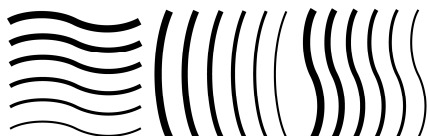


Code of Practice for the Absorbed Dose Determination in High Energy Photon and Electron Beams

NEDERLANDSE COMMISSIE VOOR STRALINGSDOSIMETRIE

Report 18 of the Netherlands Commission on Radiation Dosimetry

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Preface

The Nederlandse Commissie voor Stralingsdosimetrie (NCS, Netherlands Commission on Radiation Dosimetry, <http://ncs-dos.org>) was officially established on 3 September 1982 with the aim of promoting the appropriate use of dosimetry of ionising radiation both for scientific research and practical applications. The NCS is chaired by a board of scientists, installed upon the suggestion of the supporting societies, including the Nederlandse Vereniging voor Radiotherapie en Oncologie (Netherlands Society for Radiotherapy and Oncology), the Nederlandse Vereniging voor Nucleaire Geneeskunde (Dutch Society of Nuclear Medicine), the Nederlandse Vereniging voor Klinische Fysica (Dutch Society for Medical Physics), the Nederlandse Vereniging voor Radiobiologie (Netherlands Radiobiological Society), the Nederlandse Vereniging voor Stralingshygiëne (Netherlands Society for Radiological Protection), the Nederlandse Vereniging voor Medische Beeldvorming en Radiotherapie (Dutch Society for Medical Imaging and Radiotherapy), the Nederlandse Vereniging voor Radiologie (Radiological Society of the Netherlands) and the Belgische Vereniging voor Ziekenhuisfysici/Société Belge des Physiciens des Hôpitaux (Belgian Hospital Physicists Association).

To pursue its aims, the NCS accomplishes the following tasks: participation in dosimetry standardisation and promotion of dosimetry intercomparisons, drafting of dosimetry protocols, collection and evaluation of physical data related to dosimetry. Furthermore the commission shall maintain or establish links with national and international organisations concerned with ionising radiation and promulgate information on new developments in the field of radiation dosimetry.

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Code of Practice for the absorbed dose determination in high energy photon and electron beams

Revised edition, August 2012

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User guide

This report provides recommendations and the methodology for medical physicists to obtain the quantity absorbed dose in water under reference conditions from measurements made with an ionisation chamber in high energy photon and electron beams used in radiotherapy. A prerequisite is the traceable calibration of ionisation chambers in terms of absorbed dose to water in a ^{60}Co reference beam at a standards laboratory.

Chapter 1 gives an introduction on recent developments in dosimetric concepts and methods, measurement standards, etc. resulting in a Code of Practice for clinical photon and electron beams based on the concept of absorbed dose to water. Furthermore it introduces the basic formalism for dose determination and the concept of the beam quality correction factor.

Chapter 2 and 3 contain the Codes of Practice for the dosimetry in respectively high energy photon and electron beams generated by medical linear accelerators.

Appendices are provided giving information on methods, physical and numerical data concerning quantities, correction and conversion factors, on primary absorbed dose standards operated in Belgium and The Netherlands and on the beam quality correction factors to be used in photon and electron dosimetry. Moreover it provides an appendix concerning the differences between this Code of Practice and the previous NCS Codes based on the quantity air kerma. The last appendix deals with the estimation of uncertainty according to international guidelines recommended by ISO and EA.

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Summary

Absorbed dose to water is the quantity of interest to specify the amount of radiation to be used in radiotherapy and has the advantage that it can be measured more directly than the quantity air kerma. Advances in radiation dosimetry concepts and the development of primary measurement standards based on absorbed dose to water over the last decades offer the possibility to calibrate ionisation chambers directly in terms of absorbed dose to water. Absorbed dose standards have an uncertainty of less than 1% (1 sd) in photon beams up to 25 MV, resulting in improved accuracy in clinical reference dosimetry. Nowadays several international and national Codes of Practice (CoP) for external beam radiotherapy based on absorbed dose standards have been published and adopted in many countries worldwide.

In this report a Code of Practice is presented for the dosimetry in high energy photon and electron beams based on absorbed dose to water standards for ^{60}Co reference beams. The CoP has been written by a Subcommittee of the Netherlands Commission on Radiation Dosimetry (NCS) and corresponds to the current clinical practice in Belgium and The Netherlands. The CoP provides the concepts and methods to determine the absorbed dose in high energy photon and electron beams produced by medical linear accelerators. The CoP covers the reference dosimetry in static, open photon beams with nominal energies between 1 and 25 MV and in static, open electron beams with nominal energies between 4 and 25 MeV produced by conventional linear accelerators. Treatment modalities as intensity modulated therapy (IMRT), stereotactic radiotherapy, robotic radiotherapy, helical tomotherapy, etc. are beyond the scope of this report.

Important features are the simple basic concepts to determine absorbed dose to water under reference conditions in a water phantom and the introduction of a single, chamber dependent correction factor taking into account all effects dependent of the radiation beam quality. For photon beams these beam quality correction factors are based on experimental data, partly measured in selected clinical accelerator beams in Belgium and The Netherlands using a portable water calorimeter.

The present CoP recommends a limited number of ionisation chambers for reference dosimetry in radiotherapy, but the physical concepts outlined in the Code represent a major simplification compared to the previous Codes based on the concept of air kerma employing Bragg-Gray or Spencer-Attix cavity theory. This CoP uses updated information compared to the codes presented in TRS-398 and AAPM TG-51.

This Code of Practice replaces the protocols based on the concept of air kerma calibration coefficients described in NCS reports 2 and 5.

Abbreviations

AAPM	American Association of Physicists in Medicine
BIPM	Bureau International des Poids et Mesures
CCEMRI(I)	Comité Consultatif pour les Etalons de Mesure des Rayonnements Ionisants (Section 1) (Consultative Committee for Standards of Ionising Radiation) Since September 1997, the CCEMRI and the sections have been renamed CCRI
CCRI(I)	Comité Consultatif des Rayonnements Ionisants (Section 1) (Consultative Committee for Ionising Radiation)
CoP	Code of Practice
DIN	Deutsches Institut für Normung (German Institute for Standardisation)
EA	European Co-operation for Accreditation
ICRU	International Commission on Radiation Units and Measurements
IAEA	International Atomic Energy Agency
TRS-398	Absorbed Dose Determination in External Beam Radiotherapy: An International Code of Practice for Dosimetry Based on Standards of Absorbed Dose to Water, IAEA, Technical Report Series No. 398, Vienna, 2000
LNHB	Laboratoire National Henri Becquerel, France
ISO	International Organization for Standardization
LSDG	Laboratorium voor Standaarddosimetrie van de universiteit Gent (Laboratory for Standard Dosimetry Ghent)
NCS	Nederlandse Commissie voor Stralingsdosimetrie (Netherlands Commission on Radiation Dosimetry)
NCS-2	Nederlandse Commissie voor Stralingsdosimetrie: Code of Practice for the Dosimetry of High energy Photon Beams, NCS Report 2, December 1986
NCS-5	Nederlandse Commissie voor Stralingsdosimetrie: Code of Practice for the Dosimetry of High energy Electron Beams, NCS Report 5, December 1989
NMi	Nederlands Meetinstituut
NPL	National Physical Laboratory, United Kingdom

PMMA	Polymethyl methacrylate
PSDL	Primary Standard Dosimetry Laboratory
TG-51	AAPMs TG-51 protocol for clinical reference dosimetry of high energy photon and electron beams, <i>Med. Phys.</i> 26 1847-1870, 1999

List of symbols

In most cases the nomenclature used in publication IAEA TRS398 has been followed.

$D_{w,Q}$	Absorbed dose to water in the beam quality Q .
k_{elec}	Calibration coefficient of the electrometer, converts electrometer reading to absolute charge.
k_h	Factor to correct for the response of an ionisation chamber for the effect of humidity if the calibration coefficient of the chamber is referred to dry air.
k_{Q,Q_0}	Beam quality correction factor which corrects for the difference in the absorbed dose to water calibration coefficient of a reference beam quality Q_0 to that of a radiation beam of quality Q .
k_Q	Beam quality correction factor which corrects for the difference in the absorbed dose to water calibration coefficient of a ^{60}Co beam to that of a radiation beam of quality Q . The subscript Q_0 is omitted when the reference beam quality is ^{60}Co .
k_{pol}	Factor to correct for polarity effects affecting the response of the ionisation chamber.
k_s	Factor to correct for the response of an ionisation chamber for the lack of complete charge collection.
k_{TP}	Factor to correct for the response of an ionisation chamber for deviations of temperature and pressure from normal conditions.
$M_{\text{corr},Q}$	The electrometer reading corrected for any difference between the ambient air conditions affecting the ionisation chamber at the time of measurement and the standard ambient air conditions for which the calibration coefficient applied (air temperature, pressure and humidity), for ion recombination and for polarity effects.

M_Q	The uncorrected reading of the instrument.
$N_{D,w}$	The absorbed dose to water calibration coefficient for an ionisation chamber (or for the dosimeter assembly: ionisation chamber and electrometer) at the reference beam quality ^{60}Co .
$N_{D,w,Q}$	The absorbed dose to water calibration coefficient for an ionisation chamber (or for the dosimeter assembly: ionisation chamber and electrometer) at the beam quality Q .
$N_{D,w,Q_{\text{cross}}}$	The absorbed dose to water calibration coefficient for an ionisation chamber (or for the dosimeter assembly: ionisation chamber and electrometer) at the cross calibration quality Q_{cross} .
$\%dd(10)_x$	The photon component of the percentage depth dose at 10 cm depth in a 10 cm x 10 cm field on the surface of a water phantom at an SSD of 100 cm, used as a beam quality index in photon beams.
p_{cav}	Factor to correct for the response of an ionisation chamber for deviations from Bragg-Gray conditions due to the effect of the air cavity in the medium.
p_{cel}	Corrects for the presence of the central electrode.
p_{dis}	Corrects for the difference in ionisation at the effective point of measurement and the depth at which the absorbed dose is stated.
p_{eff}	is the effective point of measurement of an ionisation chamber. For a radiation beam incident to the chamber from one direction the effective point of measurement is shifted from the position of chamber centre towards the source by a distance depending on the beam quality and the chamber type. For plane-parallel chambers the effective point of measurement is assumed at the inner surface of the entrance window.
p_{wall}	Corrects for the difference in composition between the ionisation chamber wall and water.

Q	Symbol to indicate the quality of a radiation beam.
Q_0	Symbol to indicate the reference beam quality used for calibration of an ionisation chamber (usually in a Standards Laboratory).
rdg	Reading of a dosimeter in arbitrary units.
$R_{50,dos}$	Half-value depth of dose in water in cm, used as the beam quality index for electron beams.
$R_{50,ion}$	Half-value depth of ionisation in water in cm.
R_p	Practical range in cm for electron beams.
r_{cyl}	Cavity radius of a cylindrical ionisation chamber.
sd	Standard deviation.
SCD	Source-chamber distance.
SDD	Source-detector distance.
SSD	Source-surface distance.
$s_{m,air}$	Stopping-power ratio medium to air, defined as the ratio of the mean restricted mass stopping powers of materials m and air, averaged over an electron spectrum.
$TPR_{20,10}$	Tissue-phantom ratio in water at depths of 20 and 10 cm, for a field size of 10 cm x 10 cm and a SCD of 100 cm, used as the beam quality index for high energy photon radiation.
u_i	Standard uncertainty of a quantity.

u_c	Combined standard uncertainty of a quantity.
U	Polarising voltage applied to the ionisation chamber.
W_{air}	The mean energy expended in air per ion pair formed.
z_{max}	Depth of maximum dose in (cm).
z_{ref}	Reference depth in (cm) for in-phantom measurements.

1. Introduction

1.1. General

Until recently, it has been generally recommended to perform reference dosimetry of clinical high energy x-ray and electron beams with ionisation chambers calibrated in terms of air kerma. To this end, the NCS has published NCS report 2 [1] and NCS report 5 [2] containing Codes of Practice for the dosimetry of high energy photon beams and electron beams, respectively, based on air kerma standards. These CoP's, issued by the NCS, were based on a number of simple concepts. A strictly limited number of ionisation chambers, routinely used in the Netherlands and Belgium were selected as reference dosimeters. In addition, the concept of a single conversion factor $C_{w,u}$ or $C_{w,e}$ for each reference chamber, as function of the beam quality index (i.e. a tissue phantom ratio, $TPR_{20,10}$, for photon beams and the mean energy at the phantom surface, \bar{E}_0 , for electron beams) was applied. These recommendations were compared by Mijnheer and Wittkämper [3] with similar international recommendations of the AAPM TG-21 [4] and IAEA TRS-277 [5]. The recommendations of the NCS are applied in every radiotherapy institute in the Netherlands and Belgium since 1986. Their implementation was audited on a regular basis [6,7,11] in most of the radiotherapy centres of both countries.

In the last two decades, reference dosimetry based on absorbed dose calibration coefficients has gained much attention. Advances in radiation dosimetry concepts and the development of absorbed dose to water standards by Primary Standard Laboratories have reduced the uncertainty in absorbed dose determination in external beam therapy. New codes of practice for reference dosimetry in clinical high energy photon and electron beams, based on absorbed dose to water standards, have been published recently. They are to replace the air kerma based codes of practice applied for the past twenty years.

Task Group 51 of the AAPM developed a protocol for dosimetry of high energy photon and electron beams [9]. The IAEA published a protocol for dosimetry of all external radiotherapy beams except neutron beams: TRS-398 [10].

Since the publication of these protocols a number of studies have already been devoted to comparing the two dosimetry formalisms with each other, or with air kerma based formalisms [12-15]. In general they discuss results for a limited number of chambers and often for one or two chamber types in each study. In a recent work, Palmans et al.[16] compared within an experimental study the absorbed dose to water based dosimetry versus air kerma

based dosimetry for high energy photon beams. A set of nine ionisation chambers was calibrated both in terms of air kerma and in terms of absorbed dose to water in ^{60}Co . Graphite-walled ionisation chamber types that are recommended for reference dosimetry in recent formalisms were selected. Their objective was to determine what consistency could be expected using the new formalisms and what differences could be expected compared to air kerma based formalisms (i.e. NCS-2). In another study [17] absorbed dose to water based dosimetry was compared with air kerma based dosimetry (NCS-5) in high energy electron beams. These measurements should be seen as preparatory for the present NCS report.

The NCS subcommittee on “Uniformity Dosimetry Protocols” has written a CoP based on the absorbed dose to water concept covering reference dosimetry in clinical high energy photon beams with nominal acceleration potentials between 1 and 25 MV and high energy electron beams with nominal energies between 4 and 25 MeV. The present CoP introduces a single beam quality correction factor taking into account all effects dependent of the radiation beam quality. For photon beams the beam quality correction factors are based on experimental data, partly measured in selected clinical accelerator beams in Belgium and the Netherlands, using a portable water calorimeter. For electron beams theoretically calculated correction factors are given, based on recent literature data. Furthermore, the CoP recommends a limited number of ionisation chambers for reference dosimetry in radiotherapy and uses updated information compared to the codes presented in TRS-398 and TG-51.

This CoP will replace the codes described in NCS reports 2 and 5. The guidelines and recommendations for clinical reference dosimetry in photon and electron beams are discussed in chapters 2 and 3 of this document.

1.2. Concept k_Q

Hohlfeld [18] proposed a practical concept for high energy photon dosimetry in Germany. This concept was based on one absorbed dose to water calibration coefficient in a ^{60}Co beam and a beam quality correction factor k_{Q,Q_0} , accounting for the high energy photon beam quality Q and has further been developed by Andreo [19] and Rogers [20]. The beam quality correction factor is defined as the ratio of absorbed dose to water calibration coefficients in the clinical beam quality Q , $N_{D,w,Q}$ and the reference (calibration) beam quality Q_0 , N_{D,w,Q_0} :

$$k_{Q,Q_0} = \frac{N_{D,w,Q}}{N_{D,w,Q_0}} \quad (1)$$

The reference beam quality Q_0 , used for calibration of ionisation chambers, is ^{60}Co gamma radiation. The beam quality correction factor has been measured directly for selected reference ionisation chambers in various clinical high energy photon beams. The measurements involved a portable water calorimeter developed by NMI to determine the absorbed dose to water in the clinical photon beam. The results of this experimental work are described in appendix A.4. These measurements together with other experimental data from the literature form the basis of the present CoP for high energy photon beams of the NCS. For photon beams these data are presented as a function of $\text{TPR}_{20,10}$. For electron beams, theoretically calculated correction factors, based on recent literature data, are given as a function of the half-value depth in water $R_{50,\text{dos}}$.

1.3. Basic Formalism

The absorbed dose to water at the reference depth in water for a user beam Q in absence of the ionisation chamber is given by:

$$D_{w,Q} = M_{\text{corr},Q} N_{D,w,Q_0} k_{Q,Q_0} \quad (2)$$

where:

- $M_{\text{corr},Q}$ reading of the electrometer corrected to ambient reference conditions and for the effects of recombination, polarity and the influence of the electrometer,
- N_{D,w,Q_0} the calibration coefficient for absorbed dose to water for a reference photon beam quality Q_0 ,
- k_{Q,Q_0} accounts for the effects of the differences between the beam quality Q and the reference beam quality Q_0 .

The k_{Q,Q_0} factors listed for high energy photon and electron beams are all related to the reference beam quality of ^{60}Co gamma radiation. Therefore the symbol Q_0 in k_{Q,Q_0} will be omitted in the following sections, yielding k_Q .

1.4. Available calibration services at LSDG and NMi

LSDG and NMi operate calibration services for absorbed dose to water in ^{60}Co gamma radiation. The primary standards involved are based on sealed water calorimeters according to a design of Domen [21]. The dose response of the calorimeter in a water phantom is transferred to a set of transfer ionisation chambers. LSDG uses three NE 2571 chambers as transfer instruments, whereas NMi uses three NE 2611A chambers for the determination of the long-term stability of the absorbed dose to water value. Secondary (local) standards including plane-parallel ionisation chambers from hospitals or other users are directly calibrated against the primary standards for absorbed dose to water in both laboratories, ensuring a short traceability chain. Further details on the construction and performance characteristics of the LSDG and NMi water calorimeters are given in Appendix A.3. It should be noted that the ionisation chambers employed as secondary (local) standard should fulfil the requirements outlined in this CoP.

1.5. Equipment

1. This CoP is based on the use of ionisation chambers for reference dosimetry in clinical photon and electron beams. A typical dosimeter system consists of an ionisation chamber, an electrometer and a radioactive check source. The ionisation chambers should preferably be designed for absorbed dose measurements in water and the construction should be as homogeneous and water equivalent as possible. Cylindrical ionisation chambers may be used for measurements in ^{60}Co gamma radiation, photon beams and electron beams with energy above approximately 10 MeV. The chamber should have a volume between about 0.1 and 1 cm³. The air cavity of the chamber should not be sealed, but designed to reach rapidly equilibrium conditions with the ambient temperature and air pressure.
2. Only graphite-walled ionisation chambers are recommended for reference dosimetry in this CoP. In general, these chambers have a more uniform response and better long-term stability than plastic walled chambers. However, some plastic-walled chambers have been demonstrated to show a stable response over time. Plastic-walled chambers are more robust and therefore often preferred as field instruments to perform routine measurements. The characteristics and physical data of the graphite-walled ionisation chambers most commonly

employed as local standards for reference dosimetry and recommended as reference instruments in this CoP are given in Appendix A.1.

3. Plane-parallel chambers are recommended for the use in electron beams. The use of plane-parallel chambers as reference instrument below 10 MeV is mandatory, because the uncertainty in the perturbation correction for a Farmer type cylindrical chamber becomes significant at low electron beam energies. Well-guarded plane-parallel chambers have a small perturbation effect, which will be further discussed in Appendix A.5. The chamber samples the electron fluence through the front window, so that the contribution of electrons entering the chamber through the side walls is minimized. The effective point of measurement, p_{eff} , is taken at the centre on the inner surface of the entrance window of the plane-parallel chamber for all electron beam energies and depths. Therefore, it is convenient to choose this as the reference point of the chamber during measurements. Further details and physical data for plane-parallel chambers recommended in this CoP can be found in Appendix A.5.
4. Water is recommended as phantom material, the use of plastic phantoms is not allowed. The phantom should be a full scatter phantom extending at least 5 cm outside the beam edges and at least 10 cm beyond the centre of the ionisation chamber along the beam axis. The dimensions of the water phantom should be at least 30 cm x 30 cm x 30 cm. The beam direction is vertical, perpendicular to the water surface. The water phantom must be open in this direction. These are the reference conditions for this CoP. The use of (water equivalent) plastics as phantom material [23, 24] is not allowed for reference dosimetry, because in general they give rise to the largest discrepancies in the determination of absorbed dose to water in clinical photon and electron beams [25].
5. For non-reference conditions, where the beam direction is horizontal, the window of the phantom, usually made of plastic, should have a thickness t_{win} between 2 and 5 mm. The water equivalent thickness of the phantom, expressed in g/cm^2 , has to be taken into account when the position of the chamber is determined in the phantom. The equivalent thickness (in g/cm^2) is calculated by multiplying t_{win} by the mass density ρ_{pl} of the plastic (in g/cm^3). For plastics commonly used, like PMMA and clear polystyrene, nominal mass densities of $\rho_{\text{PMMA}} = 1.19 \text{ (g/cm}^3\text{)}$ and $\rho_{\text{polystyrene}} = 1.06 \text{ (g/cm}^3\text{)}$ may be used to calculate the water equivalent thickness of the phantom window [22]. Note, that differences in the density of water equivalent plastics and polystyrene between manufacturers may exist [25].

