Diagnostic Reference Levels based on clinical indications in computed tomography: a literature review

Graciano Paulo 1*, John Damilakis 2, Virginia Tsapaki 3, Alexander A. Schegerer 4,5, Jacques Repussard 6, Werner Jaschke 7, Guy Frija 8 and European Society of Radiology 9

Abstract

Background: In August 2017, the European Commission awarded the “European Study on Clinical Diagnostic Reference levels for X-ray Medical Imaging” project to the European Society of Radiology, to provide up-to-date Diagnostic Reference Levels based on clinical indications. The aim of this work was to conduct an extensive literature review by analysing the most recent studies published and the data provided by the National Competent Authorities, to understand the current situation regarding Diagnostic Reference Levels based on clinical indications for computed tomography.

Results: The literature review has identified 23 papers with Diagnostic Reference Levels based on clinical indications for computed tomography from 15 countries; 12 of them from Europe. A total of 28 clinical indications for 6 anatomical areas (head, cervical spine/neck, chest, abdomen, abdomen-pelvis, chest-abdomen-pelvis) have been identified.

Conclusions: In all the six anatomical areas for which Diagnostic Reference Levels based on clinical indications were found, a huge variation of computed tomography dose descriptor values was identified, providing evidence for a need to develop strategies to standardise and optimise computed tomography protocols.

Keywords: Diagnostic Reference Levels, Computed tomography, Clinical indications, Computed tomography dose descriptors

Key points

- The establishment, regular review and use of Diagnostic Reference Levels are mandatory according to the Council Directive 2013/59/EURATOM.
- Most of the existing Diagnostic Reference Levels have been established based on anatomical locations, which has some limitations as one could have several clinical indications with consequently different protocols corresponding to different exposure levels.
- In the anatomical areas for which Diagnostic Reference Levels based on clinical indications were found, a huge variation of computed tomography dose descriptors values has been identified.
- The EUCLID project aims to establish Diagnostic Reference Levels based on clinical indications.

Background

The concept of Diagnostic Reference Levels (DRLs) was introduced many years ago by the International Commission on Radiological Protection (ICRP) [1] and has been widely accepted as a practical tool for optimisation in diagnostic and interventional radiology and nuclear
medicine. DRLs should be used as a form of investigation level to identify unusually high dose levels. If DRLs are consistently exceeded, a local review usually takes place. DRLs are not intended for regulatory or commercial purposes, nor do they represent a dose constraint, nor are they linked to limits or constraints [2].

The European Union has formally introduced the concept and the mandatory use of DRLs in every Member State since 1997 [3], reinforcing the obligation for the establishment, regular review and use in 2013 through the Council Directive 2013/59/EURATOM (BSSD), on health protection of individuals against the dangers of ionising radiation in relation to medical exposure [4].

Most of the existing DRLs (independently of the imaging modality) have been established based on anatomical locations. However, some limitations of this approach were pointed out for computed tomography (CT) as, for the same anatomical location, one could have several clinical indications with consequently different protocols corresponding to different exposure levels. For example, chest CT could correspond to the work-up for pulmonary embolism, lung cancer, or even coronary calcium scoring, each of which requires corresponding image quality parameters and scan length, and hence should have different DRLs [5].

The clinical approach to DRLs was mentioned some years ago by the ICRP [6], but most of the European National Competent Authorities (NCAs) still consider DRLs for anatomical location and not for clinical indication. However, some countries have recently established DRL based on clinical indications (DRLci) and some others are planning to do so in the near future. Also the European Society of Radiology (ESR) EuroSafe Imaging Call for Action 2018 has defined the objective to develop DRLci for adults and children, under action number 2 [7].

