RADIATION SAFETY ACT 1975

DIAGNOSTIC X-RAY EQUIPMENT COMPLIANCE TESTING

PROGRAM REQUIREMENTS

2015

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DIAGNOSTIC X-RAY EQUIPMENT
COMPLIANCE TESTING

PROGRAM REQUIREMENTS

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1. PROGRAM OVERVIEW

1.1 INTRODUCTION

Commencing 1 January 1997, the Radiological Council introduced a program requiring the periodic testing of diagnostic x-ray equipment for compliance with the State’s Radiation Safety (General) Regulations and any additional criteria that the Council may apply to the equipment under test.

The need for a program of this type was indicated by —

- studies in Australia and elsewhere demonstrating that poor x-ray equipment performance is a significant contributor to unnecessary patient radiation exposure;
- a trend towards mandatory quality assurance (QA) programs for medical radiology;
- evidence of significant non-compliance documented in inspections conducted by Council’s officers.

The tests required are those primarily concerned with radiation safety. However, as patient radiation dose ultimately depends on the total imaging process, these tests should be supplemented by additional radiographic, sensitometric and image quality tests as part of a complete QA program.

The processes involved in the compliance testing program are outlined in Appendix 1. An explanation of the terms used in this document is given in Appendix 9.

1.2 EQUIPMENT SUBJECT TO TESTING

The program applies to all diagnostic x-ray equipment used on live humans for medical radiography and fluoroscopy, chiropractic and dental radiography and computed tomography.

1.3 FREQUENCY OF TESTING

The program initially required annual testing of medical and chiropractic x-ray equipment and triennial testing of dental intra-oral and panoramic equipment. However, following consideration of the first three years test results and consultation with representatives of equipment owners, users and others, the Council amended the test frequencies to those shown below. These frequencies took effect from 12 October 2000.

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1 The Radiological Council is an independent statutory body established under the Radiation Safety Act (see Section 10) for the purpose of regulating radiation safety in Western Australia. It reports directly to the Minister for Health.
Mammography 12 months  
C-arm or U-arm fluoroscopy (fixed or mobile) 12 months  
Cone Beam CT 12 months  
Other fluoroscopy 24 months  
Radiography 24 months  
CT 24 months  
Dental (intraoral, panoramic tomography, and equipment with combined panoramic tomography and cephalometry) 36 months

The Council will continue to keep test frequencies under review and may adjust them further should the need arise.

All x-ray equipment used on humans within the scope of the program (see section 1.2) must be tested and issued with a Compliance Certificate or a Conditional Compliance Certificate before it may be used on patients, unless otherwise exempted by the Radiological Council.

1.4 TEST ANNIVERSARY DATE

The continued use of x-ray equipment subject to the compliance testing program after the expiry date of its current compliance certification is an offence. The Radiological Council also may issue a formal direction to cease use of the equipment.

If the new certification expiry date is calculated from the test date, those who have their equipment tested before the current expiry date do not enjoy the full extent of their current compliance period. A fixed anniversary date has thus been adopted, according to which expiry dates are calculated as follows:

<table>
<thead>
<tr>
<th>Equipment test date</th>
<th>New expiry date</th>
</tr>
</thead>
<tbody>
<tr>
<td>Equipment tested not more than three months before the current expiry date</td>
<td>The current expiry date plus one test period</td>
</tr>
<tr>
<td>Equipment tested more than three months before the current expiry date</td>
<td>The test date plus one test period</td>
</tr>
<tr>
<td>Equipment tested not more than six months after the current expiry date</td>
<td>The current expiry date plus one test period</td>
</tr>
<tr>
<td>Equipment tested more than six months after the current expiry date</td>
<td>The test date plus one test period</td>
</tr>
</tbody>
</table>

As an example of the application of the above, take the current compliance certification expiry date of a dental x-ray machine as 1 September 2005. The owner has the machine tested on 1 June 2005. As the equipment was tested
not more three months before the expiry date, the new expiry date is 1 September 2005 plus 36 months, which is 1 September 2008.

1.5 RESPONSIBILITY FOR TESTING

Under Section 28 of the Act, the ‘owner’ of x-ray equipment, or the person having possession, must apply for its registration. Through conditions imposed on the registration under Section 36 of the Act, the registrant is legally responsible for satisfying the requirements of the compliance testing program. A copy of typical registration conditions (in this case for medical radiology) is given in Appendix 2.

Registrants must ensure that their x-ray equipment is tested by a licensed compliance tester at the prescribed frequency. Tests need to be scheduled in a timely manner to ensure that new certificates can be issued before earlier certificates expire.

A list of licensed compliance testers is available from the Radiological Council.

The conditions, restrictions and limitations imposed on registrations under Section 36 of the Radiation Safety Act make it an offence to operate or use x-ray equipment for human diagnostic radiography or fluoroscopy unless it has —

(a) a current Certificate of Compliance (see section 2.3); or
(b) a current Certificate of Conditional Compliance (see section 2.4);

‘Current’ means that the certificate was issued within the previous 12, 24 or 36 months (depending on the prescribed test frequency stated in section 1.3).

Examples of compliance and conditional compliance certificates are given in Appendix 3.

1.6 TEST PROTOCOLS

The prescribed tests assess compliance with the Radiation Safety (General) Regulations and with any other Council requirements that may apply to the equipment under test.

A series of workbooks developed by a working group of the Council describe the tests to be undertaken for each category of equipment as well as the approved test protocols. These workbooks are kept under review and updated from time to time.
Currently available are:-

Workbook 1 – Mobile Radiographic Equipment
Workbook 2 – Mammographic Equipment
Workbook 3 – Major Radiographic Equipment
Workbook 4 – Fluoroscopic Equipment
Workbook 5 – Dental Radiographic Equipment
Workbook 6 – Computed Tomography Equipment

Digital radiographic versions of the relevant workbooks\(^1\) have been issued in draft format for comment, and are:-

Workbook 7 – Mobile Digital Radiographic Equipment
Workbook 8 – Digital Mammographic Equipment
Workbook 9 – Major Digital Radiographic Equipment
Workbook 10 – Digital Fluoroscopic Equipment
Workbook 11 – Dental Digital Radiographic Equipment

Workbooks can be obtained on a cost-recovery basis from the Council through the Radiation Health Branch of the Department of Health of Western Australia.

This particular publication, the Program Requirements, is available free of charge.

Variations to the recommended test protocols may be used provided they are approved by the Radiological Council. Documentation supporting alternative test protocols must be provided to the Council.

Details of the required tests are given in section 5.

1.7 TEST PERSONNEL

Compliance testing may only be undertaken by a person who holds a licence under the Radiation Safety Act for that purpose or by a person acting under the direction and immediate personal supervision\(^2\) of a compliance testing licensee.

All tests must be assessed and certified by a qualified expert. The licensed compliance tester must ensure that the completed test report is submitted to a qualified expert within 1 month.

A qualified expert is eligible for a compliance testing licence and may both

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\(^1\) Final versions not approved by the Council at the time of publication of these Program Requirements.

\(^2\) ‘Immediate personal supervision’ requires the licensee to be physically present and to observe directly persons working under their direction and supervision.
carry out and certify tests without independent certification of the test results.

The requirements for approval as a qualified expert and the prerequisites for applicants applying for compliance testing licences are given in sections 3 and 4 respectively.

**Note:** Persons who undertake compliance testing concurrently with the service or maintenance of x-ray equipment should take particular note of section 2.8.

### 1.8 ACCURACY SPECIFICATION

Whilst the purchase of measuring equipment such as non-invasive x-ray beam analysing instruments, ion chambers etc is a matter of choice for the tester, it is a requirement that the accuracy of such equipment for the relevant parameter is ± 3% or better.
2. DOCUMENTATION, CERTIFICATION AND ASSESSMENT OF COMPLIANCE TESTS

The Radiological Council, at its discretion, may at any time reject and invalidate any compliance test or part thereof, and may require the withdrawal of any certificates, labels or notices prescribed by this program which it finds has or may have been issued on the basis of unapproved test methods, invalid, incomplete or inaccurate test data, or other information which cannot be substantiated.

