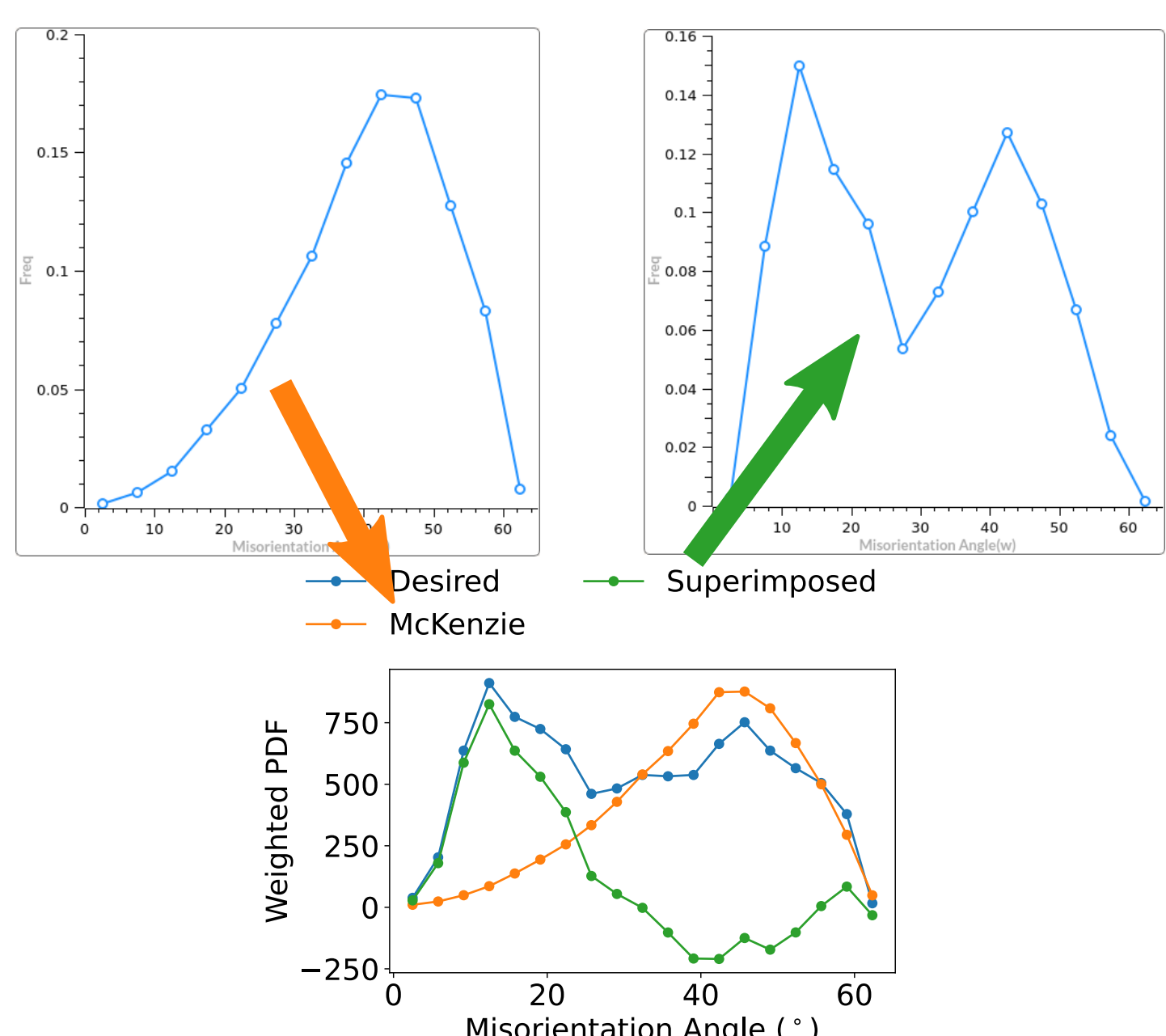


Statistics

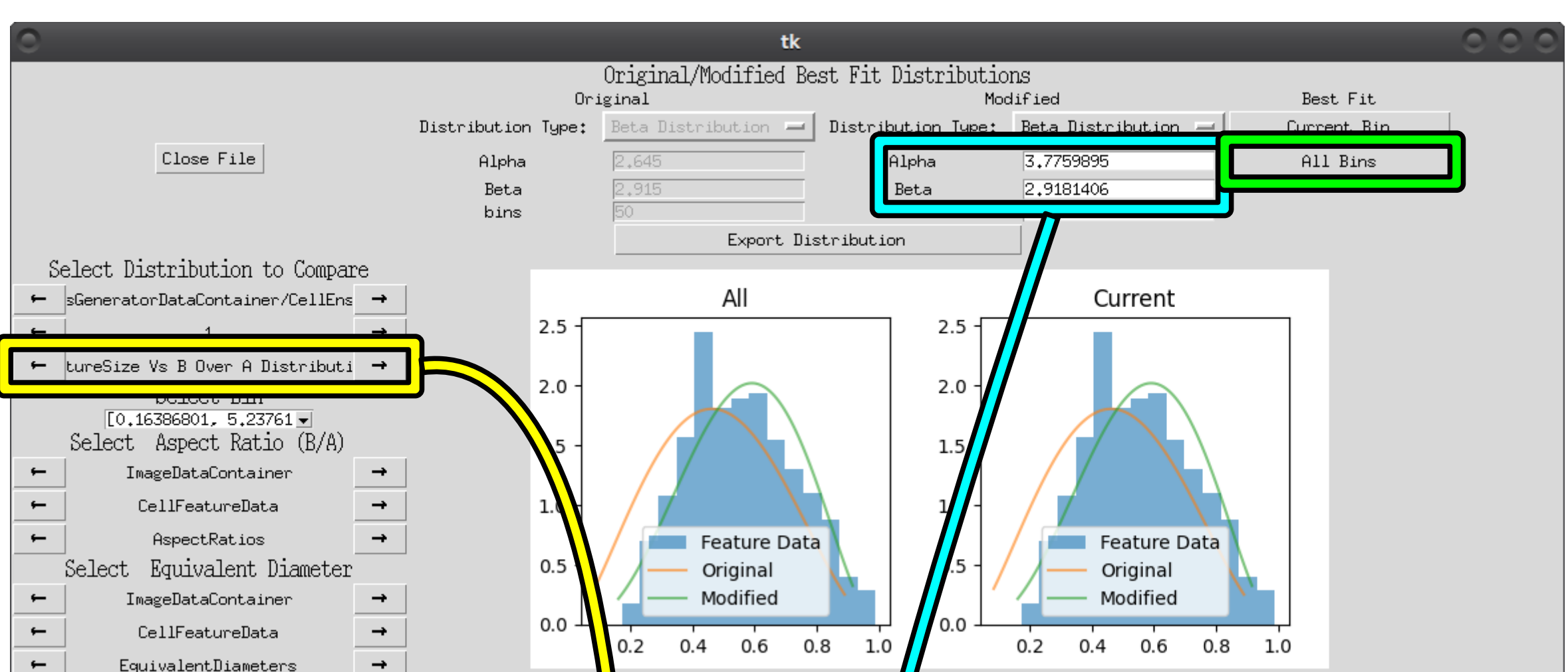
- | | |
|-------------|------------|
| BC | MAD |
| BS | Mask_Clear |
| Bands | Phases |
| Error | X |
| EulerAngles | Y |

3. Extract Feature Attributes - Segment by misorientation to re-obtain grains and recalculate morphological and crystallographic characteristics.
4. Create MDF - Create a superposition of the desired MDF and the negative of Dream3D's default McKenzie MDF.

