RC-4a

Mammography Screening

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MAMMOGRAPHY SCREENING

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Abstract

The objective of any screening programmes it to test a population group to identify a sub-group who have disease at an early stage. For a screening programme to be effective there should be a suitable test or examination. In addition there must also be a treatment which confers benefit when the sub-group is treated at an early stage. In breast screening, the test used to identify breast cancer at an early stage is x-ray examination mammography. This examination is only of benefit if early breast cancers (i.e. small tumours) are detected, this requires obtaining and maintaining high quality mammography. This may only be delivered if there is rigorous and comprehensive quality assurance of x-ray mammography. The use of ionising radiation as an intrinsic part of the screening process means that screening must be justified in radiation protection terms as well. The benefit to the screened population must exceed the risk. Thus dose assessment must be an integral part of screening. Mammography screening can only be successful in reducing fatal breast cancer deaths, thus there must be quality assurance of the process. A comprehensive quality assurance programme for breast screening is described in detail.
1 Introduction

The objective of any screening programme is to apply a test to a population group, to identify a sub-group who have disease at an early stage. For a screening programme to be effective there must be a suitable test or examination. In addition, this sub-group must also benefit from a better prognosis when detected early or at a stage before symptoms develop. In other words, if the sub-group is treated early they must survive longer than if they had been treated once symptoms have developed.

In breast screening high quality x-ray mammography is used to detect breast cancer. This examination only confers benefit on the screened population if it detects breast cancer at an early stage, where the prognosis is improved. This can only be achieved by having high quality mammography which is capable of detecting small lesions in the breast. High quality mammography must be achieved and maintained by a rigorous and comprehensive quality assurance and control programme.

The general principles of screening are (1):

1) The condition screened for should pose an important health problem.
2) The natural history of the condition should be well understood.
3) There should be a recognisable latent or early stage.
4) Treatment of disease at an early stage should be of more benefit than treatment started at a later stage.
5) There should be a suitable test or examination.
6) The examination should be acceptable to the population.
7) For diseases with an insidious onset, screening should be repeated at intervals determined by the natural history of the disease.
8) There should be adequate facilities available for the diagnosis and treatment of any abnormalities detected.
9) The chance of physical or psychological harm should be less than the chance of benefit.
10) The cost of funding (including diagnosis and subsequent treatment) should be economically balanced against the benefit it provides.

Screening, using x-ray mammography, for the detection of breast cancer at an early stage is a well establishes public health measure. In the National Health Service Breast Screening Programme, women between the ages of 50 and 70 are offered three yearly x-ray mammography. Screening above the age of 70 is available on request. Low energy x-rays are used to image the breast, in order to detected small, low contrast lesions in the breast. High image quality and low doses are demanded of x-ray mammography, as smaller cancers have a much better prognosis.

The basic principles of radiation protection are justification and optimisation. Justification applies to both the population and at an individual level. For the screened population it requires that the benefit, in terms of additional lives saved, are greater than the risk from the use of ionising radiation. In a screening programme, optimisation implies that the benefit is maximised by improving image quality and hence the cancer detection rate. This demands that screening mammography is subject to a comprehensive quality control programme.

2 Mammography

Mammography is an X-ray examination of the breast that requires specialised imaging equipment and techniques. The low inherent radiation contrast between fat and glandular tissue necessitates the use of specially filtered X-ray beams generated in an X-ray tube with a special target at a tube potential in the range of 28-32 kV. The X-ray tube must use a small focal spot (e.g. 0.1 to 0.4 mm). The most commonly used tubes have a molybdenum target...
with a 30 µm molybdenum filter. For thick, dense breasts tungsten and rhodium target X-ray tubes with appropriate beam filters may provide advantages (2).

In order to minimise radiation dose and to reduce the effect of scattered radiation on the film, the breast must be compressed. The mammography unit may also have a moving or stationary grid. In general, radiographs are acquired using a single emulsion film placed in an X-ray cassette with a single back screen to optimise image detail. Specialised, preferably dedicated, film processing is also desirable.

