Investigation of Eye Lens Dose Estimate based on AAPM Report 293 in Head Computed Tomography

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ABSTRACT

Background: Estimation of eye lens dose is important in head computed tomography (CT) examination since the eye lens is a sensitive organ to ionizing radiation.

Objective: The purpose of this study is to compare estimations of eye lens dose in head CT examinations using local size-specific dose estimate (SSDE) based on size-conversion factors of the American Association of Physicists in Medicine (AAPM) Report No. 293 with those based on size-conversion factors of the AAPM Report No. 220.

Material and Methods: This experimental study is conducted on a group of patients who had undergone nasopharyngeal CT examination. Due to the longitudinal (z-axis) dose fluctuation, the average global SSDE and average local SSDE (i.e. particular slices where the eyes are located) were investigated. All estimates were compared to the measurement results using thermo-luminescent dosimeters (TLDs). The estimated and measured doses were implemented for 14 patients undergoing nasopharyngeal CT examination.

Results: It was found that the percentage differences of the volume CT dose index (CTDIvol), average global SSDE based on AAPM No. 220 (SSDEo,g), average local SSDE based on AAPM No. 220 (SSDEo,l), average global SSDE based on AAPM No. 293 (SSDEn,g) and average local SSDE based on AAPM No. 293 (SSDEn,l) against the measured TLD doses were 22.5, 21.7, 15.0, 9.3, and 2.1%, respectively. All comparisons between dose estimates and TLD measurements gave p-values less than 0.001, except for SSDEn,l (p-value = 0.566).

Conclusion: SSDE based on AAPM Report No. 293 can be used to accurately estimate eye lens radiation doses by performing the calculations on a number of specific slices containing the eyes.

Keywords

Radiation; Ionizing; X-Rays; Computed Tomography; Algorithms; Eye Lens Dose; Organ Dose; Size-Specific Dose Estimates

Introduction

Computed tomography (CT) scanners have long been used to stage and assess patients suspected of having nasopharyngeal carcinoma (NPC) [1, 2]. The NPC detection is usually followed by injection of contrast agent to enhance differentiation among surrounding tissues to facilitate the evaluation of the lesions [3]. The NPC...
examination combined with contrast agents necessitates the scanning to be conducted twice, i.e. pre- and post-contrast agent injections. As a result, the radiation dose accumulated to the patient is higher than a general head CT examination, resulting in a concern particularly for the eye lens considered to be a high radiosensitive tissue. A recent study highlighted the risk of lens opacity as a potential consequence related to exposure, even for low doses of ionizing radiation [4]. Others have reported radiation-induced cataract formation [5, 6]. In order to assess the radiation risks, accurate estimates of the doses deposited in the eye lens of exposed individuals under the NPC examinations are urgently required.

Experimental eye lens dose measurements are often carried out using an anthropomorphic phantom and small detectors, i.e. thermoluminescent dosimeters (TLDs) [7, 8], metal-oxide-semiconductor field-effect transistors (MOSFET) [9], or optically stimulated luminescent (OSL) dosimeters [10, 11]. A computational approach can be carried out through Monte Carlo (MC) simulations using a voxel phantom [8, 12, 13] to represent a wide range of sizes [14]. MC simulations provide a closer estimation of organ dose to a patient, but it is computationally time-consuming and requires highly specialized skill intervention to simulate a voxel model [8]. An alternative is to make the use of pre-computed organ doses from MC computational phantom libraries, but several of the existing CT organ dose estimation tools are considered to be out-of-date and not entirely representative of human anatomy [15].

The dose descriptor, directly accessible at the scanning, is the volume of computed tomography dose index (CTDI\textsubscript{vol}). Studies have been conducted to correlate the CTDI\textsubscript{vol} displayed on the CT console and the eye lens dose among patients [16-18]. Zhang et al. [16] reported that the ratio of eye lens dose to CTDI\textsubscript{vol} was 59 - 63%. More recently, Lopez-Rendon et al. [8] reported that CTDI\textsubscript{vol} overestimated the eye lens dose by up to 41%. Suzuki et al. [17] reported that the ratio of eye lens doses to CTDI\textsubscript{vol} fell in a range of 81 - 103%, showing that CTDI\textsubscript{vol} can only be applied to estimate the eye lens dose in a specific clinical setting. In fact, CTDI\textsubscript{vol} provides the fundamental output radiation dose for scanners, but has limitations for estimating patient dose due to the absence of patient size in its framework [18].

For the purpose of estimating absorbed dose in patients, the American Association of Physicists in Medicine (AAPM) developed the size-specific dose estimate, SSDE (Report No. 204, 2011). The SSDE takes into consideration the CTDI\textsubscript{vol} and a conversion factor (f) describing the patient size in terms of an effective diameter (D\textsubscript{eff}) [19]. In 2014, AAPM improved the SSDE concept in Report No. 220 and introduced the water-equivalent diameter (D\textsubscript{w}), which took into account the geometrical size and patient attenuation [20]. The D\textsubscript{w} metric had previously been proposed by Wang et al. [21] and was subsequently adopted as the gold standard in representing radiological patient size for dose estimation. However, the size-conversion factors tabulated in AAPM Reports No. 204 and No. 220 were specifically modeled in the abdominal-pelvic region. Therefore, in 2019 AAPM renewed the size-conversion factor for the head region in Report No. 293, taking into account that the dose accumulated in the brain is significantly different from the doses absorbed in other parts of the body [22].

The SSDE metric may be a practical approach for estimating organ dose [18]. Several groups have explored the potential correlation between SSDE and organ doses [23-25]. The radial dose distributions at the center and the periphery of the patient may not be uniform, particularly in the body region [18]. However, the dose distribution in the center
of a 16 cm PMMA phantom was reported to be similar to the dose at the periphery [26]. In another study, Anam et al. [27] explored various physical sizes of acrylic phantoms within a range of 8-32 cm and reported that $D_w$ within a range of 12–14 cm had a homogeneous dose distribution for both central and peripheral regions. Therefore, the SSDE may be useful for organ dose estimation within the head region without any consideration of fluctuations in the radial dose distribution. Another uncertainty in estimating the organ dose from SSDE is the dose distribution along the $z$-axis. An accurate organ dose estimation needs to consider the $z$-axis distribution by employing local SSDE at particular $z$-axis values [25]. This is because the SSDE is strongly dependent on fluctuations of $D_w$ along the $z$-axis.

The concept of SSDE to estimate the eye lens dose of patients, undergoing nasopharyngeal CT examinations by taking the specific $z$-axis values within the eye region (according to AAPM Report No. 220), was investigated by Anam et al. [28]. The estimated doses were then compared to the measured doses from TLDs. The results showed that that approach provided a closer match to the measured radiation dose than the average of the global SSDE, although the discrepancy from the TLD measurement was still more than 10% [28]. With the update of the size-conversion factor provided by AAPM Report No. 293, the SSDE may estimate the organ dose more accurately. This study aims to investigate the eye lens dose by implementation of the local SSDE in $z$-axis specific regions of patients undergoing nasopharyngeal CT examination using the AAPM Report No. 293 size-conversion factors.

**Material and Methods**

**Patient preparation**

This experimental study was conducted with a group of patients, who had undergone nasopharyngeal CT examination at Prof. Dr. Margono Soekarjo Hospital, Purwokerto, Central Java, Indonesia. A total of fourteen patients (3 male and 11 female), ranging from 21 to 72 years old were included in this study. Eight patients were assigned to contiguous axial mode and the other six patients were examined in helical mode. All scans were performed with a Somatom Emotion 6 CT scanner (Siemens AG, Forchheim, Germany), operating at 120 kVp and 250 mAs. The examinations obtained anatomical images along the vertex to skull base with a total of 29 - 34 slices with thickness of each is 4 mm and reconstruction diameter of 200 mm.

**Eye lens radiation dose measurement**

The high sensitive TLD-100 chips (Harshaw Chemical Company, Solon, Ohio, USA) with dimension of $3.175 \times 3.175 \times 0.889$ m$^3$ were used for measuring the eye lens radiation dose. The TLD-100 contains 92.5% $^7$Li plus 7.5% $^6$Li and impurities, including magnesium and titanium to increase electron traps and sensitivity. Effective atomic number of the TLD-100 is nearly similar to those of the body tissues. For simplicity, the TLD-100 is written in the current report as TLD. The calibration of the TLDs was performed at Department of Dosimetry, Center of Safety Technology and Radiation Metrology, National Nuclear Energy Agency (PTKMR-BATAN) as the secondary standard dosimetry laboratory (SSDL) in Indonesia. The annealing cycle consisted of a heating phase (400 °C for 1 hour in a furnace, 200 °C for 2 hours in an oven). The TLDs were then cooled down until equilibrated to room temperature and packed into plastic bags each containing three TLDs chips. The TLDs were positioned on the eye surfaces and read by a TLD reader of 2000 A/B (Harshaw, Chemical Company, Solon, Ohio, USA) after irradiation. Accurate measurement of the eye lens dose required to put 5 mm of tissue equivalent on TLD chips.
However, measurements in the current study were conducted without additional 5 mm of tissue equivalent, hence the dose obtained was the entrance surface dose (ESD) of eye lens rather than the eye lens dose. In the current study, the term of the eye lens dose was used as an approximation of the ESD of eye lens. The final estimate of the eye lens dose was taken as the average dose from the three TLDs within a bag multiplied by the corresponding calibration factor. The measured dose from the TLDs was used as a reference for dose assessment.

**Calculation of $D_w$**

In this study, the patient size was determined using the attenuation-based size metric ($D_w$) calculated from the patient image. To determine the value of $D_w$, a derivative formula from AAPM Report No. 220 was adopted:

$$D_w = 2 \sqrt{\frac{1}{1000} CT(x,y) + 1} \frac{A}{\pi} \quad (1)$$

where $A$ is the patient area for every slice and $CT(x,y)$ is the average value of CT number inside the patient area, expressed in Hounsfield units (HU). An automated method was used to obtain the $D_w$ value [29]. Using Equation (1), the average of the global $D_w$ ($D_{wg}$) and the average of the local $D_w$ ($D_{wl}$) values characterize the dose measurement of eye organs for each patient. $D_{wg}$ is the average value of total $D_w$ from the $N$ slices along the scan axis.

$$D_{wg} = \frac{\sum_{i=1}^{N} D_{wi}}{N} \quad (2)$$

$D_{wl}$ is the average value of local $D_w$ calculated using a certain number of slices ($n$), containing the eye’s field of view. To determine the value of $D_{wl}$, the position of the eye organs along the $z$-axis is manually observed and defined by the user. Then the diameter was calculated using Equation (1) and divided by the number of slices ($n$) where the eye organs are located to give $D_{wl}$ (Equation (3)). The determination of the value of $D_{wg}$ and $D_{wl}$ is illustrated in Figure 1.

$$D_{wl} = \frac{\sum_{i}^{n} D_{wi}}{n} \quad (3)$$

**Dose Calculations**

The calculated values of global $D_w$ and local $D_w$ were then used to estimate the SSDE val-

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**Figure 1:** $D_{wg}$ represents the average value of total $D_w$ at the start to the end of scanning along the $z$-axis, while $D_{wl}$ is the average of local $D_w$ is calculated only from the image of the eye’s field of view that (a) illustrates the positions taken in calculating $D_{wg}$ and $D_{wl}$, and (b) shows the $D_w$ profiles from the 1st slice to the 24th slice. In this case, $D_{wl}$ is calculated as the average of local $D_w$ from the 11th to 17th slices ($n = 6$).
Eye Lens Dose Estimate Based on AAPM 293

value for each slice. SSDE formula was shown in Equation (4). The dose estimation in this current study used the size-conversion factor of AAPM Report No. 293 ($f_n$), and a CTDI$_{vol}$ value recorded from the CT scan console screen, rather than the conversion factors ($f_o$) from AAPM Report No. 220 used in the previous study. The $D_w$ and SSDE were calculated using the IndoseCT 20b software [30]. Plots of the size-conversion factors ($f_o$ and $f_n$) versus $D_w$ are shown in Figure 2.

The SSDE can be estimated by two approaches, i.e. the average of global SSDE ($SSDE_{n,g}$) showing the estimated dose values across all the slices during the examination, and the average of local SSDE ($SSDE_{n,l}$) represents the approximate dose at the slices where the eye organs are located. Both were shown in Equations (5) and (6), respectively. For comparison, the estimated doses calculated based on CTDI$_{vol}$ and the AAPM Report No. 220 conversion factor ($SSDE_{o,g}$ and $SSDE_{o,l}$) were calculated.

\[
SSDE = CTDI_{vol} \times f_{(size)} \quad (4)
\]

\[
SSDE_g = \frac{\sum_{i=1}^{N} SSDE_i}{N} \quad (5)
\]

\[
SSDE_l = \sum_{i=1}^{n} SSDE_i \quad (6)
\]