2.1 TEST REPORTS

In keeping with the prescribed test frequencies and when requested by the registrant (owner), a licensed compliance tester will carry out all the tests required by the Council using the protocols described in the relevant workbook or protocols otherwise approved by the Council (see section 1.6).

- To standardise reporting, test report forms are provided in each workbook. These may be reproduced as necessary. Testers may style forms to suit their own needs and record keeping systems. However, copies of test reports provided to the Council must be on the standard form included in the relevant workbook, or on any updated versions subsequently approved by the Council.

- The compliance tester must provide the qualified expert with all the test results including copies of relevant x-ray films or images, waveforms and computer printouts.

- All x-ray equipment faults determined during testing must be reported even if such faults are corrected before testing is completed. Where an item of non-compliance is corrected during the test, both the pre- and post-correction test data must be included in the report.

Compliance tests that cannot be completed because of faults with the x-ray equipment must also be reported. See also section 2.8.

2.2 CERTIFICATION AND ASSESSING

All compliance test reports must be signed initially by the tester. The test report must also be signed by the qualified expert after assessing the report.

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1 The ‘standard form’ for any report can be supplied to the tester by email. The term ‘standard form’ comprises the individual items of the report sheets, and the order in which they are listed, as shown in the workbooks, but excludes the detailed formats of the items. For example, the tester may replace ☑ with [✓], fonts or spacing may be changed etc, but additions, deletions or rearrangements of the order of identifying codes, version numbers, questions, results or other items will not be accepted. Council approval of any changes made by the tester is required prior to use of the form.
The signed test report will be used by the qualified expert as the basis for issuing, or not issuing (as the case may be), a Compliance Certificate for the equipment tested.

Note: A test report bearing only a tester’s signature indicates that the compliance test has been conducted in accordance with the approved test protocol, that the identifying information stated is correct and that the measurements obtained are accurate within the accepted errors inherent in the measuring equipment. It does not necessarily indicate that the equipment complies with the Radiological Council’s requirements.

A copy of the assessed and signed test report, together with a copy of the Compliance Certificate, must be forwarded within 2 months of the date the test commenced to —

The Secretary  
Radiological Council  
Locked Bag 2006 P O  
NEDLANDS WA 6009

The registrant, compliance tester and qualified expert should also retain copies.

Council officers will review the test reports and use the data to monitor the performance of the program. The review process may include re-testing of equipment by Council officers either on a random basis or to provide clarification of matters arising from a specific report.

2.3 CERTIFICATE OF COMPLIANCE

A Certificate of Compliance will be issued by a qualified expert when he or she is satisfied that the x-ray equipment fully satisfies the assessment criteria of the compliance test.

The certificate is valid for a period of 1, 2 or 3 years from the commencement date of the full test depending on the test frequency established for the particular equipment category (See section 1.3).

The Radiological Council’s Certificate of Compliance is the only form of approval that may be issued. Each certificate is individually numbered and must not be duplicated (by any means) until it has been completed and signed by a qualified expert.

A copy of the completed certificate must be forwarded to the Council together with a copy of the signed test report. The registrant should be given the
original compliance certificate with copies retained by both the qualified
expert and the compliance tester.

Registrants must retain a copy of the compliance documentation for all
equipment in their possession. In the case of equipment that is sold or
transferred to another owner all current compliance documents, including the
certificate, must be handed to the new owner. This does not apply when the
equipment is transferred to another site or practice belonging to the same
registrant. However, in all cases when disassembly and reassembly of
equipment that might affect its compliance status is involved, tests must be
undertaken to ensure that compliance is not adversely affected during this
process. In such circumstances, a full compliance test must be undertaken.
The registrant is responsible for ensuring the equipment remains compliant.

A sample Certificate of Compliance is shown in Appendix 3.

2.4 CERTIFICATE OF CONDITIONAL COMPLIANCE

Conditional Compliance may be issued for x-ray equipment which does not
comply fully with all program requirements but for which, in respect of the
non-complying item(s), the Council is satisfied that there is no significant
radiation risk to either patients or users.

Conditional Compliance may be applicable for:

a) x-ray equipment that was first registered before the introduction of the
compliance testing program (1 January 1997) and where its design
and/or performance could not be modified, adjusted or repaired to
comply fully with the program requirements; and

b) x-ray equipment that was issued with a full Compliance Certificate prior
to the current compliance test but which, due to 'wear and tear' or the
unavailability of parts or components, no longer fully complies and can
no longer be modified, adjusted or repaired to meet the compliance
requirements.

Conditional Compliance will not be issued if a fault directly or indirectly
relates to:

• x-ray beam quality (half value layer);
• radiographic x-ray / light beam congruency or x-ray beam dimensions;
• maximum permitted fluoroscopic dose rates;
• radiation leakage from the x-ray tube assembly (including the
collimator);
• x-ray exposure warning devices (including the fluoroscopic timer);
• interlocks which are intended to prevent x-ray exposures when
prescribed safe operating conditions are not met;
• patient dose / dose-rate limits or constraints that are prescribed by regulations or registration conditions;
• exposure reproducibility;
• any other matter that in particular circumstances the Council considers significant.

Conditional compliance is not transferable should the x-ray equipment be sold (as defined in the Act) and may be void if the x-ray equipment is relocated.

Depending on the nature of the non-compliant item(s), Council may stipulate a ‘use by’ date beyond which Conditional Compliance will not be extended. In such circumstances, the Council may give the registrant up to 6 years to plan for the equipment’s replacement or its withdrawal from service. This period may be varied on a case-by-case basis on application by the registrant. The expiry date of this period will be stated on the certificate, together with the item(s) of non-compliance.

Regular compliance testing at the prescribed frequency will still be required for the class of equipment.

The Radiological Council’s Certificate of Conditional Compliance is the only form of approval that may be issued. Each certificate is individually numbered and must not be duplicated (by any means) until it has been completed and signed. The certificate may be withdrawn or varied at any time by the Council.

Conditional Compliance is authorised by the Council and therefore will only be issued on application and on a case-by-case basis.

Registrants must retain a copy of the compliance documentation for all equipment in their possession.

A sample Certificate of Conditional Compliance is shown in Appendix 3.

### 2.5 COMPLIANCE LABELS

X-ray equipment that has been tested and certified to comply must be labelled. Only the labels approved by the Radiological Council may be used.

For full compliance, yellow self adhesive labels identifying the compliance certificate expiry date, the certificate number and the name of the tester and qualified expert are issued with the compliance certificate.

For conditional compliance, magenta self adhesive labels are issued identifying the registrant, their registration number, the compliance certificate expiry date and the certificate number.
The label for the specified x-ray equipment must be fixed in a conspicuous position on the control panel. Sample labels are shown in Appendix 4.

2.6 NON-COMPLYING EQUIPMENT

Section 2.4 deals with non-complying equipment that may be eligible for a Certificate of Conditional Compliance. However, non-complying equipment that is ineligible for conditional compliance must be repaired or modified to correct the identified faults (see also 2.7).

Immediate response to avoid unacceptable radiation exposure

Should a qualified expert believe that continued use of the x-ray equipment may place patients, the public or users at risk of significant and unnecessary radiation exposure, he or she should promptly inform the Council and provide acceptable supporting evidence for their belief. Council’s officers will review the evidence and give prompt consideration to issuing an immediate direction to the registrant to cease use of the equipment until the identified item(s) of non-compliance are corrected.

Notices to Cease Use (Appendix 5) may only be issued by an authorised person e.g. the Secretary to the Radiological Council.

