Prognostic factors and failure patterns in non-metastatic nasopharyngeal carcinoma after intensity-modulated radiotherapy

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Abstract

Background: The prognostic values of staging parameters require continual re-assessment amid changes in diagnostic and therapeutic methods. This study aimed to identify the prognostic factors and failure patterns of non-metastatic nasopharyngeal carcinoma (NPC) in the intensity-modulated radiotherapy (IMRT) era.

Methods: We reviewed the data from 749 patients with newly diagnosed, biopsy-proven, non-metastatic NPC in our cancer center (South China, an NPC endemic area) between January 2003 and December 2007. All patients underwent magnetic resonance imaging (MRI) before receiving IMRT. The actuarial survival rates were estimated using the Kaplan–Meier method, and survival curves were compared using the log-rank test. Multivariate analyses with the Cox proportional hazards model were used to test for the independent prognostic factors by backward eliminating insignificant explanatory variables.

Results: The 5-year occurrence rates of local failure, regional failure, locoregional failure, and distant failure were 5.4, 3.0, 7.4, and 17.4%, respectively. The 5-year survival rates were as follows: local relapse-free survival, 94.6%; nodal relapse-free survival, 97.0%; distant metastasis-free survival, 82.6%; disease-free survival, 75.1%; and overall survival, 82.0%. Multivariate Cox regression analysis revealed that orbit involvement was the only significant prognostic factor for local failure ($P = 0.011$). Parapharyngeal tumor extension, retropharyngeal lymph node involvement, and the laterality, longest diameter, and Ho's location of the cervical lymph nodes were significant prognostic factors for both distant failure and disease failure (all $P < 0.05$). Intracranial extension had significant prognostic value for distant failure ($P = 0.040$).

Conclusions: The key failure pattern for NPC was distant metastasis in the IMRT era. With changes in diagnostic and therapeutic technologies as well as treatment modalities, the significant prognostic parameters for local control have also been altered substantially.

Keywords: Nasopharyngeal carcinoma, Intensity-modulated radiotherapy, Prognosis, Failure pattern, Tumor staging

Background

Nasopharyngeal carcinoma (NPC) is a challenge in oncology. Due to the deep-seated anatomic location and proximity of NPCs to critical structures, radical surgical resection is extremely difficult [1]. The introduction of radiotherapy has made this otherwise lethal malignant disease curable [1]. However, radiotherapy has undergone several different periods of development, and it was not until the advent of mega-voltage machines that a 5-year overall survival (OS) rate of 25% was first achieved, which marked the first major breakthrough in the treatment of NPC and established radiotherapy as the primary modality of choice for NPC [2].
Since that time, progressive improvements in the treatment outcomes of patients with NPC have been achieved. Initially, in the 1970–1980s, conventional radiotherapy alone resulted in similar outcomes in both the endemic and non-endemic areas, with 5-year OS rates of 48%–52%, a cumulative local failure rate of 20%, a cumulative regional failure rate of 14%, and a cumulative distant metastasis rate of 19% [3, 4]. During the 1990s, rapid technological advances in imaging methods, computerized planning systems, and radiotherapy facilities and the accumulation of radiobiological knowledge that enabled schedule, dose, and fractionation optimization led to better outcomes after radiotherapy, with 5-year OS rates of 65%–74%, a cumulative local failure rate of 12%, a regional failure rate of 5%, and a distant metastasis rate of 16% [5, 6]. However, local relapse and distant metastasis remained the two major causes of failure. Intensity-modulated radiotherapy (IMRT) was introduced in the 2000s and represented a major breakthrough in the radiotherapeutic management of NPC. Due to the dosimetric advantages of IMRT combined with the use of magnetic resonance imaging (MRI)-guided tumor volume delineation and concurrent chemotherapy, local control rate in NPC patients has improved significantly [7, 8]. With these changes in diagnostic and therapeutic technologies, the significant prognostic parameters that have been identified in patients treated with conventional radiotherapy may also have been altered substantially. Therefore, caution must be applied when applying prognostic factors to predict prognosis and guide treatment strategies in patients treated using these modern, more effective treatment modalities [9]. Thus, the prognostic values of staging parameters require continual re-assessment amid these changes in diagnostic and therapeutic methods. However, few studies have systematically investigated the prognostic factors in NPC in the IMRT era. We performed a retrospective systematic evaluation of a large cohort of patients with NPC who were treated with IMRT at the Sun Yat-sen University Cancer Center in South China, an NPC endemic area. The aim of this study was to identify the key failure patterns and prognostic factors to improve the treatment of patients with NPC and provide a baseline for future studies.