In this work, the dose descriptors used to define a DRL in CT are (a) volume computed tomography dose index (CTDIvol), the standard descriptor for estimating the output dose of a CT scanner, based on measurements obtained when scanning either a 16 cm or 32 cm phantom [8] and the unit used is mGy; (b) dose length product total (DLP), which is the sum of the DLP values from each CT acquisition/phase, representing the measure of the total amount of radiation used to perform the CT examination. DLP is the product of the CTDIvol (mGy) and scan length (cm), and the unit used is mGy.cm.

Both CTDIvol and DLP, are essential tools for CT optimisation; however it is important to understand the fact that they only represent CT scanner output and consequently are not patient dose estimates [9].

In August 2017, the European Commission (EC) launched the “European Study on Clinical Diagnostic Reference Levels for X-ray Medical Imaging” (EUCLID) project, to provide up-to-date DRLci.

The main objectives of the EUCLID project, led by the ESR, were to conduct a European survey to collect data needed for the establishment of DRLci for the most important X-ray imaging tasks in Europe (from the radiation protection perspective) and to specify up-to-date DRLci for those examinations.

The aim of this work was to conduct an extensive literature review by analysing the most recent studies published and the data provided by NCAs, to understand the current situation regarding DRLci for CT, under the scope of EUCLID project.

Materials and methods

One of the goals of EUCLID was the collection of information on the status of national DRLs and DRLci in Europe from NCAs from literature and from a workshop held in December 2019 in Luxembourg. The methodology for this included contacting the NCAs of 31 European countries and asking them to provide available national data that was then discussed and confirmed during the aforementioned workshop. Additionally, a comprehensive literature review was undertaken in order to identify which clinical indications had already been specifically studied.

To perform the literature review, several databases were used, such as science direct, PubMed and Google Scholar. Multiple keywords combination was used, such as diagnostic reference levels in computed tomography, clinical diagnostic reference levels and diagnostic reference levels based on clinical indications. All publications were collected and stored in the Mendeley reference management software (www.mendeley.com).

Results

Literature review for DRLci in CT

By using the keywords defined, data from 65 papers was considered and amongst them 23 included DRLci, being that 3 of them were from countries outside Europe (United States of America, Japan and Egypt) and 12 from European countries: Austria, Denmark, Finland, France, Germany, Ireland, Italy, Norway, Sweden, Switzerland, The Netherlands, United Kingdom. In addition, data provided by the NCAs, discussed and validated during the workshop, were also included.

Considering that the concept of DRLci is a recent one, some discrepancy and inconsistency was found in the classification of the clinical indication.

The DRLci values found were for several anatomical areas and are listed in Table 1. A total of 28 clinical indications for 6 anatomical areas have been identified. The anatomical areas with the most values for DRLci were “head” and “abdomen”, with a total of 6 each.
Head CT
For head CT, 10 references with DRL\textsubscript{ci} were found for 6 clinical indications: acute stroke; haemorrhage/aneurysms/arteriovenous malformations; metastases/cerebral abscess; trauma; cholesteatoma; sinusitis. Table 2 shows the DRL\textsubscript{ci} for head CT with the CTDI\textsubscript{vol} and/or DLP values for each clinical indication. The DRL\textsubscript{ci} for trauma/sinusitis was the clinical indication with the most references found (7 out of 10). The DLP values ranged from 90 mGy.cm \cite{14} to 1000 mGy.cm \cite{18}. One publication presents DRL\textsubscript{ci} for head CT both for males and females \cite{16}, demonstrating however similar or in some cases equal values.

Cervical (spine and neck) CT
For cervical (spine and neck) CT, eight references with DRL\textsubscript{ci} were found for three clinical indications: fracture, disk pathology and adenopathy/abscess. Table 3 shows the DRL\textsubscript{ci} for cervical (spine and neck) CT with the CTDI\textsubscript{vol} and/or DLP values for each clinical indication. The DRL\textsubscript{ci} for fracture was the clinical indication with more references found (six out of eight). The DLP values ranged from 300 mGy.cm \cite{18} to 640 mGy.cm \cite{15}. One publication presents DRL\textsubscript{ci} for head CT both for males and females \cite{16}, however, with similar values. Two publications from the same country, one from 2016 \cite{11} and other from 2018 \cite{19}, show a reduction of DLP values from 600 mGy.cm to 440 mGy.cm for the same clinical indication “fracture”.