Response for non-urgent matters

If the need for correction is not urgent, the qualified expert will issue a Notice of Non-Compliance to the registrant listing items that must be corrected within a 3 month period in order for a compliance certificate to be issued (unless an alternative time period is stipulated by the Council).

In these situations, a certificate will not be issued for the equipment until the items of non-compliance have been corrected. However, the qualified expert’s written Notice of Non-Compliance identifying the items requiring correction, together with the date by which those actions must be taken, will entitle the registrant to a period of temporary exemption from compliance.

Note: Irrespective of the expiry date of the Notice of Non-Compliance, the new Compliance Certificate shall be given an expiry date derived from that of the previous Compliance Certificate expiry date plus the prescribed test frequency or, for a first test, from the date the test commenced plus the prescribed test frequency (section 1.4).

Corrective actions are the responsibility of the registrant. However, the qualified expert and the compliance tester should liaise with the registrant to ensure that non-complying items are corrected in a timely manner and, if necessary, re-tested before a compliance certificate is issued.
Notices of Non-Compliance must be provided to the registrant by the qualified expert within 1 month of receipt of the test report, with a copy to the Radiological Council. An example of the Notice of Non-Compliance is provided in Appendix 5.

**New x-ray equipment which does not comply**

New x-ray equipment with items of non-compliance that cannot be corrected may either receive an exemption for that item from the Council, or be refused registration.

It is an offence for x-ray equipment to be sold unless it complies with the regulations or other criteria specified by the Radiological Council, or unless prior written approval has been given by the Council to the seller and purchaser. Sellers may be prosecuted for breaching the Act and registration of the equipment may be refused.

However, the Council recognises that changes in technology may supersede regulatory requirements. Provided such changes can be justified in the clinical or practical context and do not give rise to unacceptable radiation exposure, Council may consider an exemption under Section 6 of the Act in respect of the non-complying item(s) and approve sale of the equipment without further modification. If approval is given, the particular equipment would be exempted from the specific regulation or criterion (although additional design and/or performance criteria may be imposed in respect of the exempted feature) and normal compliance testing procedures will apply.

X-ray equipment for which an exemption has been granted and which otherwise complies fully with the program requirements will be issued with a Compliance Certificate.

Where an exemption has general application, the Council will consider amending both the program requirements and the Regulations to reflect Council’s decision.

### 2.7 EXEMPTIONS

A registrant who possesses x-ray equipment that cannot meet either full or conditional compliance requirements but which the registrant believes serves an ongoing clinical need, may make application to the Radiological Council for an exemption.
2.8 COMPLIANCE TESTING BY SERVICE PERSONNEL

Where compliance tests are performed by persons who are also licensed to service x-ray equipment, the compliance test report must show the performance of the equipment before any service is undertaken (unless a particular fault renders further testing invalid) i.e. compliance testing, if due, must precede any routine service and maintenance procedure.

Faults found during testing must be detailed on the report form even if they are corrected before completion of testing. Some faults may be common to the particular model of x-ray equipment and failure to report them may put other users and patients at risk.

2.9 CALIBRATION OF RADIATION MEASURING INSTRUMENTS

Compliance tests involving the use of radiation measuring instruments are valid only if suitable instruments are used and they have been calibrated not more than two years before the compliance test by an organisation recognised by the Radiological Council.
3. REQUIREMENTS FOR QUALIFIED EXPERTS

3.1 FUNCTIONS

Persons approved as qualified experts by the Radiological Council must have proven competencies in compliance testing and quality assurance procedures relevant to diagnostic medical imaging. A qualified expert who also intends carrying out compliance testing must hold a licence for the purpose.

The qualified expert shall —

- verify that all radiation measuring instruments used for the tests are suitable for the purpose and have been calibrated within the 2 years preceding the compliance test;
- verify that the compliance tests have been performed by a licensed tester according to the relevant workbook or by following other protocols approved by the Council;
- issue, as appropriate, either a signed Compliance Certificate and compliance label for display on the x-ray control panel, or a Notice of Non-Compliance, within 1 month of receiving the test report from the compliance tester;
- where relevant, provide copies of the assessed and signed test report and a copy of the compliance certificate to the Radiological Council within 1 month of the date of receipt of the report. (The Council may, at any time, request the provision of the full compliance test data.);
- where relevant, ensure that Notices of Non-Compliance are issued promptly to the registrant, and a copy forwarded to the Council within 1 month of the date of receipt of the report;

**Note:** Corrective actions are the responsibility of the registrant. However, the qualified expert and the compliance tester should liaise with the registrant to ensure that non-complying items are corrected in a timely manner and, if necessary, re-tested before a compliance certificate is issue.

3.2 ROLE AND RESPONSIBILITIES

The qualified expert —

- by signing a compliance test report is responsible for determining the compliance or otherwise of the equipment on the basis of the test data provided by the licensed compliance tester.
- must either personally perform the tests
or

- must liaise appropriately with the compliance tester and exercise a sufficient level of control to ensure that the protocols are followed and that the test results are reliable.

### 3.3 QUALIFICATIONS

A university degree in physics, engineering or a related science.

In exceptional circumstances a lesser qualification may be acceptable if in combination with demonstrated expertise in a health specific radiation discipline.

### 3.4 ASSESSMENTS

Passes will be required in —

- a written examination which tests knowledge of the physics of radiology, imaging technology and radiation protection; and

- a practical examination in compliance testing.

A syllabus for the written examination is given in Appendix 6.

Applications for approval as a qualified expert should be addressed to the Secretary of the Radiological Council enclosing a statement of the applicant’s qualifications and experience.

### 3.5 EQUIVALENT ACCREDITATIONS

The Radiological Council will accept as qualified experts persons who have gained Accreditation in Radiological Physics and Quality Assurance from the Australasian College of Physical Scientists and Engineers in Medicine.

Persons holding accreditations with other authorities should apply to the Council for consideration of those accreditations.

### 3.6 LICENSING

Persons who are approved as qualified experts are also eligible to apply for a compliance testing licence. Compliance testing licences are subject to the conditions given in Appendix 7.

Qualified experts who are also licensed compliance testers may sign and forward their own reports to the Radiological Council. Certification by an independent qualified expert is not required.
4. REQUIREMENTS FOR COMPLIANCE TESTERS

4.1 FUNCTIONS

Persons carrying out compliance tests to satisfy the requirements of the Radiological Council must be licensed for this purpose or be acting under the direction and immediate personal supervision\(^1\) of a licensed compliance tester. All test reports must be assessed and certified by a qualified expert.

Licensees must —

- have a working knowledge of the x-ray equipment;

- conduct the tests according to the protocols in the relevant workbook or use other protocols approved by the Radiological Council;

- ensure that the reported test results are accurate within the accepted error of the test equipment and that all data reported, including equipment and owner identification, is a true record pertaining to the equipment under test;

- submit the test report to a qualified expert for assessment within 1 month of the commencement date of the test.

Note: Corrective actions deemed necessary by a qualified expert are the responsibility of the registrant. However, the qualified expert and the compliance tester may wish to liaise with the registrant to ensure that non-complying items are corrected in a timely manner and, if necessary, re-tested before a compliance certificate is issued.

4.2 QUALIFICATIONS

A relevant technical qualification acceptable to Council, or equivalent experience in a medical radiation field.

4.3 ASSESSMENTS

The assessment syllabus for compliance testers is given in Appendix 8.

Applicants must pass —

- a multiple choice core examination in the fundamentals of radiation safety, and

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\(^1\) ‘Immediate personal supervision’ requires the licensee to be physically present and to directly observe persons working under their direction and supervision.
• a written examination which tests knowledge of compliance testing, including physical principles and methods, and

• a practical compliance testing examination supervised by an independent\(^1\) qualified expert licensed to conduct compliance tests.

