CrySPY: a crystal structure prediction tool accelerated by machine learning

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ABSTRACT

We have developed an open-source software called CrySPY, which is a crystal structure prediction tool written in Python 3, and runs on Unix/Linux platforms. CrySPY enables anyone to easily perform crystal structure prediction simulations for materials discovery and design, and automates structure generation, structure optimization, energy evaluation, and efficiently selecting candidates using machine learning. Several searching algorithms are available such as random search, evolutionary algorithm, Bayesian optimization, and Look Ahead based on Quadratic Approximation. Machine learning is employed to efficiently select candidates for priority optimization. CrySPY does not require complex machine learning techniques for users. In the latest version of CrySPY, both atomic and molecular random structures can be generated. CrySPY supports VASP, QUANTUM ESPRESSO, OpenMX, soiap, and LAMMPS for local structure optimization and energy evaluation. CrySPY is distributed under the MIT license at https://github.com/Tomoki-YAMASHITA/CrySPY. Documentation of CrySPY is also available at https://Tomoki-YAMASHITA.github.io/CrySPY_doc.

KEYWORDS

Crystal structure prediction; Bayesian optimization; LAQA; evolutionary algorithm; first-principles calculations; machine learning; materials informatics

CLASSIFICATION

Materials data analysis

1. Introduction

Research on data-driven materials development based on computer simulations has been actively conducted for the last decades. With the development of computational capabilities, it has become possible to obtain a large amount of reliable data from first-principles calculations at high speed. However, first-principles calculations cannot be directly applied to the design of new materials for new compositions and unknown structures, because such
Quantum-mechanical approaches require their crystal structures as input. In recent years, crystal structure prediction (CSP) methods have made it possible to discover new materials in conjunction with first-principles calculations. The history of new material discovery by CSP is well summarized in the review paper by the pioneers in this field [1]. So far, a great deal of effort has been devoted to developing the searching algorithms, such as random search (RS) [2–5], simulated annealing [6,7], minima hopping [8,9], evolutionary algorithm (EA) [10–13], and particle swarm optimization (PSO) [14,15]. In particular, EA and PSO are widely used and efficient algorithms as implemented in USPEX [11–13] and CALYPSO [14,15], respectively.

Although significant progress in searching methodologies has been made over the years, CSP is quite a time-consuming task. In CSP, first-principles methods are mostly used for the structure relaxation and energy evaluation to find the global energy minimum. However, first-principles calculations are quite heavy for large systems. The development of more efficient searching algorithms is highly desired to reduce computational cost. In our previous study, we proposed two selection-type algorithms, Bayesian optimization (BO) [16,17] and Look Ahead based on Quadratic Approximation (LAQA) [18]. In the conventional methods, all generated structures are locally optimized in order, while the selection-type algorithms employ machine learning to efficiently select candidates for priority optimization. These machine learning approaches make it possible to find the most stable structure among a large number of structures with a small number of trials. Besides, it is worth mentioning that researches on the acceleration of local structure optimization have also been conducted using machine learning potential [19–23].

We have developed an open-source software called CrySPY, which enables anyone to easily perform CSP simulations for materials discovery and design. CrySPY automates structure generation, structure optimization and energy evaluation by external programs, and efficiently selecting candidates using machine learning. While some codes for CSP such as AIRSS [2–5], USPEX [11–13], CALYPSO [14,15], GAS [24], and XtalOpt [25] are released, CrySPY is unique in its feature of selecting candidates using machine learning. In Section 2, we provide an overview of the software. In Section 3, we elaborate on each searching algorithm. Some features of CrySPY are introduced in Section 4. Finally, we summarize our software in Section 5.