6. If an ionisation chamber is not designed for direct use in a water phantom, it must be used with a close-fitting waterproof sleeve, made of low-Z materials (e.g. PMMA). The wall of the sleeve should be sufficiently thin (< 1.0 mm) to allow the chamber to achieve thermal equilibrium with the phantom in typically 2 to 3 minutes per degree of temperature difference. The sleeve should be vented to allow the air pressure to reach ambient air pressure rapidly; an air gap of 0.1 - 0.3 mm between the sleeve and the chamber is sufficient. Note that the waterproof sleeve should not be left in the water longer than necessary to perform the measurement, in order to minimize the build up of water vapour around the ionisation chamber. As the sleeve is to be considered as a component of the ionisation chamber, preferably the same sleeve should be used during calibration of the chamber and measurement assembly at LSDG or NMI.
7. The measuring assembly for the measurement of current (or charge) includes an electrometer with a good long-term stability (variation less than $\pm 0.5\%$ over one year) and a power supply for the polarizing voltage of the ionisation chamber. The electrometer and the ionisation chamber may be calibrated separately. This is particularly useful in centres that have several electrometers and/or chambers.
8. Sufficient time should be allowed for the dosimeter to reach thermal equilibrium. Some mains powered electrometers are best switched on for at least 2 h before use to allow stabilization. It is always advisable to pre-irradiate an ionisation chamber with 2–5 Gy to achieve charge equilibrium in the different materials. It is especially important to operate the measuring system under stable conditions whenever the polarity or polarizing voltage are modified which, depending on the chamber and sometimes on the polarity, might require several (up to 20) minutes.
9. The leakage current should always be measured before and after irradiation, and should be small compared with the current obtained during the irradiation (less than approximately 0.1% of the measurement current and normally of the same sign). Chambers with a leakage current which is large (approximately larger than 1% of the measurement current) or variable in time should not be used.
10. Only ionometric measurements are considered in this Code of Practice for reference dosimetry. We describe the calibration of field instruments (ionisation chamber and electrometer combination). The same approach can be used when various ionisation and electrometer combinations are used in clinical practice. It is left to the user to make a protocol designed for the users practice.

11. For relative measurements with ionisation chambers the displacement of the effective point of measurement has to be corrected for. For photons this is included in the calibration coefficient in the reference situation.
12. The use of other dosimetry detectors is not within the scope of this report, although for relative measurements other detectors may be more suitable. Care must be taken to use other detectors in relative dosimetry and to apply proper correction factors in non-reference situations.
13. Although not allowed for use in reference dosimetry, plastic phantoms can be used for routine quality assurance measurements, provided the relationship between dosimeter readings in plastic and water has been established for the user beam at the time of calibration. This will involve a careful comparison with measurements in water, which should be performed prior to the routine use of the phantom, and periodic checks at reasonable intervals might be also needed to assure the validity and consistency of the original comparison result.

2. Code of Practice for high energy photon beams

2.1. General conditions

The present CoP for high energy photon beams of the NCS will follow the concept below:

1. The CoP is based on recommendations for a limited number of graphite-walled, cylindrical ionisation chambers. These are the same models as those used in the water calorimetry campaign and are listed in Table 1 (see also Appendix A.1). The recommendations regarding measurement equipment given in section 1.5 should be followed.
2. The CoP will only recommend formalisms for high energy photon beams based on a ^{60}Co calibration. The symbol used for the beam quality correction factor will be k_Q .
3. The beam quality correction factors k_Q are based on the methods described in Appendix A.4.
4. The values of k_Q obtained from the fit in Appendix A.4 are valid for $\text{TPR}_{20,10}$ values ranging from 0.62 to 0.82; this corresponds to a nominal photon energy range from about 6 MV to 25 MV. The proposed fit to the k_Q data can also be applied to lower energies without loss of accuracy. However at higher energies extrapolation of the data will increase the uncertainty of k_Q .

2.2. Beam Quality

For high energy photons produced by clinical accelerators the beam quality Q is specified by the tissue-phantom ratio, $\text{TPR}_{20,10}$ [26-29]. This parameter is defined as the ratio of the absorbed doses at depths of 20 cm and 10 cm* in a water phantom, measured with a constant source-chamber distance[†] of 100 cm and a field size of 10 cm x 10 cm at the plane of the chamber as can be seen from Figure 1. The choice of $\text{TPR}_{20,10}$ is justified in Appendix A.4 where an analysis is made of various k_Q factors as function of $\text{TPR}_{20,10}$ and $\%dd(10)_x$ [9]. The experimental set up for measuring $\text{TPR}_{20,10}$ is shown in Figure 1. This parameter is obtained as the ratio of the absorbed doses at depths of 20 cm and 10 cm under following reference conditions:

* For simplicity the depth in water is expressed in cm instead of g/cm^2 , this will lead to differences in the evaluation of $\text{TPR}_{20,10}$ amounting to 0.1% at maximum.

[†] $\text{TPR}_{20,10}$ is normally measured at a fixed SCD (source-chamber distance). This quantity is however independent of the distance of measurement between $\text{SCD} = 80$ cm and $\text{SCD} = 150$ cm for a field size of 10 cm x 10 cm at the detector plane [1].

- water phantom,
- constant SCD (source-chamber distance) of 100 cm,
- field size of 10 cm x 10 cm at the axis of the chamber.

The recommendations regarding ionisation chambers, phantoms and sleeves, given in Section 1.5 should be followed. Although the definition of $\text{TPR}_{20,10}$ is strictly made in terms of absorbed dose ratios, the use of ionisation ratios provides an acceptable accuracy due to the slow variation with depth of water/air stopping-power ratios and the assumed constancy of perturbation factors beyond the depth of dose maximum. Small variations of the recombination correction factor k_s , as defined in Appendix A.2, may appear at different depths and consequently this variation should be investigated and taken into account if there is a variation with depth.

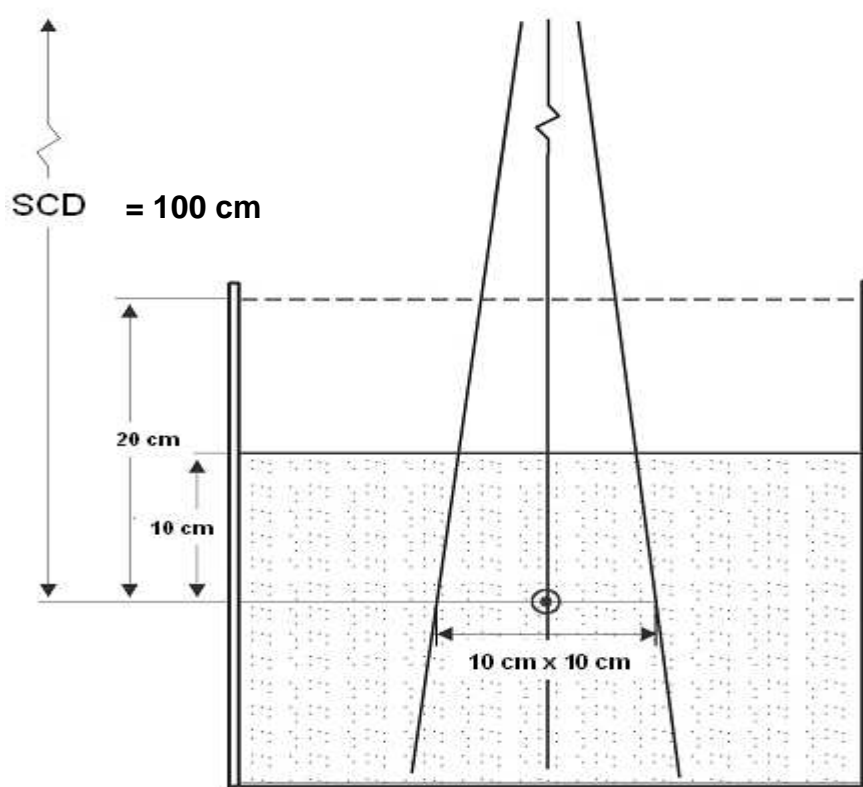


Figure 1. Experimental set-up for the determination of the beam quality index $\text{TPR}_{20,10}$. The source-to-chamber distance (SCD) is kept constant at 100 cm and measurements are made with 10 cm and 20 cm of water over the chamber. The field size at the position of the centre of the chamber, which is placed at the isocentre of the machine, is 10 cm x 10 cm.

2.3. Determination of $D_{w,Q}$ under reference conditions.

This section is based upon an absorbed dose to water calibration coefficient $N_{D,w}$ for a dosimeter in a reference ^{60}Co beam. The formalism to derive the absorbed dose to water for photon beams is described in section 1.3. Detailed information on the uncertainties, calculation of the influence quantities and numerical data needed for k_Q calculation can be found in the Appendices A.1-A.7. Moreover, these appendices will list the quantities, symbols and definitions, a summary of water calorimetry methodology and expected differences with NCS-2.

The recommendations are stated below:

1. The local standard shall be a graphite-walled, cylindrical ionisation chamber, listed in Table 1.
2. The local standard shall be calibrated in terms of absorbed dose to water, $N_{D,w}$, at a Primary Standard Dosimetry Laboratory (PSDL)* in a ^{60}Co beam under the conditions described in section 1.4.
3. A water phantom shall be used with a vertical beam; the chamber shall be fitted in the water with a thin waterproofing sheath† made of PMMA, not thicker than 1 mm. More information on phantoms and sleeves is given in section 1.5.
4. For the determination of the absorbed dose to water in the users beam, the chamber shall be placed on the beam axis with the geometrical centre of the chamber at the reference depth $z_{\text{ref}} = 10$ cm.
5. The water surface shall be positioned at the isocentre of the linear accelerator for a fixed source-surface distance (SSD = 100 cm) measurement or the detector shall be positioned at the isocentre of the linear accelerator for a fixed source-chamber distance (SCD = 100 cm) measurement. A field size of 10 cm x 10 cm at the isocentre shall be used.
6. For the local standard the absorbed dose to water $D_{w,Q}$ at the reference depth will be given by:

$$D_{w,Q} = M_{\text{corr},Q} N_{D,w} k_Q \quad (3)$$

where:

*For the Netherlands the Primary Standard Dosimetry Laboratory (PSDL) is the NMI at Delft whereas for Belgium the Primary Standard Dosimetry Laboratory is the LSDG at Ghent.

† All k_Q values including those for the waterproof Wellhöfer chamber have been determined using a waterproof sheath (≤ 1 mm). The user employing the Wellhöfer chamber without the waterproof sheath has to account for the difference in the dose determination, which is expected to be less than 0.2%.

$D_{w,Q}$ is the absorbed dose to water in the users beam Q and in absence of the chamber, at the reference depth $z_{ref} = 10$ cm,

$M_{corr,Q}$ is the electrometer reading, corrected for influence quantities as described in Appendix A.2,

$N_{D,w}$ is the absorbed dose to water calibration coefficient in the ^{60}Co reference beam quality, given by the Standards Laboratory at reference ambient conditions (normally $T_{ref} = 20^\circ\text{C}$ and $P_{ref} = 101.325$ kPa and 50% relative humidity),

k_Q is the beam quality correction factor that depends on the chamber type and radiation beam quality of the user beam. For the 4 recommended ionisation chamber types of Table 1 k_Q values as function of the quality index $\text{TPR}_{20,10}$ are obtained from equation:

$$k_Q = 0.80 + 0.20 \cdot \frac{1 + e^{C \cdot (0.57 - X_0)}}{1 + e^{C \cdot (\text{TPR}_{20,10} - X_0)}} \quad (4)$$

where the parameters C and X_0 of this equation for the recommended ionisation chambers are given in Table 1. The k_Q are valid in the photon energy range of 6 MV to 25 MV. The data obtained using equation (4) are plotted in Figure 10 (see Appendix A.4, section.A.4.5.2).

7. k_Q equals 1.000 for for determination of the absorbed dose to water in a ^{60}Co beam.
8. The electrometer reading M_Q should be corrected for all influence quantities:

$$M_{corr,Q} = M_Q k_{TP} k_h k_{pol} k_s \quad (5)$$

where:

k_{TP} is the correction factor for pressure and temperature in the air cavity at the time of measurement,

k_h is the correction for humidity in the air cavity at the time of measurement,

k_{pol} is the correction on the chamber reading due to the applied polarity of the polarizing voltage,
 k_s is the correction for incomplete charge collection in the ionisation chamber due to recombination.

These correction factors have to be determined according to the procedures explained in Appendix A.2.

9. Under these conditions the relative standard uncertainty of the absorbed dose to water $D_{w,Q}$ at the reference depth $z_{ref} = 10\text{ cm}$ for a high energy photon beam is estimated to be 1.0% (1sd). More details on the estimation of this relative combined standard uncertainty are given in Appendix A.4.

Table 1: List of the recommended graphite-walled cylindrical chambers together with the parameters of the sigmoid fit (see Appendix A.4). Using these parameters in equation 4 allows obtaining the k_Q values as a function of the beam quality $TPR_{20,10}$ for these ionisation chambers.

Chamber type	X_0	C
NE 2561/NE 2611	0.8971	15.15
NE2571 PTW30012 Wellhöfer FC65G	0.9198	11.67

2.4. Determination of $D_{w,Q}$ under non-reference conditions

Determination of $D_{w,Q}$ under non-reference conditions is not part of the CoP, however a few points should be considered. Field instruments shall be calibrated against the reference instrument at the radiation qualities at which they are to be used. This calibration shall be in terms of absorbed dose to water.

1. Only ionisation chambers are considered in this protocol.
2. Field ionisation chambers shall be calibrated in terms of absorbed dose to water against the local standard, with their central axis at the same reference depth z_{ref} , SSD and field size as stated in section 2.3. A water phantom shall be used. This calibration has to be done for each beam quality at which they are to be used. The absorbed dose to water calibration coefficient $N_{w,Q}^{field}$ at the users beam quality Q for the field ionisation chamber is then given by:

$$N_{w,Q}^{field} = D_{w,Q}^{std} / M_{corr,Q}^{field} \quad (6)$$

where:

$D_{w,Q}^{std}$ is the absorbed dose to water in the users beam at the reference depth $z_{ref} = 10$ cm determined by the local standard,
 $M_{corr,Q}^{field}$ is the electrometer reading of the field instrument, corrected for influence quantities as described in Appendix A.2.

Preferably an additional monitor chamber in the radiation field should be used to normalise the reading in order to account for beam fluctuations.

3. If the field instrument is one of the recommended ionisation chambers from Table 1, than this calibration can be considered as a cross calibration. The obtained calibration coefficient $N_{w,Q}^{field}$ is applicable for all photon beams by calculating a “ ^{60}Co calibration coefficient” for the field instrument from it. The ^{60}Co related calibration coefficient can be obtained from the ratio $N_{w,Q}^{field}$ and k_Q . This calibration coefficient can subsequently be used according to the CoP.
4. Relative measurements (i.e. determination of the absorbed dose at other points in the phantom than z_{ref}) can be performed by different types of ionisation chambers. To this end, the effect of displacement, which implicitly is already taken into account in k_Q (for photons) for dosimetry under reference conditions, should be

taken into account by positioning the effective point of measurement of the ionisation chamber to the depth of measurement. This effective point of measurement is 0.6 times the inner radius in front of the geometrical centre for cylindrical ionisation chambers [5] and on the inner site of the front window for plane parallel ionisation chambers [30].

5. The use of other dosimetry detectors (i.e. diodes, diamond detectors, TLD, Alanine, scintillator detectors, etc.) is not within the scope of this report.

3. Code of practice for high energy electron beams.

3.1. General conditions

The present CoP for high energy electron beams of the NCS will follow the concept below:

1. The CoP is based on recommendations for a limited number of graphite-walled, Farmer type and plane-parallel ionisation chambers. The Farmer type cylindrical chamber types are the same models as used in high energy photon beams and recommended only for electron energies with beam quality $R_{50,dos}$ larger than 4 cm* in water. For lower electron energies only plane-parallel chambers are recommended. The recommendations regarding measurement equipment given in section 1.5 should be followed.
2. For dosimetry in a high energy electron beam using a Farmer type cylindrical ionisation chamber (only for $R_{50,dos} > 4$ cm) the recommendation is based on a calibration in a ^{60}Co reference beam. Dosimetry using a plane-parallel ionisation chamber (for any beam quality Q) is strongly recommended to be based on a cross-calibration against a Farmer type cylindrical ionisation chamber in a high energy electron beam. This beam is referred to as the cross calibration beam quality with an $R_{50,dos}$ recommended to be larger than 7 cm. This method requires a beam quality correction factor $k_{Q,Qcross}$, which converts the calibration coefficient of the plane-parallel ionisation chamber in the cross-calibration beam quality to the beam quality Q . Only for the case this option is not available to a particular user, the protocol also provides the data necessary to perform dosimetry using a plane-parallel ionisation chamber based on a dose to water calibration in a ^{60}Co reference beam.
3. The beam quality correction factors k_Q and $k_{Q,Qcross}$ are based on the methods described in appendix A.5 and are given as a parameterised function of the beam quality index $R_{50,dos}$. These factors are based on calculations including up to date information from the literature.
4. Values of k_Q and $k_{Q,Qcross}$ are provided for $R_{50,dos}$ values ranging from 1 cm to 12 cm (although their validity extends to 20 cm).

* For simplicity $R_{50,dos}$ and the depth z in water is expressed in cm instead of g/cm², this leads to differences in the evaluation of $R_{50,dos}$ of about 0.2%.

3.2. Beam Quality

For high energy electron beams produced by clinical accelerators, the beam quality Q is specified by the 50% dose level beyond the dose maximum, $R_{50,dos}$, as illustrated in Figure 2. The parameter $R_{50,dos}$ is obtained as the depth beyond the dose maximum where the absorbed dose to water is 50% of its maximum value. The beam quality index can be determined either directly from a depth dose curve or from a depth ionisation curve. Depth ionisation curves are preferably measured using a plane-parallel ionisation chamber, with the reference point of the chamber taken as the centre of the inner surface of its front window. If a cylindrical chamber would be used, the reference point has to be taken as 0.5 times the inner radius of the chamber in front of the geometrical centre^{*}, as illustrated in Figure 3.

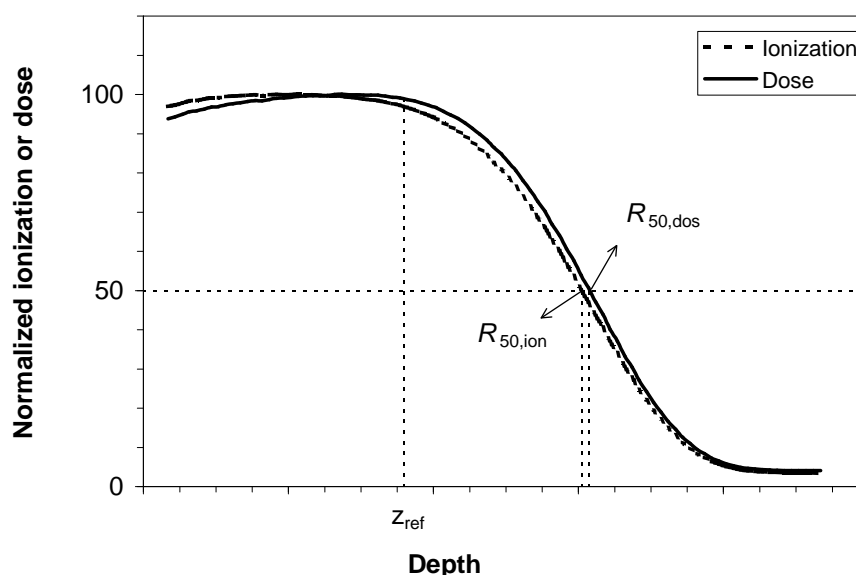


Figure 2: Definition of the beam quality index $R_{50,dos}$ for electron beams by means of an example. $R_{50,ion}$ and the reference depth z_{ref} are also indicated. Note that z_{ref} in this case does not coincide with the depth of dose maximum.

^{*} Note this means that the geometrical centre of the ionization chamber is shifted away from the beam source over a distance of 0.5 times the inner radius of the chamber's cavity.

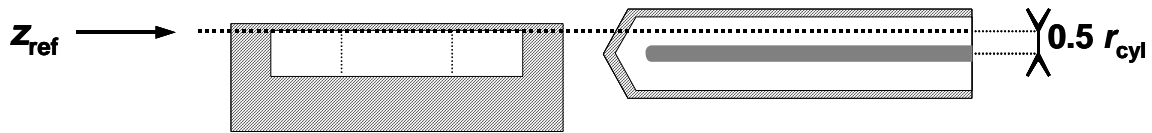


Figure 3: Plane-parallel and cylindrical ionisation chamber each at reference depth z_{ref} .

The parameter $R_{50,\text{dos}}$ is measured under the following reference conditions:

- water phantom,
- constant SSD (source-surface distance) of 100 cm,
- field size at the phantom surface of at least 10 cm × 10 cm or 20 cm × 20 cm if $R_{50,\text{dos}}$ is larger than 7 cm.

