Comparison of planning target volumes based on three-dimensional and four-dimensional CT imaging of thoracic esophageal cancer

Background and purpose: To investigate the definition of planning target volumes (PTVs) based on four-dimensional computed tomography (4DCT) compared with conventional PTV definition and PTV definition using asymmetrical margins for thoracic primary esophageal cancer.

Materials and methods: Forty-three patients with esophageal cancer underwent 3DCT and 4DCT simulation scans during free breathing. The motions of primary tumors located in the proximal (group A), middle (group B), and distal (group C) thoracic esophagus were obtained from the 4DCT scans. PTV

4D

was defined on 3DCT using the tumor motion measured based on 4DCT, PTV conventional (PTV

conv

) was defined on 3DCT by adding a 1.0 cm margin to the clinical target volume, and PTV

4D

was defined as the union of the target volumes contoured on the ten phases of the 4DCT images. The centroid positions, volumetric differences, and dice similarity coefficients were evaluated for all PTVs.

Results: The median centroid shifts between PTV

4D

and PTV

conv

and between PTV

conv

and PTV

3D

in all three dimensions were <0.3 cm for the three groups. The median size ratios of PTV

4D

to PTV

3D

were 0.80, 0.88, and 0.71, and PTV

4D

to PTV

conv

were 0.67, 0.73, and 0.76 ($\chi^2=-3.18, -2.98, and -3.06; P=0.001, 0.003, and 0.002$) for groups A, B, and C, respectively. The dice similarity coefficients were 0.87, 0.90, and 0.81 between PTV

4D

and PTV

3D

and 0.80, 0.84, and 0.83 between PTV

4D

and PTV

conv

($\chi^2=-3.18, -2.98, and -3.06; P=0.001, 0.003, and 0.002$) for groups A, B, and C, respectively. The difference between the degree of inclusion of PTV

4D

in PTV

3D

and that of PTV

4D

in PTV

conv

was <2% for all groups. Compared with PTV

conv

, the amount of irradiated normal tissue for PTV

3D

was decreased by 11.81% and 11.86% in groups A and B, respectively, but was increased by 2.93% in group C.

Conclusion: For proximal and middle esophageal cancer, 3DCT-based PTV using asymmetrical margins provides good coverage of PTV

4D

; however, for distal esophageal cancer, 3DCT-based PTV using conventional margins provides ideal conformity with PTV

4D

.

Keywords: planning target volume, 4DCT, 3DCT, esophageal carcinoma

Introduction

The incidence rate of esophageal carcinoma varies considerably among different geographic regions throughout the world and is particularly high in the People’s Republic of China, where it exceeds 1 per 1,000 individuals.1,2 Historically, chemoradiotherapy has played an important role in the management of localized esophageal cancer because it provides better palliation than does radiotherapy alone and improves the likelihood of long-term, progression-free survival.3 New technological advances in radiation techniques such as intensity-modulated radiotherapy, respiratory-gated radiotherapy, image-guided radiotherapy, and positron emission tomography (PET)-
PET/computed tomography (CT)-based radiotherapy have allowed for a selective increase in the dose delivered to the target for esophageal cancer without any significant increase in the dose delivered to the organs at risk.\(^4\,^3\)

The goal of modern radiotherapy approaches based on recent technological advances is to minimize the risk of damage to healthy tissues by improving the gross tumor volume (GTV) definition (PET or PET/CT) and reducing intrafraction motion (respiratory-gated radiotherapy; four-dimensional CT [4DCT]) and interfraction motion (image-guided radiotherapy; cone beam CT).\(^6\) Obviously, the larger the tumor and extent of involvement are, the higher the toxicity to the organs at risk will be.\(^7\,^8\) Therefore, this study focused on the definition of the planning target volume (PTV).

Tumor displacement and deformation affect the definition of PTV during the course of treatment. Creating a PTV for a moving target is an important but complicated clinical problem for various anatomical regions, such as the esophagus. There are many sources of fractional variations in internal structures that can occur (eg, physiological movements of the tumor and organs, mostly originating from respiratory or cardiac cycles and setup error).\(^9\) To account for these intrafractional and interfractional variations, large population-based margins are used. The incorporation of these uncertainties further increases the excessive irradiation of normal tissues. Because of the technical limitations of conventional PTV definition, new PTV definition methods have been investigated to decrease the size of the target volume and reduce normal tissue toxicity.

