AutoSteper

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Introduction

AutoSteper is a python package to simulate and analyze the stepwise reaction in chemistry. Currently, it well supports the exohedral fullerene systems $C_{2n}X_m(X=H,F,Cl,Br,OH,CF_3,CH_3). \label{eq:constraint} \mbox{Our lab is working intensively to cover more scenarios. Extensions and issues are greatly appreciated.}$

The stepwise reaction of exohedral fullerenes originated from the radical-addition theory, ¹ in which the functionalization of fullerenes is described to be a radical attack behavior. Functional groups attack the carbon cage one by one, and the energetically favored adducts could be isolated.

AutoSteper is designed to simulate this procedure in a fully automated fashion (denoted as growth simulation). It consists of 3 modules, namely, the generator, the optimizer, and the checker. As shown in Fig 1, the generator is designed to enumerate addition patterns and build quasi-equilibrium molecules. The optimizer is in charge of optimizing these molecules with 3rd party software. The checker is to ensure the optimized molecule maintains an intact structure.

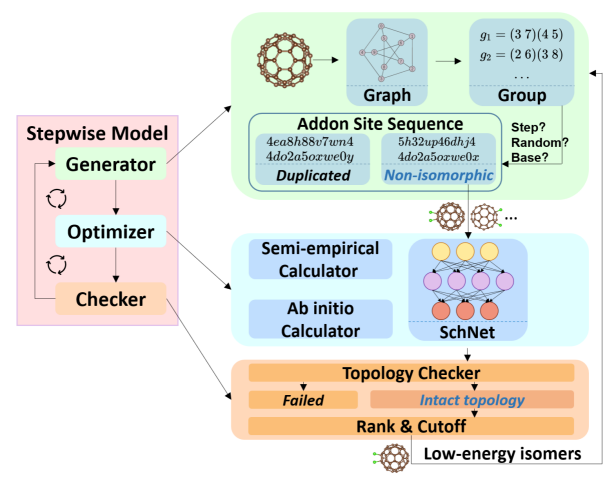


Fig 1. Overview of the stepwise model.

Despite the simulation of the stepwise addition procedure, AutoSteper provides automated topological analysis as well. Since the simulation generates valuable low-energy intermediates and the parent-son information is stored in every step, the topological linkage information could be easily extracted. Fig 2 well explains the topological linkage, that is, the relationship between $C_{2n}X_m \text{ and its parent } C_{2n}X_{m-2}, \text{ son } C_{2n}X_{m+2}.$

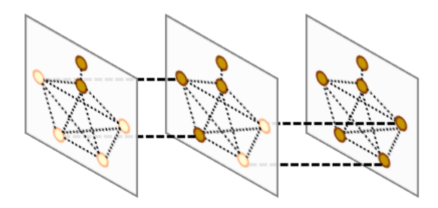


Fig 2. Illustration of the topological linkage.

This information is critical when one trying to parse pathways for a specific product, ² however, when it comes to different systems, the topological information needs to be re-generated. As an example, AutoSteper parses the Stone-Wales Rearrangement (SWR) phenomenon ³ by solving the subgraph isomorphism problem with the assistance of the networkx package. Fig 3 illustrates one of the SWR phenomena, in which a cage skeleton transformation takes place when the functional groups attack.

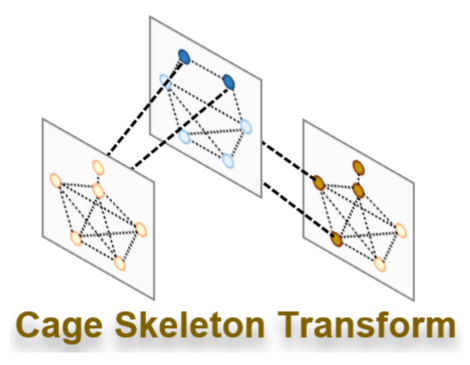


Fig 3. Illustration of an SWR scenario.

Since the subgraph isomorphism function is relatively slow, only limited isomers could be considered. Similarly, this function could also be used to re-generate pathways and draw the binding energy profile.

Installation, details of each module as well as the post-analysis functions are presented below.

Install

Enumeration

AutoSteper relies on OpenSource projects <u>FullereneDataPraser</u> and <u>usenauty</u> to properly enumerate non-isomorphic addition patterns.

<u>FullereneDataPraser</u> is an excellent python package to handle fullerene-related research problems, this project utilizes it to convert 3D coordinates to graph6str format. For install:

```
git clone https://github.com/XJTU-ICP/FullereneDataParser
cd FullereneDataParser
pip install .
```

<u>usenauty</u> is a lightweight tool to enumerate non-isomorphic addition patterns with <u>nauty</u> algorithm. The original project is in <u>usenauty</u>, here we employ a branch version of it. For install:

```
git clone https://github.com/Franklalalala/usenauty
cd usenauty
mkdir build
cd build
cmake .. -G "Unix Makefiles"
make
```

Note: The <u>CXX standard</u> is set to be **17**, which means the gcc version need to be 8 or higher, or a higher version of IDE, such as <u>Visual Studio 2017</u>. The cmake version need to be **3.1** or higher. There are two pre-compiled releases for Windows and Linux platform, respectively.

The absolute path of compiled cagesearch file corresponds to the gen_core_path button in generator module.

