X-ray Microtomography of Solid Dosage Forms

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X-ray microtomography has great potential for improving the understanding of the structural features of solid dosage forms and the changes in those features during manufacturing, handling, and storage. This article describes the basic principles of the technique and provides examples of its potential applications.

Pharmaceutical scientists have long sought the ability to see inside the solid dosage forms they produce to determine their products' structural features and to better understand their mode of action. Previous studies have used various techniques for visualizing the internal structure of solid dosage forms, including 1H NMR imaging (1), confocal microscopy (2), and conventional microscopy (optical and electron) combined with mechanical slicing of samples (i.e., microtoming) (3). One drawback of several current techniques is their invasive nature that can destroy the sample and prevent any further testing. Another is the techniques' limited penetration and resolution. Thus, it is probably fair to say that the ideal experimental approach for the three-dimensional structural imaging of pharmaceutical dosage forms has not yet been realized.

X-ray microtomography is a relatively new approach to imaging the internal structure of solid dosage forms. This technique has been widely used for the in vivo imaging of plants, insects, animals, and humans. X-ray microtomography is a non-destructive technique that has a high penetration ability and provides a reasonable level of resolution (~5–20 μm).

**Principles of X-ray microtomography**

The X-ray microtomography approach used in this work is an extension of the computer aided tomography (CAT) medical imaging technique commonly used in hospitals. X-rays are directed from a high-power source toward a sample, and a detector on the opposite side of the sample measures the intensity of the transmitted X-rays (see Figure 1). A two-dimensional "shadow" image is produced by accurately rastering the X-ray beam across the sample. The sample then is carefully moved (usually rotated) relative to the X-ray beam, and the process is repeated to produce additional two-dimensional images from various view points. Using a sophisticated Fourier transform algorithm, the two-dimensional images then are combined to generate a complete three-dimensional map of the sample.

The intensity of the X-rays reaching the detector is controlled by the sample path length and the X-ray attenuation coefficient of the material that it encounters on that path (4). The longer the path length and the greater the attenuation coefficient of the material (see http://physics.nist.gov/PhysRefData/XrayMassCoef/tab3.html), the greater the number of diffraction and scattering
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Cross section of a fast-dissolving tablet concluded that it was especially useful for sodium, chlorine, or iron) will generally create the X-ray microtomography can be used to measure the location, diameter, and depth) of the hole/hole in the coating from which the active ingredient will be expelled (see Figure 5) (3). Likewise, in a liquid-filled soft-gelatin capsule the thickness and integrity of the capsule’s solid shell and its welded seam can be readily determined (see Figure 6).

Applications

Two- and three-dimensional images of a wide range of solid dosage forms were acquired in this study and used to learn more about their function and structure. The following provides several examples of how such information can be used to aid in the design and testing of pharmaceutical tablets and in solving related technical problems.

Elucidation of structural features. Tablet shape and dimensions. X-ray microtomography can provide simple information such as the shape and size of the regions of a dosage form that are not readily accessible. For example, the technique is ideally suited for determining the thickness of layers in multilayer tablets and for determining the shape and size of the interface between these layers (see Figure 2). It also can be used to elucidate the microstructure of fast-dissolving tablets such as those manufactured using lyophilization in which the structural features reflect the size and shape of the ice crystals that were present in the tablet before drying (see Figure 3). Reports in the literature also point to the utility of this technique for assessing the morphology and pore-size distribution of pharmaceutical granules (5, 6). Farber et al. concluded that it was especially useful for studying the connectivity and shape of voids within granules made using wet granulation (5).

Coatings. X-ray microtomography can be used to measure the thickness of a functional coating (see Figure 4) and to assess variations in the film-coating thickness at the corners of the tablet to ensure the integrity of the coating is maintained over the entire surface of the dosage form. For an osmotic controlled-release dosage form it is quite simple to determine the key dimensions (e.g., location, diameter, and depth) of the hole in the coating from which the active ingredient will be expelled (see Figure 5) (3). Likewise, in a liquid-filled soft-gelatin capsule the thickness and integrity of the capsule’s solid shell and its welded seam can be readily determined (see Figure 6).

Figure 2: Conventional bilayer tablet structure.

Figure 3: Cross section of a fast-dissolving tablet manufactured by lyophilization.

Figure 4: Gel-coated analgesic tablet structure.
Embossing details, defects, damage. Embossed markings on a tablet’s surface are quite easily resolved and measured using X-ray microtomography (see Figure 7). This capability could be used to help troubleshoot tablet manufacturing problems such as sticking or picking. Defects in or damage to the internal structure of a compacted tablet also can be detected. Researchers have used X-ray microtomography to study small defects and hidden damage in a range of nonpharmaceutical materials, including metals and composites (7, 8). Figure 8 shows a region of an immediate-release tablet with an internal crack that might significantly influence its structural integrity. Although it is not currently possible to enable 100% inspection of all dosage forms from a large-scale batch, this may be feasible in the future as the instrumentation’s speed increases and accuracy improves.

