Determination of field output factors and volume averaging correction factors for different detectors in narrow Co-60 beams

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UNIVERSITY OF RIJEKA FACULTY OF PHYSICS

Nikola Šegedin

DETERMINATION OF FIELD OUTPUT FACTORS AND VOLUME AVERAGING CORRECTION FACTORS FOR DIFFERENT DETECTORS IN NARROW CO-60 BEAMS

DOCTORAL DISSERTATION

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Hvala svim divnim ljudima koji su bili prisutni kada mi je njihova podrška najviše trebala.

Determination of field output factors and volume averaging correction factors for different detectors in narrow Co-60 beams

Abstract

Accurate small photon field dosimetry is critical for the effectiveness and outcome of the treatment in radiation oncology. Small photon field dosimetry refers to the determination of an absorbed dose in a photon beam whose size is smaller than 3 cm · 3 cm. It is complex due to the finite size of the detector, loss of lateral charged particle equilibrium, and perturbation of secondary electron fluence by the detector. Absorbed dose distributions of a small photon field with nominal field sizes of 16 mm, 8 mm, or 4 mm delivered by Gamma Knife is characterised by a large dose gradient and an ellipsoidal shape. This work aims to experimentally determine detector-specific field output correction factors and measurement uncertainties in such small photon fields for different detectors commonly used for dosimetry of small Co-60 beams delivered by a Gamma Knife. Detector-specific correction factors of active detectors are determined by dividing field output factors of Monte Carlo calculated and EBT3 film measured values with values of active detectors. In addition, volume averaging correction factors and their contribution to a detector-specific field output correction factor were determined. This has been done by the simulation of the detector's geometry and position within the ellipsoid absorbed dose model and integration over the detector's volume. Finally, dose profiles were measured in non-reference geometry. An analysis of measured profiles was performed, and detectors appropriate for such dose profile measurements are recommended.

Keywords: small photon field dosimetry, Gamma Knife, volume averaging, correction factors, field output factors, dose profiles

Određivanje korekcijskih faktora za izlazne faktore polja i korekcijskih faktora na volumno usrednjavanje za različite detektore u uskim fotonskim snopovima Co-60

Prošireni sažetak

Dozimetrija malih polja obuhvaća određivanje apsorbirane doze u uskim snopovima fotona veličine manje od 3×3 cm². Raspodjele apsorbirane doze snopova Co-60 Gamma Knife uređaja karakterizirane su velikim gradijentom doze i elipsoidnim oblikom nominalnih veličina: 16, 8 ili 4 mm. Dozimetrija malih polja je složena zbog veličine detektora, narušenih uvjeta bočne elektronske ravnoteže i utjecaja detektora na tok sekundarnih elektrona. Cilj rada je eksperimentalno odrediti izlazne faktora polja, njihove korekcije u odnosu na referentne vrijednosti kao i odrediti mjerne nepouzdanosti za različite vrste i tipove detektora. Dodatno su određeni korekcijski faktori na volumno usrednjavanja za različite detektore te njihov doprinos u ukupnoj korekciji. To je učinjeno simulacijom geometrije i položaja detektora unutar elipsoidnog modela raspodjele doze. Također, izmjereni su profili doza gama noža različitim detektorima te su uspoređeni s vrijednostima dobivenim Monte Carlo simulacijama. Analizom rezultata preporučen je detektor najpogodniji za određivanje profila u nereferentnoj geometriji.

Izlazni faktori za snopove gama noža određeni su za dvanaest detektora; sedam ionizacijskih komora, četiri poluvodička detektora te jedan dijamantni detektor. Detektori su prilikom mjerenja bili postavljeni u plastičnom (*eng. solid water*) fantomu paralelno sa z-osi Gamma Knife-a. Usporedbom dobivenih rezultata s podacima dobivenim Monte Carlo simulacijama te onih određenih referentnim dozimetrom (radiokromski EBT3 film) utvrđeno je da ionizacijske komore podcjenjuju veličinu izlaznih faktora zbog narušenih uvjeta bočne elektronske ravnoteže, izraženog volumnog usrednjavanja odziva detektora te činjenice da su napravljene

od tkivu ne ekvivalentnog materijala. Sukladno očekivanjima, veličina odstupanja raste s veličinom efektivnog volumena ionizacijske komore. Zbog perturbacija detektora na snop Co-60, odziv detektora je potrebno korigirati korekcijskim faktorom specifičnim za pojedini detektor. Pokazano je da ionizacijske komore Semiflex T31010, Semiflex 3D T331021, PinPoint T31014, PinPoint 3D T3016, RAZOR i CC04, zbog veličine efektivnog volumena, nisu primjerene za dozimetriju u najmanjem polju Gamma Knife-a od 4 mm i njihovi korekcijski faktori su izvan granica prihvatljivosti. S povećanjem veličine polja smanjuje se utjecaj spomenutih perturbacija na točnost određivanja izlaznih faktora. Tako, izlazni faktori RAZORnano ionizacijske komore najmanjeg efektivnog volumena pokazuju najbolje slaganje s referentnim vrijednostima, ali i s rezultatima dobivenim EBT3 filmom. Apsorbirana doza određena na temelju mjerenja silicijevim poluvodičkim detektorom je precijenjena prvenstveno zbog većeg efektivnog atomskog broja silicija u odnosu na vodu. U 8 mm polju, izlazni faktori određeni s ovim detektorima pokazuju statistički značajno slaganje s Monte Carlo vrijednostima. Neki poluvodički detektori imaju zaštitu efektivnog volumena od raspršenih fotona niskih energija te su prvenstveno namijenjeni dozimetriji velikih polja. Pokazano je da se oni također mogu koristiti i u dozimetriji uskih snopova Co-60 kao kod Gamma Knife-a. Izlazni faktori određeni s ovim detektorima pokazuju manje slaganja s Monte Carlo vrijednostima od poluvodičkih detektora bez zaštite. Sintetički jednokristalni detektor se pokazao kao odličan izbor za dozimetriju malih polja gama noža.

Korekcijski faktori volumnog usrednjavanja detektora određeni su primjenom analitičkog elipsoidnog modela raspodjele apsorbirane doze polja Gamma Knife-a te simuliranjem geometrije efektivnog volumena detektora unutar takvih raspodjela. Integriranjem modela apsorbirane doze po efektivnom volumenu detektora izračunata je korekcija na volumno usrednjavanje odziva za četrnaest detektora: šest ionizacijskih komora, pet poluvodičkih detektora, dva plastična scintilacijska detektora te sintetički jednokristalni detektor. U pravilu se korekcijski faktor na volumeno usrednjavanje odziva s volumeno usrednjavanje odziva povećava s volumenom detektora. Međutim, rezultati ranijih istraživanja su pokazala da je osim veličine efektivnog volumena bitan i njegov oblik što je potvrđeno i ovim modelom. Primjerice, ionizacijska komora PinPoint T31014 u usporedbi s ionizacijskom komorom CC04 ima veći korekcijski faktor na volumno

detektora tj. komora CC04 je kompaktnija. Omjer duljine i promjera aktivnog volumena (I/d) CC04 komoru iznosi 0,9 dok za PinPoint T31014 on iznosi 2,5.

Profili doze određeni su dozimetrijski korištenjem različitih vrsta i tipova detektora te su dobiveni rezultati uspoređeni s profilima određeni Monte Carlo simulacijama. Ionizacijske komore većeg volumena nisu primjerene za mjerenja profila doze jer su im efektivni volumeni preveliki za točno mjerenje odziva u području velikih gradijenta doze. S druge strane, sintetički jednokristalni detektor i poluvodički detektori pokazuju manja odstupanja od referentnih vrijednosti te su dobar izbor za određivanje profila doza.

Ključne riječi: dozimetrija malih fotonskih polja, gama nož, volumno usrednjavanje, korekcijski faktori, izlazni faktori, profili apsorbirane doze

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1. Introduction and Motivation

Stereotactic radiosurgery (SRS) with narrow photon beams has started in the early 1950s by the Swedish neurosurgeon Lars Leksell who used a stereotactic frame of his own design to accurately determine a focal point of Cobalt 60 (Co-60) beams to treat small brain tumours and malformations. This has led to the development of the first Leksell Gamma Knife (LGK, Elekta Instruments AB, Stockholm, Sweden). The success of the first Gamma Knife (GK) has led to the creation of clinical radiosurgery accelerators almost simultaneously both in the United Kingdom and the United States of America, and soon spread across the world.^{1,2} From the creation of the first Gamma Knife to the present, over 1 million patients have been treated, with about 80 000 new patients each year, in more than 300 Gamma Knife centres worldwide.^{3,4} With the expected increase in the number of patients diagnosed with cancer, specialized treatments such as stereotactic radiosurgery and stereotactic body radiation therapy (SBRT) will be even more important and used as a method of choice for cancer treatment.^{5–7}

Field sizes of photon beams that traditionally spanned from 3×3 cm² to 40×40 cm² are now being regularly collimated down to 4×4 mm² making most of the available radiation detectors too large for accurate dosimetry. Therefore, the behaviour of detectors in various types of narrow photon fields is still a subject of investigation. Narrow photon beams are commonly used in modern radiation oncology and; therefore, accurate and precise dosimetry of such fields is crucial for the effective delivery of radiation therapy. Studies from the early 2000s have shown a substantial problem in the determination of an absorbed dose in narrow fields.^{8,9} Despite this, the use of narrow and non-standard fields in radiotherapy and stereotactic radiosurgery has increased significantly, creating a need for a unifying approach to standardize and enable accurate delivery of absorbed dose to the target volume while minimizing the exposure to the surrounding tissue. This has led to the collaboration of The American Association of Physicists in Medicine (AAPM) and the International Atomic Energy Agency (IAEA) in creating the first international Code of Practice (CoP), Technical Report Series 483 (TRS-483) dedicated to the dosimetry of small static field that fulfilled the need for a systematic and internationally unified approach to the dosimetry of small static fields.^{10,11}

Leksell Gamma Knife Icon is one of the radiosurgery units that use small photon fields to deliver a high-absorbed dose (up to 100 Gy). High absorbed dose delivery accuracy is achieved with a rigid Tungsten collimator providing three different field sizes: 16, 8, and 4 mm. The 16, 8, and 4 mm field can be related to the respective beam collimator aperture, and these values represent a projection of a diameter for a single collimator hole to the device's isocentre.¹³ In practice, photons are emitted by 192 Co-60 sources and are focused on a single point (isocentre) with a spatial error smaller than 0.3 mm. The resulting absorbed dose is ellipsoidal in shape. For brevity in the further text, the terms 16 mm, 8 mm, and 4 mm field size will be used. For absorbed dose calculation, the determination of field output factors (OF) and dose profiles is of paramount importance since they are input parameters for the calculation algorithm built into the system for calculation and optimization of absorbed dose distribution, also known as treatment planning system (TPS). Field output factor determination of narrow Co-60 beams is demanding due to the steep dose gradients (up to 70 %/mm), loss of lateral charged particle equilibrium (LCPE), perturbation of secondary electron fluence by the difference in detectors and tissue stopping powers, and dose volume-averaging of the detector.^{11–15} This effect is especially pronounced in the 4 mm field size being comparable with the size of detector.^{13,16} Unlike standard phantoms in which measurements are performed, most of the detectors used are not tissue equivalent and there is a perturbation of secondary electron fluence, which affects the reading of the detector. To correct perturbations, a detector-specific correction factor (DSCF) is introduced. The detector-specific correction factor accounts for the volume-averaging effect, beam perturbation due to the non-water equivalence of the detector, and spectral dependence of the photon energy absorption in the medium.^{11,14}

There are three research hypotheses and subsequent objectives of this dissertation:

- Data acquired with a perturbation-free detector (GAFChromic EBT3 radiochromic films, Ashland Advanced Materials, Bridgewater NJ) along with available data determined using Monte Carlo simulation will enable the determination of field output factors and the detector-specific correction factors for detectors not listed in the IAEA TRS-483 CoP.
- Analytically modelled ellipsoidal absorbed dose distribution of narrow Co-60 beams will enable the determination of volume-averaging correction factor for different detectors.

• Analysis of absorbed dose profiles measured with different detectors will provide the data necessary to recommend suitable detectors for dose profile measurements in non-reference geometry.

1.1. Overview of the previous research

In previous research, field output factors and detector-specific correction factors have been determined by Monte Carlo (MC) simulations ^{15–21}, using measured data acquired by different detectors ^{12,14,22–31}, analytically ^{32,33}, or some combination of these methods. ^{13,17,24}

Nowadays, an increasing number of investigations in the field is based on Monte Carlo simulations to determine various physical quantities which affect absorbed dose to quantify detector-specific perturbations.^{16–18,20,34,35} Monte Carlo simulations of particle transport follow discrete steps in the process of interactions with matter using a random number to sample from possible interactions based on probability density functions. They have been giving consistent results throughout different studies and are considered reference techniques for calculating dosimetry quantities. EGSnrc and PENELOPE code have been used for modelling different detectors in narrow photon beams used by Gamma Knife for reference dosimetry both in parallel and perpendicular orientation.¹⁸ Additionally, detectors were modelled inside three different spherical phantoms: Solid Water (Elekta, Stockholm), Acrylonitrile Butadiene Styrene (ABS, Elekta, Stockholm) and liquid water. Detector-specific correction factors for all investigated detectors were within 3% of the reference (MC) value and all had the smallest correction in a liquid water phantom and parallel orientation. Similar Monte Carlo studies support these conclusions.¹⁵

Furthermore, GAFChromic EBT3 films and diamond detectors were used to validate penEasy Monte Carlo code results. The relative dose profiles along the X and Z axis, for each investigated collimator size and field output factors, have been investigated and compared against the values acquired with the diamond detector. Results were in good agreement with the Monte Carlo simulation.¹⁷ The maximum difference in OF was 2.1% for the 4 mm collimator. The results of the EBT3 dosimetry and penEasy calculation were in excellent agreement. The difference was 0.4% for the 8 mm, and 0.7% for the 4 mm collimator.

Likewise, field output factors and detector-specific correction factors were investigated for a range of different detectors using PEGASOS, a Monte Carlo system based on the PENELOPE code.³⁶ The liquid ionization chamber (V=1.71 mm³) had a correction factor close to unity for all beams. Six air-vented ionization chambers (volume from 15.71 mm³ to 23.75 mm³) yield

detector-specific correction factors larger than 1.3. A large correction is related to volumeaveraging due to the size of the detector's effective volumes and thus are not recommended for dosimetry of Gamma Knife's photon beams. Investigated silicon diode detectors had correction of field output factors between 0.95 and 1.0 mainly contributed to the non-tissue equivalence of the detector. Using the Monte Carlo chain technique, it is possible to determine each detector-specific perturbation inside the photon beam. It has been shown that volume-averaging has a dominant contribution to overall correction for most of the studied detectors.³⁷

Experimental methods of determination of field output factors and detector–specific correction factors are performed with various detectors. ^{8,12–14,17,22,24,37–39} Field output factors are determined using detectors that can be considered perturbation-free (e.g. Exradin W1/W2, EBT3, etc.), and field output factors of studied detectors are compared in a way that the detector's specific correction is determined.

Air-vented ionization chambers are used in radiation dosimetry due to their high sensitivity, dose rate independence, low directional independence, and abundance of dosimetry data for these detectors. ^{8,13,33,37,40} However, in small photon fields, they have displayed a reduction in sensitivity, perturbation of secondary electron fluence, lower spatial resolution, and larger volume-averaging effect.¹³ Detector-specific correction factors for some ionization chambers can be larger than 1.300 and thus are not recommended for small photon field dosimetry. ¹⁴ Recently developed small-volume ionization chambers have an improved spatial resolution but require corrections up to 1.050. ¹⁴

Diode detectors are used in small photon field dosimetry due to higher sensitivity and better spatial resolution than ionization chambers but have some disadvantages such as dependence of response on dose rate, energy, and direction.³³ It is known that silicon diode detectors overrespond due to low-energy scattered photons.^{1,13,14,29} Depending on the size and design, differences in detector-specific field output factors and field output factors from the reference values can be as large as 10% in small fields.^{14,23,41}

Radiochromic film has an excellent spatial resolution. It is considered tissue equivalent and shows minimal directional dependence. ^{17,42,43} It is widely considered a perturbation-free detector. Field output factors determined using this type of detector show excellent

agreement with Monte Carlo simulations.^{17,24,26,30} A disadvantage of radiochromic film is its long saturation time (more than 48 hours), lower signal-to-noise ratio (SNR), and demanding postirradiation processing.

Diamond detectors are solid-state detectors whose effective volume is made of natural or synthetic diamond. They are water equivalent and suitable for field output factors determination and dose profile measurements due to their small size, high response, and low directional independence.^{12,13,29} Field output factors and detector-specific correction factors determined with a diamond detector have shown a discrepancy with Monte Carlo simulation data of 1.5% and 0.2% for 4 and 8 mm field sizes, respectively.⁴⁴

Plastic scintillator detectors (PSD) are a new type of detector with attractive properties like water equivalency, high spatial resolution, energy and dose rate independence, and linear dose response. The main disadvantage of PSDs is the production of Cherenkov light which contaminates the main signal. Research with PSDs on Gamma Knife has been very limited so far.⁴⁵

Researchers also use a range of different detectors, some of which have not been stated so far (e.g. alanine, TLDs, MOSFETs, etc.). In general, handling these detectors can be challenging, and they may have less desired properties (e.g., large DSCF, instability, directional dependence, etc.).^{12,13,46}

Finally, different mathematical tools are used to model the response of the detector and correct the perturbations e.g., using detectors of different sizes and extrapolating data to zero (point) size, determining the convolution kernel of detectors response, characterizing the influence of the physical properties of the detector, correcting for angular dependence etc.^{13,33,37,40}

Generally, according to recent literature, more studies of detector-specific correction factors and field output factors of Gamma Knife beams are based on Monte Carlo simulations than the experimental approach, which may indicate the direction of research in small photon field dosimetry. Some of the problems with the lack of Monte Carlo data come from the complex geometry of the Gamma Knife collimating system. With the advances in computing power, newer simulations tend to be more detailed and accurate. So far, an estimate of volume-

averaging of Co-60 beams used by Gamma Knife has been done once by printing the organic PSD detector in the shape of individual voxels.⁴⁷ Although ingenious, this solution is impractical since it requires experience and equipment for high-precision 3D printing. Moreover, such detectors are not standardized in design or response and lack an adequate research base. With this in mind, investigation of the behaviour of the detectors in small photon fields is an ongoing process.

1.2. Outline of the thesis

The thesis is based on three goals; the determination of field output factors and volumeaveraging correction factors for different detectors commonly used for small field dosimetry, coupled with dose profile measurements of Gamma Knife's narrow photon beams in nonreference geometry.

In Chapter 1 "Introduction and Motivation", a brief historical development and overview of the status of small photon field dosimetry is presented. Special emphasis was given to the field output factor determination as well as the current knowledge overview of the behaviour and performance of different detectors in small photon fields. The three hypotheses and subsequent objectives are stated.

In Chapter 2 the physics of small photon field dosimetry is introduced, and the main problems are discussed. The focus is on the main effects such as loss of charged particle equilibrium, partial source occlusion and volume-averaging of the measured signal. Since these problems are not exclusive to the dosimetry of Gamma Knife's narrow photon beam, they are discussed as general problems. The second chapter was finished with the concepts and formalism introduced by IAEA TRS-483 CoP, which will be followed through this dissertation.

Research methods and materials are presented in Chapter 3. A Gamma Knife Icon device used for Stereotactic radiosurgery is introduced. The design and capabilities of this device are discussed. Next, the list of detectors that are used in this research, as well as their design, is presented. The detector's capabilities and physical working principles are explained in more detail. For the calculation of the volume-averaging correction factors, a formalism of an ellipsoid absorbed dose model is introduced. At the end, statistical methods used for the uncertainty analysis are presented.

The experimental setup for field output factor and dose profile determination is presented in Chapter 4. Volume-averaging correction factor calculated using the ellipsoid absorbed dose model is explained along with the main characteristics and problems with the Gamma Knife's dose profiles determination.

In Chapter 5., field output factors, volume-averaging correction factors and dose profile measurements along with the results of uncertainty analysis are presented and discussed.

At the end, in Chapter 6., the main points of this research are summarized and concluded.

In Appendix A, schematics used for modelling the ionization chambers for volume-averaging calculations are presented. These schematics are provided by the vendor, PTW Freiburg and IBA Dosimetry.

Finally, in Appendix B, a MATLAB code for volume-averaging determination is provided. The MATLAB code can be used and shared by other researchers without any limitations.

2. Physics of a small field dosimetry

Dosimetry of broad high-energy photon beams used in radiotherapy is simplified by the fact that a large part of an absorbed dose distribution can be considered as a region where the requirement on charge particle equilibrium (CPE) is fulfilled (Fano's theorem). Therefore, measurements in such regions are well-established and accurate.^{2,8,13,48} Small photon fields differ from broad beams in important dosimetry properties, named and explained in the next chapter, making most of the commonly used detectors large relative to the radiation field size.¹⁰ Dosimetry of such fields refers to the measurements and characterization of an absorbed dose delivered by the photon fields smaller than 3×3 cm². The dosimetry of a small photon fields is challenging because of the presence of steep dose gradients and the lack of charged particle equilibrium making experimental procedures demanding. In the present chapter, the specific physical aspects of small photon field dosimetry are summarized.

Three physical conditions make a high-energy photon field small, and at least one of them must be fulfilled:

- 1. There is a loss in the lateral charge particle equilibrium (LCPE) on the beam axis.
- 2. There is a partial occlusion of the primary photon source on the beam's central axis due to the collimation process.
- 3. The detector's size is similar or large compared to the beam dimensions.

2.1. Charge particle equilibrium

The charge particle equilibrium (CPE) condition is fulfilled when the number of charged particles leaving the volume is equal to the number entering it, both for energy and type of particle. This is an important property since detectors, used for the absorbed dose determination, work under the assumption of the CPE, and the ratio of the absorbed dose to kinetic energy released per unit mass (collision KERMA, K_C) D/K_C is used to determine a degree of CPE. When the absorbed dose is equal to collision KERMA in a certain volume, the CPE condition is fulfilled. Lateral charge particle equilibrium range r_{LCPE} is the minimum radius of a circular field for which the absorbed dose is equal to collision KERMA at the centre of a field. If the half-width (radius) of a beam is smaller than the maximum range of secondary electrons there will be a loss of lateral charge particle equilibrium. Violation of the CPE creates difficulty in the accurate determination of an absorbed dose-to-water for non-water equivalent material.^{10,19,49} A violation of LCPE in narrow photon beams is illustrated in Figure 1. b).

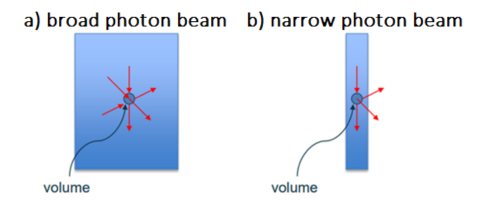
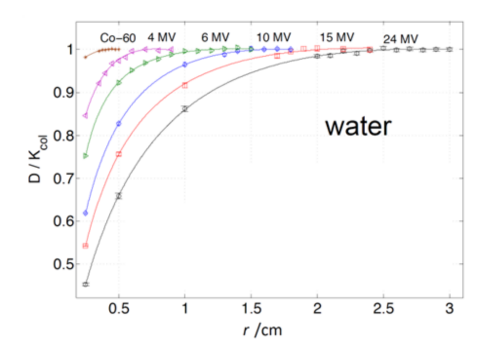


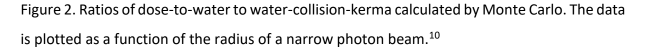
Figure 1. Charge particle equilibrium in a) broad photon beam, and b) loss of lateral charge particle equilibrium in narrow photon beam. Red arrows represent secondary electrons entering and leaving the volume.

This condition has been quantified using Monte Carlo by evaluating the necessary beam radius in centimetres for different energies to achieve CPE at the beam axis (Figure 2.) and is expressed in terms of a tissue-phantom ratio (TPR) for a 10×10 cm² field as:¹⁰

$$r_{LCPE} = 8.369 \cdot TPR_{20.10} - 4.382$$
 2.1

Where $TPR_{20,10}$ is the ratio of an absorbed dose at depth 20 cm and 10 cm. With the increase in the energy of a beam, the radius necessary for achieving LCPE also increases due to the larger mean energy of secondary electrons and as a result more penetrating power. This dependence is illustrated in Figure 2. for different photon beams.





For accurate dosimetry of small fields, for a given beam quality, the distance from the detector's outer boundary to the radiation field edge must be smaller than r_{LCPE} . This will ensure charged particle equilibrium inside the detector's effective volume because any electron that left the volume will be replenished by an electron produced outside of the volume.

2.2. Partial source occlusion

A partial source occlusion is created by the collimator which shields part of the finite primary photon source. This is illustrated in Figure 3., and it becomes important when the field is comparable in size with the primary photon source i.e. when the size of the radiation beam is equal to or less than the full width at half maximum (FWHM) of the emission spectra emitted by the source.

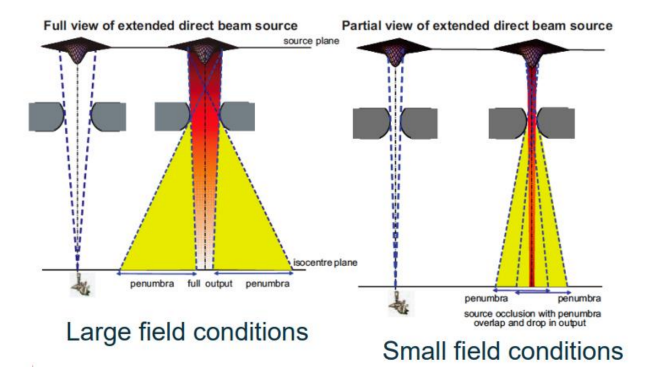


Figure 3. Schematic illustration of a partial source occlusion effect. ⁴⁹ Note: Distance from source to the isocentre plane is the same in both cases.

This effect changes the emission spectrum, and it is a cause of high-dose gradients which can have a significant effect on the detector's response.^{10,49,50} Since the source size is in general smaller than 5 mm, and LCPE begins at the same beam radius (Figure 2.), both loss of charged particle equilibrium and partial source occlusion appeared for the beams with a radius smaller than 5 mm. These effects result in the lower beam output and it is the main reason why field output factors decrease rapidly with a reduction in beam dimensions.² This drop becomes more pronounced with the increase of the photon beam energy or as the density of a medium decreases due to the increase in range of the secondary electron .^{10,19,49–51} Effects of the partial source occlusion can be summarized as overlapping of penumbras, reduction of a central-axis dose which decreases the beam output, and finally, apparent field widening causing a mismatch between true field size (FWHM) and nominal field size determined by collimator settings.⁵⁰ The effects of the partial source occlusion are illustrated in Figure 4.

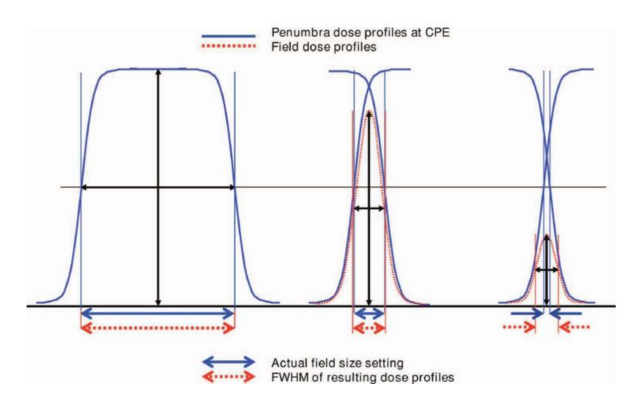


Figure 4. Effects of overlapping penumbras on the FWHM of the lateral beam profile for small fields illustrating the apparent field widening compared to the collimator settings and reduction of the central axis dose.^{9,10}

Although an important effect in small photon field dosimetry, the effect of overlapping penumbras is negligible on Gamma Knife since Co-60 sources are packed as small cylinders with a diameter of 1 mm, while the smallest beam diameter is 4 mm.

2.3. Volume-averaging

Volume-averaging of a measured response is another problem in small photon field dosimetry. A detector produces a response that is proportional to the mean absorbed dose over its effective volume and the response is affected by the homogeneity of the absorbed dose distribution over the volume of the detector. As a result, the response is averaged over the volume of the detector. ¹⁰

Volume-averaging leads to two distinct consequences:

- 1. The absorbed dose at the centre of a small field is underestimated.
- 2. The penumbra is flattened i.e. the field appears to be larger than it is.

An illustration of the volume-averaging problem is shown in Figure 5. with a schematic representation of the volume-averaging effect in the determination of the Gaussian beam profile using a detector 5 mm in length. Both underestimation of the absorbed dose at the central axes of the beam and flattening of the penumbra is visible. This effect is the subject of research primarily concentrated on X-ray beams produced by linear accelerators, CyberKnifes, and Tomotherapy units ^{10,23,32,50–53}, while research regarding volume-averaging in narrow Co-60 beams of Gamma Knife is very limited.^{47,54} It has been shown that volume averaging, besides the size of the detector's effective volume, depends on the shape of that volume as well. More compact detectors with a lower length-to-diameter ratio (I/d) will have lower volume averaging correction for the same size of an effective volume.⁵⁵

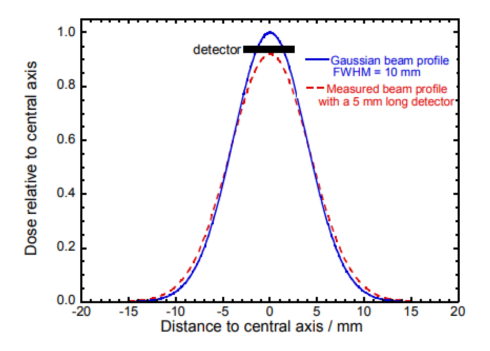


Figure 5. Schematic representation of volume-averaging effect one dimension. The blue curve is a Gaussian dose profile, the red dashed curve represents what a detector of 5 mm length measures.⁵⁰

In practice, it is impossible to eliminate volume-averaging. For field output factors determination, volume-averaging will be one of the limiting factors for the choice of detectors, especially in narrow photon beams of Gamma Knife. The detectors should be chosen in a way that its volume-averaging correction factor $(k_{vol})_{Q_{clin}}^{f_{clin}}$, for the beam quality Q_{clin}, in a clinical (clin) field of interest f_{clin}, is such that it is limited by $0.95 \leq (k_{vol})_{Q_{clin}}^{f_{clin}} \leq 1.05$.¹⁰ The volume-averaging correction factor can be calculated using the Kawachi formula⁵³ by integrating dose profiles over the detector's area A perpendicular to the beam axis:

$$(k_{vol})_{Q_{msr,clin}}^{f_{msr,clin}} = \frac{\iint_{A} w(x,y)dxdy}{\iint_{A} w(x,y)OAR(x,y)dxdy}$$
 2.2

where, x and y are the coordinates of the axes orthogonal to the beam's central axis, w(x, y) is the weighting function specific to the detector's geometry and represents the extension of the air cavity along the beam axis as a function of the beam lateral coordinates. In the Equation 2.2. msr and clin stand for machine-specific reference field and clinical field, respectively. For

most ionization chambers, this can be approximated with unity.⁴⁸ Finally, OAR(x, y) is the offaxis ratio or 2D lateral beam profile at the measuring depth normalized to the central axis.