Applicants who hold a licence for the service of diagnostic x-ray equipment may be exempted from the core examination.

A licence for compliance testing of dental x-ray equipment only is also available.

### 4.4 TRAINING

Applicants who require additional theoretical training prior to the written and practical examination may find it useful to attend relevant parts of a suitable Radiological Council approved course. Instruction might also be obtained from a qualified expert. Applicants should contact the Council for further advice on suitable courses.

To comply with Sections 25 and 36 of the Radiation Safety Act, practical experience in compliance testing can only be gained under the direction and immediate personal supervision of a licensed compliance tester i.e. the licensee must be present whenever the trainee initiates an exposure. A minimum of one machine of the type in question must be tested before considering eligibility for the practical examination.

When undertaking the practical compliance testing examination both a licensed compliance tester and the qualified expert must be present. This may be the same person if the qualified expert also holds a licence for compliance testing.

For the practical examination, the qualified expert must personally supervise the performance of the following number of complete tests applicable to the scope of the candidate’s proposed licence:

- Major Fluoroscopy/Radiography unit \(1\)
- Mammography unit \(1\)
- CT \(1\)
- Dental units \(2 \text{ (1 intraoral, 1 panoramic)}\)

For example, a person intending to provide dental compliance testing only must complete the two prescribed tests to the satisfaction of the supervising qualified expert.

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\(^1\) Independent:- having no familial, matrimonial, business etc relationships.
4.5 Licensing

Persons who satisfy the qualifications and assessment criteria are eligible to apply for a compliance testing licence.

Licences are subject to the conditions given in Appendix 7.

See also section 2 of this workbook for the requirements for a valid compliance test.
5. REQUIRED COMPLIANCE TESTS

Testing assesses the compliance of diagnostic equipment with the Radiation Safety (General) Regulations and with any additional requirements of the Radiological Council that may apply to the equipment under test. The complete test protocols, together with assessment criteria, are provided in the workbook applicable to each class of equipment.

Variations to the test protocols may be used provided they are first approved by the Radiological Council.

The following tests are required where appropriate (e.g. timer tests are not applicable to mAs units) —

5.1 MOBILE RADIOGRAPHIC EQUIPMENT

For both film and digital units:-

- Light beam collimator
  - accuracy of collimation
  - illuminance test

- Generator and x-ray tube
  - tube voltage accuracy
  - timer accuracy
  - radiation output and output linearity
  - reproducibility
  - half value layer
  - tube housing assembly leakage

See Workbook 1, sections 2 and 3.

For digital units only, add:-

- PSP calibration
  - dynamic range and dose measurement

See Workbook 7, sections 2, 3 and 4.

5.2 MAMMOGRAPHIC EQUIPMENT

For both film and digital units:-

- Light beam collimator
  - accuracy of collimation
  - illuminance test

- Generator and x-ray tube
  - tube voltage accuracy
  - timer accuracy
  - radiation output and output linearity
  - reproducibility
  - half value layer
  - tube housing assembly leakage
  - automatic exposure control
Patient dose information ➢ mean glandular dose

See Workbook 2, sections 2, 3 and 4.

For digital units only, add:-

PSP calibration ➢ dynamic range and dose measurement

See Workbook 8, sections 2, 3, 4 and 5.

5.3 MAJOR RADIOGRAPHIC EQUIPMENT

For both film and digital units:-

Light beam collimator ➢ accuracy of collimation
➢ illuminance test

Generator and x-ray tube ➢ tube voltage accuracy
➢ timer accuracy
➢ radiation output and output linearity
➢ reproducibility
➢ half value layer
➢ tube housing assembly leakage

Automatic exposure control ➢ standard density test
➢ kilovoltage and patient thickness tracking
➢ minimum response time

Patient dose information ➢ AP abdominal projection (general)
➢ AP lumbo-pelvic projection (chiropractic)

See Workbook 3, sections 2, 3, 4 and 5.

For digital units only, add:-

PSP calibration ➢ dynamic range and dose measurement

See Workbook 9, sections 2, 3, 4, 5 and 6.

5.4 FLUOROSCOPIC EQUIPMENT

For both film and digital units:-

Beam collimation ➢ collimation accuracy for fluoroscopy
➢ collimation accuracy for radiography

Generator and x-ray tube ➢ apparatus configuration
➢ tube voltage accuracy (fluoroscopic and radiographic)
➢ timer accuracy (radiographic)
➢ radiation output and output linearity (fluoroscopic and radiographic)
➢ reproducibility (radiographic)
➢ half value layer
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- tube housing assembly leakage
- standard density test
- kilovoltage and patient thickness tracking
- minimum response time

Automatic exposure control

- congruency of x-ray beam and displayed image
- image intensifier input dose rate
- image quality

Imaging system

- fluoroscopic timer
- maximum dose rate
- typical dose rates with patient equivalent phantom

Patient dose information

See Workbook 4, sections 2, 3, 4, 5 and 6.

For digital units only, add:-

- PSP calibration
  - dynamic range and dose measurement

See Workbook 10, sections 2, 3, 4, 5, 6 and 7.

5.5 DENTAL RADIOGRAPHIC EQUIPMENT

For both film and digital units:-

- Light beam collimator
  - accuracy of collimation
  - illuminance test

- X-ray beam size, beam alignment and focal spot to skin distance
  - method for intra-oral equipment
  - method for cephalometric equipment
  - method for panoramic tomographic equipment

- Control and x-ray tube
  - tube voltage accuracy
  - timer accuracy
  - radiation output and output linearity
  - reproducibility
  - half value layer
  - tube housing assembly leakage

- Patient dose information
  - typical intraoral bitewing dose

See Workbook 5, sections 2, 3, 4 and 5.

For digital units only, add:-

- PSP calibration
  - dynamic range and dose measurement

See Workbook 11, sections 2, 3, 4, 5 and 6.
## 5.6 COMPUTED TOMOGRAPHIC EQUIPMENT

| Generator, x-ray tube and scanner | ➢ tube voltage accuracy  
|                                  | ➢ radiation output and output linearity  
|                                  | ➢ half value layer  
|                                  | ➢ CT dose index  
|                                  | ➢ tube housing assembly leakage  
| Image quality                   | ➢ mean CT number/uniformity/noise  
|                                  | ➢ linearity of response  
|                                  | ➢ high contrast resolution  
|                                  | ➢ slice thickness  
|                                  | ➢ table indexing & reproducibility  
|                                  | ➢ alignment light and image slice congruence |

*See Workbook 6, sections 2 and 3.*
PUBLICATIONS

1. **Radiation Safety Act 1975 (and Amendments)**
   Schedule IX, Radiation Safety (General) Regulations 1983 (and Amendments)
   State Law Publisher

2. **Patient Dose Reduction in Diagnostic Radiology.**
   Report by the Royal College of Radiologists and the National Radiological Protection Board (UK)
   Documents of the NRPB, Vol 1 No 3 1990.

3. **Radiation Doses to Patients from Dental Radiography in New Zealand.**
   National Radiation Laboratory, Christchurch, New Zealand
   Williamson B D P, Report NRL 1990/6

4. **Variability of Medical Diagnostic X-ray Machine Parameters as determined from a National Survey.**
   National Radiation Laboratory, Christchurch, New Zealand
   Le Heron J, Report NRL 1989/1

5. **Assurance of Quality in the Diagnostic X-ray Department.**
   British Institute of Radiology 1988

6. **Quality Control in Diagnostic Imaging**
   Gray J E, Winkler N T, Stears J and Frank E D.
   University Park Press 1983

7. **Radiation Doses to Patients in Medical Diagnostic X-ray Examinations in New Zealand: a 1983-84 Survey.**
   National Radiation Laboratory, Christchurch, New Zealand
   Williamson B D P, Poletti J L, Cartwright P H and Le Heron J C
   Report NRL 1993/1

8. **Quality Assurance for Diagnostic Imaging Equipment.**
   National Council on Radiation Protection and Measurements
   NCRP Report No 99

9. **1990 Recommendations of the International Commission on Radiological Protection.**
   ICRP Publication 60, 1990

    National Health and Medical Research Council
APPENDIX 1

COMPLIANCE TESTING PROCESS FLOW CHART
Licensed compliance tester performs test. (1 month period to refer report to qualified expert (QE) begins)*

Tester refers test results, x-ray films, images, waveforms, printouts, etc. to a QE for assessment and certification

QE requests compliance tester to provide additional information and/or initiates repeat of inadequately performed tests.

