27-750, Advanced Characterization and Microstructural Analysis: Texture and its Effect on Anisotropic Properties


Last revised: 12th Jan. 2014
Microstructure-Properties Relationships

Microstructure

Processing

Design

Performance

Properties
Course Objective

• Many courses deal with microstructure-properties relationships, so what is special about this course?!

• Despite the crystalline nature of most useful and interesting materials, crystal alignment and the associated anisotropy is ignored. Yet, most properties are sensitive to anisotropy. Therefore *microstructure* should include *crystallographic orientation* (“texture”).

• The objective of this course is to provide you with the tools to understand and quantify various kinds of texture and to solve problems that involve texture and anisotropy.
Questions

• Examples of questions that you should be able to answer with the knowledge and skills provided by this course:
• What is a “fiber texture”?
• Why is a <111>//ND texture ideal for deep drawing?
• Why is obtaining a <111> fiber texture difficult in FCC metals, but straightforward in BCC?
• Why are intensity values generally much higher in the Orientation Distribution than in the corresponding pole figures?
• How is it possible to recover the full 5-parameter distribution of grain boundary character from a plane section and yet one can only measure 4 out of 5 parameters for an individual boundary in that plane section?
• What do the units “Multiples of a Random/Uniform Distribution” mean? Why are distributions scaled differently in texture than in statistics?
• Why was solving the problem of calculating an orientation distribution from pole figures a fundamental advance in texture analysis? Hint: think about the parameterization of rotations.
• Why do we need 3 (and only 3) parameters to describe a rotation?
• How do Miller indices, orthogonal matrices, Rodrigues parameters and quaternions relate to each other?
• What is epitaxy? What is apotaxy (not apoplexy!)?
• Why do textures develop during plastic deformation?
Texture refers to the physical makeup of rock—namely, the size, shape, and arrangement (packing and orientation) of the discrete grains or particles of a sedimentary rock. Two main natural textural groupings exist for sedimentary rocks: clastic (or fragmental) and nonclastic (essentially crystalline). Noncarbonate chemical sedimentary...
Websters’ Dictionary, fabric

- Main Entry: fabric
- Pronunciation: ‘fa-brik
- Function: noun
- Etymology: Middle French fabrique, from Latin fabrica workshop, structure
- Date: 15th century
- 1 a : STRUCTURE, BUILDING b : underlying structure : FRAMEWORK <the fabric of society>
- 2 : an act of constructing : ERECTION; specifically : the construction and maintenance of a church building
- 3 a : structural plan or style of construction b : TEXTURE, QUALITY -- used chiefly of textiles c : the arrangement of physical components (as of soil) in relation to each other
- 4 a : CLOTH 1a b : a material that resembles cloth
- 5 : the appearance or pattern produced by the shapes and arrangement of the crystal grains in a rock
Websters’ Dictionary,
anisotropy

- Main Entry: an·iso·trop·ic
- Pronunciation: "a-"nl-s&-'trä-pik
- Function: adjective
- Date: 1879: exhibiting properties with different values when measured in different directions <an anisotropic crystal>
- - an·iso·trop·i·cal·ly /-pi-k(&-)lE/ adverb
- - an·isot·ro·py /-(")nl-'sä-tr&-pE/ also an·isot·ro·pism /-"pi-z&m/ noun
People

- The development of the field is greatly indebted to Hans J. Bunge who passed away in 2006.
- His textbook (translated from the German original), *Texture Analysis in Materials Science*, is a very useful reference and many of his suggestions are only just now being developed into useful tools.
Microstructure

- Conventional Approach: grain structure, phase structure (*qualitative*, image based), emphasizes interfaces and boundaries between phases.
- Quantitative (conventional): grain size, aspect ratio(s), particle size, phase connectivity.
- *Modern* Quantitative: (probability) distributions of orientation of crystal axes (relative to a reference frame) of crystals or boundaries between crystals. Properties calculated from distributions and/or microstructures with orientation included.
Microstructure with Crystal Directions

Note cleavage planes within each grain: a natural indicator of crystallographic directions in a geological material.
Why study texture?