Chest CT
For chest CT, 23 references with DRL\textsubscript{ci} were found for 6 clinical indications: lung cancer, interstitial lung disease, pulmonary embolism, coronary computed tomography angiography (CCTA), calcium scoring. Table 4 shows the DRL\textsubscript{ci} for chest CT with the CTDI\textsubscript{vol} and/or DLP values for each clinical indication.
| References | Acute stroke/post fossa | Acute stroke/cerebrum | Acute stroke/brain (whole) | Acute stroke/all sequences | Haemorrhage, aneurysms, arteriovenous malformations | Metastases, cerebral abscess | Trauma, sinusitis | Cholesteatoma | Sinusitis |
|------------|-------------------------|-----------------------|---------------------------|---------------------------|--------------------------------------------------|-----------------------------|----------------|-------------|----------|
|            | CTDIvol (mGy) | DLP (mGy.cm) | CTDIvol (mGy) | DLP (mGy.cm) | CTDIvol (mGy) | DLP (mGy.cm) | CTDIvol (mGy) | DLP (mGy.cm) | CTDIvol (mGy) | DLP (mGy.cm) | CTDIvol (mGy) | DLP (mGy.cm) | CTDIvol (mGy) | DLP (mGy.cm) | CTDIvol (mGy) | DLP (mGy.cm) | CTDIvol (mGy) | DLP (mGy.cm) |
| Danish Health Authority (DK) 2015 [10] | – | – | – | – | – | – | – | – | – | – | – | – | – | – | – | – | – | – |
| Public Health England (UK) 2016 [11] | 80 | – | 60 | – | 60 | – | – | 970 | – | – | – | – | – | – | – | – | – | – | – |
| Schegerer et al. (DE) 2017 [12] | – | – | – | – | – | – | – | – | 9 | 120 | – | – | – | – | – | – | – | – | – |
| Treier et al. (CH) 2010 (22) | – | – | – | – | – | – | – | 65 | 1000 | 65 | 1000 | 25 | 350 | 50 | 250 | – | – | – | – |
| Van der Molen et al. (NL) 2013 [13] | – | – | – | – | – | – | – | 936 | – | – | – | 133 | – | – | – | – | – | – | – | – |
| Wachabauer et al. (AT) 2017 [14] | – | – | – | – | – | – | – | – | – | – | 90 | – | – | – | – | – | – | – | – |
| Geryes et al. (FR) 2019 [15] | – | – | – | – | – | – | – | 44 | 1010 | 44 | 790 | 43 | 920 | – | – | – | – | – | – |
| Ireland (IE) MERU 2017 [16] | 26 (a) | 469 (a) | – | – | – | – | – | – | – | – | 62 (a) | 918 (a) | – | – | 21 (a) | 183 (a) | – | – | – | – |
| Norway (NO) 2018 [17] | 31 (b) | 477 (b) | – | – | – | – | – | – | – | – | 64 (b) | 927 (b) | – | – | 21 (b) | 210 (b) | – | – | – | – |
| Sweden (SE) 2019 [18] | – | – | – | – | – | – | 60 | 950 | 60 | 950 | – | – | – | – | – | – | – | – | – | – |

*aFor female patients
*bFor male patients
values for each clinical indication. The DRL$_{ci}$ for CCTA was the clinical indication with more references found (11 out of 23). The DLP values ranged from 170 mGy.cm [12] to 1400 mGy.cm [22]. One publication presents DRL$_{ci}$ for CCTA [12] made with three different approaches: prospective, no padding, 170 mGy.cm (c); prospective, with padding, 280 mGy.cm (d); prospective, with gating, 380 mGy.cm (e). From the three approaches, the prospective, no padding technique is the one that provides the lowest DLP value (170 mGy.cm).