Main project

To install the main project:

```
1 | pip install AutoSteper
```

For other reliance:

```
pip install pytest-shutil, ase, numpy, pandas, networkx, tqdm, matplotlib, seaborn, dpdispatcher
```

To install from source code:

```
git clone https://github.com/Franklalalala/AutoSteper
cd AutoSteper
pip install .
pip install -r requirements.txt
```

AutoSteper

AutoSteper is the highest level of abstraction. It controls sub-modules to simulate the entire stepwise addition process. It has 3 different modes, corresponding to the 3 generator's modes. Details are presented below:

- step: the stepwise simulation.
- random: randomly generate isomers and optimize them to an equilibrium state.
- base: enumerate and evaluate all the non-isomorphic isomers for a specific addon number.

Except for these functions, AutoSteper provides engineering-related features, such as restart and error handling. For details see the engineering section.

step

The input script below is a typical case:

```
1
    import os
 2
 3
    from autosteper import AutoSteper
 4
 5
    mach_para = {
 6
        'batch_type': "x",
 7
        'context_type': "SSHContext",
 8
         'remote_root': 'xx/',
 9
        'remote_profile': {
10
            "hostname": "xx",
             "username": "xx",
11
            "password": "xx",
12
13
             "port": 22,
            "timeout": 10
14
15
        }
16
    }
17
18
    resrc_para = {
19
        'number_node': 6,
20
        'cpu_per_node': 6,
        'gpu_per_node': 0,
21
22
        'group_size': 10,
23
        'queue_name': "batch",
24
        'envs': {
             "OMP_STACKSIZE": "4G",
25
            "OMP_NUM_THREADS": "3,1",
26
27
             "OMP_MAX_ACTIVE_LEVELS": "1",
28
             "MKL_NUM_THREADS": "3"
29
        },
         'sub_batch_size': 50
30
31
    }
32
33
    para = {
34
         'pristine_path': r'geom.xyz',
35
         'root': r'./',
         'gen_para': {'group': 'Cl',
36
                      'geom_mode': 'pre_defined',
37
38
                      'gen_core_path': r'xx\cagesearch.exe'
39
                      },
         'opt_mode': 'xtb',
40
         'opt_para': {
41
42
             'cmd_list': [r'xx/xtb', '--opt', 'tight', '--json'],
43
             'out_list': ['xtbopt.xyz', 'xtbopt.log', 'xtbout.json'],
             'deal_wrong_mode': 'Report',
44
45
             'mach_para': mach_para,
             'resrc_para': resrc_para,
46
47
        },
48
         'run_para': {
```

```
49
             'start': 1,
50
             'stop': 4,
             'step': 1,
51
52
             'wht_list_para': {
53
                 'mode': 'rank',
54
                 'rank': 5
55
             }
56
         },
57
    }
58
59
    # raw folder is pre-defined
    auto = AutoSteper(para=para)
60
    auto.run()
61
```

The resrc_para and mach_para are designed to configure a suitable environment for optimizers. (see optimizer module) After that, one needs to configure the parameter of the class AutoSteper. Specifically, one needs to provide:

- pristine_path: the path to the pristine cage. It could be in any mainstream structure format, only if the <u>ASE</u> package supports it.
- root: the ABSOLUTE path to the growth simulation workbase, where the AutoSteper would make a directory in the name of the pristine cage. Fig 4 presents one of the workbase directory.

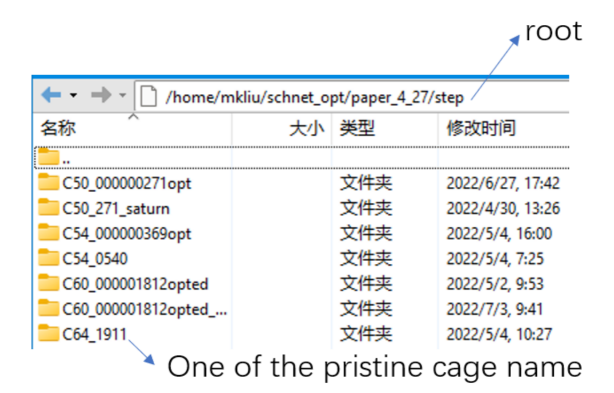


Fig 4. An AutoSteper workbase.

The gen_para and opt parameters configure a generator and an optimizer. To simulate a stepwise addition reaction, one needs to provide the run_para with the following parameters considered.

• start: the addition stage ($C_{2n}X_{start}$) when the simulation started. The first step will enumerate and evaluate all the non-isomorphic isomers without any filter.

- \bullet $\,$ stop: the addition stage $(C_{2n}X_{stop})$ when the simulation started.
- step: number of the newly attached functional groups after the first step.
- wht_list_para: parameters to control the isomers saved in every step. These isomers will serve as seeds in the next step to generate derivatives. The white list is a relatively concept to the blacklist. 7 modes could be selected, details see AutoSteper/test cutoff.py. The default mode is rank_and_value, in which the rank is 200, and the value is 1eV. rank_and_value means the lower boundary of two of them.

AutoSteper would create sub-workbases for every addon number. Fig 5 presents one of the scenarios. In this case, the start value is 1, the stop value is 10, and the step value is 1.