- Many examples exist of materials engineered to have a specific texture in order to optimize performance (single crystal turbine blades, transformer steel, magnetic thin films...).
- Control of texture achievable through control of processing but many challenges remain.
Texture examples

- Example 1. Transformer Steel
- Example 2. Anisotropic particles (whiskers) of hydroxyapatite (HA) in polyethylene (PE)
- Example 3. Earing in Deep Drawing of Cups: see slides on forming of Beer Cans
- Example 4. Anisotropy of Fatigue Properties in Aerospace Al
- Example 5. Effect of Grain Boundary Character on Pb Electrodes in Lead-Acid Batteries: see slides on grain boundaries and grain boundary engineering (GBE)
Example 1: Transformer Steel

- 1935: Goss first published his work on high permeability silicon steels.
- The most commonly used material as the soft magnetic material for transformer laminations is a highly oriented albeit polycrystalline 3%Si steel; in other words, the material is almost a *single crystal*.
- "Goss orientation" has a <110> direction normal to the sheet and a <001> parallel to the rolling direction.
- Aligns the softest magnetic direction with the direction of magnetization. Thus transformers made from the textured sheet exhibit lower electrical losses.
- Processing relies on a secondary recrystallization step in which all grains in a fine, primary recrystallized structure are pinned by second phase particles while the Goss grains grow to consume the entire volume.
- Not clearly understood what differences in grain boundary character at the perimeter of the growing Goss grains provides them with the ability to grow at the expense of the general population.
Example 2: HA particles in PE

- The figure shows (a) Spherical hydroxyapatite particles (b) Whisker hydroxyapatite particles (c) Size and frequency of the hydroxyapatite particles.
Eg 2, contd.: HA particles in PE

- The figure shows (a) An XRD orientation comparison of whisker hydroxyapatite particles and random powder (b) An XRD orientation comparison of spherical hydroxyapatite particles and random powder [Zhang et al.]. Texture is inferred from the difference between the measured powder pattern and the pattern expected for a randomly oriented material (from the powder diffraction file). This is typical in the literature as a purely qualitative measure of texture.
Texture in HA in bone: refs

- X-ray Pole Figure Analysis of Apatite Crystals and Collagen Molecules in Bone - all 3 versions, N Sasaki - Calcified Tissue International, 1997 - Springer
- ... figure analysis of mineral nanoparticle orientation in individual trabecula of human vertebral bone - all 6 versions, D Jaschouz, O Paris, P Roschger, HS Hwang, P ... - Journal of Applied Crystallography, 2003 - dx.doi.org
- Crystal alignment of carbonated apatite in bone and calcified tendon: results from quantitative ... - all 4 versions, HR Wenk, F Heidelbach - Bone, 1999 - Elsevier
- Pole figures of the orientation of apatite in bones - all 3 versions, JP Nightingale, D Lewis - Nature, 1971 - nature.com, Pole Figures of the Orientation of Apatite in Bones. ... THE orientation of the apatite and collagen in bone was first considered in this work because of its ...
- Orientation of apatite in single osteon samples as studied by pole figures, A Ascenzi, E Bonucci, P Generali, A Ripamonti, N ... - Calcified Tissue International, 1979 - Springer
- Bone Marrow Is a Reservoir of Repopulating Mesangial Cells during Glomerular Remodeling - all 4 versions, T Ito, A Suzuki, E Imai, M Okabe, M Hori - Journal of the American Society of Nephrology, 2001 - jasn.org
- Quantitative texture analysis of small domains with synchrotron radiation X-rays, F Heidelbach, C Riekel, HR Wenk - logo, 1999 - dx.doi.org
Connections

• Crystals are anisotropic.
• A collection of crystals (a polycrystal) is therefore anisotropic unless all possible orientations are present.
• Almost any processing of a material changes and biases the crystal orientations, leading to texture development.
• Anisotropy can be taken advantage of; therefore it makes sense to engineer (control, design) the texture of a material.
**Books, Links**

- **Course Textbook:** U.F. Kocks, C. Tomé, and H.-R. Wenk, Eds. (1998). *Texture and Anisotropy*, Cambridge University Press, Cambridge, UK, ISBN 0-521-79420-X. This is now available as a paperback. Relevant chapters: 1, 2, 3, 4, 5, 6, 7, 8. Note that there is more detail in each chapter than we will have time to cover.


Secondary References

• Ohser, J. and F. Mücklich (2000), *Statistical Analysis of Microstructures in Materials Science*
• Reid, C. N. (1973). *Deformation Geometry for Materials Scientists*
• Gottstein, G. and L. S. Shvindlerman (1999). *Grain Boundary Migration in Metals*
• Howe, J.M. (2000). *Interfaces in Materials*
• http://www/msm.cam.ac.uk/phase-trans/texture.html
• http://labotex.com/
Topics, Activities in Course: 1

• First major topic will be a discussion of orientations and how to represent them quantitatively with Miller indices, matrices, Rodrigues vectors and quaternions.

