Towards mid-position based Stereotactic Body Radiation Therapy using online magnetic resonance imaging guidance for central lung tumours

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ABSTRACT

Background and purpose: Central lung tumours can be treated by magnetic resonance (MR)-guided radiotherapy. Complications might be reduced by decreasing the Planning Target Volume (PTV) using mid-position (midP)-based planning instead of Internal Target Volume (ITV)-based planning. In this study, we aimed to verify a method to automatically derive patient-specific PTV margins for midP-based planning, and show dosimetric robustness of midP-based planning for a 1.5T MR-linac.

Materials and methods: Central (n=12) and peripheral (n=4) central lung tumour cases who received 8x7.5 Gy were included. A midP-image was reconstructed from ten phases of the 4D-Computed Tomography using deformable image registration. The Gross Tumor Volume (GTV) was delineated on the midP-image and the PTV margin was automatically calculated based on van Herk’s margin recipe, treating the standard deviation of all Deformation Vector Fields, within the GTV, as random error component. Dosimetric robustness of midP-based planning for MR-linac using automatically derived margins was verified by 4D dose-accumulation. MidP-based plans were compared to ITV-based plans. Automatically derived margins were verified with manually derived margins.

Results: The mean D95% target coverage in GTV + 2 mm was 59.9 Gy and 62.0 Gy for midP- and ITV-based central lung plans, respectively. The mean lung dose was significantly lower for midP-based treatment plans (difference:-0.3 Gy; p < 0.042). Automatically derived margins agreed within one millimeter with manually derived margins.

Conclusions: This retrospective study indicates that mid-position-based treatment plans for central lung Stereotactic Body Radiation Therapy yield lower OAR doses compared to ITV-based treatment plans on the MR-linac. Patient-specific GTV-to-PTV margins can be derived automatically and result in clinically acceptable target coverage.

1. Introduction

Stereotactic body radiotherapy (SBRT) is the recommended treatment for inoperable patients with early stage non-small cell lung cancer (NSCLC). These tumours require a Biologically Effective Dose (BED) of ≥100 Gy to achieve local control ([1]), but the proximity of central tumours to the mediastinum may prohibit giving such a high dose to the tumour with acceptable risk of toxicity to critical mediastinal structures. Treatment of (ultra)-central lung tumours is therefore controversial as it appears to have a higher rate of complications than treatment of peripheral tumours ([2–5]).

Thoracic radiotherapy is associated with large and complex geometrical uncertainties, such as anatomical changes over the course of treatment and respiratory motion, which must be incorporated in the treatment plan via margins around the tumour to create the planning target volume (PTV) ([6,7]). MR-guided radiotherapy enables on-table plan adaptation to the daily anatomy, thereby reducing uncertainties associated with inter-fractional anatomical changes ([8,9]). Central lung tumours are now routinely treated with MR-guided radiotherapy ([10,11,8,12]).

Respiratory motion is one of the largest sources of geometrical uncertainty for lung radiotherapy. The conventional approach, using an internal target volume (ITV), is highly conservative and leads to unnecessary exposure of healthy tissue. Gating treatment delivery to a
particular respiratory phase significantly reduces the uncertainty associated with respiratory motion and thus the irradiated volume ([91]). Clinically, improved dose conformity through gating must be balanced with reductions in duty cycle. Planning on the time-weighted mean position of the tumour, known as the mid-position (midP), also enables a significant reduction of the PTV compared to ITV-based planning whilst maintaining a 100% duty cycle and ensuring dose coverage of the tumour ([133]). The 3D tumour motion can then be regarded as a source of random error when deriving the PTV margin. This 3D motion is generally visually derived from the peak-to-peak (p2p) motion of the tumour along the three Cartesian axes, but can in principle be extracted from the deformation vector fields (DVFs) generated by the deformable image registration (DIR) used to create the mid-position image. Automated derivation of the 3D motion would also be much easier to implement in an online adaptive MR-linac workflow.

