Dual-energy CT tissue segmentation methods for Monte Carlo dose calculations in proton therapy

Citation for published version (APA):

Document status and date:
Published: 01/01/2019

DOI:
10.26481/dis.20190522ia

Document Version:
Publisher's PDF, also known as Version of record

Please check the document version of this publication:

• A submitted manuscript is the version of the article upon submission and before peer-review. There can be important differences between the submitted version and the official published version of record. People interested in the research are advised to contact the author for the final version of the publication, or visit the DOI to the publisher’s website.
• The final author version and the galley proof are versions of the publication after peer review.
• The final published version features the final layout of the paper including the volume, issue and page numbers.

Link to publication

General rights
Copyright and moral rights for the publications made accessible in the public portal are retained by the authors and/or other copyright owners and it is a condition of accessing publications that users recognise and abide by the legal requirements associated with these rights.

• Users may download and print one copy of any publication from the public portal for the purpose of private study or research.
• You may not further distribute the material or use it for any profit-making activity or commercial gain
• You may freely distribute the URL identifying the publication in the public portal.

If the publication is distributed under the terms of Article 25fa of the Dutch Copyright Act, indicated by the “Taverne” license above, please follow below link for the End User Agreement:
www.umlib.nl/taverne-license

Take down policy
If you believe that this document breaches copyright please contact us at:
repository@maastrichtuniversity.nl
providing details and we will investigate your claim.

Download date: 13 Sep. 2020
7.1 Summary and discussion
Dual-energy computed tomography (DECT) is currently an important topic of research for several applications in radiotherapy. For proton therapy, DECT is studied particularly to improve range prediction and tissue segmentation, to extract tissue compositions, to predict prompt-gamma and positron emission, and to move towards automated organ delineation. Understanding the advantages and current limitations of DECT technology is crucial for clinical implementation, which we believe will happen at many centers in the near future.

Due to the wide choice of DECT technologies available on the market, such as dual-source and twin-beam, this thesis started in Chapter 2 by quantifying and extracting, for these two different DECT systems, tissue-specific physical properties: the relative electron density ($\Omega_e$) and the effective atomic number ($Z_{\text{eff}}$) following the methods from Saito [1] and Landry et al. [2] respectively. In a quantification point-of-view, DECT benefits immensely with a system with a wide spectral separation (within the clinical range of energies), whereas a twin-beam technology with two strongly overlapping low- and high-energy spectra with the same maximum photon energy produces similar images and results in higher discrepancy between reference and calculated values for $\Omega_e$ and $Z_{\text{eff}}$ in phantom tests. For this reason, the dual-source system with the additional filtration in the high-energy x-ray source has the potential of yielding the lowest root-mean-square (RMS) errors for these physical properties for both third, but also, second generation detectors. This was also explored in Chapter 2 of this thesis, comparing two dual-source systems with different detectors.

However, the current commercially available dual-source DECT systems have a fundamental physical limitation on the size of the detectors placed inside the scanner structure, resulting in a combined field-of-view (FOV) of maximum 35 cm diameter. In the external-beam radiotherapy field, the complete image of a patient is required depending on the tumour location, which would not fit this limited FOV. The dual-spiral technology with a single x-ray tube and imaging detector appears as the compromise between a system with a good spectral separation (although no extra filtration) and the capability of producing images with the typical 50 cm FOV of SECT scanners. We studied in detail the first CT scanner specifically manufactured for radiotherapy that includes a DECT dual-spiral capability (Chapter 3). In this chapter, we extracted the $Z_{\text{eff}}$ from DECT with two purposes: assign a material per voxel using the two-dimensional [$\rho_e$,$Z_{\text{eff}}$]-space, and extract the mean excitation potential ($I$) necessary for collision stopping power calculation using the method developed by Yang et al. [3]. Additionally to analyzing this radiotherapy specific dual-spiral technology and comparing it to the twin-beam and dual-source systems, a comparison with the SECT methodology was added to complement the results of this work and investigate relevant differences in range.

A complete framework was developed to use DECT images in Monte Carlo (MC) proton dose calculations, which will be discussed further in this chapter. Two main conclusions resulted from this work: first, the dual-spiral yielded results as good as the dual-source in terms of tissue segmentation and range prediction (defined as the 80% distal fall off, R80) for the non-moving phantoms used; second, absolute differences in R80 for a single-field head & neck patient case between the DECT dual-spiral and the SECT-stoichiometric based dose calculations yielded a mean value of 1.2 ± 1.2 mm. The later was consistent
with the work presented by Wohlfahrt et al. [4] reporting a mean water-equivalent range shift of -1.1 mm between a SECT and DECT dual-spiral plans, and Hudobivnik et al. [5] who found mean differences of -1.7 mm between SECT and DECT dual-source plans, also for a head case.

