CT beam dosimetric characterization procedure for personalized dosimetry

S Rosendahl \textsuperscript{1}, L B"uermann \textsuperscript{1,2}, M Borowski \textsuperscript{1}, M Kortesniemi \textsuperscript{3}, V-M Sundell \textsuperscript{1}, A Kosunen \textsuperscript{4} and T Siiskonen \textsuperscript{4}

\textsuperscript{1} Physikalisch-Technische Bundesanstalt, Bundesallee 100, 38116 Braunschweig, Germany
\textsuperscript{2} St"{a}dtisches Klinikum Braunschweig gGmbH, Salzdahlumerstr. 90, 38126 Braunschweig, Germany
\textsuperscript{3} HUS Medical Imaging Center, Helsinki University Central Hospital, Haartmaninkatu 4, 00290 Helsinki, Finland
\textsuperscript{4} STUK—Radiation and Nuclear Safety Authority, Laippatie 4, 00880 Helsinki, Finland

E-mail: ludwig.bueermann@ptb.de

Keywords: computed tomography, personalized radiation dosimetry, Monte Carlo simulation, equivalent source models

Abstract

Personalized dosimetry in computed tomography (CT) can be realized by a full Monte Carlo (MC) simulation of the scan procedure. Essential input data needed for the simulation are appropriate CT x-ray source models and a model of the patient’s body which is based on the CT image. The purpose of this work is to develop comprehensive procedures for the determination of CT x-ray source models and their verification by comparison of calculated and measured dose distributions in physical phantoms. Mobile equipment together with customized software was developed and used for non-invasive determination of equivalent source models of CT scanners under clinical conditions. Standard and physical anthropomorphic CT dose phantoms equipped with real-time CT dose probes at five representative positions were scanned. The accumulated dose was measured during the scan at the five positions. ImpactMC, an MC-based CT dose software program, was used to simulate the scan. The necessary inputs were obtained from the scan parameters, from the equivalent source models and from the material-segmented CT images of the phantoms. 3D dose distributions in the phantoms were simulated and the dose values calculated at the five positions inside the phantom were compared to measured dose values. Initial results were obtained by means of a General Electric Optima CT 660 and a Toshiba (Canon) Aquilion ONE. In general, the measured and calculated dose values were within relative uncertainties that had been estimated to be less than 10%. The procedures developed were found to be viable and rapid. The procedures are applicable to any scanner type under clinical conditions without making use of the service mode with stationary x-ray tube position. Results show that the procedures are well suited for determining and verifying the equivalent source models needed for personalized CT dosimetry based on post-scan MC calculations.

1. Introduction

New hardware and software strategies in medical computed tomography (CT) have led to steady improvements in this type of imaging modality. Advanced x-ray tubes, new types of photon detectors for imaging, dynamic x-ray fluence as a function of the table position and the rotation angle (tube current modulation, TCM), spectral CT (e.g. dual source), spectral and spatial filtration, automatic selection of the tube potential, dynamic bow-tie (BT) filters and collimation are only some of the new hardware strategies (McCullough et al 2006, Ginat and Gupta 2014). New software modules have been developed such as iterative reconstruction algorithms and, more recently, methods based on deep learning and convolutional neural networks (Sahiner et al 2019). All these new strategies have significant potential to lower the radiation dose to the patient while maintaining the necessary application-specific image quality. However, despite these promising developments, the patient dose in CT is still a concern. CT procedures deliver approximately 50\%–60\% of the collective effective dose from medical and dental exposures in many countries due to the relatively high dose of CT procedures compared with other diagnostic imaging modalities (NCRP 2009, European Commission 2014).
A recent topical review (Kalender 2014) provides an excellent overview about the past, current and future directions of x-ray CT dosimetry. Two measurable dose quantities must be indicated on every CT scanner: the volume computed tomography dose index (CTDIvol) and the dose-length product (DLP) (Shope et al 1981, IEC 60601 2016). Both quantities are useful for quantifying x-ray tube dose output and comparing dose levels. Therefore, these two quantities are used as diagnostic reference levels (Vanó et al 2017). Although they are correlated with patient dose levels, they should not be regarded as a patient dose (McCollough et al 2011). In order to obtain generic stochastic radiation risk estimates, which are expressed in terms of an effective dose \(E\), this quantity is evaluated from the DLP by \(E = k \cdot \text{DLP}\), where \(k\) is a body-region-specific normalized effective dose conversion coefficient, usually calculated by means of MC simulations for reference scanners and reference patients. Such \(k\)-factors are tabulated in published reports, e.g. Deak et al (2010) and Brady et al (2015). However, this method of dose estimation has many shortcomings, due not only to the known limitations of the basic dose quantities (Boone 2007, Dixon and Boone 2010, 2013, Dixon and Boone 2014) if used in advanced-technology CT scanners (e.g. with TCM) or for stationary table CT but also to the fact that they are neither scanner-specific nor patient-specific.

In order to overcome the limitations given by the fixed sizes of the two CTDI phantoms, the American Association of Physicists in Medicine (AAPM) introduced the concept of size-specific dose estimates (SSDE) (Boone et al 2011, McCollough et al 2014). Although SSDE values correlate much better with the patient size and thus with the patient dose, they have the disadvantage that they are based on the conventional definition of CTDI with its known limitations. Furthermore, using SSDE to estimate organ dose is possible (e.g. Brady et al (2015)) but limited.

A promising new approach of patient-specific dose estimates (PSDE), which was described in the above-mentioned review paper (Kalender 2014), is based on fast Monte Carlo (MC) simulations and is independent of CTDI metrics. This approach makes use of the acquired patient CT data and combines it with a best-fitting pre-described voxel phantom to obtain a whole-body data set. Next, an MC dose calculation is performed by simulating the whole scan. An overlay of the calculated 3D dose distribution and the organ contours of the CT image allows organ dose values to be determined. The procedure, together with the necessary input information, is illustrated in figure 1.

A fast MC dose calculation tool for use in clinical CT on-site and in real time has been developed and validated against measurements (Schmidt and Kalender 2002, Deak et al 2008, Myronakis et al 2009, Chen et al 2012). Although this new approach has the potential to allow real-time scanner-specific and patient-specific dose estimations and indications on the scanner console to be made, it is still in the research stage. One reason for this may be the lack of standardized practical procedures for this application. Procedures for automatic segmentation of organs or anatomical structures in CT images are still challenging. Such segmentation can be performed manually but is a time-consuming process. However, new, promising procedures based on convolutional neural networks have been reported in literature (Weston 2018) and could represent a solution. Another problem is that the MC simulations require scanner-specific input data such as x-ray spectrum and subsequent filtration, including bow tie-shaped form filters, which are proprietary information that is not usually provided to the user. To overcome this restriction, some efforts have been made in recent years to develop methods that allow the determination of so-called ‘equivalent x-ray source models’ that consist of a photon energy spectrum and filtration description that are based entirely on measured values (Kruger et al 2000, Turner et al 2009, McKenney et al 2014, Randazzo and Tambasco 2015, Alikhani and Buermann 2016). Finally, even if feasible procedures for PSDE are available, it may still be a time-consuming task to validate the dose simulation software by measurements at different scanner types.

The main purpose of this work is to develop standardized practical procedures for personalized CT dosimetry under clinical conditions based on the application of the above-mentioned PSDE approach. Except for the automatic segmentation of organ tissues in CT images, all other tools necessary for the realization and validation of the PSDE approach were (in principle) already available when this work was started. These tools are as follows: fast MC simulation codes for CT applications; non-invasive procedures for the determination of equivalent source models at different scanner types; and real-time dose detectors and anthropomorphic phantoms for the validation of calculated dose values by means of measurements. However, these different tools have never been combined to a complete tool box to establish the dosimetry of the PSDE procedure at CT scanners under clinical conditions. A prototype of such a tool box was developed and is described in this paper. It consists of a newly designed mobile equipment for the non-invasive determination of equivalent CT source models, different physical phantoms equipped with different real-time dose probes and a fast MC dose calculation program. Note that the tool box does not make use of the service mode of the scanner, which has been one of the main limitations in the past to implement MC in CT.