The Equation 2.2. can be used to determine volume-averaging which will affect the detector's response in dosimetry of narrow X-ray beams. In these devices the beam is incident perpendicular to the detector surface, therefore, to determine volume-averaging one must integrate the field over the detector's surface perpendicular to the beam's central axis. However, on Gamma Knife Perfexion there are 192 beams converging to an isocentre as illustrated in Figure 6. The ellipsoid geometry of the Gamma Knife absorbed dose distribution complicates the calculation of the volume-averaging in Gamma Knife's narrow photon beams is limited even when it comes to the Monte Carlo simulations. However, if done realistically, Monte Carlo simulations should be the best tool for volume averaging correction factor estimations.

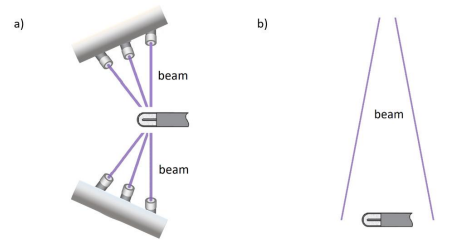


Figure 6. A schematic representation of radiation field for a) Gamma Knife Perfexion, and b) LINAC, CyberKnife, or Tomotherapy unit.

Due to the unique geometry of Gamma Knife's collimator system, Kawachi formalism must be modified from 2D to 3D to be applicable for absorbed dose distribution of Gamma Knife's narrow photon beams, and integration must be performed over the detector's effective volume, in a way:

$$(k_{vol})_{Q_{clin}}^{f_{clin}} = \frac{\iiint_{V} w(x, y, z) dx dy dz}{\iiint_{V} w(x, y, z) OAR(x, y, z) dx dy dz}$$
2.3

The effective volume of most available detectors is cylindrical. Therefore, integration was performed in the cylindrical coordinate system, with the assumption that w(x, y, z) = 1:

$$(k_{vol})_{Q_{clin}}^{f_{clin}} = \frac{\iiint_{V} sdsd\varphi dz}{\iiint_{V} OAR(s,\varphi,z)sdsd\varphi dz} = \frac{V}{\iiint_{V} OAR(s,\varphi,z)sdsd\varphi dz}$$
2.4

where V is detector's effective volume and s, φ and z are the coordinates in cylindrical coordinate system. Neglecting the weighting function can be justified by the fact that its impact on volume-averaging correction for a 2D case is approximately 0.1 %.¹⁰

The volume-averaging correction factor k_{vol} is used for a conversion of a mean dose D_{mean} determined by a detector to a point dose:

$$D_P = D_{mean} \cdot (k_{vol})_{Q_{clin}}^{f_{clin}}$$
 2.5

if the geometry of the detector is known, as well as the relative dose distribution over the detector's volume the volume-averaging correction factor can be calculated as:

$$(k_{vol})_{Q_{clin}}^{f_{clin}} = \frac{D_p}{D_{mean}} = \frac{D_p}{\frac{1}{V} \int D(\vec{r}) d^3 r} = \frac{V}{\int \left(\frac{D(\vec{r})}{D_p}\right) d^3 r} = \frac{V}{\int D_{rel}(\vec{r}) d^3 r}$$
 2.6

Using Equation 2.6. and by knowing the shape and the value of the detector's effective volume V and a normalized absorbed dose distribution over that volume $D_{rel}(\vec{r})$, a volume-averaging correction factor for the detector can be calculated.

2.4. Concepts and formalism in small photon field dosimetry

In this section, concepts and formalism for reference and relative dosimetry of narrow Co-60 beams are presented according to the IAEA TRS-483 CoP.¹¹ Reference dosimetry refers to a determination of an absorbed dose in water with an ionization chamber for which the charge is measured in the beam of the user's institution. The reference conditions used in the calibration laboratory must be reproduced, and influence quantities are measured at the time of data acquisition and the measured detector's response is corrected accordingly. In the clinical environment, various measurements are performed under non-reference conditions. These measurements are called relative.⁵⁶ For irradiation units that cannot attain a reference field, a machine-specific reference field is introduced. The machine-specific reference field has dimensions as close as possible to the conventional reference field and extends at least an r_{LCPE} distance from the outer boundary of the ionization chamber.¹⁰ That is, the size of the detector *d*, and the field size FWHM must fulfil the condition:

$$FWHM_{msr} \ge 2r_{LCPE} + d$$
 2.7

which is illustrated in Figure 7. where the yellow area represents the range of secondary electrons, and the blue is a square field over the detector's effective area.

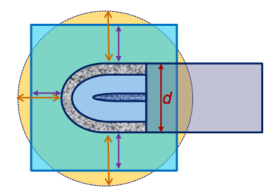


Figure 7. The illustration of a range of secondary electrons leaving the ionization chamber (yellow) and a small field superposed on the ionization chamber (blue). Charge particle equilibrium is violated.¹

In the majority of cases, the ionization chamber calibration factor in terms of an absorbed dose to water $N_{D,w,Q_0}^{f_{ref}}$ is determined in a 10×10 cm² reference field f_{ref} with beam quality Q₀. For Gamma Knife such a reference field is impossible to achieve and therefore a machine-specific reference field f_{msr} (msr) is used which is the largest 16 mm field. In such conditions, the absolute dose to water $D_{w,Q_{msr}}^{f_{msr}}$ is determined as:

$$D_{w,Q_{msr}}^{f_{msr}} = M_{Q_{msr}}^{f_{msr}} \cdot N_{D,w,Q_0}^{f_{ref}} \cdot k_{Q_{msr},Q_0}^{f_{msr},f_{ref}}$$
2.8

where $M_{Q_{msr}}^{f_{msr}}$ is the reading of the ionization chamber in a machine-specific reference field corrected for pressure, temperature, incomplete charge collection, polarity humidity etc. The beam quality correction factor $k_{Q_{msr},Q_0}^{f_{msr},f_{ref}}$ accounts for the difference between the response of the ionization chamber in the reference field f_{ref} with beam quality Q₀, and a machine-specific reference field f_{msr} with beam quality Q_{msr}, and it is defined as a ratio of ionization chamber's calibration coefficients in a machine-specific reference field and conventional reference field (Equation 2.9.).¹¹

$$k_{Q_{msr},Q_{0}}^{f_{msr,f_{ref}}} = \frac{N_{D,w,Q_{msr}}^{f_{msr}}}{N_{D,w,Q_{0}}^{f_{ref}}} = \frac{D_{w,Q_{msr}}^{f_{msr}}/M_{Q_{msr}}^{f_{msr}}}{D_{w,Q_{0}}^{f_{ref}}/M_{Q_{0}}^{f_{ref}}}$$
2.9

For most of the ionization chambers, Q_0 is the beam quality of the Co-60 beams. The beam quality correction factor is usually calculated using Monte Carlo simulations.¹⁰ For some ionization chambers that can be used for reference dosimetry of the Gamma Knife Icon beams, they are calculated and listed in IAEA TRS-483 CoP.

Field output factors $\Omega_{Q_{clin},Q_{msr}}^{f_{clin},f_{msr}}$ are measures of the difference of an absorbed dose in clinical fields compared to the machine-specific reference field. They are defined as a ratio of the absorbed dose to water in the clinical field f_{clin} with beam quality Q_{clin} the absorbed dose to water in machine-specific reference field f_{msr} with beam quality Q_{msr} (Equation 2.10.)

$$\Omega_{Q_{clin},Q_{msr}}^{f_{clin},f_{msr}} = \frac{D_{w,Q_{clin}}^{f_{clin}}}{D_{w,Q_{msr}}^{f_{msr}}}$$
2.10

To determine an absorbed dose, a response is measured in the form of an electric charge produced in the detector's effective volume. In the broad beams, a collected charge is proportional to the absorbed dose. However, in small fields due to the nonequilibrium conditions, the detector's response ratio must be multiplied by the detector-specific correction factor $k_{Q_{clin},Q_{msr}}^{f_{clin},f_{msr}}$ to determine field output factors (Equation 2.11.).

$$\Omega_{Q_{clin},Q_{msr}}^{f_{clin},f_{msr}} = \frac{M_{Q_{clin}}^{f_{clin}}}{M_{Q_{msr}}^{f_{msr}}} \cdot k_{Q_{clin},Q_{msr}}^{f_{clin},f_{msr}}$$
2.11

The detector-specific correction factor can be determined experimentally by measurements or calculated using Monte Carlo. This correction factor is specific to a given detector and must be determined for each field size, energy, and spectral distribution. For some detectors that are very small and have an energy-independent response, like radiochromic films, liquid ionization chambers, or an organic scintillator; the detector-specific correction factor is very close to unity.^{2,8,10,29,49,51} The detector-specific correction factor is a product of four different perturbations in the small fields: ratios of water-to-detector-medium stopping power in clinical and machine-specific reference fields $(s_{w,det})_{Qmsr}^{Q_{clin}}$, volume-averaging correction factor ratios $(p_{vol})_{Qmsr}^{Q_{clin}}$, ratios due to the fluence perturbations $(p_f)_{Qmsr}^{Q_{clin}}$, and, finally, ratios of spectral dependence of photon energy absorption inside the detector medium in clinical and machine specific reference field $(p_{spec})_{Qmsr}^{Q_{clin},14}$

$$k_{Q_{clin},Q_{msr}}^{f_{clin},f_{msr}} = \frac{(s_{w,det})_{Q_{clin}}}{(s_{w,det})_{Q_{msr}}} \cdot (p_{vol})_{Q_{msr}}^{Q_{clin}} \cdot (p_f)_{Q_{msr}}^{Q_{clin}} \cdot (p_{spec})_{Q_{msr}}^{Q_{clin}}$$
2.12

It has been shown that fluctuation of stopping power with beam size is negligible e.g. changing the beam dimensions from 10×10 cm² to 0.5×0.5 cm² the stopping power changes only by 0.3%.^{14,21} Therefore, Equation 2.12. can be approximated as:

$$k_{Q_{clin},Q_{msr}}^{f_{clin},f_{msr}} \approx (p_{vol})_{Q_{msr}}^{Q_{clin}} \cdot (p_f)_{Q_{msr}}^{Q_{clin}} \cdot (p_{spec})_{Q_{msr}}^{Q_{clin}}$$
2.13

A summary of formalism for small static photon field dosimetry is presented in Figure 8.

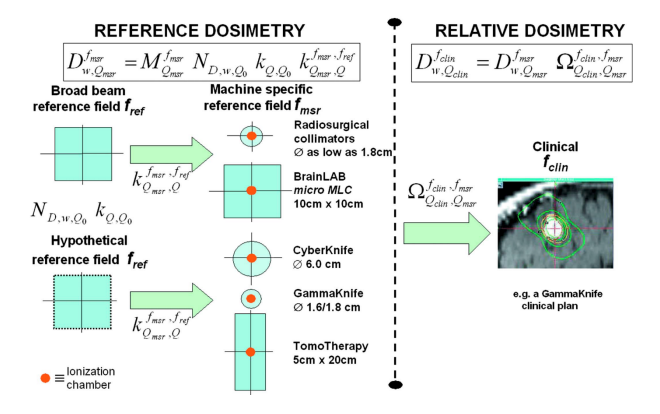


Figure 8. Schematic overview of the dosimetry of small static fields with reference to machinespecific reference fields according to Alfonso formalism and IAEA TRS-483 CoP.^{10,11}

3. Research Methods and Materials

3.1. Gamma Knife Perfexion - Icon

Leksell Gamma Knife Icon (Elekta, Stockholm, Sweden) is a model of Gamma Knife built on the Perfexion platform with the addition of Cone Beam Computed Tomography (CBCT). This model differs substantially from its predecessor, and it is the first time since Model C that its beam delivery system has been completely redesigned with 192 Co-60 sources arranged in a cone-shaped configuration compared with 201 sources in the hemispherical configuration in previous models (Figure 9.).

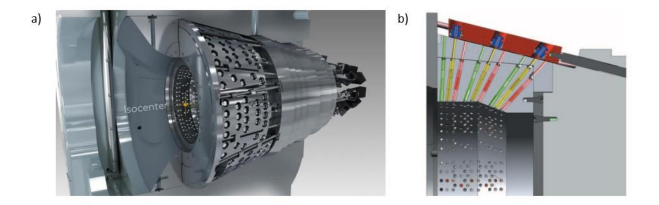


Figure 9. a) Collimator system of Gamma Knife Perfexion and b) its cross-section.

Radionuclide Co-60 decays with β^{-} decay into Ni-60 with the emission of two photons 1.17 MeV and 1.33 MeV. Since the probability for the emission of each photon is approximately equal, the mean energy of Co-60 decay is 1.25 MeV. All photons are focused to a single point - isocentre. The total absorbed dose is equal to the sum of absorbed doses delivered by each beam allowing dose escalation at the isocentre (up to 100 Gy) creating strict requirements on the geometrical and dosimetry accuracy. LGK absorbed dose distribution is ellipsoidal in shape due to the geometry of the collimating system.

On Gamma Knife Perfexion there are three available collimators: 16 mm, 8 mm, and 4 mm. The collimator array is made of a single 120 mm tick tungsten divided into eight identical regions, every with 72 collimators (24 for each of the 3 available collimators). Nominal fields are named 16 mm, 8 mm and 4 mm due to the size of the projection of a beam diameter through a single collimator hole to the isocentre. Gamma Knife absorbed dose distributions are superposition of 192 beams and therefore the actual field size is larger. Different field sizes are created by the automated sliding of Co-60 sources over the top of the collimator system (Figure 9b.).⁵⁷

Depth dose modelling is performed with the new TMR 10 water-based dose calculation algorithm (Elekta Instruments AB, Stockholm, Sweden).⁵⁸ TMR 10 calculates the absorbed dose at a point using two different attenuation parameters; first for the attenuation of the primary photon fluence and the second term where the distance from the focal point is multiplied by the virtual attenuation coefficient of the particular beam. In clinical practice, dose distributions often have to be multicentric and irregular in shape.

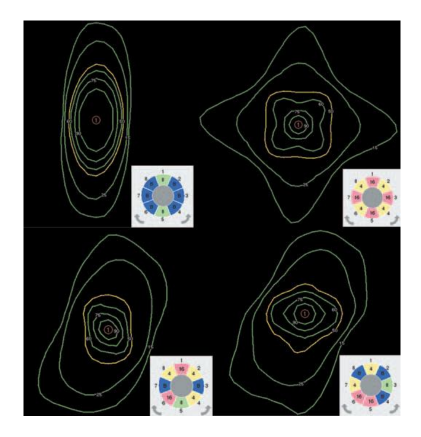


Figure 10. Examples of isodose shapes from single isocentre using composite shot features.⁵⁹



Figure 11. Leksell Gamma Knife Icon.³

3.2. Detectors

Detectors that were used for the determination of field output factors, volume-averaging correction factors and dose profile measurements are presented in Table 1.^{60–66}

Listed air–vented ionization chambers range in size of their effective volumes from 3 mm³ to 125 mm³. Some of the ionization chambers whose dosimetry properties have been investigated in this research do not have detector-specific correction factors in IAEA TRS - 483 CoP. PinPoint and PinPoint 3D have detector–specific correction factors listed in IAEA TRS - 483 CoP, except for the smallest 4 mm field in which they are not recommended for use due to the size of their effective volume (Table 1.).¹⁰

Diodes are a p-type semiconductor detectors with their effective volumes significantly smaller than the ones from air-vented ionization chambers ranging from 0.02 mm³ to 0.19 mm³. Smaller volumes of diode detectors will result in a smaller volume-averaging correction factor making them appropriate for dose profile measurements in small photon fields. Diode detectors are intended for relative dosimetry of Gamma Knife's narrow photon beams with only two detectors (Diode P and E) listed in IAEA TRS - 483 CoP.

The diamond detector is made from a synthetic diamond with correction factors for Gamma Knife Perfexion available in the literature.¹⁰ Out of all active types of detectors, it has the smallest effective volume. This detector was specially designed for the relative dosimetry of narrow photon beams.

Plastic scintillator detectors (PSD) are considered perturbation free due to their tissue equivalence and can be used for reference and relative dosimetry.⁶⁶ They are active detectors requiring correction for Cherenkov radiation. In this study, two plastic scintillator detectors are modelled in the Gamma Knife Perfexion narrow photon beams to determine its volume-averaging correction factors. Since they are not used in field output factor determination of dose profile measurements, the physics of plastic scintillator detectors and their behaviour in narrow Co-60 photon beams is omitted. No detector-specific correction factors are listed in IAEA TRS-483 CoP for these types of detectors.¹⁰

EBT3 film is a well-established detector for reference and relative dosimetry with correction factors approximated with unity. Radiochromic film is a passive detector requiring calibration of optical density against the absorbed dose.

In the present section, the physics of the listed detector types will be explained in more detail with emphasis on their advantages and disadvantages when it comes to small field dosimetry.

Name	Туре	Manufacturer	Effective	Used in research of:			Gamma Knife Perfexion:	
				OFs	k vol	Dose profiles	Dosimetry type:	DSCF available in IAEA TRS-483 CoP:
Semiflex T31010	Air – vented ion. chamber	PTW - Freiburg	125	Yes	Yes	No	Reference	Yes
Semiflex 3D T31021	Air – vented ion. chamber	PTW - Freiburg	70	Yes	Yes	No	Relative	No
PinPoint T31014	Air – vented ion. chamber	PTW – Freiburg	15	Yes	Yes	Yes	Relative	Yes, except for the 4 mm field
PinPoint 3D T31016	Air – vented ion. chamber	PTW - Freiburg	16	Yes	Yes	Yes	Reference & Relative	Yes, except for the 4 mm field
RAZOR chamber	Air – vented ion. chamber	IBA Schwarzenbruck	10	Yes	Yes	Yes	Relative	No
RAZOR nano chamber	Air – vented ion. chamber	IBA Schwarzenbruck	3	Yes	No	Yes	Relative	No
IBA CC04	Air – vented ion. chamber	IBA Schwarzenbruck	40	Yes	Yes	Yes	Reference	Yes
Diode P T60016	p – type semiconductor	PTW - Freiburg	0.030	Yes	Yes	Yes	Relative	Yes
Diode E T60017	p – type semiconductor	PTW - Freiburg	0.030	Yes	Yes	Yes	Relative	Yes
EFD 3G-pSi	p – type semiconductor	IBA Schwarzenbruck	0.190	No	Yes	Yes	Relative	No
RAZOR diode	p – type semiconductor	IBA Schwarzenbruck	0.020	Yes	Yes	Yes	Relative	No
EDGE detector	p – type semiconductor	Sun Nuclear Corp.	0.020	Yes	Yes	No	Relative	No
microDiamond T60019	Synthetic diamond	PTW - Freiburg	0.004	Yes	Yes	Yes	Relative	Yes
Exradin W2 1x1	Plastic scintillator	Standard Imaging	0.785	No	Yes	No	Reference & Relative	No
Exradin W2 1x2	Plastic scintillator	Standard Imaging	2.356	No	Yes	No	Reference & Relative	No
EBT3	Radiochromic films	GAFchromic	/	Yes	No	Yes	Reference & Relative	Approximated with unity

Table 1. List of the detectors used along their main characteristics.^{60,61,63–65,67,68}

3.2.1. Ionization chambers

Ionization chambers are established and frequently used for both relative and reference dosimetry in small photon fields. They can be air-vented or filled with nonpolar liquid. They work under the principle of ionization of atoms in a medium placed in an electric field through which incident photon beams pass. They can be cylindrical or plane-parallel in design.

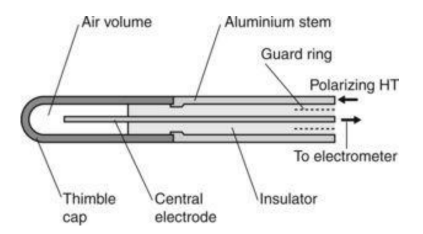


Figure 12. Schematic illustration of a cylindrical ionization chamber. HT stands for High Tension.⁶⁹

A thimble chamber (cylindrical) is the most used type of ionization chamber in narrow photon beam dosimetry. It consists of a central electrode and a cylindrical wall with a spherical or conical end mounted on a cylindrical stem (Figure 12.). When an ionization chamber is placed in the radiation beam, ionization of the medium occurs creating electrons in the walls of a chamber that transverse the air cavity, ionizing the medium inside it. A collected charge can be related to the absorbed dose through a calibration factor typically obtain from the secondary standardised dosimetry laboratory (SSDL). The wall must be thick at least as the range of secondary electrons produced in photon interactions to ensure that all electrons crossing the cavity originated in the wall itself, not surrounding material.⁶⁹ However, the chamber's walls will attenuate the photon beam and this attenuation must be appropriately considered when designing chambers. Effective volume can vary from 0.001 cm³ to 10 000 cm³ depending on the purpose of an ionization chamber. Smaller ionization chambers are typically used in medical physics while large ones are used in nuclear and reactor physics. lonization chambers are characterised by an excellent dose response, dose rate independence, low directional dependence, and an abundance of dosimetry data with these

detectors.^{9,13,29} Furthermore, the reading of air-vented ionization chambers must be corrected for the difference in pressure and temperature from the reference conditions in which the chamber was calibrated as

$$k_{Tp} = \frac{(t+273.15)p_0}{(t_0+273.15)p}$$
3.1

where *t* is temperature in °C and *p* is pressure in kPa of the air in the cavity of the ionization chamber, and t_0 and p_0 are the reference values for temperature and pressure i.e., 20 °C and 101.325 kPa. Ionization chambers are known to exhibit the polarity effect which results in the change of an intensity of collected current when the polarity of collecting potential is reversed, and this needs to be appropriately considered.^{10,70} Additional effect which may influence the accuracy of collected charge is ion recombination effect. In ionization chambers, two separate ion recombination effects can take place: the recombination of ions formed by separate ionizing particle tracks and recombination of ions formed by a single ionizing track.¹⁰ As with the polarity effect, determined charge with ionization chamber must be corrected for this influence.

So far all of the mentioned effects are influence factors that may affect the accuracy of measurements. Beside this, ionization chambers have several perturbation factors that have an impact on their accuracy in dosimetry of small fields: p_{wall} and p_{cell} are perturbations due to the non-tissue equivalence of wall and central electrode material, p_{vol} is perturbation due to the volume-averaging and p_{dis} is perturbation due to displacement effect. The total perturbation correction factor p_{tot} for air-vented ionization chambers is a product of individual perturbations (Figure 13.). The main problem with these detectors, as it will be shown in this work, is that some of them are too large to be used in a narrow photon beams due to volume-averaging and loss of lateral charged particle equilibrium.^{10,51}

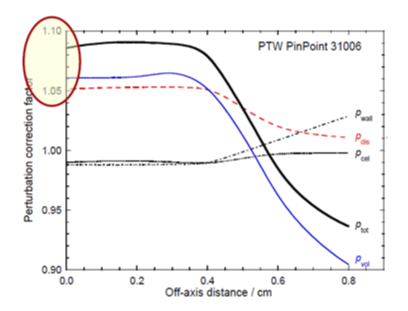


Figure 13. Perturbation correction factors as s function of off-axis distance in small fields for PinPoint T31006 chamber.⁷¹

The development of ionization chambers with small effective volumes (less than 0.1 cm³) enabled them to be used in the dosimetry of narrow photon beams. It has been shown that although small in volume, these ionization chambers still considerably underestimate field output factors in small fields.^{21,29} Some of the physical characteristics of ionization chambers listed in Table 1., are stated in Table 2., and Table 3. for ionization chambers produced by PTW and IBA, respectively.

	-	Semiflex	Semiflex 3D	PinPoint	PinPoint 3D
		T31010	T31021	T31014	T31016
Specification	Reference point / mm ^a	4.5	3.45	3.4	2.4
ecifi	Response / nC/Gy ^b	3.3	2	0.8	0.4
Sp	Chamber voltage / V	400	400	400	300
sures	Wall area density / mg/cm ²	78	84	85	84
Materials and measures	Dimension ^c	Radius 2.75 Length 6.5	Radius 2.4 Length 4.8	Radius 1 Length 5	Radius 1.45 Length 2.9
ials a	Electrode ^d	Al 99.98, 1.1	Al 99.98, 0.8	Al 99.98, 0.3	Al 99.98, 0.6
Mater	Build-up cap	PMMA, 3 mm tick	PMMA, 3 mm tick	PMMA, 3 mm tick	PMMA, 3 mm tick

Table 2. Physical characteristics of PTW ionization chambers.

^a Reference point on the chamber's axis, from the tip of a chamber.

^b Nominal response.

^c Dimensions in millimetres.

^d Material, and the diameter of an electrode in millimetres.

		RAZOR chamber	RAZORnano chamber	CC04
	Reference point / mm ^a	2.3	1.4	2.3
Specification	Response / nC/Gy ^b	0.3	0.11	1
	Chamber voltage / V	300	300	300
	Wall area density / mg/cm ²	88	88	70
Materials		Radius 1		Radius 2
and measures	Dimension ^c	Length 3.6	Diameter 2	Length 3.6
	Electrode ^d	Graphite,	Graphite – EDM3,	Shonka (C-552),
	LIEUUUUE	0.55	1	1

Table 3. Physical characteristics of IBA ionization chambers.

^a Reference point on the chamber's axis, from the tip of a chamber.

^b Nominal response.

^c Dimensions in millimetres.

^d Material, and the diameter of an electrode in millimetres.

3.2.2. Silicon detectors

Semiconductor or diode detectors are commonly used in small-field dosimetry.²⁹ They are made of silicon in a way that a layer of n-type silicon is brought into contact with p-type silicon. This allows the electrons to drift from the n-region to the p-region creating an intrinsic zone. Incident ionizing radiation liberates electrons from the intrinsic zone to the p-layer generating an electrical current, while positive charge carriers travel in the opposite direction. The produced electrical current is proportional to an absorbed dose. Semiconductor detectors do not require an external bias voltage.^{63–65} A schematic representation of the semiconductor detector's working principle is shown in Figure 14.

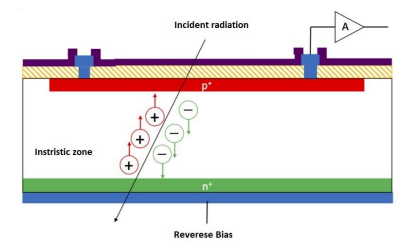


Figure 14. A schematic representation of a semiconductor detector. When traversing through a silicon detector, incoming radiation creates electron-hole pairs along its path. In the presence of an electric field, these charge carriers are separated and start drifting toward the electrodes.⁷²

A silicon detector consists of a layered silicon disk placed horizontally or vertically in the protective and/or build-up material depending on the intended application of such a detector (Figure 15.).^{63–65}

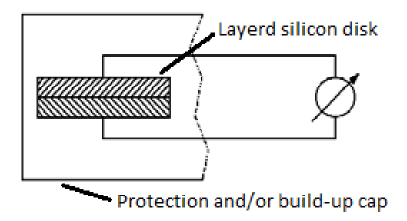


Figure 15. An illustration of a semiconductor detector.⁶³

The sensitivity of silicon detectors is larger when compared to an ionization chamber since it requires only 3.6 eV to create an electron-hole pair in silicon compared to the energy required to generate an ion pair in the air (33.97 eV). Therefore, their effective volumes can be significantly smaller, minimizing the volume-averaging effect. The silicon detectors have been widely used in small photon field dosimetry due to their small size, high resolution, and real-time readout.²⁹ They also have disadvantages such as the dependence on energy, temperature, dose rate, and direction.^{14,33,39,41}

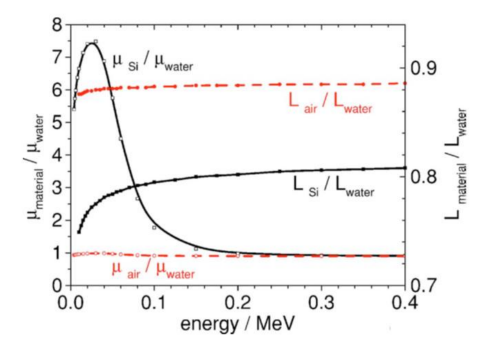


Figure 16. Energy absorption coefficient ratios (left y scale) and restricted stopping power ratios (right y scale) for air and silicon relative to the water.⁴¹

From Figure 16 one can see the main reasons for the difference in response of silicon detectors and ionization chambers. For air, both stopping power and absorption coefficients change only slightly with beam energy resulting in the uniform response of the ionization chambers. By contrast, in silicon, the absorption coefficient is increased almost eight times larger for low photon energies (at approx. 200 keV). Therefore, silicon detectors will overrespond to the low-energy photons. ^{13,33,39} Consequently, silicon detectors are not recommended for dosimetry in broad photon beam geometry due to a significant number of scattered photons. To resolve for this, shielding of an effective volume for some diode detectors is added. Some of the physical characteristics of a silicon detector listed in Table 1. used for field output determination and dose profile measurements, are stated in Table 4. for PTW detectors, and in Table 5. for IBA and Sun Nuclear detectors.

		Diode P T60016	Diode E T60017
	Reference point / mm ^a	2.42	0.77
Specification	Response / nC/Gy ^b	9	9
	Bias voltage / V	0	0
Materials	Window area density / mg/cm ²	250	140
and	Dimensions	Radius 0.56 mm	Radius 0.56 mm
measures	Dimensions	Tick 30 μm	Tick 30 μm

Table 4. Physical characteristics of PTW semiconductor detectors. Diode P and E have the same design, except Diode P's effective volume is shielded thus it is more suitable for dosimetry involving broad beam geometries.

^a Reference point on the detector's axis, from the tip of a detector.

^b Nominal response.

		EFD 3G-pSI	RAZOR diode	EDGE detector
	Reference point / mm ^a	1.2	0.8	0.3
Specification	Response / nC/Gy ^b	100	4.1	32
	Bias voltage / V	0	0	0
Materials	Window area density	Not stated	Not stated	Not stated
		Radius 0.8 mm	Radius 0.3 mm	0.8 x 0.8 mm
	Dimensions			(rectangular)
measures		Tick 0.08 mm	Tick 0.02 mm	Tick 0.03 mm

Table 5. Physical characteristics of IBA and Sun Nuclear semiconductor detectors.

^a Reference point on the chamber's axis, from the tip of a chamber.

^b Nominal response.

3.2.3. Diamond detectors

Diamond detectors are solid-state detectors and their effective volume is built of a natural or synthetic diamond. They have excellent properties such as fast response due to high electronhole mobility and tissue equivalence because of a low atomic number Z=6 and wide band gap of 5.4 eV. Consequently, they have been used in a field of high-energy physics, nuclear engineering and medical physics.⁷³ The operating principle is based on the high-resistivity diamond material sandwiched between two metal electrodes (Figure 17.). An external voltage provides an electric field across the device. Mobile charges produced by absorbed radiation drift in this electric field and generate a current in the external circuit.

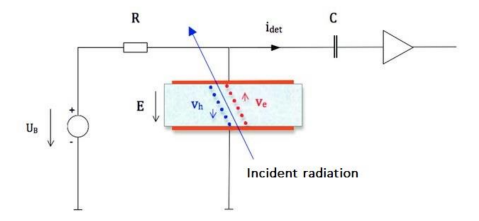


Figure 17. An operating principle of a synthetic diamond detector. The incident radiation generates charge carriers. These are separated by the electric field, thereby producing a signal

current that can be measured with an electrometer. Like silicon semiconductors, no external bias voltage is required. ⁷⁴

Diamond detectors do have disadvantages. They exhibit a substantial energy and dose rate dependence, so correction factors must be applied. In addition, although they have excellent long-term response stability, degradation of the detector's characteristic due to irradiation is inevitable.^{29,41} The physical characteristic of a TW60019 microDiamond (PTW Freiburg, Germany) detector used in this research is stated in Table 6.

