# SAnTex - Seismic Anisotropy from Texture: A Python-based library for Seismic Anisotropy Calculation

# **User Guide**

Utpal Singh<sup>1</sup>, Sinan Özaydın<sup>1</sup>, Vasileios Chatzaras<sup>1</sup>, Patrice Rey<sup>1</sup>

<sup>1</sup>The University of Sydney, School of Geosciences, Sydney, NSW, Australia

# **SAnTex Installation**

Open the terminal and go to Downloads or Documents directory by typing *cd* ~/Documents or *cd* ~/Downloads or any other directory, where you can store the repository of SAnTex. Run the following to get the updated version of SAnTex

```
git clone https://github.com/utpal-singh/SAnTex.git
cd santex
pip install .
```

# 1.0 Working with SAnTex

In a python working environment, import SAnTex's modules and Python libraries:

```
from santex import EBSD
from santex import Tensor
from santex import Material
from santex import Isotropy
import pandas as pd
```

NB: In what follows, user-defined names are in blue, e.g., ebsd.ctf

#### Load the EBSD file:

```
ebsdfile = EBSD("ebsd.ctf")
```

\*Note: Remember to change the parentheses in the python code, the Word document modifies the parentheses.

This loads the *ebsd.ctf* file in the python object *ebsdfile* of class *EBSD*, from which users are able to call methods for further processing, cleaning of the ebsd data, and ultimately calculate elastic properties.

\*Class: In Python, a class is a blueprint for creating objects. It defines the structure and behavior of objects of that type. Think of it as a template or a prototype that encapsulates data (attributes) and methods (functions) that operate on that data. Object: An object, also known as an instance, is a unique entity that is created based on the structure defined by its class. When an object is instantiated, it inherits the attributes and methods defined in its class, but can also have its own unique state and behavior.

Note: the *ebsd.ctf* should be located in the directory this notebook is being run from. But you may provide a relative path like "*data/ebsd.ctf*" or an absolute import such as "/*Users/myname/Documents/ebsd/ebsd.ctf*"

#### 1.1 Available methods in the EBSD class

To list the phases present in the ebsd file:

```
phases_available = ebsdfile.phases()
print(phases available)
```

Load the ebsd file into a pandas dataframe, via invoking the get\_ebsd\_data() method from the EBSD class:

```
df = ebsdfile.get_ebsd_data()
print(df)
```

## 2.6 Plotting Conventions

Following are the keywords to orient sample reference plane and to store in a new dataframe

```
The default is sample_ref = ["x2east", "zOutOfPlane"]
```

### Following are other available keywords:

```
sample_ref == ["x2east", "zOutOfPlane"]
sample_ref == ["x2west", "zOutOfPlane"]
sample_ref == ["x2north", "zOutOfPlane"]
sample_ref == ["x2south", "zOutOfPlane"]
sample_ref == ["x2east", "zIntoPlane"]
```

```
sample_ref == ["x2west", "zIntoPlane"]
sample_ref == ["x2north", "zIntoPlane"]
sample_ref == ["x2south", "zIntoPlane"]

rotated_df = ebsdfile.plot_rotate_ebsd(sample_ref = ["x2east", "zIntoPlane"], ebsd_df = df)
ebsdfile.plot(rotated_df)
```

To rotate the EBSD data to match the SEM orientation in any custom angles

```
angles = (180, 0, 0)
updatedebsd = ebsdfile.rotateEBSD(ebsdfile, angles)
```

This invokes the *phases* method of the *EBSD* class and returns a dictionary with percentage abundances into a variable called *phases\_available*, and then further prints the available phases and the percentage abundance within the provided ebsd file.

The above directive loads the *ebsd* dataframe into a variable called *df*, and then prints all of the pixel information, with their corresponding phases numbered from 0 to n, (n means the number of phases identified from the ebsd file), their bunge-euler angles, band contrast (BC), band slope (BS), as well as mean angular deviation (MAD), a measure of how accurate the solution assigned to each EBSD pattern is.