← ▼ → ▼								
名称 ^	大小	类型	修改时间					
<u> </u>								
addons 1addons		文件夹	2022/6/27, 17:30					
2addons		文件夹	2022/6/27, 17:30					
3addons		文件夹	2022/6/27, 17:31					
4addons		文件夹	2022/6/27, 17:34					
=== 5addons		文件夹	2022/6/27, 17:36					
addons 6addons		文件夹	2022/6/27, 17:38					
7addons		文件夹	2022/6/27, 17:39					
addons 8addons		文件夹	2022/6/27, 17:41					
9addons		文件夹	2022/6/27, 17:42					
10addons		文件夹	2022/6/27, 17:43					

Fig 5. An AutoSteper sub-workbase.

The directory of the **first** step is illustrated in Fig 6.

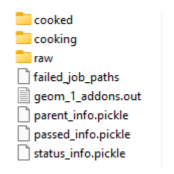


Fig 6. The first step workbase.

The functions of each file/directory are presented below.

- raw: the quasi-equilibrium isomers built in every step.
- cooked: the equilibrium state of each isomer, in xyz format.
- cooking: the real workbases for each optimization job. It typically contains more optimization details than the cooked folder.
- failed_job_paths: the absolute path of each failed optimization job as well as their corresponding failed status code.
- geom_1_addons.out: the enumerated addition patterns in the first step. geom is the name of the pristine cage, 1 is the addon number of the first step.

- passed_info.pickle: key information of the optimization jobs that **passed** the topological check. In the early version of AutoSteper, this file is called deep_yes_info.pickle, meaning information is stored in a deep chart. Fig 7 presents one of the scenarios. The meaning of each column is presented below:
 - o name: the name for each isomer, in 36 format.
 - energy: the equilibrium energy of each isomer, in units eV.
 - xyz_path: the absolute path to each isomer structure, in xyz format.
 - nimages: the number of images in each optimization trajectory.

	name	energy	xyz_path	nimages
0	4do2b19fp9mo	-3743.81058	F:\\AutoSteper_new\\AutoSteper\\tests\\test_ste	16
1	4do3ua4l8kqo	-3743.68896	F:\\AutoSteper_new\\AutoSteper\\tests\\test_ste	8
2	4do2a5oxwetc	-3743.28444	$F: \AutoSteper_new \AutoSteper \tests \test_ste$	9
3	6ki3f8jeul1c	-3743.16969	F:\\AutoSteper_new\\AutoSteper\\tests\\test_ste	9
4	4do2a5xtje9s	-3743.09186	F:\\AutoSteper_new\\AutoSteper\\tests\\test_ste	9

Fig 7. Example of the passed_info.

- parent_info.pickle: key information of the **parent-son** relationships generated during the growth simulation. In the early version of AutoSteper, this file is called flat_yes_info.pickle, meaning information is stored in a flat chart, and only the passed isomers are considered. The flat format enables a fast index when parsing the topological information.
 - \circ The first step is different from others since there is only one parent for all the $C_{2n}X_{start}$ isomers. Fig 8 presents one of the cases. The columns correspond to each $C_{2n}X_{start}$ isomer. The first row corresponds to their energy.
 - o Fig 9 presents a case in the proceeding addition stages. The columns correspond to each $C_{2n}X_m$, m>start isomer. The first row stores the names of their parent(s). Note that, isomers in $C_{2n}X_m$ addition stage could have more than one parent $C_{2n}X_{m-step}$. The second row corresponds to their energy.

	00000000000x	0000000000010	000000000014	00000000001c	0000000005m9t	0000000005m9u
0	-146.07188	-146.06557	-146.03697	-146.05130	-146.04154	-146.05752

Fig 8. Example of the parent_info in the first step.

	6ki3f8sahla8	4eweoassjg1s	6ki3f8lmrc3k	4do2a5q1uscg	6mys7ir44ni8	
0	['6ki3f8jeul1c', '4do2a5xtje9s']	['4do2a5oxwetc']	['6ki3f8jeul1c']	['4do2a5oxwetc']	['6ki3f8jeul1c']	
1	-3866.689756854536	-3866.682364908761	-3866.670989029402	-3866.666733393176	-3866.6460294948247	

Fig 9. Example of the parent_info in the proceeding addition stages.

- status_info.pickle: the status code for each optimization job, in flat chart format for indexing convenience. Note that this status code is different from the failed status code. Only three codes are available:
 - o 0: normal termination.
 - -1: topological intactness is undermined. (failed topology check)
 - -2: wrong jobs. This would happen when there are no files retrieved from computational resources, for example, the internet is broken.

	4do2a5q1uscq	4do2a5tdpwxs	4do2a5xtihfk	4do2a5xtiklc	4do2a5xtigww	4do2a5xtm7eo	4do2a5xtp0ik
0	0	0	0	0	0	0	0

Fig 10. Example of the status_info.

The directory of the **proceeding addition stages** is illustrated in Fig 11. The difference compared with the first step is presented below:

• sub_nauty: there is more than one parent that generates derivatives. Related information is dumped in this folder.

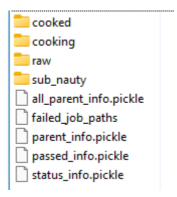


Fig 11. The workbase for the proceeding addition stages.

• all_parent_info.pickle: the parent-son information for all the $C_{2n}X_m$ isomers. (see Fig 12.) This is generated when building the quasi-equilibrium isomers. Note that the parent_info.pickle only considers the passed ones, and it contains energy info. The all_parent_info.pickle stores duplicated but more detailed information, therefore it may be useful for future development.