Meanwhile, radiotherapy clinics and hospitals rely on treatment planning systems (TPS) to perform dose calculations based on CT images, which currently are not capable of handling a set of two DECT images as input and lack the necessary tools to extract from these the different tissue properties. Since these systems rely on a single-input, a pseudo-monoenergetic image (PMI), presents itself as a hybrid solution: created from a weighted sum of the two DECT acquisitions \(\alpha\text{CT}_{\text{low}} + (1-\alpha)\text{CT}_{\text{high}}\), but using the same conversion methodology from CT# to SPR or \(\rho\) as the conventional SECT. The choice of \(\alpha\)-parameter can be optimized depending on the application: e.g. a higher contrast-to-noise ratio and edge enhancement can decrease inter-observer variability for contouring, while a smoother image with minimal beam-hardening effect can be used for SPR or \(\rho\) calculation. A 79keV-PMI was successfully implemented as clinical standard in the Universitäts Protonen Therapie Dresden for all body sites except lung and head & neck, basing this decision on the results presented by Wohlfahrt et al. [4, 6, 7] that compared 79keV-PMI- and SECT-based SPR prediction in a anthropomorphic ground-truth phantom and later evaluated the differences in a cohort of head and pelvic patients. Nevertheless, the 79keV-PMI is not able to achieve the accuracy in SPR of the full DECT based methods.

Scanner manufacturers have also been developing their own software to facilitate the radiotherapy work-flow. One example was the launching of the DirectDensity™ algorithm by Siemens Healthineers (Forchheim, Germany) in 2016, which reconstructs an image directly in scaled relative electron densities. Another product of Siemens Healthineers is the RHO/Z package from their own syngo.via software, that calculates \(\rho_e\) and \(Z_{\text{eff}}\) images from the two DECT scans.

At our institute, we are supplied with two DECT scanners (a dual-source SOMATOM Definition Drive and a dual-spiral SOMATOM Definition Confidence© RT pro), the latest software version for each scanner with the DirectDensity™ algorithm, and RHO/Z package in syngo.via. The large offer of clinically available methods of extracting \(\rho_e\) led to the on-going study in Chapter 4. Additionally, since Saito's [1] method has been recurrently shown to yield the lowest RMS for \(\rho_e\) extraction, its implementation was also added to this study, applying an additional post-processing denoising algorithm with fifteen iterations (called Saito-15it). Five \(\rho_e\)-extraction methods (CT2RED, DirectDensity™, PMI2RED, syngo.via-RHO, and Saito-15it) were tested first in two phantoms and later applied to a neurological patient scanned with both DECT and SECT protocols. The \(\rho_e\)-images for the patient case were converted to \(\rho\)-images (detailed explanation in section 4.2.6), so this could be used directly as input in the TPS. The \(\rho\)-images for the patient computed from the CT2RED, DirectDensity™ and PMI2RED methods (all methods based in a look-up-table conversion) yielded very similar \(\rho\) values, with differences within 0.005 ± 0.023 g/cm³. The two full DECT-based methods (syngo.via-RHO and Saito-15it), that start from the same equation, although following different post-processing algorithms, as discussed closely with Siemens and Saito (Appendix A), yielded higher discrepancies compared to the other three methods. Notably, opposite values where observed for the bone tissues, which was
explained after an direct correspondence with Siemens, and it was due to the fact that the syngo.via-RHO software used was not optimized for the scanner at our institution. Additionally, the effect of using the different $\rho$-images for proton dose calculations was quantified in terms of dose and range differences, where, for instance, differences of 1.5 ± 0.5 mm in R80 between CT2RED and syngo.via-RHO were found in a single patient example. We are continuing this study by adding a considerable number of patients to investigate if the variability between methods is seen in other cases and, up to now, the same trend is observed in the $\rho$-images.

**Chapters 3 and 4** confirm that proton dose calculations are sensitive to the CT modality (single or dual-energy), the DECT technology, and even the post processing algorithm used to compute the tissue properties. In this respect, we believe that MC techniques play an important role in decreasing the systematic uncertainty and in scrutinizing the numerous possibilities available.

A full framework was developed to use DECT images to create the necessary material-, $\rho$- and I-maps needed as input for Geant4 MC dose calculations. We wanted to investigate if DECT combined with a MC dose engine would yield a higher accuracy in terms of proton range compared to a material-based ground-truth simulated phantom than SECT conventional methodology. The hypothesis was confirmed and it is presented as one of the main results of Chapter 3. Furthermore, a plan calculated on the patient geometry had the individual pencil beams optimized with MC instead of using the pencil beam doses calculated by the analytical pencil beam algorithm. In this way, we account for the effect of inhomogeneities in the proton beam path.