2. Materials and methods

2.1. CT scanners
Two scanner types were used for the studies: general electric (GE) Optima CT 660 (General Electric Company, GE Healthcare) located at the PTB for research purposes and Toshiba’s Aquilion ONE (Canon Medical Systems
Europe B.V.) located at the Städtisches Klinikum Braunschweig. The 64-slice Optima CT 660 offers x-ray tube high voltages of 80 kV, 100 kV, 120 kV and 140 kV. The anode angle is 7°. The collimation widths can be varied in the range 1.25 mm to 40 mm. It is equipped with a small (head) and a large (body) BT filter. The 320-slice Aquilion ONE offers x-ray tube high voltages of 80 kV, 100 kV, 120 kV and 135 kV. The anode angle is 11°. The collimation widths can be varied in the range 2 mm to 160 mm. It is equipped with a small, medium and large BT.

2.2. Devices for dose measurements

The dose measurements are performed with various detectors. Different ionization chambers are used: a farmer type chamber (Radcal, type RC0.6) and two pencil-type CT chambers 100 mm and 300 mm in length (PTW, type 30009 and 30017, respectively). While the RC0.6 is capable for measurements of the air kerma $K_a$, the CT chambers are used to measure the air kerma length product $K_aL$. The current from the ionization chambers is measured with an electrometer (PTW, type UNIDOS). $K_a$ and $K_aL$ are given by the following expressions:

$$K_a = M_Q \cdot N_{K_a,Q_0} \cdot k_{Q,Q_0} \cdot k_\rho \quad (1)$$

$$K_aL = M_Q \cdot N_{K_a,Q_0,L} \cdot k_{Q,Q_0} \cdot k_\rho \quad (2)$$

Here, $M_Q$ is the charge measured at the electrometer at radiation quality $Q$ and $N_{K_a,Q_0}$ and $N_{K_a,Q_0,L}$ are the calibration factors at reference quality $Q_0$ determined using the primary air kerma standard at the x-ray facilities of PTB. $k_{Q,Q_0}$ is the quality correction factor that takes into account the energy dependence of the detector for different x-ray qualities that can appear in the CT, e.g. 80 kV against 140 kV. This value is set to $k_{Q,Q_0} = 1$ since the response of the used ionization chambers with respect to the air kerma is nearly independent from the spectra in the energy range between 30 keV and 150 keV (Alikhani and Büermann 2016, Siiskonen and Suutari 2018). Small deviations are taken into account via the uncertainties given in table 1, which has been validated using the methods presented in Siiskonen and Suutari (2018). Uncertainty estimations within this work were performed according to the Guide to the expression of uncertainty in measurement (GUM) (Bureau International des Poids et Mesures, BIPM 2008).

Finally, $k_\rho$ is the correction factor for the air density (see table 1) calculated from ambient pressure and temperature, which is measured using quality-controlled, high accuracy sensors.

The RC0.6 is also used in combination with a digitizer read-out (Radcal, ACCU-Gold) that allows time-resolved measurement of the dose rate to be taken.

In addition to ionization chambers, semiconductor detectors (RTI, dose profiler) are used for measurements of the air kerma. The advantage of these detectors is their high spatial resolution of less than 0.25 mm (RTI Electronics AB 2015) in longitudinal ($z$-) direction and their higher sensitivity. Furthermore, the measurement sys-

---

5 In 2016, Canon Inc. took over the medical unit of Toshiba Corp.
Table 1. Uncertainty budget for air kerma measurements on CT scanners. This table summarizes the type A (statistical) and B (evaluated by other means) uncertainties according to GUM (Bureau International des Poids et Mesures, BIPM 2008) for dose measurements on CT scanners presented in this work using different ionization chambers and semiconductor detectors. The uncertainty of $k_{Q,Q_0}$ of the semiconductor detector (0.5%) was put into brackets because it is included in $k_{Ph}$. Thus it must not be used for the combined uncertainty if the detector is used in the phantom.

| Description of component | Symbol | Uncertainty type | Pencil type chamber | Farmer type chamber | Semiconductor dose profiler |
|--------------------------|--------|------------------|---------------------|---------------------|---------------------------|
| Reproducibility (example)| $\sigma_{rep}$ | A | 0.46 | 0.28 | 0.30 | 1.13 |
| Calibration factor (RQT-9)| $N_{K,\text{is}}$ | B | 0.40 | 0.20 | 0.40 | — |
| Energy dependence (Free in air) | $k_{Q,Q_0}$ | B | 0.45 | 0.81 | 0.38 | (0.50) |
| Energy dependence (In phantom) | $k_{Ph}$ | B | — | — | — | 1.05 |
| Reproducibility (e.g. positioning) | $k_{Ph}$ | B | 0.80 | 0.43 | 0.34 | 0.59 |
| Air density correction | $k_{\rho}$ | B | 0.14 | 0.14 | 0.14 | — |
| Combined type B uncertainty ($k = 1$) | $\sigma_{B}$ | — | 1.01 | 0.95 | 0.66 | 1.21 |
| Combined uncertainty | $\sigma_{c}$ | — | 1.11 | 0.99 | 0.73 | 1.65 |

$\sqrt{\sigma_{A}^2 + \sigma_{B}^2 (k = 1)}$

\[
K_a = M_Q \cdot N_{K,\text{is}} \cdot k_{Q,Q_0} \cdot k_{Ph}.
\] (3)

In contrast to ionization chambers, air density correction is not needed; here, the correction of the energy dependence is more complicated. The Barracuda system already gives a calibrated value for air kerma $K_a = M_Q \cdot N_{K,\text{is}}$, but the manufacturer uses only one x-ray quality (RQR-9) for calibration. For tube voltages between 80 kV and 140 kV, significant differences of up to \( \approx 30\% \) can be observed due to the spectral dependence of the detector. Hence, additional calibrations with qualities from the ISO4037 narrow-spectrum series (International Organization for Standardization 1997) are performed at PTB in order to calculate correction factors $k_{Q,Q_0}$ for each tube potential using the CT-specific spectra obtained from Al attenuation measurements (compare (Siiskonen and Suutari 2018)). The comparison of a spectrally corrected dose profiler with the RQ0.6 chamber, both of which measure the air kerma successively at the isocenter of the Optima 660, leads to differences of less than 2%. This is in agreement with the uncertainties stated in table 1 and indicates that the correction factors are correct. However, the spectrum of the CT changes inside the phantom due to attenuation and scattering. For this reason, an additional factor $k_{Ph}$ is introduced, taking the spectral modulation caused by the material of the phantom into account. $k_{Ph}$ is estimated by comparing the air kerma measurement of the dose profiler after spectral correction using $k_{Q,Q_0}$ with the values measured with the ionization chamber RQ0.6 in the phantom. The direct calibration is performed for the different standard and anthropomorphic phantoms using the tube voltages that are relevant for this study. For the usage of ionization chambers inside the phantom, $k_{Ph}$ is not needed, since the energy dependence is negligible and covered by the uncertainty, as mentioned above.

2.3. Mobile measurement equipment and equivalent CT source determination

2.3.1. Characterization of x-ray spectra

X-ray spectra can be determined by measuring aluminum attenuation characteristics. This method is well established and widely used (Kruger et al 2000, Turner et al 2009, McKenney et al 2014, Randazzo and Tambasco 2015, Alikhani and Büermann 2016). Conventional spectrometry using semiconductor detectors made of materials such as germanium is complex and currently only possible by measuring Compton-scattered photons to obtain the original spectrum via deconvolution (Bazalova and Verhaegen 1988, Matscheko and Carlsson 1988, Duisterwinkel et al 2015). Measuring x-ray attenuation as a function of increasing aluminum thickness with an ionization chamber provides easy access to a very accurate approximation of the real spectrum. Typical experimental procedures using the service mode of the CT scanner are presented in Turner et al (2009) and Alikhani and Büermann (2016), while more advanced setups have been developed using rotating x-ray sources and real-time dose measurements (McKenney et al 2014, Randazzo and Tambasco 2015). For the mobile measurement system, a time-resolved read-out is used in order to measure (in spiral mode) the attenuation caused by aluminum plates of different thickness that are placed on a holding frame on the patient table, as...
shown in figure 2. For this purpose, the RC0.6 probe together with the ACCU Gold digitizer is fast enough for rotation times of ≈ 1 s.

