DEPARTMENT OF HEALTH AND HUMAN SERVICES

FOOD AND DRUG ADMINISTRATION

Form Approved: OMB No. 0910-0025 Expiration Date: August 31, 2023 See Reverse for PRA statement

Distribution List: Purchaser		REPORT OF ASSEMBLY				Assembler/Purchaser Control Number					
Assembler State Radiation Health Office	OF A	OF A DIAGNOSTIC X-RAY SYST								←	
1. EQUIPMENT LOCATION				2. ASSEMBLER INFORMATION							
a. NAME OF HOSPITAL, DOCTOR OR OFFICE WHERE INSTALLED				a. COMPANY NAME							
a. NAME OF HOSPITAL SOCION ON OF THE WILLIE HISTALLED				a. COMPANT NAME							
b. STREET ADDRESS				b. STREET ADDRESS							
c.CITY		d. STATE	c. CITY						d. STATE		
e. ZIP CODE f. TELEPHONE NUMBER			e. ZIP CODE					f. TELEPHONE NUMBER			
3. GENERAL INFORMATION	N										
a. THIS REPORT IS FOR ASSEMBLY OF CERTIFIED CO	DMPONENTS WHICH ARE (Check approp	riate box(es))									
NEW ACCURACY TO AN ACCURACY AND				REASSEMBLY - MIXED SYSTEM (Both certified and non-certified components)							
NEW ASSEMBLY - FULLY CERTIFIED SYSTEM					REPLACEMENT COMPONENTS IN AN EXISTING SYSTEM						
b. INTENDED USE(S) (Check appropriate box(s))	YSTEM				AN ADDITION	I TO AN EXISTI	NG SYSTEM				
GENERAL PURPOSE RADIOGRAPHY UROLOGY					WHOLE BODY SCANNER			RADIATION THERAPY SIMULATOR OTHER			
GENERAL PURPOSE FLUOROSCOP	\vdash	MAMMOGRAPHY			 HEAD-NECK (<i>Medical</i>)			C-ARM FLUOROSCOPIC (Specify in comments)			
TOMOGRAPHY (Other than CT)	CHEST			NTAL-INTRAORAL			DIGITA				
ANGIOGRAPHY	CHIROP	L RACTIC -	_	NTAL-CEPHALOMETRIC			\vdash	MINERAL ANALYSIS			
		DSCANNER	=	NTAL PANORAMIC			DENTA				
c. THE X-RAY SYSTEM IS (Check one)	d. THE MASTER CONTROL IS IN RO					e. DATE OF A					
STATIONARY											
MOBILE											
							(mm	****			
4. COMPONENT INFORMA this Form Number, and co			sectio	n use a	inother i	orm, rep	placing th	ne preprinted nu	mber v	with	
a. THE MASTER CONTROL IS b. CONTROL MANUFACTURER			d. CONTROL SERIAL NUMBER					e. DATE MANUFAC	TURED		
A NEW INSTALLATION											
EXISTING (Certified)	EXISTING (Certified) c. CONTROL MODEL NUMBER			f. SYSTEM MODEL NAME (CT Systems Onl							
EXISTING (Non-certified)											
Complete the following information for the For other certified components, enter in the				ting devic	es, tables a	nd CT gantri	es enter the	manufacturer and Mod	lel numbe	er in the indicated spaces.	
g. SELECTED COMPONENTS				h. OTHER CERTIFIED C (Enter number of each installed							
MANUFACTURER MODEL NUMBER		DATE MAI	DATE MANUFACTURED								
							X-RAY CONTROL			CRADLE	
MANUFACTURER	MODEL NUMBER	DATE MAI	DATE MANUFACTURED				HIGH VOLTAGE GENERATOR			FILM CHANGER	
							VERTICAL CASSETTE HOLDER			IMAGE INTENSIFIER	
MANUFACTURER	MODEL NUMBER	DATE MAI	DATE MANUFACTURED			TUBE HOUSING ASSEMBLY			INFOCE INTENSITIES		
							DENTAL TUBE HEAD			SPOT FILM DEVICE	
MANUFACTURER	MODEL NUMBER	DATE MAI	DATE MANUFACTURED						FLUOROSCOPIC IMAGING ASSEMBLY		
					CEPHALOMETRIC DEVICE			IMAGE RECEPTOR			
MANUFACTURER MODEL NUMBER		DATE MANUFACTUR		ED I		IMAGE RECEPTOR SUPPORT DEVICE					
										FLUOROSCOPIC AIR KERMA DISPLAY DEVICE	
5. ASSEMBLER CERTIFICAT	TION										
I affirm that all certified components asser	mbled or installed by me for whi	ch this report is being made v	were adio	isted and	tested by m	e according	to the instri	ictions provided by the	manufac	ture(s), were of the type	
required by the manufacturer(s), were of t	he type required by the diagnos	ic x-ray performance standard	d (21 CFR	R Part 1020), were not	modified to	adversely af	fect performance, and	were insta	alled in accordance with	
provisions of 21 CFR Part 1020. I also affirm completion of the assembly, a copy of this									iu, within	adys following	
a. PRINTED NAME b. SIGNA			RE					<u> </u>			

6. COMMENTS

This section applies only to requirements of the Paperwork Reduction Act of 1995. *DO NOT SEND YOUR COMPLETED FORM TO THE PRA STAFF EMAIL ADDRESS BELOW.*

The burden time for this collection of information is estimated to average 18 minutes per response, including the time to review instructions, search existing data sources, gather and maintain the data needed and complete and review the collection of information. Send comments regarding this burden estimate or any other aspect of this information collection, including suggestions for reducing this burden, to:

> Department of Health and Human Services Food and Drug Administration Office of Chief Information Officer Paperwork Reduction Act (PRA) Staff PRAStaff@fda.hhs.gov

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