

User manual for op_tools ver 0.2.0

Hideo Doi (doi.hideo.chemistry@gmail.com)

April 2022

目次

What is this?	2
How to install	2
data requirement	2
Common settings for all order parameters	2
Neighborhood particle setting	2
Setting for the number of times for averaging	3
List of order parameters	4
<i>A</i> : neighborhood parameters	4
<i>B</i> : bond angle analysis (BAA)	4
<i>C</i> : centrosymmetry parameter analysis (CPA)	5
<i>D</i> : neighbor distance analysis (NDA)	5
<i>F</i> : angular Fourier series (AFS) like parameter	6
<i>H</i> : angle histogram analysis (AHA)	6
<i>I</i> : tetrahedron order parameter (TOP)	7
<i>Q</i> : Bond order parameter	8
<i>W</i> : Bond order parameter	9
<i>Q2, W2</i> : function weighting Bond order parameter	10
<i>LQ, LW</i> : local Bond order parameter	11
<i>S</i> : Onsager' s parameter	12
<i>T</i> : McMillan' s Sigma	12
<i>Z</i> : user define order parameter	13
Setting for many types analysis	13
Output format	14
Reference	14

What is this?

A python module for analyzing crystal structures in molecular and particle simulations. The environment surrounding each particle or atom is evaluated and converted into a numerical value. These values are also called **order parameters**. Particles in the similar environment will have the same numerical value, while particles in the different environment will not have it. For body-centered cubic (BCC) and face-centered cubic (FCC) lattices, the order parameter is known to have a specific value, which can be used to determine. For example, the location of the boundary between solid and liquid. This type of analysis is used in simulations of melting solid metals and in the analysis of grain boundaries of crystals.

How to install

```
git clone https://github.com/hdoi/op_tools.git
cd op_tools
pip install -e .
```

data requirement

1. coord : List of coordinates of each particles. Such as [[0,0,0], [0,0,1], ...] like [[x1, y1, z1], [x2, y2, z2], ...]
2. direct (optional) : Direction vector of each particle. It may be a three-dimensional unit vector or a quaternion. Such as [[1,0,0], ...] or [[1,0,0,0], ...]. In the case of a three-dimensional unit vector, the order is [x, y, z]. In the case of quaternions, the order is [w, x, y, z]. In the case of a mass point or spherical particle, It has no direction vector. There is no problem with direct = [].
3. box_length : The length of the simulation box. Such as [3, 3, 3]
4. op_settings : Settings for calculating each order parameter

Common settings for all order parameters

Neighborhood particle setting

The numerical value of the order parameter for each particle is calculated using the coordinates of nearby particles. The particles near a particle i are called the neighborhood particles. There are three methods implemented in this program to determine the neighborhood particles. The first method is to consider a particle to be an adjacent particle of particle i if it is within a certain distance (adjacency radius) from particle i . The advantage of this method is that it is relatively easy to implement and user-friendly. The disadvantages are that the number of neighborhood particles may be different for each particle and that it is scale dependent. For example, specifying an adjacency radius of 1 nm and 1 angstrom will give different results. This is called scale-dependency. The second method is to use n particles close to particle i as neighbors. The advantage is that there is no scale-dependency and the adjacent particles do not change if the length unit changes, because the relationship of distance is unchanged. The disadvantages are that it is not easy to implement, the computational cost is slightly higher, and it is not widely used. The third method is Delaunay partitioning (**Voronoi diagram**). After performing the Voronoi partitioning, two particles sharing each Voronoi facet are considered as adjacent particles.

Setting for neighborhood radius

```
op_settings = {  
    'radius' : [1.5, 1.75] ,  
    ... ( other parameter settings ) }
```

User can specify in the list format, and it is also possible to specify multiple conditions at the same time. In the above example, the radius setting is 1.5 and 1.75. ..

Setting for the number of neighborhood particles

```
op_settings = {  
    'neighbor' : [8, 12] ,  
    ... ( other parameter settings ) }
```

User can specify in the list format, and it is also possible to specify multiple conditions at the same time. In the above example, the number of neighborhood particles setting is 8 and 12. ..

Setting for Delaunay partitioning

```
op_settings = {  
    'Delaunay': ['standard']  
    ... ( other parameter settings ) }
```

In the above example, Delaunay partitioning is used.

