

CluE Oxide (Cluster Evolution Oxide) is a open source program for simulating electron spin decoherence ab initio using the Cluster Correlation Expansion [1, 2]. CluE Oxide takes as inputs a pbd file and an options file. The pdb file specifies the coordinates and elements types of all the atoms, and the options file specifies all the additional information needed, such as the pulse sequence the applied magnetic field, the central spin coordinates and so on.

# 0.1 Overview

The first steps CluE Oxide takes is to read the options and PDB files to construct the spin system. This includes applying periodic boundary conditions, and dropping non-magnetic nuclei as well as those beyond the system radius specified in the options. Isotope substitutions can also be applied at this step, if specified in the input options. In addition to isotope substitution, atoms can be replaced with voids at a chosen rate. This can be helpful in hydrogen rich media, where the proton may be diluted, but including all the deuterons in the simulation would be too expensive. The main time cost advantage of dropping the deuterons comes in when numerically integrating the Schödinger equation, but for large systems a Bernoulli trial at each atom is also expensive. Algorithmically, time is saved by avoiding N Bernoulli trials with a p success rate. In theses cases CluE Oxide samples a random number from the binomial distribution parameterized by N and p, and then selects that many random hydrons to initialize as protons, ignoring the rest.

Once the system is set up, CluE Oxide construct the tensors for the active spins, based on the user input. CluE Oxide will default to applying the point-dipole approximation for spin-spin coupling, but the hyperfine tensors can be explicitly set to custom values. Nuclear electric quadrupole coupling tensors can also be set, but default to zero. There are several ways to specify tensors, and these are discussed later.

CluE Oxide has a grouping algorithm that allows atoms to be specified by a combination of properties found in the PDB file, such as element, serial number, or residue, among others. The groups can either require or forbid qualifiers.

If unspecified CluE Oxide will assume that the applied magnetic field points along the z-direction of the PDB frame, different orientations can be specified. Additionally, CluE Oxide can compute the orientation averaged signal. CluE Oxide can average over custom orientation with custom weights, or it use random orientations (inefficient but simple) as well as orientations from a Lebedev grid, which integrates exactly the orientation dependence of the lowest order terms in a polynomial or spherical harmonic expansion[3]. Both the spin Hamiltonian and the Lebedev grid have inversion symmetry; this is leveraged to cut the number of calculation in half.

For each orientations, the set of clusters to be included in the n-CCE simulations must be calculated. The clusters are the subsets of the system that contain the central spin (only implicitly included since it is always there), and up to n bath spins that form a connected graph, where edges are defined between two bath spin when they are within some user specified cutoff. Several cutoff options are available, including an orientation independent distance cutoff, an orientation dependent dipole-dipole coupling,  $\Delta A$ , and others based on the analytic solution of a simplified three spin system.

For large simulations, the primary time cost in CluE Oxide is often from diagonalizing and propagating the cluster Hamiltonians. In prototyping, a frequency-space method similar to that of [4] was tried, but likely due to the need to perform both a Fourier transform and inverse Fourier transform for each cluster, the time-domain integration of the Schrödinger equation is more efficient. The method CluE Oxide employ is to first diagonalize both electron spin-manifold Hamiltonians separately:

$$\hat{H}(m_{\rm S}) = \sum_{n} |\psi(m_{\rm S})\rangle E_n(m_{\rm S}) \langle \psi(m_{\rm S})|.$$
(1)

And to construct the propagators in the eigenbasis for a small time increment,  $\delta t$ .

$$\hat{U}(\delta t, m_{\rm S}) = \sum_{n} |\psi(m_{\rm S})\rangle \exp\left(-i\frac{E_n(m_{\rm S})\delta t}{\hbar}\right) \langle \psi(m_{\rm S})|.$$
 (2)

This means that each cluster Hamiltonian only needs to be diagonalized once, as the later time propagators can be found via matrix multiplication.

$$\hat{U}(n_t \delta t, m_S) = \hat{U}(\delta t, m_S)^{n_t}. \tag{3}$$

For some some of the longer time traces, it is helpful to have a small  $\delta t$  at early times and a longer  $\delta t$  at later times. The way CluE Oxide addresses this is allow multiple values of  $\delta t$  accompanied by the number of times each  $\delta t$  should be used. This requires that multiple propagators be calculated, but since the eigenvector have already been calculated, equation (2) can be used for multiple  $\delta t$ s for a single diagonalization step.

Next at each time point in the simulated experiment, a total pulse sequence propagator is calculated as

$$\hat{U}_{n_t, m_{\mathcal{S}}} = \mathscr{T} \prod_p \hat{U}(\delta t, m_{\mathcal{S}_p})^{n_t}, \tag{4}$$

where by using the  $\hat{U}(\delta t, m_{Sp})^{n_t-1}$  from the previous steps, the cost scaling with the number of timepoints is linear. of each  $n_t$  is roughly the same.

The Heisenberg picture detection operator,  $\hat{S}_{+}(t)$  in the high magnetic field limit, where the Zeeman basis is approximately the electron spin eigenbasis, can be written solely in terms of the propagators for the cluster in the as

$$\hat{S}_{+}(n_t) = \hat{U}_{n_t, m_{\rm S}'}^{\dagger} \hat{U}_{n_t, m_{\rm S}}.$$
 (5)

For square d by d matrices A and B, the inner product between them is

$$\langle A, B \rangle := \operatorname{tr}(A^{\dagger}B).$$
 (6)

To see why this definition makes sense, let n index the elements of the matrices. How exactly n indexes the matrices is unimportant as long as it is a consistent bijection between the  $d^2$  row column pairs and  $d^2$  integers. One possibility is that n = r + dc, where  $r, c \in [0, d-1]$  are the row and column indices respectfully. In element-wise summation, the trace is

$$\operatorname{tr}(A^{\dagger}B) = \sum_{r,c} A_{r,c}^* B_{c,r}. \tag{7}$$

And the inner product between vectors is

$$\langle A, B \rangle = \sum_{n} A_n^* B_n. \tag{8}$$

Since there is a bijection between n and (r, c), these expression are equivalent; however, if only the trace is desired and not the matrix AB itself, the inner product method requires fewer steps, and is how CluE Oxide evaluates  $\operatorname{tr}(\hat{\rho}\hat{S}_{+}(t))$ .

After all the cluster signals are calculated the CCE auxiliary signals and products can be constructed. By the nature of the product, when calculating the n-CCE signal, all the n'-CCE signals, n' < n are also calculated and saved.

If only a single orientation and isotopologue are requested, then CluE Oxide ends at this point. Different orientations require going back to the orientation generation step, and different isotopologues require rebuilding the system. After averaging over all the requested variations, CluE Oxide's primary returned data are the time domain signal n-CCE signal and the time axis. Depending on the input options other data may be saved, such as the clusters or individual signals from each orientation, the auxiliary signals, and system statistics including individual spin and methyl contributions.

# 0.2 Software

# 0.2.1 Installing

To start, please make sure that Rust is installed and up to date[5]. Before starting, it is a good idea to run the test suite.

```
1 cargo test
```

A few of the tests rely on random number generations and can occasionally fail. If this happens try running a second time: they should pass most of the time. To compile run the following.

```
1 cargo build --release
```

The compiled binary should be in *target/release/* and ready to use. An optional step is to make sure CluE Oxide is easy to access from the command line. One way to do this on Linux is make an alias by adding the following to the .bash aliases file.

```
1 alias clue_oxide="path/to/clue/target/release/clue_oxide"
```

#### 0.2.2 Running

To see the basic usage for CluE Oxide, run the following.

```
1 clue_oxide --help
```

The output is shown below.

```
clue_oxide input [-o "options"]
3
  Options:
      -h, --help
                        Prints help information.
6
      -H, --hide-title Do not display title information.
      -1, --license
                        Prints license.
      -0, --option
8
                        input additional options.
      -V, --version
                        Prints version information.
      -W, --warrenty
                        Prints warrenty information.
```

To run CluE Oxide from the command line, an input file is needed. The input file format is .txt file, and is supplied to CluE Oxide as shown below.

```
1 clue_oxide input.txt
```

# 0.2.3 Examples

Before detailing the syntax of the input file, here is an example input for reference.

```
Example CluE Oxide Input File
2
3
   */
4
   input_structure_file = "MD_300K_25ns-30ns_center_0001.pdb";
   detected_spin_position = centroid_over_serials([28,29]);
   detected_spin_g_matrix = [2.0097, 2.0064, 2.0025];
7
   detected_spin_g_y = diff(group(tempo_n) , group(tempo_o) );
   detected_spin_g_x = diff(group(tempo_c1) , group(tempo_c19) );
10
  radius = 25; // angstroms.
11
12
13
   pulse_sequence = hahn;
  magnetic_field = 1.2; // T.
14
15
16 number_timepoints = [101];
  time_increments = [5e-8];
17
18
19
  cluster_method = cce;
20 max_cluster_size = 4;
21
22
  neighbor_cutoff_3_spin_hahn_mod_depth = 3.23e-5;
23
   neighbor_cutoff_3_spin_hahn_taylor_4 = 1e17; // (rad/s)^4.
24
25
  orientation_grid = lebedev(170);
26
27
   #[group(tempo_h)]
28
     residues in [TEM];
     elements in [H];
29
30
  #[spin_properties(tempo_h, 1H)]
31
32
     tunnel_splitting = 80e3; // Hz.
33
34
  #[group(tempo_n)]
    residues in [TEM];
35
36
     elements in [N];
37
38
  #[group(tempo_o)]
39
    residues in [TEM];
40
     elements in [0];
41
42
   #[group(tempo_c1)]
43
     serials in [1];
44
45
  #[group(tempo_c19)]
46
     serials in [19];
47
48
  #[spin_properties(tempo_n, 14N)]
    hyperfine_coupling = [20,20,100]*1e6;
49
50
     hyperfine_x = diff(bonded(tempo_c1),bonded(tempo_c19));
51
     hyperfine_y = diff(particle,bonded(tempo_o));
52
     electric_quadrupole_coupling = [-0.28, -1.47, 1.75]*1e6;
53
     electric_quadrupole_x = diff(bonded(tempo_c1),bonded(tempo_c19));
54
     electric_quadrupole_y = diff(particle,bonded(tempo_o));
55
56
57
  #[config]
58 write_orientation_signals = "";
59 write_clusters = "";
```