During free breathing, tumor and organ motions always influence the accuracy and quality of 3DCT imaging of thoracic malignancies, including esophageal carcinoma. The detailed motion of the tumor, the various spatial positions of the tumor, and anatomical information averaged over one breathing cycle should be carefully provided when using 4DCT.\(^10\,\,^{13}\) Breathing characteristics vary greatly among individual patients, and respiratory-induced target motion and interfraction target motion are unsymmetrical.\(^10\,\,\,^{15}\) Therefore, esophageal tumor motion must be separately assessed in each individual patient because doing so may allow the tumor margins to be decreased and consequently allow for reduction in the PTV size and the radiation exposure of normal tissue. Based on this concept, we performed patient-specific PTV definition using images acquired on 4DCT, and these PTVs were compared with the conventional PTVs formed by adding clinical experience margins and individual margins to the free-breathing planning 3DCT scan. This work was a pilot study to investigate the feasibility of using 4DCT when contouring individualized PTVs for esophageal carcinoma and to assess the differences between patient-specific and population-based target volumes as well as their inherent advantages and disadvantages.

**Materials and methods**

**Ethics statement**

All patients provided signed informed consent to participate in the study and before undergoing further imaging during radiotherapy. The study design was approved by the ethics committee of Shandong Cancer Hospital and Institute (approval ID: SDTHEC20110130).

**Patient population**

Forty-three patients with esophageal carcinoma who underwent 4DCT scanning were included in this study. The mean patient age was 66 years (range: 41–83 years), and 37 of the patients were men. Nine patients had pathologically confirmed adenocarcinoma of the thoracic esophagus, and 34 patients had squamous cell carcinoma of the thoracic esophagus. Thirteen patients had primary tumors in the proximal third of the esophagus (group A), whereas 18 and 12 patients had primary tumors in the middle third and distal third of the esophagus (groups B and C), respectively. All patients exhibited regular breathing patterns.

**Image acquisition**

Patients were immobilized in the supine position with their arms above the head using a vacuum bag. Every patient underwent 3DCT and, immediately afterward, respiration-correlated 4DCT on a 16-slice CT scanner (Philips Medical System, Cleveland, OH, USA). For 3DCT, each scan (360° rotation) took 1 second to acquire followed by a 1.8 seconds dead time and had a 2.4 cm coverage. The slice thickness in the 3DCT scan was 3 mm. During 4DCT scanning, the Varian Real-time Positioning Management system (Varian Medical Systems, Palo Alto, CA, USA) was used to monitor the patients’ breathing. The Real-time Positioning Management system uses infrared beams to track the trajectory of infrared-reflecting markers placed on the epigastric region of the patient’s abdomen. The signal was sent to the scanner to label each CT image with a time tag. GE Advantage 4D software (GE Healthcare, Waukesha, WI, USA) was used to sort the reconstructed 4DCT images into ten respiratory phases, labeled as 0%–90% based on these tags, with 0% corresponding to the end of inhalation and 50% corresponding to the end of exhalation. The slice thickness was 3 mm, and
the 4DCT data set was transferred to an Eclipse treatment planning system (Eclipse 8.6; Varian Medical Systems).

Measurements of tumor motion
The same clinician contoured the GTV on both the standard free-breathing 3DCT scan and each of the ten respiratory phase volumes for each patient. Each volume was outlined using the same window and level settings. The motion amplitudes of the primary tumor for each patient throughout one respiratory cycle were measured at the center of each GTV, and the 95% upper bound of the cumulative distribution represented an extreme of motion attained by at least a portion of the GTV.

Target volume generation
The clinical target volume (CTV) was created by manually contouring the esophagus at 3 cm superior and inferior to the GTV and then adding 0.5 cm circumferentially to the created volume to account for microscopic spread. The union of all ten CTVs from the 4DCT data was used to generate the internal target volume (ITV\(_{4D}\)). ITV\(_{3D}\) was constructed by adding asymmetric margins to CTV\(_{3D}\) in each spatial direction, depending on the amount of motion measured on the 4DCT phases and the expansion necessary to cover an ITV of 95% of the tumor in all three dimensions. PTVs were generated by applying 0.5 cm expansions to ITV\(_{4D}\) and ITV\(_{3D}\) resulting in PTV\(_{4D}\) and PTV\(_{3D}\), respectively. For PTV conventional (PTV\(_{conv}\)), a 1.0 cm margin in all directions was added to CTV\(_{3D}\).

