




molify: Molecular Structure Interface

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Summary

The increasing prevalence of Machine-Learned Interatomic Potentials (MLIPs) has shifted requirements for setting up atomistic simulations. Unlike classical force fields, MLIPs primarily require atomic positions and species, thereby removing the need for predefined topology files used for classical force fields in molecular dynamics software like GROMACS ([Abraham et al., 2015](#)), LAMMPS ([Thompson et al., 2022](#)), ESPResSo ([Weik et al., 2019](#)), or OpenMM ([Eastman et al., 2024](#)). Consequently, the Atomic Simulation Environment (ASE) ([Larsen et al., 2017](#)) has become a popular Python toolkit for handling atomic structures and interfacing with MLIPs, particularly within the material science and soft matter communities. ASE originates from the electronic structure community, which shares the same setup as MLIP-driven studies. In contrast to *ab initio*, MLIPs are much faster and can run on much larger systems, making high-throughput simulations of more complex systems feasible and increasing the need for efficient initial structure generation.

Concurrently, RDKit ([Landrum et al., 2023](#)) offers extensive functionality for cheminformatics and manipulating chemical structures. However, standard RDKit workflows are not designed for MLIP-driven simulation, while typical ASE-MLIP workflows may lack rich explicit chemical information such as bond orders or molecular identities, as well as capabilities for generating different conformations or searching substructures.

The molify package bridges this gap, providing an interface between RDKit's chemical structure generation and cheminformatics capabilities and ASE's handling of 3D atomic structures. Furthermore, molify integrates with PACKMOL ([Martínez et al., 2009](#)) to facilitate the creation of complex, periodic simulation cells with diverse chemical compositions, all while preserving crucial chemical connectivity information. In addition, molify simplifies the representation of molecular structures as graphs using NetworkX ([Hagberg et al., 2008](#)), e.g., enabling traversing or comparing them. Lastly, the combination of these packages enables selection and manipulation of atomistic structures based on chemical knowledge rather than manual index handling. While designed for MLIP data, the usage of molify is not limited and can be expanded, e.g., by utilizing the bond order information in other ASE-based workflows for classical MD simulations or integrating with machine-learning driven bond order predictions.

Statement of need

molify serves as a vital link between RDKit, ASE, NetworkX, and PACKMOL, designed to aid working with molecular structures in systems without dedicated topology files and formats. While its core function is to interface these tools, it thereby unlocks new capabilities and significantly reduces the manual coding and data wrangling typically required for preparing and analyzing molecular simulations. For example, ASE has no tools for handling topological information such as bonds or molecular identities, while RDKit cannot natively interface with MLIPs.

molify simplifies workflows that previously involved laborious tasks such as sourcing individual

structure files from various databases (e.g., the Materials Project (Jain et al., 2013) or the ZINC database (Tingle et al., 2023)) and custom setups of simulation cells. With molify, more complex and chemically diverse simulation cells are easier to set up and process.

One challenge in MLIP-driven simulations is the post-simulation identification and analysis of molecular fragments or chemical changes, as explicit topological information is not available and changes in connectivity can occur. molify addresses this by enabling the use of RDKit's powerful SMILES (Weininger, 1988)/SMARTS-based substructure searching on ASE structures. In addition, the resulting molecular graph can be exported to a NetworkX (Hagberg et al., 2008) object for further analysis. This selection and handling allows for similar functionality as is provided by the MDAnalysis (Gowers et al., 2016) atom selection language, designed for simulations with a fixed topology.