The recommendations regarding ionisation chambers, phantoms and sleeves, given in Section 1.5 should be followed. When departing from an ionisation curve, $R_{50,\text{dos}}$ is derived from $R_{50,\text{ion}}$, the 50% ionisation level beyond the ionisation maximum, using the following generic expressions [32]:

$$R_{50,\text{dos}} = 1.029 \cdot R_{50,\text{ion}} - 0.06 \text{ cm} \quad (R_{50,\text{ion}} \leq 10 \text{ cm}) \quad (7)$$

$$R_{50,\text{dos}} = 1.059 \cdot R_{50,\text{ion}} - 0.37 \text{ cm} \quad (R_{50,\text{ion}} > 10 \text{ cm}) \quad (8)$$

For some plane-parallel ionisation chambers it has been reported that recombination and polarity effects vary as a function of depth. It can be concluded that these variations, as defined in Appendix A.5, can appear at different depths and consequently this variation should be investigated and taken into account if there is a variation with depth.

3.3. Determination of $D_{w,Q}$ under reference conditions.

The formalism to derive the absorbed dose to water for electron beams is described in section 1.3. Detailed information on the uncertainties, calculation of the influence quantities and numerical data needed for k_Q or $k_{Q,Q_{\text{cross}}}$ calculation can be found in the Appendices. Moreover, Appendices A.1-A.7 list the quantities, symbols and definitions and expected differences with NCS report 5.

The recommendations are stated below:

1. The local standard shall be either a Farmer type graphite-walled cylindrical ionisation chamber of one of the types listed in Table 1, or a plane-parallel ionisation chamber of type NACP02, PTW 34001 or Wellhöfer PPC-40. Note that plane-parallel chambers must be used for beam qualities $R_{50,\text{dos}} < 4 \text{ cm}$.

2. The Farmer type chamber used as a local standard shall be calibrated in terms of absorbed dose to water, $N_{D,w}$, at a Primary Standard Dosimetry Laboratory (PSDL)^{*} in a ^{60}Co beam under the conditions described in section 1.4. The plane-parallel chamber types used as a local standard shall be preferably cross calibrated against a Farmer type chamber in an electron beam with a beam quality index $R_{50,dos}$ greater than 7 cm. Only for the case this option is not available to a particular user, the protocol also provides the data necessary to perform dosimetry using a plane-parallel ionisation chamber based on a calibration in a ^{60}Co reference beam.
3. A water phantom shall be used with a vertical beam; the non-waterproof chamber shall be fitted in water with a thin waterproofing sheath made of PMMA, not thicker than 1 mm. More information on phantoms and sleeves is given in section 1.5.
4. For the determination of the absorbed dose to water in the users beam, the chamber shall be placed on the beam axis with the reference point of the chamber at the reference depth [31]:

$$z_{ref} = 0.6R_{50,dos} - 0.1\text{cm} \quad (9)$$

For a cylindrical chamber the reference point has to be taken as 0.5 times the inner radius of the chamber in front of the geometrical centre, whereas for a plane-parallel the reference point of the chamber is taken as the centre of the inner surface of its front window.

5. The water surface shall be positioned at a fixed source-surface distance of SSD = 100 cm. The field size at the phantom surface should be either 10 cm x 10 cm or the field size used for normalization of output factors, whichever is larger[†].
6. If the local standard is a Farmer type cylindrical ionisation chamber the absorbed dose to water $D_{w,Q}$ at the reference depth will be given by :

$$D_{w,Q} = M_{corr,Q} N_{D,w} k_Q \quad (10)$$

where:

$D_{w,Q}$ is the absorbed dose to water in the users beam Q at the reference depth z_{ref} ,

^{*}For the Netherlands the Primary Standard Dosimetry Laboratory (PSDL) is the NMI at Delft whereas for Belgium the Primary Standard Dosimetry Laboratory is the LSDG at Ghent.

[†] Note that this is different than the recommendation for the determination of the beam quality index $R_{50,dos}$.

- $M_{corr,Q}$ is the electrometer reading, corrected for influence quantities as described in Appendix A.2,
- $N_{D,w}$ is the absorbed dose to water calibration factor in the ^{60}Co reference beam quality, given by the Standards Laboratory at reference ambient conditions (normally $T_{ref} = 20^\circ\text{C}$ and $P_{ref} = 101.325 \text{ kPa}$ and 50% relative humidity),
- k_Q is the beam quality correction factor that depends on the chamber type and radiation beam quality of the users beam. Recommended k_Q values as function of the quality index $R_{50,dos}$ and for the three recommended ionisation chamber types are obtained from the following equation:

$$k_Q = A - B \cdot [R_{50,dos}(Q)]^C \quad (11)$$

The parameters A, B and C for this equation are given in Table 2. The k_Q for the Farmer type chambers are valid in the beam quality range from 4 cm to 12 cm. The data obtained using equation (11) are plotted in Figure 12 (see Appendix A.5 section A.5.2).

7. If the local standard is a plane-parallel ionisation chamber it is recommended that a calibration coefficient $N_{D,w,Q_{cross}}^{pp}$ is used, which is obtained in a high energy electron beam Q_{cross} as explained below. Then the absorbed dose to water $D_{w,Q}$ at the reference depth will be given by:

$$D_{w,Q} = M_{corr,Q}^{pp} \cdot N_{D,w,Q_{cross}}^{pp} \cdot k_{Q,Q_{cross}} \quad (12)$$

where:

- $D_{w,Q}$ is the absorbed dose to water in the users beam Q at the reference depth z_{ref} ,
- $M_{corr,Q}^{pp}$ is the electrometer reading, corrected for influence quantities as described in Appendix A.2,
- $N_{D,w,Q_{cross}}^{pp}$ is the absorbed dose to water calibration coefficient in the cross calibration beam quality Q_{cross} at reference ambient conditions

(normally $T_{\text{ref}} = 20^\circ\text{C}$ and $P_{\text{ref}} = 101.325 \text{ kPa}$ and 50% relative humidity). $N_{D,w,Q_{\text{cross}}}^{\text{pp}}$ is obtained from calibrating the plane-parallel chamber (pp) in the beam quality Q_{cross} against the dose measured by a Farmer type cylindrical ionisation chamber:

$$N_{D,w,Q_{\text{cross}}}^{\text{pp}} = \frac{M_{\text{corr},Q_{\text{cross}}}^{\text{cyl}}}{M_{\text{corr},Q_{\text{cross}}}^{\text{pp}}} \cdot N_{D,w}^{\text{cyl}} \cdot k_{Q_{\text{cross}}}^{\text{cyl}} \quad (13)$$

$k_{Q,Q_{\text{cross}}}$ is the beam quality correction factor that depends on the chamber type and radiation beam quality of both the cross calibration beam quality Q_{cross} and the users beam quality Q . Recommended $k_{Q,Q_{\text{cross}}}$ values as function of the quality index $R_{50,\text{dos}}$ and for the recommended ionisation chamber types are obtained from the following equation:

$$k_{Q,Q_{\text{cross}}} = \frac{A - B \cdot [R_{50,\text{dos}}(Q)]^C}{A - B \cdot [R_{50,\text{dos}}(Q_{\text{cross}})]^C} \quad (14)$$

The parameters for this equation are given in Table 2. The $k_{Q,Q_{\text{cross}}}$ are valid in the beam quality range from 1 cm to 12 cm. The data obtained using equation (14) are plotted in figure 13 (see Appendix A.5, section A.5.2).

8. If the local standard is a plane-parallel ionisation chamber, calibrated in terms of absorbed dose to water in a ^{60}Co reference beam: the same procedure as outlined in point 6 should be followed. Note, that in this case the depth has to be corrected for non-water equivalence of the entrance window of the plane-parallel chamber. To this end, the depth at which the outside front face of the plane-parallel ionisation chamber should be positioned equals the reference depth minus the water equivalent thickness of the entrance window as given in Table 3 (see Appendix A.1).
9. The electrometer reading M_Q should be corrected for all influence quantities:

$$M_{\text{corr},Q} = M_Q k_{\text{TP}} k_h k_{\text{pol}} k_s \quad (15)$$

where:

- k_{TP} is the correction factor for pressure and temperature in the air cavity at the time of measurement,
- k_h is the correction for humidity in the air cavity at the time of measurement,
- k_{pol} is the correction on the chamber reading due to the applied polarity of the polarizing voltage,
- k_s is the correction for incomplete charge collection in the ionisation chamber due to recombination.

Determination of these correction factors is explained in Appendix A.2.

10. Under these conditions the relative standard uncertainty of the absorbed dose to water $D_{w,Q}$ at the reference depth z_{ref} for a high energy electron beam is estimated to be 1.5% (1sd) when it concerns a measurement with a Farmer type chamber in an electron beam with $R_{50,dos} > 7$ cm or 1.8% (1sd) when it concerns a measurement with a cross calibrated plane-parallel chamber in any electron beam. More details on this uncertainty estimate are given in Appendix A.5.

Table 2: Parameters for the model of equations (11) and (14) to obtain k_Q and $k_{Q,Qcross}$ values as a function of the beam quality $R_{50,dos}$ for the recommended types of ionisation chambers (see Appendix A.5).

	A	B	C
Farmer types	0.9345	0.0057	0.7733
NACP02	1.1955	0.2274	0.1479
Roos types	1.1376	0.1700	0.1835

3.4. Determination of $D_{w,Q}$ under non-reference conditions

Determination of $D_{w,Q}$ under non-reference conditions is not part of the CoP, however a few points should be considered. Field instruments shall be calibrated against the reference instrument at the radiation qualities at which they are to be used. This calibration shall be in terms of absorbed dose to water.

1. Only ionisation chambers are considered in this protocol.
2. Field ionisation chambers shall be calibrated at the same reference depth z_{ref} , SSD and field size as stated in section 3.3. A water phantom shall be used. This calibration has to be done for each beam quality at which they are to be used.

The absorbed dose to water calibration coefficient $N_{w,Q}^{field}$ at the users beam quality Q for the field ionisation chamber is then given by:

$$N_{w,Q}^{field} = D_{w,Q}^{std} / M_{corr,Q}^{field} \quad (16)$$

where:

$D_{w,Q}^{std}$ is the absorbed dose to water in the users beam at the reference depth z_{ref} determined by the local standard,

$M_{corr,Q}^{field}$ is the electrometer reading of the field instrument, corrected for influence quantities as described in Appendix A.2.

Preferably an additional monitor chamber in the radiation field should be used to normalise the reading in order to account for beam fluctuations.

3. If the field instrument is one of the recommended Farmer type ionisation chambers, then this calibration can be considered as a cross calibration and the obtained calibration coefficient $N_{w,Q}^{field}$ is applicable for other electron beams with $R_{50,dos} > 4$ cm. To this end, a “ ^{60}Co calibration coefficient” for the field instrument can be back-calculated from the ratio $N_{w,Q}^{field}$ and k_Q . This calibration coefficient can subsequently be used according to the CoP. An additional uncertainty component should be taken into account.
4. Relative measurements (i.e. determination of the absorbed dose at other points in the phantom than z_{ref}) can be performed by the same ionisation chambers described above. The effective point of measurement of the ionisation chamber has to be positioned at the depth(s) of interest. Since the water to air mass stopping power ratios vary considerably as a function of electron energy the measured depth ionisation curve has to be converted into a depth dose curve by applying the water-to-air mass stopping power ratios given by the formula in section A 5.5. It has also been reported that for some ionisation chambers the recombination and polarity corrections may vary considerably as a function of depth. Consequently, these influence quantities (see Appendix A.2) should be measured as a function of depth and be corrected for if necessary.
5. The use of other dosimetry detectors (i.e. diodes, diamond detectors, TLD, Alanine, scintillator detectors, etc.) is not within the scope of this report.

Appendices

A.1 Instrument data for recommended ionisation chambers

In Table 3 the characteristics are given for the ionisation chambers recommended for reference dosimetry in this code of practice.

Table 3: Characteristics of ionisation chamber types recommended for reference dosimetry¹

¹ data taken from manufacturers

² Polymethyl Methacrylate (C₅H₈O₂), also known as acrylic. Trade names are Lucite, Perspex or Plexiglass.

Chamber type	Inner dimensions cavity			Wall material	Wall thickness	Central electrode	Waterproof
Cylindrical	nominal volume (cm ³)	length (mm)	radius (mm)	material (g/cm ²)	material		
NE 2611A	0.33	9.2	3.7	Graphite	0.090	Aluminium (hollow)	N
NE 2571, Farmer	0.69	24.1	3.15	Graphite	0.065	Aluminium	N
PTW 30012, Farmer	0.6	23.0	3.05	Graphite	0.079	Aluminium	N
Scanditronix-Wellhöfer FC65-G	0.65	23.0	3.1	Graphite	0.081	Aluminium	Y

Chamber type	Nominal volume	Materials	Window thickness water eq.		Electrode spacing	Collecting electrode diameter	Guard ring width	Waterproof
Plane-parallel	(cm ³)		(mg/cm ²)	(mm)	(mm)	(mm)	(mm)	
Scanditronix-Wellhöfer NACP02	0.16	Mylar foil and graphite window, graphited rexolite	104	0.6	2.0	10	3.0	Y
PTW 34001, Roos	0.35	PMMA ² graphited electrodes	118	1	2	16	4	Y
Scanditronix-Wellhöfer PPC-40, Roos	0.4	PMMA ²	118	1	2.0	16	4.0	Y

A.2 Influence parameters

Air density correction

The ionisation chambers recommended for reference dosimetry in this Code of Practice are open to ambient air. Consequently the mass of air in the cavity will vary with atmospheric conditions for temperature and pressure. The correction factor k_{TP} given by:

$$k_{TP} = \frac{(T + 273.15)}{(T_0 + 273.15)} \frac{P_0}{P} \quad (17)$$

accounts for this effect by converting the mass of air in the cavity to the reference conditions for temperature $T_0 = 20$ °C and pressure $P_0 = 101.325$ kPa.

Humidity

No correction is made for the relative humidity, if the ionisation chamber is used in a range of 20% to 80% relative humidity and has been calibrated at a reference condition for relative humidity of 50% [34]. It is recommended to use ionisation chambers in relative humidity conditions between 20 and 80%. In the unlikely case that the relative humidity is outside this range, a correction factor has to be applied.

Electrometer calibration coefficient k_{elec}

It is common practice in Belgium and the Netherlands to calibrate the ionisation chamber and the electrometer together, as one set. In this case the calibration coefficient $N_{D,w}$ is expressed in the unit Gy/rdg or Gy/C depending on the readout of the electrometer and no separate electrometer calibration coefficient has to be applied. If the ionisation chamber and electrometer are calibrated separately, the calibration coefficient $N_{D,w}$ for the ionisation chamber is given in units Gy/C. The calibration coefficient obtained for the electrometer converts the electrometer reading to charge and is expressed in unit rdg/C. If the reading of the electrometer is in terms of charge the electrometer calibration coefficient is dimensionless.

Polarity correction

The reading of an ionisation chamber is affected by the polarity of the chamber voltage. The polarity effect depends on the radiation quality, the (incident) beam energy, the magnitude of the polarizing voltage on the chamber, the field size and depth in the phantom, and the

design of the ionisation chamber. The effect may be considerable for plane-parallel ionisation chambers in electron beams, more in particular in low energy electron beams. The polarity effect for every reference chamber should always be measured at the reference point in the energy range where the chamber is to be used. If the polarity correction is more than a few percent the chamber is not recommended for reference dosimetry. The correction factor for polarity in a given radiation beam is defined by:

$$k_{pol} = \frac{|M^+| + |M^-|}{2M} \quad (18)$$

where the superscripts + and – in the nominator indicate the electrometer reading with a positive and negative polarizing voltage respectively. M is the electrometer reading taken at polarity routinely used during measurements. When the chamber is sent for calibration, the calibration certificate should clearly state the polarizing voltage and the polarity adopted during calibration and if or not a polarity correction has been applied by the calibration laboratory. If a chamber is used in a radiation beam with nearly the same quality as the reference beam in the calibration laboratory, and with the same magnitude and sign of the polarizing voltage, the polarity is the same and no correction is necessary. For most chamber types the correction is small in photon beams and is often neglected (The primary standards laboratories in Belgium and the Netherlands, LSDG and NMI, don't apply a polarity correction during calibration in their reference ^{60}Co beams).

When the calibration laboratory has applied a polarity correction during calibration at the reference beam quality (usually ^{60}Co) the user has to determine k_{pol} according to equation (18) for all measurements made using the routine polarity. When the calibration laboratory has not applied a polarity correction then equation (18) has to be modified to evaluate the polarity effect in a user beam with quality Q. The polarity correction for beam quality Q has to be determined relative to the polarity correction of the reference beam quality ^{60}Co using the expression:

$$k_{pol} = k_{pol,Q} / k_{pol,Co} \quad (19)$$

The correction $k_{pol,Co}$ has to be determined either by the user from known data of the chamber response to different beam qualities and polarities with a relative standard

uncertainty of less than 0.5% or by sending the chamber to a calibration laboratory in order to measure the polarity correction in a ^{60}Co beam.

Recombination correction

A correction is required to account for the lack of complete charge collection in the chamber volume, due to the recombination of ions. Several physical phenomena may contribute to recombination, but initial and volume or general recombination are commonly considered the most important effects [35]. Initial recombination refers to the forming of ion pairs along the track of a single ionising particle and therefore independent of the dose rate. Volume recombination is due to recombination of ions from different tracks and depends on the ion density and thus on the dose rate (continuous beams) or dose per accelerator pulse (pulsed beams). Both effects depend on the chamber geometry and on the polarizing voltage applied to the chamber.

For continuous radiation beams, i.e. ^{60}Co , the recombination effect is small ($< 0.2\%$) for the output rates encountered at irradiation facilities in calibration laboratories. Often initial recombination is neglected and only general recombination is considered assuming a linear relation between the inverse of the charge ($1/M$) and the inverse square of the polarizing voltage ($1/U^2$). However, when a proper correction is required initial recombination in continuous beams should not be neglected. In this case k_s can be approximated by using a method described by Boutillon [37]. This method evaluates the recombination correction including both the initial and general recombination component.

In pulsed radiation beams general recombination is the dominant effect. The recombination correction may amount to 1%-3% for a high dose rate beam produced by a medical linear accelerator and even higher for pulsed-scanned beams. An expression for the recombination factor can be derived using a theoretical approach developed by Boag [38, 39]. For pulsed beams the correction for volume recombination can be written as:

$$k_s = 1/f = u/\ln(1+u) \quad (20)$$

The dimensionless variable u is given as $u = \mu m d^2/U$, where m is the dose (charge density) per pulse, d is the electrode separation in the ionisation chamber, U the applied polarizing voltage and μ is a constant depending on the ion mobilities and recombination rate of the gas in the chamber volume. For cylindrical ionisation chambers the effective electrode separation can be calculated from expressions given by Boag [35]. However, the method does not account for chamber-to-chamber variations within a given chamber type. Under near

saturation conditions ($k_s < 1.05$) the recombination correction for pulsed and pulsed-scanned beams can be accurately determined using the two-voltage method as described by Boag and Curren [36]. The two-voltage technique involves the measurement of charge produced by a radiation beam in the cavity of an ionisation chamber when two different polarizing voltages are applied to the ionisation chamber. The method assumes a linear relation between $1/M$ and $1/U$. The recombination correction is determined from the values of charges or readings M_1 and M_2 , which are collected in the same irradiation conditions at the normal polarizing voltage U_1 and at a lower voltage U_2 respectively. The ratio (U_1/U_2) should have a value of at least two or three.

Weinhous and Meli [40] have derived numerical solutions for the evaluation of the recombination correction according to approach of Boag and Curren and have computed quadratic fits to these solutions as a function of the polarizing voltage ratio. The recombination correction factor k_s at the normal polarizing voltage U_1 is obtained from:

$$k_s = a_0 + a_1(M_1 / M_2) + a_2(M_1 / M_2)^2 \quad (21)$$

where M_1 and M_2 are the measured charge readings at polarizing voltages U_1 and U_2 , respectively. In Table 4 the coefficients a_i are given for pulsed beams as a function of voltage ratio (U_1/U_2). A voltage ratio equal to or larger than 3 is recommended. Note, that the polarity effect will change with the applied polarizing voltage and that M_1 and M_2 should be corrected for the change in this effect.

Table 4: Quadratic fit coefficients for pulsed radiation as a function of the voltage ratio U_1/U_2 . Data are taken from Weinhous and Meli [40].

(U_1/U_2)	a_0	a_1	a_2
2.0	2.337	-3.636	2.299
2.5	1.474	-1.587	1.114
3.0	1.198	-0.875	0.677
3.5	1.080	-0.542	0.463
4.0	1.022	-0.363	0.341
5.0	0.9745	-0.1875	0.2135
6.0	0.9584	-0.1075	0.1495
8.0	0.9502	-0.03732	0.08750
10.0	0.9516	-0.01041	0.05909

For values of $k_s < 1.03$ the recombination correction can be approximated to within 0.1% by the expression [41]:

$$k_s - 1 = \frac{M_1 / M_2 - 1}{U_1 / U_2 - 1} \quad (22)$$

Note that for a recombination correction < 1.03 equations (21) and (22) will also include the component for initial recombination, which is also proportional to $1/U$.