	-	microDiamond	
		T60019	
	Reference point / mm ^a	1	
Specification	Response / nC/Gy ^b	1	
	Bias voltage / V	0	
	Window area density /	101	
Materials and	mg/cm ²	101	
measures	Dimensione	Radius 1.1 mm	
	Dimensions	Tick 1 μm	

Table 6. Physical characteristics of microDiamond detector.

^a Reference point on the detector's axis, from the tip of a detector.

^b Nominal response.

3.2.4. EBT3 radiochromic films

EBT3 radiochromic film alongside ionization chambers are some of the most widely available, trusted and studied detector in medical physics used for photon beam dosimetry. It is characterised by high resolution, tissue equivalence, independence of the energy and dose rate, with the possibility of 2D measurements.⁷⁵ In general, the field output factor and determination of dose profiles are in excellent agreement with the values obtained using Monte Carlo calculations.^{17,24–26,30,46,60,75–77} Radiochromic EBT3 films are specially designed for the dosimetry of ionizing radiation and are particularly suited for dosimetry of high-energy photon beams. The EBT3 film is designed for best performance in the dose range from 0.2 to 10 Gy.

The EBT3 film is composed of an active monomer layer, nominally 28 µm tick, sandwiched between two 125 µm matte-polyester substrates. The active layer contains the active components, a marker dye, stabilizer and other components providing an almost energy-independent response.⁴² After the film has been irradiated, an effect of polymerization takes place which results in the change of the film's optical density (OD). The polymerization process of a monomer unit is carried out in three steps: chain initiation, chain propagation and chain termination. This process takes up to 48 hours, and the film must be stored in a dark place at the appropriate temperature. The result of a polymerization effect is the darkening of the film, which increases with the absorbed dose.

Postirradiation processing in the determination of an absorbed dose starts with the film scanning in the red colour channel where it has a peak brightness at λ =636 nm regardless of an absorbed dose (Figure 18.).

A net optical density (netOD) is determined in the region of interest (ROI) as:

$$netOD = \log \frac{PV_{unex} - PV_{bckg}}{PV_{exp} - PV_{bckg}}$$
3.2

where, PV_{unex} and PV_{exp} are the pixel values (measure of pixels intensity) of an exposed and unexposed film respectively, and PV_{bckg} is the pixel values of a background scan.

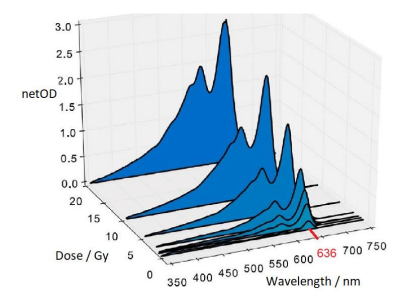


Figure 18. The visible absorption spectrum of seven EBT3 films at different absorbed doses ranging from 0 to 20 Gy after subtraction of a linear background in the range of 350-750 nm. The film has a peak net optical density at 636 nm.⁷⁷

A calibration curve relates the absorbed dose with the film's net optical density. It can be obtained by fitting the Dević fit function to the net optical density determined from films irradiated with different absorbed doses:⁷⁶

$$D(netOD) = a \cdot netOD + b \cdot netOD^n$$
 3.3

where, D (*netOD*) is an absorbed dose as a function of a film's optical density, while a, b and n are fitting parameters. Some of EBT3's disadvantages include a limited lifespan and sensitivity to environmental factors like temperature, humidity and light which can affect the response. They are not intended for very high and low dose measurements. Static electricity can affect film causing them to stick to surfaces or attract dust. Signal to noise ratio in passive detectors such as EBT3 film is larger than in active detectors. In addition, every lot needs to have its own calibration curve. When handling the film, all the above-mentioned factors must be considered. A list of the main EBT3 film's characteristics is presented in Table 7.

Configuration	The active layer (28 μ m) sandwiched between 125 μ m matte-		
Configuration	surface polyester substrate		
Dose range	0.1 – 20 Gy		
Energy dependency	<5 % difference in net optical density when exposed to 100 keV		
Energy dependency	and 18 MeV		
Doco rato rocnonco	<5% difference in net optical density for 10 Gy exposures at rates		
Dose rate response	of 3.4 Gy/min and 0.034 Gy/min.		
Stability in light	$< 5 \cdot 10^{-3}$ change in optical density per 1000 lux-day		
Ctobility in dayle	$< 5 \cdot 10^{-4}$ optical density change/day at 23°C and $< 2 \cdot 10^{-4}$		
Stability in dark	density change/day refrigerated		

Table 7. List of physical characteristics of EBT3 films as stated by the manufacturer.⁴²

3.3. Ellipsoid absorbed dose model for Gamma Knife Icon

The ellipsoid absorbed dose model assumes that a single Gamma Knife isocentre absorbed dose delivery (i.e. shot) forms an ellipsoid in three-dimensional space.⁷⁸ To create a model the dose profiles along main spatial directions have to be fitted to an analytical expression:

$$D(k) = \sum_{i=1}^{n} A_i \cdot erf\left(\frac{k+a_i}{b_i}\right) + C_0$$
3.4

where A_i , a_i , b_i , and C_0 are fitting parameters, k is the off-axis distance on any of the major axes (x, y and z), erf is an error function. The error function with a complex variable (Gauss error function) is defined as:

$$erf(z) = \frac{2}{\sqrt{\pi}} \int_0^z e^{-t^2} dt$$
 3.5

For x and y-axis dose profiles of field sizes n=4 (Equation 3.4), for the z-axis of 16 mm and 8 mm dose profiles n=5, and n=2 for the 4 mm z-axis dose profile. When building the model it is assumed that dose profiles for all field sizes on the x and y-axis are symmetrical with respect to the z-axis i.e. f(x,y)=f(-x,-y). Consequently, only positive off-axis values were considered (Equation 3.4.) to fit dose profiles on the x and y-axis (only absolute values of k are allowed). This symmetry is due to the geometry of the collimating system in which Co-60 sources are symmetrically distributed around the z-axis. By contrast, sources are asymmetrical with respect to the XY plane and both positive and negative off-axis values must be considered when fitting. Asymmetry is most pronounced for the 16 mm field, and it is gradually reduced by decreasing the field down to 4 mm.

The ellipsoid absorbed dose model states that the dose can be considered as having individual components in each spatial direction which can be calculated as:

$$D_x \cdot \cos\theta_x = f_x(|\overrightarrow{r_2} - \overrightarrow{r_1}|) \cdot (x_2 - x_1)/|\overrightarrow{r_2} - \overrightarrow{r_1}|)$$
3.6

$$D_{y} \cdot \cos\theta_{y} = f_{y}(|\vec{r_{2}} - \vec{r_{1}}|) \cdot (y_{2} - y_{1})/|\vec{r_{2}} - \vec{r_{1}}|)$$
 3.7

$$D_z \cdot \cos\theta_z = f_z(\alpha \cdot |\overrightarrow{r_2} - \overrightarrow{r_1}|) \cdot (z_2 - z_1) / (|\overrightarrow{r_2} - \overrightarrow{r_1}|)$$
3.8

where $\vec{r_1} = (x_1, y_1, z_1)$ are the coordinates of the absorbed dose distribution's centre, $\vec{r_2} = (x_2, y_2, z_2)$ are the coordinates of the point at which the absorbed dose is determined, D_x , D_y , and D_z are the dose components on each major axis, and, Θ_x , Θ_y , and Θ_z are the angles between the displacement vector and the axis, finally, α is the sign of $(z_2 - z_1)$.

The absorbed dose $D(\vec{r})$ at any point $\vec{r_2} = (x_2, y_2, z_2)$ with shot focused at $\vec{r_1} = (x_1, y_1, z_1)$ is determined as:

$$D(\vec{r}) = \sqrt{\left(D_x \cdot \frac{x_2 - x_1}{|\vec{r_2} - \vec{r_1}|}\right)^2 + \left(D_y \cdot \frac{y_2 - y_1}{|\vec{r_2} - \vec{r_1}|}\right)^2 + \left(D_z \cdot \frac{z_2 - z_1}{|\vec{r_2} - \vec{r_1}|}\right)^2}$$
 3.9

Absorbed dose D, in the volume of interest V, can be calculated by integrating Equation 3.9., or numerically performing summation over the discrete values of dose-at-point inside the volume of interest (Equation 3.10.).

$$D_V = \int_V D(\vec{r}) d^3 \vec{r} \approx \sum_{i=1}^{i \in V} D_i \Delta V$$
 3.10

In this work, an ellipsoid absorbed dose model is used for the determination of volumeaveraging correction factors for different detectors listed in Table 1. Volume-averaging correction factors are determined using Equation 2.6. A comparison of accuracy in an absorbed dose predicted by the model and determined by the system for calculation and optimization of absorbed dose distributions, Leksell GammaPlan v.11.3.1 (LGP, Elekta, Stockholm), was performed for ellipsoids with different volumes starting at the centre of dose distribution and ending at different isodose values, eight different volumes in total for all field sizes, from 100% to 20% e.g. 100%– 90 %, 100% – 80%,, 100% - 20%. To do this, a single Gamma Knife shot was simulated by LGP with the shot's centre positioned at the centre of the Leksell coordinate system and a solid water phantom with an 80 mm radius and γ -angle set to 90°. A total of 1700, 783, and 378 points were extracted from LGP for 16 mm, 8 mm, and 4 mm fields, respectively. The selection of points was taken randomly for dose values ranging from 99% to 19% of the relative dose in all spatial directions. In MATLAB (TheMathWorks Inc., USA) an ellipsoid absorbed dose distribution was simulated using fitted normalized dose profiles (Equation 3.4.) for an ellipsoid absorbed dose model (Equation 3.9.). To simplify calculations dose distributions were shifted in a way that $\vec{r_1} = (0, 0, 0)$. An examination of the model's accuracy R_V with respect to an absorbed dose distribution is done in a way that a set of points extracted from the LGP were entered in the model and an absorbed dose calculated by the LGP D_{LGP} is compared against the model's D_{model} for a different ellipsoid volume that spanned from the isocentre to a given isodose line (Equation 3.11.).

$$R_{v} = \frac{|D_{LGP} - D_{model}|}{D_{LGP}} \cdot 100\%$$
 3.11

3.4. Statistical analysis and uncertainty estimation of experimental data

Every measurement is susceptible to numerous external factors that need to be considered and properly corrected. There are two types of uncertainties. Type A uncertainty (statistical uncertainty) is determined using statistical analysis of a series of observations, while Type B uncertainty (non-statistical uncertainty) can be evaluated by means other than statistical analysis of a series of observations.⁷⁹ The pool of information for Type B uncertainty includes previously measured data, experience with or general knowledge of the behaviour and properties of relevant materials and instruments, manufacturer's specification or data provided in calibration and other certificates and uncertainty assigned to reference data taken from the handbook. IAEA TRS-483 CoP provides guidelines for the estimation based on the numerous studies performed and compiled together. However, for Gamma Knife, detailed instructions for Type B uncertainty are lacking.¹⁰

For EBT3 film dosimetry, the Type A uncertainty was calculated, while Type B uncertainty were not considered.

For dosimetry performed with the active detectors in Co-60 narrow photon beams, Type A uncertainty comes from:

- Detector short-term stability σ_{sts} determined from multiple measurements i.e. standard deviation of repeated same measurements.
- Detector absorbed dose linearity $\sigma_{lin.}$ can be determined from the different irradiation times for the same field size. These measurements are performed in the msr field (16 mm) where perturbations related to the small field dosimetry are the smallest.
- Detector dose-rate linearity $\sigma_{dr.lin.}$ can be determined by irradiating detectors with different dose rates for the same time. On Gamma Knife, this can be achieved by switching off different sectors.
- Uncertainty due to the leakage σ_l can be determined by collecting charge of detector when detector is not exposed to ionizing radiation.

Uncertainty in long-term detector stability is equal to a few percentages per year (for some detectors less than 1 %/year) and all measurements with a single detector have been performed over the period of few hours, therefore, for the purpose of this research the detector-long term stability has not been included in total uncertainty budget.^{61,63–65}

Type B uncertainties in dosimetry with active detectors on Gamma Knife come from:

- Gamma Knife's geometrical uncertainty σ_{PPS} : uncertainty in the positioning of the detector's reference point at the isocentre i.e. the accuracy of PPS (Patient Positioning System). This can be estimated from the device's declared accuracy.^{3,59}
- Uncertainty due to the electrometer stability σ_{el} . These values are determined from the electrometer specifications.^{61–65,67}

Through this study, a formalism of the Joint Committee for Guides in Metrology (JCGM) was followed for estimations and calculations of uncertainties. This formalism is presented in the JCGM publication: Evaluation of Measurement Data – Guide to the Expression of Uncertainties in Measurements.⁸⁰

3.4.1. Uncertainty in dosimetry with an EBT3 film

To estimate uncertainty $\sigma_{\left(D_{Q_0,Q_0}^{f_{clin},f_{msr}}\right)_{EBT_3}}$ in the absorbed dose determined with EBT3 film, a modified Dević formalism was used.⁷⁶ The modifications were made with weighted fitting and included uncertainties in all fitting parameters as well as correlations between fitting parameters. The total uncertainty comes from the experimental uncertainty $\sigma_{D_{exp}}$ of determined netOD, uncertainty in the fitting parameters $\sigma_{D_{fit}}$ of a calibration curve and uncertainty due to the correlation of fitting parameters σ_{corr} .

$$\sigma_{D_{exp}} = \sqrt{(a + n \cdot b \cdot netOD^{n-1})^2 \cdot \sigma_{netOD}^2}$$
3.12

$$\sigma_{D_{fit}} = \sqrt{netOD^2 \cdot \sigma_a^2 + netOD^{2n} \cdot \sigma_b^2 + 2b^2 netOD^2 \log(netOD)\sigma_n^2} \qquad 3.13$$

$$\sigma_{corr} = 3.14$$
$$= \sqrt{2netOD^{n+1}(\sigma_{ab} + \sigma_{an}blog(netOD)) + 2\sigma_{bn}bnetOD^{2n}log(netOD)}$$

where σ_a , σ_b , σ_N are uncertainties of fitting coefficients a, b, and n, σ_{netOD} is the uncertainty of determined net optical density (Equation 3.3.), σ_{ab} , σ_{an} , σ_{bn} are uncertainties in the correlation between fitting parameters and D (*netOD*) is dose determined at corresponding ROI with *netOD*.

Uncertainty in net optical density σ_{netOD} can be calculated by applying uncertainty propagation formalism on the equation for determination of net optical density (Equation 3.2.):⁷⁶

$$\sigma_{netOD} = \frac{1}{\ln 10} \sqrt{\frac{\sigma_{PV_{unex}}^2 + \sigma_{bckg}^2}{\left(PV_{unex} - PV_{bckg}\right)^2} + \frac{\sigma_{PV_{exp}}^2 + \sigma_{bckg}^2}{\left(PV_{exp} - PV_{bckg}\right)^2}}$$
3.15

where, $\sigma_{I_{unex}}, \sigma_{I_{exp}}$ are uncertainties in the pixel values of an unexposed and exposed film, while σ_{bckg} in uncertainty in the pixel value of a background scan.

Total percentage uncertainty for an absorbed dose determined with the EBT3 film $\sigma_{\left(D_{Q_0,Q_0}^{f_{clin},f_{msr}}\right)_{EBT3}}$, consisting of experimental uncertainty $\sigma_{D_{exp}}$, and uncertainty due to the fitting parameter $\sigma_{D_{fit}}$ is calculated as:

 $\sigma_{\left(D_{Q_0,Q_0}^{f_{clin},f_{msr}}\right)_{EBT_3}} = \sqrt{\sigma_{D_{exp}}^2 + \sigma_{D_{fit}}^2 + \sigma_{corr}^2}$ 3.16

Finally, since EBT3 film measurements are independent and uncorrelated, uncertainties in the field output factors determined with the EBT3 films are calculated by using Equation 3.17.⁸⁰

$$\sigma_{\left(\Omega_{Q_{0},Q_{0}}^{f_{clin},f_{msr}}\right)_{EBT_{3}}} = \sqrt{\sum_{i=1}^{2} \left(\frac{\partial \Omega_{Q_{0},Q_{0}}^{f_{clin},f_{msr}}}{\partial \left(\left(D_{Q_{0},Q_{0}}^{f_{clin},f_{msr}}\right)_{EBT_{3}}\right)_{i}} \sigma_{\left(\left(D_{Q_{0},Q_{0}}^{f_{clin},f_{msr}}\right)_{EBT_{3}}\right)_{i}}\right)^{2}}$$

$$= \sqrt{\left(\frac{1}{\left(D_{Q_{0}}^{f_{msr}}\right)_{EBT_{3}}} \sigma_{\left(D_{Q_{0}}^{f_{clin}}\right)_{EBT_{3}}}\right)^{2} + \left(-\frac{\left(D_{Q_{0}}^{f_{clin}}\right)_{EBT_{3}}}{\left(D_{Q_{0}}^{f_{msr}}\right)_{EBT_{3}}^{2}} \sigma_{\left(D_{Q_{0}}^{f_{msr}}\right)_{EBT_{3}}}\right)^{2}}$$

$$3.17$$

3.4.2. Uncertainty in dosimetry with an active detector

The uncertainty of active detector comes from Type A uncertainty and uncertainty in the alignment of the detector's reference point at the isocentre and electrometer stability (Type B). The total uncertainty is determined as:

$$\sigma_{tot} = \sqrt{\sigma_{TypeA}^2 + \sigma_{TypeB}^2}$$
 3.18

To calculate uncertainties for field output factors $\sigma_{\Omega_{Q_0,Q_0}^{f_{clin},f_{msr}}}$ and volume-averaging correction factors $\sigma_{(k_{vol})_{Q_0}^{f_{msr},f_{clin}}}$ a Gaussian theory is used.⁸⁰ An uncertainty for field output factor can be determined as:

$$\sigma_{\Omega_{Q_0,Q_0}^{f_{clin},f_{msr}}} = \sqrt{\sum_{i=1}^{2} \left(\frac{\partial \Omega_{Q_0,Q_0}^{f_{clin},f_{msr}}}{\partial \left(M_{Q_0}^{f_{clin},f_{msr}} \right)_i } \left(\sigma_{tot}^{f_{clin},f_{msr}} \right)_i \right)^2}$$
3.19

$$\sigma_{\Omega_{Q_0,Q_0}^{f_{clin},f_{msr}}} = \sqrt{\left(\frac{1}{\mathsf{M}_{Q_0}^{f_{msr}}}\sigma_{tot}^{f_{clin}}\right)^2 + \left(-\frac{\mathsf{M}_{Q_0}^{f_{clin}}}{\left(\mathsf{M}_{Q_0}^{f_{msr}}\right)^2}\sigma_{tot}^{f_{msr}}\right)^2}$$
3.20

where $\sigma_{tot}^{f_{clin}}$ and $\sigma_{tot}^{f_{msr}}$ are total uncertainty (Type A and B) of a detected charge in the clinical and machine-specific reference field.

When estimating Type B uncertainties with active detectors the most important factor is the geometrical accuracy of the PPS e.g. alignment of the detector's effective point of measurement at the isocentre. The maximum allowed geometrical error is 0.3 mm⁵⁹, and to estimate the error of the detected signal a rectangular probability density function is assumed where all outcomes are equality likely to occur. To convert an uncertainty with a rectangular distribution to a standard deviation equivalent we have divided the estimated uncertainty U_{PPS} by the square root of three.

$$\sigma_{PPS} = \frac{U_{PPS}}{\sqrt{3}}$$
 3.21

where σ_{PPS} is the standard uncertainty and U_{PPS} is the estimated uncertainty.

Additionally, electrometer uncertainty was determined from its specification. ^{81,82}

The combined standard uncertainty is multiplied by the coverage factor that indicated the level of confidence in the measurement result. This new value (combined standard uncertainty multiplied by the coverage factor) is expanded uncertainty:

$$\sigma_{exp} = k\sigma_{tot}$$
 3.22

where k is the coverage factor. The primary standard for all dosimetry quantities in medical physics uses k=2, corresponding to an interval with a 95% level of confidence.⁸⁰

The detector-specific correction factor in different fields is determined by dividing the field output of a detector $\left(\Omega_{Q_0,Q_0}^{f_{clin},f_{msr}}\right)_{detector}$ with the reference values such as EBT3 and Monte Carlo $\left(\Omega_{Q_0,Q_0}^{f_{clin},f_{msr}}\right)_{MC,EBT3}$.

$$k_{Q_0,Q_0}^{f_{clin},f_{msr}} = \frac{\left(\Omega_{Q_0,Q_0}^{f_{clin},f_{msr}}\right)_{detector}}{\left(\Omega_{Q_0,Q_0}^{f_{clin},f_{msr}}\right)_{MC,EBT3}}$$
3.23

By using the same statistical analysis as before, an uncertainty for the detector-specific correction factor $\sigma_{k_{Q_0,Q_0}^{f_{clin},f_{msr}}}$ can be determined as:

$$\sigma_{k_{Q_{0},Q_{0}}^{f_{clin},f_{msr}}} = \begin{pmatrix} \left(\frac{1}{\left(\Omega_{Q_{0},Q_{0}}^{f_{clin},f_{msr}}\right)_{detector}} \sigma_{\left(\Omega_{Q_{0},Q_{0}}^{f_{clin},f_{msr}}\right)_{MC,EBT_{3}}} \right)^{2} \\ + \left(-\frac{\left(\Omega_{Q_{0},Q_{0}}^{f_{clin},f_{msr}}\right)_{MC,EBT_{3}}}{\left(\left(\Omega_{Q_{0},Q_{0}}^{f_{clin},f_{msr}}\right)_{detector}\right)^{2}} \sigma_{\left(\Omega_{Q_{0},Q_{0}}^{f_{clin},f_{msr}}\right)_{detector}} \right)^{2} \end{pmatrix}^{2} \end{pmatrix}^{2}$$
3.24

When calculating uncertainty for the detector specific correction factor, field output values obtained with the Monte Carlo $\sigma_{\left(\Omega_{Q_0,Q_0}^{f_{clin},f_{msr}}\right)_{MC}}$ were taken as absolute i.e. without uncertainty.

Uncertainties for volume-averaging $\sigma_{(k_{vol})_{Q_0}^{f_{msr,f_{clin}}}}$ are determined similarly as for field output factors, by partially deriving an expression for volume-averaging (Equation 2.6.).

$$\sigma_{(k_{vol})_{Q_0}^{f_{msr,f}}c_{lin}} = \sqrt{\left(\frac{\partial (k_{vol})_{Q_0}^{f_{msr,f}}c_{lin}}{\partial D_{Q_0}^{f_{msr,f}}c_{lin}}\sigma_{D_{Q_0}^{f_{msr,f}}c_{lin}}\right)^2}$$
3.25

Where $D_{Q_0}^{f_{msr}, f_{clin}}$ is an absorbed dose over the detectors volume in clinical and msr field with beam quality Q_0 . When derived (Equation 3.25.):

$$\sigma_{(k_{vol})_{Q_0}^{f_{msr,f_{clin}}}} = \sqrt{\left(-\frac{V}{D_{Q_0}^{f_{msr,f_{clin}}^2}}\sigma_{D_{Q_0}^{f_{msr,f_{clin}}}}\right)^2}$$
3.26

and by substituting $\sigma_{D_{Q_0}^{fmsr,f}clin}=D_{Q_0}^{fmsr,f}clin}\cdot\overline{R_{\nu}}$,

$$\sigma_{(k_{vol})_{Q_0}^{f_{msr,f_{clin}}}} = \sqrt{\left(-\frac{V}{D_{Q_0}^{f_{msr,f_{clin}}}}\overline{R_v}\right)^2} = (k_{vol})_{Q_0}^{f_{msr,f_{clin}}} \cdot \overline{R_v}$$
 3.27

where $\overline{R_{v}}$ is the mean uncertainty in determined absorbed dose within the detector effective volume and it is determined as an average value of difference between TPS and elliptical absorbed dose model for a given field.

3.5. Uncertainties with active detectors

There are three sources of uncertainness for which measurements with active detectors had to be conduct. This uncertainness come from leakage current which negatively affect the collected charge, detector's dose rate dependence and uncertainty in the linearity of detector's response with the absorbed dose. All three factors will affect the accuracy of final result and must be appropriately considered in the uncertainty budget.

Detector's leakage current refers to the current that flows through a detector even when it's not actively detecting any signal. In various electronic devices, especially in semiconductorbased detectors, this current can arise due to imperfections or impurities in the materials used to construct the detector. The magnitude of the detector leakage current depends on factors such as the type of detector, its operating conditions, and the quality of its construction. In some applications, minimizing detector leakage current is crucial to improve the sensitivity and accuracy of measurements.

To estimate leakage current uncertainty σ_l , a charge collected when detector is outside of the field is divided by the charge collected when detector is inside different fields.

$$\sigma_l = \frac{M_0}{M_{msr,clin}}$$
 3.28

where, M_0 is the charge collected outside of the filed and $M_{msr, clin}$ is the charge collected in the clinical and machine-specific reference field.

Linearity in absorbed dose refers to the relationship between the dose of radiation delivered to a material and the response of the radiation detector used to measure that dose. In an ideal scenario, the response of the detector would be directly proportional to the absorbed dose received by the material being measured. This means that if you double the absorbed dose, the detector's response would also double. This linear relationship simplifies calibration and interpretation of measurements. However, in reality, there can be deviations from perfect linearity. Maintaining linearity in absorbed dose measurements is important for ensuring the accuracy and reliability of radiation dose assessments in various applications, including medical radiation therapy, industrial radiography, and environmental monitoring. Calibration procedures and correction factors are often employed to account for any non-linearities and ensure accurate dose measurements across a wide range of dose levels.

Uncreating σ_{lin} in the linearity of an absorbed dose can be determined from measurements as:

$$\sigma_{lin} = \sqrt{\frac{\sum_{i=1}^{n} (M_{observed} - M_{fitted})^2}{n-2}}$$
3.29

where $M_{observed}$ is the measured charge, M_{fit} is the fitted value for the same measurement, and n is the number of measurements.

Finally, dose-rate linearity $\sigma_{dr.\ lin}$ refers to the relationship between the dose rate of radiation and the response of a radiation detector. In an ideal scenario, the detector's response would be directly proportional to the dose rate, meaning that if the dose rate doubles, the detector's response (such as the electrical signal it produces) would also double. Uncertainty in the dose rate linearity can be determined by using the Equation 3.29.

4. Experimental Setup and Measurements

4.1. Determination of field output and detector-specific correction factors

Determination of field output factors $\Omega_{Q_{clin},Q_{msr}}^{f_{clin},f_{msr}}$ is most important part of a small photon field dosimetry. In this research, field output factors were determined for thirteen detectors (seven ionization chambers, four semiconductor detectors, a diamond detector and EBT3 film) specified in Table 1.

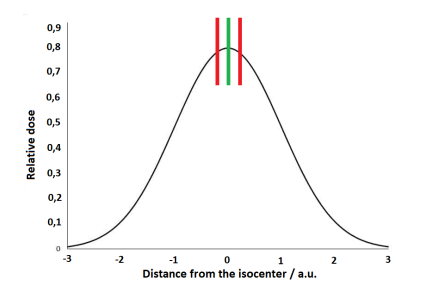
4.1.1. Determination of field output factors using EBT3 films

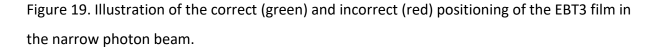
Due to the properties of an EBT3 film it can be used as a reference detector to obtain a detector-specific correction factor $k_{Q_{clin},Q_{msr}}^{f_{clin},f_{msr}}$ for other detectors. In all measurements involving EBT3 film, the same protocol for an absorbed dose determination was used:⁷⁶

- Irradiated films were wrapped with aluminium foil and stored for a minimum of 48 hours at a temperature of about 10°C so that optical density can saturate.
- Scanning was performed in transmission mode of EPSON EXPRESSION 10000XL (Epson America Inc., Long Beach) scanner in the red channel with a resolution of 150 dpi. Before the film scanning, the scanner was warmed up to ensure scanning stability.
- ImageJ software (National Institutes for Health, USA) was used for image processing and noise filtering. Filtering of noise was done by applying the filter within 3x3 pixel kernel.
- Region of interest (ROI) was placed at the centre of the film and its diameter equalled 4, 2, and 0.8 mm for 16 mm, 8 mm, and 4 mm beam sizes respectively. This ROI was used to maximise the number of points inside it and to improve the signal-to-noise ratio. The calibration curve for the film lot used in this research was created by irradiating a set of films in the XY plane using a 16 mm absorbed dose distribution with doses ranging from 0.4 to 8 Gy.

- Calibration curve was obtained by fitting the function (Equation 3.3.) to the net optical density of pixel values determined at the ROI placed at the centre of a films irradiated with different absorbed doses.
- For the calculation of field output factors, the average pixel value for all three beam sizes is determined at the ROI and related to the net optical density (netOD) (Equation 3.2.). Finally, the field output factors were calculated as a ratio of the dose in the clinical and machine-specific reference fields (Equation 2.10.).

Before irradiation, it was necessary to find the coordinates of the shot. Accurate positioning of the film inside the field is essential, especially for the 4 mm field whose dose plateau is very small, therefore it is crucial to position the film at the centre of a field. This problem is illustrated in Figure 19.





To solve this, a test film (Figure 20.) was positioned inside the solid water phantom in the XY plane and a CBCT (Cone-Beam Computed Tomography) scanning was performed to find the coordinates of the centre of the film (in the z-direction). These coordinates were used for the determination of field output factors. All films for field output factor determination were irradiated in the XY plane with shot coordinates (100.2, 100.2, 99.4). Irradiation time was 2.69 minutes which corresponds to 6 Gy of an absorbed dose for a 16 mm field size.

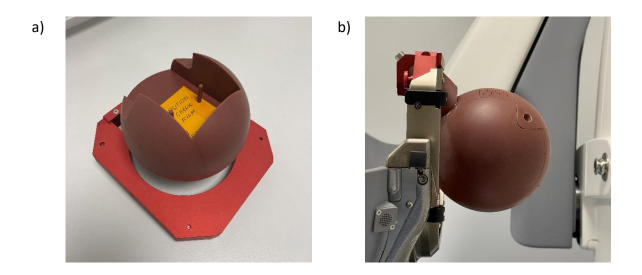


Figure 20. Open (a) and closed (b) solid water phantom with test film.

To determine field output factors with EBT3 film, a calibration curve was created by irradiating eighteen films with different doses from 0 Gy to 8 Gy. The field output factor is calculated as the ratio of absorbed doses in the clinical and machine-specific reference field as:

$$\Omega_{Q_0,Q_0}^{f_{clin},f_{msr}} = \frac{D_{Q_0}^{f_{clin}}}{D_{Q_0}^{f_{msr}}}$$
4.1

4.1.2. Determination of field output factors using active detectors

Determination of field output factors $\Omega_{Q_{clin},Q_{msr}}^{f_{clin},f_{msr}}$ with the active detectors was performed in the solid water phantom with the long axis of a detector positioned in the z direction of the Leksell coordinate system (Figure 21.).

