



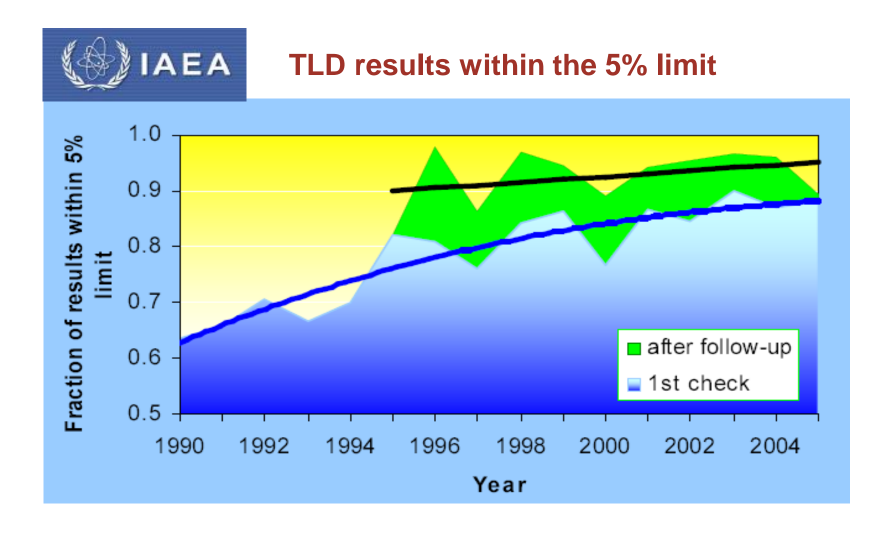
ICTP School on Medical Physics for Radiation Therapy: Dosimetry and Treatment Planning for Basic and Advanced Applications

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Dosimetry: Photon Beams

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Investigation of the quality beam calibration by IAEA Results of Quality Audits



In the following, "dosimetry" means:

- the determination of absorbed dose to water under reference conditions
- in the clinical beam of a radiation delivery unit (accelerator)
- using calibrated ionization chambers.

This is also frequently referred to as beam calibration.

Content:

- 1. Principles of a calibration procedure
- 2. Performance of a calibration procedure
- 3. Correction factors
- 4. Determination of radiation quality Q

1. Principles of the calibration procedure: Need for a Protocol

- Dosimetry protocols or codes of practice state the procedures to be followed when calibrating a clinical photon or electron beam.
- The choice of which protocol to use can be left to individual radiotherapy departments or jurisdictions of individual countries
- Dosimetry protocols are generally issued by national, regional, or international organizations.

1. Principles of the calibration procedure Protocol

Examples of dosimetry protocols

National:

- UK: Institution of Physics and Engineering in Medicine and Biology (IPEMB)
- Germany: DIN 6800-2, Deutsches Institut f
 ür Normung

Regional:

- American Association of Physicists in Medicine (AAPM) for North America: TG-51
- Nederlandse Commissie voor Stralingsdosimetrie (NCS) for Netherlands and Belgium (not used anymore)
- Nordic Association of Clinical Physics (NACP) for Scandinavia (not used anymore)

International:

International Atomic Energy Agency (IAEA): TRS 398

1. Principles of the calibration procedure Protocol

A dosimetry protocol provides three essentials:

- the formalism of determination
- the procedure (methods, prescriptions)
- all the required data, for instance in tables

which have to be used employing a calibrated ionization chamber.

The chamber calibration must be traceable to a standards laboratory for "dosimetry".

1. Principles of the calibration procedure Protocol

Two types of dosimetry protocol are available:

Not addressd in this course !!!

 Protocols based on calibration factors in absorbed dose to water.

IAEA Code of Practice TRS 398 (2000)

Conceptually, both types of protocol are similar and define the steps to be used in the process of determining absorbed dose from a signal measured by an ionization chamber.

1. Principles of the calibration procedure Calibration and calibration coefficient (factor)

Suppose the dose D_{w} is well known at 5 cm depth in a water phantom under so-called calibration conditions:

beam quality	⁶⁰ Co gamma radiation
field size:	10 cm x 10 cm
SSD:	100 cm
phantom:	water phantom
measurement depth in water:	5 cm
positioning of a cyl. chamber:	central electrode at measuring depth

1. Principles of the calibration procedure Calibration under reference conditions

- □ The cylindrical user chamber is then placed with its center at a depth of 5 cm in a water phantom
- ☐ Its calibration factor (or calibration coefficient) $N_{D,w}$ is obtained from

$$N_{D,w,Co} = \frac{D_{W}}{M}$$

where *M* is the dosimeter reading.

Unit: Gray per reading, or Gray per Coulomb

1. Principles of the calibration procedure Measurement at ⁶⁰Co gamma radiation beams

The absorbed dose to water at the reference depth z_{ref} in water for the reference beam of quality $Q_0 = \mathbf{Co}$ and in the absence of the chamber is then simply given by

$$D_{w,Q_O} = M_{Q_O} N_{D,w,Q_O}$$

where

 $M_{Q_{\mathcal{O}}}$

is the reading of the dosimeter corrected for influence quantities to the **reference conditions** as used at calibration

 N_{D,w,Q_0} is the calibration factor in terms of absorbed dose to water of the dosimeter obtained from a standards laboratory.