Fig 12. Example of the all_parent_info.

random

The random mode could be used to sample targeted configuration space, for example, building a dataset to train Neural Network Potential (NNP). The parameters for random mode are basically the same as the step mode. Differences lie in the run_para, which is replaced by random_para. Specifically, one needs to provide:

- addon_list: a list that consists of desired addon numbers, e.g, $C_{2n}X_m$, m in addon_list.
- ullet random_num: for each addon number m, the number of randomly sampled isomers $C_{2n}X_m$.
- try_times: since some systems are highly unstable, e.g, $C_{2n}X_m$, m=2n, all the isomers sampled could be unphysical and fail the topological check. In this case, the whole batch of isomers $C_{2n}X_m$ should be discarded. This parameter is highly recommended to control the failed chances. Note that, it needs <code>deal_wrong_mode</code> set as <code>Tough</code> to properly function.

Despite these parameters, the execution method of AutoSteper changed from run to random. For an example script, see AutoSteper/test random.py.

base

The base mode could be used to enumerate and evaluate all isomers for a specific system $C_{2n}X_m$. In fact, it could be viewed as the first step in the step mode. Since the base mode has only one step, its input script doesn't need <code>run_para</code>. The rest of the parameters stay the same as above. The execution method of AutoSteper changed from <code>run</code> to <code>base</code>. For an example script, see <code>AutoSteper/test_base.py</code>.

Engineering

Currently, AutoSteper provides restart and error_handling for engineering convenience. More features are under development.

restart

The restart feature is designed for the step mode in case the simulation is interrupted. To use it, simply replace the execution method of AutoSteper to restart. For example:

```
1  # auto.run()
2  auto.restart(restart_add_num=5)
```

Note that, the <code>restart</code> method will delete the original workbase for $C_{2n}X_m, m >= restart_add_num, \text{ after that, a new workbase will be created for } \\ C_{2n}X_m, m = restart_add_num. \text{ Make sure the } \\ \text{restart_add_num} \text{ equals the exact addon stage when the simulation was interrupted.}$

Besides, this feature could be used to **proceed** with a normally terminated simulation. For example, the original one terminated in add_num = 4, and the restart_add_num could be set as 4+step.

error handling

The error_handling feature is assigned to optimizers. For description convenience, details of them are presented in this section. Note that, the error mentioned here denotes an unexpected optimization task result, which is different from the failed notation.

There are 3 modes in total. Specifically:

- Report: simply report the wrong information and end out.
- Complete: recursively submit jobs in small batches to minimize the wrong jobs, then end out. Note that, the dpdispatcher submits jobs in a batch style. When there is one job ends unexpectedly, there would be no retrieval from the remote for the whole batch. The Complete mode will submit recursively with small batches until the abnormal ones are left.
- Tough: designed for random mode in case the whole batch of randomly generated isomers are unphysical.

Optimizer

The optimizer module heavily rely on open-source package <u>deepmodeling/dpdispatcher</u>. See <u>Getting Started</u> to get familiar with dpdispatcher. Here presents the usage of AutoSteper's customized version.

machine and resource

To start with, one needs to set a machine and a resource configuration. Here present some examples.

For the machine parameter, two sets of configurations are recommended. See below:

```
1 # from local (typically your win system) to clusters. Input scripts are submitted in Personal Computer (PC).
```

```
2
    mach_para = {
 3
        'batch_type': "Torque", # my cluster type
 4
        'context_type': "SSHContext",
 5
        'remote_root': '/home/test/xx/', # the remote workbase where the actual
    computation take place.
 6
        'remote_profile': {
 7
            "hostname": "2xx.2xx.xx.7x", # IP
            "username": "xx",
 8
 9
            "password": "xx",
10
            "port": 22,
            "timeout": 10
11
12
        }
    }
13
14
15
    # inside your clusters. Input scripts are submitted in the cluster.
16
    mach_para = {
        'batch_type': "Torque", # my cluster type
17
        'context_type': "LocalContext", # Do not need IP information
18
19
        'remote_root': '/home/test/xx/',
        'remote_profile': {
20
21
            None
22
        }
23
    }
```

For the resource parameter, here is an example:

```
1
    resrc_para = {
        'number_node': 6, # the sequence name for your cluster.
 2
        'cpu_per_node': 6, # computational resources for each task.
 3
 4
        'gpu_per_node': 0, # same as above
 5
        'group_size': 10, # number of tasks contained in each job (group).
        'queue_name': "batch", # queue name for my cluster
 6
                                # extra enviromental variables
 7
        'envs': {
 8
            "OMP_STACKSIZE": "4G",
 9
            "OMP_NUM_THREADS": "3,1",
10
            "OMP_MAX_ACTIVE_LEVELS": "1",
            "MKL_NUM_THREADS": "3"
11
12
        },
13
        'sub_batch_size': 50 # number of tasks contained in each batch.
14
    }
```

The machine parameters tell the dpdispatcher **which cluster** to use and **how to contact**, while the resource parameter **assigns** computation resources to each job.

The original workflow of the dpdispatcher is illustrated in Fig 13.

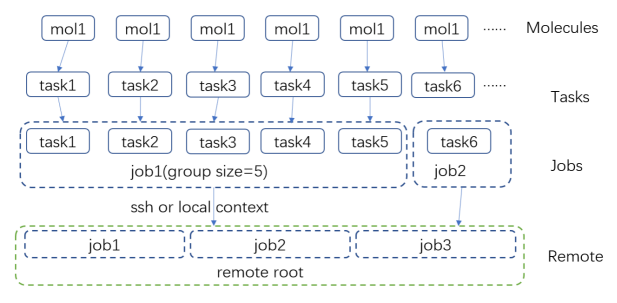


Fig 13. Simplified workflow of dpdispatcher.