Setting for the number of times for averaging

When the classification performance of the order parameter of each particle does not enough performance, to increase the performance, user can perform the averaging calculation. The average is calculated with the order parameter of particle i and the order parameter of neighborhood particles of particle i . $N_b(i)$ is a list of N neighborhood particles of particle i . $\tilde{N}_b(i)$ is a list of $N + 1$ particles that i itself is added to $N_b(i)$. The order parameter of the particles i before averaging is $X(i)$ and the order parameter after averaging is $Y(i)$. The following calculation is performed.

$$Y(i) = \frac{\sum_{j \in \tilde{N}_b(i)} X(j)}{N + 1}$$

This averaging calculation has the following advantages and disadvantages. The advantage is that the calculation cost is low, and it is possible to improve the classification performance of the same order parameter, Any numerical value can be used, and it can be averaged many times. On the other hand, the disadvantage is that more information is used to calculate the order parameters, and the resolution of the order parameters is decreased. The order parameters are calculated using the information of neighborhood particles, and the averaging calculation is calculated using the order parameters of the neighborhood particles. Each time the averaging operation is performed, the information on neighborhood particles of neighborhood particles is used. In other words, more particles are used to calculate order parameters, and information on far particles is used. This is not a problem in homogeneous systems, but it is a problem in inhomogeneous systems (such as interfaces). In inhomogeneous systems, it is likely to be used in many cases for the purpose of wanting to know the position where the abnormality is occurring or want to know the position of the interface, etc. Using distant particle information leads the indistinctness of position.

The number of times of this averaging is specified by `op_settings` specifying the calculation condition of the order parameter as follows. Note that this value starts from 0, and in the case of 0, averaging is not performed.

```
op_settings = {
    'ave_times'      : 1,
    ... ( other parameter settings )
}
```

List of order parameters

A : neighborhood parameters

common neighborhood parameter A [1], predominant common neighborhood parameter P [2], another predominant common neighborhood parameter N [2] are calculated using very similar formulas. Order parameter P and N are implemented as a another type of order parameter A .

$$A^{(a=0, type=A, m)}(i) = A^{(a=0, m)}(i) = \frac{1}{N} \sum_{j \in N_b(i)} \left| \sum_{k \in N_b(i, j)} (r_{ik} + r_{jk}) \right|^2$$

$$A^{(a=0, type=P, m)}(i) = P^{(a=0, m)}(i) = \frac{1}{N} \sum_{j \in N_b(i)} \left| \sum_{k \in N_b(i, j)} (r_{ij} + r_{kj}) \right|^2$$

$$A^{(a=0, type=N, m)}(i) = N^{(a=0, m)}(i) = \frac{1}{N} \left| \sum_{j \in N_b(i)} \sum_{k \in N_b(i, j)} (r_{ij} + r_{kj}) \right|^2$$

The variable a is the number of times to perform averaging the value among neighborhood particles. The list $N_b(i)$ is the list of N neighborhood particles of particle i . The list $N_b(i, j)$ is the list of m nearest neighborhood particles from particle i and particle j . The variable $type$ specifies the type of order parameter.

The following example shows the conditions for calculating the order parameter with 12 neighborhood particles and 2.0 adjacency radius.

```
op_param = {
    'neighbor': [12],
    'radius': [1.5],
    'ave_times': 1,
    'm_in_A': [2],
    'types_in_A': ['A', 'P', 'N'],
    'analysis_type': ['A']}

```

B : bond angle analysis (BAA)

Bond angle analysis B is calculated as follows. [3]

$$B^{(a=0, m, n, \phi)}(i) = \frac{1}{N(N-1)/2} \sum_{j > k \in N_b(i)} f^{(m, n, \phi)}(\theta_{jik})$$

$$f^{(m, n, \phi)}(\theta_{jik}) = \cos^n(m\theta_{jik} + \phi)$$

The variable a is the number of times to perform averaging the value among neighborhood particles. The list $N_b(i)$ is the list of N neighborhood particles of particle i . The variable θ_{jik} is the angle between vector

r_{ij} and vector r_{ik} . The variable m is the coefficient of the angle. The variable n is the exponent of the cos function. The variable ϕ is the intercept of angle.

The following example shows the conditions for calculating the order parameter with 12 neighborhood particles and 2.0 adjacency radius.

```
op_param = {
  'neighbor': [12],
  'radius': [1.5],
  'ave_times': 1,
  'm_in_B': [2],
  'phi_in_B': [0],
  'n_in_B': [1, 2],
  'analysis_type': ['B']}
```

