



DEPARTMENT OF HEALTH
Republic of South Africa

REQUIREMENTS FOR LICENCE HOLDERS WITH RESPECT TO QUALITY CONTROL TESTS FOR DIAGNOSTIC X-RAY IMAGING SYSTEMS

**DEPARTMENT OF HEALTH
DIRECTORATE: RADIATION CONTROL**

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I. GENERAL REQUIREMENTS

A. THE LICENCE HOLDER SHALL:

1. Display the product licence number (see list of licences from Department of Health (DoH)) on equipment.
 - 1.1. See **table 1** (row c) for which equipment this is a requirement.
2. Compile an **Individual** Equipment Record (IER) containing the information as listed in **table 1 (column 2)** (see also section VI).
3. Perform the prescribed Acceptance- and Quality Control (QC) tests listed in **table 2 by:**
 - 3.1. An Inspection Body (IB) [30](#), [31](#) approved by the Department of Health (DoH) **OR** an appropriately trained professional registered with the HPCSA as a medical physicist (see 3.2.) must be used to perform all the acceptance tests as well as the routine tests listed in **section III.2 of table 2**.
 - 3.2. If a medical physicist is used to perform the tests in 3.1, an Inspection Body approved by the Department of Health (DoH) must formally contract such person(s). Formally contracted means that the medical physicist is contracted by the IB (ISO/IEC 17020 and TR78) to perform the tests.
4. Acquire the relevant quality control manuals **or** compile in-house written protocols, which describe each test step by step to ensure that QC tests listed in section III.1 of table 2 are correctly performed.
5. Ensure that persons that perform routine tests in section III.1 of table 2 are competent to execute the tests;
6. Ensure that the required acceptance tests are performed before the diagnostic x-ray equipment listed in **table 2** is put into clinical service when:
 - 6.1. Acquired or
 - 6.2. Substantially upgraded.
 - Acceptance tests are the initial tests performed directly after installation and before the equipment is being put into clinical service.
7. Ensure that all the quality control tests are performed at the prescribed frequencies as specified in **table 2**.
 - 7.1. QC tests may be performed more frequently than specified in table 2, influenced by the age, stability, make, model, etc., of the equipment.
8. Ensure that image display monitors and reporting monitors comply with the requirements in section **V (Table 4, page 27)** of this document.
9. Establish a program to ensure that the radiation dose administered to a patient for diagnostic purposes is optimised (see **bottom of table 4, page 27** for definition of optimisation). Such program must at least use the measurements under tests 37, 76, 146, **161** and **185** to determine whether radiation protection has been optimised.
 - 9.1. Measurements (**test results**) for tests 37, 76, 146, **161** and **185** must be evaluated at the prescribed frequencies. The following documents can be used as guidance documents for **establishment of Diagnostic Reference Levels (DRLs) and for comparisons.** [3](#), [27](#), [28](#), [29](#) Inter unit comparisons must also be performed.
 - 9.1.1. A medical physicist must be appointed in writing to establish and implement an optimisation program for Interventional Radiology procedures listed in section III.2.15.4.1. This optimisation program must amongst other include the establishment of Diagnostic Reference Levels (DRLs). The appointed medical physicist must audit and review the optimisation program on a **twelve monthly cycle**.
 - 9.1.1.1. The tasks of the appointed medical physicist shall at least include the following but not limited to:
 - 9.1.1.1.1. Implementation of procedures in establishment and use of DRLs;
 - 9.1.1.1.2. Investigate and review the program when DRLs are consistently exceeded and ensure that corrective action is taken where appropriate;
 - 9.1.1.1.3. Provide suitable training to theatre staff to achieve optimisation and such training must be documented;
 - 9.1.1.1.4. Assist with the investigations of over exposure to theatre staff;

- 9.1.1.1.5. Developing a local clinical protocol for each type of interventional procedure and each x-ray unit, and this protocol must at least include the following:³⁸
- 9.1.1.1.5.1. A statement on the 'expected' radiographic images including:
 - 9.1.1.1.5.1.1. Projections; and technique factors, and
 - 9.1.1.1.5.2. The 'nominal' values for:
 - 9.1.1.1.5.2.1. Fluoroscopy times and DAP readings / or dose; and air kerma rates; and resulting cumulative dose at each skin site exposed, and
 - 9.1.1.1.6. Any other tasks that could be included under optimisation and protection of staff against unnecessary exposure to ionising radiation.
10. Keep a copy of the results of the tests mentioned in section f and g of table 1 for as long as the equipment is in use and ensure that the following information is available:
- 10.1. The measurements (raw data), Date of test(s), Summary of the results (pass or fail), Identification of product, Details of the person(s) that performed the tests, and Details of the Inspection Body.
11. Report to the Director: Radiation Control if any one or more of the following conditions (trigger values or trigger events) are observed:
- 11.1. Kerma-Air Product is greater than 500 Gy.cm² per patient
 - 11.2. Cumulative Dose at interventional reference point K_{a,r} is greater than 5 Gy per patient
 - 11.3. A radiation injury is observed;
 - 11.4. An unprescribed or erroneously prescribed procedure is performed, and
 - 11.5. The patient is pregnant and the pregnancy is unknown at the time of the procedure.
- B If the licence holder can provide sufficient proof that all QC tests as listed in Table 2 were performed on general diagnostic imaging equipment under his control for the last two years, such licence holder may apply to the Directorate; Radiation Control that the 12 month QC test cycle be extended to a 24 month cycle (Equipment excluded is: Mammography, Fluoroscopy, Computed Tomography and x-ray units installed in vehicles). **This provision will be cancelled with immediate effect if full compliance with the requirements in this document is not maintained.**

II. TABLE 1 INDIVIDUAL EQUIPMENT RECORD (IER)¹ – (see also section VI)

		General Radiography Equipment	Processor & Hardcopy device	CR Reader	DDR System	Film Viewer	Reporting Monitor	Fluoroscopy Equipment	Computed Tomography Equipment	Mammography Equipment
a)	Unit - make, model and system ID	X	X	X	X			X	X	X
b)	Generator – make, model and serial number	X						X	X	X
c)	Product Licence number, date of the latest licence & reference where a copy of the licence is kept	X		X				X	X	X
d)	Date of installation	X	X	X	X	X	X	X	X	X
e)	Operator's manual – (Indication that the operator's manual is available and reference where it is kept)	X	X	X	X			X	X	X
f)	Results of acceptance tests	X	X	X	X		X	X	X	X
g)	Results of routine quality control tests	X	X	X	X	X	X	X	X	X
h)	Date(s) of tube replacement(s)	X						X	X	X
i)	Details of repairs/maintenance and/or modification(s). The licence holder must ensure that all the applicable test(s) are performed that could be affected by the aforementioned	X	X	X	X	X	X	X	X	X
j)	Should any of the tests in table 2 indicate non-compliance or should any problems be detected (indicated), the licence holder must implement corrective maintenance (repairs), followed by re-testing	X	X	X	X	X	X	X	X	X
k)	Details of the IB and person(s) that performed the test(s)	X		X	X	X	X	X	X	X

- ❖ The following documents can be used as guidance documents for purchasing of test equipment [1](#), [5](#), [6](#), [7](#), [10](#), [12](#), [16](#), [17](#) & [23](#) or alternatively ask your IB.
- ❖ For guidelines on what tests should be performed for an application see section VII.
- ❖ For new equipment acceptance tests is the responsibility of the company that installed the equipment.

¹ The X in each cell for each category of equipment (column 3 to 11), indicates which information must be available in the IER.

