PRISMS-Fatigue: Python script installation and execution guide

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Introduction

This document describes the five Python scripts associated with the PRISMS-Fatigue toolkit. The user is required to download Python on their local computer but the scripts may also be executed on a supercomputing cluster. Several common libraries are required which are listed and imported at the top of each script (e.g., NumPy, os, pickle, pandas, etc.). The majority of these should come preinstalled with a standard Python installation package such as Anaconda or equivalent. A recent version of the open-source microstructure instantiation software DREAM.3D is also required [1]. ParaView is used for visualization.

There are several publications that should be read thoroughly by aspiring Prisms-Fatigue users [2-61. Users should watch the PRISMS-Plasticity tutorial videos located https://www.youtube.com/playlist?list=PL4yBCojM4Swqy4FRteqxHWSiM1uiOOesj. There are also a set videos tutorial associated with these scripts https://www.youtube.com/playlist?list=PL4yBCojM4Swo3CvIA57syFrzk3p1mugP5 to supplement this guide. The five Python scripts are listed below and described subsequently in this document. Users are strongly encouraged to examine each script in detailed to gain a fundamental understanding of the workflows! Each script considers the calculation of Fatigue Indicator Parameters (FIPs) [2].

- 1. generate_microstructures.py
- 2. calculate FIPs.py
- 3. volume_average_FIPs.py
- 4. compule_and_plot_FIPs.py
- 5. gamma_plane.py

The raw data associated with the PRISMS-Fatigue manuscript is available for download from Materials Commons at: https://doi.org/10.13011/m3-rcyy-gx13. The results and figures in the PRISMS-Fatigue manuscript can be entirely replicated by downloading this raw data and executing the last four Python scripts. The video tutorials do just this, so users are encouraged to go through this document as well as the online videos. Finally, the manuscript associated with this framework is available for download at https://doi.org/10.1038/s41524-021-00506-8 [2].

Python installation

Python can be downloaded and installed at https://www.anaconda.com/products/individual for several operating systems as listed below, but the user is free to install Python in whichever way is most convenient. Various Python packages are required to execute the scripts that are straightforward to install. The scripts are backwards compatible with older 2.X versions of Python.

Anaconda Installers							
Windows #	MacOS É	Linux 🛆					
Python 3.8 64-Bit Graphical Installer (466 MB)	Python 3.8 64-Bit Graphical Installer (462 MB)	Python 3.8 64-Bit (x86) Installer (550 MB)					
32-Bit Graphical Installer (397 MB)	64-Bit Command Line Installer (454 MB)	64-Bit (Power8 and Power9) Installer (290 MB)					

Fig. 1. Python installation options at https://www.anaconda.com/products/individual.

ParaView

The open-source software ParaView is used to visualize microstructures and PRISMS-Plasticity simulation results. It is available at https://www.paraview.org/download/ for users to download.

Get the Software

You can either download binaries or source code archives for the latest stable or previous release or access the current development (aka nightly) distribution through Git. Specific license information can be found here. This software may not be exported in violation of any U.S. export laws or regulations. For more information regarding Export Control matters please go to https://kitware.com/export_control/index.html.

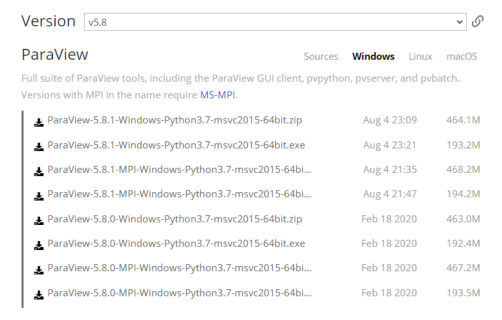


Fig. 2. ParaView installation options at https://www.paraview.org/download/.

DREAM.3D installation

DREAM.3D is a powerful, open-source software with many capabilities including (but not limited to) generation of synthetic microstructures, microstructure reconstruction using data sets such as electron back scatter diffraction (EBSD) or high energy x-ray diffraction microscopy (HEDM), and analysis of statistical information [1]. It is available for download at http://dream3d.blueguartz.net/. Prospective PRISMS-Fatigue users should read the work by Groeber and Jackson [1] to understand the structure of the program, which is briefly reviewed here. In addition to the PRISMS-Fatigue video tutorials, DREAM.3D specific tutorials for more advanced users are available https://www.youtube.com/channel/UCjeF8pFMzET5ZN3vsBHATpg. A screenshot of the program files and folders is shown in Fig. 3. The executable DREAM3D.exe opens a blank Graphical User Interface (GUI) as shown in Fig. 4.

Name	Date modified	Type	Size	^
Data	11/2/2020 8:36 AM	File folder		
Help	11/2/2020 8:36 AM	File folder		
lib	11/2/2020 8:36 AM	File folder		
Plugins	11/2/2020 8:36 AM	File folder		
PrebuiltPipelines	11/2/2020 8:36 AM	File folder		
Ø DREAM3D.exe	11/2/2020 8:36 AM	Application	1,724 KB	
PipelineRunner.exe	11/2/2020 8:36 AM	Application	91 KB	
RESTServer.exe	11/2/2020 8:36 AM	Application	86 KB	
oncrt140.dll	11/2/2020 8:36 AM	Application exten	325 KB	
EbsdLib.dll	11/2/2020 8:36 AM	Application exten	1,698 KB	
EMMPMLib.dll	11/2/2020 8:36 AM	Application exten	90 KB	
H5Support.dll	11/2/2020 8:36 AM	Application exten	114 KB	
hdf5.dll	11/2/2020 8:36 AM	Application exten	2,914 KB	
hdf5_cpp.dll	11/2/2020 8:36 AM	Application exten	332 KB	
ITKCommon-4.13.dll	11/2/2020 8:36 AM	Application exten	820 KB	
MIN 11 VI	11/2/2020 8:36 AM	Annlication exten	76 KR	~

Fig. 3. DREAM.3D program folder (version 6.5.141 downloaded November 2020). The path of the *PipelineRunner.exe* executable is an input in the first Python script.

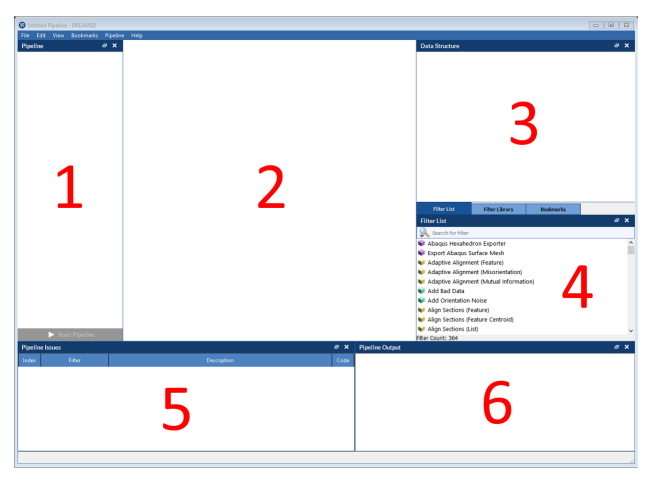


Fig. 4. Empty DREAM.3D GUI. Filters from region 4 are "dragged and dropped" into region 1 for sequential execution. Region 2 contains unique settings for each filter. The structure of the data is shown in region 3. Regions 5 and 6 display pipeline issues/warnings/errors and pipeline output, respectively.

Fig. 5 shows a screenshot of what will be referred to as the microstructure instantiation pipeline "input file." This file consists of only two filters. The first specifies the desired microstructure statistics that include crystal symmetry, grain size distribution, crystallographic texture, grain morphology, phase volume fraction, etc. Once these options are set, the user must click on "Create Data"! The second filter simply writes this information to a ".dream3d" type file. At this point, only the desired *statistics* are stored in this file. As part of the initial PRISMS-Fatigue release, six input files are available for face centered cubic microstructures. These include equiaxed and elongated/rolled grain morphologies, and cubic, random, and rolled target crystallographic textures. Sample microstructure instantiations are depicted in Fig. 6. There are several other DREAM.3D files available as part of PRISMS-Plasticity.

An important note is that any changes made to the DREAM.3D "input" file shown in Fig. 5 cannot be simply saved by clicking "File \rightarrow save"! The user must instead click on "Create Data" in the middle of the screen on the "StatsGenerator" options and then click "Start Pipeline" to execute these two filters in sequence. The name of the resultant .dream3d file in the second filter should also be updated.

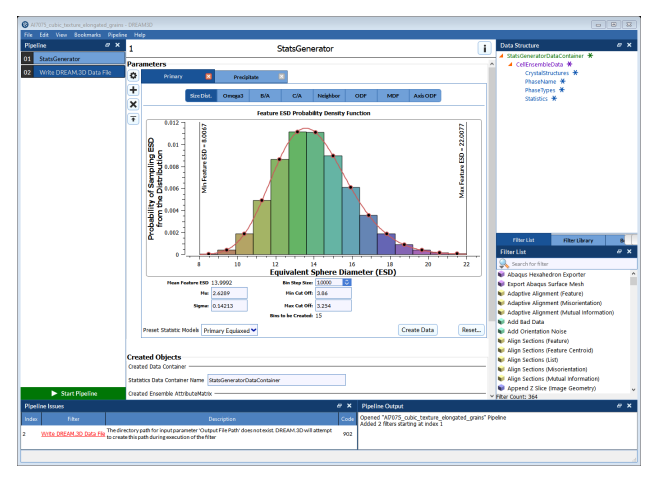


Fig. 5. DREAM.3D "input file" consisting of two filters.