The calibration certificate issued by the standards laboratory should state whether or not a recombination correction has been applied. If a calibration laboratory has not applied a recombination correction, the correction factor for the user beam quality Q has to be derived relative to the appropriate correction factor for the reference beam quality ^{60}Co used in the calibration laboratory. The recombination correction is then obtained by:

$$k_s = k_{s,Q} / k_{s,\text{Co}} \quad (23)$$

For ^{60}Co the recombination correction will be relatively small compared to the correction for the user beam quality Q and is therefore neglected in most cases. Parallel-plate chambers used for electron beams having typically plate separations of 1 - 2 mm would be expected to have a low recombination effect, when employed in clinical beams. However, some recombination corrections measured using the two-voltage method are larger than expected. It has been demonstrated that for some plane-parallel chambers the linear relationship between $1/M$ and $1/U$ is not satisfied in the voltage region used for the two-voltage method [36, 40, 42-44]. It is recommended only to apply the two-voltage method to plane parallel chambers, if the response of the chamber as function of the polarizing voltage has been measured up to its maximum value specified by the manufacturer in order to establish a range of linearity. The chamber should be operated at voltages to remain within the linear range, ensuring that the use of the two-voltage method is valid.

A.3 Absorbed dose to water standards at NMI and LSDG

The Belgian and Dutch standard dosimetry laboratories provide absorbed dose to water calibration coefficients in ^{60}Co relying on water calorimeters. For high energy photon beams k_Q values, NCS decided to base these on experimental data. To this end experimental k_Q 's were measured in nine clinical photon beams using the NMI portable water calorimeter.

This section describes the water calorimeters of both laboratories. The experimental determination of k_Q values with the NMI water calorimeter and the analysis of the experimental k_Q values in the context of this NCS protocol are described in Appendix A.4.

A.3.1 The NMI water calorimeter

The design of the water calorimeter is based on the sealed-water calorimeter of Domen [21]. The water calorimeter consists of a water tank surrounding a high-purity water cell, which contains two thermistor probes. The high-purity water cell is a sealed, thin-walled glass vessel enclosing a volume of water. The temperature rise due to irradiation by high energy photons is measured inside this volume of water. The quality of the water is carefully controlled by purifying and by saturating the water with selected gasses, before filling the glass vessel. The water calorimeter is compact and transportable, weighing 60 kg when empty, and having outer dimensions of only 60 cm x 60 cm x 70 cm. The dimensions of the water phantom are standard 30 cm x 30 cm x 30 cm. A small spacing between the inner and outer polystyrene foam insulation boxes encloses a copper heat exchange system. The six copper walls are connected in a parallel manner, which helps to reduce temperature gradients inside the water phantom. Cooling is performed by a computer-controlled water-cooling thermostat, which uses two PT100 thermometers (one mounted on one of the copper walls and one mounted inside the water phantom) to control and monitor the calorimeter temperature. A built-in magnetic stirrer enables a reduction of temperature drifts due to conduction in between irradiation runs. A cold finger placed inside the water phantom can be switched into the cooling circuit to reduce the time needed to cool down from room temperature to 4 °C.

The water calorimeter can be used in both horizontal and vertical beams. A schematic drawing of the NMI water calorimeter is given in Figure 4 below.

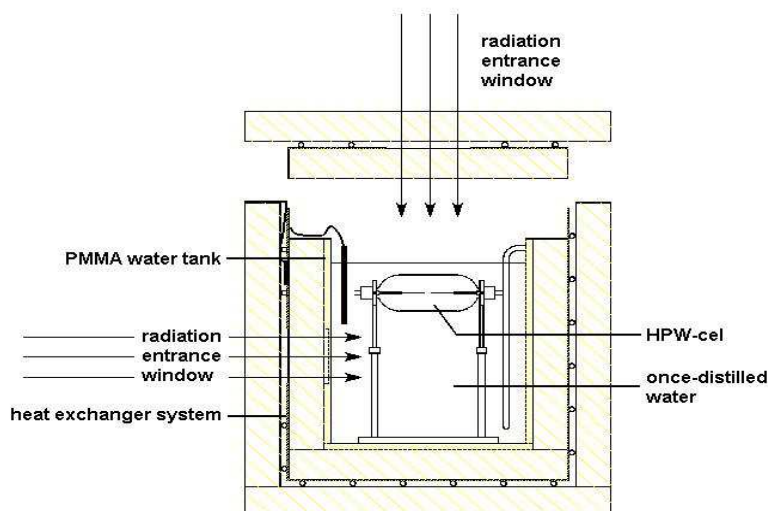


Figure 4: Schematic diagram of the NMI portable water calorimeter which was used for the measurement of k_Q values described in appendix A.4.

The calorimeter is equipped with an electrical measurement assembly comprising high precision digital multimeters (DMM) as read-out devices for the determination of the resistance changes of the thermistors during irradiation. Further details are given in [45, 46]. The water calorimeter has been compared with the graphite calorimeter of NMI for ^{60}Co gamma radiation. To convert the dose to graphite measured with the graphite calorimeter to dose to water the fluence scaling theorem is used [45]. Various correction factors of the water calorimeter as a function of photon energy have been determined experimentally and with the use of a computer program to simulate heat transport. Further test measurements were carried out in clinical photon beams in several Belgian hospitals and at NKI with the water calorimeter.

A.3.2 The LSDG water calorimeter

The construction of the sealed water calorimeter at the university of Ghent is based on the design of Domen [21] and has been described in detail in [47-50]. A short description is given here. The calorimeter tank consists of a 30 cm × 30 cm × 30 cm water phantom thermally isolated by polystyrene foam as shown in Figure 5a.

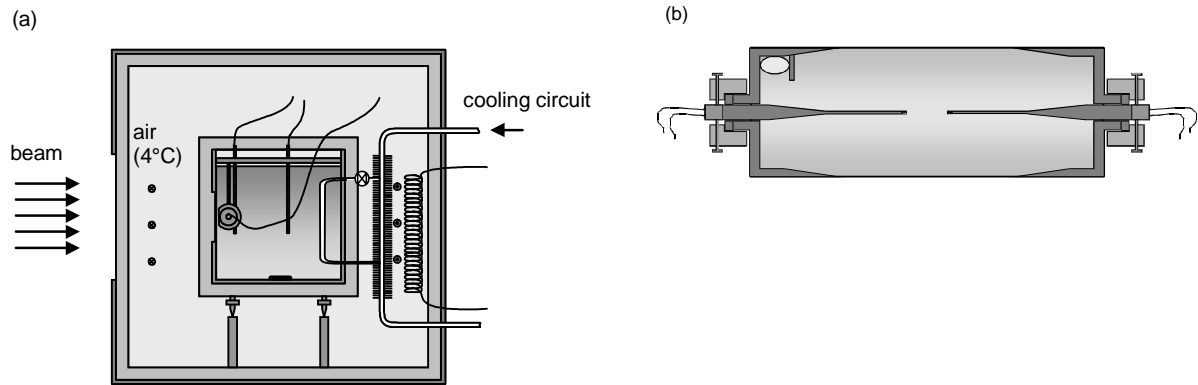


Figure 5: (a) SWC phantom and wooden enclosure of the Ghent water calorimeter and (b) cylindrical glass vessel with thermistor probes positioned (adapted from ref [50]).

Isolation of the enclosure allows temperature stabilization of the air surrounding the calorimeter phantom. The operating temperature is 4 °C in order to remove any concern related to convection. The water temperature is continuously measured with calibrated Pt-resistor probes inserted at different positions in the water phantom. The temperature increase due to irradiation is measured at the centre of a cylindrical vessel using small thermostats that are embedded in the tip of small glass probes as shown in Figure 5b. The probe consists of thermostats embedded in glass rods with a diameter of 0.5 mm. The vessel contains de-ionised and 3 times distilled water and has a diameter of 4 cm, a length of 14 cm and a wall thickness varying from 0.3 mm in the central part corresponding with the central axis of the beam to 1.6 mm at the lateral edges. The water inserted in the vessel is saturated with argon, nitrogen or hydrogen, three systems that do not result in a chemical heat defect. The electronics and the methods to determine the measured temperature increase are described in [47, 48].

A.3.3 Experimental Calorimetric determination of absorbed dose to water

Absorbed dose to water D_w is derived from the temperature increase ΔT due to irradiation at the location of the thermistors [50] as:

$$D_w = c_w \cdot \Delta T \cdot k_c \cdot k_{sc} \cdot k_{dd} \cdot \frac{1}{1-h} \quad (24)$$

where c_w is the specific heat capacity of water at the measuring temperature, k_c a correction factor for conductive heat effects (which is calculated by numerically solving the heat transport equation), k_{sc} a correction factor for scattering and attenuation induced by probes and vessel (which for the thermistors is negligible and for the vessel is determined by a perturbation measurement), k_{dd} a correction factor for the non-uniformity of the lateral dose distributions and h the chemical heat defect which corrects for the heat that is released or absorbed in chemical reactions by radiation induced species with each other or with impurities present in the water. The latter is calculated by numerically solving the set of coupled differential equations that describe all chemical reactions taking place and requires knowledge of the enthalpy of formation of each species involved in the reactions and is studied in detail in [47, 48]. In Tables 5a and 5b the uncertainty budgets for the determination of absorbed dose to water in a ^{60}Co photon beam for respective the LSDG and the NMi water calorimeter are shown.

Table 5a: Uncertainty budget for the determination of absorbed dose to water $D_{w,Q}$ in a ^{60}Co photon beam with the LSDG water calorimeter

	Input quantity x_i Source of uncertainty	Uncertainty type	Standard uncertainty
			u_i
Dose determination water calorimeter D_w in ^{60}Co			
	Reproducibility calorimeter response	A	0.20 %
	Bridge calibration	A	0.20 %
	Thermistor calibration	B	0.15 %
	Thermistor positioning	B	0.13 %
	Excess heat correction factor k_c	B	0.25 %
	Scatter correction factor k_{sc}	B	0.04 %
	Chemical heat defect h	B	0.50 %
	Combined Standard Uncertainty in $D_{w,Q}$		0.66 %

Note that the uncertainty analysis for the LSDG water calorimeter is still based on the comparison held at the BIPM in September 1999 [66]. Further work is foreseen to re-evaluate the uncertainty budget in the future.

Table 5b: Uncertainty budget for the determination of absorbed dose to water $D_{w,Q}$ in a ^{60}Co photon beam with the NMI portable water calorimeter

	Input quantity x_i Source of uncertainty	Uncertainty type	Standard uncertainty
			u_i
Dose determination water calorimeter D_w in ^{60}Co			
	Reproducibility calorimeter response	A	0.26 %
	Long-term stability in ^{60}Co	A	0.18 %
	Thermistor calibration	B	0.14 %
	Thermistor positioning	B	0.10 %
	Excess heat correction factor k_c	B	0.10 %
	Scatter correction factor k_{sc}	B	0.15 %
	Chemical heat defect h	B	0.20 %
	Combined Standard Uncertainty in $D_{w,Q}$		0.45 %

A.4 Experimental k_Q values and data for clinical high energy photon beams

A.4.1 Description of beams and ionisation chambers

k_Q values were measured for four cylindrical ionisation chamber types in 9 clinical high energy photon beams using the NMI portable water calorimeter. Details of these beams are shown in Table 6. The beam qualities were measured using a uniform protocol that is consistent with the recommendations of this CoP and TRS-398 for $\text{TPR}_{20,10}$ and TG-51 for $\%dd(10)_x$.

For each chamber type, 6 individual chambers were used. Details of the chamber types can be found in Table 3 (appendix A.1). They are all graphite-walled ionisation chambers with an aluminum central electrode. These characteristics fulfil the criteria for robustness and stability for reference ionisation chambers as described in TRS-398.

Table 6: The clinical beams in which k_Q were measured with their photon beam quality specifiers $TPR_{20,10}$ and $\%dd(10)_x$.

Date	Accelerator	Photon Energy (MV)	Institute	$TPR_{20,10}$	$\%dd(10)_x$
03/03	Varian 2300 C/D	15	Jolimont, Haine Saint-Paul, BE	0.7576	77.97
06/03	Elekta SI15	15	AMC, Amsterdam, NL	0.7287	73.06
06/03	Elekta SI25	25	UCL, Brussels, BE	0.7993	84.38
06/03	Elekta SI18	15	AZ St Luc, Gent, BE	0.7583	76.84
09/03	Siemens Primus	18	Virga Jesse Ziekenhuis, Hasselt, BE	0.7674	79.52
09/03	Elekta SI15	6	NKI/AvL, Amsterdam, NL	0.6732	66.24
12/03	Siemens Primus	23	Service de Radiotherapie, CHU de Liège, BE	0.7829	81.39
01/04	Varian 2300 C/D	6	VU, Amsterdam, NL	0.6594	65.82
11/04	Varian 2300 C/D	15	Jolimont, Haine Saint-Paul, BE (repeated measurement)	0.7563	78.05

A 4.2 Experimental method

The beam quality correction factor k_Q is defined as the ratio of absorbed dose to water calibration coefficients in the clinical beam quality Q , $N_{D,w,Q}$ and the reference (calibration) beam quality Q_0 , $N_{D,w}$ (^{60}Co in this case and therefore Q_0 is omitted):

$$k_Q = \frac{N_{D,w,Q}}{N_{D,w}} \quad (25)$$

The absorbed dose to water calibration coefficients are determined with the portable water calorimeter. For the beam quality Q :

$$N_{D,w,Q} = \frac{D_{w,Q}}{M_{\text{corr},Q}} \quad (26)$$

where:

- $N_{D,w,Q}$ calibration coefficient for reference chamber in beam quality Q ,
- $D_{w,Q}$ absorbed dose to water per monitor unit for beam quality Q ,
- $M_{\text{corr},Q}$ reference chamber signal per monitor unit.

Absorbed dose to water per monitor unit in the accelerator beam is derived as:

$$D_{w,Q} = \frac{1}{M_{mon,Q} \cdot k_{TP,mon,Q}} \cdot \Delta R_{midrun,Q} \cdot \frac{dT}{dR}(T_Q) \cdot c_w(T_Q) \cdot (\Pi k_i)_Q \quad (27)$$

where (for the beam quality Q):

$M_{mon,Q} \cdot k_{TP,mon,Q}$	monitor signal corrected for T and P,
$\Delta R_{midrun,Q}$	thermistor resistance change as a result of irradiation,
T_Q	calorimeter operating temperature at the point of measurement,
dT/dR	thermistor response as function of thermistor resistance,
$c_w(T_Q)$	specific heat capacity of water at the working temperature,
$(\Pi k_i)_Q$	product of correction factors for the chemical heat defect, for excess heat due to the presence of non-water materials, for the perturbation due to presence of the glass cell and for deviations from the reference SDD and measurement depth.

The overall measurement equation to derive k_Q thus becomes:

$$k_Q = \frac{M_{corr,Q}}{M_{corr,60Co}} \cdot \frac{M_{mon,60Co} \cdot k_{PT,mon,60Co}}{M_{mon,Q} \cdot k_{PT,mon,Q}} \cdot \frac{\Delta R_{midrun,Q} \cdot \frac{dT}{dR}(T_Q) \cdot c_w(T_Q)}{\Delta R_{midrun,60Co} \cdot \frac{dT}{dR}(T_{60Co}) \cdot c_w(T_{60Co})} \cdot \frac{(\Pi k_i)_Q}{(\Pi k_i)_{60Co}} \quad (28)$$

Note that a number of the factors occur both in numerator and denominator and these pairs are strongly correlated.

A.4.3 Uncertainties

Table 7 summarizes the uncertainty contributions in the experimental determined k_Q . A expanded standard uncertainty (coverage factor $k=2$) in k_Q of 1.1% is estimated. These uncertainty estimates apply to the portable water calorimeter, which was used during the measurement campaign in clinical photon beams in Belgium and The Netherlands. Correlations due to common uncertainty contributions are removed from the experimental data presented in Table 7.

Table 7: Combined standard uncertainty for experimental determined k_Q values.

Step 1: Absorbed dose determination in ^{60}Co		<i>Relative Standard Uncertainty (%)</i>
<i>Type of physical quantity or procedure</i>		
D_w determination with water calorimeter		0.40
Charge measurement and experimental procedure		0.16
<i>Combined uncertainty in $D_{w,\text{Co-60}}$</i>		<i>0.43</i>
Step 2: Absorbed dose determination in high energy photon beam		
<i>Type of physical quantity or procedure</i>		
D_w determination with water calorimeter		0.27
Charge measurement and experimental procedure		0.19
<i>Combined uncertainty in Step 2</i>		<i>0.33</i>
Combined standard uncertainty in k_Q (Steps 1 + 2)		0.54

A.4.4 Results of NCS measurement data

Table 8 lists the final data of the measurements. Each data point is the average value for the six ionisation chambers of the same type. The standard deviations on these six values was lower than 0.1% for all data points with the exception of two for which it was 0.2%. The measurement in Jolimont was performed twice and the agreement was good as can be seen in Table 8. In the further analysis the average result was used.

Table 8: Average k_Q per chamber type measured with the NMI portable water calorimeter in each of the high energy clinical photon beams.

Institute	TPR _{20,10}	%dd(10) _x	NE 2571	NE 2611	PTW 30012	Welhöfer FC65G
VU, Amsterdam, NL	0.6594	65.82	0.9940	1.0013	0.9932	0.9943
AvL/NKI, Amsterdam, NL	0.6732	66.24	0.9966	1.0006	0.9963	0.9996
AMC, Amsterdam, NL	0.7287	73.06	0.9840	0.9856	0.9778	0.9837
Jolimont, Haine Saint-Paul, BE	0.7563	78.05	0.9667	0.9703	0.9643	0.9679
Jolimont, Haine Saint-Paul, BE	0.7576	77.97	0.9681	0.9681	0.9655	0.9690
AZ St Luc, Gent, BE	0.7583	76.84	0.9789	0.9786	0.9771	0.9803
Virga Jesse Ziekenhuis, Hasselt, BE	0.7674	79.52	0.9769	0.9752	0.9738	0.9752
Service de Radiothérapie, CHU de Liège, BE	0.7829	81.39	0.9636	0.9690	0.9617	0.9632
UCL, Brussels, BE	0.7993	84.38	0.9654	0.9676	0.9634	0.9661

The data in Table 8 show no significant systematic trends in the differences between the Farmer types: NE 2571, PTW 30012 and Wellhöfer FC65G, which is not unexpected given the similarity in geometry and design of those three chamber types. In the subsequent analysis which will be discussed below, it was also found that the fits to the data for these three chamber types did not show significant differences. It was decided to pool the data for these three chamber types and to recommend a single data-set. An additional reason for that was that for the PTW 30012 and Wellhöfer FC65G chamber types no other data are available from the literature whereas for the NE 2571 there is a substantial amount of data available. The experimental k_Q data obtained in that way are presented in Figure 6.

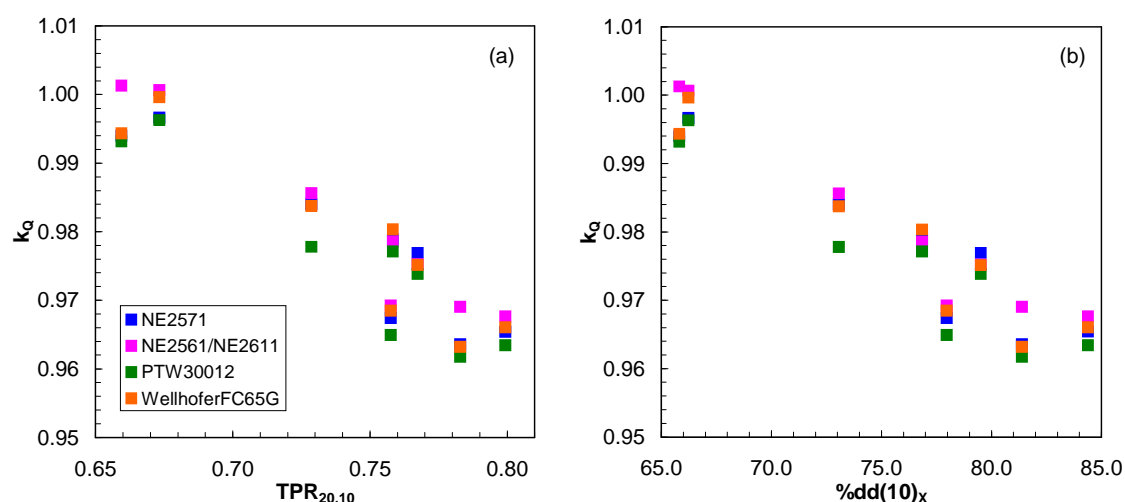


Figure 6: Results of the k_Q measurements for the Farmer types and the NE 2611A / NE 2561 types plotted (a) as a function of $TPR_{20,10}$ and (b) as a function of $\%dd(10)_x$.

A.4.5 Analysis of the experimental k_Q values

The data in Figure 6 are found to be insufficient and show too much scatter to be proposed as the single database for this protocol. Therefore, they are analysed in conjunction with other experimental data from the literature.