III. TABLE 2 ACCEPTANCE AND ROUTINE QUALITY CONTROL TESTS ³⁷

III.1. Routine Tests in this section are to be performed by the licence holder or person(s) appointed by the licence holder and Acceptance Tests in this section must be performed by an Inspection Body approved by Department of Health.				
	Physical parameter (required test)	Frequency	Standard	Reference
III.1.1. General Tests				
1.	Indicators, mechanical and other safety checks & warm-up	On acceptance & Daily	Results must be documented at least once every 3 months	page 30 ⁵
2.	Gonad shields, lead rubber aprons and gloves	3 monthly	Available and free from holes or cracks (Visual check and if suspect perform an x-ray test)	
3.	Appropriate technique chart displayed at x-ray unit	6 monthly	Available, applicable and compliant with ALARA principle	
III.1.2. X-ray Tubes and Generators				
4.	Alignment of the centre of the X-ray field and the centre of the bucky	On acceptance & 3 monthly	Deviation must be $\leq \pm 1$ cm @1m SID	RAD03 ¹⁰ & (A2.1, A2.2) ⁵
5.	The X-ray field dimensions in the plane of the image receptor must correspond with those indicated by the beam-limiting device	On acceptance & 3 monthly	Deviation must be $\leq \pm 1$ cm @1m SID	RAD04 ¹⁰ & (A2.1, A2.2) ⁵
6.	Congruence between the X-ray field and light field	On acceptance & 3 monthly	For any one side deviation must be $\leq \pm 1$ cm misalignment @1 m SID	RAD01 ¹⁰ & (A2.1, A2.2) ⁵
7.	X-ray/light beam centring	On acceptance & 3 monthly	Deviation must be $\leq \pm 1$ cm @1m SID	RAD02 ¹⁰ & (A2.1, A2.2) ⁵
8.	Alignment and collimation to film changer / bucky	On acceptance & 6 monthly	Any side ± 1 cm @1m	RAD06 ¹⁰ & (A2.1, A2.2) ⁵
III.1.2.1. Automatic Exposure Control (AEC) Device				
9.	Constancy (reproducibility) (test all chambers)	At 4 months intervals between annual tests	Baseline $\pm 20\%$ mAs or if mAs readout not available, Baseline ± 0.3 OD (use baseline of test 86 or 87)	(A4.2 or A4.1) ⁵

Table 2 continued

	Physical parameter (required test)	Frequency	Acceptance Standard	Reference
III.1.3. Processor Monitoring				
Tests must be performed before diagnostic films are processed. All measurements must be plotted on graph paper (Ref 15)				
10.	Processing temperature	Daily	Baseline \pm 1°C	IFSP01 10 & (D1) 5
11.	Base + Fog (B+F)	Daily	Variance \leq +0.03 OD. Maximum OD < 0.3	FSP02 10 & 17
12.	Mid-density (MD) step (speed index)	Daily	Variance \leq \pm 0.15 OD	FSP03 10 & 17
13.	Density difference (DD) (contrast index)	Daily	Variance \leq \pm 0.15 OD	FSP04 10 & 17
III.1.4. Intensifying Screens and Darkroom				
14.	Cleanliness of darkroom and screens	Written protocol for maintaining darkroom cleanliness, cassettes and screens clean, free from blemishes		
15.	Condition of cassettes and screens	12 monthly	Screen type, speed and date of installation Identification (cassette no.) and light tightness	FSP08 10 , (B1 & B3) 5 & 17
16.	Darkroom fog	Acceptance & 6 monthly & when fault reported	Density difference \leq 0.05 for 2 minutes	(C1 & C2) 5 & 17
17.	Relative speed of intensifying screens	Before initial use & 24 monthly	Baseline minus 10%	FSP09 10 & 17
III.1.5. CR Reader 2 & 12				
18.	Detector dose indicator monitoring (exposure index monitoring)	On acceptance & 3 monthly	Baseline \pm 20%	CR01 10 , (K1) 5 & (1) 12.2
19.	Image uniformity	On acceptance & 3 monthly	Free from dots and lines	CR02 10 & (2) 12.2
20.	Condition of cassettes and image plates	Supplier's recommendation	Free of dirt or damage	CR03 10 & Supplier's maintenance manual
21.	Test is not required – see test 93		CR04 10 & (3) 12.2	
22.	Test is not required – see test 95		CR05 10 & (4) 12.2	

Table 2 continued

	Physical parameter (required test)	Frequency	Acceptance Standard	Reference
III.1.5.1. AEC Device				
23.	Sensitivity	On acceptance & 3 monthly	Baseline \pm 30%	CR14 ¹⁰ & (K5) ⁵
III.1.6. DDR System				
24.	Detector dose indicator monitoring	On acceptance & 3 monthly	Baseline \pm 20%	DDR01 ¹⁰ & (1) ^{12.4}
25.	Image uniformity	On acceptance & 3 monthly	Lines or rectangles not apparent	DDR02 ¹⁰ & (2) ^{12.4}
26.	Test is not required – see test 106			DDR03 ¹⁰ & (3) ^{12.4}
III.1.6.1. AEC Device				
27.	Sensitivity	On acceptance & 3 monthly	Baseline \pm 25%	DDR13 ¹⁰ & (K5) ⁵
III.1.7. Film Viewing				
28.	Film viewer condition	6 monthly	Perceived brightness, colours and must be clean and uniformly illuminated	DD01 ¹⁰ & (M1) ⁵
III.1.8. Image Display Monitor & Reporting Monitor²				
29.	a) Condition of Image Display Monitor b) Condition of Reporting Monitor – Each reporting monitor must be labelled “REPORTING MONITOR”	a) At least 6 monthly. b) On acceptance & as required or at least weekly	a) Image display monitors should be clean & free from flicker b) Reporting monitors should be clean, and the perceived contrast of the test pattern should be consistent between monitors. Ensure that the 5% & 95% details superimposed on the 0% and 100% squares, respectively, are visible	IDD06 ¹⁰ & Use SMPTE or TG18 ¹⁸
30.	Test is not required – see test 119.2		IPEM 91 IDD07& TG 18	

² Reporting monitors refer to primary display systems used for the interpretation of medical images – i.e. excludes systems used by general medical staff & specialists after a report has been provided as well as operators' consoles, QC workstations and monitors used with fluoroscopy units, which are all classified as Display monitors (page 49 ¹⁰)

Table 2 continued

	Physical parameter (required test)	Frequency	Acceptance Standard	Reference
31.	Distance and angle calibration (<u>Comment</u> : This test is intended for applications where measurements of distance and angle are performed using image display monitor & diagnostic workstation)	On acceptance & 3 monthly	± 5 mm $\pm 3^\circ$ (degrees)	IDD08 ¹⁰
32.	Reporting monitors – Resolution	On acceptance & 3 monthly	Visual inspection of SMPTE or TG18-QC. Review both low contrast and high contrast resolution patterns. Check resolution at centre and periphery is consistent and similar to baseline image. Must be visible	IDD09 ¹⁰ & SMPTE or TG18 ¹⁸
III.1.9. Hardcopy Device (Only applicable if prints are used for reporting (interpretation of medical images))				
33.	Self – calibration	On acceptance & Weekly	Manufacturer's specification	IDD15 ¹⁰ & (N1) ⁵
34.	Optical density consistency	On acceptance & 3 monthly	Baseline OD ± 0.20	IDD16 ¹⁰ & (N2) ⁵
35.	Image quality	On acceptance & 3 monthly	Based on visual inspection	IDD17 ¹⁰ & (N3) ⁵
III.1.10. Reject Analysis				
36.	Reject analysis - Digital: Must use software supplied by vendor or implement effective procedure (general radiography)	3 monthly	May not increase with more than 2% from the previous determined rate and total rate should not exceed 10%	For film Screen use (Ch 2) ⁵ (4.10) ⁷ & ¹⁷
III.1.11. Fluoroscopy Equipment				
37.	Fixed fluoroscopic x-ray units must be equipped with a Dose Area Product (DAP) meter or a device that provide a dose read-out during fluoroscopy. DAP readings or dose read-out must be recorded in a book/register. The book/register must include the procedure, date of procedure, patient details, operator, specialist performing the procedure, the total dose (DAP reading/or dose) and the total fluoroscopy time. For each procedure in section III.2.15.4 the average DAP reading / average dose and average time must be calculated for a 12 month cycle by the licence holder and be recorded (see also test 216).			
38.	Radiation warning light at entrance, excluding theatres	On acceptance & Daily	Must work when beam is activated	
39.	Dose rate reproducibility under automatic exposure control	On acceptance & 3 monthly	Baseline $\pm 25\%$ (Use water container filled with water – approximately 30 cm x 30cm wide and 20cm thick)	FLU01 ¹⁰ & (H3) ⁵
III.1.11.1. Fluorography (For this section use IPEM Report 77) ⁵				
40.	Dose per frame reproducibility under automatic exposure control	On acceptance & 3 monthly	Baseline $\pm 25\%$ (For equipment with DAP meter)	⁹ & (B 11.1) ⁵