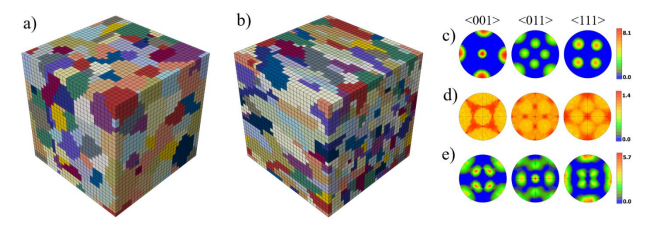


Fig. 6. Sample microstructure instantiations with (a) equiaxed and (b) elongated/rolled grain (grain elongation ratio of 5:1:1) morphology. The different target crystallographic textures include (c) cubic, (d) random, and (e) rolled.

The second DREAM.3D file is shown in Fig. 7 and is referred to as the DREAM.3D "pipeline." Note that the first filter in this pipeline is "Read DREAM.3D Data File" which reads in a ".dream3D" file, for instance the one created by the "input file" shown in Fig. 5. Although a single .json "pipeline" can be used to instantiate microstructures, the scripted workflows presented here split these into an "input" file and a "pipeline." The latter remains unchanged and contains the necessary filters to instantiate microstructures and create other necessary files whereas the former is a simple file that users can edit to change target microstructure statistics. The pipeline file has the .json extension and can be edited as a text file. In fact, the *generate_microstructures.py* script edits the .json pipeline file before execution with user inputs in the main() function.

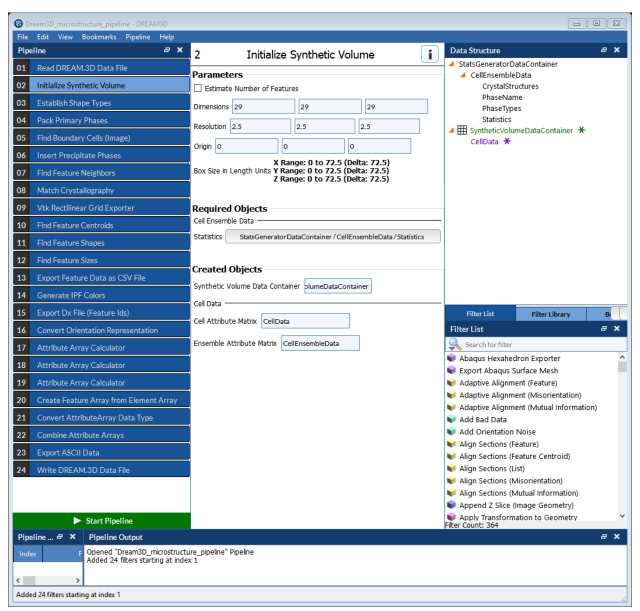


Fig. 7. DREAM.3D "pipeline file" with the filters necessary to instantiate microstructures for CPFE simulation.

generate microstructure.py

The first script generates microstructures for CPFE simulation in PRISMS-Plasticity. A screenshot of the "main()" input function to be edited by the user is shown below. This and the other four Python scripts can be called either from a command prompt window as "python generate_microstructure.py" or using an interactive version of Python (i.e., executing the "ipython" command in command prompt and importing the script). The latter method is particularly useful because of its debugging capabilities. Fig. 9 and Fig. 10 depict both of these methods. The terms voxel and element are used interchangeably in this guide since the microstructure voxels directly represent the elements for CPFE simulations.

```
def main():
     # Run from command prompt
     ''' Specify directories and paths '''
     # Directory where microstructure data should be generated and pre-processed
     # This command creates a directory in the same location as the "PRISMS-Fatigue" directory with python scripts and DREAM.3D files
     directory = os.path.dirname(DIR LOC) + '\\tutorial\\test run 1
     \mbox{\sharp} Alternatively, the directory can be expressed as an absolute path as:
     # directory = r'C:\Users\stopk\Documents\GitHub\PRISMS-Fatigue\tutorial\test run 1
     # Location of DREAM.3D input file; should consist of only the "StatsGenerator" and "Write DREAM.3D Data File"
     # "StatsGenerator" inputs include grain size distribution, crystallographic texture, grain morphology, etc.
     # Six ".dream3d" files are included in PRISMS-Fatigue
     d3d_input_file = os.path.abspath(DIR_LOC) + '\\Al7075_random_texture_equiaxed_grains.dream3d'
      # Once again, this may be specified using an absolute path as:
     # d3d_input_file = r'C:\Users\stopk\Documents\GitHub\PRISMS-Fatigue\A17075 cubic texture equiaxed grains.dream3d'
     # Average grain size as determined in the "StatsGenerator" filter of the .dream3d file above
     # Used for automated band and sub-band sizing as shown below but which can be overwritten by the user
     # Therefore, this does NOT change the grain size generated and only affects the way in which microstructures are banded and sub-banded!
     avg_grain_size = 0.014 # millimeters
     # Location of DREAM.3D .json pipeline
     # This can be modified by the user to include additional outputs
     d3d_pipeline_path = os.path.abspath(DIR_LOC) + '\\Dream3D_microstructure_pipeline.json'
     # Once again, this may be specified using an absolute path as:
     # d3d_pipeline_path = r'C:\Users\stopk\Documents\GitHub\PRISMS-Fatigue\Dream3D_microstructure_pipeline.json'
     # Location of DREAM.3D 'PipelineRunner.exe' file; this should be in the DREAM.3D program folder
     d3d_executable_path = r'C:\Users\stopk\Desktop\DREAM3D-6.5.141-Win64\PipelineRunner.exe
     ''' Specify desired microstructure size and shape '''
     # Size of microstructure instantiations in millimeters, in the X, Y, and Z directions, respectively.
     size = np.asarray([.0725,.0725,.0725])
     # Shape of microstructure instantiations (number of voxels/elements), in the X, Y, and Z directions, respectively.
     # IMPORTANT: at this point, only CUBIC voxel functionality supported even with a non-cubic microstructure
     # I.e., size = [.05, .1, .025] and shape = [50, 100, 25] is acceptable
     shape = np.asarray([29,29,29])
     # Number of microstructure instantiations to generate using DREAM.3D
     num instantiations = 5
     ''' Specify details of banding and sub-banding process '''
     # Specify the number of elements in each sub-band for volume averaging
     # Please see the references below for more information
     # WARNING!: If this is too low for very refined grains, this module will take a long time to run!
     # This is because it attempts to determine all UNIQUE combinations of some number of neighboring elements
     \sharp NOTE: Aim for ~8-10% of the average grain volume as the num_vox, as specified below
     \sharp This is especially important when comparing microstructures with different grain sizes! num_vox_percentage = 0.10
```

Fig. 8. Screenshot of the top of the main() function in generate microstructure.py.

```
| Anaconda Powershell Prompt (anaconda3)

(base) PS C:\Users\stopk\ cd C:\Users\stopk\Documents\GitHub\PRISMS-Fatigue
(base) PS C:\Users\stopk\Documents\GitHub\PRISMS-Fatigue> python .\generate_microstructures.py
Current output microstructure file: C:/Users/stopk/Documents/GitHub/test_ms_gen_2/Output_FakeMatl_0.vtk
Calculate realistic grain centroids

Working on grain 9
Working on grain 150
Working on grain 150
Working on grain 150
Working on grain 150
Sub-banding on grain 200
Amount of bands in total is 4529
Time to band microstructure: 0.22 seconds
Sub-banding grain: 9
Sub-banding grain: 100
Sub-banding grain: 150
Sub-banding grain: 150
Sub-banding grain: 200
Sub-banding grain: 200
Sub-banding grain: 200
Sub-banding complete
Sub-band generation with .inp file: 7.75 seconds
Total program: 20.38 seconds
(base) PS C:\Users\stopk\Documents\GitHub\PRISMS-Fatigue>
```

Fig. 9. Command prompt execution of the *generate microstructure.py* script.

```
- · X
 base) PS C:\Users\stopk\Documents\GitHub\PRISMS-Fatigue> ipythor
Python 3.7.6 (default, Jan 8 2020, 20:23:39) [MSC v.1916 64 bit (AMD64)]
Type 'copyright', 'credits' or 'license' for more information
IPython 7.12.0 -- An enhanced Interactive Python. Type '?' for help.
    [1]: import generate_microstructures as gen
          gen.main()
 urrent output microstructure file: C:/Users/stopk/Documents/GitHub/test_ms_gen_2/Output_FakeMatl_0.vtk
 Calculate realistic grain centroids
Working on grain 0
Working on grain 50
Working on grain 100
Working on grain 150
 Orking on grain 200
 mount of bands in total is 4743
Time to band microstructure: 0.23 seconds
 Sub-banding grain: 0
Sub-banding grain: 50
Sub-banding grain: 100
Sub-banding grain: 150
Sub-banding grain: 200
Sub-banding complete
 oub-band generation with .inp file: 7.53 seconds
Total program: 60.68 seconds
```

Fig. 10. Command prompt initiation of interactive Python session (i.e., "ipython") to execute Python scripts.