Abdominal CT
For abdominal CT, 11 references with DRL$_{ci}$ were found for 6 clinical indications: liver metastasis, abscess, kidney stones/colic, kidney tumour/colic, acute abdomen and pancreas adenocarcinoma. Table 5 shows the DRL$_{ci}$ for abdominal CT with the CTDI vol and/or DLP values for each clinical indication. The DRL$_{ci}$ for liver metastasis and kidney stone/colic were the clinical indications with more references found (6 out of 12). For liver metastasis, the DLP values ranged from 400 mGy.cm [15, 19] to 1423 mGy.cm [27]. For kidney stone/colic, the DLP values ranged from 280 mGy.cm [33] to 600 mGy.cm [18]. One publication presents DRL$_{ci}$ for abdominal CT, both for males and females [16], demonstrating however similar values.

Abdominopelvic CT
For abdominopelvic CT, five references with DRL$_{ci}$ were found for five clinical indications: abscess/lymphadenopathy, virtual colonoscopy (VC)/polyps/tumour, CT for abdominal aortic aneurysms (AAA), colic and occlusion. Table 6 shows the DRL$_{ci}$ for abdominopelvic CT with the CTDI vol and/or DLP values for each clinical indication. The DRL$_{ci}$ for abscess/lymphadenopathy was the clinical indication with more references found (four out of five). The DLP values ranged from 650 mGy.cm [11, 15, 34] to 750 mGy.cm [18].

Chest-abdominopelvic CT
For chest-abdominopelvic CT, three references with DRL$_{ci}$ were found for three clinical indications: tumour, infectious and oncologic follow-up. Table 7 shows the DRL$_{ci}$ for chest-abdominopelvic CT with the CTDI vol and/or DLP values for each clinical indication. The DRL$_{ci}$ for tumour and oncologic follow-up were the clinical indications with more references found (two out of three). For tumour, the DLP values ranged from 870 mGy.cm [15] to 950 mGy.cm [35]. For oncologic follow-up, the DLP values ranged from 605 mGy.cm [36] to 970 mGy.cm [34]. One publication presents DRL$_{ci}$ for abdominal CT, both for males and females [16], demonstrating however similar values.

Discussion
To our knowledge, this is the first article to perform literature review for DRL$_{ci}$. Considering that DRL$_{ci}$ is a recent concept, it is understandable that only a limited number of papers was found in the literature and most of the proposed DRL$_{ci}$ came from the NCAs of 12 European countries.

In all the six anatomical areas where DRL$_{ci}$ were found, a huge variation of CT dose descriptors values was identified, providing evidence that different approaches/protocols are used to perform the CT procedure for the same clinical indication.

In the 28 clinical indications identified in the literature, the procedures with the highest differences in DLP values were head trauma (11-fold), CCTA (9-fold), liver metastasis (3-fold) and cervical fracture (2-fold).

The huge variations in the reported CT dose descriptors values for almost all the clinical indications

### Table 3 DRL$_{ci}$ for cervical CT

| References                              | Fracture CTDI$_{vol}$ (mGy) | DLP (mGy.cm) | Disk pathology CTDI$_{vol}$ (mGy) | DLP (mGy.cm) | Adenopathy, abscesses CTDI$_{vol}$ (mGy) | DLP (mGy.cm) |
|-----------------------------------------|----------------------------|-------------|-----------------------------------|-------------|----------------------------------------|-------------|
| Schegerer et al. (DE) 2019 [20]         | 20                        | –           | 25                                | –           | –                                      | –           |
| Public Health England (UK) 2016 [11]    | 26                        | 600         | –                                 | –           | –                                      | –           |
| Treier et al. (CH) 2010 [21]            | –                         | –           | –                                 | –           | 30                                     | 600         |
| Geryes et al. (FR) 2019 [15]            | 31                        | 640         | –                                 | –           | –                                      | –           |
| Ireland (IE) MERU 2017 [16]             | 26 (a)                    | 469 (a)     | –                                 | –           | –                                      | –           |
|                                       | 31 (b)                    | 477 (b)     | –                                 | –           | –                                      | –           |
| Norway (NO) 2018 [17]                   | 15                        | 350         | –                                 | –           | –                                      | –           |
| Sweden (SE) 2019 [18]                   | 13                        | 300         | –                                 | –           | 30                                     | 600         |
| Public Health England (UK) 2018 [19]    | 21                        | 440         | –                                 | –           | –                                      | –           |

