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Calculation of x-ray intensity from a rough sample based on a statistical model

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An x-ray intensity correction is developed which begins with a roughness model that is often used to describe real surfaces. This is based upon a normal distribution of surface asperities relative to a mean plane. Pair correlation between absorbing elements along x-ray paths either entering or leaving the sample with respect to the signal producing element is accomplished by means of an exponential autocorrelation function. This allows the degree of roughness to be varied on a local scale to fit specific surfaces using statistical data. Equations are developed to describe x-ray fluorescence and diffraction signals for symmetric and asymmetric beam optics. Theory is compared with experiment using a roughened, fully stabilized zirconia sample.

INTRODUCTION

A correction by Harrison and Paskin\(^1\) deals with the effect of granularity on the diffracted x-ray intensity from powders. The absorption lengths associated with a signal-producing element in powders is changed from the ideal value for a theoretically dense sample to a smaller value due to the presence of pores. Partial correlations of the absorption lengths for the incident and scattered rays produce an intensity correction relative to a theoretically dense sample. Likewise, for a rough sample, the absorption lengths for pairs of incident and scattered rays are altered by surface roughness and the intensity falls below that of an ideally flat sample. Granularity and roughness effects are similar in the sense that both produce a distribution of path lengths at each fixed depth below the sample surface. Harrison and Paskin’s work is restricted to symmetrical x-ray optics, i.e., the incident and scattered rays must make the same angle with the mean sample surface. Also, it is not easily applied to surfaces described by statistical functions.

An exact calculation for a 45° sawtooth surface was made by Borie.\(^2\) Although such a surface differs greatly from reality, this calculation has the value in providing physical insight to the roughness problem and in establishing an upper limit for the correction over a full range of scattering angles. As one might expect, the maximum correction occurs when both rays are parallel to the sawtooth making an angle of 45° with the mean surface. Again, symmetrical x-ray optics as well as a common absorption coefficient for incident-signal pairs were assumed.

This work makes use of a generally accepted statistical model to describe surface asperities,\(^3\) and thereby represents a first treatment for the roughness effect from real surfaces. We assume that the heights of the surface profile are normally distributed with a standard deviation \(\sigma\), and the pair correlation between different asperities is determined by an autocorrelation function which decays exponentially with distance. This latter parameter controls the degree of roughness on a local scale. The restrictions of symmetrical optics and a common absorption coefficient for the incident and the signal rays are avoided. The latter allows a treatment of x-ray fluorescence as well as diffraction problems for a range of optical paths. Data are presented for a severely ground zirconia sample that are in agreement with the statistical model developed herein.

THE STATISTICAL ROUGHNESS MODEL—ABSORPTION

We use a normal distribution function to describe the heights of surface excursions:

\[
p(Z_s) = \frac{1}{\sqrt{2\pi \sigma^2}} e^{-Z_s^2},
\]

where \(\sigma\) is the standard deviation and \(Z_s\) is the height measured from the mean surface scaled in units of \(\sqrt{2}\sigma\). A pair of excursion heights at two positions separated by a horizontal distance \(\tau\) are distributed according to the following two-dimensional normal distribution function:

\[
p(Z_{1s}, Z_{2s}, \tau) = \frac{1}{2\pi \sigma^2 \sqrt{1 - \rho(\tau)^2}} \exp\left\{-\frac{1}{2} \rho(\tau)^2 (Z_{1s} - Z_{2s})^2\right\},
\]

where \(Z_{1s}\) and \(Z_{2s}\) are the scaled heights at two positions and \(\rho(\tau)\) is the autocorrelation coefficient function which determines the degree of likeness between excursions \(Z_{1s}\) and \(Z_{2s}\). \(\rho(\tau)\) is the normalized form of the autocorrelation function defined by

\[
R(\tau) = \lim_{L \to \infty} \frac{1}{2L} \int_{-L}^{L} Z(X)Z(X + \tau) dX,
\]

where \(X\) is parallel to the mean surface plane and \(-Z\) is the distance below the mean surface. By definition the variance of the height distribution is

\[
R(0) = \lim_{L \to \infty} \frac{1}{2L} \int_{-L}^{L} Z^2(X) dX = \sigma^2
\]

and

\[
\rho(\tau) = \frac{R(\tau)}{R(0)} = \frac{R(\tau)}{\sigma^2}.
\]