• The next major topic will be x-ray pole figures and their analysis
  – Every student will obtain his/her own data set
  – We will first perform a standard analysis using popLA to generate an orientation distribution; then each student will measure their own pole figures and analyze the results
  – The emphasis will be on development of practical skills followed up by discussion of the underlying concepts
  – The objective will be to have students be competent and comfortable with pole figure analysis

• Each student will report on their analyses as their project presentation at the end of the course.
Topics, Activities: 2

• The next major topic will be the analysis of orientation distributions
  – This will involve understanding the relationships between the different methods of describing orientations, especially Euler angles and Miller indices
  – We will explore the mathematical aspects of orientation space and the impact of crystal symmetry and sample symmetry
  – The objective will be to develop students’ quantitative skills with orientation information so that they understand the physical meaning of orientation and texture
Topics, Activities: 3

- The next major topic will be to investigate orientation imaging microscopy (OIM) based on automated indexing of electron back scatter diffraction (EBSD) patterns in the scanning electron microscope (SEM)
  - As with x-ray pole figures, students will first analyze a standard data set and then will make their own scan (if they are not already using EBSD) for further analysis
  - The objective will be to understand the differences between sampling discrete orientations in a limited area (EBSD) and measurement of the average orientation (distribution) over a large area
Topics, Activities: 4

• The next major topic is grain boundaries, whose crystallography can be easily characterized by electron microscopy
  – We will discuss the physical characteristics of grain boundaries, e.g. energy, mobility, together with the additional complications for symmetry and descriptions (Rodrigues vectors, quaternions)
  – The objective is for students to become familiar with both the properties of grain boundaries and the methods for quantitative characterization
Topics, Activities: 5

• The next major topic is microstructure-property relationships using texture information
  – Students will explore percolation analysis using electrical conductivity in superconductors as an example of a case where the crystal properties are (strongly) anisotropic and the grain boundaries are also anisotropic.
  – This exercise will teach students how to develop a computer model on a discrete grid. Programming will be required, although any of the following languages may be used: C, C++, Fortran, VisualBasic.
The next major topic is microstructural measurement and stereology. Stereology is the science of obtaining 3D information about microstructure from 2D sections.

- Even something as seemingly straightforward as grain size is interesting because we lack quantitative models for the size distribution. GSDs are approximately log-normal, but not exactly.
- Stereology is necessary because characterization is most readily available on plane cross sections. Therefore for most microstructures, we need tools to infer the true 3D image from the 2D slices through the material
- The objective is to equip students to understand and use stereological tools, e.g. reconstruction of particle size distributions from cross sections, or, use of Microstructure Builder
The next major topic is elastic and plastic anisotropy

- Plastic deformation in metals (and ceramics at high temperatures, and some polymers) is governed by the motion of line defects - dislocations. The crystallographically restricted slip directions (Burgers vector) and slip planes mean that any degree of texture results in an anisotropic response, e.g. a multi-axial strain from an imposed uniaxial stress

- The objective is to equip students to understand and use polycrystal analysis + modeling, e.g. LApp
Lecture List (abbreviated)

1. Introduction
2. Texture components, Euler angles
3. X-ray diffraction
4. Calculation of ODs from pole figure data, popLA
5. Orientation distributions
6. Microscopy, SEM, electron diffraction
7. Texture in bulk materials
8. EBSD/OIM
9. Misorientation at boundaries
10. Continuous functions for ODs
11. Stereology
12. Graphical representation of ODs
13. Symmetry (crystal, sample)
14. Euler angles, variants
15. Volume fractions, Fiber textures
16. Grain boundaries
17. Rodrigues vectors, quaternions
18. CSL boundaries
19. GB properties
20. 5-parameter descriptions of GBs
21. Herring’s relations
22. Elastic, plastic anisotropy
23. Taylor/Bishop-Hill model
24. Yield Surfaces
Learning Approach

1. Overall Concept
2. Phenomenology
3. Cause-and-Effect
4. Required Math+Physics+Chemistry
5. Measurement Technique, data
6. Analysis
7. Interpretation
Anisotropy-Texture

1. **Overall Concept:**
   materials behave anisotropically and, regarding texture as part of microstructure, this is another microstructure-property relationship

2. **Phenomenology:**
   anisotropy is correlated with non-random grain alignment.

3. **Cause-and-Effect:**
   the cause of anisotropic behavior is the crystallographic preferred orientation (texture) of the grains in a polycrystal.

4. **Required Math:**
   Crystal orientation is described by a (3D) rotation; therefore texture requires distributions of rotations to be described.