*For female patients
*bFor male patients
addressed above are likely to be explained by differences in protocols (exposure parameters and scan length), type and age of scanner, number of acquisition series and, in the specific case of CCTA, by the option of performing either prospective or retrospective acquisitions. The same variations in radiation doses for CT across patients is described in the literature, and the reasons are primarily related on how CT scanners are used [34], the differences in patient’s size (weight and height) [36] and to the level of image quality required to answer the clinical question [35]. Although the DRLs are defined for standard patients [4], taking into consideration that the weight and height of patients are also a determining factor.

Table 4 DRLci for chest CT

| References | Lung cancer | Interstitial lung disease (axial) | Interstitial lung disease (helical) | Pulmonary embolism | CCTA | Calcium scoring |
|------------|-------------|----------------------------------|------------------------------------|-------------------|------|----------------|
|            | ctdi vol (mgy) | dlp (mgy.cm) | ctdi vol (mgy) | dlp (mgy.cm) | ctdi vol (mgy) | dlp (mgy.cm) | ctdi vol (mgy) | dlp (mgy.cm) | ctdi vol (mgy) | dlp (mgy.cm) | ctdi vol (mgy) | dlp (mgy.cm) |
| Castellano et al. (UK) 2017 [22] | – | – | – | – | – | – | 173 | – | – | – | – | – |
| Danish Health Authority (DK) 2015 [10] | 16 | 620 | – | – | 13 | 500 | – | – | 29 | 230 | – | – | – |
| Foley et al. (IE) 2012 [23] | – | – | 7 | 276 | – | – | 13 | 432 | – | – | – | – | – |
| Fukushima et al. (JP) 2012 [24] | – | – | – | – | – | – | – | – | – | – | – | – | – |
| Schegerer et al. (DE) 2019 [20] | – | – | – | – | – | – | – | – | – | – | – | – | – |
| Hausleiter et al. 2009 [25] | – | – | – | – | – | – | 696 | 1152 | – | – | – | – | – |
| Japan Network for Research on Medical Exposures (JP) 2015 [26] | – | – | – | – | – | – | 90 | 1400 | – | – | – | – | – |
| Kanai et al. (USA) 2017 [27] | – | – | – | – | – | – | 19 | 557 | – | – | – | – | – |
| Mafalanka et al. (FR) 2015 [28] | – | – | – | – | – | – | – | – | – | 870 | – | – | – |
| Palorini et al. (IT) 2014 [29] | – | – | – | – | – | – | – | – | – | 1208 | – | 131 | – |
| Public Health England (UK) 2016 [11] | 12 | 610 | 4 | 140 | 12 | 350 | 13 | 440 | – | – | – | – | – |
| Radiation and Nuclear Safety Authority (FI) 2013 [30] | 11 | 430 | – | – | – | – | – | – | – | – | – | – | – |
| Salama et al. (EG) 2017 [31] | – | 22 | 421 | – | – | – | – | – | – | – | – | – | – |
| Schegerer et al. (DE) 2017 [12] | – | – | – | – | – | – | 15 | 300 | 36 | 551 | 19 | 270 | 8 | 119 |
| Treier et al. (CH) 2010 [21] | – | – | – | – | – | – | – | – | – | 1000 | – | 150 | – |
| Van der Molen et al. (NL) 2013 [13] | – | – | 276 | – | 371 | – | 671 | – | 51 | – | – | – | – |
| Wachabauer et al. al (AT) 2017 [14] | – | – | – | – | – | – | 400 | – | – | – | – | – | – |
| Geryes et al. (FR) 2019 [15] | – | 7 (a) | 241 (a) | – | 8 | 310 | – | – | – | – | – | – | – |
| Ireland (IE) MERU 2017 [16] | 7 (b) | 272 (b) | 7 (b) | 249 (b) | 12 (b) | 278 (b) | – | – | – | – | – | – | – |
| Norway (NO) 2018 [17] | 9 | 350 | – | 9 | 300 | – | – | – | – | – | – | – | – |
| Sweden (SE) 2019 [18] | 9 | 350 | – | – | – | – | – | – | – | – | – | – | – |
| Public Health England (UK) 2018 [19] | – | – | – | – | – | – | 170 (c) | – | – | – | – | – | – |
| Netherlands (NL) 2012 [32] | – | – | – | – | 10 | 350 | – | – | – | – | – | – | – |