The input script is not Turing complete, but is sufficient for starting CluE Oxide, and as will be discussed later, CluE Oxide can run from Turing complete languages like Bash, Python, and Rust. The syntax is inspired by C, C++, Rust, MATLAB and Python: all normal lines end in semicolons and double slashes indicate that the rest of line is a comment. Block commenting with "/\*" and "\*/" also works. Pound signs

indicate that the following lines are to be interpreted differently. Line 5 specifies the input structure as a PDB file. Line 7 places the detected electron between the nitrogen and oxygen of TEMPO. Line 8–10 specifies the electron g-matrix. Line 12 says to only use a 25 Å radius sphere of the PDB, applying periodic boundary conditions if need be. Line 14 specifies the pulse sequence to be simulated. Line 15 declares the applied magnetic field strength in tesla: Line 18 defines the time increments used evaluating propagators; for Carr-Purcell sequences with  $n_{\pi}$   $\pi$ -pulses, the time increment on the returned time-axis will be  $(n_{\pi}+1)\Delta t$ . the applied magnetic field is always in along the z-axis. Line 20 determines which cluster approximation to use, and line 21 specifies the maximum cluster size to be used. Lines 23 and 24 specify neighbor cutoffs for determining clusters. Line 27 selects a 170 point Lebedev grid[3]. Lines 29–31 declare a group selecting the TEMPO hydrogens.

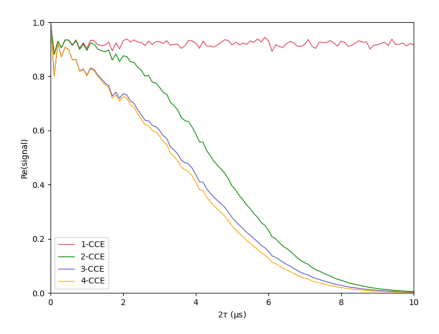


Figure 1: These Hahn echo simulation show the output of the previous example.

Each group is required to starts with a  $\#[group(my\_label)]$ , followed by various conditions; in this case the residue (as indicated in the PDB file), and the element. Line 33 and 34 set the methyl tunnel splitting for the TEMPO to 80 kHz, by referencing the previous group. This is done by using  $\#[spin\_properties(my\_label, my\_isotope)]$ , where  $my\_label$  matches the group selecting the TEMPO hydrogens. Lines 36–48 declare more groups. Lines 50–57 sets the TEMPO nitrogen's hyperfine tensor and electric quadrupole coupling tensor. Note that one group is used in  $\#[spin\_properties(tempo\_n, 14N)]$  to select the TEMPO nitrogen and other groups are used to specify atoms for orientating the coupling tensors. Line 59 sets the mode back to #[config]; there is an implied #[config] at the start of every CluE Oxide file. Line 60 and 61 turn off saving orientation dependent signals an clusters, respectively.

# 0.3 Input File

# 0.3.1 #[config]

The main section of the input file is #[config], and is the default when no mode is declared; however, the line

1 #[config]

will set CluE Oxide to read in #[config] mode and is needed to exit other modes back to #[config]. The available fields are explained below.

