Estimation of radiotherapy modalities for patients with stage I-II nasal natural killer T-Cell lymphoma

Xianfeng Liu¹, Furong Wu¹, Qishuai Guo¹, Ying Wang¹, Yanan He¹, Huanli Luo¹, Qicheng Li¹, Mingsong Zhong¹, Chao Li¹, Han Yang¹, Juan Zhou², Fu Jin¹

¹Department of Radiation Oncology, Chongqing University Cancer Hospital, Chongqing Cancer Institute, Chongqing Cancer Hospital, Chongqing, People’s Republic of China; ²Forensic Identification Center, College of Criminal Investigation, Southwest University of Political Science and Law, Chongqing, People’s Republic of China

Purpose: The objective of this study is to estimate radiotherapy (RT) modalities for patients with stage I-II nasal natural killer T-Cell lymphoma (NNKTCL), including plan quality, radiation delivery efficiency, cost of RT and excess absolute risk (EAR).

Materials and methods: Twenty-four representative patients with stage I-II NNKTCL treated with fix-field intensity-modulated radiotherapy (FF-IMRT) were re-planned for volumetric modulated arc therapy (VMAT), TomoDirect (TD) and TomoHelical (TH) on the TomoHDA system, respectively. Plan characteristics, cost of RT and EAR were compared.

Results: Compared with IMRT, TD and TH showed significant improvement in terms of $D_{\text{max}}$, $D_{2%}$, cold spot volume and homogeneity index (HI) of planning target volume (PTV), while achieving worse $D_{\text{mean}}$ and conformity index (CI). The mean dose of oropharynx, thyroid and left salivary, and the maximum dose of right salivary by TD (249.20%, $p=0.000$; 52.94%, $p=0.000$; 160.23%, $p=0.022$; 122.67%, $p=0.027$), VMAT (15.76%, $p=0.000$; 23.53%, $p=0.000$; 34.09%, $p=0.000$; 31.33%, $p=0.000$) and TH (250.32%, $p=0.000$; 58.82%, $p=0.000$; 120.45%, $p=0.020$; 117.33%, $p=0.032$) increased significantly compared to IMRT. VMAT reduced treatment time ($p=0.000$; 0.000; 0.000) and monitor units (MUs) ($p=0.000$; 0.000; 0.000) obviously compared with IMRT, TD and TH. The cost of RT for TD and TH increased 150% compared with IMRT and VMAT. IMRT obtained the lowest EAR to oropharynx, thyroid, left and right salivary gland in the four treatment modalities.

Conclusion: The results show that both TD and TH can achieve higher conformal target quality while getting worse organs at risk (OARs) sparing and EAR to some organs than IMRT for patients with stage I-II NNKTCL. IMRT delivers the lowest dose to most OARs, VMAT requires the lower cost of RT and shortest delivery time, and TH obtained the optimal target coverage. The results could provide direction for selecting proper RT modalities for different cases.

Keywords: nasal natural killer T-Cell lymphoma, excess absolute risk, IMRT, VMAT, TomoHelical

Background

Nasal natural killer T-Cell lymphoma (NNKTCL) usually occurs in Eastern Asia, Mexicans and South Americans, and accounts for <10% of primary non-Hodgkin lymphoma disease.¹⁻³ Radiotherapy (RT) is recommended as the main therapy method for stage I-II NNKTCL, and previous research has shown an overall survival benefit in favor of RT for patients with the NNKTCL.²⁻⁴⁻⁵

To achieve better target coverage and organs at risk (OARs) sparing, and improve the control rates and reduce the treatment morbidity subsequently, various RT methods such as medical linear accelerator-based fix-field intensity-modulated radiotherapy (FF-IMRT) and volumetric modulated arc therapy (VMAT), TomoHDA system-
based TomoHelical (TH) are all used to treat NNKTCL, showing that each method has both advantage and disadvantage at target dose coverage, OARs sparing, and therapeutic efficiency, respectively, of which the out-of-field dose also differs significantly.3,6,7

Exposure to therapeutic doses of RT unavoidably leads to radiation damage and toxicity for normal tissue, and previous literature have shown therapy-related second cancer risk (SCR) as a late complication of treatment becomes obvious for patients who received RT previously, although the precise-related data remain unknown.8-10 Model excess absolute risk (EAR) developed by Schneider can be utilized to estimate SCR for normal tissues, which are usually unavoidably irradiated during a radiation treatment.11,12 Currently, advances in RT have resulted in large cases of long-term cancer survivors, and late sequelae of RT are becoming the next core concern. Hence, SCR remains studying further, especially involving younger patients who potentially have a longer life expectancy.13,14