3 Quality Assurance and Quality Control

As stated by the WHO quality assurance in diagnostic radiology is: “All those planned and systematic actions necessary to provide adequate confidence that a structure, system or component will perform satisfactorily in service (3). Satisfactory performance in service implies the optimum quality of the entire diagnostic process, i.e. the consistent production of adequate diagnostic information with the minimum exposure of both patients and personnel.” Thus the main objectives of a quality assurance (QA) programme are to improve diagnostic accuracy without unnecessary radiation and to minimise costs.

3.1 Quality Assurance Programme

The responsibilities for performing the various QA and quality control (QC) procedures are delegated to X-ray operators, radiographers, and medical or health physicists depending on the size of the facility.

Three levels of testing are usually performed.
1. Acceptance tests.
2. Status tests.
3. Constancy tests.

Acceptance tests are performed when equipment is purchased to ensure that it meets its contractual specification. Status tests are undertaken to determine the absolute performance of equipment and may be included in the acceptance test. The purpose of the constancy test is to monitor the consistency of performance of the equipment. What constitutes an acceptance test, status test or constancy test is dependent on the type of equipment. It is difficult to be prescriptive about QA and QC test methods and frequencies that are applicable in all situations in diagnostic radiology. An automatic film processor may require monitoring on a daily basis, whereas it may only be necessary to check the tube filtration when an X-ray tube has been replaced. Similarly, the tests performed on a new image intensifier fluoroscopy unit will differ from those undertaken if the consideration is whether an old unit is to be taken out of service or not. Advice on test frequencies for mammographic QC testing is given in Appendix 1 which is adapted from a previously published protocol (4).

The Basic Safety Standards (5) state that registrants and licensees shall establish a comprehensive quality assurance programme for medical exposures with the participation of qualified experts in appropriate fields (i.e. radiodiagnostic physics), “taking into account the principles established by the WHO and the PAHO”.

In a manual of this kind it is impractical to describe in detail QA and QC tests that should be performed on all types of diagnostic radiology equipment. There have been numerous publications on QA and QC test methods and it is suggested that the reader refer to these manuals for further guidance.
Appendix 1

Test Frequencies

This Appendix lists all the tests in the order given in the protocol, together with the suggested frequencies at which they might be undertaken. The list may not be exhaustive, but will certainly be exhausting and readers may need to be selective. Frequencies should be regarded as tentative and may need to be altered in the light of experience or according to local circumstances. All the tests are regarded as being part of the commissioning process; a frequency is not shown if the test does not need to be repeated after commissioning. Some of the safety tests may need to be repeated more frequently for equipment fitted in mobile trailers. Repairs and maintenance may necessitate additional tests.

(Key: D=Daily to Weekly, M=3 to 6 Monthly, A=Annually)

Electrical Safety

Mechanical Safety
1. Table movement prevented under compression A
2. Compression auto-release M
3. Auto-release override M
4. Emergency release M
5. Maximum compression force M
6. No sharp edges A
7. Field light A
8. Screen edges marked -
9. Adequate retraining devices on mobiles A

Mechanical Functioning
1. Equipment complete A
2. Markings A
3. Free movements M
4. Brakes A
5. Scale markings A
6. Vertical movement A
7. Foot switches A
8. Attachments A
9. Field sizes marked A
10. AEC detector A
11. Cassette movement A
12. Cassette interlock A
13. Light intensity A
14. Compression plate movement M
15. Breast thickness scale A

Radiation Safety Inspection
1. Mains isolator position -
2. Clear control markings A
3. Mains-on light A
4. X-rays-on light A
5. Total filtration A
6. Added filter interlock A
7. Diaphragm interlock A
8. Exposure termination A
9. Exposure control position/lead A
10. Exposure control design -
11. Exposure control function A
12. Entrance warning light       A
13. Lead equivalence markings    -
14. Lead equivalence             -
15. Protective screen gap        A
16. Visibility                   A

Radiation Safety Measurements
- Tube leakage                    (a)
- Lead equivalence of screen
- Table transmission
- Separation of film/table edge   A
- Alignment of x-ray field to film/cassette   M
- Alignment of light/x-ray field
- Additional checks for mobiles      As required

X-ray Measurements
- X-ray field non-uniformity       A

Dimensions of focal spot
- Slit camera                      (b)
- Star resolution grid             M
- Pinhole                          As required

Tube kilovoltage
- Brief check                      M
- Full check                       A
- HVL filtration                   A
- Exposure time                    A

