

Imperfections – deformation and microstructures in polycrystals

Sébastien Merkel
Professor, Physics Department
UMET Laboratory (Unité Matériaux et Transformations)
sebastien.merkel@univ-lille.fr

Imperfections - deformation and microstructures in polycrystals

2- Grain orientation

Objectives

How to represent a grain orientation?

- Coordinate systems
- Euler angles
 - Presentation
 - Bunge Euler angle
- Matrix representation
 - Definition
 - Properties
- Graphical representation :
 - Introduction to pole figures
 - Examples and applications.

Imperfections - deformation and microstructures in polycrystals

2- Grain orientation a- Coordinate systems

Representation of orientations

Need to define several coordinate systems (or *reference frames*)

- A reference frame for each crystallite
- A reference frame for the polycrystal, the sample

In texture analysis, we use Cartesian coordinate systems

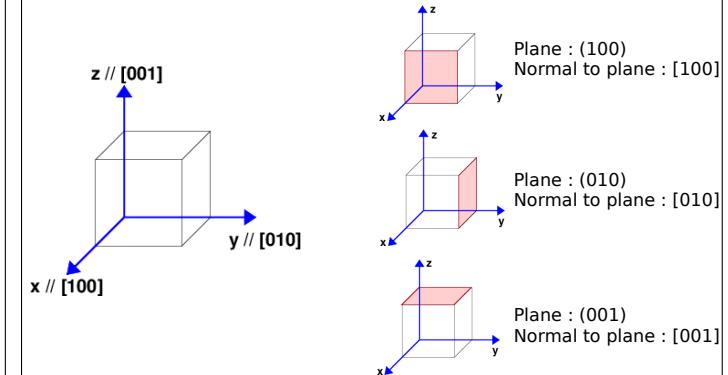
Grain orientation

- Operation *sample reference frame → grain reference frame*

In 3D, the transformation between 2 Cartesian coordinate systems can be seen as

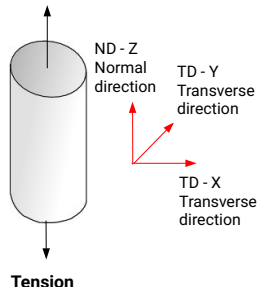
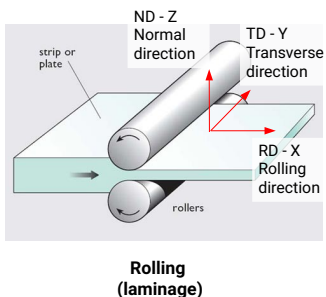
- Rotation matrices (in 3D)
- 3 subsequent rotations, defining Euler angles

Crystal reference frame Cubic symmetry



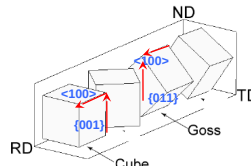
Sample reference frame

Depends on process applied to the sample

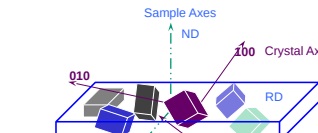


Crystal orientation

Defined with "standard" component or Euler angles to rotate from one coordinate system to the other



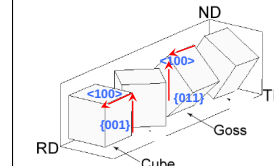
Component	RD	ND
Cube	<100>	{001}
Goss	<100>	{011}
Brass (laiton)	<112>	{110}
Copper (cuivre)	<111>	{112}



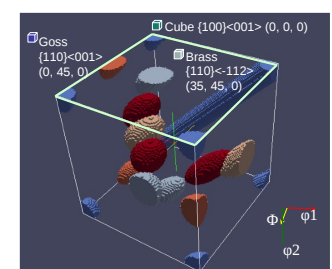
C. N. Tomé and R. A. Lebensohn, Crystal Plasticity, presentation at Pohang University of Science and Technology, Korea, 2009

Orientation space

Defined with "standard" component or Euler angles to rotate from one coordinate system to the other



