Identification of thorium dioxide in human liver cells by electron microscopic x-ray microanalysis

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SUMMARY Thirty-two years after injection of thorium dioxide (Thorotrast) for diagnostic x-ray studies in a female patient deposits were found by light microscopy in the liver macrophages (Kupffer cells). They were shown by electron microscopy to be located inside secondary lysosomes, and by autoradiography and x-ray microanalysis they were identified as thorium.

The aim of the present study was to undertake an ultrastructural examination of liver cells obtained by biopsy from a 47-year-old woman who in 1945 had received an unknown amount of thorium dioxide (Thorotrast) intravenously for diagnostic x-ray purposes before the removal of a brain tumour. The deposits of Thorotrast in the liver and spleen were discovered incidentally during x-ray examination of the kidney for an intercurrent disease. The biopsy was taken specifically for morphological study of the liver. The intracellular presence and localisation of aggregates of thorium dioxide particles was studied using light microscopic autoradiography and electron microscopic x-ray microanalysis. The latter technique has proved a valuable tool for the identification of elements naturally occurring in biological samples as well as in exogenously introduced elements. To our knowledge, this is the first study demonstrating thorium dioxide (Thorotrast) in liver cells by x-ray microanalysis.

Material and methods

Tissue from needle biopsy of the liver was fixed immediately in 4% glutaraldehyde in 0.1 M cacodylate buffer at pH 7-2. One-half of the tissue was prepared for electron microscopy while the other was transferred to buffered formalin and processed by routine histological methods.

Paraffin-embedded sections, 5 μm thick, were deparaffinised, dehydrated, and coated with Kodak nuclear track emulsion NTB 2 for autoradiography. The emulsion-coated slides were stored in darkness at 4°C. After four weeks they were developed and stained with haematoxylin and eosin.

The material for electron microscopic studies was transferred to a mixture with a final concentration of 2% glutaraldehyde and 2% osmium tetroxide buffered to pH 7-2 by 0.1 M cacodylate. After one hour the tissue was washed in buffer and postosmification was carried out in 2% OsO4 for one hour (Franke et al., 1969). Subsequently the tissue was dehydrated and embedded in Epon 812. All substances used for preparation were of high chemical purity with an insignificant amount of thorium. Sections approximately 60 nm thick, judged from interference colours, were cut with glass knives, mounted on copper grids, and examined in a Zeiss EM-9S-2 electron microscope at 60 keV. Some sections were examined without prior staining, others after staining with uranyl acetate and lead citrate (Reynolds, 1963). For electron microscopic x-ray microanalysis unstained sections, about 150 nm thick, were mounted on copper grids and analysed in a Transmission Electron Microscope (Phillips 300) equipped with an energy dispersive x-ray analysis system (EDAX). The electron microscope was operated at 100 kV with a specimen tilt of 42 degrees. Analysing time was 55 seconds.

Results

MORPHOLOGICAL FINDINGS

Light microscopy (Fig. 1)
The hepatic architecture was preserved. A variable amount of granular material was observed in the sinusoidal macrophages (Kupffer cells).

Electron microscopy (Figs 2 and 3)
In the ultrastructural study one or more electron-dense cytoplasmic inclusions were found within each liver macrophage. The study of 60 nm thick stained sections revealed that the inclusions lay...
inside single-membrane bound cell organelles, identified as residual bodies (Fig. 2). Some of these bodies were completely filled with the dense material. In some instances electron-dense inclusions could be demonstrated inside autophagic vacuoles (Fig. 3).

**Electron microscopic x-ray microanalysis (Figs 4 and 5)**

In the semithick unstained sections prepared for x-ray microanalysis little cellular detail and no membranes could be identified. However, the electron dense inclusions were easily located (Fig. 4). Figure 5 shows the x-ray spectrum from microanalysis of the heavily loaded cells shown in Figure 4. The spectrum shows prominent thorium peaks (M-line at 3.06, Lxα at 12.96, Lxβ1 at 16.19, Lxβ2 at 15.62, and Lxγ at 18.92 keV, respectively). None of these peaks was present in the spectrum from an adjacent cell without electron-dense inclusions. The other peaks present in Fig. 5 represent elements (P, Cl, Cr, Ni, Cu, Zn, Os) in the fixative, embedding material, and metal-containing parts of the ana-
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Fig. 3 Electron micrograph showing autophagic vacuole with heterogeneous granules. Liver macrophage. Uranyl acetate and lead citrate × 14 000

Fig. 4 Semithick section (150 nm) of liver macrophage heavily loaded with inclusions of Thorotrast. Unstained × 14 000

lysing instrument. These peaks were present in all the x-ray spectra.

Autoradiography disclosed some short, straight tracks emanating in a spokelike fashion from the thorium inclusions, typical of those produced by alpha particles.

The patient's present condition is excellent, without signs of liver disease.

Discussion

The ultimate fate of phagocytosed material depends not only on the function of the lysosomal apparatus but also on the nature of the ingested material. Digestible organic substances are broken down and may enter the metabolism as simpler compounds. On the other hand, phagocytosed material resistant to the lysosomal enzymes is stored indefinitely in residual bodies. Depending on the quantity and type, such material may be injurious. An example of a potentially hazardous material is radioactive Thorotrast particles. Thorotrast, a colloidal solution of thorium dioxide, was used widely some 30 years ago as a contrast medium for diagnostic X-ray purposes. Thorotrast contains the radioactive isotope, thorium-232, which is a nearly pure alpha
particle emitter. Its biological half-life has been calculated to be over 400 years (Hurst et al., 1957).

There is a voluminous literature on Thorotrast and the liver. The induction of liver cancer in individuals with Thorotrast deposits has gained particular attention (Smoron and Battifora, 1972; Kiely et al., 1973).

In these studies the presence of Thorotrast has been proved by light microscopic autoradiography and by morphological studies with the electron microscope. However, only analytical electron microscopy can determine in situ the exact elementary composition of the deposits.

In this study we identify and demonstrate the sub-cellular localisation of Thorotrast particles injected more than 30 years ago. The Thorotrast particles were mainly located in liver macrophages (Kupffer cells). The liver tissue appeared otherwise undamaged. Electron microscopic examination showed that the particles were retained inside residual bodies. With the analytical electron microscope it was possible to identify thorium-containing deposits.

A major limiting factor in x-ray microanalysis is the possible loss and redistribution of elements during conventional preparation techniques (Morgan et al., 1975). Fixation and embedding of the tissue may further introduce new elements into the substrate to be analysed. Nevertheless, when firmly bound elements are investigated, the routine preparation for transmission electron microscopy may be adequate (Yarom et al., 1976). As shown in this study, thorium deposits in the liver are easily detected by this method.

References


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J Clin Pathol 1978 31: 893-896
doi: 10.1136/jcp.31.9.893

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