Cone-Beam CT-based position verification for oesophageal cancer: Evaluation of registration methods and anatomical changes during radiotherapy

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Abstract

Purpose: To evaluate different registration methods, setup margins and number of corrections for CBCT-based position verification for oesophageal cancer and to evaluate anatomical changes during the course of radiotherapy treatment.

Methods: From 50 patients, 440 CBCT-scans were registered automatically using a soft tissue or bone registration algorithm and compared to the clinical match. Moreover, relevant anatomical changes were monitored. A sub-analysis was performed to evaluate if tumour location influenced setup variations. Margin calculation was performed and the number of setup corrections was estimated. Results were compared to a patient group previously treated with MV-EPID based position verification.

Results: CBCT-based setup variations were smaller than EPID-based setup variations, resulting in smaller setup margins of 5.9 mm (RL), 7.5 mm (CC) and 4.7 mm (AP) versus 6.0 mm, 7.8 mm and 5.5 mm, respectively. A reduction in average number of setup corrections per patient was found from 0.75 to 0.36. From all automatically registered CBCT-scans, a clipbox around PTV and vertebrae combined with soft tissue registration resulted in the smallest setup margins of 5.9 mm (RL), 7.7 mm (CC), 4.8 mm (AP) and smallest average number of corrections of 0.38. For distally located tumours, a setup margin of 7.7 mm (CC) was required compared to 5.6 mm for proximal tumours. Reduction of GTV volume, heart volume and change in diaphragm position were observed in 16, 10 and 15 patients, respectively.

Conclusions: CBCT-based set-up variations are smaller than EPID-based variations and vary according to tumour location. When using KV-CBCT a large variety of anatomical changes is revealed, which cannot be observed with MV-EPID.

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Introduction

The incidence of oesophageal cancer is increasing rapidly. Oesophageal cancer is the eight most common cancer worldwide and the sixth most common cause of death from cancer [1].

Radiotherapy is frequently used as part of the multimodality treatment for oesophageal cancer. For patients with potentially curable oesophageal or oesophago gastric junction cancer, the preferred treatment is neoadjuvant chemoradiotherapy in combination with surgery [2,3]. For inoperable oesophageal or oesophagogastric cancer, definitive chemoradiotherapy is the treatment of choice [4]. The toxicity of chemoradiotherapy depends, among others, on the size of the treated volume and dose to the organs at risk (OAR) [5].

Accurate delineating of radiotherapy target volumes is needed to improve local tumour control and reduce toxicity [6]. The Gross Tumour Volume (GTV) is expanded with anisotropic margins to a Clinical Target Volume (CTV), mainly in cranio-caudal (CC) direction. Subsequently, the CTV is expanded to a Planning Target Volume (PTV) with an additional margin. This margin is used to account for uncertainties in delineation and variation in tumour position [7]. Smaller margins will result in less toxicity but will...
increase the risk of under-dosage of the clinical target volume [5]. Therefore Image Guided Radiotherapy (IGRT) using sophisticated imaging methods is crucial during treatment to improve the precision and accuracy of treatment delivery. For position verification, different methods are available, such as MV-EPID and kV-ConeBeam CT (CBCT). In 2013 the Catharina Hospital in Eindhoven introduced CBCT-based position verification for oesophageal and oesophagogastric junction cancer. kV-CBCT based imaging provides additional anatomical information compared to using EPID imaging. The aim of this study was to evaluate different registration methods, resulting setup margins and number of corrections for CBCT-based position verification of oesophageal cancer. Moreover, it was aimed to monitor anatomical changes during the course of the radiotherapy treatment and evaluate its influence on the position verification image registration process.

**Materials and methods**

**Patients**

Between January 2013 and August 2013, 50 consecutive patients treated in combination with CBCT-based position verification were included and data of these patients were analysed retrospectively. Patients had a histologically confirmed adenocarcinoma (AC) or Squamous Cell Carcinoma (SCC) of the oesophagus or oesophagogastric junction and no clinical evidence of distant metastatic spread.

