Estimation of patient organ doses from CT examinations in Tanzania

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Although the use of CT in medical diagnosis delivers radiation doses to patients that are higher than those from other radiological procedures, lack of optimized protocols could be an additional source of increased dose in developing countries. The aims of this study are, first, to determine the magnitude of radiation doses received by selected radiosensitive organs of patients undergoing CT examinations and compare them with other studies, and second, to assess how CT scanning protocols in practice affect patient organ doses. In order to achieve these objectives, patient organ doses from five common CT examinations were obtained from eight hospitals in Tanzania. The patient organ doses were estimated using measurements of CT dose indexes (CTDI), exposure-related parameters, and the ImPACT spreadsheet based on NRPB conversion factors. A large variation of mean organ doses among hospitals was observed for similar CT examinations. These variations largely originated from different CT scanning protocols used in different hospitals and scanner type. The mean organ doses in this study for the eye lens (for head), thyroid (for chest), breast (for chest), stomach (for abdomen), and ovary (for pelvis) were 63.9 mGy, 12.3 mGy, 26.1 mGy, 35.6 mGy, and 24.0 mGy, respectively. These values were mostly comparable to and slightly higher than the values of organ doses reported from the literature for the United Kingdom, Japan, Germany, Norway, and the Netherlands. It was concluded that patient organ doses could be substantially minimized through careful selection of scanning parameters based on clinical indications of study, patient size, and body region being examined. Additional dose reduction to superficial organs would require the use of shielding materials.

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I. INTRODUCTION

It is well known that patient doses from CT procedures are relatively higher than doses from other imaging modalities based on ionizing radiation. For example, one CT examination of the chest delivers about 400 times the dose delivered by a conventional chest X-ray examination.\textsuperscript{(1,2)} Therefore, although CT represents only 5\% of the total number of medical X-ray procedures worldwide, this high dose procedure contributes about 34\% of the annual collective dose from all medical X-ray examinations to the population.\textsuperscript{(1)} This contribution is inevitable because it results from a combination of high dose per examination and frequent use of CT examination in diagnosis.\textsuperscript{(1–3)} This trend has started to appear in Tanzania, where use of CT for medical examination started as early as 1996. Currently, there are about 12 CT scanners in the country, and nearly 10 000 X-ray CT examinations are now performed annually. This constitutes about 1\% of all medical X-ray procedures performed in Tanzania.\textsuperscript{(1,4)}
Increased use of this high dose procedure has been of great concern globally because of the high possibility of inducing undesired health effects, such as induction of cancer, in patients. Of prime concern is the significant radiation dose delivered to superficial radiosensitive organs such as the eye lens, breast, and thyroid, which are, unfortunately, irradiated during radiological procedures of the head, chest, and cervical spine. The implication of some of these exposures, for example, to the breast and eyes, is the potential increase in the risk of breast cancer and cataract formation in the population. Since radiation exposure of different organs leads to different health effects, it has been of interest in this work to determine the actual doses delivered to individual organs.

The most useful way to assess organ doses is either by direct measurement (on patients using thermoluminescent dosimeters (TLDs) or on phantom using either an ionization chamber or TLDs) or by indirect measurement through measurement of CT dose indexes (CTDI) and published conventional factors obtained from Monte Carlo simulation and mathematical phantoms. Since dose measurement using TLDs is laborious and time-consuming for a wide survey, the patient organ doses in this study were determined using measurements of CTDI and published conversion factors. With the knowledge of organ doses it would be easy to identify the organs at greatest radiation risk, which requires immediate protective measures. In addition, countries with insufficient experience in the use of this modality may have additional sources of increased dose to patients, such as lack of optimized techniques. The aims of this study, therefore, are to assess the magnitude of organ dose imparted to patients undergoing CT examinations in Tanzania and to compare them with other studies from developed countries, and to assess the effect of CT scanning protocols on patient doses.

II. MATERIALS AND METHODS

A. Data collection

The data used in this study were collected from eight hospitals in Tanzania with CT scanners: Muhimbili National Hospital (MNH), Kilimanjaro Christian Medical Centre (KCMC), Bugando Medical Centre (BMC), the Agha Khan Hospital (AGKH), TMJ Hospital (TMJH), the Mission Mikocheni Hospital (MMH), Regency Medical Centre (RMC), and Shree Hindu Mandal Hospital (SHMH). Detailed specifications of the scanner used at each of these hospitals are contained in Table 1. Patient data collected from these centers were further classified in two categories. In the first, data were collected to study the effects of patient-related parameters (e.g., age, sex, diagnostic purpose of examination, body region, and use of contrast media) on organ dose. In the second, data were collected to investigate the effect of exposure-related parameters (gantry tilt, kilovoltage (kV), tube current (mA), exposure time, slice thickness, table increment, number of slices, and start and end positions of scans) on organ dose. The selected investigations used in this study represented over 90% of the total CT examinations conducted in Tanzania today. The collection of patient exposure parameters was done using patient dose survey forms prepared for collection of patient exposure-related parameters, hospital, and scanner used in each hospital. This form also includes a diagram of the human skeleton, on which the reported upper and lower extent of each scan was marked. A minimum number of 10 patients were used for each selected CT examination per hospital. The sum of 191, 74, 84, 88, and 54 (for 6 hospitals only) CT examinations of head, chest, abdomen, lumbar spine, and pelvis, respectively, was collected, making a total collection of 491 (about 500) examinations.
Table 1. Specifications of CT scanners used at each hospital in Tanzania

| Hospital | Manufacturer                  | Scanner model/ Scan mode                                   | Focal axial distance (FAD)/ Detector type |
|----------|-------------------------------|-----------------------------------------------------------|------------------------------------------|
| MNH      | Philips Medical Systems       | Tomoscan SR 4000 single slice, axial and helical modes    | FAD: 606 mm                              |
| KCMC     | Philips Medical Systems       | Tomoscan SR 4000 single slice, axial and helical modes    | FAD: 606 mm                              |
| RMC      | Philips Medical Systems       | Tomoscan M-EG single slice, axial and helical modes       | FAD: 480 mm detector: solid state        |
| BMC      | Philips Medical Systems       | Tomoscan M-EG single slice, axial and helical modes       | FAD: 480 mm detector: solid state        |
| TMJH     | Siemens Medical Systems       | Somatom Plus 4 single slice, axial and helical modes      | FAD: 570 mm detector: xenon gas          |
| MMH      | Siemens Medical Systems       | Somatom AR.Star single slice, axial and helical modes     | FAD: 510 mm detector: xenon gas          |
| AGKH     | GE Medical Systems            | CT/e single slice, axial and helical modes                 | FAD: 525 mm detector: solid state        |
| SHMH     | GE Medical Systems            | CT Max 640 single slice, axial mode only                   | FAD: 525 mm detector: solid state        |

B. CT dose measurements

The patient dose estimation from CT examination using the Monte Carlo technique requires measurements of CTDI and conversion coefficient data packages.\(^{9–11}\) In theory, the CTDI, which is a measure of the dose from single-slice irradiation, is defined as the integral along a line parallel to the axis of rotation (z) of the dose profile, \(D(z)\), divided by the nominal slice thickness, \(t\).\(^{9,11–13}\) In this study, CTDI was obtained from a measurement of dose, \(D(z)\), along the z-axis made in air using a special pencil-shaped ionization chamber (Diados, type M30009, PTW-Freiburg) connected to an electrometer (Diados, type 11003, PTW-Freiburg). The calibration of the ion chamber is traceable to the standards of the German National Laboratory and was calibrated according to the International Electrical Commission standards.\(^{14}\) The overall accuracy of ionization chamber measurements was estimated to be ±5%. Measurements of CTDI in air (CTDI\(_{100, \text{air}}\)) were made as recommended by the EUR 16262EN based on each
combination of typical scanning parameters obtained from each hospital.\(^{(11)}\) Unfortunately, the CTDI\(_{100,\text{air}}\) for CT scanner model Tomoscan M-EG from RMC were not determined due to a malfunction of the scanner. Hence, the required organ doses for this particular hospital were estimated using normalized CTDI values published by the ImPACT group.\(^{(15)}\) For the sake of simplicity, the CTDI\(_{100,\text{air}}\) will henceforth be abbreviated as CTDI\(_{\text{air}}\).

C. Organ dose determinations

Doses to selected organs (e.g., eye lens, thyroid, red bone marrow, breast, lung, and ovaries) were determined using the CTDI\(_{\text{air}}\) described in the previous section and organ dose conversion coefficients.\(^{(16)}\) Since the organ dose data are expressed in terms of absorbed dose to tissue, the measured values of dose in air (CTDI\(_{\text{air}}\)) were converted to dose to tissue (CTDI\(_{\text{tissue}}\)) using the ratio of the mass–energy absorption coefficients \(\left(\frac{\mu_{\text{en}}}{\rho}\right)\) of tissue to air:\(^{(9,10,12)}\)

\[
\text{CTDI}_{\text{tissue}} = \left[\frac{\left(\frac{\mu_{\text{en}}}{\rho}\right)_{\text{tissue}}}{\left(\frac{\mu_{\text{en}}}{\rho}\right)_{\text{air}}}\right] \times \text{CTDI}_{\text{air}}. 
\]

Despite the fact that the mass–energy absorption coefficient depends significantly on photon energy, the ratio of the mass–energy absorption coefficient of soft tissue to air is assumed to be constant for all typical X-ray spectra produced by the CT scanners examined with a value of 1.06 (with an error of no more than ±1\%).\(^{(10,12,15)}\) By using the scanner-specific organ dose conversion coefficient, the typical average organ dose, \(D_{\text{org,T}}\), for individual examination can be estimated by summation of the following form:\(^{(6,9,13)}\)

\[
D_{\text{org,T}} = \text{CTDI}_{\text{tissue}} \sum_{z_1}^{z_2} f(\text{organ}, z). 
\]

where \(z_1\) and \(z_2\) are the start and end positions of the scanned region, respectively. The organ dose conversion factor \(f(\text{organ}, z)\) was obtained from the NRPB datasets (NRPB-SR250) based on the Monte Carlo simulations.\(^{(6,16,17)}\) The CTDOSE software supplied by the ImPACT group\(^{(15)}\) was used to implement Eq. (2). The CTDI\(_{\text{air}}\) normalized to 100 mAs \(\left(\text{CTDI}_{\text{air}}\right)\), CT scanner manufacturer and model, and typical scanning parameters such as kV, mA, exposure time, pitch, slice thickness, gender, and start and end positions of each scan were used as input data to the CTDOSE spreadsheet in organ dose estimations.\(^{(10,15,16)}\)

Due to the fact that the software does not take into account the patient size, that is, the software was not discriminate between tall and short patients, it was necessary to adjust the scan region indicated on the human skeleton from each patient survey form in NRPB’s mathematical phantom (Figs. 1 and 2 in NRPB-R250)\(^{(10,15)}\) for each individual examination. This was done by first marking the start and end positions of the scan region and then determining the scan length from the number of slices, the slice thickness, and the table increment. This information was used in the selection of the part of the phantom irradiated in order to improve the correspondence between the organs irradiated in the patient and the phantom.\(^{(10)}\) Since the scanners used in this study (Table 1) were not in use at the time of the NRPB survey, the estimation of organ dose has to rely on the attributes of the new model compared to that of older designs. This was done using scanner-matching data published by the ImPACT group, and may lead to uncertainty of not more than 15\% of organ dose measurements.\(^{(13,15)}\) The overall uncertainties associated with estimation of organ doses, calculated by quadratic propagation of errors, were estimated to range from ±17\% to ±24\%. These uncertainties arose mainly from patient variations, uncertainty in the start and end positions of the scan region, calibration of the ionization chamber used to measure CTDI\(_{\text{air}}\), statistical uncertainties in normalized organ doses, and scanner matching data. In order to evaluate how well the hospitals in Tanzania are performing in terms of minimization of organ doses associated with CT imaging, it was
useful to compare mean organ dose per examination for all the hospitals that participated in the study. This was done by finding the mean organ dose from the typical patient organ dose weighted by the number of scans performed per given examination for each hospital. On the other hand, the mean value of the typical patient organ doses weighted by the number of scans per given examination, based on all hospitals in the survey, was taken as the country mean organ dose. The country mean organ doses were determined in order to compare with other studies from developed countries.

