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Ceramic Microstructures and Their Elucidation by Imaging, Diffraction and Spectroscopic Methods

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7-11 Sept. 1992

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CERAMIC MICROSTRUCTURES AND THEIR ELUCIDATION BY IMAGING, DIFFRACTION AND SPECTROSCOPIC METHODS.

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The development and potential utilization of ceramic materials is dependent on a systematic effort involving processing, characterization and appropriate property measurements. The methods of characterization are numerous and it is important to employ the one that is appropriate to the problem both in terms of its information content and the achievable level of resolution. With the incorporation of fine probe forming capabilities in a transmission electron microscope and the development of related diffraction, imaging and spectroscopic methods, it is now possible to obtain structural and chemical information from the same region of the sample at high spatial resolution. In this review, recent advances along with representative examples in the application of high resolution electron microscopy (HREM), convergent beam electron diffraction (CBED), low atomic number element microanalysis by x-ray emission spectroscopy (XES), fine structures in electron energy-loss spectroscopy (EELS) and specific site occupancy determination by channelling experiments are discussed.

ALUMINIUM OXYNITRIDE CERAMICS: Pure single crystal aluminium oxide is an anisotropic material and exhibits directional variation in optical and thermal properties. True optical transparency of polycrystalline Al₂O₃ is impossible unless all the grains are identically oriented. An alternative but simpler approach is the stabilization of cubic Al₂O₃ by the addition of nitrogen, in the form of AIN, to form defective spinels. Based on the relatively wide range of compositional stability[1], roughly centered at 35.7 mol% AlN, fully dense materials were sintered for microstructural evaluation. Comparing the symmetries (figure 1) of the whole pattern (4mm), the central transmitted disc (4mm) and the (400) dark field order when set at the Bragg condition (2mm), with the tables of Buxton et al [2], it can be concluded that this material has a projected diffraction group of 4mm₁ᵣ and a point group of m₃m. The displacements of the position of the FOLZ discs with respect to the ZOLZ disc confirms that the material is indeed face centered cubic with a spinel structure. Unfortunately, most AlN powders contain some oxygen impurities and therefore such sintering processes result in materials with polytype-like phases of AlN [3]. The periodicity of the faulting, that defines the character of this modulated structure, is a function of the overall composition and constrained by charge balance. Figure 2 shows a SAD pattern, conventional lattice image and an optical diffractogram from a region of the 32H polytypoid structure. The position of the (010) spot with respect to the transmitted beam in the SAD confirms the hexagonal stacking sequence. From this SAD pattern, the repeat distance along the c-axis can be calculated to be 8.3nm for this structure. However, both the lattice
image and the accompanying optical diffractogram show a spacing of 4.3nm. This is not a surprising
discrepancy, for the nH polytypoids in this system normally consist of two blocks of n/2 layers related
by a c-glide plane. This is verified in figure 3, which shows a CBED pattern obtained with a 60nm
diameter probe (information is averaged over a few unit cells) oriented such that the c-axis is parallel to
the incident beam. From the radius of the FOLZ ring, the periodicity along the c-axis is calculated to be
4.2nm. If these polytypoids are compositionally stabilized, it is predicted that the anion/cation ratio
should be 17/16 or 1.06. A typical EDX spectrum obtained from a 32H region of the sample (figure 4)
analyzed by using experimentally measured K-factors [4,5], confirms a composition of 47.9 at.% Al,
13.8 at.% O and 38.3 at.% N with an experimental error of ±2 %. The anion/cation ratio obtained from
this measurement, i.e. 1.08, agrees well with the predicted value, confirming the stability of such a
long period polytypoid. However, if direct imaging of the structure with atomic resolution is used
(figure 5), the periodicity along the c-direction can be shown to be a sequence of alternate 7:9:7:9 ...
repeats. This result can be interpreted as parallel intergrowths of the 27R (c = 7.2 nm) and 21R (c =
5.7 nm) polytypoids. Clearly adjacent units of 21R and 27R add up to the 32H repeat units. A detailed
interpretation of these images would require simulations of the image based on a predicted intergrowth
structure.

NOVEL GRAPHITES: A novel graphite material, BC3, has been synthesized by the interaction
of benzene and boron trichloride, at 800 °C. These reactions: 2 BCl3 + C6H6 -> 2 BC3 + 6 HCl , are
driven by favourable TDS values associated with HCl elimination. For BC3, an ordered structure with
periodic substitution of B for every other C in the graphite lattice has been suggested. Recently,
important differences in the density of states (DOS) of BC3 from graphite, for the proposed structure
has been suggested[6]. Two sharp peaks in the DOS associated with the p band below the conducting s
bands(figure 6a), unlike graphite with one peak, are predicted. Representative K-edge spectra of B and
C obtained from a thin foil of BC3 are shown in figure 6b. The fine structure of the B K-edge shows
two sharp peaks at 187.5eV and 190.1eV. These peaks can be assigned to the excitation of the 1s
electrons to unoccupied levels in the conduction band. Observation of these features provides direct
confirmation for the atomic arrangement in this material [7].