The detector is placed on the x-axis away from the isocenter to match \( z = y = 0 \) with a distance between the source and the isocenter of \( x = r_p \approx 25 \) cm. Hence, contamination of the signal by scattered photons from the aluminum sheets on the one hand and back-scattered photons from the detector array of the CT on the other hand are reduced. The collimation in z-direction \( \Delta_z \) should be chosen in such a way that the ionization chamber is fully covered by the beam, e.g. \( \Delta_z \approx 2 \) cm. A larger collimation produces additional scattered photons that may influence the results. The measurement procedure includes an initial overview scan in order to define scan areas centered at the single aluminum plates. This simplifies the data analysis and ensures that a certain distance between the scan area and the holding frame is maintained in order to reduce scattering from the frame. Typically, nine scan areas are defined: one free in air without aluminum as a reference and eight aluminum sheets with thicknesses between 1 mm and 20 mm and a purity of 99.999%. The homogeneity in the thickness of the plates, as well as their absolute thicknesses, have been measured on site and shown to be better than 1%. The parameters normally chosen for the scan are a rotation time of 1 s and a pitch of ≈ 0.5. The pitch depends on the table velocity and collimation and the possible values are scanner-specific. Hence, the total scan time varies for the different manufacturers, but is in the range of ≈ 2 min.

In figure 3, the time-resolved dose rate for one rotation of the x-ray source is shown. The characteristic behavior is caused by the inverse square law and the attenuation of the BT Filter. The maximum of intensity at 1 s and 2 s are the position where the source and the detector are at their closest position. The region of interest where the aluminum sheet, the detector and the source are aligned (at \( \sim 1.5 \) s) shows the attenuation and is gained after a half rotation when the source, the aluminum sheet and the detector are aligned (compare schematic drawing in figure 2). It is expressed as the ratio \( \kappa_{\text{Meas}} \) of \( K_x \) to \( K_0 \), which are the air kerma with and without the added aluminum sheet of the thickness \( x_{\text{Al}} \):

\[
\kappa_{\text{Meas}} = \frac{K_x}{K_0}, \quad (4)
\]

The signal inhomogeneities after the attenuation from the Al layer are caused by the patient table and the holding frame. Scattering from the table and the holding frame may bias the attenuation measurements. The influence of this effect was studied by comparison with results obtained from conventional static measurements with stationary x-ray tube at the 6 o’clock position and Al sheets positioned close to the gantry (as described in our previous work Alikhani and Büermann (2016)). It turned out that attenuation curves measured with the static and dynamic method were indistinguishable. Therefore, it was not necessary to include this effect as an additional uncertainty component. In a further analysis, \( \kappa_{\text{Meas}} \) will be approximated by a corresponding calculated value, \( \kappa_{\text{Calc}} \), according to equation (5) by varying the thickness of the total filtration \( d_{\text{Al}} \).

\[
\kappa_{\text{Calc}} = \frac{K_x^{\text{Calc}}(x_{\text{Al}})}{K_0^{\text{Calc}}} = \frac{\int \Phi_E e^{-\mu_{\text{Al}}(E)} dx e^{-\mu_{\text{Al}}(E)} s_{\text{Al}} E(\frac{\mu_{\text{Al}}}{\rho})_{\text{Al}} dE}{\int \Phi_E e^{-\mu_{\text{Al}}(E)} dx e^{-\mu_{\text{Al}}(E)} s_{\text{Al}} E(\frac{\mu_{\text{Al}}}{\rho})_{\text{Al}} dE}, \quad (5)
\]

where \( \Phi_E \) is the non-filtered x-ray spectrum and \( d_{\text{Al}} \) is the quality equivalent filtration thickness of aluminum. Note that \( d_{\text{Al}} \) is the aluminium equivalent filtration of all non-removable filtration parts in the beam composed of the inherent filtration of the tube, the centre of the BT filter and the Mylar window. \( \Phi_E \) is calculated using the
SpecCalc software tool (Poludniowski and Evans 2007a, 2007b, Poludniowski et al 2009), which requires the tungsten anode angle and the tube high voltage as inputs. $d_{Al}$ is obtained by minimizing $\Delta(d_{Al})$, given as

$$\Delta(d_{Al}) = \sum_{i} |\kappa_{Meas}(x_i) - \kappa_{Calc}(d_{Al}, x_i)|.$$  

(6)

In-house analysis software reads the measured data and identifies the regions of interest for the calculation of the attenuation curve. Due to the small pitch factor and collimation, multiple rotations per Al sheet can be used to calculate the average $\kappa_{Meas}(x_i)$.

2.3.2. Characterization of BT filters

The BT filter is characterized using the COBRA formalism (characterization of bow-tie relative attenuation) proposed by Boone (2010). The geometry for the mobile measurement system has been chosen in order to additionally collect the data relevant for the determination of the attenuation profile without additional hardware re-arrangements, which represents a significant advantage of the system. The attenuation is measured performing a dedicated free-in-air scan with a collimation of $\Delta_Z = 4$ cm. A smaller collimation is not possible because it prevents full coverage of the detector when the distance between the source and the detector is too small. For the following COBRA analysis, which was performed to determine the attenuation characteristic of the BT filter, we strictly followed the calculation process of Boone (2010), transforming the measured dose from the time domain $I(t)$ into the rotation domain $I(\alpha)$, with $\alpha$ being the rotation angle, as shown in figure 4.

It is assumed that, at rotation angles $\alpha = 0$ and $\alpha = \pi$, where the x-ray source, the probe and the isocenter are aligned (collinear), the thickness of the BT filter is zero for the calculations, as the attenuation in the center of the BT (fan angle $\theta = 0$) is at its minimum (Boone 2010, McKenney 2011). The detector measures the highest possible intensity $I_0(0)$ at $\alpha = 0$, which modulates according to the inverse square law while the source rotates. $I_0(\alpha)$ is calculated using the distances between the source and the isocenter $s_{iso}$ and between the source and the detector $g(\alpha)$. The attenuation $F(\theta)$, caused by the BT, is calculated as the ratio between the measured intensity $I_{att}(\alpha)$ and $I_0(\alpha)$ according to equation (7), where $r_P$ is the distance between the detector and the isocenter (Boone 2010, McKenney 2011).

$$F(\theta) = \frac{I_{att}(\alpha)}{I_0(\alpha)}$$  

with  

$$I_0(\alpha) = I_0(0) \left( \frac{s_{iso} - r_P}{s_{iso}} \right)^2 \cdot \left( \frac{s_{iso}}{g(\alpha)} \right)^2.$$  

(7)

In figure 4 (right), the relative attenuation profile measured on an Optima 660 is shown for the large BT. In order to perform a quality check of the implementation, the attenuation curve obtained from the dynamic COBRA mode (blue dots) is compared with the curve obtained from static dose profile measurements (red dots) where the x-ray tube is fixed at the 12 o’clock position and the chamber is moved along the x-axis using the time-resolved integration of the charge (TRIC) method described in Alikhani and Büermann (2016), which is in good agreement. Note that the static TRIC method needs a fixed x-ray source which can only be achieved in the service mode of the CT scanner. The analysis was performed using rotation angles in the range of $\alpha_{min} < \alpha < \alpha_{max}$,
while the minimum and maximum rotation angles $\alpha_{\text{min}}$ and $\alpha_{\text{max}}$ depend on $r_P$. The measurements from the other angles are neglected in this work, since the distance between the probe and the detector is very small; this leads to significant contamination, disturbing the attenuation profile, which should be measured in narrow beam geometry.

In the standard COBRA formalism, the thickness at the center of the BT is assumed to be zero for the calculations. This initial thickness at the center is usually included in the equivalent filtration of the spectrum that is achieved in dynamic measurements of the aluminum attenuation characteristics. Consequently, in the MC simulations, the equivalent BT filter can be used only in combination with the associated x-ray spectrum containing the correct equivalent filtration, which is a potential source of errors when incorrect files are combined. To reduce this uncertainty, the measured total filtration $d_{\text{Al}}$ is treated as part of the BT filter, allowing the unfiltered spectrum to be used for the simulation. Here, the advantage is that the Al-equivalent filter geometry derived in this way contains all the specific information from the bow tie as well as from the total filtration. The unfiltered x-ray spectrum used in the simulations is independent from the BT filter used. From the relative attenuation $F(\theta)$, an equivalent Al filter is calculated as described in e.g. Alikhani and Büermann (2016) and Boone (2010).