C : centrosymmetry parameter analysis (CPA)

centrosymmetry parameter C is calculated as follows. [4]

$$C^{(a=0, type='half', mode)}(i) = \sum_{j \in M_b(i)} |r_{ij} + r_{ik}|^2$$

$$C^{(a=0, type='all', mode)}(i) = \sum_{j \in N_b(i)} |r_{ij} + r_{ik}|^2$$

The variable a is the number of times to perform averaging the value among neighborhood particles. The list $N_b(i)$ is the list of N neighborhood particles of particle i . The variable $M_b(i)$ is the list of $N/2$ nearest neighbors of particle i . The variable $type$ is a variable that specifies the number of times to add vectors. The variable $mode$ is indicating how to determine particle K . When the variable $mode$ is 'dist', particle k is the particle in the list $N_b(i)$ nearest from the coordinate r'_j , opposite side of particle r_i . When the variable $mode$ is 'angle', particle k is the particle in the list $N_b(i)$ whose angle θ_{jik} is closest to π . Two methods were implemented for use with liquids as well as solids.

The following example shows the conditions for calculating the order parameter with 12 neighborhood particles and 2.0 adjacency radius.

```
op_param = {
  'neighbor': [12],
  'radius': [2.0],
  'ave_times': 1,
  'types_in_C' : ['half'],
  'modes_in_C' : ['dist'],
  'analysis_type': ['C']}
```

D : neighbor distance analysis (NDA)

neighbor distance analysis D is calculated as follows. [5]

$$D_i^{(a=0, f_{ij}, f_{ik}, f_{jk})} = \frac{1}{N(N-1)/2} \sum_{j>k \in N_b(i)} f_{ij}(r_{ij}) f_{ik}(r_{ik}) f_{jk}(r_{jk})$$

The variable a is the number of times to perform averaging the value among neighborhood particles. The list $N_b(i)$ is the list of N neighborhood particles of particle i . The functions $f_{ij}(r_{ij}), f_{ik}(r_{ik}), f_{jk}(r_{jk})$ are some function with distance as argument and return value. The variables r_{ij}, r_{ik}, r_{jk} are the distances of the vectors r_{ij}, r_{ik}, r_{jk} respectively. The distances also satisfy $r_{ij} \leq r_{ik}$.

The following example shows the conditions for calculating the order parameter with 12 neighborhood particles and 2.0 adjacency radius. The following example is for the function $\{f_{ij}(r), f_{ik}(r), f_{jk}(r)\} = r$.

```
def f_1(r):
    return r

op_settings = {
    'neighbor': [12],
    'radius': [2.0],
    'ave_times': 1,
    'function': [f1],
    'analysis_type': ['D'] }
```

F : angular Fourier series (AFS) like parameter

angular Fourier series parameter F is calculated as follows. [6][7]

$$F_i^{(a=0, f_1, f_2, m)} = \frac{1}{N(N-1)/2} \sum_{j>k \in N_b(i)} f_1(r_{ij}) f_2(r_{ik}) \cos(l\theta_{jik})$$

The variable a is the number of times to perform averaging the value among neighborhood particles. The list $N_b(i)$ is the list of N neighborhood particles of particle i . The function $f^1(r), f^2(r)$ is some function with distance as argument and value as return value. The variable l is the coefficient of the angle. The variable θ_{jik} is the angle between vector r_{ij} and vector r_{ik} . The distance satisfies $r_{ij} \leq r_{ik}$.

The following example shows the conditions for calculating the order parameter with 12 neighborhood particles and 2.0 adjacency radius. The following example is for the function $\{f_{ij}(r), f_{ik}(r)\} = r$.

```
def f_1(r):
    return r

op_settings = {
    'neighbor': [12],
    'radius': [2.0],
    'ave_times': 1,
    'l_in_F': [1],
    'function': [f1],
    'analysis_type': ['F']}
```

H : angle histogram analysis (AHA)

angle histogram analysis H は次の式で計算される。

$$\begin{aligned}
h^{(b=0,bin)}(i) &= \frac{1}{N(N-1)/2} \sum_{j>k \in N_b(i)} histogram^{(bin)}(\theta_{jik}) \\
h^{(b,bin)}(i) &= \frac{1}{N+1} \sum_{j \in \tilde{N}_b(i)} h^{(b-1,bin)}(j) \\
H^{(a=0,b,bin,\nu)}(i) &= FFT_{amplitude}(h^{(b,bin)}(i))\delta(X-\nu)
\end{aligned}$$

The variable a is the number of times to perform averaging the value among neighborhood particles. The list $N_b(i)$ is the list of N neighborhood particles of particle i . The variable $\tilde{N}_b(i)$ is a list of $N+1$ particles with particle i itself added to the variable $N_b(i)$. The variable b is the number of times the histogram of each particle is averaged with the histograms of its neighboring particles. The variable θ_{jik} is the angle between the vector r_{ij} and the vector r_{ik} . The function *histogram* creates a histogram of the number of bins bin . The function *FFT_{amplitude}* computes amplitude by Fourier transforming the histogram. The function δ is the Dirac delta function. The variable ν is the frequency component of the histogram after the Fourier transform. This is because the delta function δ becomes 1 only when the frequency component X is the same as ν .