D. Data analysis
A summary of organ dose, \( D_{\text{org}, T} \), described by Eq. (2) was estimated from about 500 CT examinations using CTDOSE software. The summary also consisted of scanning parameters (e.g., \( kV \), \( mA \), slice thickness, table increment, and number of slices with and without contrast) used for each typical CT examination and \( \text{CTDI}_{\text{air}} \). For each examination type per hospital, the scan parameters were registered for up to six scan sequences together with the corresponding selected organ dose. From this summary, the total organ dose for the selected organs for each examination was calculated by the summation guided by its respective scan sequences. The total organ dose for selected organ per examination, together with corresponding \( \text{CTDI}_{\text{air}} \) and scanning parameters, was then used as input to the statistical software for analysis.

III. RESULTS AND DISCUSSION

A. CT scanning protocols
The mean values of scanning parameters and \( \text{CTDI}_{\text{air}} \) conducted in each hospital for CT examinations of (a) head, (b) chest, (c) abdomen, (d) lumbar spine, and (e) pelvis were analyzed to obtain the respective mean and related statistics. The results of the analysis are presented in Table 2. From the table, it is observed that CT scanning protocols used for a CT examination per hospital for most hospitals are highly standardized, similar to what has been experienced in other studies.(3,17) For example, children, a thick patient, and a thinner patient were examined with the same exposure settings, such as \( kV \), \( mA \), tube rotation time (s), table increment, pitch factor, and slice thickness. Use of exposure settings depending on the size of the patient was an occasional event in a few hospitals. In clinical practice, the patient dose increases with decreasing body size.(3,17) Although scanning protocols are standardized for most hospitals, it is evident from Table 2 that the standardization is not uniform across hospitals. However, the \( kV \) that is not indicated in Table 2 was an exception, since it was the same (120 kV) for all hospitals, except MMH (130 kV). The trend was almost the opposite for \( mA \) product, which showed a large variation ranging from 60 \( mA \) to 360 \( mA \) for head, 60 \( mA \) to 375 \( mA \) for chest, 60 \( mA \) to 355 \( mA \) for abdomen, 60 \( mA \) to 480 \( mA \) for lumbar spine, and 100 \( mA \) to 247 \( mA \) for pelvis. In general, the lowest value was consistently used by one hospital (MMH), while the highest values varied from one hospital to another. The variation of \( mA \) per given examination would be expected because of the difference in focus to isocenter distance among scanners. This is due to the fact that radiation intensity is inversely proportional to the square of the distance between the focus and the patient. That is, the shorter the distance, the lower the \( mA \) values, while the longer the distance, the higher the \( mA \) values. Of great concern in this study is the variation of \( mA \) of up to a factor of 3.3 per examination for scanners of the same model (such as of BMC and RMC).
Table 2. Summary of scanning parameters and $\mu$CTDI$_{air}$ used by each hospital