IRON OXIDES: The ability to resolve the oxidation state and the coordination of iron in various
spinels is important in understanding the magnetic behaviour of a wide range of magnetic oxides. Four
different model oxide mineral sample, each containing iron in a specific oxidation state and
coordination, were investigated. They were franklinite ( nominal composition [Zn,Mn]Fe3+204 )
where Fe(III) is in octahedral sites; hercynite (nominal composition Fe2+Al2O4 ) where Fe(II) is in
tetrahedral sites; fayalite (nominal composition Fe2Si04 ) where Fe(II) is in octahedral coordination;
and an iron analogue of leucite (nominal composition KFeSi2O6 ) where Fe(III) is in tetrahedral
coordination. Figures 7a and b show the L3/L2 edges for Fe(III) in octahedral and tetrahedral
coordination respectively. In the former case, both the L3 and L2 edges show distinct splittings,
separated by 1.6eV, at the onset of the edge. No such splitting is observed in the latter. For Fe(II) in
octahedral and tetrahedral sites, the \( L_3/L_2 \) peak height ratio is lower when compared to Fe(III). The observed fine structures can be interpreted in terms of a simple ligand field approach using the commonly observed low spin state for iron[8]. For Fe(III) in octahedral coordination, in addition to the empty \( e_g \) levels there is an unpaired electron in the \( t_{2g} \) level. This explains the splitting of the \( L_3 \) and \( L_2 \) white line features in Figure 7a. For Fe(III) in tetrahedral coordination, the energies of the ligand field split levels, \( e_2 \) and \( t_2 \), are reversed. The lower \( e_2 \) level is completely filled and therefore one observes a single peak (fig.7b).

ATOM LOCATION BY CHANNELLING ENHANCED MICROANALYSIS: For crystalline materials both the characteristic energy-loss edges and x-ray intensities are governed by phenomena that are highly localized at the atomic sites. They show strong dependence on the orientation of the incident beam and this has been developed into a powerful crystallographic technique for specific site occupancy/valence determinations [9]. For the dolomite structure in the [1210] orientation (figure 8) when the (0001) systematic row is excited the wave field of the dynamically diffracted electrons in the crystal is two-dimensional, i.e., constant in a direction normal to the c-axis. In this structure, the candidate sites of interest (Mg and Ca) occupy alternating planes at one third the unit cell parameter along the c-axis. For the (0001) systematic row, at orientations corresponding to small positive excitation errors (s>o) (figure 9) of the third order Bragg diffraction condition (3g) an enhancement of the Mg Ka is observed. For negative excitation errors (s<o) an enhancement of the Ca Ka is observed. The impurity Fe Ka intensity follows that of Mg Ka, suggesting similar site occupancy. The exact degree of substitution of the impurity atoms (Fe) can be obtained by an elegant method of ratios with respect to the intensities of the reference elements (Mg & Ca) in the host lattice [9]. Site occupancies in a variety of ceramic materials have been measured using this technique. In the case of EELS, combining these orientation dependence with detectable changes in oxidation states (by the small chemical shifts observed in the onset of the core-loss edges) it is also possible to obtain specific site valence information[10].

A NOTE ON LOW-Z MICROANALYSIS: For the characterization of ceramic materials, improvement in the accuracy of microanalysis of low-z elements is desired. For XES, using an UTW detector best results are obtained using experimentally determined k-factors, accurate measurement of sample thickness and correction for absorption in the specimen [5]. For EELS, comparison of experimentally measured partial cross-sections for four different scattering geometries with theoretical calculations using both the standard hydrogenic model and parametrized Hartree-Slater cross-sections show best agreement when experiments are performed in diffraction mode (image coupling) with the probe convergence angle (0.84 mrad) much smaller than the spectrometer collection angle (6.84 mrad) [11].
REFERENCES

FIGURE CAPTIONS
1. CBED patterns in the <100> orientation confirming the spinel structure of AlON.
2. SAD pattern, lattice image and optical diffractogram from the hexagonal long period 32H polytypoid structure.
3. A CBED pattern in the <0001> orientation for the 32H polytypoid. The structure with c=8.4nm has a glide plane at half the unit cell parameter.
4. XES spectrum with the UTW detector and analysis from 32H.
5. A HREM image of the 32H obtained on the atomic resolution microscope. The 16 layer structure is shown to be an intergrowth of 9 and 7 layer repeats.
6a. Total DOS of a BC3 monolayer showing the splitting of the p band.
6b. B-K and C-K edges from BC3 with the C-K edge for graphite superimpose for comparison.
7. L3 and L2 edges for Fe(III) in (a) octahedral and (b) tetrahedral coordination along with the ligand field splitting.
8. Crystallographic structure of dolomite projected along the c-axis.
Figure 2
Figure 3
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TOTAL 100.00

Figure 4
Figure 5
Figure 6a
Figure 6b
Figure 7
Figure 8
Figure 9