5. **Measurement Technique, data:**
   see next page

6. **Analysis:**
   3D distributions have to be reconstructed from 2D projections

7. **Interpretation:**
   Although pole figures often provide easily recognized patterns, orientation distributions provide quantitative information.
Crystal Orientations – Euler angles

Component | RD  | ND   |
------------|-----|------|
Cube        | <100>| {001}|
Goss        | <100>| {011}|
Brass       | <112>| {110}|
Copper      | <111>| {112}|

Rotation 1 ($\varphi_1$): rotate sample axes about ND

Rotation 2 ($\Phi$): rotate sample axes about rotated RD

Rotation 3 ($\varphi_2$): rotate sample axes about rotated ND

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

<table>
<thead>
<tr>
<th>Component</th>
<th>Euler Angles (°)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cube</td>
<td>(0, 0, 0)</td>
</tr>
<tr>
<td>Goss</td>
<td>(0, 45, 0)</td>
</tr>
<tr>
<td>Brass</td>
<td>(35, 45, 0)</td>
</tr>
<tr>
<td>Copper</td>
<td>(90, 45, 45)</td>
</tr>
</tbody>
</table>

[1] C. N. Tome and R. A. Lebensohn, crystal plasticity, presentation at Pohang University of Science and Technology, Korea, 2009
Crystal Orientations – ODF

ODF gives the density of grains having a particular orientation.

ODF is the Orientation Distribution Function, denoted as $f(g)$, which gives the density of grains having a particular orientation. In the context of crystallography, the orientation of a crystal can be represented by a set of angles $g = \{\phi_1, \Phi, \phi_2\}$.

- **Goss**: {110}<001> (0, 45, 0)
- **Cube**: (100)<001> (0, 0, 0)
- **Brass**: {110}<-112> (35, 45, 0)

Contours at $0.500, 1.000, 2.000, 4.000, 8.000, 16.000$
A {111} pole figure of rolled copper, showing the typical distribution of intensity for moderate to large strains. The rolling plane normal (ND) is perpendicular to the plane of the figure and the rolling (RD) and transverse (TD) directions are vertical and horizontal, respectively, in the plane of the figure. The contours indicate the diffracted intensity in units of Multiples of a Random Density (MRD). High frequencies of <111> directions are found close to the RD, for example, and also inclined 20° away from the ND towards the RD [Hirsch, J. and K. Lücke. Mechanism of Deformation and Development of Rolling Textures in Polycrystalline FCC Metals 1. Description of Rolling Texture Development in Homogeneous CuZn Alloys. *Acta Metallurgica*, 36 (11): 2863-2882, 1988].
**Effect of Alloying: Cu-Zn (brass);**

*the texture transition*

Check contour levels: 1, 2, 3 ...?

Zn content: (a) 0%, (b) 2.5%, (c) 5%, (d) 10%, (e) 20% and (f) 30% [Stephens 1968]
Texture: Quantitative Description

- **Three (3) parameters** needed to describe the orientation [of a crystal relative to the embedding body or its environment] because it is a 3D rotation.
- Most common description: 3 [rotation] **Euler angles**
- Other descriptions include: (orthogonal) rotation matrix (or axis transformation matrix), Rodrigues-Frank vector, unit quaternion.
- A common misunderstanding: although 2 parameters are sufficient to describe the position of a vector, a 3D object such as a crystal requires **3 parameters** to describe its (angular) position
- Most experimental methods [X-ray pole figures included] do not measure all 3 angles, so **orientation distribution** must be calculated. An orientation distribution is just a probability distribution: it tells you how likely you are to find a crystal that has the orientation specified by the coordinates (Euler angles) of the point
Euler Angles, Animated

$e'_3 = e_3 = z_{sample} = ND$

$z_{crystal} = e''_3$

$e''_3$

$\phi_1$

$\phi_2$

$e'_2$

$e''_2$

$x_{crystal} = e'_1 = e''_1$

$e_1 = X_{sample} = RD$

$e_2 = Y_{sample} = TD$

$e_3 = Z_{sample} = ND$

$e''_3$

$y_{crystal} = e''_2$

$e'_3$

$[010]$

$[001]$

$[100]$

3rd position (final)

2nd position

1st position

Sample Axes

Crystal

RD

TD
Sections through an OD

\[ \phi_2 = 0^\circ \quad \phi_2 = 5^\circ \quad \phi_2 = 15^\circ \]

This example of the texture of rolled copper, taken from Bunge’s book, uses the Bunge definition of the Euler angles so that each possible orientation is defined by \((\phi_1, \Phi, \phi_2)\)
Transformations of Axes

• A 3D object such as a crystal requires 3 parameters to describe its (angular) position.

