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Argentum EX1040

## METHOD II (AZEOTROPIC-TOLUENE DISTILLATION)

Apparatus-Use a $500-\mathrm{mL}$ glass flask $A$ connected by means of a trap $B$ to a reflux condenser $C$ by ground glass joints (see figure).


## Toluene Moisture Apparatus

The critical dimensions of the parts of the apparatus are as follows. The connecting tube $D$ is 9 to 11 mm in internal diameter. The trap is 235 to 240 mm in length. The condenser, if of the straight-tube type, is approximately 400 mm in length and not less than 8 mm in bore diameter. The receiving tube $E$ has a $5-\mathrm{mL}$ capacity, and its cylindrical portion, 146 to 156 mm in length, is graduated in $0.1-\mathrm{mL}$ subdivisions, so that the error of reading is not greater than 0.05 mL for any indicated volume. The source of heat is preferably an electric heater with rheostat control or an oil bath. The upper portion of the flask and the connecting tube may be insulated.

Clean the receiving tube and the condenser with chromic acid cleansing mixture, thoroughly rinse with water, and dry in an oven. Prepare the toluene to be used by first shaking with a small quantity of water, separating the excess water, and distilling the toluene.
Procedure-Place in the dry flask a quantity of the substance, weighed accurately to the nearest centigram, which is expected to yield 2 to 4 mL of water. If the substance is of a pasty character,
of water adhere to the walls of the receiving tube, scrub them wire and wetted with toluene. When the water and toluene separated completely, read the volume of water, and calcula percentage that was present in the substance.

## METHOD III (GRAVIMETRIC)

Procedure for Chemicals-Proceed as directed in the indi monograph preparing the chemical as directed under Loss on ing (731).
Procedure for Biologics-Proceed as directed in the indi monograph.
Procedure for Vegetable Drugs--Place about 10 g of the prepared as directed (see Vegetable Drugs-Methods of An (561)) and accurately weighed, in a tared evaporating dish. I $105^{\circ}$ for 5 hours, and weigh. Continue the drying and weigh 1-hour intervals until the difference between two successive y ings corresponds to not more than $0.25 \%$.

## 〈941〉 X-RAY DIFFRACTION

Every crystal form of a compound produces its own charact X-ray diffraction pattern. These diffraction patterns can be de either from a single crystal or from a powdered specimen (co ing numerous crystals) of the material. The spacings betwee the relative intensities of the diffracted maxima can be use qualitative and quantitative analysis of crystalline materials. P diffraction techniques are most commonly employed for ro identification and the determination of relative purity of cryst materials. Small amounts of impurity, however, are not nor detectable by the X-ray diffraction method, and for quant measurements it is necessary to prepare the sample carefu avoid preferred orientation effects.
The powder methods provide an advantage over other mea analysis in that they are usually nondestructive in nature (spe preparation is usually limited to grinding to ensure a randoml ented sample, and deleterious effects of X -rays on solid ph ceutical compounds are not commonly encountered). The pri use of single-crystal diffraction data is for the determination o ecular weights and analysis of crystal structures at the atomic However, diffraction established for a single crystal can be us support a specific powder pattern as being truly representativ single phase.
Solids-A solid substance can be classified as being cryst noncrystalline, or a mixture of the two forms. In crystalline rials, the molecular or atomic species are ordered in a three-d sional array, called a lattice, within the solid particles. This or of molecular components is lacking in noncrystalline material. crystalline solids sometimes are referred to as glasses or amor solids when repetitive order is nonexistent in all three dimen It is also possible for order to exist in only one or two dimen resulting in mesomorphic phases (liquid crystals). Although talline materials are usually considered to have well-defined $v$ external morphologies (their habits), this is not a necessity $f$ ray diffraction analysis.
The relatively random arrangement of molecules in noncr line substances makes them poor coherent scatterers of X-ray sulting in broad, diffuse maxima in diffraction patterns. The
acteristic X-ray patterins, so does every solvate. Sometimes the differences in the diffraction patterns of different polymorphs are relatively minor, and must be very carefully evaluated before a definitive conclusion is reached. In some instances, these polymorphs and or solvates show varying dissolution rates. Therefore, on the time scale of pharmaceutical bioavailability, different total amounts of drug are dissolved, resulting in potential bioinequivalence of the several forms of the drug.
Fundamental Principles-A collimated beam of monochromatic X-rays is diffracted in various directions when it impinges upon a rotating crystal or randomly oriented powdered crystal. The crystal acts as a three-dimensional diffraction grating to this radiation. This phenomenon is described by Bragg's law, which states that diffraction (constructive interference) can occur only when waves that are scattered from different regions of the crystal, in a specific direction, travel distances differing by integral numbers ( $n$ ) of the wayelength $(\lambda)$. Under such circumstances, the waves are in phase. This condition is described by the Bragg equation:

$$
\frac{n \lambda}{2 \sin \theta}=d_{u k l},
$$

in which $d_{h k}$ denotes the interplanar spacings and $\theta$ is the angle of diffraction.