The main() function contains all the necessary parameters for the user to edit to generate microstructures, as well as descriptions of each variable. This script edits the .json pipeline based on user inputs for the size and shape of instantiations, the number of instantiations to generate, and the locations of the input, pipeline, and executable DREAM.3D files, and then calls DREAM.3D [1] as a subprocess. This and the other Python scripts contain many other functions that the user is encouraged to inspect for a more fundamental understanding of the scripts. A more detailed explanation for each variable is included in Table 1 below. Table 2 describes the generated files for each instantiation.

Table 1. List of variables and settings in the *generate_microstructures.py* script.

Variable/setting (type)

Description

Path to folder where all microstructure instantiation data should be stored.
This is referred to as a single "batch simulation folder." If this folder does not
exist, it will be created.
Users will note two ways to define the directory as shown in Fig. 8. The
default option uses the <i>DIR_LOC</i> variable which is the location of the folder
that contains the Python scripts and DREAM.3D files. The top of each script
defines this with the command below wherefile is the Python script file:
DIR_LOC = os.path.dirname(os.path.abspath(file))
NOTE : If this script is executed twice, the second instance of microstructure
instantiations will overwrite the first set (see last variable in this table)!
Location of the desired DREAM.3D "input" file, which contains two filters:
"StatsGenerator" and "Write DREAM.3D Data File."
This and the next variable can be specified in the manner above or by using
an absolute path to the file location.
Average grain size as calculated in the "StatsGenerator" filter in the .dream3d
input file. This is solely used for automatic band and sub-band sizing and does
NOT affect the instantiation of microstructures
Location of the DREAM.3D .json pipeline that reads in the microstructure
statistics defined by d3d_input_file and executes the pipeline.
Location of the DREAM.3D <i>PipelineRunner.exe</i> executable file (see Fig. 3)
required for Python during the subprocess call.
Size of the microstructure instantiation in millimeters in the X, Y, and Z
directions, respectively.
Number of voxels in the X, Y, and Z directions, respectively.
NOTE: At the time of this release, only CUBIC voxels are supported, so the
user must ensure that the size divided by shape in each of the X, Y, and Z
directions results in an identical voxel/element size!
Number of microstructure instantiations to generate in this batch simulation
folder. Each unique microstructure file contains the suffix "_#" where # is the
number of each instantiation and is indexed at 0 (see Fig. 12 and Table 2).
Desired percentage of grain volume that should be assigned to each sub-
band during automated calculation of sub-band volumes, i.e., how many
elements should be assigned to each sub-band. This is set to 10% by default.
This is especially important when comparing FIPs from microstructures with
different average grain sizes because the averaging volume must be
consistent!
Number of voxels in each sub-band region for FIP volume averaging. Please
see references [4-6] and the volume_average_FIPs.py section below for

	more information. In these references, the sub-band regions consist of 8
	elements which is ~8-10% of the grain volume.
	Users may specify this number manually or allow the default calculation
	using the "num_vox_percentage" variable as specified above.
band_thickness	Specifies the thickness of bands as a multiple of element width. This may be
integer	automatically calculated using the "avg_grain_size" specified above or users
	may specify their own values.
	NOTE: Users may need to iteratively generate microstructures and then
	visualize the bands using the "Output_FakeMatl_0_bands.vtk" file to fine
	tune this variable. The default setting will generate ~5-6 bands for each plane
	of each grain.
num_planes	Number of unique crystallographic slip planes in the material system under
integer	investigation. In the first PRISMS-Fatigue release, the face centered cubic
	(fcc) aluminum alloy Al 7075-T6 is simulated which has four crystallographic
	slip planes over which FIPs can be volume averaged.
create_sub_bands	Specify whether bands should be further designated into unique sub-band
boolean	volumes for the current microstructure instantiation(s). Default is "True".
	NOTE: Generating sub-bands may take excessively long for microstructures
	with a relatively high number of elements per grain (i.e., more than ~100
	elements/grain). This variable can be set to False and the subsequent
	microstructures will not underdo the sub-banding process. However, users
	can rerun this script with this variable set to True and the variable
	generate_new_microstructure_files set to False so that the already
	instantiated microstructures can be sub-banded. In this way, microstructures
	will not be overwritten.
	Additionally, users may not be interested in averaging FIPs over sub-bands
	because band or grain averaged values will suffice. Setting this variable to
	False will massively speed up the microstructure instantiation time.
compute kecher	Specify whether the grains that are split by the periodic microstructure
compute_kosher_ grain centroids	boundary (only for microstructures that are generated as fully-periodic in
° =	
boolean	DREAM.3D) should undergo additional preprocessing to more accurately
	determine grain centroids. This is important to more accurately average FIPs
	over grains, bands, or sub-bands. This can be set to False to speed up the
	microstructure instantiation process but will reduce the accuracy of FIP
	calculations at grains that are in contact with the microstructure boundary!
	Users should leave this as True but can change this to False if very large
	microstructures take much too long to process. Like the variable directly
	above, this can be initially set to False to generate multiple instantiations and
	then set to True with the variable generate_new_microstructure_files set to
	False so that the already instantiated microstructures can be more accurately

	preprocessed for FIP volume averaging. This variable is only relevant for the
	two latter face boundary conditions as described below.
face_bc	Defines the boundary conditions for microstructure periodicity in DREAM.3D
string array	and controls how FIP averaging volumes are determined. There are three
	options for users:
	1) Set all face boundary conditions to "free":
	 i. Microstructures are instantiated without periodicity in any direction. Afterwards, each grain is "banded" and then "subbanded" to determine FIP volume averaging regions for band-averaged and sub-band averaged FIPs, respectively. This is the simplest type of instantiation.
	2) Set all face boundary conditions to "periodic":
	 i. Microstructures are instantiated with periodicity in all three directions, i.e., grain tessellations that reach the boundary of the synthetic volume will "wrap around" and appear on the other side of the instantiation. These instantiations are simulated with periodic boundary conditions in PRISMS-Plasticity. However, additional processing is required before FIPs may be volume averaged. In the process of "banding" and "sub-banding" grains in a microstructure, the grain center of mass (COM) is required. Grains split by instantiation boundaries will have erroneous COM calculations based on their voxel locations. The Python script detects grains split by SVE boundaries and adjusts grain COMs for proper banding and sub-banding. 3) Set any two face boundary conditions to "periodic" and set the third to "free":
	 i. This boundary condition mimics a thin sheet of material [4-6]. Microstructures are instantiated with periodicity in all three directions. Grains split by one pair of parallel faces (determined by which of the face boundary conditions is set to "free") are detected and indexed as new grains. Grain periodicity is retained in the other two directions and so this emulates a thin sheet of material with a high surface area to
	volume ratio. <u>NOTE:</u> It is crucial that the subsequent CPFE simulations in PRISMS-Plasticity employ appropriate boundary conditions
	(i.e., with periodicity enforced along two directions)! DREAM.3D has the option to generate microstructures as either periodic or non-periodic. The third option described here provides users advanced

	control of subsequent FIP averaging. Fig. 11b depicts these three types of
	boundary conditions.
generate_new_	Specify whether new microstructures should be instantiated (True) or
microstructure_files	whether previous _#.csv and GrainID_#.txt files should be read to band and
Boolean	sub-band microstructures (False).

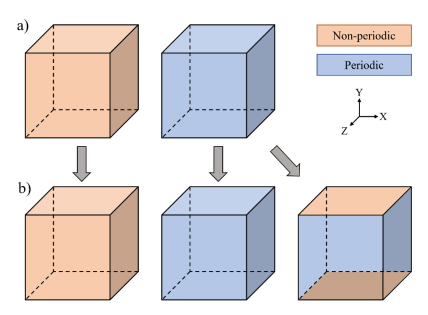


Fig. 11. Options for a) microstructure generation in DREAM.3D [1] and b) CPFEM simulation in PRISMS-Fatigue [2] in terms of periodicity. DREAM.3D instantiates synthetic microstructures by packing a volume with either non-periodic or periodic options. In the latter, grain tessellations that reach the boundary of a microstructure begin to grow from the opposite face. In b), the third option represents a "thin film" condition where one set of parallel faces (in this case, those perpendicular to the Y direction) can deform freely. The Python pre-processing module *generate_microstructures.py* in PRISMS-Fatigue detects and modifies the grains split by the non-periodic direction and indexes them as new grains for appropriate FIP volume averaging. Depiction of the three combinations of boundary conditions as described in Table 1.

Name	Date modified	Туре	Size
and_plane_normals_0.txt	5/27/2021 7:42 AM	TXT File	58 KB
and_sets_0.txt	5/27/2021 7:42 AM	TXT File	844 KB
Oream3D_microstructure_pipeline.json	5/27/2021 7:42 AM	JSON File	29 KB
🔁 El_pos_0.p	5/27/2021 7:42 AM	MATLAB P-code	572 KB
🔁 element_band_sets_0.p	5/27/2021 7:42 AM	MATLAB P-code	526 KB
🔁 element_grain_sets_0.p	5/27/2021 7:42 AM	MATLAB P-code	205 KB
elements_per_band_0.txt	5/27/2021 7:42 AM	TXT File	92 KB
FeatureData_FakeMatl_0.csv	5/27/2021 7:42 AM	Microsoft Excel C	96 KB
grainID_0.txt	5/27/2021 7:42 AM	TXT File	88 KB
microstructure_parameters.txt	5/27/2021 7:42 AM	TXT File	1 KB
orientations_0.txt	5/27/2021 7:42 AM	TXT File	9 KB
output.txt	5/27/2021 7:42 AM	TXT File	15 KB
Output_FakeMatl_0.dream3d	5/27/2021 7:42 AM	DREAM3D File	1,277 KB
Output_FakeMatl_0.vtk	5/27/2021 7:42 AM	VTK File	722 KB
Output_FakeMatl_0.xdmf	5/27/2021 7:42 AM	XDMF File	3 KB
Output_FakeMatl_0_bands.vtk	5/27/2021 7:43 AM	VTK File	923 KB
🔝 sub_band_info_0.p	5/27/2021 7:43 AM	MATLAB P-code	4,377 KB
sub_band_regions_0.inp	5/27/2021 7:43 AM	INP File	9,483 KB

Fig. 12. Screenshot of the files generated for each microstructure instantiation using the *generate_microstructure.py*. Each file is described in Table 2.