Example of an Calibration Certificate providing the calibration factor N_{D.w}

Calibration Certificate

000877

Calibration laboratory for ionising radiation quantities

Calibration mark

04-06

Object: Ionization chamber

Manufacturer: Scanditronix Wellhöfer, Germany

CC04 Type: Serial number: 6602

Beam quality:

Absorbed dose to water calibration factor:

Co-60

Measurement uncertainty:

Reference conditions: T₀: 20.0 °C U = 2.2 %

R.H.: 50 %

The reported expanded uncertainty is based on a standard uncertainty multiplied by a coverage factor k = 2, which for a normal distribution provides a level of confidence of approximately 95%.

The secondary standard of this laboratory is traceable to the PTB in Braunschweig (German Federal Institute of Physics

Calibration reported in this certificate was carried out in accordance with the procedures described in the IAEA TRS 398 Code of Practice.

Measuring conditions:

Phantom size

30 cm × 30 cm × 30 cm

Phantom material

water

Source to phantom surface distance (SSD)

100 cm

Field size at the phantom surface

Depth in phantom of the reference point of the chamber

Reference point of the IC:

on the chamber axis at the centre of the cavity volume

Chamber orientation

the beam axis perpendicular to the chamber axis

If the chamber stem has a mark, the mark is oriented towards the radiation source

Waterproof sleeve (PMMA)

Sleeve Serial Number:

Polarizing potential of collecting (central) electrode

1.0 Gy-min

Dose rate

Recombination correction has not been applied

Date of calibration

Head of the Dosimetry Laboratory

Calibration performed by

28.04.2006

1. Principles of the calibration procedure Measurement at other (which means) user qualities

The chamber is now to be used in a beam with a another quality Q such as

- high energy photons
- high energy electrons
 that differs from the ⁶⁰Co quality used in the chamber calibration at the standards laboratory
- Then the formula for the determination of absorbed dose to water is changed
 Beam quality

from
$$D_{W,Qo} = M_{Q_O} N_{D,W,Q_O}$$
 correction factor $D_{W,Q} = M_{Q} N_{D,W,Q_O} K_{Q,Q_O}$

$$D_{w,Q} = M_Q N_{D,w,Q_o} k_{Q,Q_o}$$

is the **chamber reading in beam of quality Q** and corrected for influence quantities to the reference conditions used in the standards laboratory.

 N_{D,w,Q_0} is the water dose calibration coefficient provided by the standards laboratory for reference beam quality Q_0 .

 k_{Q,Q_o} is a factor correcting for the differences between the reference beam quality Q_o and the actual user quality Q_o .

Frequently, the **common reference** quality Q_o used for the calibration of ionization chambers is the cobalt-60 gamma radiation and the symbol k_Q is normally used to designate the beam quality correction factor:

$$K_{Q,Qo} = K_{Q,Co-60} = K_Q$$

How to get the beam quality correction factor k_{Q} ???

- ☐ First choice:
 - An experimentally obtained k_{Q} is available from the calibration laboratory.
- ☐ Second choice:
 - When no experimental data are available, or it is difficult to measure $K_{\mathbb{Q}}$ directly for realistic clinical beams, calculated correction factors can be used.
- ☐ Such calculated correction factors are normally provided in dosimetry protocols.

- \Box General properties of k_{\Box}
 - Values for k_Q are dependent on the quality of radiation (type, energy, machine).
 - Each **type** of ionization chamber needs a particular k_{Q}
 - Values for k_Q are given in protocol tables for a large variety of beam qualities and chambers (e.g.in TRS 398)

TABLE 14. CALCULATED VALUES OF k_Q FOR HIGH ENERGY PHOTON BEAMS FOR VARIOUS CYLINDRICAL IONIZATION CHAMBERS AS A FUNCTION OF BEAM QUALITY $\text{TPR}_{20,10}$ (adapted from Andreo [20])

.	Beam quality														
Ionization chamber type ^a	0.50	0.53	0.56	0.59	0.62	0.65	0.68	0.70	0.72	0.74	0.76	0.78	0.80	0.82	0.84
Capintec PR-05P mini															
Capintec PR-05 mini															
Capintec PR-06C/G															
Farmer															
Exradin A2 Spokas															
Exradin T2 Spokas															
Exradin A1 mini Shonka															
Exradin T1 mini Shonka															
Exradin A12 Farmer															
Far West Tech. IC-18															
FZH TK 01															
Nuclear Assoc. 30-750															
Nuclear Assoc. 30-749															
Nuclear Assoc. 30-744															
Nuclear Assoc. 30-716															
Nuclear Assoc. 30-753															
Farmer shortened															
Nuclear Assoc. 30-751															
Farmer															
Nuclear Assoc. 30-752															
Farmer															

Note:

The quality correction factors k_Q are always and exclusively valid for the

reference conditions of beam calibration

For instance at the reference depth in water.