| clash_distance_pbc  clash distance_pbc  cluster batch size  cluster density_matrix  cluster_method  rzece, cce  detected_g_natrix  detected_g_x  vector  schedected_g_y  vector  spin coor  cluster_field  max_cluster_size  magnetic_field  max_cluster_size  magnetic_field  max_cluster_size  minimum_distance in \( \triangle \) for the detected spin g-matrix  shape of the spin system  method to calculate the density matrix of each  cluster;  method of evaluating clusters  eigenvalues or matrix of the detected spin g-matrix  and detected_g_x  vector   | field  | values     | description   |
|---|--|------------|---|
| clash_distance_pbe  cluster_batch_size  int  mumber of clusters to evaluate between saves method to calculate the density matrix of each cluster_method  r2cce, cee  detected_g_x  vector  ve   |  |            |   |
| cluster batch size   int   number of clusters to evaluate between saves   method to calculate the density matrix of each cluster   cluster   method to calculate the density matrix of each cluster   method of evaluating clusters   matrix  | clash distance pho   | float      | <del>-</del>  |
| cluster batch size  cluster_density_matrix  cluster_density_matrix  den_mat  cluster_method  r2ccc, ccc  detected_g_matrix  float]  detected_g_x  vector  vector  vector  vector z-direction of the detected spin g-matrix  detected_g_z  vector  vector  vector z-direction of the detected spin g-matrix  detected_g_z  vector  vector z-direction of the detected spin g-matrix  path to the input PDB  shape of the spin system  magnetic_field  max_cluster_size  int  magnetic_field strength in T of the applied magnetic field  max_cluster_size  int  magnetic_field strength in T of the applied magnetic field  max_cluster_size  int  magnetic field strength in T of the applied magnetic field  max_cluster_size  int  magnetic field strength in T of the applied magnetic field  max_cluster_size  int  magnetic field strength in T of the applied magnetic field  max_cluster_size  int  magnetic field strength in T of the applied magnetic field  max_cluster_size int  magnetic field strength in T of the applied magnetic field  max_cluster_size int  magnetic field strength in T of the applied magnetic field  max_cluster_size to be evaluated  minimum modulation depth for the three spin halm echo for an edge to be placed between two spins  minimum interpolate of the $\mathcal{O}(27^{\circ})^{\circ}$ coefficient, in the three spin halm echo Taylor series, $ k_{s} ^{s} _{s}$ for an edge to be placed between two spins  minimum $ k_{s} _{s} _{s} _{s} _{s} _{s}$ for an edge to be placed between two spins  minimum $ k_{s} _{s} _{s} _{s} _{s} _{s} _{s} _{s} $   |  | 11000      |   |
| cluster density matrix  cluster_method  r2ccc, ccc  detected_g_matrix  [float]  respectively  detected_g_x  vector  vedirection of the detected spin g-matrix  vector  vedirection of the detected spin g-matrix  vector  shape of the spin system  magnetic field  manumatic field  manumatic user size to be valuated  minimum langual in  | cluster batch size   | int        |   |
| custer_method   | Clustel_batell_size  | 1110       |   |
| cluster_method  | cluster_density_matrix   | den_mat    |   |
| detected $g$ matrix  detected $g$ x  vector  detected $g$ y  vector  vedirection of the detected spin g-matrix  valueffer  float  magnetic field  maximum cluster size to be valuated  minimum belap of the spin system  in the three spin hahe for the three spin hahe for spin spin spin spin spin spin spin spin   | cluster method   | r2cce, cce |   |
| $\begin{array}{c cccccd} \operatorname{detected} & \operatorname{matrix} & \operatorname{intoat} \\ \operatorname{detected} & \operatorname{g} & \operatorname{vector} \\ \operatorname{detected} & \operatorname{g} & \operatorname{vector} \\ \operatorname{detected} & \operatorname{g} & \operatorname{vector} \\ \operatorname{detected} & \operatorname{gr} & \operatorname{vector} \\ \operatorname{detected} & \operatorname{spin} & \operatorname{position} \\ \operatorname{input} & \operatorname{string} \\ \operatorname{load} & \operatorname{geometry} \\ \operatorname{magnetic} & \operatorname{field} \\ \operatorname{max} & \operatorname{cluster} & \operatorname{size} \\ \operatorname{magnetic} & \operatorname{field} \\ \operatorname{max} & \operatorname{cluster} & \operatorname{size} \\ \operatorname{meighbor} & \operatorname{cutoff} & \operatorname{geometry} \\ \operatorname{magnetic} & \operatorname{field} \\ \operatorname{max} & \operatorname{cluster} & \operatorname{size} \\ \operatorname{meighbor} & \operatorname{cutoff} & \operatorname{geometry} \\ \operatorname{meighbor} & \operatorname{cutoff} & \operatorname{geometry} \\ \operatorname{meighbor} & \operatorname{magnetic} & \operatorname{field} \\ \operatorname{max} & \operatorname{cluster} & \operatorname{size} \\ \operatorname{meighbor} & \operatorname{cutoff} & \operatorname{geometr} \\ \operatorname{meighbor} & \operatorname{geometr} \\ \operatorname{geometr} geo$ | <del></del>  |            |   |
| $\begin{array}{c ccccccccccccccccccccccccccccccccccc$   | detected_g_matrix  | [float]    |   |
| $\begin{array}{c ccccccccccccccccccccccccccccccccccc$   |  | vector     |   |
| $\begin{array}{c ccccccccccccccccccccccccccccccccccc$   |  | vector     |   |
| input_structure_file string cube, shape of the spin system magnetic_field float max_cluster_size int max_cluster_size int max_mum_cluster_size to be evaluated minimum modulation depth for the three spin hahn echo for an edge to be placed between two spins minimum $ b_{\perp} $ in the three spin hahn echo from the three spin hahn echo for an edge to be placed between two spins minimum $ b_{\perp} $ in the three spin hahn echo frame dege to be placed between two spins minimum $ b_{\perp} $ in the three spin hahn echo frame dege to be placed between two spins minimum $ b_{\perp} $ in the three spin hahn echo frame dege to be placed between two spins minimum $ b_{\perp} $ in the for an edge to be placed between two spins minimum $ b_{\perp} $ in the for an edge to be placed between two spins minimum $ b_{\perp} $ in the for an edge to be placed between two spins minimum $ b_{\perp} $ in the for an edge to be placed between two spins minimum $ b_{\perp} $ in the for an edge to be placed between two spins maximum distance in $A$ for an edge to be placed between two spins maximum distance in $A$ for an edge to be placed between two spins maximum distance in $A$ for an edge to be placed between two spins maximum distance in $A$ for an edge to be placed between two spins maximum distance in $A$ for an edge to be placed between two spins maximum distance in $A$ for an edge to be placed between two spins maximum distance in $A$ for an edge to be placed between two spins maximum distance in $A$ for an edge to be placed between two spins maximum distance in $A$ for an edge to be placed between two spins maximum distance in $A$ for an edge to be placed between two spins maximum distance in $A$ for an edge to be placed between two spins maximum distance in $A$ for an edge to be placed between two spins maximum distance in $A$ for an edge to be placed between two spins maximum distance in $A$ for an edge to be placed between two spins maximum distance in $A$ for an edge to be placed between two spins maximum distance in $A$ for an edge to be placed between two  | $\det \underline{\operatorname{detected}} \underline{\operatorname{g}} \underline{\operatorname{z}}$ | vector     | z-direction of the detected spin g-matrix               |
| load geometry         cube, sphere         shape of the spin system           magnetic_field         float         magnetic field strength in T of the applied magnetic field           max_cluster_size         int         maximum cluster size to be evaluated magnetic field           neighbor_cutoff_3_spin_hahn_mod_depth         float         minimum modulation depth for the three spin hahn echo for an edge to be placed between two spins           neighbor_cutoff_3_spin_hahn_taylor_4         float         minimum magnitude of the $\mathcal{O}((2\tau)^4)$ coefficient, in the three spin hahn echo Taylor series, $[kω^4]$ , in $(rad/s)^4$ , for an edge to be placed between two spins           neighbor_cutoff_coupling         float         minimum $[bz_z - 2J]$ in Hz for an edge to be placed between two spins           neighbor_cutoff_dielta_hyperfine         float         minimum $[ΔA_{nn}] := [A_{zz,n} - A_{zz,m}]$ in Hz for an edge to be placed between two spins           neighbor_cutoff_distance         float         minimum $[b_L]$ in Hz for an edge to be placed between two spins           number_timepoints         [int]         maximum distance in Å for an edge to be placed between two spins           number_system_instances         int         number of timepoints used for each time increment           number_system_instances         int         number of times the system will be simulated orientation_grid         ori_grid         ori_grid         ori_grid         ori_grid         ori_grid <t< td=""><td>detected_spin_position</td><td>spin_coor</td><td>coordinates in Å of the detected spin</td></t<>   | detected_spin_position   | spin_coor  | coordinates in Å of the detected spin                   |
| load geometry         cube, sphere         shape of the spin system           magnetic_field         float         magnetic field strength in T of the applied magnetic field           max_cluster_size         int         maximum cluster size to be evaluated magnetic field           neighbor_cutoff_3_spin_hahn_mod_depth         float         minimum modulation depth for the three spin hahn echo for an edge to be placed between two spins           neighbor_cutoff_3_spin_hahn_taylor_4         float         minimum magnitude of the $\mathcal{O}((2\tau)^4)$ coefficient, in the three spin hahn echo Taylor series, $[kω^4]$ , in $(rad/s)^4$ , for an edge to be placed between two spins           neighbor_cutoff_coupling         float         minimum $[bz_z - 2J]$ in Hz for an edge to be placed between two spins           neighbor_cutoff_dielta_hyperfine         float         minimum $[ΔA_{nn}] := [A_{zz,n} - A_{zz,m}]$ in Hz for an edge to be placed between two spins           neighbor_cutoff_distance         float         minimum $[b_L]$ in Hz for an edge to be placed between two spins           number_timepoints         [int]         maximum distance in Å for an edge to be placed between two spins           number_system_instances         int         number of timepoints used for each time increment           number_system_instances         int         number of times the system will be simulated orientation_grid         ori_grid         ori_grid         ori_grid         ori_grid         ori_grid <t< td=""><td>input structure file</td><td></td><td></td></t<>   | input structure file   |            |   |
| magnetic_field  minimum modulation depth for the three spin hahn cho for an edge to be placed between two spins  minimum magnitude of the $\mathcal{O}(2\tau)^4$ coeffi- cient, in the three spin hahn cho for an edge to be placed between two spins  minimum $b_{bz} - 2J$ in Hz for an edge to be placed between two spins  minimum $b_{bz} - 2J$ in Hz for an edge to be placed between two spins  neighbor_cutoff_delta_hyperfine  float  minimum $b_{bz} - 2J$ in Hz for an edge to be placed between two spins  minimum $b_{bz} - 2J$ in Hz for an edge to be placed between two spins  minimum $b_{bz} - 2J$ in Hz for an edge to be placed between two spins  minimum be_i in Hz for an edge to be placed between two spins  minimum be_i in Hz for an edge to   | <del>-</del>   | _          |   |