• The 3 parameters can be one of a great variety of types, including Euler angles (best known in Materials Science), unit quaternions (popular in robotics), Rodrigues vectors (useful for grain boundaries), or rotation matrices.

• Most often in materials science, we prefer to describe properties and field quantities (stress, strain, current, heat flux ...) in terms of a convenient local frame.

• Since some quantities are most easily described in, say, a frame associated with a specimen (e.g. rolling-transverse-normal) and others (such as single crystal properties) are associated with the local crystal frame, we need a method to transform quantities from one frame to another.

• There is a standard procedure known as “transformation of axes” that we will use. Later on we will see that it is implicit in the Tensor Transformation Rule. Mathematicians know this as a “passive rotation”. It is complementary to the “active rotation” commonly used in solid mechanics.

 Scalars, Vectors, Tensors: Notation

• General case: three dimensions
• Vector: needs 3 numbers or coefficients to quantify its $x$, $y$ and $z$ components.
• Two notations for vectors: “vector-tensor notation” where bold-face implies higher-than-scalar nature; “component notation” where a suffix(-es) show how many coefficients are needed.
• Vector: either $\mathbf{b}$ or $b_i, i \in \{1,2,3\}$, or, $i \in \{x,y,z\}$.
• 2nd rank tensor: either $T$ or $T_{ij}, i,j \in \{1,2,3\}$
• Advantage of vector-tensor notation is that the equations work (or, are valid) in any reference frame. By contrast, when component notation is used, the actual values of the coefficients depend on which reference frame is used.
• If you see subscripts attached to a quantity, it is (almost always) a tensor and the Einstein summation convention is assumed. The Einstein summation convention says that a repeated index (on the RHS) implies summation over that index (typically 1,2, and 3 in 3D). If the same, non-repeated index is found on both sides of an equation, then no summation is performed.
Changing the Coordinate System

• Many different choices are possible for the orthonormal base vectors and origin of the Cartesian coordinate system. A vector is an example of an entity which is independent of the choice of coordinate system. Its direction and magnitude must not change (and are, in fact, invariants), although its components will change with this choice.

• Why would we want to do something like this? For example, although the properties are conveniently expressed in a crystal reference frame, experiments often place the crystals in a non-symmetric position with respect to an experimental frame. Therefore we need some way of converting the coefficients of the property into the experimental frame.

• Changing the coordinate system is also known as axis transformation.
Motivation for Axis Transformation

- One motivation for axis transformations is the need to solve problems where the specimen shape (and the stimulus direction) does not align with the crystal axes. Consider what happens when you apply a force parallel to the sides of this specimen ...

The direction parallel to the long edge does not line up with any simple, low index crystal direction. Therefore we have to find a way to transform the properties that we know for the material into the frame of the problem (or vice versa).

*Image of Pt surface from www.cup.uni-muenchen.de/pwintterlin/IMGs/pt10p3.jpg*
New Axes

• Consider a new orthonormal system consisting of right-handed base vectors: \( \hat{e}'_1, \hat{e}'_2 \text{ and } \hat{e}'_3 \)
  These all have the same origin, \( o \), associated with the original axes: \( \hat{e}_1, \hat{e}_2 \text{ and } \hat{e}_3 \)

• The vector \( \mathbf{v} \) is clearly expressed equally well in either coordinate system:

\[
\mathbf{v} = v_i \hat{e}_i = v'_i \hat{e}'_i
\]

Note - same physical vector but different values of the components.

• We need to find a relationship between the two sets of components for the vector.
Direction cosines

Each direction cosine is the length of the unit vector, $\mathbf{a}$, projected onto each axis in turn. The second direction cosine, $\alpha_2$ is shown.