Target volume analysis
PTV\(_{3D}\), PTV\(_{conv}\), and PTV\(_{4D}\) were compared with respect to their centroid positions, volumes, dice similarity coefficients (DSCs), and degrees of inclusion (DIs). The DSC can be used to determine the extent of spatial overlap between two regions of interest and takes values ranging from 0 (no overlap) to 1 (perfect overlap).

\[
\text{DSC} = \frac{2 | A \cap B |}{| A \cap B | + | A \cup B |} \quad (1)
\]

The DI of volume A included in volume B [DI (A in B)] is given by the following equation:

\[
\text{DI} = \frac{| A \cap B |}{A} \quad (2)
\]

From these data, assuming that B is the reference for the standard target volume, for treatment planning based on A, 1 – DI (A in B) of A will be unnecessarily irradiated and 1 – DI (B in A) of B will be lacking irradiation.

Statistical analyses
The Friedman Z test was performed to detect the differences among the GTV centroid displacements in all three dimensions. The Wilcoxon signed-rank test was performed to determine significant differences in variability in the centroid positions and volumes of the PTVs. Additionally, comparison of the PTV volume ratios and DSCs at different locations was performed based on the Kruskal–Wallis H test. A P-value < 0.05 was considered to be significant.

Results
Measurement of primary tumor motion
The average three-dimensional centroid motion amplitude of the GTVs caused by respiration was highest in the superoinferior (SI) direction for each group. The mean motions in the anteroposterior (AP) and lateral directions for tumors located in the distal third of the esophagus were larger than those for upper and middle esophageal tumors (\(\chi^2_{\text{LR}} = 9.72, P = 0.008; \chi^2_{\text{AP}} = 9.08, P = 0.011\)). In the SI direction, the median motion was 0.44 cm for distal tumors compared with 0.31 cm and 0.27 cm for upper and middle esophageal tumors, respectively, although this difference was not statistically significant (\(\chi^2 = 3.33, P = 0.189\)). The 95th percentile values from the cumulative distribution were used to define minimum margins to account for GTV motion during target volume generation (Table 1).

PTV conformity
The differences in the PTV centroid positions between PTV\(_{3D}\) and PTV\(_{4D}\) and between PTV\(_{conv}\) and PTV\(_{4D}\) were < 0.2 cm in the lateral and AP directions and < 0.3 cm in the SI direction. The median tumor volume ratios of PTV\(_{4D}\) to PTV\(_{3D}\) (PTV\(_{4D}\)/PTV\(_{3D}\)) in the upper, middle, and distal thirds

| Parameter | Group A | Group B | Group C |
|-----------|---------|---------|---------|
| LR | AP | SI | LR | AP | SI | LR | AP | SI |
| Minimum | 0.04 | 0.03 | 0.05 | 0.04 | 0.02 | 0.13 | 0.06 | 0.06 | 0.12 |
| Maximum | 0.22 | 0.18 | 1.30 | 0.15 | 0.12 | 0.88 | 0.67 | 1.22 | 1.21 |
| Median | 0.14 | 0.10 | 0.31 | 0.09 | 0.08 | 0.27 | 0.20 | 0.19 | 0.44 |
| Mean | 0.14 | 0.10 | 0.33 | 0.10 | 0.08 | 0.33 | 0.25 | 0.25 | 0.52 |
| SD | 0.06 | 0.05 | 0.33 | 0.04 | 0.03 | 0.21 | 0.18 | 0.32 | 0.32 |
| 95th percentile | 0.18 | 0.13 | 0.54 | 0.12 | 0.10 | 0.46 | 0.36 | 0.47 | 0.72 |
| \(\chi^2\) | 14.63 | 28.90 | 11.62 | 0.001 | 0.000 | 0.003 |

Note: Group A, B, and C patients had primary tumors in the proximal third, the middle third, and distal third of the esophagus, respectively.