Clinically, selecting the optimal treatment modality for patients with stage I-II NNKTCL has crucial importance, previous studies showed that each modality has its own characteristic in plan quality.3,6,7 However, the SCR and cost of RT triggered by various RT modalities have not studied. The purpose of this study is to compare the plan quality, radiation delivery efficiency, cost of RT and SCR among four treatment modalities (FF-IMRT, VMAT, TH and TomoDirect[TD]) for patients with stage I-II NNKTCL, and provide guide for choosing reasonable treatment modalities.

Materials and methods

Patients and materials

Planning computer tomography (CT) datasets of 24 different patients with stage I-II NNKTCL who were recently treated at our Institution were used for this study. This study was approved by the Chongqing Cancer Hospital’s ethics committee, and written informed consent was obtained from each patient. Diversity in shape and stage of the treated NNKTCL in the selected patients were ascertained, with 22 stage I and 2 stage II cases. The 24 patients were without distant metastases and had not received prior RT.

The CT scans were acquired on a Philips Brillance Big Bore CT (Philips, Holland) simulation in 3 mm slice thickness, in the supine position with the scan scope from the inferior margin of the clavicular heads to the vertex of the skull. Image registration and delineation of the target and OARs were performed using the Eclipse treatment planning system (TPS, Varian Medical Systems, Version 11.0, Inc.). Gross tumor volume (GTV) contained primary tumor and regional metastatic lymph nodes, diagnosed by magnetic resonance imaging or CT, endoscopic examination and physical examination. Clinical target volume (CTV) included GTV and adjacent tissues at risk for contiguous spread. Planning target volume (PTV) was obtained from the respective CTV expanding symmetrically by 3 mm in all dimensions to offset setup errors. The mean volume of PTV was 248.8 cm3, ranging from 60.7 to 471.3 cm3. Surrounding OARs of lenses, eyes, optic nerves, optic chiasm, brainstem, spinal cord, parotid glands, salivary gland, oropharynx and thyroid gland were contoured. All CT images and delineation of the target and OARs in the Eclipse TPS were conveyed to the tomotherapy TPS (Tomotherapy, Madison, WI, USA).

RT plans

For each of the 24 cases, four different planning modalities were adopted: FF-IMRT, VMAT, TH and TD. FF-IMRT and VMAT plans were designed on the Eclipse TPS. For all Eclipse plans, dose calculations were done with 6 MV photon beams generated by Varian IX linear accelerator, Progressive Resolution Optimizer and Dose–Volume Optimizer algorithms were utilized for VMAT and FF-IMRT optimizations, respectively.15 Anisotropic analytical algorithm was used for final dose calculations.16 TH and TD plans were designed on tomotherapy TPS, utilizing 6 MV photon beams, a fixed dose rate (DR) of 877 monitor units (MUs)/min and least squares optimization method.17,18 A 0.5 cm bolus was added for all plans optimizations, the inner edges of the paropias on both sides were the right and left bounds of the bolus, while the lower edge of the palate and the upper edge of the frontal sinus were the lower and upper bounds of the bolus, respectively.

FF-IMRT

All FF-IMRT plans owned 10 coplanar fields, where the gantry angles were 200°, 240°, 280°, 320°, 0°, 0°, 40°, 80°, 120° and 160°, respectively. The angles of collimator and couch rotation for all fields were set to 0°. To minimize the dose to the lenses and eyes, the position of jaws in the two fields of 0° should be adjusted. After fitting the jaws to the structure of PTV with a margin of 6 mm, the position of the upper jaw of one field with gantry angle 0° was adjusted to 3 mm below the lower edge of the lenses,
and the position of the left and right jaws of the other field
with gantry angle 0° was adjusted to keep the lenses out of
radiation field. And dynamic sliding-window IMRT deliv-
ery technique and a fixed DR of 400 MUs/min were
adopted to deliver the dose.

VMAT
All VMAT plans were set as two coplanar arcs of 360°, with
the collimator rotation of 45° and 315°, respectively, and the
couch rotation of 0°. The maximum DR of all plans was set to
600 MUs/min.

