Performance Standards and Test Protocols for Radiological Equipment

Number 2 — Dose Rate Instrumentation

Publication No. 2B/10
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Abstract

This document is the second in a set of five performance standards. The purpose of the standards is to ensure best current capability across a broad range of radiological equipment and provide a national benchmark against which any radiological equipment can be assessed. This assessment will help to improve the quality and consistency of radiological equipment used by emergency services in the event of CBRN incidents. The primary purpose of this equipment is to alert the user to the presence of radiation.

The first element of this document establishes appropriate and targeted technical performance criteria against which radiological equipment can be assessed. These criteria are identified as ‘essential’ requirements. A second element of this document will aim to stimulate the development of radiological equipment beyond current best measurement capability, with particular focus on end user requirements. These criteria are identified as ‘desirable’ requirements.

The test methodologies to be used are specified. If at any stage the equipment doesn’t meet the requirement stipulated, then testing may be terminated and the equipment would be defined as not satisfying the required standard.

When submitting radiological equipment for assessment, the manufacturer is required to provide at least three fully operational production instruments for the duration of the tests. A user manual for the instruments shall also be provided. It should be noted that some tests might cause significant damage to the instrument and so agreement from the manufacturer should be sought before any destructive tests are performed. Where possible, destructive tests should only be performed on a single instrument. If any tests are excluded then these shall be stated.

The five standards in the series each relate to one of the following categories: dose rate instrumentation, contamination-monitoring instrumentation, electronic personal dose meters, portable spectrometry/identification systems and detection/alert devices. Each standard addresses the following broad areas: radiological, environmental, electrical and ergonomic aspects of performance.
Acknowledgements

The standards were written and produced by the Radiation Metrology Group of the Health Protection Agency and reviewed by representatives of the Defence Science and Technology Laboratory (Dstl), the National Physical Laboratory (NPL) and the Atomic Weapons Establishment (AWE).
1 Introduction

To assess an instrument’s suitability for its use by emergency services in the event of CBRN incidents, a number of tests are required. The tests detailed in this report are designed to not only assess the instrument’s radiological performance under laboratory conditions, but also test the environmental, electrical and mechanical aspects, which may affect the instrument’s performance out in the field.

Instruments that are designed to measure photon or beta ($\beta$) dose rates can utilise a variety of different types of detector, the most common being the Geiger-Muller (GM) tube, the ionisation chamber and the scintillation detector.

1.1 Dose Rate Instrumentation

The requirement to make a measurement of the radiation dose rate is present in most scenarios where there is a penetrating gamma emitting nuclide or a high-energy beta. Typical and widely used nuclides for which this is likely to be applicable are: $^{241}$Am, $^{137}$Cs, $^{60}$Co, $^{192}$Ir, $^{226}$Ra, $^{75}$Se and possibly $^{90}$Sr, as this nuclide is likely to be present with its daughter nuclide of $^{90}$Y.
2 Reference Documents

This document has been compiled with reference to the following documents:

IEC 60846:2004 Radiation protection instrumentation. Ambient and/or directional dose equivalent (rate) meters and/or monitors for beta, X and gamma radiation.

IEC 60529:1992 Degrees of protection provided by enclosures (IP code).


ISO 4037-2:1997 X and gamma reference radiation for calibrating dose meters and dose rate meters and for determining their response as a function of energy. Part 2: Dosimetry for radiation protection over the energy ranges 8 to 1.3 MeV and 4 to 9 MeV.

ISO 4037-3:1999 X and gamma reference radiation for calibrating dose meters and dose rate meters and for determining their response as a function of energy. Part 3: Calibration of area and personal dose meters and the measurement of their response as a function of energy and angle of incidence.


RPD-OP-004-2006 Suitability of Radiation Monitoring Equipment — Comparison of Type-test Data. (RESTRICTED — Commercial).
3 Terminology

3.1 Special Word Usage
The following word usage applies:

- The word “shall” signifies a mandatory requirement.
- The word “should” signifies a recommended specification or method.
- The word “may” signifies an acceptable method or an example of good practice.

3.2 Definitions

3.2.1 SI Units
The units of the ‘International System of Units’. Multiples and sub-multiples of the SI units will be used in accordance with the SI.

3.2.2 Sievert
The SI unit of dose equivalent is the joule per kilogram (J kg⁻¹), which has been named the Sievert (Sv) by the International Commission on Radiological Protection (ICRP).

\[ 1 \text{ Sv} = 1 \text{ J kg}^{-1} \]

3.2.3 Ambient dose equivalent \( H^*(10) \)
Dose equivalent at a point in a radiation field that would be produced by the corresponding aligned and expanded field; in the ICRU sphere at a depth of 10 mm and on the radius opposing the direction of the aligned field. The ICRU sphere (ICRU report 33, 1980) is a 30 cm diameter, tissue equivalent sphere with a density of 1 g cm⁻³.

3.2.4 Ambient dose equivalent rate \( \dot{H}^*(10) \)
Ratio of \( dH^*(10) \) by \( dt \), where \( dH^*(10) \) is the increment of ambient dose equivalent in the time interval \( dt \).

\[ H^*(10) = \frac{dH^*(10)}{dt} \]

The SI unit of ambient dose equivalent rate is the Sievert per second (Sv s⁻¹). Units of ambient dose equivalent rate are any quotient of the Sievert, or its multiples or sub-multiples, by a suitable unit of time (e.g. \( \mu \text{Sv h}^{-1} \)).
3.2.5 **Directional dose equivalent** $\dot{H}(0.07; \Omega)$

Dose equivalent at a point in a radiation field that would be produced by the corresponding expanded field, in the ICRU sphere at a depth of 0.07 mm, on the radius in a specified direction $\Omega$.

The ICRU sphere (ICRU report 33, 1980) is a 30 cm diameter, tissue equivalent sphere with a density of 1 g cm$^{-3}$.

3.3 **Operating Modes**

3.3.1 **User mode (Routine)**

The default operating mode whilst the instrument is being operated by non-expert users. Any parameters that may affect the operation of the instrument shall be protected via password or other appropriate security measures. The ability to view these parameter settings is desirable, but they shall be protected to prevent any changes. This mode may also be referred to as 'simple' mode.

3.3.2 **Supervisor mode (Restricted)**

An advanced operating mode that can only be accessed by an expert user, via password or other appropriate security measures, to edit parameters that will affect the operation of the instrument, i.e. calibration parameters, alarm thresholds, etc. This mode may also be referred to as 'advanced' or 'expert' mode.

3.4 **Test Nomenclature**

3.4.1 **Acceptance test (Pre-use test / Test before first use)**

The acceptance test shall demonstrate that the instrument conforms to type-test data. The acceptance test checks for any potential faults and provides a reference of performance for comparison with subsequent routine tests for the lifetime of the instrument. Further information on the tests required can be found in current UK guidance, such as the NPL Measurement Good Practice Guide 14 (GPG14).

3.4.2 **Routine test (Periodic test)**

This test confirms that the performance of the instrument has not deteriorated since the acceptance test. It is more than a simple check. Further information on the tests required can be found in current UK guidance, such as the NPL Measurement Good Practice Guide 14 (GPG14).

It is recommended that the performances of the instrument's electrical and mechanical systems are also inspected during the routine test. For example, batteries, cables, connectors and controls shall be inspected and repaired or replaced where necessary. Depending on the severity
of the repair, it may be necessary to repeat the acceptance tests if, for example, the detector has been repaired or replaced.

3.4.3 Type-test
This test is performed on at least one or more standard production instruments picked at random. Ideally, all non-destructive tests should be performed on at least three standard production instruments. Destructive tests, however, may be performed on just a single instrument. The type-test investigates all aspects of the instrument’s design to show the extent of compliance with pre-defined specifications.
4 General Requirements

4.1 Quantities and Units
The instrument shall be scaled in appropriate units. The SI unit of dose equivalent is the joule per kilogram (J kg⁻¹) which has been named the Sievert (Sv) by the International Commission on Radiological Protection (ICRP). It is recommended that this unit is prefixed as required; for example, ‘μSv’ (micro Sievert: 10⁻⁶ Sv) or ‘mSv’ (millisievert: 10⁻³ Sv). For dose rate instrumentation, a suitable unit of time is also required and in most cases per hour (h⁻¹) is the most appropriate.

Instruments that are designed to measure photon ambient dose equivalent shall use the measuring quantity H*(10).