These studies on the accuracy of DECT were especially relevant in our radiotherapy clinic, since its recent construction of the Zuid-Oost Nederland Protonen Therapie Centrum (ZONPTC), adding proton therapy as another available treatment modality. Our institution will start to treat patients either with intensity modulated radiotherapy (IMRT) or with scanned proton beam therapy. The choice of radiotherapy modality (IMRT or proton therapy) will be made according to a *model-based approach* from official Dutch scientific and health care governance bodies, which uses different normal tissue complication probability models to calculate the tissue-toxicity predicted for each treatment and, with these results and based on some thresholds, selects the modality that most benefits the patient [8]. For this reason, every patient is scanned with a single imaging protocol (depending on the body site), which has to suit both radiotherapy modalities.

ZONPTC decided to acquire the MEVION S250i with HYPERSCAN compact proton system (Mevion Medical Systems Inc., Littleton, MA) with the Adaptive Aperture™ (AA) beam collimation system. Aiming to have an independent dose (re)calculation engine, the MC beam model of this system was developed and it is described in detail in Chapter 5, showing a very good agreement for the integrated depth dose curves and the spot dose profiles for fifteen selected energies, for a single 10 x 10 cm² field at the isocenter and a 3D geometry tested with and without the AA. We are already partially integrating the framework developed in Chapter 3 with the beam model of our center, in particular, the implementation of DECT maps, so that DECT and SECT-based simulations can be performed and, additionally, compared with experiments measured in-house.
The AA component of our system makes it theoretically possible to create very small fields, either for treatment of small lesions (Appendix C), or even for pre-clinical in vivo small animal research. A first proof of concept to show the feasibility of performing pre-clinical research using the S250i with HYPERSCAN system was done in silico and it is presented in Chapter 6. This study helped us to understand the implications of degrading the nominal 230 MeV proton beam to very low energies and its effect on spot size and dose homogeneity, and also to investigate the system’s dose rate. We proved that this system can deliver doses of 2 Gy to a target volume of 27 mm$^3$ in under half a minute, achieving a good tumor coverage for several research scenarios.

7.2 Considerations and future perspectives

7.2.1 Advances in proton range prediction techniques

As shown for a subset of centers by the survey done by Taasti et al. [9], most particle therapy centers worldwide base their CT# to SPR (or $\Omega$) curve on one of two methods: interpolation between measured tissue mimicking plastics or CT# prediction by means of the stoichiometric calibration proposed by Schneider et al. [10]. For the latter, a scanner specific parameterized model is created based on the compositions of tissue equivalent materials and it is used to predict the CT# of tabulated human tissues.

A recent study in which we participated (Gomà et al.) looked carefully at the statements made in the 1996 publication from Schneider et al. and presented a critical review of the method [11]. First, for the stoichiometric calibration, Schneider claimed that a set of materials with known compositions, not necessarily biological tissue-like, could be used for the calibration and extraction of $k_1$ and $k_2$ fit-coefficients for the CT scanner model (equation A.20 from [11]). Gomà et al. shed new light on this by showing that the use of different phantoms, scanned with the same protocol and scanner, yields different fit results and, as consequence, different CT# prediction for the set of tabulated tissues (see table 2 from [11]). Secondly, the stoichiometric calibration seems to be a redundant step in converting CT# to $\rho_e$ when using an interpolation between the tissue mimicking inserts from the RMI 467 phantom (Gammex Inc. Middleton, WI), as shown in Figure 7.1. The same is not observed for the 062 M phantom, since it contains Barium in the elemental compositions of the bone inserts and this element is not present in biological tissues. These results were confirmed by an independent implementation of the stoichiometric method and using another set of CT images scanned at our institution with a different CT scanner, different protocols and reconstruction methods. On the other hand, the results published by Hansen et al. [12] do not reflect the same agreement between SPR extracted from the stoichiometric calibration versus the interpolation between the RMI 467 tissues.

Additionally, the list of human tissues used for the stoichiometric calibration and numerous other studies, reported in the International Commission on Radiation Units and Measurements (ICRU) and the International Commission on Radiological Protection (ICRP) publications, refer back to the early 50s, and have since been recompiled, republished and re-normalized by different sources. The true accuracy of these tabulated human tissues is unknown and it will have a direct impact on the accuracy of the CT# conversion to SPR, because mass-densities and compositions are used for SPR calculation methods for: