

### **BOISE STATE UNIVERSITY COLLEGE OF ENGINEERING**

### ABSTRACT

Coarse-graining---simplifying models of molecules by representing a collection of atoms with a simulation element like a sphere or ellipsoid---can significantly increase the timescales accessible to simulations without loss of structural accuracy. Spherical simulation elements are inaccurate representations of flat molecular structures, though, which are better represented with anisotropic shapes like ellipsoids. In this work we debug and extend open-source software (GRiTS) for calculating the shapes and orientations of an ellipsoid representing a collection of atoms. These functionalities are useful for both validating the correctness of coarse-grained models and for training advanced anisotropic potentials that can be used in accelerated molecular simulations.

### OBJECTIVE

The objective of this project is to create an anisotropic bead option to the existing open-source software (GRiTS) in order to visualize orientational phenomena.





# Coarse-graining of Anisotropic Molecules for Energy Materials Simulations

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### COARSE-GRAIN PARAMETERS (CONT.)

- In order to the calculate the orientation, a series of steps must first be completed: 1) A plane must be created using the dimensions of the monomer. In order to do this a second vector which is unique and non-parallel to the direction vector is used, called
- A CoM. This will be referred to as the reference plane. 2) A second monomer is created, which is meant to represent any other orientation than the reference plane.
- 3) The cross product of the normal vector of the reference plane and the normal vector of the particle's reference plane is calculated to produce an axis of rotation.
- 4) The reference plane is rotated by some angle theta along this axis in order to transform the reference plane into the other plane.
- 5) The axis angle then can be transformed into a quaternion using a python package called rowan, so that they are compatible with our simulation engine, HOOMD-blue.





Figure 3. The formula on the left shows the math required in order to perform the rotation between the reference plane  $(n_0)$  and the new plane  $(n_1)$ .  $\overrightarrow{v_{ax}}$  represents the axis about which the reference plane is being rotated in order to create the new plane, and  $\theta_{rot}$ represents the amount that the reference plane needs to be rotated in order to create the new plane

### def get\_quaternion(n1, n0=np.array([0, 0, 1])): """Calculates axis and angle of rotation given two planes normal vectors""" V\_axis = np.cross(n0, n1) theta\_numerator = np.dot(n0, n1) theta\_denominator = np.linalg.norm(n0) \* np.linalg.norm(n1) theta\_rotation = np.arccos(theta\_numerator / theta\_denominator) quaternion = rowan.from\_axis\_angle(V\_axis, theta\_rotation) return quaternion

Figure 4. The function above converts the two planes into a rotational quaternion.

Why use quaternions over Euler angles? Quaternions are more compact, efficient, and stable than Euler angles, but are harder to understand. They extend to imaginary numbers, and rather than returning a typical (X, Y, Z) they return (C, XS, YS, ZS) where C represents  $\cos(\theta/2)$ , and S represents  $\sin(\theta/2)$ . Because of this added fourth element, they are resistant to gimbal lock. Gimbal lock is the alignment of rotational planes, and once two separate rotational planes align, they can no longer be represented uniquely using normal vectors.

Carbon

= Sulfur

Center of Mass



Figure 5. The image above is a rendering of four PPS monomers with different orientations using a visualization software called OVITO. The transparent ellipsoids enveloping each of the four PPS monomers represent the anisotropic coarse-grained beads for each respective monomer, reflective of their orientation.

### Conclusions

- parameters at the time.

Future work

- $\succ$  Write tests to ensure that code will apply to any input.
- coarse-grain systems.

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### APPLICATION VISUALIZED

### CONCLUSIONS AND FUTURE WORK

> Anisotropic coarse-grained particles can be mapped from arbitrary sets of atoms. > The mapping can be done while remaining agnostic of the bead dimensions/forcefield

> This mapping can be performed on all atom trajectory files in one workflow step.

> These coarse-grain mappings can be used to train those forcefield parameters with various techniques (e.g., machine learning, iterative Boltzmann inversion, force-matching, etc.). ➤ Revise and integrate code with existing workflows (GRiTS).

> Compare performance and accuracy of anisotropic-mapped and traditional sphere-mapped

➤ Implement additional mapping schemes (e.g., 3-axis anisotropic beads).  $\blacktriangleright$  Create solutions for edge cases (e.g., linear monomers, n<3 particle systems, etc.).

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