### 2.4. MC simulation with ImpactMC

#### 2.4.1. Material description

ImpactMC is a simulation tool that traces its origins to works by Schmidt and Kalender (2002). For this study, Version 1.5.1 of ImpactMC in GPU mode is used. The user can control a wide range of parameters that allow scanner-specific x-ray spectra and BT-filter geometries to be included. Furthermore, asymmetric collimation and tube current modulation can be incorporated (CT Imaging GmbH 2014), although these techniques are not investigated here. Of crucial importance for obtaining accurate simulation results is the material conversion feature of ImpactMC, which is needed in order to link the Hounsfield units (HU) from the DICOM file to the specific material.

The validation process presented in this work was carried out with two different phantoms: the standard CTDI phantoms made of polymethylmethacrylate (PMMA) with a diameter of 32 cm (body) and a ‘Thorax-Medium Adult’ anthropomorphic phantom (Mod. Nr. 007TE-17) manufactured by the CIRS company. Details of the physical dimensions can be found online (Computerized Imaging Reference Systems Inc. 2013). Both the homogenous and the anthropomorphic phantom are successively equipped with the 100 mm pencil chamber to measure conventional CTDI values and with five dose profilers for the simultaneous measurement of point doses.

Since both PMMA and anthropomorphic phantoms are used in this study, the material conversion file contains values for PMMA as well as for lung tissue, bone and water. The ranges for material conversion used in this work are given in table 2. The water equivalent material of the phantom represents, to some extent, the soft tissue. Hence, the range for the Hounsfield units of water is enlarged. In addition, the ImpactMC software requires the consecutive definitions of the material conversion; changing the minimum HU for water to a value of e.g. $-30$ would enlarge the range for lung tissue. Furthermore, materials related to the patient table such as carbon-fiber and aluminum are included in the conversion file as well. Since the composition of the carbon fiber is unknown, an equivalent material has been defined that is made from carbon ($\approx 92\%$ mass fraction), hydrogen ($\approx 5\%$) and
lead (≈3%). The fraction of lead has been calculated using the attenuation of the table (as measured with an ionization chamber), the geometry of the table from the DICOM file, the CT spectra and the mass attenuation coefficients of the materials mentioned above. In summary, an equivalent table material was calculated that mimics the attenuation properties of the table in order to achieve accurate simulation results.

2.4.2. Dose output measurements for normalization

In order to normalize the simulated values in absolute values of air kerma, the scanner output is used. To this end, the air kerma is measured free-in-air at the isocenter of the scanner using the ROC0.6 probe mounted at the front of the aluminum frame. A large collimation (±40 mm) ensures the full coverage of the detector and gives the peak air kerma of the beam \( K_a \). This value is used in ImpactMC for normalization and is inserted into the ‘AirKerma (mGy/100 mAs)’ field. Unfortunately, taking the average air kerma alone and assuming a rectangular beam profile with a nominal collimation are not sufficient to achieve correct results. Doing so would lead to an underestimation of the dose in the simulation, since Heel and penumbra effects as well as scattered radiation are not included. These effects deform the assumed rectangular shape; in particular, the penumbra effect, which is caused by a non-perfect collimation, enhances the deposited dose and needs to be taken into account.

Hence, additional measurements were made using a 100 mm pencil-type chamber to measure the free-in-air CTDI \( \text{CTDI}_{\text{free air}} \) for different collimations of interest in the isocenter. \( \text{CTDI}_{\text{free air}} \) is defined as the air kerma length product measured free-in-air at the isocenter of the CT and normalized to the nominal collimation. It includes the different effects, mentioned above, as well as scattered radiation from the BT filter. In order to realize the proper normalization from measuring \( \text{CTDI}_{\text{free air}} \), the property labeled as ‘total beam collimation’ in ImpactMC is set. For our purposes, the total beam collimation \( z_{\text{tot}} \) is calculated from the nominal beam width \( z_N \) using the ratio of \( \text{CTDI}_{\text{free air}} \) and peak air kerma \( K_a \):

\[
z_{\text{tot}} = z_N \cdot \frac{\text{CTDI}_{\text{free air}}}{K_a^P} = \frac{K_a L}{K_a^P}. \tag{8}
\]

The total beam width is typically larger than the nominal beam width. For example, using \( z_N = 10 \) mm and measured values for the peak dose and the \( \text{CTDI}_{\text{free air}} \) at 120 kV of \( K_a^P = 21.3 \) mGy/100 mAs and \( \text{CTDI}_{\text{free air}} = 28.26 \) mGy/100 mAs leads to \( z_{\text{tot}} = 13.3 \) mm (compare table 6).

For broader collimations (e.g. 160 mm), a longer, commercially available pencil-type chamber (300 mm) was used.

2.4.3. Uncertainty of simulated dose

The uncertainty of the dose simulation is estimated by varying single input parameters while keeping the rest of the parameters fixed. In the first attempt, it is assumed that the different parameters have negligible correlations. The summary of these studies is presented in table 3. The statistical uncertainty \( \sigma_{\text{Stat}} \) is defined by the number of photons \( (4 \times 10^9) \) and the number of voxels, which is \( 512 \times 512 \times 201 \) (voxel dimensions 0.98 × 0.98 × 2.5 mm\(^3\)) for all geometries and was estimated by repeating the same simulation multiple times at \( \sigma_{\text{Stat}} = 0.8\% \) for \( k = 1 \). Other effects such as the uncertainty in the total collimation \( z_{\text{tot}} \) and the normalization related to the air kerma using \( K_a^P \) lead to similar values below 1%. Except for the uncertainty in the simulation of the air kerma length product, the uncertainty from \( z_{\text{tot}} \) is slightly higher, which is expected due to the penumbra effects. The construction of the equivalent BT filter leads to a much higher contribution to the final uncertainty. From the multiple measurements performed on the Optima 660, the filters with minimum and maximum thickness are extracted and compared to the dose results performed with the average filter. The resulting uncertainty is calculated to be 2.5%, which is a conservative estimation. The major contributor in this example is related to the rotation angle \( \alpha \) of the x-ray tube. The starting angle of the exposure is an important input parameter, especially for helical scans with multiple rotations. Since the information of the angle is not given by the CT console nor by the header of the DICOM file, its uncertainty has been estimated by performing
multiple simulations with different starting angles between 0° and 270° and averaging the results. The standard deviation was calculated to be 0.9% for the axial mode and for one rotation in the first measurements. However, the standard deviation increases drastically to 7.5% when helical scan modes and multiple rotations are used. For the latter case, the combined uncertainty \( \sigma_c \) is calculated by means of a quadratic summation to \( \sigma_c = 8.0\% \) for point dose simulations and 8.1% for simulated values of the air kerma length product \( (k = 1) \). For the axial mode, the combined uncertainty is calculated to be \( \sigma_c = 2.9\% \) for point dose simulations and 3.1% \( (k = 1) \) for simulated values of the air kerma length product.

### 2.5. Timing and duration of the procedures

The duration of the overall procedure may be estimated roughly from the following time frames: set-up of the mobile equipment (shown in figure 2): 1 h, total time covering the measurement scans with each BT filter and tube voltage setting (followed by immediate source model calculations by automated software): 0.5–1 h, removal and replacement of the mobile equipment by verification phantom with real-time dosimeters: 1 h, phantom scan with dose measurements and final dismantlement of the system: 1 h. In another session, the measured equivalent source models, the geometry of the verification phantom and the scan parameters are applied in the ImpactMC (GPU version) simulation: 1 h (for details in computing time see Chen et al (2012)), and finally compared with measurements in verification analysis: 1 h. Combining these times together, the entire beam characterization for a new CT scanner model takes about one or two days. Note: these procedures need to be done only once per single CT scanner type for future patient specific dose estimations.

### 3. Measurements and results

#### 3.1. Determination of equivalent source models

##### 3.1.1. GE Optima CT 660

The average Al thicknesses \( d_{Al} \) from the determination of the total filtration, as described in section 2, are given in table 4, organized according to the different BT filters and tube voltages. In total, eight measurements have been taken to date for the Optima 660 using the mobile measurement setup. The standard deviation for the small filter varies between 1.3% and 2.7%, while the corresponding values for the large filter are slightly higher, ranging between 2.1% and 3.8%.