**PET/CT and treatment planning**

Before treatment, a 18F-FDG PET/CT without intravenous contrast was made (Philips Gemini PET/CT, Best, The Netherlands). For this scan, patients were immobilised using an arm support and knee support (Sinmed Radiotherapy Products, Reeuwijk, The Netherlands). All patients with a tumour in the proximal oesophagus were immobilised in a mask (Orfit Industries, Wijinzeem, Belgium). Scan area was from cricoid to kidneys and slice thickness was 3 mm. Delineation of GTV and CTV was performed by the radiation oncologist. GTV was defined by primary tumour and suspect regional lymph nodes. All available information was used to delineate the GTV: physical examination, endoscopy, Endoscopic Ultrasound (EUS) and PET/CT. CTV was defined by the GTV plus the area of the regional lymph nodes up to at least 3 cm in cranial and caudal extension of the oesophagus from the GTV. For distal tumours, the caudal margin should follow the oesophageal and cardia wall and the margin in direction of stomach wall was limited to 2 cm. In case of pathological lymph nodes, the CTV was extended up to the level of the pathologic nodes in all directions. PTV consisted of CTV with an isotropic margin of 1 cm. OAR consisted of lungs, heart, spinal cord and kidneys.

A seven beam Intensity Modulated Radiotherapy plan (IMRT) was created using the Pinnacle Treatment Planning System (Pinnacle 3 TPS, version 9.8), with a photon energy of 6 MV for all beams. The prescribed dose was 41.4 Gy or 50.4 Gy in 23 or 28 fractions. Patients were treated in combination with weekly carboplatin and paclitaxel.

**Radiotherapy treatment and position verification**

Before each treatment patients were positioned using tattoos and skin marks. During the treatment course, position verification was performed using CBCT (Elekta Synergy XVI system, version 4.5). CBCT was performed with a full 360° rotation and a M20 collimator resulting in a Field of View (FOV) of 41 cm and a scan length of 28 cm. Standardly, setup corrections were applied using an offline shrinking action level (SAL) protocol with an initial action level of 10 mm and a maximum number of three measurements (n = 3, \( \alpha = 10 \)) [8]. For patients with large setup variations in the first few fractions, a switch was made to an online correction protocol. For patients with relatively small setup variations, an offline correction protocol accommodated the setup variations adequately within the applied margins.

**Data acquisition**

440 CBCT-scans were analysed retrospectively and reviewed systematically. For evaluating the consistency of CBCT and the required setup margins, different registration methods were used (Fig. 1a). A clipbox around PTV and vertebras was used for clinical registration, which is referred to as the ‘original clipbox’. Clinically, the registration was made such that the best correspondence between CBCT-scan and reference CT-scan was obtained, with respect to all visible anatomical structures: tumour, oesophagus, trachea and tracheal bifurcation, heart, mediastinum, diaphragm and vertebras. This was obtained by soft tissue registration and if necessary manual adjustments were applied. Obviously not always all structures were equally visible, but in general it was sufficient to assess the accuracy of the registration. The longitudinal displacement depended on the tumour location which visible structure was best suited to assess the accuracy in this direction. In addition, the consistency of a registration method using an extended clipbox around PTV, vertebras and sternum and a registration method using a mask of PTV + 1 cm was investigated. All CBCT-scans were registered automatically using a soft tissue or a bone registration algorithm. In contrast to the clinical registration, no manual adjustments were allowed in the automatic registrations.

**Data analysis**

Using the setup results obtained with the different registration methods and algorithms, required setup margins for each direction were calculated according to the Van Herk Formula: margin [mm] = 2.5Σ + 0.7σ [9]. In this formula, Σ indicates the systematic error and σ the random error for the analysed group of patients. Furthermore, the clinically applied SAL correction protocol was simulated for each registration method to estimate the number of setup corrections for each individual patient. Before simulation of the SAL correction protocol, the clinically performed corrections were made undone. For comparison, the results of the required setup margins and number of setup corrections were compared with data of 196 patients with oesophageal or oesophagogastric junction cancer previously treated with MV EPID-based position verification in our hospital. For this, manual registration based on bony anatomy was performed using two orthogonal EPID images.

