

2. DIFFRACTION GEOMETRY AND ITS PRACTICAL REALIZATION

AT-cut quartz plate, or on Mylar for transmission. It is essential to rotate the specimen and increase the count time. A Gandolfi camera has also been used for very small specimens (see Section 2.3.4). A high-brilliance microfocus X-ray source has been used with a collimator made of 10 to 100 μm internal-diameter capillary tube. An X-Y stage is used with an optical microscope to locate selected areas of the specimen.

A microdiffractometer has been designed for microanalysis, Fig. 2.3.1.15 (Rigaku Corporation, 1990). It has been used to determine phases and stress in areas $< 10^4 \mu\text{m}^2$ (Goldsmith & Walker, 1984). The key to the method is the use of an annular-ring receiving slit, which transmits the entire diffraction cone to the detector instead of a small chord as in conventional diffractometry, thereby utilizing all the available intensity. The pattern is scanned by translating the ring and detector along the direct-beam path so that

$$2\theta = \arctan(R_{\text{RS}}/L), \quad (2.3.1.26)$$

where R_{RS} is the radius of the ring slit and L the distance from the fixed specimen. For $R_{\text{RS}} = 15 \text{ mm}$, L varies from 171 to 9 mm in the transmission range 5 to $60^\circ 2\theta$; a 50 mm diameter scintillation counter is used. A doughnut-shaped proportional counter (3/4 of a full circle) is used for the 30 to 150° reflection specimen mode. The slit width is 0.2 mm and the aperture varies with 2θ . The intensities fall off at the higher 2θ 's because of the small incidence angles to the slit. An alternative method uses a position-sensitive proportional counter. Steinmeyer (1986) has described applications of microdiffractometry.

By using synchrotron radiation (Section 2.3.2), single-crystal data for structure determination can now be obtained from a microcrystal about 5–10 μm in size; see Andrews *et al.* (1988), Bachmann, Kohler, Schultz & Weber (1985), Harding (1988), Newsam, King & Liang (1989), Cheetham, Harding, Mingos & Powell (1993), Harding & Kariuki (1994), and Harding, Kariuki, Cernik & Cressey (1994).

2.3.2. Parallel-beam geometries, synchrotron radiation

The radiation from the X-ray tube is divergent and various methods can be used to obtain a parallel beam as shown in Fig. 2.3.2.1. Symmetrical reflection from a flat crystal is the usual method. An asymmetric reflecting monochromator with small incidence angle and large exit angle expands the beam, or in reverse condenses it (§2.3.5.4.1). A channel monochromator has the advantage of not changing the beam direction. A receiving slit or preferably Soller slits can be used to define the diffracted beam. A graphite monochromator in the diffracted beam or a solid-state detector eliminates fluorescence. The incident-beam

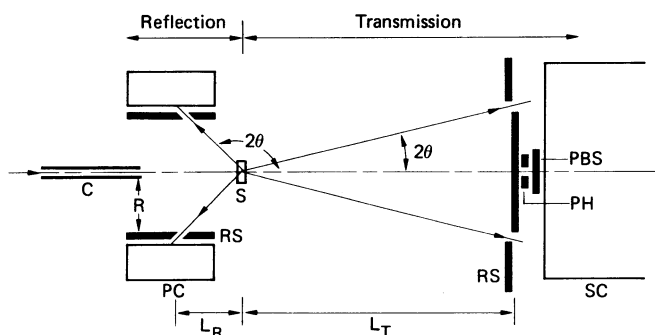


Fig. 2.3.1.15. Rigaku microdiffractometer for microanalysis. *C* collimator, *PC* ring proportional counter, *RS* ring slit with radius r , *S* specimen, *SC* scintillation counter, *PBS* primary beam stop, *PH* pinhole for alignment, L specimen-to-receiving-slit distance.

parallel slits limit vertical divergence. However, all the methods result in a large loss of intensity compared with conventional focusing. In contrast, the storage ring produces a virtually parallel beam with very small vertical divergence of about 0.1 mrad, and the monochromator is used only to select the wavelength. The rest of this section assumes a synchrotron-radiation source.

Storage-ring X-ray sources have a number of unique properties that are of great importance for powder diffraction. The advantages of synchrotron powder diffraction have been described by Hastings, Thomlinson & Cox (1984), Parrish & Hart (1987), Parrish (1988), and Finger (1989). Excellent patterns with high resolution and high peak-to-back ground ratio have been reported. These include the orders-of-magnitude higher intensity and nearly uniform spectral distribution compared with X-ray tubes, the wide continuous range of selectable wavelengths, and the single profile that avoids the problems caused by $K\alpha$ doublets and β filters. Owing to major differences in the diffractometer geometries, comparisons of intensities with X-ray tube focusing methods cannot be predicted simply from the number of source photons.

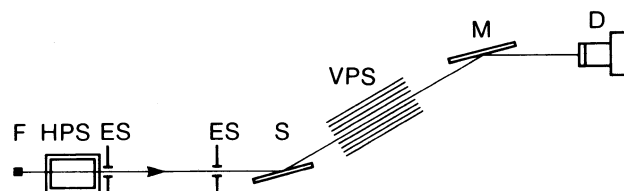


Fig. 2.3.2.1. Method to obtain parallel beam from X-ray tube for powder diffraction. HPS parallel slits to limit axial divergence, ES entrance slits (can be replaced by pair of flat parallel steel bars), *S* specimen, VPS parallel slits to define diffracted beam, *M* flat monochromator (can be omitted), *D* detector. See also Fig. 2.3.2.4(a).

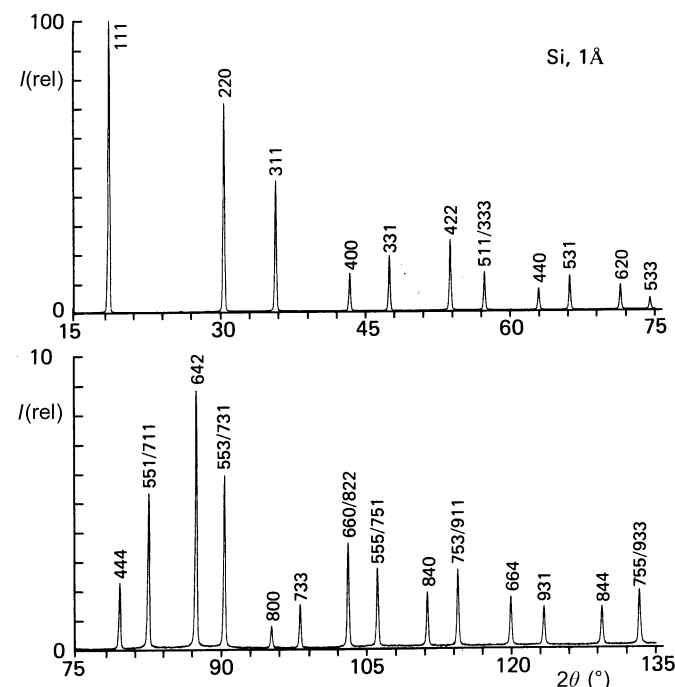


Fig. 2.3.2.2. Silicon powder pattern with 1 \AA synchrotron radiation using method shown in Fig. 2.3.2.4(a). The 444 reflection is the limit for Cu $K\alpha$ radiation.