Table 2. List and description of files generated for each microstructure instantiation using *generate_microstructures.py*. The "#" indicates the instantiation number and is indexed at 0. Files with the ".p" extension indicate "pickle" files that are easily written and read by Python.

File name Description
band_plane_normals_#.txt For each grain, the direction normal

band_plane_normals_#.txt	For each grain, the direction normal to each of the four
	crystallographic slip planes.
band_sets_#.txt	Complete list of which elements belong to each band,
	indexed at 1.
Dream3D_microstructure_pipeline.json	A copy of the DREAM.3D pipeline executed by
	generate_microstructures.py. Certain filters are modified
	within Python (i.e., size and shape of instantiations) based
	on user setting in the main() function.
elements_per_band_#.txt	List of number of elements in each slip band for each grain,
	slip plane, and band layer number.
element_band_sets_#.p	List of elements in each band. Read by the
	volume_average_FIPs.py script to average FIPs over bands.
element_grain_sets_#.p	List of elements in each grain. Read by the
	volume_average_FIPs.py script to average FIPs over grains.
El_pos_0.p	List of each element's centroid. Read by the
	compile_and_plot_FIPs.py script to plot the highest FIPs as
	a function of their distance to the free-surface (only for

	simulations with one of three "face_bc" boundary condition
	set to "free" as described in Table 1). Only one instance is
	generated because element centroids are identical across
	different microstructures in a single folder.
FeatureData_FakeMatl_#.csv	Detailed list of grain information generated in DREAM.3D
	including centroids, volumes, number of voxels/elements,
	Euler angles, aspect ratios, etc.
grainID_#.txt	Specifies to which grain each element belongs. One of the
	two microstructure files required for CPFE simulation in
	PRISMS-Plasticity.
microstructure_parameters.txt	Text file that lists the settings in the main() function of
	generate_microstructures.py, e.g., size and shape of
	microstructure, which .dream3d input file was used, etc.
orientations_#.txt	Specifies grain orientations using Rodrigues vectors. The
	second microstructure file for CPFE simulation.
output.txt	Prints the DREAM.3D pipeline output to a text file for the
	last instantiation (see Region 6 in Fig. 4).
Output_FakeMatl_#.dream3d	Microstructure instantiation data stored in the ".dream3d"
	file format. This file is not used by subsequent Python
	scripts and is not required for CPFE simulations but is
	versatile and can be read by other .json DREAM.3D
	pipelines for further analysis.
Output_FakeMatl_#.xdmf	Generated alongside the ".dream3d" microstructure
	instantiation file in the "Write DREAM.3D Data File" filter. It
	allows the user to view the associated microstructure
	instantiation when opened in ParaView.
Output_FakeMatl_ # .vtk	This file also allows the user to view the microstructure
	instantiation in ParaView but the .json pipeline filter
	entitled "Vtk Rectilinear Grid Exporter" can be modified to
	export additional data for visualization.
Output_FakeMatl_#_bands.vtk	Additional .vtk file with visualization of grain bands.
sub_band_info_#.p	List of elements in each sub-band region. Read by the
	volume_average_FIPs.py script to average FIPs over sub-
	band regions.
sub_band_regions_#.inp	Complete list of which elements belong to each sub-band

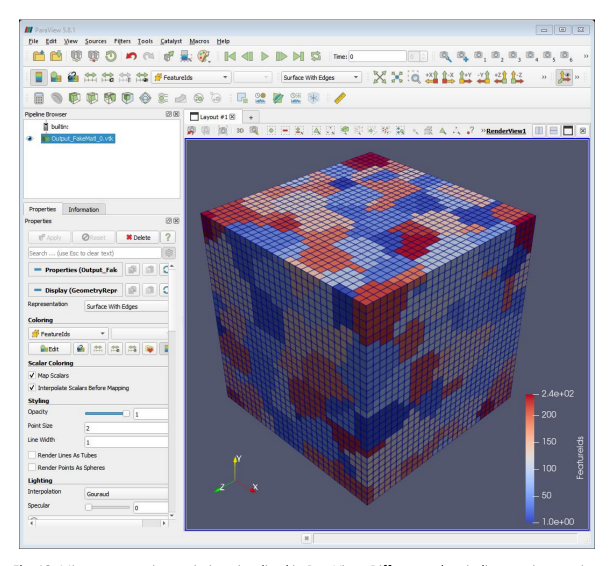


Fig. 13. Microstructure instantiation visualized in ParaView. Different colors indicate unique grains.

Crystal Plasticity Finite Element Method (CPFEM) Simulations

The files necessary to perform CPFEM simulations are available on the GitHub at this link: https://github.com/prisms-center/Fatigue/tree/main/src/MaterialModels. There are three .cc files specific to PRISMS-Fatigue that users must copy and paste into the PRISMS-Plasticity folders and then recompile PRISMS-Plasticity. More specifically, the three .cc files at the link above should replace the three .cc files with the same name in the PRISMS-Plasticity folder at: https://github.com/prisms-center/plasticity/tree/master/src/materialModels/crystalPlasticity, after which the package should be recompiled. This will ensure that the same quadrature outputs are written to the output .csv files and that users can arrive at identical results and figures.

Materials Commons

The entire dataset associated with the PRISMS-Fatigue manuscript is available for download at Materials Commons at this link: https://doi.org/10.13011/m3-rcyy-gx13. Users are encouraged to download these files to recreate the results and figures. There are four folders with microstructure and CPFE simulation results that correspond to the four applications described in the manuscript. Each folder contains the necessary prm_0.prm and other input files to perform the same CPFE simulations as described in the manuscript. The PRISMS-Fatigue videos describe this in detail.

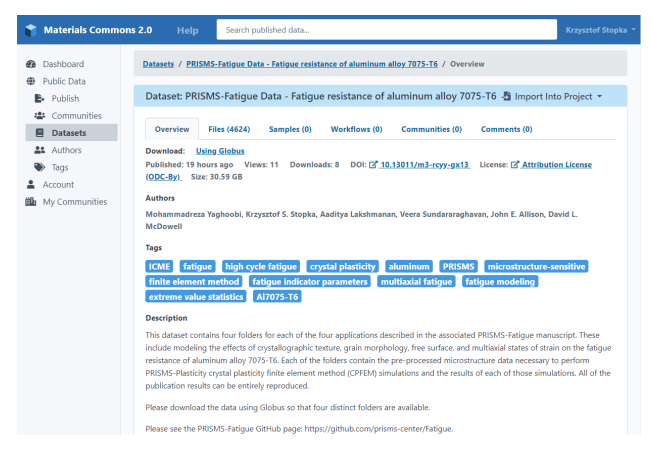


Fig. 14. Screenshot of PRISMS-Fatigue manuscript data available for public download.

calculate_FIPs.py

The second Python script calculates FIPs using simulation data exported from PRISMS-Plasticity. A screenshot of the main() function is shown in Fig. 15.

```
∃def main():
      # Directory where PRISMS-Plasticity results .csv files are stored
     # In the case of gamma plane simulations, this should contain all the folders numbered 0 thru (number of folder - 1),
     each of which contains some number of instantiations simulated at some combination of strain state and magnitude
     directory = os.path.dirname(DIR_LOC) + '\\tutorial\\MultiaxialFatigue_Al7075T6'
     r'C:\Users\stopk\Documents\GitHub\PRISMS-Fatigue\tutorial\Texture_Effect_A17075T6\rolled_equiaxed_periodic'
     # Shape of microstructure instantiations (at this point, only CUBIC voxel functionality supported)
     # THIS MUST MATCH THE SHAPE FROM THE 'generate microstructure.py' SCRIPT!
     shape = np.asarray([29,29,29])
     # Number of microstructure instantiations for which to calculate FIPs
     # THIS MUST MATCH THE SHAPE FROM THE 'generate_microstructure.py' SCRIPT! Otherwise, FIPs will not be computed for
     all instantiations!
     num_instantiations = 10
     # Specify the FIP to be calculated; default is "FS FIP"
     # The other FIP that can be calculated uses the plastic shear strain range on each slip system; FIP_type =
      'plastic_shear_strain_range'
     FIP_type = 'FS_FIP'
     # Define the two Fatemi-Socie (FS) FIP parameters; Please see the references below for more information
     k_fip = 10.0
     sigma_y = 517.0
     FIP_params = [k_fip, sigma_y]
     # For an fcc material, there are 12 slip systems for each element and therefore 12 FIPs per element
     num_slip_systems = 12
     # Specify whether this folder contain multiple instantiations to generate the multiaxial Gamma plane
     # This requires additional post-processing of PRISMS-Plasticity output files to calculate macroscopic plastic strain
     # Specify whether to append the .vtk file generated by DREAM.3D to visualize the highest FIP per element
     vtk visualize FIPs = False
      # Number of multiaxial strain state and magnitude folders, i.e., strain states simulated
      # This is only required if calculating the necessary files to generate a multiaxial gamma plane
     num_gamma_plane_folders = 1
     if gamma_plane_simulations:
          # Iterate through folders containing result files from multiaxial simulations
         for jj in range(num_gamma_plane_folders):
             dirr = os.path.join(directory, str(jj))
             print(dirr)
             for ii in range (num instantiations):
                 # print('Instantiation number %d' % ii)
                 import_PRISMS_data(dirr, shape, ii, FIP_type, num_slip_systems, FIP_params, gamma_plane_simulations)
          # Otherwise, compute FIPs for a single folder
         for ii in range (num instantiations):
             import_PRISMS_data(directory, shape, ii, FIP_type, num_slip_systems, FIP_params, gamma_plane simulations)
     if vtk visualize FIPs and not gamma plane simulations:
         for ii in range(num_instantiations):
             # Write FIPs to .vtk file for visualization
             append_FIPs_to_vtk(directory, shape, ii, FIP_type, num_slip_systems)
```

Fig. 15. Screenshot of the main() function in calculate FIPs.py.