A family of planes in space can be indexed by three whole numbers, usually referred to as Miller indices. These indices are the reciprocals, reduced to smallest integers, of the intercepts that a plane makes along the axes corresponding to three nonparallel edges of the unit cell (basic crystallographic unit). The unit cell dimensions ate given by the lengths of the spacings along the three axes, $a, b, c$, and the angles between them, $\alpha, \beta$, and $\gamma$. The interplanar spacing for a specific set of parallel planes $h k l$ is denoted by $d_{h k l}$. Each such family of planes may show higher orders of diffraction where the $d$ values for the related families of planes $n h, n k, n l$ are diminished by the factor $1 / n$ ( $n$ being an integer: $2,3,4$, etc.). Every set of planes throughout a crystal has a corresponding Bragg diffraction angle associated with it (for a specific $\lambda$ )
The amplitude of a diffracted X-ray beam from any set of planes is dependent upon the following atomic properties of the crystal: (1) position of each atom in the unit cell; (2) the respective atomic scattering factors; and (3) the individual thermal motions. Other factors that directly influence the intensities of the diffracted beam are: (1) the intensity and wavelength of the incident radiation; (2) the volume of crystalline specimen; (3) the absorption of the Xradiation by the specimen; and (4) the experimental arrangement utilized to record the intensity data. Thus, the experimental conditions are especially important for measurement of diffraction intensities.

Only a limited numbet of Bragg planes are in a position to diffract when monochromatized X-rays pass through a single crystal. Techniques of recording the intensities of all of the possible diffracting $h k l$ planes involve motion of the single crystal and the recording media. Recording of these data is accomplished by photographic techniques (film) or with radiation detectors
A beam passing through a very large number of small, randomly oriented crystals produces continuous cones of diffracted rays from each set of lattice planes. Each cone corresponds to the diffraction from various planes having a similar interplanar spacing. The intensities of these Bragg reflections are recorded by either film or radiation detectors. The Bragg angle can be measured easily from a film, but the advent of radiation detectors has made possible the construction of diffractometers that read this angle directly. The intensities and $d$ spacings are more conveniently determined with powder diffractometers employing radiation detectors than by film


Several specialized handing techniques may be employed to Sinimize prefered orientation, but further reduction of particle size s often the best approach.
Where very accurate measurement of the Bragg angles is necessary, a small amount of an intemal standard can be mixed into the specimen. This enables the film or recorder tracing to be calibrated. If comparisons to literature values (including compendial limits) of $d$ are being made, calibrate the diffractometer. NIST stan dards are available covering to a $d$-value of 0.998 nm . Tetradecano may be used ( $d$ is 3.963 nm ) for larger spacing.
The absorption of the radiation by any specimen is determined by the number and kinds of atoms through which the X-ray beam passes. An organic matrix usually absorbs less of the diffracted radiation than does an inorganic matrix. Therefore, it is important in quantitative studies that standard curves relating amount of material to the intensity of certain $d$ spacings for that substance be
${ }^{\text {' Brindley, GW and Brown, G, eds., Crystal Structures of Clay }}$ Minerals and their X-ray Identification, Mineralogical Society Monograph No. 5, London, 1980, pp. 318 ff
(ratio of the peak intensity of a particular $d$ spacing to the int of the strongest maxima in the diffraction pattern) and the $d$ s are used in the comparison. If a reference material (e.g., US erence Standard) is available, it is preferable to generate a pr reference pattern on the same equipment used for running tt known sample, and under the same conditions. For most o crystals, it is appropriate to record the diffraction pattern to ir values for $2 \theta$ that range from as near zero degrees as possi 40 degrees. Agreement between sample and reference sho within the calibrated precision of the diffractometer for diffr angle ( $2 \theta$ values should typically be reproducible to $\pm 0.10$ o degrees), while relative intensities between sample and ref may vary considerably. For other types of samples (e.g., ino salts), it may be necessary to extend the $2 \theta$ region scanned t beyond 40 degrees. It is generally sufficient to scan past t strongest reflections identified in the Powder Diffraction Fil
${ }^{2}$ The International Centre for Diffraction Data, Newtown S Corporate Campus, 12 Campus Boulevard, Newtown Squar 19073, maintains a file on more than 60,000 crystaline mat both organic and inorganic, suitable for such comparisons.
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