Instruments that are designed to measure photon or beta directional dose equivalent shall use the measuring quantity H’(0.07).

4.2 Measuring Ranges
The instrument shall be able to determine dose equivalent rate over a wide range of values as specified below.

4.2.1 Essential
The instrument shall be capable of determining dose equivalent rates within the range from 1 μSv h⁻¹ to approximately 10 mSv h⁻¹.

4.2.2 Desirable
The instrument should be capable of determining dose equivalent rates within the range from 1 μSv h⁻¹ to approximately 10 Sv h⁻¹.

4.3 Storage and Transport

4.3.1 Essential
The instrument shall be supplied in a bespoke foam-lined carry-case. The case shall be waterproof and impact resistant. The case shall have provision to store the batteries when removed from the instrument. The instrument should not be stored for long periods of time with the batteries installed.

The instrument shall be designed to operate within the requirements of this document following storage or transport during a period of at least 3 months in the supplied carry case at any temperature between -25 °C and +50 °C.
4.3.2 Desirable
It is desirable that the carry-case can protect the instrument during any possible air transport at low ambient pressure. Where this is not possible and the instrument could be damaged by air transport, this shall be clearly stated on the instrument and the case.
5 Standard Test Conditions

5.1 Reference Conditions

Reference conditions are given in the table below. Except where otherwise specified, the tests in this standard shall be carried out under the standard test conditions as indicated in the table below.

Table 1. Reference conditions and standard test conditions

<table>
<thead>
<tr>
<th>Influence quantity</th>
<th>Reference conditions</th>
<th>Standard test conditions</th>
</tr>
</thead>
<tbody>
<tr>
<td>Photon radiation for:</td>
<td></td>
<td></td>
</tr>
<tr>
<td>1. Ambient dose equivalent $H'(10)$</td>
<td>Gamma radiation from $^{137}\text{Cs}$ N-20 (ISO 4037-3)</td>
<td>Gamma radiation from $^{137}\text{Cs}$ or $^{60}\text{Co}$. Photon radiation from X-rays. N-20 (ISO 4037-3)</td>
</tr>
<tr>
<td>2. Directional dose equivalent $H'(0.07)$</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Beta radiation for:</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Directional dose equivalent $H'(0.07)$</td>
<td>$^{90}\text{Sr}/^{90}\text{Y}$</td>
<td>$^{90}\text{Sr}/^{90}\text{Y}$</td>
</tr>
<tr>
<td>Neutron radiation for:</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Neutron dose equivalent, $H'(10)$</td>
<td>Neutron radiation from $^{241}\text{Am(Be)}$ or $^{252}\text{Cf}$</td>
<td>Neutron radiation from $^{241}\text{Am(Be)}$ or $^{252}\text{Cf}$</td>
</tr>
<tr>
<td>Stabilisation time</td>
<td>15 minutes</td>
<td>Minimum of 15 minutes</td>
</tr>
<tr>
<td>Ambient temperature</td>
<td>20 °C</td>
<td>18 °C to 22 °C (1)</td>
</tr>
<tr>
<td>Relative humidity</td>
<td>65%</td>
<td>55% to 75% (1)</td>
</tr>
<tr>
<td>Atmospheric pressure</td>
<td>101.3 kPa</td>
<td>86.0 to 106.6 kPa (1)</td>
</tr>
<tr>
<td>Power supply voltage</td>
<td>Nominal power supply voltage</td>
<td>Nominal power supply voltage ±10%</td>
</tr>
<tr>
<td>Angle of incidence of radiation</td>
<td>Calibration direction supplied by manufacturer.</td>
<td>Direction given ±5°</td>
</tr>
<tr>
<td>Orientation of instrument</td>
<td>To be stated by the manufacturer.</td>
<td>Stated orientation ±5°</td>
</tr>
<tr>
<td>Instrument controls</td>
<td>Set up for normal operation.</td>
<td>Set up for normal operation.</td>
</tr>
<tr>
<td>Radiation background</td>
<td>1 μSv h$^{-1}$ or less if practical.</td>
<td>Known field less than 1 μSv h$^{-1}$</td>
</tr>
<tr>
<td>Contamination by radioactive elements</td>
<td>Negligible</td>
<td>Negligible</td>
</tr>
</tbody>
</table>

(1) The actual values of these quantities at the time of test shall be stated.
5.2 Reference Radiations
Unless specified otherwise in individual methods within this document, the nature, construction and conditions of use of radiation sources shall be accordance with the relevant parts of ISO 4037 and ISO 6980.

If the complete range of dose rates is not available from a single source of radiation, then additional sources normalised to the original source of radiation may be used.

Where an X-ray set is used to generate reference radiations, its output should be monitored by means of a monitor ionisation chamber that is permanently mounted in the radiation beam as specified in ISO 4037-2:1997. This is to ensure that the X-ray set is stable. Where a monitor chamber is not available, the X-ray set output shall be measured immediately before and after any irradiation, and any fluctuations found shall be accounted for.

The value of the quantity to be measured at the point of test shall be known with an uncertainty of less than 10%. All dose rates shall be traceable to national standards.

The radiation beam diameter shall be sufficient to irradiate the detector(s) of the instrument to be tested. The beam shall be uniform to within 10% over the useful beam area.

5.3 Low Dose Rates
For the measurement of low dose rates, it may be necessary to account for the contribution of natural background at the exact location of the test. Other contributing effects shall also be considered, such as electronic noise or inherent radiation from the construction materials of the instrument.

5.4 Variation of Influence Quantities
To properly assess each element of the required tests individually, it is important that only one influence quantity is changed at a time. Unless otherwise specified, all other influence quantities should be maintained at fixed values within the standard test conditions.

5.5 Instrument Orientation
The instrument under test shall be placed in an orientation with respect to the direction of the radiation field as indicated by the manufacturer and with the marked reference point accurately positioned at the point of test in the radiation field. In the absence of a marked reference point the geometric centre of the detector shall be used. The instrument orientation and reference point used must be clearly stated.
6 Radiological Performance Requirements

6.1 Linearity of Response

6.1.1 Essential
With the instrument set up as specified by the manufacturer’s instructions and under standard test conditions, the relative intrinsic error in the response to the reference gamma radiation (linearity) shall not exceed ±30% over the entire effective range of ambient dose equivalent rates.

6.1.2 Desirable
With the instrument set up as specified by the manufacturer’s instructions and under standard test conditions, the relative intrinsic error in the response to the reference gamma radiation (linearity) shall not exceed ±20% over the entire effective range of ambient dose equivalent rates.

6.1.3 Test radiation
The recommended reference radiation source for photon and β dose rate instruments is $^{137}$Cs. If $^{137}$Cs is not available then $^{60}$Co can be used, although corrections will need to be applied to relate the response to $^{137}$Cs.

For β dose rate instruments, the linearity measurements may also be performed with a set of β sources with various activities. Each source used must be of the same radionuclide and construction. In most instances $^{90}$Sr/$^{90}$Y will be a suitable radionuclide.

For the purpose of this test, the value of the true ambient dose equivalent rate produced by the reference gamma radiation at the point of test shall be known with an uncertainty of less than 10% at a confidence level of 95%.

Where higher dose equivalent rates are not available using a $^{137}$Cs source, then an X-ray generator may be used. Suitable corrections shall be made for any difference in the instrument’s response to the radiation generated to enable reference back to the reference radiation of $^{137}$Cs.

6.1.4 Method of test
For instruments with linear scales, measurements shall be performed on all scale ranges and at a minimum of three instrument indications in each range. It is recommended these indications be at approximately 30%, 60% and 90% of the maximum scale indication for each range.
For instruments with logarithmic scales or a digital display, measurements shall be performed on a minimum of three instrument indications in each order of magnitude. It is recommended these indications be at approximately 20%, 40% and 80% of each order of magnitude.

Any ranges or decades that have been untested must be clearly indicated in the test report.

Sufficient readings should be taken to establish a mean indication with an accuracy of ±10% of the standard deviation of the mean.

Where the dose equivalent rate meter utilises more than one radiation detector to cover the full range of dose equivalent rates indicated by the meter, these requirements apply to the relevant ranges for each detector separately.

The relative intrinsic error, \( I \), is defined as the ratio of the dose equivalent rate meter’s value to the conventionally true value.

**Equation 1.** Dose rate instrumentation linearity of response:

\[
I = \frac{H_i}{H_t}
\]

where \( H_i \) is the (mean) indicated dose equivalent rate value above background and \( H_t \) is the conventionally true dose equivalent rate value at the point of reference.