Each optimization job corresponds to a task. Then, tasks are grouped (group_size) into jobs. These jobs are submitted through ssh or local context (context_type) to remote (remote_root), where the remote would assign computational resources to each job (All_cpu_cores /cpu_per_node) and execute them in parallel.

However, when it comes to huge task sequences, the number of groups in line may put pressure on the cluster. And when something wrong happened in a single job, the whole batch would be undermined. (For example, no retrieval from remote.) Therefore, we proposed the sub_batch_size parameter to perform job dispatch in a mini-batch style. An illustration of the modified dpdispatcher is presented in Fig 14.

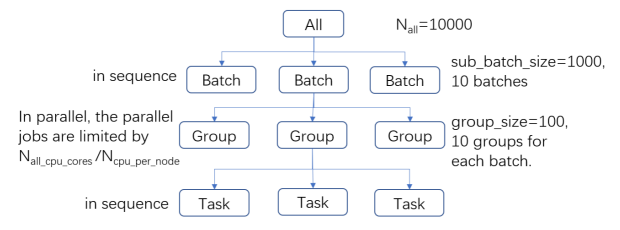


Fig 14. A top-down illustration of the modified dpdispatcher.

More details could be found in the documentation of <u>Machine parameters</u> and <u>Resources</u> <u>parameters</u>.

opt mode and parameter

Currently, AutoSteper provides interfaces for 3 software, namely, the <u>xTB</u> program, the <u>Gaussian</u> software, and the <u>ASE</u> python library. Examples could be found in <u>test ASE Optimizer</u>, <u>test Gaussian Optimizer</u>, and <u>test XTB Optimizer</u>.

In addition, AutoSteper provides the Multi_Optimizer to properly integrate different software or employ the same software repeatly. Examples could be found in test multi_optimizer.

The details about related parameters are presented below:

- 1. opt_mode: tells the AutoSteper class or the switch_optimizers function which optimizer to choose.
- 2. cmd_list: the actual command line in the final workbase (without the filename). It consists of the call of the program, options, flags, and so on.
- 3. out_list: the names of output files that need to be downloaded.
- 4. deal_wrong_mode: how to deal with wrong jobs, details see the engineering section.

Generator

The generator module is in charge of building molecules. Details of parameters are presented below:

- group: the name of functional groups. Currently, AutoSteper supports $C_{2n}X_m(X=H,F,Cl,Br,OH,CF_3,CH_3)$.
- gen_core_path: the absolute path to the executable binary file cagesearch, which is compiled from the Franklalalala/usenauty repository.
- geom_mode: decides how to build quasi-equilibrium isomers. This parameter is highly recommended to be set as pre_defined. The pre-defined geometry parameters are chosen from thousands of randomly sampled isomers. If one needs to change these parameters, set geom_mode to another value and assign new parameters through geom_para. Note that, the new format needs to stay consistent with the original.

Note that, the generator module could be used alone to build hand-tuned structures. See <u>AutoSteper/test build unit.py</u>.

Checker

The checker module will check optimized isomers to ensure an undermined topology. 7 scenarios could be detected, their corresponding failed status codes are presented below.

- 1. At least one functional group breaks the bond with the cage and becomes a radical.
- 2. At least one functional group deviates from the initial addition site and moves to another.
- 3. At least one 3-membered carbon ring is formed during optimization, meaning the pristine cage is squeezed by functional groups.
- 4. At least one carbon atom only has 2 neighboring carbon atoms or less, meaning the cage is broken.
- 5. At least one functional group binds with 2 or more carbon atoms, which is unstable for currently supported functional groups.
- 6. At least one carbon atom binds with 5 or more atoms, which means a small cluster or a coordination is formed.
- 7. The inner intactness of at least one functional group (OH, CF_3, CH_3) is undermined.

These status codes will be reported in the failed_job_paths file.

Need to mention that, the AutoSteper module doesn't need any specific input parameters for the checker module, though it could also be used alone. See <u>AutoSteper/test_checker.py</u>.

Blacklist

The concept of the blacklist is based on the assumption that high-energy isomers probably contain local instability motifs, therefore their derivatives will unlikely to become stable ones since they still contain those instability motifs. This is a dual concept to the low-energy configuration space, which is treated as seeds to generate derivatives. See Fig 15.

AutoSteper collects two kinds of isomers into the blacklist.

- 1. The isomers that failed the topological check. (denote as failed)
- 2. The high-energy isomers within certain reverse cutoff. (denote as unstable)

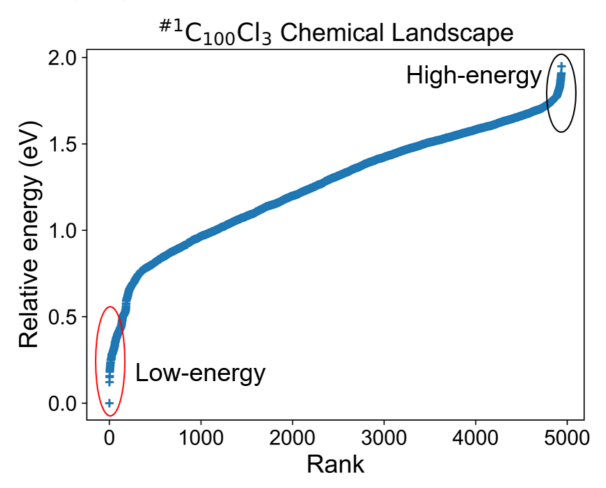


Fig 15. Illustration of the high-energy configuration space.