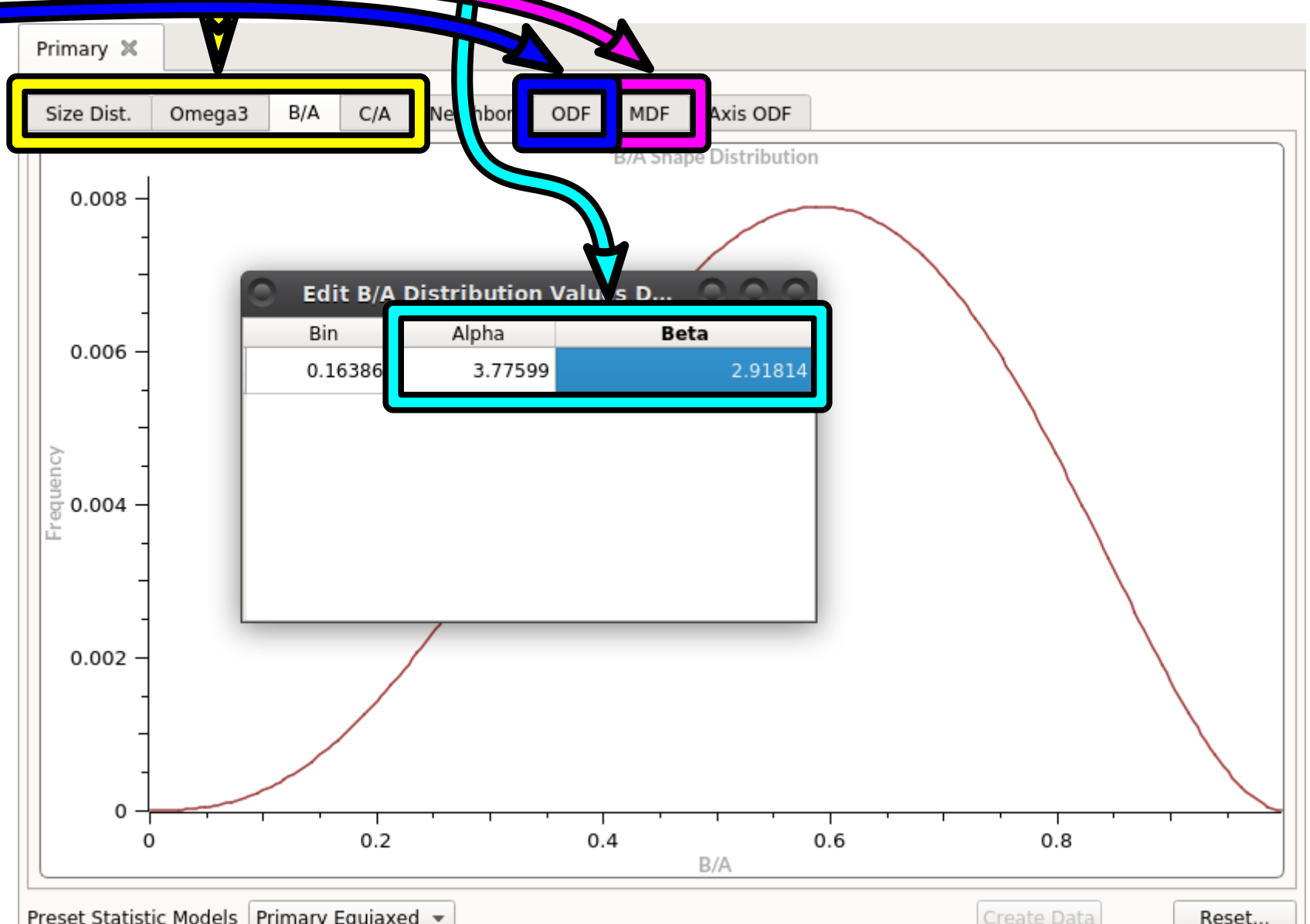







5. **Generate Initial Statistics** - Create a **CellEnsembleData/Statistics** directory in **Dream3D** file for **utils** packing to access.

6. **Grain Packing Utils** - Generate the statistical distribution parameters of the morphological characteristics of the CGs.