In this work, we aimed to verify automatic derivation of margins for central lung tumours and to demonstrate the dosimetric robustness of midP PTV creation, the non-linear van Herk margin recipe ([15]), with reductions in duty cycle. Planning on the time-weighted mean position of the tumour, known as the mid-position (midP), also enables a significant reduction of the PTV compared to ITV-based planning whilst maintaining a 100% duty cycle and ensuring dose coverage of the tumour ([133]). The 3D tumour motion can then be regarded as a source of random error when deriving the PTV margin. This 3D motion is generally visually derived from the peak-to-peak (p2p) motion of the tumour along the three Cartesian axes, but can in principle be extracted from the deformation vector fields (DVFs) generated by the deformable image registration (DIR) used to create the mid-position image. Automated derivation of the 3D motion would also be much easier to implement in an online adaptive MR-linac workflow.

2. Materials and methods

2.1. Patient selection and imaging

In this retrospective study, 17 patients (median age: 70, range: 30–86; four female) with NSCLC were included. All patients underwent SBRT on a conventional cone-beam CT-guided linac. Thirteen patients with central lung tumour were selected in chronological order between 12–2017 and 11–2019, of which one patient was excluded due to poor image quality. Another four patients with peripheral lung tumours exhibiting large respiratory motion (> 10 mm) p2p were included to investigate robustness of the derived margins for large tumour movement. Patient data were collected retrospectively under the FAST-ART protocol (IRB reference: 20–519/C). Patients underwent 4D-CT imaging with 10 respiratory phases, from which an averaged 3D-CT image and a midP image ([13,14]) were reconstructed for ITV-based and midP-based dose planning, respectively (Supp. 6.1).

2.2. Target definition and margin

For each patient, the ITV was delineated on the averaged 3D-CT image taking 4D tumour excursion into account, while the GTV was delineated on the midP-CT image. The ITVs, which were taken from the clinical plans, were delineated by several radiation oncologists (in training) and verified by a second specialized radiation oncologist, whereas the GTVs were delineated by a single radiation oncologist (in training) focused on lung radiotherapy. For the ITV-based treatment plans an isotropic ITV-PTV margin of 3 mm was used following institutional practices. For midP PTV creation, the non-linear van Herk margin recipe was used ([15]),

$$M = 2.5\Sigma + 0.67\sqrt{\left(\sigma_p^2 + \sigma_b^2 + \sigma_t^2\right) - 0.67\sigma_p}$$

with M the GTV-PTV margin to get target coverage in 90% of the population with the 75% isoline. Furthermore $\sigma_b$ (6.4 mm) is the beam penumbra in lung tissue, $\sigma_\Sigma$ the random error component due to breathing, and $\sigma_t$ the residual systematic and random error components, respectively ([16]) (Table S2). Note that the 2 mm delineation uncertainty ([17]) was accounted for a random error (\(\sigma_t\)) to better reflect the daily target redelineation in MR-linac workflows. A similar delineation uncertainty was assumed for the ITV plans which in clinical practice is re-delineated on the daily motion-averaged MRI ([18]).

In this study, $\sigma_b$ was directly calculated per voxel from the ten DVFs generated for midP reconstruction (Fig. 1). For each axis, i.e. cranio-caudal (CC), anterior-posterior (AP) and left-right (LR), the standard deviation (SD) of the ten DVFs was calculated. The automatically-derived median SD of all voxels within the GTV ($\sigma_{\text{GTV}}$) was used as input for the random error due to tumour motion in the aforementioned margin recipe, which yields automatically-derived anisotropic GTV-PTV margins ($\text{Margin}_{\text{GTV}}$) specific to each patient.

2.3. Treatment planning

Treatment plans were generated in Monaco v5.40.01 for the Unity
2.4. Validation of GTV-PTV margins

In order to validate the automatically derived GTV-PTV margins for the midP planning process, the automatically derived $\sigma_{\text{auto}}^{SD}$ was compared to a manually derived $\sigma_{\text{man}}^{SD}$. The $\sigma_{\text{man}}^{SD}$ was derived by manual rigid registration of the tumour region of each 4D-CT phase to the end-exhale phase, resulting in ten displacement vectors. The SD of these displacement vectors is defined as the $\sigma_{\text{man}}^{SD}$, and should be approximately the same as the $\sigma_{\text{auto}}^{SD}$.