$\alpha_i = \hat{\mathbf{a}} \cdot \hat{x}_i$

$\alpha_1 = u = \cos \theta_1$
$\alpha_2 = v = \cos \theta_2$
$\alpha_3 = w = \cos \theta_3$
Rotation of axes in the x-y plane

Passive Rotation/Transformation of Axes

\[ v' = a v = \begin{pmatrix} \cos \theta & \sin \theta \\ -\sin \theta & \cos \theta \end{pmatrix} v \]

General rule for determining the coefficients of a transformation matrix, using dot-products:

\[ a_{ij} = \hat{x}_i' \cdot \hat{x}_j \]

\( x, y = \text{old axes}; \ x', y' = \text{new axes} \)

x, y = old axes; x’, y’ = new axes

Passive Rotation/Transformation of Axes
Example: rotation angle = 30°

\[
v' = a v = \begin{pmatrix} \cos 30° & \sin 30° \\ -\sin 30° & \cos 30° \end{pmatrix} v = \begin{pmatrix} \sqrt{3}/2 & 1/2 \\ -1/2 & \sqrt{3}/2 \end{pmatrix} v
\]

\[a_{ij} = \hat{\mathbf{x}}'_i \cdot \hat{\mathbf{x}}_j\]

\[x, y = \text{old axes}; \quad x', y' = \text{new axes}\]

Passive Rotation/Transformation of Axes
Tensor: definition, contd.

- In order for a quantity to “qualify” as a tensor it has to obey the axis transformation rule, as discussed in the previous slides.

- The transformation rule defines relationships between transformed and untransformed tensors of various ranks.

Vector:

- 2nd rank
  
  \[ V'_i = a_{ij} V_j \]
  
  \[ T'_{ij} = a_{ik} a_{jl} T_{kl} \]

- 3rd rank
  
  \[ T'_{ijk} = a_{il} a_{jm} a_{kn} T_{lmn} \]

- 4th rank
  
  \[ T'_{ijkl} = a_{im} a_{jn} a_{ko} a_{lp} T_{mnop} \]

- This rule is a critical piece of information, which you must know how to use.
3D Axis Transformation

Sample to Crystal (primed)

\[ a_{ij} = \hat{e}_i \cdot \hat{e}_j \]

\[
\begin{pmatrix}
  a_{11} & a_{12} & a_{13} \\
  a_{21} & a_{22} & a_{23} \\
  a_{31} & a_{32} & a_{33}
\end{pmatrix}
\]

\( e = \) old (sample) axes;
\( e' = \) new (crystal) axes
**Rodrigues-Frank vector definition**

- We write the axis-angle representation as: \((\hat{r}, \alpha)\)
  where the rotation axis = \(\overrightarrow{OQ}/|\overrightarrow{OQ}|\)
- The Rodrigues vector is defined as:

\[
\rho = \hat{r} \tan(\alpha/2) = \tan\left(\frac{\alpha}{2}\right)[r_x, r_y, r_z]
\]

The vector is parallel to the rotation axis, and the rotation angle is \(\alpha\), and the magnitude of the vector is scaled by the *tangent* of the *semi*-angle.
Quaternion: definition

- \( q = q(q_1,q_2,q_3,q_4) \)
- For a unit quaternion, representing a rotation, we have:
  \[
  q(r \sin \theta/2, \cos \theta/2) \\
  q(u \sin \theta/2, v \sin \theta/2, w \sin \theta/2, \cos \theta/2)
  \]
- Here, the rotation axis is \( r=[u,v,w] \), as a unit vector, and the rotation angle is \( \theta \).
- Alternative notation puts cosine term in 1st position, \( q(q_0,q_1,q_2,q_3) : \)
  \[
  q = ( \cos \theta/2, u \sin \theta/2, v \sin \theta/2, w \sin \theta/2 ).
  \]
Summary

• Microstructure contains far more than qualitative descriptions (images) of cross-sections of materials.

• Most properties are anisotropic which means that it is critically important for quantitative characterization to include orientation information (texture).

• Many properties can be modeled with simple relationships, although numerical implementations are (almost) always necessary.
Supplemental Slides
Websters’ Dictionary, texture

- Pronunciation: 'teks-ch&r
- Function: noun
- Etymology: Latin textura, from textus, past participle of texere to weave -- more at TECHNICAL
- Date: 1578
- 1 a : something composed of closely interwoven elements; specifically : a woven cloth b : the structure formed by the threads of a fabric
- 2 a : essential part : SUBSTANCE b : identifying quality : CHARACTER
- 3 a : the disposition or manner of union of the particles of a body or substance b : the visual or tactile surface characteristics and appearance of something <the texture of
- an oil painting>
- 4 a : a composite of the elements of prose or poetry <all these words... meet violently to form a texture impressive and exciting -- John Berryman> b : a pattern of
- musical sound created by tones or lines played or sung together
- 5 a : basic scheme or structure b : overall structure
What do we need to learn?