| Hospital | Exposure setting (mAs) | Slice width (mm) | Table increment (mm) | Slice No. without contrast | Slice No. with Contrast | Scan length (cm) | $\mu$CTDI$_{air}$ (mGy/100 mAs) |
|----------|------------------------|------------------|----------------------|---------------------------|------------------------|-----------------|-----------------------------|
| head     | AGKH 247.0±26.0        | 6.3±0.3          | 17.9±2.2             | 22.9±2.2                  | 27.5±1.1               | 2.2±1.2         |
| BMC      | 191.7±17.6             | 9.1±1.4          | 14.0±2.4             | 13.0±0.8                  | 47.1±0.3               | 19.7±6.0        |
| KCMC     | 309.1±50.0             | 7.3±0.7          | 18.9±3.0             | 18.9±3.0                  | 16.9±0.3               | 23.1±7.4        |
| MMH      | 275.3±32.9             | 3.9±0.3          | 32.9±6.4             | 32.8±6.9                  | 36.8±0.1               | 24.4±9.8        |
| MNH      | 241.7±25.3             | 7.6±0.5          | 17.9±1.3             | 18.0±1.0                  | 17.5±0.6               | 26.0±3.0        |
| RMC      | 60.0±2.0               | 6.3±0.4          | 18.9±3.1             | 23.9±1.4                  | 58.4±2.0               | 24.2±3.7        |
| SHMH     | 275.0±2.0              | 7.6±0.6          | 17.1±1.0             | 17.1±1.0                  | 27.2±2.3               | 24.9±4.2        |
| TMJH     | 360.0±2.0              | 6.1±0.7          | 16.5±3.7             | 22.1±0.5                  | 18.5±0.1               | 19.4±5.0        |
| chest    | AGKH 160.0±2.0         | 8.5±1.8          | 12.7±2.7             | 40.6±8.3                  | 22.6±0.4               | 81.4±2.0        |
| BMC      | 90.0±2.0               | 10.0±2.0         | 30.0±0.0             | 30.0±0.0                  | 48.1±0.0               | 60.0±0.0        |
| KCMC     | 280.0±2.0              | 10.0±2.0         | —                    | 31.0±2.8                  | 16.9±0.0               | 31.0±2.8        |
| MMH      | 124.5±2.0              | 15.0±2.0         | 35.5±2.1             | 76.9±3.3                  | 35.2±0.0               | 41.0±1.15       |
| MNH      | 280.0±2.0              | 8.3±2.6          | 25.5±3.1             | 35.5±3.1                  | 17.8±0.2               | 41.7±11.6       |
| RMC      | 60.0±2.0               | 10.0±2.0         | 20.0±2.0             | 19.5±0.7                  | 58.4±2.0               | 39.5±0.7        |
| SHMH     | 375.0±2.0              | 10.0±2.0         | 22.0±2.4             | 22.5±2.1                  | 25.1±0.0               | 44.5±3.5        |
| TMJH     | 165.0±2.0              | 10.0±2.0         | 29.4±3.2             | 30.0±2.7                  | 18.9±0.0               | 59.4±5.4        |
| abdomen  | AGKH 161.2±4.9         | 9.3±1.6          | 14.0±2.4             | 35.6±12.5                 | 25.6±0.6               | 102.9±38.0      |
| BMC      | 120.0±4.0              | 9.4±1.3          | 33.3±8.7             | 36.0±12.7                 | 47.9±0.0               | 63.9±14.8       |
| KCMC     | 310.0±24.4             | 10.0±2.0         | —                    | 34.5±13.4                 | 17.1±0.1               | 34.5±13.4       |
| MMH      | 124.7±2.0              | 14.0±2.0         | 41.0±1.5             | 35.2±0.0                  | 17.9±0.0               | 14.0±2.0        |
| MNH      | 280.0±2.0              | 10.0±2.0         | 24.9±2.7             | 25.1±2.7                  | 17.9±0.0               | 24.9±2.7        |
| RMC      | 60.0±2.0               | 10.0±2.0         | 22.5±2.1             | 46.5±7.8                  | 58.4±2.0               | 19.5±0.7        |
| SHMH     | 355.0±2.0              | 10.0±2.0         | 28.8±10.3            | 55.6±21.4                 | 25.1±0.0               | 30.1±10.0       |
| TMJH     | 180.0±2.0              | 12.0±2.0         | 30.6±8.3             | 76.6±21.6                 | 18.7±0.0               | 31.9±9.7        |
| l/spine  | AGKH 480.0±0.0         | 3.0±0.0          | 42.2±9.0             | —                         | 26.3±0.1               | 12.7±2.7        |
| BMC      | 200.0±2.0              | 3.0±0.0          | 27.0±5.0             | —                         | 48.4±0.0               | 8.1±3.4         |
| KCMC     | 440.0±39.7             | 3.0±0.0          | 23.1±4.6             | —                         | 16.6±0.2               | 7.0±1.4         |
| MMH      | 200.0±2.0              | 4.0±0.0          | 50.4±3.3             | —                         | 36.3±0.0               | 20.1±1.3        |
| MNH      | 310.0±37.1             | 3.0±0.0          | 38.6±16.0            | —                         | 58.4±2.0               | 11.4±4.9        |
| RMC      | 60.0±2.0               | 5.0±2.0          | 27.3±1.5             | —                         | 17.9±0.0               | 13.7±0.8        |
| SHMH     | 495.0±95.3             | 3.0±0.0          | 32.3±6.4             | —                         | 23.9±0.0               | 9.7±1.9         |
| TMJH     | 360.0±2.0              | 5.0±2.0          | 34.8±6.0             | —                         | 18.4±0.0               | 17.2±3.0        |
| pelvis   | AGKH 160.0±0.0         | 8.7±2.2          | 13.1±3.3             | 26.6±8.7                  | 25.7±11.5              | 56.1±26.7       |
| BMC      | 100.0±2.0              | 10.0±2.0         | 30.0±0.0             | 30.0±0.0                  | 47.9±0.0               | 60.0±0.0        |
| KCMC     | 230.0±70.7             | 10.0±2.0         | —                    | 38.5±3.5                  | 16.8±0.1               | 38.5±3.5        |
| MMH      | 157.7±2.0              | 12.0±2.0         | 32.0±4.3             | —                         | 39.4±5.2               | 39.4±5.2        |
| MNH      | 220.0±2.0              | 10.0±2.0         | 27.0±3.0             | —                         | 17.9±0.0               | 54.0±4.2        |
| TMJH     | 246.7±57.7             | 11.3±1.2         | 36.3±4.0             | —                         | 18.8±0.1               | 36.3±4.0        |

It is further evident from Table 2 that the scan length (calculated from slice thickness, table increment, and number of slices) among scanners varied up to a factor of 1.5, 2.6, 3.0, 2.9, and 2.1 for head, chest, abdomen, lumbar spine, and pelvis CT examination, respectively. These
variations as also observed elsewhere were largely caused by different scanning protocols (e.g., slice thickness, number of slices, and use of contrast) used by the hospitals. Large scan length was mainly attributed to the use of contrast media, since in most cases this procedure involves a repeated scan of the same scan length. Another scanning parameter that is not shown in Table 2 is gantry tilt, which was often done for head and lumbar spine. For about 60% of CT examinations of head and lumbar spine for all hospitals, the gantry was tilted to have slices parallel to supraorbitomeatal (SOM) and intravertebral disk lines, respectively. However, this parameter was not used as input to the software since the software does not take into account gantry tilt. Lack of this capability inevitably makes accurate estimation of organ dose difficult to achieve. Further, not taking angulation into account is expected to exaggerate dose estimation, especially dose to the lens of the eyes. Alternatively, the entrance surface dose by direct measurements on patients using TLDs could accurately estimate the dose to this superficial organ. The angulation of gantry parallel to the SOM has proved to some extent to protect the lens of eyes from CT examination of the brain. Apart from angulation of the gantry, no hospital in this study was found protecting superficial radiosensitive organs such as the lens of the eyes, thyroid, and breast, which are rarely the organs of clinical concern in the CT examinations. It has been shown that by using radioprotective material such as bismuth for shielding superficial radiosensitive organs, the radiation dose to the patient can be reduced significantly without loss of image quality.