6.2 **Response Time**

The manufacturer shall state the maximum response time of the instrument.

When the instrument is subjected to a step increase or decrease in dose equivalent rate, its indication shall reach the value given by the response time equation.

**Equation 2.** Response time:

\[
H_i + \frac{90}{100} (H_f - H_i)
\]

Where \( H_i \) is the initial indication and \( H_f \) is the final indication.

6.2.1 **Essential**

When the instrument is subjected to a step increase or decrease in dose equivalent rate, its indication shall reach the value given by the response time equation in less than 10 seconds for values of \( H_f \) between 1 \( \mu \)Sv h\(^{-1}\) and 10 mSv h\(^{-1}\).

For values of \( H_f \) above this, the time shall be 2 seconds or less.
6.2.2 Desirable
When the instrument is subjected to a step increase or decrease in dose equivalent rate, its indication shall reach the value given by the response time equation in less than 3 seconds for values of \( H_f \) between 1 μSv h\(^{-1}\) and 10 mSv h\(^{-1}\).

For values of \( H_f \) above this, the time shall be 2 seconds or less.

6.2.3 Test radiation
The recommended reference radiation source for photon and β dose rate instruments is \(^{137}\)Cs. If \(^{137}\)Cs is not available then the test may be performed by injecting a suitable electrical signal into the input of the instrument. The type of method used shall be stated.

6.2.4 Method of test
The initial and final dose equivalent rates shall differ by at least a factor of 10. This test shall be performed for both an increase and a decrease in the dose equivalent rate by this factor.

Where instruments have a digital or a logarithmic display, measurements shall be made over every order of magnitude of indication. Where instruments have a linear display, measurements shall be made on each scale range.

For linear scale instruments, the lower dose equivalent rate may be taken as zero (or background) and the upper as that corresponding to at least 90% of the scale maximum on each range.

For the increasing dose equivalent rate test, the instrument shall first be subjected to the higher dose equivalent rate for a time sufficient for the indication to reach a steady value and the indication \( H_f \) recorded. Next, the instrument shall be subjected to the lower dose equivalent rate for a time sufficient for the indication \( H_i \) to reach a steady value and this indication noted.

With the instrument still subjected to the lower dose equivalent rate, this rate shall then be increased as quickly as possible (less than 1 second) to that corresponding to the indication \( H_f \). The time taken to reach the value given by the response time equation shall be recorded.

For the decreasing dose equivalent rate test, the test shall be performed in the same way but with the values of dose equivalent rates corresponding to \( H_i \) and \( H_f \) interchanged.

6.3 Statistical Fluctuations
Due to the random nature of the emissions from radioactive materials, any test of the statistical variation of an instrument reading will have an additional fluctuation superimposed. Where this additional fluctuation is a significant fraction of the variation of indication, then a significant number of readings shall be taken to ensure that the mean value of the readings is
obtained with sufficient accuracy. In order to ensure these readings are statistically independent, the interval between such readings shall be a minimum of three times the instrument’s response time.

6.3.1 **Essential**
The statistical fluctuation shall be less than ±10% at approximately 10 μSv h⁻¹.

6.3.2 **Desirable**
The statistical fluctuation should be less than ±5% at approximately 10 μSv h⁻¹.

6.3.3 **Test radiation**
The recommended reference radiation source for photon and β dose rate instruments is ¹³⁷Cs.

6.3.4 **Method of test**
The limit of variation of the statistical fluctuation shall be determined at approximately 10 μSv h⁻¹, which is considered a level that would be unlikely to arise due to the effects of natural background only.

6.4 **Background Indications**
Variation in background readings can arise for a variety of reasons, some of which are more applicable for certain types of radiation monitoring equipment. Radioactive decay is a random process and therefore the number of events ‘seen’ by a detector can vary over short time periods; for this reason the displayed background can vary significantly. Also, for some types of detector, variations in background material can lead to easily observable differences in displayed dose rates.

6.4.1 **Essential**
The statistical fluctuation shall be less than ±30% at less than 1 μSv h⁻¹.

6.4.2 **Desirable**
The statistical fluctuation shall be less than ±15% at less than 1 μSv h⁻¹.

6.4.3 **Test radiation**
The background indication shall be determined for an ambient dose equivalent rate of less than 1 μSv h⁻¹.
6.4.4 **Method of test**

The background indication of an instrument shall be determined by taking sufficient readings, at time intervals that exceed three time constants, in order to ensure that each reading is statistically independent.

6.5 **Overload Performance**

6.5.1 **Essential**

On all ranges, when exposed to dose rates greater than its measuring range, the instrument shall remain ‘off scale’ at the higher end of the scale (or display an overload indication). The instrument must remain in this overload state whilst exposed to that, or a greater radiation field, and should not return ‘on scale’ or display any indication of dose rate until the exposed dose rate is within measuring range. On completion of the overload test, the instrument shall function correctly in a time period of less than 5 minutes.

NOTE: The overload test may be ignored for instruments that are likely to be damaged by performing this test. If this test is omitted, this must be clearly recorded in the test report.

6.5.2 **Desirable**

As requirement above, but with the following change: on completion of the overload test, the instrument shall function correctly in a time period not exceeding 2 minutes.

6.5.3 **Test radiation**

The recommended reference radiation source is $^{137}$Cs.

For the purpose of this test, the value of the true ambient dose equivalent rate produced by the reference gamma radiation at the point of test shall be known with an uncertainty of less than 10% at a confidence level of 95%.

6.5.4 **Method of test**

The instrument shall be exposed to the following rates as appropriate for a period of at least 5 minutes. A minimum of 10 mSv h$^{-1}$ should be used.

- 100 times the range maximum for ranges with a maximum indication up to and including 100 mSv h$^{-1}$.
- 10 times the range maximum for ranges with a maximum indication greater than 100 mSv h$^{-1}$.

After the overload test, the instrument’s performance shall return back to normal within the time specified.
6.6 **Photon Energy Response**

The photon energy response of the instrument shall be determined using the narrow spectrum series of X-radiation qualities as defined in ISO 4037-1. In addition, the gamma radiations of $^{137}$Cs (662 keV), $^{241}$Am (59.5 keV) and $^{60}$Co (1.25 MeV) shall be used, depending upon sufficient dose rate range.

6.6.1 **Essential**

The instrument’s ambient dose equivalent response to the $^{137}$Cs (662 keV) reference gamma radiation source shall be approximately unity.

The instrument’s response to incident photon radiation between 60 keV and 1.25 MeV shall not vary by more than ±30% from the response to the $^{137}$Cs (662 keV) reference gamma radiation source.

6.6.2 **Desirable**

The instrument’s response to incident photon radiation between 20 keV and 1.25 MeV should not vary by more than ±20% from the response to the $^{137}$Cs (662 keV) reference gamma radiation source.

6.6.3 **Test radiation**

The recommended reference radiation sources are:

- Narrow series of X-radiation qualities as defined in ISO 4037-1
- $^{241}$Am or filtered X-rays of approximately 60 keV
- Gamma (γ) radiation from $^{137}$Cs at 662 keV
- $^{60}$Co at 1.25 MeV (mean energy).

6.6.4 **Method of test**

Ideally, the same dose equivalent rate should be used for each radiation energy used. Where this isn’t possible, the instrument indication for each energy used shall be corrected for intrinsic error of the indicated rate.

Measurements shall be performed starting with high energies and repeated with lower energies, until the first energy where the response normalised to that obtained for $^{137}$Cs provides a normalised response of less than 0.5.

For dose equivalent rate meters, the response normalised to $^{137}$Cs is defined as:

**Equation 3.** Photon energy response, calculation of normalised response:

\[
I_{\text{norm}} = \frac{I_{\text{energy}_x}}{I_{Cs^{137}}}
\]
where:

\[ I_{\text{energy}} = \text{response at energy } n \]
\[ n = 20 \text{ keV to } 1.25 \text{ MeV} \]
\[ I_{\text{Cs}} = \text{response at } ^{137}\text{Cs} \]

6.7 Polar Response

6.7.1 Essential
For angles of ±180°, in both horizontal and vertical planes, the response normalised to the reference radiation at 0° shall be unity ±30%.

6.7.2 Desirable
For angles of ±180°, in both horizontal and vertical planes, the response normalised to the reference radiation at 0°, shall be unity ±20%.