A.4.5.1 Compilation of experimental k_Q values from the literature

Only for the NE 2571 and NE 2561 / NE 2611A type chambers there is information available from the literature. All the k_Q values that were used in this analysis are listed in Table 9. It was decided that only data with an agreed quality standard and for heavily filtered (“clinic-like”) beams should be used for the determination of generic experimental k_Q data. The first criterion has been interpreted in TRS-398 as data measured by a primary standard

laboratory for a substantial number of chambers. This requirement is relaxed here such as to include the data measured by Palm et al. [51] and Medin et al. [52] since we did not find an argument that they would not have been measured up to the metrological quality achieved in a standards laboratory. On the contrary, given that they are measured in clinical beams they are more relevant to be included than some data measured in standard laboratories and their inclusion makes the generic set of k_Q values more robust. There has been some argument that data measured for a large number of chambers should be assigned a higher weight than other data for smaller sets of chambers. However, the NPL data as well as the NCS data show that chamber to chamber variations are small and that the spread of data in the entire set is more likely due to the differences in measurement method (within similar uncertainty limits) and the variety of beams. Hence, all the data points are given an equal weight in the further analysis. The data for the NE 2561 chambers from Palmans (1999) [49] were excluded since they were not obtained by direct measurement against a water calorimeter. The second criterion (to accept only heavily filtered beam data) excludes three data points from Ross et al. (1994) [54] and the data points by Seuntjens et al. (2000) [55]. Some of the data were measured using the Fricke ferrous sulphate chemical dosimeter of which Seuntjens et al. [55] have demonstrated that its response is not energy independent in the high energy x-ray range. Those results (Shortt et al. (1992) [56], Boutillon et al. (1994) [57], Ross et al. (1994) [54] and Guerra et al. (1995) [58]) have been corrected for this dependence with the equation presented by Seuntjens et al. [55]. For the LNHB data taken from Andreo (2000) [60] the values were back-corrected with the expression given by Andreo (2000) [60] and corrected with the expression given by Seuntjens et al. [55]. The Palm et al. (2002) [51] data have also been measured with Fricke but the results were already corrected in the same way by the authors.

For most of the data in this compilation the sleeve material was PMMA and the sleeve thickness was 1 mm but not for all data this information is given in the cited publications. Also, for at least three references the sleeve thickness was only 0.5 mm [53, 54, 58] and in one paper a nylon sleeve of 0.3 mm thickness was used [56]. These differences can make a small contribution to the spread in the data at the highest photon energies since it has been demonstrated that a 1 mm thick PMMA sleeve reduces the response of a Farmer type ionisation chamber by $0.2\% \pm 0.1\%$ in a 20 MV beam [92]. On the other hand, the uncertainty on this sleeve perturbation is substantial and correcting certain data points for this would have only a minor influence. The effect of these differences was ignored in the further analysis.

Recently, Krauss and Kapsch [72] reported experimental determined k_Q values in two high energy photon beams, which were not incorporated in the final analysis presented here. The k_Q values they reported in photon beams with $\text{TPR}_{20,10}$ values of 0.716 and 0.762 were 0.9889 and 0.9743, respectively, for the NE2571 and 0.9885 and 0.9759 for the NE2561, respectively.

Table 9: Experimental k_Q values for the NE 2571 and the NE 2561 / NE 2611A chamber from the literature.

Source reference	$\text{TPR}_{20,10}$	$\%dd(10)_x$	k_Q NE 2571	u (1sd)	k_Q NE 2561 /NE 2611A	u (1sd)
Shortt et al. (1992) [56]	0.7720 0.7850		0.9715 0.9660	0.0098 0.0097		
Boutillon et al. (1994) [57]	0.7110 0.7250 0.7640 0.7720				0.9924 0.9919 0.9827 0.9725	0.0100 0.0100 0.0099 0.0098
Ross et al. (1994) [54]	0.7580 0.7800 0.7910 0.7950 0.8150 0.8210	82.70 86.60 85.20 90.10 89.60 92.30	0.9729 0.9628 0.9672 0.9577 0.9602 0.9570	0.0039 0.0039 0.0039 0.0039 0.0039 0.0039		
Guerra et al. (1995) [58]	0.6730 0.7570	67.00 75.90	0.9946 0.9769	0.0065 0.0064	0.9956 0.9759	0.0065 0.0064
Palmans et al. (1999, 2002) [53, 59] Palmans (1999) [49]	0.6780 0.7440	65.60 74.20	0.9954 0.9786	0.0050 0.0052	0.9882 0.9729	0.0061 0.0071
Seuntjens et al. (2000) [55]	0.6820 0.7580 0.7940	69.60 80.50 88.40	0.9904 0.9723 0.9557	0.0038 0.0036 0.0036	0.9944 0.9724 0.9560	0.0038 0.0036 0.0036
LNHB data ($\text{TRP}_{20,10}$ and k_Q) from Andreo 2000) [60] ($\%dd10x$ from Ross 2005) [61] (corrected for difference Fricke correction Andreo and Seuntjens) [55]	0.6750 0.7170 0.7430 0.7490 0.7650 0.7670 0.7840 0.7850	67.36 71.63 75.02 75.63 78.43 78.75 81.96 82.15	0.9920 0.9880 0.9810 0.9810 0.9740 0.9730 0.9710 0.9680	0.0070 0.0079 0.0089 0.0089 0.0098 0.0098 0.0098 0.0097		
Palm et al. (2002) [51]	0.6210	63.00	0.9940	0.0070		

(Fricke corrected in the paper)	0.6720	66.80	0.9898	0.0069		
	0.7640	78.00	0.9705	0.0068		
	0.7930	82.20	0.9680	0.0068		
NPL (2003) [62]	0.6210	63.40	0.9958	0.0070	0.9986	0.0070
	0.6700	66.80	0.9910	0.0069	0.9941	0.0070
	0.7170	71.10	0.9865	0.0069	0.9885	0.0069
	0.7460	74.50	0.9808	0.0069	0.9815	0.0069
	0.7580	77.00	0.9766	0.0068	0.9781	0.0068
	0.7790	80.50	0.9695	0.0068	0.9717	0.0068
	0.7900	82.60	0.9635	0.0067	0.9670	0.0068
Stucki (2003) [63] From EUROMET-605 (data for $TPR_{20,10}=0.7847$ combined)	0.6264	62.84	0.9978	0.0055	0.9997	0.0055
	0.6392	63.71	1.0034	0.0055	1.0034	0.0055
	0.6695	66.29	0.9923	0.0054	0.9962	0.0055
	0.6741	66.77	0.9928	0.0055	0.9949	0.0055
	0.7160	70.75	0.9858	0.0054	0.9904	0.0054
	0.7478	74.29	0.9792	0.0054	0.9812	0.0054
	0.7592	75.39	0.9786	0.0054	0.9810	0.0054
	0.7627	76.18	0.9742	0.0053	0.9757	0.0054
	0.7847	81.83	0.9672	0.0053	0.9686	0.0053
	0.7981	83.28	0.9606	0.0053	0.9655	0.0053
McEwen (2006) (private communication) [64]	0.6810	67.20	0.9932	0.0040	0.9971	0.0040
	0.7314	72.60	0.9849	0.0039	0.9868	0.0039
	0.7998	84.40	0.9660	0.0039	0.9699	0.0039
LNHB (2006) From EUROMET-605 [65]	0.6780				0.9932	
	0.7520				0.9815	
	0.7830				0.9723	
Medin (2006) [52]	0.7340		0.9840	0.0069		

A.4.5.2 Analysis of the NCS data together with the data from the literature

The data measured by NCS, presented in Table 8, were added to this collection of data and as a function of $TPR_{20,10}$ sigmoidal fits were made of the form:

$$k_Q = A + \frac{B}{1 + \exp(C \cdot (TPR_{20,10} - X_0))} \quad (29)$$

where A, B, C and X_0 are fit parameters.

The justification for this is that such fits give an excellent agreement with the stopping power ratios from Andreo (1994) [73] that form at the basis of the calculated k_Q values in TRS-398. Also the second largest contribution to the beam quality dependence of k_Q for a graphite-walled ionisation chamber, the wall correction factor p_{wall} , can be accurately modelled with such a function. Although third order polynomials are found to be accurate for these theoretical data as well, they have the disadvantage that for experimental data, modest fluctuations can give rise to substantial changes in shape of the curve. A sigmoid on the other hand will still preserve its smooth character.

The model is first normalised such that the value of k_Q for a $TPR_{20,10}$ of 0.57 is unity. Here the assumption is made that for a high energy photon beam with a $TPR_{20,10}$ similar to the one for ^{60}Co k_Q is unity. An uncertainty is associated with this assumption. This results in the reduction of the number of free parameters in the model with one. In order to decide which of the four parameters to express in function of the other three, a fit to the NE 2571 data was performed and the parameter to which the fit is least sensitive is selected. It was found that this is parameter B, leading to

$$k_Q = A + (1 - A) \cdot \frac{1 + \exp(C \cdot (0.57 - X_0))}{1 + \exp(C \cdot (TPR_{20,10} - X_0))} \quad (30)$$

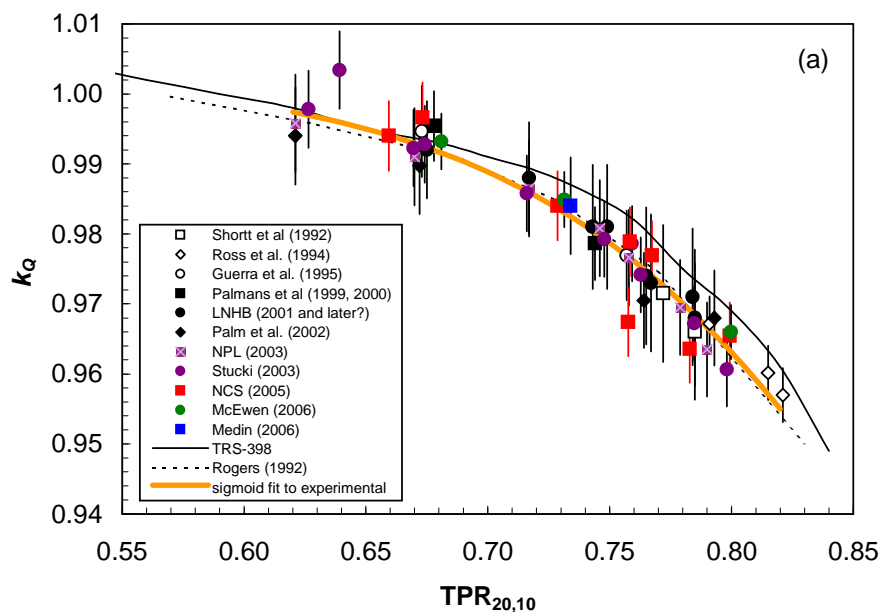
The parameters in the model are optimised with a least squares fitting algorithm (Levenberg-Marquardt) [70, 71].

The k_Q data as a function of $\%dd(10)_x$ are fitted with a linear curve. The justification of this is similar as for the dependence as function of $TPR_{20,10}$, i.e. that the stopping powers in the high energy x-ray beams are linearly dependent on $\%dd(10)_x$ [69]. Some higher polynomial fits were tried to investigate if they lead to a better fit.

Figures 7 and 8 show all the data for the NE 2571 and the NE 2561/NE 2611A as a function of both beam quality specifiers together with the sigmoid and linear fits as described above. In these sigmoid fits A was given the constant value of 0.80. The reason for this will be explained below. Theoretical data from TRS-398, TG-51 and Rogers (1992) [20] are shown as well.

The root mean square of the deviation from both models and both ionisation chamber types is comparable (around 0.3%). Performing quadratic and cubic polynomial fits to the data as a function of $\%dd(10)_x$ did not significantly alter this, so the linear model is deemed accurate enough. We conclude that there is no obvious preference of one beam quality specifier above another. It was decided to use $TPR_{20,10}$ as beam quality specifier and the sigmoid fit is

the best model we can find to represent the beam quality dependence of the data. The most important characteristic of the beam quality index $\text{TPR}_{20,10}$ is its independence on the electron contamination in the incident beam. It is also a measure of the effective attenuation coefficient describing the approximately exponential decrease of a photon depth-dose curve beyond the depth of maximum dose. $\text{TPR}_{20,10}$ is obtained as a ratio of doses; it does not require the use of displacement correction factors at two depths when cylindrical chambers are used. For this reason, $\text{TPR}_{20,10}$ is practically not affected by systematic errors in positioning the chamber at each depth, as the settings in the two positions will be affected in a similar manner. Finally, it is a parameter that is normally available through commissioning measurements and does not require special measurements using lead attenuators as it is the case for the $\%dd(10)_x$ parameter [67-69].



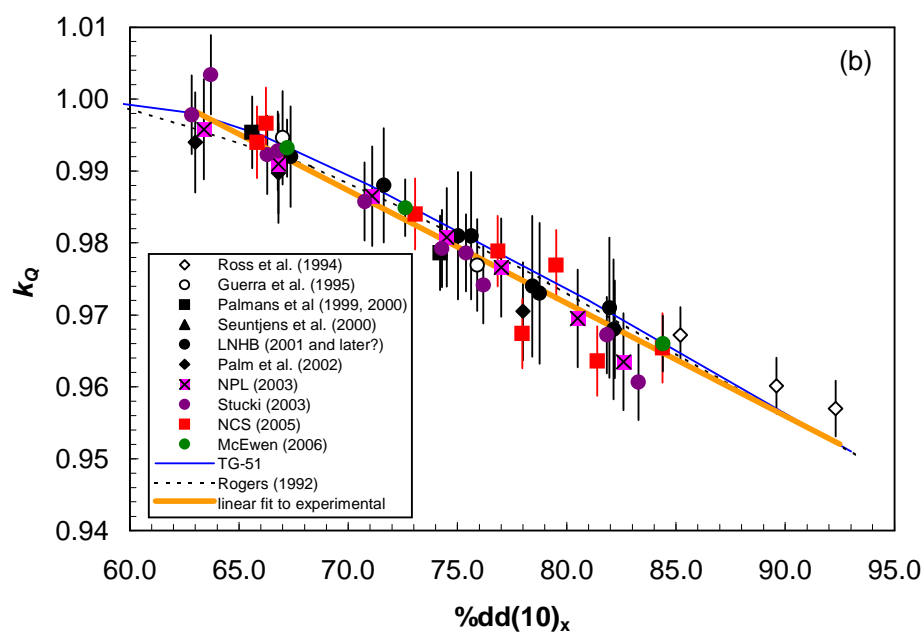


Figure 7: k_Q data from the literature together with the data measured by NCS for the NE 2571 type ionisation chamber (a) as function of $TPR_{20,10}$ with a sigmoid fit (with $A=0.80$) and (b) as function of $\%dd(10)_x$ with a linear fit.

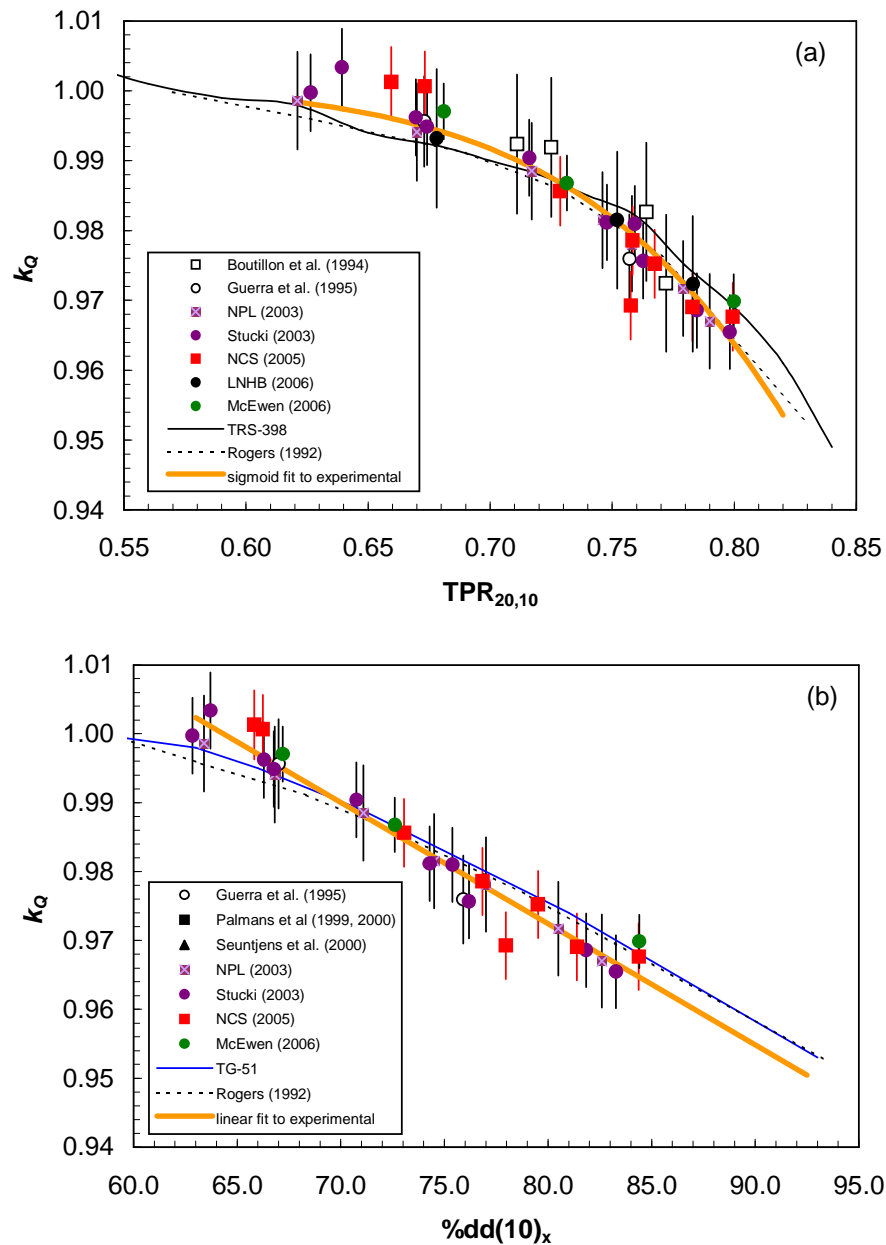


Figure 8: k_Q data from the literature together with the data measured by NCS for the NE 2561 and NE 2611A type ionisation chamber (a) as function of $TPR_{20,10}$ with a sigmoid fit (with $A=0.80$) and (b) as function of $\%dd(10)_x$ with a linear fit.

Figure 9 shows all the data points for the NE 2571 and the NE 2561/NE 2611A with various sigmoid fits in which none, one or two parameters are kept constant. Curves defining 1sd levels were calculated for the model by propagating through the model the 1sd values of the fit parameters obtained from the Levenberg-Marquardt algorithm [70, 71]. In addition an uncertainty of 0.02 (1sd) was assigned to the $TPR_{20,10}$ value of 0.57 where k_Q was assumed to be unity. This is consistent with the spread of $TPR_{20,10}$ values in TRS-398 where k_Q is unity

for cylindrical ionisation chambers. It can be seen in Figure 9 that the contribution to the overall uncertainty is not large given that in the region around the $\text{TPR}_{20,10}$ value of 0.57 the curve is fairly flat. The overall standard deviation is obviously energy dependent; it is small at $\text{TPR}_{20,10} = 0.57$ and grows as the beam quality increases.

It is obvious that leaving three parameters free leads to uncertainty estimates that are far too large, also the model tends to curve upward at the higher beam qualities which is unrealistic, compared with the theoretical beam quality dependence of the stopping powers. It was concluded that three free parameters in the model leaves too much degrees of freedom and the (remaining) parameter to which the model was least sensitive, which was found to be A, was kept constant. A range of values for A between 0.5 and 0.9 were tried and it was found that at $A=0.80$ the standard deviation of the model was such that about 2/3 data points fall within these 1sd curves. The tails of the distributions were less conform to a normal distribution but about 90% within the 2sd curves and about 95% within the 3sd curves for both the NE 2571 and NE 2561/NE 2611A. Hence it was decided that the sigmoid models with $A=0.80$ give a good representation of the experimental data with their experimental noise.

The lower graphs in Figure 9 show that keeping only one parameter free leads to an underestimation of the uncertainties.

The resulting values of the parameters C, A and X_0 in the sigmoid model are given in Table 10. The amount of digits given is sufficient to reproduce the fits presented above to within 0.01%. The uncertainties on the parameters in Table 10 which were determined by the fit algorithm result in a relative standard uncertainty in the k_Q values varying from 0.1% to 0.4%. In order to enable the user a way of verifying the implementation of this parameterized model, the data are plotted in Figure 10.

The recent data of Krauss and Kapsch [72] are close (and within the uncertainties equal) to the values obtained with the parameterised model presented in this work. Therefore it is assumed that the addition of those data in a new analysis will not significantly affect the outcome of the fits.