When \(\tau = 0\), \(Z_{1s}\) and \(Z_{2s}\) are coincident, therefore, \(\rho(0) = 1\). For \(\tau = \infty\), \(Z_{1s}\) and \(Z_{2s}\) are completely uncor-
lated, and \( \rho(\omega) = 0 \). Between the two extremes of \( r = 0 \) and \( r = \infty \), \( Z_s \) and \( Z_{s'} \) are partially correlated with decreasing correlation for increasing \( r \). The function

\[
\rho(r) = e^{-|r'|/\tau_s}
\]

is commonly used in tribology and will be used in this development. \( \tau_s \) is a correlation distance which indicates how quickly a surface excursion loses correlation with increasing distance \( r \) from a neighbor.

The scaled form of the conditional probability density function based upon a normal distribution is

\[
p(Z_{s'}/Z_s,\tau) = \frac{p(Z_{s'}/Z_s,\tau)}{p(Z_s)} = \frac{1}{\sqrt{2\pi}\sigma(1 - \rho^2(\tau))^{1/2}} \times e^{-\left[\frac{(Z_{s'} - \mu(\tau)Z_s)^2}{2(1 - \rho^2(\tau))}\right]}.
\]

(7)

This represents the probability of an excursion \( Z_{s'} \) under the condition that a prescribed excursion \( Z_s \) exists at a horizontal distance \( \tau \) away.

The kinematic intensity from a polycrystalline solid is taken as a sum of intensities produced at each volume element. Consider a sample consisting of planes of signal producing elements \( dV \) (Fig. 1) located at a depth \( -Z \) and extending to \( -Z_0 - dZ \). If the \( X \) axis is parallel to the mean surface plane, the volume element in a two-dimensional model is \( dX \times dZ \). Incident ray \( A' \rightarrow A \) produces a signal at \( A \) which follows a signal path \( A \rightarrow A'' \). The distance \( \tau \) is measured from each element \( A \) along \( \pm X \). Since there are two rays that influence the detected signal, two variables, \( \tau = \tau_s \) and \( \tau = \tau_s \), are required, extending in \( \pm X \) directions from \( A \). Two correlated conditional functions \( p(Z_{s'}/Z_s,\tau) \) and \( p(Z_{s''}/Z_{s'},\tau_s) \) are used to describe absorption along \( A' \rightarrow A \) and \( A \rightarrow A'' \) with correlation determined by \( |\tau'|/\tau_s \) in Eq. (6).

As shown in Fig. 1, volume elements within the same horizontal layer can be divided into groups. Each group is defined by a specific excursion height above the signal generating element and the location of the layer. For example, volume elements \( A, B, C, D \) shown in Fig. 1, are in a common layer, and belong to the same group because the excursion heights are the same. This group is designated by \( (Z_s, Z_{s0}) \).

Clearly the strength of \( x \)-ray fluorescence or diffraction signals depends upon incident and signal ray pairs. In the present model, these are coupled relative to the intermediate signal element through \( \rho(\tau) \). This approaches unity as either \( \tau_s/\tau \) or \( \tau_s/\tau \) approach zero. And, coupling increases to the extent that neighboring surface asperities must have the same \( Z_{s0} \). A large \( \tau_s \) causes neighboring points along the surface to look alike. At the other extreme, a small \( \tau_s \) causes nearby points to be unrelated. Here the surface asperities vary rapidly with \( X \).

For the present, consider only the signal ray. The same development and result will be used for the incident ray. A signal ray from an element of the \( (Z_s, Z_{s0}) \) group makes an angle of \( \theta_s \) with the horizontal plane (Fig. 2). Surface traces corresponding to all "elements" in this group are superimposed to illustrate the distribution of material above the signal ray. If each excursion intersects the signal ray only once, then the signal ray is completely within material below the intersection and completely outside above. The distribution of the intersections along the signal ray determines the distribution of its absorption lengths. By restricting this development to one intersection, the distribution of the intersections along the signal ray can be determined rather simply. Although a single intersection model is not valid for small \( \theta_s \) and \( \theta_s \), it treats most cases normally encountered with conventional x-ray optics.