The reproducibility from our previous work Rosendahl and Büermann (2017) has been preserved, although the procedure has been changed to include a smaller collimation. The uncertainty of the fit has been estimated from an \( \chi^2 \)-analysis of the single measurements, with \( 1\sigma_{\chi^2} \)-uncertainty being defined as the increase of \( \chi^2 \) to \( \chi^2 + 1 \). From the measurements, the average uncertainty is estimated to be in the range of \( 1.3\% \leq \sigma_{\chi^2} \leq 2.8\% \) for the large and small filters, as it is of the same magnitude as the statistical deviation, as shown in table 4.

Using the total Al-filtering \( d_{Al} \), aluminum equivalent BT filter geometries for different tube voltages are constructed for both the small filter and the large filter. From the eight different measurements, the average has been calculated and is shown graphically in figure 5. For large fan angles, a voltage dependent filter thickness appears. This is related to the assumption of only one filter material. In reality, the filter might be constructed from materials other than aluminum, or even from combinations of materials. The energy-dependent photon cross-sections of these materials differ from that of Al, which leads to this spreading. The uncertainty in the determination...
of the total filtration directly affects the modeling of the equivalent filter; its influence is shown graphically in figure 6. The average large BT filter is plotted for 80 kV and 120 kV in comparison to filter geometries with the highest and lowest values of $d_{\text{Al}}$ obtained from the measurement. The comparison with the standard deviation in table 4 shows that the higher values for 80 kV are also visible in the spreading of the filter construction, while the deviation for 120 kV is much smaller.

3.1.2. Toshiba Aquilion ONE

For the determination of the equivalent source model, a collimation of 40 mm with a pitch of 0.637 is used. The total filtration has been determined under different conditions, and the results are summarized in table 5. In order to minimize the measurement time in the hospital, not all voltage steps are carried out for each filter. Instead, the Al attenuation measurements are repeated three times for 120 kV, leading to a standard deviation of $\approx 2\%$, which verifies the reproducibility. While the $\chi^2$-analysis of the fitting gives uncertainties of less than 5% in most cases, some single measurements show higher uncertainties of up to 8%. From the values, it is shown that the medium and large filters have the same total thickness at the center of the BT filter, while the small filter is thinner. Within the uncertainty, it is not possible to distinguish between the medium and large filters from the Al-equivalent thickness alone.

Since the large BT filter is used for the validation process in this work, we focus on the determination of the equivalent thickness of this filter while the results for the other filters are planned for publication elsewhere. In figure 7, the Al-equivalent geometry of the large filter is presented for different tube voltages (left). In contrast to the results from the Optima 660 shown in figure 5, the geometry does not split for the different tube voltages at large fan angles. Hence, it can be assumed that the filter is made from materials whose attenuation characteristics are similar to aluminum. Consequently, an average filter from the different tube voltages can be calculated.

Compared to the Optima 660, the filtration at the center of the BT is lower, while at the edges, the opposite is true. This should lead to higher values for air kerma on-axis at the isocenter. For the 80 kV filter geometry, some conspicuous peaks occur at high fan angles. These artifacts are related to the low signal intensity of 5 $\mu$Gy s$^{-1}$ to the high filtering at large fan angles. This effect has not been observed at the Optima 660 since the absolute thickness of the filter in the Aquilion ONE is higher at the edges compared to the Optima 660.

3.2. Validation of GE Optima CT 660 equivalent source models

For the validation, different simulations have been performed using equivalent source models obtained from the dynamic method and the TRIC method. In this work, the results for tube voltages of 80 kV and 120 kV at a

| Voltage (kV) | Small filter |  | Large filter |  |
|-------------|--------------|---------------|--------------|---------------|
|             | $d_{\text{Al}}$ (mm) | St. dev. (%) | $1\sigma_{\chi}$ (%) | $d_{\text{Al}}$ (mm) | St. dev. (%) | $1\sigma_{\chi}$ (%) |
| 80          | 7.464        | 2.7           | 2.6          | 10.30        | 3.8           | 2.8          |
| 100         | 7.376        | 2.2           | 1.9          | 10.26        | 2.1           | 2.2          |
| 120         | 7.398        | 1.7           | 1.4          | 10.26        | 2.6           | 1.5          |
| 140         | 7.346        | 1.3           | 1.3          | 10.15        | 2.6           | 1.3          |

Figure 5. Aluminum equivalent bow tie filters of an Optima 660. From the eight measurements with the mobile setup, the average aluminum equivalent BT filters for different tube voltages are calculated. The Optima 660 is equipped with a large and a small BT filter.

Table 4. Total aluminum filtration of the Optima 660. The values for the total filtration $d_{\text{Al}}$ are given for the different tube voltages and BT filters and are average values from eight single determinations using the mobile setup.
nominal collimation of 10 mm are presented for the Body PMMA CT phantom and the thorax phantom in order to provide a manageable number of values.

### 3.2.1. Dose output

For the proper normalization, the air kerma at the isocenter has been measured free-in-air as described above. The values for the \( CTDI_{\text{free air}} \) and the peak air kerma, \( K_P^a \), as well as the total beam width \( z_{\text{tot}} \) for the BT filter and tube voltages used in the Optima 660 are presented in table 6. The estimated uncertainty budget for air kerma measurements are compiled in table 1 for the different detectors. The relative uncertainty of \( z_{\text{tot}} \) is calculated as 1.2% for \( k = 1 \); its influence on the simulation results is investigated in the following.

### 3.2.2. Comparison of measured and simulated dose in the PMMA phantom

Air kerma measurements are performed inside the homogenous PMMA body phantom using the axial scan mode with only one rotation of 1 s and a current of 100 mA. Five calibrated dose profilers were used for the measurement. In table 7, the results from the measurement and from the simulation are compiled.
The mean average difference between the measured and the simulated doses is determined to be 3.6\% for 80 kV and 3.2\% for 120 kV with a maximum difference of 5.6\% and 8.0\%, respectively; the dynamic determination of equivalent source models show good agreement. The combined uncertainty for the measurement of the air kerma length product is measured using a 100 mm pencil-type ionization chamber and the RC0.6, respectively. The values are used to calculate the total beam width $z_{\text{tot}}$. The combined uncertainty $\sigma_c$ in the dose measurement and total beam width was calculated for a coverage factor of $k = 1$.

| Filter  | Voltage (kV) | $K_x^P$ (mGy/100 mAs) | $CTD_{\text{free air}}$ (mGy/100 mAs) | $z_{\text{tot}}$ (mm) | $CTD_{\text{free air}}$ (mGy/100 mAs) | $z_{\text{tot}}$ (mm) |
|---------|--------------|------------------------|--------------------------------------|-----------------------|--------------------------------------|-----------------------|
|         |              |                        | 10 mm                                | 40 mm                 | 10 mm                                | 40 mm                 |
|         |              | $\sigma_c = 0.7\%$     | $\sigma_c = 1.0\%$                   | $\sigma_c = 1.2\%$    | $\sigma_c = 1.0\%$                   | $\sigma_c = 1.2\%$    |
| Large   | 80           | 7.575                  | 10.10                                | 13.3                  | 8.496                                | 44.9                  |
|         | 120          | 21.30                  | 28.26                                | 13.3                  | 23.74                                | 44.5                  |

The mean average difference between the measured and the simulated doses is determined to be 3.6\% for 80 kV and 3.2\% for 120 kV with a maximum difference of 5.6\% and 8.0\%, respectively; the dynamic determination of equivalent source models show good agreement. The combined uncertainty for the measurement of the air kerma length product is measured using a 100 mm pencil-type ionization chamber and the RC0.6, respectively. The values are used to calculate the total beam width $z_{\text{tot}}$. The combined uncertainty $\sigma_c$ in the dose measurement and total beam width was calculated for a coverage factor of $k = 1$.