Table 10: Fit parameters X_0 , C and A in the sigmoid model resulting from this work

	X_0	C	A
NE 2571/ PTW 30012 / Wellhöfer FC65G	0.9198 ± 0.0097	11.67 ± 0.92	0.80
NE 2561/NE 2611A	0.8971 ± 0.0125	15.15 ± 1.60	0.80

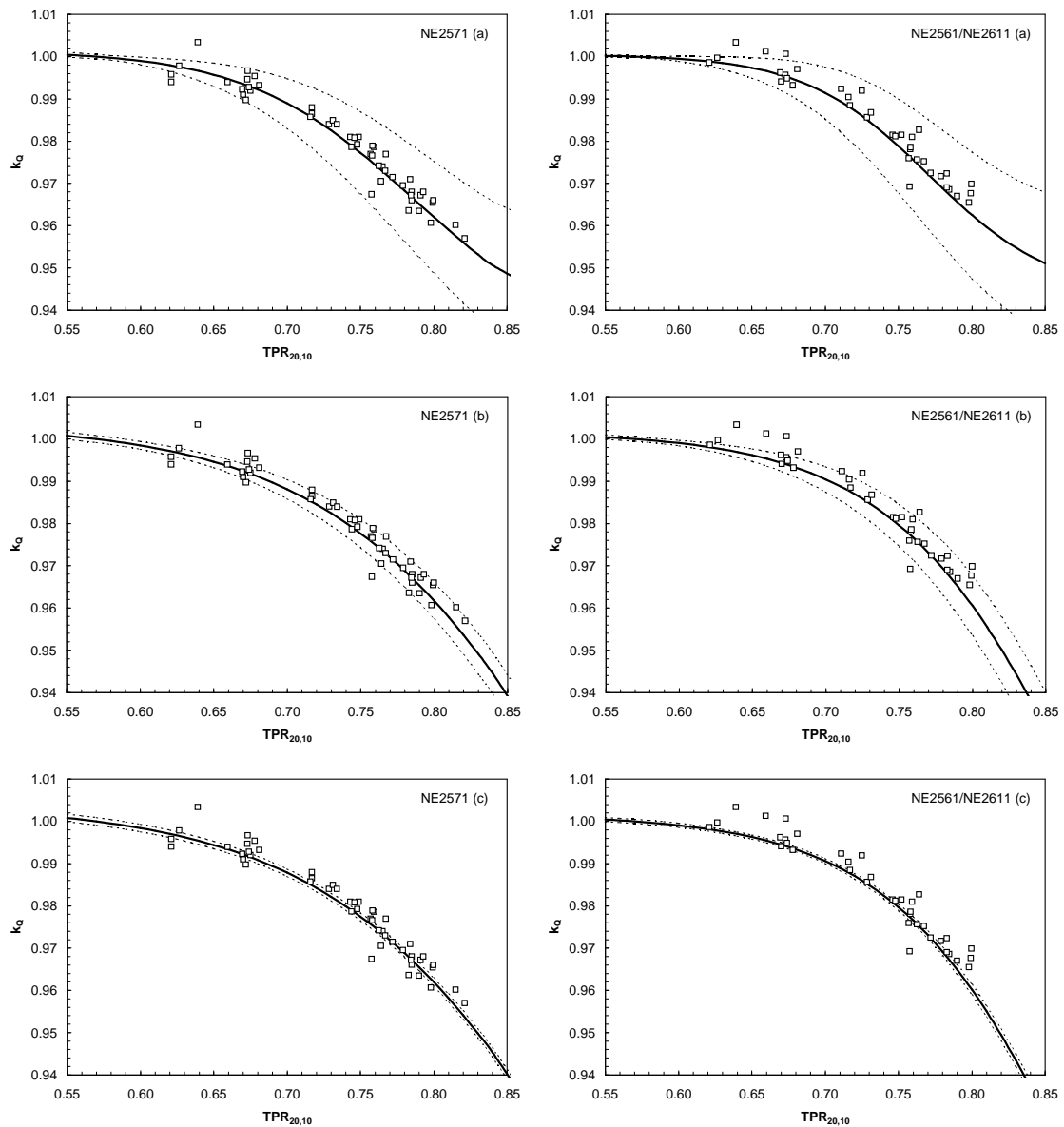


Figure 9: Sigmoid fit (solid lines) to all the data points (a) with X_0 , C and A as free parameters, (b) with X_0 and C as free parameters and $A=0.80$ and (c) with only X_0 as free parameter and C and A as in (b) for the NE 2571 (left) and for the NE 2561/NE 2611A (right). The dotted lines give the one standard deviations of the model for each fit.

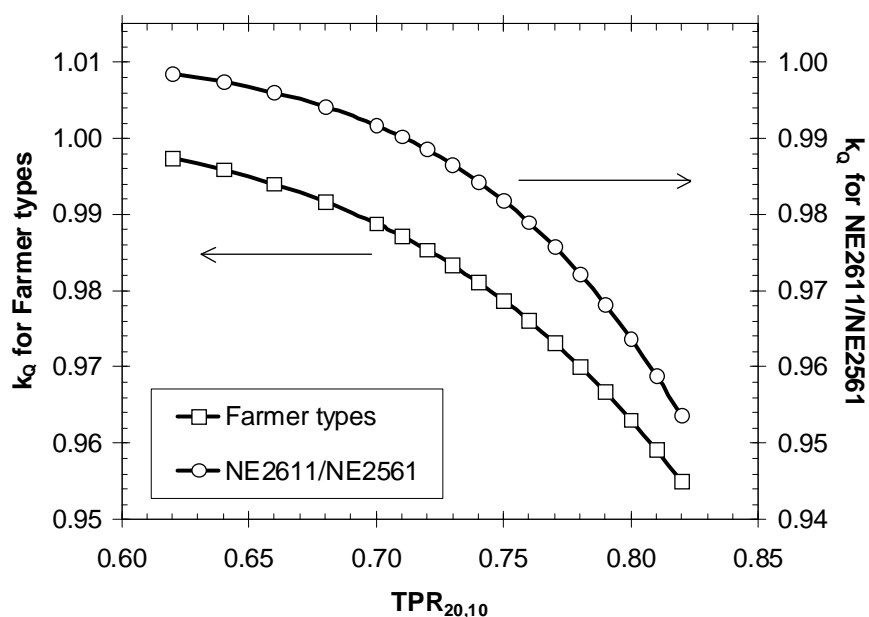


Figure 10: k_Q data calculated according to equation (30) and the data of Table 10. This plot is given for verification purposes.

A.4.6 Limiting value for the expected uncertainty

An estimate of the uncertainties in the calibration of a high energy photon beam is given in Table 11*. In order to determine the uncertainty on the absorbed dose to water in the user's beam by using a reference ionisation chamber, one must evaluate the uncertainties in the different physical quantities or procedures that contribute to the dose determination. These can be divided into two steps. Step 1 considers uncertainties up to the calibration of the reference dosimeter in terms of $N_{D,w}$ at the primary standard dosimetry laboratory. Step 2 deals with the calibration of the user's beam and includes the uncertainties associated with the measurements at the reference point in a water phantom. Step 2 also includes the uncertainty of the k_Q value. A more detailed evaluation of the uncertainty contribution, due to k_Q with the use of the sigmoid fit, is described in appendix A.4, section A.4.5.2.

* The uncertainty estimates given in Table 11 should be considered typical values; these values may vary depending on the expanded standard uncertainty quoted by Standards Laboratories for calibration coefficients and on the experimental uncertainty determined at the user's institution.

Table 11: Estimated combined standard uncertainty on $D_{w,Q}$ at the reference depth in water for a high energy photon beam.

Step 1: Primary Standard Dosimetry Laboratory (NMI, LSDG)	<i>Relative Standard Uncertainty (%)</i>
<i>Type of physical quantity or procedure</i>	
$N_{D,w}$ calibration of the secondary (local) standard at PSDL	0.5
Long term stability of the secondary standard	0.3
<i>Combined uncertainty in Step 1</i>	0.6
Step 2: User beam	
<i>Type of physical quantity or procedure</i>	
Dosimeter reading $M_{\text{corr},Q}$ (incl. influence quantities corrections $\sum k_i$)	0.5
Experimental procedure and stability of the dosimetry system	0.5
Beam quality correction, k_Q (based on NCS and literature data)	0.4
<i>Combined uncertainty in Step 2</i>	0.8
Combined standard uncertainty in $D_{w,Q}$ (Steps 1 + 2)	1.0

A.5 k_Q values and data for clinical high energy electron beams

For electron beams few experimental k_Q or $k_{Q,Q_{\text{cross}}}$ data are available in the literature. Guerra et al. [58] measured k_Q in a 14 MeV electron beam with a beam quality of $R_{50} = 5.27$ cm. The measurement was however performed at the depth of maximum dose $z_{\text{max}} = 2.8$ cm, which used to be the recommended reference depth in earlier codes of practice. With this NCS protocol the measurement should have been performed at a depth of $z_{\text{ref}} = 3.06$ cm. For the NE 2571 and NE 2561 they found k_Q to be 0.906 and 0.904 respectively. Experimental k_Q data-sets for plane-parallel chambers were reported by Stewart and Seuntjens [75] and by Stucki et al. [63] but in both cases no final published values are available. NPL has a substantial set of $k_{Q,Q_{\text{cross}}}$ data for plane-parallel chambers. However, they are measured using graphite calorimetry and the conversion procedure from absorbed dose to graphite to absorbed dose to water introduces a substantial uncertainty in these values. Furthermore, this conversion procedure uses NACP02 plane-parallel ionisation chambers as transfer instrument assuming that wall and cavity perturbation factors in electron beams both in graphite and in water are unity for this chamber types. This assumption is likely to be invalid as shown in a number of recent papers by Verhaegen et al, Buckley and Rogers, and McEwen et al. [76-78].

A.5.1 Calculation of k_Q and $k_{Q,Q_{cross}}$

As shown before, k_Q and $k_{Q,Q_{cross}}$ are per definition ratios of calibration coefficients in terms of absorbed dose to water in two beam qualities:

$$k_Q = \frac{N_{D,w,Q}}{N_{D,w,^{60}\text{Co}}} \text{ and } k_{Q,Q_{cross}} = \frac{N_{D,w,Q}}{N_{D,w,Q_{cross}}} \quad (31)$$

where the beam quality index ^{60}Co is shown explicitly even though it is omitted in the rest of this report. This notation does not conform to the CoP but is used here for clarity to stress that k_Q concerns a ratio of calibration coefficients in an electron and a photon beam.

Both these relations may be further generalized to:

$$k_{Q,Q_0} = \frac{N_{D,w,Q}}{N_{D,w,Q_0}} = \frac{\frac{D_{w,Q}}{M_{Q,corr}}}{\frac{D_{w,Q_0}}{M_{Q_0,corr}}} \quad (32)$$

where Q_0 stands for either ^{60}Co or the cross calibration beam quality Q_{cross} . Given the Bragg-Gray relation between absorbed dose to water and absorbed dose to air in the ionisation chamber's air cavity:

$$k_{Q,Q_0} = \frac{\frac{D_{air,Q}}{M_{Q,corr}} (s_{w,air})_Q p_Q}{\frac{D_{air,Q_0}}{M_{Q_0,corr}} (s_{w,air})_{Q_0} p_{Q_0}} = \frac{\frac{(W_{air}/e)_Q}{m_{air}} (s_{w,air})_Q p_Q}{\frac{(W_{air}/e)_{Q_0}}{m_{air}} (s_{w,air})_{Q_0} p_{Q_0}} \approx \frac{(s_{w,air})_Q p_Q}{(s_{w,air})_{Q_0} p_{Q_0}} \quad (33)$$

since the mean energy required to produce an ion pair in dry air, W_{air} , is assumed to be independent of energy in photon and electron beams. $(s_{w,air})_Q$ in equation (33) is the Spencer-Attix water to air collision mass stopping power ratio for the beam quality Q and the overall perturbation correction factor p_Q conventionally consists of four contributions:

$$p_Q = (p_{dis} p_{cav} p_{wall} p_{cel})_Q \quad (34)$$

where p_{dis} corrects for the difference in ionisation at the effective point of measurement and the depth at which the absorbed dose is stated, p_{cav} is a factor to correct for the response of an ionisation chamber for deviations from Bragg-Gray conditions due to the effect of the air cavity in the medium, p_{wall} corrects for the difference in composition between the ionisation chamber wall and water and p_{cel} corrects for the presence of the central electrode.

For stopping power data for ^{60}Co and for clinical electron beams there is no new information available since the publication of TRS-398. For the ^{60}Co calibration beam $s_{w,\text{air}} = 1.133$ and for electron beams the data are taken from Burns et al. [31], i.e. at the measurement depth z_{ref} ($= 0.6 R_{50,\text{dos}} - 0.1$ cm) the water to air mass stopping power ratio as a function of the beam quality $R_{50,\text{dos}}$ (in cm) is given by:

$$s_{w,\text{air}} = 1.253 - 0.1487 \cdot (R_{50,\text{dos}})^{0.214} \quad (35)$$

The choice of $R_{50,\text{dos}}$ as the beam quality index is related to these considerations on the stopping power ratios. This choice is justified since it results in a simplification of the procedures in NCS-5 involving the derivation of the mean energy at the phantom surface based on $R_{50,\text{dos}}$ as an intermediate step. It also directly relates the beam quality specifier to the penetration characteristics of the electron beam. The definition of $R_{50,\text{dos}}$ is made in terms of absorbed dose levels, whereas usually ionisation curves are measured. Contrary to the situation in high energy photon beams, the variation of the water/air stopping-power ratios with depth is not negligible. Therefore, strictly, the depth ionisation curve should be converted to a depth dose curve using an accelerator dependent water-to-air stopping-power function of depth as was done in NCS-5. However generic expressions to derive $R_{50,\text{dos}}$ from $R_{50,\text{ion}}$ for a large number of accelerator types has proven to be sufficient.

A.5.1.1 Farmer-type chambers

For the perturbation correction factors for Farmer type chambers there is no new data available since the publication of TRS-398. Hence, for Farmer type chambers k_Q is calculated as in equation (33) using data taken from TRS-398. Differences between the three Farmer type chambers were found to be negligible. The product $(s_{w,\text{air}})_{60\text{Co}} p_{60\text{Co}} = 1.102$.

The only perturbations that are considered in the electron beam are p_{cav} and p_{cel} . p_{cav} depends on the beam quality and the cylindrical cavity radius r_{cyl} (in mm) as [87]:

$$p_{cav} = 1 - 0.0217 \cdot r_{cyl} \cdot e^{-0.153 \cdot R_{50,dos}} \quad (36)$$

A mean radius of 3.15 mm was assumed for the three Farmer type chambers.

For p_{cel} a uniform value of 0.998 was taken independent of electron beam quality [88].

The displacement correction factor p_{dis} is inherently accounted for by using the concept of an effective point of measurement (see section 3.4). p_{wall} is assumed to be unity for Farmer type chambers.

The k_Q values thus obtained for the Farmer type chambers can be summarized into a single formula of the form:

$$k_Q = A - B \cdot [R_{50,dos}(Q)]^C \quad (37)$$

The rationale behind this expression is that the underlying water to air mass stopping powers follow such a relationship (see equation 35). The parameters A, B and C for calculating k_Q with this expression for the Farmer type chambers are given in Table 13. The maximum deviation between the calculated values for k_Q and the values obtained with equation 37 is 0.05% in the beam quality index range from 4 to 20 cm.

A.5.1.2 Plane-parallel chambers

For plane-parallel chambers only p_{cav} and p_{wall} play a role, under the assumption that gradient corrections are taken care of adequately by putting the effective point of measurement at the reference depth, hence:

$$k_{Q,Q_{cross}} = \frac{(s_{w,air})_Q}{(s_{w,air})_{Q_{cross}}} \cdot \frac{(p_{cav}p_{wall})_Q}{(p_{cav}p_{wall})_{Q_{cross}}} \quad (38)$$

For well-guarded plane-parallel chambers, such as the one recommended in this report, both TRS-398 and TG-51 assume that both perturbations factors are unity. Regarding p_{cav} , various papers indicate that it is constant as a function of energy (Ma et al. [79], Verhaegen et al. [76]) for well-guarded chambers. So even if the value is non-unity it would not influence the $k_{Q,Q_{cross}}$ values since only ratios of p_{cav} are needed. Regarding p_{wall} , however, several recent papers have demonstrated a variation of p_{wall} at the reference depth with beam quality

(Sempau et al. [80], Verhaegen et al. [76], Buckley and Rogers [77] and McEwen et al. [78]). Significant is that two different Monte Carlo codes were used (PENELOPE by Sempau et al. [80] and EGSnrc by Verhaegen et al. [76] and Buckley and Rogers [77]) and one semi-empirical model (by McEwen et al. [78]). All of them predict a similar tendency of p_{wall} values slightly above unity at low electron energies and increasing with decreasing energy. It was decided to estimate improved p_{wall} values from these data.

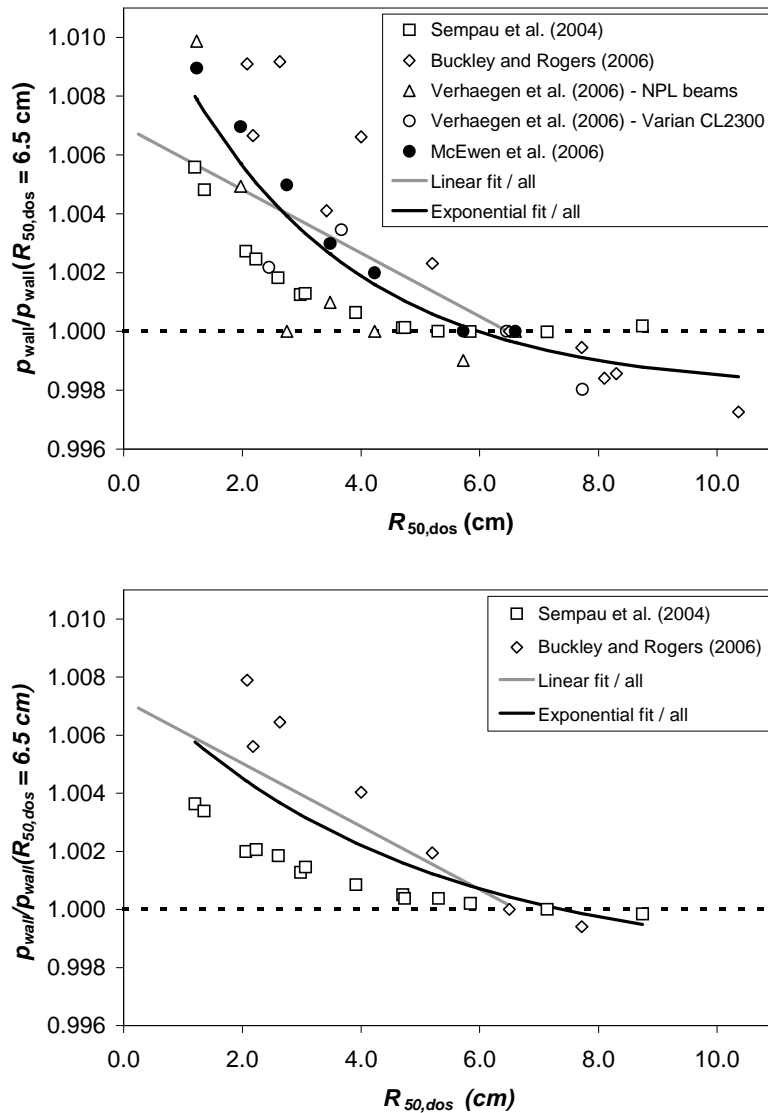


Figure 11: p_{wall} values (relative to the values at $R_{50,\text{dos}} = 6.5 \text{ cm}$) obtained from Monte Carlo simulations and semi-empirical model calculations for the NACP02 and the Roos chambers. Linear and exponential fits are shown.

Figure 11 shows these p_{wall} values (relative to the values at $R_{50,dos} = 6.5$ cm) for the NACP02 and the Roos chambers. The normalization at $R_{50,dos} = 6.5$ cm is a bit arbitrary but is not important for $k_{Q,Qcross}$ since again only ratios will be considered. Linear fits to all the data are shown as well as exponential fits of the form:

$$p_{wall}(z_{ref}) = a + b \cdot e^{c \cdot R_{50,dos}} \quad (39)$$

Note that the fit parameters a,b and c are constant. It is obvious that there is considerable spread on the data points, possibly reflecting the variability of the p_{wall} values for beams of varying type, and that the differences between the various fit models are not significant. The exponential fits were used for the further evaluation of $k_{Q,Qcross}$ data.

The $k_{Q,Qcross}$ values calculated in that way can again be represented by a single equation of the form:

$$k_{Q,Qcross} = \frac{A - B \cdot [R_{50,dos}(Q)]^C}{A - B \cdot [R_{50,dos}(Q_{cross})]^C} \quad (40)$$

The parameters A, B and C for calculating $k_{Q,Qcross}$ with this expression for the NACP02 and Roos type chambers is given in Table 13. The maximum truncation error introduced by this approximation is 0.05% in the beam quality range from 1 to 20 cm.

Although we emphasize once again that it is not recommended to use plane-parallel chambers with ^{60}Co calibration coefficients the option is left open in case an electron beam with sufficiently high energy to perform a cross-calibration is not available. In this case k_Q values for the plane-parallel chambers need to be calculated as follows:

$$k_Q = \frac{(s_{w,air})_Q}{(s_{w,air})_{^{60}\text{Co}}} \cdot \frac{(p_{cav} p_{wall})_Q}{(p_{cav} p_{wall})_{^{60}\text{Co}}} \quad (41)$$

The main problem here is that the perturbation factors for the plane-parallel chamber in the ^{60}Co calibration beam have a considerable uncertainty. Various papers in the literature have reported that these perturbation factors can vary by 3% for chambers of the same type so it is difficult to recommend accurate generic data per chamber type. Nevertheless, for NACP02 and Roos type chambers the variability of these reported perturbation factors is not so dramatic and since the publication of TRS-398 various investigations have resulted in more

accurate data for p_{wall} in ^{60}Co by comparing dosimetry using both calibration routes (cross calibration in a high energy electron beam or calibration in ^{60}Co). These data are shown in Table 12.