The following example shows the conditions for calculating the order parameter with 12 neighborhood particles and 2.0 adjacency radius.

```

op_settings = {
    'neighbor': [12],
    'radius': [2.0],
    'ave_times': 1,
    'b_in_H': 1,
    'bin_in_H': [24],
    'nu_in_H': [3],
    'analysis_type': ['H'] }

```

I : tetrahedron order parameter (TOP)

Tetrahedron order parameter I is calculated as follows. [8][9]

$$I^{(a=0)}(i) = 1 - \frac{3}{8} \sum_{j>k \in N_b(i)} \{\cos(\theta_{jik}) + 1/3\}^2$$

The variable a is the number of times to perform averaging the value among neighborhood particles. The list $N_b(i)$ is the list of N neighborhood particles of particle i . The variable θ_{jik} is the angle between the vector r_{ij} and the vector r_{ik} .

The following example shows the conditions for calculating the order parameter with 4 neighborhood particles and 2.0 adjacency radius.

```

op_settings = {
    'neighbor': [4],
    'radius': [2.0],
    'ave_times': 1,
    'analysis_type': ['I'] }

```

Q : Bond order parameter

Bond order parameter Q is calculated as follows. [10][11]

$$Q_i^{(l,a=0,b)} = \sqrt{\frac{4\pi}{2l+1} \sum_{m=-l}^l |q_{lm}^{(l,a=0,b)}(i)|^2}$$

$$q_{lm}^{(l,a=0,b)}(i) = \frac{1}{N+1} \sum_{j \in \tilde{N}_b(i)} q_{lm}^{(l,a=0,b-1)}(j)$$

$$q_{lm}^{(l,a=0,b-1)}(i) = \frac{1}{N} \sum_{j \in N_b(i)} Y_{lm}(\theta_{ij}, \phi_{ij})$$

The variable a is the number of times to perform averaging the value among neighborhood particles. The list $N_b(i)$ is the list of N neighborhood particles of particle i . The variable $\tilde{N}_b(i)$ is a list of $N+1$ particles with particle i itself added to $N_b(i)$. The variable b is the number of times the spherical harmonic function is averaged over adjacent particles. The function Y is the spherical harmonic function. The variables θ_{ij}, ϕ_{ij} are angles in the spherical coordinate system representation of vector r_{ij} , where θ is the angle from the z axis and ϕ is the angle from the x axis.

The following example shows the conditions for calculating the order parameter with 12 neighborhood particles and 2.0 adjacency radius.

```
op_settings = {
    'neighbor': [12],
    'radius': [2.0],
    'ave_times': 1,
    'l_in_Q': [4],
    'b_in_Q': 1,
    'analysis_type': ['Q'] }
```

Steinhardt's order parameter [10] can be calculated with the following setting.

```
op_settings = {
    'neighbor': [12],
    'radius': [2.0],
    'ave_times': 0,
    'l_in_Q' : [4,6],
    'b_in_Q' : 0,
    'analysis_type' : ['Q'] }
```

Lechner's order parameter [11] can be calculated with the following setting.

```
op_settings = {
    'neighbor': [12],
    'radius': [2.0],
    'ave_times': 0,
    'l_in_Q' : [4,6],
    'b_in_Q' : 1,
    'analysis_type' : ['Q'] }
```


W : Bond order parameter

Bond order parameter W is calculated as follows. [10][11]

$$W_i^{(l,a=0,b)} = \frac{\sum_{m_1+m_2+m_3=0} \begin{pmatrix} l & l & l \\ m_1 & m_2 & m_3 \end{pmatrix} q_{lm_1}^{(l,a=0,b)}(i) q_{lm_2}^{(l,a=0,b)}(i) q_{lm_3}^{(l,a=0,b)}(i)}{\left(\sum_{m=-l}^l |q_{lm}^{(l,a=0,b)}(i)|^2\right)^{3/2}}$$

$$q_{lm}^{(l,a=0,b)}(i) = \frac{1}{N+1} \sum_{j \in \tilde{N}_b(i)} q_{lm}^{(l,a=0,b-1)}(j)$$

$$q_{lm}^{(l,a=0,b=0)}(i) = \frac{1}{N} \sum_{j \in N_b(i)} Y_{lm}(\theta_{ij}, \phi_{ij})$$