In current clinical practice, $\sigma_{\text{man}}$ for midP is typically based on the p2p amplitude of the tumour. The p2p amplitude is manually measured and converted to a random error component by using the rule of thumb that $\sigma_{\text{man}}^{p2p}$ can be approximated as one third of the p2p amplitude ([20,21]).

All $\sigma$’s were used as input for the $\sigma_{\text{man}}^{SD}$ in the van Herk recipe. Bland–Altman analyses ([22]) were performed to compare the margins derived by manual rigid registration ($\text{Margin}_{\text{man}}^{SD}$), and the margins determined from the p2p motion ($\text{Margin}_{\text{p2p}}^{SD}$), with the automatically-derived margins ($\text{Margin}_{\text{auto}}^{SD}$) ([22]). The peripheral lung tumour cases were processed in the margin analysis and shown the Bland–Altman plot but were excluded from statistical analysis.

2.5. 4D-dose accumulation

A 4D-dose accumulation was performed to evaluate the dosimetric robustness of each plan under respiratory motion conditions (Fig. 1). For each treatment plan, the dose was recalculated on the ten phases of the 4D-CT using a standalone GPU-Monte-Carlo Dose (GPUMCD) engine ([23]). The dose was then warped and accumulated to the midP-CT using the DVFs generated by the midP reconstruction process. The dose accumulation was regularized to keep energy-mass transfer (EMT) constant ([24]).

2.6. Dosimetric evaluation

Target coverage was evaluated on the GTV expanded with a 2 mm margin (GTV + 2 mm) for both the ITV-based and the midP-based plan. The evaluation margin was used to incorporate one SD of delineation uncertainty ([17]), which is otherwise not accounted for in the 4D-dose accumulation workflow. Lung volume for mean lung dose (MLD) calculation was defined as lung minus GTV. For the central lung tumours, a statistical analysis with a pairwise Wilcoxon-signed rank test was performed to compare the metrics determined from the 4D-accumulated dose. Given the limited number of patients, the statistical analysis of the organ-at-risk (OAR) DVH-parameters was only used for hypothesis generation to provide an overview of the differences between both planning methods.

We hypothesized that a larger PTV would correspond to a higher MLD, and that a midP-based treatment would be most beneficial in terms of reduction of MLD for central lung patients with larger p2p tumour motion ([25]). MLD difference between midP- and ITV-based plans was therefore compared with the difference in PTV and p2p-motion. The four peripheral cases were not analyzed statistically, but were included in the figures to show the trend for larger tumour motion.

2.7. DDM and $\delta$-index

The distance discordance metric (DDM) ([26,27]) was calculated for each patient to assess the precision of the registration used for midP image reconstruction and dose accumulation. Registration of the end-exhale phase to the end-inhale phase via an intermediate registration to each in-between phase of the 4D-CT results in eight DVFs. The SD of the eight voxel positions in the end-exhale phase gives the DDM value for each voxel, with a lower score indicating higher DIR consistency. The resulting DDM map was then warped to the midP image ([27]).

To determine the uncertainty in the accumulated dose distributions the $\delta$-index was calculated for all voxels with a dose difference criterion of 3%. The $\delta$-index is a novel metric to evaluate the uncertainty associated with the dose accumulation process, by incorporating the uncertainty of the registration (i.e., the DDM) and the spatial variation in the 4D-accumulated dose distribution in the neighbourhood of each voxel ([28]). In analogy to the gamma-index, the uncertainty of the 4D-dose accumulation is defined to be acceptable when the $\delta$-index is <1.