B. CT dose measurements

Of interest in this study is the determination of organ doses. However, as described earlier, this quantity cannot be determined without the knowledge of normalized CTDI (nCTDI). The nCTDI to mAs for CT scan under typical operation conditions for each scanner model were determined as described earlier from measurements of CTDI in air (nCTDIair). The results of the measurements of nCTDIair for CT scans under typical operation conditions for each scanner are presented in Fig. 1. Derived estimates of CTDI normalized to the exposure setting, nCTDIair (mGy (mAs)−1) have been analyzed by scanner type, nominal slice width, applied potential, and focus-to-axial distance. Overall, individual measurements revealed that a variation as large as a factor of 2.8 (0.17 to 0.48 mGy (mAs)−1) was obtained between the mean values of nCTDIair per scanner model.

Fig. 1. Normalized CT dose index free in air: Comparison between types of CT scanner models
It is interesting to observe in Fig. 1 that similar makes but different models of scanners can have such large deviations of $n_{CTDI_{air}}$ and, hence, organ doses. The relatively high values shown for Tomoscan M-EG are probably a function of their short focus-to-axial distance as shown in Table 1. This is due to the fact that radiation intensity varies as the inverse of the squared distance between the source of the radiation and the object (patient). As a result, if all other scanning parameters are held constant, the scanner with the shorter distance between the X-ray tube focal spot and the isocenter of the gantry aperture can produce more radiation exposure than the long geometry scanner. On the other hand, the variation of source-to-detector distance among scanners might affect the image quality. This is due to the fact that image noise in CT is known to be inversely proportional to the square root of the number of photons received by the detector, whereas the number of photons (dose) is inversely proportional to the squared distance between the source of the radiation and the detector. As a result, if all other scanning parameters are held constant, the scanner with the long distance between source and detector can have higher image noise than the short geometry scanner. The difference in the case of Somatom AR.Star and Somatom Plus 4 is probably the result of the use of different kV and focus-to-axial distances. The $n_{CTDI_{air}}$ are also presented in Fig. 1 for different scanners using different slice thicknesses. From this figure, it is evident that the effect of slice thickness on $n_{CTDI_{air}}$ and, hence, organ dose is marginal for some scanners (i.e., Somatom Plus 4 and Tomoscan SR 4000) and significant for others (i.e., CT/e and Somatom AR.Star).

C. Doses to selected organs

The total mean organ doses of selected organs were determined as described earlier for different examinations including the head, chest, abdomen, lumbar spine, and pelvis using patient exposure parameters specific to hospitals and scanners used. The results of estimated total mean organ doses for selected organs per examination are presented in Fig. 2 and Table 3. It is evident from the table that large variations of organ dose exist within and among hospitals. For instance, the mean organ doses per hospital for the eye lens (for head) and breast (for chest examination) varied up to a factor of 2.7 and 2.4, respectively, while for the stomach (for abdomen) and ovary (for pelvis), the variation was up to a factor of 2.1 and 2.4, respectively. The observed wide variation of organ doses in Figs. 3 to 6 for all CT examinations of head, chest, and abdomen, respectively, is an indication that different scanning parameters (i.e., kV, mAs, scan length, and use of contrast media) used in hospitals have a significant influence on organ dose determinations. It was observed that MMH always had the highest organ doses, followed by the TMJH. MMH, for example, uses CT scanners operated at a higher voltage (130 kV), while the rest use 120 kV. Other factors include a contribution of use of larger scan length, exposure settings (mAs), and contrast media. The use of contrast media by some hospitals was done as a routine procedure without clinical justification of using it. In a few hospitals, such as AGKH and TMJH, the radiologist was consulted before IV administration of contrast. KCMC appears to consistently have the lowest organ doses for the chest (Table 3(b)), abdomen (Table 3(c)), and pelvis (Table 3(e)). This was expected because the protocols used exclude scan sequence without contrast. The observed high mean organ doses at TMJH for pelvis examination relative to other hospitals is an indication that large scan length has a significant impact on organ dose.
Fig. 2. Absorbed dose for selected organs from CT examination type

Fig. 3. Histogram of dose distribution to the lenses of the eyes for all CT examinations of the head

Fig. 4. Histogram of dose distribution to the breast for all CT examinations of the chest

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Fig. 5. Histogram of dose distribution to the stomach for all CT examinations of the abdomen

Fig. 6. Histogram of dose distribution to the gonads for all CT examinations of the pelvis
Despite the fact that CT scanning protocols were highly standardized for most of the hospitals, wide variation doses among patients up to a factor of 3 were also observed within the hospitals. For instance, the doses to the eye lenses for TMJH and KCMC varied from 52.0 mGy to 130.6 mGy and from 30.5 mGy to 109.4 mGy, respectively, while for MNH and MMH, the doses varied from 30.8 mGy to 82.0 mGy and from 49.5 mGy to 149.0 mGy, respectively. These variations were largely attributed to variation of clinical indications among patients, use of contrast, and number of slices used depending on patient size. It was further observed that organ dose distributions (shown in Figs. 3 to 6) for a given examination vary to an order of magnitude with a broad and relatively flat distribution. Such a wide spread in organ doses observed in this study is comparable to similar surveys performed in other countries.\(^{(6,9,10,13,18,19)}\) For instance, the wide spread of doses to the lenses of the eyes in this study ranged from 12.8