2. Overview

2.1. System requirement

CrySPY is written in the Python 3 programming language, and runs on Unix/Linux platforms including macOS. CrySPY requires the following external Python libraries:

- COMBO
- pymatgen
- PyXtal

COMBO [26] is an efficient Bayesian optimization library, pymatgen [27] has many features in materials analysis, and PyXtal [28] is useful for crystal structure generation and symmetry analysis. First-principles or classical interatomic potential codes are also needed for energy evaluation and local structure optimization. CrySPY is currently interfaced with the following program packages for such purposes:

- VASP
- QUANTUM ESPRESSO
- OpenMX
- soiap
- LAMMPS

VASP [29], QUANTUM ESPRESSO [30], and OpenMX [31] are first-principles program packages, and soiap [32] and LAMMPS [33] are classical molecular dynamics packages.

2.2. Simple execution

CrySPY is simple to run. CrySPY itself is available at https://github.com/Tomoki-YAMASHITA/CrySPY. Users only need to provide the input file of CrySPY, cryspy.in, and input files required for the structure optimizer. An example of cryspy.in is shown below:

```
[basic]
algo = RS
calc_code = VASP
tot_struc = 10
nstage = 2
njob = 5
jobcmd = qsub
jobfile = job_cryspy

[structure]
atot = 8
atype = Si
nat = 8

[VASP]
kppvol = 80 100
```

If VASP code is used, for example, INCAR and POTCAR files are needed as input files. Structure data and k-point input files such as POSCAR and KPOINTS are automatically generated by CrySPY with pymatgen. A job script to submit structure optimization jobs according to the system is also needed. Once the input files are prepared, the CSP simulation can be easily started by executing the following command:

```
$ python3 /path/to/cryspy.py
```
2.3. Job automation

Structure optimization jobs are managed automatically by CrySPY. The number of the jobs to be submitted at a time is controlled by the input variable, njob, in cryspy.in. We adopt a stage-based system for structure optimization calculations. As an example of this, users can configure the following settings. In the first stage, only the ionic positions are relaxed, fixing the cell shape. Next, the ionic positions and cell shape are fully relaxed in the second stage. When a job is finished, the optimized structure data and energy can be automatically collected and a new job can be submitted.

![Diagram](image_url)

**Figure 1.** Example of CSP result by BO. Energies of optimized $Y_2Co_{17}$ structures are shown. Each selection by BO is color-coded.

2.4. Analysis and visualization

CSP simulation data are serialized and saved using the pickle module of Python. The data can be easily analyzed using Python codes, and jupyter notebook templates for this purpose are available at [https://github.com/Tomoki-YAMASHITA/CrySPY_utility](https://github.com/Tomoki-YAMASHITA/CrySPY_utility). Figure 1 is an example of CSP results drawn using the template. The energies of optimized structures of $Y_2Co_{17}$ obtained by CSP with BO are shown here. Each selection (five structure candidates) by machine learning is visualized in different colors. The result summary is also output as a text file, including the structure indices, the selection number, space group of the initial and optimized structures, energies, and so on, for quick checking as listed in Table 1.

In addition, initial and optimized structure data are written as text files in POSCAR and CIF format, so that users can see the structures using a crystal structure visualization tool such as VESTA [34].

### 3. Searching algorithms

The following searching algorithms are implemented in CrySPY:

- RS
- EA
- BO
- LAQA

RS relies on random structure generation and local structure optimization. Random structure generation is fundamental and used in other searching algorithms. EA puts emphasis on effective structure generation modeled after the theory of evolution, while BO and LAQA are the selection-type algorithms that can efficiently select potential candidates from a large number of structures by machine learning, reducing the number of searching trials. The key point here is that the selection-type algorithm can be used in combination with the structure generation methods by RS and EA.