3. Results

3.1. Comparison of margin derivation methods

Bland–Altman plots showing the comparisons between automatically-derived margins ($\text{Margin}_{\text{auto}}^{SD}$) with corresponding manually derived margins based on the p2p motion ($\text{Margin}_{\text{p2p}}^{SD}$) or the SD ($\text{Margin}_{\text{SD}}^{\delta}$) are shown in Fig. 2. For the central lung tumours, the PTV margins in CC direction for $\text{Margin}_{\text{auto}}^{SD}$, $\text{Margin}_{\text{p2p}}^{SD}$, and $\text{Margin}_{\text{SD}}^{\delta}$ ranged between 4.0–6.1 mm, 4.0–7.1 mm and 4.0–7.1 mm respectively. The margins in AP and RL direction were all four and three millimeter when rounded. The differences between $\text{Margin}_{\text{auto}}^{SD}$ and $\text{Margin}_{\text{p2p}}^{SD}$, or $\text{Margin}_{\text{SD}}^{\delta}$ were mainly along the CC axis, as expected with the motion predominantly in cranio-caudal direction. Differences between $\text{Margin}_{\text{auto}}^{SD}$ and $\text{Margin}_{\text{p2p}}^{SD}$, and between $\text{Margin}_{\text{auto}}^{SD}$ and $\text{Margin}_{\text{SD}}^{\delta}$, ranged between −1.0
mm to +0.2 mm and −1.1 mm to +0.3 mm respectively for the central lung tumour cases, with a trend of larger variation between margin derivation methods for larger target motion. For the peripheral lung tumours with large motion, a similar larger variation was observed between Margin$^{SD}_{\text{auto}}$ (range margins: 4.7–5.8 mm) and Margin$^{SD}_{\text{man}}$ or Margin$^{p2p}_{\text{man}}$, with differences from −0.5 to 0.3 mm and −0.9 to −0.04 mm, respectively.

After rounding the margins to the nearest millimeter, to mimic the precision of treatment planning systems, the automatically-derived margin was in two instances smaller than the p2p-margin by 1 mm, and in one instance larger by 1 mm. All differences were along the CC axis. No differences were found between margins determined along the AP and RL axes.

### 3.2. Plan comparison

For central lung tumour cases, the mean (SD) PTVs for the ITV-based plans was significantly larger ($p = 0.0037$) than for the midP-based plans with 40.0 cm$^3$ (41.4 cm$^3$) and 28.9 cm$^3$ (40.7 cm$^3$), respectively (Table S3). The metrics for the target and OAR doses determined for the 4D-dose accumulations for the midP treatment plans were compared to

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Fig. 3. Example patient with 4D-accumulated midP and ITV doses, dose differences to planned dose, and the midP versus ITV dose difference.
the corresponding 4D-dose accumulations for the ITV-based plans. In the example patient with respiratory motion of 23 mm along the CC-axis, it can be seen that dose was mainly reduced in the CC-direction, by up to 30 Gy, for midP-based plans compared to ITV-based plans (Fig. 3). The mean (SD) D95% coverage of the GTV + 2 mm was 59.9 Gy (4.1 Gy) and 62.0 Gy (4.7 Gy) for the midP and ITV plans for the central lung tumours respectively. The four peripheral lung tumours all got acceptable dose coverage with D95% ranging between 60.1–63.1 Gy and 63.7–68.1 Gy for midP-based and ITV-based dose distributions, respectively.

The dose delivered to the target for central lung tumours in midP plans was significantly lower for D95% (-2.9 Gy, p = 0.032) compared to the ITV plan, but not for D98% (-2.3 Gy, p = 0.057) and D0.1 cm³ (+1.3 Gy, p = 0.73) (Fig. 4). The midP-based accumulated doses showed a D95% and D98% closer to the prescribed dose (60 Gy), with the spread (SD) in dose two-thirds of the ITV-based accumulated plans. Several DVH parameters for OARs were significantly lower, including the MLD, which showed a small yet significant reduction for midP-based versus ITV-based treatment planning of median 0.3 Gy (range:-3.0 Gy + 0.4 Gy; p = 0.042). The aorta DVH parameters show that in two cases the dose was above the planning constraints for the ITV dose distributions whereas the midP dose distribution was below the planning constraint.

4. Discussion

In this retrospective study, we provided the first successful demonstration of the dosimetric suitability of mid-position-based treatments for central lung SBRT on an MR-linac. Using 4D dose calculation on CT, and EMT mapping, we showed that mid-position-based treatment plans maintain target coverage with slightly reduced dose compared to ITV-based reference plans, while achieving a variable amount of OAR sparing depending mostly on respiratory amplitude and target volume. The four peripheral cases with large tumour motion indicate that with larger motion amplitudes, the target is still covered using midP based treatment planning. Importantly, our dosimetric analysis supported the automatic derivation of patient-specific GTV-to-PTV margins based on the DVFs inherent to the generation of the mid-position CT image. By directly inserting the SD of motion into the margin equation, motion data from multiple deformable registration is evaluated, which increases the robustness against registration outliers. In addition, the simplification that the SD of motion typically corresponds to one-third of the p2p motion is retired. Automating margin calculations in this way speeds up
margin creation, and thus opens opportunities for daily margin adaption for MR-linac systems.