The variable a is the number of times to perform averaging the value among neighborhood particles. The list $N_b(i)$ is the list of N neighborhood particles of particle i . The variable $\tilde{N}_b(i)$ is a list of $N+1$ particles with particle i itself added to $N_b(i)$. The variable b is the number of times the spherical harmonic function is averaged over adjacent particles. The function Y is the spherical harmonic function. Variables θ_{ij}, ϕ_{ij} are angles in the spherical coordinate system representation of vector r_{ij} , where θ is the angle from the z axis and ϕ is the angle from the x axis. The variables m_1, m_2, m_3 take values from $-l$ to l , but are calculated only when $m_1 + m_2 + m_3 = 0$. The matrix $\begin{pmatrix} l & l & l \\ m_1 & m_2 & m_3 \end{pmatrix}$ is a Wigner 3- j symbol.

The following example shows the conditions for calculating the order parameter with 12 neighborhood particles and 2.0 adjacency radius.

```
op_settings = {
    'neighbor': [12],
    'radius': [2.0],
    'ave_times': 1,
    'l_in_Q': [4],
    'b_in_Q': 1,
    'analysis_type': ['W'] }
```

Steinhardt' s order parameter [10] can be calculated with the following setting.

```
op_settings = {
    'neighbor': [12],
    'radius': [2.0],
    'ave_times': 0,
    'l_in_Q': [4,6],
    'b_in_Q': 0,
    'analysis_type': ['W'] }
```

Lechner' s order parameter [11] can be calculated with the following setting.

```
op_settings = {
    'neighbor': [12],
    'radius': [2.0],
    'ave_times': 0,
    'l_in_Q': [4,6],
```

```
'b_in_Q': 1,
'analysis_type': ['W'] }
```

$Q2, W2$: function weighting Bond order parameter

$$Q2_i^{(l,a=0,b)} = \sqrt{\frac{4\pi}{2l+1} \sum_{m=-l}^l |q_{lm}^{(l,a=0,b)}(i)|^2}$$

$$q_{lm}^{(l,a=0,b)}(i) = \frac{1}{N+1} \sum_{j \in \tilde{N}_b(i)} q_{lm}^{(l,a=0,b-1)}(j)$$

$$q_{lm}^{(l,a=0,b=0)}(i) = \sum_{j \in N_b(i)} F(i,j) Y_{lm}(\theta_{ij}, \phi_{ij})$$

A function $F(i,j)$ can be used to weight between particles i and j .

The variable a is the number of times to perform averaging the value among neighborhood particles. The list $N_b(i)$ is the list of N neighborhood particles of particle i . The variable $\tilde{N}_b(i)$ is a list of $N+1$ particles with particle i itself added to $N_b(i)$. The variable b is the number of times the spherical harmonic function is averaged over adjacent particles. The function Y is the spherical harmonic function. The variables θ_{ij}, ϕ_{ij} are the angles in the spherical coordinate system representation of the vector r_{ij} , where θ is the angle from the z axis and ϕ is the angle from the x axis.

The order parameter $W2$ is calculated as follows.

$$W2_i^{(l,a=0,b)} = \frac{\sum_{m_1+m_2+m_3=0} \binom{l}{m_1} \binom{l}{m_2} \binom{l}{m_3} q_{lm_1}^{(l,a=0,b)}(i) q_{lm_2}^{(l,a=0,b)}(i) q_{lm_3}^{(l,a=0,b)}(i)}{\left(\sum_{m=-l}^l |q_{lm}^{(l,a=0,b)}(i)|^2 \right)^{3/2}}$$

If you use Delaunay partitioning, you can calculate Mickle' s order parameter as follows. [12]

```
def f1(j, voronoi_area_list, distance_list):
    weight = voronoi_area_list[j] / np.sum(voronoi_area_list)
    return weight
```

```
op_settings = {
    'Delaunay': ['standard'],
    'ave_times': 1,
    'l_in_Q': [4],
    'function_in_Q2': [f1],
    'analysis_type': ['Q2', 'W2'] }
```

In addition, you can use other weighting function, such as weighting based on distance. The following example is a setting weighting function based on the distance. The more distance is the more less weight.

```
def f1(j, voronoi_area_list, distance_list):
    sum_weight = 0
    for dist in distance_list:
        sum_weight += 1/dist
    weight = (1/distance_list[j]) / sum_weight
```

```
return weight
```

```
op_settings = {
    'Delaunay': ['standard'],
    'radius': [2.0],
    'ave_times': 1,
    'l_in_Q': [4],
    'function_in_Q2': [f1],
    'analysis_type': ['Q2'] }
```

