, molecule, charge_mols)`

Given an input molecule, checks against a list of molecules for an isomorphic match. If found, assigns partial charges from the match to the input molecule.

### Parameters

`molecule` [an openforcefield.topology.FrozenMolecule] The molecule to have partial charges assigned if a match is found.

`charge_mols` [list of [openforcefield.topology.FrozenMolecule]] A list of molecules with charges already assigned.

### Returns

`match_found` [bool] Whether a match was found. If True, the input molecule will have been modified in-place.

`postprocess_system(self, system, topology, **kwargs)`

Allow the force to perform a final post-processing pass on the System following parameter assignment, if needed.

**Parameters**

**topology** [openforcefield.topology.Topology] The Topology for which parameters are to be assigned. Either a new Force will be created or parameters will be appended to an existing Force.

**system** [simtk.openmm.System] The OpenMM System object to add the Force (or append new parameters) to.

**add\_cosmetic\_attribute(self, attr\_name, attr\_value)**

Add a cosmetic attribute to this ParameterHandler object. This attribute will not have a functional effect on the object in the Open Force Field toolkit, but can be written out during output.

**Parameters**

**attr\_name** [str] Name of the attribute to define for this ParameterType object.

**attr\_value** [str] The value of the attribute to define for this ParameterType object.

**add\_parameter(self, parameter\_kwargs)**

Add a parameter to the forcefield, ensuring all parameters are valid.

**Parameters**

**parameter\_kwargs** [dict] The kwargs to pass to the ParameterHandler.INFOTYPE (a ParameterType) constructor

**assign\_parameters(self, topology, system)**

Assign parameters for the given Topology to the specified System object.

**Parameters**

**topology** [openforcefield.topology.Topology] The Topology for which parameters are to be assigned. Either a new Force will be created or parameters will be appended to an existing Force.

**system** [simtk.openmm.System] The OpenMM System object to add the Force (or append new parameters) to.

**check\_parameter\_compatibility(self, parameter\_kwargs)**

Check to make sure that the fields requiring defined units are compatible with the required units for the Parameters handled by this ParameterHandler

**Parameters**

**parameter\_kwargs: dict** The dict that will be used to construct the ParameterType

**Raises**

**Raises a ValueError if the parameters are incompatible.**

**delete\_cosmetic\_attribute(self, attr\_name)**

Delete a cosmetic attribute from this ParameterHandler object.

**Parameters**

**attr\_name** [str] Name of the attribute to delete.

**find\_matches(self, entity)**

Find the elements of the topology/molecule matched by a parameter type.

**Parameters**

**entity** [openforcefield.topology.Topology] Topology to search.

**Returns**

---

**matches** [ValenceDict[Tuple[int], ParameterHandler.Match]]  
 matches[particle\_indices] is the ParameterType object matching the tuple  
 of particle indices in entity.

**get\_parameter(self, parameter\_attrs)**

Return the parameters in this ParameterHandler that match the parameter\_attrs argument

**Parameters**

**parameter\_attrs** [dict of {attr: value}] The attrs mapped to desired values (for example {"smirks": "[1]~[#16:2]=,:[#6:3]~[:4]", "id": "t105"} )

**Returns**

**list of ParameterType objects** A list of matching ParameterType objects

**property known\_kwargs**

List of kwargs that can be parsed by the function.

**property parameters**

The ParameterList that holds this ParameterHandler's parameter objects

**to\_dict(self, discard\_cosmetic\_attributes=False)**

Convert this ParameterHandler to an OrderedDict, compliant with the SMIRNOFF data spec.

**Parameters**

**discard\_cosmetic\_attributes** [bool, optional. Default = False.] Whether to discard non-spec parameter and header attributes in this ParameterHandler.

**Returns**

**smirnoff\_data** [OrderedDict] SMIRNOFF-spec compliant representation of this ParameterHandler and its internal ParameterList.

## Parameter I/O Handlers

ParameterIOHandler objects handle reading and writing of serialized SMIRNOFF data sources.

---

ParameterIOHandler	Base class for handling serialization/deserialization of SMIRNOFF ForceField objects
XMLParameterIOHandler	Handles serialization/deserialization of SMIRNOFF ForceField objects from OFFXML format.

---

### openforcefield.typing.engines.smirnoff.io.ParameterIOHandler

**class openforcefield.typing.engines.smirnoff.io.ParameterIOHandler**

Base class for handling serialization/deserialization of SMIRNOFF ForceField objects

**Methods**

---

**parse\_file(self, file\_path)****Parameters**


---

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<code>parse_string(self, data)</code>	Parse a SMIRNOFF force field definition in a serialized format
<code>to_file(self, file_path, smirnoff_data)</code>	Write the current forcefield parameter set to a file.
<code>to_string(self, smirnoff_data)</code>	Render the forcefield parameter set to a string

**\_\_init\_\_(self)**

Create a new ParameterIOHandler.

**Methods**

<code>__init__(self)</code>	Create a new ParameterIOHandler.
<code>parse_file(self, file_path)</code>	<b>Parameters</b>
<code>parse_string(self, data)</code>	Parse a SMIRNOFF force field definition in a serialized format
<code>to_file(self, file_path, smirnoff_data)</code>	Write the current forcefield parameter set to a file.
<code>to_string(self, smirnoff_data)</code>	Render the forcefield parameter set to a string

**parse\_file(self, file\_path)****Parameters****file\_path****parse\_string(self, data)**

Parse a SMIRNOFF force field definition in a serialized format

**Parameters****data****to\_file(self, file\_path, smirnoff\_data)**

Write the current forcefield parameter set to a file.

**Parameters****file\_path** [str] The path to the file to write to.**smirnoff\_data** [dict] A dictionary structured in compliance with the SMIRNOFF spec**to\_string(self, smirnoff\_data)**

Render the forcefield parameter set to a string

**Parameters****smirnoff\_data** [dict] A dictionary structured in compliance with the SMIRNOFF spec**Returns****str**

`openforcefield.typing.engines.smirnoff.io.XMLParameterIOHandler``class openforcefield.typing.engines.smirnoff.io.XMLParameterIOHandler`

Handles serialization/deserialization of SMIRNOFF ForceField objects from OFFXML format.

**Methods**

<code>parse_file(self, source)</code>	Parse a SMIRNOFF force field definition in XML format, read from a file.
<code>parse_string(self, data)</code>	Parse a SMIRNOFF force field definition in XML format.
<code>to_file(self, file_path, smirnoff_data)</code>	Write the current forcefield parameter set to a file.
<code>to_string(self, smirnoff_data)</code>	Write the current forcefield parameter set to an XML string.

`__init__(self)`

Create a new ParameterIOHandler.

**Methods**

<code>__init__(self)</code>	Create a new ParameterIOHandler.
<code>parse_file(self, source)</code>	Parse a SMIRNOFF force field definition in XML format, read from a file.
<code>parse_string(self, data)</code>	Parse a SMIRNOFF force field definition in XML format.
<code>to_file(self, file_path, smirnoff_data)</code>	Write the current forcefield parameter set to a file.
<code>to_string(self, smirnoff_data)</code>	Write the current forcefield parameter set to an XML string.

`parse_file(self, source)`

Parse a SMIRNOFF force field definition in XML format, read from a file.

**Parameters**

**source** [str or io.RawIOWrapper] File path of file-like object implementing a `read()` method specifying a SMIRNOFF force field definition in [the SMIRNOFF XML format](#).

**Raises****ParseError** If the XML cannot be processed.**FileNotFoundException** If the file could not be found.`parse_string(self, data)`

Parse a SMIRNOFF force field definition in XML format.

A ParseError is raised if the XML cannot be processed.

