Radiobiological evaluation of simultaneously dose-escalated versus non-escalated intensity-modulated radiation therapy for patients with upper thoracic esophageal cancer

Bao-Tian Huang1,2,*
Li-Li Wu1,3,*
Long-Jia Guo1,2,*
Liang-Yu Xu1,2,*
Rui-Hong Huang1
Pei-Xian Lin2
Jian-Zhou Chen1,3
De-Rui Li1
Chuang-Zhen Chen1
1Department of Radiation Oncology, Cancer Hospital of Shantou University Medical College, Shantou, 2Department of Nosocomial Infection Management, The Second Affiliated Hospital of Shantou University Medical College, Shantou, People’s Republic of China; 3CRUK/MRC Oxford Institute for Radiation Oncology, University of Oxford, Oxford, United Kingdom
*These authors contributed equally to this work

Objective: To compare the radiobiological response between simultaneously dose-escalated and non-escalated intensity-modulated radiation therapy (DE-IMRT and NE-IMRT) for patients with upper thoracic esophageal cancer (UTEC) using radiobiological evaluation.

Methods: Computed tomography simulation data sets for 25 patients pathologically diagnosed with primary UTEC were used in this study. DE-IMRT plan with an escalated dose of 64.8 Gy/28 fractions to the gross tumor volume (GTV) and involved lymph nodes from 25 patients pathologically diagnosed with primary UTEC, was compared to an NE-IMRT plan of 50.4 Gy/28 fractions. Dose-volume metrics, tumor control probability (TCP), and normal tissue complication probability for the lung and spinal cord were compared. In addition, the risk of acute esophageal toxicity (AET) and late esophageal toxicity (LET) were also analyzed.

Results: Compared with NE-IMRT plan, we found the DE-IMRT plan resulted in a 14.6 Gy dose escalation to the GTV. The tumor control was predicted to increase by 31.8%, 39.1%, and 40.9% for three independent TCP models. The predicted incidence of radiation pneumonitis was similar (3.9% versus 3.6%), and the estimated risk of radiation-induced spinal cord injury was extremely low (<0.13%) in both groups. Regarding the esophageal toxicities, the estimated grade ≥2 and grade ≥3 AET predicted by the Kwint model were increased by 2.5% and 3.8%. Grade ≥2 AET predicted using the Wijsman model was increased by 14.9%. The predicted incidence of LET was low (<0.51%) in both groups.

Conclusion: Radiobiological evaluation reveals that the DE-IMRT dosing strategy is feasible for patients with UTEC, with significant gains in tumor control and minor or clinically acceptable increases in radiation-induced toxicities.

Keywords: radiobiological evaluation, dose-escalated, non-escalated, intensity-modulated radiation therapy, esophageal cancer

Introduction

Esophageal cancer (EC) is the eighth most common cancer and the sixth leading cause of cancer-related mortality worldwide.1,2 Concurrent chemoradiotherapy (CCRT) has been established as the standard definitive treatment for locally advanced EC,3,4 with a total dose of 50 to 50.4 Gy delivered in a standard dose of 1.8 to 2 Gy per fraction.5,6 Unfortunately, the standard dose strategy is clinically disappointing because 90% of local failures still occur in the gross tumor volume (GTV),7 suggesting that a higher radiation dose to GTV might be more beneficial for improving local control.
The implementation of intensity-modulated radiation therapy (IMRT) can modify dose distribution and improve normal tissue sparing.\textsuperscript{8,9} Dose escalation using a simultaneous integrated boost (SIB) is a novel technique in which an accelerated higher dose, and a conventional lower dose can be delivered to the primary tumor and the subclinical disease simultaneously.\textsuperscript{10} Recently, an increasing number of studies have demonstrated its feasibility, efficacy, and safety for the treatment of EC patients with the combination of IMRT and dose escalation techniques.\textsuperscript{11–13} However, to the best of our knowledge, only several studies investigated the potential benefits of dose-escalated IMRT (DE-IMRT) to offer a dose escalation to the primary tumor, and improve the sparing of normal tissues compared with non-escalated IMRT (NE-IMRT) for EC patients.\textsuperscript{14–16} Unfortunately, the studies failed to provide further information on esophageal toxicity, which is a common radiation-induced side effect in the treatment of EC patients.\textsuperscript{11,12,17,18}

Radiobiological evaluation is a method to predict the outcome for the tumor and normal tissue using mathematical analysis with parameters derived from clinical trials. The method has an advantage in translating the dosimetric changes into radiobiological responses, and thus has been widely used to predict the feasibility of dose escalation for EC and primary prostate cancer.\textsuperscript{16,19}

In this study, we aimed to compare the radiobiological dose response between the DE-IMRT and NE-IMRT strategies across a group of representative patients suffering from EC, using radiobiological evaluation. We sought to evaluate the feasibility of DE-IMRT treatment for EC patients.