| Select | Spg_num | Spg_sym | Spg_num_opt | Spg_sym_opt | E_eV_atom | Magmom | Opt   |
|--------|---------|---------|-------------|-------------|-----------|--------|-------|
| 251    | 1       | 107     | H4mm        | 0           | None      | NaN    | NaN   | skip  |
| 14     | 1       | 175     | P6/m        | 191         | P6/m/mmm  | −6.10057 | 27.5053 | done  |
| 273    | 1       | 44      | Imm2        | 44          | Imm2      | −6.54106 | 26.9209 | done  |
| 107    | 1       | 195     | P23         | 200         | Pm-3      | −6.49809 | 26.6711 | done  |
| 190    | 1       | 38      | Amm2        | 38          | Amm2      | −6.68704 | 25.6406 | done  |
| 111    | 2       | 71      | Immm        | 71          | Immm      | −6.50534 | 26.2337 | done  |
| 59     | 2       | 16      | P222        | 16          | P222      | −6.47951 | 26.1596 | done  |
| 288    | 2       | 12      | C2/m        | 12          | C2/m      | −6.79212 | 25.1641 | done  |
| 133    | 2       | 71      | Immm        | 71          | Immm      | −6.56943 | 25.576  | done  |
| 262    | 2       | 156     | P3m1        | 0           | None      | NaN    | NaN   | skip  |
| 93     | 3       | 150     | P321        | 0           | None      | NaN    | NaN   | skip  |
| 294    | 3       | 155     | R32         | 155         | R32       | −6.28959 | 26.4513 | not_yet |
| 172    | 3       | 177     | P622        | 191         | P6/m/mmm  | −5.96608 | 28.2994 | done  |
| 145    | 3       | 177     | P622        | 177         | P622      | −6.60823 | 24.9556 | done  |
| 176    | 3       | 115     | P-4 m2      | 115         | P-4 m2    | −6.56049 | 26.7792 | done  |
3.1. Random search

A specified number of structures are generated randomly at the beginning, and local structure optimization is performed one by one. CrySPY has two modes of random structure generation: atomic crystal structure generation and molecular crystal structure generation. The latest version of CrySPY employs PyXtal library [28] as default to generate such random structures.

3.1.1. Atomic crystal structure generation

CrySPY can generate crystal structures with specific symmetry and chemical compositions using PyXtal. Optionally, it is also possible to generate the crystal structures using find_wy program [35] that we have been using. As default, a space group is randomly selected and the lattice parameters are automatically set to appropriate values. The volume of the unit cell can be specified within an arbitrary range in CrySPY.

Non-symmetric random structures can also be generated using a built-in function of CrySPY for low-symmetry CSP. In this case, a crystal system (not a space group) and lattice parameters are randomly selected within a physically appropriate range, and a specified number of atoms are arranged in the cell at random.

3.1.2. Molecular crystal structure generation

Interfacing with PyXtal enables symmetric molecular crystal generation. When the molecular geometry is given, random crystal structures with molecules occupying both general and special Wyckoff positions according to a specified space group can be generated. The following is an example of the structure section of the input file, cryspy.in, for two formula units of Li₅PS₆:

```
[structure]
struc_mode = mol
natot = 16
atype = Li P S
nat = 6 2 8
mol_file = ./Li.xyz ./PS4.xyz
nmol = 6 2
```

Here, Li and PS₄ units are defined in ./Li.xyz and ./PS4.xyz files, respectively. A single atom is treated as a molecule in the molecular crystal structure generation mode. In this example, a random molecular crystal structure is composed of six Li molecules (atoms) and two PS₄ molecules as specified in the input variable, nmol.

We generated 50 random molecular crystal structures using the above settings to validate whether a stable structures can be obtained. Total energy calculations and local structure optimizations were carried out using the density functional theory (DFT) with the projector-augmented wave (PAW) method [36] with the VASP code [29]. The generalized gradient approximation (GGA) by Perdew, Burke, and Ernzerhof [37] was employed for exchange-correlation functional. A cutoff energy of 625 eV for the plane-wave expansion of the wave function and k-point mesh density of 80 Å⁻³ were used. The atomic coordinates and cell parameters were fully optimized until forces acting on every atom became at least smaller than 0.01 eV/Å. Figure 2 shows the most and second most stable structures predicted by our CSP simulation. Space group of the most and second most stable structures are P43m and Pmn2₁, respectively. The total energy difference between them is only 10 meV/atom. Unfortunately, the former (P43m) was not observed in the experiment, while the latter (Pmn2₁) was obtained and is known as a stable structure, γ-Li₅PS₆ [38]. This can happen because the calculated energy difference is much smaller than the chemical accuracy (1 kcal/mol = 43 meV/atom). We might evaluate the total energy of the former structure a little too low with the DFT calculation. The predicted and experimental crystal parameters of γ-Li₅PS₆ were listed in Table 2. It is an almost perfect match except for the tiny amount of differences in lattice constants and atomic positions. These results demonstrate that the symmetric molecular structure generation method is quite useful for stable structure predictions of molecular crystals.