When it comes to a new step, the new addition patterns will check through the blacklist at first. If a pattern contains any of the recorded patterns, it will be directly skipped.

To control the influence of a high-energy pattern, AutoSteper provides a queue to store high-energy patterns. See Fig 16.

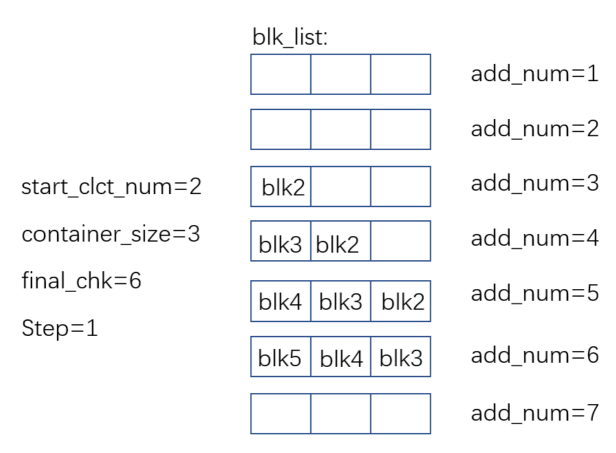


Fig 16. Illustration of the queue maintained by AutoSteper.

AutoSteper starts collecting high-energy isomers in start_clct_num. These patterns start functioning in the next step and will continue to function till start_clct_num+container_size*step. The blacklist system will shut down after final_chk.

To enable the blacklist feature, one needs to provide a <code>blk_para</code>. Here is an example of <code>blk_para</code>, for example of an input script, see AutoSteper/test-blk-list.py

```
b1k_para = {
1
2
         'start_clct_num': 2,
3
         'final_chk_num': 8,
4
         'clct_unstb': True,
5
         'unstb_para': {
             'mode': 'rank',
6
7
             'rank': 10,
8
        },
9
         'container_size': 3
10
    }
```

Note that, all failed addition patterns are collected by default as long as the blacklist system functions. Another kind of high-energy isomers is collected when the clct_unstb is Ture. The unstb_para controls the reversed cutoff range, details see AutoSteper/test cutoff.py.

Pre-scan

The pre-scan feature takes the quasi-equilibrium geometry to approximate the equilibrium state isomer. Since AutoSteper builds quasi-equilibrium isomers in a python environment, currently only the python package <u>ASE</u> is supported as the single-point evaluator.

The generated isomer (in atom class) would go through a single-point evaluation before dumping to a xyz format file. After the generation of all isomers, the low-energy ones will be selected and re-dumped into the post_pre_scan_raw folder. These isomers would undergo geometry optimization with optimizers. Fig 17 presents a working folder when the pre-scan feature is enabled. It's basically the same as the step mode workbase.

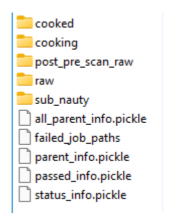


Fig 17. The workbase when the pre-scan feature enabled.

To enable a pre-scan feature, one needs to provide a pre_scan_para. Here is an example of pre_scan_para, for example of an input script, see AutoSteper/test_pre_scan.py.

```
pre_scan_para = {
1
2
       'start_ps_para': 2, # when the pre-scan feature enabled
3
       'final_ps_para': 3, # when the last addition stage that the pre-scan
   feature functions
       'calculator': calculator, # the calculator in ASE format
4
5
       'ps_cut_para': {  # to control the cutoff range
6
           'mode': 'rank',
7
          'rank': 80
8
       }
9
  }
```

Note:

- 1. The calculator needs to stay in ASE format.
- 2. The ps_cut_para controls the cutoff range for the isomers that need geometry optimization.

Post-analysis

AutoSteper provides multiple functions and abstractions to perform the post-analysis. Here presents a brief introduction.

refine

When one needs to improve computational accuracy, the refine function in the parser module presents a nice solution. Only 3 parameters are needed to perform a refinement procedure. That is:

- old_workbase: the original workbase.
- new_workbase: the new workbase.
- ref_para: the same format as the optimizer's parameter to configure an optimizer.

That's it. AutoSteper will refine the original data and dump them into the new workbase. For details, see AutoSteper/test refine.py.

isomorphism test

AutoSteper provides 3 functions to perform the isomorphism test. For example, see <u>AutoSteper/test_iso_relatives.py</u>.

Details are presented below:

simple_test_iso

This function is designed to test whether a specific isomer $^{\#M}C_{2n}X_m$ is within the simulation results $^{\#M}C_{2n}X_{m}$ -q, $0 < q <= max_rank$. If it's indeed within the results, this function will output its corresponding rank q. One needs to provide:

- q_atoms: the queried isomer, in <u>ASE Atoms format</u>.
- passed_info_path: the absolute path to the queried passed_info.pickle.
- top_k: a cutoff performed on the passed_info.pickle, rank mode only. If none, AutoSteper will scan all the simulation results.

simple_log_relatives

This function is designed to quickly find relatives of a specific isomer $^{\#M}C_{2n}X_m$ and log key information to a writeable path. Here relatives mean the intermediates ($^{\#M}C_{2n}X_q,q< m$), isomer ($^{\#M}C_{2n}X_q,q=m$) and derivatives ($^{\#M}C_{2n}X_q,q>m$) of the queried isomer. To ensure a fast test, here use the addition patterns as a criterion.