1. How to measure texture:
   - Method 1: x-ray pole figures
   - Method 2: electron back scatter diffraction (EBSD)
   - Method 3: transmission electron microscopy (TEM)
   - Stereology: sections through 3D materials

2. What causes texture to develop in materials, and how does it depend on material type and the processing history?
   - Deformation of bulk metals: rolling vs. torsion etc.
   - Annealing: grain growth, recrystallization
   - Thin films
What do we need to learn? (contd.)

3. How to describe texture quantitatively, how to plot textures, and how to understand texture:
   Method 1: pole figures
   Method 2: orientation distributions (OD)
   Symmetry: crystal symmetry, sample symmetry
   Components
   Fibers
   How to obtain ODs from pole figures

4. How does anisotropy depend on texture?
   Elastic anisotropy
   Plastic anisotropy; yield surfaces
   Corrosion (grain boundaries)
What do we need to learn? (contd.)

5. Grain Boundaries
   - Grain boundary atomic structure: low angle vs. high angle boundaries
   - Special grain boundaries: Coincident Site Lattice boundaries (CSL)
   - How to describe grain boundary crystallography: axis-angle, Rodrigues vectors
   - How to measure grain boundaries

6. Underlying Concepts
   - Different descriptions of rotations: *Miller indices, Euler angles, matrices, axis-angle pairs, Rodrigues vectors, quaternions*
   - How to work with distributions
   - Spherical harmonics (series expansions)
   - Discretization of distributions
   - Volume fractions
What is the result that we want? For a solved problem, we quote the equation or concept.

How do we find solutions for the differential equations, and what are they?

What are the variables?

How do we visualize the solution - what graphs are appropriate?

How do we set up the differential equations?

What do worked solutions corresponding to physical situations look like?

How do we determine the boundary conditions?
How to Measure Texture

- X-ray diffraction; pole figures; measures \textit{average} texture at a surface (\(\mu\)ms penetration); projection (2 angles).
- Neutron diffraction; type of data depends on neutron source; measures \textit{average} texture in bulk (cms penetration in most materials); projection (2 angles).
- Electron [back scatter] diffraction; easiest [to automate] in scanning electron microscopy (SEM); \textit{local} texture; complete orientation (3 angles).
- Optical microscopy: optical activity (plane of polarization); limited information (one angle)
**X-ray Pole Figures**

- X-ray pole figures are the most common source of texture information; cheapest, easiest to perform.
- Pole figure: variation in diffracted intensity with respect to direction in the specimen.
- Representation: map in projection of diffracted intensity.
- Each PF is equivalent to a geographic map of a hemisphere (North pole in the center).
- *Map of crystal directions w.r.t. sample reference frame.*
Anisotropy Example 2:
Drawn Aluminum Cup with Ears

Figure shows example of a cup that has been deep drawn. The plastic anisotropy of the aluminum sheet resulted in non-uniform deformation and “ears.”

Randle, Engler, p.340
Challenges in Microstructure

- **Annealing textures:** where does the cube texture come from in annealed fcc metals? Goss texture in bcc metals?
- **Processing:** how can we produce large crystals of ceramics by abnormal grain growth?
- **Plastic deformation:** how can we explain the “break-up” of grains during deformation?
- **Simulation, numerical representation:** how can we generate faithful 3D representations of microstructure?
- **Constitutive relations:** what are the properties of defects such as grain boundaries?
**Notation: vectors, matrices**

- **Vector-Matrix**: $\mathbf{v}$ is a vector, $\mathbf{A}$ is a matrix (always a square matrix in this course).
- **Index notation**: explicit indexes (Einstein convention): $v_i$ is a vector, $A_{jk}$ is a matrix (maybe tensor, though not necessarily).
- **Scalar (dot) product**: $c = \mathbf{a} \cdot \mathbf{b} = a_i b_i$; zero dot product means vectors are perpendicular. For two unit vectors, the dot product is equal to the cosine of the angle between them.
- **Vector (cross) product**: $c = c_i = \mathbf{a} \times \mathbf{b} = \mathbf{a} \wedge \mathbf{b} = \varepsilon_{ijk} a_j b_k$; generates a vector that is perpendicular to the first two. Two vectors that are perpendicular have a zero length cross product. The cross product defines a rotation axis that carries one vector into another. The magnitude of the cross product is the product of the magnitudes (lengths) of the vectors multiplied by the sine of the angle between them.
- **Permutation or alternating tensor**, $\varepsilon_{ijk}$, is $+1$ for $ijk=123, 231, 312$, and $-1$ for $ijk=132, 213$ and $321$. 

**Obj/notation** AxisTransformation Matrix EulerAngles Components
Consider a right-handed set of axes defined by a set of three unit basis vectors, $e$.