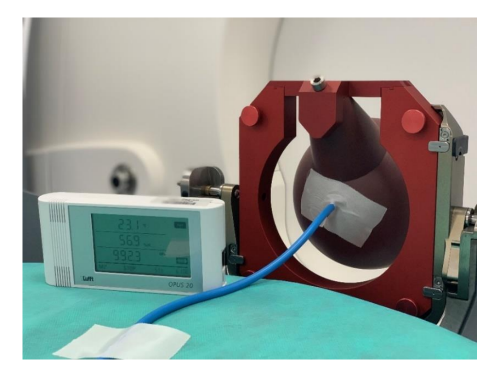


Figure 21. Experimental setup for field output factors determination using an ionization chamber.

Custom 3D-printed inserts were manufactured out of tissue-equivalent plastics for detectors that do not have inserts provided by the manufacturer. Also, any remaining air gaps were filled with tissue-equivalent gel. The presence of air gaps was checked with CBCT. Coordinates of reference point were determined firstly by performing a CBCT scan (Figure 22.) and manually positioning the detector at the shot centre along with additional adjustments according to measured dose profiles of three principal axes with the resolution of 0.5 mm and 15 seconds of signal acquisition M. For IBA's detectors, the DOSE 1 High Performance Reference Class Electrometer was used, and for the remaining detectors, the PTW UNIDOS E electrometer was used.^{81,82}

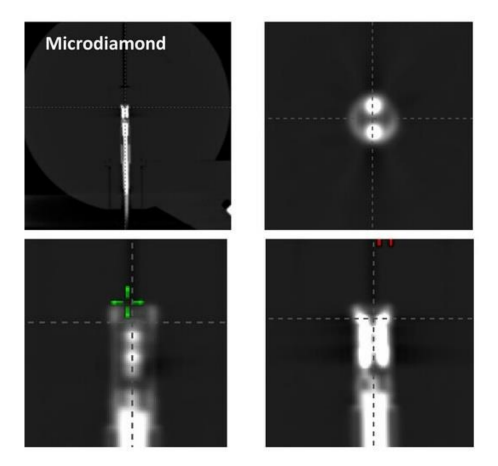


Figure 22. A CBCT scan of PTW microDiamond T60019 inside the solid water phantom. The image shows a coronal image of the entire phantom with detectors inside it, (top left), axial (top right), longitudinal (bottom left) and sagittal (bottom right) reconstruction.⁸³

The x coordinate of a reference point can be determined as a weighted average of the detector position from dose profile measurements as:

$$x = \frac{\sum_{i=1}^{n} x_i M_i}{\sum_{i=1}^{n} M_i}$$
 4.2

where x_i is the coordinate of a point on a profile's axis at which M_i charge is determined. The same procedure was done for the y and z coordinates of the reference point. This is the most accurate method of detector positioning, and it assures that the detector's reference point is at the shot's centre (Table 8.).

Detector	x / mm	y / mm	y / mm
Semiflex T31010	100.1	99.7	99.4
Semiflex 3D T31021	100.2	99.7	99.8
PinPoint T31014	100.1	99.7	99.8
PinPoint 3D T31016	100.2	99.8	99.8
RAZOR chamber	100.2	99.6	100.4
RAZORnano chamber	100.3	99.7	99.4
IBA CC04	100.3	99.8	102.5
Diode P T60016	100.2	99.7	100.9
Diode E T60017	100.2	99.6	100.2
RAZOR diode	100.4	99.8	99.5
EDGE detector	101.6	100.2	101.4
microDiamond T60019	100.2	99.2	100.1
EBT3	100.2	100.2	99.4

Table 8. Coordinates of the detector's effective point of measurement for field output factor determination on the Gamma Knife Icon.

To determine field output factors, ten measurements of response during 60 seconds of acquisition time were averaged for each beam size. Temperature and pressure were monitored when using air-vented ionization chambers.

Therefore, the response determined with vented ionization chambers must be corrected for pressure and temperature changes which affect field output factors (Equation 4.3.).

$$\Omega_{Q_{0},Q_{0}}^{f_{clin},f_{msr}} = \frac{M_{Q_{0}}^{f_{clin}} \cdot (k_{t,p})_{clin}}{M_{Q_{0}}^{f_{msr}} \cdot (k_{t,p})_{msr}}$$
4.3

Temperature and pressure were monitored with an OPUS 20 THI device (G. Luffi Mess – und Regeltechnik GmbG, Deutschland).⁸⁴ A detector-specific correction factor $k_{Q_0,Q_0}^{f_{clin},f_{msr}}$ for active detectors is determined by comparing the measured output with values obtained using Monte Carlo and EBT3 film. For clarity, the notation was changed from Q_{msr} and Q_{clin} to Q_0 to indicate the same beam quality in the machine-specific reference field and clinical field.

$$k_{Q_0,Q_0}^{f_{clin},f_{msr}} = \frac{\left(\Omega_{Q_0,Q_0}^{f_{clin},f_{msr}}\right)_{MC,EBT3}}{\left(\Omega_{Q_0,Q_0}^{f_{clin},f_{msr}}\right)_{detector}}$$
4.4

To determine the difference in field output factors $\Omega_{Q_0,Q_0}^{f_{clin},f_{msr}}$ of active detectors from values obtained with the Monte Carlo and EBT3 film, a relative difference R_{EBT3, MC} is calculated (Equation 4.5.). This value gives information on whether it is appropriate to use a detector in a given field size. Ideally, $R_{EBT3,MC} = 0$ %.

$$R_{EBT3,MC} = \frac{\left(\Omega_{Q_{0},Q_{0}}^{f_{clin},f_{msr}}\right)_{detector} - \left(\Omega_{Q_{0},Q_{0}}^{f_{clin},f_{msr}}\right)_{EBT3,MC} \cdot 100\%$$
4.5

4.2. Volume-averaging correction factors

To determine a volume-averaging correction factor $(k_{vol})_{Q_0}^{f_{msr},f_{clin}}$ in clinical and machinespecific reference fields, an ellipsoid absorbed dose model was used for simulation of absorbed dose distribution along with geometries of different detectors. The formalism of an ellipsoid absorbed dose model of Gamma Knife Icon was explained in section 3.3., while simulation of that model in MATLAB was performed (Equation 3.9.) with a fitted dose profile (Equation 3.4.).

The volume-averaging correction factor was determined by integrating the relative dose distribution over the detector's effective volume using numerical integration; the Simpson's method with hundred integration steps in each spatial direction e.g., the volume of the detector was divided into one million parts. The volume-averaging correction factor was determined as a ratio of the detector's effective volume with the value of a normalized field integrated over that volume (Equation 2.6.). When simulating the detector's geometry, detectors were positioned inside the field in a way that their reference point is placed at the centre of the Leksell coordinate system with coordinates (100, 100, 100) with the detector's long axis parallel with the z direction. The code written in MATLAB used for the determination of the volume-averaging correction factor is provided in Appendix B. A list of the examined detectors can be found in Table 1. To calculate the volume-averaging correction factor, it was necessary to know the detailed geometry of each detector and its position inside the different absorbed dose distributions. Detailed geometry implies data on the boundaries and shape of the effective volume and size of a central electrode, if any. This was determined using schematics provided by the manufacturers.^{61,63–65} Since most detector's effective volumes are cylindrical, the integration in the cylindrical coordinate system was performed. Diamond and most of the semiconductor detectors have a cylindrical effective volume except for the EDGE detector which is rectangular. An illustration of different detector geometries can be seen in Figure 23. All investigated ionization chambers are cylindrical in shape with a curved top and cylindrical central electrode. Detailed schematics of ionization chambers can be found in Appendix A.

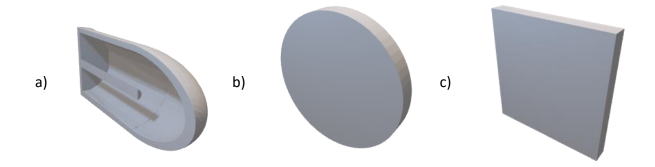


Figure 23. Illustration of the geometry of effective volume for studied detectors: a) ionization chambers, b) diamond, plastic scintillator and semiconductor detectors except, c) EDGE SunNuclear detector.

The design of an ionization chamber can be subdivided into three parts:

- 1. Cylindrical part from the bottom until the beginning of a cap
- 2. Cap on the top with the radius of curvature R, and
- 3. Cylindrical electrode.

To accurately calculate volume-averaging over the hemispherical part of an ionization chamber it was necessary to find the radius of curvature i.e. the equation for the curvature of a cap in cylindrical coordinates. For some ionization chambers, the top part was a spherical cap. Therefore, it was necessary to solve a system of equations (Equations 4.7. and 4.8.) to calculate the volume of a spherical cap. In general, for cylindrical ionization chambers the size of the effective volume is calculated as:

$$V = V_{cylin} + V_{cap} - V_{elec}$$
 4.6

where, V_{cylin} , V_{cap} and V_{elec} are the volumes of a cylinder, cap and electrode respectively.

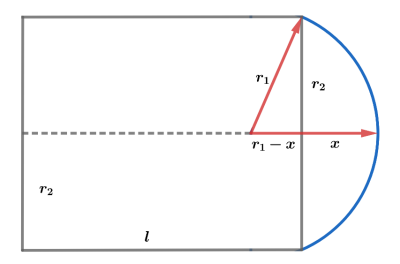


Figure 24. The geometry of an ionization chamber.

From Figure 24. it follows:

$$V = V_{cylin} + V_{cap} - V_{elec} = r_2^2 \pi l + \frac{\pi x^2}{3} (3r_1 - x) - V_{elec}$$

$$r_2^2 - 2r_1 x + x^2 = 0$$
4.8

and by solving Equations 4.7. and 4.8., we can obtain r_1 and x values. An equation of a cap's curvature can be expressed in the cylindrical coordinate system as:

$$s^{2} + (z + r_{1} - x)^{2} = r_{1}^{2}$$
 4.9

where s and z are the coordinates in the cylindrical coordinate system. Dimensions of ionization chambers used in this research are listed in Table 9. Numerical calculations of volume-averaging correction factors for ionization chambers were done in three steps over three different chamber volumes:

- 1. Integral over the cylindrical part J_1
- 2. Integral over the cap part J_2
- 3. Integral over the volume of the electrode $-J_3$

The total integral value J_{tot} of an ionization chamber, the denominator of Equation 2.6., is determined as:

$$J_{tot} = J_1 + J_2 - J_3$$
 4.10

Ionization chambers	r ₁ / mm	r₂ / mm	l / mm	r1 - x / mm
Semiflex T31010	2.50	2.75	4.00	0.00
Semiflex 3D T31021	2.40	2.40	2.80	0.40
PinPoint T31014	1.98	1.00	4.73	0.27
PinPoint 3D T31016	1.58	1.45	1.94	0.96
RAZOR chamber	1.00	1.00	2.60	0.00
CC04	2.00	2.00	1.60	0.10

Table 9. Dimensions of ionization chambers used for volume-averaging determination.

Detectors	Length / µm	Radius / mm
Diode P T60016	30	0.56
Diode E T60017	30	0.56
EFD 3G-pSI	80	1.60
RAZOR diode	20	0.30
microDiamond	1	1.10
Exradin W2 1x1	1000	1.00
Exradin W2 1x3	3000	1.00
EDGE	30	0.8 (square)

Table 10. Dimensions of cylindrically shaped effective volumes of detectors used for volumeaveraging determination. EDGE detector has s square effective volume.

A volume-averaging perturbation $(k_{vol})_{Q_0,Q_0}^{f_{msr},f_{clin}}$, a part of detector-specific field output correction factor $k_{Q_0,Q_0}^{f_{msr},f_{clin}}$, was determined as:

$$(k_{vol})_{Q_0,Q_0}^{f_{msr,f}clin} = \frac{(k_{vol})_{Q_0}^{f_{msr,clin}}}{(k_{vol})_{Q_0}^{f_{msr}}}$$
4.11

where $(k_{vol})_{Q_0}^{f_{msr,clin}}$ is the volume-averaging correction factor in msr and clin field and $(k_{vol})_{Q_0}^{f_{msr}}$ is the volume averaging correction factor in msr (16 mm) field. The contribution of a volume-averaging to a field output correction factor as:

$$k_{\%vol} = \frac{(k_{vol})_{Q_0,Q_0}^{f_{msr},f_{clin}}}{k_{Q_0,Q_0}^{f_{msr},f_{clin}}} \cdot 100\%$$
4.12

4.3. Gamma Knife's Perfexion dose profiles

Relative dose profile determination, also known as off-axis ratios, refers to the measurements of the absorbed dose or relative dose distribution along a particular axis. Dose profiles on Gamma Knife are typically measured along three principal axes due to the device's collimator system (Figure 6.).^{85,86}

Dose profiles are usually obtained by moving a detector along the axis and measuring the response at various points. They can be also measured with planar detectors e.g., EBT3 or an array of detectors. On the other hand, active detectors such as diodes, and ionization chambers may average the response over the detectors' effective volume affecting the dose profiles (Figure 5.). Reference geometry for dose profile determination on Gamma Knife implies using a film detector placed inside a spherical phantom which is positioned at the isocentre. By contrast, in non-reference geometry the active detector is positioned in the phantom and moved along the desired axis to measure detector's response at various points. Choosing the right detector is a nontrivial task requiring a profound knowledge of the detector's performance and design, with volume-averaging usually being the main limiting factor. A detailed process of profile determination with film detectors on Gamma Knife is explained in the Report of the American Association of Physicists in Medicine (AAPM) Task Group 178 (TG 178).⁸⁶ The resulting data is used to create a graph of a measured detector's response as a function of distance from the isocentre on the principal axis of the Gamma Knife. Dose profiles can be used to verify photon beam shape and alignment of the beam's central axis with the target. According to the TG 178, the tolerance of all FWHM values of measured dose profiles is ±1 mm of the TPS data calculated by the Monte Carlo.

The dose profile, illustrated in Figure 25, is characterized by:

- full width at half maximum (FWHM).
- width of penumbra region; between 80% 20% of isodose value.
- dose gradient up to 70%/mm for 4 mm field. In this work, a mean dose gradient is calculated in the penumbra region.

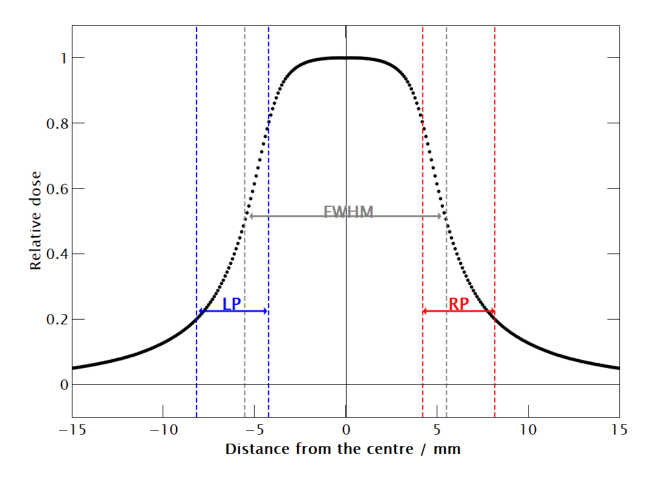


Figure 25. An illustration of a dose profiles with its main characteristics. LP and RP stands for the width of left and right penumbra, respectively.

Dose profiles were determined with active detectors (Table 1.) positioned in the spherical solid water phantom the same way as for the field output factor determination (Figure 21.).

Dose profiles were determined with resolution of 0.2 mm and 15 s of detector acquisition time for each spatial point. Afterwards, measured profiles are normalized to a maximum value of an absorbed dose and imported into CurveExpert Professional software where Savitzky – Golay smoothing is applied to the data points (Figure 26.).

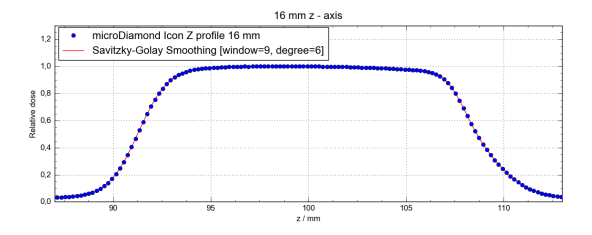


Figure 26. Example of Savitzky - Golay smoothing (red line) for 16 mm z-axis dose profile measured with microDiamond T60019.

For the determination of dose profiles with EBT3 films, a total of six films were used, two for each collimator. Films were positioned at the XY and XZ plain and irradiated with the absorbed dose of 6 Gy at the maximum, for a 16 mm collimator. From previous experience, we found that two planes are enough to determine all three dose profiles. Piercing of the film is performed to create fiducial holes as references for profile determination in each plane. The same postprocessing procedure of EBT3 films for dose profile determination was followed as for the field output factor determination.

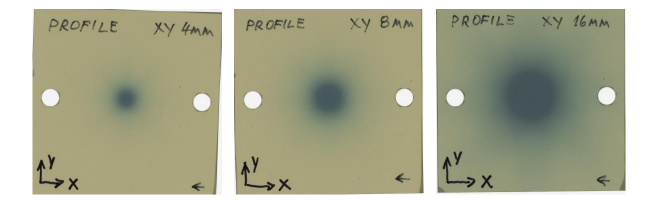


Figure 27. Example of EBT3 film irradiated in XY plane with fiducial holes for dose profile determination. All EBT3 films were scanned in the same direction (bottom right corner arrow) to eliminate variations in measured relative optical density due to film's orientation relative to the scanning direction.⁸⁷

After saturation of the film's optical density, the scanning was performed and dose profiles were determined using ImageJ software with the help of fiducial points as a reference for each plane. The resulting optical density was normalized at the values 6 Gy, 5.334 Gy and 4.956 Gy for 16 mm, 8 mm and 4 mm field sizes respectively. An example of a 16 mm z-axis dose profile measured with EBT3 film is shown in Figure 28.

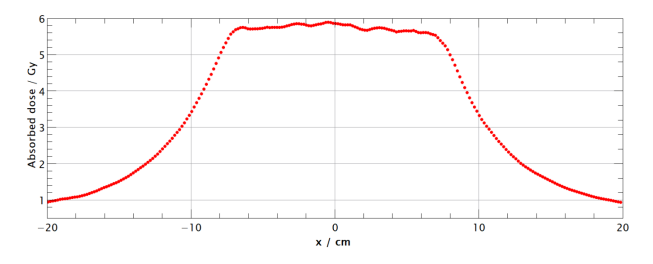


Figure 28. An example of an ImageJ filtered 16 mm x-axis dose profile determined with EBT3 film.

To find a centre of the film, Equation 4.2. was used only in the penumbra region as explained in TG 178.⁸⁸ From there, the film centre x_{cent} was determined as:

$$x_{cent} = \frac{x_1 + x_2}{2}$$
 4.13

where x_1 and x_2 are the centres of the left and right penumbras respectively. Afterwards, the resulting profiles were imported into CurveExpert Professional and the same analysis is done as for the profiles obtained with active detectors.

5. Results and Discussion

	Field size / mm	Value / %			
	16: σ _{net0D16}	0.16			
ncertainty in netOD: σ_{netOD}	8: σ_{netOD_8}	0.10			
	4: σ_{netOD_4}	0.10			
Combined uncertainties					
Uncertainty in the absorbed dose for	16: $\sigma_{(D^{16}_{Q_0})_{EBT_3}}$	3.17			
16 mm, 8 mm and 4 mm field size:	8: $\sigma_{\left(D^{8}_{\boldsymbol{Q}_{0}}\right)_{EBT_{3}}}$	2.99			
$\sigma_{\left(D_{Q_0,Q_0}^{f_{msr,f_{clin}}}\right)_{EBT_3}}$	$4: \sigma_{\left(D^{4}_{Q_{0}}\right)_{EBT_{3}}}$	2.91			
Incontainty in the field output factors	16: $\sigma_{\left(\Omega_{O_0}^{f_{16}}\right)_{\text{FDTO}}}$	4.49			
Jncertainty in the field output factor: $\sigma_{\left(\Omega_{Q_0,Q_0}^{f_{msr,f_{clin}}}\right)_{EBT_3}}$	8: $\sigma_{\left(\Omega_{O_{2}}^{f_{8}}\right)}$	4.36			
$\left(\Pi_{Q_0,Q_0} \right)_{EBT_3}$	$\frac{(q_0)_{EBT_3}}{4:\sigma_{\left(\Omega_{Q_0}^{f_4}\right)_{EBT_3}}}$	4.31			

5.1. Uncertainties in dosimetry of Co-60 narrow photon beams

Table 11. Uncertainty analysis for the dosimetry of Gamma Knife's Perfexion narrow photon beams with the EBT3 film (field output factors). Values are rounded to two significant digits.

In this section, the results of uncertainty analysis are presented for passive (Table 11.) and active detectors (Table 12.).

The determination of an absorbed dose with EBT3 film is subjected to uncertainties that come from three basic sources; from an experimental nature in determining dose using netOD σ_{Dexp} using the fit function, from the fit process σ_{fit} , and from the correlation between fit parameters σ_{corr} .⁷⁶ The experimental uncertainty is primarily caused by the contribution from netOD measurement uncertainty σ_{netOD} and less by other means (e.g. registration of a source with a film, uncertainty in the Gamma Knife calibration, mismatch in the temporal and thermal history of a film, the difference in mean response from one piece of a film to another etc.). For calculation of uncertainties in net optical density σ_{netOD} an error propagation expression is used (Equation 3.15.). When examining uncertainty in the absorbed dose determined with the EBT3 film $\sigma_{\left(D_{Q_0,Q_0}^{f_{msr},f_{clin}}\right)_{EBT3}}$ the largest contribution is due to the uncertainty of the fitting

parameters σ_{fit} (Table 11.). Experimental uncertainty in the dose determined $\sigma_{D_{exp}}$ can be reduced by irradiating more films. When combined, the uncertainty in the absorbed doses $\sigma_{\left(D_{Q_0,Q_0}^{f_{msr.f_{clin}}}\right)_{EBT_3}}$ are 3.17%, 2.98% and 8.91% for 16 mm, 8 mm, and 4 mm field size. Finally, as uncertainty propagate thorough calculations, the uncertainties in the field output factor $\sigma_{\left(\Omega_{Q_0,Q_0}^{f_{msr.f_{clin}}}\right)_{EBT_3}}$ are increased to 4.49%, 4.36%, and 4.31% for 16 mm, 8 mm, and 4 mm field, respectively.

From this analysis, one can see that although EBT3 film has desirable qualities such as tissue equivalence, energy and dose rate independence, it is burdened with relatively large uncertainties when used for absorbed dose determination. This uncertainty comes from the film's design, post-irradiation processing and experience in handling such detectors.⁷⁶ This is the main reasons why measurements obtained with an EBT3 film detector should always be compared against measurements with an active detector to ensure the best accuracy of the result.

	Type A uncertainties					Type B uncertainties						
		cor's shor cor's shor cor's shor contract σ_{sts}		Linearity in absorbed dose σ_{lin} / %	Dose-rate linearity $\sigma_{dr. \ lin}$ / %	Lea	kage σ_l	/%	Electrometer Stability / %		etector a phanton ioning σ_{j}	m
Field size / mm Detector name	16	8	4	All fields	All fields	16	8	4	All fields	16	8	4
Semiflex T31010	0.015	0.009	0.021	0.007	0.130	0.010	0.013	0.030	0.5	0.06	0.30	0.48
Semiflex 3D T31021	0.011	0.020	0.019	0.120	0.130	0.019	0.022	0.042	0.5	0.06	0.30	0.48
PinPoint T31014	0.008	0.029	0.011	0.024	0.094	0.077	0.088	0.120	0.5	0.06	0.30	0.48
PinPoint 3D T31016	0.013	0.014	0.017	0.083	0.120	0.072	0.082	0.110	0.5	0.06	0.30	0.48
RAZOR chamber	0.033	0.066	0.057	0.160	0.150	0.270	0.300	0.370	0.2	0.06	0.30	0.48
IBA CC04	0.025	0.025	0.050	0.044	0.180	0.091	0.110	0.160	0.2	0.06	0.30	0.48
RAZORnano chamber	0.480	0.200	0.260	0.310	0.610	0.048	0.053	0.059	0.2	0.06	0.30	0.48
Diode P T60016	0.020	0.052	0.024	0.180	0.590	0.010	0.011	0.011	0.5	0.06	0.30	0.48
Diode E T60017	0.037	0.046	0.017	0.099	0.620	0.028	0.031	0.033	0.5	0.06	0.30	0.48
RAZOR diode	0.190	0.053	0.051	0.059	0.210	0.046	0.052	0.055	0.5	0.06	0.30	0.48
EDGE detector *	0.026	0.026	0.032	/	/	/	/	/	0.5	0.06	0.30	0.48
microDiamond T60019	0.055	0.041	0.017	0.480	0.210	0.053	0.060	0.065	0.5	0.06	0.30	0.48

Table 12. Results of an uncertainty analysis for the field output determined with active detectors. Values are rounded to two significant digits.

* Uncertainties of EDGE detector were not examined because of the unavailability of that detector.

Field size / mm	D(0,0,0) / a.u.	D(0, 0, 0.3) / a.u.
4 mm	1.000	0.992
8 mm	1.000	0.995
16 mm	1.000	0.999

Table 13. Reduction in detected signal if the detector's reference point is misaligned by 0.3 mm from the isocentre in the z direction.

The main source of uncertainties in the relative dosimetry of Gamma Knife's narrow photon beams with active detectors comes from the uncertainty in the positioning of the detector's effective point of measurements at the isocentre (Type B uncertainty) (Table 12.). The assumption is made, that for every measurement, a detector's effective point of measurement was misaligned from the device's isocentre by 0.3 mm in the z direction. This displacement represents the maximum allowed geometrical error of the Gamma Knife Perfexion device and displacement in the z direction was assumed since maximum uncertainty is expected in that direction. It is shown that the difference in the detector's response is increasing with a decrease in the field size (Table 13.). Because 16 mm and 8 mm fields do have dose plateau, a displacement from the isocentre will affect the absorbed dose by 0.1% and 0.51% respectively. On the other hand, the dose plateau of the 4 mm field is non-existent therefore displacement from the field centre will result in the 0.83% reduction from the maximum absorbed dose. To calculate the uncertainty in the detector positioning σ_{PPS} the uniform probability distribution was used (Equation 3.21.).

In the IAEA TRS-483 CoP it is stated that the selection of detectors for small field dosimetry should be such that its uncertainty in short-term stability, dose and dose-rate linearity and leakage current all should be lower than 0.1%.¹⁰ From our data (Table 12.), detectors stability (Type A uncertainty) is, in general, higher for ionization chambers comparing to semiconductor or diamond detectors, with very few exceptions. Moreover, the short-term stability of an ionization chamber increases with the size of its effective volume. This can be noticed in the RAZORnano chamber with the smallest effective volume being a single detector that does not satisfy IAEA criteria on detector short-term stability (<0.1%) which is in line with the recent findings of multicentre experimental study.⁸⁹ Interestingly, detector stability i.e. measurement reproducibility, cannot be correlated with the field size, with a significant

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number of detectors having the largest uncertainty in the 16 mm or 8 mm field size. In addition, detectors have shown a higher uncertainty in dose-rate dependence than in absorbed dose linearity.

Out of eleven investigated detectors in narrow Co-60 beams, five of them have values in linearity of absorbed dose higher than IAEA recommends (<0.1%), and ten have their values higher when it comes to the dose-rate linearity. In previous small field dosimetry study involving high energy X-rays, less than 50% of investigated detectors have satisfied IAEA criteria for linearity as well as dose-rate dependence. It has been shown that if 90% of investigated detectors were to satisfy criteria the tolerance level must be increased.⁸⁹ Relatively high values in the absorbed dose linearity in narrow Co-60 beams for microDiamond detector could be a symptom of the degradation of the detector's characteristic due to irradiation that these detectors are known for.^{29,74} Values of leakage current are increasing with the decrease in the field size. This is expected since less charge is detected in smaller field therefore the relative contribution of leaked charge is increasing (Equation 3.28.). Out of thirty-three leakage current measurements, (eleven detectors in three fields), six are over 0.1% with four of them being in the smallest 4 mm field size (Table 12.), which is in line with the results of the multicentre experiment study.⁸⁹

Finally, results of this study confirm previous findings involving detector's linearity and doserate dependence in high energy X-rays. Notably, the fact that the limit of 0.1% is too low for much of the investigated detectors. Therefore, the revision and increase of this limit in order to be achievable in experimental studies should be considered.⁸⁹

Combined uncertainties were calculated as square root of quadratically added individual uncertainties (Equation 3.18.), and to calculate expanded uncertainty it was multiplied by the coverage factor (k=2) to ensure a confidence of 95% (Table 14.).⁸⁰

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-	Combined uncertainties / %		Expanded uncertainties (k=2) / %			
Fields size / mm Detector name	16	8	4	16	8	4
Semiflex T31010	0.52	0.60	0.71	1.04	1.20	1.41
Semiflex 3D T31021	0.53	0.61	0.72	1.07	1.22	1.43
PinPoint T31014	0.52	0.60	0.71	1.04	1.20	1.42
PinPoint 3D T31016	0.53	0.61	0.72	1.06	1.21	1.43
RAZOR chamber	0.41	0.52	0.68	0.81	1.05	1.35
IBA CC04	0.29	0.42	0.58	0.59	0.84	1.15
RAZORnano chamber	0.86	0.80	0.90	1.72	1.59	1.79
Diode P T60016	0.79	0.85	0.93	1.56	1.69	1.85
Diode E T60017	0.81	0.86	0.94	1.62	1.72	1.88
RAZOR diode	0.58	0.63	0.73	1.16	1.25	1.46
EDGE detector*	0.50	0.58	0.69	1.01	1.17	1.39
microDiamond T60019	0.73	0.79	0.87	1.46	1.57	1.74

Table 14. Combined and expanded uncertainty for the field output determined with active detectors. Values are rounded to the second significant digit.

*Calculated uncertainties of EDGE detector may be smaller than they actually are, since some uncertainties were not examined due to the unavailability of this detector.

5.2. Charged particle equilibrium for different detectors in narrow Co-60 beams used by Gamma Knife

In this section, investigated detectors are evaluated in different Gamma Knife's absorbed dose distributions with respect to the charged particle equilibrium. Tissue-phantom ratio TPR_{20, 10} for Co-60 (k_Q=1) is taken to be 0.56^{48} , and calculated lateral range of secondary electron r_{LCPE} (Equation 2.1.) is 3.883 mm.

If charged particle equilibrium exists at the center of a detector's effective volume with detector's diameter d placed inside absorbed dose distribution with dimensions $FWHM_{msr,clin}$ then $FWHM_{msr,clin} \ge 2r_{LCPE} + d$. $FWHM_{msr,clin}$ values are determined by the device's manufacturer with Monte Carlo simulations (Pegasos based system).