![Figure 2](image_url)

*Figure 2.* Two stable structures of Li₅PS₆ predicted by a CSP simulation drawn by VESTA [34]. (a) Space group P43m and (b) space group Pmn2₁. Green, purple, and yellow spheres represent Li, P, and S atoms, respectively. The structure of Pmn2₁ corresponds to γ-Li₅PS₆ that is known as the most stable structure [38].
Table 2. Crystal parameters of predicted structure, and experimental ones [38] for γ-Li₃PS₄.

| Atom | Site | x    | y    | z    |
|------|------|------|------|------|
| Li   | 4b   | 0.2432| 0.3151| 0.9970|
| Li   | 2a   | 0    | 0.1450| 0.4853|
| P    | 2a   | 0    | 0.8173| 0.9956|
| S    | 4b   | 0.2178| 0.6722| 0.8867|
| S    | 2a   | 0    | 0.1108| 0.8849|
| S    | 2a   | 0    | 0.8090| 0.3276|

| Atom | Site | x    | y    | z    |
|------|------|------|------|------|
| Li   | 4b   | 0.2502| 0.3309| 0.0179|
| Li   | 2a   | 0    | 0.1450| 0.4866|
| P    | 2a   | 0    | 0.8218| 0.9942|
| S    | 4b   | 0.2185| 0.6717| 0.8857|
| S    | 2a   | 0    | 0.1083| 0.8886|
| S    | 2a   | 0    | 0.8049| 0.3230|

Even in the molecular crystal structure generation mode, CrySPY can specify the volume of the unit cell within an arbitrary range, keeping the molecular geometry. The feature to generate non-symmetric molecular crystal structures is also implemented in CrySPY for low-symmetry CSP. After randomly selecting a space group, the molecule is placed at a Wyckoff position and rotated randomly without considering the space group, leading to breaking the symmetry.

3.1.3. Constraint of minimum interatomic distance
In generating random structures, CrySPY can restrict the interatomic distance so that it is not too close. Here is an example of the input file to limit minimum interatomic distance for a Y-Co system:

```
[structure]
mindist_1 = 2.018
mindist_2 = 1.815
```

This means that minimum interatomic distances of Y-Y, Y-Co, and Co-Co are limited to 2.0, 1.8, and 1.5 Å, respectively. Structures with interatomic distances shorter than these values are automatically eliminated.

3.1.4. Constraint of space group
Users can limit space group. As an example, generated structures are limited to cubic systems with the following setting:

```
[structure]
spgnum = 195-230
```

3.2. Evolutionary algorithm
EA can be more effective than RS in generating structure. We implemented EA in CrySPY, referring to USPEX [11–13] and XtalOpt [25]. EA is based on that a population of structures is evolved, driven by survival of the fittest. Some lower-energy structures can become parents of a new generation of structures. CrySPY employs tournament selection and roulette selection for selecting parents from a population of surviving parents. Evolutionary operations for producing offspring from parents are of key importance. Crossover, permutation, and strain operations are available in CrySPY. The new generation includes not only structures produced by these evolutionary operations but also random structures and structures taken over by elite selection. In the elite selection, a part of the surviving parents are passed on to the next generation without mutation to keep the current best structures. An example of the EA section of the input file for a binary system is following:

```
[EA]
n_pop = 20
n_crssov = 10
n_perm = 3
n_strain = 5
n_rand = 2
n_elite = 2
n_fittest = 10
slct_func = TNM
t_size = 2
maxgen_ea = 5
mindist_ea_1 = 2.018
mindist_ea_2 = 1.815
```

Minimum interatomic distances can be also limited in producing offspring.