| Voltage (kV) | Position | $k_{Q,ph}$ | $\sigma_c = 1.3\%$ (k = 1) | $CTD_{\text{free air}}$ (mGy/100 mAs) | $\sigma_{\text{rep, max}}$ (%) | Difference (mGy/100 mAs) | TRIC method (mGy/100 mAs) | Difference (%) |
|-------------|----------|------------|----------------------------|--------------------------------------|-------------------------------|--------------------------|--------------------------|----------------|
| 80          | 12 o’c   | 0.9699     | 2.789                      | 2.632                                | -5.6                          | 2.783                    | -0.2                     |
|             | 3 o’c    | 0.9657     | 2.611                      | 2.588                                | -0.9                          | 2.754                    | 5.5                      |
|             | 6 o’c    | 0.9671     | 2.035                      | 1.953                                | -4.1                          | 2.172                    | 6.7                      |
|             | 9 o’c    | 0.9785     | 2.629                      | 2.708                                | 3.0                           | 2.804                    | 6.7                      |
|             | Center   | 0.9830     | 0.4129                     | 0.4303                               | 4.2                           | 0.4265                   | 3.3                      |
| 120         | 12 o’c   | 0.9384     | 8.178                      | 7.997                                | -2.2                          | 8.326                    | 1.8                      |
|             | 3 o’c    | 0.9352     | 7.682                      | 7.895                                | 2.8                           | 8.094                    | 5.4                      |
|             | 6 o’c    | 0.9369     | 6.193                      | 6.246                                | 0.9                           | 6.437                    | 3.9                      |
|             | 9 o’c    | 0.9445     | 7.855                      | 8.025                                | 2.2                           | 8.318                    | 5.9                      |
|             | Center   | 0.9131     | 1.511                      | 1.632                                | 8.0                           | 1.593                    | 5.4                      |

### 3.2.3. Comparison of measured and simulated dose in the anthropomorphic phantom

In the next step, point dose measurements with semiconductor sensors are performed in an anthropomorphic phantom using the ‘Thorax-Medium Adult’ phantom. The scan parameters are set to 100 mA, one rotation and a rotation time of 1 s, which are the same as in previous measurements. The material conversion information for lungs, bones and water from the ImpactMC manual (CT Imaging GmbH 2014) were used, which yield the simulated dose values compiled in table 9. In this example, the simulated doses are larger than those measured using the RTI dose profilers. For the 80 kV data, the average difference observed is 1.9\%, with a maximum
difference of 3.7%. For the 120 kV measurement, the average difference is slightly higher, being 3.9% with a maximum difference of 5.8% in the central position. For the air kerma measurements, a combined uncertainty has been estimated to be $\sigma_{k} = 2.0\%$ for a coverage factor ($k = 1$), using the maximum deviation observed from all positions and voltages after three repeated measurements of $\sigma_{rep,\ max} = 1.7\%$.

### 3.2.4. Routine chest protocol applied to the anthropomorphic phantom

In order to validate the concept for scan parameters that are closer to those used in clinical practice, the thorax phantom is scanned with the ‘routine chest 0.6 s 5 mm SMARTmA’ GE protocol. This helical scan has a pitch of 1.375 and a source rotation time of 0.6 s. The voltage is set to 120 kV and the large BT filter is used. Although this scan protocol normally uses a collimation of 40 mm, a smaller collimation of 20 mm is chosen due to the shortness of the phantom. The dose measurement is improved and additionally the uncertainties from the edges of the phantom are reduced. Furthermore, the ‘SMARTmA’ current modulation technique, which adjusts the current to the patient, was switched off to simplify the simulation. The current was has been fixed to 300 mA for the whole scan range of 6 cm, resulting in a measurement time of 2.01 s. In figure 8, the ImpactMC user display for the 3D dose simulation is shown.

Although only 6 cm of the phantom are reserved for projection, a total scan length of $\approx 9.2$ cm is calculated from the $\text{CTDI}_{\text{vol}}$ and DLP values given by the system (see table 10). The difference is caused by the well-known over-beaming effect related to the spiral scan modality. Hence, to achieve good dose results, the complete geometry of the phantom is needed for the simulation. Otherwise, the simulated dose will be drastically underestimated in the way it is illustrated in figure 8, where the dose maps (b) and (c) are compared. For the studies in this work, the whole phantom and additional space before and after were scanned to address possible scattering from the table. In reality, because only a part of the patient is scanned, the border areas need to be modeled. The results from the simulation and from the measurement of the air kerma length product are compiled in table 10.

![Image](https://example.com/image.png)

**Figure 8.** The ImpactMC user display for the 3D dose simulation is shown.

The combined uncertainty of the measurement has been determined to be $\sigma_{k} = 2.7\%$ and is dominated by the statistical uncertainty that occurs during the measurement of $\sigma_{rep} = 2.5\%$. The measured and simulated values agree within the uncertainties of the measurements and the simulation. The average difference is calculated to be 2.8% with a maximum difference of 5%. For this simulation, the starting angle, which is unknown, has been varied between $0^\circ$ and $270^\circ$ and the average of these simulations is compiled in table 10. This reduces the uncertainty in the starting angle for helical scans from 7.5% down to 3.9% for a coverage factor $k = 1$. In summary, the test validates the notion that more realistic scan settings can also be simulated with small differences between the measurement and the simulation.

### 3.3. Validation of Toshiba Aquilion ONE equivalent source models

#### 3.3.1. Dose output

For the Aquilion ONE, the air kerma $K_{p}^{a}$ and the $\text{CTDI}_{\text{free air}}$ were measured with the RC0.6 chamber and with a 300 mm pencil-type chamber, respectively. In table 11, the results from the measurements and the corresponding values for the total beam collimation are compiled. The absolute values for $K_{p}^{a}$ and for $\text{CTDI}_{\text{free air}}$ are higher than for the GE Optima CT under comparable conditions, which is expected due to the lower total filtration from the equivalent source model.

| Voltage (kV) | Position | Measured $K_{p}^{a}$ | Dynamic method | Difference (%) | TRIC method | Difference (%) |
|-------------|----------|----------------------|-----------------|---------------|-------------|---------------|
| 120 6 o’c  | 1.343 | 12.75 | -5.1 | 12.59 | -6.3 |
| 3 o’c      | 1.233 | 12.67 | -4.4 | 12.43 | -6.1 |
| 80 6 o’c   | 11.89 | 10.99 | -7.5 | 11.34 | -4.6 |
| 9 o’c      | 13.26 | 12.64 | -4.7 | 12.45 | -6.1 |
| Center     | 6.425 | 6.876 | 7.0 | 6.565 | 2.2 |

**Table 8.** Results of dose measurements with 100 mm pencil-type chamber and simulations performed on the Optima 660. The air kerma length product $K_{p}^{a}L$ at the five positions of the PMMA 32 cm phantom are measured with a 100 mm CT pencil-type chamber (PTW) and compared to simulated values from ImpactMC. The scan parameters chosen are one rotation with a current of 100 mA, a rotation time of 1 s and a nominal collimation of 10 mm.
3.3.2. Comparison of measured and calculated dose in the anthropomorphic phantom

For the validation of the equivalent source model, the measured air kerma length product is compared to a simulation in an anthropomorphic phantom. In table 12, the results for one tube rotation at 120 kV tube voltage are shown. It has been observed that, after repeating the measurement three times, the values of measured air kerma length product using the pencil-type ionization chamber can vary up to 8.5%, leading to a very high combined uncertainty of $\sigma_c \approx 8.6\%$ ($k = 1$) in the dose measurement. The reason for the fluctuation is related to the rotation angle of the tube when starting the irradiation, which varies over time. The effect is much more pronounced for the Toshiba Aquilion ONE than for the GE Optima 660. This fact may be due to the construction of the CT.

**Table 9.** Results of point dose measurements and simulations in an anthropomorphic phantom performed on the Optima 660. The dose values at the five positions of the ‘Thorax-Medium Adult’ anthropomorphic phantom are measured with semiconductor detectors (RTI) and compared to simulated values from ImpactMC. The scan parameters chosen are one rotation with 100 mA, a rotation time of 1 s and a nominal collimation of 10 mm.

| Voltage (kV) | Position | Measured dose (mGy/100 mAs) | Dynamic method (mGy/100 mAs) | Difference (%) |
|-------------|----------|-----------------------------|------------------------------|----------------|
| 80          | 12 o’c   | 2.889                       | 2.813                        | −2.6           |
|             | 3 o’c    | 2.437                       | 2.426                        | −0.4           |
|             | 6 o’c    | 2.204                       | 2.254                        | 2.3            |
|             | 9 o’c    | 2.426                       | 2.335                        | −3.7           |
|             | Center   | 0.7085                      | 0.7067                       | −0.2           |
| 120         | 12 o’c   | 8.451                       | 8.791                        | 4.0            |
|             | 3 o’c    | 7.169                       | 7.394                        | 3.1            |
|             | 6 o’c    | 6.627                       | 6.963                        | 5.1            |
|             | 9 o’c    | 7.407                       | 7.291                        | −1.6           |
|             | Center   | 2.545                       | 2.692                        | 5.8            |

**Figure 8.** Dose simulation of a routine chest protocol (GE). A standard patient protocol provided by General Electric, ‘routine chest 0.6 s 5 mm SMARTmA’, is applied to the ‘Thorax-Medium Adult’ anthropomorphic phantom and the dose simulation is performed using the equivalent source model of the Optima 660. The 3D dose distribution is shown for full geometry of the phantom (a) + (b) and for the 6 cm scan length (c), which has been defined as the scan area. The images in (b) + (c) show the projection in z-direction (‘Coronal’ window of ImpactMC).