6.7.3 Test radiation
This test shall be performed with \(^{137}\text{Cs}\), \(^{241}\text{Am}\) (or the equivalent filtered X-radiation) and also with the lowest energy found during the energy response measurements that produced a normalised response greater than 0.6.

The conventionally true ambient dose equivalent rate produced by the reference radiation at the point of test shall be known with an uncertainty of less than 10% at the 95% confidence level.

6.7.4 Method of test
The instrument shall be mounted in its calibration orientation as specified by the manufacturer. The instrument indication and its response to the reference radiation in this orientation (0°) shall be recorded.

Next, the instrument shall be rotated about its reference point from angles 0° to ±180° in 10° steps. This shall be performed in two planes: vertical and horizontal. At each step, expose the instrument to the same dose rate as at 0° and record the instrument indication and its response to the reference radiation. These responses shall then be normalised to the reference radiation at 0°.

6.8 Accuracy of Dose Equivalent Rate Alarm

6.8.1 Essential
When the instrument is subjected to a dose rate that is 20% lower than the dose rate to which the alarm is set, the alarm shall not be triggered more than 10% of the time.
When the instrument is subjected to a dose rate that is 20% greater than the dose rate to which the alarm is set, the alarm shall remain triggered more than 90% of the time.

6.8.2 Desirable
When the instrument is subjected to a dose rate that is 10% lower than the dose rate to which the alarm is set, the alarm shall not be triggered more than 5% of the time.

When the instrument is subjected to a dose rate that is 10% greater than the dose rate to which the alarm is set, the alarm shall remain triggered more than 95% of the time.

6.8.3 Test radiation
The conventionally true ambient dose equivalent rate produced by the reference radiation at the point of test shall be known with an uncertainty of less than 10% at the 95% confidence level.

For measurements performed in terms of $H^*(10)$, the reference radiation shall be $^{137}\text{Cs}$ gamma radiation.

6.8.4 Method of test
With the instrument mounted in its calibration orientation as specified by the manufacturer, subject the instrument to the specified lower dose equivalent rate. During a period of 10 minutes, record the total length of time that the alarm is triggered.

Next, subject the instrument to the specified greater dose equivalent rate. During a period of 10 minutes, record the total length of time that the alarm is triggered.

6.9 Response to Neutron Radiation
The testing of instruments with neutron radiation is not covered in this document.

6.9.1 Essential
The manufacturer shall state the instrument’s response to neutron radiation.

6.9.2 Desirable
Not currently defined.

6.10 Response to Beta Radiation
The testing of instruments with beta radiation is not covered in this document.
6.10.1 Essential
The manufacturer shall state the instrument’s response to beta radiation.

6.10.2 Desirable
Not currently defined.
7 Alarms

7.1 General
For the purpose of this standard, an alarm is taken to be anything that draws the user’s attention to the device.

If alarm trigger points are settable, they shall be protected from unauthorised and accidental changes.

If any alarm can be switched off or reduced in intensity, they shall be protected from unauthorised and accidental changes.

Where it is possible to turn off or reduce intensity of any alarm, the instrument shall have a vibrating alarm.

7.2 Audible Alarm

7.2.1 Essential
Audible alarms shall exceed 100 dBA at a distance of 30 cm and be within the frequency range of 1 to 4 kHz. The maximum sound level for audible alarms shall not be greater than 120 dBA at the minimum hearing distance from the instrument.

An option to mute the alarm shall be provided, but only available through the supervisor mode to prevent inadvertent muting of the alarm.

Where multiple alarms are available, each alarm shall have a unique sound.

Any alarm shall not sound similar to any other emergency service equipment alarms, such as distress signals or low oxygen alarms. Alarms should be distinguishable from these by amplitude, frequency modulation or pattern where possible.

Examples of emergency service equipment alarms currently in use are defined below:

- Honeywell O2 meter: audible alarm with a pure tone at 3875 Hz;
- Diktron: various audible alarms with an upper frequency of 2900 Hz (±200 Hz).

7.2.2 Desirable
The intensity of the alarm should be fully adjustable from off to the maximum sound level. This is particularly important for covert operations.
Where it is possible to adjust the alarm intensity, the instrument shall have a vibrating alarm.

The instrument should have the ability to attach earphones.

7.3 Visual Alarm

7.3.1 Essential
The visual alarm, such as flashing light or display indication, shall be positioned so that, if triggered, it the user will easily notice it.

7.3.2 Desirable
Where the visual alarm is of high intensity, such as a beacon or similar, then the intensity of the visual alarm should be fully adjustable from off to the maximum brightness level. This is particularly important for covert operations or during use in dark conditions.

Where it is possible to adjust the alarm intensity, the instrument shall have a vibrating alarm.

7.4 Vibrating Alarm
A vibrating alarm is desirable for use in covert operations or in situations where other alarms may cause distress.

7.4.1 Essential
The vibrating alarm shall have sufficient intensity to be easily felt by the user through gloved hands.

7.4.2 Desirable
If it is possible to turn off or reduce the intensity of any visual or audible alarms, the dose meter shall have a vibrating alarm.
8 Electric Performance Requirements

8.1 Power Supply (Batteries)
The object of this element of the test is to evaluate the battery test mode of the instrument, so as to demonstrate that it provides a valid check on the state of the batteries and connectors. In addition to low voltage checks, the test simulates the presence of a good battery, but with significant corrosion on the battery terminals or connectors. This can be achieved by connecting the instrument under test to a variable power supply and a variable resistor in series.

An estimation of the battery life shall be determined from information on the current drawn by the instrument under realistic operating conditions.

The total number of batteries or cells required to power the instrument shall be noted, and the ease of their supply and replacement determined.

8.1.1 Voltage dependence
This test is designed to simulate any problems that may occur as the instrument’s batteries discharge through normal use.

8.1.1.1 Essential
Unless the instrument is displaying the low battery indication, all functions shall operate correctly and the mean instrument indication will not vary by more than ±20% from the indication recorded with an optimal battery voltage.

8.1.1.2 Desirable
Unless the instrument is displaying the low battery indication, all functions shall operate correctly and the mean instrument indication will not vary by more than ±10% from the indication recorded with an optimal battery voltage.

8.1.1.3 Test radiation
The recommended reference radiation source for photon and β dose rate instruments is $^{137}$Cs.

8.1.1.4 Method of test
With the instrument’s internal batteries removed, connect the instrument to a variable power supply, a variable resistance and a means to monitor the voltage and current in the circuit. The power supply shall be set to supply the optimal battery operating voltage, and the variable resistance shall be negligible.
With the instrument in background conditions, decrease the supply voltage in small decrements and record the supply voltage, instrument indication and any other observations, until the instrument switches off. Particular attention should be made as to when the low battery indication is triggered or if the indication varies by more than ±10% from the indication recorded with an optimal battery voltage.

This test should then be repeated with the instrument exposed to a dose equivalent rate of between 10 μSv h⁻¹ and 100 μSv h⁻¹. If possible, a dose rate should be chosen that doesn't trigger any audible or visual alarms.

Finally, this test should be repeated with the instrument exposed to a dose equivalent rate that triggers the audible or visual alarms.

8.1.2 Current dependence

This test is designed to simulate the presence of a good battery, but with significant corrosion on the battery terminals or connectors.

8.1.2.1 Essential

The current drawn by the instrument shall be as low as possible.

Unless the instrument is displaying the low battery indication, all functions shall operate correctly and the mean instrument indication will not vary by more than ±20% from the indication recorded with an optimal battery voltage.

8.1.2.2 Desirable

Unless the instrument is displaying the low battery indication, all functions shall operate correctly and the mean instrument indication will not vary by more than ±10% from the indication recorded with an optimal battery voltage.

8.1.2.3 Test radiation

The recommended reference radiation source for photon and β dose rate instruments is ¹³⁷Cs.

8.1.2.4 Method of test

With the instrument’s internal batteries removed, connect the instrument in series to a variable power supply, a variable resistance and a means to monitor the voltage and current in the circuit. The power supply shall be set to supply the optimal battery operating voltage, and the variable resistance shall be negligible.

With the instrument in background conditions, increase the series resistance in small increments and record the supply voltage, resistance, current drawn, instrument indication and any other observations until the
instrument switches off. Particular attention should be made as to when the low battery indication is triggered or if the indication varies by more than ±10% from the indication recorded with an optimal battery voltage.