3.2.1. Parents selection
The number of surviving structures is specified in the input variable, n_fittest. The specified number of lower-energy structures excluding duplicates can be candidates to produce offspring. CrySPY has two methods to select the candidates. One is the tournament selection. A specified number of candidates (t_size) are randomly selected, and the candidate with the lowest energy among them is chosen as the parent. The other is the roulette selection. In this method, the probability of being selected as a parent is determined as:

$$p_i = \frac{f_i'}{\sum_k f_k'},$$

(1)

where $p_i$ and $f_i'$ are the probability and linear-scaled fitness of $i$th candidate, respectively. The linear scaling of fitness is done by:

$$f_i' = \frac{a - b}{f_{max} - f_{min}} f_i + \frac{b f_{max} - a f_{min}}{f_{max} - f_{min}},$$

(2)

where fitness $f_i$ represents sign-reversed energy of $i$th candidate, $a$ and $b$ are parameters that define the range.
3.2. Crossover

Figure 3 depicts a schematic crossover operation. Selected two parents are sliced near the center in a lattice vector after randomly translating their origin. Two offspring are produced by swapping the sliced parents, each carrying some genetic information. The number of atoms in the offspring is not necessarily the same as that of the parents at this time. Only a structure with a larger number of atoms is adopted. The number of atoms in the adopted structure is adjusted by removing or adding atoms near the border. Finally, the interatomic distances are checked.

3.2.3. Mutation

In permutation operation, two atoms of different elements are swapped in a selected parent. The number of times for permutation can be specified in the input file. Strain operation is also used as mutation. Lattice vectors $a$ of a selected parent are transformed to $a'$ by applying a strain matrix:

$$a' = \begin{pmatrix} 1 + \eta_1 & \frac{1}{2} \eta_6 & \frac{1}{2} \eta_5 \\ \frac{1}{2} \eta_4 & 1 + \eta_2 & \frac{1}{2} \eta_4 \\ \frac{1}{2} \eta_3 & \frac{1}{2} \eta_4 & 1 + \eta_3 \end{pmatrix} a,$$

where $\eta_i$ are given by normal distribution with a mean of zero and a standard deviation.

3.3. Bayesian optimization

Our BO algorithm has been implemented in CrySPY. CSP using BO has been successfully applied to the known systems such as NaCl and Y$_2$CO$_3$ [16]. The results have demonstrated that BO can significantly reduce the number of searching trials required to find the global minimum structure by 30–40% in comparison with pure RS, leading to much less computational cost. Although the candidate structures were created by random structure generation in the above study, it is possible to select from candidate structures generated by other methods such as EA.

BO is a well-established technique for black-box optimizations [39–41]. Figure 4 shows a CSP simulation procedure by BO, which is automated in CrySPY. First, a specified number of initial structures are randomly generated. We have to calculate the structure descriptors of the initial structures to learn the data and predict promising candidates. In the first selection, a specified number of structures are randomly selected and locally optimized to prepare the first training data. The descriptors for the optimized structures are updated. Then, the next candidates are selected by BO, and this sequence of the procedure is repeated. The below is an example of the BO section of the input file, cryspy.in:

```plaintext
[BO]
  nselect_bo = 5
dscrt = FP
  score = TS
  fp_rmin = 0.5
  fp_rmax = 5.0
  fp_npoints = 20
  fp_sigma = 1.0
  max_select_bo = 20
```

Here, one can specify the number of structures to be selected at a time, acquisition function, and parameters for calculating the structure descriptors.

