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Management of Medical Equipment Inspection Checklist- Random

Name of the Facility:		_	
Date of Inspection:	/	_/_	

Ref.	Description	Yes	No	N/A	Remarks			
5	STANDARD ONE: HEALTH FACILITIES DOCUMENTED PROGRAM/PLAN FOR MEDICAL EQUIPMENT							
5.1.	Health facilities shall cover the entire range of medical equipment installed at the facility as well as its affiliate services (such as homecare, ambulances etc.).							
6	STANDARD TWO: NEW MEDICAL EQUIPMENT ARRIVE NONFACILITY EQUIPMENT)	ES TO THE	FACILITY	(PURCHA	ASED/			
6.1.	All medical equipment in the health facilities should be registered in Ministry of Health and Prevention (MOHAP).							
6.3.	For new purchase, all installation documents should be available with supply chain document to ensure the safety of the supply chain and therefore, protecting patients and staff from unstable, contaminated, defective, and counterfeit supplies. Documents include but are not limited to:							
6.3.1.	Purchase order.							
6.3.2.	Country of origin.							
6.3.3.	Installation report							
6.3.4.	PPM schedule.							
6.3.5.	Warranty certificate.							
6.3.7.	Delivery note.							

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6.3.9.	User training certificates or attendance sheet document.			
	All medical equipment (purchased, demonstration,			
6.5.	loaned etc.) should be tested as applicable for safety			
0.5.	(electrical safety test, mechanical, radiation etc.), QC and			
	calibration is done upon installation.			
	Availability of operating manual accessible by clinical			
6.6.	staff of the department and service manuals should be			
	available for all inhouse service equipment.			
6.8.	To eliminate the use of extension cords			
6.9.	For non-Facility medical equipment			
0.9.	(DEMO/LOANED/PATIENT PROPERTY):			
6.9.1.	Should be tagged with DEMO/LOANED/PATIENT			
6.9.1.	PROPERTY stickers.			
	A file containing all non-facility equipment details and			
	checklist shall be checked and verified by Biomedical			
6.9.2.	Engineering, insuring they are properly checked and are			
	safe for all patients, staff, and visitors, prior to use in the			
	health facility.			
	For the radiology equipment with radiation, the facility			
6.10.	must ensure that they have the required credentials and			
0.10.	licenses from FANR for all equipment producing			
	radiation and falling in the scope of FANR licensing.			
7	STANDARD THREE: MEDICAL EQUIPMENT INVENTOR	Y MANAG	EMENT	
	Maintaining proper inventory through inventory system			
7.1.	for all the medical equipment available and to be			
7.1.	updated every time a new equipment arrives or removed			
	from service			
	Critical equipment is identified in the inventory, there is			
7.2.	provision for back-up/ alternative for critical equipment			
	during their failure or maintenance.			

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lical equipment should be labelled and tagged
omedical asset number.
ails of medical equipment (manufacturer/ model/
umber/ local supplier/ date of purchase and
n at the facility) should be available.
DARD FOUR: CORRECTIVE MAINTENANCE
ility of biomedical workshop and biomedical
er is mandatory at general hospitals.
cility should tag and remove any defective
ent from use.
DARD FIVE: PLANNED PREVENTATIVE MAINTENANCE OF MEDICAL EQUIPMENT
hedule is available for all medical equipment
needs PPM.
to ensure that PPM is performed according to
se and as per recommendations of their
acturers. The PPM period should adhere at least
nufacturer recommendation or better.
ickers should be available physically on the
l equipment and matching the PPM date on
l equipment management system in the facility
the PPM check list.
DARD SIX: MEDICAL EQUIPMENT RISK MANAGEMENT
's process should follow the below steps to
e a medical device recall:
the availability of the equipment in the facility.
nent recall record is to be prepared and updated
у.
nent to be removed from the use and informed to
oplier/manufacturer.
Tan Di C b ee io m C so r y is fa a it C y go t m ri m

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	equipment file.							
	' '							
10.5.5.	Necessary repair/ren	placement should be done.						

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