This test should then be repeated with the instrument exposed to a dose equivalent rate of between $10 \mu$Sv h$^{-1}$ and $100 \mu$Sv h$^{-1}$. If possible, a dose rate should be chosen that doesn't trigger any audible or visual alarms.

Finally, this test should be repeated with the instrument exposed to a dose equivalent rate that triggers the audible or visual alarms.

### 8.1.3 Battery test function (applicable to all instruments)

The instrument shall have a means to assess the battery condition. The battery condition shall be indicated to enable the operator to assess when the battery condition is no longer suitable.

The rating for the battery test function should be based upon data provided by the voltage and current dependence tests.

#### 8.1.3.1 Essential

The battery test shall provide an accurate assessment of the condition of the cells and their suitability for further powering the instrument.

#### 8.1.3.2 Desirable

Not currently defined.

#### 8.1.3.3 Failure

Indicates a battery test with serious deficiencies that could result in incorrect measurements being made.

### 8.1.4 Battery test function (applicable only to digital instruments)

The instrument shall have a means to estimate the remaining battery life under the normal and maximum load conditions expected during use. The battery condition shall be indicated and the operator alerted when the battery condition is becoming unsuitable for the instrument to meet the requirements in this document.

The rating for the battery test function should be based upon data provided by the voltage and current dependence tests.

#### 8.1.4.1 Essential

During normal operation the battery condition shall be monitored, such that the operator is alerted when the expected remaining life of the battery falls below 30 minutes.
8.1.4.2 Desirable
During normal operation the battery condition shall be monitored, such that the operator is alerted when the expected remaining life of the battery falls below 1 hour.

8.2 Batteries

8.2.1 General
Consideration shall be given to the fact that below -10 °C the capacity of most types of batteries significantly decreases with decreasing temperature.

8.2.1.1 Essential
Batteries shall be installed in separate compartments to the instrument electronics.

Batteries shall be easily accessible for replacement and routine maintenance. The correct polarity shall be clearly indicated on the instrument.

In the event of a CBRN incident, instrumentation will be required to operate in the field. The instrument, therefore, shall be capable of running solely on a battery supply. Any chargers supplied should be capable of recharging the batteries from a vehicle 12 volt cigarette lighter socket as well as a standard 240 volt mains supply.

8.2.1.2 Desirable
All batteries should be easy to change in the field without a special tool.

Batteries should be of a standard recognised type and be easy to obtain off the shelf from retail suppliers. The preferred commercially available battery size is AA (also known internationally as LR6 or MN1500). Standard type rechargeable batteries are acceptable (see requirements for rechargeable batteries in section 8.2.3).

8.2.2 Bespoke batteries
Bespoke batteries are acceptable if they can sustain the operation of the dose meter for a significant increase of time over standard batteries. If bespoke batteries are required then a second (spare) battery must be supplied and, in addition, a reliable supply of additional batteries shall be guaranteed for a minimum of 10 years.
8.2.3 Rechargeable batteries

8.2.3.1 Essential
All rechargeable batteries shall be able to be charged independently of the instrument. A second set of rechargeable batteries and a fast charger shall be supplied. A fully discharged battery shall be able to be fully recharged within 2 hours.

8.2.3.2 Desirable
It should be possible to operate the instrument whilst its installed batteries are recharging. There should be an indication of the current status of charging.

8.2.4 Battery lifetime

8.2.4.1 Essential
After fresh or fully recharged batteries have been fitted, the instrument must be capable of operating under standard test conditions with no alarms or illumination features in operation for at least 12 hours.

After fresh or fully recharged batteries have been fitted, the instrument must be capable of operating under standard test conditions with ALL alarms and illumination features in operation at their maximum intensity for at least 1 hour.

8.2.4.2 Desirable
After fresh or fully recharged batteries have been fitted, the instrument must be capable of operating under standard test conditions with no alarms or illumination features in operation for at least 24 hours.

After fresh or fully recharged batteries have been fitted, the instrument must be capable of operating under standard test conditions with ALL alarms and illumination features in operation at their maximum intensity for at least 2 hours.

8.3 External DC or AC Power Supplies

8.3.1 Essential
For the majority of uses the instrument will be powered solely by a battery supply. If rechargeable batteries are supplied then some means of charging these batteries shall also be supplied. Power supplies for charging the batteries shall include one or more power adapters to enable charging from a standard UK 240 volt mains supply and from a nominal 12 volt vehicle electrical system. Protection against over voltage and reverse polarity shall be provided.
8.4 **Electromagnetic Compatibility**

The instrument shall be electronically compatible and not interfere with emergency service communications equipment including UHF hand held radios, VHF main schemes radios and mobile radios. The electronic interference tolerance of the unit shall be quoted.

An example of frequencies currently utilised by emergency service communication equipment is defined below:

- 380 to 400 MHz
9 Mechanical Requirements

9.1 Mechanical Shock (Drop Test)

9.1.1 Essential
The instrument shall be able to withstand drops from heights of 1 metre on to a hardwood surface without severe mechanical damage. The drop shall not affect the indicated dose rate reading by more than 20%.

9.1.2 Desirable
The instrument shall be able to withstand drops from heights of 1 metre on to a concrete surface without severe mechanical damage. The drop shall not affect the indicated dose rate reading by more than 10%.

9.1.3 Test radiation
The recommended reference radiation source is $^{137}$Cs.

For the purpose of this test, the value of the true ambient dose equivalent rate produced by the reference gamma radiation at the point of test shall be known with an uncertainty of less than 10% at a confidence level of 95%.

9.1.4 Method of test
First, the instrument’s response to the appropriate reference nuclide shall be determined in a reproducible geometry.

Next, the instrument shall be dropped on to each face in turn from a height of 1 metre on to the specified surface. After each drop, the instrument’s response to the appropriate reference nuclide shall be confirmed and recorded. The instrument shall also be checked for any mechanical damage or loose fittings and any observations recorded.

9.2 Vibration
The physical condition of the instrument shall not be affected by harmonic loadings of 2 g applied for 15 minutes in the frequency range 10 to 33 Hz, i.e. all electrical connections and mechanical fastenings shall hold and not become loose.

9.2.1 Essential
The mean response of the instrument shall not vary by more than 20% as a result of these vibrations.
9.2.2 Desirable
The mean response of the instrument shall not vary by more than 10% as a result of these vibrations.

9.2.3 Test radiation
The recommended reference radiation source for photon and β dose rate instruments is $^{137}\text{Cs}$. If $^{137}\text{Cs}$ is not available, then $^{60}\text{Co}$ can be used although corrections will need to be applied to relate the response to $^{137}\text{Cs}$.

9.2.4 Method of test
The instrument shall be exposed to photon radiation in reproducible geometry and the mean indication determined. The instrument shall then be subjected to harmonic loadings of 2 g applied for 15 minutes in each of three planes. At least one test shall be performed within each of the ranges 10 to 21 Hz and 22 to 33 Hz. After each vibration the mean indication will be determined using the same radiation conditions and geometry as before. All pre- and post-vibration readings shall be recorded as well as the physical condition of the instrument.
10 Environmental Performance Requirements

Instruments shall be so designed and constructed as to be capable of performing their intended function in full safety in changing environmental situations.

10.1 Environmental Protection

The manufacturer should state the environmental protection classification of the instrument. Where this is not supplied, a visual inspection of probable ingress shall be performed and recorded.

10.1.1 Essential

The instrument shall have an IP rating of at least IP54. Instruments shall be designed to resist ingress from dust, wind driven rain, high humidity or condensation. If the instrument has been disassembled for any reason the manufacturer shall state which seals or gaskets would need to be replaced to retain acceptable weather protection.

[IP 5x — Dust protected. Ingress of dust is not totally prevented, but dust shall not penetrate in a quantity to interfere with the satisfactory operation of the apparatus or to impair safety.]

[IP x4 — Protected against splashing water. Water splashed against the enclosure from any direction shall have no harmful effects.]

10.1.2 Desirable

The instrument should have an IP rating close to IP67. Instruments should be designed to resist water ingress from temporary immersion.

[IP 6x — Dust tight. No ingress of dust.]

[IP x7 — Protected against temporary immersion. Ingress of water in harmful quantity shall not be possible when the enclosure is immersed in water under defined conditions of pressure and time.]

10.2 Temperature Stability

The object of this test is to determine the dependency of the instrument response on temperature.