Materials and methods

Ethics statement

A total of 25 patients with pathologically confirmed upper thoracic esophageal cancer (UTEC) receiving CCRT were used in the study (ClinicalTrial.gov number, NCT01670409, and Chinese Clinical Research Registry number, ChiCTR-ONC-12002356). The Clinical Research Ethics Review Committee of Cancer Hospital of Shantou University Medical College approved this study. Only the computed tomography (CT) images of these patients were used in this study. All the patients have given their written informed consent for their CT images to be used before the study started. The patients' information in the CT images was de-identified to avoid any ethical concerns.

Immobilization and CT simulation

Patients were immobilized in the supine position with the head and shoulders wrapped in a thermoplastic mask and shoulders wrapped in a thermoplastic mask.

Target volume and OAR delineation

We have previously reported the approaches for target volume and OAR contouring in EC patients.\textsuperscript{14} Briefly, the GTV consisted of the primary tumor (GTV\textsubscript{p}) and the positive regional lymph nodes (GTV\textsubscript{LN}). The contouring of GTV was identified using CT images, endoscopic reports or barium swallow fluoroscopy. GTV\textsubscript{LN} included mediastinal or supraclavicular lymph nodes if the shortest axis was $\geq 10 \text{ mm}$. The clinical target volume (CTV) was delineated with 20 mm longitudinal and 5–10 mm radial margins with respect to the GTV\textsubscript{p} and a 5 mm uniform margin from GTV\textsubscript{LN}. Paraeosophageal or tracheoesophageal groove lymph nodes that did not meet the criteria of positive lymph nodes, but with shortest axis $\geq 5 \text{ mm}$, were also encompassed in CTV. Two planning target volumes (PTVs) PTV64.8 and PTV50.4 were generated with a uniform 5 mm margin expansion from CTV and GTV\textsubscript{LN}. OAR contours, including the spinal cord and lung, were created as previously described.\textsuperscript{20} Briefly, the lung contour was limited to the air-inflated lung parenchyma without inclusion of fluid and atelectasis visible on the CT images, excluding the proximal bronchial tree. The contour of the spinal cord started at the same cranial level as the esophagus to the bottom of L2 or at the level in which the cord ended. The planning OAR volume (PRV) for the spinal cord was generated by expanding the spinal cord to a 5 mm margin. The esophagus contour began from the level of cricoid cartilage on every CT image to the gastroesophageal junction.\textsuperscript{21–23}

Planning objectives

The dose constraints for OAR were as follows: spinal cord, maximum dose (D\textsubscript{max}) $< 45 \text{ Gy}$; PRV for spinal cord, V\textsubscript{50} $\leq 1 \text{ cc}$; lung, V\textsubscript{5} $< 60\%$, V\textsubscript{10} $< 50\%$, V\textsubscript{20} $< 30\%$ and mean lung dose $< 15 \text{ Gy}$, where V\textsubscript{x} is percentage of the target volume receiving $\geq x \text{ Gy}$. The dose was normalized to ensure that 95\% of the PTV received 100\% of the prescribed dose.
Treatment planning

The prescribed dose for the DE-IMRT plan was 64.8 Gy in 28 fractions for PTV64.8 (delivered in 2.31 Gy/fraction), and 50.4 Gy in 28 fractions for PTV50.4 (delivered in 1.8 Gy/fraction) according to Welsh et al. The prescribed dose for the NE-IMRT plan was set to 50.4 Gy in 28 fractions (delivered in 1.8 Gy/fraction) for the target of PTV50.4. Plans with different prescribed doses were generated using five coplanar sliding-window based IMRT fields with beam arrangements of 210°, 300°, 0°, 60°, and 150°. All plans were designed using a 6 MV photon beam from a TrueBeam linear accelerator (Varian Medical Systems). Optimization was performed using the dose volume optimizer (version 10.0.28) algorithm. Dose calculation was performed using the anisotropic analytical algorithm (version 10.0.28), considering the heterogeneity correction. Several dose-limiting structures were created to make the dose more conformal to the target. The method termed base dose function (BDF) described in our previous study was used to acquire more homogeneous dose distribution. Briefly, the fractions of the original plan were modified to half (from 28 to 14 fractions in the study), and then the half-prescribed plan was copied and re-optimized using the BDF incorporated in Eclipse with the half-prescribed plan as the base dose plan. When the dose was finally calculated, the fractions of the plan were doubled to generate the target plan.