		Dose profile FWHM / mm						
		16 x	16 z	8 x	8 z	4 x	4 z	
Detector name	$2r_{LCPE} + d$ / mm	21.75	17.44	11.06	9.80	6.16	5.04	
Semiflex T31010	14.67	СРЕ	СРЕ	No CPE	No CPE	No CPE	No CPE	
Semiflex 3D T31021	12.56	СРЕ	СРЕ	No CPE	No CPE	No CPE	No CPE	
PinPoint T31014	11.16	СРЕ	СРЕ	No CPE	No CPE	No CPE	No CPE	
PinPoint 3D T31016	12.01	СРЕ	СРЕ	No CPE	No CPE	No CPE	No CPE	
RAZOR ch.	10.76	СРЕ	СРЕ	No CPE	No CPE	No CPE	No CPE	
IBA CC04	12.56	СРЕ	СРЕ	No CPE	No CPE	No CPE	No CPE	
RAZORnano ch.	10.76	СРЕ	СРЕ	СРЕ	No CPE	No CPE	No CPE	
Diode P T60016	8.32	СРЕ	СРЕ	СРЕ	СРЕ	No CPE	No CPE	
Diode E T60017	8.32	СРЕ	СРЕ	СРЕ	СРЕ	No CPE	No CPE	
RAZOR diode	8.36	СРЕ	СРЕ	СРЕ	СРЕ	No CPE	No CPE	
EDGE detector	8.56	СРЕ	СРЕ	СРЕ	СРЕ	No CPE	No CPE	
microDiamond T60019	9.96	СРЕ	СРЕ	СРЕ	No CPE	No CPE	No CPE	

Table 15. Results of CPE analysis for different detectors.

Absorbed dose distribution of Gamma Knife is ellipsoidal, therefore, to evaluate different detectors inside such distributions it is necessary to do it in two planes: XY and XZ. Moreover, the XY plane is circular while XZ and YZ planes are elliptical. Because of this, an evaluation of CPE for a given detectors in XY plane can be done with the respect to one dose profile (i.e. x-axis). On the other hand, to evaluate CPE in XZ plane, an evaluation must be done with the respect to z-axis dose profile. In addition, due to the difference in dimensions of dose profiles, if CPE is violated in XY plane it is also violated in XZ plain. Only detectors used for the field output factor determination are considered in this analysis, and their dimensions are determined from the detector's schematics.^{61,63–65}

From CPE condition, the maximum allowed diameter of a detector for CPE to exist at the centre is: 9.674 and 2.034 mm for 16 mm an 8 mm field size respectively. So far, not a single detector can fulfil the CPE condition for a 4 mm field since the lateral range of secondary electrons is larger than the FWHM value.

5.3. Ellipsoid absorbed dose model of Gamma Knife Perfexion

In this section results of an ellipsoid absorbed dose model used for the determination of volume-averaging correction factor are presented. Normalized fitted dose profiles (Equation 3.4.) can be seen in Figure 29.

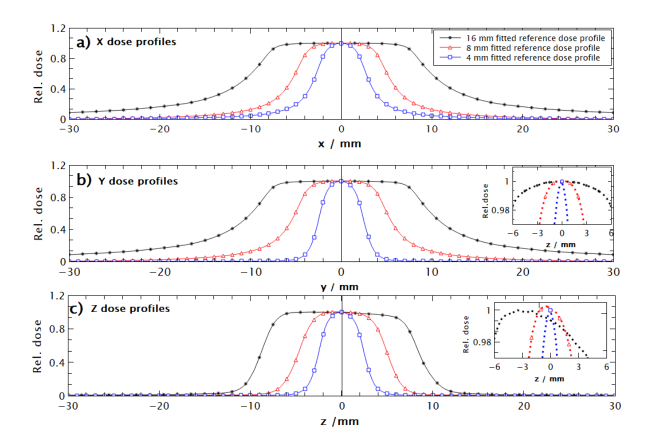


Figure 29. Normalized fitted dose profiles for a 16 mm, 8 mm and 4 mm field size on LGK Perfexion. Data were calculated using Monte Carlo calculation by the device's manufacturer for a shot at the centre of a spherical phantom with a radius of 80 mm. a) dose profile on the x-axis, b) dose profile on the y-axis with the insert showing symmetry in y dose profile, and c) dose profile on the z-axis with insert of enlarged centre of the profile showing asymmetry.

In MATLAB (TheMathWorks Inc, USA), using Equation 3.9., the visualization of different Gamma Knife's absorbed dose distributions was created as shown in Figure 30.

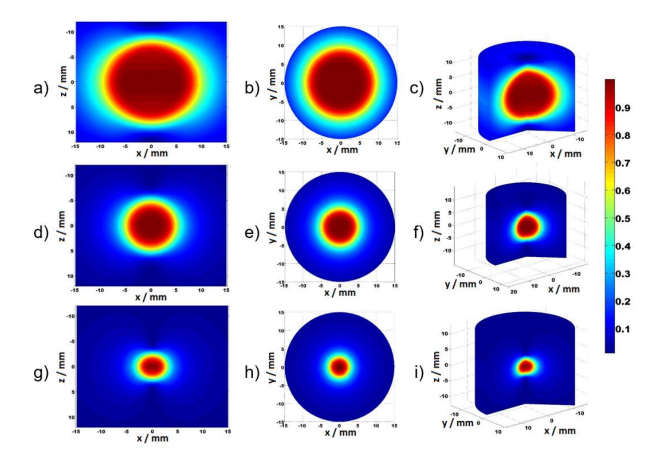


Figure 30. A Gamma Knife Perfexion field size simulation using the ellipsoid absorbed dose model. From a)-c): XZ, XY plane and 3D simulation of 16 mm absorbed dose distribution, from d)-f): XZ, XY plane and 3D simulation of 8 mm absorbed dose distribution, from g)-i): XZ, XY plane and 3D simulation of 4 mm absorbed dose distribution. For the XZ plane y=0, and z=0 for the XY plane.

An ellipsoid absorbed dose model accuracy was verified against the LGP for 1700, 783 and 378 points for 16 mm, 8 mm, and 4 mm field sizes ranging from 99% to 19% of the relative dose. The accuracy of a model with respect to an absorbed dose was the highest for the 4 mm field size, with the 16 mm field having the largest difference from the LGP (Figure 31.). For 4 mm field size the largest difference of all analysed points was around 0.5%. The difference in the absorbed doses for an 8 mm field size is 1.5% at the 80% isodose and it is gradually decreasing for lower isodoses. For 16 mm field size, the difference is the smallest at the 90% isodose volume and it is gradually increased to 2.25% for 30% isodose volume. Interestingly, the model displays an increase in accuracy with a decrease in the field size.⁵⁴ This can be contributed to the lower asymmetry of the 4 mm z-axis dose profile and consequently better fitting.

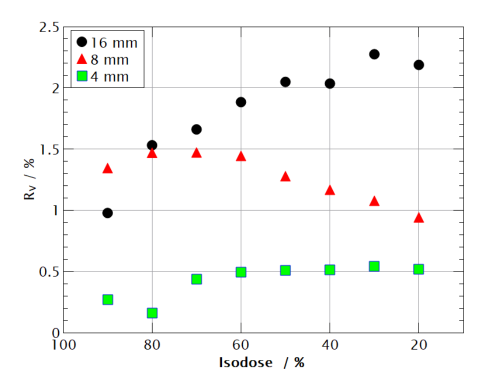


Figure 31. A difference in an absorbed dose between the LGP and the model for different ellipsoid volumes beginning at the isocentre and ending at the givens dose value.

It is beneficial that the model increases its accuracy with a decrease in the field size because the largest volume-averaging is expected in the smallest field sizes. In this analysis, we conclude that the ellipsoids absorbed dose model can be used for the determination of volume-averaging correction factors.

5.4. Volume-averaging correction factors

Volume-averaging correction factors $(k_{vol})_{Q_0}^{f_{msr,clin}}$ for detectors listed in Table 1. are calculated using the Equation 2.6. and are presented in Table 16. and Figure 32.⁵⁴ Mean uncertainties in the absorbed dose predicted by the model are 1.79%, 1.15% and 0.43% for 16 mm, 8 mm and 4 mm field size and corresponding to Type A uncertainties (Figure 31). Since volume-averaging correction is calculated using a mathematical model, and not measured, Type B uncertainties were not examined.

	$(k_{vol})_{Q_0}^{f_{msr,clin}}$				
Detector name	16 mm	8 mm	4 mm		
Semiflex T31010	1.006±0.018	1.042±0.012	1.627±0.007		
Semiflex 3D T31021	1.005±0.018	1.019±0.012	1.304±0.006		
PinPoint T31014	1.005±0.018	1.011±0.012	1.181±0.005		
PinPoint 3D T31022	1.003±0.018	1.011±0.012	1.101±0.005		
Diode P T60016	1.001±0.018	1.001±0.012	1.008±0.004		
Diode E T60017	1.001±0.018	1.001±0.012	1.008±0.004		
microDiamond T60019	1.000±0.018	1.002±0.012	1.021±0.004		
RAZOR diode	1.001±0.018	1.000±0.012	1.004±0.004		
EFD 3G-pSi	1.001±0.018	1.001±0.012	1.018±0.004		
EDGE detector	1.004±0.018	1.003±0.012	1.007±0.004		
RAZOR chamber	1.003±0.018	1.006±0.012	1.074±0.005		
IBA CC04	1.007±0.018	1.014±0.012	1.156±0.005		
Exradin W2 1x1	1.003±0.018	1.002±0.012	1.010±0.004		
Exradin W2 1x3	1.005±0.018	1.005±0.012	1.042±0.004		

Table 16. A result of volume-averaging correction factors $(k_{vol})_{Q_0}^{f_{msr,clin}}$ with uncertainties $\sigma_{(k_{vol})_{Q_0}^{f_{msr,clin}}}$ for different detectors in different fields of Gamma Knife Perfexion. Detectors whose volume-averaging correction factors are marked in red are not recommended by IAEA TRS-483 CoP for dosimetry at that field size.

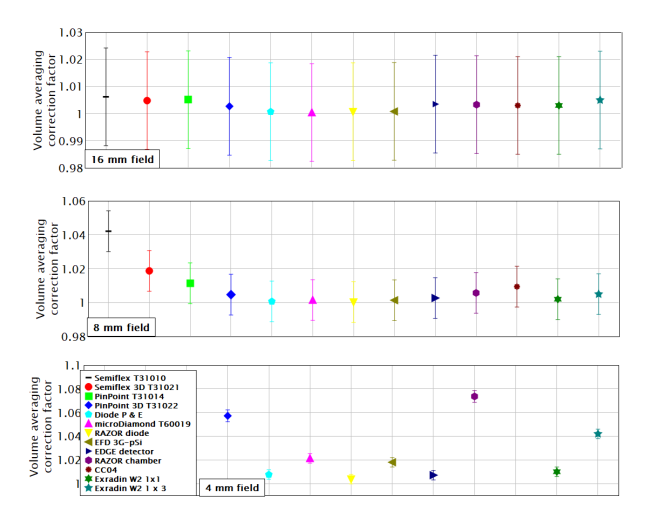


Figure 32. Volume-averaging correction factors $(k_{vol})_{Q_0}^{f_{msr,clin}}$ in msr and clin fields for a different detector with uncertainties $\sigma_{(k_{vol})_{Q_0}^{f_{msr,clin}}}$. Diode P T60016 and Diode E T60017 are represented by one data point since their effective volumes are identical. Detectors with volume-averaging correction factors larger than 1.1 are excluded from 4 mm field graph for better graph readability.

From the theory it is expected that for all detectors volume-averaging correction factor will decrease with the increase of the field size and any deviations from this are due to the imperfection in the ellipsoid absorbed dose model. This explains why Diode P T60016, Diode E T60017, microDiamond T60019, RAZOR diode, EDGE detector and Exradin W2 1x1 have larger volume-averaging correction factor for 16 mm field size than for 8 mm field (Table 16.). This increase in the volume-averaging for larger field size is rather small and can be neglected (Table 16.).

Presented results (Table 16. and Figure 32.) confirm that ionization chambers have a larger volume-averaging correction factor than other types of detectors due to the size of their effective volume. Semiflex T31010, a reference class ionization chamber, has the largest volume-averaging correction factor for all field sizes (V=125 mm²), followed by Semiflex 3D T31021 (V=70 mm³); 1.006, 1.042, 1.627, and 1.005, 1.019, 1.304 for 16 mm, 8 mm, and 4 mm field respectively. Although it has a larger volume, PinPoint 3D T31022 has lover volumeaveraging correction than PinPoint T31014 (16 mm³ vs. 15 mm³) for all field sizes. This is due to the geometry of these chambers where PinPoint T31014 is longer and narrower, and PinPoint 3D T31022 is more compact. A length-to-diameter ratio R, is 2.5 and 1.23 for PinPoint and PinPoint 3D respectively. Positioning the detector with its long axis in the z direction makes it more influenced by the asymmetry, and dose gradient of the z-axis dose profile over its volume, therefore, increasing the volume-averaging correction factor (Figure 33.). Importantly, the z-dose profile also has the lowest FWHM value (Figure 29.). By contrast, at the radial dimensions of these chambers x, and y-axis dose profiles are identical, therefore they could not be the reason behind the difference in volume-averaging (Figure 34.). Moreover, the radius of examined chambers is similar therefore the difference in the volumeaveraging must be because of the z-axis dose profile. The larger volume-averaging correction factor at the 16 mm field compared with the 8 mm field for some detectors is because of the lower accuracy of the model for that field size (Figure 31.) with a difference of around 0.1% (Table 16.). For a 4 mm field size, IBA CC04 has volume-averaging comparable with PinPoint although its effective volume is almost three times larger (Table 16.). Again, this is due to the longer effective volume in the z-direction of PinPoint T31014 than IBA CC04 (5 mm compared to 3.6 mm) as shown in Table 16. i.e., IBA CCO4 is a more compact chamber with an active length-to-diameter ratio equal to 0.9. These results are in line with the similar findings of investigation on detector-specific correction factor with respect to different orientation in high-energy X-ray beams.⁵⁵

RAZOR chamber is the most compact ionization chamber in this study with an effective volume of 10 mm³ and, as a result, it has the smallest volume-averaging correction factor out of all ionization chambers, for all field sizes. RAZORnano chamber has a unique semi-spherical shape with 3 mm³ of an effective volume. Modelling such geometry inside ellipsoid absorbed dose

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model resulted in significant uncertainties in volume averaging correction factor, therefore, it was left out of this research.

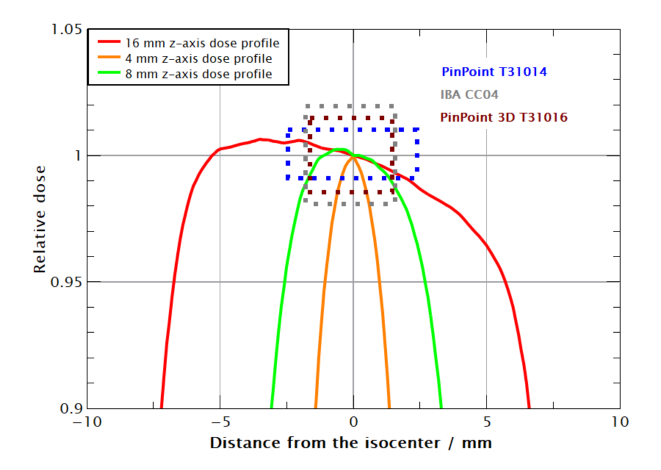


Figure 33. The relative dose profiles on the z-axis with geometries of three detectors. The gradient on the z-axis, for detectors that are longer, creates an increase in the volume-averaging correction factor. Dose profiles are normalized at z=0 mm.

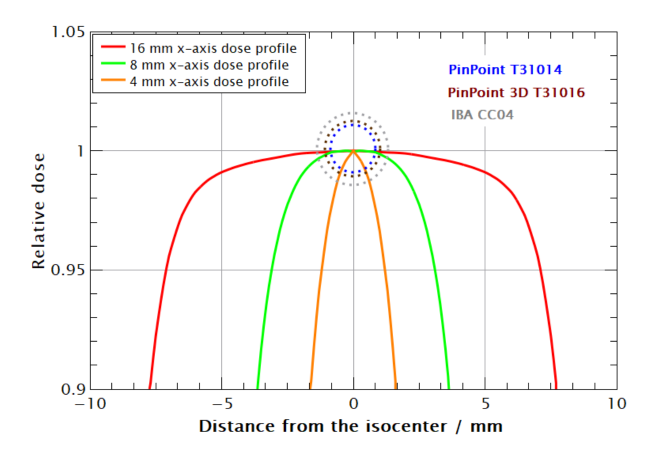


Figure 34. The relative dose profiles on the x-axis with geometries of three detectors. The gradient on the z-axis more influences detectors that are longer creating an increase in the volume-averaging correction factor. Dose profiles are normalized at x=0 mm.

There is an open issue whether the detectors should be placed in a phantom parallel or perpendicular to the x-axis when performing dosimetry of Gamma Knife's narrow photon beams. Therefore, an investigation related to the proper orientation of the detector in the phantom with respect to the volume-averaging should be performed. Since the z-axis dose profile contributes the most to volume-averaging, a detector orientation perpendicular to the z-axis should be considered to limit its influence. However, some studies using Monte Carlo simulations have calculated detector-specific correction factors for the ionization chambers that were placed parallel and perpendicular to the z-axis in the ABS and solid water phantom. Lower detector-specific correction factors were found in the parallel orientation.⁹⁰ Although relevant for the dosimetry in small fields, for dose profile measurements our research into volume-averaging indicated that positioning detectors perpendicular to the scanning dose profile is better since volume-averaging is the only effect influencing these types of

measurements. A difference in the geometries of the detector placement in the ABS and solid water phantom can be seen in Figure 35.

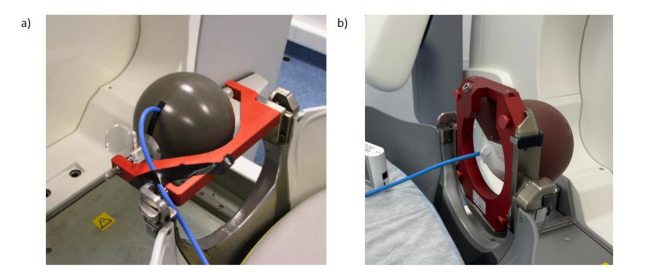


Figure 35. Measuring geometries on Gamma Knife a) dosimetry in ABS phantom with detector places in XY plane perpendicular to the z- axis, and, b) detector in solid water phantom positioned parallel to the z-axis.

Due to the small size of an effective volume and consequently smaller volume-averaging effect, semiconductor detectors are the ideal type of detectors for dosimetry in small fields.²⁰ This fact is supported by this study for a large majority of explored semiconductor detectors. The single exception is the EFD diode which has a relatively large volume (0.16 mm³), therefore the correction factor for the 4 mm field size is 1.018. Diode E T60016 and Diode P T60017 have equal volume-averaging correction factors since their effective volumes are identical, and the only difference is shielding for the filtration of low-energy photons. The microDiamond T60019 is characterized by the smallest volume out of all studied detectors, however, its correction factor for the 4 mm field is relatively large being 1.021. This is due to the shape of its effective volume. It is unique in a way that it is a very thin $(1 \mu m)$ disk with a large diameter of 2.2 mm. This design makes it more sensitive to dose gradients in the XY plane which starts to be noticeable at a 4 mm field increasing the correction for volume-averaging. From this, one can see that although the magnitude of the detector's effective volume is important, when it comes to dosimetry of GK narrow photon beams, the shape of that volume also has an important role. More precisely, the length-to-diameter ratio of a detector has s fundamental role in volume averaging correction factor⁵⁵. The manufacturer denotes plastic scintillator detectors Exradin W2 1x1 and 1x3 as perturbation-free detectors since they are tissue–equivalent. However, the effect of volume-averaging could not be neglected for them (Table 16.). In addition, following IAEA TRS-483 CoP we have selected detectors that are not suitable for dosimetry of Gamma Knife Perfexion photon beams according to their volume-averaging correction factor. Therefore, Semiflex T31010, Semiflex 3D T31021, PinPoint T31014, PinPoint 3D T31022, CC04, and RAZOR chamber are not recommended for dosimetry in 4 mm field since their volume-averaging correction factor is larger than 1.05 (Table 16.). Finally, we have calculated the contribution of the volume-averaging correction factor in the field output correction factor $k_{\%vol}$ for detectors listed in IAEA TRS-483 CoP. For all studied detectors, volume-averaging contribution is larger than the total field output correction factor they over-respond to low energy scattered photons and their total correction factors are lower than 1.000 due to the fluence perturbations of these detectors.

A volume-averaging perturbation correction factor $(k_{vol})_{Q_0,Q_0}^{f_{msr},f_{clin}}$ calculated (Equation 4.11.) for the detectors that are listed in IAEA TRS-483 CoP and the results are shown in Table 17. As previously, detectors that are marked with red are not suitable for dosimetry at that field size.

	$(k_{vol})_{Q_0,Q_0}^{fmsr,f_{clin}}$					
Detector name	16 mm	8 mm	4 mm			
Semiflex T31010	1.000	1.036±0.022	1.617±0.007			
Semiflex 3D T31021	1.000	1.014±0.022	1.298±0.006			
PinPoint T31014	1.000	1.006±0.022	1.175±0.005			
PinPoint 3D T31022	1.000	1.001±0.022	1.091±0.005			
Diode P T60016	1.000	0.999±0.022	1.007±0.004			
Diode E T60017	1.000	0.999±0.022	1.007±0.004			
microDiamond T60019	1.000	1.001±0.022	1.020±0.004			
RAZOR diode	1.000	0.999±0.022	1.003±0.004			
EFD 3G-pSi	1.000	1.000±0.022	1.017±0.004			
EDGE detector	1.000	0.999±0.022	1.004±0.004			
RAZOR chamber	1.000	1.003±0.022	1.071±0.005			
IBA CC04	1.000	1.007±0.022	1.147±0.005			
Exradin W2 1x1	1.000	0.999±0.022	1.007±0.004			
Exradin W2 1x3	1.000	1.000±0.022	1.037±0.004			

Table 17. A result of a volume-averaging perturbation in clinical and machine-specific reference fields $(k_{vol})_{Q_0,Q_0}^{f_{msr},f_{clin}}$ for different detectors with uncertainties $\sigma_{(k_{vol})_{Q_0,Q_0}^{f_{msr},f_{clin}}}$. Investigated detectors marked in red are not recommended for dosimetry at that field size according to IAEA TRS-483 CoP guidelines for volume-averaging criteria.¹⁰

	$k_{Q_0,Q_0}^{fmsr,f_{clin}}$			$k_{\% vol}/\%$			
Detector name	16 mm	8 mm	4 mm	16 mm	8 mm	4 mm	
Semiflex T31010	1.004	N.A.	N.A.	99.6	N.A.	N.A.	
PinPoint T31014	1.000	1.030	N.A.	100.0	98.4	N.A.	
Diode P T60016	1.000	0.981	0.965	100.0	101.8	104.4	
Diode E T60017	1.000	0.996	0.985	100.0	100.3	102.2	
microDiamond T60019	1.000	1.005	0.993	100.1	99.6	102.9	
IBA CC04	1.021	N.A.	N.A.	97.9	N.A.	N.A.	

Table 18. Field output correction factors $k_{Q_0,Q_0}^{fmsr,f_{clin}}$ from IAEA TRS-483 CoP and contribution of volume-averaging perturbation to a total correction $k_{\%vol}$ for different detectors. N.A. stands for not available in IAEA TRS-483 CoP.

5.5. Field output factors and detector-specific correction factors

A set of EBT3 film irradiated in the XY plain with different doses using a 16 mm field size for creation of a calibration curve can be seen in Figure 36.

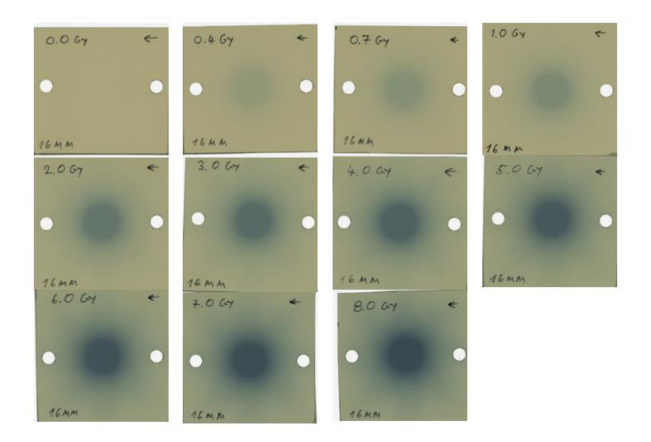


Figure 36. EBT3 films irradiated in the XY plane with different absorbed doses for creation of a calibration curve.

The calibration curve using Dević fit function relating netOD with an absorbed dose can be seen in Figure 37. with fit values in Table 19.

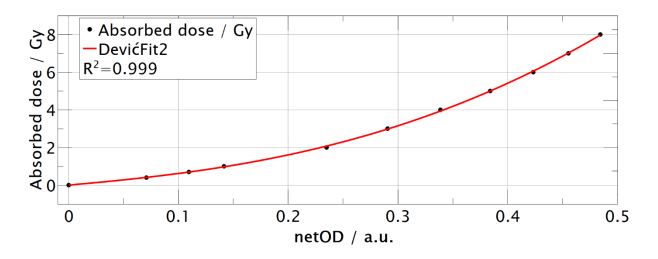


Figure 37. Calibration curve for EBT3 films.

Coefficients	Value	Uncertainties			
а	5.17	0.20			
b	35.30	2.17			
n	2.514	0.088			
$\frac{1}{D(netOD)} = a \cdot netOD + b \cdot netOD^{n}$					

Table 19. Fit coefficients for EBT3 calibration curve.

Films irradiated for the determination of field output factors are presented in Figure 38. Field output factor $\Omega_{Q_0,Q_0}^{f_{clin},f_{msr}}$ were determined on ROI: 4, 2, and 0.8 mm for 16 mm, 8 mm and 4 mm field size respectively. Since EBT3 film is considered perturbation free, the detector-specific correction factor is equal to one, field output factors were determined by converting the mean pixel value on ROI to absorbed dose and by dividing the absorbed dose in the clinical

field with dose in msr field using $\Omega_{Q_0,Q_0}^{f_{clin},f_{msr}} = \frac{D_{w,Q_0}^{f_{clin}}}{D_{w,Q_0}^{f_{msr}}}$. Result of field output factors and its difference from Monte Carlo calculated values²⁰, values are presented in Table 20.

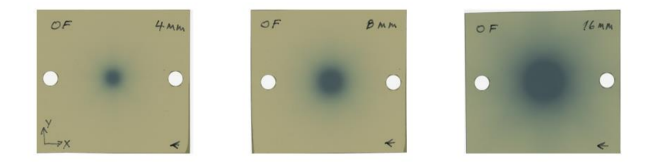


Figure 38. EBT3 films irradiated 2.69 minutes in XY plane for field output factor determination.

Field size / mm	$\left(\Omega_{Q_0,Q_0}^{f_{clin},f_{msr}}\right)_{EBT3}$	$\left(\Omega_{Q_0,Q_0}^{f_{clin},f_{msr}}\right)_{MC}$	EBT3 diff. from MC / %
4	0.826±0.064	0.814	1.474
8	0.889±0.053	0.900	-1.222

Table 20. Field output factors $\Omega_{Q_0,Q_0}^{f_{clin},f_{msr}}$ determined using EBT3 film, Monte Carlo²⁰ simulations and difference of EBT3 values from Monte Carlo calculated values.

Field output factors determined using EBT3 film have shown a good agreement with the Monte Carlo calculated values, with a mean absolute difference of less than 2%. However, it should be pointed out that field output factors determined with the EBT3 film have uncertainty up to 6% due to the noise and postirradiation handling (Table 11.).^{17,42,43,76}

To analyse the performance of different detectors in narrow Co-60 beams, field output factors are determined with twelve different detectors. The results of such measurements are presented in Table 21. Comparison analysis of field output factors obtained with the active detectors and the ones obtained with the EBT3 film and Monte Carlo is presented in Table 22., together with the calculated detectors specific field output correction factors $k_{Q_0,Q_0}^{f_{clin},f_{msr}}$.

	$M_{Q_0}^{fmsr,clin}/nC$			$\Delta M_{Q_0}^{f m sr, clin} / nC$			$\Omega^{f_{clin},f_{msr}}_{Q_0,Q_0}\pm\sigma_{\Omega^{f_{clin},f_{msr}}_{Q_0,Q_0}}$			
Field size / mm Detector name	16	8	4	16	8	4	16	8	4	
Semiflex T31010	7.273	5.717	2.430	0.077	0.083	0.048	1.000	0.786±0.014	0.334±0.007	
Semiflex 3D T31021	3.669	3.131	1.645	0.040	0.046	0.032	1.000	0.854±0.016	0.449±0.010	
PinPoint T31014	0.913	0.799	0.580	0.010	0.012	0.011	1.000	0.877±0.016	0.636±0.014	
PinPoint 3D T31016	0.856	0.756	0.578	0.009	0.011	0.011	1.000	0.880±0.016	0.675±0.015	
RAZOR chamber	0.596	0.528	0.433	0.005	0.007	0.008	1.000	0.885±0.014	0.726±0.015	
RAZORnano chamber	0.208	0.187	0.170	0.034	0.031	0.026	1.000	0.900±0.022	0.815±0.023	
IBA CC04	1.972	1.716	1.140	0.001	0.002	0.003	1.000	0.870±0.012	0.578±0.011	
Diode P T60016	20.568	18.676	17.548	0.328	0.352	0.333	1.000	0.908±0.022	0.853±0.024	
Diode E T60017	21.428	19.131	18.139	0.348	0.365	0.420	1.000	0.893±0.022	0.847±0.024	
RAZOR diode	7.549	6.774	6.397	0.088	0.102	0.127	1.000	0.897±0.017	0.847±0.020	
EDGE detector	57.490	51.070	48.790	0.587	0.730	0.946	1.000	0.888±0.016	0.849±0.019	
microDiamond T60019	2.254	1.997	1.851	0.033	0.035	0.041	1.000	0.886±0.020	0.821±0.022	

Table 21. Results of an average detector's response in clinical and machine-specific reference fields $M_{Q_0}^{f msr,clin}$, a calculated value of signal uncertainty $\Delta M_{Q_0}^{f msr,clin}$ (Table 14.), uncorrected field output factors $\Omega_{Q_0,Q_0}^{f_{clin},f_{msr}}$ with total uncertainties $\sigma_{\Omega_{Q_0,Q_0}^{f_{clin},f_{msr}}}$.

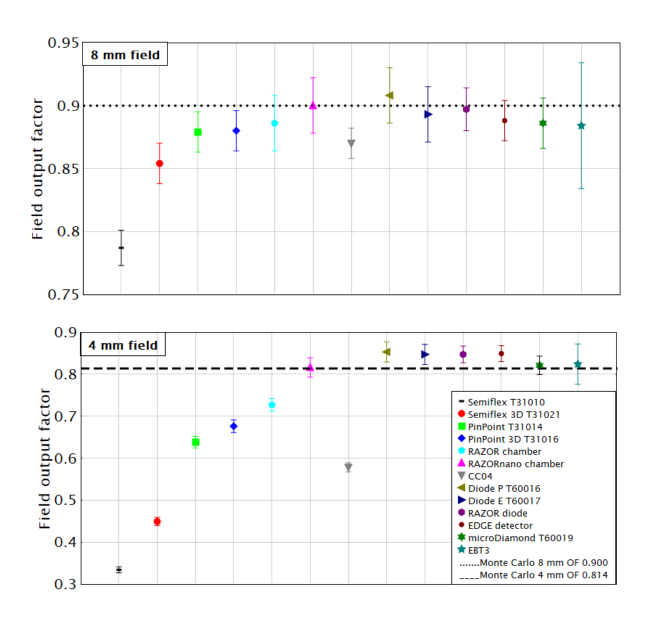


Figure 39. Results of uncorrected field output factors $\Omega_{Q_0,Q_0}^{f_{clin},f_{msr}}$ for all investigated detectors with uncertainties $\sigma_{\Omega_{Q_0,Q_0}^{f_{clin},f_{msr}}}$ in 8, and 4 mm field. Dotted and dashed lines represent Monte Carlo values for 8 mm field (0.900) and 4 mm field (0.814) respectively.