10.2.1 Essential

The worst case percentage change of the indications at -10 °C and +40 °C compared to the indication at 20 °C for both radiation levels shall be less than 50%. 
10.2.2 Desirable
The worst case percentage change of the indications at $-10 \, ^\circ\text{C}$ and $+60 \, ^\circ\text{C}$ compared to the indication at $20 \, ^\circ\text{C}$ for both radiation levels shall be less than 20%.

10.2.3 Test radiation
The recommended reference radiation source for photon and $\beta$ dose rate instruments is $^{137}\text{Cs}$.

10.2.4 Method of test
The instrument shall be placed in a climatic chamber initially set to an operating temperature of $20 \, ^\circ\text{C}$ and allowed to stabilise for a minimum of 60 minutes. Both the instrument background reading and a higher indication produced by a radioactive source shall be recorded.

The chamber temperature shall be increased to the specified upper temperature and the instrument left for a minimum of four hours to achieve thermal equilibrium. The instrument response at background and the higher indication shall be recorded.

This test shall then be repeated with a temperature of $-10 \, ^\circ\text{C}$.

10.3 Temperature shock

10.3.1 Essential
The instrument should be capable of working up to temperatures of $+60 \, ^\circ\text{C}$.

10.3.2 Desirable
Not currently defined.

10.3.3 Test radiation
The recommended reference radiation source for photon and $\beta$ dose instruments is $^{137}\text{Cs}$.

10.3.4 Method of test
The instrument shall be placed in a climatic chamber initially set to an operating temperature of $20 \, ^\circ\text{C}$ and allowed to stabilise for a minimum of 60 minutes. Both the instrument background reading and a higher indication produced by a radioactive source shall be recorded.

Next, the chamber temperature shall be increased to the specified upper temperature within 5 minutes. The readings at background and the higher indication shall be repeated and recorded every 15 minutes for a duration 2 hours.
Next, the chamber temperature shall be returned to the original temperature of 20 °C within 5 minutes. The readings at background and the higher indication shall be repeated and recorded every 15 minutes for a duration 2 hours.

Finally, the chamber temperature shall be decreased to -10 °C within 5 minutes. The readings at background and the higher indication shall be repeated and recorded every 15 minutes for a duration 2 hours.

10.4 Low Temperature Start-up

10.4.1 Essential
The instrument shall switch on and operate correctly at -10 °C.

10.4.2 Desirable
Not currently defined.

10.4.2.1 Test radiation
The recommended reference radiation source for photon and β dose instruments is $^{137}$Cs.

10.4.2.2 Method of test
The instrument shall be placed in a climatic chamber initially set to an operating temperature of -10 °C and allowed to stabilise for a minimum of 4 hours. The instrument shall then be switched on.

Both the instrument background reading and a higher indication produced by a radioactive source shall be recorded. These indications shall be within ±20% of those recorded during the temperature stability tests at 20 °C.

Any additional observations shall also be recorded.

10.5 Humidity Stability

10.5.1 Essential
The instrument shall be capable of working at relative humidity levels between 20% RH and 90% RH. The variation of the relative response due to humidity shall be less than ±20%.

10.5.2 Desirable
The variation of the relative response due to humidity shall be less than ±10%.
10.5.3 Test radiation
The recommended reference radiation source for photon and β dose rate instruments is $^{137}\text{Cs}$.

10.5.4 Method of test
The instrument shall be placed in a climatic chamber initially set to an operating temperature of +35 °C and a relative humidity of 65% RH. The instrument shall be left switched off in these conditions for a minimum of 24 hours. In the last 30 minutes of this period, the instrument should be switched on and both the instrument background reading and a higher indication produced by a radioactive source shall be recorded.

Keeping the temperature at 35 °C, increase the relative humidity inside the chamber to 90% RH. The instrument shall be left switched off in these conditions for a minimum of 24 hours. In the last 30 minutes of this period the instrument should be switched on and both the instrument background reading and a higher indication produced by a radioactive source shall be recorded.

This test shall then be repeated with a relative humidity of 20% RH.

NOTE: For this test the reference response is determined at +35 °C and not +20 °C.

10.6 Submersion

10.6.1 Essential
The level of water resistance shall be clearly stated in the manual. Where the level of water resistance has not been tested then this shall be stated in the manual.

10.6.2 Desirable
The instrument should be capable of satisfactory operation after being fully submerged under water for 5 minutes.

10.6.3 Method of test
The instrument shall be submerged under water at a depth of approximately 30 mm for a period of at least 5 minutes and then thoroughly dried. Both the instrument's background reading and a higher indication produced by a radioactive source shall be recorded.

10.7 Explosive Atmospheres
Circumstances can be foreseen where it is necessary for instruments to be used in flammable atmospheres or close to explosive devices.
10.7.1 **Essential**
The manufacturer shall clearly state the level of intrinsic safety.

10.7.2 **Desirable**
The instrument should be intrinsically safe. Potential ignition sources, such as sparks, electric arcs or high surface temperatures, should not occur.

Instruments shall be designed so that the opening of equipment parts, which may be sources of ignition, is only possible under non-active or intrinsically safe conditions. Where it is not possible to render the equipment non-active, a warning label shall be affixed to the opening part of the equipment.
11 Maintenance Requirements

The manufacturer must define the time limit that the instrument will be supportable for spares and repairs. This shall be a minimum of 10 years. The manufacturer shall provide details of technical support and advice options, as well as recommendations and information on the testing and maintenance regime required.

The instrument shall be supplied with a comprehensive instruction manual. In addition, a maintenance manual shall be available upon request.

The instrument shall contain a sufficient amount of easily accessible test points to facilitate fault location. Any maintenance aids, such as fault diagnosis software, extension leads and special maintenance tools, shall be available from the manufacturer upon request.

Unauthorised access to all the set-up functions of the equipment shall be prevented (see paragraph 3.3 Operating modes).
12 Ergonomic Requirements

12.1 General
The object of this assessment is to give some idea of the ease of regular use of the instrumentation. Owing to the nature of the parameters assessed, the ratings are, to a large extent, subjective and should be based on impressions gained during the testing and handling of the instruments. Factors to be considered are:

- ease of operation;
- clarity of the display;
- ease of decontamination;
- susceptibility to damage.

12.1.1 Essential
Features are reasonably satisfactory.

12.1.2 Desirable
Features are fully satisfactory and could not be usefully improved.

12.1.3 Failure
Indicates a poor feature that could be irritating or inconvenient during regular use.

12.2 Size

12.2.1 Essential
The instrument shall fit into a volume of less than 2 litres, excluding any external probes.

12.2.2 Desirable
The instrument should fit into a volume of less than 1 litre, excluding any external probes.

12.3 Weight

12.3.1 Essential
The instrument shall be as light as possible with a maximum weight of 2 kg. The instrument shall be well-balanced.
12.3.2 Desirable
The instrument should be as light as possible with a maximum weight of 1 kg.

12.4 Case Construction

12.4.1 Essential
The instrument case shall be smooth, rigid, shock resistant, splash proof and dust resistant. An additional rubberised outer to minimise damage can be utilised, but where this is removable it shall not affect the instrument’s radiological performance.

Care shall be taken to ensure that any control, display or visual alarm is located such that they are visible to the operator.

The case shall be provided with the facility to attach support straps. These will allow for freedom of movement whilst carrying out strenuous work. Any support straps provided must be adjustable and compatible for use whilst wearing personal protection equipment (PPE); e.g. a full gas-tight suit.

12.4.2 Desirable
Not currently defined.

12.5 Resistance to Contamination (Ease of Decontamination)

12.5.1 Essential
The instrument shall be easy to decontaminate, ideally without any areas where contaminants could become difficult to remove. A smooth non-porous external surface that is free from crevices is recommended. The instrument may be fitted with an additional protective cover providing this doesn’t affect any aspects of the instrument’s performance.

12.5.2 Desirable
Not currently defined.

12.6 Transportation

12.6.1 Essential
Not currently defined.

12.6.2 Desirable
Capable of surviving high altitude air transport.
12.7 Cabling and Connections

12.7.1 Essential
All cabling shall be substantial and have strain relief where it is terminated. Rugged connectors shall be used and these should be securable. The integrity of non-securable connectors shall be protected.

12.7.2 Desirable
Not currently defined.

12.8 Switches and Controls

12.8.1 Essential
All switches and other controls shall be designed to ensure that the instrument can be properly operated while minimising accidental operation of any controls. All switches and controls shall have a positive feel that they have been operated.

12.8.2 Desirable
All switches and controls should be illuminated such that their location and function can be identified in dark conditions. The spacing between each switch or control should at least 15 mm so as to increase the ease of operation of the instrument while the operator is in full gas-tight PPE.