*aFor female patients
*bFor male patients
*cProspective, no padding
*dProspective, with padding
*eRetrospective, with gating
for dose increase, categorising patients by body mass index should be considered in the near future [36].

Several other factors may also contribute to the heterogeneity of results shown in the DRLci tables. DLP values may refer to individual sequences or to a complete examination (total DLP), and in some cases, this information is not included in the paper/report.

In addition, different names have been used for what is likely to have been the same indication (e.g. abscess versus acute abdomen), and the question of whether these differences are related to various interpretations of the name of the clinical indication or to different practices remains open. A semantic refinement, with the precise description of the clinical indication, should be made in the future in order to minimise any variation related to the meaning of the clinical indication.

For liver metastases and a few other clinical indications, DRLci in terms of CTDIvol are similar, but DRLs in terms of DLP differ considerably. The difference between results in values of total DLP (yet similar levels of

Table 5 DRLci for abdominal CT

| Reference | Liver metastases | Abscess | Kidney stones/ colic | Kidney tumour/ colic | Acute abdomen Ca | Pancreas adeno Ca |
|-----------|------------------|---------|----------------------|----------------------|-----------------|------------------|
|           | CTDIvol (mGy)   | DLP (mGy.cm) | CTDIvol (mGy) | DLP (mGy.cm) | CTDIvol (mGy) | DLP (mGy.cm) | CTDIvol (mGy) | DLP (mGy.cm) | CTDIvol (mGy) | DLP (mGy.cm) |
| Danish Health Authority (DK) 2015 [10] | – | – | – | – | 17 | 700 | – | – | – | – |
| Public Health England (UK) 2016 [11] | 14 | 910 | 15 | 745 | 10 | 460 | 13 | 1150 | – | – | – | – |
| Radiation and Nuclear Safety Authority (FI) 2013 [30] | – | – | – | 7 | 330 | – | – | – | – | – | – |
| Salama et al. (EG) 2017 [31] | 31 | 1423 | – | – | – | – | – | – | – | – | – |
| Treier et al. (CH) 2010 [21] | 15 | 400 | – | – | – | – | – | – | – | – | – |
| Van der Molen et al. (NL) 2013 [13] | – | – | – | 329 | – | 1371 | – | – | – | 1000 |
| Wachabauer et al. (AT) 2017 [14] | – | 400 | – | – | – | – | – | – | – | – | – |
| Ireland (IE) MERU 2017 [16] | 9 (a) | 554 (a) | – | – | 6 (a) | 254 (a) | – | – | – | – | – | – |
| | 10 (b) | 515 (b) | – | – | 8 (b) | 291 (b) | – | – | – | – | – | – |
| Norway (NO) 2018 [17] | – | – | – | 5 | 250 | 13 | 1300 | – | – | – | – |
| Sweden (SE) 2019 [18] | 11 | 550 | – | 5 | 200 | 12 | 1000 | – | – | – | – |
| Netherlands (NL) 2012 [32] | – | – | – | – | – | – | 15 | 700 | – | – | – | – |