To investigate if tumour location influenced the set-up variation and thus the required setup margins, a sub-analysis was performed by dividing the current patient group in three categories: tumours in the proximal 1/3 part of the oesophagus, tumours in the middle 1/3 part of the oesophagus and tumours in the distal 1/3 part of the oesophagus including oesophagogastric junction (Fig. 1b). For these different tumour locations, a margin calculation was performed for results based on a bone and a soft tissue registration when using the original clipbox.

**Anatomical changes during radiotherapy treatment**

Since all CBCT-scans were reviewed systematically, anatomical changes during radiotherapy treatment were described and monitored. Subsequently, it was evaluated as to what extent anatomical changes influenced the location of the GTV with respect to applied PTV margins. All findings were discussed with a radiation oncologist and a clinical physicist, in order to assess possible implications for treatment adaptation.
Results

Patient characteristics are summarised in Table 1. Setup results, required setup margins and estimated number of corrections for different registration strategies are summarised in Table 2. These results were adjusted for possible setup corrections. Clinically, CBCT-based setup variations were somewhat smaller than EPID-based setup variation, resulting in smaller setup margins of 5.86 mm (RL), 7.54 mm (CC) and 4.67 mm (AP) versus 6.04 mm (RL), 7.80 mm (CC) and 5.37 mm (AP), respectively. There was a

![Fig. 1. Different regions of interest and tumour sites. (Blue line: GTV, red line: PTV, CTV: not shown).](image)

a: Different regions of interest used for automatic CBCT registrations: original clipbox (left), extended clipbox (middle) and mask of PTV + 1 cm (right).

b: Different tumour sites.
reduction in average number of setup corrections per patient from 0.75 with EPID (i.e. a total of 147 corrections in 196 patients) to 0.36 with CBCT (i.e. a total of 16 corrections in 45 patients).

Using the original clipbox, setup variations, required margins and estimated setup corrections increased slightly when applying a soft tissue algorithm compared to the clinical registration. There was a setup margin increase when applying a bone match algorithm from 5.86 mm to 6.23 mm (RL), 7.54 mm to 8.94 mm (CC) and 4.67 mm to 5.13 mm (AP). Using an extended clipbox, setup variations and margins were comparable to original clipbox registrations for both registration algorithms; the number of corrections increased slightly when applying a soft tissue algorithm (from 0.38 to 0.49), and decreased slightly when applying a bone match algorithm (from 0.69 to 0.56). Using a mask of PTV + 1 cm, setup variations, margins and corrections increased for both soft tissue and bone registration algorithms.

Setup results and required setup margins for different tumour locations are summarised in Table 3. All results were obtained using the original clipbox because this registration method resulted in the least variation and smallest number of setup corrections in the entire group of patients (Table 2).

When applying a soft tissue registration for tumours in the proximal part of the oesophagus, the required margins decreased slightly compared to the clinical registration for the whole patient group. The main decrease was in cranio-caudal (CC) direction from 7.54 mm to 5.61 mm, i.e. longitudinal direction. When applying a bone registration for this tumour site there was a margin increase in lateral (RL) direction from 5.86 mm to 6.24 mm and in vertical (AP) direction from 4.67 mm to 6.71 mm. In longitudinal (CC) direction a margin decrease was found from 7.54 mm to 4.74 mm.

For tumours in the middle part of the oesophagus, the required setup margin was comparable in RL and AP direction when applying a soft tissue or a bone match algorithm. In contrast, setup margin in CC direction increased from 7.54 mm to 8.29 mm when applying a soft tissue match algorithm and decreased from 7.54 mm to 6.29 mm when applying a bone match algorithm. For distally located tumours, the largest required setup margin was also in CC direction, 7.66 mm with soft tissue registration and 9.69 mm with bone registration. In contrast to tumours in proximal or middle part of the oesophagus, the required setup margins increased when applying a bone registration compared to a soft tissue registration from 5.88 mm to 6.24 mm (RL), 7.66 mm to 9.69 mm (CC) and 4.71 mm to 4.75 mm (AP).