Abbreviations: LR, lateral; AP, anteroposterior; SI, superoinferior; SD, standard deviation.
of the esophagus were 0.80, 0.88, and 0.71, respectively. However, the volume ratios of PTV_{4D} to PTV_{conv} (PTV_{4D} / PTV_{conv}) had median values of 0.67, 0.73, and 0.76, respectively. Significant differences were observed in the ratios of the target volumes at different locations, particularly for the upper esophageal tumors ($\chi^2 = -3.18, -2.98$, and $-3.06$, respectively; $P=0.001, 0.003$, and 0.002, respectively).

The median DSCs between PTV_{3D} and PTV_{4D} were 0.87 (mean: 0.87; range: 0.79–0.91), 0.90 (mean: 0.89; range: 0.79–0.91), and 0.81 (mean: 0.81; range: 0.76–0.88) for the upper, middle, and distal esophageal tumors, respectively. The median DSCs between PTV_{conv} and PTV_{3D} were 0.80 (mean: 0.79; range: 0.72–0.84) for group A, 0.84 (mean: 0.83; range: 0.72–0.89) for group B, and 0.83 (mean: 0.83; range: 0.78–0.90) for group C. The Kruskal–Wallis $H$ test indicated that the DSC between PTV_{3D} and PTV_{4D} was significantly larger than that between PTV_{conv} and PTV_{4D} for group A ($\chi^2 = -3.18; P=0.001$) and group B ($\chi^2 = -2.98; P=0.003$) but smaller for group C ($\chi^2 = -3.06; P=0.002$).

The DI of PTV_{3D} in PTV_{4D} exhibited median values of 0.98, 0.98, and 0.99, whereas the DI of PTV_{conv} in PTV_{4D} exhibited median values of 1.00, 1.00, and 0.99 for the upper, middle, and distal esophageal tumors, respectively (Table 2). The median disparities in the DI were only $0.98\%$ for group A, $0.98\%$ for group B, and $0.99\%$ for group C. In the treatment planning based on PTV_{3D} compared with that based on PTV_{conv}, the amount of normal tissue that unnecessarily irradiated was decreased by nearly 11.81% and 11.86% for upper and middle esophageal tumors, respectively. However, for the distal esophageal tumor patients, the average percentage of normal tissue that was unnecessarily irradiated was increased by nearly 2.93%.

**Discussion**

For thoracic esophageal carcinoma, several factors lead to uncertainties in target displacement. The most important causes are patient positioning variability, breathing motion, and changes in the shape of the tumor. Recently, Hawkins et al.\(^9\) have demonstrated that the alignment “clipbox” and selected registration method can affect the displacements obtained. Additionally, for lung tumors, the three-dimensional tumor trajectory exhibits hysteresis ranging from 1 mm to 5 mm.\(^9\) Therefore, the previously described optimal multidisciplinary approach to the measurement of tumor movement using multiple CT scanning might introduce significant artifacts and inaccuracies in the 3DCT images. The acquisition time of 4DCT is $>60$ seconds, allowing for the capture of CT data in separate phases of the respiratory cycle. Additionally, the coregistration of all phases provides precise information regarding the amplitude of the structure motion as well as the position and anatomic deformation information of the structure in each phase of the breathing cycle.

We initially analyzed the respiratory motion of primary esophageal cancers using 4DCT and found that the tumor motion caused by respiration was greatest in the SI direction for all three groups of tumor patients. Compared with upper and middle esophageal cancers, a large intrafractional radial margin (0.36 cm in the lateral direction and 0.47 cm in the AP direction) for distal esophageal cancer would provide tumor motion coverage for 95% of the cases in our study population. These results are comparable to those of previous studies that have investigated three-dimensional tumor motion using time-resolved 4DCT.\(^10\)–\(^13\) Thus, asymmetric margins are recommended because of variations in tumor central displacement in different directions and in different regions caused by respiratory or cardiac cycles.

Delineation accuracies in 3DCT images are influenced by artifacts and partial volume effects that arise with the motion of the thoracic contents caused by respiration.\(^20\)–\(^21\) PTV_{3D} based on 3DCT CTV allows for the definition of an asymmetric margin to increase the target volume. We observed mean volumetric differences between PTV_{3D} and PTV_{4D} of 20%, 12%, and 29% for groups A (proximal esophagus cancer), B (middle esophagus cancer), and C (distal esophagus cancer), respectively. However, the centers of the masses differed by $<0.3$ cm.
4DCT imaging for target volume definition and motion has been well studied in non-small-cell lung cancer, but it is not quite ready for routine clinical use for esophageal cancer PTV definition for radiotherapy. In their non-small-cell lung cancer study, Li et al. evaluated the positional and volumetric differences between PTV and PTV vector in 28 patients. Their results indicated a necessity to expand the internal margin isotropically in a single direction for 3DCT treatment planning. The PTVs derived from 3DCT encompassed a relatively large proportion of normal tissues. To the best of our knowledge, ours is the first study to compare DSC and DI values between 3D and PTV values determined by adding conventional margins and individual margins using 3DCT to delineate the esophageal cancer target volume.