As described in the PRISMS-Fatigue manuscript, a particular FIP is employed to demonstrate the capabilities of these scripts. However, prospective users can define their own FIPs which may require new and unique CPFE simulation outputs.

One of the output files of PRISMS-Plasticity CPFE simulations is the "Quadrature Output" as described in the fourth video tutorial (https://www.youtube.com/watch?v=YqaMhRLzqu0). It contains local (i.e., for each element integration point) user-defined state variables. Fig. 16 shows a sample file as used by PRISMS-Fatigue with a header row to distinguish each column. The columns include grain number, position in the X, Y, and Z directions, the current value of plastic shear strain for the 12 slip systems, and the stress normal to each slip system slip plane. These state variables are reported because of the desired FIP formulation. If the user's FIP formulation requires different state variables, then the "updateAfterIncrement.cc" file must be updated accordingly. These .csv files are read by this script using the "pandas" Python package.

Any FIP formulation must carefully consider the point at which FIPs are calculated. In the default FIP considered here (i.e., the crystallographic version of the Fatemi-Socie FIP), state variables at the points of maximum applied compression and tension across the final fully reversed loading/straining cycle are considered. Thus, the *calculate_FIPs.py* script searches for files entitled *MaxComp_#.csv* and *MaxTen_#.csv* from which state variables are read. Prospective users will heavily modify the import_PRISMS_data() function in this script to define unique FIP formulations. These must be saved to the *PRISMS_*FIP_type*_FIPs_#.p* file for subsequent FIP volume averaging. The following script requires that each element contain a FIP for every slip system. If a FIP formulation does not consider unique slip systems and instead provides a single scalar value at each element, this value can simply be repeated.

	sSave Off						x:Comp-0.csv *			∠ Search						7	A .	Stopka, Krzysztof S	SS EE -	
e	Home	Insert	Page	Layout Fr	ormulas Dat	ta Review	View Help	Acrobat											d Share □	Commer
1	Α			C	D	E	F	G	Н	1	J	K	L	M	N	0	P	Q	R	
8	grain_ID			У	Z	slip_1	slip_2	slip_3	slip_4	slip_5		slip_7	slip_8	slip_9		slip_11		normal_stress_1	_	_
H										4.60E-05									-1.14E	
L										3.69E-13									-4.50E	
L										5.18E-09									-7.82E	
L										3.32E-05									-1.38E	+02
	669	8 3.63	E-03	4.03E-04	4.03E-04	-3.24E-05	4.48E-04	-2.78E-05	2.79E-05	3.35E-05	-4.69E-04	-1.90E-72	2.67E-04	-2.87E-04	-2.89E-39	2.21E-05	-1.63E-05	-1.23E+02	-1.23E	+02
	669	8 4.43	E-03	4.03E-04	4.03E-04	-3.08E-05	4.58E-04	-3.00E-05	2.66E-05	3.64E-05	-4.92E-04	-4.04E-70	2.68E-04	-3.02E-04	-7.23E-36	2.44E-05	-1.80E-05	-1.29E+02	-1.29E	+02
L	669	8 5.24	E-03	4.03E-04	4.03E-04	-3.14E-05	4.84E-04	-3.10E-05	2.83E-05	3.76E-05	-5.39E-04	1.84E-55	2.65E-04	-3.21E-04	-1.61E-34	2.39E-05	-1.72E-05	-1.17E+02	-1.17E	+02
	669	8 6.04	E-03	4.03E-04	4.03E-04	-2.88E-05	4.23E-04	-2.98E-05	3.04E-05	3.47E-05	-5.27E-04	5.92E-35	2.25E-04	-3.29E-04	-2.85E-44	2.22E-05	-1.73E-05	-1.25E+02	-1.25E	+02
	669	8 6.85	E-03	4.03E-04	4.03E-04	-2.52E-05	4.04E-04	-3.22E-05	3.29E-05	3.20E-05	-5.24E-04	-4.88E-30	1.98E-04	-3.18E-04	-4.03E-91	1.91E-05	-1.92E-05	-1.25E+02	-1.25E	+02
	369	4 7.65	E-03	4.03E-04	4.03E-04	6.48E-05	1.13E-37	-4.56E-05	-3.18E-05	3.13E-05	6.39E-76	-1.47E-41	-1.57E-04	7.23E-05	3.86E-10	-4.66E-06	-2.82E-38	-3.68E+01	-3.68E	+01
	369	4 8.46	E-03	4.03E-04	4.03E-04	9.05E-05	-1.90E-22	-4.34E-05	-3.34E-05	3.10E-05	3.68E-67	5.16E-32	-1.83E-04	6.61E-05	-1.74E-14	-2.75E-06	-4.60E-39	-5.80E+01	-5.80E	+01
	369	4 9.26	E-03	4.03E-04	4.03E-04	1.50E-04	-1.21E-22	-4.73E-05	-3.12E-05	3.05E-05	3.21E-86	3.71E-25	-2.20E-04	7.81E-05	1.27E-06	-5.69E-06	1.02E-48	-5.59E+01	-5.59E	+01
ı	307	4 1.01	E-02	4.03E-04	4.03E-04	-4.23E-05	4.76E-05	-6.85E-42	1.92E-04	-1.38E-64	-2.14E-04	1.29E-12	1.74E-13	-1.05E-05	3.49E-52	-3.35E-05	2.96E-05	-5.49E+01	-5.49E	+01
Ι	307	4 1.09	E-02	4.03E-04	4.03E-04	-4.19E-05	4.73E-05	-2.23E-41	2.17E-04	-1.38E-64	-2.43E-04	1.30E-07	5.79E-14	-1.25E-05	3.00E-53	-3.31E-05	2.94E-05	-4.81E+01	-4.81E	+01
	307	4 1.17	E-02	4.03E-04	4.03E-04	-4.31E-05	4.69E-05	-4.38E-44	2.34E-04	-1.33E-62	-2.60E-04	1.51E-06	3.98E-16	-1.32E-05	4.80E-66	-3.34E-05	3.13E-05	-4.15E+01	-4.15E	+01
ı	307	4 1.25	E-02	4.03E-04	4.03E-04	-4.13E-05	4.65E-05	-1.05E-41	1.98E-04	-5.37E-60	-2.38E-04	5.59E-07	8.22E-24	-9.78E-06	7.55E-64	-3.47E-05	3.21E-05	-4.38E+01	-4.38E	+01
ı	307	4 1.33	E-02	4.03E-04	4.03E-04	-4.10E-05	4.77E-05	-4.21E-34	1.72E-04	1.95E-44	-2.50E-04	1.10E-07	8.75E-28	-8.13E-06	7.90E-68	-3.47E-05	3.30E-05	-3.47E+01	-3.47E	+01
ı	134	9 1.41	E-02	4.03E-04	4.03E-04	1.05E-05	1.32E-59	-1.36E-05	-4.02E-04	2.07E-04	1.13E-14	-9.22E-08	-4.15E-05	1.50E-04	-3.14E-05	3.69E-07	1.55E-05	2.56E+00	2.56E	+00
ı	134	9 1.49	E-02	4.03E-04	4.03E-04	1.28E-05	2.77E-32	-2.00E-05	-3.81E-04	2.41E-04	3.20E-25	-6.13E-07	-4.15E-05	1.60E-04	-2.71E-05	1.38E-10	1.31E-05	-6.81E+00	-6.81E	+00
l	134	9 1.57	E-02	4.03E-04	4.03E-04	1.75E-05	3.84E-40	-2.31E-05	-3.60E-04	2.32E-04	3.12E-28	-4.13E-11	-4.17E-05	1.27E-04	-2.34E-05	7.02E-09	8.71E-06	-2.53E+00	-2.53E	+00
t	134	9 1.65	E-02	4.03E-04	4.03E-04	1.87E-05	-4.27E-54	-2.26E-05	-3.41E-04	1.69E-04	9.00E-19	-4.89E-09	-4.38E-05	1.75E-04	-2.00E-05	3.75E-20	9.58E-06	-1.72E+01	-1.72E	+01
t	134	9 1.73	E-02	4.03E-04	4.03E-04	1.68E-05	1.16E-54	-2.03E-05	-3.07E-04	1.50E-04	1.17E-21	-1.44E-11	-4.41E-05	1.61E-04	-2.10E-05	5.86E-24	1.18E-05	-2.12E+01	-2.12E	+01
t	721	0 1.81	E-02	4.03E-04	4.03E-04	2.77E-05	2.12E-35	-3.42E-05	-2.64E-04	3.03E-04	1.91E-67	9.18E-45	-3.01E-05	3.41E-05	-1.68E-05	1.45E-05	1.62E-68	-3.02E+01	-3.02E	+01
t	721	0 1.89	E-02	4.03F-04	4.03E-04	2.58F-05	1.41F-33	-3.27E-05	-2.84E-04	2.89E-04	********	-9.03E-35	-2.75E-05	3.41F-05	-1.75E-05	1.57E-05	-4.21F-73	-3.26E+01	-3.26E	+01
t										2.70E-04									-2.19E	
t										2.56E-04									-3.36E	
H										3.08E-04									-6.35E	
ı										6.11E-04									-5.85E	
l										6.18E-04									-6.18E	
H										5.74E-04									-7.98E	
H										-2.83F-05									-6.30F	

Fig. 16. Sample PRISMS-Plasticity Quadrature output file. The header row was added manually and is not required by *calculate_FIPS.py*. Values are instead read in by column number.