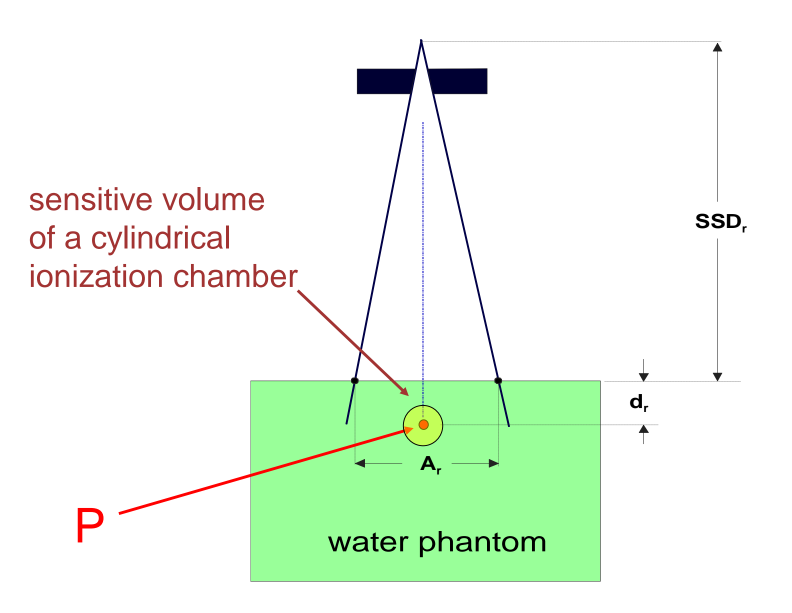
- □ The absorbed dose to water is to be determined at a point P in water at the reference depth z_{ref}.
- ☐ In the absence of the chamber the dose is given by

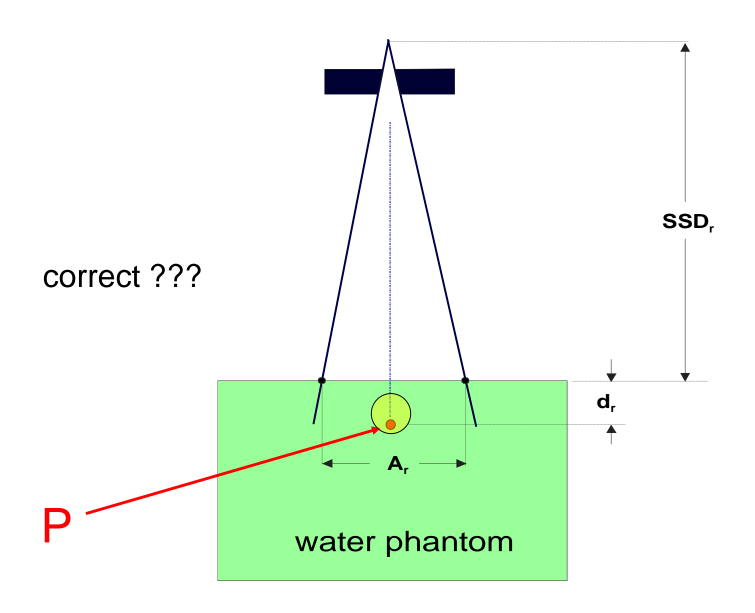
$$D_{w,Q}(P=z_{ref})$$

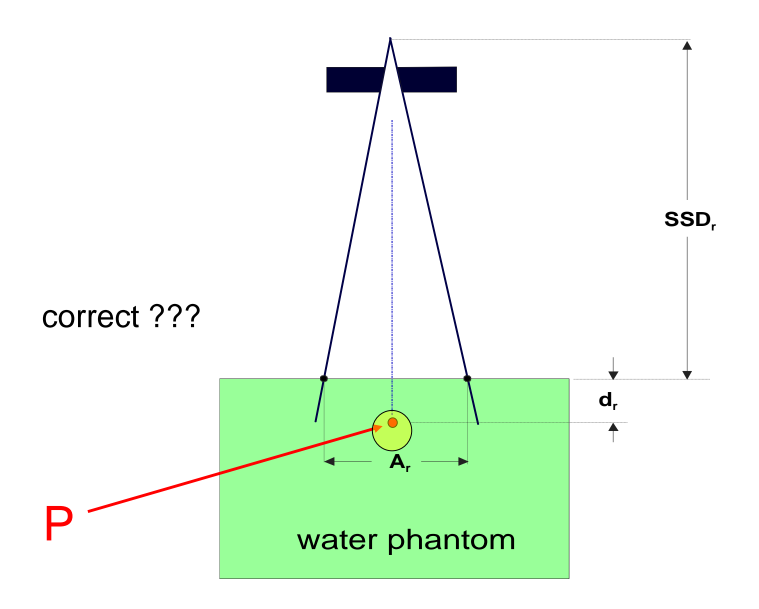
☐ Using the chamber, the dose is given by

$$D_{w,Q}(P) = M_Q N_{D,w,Q_Q} k_Q$$

■ How the chamber must be positioned??