LQ, LW : local Bond order parameter

These order parameters were implemented in imitation of [13][14][15].

$$\begin{aligned}
 LQ_i^{(l,a=0,b)} &= \frac{1}{N} \sum_{j \in N_b(i)} \frac{q_{lm}^{(l,a,b)}(i,j)}{|q_{lm}^{(l,a,b)}(i)| |q_{lm}^{(l,a,b)}(j)|} \\
 q_{lm}^{(l,a,b)}(i,j) &= \sum_{m=-l}^l q_{lm}^{(l,a,b)}(i) q_{lm}^{*(l,a,b)}(j) \\
 q_{lm}^{(l,a=0,b)}(i) &= \frac{1}{N+1} \sum_{j \in \tilde{N}_b(i)} q_{lm}^{(l,a=0,b-1)}(j) \\
 q_{lm}^{(l,a=0,b=0)}(i) &= \frac{1}{N} \sum_{j \in N_b(i)} Y_{lm}(\theta_{ij}, \phi_{ij})
 \end{aligned}$$

The variable a is the number of times to perform averaging the value among neighborhood particles. The list $N_b(i)$ is the list of N neighborhood particles of particle i . The variable $\tilde{N}_b(i)$ is a list of $N+1$ particles with particle i itself added to $N_b(i)$. The variable b is the number of times the spherical harmonic function is averaged over adjacent particles. The function Y is the spherical harmonic function. The variables θ_{ij}, ϕ_{ij} are the angles in the spherical coordinate system representation of the vector r_{ij} , where θ is the angle from the z axis and ϕ is the angle from the x axis.

$$\begin{aligned}
 LW_i^{(l,a=0,b)} &= \frac{\sum_{m_1+m_2+m_3=0} \begin{pmatrix} l & l & l \\ m_1 & m_2 & m_3 \end{pmatrix} l q_{lm_1}^{(l,a=0,b)}(i) l q_{lm_2}^{(l,a=0,b)}(i) l q_{lm_3}^{(l,a=0,b)}(i)}{\left(\sum_{m=-l}^l |l q_{lm}^{(l,a=0,b)}(i)|^2 \right)^{3/2}} \\
 l q_{lm}^{(l,a,b)}(i) &= \frac{1}{N} \sum_{j \in N_b(i)} \frac{q_{lm}^{(l,a,b)}(i) q_{lm}^{*(l,a,b)}(j)}{|q_{lm}^{(l,a,b)}(i)| |q_{lm}^{(l,a,b)}(j)|}
 \end{aligned}$$

The following example shows the conditions for calculating the order parameter with 12 neighborhood particles and 2.0 adjacency radius. That calculate the similar order parameters in [13].

```
op_settings = {
    'neighbor': [12],
    'radius': [2.0],
    'ave_times': 0,
    'l_in_Q': [4,6],
    'b_in_Q': 0,
    'analysis_type': ['LQ', 'LW'] }
```

S : Onsager' s parameter

Onsager' s order parameter S is calculated as follows. [16][17]

$$S^{(a=0,n)}(i) = \frac{\sum_{j \in N_b(i)} P_n(\cos(\theta))}{N}$$

The variable a is the number of times to perform averaging the value among neighborhood particles. The list $N_b(i)$ is the list of N neighborhood particles of particle i . The function P_n is **Legendre polynomial**. The variable N is Degree of the polynomial and is even. The variable θ is the angle between the direction vector $u(i)$ of particle i and the direction vector $u(j)$ of particle j . The variable $\cos(\theta)$ is usually calculated by $u(i) \cdot u(j)$.

When $n = 2, 4$, the order parameter S is calculated by the following equations, respectively.

$$S^{(a=0,n=2)}(i) = \frac{\sum_{j \in N_b(i)} [3 \cos^2(\theta) - 1]/2}{N}$$

$$S^{(a=0,n=4)}(i) = \frac{\sum_{j \in N_b(i)} [35 \cos^4(\theta) - 30 \cos^2(\theta) + 3]/8}{N}$$

The following example shows the conditions for calculating the order parameter with 12 neighborhood particles and 2.0 adjacency radius. Note that this order parameter requires the direction vector.

```
op_settings = {
    'neighbor': [12],
    'radius': [2.0],
    'ave_times': 1,
    'n_in_S' : [2],
    'analysis_type' : ['S']
}
```

T : McMillan' s Sigma

McMillan' s order parameter T is calculated as follows. [18]

$$T^{(a=0,n)}(i) = \frac{\sum_{j \in N_b(i)} \cos(2\pi z(i,j)/d) P_n(\cos(\theta))}{N}$$

The variable a is the number of times to perform averaging the value among neighborhood particles. The list $N_b(i)$ is the list of N neighborhood particles of particle i . The function P_n is **Legendre polynomial**. The variable N is Degree of the polynomial and is even. The variable θ is the angle between the direction vector u_i of particle i and the direction vector u_j of particle j . The variable $Z(i,j)$ is the distance from the plane P , which passes through particle i and is perpendicular to the direction vector u_i , to particle j . The variable D is the distance from layer to layer of the smectic phase of the liquid crystal.