| magnetic field  max_cluster_size  int  maximum cluster size to be evaluated  max_cluster_size  int  maximum cluster size to be evaluated  minimum modulation depth for the three spin hahn echo for an edge to be placed between two spins  minimum magnitude of the $\mathcal{O}((2\tau)^4)$ coefficient, in the three spin hahn echo Taylor series, $ k\omega^4 $ , in $(rad/s)^4$ , for an edge to be placed between two spins  neighbor_cutoff_coupling  float  neighbor_cutoff_delta_hyperfine  neighbor_cutoff_dipole_perpendicular  neighbor_cutoff_dipole_perpendicular  float  float  float  float  float  float  float  maximum $ b_\perp $ in $ a_{2,n} - a_{2,z,m} $ in $ $   | load_geometry  |            | shape of the spin system                                |
| magnetic field  max_cluster_size  int  maximum cluster size to be evaluated  max_cluster_size  int  maximum cluster size to be evaluated  minimum modulation depth for the three spin hahn echo for an edge to be placed between two spins  minimum magnitude of the $\mathcal{O}((2\tau)^4)$ coefficient, in the three spin hahn echo Taylor series, $ k\omega^4 $ , in $(rad/s)^4$ , for an edge to be placed between two spins  neighbor_cutoff_coupling  float  neighbor_cutoff_delta_hyperfine  neighbor_cutoff_dipole_perpendicular  neighbor_cutoff_dipole_perpendicular  float  float  float  float  float  float  float  maximum $ b_\perp $ in $ a_{2,n} - a_{2,z,m} $ in $ $   |  | 0 4        | magnetic field strength in T of the applied             |
| max_cluster_size int maximum cluster size to be evaluated minimum modulation depth for the three spin hahn echo for an edge to be placed between two spins minimum magnitude of the $\mathcal{O}((2\tau)^4)$ coefficient, in the three spin hahn echo Taylor series, $ k\omega^4 $ , in $(rad/s)^4$ , for an edge to be placed between two spins neighbor_cutoff_coupling float minimum $ b_{2z}-2J $ in Hz for an edge to be placed between two spins minimum $ b_{2z}-2J $ in Hz for an edge to be placed between two spins minimum $ \Delta A_{mn}  :=  A_{zz,n} - A_{zz,m} $ in Hz for an edge to be placed between two spins minimum $ b_{\perp} $ in Hz for an edge to be placed between two spins minimum $ b_{\perp} $ in Hz for an edge to be placed between two spins minimum $ b_{\perp} $ in Hz for an edge to be placed between two spins minimum $ b_{\perp} $ in Hz for an edge to be placed between two spins number_cutoff_distance float maximum distance in $A$ for an edge to be placed between two spins number_system_instances int number of timepoints used for each time increment number_system_instances int number of times the system will be simulated orientation_grid ori_grid grid used for orientation averaging pulse_sequence $ a_{\perp} $ pulse sequence to simulate with $n_{\pi}$ $\pi$ -pulses radius float from the average position of the detected spin to use save_dir time_increments [float] list of $\Delta \tau$ s in s used for propagation file name to save cluster auxiliary signals under write_auxiliary_signals write_lauxiliary_signals write_lauxilia   | magnetic_field   | поат       |   |
| meighbor_cutoff_3_spin_hahn_mod_depth float minimum modulation depth for the three spin hahn echo for an edge to be placed between two spins minimum magnitude of the $\mathcal{O}((2\tau)^4)$ coefficient, in the three spin hahn echo Taylor series, $ k\omega^4 $ , in (rad/s) $^4$ , for an edge to be placed between two spins minimum $ \Delta_{zz} - 2J $ in Hz for an edge to be placed between two spins minimum $ \Delta_{Ann}  :=  A_{zz,n} - A_{zz,m} $ in Hz for an edge to be placed between two spins minimum $ \Delta_{Ann}  :=  A_{zz,n} - A_{zz,m} $ in Hz for an edge to be placed between two spins minimum $ \Delta_{Ann}  :=  A_{zz,n} - A_{zz,m} $ in Hz for an edge to be placed between two spins minimum $ \Delta_{Ann}  :=  A_{zz,n} - A_{zz,m} $ in Hz for an edge to be placed between two spins minimum $ \Delta_{Ann}  :=  A_{zz,n} - A_{zz,m} $ in Hz for an edge to be placed between two spins minimum $ \Delta_{Ann}  :=  A_{zz,n} - A_{zz,m} $ in Hz for an edge to be placed between two spins minimum $ \Delta_{Ann}  :=  A_{zz,n} - A_{zz,m} $ in Hz for an edge to be placed between two spins minimum $ \Delta_{Ann}  :=  A_{zz,n} - A_{zz,m} $ in Hz for an edge to be placed between two spins minimum $ \Delta_{Ann}  :=  A_{zz,n} - A_{zz,m} $ in Hz for an edge to be placed between two spins minimum $ \Delta_{Ann}  :=  A_{zz,n} - A_{zz,m} $ in Hz for an edge to be placed between two spins minimum $ \Delta_{Ann}  :=  A_{zz,n} - A_{zz,m} $ in Hz for an edge to be placed between two spins minimum $ \Delta_{Ann}  :=  A_{zz,n} - A_{zz,m} $ in Hz for an edge to be placed between two spins minimum $ \Delta_{Ann}  :=  A_{zz,n} - A_{zz,m} $ in Hz for an edge to be placed between two spins minimum $ \Delta_{Ann}  :=  A_{zz,n} - A_{zz,m} $ in Hz for an edge to be placed between two spins minimum $ \Delta_{Ann}  :=  A_{zz,n} - A_{zz,m} $ in Hz for an edge to be placed between two spins minimum $ \Delta_{Ann}  :=  A_{zz,n} - A_{zz,m} $ in Hz for an edge to be placed between two spins minimum $ \Delta_{Ann}  :=  A_{zz,n} - A_{zz,m} $ in Hz for an edge to be placed between two spins minimum $ \Delta_{Ann}  :=  A_{zz,n} - A_{zz,m} $ in Hz for an edge to be  | max cluster size   | int        |   |
| neighbor_cutoff_3_spin_hahn_mod_depth neighbor_cutoff_3_spin_hahn_taylor_4 float when two spins minimum magnitude of the $\mathcal{O}((2\tau)^4)$ coefficient, in the three spin hahn echo Taylor series, $ k\omega^4 $ , in $(\operatorname{rad/s})^4$ , for an edge to be placed between two spins neighbor_cutoff_coupling float minimum $ b_{zz} - 2J $ in Hz for an edge to be placed between two spins neighbor_cutoff_delta_hyperfine float minimum $ \Delta A_{mn}  :=  A_{zz,n} - A_{zz,m} $ in Hz for an edge to be placed between two spins neighbor_cutoff_dipole_perpendicular float minimum $ \Delta A_{mn}  :=  A_{zz,n} - A_{zz,m} $ in Hz for an edge to be placed between two spins neighbor_cutoff_distance float maximum distance in $A$ for an edge to be placed between two spins maximum distance in $A$ for an edge to be placed between two spins number_string float placed between two spins number_string float grid used for orientation averaging cp- $n_{\pi}$ , hahn pulse sequence from the average position of the detected spin to use save_dir string directory where data will be saved file name to save cluster auxiliary signals under write_auxiliary_signals string file name to save cluster auxiliary signals under write_clusters string file name to save custers under directory name to save general information under string file name to save exchange groups (such as methyls) under   |  |            | minimum modulation depth for the three spin             |
| meighbor_cutoff_3_spin_hahn_taylor_4 float minimum magnitude of the $\mathcal{O}((2\tau)^4)$ coefficient, in the three spin hahn echo Taylor series, $ k\omega^4 $ , in $(\operatorname{rad/s})^4$ , for an edge to be placed between two spins minimum $ b_{zz}-2J $ in Hz for an edge to be placed between two spins neighbor_cutoff_delta_hyperfine float minimum $ \Delta A_{mn}  :=  A_{zz,n} - A_{zz,m} $ in Hz for an edge to be placed between two spins neighbor_cutoff_dipole_perpendicular float minimum $ b_\perp $ in Hz for an edge to be placed between two spins minimum $ b_\perp $ in Hz for an edge to be placed between two spins neighbor_cutoff_distance float maximum distance in $\bar{A}$ for an edge to be placed between two spins number_timepoints [int] number of timepoints used for each time increment number_system_instances in the number of times the system will be simulated orientation_grid ori_grid grid used for orientation averaging pulse_sequence contains a string directory where data will be saved time_increments float list of $\Delta \tau$ s in s used for propagation file name to save cluster auxiliary signals under write_auxiliary_signals string file name to save cluster auxiliary signals under write_clusters string file name to save exchange groups (such as methyls) under  | neighbor cutoff 3 spin hahn mod depth  | float      |   |
| minimum magnitude of the $\mathcal{O}((2\tau)^4)$ coefficient, in the three spin hahn echo Taylor series, $ k\omega^4 $ , in $(rad/s)^4$ , for an edge to be placed between two spins  neighbor_cutoff_coupling  float  neighbor_cutoff_delta_hyperfine  neighbor_cutoff_dipole_perpendicular  neighbor_cutoff_dipole_perpendicular  neighbor_cutoff_distance  neighbor_cutoff_distance  neighbor_cutoff_distance  float  | noignooi_outon_o_spin_noin_inou_uopen  | 11000      |   |
| neighbor_cutoff_3_spin_hahn_taylor_4  float   |  |            |   |
| neighbor_cutoff_aspin_nann_taylor_4  neighbor_cutoff_coupling  float  float  minimum $ b_{zz} - 2J $ in Hz for an edge to be placed between two spins  minimum $ \Delta A_{mn}  :=  A_{zz,n} - A_{zz,m} $ in Hz for an edge to be placed between two spins  meighbor_cutoff_dipole_perpendicular  float  flo  |  |            |   |
| tween two spins $\begin{array}{c} \text{meighbor\_cutoff\_coupling} & \text{float} & \text{minimum} \mid b_{zz} - 2J \mid \text{in Hz for an edge to be placed between two spins} \\ \text{meighbor\_cutoff\_delta\_hyperfine} & \text{float} & \text{minimum} \mid \Delta A_{mn} \mid := \mid A_{zz,n} - A_{zz,m} \mid \text{in Hz for an edge to be placed between two spins} \\ \text{meighbor\_cutoff\_dipole\_perpendicular} & \text{float} & \text{minimum} \mid b_{\perp} \mid \text{in Hz for an edge to be placed between two spins} \\ \text{meighbor\_cutoff\_distance} & \text{float} & \text{maximum distance in Å for an edge to be placed between two spins} \\ \text{number\_timepoints} & \text{int} & \text{maximum distance in Å for an edge to be placed between two spins} \\ \text{number\_system\_instances} & \text{int} & \text{number of timepoints used for each time increment} \\ \text{number\_system\_instances} & \text{int} & \text{number of times the system will be simulated orientation\_grid} & \text{ori\_grid} & \text{grid used for orientation averaging} \\ \text{cp-}n_{\pi}, & \text{pulse sequence to simulate with } n_{\pi} \pi \text{-pulses} \\ \text{radius} & \text{float} & \text{radius in Å from the average position of the detected spin to use} \\ \text{save\_dir} & \text{string} & \text{directory where data will be saved} \\ \text{time\_increments} & \text{[float]} & \text{list of } \Delta \tau \text{s in s used for propagation} \\ \text{file name to save cluster auxiliary signals under} \\ \text{write\_bath} & \text{string} & \text{file name to save bath spins under} \\ \text{write\_lusters} & \text{string} & \text{file name to save general information} \\ \text{under} & \text{write\_info} & \text{string} & \text{file name to save exchange groups (such as methyls) under} \\ \end{array}$   | neighbor_cutoff_3_spin_hahn_taylor_4   | float      |   |
