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FREEDOM OF INFORMATION ACT REQUEST

October 12, 1984

Mr. J.M. Felton, Director
Division of Rules and Records
Office of Administration
United States Nuclear Regulatory Commission
7735 Old Georgetown Road
Bethesda, Maryland

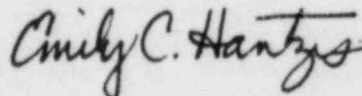
FREEDOM OF INFORMATION
ACT REQUEST
FOIA-84-801
Rec'd 10-15-84

Dear Mr. Felton,

I am interested in obtaining a copy of the original and subsequent renewal license applications and supporting documentation filed by the Medical Products Division of 3M, St. Paul, Minn., or the Lawrence Soft Ray Corporation, San Jose, Calif., concerning an implantable medical device known as "I-125 seeds". These "seeds" consist of a welded titanium capsule containing iodine-125 adsorbed onto a silver rod or onto two anion exchange resin spheres. They are intended for interstitial radiation treatment of tumors. In addition, I am interested in any staff review documentation related to this device.

I have spoken with David Meyer about this. Thank you in advance for any information your office can provide.

Very truly yours,



Emily C. Hantzes

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PDR FOIA
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