Table 12: Experimental p_{wall} values in ^{60}Co for the NACP02 and Roos plane-parallel chamber types recommended in this report as reported in the literature.

Reference	Based on	p_{wall}	
		NACP02	PTW~Roos
Ding et al. (1998) [81]	TG-51	1.026	
Palm et al. (2000) [82]	TRS-381	1.020	1.014
Dohm et al. (2001) [83]	DIN 6800-2 [90]		1.024
Dohm et al. (2001) [83]	TG-51		1.020
Palm et al. (2002) [84]	TRS-381		1.018 ^(*)
Stewart and Seuntjens (2002) [85]	TG-51	1.020	1.015
Stewart and Seuntjens (2002) [85]	TRS-398	1.020	1.016
Christ et al. (2002) [86]	DIN 6800-2 [90]		1.024
Christ et al. (2002) [86]	TG-51		1.020
Palmans et al. (2003) [17]	TRS-398	1.015	
Palmans et al. (2003) [17]	TG-51	1.016	
Kapsch et al. (2007) [89]	DIN 6800-2 [90]		1.020
	TRS-398/TRS-381 (average)	1.018	1.016
	u (1sd)	0.3%	0.2%
	TG-51 (average)	1.021	1.018
	u (1sd)	0.5%	0.3%

^(*)Data point for the Wellhöfer-PPC35 ionisation chamber which is assumed to be of the same design

One comment on these data is that, although all these authors report to have measured a value of p_{wall} in ^{60}Co for the plane-parallel chamber, they have actually measured a ratio of overall perturbation factors in ^{60}Co and the high energy beam assuming that the perturbation factors for the cylindrical chamber are perfectly known. The latter explains why the values determined based on TRS-398 are different from the values based on TG-51 and DIN 6800-2 since different data for the cylindrical chambers are recommended in different protocols.

Hence for consistency, in the further analysis the perturbation correction factor data from TRS-398 for cylindrical chambers have been used.

Also noteworthy is that recent Monte Carlo simulated values have been published for p_{wall} by Mainegra-Hing et al. [91] who found values of 1.021 for the NACP02 chamber and 1.009 for the Roos chamber. Even though they may represent more accurate values for p_{wall} itself, it was decided to adopt the values of Table 12 for consistency with experimental data on the ratio of overall perturbations as mentioned above. These data can then be used to come up with a data set of k_Q values for the plane-parallel chambers consistent with the Farmer data and the $k_{Q,Qcross}$ data. Equation (41) can be modified to:

$$k_Q = \frac{(s_{w,air})_Q}{(s_{w,air})_{Co}} \cdot \frac{(p_{cav} p_{wall})_Q}{(p_{cav} p_{wall})_{high-e}} \cdot \frac{(p_{cav} p_{wall})_{high-e}}{(p_{cav} p_{wall})_{Co}} \quad (42)$$

where the first two terms have been discussed already before where the high-e quality index refers to the cross calibration beam quality. The third term in equation (42) equals the reciprocal of the values presented in Table 12.

The resulting k_Q values can again be summarized into a single formula of the form of equation (37) with the coefficients given in Table 13. The maximum truncation error introduced by this approximation is 0.05% in the beam quality range from 1 to 20 cm.

A.5.2 Calculated k_Q values for high energy electron beams

The resulting coefficients to calculate k_Q and $k_{Q,Qcross}$ with equations (37) and (40) are given in Table 13. In Figure 12 and Figure 13 the data for k_Q and $k_{Q,Qcross}$ as a function of $R_{50,dos}$ are presented.

Table 13: coefficients to calculate k_Q and $k_{Q,Qcross}$ with equations (37) and (40).

	A	B	C
Farmer types	0.9345	0.0057	0.7733
NACP02	1.1955	0.2274	0.1479
Roos types	1.1376	0.1700	0.1835

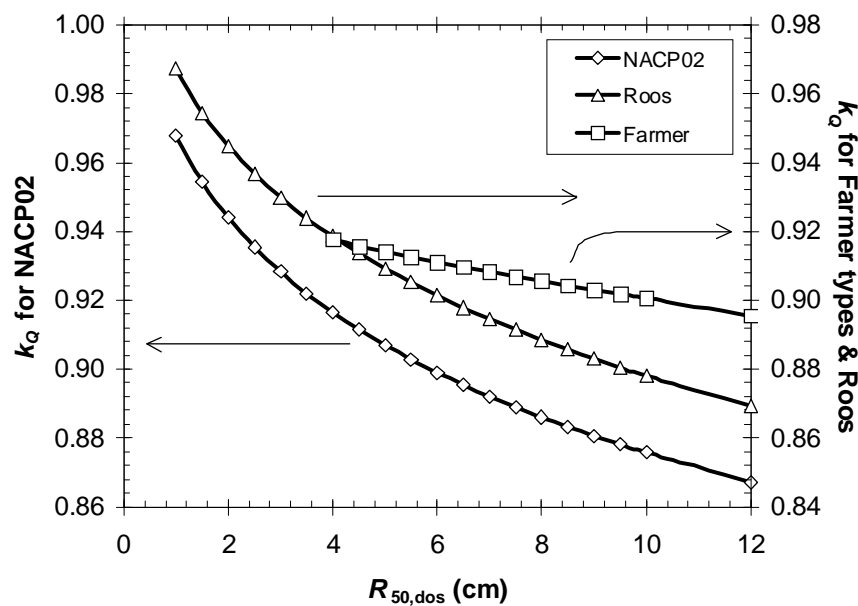


Figure 12: k_Q data as a function of $R_{50,dos}$ calculated according to equation (37) for the chamber types considered in this code of practice. This plot is given for verification purposes.

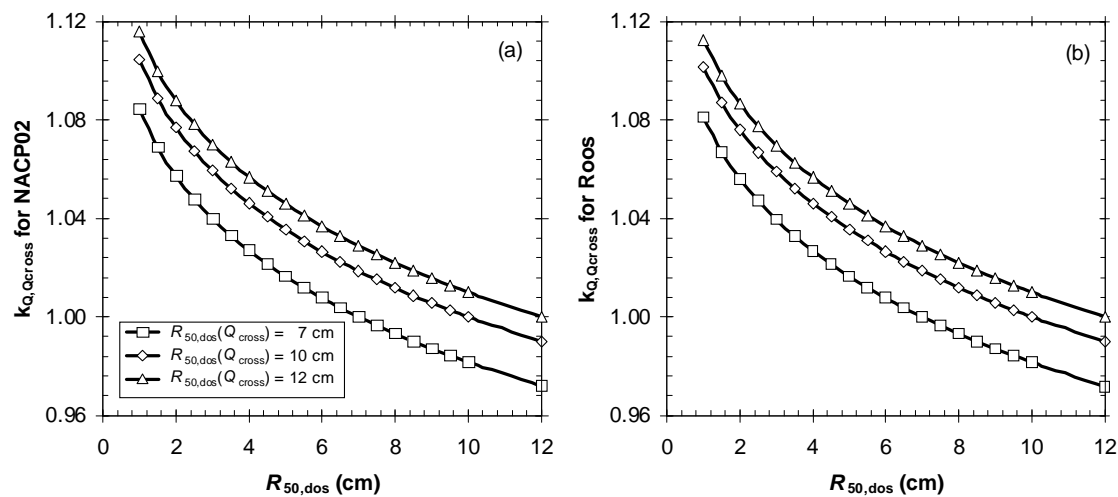


Figure 13: $k_{Q,Qcross}$ data as a function of $R_{50,dos}$ calculated according to equation (40) for (a) NACP02 chambers and (b) Roos chambers. These plots are given for verification purposes.

A.5.3 Comparison with experimental data

Experimental data regarding k_Q in electron beams are very scarce. Guerra et al. [58] reported an experimental k_Q value of 0.904 for a NE 2571 in a 14 MeV electron beam with an $R_{50} = 5.27$ cm. The present NCS report predicts a value of 0.914 for these conditions. This is a reasonable agreement given the difference in conditions (the measurement was done at the depth of dose maximum). Also the use of a different standard (Fricke) might influence the result.

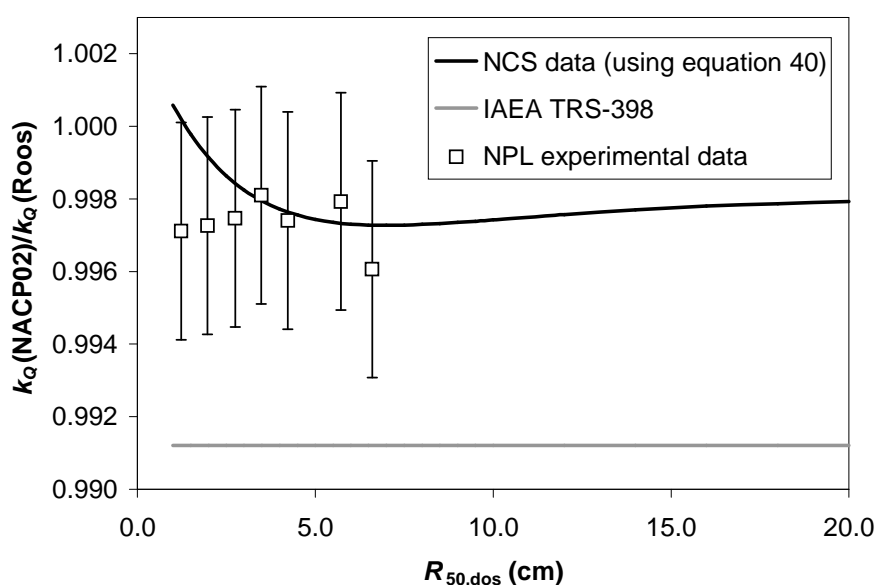


Figure 14: Ratio of k_Q data for the NACP02 and Roos type chambers compared with the experimental values from the NPL database.

NPL has a substantial set of calibration data for NACP02 and Roos type chambers. Even though there are at present question marks about the absolute calibration coefficients (the non-unity perturbation factors for the NACP02 mentioned before influence the conversion procedure from dose to graphite to dose to water, needed because the NPL calibrations are based on graphite calorimetry) the ratios of the k_Q factors can be determined with a good accuracy. Figure 14 shows that the k_Q ratios between the NACP02 and Roos type chambers calculated in this report are in better agreement with the NPL data than the data from TRS-398. The curl-up effect in the NCS data shown in figure 14 is due to small differences

between p_{wall} calculations for NACP02 and Roos chambers derived from the data presented in Figure 11.

A.5.4 Limiting value for the expected uncertainty

An estimate of the uncertainties in the calibration of a high energy electron beam is given in Tables 14, 15 and 16*. In order to determine the uncertainty on the absorbed dose to water in the user's beam by using a reference ionisation chamber, one must evaluate the uncertainties in the different physical quantities or procedures that contribute to the dose determination. These can be divided into two or three contributions depending on the calibration procedure followed. Contribution one (Step 1 in Tables 14 and 16) considers uncertainties up to the calibration of the reference dosimeter in terms of $N_{D,w}$ at the standard dosimetry laboratory. Contribution two deals with either the absorbed dose determination in a electron beam with a Farmer type cylindrical ionisation chamber (Step 2 in Table 14) or the calibration of a plane-parallel chamber in the cross-calibration beam (Step 1 in Table 15), both including the uncertainties associated with the measurements at the reference point in a water phantom. Contribution three (Step 2 in Table 15 and 16) deals with the absorbed dose determination in a electron beam with a plane-parallel ionisation chamber and includes the uncertainties associated with the measurements at the reference point in a water phantom. Contributions two and three also include the uncertainty of the k_Q and $k_{Q,Q_{\text{cross}}}$ values. A more detailed evaluation of the uncertainty budget, including the uncertainty on k_Q and $k_{Q,Q_{\text{cross}}}$ is described in appendix A 7.

Although the estimated uncertainty on dosimetry using a plane-parallel ionisation chamber using a cross calibration is not much lower than using a ^{60}Co calibration beam, it has been demonstrated by various audits that the consistency of clinical reference dosimetry is substantially improved when implementing the cross-calibration procedure [33].

* The uncertainty estimates given in Tables 14,15 and 16 should be considered typical values; these values may vary depending on the expanded standard uncertainty quoted by Standards Laboratories for calibration coefficients and on the experimental uncertainty determined at the user's institution.

Table 14: Estimated combined standard uncertainty on $D_{w,Q}$ measured with a Farmer type chamber at the reference depth in water for a high energy electron beam.

Step 1: Primary Standard Dosimetry Laboratory	<i>Relative Standard Uncertainty (%)</i>
<i>Type of physical quantity or procedure</i>	
$N_{D,w}$ calibration of the secondary (local) standard at PSDL	0.5
Long term stability of the secondary standard	0.3
<i>Combined uncertainty in Step 1</i>	0.6
 Step 2: Farmer with ^{60}Co calibration coefficient in an electron beam with $R_{50} > 4$ cm	
<i>Type of physical quantity or procedure</i>	
Dosimeter reading (incl. influence quantities corrections $\prod k_i$), $M_{\text{corr}, Q}$	0.5
Experimental procedure and stability of the dosimetry system	0.5
Beam quality correction, k_Q	1.2
<i>Combined uncertainty in Step 2</i>	1.4
Combined standard uncertainty in $D_{w,Q}$ (Steps 1 + 2)	1.5

Table 15: Estimated combined standard uncertainty on $D_{w,Q}$ measured with a plane-parallel ionisation chamber at the reference depth in water for a high energy electron beam based on a cross calibration in a high energy electron beam.

Step 1: Cross calibration of plane-parallel chamber in a beam with $R_{50} > 7$ cm	
<i>Type of physical quantity or procedure</i>	<i>Relative Standard Uncertainty (%)</i>
Dose determination using the cylindrical chamber in a high energy electron beam (from Table 14)	1.5
Dosimeter reading (incl. influence quantities corrections $\sum k_i$), $M_{corr, Q}$ *	0.0
Experimental procedure and stability of the dosimetry system	0.5
<i>Combined uncertainty on $N_{D,w,Qcross}$ in Step 1</i>	<i>1.6</i>
 Step 2: Plane-parallel chamber with cross-calibration coefficient in an electron beam	
<i>Type of physical quantity or procedure</i>	
Dosimeter reading (incl. influence quantities corrections $\sum k_i$), $M_{corr, Q}$	0.5
Experimental procedure and stability of the dosimetry system	0.5
Beam quality correction, $k_{Q,Qcross}$	0.6
<i>Combined uncertainty in Step 2</i>	<i>0.9</i>
Combined standard uncertainty on $D_{w,Q}$ (Steps 1 + 2)	1.8

* The dosimeter reading in step 2 in Table 14 and the dosimeter reading in step 1 in table 15 are taken in the same beam quality and are therefore highly correlated. We assumed for the combined uncertainty contribution of these two dosimeter readings a value of 0.5 %

Table 16: Estimated combined standard uncertainty on $D_{w,Q}$ measured with a plane-parallel ionisation chamber at the reference depth in water for a high energy electron beam based on a ^{60}Co calibration coefficient in a PSDL.

Step 1: Primary Standard Dosimetry Laboratory	<i>Relative Standard Uncertainty (%)</i>
<i>Type of physical quantity or procedure</i>	
$N_{D,w}$ calibration of the secondary (local) standard at PSDL	0.5
Long term stability of the secondary standard	0.3
<i>Combined uncertainty in Step 1</i>	0.6
 Step 2: Plane-parallel chamber with ^{60}Co calibration coefficient in an electron beam	
<i>Type of physical quantity or procedure</i>	
Dosimeter reading (incl. influence quantities corrections $\sum k_i$), $M_{corr,Q}$	0.5
Experimental procedure and stability of the dosimetry system	0.5
Beam quality correction, k_Q	1.7
<i>Combined uncertainty in Step 2</i>	1.8
Combined standard uncertainty in $D_{w,Q}$ (Steps 1 + 2)	2.0

A.5.5 Data for measurement in non-reference conditions

The water to air mass stopping power ratios vary considerably with depth and in order to derive a depth dose curve from a measured depth ionisation curve with an ionisation chamber stopping power data as a function of beam quality and depth are needed. Burns et al. [31] derived from the same Monte Carlo simulations that led to equation (35) a more complicated equation for these data:

$$S_{w,air}(z) = \frac{a + b \cdot \ln(R_{50}) + c \cdot [\ln(R_{50})]^2 + d \cdot \frac{z}{R_{50}}}{1 + e \cdot \ln(R_{50}) + f \cdot [\ln(R_{50})]^2 + g \cdot [\ln(R_{50})]^3 + h \cdot \frac{z}{R_{50}}} \quad (43)$$

with $a = 1.075$, $b = -0.5087$, $c = 0.0887$, $d = -0.084$, $e = -0.4281$, $f = 0.0646$, $g = 0.00309$ and $h = -0.125$.

Rogers [74] showed that for a wide range of accelerators these data are accurate to within 1.0% except at very shallow depths or depths beyond $1.1 z/R_{50}$.

A.6 Expected differences with NCS-2 and NCS-5

The ratio of dose to water values obtained using this protocol and the old protocols is given by:

$$\frac{D_{w,Q}(NCS-18)}{D_{w,u}(NCS-2)} = \frac{N_{D,w}}{N_K} \times \frac{k_Q}{C_{w,u}(Q)} \quad (44)$$

for photon beams and

$$\frac{D_{w,Q}(NCS-18)}{D_{w,e}(NCS-5)} = \frac{N_{D,w,Q_0}}{N_K} \times \frac{k_{Q,Q_0}}{C_{w,e}(Q)} \quad (45)$$

for electron beams, assuming that the corrected ionisation chamber readings are the same for both protocols. In equation (45) k_{Q,Q_0} stands for either k_Q or $k_{Q,Q_{cross}}$ depending on the calibration route and correspondingly N_{D,w,Q_0} stands for the absorbed dose to water calibration coefficient in either a cross calibration beam Q_{cross} or a ^{60}Co photon beam.

It is clear that the ratio of dose to water values obtained using this protocol and the old protocols is determined by two components: the ratio of absorbed dose to water to air kerma calibration coefficients and the ratio of conversion factors. The former is dependent on the calibration laboratory and chamber type whereas the second is purely dependent on the data used for a particular chamber type.

The interpretation of these differences can be made easier by introducing a 'calculated' absorbed dose to water calibration coefficient $N_{D,w}^*$ and k_Q values derived from the old protocols.

The 'calculated' absorbed dose to water calibration coefficient is calculated as:

$$N_{D,w}^* = N_K \times C_{w,u}(^{60}\text{Co}) \quad (46)$$

and the k_Q derived from the old NCS protocols can be obtained by multiplying the numerator and denominator of expression (33) with the product $k_{att}k_m k_{ce}(1-g)$, which is entirely related to in-air perturbations during an air kerma calibration. For photon beams this gives:

$$k_Q(NCS-2) = \frac{(s_{w,air})_Q p_Q}{(s_{w,air})_{^{60}\text{Co}} p_{^{60}\text{Co}}} \times \frac{k_{att} k_m k_{ce} (1-g)}{k_{att} k_m k_{ce} (1-g)} = \frac{C_{w,u}(Q)}{C_{w,u}(^{60}\text{Co})} \quad (47)$$

and for a plane-parallel chamber in a electron beam ($k_{ce}=1$):

$$k_{Q,Q_{cross}}(NCS-5) = \frac{(s_{w,air})_Q p_Q}{(s_{w,air})_{Q_{cross}} p_{Q_{cross}}} \times \frac{k_{att} k_m (1-g)}{k_{att} k_m (1-g)} = \frac{C_{w,e}(Q)}{C_{w,e}(Q_{cross})} \quad (48)$$

For photons, equation (44) can then be re-written as:

$$\frac{D_{w,Q}(NCS-18)}{D_{w,u}(NCS-2)} = \frac{N_{D,w}}{N_{D,w}^*} \times \frac{k_Q(NCS-18)}{k_Q(NCS-2)} \quad (49)$$

The expression is thus split in two factors. The first one is the ratio of a measured and a calculated absorbed dose to water calibration coefficient and its deviation from unity indicates an inconsistency between the theoretical data in the old protocol and the measured ratio of absorbed dose to water and air kerma calibration coefficients. This factor is independent of the data in the new code of practice. The second one reflects the change of chamber dependent data for the clinical beams in the new code compared to the old code. The ratio of absorbed dose to water values can thus entirely be studied in terms of calibration data for a particular chamber in the standard laboratory and data used in the protocol.

A similar expression can be written for electron dosimetry based on a calibration in ^{60}Co with a Farmer type chamber (or the unfavourable option of a plane-parallel chamber calibrated in ^{60}Co):

$$\frac{D_{w,Q}(NCS-18)}{D_{w,e}(NCS-5)} = \frac{N_{D,w}}{N_{D,w}^*} \times \frac{k_Q(NCS-18)}{k_Q(NCS-5)} \quad (50)$$

In this case the expression for $k_Q(NCS-5)$ is analogous to equation (47) using the data of the old electron code of practice (where for a plane-parallel chamber k_{ce} equals 1).