In Fig. 3, a point absorber on the signal ray is shown located at a scaled height of \( Z_{a0} = Z/\sqrt{2}\sigma \) and separated by a scaled horizontal distance \( \tau_s = \tau/\sqrt{2}\sigma \) from the signal generating element. Vertical lines passing through this point and the signal generating element are shown as an absorption line and a signal line. Since every excursion of the \( (Z_s, Z_{s0}) \) group crosses the absorption line once, the fraction of crossings located at or above \( Z_{a0} \) is the cumulative probability function from \( Z_{s0} \) to \( \infty \), i.e.,

\[
P(Z_{a'}/Z_{s0}, \tau) = \sqrt{2}\sigma \int_{Z_{s0}}^{\infty} p(Z_{a'}/Z_{s0}, \tau_s) dZ_{a'}
\]

(8)

Equation (8) is valid for any point because the point absorber on the signal ray at \( Z_{a0} \) is arbitrarily chosen along the signal ray. The behavior of Eq. (8) is as expected. With ideal correlation, \( \rho \) becomes unity and erfc is either unity or zero.

\[
P(Z_{a'>Z_{s0}}/Z_{s0}, \tau) = 0 \quad \text{or} \quad P(Z_{a'<Z_{s0}}/Z_{s0}, \tau) = 1.
\]

Figure 3 illustrates two closely spaced absorption lines.
intersecting the signal ray at heights of \( Z_{i,s} \) and \( Z_{i,s} + dZ_{i,s} \), respectively. If the total number of signal generating elements in the \((Z_{i,s},Z_{0,i})\) group is \( N \), then the total number of excursions starting from the signal line is also \( N \). \( NP(Z_{i,s}/Z_{0,i},\tau_{s}) \) and \( NP(Z_{i,s} + dZ_{i,s}/Z_{0,i},\tau_{s} + d\tau_{s}) \) excursions are located above the signal ray at the two absorption lines, respectively. The difference

\[
n = NP(Z_{i,s}/Z_{0,i},\tau_{s}) - NP(Z_{i,s} + dZ_{i,s}/Z_{0,i},\tau_{s} + d\tau_{s})
\]
gives the number of excursions passing through the signal ray in the differential spacing between the two absorption lines. Because the total number of intersections between the excursions and the signal ray is also \( N \), \( n \) out of \( N \) intersections gives an absorption length of \( l_{i} = \sqrt{2\sigma(Z_{i,s} - Z_{i})/\sin \theta_{i}} \). Consequently, the probability density function \( p(l_{i}) \) for the absorption length distribution becomes

\[
p(l_{i})dl_{i} = n/N = - \left\{ P(Z_{i,s} + dZ_{i,s}/Z_{0,i},\tau_{s} + d\tau_{s}) \right\} \left\{ - P(Z_{i,s}/Z_{0,i},\tau_{s}) \right\}.
\]

(9)

Because

\[
P(Z_{i,s} + dZ_{i,s}/Z_{0,i},\tau_{s} + d\tau_{s})
\]

\[
\approx P(Z_{i,s}/Z_{0,i},\tau_{s}) + P'(Z_{i,s}/Z_{0,i},\tau_{s})dZ_{i,s},
\]

\[
p(l_{i})dl_{i} = - P'(Z_{i,s}/Z_{0,i},\tau_{s})dZ_{i,s},
\]

(10)

the derivative of \( P(Z_{i,s}/Z_{0,i},\tau_{s}) \) with respect to \( Z_{i,s} \) gives the required probability density for absorption paths of length \( l_{i} \), but written as a function of the \( Z \) coordinate. It can be shown that

\[
p(l_{i})dl_{i} = \frac{1}{\sqrt{\pi}} \left[ \exp \left( - \frac{(Z_{i,s} - \rho Z_{0,i})^2}{1 - \rho^2} \right) \right] \times \left( \frac{1}{1 - \rho^2} - \frac{\rho^2 Z_{i,s} - \rho Z_{0,i}}{\tau_{s} \tan \theta_{i}(1 - \rho^2)^{3/2}} \right)dZ_{i,s}.
\]

(11)