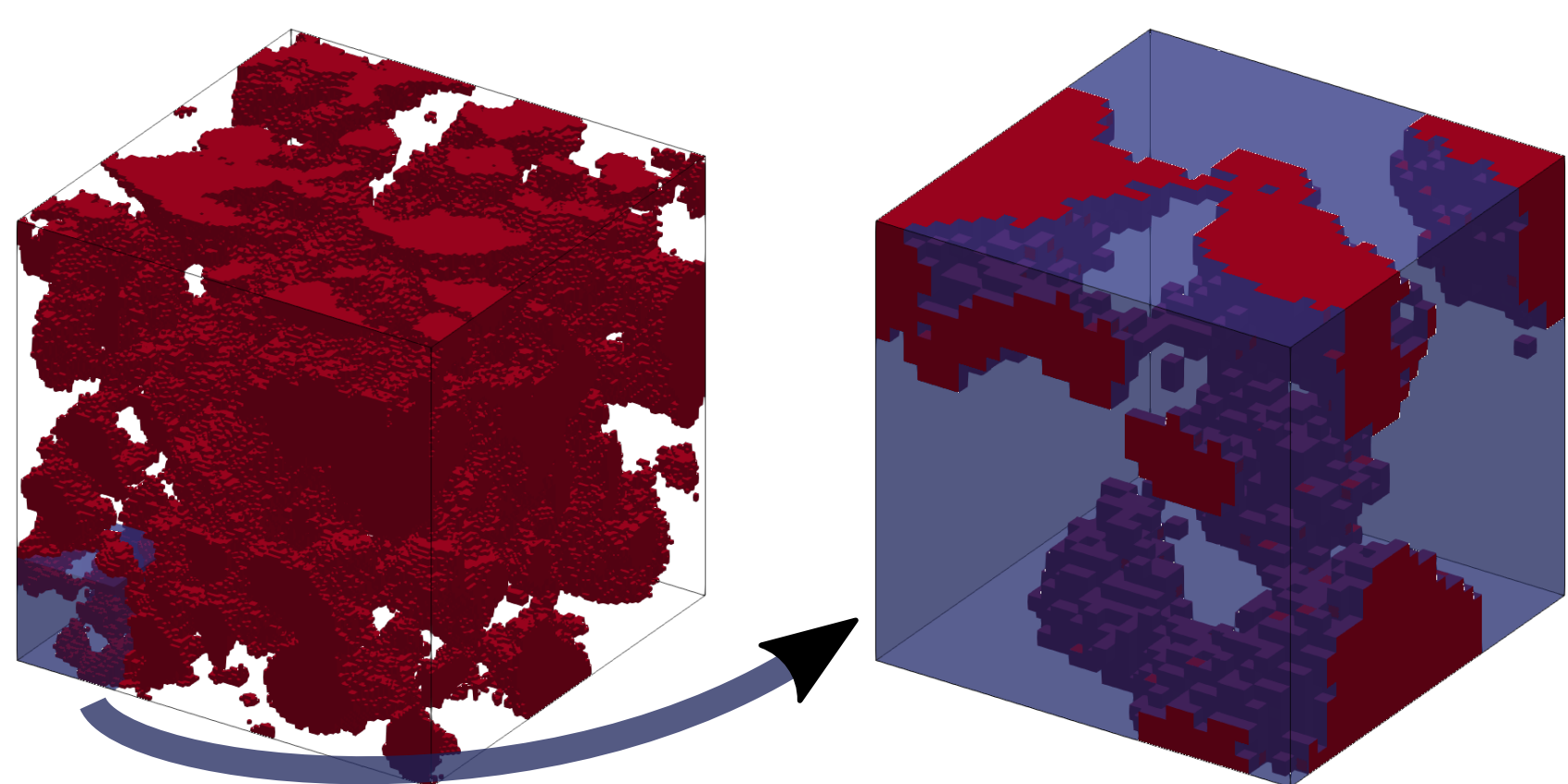
7. **Generate Final Statistics** - Create and populate the CellEnsembleData/Statistics directory that will be used to pack the SEVM with a statistically equivalent coarse grain structure.