Statistical analysis

A comparison of doses indicators, i.e. CTDI$_{vol}$, SSDE$_{o,g}$, SSDE$_{o,l}$, SSDE$_{n,g}$, and SSDE$_{n,l}$ against the measured doses of TLDs was obtained, along with the average and standard deviation values of their percentage differences. A statistical test using Wilcoxon Mann-Witney U test at a significance level of 0.05 was performed to find if there is a statistically significant difference. All statistical analysis was conducted using the Matlab software (Mathworks Inc., Natick, Massachusetts, USA).

Results

Values of $D_{w,g}$ and $D_{w,l}$

$D_{w,g}$ (indicated by circular markers) and $D_{w,l}$ (indicated by diamond markers) showed a similar trend (Figure 3a). The values of $D_{w,g}$ (16.41 ± 1.04 cm) were smaller than the values of $D_{w,l}$ (17.82 ± 0.79 cm) for all patients included in this study. The relationship between them shows a statistically significant difference ($p$-value = 0.001).

Comparisons of eye dose estimations against TLDs

The measured doses using TLDs were used as the gold standard in this current study. The comparisons of the CTDI$_{vol}$, SSDE$_{o,g}$, SSDE$_{o,l}$, SSDE$_{n,g}$, and SSDE$_{n,l}$ against TLDs doses are presented in Figure 4 and Table 1. As expected, the CTDI$_{vol}$ value was constant for each patient. The four estimated doses in terms of SSDE had a similar trend as the TLDs dose, but the ones that come closest to the measurement results were SSDE$_{n,l}$ (percentage difference of 2.1 ± 4.2%), followed by SSDE$_{n,g}$ (9.3 ± 4.3%), SSDE$_{o,l}$ (15.0 ± 5.7 %), and SSDE$_{o,g}$ (21.7 ± 5.3 %). Statistical analysis indicates that there was no significant difference be-
It has been reported that the radial distribution of dose in a head CT examination is homogeneous [26], so that the SSDE, which is an average dose within a patient, can be used to estimate the dose even in the organs at the edge of the head such as the eye. However, the diameter of the head of every patient fluctuates along the $z$-axis, and an accurate estimate of organ dose should consider this fluctuation. The SSDE along the $z$-axis depends on $D_{w,g}$ in fixed tube current (FTC) and depends on both $D_{w}$ and tube current in tube current modulation (TCM) [27].

A previous study [28] reported estimated eye lens doses by calculating the average of the global SSDE ($SSDE_{n,g}$) and the average of the local SSDE ($SSDE_{o,l}$) using AAPM Report No. 220 (16 cm CTDI phantom). The results showed that $SSDE_{o,l}$ was closer than $SSDE_{n,g}$ to the results measured using TLDs. The discrepancies between the $SSDE_{o,l}$ results and

**Discussion**

Figure 3: (a) The values of $D_{w,g}$ and $D_{w,l}$ for all 14 patients, and (b) box-plot diagram indicates the distribution (the median, maximum, minimum, and interquartile range) of patient size in terms of $D_{w,g}$ and $D_{w,l}$. The $D_{w,l}$ gives greater values than the $D_{w,g}$ ($p$-value = 0.0012).

Figure 4: (a) Estimated eye lens doses radiation (CTDI$_{vol}$ volume computed tomography dose index, $SSDE_{n,g}$ average of global size-specific dose estimate based on new report of the American Association of Physicists in Medicine (AAPM) No. 293, $SSDE_{o,g}$ average of global size-specific dose estimate based on previously report of the AAPM No. 220, $SSDE_{n,l}$: average of local size-specific dose estimate based on new report of AAPM No. 293, $SSDE_{o,l}$: average of local size-specific dose estimate based on report of the AAPM No. 220) and measured doses using the thermo-luminescent dosimeters (TLDs), and (b) the dose distributions of the six descriptors in a box-plot diagram. $SSDE_{o,l}$ gave a more closely match estimation of eye lens doses to the measured doses.
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Table 1: Percentage differences and p-values of eye radiation dose between the average of local size specific dose estimate (SSDE), average global SSDE, and volume computed tomography dose index (CTDI\(_{vol}\)) and the measured doses by the thermo-luminescent dosimeters (TLDs).