For esophageal cancer, PTVs constructed by applying asymmetric margins to standard 3DCT scans (PTV) provide good coverage of patient-specific PTVs based on the unions of 4DCT CTVs (PTV). PTV allows for an average reduction of 2% in the unirradiated target volume compared with PTV. These results indicate that the use of PTV to permit customization of the target volumes leads to a geographic miss of only 2%. In addition, we found that mean volumes of 22%, 16%, and 31% of the surrounding healthy tissues were unnecessarily irradiated when using PTV compared with PTV for groups A, B, and C, respectively. The latter findings demonstrated that PTV could provide good coverage of PTV but the risk of increasing the affected volume of normal tissues should be noted, particularly for upper and distal esophageal cancers, for treatment planning performed using PTV. 4DCT generates up to ten times more data than conventional 3DCT, and 4DCT also incurs an increased workload by requiring the contouring of multiple target volumes. Our study demonstrated that PTV definition using asymmetric margins applied to planning 3DCT scans (PTV) requires an amount of time that is shorter by a factor of 8.75, on average, than that required for the delineation of ten scans and the creation of a composite PTV.

Conventional PTV definition for esophageal tumors is often based on helical treatment-planning CT, in which the visible primary tumor volume is enlarged by individual isotropic margins for the CTV. The coverage of PTV by PTV was very high, with mean values of 99%, 99%, and 97% for groups A, B, and C, respectively. The amount of normal tissue that was unnecessarily irradiated was decreased to nearly 12% for upper and middle esophageal tumors; however, for the distal esophageal tumor patients, the average percentage of the adjacent normal tissue to be included within the irradiation field was actually slightly increased, by 2.93%. These findings suggest that PTV can compensate for the additional effects that the implementation of 4DCT in target volume definition can offer, but only in the case of distal esophageal cancer.

Large population-based margins further increase the excessive irradiation of normal tissues; however, there is no benefit of a larger PTV with respect to minimizing local regional recurrence and limiting the toxicity to normal surrounding tissues or increasing overall survival. Currently, there are several lines of clinical evidence to suggest a local control and survival advantage with radiation dose escalation. However, radiation-induced lung injury has been a hindering factor in dose escalation, particularly for patients with abnormal heart and pulmonary functions. As we have demonstrated, esophageal tumors move substantially during the respiratory cycle, particularly in the SI direction of the distal thoracic esophagus. Wang et al. have corroborated that systematic gastroesophageal junction displacements in all three dimensions are correlated, to varying degrees, with variations in tidal volume and diaphragmatic excursion during treatment. Intrapatient variability may be caused by the effect of changes in breathing patterns, and interpatient variability is most likely induced by inherent physical differences among patients, for example, tumor lengths and pulmonary functions. Therefore, normal, free-breathing conditions with regular respiratory rhythms are of fundamental importance to the improvement of PTV definition for esophageal cancer. The primary objective of performing a pretreatment CT scan is to reduce the uncertainty in the tumor location resulting from respiration, swallowing, and patient positioning variability.

**Conclusion**

We explored the methods of PTV delineation to minimize the damage to sensitive normal tissues within irradiated fields without sparing the primary cancer for esophageal patients. Asymmetric margins are recommended for proximal and middle thoracic esophagus cancer because of tumor displacement and deformation during the respiratory cycle. This approach reduces the time required for planning and not only provides adequate coverage of PTV but also encompasses a relatively large volume of surrounding healthy tissues. In addition, for distal esophageal cancer, adequate coverage of the moving target within the radiation field can be achieved without excessive irradiation of the surrounding normal tissue by applying clinical experience or the published margin guidelines reported in the literature for PTV definition. Therefore, the thoracic esophageal cancer target volume must be separately assessed because it may influence the size and spatial location of the tumors.
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Disclosure

The authors report no conflicts of interest in this work.

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