Right-handed means that the scalar triple product, $e_1 \times e_2 \cdot e_3 = +1$
Direction cosines

\[ \alpha_i = \hat{a} \cdot \hat{x}_i \]

\[ \alpha_1 = u = \cos \theta_1 \]
\[ \alpha_2 = v = \cos \theta_2 \]
\[ \alpha_3 = w = \cos \theta_3 \]
Consider a new orthonormal system consisting of right-handed base vectors $\hat{e}'_1$, $\hat{e}'_2$ and $\hat{e}'_3$

with the same origin, $o$, associated with the basis vectors. The vector $\vec{v}$ is clearly expressed equally well in either coordinate system:

$$\vec{v} = v_i \hat{e}_i = v'_i \hat{e}'_i$$

Note - same vector, different values of the components. We need to find a relationship between the two sets of components for the vector.
The two systems are related by the nine direction cosines, $a_{ij}$, which fix the cosine of the angle between the $i^{th}$ primed and the $j^{th}$ unprimed base vectors:

$$a_{ij} = \hat{e}_i' \cdot \hat{e}_j$$

Equivalently, $a_{ij}$ represent the components of $\hat{e}_i'$ in $\hat{e}_j$ according to the expression

$$\hat{e}_i' = a_{ij} \hat{e}_j$$
Rotation of axes in the x-y plane

\[ \mathbf{v}' = a\mathbf{v} = \begin{pmatrix} \cos \theta & \sin \theta \\ -\sin \theta & \cos \theta \end{pmatrix} \mathbf{v} \]

\[ a_{ij} = \hat{x}'_i \cdot \hat{x}_j \]

\( x, y = \text{old axes}; \ x', y' = \text{new axes} \)

Passive Rotation/Transformation of Axes
Example: rotation angle = 30°

\[ \mathbf{v}' = a\mathbf{v} = \begin{pmatrix} \cos 30° & \sin 30° \\ -\sin 30° & \cos 30° \end{pmatrix} \mathbf{v} = \begin{pmatrix} \sqrt{3}/2 & 1/2 \\ -1/2 & \sqrt{3}/2 \end{pmatrix} \mathbf{v} \]

\[ a_{ij} = \hat{x}'_i \cdot \hat{x}_j \]

\[ x, y = \text{old axes}; x', y' = \text{new axes} \]

Passive Rotation/Transformation of Axes
Rotation Matrices

\[
a_{ij} = \begin{pmatrix}
a_{11} & a_{12} & a_{13} \\
a_{21} & a_{22} & a_{23} \\
a_{31} & a_{32} & a_{33}
\end{pmatrix}
\]

Since an orthogonal matrix merely rotates a vector but does not change its length, the determinant is one, \( \det(a) = 1 \).
Moreover, each row and each column is a unit vector, so these six relations apply, resulting in only 3 independent parameters:

\[
\sqrt{a_{11}^2 + a_{12}^2 + a_{13}^2} = 1
\]
\[
\sqrt{\sum_i a_{1i}^2} = 1 , \sqrt{\sum_i a_{3i}^2} = 1 \quad , \text{etc.}
\]
Scalars, Vectors, Tensors

- **Scalar**: quantity that requires only one number, e.g. density, mass, specific heat.
- **Vector**: quantity that has direction as well as magnitude, e.g. velocity, current, magnetization; requires 3 numbers or *coefficients* (in 3D).
- **Tensor**: quantity that requires higher order descriptions but is the same, no matter what coordinate system is used to describe it, e.g. stress, strain, elastic modulus; requires 9 (or more, depending on rank) numbers or *coefficients*. 
Scalars, Vectors, Tensors: NOTATION

• General case: three dimensions
• Vector: needs 3 numbers or coefficients to quantify its x, y and z components.
• Two notations for vectors: “vector-tensor notation” where bold-face implies higher-than-scalar nature; “component notation” where a suffix(-es) show how many coefficients are needed.
• Vector: either \( b \) or \( b_i, i \in \{1,2,3\} \), or, \( i \in \{x,y,z\} \).
• 2nd rank tensor: either \( T \) or \( T_{ij}, i,j \in \{1,2,3\} \)
• Advantage of vector-tensor notation is that the equations work in any reference frame. By contrast, when component notation is used, the actual values of the coefficients depend on which reference frame is used.
• If you see subscripts attached to a quantity, it is (almost always) a tensor and the Einstein summation convention is assumed. The Einstein summation convention says that a repeated index (on the RHS) implies summation over that index (typically 1,2, and 3 in 3D).