Table 3. Summary of mean organ doses per hospital for each CT examination

| Selected organ | AGKH | BMC | KCMC | MMH | MNH | RMC | SHMH | TMJH |
|----------------|------|-----|------|-----|-----|-----|------|------|
| (a) Head       |      |     |      |     |     |     |      |      |
| RBM\(^a\)      | 2.4±1.0 | 3.9±1.5 | 3.4±1.1 | 5.8±1.7 | 3.3±0.5 | 2.1±0.3 | 2.9±0.5 | 4.5±1.1 |
| brain          | 37.5±16.1 | 62.6±23.4 | 52.0±1.0 | 84.4±22.6 | 51.2±9.5 | 32.5±4.8 | 48.4±8.5 | 72.1±16.5 |
| eye lens       | 41.0±21.0 | 75.0±28.1 | 57.0±10.1 | 110.3±35.2 | 53.6±11.1 | 40.2±6.3 | 92.8±9.6 | 92.5±21.0 |
| thyroid        | 1.6±0.8 | 2.4±1.0 | 2.5±1.0 | 4.7±1.5 | 2.3±0.4 | 1.5±0.4 | 2.3±1.1 | 3.0±1.0 |
| (b) Chest      |      |     |      |     |     |     |      |      |
| RBM\(^a\)      | 7.8±3.2 | 12.4±4.4 | 7.0±0.5 | 10.7±6.0 | 10.3±2.6 | 7.9±0.1 | 11.4±0.8 | 11.6±1.0 |
| lung           | 20.1±6.3 | 44.0±7.9 | 24.0±0.0 | 25.3±10.1 | 34.6±12.7 | 31.5±3.5 | 36.0±5.7 | 38.9±2.1 |
| breast         | 14.8±5.0 | 36.0±8.1 | 20.0±0.0 | 21.0±8.1 | 29.6±17.1 | 29.5±7.0 | 31.0±14 | 33.1±10 |
| thyroid        | 21.5±5.0 | 9.0±2.7 | 4.9±0.0 | 4.6±0.0 | 8.7±2.1 | 3.5±1.9 | 12.1±1.0 | 9.3±1.2 |
| esophagus      | 23.3±0.0 | 50.0±5.5 | 27.0±0.0 | 27.0±0.0 | 52.0±5.5 | 42.5±0.7 | 68.0±0.0 | 43.6±0.8 |
| (c) Abdomen    |      |     |      |     |     |     |      |      |
| liver          | 26.1±10.1 | 42.8±5.2 | 21.0±1.4 | 28.0±4.2 | 41.6±6.2 | 33.0±1.4 | 40.5±10.2 | 30.7±6.5 |
| stomach        | 28.5±10.9 | 46.4±6.0 | 22.5±0.7 | 28.0±3.5 | 42.4±6.5 | 36.0±0.0 | 42.7±11.6 | 32.3±6.9 |
| pancreas       | 24.9±9.8 | 39.6±4.9 | 18.5±0.7 | 26.0±6.0 | 37.9±5.5 | 31.0±1.4 | 44.7±13.5 | 28.7±6.2 |
| spleen         | 26.8±10.4 | 42.1±5.3 | 20.5±0.7 | 28.0±1.9 | 41.5±6.0 | 33.0±2.3 | 37.9±9.4 | 30.5±6.2 |
| kidney         | 32.0±12.6 | 51.6±7.5 | 25.0±0.5 | 30.0±5.0 | 44.6±7.2 | 39.0±1.7 | 47.3±13.6 | 47.3±7.2 |
| adrenal        | 26.4±10.4 | 40.1±5.2 | 18.5±0.7 | 26.0±2.8 | 40.3±5.6 | 32.0±1.4 | 49.4±14.3 | 29.7±6.1 |
| (d) Lumbar spine |      |     |      |     |     |     |      |      |
| large intestine | 32.4±2.4 | 29.0±9.2 | 9.6±3.0 | 21.1±1.1 | 9.4±3.8 | 11.5±0.5 | 17.9±0.0 | 17.4±7.3 |
| small intestine | 32.9±3.5 | 28.0±6.2 | 10.5±2.7 | 22.4±1.3 | 11.9±3.9 | 11.5±0.7 | 18.1±0.0 | 18.0±6.2 |
| colon          | 17.8±4.6 | 8.8±3.2 | 8.3±2.3 | 18.3±1.4 | 8.5±2.3 | 5.2±0.6 | 9.3±3.4 | 12.0±2.8 |
| gonads         | 8.2±18.8 | 7.4±2.6 | 11.7±6.9 | 11.5±11.2 | 8.1±5.7 | 9.0±1.8 | 2.6±3.1 | 12.6±10.4 |
| uterus         | 28.6±7.3 | 6.2±2.1 | 15.6±8.2 | 27.3±2.8 | 11.9±5.9 | 8.7±3.1 | 14.4±6.5 | 19.0±5.8 |
| (e) Pelvis     |      |     |      |     |     |     |      |      |
| ovaries        | 21.0±1.0 | 23.0±0.0 | 16.4±5.1 | 21.8±0.0 | 32.2±0.0 | — | — | 39.7±16.2 |
| bladder        | 26.0±13.5 | 26.0±0.0 | 19.5±6.4 | 25.0±0.0 | 38.0±0.0 | — | — | 38.6±24.7 |
| uterus         | 23.4±11.8 | 26.0±0.0 | 17.5±4.9 | 23.0±0.0 | 34.0±0.0 | — | — | 42.7±18.1 |
| testis         | 11.2±16.0 | 5.0±0.0 | — | 7.6±0.0 | 7.0±0.0 | — | — | 16.7±11.4 |

\(^a\) RBM = red bone marrow.
mGy to 149 mGy (median 53.8 mGy), while in Australia, Norway, and Japan the spread ranged from 10 mGy to 160 mGy (median of 61 mGy), 39.1 mGy to 108.6 mGy, and 8.7 mGy to 47.2 mGy, respectively.\textsuperscript{13,18,19} The wide spread of organ dose distribution is again influenced by variation of techniques used among scanners.