Output
- Consistency                      M
- With change in kV                 M
- With change in tube current/focus A

Magnification
- Grid factor/grid system factor   -
- Grid film                        A

Automatic Exposure Control
- Consistency                      M
- Sensitive area of AEC detector   -
- Phantom thickness                M
- Tube voltage                     M
- Tube current                     A
- Other parameters                 A
- Calibration of density control   A
- Guard timer                      A
- Regular test                     D

Automatic processing unit (APM)
- Sensitometry                      D
- APU temperature                  D
- Transport speed                  As required
- Replenishment rate               As required
- Specific gravity/pH               As required
- Residual hypo                     As required
- Silver recovery                   As required

Screen-film system
- Cassette and screen identification M
- Screen-film contact               M
- Light-tightness of cassette       M
- Relative sensitivity of screen-cassette M
- Characteristic curve of screen-film As required
Dark room and film storage
- Light-tight darkroom: A
- Safelights/warning lights: A
- Temperature: A
- Humidity: A
- Stock control: D

Illuminators and viewing room
- Visual check: M
- Illuminator light level: A
- Ambient light level: A

Breast dose
- Dose to standard breast: M
- Alternative method: A
- Routine monitoring: D

Image quality
- Optimisation: As required
- Routine check: D
- Stereotactic systems: A

Specimen x-ray cabinets
- All tests: A

Notes:
(a) = 3 yearly
(b) = may need to be checked more frequently
5 References


International Radiation Protection Association

11th International Congress

Madrid, Spain - May 23-28, 2004

RC-4a Screening Mammography
Including Quality Assurance

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NHSBSP

- Women aged 50 – 70 years screened
- X-ray mammography
- Two views all rounds
- Mainly double read
High Quality Images Required

This means rigorous QA
Breast Cancer Detection Rates (per 1,000 England)

<table>
<thead>
<tr>
<th>Age Band</th>
<th>94-95</th>
<th>95-96</th>
<th>96-97</th>
<th>97-98</th>
<th>98-99</th>
<th>Mean</th>
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<td>6.1</td>
<td>6.1</td>
<td>6.8</td>
<td>6.2</td>
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</table>
A quality assurance program may be defined (WHO definition) as an organized effort by the staff operating a facility to ensure that the diagnostic images produced by the facility are of sufficiently high quality so that they consistently provide adequate diagnostic information at the lowest possible cost and with the least possible exposure of the patient to radiation.
QUALITY ASSURANCE PROGRAMS (II)

- Radiology imaging equipment should produce images that meet the needs of the radiologist or other interpreters without involving unnecessary irradiation of the patient.
- Quality assurance actions contribute to the production of diagnostic images of a consistent quality by reducing the variations in performance of the imaging equipment.
QA Objectives

- The aim of quality assurance in the breast screening programme is the maintenance of minimum standards and the continuous improvement in performance.
QA Objectives

- To review the performance and outcomes of breast screening and individual units
- To provide advice and continuing professional education for individuals
- To support health authorities and trust in the specification, commissioning and delivery of screening to meet national standards
QA Team

- Professional members are appointed with a clear job description and a paid commitment
- Accountable to the regional QA director
- QA Director is accountable to the RDPH
QA Visit

Visiting Team

- Radiology
- Radiography
- Pathology
- Surgery
- Breast care nursing
- Administration and clerical
- Medical physics
National QA Guidelines

- NHSBSP documents
- Revised Pritchard standards
- Professional QA guidance
- QA visit protocol
QA Visit

- Verifies the achievement of national standards and identify variance from these standards
- Support professionals working in the programme to maintain and improve standards of professional performance
QA Visit

- Take place every three years
- Multidisciplinary
- Take place in the breast screening unit
Comment on screening outcomes and interval cancer rates

Identify strengths and weaknesses in the unit

Recommend actions and a timescales for their implementation
Incident Investigation

- QA Director informed of potential incident
- Set up a team to determine nature and extent of incident
- Follow NHSBSP protocol
- Establish if it is an incident
- Hand it over to the Trust
- Keep RDPH and National Office informed
Physics QA Programme

- Acceptance testing
- Constancy testing
- Status testing
Electrical Safety

- Responsibility of the supplier
- IEC 601-1
- Department of Health TRS 89 Technical requirements for the supply and installation of apparatus for diagnostic imaging and radiotherapy
Mechanical Safety