	$\Omega^{f_{clin},f_{msr}}_{Q_0,Q_0}$		$\Omega^{f_{clin},f_{msr}}_{Q_0,Q_0}$ diff. from MC / %		$ \begin{pmatrix} k_{Q_0,Q_0}^{f_{msr},f_{clin}} \end{pmatrix}_{MC} \\ \pm \sigma_{\left(k_{Q_0,Q_0}^{f_{msr},f_{clin}}\right)_{MC}} $		$\Omega^{f_{clin},f_{msr}}_{Q_0,Q_0}$ diff. from EBT3 / %		$ \begin{pmatrix} k_{Q_0,Q_0}^{f_{msr},f_{clin}} \end{pmatrix}_{EBT3} \\ \pm \sigma_{\left(k_{Q_0,Q_0}^{f_{msr},f_{clin}}\right)_{EBT3}} $	
Field size / mm Detector name										
	8	4	8	4	8	4	8	4	8	4
Semiflex T31010	0.786	0.334	-12.6	-58.9	1.145±0.021	2.436±0.054	-11.5	-59.6	1.131±0.054	2.470±0.130
Semiflex 3D T31021	0.854	0.449	-5.1	-44.8	1.054±0.019	1.814±0.041	-3.9	-45.6	1.041±0.050	1.840±0.093
PinPoint T31014	0.877	0.636	-2.3	-21.7	1.026±0.018	1.279±0.028	-1.1	-22.8	1.013±0.049	1.298±0.066
PinPoint 3D T31016	0.880	0.675	-2.2	-17.0	1.023±0.019	1.205±0.027	-1.0	-18.2	1.010±0.049	1.223±0.062
RAZOR chamber	0.885	0.726	-1.6	-10.7	1.016±0.016	1.121±0.023	-0.3	-12.0	1.004±0.046	1.137±0.054
RAZORnano chamber	0.900	0.816	0	0.2	1.000±0.025	0.998±0.028	1.2	-1.2	0.988±0.047	1.013±0.050
IBA CC04	0.870	0.578	-3.2	-29.0	1.034±0.014	1.408±0.026	-2.1	-30.0	1.022±0.047	1.428±0.068
Diode P T60016	0.908	0.853	0.9	4.8	0.991±0.024	0.954±0.027	2.1	3.3	0.979±0.047	0.968±0.049
Diode E T60017	0.893	0.847	-0.8	4.0	1.008±0.025	0.962±0.027	0.4	2.5	0.996±0.048	0.976±0.050
RAZOR diode	0.897	0.847	-0.3	4.1	1.003±0.019	0.961±0.022	0.9	2.5	0.991±0.048	0.975±0.050
EDGE detector	0.888	0.849	-1.3	4.3	1.013±0.018	0.959±0.021	-0.1	2.8	1.001±0.048	0.973±0.050
microDiamond T60019	0.886	0.821	-1.6	0.9	1.016±0.023	0.991±0.026	-0.3	-0.6	1.003±0.048	1.006±0.051

Table 22. Results in difference in uncorrected field output factors $\Omega_{Q_0,Q_0}^{f_{clin},f_{msr}}$ and detector-specific field output correction factor $k_{Q_0,Q_0}^{f_{msr},f_{clin}}$ with their uncertainties for different detectors when compared with Monte Carlo and EBT3 film values. Result are rounded to the second significant digit.

For the discussion of data from Table 22., detectors will be subdivided into two groups; firstly, ionization chambers will be discussed and then the remaining detectors. Similarly, the results of the field output factors for the 4 mm field will be discussed first since all perturbation effects are noticeable at that field size, afterwards, the results for the 8 mm field will be examined. Differences in field output factors determined using active detectors from values determined with the EBT3 film and Monte Carlo are due to three effects that add up, these are: lack of lateral charged particle equilibrium, over-response/under-response of high/low Z material, and volume-averaging effect.

5.5.1. Field output factors of Gamma Knife's 4 mm field

Results of a field output factors $\Omega^{f_{clin},f_{msr}}_{Q_0,Q_0}$, for 4 mm field, obtained with the ionization chambers; Semiflex T31010 (V=125 mm³), Semiflex 3D T31021 (V=70 mm³), PinPoint T31014 (V=15 mm³), PinPoint 3D T31016 (V=16 mm³), RAZOR chamber (V=10 mm³) and CC04 (V=40 mm³) shown a substantial difference both from the Monte Carlo and EBT3 values (Table 18.). For all mentioned detectors field output factors are underestimated in a range of -60% for Semiflex T31010 to -12% for the RAZOR chamber. Moreover, the underestimation of a response is directly proportional to the size of a chamber with RAZORnano chamber, with the smallest volume (V=3 mm³), having the best result. For these detectors, in 4 mm field size, all the abovementioned perturbations have a part in the resulting underestimation of field output factors. Loss of charged particle equilibrium is one of those reasons. Presented data (Table 15.) shows that the field must expand at least 3.883 mm from the chamber's edges (Figure 7.) to preserve CPE in water for CO-60 beams. For all studded detectors, in 4 mm field, this condition in not fulfilled. The second perturbation is the significant volume-averaging effect $(k_{vol})_{Q_0,Q_0}^{f_{msr},f_{clin}}$ (Table 17.) All ionization chambers, except the RAZORnano chamber, have large effective volumes relative to the size of a 4 mm field. Because of this, the detector will average its signal over the volume, resulting in a decrease in measured detector response. This is most noticeable for the largest chambers such as Semiflex, Semiflex 3D and CC04 with differences of -58.9%, -44.8% and -29.0% respectively, compared to the Monte Carlo values. They have the largest detector-specific correction factors $k_{Q_0,Q_0}^{f_{msr,f_{clin}}}$ and volume-averaging is one of the main reasons behind this difference (Table 17.). Ionization chambers presented in this study, except RAZORnano, are not recommended for dosimetry of Gamma Knife's 4 mm field because of the extreme perturbations that they demonstrate. RAZORnano chamber is the best performing one with 0.2% and -1.2% difference from field output factor determined with Monte Carlo and EBT3 films respectively (Table 22.). This is expected since it has the smallest effective volume which will result in lower volume averaging regardless of the loss of CPE (Table 15.).

The next group of detectors include remaining semiconductor detectors as well as a diamond detector. For these detectors, volume-averaging is negligible due to its small volume. Volumeaveraging correction factors $(k_{vol})_{Q_0,Q_0}^{f_{msr},f_{clin}}$ calculated for these detectors and are given in Table 16. In addition, in the 4 mm field size, loss of CPE is present for semiconductor and diamond detectors and should be taken into consideration (Table 15.) However, due to the smaller dimensions of an effective volumes of these detectors one can expect that loss of CPE is not significant as with ionization chambers. It should be emphasised that all semiconductor detectors and diamond detector show a better agreement with the field output factor $\Omega^{f_{clin},f_{msr}}_{Q_0,Q_0}$ determined with the EBT3 film than with the Monte Carlo determined values. Similar studies support these conclusions.²⁰ Moreover, all semiconductor detectors overresponded in a 4 mm field. This is due to the high atomic number of silica Z=14, compared with the effective atomic number of the water Zeff=7.42. Furthermore, semiconductor detectors will additionally over-respond to the low energy scattered photon (Figure 16.). This effect is not emphasized as much in narrow beam geometries of Gamma Knife's collimating system due to the lack of low energy photons, however, it is the reason behind the difference in field output factors between Diode P and E (0.966 and 0.973). They are identical detectors, except for the fact that Diode P has a high density shielding of its effective volume that filters out the low-energy photons. Therefore, Diode P will overrespond by 4.8% and 3.3%, and Diode E by 4.0% and 2.5% when the field output values are compared with the Monte Carlo and EBT3 film for 4 mm field size. In general, Diode E is better suited for the dosimetry of narrow photon beams used by Gamma Knife. Unshielded RAZOR diode and EDGE detector have their detector-specific correction factors close to the ones obtained with Diode E (0.973 and 0.971).

From all detectors used for dosimetry of 4 mm field size, the microDiamond detector has shown the best agreement with the Monte Carlo and EBT3 film results. It is characterised by the small effective volume, high dose-response and tissue equivalence. Therefore, it has the lowest perturbation of all detectors resulting in the detector-specific correction factor being close to unity (Table 22.).¹⁶ From these results, we can conclude that microDiamond T60019 is the best-performing detector in dosimetry of 4 mm Gamma Knife's field.

5.2.2. Field output factors of Gamma Knife's 8 mm field.

The results of field output factors $\Omega^{f_{clin},f_{msr}}_{Q_0,Q_0}$ determined for the 8 mm field are presented in Table 22. All investigated ionization chambers have loss of charged particle equilibrium (Table 15.) and high volume-averaging (Table 16.). CPE is not violated for semiconductor detectors. Diamond detectors violets CPE only if positioned perpendicular to the device's z-axis due to large diameter of its effective volume. Volume-averaging correction $(k_{vol})_{Q_0 Q_0}^{f_{msr,f}c_{lin}}$ for semiconductor and diamond detector in an 8 mm field is negligible. The result of field output factors obtained with the ionization chambers still show a significant difference from the reference values. On the other hand, for small-volume chambers this difference is low, so they can be used for dosimetry of such field sizes. Semiflex T31010 and Semiflex 3D T31021 still have large differences from the Monte Carlo values and are underestimating field output factors by 12.6% and 5.1%, respectively. They are not suited for the dosimetry in an 8 mm field size. On the other hand. PinPoint, PinPoint 3D and RAZOR chamber have shown a good agreement with the EBT3 film with all underestimating field output factor by less than 1.1%. CC04 has a difference in field output factor values of -3.2% and -1.1% from Monte Carlo and EBT3 film values. All ionization chambers have better agreement with the EBT3 film than with the Monte Carlo, except for the RAZORnano chamber, which over-responded by 1.2% when compared with the EBT3 film.

Field output factors determined with Diode P and RAZOR diode are closer in value to Monte Carlo values, while the remaining semiconductor detectors are more consistent with the EBT3 film results. Overresponse of Diode P and RAZOR diode is due to high density shielding of its effective volume. This can explain why they are ones of the few detectors with their values closer to Monte Carlo results. Diode E and EDGE have similar values of detector-specific correction factor $k_{Q_0,Q_0}^{f_{msr},f_{clin}}$, when compared to the EBT3 film, being 0.996 and 1.001, respectively. For 8 mm field, ionization chambers have their field output factors in better agreement with the Monte Carlo values. As previously, microDiamond has shown a good agreement with reference EBT3 film values with a correction factor close to unity, 1.003, for 8 mm field.

In this study of field output factor determination, one can see how different detector types behave in different field sizes. Ionization chambers are well-established detectors with an excellent signal-to-noise ratio, however, most of them are not suitable for the dosimetry of small fields.^{12,13,20,26,33} This is due to loss of charged particle equilibrium (Table 15.) and high volume-averaging effect in such fields (Table 16.). Semiconductor detectors are a good selection for small field dosimetry, however, they do have perturbations that must be appropriately considered.^{13,16,17}. The diamond detector has outperformed all detectors with values closest to the reference ones due to its properties.^{17,27,73} Furthermore, most of the detectors agree better in results with the EBT3 films than with the Monte Carlo values. One should keep in mind that uncertainties in the field output factors determined with the EBT3 film are larger than with the active detectors (Table 11. and Table 12.).

5.6. Gamma Knife's Perfexion dose profiles

Different detector will provide dose profile data which may differ significantly. The main cause is the volume-averaging. An increase in detector volume-averaging with a decrease in the size of a dose profile can be seen on all z-axis dose profiles when compared with the dose profiles on x and y - axis for a given field size. The effect is more emphasized as field is decreased down to 4 mm. It is obvious from the Figure 40. to Figure 48. that the relative dose is underestimated in the region of 100-50% of the relative dose and overestimated for the rest of the profile. This is because the volume averaging is proportional to the second derivative of a dose profile. Where a second derivative of dose profile is lower than zero, the detector will underestimate absorbed dose. Similarly, if the value of a second derivative is large than zero, the detector will overestimate the absorbed dose. In inflection points, there is no volume averaging.⁵¹

The plotted dose profiles determined with different detectors are shown from Figure 40. to Figure 48. Results of an analysis of determined dose profile with different detectors are shown from Table 23. to Table 26., and from Figure 49. to Figure 51.

Detector	Profile	LP /	RP /	FWHM /	LP diff. from	RP diff. from	FWHM diff. from	•	Mean dose gradient RP
		mm	mm	mm	MC / mm	MC / mm	MC / mm	/ % per mm	/ % per mm
	16 x	9.02	9.03	21.75				6.65	6.65
	16 y	9.02	9.02	21.75				6.65	6.65
<u>0</u>	16 z	2.30	2.83	17.48				26.13	21.24
Monte Carlo	8 x	3.95	3.95	11.06				15.20	15.20
te	8 y	3.95	3.95	11.06				15.20	15.19
lon	8 z	2.36	2.21	9.80				25.47	27.11
≥	4 x	2.82	2.82	6.16				21.25	21.25
	4 y	2.82	2.82	6.16				21.25	21.25
	4 z	1.49	1.54	5.04				40.30	39.06
	16 x	9.05	9.00	21.29	0.02	-0.03	-0.46	6.63	6.67
_	16 y	9.45	9.39	21.49	0.43	0.37	-0.26	6.35	6.39
014	16 z	3.87	4.16	17.53	1.57	1.34	0.05	15.52	14.41
PinPoint T31014	8 x	4.67	4.63	10.62	0.72	0.69	-0.43	12.86	12.95
	8 y	4.69	4.65	10.67	0.74	0.70	-0.39	12.79	12.90
	8 z	3.76	3.77	9.77	1.41	1.56	-0.03	15.95	15.90
lui	4 x	3,74	3.72	6.23	0.91	0.90	0.070	16.06	16.13
-	4 y	3.76	3.72	6.29	0.93	0.90	0.13	15.97	16.12
	4 z	2.89	2.83	5.98	1.40	1.29	0.95	20.78	21.20
	16 x	9.27	9.30	21.66	0.25	0.28	-0.10	6.47	6.450
16	16 y	9.25	9.23	21.53	0.22	0.21	-0.22	6.49	6.50
PinPoint 3D T31016	16 z	3.25	3.61	17.37	0.95	0.78	-0.11	18.48	16.64
	8 x	4.72	4.68	10.92	0.77	0.73	-0.13	12.71	12.84
30	8 y	4.72	4.70	10.92	0.78	0.76	-0.13	12.71	12.76
aint	8 z	3.27	3.26	9.54	0.92	1.04	-0.26	18.34	18.42
nPc	4 x	3.72	3.67	6.49	0.90	0.85	0.33	16.14	16.34
Pil	4 y	3.68	3.71	6.48	0.86	0.89	0.32	16.29	16.16
	4 z	2.54	2.49	5.24	1.05	0.95	0.20	23.67	24.14

Table 23. Results of Gamma Knife's Perfexion dose profile measurements obtained with Monte Carlo simulations, PinPoint T31014 and PinPoint 3D T31016 chambers for FWHM, LP (left penumbra) and RP (right Penumbra) values and their difference from Monte Carlo values as well as mean dose gradients in the penumbra region. Values are rounded to one hundredth of a millimetre.

Detector	Profile	LP / mm	RP / mm	FWHM / mm	LP diff. from MC / mm	RP diff. from MC / mm	FWHM diff. from MC / mm	Mean dose gradient LP / % per mm	Mean dose gradient RP / % per mm
	16 x	8.72	8.55	21.62	-0.30	-0.48	-0.13	6.88	7.02
	16 y	9.08	9.04	21.81	0.05	0.01	0.06	6.61	6.64
E T60016	16 z	2.16	2.72	17.44	-0.14	-0.11	-0.03	27.78	22.07
600	8 x	3.89	3.83	11.05	-0.06	-0.12	-0.01	15.44	15.67
н	8 y	3.88	3.89	11.06	-0.07	-0.06	0.01	15.46	15.44
de	8 z	2.17	2.09	9.80	-0.19	-0.12	0.01	27.65	28.72
Diode	4 x	2.72	2.66	6.20	-0.10	-0.17	0.04	22.06	22.59
	4 y	2.70	2.72	6.20	-0.13	-0.11	0.03	22.25	22.10
	4 z	1.28	1.36	5.01	-0.21	-0.18	-0.03	46.86	44.28
	16 x	8.72	8.55	21.62	-0.30	-0.48	-0.13	6.88	7.02
	16 y	9.08	9.04	21.81	0.05	0.01	0.06	6.61	6.64
Р Т60017	16 z	2.16	2.72	17.44	-0.14	-0.11	-0.03	27.78	22.07
	8 x	3.89	3.83	11.05	-0.06	-0.12	-0.01	15.44	15.67
	8 y	3.88	3.89	11.06	-0.07	-0.06	0.01	15.46	15.44
Diode	8 z	2.17	2.09	9.80	-0.19	-0.12	0.01	27.65	28.72
Dio	4 x	2.72	2.66	6.20	-0.10	-0.17	0.04	22.06	22.59
	4 y	2.70	2.72	6.20	-0.13	-0.11	0.03	22.25	22.10
	4 z	1.28	1.36	5.01	-0.21	-0.18	-0.03	46.86	44.28
	16 x	8.81	8.83	21.63	-0.21	-0.19	-0.12	6.81	6.79
	16 y	8.96	8.82	21.67	-0.06	-0.20	-0.08	6.70	6.80
:=	16 z	2.29	2.82	17.44	-0.00	-0.00	-0.04	26.18	21.24
-bS	8 x	3.98	3.98	11.06	0.03	0.04	0.00	15.07	15.06
30	8 y	4.00	3.96	11.03	0.05	0.01	-0.03	14.99	15.15
EFD 3G-pSi	8 z	2.31	2.25	9.76	-0.05	0.03	-0.03	26.01	26.71
ш	4 x	2.86	2.84	6.24	0.04	0.01	0.08	20.96	21.15
	4 y	2.85	2.86	6.23	0.02	0.04	0.07	21.09	20.98
	4 z	1.39	1.49	4.99	-0.10	-0.05	-0.05	43.18	40.35

Table 24. Results of Gamma Knife's Perfexion dose profile measurements obtained with Diode E T60016, Diode P T60017 and EFD 3G-pSi detectors for FWHM, LP (left penumbra) and RP (right Penumbra) values and their difference from Monte Carlo value as well as mean dose gradients in the penumbra regions. Values are rounded to one hundredth of a millimetre.

Detector Profile III / IIIIII/III / IIIIIII/IIIIII/IIIIII	radient
Image: Note of the system 16 y 9.56 9.56 21.72 0.55 0.53 -0.03 6.28 6.28 16 z 3.64 3.95 17.31 1.34 1.12 -0.17 16.49 15.20 8 x 5.05 5.04 10.88 1.10 1.09 -0.18 11.89 11.90 8 y 5.06 5.07 10.86 1.12 1.12 -0.19 11.85 11.84 8 z 3.63 3.68 9.44 1.27 1.46 -0.35 16.53 16.32 4 x 4.09 4.06 7.03 1.27 1.24 0.87 14.67 14.77 4 y 4.07 4.09 7.02 1.25 1.27 0.86 14.72 14.66 4 z 2.93 2.71 5.65 1.45 1.17 0.61 20.45 22.18 16 x 8.70 8.78 21.42 -0.32 -0.24 -0.33 6.90 6.83	mm
by 16 z 3.64 3.95 17.31 1.34 1.12 -0.17 16.49 15.20 8 x 5.05 5.04 10.88 1.10 1.09 -0.18 11.89 11.90 8 y 5.06 5.07 10.86 1.12 1.12 -0.19 11.85 11.84 8 z 3.63 3.68 9.44 1.27 1.46 -0.35 16.53 16.32 4 x 4.09 4.06 7.03 1.27 1.24 0.87 14.67 14.77 4 y 4.07 4.09 7.02 1.25 1.27 0.86 14.72 14.66 4 z 2.93 2.71 5.65 1.45 1.17 0.61 20.45 22.18 16 x 8.70 8.78 21.42 -0.32 -0.24 -0.33 6.90 6.83	
8x 5.05 5.04 10.88 1.10 1.09 -0.18 11.89 11.90 8y 5.06 5.07 10.86 1.12 1.12 -0.19 11.85 11.84 8z 3.63 3.68 9.44 1.27 1.46 -0.35 16.53 16.32 4x 4.09 4.06 7.03 1.27 1.24 0.87 14.67 14.77 4y 4.07 4.09 7.02 1.25 1.27 0.86 14.72 14.66 4z 2.93 2.71 5.65 1.45 1.17 0.61 20.45 22.18 16x 8.70 8.78 21.42 -0.32 -0.24 -0.33 6.90 6.83 <td></td>	
4 x 4.09 4.06 7.03 1.27 1.24 0.87 14.67 14.77 4 y 4.07 4.09 7.02 1.25 1.27 0.86 14.72 14.66 4 z 2.93 2.71 5.65 1.45 1.17 0.61 20.45 22.18 16 x 8.70 8.78 21.42 -0.32 -0.24 -0.33 6.90 6.83	
4 x 4.09 4.06 7.03 1.27 1.24 0.87 14.67 14.77 4 y 4.07 4.09 7.02 1.25 1.27 0.86 14.72 14.66 4 z 2.93 2.71 5.65 1.45 1.17 0.61 20.45 22.18 16 x 8.70 8.78 21.42 -0.32 -0.24 -0.33 6.90 6.83	
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4 y 4.07 4.09 7.02 1.25 1.27 0.86 14.72 14.66 4 z 2.93 2.71 5.65 1.45 1.17 0.61 20.45 22.18 16 x 8.70 8.78 21.42 -0.32 -0.24 -0.33 6.90 6.83	
4 z 2.93 2.71 5.65 1.45 1.17 0.61 20.45 22.18 16 x 8.70 8.78 21.42 -0.32 -0.24 -0.33 6.90 6.83	
16 x 8.70 8.78 21.42 -0.32 -0.24 -0.33 6.90 6.83	
16 y 9.23 9.19 21.68 0.20 0.17 -0.07 6.50 6.53	
E 16 z 3.04 2.66 17.38 0.75 -0.16 -0.10 19.73 22.55	
8 x 4.23 4.29 10.94 0.28 0.34 -0.12 14.19 13.99	
8 y 4.28 4.43 10.95 0.33 0.48 -0.10 14.01 13.56	
8 z 2.58 2.79 9.72 0.23 0.58 -0.08 23.23 21.52	
4 x 3.27 3.24 6.13 0.45 0.42 -0.03 18.34 18.50	
4 y 3.16 3.18 6.16 0.33 0.36 -0.01 19.01 18.86	
4 z 1.97 1.80 4.93 0.48 0.26 -0.11 30.44 33.37	
16 x 9.13 9.20 21.50 0.10 0.17 -0.25 6.57 6.52	
16 y 9.24 9.24 21.53 0.22 0.22 -0.22 6.49 6.49	
<u>a</u> 16 z 3.30 3.69 17.51 1.01 0.87 0.04 18.16 16.25	
8 x 4.44 4.44 10.83 0.49 0.49 -0.23 13.53 13.51	
8 y 4.45 4.47 10.81 0.50 0.52 -0.25 13.50 13.43	
If z 3.30 3.69 17.51 1.01 0.87 0.04 18.16 16.25 8 x 4.44 4.44 10.83 0.49 0.49 -0.23 13.53 13.51 8 y 4.45 4.47 10.81 0.50 0.52 -0.25 13.50 13.43 8 z 3.26 3.28 9.72 0.90 1.06 -0.08 18.42 18.30 4 x 3.44 3.45 6.08 0.62 0.63 -0.08 17.44 17.38	
4 x 3.44 3.45 6.08 0.62 0.63 -0.08 17.44 17.38	
4 y 3.43 3.47 6.09 0.61 0.65 -0.08 17.47 17.29	
4 z 2.50 2.57 5.41 1.01 1.04 0.37 24.00 23.32	

Table 25. Results of Gamma Knife's Perfexion dose profile measurements obtained with CC04, RAZORnano and RAZOR chambers for FWHM, LP (left penumbra) and RP (right Penumbra) values and their difference from Monte Carlo values as well as mean dose gradients in the penumbra regions. Values are rounded to one hundredth of a millimetre.

Detector	Profile	LP / mm	RP / mm	FWHM / mm	LP diff. from MC / mm	RP diff. from MC / mm	FWHM diff. from MC / mm	Mean dose gradient LP / % per mm	Mean dose gradient RP / % per mm
	16 x	8.95	8.90	21.70	-0.08	-0.13	-0.05	6.70	6.74
	16 y	8.93	8.92	21.68	-0.09	-0.10	-0.07	6.72	6.73
e	16 z	2.34	2.85	17.46	0.04	0.02	-0.01	25.67	21.08
lioe	8 x	3.97	3.95	11.00	0.02	0.00	-0.06	15.12	15.19
BR C	8 y	3.99	3.99	10.98	0.04	0.04	-0.08	15.04	15.05
RAZOR diode	8 z	2.32	2.27	9.80	-0.04	0.06	0.00	25.86	26.42
R/	4 x	2.78	2.76	6.19	-0.04	-0.07	0.03	21.56	21.77
	4 y	2.77	2.79	6.19	-0.06	-0.03	0.02	21.70	21.51
	4 z	1.40	1.48	5.08	-0.09	-0.06	0.04	42.87	40.64
•	16 x	8.78	8.73	21.64	-0.25	-0.30	-0.11	6.84	6.87
019	16 y	9.17	9.16	21.85	0.15	0.14	0.10	6.54	6.55
microDlamond T60019	16 z	2.23	2.72	17.38	-0.07	-0.10	-0.10	26.92	22.02
	8 x	4.09	4.02	11.11	0.14	0.07	0.05	14.69	14.93
	8 y	4.03	4.07	11.09	0.08	0.12	0.04	14.88	14.74
	8 z	2.28	2.18	9.70	-0.07	-0.03	-0.10	26.27	27.52
2	4 x	2.93	2.88	6.25	0.11	0.05	0.09	20.45	20.87
nic	4 y	2.90	2.96	6.27	0.07	0.12	0.11	20.70	20.36
<u> </u>	4 z	1.39	1.46	4.92	-0.10	-0.07	-0.12	43.02	40.92
	16 x	8.39	8.52	21.24	-0.63	-0.51	-0.51	7.15	7.05
	16 y	8.75	8.70	21.73	-0.28	-0.33	-0.02	6.86	6.90
	16 z	2.35	2.90	17.52	0.05	0.08	0.05	25.54	20.68
<u>n</u>	8 x	3.94	3.96	10.86	-0.01	0.01	-0.20	15.25	15.16
EBT3 film	8 y	4.12	4.02	11.02	0.17	0.07	-0.02	14.58	14.93
EB	8 z	2.53	2.42	9.07	0.17	0.21	-0.73	23.72	24.79
	4 x	2.86	2.79	5.95	0.04	-0.04	-0.21	20.95	21.55
	4 y	2.77	2.79	6.18	-0.06	-0.02	0.02	21.70	21.48
	4 z	1.62	1.61	5.04	0.13	0.08	0.00	36.96	37.20

Table 26. Results of Gamma Knife's Perfexion dose profile measurements obtained with RAZOR diode, microDiamond and EBT3 film for FWHM, LP (left penumbra) and RP (right Penumbra) values and their difference from Monte Carlo values as well as mean dose gradients in the penumbra regions. Values are rounded to one hundredth of a millimetre.

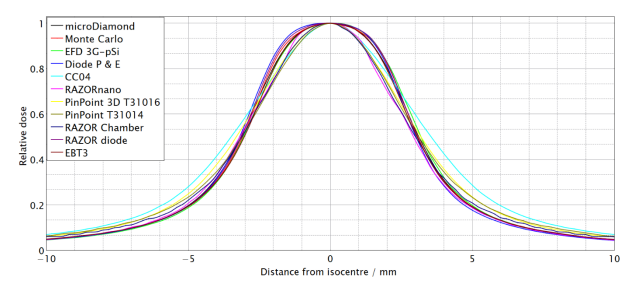


Figure 40. 4 mm x - axis dose profile determined with Monte Carlo, microDiamond, EFD 3GpSi, Diode P & E, CC04 chamber, RAZORnano, PinPoint, PinPoint 3D, RAZOR chamber, RAZOR diode and EBT3 film.

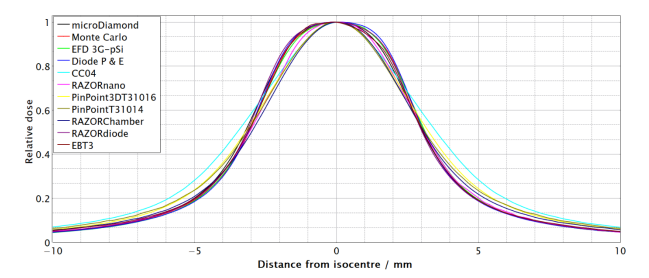


Figure 41. 4 mm y - axis dose profile determined with Monte Carlo, microDiamond, EFD 3GpSi, Diode P & E, CC04 chamber, RAZORnano, PinPoint, PinPoint 3D, RAZOR chamber, RAZOR diode and EBT3 film.

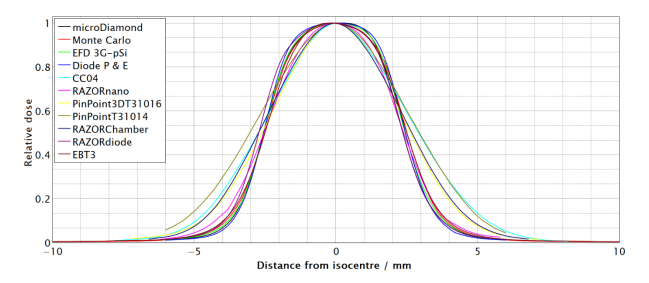


Figure 42. 4 mm z - axis dose profile determined with Monte Carlo, microDiamond, EFD 3GpSi, Diode P & E, CC04 chamber, RAZORnano, PinPoint, PinPoint 3D, RAZOR chamber, RAZOR diode and EBT3 film.

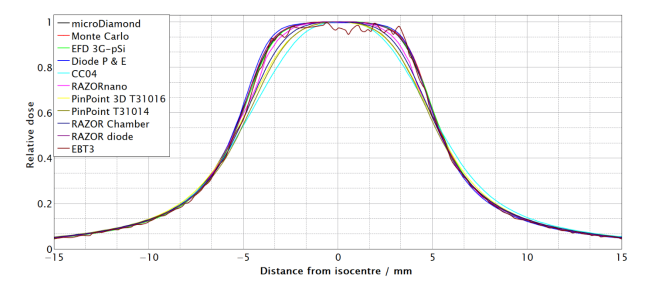


Figure 43. 8 mm x - axis dose profile determined with Monte Carlo, microDiamond, EFD 3GpSi, Diode P & E, CC04 chamber, RAZORnano, PinPoint, PinPoint 3D, RAZOR chamber, RAZOR diode and EBT3 film.