Table 3. List of variables and settings in the *calculate_FIPs.py* script.

Variable/setting (type)

Description

directory	Path to folder with microstructure data and simulation results. Like the
Path to folder	previous script, this can be defined with a relative or absolute path.
shape	Number of voxels in the X, Y, and Z directions, respectively.
integer array	
num_instantiations	Number of microstructure instantiations for which FIPs should be
integer	calculated. This should match the number specified in
	generate_microstructure.py.
FIP_type	The type of FIP that should be computed for each voxel/element using
string	the CPFE simulation output data. The default is "FS_FIP" which calculates
	a crystallographic version of the Fatemi-Socie FIP. Another option
	included in the initial release of PRISMS-Fatigue is
	"plastic_shear_strain_range" that considers solely the plastic shear
	strain range on each slip system at every element.
	This script will be updated with other types of FIPs in the future. The user
	is encouraged to define their own FIPs.
k_fip	Parameter to calculate "FS_FIP"; controls the influence of the stress
float	normal to the slip plane on the FIP magnitude.
sigma_y	Parameter to calculate "FS_FIP"; yield strength of the material system
float	modeled; normalizes the influence of the stress normal to the slip plane.
FIP_params	FIP parameters compiled into a list for more convenient function calls.
num_slip_systems	Number of slip systems in the material of interest. The FS_FIP is
integer	calculated for each slip system but other FIP definitions may not require
	this or the previous two variables (i.e., k_fip and sigma_y).
gamma_plane_simulations	Specifies whether the FIPs to be calculated will be used for a multiaxial
Boolean	gamma plane [6] (explained further in the gamma_plane.py section).
vtk_visualize_FIPs	Specifies whether the highest elemental FIP per element should be
Boolean	appended to the .vtk file for visualization of FIPs in ParaView.
num_gamma_plane_folders	Number of unique simulation folders to populate the multiaxial gamma
integer	plane.

Table 4. List and description of files generated for each microstructure by *calculate_FIPs.py*. The "#" indicates the instantiation number and is indexed at 0. Files with the ".p" extension indicate "pickle" files that are easily written and read by Python.

File name	Description
PRISMS_*FIP_type*_FIPs_#.p	List of FIPs for each element. Read by the volume_average_FIPs.py
	script to average FIPs over grains, bands, or sub-band regions.

volume average FIPs.py

The third Python script averages FIPs over one of three types of volumes: grains, bands, or subband regions. Fig. 17 depicts each type of volume. A screenshot of the main() function is shown in Fig. 18 and each user setting is described in Table 5.

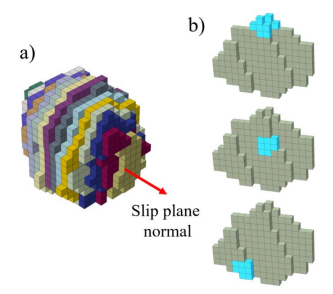


Fig. 17. (a) A single grain with slip bands representative of crystallographic slip planes. (b) A single slip band from the grain in (a) in which different sub-band regions comprised of eight elements each (controlled by the variable *num_vox* in *generate_microstructures.py*) are highlighted.

```
□def main():
     # Directory where microstructure FIP .p files are stored:
directory = os.path.dirname(DIR_LOC) + '\\tutorial\\MultiaxialFatigue_A17075T6'
     # directory = r'C:\Users\stopk\Documents\GitHub\PRISMS-Fatique\tutorial\test run 1'
     # Number of microstructure instantiations in each folder for which to volume average FIPs
     num instantiations = 10
     # Specify the FIP to be volume averaged
     FIP_type = 'FS_FIP'
     # Specify type of FIP averaging: 'sub_band', 'band', or 'grain'
     averaging_type = 'sub_band'
     # Specify whether this folder contain multiple instantiations to generate the multiaxial
     Gamma plane
     gamma_plane_simulations = False
     # Number of multiaxial strain state and magnitude folders, i.e., strain states simulated
     # This is only required if calculating necessary files to generate multiaxial gamma plane
     num_gamma_plane_folders = 1
     if gamma_plane_simulations:
         # Iterate through folders containing result files from multiaxial simulations
         for jj in range(num_gamma_plane_folders):
             dirr = os.path.join(directory, str(jj))
             print(dirr)
             for ii in range(num instantiations):
                 call_averaging(ii, dirr, FIP_type, averaging_type, gamma_plane_simulations)
         # Call function to perform averaging for each instantiation in folder
         for ii in range(num_instantiations):
             call_averaging(ii, directory, FIP_type, averaging_type, gamma_plane_simulations)
```

Fig. 18. Screenshot of the main() function in *volume_average_FIPs.py*.

Table 5. List of variables and settings in the *volume_average_FIPs.py* script.

Variable/setting (type) Description

directory	Path to folder where all microstructure instantiation data and simulation
path to folder	results are stored. Like the previous script, this can be defined with a
	relative or absolute path.
num_instantiations	Number of microstructure instantiations for which FIPs should be
integer	volume averaged.
FIP_type	The type of FIP that should be volume averaged.
string	NOTE: The desired FIP that should be volume averaged should have
	been calculated using the previous script!
averaging_type	Desired type of FIP volume averaging. Options include "sub_band",
string	"band", and "grain".
	NOTE: This script will fail if this is set to "sub_band" but microstructures
	have not undergone the sub-banding process.
gamma_plane_simulations	Specifies whether the FIPs to be calculated will be used for a multiaxial
Boolean	gamma plane [6] (explained further in the gamma_plane.py section).
num_gamma_plane_folders	Number of unique simulation folders to populate the multiaxial gamma
integer	plane.

The "averaging_type" specified by the user determines which of the three pairs of files listed in Table 6 will be generated. The pickle files are read by subsequent scripts whereas the .csv files allow the user to quickly read through the averaged FIPs. Microstructures with a very large number of elements may produce excessively large files. The user may modify the scripts to suppress generation of certain files (e.g., .csv files) to reduce memory usage. The user may also run this script three times and average FIPs over all three types of volumes.

Table 6. List and description of files generated for each microstructure by *volume_average_FIPs.py*. The "#" indicates the instantiation number and is indexed at 0. Files with the ".p" extension indicate "pickle" files that are easily written and read inside Python.

File name Description

sub_band_averaged_*FIP_type*_pickle_#.p	List of sub-band averaged FIPs. Read by
	compile_and_plot_FIPs.py and gamma_plane.py.
sub_band_averaged_*FIP_type*_#.csv	List of sub-band averaged FIPs written to readable .csv
	file.
band_averaged_*FIP_type*_pickle_#.p	List of band averaged FIPs. Read by
	compile_and_plot_FIPs.py.
band_averaged_*FIP_type*_#.csv	List of band averaged FIPs written to readable .csv file.
grain_averaged_*FIP_type*_pickle_#.p	List of grain averaged FIPs. Read by
	compile_and_plot_FIPs.py.
grain_averaged_*FIP_type*_#.csv	List of grain averaged FIPs written to readable .csv file.