**Table 10.** Results of measurements of the air kerma length product and simulations in an anthropomorphic phantom using routine chest protocol. The air kerma length product $K_a L$ at the five positions of the ‘Thorax-Medium Adult’ anthropomorphic phantom are measured with a pencil-type ionization chamber (100 mm) and are compared to simulated values from ImpactMC. A routine chest protocol with a 5 mm slice thickness and 120 kV in the Optima 660 is used.

| Voltage (kV) | Position | $K_a L$ (mGy · cm) | $\sigma_c = 3.3\%$ | Dynamic method (mGy · cm) | Differences (%) |
|-------------|----------|--------------------|---------------------|--------------------------|----------------|
| 120         | 12 o’c   | 147.2              | 150.4               | 2.2                      |
|             | 3 o’c    | 116.2              | 118.4               | 1.9                      |
|             | 6 o’c    | 125.1              | 129.5               | 3.5                      |
|             | 9 o’c    | 119.0              | 120.5               | 1.3                      |
|             | Center   | 84.72              | 88.97               | 5.0                      |

3.3.2. Comparison of measured and calculated dose in the anthropomorphic phantom

For the validation of the equivalent source model, the measured air kerma length product is compared to a simulation in an anthropomorphic phantom. In table 12, the results for one tube rotation at 120 kV tube voltage are shown. It has been observed that, after repeating the measurement three times, the values of measured air kerma length product using the pencil-type ionization chamber can vary up to 8.5%, leading to a very high combined uncertainty of $\sigma_c \approx 8.6\%$ ($k = 1$) in the dose measurement. The reason for the fluctuation is related to the rotation angle of the tube when starting the irradiation, which varies over time. The effect is much more pronounced for the Toshiba Aquilion ONE than for the GE Optima 660. This fact may be due to the construction of the CT.
The average difference between the measurement and the simulation has been determined to be 3.8% with the maximum difference of 9.2%. The overall results are in agreement with the studies performed on the Optima 660, indicating a slight overestimation of the air kerma in the simulations for the 6 o’clock and central positions of the phantom.

Finally, more realistic scan settings are presented. The anthropomorphic phantom is scanned using a spiral scan mode with a pitch of 0.813 and a nominal collimation of 40 mm, covering a scan range of 8 cm around the center of the phantom. The tube voltage was set to 120 kV with a constant current of 100 mA without the tube current modulation. The air kerma length product measured using the 100 mm pencil-type ionization chamber (PTW) and compared to simulated values from ImpactMC. The scan parameters have been set to one rotation with 150 mA a rotation time of 1 s and a nominal collimation of 12 mm.

The results of the measurement and the simulation show differences of less than 8% with an average difference of 3.4% over all positions. During this measurement, the uncertainty from repetition (three times) was very small, of the order of $\sigma_{\text{Rep}} \approx 1.0\%$ ($k = 1$). Hence, the combined uncertainty for the dose measurement is calculated to be $\sigma_{\text{c, meas}} = 1.4\%$ for coverage factor ($k = 1$) using the Type B uncertainties from table 1. The reason for the large difference in the statistical uncertainty in the tube output between measurements with one rotation and measurements with several rotations is unknown.

### 4. Discussion of the results

In summary, the measured and simulated values for the Optima 660 in both standard CTDI and anthropomorphic phantoms using equivalent source models from a dynamic determination agree within a few percent, with only a few examples of larger differences. In different measurements, it was observed that the simulated dose in the central position was overestimated in the simulation, which indicates that the value for total filtration $d_{\text{tot}}$ is slightly overestimated, thus causing a harder spectrum. Additional measurements done in this work by using static methods with lead shielding of scattered radiation from the x-ray tube confirm this assumption. Alternatively, the phantom and the detector positions relative to the beam and non-perfect collimation may have an influence. However, this has not been exhaustively investigated and other measurements have not shown this effect. Here, additional studies are needed. The maximum difference observed was slightly above 10%. The majority of observables are in agreement with the estimated uncertainties of the dose measurement and the simulation. Although material assignment becomes more complicated with anthropomorphic phantoms, it was demonstrated that, for both simple scan parameters and more advanced scan protocols, equivalent source models and the MC simulation are well suited for verification of personalized CT dosimetry techniques. This was confirmed after using the technique at a different CT in a clinical environment at the Städtisches Klinikum.

### Table 11. CT dose output of the Toshiba Aquilion ONE. The values for the $\text{CTDI}_{\text{free air}}$ and the average air kerma $K_a^c$ are measured using a 300 mm pencil-type chamber and the RC0.6, respectively. The values are used to calculate the total beam width $z_{\text{list}}$. The combined uncertainty $\sigma_c$ in the dose measurement and total beam width is calculated using the GUM workbench for a coverage factor $k = 1$.

| Voltage (kV) | $K_a^c$ (mGy/100 mAs) | $\text{CTDI}_{\text{free air}}$ (mGy/100 mAs) | $z_{\text{list}}$ (mm) |
|-------------|------------------------|---------------------------------|------------------|
| 80          | 9.670                  | 12.85                           | 15.94            |
| 120         | 25.02                  | 32.60                           | 15.63            |

### Table 12. Results of measurements and simulations of the air kerma length product in anthropomorphic phantom at the Toshiba Aquilion ONE. The air kerma at the five positions of the ‘Thorax-Medium Adult’ anthropomorphic phantom are measured with a 100 mm pencil-type ionization chamber (PTW) and compared to simulated values from ImpactMC. The scan parameters have been set to one rotation with 150 mA a rotation time of 1 s and a nominal collimation of 12 mm.

| Voltage (kV) | Position | Measured $K_aL$ (mGy · cm/100 mAs) | Simulated $K_aL$ (mGy · cm/100 mAs) | Differences (%) |
|-------------|----------|----------------------------------|----------------------------------|-----------------|
| 120         | 12 o’c   | 18.23                            | 18.21                            | −0.2            |
|             | 3 o’c    | 14.29                            | 14.54                            | 1.7             |
|             | 6 o’c    | 14.71                            | 15.60                            | 6.1             |
|             | 9 o’c    | 14.42                            | 14.65                            | 1.6             |
|             | Center   | 10.66                            | 11.63                            | 9.2             |
Braunschweig. The equivalent source model of the Toshiba Aquilion ONE has been derived from measurements with the mobile measurement setup; the shape of the BT filters differs significantly from the BT of the Optima 660. The validation measurements of one rotation in axial mode yield differences between the simulation and the measurement of less than 10%. The same results were obtained for corresponding additional measurements done in this work by using multiple rotations in spiral mode. The major difference between the two scanners was related to large fluctuations of 8.5% in the dose measurement when repeating single measurements. Since the different detector systems have been extensively tested and calibrated, the uncertainty of the air kerma measurement has been shown to be lower than 2% ($k = 1$) for the different detector systems. Hence, the fluctuation is very likely caused by the CT itself. The fluctuation of the tube output is evidently a disadvantage for any scan-specific or patient-specific dosimetry approach and leads to very high uncertainties. Additional studies are needed to investigate this effect and whether other clinical CT machines have comparable characteristics.

The major sources of uncertainty for the simulation are related to the shape of the BT filter and the rotation angle of the x-ray tube when starting the irradiation. The lack of knowledge of the starting angle was identified to be one of the crucial challenges in order to achieve accurate simulation results. To overcome this problem, several simulations with different starting angles were performed. The results were averaged, which reduced the uncertainty. The effect of the geometry of the BT filter has been investigated by modifying the filter thickness with respect to the estimated uncertainties from the measurement of the total filtering. This effect is estimated to be less than 3%, which is the same size as the uncertainty of the total filtration.