Features and Implementation

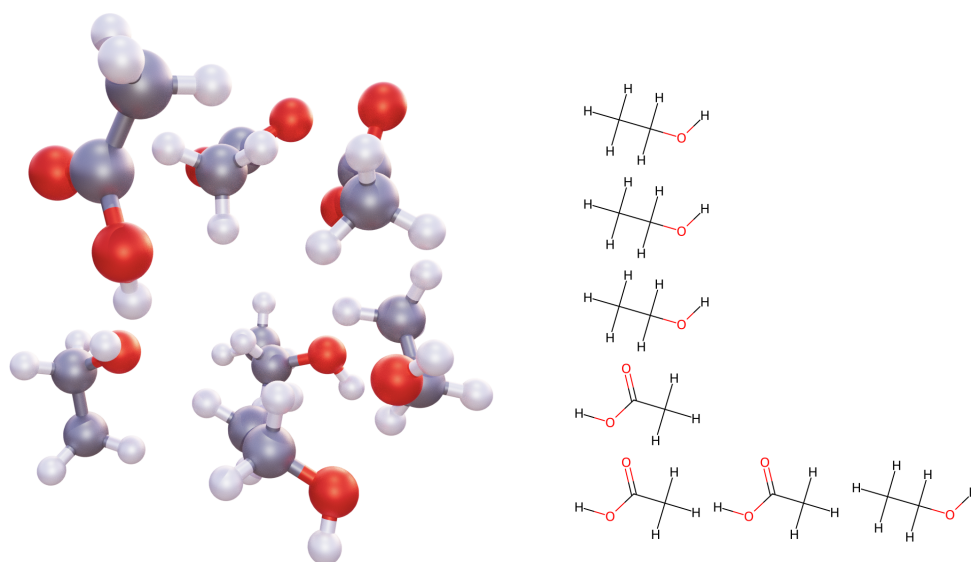


Figure 1: Visualization of a 3D structure from ASE, visualized with ZnDraw (Elijošius et al., 2024) (left) and its corresponding RDKit 2D chemical structure representation (right).

The generation of atomic configurations in molify is centered around SMILES for defining molecular species. A typical workflow often follows these steps:

1. Generating 3D conformers for individual molecular species from their SMILES using RDKit.
2. Packing these conformers into a simulation box with a target density using PACKMOL and returning the simulation box as an ASE Atoms object.
3. Running MLIP-based simulations using ASE.
4. Post-processing the simulation data by identifying and selecting structures based on SMARTS.

```
from molify import pack, smiles2conformers
from ase.optimize import LBFGS
from mace.calculators import mace_mp
```

```
water = smiles2conformers("O", numConfs=2)
print(water[0].info['connectivity'])
```

```
>>> [(0, 1, 1.0), (0, 2, 1.0)] # (atom_idx1, atom_idx2, bond_order)
ethanol = smiles2conformers("CCO", numConfs=5)
density = 1000 # kg/m^3
box = pack([water, ethanol], [7, 5], density)
print(box)
>>> Atoms(symbols='C10H44O12', pbc=True, cell=[8.4, 8.4, 8.4])
box.calc = mace_mp() # MLIP calculator
opt = LBFGS(box)
opt.run(fmax=0.01)
```

All ASE Atoms objects generated or processed by molify store connectivity information (bonds and their orders) within the ase.Atoms.info dictionary. If this information is available, molify uses it to convert between ASE, NetworkX and RDKit. If an ASE structure is converted to an RDKit molecule without pre-existing connectivity, molify leverages RDKit's bond perception algorithms (Kim & Kim, 2015) to estimate this information.

A visualisation of the 2D and 3D structure from the simulation is shown in Figure 1.

```
from molify import ase2rdkit
from rdkit.Chem import Draw
```

```
mol = ase2rdkit(box)
img = Draw.MolToImage(mol)
```

Additionally, molify can convert structures to and from NetworkX graphs, enabling graph-based algorithms such as traversal, isomorphism checks, or component analysis:

```
from molify import ase2networkx
```

```
graph = ase2networkx(box) # NetworkX graph with atomic numbers and bond orders
```

This bidirectional conversion capability allows the use of RDKit's chemical analysis tools together with ASE for MLIP-based simulations.

For example, after atomic positions in an ASE Atoms object are updated, molify can convert this structure back to an RDKit molecule allowing for further analysis or visualization. One common example is the extraction of substructures based on SMILES or SMARTS to track their structure and dynamics within a simulation. For example, molify streamlines the extraction of the CH₃ alkyl group from the ethanol molecules inside the simulation cell, without manual index lookup.

```
from molify import get_substructures

frames: list[ase.Atoms] = get_substructures(
    atoms=box,
    smiles="[C]([H])([H])[H]"
)
```

Related software

The functionality of molify relies critically on the following packages:

- **RDKit**: For cheminformatics tasks, SMILES parsing, conformer generation, and substructure searching.
- **ASE**: For representing and manipulating atomic structures, and interfacing with simulation engines.
- **PACKMOL**: For optimizing the placement of molecules within a specified simulation box size.
- **NetworkX**: For the handling and analysis of molecular graphs.

The molify package is currently a crucial part of the following software packages:

- [IPSuite](#): For generating structures for training MLIPs.
- [ZnDraw](#): Interactive generation of simulation boxes and selection of substructures through a graphical user interface inside a web-based visualization package.
- [mlipx](#): Creating initial structures for benchmarking different MLIPs on real-world test scenarios.

The OpenBabel ([O'Boyle et al., 2011](#)) package provides similar cheminformatics functionality to RDKit along with extensive file format support. However, OpenBabel is primarily designed as a format conversion tool with a focus on command-line usage and file I/O, while molify is designed for Python-native workflows with in-memory object conversions. Currently, OpenBabel does not provide direct support for ASE Atoms objects, ASE calculators (including MLIPs), or seamless RDKit-ASE interconversion within Python. Furthermore, molify's integration with PACKMOL and NetworkX provides capabilities beyond OpenBabel's core focus on format conversion.

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