3.3.1. Structure descriptor

To utilize machine learning, we have to convert crystal structures into vector data, called structure descriptor. Structure descriptors should measure the similarity of crystal structures and correlation between energy and structure. Although various structure descriptors have been developed recently [19,42–47], CrySPY currently supports only the $F$-fingerprint of Oganov and Valle [48] expressed as

$$F_{AB}(R) = \sum_{A_i, cell} \sum_{b_i} \frac{\delta(R - R_{ij})}{4\pi R_j^3 N_A N_B \Delta} - 1.$$  (4)

$F_{AB}(R)$ is a function of distance $R$, where $A$ and $B$ are chemical elements, $R_{ij}$ is the interatomic distance between atoms $i$ and $j$, $N_A$ and $N_B$ are the number of $A$ and $B$ atoms, and $V$ is the unit cell volume. $\delta(R - R_{ij})$ is a Gaussian-smearred delta function, and the double sum runs over all atoms $i$ within the unit cell and all $j$ within the distance between minimum and maximum cutoff distance. The $F$-fingerprint is discretized over bins of width $\Delta$ to be treated as a vector.

The parameters for structure descriptors would affect the CSP performance. A common problem in BO is the need to predetermine the parameters. We
also proposed an information measure of the distribution to estimate an appropriate parameter value, which can be predetermined [17]. CrySPY will be further developed to include other structure descriptors such as atom-centered symmetry functions [19,42,43], Smooth Overlap of Atomic Positions (SOAP) [44], orbital-field matrix (OFM) [45], and crystal graph [46]. Although various descriptors have been developed, selecting promising candidates in multi-specie systems becomes more difficult as the dimensions of the descriptors increase and become more complex. In general, BO has a problem with searching in high-dimensional space [49,50]. We cannot tell at this time whether BO can work efficiently in ternary or quaternary systems. To tackle this problem, investing the searching efficiency in multi-specie systems with the same atomic configuration but different number of atomic species such as diamond, zinc blende, and chalcopyrite structures will be a key task for us. Moreover, we will investigate the efficiency of BO when dimensionality reduction such as principal component analysis is applied to complicated descriptors for future work.

3.3.2. COMBO in CrySPY
CrySPY is interfaced with COMBO [26], which is a Python library designed for BO that employs Thompson sampling (TS) [41], random feature maps [51], and automatic hyperparameter tuning. CrySPY does not require complex machine learning techniques for users. A Gaussian process [52] is commonly used in BO to predict a true function. The uncertainty of the predicted function is also evaluated as predictive variance. The next candidates are selected based on the

Figure 4. Simulation procedure in BO, which is automated in CrySPY.
acquisition function computed from the predictive value and variance. The following acquisition functions are available in CrySPY: probability of improvement (PI) [53], expected improvement (EI) [39], and TS. BO enables us to balance the tradeoff between exploration and exploitation of the search space. The candidates with lower predicted values and large variance are preferentially selected. CrySPY has an additional feature to output the predictive value and variance as a text file during CSP simulations.

A random feature map can be used instead of a Gaussian process to speed up the computation. It allows us to approximate a Gaussian process with a Bayesian linear model. The number of features in the random feature map, which determines the accuracy and computational complexity, is specified in the input file as:

```plaintext
[BO]
  num_rand_basis = 2000
```

### 3.4. LAQA

As mentioned above, BO requires a structure descriptor, and the searching efficiency strongly depends on the structure descriptor. For non-experts, there are many difficulties such as how to predetermine the parameters for the structure descriptor. Besides, there are many cases where the selected candidate with a high energy structure does not provide important information for learning, and the local structure optimization of that candidate takes inefficient time. To simplify and improve the efficiency, another selection-type algorithm, LAQA, has been developed [18]. LAQA allows us to find the most stable structure with a minimum number of total local optimization steps. In our previous study, the computational cost can be significantly reduced by 50–90% compared to RS [18]. In the case of LAQA as well as BO, the candidate structures can be generated by any methods such as EA.