The absolute difference (mean ± SD) between soft tissue and bone registration was on average small for tumours in proximal or middle part of the oesophagus, 0.43 ± 0.22 mm (RL), 0.29 ± 0.25 mm (CC), 0.62 ± 0.60 mm (AP). The absolute differences were larger for tumours in distal part or at the oesopha-gogastric junction, 0.79 ± 0.63 mm (RL), 1.82 ± 1.29 mm (CC), 0.57 ± 0.44 mm (AP) (Table 4).

An overview of observed anatomical changes is summarised in Table 5. Frequent anatomical changes were GTV and heart volume reduction and changes in diaphragm position in 16, 10 and 15 patients, respectively. The average heart volume reduction was 6%. All findings were discussed with a radiation oncologist and a clinical physicist to estimate the impact on CTV coverage and dose in OAR.

**Table 1**

| Number of patients | 50 |
|---|---|
| Gender (male/ female) | 38/12 |
| Mean age in years (range) | 69 (49–85) |
| Tumour characteristics |  |
| Pathology (AC/SCC) | 34/16 |
| Tumour location (proximal/middle/distal) | 8/7/35 |
| Treatment | Chemoradiotherapy + Surgery 27 Chemoradiotherapy 23 |
| Position verification protocol (online/ offline) | 5/45 |
correction protocol, the required margins for setup variation may theoretically be reduced to 0 mm. However, other uncertainties such as rotations, intrafraction variation, delineation and treatment uncertainties still exist, therefore a CTV-PTV margin is still required. Even when using an online correction protocol, it is important to select the most appropriate registration method. Registration with original clipbox and a soft tissue algorithm resulted in smallest setup margins and number of corrections. Hence, the original clipbox was used for sub-analysis. The differences between a soft tissue and bone match were small for the included patients in this study when using a certain registration method e.g. the original clipbox. It is difficult to determine which registration (bone or soft tissue) is correct, since there is no gold standard. This way, the automatic registration algorithms can merely be seen as a tool to limit the amount of variation between different observers. When applying clinically, it remains important to review the registration and not only rely on the automatic algorithm, and adjust the registration manually if necessary.

Performing the registration method with an extended clipbox resulted in larger setup variations. The position of oesophagus and vertebrae is obviously not related to the sternum position. Motion due to breathing could be a possible explanation. When performing registration with an extended clipbox, sometimes the automatic registration focuses more on the sternum. In case this was seen, a mismatch between relevant visible structures such as tumour, oesophagus, trachea and tracheal bifurcation, heart, mediastinum and vertebrae could be observed, leading to inaccurate registrations.

When applying the registration with a mask of PTV + 1 cm, it resulted in the most setup variation possibly caused by limited soft tissue contrast in the registration volume.

For tumours in the proximal and middle part of the oesophagus, setup margins increased when applying a soft tissue algorithm. For tumours in distal part, setup margins increased when applying a bone match algorithm. The position of the skin marks and tattoos in relation to the anatomy could be a possible explanation. However, evidence of these results is limited due to a small patient group with proximal or middle oesophagus tumours (n = 15).

It may be advisable to use region specific margins. The largest setup variations and required setup margins were found for