NOTE: The finger diameter of the gloves used with full gas-tight PPE is approximately 25 to 30 mm.

12.9 Ease of Operation

12.9.1 Essential
The instrument must be designed so that it can be used safely and efficiently without a high level of specialist knowledge.

The instrument must be easy to operate with the operator in full gas-tight PPE.

12.9.2 Desirable
The instruments should be controlled via a menu operation with “soft-keys”. One-handed operation is possible.

12.10 Detector Location
For the majority of dose rate instrumentation the detector should be integral to the instrument. However, instruments specifically designed
to measure high dose rates shall have an external extendable probe to help reduce operator doses.

Other than an external probe, the instrument shall not require any other external devices to be attached for it to function normally in the field. The provision for the attachment of an earphone is acceptable for use in noisy environments.

12.11 External Markings
All external markings shall remain permanently fixed under both normal conditions and those during normal decontamination procedures.

The instrument shall be clearly marked with the following:

- manufacturer’s name;
- model type;
- unique serial number;
- the function of all controls (that are not displayed via soft menus) and indicators.

Instruments with internal radiation detectors shall have markings to clearly indicate the calibration reference point of each detector in at least two planes.

Probes should be clearly marked to show their intended function.

12.12 Visual Display

12.12.1 Essential
Whether the instrument has an analogue or digital display (or a hybrid of the two), for indicating the dose rate, the display shall react instantly to any change of measuring range. In addition, the display shall clearly indicate the measuring quantity.

The display must be clear and easy to read under normal and extreme conditions, which include the use in bright sunlight and in total darkness. The display shall have an illumination function that can be turned on and off and this must not time out. A provision to test for failure of the display shall be installed. The display should not be influenced by gravity.

In addition to the visual indication of dose rate, an audible indication of dose rate shall be provided. A facility for muting this indication shall also be provided. Where the equipment has been designed for use where ambient noise levels could be high, provision for the connection of a head set shall be made.

The size of the display shall be at least 45 x 15 mm.
12.12.2 Desirable
The brightness of the display illumination should be fully adjustable from off to the maximum brightness. There should be an option for the illumination brightness to adjust automatically dependent on the ambient lighting conditions.

The size of the display shall be at least 70 x 40 mm.

12.13 Additional Indications

12.13.1 Low battery

12.13.1.1 Essential
An indication of the battery condition shall be displayed on the display.

12.13.2 Detector failure

12.13.2.1 Essential
The instrument shall detect when the detector has failed and alert the operator.

12.14 Firmware

12.14.1.1 Essential
In the design of any firmware-controlled instrumentation, special account shall be taken of the risks arising from faults in the program.

The software shall have a version number for identification. It shall be possible to display this identification whilst the software is running.

All commands or parameters shall be defined; i.e. they shall have a clearly defined function that can be processed by the instrument, otherwise the instrument shall identify them as invalid. Invalid commands shall not affect any data or functions of the instrument.

12.14.1.2 Desirable
If applicable, firmware stored in the dose meter should be easy to update. Any updates shall be possible using only a PC with USB or other standard communications port. The dose meter shall request re-calibration after any update and this shall be made clear to the operator on every occasion before any updates are made.
12.15 Data Logging

12.15.1 Essential
Not currently defined.

12.15.2 Desirable
The instrument should have a data logging capability, with a download facility for extracting and/or interrogating data and logging faults to a computer. In the event of a power/battery failure the instrument should retain this data.

12.16 Communication Interface

12.16.1 Essential
Not currently defined.

12.16.2 Desirable
The instrument should be able to communicate data to an external device, such as a computer. The type of data to transfer could include dose equivalent rate indication history with time and date and/or GPS location.

The data transfer shall be via a bi-directional serial port that meets the requirements of Ethernet, USB or other, by other electronic means, such as a standard removable media device (e.g. SD Card). The protocol used shall conform to applicable IEEE protocols (e.g. IEEE 802) and proprietary protocols shall not be used. The transferred data shall be of a format (e.g. ASCII) that can be easily imported into common analysis programs. The manufacturer shall provide a full description of the transfer format and, if required, proprietary software for data interpretation.

Whilst it is acceptable in some circumstances for data to be communicated wirelessly (e.g. via Bluetooth® or Wi-Fi), all instruments shall have the option to fully disable its radio communications.
13 Documentation

13.1 Type-test Report
The manufacturer shall make the relevant type-test report available to any user or potential user of the instrument. If requested, the type-test report shall be supplied in its entirety.

13.2 Calibration Certificate Requirements
A certificate shall be provided giving at least the following information:

- manufacturer’s name or registered trade mark;
- instrument type and serial number;
- probe type and serial number (if applicable);
- types and energies of radiations for which the instrument is intended;
- reference point of device;
- calibration orientation relative to radiation sources;
- effective range of use.

13.3 Operation and Maintenance Manual

13.3.1 Essential
The manufacturer shall supply an operational and maintenance manual containing a minimum of following information:

- operating instructions and restrictions;
- schematic electrical diagrams, spare parts list and specifications;
- troubleshooting guide;
- contact information for the manufacturer.

13.3.2 Desirable
The manufacturer should supply a quick reference guide that explains the basic operations.
14 Training

14.1.1 Essential
Not currently defined.

14.1.2 Desirable
Training simulator options should be available for realistic training.
Appendix A: Summary of Performance Criteria

A.1 Radiological

Table 2. Summary of radiological performance criteria

<table>
<thead>
<tr>
<th>Requirement</th>
<th>Essential</th>
<th>Desirable</th>
</tr>
</thead>
<tbody>
<tr>
<td>Linearity of response</td>
<td>Linearity better than ±30%.</td>
<td>Linearity better than ±20%.</td>
</tr>
<tr>
<td>Response time</td>
<td>Less than 10 seconds below 10 mSv h⁻¹.</td>
<td>Less than 3 seconds below 10 mSv h⁻¹.</td>
</tr>
<tr>
<td></td>
<td>Less than 2 seconds above 10 mSv h⁻¹.</td>
<td>Less than 2 seconds above 10 mSv h⁻¹.</td>
</tr>
<tr>
<td>Statistical fluctuations</td>
<td>Statistical fluctuations less than ±10% at approximately 10 μSv h⁻¹.</td>
<td>Statistical fluctuations less than ±5% at approximately 10 μSv h⁻¹.</td>
</tr>
<tr>
<td>Background indications</td>
<td>Statistical fluctuations less than ±30% at less than 1 μSv h⁻¹.</td>
<td>Statistical fluctuations less than ±15% at less than 1 μSv h⁻¹.</td>
</tr>
<tr>
<td>Overload performance</td>
<td>Satisfactory overload and return to normal function in less than 5 minutes.</td>
<td>Satisfactory overload and return to normal function in less than 2 minutes.</td>
</tr>
<tr>
<td>Photon energy response</td>
<td>Across the specified energy range of the instrument the response is unity ±30%.</td>
<td>Across the specified energy range of the instrument the response is unity ±20%.</td>
</tr>
<tr>
<td>Polar response</td>
<td>For angles of ±180°, in both horizontal and vertical planes, the response normalised to the reference radiation at 0°, shall be unity ±30%.</td>
<td>For angles of ±180°, in both horizontal and vertical planes, the response normalised to the reference radiation at 0°, shall be unity ±20%.</td>
</tr>
<tr>
<td>Accuracy of dose equivalent rate alarm</td>
<td>When subjected to a dose rate that is 20% lower than the dose rate to which the alarm is set, the alarm shall not be triggered more than 10% of the time.</td>
<td>When subjected to a dose rate that is 10% lower than the dose rate to which the alarm is set the alarm shall not be triggered more than 5% of the time.</td>
</tr>
<tr>
<td></td>
<td>When subjected to a dose rate that is 20% greater than the dose rate to which the alarm is set the alarm shall remain triggered more than 90% of the time.</td>
<td>When subjected to a dose rate that is 10% greater than the dose rate to which the alarm is set the alarm shall remain triggered more than 95% of the time.</td>
</tr>
<tr>
<td>Response to neutron radiation</td>
<td>The manufacturer shall state the instrument’s response to neutron radiation.</td>
<td>Not currently defined.</td>
</tr>
<tr>
<td>Response to beta radiation</td>
<td>The manufacturer shall state the instrument’s response to beta radiation.</td>
<td>Not currently defined.</td>
</tr>
</tbody>
</table>
### A.2 Alarms