TH
All the TH plans were optimized with a field width of 2.5 cm,
pitch values of 0.287 and modulation factor of 3, and dose
calculated with fine dose calculation grid. In addition, the
function of complete block was adopted to protect the lenses.

TD
The TD plans had nine equally distributed beam, with the
angel of 200°, 240°, 280°, 320°, 0°, 40°, 80°, 120° and
160°, respectively. The optimization and calculation set-
tings of the TD plans were the same as the TH plans.
Similarly, the function of complete block was used to
minimize the dose to the lenses.

For the PTV of all plans, the prescribed dose was 50 Gy in
25 fractions, the prescribed 95% isodose covered no <95% of
the PTV, and the percentage volume of PTV receiving over
107.5% of the prescription dose was no more than 2%.7,19
Based on the Radiation Therapy Oncology Group 0615
Protocol,20 dose constraints for the four types of plans were
slightly modified. Dose constraint for OARs was defined as
follows: maximum dose (Dmax) of eyes, optic nerves, optic
chiasm and brainstem were limited to <50 Gy; Dmax of spinal
cord and lenses were limited to 45 and 15 Gy, respectively;
mean dose (Dmean) of parotid glands was limited to <26 Gy;
Dmean of salivary gland, oropharynx and thyroid gland were
limited to <40 Gy.

Treatment plan evaluation
All the data acquired from Dose–Volume Histogram
(DVH) of the 96 plans were analyzed. Figures 1 and 2
show representative dose distribution and DVHs for the
four types of modalities, respectively. The plan evaluation
focused on the items as in the following:

PTV

The dose received by 2%, and 98% (D2% and D98%) of the
PTV volume were defined as the maximum and minimum
dose of PTV, respectively. The Dmean of PTV was also used.
The percentage volumes of PTV receiving <93% of the pre-
scribed dose were defined as the cold spot volume of PTV,
which should be <1%. The homogeneity index (HI) and con-
formity index (CI) of the PTV were compared by treatment
modalities. CI was evaluated using the Paddick conformity
index, defined as CI=TV2PTV/(TV×PIV), where TV was the
PTV volume, PIV was the body volume receiving 95% of the
prescription dose and TVPTV was the PTV volume covered by
95% of the prescribed dose. Ranging from 0 to 1, the value of
CI approaching 1 meant fine conformity of PTV. HI repre-
sented dose uniformity, calculated with D5% minus D95%÷
divided by Dmean, where D5% and D95% was the minimum
dose delivered to 5% and 95% of the PTV. Lower HI indicated
better uniformity irradiation of the target volume.

Organs at risk
For all the plans, dosimetric analysis was executed for the
Dmax, and Dmean, of OARs, including left and right lens, left
and right eye, left and right optic nerve, left and right
parotid gland, left and right salivary gland, thyroid gland,
optic chiasm, oropharynx, spinal cord and brainstem.

MUs and treatment time
The total MUs and treatment time were compared for the
four kinds of modalities. Treatment time was calculated
only from beam-on to the end of total MUs delivery,
including the time of gantry rotation and dose delivery.

Cost of RT
The cost of RT through the whole radiation treatment process
includes the expense of simulation, delineation, dose calcula-
tion, treatment delivery and quality assurance. In People's
Republic of China, different provinces have different RT
costs. In our western region, under the policies regulation
of the Chongqing price bureau, the four treatment modalities
only have different cost of treatment delivery in our institute,
so we only compare the cost of treatment delivery among
four treatment modalities in this study.

EAR calculations
For a person exposed to a radiation dose, the SCR of an
organ can be obtained from Equation (1) utilizing the EAR
model.11,21

\[
\text{EAR}^\text{org} = \frac{1}{V_T} \sum_i V(D_i) \beta_{\text{EAR}} \text{RED}(D_i) \mu(x, a)
\]  

where \( V_T \) represented the total organ volume, \( V(D_i) \) was the
organ volume to receive a dose \( D_i \), and \( \beta_{\text{EAR}} \) represented the
slope of the dose–response curve at low dose. RED(D) was the dose–response mechanistic model which accounts for cell killing and fractionation effects described as Equation (2), \( \mu(x, a) \) was the modifying function given by Equation (3).