- Prevention of powered movement under compression
- Automatic release of compression plate after an exposure
- 200N maximum powered compression force
- No sharp edges
Marking and Labelling

- Focal spot size and position
- Inherent, added and total filtration
- AEC position
- Magnification factor
- Function of all controls
Mechanical Function Checks

- All manually controlled movements
- Mechanical/electromechanical brakes
- Scales/indicators
- Beam diaphragms
- Foot switches
- Attachments
- AEC position selector
Radiation Safety

- Mains isolator switch
- Mains on light
- Exposure light
- Total filtration 0.5mmAl/0.03mmMo
- Diaphragm interlock
- Exposure termination if button released
Alignment

- Light field to X-ray field
- X-ray field to film
- Field edge and edge of breast support platform
- Imaged area for digital systems
X-ray field to film

Alignment >0mm and <5mm
Compression Force

- Maximum force $>150\text{N}$ and $<200\text{N}$
- Thickness indicator accurate to 5mm
$l_\omega = l\left[\tan \phi - \tan \omega\right]/\left[\tan \phi - \tan \theta\right]$
Measurement Methods

- Slit camera $f = \frac{d}{(M-1)}$
- Pin hole
- Star pattern $f = \pi \theta \times \frac{D}{180(M-1)}$
Tube Voltage Measurement

- Accurate to 1 kV
- Remedial level 2 kV
- Digital kV meter
HVL Measurement

Compression Paddle
Absorbers

Focus to Detector Distance of 40-55 cm

Detector

HVL > 0.3 mmAl, < 0.4 mmAl
Estimated Tube Filtration

Half-value layer (mm Al)

Added filtration (μm Mo)
Tube Output

- Repeatability
- Specific output
- Specific output rate
- Variation of output with kV
- Variation of output with mAs
X-ray Uniformity

Diagram:
- 24 cm
- 18 cm
- 12 cm
- 10 cm
- 4 cm

Diagram:
- 30 cm
- 24 cm
- 15 cm
- 10 cm
- 4 cm
AEC System

- Target density (1.5-1.9)
- Repeatability (5%)
- Constancy with change in thickness (>0.2, range >0.3)
- Constancy with change in kV (>0.2, range >0.3)
- Density control
- Guard timer
- Exposure timer (>1s)
Sensitometric Curve

B = base + fog
Speed = D1 - B
C = D2 - D1
Processing Control Charts

Base + fog
Processing Control Charts
Processing Control Charts
Effect of Delay in Processing

![Graph showing the effect of delay in processing on relative film speed and optical density.](image)

- **Relative film speed**
- **Optical density, 4 cm perspex**

The graph illustrates the decrease in relative film speed and optical density over time after exposure, highlighting the impact of delay in processing.
Automatic Film Processor

- Sensitometry
- Temperature (0.5 C)
- Speed (5%)
Film Cassette

- Screen-Film contact
- Sensitivity (0.05)
Illuminators

- Subjective visual check
- Luminance (3000 cdm\(^{-2}\))
- Luminance (variation <15% between panels)
- Ambient light (50 lux)
Breast Dose

\[ D_{\text{old}} = K_{\text{pgs}} \]

\( p \) converts incident air Kerma \( K \) for perspex phantom to that of the standard breast.

\( g \) converts incident air Kerma for standard breast to mean glandular dose.

\( S \) is a spectral conversion factor.
Breast Dose

\[ D_{\text{new}} = K_{45} \cdot g_{53} \cdot c_{53} \cdot S \]

- \( K_{4.5} \) is the entrance air Kerma for 4.5cm perspex
- \( G_{5.3} \) is the g factor for 5.3cm standard breast
- \( C_{5.3} \) is the glandularity factor for 5.3cm
- \( S \) is a spectral conversion factor
## Conversion Factors

<table>
<thead>
<tr>
<th>HVL (mm Al)</th>
<th>$g$ (mGy/mGy)</th>
<th>$c$</th>
<th>product of $g$ and $c$</th>
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<tr>
<td>0.30</td>
<td>0.155</td>
<td>1.109</td>
<td>0.172</td>
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<tr>
<td>0.35</td>
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<td>0.60</td>
<td>0.295</td>
<td>1.088</td>
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</table>
Mean Glandular Dose Standard Breast