To finalize the complete procedure of personalized dosimetry, techniques for organ segmentation need to be developed and implemented as well as the correction for missing scattered radiation due to the shortness of the phantom. Additional verifications of the procedure are necessary using different CTs from other manufacturers (e.g. Siemens) and more advanced phantoms such as CIRS: Model 600 ‘3D Sectional Torso Phantom’, as well as patient scan data for the final test. This will allow more realistic geometries and materials to be incorporated.

5. Conclusion

Viable procedures were developed that allow the CT scan dose to be measured and calculated at five positions inside physical anthropomorphic phantoms. The relative standard uncertainties of the dose measurements were in the range 1% to 2% and those of the calculations between 3% for axial and 8% for helical scans. In general, measured and calculated dose values inside the phantoms were within the range of uncertainties of 5% to 10%. The procedures are applicable to any scanner type under clinical conditions. Results show that these are well suited for verifying CT x-ray source models needed for personalized CT dosimetry based on post-scan Monte Carlo calculations. The procedures could be part of a potential acceptance test if personalized CT dosimetry is integrated in future CT scanners.

Acknowledgments

The authors thank Reinulf Böttcher, Hartmut Drehsler, Marvin Wenzel and Lars Herrman (PTB) for supporting the measurements and mechanical construction of the mobile measurement equipment. This project has received funding from the EMPIR programme 15HLT05, ‘Metrology for multi-modality imaging of impaired tissue perfusion’. The EMPIR initiative is co-funded by the European Union’s Horizon 2020 research and innovation programme and the EMPIR Participating States.
References

Alikhani B and Büermann L 2016 Non-invasive experimental determination of a CT source model Phys. Med. 32 59–66

Bazalova M and Verhaegen F 1988 Monte Carlo simulation of a computed tomography x-ray tube Phys. Med. Biol. 32 5945–55

Boone J M 2007 The trouble with CTDI Med. Phys. 34 1364–71

Boone J M 2010 Method for evaluating bow-tie filter angle dependent attenuation in CT: theory and simulation results Med. Phys. 37 40–8

Boone J M et al 2011 Size-specific dose estimates (SSDE) in pediatric and adult body CT examinations AAPM Task Group Report No. 204 (College Park, MD: AAPM) (https://www.aapm.org/pubs/reports/rpt_204.pdf)

Brady S L et al 2015 How to appropriately calculate effective dose for CT using either size-specific dose estimates or dose-length product AJR 204 953–8

Bureau International des Poids et Mesures, BIPM 2008 Evaluation of measurement data guide to the expression of uncertainty in measurement JCGM 100 2008 (https://www.bipm.org/utils/common/documents/jcgm/JCGM_100_2008_E.pdf)

Chen W et al 2012 Fast on-site Monte Carlo tool for dose calculations in CT applications Med. Phys. 39 2985–96

Computerized Imaging Reference Systems Inc. 2013 Tissue equivalent CT dose phantoms—data sheet http://cirscinc.com/products/all/17/tissue-equivalent-ct-dose-phantoms (Accessed: 06 March 2018)

CT Imaging GmbH 2014 ImpactMC—user guide document version 1.17

Deak P et al 2008 Validation of a Monte Carlo tool for patient-specific dose simulations in multi-slice computed tomography Eur. Radiol. 18 759–72

Deak P et al 2010 Multisection CT protocols: sex- and age-specific conversion factors used to determine effective dose from dose-length product Radiology 257 158–68

Dixon R L and Boone J M 2010 Cone beam CT dosimetry: a unified and self-consistent approach including all scan modalities—with or without phantom motion Med. Phys. 37 2703–18

Dixon R L and Boone J M 2013 Dose equations for tube current modulation in CT scanning and the interpretation of the associated CTDIvol Med. Phys. 40 111920

Dixon R L and Boone J M 2014 Stationary table CT dosimetry and anomalous scanner-reported values of CTDIvol Med. Phys. 41 011907

Duisterwinkel H A et al 2015 Spectra of clinical CT scanners using a portable Compton spectrometer Med. Phys. 42 1884–94

European Commission (EC) 2014 Medical Radiation Exposure of the European Population Radiation Protection Report No. 180 (Part 1/2) (Brussels, Belgium: European Commission) p66

Ginat D T and Gupta R 2014 Advances in computed tomography imaging technology Ann. Rev. Biomed. Eng. 16 431–53

IEC 60601-2-44:2009 x and Gamma Reference Radiation for Calibrating Dosimeters and Doserate Meters and for Determining their Response as a Function of Photon Energy—Part 1: Radiation Characteristics and Production Methods (Geneva, Switzerland: International Organisation for Standardization)

International Organization for Standardization 1997 International Standard ISO 4037-1 Radiation Protection and Production Methods (Geneva, Switzerland: International Organisation for Standardization)

Kalverda A W 2014 Dose in x-ray computed tomography Phys. Med. Biol. 59 R129–50

Kruger R L, McCollough C H and Zink F E 2000 Measurement of half-value layer in x-ray CT: a comparison of two non-invasive techniques Med. Phys. 27 1915–9

Matscheko G and Carlsson G A 1988 Measurement of absolute energy spectra from a clinical CT machine under working conditions using a Compton spectrometer Phys. Med. Biol. 34 209–22

McCollough C H, Bruesewitz M R and Koffler J M 2006 CT dose reduction and dose management tools: overview of available options RadioGraphics 26 503–12

McCollough C H et al 2011 CT dose index and patient dose: they are not the same thing Med. Phys. 38 1806–15

McNab M and Poludniowski G 2014 A method to generate equivalent energy spectra and filtration models based on measurement for multidetector CT Med. Phys. 41 3633–40

McNabb M and Poludniowski G 2015 A rapid noninvasive characterization of CT x-ray sources Med. Phys. 42 3960–8

Myronakis M et al 2009 Evaluation of a patient-specific Monte Carlo software for CT dosimetry Radiat. Prot. Dosimetr. 133 248–55

NCRP 2009 Ionizing radiation exposure of the population of the United States NCRP Report 160 (Bethesda, MD: National Council on Radiation Protection and Measurements)

Poludniowski G and Evans P M 2007a Calculation of x-ray spectra emerging from an x-ray tube. Part I. Electron penetration characteristics in x-ray targets Med. Phys. 34 2164–74

Poludniowski G and Evans P M 2007b Calculation of x-ray spectra emerging from an x-ray tube. Part II. X-ray production and filtration in x-ray targets Med. Phys. 34 2175–86

Poludniowski G et al 2009 SpekCalc: a program to calculate photon spectra from tungsten anode x-ray tubes Phys. Med. Biol. 54 433–38

Randazzo M and Tambasco M 2015 A rapid noninvasive characterization of CT x-ray sources Med. Phys. 42 1884–94

Rosendahl S and Büermann L 2017 Dynamic determination of equivalent CT source models for personalized dosimetry J. Med. Imaging 4 043005

Sahiner B, Pezetlik A, Hadijski I. M, Wang X, Drukker K, Cha K H and Giger M L 2019 Deep learning in medical imaging and radiation therapy Med. Phys. 46 e1–36

Schmidt B and Kalender W A 2002 A fast voxel-based Monte Carlo method for scanner- and patient-specific dose calculation in computed tomography Phys. Med. 18 45–53

Shope T B, Gagne R M and Johnson G C 1981 A method for describing the dose delivered by transmission x-ray computed tomography Med. Phys. 8 488–95

Siiskonen T and Suutari J 2018 Calibration and use of a pencil-type ct chamber: effect of the PMMA body phantom Radiat. Prot. Dosim. 178 272–5

Váňo E et al 2017 ICRP publication 135: diagnostic reference levels in medical imaging Ann. ICRP 46 1–144

Weston A D et al 2018 Automated abdominopelvic segmentation of CT scans for body composition analysis using deep learning Radiology 290 669–79

Turner A C et al 2009 A method to generate equivalent energy spectra and filtration models based on measurement for multidetector CT Monte Carlo dosimetry simulations Med. Phys. 36 2154–64