Radiobiological evaluation

Both the tumor control probability (TCP) and normal tissue complication probability (NTCP) calculations were performed using in-house developed programs designed with MATLAB 7.0 (MathWorks, Natick, MA, USA). Three radiobiological models, including the Geh model, Webb-Nahum model (WN model), and equivalent uniform dose-based model (EUD model) were used to predict the TCP value. We emphasize that the Geh is a multivariate logistic based model (EUD model) were used to predict the TCP and NTCP, respectively. A comparison of the data was performed using a paired two-tailed Student’s t-test. A Wilcoxon matched-pair signed-rank test was alternatively used when the data did not follow a normal distribution. These results were considered statistically significant when P-value <0.05. The parameters used were n = 0.03, m = 0.03, and TD50 = 76.1 Gy in this model. All physical doses were converted to a biologically equivalent dose in 2 Gy fraction (EQD2) before modeling. An α/β value of 4.9 Gy was used for TCP prediction. The α/β values of 2 Gy and 3 Gy were used for NTCP calculation for the spinal cord and lung, respectively. With regard to the esophageal toxicity estimation, the α/β of 10 Gy and 3 Gy were employed for AET and LET prediction, respectively. A detailed procedure of modeling has been published in our previous work.

Statistical analysis

A comparison of the data was performed using a paired two-tailed Student’s t-test. A Wilcoxon matched-pair signed-rank test was alternatively used when the data did not follow a normal distribution. These results were considered statistically significant when P-value <0.05.

Results

Patient characteristics

CT simulation data sets of 25 UTEC patients, treated in our department from August 2012 to March 2014, were used in this study. The patients were between 49 and 73 years old.
The median volumes of GTV, PTV64.8, and PTV50.4 were 25.1±15.4 cc, 72.9±36.0 cc, and 201.3±62.5 cc, respectively. The patients’ characteristics are listed in Table 1.

### Dose difference between the DE-IMRT and NE-IMRT strategies

The mean dose-volume parameters averaging from 25 patients for the two dosing strategies are listed in Table 2. The DE-IMRT plan resulted in a 28.4% dose escalation to the GTV on average compared with the NE-IMRT plan (65.9 Gy versus 51.3 Gy). The $V_{10}\%$, $V_{20}\%$, and mean dose ($D_{\text{mean}}$) of the lung were increased by 1.0%, 1.3%, and 0.4 Gy, respectively ($P<0.05$). $D_{\text{max}}$ of the spinal cord was 2 Gy higher in the DE-IMRT group. With regard to the esophageal dose, the $V_{60}\%$, $V_{40}\%$, and $V_{50}\%$ of the esophagus were comparable between the DE-IMRT and NE-IMRT groups ($P>0.05$). However, the $V_{50}\%$, $D_{\text{mean}}$, and $D_{\text{max}}$ were dramatically increased in the DE-IMRT group ($P<0.001$). The dose volume histogram of the GTV, lung, spinal cord, and esophagus is presented in Figure 1. The dose distribution in the transversal, coronal, and sagittal views from one representative case is shown in Figure 2.

### Table 1 Characteristics of 25 patients with UTEC

| Patients | Gender | Age | Stage | GTV (cc) | PTV64.8 (cc) | PTV50.4 (cc) |
|----------|--------|-----|-------|----------|--------------|--------------|
| 1        | M      | 53  | T3N1M0 | 25.4     | 75.0         | 191.6        |
| 2        | M      | 64  | T3N1M1 | 38.8     | 102.1        | 253.3        |
| 3        | M      | 49  | T3N1M0 | 16.9     | 47.8         | 196.4        |
| 4        | M      | 64  | T3N0M1 | 16.1     | 45.5         | 161.9        |
| 5        | M      | 55  | T3N1M0 | 22.5     | 70.3         | 180.0        |
| 6        | M      | 73  | T2N0M1 | 13.8     | 38.3         | 168.2        |
| 7        | M      | 61  | T3N1M0 | 11.3     | 34.1         | 158.9        |
| 8        | M      | 59  | T2N0M1 | 22.4     | 58.5         | 174.7        |
| 9        | M      | 61  | T4N0M1 | 61.9     | 126.5        | 325.2        |
| 10       | M      | 59  | T3N1M0 | 13.3     | 46.3         | 160.2        |
| 11       | M      | 56  | T4N1M0 | 21.7     | 62.1         | 199.4        |
| 12       | F      | 53  | T3N1M0 | 8.4      | 27.9         | 110.5        |
| 13       | M      | 60  | T4N1M0 | 23.5     | 70.7         | 178.4        |
| 14       | M      | 64  | T3N0M1 | 20.8     | 60.4         | 200.7        |
| 15       | M      | 72  | T3N0M0 | 21.8     | 56.2         | 162.8        |
| 16       | M      | 66  | T4N0M0 | 67.0     | 137.1        | 359.2        |
| 17       | M      | 59  | T4N0M0 | 38.3     | 85.9         | 233.7        |
| 18       | M      | 59  | T3N1M0 | 11.9     | 146.2        | 35.9         |
| 19       | M      | 51  | T3N1M0 | 32.4     | 222.7        | 174.6        |
| 20       | M      | 67  | T3N0M0 | 44.1     | 267.8        | 98.9         |
| 21       | F      | 67  | T2N1M1 | 14.3     | 224.4        | 98.4         |
| 22       | F      | 65  | T3N0M0 | 14.9     | 135.9        | 83.3         |
| 23       | M      | 69  | T2N1M0 | 7.0      | 104.5        | 32.8         |
| 24       | M      | 61  | T3N1M0 | 36.8     | 299.6        | 95.0         |
| 25       | M      | 66  | T2N1M0 | 22.4     | 216          | 57.8         |