Component	RD	ND
Cube	<100>	{001}
Goss	<100>	{011}
Brass (laiton)	<112>	{110}
Copper (cuivre)	<111>	{112}

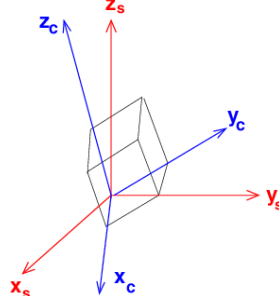


Imperfections - deformation and microstructures in polycrystals

2- Grain orientation b- Euler angles

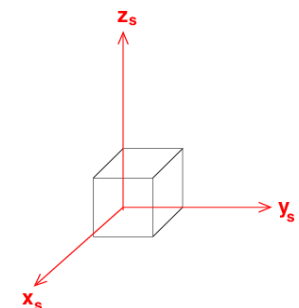
Reference frame transformation Bunge convention

Transformation from the sample reference frame to the crystal reference frame



Reference frame transformation Bunge convention

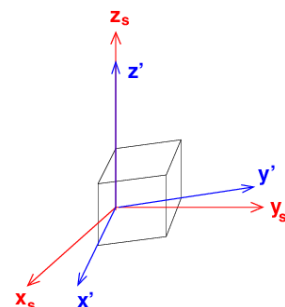
Transformation from the sample reference frame to the crystal reference frame



Reference frame transformation Bunge convention

Transformation from the sample reference frame to the crystal reference frame

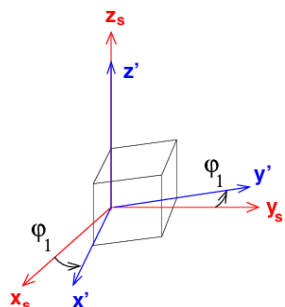
- first rotation



Reference frame transformation Bunge convention

Transformation from the sample reference frame to the crystal reference frame

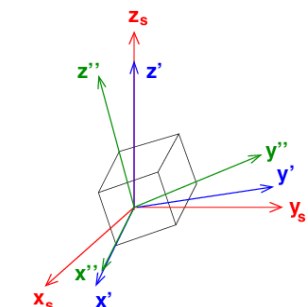
- first rotation
 φ_1 – rotation around z_s



Reference frame transformation Bunge convention

Transformation from the sample reference frame to the crystal reference frame

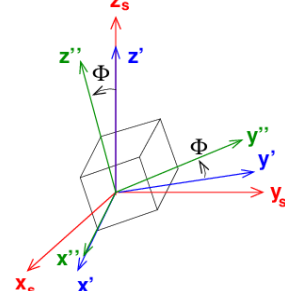
- first rotation
 φ_1 – rotation around z_s
- second rotation



Reference frame transformation Bunge convention

Transformation from the sample reference frame to the crystal reference frame

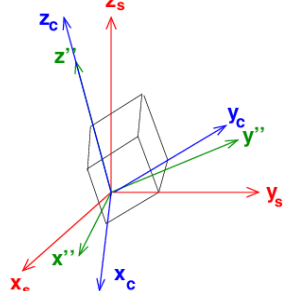
- first rotation
 φ_1 – rotation around z_s
- second rotation
 Φ – rotation around x'



Reference frame transformation Bunge convention

Transformation from the sample reference frame to the crystal reference frame

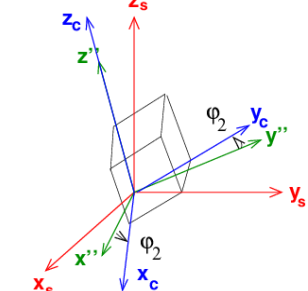
- first rotation
 φ_1 – rotation around z_s
- second rotation
 Φ – rotation around x'
- third rotation



Reference frame transformation Bunge convention

Transformation from the sample reference frame to the crystal reference frame

- first rotation
 φ_1 – rotation around z_s
- second rotation
 Φ – rotation around x'
- third rotation
 φ_2 – rotation around z''