\[
\text{RED}(D) = \frac{e^{-D/\alpha}}{\alpha R} \left( 1 - 2R + R^2 e^{D/\alpha} - (1 - R)^2 e^{-D/\alpha} \right)
\]

where \( R \) was the repopulation/repair parameter, which characterized the repopulation/repairability of the tissue between two dose fractions, and \( \alpha \) was given by Equation (4).

\[
\mu(x, a) = e^{x\gamma_e + y\gamma_a / 70}
\]

where \( \gamma_e \) and \( \gamma_a \) were the age modifying parameters.

\[
\alpha' = \alpha + \beta d = \alpha + \beta D_T / D_T d_T
\]

where \( D_T \) and \( d_T \) represented the prescribed dose to the target volume with the corresponding fractionation dose, respectively.

The EAR has been investigated to the organs of oro-pharynx, thyroid, left and right salivary gland. The reported values of \( \alpha \) and \( \alpha/\beta \) were applied in this study. As far as we know, there is no published value of \( R \) for thyroid, therefore, we set \( R=0.5 \) which represented intermediate repopulation just like the previous literature did. \( R \) is 0.5 which represented intermediate repopulation just like the previous literature did. \( R \) was 0.5 which represented intermediate repopulation just like the previous literature did. All other parameters utilized in EAR calculations in this study were selected from published research. \( R \) is 0.5 which represented intermediate repopulation just like the previous literature did. All the parameters are shown in Table 1.

**Statistical analysis**

Utilizing the SPSS statistical software (SPSS, Chicago, IL, USA), paired t-test was carried out for determining if there was a significant difference in each of the pair parameters. A \( p<0.05 \) was considered statistically significant.

**Results**

PTV coverage

\( D_{98\%} \), \( D_{2\%} \), \( D_{\text{mean}} \), cold spot volume, CI and HI were compared for assessing the quality of PTV coverage. TD and TH could provide better \( D_{98\%} \) and \( D_{2\%} \) than IMRT and VMAT \( (p<0.05) \), while achieving worse \( D_{\text{mean}} \) than IMRT and VMAT \( (p<0.05) \). Compared with IMRT, cold spot volume (%) of PTV with TD, VMAT and TH decreased by 68.45% \( (p=0.000) \), 53.57% \( (p=0.000) \) and 76.79% \( (p=0.000) \). The CI was worse with TD \( (0.830\pm0.007) \) and TH \( (0.850\pm0.008) \) than with IMRT \( (0.905\pm0.004) \) and

**Figure 1** Example of the dose distribution using IMRT, TD, VMAT and TH for the same patient.

**Abbreviations:** IMRT, intensity-modulated radiotherapy; TD, TomoDirect; VMAT, volumetric modulated arc therapy; TH, TomoHelical.
VMAT (0.920±0.004), while achieving better HI than IMRT and VMAT (p<0.05). The findings on PTV are listed in Table 2.

OARs
The radiation dose to the OARs is listed in Table 3. In comparison with TD and TH, IMRT and VMAT reduced the maximum and the mean dose of optic chiasm (p<0.05), left and right optic nerve (p<0.05) and spinal cord (p<0.05). Similarly, compared with TD, VMAT and TH, the maximum to right eye with IMRT was decreased by 5.04%, 4.35% and 6.28% (p=0.028; 0.019; 0.014). Compared with TH, the maximum dose to left lens with IMRT, TD and VMAT were significantly decreased by 11.65%, 28.19% and 9.13% (p=0.021; 0.000; 0.000), the mean dose to left lens with TD and VMAT were decreased by 26.62% and 15.82% (p=0.000; 0.001), and the maximum dose to right lens with TD and VMAT were decreased by 28.15% and 13.96% (p=0.000; 0.035). In comparison with IMRT, the mean dose to oropharynx with TD, VMAT and TH were significantly increased by 249.20%, 15.76% and 250.32% (p=0.000; 0.000; 0.000), the mean dose to thyroid with TD, VMAT and TH were significantly increased by 52.94%, 23.53% and 58.82% (p=0.000; 0.000; 0.000), the

Table 1 All the parameters utilized in EAR calculations in this study

| Site          | β_EAR | γ_e | γ_a | α  | α/β | R   |
|---------------|-------|-----|-----|----|-----|-----|
| Oropharynx    | 0.56  | -0.024 | 2.38 | 0.043 | 10  | 0.97|
| Thyroid       | 1.2   | -0.046 | 0.6 | 0.033 | 3   | 0.5 |
| Salivary gland| 0.56  | -0.024 | 2.38 | 0.087 | 3   | 0.23|

Abbreviation: EAR, excess absolute risk.
mean dose to left salivary with TD, VMAT and TH were significantly increased by 160.23%, 34.09% and 120.45% \((p=0.022; 0.000; 0.020)\), and the maximum dose to right salivary with TD, VMAT and TH were significantly increased by 122.67%, 31.33% and 117.33% \((p=0.027; 0.000; 0.032)\).