**Parameters**

**data** [str] A SMIRNOFF force field definition in [the SMIRNOFF XML format](#).

**to\_file(self, file\_path, smirnoff\_data)**  
Write the current forcefield parameter set to a file.

**Parameters**

**file\_path** [str] The path to the file to be written. The *.offxml* or *.xml* file extension must be present.

**smirnoff\_data** [dict] A dict structured in compliance with the SMIRNOFF data spec.

**to\_string(self, smirnoff\_data)**  
Write the current forcefield parameter set to an XML string.

**Parameters**

**smirnoff\_data** [dict] A dictionary structured in compliance with the SMIRNOFF spec

**Returns**

**serialized\_forcefield** [str] XML String representation of this forcefield.

## 2.3 Utilities

### 2.3.1 Toolkit wrappers

The toolkit wrappers provide a simple uniform API for accessing minimal functionality of cheminformatics toolkits.

These toolkit wrappers are generally used through a `ToolkitRegistry`, which can be constructed with a desired precedence of toolkits:

```
from openforcefield.utils.toolkits import ToolkitRegistry
toolkit_registry = ToolkitRegistry()
toolkit_precedence = [OpenEyeToolkitWrapper, RDKitToolkitWrapper, AmberToolsToolkitWrapper]
[ toolkit_registry.register(toolkit) for toolkit in toolkit_precedence if toolkit.is_available() ]
```

Alternatively, the global toolkit registry (which will attempt to register any available toolkits) can be used:

```
from openforcefield.utils.toolkits import DEFAULT_TOOLKIT_REGISTRY as toolkit_registry
```

The toolkit wrappers can then be accessed through the registry:

```
molecule = Molecule.from_smiles('Cc1ccccc1')
smiles = toolkit_registry.call('to_smiles', molecule)
```

<code>ToolkitRegistry</code>	Registry for <code>ToolkitWrapper</code> objects
<code>ToolkitWrapper</code>	Toolkit wrapper base class.
<code>OpenEyeToolkitWrapper</code>	OpenEye toolkit wrapper
<code>RDKitToolkitWrapper</code>	RDKit toolkit wrapper
<code>AmberToolsToolkitWrapper</code>	AmberTools toolkit wrapper

**openforcefield.utils.toolkits.ToolkitRegistry**

```
class openforcefield.utils.toolkits.ToolkitRegistry(register_imported_toolkit_wrappers=False,
                                                 toolkit_precedence=None,
                                                 exception_if_unavailable=True)
```

Registry for ToolkitWrapper objects

**Examples**

Register toolkits in a specified order, skipping if unavailable

```
>>> from openforcefield.utils.toolkits import ToolkitRegistry
>>> toolkit_registry = ToolkitRegistry()
>>> toolkit_precedence = [OpenEyeToolkitWrapper, RDKitToolkitWrapper, AmberToolsToolkitWrapper]
>>> for toolkit in toolkit_precedence:
...     if toolkit.is_available():
...         toolkit_registry.register_toolkit(toolkit)
```

Register specified toolkits, raising an exception if one is unavailable

```
>>> toolkit_registry = ToolkitRegistry()
>>> toolkits = [OpenEyeToolkitWrapper, AmberToolsToolkitWrapper]
>>> for toolkit in toolkits:
...     toolkit_registry.register_toolkit(toolkit)
```

Register all available toolkits in arbitrary order

```
>>> from openforcefield.utils import all_subclasses
>>> toolkits = all_subclasses(ToolkitWrapper)
>>> for toolkit in toolkit_precedence:
...     if toolkit.is_available():
...         toolkit_registry.register_toolkit(toolkit)
```

Retrieve the global singleton toolkit registry, which is created when this module is imported from all available toolkits:

```
>>> from openforcefield.utils.toolkits import GLOBAL_TOOLKIT_REGISTRY as toolkit_registry
>>> available_toolkits = toolkit_registry.registered_toolkits
```

**Warning:** This API is experimental and subject to change.

**Attributes**

`registered_toolkits` List registered toolkits.

**Methods**

<code>add_toolkit(self, toolkit_wrapper)</code>	Append a ToolkitWrapper onto the list of toolkits in this ToolkitRegistry
<code>call(self, method_name, \*args, \*\*kwargs)</code>	Execute the requested method by attempting to use all registered toolkits in order of precedence.

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<code>register_toolkit(self, toolkit_wrapper[, ...])</code>	Register the provided toolkit wrapper class, instantiating an object of it.
<code>resolve(self, method_name)</code>	Resolve the requested method name by checking all registered toolkits in order of precedence for one that provides the requested method.

`__init__(self, register_imported_toolkit_wrappers=False, toolkit_precedence=None, exception_if_unavailable=True)`  
Create an empty toolkit registry.

#### Parameters

`register_imported_toolkit_wrappers` [bool, optional, default=False]

If True, will attempt to register all imported ToolkitWrapper subclasses that can be found, in no particular order.

`toolkit_precedence` [list, optional, default=None] List of toolkit wrapper classes, in order of desired precedence when performing molecule operations. If None, defaults to [OpenEyeToolkitWrapper, RDKitToolkitWrapper, AmberToolsToolkitWrapper].

`exception_if_unavailable` [bool, optional, default=True] If True, an exception will be raised if the toolkit is unavailable

#### Methods

<code>__init__(self[, ...])</code>	Create an empty toolkit registry.
<code>add_toolkit(self, toolkit_wrapper)</code>	Append a ToolkitWrapper onto the list of toolkits in this ToolkitRegistry
<code>call(self, method_name, *args, **kwargs)</code>	Execute the requested method by attempting to use all registered toolkits in order of precedence.
<code>register_toolkit(self, toolkit_wrapper[, ...])</code>	Register the provided toolkit wrapper class, instantiating an object of it.
<code>resolve(self, method_name)</code>	Resolve the requested method name by checking all registered toolkits in order of precedence for one that provides the requested method.

#### Attributes

<code>registered_toolkits</code>	List registered toolkits.
----------------------------------	---------------------------

#### `property registered_toolkits`

List registered toolkits.

**Warning:** This API is experimental and subject to change.

#### Returns

`toolkits` [iterable of toolkit objects]

**register\_toolkit(self, toolkit\_wrapper, exception\_if\_unavailable=True)**  
 Register the provided toolkit wrapper class, instantiating an object of it.

**Warning:** This API is experimental and subject to change.

#### Parameters

**toolkit\_wrapper** [instance or subclass of ToolkitWrapper] The toolkit wrapper to register or its class.

**exception\_if\_unavailable** [bool, optional, default=True] If True, an exception will be raised if the toolkit is unavailable

**add\_toolkit(self, toolkit\_wrapper)**

Append a ToolkitWrapper onto the list of toolkits in this ToolkitRegistry

**Warning:** This API is experimental and subject to change.

#### Parameters

**toolkit\_wrapper** [openforcefield.utils.ToolkitWrapper] The ToolkitWrapper object to add to the list of registered toolkits

**resolve(self, method\_name)**

Resolve the requested method name by checking all registered toolkits in order of precedence for one that provides the requested method.

#### Parameters

**method\_name** [str] The name of the method to resolve

#### Returns

**method** The method of the first registered toolkit that provides the requested method name

#### Raises

**NotImplementedError** if the requested method cannot be found among the registered toolkits

## Examples

Create a molecule, and call the toolkit to\_smiles() method directly

```
>>> from openforcefield.topology import Molecule
>>> molecule = Molecule.from_smiles('Cc1ccccc1')
>>> toolkit_registry = ToolkitRegistry(register_imported_toolkit_wrappers=True)
>>> method = toolkit_registry.resolve('to_smiles')
>>> smiles = method(molecule)
```

**call(self, method\_name, \*args, \*\*kwargs)**

Execute the requested method by attempting to use all registered toolkits in order of precedence.