| neighbor_cutoff_couplingfloatminimum $ b_{zz}-2J $ in Hz for an edge to be placed between two spinsneighbor_cutoff_delta_hyperfinefloat $\min  \Delta A_{mn}  :=  A_{zz,n} - A_{zz,m} $ in Hz for an edge to be placed between two spinsneighbor_cutoff_dipole_perpendicularfloat $\min  \Delta A_{mn}  :=  A_{zz,n} - A_{zz,m} $ in Hz for an edge to be placed between two spinsneighbor_cutoff_distancefloat $\min  \Delta A_{mn}  :=  A_{zz,n} - A_{zz,m} $ in Hz for an edge to be placed between two spinsnumber_cutoff_distancefloat $\min  \Delta A_{mn}  :=  A_{zz,n} - A_{zz,m} $ in Hz for an edge to be placed between two spinsnumber_cutoff_distancefloat $\max  \Delta A_{mn}  :=  A_{zz,n} - A_{zz,m} $ in Hz for an edge to be placed between two spinsnumber_cutoff_distancefloat $\max  \Delta A_{mn}  :=  A_{zz,n} - A_{zz,m} $ in Hz for an edge to be placed between two spinsnumber_cutoff_distancefloat $\max  \Delta A_{mn}  :=  A_{zz,n} - A_{zz,m} $ in Hz for an edge to be placed between two spinsnumber_cutoff_distancefloat $\max  \Delta A_{mn}  :=  A_{zz,n} - A_{zz,m} $ in Hz for an edge to be placed between two spinsnumber_dutoff_distancefloat $\max  \Delta A_{mn}  :=  A_{zz,n} - A_{zz,m} $ in Hz for an edge to be placed between two spinsnumber_dutoff_distancefloatnumber of time the zore distancepulse_sequencefile name to save eluster with $n_n \pi$ -pulsesradius_sequencefloatlist of $\Delta T$ in sused for propagationstring_distancefile name to save clusters underwrite_bathstring_directory_name to save general information underwrite_exchange_groupsstring_directory_name to save exchange   |  |            |   |
| placed between two spins  neighbor_cutoff_delta_hyperfine  float   |  |            |   |
| neighbor_cutoff_delta_hyperfinefloat $\min \min  \Delta A_{mn}  :=  A_{zz,n} - A_{zz,m}  $ in Hz for an edge to be placed between two spinsneighbor_cutoff_dipole_perpendicularfloat $\min \lim_{ b_{\perp} } \ln Hz$ for an edge to be placed between two spinsneighbor_cutoff_distancefloat $\max \lim_{ a  \to a }  a  =  A_{zz,n} - A_{zz,m}  $ in Hz for an edge to be placed between two spinsnumber_ctimepointsfloat $\max \lim_{ a  \to a }  a  =  A_{zz,n} - A_{zz,m}  $ in Hz for an edge to be placed between two spinsnumber_ctimepoints $\max \lim_{ a  \to a }  a  =  A_{zz,n} - A_{zz,m}  $ in Hz for an edge to be placed between two spinsnumber_ctimepoints $\min \lim_{ a  \to a }  a  =  A_{zz,n} - A_{zz,m}  $ in Hz for an edge to be placed between two spinsnumber_stimepoints $\min \lim_{ a  \to a }  a  =  A_{zz,n} - A_{zz,m}  $ in Hz for an edge to be placed between two spinsnumber_stime_number_stimes $\min \lim_{ a  \to a }  a  =  A_{zz,n} - A_{zz,m}  $ in Hz for an edge to be placed between two spinsnumber_of times the system will be two spinsnumber of times the system will be simulated orientation averagingpulse_sequence $\min \lim_{ a  \to a }  a  =  a$  | neighbor_cutoff_coupling   | float      | minimum $ 0_{zz} - 2J $ in Hz for an edge to be         |
| neighbor_cutoff_dipole_perpendicular float $  harmonterm{minimum}   h_{\perp}                                      $  |  |            |   |
| neighbor_cutoff_dipole_perpendicular  float  minimum $ b_{\perp} $ in Hz for an edge to be placed between two spins  minimum $ b_{\perp} $ in Hz for an edge to be placed between two spins  maximum distance in Å for an edge to be placed between two spins  maximum distance in Å for an edge to be placed between two spins  number_timepoints  [int]  number of timepoints used for each time increment  number_system_instances  int  number of times the system will be simulated orientation_grid  ori_grid  grid used for orientation averaging  pulse_sequence $cp-n_{\pi}$ , hahn  pulse sequence to simulate with $n_{\pi}$ $\pi$ -pulses  float  float  float  radius in Å from the average position of the detected spin to use  save_dir  time_increments  [float]  list of $\Delta \tau$ s in s used for propagation  file name to save cluster auxiliary signals under  write_auxiliary_signals  string  file name to save bath spins under  write_clusters  write_info  write_exchange_groups  string  file name to save exchange groups (such as methyls) under  | neighbor cutoff delta hyperfine  | float      |   |
| heighbor_cutoff_distance   he maximum distance in Å for an edge to be placed between two spins   heighbor_cutoff_distance   heig   |  |            |   |
| neighbor_cutoff_distance float maximum distance in Å for an edge to be placed between two spins number_timepoints [int] mumber of timepoints used for each time increment number_system_instances int number of times the system will be simulated orientation_grid ori_grid grid used for orientation averaging pulse_sequence   | neighbor cutoff dipole perpendicular   | float      | ' '   |
| neighbor_cutorr_distance placed between two spins number_timepoints   | 0 = 1 = 1  |            |   |
| number_timepoints $ [int]                                    $  | neighbor cutoff distance   | float      |   |
| number_timepoints     int   |  |            |   |
| number_system_instances int number of times the system will be simulated orientation_grid ori_grid grid used for orientation averaging pulse_sequence $ \begin{array}{c} \operatorname{cp-}n_{\pi}, \\ \operatorname{hahn} \end{array} $ pulse sequence to simulate with $n_{\pi}$ $\pi$ -pulses radius $ \begin{array}{c} \operatorname{radius} & \operatorname{float} \end{array} $ radius in Å from the average position of the detected spin to use $ \operatorname{save\_dir} $ string directory where data will be saved $ \operatorname{time\_increments} $ [float] list of $\Delta \tau s$ in s used for propagation $ \operatorname{write\_auxiliary\_signals} $ string file name to save cluster auxiliary signals under write\_clusters $ \operatorname{string}  $ file name to save clusters under $ \operatorname{write\_info} $ string file name to save general information under $ \operatorname{write\_info} $ string file name to save exchange groups (such as methyls) under $ \operatorname{write\_exchange\_groups} $   | number timepoints  | [int]      | number of timepoints used for each time in-             |
| orientation_grid ori_grid grid used for orientation averaging pulse_sequence $ \begin{array}{c} \text{cp-}n_{\pi}, \\ \text{hahn} \end{array}  \text{pulse sequence to simulate with } n_{\pi} \; \pi\text{-pulses} \\ \text{radius}  \text{float}  \text{radius in Å from the average position of the detected spin to use} \\ \text{save\_dir}  \text{string}  \text{directory where data will be saved} \\ \text{time\_increments}  [\text{float}]  \text{list of $\Delta \tau \text{s in s used for propagation}} \\ \text{write\_auxiliary\_signals}  \text{string}  \text{file name to save cluster auxiliary signals under} \\ \text{write\_bath}  \text{string}  \text{file name to save bath spins under} \\ \text{write\_clusters}  \text{string}  \text{file name to save clusters under} \\ \text{write\_info}  \text{string}  \text{file name to save general information under} \\ \text{write\_exchange\_groups}  \text{string}  \text{file name to save exchange groups (such as methyls) under} \\ \end{array}$  |  | [2220]     |   |
| pulse_sequence $ \begin{array}{c} \operatorname{cp-}n_{\pi}, \\ \operatorname{hahn} \end{array}  \text{pulse sequence to simulate with } n_{\pi} \; \pi\text{-pulses} \\ \operatorname{radius} \qquad \qquad$  |  |            | v   |
| radius float radius in Å from the average position of the detected spin to use save_dir string directory where data will be saved list of $\Delta \tau$ s in s used for propagation write_auxiliary_signals string file name to save cluster auxiliary signals under write_clusters string file name to save clusters under file name to save clusters under write_info string file name to save general information under write_exchange_groups string file name to save exchange groups (such as methyls) under   | orientation_grid   | ori_grid   | grid used for orientation averaging                     |
| radius float radius in Å from the average position of the detected spin to use save_dir string directory where data will be saved time_increments [float] list of $\Delta \tau$ s in s used for propagation write_auxiliary_signals string file name to save cluster auxiliary signals under write_bath string file name to save bath spins under write_clusters string file name to save clusters under directory name to save general information under write_info string file name to save exchange groups (such as methyls) under   | pulse sequence   |            | pulse sequence to simulate with $n_{\pi}$ $\pi$ -pulses |
| $\begin{array}{cccccccccccccccccccccccccccccccccccc$  | paise_coquence   | hahn       |   |
| $\begin{array}{c ccccccccccccccccccccccccccccccccccc$   | radius   | float      |   |
| time_increments       [float]       list of $\Delta \tau s$ in s used for propagation         write_auxiliary_signals       string       file name to save cluster auxiliary signals under         write_bath       string       file name to save bath spins under         write_clusters       string       file name to save clusters under         write_info       string       directory name to save general information under         write_exchange_groups       string       file name to save exchange groups (such as methyls) under  |  |            |   |
| write_auxiliary_signals  string  file name to save cluster auxiliary signals under  write_bath  write_clusters  string  file name to save bath spins under  file name to save clusters under  file name to save general information  under  write_exchange_groups  string  file name to save cluster auxiliary signals under  file name to save bath spins under  file name to save clusters under  directory name to save general information  under  file name to save exchange groups (such as methyls) under  |  | _          |   |
| write_auxmary_signals  write_bath  write_clusters  string  file name to save bath spins under  file name to save clusters under  directory name to save general information  under  write_exchange_groups  string  file name to save exchange groups (such as methyls) under  | time_increments  | [float]    |   |
| write_bath string file name to save bath spins under write_clusters string file name to save clusters under write_info string directory name to save general information under write_exchange_groups string file name to save exchange groups (such as methyls) under   | write auxiliary signals  | string     |   |
| write_clusters string file name to save clusters under write_info string directory name to save general information under  write_exchange_groups string file name to save exchange groups (such as methyls) under   |  |            |   |
| write_info string directory name to save general information under write_exchange_groups string file name to save exchange groups (such as methyls) under   | _  |            |   |
| write_inio under  write_exchange_groups string under  file name to save exchange groups (such as methyls) under   | write_clusters   | string     |   |
| write_exchange_groups  string  under  file name to save exchange groups (such as methyls) under   | write_info   | string     |   |
| write_exchange_groups string methyls) under   |  |            |   |
| metnyis) under  | write exchange groups  | string     | `   |
| write_structure_pdb string PDB file name to save the spin system under  |  | 5011118    | · /   |
|   | write_structure_pdb  | string     | PDB file name to save the spin system under             |