Table 2 continued

	Physical parameter (required test)	Frequency	Acceptance Standard	Reference
41.	Resultant film density	On acceptance & 3 monthly	Baseline ± 0.3 OD (Optical density)	9 & (B I1.2) 5
42.	Film density reproducibility	On acceptance & 3 monthly	Baseline ± 0.3 OD	9 & (B I1.3) 5
III.1.11.2. Digital Fluorography				
43.	Test is not required – see test 137			FLG01 10
44.	Test is not required – see test 138			FLG02 10
45.	Test is not required – see test 139.1			FLG03 10
III.1.12. Computed Tomography				
46.	Indicators, radiation warning light at entrance, mechanical and other safety checks	On acceptance & Daily	Must work properly	
47.	Image noise	On acceptance & Daily	Baseline $\pm 10\%$	CT01 10 & (B J1) 5
48.	CT number values	On acceptance & Daily	Water baseline ± 5 HU. Other material: baseline ± 10 HU	CT02 10 & (B J2) 5
49.	Scan plane localisation from alignment lights	On acceptance & 3 monthly	$\leq \pm 2$ mm	CT03 10 & (3.5.1) 18
III.1.13. Screen Film Mammography - For this section use ACR manual 4 or 6 as a guideline				
50.	Image quality evaluation (phantom images)	Weekly	At a minimum, the 4 largest fibers, the 3 largest speck groups, and the 3 largest masses must be visible. The background optical density must be at least 1.4 and the density difference should be at least 0.4 for a 4-mm thick acrylic disk. Maximum allowable changes are: mAs $\pm 15\%$; background density ± 0.2 ; density difference ± 0.05 ; fiber, speck groups or mass score decrease by 0.5. (Check manual for correct procedure)	Page 167 4

Table 2 continued

	Physical parameter (required test)	Frequency	Acceptance Standard	Reference
51.	Compression	On acceptance & 6 monthly	The maximum compression force must be between 111 Newton (11.3 kg) and 200 Newton (20.4 kg)	page 199 ⁴
52.	Repeat and reject analysis	3 monthly	May not increase with more than 2% from the previous determined rate and total rate shall not exceed 5%	page 202 ⁴ , (Chapter 2) ⁵ & (4.10) ⁷
53.	Accuracy of stereotactic device	On acceptance & Weekly or as used	Errors of ≤ 1 mm in X or Y or ≤ 3 mm in Z	MAM10 ¹⁰ & page 118 ¹¹
54.	Appropriate exposure technique chart (automatic and manual exposures) displayed near the control panel of the unit	6 monthly	Available and applicable	page 145 ⁴
55.	Analysis of fixer retention in film	6 monthly	The residual fixer retention shall be ≤ 5 micrograms per square cm	page 210 ⁴
III.1.14. Digital Mammography - For this section use European guidelines for quality assurance in breast cancer screening and diagnosis ⁶ & ³⁵				
56.	Repeat and reject analysis	3 monthly	May not increase with more than 2% from the previous determined rate and total rate shall not exceed 5%	(Chapter 2) ⁵ page 202 ⁴ & (4.10) ⁶
57.	AEC device: Long term reproducibility	On acceptance & weekly	Variation of SNR in the reference ROI and dose $< \pm 10\%$.	2b.2.1.3.4 ⁶ & 0604 ¹⁴
58.	Image receptor homogeneity	On acceptance & Weekly	Variation in mean pixel value $< \pm 15\%$ (on images); Maximum deviation in SNR $< \pm 15\%$ of mean SNR (on images); Maximum variation of the mean SNR between weekly images $\leq \pm 10\%$ (between images); Entrance surface air kerma OR tube loading (mAs) between weekly images $\leq \pm 10\%$	2b.2.2.3. ⁶ & (7.2.3) ²³

Table 2 continued

	Physical parameter (required test)	Frequency	Acceptance Standard	Reference
59.	Image quality evaluation (phantom images – RMI 156)	Weekly	At a minimum, the 5 largest fibers, the 4 largest speck groups, and the 4 largest masses must be visible. The background optical density must be at least 1.4 for hard copy. Maximum allowable changes are: mAs $\pm 10\%$ (EI tolerances for CR see table 7 of Ref 21); fiber, speck groups or mass score decrease by 0.5 (Check manual for correct procedure) and there shall be no blotches, lines and bright or dark pixels (Ref 21 par 7.2.4.4 and 7.3.2)	page 167 4 & (7.2.4) 23
60.	Uncorrected defective detector elements (DR systems)	On acceptance & Weekly	Limits of the manufacturer.	2b.2.2.3.3 6
61.	Monitors: Geometrical distortion (CRT displays)	On acceptance & Daily	Borders should be completely visible, lines should be straight, and the active display area should be centred on the screen.	2b.4.1.2 6
62.	Monitors: Contrast visibility	On acceptance & Daily	All corner patches shall be visible; the 5% and 95% pixel value squares shall be clearly visible.	2b.4.1.3 6
63.	Monitors: Display artefacts	On acceptance & Daily	No disturbing artefacts should be visible.	2b.4.1.5 6
64.	Printers: Geometrical distortion	On acceptance & Daily	Borders should be completely visible, lines should be straight.	2b.4.2.1 6
65.	Printers: Contrast visibility	On acceptance & Daily	All corner patches should be visible; the 5% and 95% pixel value squares should be clearly visible.	2b.4.2.2 6
66.	Printers: Printer artefacts	Daily	No disturbing artefacts should be visible.	2b.4.2.4 6

Table 2 continued

	Physical parameter (required test)	Frequency	Acceptance Standard	Reference
III.1.15. Small Field Digital Mammography System ¹⁵				
67.	Image quality evaluation (phantom images – RMI 156S)	Weekly (at least) or before use	At a minimum, the 3 largest fibers, the 3 largest speck groups, and the 2.5 largest masses must be visible. The background optical density must be at least 1.4 for hard copy. Maximum allowable changes are: mAs $\pm 10\%$ (EI tolerances for CR see table 7 of Ref 21); background density variation if hardcopy is produced is ± 0.2 ; fiber, speck groups or mass score decrease by 0.5 (Check manual for correct procedure).	page 167 ⁴ & (7.2.4) ²³
68.	Accuracy of stereotactic device	On acceptance & Weekly or as used	Errors of $\leq 1\text{mm}$ in X or Y or $\leq 3\text{mm}$ in Z	MAM10 ¹⁰ & page 118 ¹¹
III.1.16. Additional tests for mobile Mammography Systems ¹¹				
69.	Must ensure that all freely moveable objects/equipment are firmly locked or strapped down	Before moving		par 5.5.1 ¹¹
70.	Perform visual check of breast support and associated equipment for possible damage	After moving		par 5.5.2 ¹¹
71.	Compression device	After moving	Mechanical function and safety aspects must be checked	
72.	Alignment of x-ray beam to image receptor	After moving	For screen film see tests 149, 150 & 151 of this document; For digital see test 166 of this document	
73.	AEC system	After moving	For screen film see tests 153 & 154 of this document; For digital see test 57 and 173 of this document	
74.	Image quality	After moving	For screen film see test 50 of this document; For digital see test 59 of this document	