*According to American Joint Committee on Cancer (AJCC), 6th edition.

**Abbreviations:** UTEC, upper thoracic esophageal cancer; GTV, gross tumor volume; PTV, planning target volume; M, male; F, female.

### Table 2 Physical dose difference between the DE-IMRT and NE-IMRT plans

| Structures | Parameter | DE-IMRT | NE-IMRT | $P$-value |
|------------|-----------|---------|---------|-----------|
| GTV        | $D_{\text{max}}$ (Gy) | 65.9±0.3 | 51.3±0.3 | <0.001    |
|            | $V_{10}\%$ (%)     | 23.4±7.0 | 22.4±6.8 | <0.001    |
| Lung       | $V_{20}\%$ (%)     | 11.4±4.3 | 10.1±4.1 | 0.002     |
| Spinal cord| $D_{\text{max}}$ (Gy) | 39.2±3.2 | 37.2±3.2 | <0.001    |
| Esophagus  | $V_{60}\%$ (%)     | 64.7±1.7 | 64.6±1.6 | 0.852     |
|            | $V_{40}\%$ (%)     | 63.8±1.6 | 63.7±1.6 | 0.200     |
|            | $V_{50}\%$ (%)     | 62.2±1.7 | 62.1±1.7 | 0.058     |

*Notes: $P$-value was shown for the dosimetric differences between the DE-IMRT and NE-IMRT techniques. $P<0.05$ was identified as statistically significant.

**Abbreviations:** DE-IMRT, dose-escalated intensity-modulated radiation therapy; NE-IMRT, non-escalated intensity-modulated radiation therapy; GTV, gross tumor volume; PTV, planning target volume; MLD, mean lung dose; $D_{\text{mean}}$, mean dose; $D_{\text{max}}$, maximum dose; $V_{x}$, percentage of the target volume receiving $\leq x$ Gy.

### Table 3. TCP in the DE-IMRT plan was increased by 31.8% compared with the NE-IMRT plan

| Structures | Parameter | DE-IMRT | NE-IMRT | $P$-value |
|------------|-----------|---------|---------|-----------|
| GTV        | TCP (%) | 85.9±1.7 | 66.2±1.7 | <0.001    |

**Notes:** $P$-value was shown for the TCP differences between the DE-IMRT and NE-IMRT techniques. $P<0.05$ was identified as statistically significant.

**Abbreviations:** DE-IMRT, dose-escalated intensity-modulated radiation therapy; NE-IMRT, non-escalated intensity-modulated radiation therapy; GTV, gross tumor volume; TCP, tumor control probability; PTV, planning target volume; $D_{\text{max}}$, maximum dose; $V_{x}$, percentage of the target volume receiving $\leq x$ Gy.

### Discussion

After utilizing a standard dose of 50.4 Gy for definitive chemoradiation treatment for EC patients, 90% of local failures still occurred in the GTV. Therefore, the question was raised whether it is feasible to escalate the dose to the...
tumor to improve local control, while maintaining clinically acceptable side effects to normal tissues. Increasing evidence has demonstrated that the SIB technique is more clinically beneficial for EC patients;\textsuperscript{11–13} however, the difference in the radiobiological response, particularly for esophageal toxicity between the DE-IMRT and NE-IMRT strategies for UTEC patients, remains unknown. To the best of our knowledge, this is the first study to compare tumor control and esophageal toxicity between two dosing strategies using radiobiological evaluation. We found that it is feasible to apply the DE-IMRT technique for UTEC patients, with significant gains (31.8\%–40.9\%) for tumor control, while minor increased toxicity was observed for the lung and spinal cord, and acceptable toxicities were found for the esophagus.