Air Kerma \( (K) = T \cdot mA_{\text{exp}} \cdot (50/d)^2 \)

The mAs/exposure is determined using the perspex phantom

\( T \) is the tube output at 50cm

\( D \) is the focus phantom distance
Mean Glandular Dose
Real Breasts

\[ D = K g c s \]

K is the incident air Kerma at the upper surface of the breast

g is the glandularity conversion factor

c converts from 50% glandularity

s is the spectral conversion factor
## Conversion Factors g

<table>
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<tr>
<th>Breast Thickness cm</th>
<th>HVL mm Al</th>
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<th>0.35</th>
<th>0.40</th>
<th>0.45</th>
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## Conversion Factors g

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## Conversion Factor $c$

### 50-64 years

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<th>HVL (mm Al)</th>
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<td>1.307</td>
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<tr>
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Standard Breast

- 50:50 adipose/glandular tissue, superficial region of 0.5cm adipose tissue
- 4.5 cm compressed thickness
- Area 100cm²
## Conversion Factors

<table>
<thead>
<tr>
<th>HVL (mm Al)</th>
<th>p</th>
<th>g (mGy/mGy)</th>
</tr>
</thead>
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<td>0.25</td>
<td>1.12</td>
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<tr>
<td>0.30</td>
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<td>0.183</td>
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</tr>
<tr>
<td>0.50</td>
<td>1.09</td>
<td>0.285</td>
</tr>
<tr>
<td>0.55</td>
<td>1.07</td>
<td>0.311</td>
</tr>
<tr>
<td>0.60</td>
<td>1.06</td>
<td>0.339</td>
</tr>
</tbody>
</table>
## Conversion Factors

<table>
<thead>
<tr>
<th>Spectrum</th>
<th>$s$-factor</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mo/Mo</td>
<td>1.000</td>
</tr>
<tr>
<td>Mo/Rh</td>
<td>1.017</td>
</tr>
<tr>
<td>Rh/Rh</td>
<td>1.061</td>
</tr>
<tr>
<td>Rh/Al</td>
<td>1.044</td>
</tr>
<tr>
<td>W/Rh</td>
<td>1.042</td>
</tr>
</tbody>
</table>
TOR (MAM)

Six groups of filaments in a range of diameters. Filaments are placed parallel, perpendicular and at 45° to the anode-cathode axis.

Calcium-based particles in six size ranges.

Low-contrast details, 3 mm diameter. Six contrast values in mixed groups of three.

Background density

Simulated breast tissue including 'micro-calcifications'
Contrast Detail Diagram

Graph showing the relationship between threshold contrast (%) and detail size (mm) for typical analogue and digital systems.
Image Quality CDMAM Phantom Detection Index

![Graph showing detection index versus detail size for typical analogue and digital systems.](image)

- **Typical analogue system**
- **Typical Digital System**
CDMAM Test Object
TOR(MAX)
High Contrast Resolution

![Graph showing the proportion of systems parallel to the tube axis and perpendicular to the tube axis at different resolutions (lp/mm)].

- **Proportion of systems**
  - Parallel to the tube axis
  - Perpendicular to the tube axis

- **Resolution (lp/mm)**: 7.1, 8, 8.9, 10, 11.1, 12.5, 14.3, 16.6, 20
Threshold Contrast Measurements (0.25mm)
Threshold Contrast (0.5mm)