The average absorption factor for group \((Z_{i,s},Z_{0,i})\) is calculated from

\[
\langle e^{-\mu l_{i}} \rangle = \int_{0}^{\infty} e^{-\mu l_{i}}p(l_{i})dl_{i}
\]

\[
= - \int_{Z_{i,s}}^{\infty} e^{-\left[ \frac{\sqrt{2\sigma(Z_{i,s} - Z_{i})/\sin \theta_{i}}}{\sqrt{1 - \rho^2}} \tan \theta_{i}(1 - \rho^2)^{3/2} \right]}dZ_{i,s}
\]

\[
\times P'(Z_{i,s}/Z_{0,i},\tau_{s})dZ_{i,s},
\]

(12)

where \( \mu \) is the linear absorption coefficient of the sample for the signal ray.

A similar result is obtained for the incident beam by replacing \( \theta_{i} \) with \( \theta_{s} \) (the incident angle) and \( \mu_{s} \) with \( \mu_{i} \) (the linear absorption coefficient for the incident ray). With these substitutions in Eqs. (11) and (12), we obtain the average absorption attenuation factor \( \langle e^{-\mu l_{i}} \rangle \) for the incident ray.

The average absorption attenuation factor for one group which includes both the incident and signal rays is taken as

\[
A(Z_{i,s},Z_{0,i}) = \langle e^{-\left(\mu_{i} + \mu_{s}\right)l_{i}} \rangle
\]

\[
= \int_{0}^{\infty} \int_{0}^{\infty} p(l_{i},l_{s})e^{-\left(\mu_{i} + \mu_{s}\right)l_{i}}dl_{i}dl_{s},
\]

(13)

where \( l_{i},l_{s} \) are related to the \( Z \) coordinate by \( l_{i} = \sqrt{2\sigma(Z_{i,s} - Z_{i})/\sin \theta_{i}} \), etc. \( p(l_{i},l_{s}) \) is the probability density for \( l_{i} \) and \( l_{s} \), and is determined by taking the product of two functions which are correlated through the excursion height at the intermediate signal line. The mutual distribution of the intersections at the extreme ends is ignored. This approximation gives

\[
A(Z_{i,s},Z_{0,i}) = \langle e^{-\left(\mu_{i} + \mu_{s}\right)l_{i}} \rangle
\]

\[
= \int_{0}^{\infty} p(l_{i})p(l_{s})e^{-\left(\mu_{i} + \mu_{s}\right)l_{i}}dl_{i}dl_{s}
\]

\[
= \int_{0}^{\infty} p(l_{i})e^{-\mu_{i}l_{i}}dl_{i}\int_{0}^{\infty} p(l_{s})e^{-\mu_{s}l_{s}}dl_{s}.
\]

(14)

Strong pair correlation between \( l_{i} \) and \( l_{s} \) occurs when both \( \theta_{i} \) and \( \theta_{s} \) are close to 90°, while weak correlation can be associated with either \( \theta_{i} \) or \( \theta_{s} \) becoming small. Note that for the more common symmetrical case \( \theta_{i} = \theta_{s} \) and \( p(l_{i}) = p(l_{s}) \).

It is important to note that when incident and signal rays are on the same side of the sample normal, the signal line does not serve as an intermediate point. Under these conditions, mutual correlation between beam paths must also be considered. Therefore, \( \theta_{i} \) and \( \theta_{s} \) are restricted to a range extending from 0° to 90°.

X-RAY INTENSITY FROM THE STATISTICAL ROUGHNESS MODEL

The signal intensity contributed by a specific group \((Z_{i,s},Z_{0,i})\) is

\[
\Delta I(Z_{i,s},Z_{0,i}) = I_{0}qA_{V}(Z_{i,s},Z_{0,i})A(Z_{i,s},Z_{0,i}),
\]

(15)

where \( I_{0} \) is the incident intensity. For a fluorescence signal from a specific element \( i \),

\[
Q = Q_{i} = N_{i}(D_{i}/4\pi R^{2}).
\]