The results of this study are consistent with previously reported dosimetric data, which identified an improvement in dose escalation to the tumor using the DE-IMRT strategy. Welsh et al reported a planning study in distal EC patients using the SIB technique to deliver a boosted dose to the GTV, using the same dose prescription employed in our study. They found that the use of DE-IMRT produced a 28\% dose escalation (from 52 Gy to 66.9 Gy) to the GTV, while providing a comparable dose to the lungs and a slightly higher dose to the spinal cord compared with the conventional dose plan.\textsuperscript{15} Warren et al administered a boosted dose (62.5 Gy in 25 fractions) to the tumor in mid-EC patients, and found that significant gains (>18\%) in tumor control were achieved compared with the non-escalated plan using radiobiological evaluation.\textsuperscript{16} Similar to patients with middle and distal EC, our results demonstrated that the DE-IMRT technique was also applicable for UTEC patients. Apart from dosimetric investigations, an increase in the number of clinical trials has also

---

**Figure 1** DVH of the GTV, lung, spinal cord, and esophagus.

**Notes:** (A) DVH of GTV; (B) DVH of lung; (C) DVH of spinal cord; (D) DVH of esophagus.

**Abbreviations:** DVH, dose volume histogram; GTV, gross tumor volume; DE-IMRT, dose-escalated intensity-modulated radiation therapy; NE-IMRT, non-escalated intensity-modulated radiation therapy.
Table 3 Prediction of tumor control and toxicities in the DE-IMRT and NE-IMRT plans

| Structures     | Model     | DE-IMRT | NE-IMRT | P-value |
|----------------|-----------|---------|---------|---------|
| Tumor          | Geh (%)   | 67.6±12.2| 35.8±13.0| <0.001 |
|                | WN (%)    | 91.6±1.6 | 52.5±5.5 | <0.001 |
|                | EUD (%)   | 95.1±0.2 | 54.2±1.4 | <0.001 |
| Lung           | LKB (%)   | 3.9±0.0  | 3.6±0.9  | <0.001 |
| Spinal cord    | LKB (%)   | (7.2±0.2)×10⁻²| (4.2±3.2)×10⁻²| <0.001|
|                | Kwint (%) | 75.6±5.8 | 73.1±6.1 | <0.001 |
|                | Kwint (%) | 38.2±8.6 | 34.4±8.3 | <0.001 |
| Esophagus      | Wijsman (%)| 64.7±13.9| 49.8±11.1| <0.001 |
|                | Chen (%)  | (2.2±2.9)×10⁻¹| (1.7±1.8)×10⁻¹| <0.001|

Notes: *indicates TCP models for predicting tumor local control; †indicates LKB model for predicting grade ≥2 radiation pneumonitis; ‡indicates LKB model for predicting radiation-induced myelitis/necrosis; ††indicates Kwint model for predicting grade ≥2 acute esophageal toxicity; †‡indicates Kwint model for predicting grade ≥3 acute esophageal toxicity; †††indicates Wijsman model for predicting grade ≥2 acute esophageal toxicity; ‡‡‡indicates Chen model for predicting late esophageal toxicity. P-value was shown for the tumor control and toxicity differences between the DE-IMRT and NE-IMRT techniques. P<0.05 was identified as statistically significant.

Abbreviations: DE-IMRT, dose-escalated intensity-modulated radiation therapy; NE-IMRT, non-escalated intensity-modulated radiation therapy; TCP, tumor control probability; Geh, Geh model; WN, Webb-Nahum model; EUD, equivalent uniform dose model; LKB, Lyman-Kutcher-Burman model; Kwint, Kwint model; Wijsman, Wijsman model; Chen, Chen model.

Figure 2 Dose distributions in the transversal, coronal, and sagittal views from one representative case.

Notes: (A) NE-IMRT. (B) DE-IMRT. The dark green line represents PTV50.4, the red line represents PTV64.8, and the orange line represents GTV.

Abbreviations: PTV, planning target volume; GTV, gross tumor volume; DE-IMRT, dose-escalated intensity-modulated radiation therapy; NE-IMRT, non-escalated intensity-modulated radiation therapy.

Our analysis was partly dependent on the choice of radio-biological models and the parameters used. To strengthen the reliability of our data, we used three TCP predicting models and two esophageal toxicity models from the literature, to predict the tumor control and likelihood of grade ≥2 AET. Interestingly, three independent models exhibited a similar trend of TCP increase using the DE-IMRT method, although demonstrated the feasibility and efficacy of the DE-IMRT technique for EC patients, with encouraging improvement in local control and well tolerated acute toxicities, although the escalated dose was slightly different. Both Yu et al and Chen et al initiated a phase II study to investigate the efficacy of radiation dose escalation using the DE-IMRT technique combined with chemotherapy, for the treatment of EC patients. The results showed a similar trend of improvement in local-regional control and overall survival, indicating that the method of theoretical evaluation could obtain results that were similar to those obtained in clinical practices. Taken together, these described studies theoretically and practically validate the efficacy of dose escalation for EC patients.