Two ways to decide the queried addition pattern.

- The recommended way to get the addition pattern:
 - q_atoms: the queried isomer, in <u>ASE Atoms format</u>. Note that, it needs to have an identical pristine cage to the target. This ensures an identical sequence.
 - o group: the symbol of the functional group.
 - o cage_size: the size of the pristine cage.
- The second way to get the addition pattern:
 - o q_seq: the 36-base format name. See the 36 base function.
 - q_cage: the key to decipher the 36-base name to a sequence, in <u>AutoSteper/cage</u> format.

After that, one needs to provide:

- fst_add_num: the smallest addon number to be scanned.
- final_add_num: the biggest addon number to be scanned.
- step: the step that used in the growth simulation.
- workbase: the original workbase.
- dump_log_path: the absolute path to dump the related information.

Here is an example of the dumped log.

strict_scatter_relatives

This function is designed to strictly find relatives of a specific isomer. It implements the <u>subgraph is isomorphic</u> function to perform the isomorphism test and dump information in a png format (see Fig 18). The input parameters are basically the same as the above function. The difference is that it needs a folder to dump information.

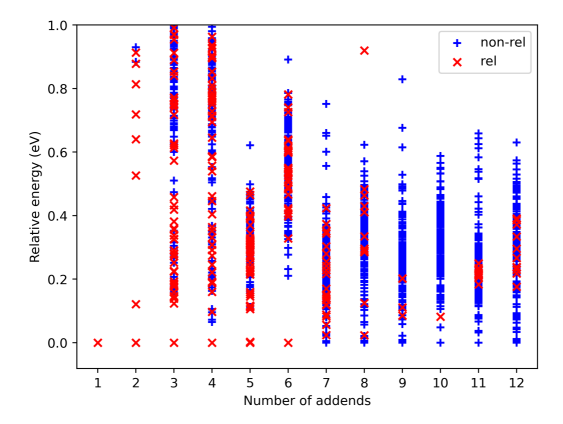


Fig 18. Example of the dumped information. The red 'x' presents a relative, blue '+' is a non-isomerphic one.

In addition, AutoSteper dumps the relative energy of each scanned isomer, and groups them into the non_rel_e and rel_e.

cook_disordered

To better analyze the low-energy configuration space, AutoSteper provides the cook_disordered function to translate disordered log files into structured information. Two folders and one pickle file will be generated. Details are presented below:

- pathway_info.pickle: information for all the pathways. name column meaning the pathway-related names, rel_e corresponds to the relative energy of isomers, e_area means the relative energy of pathways.
- ./sorted : sorted information for isomer rankings. And this is a standard output format.
 - ./sorted/log: the final optimization logs for a specific isomer. File names contain two
 metrics. The first number means the number of addends, the last number means the
 ranking of the specific isomer. For example, l_addons_1 means it contains 1 addend
 and its energy rank is 1 (the lowest energy one).
 - ./sorted/xyz: the final image of the optimization trajectory. The name convention is the same as above.

- ./sorted/info: energy information.
- ./sorted/connection: connection information. 1_addons_1.npy corresponds to the isomer, whose geometry information is stored in 1_addons_1.xyz. This isomer has connection relationships with higher addends, here in this case, it means 2 addends.
 1_addons_1.npy stores this information, 1 meaning connected, 0 for not.
- ./pathways: pathway-related isomers. traj.log for the whole trajectory.

One needs to provide following parameters:

- disordered_root: path to the disordered root.
- dump_root: path to dump information.
- keep_top_k_pathway: how many generated pathways to keep. (in rank mode)
- step: the step of the original growth simulation.
- log_mode: two log formats are supported. 1. the gaussian format, type key word gauss. 2. the xyz format, type key word xyz.

For an example, see AutoSteper/test cook disordered.py.

On the bases of structured information, one can easily perform topological analysis. See find_swR and get_binding_e in below.

find_SWR

For description convenience, here we denote an isomer before and after an SWR as $^{\#1}C_{2n}Cl_{2m}$ and $^{\#2}C_{2n}Cl_{2(m+1)}$. The $^{\#1}C_{2n}$ system is represented as the query system and $^{\#2}C_{2n}$ is the target system.

One needs to provide the following parameters:

- q_sorted_root: the sorted information provided by the cook_disordered function. This root corresponds to the query system.
- tgt_sorted_root: same as above, for the target system.
- swr_dump_path: where to dump the information.
- step: the step of the original growth simulation.
- is_unique: if true, for every atoms in q_root, only one SWR target is outputted, typically for the lowest energy isomer, here we take the rank info in the name as criteria.
- is_low_e: if true, for every atoms in q_root, only one SWR target is outputted, and it should have lower energy than the 'ought to be' parents.

Details about the output are presented below:

./swr_dump_path: Automated generated SWR pairs.