Information on the dataframe and the operations that can be performed can be found in the official documentation of the pandas <a href="https://pandas.pydata.org/pandas-docs/stable/reference/api/pandas.DataFrame.html">https://pandas.pydata.org/pandas-docs/stable/reference/api/pandas.DataFrame.html</a>

\*A dataframe is like a python based object for Excel based calculations, the dataframe stores tabular forms of data in which many operations can be done.

To print the header information, which includes the acquisition details of the ebsd file, first load the ebsd data header into a dataframe variable here called *df\_header*.

```
df_header = ebsdfile.get_ebsd_data_header()
print(df header)
```

# Maybe follow Import - ROtation - Cleaning - ETC

To rotate the EBSD data to match the SEM orientation

```
angles = (180, 0, 0)
ebsdfile_downsample = ebsdfile.downsampleEBSD()
updatedebsd = ebsdfile.rotateEBSD(ebsdfile downsample, angles)
```

# 2.0 Cleaning EBSD data

To clean the EBSD dataset we successively remove grains with large mean angular deviation (MAD), reconstruct grains with grain boundaries misorientation ≥ 10 degrees, and remove grains smaller than 7 pixels.

# 2.1 Listing of the phases

Firstly, derive the phase names from the ebsd file using:

```
from santex import EBSD
ebsdfile = EBSD("ebsd.ctf")
phases_names = ebsdfile.phases_names()
print(phases_names)
```

# 2.2 Remove pixel data with MAD higher than a user specified value

To remove pixels with mean angular deviation (MAD) > e.g., 0.8, and store the cleaned dataset into a new dataframe called *filtered\_df*:

```
filtered_df = ebsdfile.filterMAD(df, 0.8)
print(filtered df)
```

## 2.3 Reconstruct grains with boundaries misorientation

Reconstruct grains with misorientation ≥ 10 degrees and store the cleaner dataset into a new dataframe called *df\_grain\_boundary*:

```
df_grain_boundary = ebsdfile.calcGrains(df = filtered_
df, threshold = 10, phase_names=phases_names,
downsampling_factor=100)
```

## 2.4 Remove small grains

Remove tiny grains smaller than e.g. 7 pixels, and store the cleaner dataset into a new dataframe called *filtered df grain boundary*:

```
filtered_df_grain_boundary =
ebsdfile.filterByGrainSize(df_grain_boundary, phases_names,
min grain size=7)
```

# 2.5 Compare original and clean datasets

To compare the original ebsd dataset (*ebsdfile*) and the clean dataset (*filtered\_df\_grain\_boundary*) users can plot them:

```
ebsdfile.plot()
ebsdfile.plot(df = filtered df grain boundary)
```

# 2.6 Plotting Conventions

```
ebsdfile.plot(data = ebsdfile.downsampleEBSD(),
rotation angle=90, inside plane=False)
```

# 2.6 Sample reference alignment

Following directive loads the EBSD data with x to east, y to north and z out of plane

ebsdfile.plot(df)

# **ODF, PDF and IPF analysis**

```
ebsdfile.odf(df, phase=1, crystal symmetry='D2', random val=True,
miller=[1, 0, 0], hemisphere = 'both', axes_labels=["Xs", "Ys"],
alpha = 0.01, figure = None, vector_labels = None,
reproject=False, show hemisphere label = None,
       grid = None, grid resolution = None, return figure = None
)
ebsdfile.pdf(df, phase=1, crystal_symmetry='D2', random_val=True,
miller=[1, 0, 0], hemisphere = 'both',
       axes labels=["Xs", "Ys"], alpha = 0.01, figure = None,
vector labels = None, reproject=False, show hemisphere label =
None,
      grid = None, grid resolution = None, return figure = None
)
ebsdfile.ipf(df, phase=1, vector sample=[0, 0, 1],
random val=True,
           vector title='Z', projection='ipf',
crystal symmetry='D2'
```

# 3.0 Calculation and visualisation of seismic anisotropies

# 3.1 Prepare Dataframes

```
from santex import Material
```

Create a material instance from material class:

```
material_instance = Material()
phase = 'Diopside'
voigtMatrix = material_instance.voigthighPT(phase, PRESSURE =
3, TEMP = 1000)
```

Alternatively, if we want standard reference tensors:

```
voigtMatrix = material instance.get voigt matrix(phase)
```