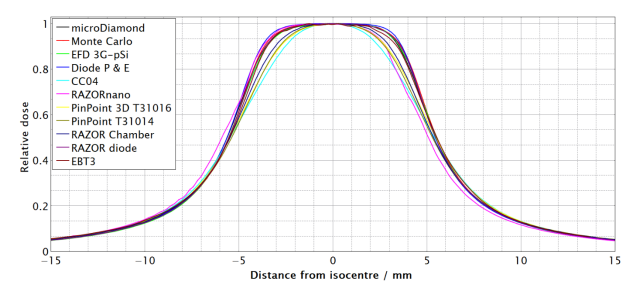


Figure 44. 8 mm y - axis dose profile determined with Monte Carlo, microDiamond, EFD 3GpSi, Diode P & E, CC04 chamber, RAZORnano, PinPoint, PinPoint 3D, RAZOR chamber, RAZOR diode and EBT3 film.

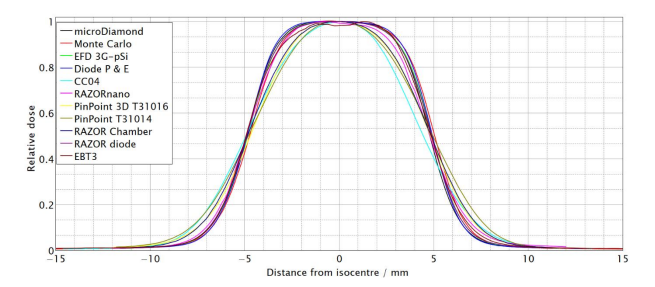


Figure 45. 8 mm z - axis dose profile determined with Monte Carlo, microDiamond, EFD 3GpSi, Diode P & E, CC04 chamber, RAZORnano, PinPoint, PinPoint 3D, RAZOR chamber, RAZOR diode and EBT3 film.

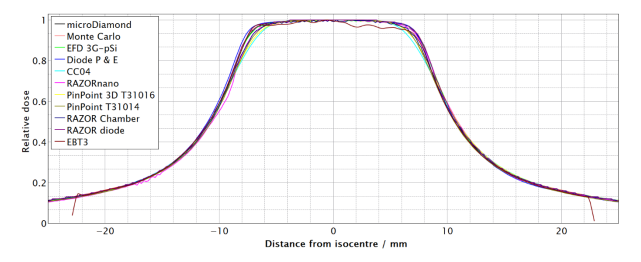


Figure 46. 16 mm x - axis dose profile determined with Monte Carlo, microDiamond, EFD 3G-Si, Diode P & E, CC04 chamber, RAZORnano, PinPoint, PinPoint 3D, RAZOR chamber, RAZOR diode and EBT3 film.

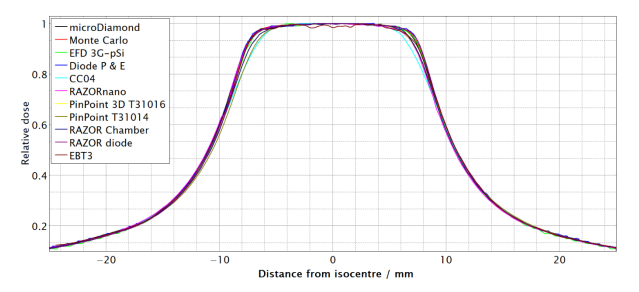


Figure 47. 16 mm y - axis dose profile determined with Monte Carlo, microDiamond, EFD 3GpSi, Diode P & E, CC04 chamber, RAZORnano, PinPoint, PinPoint 3D, RAZOR chamber, RAZOR diode and EBT3 film.

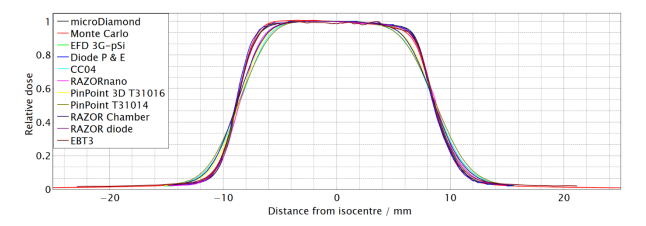


Figure 48. 16 mm z - axis dose profile determined with Monte Carlo, microDiamond, EFD 3GpSi, Diode P & E, CC04 chamber, RAZORnano, PinPoint, PinPoint 3D, RAZOR chamber, RAZOR diode and EBT3 film.

In Figure 49. and Figure 50. the difference from the Monte Carlo values in the width of the left and right penumbra for all dose profiles is presented. All ionization chambers have a larger difference from Monte Carlo values than semiconductor or diamond detectors due to the more dominant volume-averaging effect (Table 16.). RAZORnano chamber, ionization chamber with the smallest volume (V=3 mm³), is in between these two groups of detectors, therefore, one can expect its volume-averaging correction factor to be the smallest of all ionization chambers. A spike in the difference of ionization chambers at the z-axis dose profile for all field sizes is noticeable while the difference for the x and y -axis is not as pronounced. This can be explained by the difference in the length of these profiles, increasing the volumeaveraging i.e. z-axis dose profile is shorter than the x and y-axis. One should keep in mind that detectors were placed parallel with the z-axis of the device because of the phantom default geometry making them more influenced by the z-axis dose gradient. This effect is noticeable for RAZORnano chamber as well, being spherical in shape. Although the PinPoint T31014 chamber is not the largest (V=15 mm³), its difference in the width of the penumbras from the Monte Carlo values is the largest for the z-axis dose profile. This can be explained by the volume-averaging correction factors previously calculated (Table 16.). Because of the shape of the PinPoint T31014 chamber and its position with the long axis parallel to the z-axis dose profile while scanning the profile, it is more influenced by the gradient of the z-axis dose profile, increasing volume-averaging and flattening the dose profile. Both CC04 and PinPoint 3D T31016 chambers have larger volumes than PinPoint, however, they are more compact reducing therefore their volume-averaging (the length-to-diameter ratio is closer to one). The differences from the Monte Carlo results on the x and y – axis dose profile are largest for the CC04 because of its volume (V=40 mm³), followed by the PinPoint 3D (V=16 mm³), PinPoint (V=15mm³), RAZOR chamber (V=10 mm³) and RAZORnano chamber (V=3 mm³) (Figure 49. and Figure 50.). This supports previous findings that the z-axis dose profile plays a fundamental role in volume-averaging on Gamma Knife and that besides volume, the shape of the detector has an important role as well as explained in section 5.4. From this study, we conclude that the semiconductor and diamond detector, alongside EBT3 film are a better option for dose profile determination than ionization chambers.

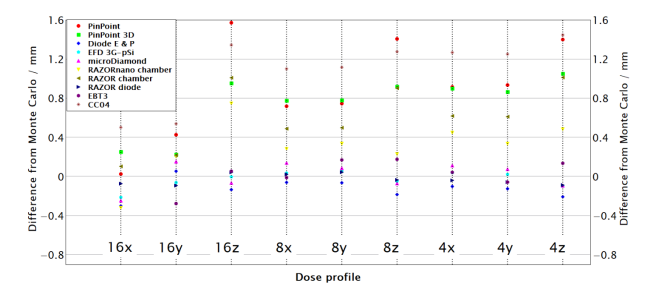


Figure 49. The difference from Monte Carlo results in the widths of left penumbra (LP) measured with different detectors.

The semiconductor detector, microDiamond and EBT3 films have much smaller differences in the width of left and right penumbras from Monte Carlo values compared with the ionization chambers (Figure 49. and Figure 50.). This is, again, due to its volume which is several orders of magnitude smaller than the ionization chamber's one, therefore, minimizing the volumeaveraging effect. The spike at the z-axis dose profile is smaller since the spatial dimensions of the detector in that direction are small limiting the influence of the z-axis dose gradient. Due to the small thickness of these detectors many of them show a decrease in the difference from Monte Carlo values when scanning z -the axis dose profile and an increase in the x and y direction. However, these differences are not as pronounced as they are for ionization chambers and when performing dose profile measurements, they should be the detectors of choice.

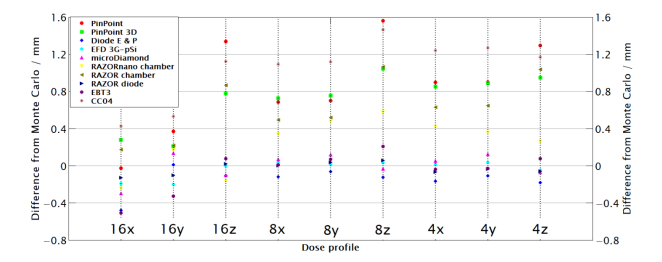


Figure 50. Difference from Monte Carlo in the widths of right penumbra (RP) measured with different detectors.

The results of an analysis for FWHM value can be seen in Figure 51. The difference from Monte Carlo values is smaller than it is for the width of penumbra regions. This is expected since volume-averaging should not impact the FHWM values. It can be seen from Figure 40. to Figure 48., that the measured dose profile passes through or nearby the point of 50% isodose value resulting in a minimal difference in FWHM value. The exceptions are CC04 and PinPoint chambers which show a substantial increase at 4 mm field size.

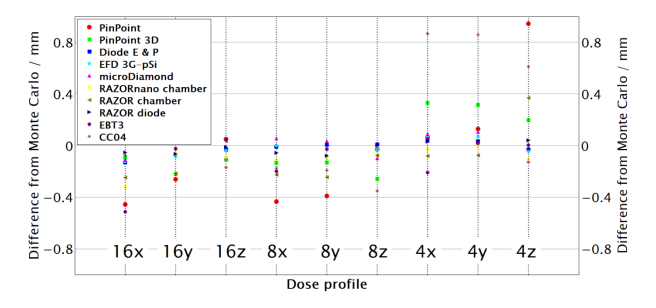


Figure 51. The difference from Monte Carlo results in the FWHM value measured with different detectors.

6. Conclusion

A first hypothesis of this dissertation is directed towards the problem of finding the suitable detector for dosimetry of Gamma Knife's narrow photon beams. Therefore, field output factors $\Omega_{Q_0,Q_0}^{f_{clin},f_{msr}}$ were determined with: seven ionization chambers, four semiconductor detectors, and diamond detectors. Detector's data were compared against Monte Carlo and EBT3 film values. Additionally, a detectors-specific correction factors $k_{Q_0,Q_0}^{f_{clin},f_{msr}}$ were determined for investigated detectors, all widely used for relative dosimetry.

EBT3 film overestimated the field output factor for the smallest 4 mm field by 1.4% and underestimated the 8 mm field by 1.2%. As shown in this research EBT3 film does have disadvantages such as the low signal-to-noise ratio, demanding handling and postirradiation processing. Most importantly, the uncertainty of the absorbed dose determined with the EBT3 film detector can be substantial. Ionization chambers have consistently underestimated field output factors due to perturbation which are more emphasized in smaller fields. Large ionization chambers such as Semiflex T31010, Semiflex 3D T31021, PinPoint T31014, PinPoint 3D T31016, RAZOR chamber and CC04, are not suitable for dosimetry in the smallest 4 mm field. The exception is RAZORnano chamber. Semiconductor and diamond detector have performed better than ionization chambers in the 4 mm field size and have substantially lower detector-specific correction factors. This is primarily due to smaller volume-averaging effect. It has been shown that not a single investigated detector can fulfil the CPE condition in the 4 mm field (Table 15.).

All ionization chambers have shown better agreement of their field output factors with the reference values in the 8 mm field. This is due to the preservation of lateral charged particle equilibrium (Table 15.) and decrease of volume-averaging effect (Table 16.). Large ionization chambers (e.g., Semiflex and Semiflex 3D) have significant perturbations, thus, they are not recommended for the dosimetry of 8 mm field. CCO4 has underestimated the doze by 3.2%. Remaining ionization chambers have their detector-specific correction factors within 1% from reference values. Semiconductor detectors have shown better agreement with field output factors determined with the Monte Carlo calculations and EBT3 film than ionization chambers in the 4 mm field due to their small volume and high sensitivity. On the other hand, all of them

over-responded because of the high atomic numbers of silicon. The microDiamond detector has proven to be the best active detector with values of field output factors within 1.6% for all field sizes. Uncertainty analysis has shown that EBT3 film, although tissue-equivalent, has large uncertainty (Table 11.). Active detectors are more stable with lower uncertainties, but they require correction factors (Table 12.).

Given the fact that there is a small number of detectors with the detectors-specific correction factors for narrow Co-60 beams reported in the IAEA TRS-483 CoP, this research can be considered as a valuable supplement to the results of previous investigations^{12,15–18,20,24,31,39,90,91} already published and complied as IAEA TRS-483 CoP dataset. ¹⁰

Our second hypothesis is aimed towards the problem of accurate determination of volumeaveraging correction factor in the narrow Co-60 beams of Gamma Knife. It is assumed that volume-averaging correction factors $(k_{vol})_{Q_0,Q_0}^{fmsr,f_{clin}}$ can be calculated by simulating the ellipsoid absorbed dose model and the detector geometries inside it. Applying numerical integration of the absorbed dose distribution over the detector's volume, a volume-averaging correction factor is calculated in all fields for: seven ionization chambers, four semiconductor detectors and a diamond detector.⁵⁴ To the best of our knowledge, no volume averaging correction factors for clinically used detectors in different Gamma Knife absorbed dose distributions have been previously reported in the literature.

It is confirmed that the volume-averaging correction factor is inversely proportional to the field size, and in general, it increases with the detector's size. It can be noticed that some detectors have more prominent correction, although smaller in volume. The reason for this is the z-axis dose profile which is asymmetrical and shorter compared with the x and y-axis. Placing the detectors with the long axis in the z-direction makes them more influenced by the asymmetry of the shortest z-axis dose profile. Consequently, the larger change of a dose gradient over their effective volume will result in increased volume-averaging. Because of this, an ionization chamber such as PinPoint T31014 has higher correction than CC04 for the smallest 4 mm field although is almost three times smaller in volume. It can be concluded that although the size of an effective volume of the detector is important the shape of the detectors plays an important role as well. These findings are supported by previous studies

involving dosimetry of high-energy X-rays with detectors positioned in parallel and perpendicular orientations.⁵⁵

Additionally, it has been shown that volume-averaging is the dominant perturbation in the narrow Co-60 beams used by Gamma Knife for the detectors that have their detector-specific correction factor reported in the IAEA TRS-483 CoP (Table 17.). Presented volume-averaging data can be considered as valuable supplement to the literature.

Finally, third hypothesis aimed at the problem of dose profile determination on Gamma Knife and finding the suitable detectors for such measurements. In this research, Gamma Knife's dose profiles are determined with EBT3 film and ten active detectors; five ionization chambers, four semiconductor and diamond detector. Obtained results are compared with Monte Carlo calculated values. Some detectors were excluded from this study due to their large volume while others were unavailable (Table 16). The difference in the widths of penumbras and FWHM values for different detectors was examined. Since volume-averaging is the only effect influencing the dose profile, ionization chambers with large volumes are not suitable for these types of measurements. They significantly underestimate the absorbed dose in the region of 100%-50% of relative value, and overestimate for the remaining dose profile. Results of smaller ionization chamber, e.g. RAZORnano, agree better with Monte Carlo calculated profiles. Moreover, it is noticed that large ionization chambers have a spike in difference for the z-axis dose profile. Reasons for this can be found in the shorter length (FWHM) of the z-axis dose profile and detector's position. Semiconductor detectors, and diamond detectors due to their small size have results in better agreement with the Monte Carlo calculated profiles. They, together with the EBT3 film should be the detectors of choice for dose profile determination.

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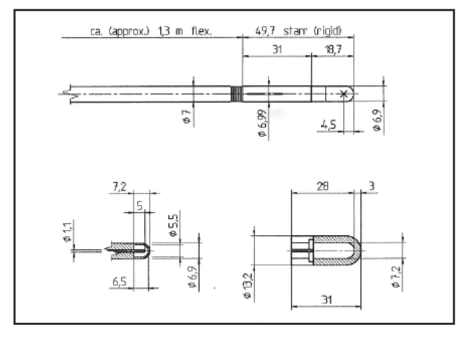
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Appendix A

Detector schematics presented in this Appendix are relevant to the calculation of volumeaveraging and are presented only for fallowing detectors:

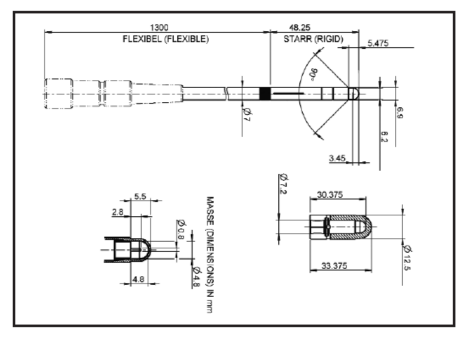
- PTW Semiflex T31010
- PTW Semiflex 3D T31021
- PTW PinPoint T31014
- PTW PinPoint 3D T31016
- RAZOR chamber
- RAZORnano chamber
- CC04 chamber

These schematics are kindly provided by the PTW Freiburg, Germany as well as IBA Dosimetry Schwarzenbruck. The remaining detectors have simpler geometers being disk (cylinder) or rectangular shaped. For these detectors data provided in Table 10. are sufficient.

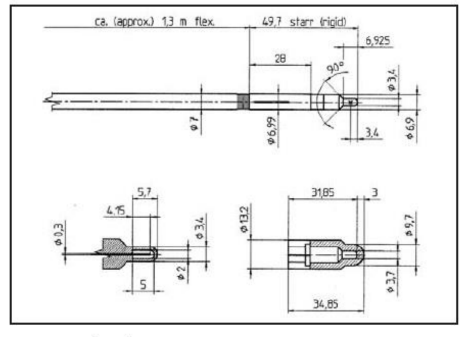


0.125 cm³ Semiflex Chamber



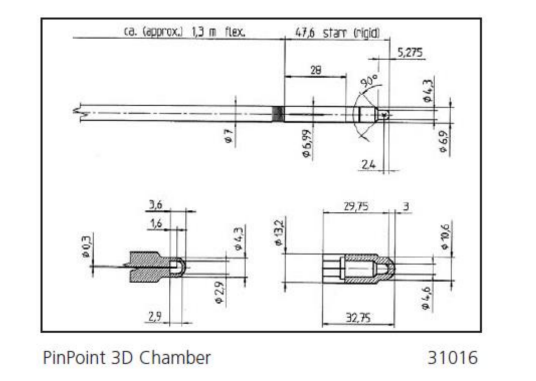


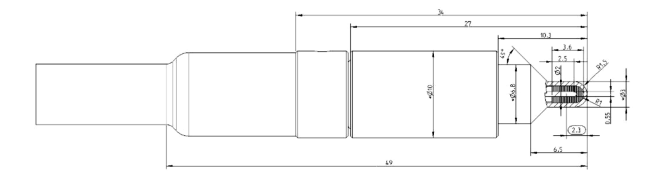
0.07 cm³ Semiflex 3D Chamber



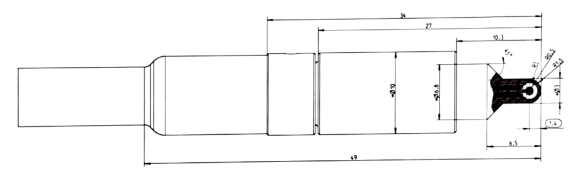
PinPoint Chamber

31014

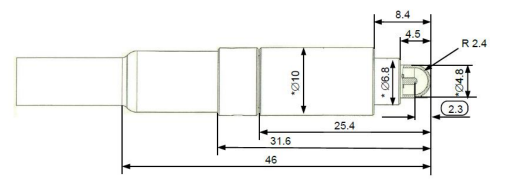




Dimensions of the RAZOR^{Chamber} (dimensions in mm, add 0.2 mm for tolerance if you intend to make adapters)



Dimensions of the RAZOR Nano ^{Chamber} (dimensions in mm, add 0.2 mm tolerance if you intend to make adapters)



Dimensions of CC04 (dimensions in mm, add 0.2 mm for tolerance if you intend to make adapters)

Appendix B

A computer code written in MATLAB for the calculation of volume-averaging on Gamma Knife Perfexion for detectors studied in this thesis is provided in this Appendix. The written program calculates volume-averaging correction factor for the following detectors:

- PTW Semiflex T31010
- PTW Semiflex 3D T31021
- PTW PinPoint T31014
- PTW PinPoint 3D T31022
- PTW Diode P T60016
- PTW Diode E T60017
- PTW microDiamond T60019
- IBA RAZOR diode
- IBA RAZOR chamber
- IBA CC04
- EDGE detector
- EFD diode

- 1 %Calculation of a volume averaging on Gamma Knife
- 2 digits(32)
- 3 format short
- 4 polje=menu('Please select the field size for which volume averaging is calculated',' 16 mm','8
- 5 mm','4 mm'); %select the field size
- 6 if polje == 1 %16 mm field
- 7 %Coefficients for s direction in a cylindrical coordinate system x direction is fitted from
- 8 Leksell coordinate system
- 9 cOs = 9.69539388004509E-01;
- 10 A1s = 3.81762920289576E-01;
- 11 a1s = 2.49913438196976E+01;
- 12 b1s = -1.67550520977859E+00;
- 13 A2s = -2.03591059391798E-01;
- 14 a2s = -8.94687901300991E+00;
- 15 b2s = 1.926266666629579E+00;
- 16 A3s = -1.02427191560397E-02;
- 17 a3s = 5.32940693032982E+00;
- 18 b3s = 4.91979665382671E+01;
- 19 A4s = 2.09537779074513E-01;
- 20 a4s = -1.24942845009386E+01;
- 21 b4s = -4.58390178363538E+00;
- 22

23 %Coefficients for z direction in a cylindrical coordinate system - z direction is fitted from

- 24 Leksell coordinate system
- 25 cOz = 6.23439585197441E-O2;
- 26 A1z = 1.65016076148551E+00;
- 27 a1z = -6.64266771664852E+00;
- 28 b1z = -2.54636160615763E+00;
- 29 A2z = 1.75606701707450E+00;
- 30 a2z = -5.60760279869319E+00;
- 31 b2z = -1.76766384666806E+00;
- 32 A3z = 2.97263968316475E+00;
- 33 a3z = -5.79821291403295E+00;
- 34 b3z = 1.93643475942725E+00;
- 35 A4z = -3.39734861765111E-03;
- 36 a4z = 1.89356823656770E-01;
- 37 b4z = 1.17416975626696E+00;
- 38 A5z = 4.67330339984927E-01;
- 39 a5z = 8.76144173778636E+00;
- 40 b5z = 1.80287660180637E+00;
- 41 A0z = 3.12810092663561E-02;
- 42 az = -4.71330838427608E+00;
- 43 bz = -7.59998845310368E-01;
- 44 else if polje == 2 %8 mm polje
- 45 %Coefficients for s direction in a cylindrical coordinate system x direction is fitted from
- 46 Leksell coordinate system
- 47 cOs = 5.03635710695764E-01;

48	A1s = 9.88325720443460E-02;
49	a1s = -8.01982219971946E+00;
	b1s = -4.03898164613522E+00;
51	A2s = 5.27104129024405E-02;
52	a2s = -1.36396115774001E+01;
53	$b_{2s} = -9.40368469738407E+00;$
55 54	A3s = 2.35344836867871E-01;
55	a3s = -5.37881836259153E+00;
56	$b_{3s} = -2.18398489903176E+00;$
57	A4s = 1.12806315029468E-01;
58	a4s = -4.58912556690278E+00;
59	b4s = -1.06379030441735E+00;
60	543 - 1.003750504417552400,
61	%Coefficients for z direction in a cylindrical coordinate system - z direction is fitted from
62	Leksell coordinate system
63	c0z = 1.15950725257977E+00;
64	A1z = -5.92978414412138E+00;
65	a1z = -5.36550589710118E+00;
66	b1z = -1.08424553524007E+02;
67	A2z = -4.67178392213500E-01;
68	a2z = 4.67455624404566E+00;
69	b2z = -1.84823205607355E+00;
70	A3z = 4.83205221309876E-01;
71	a3z = -5.04296136055748E+00;
72	b3z = -1.84831005046076E+00;
73	A4z = -4.78929405327008E+00;
74	a4z = -4.30280110051637E+00;
75	b4z = 8.53827968025802E+01;
76	A5z = -3.69804960557376E-02;
77	a5z = 7.56801950098436E+00;
78	b5z = -3.60316445872631E+00;
79	A0z = 1.34532865234678E+00;
80	az = -2.51279085600255E+00;
81	bz = 2.69871447884177E+00;
82	else if polje == 3 %4 mm polje
83	%Coefficients for s direction in a cylindrical coordinate system - x direction is fitted from
84	Leksell coordinate system
85	c0s = 5.33262345742450E-01;
86	A1s = 2.66410381382869E-01;
87	a1s = -2.59032772830263E+00;
88	b1s = -1.19772081535829E+00;
89	A2s = 8.47103711400306E-02;
90	a2s = -5.30783226357609E+00;
91	b2s = -7.40969527700572E+00;
92	A3s = 1.49732918683219E-01;
93	a3s = -4.03070534843962E+00;
94	b3s = -2.33322160403931E+00;

95 A4s = -6.15704845438336E-02;96 a4s = 5.29242467969531E+00;b4s = 7.00442818850188E+01;97 98 %Coefficients for z direction in a cylindrical coordinate system - z direction is fitted from 99 100 Leksell coordinate system cOz = 3.81570813736090E-02; 101 A1z = 4.80640052626926E-01; 102 103 a1z = -2.48998145956572E+00;104 b1z = -1.24629020543953E+00;105 A2z = 4.79647659835712E-01; 106 a2z = 2.50218824713077E+00; 107 b2z = 1.20777444317923E+00; 108 end 109 end end 110 %List of variables: 111 rr1 - Vector in cylindrical coordinate system 112 % fss - Dose profile in s direction with "s" variable 113 % fs1 - Dose profile in s direction with "rr1" variable 114 % % fzz - Dose profile in z direction with "z" variable 115 fz1 - Dose profile in +z direction with "rr1&rr2" variable 116 % 117 % fz2 - Dose profile in -z direction with "rr1&rr2" variable raspodjela - distribution in an elliptical model 118 % s0 - shift in field in s direction 119 % z0 - shift in field in z direction % 120 z1(s) - initial point of integration of an cylindrical part of 121 % 122 % detector z2(s) - final point of integration of an cylindrical part of an 123 % detector 124 % 125 % $z_3(s)$ - length of an electrode above the xy plain determined by % the reference point 126 127 % s1 - initial point of integration of an detector/electrode 128 % s2 - detector radius 129 s3 - radius of an electrode % z00(s) - defines zero so we can integrate on both z profiles 130 % zf - function that describes the curvature of an chamber 131 % % n - number of steps of integration in s direction 132 % m - number of steps of integration in z direction 133 f1 - distribution for z>0 and differential for integration in 134 % % cylindrical coordinate system is added at the end 135 f2 - distribution for z<0 and differential for integration in % 136 137 % cylindrical coordinate system is added at the end 138 % Juk1 - value of an integral of a cylinder in the area z>0 to the 139 % curved part of chamber Juk2 - value of an integral of a cylinder in the area z<0 to the 140 % 141 % curved part of chamber

```
Juk3 - integral of a cylindrical electrode in the area z>0
142
      %
            Juk4 - integral of a cylindrical electrode in the area z<0
143
      %
      %
            Juk5 - integral of a curved part
144
145
      syms s
      z00=@(s) 0;
146
147
      n=100;
148
      m=100;
149
      k=menu('Select the detector to calculate volume averaging','Semiflex TM31010', 'PinPoint
150
      TM31014', 'Dioda P T60016', 'Dioda E T60017', 'PinPoint 3D T31022',...
151
152
         'microDiamond T60019', 'EDGE Detector', 'EFD diode', 'RAZOR diode detector', 'RAZOR
      chamber', '%RAZORnano chamber', 'Semiflex 3D', 'IBA CC04', 'Exradin W2 1X1', 'Excradin W3
153
154
      1X3');
      if k==1 %Semiflex T31010
155
         z1=@(s) -2.7; %length of a detector in z<0
156
         z2=@(s) 1.05; %length of a detector in z>0 (only cylindrical part)
157
         z3=@(s) 2.3; %length of an electrode above z>0
158
         s1=0.0001; %initial point of an integration of a detector
159
         s2=2.75; %detector's radius
160
         s3=0.55; %radius of an electrode
161
         zf=@(s) sqrt(7.5625-s.^2); %function that describes the radius of curvature of a top part
162
         V=128; %volume in mm3
163
      else if k==2 %PinPoint T31014
164
           z1=@(s) -2.3;
165
166
           z2=@(s) 2.4294;
           z3=@(s) 1.85;
167
           s1=0.0001;
168
169
           s2=1;
           s3=0.15;
170
           zf=@(s)sqrt(3.9326-s.^2)-1.71;
171
172
           V=15;
173
      else if k==3 %Diode P T60016
174
           z1=@(s) -0.015;
175
           z2=@(s) 0.015;
           z3=@(s) 0;
176
177
           s1=0.0001;
178
           s2=0.56418;
179
           s3=0;
           zf=@(s) 0;
180
           V=0.03;
181
         else if k==4 %Diode E T60017
182
           z1=@(s) -0.015;
183
184
           z2=@(s) 0.015;
185
           z3=@(s) 0;
           s1=0.0001;
186
           s2=0.56418;
187
188
           s3=0;
```

189	zf=@(s)0;
190	V=0.03;
191	else if k==5 %PinPoint 3D TM31022
192	z1=@(s) -0.175;
193	z2=@(s) 1.7676;
194	z3=@(s) 1.425;
195	zf=@(s) sqrt(2.4649-s.^2)-0.6192;
196	s1=0.0001;
197	s2=1.45;
198	s3=0.3;
199	V=16;
200	else if k==6 %microDiamond TM60019
200	z1=@(s) -5*10^(-7);
201	z2=@(s) 5*10^(-7);
202	z3=@(s) 0;
204	s1=0.0001;
205	s2=1.1;
206	s3=0;
207	zf=@(s) 0;
208	V=3.801*10^(-6);
209	else if k==7 %EDGE detector
210	xmin=-0.4;
211	xmax=0.4;
212	ymin=-0.015;
213	ymax=0.015;
214	zmin=-0.4;
215	zmax=0.4;
216	V=0.0192;
217	else if k==8 %EFD diode
218	z1=@(s) -0.03;
219	z2=@(s) 0.03;
220	z3=@(s) 0;
221	s1=0.0001;
222	s2=1;
223	s3=0;
224	zf=@(s) 0;
225	V=0.1885;
226	else if k==9 %RAZOR diode
227	z1=@(s) -0.01;
228	z2=@(s) 0.01; z2=@(s) 0.01;
229	z3=@(s) 0;
	s1=0.0001;
230	
231	s2=0.3;
232	s3=0;
233	zf=@(s) 0;
234	V=5.655*10^(-3);
235	else if k==10 %RAZOR chamber