QE satisfied that the test is accurate (within test equipment specifications) and performed according to approved protocols?

NO

Can the fault(s) be rectified?

YES

Equipment complies?

NO

YES

QE certifies (signs) the compliance test report and issues a Compliance Certificate and compliance label for the x-ray equipment within 1 month of receipt of the test report.

Original Compliance Certificate, label and copy of the test report provided to the client (registrant). QE promptly (within 2 months of the date the test commenced) provides a copy of the assessed and signed test report and a copy of the Compliance Certificate to the Radiological Council.

QE issues the Registrant with a Notice of Non-Compliance (period not to exceed 3 months).

NO

YES

QE promptly forwards a copy of the Non-Compliance Notice to the Radiological Council (within 2 months of the date the test commenced. If the fault is such that the QE believes there is significant radiation risk to patients, users or the public, the QE shall immediately advise the Council that a direction to stop use should be issued.

The client (registrant) is responsible for acting on the Notice of Non-Compliance but the QE and tester should liaise with the client to ensure follow-up action is timely.

QE advises client to withdraw the x-ray equipment from use or to apply to the Radiological Council if there are possible grounds for Conditional Compliance or an Exemption. QE promptly informs the Radiological Council of the action taken.

* When a Notice of Non-Compliance is issued, the registrant (the owner) has until expiry of the Notice or 3 months from expiration of the previous Compliance Certificate (whichever occurs earlier) to effect the required repairs. The tester and QE therefore should liaise with the client to complete any necessary re-testing and assessment without delay. The use of non-complying equipment after the specified period may breach the Act.

Overview of the Processes in Compliance Testing
APPENDIX 2

REGISTRATION CONDITIONS
RADIATION SAFETY ACT

CONDITIONS, RESTRICTIONS AND LIMITATIONS (SECTION 36)

MEDICAL RADIOLOGY

1. This registration provides for the possession and use of the specified x-ray apparatus for the purpose of human diagnostic radiography and fluoroscopy, performed under the direction and supervision of a licensed medical practitioner. Self referred screening x-ray examinations are excluded unless the registration has been otherwise endorsed.

2. Subject to the restrictions and limitations that may be imposed by these conditions and provided that all x-ray examinations are performed on x-ray equipment designed for the purpose, the range of x-ray examinations permitted is unrestricted.

3. The registrant is directed to ensure that —

3.1 x-ray apparatus on the premises is not operated or used for human diagnostic radiography or fluoroscopy unless it has —

(a) a current Certificate of Compliance;
(b) a current Certificate of Conditional Compliance; or
(c) an exemption from compliance granted by the Council.

Any x-ray apparatus which has been tested but which requires service before a Certificate of Compliance can be issued, can continue to be used for a period of 3 months after expiry of the current certificate provided —

• a qualified expert is satisfied that the fault(s) do not pose a significant radiation risk to users and/or patients; and

• the qualified expert has issued a written order (Notice of Non-Compliance) to the registrant for correction of the fault(s).

3.2 fluoroscopic x-ray apparatus is used only by —

3.2.1 licensed radiologists;

3.2.2 medical practitioners training for qualifications in diagnostic radiology and working under the direction and general supervision of a licensed radiologist;

3.2.3 medical practitioners with specialist qualifications (other than a licensed radiologist) who hold a licence or an exemption from licence and who are using the apparatus -

(i) for a purpose relevant to those qualifications; and

(ii) in the presence of a radiographer who has responsibility for positioning and manipulating the apparatus, minimising patient and personnel radiation exposure;

3.2.4 medical practitioners training for specialist qualifications (other than a person to whom paragraph 3.2.2 applies) who -

(i) has attended an approved course of training and has passed an examination in
radiation safety;

(ii) is using the apparatus under the direction and general supervision of a medical practitioner who holds a licence or an exemption from licence; and

(iii) is using the apparatus for a purpose relevant to those qualifications in the presence of a radiographer who has responsibility for positioning and manipulating the apparatus, minimising patient and personnel radiation exposure;

3.2.5 radiographers using the apparatus as part of a procedure —

(i) while working under the direction and personal supervision of a licensed radiologist who is responsible for that procedure; or

(ii) where -

(a) conventional radiography is inadequate; and

(b) the licensed radiologist responsible for that procedure has given permission for such use.

3.3 radiographic x-ray apparatus is used only by —

3.3.1 radiographers,

3.3.2 persons enrolled in a recognised course of training in diagnostic radiography and working under —

(i) the direction and general supervision of a licensed radiologist; and

(ii) the personal supervision of a radiographer.

3.3.3 approved x-ray operators for chest and extremity radiography using low powered x-ray equipment only. X-ray operators are restricted to providing services outside the metropolitan area only.

3.4 except where the registration is otherwise endorsed, radiographic x-ray examinations are performed only on receipt of a written request authorised by -

3.4.1 a medical practitioner; or

3.4.2 a dentist, for whom such authorisation is restricted to —

(i) examinations of the teeth and jaws and other maxillary or mandibular structures (by means of either plain radiography or CT, with or without the use of contrast media); and

(ii) a plain chest x-ray examination, where the dentist believes that a patient may have aspirated a foreign body during, or as a result of, dental treatment.

*Note:* In exceptional circumstances, a verbal request from the practitioner directly to the person expected to perform the examination will suffice but full authorisation must be provided at the earliest opportunity.

3.5 unless exempted by the regulations or by Council, each person on the premises who is occupationally exposed to radiation, is individually issued with and wears an approved radiation monitoring device to record their cumulative radiation dose and that —
3.5.1 each device is used only by the person to whom it was issued;
3.5.2 the device(s) are returned for assessment at the intervals directed by the Council;
3.5.3 continuing records are maintained of all personal monitoring, and
3.5.4 the results of this monitoring are made available forthwith to the individuals concerned.

Note: An exemption from monitoring applies to persons who must necessarily be present during the use of mobile fluoroscopic x-ray apparatus, provided —

- they stand no closer than 3 metres to the patient and x-ray tube during exposures, and
- they wear a 0.25 mm (minimum) lead equivalent apron or are standing behind an equivalent approved barrier.

3.6 these conditions are brought to the attention of all personnel authorised to use the x-ray apparatus and are a copy displayed or kept available within the x-ray room, department or practice to which such personnel have access. 

NOTES:

a “current” means a certificate that has not exceeded the date of expiry.