**MUs and treatment time**

The MUs and treatment time of the four treatment modalities are shown in Table 2. The mean MUs of IMRT, TD, VMAT and TH were 1191.08, 3415.33, 451.42 and 3441.42. Compared with VMAT, the mean MUs of IMRT, TD and TH were significantly increased by 163.85%, 656.57% and 662.35% \((p=0.000; 0.000; 0.000)\). The mean treatment time of IMRT, TD, VMAT and TH were 421.66, 327.10, 189.27 and 245.46 s. In comparison with VMAT, the mean treatment time of IMRT, TD and TH were significantly increased by 122.78%, 72.82% and 29.69% \((p=0.000; 0.000; 0.000)\).

**Cost of RT**

The cost of per treatment delivery for IMRT and VMAT are ~1200 RMB, and for TH and TD are ~3000 RMB, respectively. For all RT plans, the prescribed dose was 50 Gy in 25 fractions, so the total cost of treatment delivery for IMRT and VMAT are ~3000 RMB, while TD and TH are ~75,000 RMB, enhancing 150% compared to IMRT and VMAT.

**EAR calculations**

The EAR to the organs of oropharynx, thyroid, left and right salivary gland with four treatment modalities are shown in Table 4. Compared with IMRT, the EAR to the organs of oropharynx with TD, VMAT and TH were increased by 83.67%, 14.97% and 82.31% \((p=0.000; 0.000; 0.000)\), the EAR to the organ of thyroid with TD, VMAT and TH were increased by 36.36%, 18.18% and 36.36% \((p=0.002; 0.000; 0.001)\), the EAR to the organ of left salivary gland with TD, VMAT and TH were increased by 38.71%, 29.03% and 41.94% \((p=0.000; 0.000; 0.000)\) and the EAR to the organ of right salivary gland with TD, VMAT and TH were increased by 35.48%, 29.03% and 35.48% \((p=0.000; 0.000; 0.000)\).

**Discussion**

The study concerning estimation of various RT modalities for stage I-II NNKTCCL is rare, hence, this study reporting a comparison about plan quality, radiation delivery
Table 3 Results of dosimetric comparisons for OARs from DVH (x±S)