Figure 5 illustrates a CSP simulation procedure by LAQA, which is automated in CrySPY. Unlike conventional CSP methods, in LAQA, local structure optimization of the selected candidates is not performed all at once. We proceed only a few optimization steps for the selected candidates, and pause the simulation. LAQA can roughly estimate the final energy during local optimization, and the total computational cost can be reduced by controlling the local optimization step based on that estimated energy. A rough estimate of the final energy is used as the score $L$, and predicted using a quadratic approximation from the current energy and force acting each atom as:

$$ L = -E + \frac{F^2}{2\Delta F}, $$

where $E$ is the energy per atom, $F$ is the averaged norm of the atomic force, $\Delta F$ is the force difference from the previous step, and $c$ is the bias parameter. The next candidates are selected based on the score $L$ and preferentially optimized, thereby enabling an efficient search for the global minimum without unnecessary optimization calculations. The below is an example of the LAQA section of the input file, `crys.py.in`:

```plaintext
[LAQA]
  nselect_laqa = 5
```

The input for LAQA is quite simple. One needs to specify the number of candidates to be selected at a time. Optionally, the bias parameter $c$ can be specified. It is important to note that LAQA does not require any structure descriptors. One does not have to worry about the descriptors getting complicated in large and multi-species systems.

### 4. Other features

#### 4.1. Loading initial structures

User-defined initial structures can be used instead of random structures in CrySPY. The initial structure data should be put in the relative path from the input file) before starting the CSP simulation. The loading flag also should be turned on in the input file as:

```plaintext
[option]
  load_struc_flag = True
```

#### 4.2. Appending structures

Initial structures can be appended at any time during the CSP simulation, except when using EA. By simply increasing the value of the input variable, `tot_struc`, in the basic section, CrySPY can append random structures to the initial structure data. Furthermore, CrySPY has a feature to generate and append structures using EA from already optimized structures and their energy data by turning on the flag as:

```plaintext
[option]
  append_struc_ea = True
```

The input variables for EA as listed in Section 3.2 are also needed. This allows us to use the hybrid algorithms of EA and BO, or EA and LAQA, which could be more efficient. We will investigate searching efficiency of the hybrid algorithms in several materials for future work.

#### 4.3. Skip and recalculating

Sometimes first-principles calculations do not work and give strange energies with wrong electronic structures. In such a case, CrySPY can skip the optimization and energy evaluation by manually editing the file, `./work/000016/stat_job` (relative path from the input file as an example of structure ID 16). If
necessary, CrySPY can recalculate the structure by specifying the structure ID in the input file as:

```
[option]
recalc = 16
```

4.4. Step data

Optionally, CrySPY can save the data in the process of local structure optimization for the structure, energy, force, and stress, by turning on the corresponding flag in the input file as:

```
[option]
energy_step_flag = True
struc_step_flag = True
force_step_flag = True
stress_step_flag = True
```

These can be used to generate a database for various applications such as machine learning potential generation.

5. Conclusion

We introduced a CSP tool CrySPY written in Python 3, which runs on Unix/Linux platforms. This open-source code allows us to automate structure generation, structure optimization, energy evaluation, and efficiently selecting candidates using machine learning. The searching algorithms available in CrySPY are highlighted. In particular, the
selection-type algorithms can efficiently select candidates for priority optimization, reducing the computational cost significantly. Non-experts can conduct CSP simulations using CrySPY without the need for special skills. CrySPY is interfaced with several structure optimizers such as VASP, QUANTUM ESPRESSO, OpenMX, soiap, and LAMMPS. The current version of the software is 0.9.1 at the time of writing. CrySPY will be further developed to include more structure descriptors and new features. The open-source code is developed in github. The documentation and utility of CrySPY are also available. We hope our open-source software will help to standardize research with CSP for materials discovery and design.

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Disclosure statement

No potential conflict of interest was reported by the authors.

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