| Table 2 | Setup results, required setup margins and estimated number of setup corrections. |
|---------|----------------------------------------------------------------------------------|
| EPID    | CBCT (original clipbox) | CBCT (extended clipbox) | CBCT (mask = PTV+1 cm) |
| Group mean [mm] | | | |
| Lateral (RL) | Bone match | Clinical match | Soft tissue match | Bone match | Soft tissue match | Bone match | Soft tissue match | Bone match |
| -0.40 | 0.13 | 0.11 | 0.20 | 0.10 | -0.08 | 0.03 | -0.01 |
| Longitudinal (CC) | -0.03 | 0.33 | 0.27 | 0.28 | 0.01 | 0.33 | 0.58 | 0.46 |
| Vertical (AP) | 0.33 | 0.28 | 0.25 | 0.31 | 0.15 | 0.33 | 0.03 | -0.11 |
| Lateral | 1.44 | 1.78 | 1.78 | 1.90 | 1.69 | 1.89 | 1.82 | 2.47 |
| Longitudinal | 2.01 | 2.07 | 2.14 | 2.63 | 1.93 | 2.33 | 3.12 | 2.68 |
| Vertical | 1.49 | 1.42 | 1.46 | 1.64 | 1.88 | 1.73 | 1.77 | 1.51 |
| Lateral | 3.47 | 2.03 | 2.04 | 2.10 | 1.97 | 2.05 | 2.11 | 2.27 |
| Longitudinal | 3.97 | 3.38 | 3.38 | 3.40 | 3.21 | 3.34 | 3.53 | 3.09 |
| Vertical | 2.35 | 1.59 | 1.62 | 1.45 | 1.68 | 1.46 | 1.92 | 1.37 |
| Setup margin [mm] | | | | |
| Lateral | 6.04 | 5.86 | 5.88 | 6.23 | 5.62 | 6.17 | 6.02 | 7.78 |
| Longitudinal | 7.80 | 7.54 | 7.73 | 8.94 | 7.07 | 8.17 | 10.27 | 8.87 |
| Vertical | 5.37 | 4.67 | 4.79 | 5.13 | 5.88 | 5.36 | 5.76 | 4.72 |
| Average number of corrections using offline SAL protocol | 0.75 | 0.36 | 0.38 | 0.69 | 0.49 | 0.56 | 0.64 | 0.71 |

| Table 3 | Analysis of setup results and margins for different tumour sites. |
|---------|------------------------------------------------------------------|
| CBCT    | Clinical match |
|         | Proximal n = 8 | Middle n = 7 | Distal n = 35 |
| Soft tissue match | Bone match | Soft tissue match | Bone match | Soft tissue match | Bone match |
| Lateral | 1.78 | 1.52 | 2.00 | 1.21 | 1.13 | 1.78 | 1.90 |
| Longitudinal | 2.07 | 1.56 | 1.48 | 2.61 | 1.82 | 2.05 | 2.83 |
| Vertical | 1.42 | 1.41 | 2.23 | 1.70 | 1.82 | 1.42 | 1.49 |
| Lateral | 2.03 | 1.76 | 1.78 | 2.18 | 2.10 | 2.05 | 2.14 |
| Longitudinal | 3.38 | 2.42 | 1.51 | 2.51 | 2.47 | 3.62 | 3.72 |
| Vertical | 1.59 | 1.73 | 1.62 | 1.19 | 1.18 | 1.67 | 1.47 |
| Setup margin [mm] | | | | |
| Lateral | 5.86 | 5.03 | 6.24 | 4.54 | 4.29 | 5.88 | 6.24 |
| Longitudinal | 7.54 | 5.61 | 4.74 | 8.29 | 6.29 | 7.66 | 9.69 |
| Vertical | 4.67 | 4.73 | 6.71 | 5.09 | 5.37 | 4.71 | 4.75 |

| Table 4 | Analysis of differences between registration algorithms (mean ± SD). |
|---------|------------------------------------------------------------------|
| CBCT (original clipbox) | Proximal n = 8 | Middle n = 7 | Distal n = 35 |
| Absolute difference between soft tissue and bone match [mm] | | | |
| Lateral | 0.43 ± 0.22 | 0.68 ± 0.31 | 0.79 ± 0.63 |
| Longitudinal | 0.29 ± 0.25 | 1.03 ± 0.74 | 1.82 ± 1.29 |
| Vertical | 0.62 ± 0.60 | 0.41 ± 0.49 | 0.57 ± 0.44 |

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patients with distally located tumours. The absolute difference between soft tissue and bone registration were predominantly determined by patients with distal oesophageal tumours, where variations in breathing pattern and diaphragm position resulted in large differences, mainly in longitudinal direction. One frequently observed anatomical change was a change in diaphragm position. The possible cause for this change is a different breathing pattern. Jin et al. [13] recently investigated the respiration-induced oesophageal tumour motion and concluded that the largest tumour motion was in the distal part of the oesophagus, that tumour motion in the proximal part was limited, and that there is some need for individualised internal margins. Nowadays the Catharina Hospital standardly uses four-dimensional CT-scans for patients with a tumour of the distal part of the oesophagus or oesophagogastric junction and the individualised margins are based on tumour motion.