| Parameters           | IMRT | TD    | VMAT   | TH    | p-value          |
|----------------------|------|-------|--------|-------|------------------|
|                      |      | D<sub>max</sub> (Gy) |        |        | IMRT vs TD | IMRT vs VMAT | IMRT vs TH | TD vs VMAT | TD vs TH | VMAT vs TH |
| Optic chiasm         | 41.50±1.40 | 45.69±1.45 | 42.10±1.42 | 45.02±1.31 | 0.000 | 0.030 | 0.000 | 0.000 | 0.181 | 0.000 |
|                      | 30.43±1.10 | 34.25±1.13 | 31.26±1.04 | 34.24±1.02 | 0.000 | 0.027 | 0.000 | 0.000 | 0.990 | 0.000 |
| Left optic nerve     | 46.48±1.38 | 51.21±0.07 | 47.00±1.45 | 49.98±0.88 | 0.000 | 0.218 | 0.001 | 0.000 | 0.002 | 0.002 |
|                      | 35.08±1.09 | 44.65±0.09 | 38.35±1.37 | 43.78±1.07 | 0.000 | 0.000 | 0.000 | 0.000 | 0.205 | 0.000 |
| Right optic nerve    | 46.80±1.04 | 50.85±0.53 | 46.86±1.12 | 49.81±0.46 | 0.000 | 0.820 | 0.000 | 0.000 | 0.000 | 0.001 |
|                      | 35.46±1.32 | 42.86±0.92 | 37.26±1.30 | 41.91±0.96 | 0.000 | 0.000 | 0.000 | 0.000 | 0.137 | 0.000 |
| Left eye             | 47.55±1.28 | 48.57±0.83 | 48.32±1.23 | 49.07±0.97 | 0.171 | 0.161 | 0.066 | 0.703 | 0.215 | 0.238 |
|                      | 21.63±0.83 | 26.91±1.09 | 21.09±0.88 | 27.44±0.57 | 0.000 | 0.023 | 0.000 | 0.000 | 0.251 | 0.000 |
| Right eye            | 43.75±1.72 | 46.07±1.23 | 45.74±1.33 | 46.68±1.05 | 0.028 | 0.019 | 0.014 | 0.637 | 0.277 | 0.323 |
|                      | 21.47±1.41 | 25.09±0.68 | 19.42±0.93 | 25.70±0.61 | 0.012 | 0.124 | 0.008 | 0.000 | 0.056 | 0.000 |
| Left lens            | 12.66±0.33 | 10.29±0.60 | 10.95±0.29 | 14.33±0.62 | 0.001 | 0.000 | 0.021 | 0.277 | 0.000 | 0.000 |
|                      | 10.13±0.27 | 7.47±0.38 | 8.57±0.24 | 10.18±0.40 | 0.000 | 0.000 | 0.015 | 0.000 | 0.000 | 0.001 |
| Right lens           | 12.50±0.57 | 9.11±0.63 | 10.91±0.44 | 12.68±0.81 | 0.000 | 0.000 | 0.818 | 0.011 | 0.000 | 0.035 |
|                      | 9.92±0.38 | 6.66±0.41 | 8.56±0.34 | 8.87±0.53 | 0.000 | 0.000 | 0.061 | 0.001 | 0.000 | 0.639 |
| Spinal cord          | 11.79±1.93 | 17.86±2.00 | 12.10±1.88 | 16.99±1.85 | 0.000 | 0.000 | 0.599 | 0.000 | 0.000 | 0.000 |
|                      | 1.17±0.22 | 8.97±1.00 | 1.46±0.30 | 8.53±0.93 | 0.000 | 0.000 | 0.003 | 0.000 | 0.000 | 0.000 |
| Brainstem            | 35.20±1.07 | 36.10±1.29 | 33.60±1.24 | 32.40±1.63 | 0.013 | 0.013 | 0.014 | 0.001 | 0.000 | 0.145 |
|                      | 23.64±0.82 | 23.06±0.61 | 23.31±0.82 | 21.75±1.10 | 0.038 | 0.338 | 0.012 | 0.621 | 0.000 | 0.242 |
| Left parotid         | 24.01±1.98 | 24.13±1.72 | 21.94±1.73 | 23.64±1.52 | 0.090 | 0.021 | 0.753 | 0.036 | 0.538 | 0.045 |
|                      | 6.45±0.98 | 12.59±0.92 | 6.48±0.89 | 12.62±0.87 | 0.000 | 0.878 | 0.000 | 0.000 | 0.943 | 0.000 |
| Right parotid        | 21.97±2.02 | 21.13±1.80 | 19.79±1.95 | 21.25±1.79 | 0.416 | 0.020 | 0.489 | 0.125 | 0.811 | 0.031 |
|                      | 6.41±1.15 | 11.20±0.98 | 6.57±1.05 | 11.27±0.99 | 0.000 | 0.096 | 0.000 | 0.000 | 0.809 | 0.000 |
| Oropharynx           | 32.68±3.78 | 42.15±2.21 | 33.46±3.70 | 42.52±2.12 | 0.000 | 0.242 | 0.000 | 0.000 | 0.444 | 0.000 |
|                      | 6.22±1.08 | 21.72±1.16 | 7.20±1.18 | 21.79±1.09 | 0.000 | 0.000 | 0.000 | 0.000 | 0.808 | 0.000 |
| Thyroid              | 0.34±0.03 | 0.39±0.02 | 0.38±0.03 | 0.39±0.03 | 0.000 | 0.000 | 0.000 | 0.000 | 0.305 | 0.000 |
|                      | 0.17±0.02 | 0.26±0.01 | 0.21±0.01 | 0.27±0.02 | 0.000 | 0.000 | 0.000 | 0.000 | 0.305 | 0.000 |
| Left salivary        | 1.43±0.12 | 3.21±1.04 | 1.95±0.21 | 3.18±0.92 | 0.076 | 0.001 | 0.046 | 0.177 | 0.828 | 0.129 |
|                      | 0.88±0.07 | 2.29±0.61 | 1.18±0.11 | 1.94±0.47 | 0.022 | 0.000 | 0.020 | 0.064 | 0.327 | 0.076 |
Table 3 (Continued).