Note: The expiry date for certificates that bear a “Date of issue” alone will be determined by adding the test period prescribed below to that date —

- 12 months for mammographic and C-arm or U-arm fluoroscopic apparatus (fixed or mobile);
- 24 months for other fluoroscopic, general radiographic and CT apparatus; and
- 36 months for dental x-ray apparatus

b “compliance” means compliance with the Radiation Safety (General) Regulations 1983, with any subsequent amendments to those regulations, and with any additional requirements of the Council applicable to that class of x-ray apparatus

c “conditional compliance” may be granted to non-complying equipment if the equipment has an existing registration and was manufactured before the particular regulations or standards applying to the non-compliance came into effect, and if the non-compliance, in the opinion of Council —

- cannot reasonably be rectified; and
- does not cause an unacceptable increase in radiation dose.

d “radiologist” means a medical practitioner with qualifications in diagnostic radiology that are recognised by the Royal Australian and New Zealand College of Radiologists

e “radiographer” means a person with qualifications in diagnostic radiography who is eligible for ordinary membership of the Australian Institute of Radiography

f “recognised” means recognised by the Australian Institute of Radiography

g “general supervision” means the exercise of control over radiation safety without the person exercising such control necessarily being present at the registered premises or field site

h “personal supervision” means the exercise of control over radiation safety by the person exercising such control being present on the registered premises or field site

i “x-ray operator” means a person who has attended an approved course of training and has passed an examination in radiation safety and radiographic techniques for plain radiography of the chest and extremities

j “low powered irradiating apparatus” means irradiating apparatus registered for the purpose of medical radiography (other than mammography or dental radiography) that has a maximum rating of 100 kV(peak) at 100 mA or 100 kV(peak) and 1 microfarad

k “authorised” means by personal signature or by any other method approved by the Council.

l Regulations 18 and 19 of the Radiation Safety (General) Regulations 1983 require the registrant and Radiation Safety Officer to
ensure that training and instruction is given to all persons working with x-ray equipment on the premises.

"approved" means approved in writing by the Council

"plain radiography" means an x-ray examination during the course of which the x-ray tube and film remain stationary and no contrast medium is introduced into the patient

CONDITION NO: 2
CERTIFICATE NO: RX
EXPIRES ON:
APPENDIX 3

COMPLIANCE CERTIFICATES
RADIATION SAFETY ACT (WA)

CERTIFICATE of COMPLIANCE

for the following x-ray equipment:-

Manufacturer

Model

Serial Number

Equipment Use

Registrant

Premises Location

Location on Premises

Tested by

Licence No.

Test Date

I certify that, at the date of test, the equipment satisfied the compliance assessment criteria stipulated by the Radiological Council in the Workbook applicable to this class of equipment

Qualified Expert\(^1\)

Signature

Certificate Number

Expiry Date

\(^1\) A ‘qualified expert’ means an expert whose qualifications are approved by the Radiological Council.
**RADIATION SAFETY ACT (WA)**

**CERTIFICATE of CONDITIONAL\(^1\) COMPLIANCE**

*for the following x-ray equipment:*

<table>
<thead>
<tr>
<th>Manufacturer</th>
<th>Model</th>
<th>Serial Number</th>
<th>Equipment Use</th>
<th>Registrant</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Tested by **Test Date**

I certify that, at the date of test, the equipment satisfied the compliance assessment criteria stipulated by the Radiological Council in the Workbook applicable to this class of equipment, except in respect of the following item(s):-

<table>
<thead>
<tr>
<th>Qualified Expert(^2)</th>
<th>Signature</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Certificate Number **_EXPIRY DATE**\(^3\)

---

1. This certificate is not transferable and is valid only for the current compliance period. Compliance has been issued on the grounds that faults identified are unlikely to impact on patient or occupational doses and inherent design features of the unit make corrective modification infeasible.

2. A 'qualified expert' means an expert whose qualifications are approved by the Radiological Council.

3. The Radiological Council has stipulated that Conditional Compliance applies for a maximum period of 6 years. Routine compliance testing at the specified frequency for this class of equipment must continue during this period. Conditional Compliance certificates will not be issued for this machine after .................
APPENDIX 4

SAMPLE COMPLIANCE LABELS
Compliance label - yellow

Conditional compliance label - magenta

Shown larger than actual size
APPENDIX 5

NOTICES OF NON-COMPLIANCE AND CEASE USE
# RADIATION SAFETY ACT 1975

## NOTICE OF NON-COMPLIANCE

for the following x-ray equipment:-

<table>
<thead>
<tr>
<th>Manufacturer</th>
<th>Model</th>
</tr>
</thead>
<tbody>
<tr>
<td>Serial No.</td>
<td>Location</td>
</tr>
<tr>
<td>Equipment Use</td>
<td>Test Date</td>
</tr>
<tr>
<td>Tested by</td>
<td>Lic No.</td>
</tr>
<tr>
<td>Registrant</td>
<td></td>
</tr>
</tbody>
</table>

The following corrective actions must be undertaken by\(^1\):-

1. 
2. 

Signature __________________________  Date __________________________

Qualified Expert\(^2\)

Copies of this document must be provided forthwith to the Registrant and the Radiological Council\(^3\)

---

\(^1\) If the test date is prior to the current compliance certificate expiry date, add 3 calendar months to the test date. If the test date is on or later than the expiry date, add 3 calendar months to the expiry date.

\(^2\) Radiation Safety (General) Regulations 1983, Sections (5) and (6) of regulation 23. A ‘qualified expert’ means an expert whose qualifications are approved by the Radiological Council.

\(^3\) Locked Bag 2006 P O, NEDLANDS WA 6009. Telephone: (08) 9346 2260 Facsimile: (08) 9381 1423
RADIATION SAFETY ACT 1975

NOTICE UNDER SECTION 46

to CEASE USE of the following x-ray equipment:-

<table>
<thead>
<tr>
<th>Manufacturer</th>
<th>Model</th>
</tr>
</thead>
<tbody>
<tr>
<td>Serial No.</td>
<td>Location</td>
</tr>
<tr>
<td>Equipment Use</td>
<td>Test Date</td>
</tr>
<tr>
<td>Tested by</td>
<td>Lic No.</td>
</tr>
</tbody>
</table>

Registrant

TO:………………………..

I, ……………………………….., being a duly appointed ‘authorised officer’ for the purposes of the Radiation Safety Act 1975 (the Act), and being of the opinion that the irradiating equipment specified above may affect the safety or health of any person if the equipment is permitted to continue to be used without a current certificate of compliance (as required as part of the conditions of registration), hereby direct you -

(a) to stop using the equipment immediately, and
(b) to not resume using the equipment until –
   (i) the following non-compliant items have been corrected:-

(ii) the equipment has been retested for compliance
(iii) a certificate of compliance has been issued in respect of the equipment
(iv) notice of completion of the requirements set out at (i), (ii) and (iii) has been given to the Radiological Council
(v) notice has been received that this direction has been withdrawn.

Signature __________________________ Date ____________

Authorised Officer

Note that in accordance with provisions of Section 12 of the Act, you have a right to apply to the State Administrative Tribunal (the SAT) for a review of this decision within 28 days from the day on which the decision was made. The SAT can be contacted –

Online: Forms can be completed electronically using the SAT Wizard
By telephone: (08) 9219 3111 or 1300 306 017
In person: Level 4, 12 St Georges Terrace, Perth, WA
By post: GPO Box U1991, Perth, WA 6845
APPENDIX 6

QUALIFIED EXPERT SYLLABUS
WRITTEN ASSESSMENT

1. Interactions between x-rays and matter

   1.1 Nature of x-radiation.

   1.2 Interaction processes:-
       - photoelectric effect
       - characteristic radiation
       - Compton scattering
       - bremsstrahlung
       - x-ray spectrum

   1.3 Attenuation:-
       - monoenergetic attenuation
       - linear attenuation coefficient
       - half-value layer
       - factors affecting attenuation

   1.4 Scattered radiation:-
       - effect of kV, field size, thickness

2. Production of x-rays

   2.1 X-ray spectrum:-
       - general radiation
       - characteristic radiation

   2.2 Effect of variation of:-
       - kV
       - mA
       - filtration
       - voltage waveform

   2.3 X-ray tubes:-
       - principal types and construction
       - line focus principle
       - heel effect
       - causes of failure
       - HT cables
       - tube ratings
       - tube housing leakage

   2.4 Types of generators:-
       - rectification
       - 3-phase, 6- and 12 pulse
       - medium frequency
       - capacitor discharge
       - battery powered