When $n = 2$, the order parameter T is calculated as follows.

$$T^{(a=0,n=2)}(i) = \frac{\sum_{j \in N_b(i)} \cos(2\pi z(i,j)/d) [3 \cos^2(\theta) - 1]/2}{N}$$

The following example shows the conditions for calculating the order parameter with 12 neighborhood particles and 2.0 adjacency radius. Note that this order parameter requires the direction vector.

```
op_settings = {
    'neighbor': [12],
    'radius': [2.0],
    'ave_times': 1,
    'n_in_T': [2],
    'd_in_T': [1.0],
    'analysis_type' : ['T']
}
```

Z : user define order parameter

An order parameter Z is provided for the user to define.

The following example shows the conditions for calculating the order parameter with 12 neighborhood particles and 2.0 adjacency radius.

```
op_settings = {
    'neighbor': [12],
    'radius': [2.0],
    'analysis_type': ['Z'] }
```

The calculation part of the order parameter is op_tools/op_z_user_define.py. It is possible to edit this file and implement the order parameter that the user has thought of.

Setting for many types analysis

uj This takes much time.

```
def f_1(r):
    return r
op_settings = {
    # neighborhood particles settings
    'neighbor'      : [8],                # number of neighborhood particles
    'radius'        : [1.5],              # adjacency radius
    'ave_times'     : 1,                  # average time
    # A
    'op_types'      : ['A', 'P', 'N'],    # specify the type of analysis in order parameter A
    'm_in_A'        : [2, 4],             # number of neighborhood particles from particle i and j
    # B
    'm_in_B'        : [2],                # factor of angle
    'n_in_B'        : [1, 2],             # exponent of cos function
    'phi_in_B'      : [0],                # intercept of angle
    # C
    'types_in_C'    : ['half'],           # number of vector adding
    'modes_in_C'    : ['dist'],           # search method for opposite side particle
    # D
    'function'      : [f1]                # function
    # F
    'l_in_F'        : [1],                # factor of angle
```

```

# H
'b_in_H' : 1,          # number of averaging of angle histogram
'bin_in_H' : [24],     # bin of histogram
'nu_in_H' : [3],       # frequency components of angle, (pi/3)
# I
# Q
'b_in_Q'      : 1,      # number of averaging of spherical harmonic function
'l_in_Q'      : [2, 4, 6], # parameter for spherical harmonic
# Q2 W2
'function_in_Q2' : [f2], # weighting function
# LQ LW
# S
'n_in_S'      : [2],    # degree of Legendre polynomials
# T
'n_in_T'      : [2],    # degree of Legendre polynomials
'd_in_T'      : [1.0],  # distance between layers of smectic phase
'analysis_type': ['A', 'B', 'C', 'D', 'F', 'H', 'I',
'Q', 'W', 'Q2', 'W2', 'LQ', 'LW', 'S', 'T']] # Order parameter types

```

Output format

User can use as follows.

```

import order_tools
order_param_data = \
    op_analyze(coord, direct, box_length, op_settings)

```

To access the order parameters for each particle,

```

order_param_data['Q_N6_a=1_b=1']

```

- Q : Types of order parameter
- N6 : setting of neighborhood particles
- 'a=1_b=1' : parameter of order parameter calculation

Reference

- [1] J.D. Honeycutt, H.C. Andersen, Molecular dynamics study of melting and freezing of small Lennard-Jones clusters, The Journal of Physical Chemistry. 91 (1987) 4950-4963. doi:[10.1021/j100303a014](https://doi.org/10.1021/j100303a014).
- [2] A. Radhi, K. Behdian, Identification of crystal structures in atomistic simulation by predominant common neighborhood analysis, Computational Materials Science. 126 (2017) 182-190. doi:[10.1016/j.commatsci.2016.09.035](https://doi.org/10.1016/j.commatsci.2016.09.035).
- [3] G.J. Ackland, A.P. Jones, Applications of local crystal structure measures in experiment and simulation, Physical Review B - Condensed Matter and Materials Physics. 73 (2006) 1-7. doi:[10.1103/PhysRevB.73.054104](https://doi.org/10.1103/PhysRevB.73.054104).
- [4] C.L. Kelchner, S.J. Plimpton, J.C. Hamilton, Dislocation nucleation and defect structure during surface indentation, Physical Review B. 58 (1998) 11085-11088. doi:[10.1103/PhysRevB.58.11085](https://doi.org/10.1103/PhysRevB.58.11085).
- [5] A. Stukowski, Structure identification methods for atomistic simulations of crystalline materials, Modelling and Simulation in Materials Science and Engineering. 20 (2012) 045021. doi:[10.1088/0965-0393/20/4/045021](https://doi.org/10.1088/0965-0393/20/4/045021).