| Parameters          | IMRT  | TD   | VMAT  | TH    | p-value |
|---------------------|-------|------|-------|-------|---------|
|                     |       |      |       |       | IMRT vs TD | IMRT vs VMAT | IMRT vs TH | TD vs VMAT | TD vs TH | VMAT vs TH |
| Right salivary      | D<sub>max</sub> (Gy) | 1.50±0.21 | 3.34±0.98 | 1.97±0.21 | 3.26±0.97 | 0.027 | 0.000 | 0.032 | 0.112 | 0.515 | 0.131 |
|                     | D<sub>mean</sub> (Gy) | 1.14±0.24 | 2.00±0.51 | 1.41±0.25 | 1.97±0.51 | 0.089 | 0.424 | 0.099 | 0.204 | 0.609 | 0.227 |

Abbreviations: OARs, organs at risk; DVH, Dose–Volume Histogram; IMRT, intensity-modulated radiotherapy; TD, TomoDirect; VMAT, volumetric modulated arc therapy; TH, TomoHelical.

Table 4 The EAR of oropharynx, thyroid, left and right salivary gland with four treatment modalities

| EAR              | IMRT | TD   | VMAT  | TH    | p-value |
|------------------|------|------|-------|-------|---------|
|                   |      |      |       |       | IMRT vs TD | IMRT vs VMAT | IMRT vs TH | TD vs VMAT | TD vs TH | VMAT vs TH |
| Oropharynx        | 1.47±0.20 | 2.70±0.29 | 1.69±0.22 | 2.68±0.28 | 0.000 | 0.000 | 0.000 | 0.000 | 0.607 | 0.000 |
| Thyroid           | 0.11±0.02 | 0.15±0.02 | 0.13±0.02 | 0.15±0.02 | 0.002 | 0.000 | 0.001 | 0.077 | 0.989 | 0.045 |
| Left salivary gland | 0.31±0.03 | 0.43±0.05 | 0.40±0.04 | 0.44±0.05 | 0.000 | 0.000 | 0.000 | 0.050 | 0.669 | 0.052 |
| Right salivary gland | 0.31±0.03 | 0.42±0.04 | 0.40±0.04 | 0.42±0.04 | 0.000 | 0.000 | 0.000 | 0.045 | 0.832 | 0.031 |

Abbreviations: EAR, excess absolute risk; IMRT, intensity-modulated radiotherapy; TD, TomoDirect; VMAT, volumetric modulated arc therapy; TH, TomoHelical.
efficiency, cost of RT and SCR among IMRT, TD, VMAT and TH for treating stage I-II NNKTCI is significant. The purpose of the study is to estimate the properties of four types of RT modalities, and further provide guidance for selecting the optimal RT modality in clinical practice for stage I-II NNKTCI.

The reported researches confirmed that IMRT therapy has demonstrated excellent target dose distribution and OARs sparing in stage I-II NNKTCI.\(^3\)\(^6\)\(^7\) Compared with IMRT, TD, VMAT and TH were the hotter issues in recent years. VMAT therapy in stage I-II NNKTCI has demonstrated excellent target coverage quality, improvements in OARs sparing and delivery time.\(^3\)\(^7\) TH therapy in stage I-II NNKTCI also has demonstrated its advantages in target coverage quality.\(^5\)\(^7\) However, only this study utilized TD for stage I-II NNKTCI up to present. In the study, we generally found that TD, VMAT and TH offered better D\(_{95}\), cold spot volume and HI than IMRT, and TH plan had the optimal cold spot volume and HI, while CI with IMRT was slightly superior to with TD and TH. VMAT provided the optimal MUs and radiation delivery time, which has confirmed in previous studies.\(^3\)\(^7\)

However, the unexpected results showed that IMRT obtained the optimal sparing of OARs in current study, and TD, VMAT and HD had higher maximum and mean radiation dose in most OARs. But previous researches confirmed that VMAT and HT offered improved OARs sparing compared with IMRT, and especially reduced side effects of parotid glands.\(^27\)\(^28\) The opposite findings about OARs sparing could be caused by the different spatial relationship of OARs and PTV. In general, the target volume of NNKTCI is not just close to lenses, optic nerves, optic chiasm and eyes, but also surrounded by them in space. Once TD, VMAT and TH pursue deceased cold spot volume or improved HI in target volume, the radiation dose to OARs would unavoidably increase. This implies that a good treatment plan should achieve a balance between target coverage and critical organ protection.

