## **DEPARTMENT OF HEALTH AND HUMAN SERVICES** PUBLIC HEALTH SERVICE

FOOD AND DRUG ADMINISTRATION

## **DECLARATION FOR IMPORTED**

Form Approved OMB No. 0910-0025 Expiration Date: 10/31/2000

INSTRUCTIONS

- 1. If submitting entries electronically through ACS/ABI, hold FDA-2877 in entry file. Do not submit to FDA unless requested.
- 2. If submitting paper entry documents, submit the following to FDA:

RADIATION CONTROL STANDARDS		a. 2 copies of customs Entry Point (e.g. CF 3461, CF 3461 Att, CF 7501, etc.) b. 1 copy of FDA 2877 c. Commercial Invoice(s) in English.	
U.S. CUSTOMS PORT OF ENTRY		ENTRY NUMBER	DATE OF ENTRY
NAME & ADDRESS OF MANUFACTURING SITE; COUNTRY OF ORIGIN		NAME & ADDRESS OF IMPORTER & ULTIMATE CONSIGNEE (if not importer)	
PRODUCT DESCRIPTION	QUANTITY (Items/Containers)	MODEL NUMBER(S) & BRAN	D NAME(S)
DECLARATION: I / WE DECLARE THAT TI	HE PRODUCTS IDENTIFIED A	BOVE: (Mark applic	able statements, fill in blanks, & sign)
$\ \square$ A. ARE NOT SUBJECT TO RADIATI	ON PERFORMANCE STAN	IDARDS BECAUSE THEY:	
1. Were manufactured prior to the effect     2. Are excluded by the applicability claused by the applicability clauses are specify reason for exclusion			·
destroyed, or held for future testing ( 7. Are being reprocessed in accordance or transferred without FDA approval.	o be used in manufacturing or as product development by the imp i.e., not distributed). (Quantities with P.L. 104-134 or other FDA	replacement parts (NOT APPL porting firm, are labeled "FOR T Limited - see reverse.) a guidance, are labeled "FOR EX	ICABLE to diagnostic x-ray parts). 'EST/EVALUATION ONLY," and will be exported,  XPORT ONLY," and will not be sold, distributed,
☐ B. COMPLY WITH THE PERFORMA CERTIFICATION LABEL OR TAG ☐ 1. Last annual report or Product/Initial r	TO THIS EFFECT IS AFFIX		COMPLIANCE DOCUMENTED IN:
ACCESSION NUMBER of Report  Name of MANUFACTURER OF RECORD (Filed report with FDA/CDRH)  2. Unknown manufacturer or report number; State reason:			
	RCE; WILL BE USED UNDE STOMS SUPERVISION WH Training (attach Form FDA 766)	R A RADIATION PROTEC	A TEMPORARY IMPORT BOND; WILL NOT ETION PLAN; AND WILL BE DESTROYED SSION IS COMPLETE:
□ D. DO NOT COMPLY WITH PERFORM INTRODUCED INTO COMMERCE INTO COMPLIANCE IN ACCORD. □ 1. Approved Petition is attached.	UNTIL NOTIFICATION IS	RECEIVED FROM FDA TH ROVED PETITION. <i>(See F</i>	HAT PRODUCTS HAVE BEEN BROUGHT
WARNING: Any person who knowingly declaration may be fined not more thimprisoned not more than 5 years or both,	makes a false SIGNATURE tan \$10,000 or pursuant to Title	OF IMPORTER OF RECORD	
18 U.S.C. 1001. Any person importing a electronic product may also be subject to \$1000 per violation, up to a maximum \$300 violations pursuant to Title 21 U.S.C. 360pp.	civil penalties of	ITLE OF RESPONSIBLE PERSON	
Public reporting burden for this collection of			

this burden estimate or any other aspect of this collection of information, including suggestions for reducing this burden to:

Food and Drug Administration CDRH (HFZ-342) 2094 Gaither Road Rockville, MD 20850

An agency may not conduct or sponsor, and a person is not required to respond to, a collection of information unless it display currently valid OMB control number.