D. Comparison with other studies

To facilitate the comparison between nations, the mean values of number of slice ($n$) and mean organ doses of selected organs per examination in this study and from reported values from the literature for the United Kingdom, Germany, the Netherlands, and Japan have been presented in Table 4.\textsuperscript{6,9,10,19} It is clear from the table that with the exception of values reported from Japan, the mean organ doses per given examination were mostly comparable with those from other studies. For instance, the variation of organ doses between this study and reported values from the Netherlands and the United Kingdom mostly varied by up to a factor of 1.0 and 1.7, respectively, while for Germany and Japan, the organ doses mostly varied by up to a factor of 2.3 and 3.0, respectively. On the one hand, the higher organ doses observed in this study relative to that reported from Japan might be attributed to the different method used for estimation of organ doses, whereby that of Nishizawa et al.\textsuperscript{19} used TLD in a female Rando Alderson phantom. In addition, the higher organ doses in this study relative to those of Japan might also have been attributed to the fact that Nishizawa et al. estimated the organ dose from a single scan at a fixed number of slices for a given examination, while in this study organ doses were estimated from an examination with either one scan (i.e., without or with contrast) or with two scans, one with and one without contrast.

Table 4. Comparison of mean organ doses in this study and other studies

| CT examination | Selected organ | This study | UK\textsuperscript{(10)} | Germany\textsuperscript{(6)} | Netherlands\textsuperscript{(9)} | Japan\textsuperscript{(19)} |
|----------------|---------------|-----------|-------------------|-------------------|-------------------|-------------------|
|                | $n^a$ | $D$ (mGy) | $n^a$ | $D$ (mGy) | $n^a$ | $D$ (mGy) | $n^a$ | $D$ (mGy) | $n^a$ | $D$ (mGy) |
| head | RBM\textsuperscript{b} | 42 | 3.5±1.6 | 25 | 2.7 | — | — | 9 | 1.5 |
|        | eye lens | 42 | 63.9±32.6 | — | — | 24 | 24.8 | — | 9 | 22.4 |
|        | thyroid | 42 | 2.5±1.3 | 25 | 1.9 | — | — | 9 | 0.6 |
| chest | RBM\textsuperscript{b} | 60 | 9.9±2.8 | 34 | 5.9 | — | — | 28 | 10 | 16 | 5.7 |
|        | lung | 60 | 31.5±10.6 | 34 | 22.4 | 49 | 20.5 | 28 | 37 | 16 | 19.6 |
|        | breast | 60 | 26.1±10.8 | 34 | 21.4 | 49 | 22.6 | 28 | 32 | 16 | 15.9 |
|        | thyroid | 60 | 12.3±8.5 | 34 | 2.3 | — | — | 28 | 7 | 16 | 1.9 |
| abdomen | liver | 61 | 34.1±10.3 | 31 | 20.4 | 32 | 15.0 | 31 | 35.5 | 13 | 27.8 |
|        | stomach | 61 | 35.6±10.7 | 31 | 22.2 | 32 | 15.4 | 31 | 38.5 | 13 | 26.9 |
| pelvis | ovary | 62 | 24±17.1 | 25 | 22.7 | 52 | 14.9 | — | 12 | 15.1 |
|        | bladder | 62 | 28.8±21.2 | 25 | 23.2 | 52 | 16.1 | — | 12 | 10.6 |
|        | uterus | 62 | 26.5±18.6 | 25 | 25.5 | 52 | 14.6 | — | — | — |
|        | testis | 62 | 12.5±19.9 | 25 | 1.7 | — | — | — | 12 | 1.0 |

\textsuperscript{a} $n$ is the number of slices (with and without contrast).

\textsuperscript{b} RBM = red bone marrow.
On the other hand, the differences in organ doses between this study and those reported in the literature for the United Kingdom, Germany, and the Netherlands with NRPB organ dose conversion factors were mainly attributed to the variation in CT scanning protocols (i.e., kV, mAs, slice thickness, number of slices, use of contrast media, etc.) and type of scanners used. For instance, the average number of slices used in this study was higher by factor of 1.7 to 2.5, 1.2 to 1.9, and 2.1 than the average number of slices reported in the literature for the United Kingdom, Germany, and the Netherlands, respectively. This further suggests that there exists potential for dose reduction in Tanzanian hospitals through decreasing numbers of slices (hence scan length). Moreover, the mean doses for selected organ per examination in this study were found to be comparable to those reported by Olerud of Norway.\(^{(18)}\) For example, the mean organ dose in this study for the lenses of the eyes, red bone marrow (for head), ovary (for pelvis), and testis (for pelvis) were 63.9 mGy, 3.5 mGy, 24.0 mGy, and 12.5 mGy, respectively, while for Norway they were 80.9 mGy, 3.2 mGy, 26.0 mGy and 7.0 mGy, respectively.

In view of the observed causes of organ dose variations and similar experience observed elsewhere,\(^{(6,10,16–18)}\) further studies on the optimization of organ doses to patients undergoing CT examinations are needed. There are a number of observed parameters that greatly need optimization:

1. The first is the minimization of the number of slices (hence scan length) as much as possible, without missing any vital anatomical region. Several studies have recommended that with the reduction of irradiation volume depending on body region being scanned, radiation dose to patients can be significantly reduced.\(^{(3,9,10,13,17,18)}\)

2. The second is minimization of tube current (mA) based on indication of study. Some studies have already revealed that adjustment of mA based on indication of study dose to patient can be reduced to 50\% without significantly affecting the image quality.\(^{(20,21)}\)

3. The third approach is through modulation of exposure parameters (i.e., kV, mA, exposure time, and slice thickness) based on patient size and age, while maintaining a constant contrast to image-to-noise ratio. Some studies have demonstrated that by adjusting the exposure parameters based on patient age, weight, or transverse diameter of the body part imaged, patient radiation doses can be reduced significantly.\(^{(17,21,22)}\)

4. Another possible method is through use of contrast media only to optimize diagnostic yield. As mentioned earlier, use of contrast media by some of the hospitals was done as a routine procedure without clinical justification of using it. In only a few hospitals (AGKH, TMJH, and MMH) were the radiologists consulted before the IV administration of contrast.

