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## Anisotropic Elasticity

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Texture, Microstructure \& Anisotropy A.D. Rollett \& M. De Graef

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## Bibliography

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1. How do we write the relationship between (tensor) stress and (tensor) strain? $\sigma=\mathrm{C}: \varepsilon$. How about the other way around? $\varepsilon=S: \sigma$. What are "stiffness" and "compliance" in this context? The stiffness tensor is the collection of coefficients that connect all the different stress coefficients/components to all the different strain coefficients/components. How do we express this in Voigt or vector-matrix notation? The only difference is that the stress and strain are vectors and the stiffness and compliance are matrices. If indices are used then stress and strain each have two indices and the stiffness and compliance each have four.
2. What are the relationships between the coefficients of the ( $4^{\text {th }}$ rank) stiffness tensor and the stiffness matrix ( $6 \times 6$ )? See the notes for details but, e.g., $\{11,22,33\}_{\text {tensor }}$ correspond to $\{1,2,3\}_{\text {matrix }}$. E.g. $\mathrm{C}_{12}$ (matrix) $=\mathrm{C}_{1122}$ (tensor). What about the compliance tensor and matrix? Here, more care is required because certain coefficients have factors of 2 or 4.
3. What does work conjugacy mean? The energy stored in a body when elastic strains and stresses are present is calculated as the product of the stress and strain, which means that the work done makes the strain and stress conjugate (joined) variables. What does this mean for the relationships between ( $2^{\text {nd }}$ rank) tensor stress and its vector form? What about strain? Answering these two together, we note that work conjugacy means that whatever notation is used to express stress and strain, the product of the two must be the same because of conservation of energy. This then explains why factors of two are used in the conversion to/from matrix to tensor representations of the shear components of strain (but not the normal strain components). These factors of two could have been applied to stress, but by convention we do this for strain.
4. How do we write the tensor transformation rule in vector-matrix notation? See the notes for details but the basic idea is that a $6 \times 6$ matrix (that can be applied to a stiffness or compliance tensor) is formed from the coefficients of the transformation matrix.
5. How do we apply crystal symmetry to elastic moduli (e.g. the stiffness tensor)? We apply a symmetry operator to the (stiffness) tensor and set the new and old versions of the tensor equal to each other, coefficient by coefficient. What net effect does it have on the stiffness matrix for cubic materials? Applying the cubic crystal symmetry to the stiffness tensor reduces most of the coefficients to zero and there are only 3 independent coefficients that remain.

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## Q\&A, part 2

6. How do we convert from stiffness to compliance (and vice versa)? The detailed mathematics is out of scope for this course. It is sufficient to know that the two tensors combine to form a $4^{\text {th }}$ rank identity tensor, from which one can obtain algebraic relationships as given in the notes. Be aware that these formulae depend on the crystal symmetry (as do the compliance \& stiffness tensors themselves).
7. How do we apply symmetry (and transformations of axes in general) to the property of anisotropic elasticity? There are two answers. The first answer is that one can apply the tensor transformation rule, just as explained in previous lectures. Generate the transformation matrix with any the methods described (i.e. dot products between old and new axes, or using the combination of axis and angle). Then write out the transformation with 4 copies of the matrix taking care to specify the indices correctly. The alternative answer is to generate a $6 \times 6$ transformation matrix that can be used with vector-matrix (Voigt) notation for either the stress, strain ( $6 \times 1$ ) vectors or the modulus ( $6 \times 6$ ) matrix.
8. How do we show that symmetry reduces the number of independent coefficients in an anisotropic elasticity modulus tensor? Given a symmetry matrix, one proceeds just as in the previous examples i.e. apply symmetry and then equate individual coefficients to find the cases of either zero or equality(between different coefficients).
9. How do we calculate the (anisotropic) elastic (Young's) modulus in an arbitrary direction? This looks ahead to the next lecture. The idea is to realize that a tensile test is such that there is only one nonzero coefficient in the stress tensor (or vector); the strain tensor, however, has to have more than one non-zero coefficient (because of the Poisson effect). Therefore one uses the relationship that strain = compliance $x$ stress. By rotating the compliance tensor such that one axis (usually $x$ ) is parallel to the desired direction, one obtains the Young's modulus in that direction as $1 / \mathrm{S}_{11}$.

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## Notation

| F | Stimulus (field) | $a$ |
| :--- | :--- | :--- |
| transformation matrix |  |  |
| R | Response | $W$ |
| work done (energy) |  |  |
| P | Property | $d W$ work increment |
| $\mathbf{j}$ | electric current | $I$ |
| identity matrix |  |  |
| E | electric field | $O$ |
| D | electric polarization | $Y$ |
| Young's modulus |  |  |
| $\varepsilon$ | Strain (also, permutation | $\delta$ |
| Kronecker delta |  |  |
|  | tensor) | $e$ |
| $\sigma$ axis (unit) vector |  |  |
| $\rho$ | Stress (or conductivity) | $T$ |
| tensor |  |  |

## Objective

- The objective of this lecture is to provide a mathematical framework for the description of properties, especially when they vary with direction.
- A basic property that occurs in almost applications is elasticity. Although elastic response is linear for all practical purposes, it is often anisotropic (composites, textured polycrystals etc.).
- Why do we care about elastic anisotropy? In composites, especially fibre composites, it is easy to design in substantial anisotropy by varying the lay-up of the fibres.
- Geologists are very familiar with elastic anisotropy and exploit it for understanding seismic results; see, e.g., https://en.wikipedia.org/wiki/Seismic_anisotropy .


## In Class Questions

1. Why is plastic yielding a non-linear property, in contrast to elastic deformation?
2. What is the definition of a tensor?
3. Why is stress is $2^{\text {nd }}$-rank tensor?
4. Why is elastic stiffness a $4^{\text {th }}$-rank tensor?
5. What is "matrix notation" (in the context of elasticity)?
6. What are the relationships between tensor and matrix coefficients for stress? Strain? Stiffness? Compliance?
7. Why do we need factors of 2 and 4 in some of these conversion factors?
8. How do we use crystal symmetry to decrease the number of coefficients needed to describe stiffness and compliance?
9. How many independent coefficients are needed for stiffness (and compliance) in cubic crystals? In isotropic materials?
10. How do we express the directional dependence of Young's modulus?
11. What is Zener's anisotropy factor?

## Anisotropy: Practical Applications

- The practical applications of anisotropy of composites, especially fiber-reinforced composites are numerous.
- The stiffness of fiber composites varies tremendously with direction. Torsional rigidity is very important in car bodies, boats, aeroplanes etc.
- Even in monolithic polymers (e.g. drawn polyethylene) there exists large anisotropy because of the alignment of the long-chain molecules.


## What is a Material Property?

- When we "do something" to a material, the material will "react" in a particular way
- More formally: when we apply a field, we obtain a response
- We can "do something" to a material by applying a field, ( $F$ ) then measuring its "reaction" or response (R)

This is one of the most important and central problems in MSE:

What is the response is for a given field?

## What is a Material Property?

## One of the most important problems in MSE: What is the response is for a given field?

- Response (R) is a functional of the applied field (F): R=R(F)
- By using a finite number of terms of its Taylor series, we can approximate the response (R) of a given material under an applied field (F)

$$
\begin{aligned}
& \text { "State" of the Response is proportional to Response is NOT proportional to } \\
& \text { material at zero field } \\
& \text { Response is proportional to } \\
& \text { field (Linear material property) } \\
& \text { Response is NOT proportional to } \\
& \text { field (Non-linear material properties) } \\
& \mathcal{R}=\mathcal{R}_{0}+\frac{1}{1!} \frac{\partial \mathcal{R}}{\partial \mathcal{F}}{\underset{\mathcal{F}}{ }=0}_{\mathcal{F}}+\frac{1}{2!} \frac{\partial^{2} \mathcal{R}}{\partial \mathcal{F}^{2}}{ }_{\mathcal{F}=0} \mathcal{F}^{2}+\frac{1}{3!} \frac{\partial^{3} \mathcal{R}}{\partial \mathcal{F}^{3}}{ }_{\mathcal{F}=0} \mathcal{F}^{3}+\ldots \\
& =\mathcal{R}_{0}+\left.\sum_{n=1}^{\infty} \frac{1}{n!} \frac{\partial^{n} \mathcal{R}}{\partial \mathcal{F}^{n}}\right|_{\mathcal{F}=0} \mathcal{F}^{n}
\end{aligned}
$$

If the complete Taylor series is needed, then we use a more general "constitutive relation" to describe the response in terms of material properties (i.e., hysteresis)

## Linear properties

- Certain properties, such as elasticity in most cases, are linear which means that we can simplify even further to obtain

$$
R=R_{0}+\mathbf{P} F
$$

or if $R_{0}=0$,


In tension, $C \equiv$ Young's modulus, $Y$ or $E$.

## Non-Linear properties, example

- Another important example of non-linear anisotropic properties is plasticity, i.e. the irreversible deformation of solids.
- A typical description of the response at plastic yield (what happens when you load a material to its yield stress) is elastic-perfectly plastic. In other words, the material responds elastically until the yield stress is reached, at which point the stress remains constant (strain rate unlimited).

- A more realistic description is a power-law with a large exponent, $\mathrm{n}^{\sim 50}$. The stress is scaled by the crss, and be expressed as either shear stressshear strain rate [graph], or tensile stress-tensile strain [equation].
$\dot{\varepsilon}=\left(\frac{\sigma}{\sigma_{\text {yield }}}\right)^{n}$
[Kocks]


## General Anisotropic Properties

- There is no a-priori reason for a property to be the same in all directions.
- There is no a-priori reason for a response to be in the same direction as the applied field.
- Linear and non-linear properties can be anisotropic; a constitutive relation can also be anisotropic.
- Both single and poly-crystals can be anisotropic


## General Anisotropic Properties

- Many different properties of crystals can be described as tensors.
- The rank of each tensor property depends, naturally, on the nature of the quantities related by the property.


## Application example: quartz oscillators

- Piezoelectric quartz crystals are commonly used for frequency control in watches and clocks. Despite having small values of the piezoelectric coefficients, quartz has positive aspects of low losses and the availability of orientations with negligible temperature sensitivity. The property of piezoelectricity relates strain to electric field, or polarization to stress.
- $\quad \varepsilon_{i j}=d_{i j k} E_{k}$
- PZT , lead zirconium titanate $\mathrm{PbZr}_{1-\mathrm{x}} \mathrm{Ti}_{\mathrm{x}} \mathrm{O}_{3}$, is another commonly used piezoelectric material.



## Piezoelectric Devices

- The property of piezoelectricity relates strain to electric field, or polarization to stress.

$$
\varepsilon_{i j}=d_{i j k} E_{k}
$$

- PZT, lead zirconium titanate $\mathrm{PbZr}_{1-\mathrm{x}} \mathrm{Ti}_{\mathrm{x}} \mathrm{O}_{3}$, is another commonly used piezoelectric material.


Note: Newnham consistently uses vector-matrix notation, rather than tensor notation. We will explain how this works later on.

Fig. 12.12 Ceramic multilayer actuators consist of thin layers of piezoelectric ceramic and metal electrodes. In contrast to traditional piezoelectrics, even low voltages produce large forces and substantial displacements. A tradeoff exists between force and displacement. The multilayer stack utilizing the $d_{33}$ coefficient give kilonewton forces capable of pushing heavy weights through small distances. Bimorph benders make use of the smaller transverse of $d_{31}$ coefficients to give larger displacements in the millimeter range, but only small forces.

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## Piezoelectric Crystals

- How is it that crystals can be piezoelectric?
- The answer is that the bonding must be ionic to some degree (i.e. there is a net charge on the different elements) and the arrangement of the atoms must be non-centrosymmetric.
- PZT is a standard piezoelectric material. It has Pb atoms at the cell corners ( $a \sim 4 \AA$ ), O on face centers, and a Ti or Zr atom near the body center. Below a certain temperature (Curie T), the cell transforms from cubic (high T) to tetragonal (low T). Applying stress distorts the cell, which changes the electric displacement in different ways (see figure).
- Although we can understand the effect at the single crystal level, real devices (e.g. sonar transducers) are polycrystalline. The operation is much complicated than discussed here, and involves "poling" to maximize the response, which in turns involves motion of domain walls.
(a)

(b)

(c)

(d)


Fig. 12.11 Structure-property relations for the intrinsic piezoelectric effect in $\mathrm{PbTiO}_{3}$ In the unstressed state there is an electric dipole associated with the off-center shift of the titanium atom. Under stress, this dipole can be increased $\left(d_{33}\right)$, decreased $\left(d_{31}\right)$, or tilted ( $d_{15}$ ).
[Newnham]

## Mathematical Descriptions

- Mathematical descriptions of properties are available.
- Mathematics, or a type of mathematics provides a quantitative framework. It is always necessary, however, to make a correspondence between mathematical variables and physical quantities.
- In group theory one might say that there is a set of mathematical operations \& parameters, and a set of physical quantities and processes: if the mathematics is a good description, then the two sets are isomorphous.
- This lecture makes extensive use of tensors. A tensor is a quantity that can be transformed from one set of axes to another via the tensor transformation rule (next slide).


## 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.
- It says that any tensor quantity can be transformed from one reference frame to another; this transformation of axes is sometimes called a passive rotation.

Vector:
$2^{\text {nd }}$ rank
$3^{\text {rd }}$ rank
$4^{\text {th }}$ rank

$$
\begin{aligned}
& V_{i}^{\prime}=a_{i j} V_{j} \\
& T_{i j}^{\prime}=a_{i k} a_{j l} T_{k l} \\
& T_{i j k}^{\prime}=a_{i l} a_{j m} a_{k n} T_{l m n} \\
& T_{i j k l}^{\prime}=a_{i m} a_{j n} a_{k o} a_{l p} T_{m n o p}
\end{aligned}
$$

This rule is a critical piece of information, which you must know how to use.

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## Tensor: definition, contd.

- We will denote tensors by subscripted variables, e.g.,
- $\varepsilon_{\mathrm{ij}}$ (rank-2)
- $\mathrm{d}_{\mathrm{pqr}}$ (rank-3)
- C Imno $^{\text {(rank-4) }}$
- The number of subscripts represents the rank of the tensor. Vectors = rank-1 tensors, scalars = rank-0 tensors.
- Subscripts are known as dummy indices; it doesn't matter which character you use . . . as long as you use it consistently.
- Two (2) subscript "rules":
- Conservation of subscripts
- Summation implied over each pair of subscripts on one side of an equation


## Einstein Convention

- The Einstein Convention, or summation rule for suffixes looks like this:

$$
A_{i}=B_{i j} C_{j}
$$

where " i " and " j " both are integer indexes whose range is $\{1,2,3\}$. So, to find each " $i$ th" component of $A$ on the LHS, we sum up over the repeated index, "j", on the RHS:

$$
\begin{aligned}
& A_{1}=B_{11} C_{1}+B_{12} C_{2}+B_{13} C_{3} \\
& A_{2}=B_{21} C_{1}+B_{22} C_{2}+B_{23} C_{3} \\
& A_{3}=B_{31} C_{1}+B_{32} C_{2}+B_{33} C_{3}
\end{aligned}
$$

## Examples of Materials Properties as Tensors

- Table 1 shows a series of tensors that are of importance for material science. The tensors are grouped by rank and are also labeled (in the last column) by $E$ (equilibrium property) or $T$ (transport property). The number following this letter indicates the maximum number of independent, nonzero elements in the tensor, taking into account symmetries imposed by thermodynamics.
- The Field and Response columns contain the following symbols: $\Delta T=$ temperature difference, $\Delta S=$ entropy change, $E_{i}=$ electric field components, $H_{i}=$ magnetic field components, $\varepsilon_{i \mathrm{ij}}=$ mechanical strain, $D_{i}=$ electric displacement, $B_{i}=$ magnetic induction, $\sigma_{\mathrm{ij}}=$ mechanical stress, $\Delta \beta_{\mathrm{ij}}=$ change of the impermeability tensor, $j_{\mathrm{i}}=$ electrical current density, $\nabla_{\mathrm{j}} T=$ temperature gradient, $h_{\mathrm{i}}=$ heat flux, $\nabla_{\mathrm{j}} c=$ concentration gradient, $m_{\mathrm{i}}=$ mass flux, $\rho_{\mathrm{i}}^{\mathrm{a}}=$ anti-symmetric part of resistivity tensor, $\rho_{\mathrm{i}}^{\mathrm{s}}=$ symmetric part of resistivity tensor, $\Delta \rho_{\mathrm{ij}}=$ change in the component $i j$ of the resistivity tensor, $l_{i}=$ direction cosines of wave direction in crystal, $G=$ gyration constant,


## Examples of Materials Properties as Tensors

| Property | Symbol | Field | Response | Type\# |
| :---: | :---: | :---: | :---: | :---: |
| Tensors of Rank 0 (Scalars) |  |  |  |  |
| Specific Heat | C | $\Delta T$ | $T \Delta S$ | E1 |
| Tensors of Rank 1 (Vectors) |  |  |  |  |
| Electrocaloric | $p_{i}$ | $E_{i}$ | $\Delta S$ | E3 |
| Magnetocaloric | $q_{i}$ | $H_{i}$ | $\Delta S$ | E3 |
| Pyroelectric | $p_{i}^{\prime}$ | $\Delta T$ | $D_{i}$ | E3 |
| Pyromagnetic | $q_{i}^{\prime}$ | $\Delta T$ | $B_{i}$ | E3 |
| Tensors of Rank 2 |  |  |  |  |
| Thermal expansion | $\alpha_{i j}$ | $\Delta T$ | $\epsilon_{i j}$ | E6 |
| Piezocaloric effect | $\alpha_{i j}^{\prime}$ | $\sigma_{i j}$ | $\Delta S$ | E6 |
| Dielectric permittivity | $\kappa_{i j}$ | $E_{j}$ | $D_{i}$ | E6 |
| Magnetic permeability | $\mu_{i j}$ | $H_{j}$ | $B_{i}$ | E6 |
| Optical activity | $g_{i j}$ | $l_{i} l_{j}$ | $G$ | E6 |
| Magnetoelectric polarization | $\lambda_{i j}$ | $H_{j}$ | $D_{i}$ | E9 |
| Converse magnetoelectric polarization | $\lambda_{i j}^{\prime}$ | $E_{j}$ | $B_{i}$ | E9 |
| Electrical conductivity (resistivity) | $\sigma_{i j}\left(\rho_{i j}\right)$ | $E_{j}\left(j_{j}\right)$ | $j_{i}\left(E_{i}\right)$ | T6 |
| Thermal conductivity | $K_{i j}$ | $\nabla_{j} T$ | $h_{i}$ | T6 |
| Diffusivity | $D_{i j}$ | $\nabla_{j} c$ | $m_{i}$ | T6 |
| Thermoelectric power | $\Sigma_{i j}$ | $\nabla_{j} T$ | $E_{i}$ | T9 |
| Hall effect | $R_{i j}$ | $B_{j}$ | $\rho_{i}^{a}$ | T9 |

[^0] direction in crystal, $G=$ gyration constant

| Property | Symbol | Field | Response | Type\# |  |
| :--- | :---: | :---: | :---: | :---: | :---: |
| Tensors of Rank 3 |  |  |  |  |  |
| Piezoelectricity | $d_{i j k}$ | $\sigma_{j k}$ | $D_{i}$ | E18 |  |
| Converse piezoelectricity | $d_{i j k}^{\prime}$ | $E_{k}$ | $\epsilon_{i j}$ | E18 |  |
| Piezomagnetism | $Q_{i j k}$ | $\sigma_{j k}$ | $B_{i}$ | E18 |  |
| Converse piezomagnetism | $Q_{i j k}^{\prime}$ | $H_{k}$ | $\epsilon_{i j}$ | E18 |  |
| Electro-optic effect | $r_{i j k}$ | $E_{k}$ | $\Delta \beta_{i j}$ | E18 |  |
| Nernst tensor | $\Sigma_{i j k}$ | $\nabla_{j} T B_{k}$ | $E_{i}$ | T27 |  |
|  |  |  |  |  |  |
| Elasticity | Tensors of Rank 4 | $\epsilon_{i j}\left(\sigma_{i j}\right)$ | E21 |  |  |
| Electrostriction | $s_{i j k l}\left(c_{i j k l}\right)$ | $\sigma_{k l}\left(\epsilon_{k l}\right)$ | $\epsilon_{i j}$ | E36 |  |
| Photoelasticity | $\gamma_{i j k l}$ | $E_{k} E_{l}$ | $\epsilon_{i j}$ | E36 |  |
| Kerr effect | $q_{i j k l}$ | $\sigma_{k l}$ | $\Delta \beta_{i j}$ | $\Delta \beta_{i j}$ |  |
| Magnetoresistance | $p_{i j k l}$ | $E_{k} E_{l}$ | E36 |  |  |
| Piezoresistance | $\xi_{i j k l}$ | $B_{k} B_{l}$ | $\rho_{i j}^{s}$ | T36 |  |
| Magnetothermoelectric power | $\Pi_{i j k l}$ | $\Sigma_{i j k l}$ | $\nabla_{j l} T B_{k} B_{l}$ | $\Delta \rho_{i j}$ |  |
| Second order Hall effect | $\rho_{i j k l}$ | $B_{j} B_{k} B_{l}$ | $\rho_{i}^{2}$ | T54 |  |
|  |  |  |  |  |  |
| Third order elasticity | Tensors of Rank 6 | $\sigma_{i j}$ |  |  |  |

Table 1: Materials property and transport tensors (adapted from Nowick (1995) Crystal Properties via Group Theory, Cambridge University Press.
$\left(\begin{array}{c}\Delta S \\ D_{x} \\ D_{y} \\ D_{z} \\ B_{x} \\ B_{y} \\ B_{z} \\ \epsilon_{x x} \\ \epsilon_{y y} \\ \epsilon_{z z} \\ \epsilon_{y z} \\ \epsilon_{x z} \\ \epsilon_{x y}\end{array}\right)=\left(\begin{array}{c|ccc|ccc|cccccc}\frac{C}{T} & p_{x} & p_{y} & p_{z} & q_{x} & q_{y} & q_{z} & \alpha_{x x}^{\prime} & \alpha_{y y}^{\prime} & \alpha_{z z}^{\prime} & \alpha_{y z}^{\prime} & \alpha_{x z}^{\prime} & \alpha_{x y}^{\prime} \\ \hline p_{x}^{\prime} & \kappa_{x x} & \kappa_{x y} & \kappa_{x z} & \lambda_{x x} & \lambda_{x y} & \lambda_{x z} & d_{x x x} & d_{x y y} & d_{x z z} & d_{x y z} & d_{x x z} & d_{x x y} \\ p_{y}^{\prime} & \kappa_{y x} & \kappa_{y y} & \kappa_{y z} & \lambda_{y x} & \lambda_{y y} & \lambda_{y z} & d_{y x x} & d_{y y y} & d_{y z z} & d_{y y z} & d_{y x z} & d_{y x y} \\ p_{z}^{\prime} & \kappa_{z x} & \kappa_{z y} & \kappa_{z z} & \lambda_{z x} & \lambda_{z y} & \lambda_{z z} & d_{z x x} & d_{z y y} & d_{z z z} & d_{z y z} & d_{z x z} & d_{z x y} \\ \hline q_{x}^{\prime} & \lambda_{x x}^{\prime} & \lambda_{x y}^{\prime} & \lambda_{x z}^{\prime} & \mu_{x x} & \mu_{x y} & \mu_{x z} & Q_{x x x} & Q_{x y y} & Q_{x z z} & Q_{x y z} & Q_{x x z} & Q_{x x y} \\ q_{y}^{\prime} & \lambda_{y x}^{\prime} & \lambda_{y y}^{\prime} & \lambda_{y z}^{\prime} & \mu_{y x} & \mu_{y y} & \mu_{y z} & Q_{y x x} & Q_{y y y} & Q_{y z z} & Q_{y y z} & Q_{y x z} & Q_{y x y} \\ q_{z}^{\prime} & \lambda_{z x}^{\prime} & \lambda_{z y}^{\prime} & \lambda_{z z}^{\prime} & \mu_{z x} & \mu_{z y} & \mu_{z z} & Q_{z x x} & Q_{z y y} & Q_{z z z} & Q_{z y z} & Q_{z x z} & Q_{z x y} \\ \hline \alpha_{x x} & d_{x x x}^{\prime} & d_{x x y}^{\prime} & d_{x x z}^{\prime} & Q_{x x x}^{\prime} & Q_{x x y}^{\prime} & Q_{x x z}^{\prime} & s_{x x x x} & s_{x x y y} & s_{x x z z} & s_{x x y z} & s_{x x x z} & s_{x x x y} \\ \alpha_{y y}^{\prime} & d_{y y x}^{\prime \prime} & d_{y y y}^{\prime} & d_{y y z}^{\prime} & Q_{y y x}^{\prime} & Q_{y y y}^{\prime} & Q_{y y z}^{\prime} & s_{y y x x} & s_{y y y y} & s_{y y z z} & s_{y y y z} & s_{y y x z} & s_{y y x y} \\ \alpha_{z z} & d_{z z x}^{\prime} & d_{z z y}^{\prime} & d_{z z z}^{\prime} & Q_{z z x}^{\prime} & Q_{z z y}^{\prime} & Q_{z z z}^{\prime} & s_{z z x x} & s_{z z y y} & s_{z z z z} & s_{z z y z} & s_{z z x z} & s_{z z x y}^{\prime} \\ \alpha_{y z}^{\prime} & d_{y z x}^{\prime \prime} & d_{y z y}^{\prime} & d_{y z z}^{\prime} & Q_{y z x}^{\prime} & Q_{y z y}^{\prime} & Q_{y z z}^{\prime} & s_{y z x x} & s_{y z y y} & s_{y z z z} & s_{y z y z} & s_{y z x z} & s_{y z x y} \\ \alpha_{x z} & d_{x z x}^{\prime} & d_{x z y}^{\prime} & d_{x z z}^{\prime} & Q_{x z x}^{\prime} & Q_{x z y}^{\prime} & Q_{x z z}^{\prime} & s_{x z x x} & s_{x z y y} & s_{x z z z} & s_{x z y z} & s_{x z x z} & s_{x z x y} \\ \alpha_{x y}^{\prime} & d_{x y x}^{\prime \prime} & d_{x y y}^{\prime} & d_{x y z}^{\prime} & Q_{x y x}^{\prime} & Q_{x y y}^{\prime} & Q_{x y z}^{\prime} & s_{x y x x} & s_{x y y y} & s_{x y z z} & s_{x y y z} & s_{x y x z} & s_{x y x y}\end{array}\right)\left(\begin{array}{l}\Delta T \\ E_{x} \\ E_{y} \\ E_{z} \\ H_{x} \\ H_{y} \\ H_{z} \\ \sigma_{x x} \\ \sigma_{y y} \\ \sigma_{z z} \\ \sigma_{y z} \\ \sigma_{x z} \\ \sigma_{x y}\end{array}\right)$

## Principal Effects

Courtesy of Prof. M. De Graef
Electrocaloric = pyroelectric

- Magnetocaloric = pyromagnetic

Thermal expansion = piezocaloric
Magnetoelectric and converse magnetoelectric
Piezoelectric and converse piezoelectric
Piezomagnetic and converse piezomagnetic
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$$
\begin{aligned}
\Delta S & =\frac{C}{T} \Delta T+p_{i} E_{i}+q_{i} H_{i}+\alpha_{i j}^{\prime} \sigma_{i j} ; \\
D_{i} & =p_{i}^{\prime} \Delta T+\kappa_{i j} E_{j}+\lambda_{i j} H_{j}+d_{i j k} \sigma_{j k} ; \\
B_{i} & =q_{i}^{\prime} \Delta T+\lambda_{i j} E_{j}+\mu_{i j} H_{j}+Q_{i j k} \sigma_{j k} ; \\
\epsilon_{i j} & =\alpha_{i j} \Delta T+d_{i j k}^{\prime} E_{k}+Q_{i j k}^{\prime} H_{k}+s_{i j k l} \sigma_{k l} .
\end{aligned}
$$

# Principal Effects 

Courtesy of Prof. M. De Graef
1st rank cross effects

2nd rank cross effects

## 3rd rank cross effects

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## Examples of Materials Properties as Tensors

- A tensor of rank $n$ has $3 n$ components; but they are not all independent due to the presence of symmetry which reduces the number of independent components
- Two types of symmetry to consider:
- Thermodynamic symmetry (from Helmholtz free energy)

$$
\mathrm{K}_{\mathrm{ij}}=\mathrm{K}_{\mathrm{ji}}
$$

True for tensors which connect conjugate variables (e.g. $\sigma-\varepsilon, H-B$, etc. )

- Crystallographic symmetry (Group Symmetry)
- Neumann's Principle: "The symmetry of any physical property of a crystal must include the symmetry elements of the point group of the crystal."
- Definition \#1: We call a tensor "symmetric" if interchanging two subscripts does not change the value ( $\varepsilon_{\mathrm{ij}}=\varepsilon_{\mathrm{ij}}$ )
- Definition \#2: We call a tensor is "antisymmetric" if the interchange of two subscripts causes a sign change $\left(\varepsilon_{\mathrm{ij}}=-\varepsilon_{\mathrm{j}}\right)$

Note the minus (-) sign!
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## Elasticity

- Elasticity: example of a property that requires tensors to describe it fully.
- Even in cubic metals, a crystal is quite anisotropic. The [111] in many cubic metals is stiffer than the [100] direction.
- Even in cubic materials, 3 numbers/coefficients/moduli are required to describe elastic properties; isotropic materials only require 2.
- Familiarity with Miller indices, suffix notation, Einstein convention, Kronecker delta, permutation tensor, and tensors is assumed.


## Elastic Anisotropy: 1

- First, we restate the linear elastic relations for the properties Compliance, written S, and Stiffness, written C (admittedly not very logical choice of notation), which connect stress, $\sigma$, and strain, $\varepsilon$. We write it first in vector-tensor notation with ":" signifying inner product (i.e., add up terms that have a common suffix or index in them):

$$
\begin{aligned}
& \sigma=C: \varepsilon \\
& \varepsilon=S: \sigma
\end{aligned}
$$

- In component form (with suffixes),

$$
\begin{aligned}
& \sigma_{\mathrm{ij}}=C_{\mathrm{ijk} \mid} \varepsilon_{\mathrm{kl}} \\
& \varepsilon_{\mathrm{ij}}=S_{\mathrm{ij} \mathrm{j} \mathrm{kl}} \sigma_{\mathrm{kl}}^{\mathrm{ol}}
\end{aligned}
$$

## Elastic Anisotropy: 2

The definitions of the stress and strain tensors mean that they are both symmetric (second rank) tensors. Therefore we can see that

$$
\begin{aligned}
& \varepsilon_{23}=S_{2311} \sigma_{11} \\
& \varepsilon_{32}=S_{3211} \sigma_{11}=\varepsilon_{23}
\end{aligned}
$$

which means that,

$$
S_{2311}=S_{3211}
$$

and in general,

$$
S_{\mathrm{ijkl}}=S_{\mathrm{j} \mathrm{jkl}}
$$

This reduces considerably the number of different coefficients needed

## Elastic Anisotropy: 3

For $4^{\text {th }}$-order symmetric tensors, the 81 components (i.e. $3^{4}$ ) reduces to just 36 components since:

$$
S_{\mathrm{ijk} \mid}=S_{\mathrm{j} \mathrm{jk} \mid}=S_{\mathrm{ijlk}}=S_{\mathrm{jilk}}
$$

Becomes only:

$$
S_{i j}
$$

# Also known as the 6x6 Stiffness (C) or Compliance (S) Matrix 

## Stiffness Matrix

$$
C=\left[\begin{array}{llllll}
C_{11} & C_{12} & C_{13} & C_{14} & C_{15} & C_{16} \\
C_{12} & C_{22} & C_{23} & C_{24} & C_{25} & C_{26} \\
C_{13} & C_{23} & C_{33} & C_{34} & C_{35} & C_{36} \\
C_{14} & C_{24} & C_{34} & C_{44} & C_{45} & C_{46} \\
C_{15} & C_{25} & C_{35} & C_{45} & C_{55} & C_{56} \\
C_{16} & C_{26} & C_{36} & C_{46} & C_{56} & C_{66}
\end{array}\right]
$$

Vector-matrix notation (two indices for the moduli, one index for stress or strain); note that this matrix is symmetric, therefore there are only 21 independent coefficients, even for triclinic crystals (see later slides).

## Stiffness in sample coords.

- Consider how to express the elastic properties of a single crystal in the sample coordinates. In this case we need to rotate the ( $4^{\text {th }}$ rank) tensor stiffness from crystal coordinates to sample coordinates using the orientation (matrix), $a$ :

$$
c_{\mathrm{ijkl}}^{\prime}=a_{\mathrm{im}} a_{\mathrm{jn}} a_{\mathrm{ko}} a_{\mathrm{lp}} c_{\mathrm{mnop}}
$$

- Note how the transformation matrix appears four times because we are transforming a $4^{\text {th }}$ rank tensor!
- The axis transformation matrix, $a$, is sometimes also written as $\lambda$, also as the orientation matrix $g$.


## Young's modulus from compliance

- Young's modulus as a function of direction can be obtained from the compliance tensor as:

$$
E=1 / s_{1111}^{\prime}
$$

Using compliances and a stress boundary condition (only $\sigma_{11} \neq 0$ ) is most straightforward. To obtain $s_{1111}^{\prime}$, we simply apply the same transformation rule,

$$
s_{\mathrm{ijkl}}^{\prime}=a_{\mathrm{im}} a_{\mathrm{jn}} a_{\mathrm{ko}} a_{\mathrm{lp}} s_{\mathrm{mnop}}
$$

## Matrix, Miller Indices

- The general Rotation Matrix, $a$, can be represented as in the following:

- Here the Rows are the direction cosines for the 3 crystal axes, [100], [010], and [001] expressed in the sample coordinate system (pole figure).


## "Voigt" or "matrix" notation

- It is useful to re-express the three quantities involved in a simpler format. The stress and strain tensors are vectorized, i.e. converted into a $1 \times 6$ notation and the elastic tensors are reduced to $6 \times 6$ matrices.

$$
\begin{aligned}
& \left(\begin{array}{lll}
\sigma_{11} & \sigma_{12} & \sigma_{13}
\end{array}\right) \quad\left(\begin{array}{lll}
\sigma_{1} & \sigma_{6} & \sigma_{5}
\end{array}\right) \\
& \left(\begin{array}{lll}
\sigma_{21} & \sigma_{22} & \sigma_{23} \\
\sigma_{31} & \sigma_{32} & \sigma_{33}
\end{array}\right) \longleftrightarrow\left(\begin{array}{lll}
\sigma_{6} & \sigma_{2} & \sigma_{4} \\
\sigma_{5} & \sigma_{4} & \sigma_{3}
\end{array}\right) \\
& \longleftrightarrow\left(\sigma_{1}, \sigma_{2}, \sigma_{3}, \sigma_{4}, \sigma_{5}, \sigma_{6}\right)
\end{aligned}
$$

## "matrix notation", contd.

- Similarly for strain:

$$
\begin{aligned}
& \left(\begin{array}{lll}
\varepsilon_{11} & \varepsilon_{12} & \varepsilon_{13}
\end{array}\right) \\
& \left(\begin{array}{lll}
\varepsilon_{21} & \varepsilon_{22} & \varepsilon_{23} \\
\varepsilon_{31} & \varepsilon_{32} & \varepsilon_{33}
\end{array}\right) \longleftrightarrow\left(\begin{array}{ccc}
\varepsilon_{1} & \frac{1}{2} \varepsilon_{6} & \frac{1}{2} \varepsilon_{5}
\end{array}\right) \\
& \longleftrightarrow\left(\begin{array}{ccc}
\frac{1}{2} \varepsilon_{6} & \varepsilon_{2} & \frac{1}{2} \varepsilon_{4} \\
\frac{1}{2} \varepsilon_{5} & \frac{1}{2} \varepsilon_{4} & \varepsilon_{3}
\end{array}\right) \\
& \longleftrightarrow\left(\varepsilon_{1}, \varepsilon_{2}, \varepsilon_{3}, \varepsilon_{4}, \varepsilon_{5}, \varepsilon_{6}\right)
\end{aligned}
$$

The particular definition of shear strain used in the reduced notation happens to correspond to that used in mechanical engineering such that $\varepsilon_{4}$ is the change in angle between direction 2 and direction 3 due to deformation.

## Work conjugacy, matrix inversion

- The more important consideration is that the reason for the factors of two is so that work conjugacy is maintained.

$$
d W=\sigma \cdot d \varepsilon=\sigma_{\mathrm{ij}}: d \varepsilon_{\mathrm{ij}}=\sigma_{\mathrm{k}} \bullet d \varepsilon_{\mathrm{k}}
$$

Also we can combine the expressions

$$
\sigma=C \varepsilon \text { and } \varepsilon=S \sigma \text { to give: }
$$

$$
\begin{aligned}
& \sigma=C S \sigma, \text { which shows: } \\
& I=C S, \text { or, } C=S^{-1}
\end{aligned}
$$

## Tensor conversions: stiffness

- Lastly we need a way to convert the tensor coefficients of stiffness and compliance to the matrix coefficients. For stiffness, it is very simple because one substitutes values according to the following table, such that [vector-matrix] $C_{11}$
$=C_{1111}$ [tensor] for example.

| Tensor | 11 | 22 | 33 | 23 | 32 | 13 | 31 | 12 | 21 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Matrix | 1 | 2 | 3 | 4 | 4 | 5 | 5 | 6 | 6 |

## Axis Transformations

- It is still possible to perform axis transformations, as allowed for by the Tensor Rule. The coefficients can be combined together into a 6 by 6 matrix that can be used for $2^{\text {nd }}$ rank tensors such as stress and strain, below.

Table 10.1 Transformation matrices for stresses and strains written in matrix form

- Stress (in vector notation) transforms as: $X_{i}^{\prime}=\alpha_{i j} X_{j}$
- Strain (in vector notation) transforms as:
$x_{i}^{\prime}=\left(\alpha^{-1}{ }_{i j}\right)^{T} x_{j}$ where superscript "T" signifies transpose of the matrix.

| ( $\alpha$ ) | $\left(a_{11}^{2}\right)$ | $\left(a_{12}^{2}\right)$ | $\left(a_{13}^{2}\right)$ | (2a $a_{12} a_{13}$ ) | ( $2 a_{13} a_{11}$ ) | ( $2 a_{11} a_{12}$ ) |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  | $\left(a_{21}^{2}\right)$ | $\left(a_{22}^{2}\right)$ | $\left(a_{23}^{2}\right)$ | (2a $22 a_{23}$ ) | (2a $23 a_{21}$ ) | ( $2 a_{21} a_{22}$ ) |
|  | $\left(a_{31}^{2}\right)$ | $\left(a_{32}^{2}\right)$ | $\left(a_{33}^{2}\right)$ | (2a ${ }_{32} a_{33}$ ) | (2a $a_{33} a_{31}$ ) | ( $2 a_{31} a_{32}$ ) |
|  | ( $a_{21} a_{31}$ ) | ( $a_{22} a_{32}$ ) | $\left(a_{23} a_{33}\right)$ | $\left(a_{22} a_{33}+a_{23} a_{32}\right)$ | $\left(a_{21} a_{33}+a_{23} a_{31}\right)$ | $\left(a_{22} a_{31}+a_{21} a_{32}\right)$ |
|  | $\left(a_{31} a_{11}\right)$ | $\left(a_{32} a_{12}\right)$ | $\left(a_{33} a_{13}\right)$ | $\left(a_{12} a_{33}+a_{13} a_{32}\right)$ | $\left(a_{13} a_{31}+a_{11} a_{33}\right)$ | $\left(a_{11} a_{32}+a_{12} a_{31}\right)$ |
|  | ${ }_{\left(a_{11} a_{21}\right)}$ | ( $a_{12} a_{22}$ ) | ( $a_{13} a_{23}$ ) | $\left(a_{12} a_{23}+a_{13} a_{22}\right)$ | $\left(a_{13} a_{21}+a_{11} a_{23}\right)$ | $\left(a_{11} a_{22}+a_{12} a_{21}\right)$ |
| $\left(\alpha^{-1}\right)$ | $\left(a_{11}^{2}\right)$ | $\left(a_{21}^{2}\right)$ | $\left(a_{31}^{2}\right)$ | ( $2 a_{21} a_{31}$ ) | (2a $a_{31} a_{11}$ ) | $\left(2 a_{11} a_{21}\right)$ |
|  | $\left(a_{12}^{2}\right)$ | $\left(a_{22}^{2}\right)$ | $\left(a_{32}^{2}\right)$ | (2a $a_{22} a_{32}$ ) | (2a $a_{32} a_{12}$ ) | (2a $a_{12} a_{22}$ ) |
|  | $\left(a_{13}^{2}\right)$ | $\left(a_{23}^{2}\right)$ | $\left(a_{33}^{2}\right)$ | ( $2 a_{23} a_{33}$ ) | ( $2 a_{33} a_{13}$ ) | (2a $a_{13} a_{23}$ ) |
|  | ( $a_{12} a_{13}$ ) | $\left(a_{22} a_{23}\right)$ | $\left(a_{32} a_{33}\right)$ | $\left(a_{22} a_{33}+a_{32} a_{23}\right)$ | $\left(a_{12} a_{33}+a_{32} a_{13}\right)$ | $\left(a_{22} a_{13}+a_{12} a_{23}\right)$ |
|  | $\left(a_{13} a_{11}\right)$ | $\left(a_{23} a_{21}\right)$ | $\left(a_{33} a_{31}\right)$ | $\left(a_{21} a_{33}+a_{31} a_{23}\right)$ | $\left(a_{31} a_{13}+a_{11} a_{33}\right)$ | $\left(a_{11} a_{23}+a_{21} a_{13}\right)$ |
|  | ${ }_{\left(a_{11} a_{12}\right)}$ | $\left(a_{21} a_{22}\right)$ | $\left(a_{31} a_{32}\right)$ | $\left(a_{21} a_{32}+a_{31} a_{22}\right)$ | $\left(a_{31} a_{12}+a_{11} a_{12}\right)$ | $\left(a_{11} a_{22}+a_{21} a_{12}\right)$ |

[Newnham]

## Tensor conversions: compliance

- For compliance some factors of two are required and so the rule becomes:

$$
\begin{array}{rlr}
p S_{i j k l}=S_{m n} & \\
p=1 & & \text { m.AND.n } \in[1,2,3] \\
p=2 & \text { m.XOR.n } \in[1,2,3] \\
p=4 & \text { m.AND.n } \in[4,5,6]
\end{array}
$$

## Relationships between coefficients: $C$ in terms of $S$

Some additional useful relations between coefficients for cubic materials are as follows. Symmetrical relationships exist for compliances in terms of stiffnesses (next slide).

$$
\begin{aligned}
& \mathrm{C}_{11}=\left(\mathrm{S}_{11}+\mathrm{S}_{12}\right) /\left\{\left(\mathrm{S}_{11}-\mathrm{S}_{12}\right)\left(\mathrm{S}_{11}+2 \mathrm{~S}_{12}\right)\right\} \\
& \mathrm{C}_{12}=-\mathrm{S}_{12} /\left\{\left(\mathrm{S}_{11}-\mathrm{S}_{12}\right)\left(\mathrm{S}_{11}+2 \mathrm{~S}_{12}\right)\right\} \\
& \mathrm{C}_{44}=1 / \mathrm{S}_{44}
\end{aligned}
$$

## $S$ in terms of $C$

The relationships for $S$ in terms of $C$ are symmetrical to those for stiffnesses in terms of compliances (a simple exercise in algebra).

$$
\begin{aligned}
& \mathrm{S}_{11}=\left(\mathrm{C}_{11}+\mathrm{C}_{12}\right) /\left\{\left(\mathrm{C}_{11}-\mathrm{C}_{12}\right)\left(\mathrm{C}_{11}+2 \mathrm{C}_{12}\right)\right\} \\
& \mathrm{S}_{12}=-\mathrm{C}_{12} /\left\{\left(\mathrm{C}_{11}-\mathrm{C}_{12}\right)\left(\mathrm{C}_{11}+2 \mathrm{C}_{12}\right)\right\} \\
& \mathrm{S}_{44}=1 / \mathrm{C}_{44}
\end{aligned}
$$

$$
\mathrm{S}_{11}-\mathrm{S}_{12}=\left(\mathrm{C}_{11}+2 \mathrm{C}_{12}\right) /\left\{\left(\mathrm{C}_{11}-\mathrm{C}_{12}\right)\left(\mathrm{C}_{11}+2 \mathrm{C}_{12}\right)\right\}
$$

$$
\mathrm{S}_{11}-\mathrm{S}_{12}=1 /\left(\mathrm{C}_{11}-\mathrm{C}_{12}\right)
$$

## Neumann's Principle

- A fundamental natural law: Neumann's Principle: the symmetry elements of any physical property of a crystal must include the symmetry elements of the point group of the crystal. The property may have additional symmetry elements to those of the crystal (point group) symmetry. There are 32 crystal classes for the point group symmetry.
- F.E. Neumann 1885.


## Neumann, extended

- If a crystal has a defect structure such as a dislocation network that is arranged in a non-uniform way then the symmetry of certain properties may be reduced from the crystal symmetry. In principle, a finite elastic strain in one direction decreases the symmetry of a cubic crystal to tetragonal or less. Therefore the modified version of Neumann's Principle: the symmetry elements of any physical property of a crystal must include the symmetry elements that are common to the point group of the crystal and the defect structure contained within the crystal.


## Effect of crystal symmetry

- Consider an active rotation of the crystal, where $O$ is the symmetry operator. Since the crystal is indistinguishable (looks the same) after applying the symmetry operator, the result before, $R^{(1)}$, and the result after, $R^{(2)}$, must be identical:

$$
\left.\begin{array}{c}
R^{(1)}=\mathbf{P} F \\
R^{(2)}=O \mathbf{P} O^{T} F \\
R^{(1)} \stackrel{=}{\longleftrightarrow} R^{(2)}
\end{array}\right\}
$$

The two results are indistinguishable and therefore equal. It is essential, however, to express the property and the operator in the same (crystal) reference frame.

## Symmetry, properties, contd.

- Expressed mathematically, we can rotate, e.g. a second rank property tensor thus:

$$
\begin{aligned}
\mathbf{P}^{\prime} & =O \mathbf{P} O^{\mathrm{T}}=\mathbf{P}, \text { or, in coefficient notation, } \\
P_{i j}^{\prime} & =O_{i k} O_{i l} P_{k l}
\end{aligned}
$$

where $O$ is a symmetry operator.

- Since the rotated (property) tensor, $\mathbf{P}^{\prime}$, must be the same as the original tensor, $\mathbf{P}$, then we can equate coefficients:

$$
P_{i j}^{\prime}=P_{i j}
$$

- If we find, for example, that $P^{\prime}{ }_{21}=-P_{21}$, then the only value of $P_{21}$ that satisfies this equality is $P_{21}=0$.
- Remember that you must express the property with respect to a particular set of axes in order to use the coefficient form. In everything related to single crystals, always use the crystal axes as the reference frame!
- Homework question: based on cubic crystal symmetry, work out why a second rank tensor property can only have one independent coefficient.


## Effect of symmetry on stiffness matrix

- Why do we need to look at the effect of symmetry? For a cubic material, only 3 independent coefficients are needed as opposed to the 81 coefficients in a 4th rank tensor. The reason for this is the symmetry of the material.
- What does symmetry mean? Fundamentally, if you pick up a crystal, rotate [mirror] it and put it back down, then a symmetry operation [rotation, mirror] is such that you cannot tell that anything happened.
- From a mathematical point of view, this means that the property (its coefficients) does not change. For example, if the symmetry operator changes the sign of a coefficient, then it must be equal to zero.


## 2nd Rank Tensor Properties \& Symmetry

Table 3
The effect of crystal symmetry on properties represented by symmetrical second-rank tensors


[^1]- The table from Nye shows the number of independent, non-zero coefficients allowed in a 2nd rank tensor according to the crystal symmetry class.

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## Effect of symmetry on stiffness matrix

- Following Reid, p. 66 et seq.:

Apply a $-90^{\circ}$ rotation about the crystal-z axis (axis 3 )*,
$C^{\prime}{ }_{i j k l}=O_{i m} O_{j n} O_{k o} O_{l p} C_{\text {mпор }}:$
$C^{\prime}=C$

## Effect of symmetry, 2

- Using $\mathrm{P}^{\prime}=\mathrm{P}$, we can equate all the coefficients in the $6 \times 6$ matrix and find that:

$$
\begin{aligned}
& \mathrm{C}_{11}=\mathrm{C}_{22}, \mathrm{C}_{13}=\mathrm{C}_{23}, \mathrm{C}_{44}=\mathrm{C}_{35}, \mathrm{C}_{16}=-\mathrm{C}_{26}, \\
& \mathrm{C}_{14}=\mathrm{C}_{15}=\mathrm{C}_{24}=\mathrm{C}_{25}=\mathrm{C}_{34}=\mathrm{C}_{35}=\mathrm{C}_{36}=\mathrm{C}_{45}=\mathrm{C}_{46}= \\
& \mathrm{C}_{56}=0 .
\end{aligned}
$$

$$
C^{\prime}=\left[\begin{array}{cccccc}
C_{11} & C_{12} & C_{13} & 0 & 0 & C_{16} \\
C_{12} & C_{11} & C_{13} & 0 & 0 & -C_{16} \\
C_{13} & C_{13} & C_{33} & 0 & 0 & 0 \\
0 & 0 & 0 & C_{44} & 0 & 0 \\
0 & 0 & 0 & 0 & C_{44} & C_{46} \\
C_{16} & -C_{16} & 0 & 0 & C_{46} & C_{66}
\end{array}\right]
$$

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## Effect of symmetry, 3

- Thus by repeated applications of the symmetry operators, one can demonstrate (for cubic crystal symmetry) that one can reduce the 81 coefficients down to only 3 independent quantities. These become two in the case of isotropy.

$$
\left[\begin{array}{cccccc}
C_{11} & C_{12} & C_{12} & 0 & 0 & 0 \\
C_{12} & C_{11} & C_{12} & 0 & 0 & 0 \\
C_{12} & C_{12} & C_{11} & 0 & 0 & 0 \\
0 & 0 & 0 & C_{44} & 0 & 0 \\
0 & 0 & 0 & 0 & C_{44} & 0 \\
0 & 0 & 0 & 0 & 0 & C_{44}
\end{array}\right]
$$

$$
\left(\begin{array}{l}
\Delta S \\
D_{x} \\
D_{y} \\
D_{z} \\
B_{x} \\
B_{y} \\
B_{z} \\
\epsilon_{x x} \\
\epsilon_{y y} \\
\epsilon_{z z} \\
\epsilon_{y z} \\
\epsilon_{x z} \\
\epsilon_{x y}
\end{array}\right)=\left(\begin{array}{ccccccccccccc}
\frac{C}{T} & p_{x} & p_{y} & p_{z} & q_{x} & q_{y} & q_{z} & \alpha_{x x}^{\prime} & \alpha_{y y}^{\prime} & \alpha_{z z}^{\prime} & \alpha_{y z}^{\prime} & \alpha_{x z}^{\prime} & \alpha_{x y}^{\prime} \\
p_{x}^{\prime} & \kappa_{x x} & \kappa_{x y} & \kappa_{x z} & \lambda_{x x} & \lambda_{x y} & \lambda_{x z} & d_{x x x} & d_{x y y} & d_{x z z} & d_{x y z} & d_{x x z} & d_{x x y} \\
p_{y}^{\prime} & \kappa_{y x} & \kappa_{y y} & \kappa_{y z} & \lambda_{y x} & \lambda_{y y} & \lambda_{y z} & d_{y x x} & d_{y y y} & d_{y z z} & d_{y y z} & d_{y x z} & d_{y x y} \\
p_{z}^{\prime} & \kappa_{z x} & \kappa_{z y} & \kappa_{z z} & \lambda_{z x} & \lambda_{z y} & \lambda_{z z} & d_{z x x} & d_{z y y} & d_{z z z} & d_{z y z} & d_{z x z} & d_{z x y} \\
q_{x}^{\prime} & \lambda_{x x}^{\prime} & \lambda_{x y}^{\prime} & \lambda_{x z}^{\prime} & \mu_{x x} & \mu_{x y} & \mu_{x z} & Q_{x x x} & Q_{x y y} & Q_{x z z} & Q_{x y z} & Q_{x x z} & Q_{x x y} \\
q_{y}^{\prime} & \lambda_{y x}^{\prime} & \lambda_{y y}^{\prime} & \lambda_{y z}^{\prime} & \mu_{y x} & \mu_{y y} & \mu_{y z} & Q_{y x x} & Q_{y y y} & Q_{y z z} & Q_{y y z} & Q_{y x z} & Q_{y x y} \\
q_{z}^{\prime} & \lambda_{z x}^{\prime} & \lambda_{z y}^{\prime} & \lambda_{z z}^{\prime} & \mu_{z x} & \mu_{z y} & \mu_{z z} & Q_{z x x} & Q_{z y y} & Q_{z z z} & Q_{z y z} & Q_{z x z} & Q_{z x y} \\
\alpha_{x x} & d_{x x x}^{\prime} & d_{x x y}^{\prime} & d_{x x z}^{\prime} & Q_{x x x}^{\prime} & Q_{x x y}^{\prime} & Q_{x x z}^{\prime} & s_{x x x x} & s_{x x y y} & s_{x x z} & s_{x x y z} & s_{x x x z} & s_{x x x y} \\
\alpha_{y y} & d_{y y x}^{\prime} & d_{y y y}^{\prime} & d_{y y z}^{\prime} & Q_{y y x}^{\prime} & Q_{y y y}^{\prime} & Q_{y y z}^{\prime} & s_{y y x x} & s_{y y y y}^{\prime} & s_{y y z z}^{\prime} & s_{y y y z}^{\prime} & s_{y y x z}^{\prime} & s_{y y x y}^{\prime} \\
\alpha_{z z}^{\prime} & d_{z z x}^{\prime} & d_{z z y}^{\prime} & d_{z z z}^{\prime} & Q_{z z x}^{\prime} & Q_{z z y}^{\prime} & Q_{z z z}^{\prime} & s_{z z x x} & s_{z z y y} & s_{z z z z} & s_{z z y z} & s_{z z x z} & s_{z z x y} \\
\alpha_{y z} & d_{y z x}^{\prime} & d_{y z y}^{\prime} & d_{y z z}^{\prime} & Q_{y z x}^{\prime} & Q_{y z y}^{\prime} & Q_{y z}^{\prime} & s_{y z x x} & s_{y z y y} & s_{y z z} & s_{y z y} & s_{y z x z} & s_{y z x y} \\
\alpha_{x z} & d_{x z x}^{\prime} & d_{x z y}^{\prime} & d_{x z z}^{\prime} & Q_{x z x}^{\prime} & Q_{x z y}^{\prime} & Q_{x z z}^{\prime} & s_{x z x x}^{\prime} & s_{x z y y} & s_{x z z z}^{\prime} & s_{x y y}^{\prime} & s_{x z x z} & s_{x z x y}^{\prime} \\
d_{x y z}^{\prime} & Q_{x y x}^{\prime} & Q_{x y y}^{\prime} & Q_{x y z}^{\prime} & s_{x y x x} & s_{x y y y} & s_{x y z z} & s_{x y y z} & s_{x y x z} & s_{x y x y}
\end{array}\right)\left(\begin{array}{l}
\Delta T \\
E_{x} \\
E_{y} \\
E_{z} \\
H_{x} \\
H_{y} \\
H_{z} \\
\sigma_{x x} \\
\sigma_{y y} \\
\sigma_{z z} \\
\sigma_{y z} \\
\sigma_{x z} \\
\sigma_{x y}
\end{array}\right)
$$

General crystal symmetry shown above.

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$$
\left(\begin{array}{c}
\Delta S \\
D_{x} \\
D_{y} \\
D_{z} \\
B_{x} \\
B_{y} \\
B_{z} \\
\epsilon_{x x} \\
\epsilon_{y y} \\
\epsilon_{z z} \\
\epsilon_{y z} \\
\epsilon_{x z} \\
\epsilon_{x y}
\end{array}\right)=\left(\begin{array}{cccccccccccc}
C & p_{x} & p_{x} & p_{z} & q_{x} & q_{x} & q_{z} & \alpha_{x x} & \alpha_{x x} & \alpha_{z z} & 0 & 0 \\
p_{x} & \kappa_{x x} & 0 & 0 & \lambda_{x x} & \lambda_{x y} & 0 & 0 & 0 & 0 & d_{x y z} & d_{x x z} \\
p_{x} & 0 & \kappa_{x x} & 0 & -\lambda_{x y} & \lambda_{x x} & 0 & 0 & 0 & 0 & -d_{x x z} & d_{x y z} \\
p_{z} & 0 & 0 & \kappa_{z z} & 0 & 0 & \lambda_{z z} & d_{z x x} & d_{z x x} & d_{z z z} & 0 & 0 \\
q_{x} & \lambda_{x x} & -\lambda_{x y} & 0 & \mu_{x x} & 0 & 0 & 0 & 0 & 0 & Q_{x y z} & Q_{x x z} \\
q_{x} & \lambda_{x y} & \lambda_{x x} & 0 & 0 & \mu_{x x} & 0 & 0 & 0 & 0 & -Q_{x x z} & Q_{x y z} \\
q_{z} & 0 & 0 & \lambda_{z z} & 0 & 0 & \mu_{z z} & Q_{z x x} & Q_{z x x} & Q_{z z z} & 0 & 0 \\
\alpha_{x x} & 0 & 0 & d_{z x x} & 0 & 0 & Q_{z x x} & s_{x x x x} & s_{x x y y} & s_{x x z} & 0 & 0 \\
\alpha_{x x} & 0 & 0 & d_{z x x} & 0 & 0 & Q_{z y y} & s_{x x y y} & s_{x x x x} & s_{x x z z} & 0 & 0 \\
\alpha_{z z} & 0 & 0 & d_{z z z} & 0 & 0 & Q_{z z z} & s_{x x z z} & s_{x x z z} & s_{z z z z} & 0 & 0 \\
0 & d_{x y z} & -d_{x x z} & 0 & Q_{x y z} & -Q_{x x z} & 0 & 0 & 0 & 0 & s_{x z x z} & -s_{x z y z} \\
0 & d_{x x z} & d_{x y z} & 0 & Q_{x x z} & Q_{x y z} & 0 & 0 & 0 & 0 & s_{x z y z} & s_{x z x z} \\
0 & 0 & 0 & 0 & 0 & 0 & 0 & s_{x x x y} & -s_{x x x y} & 0 & 0 & 0 \\
s_{x y x y}
\end{array}\right)\left(\begin{array}{l}
\Delta T \\
E_{x} \\
E_{y} \\
E_{z} \\
H_{x} \\
H_{y} \\
H_{z} \\
\sigma_{x x} \\
\sigma_{y y} \\
\sigma_{z z} \\
\sigma_{y z} \\
\sigma_{x z} \\
\sigma_{x y}
\end{array}\right)
$$

## Point group 4

## Courtesy of Prof. M. De Graef

$$
\left(\begin{array}{c}
\Delta S \\
D_{x} \\
D_{y} \\
D_{z} \\
B_{x} \\
B_{y} \\
B_{z} \\
\epsilon_{x x} \\
\epsilon_{y y} \\
\epsilon_{z z} \\
\epsilon_{y z} \\
\epsilon_{x z} \\
\epsilon_{x y}
\end{array}\right)=\left(\begin{array}{ccccccccccccc}
\frac{C}{T} & 0 & 0 & 0 & 0 & 0 & 0 & \alpha & \alpha & \alpha & 0 & 0 & 0 \\
0 & \kappa & 0 & 0 & \lambda & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 \\
0 & 0 & \kappa & 0 & 0 & \lambda & 0 & 0 & 0 & 0 & 0 & 0 & 0 \\
0 & 0 & 0 & \kappa & 0 & 0 & \lambda & 0 & 0 & 0 & 0 & 0 & 0 \\
0 & \lambda & 0 & 0 & \mu & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 \\
0 & 0 & \lambda & 0 & 0 & \mu & 0 & 0 & 0 & 0 & 0 & 0 & 0 \\
0 & 0 & 0 & \lambda & 0 & 0 & \mu & 0 & 0 & 0 & 0 & 0 & 0 \\
\alpha & 0 & 0 & 0 & 0 & 0 & 0 & s_{x x x x} & s_{x x y y} & s_{x x y y} & 0 & 0 & 0 \\
\alpha & 0 & 0 & 0 & 0 & 0 & 0 & s_{x x y y} & s_{x x x x} & s_{x x y y} & 0 & 0 & 0 \\
\alpha & 0 & 0 & 0 & 0 & 0 & 0 & s_{x x y y} & s_{x x y y} & s_{x x x x} & 0 & 0 & 0 \\
0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & s_{y z y z} & 0 & 0 \\
0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & s_{y z y z} & 0 \\
0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & s_{y z y z}
\end{array}\right)\left(\begin{array}{c}
\Delta T \\
E_{x} \\
E_{y} \\
E_{z} \\
H_{x} \\
H_{y} \\
H_{z} \\
\sigma_{x x} \\
\sigma_{y y} \\
\sigma_{z z} \\
\sigma_{y z} \\
\sigma_{x z} \\
\sigma_{x y}
\end{array}\right)
$$

Point group $\mathrm{m} \overline{3} \mathrm{~m}$
Note how many fewer independent coefficients there are! Note how the center of symmetry eliminates many of the properties, such as pyroelectricity

Courtesy of Prof. M. De Graef

## Cubic crystals: anisotropy factor

- If one applies the symmetry elements of the cubic system, it turns out that only three independent coefficients remain: $C_{11}, C_{12}$ and $C_{44,}$ (similar set for compliance). From these three, a useful combination of the first two is

$$
C^{\prime}=\left(C_{11}-C_{12}\right) / 2
$$

- See Nye, Physical Properties of Crystals


## Zener's anisotropy factor

- $C^{\prime}=\left(C_{11}-C_{12}\right) / 2$ turns out to be the stiffness associated with a shear in a <110> direction on a plane. In certain martensitic transformations, this modulus can approach zero which corresponds to a structural instability.
- Zener (Physics, Carnegie Tech. Inst.) proposed a measure of elastic anisotropy based on the ratio $C_{44} / C^{\prime}$. This turns out to be a useful criterion for identifying materials that are elastically anisotropic, i.e., via the extent to which $C_{44} / C^{\prime}$ varies from unity.
- Note that this provides a way to convert an anisotropic elastic stiffness into an isotropic one. One can, e.g., adjust $\mathrm{C}_{12}$ until the Zener ratio=1. Some care is required, however, because one might want to match some average Young's modulus, for example.


## Rotated compliance (matrix)

- Given an orientation $a_{i j}$, we transform the compliance tensor, using cubic point group symmetry, and find that:

$$
\begin{aligned}
S_{11}^{\prime}= & S_{11}\left(a_{11}^{4}+a_{12}^{4}+a_{13}^{4}\right) \\
& +2 S_{12}\left(a_{12}^{2} a_{13}^{2}+a_{11}^{2} a_{12}^{2}+a_{11}^{2} a_{13}^{2}\right) \\
& +S_{44}\left(a_{12}^{2} a_{13}^{2}+a_{11}^{2} a_{12}^{2}+a_{11}^{2} a_{13}^{2}\right)
\end{aligned}
$$

## Rotated compliance (matrix)

- This can be further simplified with the aid of the standard relations between the direction cosines, $a_{\mathrm{ik}} a_{\mathrm{jk}}=1$ for $i=j ; a_{\mathrm{ik}} a_{\mathrm{jk}}$ $=0$ for $i \neq j,\left(a_{\mathrm{ik}} a_{\mathrm{jk}}=\delta_{\mathrm{ij}}\right)$ to read as follows.

$$
\begin{aligned}
& S_{11}^{\prime}=s_{11}- \\
& 2\left(s_{11}-s_{12}-\frac{s_{44}}{2}\right)\left\{\alpha_{1}^{2} \alpha_{2}^{2}+\alpha_{2}^{2} \alpha_{3}^{2}+\alpha_{3}^{2} \alpha_{1}^{2}\right\}
\end{aligned}
$$

- By definition, the Young's modulus in any direction is given by the reciprocal of the compliance, $E=1 / S_{11}^{\prime}$.
- By definition, the Young's modulus along <100> is given by the reciprocal of the compliance for $\langle 100>$ :
$E_{100}=1 / S_{11}=\left\{\left(C_{11}-C_{12}\right)\left(C_{11}+2 C_{12}\right)\right\} /\left(C_{11}+C_{12}\right)$.


## Anisotropy in cubic materials

- Thus the second term on the RHS is zero for $\langle 100\rangle$ directions and, for $C_{44} / C^{\prime \prime}>1$, a maximum in $<111>$ directions (conversely a minimum for $C_{44} / C^{\prime}<1$ ).
The following table shows that most cubic metals have positive values of Zener's coefficient so that $<100>$ is soft and <111> is hard, with the exceptions of $V$ and NaCl .
- See the supplemental slides for how to go between $C$ values and the Lamé constants used to describe isotropic materials.

| Material | $C_{44} / C^{\prime}$ | $\mathrm{E}_{111} / \mathrm{E}_{100}$ |
| :---: | :---: | :---: |
| Cu | 3.21 | 2.87 |
| Ni | 2.45 | 2.18 |
| A 1 | 1.22 | 1.19 |
| Fe | 2.41 | 2.15 |
| Ta | 1.57 | 1.50 |
| $\mathrm{~W}(2000 \mathrm{~K})$ | 1.23 | 1.35 |
| $\mathrm{~W}(\mathrm{R} . \mathrm{T})$. | 1.01 | 1.01 |
| V | 0.78 | 0.72 |
| Nb | 0.55 | 0.57 |
| $\beta-\mathrm{CuZn}$ | 18.68 | 8.21 |
| spinel | 2.43 | 2.13 |
| MgO | 1.49 | 1.37 |
| NaC 1 | 0.69 | 0.74 |

## Stiffness coefficients, cubics

| Material class | Material | $\boldsymbol{C}_{\mathbf{1}}$ <br> $\left(\mathbf{1 0}^{\mathbf{1 0}} \mathbf{N} / \mathbf{m}^{\mathbf{2}}\right)$ | $\boldsymbol{C}_{\mathbf{1 2}}$ <br> $\left(\mathbf{1 0}^{\mathbf{1 0}} \mathbf{N} / \mathbf{m}^{\mathbf{2}}\right)$ | $\boldsymbol{C}_{\mathbf{4 4}}$ <br> $\left(\mathbf{1 0}^{\mathbf{0}} \mathbf{N} / \mathbf{m}^{\mathbf{2}}\right)$ | Anisotropy ratio <br> $\left(\boldsymbol{C}_{\mathbf{1 1}}-\boldsymbol{C}_{\mathbf{1 2}}\right) / \mathbf{2} \boldsymbol{C}_{\mathbf{4 4}}$ |
| :--- | :--- | :---: | :---: | :---: | :---: |
| Metals | Ag | 12.4 | 9.3 | 4.6 | 0.34 |
|  | Al | 10.8 | 6.1 | 2.9 | 0.81 |
|  | Au | 18.6 | 15.7 | 4.2 | 0.35 |
|  | Cu | 16.8 | 12.1 | 7.5 | 0.31 |
|  | $\alpha-\mathrm{Fe}$ | 23.7 | 14.1 | 11.6 | 0.41 |
|  | Mo | 46.0 | 17.6 | 11.0 | 1.29 |
|  | Na | 0.73 | 0.63 | 0.42 | 0.12 |
|  | Ni | 24.7 | 14.7 | 12.5 | 0.40 |
|  | Pb | 5.0 | 4.2 | 1.5 | 0.27 |
| Covalent | W | 50.1 | 19.8 | 15.1 | 1.00 |
| solids | Si | 16.6 | 6.4 | 8.0 | 0.64 |
|  | Diamond | 107.6 | 12.5 | 57.6 | 0.83 |
|  | TiC | 51.2 | 11.0 | 17.7 | 1.14 |
| Ionic solids | LiF | 11.2 | 4.6 | 6.3 | 0.52 |
|  | MgO | 29.1 | 9.0 | 15.5 | 0.65 |
|  | NaCl | 4.9 | 1.3 | 1.3 | 1.38 |

Table 2.2
Stiffness coefficients for selected cubic materials
[Courtney]
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## Anisotropy in terms of moduli

- Another way to write the above equation is to insert the values for the Young's modulus in the soft and hard directions, assuming that the $<100>$ are the most compliant direction(s). (Courtney uses $\alpha, \beta$, and $\gamma$ in place of my $\alpha_{1}, \alpha_{2}$, and $\alpha_{3}$.) The advantage of this formula is that moduli in specific directions can be used directly.

$$
\frac{1}{E_{u v w}}=\frac{1}{E_{100}}-3\left\{\frac{1}{E_{100}}-\frac{1}{E_{111}}\right\}\left(\alpha_{1}^{2} \alpha_{2}^{2}+\alpha_{2}^{2} \alpha_{3}^{2}+\alpha_{3}^{2} \alpha_{1}^{2}\right)
$$

## Example Problem

2.11 a Sketch a (001) plane in a face-centered cubic material and an arbitrary vector within it making an angle $\theta$ with the [100] direction. Plot the Young's modulus for copper as a function of $\theta$ for directions between [110] and [100].
b Sketch a (110) plane in Cu and a vector in the plane making an angle $\alpha$ with the [110] direction. Plot E vs. $\alpha$ for directions between [110] and [001].

Solution: (a) The plane is illustrated to the right Use Eq. (3.22).
$1 / E_{\{n k \mid]}=1 / E_{<100\rangle}-3\left\{1 / E_{<100\rangle}-1 / E_{<111>}\right\}\left(\alpha^{2} \beta^{2}+\alpha^{2} \gamma^{2}+\beta^{2} \gamma^{2}\right)$
where $\alpha$ is the cosine of the angle between the direction [100] and [hkl], $\beta$ is the cosine of the angle between [010] and [hkl], and $\gamma$ is the like cosine between [001] and [hkl]. From the sketches provided we see that $\alpha=\cos \theta, \beta=\cos \left(90^{\circ}-\theta\right)$ $=\sin \theta$ and $\gamma=0$. Employing moduli in units of $10^{10} \mathrm{~N} / \mathrm{m}^{2}$, with $\mathrm{E}_{<100\rangle}=6.7, \mathrm{E}_{<111>}=11.2$ the above equation becomes
$1 / E_{[n k]]}=0.149-0.2915 \cos ^{2} \theta \sin ^{2} \theta$
The table below presents results obtained with the
 above formula; the figure to the right graphs these results.

| $\theta\left({ }^{\circ}\right)$ | $\cos ^{2} \theta \sin ^{2} \theta$ | $E\left(10^{10} \mathrm{~N} / \mathrm{m}^{2}\right)$ |
| :--- | :--- | :--- |
| 0 | 0 | 6.7 |
| 5 | 0.0075 | 6.81 |
| 10 | 0.0292 | 7.12 |
| 15 | 0.0625 | 7.65 |
| 20 | 0.1033 | 8.41 |
| 25 | 0.1467 | 9.41 |
| 30 | 0.1875 | 10.6 |
| 35 | 0.2207 | 11.82 |
| 40 | 0.2425 | 12.77 |
| 45 | $(=[110])$ | 0.25 |



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## Transformations of Stress \& Strain Vectors

- It is useful to be able to transform the axes of stress tensors when written in Voigt notation or matrix/vector form (equation on the left). The table (right) is taken from Newnham's book; the "a" coefficients are from the standard transformation matrix. In vector-matrix form,

$$
\begin{aligned}
& \sigma_{i}^{\prime}=\alpha_{i j} \sigma_{j} \\
& \sigma_{i}=\alpha_{i j}^{-1} \sigma_{j}^{\prime} \\
& \varepsilon_{i}^{\prime}=\alpha_{i j}^{-1 T} \varepsilon_{j} \\
& \varepsilon_{i}=\alpha_{i j}^{T} \varepsilon_{j}^{\prime}
\end{aligned}
$$ the transformations are:

Table 10.1 Transformation matrices for stresses and strains written in matrix form

$$
\left(\begin{array}{c}
\sigma_{1}^{\prime} \\
\sigma_{2}^{\prime} \\
\sigma_{3}^{\prime} \\
\sigma_{4}^{\prime} \\
\sigma_{5}^{\prime} \\
\sigma_{6}^{\prime}
\end{array}\right)=\left[\begin{array}{llllll}
\alpha_{11} & \alpha_{12} & \alpha_{13} & \alpha_{14} & \alpha_{15} & \alpha_{16} \\
\alpha_{21} & \alpha_{22} & \alpha_{23} & \alpha_{24} & \alpha_{25} & \alpha_{26} \\
\alpha_{31} & \alpha_{32} & \alpha_{33} & \alpha_{34} & \alpha_{35} & \alpha_{36} \\
\alpha_{41} & \alpha_{42} & \alpha_{43} & \alpha_{44} & \alpha_{45} & \alpha_{46} \\
\alpha_{51} & \alpha_{52} & \alpha_{53} & \alpha_{54} & \alpha_{55} & \alpha_{56} \\
\alpha_{62} & \alpha_{63} & \alpha_{64} & \alpha_{65} & \alpha_{66}
\end{array}\right]\left(\begin{array}{c}
\sigma_{1} \\
\sigma_{2} \\
\sigma_{3} \\
\sigma_{4} \\
\sigma_{5} \\
\sigma_{6}
\end{array}\right)
$$

| $(\alpha) \quad\left(\begin{array}{cccccc}\left(a_{11}^{2}\right) & \left(a_{12}^{2}\right) & \left(a_{13}^{2}\right) & \left(2 a_{12} a_{13}\right) & \left(2 a_{13} a_{11}\right) & \left(2 a_{11} a_{12}\right) \\ \left(a_{21}^{2}\right) & \left(a_{22}^{2}\right) & \left(a_{23}^{2}\right) & \left(2 a_{22} a_{23}\right) & \left(2 a_{23} a_{21}\right) & \left(2 a_{21} a_{22}\right) \\ \left(a_{31}^{2}\right) & \left(a_{32}^{2}\right) & \left(a_{33}^{2}\right) & \left(2 a_{32} a_{33}\right) & \left(2 a_{33} a_{31}\right) & \left(2 a_{31} a_{32}\right) \\ \left(a_{21} a_{31}\right) & \left(a_{22} a_{32}\right) & \left(a_{23} a_{33}\right) & \left(a_{22} a_{33}+a_{23} a_{32}\right) & \left(a_{21} a_{33}+a_{23} a_{31}\right) & \left(a_{22} a_{31}+a_{21} a_{32}\right) \\ \left(a_{31} a_{11}\right) & \left(a_{32} a_{12}\right) & \left(a_{33} a_{13}\right) & \left(a_{12} a_{33}+a_{13} a_{32}\right) & \left(a_{13} a_{31}+a_{11} a_{33}\right) & \left(a_{11} a_{32}+a_{12} a_{31}\right) \\ \left(a_{11} a_{21}\right) & \left(a_{12} a_{22}\right) & \left(a_{13} a_{23}\right) & \left(a_{12} a_{23}+a_{13} a_{22}\right) & \left(a_{13} a_{21}+a_{11} a_{23}\right) & \left(a_{11} a_{22}+a_{12} a_{21}\right)\end{array}\right)$ |
| :---: |
| $\left(\alpha^{-1}\right) \quad\left(\begin{array}{cccccc}\left(a_{11}^{2}\right) & \left(a_{21}^{2}\right) & \left(a_{31}^{2}\right) & \left(2 a_{21} a_{31}\right) & \left(2 a_{31} a_{11}\right) & \left(2 a_{11} a_{21}\right) \\ \left(a_{12}^{2}\right) & \left(a_{22}^{2}\right) & \left(a_{32}^{2}\right) & \left(2 a_{22} a_{32}\right) & \left(2 a_{32} a_{12}\right) & \left(2 a_{12} a_{22}\right) \\ \left(a_{13}^{2}\right) & \left(a_{23}^{2}\right) & \left(a_{33}^{2}\right) & \left(2 a_{23} a_{33}\right) & \left(2 a_{33} a_{13}\right) & \left(2 a_{13} a_{23}\right) \\ \left(a_{12} a_{13}\right) & \left(a_{22} a_{23}\right) & \left(a_{32} a_{33}\right) & \left(a_{22} a_{33}+a_{32} a_{23}\right) & \left(a_{12} a_{33}+a_{32} a_{13}\right) & \left(a_{22} a_{13}+a_{12} a_{23}\right) \\ \left(a_{13} a_{11}\right) & \left(a_{23} a_{21}\right) & \left(a_{33} a_{31}\right) & \left(a_{21} a_{33}+a_{31} a_{23}\right) & \left(a_{31} a_{13}+a_{11} a_{33}\right) & \left(a_{11} a_{23}+a_{21} a_{13}\right) \\ \left(a_{11} a_{12}\right) & \left(a_{21} a_{22}\right) & \left(a_{31} a_{32}\right) & \left(a_{21} a_{32}+a_{31} a_{22}\right) & \left(a_{31} a_{12}+a_{11} a_{32}\right) & \left(a_{11} a_{22}+a_{21} a_{12}\right)\end{array}\right)$ |

## Alternate Vectorization

$$
\begin{aligned}
& \mathbf{b}^{(1)}=\frac{1}{\sqrt{6}}\left(\begin{array}{ccc}
-1 & 0 & 0 \\
0 & -1 & 0 \\
0 & 0 & 2
\end{array}\right) ; \mathbf{b}^{(2)}=\frac{1}{\sqrt{2}}\left(\begin{array}{ccc}
-1 & 0 & 0 \\
0 & 1 & 0 \\
0 & 0 & 0
\end{array}\right) ; \mathbf{b}^{(3)}=\frac{1}{\sqrt{2}}\left(\begin{array}{lll}
0 & 0 & 0 \\
0 & 0 & 1 \\
0 & 1 & 0
\end{array}\right) \\
& \mathbf{b}^{(4)}=\frac{1}{\sqrt{2}}\left(\begin{array}{lll}
0 & 0 & 1 \\
0 & 0 & 0 \\
1 & 0 & 0
\end{array}\right) ; \quad \mathbf{b}^{(5)}=\frac{1}{\sqrt{2}}\left(\begin{array}{lll}
0 & 1 & 0 \\
1 & 0 & 0 \\
0 & 0 & 0
\end{array}\right) ; \quad \mathbf{b}^{(6)}=\frac{1}{\sqrt{3}}\left(\begin{array}{lll}
1 & 0 & 0 \\
0 & 1 & 0 \\
0 & 0 & 1
\end{array}\right)
\end{aligned}
$$

An alternate vectorization, discussed by Tomé on p287 of the Kocks et al. textbook, is to use the above set of eigentensors. For both stress and strain, one can matrix multiply each eigentensor into the stress/strain tensor in turn and obtain the coefficient of the corresponding stress/strain vector. Work conjugacy is still satisfied. The first two eigentensors represent shears in the $\{110\}$ planes; the next three are simple shears on $\{110\}<110>$ systems, and the last ( $6^{\text {th }}$ ) is the hydrostatic component. The same vectorization can be used for plastic anisotropy, except in this case, the sixth, hydrostatic component is (generally) ignored.

## Homogeneity

- Stimuli and responses of interest are, in general, not scalar quantities but tensors. Furthermore, some of the properties of interest, such as the plastic properties of a material, are far from linear at the scale of a polycrystal. Nonetheless, they can be treated as linear at a suitably local scale and then an averaging technique can be used to obtain the response of the polycrystal. The local or microscopic response is generally well understood but the validity of the averaging techniques is still controversial in many cases. Also, we will only discuss cases where a homogeneous response can be reasonably expected.
- There are many problems in which a non-homogeneous response to a homogeneous stimulus is of critical importance. Stress-corrosion cracking, for example, is a wildly non-linear, non-homogeneous response to an approximately uniform stimulus which depends on the mechanical and electro-chemical properties of the material.


## Homogeneity

- How do we go from a single crystal to a poly-crystal ?
- How do we average properties over a microstructure? What are the relevant parameters? Orientation, grain size, grain boundaries, ...

This is a very complex problem!<br>How do we "solve this"?

## Homogeneity

- Take the effective properties of a microstructure in the form of bounds
- Bounds are typically estimates of the upper and lower limiting values of a given property for a given microstructure.
- Many ways to derive such bounds at various levels of complexity:
- Simple: (i.e. Voigt-Reuss Bounds)

- More complex: Hashin-Shtrickman bounds; Self-consistent methods (i.e. elastic-plastic polycrystals VPSC/EPSC)


## Summary

- We have covered the following topics:
- Linear properties
- Non-linear properties
- Examples of properties
- Tensors, vectors, scalars, tensor transformation law.
- Elasticity, as example as of higher order property, also as example as how to apply (crystal) symmetry.


## Supplemental Slides

- The following slides contain some useful material for those who are not familiar with all the detailed mathematical methods of matrices, transformation of axes, tensors etc.


## Matrix Multiplication

- Take each row of the LH matrix in turn and multiply it into each column of the RH matrix.
- In suffix notation, $a_{\mathrm{ij}}=b_{\mathrm{ik}} c_{\mathrm{kj}}$

$$
\begin{aligned}
& {\left[\begin{array}{lll}
a \alpha+b \delta+c \gamma & a \beta+b \varepsilon+c \mu & a \gamma+b \phi+c v \\
d \alpha+e \delta+f \gamma & d \beta+e \varepsilon+f \mu & d \gamma+e \phi+f v \\
l \alpha+m \delta+n \gamma & l \beta+m \varepsilon+n \mu & l \gamma+m \phi+n v
\end{array}\right]} \\
& =\left[\begin{array}{ccc}
a & b & c \\
d & e & f \\
l & m & n
\end{array}\right] \times\left[\begin{array}{lll}
\alpha & \beta & \gamma \\
\delta & \varepsilon & \phi \\
\lambda & \mu & v
\end{array}\right]
\end{aligned}
$$

## Properties of Rotation Matrix

- The rotation matrix is an orthogonal matrix, meaning that any row is orthogonal to any other row (the dot products are zero). In physical terms, each row represents a unit vector that is the position of the corresponding (new) old axis in terms of the (old) new axes.
- The same applies to columns: in suffix notation -

$$
a_{\mathrm{ij}} a_{\mathrm{kj}}=\delta_{\mathrm{ik}}, a_{\mathrm{ji}} a_{\mathrm{jk}}=\delta_{\mathrm{ik}}
$$



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## Direction Cosines,

## contd.

- That the set of direction cosines are not independent is evident from the following construction:
$\hat{e}_{i}^{\prime} \cdot \hat{e}_{j}^{\prime}=a_{i k} a_{j l} \hat{e}_{k} \cdot \hat{e}_{l}=a_{i k} a_{j l} \delta_{k l}=a_{i k} a_{j k}=\delta_{i j}$
Thus, there are six relationships (itakes values from 1 to 3 , and $j$ takes values from 1 to 3) between the nine direction cosines, and therefore, as stated above, only three are independent, exactly as expected for a rotation.
- Another way to look at a rotation: combine an axis (described by a unit vector with two parameters) and a rotation angle (one more parameter, for a total of 3).


## Orthogonal Matrices

- Note that the direction cosines can be arranged into a $3 \times 3$ matrix, $\Lambda$, and therefore the relation above is equivalent to the expression

$$
\Lambda \Lambda^{T}=\mathrm{I}
$$

where $\Lambda^{\top}$ denotes the transpose of $\Lambda$. This relationship identifies $\Lambda$ as an orthogonal matrix, which has the properties

$$
\Lambda^{-1}=\Lambda^{T} \quad \operatorname{det} \Lambda= \pm 1
$$

## Relationships

- When both coordinate systems are right-handed, $\operatorname{det}(\Lambda)=+1$ and $\Lambda$ is a proper orthogonal matrix. The orthogonality of $\Lambda$ also insures that, in addition to the relation above, the following holds:

$$
\hat{e}_{j}=a_{i j} \hat{e}_{i}^{\prime}
$$

Combining these relations leads to the following interrelationships between components of vectors in the two coordinate systems:

$$
v_{i}=a_{j i} v_{j}^{\prime}, v_{j}^{\prime}=a_{j i} v_{i}
$$

## Transformation Law

- These relations are called the laws of transformation for the components of vectors. They are a consequence of, and equivalent to, the parallelogram law for addition of vectors. That such is the case is evident when one considers the scalar product expressed in two coordinate systems:

$$
\begin{array}{r}
\vec{u} \cdot \vec{v}=u_{i} v_{i}=a_{j i} u_{j}^{\prime} a_{k i} v_{k}^{\prime}= \\
\delta_{j k} u_{j}^{\prime} v_{k}^{\prime}=u_{j}^{\prime} v_{j}^{\prime}=u_{i}^{\prime} v_{i}^{\prime}
\end{array}
$$

## Invariants

Thus, the transformation law as expressed preserves the lengths and the angles between vectors. Any function of the components of vectors which remains unchanged upon changing the coordinate system is called an invariant of the vectors from which the components are obtained. The derivations illustrate the fact that the scalar product $\vec{u} \cdot \vec{v}$ is an invariant of $\vec{u}$ and $\vec{v}$. Other examples of invariants include the vector product of two vectors and the triple scalar product of three vectors. The reader should note that the transformation law for vectors also applies to the components of points when they are referred to a common origin.

## Orthogonality

- A rotation matrix, $\Lambda$, is an orthogonal matrix, however, because each row is mutually orthogonal to the other two.

$$
a_{k i} a_{k j}=\delta_{i j}, \quad a_{i k} a_{j k}=\delta_{i j}
$$

- Equally, each column is orthogonal to the other two, which is apparent from the fact that each row/column contains the direction cosines of the new/old axes in terms of the old/new axes and we are working with [mutually perpendicular] Cartesian axes.


## Anisotropy

- Anisotropy as a word simply means that something varies with direction.
- Anisotropy is from the Greek: aniso = different, varying; tropos = direction.
- Almost all crystalline materials are anisotropic; many materials are engineered to take advantage of their anisotropy (beer cans, turbine blades, microchips...)
- Older texts use trigonometric functions to describe anisotropy but tensors offer a general description with which it is much easier to perform calculations.
- For materials, what we know is that some properties are anisotropic. This means that several numbers, or coefficients, are needed to describe the property - one number is not sufficient.
- Elasticity is an important example of a property that, when examined in single crystals, is often highly anisotropic. In fact, the lower the crystal symmetry, the greater the anisotropy is likely to be.
- Nomenclature: in general, we need to use tensors to describe fields and properties. The simplest case of a tensor is a scalar which is all we need for isotropic properties. The next "level" of tensor is a vector, e.g. electric current.


## Scalars, Vectors, Tensors

- Scalar:= quantity that requires only one number, e.g. density, mass, specific heat. Equivalent to a zero-rank tensor.
- Vector:= quantity that has direction as well as magnitude, e.g. velocity, current, magnetization; requires 3 numbers or coefficients (in 3D). Equivalent to a first-rank tensor.
- 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.


## Vector field, response

- If we have a vector response, $R$, that we can write in component form, a vector field, $F$, that we can also write in component form, and a property, $P$, that we can write in matrix form (with nine coefficients) then the linearity of the property means that we can write the following ( $R_{0}=0$ ):

$$
R_{i}=P_{\mathrm{ij}} F_{j}
$$

- A scalar (e.g. pressure) can be called a zero-rank tensor.
- A vector (e.g. electric current) is also known as a first-rank tensor.


## Linear anisotropic property

- This means that each component of the response is linearly related to each component of the field and that the proportionality constant is the appropriate coefficient in the matrix. Example:

$$
R_{1}=P_{13} F_{3},
$$

which says that the first component of the response is linearly related to the third field component through the property coefficient $P_{13}$.


## Example: electrical conductivity

- An example of such a linear anisotropic (second order tensor, discussed in later slides) property is the electrical conductivity of a material:
- Field: Electric Field, $E$
- Response: Current Density,J
- Property: Conductivity, $\sigma$
- $J_{i}=\sigma_{i j} E_{j}$


## Anisotropic electrical conductivity

- We can illustrate anisotropy with Nye's example of electrical conductivity, $\sigma$.


b

Fig. 1.1. The relation between the electric current density $\mathbf{j}$ and the electric field $\mathbf{E}$ in ( $a$ ) an isotropic conductor and (b) an anisotropic conductor.


Fig. 1.2. The components of current density when a field is applied along $O x_{1}$.

Stimulus/ Field: $E_{1} \neq 0, E_{2}=E_{3}=0$ Response: $j_{1}=\sigma_{11} E_{1}, j_{2}=\sigma_{21} E_{1}, j_{3}=\sigma_{31} E_{1}$,

## 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 nonsymmetric 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/pc/wintterlin/IMGs/pt10p3.jpg
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## New Axes

- Consider a new orthonormal system consisting of righthanded base vectors: $\hat{e}_{1}^{\prime}, \hat{e}_{2}^{\prime}$ and $\hat{e}_{3}^{\prime}$ These all have the same origin, $o$, associated with $\hat{e}_{1}^{\prime}, \hat{e}_{2}^{\prime}$ and $\hat{e}_{3}^{\prime}$
- The vector $v$ is clearly expressed equally well in either coordinate system:

$$
\vec{v}=v_{i} \hat{e}_{i}=v_{i}^{\prime} \hat{e}_{i}^{\prime}
$$

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.


## Anisotropy in Composites

- The same methods developed here for describing the anisotropy of single crystals can be applied to composites.
- Anisotropy is important in composites, not because of the intrinsic properties of the components but because of the arrangement of the components.
- As an example, consider (a) a uniaxial composite (e.g. tennis racket handle) and (b) a flat panel cross-ply composite (e.g. wing surface).


## Fiber Symmetry



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## Fiber Symmetry

- We will use the same matrix notation for stress, strain, stiffness and compliance as for single crystals.
- The compliance matrix, s, has 5 independent coefficients.

$$
\left[\begin{array}{cccccc}
s_{11} & s_{12} & s_{13} & 0 & 0 & 0 \\
s_{12} & s_{11} & s_{13} & 0 & 0 & 0 \\
s_{13} & s_{13} & s_{33} & 0 & 0 & 0 \\
0 & 0 & 0 & s_{44} & 0 & 0 \\
0 & 0 & 0 & 0 & s_{44} & 0 \\
0 & 0 & 0 & 0 & 0 & 2\left(s_{11}-s_{12}\right)
\end{array}\right]
$$

## Relationships

- For a uniaxial stress along the $z(3)$ direction,

$$
E_{3}=\frac{\sigma_{3}}{\varepsilon_{3}}=\frac{1}{s_{33}}\left(=\frac{\sigma_{z z}}{\varepsilon_{z z}}\right)
$$

- This stress causes strain in the transverse plane: $e_{11}=e_{22}=s_{12} \sigma_{33}$. Therefore we can calculate Poisson's ratio as:

$$
v_{13}=\frac{e_{1}}{e_{3}}=\frac{s_{13}}{s_{33}}\left(=\frac{e_{x x}}{e_{z z}}\right)
$$

- Similarly, stresses applied perpendicular to z give rise to different moduli and Poisson's ratios.

$$
E_{1}=\frac{\sigma_{1}}{\varepsilon_{1}}=\frac{1}{s_{11}}, \quad v_{21}=\frac{-s_{12}}{s_{11}}, \quad v_{31}=\frac{-s_{13}}{s_{11}}
$$

## Relationships, contd.

- Similarly the torsional modulus is related to shears involving the $z$ axis, i.e. $y z$ or $x z$ shears:

$$
s_{44}=s_{55}=1 / G
$$

- Shear in the $x-y$ plane (1-2 plane) is related to the other compliance coefficients:

$$
s_{66}=2\left(s_{11}-s_{12}\right)=1 / G_{x y}
$$

## Plates: Orthotropic Symmetry

- Again, we use the same matrix notation for stress, strain, stiffness and compliance as for single crystals.
- The compliance matrix, s, has 9 independent coefficients.
- This corresponds to othorhombic sample symmetry: see the following slide with Table from Nye's book.
$\left[\begin{array}{cccccc}s_{11} & s_{12} & s_{13} & 0 & 0 & 0 \\ s_{12} & s_{22} & s_{23} & 0 & 0 & 0 \\ s_{13} & s_{23} & s_{33} & 0 & 0 & 0 \\ 0 & 0 & 0 & s_{44} & 0 & 0 \\ 0 & 0 & 0 & 0 & s_{55} & 0 \\ 0 & 0 & 0 & 0 & 0 & s_{66}\end{array}\right]$


## Plates: $0^{\circ}$ and $90^{\circ}$ plies

- If the composite is a laminate composite with fibers laid in at $0^{\circ}$ and $90^{\circ}$ in equal thicknesses then the symmetry is higher because the $x$ and $y$ directions are equivalent.
- The compliance matrix, $\mathbf{s}$, has 6 independent coefficients.
- This corresponds to (tetragonal) $4 m m$ sample symmetry: see the following slide with Table from Nye's book.
$\left[\begin{array}{cccccc}s_{11} & s_{12} & s_{13} & 0 & 0 & 0 \\ s_{12} & s_{11} & s_{13} & 0 & 0 & 0 \\ s_{13} & s_{13} & s_{33} & 0 & 0 & 0 \\ 0 & 0 & 0 & s_{44} & 0 & 0 \\ 0 & 0 & 0 & 0 & s_{44} & 0 \\ 0 & 0 & 0 & 0 & 0 & s_{66}\end{array}\right]$

Table 9
Form of the $\left(s_{i j}\right)$ and $\left(c_{i j}\right)$ matrices


| Triclinic <br> Both classes |  |
| :---: | :---: |
|  | NIC |
| Orthorhombic <br> All classes |  |

## Effect of Symmetry on the Elasticity Tensors, S, C



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## Lamé constants (isotropic elasticity)

For an elastically isotropic body, there are only 2 elastic moduli, known as the Lamé constants.

$$
\left(\begin{array}{c}
\sigma_{11} \\
\sigma_{22} \\
\sigma_{33} \\
\sigma_{23} \\
\sigma_{13} \\
\sigma_{12}
\end{array}\right)=\left(\begin{array}{cccccc}
\lambda+2 G & \lambda & \lambda & 0 & 0 & 0 \\
\lambda & \lambda+2 G & \lambda & 0 & 0 & 0 \\
\lambda & \lambda & \lambda+2 G & 0 & 0 & 0 \\
0 & 0 & 0 & G & 0 & 0 \\
0 & 0 & 0 & 0 & G & 0 \\
0 & 0 & 0 & 0 & 0 & G
\end{array}\right)\left(\begin{array}{c}
\epsilon_{11} \\
\epsilon_{22} \\
\epsilon_{33} \\
2 \epsilon_{23} \\
2 \epsilon_{13} \\
2 \epsilon_{12}
\end{array}\right)
$$

This means that, if you know the Lamé constants, then you can obtain the stiffness values thus:

$$
\begin{array}{r}
C_{11}=\lambda+2 G \\
C_{12}=\lambda \\
C_{44}=G
\end{array}
$$

## Stiffnesses in terms of E and v

$$
\begin{aligned}
C_{11} & =\frac{E}{(1+\nu)}\left(1+\frac{\nu}{1-2 \nu}\right) \\
C_{12} & =\frac{E}{(1+\nu)} \frac{\nu}{(1-2 \nu)} \\
C_{44} & =\frac{E}{2(1+\nu)}
\end{aligned}
$$

$$
\begin{aligned}
& \mathrm{S}_{11}=\left(\mathrm{C}_{11}+\mathrm{C}_{12}\right) /\left\{\left(\mathrm{C}_{11}-\mathrm{C}_{12}\right)\left(\mathrm{C}_{11}+2 \mathrm{C}_{12}\right)\right\} \\
& \mathrm{S}_{11}=(\lambda+2 G+\lambda) /\{(\lambda+2 G-\lambda)(\lambda+2 G+2 \lambda)\} \\
& \mathrm{S}_{11}=(\lambda+G) /\{G(2 G+3 \lambda)\} \\
& \mathrm{S}_{12}=-\mathrm{C}_{12} /\left\{\left(\mathrm{C}_{11}-\mathrm{C}_{12}\right)\left(\mathrm{C}_{11}+2 \mathrm{C}_{12}\right)\right\} \\
& \mathrm{S}_{12}=-\lambda /\{2 G(2 G+3 \lambda)\} \\
& \mathrm{S}_{44}=1 / G \quad \text { Please acknowledge Carnegie Mellon if you make public use of these slides }
\end{aligned}
$$

## Young's, Bulk moduli, Poisson's ratio

1. Young's modulus: the ratio of longitudinal stress to longitudinal strain under uniaxial normal loading in the longitudinal direction.

$$
E=2 G+\frac{G \lambda}{G+\lambda} \quad E=2 G(1+\nu)=>G=\frac{E}{2(1+\nu)}
$$

2. Poisson's ratio : minus the ratio of lateral strain to longitudinal strain.

$$
2 \nu=\frac{\lambda}{\lambda+G} \quad \nu=\frac{\lambda}{2(\lambda+G)} \quad \lambda=\frac{\nu E}{(1+\nu)(1-2 \nu)}
$$

3. Bulk modulus : minus the ratio of pressure to dilatation.

$$
\begin{aligned}
& K=-\frac{1}{3} \frac{\sigma_{x x}+\sigma_{y y}+\sigma_{z z}}{\epsilon_{x x}+\epsilon_{y y}+\epsilon_{z z}}=\frac{E}{3(1-2 \nu)} \\
& v=\lambda / 2(G+\lambda)=>1-v=\{2(G+\lambda)-\lambda\} / 2(G+\lambda) \\
& E=2\{2 G(G+\lambda)+G \lambda\} / 2(G+\lambda) \\
& 1-v=\{2 G+\lambda)\} / 2(G+\lambda) \\
& E=2\left\{2 G^{2}+3 G \lambda\right\} / 2(G+\lambda) \\
& 1-2 v=\{2(G+\lambda)-2 \lambda\} / 2(G+\lambda)=G /(G+\lambda) \\
& \mathrm{E} / 1-2 v=2 G\{2 G+3 \lambda\} / 2 G=2 G+3 \lambda \\
& 1-(1-2 v) /(1-2 v)=\lambda / G \\
& \mathrm{E} / 1-2 v=2 G\{2 G+3 \lambda\} / 2 G=2 G+3 G\{2 v /(1-2 v)\} \\
& 2 v /(1-2 v)=\lambda / G \\
& \mathrm{E} / 1-2 v=2 G\{1+v\} /(1-2 v) \\
& G 2 v /(1-2 v)=\lambda \\
& \mathrm{E}=2 G\{1+v\}=>G=E / 2\{1+v\} \\
& \lambda=G 2 v /(1-2 v) \\
& \lambda=E / 2\{1+v\}^{*} 2 v /(1-2 v) \\
& \mathrm{C}_{11}=E /\{1+v\}+v E /\{(1+v)(1-2 v)\} \\
& \lambda=v E /\{(1+v)(1-2 v)\} \\
& \mathrm{C}_{11}=E[1+(v /(1-2 v))] /\{1+v\} \\
& \lambda=E /(1+v)\{v /(1-2 v)\}
\end{aligned}
$$

## General Anisotropic Properties

- Many different properties of crystals can be described as tensors.
- The rank of each tensor property depends, naturally, on the nature of the quantities related by the property.


## Examples of Materials Properties as Tensors

- Table 1 shows a series of tensors that are of importance for material science. The tensors are grouped by rank, and are also labeled (in the last column) by $E$ (equilibrium property) or $T$ (transport property). The number following this letter indicates the maximum number of independent, nonzero elements in the tensor, taking into account symmetries imposed by thermodynamics.
- The Field and Response columns contain the following symbols: $\Delta T=$ temperature difference, $\Delta S=$ entropy change, $E_{i}=$ electric field components, $H_{i}=$ magnetic field components, $\varepsilon_{i \mathrm{ij}}=$ mechanical strain, $D_{i}=$ electric displacement, $B_{i}=$ magnetic induction, $\sigma_{\mathrm{ij}}=$ mechanical stress, $\Delta \beta_{\mathrm{ij}}=$ change of the impermeability tensor, $j_{\mathrm{i}}=$ electrical current density, $\nabla_{\mathrm{j}} T=$ temperature gradient, $h_{\mathrm{i}}=$ heat flux, $\nabla_{\mathrm{j}} c=$ concentration gradient, $m_{\mathrm{i}}=$ mass flux, $\rho_{\mathrm{i}}^{\mathrm{a}}=$ anti-symmetric part of resistivity tensor, $\rho_{\mathrm{i}}^{\mathrm{s}}=$ symmetric part of resistivity tensor, $\Delta \rho_{\mathrm{ij}}=$ change in the component $i j$ of the resistivity tensor, $I_{i}=$ direction cosines of wave direction in crystal, $G=$ gyration constant,

| Property | Symbol | Field | Response | Type\# |
| :---: | :---: | :---: | :---: | :---: |
| Tensors of Rank 0 (Scalars) |  |  |  |  |
| Specific Heat | C | $\Delta T$ | $T \Delta S$ | E1 |
| Tensors of Rank 1 (Vectors) |  |  |  |  |
| Electrocaloric | $p_{i}$ | $E_{i}$ | $\Delta S$ | E3 |
| Magnetocaloric | $q_{i}$ | $H_{i}$ | $\Delta S$ | E3 |
| Pyroelectric | $p_{i}^{\prime}$ | $\Delta T$ | $D_{i}$ | E3 |
| Pyromagnetic | $q_{i}^{\prime}$ | $\Delta T$ | $B_{i}$ | E3 |
| Tensors of Rank 2 |  |  |  |  |
| Thermal expansion | $\alpha_{i j}$ | $\Delta T$ | $\epsilon_{i j}$ | E6 |
| Piezocaloric effect | $\alpha_{i j}^{\prime}$ | $\sigma_{i j}$ | $\Delta S$ | E6 |
| Dielectric permittivity | $\kappa_{i j}$ | $E_{j}$ | $D_{i}$ | E6 |
| Magnetic permeability | $\mu_{i j}$ | $H_{j}$ | $B_{i}$ | E6 |
| Optical activity | $g_{i j}$ | $l_{i} l_{j}$ | $G$ | E6 |
| Magnetoelectric polarization | $\lambda_{i j}$ | $H_{j}$ | $D_{i}$ | E9 |
| Converse magnetoelectric polarization | $\lambda_{i j}^{\prime}$ | $E_{j}$ | $B_{i}$ | E9 |
| Electrical conductivity (resistivity) | $\sigma_{i j}\left(\rho_{i j}\right)$ | $E_{j}\left(j_{j}\right)$ | $j_{i}\left(E_{i}\right)$ | T6 |
| Thermal conductivity | $K_{i j}$ | $\nabla_{j} T$ | $h_{i}$ | T6 |
| Diffusivity | $D_{i j}$ | $\nabla_{j} c$ | $m_{i}$ | T6 |
| Thermoelectric power | $\Sigma_{i j}$ | $\nabla_{j} T$ | $E_{i}$ | T9 |
| Hall effect Pleaseacknowte | $\xrightarrow{R_{i j}}$ | ${ }^{B_{j}}$ | $\begin{gathered} \rho_{i}^{a} \\ \text { esestides } \end{gathered}$ | T9 |


| Tensors of Rank 3 |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: |
| Piezoelectricity | $d_{i j k}$ | $\sigma_{j k}$ | $D_{i}$ | E18 |
| Converse piezoelectricity | $d_{i j k}^{\prime}$ | $E_{k}$ | $\epsilon_{i j}$ | E18 |
| Piezomagnetism | $Q_{i j k}$ | $\sigma_{j k}$ | $B_{i}$ | E18 |
| Converse piezomagnetism | $Q_{i j k}^{\prime}$ | $H_{k}$ | $\epsilon_{i j}$ | E18 |
| Electro-optic effect | $r_{i j k}$ | $E_{k}$ | $\Delta \beta_{i j}$ | E18 |
| Nernst tensor | $\Sigma_{i j k}$ | $\nabla_{j} T B_{k}$ | $E_{i}$ | T27 |
| Tensors of Rank 4 |  |  |  |  |
| Elasticity | $s_{i j k l}\left(c_{i j k l}\right)$ | $\sigma_{k l}\left(\epsilon_{k l}\right)$ | $\epsilon_{i j}\left(\sigma_{i j}\right)$ | E21 |
| Electrostriction | $\gamma_{i j k l}$ | $E_{k} E_{l}$ | $\epsilon_{i j}$ | E36 |
| Photoelasticity | $q_{i j k l}$ | $\sigma_{k l}$ | $\Delta \beta_{i j}$ | E36 |
| Kerr effect | $p_{i j k l}$ | $E_{k} E_{l}$ | $\Delta \beta_{i j}$ | E36 |
| Magnetoresistance | $\xi_{i j k l}$ | $B_{k} B_{l}$ | $\rho_{i j}^{s}$ | T36 |
| Piezoresistance | $\Pi_{i j k l}$ | $\sigma_{k l}$ | $\Delta \rho_{i j}$ | T36 |
| Magnetothermoelectric power | $\Sigma_{i j k l}$ | $\nabla_{j} T B_{k} B_{l}$ | $E_{i}$ | T54 |
| Second order Hall effect | $\rho_{i j k l}$ | $B_{j} B_{k} B_{l}$ | $\rho_{i}^{2}$ | T30 |
| Tensors of Rank 6 |  |  |  |  |
| Third order elasticity | $c_{i j k l m n}$ | $\epsilon_{k l} \epsilon_{m n}$ | $\sigma_{i j}$ | E56 |

## Courtesy of Prof. M. De Graef

$\left(\begin{array}{c}\Delta S \\ D_{x} \\ D_{y} \\ D_{z} \\ B_{x} \\ B_{y} \\ B_{z} \\ \epsilon_{x x} \\ \epsilon_{y y} \\ \epsilon_{z z} \\ \epsilon_{y z} \\ \epsilon_{x z} \\ \epsilon_{x y}\end{array}\right)=\left(\begin{array}{c|ccc|ccc|cccccc}\frac{C}{T} & p_{x} & p_{y} & p_{z} & q_{x} & q_{y} & q_{z} & \alpha_{x x}^{\prime} & \alpha_{y y}^{\prime} & \alpha_{z z}^{\prime} & \alpha_{y z}^{\prime} & \alpha_{x z}^{\prime} & \alpha_{x y}^{\prime} \\ \hline p_{x}^{\prime} & \kappa_{x x} & \kappa_{x y} & \kappa_{x z} & \lambda_{x x} & \lambda_{x y} & \lambda_{x z} & d_{x x x} & d_{x y y} & d_{x z z} & d_{x y z} & d_{x x z} & d_{x x y} \\ p_{y}^{\prime} & \kappa_{y x} & \kappa_{y y} & \kappa_{y z} & \lambda_{y x} & \lambda_{y y} & \lambda_{y z} & d_{y x x} & d_{y y y} & d_{y z z} & d_{y y z} & d_{y x z} & d_{y x y} \\ p_{z}^{\prime} & \kappa_{z x} & \kappa_{z y} & \kappa_{z z} & \lambda_{z x} & \lambda_{z y} & \lambda_{z z} & d_{z x x} & d_{z y y} & d_{z z z} & d_{z y z} & d_{z x z} & d_{z x y} \\ \hline q_{x}^{\prime} & \lambda_{x x}^{\prime} & \lambda_{x y}^{\prime} & \lambda_{x z}^{\prime} & \mu_{x x} & \mu_{x y} & \mu_{x z} & Q_{x x x} & Q_{x y y} & Q_{x z z} & Q_{x y z} & Q_{x x z} & Q_{x x y} \\ q_{y}^{\prime} & \lambda_{y x}^{\prime} & \lambda_{y y}^{\prime} & \lambda_{y z}^{\prime} & \mu_{y x} & \mu_{y y} & \mu_{y z} & Q_{y x x} & Q_{y y y} & Q_{y z z} & Q_{y y z} & Q_{y x z} & Q_{y x y} \\ q_{z}^{\prime} & \lambda_{z x}^{\prime} & \lambda_{z y}^{\prime} & \lambda_{z z}^{\prime} & \mu_{z x} & \mu_{z y} & \mu_{z z} & Q_{z x x} & Q_{z y y} & Q_{z z z} & Q_{z y z} & Q_{z x z} & Q_{z x y} \\ \hline \alpha_{x x} & d_{x x x}^{\prime} & d_{x x y}^{\prime} & d_{x x z}^{\prime} & Q_{x x x}^{\prime} & Q_{x x y}^{\prime} & Q_{x x z}^{\prime} & s_{x x x x} & s_{x x y y} & s_{x x z z} & s_{x x y z} & s_{x x x z} & s_{x x x y} \\ \alpha_{y y}^{\prime} & d_{y y x}^{\prime \prime} & d_{y y y}^{\prime} & d_{y y z}^{\prime} & Q_{y y x}^{\prime} & Q_{y y y}^{\prime} & Q_{y y z}^{\prime} & s_{y y x x} & s_{y y y y} & s_{y y z z} & s_{y y y z} & s_{y y x z} & s_{y y x y} \\ \alpha_{z z} & d_{z z x}^{\prime} & d_{z z y}^{\prime} & d_{z z z}^{\prime} & Q_{z z x}^{\prime} & Q_{z z y}^{\prime} & Q_{z z z}^{\prime} & s_{z z x x} & s_{z z y y} & s_{z z z z} & s_{z z y z} & s_{z z x z} & s_{z z x y}^{\prime} \\ \alpha_{y z}^{\prime} & d_{y z x}^{\prime \prime} & d_{y z y}^{\prime} & d_{y z z}^{\prime} & Q_{y z x}^{\prime} & Q_{y z y}^{\prime} & Q_{y z z}^{\prime} & s_{y z x x} & s_{y z y y} & s_{y z z z} & s_{y z y z} & s_{y z x z} & s_{y z x y} \\ \alpha_{x z} & d_{x z x}^{\prime} & d_{x z y}^{\prime} & d_{x z z}^{\prime} & Q_{x z x}^{\prime} & Q_{x z y}^{\prime} & Q_{x z z}^{\prime} & s_{x z x x} & s_{x z y y} & s_{x z z z} & s_{x z y z} & s_{x z x z} & s_{x z x y} \\ \alpha_{x y}^{\prime} & d_{x y x}^{\prime \prime} & d_{x y y}^{\prime} & d_{x y z}^{\prime} & Q_{x y x}^{\prime} & Q_{x y y}^{\prime} & Q_{x y z}^{\prime} & s_{x y x x} & s_{x y y y} & s_{x y z z} & s_{x y y z} & s_{x y x z} & s_{x y x y}\end{array}\right)\left(\begin{array}{l}\Delta T \\ E_{x} \\ E_{y} \\ E_{z} \\ H_{x} \\ H_{y} \\ H_{z} \\ \sigma_{x x} \\ \sigma_{y y} \\ \sigma_{z z} \\ \sigma_{y z} \\ \sigma_{x z} \\ \sigma_{x y}\end{array}\right)$

## Principal Effects

Courtesy of Prof. M. De Graef
Electrocaloric = pyroelectric

- Magnetocaloric = pyromagnetic

Thermal expansion = piezocaloric
Magnetoelectric and converse magnetoelectric
Piezoelectric and converse piezoelectric
Piezomagnetic and converse piezomagnetic
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$$
\begin{aligned}
\Delta S & =\frac{C}{T} \Delta T+p_{i} E_{i}+q_{i} H_{i}+\alpha_{i j}^{\prime} \sigma_{i j} ; \\
D_{i} & =p_{i}^{\prime} \Delta T+\kappa_{i j} E_{j}+\lambda_{i j} H_{j}+d_{i j k} \sigma_{j k} ; \\
B_{i} & =q_{i}^{\prime} \Delta T+\lambda_{i j} E_{j}+\mu_{i j} H_{j}+Q_{i j k} \sigma_{j k} ; \\
\epsilon_{i j} & =\alpha_{i j} \Delta T+d_{i j k}^{\prime} E_{k}+Q_{i j k}^{\prime} H_{k}+s_{i j k l} \sigma_{k l} .
\end{aligned}
$$