\*args and \*\*kwargs are passed to the desired method, and return values of the method are returned

This is a convenient shorthand for `toolkit_registry.resolve_method(method_name)(*args, **kwargs)`

#### Parameters

`method_name` [str] The name of the method to execute

#### Raises

`NotImplementedError` if the requested method cannot be found among the registered toolkits

#### Examples

Create a molecule, and call the toolkit `to_smiles()` method directly

```
>>> from openforcefield.topology import Molecule
>>> molecule = Molecule.from_smiles('Cc1ccccc1')
>>> toolkit_registry = ToolkitRegistry(register_imported_toolkit_wrappers=True)
>>> smiles = toolkit_registry.call('to_smiles', molecule)
```

## openforcefield.utils.toolkits.ToolkitWrapper

```
class openforcefield.utils.toolkits.ToolkitWrapper
    Toolkit wrapper base class.
```

**Warning:** This API is experimental and subject to change.

#### Attributes

`toolkit_file_read_formats` List of file formats that this toolkit can read.

`toolkit_file_write_formats` List of file formats that this toolkit can write.

`toolkit_installation_instructions` classmethod(function) -> method

`toolkit_name` The name of the toolkit wrapped by this class.

#### Methods

<code>from_file(self, file_path, file_format[, ...])</code>	Return an <code>openforcefield.topology.Molecule</code> from a file using this toolkit.
<code>from_file_obj(self, file_obj, file_format[, ...])</code>	Return an <code>openforcefield.topology.Molecule</code> from a file-like object (an object with a “ <code>.read()</code> ” method using this
<code>is_available()</code>	Check whether the corresponding toolkit can be imported
<code>requires_toolkit()</code>	

`__init__(self, /, *args, **kwargs)`

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

## Methods

<code>__init__(self, /, *args, **kwargs)</code>	Initialize self.
<code>from_file(self, file_path, file_format[, ...])</code>	Return an openforcefield.topology.Molecule from a file using this toolkit.
<code>from_file_obj(self, file_obj, file_format[, ...])</code>	Return an openforcefield.topology.Molecule from a file-like object (an object with a “.read()” method) using this
<code>is_available()</code>	Check whether the corresponding toolkit can be imported
<code>requires_toolkit()</code>	

## Attributes

<code>toolkit_file_read_formats</code>	List of file formats that this toolkit can read.
<code>toolkit_file_write_formats</code>	List of file formats that this toolkit can write.
<code>toolkit_installation_instructions</code>	classmethod(function) -> method
<code>toolkit_name</code>	The name of the toolkit wrapped by this class.

### `property toolkit_name`

The name of the toolkit wrapped by this class.

### `property toolkit_installation_instructions`

classmethod(function) -> method

Convert a function to be a class method.

A class method receives the class as implicit first argument, just like an instance method receives the instance. To declare a class method, use this idiom:

```
class C: @classmethod def f(cls, arg1, arg2, ...):
```

```
...
```

It can be called either on the class (e.g. `C.f()`) or on an instance (e.g. `C().f()`). The instance is ignored except for its class. If a class method is called for a derived class, the derived class object is passed as the implied first argument.

Class methods are different than C++ or Java static methods. If you want those, see the `staticmethod` builtin.

### `property toolkit_file_read_formats`

List of file formats that this toolkit can read.

### `property toolkit_file_write_formats`

List of file formats that this toolkit can write.

### `static is_available()`

Check whether the corresponding toolkit can be imported

#### Returns

`is_installed` [bool] True if corresponding toolkit is installed, False otherwise.

### `from_file(self, file_path, file_format, allow_undefined_stereo=False)`

Return an openforcefield.topology.Molecule from a file using this toolkit.

#### Parameters

**file\_path** [str] The file to read the molecule from  
**file\_format** [str] Format specifier, usually file suffix (eg. ‘MOL2’, ‘SMI’) Note that not all toolkits support all formats. Check ToolkitWrapper.toolkit\_file\_read\_formats for details.  
**allow\_undefined\_stereo** [bool, default=False] If false, raises an exception if any molecules contain undefined stereochemistry.

#### Returns

---

**molecules** [Molecule or list of Molecules] a list of Molecule objects is returned.

**from\_file\_obj(self, file\_obj, file\_format, allow\_undefined\_stereo=False)**

Return an openforcefield.topology.Molecule from a file-like object (an object with a “.read()” method using toolkit.

#### Parameters

**file\_obj** [file-like object] The file-like object to read the molecule from  
**file\_format** [str] Format specifier, usually file suffix (eg. ‘MOL2’, ‘SMI’) Note that not all toolkits support all formats. Check ToolkitWrapper.toolkit\_file\_read\_formats for details.  
**allow\_undefined\_stereo** [bool, default=False] If false, raises an exception if any molecules contain undefined stereochemistry. If false, the function skips loading the molecule.

#### Returns

**molecules** [Molecule or list of Molecules] a list of Molecule objects is returned.

## openforcefield.utils.toolkits.OpenEyeToolkitWrapper

**class** openforcefield.utils.toolkits.OpenEyeToolkitWrapper  
OpenEye toolkit wrapper

**Warning:** This API is experimental and subject to change.

#### Attributes

**toolkit\_file\_read\_formats** List of file formats that this toolkit can read.  
**toolkit\_file\_write\_formats** List of file formats that this toolkit can write.  
**toolkit\_installation\_instructions** classmethod(function) -> method  
**toolkit\_name** The name of the toolkit wrapped by this class.

#### Methods

---

**compute\_partial\_charges(self, molecule[, ...])** Compute partial charges with OpenEye quacpac  
Continued on next page

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<code>compute_partial_charges_am1bcc(self, molecule)</code>	Compute AM1BCC partial charges with OpenEye quacpac
<code>compute_wiberg_bond_orders(self, molecule[, ...])</code>	Update and store list of bond orders this molecule.
<code>find_smarts_matches(self, molecule, smarts)</code>	Find all SMARTS matches for the specified molecule, using the specified aromaticity model.
<code>from_file(self, file_path, file_format[, ...])</code>	Return an openforcefield.topology.Molecule from a file using this toolkit.
<code>from_file_obj(self, file_obj, file_format[, ...])</code>	Return an openforcefield.topology.Molecule from a file-like object (an object with a “.read()” method using this toolkit.
<code>from_object(self, object)</code>	If given an OEMol (or OEMol-derived object), this function will load it into an openforcefield.topology.molecule. Otherwise, it will return False.
<code>from_openeye(oemol[, low_undefined_stereo])</code>	al- Create a Molecule from an OpenEye molecule.
<code>from_smiles(self, smiles[, ...])</code>	Create a Molecule from a SMILES string using the OpenEye toolkit.
<code>generate_conformers(self, molecule[, ...])</code>	Generate molecule conformers using OpenEye Omega.
<code>is_available([oetools])</code>	Check if the given OpenEye toolkit components are available.
<code>requires_toolkit()</code>	
<code>to_file(self, molecule, file_path, file_format)</code>	Writes an OpenFF Molecule to a file-like object
<code>to_file_obj(self, molecule, file_obj, ...)</code>	Writes an OpenFF Molecule to a file-like object
<code>to_openeye(molecule[, aromaticity_model])</code>	Create an OpenEye molecule using the specified aromaticity model
<code>to_smiles(molecule)</code>	Uses the OpenEye toolkit to convert a Molecule into a SMILES string.

**`__init__(self, /, *args, **kwargs)`**

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

**Methods**

<code>__init__(self, /, *args, **kwargs)</code>	Initialize self.
<code>compute_partial_charges(self, molecule[, ...])</code>	Compute partial charges with OpenEye quacpac
<code>compute_partial_charges_am1bcc(self, molecule)</code>	Compute AM1BCC partial charges with OpenEye quacpac
<code>compute_wiberg_bond_orders(self, molecule[, ...])</code>	Update and store list of bond orders this molecule.
<code>find_smarts_matches(self, molecule, smarts)</code>	Find all SMARTS matches for the specified molecule, using the specified aromaticity model.
<code>from_file(self, file_path, file_format[, ...])</code>	Return an openforcefield.topology.Molecule from a file using this toolkit.
<code>from_file_obj(self, file_obj, file_format[, ...])</code>	Return an openforcefield.topology.Molecule from a file-like object (an object with a “.read()” method using this toolkit.