Table 2 continued

III.2. Acceptance tests and Routine tests listed in this section must be performed by an <u>Inspection Body (IB)</u> approved by the Department of Health				
	Physical parameter (required test)	Frequency	Acceptance Standard	Reference
III.2.1. General Tests				
75.	Safety of premises – On acceptance, when the workload increase or technique factors change that may jeopardise premises safety	On acceptance & changes that jeopardise safety	Controlled areas $\leq 5\text{mSv/year}$, for uncontrolled areas $\leq 1\text{mSv/year}$	NCRP 147 ¹³ & ²¹
76.	Entrance Surface Exposure (ESE) in air without backscatter for Chest, Lumbar Spine, Abdomen, Skull and Foot (ANSI phantom) For Paediatric – Perform measurements without phantom and on manual setting (technique factors used by radiographer) See section III.2.15.1 page 28	First set of ESE results must be reported after 12 months & thereafter every 24 months	ESE shall be evaluated in accordance with the guideline For Paediatric measurements the detector must be positioned at table to detector distance (TDD) – See section III.2.15.1 page 28 (IB report results on Electronic Submission)	Patient Dose Measurements ¹⁶ ; Paediatric, Table 19, Page 97 ³⁶
III.2.2. X-ray Tubes and Generators				
77.	Accuracy of the source (focal spot)-to-image distance (SID) indicators	On acceptance & 12 monthly	The difference between the indicated focus to film distance (FFD) and the actual FFD must be $\leq 2\%$	RAD05 ¹⁰ & par 3.4 ⁷
78.	Brightness of the light field, which defines the x-ray field.	On acceptance & 12 monthly	Average illuminance must be ≥ 100 lux at 100 centimetres or at the maximum FFD, whichever is less	par 2.11 ⁷
79.	Radiation output: repeatability	On acceptance & 12 monthly	Mean $\pm 10\%$	RAD09 ¹⁰
80.	Radiation output: reproducibility	On acceptance & 12 monthly	Baseline $\pm 20\%$	RAD10 ¹⁰
81.	The accuracy of the timer for different settings	On acceptance & 12 monthly	Manufacturers' specifications for specific model or if not available $\leq 10\%$	RAD11 ¹⁰
82.	The accuracy of the kV for different settings	On acceptance & 12 monthly	Manufacturers' specifications for specific model or if not available $\leq 10\%$	RAD12 ¹⁰
83.	Beam quality (half value layer (HVL))	On acceptance & Only to be tested when the x-ray tube or collimator is replaced	See section IV table 3	par 2.3 ⁷
84.	Leakage radiation from the diagnostic source assembly (x-ray tube)	Acceptance, tube replacement or after intervention of tube housing.	< 1 mGy in 1 hour at 1 m from the focus	Tube leakage ²⁰

Table 2 continued

	Physical parameter (required test)	Frequency	Acceptance Standard	Reference
III.2.2.1. Automatic Exposure Control (AEC) Device				
85.	Consistency between chambers	On acceptance & 12 monthly	Mean \pm 0.3 OD	FSP 14 10
86.	Repeatability (post-exposure mAs readout available, if not perform 87) (86 or 87)	On acceptance & 12 monthly	Mean \pm 20%	FSP15 10
87.	Repeatability	On acceptance & 12 monthly	Mean \pm 0.2 OD	FSP16 10
88.	Reproducibility (test all chambers) (as FSP13 but for different technique values – more extensive)	On acceptance & 12 monthly	Baseline \pm 0.3 OD	FSP17 10
89.	Image receptor dose	On acceptance & 12 monthly	Baseline \pm 30%	FSP18 10
III.2.3. CR Reader (see also Ref 1.1 & KCARE (Ref 10))				
90.	Detector dose indicator repeatability	On acceptance & 12 monthly	Baseline \pm 10%	CR06 10
91.	Detector dose indicator reproducibility	On acceptance & 12 monthly	Baseline \pm 20%	CR07 10
92.	Measured uniformity	On acceptance & 12 monthly	Mean \pm 10%	CR08 10
93.	Threshold contrast detailed detectability	On acceptance & 12 monthly	See comments CR09	CR09 10
94.	Erase cycle efficiency	On acceptance & 12 monthly	Blocker not visible in second image	CR10 10
95.	Limiting spatial resolution	On acceptance & 12 monthly	Baseline minus 25%	CR11 10
96.	Scaling errors	On acceptance & 12 monthly	\leq 2%	CR12 10
97.	Dark Noise	On acceptance & 12 monthly	Baseline + 50%	CR13 10
III.2.3.1. AEC Device				
98.	Consistency between chambers (Sensitivity / reproducibility)	On acceptance & 12 monthly	Baseline \pm 30% Mean \pm 20%	CR16 10
99.	Repeatability	On acceptance & 12 monthly	Mean \pm 20%	CR17 10

Table 2 continued

	Physical parameter (required test)	Frequency	Acceptance Standard	Reference
100.	Reproducibility	On acceptance & 12 monthly	Baseline \pm 30%	CR18 10
101.	Image receptor dose	On acceptance & 12 monthly	Baseline \pm 30%	CR19 10
III.2.4. DDR System (KCARE 10)				
102.	Test not required - see 107			DDR04 10
103.	Detector dose indicator repeatability	On acceptance & 12 monthly	Baseline \pm 10%	DDR05 10
104.	Detector dose indicator reproducibility	On acceptance & 12 monthly	Baseline \pm 20%	DDR06 10
105.	Measured uniformity	On acceptance & 12 monthly	Mean \pm 5%	DDR07 10
106.	Threshold contrast detail detectability	On acceptance & 12 monthly	See comments in report 91	DDR08 10
107.	Limiting spatial resolution	On acceptance & 12 monthly	Baseline minus 25%	DDR09 10
108.	Uniformity of resolution	On acceptance & 12 monthly	No increase in blurring from baseline	DDR10 10
109.	Scaling errors	On acceptance & 12 monthly	\leq 2%	DDR11 10
110.	Dark noise	On acceptance & 12 monthly	Baseline \pm 50%	DDR12 10
III.2.4.1. AEC Device				
111.	Consistency between chambers (sensitivity reproducibility)	On acceptance & 12 monthly	Baseline \pm 30% Mean \pm 20%	DDR15 10
112.	Repeatability	On acceptance & 12 monthly	Mean \pm 20%	DDR16 10
113.	Reproducibility	On acceptance & 12 monthly	Baseline \pm 30%	DDR17 10

Table 2 continued

	Physical parameter (required test)	Frequency	Acceptance Standard	Reference
114.	Image receptor dose	On acceptance & 12 monthly	Baseline \pm 30%	DDR18 ¹⁰
III.2.5. Film Viewing (Viewing boxes used for Reporting/Interpretation of medical images - see Chapter 7 of IPEM 91) & Film processing				
115.	Film viewer luminance	On acceptance & 12 monthly	\geq 1500 cd/m ² for general radiography	IDD02 ¹⁰
116.	Film viewer uniformity	On acceptance & 12 monthly	\leq 20%	IDD03 ¹⁰
117.	Film viewer variation	On acceptance & 12 monthly	\leq 20% difference from the mean value in bank	IDD04 ¹⁰
118.	Room illumination	On acceptance & 12 monthly	\leq 100 lux for general radiography	IDD05 ¹⁰
118.1.	Film processing evaluation – STEP	12 monthly	Processing speed between 80% to 120%	STEP ²⁵
III.2.6. Reporting Monitor				
119.	DICOM greyscale calibration	On acceptance & 12 monthly	GSD \pm 10%	IDD11 ¹⁰
119.1.	Minimum requirements for monitors	On acceptance & 12 monthly	Comply with table 4	table 1 & 2 ²⁴
119.2.	Reporting monitors – Greyscale (luminance response)	On acceptance & 12 monthly	Ratio white to black \geq 250	IDD07 ¹⁰ & ¹⁸
120.	Luminance uniformity	On acceptance & 12 monthly	Maximum variation \leq 30%	IDD12 ¹⁰
121.	Variation between monitors	On acceptance & 12 monthly	\leq 30%	IDD13 ¹⁰
122.	Room illumination	On acceptance & 12 monthly	\leq 15 lux for CRT displays & < 20 lux for LCD displays	IDD14 ¹⁰ + test 190
III.2.7. Fluoroscopy Equipment				
123.	Display monitor set-up	On acceptance & 12 monthly	All steps visible and black/white circles	FLU02 ¹⁰
124.	Minimum requirements for monitors	On acceptance & 12 monthly	Comply with table 4	²⁴
125.	Test is not required – see test 130		IPEM 91 FLU04 & BIR (B, H2)	