Another frequent anatomical change was reduction of GTV volume. Patients with different types of tumours, SCC and AC were included. Reduction of GTV volume was frequently observed for patients with a SCC. It is likely that this reduction is a result of tumour regression, but it is difficult to distinguish tumour from normal oesophagus on CBCT. Jin et al. [14] investigated the use of markers for setup verification. They concluded that markers can be useful to determine whether the coverage of the tumour is adequate or not, but the use of markers for setup verification is not feasible due to large tissue deformations during the treatment period.

The third frequent change observed was a reduction of heart volume. Lutkenhaus et al. [15] and Mohammad et al. [16] evaluated this reduction and found a significant reduction of the median heart volume of 8% over the radiotherapy treatment course. In the current study the reduction of heart volume was estimated in a similar way as Lutkenhaus et al. and Mohammad et al., and was found to be 6%.

Despite all possible anatomical variations, CBCT registration was in all cases considered to be accurate enough in order to confidently and accurately deliver the prescribed dose.

If reduction of GTV volume was observed, plan adaptation was considered not necessary because the CTV remained roughly unchanged. Only in a few cases, a new CT-scan was made to estimate the impact of the anatomical change on the treatment plan, but it never resulted in an actual adjustment or adaptation of the delivered plan. Nyeng et al. [17] investigated the dosimetric consequences of anatomical changes during treatment. An extra CT-scan was acquired during the treatment period at (median) fraction 10. These scans were deformably registered to the original planning-CT. A plan adaptation was performed when CTV coverage decreased >1% or PTV coverage decreased >3%. For nine out of twenty-nine patients, a plan adaptation was made. Main causes of plan adaptation were change in diaphragm position, mediastinal changes and bowel filling changes. Applied CTV-PTV margins in the study of Nyeng et al. were 5 mm in RL and AP direction and 8 mm in CC direction. In the current study isotropic margins of 1 cm were applied. A CTV coverage reduction is observed earlier when applying smaller margins. Moreover, Nyeng et al. performed registration on bony structures but in the current study soft tissue registration was performed. These differences could explain the need for plan adaptation in Nyeng’s study.

Another anatomical change in the current study and the study of Nyeng et al. was change in bowel filling and air in the oesophagus for some patients. All patients received concurrent chemotherapy, after which patients often drink carbonated drinks, which could explain the change in bowel filling and air in the oesophagus. Now patients are recommended to consume carbonated drinks only after the radiotherapy session.

In our study, the impact of the observed anatomical changes on the CTV coverage and dose in OAR was only a rough estimate based on clinical experience. For a more detailed analysis of the actually delivered dose to targets and OAR, further investigation is needed. A sophisticated method for this would be to apply dose calculations on the CBCT scans. This is currently not yet implemented in our hospital, and needs further refinement to accurately assess the impact of anatomical changes on the accuracy of the delivery of the prescribed dose.

### Conclusions

CBCT-based position verification reduces setup variations and corrections, resulting in a more consistent IGRT method for oesophageal cancer. Registration with a clipbox around PTV and vertebras and a soft tissue algorithm resulted in the smallest setup
margins and number of corrections. It is difficult to determine which registration (bone or soft tissue) is correct, since there is no gold standard.

Moreover, tumour location influences the required setup margins. For tumours of distal oesophagus or oesophagogastric junction, the required setup margin should be larger, especially in longitudinal direction, in comparison with tumours of more proximal parts of the oesophagus.

A large variety of anatomical changes is revealed when using kV-CBCT for oesophageal cancer patients, which cannot be observed using MV-EPID images. The most common anatomical changes were reduction of GTV volume, heart volume reduction and change in diaphragm position. The influence on tumour position was limited and applied PTV margins were still accurate. However, monitoring the influence of anatomical changes may become more important e.g. when applying smaller margins or plan adaptation.

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