(16)

where \( N_{i} \) = number of \( i \) atoms per volume, \( \sigma_{i} \) = atomic cross section of atom \( i \) for producing fluorescence of one type, \( D_{i} \) = dimensionless absorption factor for fluorescence
radiation in the path from sample surface to detector times the detector efficiency, and \( R = \text{sample to detector distance} \). For a diffraction signal from a powder at a specific reflection,\(^5\)

\[
Q = Q_d = \text{const} \left( \frac{2}{v_r^2} \right) \frac{1 + \cos^2 2\theta \cos^2 2\theta'}{\sin \theta \sin 2\theta} f^2, \tag{17}
\]

where \( Q_d \) = reflectivity, \( \lambda \) = wavelength of the incident beam, \( v_r \) = volume of the unit cell, \( 2\theta = \text{Bragg angle}, 2\theta' = \text{Bragg angle for monochromator}, j = \text{multiplicity}, \) and \( F = \text{structure factor}. \Delta V(Z_s, Z_o) \) is the total volume of all elements in group \((Z_s, Z_o)\). Because the height distribution of excursions is normal,

\[
\Delta V(Z_s, Z_o) = \Delta V(2\theta \sigma(Z_o))dZ_o, \tag{18}
\]

where \( \sigma(Z_o) \) is the normal probability density function given in Eq. (1). Here \( \Delta V \) is given by (see Fig. 1)

\[
\Delta V = (\sqrt{2\pi} \sigma \sin \theta) dZ_s, \tag{19}
\]

where \( A_j \) is the cross-sectional area of the incident beam. Combining Eqs. (1), (15), (18), and (19),

\[
dl(Z_s, Z_o) = \frac{\sqrt{2\pi} \sigma_l Q_{AO}}{\sqrt{\pi} \sin \theta} e^{-Z_s^2} A(Z_s, Z_o) dZ_o dZ_s. \tag{20}
\]

The intensity contributed by a layer at \( Z_s \) is simply the integral over all asperities at the signal line above \( Z_s \):

\[
dl(Z_s) = \frac{\sqrt{2\pi} \sigma_l Q_{AO}}{\sqrt{\pi} \sin \theta} \int_{-\infty}^{\infty} e^{-Z_s^2} A(Z_s, Z_o) dZ_o dZ_s. \tag{21}
\]

Although the upper limit should be infinity, a value of 3 causes the integrand to vanish and is readily treated in numerical calculations.

The total intensity is the summation of the intensities contributed by each layer. Integrating Eq. (21) over all possible \( Z_s \),

\[
I_n = \frac{\sqrt{2\pi} \sigma_l Q_{AO}}{\sqrt{\pi} \sin \theta} \int_{-\infty}^{\infty} e^{-Z_s^2} A(Z_s, Z_o) dZ_o dZ_s. \tag{22}
\]

The intensity from an ideally flat sample provides a useful reference and can be obtained by integrating the intensity scattered by each complete layer

\[
I_f = \int_{-\infty}^{\infty} I_0 Q_{AO} \frac{1}{[\sin \theta_i + \sin \theta_o]} \exp \left( \frac{\mu_i - Z_s}{\sin \theta_i} + \frac{\mu_o - Z_s}{\sin \theta_o} \right) dZ_s
\]

\[
= I_0 Q_{AO} \frac{1}{[\sin \theta_i + \sin \theta_o]} \exp \left( \frac{\mu_i - Z_s}{\sin \theta_i} + \frac{\mu_o - Z_s}{\sin \theta_o} \right) dZ_s. \tag{23}
\]

Again, the numerical integration of Eq. (22) is conveniently taken over \( \pm 3 \) without significant truncation error. For material located below \( Z_s = -3 \), the surface asperities are negligible and a flat sample calculation is used:

\[
\frac{I_0 Q_{AO}}{\sin \theta_i} \int_{-\infty}^{\infty} \exp \left( \frac{\mu_i - Z_s}{\sin \theta_i} + \frac{\mu_o - Z_s}{\sin \theta_o} \right) dZ_s
\]

\[
= I_0 Q_{AO} \left( \frac{\mu_i}{\sin \theta_i} + \frac{\mu_o}{\sin \theta_o} \right) e^{-Z_s^2} \left( \frac{1}{\sin \theta_i} + \frac{1}{\sin \theta_o} \right) \tag{24}
\]