#### ClusterDensityMatrix

If no  $cluster\_density\_matrix$  is specified, the infinite temperature limit is assumed for the bath spin, so that the density matrix is proportional to the identity matrix. To specify a particular temperature, T, in kelvin, use

```
cluster_density_matrix = thermal(T);
```

## ClusterMethod

There are two cluster evolutions schemes available in CluE Oxide, cce, and r2cce. The cce method implements ensemble CCE [1, 2] and has no explicit limit on the maximum cluster size. The r2cce method is a restricted version of CCE, using the analytic solution to the simplified three spin model from [6], and is therefore limited to a cluster size of two.

```
1 temperature = 20; // K.
```

This will switch CluE Oxide from using the identity matrix to the thermal density matrix.

# **DetectedSpinCoordinates**

The detected spin coordinates have several ways to specified. They can be specified directly.

```
1 detected_spin_position = [1.2, -3, 0];
```

They can be specified as the centroid of atoms in the PDB file.

```
detected_spin_position = centroid_over_serials([28,29]);
```

Both of these methods assume that the detected spin is localized to a single point. If the spin is delocalized of multiple points, a csv file can be used to provide  $|\Psi(x,y,z)|^2$ .

```
1 detected_spin_position = read_csv("xyzw.csv");
```

Each row of the file "xyzw.csv" should be four comma separated floating point numbers, specifying the x, y, z and weight,  $w = |\Psi(x, y, z)|^2 dx dy dz$ , for the coordinates.

## LoadGeometry

The  $load\_geometry$  can be either sphere, which is the default, or cube. The  $sphere\ load\_geometry$  includes only bath particle that are within the specified radius of the detected spin. The  $cube\ load\_geometry$  includes all bath particles from all periodic boundary duplicates. Periodic boundary duplicates are constructed as rectangular prism where there are 1 + 2ceil(r/a) cell along each dimension, where r is the radius and a in the cell edge length along the dimension in question.

# OrientationGrid

The orientation grid specifies the directions and weights for the applied static magnetic field. If no grid is given, the magnetic field is directed along the z-axis of the PDB file. A lebedev grid [3] can be specified.

```
1 orientation_grid = lebedev(302);
```

A random grid is also an option. Here is an example of using a single random direction.

```
1 orientation_grid = random(1);
```

And as with the detected spin position, a csv file x, y, z and weight entries can be used for a custom grid.

```
1 orientation_grid = read_csv("xyzw.csv");
```

#### PulseSequence

To simulate a Carr-Purcell pulse sequence with two  $\pi$ -pulses

```
1 pulse_sequence = cp-2;
```

A Hahn echo can be selected as shown.

```
pulse_sequence = hahn;
```

Or a Hahn echo is also a Carr-Purcell pulse sequence with one  $\pi$ -pulse.

```
pulse_sequence = cp-1;
```

## SystemInstance

For simulations with random elements, such as a randomized isotope distribution, running the over multiple instances can be important. For example to run the simulation 1000 times each with a random direction, use the following.

```
1 number_system_instances = 1000;
2 orientation_grid = random(1);
```

Note that this is different from the following.

```
number_system_instances = 1;
orientation_grid = random(1000);
```

The former chooses a random isotopologue and a random direction 1000 times, while the latter uses a single random isotopologue but 10000 random directions. As an in between, the following chooses 900 random isotopologues, and for each of them chooses 100 random orientations.

```
number_system_instances = 900;
orientation_grid = random(100);
```

#### **Tensor Specification**

Tensors such as  $detected\_g\_matrix$  or various bath spin coupling tensors are specified with an array of floating point numbers. Isotropic matrices can be specified with a single number, as an unwritten identity matrix is assumed.

```
1 detected_g_matrix = 2;
```

The eigenvalues can be specified by an array of three numbers; however, two eigenvectors are also needed under this method. See the section on *VectorSpecifiers* for details. Since CluE Oxide only uses symmetric tensors, six value can be used to specify the upper triangular part of a tensor, and the remaining values will be inferred.

Since the numerical values on the right hand side are for illustrative purposes only,  $some\_matrix$  is being used as a generic name. Finally, nine values can be used to specify a tensor; however, the redundant entries in the lower triangular section of the matrix are discarded. As such the following example is equivalent to the previous example.

```
1 some_matrix = [1, 2, 3, 4. 5. 6. 7, 8, 9];
```

CluE Oxide will interpret both the six and nine value examples as

$$\begin{bmatrix} 1 & 2 & 3 \\ 2 & 5 & 6 \\ 3 & 6 & 9 \end{bmatrix}$$

in the frame of the input PDB. When orientation averaging, CluE Oxide automatically rotates the tensors as well.

#### **TimeAxis**

The time-axis is specified in two part, the number of timepoints, and the time increments. For example, the CluE Oxide input file could include the following.

```
number_timepoints = [101,90];
time_increments = [5,50]*1e-9; // seconds.
```

CluE Oxide will read this and use  $\Delta t = 5$  ns for the first 101 timepoints, and then  $\Delta t = 50$  ns for the remaining 90 timepoints. There will be 191 timepoints total. Note that these are the  $\Delta t$ s for the smallest propagator unit, so a Carr-Purcell pulse sequence with  $n_{\pi}$   $\pi$ -pulses will have a  $\Delta t_{\text{axis}} = (n_{\pi} + 1)\Delta t$ . So if the above example is for a Hahn echo  $(n_{\pi} = 1)$ ,  $\Delta t_{\text{axis}} = 10$  ns for the first 101 timepoints  $\in [0, 1]$   $\mu$ s and then  $\Delta t_{\text{axis}} = 100$  ns for the next 90 timepoints  $\in [1.1, 10]$   $\mu$ s.

# VectorSpecifiers

The fields that require values of "vector" have several options. First, a vector can be a list.

```
1 detected_g_x = [1,0,0];
2 detected_g_z = [0,0,1];
```

Note that exactly two axes should be specified: the third axis is found with the appropriate cross product. The specified axes do not need to be normalized or perfectly orthogonal, since CluE Oxide will orthonormalize the vectors itself. The reason for this is allow for directions to be specified by properties of the system. The following lines say to use the difference between the atoms in the  $group(tempo\_c1)$ , and the atoms in  $group(tempo\_c1)$  for the x-direction, and to use difference between the atoms in the  $group(tempo\_c1)$ , and the atoms in  $group(tempo\_o)$  for the y-direction. Filters are explained in the next section, but the union of the atoms within the diff() must contain exactly two atoms for a unique vector to be specified.

```
detected_g_x = diff(group(tempo_c1) , group(tempo_c19));;
detected_g_y = diff(group(tempo_n) , group(tempo_o));;
```

# 0.3.2 #[group(my\_label)]

The options file can have zero, one, or multiple  $\#[group(my\_label)]$  sections. Each  $\#[group(my\_label)]$  section must have a unique label, by replacing  $my\_label$  with the desired label. Under the  $\#[group(my\_label)]$  header, the group conditions that can have multiple matches are listed as

```
1 property in [allowed_value_1, allowed_value_2, ... allowed_value_n];
```

or

```
property not in [forbidden_value_1, forbidden_value_2, ... forbidden_value_n];
```

The distance group only takes maximum and minimum values and is specified as

```
distance <= r_max; // angstroms
distance >= r_min; // angstroms
```

where  $r\_max$  and  $r\_min$  are floating point numbers specifing the maximum and minimum distances in Å from the detected spin that the particles in the group can be. CluE Oxide will find all the spins that fit all specified criteria. Note that depending upon how the criteria are defined it is possible that some spins will fall under multiple groups. CluE Oxide treats this case as an error and will indicate as such. To resolve this error, the group criteria should be refined. The following table shows the possible group criteria.

| field                           | values  | description  |
|---------------------------------|---------|--|
| bonded_elements                 | [H,He,] | elements of bonded atoms to match (in) or avoid (not in)                               |
| bonded_residues                 | [GLY,]  | residues of bonded atoms as specified in the PDB file to match (in) or avoid (not in)  |
| bonded_serials                  | [int]   | PDB serial numbers of bonded atoms to match (in) or avoid (not in)                     |
| bonded_residue_sequence_numbers | [int]   | PDB residues sequence numbers of bonded atoms to match (in) or avoid (not in)          |
| distance                        | float   | maximum (<=) or minimum (>=) allowed distance in Å from the detected spin              |
| elements                        | [H,He,] | elements to match (in) or avoid (not in)   |
| extracells                      | none    | restrict to unit cells other than the primary cell                                     |
| primary_cell                    | none    | restrict to the primary unit cell  |
| residues                        | [GLY,]  | residues as specified in the PDB file to match (in) or avoid (not in)                  |
| residue_sequence_numbers        | [int]   | residues sequence numbers as specified in the PDB file to match (in) or avoid (not in) |
| serials                         | [int]   | serial numbers as specified in the PDB file to match (in) or avoid (not in)            |

# $0.3.3 \quad \#[structure\_properties(my\_label)]$

The options file can have zero, one, or multiple  $\#[structure\_properties(my\_label)]$  sections, where  $my\_label$  should match to a  $\#[group(my\_label)]$ . The properties specified under  $\#[structure\_properties(my\_label)]$  will be applied to all spins that fall under the corresponding  $\#[group(my\_label)]$ . The following table shows the available properties.

| field              | values                | description                               |
|--------------------|-----------------------|---|
| isotope_abundances | {1H: 0.9,<br>2H: 0.1} | possible isotopes and probabilities       |
| void_probability   | float                 | chance that each particle will be removed |

# $0.3.4 \quad \#[\text{spin\_properties}(\text{my\_label}, \, \text{my\_isotope})]$

The options file can have zero, one, or multiple  $\#[spin\_properties(my\_label, my\_isotope)]$  sections, where  $my\_label$  should match to a  $\#[group(my\_label)]$ . The properties specified under  $\#[spin\_properties(my\_label, my\_isotope)]$  will be applied to all spins of isotope  $my\_isotope$  that fall under the corresponding  $\#[group(my\_label)]$ . The reason to differentiate between  $\#[spin\_properties(my\_label, my\_isotope)]$  and  $\#[structure\_properties(my\_label)]$  is isotope specificity:  $\#[structure\_properties(my\_label)]$  applies to all particles in the corresponding group, while  $\#[spin\_properties(my\_label, my\_isotope)]$  only applies to the specified isotope in the corresponding group. The following table shows the available properties.