Our analysis was partly dependent on the choice of radio-biological models and the parameters used. To strengthen the reliability of our data, we used three TCP predicting models and two esophageal toxicity models from the literature, to predict the tumor control and likelihood of grade ≥2 AET.
the changes were different among the models. Given that the Geh model is a multivariate logistic regression model fitted to the data from 26 chemoradiotherapy trials that considers many factors, such as the dose of radiotherapy, duration, chemotherapy (cisplatin and 5-FU) dose, and age, we believe that our data in TCP prediction is reliable. Regarding esophageal toxicity, both the Kwint and Wijsman models displayed an increase in grade $\geq 2$ AET using the DE-IMRT technique. However, we found that when using the Wijsman model, this increase was much more significant than that obtained in the Kwint model. This finding might be partly due to the different chemotherapy administration and radiation techniques in the two independent investigations: CCRT was used for all the patients in the Kwint model, while only selective patients with good performance status received CCRT and most patients underwent sequential treatment or radiotherapy alone in the Wijsman model. Moreover, the concurrent chemotherapy regimen consisted of low-dose intravenous cisplatin in the Kwint model, whereas gemcitabine combined with cisplatin or etoposide combined with cisplatin were delivered in sequential chemotherapy in the Wijsman model. Since CCRT has been widely reported to be associated with the incidence of AET in many studies, the difference in chemotherapy practice might greatly have affected our esophageal toxicity prediction. On the other hand, all of the patients were treated with IMRT in the Kwint model, whereas both IMRT and volumetric-modulated arc therapy were used in the Wijsman model. Radiotherapy technique was reported be significantly correlated with grade $\geq 2$ AET and it might partly influence the esophageal toxicity prediction. We preferred the prediction obtained from the Kwint model, because CCRT with IMRT administration is the recommended strategy for the definitive treatment of EC patients in most publications, however, further clinical data are needed to validate the model.

AET and LET characterized by dysphagia, odynophagia, stenosis, and perforation are common radiation-induced adverse events that significantly affect quality-of-life, and negatively affect long-term survival when receiving thoracic irradiation. In the definitive treatment of EC patients, the esophagus is more prone to developing these symptoms because a part of the esophagus is located in the radiation field, resulting in high-dose exposure during treatment. It was reported that approximately 40% of patients developed grade 3 AET after SIB radiation dose treatment. Unfortunately, modeling of the incidence of esophageal toxicity in EC patients is challenging due to the following two reasons. 1) Extensive publications have developed dosimetric predictors or predicting models to estimate the incidence of AET or LET in NSCLC patients; however, none of these studies have yet developed models derived from EC cases. Our group completed a phase II study, and the establishment of an esophagitis-predicting model first derived from EC patients is still ongoing. 2) There is no consensus on the contouring of the esophagus in EC patients. Whether the tumor and the esophagus outside the treatment field should be included, still remains unclear. Caglar et al found that the esophagus in-field is a new predictor for esophagitis in NSCLC patients. However, other studies have used the entire esophagus as the predictor. Given that both the Kwint and Wijsman models were generated from the entire esophagus, our study used this information to maintain consistency with the original work. However, whether different definitions of the esophagus will affect the evaluation of esophageal toxicity remains an important issue that needs to be addressed in our future work.

In the past few years, many studies have proposed dosimetric predictors to correlate with the occurrence of grade $\geq 2$ or grade $\geq 3$ AET. Unfortunately, only several studies have developed models to predict their incidence. Furthermore, most of the developed models were generated from the 3-dimensional conformal radiation therapy (3DCRT) technique. Because IMRT has been reported to be more superior in delivering a conformal dose and improving normal tissue sparing than 3DCRT, the models generated from 3DCRT might potentially limit the evaluation of AET for IMRT-treated patients. To date, AET predicting models derived from patients undergoing IMRT and chemotherapy are scarce, except for the Kwint and Wijsman models. Consistent with this information, we had to use these models for the prediction of esophageal toxicity in the study.

Several SIB dosing regimens, including 62.5 Gy/25 fractions, 63 Gy/28 fractions, 64.8 Gy/28 fractions, and 66 Gy/30 fractions, have been demonstrated to provide improved local control and clinically acceptable side effects for EC patients. However, whether a variety of dosing regimens may translate into dose-response differences, is still questionable. Consequently, we calculated the EQD$_2$ for the tumor and esophagus from different dosing regimens. We found that the EQD$_2$ for the tumor was 67 Gy, 65.3 Gy, 67.7 Gy, and 67.9 Gy, respectively, using an $\alpha/\beta$ value of 4.9 Gy. Similarly, the calculated EQD$_2$ for the esophagus was 65.1, 64.3, 66.5, and 67.1 Gy, respectively, using an $\alpha/\beta$ value of 10 Gy. This result suggests that comparable EQD$_2$ is acquired for both the tumor and the esophagus among the...
Conclusion
In summary, our study has demonstrated the benefits of using the DE-IMRT strategy to significantly improve tumor control with minor side effects, or clinically acceptable toxicities for UTEC patients. These findings require further validation in clinical samples.