-  Auto-Fit via SciPy
-  Fitting Parameters
-  Morphological
(Copy From 6)
-  Crystallographic
(Load from 3.4)
- 

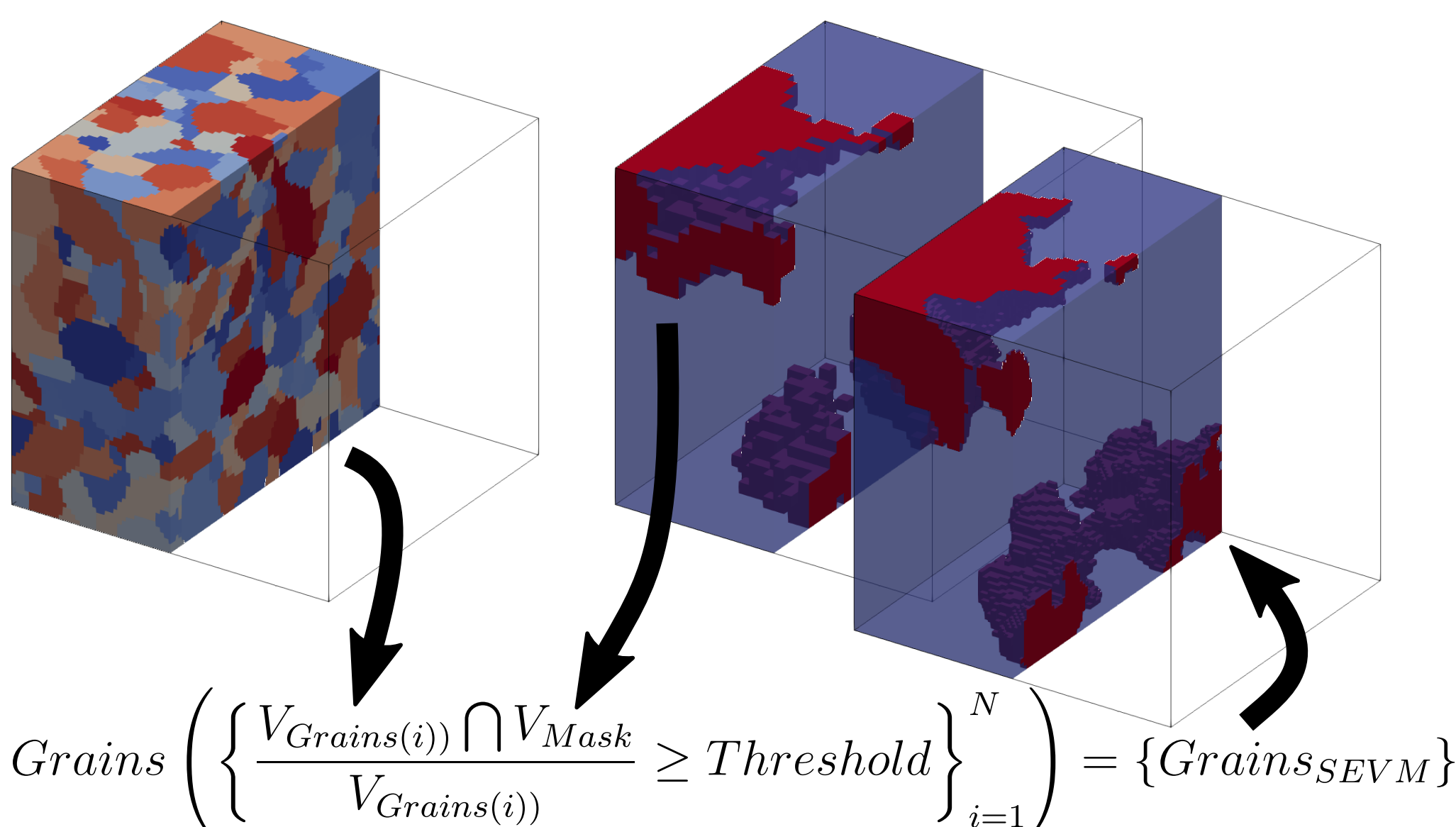
Morphology

8. Choose Subdomain - Sample a rectangular subdomain of the SliceGAN generated 3D bimodal mask.



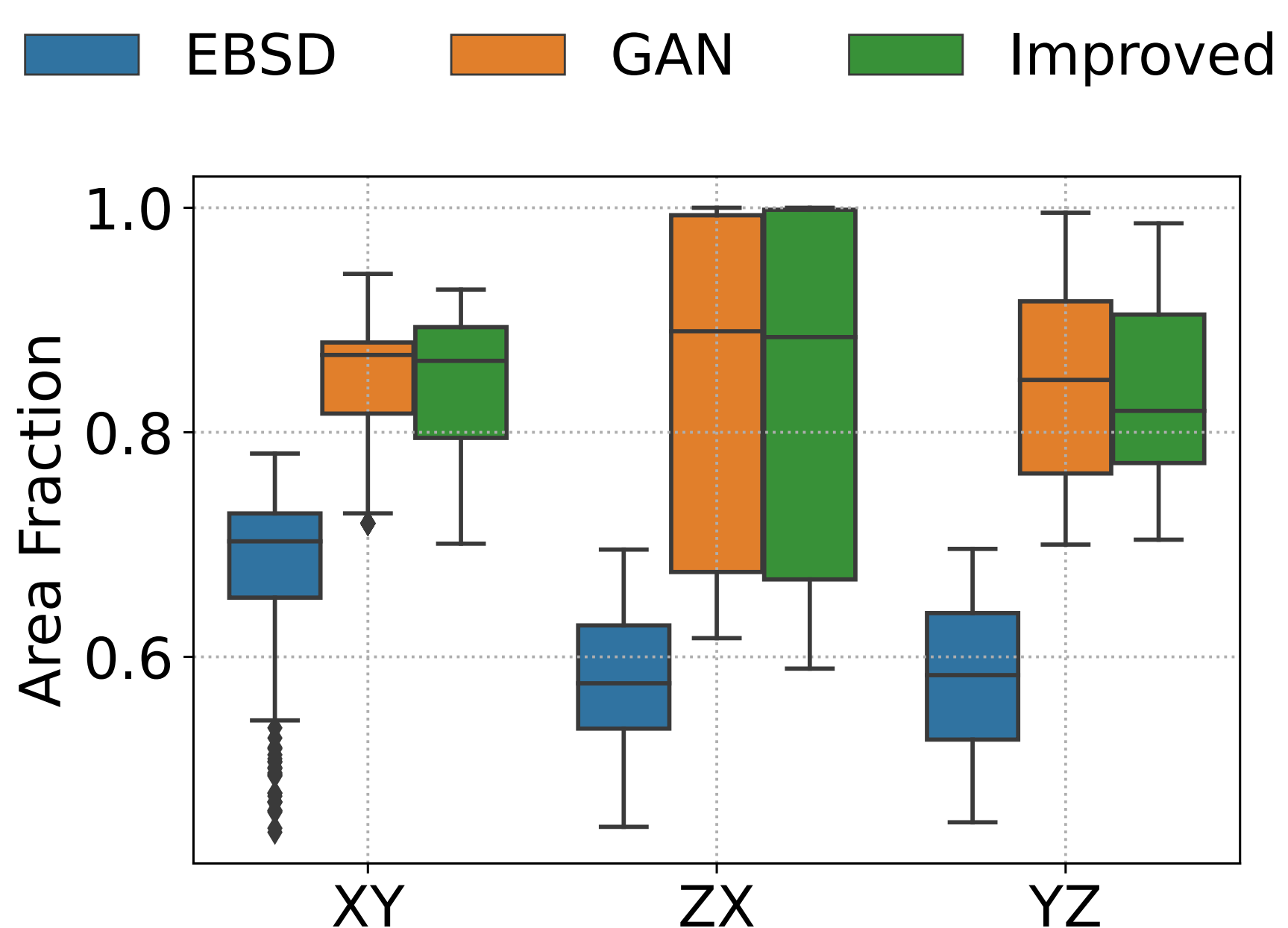
9. **Format Boundaries** - Remove unnecessary data from bimodal mask, and copy over the CellEnsembleData folder from 7.
10. **Generate Grains** - Generate a full cube of statistically equivalent CGs.
11. **Create Improved Mask** - Generate a new mask from the intersection of the CGs

11. **Create Improved Mask -**
Generate a new mask from the intersection of the CGs and the SlicaGAN mask.