| field                        | values  | description                                       |
|------------------------------|---------|---|
| active                       | bool    | specifies if the particle should be used for spin |
|                              |         | dynamics  |
| electric quadrupole coupling | [float] | eigenvalues in Hz of the electric quadrupole      |
| electric_quadrupole_coupling |         | tensor  |
| electric_quadrupole_x        | vector  | x-direction of the electric quadrupole tensor     |
| electric_quadrupole_y        | vector  | y-direction of the electric quadrupole tensor     |
| electric_quadrupole_z        | vector  | z-direction of the electric quadrupole tensor     |
| g_matrix                     | [float] | eigenvalues in of the g-matrix                    |
| g_x                          | vector  | x-direction of the g-matrix                       |
| g_y                          | vector  | y-direction of the g-matrix                       |
| g_z                          | vector  | z-direction of the g-matrix                       |
| hyperfine_coupling           | [float] | eigenvalues in Hz of the hyperfine tensor         |
| hyperfine_x                  | vector  | x-direction of the hyperfine tensor               |
| hyperfine_y                  | vector  | y-direction of the hyperfine tensor               |
| hyperfine_z                  | vector  | z-direction of the hyperfine tensor               |
| tunnel_splitting             | float   | tunnel splitting in Hz for particle in an ex-     |
|                              |         | change group                                      |

# VectorSpecifiers

Two tensor axes must be specified in addition to the coupling. For example, the following sets the nitrogen hyperfine coupling to  $A_{xx} = A_{yy} = 20$  MHz and  $A_{zz} = 100$  MHz. It defines the x-direction of the tensor as the vector between the two carbons bonded to the nitrogen: since the nitrogen in TEMPO in bonded to only two carbon the vector is uniquely defined. The y-direction is defined as between the particle, the nitrogen in this case, and the oxygen that is bonded to the particle.

```
#[spin_properties(tempo_n, 14N)]
hyperfine_coupling = [20,20,100]*1e6; // Hz.
hyperfine_x = diff(bonded(tempo_c),bonded(tempo_c));
hyperfine_y = diff(particle,bonded(tempo_o));;
```

The arguments diff() can take are tabulated below. The table assumes that the user defined  $\#[group(my\_group)]$ .

| argument                | description   |
|-------------------------|---|
| bonded(my_group)        | particles that are are in my_group and bonded to reference particle                   |
| group(my_group)         | particles that are are in my_group  |
| particle                | the reference particle itself   |
| same_molecule(my_group) | particles that are are in my_group and in the same molecule as the reference particle |

# 0.4 Shell Scripting

The previous interface is useful for an isolated simulation, but to run multiple simulations that are nearly the same except for a few parameters being investigated, running CluE Oxide as a scrip is helpful. To this end there are several methods available. The first is via a shell script: CluE Oxide allows additional option to be appended to a base options file with the "-O" flag. Here is an example Bash script to run several different input structures with the same setting.

```
#!/usr/bin/bash
clue="path/to/clue"

options="path/to/options.txt"
```

```
7 files=(
8    structure_1.pdb
9    structure_2.pdb
10    structure_n.pdb
11 )
12 for file in "${files[@]}"; do
13    $clue $options -0 "\"input_structure_file = $file;\""
14 done
```

Note that CluE Oxide interprets this the same as just appending the options to the end of the options file, the command line options must be be compatible with the options file. This means that the option should not already be set in the options file, and that the options file leaves off in the correct mode. For example if "#[config]" options are to be set, the options file should not end in another mode like "#[group(label=my\_label)]".

# 0.5 pyCluE

Additionally, CluE Oxide has a Python interface.

# 0.5.1 Installing

Within the CluE Oxide source directory, navigate to the pyclue directory.

```
1 cd pyclue
```

The Python interface uses maturin[7] to compile.

```
cd pyclue
python3 -m venv <path/to/virtual/environment>
source <path/to/virtual/environment/bin/activate>
pip install maturin
maturin build
```

One potential issue is that when maturin tries to compile CluE Oxide, it can fail to see the operating system unique flags, and will try to use them all. To account for this, open CluE Oxide's Cargo.toml file and comment out everything under the unneeded operating system section. For example, to compile on Linux add comments as shown below.

```
[target.'cfg(unix)'.dependencies]
ndarray-linalg = { version = "0.15", features = ["openblas-static"] }

#[target.'cfg(windows)'.dependencies.ndarray-linalg]
#version = '0.15.0'
#features = ['intel-mkl']
```

Once built, navigate to target/wheels, source the desired Python environment, and use pip to install the wheel.

```
cd target/wheels
source <path/to/installation/environment/bin/activate>
pip install <pyclue.whl>
```

## 0.5.2 Running

Running CluE Oxide via Python is done through the CluEOptions class.

| parameter                 | values                | description                           |
|---------------------------|-----------------------|---------------------------------------|
| config                    | dictionary of strings | general config input                  |
| group_list                | [Group]               | group input                           |
| structure_properties_list | [StructureProperties] | non-isotope specific properties input |
| spin_properties_list      | [SpinProperties]      | isotope specific properties input     |

The *Group* class signature is shown below.

```
1 class clue_oxide.Group(label, criteria = {})
```

The *label* parameter should be a string and serves the same purpose as the label in  $\#[group(my\_label)]$  discussed earlier. The *criteria* parameter is a dictionary of strings. Similarly StructureProperties in analogous to a  $\#[structure\_properties(my\_label)]$  section.

```
1 class clue_oxide.StructureProperties(label, properties = {})
```

And SpinProperties in analogous to a #[spin\_properties(my\_label, my\_isotope)] section.

```
1 class clue_oxide.SpinProperties(label, isotope, properties = {})
```

Note that the additional string, *isotope*, is required.

Below in an example where an MD frame of TEMPO solvated in water is run with different tunnel splittings. Note that the pyCluE specific syntax is similar is similar to the standard CluE Oxide input files.

```
import clue_oxide as clue
  import matplotlib
   from matplotlib import pyplot as plt
4
  import numpy as np
5
6
  def main():
7
     # Initialize options.
8
9
     options = clue.CluEOptions(
10
         config = {
           "input_structure_file": "\"MD_300K_25ns-30ns_center_0001.pdb\"",
11
12
           "radius": "25", # angstroms
13
           "detected_spin_position": "centroid_over_serials([28,29])",
           "detected_spin_g_matrix": "[2.0097, 2.0064, 2.0025]",
14
           "detected_spin_g_y": "diff(group(tempo_n) , group(tempo_o) )",
15
16
           "detected_spin_g_x": "diff(group(tempo_c1) , group(tempo_c19) )",
           "number_timepoints": "[101]",
17
           "time_increments": "[5e-8]", # s
18
           "cluster_method": "cce",
19
20
           "max_cluster_size": "3",
           "magnetic_field": "1.2", # T
21
22
           "pulse_sequence": "hahn",
           "neighbor_cutoff_3_spin_hahn_mod_depth": "3.23e-5",
23
           "neighbor_cutoff_3_spin_hahn_taylor_4": "1e17", # (rad/s)^4
24
25
         },
         group_list = [
26
           clue.Group("tempo_h", criteria = {
27
             "residues in": "[TEM]",
28
             "elements in": "[H]",
29
30
31
           clue.Group("tempo_n", criteria = {
             "residues in": "[TEM]",
32
             "elements in": "[N]",
33
34
           }),
35
           clue.Group("tempo_o", criteria = {
             "residues in": "[TEM]",
36
37
             "elements in": "[0]",
38
           F).
39
           clue.Group("tempo_c1", criteria = {
40
             "residues in": "[TEM]",
             "elements in": "[C]",
41
42
             "serials in": "[1]"
```

```
43
            }),
44
            clue.Group("tempo_c19", criteria = {
              "residues in": "[TEM]",
45
              "elements in": "[C]",
 46
              "serials in": "[19]"
47
48
            }),
          ],
49
          spin_properties_list = [
50
            clue.SpinProperties("tempo_n", "14N", properties = {
51
52
              "hyperfine_coupling": "[20,20,100]*1e6", # Hz
 53
              "hyperfine_x": "diff(bonded(tempo_c1),bonded(tempo_c19))",
              "hyperfine_y": "diff(particle,bonded(tempo_o))",
54
              "electric_quadrupole_coupling": "[-0.28, -1.47, 1.75]*1e6", # Hz
55
              "electric_quadrupole_x": "diff(bonded(tempo_c1),bonded(tempo_c19))",
 56
              "electric_quadrupole_y": "diff(particle,bonded(tempo_o))",
57
58
            }),
          1)
59
60
61
62
      # Initialize figure.
     plt.rc('font', size=12)
plt.rc('axes', labelsize=12)
63
64
65
      fig, (ax) = plt.subplots(1,1,figsize=(8, 6) );
66
67
      cmap = matplotlib.colormaps['viridis']
68
69
      # Set desired methyl tunnel splittings.
70
      tunnel_splittings = np.array([0,20,40,80,100])*1e3;
71
72
      # Loop over tunnel splittings.
73
      for nut in tunnel_splittings:
74
75
        # Set tunnel splitting.
        options.set_spin_properties("tempo_h", "1H",
76
 77
            key = "tunnel_splitting", value = f"{nut}")
78
79
        # Print CluE input file.
80
        options.print()
81
82
        # Run CluE.
        time , signal = options.run()
83
84
 85
        # Plot output.
86
        ax.plot(np.array(time)*1e6,np.real(np.array(signal)),
87
          color = cmap(nut/max(tunnel_splittings)), linewidth=1);
88
89
90
      # Polish the plot.
91
     norm = matplotlib.colors.Normalize(vmin=0, vmax=max(tunnel_splittings)*1e-3)
92
      sm = plt.cm.ScalarMappable(cmap=cmap, norm=norm)
93
94
      sm.set_array([])
95
      plt.colorbar(sm, label=r"$\nu_\mathrm{t}$ (kHz)", orientation="vertical")
96
      ax.set_xlabel(r"$2\tau$ (s)");
97
      ax.set_ylabel("normalized echo amplitude");
98
99
      ax.axis([0,10,0,1.0])
100
101
      # Save figure.
      plt.savefig('CluE-sims.png')
102
103
104 if __name__ == "__main__":
105
     main();
```

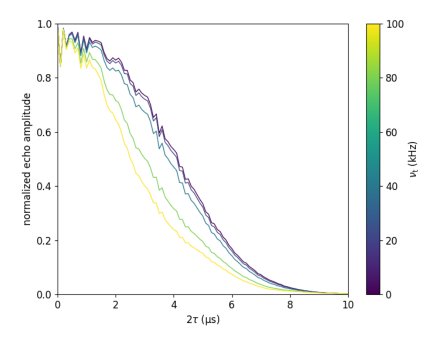


Figure 2: These Hahn echo simulation show the output of the previous example. The color-axis indicates the tunnel splitting.