It should be noted that clinical experience with stereotactic lung treatments on MR-linac devices is still limited ([9,29,11]). Intra-fractional motion management based on MRI guidance holds great promise ([30,31]) but has not yet achieved its full potential. Currently, ITV-based lung treatments are still the only vendor-supported motion management strategy on the Unity MR-linac. For MRIdian MR-linacs, respiratory gating ensures tight PTV margins to reduce dose to organs-at-risk, with a trade-off on treatment efficiency, which might lead to longer treatment times ([32]). In future, dynamic multi-leaf collimator (MLC) tracking is expected to combine the dosimetric advantages of gating with the 100% duty cycle of ITV/midP deliveries ([33]). In the meantime, this study demonstrates that mid-position-based treatments offer a viable dosimetric improvement over ITV-based deliveries while maintaining a 100% duty cycle. For liver SBRT, it was previously shown that a mid-position-based treatment strategy is feasible on the Unity MR-linac ([34]).

Given the small sample size in this study, the statistical analysis on OAR dose should be considered as exploratory and hypothesis generating. Multiple testing was not performed, given the sample size constraint, and the explorative nature of the study. Validation of these data is needed in a larger cohort. This study is limited to analysing 4D-accumulated dose based on the pre-treatment 4D-CT. Snap-shot 4D-CTs poorly correlate with motion during treatment ([35–37]). This limitation can be overcome by creating a mid-position MRI based on 4D-MRI for daily plan adaption ([27]), especially when combined with (automatically) extracted daily GTV-to-PTV margins from the 4D-MRI. MidP MR-image reconstruction was shown to be feasible for daily plan adaptation on an MR-linac ([38]). Underprediction of peak-to-peak target motion during treatment on pre-treatment 4D-CTs is known issue([36,37]), which will affect the ITV-based more than the midP-based treatment plans as the midP-margin only weakly depends on peak-to-peak motion. 4D-CTs are also prone to image artefacts induced by irregular or slow breathing ([39]). As our mid-position CTs were derived from 4D-CT data, any such artefact could propagate to the midP-CT, thus causing dose calculation and warping errors. Artefacts outside the beam path were deemed acceptable. In addition, the DDM, a quantitative metric for DIR uncertainty, was calculated. The deformable registration algorithm used in this study does not explicitly account for sliding motion, therefore registration accuracy is limited near those regions. The novel \( \delta \)-index generally yielded very high pass rates in the high dose region, indicating a high level of confidence in our 4D-accumulated dose distributions. For one (excluded) patient, the \( \delta \)-index showed a very low passing rate, in agreement with the visually assessed low image quality of the underlying 4D-CT. Interestingly, the DDM and \( \delta \)-index were able to detect this case with suboptimal midP-image quality, which might in future help to automatically identify patients that could benefit from a new 4D-CT planning scan. In our margin calculation, delineation uncertainty was assumed to be a random error as the target is re-delineated during each treatment fraction. Further investigation of inter- and intra-observer delineation uncertainty during treatment on an MRI-linac is needed to confirm this assumption. Similar interobserver variation error was assumed for ITV and GTV delineation, as the difference for this error is within one mm between different types of CT reconstructions ([40]). Importantly, adapting plans on daily MRI and accumulating dose across fractions was considered outside the scope of this study. While an in-depth analysis of inter-fractional anatomical changes and plan adaptation is valuable, it is not expected to undermine our findings relating to target coverage under periodic intra-fractional.

Fig. 6. Central lung tumour case with 4D-accumulated dose for a midP-based plan, and corresponding \( \delta \)-index pass-fail map for the \( \geq 10\% \)-prescribed dose region, and the DDM map.
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.

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Appendix A. Supplementary data

Supplementary data associated with this article can be found, in the online version, at https://doi.org/10.1016/j.phro.2022.05.002.

References