This result is combined with Eq. (22) according to the previous discussion, and divided by Eq. (23). Consequently, the ratio \( R \), giving the intensity reduction for a rough sample, is

\[
R = \frac{I_{\text{rough sample}}}{I_{\text{flat sample}}} = \sqrt{2\pi} \sigma \left( \frac{\mu_i}{\sin \theta_i} + \frac{\mu_o}{\sin \theta_o} \right) \times \int_{-\infty}^{\infty} \int_{-\infty}^{\infty} e^{-Z_s^2} A(Z_s, Z_o) dZ_o dZ_s
\]

\[
+ e^{-3Z_s^2} \left[ \frac{\mu_i}{\sin \theta_i} + \frac{\mu_o}{\sin \theta_o} \right]. \tag{25}
\]

A computer program with repeated use of 12-point Gauss-Legendre quadrature integrations is used to evaluate Eq. (25).

EXPERIMENTAL VERIFICATION OF THE STATISTICAL MODEL

The statistical model, which we have developed, is compared with measured intensities from a roughly fully stabilized zirconia (FSZ), containing 8 mol % \( Y_2O_3 \), and one which is identical but is well polished. For the ground (rough) sample, we obtained \( \sigma = 6.2 \mu m \) and \( \tau_r = 82 \mu m \) by profilometer measurements. A typical surface profile for the rough sample is shown in Fig. 4, while the corresponding autocorrelation function calculated directly from the complete digitized profile, is shown in Fig. 5.

The intensity ratio given by Eq. (25) is evaluated using the \( \sigma \) and \( \tau_r \) values of the rough sample for symmetrical diffraction and \( CuK \alpha \) radiation. The linear absorption coefficient for the FSZ and this radiation is 650 cm\(^{-1}\). The calculated result is shown in Fig. 6 as the solid curve while the dashed line represents an extrapolation which is likely to be influenced by multiple surface intersections. Experimental ratios of integrated intensities measured from the flat and rough samples as shown in Fig. 6. These measurements were carried out using a symmetrically aligned Siemens diffractometer, and a diffracted beam quartz \( K \alpha \) monochromator. Two overlapping slow scans (0.1 min) were made over each peak to obtain a better statistical average. This gave a total of 12 peaks for each sample. Peak areas,
corrected for background, were measured using a planimeter. The theoretical calculations based on our statistical model follow the same trend as the experimental data.

**DISCUSSION**

An examination of the theoretical curve of Fig. 6 illustrates that no sharp minimum exists over the range of experimental observations. For a 45° sawtooth a sharp minimum is found at $\theta_i = \theta_s = 45°$. Rays traveling parallel to the edge of a sawtooth experience large fluctuations in the absorption pathlengths from point to point along the $X$ axis. It is known that these rays introduce a large correction to the intensity. Changing the sawtooth angle would shift the $\theta$ position for a maximum correction. With real surfaces one finds a distribution of sawtoothlike fluctuations having nearly linear regions for either the incident beam or the signal beam to follow. When this parallel tuning between the free surface and either (or both) beam(s) occurs, the fluctuations in absorption path length produce an increase in the intensity correction. The broad intensity correction curve shown in Fig. 5 may be explained on this basis. In other words, a real surface may be treated as a distribution of slopes. If one numerically calculates the average absolute slope from the profilometer data (Fig. 1), a value of 9° is obtained. This agrees almost exactly with the minimum of Fig. 6. We point this out with a degree of caution because below this angle multiple intersections of the surface with the incident and signal rays become more likely. The theory presented here is based upon a single intersection model. If the fluctuation in path lengths is determined largely by local fluctuations in path length near the signal element, then the minimum may be approximated by a single intersection model. A simple formula for the minimum based upon one example is given by

$$\theta_{\text{min}} = \tan^{-1}(2.3\sigma/\tau_s).$$

This result is given as a rough guideline in future studies.

If one refers back to Fig. 6, one sees that as $\theta_i = \theta_s \rightarrow 90°$, the integrated intensity equals that of the ideally flat sample. This is due to complete correlation between the incident and signal rays. Also, one observes that the experimental points are systematically below the theoretical curve. This is believed to result from residual stress gradients produced in the severely ground sample. Residual stresses near the surface were found to produce a range of interplanar spacings and an associated redistribution of the Bragg intensity into weak tails which are difficult to measure. A few percent of the intensity in the ground sample is lost because of this effect, causing the data points to be slightly lower than theory.

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