compile and plot FIPs.py

The fourth Python script compiles and plots FIPs from different simulation batch folders. Two types of figures are supported. The first fits the highest FIPs to an EVD. The second plots FIPs as a function of their distance to the microstructure boundary (i.e., to the free surface for "thin sheet" simulations as described previously). Fig. 19 depicts a screenshot of the variables in the main() function which are described in Table 7.

```
□def main():
     # Specify type of analysis: either 'fip_evd' to plot FIP extreme value distributions (EVDs)
     or 'surf dist' to plot the highest FIPs as a function of their distance to the free surface
     analysis type = 'fip evd'
     # Specify folder which contains all of the simulation batch folders
     # IMPORTANT: this directory should contain the locations specified at the top of this python
     script, NOT the individual folders with the instantiations!
     # This script goes through EACH ONE of the folders specified and extracts the relevant
     information for plotting purposes
     directory = os.path.dirname(DIR LOC) + '\\tutorial'
     # directory = r'C:\Users\stopk\Documents\GitHub\PRISMS-Fatigue\tutorial\test_run_1'
     # Specify which 'material' to plot, i.e., which combination of microstructure folders
     # Please see the top of the "plot_FIPS" function
     mat = 'prisms compare bcs'
     # Specify which FIP to import
     FIP_type = 'FS_FIP'
     # Specify which volume averaging scheme to import. By default, the sub-band averaged (SBA)
     FIPs are used
     # IMPORTANT: this will fail if the FIPs have not yet been averaged in the desired manner
     (see 'volume_average_FIPs.py' script)
     # Options: 'sub_band', 'band', and 'grain'
     averaging_type = 'sub_band'
     if analysis_type == 'fip_evd':
         # Specify how many of the highest FIPs should be plotted
         n_{j} = 50
         # Specify whether FIPs should be fit to the "gumbel" or "frechet" EVD
         plt_type = 'gumbel'
         # Call function to plot FIPs
         plot_FIPS(directory, plt_type, mat, n_fip_plot, FIP_type, averaging_type, save_fig = True)
     elif analysis_type == 'surf_dist':
         # Plot SBA FIP vs distance from free surface:
         # Specify how many of the highest FIPs should be plotted
         n_fip_plot = 10
         # Specify the non-periodic direction (i.e., the set of parallel faces set to
         traction-free conditions)
         # In the references associated with these scripts (see below), this is set to the 'Y'
         direction
         non_periodic_dir = 'Y'
         # Specify the size/length of the SVE in the non-periodic direction in mm, to properly
         locate the FIPs within the microstructure
         SVE non periodic length = 0.0725
         # Call function to plot FIPs
         plot EVD FIP distance (directory, mat, n fip plot, FIP type, averaging type,
         non_periodic_dir, SVE_non_periodic_length)
```

Fig. 19. Screenshot of the main() function in *compile_and_plot_FIPs.py*.

Table 7. List of variables and settings in the main() function in *compile_and_plot_FIPs.py*.

Variable/setting (type)

Description

(0) 0	
analysis_type	Type of plot to generate. Options include "fip_evd" and "surf_dist" and
string	correspond to FIP EVDs and FIPs plotted as a function of distance to the
	microstructure boundary, respectively.
directory	NOTE: This directory should be different than previously defined
path to folder	directories and will store the resultant plots and other generated files.
	The previous scripts defined the directory as the location with
	microstructure and simulation data. In contrast, this script will read FIP
	data from locations specified at the top of the script, as show in Fig. 20.
mat	This specifies additional settings in the plot_FIPs() and
string	plot_EVD_FIP_distance() functions including figure legend labels, marker
	shapes and colors, number of columns in the figure legend, etc., and where
	all FIP data is stored ("locs" variable).
FIP_type	The type of FIP that should be plotted.
string	NOTE: An error will occur if the user specifies a FIP for which volume
	averaged files do not exist.
averaging_type	The type of FIP that should be read, compiled, and used in figures. Options
string	include "sub_band", "band", and "grain".
	NOTE: An error will occur if the user has not averaged FIPs in the previous
	script in the same manner as desired by this script.
n_fip_plot	Number of FIPs to plot for either type of figure. The default is 50 and 10 for
integer	FIP EVDs and FIP vs. distance to microstructure boundary figures,
	respectively.
plt_type	Specifies whether FIPs should be fit to the Gumbel or Fréchet EVD with
string	options as "gumbel" and "frechet", respectively.
non_periodic_dir	Specifies which pair of microstructure instantiation faces are non-periodic
string	(i.e., free surfaces as set in generate_microstructure.py; option 3 for
	"face_bc").
SVE_non_periodic_length	The length of the microstructure in the non-periodic direction. Required to
float	correctly calculate distance of the highest FIPs to the free surface or
	instantiation boundary.

```
| Cos.path.dirname(DIR_LOC) + '\tutorial\\Texture_Effect_Al7075T6\\cubic_equiaxed_periodic',
| os.path.dirname(DIR_LOC) + '\tutorial\\Texture_Effect_Al7075T6\\cubic_equiaxed_free_surface',
| os.path.dirname(DIR_LOC) + '\tutorial\\Texture_Effect_Al7075T6\\random_equiaxed_periodic',
| os.path.dirname(DIR_LOC) + '\tutorial\\Texture_Effect_Al7075T6\\random_equiaxed_free_surface',
| os.path.dirname(DIR_LOC) + '\tutorial\\Texture_Effect_Al7075T6\\rolled_equiaxed_periodic',
| os.path.dirname(DIR_LOC) + '\tutorial\\Texture_Effect_Al7075T6\\rolled_equiaxed_free_surface']

| Dicats_Al7075_compare_shape_prisms = [os.path.dirname(DIR_LOC) + '\tutorial\\Texture_Effect_Al7075T6\\roubic_equiaxed_periodic',
| os.path.dirname(DIR_LOC) + '\tutorial\\Texture_Effect_Al7075T6\\roubic_equiaxed_periodic',
| os.path.dirname(DIR_LOC) + '\tutorial\\Texture_Effect_Al7075T6\\random_equiaxed_periodic',
| os.path.dirname(DIR_LOC) + '\tutorial\\Texture_Effect_Al7075T6\\rolled_equiaxed_periodic',
| os.path.dirname(DIR_LOC) + '\tutorial\\Te
```

Fig. 20. Screenshot of the top of *compile_and_plot_FIPs.py* where the paths of individual simulation folder are specified.

This script is unique in that users must also modify another function entitled plot_FIPs() and/or plot_EVD_FIP_distance(). The "mat" variable specified by users in the main() function determines which combination of individual simulation batch folders to plot. As an example, the screenshot in Fig. 21 shows two instances of "mat" that plot and compare FIP EVDs that vary in boundary condition (free surface vs. fully periodic CPFE microstructure and boundary conditions, mat = "prisms_compare_bcs") and grain morphology (equiaxed vs. elongated, mat = "prisms_compare_shape") as described in the PRISMS-Fatigue manuscript. The variable assignments under a specific "mat" definition control figure properties (e.g., name in legend, marker styles, etc.) and the location of each folder as indicated in Fig. 20.

At this point, the "surf_dist" option only supports sub-band averaged FIPs. This is because attempting to locate the grains with the highest grain averaged FIPs with respect to the microstructure boundaries may too excessively smear interpretation of this plot. More specifically, it makes more sense to locate the sub-band region with the highest FIP with respect to the microstructure boundary because it more accurately reflects the fact that fatigue crack formation occurs at the sub-grain scale, and not across an entire grain instantaneously. Even the grain which contains the highest grain averaged FIP may have regions which undergo little local plastic deformation.

```
def plot_FIPS(base_directory, plt_type, mat, num_fips_plot, fip_type, save_fig = True):
     # Function to plot FIPs from bulk (i.e., fully periodic) and surface (i.e., traction-free/free
     surface) simulations, or other types of simulations
     print('Plotting volume averaged FIPs')
     os.chdir(base_directory)
      # Specify which FIPS to plot
     # This will plot FIPs from subsurface/bulk and free-surface simulations
     if mat == 'prisms_compare_bcs':
          # Names to use for figure legend
         # These must match the order of simulation folders specified at the top of this script
         names = ["Cubic Periodic", "Cubic Free Surface", "Random Periodic", "Random Free Surface", "Rolled
         Periodic". "Rolled Free Surface"1
         # Locations of actual FIP files
         locs = locats Al7075 compare BC prisms
         # Number of columns for figure legend
         plot col = 1
         # Marker colors
         cfm = ['r','r','b','b','g','g']
         # Marker shapes
         sfm = ['0','0','s','s','^','^']
         # If set to true, FIPs from periodic and free surface simulations will be hollow and full,
         hollow and full = True
     # This will plot FIPs from fully periodic simulations with different crystallographic textures and
     grain morphologies
     elif mat == 'prisms compare shape':
         # Names to use for figure legend
         names = ["Cubic Equiaxed","Cubic Elongated","Random Equiaxed","Random Elongated","Rolled Equiaxed",
         "Rolled Elongated"]
         # Locations of actual FIP files
         locs = locats A17075 compare shape prisms
         # Number of columns for figure legend
         plot_col = 1
         # Marker colors
         cfm = ['r','r','b','b','g','g']
         # Marker shapes
         sfm = ['0','0','s','s','^','^']
         # If set to true, FIPs from periodic and free surface simulations will be hollow and full,
         respectively
         hollow_and_full = True
```

Fig. 21. Screenshot of the top of the plot FIPs() function in compile and plot FIPs.py.

Several files are generated by this script, as shown in Fig. 22 and described in Table 8. The pickle ".p" files are generated to increase efficiency. Each folder may contain dozens of microstructures and extracting FIPs from each volume averaged pickle file may take considerable time, especially for those with many elements and for the more complex sub-band averaged FIPs. Therefore, the highest FIPs per grain for all microstructures in a single folder are stored into a single array which is then stored in the compiled_*.p files shown in this folder. The script first checks whether these files already exist and if this is not true, it generates them by reading the appropriate FIP files. Once the files are present, they are quickly read to produce plots. This allows the user to rapidly change figure properties and spend more time interpreting data.

For the first type of plot, users can specify whether FIPs should be fit to the Gumbel or Fréchet EVD and plots are saved with corresponding file names. A .csv file contains properties of the best fit line for each plot along with the value of the largest FIP. By default, the highest 50 FIPs from each data set are fit to the EVD. The second type of figure by default plots the 10 highest FIPs as a function of their distance to the instantiation boundary. An associated text file contains the average distance of these FIPs to the instantiation boundary.