Table 3. Summary of alarm performance criteria

<table>
<thead>
<tr>
<th>Requirement</th>
<th>Essential</th>
<th>Desirable</th>
</tr>
</thead>
<tbody>
<tr>
<td>Audible alarm</td>
<td>Audible alarms shall exceed 100 dBA at a distance of 30 cm and have a unique sound, which shall not sound similar to any other emergency service equipment.</td>
<td>Adjustable alarm intensity in conjunction with vibrating alarm.</td>
</tr>
<tr>
<td>Visual alarm</td>
<td>Positioned so that, if triggered, the user will easily notice it.</td>
<td>Adjustable alarm intensity in conjunction with vibrating alarm.</td>
</tr>
<tr>
<td>Vibrating alarm</td>
<td>The vibrating alarm shall have sufficient intensity to be easily felt by the user through gloved hands.</td>
<td>If it is possible to turn off or reduce the intensity of any visual or audible alarms, then the dose meter shall have a vibrating alarm.</td>
</tr>
</tbody>
</table>

### A.3 Electrical

Table 4. Summary of electrical performance criteria

<table>
<thead>
<tr>
<th>Requirement</th>
<th>Essential</th>
<th>Desirable</th>
</tr>
</thead>
<tbody>
<tr>
<td>Voltage dependence</td>
<td>Unless the instrument is displaying the low battery indication, all functions shall operate correctly and the mean instrument indication will not vary by more than ±20% from the indication recorded with an optimal battery voltage.</td>
<td>Unless the instrument is displaying the low battery indication, all functions shall operate correctly and the mean instrument indication will not vary by more than ±10% from the indication recorded with an optimal battery voltage.</td>
</tr>
<tr>
<td>Current dependence</td>
<td>Unless the instrument is displaying the low battery indication, all functions shall operate correctly and the mean instrument indication will not vary by more than ±20% from the indication recorded with an optimal battery voltage.</td>
<td>Unless the instrument is displaying the low battery indication, all functions shall operate correctly and the mean instrument indication will not vary by more than ±10% from the indication recorded with an optimal battery voltage.</td>
</tr>
<tr>
<td>Battery test function (applicable to all instruments)</td>
<td>The battery test shall provide an accurate assessment of the condition of the cells and their suitability for further powering of the instrument.</td>
<td>Not currently defined.</td>
</tr>
<tr>
<td>Battery test function (applicable only to digital instruments)</td>
<td>Operator alerted when the expected remaining life of the battery falls below 30 minutes.</td>
<td>Operator alerted when the expected remaining life of the battery falls below 1 hour.</td>
</tr>
</tbody>
</table>
## A.4 Mechanical

Table 5. Summary of mechanical performance criteria

<table>
<thead>
<tr>
<th>Requirement</th>
<th>Essential</th>
<th>Desirable</th>
</tr>
</thead>
<tbody>
<tr>
<td>Battery lifetime</td>
<td>At least 12 hours with no alarms or illumination features in operation.</td>
<td>At least 24 hours with no alarms or illumination features in operation.</td>
</tr>
<tr>
<td></td>
<td>At least 1 hour with ALL alarms or illumination features in operation.</td>
<td></td>
</tr>
</tbody>
</table>

Table 6. Summary of environmental performance criteria

<table>
<thead>
<tr>
<th>Requirement</th>
<th>Essential</th>
<th>Desirable</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mechanical shock (drop test)</td>
<td>Withstand drops from 1 metre on to a hardwood surface without severe mechanical damage. The drop shall not affect the indicated dose rate reading by more than 20%.</td>
<td>Withstand drops from 1 metre on to a concrete surface without severe mechanical damage. The drop shall not affect the indicated dose rate reading by more than 10%.</td>
</tr>
<tr>
<td>Vibration</td>
<td>Response shall not vary by more than 20% as a result of the specified vibrations.</td>
<td>Response shall not vary by more than 10% as a result of the specified vibrations.</td>
</tr>
</tbody>
</table>

## A.5 Environmental

Table 6. Summary of environmental performance criteria

<table>
<thead>
<tr>
<th>Requirement</th>
<th>Essential</th>
<th>Desirable</th>
</tr>
</thead>
<tbody>
<tr>
<td>Environmental protection</td>
<td>IP rating of at least IP54.</td>
<td>IP rating close to IP67.</td>
</tr>
<tr>
<td>Temperature stability</td>
<td>A change of less than 50% for indications at -10 °C and +40 °C compared to the indication at 20 °C.</td>
<td>A change of less than 20% for indications at -10 °C and +60 °C compared to the indication at 20 °C.</td>
</tr>
<tr>
<td>Temperature shock</td>
<td>Capable of satisfactory operation up to +60 °C.</td>
<td>Not currently defined.</td>
</tr>
<tr>
<td>Low temperature start-up</td>
<td>Shall switch on and operate correctly at –10 °C.</td>
<td>Not currently defined.</td>
</tr>
<tr>
<td>Humidity stability</td>
<td>A change of less than 20% in indications for humidity levels between 20% and 90%.</td>
<td>A change of less than 10% in indications for humidity levels between 20% and 90%.</td>
</tr>
<tr>
<td>Submersion</td>
<td>The level of water resistance shall be clearly stated in the manual.</td>
<td>Capable of satisfactory operation after being fully submerged for 5 minutes.</td>
</tr>
<tr>
<td>Explosive atmospheres</td>
<td>The manufacturer shall clearly state the level of intrinsic safety.</td>
<td>Intrinsically safe.</td>
</tr>
</tbody>
</table>
A.6  
Ergonomic

Table 7. Summary of ergonomic performance criteria

<table>
<thead>
<tr>
<th>Requirement</th>
<th>Essential</th>
<th>Desirable</th>
</tr>
</thead>
<tbody>
<tr>
<td>General</td>
<td>Features are reasonably satisfactory.</td>
<td>Features are fully satisfactory and could not be usefully improved.</td>
</tr>
<tr>
<td>Size</td>
<td>Fits in to a volume of less than 2 litres excluding any extendable probes.</td>
<td>Fits in to a volume of less than 1 litre excluding any extendable probes.</td>
</tr>
<tr>
<td>Weight</td>
<td>Maximum weight of 2 kg.</td>
<td>Light as possible with a maximum weight of 1 kg.</td>
</tr>
<tr>
<td>Case construction</td>
<td>Instrument case shall be smooth, rigid, shock resistant, splash proof and dust resistant.</td>
<td>Any supporting straps/lanyards should be adjustable and compatible for use while wearing PPE, e.g. a full gas-tight suit.</td>
</tr>
<tr>
<td>Resistance to contamination (ease of de-contamination)</td>
<td>Should not have any areas where contaminants could become difficult to remove.</td>
<td>Not currently defined.</td>
</tr>
<tr>
<td>Transportation</td>
<td>Not currently defined.</td>
<td>Capable of surviving high altitude air transport.</td>
</tr>
<tr>
<td>Cabling and connectors</td>
<td>Substantial cabling with strain relief where terminated.</td>
<td>Not currently defined.</td>
</tr>
<tr>
<td>Switches and controls</td>
<td>Designed to ensure that the instrument can be properly operated while minimising accidental operation of any controls.</td>
<td>Illuminated such that their location and function can be identified in dark conditions. The spacing between each switch or control should at least 15 mm so as to increase the ease of operation of the dose meter while the operator is in full gas-tight PPE.</td>
</tr>
<tr>
<td>Ease of operation</td>
<td>Designed so that it can be used safely and efficiently without a high level of specialist knowledge.</td>
<td>The instruments should be controlled via a menu operation with “soft-keys”. One-handed operation is possible.</td>
</tr>
<tr>
<td>Visual display</td>
<td>Clear and easy to read under normal and extreme conditions. Minimum size of 45 x 15 mm.</td>
<td>Intensity of the display illumination should be fully adjustable. Minimum size of 70 x 40 mm.</td>
</tr>
<tr>
<td>Firmware / software</td>
<td>The software shall have a version number for identification.</td>
<td>Firmware stored in the instrument should be easy to update.</td>
</tr>
<tr>
<td>Data logging</td>
<td>Not currently defined.</td>
<td>Yes</td>
</tr>
<tr>
<td>Communication interface</td>
<td>Not currently defined.</td>
<td>Be able to communicate with an external device such as a computer.</td>
</tr>
</tbody>
</table>