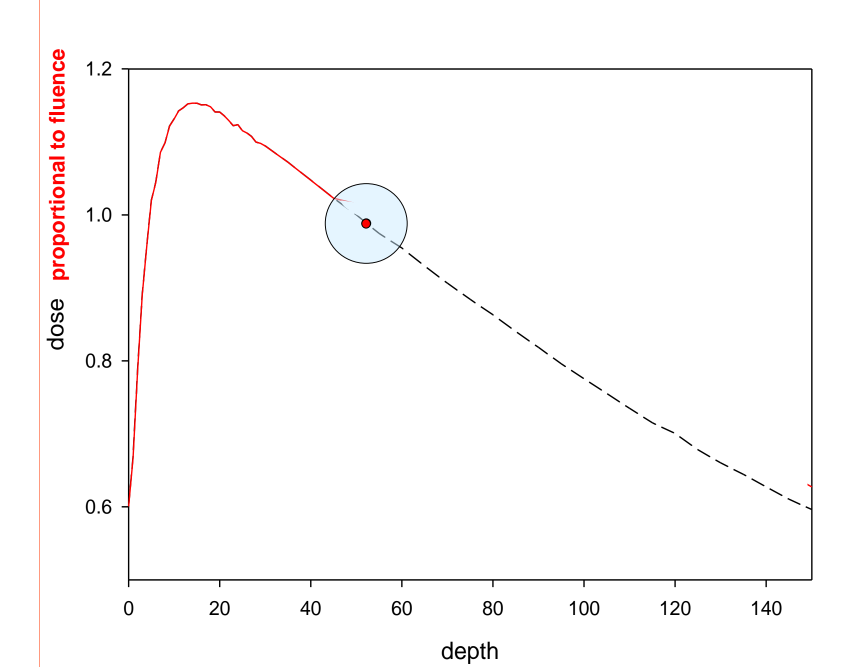


- 2. Performance of a calibration procedure Positioning of the ionization chamber in water
- □ Remember the Bragg-Gray Condition (1):

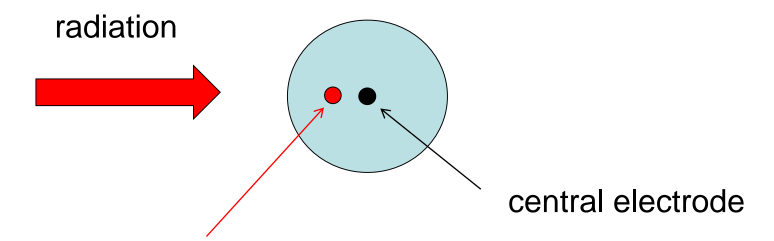
The cavity must be small when compared with the range of charged particles, so that its **presence does not perturb the fluence** of charged particles in the water.

☐ However:

A chamber positioned with its cavity center at the point P does not sample the same electron fluence which is present at P in the undisturbed phantom, i.e. without the chamber.



- Which positioning is correct?
- One may think that the correct way is the positioning of the chamber with its effective point at the reference.



effective point of measurement

However:

"Correct" positioning must strictly follow the prescription in the dose protocol!

There are different prescriptions in different protocols.

There are also different prescriptions in the same protocol for different purposes.

And even more important:

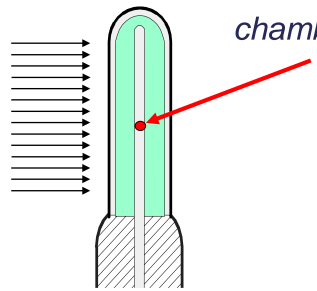
How can the positioning unambiguously be described?

Positioning for the calibration geometry setup:

- Positioning must refer to a well defined point within the chamber which must then be set to coincide with the desired point of measurement.
- This well defined point is the so-called reference point of the chamber.

Positioning of a cylindrical ionization chamber in water

cylindrical chamber



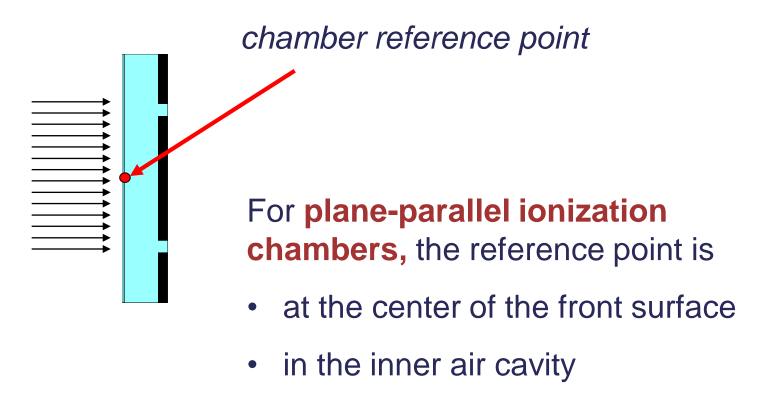
chamber reference point

For **cylindrical chambers** the reference point is

- at the centre of the cavity volume of the chamber
- on the chamber axis.