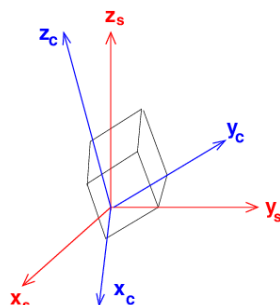
Reference frame transformation Bunge convention

Transformation from the **sample reference frame** to the **crystal reference frame**

- first rotation
 φ_1 – rotation around z_s
- second rotation
 Φ – rotation around x'
- third rotation
 φ_2 – rotation around z''

Crystal orientation defined by 3 Euler angles φ_1 , Φ , et φ_2 .

Bunge convention (there are others, Matthies, Roe, etc)

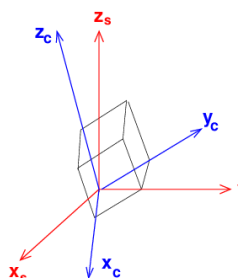


Why 3 angles?

φ_1 – rotation around z_s
 Φ – rotation around x'
 φ_2 – rotation around z''

Angles φ_1 and Φ are used to locate [001] (z_c) in relation to the sample reference frame

Angle φ_2 adds a supplementary rotation to locate x_c and y_c



2- Grain orientation b- Marix representation

Matrix representation

Rotations can be expressed as matrices

$$R_x(\theta) = \begin{pmatrix} 1 & 0 & 0 \\ 0 & \cos\theta & -\sin\theta \\ 0 & \sin\theta & \cos\theta \end{pmatrix}, \quad R_y(\theta) = \begin{pmatrix} \cos\theta & 0 & \sin\theta \\ 0 & 1 & 0 \\ -\sin\theta & 0 & \cos\theta \end{pmatrix}, \quad R_z(\theta) = \begin{pmatrix} \cos\theta & -\sin\theta & 0 \\ \sin\theta & \cos\theta & 0 \\ 0 & 0 & 1 \end{pmatrix}.$$

Direct θ rotation around X

Direct θ rotation around Y

Direct θ rotation around Z

To combine rotations: multiplication of corresponding matrices.

For instance, a rotation of α around z , followed by a rotation β around x , and then a rotation γ around z will be:

$$g(\alpha, \beta, \gamma) = g_z(\gamma) * g_x(\beta) * g_z(\alpha)$$

Pay attention to the order! The first rotation is on the right side of the equation!

Properties of rotation matrices

The inverse of a rotation matrix is its transpose

$$g^{-1} = {}^t g$$

The product of 2 rotations is a rotation

$$g_3 = g_2 * g_1$$

In dimension > 2 , order matters when multiplying rotation matrices

$$g_1 * g_2 \neq g_2 * g_1$$

They are orthogonal (they do not change vector length)

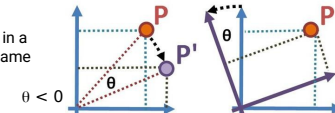
$$\det(g) = 1$$

$$\sqrt{(g_{11}^2 + g_{12}^2 + g_{13}^2)} = 1$$

$$\sqrt{(g_{21}^2 + g_{22}^2 + g_{23}^2)} = 1$$

$$\sqrt{(g_{31}^2 + g_{32}^2 + g_{33}^2)} = 1$$

Active rotation:
The point moves in a fixed reference frame
 $\theta < 0$



Passive rotation:
Le point is fixed and the reference frame moves
 $\theta > 0$

In materials science, we express physical properties (stress, elasticity, etc) in different reference frames

- The sample reference frame
- The crystal reference frame
- Etc

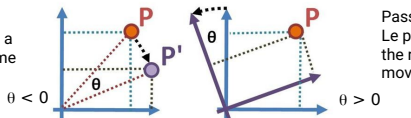
The sample, itself, does not move.

In mechanics, we study movements. The sample moves. Mechanics uses active rotations.

In texture analysis, we are moving reference frames, not the sample : texture analysis uses passive rotations.

