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To cite this version:
Igor Bessières, Bénédicte Poumarède, Jean-Marc Bordy. Development and experimental validation of TPS software to determine the out-of-field dose in radiotherapy beams. Third European Workshop on Monte Carlo Treatment Planning (EWG-MCTP 2012), May 2012, Sevilla, Spain. cea-02620216

HAL Id: cea-02620216
https://hal-kea.archives-ouvertes.fr/cea-02620216
Submitted on 25 May 2020

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Development and experimental validation of TPS software to determine the out-of-field dose in radiotherapy beams

Igor Bessières, Bénédicte Poumarède, Jean-Marc Bordy

CEA, LIST, Laboratoire Modélisation, Simulation et Systèmes, 91191 Gif-sur-Yvette CEDEX, France

igor.bessieres@cea.fr

I. INTRODUCTION

Even if the IMRT modality allows a more accurate definition of the target volume in radiotherapy, low doses are still delivered around the tumour to organs at risk. Epidemiological studies [1] demonstrated the relationship between peripheral doses and second cancers or heart diseases. Many experimental studies measuring the variation of the peripheral dose with the treatment parameters have been undertaken [2]. Up to now, very few studies have been performed on out-of-field Monte Carlo (MC) simulations. To our knowledge, MCNPX is the only one MC code that has been used to simulate the peripheral dose in radiotherapy [3]. Nowadays, there is thus no specific and accurate tool predicting the peripheral dose. Our aim is to develop a MC tool based on the PENELOPE code to compute the dose at the target volume and the organs at risk in order to enable a decrease of the peripheral dose by adapting the treatment's parameters. This tool will be implemented in a Treatment Planning System (TPS). In this paper we expose the first validation step of the out-of-field MC calculations, using a comparison with measurements in a specific large water tank.

II. MATERIALS AND METHODS

a. Experimental validation

i. Irradiation configuration

The irradiations have been performed at LNHB (French Primary Standard Laboratory) on a GE Saturne 43 linear accelerator for 6, 12 and 20 MV beam qualities. We followed the configuration of the IAEA 398 protocol which implies the use of a 10 × 10 cm² field, a skin surface distance (SSD) of 90 cm and a measurement at 10 cm depth in water. The only difference with the irradiation conditions proposed in this protocol is the use of a large water phantom (60 x 30 x 30 cm³) specially designed for this experiment. We considered a symmetric situation so only one side of the field has been explored.

ii. OSL dosimeter

Another aim of this experiment is to validate the use of Al₂O₃:C detectors (Nanodots, Landauer) and the optically stimulated luminescence (OSL) technique for out-of-field dose measurements. The OSL technique presents advantages over the thermoluminescent technique (TL): no sample heating is required and the detector can be read several times. Moreover, OSL measurements are relatively cheaper than TL measurements [5]. In this work, OSL dosimeters will be useful for the clinical validation of the code on an anthropomorphic phantom. Measurements performed with Nanodots have been compared to NE2571 ionization chambers (IC) measurements, considered as the reference data here.
Fig. 1: Energy response of the OSL in Air Kerma. The ratio $R/R_{C0}$ corresponds to the raw response of the reader divided by the raw response coming from the Cobalt.

Four main factors are applied to the raw reading of the reader: particular sensitivity for each Nanodot, the measured calibration factor, air calibration factor $D_{water}/K_{air}$ correcting the fact that the calibration has been done in air whereas the measurements have been performed in water and finally the energy dependence correction factor.

Indeed, we measured a high over-estimation of the dose (by a factor 3 or 4) for photon belonging to a low energy range ($< 100$ keV) [Fig. 1]. Consequently, we developed an energy dependence correction protocol. It combines the results of the experimental over-response curve with PENELOPE spectra calculations at measurement points. The influence of this correction is described in the next part.

b. Calculation tool

The tool we are developing is based on the 2006 release of PENELOPE [4]. This code has been parallelized to save computation time by running the calculations on a cluster of 372 processors. The simulations are run from specific large Phase Space File (PSF) recorded in a $60 \times 30$ cm$^2$ plane located at 90 cm from the source, after the jaws of the modelled accelerator. An additional comparison has been done using the tally F6 of the MCNPX MC code, the unique MC code used for this kind of studies until now. A specific feature of MCNPX is the availability of the DXTRAN variance reduction method. It is specially adapted to the calculation far from the primary beam and was tested in order to assess the gain to be expected on the out-of-field dose calculation efficiency.

III. RESULTS

a. MC comparison

As results obtained for 6, 12 and 20 MV quality beams are quite similar, we have reported on Fig. 2 the experimental and calculated out-of-field dose profiles obtained for 20 MV. These dose profiles have been normalized to the maximum of dose.
Fig. 2: Experimental and calculated 20 MV dose profiles

As expected, PENELOPE and MCNPX results in the field are in good agreement with OSL and IC measurements (error < 3%).

Out of the field, PENELOPE data are in good agreement with the IC measurements from the centre of the beam up to most distant points (40 cm). The statistical uncertainty associated to the most peripheral points is quite low (~15 %) regarding to the level of dose. The global mean error is about 3 % between these two sets of results.

Unlike the PENELOPE code, the MCNPX calculations over-estimate more and more the dose from 18 cm from the centre of the beam to the further positions. The global mean error is about 30 % between MCNPX and the IC values.

Consequently, compared to MCNPX, PENELOPE seems to be the best suited to simulate the out-of-field dose deposition with a high level of accuracy. To analyse the difference between the two MC codes, different cross-sections libraries were tested in MCNPX without significant improvement in the results. Discrepancies observed between both codes are still under investigation.

b. OSL comparison

On Fig. 2, one can observe that uncorrected OSLs always over-estimate the dose. Before applying the correction, the global mean error between OSL and IC measurements is around 5.6 %, after correction, it decreases to 1.1 %. This result demonstrates the relevance of OSL dosimeters to measure low doses in out-of-field regions and thus to validate the calculations of the MC calculation tool in a real IMRT configuration with an anthropomorphic phantom.

c. DXTRAN tests

PENELOPE simulations conducted in this study are time consuming (more than 5 days of computation on 48 processors for the whole water tank dose calculation). The code converges slowly for these calculations because of the low number of particles present out-of-field. Preliminary results obtained about the use of DXTRAN in MCNPX show an increase of the efficiency by a factor of 300. Therefore the implementation of such a tool in PENELOPE could significantly reduce the computation time of out-of-field dose and make enable the implementation of our tool in clinical practice.

IV. CONCLUSIONS
The comparison of the MC tool based on PENELOPE with measurements gives satisfying agreements for out-of-field doses. The next step is the clinical validation with the OSL dosimeters within an anthropomorphic phantom and with an IMRT step-and-shoot treatment plan.

At the same time we are working on the acceleration of the calculations. The implementation of reduction variance techniques such as DXTRAN in the MC tool should be helpful by increasing calculation efficiency out of the beam. This implementation is undergoing and we will soon be able to give our first results of the acceleration of the PENELOPE code.

Part of the work has been done within the framework of EURADOS WG9.

V. REFERENCES