Positioning of a plane-paralle ionization chamber in water

plane-parallel chamber



Positioning of the ionization chamber in water

Positioning can now be defined as the adjustment of the **reference point** of a chamber with respect to the **measuring depth**.

Positioning of the **reference point** of a **cylindrical chamber** according to the International Code of Practice of the IAEA, TRS 398 and according to the purpose:

	Purpose		
	Beam calibration		
Co-60	at measuring depth		
HE photons	at measuring depth		
HE electrons	Half the radius deeper than at measuring depth		

Positioning of the ionization chamber in water

Positioning can now be defined as the adjustment of the **reference point** of a chamber with respect to the **measuring depth**.

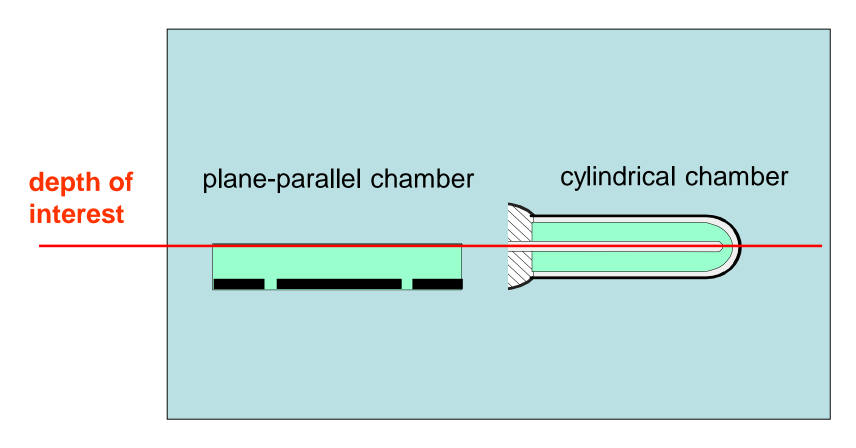
Positioning of the **reference point** of a **cylindrical chamber** according to the International Code of Practice of the IAEA, TRS 398 and according to the purpose:

	Purpose					
	Beam calibration	Depth dose measurement				
Co-60	at measuring depth	0.6 <i>r</i> deeper than measuring depth				
HE photons	at measuring depth	0.6 <i>r</i> deeper than measuring depth				
HE electrons	Half the radius deeper than at measuring depth	0.5 <i>r</i> deeper than measuring depth				

Positioning of the reference point of a **plane parallel chamber** according to the International Code of Practice of the IAEA, TRS 398:

	Purpose				
	Beam calibratio	n Depth dose measurement			
Co-60					
HE photons	always at measuring depth				
HE electrons					

Illustration of positioning for ⁶⁰Co radiation and high energy photons for calibration



2. Performance of a calibration procedure Main procedure

The procedure of a calibration measurement now appears to be quite simple:

- Take an ionization chamber for which a calibration factor from a certificate is available
- Adjust the chamber in the water phantom following the positioning prescription in the protocol
- Obtain charge under reference conditions
- Obtain k_0 from an appropriate look-up table (e.g. protocol)
- Multiply charge reading M, calibration factor and quality correction factor $k_{\rm O}$ to get the absorbed dose to water

$$D_{w,Q} = M_Q N_{D,w} k_Q$$

2. Performance of a calibration procedure Main procedure

There are only two points left to do:

- (1) What exactly means "Obtain charge under reference conditions"?
- (2) We have a lookup table for k_{Q} , but how we get a quantitative value for the quality Q?

We need a procedure to determine a quantitative measure for the beam quality

How to perform a measurement of charge under reference conditions

Why?

The numerical value of the calibration factor $N_{D,w}$ and that of the quality correction factor k_Q are applicable only if the **reference** conditions are fulfilled.

What are the reference conditions? Reference conditions are described by a set of values for influence quantities.

(1) Measurement of charge under reference conditions

Reference conditions for the calibration of ionization chambers

Influence quantity	Reference value or reference characteristic					
Phantom material	water					
Phantom size	30 cm x 30 cm x 30 cm (approximately)					
Source-chamber distance (SCD)	100 cm					
Air temperature	<i>T₀</i> = 20 °C °					
Air pressure	$P_0 = 101.3 \text{ kPa}$					
Reference point of the ionization chamber	for cylindrical chambers, on the chamber axis for plane-parallel chambers on the inner surface of the entrance window,					
Depth in phantom of the reference point of the chamber	5 g cm ⁻²					
Field size at the position of the reference point of the chamber	10 cm x 10 cm					
Relative humidity	50%					
Polarizing voltage and polarity	as in the calibration certificate					
Dose rate	no reference values are recommended but the dose rate used should always be stated in the calibration certificate. It should also be stated whether a recombination correction has or has not been applied and if so, the value should be stated					

(1) Measurement of charge under reference conditions

- In calibrating an ionization chamber or a dosimeter, as many influence quantities as practicable are kept under control.
- However, some influence quantities cannot be controlled, for example air pressure and humidity, and dose rate in ⁶⁰Co gamma radiation.
- If those influence quantities cannot be adjusted to the reference conditions, their departure can be taken into account by applying appropriate correction factors.
- Assuming that influence quantities act independently from each other, a product of correction factors can be applied:

$$M_{\rm Q} = M_{\rm Q}^{raw} \cdot \prod k_{\rm i}$$

where k_i refers to different influence quantities

(1) Measurement of charge under reference conditions

Air temperature and air pressure

- □ T₀ and P₀ are the reference conditions for chamber air temperature (in °C) and pressure.
- T and P are the actual air temperature (in °C) and pressure.
- Then in the user's beam, the correction factor for air temperature and air pressure k_{T,P} is:

$$k_{\mathsf{T,P}} = \frac{(273.2 + T)}{(273.2 + T_0)} \frac{P_0}{P}$$

(1) Measurement of charge under reference conditions

Polarity effect

- Under identical irradiation conditions the use of potentials of opposite polarity in an ionization chamber may yield different readings. This phenomenon is called the polarity effect.
- If the used polarity differs from that at calibration, the following correction factor must be applied:

$$k_{\text{pol}}(V) = \frac{\left| M_{+}(V) \right| + \left| M_{-}(V) \right|}{2M}$$

- M_{+} is the chamber signal obtained at positive chamber polarity
- *M*₋ is the chamber signal obtained at negative chamber polarity
- *M* is the chamber signal obtained at the polarity used routinely (either positive or negative).

(1) Measurement of charge under reference conditions

Polarity effect: Has the calibration laboratory really corrected for the polarity effect ??

Reference conditions: To: 20.0 °C p₀: 101.325 kPa R.H.: 50 % The reported expanded uncertainty is based on a standard uncertainty multiplied by a coverage factor k = 2, which for a normal distribution provides a level of confidence of approximately 95%. The secondary standard of this laboratory is traceable to the PTB in Braunschweig (German Federal Institute of Physics and Metrology) Calibration reported in this certificate was carried out in accordance with the procedures described in the IAEA TRS 398 Code of Practice Measuring conditions: Phantom size 30 cm × 30 cm × 30 cm Phantom material water Source to phantom surface distance (SSD) 100 cm Field size at the phantom surface 10 cm × 10 cm Depth in phantom of the reference point of the chamber 5 a cm Reference point of the IC on the chamber axis at the centre of the cavity volume Chamber orientation the beam axis perpendicular to the chamber axis If the chamber stem has a mark, the mark is oriented towards the radiation source Waterproof sleeve (PMMA) NO Sleeve Serial Number: Polarizing potential of collecting (central) electrode 300 V Dose rate 1.0 Gy min Recombination correction has not been applied

If not then do the following:

- 3. Correction factors
 - (1) Measurement of charge under reference conditions
- If no polarity correction is performed during calibration, it is included in N:
- It follows: If the
 - user beam quality is the same as the calibration quality (normally Co-60)
 - and the chamber is used at the same polarizing potential and polarity as used during the calibration,

then k_{pol} will be the same at **calibration laboratory** and at the **user beam**

Therefore the user must *not* apply a polarity correction for that particular beam.

(1) Measurement of charge under reference conditions

If the user beam quality is not the same as the calibration quality, one should:

- 1) reproduce the calibration quality
- 2) estimate the polarity correction $[k_{pol}]_{Qo}$ that was not applied at the time of calibration using the **same polarizing potential and polarity** as was used at the calibration laboratory.

$$\left[k_{pol}\right]_{Qo} = \frac{\left|M_{+}\right| + \left|M_{-}\right|}{2M}$$

In the same way, the polarity effect at the user beam quality, k_{pol} must be determined.

3. Correction factors Polarity correction factor

The correct polarity correction then is:

$$k_{pol} = \frac{k_{pol}}{[k_{pol}]_{Q_o}}$$

(1) Measurement of charge under reference conditions

There can be a difference between the charge produced by the radiation and actually measured one

- Most important is an incomplete collection of charge in an ionization chamber cavity owing to the recombination of ions.
- Two main separate effects take place:
 - (i) the recombination of ions formed by separate ionizing particle tracks
 - It is termed **general** (or volume) **recombination**, which is dependent on the density of ionizing particles and therefore on the dose rate;
 - (ii) the recombination of ions formed by a single ionizing particle track.
 - It is referred to as **initial recombination**, which is independent of the dose rate.

(1) Measurement of charge under reference conditions

Recombination effect

- In pulsed radiation (i.e. at any linear accelerator!), the dose rate during a pulse is relatively high and general recombination is often significant.
- In the IAEA Code of Practice it is recommended, that the correction factor k_s for pulsed beams be derived using the two voltage method:

$$k_{s} = a_{o} + a_{1} \left(\frac{M_{1}}{M_{2}} \right) + a_{2} \left(\frac{M_{1}}{M_{2}} \right)^{2}$$

- where the values of the collected charges M1 and M2 are measured at the polarizing voltages V1 and V2, respectively.
- V1 is the normal operating voltage and V2 a lower voltage.
- The ratio V1/V2 should ideally be equal to or larger than 3.
- the constants a_0 , a_1 , and a_2 are given in the following slide.