Table 2 continued

	Physical parameter (required test)	Frequency	Acceptance Standard	Reference
126.	Field limitation requirement. X-Ray field/Image intensifier	On acceptance & 12 monthly	The ratio of the areas ≤ 1.15 .	FLU05 ¹⁰
127.	Dose rate at <u>entrance</u> surface of phantom	On acceptance & 12 monthly	≤ 50 mGy/min (entrance air kerma) and <u>baseline $\pm 25\%$</u>	FLU06 ¹⁰
128.	Entrance exposure rate to image intensifier	On acceptance & 12 monthly	<u>Baseline $\pm 25\%$</u>	FLU07 ¹⁰
129.	Limiting spatial resolution	On acceptance & 12 monthly	36-40 cm: ≥ 0.7 line pairs mm^{-1} ; 30-35 cm: ≥ 0.8 line pairs mm^{-1} 25-29 cm: ≥ 0.9 line pairs mm^{-1} ; 20-24 cm: ≥ 1.0 line pairs mm^{-1} 15-18 cm ≥ 1.25 line pairs mm^{-1} .	FLU09 ¹⁰
130.	Threshold contrast	On acceptance & 12 monthly	See comments Flu10	FLU10 ¹⁰
131.	Image resolution uniformity	On acceptance & 12 monthly	See Comments FLU11	FLU11 ¹⁰
132.	Calibration of Dose area product meter (DAP/KAP meter) or the device that provides a dose read-out during fluoroscopy (total dose)	On acceptance & 12 monthly	Calibration of DAP/KAP meter or dose read out device according to manufacturer's specifications	(Page 336 – 340) ²²
III.2.7.1. Fluorography				
133.	Overall Image quality	On acceptance & 12 monthly	Manufacturer's specifications for a specific model	(B 11.4) ⁵
134.	Resultant film density	On acceptance & 12 monthly	Baseline ± 0.3 OD	(B 11.3) ⁵
135.	Dose per frame at the input face of the image intensifier under automatic exposure control	On acceptance & 12 monthly	Baseline $\pm 25\%$ or ≤ 1 μGy per frame (Largest field)	(B 11.1) ⁵
136.	Image quality: limiting spatial resolution	On acceptance & 12 monthly	1.6 line-pairs/mm for 30-35 cm systems; 2.5 line-pairs/mm for 23-25 cm systems, and 3 line-pairs/mm for 15-18 cm systems.	(B 11.4) ⁵ & ⁹
III.2.7.2. Digital Fluorography				
137.	Dose per image at the input face of the image receptor under automatic exposure control	On acceptance & 12 monthly	Baseline $\pm 25\%$	FLG04 ¹⁰
138.	Limiting spatial resolution	On acceptance & 12 monthly	Baseline reduced by 2 groups	FLG05 ¹⁰
139.	Dynamic range	On acceptance & 12 monthly	See Comments FLG07	FLG07 ¹⁰
139.1.	Threshold contrast	On acceptance & 12 monthly	Baseline ± 2 discs	FLG06 ¹⁰

Table 2 continued

	Physical parameter (required test)	Frequency	Acceptance Standard	Reference
III.2.8. Computed Tomography				
140.	Image noise	On acceptance & 12 monthly	Baseline \pm 10% Inter –slice variation; Mean \pm 10%	CT06 10
141.	CT number values	On acceptance & 12 monthly	Water baseline \pm 5 HU Other materials: baseline \pm 10HU	CT07 10
142.	CT number uniformity	On acceptance & 12 monthly	Head phantom: \leq \pm 10HU Body phantom: \leq \pm 20HU	CT08 10
143.	High contrast spatial resolution	On acceptance & 12 monthly	Baseline \pm 20%	CT09 10
144.	Computed tomography dose index (CTDI)	On acceptance & 12 monthly	Baseline \pm 15%	CT10 10
145.	Image slice thickness	On acceptance & 12 monthly	Baseline \pm 20% or \pm 1mm, whichever is greater	CT13 10
146.	CTDI _{vol} for technique factors used for groups specified in section III.2.15.3 page 28	On acceptance & 12 monthly	\leq Reference dose - table 3 of reference 1.2 and reference 24 (IB report results on Electronic Submission)	CT11 10
III.2.9. Screen Film Mammography ⁴				
147.	Screen-film systems – Image receptors	On acceptance & 12 monthly	Must have image receptors of 18x24 cm and 24x30 cm with matching moving grids	
148.	Assessment of locks, detents, angulation indicators, and mechanical support devices for X-ray tube and image receptor holder assembly	On acceptance & 12 monthly	Must function properly	Page 231 4
149.	Collimation assessment: Deviation between X-ray field and light field	On acceptance & 12 monthly	The sum of left plus right edge deviations or anterior plus chest edge deviations must be \leq 2% of SID	Page 233 4
150.	Collimation assessment: Deviation between X-ray field and edges of the image receptor	On acceptance & 12 monthly	The X-ray field may not exceed the image receptor at any side by more than 2% of SID and the X-ray field may not fall within the image receptor on the chest wall side	Page 233 4
151.	Collimation assessment: Alignment of chest-wall edges of compression paddle and film	On acceptance & 12 monthly	The chest-wall edge of the compression paddle may not fall within the image receptor or project beyond the chest-wall edge of the image receptor by more than 1% of SID	Page 233 4

Table 2 continued

	Physical parameter (required test)	Frequency	Acceptance Standard	Reference
152.	Evaluation of system resolution	On acceptance & 12 monthly	The resolution with the bars parallel to the anode-cathode axis must be ≥ 13 line-pairs/mm or with the bars perpendicular to the anode-cathode axis must be ≥ 11 line-pairs/mm	Page 238 4
153.	Automatic exposure control (AEC) system performance: Thickness tracking, kVp tracking and image mode tracking	On acceptance & 12 monthly	<u>Equipment sold prior to 01/01/2003:</u> The AEC system must maintain the film optical density within ± 0.3 of the mean when the thickness of the phantom is varied over 2-6 cm and the kVp is varied over the range of those used clinically for these thickness. If this requirement cannot be met, a technique chart shall be developed showing appropriate techniques (kVp and density control settings) for different breast thickness and compositions that must be used so that optical densities within ± 0.3 of the average can be produced under photo timed conditions. <u>Equipment sold after 01/01/2003:</u> The AEC system must maintain the film optical density within ± 0.15 of the mean when the thickness of the phantom is varied over 2-6 cm and the kVp is varied over the range of those used clinically for these thickness.	Page 241 4
154.	Automatic exposure control (AEC) system performance: Density control	On acceptance & 12 monthly	Each step (density setting) shall result in a 12-15% change in mAs, or approximately a 0.15 change in film optical density	Page 241 4
155.	Uniformity of screen speed (for all cassette sizes)	On acceptance & 12 monthly	The standard deviation for the control cassette densities must be less than 0.05 and density range for all cassettes (of the same size) must be ≤ 0.30	Page 246 4
156.	Image quality evaluation	On acceptance & 12 monthly	At a minimum, the 4 largest fibers, the 3 largest speck groups, and the 3 largest masses must be visible. The background optical density must be at least 1.4 and the density difference should be at least 0.4 for a 4-mm thick acrylic disk.	Page 258 4
157.	Artefact evaluation	On acceptance & 12 monthly	No significant artefacts must be visible	Page 249 4
158.	kVp accuracy and reproducibility	On acceptance & 12 monthly	The mean kVp may not differ from the nominal kVp (set value) with more than $\pm 5\%$, or the coefficient of variation may not exceed 0.02	Page 271 4
159.	Beam quality (HVL) measurement	On acceptance & 12 monthly	The measured HVL must be $\geq \text{kVp}/100$ (mm Al) (Please note 0.03 must be added when filtration is performed with compression paddle (see page 275) of 1999 addition, ACR)	Page 273 4