For electron dosimetry using a plane-parallel chamber with a cross calibration, the differences are more complicated since the N_K and $N_{D,w}$ calibration coefficients are for different chambers, but it can be easily shown, that:

$$\frac{D_{w,Q}(NCS-18)}{D_{w,e}(NCS-5)} = \frac{N_{D,w}^{Farmer}}{N_K^{pp}} \times \frac{M_{Q_{cross}}^{Farmer}}{M_{Q_{cross}}^{pp}} \times \frac{k_{Q_{cross}}^{Farmer}}{C_{w,e}^{pp}(Q_{cross})} \times \frac{k_{Q,Q_{cross}}^{pp}(NCS-18)}{k_{Q,Q_{cross}}^{pp}(NCS-5)} \tag{51}$$

so the ratio can again be assessed in terms of a ratio of calibration coefficients and protocol data, but in addition there is a ratio of two chamber readings in the clinical beam involved. The ratio is thus not solely due to measurements in the standards laboratory and protocol data but can be influenced by the local set-up in the clinical beam. This could for example lead to a different ratio being observed by different users with the same equipment in the same beam.

The expected differences based on these equations for the chambers recommended in this report are discussed below for photon and electron beams.

A.6.1 High energy photon beams

The data to evaluate equation (49) for high energy photon beams are given in Table 17. The ratio of the measured and the calculated calibration coefficient is a generic value derived from the paper by Palmans et al. (2002) [16]. Hence, the values in this example are indicative and the user who wants to know the change for his particular therapy unit and equipment should evaluate this ratio for each individual calibrated ionisation chamber. The ratio of the k_Q data is also shown as a function of $TPR_{20,10}$ in Figure 15. The two contributions to the difference are tabulated separately but since the ratio of the measured and the calculated calibration coefficient is very close to unity, it is easy to see that the overall differences are smaller than 1% in all cases.

Table 17: Components according to equation (49) of typical differences in absorbed dose to water in high energy photon beams measured with this code of practice and NCS-2.

(1) From Palmans et al. (2002) taking into account a 0.3% difference between the disseminated air kerma calibrations and the air kerma calibrations reported in that paper.
(2) For the PTW 30012 and the Wellhöfer-FC65G no data are given in NCS-2. The same data as for the NE 2571 were assumed.

	NE 2571	PTW 30012	Wellhöfer FC65G	NE 2611A/NE 2561
PSDL	$\frac{N_{D,w}}{N_{D,w}^*}$			
LSDG ⁽¹⁾	0.999	0.999	0.997	—

$TPR_{20,10}$	$\frac{k_Q(NCS-18)}{k_Q(NCS-2)}^{(2)}$			
0.60	1.001	1.001	1.001	1.002
0.64	1.001	1.001	1.001	1.004
0.68	1.001	1.001	1.001	1.003
0.72	0.998	0.998	0.998	1.002
0.75	0.999	0.999	0.999	1.001
0.78	0.999	0.999	0.999	0.999
0.81	1.000	1.000	1.000	0.998

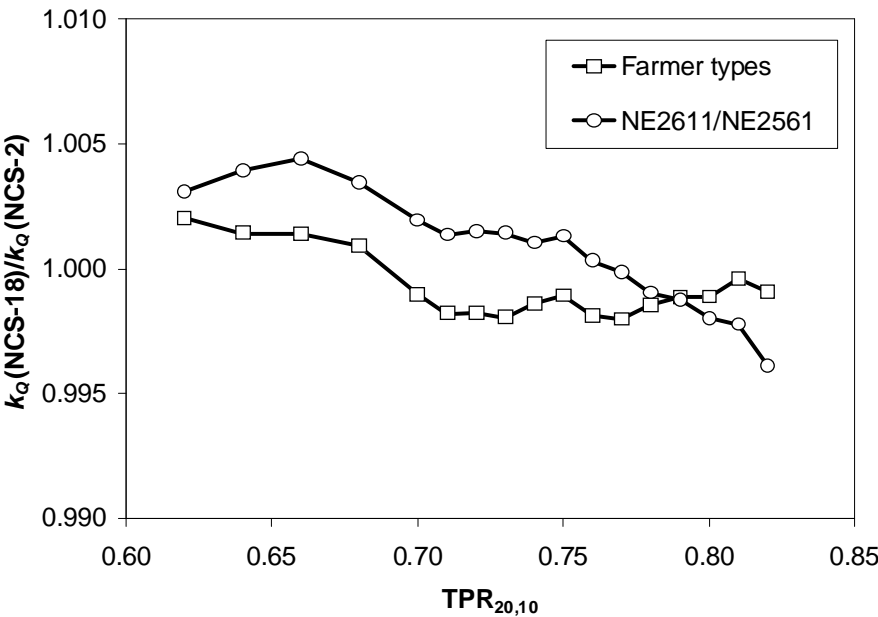


Figure 15: Ratio of k_Q values for photon beams recommended in this protocol and derived from NCS-2.

A.6.2 High energy electron beams

The data to evaluate equation (50) for high energy electron beams are given in Table 18 as well as the overall resulting absorbed dose to water ratios obtained with the new protocol and NCS-5. Here, the ratio of the measured and the calculated calibration coefficient are taken from the paper by Palmans et al. (2003) [17] and thus, again, the values are indicative and should be investigated by the user for his particular set-up. For the NACP02 chamber the air kerma calibration coefficient N_K was determined according the procedures in NCS-5, i.e. with a 3 mm graphite build-up plate. The ratios of k_Q data as a function of beam quality are also shown graphically in Figure 16. One complication in such a comparison exercise is that the

reference depths in both codes of practice can be different. This is in general the case for higher energies since the new reference depth can be deeper than the dose maximum. To make a generic comparison possible as an example, the $C_{w,e}$ values for NCS-5 were calculated at the new z_{ref} (derived from R_{50}) according to the tables of NCS-5. In principle, measurements at both depths should be taken and the ionisations converted with the different water to air collision mass stopping powers and perturbation correction factors at both depths to allow a rigorous comparison of dosimetry using the two protocols. The presented values are thus only indicative and a proper comparison should be performed for each electron beam unit and dosimetry system. The differences for the NE 2571 and the NACP02 are in general smaller than 1% over the entire range of beam qualities. Differences would not be expected to be larger than 1% so any larger deviation could be an indication that further investigation of the source of the difference is needed. The differences observed in Table 18 are consistent with the findings of Palmans et al. (2003) [17] for a limited set of three electron beam energies.

Table 18: Components and total values according to equation (50) of typical differences in absorbed dose to water in high energy electron beams measured with this code of practice and NCS-5 using a calibration coefficient in ^{60}Co .

⁽¹⁾ From Palmans et al. (2003) taking into account a 0.3% difference between the disseminated air kerma calibrations and the air kerma calibrations reported in that paper

⁽²⁾ The newest p_{wall} value for the NACP02 chamber in ^{60}Co (from this report) is used. This affects the ratio of k_Q data, but also the ratio of measured and "calculated" calibration coefficients over an amount which is equal but which corrects in the opposite direction. The overall resulting absorbed dose to water ratios are thus unaffected by this choice.

	NE 2571	NACP02	NE 2571	NACP02
PSDL	$\frac{N_{D,w}}{N_{D,w}^*}$			
LSDG ⁽¹⁾	0.999	0.994 ⁽²⁾		
R_{50}	$\frac{k_Q(NCS-18)}{k_Q(NCS-5)}$		$\frac{D_{w,Q}(NCS-18)}{D_{w,e}(NCS-5)}$	
1.0	N/A	1.011	N/A	1.005
2.0	N/A	1.005	N/A	0.999
3.0	N/A	1.001	N/A	0.995
4.0	0.995	1.000	0.994	0.994
5.0	0.998	1.002	0.997	0.996

7.5	1.005	1.001	1.004	0.995
10.0	1.007	1.001	1.006	0.995
12.0	1.004	1.001	1.003	0.995

A specific example of the evaluation of equation (51) for high energy electron beams in the case of a cross calibration beam quality of $R_{50} = 7$ cm is given in Table 19. The ratios of $k_{Q,Q_{cross}}$ data for this specific case as a function of beam quality are also shown graphically in Figure 16. The same remark as above on the different measurement depths in both codes, and thus on the non-generality of the data, applies here. This example is again based on the calibration data in Palmans et al. (2003) [17]. It must be noted that not only the measurement data can vary according to the conditions of the experiment. In addition, the ratio of $k_{Q,Q_{cross}}^{pp}(NCS-18)$ and $k_{Q,Q_{cross}}^{pp}(NCS-5)$ will show a slight dependence on the choice of the cross calibration beam quality since the data for the cross calibration beam are not the same in both protocols. For the beam quality range considered in this protocol, however, this variation was found to be smaller than 0.2%. The differences for the NACP02 are again smaller than 1% over the entire range of beam qualities. Differences would not be expected to be larger than 1% and a larger deviation could be an indication that further investigation of the source of the difference is needed. The differences observed in Table 19 are consistent with the findings of Palmans et al. (2003) [17] for a limited set of three electron beam energies.

Table 19: Components and total values according to equation (51) of typical differences in absorbed dose to water in high energy electron beams measured with this code of practice and NCS-5 using a plane-parallel chamber with a cross calibration coefficient.

	NACP02		
PSDL	$\frac{N_{D,w}^{Farmer}}{N_K^{pp}}$		
LSDG	0.3211		
	$\frac{M_{Q_{cross}}^{Farmer}}{M_{Q_{cross}}^{pp}}$	$\frac{k_{Q_{cross}}^{Farmer}}{C_{w,e}^{pp}(Q_{cross})}$	$\frac{N_{D,w}^{Farmer}}{N_K^{pp}} \times \frac{M_{Q_{cross}}^{Farmer}}{M_{Q_{cross}}^{pp}} \times \frac{k_{Q_{cross}}^{Farmer}}{C_{w,e}^{pp}(Q_{cross})}$
$R_{50}(Q_{cross}) \approx 7 \text{ cm}$	3.425	0.906	0.997
R_{50}		$\frac{k_{Q,Q_{cross}}^{pp}(NCS-18)}{k_{Q,Q_{cross}}^{pp}(NCS-5)}$	$\frac{D_{w,Q}(NCS-18)}{D_{w,e}(NCS-5)}$
1.0		1.009	1.005
2.0		1.003	1.000
3.0		0.999	0.996
4.0		0.998	0.995
5.0		1.000	0.996
7.5		0.999	0.996
10.0		0.999	0.996
12.0		0.999	0.995

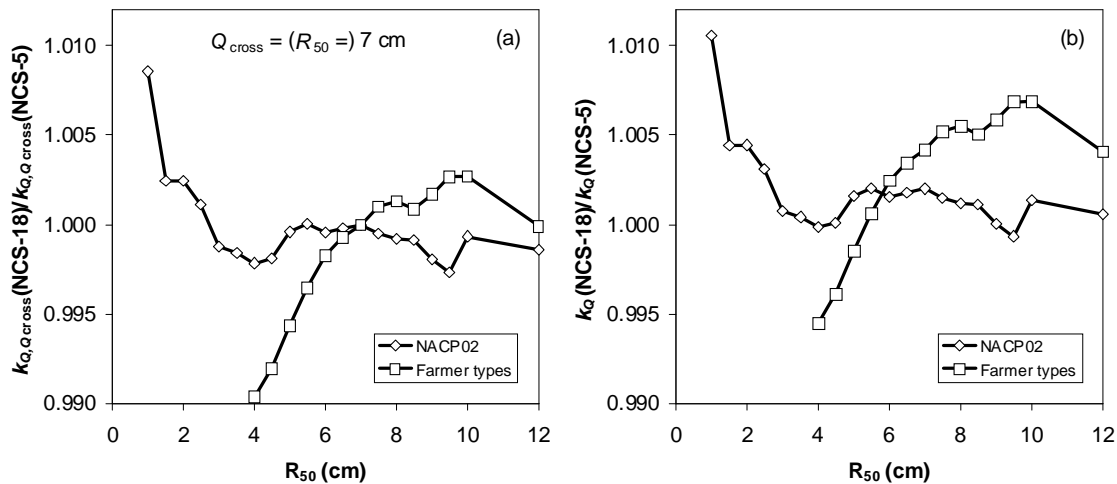


Figure 16: Ratio of (a) $k_{Q,Q_{cross}}$ values (b) k_Q values for electron beams recommended in this protocol and derived from NCS-5.

A.7 Expression of Measurement Uncertainty

A.7.1 Introduction

It is recommended to base the estimation of measurement uncertainty on the guidelines presented in the ISO *Guide to the Expression of Uncertainty in Measurement* [93] and in document EA-4/02 of The European Co-operation for Accreditation [94]. The uncertainty budgets reported in this CoP follow the principles and analysis methods set out in these documents and the assessment of the uncertainties associated to clinical measurement results should be carried out in accordance with the recommendations of the ISO Guide and EA-4/02. This appendix provides an example of an uncertainty analysis for an absorbed dose to water determination in a clinical photon beam produced by a medical linear accelerator. Usually the uncertainty in an absorbed dose determination is dominated by a number of uncertainty contributions, while others have a small or negligible effect. For practical reasons a value for the smallest uncertainty is set, which has to be considered in the uncertainty analysis. Contributions below this value will be ignored. However, these contributions have to be evaluated and documented. In the example given below, the limiting value for an uncertainty contribution (relative standard uncertainty) to be considered is set to 0.1%.

A.7.2 General procedure of the uncertainty estimation

The result of the $D_{w,Q}$ (output quantity) has to be calculated from a number of input quantities according to the equation:

$$D_{w,Q} = M_{corr,Q} N_{D,w} k_Q \quad (52)$$

where:

- $D_{w,Q}$ is the absorbed dose to water in the users beam Q at the reference depth
- $M_{corr,Q}$ the electrometer reading corrected for influence quantities, e.g. k_{TP} k_h k_{pol} k_s
- $N_{D,w}$ is the absorbed dose to water calibration factor in the ^{60}Co reference beam quality at $T_{ref} = 20^\circ\text{C}$ and $P_{ref} = 101.325 \text{ kPa}$ and 50% relative humidity.
- k_Q is the beam quality correction factor that depends on the chamber type and radiation beam quality of the users beam.

The output quantity $D_{w,Q}$ is considered a function of all input quantities (model equation) and consequently the uncertainty in $D_{w,Q}$ is determined from the standard uncertainties of all input quantities. The standard uncertainty of a corresponding input quantity is estimated in terms of one standard deviation associated with the assumed probability distribution. The standard uncertainties, denoted by the symbol u_i , are classified into type A and type B. Type A uncertainties are evaluated on the basis of a statistical analysis of a series of observations. Type B uncertainties are obtained using non-statistical methods. Examples of type B uncertainty contributions are: uncertainties stated in calibration certificates, uncertainties and tolerances stated in specifications given by the manufacturer, uncertainties obtained from literature (reference data). Usually the standard uncertainties for all components are expressed in a relative form. The standard uncertainty in $D_{w,Q}$ is termed combined standard uncertainty and denoted by u_c , which is determined from all uncertainty contributions of the input quantities u_i using the following equation:

$$u_c^2 = \sum_{i=1}^N (c_i \cdot u_i)^2 \quad (53)$$

where c_i are the sensitivity coefficients, defined by:

$$c_i = \partial D_{w,Q} / \partial x_i, \text{ with } i = 1, \dots, N \quad (54)$$

Note, that it is assumed that all sources of uncertainties are uncorrelated. Finally, the expanded uncertainty U for $D_{w,Q}$ is calculated by multiplying the combined standard uncertainty u_c by a coverage factor k , which is mostly chosen $k=2$.

A.7.3 Example of an uncertainty budget for the determination of absorbed dose to water $D_{w,Q}$ in a clinical photon beam Q

In this example the complete dosimetry system (ionisation chamber and electrometer) has been calibrated at the standards laboratory. Therefore $k_{\text{elec}} = 1.00$ and no uncertainty contribution is assumed. Before starting the measurement the dosimeter system was allowed sufficient time to stabilize after switching on and to reach temperature stabilization of the ionisation chamber in the water phantom. In addition the chamber is pre-irradiated to a dose of several gray (2-5 Gy) in order to settle adequately achieving charge equilibrium in the different materials involved. The effect of non-linearity in the response of the dosimeter system is considered to be very small and the resulting uncertainty is neglected. No

corrections are applied for the chamber orientation in the beam, for any departure of the field size or the radial non-uniformity of the beam and the resulting uncertainties are assumed to be negligible. It is assumed that during the measurements the ambient conditions of temperature, pressure and humidity are constant. As a consequence no uncertainty contributions arise.

A.7.3.1 Notes on the uncertainty contributions of the input quantities

All relevant uncertainty sources are considered individually and are presented in Table 20. Explanatory notes concerning the different uncertainty sources are given below:

Uncertainty in calibration certificate

The expanded uncertainty stated on the calibration certificate from a Primary Standard Dosimetry Laboratory is estimated to be 0.9% for a coverage factor $k=2$, which corresponds to a confidence level of about 95% assuming a normal distribution. Usually the calibration coefficient of a dosimeter changes during a time period of several years (typically 2-3 years) between calibrations. The relative change is about 0.3%. Taking this value as a limit and assuming a rectangular distribution a value of 0.173% for the relative standard uncertainty is determined.

Dose meter reading

The mean and the standard deviation are obtained from a series of readings (preferably at least ten readings should be taken). The leakage current should be measured before and after the irradiation. Typically the leakage current is less than 0.1% of the current measured during irradiation and can often be ignored or taken into account in the resulting uncertainty. In this example the dosimeter reading is corrected for the leakage current.

Often the readings are digitized and a resolution uncertainty has to be assigned of ± 0.5 to the last digit/count on the display assuming a rectangular distribution. In the example a resolution uncertainty of 0.10% is adopted.

Temperature and pressure correction k_{TP}

The correction factor k_{TP} in the air volume of the ionisation chamber is calculated according to equation (17) from the measured values of the temperature T (K) and the pressure P (hPa). Typical values of relative expanded uncertainties of 0.17% (0.5 K at a temperature of about 293.15 K) and 0.10% are taken from the certificates of the temperature and barometer calibration respectively.

As the temperature and pressure do not vary during the measurements no type A uncertainties arise. Type B uncertainties arising from the thermometer and barometer resolution result in very small relative standard uncertainties, which can often be neglected in practice. It should be noted that the temperature is not measured in the chamber cavity but close to the ionisation chamber in the water phantom. In this example the difference in temperature between the air cavity and the position of the sensing part of the thermometer in the water phantom is neglected.

Humidity correction k_h

In a range between 20%-80% relative humidity the response of an ionisation chamber changes not more than 0.1%. No correction for the humidity effect is made, but an uncertainty contribution assuming a rectangular distribution is assigned to k_h .

Recombination correction k_s

Using expression (21) k_s is calculated with a relative standard uncertainty of 0.1%.

Polarity correction k_{pol}

For most ionisation chamber types the polarity effect is small and often neglected in photon beams. The relative standard uncertainty in k_{pol} is typically estimated 0.1%.

Deviation of chamber position (depth) in phantom

The relative change in absorbed dose to water in the vicinity of the reference depth in the water phantom is 0.6%/mm. The uncertainty in positioning at the reference depth is 0.1 mm in terms of one standard deviation resulting in a relative standard uncertainty of 0.06%.

Beam quality correction factor k_Q

According to section A.4.5.2 of appendix A.4 the relative standard uncertainty in k_Q is estimated 0.4%.

Table 20: Uncertainty budget for the determination of absorbed dose to water $D_{w,Q}$ in a clinical photon beam Q

All individual uncertainty contributions are summed in quadrature and from the total sum the root is taken to give the combined standard uncertainty of the result: $u(D_{w,Q})/D_{w,Q} \approx 0.8\%$. The expanded uncertainty is determined by multiplying the combined standard uncertainty with a coverage factor $k=2$.

	Input quantity x_i Source of uncertainty	Relative (expanded) uncertainty	Uncertainty type	Uncertainty distribution	Coverage (reducing) factor	Relative standard uncertainty	Sensitivity coefficient	Uncertainty contribution
		U_i			k	u_i	c_i	$ c_i u_i $
Calibration coefficient, $N_{D,w}$								
	Uncertainty in calibration certificate	0.90%	B	normal	2	0.450%	1	0.450%
	Long term stability of calibration coefficient	0.30%	B	rectangular	1.73	0.173%	1	0.173%
Dosimeter reading, corrected for leakage, M								
	Repeatability	0.50%	A	normal	1	0.500%	1	0.500%
	Resolution	0.10%	B	rectangular	1.73	0.058%	1	0.058%
Temperature during measurement, T (K)								
	Thermometer calibration	0.34%	B	normal	2	0.170%	1	0.170%
	Thermometer resolution	0.03%	B	rectangular	1.73	0.020%	1	0.020%
Pressure during measurement, P (hPa)								
	Barometer calibration	0.10%	B	normal	2	0.050%	1	0.050%
	Barometer resolution	0.005%	B	rectangular	1.73	0.003%	1	0.003%
Humidity correction k_h								
	Deviation from reference condition of 50%	0.10%	B	rectangular	1.73	0.058%	1	0.058%
Recombination correction k_s								
	Uncertainty in recombination correction factor	0.10%	B	normal	1	0.100%	1	0.100%
Polarity correction k_{pol}								
	Uncertainty in polarity correction factor	0.10%	B	normal	1	0.100%	1	0.100%
Depth positioning ionisation chamber in phantom								
	Uncertainty due to deviation from ref. position	0.12%	B	normal	2	0.060%	1	0.060%
Beam quality correction factor k_Q								
	Uncertainty in the quality correction factor	0.8%	B	normal	2	0.400%	1	0.400%

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