Passive rotation - active rotation

Active rotation:
The point moves in a fixed reference frame



Passive rotation:
Le point is fixed and the reference frame moves

$$\begin{cases} x' = x \cos \theta - y \sin \theta \\ y' = x \sin \theta + y \cos \theta \end{cases}$$

$$g = \begin{bmatrix} \cos \theta & -\sin \theta \\ \sin \theta & \cos \theta \end{bmatrix}$$

$$\begin{cases} x' = x \cos \theta + y \sin \theta \\ y' = -x \sin \theta + y \cos \theta \end{cases}$$

$$g = \begin{bmatrix} \cos \theta & \sin \theta \\ -\sin \theta & \cos \theta \end{bmatrix}$$

2- Grain orientation c- Graphical representation – Pole figures

Bunge angles - Rotation matrices

Rotation 1 (φ_1): rotates axis 3 (z) of the sample reference frame

$$g_1 = \begin{bmatrix} \cos \varphi_1 & \sin \varphi_1 & 0 \\ -\sin \varphi_1 & \cos \varphi_1 & 0 \\ 0 & 0 & 1 \end{bmatrix}$$

Rotation 2 (Φ): rotation around axis 1 (x) of the newly formed reference frame

$$g_2 = \begin{bmatrix} 1 & 0 & 0 \\ 0 & \cos \Phi & \sin \Phi \\ 0 & -\sin \Phi & \cos \Phi \end{bmatrix}$$

Rotation 3 (φ_2): rotation around axis 3 (z) of the newly formed reference frame

$$g_3 = \begin{bmatrix} \cos \varphi_2 & \sin \varphi_2 & 0 \\ -\sin \varphi_2 & \cos \varphi_2 & 0 \\ 0 & 0 & 1 \end{bmatrix}$$

$$g(\varphi_1, \Phi, \varphi_2) = g_3 \cdot g_2 \cdot g_1$$

$$= \begin{bmatrix} \cos \varphi_1 \cos \varphi_2 - \sin \varphi_1 \sin \varphi_2 \cos \Phi & \sin \varphi_1 \cos \varphi_2 + \cos \varphi_1 \sin \varphi_2 \cos \Phi & \sin \varphi_2 \sin \Phi \\ -\cos \varphi_1 \sin \varphi_2 - \sin \varphi_1 \cos \varphi_2 \cos \Phi & -\sin \varphi_1 \sin \varphi_2 + \cos \varphi_1 \cos \varphi_2 \cos \Phi & \cos \varphi_2 \sin \Phi \\ \sin \varphi_1 \sin \Phi & -\cos \varphi_1 \sin \Phi & \cos \Phi \end{bmatrix}$$

Geometric interpretation

Crystal x axis in the sample reference frame

$$g = \begin{bmatrix} g_{11} & g_{12} & g_{13} \\ g_{21} & g_{22} & g_{23} \\ g_{31} & g_{32} & g_{33} \end{bmatrix}$$

Crystal y axis in the sample reference frame

$$g = \begin{bmatrix} g_{11} & g_{12} & g_{13} \\ g_{21} & g_{22} & g_{23} \\ g_{31} & g_{32} & g_{33} \end{bmatrix}$$

Sample y axis in the crystal reference frame

$$g = \begin{bmatrix} g_{11} & g_{12} & g_{13} \\ g_{21} & g_{22} & g_{23} \\ g_{31} & g_{32} & g_{33} \end{bmatrix}$$

Sample z axis in the crystal reference frame

Imperfections - deformation and microstructures in polycrystals

Issue with 3D rotation matrices

(Almost) no-one can visualize 3D Euler angles.

Most figures in science use 2D projections. This is what people are used to. How to represent rotations in 3D projected onto a 2D figure?

A true material is made of thousands of crystallites: nearly impossible to look at all Euler angles individually.

Crystals have symmetries. Not easy (at all) to account for crystal symmetries with Euler angles.

Solution: *pole figures*.