Here the input is modified before sending it to CluE Oxide, so settings can be overwritten in the script. Figure 2 shows the output. Note that to save on computation time, these simulation are only at the 3-CCE level, and only use one frame at a single orientation.

# 0.6 Rust

Finally, since CluE Oxide is written in Rust, it is fairly straight forward to integrate CluE Oxide into a Rust program. Here is a script to run CluE Oxide over different proton concentrations. The output is shown in figure 3.

```
use clue_oxide as clue;
   use clue_oxide::{
3
     physical_constants::Isotope,
4
     \verb|config::particle_config::\{IsotopeAbundance,IsotopeDistribution\}|
5
6
   fn main() {
9
        // Initialize options.
10
        let config = match clue::config::Config::from("
            input_structure_file = \"TEMPO_wat_gly_70A.pdb\";
11
12
            radius = 25; // angstroms.
13
            detected_spin_position = centroid_over_serials([28,29]);
14
15
            detected_spin_g_matrix = [2.0097, 2.0064, 2.0025];
            detected_spin_g_y = diff(group(tempo_n) , group(tempo_o) );
detected_spin_g_x = diff(group(tempo_c1) , group(tempo_c19) );
16
17
            number_timepoints = [101,140];
18
19
20
            time_increments = [50,500]*1e-9;
21
            cluster_method = cce;
22
            max_cluster_size = 2;
```

```
23
            magnetic_field = 1.2; // T.
24
            pulse_sequence = hahn;
25
26
           number_system_instances = 10;
           orientation_grid = random(1);
27
28
29
           neighbor_cutoff_distance = 4; // angstroms.
30
31
         #[group(tempo_n)]
32
           residues in [TEM];
33
            elements in [N];
34
35
         #[group(tempo_o)]
36
           residues in [TEM];
37
           elements in [0];
38
39
         #[group(tempo_c1)]
40
           serials in [1];
41
42
         #[group(tempo_c19)]
43
            serials in [19];
44
45
         #[spin_properties(tempo_n, 14N)]
           hyperfine_coupling = [20,20,100]*1e6;
46
           hyperfine_x = diff(bonded(tempo_c1),bonded(tempo_c19));
hyperfine_y = diff(particle,bonded(tempo_o));
47
48
49
50
            electric_quadrupole_coupling = [-0.28, -1.47, 1.75]*1e6;
51
           electric_quadrupole_x = diff(bonded(tempo_c1),bonded(tempo_c19));
            electric_quadrupole_y = diff(particle,bonded(tempo_o));
52
53
54
         #[group(solvent_h)]
55
           residues not in [TEM];
           elements in [H];
56
57
         #[structure_properties(solvent_h)]
58
           isotope_abundances = {1H: 1, 2H: 0};
59
60
61
       "){
62
         Ok(cfg) => cfg,
         Err(err) => {
63
           eprintln!("CluE Error: {}.",err);
64
65
66
         },
67
       };
68
69
70
       // Find where the solvent properties are stored.
       let mut solvent_h_idx: Option <usize> = None;
71
72
       for (idx, p_cfg) in config.particles.iter().enumerate(){
73
74
         if p_cfg.label == "solvent_h"{
           solvent_h_idx = Some(idx);
75
76
            break:
         }
77
       }
78
79
80
       let solvent_h_idx = solvent_h_idx.expect("Could not find solvent_h_idx.");
81
82
83
       // Initialize desired mole fractions.
84
       let proton_fractions = vec![1.0, 0.75, 0.5, 0.25, 0.0];
85
86
       // Loop over mole fractions.
87
       for &frac in proton_fractions.iter(){
88
89
         let mut frac_config = config.clone();
90
```

```
91
          frac_config.save_name = Some(format!("CluE-proton_frac_{}",frac));
92
93
          let properties = frac_config.particles[solvent_h_idx]
94
            .properties.as_mut().expect("No properties for solvent_h");
95
96
          // Set isotope mole fractions.
97
          let isotope_abundances = vec![
98
            IsotopeAbundance{isotope: Isotope::Hydrogen1, abundance: frac},
99
            IsotopeAbundance{isotope: Isotope::Hydrogen2, abundance: 1.0 - frac},
100
101
          properties.isotopic_distribution = IsotopeDistribution{
102
103
              isotope_abundances,
104
              void_probability: None
105
          };
106
          // Run CluE.
107
108
          match clue::run(frac_config){
109
            Ok(_) => (),
110
            Err(err) => eprintln!("CluE Error: {}.",err),
111
112
113 }
```

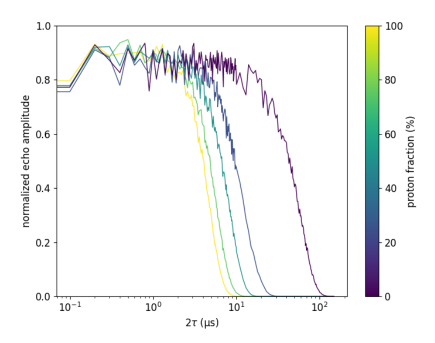


Figure 3: These Hahn echo simulation show the output of the previous example. The color-axis indicates the proton fraction. Note that the  $2\tau$ -axis is logarithmic.

The primary way to use CluE Oxide with Rust is to create an instance of Config and feed it to the run function. The easiest way to initialize an instance of Config is to use Config::from(input: &str); this uses the same syntax as the CluE Oxide input files. Note that because the each new  $frac\_config$  is modified after it has been constructed, the safety check to ensure values are not set multiple times is passed and properties can be modified directly. Every field of Config has a data type that is either an Option < T > or a Vec::< T >, for generic type T; values of None or an empty vector are considered unset. Which entries are required depends on the details of the simulation. CluE Oxide will return an error explaining what is missing if it needs an option that is unset. Several fields have default values, which can be set with the set defaults

method. Note that set\_defaults will not overwrite user set values. This method is called automatically when using the command line or python interface. The *Config* struct is defined as shown below.

```
#[derive(Debug, Clone, Default)]
  pub struct Config{
    pub clash_distance: Option<f64>,
    pub clash_distance_pbc: Option<f64>,
    pub cluster_batch_size: Option<usize>,
    pub cluster_method: Option<ClusterMethod>,
    pub density_matrix: Option<DensityMatrixMethod>,
     pub detected_spin_g_matrix: Option<TensorSpecifier>,
     pub detected_spin_identity: Option<Isotope>,
     pub detected_spin_multiplicity: Option<usize>,
10
     pub detected_spin_position: Option<DetectedSpinCoordinates>,
12
     pub detected_spin_transition: Option<[usize;2]>,
13
     pub extracell_particles: Vec::<ParticleConfig>,
14
     pub input_structure_file: Option<String>,
     pub load_geometry: Option<LoadGeometry>,
15
16
     pub magnetic_field: Option<Vector3D>,
     pub max_cluster_size: Option<usize>,
17
18
     pub neighbor_cutoff_coupling: Option<f64>,
19
     pub neighbor_cutoff_delta_hyperfine: Option<f64>,
20
     pub neighbor_cutoff_dipole_perpendicular: Option<f64>,
21
     pub neighbor_cutoff_distance: Option<f64>,
22
     pub neighbor_cutoff_3_spin_hahn_mod_depth: Option<f64>,
23
     pub neighbor_cutoff_3_spin_hahn_taylor_4: Option<f64>,
24
     pub number_system_instances: Option<usize>,
25
    pub number_timepoints: Vec::<usize>,
26
     pub orientation_grid: Option<OrientationAveraging>,
     pub particles: Vec::<ParticleConfig>,
27
28
     pub pdb_model_index: Option<usize>;
     pub pulse_sequence: Option<PulseSequence>,
29
     pub remove_partial_methyls: Option<bool>,
30
31
     pub root_dir: Option<String>,
     pub radius: Option<f64>,
32
33
     pub rng_seed: Option<u64>,
34
     pub save_name: Option<String>,
35
     time_axis: Vec::<f64>,
    pub time_increments: Vec::<f64>,
36
37
     pub write_auxiliary_signals: Option<String>,
38
     pub write_bath: Option<String>,
39
     pub write_clusters: Option<String>,
    pub write_info: Option<String>,
    pub write_exchange_groups: Option<String>,
41
42
     pub write_orientation_signals: Option<String>,
43
     pub write_structure_pdb: Option<String>,
    pub write_tensors: Option<String>,
44
45 }
```

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