*For female patients
*For male patients

Table 6 DRLci for abdomino-pelvis CT

| Abdomino-pelvis CT | Abscess lymphadenopathy | VC-polyps/tumour CT angiography (AAA) | Colic | Occlusion |
|-------------------|-------------------------|--------------------------------------|--------|----------|
| CTDIvol (mGy)   | DLP (mGy.cm) | CTDIvol (mGy) | DLP (mGy.cm) | CTDIvol (mGy) | DLP (mGy.cm) | CTDIvol (mGy) | DLP (mGy.cm) |
| Public Health England (UK) 2016 [11] | 15 | 745 | 11 | 950 | – | – | – | – |
| Treier et al. (CH) 2010 [21] | 15 | 650 | – | – | 15 | 650 | – | – | – | – |
| Van der Molen et al. (NL) 2013 [13] | – | – | – | – | – | 727 | – | – | – | – |
| Wachabauer et al. (AT) 2017 [14] | – | 650 | – | – | – | – | – | – | – | – |
| Geryes et al. (FR) 2019 [15] | – | 650 | – | – | – | – | 8 | 400 | 12 | 880 |
CTDIvol) for examinations of the lower trunk could be a consequence of the present use of increased scan lengths and/or number of sequences (particularly in relation to imaging for different phases in the distribution of contrast medium). The substantial variations in CT protocols, for the same clinical indication, delivers several folds higher radiation than necessary [33].

Although a large number of research studies have shown that dose optimisation tools such as tube current modulation can reduce patient dose considerably, it is not known how these tools are being used in everyday clinical practice. Large differences in dose descriptors for the same clinical indication and, sometimes, for the same CT scanner model may be addressed by standardising acquisition protocols, using dose reduction tools properly and improving education of practitioners in medical radiation protection.

Conclusions

From this literature review, it is obvious that there is a lot of space for improvement in terms of standardising the CT protocols for each clinical indication and that the development of European guidelines on this topic would be very useful as a tool to implement dose reduction strategies in CT procedures.

Continuing to develop DRLs for CT based in anatomical areas without taking into consideration the clinical indication will probably meet the minimum standard of the BSSD but will insufficiently contribute to fulfil the main purpose of the existence of DRLs: a tool for optimisation.

We expect that the results of this work can stimulate the radiological community and the NCAs to move toward the establishment of DRLci in a more harmonised and consistent way.

Abbreviations

BSSD: Council Directive 2013/59/EURATOM; CT: Computed tomography; CTDIvol: Volume computed tomography dose index; DLP: Dose length product total; DRLci: DRL based on clinical indications; DRLs: Diagnostic Reference Levels; EC: European Commission; ESR: European Society of Radiology; EUCLID: European Study on Clinical Diagnostic Reference Levels for X-ray medical Imaging; ICRP: International Commission on Radiological Protection; NCAs: National Competent Authorities

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Authors’ contributions

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Author details

1ESTESC - Coimbra Health School, Medical Imaging and Radiotherapy Department, Instituto Politécnico de Coimbra, Rua 5 de Outubro, S. Martinho do Bispo, 3046-854 Coimbra, Portugal. 2School of Medicine, University of Crete, Iraklion, Crete, Greece. 3Konstantopoulio General Hospital, Athens, Greece. 4Department of Medical and Occupational Radiation Protection, Federal Office for Radiation Protection, Neuenheim, Germany. 5Radiation Protection and Image Processing Systems, Hirslanden AG, Glattpark, Switzerland. 6International Consultant, St Norn la Breêche, France. 7Department of Radiology, Medical University Innsbruck, Innsbruck, Austria. 8European Society of Radiology, Vienna, Austria.

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