Short Communication

An efficient planning technique for low dose whole lung radiation therapy for covid-19 pandemic patients

Lulin Yuan a, Siyong Kim a,*, Jatinder Palta a, b, Michael P. Hagan a, b

a Department of Radiation Oncology, Virginia Commonwealth University, Richmond, VA 23219, United States
b VA National Radiation Oncology Program Office, Richmond, VA 23249, United States

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ABSTRACT

This study aimed to establish an efficient planning technique for low dose whole lung treatment that can be implemented rapidly and safely. The treatment technique developed here relied only on chest radiograph and a simple empirical monitor unit calculation formula. The 3D dose calculation in real patient anatomy, including both nonCOVID and COVID-19 patients, which took into account tissue heterogeneity showed that the dose delivered to lungs had reasonable uniformity even with this simple and quick setup.

1. Introduction

Low dose X-ray radiation (LDIR) has been shown to have immunosuppressive effect in human. Prior to the introduction in 1939 of antibiotic therapy for pneumonia, LDIR showed promising effect for a wide spectrum of community acquired pneumonias [1]. Reviewing this experience in 2013, Calabrese and Dhawan summarized successful studies involving more than 700 patients whose pneumonias were treated with X-rays [2]. While for the past seven decades clinical evaluation of the use of ionizing radiation has centered on high-dose cancer therapies, LDIR is still being used to treat many “non-malignant disorders”, such as painful arthrosis, plantar fasciitis, keloids and heterotopic ossification, etc.[3].

Many deaths from the COVID-19 pandemic were related to the lung injury caused by the inflammatory reaction of the lung [4]. Immunosuppression was suggested as possible therapy that could improve mortality for severe COVID-19 patients [5–8]. Currently, low dose radiation therapies of three fractions of 0.35–0.50 Gy up to 1 Gy in a single fraction are under investigation in clinical trials to reduce the hyperinflammatory reaction in the lung (see, e.g., [9]).

An radiation therapy treatment planning technique for such low dose whole lung treatment was introduced in this work. Due to the emergency nature of the COVID-19 pandemic and the need to initiate and deliver the LDIR in the very short time window to get possible antiinflammatory effect [10], the goal of the study was to establish an efficient treatment technique that could be implemented rapidly and safely. A simple parallel opposed fields setup for LDIR was proposed by Kirkby and Mackenzie [6] as a proof-of-principle technique without implementation details. In this work, a treatment technique was independently presented. In addition, we proposed an monitor unit (MU) calculation method which could better account for varying patient anatomy and levels of lung consolidation, and validated it on clinical patients.

2. Materials and methods

2.1. Patient data

Twelve clinical patients (six males and six females) including two pediatric patients who had undergone cancer treatment in our institution were randomly selected in this study under an Institutional Review Board approved protocol. These patients (nonCOVID patients) had a wide range of anatomy. Among them, six patients were used to develop an MU calculation method (training patients); the other six for validation (nonCOVID validation patients). In addition, our planning method was validated on six COVID-19 patients whose computed tomography (CT) image sets were downloaded from a publicly accessible data source [10] (see the Supplementary Material for more information).

2.2. Treatment field setup

To enable rapid implementation, the treatment plan utilized

* Corresponding author at: Department of Radiation Oncology, Virginia Commonwealth University, 1250 E. Marshall Street, Richmond, VA 23219, United States.
E-mail address: siyong.kim@vcuhealth.org (S. Kim).

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anterior-posterior parallel opposed fields with equal weighting. A 6MV photon beam was chosen as treatment modality. The isocenter of the fields was at the mid-plane of the total lung volume along the left-right, anterior-posterior and superior-inferior directions, based on an imaging method available in the treatment room, e.g., on-board imager (OBI). The field aperture was shaped by multileaf collimator (MLC) leaves. They were drawn on the 2D radiograph to encompass the whole lung including the mediastinum with a 2 cm aperture margin. The collimator was rotated 90 degree in our system so that the MLC leaves could better conform to the outer contours of the lung. The Beam’s Eye view image of the anterior treatment field for patient #7 is shown in Fig. 1(a) along with the DRR reconstructed from the CT to simulate the OBI image. Similar image for a COVID-19 patient (#6) is also shown (Fig. 1(b)). The whole planning process can be performed in the radiation therapy record and verify system while the patient is waiting.

### 2.3. MU calculation

The treatment planning goal was to deliver a prescription dose of 50 cGy to the mid-plane of both lungs. The dose should cover the whole lung with reasonable uniformity (approximately 15% as typically in Total Body Irradiation (TBI)). A reference point was chosen at the center of the right lung assuming both of them were similar in their dimension. MU was calculated to deliver 100% dose to the reference point. The traditional MU calculation method based on depth dose in water would not be accurate enough due to the lower density of lung tissue. The nomograph method provided in [11] to relate lung dose correction factors with patient thickness in the context of TBI cannot account for the wide variation of lung density caused by the pathophysiological changes in lung tissue seen in COVID-19 patients [12]. In this work, 3D dose calculation was performed in our clinical treatment planning system (Eclipse V13.7, Varian Medical System) using AcurosXB algorithm with inhomogeneity correction turned on. AcurosXB is an implementation of the numerical solver to the linear Boltzmann transport equation [13]. It has been shown to have less than 3% difference from Monte Carlo type of algorithm in heterogeneous medium [13,14].