12. **Apply Improved Mask** - Delete all grains outside of the improved mask.

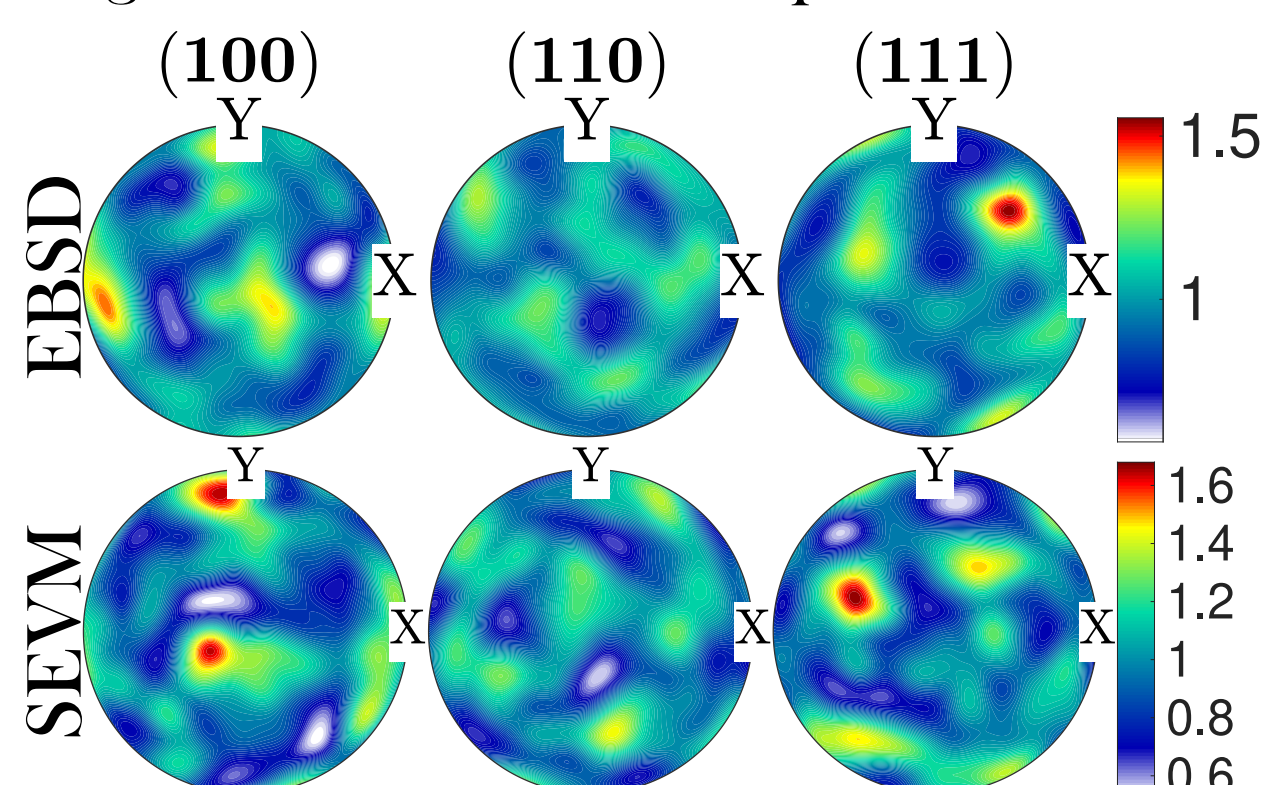
13. **Compare Area Fractions** - Compares the area fractions of orthogonal slices of the improved mask to investigate the anisotropic density.



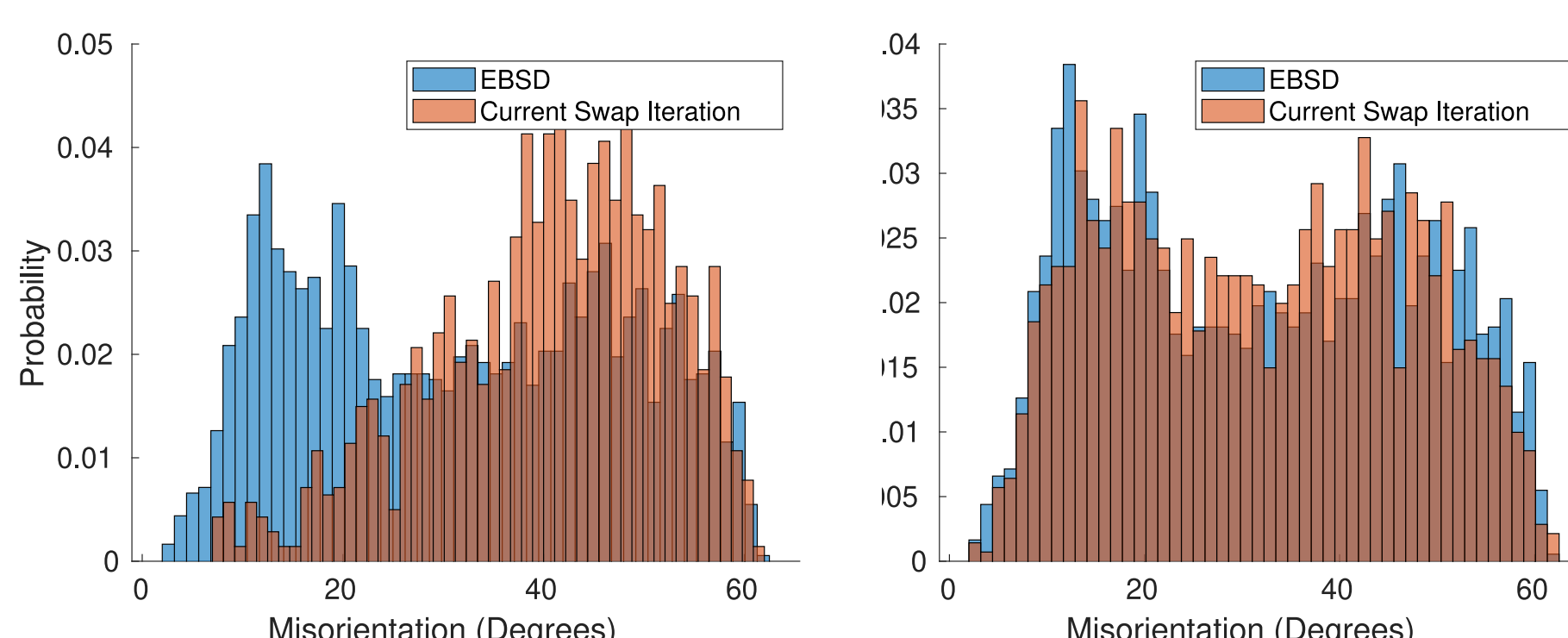
Crystallography

14. **Generate Crystallography** - Assigns crystallographic orientations from the ODF and MDF created by 3, 4. Created crystallography found to be unreliable. Used here only to initialize crystallography variables.

15. Replace Crystallography - Use MTEX to build an ODF from the EBSD file, then replace the existing Euler angles with random samples from the ODF.



16. Grain Swap - Swap the crystallographic orientations of two random grains until the Kolmogorov Smirnov (**KS**) statistic of the **MDF** is less than some given threshold.



17. Grain Swap Import - Import the results of 16 to the *.dream3d file.

18. Lattice Constants Import - Import the crystallographic lattice constants from 3 that match the remaining materials.

Compare

19. **Extract Feature Attributes Synthetic** - Calculate all the dependent grain characteristic attributes.
20. **Compare Stats** - Compare the grain characteristic attributes between the EBSD file and the SEVM