Acknowledgments
This work was sponsored by National Natural Science Foundation of China (81602667), Science and Technology Planning Project of Guangdong Province (2016ZC0166), Medical Scientific Research Foundation of Guangdong Province (A2015534), Shantou University Medical College Clinical Research Enhancement Initiative (201424), Collaborative and Creative Center, Molecular Diagnosis and Personalized Medicine, Shantou University, Guangdong Province, People’s Republic of China.

Author contributions
Each author had participated sufficiently in the work to take public responsibility for appropriate portions of the content. BTH and CZC conceived and designed the experiments. LLW, LJG, LYX, and JZC performed the experiments. BTH, LLW, LJG, LYX, and RHH collected the data. PXL analyzed the data. BTH wrote the paper. DRL and CZC revised the paper. All authors contributed toward data analysis, drafting and revising the paper and agree to be accountable for all aspects of the work.

Disclosure
The authors report no conflicts of interest in this work.

References
1. Siegel R, Naishadham D, Jemal A. Cancer statistics, 2013. CA Cancer J Clin. 2013;63(1):11–30.
2. Pennathur A, Gibson MK, Jobe BA, Luketich JD. Oesophageal carcinoma. Lancet. 2013;381(9864):400–412.
3. Cooper JS, Guo MD, Herskovic A, et al. Chemoradiotherapy of locally advanced esophageal cancer: long-term follow-up of a prospective randomized trial (RTOG 85-01). Radiation Therapy Oncology Group. JAMA. 1999;281(17):1623–1627.
4. Herskovic A, Martz K, al-Sarraf M, et al. Combined chemotherapy and radiotherapy compared with radiotherapy alone in patients with cancer of the esophagus. N Engl J Med. 1992;326(24):1593–1598.
5. Minsky BD, Pajak TF, Ginsberg RJ, et al. INT 0123 (Radiation Therapy Oncology Group 94-05) phase III trial of combined-modality therapy for esophageal cancer: high-dose versus standard-dose radiation therapy. J Clin Oncol. 2002;20(5):1167–1174.
6. National Comprehensive Cancer Network. Esophageal and Esophagogastric Junction Cancers. National Comprehensive Cancer Network; 2015. Available from: https://www.nccn.org/store/login/login.aspx?ReturnURL=https://www.nccn.org/professionals/physician_gls/pdf/esophageal.pdf
7. Welsh J, Settle SH, Amini A, et al. Failure patterns in patients with esophageal cancer treated with definitive chemoradiation. Cancer. 2012;118(10):2632–2640.
8. Chandra A, Guerrero TM, Liu HH, et al. Feasibility of using intensity-modulated radiotherapy to improve lung sparing in treatment planning for distal esophageal cancer. Radiother Oncol. 2005;77(3):247–253.
9. Fenkell L, Kaminsky I, Breen S, Huang S, Van Proosijen M, Ringash J. Dosimetric comparison of IMRT vs. 3D conformal radiotherapy in the treatment of cancer of the cervical esophagus. Radiother Oncol. 2008;89(3):287–291.
10. Leclerc M, Maingon P, Hamoir M, et al. A dose escalation study with intensity modulated radiation therapy (IMRT) in T2N0, T2N1, T3N0 squamous cell carcinomas (SOC) of the oropharynx, larynx and hypopharynx using a simultaneous integrated boost (SIB) approach. Radiother Oncol. 2013;106(3):333–340.
11. Yu WW, Zhu ZF, Fu XL, et al. Simultaneous integrated boost intensity-modulated radiotherapy in esophageal carcinoma: early results of a phase II study. Strahlenther Onkol. 2014;190(11):979–986.
12. Yu W, Cai XW, Liu Q, et al. Safety of dose escalation by simulta neous integrated boosting radiation dose within the primary tumor guided by (18)FDG-PET/CT for esophageal cancer. Radiother Oncol. 2015;114(2):195–200.
13. Chen J, Guo H, Zhai T, et al. Radiation dose escalation by simultaneous modulated accelerated radiotherapy combined with chemotherapy for esophageal cancer: a phase II study. Oncotarget. 2016;7(16):22711–22719.
14. Zhang WZ, Chen IZ, Li DR, et al. Simultaneous modulated accelerated radiation therapy for esophageal cancer: a feasibility study. World J Gastroenterol. 2014;20(38):13973–13980.