For example, if we want to get stiffness tensors and densities for Forsterite, diopside and enstatite, we write:

```
cij_Forsterite = material_instance.voigthighPT('Forsterite',
PRESSURE = 3, TEMP = 1000)
cij_Enstatite = material_instance.voigthighPT('Enstatite',
PRESSURE = 3, TEMP = 1000)
cij_Diopside = material_instance.voigthighPT('Diopside', PRESSURE
= 3, TEMP = 1000)

density_Forsterite = material_instance.load_density("Forsterite")
density_Enstatite = material_instance.load_density("Enstatite")
density_Diopside = material_instance.load_density("Diopside")
```

Similarly we can invoke print keyword to look at the cij Forsterite.

Store the densities in the list format for both cij and density. This will assemble all the stiffness tensors and densities into one variable.

```
cij = [cij_Forsterite, cij_Enstatite, cij_Diopside]
density = [density_Forsterite, density_Enstatite,
density_Diopside]
```

## 3.2 Calculation of Anisotropy from EBSD File

Get all the euler angles for each phases and store them in a variable called euler\_angles:

```
forsterite = ebsdfile.get_euler_angles(phase = 1, data =
filtered_df_grain_boundary)
enstatite = ebsdfile.get_euler_angles(phase = 2, data =
filtered_df_grain_boundary)
diopside = ebsdfile.get_euler_angles(phase = 3, data =
filtered_df_grain_boundary)
euler_angles = [forsterite, enstatite, diopside]
```

To calculate the average tensor and density:

```
from santex import Anisotropy
from santex import EBSD

average_tensor, average_density =
ebsdfile.getAnisotropyForEBSD(cij = cij, euler_angles=
euler angles, density=density)
```

The average\_tensor and average\_density is calculated internally, where we can parse this as input to SAnTex's Anisotropy class, and then we can generate the anisotropy plot for the EBSD file

```
anis = Anisotropy(average_tensor*10**9, average_density)
anis.plot()
```

If you want to include melt in the calculation, the modification will be

```
average_tensor, y, average_density =
ebsdfile.getAnisotropyForEBSD(cij = cij, euler_angles=
euler_angles, density=density, melt = 2)
```

# 3.3 Anisotropy Calculations and Visualisation

The SAnTex module *Anisotropy* requires a stiffness tensors (here *stiffness\_matrix*) as an numpy array, and a density in kg.m<sup>-3</sup> and returns an instance of the Anisotropy class from which seismic anisotropy is calculated.

```
[9.735, 6.295, 33.85, 0., 60.23, 0.],
[0., 0., 0., 6.415, 0., 65.18]]) * 10**9

density = 3500

# Create an instance of the Anisotropy class
anisotropy_instance = Anisotropy(stiffness_matrix, density)
anisotropy_instance.plot()
```

# 3.4 Anisotropy Calculations for Modal Rock

```
from santex import ModalAnisotropy
modal_rock = { 'diopside : 0.5', 'forsterite : 0.3',
   'enstatite' : 0.2}
anisotropy_instance = AnisotropyModal(modal_rock, pressure =
2, temperature = 1000)
anisotropy_instance.plot()
```

#### 3.5 3-D Visualisations

To visualise the seismic anisotropies vp, vs1, vs2, and shear wave splitting one calls the anisotropy\_instance.plot():

#### For vp

```
anisotropy instance.plotter vp(density, stiffness matrix)
```

# For vs1:

```
anisotropy instance.plotter vs1(density, stiffness matrix)
```

#### For vs2:

```
anisotropy instance.plotter vs2(density, stiffness matrix)
```

## For vs splitting:

```
anisotropy_instance.plotter_vs_splitting(density,
stiffness_matrix)
```

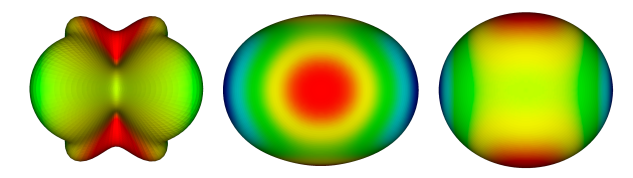


Figure 1: (a) Olivine shear wave splitting (b) Olivine Vp (c) Olivine Vs1