   2.5 Exposure timers.

   2.6 Automatic exposure control.

3. Filters, collimators, grids

   3.1 Filtration:-
       - inherent
       - added
       - K-edge (erbium, hafnium)
3.2 Types of collimators:
- radiography
- fluoroscopy

3.3 Scatter reduction techniques:
- collimation
- compression
- grids (grid ratio)
- air gaps

4. Radiographic films, screens and processing

4.1 Film:
- structure of x-ray film
- latent image formed by light or x-rays
- photographic density
- characteristic curve and film contrast, latitude
- speed
- spectral sensitivity

4.2 Luminescent screens:
- general principles
- intensification factor
- speed
- types of phosphor
- emission spectrum
- resolution
- response to kV

4.3 Film processing:
- darkroom safelights
- manual processing
- automatic processors

5. Radiographic image

5.1 Contrast:
- subject contrast
- film contrast
- fog

5.2 Image quality:
- quantum mottle - noise
- sharpness
- limiting resolution
- line spread function
- modulation transfer function
- Weiner power spectrum

5.3 Geometrical considerations:
- effect of magnification
- effect of focal spot size
- distortion

6. Image intensification and TV chain

6.1 Principles of system.
6.2 Design and operation.
6.3 Performance characteristics:
- contrast, resolution, Gx
- distortion
- veiling glare
- MTF

6.4 Automatic brightness control.

7. Digital radiographic systems

7.1 Modalities:
- computed radiography, principle of photostimulable plate
- charge coupled devices, principle of CCD chip, CMOS and uses in fluoroscopy, mammography and dental radiography
- indirect flat panel, principle of thin film transistor array
- direct flat panel, principle

7.2 Image production and processing:
- general principles of laser read out in computed radiography and display in other modalities
- image processing techniques available

7.3 System performance:
- correlation of ‘exposure index’ and incident plate dose for various manufacturers, e.g. Agfa ‘lgM’, Fuji ‘S’, etc
- relationship between incident dose and dynamic range of image
- calibration of dynamic range
- other aspects of system performance including:– erasure cycle efficiency, sensitivity index consistency/sensitivity, uniformity, blurring, limiting resolution, threshold contrast detail detectability

8. Types of x-ray machines

8.1 Familiarity with:
- dental, mobile radiographic, fluoroscopic, CD units
- chest and general x-ray units
- fluoroscopic tables, C-arms and special units
- mammography units
- CT scanners

9. Instrumentation

9.1 Ionisation chambers:
- principles of ionisation chambers
- types of ionisation chambers
- calibration of chambers
9.2 Non-invasive testers e.g. NERO:-

- principles of the test device
- familiarity with and ability to use test device and associated software for kV, mA, timer, reproducibility checks, HVL measurement etc

9.3 Image quality test objects e.g. Leeds test object:-

- principles of the device
- familiarity with and ability to use the test object correctly according to the manufacturer's instructions regarding test factors

10. Compliance tests

10.1 Workbook tests:-

- knowledge of the physical basis of the tests
- assessment of alternative methods of performance of the tests
- ability to perform the tests

11. Physical concepts

11.1 Radiation units:-

- exposure (air kerma)
- absorbed dose

12. Radiation protection concepts

12.1 ICRP 60:-

- principles of justification
- optimisation
- dose limits
- dose constraints
- occupational exposure
- medical exposure
- population exposure
- radiation weighting factor
- equivalent dose
- tissue weighting factor
- effective dose

12.2 Protection of the patient:-

- summary of principles given in ICRP 62 (in Summary)

13. Practical radiation protection

13.1 Inverse square law and distance.

- control area barriers
- personal aprons, lead effectiveness etc

13.2 Shielding:-

- basic understanding of personal radiation monitoring dosimetry
14. Regulations

14.1 ICRP 57:-
   - familiarity with section on requirements for diagnostic imaging equipment

14.2 State Regulations:-
   - familiarity with sections dealing with diagnostic imaging equipment

PRACTICAL ASSESSMENT

- A pass in the written paper is normally required before candidates without compliance testing experience may attempt the practical assessment.

- The assessment involves the candidate performing a compliance test under the direct supervision of a qualified expert who holds a licence for compliance testing.

- The documented test and the qualified expert’s report on the candidate’s performance are then submitted to the Council for consideration.

- If, by mutual agreement with a qualified expert, a candidate has taken and passed a practical assessment before applying to sit the written examination, no additional practical assessment will be required.

- Candidates who require tuition before the practical assessment must be under the immediate personal supervision of a person who holds a licence for compliance testing. This person does not have to be a qualified expert.

References


Radiological Protection of the Worker in Medicine and Dentistry.  
ICRP Publication 57  
Pergamon Press, 1990

Radiological Protection in Biomedical Research (includes Summary of the Current ICRP Principles for Protection of the Patient in Diagnostic Radiology)  
ICRP Publication 62  
Pergamon Press, 1993

Diagnostic X-Ray Equipment Compliance Testing.  
Workbooks 1 to 6.  
Radiological Council of Western Australia 2005

Radiation Safety (General) Regulations 1983 (and amendments)  
State Law Publisher

Acceptance Testing and Quality Control of Photostimulable Storage Phosphor Imaging Systems  
Report of Task Group #10, American Association of Physicists in Medicine, 1998

Protocol for the QA of Computed Radiography Systems: Routine QA Tests  
Kings Centre for the Assessment of Radiological Equipment (K CARE), London, 2004.

Specification, acceptance testing and quality control of diagnostic X-ray imaging equipment.  
Seibert J A, Barnes G T and R G Gould  
AAPM Medical Physics Monograph No 20, 1994, American Institute of Physics.
APPENDIX 7

LICENCE CONDITIONS FOR COMPLIANCE TESTERS
RADIATION SAFETY ACT

CONDITIONS, LIMITATIONS AND RESTRICTIONS (SECTION 36)

COMPLIANCE TESTING OF DIAGNOSTIC X-RAY EQUIPMENT

1. This licence permits the holder, and persons acting under the direction and immediate personal supervision of the licensee, to operate diagnostic x-ray equipment registered for human dental, medical and chiropractic diagnosis for the purpose of compliance testing.

2. The licensee is directed —
   2.1 to ensure that no person is exposed to the direct x-ray beam for any purpose during test procedures;
   2.2 to use protective barriers and/or lead equivalent drapes and aprons to minimise the radiation dose to themselves and any persons in the vicinity (except in the case of intra-oral and panoramic tomographic x-ray equipment when distance from the x-ray source and appropriate beam orientation will provide adequate protection); and
   2.3 to wear a radiation monitoring film or other approved personal monitoring device, issued to the licensee for his or her exclusive use, whenever x-ray equipment is used.

3. For compliance testing of x-ray equipment which has been imposed as a statutory requirement by the Radiological Council, the licensee is directed to —
   3.1 test the equipment according to the protocols in the approved workbook relevant to the class of equipment under test, or by following other approved test procedures, and by using appropriate instruments that have been calibrated within the 2 years prior to the commencement of the test by an organisation recognised by the Radiological Council;
   3.2 ensure that the reported test results are accurate within the accepted error of the test equipment and that all data reported, including equipment and owner identification, is a true record pertaining to the equipment under test;
   3.3 ensure that any faults found, or found and corrected, during testing are detailed in the test report; and
   3.4 within 1 month of the date the compliance test commenced provide a copy of all the test results including relevant x-ray images, waveforms and computer printouts to a qualified expert for assessment.

a ‘Immediate personal supervision’ means maintaining direct visual supervision of the person concerned.

b ‘Compliance testing’ means testing x-ray equipment for compliance with the regulations under the Act and with other standards that may have been adopted by the Radiological Council for that class of equipment.

c ‘Qualified expert’ means an expert whose qualifications are approved by the Radiological Council.
NOTES:

For compliance testing imposed as a statutory requirement —

- The qualified expert must sign a compliance certificate and provide it to the registrant (the ‘owner’ of the equipment) for each item of x-ray equipment tested and found to be in compliance.