With the rising cost of RT, the cost-effectiveness of the treatment modality must be concerned.\(^29\) In this study, we only compare the cost of treatment delivery among the four treatment modalities, and find that the total cost of TD and TH dramatically increased 150% compared with IMRT and VMAT. Andrea Peeters et al\(^30\) evaluated the cost of external beam RT with carbon-ions, protons and photons and illuminated the higher cost of particle therapy, which might reduce by fewer fractions. However, in our study, the four treatment modalities had the same prescribed dose of 50 Gy in 25 fractions, the difference of RT cost for the four treatment modalities was determined by the cost of per treatment delivery.

For decades, improving curative effects in RT results in longer survival time for cancer patients, the assessment of SCR caused by RT has become increasingly important.\(^14\) The second cancer exists highly for patients who received RT, which have demonstrated by numerous clinical studies, and usually occur in volumes receiving total dose irradiation or nearby areas radiated with the doses from 2 to 50 Gy.\(^31\)\(^32\) Thyroid is known to have a low dose threshold for radiation-induced cancer, which is as low as 0.05 Gy in children and young adults.\(^33\)\(^34\) Hence, in RT treatment, second cancer for thyroid or other organs may be affected by out-of-field dose, which is likely to be highly associated with treatment facility due to leakage and scattering from head and accessories.\(^35\) Featured with more field and MUs, intensity-modulated techniques will increase leakage and scattering radiation unavoidably.\(^36\) Recently, particle therapy is introduced into clinical routine, which can often obtain excellent target converge while delivering minimal dose to normal tissue nearby tumor targets, and significantly decrease the risk of second cancer.\(^37\) However, due to the high cost of the particle therapy confirmed by Andrea Peeters et al,\(^30\) the total benefit still needs to be discussed.

For predicting cancer induction, based on the DVH data of the RT plan and some parameters, model EAR and model excess relative risk or a mixture of the two models were developed to calculate SCR.\(^11\)\(^38\)\(^39\) Previous literature\(^26\)\(^40\) compared SCR of RT modalities in whole-breast irradiation and rectal cancer and showed that RT modalities had obvious difference in SCR to some organs, but similar study was rare in head and neck cancer. In this study, owing to the insufficient data of survival rate for NNKTCI, we used EAR to determine SCR. Our study showed that IMRT obtained the lowest EAR to the organs of oropharynx, thyroid, the left and right salivary gland, hence, IMRT was suitable for the young patients who probably have a longer life expectancy. However, uncertainties inevitably existed in the accuracy of the commercially TPS models and the risk models, the results about EAR evaluating carried with inherent limitation.

In general, each RT modality had its own characteristics, adapting to individualized patient treatment. The optimal treatment modality for every individual patient depended on the balance between all relevant factors, such as target converge, normal tissue damage, economic aspects and long-term risk effects.
Conclusion

Both TD and TH showed slight improvements in target quality compared with IMRT for stage I-II NNKTCL. However, IMRT could offer lower cost of RT, lowest EAR in some organs and lowest dose to most OARs, and VMAT featured with lower cost of RT, lowest MUs and shortest delivery time. Each treatment modality has its own advantage, so selecting appropriate RT modality for individualized patient is necessary.

Abbreviation list

RT, radiotherapy; NNKTCL, nasal natural killer T-Cell lymphoma; EAR, excess absolute risk; FF-IMRT, fixed-field intensity-modulated radiotherapy; VMAT, volumetric-modulated arc therapy; TD, TomoDirect; TH, TomoHelical; HI, homogeneity index; PTV, planning target volume; CI, conformity index; MUs, monitor units; OARs, organs at risk; SCR, second cancer risk; TPS, treatment planning system; CT, computer tomography; GTV, gross tumor volume; CTV, clinical target volume; DR, dose rate; DVH, Dose–Volume Histogram.

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Disclosure

The authors report no conflicts of interest in this work.

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