Table 2 continued

	Physical parameter (required test)	Frequency	Acceptance Standard	Reference
160.	AEC reproducibility	On acceptance & 12 monthly	The coefficient of variation for R (exposure) or mAs must be ≤ 0.05	Page 277 4
161.	Average glandular dose	On acceptance & 12 monthly	The dose must be ≤ 300 mRad (3 mGy) for 4.2 cm effective breast thickness (IB report results on Electronic Submission)	Page 277 4
162.	Radiation output rate	On acceptance & 12 monthly	The output must be ≥ 800 mR/s (7.0 mGy/s) at maximum SID ³	Page 277 4
163.	View box luminance, room illuminance and masking	On acceptance & 12 monthly	Luminance of the view box shall be ≥ 3000 cd/m ² and illuminance of the room shall be ≤ 50 lux. Viewboxes must be masked to the exposed area of the film	Page 286 4
III.2.10. DDR & CR Mammography ^{6 & 35} (Reference 35 must be consulted)				
164.	Assessment of locks, detents, angulation indicators, and mechanical support devices for X-ray tube and image receptor holder assembly	On acceptance & 12 monthly	Comply to par 8.2.1 of Ref 21	(8.2.1) 23
165.	X-ray source: Source to image distance Only if adjustable	On acceptance & 12 monthly	Manufacturers specification, typical 600-650 mm.	2b.2.1.1.2 6
166.	X-ray source: Alignment of X-ray field/image receptor	On acceptance & 12 monthly	All sides: X-rays must cover the film by no more than 5 mm outside the film. On chest wall edge: distance between film edge and edge of the bucky must be ≤ 5 mm.	2b.2.1.1.3 6
167.	X-ray source: Radiation leakage	On acceptance and after intervention on the tube housing.	≤ 1 mGy in 1 hour at 1 m from the focus	2b.2.1.1.4 6
168.	X-ray source: Tube output	On acceptance & 12 monthly	> 30 μ Gy/mAs at 1 metre and $> 70\%$ of value at acceptance	2b.2.1.1.5 6
169.	Tube voltage reproducibility and accuracy	On acceptance & 12 monthly	Accuracy for the range of clinically used tube voltages: $< \pm 1$ kV Reproducibility $< \pm 0.5$ kV	2b.2.1.2.1 6
170.	Half Value Layer (HVL)	On acceptance and after intervention on the tube housing.	4 th edition supplement standard.	2b.2.1.2.2 35

³ Test 168 - Units manufactured after 01-01-2003. Units manufactured prior to 01-01-2003 and that do not comply may not be resold.

Table 2 continued

	Physical parameter (required test)	Frequency	Acceptance Standard	Reference
171.	AEC-system: Optical density control setting: central value and difference per step (if applicable)	On acceptance & 12 monthly	Measure increase in exposure per step and inform user - must be displayed at technique chart	2b.2.1.3.1 6 35
172.	AEC-system: Short term reproducibility	On acceptance & 12 monthly	$< \pm 5\%$	2b.2.1.3.3 6
173.	AEC-system: Object thickness and tube voltage compensation	On acceptance & 12 monthly	Thickness indicator $< \pm 0.5$ cm and 4 th edition supplement standard.	2b.2.1.3.5, 6 , 35
174.	Compression force	On acceptance & 12 monthly	130 - 200 N (13-20 kg) maintained unchanged for at least 1 minute and indicated compression force should be within ± 20 N of the measured value	2b.2.1.4 6
175.	Compression plate alignment	On acceptance & 12 monthly	≤ 5 mm	2b.2.1.4 6 & (8.9) 23
176.	Local dense area (only DR systems)	On acceptance & 12 monthly	The SNR of each image should be within 20% of the average SNR	2b.2.1.3.6 35
177.	Grid imaging	On acceptance & 12 monthly	No significant non-uniformity	2b.2.1.5.2 6
178.	Image receptor response function	On acceptance & 12 monthly	$R^2 > 0.99$, results at acceptance are used as reference.	2b.2.2.1.1 6
179.	Image receptor Noise evaluation	On acceptance & 12 monthly	Results at acceptance are used as reference	2b.2.2.1.2 6
180.	Missed tissue at chest wall side	On acceptance	Width of missed tissue at chest wall side ≤ 5 mm	2b.2.2.2.4 6 & (8.9) 23
181.	Image receptor homogeneity and Image receptor detector element failure (DR systems)	On acceptance & 12 monthly	Variation in mean pixel value $< \pm 30\%$ (on images); Maximum deviation in SNR $< \pm 15\%$ of mean SNR (on images); Maximum variation of the mean SNR between weekly images $\leq \pm 10\%$ (between images); Entrance surface air kerma OR tube loading (mAs) between annual images $\leq \pm 10\%$ Limits of the manufacturer.	2b.2.2.3.1 35 & 2b.2.2.3.2 6
182.	Inter plate sensitivity variations (CR systems)	On acceptance & 12 monthly	SNR variation $\leq \pm 10\%$. Variation in entrance surface air kerma OR tube loading (mAs) $\leq \pm 10\%$,	2b.2.2.4 6 , 35

Table 2 continued

	Physical parameter (required test)	Frequency	Acceptance Standard	Reference
183.	Influence of other sources of radiation (CR)	On acceptance	The coins should not be visible.	2b.2.2.5 6
184.	Fading of latent image (CR)	On acceptance	Results at acceptance are used as reference.	2b.2.2.6 6
185.	Dosimetry	On acceptance & 12 monthly	< 2.5 mGy for 4.5 cm PMMA - see 2a.2.5.1 for rest of values (IB report results on Electronic Submission)	2b.2.3 6
186.	Threshold contrast visibility	On acceptance & 12 monthly	See table in 2b.2.4.1 for limiting values	2b.2.4.1 6
187.	Exposure time	On acceptance & 12 monthly	< 2 s	2b.2.4.3 6
188.	Geometric distortion and artefact evaluation	On acceptance & 12 monthly	No disturbing artefacts, no visible distortion.	2b.2.4.4 6
189.	Ghost image/erasure thoroughness	On acceptance & 12 monthly	"Ghost image"-factor < 0.3	2b.2.4.5 6
190.	Monitors : Ambient light	On acceptance & 12 monthly	< 10 lux for CRT displays & < 20 lux for LCD displays	2b.4.1.1 6 +35
191.	Monitors: Resolution	On acceptance & 12 monthly	All line patterns should be discernible.	2b.4.1.4 6
192.	Monitors: Luminance range: Maximum to minimum luminance ratio	On acceptance & 12 monthly	Primary display devices ≥ 250 Secondary display devices ≥ 100 ; Displays belonging to one displaying station should not exceed 5% of the lowest.	2b.4.1.6 6
193.	Monitors: Greyscale Display Function	On acceptance & 12 monthly	$< \pm 10\%$ of the GSDF for primary class displays and $< \pm 20\%$ of the GSDF for secondary class displays	2b.4.1.7 6
194.	Monitors: Luminance uniformity	On acceptance & 12 monthly	Maximum luminance deviation of a display device should be less than 30% for CRT displays and LCD displays $((L_{max}-L_{min})/L_{centre} < 0.3)$.	2b.4.1.8 6
195.	Printers: Resolution	On acceptance	All line patterns should be discernible	2b.4.2.3 6
196.	Printers: Greyscale Display Function	On acceptance & 12 monthly	The calculated contrast response should fall within $\pm 10\%$ of the GSDF contrast response.	2b.4.2.6 6
197.	Printers: Density uniformity	On acceptance & 12 monthly	Maximum optical density deviation should be less than 10% $((D_{max}-D_{min})/D_{centre} < 0.1)$	2b.4.2.7 6