15. Welsh J, Palmer MB, Ajani JA, et al. Esophageal cancer dose escalation using a simultaneous integrated boost technique. Int J Radiat Oncol Biol Phys. 2012;82(1):468–474.
16. Warren S, Partridge M, Carrington R, Hure C, Crosby T, Hawkins MA. Radiobiological determination of dose escalation and normal tissue toxicity in definitive chemoradiation therapy for esophageal cancer. Int J Radiat Oncol Biol Phys. 2014;90(2):423–429.
17. Higuchi K, Komori S, Tanabe S, et al. Definitive chemoradiation therapy with docetaxel, cisplatin, and 5-fluorouracil (DCF-R) in advanced esophageal cancer: a phase 2 trial (KODO 0501-P2). Int J Radiat Oncol Biol Phys. 2014;89(4):872–879.
18. Roeder F, Nicolay NH, Nguyen T, et al. Intensity modulated radiotherapy (IMRT) with concurrent chemotherapy as definitive treatment of locally advanced esophageal cancer. Radiat Oncol. 2014;9:191.
19. Kuang Y, Wu L, Hirata E, Miyazaki K, Sato M, Kwee SA. Volumetric modulated arc therapy planning for primary prostate cancer with selective intraprostatic boost determined by 18F-choline PET/CT. Int J Radiat Oncol Biol Phys. 2015;91(5):1017–1025.
20. Kong FM, Ritter T, Quint DJ, et al. Consideration of dose limits for organs at risk of thoracic radiotherapy: atlas for lung, proximal bronchial tree, esophagus, spinal cord, ribs, and brachial plexus. Int J Radiat Oncol Biol Phys. 2011;81(5):1442–1457.
21. Caglar HB, Othus M, Allen AM. Esophagus in-field: a new predictor for esophagitis. *Radiother Oncol*. 2010;97(1):48–53.
22. Kwint M, Uyterlinde W, Nijkamp J, et al. Acute esophagus toxicity in lung cancer patients after intensity modulated radiation therapy and concurrent chemotherapy. *Int J Radiat Oncol Biol Phys*. 2012;84(2):e223–e228.
23. Wijisman R, Dankers F, Troost EG, et al. Multivariable normal-tissue complication modeling of acute esophageal toxicity in advanced-stage non-small-cell lung cancer patients treated with intensity-modulated (chemo-)radiotherapy. *Radiother Oncol*. 2015;117(1):49–54.
24. Lu JY, Cheung ML, Huang BT, et al. Improving target coverage and organ-at-risk sparing in intensity-modulated radiotherapy for cervical oesophageal cancer using a simple optimisation method. *PLoS One*. 2015;10(3):e0121679.
25. Geh JI, Bond SJ, Bentzen SM, Glynn-Jones R. Systematic overview of preoperative (neoadjuvant) chemoradiotherapy trials in oesophageal cancer: evidence of a radiation and chemotherapy dose response. *Radiother Oncol*. 2006;78(3):236–244.
26. Webb S, Nahum AE. A model for calculating tumour control probability in radiotherapy including the effects of inhomogeneous distributions of dose and clonogenic cell density. *Phys Med Biol*. 1993;38(6):653–666.
27. Bedford JL, Viviers L, Guzel Z, Childs PJ, Webb S, Tait DM. A quantitative treatment planning study evaluating the potential of dose escalation in conformal radiotherapy of the oesophagus. *Radiother Oncol*. 2000;57(2):183–193.
28. OkuneifTP, Morgan D, Niemierko A, Suit HD. Radiation dose-response of human tumors. *Int J Radiat Oncol Biol Phys*. 1995;32(4):1227–1237.
29. Seppenwoolde Y, Lebesque JV, de Jaeger K, et al. Comparing different NTCP models that predict the incidence of radiation pneumonitis. Normal tissue complication probability. *Int J Radiat Oncol Biol Phys*. 2003;55(3):724–735.
30. Burman C, Kucher GJ, Emami B, Goitein M. Fitting of normal tissue tolerance data to an analytic function. *Int J Radiat Oncol Biol Phys*. 1991;21(1):123–135.
31. Chen C, Uyterlinde W, Sonke JJ, de Bois J, van den Heuvel M, Belderbos J. Severe late esophagus toxicity in NSCLC patients treated with IMRT and concurrent chemotherapy. *Radiother Oncol*. 2013;108(2):337–341.
32. Huang BT, Lu JY, Lin PX, Chen JZ, Li DR, Chen CZ. Radiobiological modeling analysis of the optimal fraction scheme in patients with peripheral non-small-cell lung cancer undergoing stereotactic body radiotherapy. *Sci Rep*. 2015;5:18010.