- Equipment certified to be in compliance with the regulations and other Council requirements must bear an approved label showing the test date, the certificate number and the name of the tester and qualified expert.

Workbooks, compliance certificates and equipment labels are available from the Radiological Council.

Radiological Council
18 Verdun Street
NEDLANDS WA 6009

Locked Bag 2006 P O NEDLANDS WA 6009
Telephone (08) 9346 2260 Fax (08) 9381 1423
APPENDIX 8

COMPLIANCE TESTER SYLLABUS
WRITTEN ASSESSMENT

The written assessment comprises two sections:

- **Core Paper**
  *Closed book, one hour paper covering general radiation safety*

- **Main Paper**
  *Open book, two hour paper covering compliance testing*

The main paper is based on material contained in the Radiological Council’s workbooks for the following equipment categories —

1. Mobile Radiographic
2. Mammographic
3. Major Radiographic
4. Fluoroscopic
5. Dental Radiographic
6. Computed Tomography

Two main papers are available depending on the applicant’s needs. One deals with dental x-ray equipment (*with reference to workbook 5 only*) and the other with all medical and dental categories (*with reference to workbooks 1-6,*).

1. **Core Paper - Closed book**

   1.1 Legislation:-
      - Radiation Safety Act 1975
      - Radiation Safety (General) Regulations 1983
   
   1.2 Dose limits:-
      - workers
      - members of the public

   1.3 Radiation types & properties.
   1.4 Background radiation.
   1.5 Quantities and units of measurement.
   1.6 Biological effects.
   1.7 Radiation risk.
   1.8 Basic radiation safety calculations.
   1.9 Inverse square law.
   1.10 Pro rata dose calculations.
   1.11 Personal radiation monitoring.
1.12 Principles of protection:-
   - time
   - distance
   - shielding

2. **Main Paper.** *Open book.* The syllabus includes —

2.1 Half value layer.

2.2 Leakage:-
   - tube housing
   - light beam diaphragm

2.3 Light beam diaphragm:-
   - alignment
   - congruency
   - illuminance

2.4 CD leakage requirements.

2.5 Tube voltage accuracy.

2.6 Exposure time accuracy.

2.7 Radiation dose measurement.

2.8 Radiation output linearity with tube current.

2.9 Reproducibility of outputs:-
   - coefficient of variation

2.10 AEC tests:-
   - standard
   - subject thickness tracking
   - tube voltage tracking
   - minimum response time

2.11 Fluoroscopic doserates:-
   - maximum
   - typical
   - high doserate restrictions

2.12 Mammography:-
   - dose measurements
   - mean glandular dose calculation
   - QA measurements (focal spot size, sensitometry)

2.13 CT:-
   - CT dose index
   - noise
   - mean CT number
   - uniformity
   - resolution
   - slice thickness
PRACTICAL ASSESSMENT

- A pass in the written paper is normally required before candidates without compliance testing experience may attempt the practical assessment.

- The assessment involves the candidate performing a compliance test under the direct supervision of an independent\(^1\) qualified expert who holds a licence for compliance testing.

- The documented test and the qualified expert’s report on the candidate’s performance are then submitted to the Council for consideration.

- If, by mutual agreement with an independent qualified expert, a candidate has taken and passed a practical assessment before applying to sit the written examination, no additional practical assessment will be required.

Candidates who require practical tuition before the assessment must work under the immediate personal supervision of a person who holds a licence for compliance testing. This person does not have to be a qualified expert.

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\(^{1}\) Independent: having no familial, matrimonial, business etc relationships
APPENDIX 9

DEFINITIONS
## DEFINITIONS

The following terms and definitions have been extracted from the Radiation Safety Act, the regulations and relevant registration conditions. They may assist in interpreting the requirements of the compliance testing program.

<table>
<thead>
<tr>
<th>Term or Definition</th>
<th>Meaning</th>
</tr>
</thead>
<tbody>
<tr>
<td>‘approved’</td>
<td>approved in writing by the Radiological Council</td>
</tr>
<tr>
<td>‘authorized officer’</td>
<td>a person who is appointed in writing by the Executive Director (of Public Health), either generally or in a particular case, to perform duties under the Radiation Safety Act 1975 and is thereby authorised to exercise the powers conferred by or under the Act, and also includes any member of the Council</td>
</tr>
<tr>
<td>‘compliance’</td>
<td>means compliance with the Radiation Safety (General) Regulations, with any subsequent amendments to those regulations, and with any additional requirements of the Council applicable to that class of x-ray apparatus</td>
</tr>
<tr>
<td>‘compliance tester’</td>
<td>a person licensed under the State’s Radiation Safety Act for that purpose</td>
</tr>
<tr>
<td>‘condition’</td>
<td>conditions, restrictions and limitations imposed under Section 36 of the Radiation Safety Act</td>
</tr>
<tr>
<td>‘conditional compliance’</td>
<td>non-transferable compliance which may be granted to non-complying equipment if the equipment has an existing registration and if the non-compliance, as assessed by Council officers cannot reasonably be rectified; and does not cause an unacceptable increase in radiation dose.</td>
</tr>
<tr>
<td>‘Council’</td>
<td>the Radiological Council established pursuant to Section 13 of the Radiation Safety Act</td>
</tr>
<tr>
<td>‘current’</td>
<td>in relation to a compliance certificate means that the certificate was issued within the past 12, 24 or 36 months, depending on the test frequency established by the Radiological Council for that class of x-ray equipment</td>
</tr>
<tr>
<td>‘exemption’</td>
<td>exemption referred to in Section 6 of the Radiation Safety Act 1975</td>
</tr>
<tr>
<td>‘fluoroscopy’</td>
<td>the use of a continuous or pulsed x-ray beam to produce a dynamic real time image, the duration of which is not predetermined before the exposure is initiated;</td>
</tr>
<tr>
<td>‘general supervision’</td>
<td>the exercise of control over radiation safety without the person exercising such control necessarily being present at the registered premises or field site</td>
</tr>
<tr>
<td>Term or Definition</td>
<td>Meaning</td>
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<td>---------------------------------</td>
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</tr>
<tr>
<td>‘image receptor’</td>
<td>x-ray film, fluorescent screen, image intensifier input phosphor or electronic device in or from which an image is created following exposure to x-rays</td>
</tr>
<tr>
<td>‘immediate personal supervision’</td>
<td>the exercise of control over radiation safety by the person exercising such control being in the company of and directly observing the person under supervision.</td>
</tr>
<tr>
<td>‘irradiating apparatus’</td>
<td>any apparatus capable of producing ionising radiation of any prescribed type, or capable of accelerating atomic particles under any prescribed conditions</td>
</tr>
<tr>
<td>‘licence’</td>
<td>licence granted under the Act</td>
</tr>
<tr>
<td>‘licensed’</td>
<td>in relation to a person, means that the person is the holder of a relevant licence under the Act</td>
</tr>
<tr>
<td>‘licensee’</td>
<td>holder of a licence</td>
</tr>
<tr>
<td>‘owner’</td>
<td>used in relation to any substance, apparatus, product, article or premises, means the person to whom it belongs or the hirer, lessee, borrower, bailee, or mortgagee in possession, thereof, and includes any attorney, agent, manager, foreman, supervisor or other person in charge or having control of management thereof</td>
</tr>
<tr>
<td>‘qualified expert’</td>
<td>expert whose qualifications are approved</td>
</tr>
<tr>
<td>‘registered’</td>
<td>registered under the Act</td>
</tr>
<tr>
<td>‘registrant’</td>
<td>person in whose name premises are registered</td>
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</tbody>
</table>