(1) Measurement of charge under reference conditions

Fit coefficients $\mathbf{a_i}$, for the calculation of k_s by the "TWO-VOLTAGE" technique in pulsed radiation, as a function of the voltage ratio V_1/V_2

V_{1}/V_{2}									
	a_o	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	0.975	-0.188	0.214						

Example useful for the Farmer chamber:

Measure charge under reference condition with $V_1 = 400 \text{ V}$: M_1 Measure charge under reference condition with $V_2 = 100 \text{ V}$: M_2

$$k_S = 1.022 - 0.363 \frac{M_1}{M_2} + 0.341 \left(\frac{M_1}{M_2}\right)^2$$

V_{1}/V_{2}			
	a_o	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	0.975	-0.188	0.214

(1) Measurement of charge under reference conditions

In continuous radiation, for instance in ⁶⁰Co gamma rays, the two voltage method may also be used and a correction factor derived using the relation:

$$k_{s} = \frac{(V_{1}/V_{2})^{2} - 1}{(V_{1}/V_{2})^{2} - (M_{1}/M_{2})}$$

Summary:

If the chamber is used under conditions that differ from the reference conditions, then the measured charge must be corrected for the influence quantities by so-called influence correction factors **k**.

The three most import correction factors are:

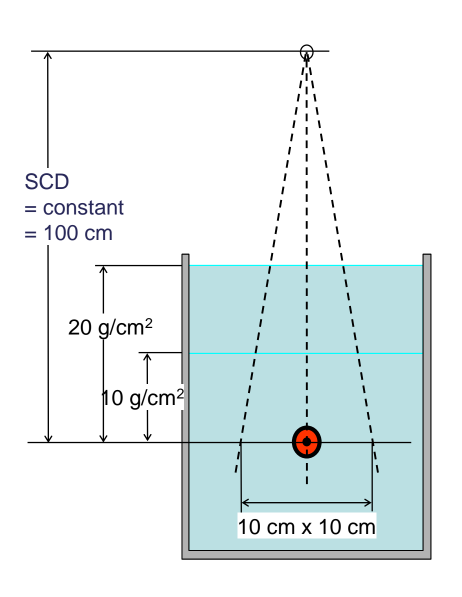
- k_{T.P} for air density
- k_{pol} for polarity effects
- **k**_{sat} for missing saturation effects

- Radiation quality may refer to:
 - a) Low energy X rays with generating potentials up to 100 kV and HVL of 3 mm AI (the lower limit is determined by the availability of standards);
 - **b)** Medium energy X rays with generating potentials above 80 kV and HVL of 2 mm AI;
 - c) ⁶⁰Co gamma radiation;
 - **d) High energy photons** generated by electrons with energies in the interval 1–50 MeV;
 - **e) Electrons** in the energy interval 3–50 MeV;
 - **f) Protons** in the energy interval 50–250 MeV, with a practical range, R_p , between 0.25 and 25 g/cm²;
 - g) Heavy ions with Z between 2 (He) and 18 (Ar) having a practical range in water, R_p , of 2 to 30 g/cm²

- Within each category of radiation type, a particular quantitative parameter, the so-called quality parameter is defined.
- \Box Values of k_Q are tabulated as a function of this quality parameter.
- The selection of the correct value of k_Q therefore requires the determination of the quality parameter.
- The method to determine the quality parameter differs from one radiation type to another.

Definition of the quality index Q for HE photons

- For high energy photons produced by clinical accelerators the beam quality Q is specified by the tissue phantom ratio TPR_{20,10}.
- This is the ratio of the absorbed doses at depths of 20 and 10 cm in a water phantom, measured with a constant SCD of 100 cm and a field size of 10 cm × 10 cm at the plane of the chamber.
- The most important characteristic of the beam quality index TPR_{20,10} is its independence of the electron contamination in the incident beam.



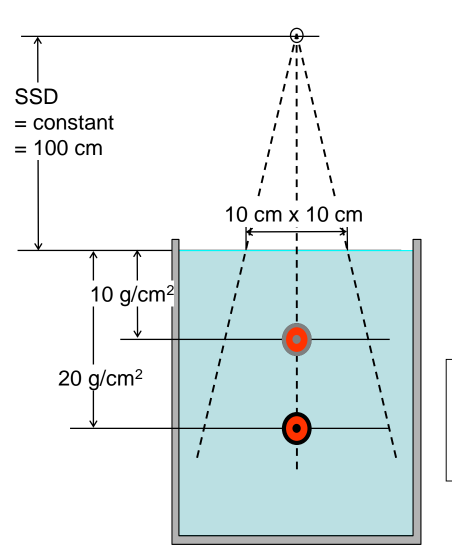
$$\Rightarrow M_{10}$$

$$\Rightarrow M_{20}$$

$$\Rightarrow M_{20}$$

$$\mathsf{TPR}_{20,10} = \frac{D_{20}}{D_{10}} \approx \frac{M_{20}}{M_{10}}$$

Alternative method and easier to perform: the PDD method



$$\Rightarrow M_{10}$$

$$\Rightarrow M_{20}$$

$$PDD_{20,10} = \frac{M_{20}}{M_{10}}$$

$$TPR_{20,10} =$$

$$1.2661 \cdot PDD_{20,10} - 0.0595$$

TABLE 14. CALCULATED VALUES OF k_Q FOR HIGH ENERGY PHOTON BEAMS FOR VARIOUS CYLINDRICAL IONIZATION CHAMBERS AS A FUNCTION OF BEAM QUALITY $\text{TPR}_{20,10}$ (adapted from Andreo [20])