NHSBSP Standard
≥ 8 details

number of 0.5mm details seen
Stereotactic Assessment
## Performance Measurements

<table>
<thead>
<tr>
<th>Parameter</th>
<th>minimum</th>
<th>25th percentile</th>
<th>mean</th>
<th>75th percentile</th>
<th>maximum</th>
<th>unit</th>
</tr>
</thead>
<tbody>
<tr>
<td>measured kV at 28 kV set</td>
<td>26.9</td>
<td>27.7</td>
<td>27.9</td>
<td>28.2</td>
<td>29.5</td>
<td>kVp</td>
</tr>
<tr>
<td>output at broad focus at 28 kV Mo/Mo (at 50 cm)</td>
<td>136</td>
<td>182</td>
<td>201</td>
<td>220</td>
<td>282</td>
<td>Gy/mAs</td>
</tr>
<tr>
<td>output per second at FFD at 28 kV Mo/Mo</td>
<td>7.81</td>
<td>11.5</td>
<td>15.5</td>
<td>19.2</td>
<td>29.2</td>
<td>mGy/s</td>
</tr>
<tr>
<td>HVL at 28kV with compression paddle</td>
<td>.30</td>
<td>.34</td>
<td>.35</td>
<td>.365</td>
<td>.41</td>
<td>mm Al</td>
</tr>
<tr>
<td>broad focus width</td>
<td>.11</td>
<td>.30</td>
<td>.35</td>
<td>.39</td>
<td>.63</td>
<td>mm</td>
</tr>
<tr>
<td>broad focus length</td>
<td>.19</td>
<td>.42</td>
<td>.53</td>
<td>.62</td>
<td>1.09</td>
<td>mm</td>
</tr>
<tr>
<td>fine focus width</td>
<td>.05</td>
<td>.10</td>
<td>.13</td>
<td>.16</td>
<td>.45</td>
<td>mm</td>
</tr>
<tr>
<td>fine focus length</td>
<td>.05</td>
<td>.12</td>
<td>.16</td>
<td>.19</td>
<td>.37</td>
<td>mm</td>
</tr>
<tr>
<td>Separation between film edge and table edge</td>
<td>1</td>
<td>3</td>
<td>3.2</td>
<td>4</td>
<td>6</td>
<td>mm</td>
</tr>
<tr>
<td>Overlap of X-ray field at chest wall edge</td>
<td>-2</td>
<td>1</td>
<td>2.0</td>
<td>3</td>
<td>11.2</td>
<td>mm</td>
</tr>
<tr>
<td>Maximum compression force (automatically applied)</td>
<td>90</td>
<td>150</td>
<td>170</td>
<td>190</td>
<td>255</td>
<td>N</td>
</tr>
<tr>
<td>AEC consistency (a)</td>
<td>.00%</td>
<td>.01%</td>
<td>.75%</td>
<td>1.01%</td>
<td>7.7%</td>
<td>%</td>
</tr>
<tr>
<td>AEC error with 2 cm Perspex (b)</td>
<td>-3.5</td>
<td>-.09</td>
<td>-.03</td>
<td>.02</td>
<td>0.22</td>
<td>OD</td>
</tr>
<tr>
<td>AEC error with 6 cm Perspex (b)</td>
<td>-.70</td>
<td>-.08</td>
<td>-.02</td>
<td>.05</td>
<td>0.35</td>
<td>OD</td>
</tr>
<tr>
<td>film density for 4cm Perspex using AEC at clinical settings</td>
<td>1.31</td>
<td>1.58</td>
<td>1.66</td>
<td>1.74</td>
<td>2.23</td>
<td>OD</td>
</tr>
<tr>
<td>Exposure times for 4cm Perspex</td>
<td>.16</td>
<td>.32</td>
<td>.53</td>
<td>.64</td>
<td>1.74</td>
<td>s</td>
</tr>
<tr>
<td>Exposure times for 6cm Perspex</td>
<td>.58</td>
<td>1.21</td>
<td>1.78</td>
<td>2.22</td>
<td>4.12</td>
<td>s</td>
</tr>
<tr>
<td>mean glandular dose to old standard breast (at 28 kV Mo/Mo)</td>
<td>.65</td>
<td>1.13</td>
<td>1.36</td>
<td>1.61</td>
<td>2.60</td>
<td>mGy</td>
</tr>
<tr>
<td>mean glandular dose to old standard breast (at clinical settings)</td>
<td>.69</td>
<td>1.20</td>
<td>1.40</td>
<td>1.56</td>
<td>2.60</td>
<td>mGy</td>
</tr>
<tr>
<td>high contrast resolution parallel to tube axis (c)</td>
<td>10.0</td>
<td>13.0</td>
<td>14.0</td>
<td>14.5</td>
<td>20.0</td>
<td>lp/mm</td>
</tr>
<tr>
<td>high contrast resolution perpendicular to tube axis (c)</td>
<td>11.0</td>
<td>15.0</td>
<td>16.1</td>
<td>17.0</td>
<td>20.0</td>
<td>lp/mm</td>
</tr>
</tbody>
</table>