| Metrics          | Percentage differences (%) | p-values         |
|------------------|-----------------------------|------------------|
| CTDI\(_{vol}\)   | 22.5 ± 8.9                  | < 0.001          |
| SSDE\(_{o.g}\)   | 21.7 ± 5.3                  | < 0.001          |
| SSDE\(_{o.l}\)   | 15.0 ± 5.7                  | < 0.001          |
| SSDE\(_{n.g}\)   | 9.3 ± 4.3                   | 0.001            |
| SSDE\(_{n.l}\)   | 2.1 ± 4.2                   | 0.566 (> 0.05)   |

CTDI\(_{vol}\): Computed tomography dose index; SSDE\(_{o.g}\): Average of global size-specific dose estimate based on new report of the American Association of Physicists in Medicine (AAPM) No. 293; SSDE\(_{o.l}\): Average of global size-specific dose estimate based on previously report of the AAPM No. 220; SSDE\(_{n.g}\): Average of local size-specific dose estimate based on new report of AAPM No. 293; SSDE\(_{n.l}\): Average of local size-specific dose estimate based on report of the AAPM No. 220.

The TLD measurements were about 16% [28]. However, size-conversion factors provided in that report were specifically intended for the abdominal-pelvic region. In this current study, we calculated the average of global and local SSDEs (denoted as SSDE\(_{o.g}\) and SSDE\(_{o.l}\)) using \(f_n\), the size-conversion factors for head CT examinations reported in AAPM Report No. 293.

The distributions of the six doses indicators observed in the current study are shown in Figure 4. The measured doses from calibrated TLDs were assigned as the gold standard. All patients had a CTDI\(_{vol}\) value greater than the TLD doses by 22.5 ± 8.9% that is consistent with the results reported by previous investigators [8, 16]. There was a statistically significant difference between CTDI\(_{vol}\) and TLDs doses (p-value < 0.001). The value of CTDI\(_{vol}\) was constant for all patients, independent of patient size, so that it only characterizes the output radiation dose from a scanner [31]. This is indicative of fixed tube current (TCM) scans.

We then calculated the SSDE in terms of SSDE\(_{o.g}\) and SSDE\(_{o.l}\). Similar trends to the TLD doses were obtained, but the percentage differences in both cases were relatively high (SSDE\(_{o.g}\) = 21.7 ± 5.3% and SSDE\(_{o.l}\) = 15.0 ± 5.7%). A statistically significant difference was also observed from the resulting p-value < 0.001. We then determined the values of SSDE\(_{n.g}\) and SSDE\(_{n.l}\). These metrics provided the closest estimated doses to the TLD measurements, viz. SSDE\(_{n.l}\) (2.1 ± 4.2%) rather than SSDE\(_{n.g}\) (9.3 ± 4.3%). SSDE\(_{n.l}\) did not show a statistically significant difference from the TLD doses (p-value = 0.566).

An accurate eye dose estimation in patients suspected of NPC is very important. One of the main treatments for patients with NPC is high-dose radiotherapy, and the eye is one of organs at risk. Therefore, calculating the dose received by the eye in the diagnostic stages (diagnostic CT scan and/or CT simulation) and adding it to the dose received in radiation therapy will cause a more accurate calculation of the total dose of the eye.

This current study employed a very limited number of patients. Although it provides proof of concept. A further study with a larger cohort will be needed. Although the current study is only focused on eye lens dose, other organ doses within the head region can be treated similarly using the proposed approach since the radial dose within the head region is relatively homogeneous [28]. Estimates of organ doses outside the head would require
the appropriate radial dose distribution within patient [26].

Conclusion
We estimated the eye lens dose using the average local SSDE within the eye location based on the size-conversion factors of AAPM Report No. 293 (SSDE_{n,l}). We compared these estimates with other metrics such as CTDI_{vol}, SSDE_{o,g}, SSDE_{o,l}, and SSDE_{n,g}. All the estimates were compared to measurements using TLDs. The differences from the TLDs were 22.5, 21.7, 15.0, 9.3, and 2.1% for CTDI_{vol}, SSDE_{o,g}, SSDE_{o,l}, and SSDE_{n,g}, respectively. Thus, the average of local SSDE calculated using size-conversion factors of AAPM Report No. 293 gives the highest accuracy for estimating eye lens radiation doses.

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Conflict of Interest
None

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