Distribution of components. If the material of interest generates sufficient X-ray contrast, then it is feasible to determine the distribution of an active ingredient or functional excipient in a tablet or capsule using X-ray microtomography. This may provide vital information about the performance of a novel manufacturing process or, in the future, may enable at-line process monitoring to occur. The distribution of particles (citric acid) in a fast-dissolve tablet is shown in Figure 9, and the distribution of an inorganic excipient incorporated into an osmotic controlled-release tablet is shown in Figure 10. This approach can provide complementary information to spectroscopic chemical images (9). With the rigorous use of three-dimensional image analysis tools, quantitative information about the uniformity of pharmaceutical samples can be obtained.

Density distribution determinations
The intensity of the gray scale that appears in the X-ray images is partly a function of the sample’s density. The density of standard uni-axially compressed tablets is not uniform (1, 10), and X-ray microtomography may provide a means to study and quantify this density inhomogeneity in tablets. Density differences have been detected in immediate-release tablets (see Figure 11). By using standards of known density, researchers interested in the density of bone samples quantitatively computed this information directly from X-ray microtomography data (11). A similar approach with other materials also has been successful (12, 13). Quantitative studies of this type have been reported for pharmaceutical compacts. One study showed that the tablet shape and the presence of embossing resulted in nonuniformity in the density of compacts made from microcrystalline cellulose (14). In addition, researchers showed that die lubrication and the mode of compression each influenced the degree of this nonuniformity. In another study, X-ray microtomography of several compacted excipients showed how density variations in various regions of a compact control the mechanical response of the sample during tensile failure and indentation hardness tests (15). Researchers also demonstrated in this work that the degree of structural and mechanical anisotropy in a compact is significantly less for highly deformable (plastic) materials.

Foreign matter detection
A unique application of X-ray microtomography is the detection and location of foreign matter such as a metal particle within a solid dosage form. Such atypical substances show up very clearly as being different from the materials commonly used to manufacture pharmaceutical tablets. A metal particle within a tablet creates a hot spot in the tomographic image that shows it is unmistakably foreign (see Figure 12). This informa-
tion enables the location of the particle to be exactly pinpointed even though it is entirely contained within the dosage form and its size is quite small (~50 μm width in this example).

**Counterfeit product detection**

Counterfeit tablets can be difficult to distinguish from authentic dosage forms on the basis of external appearances. By using X-ray microtomography, however, it is possible to compare the internal structures and dimensions of two tablets without destroying the samples, which may be important if the tablet is to be used as evidence in patent litigation or other legal proceedings. Figure 13 shows the images of two tablets, an innovator’s patented dosage form and an unofficial copy, in which clear differences in the internal structures (e.g., layered core structure) can be seen easily.

**Future applications**

**Compaction studies.** Several researchers have used X-ray microtomography to study changes in the structure of porous composites and foams under applied external stresses (16–18). It is implicit from these studies that the technique also has potential use in research into powder compaction and the formation of compressed tablet dosage forms (12, 17). It is likely that the time-dependent structural information obtained from such studies would provide considerable insight into the pharmaceutical manufacturing processes of tablet manufacture and dry granulation. Current measurement systems cannot collect data rapidly enough to enable powder compaction to be studied under current pharmaceutical manufacturing conditions, nor do they enable the behavior of the smallest individual particles to be resolved. The response time of research instruments is improving rapidly, however, and the need for studying short-lived events and even smaller samples has been recognized by commercial instrument designers and manufacturers. Using synchrotron X-ray sources can provide some help in this regard (see http://www.esrf.fr/Industry/Applications/Tomography/), and enhanced X-ray sources, detector elements, control systems, and analysis software will likely lead to significant improvements in both response time and resolution for commercial instruments within the next 5–10 years.

**Chemical mapping.** Recent developments in instrumentation suggest that in the future, X-ray microtomography studies may provide information about the chemical nature of samples (9, 19). This is an exciting advance that could address one of the current limitations of the technique for fully characterizing pharmaceutical products (i.e., a lack of information about the chemical nature of the sample). Commercial instruments capable of this type of multifaceted analysis are still a long way away, but systems in which X-ray microtomography is combined with more conventional spectroscopic mapping techniques in a single integrated instrument are already under development. Currently these systems require the destruction of the sample to facilitate the spectroscopic mapping of the samples, but they do have the potential for providing a detailed three-dimensional map of the physical and chemical nature of a product based on the analysis of just one or two dosage forms. This is clearly a significant improvement over the capabilities of current nonintegrated systems that each require a comparable number of dosage units to complete a full analysis.

**Conclusion**

X-ray microtomography can obtain useful qualitative and quantitative information about the structure of pharmaceutical dosage forms. In the future, it is likely that X-ray microtomography also will be able to provide chemical mapping capabilities and improved spatial resolution of small structural features within these types of samples (9, 19, 20).

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References


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