Continued on next page

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<code>from_object(self, object)</code>	If given an OEMol (or OEMol-derived object), this function will load it into an openforcefield.topology.molecule. Otherwise, it will return False.
<code>from_openeye(oemol[, low_undefined_stereo])</code>	Create a Molecule from an OpenEye molecule.
<code>from_smiles(self, smiles[, ...])</code>	Create a Molecule from a SMILES string using the OpenEye toolkit.
<code>generate_conformers(self, molecule[, ...])</code>	Generate molecule conformers using OpenEye Omega.
<code>is_available([oetools])</code>	Check if the given OpenEye toolkit components are available.
<code>requires_toolkit()</code>	
<code>to_file(self, molecule, file_path, file_format)</code>	Writes an OpenFF Molecule to a file-like object
<code>to_file_obj(self, molecule, file_obj, ...)</code>	Writes an OpenFF Molecule to a file-like object
<code>to_openeye(molecule[, aromaticity_model])</code>	Create an OpenEye molecule using the specified aromaticity model
<code>to_smiles(molecule)</code>	Uses the OpenEye toolkit to convert a Molecule into a SMILES string.

## Attributes

<code>toolkit_file_read_formats</code>	List of file formats that this toolkit can read.
<code>toolkit_file_write_formats</code>	List of file formats that this toolkit can write.
<code>toolkit_installation_instructions</code>	classmethod(function) -> method
<code>toolkit_name</code>	The name of the toolkit wrapped by this class.

**static is\_available(oetools=(‘oechem’, ‘oequacpac’, ‘oeiupac’, ‘oeomega’))**

Check if the given OpenEye toolkit components are available.

If the OpenEye toolkit is not installed or no license is found for at least one the given toolkits , False is returned.

### Parameters

**oetools** [str or iterable of strings, optional, default=(‘oechem’, ‘oequacpac’, ‘oeiupac’, ‘oeomega’)] Set of tools to check by their Python module name. Defaults to the complete set of tools supported by this function. Also accepts a single tool to check as a string instead of an iterable of length 1.

### Returns

**all\_installed** [bool] True if all tools in oetools are installed and licensed, False otherwise

`from_object(self, object)`

If given an OEMol (or OEMol-derived object), this function will load it into an openforcefield.topology.molecule. Otherwise, it will return False.

### Parameters

**object** [A molecule-like object] An object to be type-checked.

### Returns

**Molecule** An openforcefield.topology.molecule Molecule.

**Raises**

**NotImplementedError** If the object could not be converted into a Molecule.

**from\_file(self, file\_path, file\_format, allow\_undefined\_stereo=False)**

Return an openforcefield.topology.Molecule from a file using this toolkit.

**Parameters**

**file\_path** [str] The file to read the molecule from

**file\_format** [str] Format specifier, usually file suffix (eg. ‘MOL2’, ‘SMI’) Note that not all toolkits support all formats. Check ToolkitWrapper.toolkit\_file\_read\_formats for details.

**allow\_undefined\_stereo** [bool, default=False] If false, raises an exception if oemol contains undefined stereochemistry.

**Returns**

**molecules** [List[Molecule]] The list of Molecule objects in the file.

**Raises**

**GAFFAtomTypeWarning** If the loaded mol2 file possibly uses GAFF atom types, which are not supported.

**Examples**

Load a mol2 file into an OpenFF Molecule object.

```
>>> from openforcefield.utils import get_data_file_path
>>> mol2_file_path = get_data_file_path('molecules/cyclohexane.mol2')
>>> toolkit = OpenEyeToolkitWrapper()
>>> molecule = toolkit.from_file(mol2_file_path, file_format='mol2')
```

**from\_file\_obj(self, file\_obj, file\_format, allow\_undefined\_stereo=False)**

Return an openforcefield.topology.Molecule from a file-like object (an object with a “.read()” method using this toolkit.

**Parameters**

**file\_obj** [file-like object] The file-like object to read the molecule from

**file\_format** [str] Format specifier, usually file suffix (eg. ‘MOL2’, ‘SMI’) Note that not all toolkits support all formats. Check ToolkitWrapper.toolkit\_file\_read\_formats for details.

**allow\_undefined\_stereo** [bool, default=False] If false, raises an exception if oemol contains undefined stereochemistry.

**Returns**

**molecules** [List[Molecule]] The list of Molecule objects in the file object.

**Raises**

**GAFFAtomTypeWarning** If the loaded mol2 file possibly uses GAFF atom types, which are not supported.

**to\_file\_obj(self, molecule, file\_obj, file\_format)**

Writes an OpenFF Molecule to a file-like object

**Parameters**

**molecule** [an OpenFF Molecule] The molecule to write

**file\_obj** The file-like object to write to

**file\_format** The format for writing the molecule data

**to\_file(self, molecule, file\_path, file\_format)**

Writes an OpenFF Molecule to a file-like object

#### Parameters

**molecule** [an OpenFF Molecule] The molecule to write

**file\_path** The file path to write to.

**file\_format** The format for writing the molecule data

**static from\_openeye(oemol, allow\_undefined\_stereo=False)**

Create a Molecule from an OpenEye molecule.

**Warning:** This API is experimental and subject to change.

#### Parameters

**oemol** [openeye.oecchem.OEMol] An OpenEye molecule

**allow\_undefined\_stereo** [bool, default=False] If false, raises an exception if oemol contains undefined stereochemistry.

#### Returns

**molecule** [openforcefield.topology.Molecule] An openforcefield molecule

### Examples

Create a Molecule from an OpenEye OEMol

```
>>> from openeye import oecchem
>>> from openforcefield.tests.utils import get_data_file_path
>>> ifs = oecchem.oemolistream(get_data_file_path('systems/monomers/ethanol.mol2'))
>>> oemols = list(ifs.GetOEGraphMols())
```

```
>>> toolkit_wrapper = OpenEyeToolkitWrapper()
>>> molecule = toolkit_wrapper.from_openeye(oemols[0])
```

**static to\_openeye(molecule, aromaticity\_model='OEArroModel\_MDL')**

Create an OpenEye molecule using the specified aromaticity model

**Warning:** This API is experimental and subject to change.

#### Parameters

**molecule** [openforcefield.topology.molecule.Molecule object]

The molecule to convert to an OEMol

**aromaticity\_model** [str, optional, default=DEFAULT\_AROMATICITY\_MODEL]  
The aromaticity model to use

**Returns**

**oemol** [openeye.oecchem.OEMol] An OpenEye molecule

**Examples**

Create an OpenEye molecule from a Molecule

```
>>> from openforcefield.topology import Molecule
>>> toolkit_wrapper = OpenEyeToolkitWrapper()
>>> molecule = Molecule.from_smiles('CC')
>>> oemol = toolkit_wrapper.to_openeye(molecule)
```

**static to\_smiles(molecule)**

Uses the OpenEye toolkit to convert a Molecule into a SMILES string.

**Parameters**

**molecule** [An openforcefield.topology.Molecule] The molecule to convert into a SMILES.

**Returns**

**smiles** [str] The SMILES of the input molecule.

**from\_smiles(self, smiles, hydrogens\_are\_explicit=False)**

Create a Molecule from a SMILES string using the OpenEye toolkit.

**Warning:** This API is experimental and subject to change.

**Parameters**

**smiles** [str] The SMILES string to turn into a molecule

**hydrogens\_are\_explicit** [bool, default = False] If False, OE will perform hydrogen addition using OEAddExplicitHydrogens

**Returns**

**molecule** [openforcefield.topology.Molecule] An openforcefield-style molecule.

**generate\_conformers(self, molecule, n\_conformers=1, clear\_existing=True)**

Generate molecule conformers using OpenEye Omega.