- ullet For example, [q_14_to_tgt_15] means SWRs between $^{\#14}C_{2n}Cl_{2m}$ and $^{\#15}C_{2n}Cl_{2(m+1)}$
- [q_14_to_tgt_15/0_to_2_swr_1] means SWRs between $^{\#14}C_{2n}Cl_0$ and $^{\#15}C_{2n}Cl_2$. $^{\#14}C_{2n}Cl_0$ means the pristine cage.
- The 1 in 0_to_2_swr_1 means this query atoms has an energy rank of 1 among other query atoms.
- In the folder of <code>q_14_to_tgt_15/0_to_2_swr_1</code>, <code>q_atoms.xyz</code> is the geometry structure for the query atoms $^{\#14}C_{2n}Cl_0$, <code>tgt_atoms_rank_1.xyz</code> is the corresponding target atoms $^{\#15}C_{2n}Cl_2$, since there may be more than one target atoms (if the <code>is_unique</code> option is <code>False</code>), the <code>1</code> in <code>tgt_atoms_rank_1.xyz</code> means the energy rank of target atom.

For an example, see <u>AutoSteper/test_find_SWR</u>.

get_binding_e

The binding energy well explains the reaction activity. Based on the structured topological information provided by the <code>cook_disordered</code> function, one can easily parse the binding energy information. Set hydrofullerene as an example, AutoSteper following this equation to calculate binding energy.

$$E_{binding} = E_{C_{66}H_{2m}} - E_{C_{66}H_{2(m-1)}} - E_{H_2} \tag{1}$$

One needs provide the following parameters:

- sorted_root: the structured source folder.
- cage_e: the energy of the pristine cage.
- addends_e: the energy of the simple substance of addons. Here in this case, its Hydrogen.

Note that, the cage_e and addends_e need to be calculated under the same computational level as the general isomers.

The output of this function is dumped into the <code>sorted_root/info/</code>, in the format of <code>pickle</code> and <code>xlsx</code>.

For an example, see AutoSteper/test binding e.py.

Path_Parser

The Path_Parser is an abstraction designed to get pathways from the parent-son information, which is generated along with the **Growth Simulation**. This feature is considered since the very beginning of AutoSteper. The parent-son information could be viewed as a by-product of the simulation, therefore the calculation of pathways is extremely fast.

One needs to provide the following parameters for a basic configuration:

```
1
   path_para = {
2
        'step': 1, # the step that used in the growth simulation.
        'start': 1, # when the growth simulation started
3
4
        'q_add_num': 4,
5
        'q_path_rank': 10,
6
        'q_isomer_rank': 5,
7
        'log_low_e_num': 10,
8
  }
```

The configuration will generate pathways for $C_{2n}X_{m}$ i. Here the <code>q_add_num</code> is the m, the <code>q_isomer_rank</code> is i. <code>log_low_e_num</code> will dump low-energy isomers $C_{2n}X_{m}$ i, $i < log_low_e_num$ into a log. <code>q_path_rank</code> will decide how many low-energy pathways to be dumped.

This configuration parses pathways for the specific <code>q_isomer_rank</code> . All the pathways end to the $C_{2n}X_{m_i}, i=q_isomer_rank$.

If one needs a higher accuracy of pathways, there is a refiner option in Path_Parser. See AutoSteper/test path path parser 2 ref.py.

If one needs to control the number of pathways, set a ctl_path_para as below:

```
path_para = {
 2
        'step': 1, # the step that used in the growth simulation.
 3
        'start': 1, # when the growth simulation started
 4
        'q_add_num': 22,
 5
        'q_path_rank': 10,
        'q_isomer_rank': 5,
 6
 7
        'log_low_e_num': 10,
 8
        'ctl_path_para': {
             'ctl_parent_num': 3, # Control the number of parents for each
    isomer.
             'max_path_num': 10000, # Control the maximum number of pathways.
10
        }
11
12
    }
```

This is for the case when the queried addon number is very high. Since the Path_Parser functions in a DFS way, the low-energy pathways will be well preserved.

The generated pathways are highly structured and informative, see <u>AutoSteper/test_path_parser</u>. Here presents the well-designed heatmap. See Fig 19.

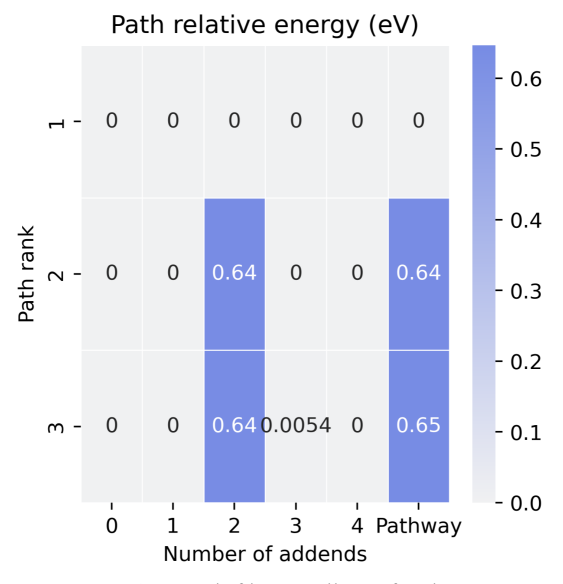


Fig 19. Example of the generated heatmap for pathways.

It well-explained the ranking of pathways. For example, all the related isomers for the pathway in row 1 is the lowest-energy one (relative energy is 0), therefore it's the lowest-energy pathway.

Reference

In DOI format.

- 1. 10.1039/A905719F 🔁
- 2. 10.1038/nchem.549 🔁
- 3. 10.1021/acs.accounts.9b00175 🔁