It has been observed that an important symptom of COVID-19 patients is the inflammation of the lung tissue. The inflammation causes diffuse ground-glass infiltration in the lung which increases the density of the lung to a varying degree [12]. To better represent this pathologic change without 3D information, we artifically altered the lung density uniformly in the treatment planning system. Radiologically, the degree of fibrosis of the lung tissue can be quantified by consolidation, with normal lung density (assumed to be ~0.2 gm/cm$^3$) equivalent to zero (0) consolidation and a lung full of fluid (water) 1.0 (i.e., 100%). In this study, we assumed a linear relationship between the consolidation obtained from planar lung x-ray and the average density of the lung. Thus, we used the following formula to convert the density (in gm/cm$^3$) to consolidation:

$$\text{Consolidation} = 1.25 \times (\text{Density} - 0.2) \quad (1)$$

We assigned the lung tissue to a number of different densities to simulate lung consolidation at 25%, 50% and 75%, in addition to the clinical lung density from CT.

To enable rapid implementation of this treatment technique, an empirical formula was derived to calculate MUs directly from a simple measurement of patient thickness as the anterior-posterior separation at mid-sternum level and a chest radiograph to clinically assess lung consolidation. However, since this study was performed on retrospective dataset, the thickness was measured equivalently from CT images. A linear multiple regression method was used to fit a correlation between the MU and the patient body thickness and lung consolidation from six training patients (24 training cases in total after density override). The treatment planning method and the MU calculation formula were validated with the remaining six nonCOVID patients with various levels of lung density override and six COVID-19 patients. The dose at the center of the lung, as well as the maximum and mean lung dose were extracted to evaluate the dosimetric quality of the treatment technique.

### 3. Results

The MUs for the training patients with clinical CT lung density and altered lung densities were in the range from 24.3 to 30.4. Within the nonCOVID validation patients, the actual lung consolidations were in the range of 0.03 to 0.35. Both the two pediatric patients had higher lung consolidation than all the adult patients. The lung consolidation within the COVID-19 patients was from 0.0 to 0.19.

The formula to calculate MU per field is given below in terms of MU per cGy of prescription dose so it can be applied to different prescription dose levels:

$$\text{MU/cGy} = 0.426 + 0.0036 \times \text{Body Thickness} \times \text{Consolidation} \times 0.086$$

The regression coefficient of determination was 0.83 and the largest relative MU error was 4.5% with this formula. The MU, body thickness and lung consolidation for these plans are shown as scatter plots on Fig. 2.

Among the nonCOVID validation patients, the maximum deviation for the mid-lung dose was 3% (range: 98–103%) and the range of maximum and mean lung dose were from 106% to 113% and from 94%.

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![Fig. 1. (a) Beam’s Eye view image of the AP treatment field for patient #7 along with the DRR reconstructed from the CT image set to simulate the x-ray radiograph. Blue lines indicate MLC positions. Each division on the field graticule represents one cm. (b) The AP treatment field and DRR for COVID-19 patient #6.](image-url)
COVID-19 patients are shown on their CT images in Supplementary Table S1 for the non-COVID validation patients and Table S2 for the COVID-19 patients, respectively. The isodose distributions for the patients with very small lung volume. Secondly, the thicknesses of the breast and chest wall were not taken into account in the MU calculation. We can see that these are reasonable assumptions based on the small variation in the actual mid-lung dose among the validation patients.

In this work, the AcurosXB algorithm was used and the lung consolidation was considered to improve dose calculation accuracy inside lung. AcurosXB has been shown to be more accurate than the convolution-superposition type of algorithms, especially in the low density lung tissues and the lung soft tissue interface [16,17]. However, we realized that the determination of lung consolidation level from either chest X-ray or an OBI image may have large uncertainty. Another limitation with the 2D projective imaging is that it lacks information on the 3D distribution of lung density. Thus we made the approximation that the consolidation is uniform over the whole lung. The approximation is based on the observation from Eq. (2) that the delivered dose is not very sensitive to the variation in lung consolidation: a 10% change in consolidation results in about 2% deviation on mid-lung dose. Similar observation has been made in a previous study which concluded that the dosimetric impact of the CT number variation is limited in low density medium [18]. This approximation is also supported by the radiographic studies of COVID-19 patients that show extensive diffuse ground-glass opacities with consolidation in the lungs of later stage patients [12].

It should be pointed out that the MU calculations in Eq. (2) are associated with the output calibration condition for the linac in our clinic, which is described in detail in the Supplementary Material. Therefore, readers should validate them on their own machines and under their own setup conditions, and establish the equivalent formula for these conditions [19,20]. For instance, in the case of different calibration condition for reference dosimetry, the derived MU will have to be corrected using the well-established correlation between two different setup conditions for output calibration [14].

In conclusion, an efficient planning technique for low dose whole lung treatment was presented in this work. 3D dose calculation in real patient anatomy, including both non-COVID and COVID-19 patients, showed that the dose delivered to the lungs had reasonable uniformity even with this simple and quick setup.
Declaration of Competing Interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

Appendix A. Supplementary data

Supplementary data to this article can be found online at https://doi.org/10.1016/j.phro.2020.10.004.

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