**Warning:** This API is experimental and subject to change.

**molecule** [a Molecule ] The molecule to generate conformers for.

**n\_conformers** [int, default=1] The maximum number of conformers to generate.

**clear\_existing** [bool, default=True] Whether to overwrite existing conformers for the molecule

```
compute_partial_charges(self, molecule, quantum_chemical_method='AM1-BCC', partial_charge_method='None')
Compute partial charges with OpenEye quacpac
```

**Warning:** This API is experimental and subject to change.

#### Parameters

**molecule** [Molecule] Molecule for which partial charges are to be computed  
**charge\_model** [str, optional, default=None] The charge model to use. One of ['noop', 'mmff', 'mmff94', 'am1bcc', 'am1bccnosymspt', 'amber', 'amberff94', 'am1bccelf10'] If None, 'am1bcc' will be used.

#### Returns

**charges** [numpy.array of shape (natoms) of type float] The partial charges

```
compute_partial_charges_am1bcc(self, molecule)
Compute AM1BCC partial charges with OpenEye quacpac
```

**Warning:** This API is experimental and subject to change.

#### Parameters

**molecule** [Molecule] Molecule for which partial charges are to be computed

#### Returns

**charges** [numpy.array of shape (natoms) of type float] The partial charges

```
compute_wiberg_bond_orders(self, molecule, charge_model=None)
Update and store list of bond orders this molecule. Can be used for initialization of bondorders list, or for updating bond orders in the list.
```

**Warning:** This API is experimental and subject to change.

#### Parameters

**molecule** [openforcefield.topology.molecule Molecule] The molecule to assign wiberg bond orders to  
**charge\_model** [str, optional, default=None] The charge model to use. One of ['am1', 'pm3']. If None, 'am1' will be used.

```
find_smarts_matches(self, molecule, smarts, aromaticity_model='OEArroModel_MDL')
Find all SMARTS matches for the specified molecule, using the specified aromaticity model.
```

**Warning:** This API is experimental and subject to change.

#### Parameters

---

**molecule** [openforcefield.topology.Molecule] The molecule for which all specified SMARTS matches are to be located

**smarts** [str] SMARTS string with optional SMIRKS-style atom tagging

**aromaticity\_model** [str, optional, default='OEArroModel\_MDL'] Aromaticity model to use during matching

**.. note ::** Currently, the only supported “aromaticity\_model“ is “OEArroModel\_MDL“

**property toolkit\_file\_read\_formats**

List of file formats that this toolkit can read.

**property toolkit\_file\_write\_formats**

List of file formats that this toolkit can write.

**property toolkit\_installation\_instructions**

classmethod(function) -> method

Convert a function to be a class method.

A class method receives the class as implicit first argument, just like an instance method receives the instance. To declare a class method, use this idiom:

```
class C: @classmethod def f(cls, arg1, arg2, ...):
```

...

It can be called either on the class (e.g. C.f()) or on an instance (e.g. C().f()). The instance is ignored except for its class. If a class method is called for a derived class, the derived class object is passed as the implied first argument.

Class methods are different than C++ or Java static methods. If you want those, see the statmethod builtin.

**property toolkit\_name**

The name of the toolkit wrapped by this class.

**openforcefield.utils.toolkits.RDKitToolkitWrapper**

```
class openforcefield.utils.toolkits.RDKitToolkitWrapper
    RDKit toolkit wrapper
```

**Warning:** This API is experimental and subject to change.

**Attributes**

**toolkit\_file\_read\_formats** List of file formats that this toolkit can read.

**toolkit\_file\_write\_formats** List of file formats that this toolkit can write.

**toolkit\_installation\_instructions** classmethod(function) -> method

**toolkit\_name** The name of the toolkit wrapped by this class.

**Methods**

<code>find_smarts_matches(self, molecule, smarts)</code>	Find all SMARTS matches for the specified molecule, using the specified aromaticity model.
<code>from_file(self, file_path, file_format[, ...])</code>	Create an openforcefield.topology.Molecule from a file using this toolkit.
<code>from_file_obj(self, file_obj, file_format[, ...])</code>	Return an openforcefield.topology.Molecule from a file-like object (an object with a “.read()” method using this toolkit.
<code>from_object(self, object)</code>	If given an rdchem.Mol (or rdchem.Mol-derived object), this function will load it into an openforcefield.topology.molecule.
<code>from_rdkit(self, rdmol[, low_undefined_stereo])</code>	Create a Molecule from an RDKit molecule.
<code>from_smiles(self, smiles[, ...])</code>	Create a Molecule from a SMILES string using the RDKit toolkit.
<code>generate_conformers(self, molecule[, ...])</code>	Generate molecule conformers using RDKit.
<code>is_available()</code>	Check whether the RDKit toolkit can be imported
<code>requires_toolkit()</code>	
<code>to_file(self, molecule, file_path, file_format)</code>	Writes an OpenFF Molecule to a file-like object
<code>to_file_obj(self, molecule, file_obj, ...)</code>	Writes an OpenFF Molecule to a file-like object
<code>to_rdkit(molecule[, aromaticity_model])</code>	Create an RDKit molecule
<code>to_smiles(molecule)</code>	Uses the RDKit toolkit to convert a Molecule into a SMILES string.

### `__init__(self, /, *args, **kwargs)`

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

## Methods

<code>__init__(self, /, *args, **kwargs)</code>	Initialize self.
<code>find_smarts_matches(self, molecule, smarts)</code>	Find all SMARTS matches for the specified molecule, using the specified aromaticity model.
<code>from_file(self, file_path, file_format[, ...])</code>	Create an openforcefield.topology.Molecule from a file using this toolkit.
<code>from_file_obj(self, file_obj, file_format[, ...])</code>	Return an openforcefield.topology.Molecule from a file-like object (an object with a “.read()” method using this toolkit.
<code>from_object(self, object)</code>	If given an rdchem.Mol (or rdchem.Mol-derived object), this function will load it into an openforcefield.topology.molecule.
<code>from_rdkit(self, rdmol[, low_undefined_stereo])</code>	Create a Molecule from an RDKit molecule.
<code>from_smiles(self, smiles[, ...])</code>	Create a Molecule from a SMILES string using the RDKit toolkit.
<code>generate_conformers(self, molecule[, ...])</code>	Generate molecule conformers using RDKit.
<code>is_available()</code>	Check whether the RDKit toolkit can be imported
<code>requires_toolkit()</code>	
<code>to_file(self, molecule, file_path, file_format)</code>	Writes an OpenFF Molecule to a file-like object
<code>to_file_obj(self, molecule, file_obj, ...)</code>	Writes an OpenFF Molecule to a file-like object

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<code>to_rdkit(molecule[, aromaticity_model])</code>	Create an RDKit molecule
<code>to_smiles(molecule)</code>	Uses the RDKit toolkit to convert a Molecule into a SMILES string.

## Attributes

<code>toolkit_file_read_formats</code>	List of file formats that this toolkit can read.
<code>toolkit_file_write_formats</code>	List of file formats that this toolkit can write.
<code>toolkit_installation_instructions</code>	classmethod(function) -> method
<code>toolkit_name</code>	The name of the toolkit wrapped by this class.

**static is\_available()**

Check whether the RDKit toolkit can be imported

### Returns

**is\_installed** [bool] True if RDKit is installed, False otherwise.

**from\_object(self, object)**

If given an rdchem.Mol (or rdchem.Mol-derived object), this function will load it into an open-forcefield.topology.molecule. Otherwise, it will return False.

### Parameters

**object** [A rdchem.Mol-derived object] An object to be type-checked and converted into a Molecule, if possible.

### Returns

**Molecule or False** An openforcefield.topology.molecule Molecule.

### Raises

**NotImplementedError** If the object could not be converted into a Molecule.

**from\_file(self, file\_path, file\_format, allow\_undefined\_stereo=False)**

Create an openforcefield.topology.Molecule from a file using this toolkit.

### Parameters

**file\_path** [str] The file to read the molecule from

**file\_format** [str] Format specifier, usually file suffix (eg. ‘MOL2’, ‘SMI’) Note that not all toolkits support all formats. Check ToolkitWrapper.toolkit\_file\_read\_formats for details.

**allow\_undefined\_stereo** [bool, default=False] If false, raises an exception if oemol contains undefined stereochemistry.

### Returns

**molecules** [iterable of Molecules] a list of Molecule objects is returned.

**from\_file\_obj(self, file\_obj, file\_format, allow\_undefined\_stereo=False)**

Return an openforcefield.topology.Molecule from a file-like object (an object with a “.read()” method using this toolkit.

**Warning:** This API is experimental and subject to change.

**Parameters**

**file\_obj** [file-like object] The file-like object to read the molecule from

**file\_format** [str] Format specifier, usually file suffix (eg. ‘MOL2’, ‘SMI’) Note that not all toolkits support all formats. Check ToolkitWrapper.toolkit\_file\_read\_formats for details.

**allow\_undefined\_stereo** [bool, default=False] If false, raises an exception if oemol contains undefined stereochemistry.

**Returns**

**molecules** [Molecule or list of Molecules] a list of Molecule objects is returned.

**to\_file\_obj(self, molecule, file\_obj, file\_format)**  
Writes an OpenFF Molecule to a file-like object

**Parameters**

**molecule** [an OpenFF Molecule] The molecule to write

**file\_obj** The file-like object to write to

**file\_format** The format for writing the molecule data

**to\_file(self, molecule, file\_path, file\_format)**  
Writes an OpenFF Molecule to a file-like object

**Parameters**

**molecule** [an OpenFF Molecule] The molecule to write

**file\_path** The file path to write to

**file\_format** The format for writing the molecule data

**Returns**

---

**classmethod to\_smiles(molecule)**  
Uses the RDKit toolkit to convert a Molecule into a SMILES string.

**Parameters**

**molecule** [An openforcefield.topology.Molecule] The molecule to convert into a SMILES.

**Returns**

**smiles** [str] The SMILES of the input molecule.

**from\_smiles(self, smiles, hydrogens\_are\_explicit=False)**  
Create a Molecule from a SMILES string using the RDKit toolkit.

**Warning:** This API is experimental and subject to change.

**Parameters**

**smiles** [str] The SMILES string to turn into a molecule

**hydrogens\_are\_explicit** [bool, default=False] If False, RDKit will perform hydrogen addition using Chem.AddHs

**Returns**

**molecule** [openforcefield.topology.Molecule] An openforcefield-style molecule.

**generate\_conformers(self, molecule, n\_conformers=1, clear\_existing=True)**  
Generate molecule conformers using RDKit.

**Warning:** This API is experimental and subject to change.

**molecule** [a Molecule ] The molecule to generate conformers for.

**n\_conformers** [int, default=1] Maximum number of conformers to generate.

**clear\_existing** [bool, default=True] Whether to overwrite existing conformers for the molecule.

**from\_rdkit(self, rdmol, allow\_undefined\_stereo=False)**  
Create a Molecule from an RDKit molecule.

Requires the RDKit to be installed.

**Warning:** This API is experimental and subject to change.

**Parameters**

**rdmol** [rkit.RDMol] An RDKit molecule

**allow\_undefined\_stereo** [bool, default=False] If false, raises an exception if rdmol contains undefined stereochemistry.

**Returns**

**molecule** [openforcefield.Molecule] An openforcefield molecule

**Examples**

Create a molecule from an RDKit molecule

```
>>> from rdkit import Chem
>>> from openforcefield.tests.utils import get_data_file_path
>>> rdmol = Chem.MolFromMolFile(get_data_file_path('systems/monomers/ethanol.sdf'))
```

```
>>> toolkit_wrapper = RDKitToolkitWrapper()
>>> molecule = toolkit_wrapper.from_rdkit(rdmol)
```

**classmethod to\_rdkit(molecule, aromaticity\_model='OEAroModel\_MDL')**  
Create an RDKit molecule

Requires the RDKit to be installed.

**Warning:** This API is experimental and subject to change.

**Parameters**

**aromaticity\_model** [str, optional, default=DEFAULT\_AROMATICITY\_MODEL] The aromaticity model to use

**Returns**

**rdmol** [rkit.RDMol] An RDKit molecule

**Examples**

Convert a molecule to RDKit

```
>>> from openforcefield.topology import Molecule
>>> ethanol = Molecule.from_smiles('CCO')
>>> rdmol = ethanol.to_rdkit()
```

**find\_smarts\_matches**(self, molecule, smarts, aromaticity\_model='OEArroModel\_MDL')

Find all SMARTS matches for the specified molecule, using the specified aromaticity model.

**Warning:** This API is experimental and subject to change.

**Parameters**

**molecule** [openforcefield.topology.Molecule] The molecule for which all specified SMARTS matches are to be located

**smarts** [str] SMARTS string with optional SMIRKS-style atom tagging

**aromaticity\_model** [str, optional, default='OEArroModel\_MDL'] Aromaticity model to use during matching

.. note :: Currently, the only supported “aromaticity\_model“ is “OEArroModel\_MDL“

**property toolkit\_file\_read\_formats**

List of file formats that this toolkit can read.

**property toolkit\_file\_write\_formats**

List of file formats that this toolkit can write.

**property toolkit\_installation\_instructions**

classmethod(function) -> method

Convert a function to be a class method.

A class method receives the class as implicit first argument, just like an instance method receives the instance. To declare a class method, use this idiom:

```
class C: @classmethod def f(cls, arg1, arg2, ...):
```

...

It can be called either on the class (e.g. C.f()) or on an instance (e.g. C().f()). The instance is ignored except for its class. If a class method is called for a derived class, the derived class object is passed as the implied first argument.

Class methods are different than C++ or Java static methods. If you want those, see the staticmethod builtin.

**property toolkit\_name**

The name of the toolkit wrapped by this class.

**openforcefield.utils.toolkits.AmberToolsToolkitWrapper**

```
class openforcefield.utils.toolkits.AmberToolsToolkitWrapper
    AmberTools toolkit wrapper
```

**Warning:** This API is experimental and subject to change.

**Attributes**

- toolkit\_file\_read\_formats** List of file formats that this toolkit can read.
- toolkit\_file\_write\_formats** List of file formats that this toolkit can write.
- toolkit\_installation\_instructions** classmethod(function) -> method
- toolkit\_name** The name of the toolkit wrapped by this class.

**Methods**

<code>compute_partial_charges(self, molecule[, ...])</code>	Compute partial charges with AmberTools using antechamber/sqm
<code>compute_partial_charges_am1bcc(self, molecule)</code>	Compute partial charges with AmberTools using antechamber/sqm.
<code>from_file(self, file_path, file_format[, ...])</code>	Return an openforcefield.topology.Molecule from a file using this toolkit.
<code>from_file_obj(self, file_obj, file_format[, ...])</code>	Return an openforcefield.topology.Molecule from a file-like object (an object with a “.read()” method using this
<code>is_available()</code>	Check whether the AmberTools toolkit is installed
<code>requires_toolkit()</code>	

**\_\_init\_\_(self)**

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

**Methods**

<code>__init__(self)</code>	Initialize self.
<code>compute_partial_charges(self, molecule[, ...])</code>	Compute partial charges with AmberTools using antechamber/sqm
<code>compute_partial_charges_am1bcc(self, molecule)</code>	Compute partial charges with AmberTools using antechamber/sqm.
<code>from_file(self, file_path, file_format[, ...])</code>	Return an openforcefield.topology.Molecule from a file using this toolkit.
<code>from_file_obj(self, file_obj, file_format[, ...])</code>	Return an openforcefield.topology.Molecule from a file-like object (an object with a “.read()” method using this
<code>is_available()</code>	Check whether the AmberTools toolkit is installed
<code>requires_toolkit()</code>	

## Attributes

<code>toolkit_file_read_formats</code>	List of file formats that this toolkit can read.
<code>toolkit_file_write_formats</code>	List of file formats that this toolkit can write.
<code>toolkit_installation_instructions</code>	classmethod(function) -> method
<code>toolkit_name</code>	The name of the toolkit wrapped by this class.

### `static is_available()`

Check whether the AmberTools toolkit is installed

#### Returns

`is_installed` [bool] True if AmberTools is installed, False otherwise.

### `compute_partial_charges(self, molecule, charge_model=None)`

Compute partial charges with AmberTools using antechamber/sqm

**Warning:** This API experimental and subject to change.

#### Parameters

`molecule` [Molecule] Molecule for which partial charges are to be computed

`charge_model` [str, optional, default=None] The charge model to use. One of ['gas', 'mul', 'bcc']. If None, 'bcc' will be used.

#### Raises

`ValueError` if the requested charge method could not be handled

#### Notes

Currently only sdf file supported as input and mol2 as output <https://github.com/choderalab/openmoltools/blob/master/openmoltools/packmol.py>

### `compute_partial_charges_am1bcc(self, molecule)`

Compute partial charges with AmberTools using antechamber/sqm. This will calculate AM1-BCC charges on the first conformer only.

**Warning:** This API experimental and subject to change.

#### Parameters

`molecule` [Molecule] Molecule for which partial charges are to be computed

#### Raises

`ValueError` if the requested charge method could not be handled

### `from_file(self, file_path, file_format, allow_undefined_stereo=False)`

Return an openforcefield.topology.Molecule from a file using this toolkit.

#### Parameters

`file_path` [str] The file to read the molecule from

**file\_format** [str] Format specifier, usually file suffix (eg. ‘MOL2’, ‘SMI’) Note that not all toolkits support all formats. Check ToolkitWrapper.toolkit\_file\_read\_formats for details.

**allow\_undefined\_stereo** [bool, default=False] If false, raises an exception if any molecules contain undefined stereochemistry.

### Returns

---

**molecules** [Molecule or list of Molecules] a list of Molecule objects is returned.

**from\_file\_obj(self, file\_obj, file\_format, allow\_undefined\_stereo=False)**

Return an openforcefield.topology.Molecule from a file-like object (an object with a “.read()” method using toolkit.

### Parameters

**file\_obj** [file-like object] The file-like object to read the molecule from

**file\_format** [str] Format specifier, usually file suffix (eg. ‘MOL2’, ‘SMI’) Note that not all toolkits support all formats. Check ToolkitWrapper.toolkit\_file\_read\_formats for details.

**allow\_undefined\_stereo** [bool, default=False] If false, raises an exception if any molecules contain undefined stereochemistry. If false, the function skips loading the molecule.

### Returns

**molecules** [Molecule or list of Molecules] a list of Molecule objects is returned.

**property toolkit\_file\_read\_formats**

List of file formats that this toolkit can read.

**property toolkit\_file\_write\_formats**

List of file formats that this toolkit can write.

**property toolkit\_installation\_instructions**

classmethod(function) -> method

Convert a function to be a class method.

A class method receives the class as implicit first argument, just like an instance method receives the instance. To declare a class method, use this idiom:

**class C: @classmethod def f(cls, arg1, arg2, ...):**

...

It can be called either on the class (e.g. C.f()) or on an instance (e.g. C().f()). The instance is ignored except for its class. If a class method is called for a derived class, the derived class object is passed as the implied first argument.

Class methods are different than C++ or Java static methods. If you want those, see the staticmethod builtin.

**property toolkit\_name**

The name of the toolkit wrapped by this class.

### 2.3.2 Serialization support

Serializable	Mix-in to add serialization and deserialization support via JSON, YAML, BSON, TOML, MessagePack, and XML.
--------------	---

#### openforcefield.utils.serialization.Serializable

```
class openforcefield.utils.serialization.Serializable
```

Mix-in to add serialization and deserialization support via JSON, YAML, BSON, TOML, MessagePack, and XML.

For more information on these formats, see: [JSON](#), [BSON](#), [YAML](#), [TOML](#), [MessagePack](#), and [XML](#).

To use this mix-in, the class inheriting from this class must have implemented `to_dict()` and `from_dict()` methods that utilize dictionaries containing only serializable Python objects.

**Warning:** The serialization/deserialization schemes used here place some strict constraints on what kinds of dict objects can be serialized. No effort is made to add further protection to ensure serialization is possible. Use with caution.

#### Examples

Example class using `Serializable` mix-in:

```
>>> from openforcefield.utils.serialization import Serializable
>>> class Thing(Serializable):
...     def __init__(self, description):
...         self.description = description
...
...     def to_dict(self):
...         return { 'description' : self.description }
...
...     @classmethod
...     def from_dict(cls, d):
...         return cls(d['description'])
...
>>> # Create an example object
>>> thing = Thing('blobb')
```

Get `JSON` representation:

```
>>> json_thing = thing.to_json()
```

Reconstruct an object from its `JSON` representation:

```
>>> thing_from_json = Thing.from_json(json_thing)
```

Get `BSON` representation:

```
>>> bson_thing = thing.to_bson()
```

Reconstruct an object from its `BSON` representation:

```
>>> thing_from_bson = Thing.from_bson(bson_thing)
```

Get [YAML](#) representation:

```
>>> yaml_thing = thing.to_yaml()
```

Reconstruct an object from its [YAML](#) representation:

```
>>> thing_from_yaml = Thing.from_yaml(yaml_thing)
```

Get [MessagePack](#) representation:

```
>>> messagepack_thing = thing.to_messagepack()
```

Reconstruct an object from its [MessagePack](#) representation:

```
>>> thing_from_messagepack = Thing.from_messagepack(messagepack_thing)
```

Get [XML](#) representation:

```
>>> xml_thing = thing.to_xml()
```

## Methods

<code>from_bson(serialized)</code>	Instantiate an object from a BSON serialized representation.
<code>from_json(serialized)</code>	Instantiate an object from a JSON serialized representation.
<code>from_messagepack(serialized)</code>	Instantiate an object from a MessagePack serialized representation.
<code>from_pickle(serialized)</code>	Instantiate an object from a pickle serialized representation.
<code>from_toml(serialized)</code>	Instantiate an object from a TOML serialized representation.
<code>from_xml(serialized)</code>	Instantiate an object from an XML serialized representation.
<code>from_yaml(serialized)</code>	Instantiate from a YAML serialized representation.
<code>to_bson(self)</code>	Return a BSON serialized representation.
<code>to_json(self[, indent])</code>	Return a JSON serialized representation.
<code>to_messagepack(self)</code>	Return a MessagePack representation.
<code>to_pickle(self)</code>	Return a pickle serialized representation.
<code>to_toml(self)</code>	Return a TOML serialized representation.
<code>to_xml(self[, indent])</code>	Return an XML representation.
<code>to_yaml(self)</code>	Return a YAML serialized representation.

<code>from_dict</code>	
<code>to_dict</code>	

`__init__(self, /, *args, **kwargs)`

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

## Methods

<code>__init__(self, /, *args, **kwargs)</code>	Initialize self.
<code>from_bson(serialized)</code>	Instantiate an object from a BSON serialized representation.
<code>from_dict(d)</code>	
<code>from_json(serialized)</code>	Instantiate an object from a JSON serialized representation.
<code>from_messagepack(serialized)</code>	Instantiate an object from a MessagePack serialized representation.
<code>from_pickle(serialized)</code>	Instantiate an object from a pickle serialized representation.
<code>from_toml(serialized)</code>	Instantiate an object from a TOML serialized representation.
<code>from_xml(serialized)</code>	Instantiate an object from an XML serialized representation.
<code>from_yaml(serialized)</code>	Instantiate from a YAML serialized representation.
<code>to_bson(self)</code>	Return a BSON serialized representation.
<code>to_dict(self)</code>	
<code>to_json(self[, indent])</code>	Return a JSON serialized representation.
<code>to_messagepack(self)</code>	Return a MessagePack representation.
<code>to_pickle(self)</code>	Return a pickle serialized representation.
<code>to_toml(self)</code>	Return a TOML serialized representation.
<code>to_xml(self[, indent])</code>	Return an XML representation.
<code>to_yaml(self)</code>	Return a YAML serialized representation.

`to_json(self, indent=None)`

Return a JSON serialized representation.

Specification: <https://www.json.org/>

### Parameters

`indent` [int, optional, default=None] If not None, will pretty-print with specified number of spaces for indentation

### Returns

`serialized` [str] A JSON serialized representation of the object

`classmethod from_json(serialized)`

Instantiate an object from a JSON serialized representation.

Specification: <https://www.json.org/>

### Parameters

`serialized` [str] A JSON serialized representation of the object

### Returns

`instance` [cls] An instantiated object

`to_bson(self)`

Return a BSON serialized representation.

Specification: <http://bsonspec.org/>

**Returns**

**serialized** [bytes] A BSON serialized representation of the object

**classmethod from\_bson(serialized)**

Instantiate an object from a BSON serialized representation.

Specification: <http://bsonspec.org/>

**Parameters**

**serialized** [bytes] A BSON serialized representation of the object

**Returns**

**instance** [cls] An instantiated object

**to\_toml(self)**

Return a TOML serialized representation.

Specification: <https://github.com/toml-lang/toml>

**Returns**

**serialized** [str] A TOML serialized representation of the object

**classmethod from\_toml(serialized)**

Instantiate an object from a TOML serialized representation.

Specification: <https://github.com/toml-lang/toml>

**Parameters**

**serialized** [str] A TOML serialized representation of the object

**Returns**

**instance** [cls] An instantiated object

**to\_yaml(self)**

Return a YAML serialized representation.

Specification: <http://yaml.org/>

**Returns**

**serialized** [str] A YAML serialized representation of the object

**classmethod from\_yaml(serialized)**

Instantiate from a YAML serialized representation.

Specification: <http://yaml.org/>

**Parameters**

**serialized** [str] A YAML serialized representation of the object

**Returns**

**instance** [cls] Instantiated object

**to\_messagepack(self)**

Return a MessagePack representation.

Specification: <https://msgpack.org/index.html>

**Returns**

**serialized** [bytes] A MessagePack-encoded bytes serialized representation of the object

**classmethod from\_messagepack(serialized)**

Instantiate an object from a MessagePack serialized representation.

Specification: <https://msgpack.org/index.html>

**Parameters**

**serialized** [bytes] A MessagePack-encoded bytes serialized representation

**Returns**

**instance** [cls] Instantiated object.

**to\_xml(self, indent=2)**

Return an XML representation.

Specification: <https://www.w3.org/XML/>

**Parameters**

**indent** [int, optional, default=2] If not None, will pretty-print with specified number of spaces for indentation

**Returns**

**serialized** [bytes] A MessagePack-encoded bytes serialized representation.

**classmethod from\_xml(serialized)**

Instantiate an object from an XML serialized representation.

Specification: <https://www.w3.org/XML/>

**Parameters**

**serialized** [bytes] An XML serialized representation

**Returns**

**instance** [cls] Instantiated object.

**to\_pickle(self)**

Return a pickle serialized representation.

**Warning:** This is not recommended for safe, stable storage since the pickle specification may change between Python versions.

**Returns**

**serialized** [str] A pickled representation of the object

**classmethod from\_pickle(serialized)**

Instantiate an object from a pickle serialized representation.

**Warning:** This is not recommended for safe, stable storage since the pickle specification may change between Python versions.

**Parameters**

**serialized** [str] A pickled representation of the object

**Returns**

**instance** [cls] An instantiated object

### 2.3.3 Miscellaneous utilities

Miscellaneous utility functions.

<code>inherit_docstrings</code>	Inherit docstrings from parent class
<code>all_subclasses</code>	Recursively retrieve all subclasses of the specified class
<code>temporary_cd</code>	Context to temporary change the working directory.
<code>temporary_directory</code>	Context for safe creation of temporary directories.
<code>get_data_file_path</code>	Get the full path to one of the reference files in test-systems.
<code>convert_0_1_smirnoff_to_0_2</code>	Convert an 0.1-compliant SMIRNOFF dict to an 0.2-compliant one.
<code>convert_0_2_smirnoff_to_0_3</code>	Convert an 0.2-compliant SMIRNOFF dict to an 0.3-compliant one.

#### `openforcefield.utils.utils.inherit_docstrings`

`openforcefield.utils.utils.inherit_docstrings(cls)`

Inherit docstrings from parent class

#### `openforcefield.utils.utils.all_subclasses`

`openforcefield.utils.utils.all_subclasses(cls)`

Recursively retrieve all subclasses of the specified class

#### `openforcefield.utils.utils.temporary_cd`

`openforcefield.utils.utils.temporary_cd(dir_path)`

Context to temporary change the working directory.

**Parameters**

**dir\_path** [str] The directory path to enter within the context

**Examples**

```
>>> dir_path = '/tmp'
>>> with temporary_cd(dir_path):
...     pass # do something in dir_path
```

## `openforcefield.utils.utils.temporary_directory`

`openforcefield.utils.utils.temporary_directory()`  
Context for safe creation of temporary directories.

## `openforcefield.utils.utils.get_data_file_path`

`openforcefield.utils.utils.get_data_file_path(relative_path)`  
Get the full path to one of the reference files in testsystems. In the source distribution, these files are in `openforcefield/data/`, but on installation, they're moved to somewhere in the user's python site-packages directory. Parameters ——— name : str  
Name of the file to load (with respect to the repex folder).

## `openforcefield.utils.utils.convert_0_1_smirnoff_to_0_2`

`openforcefield.utils.utils.convert_0_1_smirnoff_to_0_2(smirnoff_data_0_1)`  
Convert an 0.1-compliant SMIRNOFF dict to an 0.2-compliant one. This involves renaming several tags, adding Electrostatics and ToolkitAM1BCC tags, and separating improper torsions into their own section.

### Parameters

`smirnoff_data_0_1` [dict] Hierarchical dict representing a SMIRNOFF data structure according the the 0.1 spec

### Returns

`smirnoff_data_0_2` Hierarchical dict representing a SMIRNOFF data structure according the the 0.2 spec

## `openforcefield.utils.utils.convert_0_2_smirnoff_to_0_3`

`openforcefield.utils.utils.convert_0_2_smirnoff_to_0_3(smirnoff_data_0_2)`  
Convert an 0.2-compliant SMIRNOFF dict to an 0.3-compliant one. This involves removing units from header tags and adding them to attributes of child elements. It also requires converting ProperTorsions and ImproperTorsions potentials from “charmm” to “fourier”.

### Parameters

`smirnoff_data_0_2` [dict] Hierarchical dict representing a SMIRNOFF data structure according the the 0.2 spec

### Returns

`smirnoff_data_0_3` Hierarchical dict representing a SMIRNOFF data structure according the the 0.3 spec

## 2.3.4 Structure tools

Tools for manipulating molecules and structures

**Warning:** These methods are deprecated and will be removed and replaced with integrated API methods. We recommend that no new code makes use of these functions.

---

<code>get_molecule_parameterIDs</code>	Process a list of molecules with a specified SMIRNOFF fxml file and determine which parameters are used by which molecules, returning collated results.
--	---

---

## `openforcefield.utils.structure.get_molecule_parameterIDs`

`openforcefield.utils.structure.get_molecule_parameterIDs(molecules, forcefield)`

Process a list of molecules with a specified SMIRNOFF fxml file and determine which parameters are used by which molecules, returning collated results.

### Parameters

**molecules** [list of `openforcefield.topology.Molecule`] List of molecules (with explicit hydrogens) to parse

**forcefield** [`openforcefield.typing.engines.smirnoff.ForceField`] The ForceField to apply

### Returns

**parameters\_by\_molecule** [dict] Parameter IDs used in each molecule, keyed by isomeric SMILES generated from provided OEMols. Each entry in the dict is a list which does not necessarily have unique entries; i.e. parameter IDs which are used more than once will occur multiple times.

**parameters\_by\_ID** [dict] Molecules in which each parameter ID occur, keyed by parameter ID. Each entry in the dict is a set of isomeric SMILES for molecules in which that parameter occurs. No frequency information is stored.



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