Indication should be to a	Beam Quality TPR _{20,10}														
Ionization chamber type ^a	0.50	0.53	0.56	0.59	0.62	0.65	0.68	0.70	0.72	0.74	0.76	0.78	0.80	0.82	0.84
Capintec PR-05P mini	1.004	1.003	1.002	1.001	1.000	0.998	0.996	0.994	0.991	0.987	0.983	0.975	0.968	0.960	0.949
Capintec PR-05 mini	1.004	1.003	1.002	1.001	1.000	0.998	0.996	0.994	0.991	0.987	0.983	0.975	0.968	0.960	0.949
Capintec PR-06C/G	1.001	1.001	1.000	0.998	0.998	0.995	0.992	0.990	0.988	0.984	0.980	0.972	0.965	0.956	0.944
Farmer															
Exradin A2 Spokas	1.001	1.001	1.001	1.000	0.999	0.997	0.996	0.994	0.992	0.989	0.986	0.979	0.971	0.962	0.949
Exradin T2 Spokas	1.002	1.001	0.999	0.996	0.993	0.988	0.984	0.980	0.977	0.973	0.969	0.962	0.954	0.946	0.934
Exradin A1 mini Shonka	1.002	1.002	1.001	1.000	1.000	0.998	0.996	0.994	0.991	0.986	0.982	0.974	0.966	0.957	0.945
Exradin T1 mini Shonka	1.003	1.001	0.999	0.996	0.993	0.988	0.984	0.980	0.975	0.970	0.965	0.957	0.949	0.942	0.930
Exradin A12 Farmer	1.001	1.001	1.000	1.000	0.999	0.997	0.994	0.992	0.990	0.986	0.981	0.974	0.966	0.957	0.944
Far West Tech. IC-18	1.005	1.003	1.000	0.997	0.993	0.988	0.983	0.979	0.976	0.971	0.966	0.959	0.953	0.945	0.934
FZH TK 01	1.002	1.001	1.000	0.998	0.996	0.993	0.990	0.987	0.984	0.980	0.975	0.968	0.960	0.952	0.939
Nuclear Assoc. 30-750	1.001	1.001	1.000	0.999	0.998	0.996	0.994	0.991	0.988	0.984	0.979	0.971	0.963	0.954	0.941
Nuclear Assoc. 30-749	1.001	1.000	1.000	0.999	0.998	0.996	0.994	0.992	0.989	0.984	0.980	0.972	0.964	0.956	0.942
Nuclear Assoc. 30-744	1.001	1.000	1.000	0.999	0.998	0.996	0.994	0.992	0.989	0.984	0.980	0.972	0.964	0.956	0.942
Nuclear Assoc. 30-716	1.001	1.000	1.000	0.999	0.998	0.996	0.994	0.992	0.989	0.984	0.980	0.972	0.964	0.956	0.942
Nuclear Assoc. 30-753	1.001	1.000	1.000	0.999	0.998	0.996	0.994	0.992	0.989	0.985	0.980	0.973	0.965	0.956	0.943
Farmer shortened															
Nuclear Assoc. 30-751	1.002	1.002	1.000	0.999	0.997	0.994	0.991	0.989	0.985	0.981	0.977	0.969	0.961	0.953	0.940
Farmer															
Nuclear Assoc. 30-752	1.004	1.003	1.001	1.00	0.99	98 0.9	996 0.9	93 0.99	0.9	89 0.98	85 0.9	81 0.9	74 0.967	7 0.959	0.947
Farmer															

Summary: Beam Calibration of Photon Beams TRS 398

1) Calibration formula:

$$D_{w,Q} = M_Q N_{D,w,Q_o} k_{Q,Q_o}$$

2) Follow the positioning instruction of the protocol:

For depth dose measurements:

Position the effective point of the chamber at the measuring depth

For beam calibration measurements:

Position the reference point of the chamber at measuring depth

- 3) The most important correction factors required to meet the reference conditions are:
 - k_{T.P} for air density
 - k_{pol} for polarity effects
 - k_{sat} for missing saturation effects

Summary: Beam Calibration of Photon Beams TRS 398

- 4) The quality correction factor $k_{\mathbb{Q}}$ is given in tables provided in the TRS document, Chapter 6.
- 5) For high energy photons produced by clinical accelerators, the beam quality *Q* is specified by the **tissue phantom ratio TPR**_{20,10} photons produced This parameter can be measured directly or determined by the depth dose methods