5. The last possible method is the use of radioprotective materials for protecting superficial radiosensitive organs such as the lenses of the eyes, thyroid, and breast. It has been demonstrated that in-plane bismuth shields could significantly reduce absorbed dose to these superficial organs without loss of image quality.\(^{(3,7,8)}\)

However, in order to achieve the above optimization strategies, there is a great demand to educate CT personnel on the effects of scan parameter settings on radiation dose to patients and image quality required for accurate diagnosis.

**IV. CONCLUSIONS**

The assessment of radiation dose to selected radiosensitive organs of patients undergoing CT examinations in Tanzania was investigated. In this study, large variations of radiation dose to various organs were observed. Different scanning protocols used among hospitals and variation in equipment design among manufacturers and models were responsible for these variations.
The mean organ doses in this study were mostly comparable to and slightly higher than reported values from the United Kingdom, Germany, the Netherlands, Norway, and Japan. The main contributor for this difference was the use of a larger scan length in Tanzania than that used in some of these countries. The large observed variations of organ doses among hospitals and relatively high organ doses in Tanzanian hospitals call for the need to optimize CT scanning protocols. This can be achieved through optimal selection of scanning parameters based on indication of study, body region of interest being scanned, and patient size. In addition, further studies should be done to investigate the potential for using radioprotective materials to protect superficial radiosensitive organs.

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REFERENCES

1. United Nations Scientific Committee on Effects of Radiation Atomic Radiation (UNSCEAR). Sources and effects of ionising radiation report to the general assembly. New York: United Nations; 2000.
2. International Atomic Energy Agency (IAEA). Protection of patients in diagnostic and intervention radiology, nuclear medicine and radiotherapy. Proceeding of International Conference, Malaga, Spain, 26–30 March 2001; Vienna.
3. International Commission on Radiological Protection (ICRP). Managing patient dose in computed tomography annals of ICRP publication 87; 30(4). Oxford: Pergamon Press; 2000.
4. Muhogora WE, Nyanda AM, Lema US, Ngaile JE. Typical radiation doses to patients from some common x-ray examinations in Tanzania. Radiat Prot Dosim. 1999;82(4):301–305.
5. Brenner DV, Elliston CD, Hall EJ, Berdon WE. Estimated risks of radiation fatal cancer from paediatric CT. Am J Roentgenol. 2001;176:289–296.
6. Hidajat N, Maurer J, Schroder RJ, et al. Relationships between physical dose quantities and patient dose in CT. Br J Radiol. 1999;72:556–561.
7. Hopper KD, Neuman JD, King SH, Kunselman AR. Radioprotection to the eye during CT scanning. Am J Neuroradiol. 2001;22:1194–1198.
8. Hopper KD, King SH, Lobell ME, TenHave TR, Weaver JS. The breast: In-plane X-ray protection during diagnostic thoracic CT—shielding with bismuth garments. Radiology 1997;205:853–858.
9. Geleijns J, Van Unnik JG, Zoetelief J, Zweers D, Broeze JJ. Comparison of two methods for assessing patient dose from computed tomography. Br J Radiol. 1994;67:360–365.
10. Shrimpton PC, Jones DG, Hillier MC, et al. Survey of CT practice in the UK. Part 2: Dosimetric aspects. Chilton, NR PB-R249. London: HMSO; 1991.
11. European Commission. European guidelines on quality criteria for computed tomography EUR 16262 EN, Luxemburg: Office for Official Publications of the European Communities; 1999.
12. Jessen KA, Shrimpton PC, Geleijn J, Panzer W, Tosi G. Dosimetry for optimisation of patient protection in CT. Appl Radiat Isot. 1999;50:165–172.
13. Thomas JEM, Tingeys DRC. Radiation doses from computed tomography in Australia. Australian Radiation Laboratory Report, ARL/1R123; 1997.
14. International Electrotechnical Commission (IEC). Medical electrical equipment. Part 2–44: Particular requirements for the safety of X-ray equipment or computed tomography. IEC 60601-2-44. Geneva, Switzerland; 1999.
15. Imaging Performance Assessments of CT (ImPACT). CT patient dosimetry spreadsheet (v0.99u, 12/12/2003). Retrieved April 2004 from www.impactscan.org/ctdosimetry.htm.
16. Jones DG, Shrimpton PC. Normalised organ doses for X-ray computed tomography calculated using Monte Carlo techniques. NR PB-SR250. National Radiological Protection Board (NRPB). Chilton, UK; 1993.
17. Huda W, Scalzetti EM, Roskopf M. Effective dose to patients undergoing thoracic computed tomography examinations. Med Phys. 2000;27(5):835–844.
18. Olerud HM. Analysis of factors influencing patient dose from CT in Norway. Radiat Prot Dosim. 1997;71(2):123–133.
19. Nishizawa K, Maruyama T, Takayama M, Okada M, Hachiya J, Furuya Y. Determination of organ doses and effective dose equivalents from computed tomographic examination. Br J Radiol. 1991;64:20–28.
20. Mayo JR, Hartman TE, Lee KS, Primack SL, Vedal S, Muller NL. CT of the chest: Minimal tube current required for good image quality with the least radiation dose. Am J Roentgenol. 1995;164:603–607.
21. Donnelly LF, Emery KH, Brody AS, et al. Minimizing radiation dose for paediatric body applications of single-detector helical CT. Am J Roentgenol. 2001;176:303–306.
22. Starck G, Lonn L, Cederblad A, Forsell-Aronsson E, Sjostrom L, Alpsten M. A method to obtain the same levels of CT image noise for patients of various sizes to minimize radiation dose. Br J Radiol. 2002;75:140–150.