236	z1=@(s) -1.8;
237	z2=@(s) 0.8;
238	z3=@(s) 0.7;
239	s1=0.0001;
240	s2=1;
241	s3=0.275;
242	zf=@(s) sqrt(1.001-s.^2);
243	V=9.668;
244	% else if k==11; %RAZORnano chamber
245	z1=@(s) 0;
246	z2=@(s) 0;
247	z3=@(s) 0;
248	s1=0.0001;
249	s2=1;
250	s3=0.5;%radius of an spherical electrode
251	zf=@(s) sqrt(1-s.^2);
252	zf2=@(s) sqrt(0.5.^2-s.^2);
253	V=3.665;
254	else if k==12 %Semiflex 3D T31021
255	z1=@(s) -2.05;
256	z2=@(s) 0.75;
257	z3=@(s)0.75;
258	s1=0.00001;
259	s2=2.4;
260	s3=0.4;
261	zf=@(s) sqrt(5.76-s.^2)-0.4;
262	V=71.04;
263	else if k==13 %IBA CC04
264	z1=@(s) -1.6;
265	z2=@(s) 0;
266	z3=@(s) 0.5;
267	s1=0.00001;
268	s2=2;
269	s3=0.5;
270	zf=@(s) sqrt(4-s.^2);
271	V=35.37;
272	else if k==14; %Exradin W2 1X1
273	z1=@(s) -0.5;
274	z2=@(s) 0.5;
275	z3=@(s) 0;
276	s1=0.0001;
277	s2=0.5;
278	s3=0;
279	zf=@(s)0;
280	V=0.7854;
281	else if k==15 %Exradin W2 1X3
282	z1=@(s) -1.5;

283	z2=@(s) 1.5;
284	z3=@(s) 0;
285	s1=0.0001;
286	s2=0.5; s3=0;
287 288	zf=@(s)0;
288 289	V=2.3562;
289	end
291	end
292	end
293	end
294	end
295	end
296	end
297	end
298	end
299	end
300	end
301	end
302	end
303	end
304	end
305	
306	if polje ==1
307	disp('16 mm field')
308	else if polje ==2
309	disp('8 mm field')
310 311	else if polje ==3 disp('4 mm field')
312	end
313	end
314	end
315	
316	
317	x0=0; y0=0; z0=0;
318	
319	if polje == 1 polje == 2
320	
321	if k==7
322	fx=@(x)
323	(c0s+A1s.*erf((x+a1s)./b1s)+A2s.*erf((x+a2s)./b2s)+A3s.*erf((x+a3s)./b3s)+A4s.*erf((x+a4s)./
324	b4s));
325	fy=@(y)
326	(c0s+A1s.*erf((y+a1s)./b1s)+A2s.*erf((y+a2s)./b2s)+A3s.*erf((y+a3s)./b3s)+A4s.*erf((y+a4s).
327	/b4s));
328	fzz=@(z)(c0z+A1z.*erf((z+a1z)./b1z)+A2z.*erf((z+a2z)./b2z)+A3z.*erf((z+a3z)./b3z)+A4z.*erf(
329	(z+a4z)./b4z)+A5z.*erf((z+a5z)./b5z)+A0z.*erf(az./bz));

```
330
            rr2=@(x,y,z)sqrt((x-x0).^2+(y-y0).^2+(z-z0).^2);
331
            fxx=@(x,y,z)
            (c0s+A1s.*erf((rr2(x,y,z)+a1s)./b1s)+A2s.*erf((rr2(x,y,z)+a2s)./b2s)+A3s.*erf((rr2(x,y,z)+a3s)./
332
            b3s)+A4s.*erf((rr2(x,y,z)+a4s)./b4s));
333
334
            fyy=@(x,y,z)
            (c0s+A1s.*erf((rr2(x,y,z)+a1s)./b1s)+A2s.*erf((rr2(x,y,z)+a2s)./b2s)+A3s.*erf((rr2(x,y,z)+a3s)./
335
336
            b3s)+A4s.*erf((rr2(x,y,z)+a4s)./b4s));
            fz1=@(x,y,z)(c0z+A1z.*erf((rr2(x,y,z)+a1z)./b1z)+A2z.*erf((rr2(x,y,z)+a2z)./b2z)+A3z.*erf((rr2
337
            (x,y,z)+a3z)./b3z)+A4z.*erf((rr2(x,y,z)+a4z)./b4z)+A5z.*erf((rr2(x,y,z)+a5z)./b5z)+A0z.*erf(az.
338
339
            /bz));
340
            fz2=@(x,y,z)(c0z+A1z.*erf((-rr2(x,y,z)+a1z)./b1z)+A2z.*erf((-rr2(x,y,z)+a2z)./b2z)+A3z.*erf((-
341
            rr2(x,y,z)+a3z)./b3z)+A4z.*erf((-rr2(x,y,z)+a4z)./b4z)+A5z.*erf((-
342
            rr2(x,y,z)+a5z)./b5z)+A0z.*erf(az./bz));
            fedgeplus=@(x,y,z) sqrt((fxx(x,y,z).*abs(x-x0)./rr2(x,y,z)).^2+(fyy(x,y,z).*abs(y-x0)./rr2(x,y,z)).^2+(fyy(x,y,z).*abs(y-x0)./rr2(x,y,z)).^2+(fyy(x,y,z).*abs(y-x0)./rr2(x,y,z)).^2+(fyy(x,y,z).*abs(y-x0)./rr2(x,y,z)).^2+(fyy(x,y,z).*abs(y-x0)./rr2(x,y,z)).^2+(fyy(x,y,z).*abs(y-x0)./rr2(x,y,z)).^2+(fyy(x,y,z).*abs(y-x0)./rr2(x,y,z)).^2+(fyy(x,y,z).*abs(y-x0)./rr2(x,y,z)).^2+(fyy(x,y,z).*abs(y-x0)./rr2(x,y,z)).^2+(fyy(x,y,z).*abs(y-x0)./rr2(x,y,z)).^2+(fyy(x,y,z).*abs(y-x0)./rr2(x,y,z)).^2+(fyy(x,y,z).*abs(y-x0)./rr2(x,y,z)).^2+(fyy(x,y,z).*abs(y-x0)./rr2(x,y,z)).^2+(fyy(x,y,z).*abs(y-x0)./rr2(x,y,z)).^2+(fyy(x,y,z).*abs(y-x0)./rr2(x,y,z)).^2+(fyy(x,y,z).*abs(y-x0)./rr2(x,y,z)).^2+(fyy(x,y,z).*abs(y-x0)./rr2(x,y,z)).^2+(fyy(x,y,z).*abs(y-x0)./rr2(x,y,z)).^2+(fyy(x,y,z).*abs(y-x0)./rr2(x,y,z)).^2+(fyy(x,y,z).*abs(y-x0)./rr2(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(
343
            y0)./rr2(x,y,z)).^2+(fz1(x,y,z).*abs(z-z0)./rr2(x,y,z)).^2);
344
345
            fedgeminus=@(x,y,z) sqrt((fxx(x,y,z).*abs(x-x0)./rr2(x,y,z)).^2+(fyy(x,y,z).*abs(y-
            y0)./rr2(x,y,z)).^2+(fz2(x,y,z).*abs(z-z0)./rr2(x,y,z)).^2);
346
347
            integral1=triplequad(fedgeplus,xmin,xmax,ymin,ymax,0.0000001,zmax);
348
349
            integral2=triplequad(fedgeminus,xmin,xmax,ymin,ymax,zmin,-0.0000001);
350
            disp('Volume averaging for EDGE detector is: ')
351
            kvol=V/(integral1+integral2) %kvol for EDGE detector
352
            return
353
                end
            s0=sqrt(x0.^2+y0.^2);
354
355
            rr1=@(s,z) sqrt((s-s0).^2+(z-z0).^2);
356
            %function for s direction for 16 & 8 mm field
357
            fss=@(s)
            (c0s+A1s.*erf((s+a1s)./b1s)+A2s.*erf((s+a2s)./b2s)+A3s.*erf((s+a3s)./b3s)+A4s.*erf((s+a4s)./
358
359
            b4s));
360
            fs1=@(s,z)
            (c0s+A1s.*erf((rr1(s,z)+a1s)./b1s)+A2s.*erf((rr1(s,z)+a2s)./b2s)+A3s.*erf((rr1(s,z)+a3s)./b3s)+
361
362
            A4s.*erf((rr1(s,z)+a4s)./b4s));
363
            %function for z direction for 16%8 mm field size
364
            fzz=@(z)(c0z+A1z.*erf((z+a1z)./b1z)+A2z.*erf((z+a2z)./b2z)+A3z.*erf((z+a3z)./b3z)+A4z.*erf(
365
366
            (z+a4z)./b4z)+A5z.*erf((z+a5z)./b5z)+A0z.*erf(az./bz));
            fz1=@(s,z)(c0z+A1z.*erf((rr1(s,z)+a1z)./b1z)+A2z.*erf((rr1(s,z)+a2z)./b2z)+A3z.*erf((rr1(s,z)+
367
            a3z)./b3z)+A4z.*erf((rr1(s,z)+a4z)./b4z)+A5z.*erf((rr1(s,z)+a5z)./b5z)+A0z.*erf(az./bz));
368
            fz2=@(s,z)(c0z+A1z.*erf((-rr1(s,z)+a1z)./b1z)+A2z.*erf((-rr1(s,z)+a2z)./b2z)+A3z.*erf((-
369
370
            rr1(s,z)+a3z)./b3z)+A4z.*erf((-rr1(s,z)+a4z)./b4z)+A5z.*erf((-
371
            rr1(s,z)+a5z)./b5z)+A0z.*erf(az./bz));
372
373
            %distribution
374
            f1=@(s,z) sqrt((fs1(s,z).*abs(s-s0)./rr1(s,z)).^2+(fz1(s,z).*abs(z-z0)./rr1(s,z)).^2)*s;
            f2=@(s,z) sqrt((fs1(s,z).*abs(s-s0)./rr1(s,z)).^2+(fz2(s,z).*abs(z-z0)./rr1(s,z)).^2)*s;
375
```

```
377
                    else if polje == 3
378
379
                           if k==7
380
                    fx=@(x)
                    (c0s+A1s.*erf((x+a1s)./b1s)+A2s.*erf((x+a2s)./b2s)+A3s.*erf((x+a3s)./b3s)+A4s.*erf((x+a4s)./
381
382
                   b4s));
383
                   fy=@(y)
                   (c0s+A1s.*erf((y+a1s)./b1s)+A2s.*erf((y+a2s)./b2s)+A3s.*erf((y+a3s)./b3s)+A4s.*erf((y+a4s).
384
385
                   /b4s));
                   fzz=@(z)(c0z+A1z.*erf((z+a1z)./b1z)+A2z.*erf((z+a2z)./b2z));
386
387
                   rr2=@(x,y,z)sqrt((x-x0).^{2}+(y-y0).^{2}+(z-z0).^{2});
388
                   fxx=@(x,y,z)
389
                    (c0s+A1s.*erf((rr2(x,y,z)+a1s)./b1s)+A2s.*erf((rr2(x,y,z)+a2s)./b2s)+A3s.*erf((rr2(x,y,z)+a3s)./
                    b3s)+A4s.*erf((rr2(x,y,z)+a4s)./b4s));
390
                    fyy=@(x,y,z)
391
                   (c0s+A1s.*erf((rr2(x,y,z)+a1s)./b1s)+A2s.*erf((rr2(x,y,z)+a2s)./b2s)+A3s.*erf((rr2(x,y,z)+a3s)./
392
393
                    b3s)+A4s.*erf((rr2(x,y,z)+a4s)./b4s));
                    fz1=@(x,y,z)(c0z+A1z.*erf((rr2(x,y,z)+a1z)./b1z)+A2z.*erf((rr2(x,y,z)+a2z)./b2z));
394
                   fz2=@(x,y,z)(c0z+A1z.*erf((-rr2(x,y,z)+a1z)./b1z)+A2z.*erf((-rr2(x,y,z)+a2z)./b2z));
395
                   fedgeplus=@(x,y,z) sqrt((fxx(x,y,z).*abs(x-x0)./rr2(x,y,z)).^2+(fyy(x,y,z).*abs(y-x0)./rr2(x,y,z)).^2+(fyy(x,y,z).*abs(y-x0)./rr2(x,y,z)).^2+(fyy(x,y,z).*abs(y-x0)./rr2(x,y,z)).^2+(fyy(x,y,z).*abs(y-x0)./rr2(x,y,z)).^2+(fyy(x,y,z).*abs(y-x0)./rr2(x,y,z)).^2+(fyy(x,y,z).*abs(y-x0)./rr2(x,y,z)).^2+(fyy(x,y,z).*abs(y-x0)./rr2(x,y,z)).^2+(fyy(x,y,z).*abs(y-x0)./rr2(x,y,z)).^2+(fyy(x,y,z).*abs(y-x0)./rr2(x,y,z)).^2+(fyy(x,y,z).*abs(y-x0)./rr2(x,y,z)).^2+(fyy(x,y,z).*abs(y-x0)./rr2(x,y,z)).^2+(fyy(x,y,z).*abs(y-x0)./rr2(x,y,z)).^2+(fyy(x,y,z).*abs(y-x0)./rr2(x,y,z)).^2+(fyy(x,y,z).*abs(y-x0)./rr2(x,y,z)).^2+(fyy(x,y,z).*abs(y-x0)./rr2(x,y,z)).^2+(fyy(x,y,z).*abs(y-x0)./rr2(x,y,z)).^2+(fyy(x,y,z).*abs(y-x0)./rr2(x,y,z)).^2+(fyy(x,y,z).*abs(y-x0)./rr2(x,y,z)).^2+(fyy(x,y,z).*abs(y-x0)./rr2(x,y,z)).^2+(fyy(x,y,z).*abs(y-x0)./rr2(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(
396
397
                   y0)./rr2(x,y,z)).^2+(fz1(x,y,z).*abs(z-z0)./rr2(x,y,z)).^2);
398
                   fedgeminus=@(x,y,z) sqrt((fxx(x,y,z).*abs(x-x0)./rr2(x,y,z)).^2+(fyy(x,y,z).*abs(y-x0)./rr2(x,y,z)).^2+(fyy(x,y,z).*abs(y-x0)./rr2(x,y,z)).^2+(fyy(x,y,z).*abs(y-x0)./rr2(x,y,z)).^2+(fyy(x,y,z).*abs(y-x0)./rr2(x,y,z)).^2+(fyy(x,y,z).*abs(y-x0)./rr2(x,y,z)).^2+(fyy(x,y,z).*abs(y-x0)./rr2(x,y,z)).^2+(fyy(x,y,z).*abs(y-x0)./rr2(x,y,z)).^2+(fyy(x,y,z).*abs(y-x0)./rr2(x,y,z)).^2+(fyy(x,y,z).*abs(y-x0)./rr2(x,y,z)).^2+(fyy(x,y,z).*abs(y-x0)./rr2(x,y,z)).^2+(fyy(x,y,z).*abs(y-x0)./rr2(x,y,z)).^2+(fyy(x,y,z).*abs(y-x0)./rr2(x,y,z)).^2+(fyy(x,y,z).*abs(y-x0)./rr2(x,y,z)).^2+(fyy(x,y,z).*abs(y-x0)./rr2(x,y,z)).^2+(fyy(x,y,z).*abs(y-x0)./rr2(x,y,z)).^2+(fyy(x,y,z).*abs(y-x0)./rr2(x,y,z)).^2+(fyy(x,y,z).*abs(y-x0)./rr2(x,y,z)).^2+(fyy(x,y,z).*abs(y-x0)./rr2(x,y,z)).^2+(fyy(x,y,z).*abs(y-x0)./rr2(x,y,z)).^2+(fyy(x,y,z).*abs(y-x0)./rr2(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+(fyy(x,y,z)).^2+
399
                   y0)./rr2(x,y,z)).^2+(fz2(x,y,z).*abs(z-z0)./rr2(x,y,z)).^2);
400
401
                    integral1=triplequad(fedgeplus,xmin,xmax,ymin,ymax,0.0000001,zmax);
402
                   integral2=triplequad(fedgeminus,xmin,xmax,ymin,ymax,zmin,-0.0000001);
403
                    disp('Volume avearging for EDGE SunNuclear detector is: ')
404
                    kvol=V/(integral1+integral2) %kvol for EDGE detector
405
                         return
406
                           end
407
408
                    s0=sqrt(x0.^2+y0.^2);
                    rr1=@(s,z) sqrt((s-s0).^2+(z-z0).^2);
409
410
                   %functions for s direction foe 4 mm field
411
                   fss=@(s)
                   (c0s+A1s.*erf((s+a1s)./b1s)+A2s.*erf((s+a2s)./b2s)+A3s.*erf((s+a3s)./b3s)+A4s.*erf((s+a4s)./
412
413
                    b4s));
414
                   fs1=@(s,z)
415
                   (c0s+A1s.*erf((rr1(s,z)+a1s)./b1s)+A2s.*erf((rr1(s,z)+a2s)./b2s)+A3s.*erf((rr1(s,z)+a3s)./b3s)+
                    A4s.*erf((rr1(s,z)+a4s)./b4s));
416
417
                    %functions for z direction for 4 mm field
418
419
                   fzz=@(z)(c0z+A1z.*erf((z+a1z)./b1z)+A2z.*erf((z+a2z)./b2z));
420
                   fz1=@(s,z)(c0z+A1z.*erf((rr1(s,z)+a1z)./b1z)+A2z.*erf((rr1(s,z)+a2z)./b2z));
421
                    fz2=@(s,z)(c0z+A1z.*erf((-rr1(s,z)+a1z)./b1z)+A2z.*erf((-rr1(s,z)+a2z)./b2z));
422
423
                    %rdistributions
```

424	f1=@(s,z) sqrt((fs1(s,z).*abs(s-s0)./rr1(s,z)).^2+(fz1(s,z).*abs(z-z0)./rr1(s,z)).^2)*s;
425	f2=@(s,z) sqrt((fs1(s,z).*abs(s-s0)./rr1(s,z)).^2+(fz2(s,z).*abs(z-z0)./rr1(s,z)).^2)*s;
426	12 = (3,2) $3(1((13+(3,2)), 30)(3,30))(11+(3,2)), 2((122(3,2)), 30)(2,20))(11+(3,2)), 2(-3)$
427	end
428	end
429	
430	if k==1
431	disp('Volume averaging for Semiflex TM31010 is: ')
432	else if k==2
433	disp('Volume averaging for PinPoint T31014 is: ')
434	else if k==3
435	disp('Volume averaging for Diode P T60016 is: ')
436	else if k==4
437	disp('Volume averaging for Diode E T60017 is: ')
438	else if k==5
439	disp('Volume averaging for PinPoint 3D T31022 is: ')
440	else if k==6
441	disp('Volume averaging for microDiamond T60019 is: ')
442	else if k==8
443	disp('Volume averaging for EFD is: ') else if k==9
444 445	disp('Volume averaging for RAZOR diode is: ')
446	else if k==10
447	disp('Volume averaging for RAZOR chamber is: ')
448	else if k==11
449	disp('Volume averaging for RAZORnano chamber is: ')
450	else if k==12
451	disp('Volume averaging for Semiflex 3D T31021 is: ')
452	else if k==13
453	disp('Volume averaging for IBA CC04 is: ')
454	else if k==14
455	disp('Volume averaging for Exradin W2 1X1 is: ')
456	else if k==15
457	disp('Volume averaging for Exradin W2 1X3 is: ')
458	end
459 460	end end
460 461	end
462	end
463	end
464	end
465	end
466	end
467	end
468	end
469	end
470	end

471	end
472	
473	
474	
475	if k~=11 % to exclude nanochamber since it had different geometry
476	
477	h=(s2-s1)/n; % integral for the cylinder in the area z>0
478	J1=0; J2=0; J3=0;
479	for i=0:n
480	s = s1 + i*h;
481	HX = (z2(s)-z00(s))/m;
482	K1 = f1(s, z00(s)) + f1(s, z2(s));
483	K2 = 0; K3 =0;
484	for j=1:m-1
485	z = z00(s) + j*HX;
486	Q = f1(s,z);
487 488	if rem(j,2) == 0 K2 = K2 + Q;
480 489	else
489 490	K3 = K3 + Q;
491	end
492	end
493	L = (K1 + 2*K2 + 4*K3)*HX/3;
494	if i == 0 i == n
495	J1 = J1 + L;
496	else if $rem(i,2) == 0$
497	J2 = J2 + L;
498	else
499	J3 = J3 + L;
500	end
501	end
502	end
503	Juk1 = 2 * pi* h * (J1 + 2*J2 + 4*J3)/3;
504	
505	h=(s2-s1)/n; %integral for the cylinder in the area z<0
506	J1=0; J2=0; J3=0;
507	for i=0:n
508	$s = s1 + i^{*}h;$
509	HX = (z00(s)-z1(s))/m;
510	K1 = f2(s, z1(s))+f2(s, z00(s));
511	K2 = 0; K3 =0;
512	$\frac{\text{for } j=1:m-1}{m-1}$
513 514	$z = z1(s) + j^{*}HX;$
514 515	Q = f2(s,z);
515 516	if rem(j,2) == 0 K2 = K2 + Q;
516	Rz = Rz + Q; else
21/	

518	K3 = K3 + Q;
519	end
520	end
521	L = (K1 + 2*K2 + 4*K3)*HX/3;
522	if i == 0 i == n
523	J1 = J1 + L;
524	else if rem(i,2) == 0
525	J2 = J2 + L;
526	else
527	J3 = J3 + L;
528	end
529	end
530	end
531	Juk2 = 2 * pi * h * (J1 + 2*J2 + 4*J3)/3;
532	
533	h=(s3-s1)/n; %integral for the electrode in the area z>0
534	J1=0; J2=0; J3=0;
	for i=0:n
535	
536	$s = s1 + i^{*}h;$
537	HX = (z3(s)-z00(s))/m;
538	K1 = f1(s, z00(s))+f1(s, z3(s));
539	K2 = 0; K3 =0;
540	for j=1:m-1
541	z = z00(s) + j*HX;
542	Q = f1(s,z);
543	if rem(j,2) == 0
544	K2 = K2 + Q;
545	else
546	K3 = K3 + Q;
547	end
548	end
549	L = (K1 + 2*K2 + 4*K3)*HX/3;
550	if i == 0 i == n
551	J1 = J1 + L;
552	else if rem(i,2) == 0
553	J2 = J2 + L;
554	else
555	J3 = J3 + L;
556	end
557	end
558	end
559	Juk3 = 2 * pi* h * (J1 + 2*J2 + 4*J3)/3;
560	
561	h=(s3-s1)/n; %integral for the electrode in the area z<0
562	J1=0; J2=0; J3=0;
563	for i=0:n
564	s = s1 + i*h;

```
565
         HX = (z00(s)-z1(s))/m;
         K1 = f2(s, z1(s))+f2(s, z00(s));
566
         K2 = 0; K3 =0;
567
         for j=1:m-1
568
         z = z1(s) + j^{*}HX;
569
570
          Q = f2(s,z);
          if rem(j,2) == 0
571
            K2 = K2 + Q;
572
573
          else
            K3 = K3 + Q;
574
575
          end
576
         end
577
         L = (K1 + 2*K2 + 4*K3)*HX/3;
         if i == 0 || i == n
578
579
           J1 = J1 + L;
         else if rem(i,2) == 0
580
              J2 = J2 + L;
581
582
         else
           J3 = J3 + L;
583
584
           end
585
         end
         end
586
         Juk4 = 2 * pi * h * (J1 + 2*J2 + 4*J3)/3;
587
588
        if k==1 % moving the field so the curvature part of a chamber could be in the right place
589
         z0=-z0-1.05;
590
       else if k==2
591
              z0=-z0-2.4294;
592
593
           else if k==5
                z0=-z0-0.275;
594
             else if k==10
595
596
                  z0=-z0-0.8;
597
               else if k==12
598
                    z0=-z0-0.75;
                  end
599
600
               end
601
             end
602
             end
        end
603
604
        rr1=@(s,z) sqrt((s-s0).^2+(z-z0).^2);
605
606
        fs1=@(s,z)
607
       (c0s+A1s.*erf((rr1(s,z)+a1s)./b1s)+A2s.*erf((rr1(s,z)+a2s)./b2s)+A3s.*erf((rr1(s,z)+a3s)./b3s)+
608
       A4s.*erf((rr1(s,z)+a4s)./b4s));
609
610
        if polje == 1 || polje == 2
```

611 612 fz1=@(s,z)(c0z+A1z.*erf((rr1(s,z)+a1z)./b1z)+A2z.*erf((rr1(s,z)+a2z)./b2z)+A3z.*erf((rr1(s,z)+ a3z)./b3z)+A4z.*erf((rr1(s,z)+a4z)./b4z)+A5z.*erf((rr1(s,z)+a5z)./b5z)+A0z.*erf(az./bz)); 613 else if polje == 3 614 fz1=@(s,z)(c0z+A1z.*erf((rr1(s,z)+a1z)./b1z)+A2z.*erf((rr1(s,z)+a2z)./b2z)); 615 616 end 617 end 618 f1=@(s,z) sqrt((fs1(s,z).*abs(s-s0)./rr1(s,z)).^2+(fz1(s,z).*abs(z-z0)./rr1(s,z)).^2)*s; 619 620 621 h=(s2-s1)/n; %calculation of an integral of top helispherical part 622 J1=0; J2=0; J3=0; 623 for i=0:n $s = s1 + i^{*}h;$ 624 625 HX = (zf(s)-z00(s))/m;626 K1 = f1(s, z00(s))+f1(s, zf(s)); K2 = 0; K3 =0; 627 for j=1:m-1 628 z = z00(s) + j*HX;629 Q = f1(s,z);630 631 if rem(j,2) == 0 K2 = K2 + Q;632 633 else K3 = K3 + Q;634 635 end 636 end L = (K1 + 2*K2 + 4*K3)*HX/3;637 if i == 0 || i == n 638 J1 = J1 + L;639 else if rem(i,2) == 0 640 641 J2 = J2 + L;642 else 643 J3 = J3 + L;644 end end 645 646 end 647 Juk5 = 2 * pi* h * (J1 + 2*J2 + 4*J3)/3; 648 649 kvol=V/(Juk1+Juk2+Juk5-Juk3-Juk4) 650 651 else if k==11 %RAZORnano chamber 652 653 h=(s2-s1)/n; %calculation of an integral of top half spherical part 654 J1=0; J2=0; J3=0; for i=0:n 655 $s = s1 + i^{*}h;$ 656 HX = (zf(s)-z00(s))/m;657

```
658
         K1 = f1(s, z00(s))+f1(s, zf(s));
659
         K2 = 0; K3 =0;
         for j=1:m-1
660
          z = z00(s) + j*HX;
661
662
          Q = f1(s,z);
663
          if rem(j,2) == 0
            K2 = K2 + Q;
664
665
          else
            K3 = K3 + Q;
666
          end
667
668
         end
669
         L = (K1 + 2*K2 + 4*K3)*HX/3;
         if i == 0 || i == n
670
            J1 = J1 + L;
671
672
         else if rem(i,2) == 0
              J2 = J2 + L;
673
         else
674
            J3 = J3 + L;
675
            end
676
677
         end
678
         end
679
         Juk1RAZOR = 2 * pi* h * (J1 + 2*J2 + 4*J3)/3;
680
681
         h=(s2-s1)/n; %calculation of an integral of bottom half spherical part
         J1=0; J2=0; J3=0;
682
         for i=0:n
683
684
         s = s1 + i^{*}h;
         HX=(z00(s)-zf(s))/m;
685
686
         K1 = f2(s, z00(s))+f2(s, zf(s));
         K2 = 0; K3 =0;
687
         for j=1:m-1
688
          z = z00(s) + j^{*}HX;
689
690
          Q = f2(s,z);
691
          if rem(j,2) == 0
692
            K2 = K2 + Q;
693
          else
            K3 = K3 + Q;
694
695
          end
         end
696
         L = (K1 + 2*K2 + 4*K3)*HX/3;
697
         if i == 0 || i == n
698
            J1 = J1 + L;
699
700
         else if rem(i,2) == 0
701
              J2 = J2 + L;
702
         else
703
            J3 = J3 + L;
704
            end
```

705 end 706 end Juk2RAZOR = 2 * pi* h * (J1 + 2*J2 + 4*J3)/3; 707 708 709 h=(s3-s1)/n; %calculation of an integral of top half spherical part of an electrode 710 J1=0; J2=0; J3=0; 711 for i=0:n 712 $s = s1 + i^{*}h;$ 713 HX = (zf2(s)-z00(s))/m;714 K1 = f1(s, z00(s))+f1(s, zf2(s)); 715 K2 = 0; K3 =0; 716 for j=1:m-1 717 $z = z00(s) + j^{*}HX;$ 718 Q = f1(s,z);719 if rem(j,2) == 0 720 K2 = K2 + Q;721 else K3 = K3 + Q;722 723 end 724 end 725 L = (K1 + 2*K2 + 4*K3)*HX/3;726 if i == 0 || i == n J1 = J1 + L;727 else if rem(i,2) == 0 728 729 J2 = J2 + L;else 730 731 J3 = J3 + L;end 732 733 end 734 end Juk3RAZOR = 2 * pi* h * (J1 + 2*J2 + 4*J3)/3; 735 736 737 h=(s3-s1)/n; %calculation of an integral of bottom half spherical part of an electrode 738 J1=0; J2=0; J3=0; for i=0:n 739 $s = s1 + i^{*}h;$ 740 HX = (z00(s) - zf2(s))/m;741 K1 = f2(s, z00(s))+f2(s, zf2(s)); 742 743 K2 = 0; K3 =0; 744 for j=1:m-1 $z = z00(s) + j^{*}HX;$ 745 746 Q = f2(s,z);747 if rem(j,2) == 0 748 K2 = K2 + Q;749 else 750 K3 = K3 + Q;751 end

752	end
753	L = (K1 + 2*K2 + 4*K3)*HX/3;
754	if i == 0 i == n
755	J1 = J1 + L;
756	else if rem(i,2) == 0
757	J2 = J2 + L;
758	else
759	J3 = J3 + L;
760	end
761	end
762	end
763	Juk4RAZOR = 2 * pi* h * (J1 + 2*J2 + 4*J3)/3;
764	
765	kvol=V/(Juk1RAZOR + abs(Juk2RAZOR)-abs(Juk3RAZOR)-abs(Juk4RAZOR))
766	end
767	end

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List of abbreviations

Α

AAPM
American Association of Physicists in Medicine11
ABS
Acrylonitrile Butadiene Styrene14

С

CBCT
Cone-Beam Commputed Tomography63
clin
Clinical field26
Co-60
Cobalt 6011
СоР
Code of practice11
CPE
Charge particle equilibrium21

D

DSCF	
Detector - specific correction factor12	2

Ε

EBT3	
External Beam Therapy 348	

F

f _{msr}	
machine-specific reference field	0
fref	
reference field3	0
FWHM	
Full width at half maxima2	3

G

GK
Gamma Knife11

I

IAEA
IALA
International Atomic Energy Agency11

J

JCGM	JC
Joint Committee for Guides in Metrology55	

Κ

k _{vol}	
Volume averaging correction factor	28

L

LCPE	
Lateral charge particle equilibrium	20
LGK	
Leksell Gamma Knife	11

Μ

MC	
Monte Carlo	14
msr	
machine-specific reference field	30

Ν

netOD	
Net optical density	

0

	OAR	
	Off-axis ratio	27
OF	7	
	field output factors	12

Ρ

PSD	
Plastic scintiator detector	. 16

Q

Q_0	
Beam quality with Co-60 3	0
Q _{clin}	
Beam qualiy in clinical fields 2	6
Q _{msr}	
Beam quality in machine-specific reference field 3	0

R

ROI	
Region of interest	48

S	Transient charge particle equilibrium
	TPR
SRS	Tissue-phantom ratio 21
Stereotactic radiosurgery11	TPS
	The Planning System 12
	TRS
	Technical report series 11

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