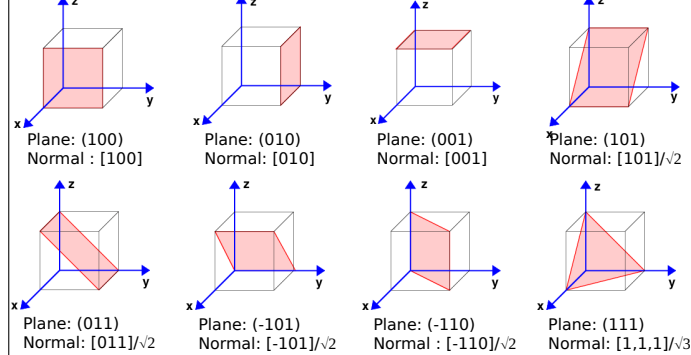
Crystallography reminders

- (hkl) is the plane of *Miller indices* h , k , and l
- $[hkl]$ is the direction of *direction indices* h , k , and l
- $\{hkl\}$ is a family of planes which are *equivalent* due to *symmetry operations*
- $\langle hkl \rangle$ is a family of directions which are *equivalent* due to *symmetry operations*
- hkl indices (with no parenthesis) indicate *Bragg reflection* from the (hkl) planes. They are called *Laue indices*.

Pole = intersection between a line and a sphere around the crystal

- Planes: line perpendicular to a diffracting plane. Defined with Laue indices (i.e. no parenthesis) as we are very often referring to diffraction.
- Directions: line parallel to a crystal direction. Better use $\langle hkl \rangle$ pole in this case.

In a cubic structure



Université de Lille

© S. Merkel
Faculté des Sciences et Technologies
Unité Matériaux et Transformations

44

Tips and tricks

- Pole = intersection between a line and a sphere around the crystal
- Planes: line perpendicular to a diffracting plane. Defined with Laue indices (i.e. no parenthesis) as we are very often referring to diffraction.
 - Directions: line parallel to a crystal direction. Better use $\langle hkl \rangle$ pole in this case.

For a cubic crystal

- $\langle hkl \rangle$ plane is orthogonal to $[hkl]$ direction
- Whatever pole definition you use, it will be the same direction

Pay attention !!!

This is not true for other symmetries than cubic.

The $[hkl]$ direction is not always perpendicular to $\langle hkl \rangle$.

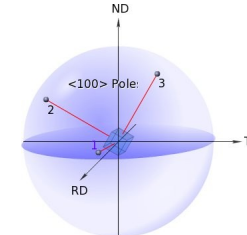
The hkl Laue indices refer to a Bragg reflection, it is perpendicular to $\langle hkl \rangle$.

Université de Lille

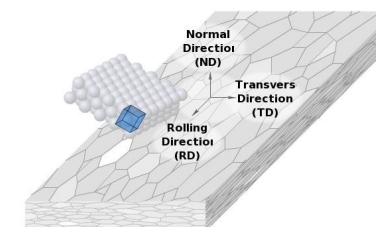
© S. Merkel
Faculté des Sciences et Technologies
Unité Matériaux et Transformations

45

Pole figure



Orientation of poles [100],
[010], [001] : <100> poles



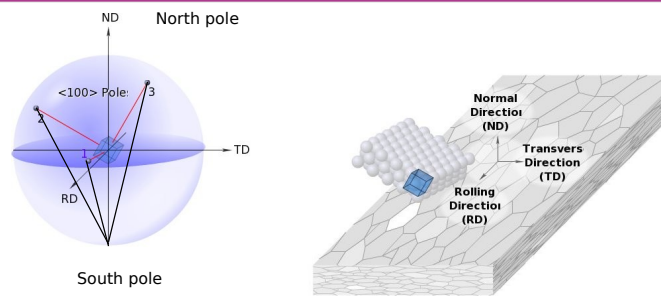
Cubic-structured crystallite in a deformed sample

Illustrations : aluMatter
Université de Lille

© S. Merkel
Faculté des Sciences et Technologies
Unité Matériaux et Transformations