- [6] A.P. Bartók, R. Kondor, G. Csányi, On representing chemical environments, *Physical Review B - Condensed Matter and Materials Physics*. 87 (2013) 1-16. doi:[10.1103/PhysRevB.87.184115](https://doi.org/10.1103/PhysRevB.87.184115).
- [7] A. Seko, A. Togo, I. Tanaka, Descriptors for Machine Learning of Materials Data, in: *Nanoinformatics*, Springer Singapore, Singapore, 2018: pp. 3-23. doi:[10.1007/978-981-10-7617-6_1](https://doi.org/10.1007/978-981-10-7617-6_1).
- [8] P.-L. CHAU, A.J. HARDWICK, A new order parameter for tetrahedral configurations, *Molecular Physics*. 93 (1998) 511-518. doi:[10.1080/002689798169195](https://doi.org/10.1080/002689798169195).
- [9] E. Duboué-Dijon, D. Laage, Characterization of the Local Structure in Liquid Water by Various Order Parameters, *The Journal of Physical Chemistry B*. 119 (2015) 8406-8418. doi:[10.1021/acs.jpcb.5b02936](https://doi.org/10.1021/acs.jpcb.5b02936).
- [10] P.J. Steinhardt, D.R. Nelson, M. Ronchetti, Bond-orientational order in liquids and glasses, *Physical Review B*. 28 (1983) 784-805. doi:[10.1103/PhysRevB.28.784](https://doi.org/10.1103/PhysRevB.28.784).
- [11] W. Lechner, C. Dellago, Accurate determination of crystal structures based on averaged local bond order parameters, *Journal of Chemical Physics*. 129 (2008) 114707. doi:[10.1063/1.2977970](https://doi.org/10.1063/1.2977970).
- [12] W. Mickel, S.C. Kapfer, G.E. Schröder-Turk, K. Mecke, Shortcomings of the bond orientational order parameters for the analysis of disordered particulate matter, *Journal of Chemical Physics*. 138 (2013) 044501. doi:[10.1063/1.4774084](https://doi.org/10.1063/1.4774084).
- [13] E.B. Moore, E. De La Llave, K. Welke, D.A. Scherlis, V. Molinero, Freezing, melting and structure of ice in a hydrophilic nanopore, *Physical Chemistry Chemical Physics*. 12 (2010) 4124-4134. doi:[10.1039/b919724a](https://doi.org/10.1039/b919724a).
- [14] M. Fitzner, P. Pedevilla, A. Michaelides, Predicting heterogeneous ice nucleation with a data-driven approach, *Nature Communications*. 11 (2020) 1-9. doi:[10.1038/s41467-020-18605-3](https://doi.org/10.1038/s41467-020-18605-3).
- [15] G.A. Tribello, F. Giberti, G.C. Sossa, M. Salvalaglio, M. Parrinello, Analyzing and Driving Cluster Formation in Atomistic Simulations, *Journal of Chemical Theory and Computation*. 13 (2017) 1317-1327. doi:[10.1021/acs.jctc.6b01073](https://doi.org/10.1021/acs.jctc.6b01073).
- [16] L. Onsager, THE EFFECTS OF SHAPE ON THE INTERACTION OF COLLOIDAL PARTICLES, *Annals of the New York Academy of Sciences*. 51 (1949) 627-659. doi:[10.1111/j.1749-6632.1949.tb27296.x](https://doi.org/10.1111/j.1749-6632.1949.tb27296.x).
- [17] C. Zannoni, Distribution function and order parameters, in: G.R. Luchhurst, G.W. Gray (Eds.), *The Molecular Physics of Liquid Crystals*, Academic Press Inc. (London) Ltd., 1979: p. 51.
- [18] W. McMillan, Simple Molecular Model for the Smectic A Phase of Liquid Crystals, *Physical Review A*. 4 (1971) 1238-1246. doi:[10.1103/PhysRevA.4.1238](https://doi.org/10.1103/PhysRevA.4.1238).