Table 8. List and description of files generated by *compile_and_plot_FIPs.py*.

Description File name compiled_*FIP_type*_*averaging_type* List of compiled FIPs for each simulation folder. *folder*.p compiled_surf_dist_*FIP_type* List of compiled FIPs and distance to microstructure _*averaging_type**folder*.p boundary for each simulation folder. r squared *plt type* *FIP type* Line of best fit properties of FIPs fit to EVD. *averaging_type*_* n_fip_plot *_*mat*.csv *FIP_type*_EVD_* averaging_type * Plot of FIPs fit to either Gumbel or Fréchet EVD. _*plt_type*_*n_fip_plot*_*mat*.png Avg_surf_dist_for_top_*n_fip_plot*_*mat.txt Average distance of FIPs to microstructure boundary. SBA_FIP_surface_dist_*n_fip_plot*_*mat*.png Plot of FIPs vs. distance to microstructure boundary.

🔝 compiled_FS_FIP_grain_random_elongated_periodic.p
🔝 compiled_FS_FIP_grain_random_equiaxed_periodic.p
🔝 compiled_FS_FIP_grain_rolled_elongated_periodic.p
🔝 compiled_FS_FIP_grain_rolled_equiaxed_periodic.p
🔝 compiled_FS_FIP_sub_band_cubic_elongated_periodic.p
🔝 compiled_FS_FIP_sub_band_cubic_equiaxed_free_surface.p
🔝 compiled_FS_FIP_sub_band_cubic_equiaxed_periodic.p
🔝 compiled_FS_FIP_sub_band_random_elongated_periodic.p
🔝 compiled_FS_FIP_sub_band_random_equiaxed_free_surface.p
🔝 compiled_FS_FIP_sub_band_random_equiaxed_periodic.p
🔝 compiled_FS_FIP_sub_band_rolled_elongated_periodic.p
🔝 compiled_FS_FIP_sub_band_rolled_equiaxed_free_surface.p
🔝 compiled_FS_FIP_sub_band_rolled_equiaxed_periodic.p
🔝 compiled_surf_dist_FS_FIP_sub_band_cubic_equiaxed_free_surface.p
🔝 compiled_surf_dist_FS_FIP_sub_band_cubic_equiaxed_periodic.p
🔝 compiled_surf_dist_FS_FIP_sub_band_random_equiaxed_free_surface.p
🔝 compiled_surf_dist_FS_FIP_sub_band_random_equiaxed_periodic.p
🔝 compiled_surf_dist_FS_FIP_sub_band_rolled_equiaxed_free_surface.p
🔝 compiled_surf_dist_FS_FIP_sub_band_rolled_equiaxed_periodic.p
FS_FIP_EVD_band_gumbel_50_prisms_compare_shape.png
FS_FIP_EVD_grain_gumbel_50_prisms_compare_shape.png
FS_FIP_EVD_sub_band_gumbel_50_prisms_compare_bcs.png
FS_FIP_EVD_sub_band_gumbel_50_prisms_compare_shape.png
r_squared_gumbel_FS_FIP_band_50_prisms_compare_shape.csv
r_squared_gumbel_FS_FIP_grain_50_prisms_compare_shape.csv
r_squared_gumbel_FS_FIP_sub_band_50_prisms_compare_bcs.csv
r_squared_gumbel_FS_FIP_sub_band_50_prisms_compare_shape.csv
SBA FIP surface dist 10 prisms compare bcs.png

Fig. 22. Folder structure to execute compile and plot FIPs.py and output files. See Fig. 20.

gamma plane.py

The fifth and final Python script creates a multiaxial gamma (Γ) plane. It requires a single, large microstructure or a set of microstructures simulated under multiple combinations of strain states (e.g., uniaxial, biaxial, and pure shear) and magnitudes [6]. A screenshot of the main() function is shown in Fig. 23 and Table 9 describes each input variable.

```
□def main():
     # Specify name of directory with simulation folders, each with the same X number of
     microstructures simulated to some combination of strain state and magnitude, and
     numbered 0 thru (number of folder - 1)
     directory = os.path.dirname(DIR_LOC) + '\\tutorial\\MultiaxialFatigue A17075T6'
     # Specify the number of FIPs to extract from each simulation folder
     num_FIPs_extract = 50
     # Specify how many of the highest FIPs from each batch of simulations should be
     considered to plot the ISO-FIP contours
     num_read_top_FIPs = 10
     # Specify which FIP to import
     FIP_type = 'FS_FIP'
     # Specify type of FIP averaging: 'sub band', 'band', or 'grain'
     averaging_type = 'sub_band'
     # Plot the locations of each simulation batch folder in terms of only the response
     coordinates (x and y position on the gamma plane)
     plot gamma plane locations (directory)
     # This function will extract the 50 highest fips from each folder and store these in
     a single pickle file.
     # This function is run once to speed up subsequent analysis and plotting
     get_gamma_FIPs(directory, num_FIPs_extract, FIP_type, averaging_type)
     # Plot the gamma plane
     plot_gamma(directory, num_read_top_FIPs, FIP_type, averaging_type)
```

Fig. 23. Screenshot of the main() function in gamma_plane.py.

Table 9. List of variables and settings in *gamma plane.py*.

Variable/setting Description

directory	Path to directory which contains folders numbered 0 thru (number of
path to folder	total simulated strain states and magnitudes). Each of these numbered
	folders must contain pairs of result .csv files as described previously.
num_read_top_FIPs	The amount of the highest volume averaged FIPs that should be used to
integer	calculate the ISO-FIP contours for the gamma plane (i.e., the number of
	FIPs that should be averaged and used in the Gaussian Process
	Regression (GPR) model to interpolate across the Gamma plane).
FIP_type	The type of FIP that should be considered.
string	NOTE: An error will occur if the user specifies a FIP for which volume
	averaged files do not exist.
averaging_type	The type of FIP that should be read, compiled, and used in figures.
string	NOTE: An error will occur if the user has not averaged FIPs in the
	previous script in the same manner as desired by this script.

The steps required to generate the gamma plane are as follows:

- 1) Generate a single large microstructure or an ensemble of microstructures using the *generate_microstructures.py* script.
- 2) Simulate these at various combinations of strain states and magnitudes.
 - a. The download from Materials Commons includes an excel sheet that lists all the combinations of strain states and magnitudes that were used to populate the Gamma plane presented in the PRISMS-Fatigue manuscript. This download also includes the necessary prm_0.prm and microstructure file to apply the appropriate boundary/simulation conditions to the 10 microstructures.
- 3) Calculate FIPs for each microstructure in each simulation folder numbered 0 thru (number of simulated strain states) using the *calculate FIPs.py* script.
- 4) Volume average FIPs over the desired volume using the volume_average_FIPs.py script.
- 5) Execute the *gamma_plane.py* script.

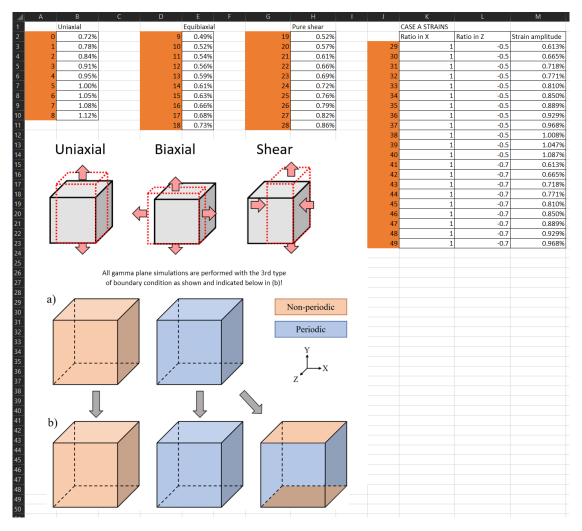


Fig. 24. Screenshot of the excel sheet included in the Materials Commons dataset that specifies each combination of strain state and magnitude used to generate the Gamma plane for Al7075-T6.

Sample tutorial

This playlist: https://www.youtube.com/playlist?list=PL4yBCojM4Swo3CvlA57syFrzk3p1mugP5 guides users through the entire framework to reproduce the results and figures in the PRISMS-Fatigue manuscript.

Useful links:

GitHub Repository: https://github.com/prisms-center/Fatigue

Materials Commons Dataset: https://doi.org/10.13011/m3-rcyy-gx13

PRISMS-Plasticity Forum: https://groups.google.com/g/prisms-cpfe-users

PRISMS Center: http://www.prisms-center.org/#/home

Support/current developers:

Krzysztof S. Stopka: stopka:stopka:gmail.com, kstopka:gmail.com, <a href="mailto:kstopka:kstopka:gstopka:kstopka:gstopka:gstopka:kstopka:gstopka:kstopka:gstopka:kstopka:gstopka:kstopka:gstopka:kstopka:gstopka:kstopka:gstopka:kstopka:gstopka:kstopka:gstopka:kstopka:gstopka:kstopka:gstopka:kstopka:gstopka:kstopka:gstopka:kstopka:gstopka:kstopka:gstopka:kstopka:gstopka:kstopka:gstopka:kstopka:kstopka:gstopka:kstopka:gstopka:kstopka:gstopka:kstopka:gstopka:kstopka:kstopka:gstopka:k

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