# Principal Effects 

Courtesy of Prof. M. De Graef
1st rank cross effects

2nd rank cross effects

## 3rd rank cross effects

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$$
\left(\begin{array}{l}
\Delta S \\
D_{x} \\
D_{y} \\
D_{z} \\
B_{x} \\
B_{y} \\
B_{z} \\
\epsilon_{x x} \\
\epsilon_{y y} \\
\epsilon_{z z} \\
\epsilon_{y z} \\
\epsilon_{x z} \\
\epsilon_{x y}
\end{array}\right)=\left(\begin{array}{ccccccccccccc}
\frac{C}{T} & p_{x} & p_{y} & p_{z} & q_{x} & q_{y} & q_{z} & \alpha_{x x}^{\prime} & \alpha_{y y}^{\prime} & \alpha_{z z}^{\prime} & \alpha_{y z}^{\prime} & \alpha_{x z}^{\prime} & \alpha_{x y}^{\prime} \\
p_{x}^{\prime} & \kappa_{x x} & \kappa_{x y} & \kappa_{x z} & \lambda_{x x} & \lambda_{x y} & \lambda_{x z} & d_{x x x} & d_{x y y} & d_{x z z} & d_{x y z} & d_{x x z} & d_{x x y} \\
p_{y}^{\prime} & \kappa_{y x} & \kappa_{y y} & \kappa_{y z} & \lambda_{y x} & \lambda_{y y} & \lambda_{y z} & d_{y x x} & d_{y y y} & d_{y z z} & d_{y y z} & d_{y x z} & d_{y x y} \\
p_{z}^{\prime} & \kappa_{z x} & \kappa_{z y} & \kappa_{z z} & \lambda_{z x} & \lambda_{z y} & \lambda_{z z} & d_{z x x} & d_{z y y} & d_{z z z} & d_{z y z} & d_{z x z} & d_{z x y} \\
q_{x}^{\prime} & \lambda_{x x}^{\prime} & \lambda_{x y}^{\prime} & \lambda_{x z}^{\prime} & \mu_{x x} & \mu_{x y} & \mu_{x z} & Q_{x x x} & Q_{x y y} & Q_{x z z} & Q_{x y z} & Q_{x x z} & Q_{x x y} \\
q_{y}^{\prime} & \lambda_{y x}^{\prime} & \lambda_{y y}^{\prime} & \lambda_{y z}^{\prime} & \mu_{y x} & \mu_{y y} & \mu_{y z} & Q_{y x x} & Q_{y y y} & Q_{y z z} & Q_{y y z} & Q_{y x z} & Q_{y x y} \\
q_{z}^{\prime} & \lambda_{z x}^{\prime} & \lambda_{z y}^{\prime} & \lambda_{z z}^{\prime} & \mu_{z x} & \mu_{z y} & \mu_{z z} & Q_{z x x} & Q_{z y y} & Q_{z z z} & Q_{z y z} & Q_{z x z} & Q_{z x y} \\
\alpha_{x x} & d_{x x x}^{\prime} & d_{x x y}^{\prime} & d_{x x z}^{\prime} & Q_{x x x}^{\prime} & Q_{x x y}^{\prime} & Q_{x x z}^{\prime} & s_{x x x x} & s_{x x y y} & s_{x x z} & s_{x x y z} & s_{x x x z} & s_{x x x y} \\
\alpha_{y y} & d_{y y x}^{\prime} & d_{y y y}^{\prime} & d_{y y z}^{\prime} & Q_{y y x}^{\prime} & Q_{y y y}^{\prime} & Q_{y y z}^{\prime} & s_{y y x x} & s_{y y y y}^{\prime} & s_{y y z z}^{\prime} & s_{y y y z}^{\prime} & s_{y y x z}^{\prime} & s_{y y x y}^{\prime} \\
\alpha_{z z}^{\prime} & d_{z z x}^{\prime} & d_{z z y}^{\prime} & d_{z z z}^{\prime} & Q_{z z x}^{\prime} & Q_{z z y}^{\prime} & Q_{z z z}^{\prime} & s_{z z x x} & s_{z z y y} & s_{z z z z} & s_{z z y z} & s_{z z x z} & s_{z z x y} \\
\alpha_{y z} & d_{y z x}^{\prime} & d_{y z y}^{\prime} & d_{y z z}^{\prime} & Q_{y z x}^{\prime} & Q_{y z y}^{\prime} & Q_{y z}^{\prime} & s_{y z x x} & s_{y z y y} & s_{y z z} & s_{y z y} & s_{y z x z} & s_{y z x y} \\
\alpha_{x z} & d_{x z x}^{\prime} & d_{x z y}^{\prime} & d_{x z z}^{\prime} & Q_{x z x}^{\prime} & Q_{x z y}^{\prime} & Q_{x z z}^{\prime} & s_{x z x x}^{\prime} & s_{x z y y} & s_{x z z z}^{\prime} & s_{x y y}^{\prime} & s_{x z x z} & s_{x z x y}^{\prime} \\
d_{x y z}^{\prime} & Q_{x y x}^{\prime} & Q_{x y y}^{\prime} & Q_{x y z}^{\prime} & s_{x y x x} & s_{x y y y} & s_{x y z z} & s_{x y y z} & s_{x y x z} & s_{x y x y}
\end{array}\right)\left(\begin{array}{l}
\Delta T \\
E_{x} \\
E_{y} \\
E_{z} \\
H_{x} \\
H_{y} \\
H_{z} \\
\sigma_{x x} \\
\sigma_{y y} \\
\sigma_{z z} \\
\sigma_{y z} \\
\sigma_{x z} \\
\sigma_{x y}
\end{array}\right)
$$

General crystal symmetry shown above.

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$$
\left(\begin{array}{c}
\Delta S \\
D_{x} \\
D_{y} \\
D_{z} \\
B_{x} \\
B_{y} \\
B_{z} \\
\epsilon_{x x} \\
\epsilon_{y y} \\
\epsilon_{z z} \\
\epsilon_{y z} \\
\epsilon_{x z} \\
\epsilon_{x y}
\end{array}\right)=\left(\begin{array}{cccccccccccc}
C \\
T & p_{x} & p_{x} & p_{z} & q_{x} & q_{x} & q_{z} & \alpha_{x x} & \alpha_{x x} & \alpha_{z z} & 0 & 0 \\
p_{x} & \kappa_{x x} & 0 & 0 & \lambda_{x x} & \lambda_{x y} & 0 & 0 & 0 & 0 & d_{x y z} & d_{x x z} \\
p_{x} & 0 & \kappa_{x x} & 0 & -\lambda_{x y} & \lambda_{x x} & 0 & 0 & 0 & 0 & -d_{x x z} & d_{x y z} \\
p_{z} & 0 & 0 & \kappa_{z z} & 0 & 0 & \lambda_{z z} & d_{z x x} & d_{z x x} & d_{z z z} & 0 & 0 \\
q_{x} & \lambda_{x x} & -\lambda_{x y} & 0 & \mu_{x x} & 0 & 0 & 0 & 0 & 0 & Q_{x y z} & Q_{x x z} \\
q_{x} & \lambda_{x y} & \lambda_{x x} & 0 & 0 & \mu_{x x} & 0 & 0 & 0 & 0 & -Q_{x x z} & Q_{x y z} \\
q_{z} & 0 & 0 & \lambda_{z z} & 0 & 0 & \mu_{z z} & Q_{z x x} & Q_{z x x} & Q_{z z z} & 0 & 0 \\
\alpha_{x x} & 0 & 0 & d_{z x x} & 0 & 0 & Q_{z x x} & s_{x x x x} & s_{x x y y} & s_{x x z z} & 0 & 0 \\
\alpha_{x x} & 0 & 0 & d_{z x x} & 0 & 0 & Q_{z y y} & s_{x x y y} & s_{x x x x} & s_{x x z z} & 0 & 0 \\
\alpha_{z z} & 0 & 0 & d_{z z z} & 0 & 0 & Q_{z z z} & s_{x x z z} & s_{x x z z} & s_{z z z z} & 0 & 0 \\
0 & d_{x y z} & -d_{x x z} & 0 & Q_{x y z} & -Q_{x x z} & 0 & 0 & 0 & 0 & s_{x z x z} & -s_{x z y z} \\
0 & d_{x x z} & d_{x y z} & 0 & Q_{x x z} & Q_{x y z} & 0 & 0 & 0 & 0 & s_{x z y z} & s_{x z x z} \\
0 & 0 & 0 & 0 & 0 & 0 & 0 & s_{x x x y} & -s_{x x x y} & 0 & 0 & 0 \\
s_{x y x y}
\end{array}\right)\left(\begin{array}{l}
\Delta T \\
E_{x} \\
E_{y} \\
E_{z} \\
H_{x} \\
H_{y} \\
H_{z} \\
\sigma_{x x} \\
\sigma_{y y} \\
\sigma_{z z} \\
\sigma_{y z} \\
\sigma_{x z} \\
\sigma_{x y}
\end{array}\right)
$$

## Point group 4

## Courtesy of Prof. M. De Graef

$$
\left(\begin{array}{c}
\Delta S \\
D_{x} \\
D_{y} \\
D_{z} \\
B_{x} \\
B_{y} \\
B_{z} \\
\epsilon_{x x} \\
\epsilon_{y y} \\
\epsilon_{z z} \\
\epsilon_{y z} \\
\epsilon_{x z} \\
\epsilon_{x y}
\end{array}\right)=\left(\begin{array}{ccccccccccccc}
\frac{C}{T} & 0 & 0 & 0 & 0 & 0 & 0 & \alpha & \alpha & \alpha & 0 & 0 & 0 \\
0 & \kappa & 0 & 0 & \lambda & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 \\
0 & 0 & \kappa & 0 & 0 & \lambda & 0 & 0 & 0 & 0 & 0 & 0 & 0 \\
0 & 0 & 0 & \kappa & 0 & 0 & \lambda & 0 & 0 & 0 & 0 & 0 & 0 \\
0 & \lambda & 0 & 0 & \mu & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 \\
0 & 0 & \lambda & 0 & 0 & \mu & 0 & 0 & 0 & 0 & 0 & 0 & 0 \\
0 & 0 & 0 & \lambda & 0 & 0 & \mu & 0 & 0 & 0 & 0 & 0 & 0 \\
\alpha & 0 & 0 & 0 & 0 & 0 & 0 & s_{x x x x} & s_{x x y y} & s_{x x y y} & 0 & 0 & 0 \\
\alpha & 0 & 0 & 0 & 0 & 0 & 0 & s_{x x y y} & s_{x x x x} & s_{x x y y} & 0 & 0 & 0 \\
\alpha & 0 & 0 & 0 & 0 & 0 & 0 & s_{x x y y} & s_{x x y y} & s_{x x x x} & 0 & 0 & 0 \\
0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & s_{y z y z} & 0 & 0 \\
0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & s_{y z y z} & 0 \\
0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & s_{y z y z}
\end{array}\right)\left(\begin{array}{c}
\Delta T \\
E_{x} \\
E_{y} \\
E_{z} \\
H_{x} \\
H_{y} \\
H_{z} \\
\sigma_{x x} \\
\sigma_{y y} \\
\sigma_{z z} \\
\sigma_{y z} \\
\sigma_{x z} \\
\sigma_{x y}
\end{array}\right)
$$

Point group $m \overline{3} m$
Note how many fewer independent coefficients there are! Note how the center of symmetry eliminates many of the properties, such as pyroelectricity

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## Homogeneity

- Stimuli and responses of interest are, in general, not scalar quantities but tensors. Furthermore, some of the properties of interest, such as the plastic properties of a material, are far from linear at the scale of a polycrystal. Nonetheless, they can be treated as linear at a suitably local scale and then an averaging technique can be used to obtain the response of the polycrystal. The local or microscopic response is generally well understood but the validity of the averaging techniques is still controversial in many cases. Also, we will only discuss cases where a homogeneous response can be reasonably expected.
- There are many problems in which a non-homogeneous response to a homogeneous stimulus is of critical importance. Stress-corrosion cracking, for example, is a wildly non-linear, non-homogeneous response to an approximately uniform stimulus which depends on the mechanical and electro-chemical properties of the material.


## Use of MuPAD inside Matlab

- Note that the 6x6 transformation matrix can be programmed inside Matlab just as a $3 \times 3$ can.
- In order to apply a transformation (e.g. a symmetry operator) to a $6 \times 6$ stiffness or compliance matrix, the formula is the same as before, i.e.:

$$
C^{\prime}=O C O^{T}
$$

## Matrix

 representation of therotation point groups
What is a group? A group is a set of objects that form a closed set: if you combine any two of them together, the result is simply a different member of that same group of objects. Rotations in a given point group form closed sets - try it for yourself!

Note: the 3rd matrix in the 1 st column (x-diad) is missing a "-" on the 33 element; this is corrected in this slide. Also, in the 2nd from the bottom, last column: the 33 element should be +1 , not -1 . In some versions of the book, in the last matrix (bottom right corner) the 33 element is incorrectly given as -1 ; here the +1 is correct.

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The dashed boxes in this column make up group 32.
cubic branch

|  |  |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: |
| ; 1 | 0 |  | 0 | 0 | -1 |
| ! 0 | 1 | $0 \vdots$ | 0 | -1 | 0 |
| : 0 | 0 | 1 : | -1 | 0 | 0 |
| : 0 | 0 | $1:$ | 0 | 0 | 1 |
| ! 1 | 0 | $0 \vdots$ | 0 | -1 | 0 |
| ! 0 | 1 | $0 \vdots$ | 1 | 0 | 0 |
| - 0 | 1 | $0 \vdots$ | 0 | 0 | 1 |
| : 0 | 0 | 1 | 0 | 1 | 0 |
| ! 1 | 0 | $0 \vdots$ | -1 | 0 | 0 |
| : 0 | -1 | $0:$ | 0 | 0 | -1 |
| : 0 | 0 | $1:$ | 0 | 1 | 0 |
| :-1 | 0 | 0 : | 1 | 0 | 0 |
| $\cdots 0$ | -1 | $0 \vdots$ | -1 | 0 | 0 |
| : 0 | 0 | -1: | 0 | 0 | -1 |
| : 1 | 0 | $0 \vdots$ | 0 | -1 | 0 |
| : 0 | 1 | $0!$ | 1 | 0 | 0 |
| : 0 | 0 | -1: | 0 | 0 | -1 |
| :-1 | 0 | $0:$ | 0 | 1 | 0 |
| $\therefore 0$ | 0 | -1 | 1 | 0 | 0 |
| : 1 | 0 | $0 \vdots$ | 0 | 0 | 1 |
| $\because 0$ | -1 | $0 \vdots$ | 0 | -1 | 0 |
| \% 0 | 0 | -1 | -1 | 0 | 0 |
| :-1 | 0 | 0 : | 0 | 0 | 1 |
| : 0 | 1 |  | 0 | 1 | 0 |
| $\vdots 0$ | 0 | $1 \vdots$ | 0 | -1 | 0 |
| : -1 | 0 | $0 \vdots$ | -1 | 0 | 0 |
| $\bigcirc 0$ | 1 | 0 : | 0 | 0 | -1 |
| -1 | 0 | 0 | 0 | 1 | 0 |
| 0 | 1 | 0 | -1 | 0 | 0 |
| 0 | 0 | -1 | 0 | 0 | (1) |
| -1 | 0 | 0 | 0 | 1 | 0 |
| 0 | -1 | 0 | 1 | 0 | 0 |
| 0 | 0 | 1 | 0 | 0 | -1 |
| 1 | 0 | 0 | 0 | -1 | 0 |
| 0 | -1 | 0 | 1 | 0 | 8 |
| 0 | 0 | -1 | 0 | 0 | (1) |
| 23 |  |  |  |  | ( |
|  |  |  | 432 |  |  |

The dashed box
in this column comprises the 3 -fold axes only.


[^0]:    Table 1 shows a series of tensors that are of importance for material science
    The tensors are grouped by rank, and are also labeled (in the last column) by $E$ (equilibrium property) or $T$ (transport property). The number following this
    letter indicates the maximum number of independent, non-zero elements in the tensor, taking into account symmetries imposed by thermodynamics. The Field and Response columns contain the following symbols: $\Delta T=$ temperature differ ence, $\Delta S=$ entropy change, $E_{i}=$ electric field components, $H_{i}=$ magnetic field components, $\epsilon_{i j}=$ mechanical strain, $D_{i}=$ electric displacement, $B_{i}=$ mag netic induction, $\sigma_{i j}=$ mechanical stress, $\Delta \beta_{i j}=$ change of the impermeability tensor, $j_{i}=$ electrical current density, $\nabla_{j} T=$ temperature gradient, $h_{i}=$ heat flux, $\nabla_{j} c=$ concentration gradient, $m_{i}=$ mass flux, $\rho_{i}^{a}=$ anti-symmetric part of resistivity tensor, $\rho_{i}^{s}=$ symmetric part of resistivity tensor, $\Delta \rho_{i j}=$ change in the component $i j$ of the resistivity tensor, $l_{i}=$ direction cosines of wav

[^1]:    $\dagger$ Axes of symmetry may be rotation axes or inversion axes.
    $\ddagger$ The setting of the reference axes $x_{1}, x_{2}, x_{3}$ in column 6 in relation to the crystallographic axes $x, y, z$ and to the symmetry elements is that shown in column 4. For further notes on axial conventions, see Appendix B.