46

Pole figure



Lines between <100> poles and south pole
Intersection with equatorial plane

Cubic-structured crystallite in a deformed sample

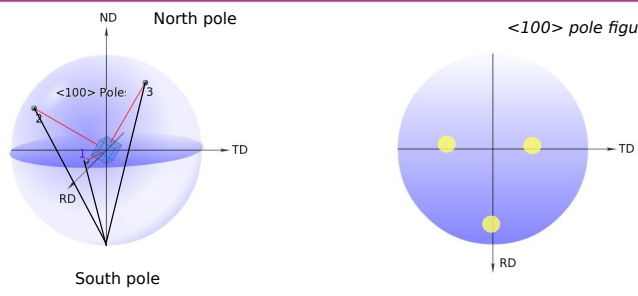
Illustrations : aluMatter

Université de Lille

© S. Merkel
Faculté des Sciences et Technologies
Unité Matériaux et Transformations

47

Pole figure



Lines between <100> poles and south pole
Intersection with equatorial plane

View from North pole

Illustrations : aluMatter

Université de Lille

© S. Merkel
Faculté des Sciences et Technologies
Unité Matériaux et Transformations

48

PF training 1

- Take a cubic crystal with
- $[100] // RD$
 - $[010] // TD$
 - $[001] // ND$

Plot the <100> pole figure
Plot the <111> pole figure

Illustrations : aluMatter

Université de Lille

© S. Merkel
Faculté des Sciences et Technologies
Unité Matériaux et Transformations

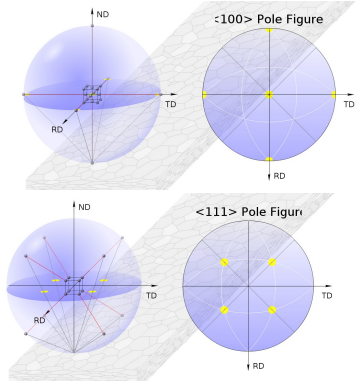
49

PF training 1

Take a cubic crystal with

- [100] // RD
- [010] // TD
- [001] // ND

Plot the <100> pole figure
Plot the <111> pole figure



Illustrations : aluMatter



© S. Merkel
Faculté des Sciences et Technologies
Unité Matériaux et Transformations

50

PF training 2

Take a cubic crystal with

- [100] // RD
- [010] // TD
- [001] // ND

Rotate it by 45° around ND
Plot the <100> pole figure
Plot the <111> pole figure

Illustrations : aluMatter



© S. Merkel
Faculté des Sciences et Technologies
Unité Matériaux et Transformations

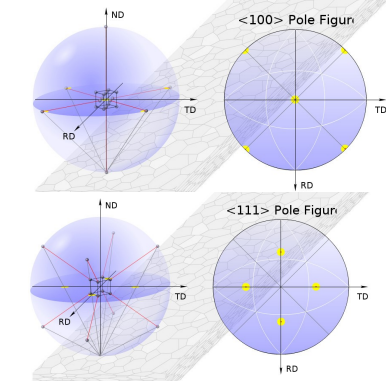
51

PF training 2

Take a cubic crystal with

- [100] // RD
- [010] // TD
- [001] // ND

Rotate it by 45° around ND
Plot the <100> pole figure
Plot the <111> pole figure



Illustrations : aluMatter



© S. Merkel
Faculté des Sciences et Technologies
Unité Matériaux et Transformations

52

PF training 3

Take a cubic crystal with

- [100] // RD
- [010] // TD
- [001] // ND

Rotate it by 45° around TD
Plot the <100> pole figure
Plot the <111> pole figure

Illustrations : aluMatter



© S. Merkel
Faculté des Sciences et Technologies
Unité Matériaux et Transformations

53

PF training 3

Take a cubic crystal with

- [100] // RD
- [010] // TD
- [001] // ND

Rotate it by 45° around TD
Plot the <100> pole figure
Plot the <111> pole figure

Illustrations : aluMatter



© S. Merkel
Faculté des Sciences et Technologies
Unité Matériaux et Transformations

54