Table 2 continued

	Physical parameter (required test)	Frequency	Acceptance Standard	Reference
198.	Image quality evaluation (phantom images – RMI 156)	On acceptance & 12 monthly	At a minimum, the 5 largest fibers, the 4 largest speck groups, and the 4 largest masses must be visible. The background optical density must be at least 1.4 for hard copy. Maximum allowable changes are: mAs $\pm 10\%$ (EI tolerances for CR see table 7 of Ref 21); fiber, speck groups or mass score decrease by 0.5 and there shall be no blotches, lines and bright or dark pixels (Ref 21 par 7.2.4.4 and 7.3.2)	Page 167 ⁴ & 23
199.	Viewing boxes	On acceptance & 12 monthly	If mammograms are read on printed images, use the method and limiting values described in section 2a.2.4.1	2b.4.3 ⁶
III.2.11. Small Field Digital Mammography System ¹⁵				
200.	For dedicated small field digital imaging systems the applicable quality control tests specified in section III.2.10 and III.1.14 must be included. For image quality use RMI 156S phantom			
201.	Beam alignment: Alignment of the light field to the x-ray field	On acceptance & 12 monthly	± 10 mm on all sides	3.1 ¹⁵
202.	Beam alignment: Alignment of the x-ray field to the imaged field	On acceptance & 12 monthly	0 to + 10 mm on all sides	3.1.1 ¹⁵
203.	Size of image field	On acceptance	Each dimension should be within 5% of specified value	3.1.2 ¹⁵
204.	X-ray field non-uniformity	On acceptance & 12 monthly	Variation in <i>pixel value</i> $\leq 10\%$ from the value measured in the centre of the image	3.2 ¹⁵
205.	Automatic exposure control: Overall repeatability	On acceptance & 12 monthly	Maximum deviation in mAs $\leq 5\%$ from the mean	3.3.1 ¹⁵
206.	Constancy with change in phantom thickness	On acceptance & 12 monthly	Maximum deviation in <i>pixel values</i> $\leq 10\%$ of the mean	3.3.2 ¹⁵
207.	Constancy with change in tube voltage	On acceptance & 12 monthly	Maximum deviation in <i>pixel values</i> should not exceed 10% of the mean	3.3.3 ¹⁵
208.	Display devices: Greyscale	On acceptance & 12 monthly	Monitor – 5% steps from 0% and 100% grey levels equally visible Hardcopy – baseline greyscale step ± 0.15 OD (± 0.05 OD for minimum density step)	3.4.1 ¹⁵
209.	Display devices: Resolution	On acceptance & 12 monthly	Frequency high contrast resolution pattern resolved	3.4.1 ¹⁵

Table 2 continued

	Physical parameter (required test)	Frequency	Acceptance Standard	Reference
210.	Hardcopy printer: Greyscale	On acceptance & 12 monthly	Greyscale must match the image display monitor and the greyscale steps selected shall be within the following tolerances: Step 1: ± 0.05 ; Step 2: ± 0.15 ; Step 3: ± 0.15	3.4.2 15
211.	Hardcopy printer: Resolution	On acceptance & 12 monthly	Maximum frequency in the high contrast patterns should be resolved	3.4.2 15
212.	Image quality: Limiting spatial resolution	On acceptance & 12 monthly	Should be at least 70% of the Nyquist frequency of the detector. Should be at least 75% of the value determined at commissioning	3.5.1 15
213.	Image quality evaluation (phantom images – RMI 156S)	On acceptance & 12 monthly	≥ 3 largest fibers, ≥ 3 largest speck groups, and ≥ 2.5 largest masses be visible. The background optical density ≥ 1.4 for hard copy. Maximum allowable changes are: mAs $\pm 15\%$; fiber, speck groups or mass score decrease by 0.5.	Page 167 ⁴
214.	Measurement of dose: Dose to the standard breast at the clinical setting	On acceptance & 12 monthly	Variation within $\pm 25\%$ of value determined at commissioning and the dose must be ≤ 3 mGy for 4.2 cm effective breast thickness	3.6.1 15
III.2.12. DRLs				
215.	A Medical physicist is appointed in writing and an optimisation program is implemented	12 monthly	Comply with requirements in paragraph I.A.9.1.1	
216.	For each procedure in section III.2.15.4 the average DAP reading / average dose and average time was calculated by licence holder, documented and reported by IB	12 monthly	The Inspection Body must report these results on the Electronic Submission	Fluoroscopy 27, 28 & 29

III.2.13. TABLE 3 – HVL values										
X-ray tube voltage (kilovolt peak)	71	80	90	100	110	120	130	140	150	> 150
Minimum HVL (mm of Al)	2.1	2.3	2.5	2.7	3.0	3.2	3.5	3.8	4.1	See note 2
Minimum HVL (mm of Al), manufactured after June 2006	2.5	2.9	3.2	3.6	3.9	4.3	4.7	5.0	5.4	

1. HALF-VALUE LAYERS for intermediate selected voltages are to be obtained by linear interpolation.

2. Linear extrapolation is to be used.

III.2.14. Table 4 - MINIMUM REQUIREMENTS FOR MONITORS ²⁴

Description and application	Monitors purchased on or after 1 March 2012		
	Licensed with Department of Health as a medical device for import	Minimum resolution	CE medical device
01. Diagnostic (reporting) monitor used in mammography	Yes	5 Megapixel	Yes
02. Diagnostic (reporting) monitor used in conventional radiology	Yes	3 Megapixel	Yes
03. Diagnostic (reporting) monitor used in Computed Tomography	Yes	1.3 Megapixel	Yes
04. All monitors not covered under 01, 02 and 03 (e.g. Image display monitor not to be used for diagnosis but images viewed only in <u>conjunction with the report</u> – ward, clinic, theatre, etc; Workstations; Image review monitors (<u>not</u> used for immediate feedback to clinical activity); Fluoroscopy (production of dynamic x-ray images which are displayed in real time); etc)	No	1 Megapixel	Yes

- Optimisation** in diagnostic radiology means that equipment and methods must be selected to ensure that radiation administered to a patient for diagnostic purposes, is sufficient to enable the procedure to provide the required information; and not greater than is necessary to provide that information.
- All diagnostic image interpretation shall be performed by making use of the application software which includes, zoom, pan, magnification and windowing tools to optimise spatial and contrast resolution.

III.2.15. TABLE 5 - DIAGNOSTIC REFERENCE LEVELS

III.2.15.1. GENERAL RADIOGRAPHY											
Report ESD per radiograph (mGy) as determined with test 76											
250	Chest (PA) Grid	251	Abdomen (AP) Grid	252	Lumbar Spine (AP) Grid	253	Skull (Lateral) Grid	254	Foot - Non-grid		
255	Chest 1 year (TDD 13 cm)	256	Abdomen 1 year (TDD 13 cm)	257	Chest 5 year (TDD 15 cm)	258	Abdomen 5 year (TDD 15 cm)				
259	Chest (PA) 10 year (TDD 16.8 cm)	260	Abdomen 10 year (TDD 16.8 cm)								
III.2.15.2. MAMMOGRAPHY											
Report ESD per radiograph (mGy) as determined with test 161 & 185 – Dose values for a 4.5 cm phantom must be reported											
261	Mammography Average glandular dose										
III.2.15.3. COMPUTED TOMOGRAPHY											
Report average CTDI_{vol} per examination (mGy) as determined with test 146											
Paediatric is between 1 to 5 years.											
262	Adult head	263	Adult chest	264	Adult abdomen-pelvis	265	IVP	266	Paediatric abdomen	267	Paediatric head

III.2.15.4. FLUOROSCOPIC EXAMINATIONS

Report average DAP value per examination (**Gy.cm²**) and average fluoroscopy time per examination as determined with test 37 for **each** fixed fluoroscopic unit. Pediatric is between 1 to 5 years. The correction factor as determined with test 132 must be used to correct all average values for reporting

282	Barium (or water soluble) swallow	283	Barium meal	284	Barium follow through	285	Barium (or water soluble) enema
286	Small bowel enema	287	MCU	288	MCU - Pediatric		

III.2.15.4.1. INTERVENTIONAL EXAMINATIONS

	Procedure Name	Also known as		Procedure Name	Also known as
289	Coronary Angiography	CA (Coronary Angiogram) / Cardiac Catheterization	290	Cerebral Angiography	Cerebral Angiogram / Neuro-Angiogram
291	Cerebral Angiography + Interventions	Neuro-Angiogram + Interventions	292	Renal Angiography	Renal Angiogram / Renal Arteriography
293	Peripheral Angiography	Peripheral Angiogram / Peripheral Arteriogram	294	CA + EPS	Coronary angiogram + Electro Physiology Study
295	Ablation (RF)	Catheter Ablation - Radio Frequency / Cardiac RF ablation / Radio Frequency ablation	296	CA + LV function	CA + LV function; Left Ventriculography
297	EPS	Electrophysiology Study	298	Pacemaker (Bi Vent)	Biventricular Pacemaker / CRT (Cardiac Resynchronization Therapy)
299	CA + EPS + Ablation	CA + EPS + Catheter ablations	300	TAVI	Valve placement; Transcatheter Aortic Valve Implantation
301	Pacemaker (Permanent)	Pacemaker/ PPM	302	Pediatric Diagnostic heart caths	Pediatric Diagnostic left & right heart catheters
303	EVAR	Endovascular Aneurysm Repair / Endovascular Aortic Repair	304	Femoral angiogram	
305	Uterine Artery Embolisation (UAE)		306	ERCP (Endoscopic retrograde cholangiopancreatography)	
307	Percutaneous transhepatic cholangiography (PTHC or PTC)				

VI. EXAMPLE OF A FORM THAT SHOULD BE INCLUDED IN IER

A UNIT PARTICULARS								
DoH ref. no.:		Date of latest DoH document			Copy is available at:			
Product Licence no.:			Appointed person responsible for QC tests:					
Inspection Body ⁴ :								
<i>X = Indicate Applicability</i>	General Radiography	Processor & Hardcopy device	CR System	DDR System	Reporting Monitor	Fluoroscopy Equipment	CT	Mammo
Date of installation		Operator's manual(s) is available & where is it kept?			Results of acceptance tests is available & date			
Date(s) of replacement(s) / upgrading								
B COMPONENT PARTICULARS								
		Make		Model			ID number /Serial number	
Unit - make, model and system ID								
Generator								
X-ray Tube(s)								
Comments								

⁴ An Inspection Body is an Organisation that is accredited by SANAS (www.sanas.co.za) and approved by the Department of Health

C TESTS APPLICABLE ON MACHINE			DoH Licence no.		
Table 2 Section III.1		<u>The licence holder must perform Routine Tests. An Inspection Body must perform acceptance tests.</u>	Table 2 Section III.2		<u>An Inspection Body must perform acceptance tests and Routine tests⁵.</u>
Ref. No.	Physical parameter to be tested	X = Indicate Applicability	Ref. No.	Physical parameter to be tested	X = Indicate Applicability
III.1.1	General Tests		III.2.1	General Tests	
III.1.2	X-Ray Tubes and Generators		III.2.2	X-Ray Tubes and Generators	
III.1.2.1	Automatic Exposure Control (AEC) Device		III.2.2.1	Automatic Exposure Control (AEC) Device	
III.1.3	Processor Monitoring				
III.1.4	Intensifying Screens and Darkroom				
III.1.5	CR Reader		III.2.3	CR Reader	
III.1.5.1	AEC Device		III.2.3.1	AEC Device	
III.1.6	DDR System		III.2.4	DDR System	
III.1.6.1	AEC Device		III.2.4.1	AEC Device	
III.1.7	Film Viewing		III.2.5	Film Viewing	
III.1.8	Image Display Monitor		III.2.6	Image Display Monitor	
III.1.9	Hardcopy Device				
III.1.10	Repeat and Reject Analysis				
III.1.11	Fluoroscopy Equipment		III.2.7	Fluoroscopy Equipment	
III.1.11.1	Fluorography		III.2.7.1	Fluorography	
III.1.11.2	Digital Fluorography		III.2.7.2	Digital Fluorography	
III.1.12	Computed Tomography		III.2.8	Computed Tomography	
III.1.13	Screen Film Mammography		III.2.9	Screen Film Mammography	
III.1.14	Digital Mammography		III.2.10	Digital Mammography	
III.1.15	Small Field Digital Mammography System		III.2.11	Small Field Digital Mammography System	
III.1.16	Additional Tests for mobile Mammography Systems				

⁵ An Inspection Body is an Organisation that is accredited by SANAS (www.sanas.co.za) and approved by the Department of Health
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VII. TEST GUIDELINES

- X-Ray Tubes and Generators – Conventional film systems; (tests 1-17, 28, 33-36, 75-89, 115-118 ~~118-1~~);
- Computerised Radiography Reader (tests 18-20, 23, 29-35, 90-101, 119-122);
- Direct Digital Radiography System, (tests 1-8, 24-25, 27, 29-36, 75-84, 102-114, 119-122);
- Fixed Fluoroscopy Equipment (tests 1-3, 37-42, 75, 79-84, 123-139.1, ~~215~~, ~~216~~);
- Mobile Fluoroscopy and X-Ray Tubes and Generators (tests 1-3, 39, 79-84, 123-131, 137-139.1)
- Computed Tomography (tests 1-3, 29-32, 36, 46-49, 75, 119-122, 140-146)
- Screen film Mammography (tests 1, 10-17, 28, 50-55, 69-75, 116-117, ~~118-1~~, 147-163)
- Digital Mammography (tests 1, 56-75, 119-122, 164-214)

VIII. REFERENCES

- References listed below can/should be used as guidelines. Purchasing of these documents is not a requirement. Other sources could be consulted in obtaining the relevant information.
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 4. **ACR** (American College of Radiology), Mammography Quality Control Manual (1999) www.acr.org → ACR Store → Quality and Safety → Quality Control Manuals
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 8. **IMPACT**, CT Scanner Acceptance Testing, www.impactscan.org → Reports & info → acceptance testing of CT
 9. **IPEM** (Institute of Physics and Engineering in Medicine) 1997, Recommended Standards for the Routine Performance Testing of Diagnostic X-ray Imaging Systems, **Report no. 77**, www.IPEM.org.uk
 10. **IPEM** (Institute of Physics and Engineering in Medicine) 2005, Recommended Standards for the Routine Performance Testing of Diagnostic X-ray Imaging Systems, Report no. 91, www.IPEM.org.uk
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 12. **KCARE**; Protocols for QA of CR System – Routine and Annual; Protocols for QA of DDR Systems – Routine and Annual, <http://www.kcare.co.uk> → Education → Protocols

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- 12.1. CR system: Commissioning & Annual tests;
- 12.2. CR system: Routine QA tests;
- 12.3. DDR system: Commissioning & Annual tests, and
- 12.4. DDR system: Routine QA tests.
13. **NCRP** (National Council on Radiation Protection and Measurements) 2004, Structural Shielding Design for Medical X-Ray Imaging Facilities, NCRP Report No.147, <http://www.ncrponline.org>
14. **NHSBSP** Equipment Report **0604**, June 2006, Commissioning and Routine testing of full field digital mammography systems, www.cancerscreening.nhs.uk → Search this site for → *Report 0604*
15. **NHSBSP** Equipment Report **0705**, May 2007, Commissioning and Routine testing of small field digital mammography systems, www.cancerscreening.nhs.uk → Search this site for → *Report 0705*
- 15.1. Quality Assurance Guidelines for Mammography Including Radiographic Quality Control, Publication No 63, www.cancerscreening.nhs.uk → Search this site for → *Quality Assurance Guidelines for Mammography Including Radiographic*
16. Patient Dose Measurements in Diagnostic Radiology <https://sites.google.com/site/radiationcontroldoh/> → **Electronic devices – Use → Electronic devices - ionising radiation → Guidelines**
17. Test procedures for film processing and intensifying screens, <https://sites.google.com/site/radiationcontroldoh/> → **Electronic devices – Use → Electronic devices - ionising radiation → Guidelines**
18. **TG18** by **AAPM** (American Association of Physicists in Medicine), Task Group 18, <http://deckard.mc.duke.edu/~samei/tg18> OR www.aapm.org → Publications → Reports → **OR-03**
19. Measurement of the Performance Characteristics of Diagnostic X-Ray Systems: Digital Imaging Systems, www.IPEM.org.uk
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22. www-pub.iaea.org/MTCD/publications/PDF/TRS457_web.pdf OR <http://www.radcal.com/PDC.html>
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28. NCRP Report no. 172, Reference Levels and Achievable Doses in Medical and Dental Imaging: Recommendations for the United States
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30. TR 78-03, Technical requirements for the application of SANS/ISO/IEC 17020: 2012 for testing of Diagnostic X-ray Imaging Systems by Inspection Bodies, www.Sanas.co.za;
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36. Dosimetry in Diagnostic Radiology for Paediatric Patients, Pub1609, IAEA
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