**Methods**

**Patient cohort**

We reviewed the medical records of all patients with NPC treated with IMRT at the Sun Yat-sen University Cancer Center between January 2003 and December 2007. We included all patients with newly diagnosed, non-metastatic, histologically confirmed disease. All patients were staged using a standard protocol [10] comprising a complete disease history, physical examination, hematology and biochemistry profiles, fiberoptic nasopharyngoscopy, MRI of the neck and nasopharynx, chest radiography, abdominal sonography, and a whole-body bone scan using single-photon emission computed tomography (SPECT). All MRI materials and clinical records were reviewed to minimize heterogeneity in restaging. The study was reviewed and approved by the ethics committee of the Sun Yat-sen University Cancer Center.

**Image assessment and criteria for staging parameters**

Two radiologists specializing in head and neck cancers evaluated all scans separately. All parameters included in the TNM staging were re-assessed, and any disagreements were resolved by consensus. Parapharyngeal tumor extension was identified as a direct tumor extension through the pharyngobasilar fascia [11]. Nasal cavity extension was defined as tumor extension beyond the bony nasal septum [12]. Oropharyngeal extension was defined as the primary tumor extension of the mucosa or submucosal tissue below the plane of the superior surface of the soft palate or the C1–C2 interspace [13]. Hypopharyngeal extension was defined as the primary tumor extension of the mucosa or submucosal tissue to the superior border of the epiglottis or lower margin of the C3 interspace [14]. The skull base included the clivus, pterygoid bones, body of the sphenoid, apices of the petrous temporal bones, sphenoid wings, upper cervical spine, and skull base foramina and fissures [15]. The criterion for orbit extension was tumor extension into the orbital apex, inferior orbital fissure, or superior orbital fissure [16]. Involvement of the cavernous sinus, brain tissue, cistern, or dural meninges was interpreted as intracranial extension [17].

The diagnostic criteria for metastatic lymphadenopathy [18] include the following: (a) lateral retropharyngeal lymph nodes with a minimal axial diameter in the largest plane of an individual node of at least 5 mm and any node observed in the median retropharyngeal group, lymph nodes with a minimal axial diameter of at least 11 mm in the jugulodigastric region and 10 mm for all other cervical nodes; (b) lymph nodes of any size with central necrosis or a contrast-enhanced rim; (c) nodal groups, defined as the presence of three or more contiguous and confluent lymph nodes, each of which with a minimal axial diameter of 8–10 mm; and/or (d) lymph nodes of any size with extracapsular spread, the presence of indistinct nodal margins, irregular nodal capsular enhancement, or infiltration into the adjacent fat or muscle.

**Treatment**

All patients underwent radical radiotherapy. The nasopharyngeal and upper neck tumor volumes were treated using IMRT for the entire treatment course. The lower
neck was treated with a conventional anterior or anter- 
oposterior opposing cervical technique. The prescribed 
doses were 68 Gy to the planning target volume of the 
primary tumor (PTV-P), 60–68 Gy to the PTV of the 
nodes (PTV-N), 60 Gy to the high-risk regions (PTV1), 
and 54 Gy to the low-risk regions (PTV2) in 30 frac- 
tions over 6 weeks. The anterior split-neck field received 
50–60 Gy irradiation in 2-Gy fractions. The irradiation 
was delivered once daily for 5 days per week. Further 
details of the IMRT techniques used at our cancer center 
have been previously reported [10].

During the study period, concurrent chemoradiother- 
apy was recommended for stage III to IVa-b disease, and 
no chemotherapy was recommended for stage I to IIa 
disease (as defined using the 6th edition of the Interna- 
tional Union against Cancer/American Joint Committee 
on Cancer [UICC/AJCC] staging system). However, some 
patients with stage III to IVa–b disease also received neo-
adjuvant or adjuvant chemotherapy, and some patients 
with stage IIb disease received concurrent chemora-
diotherapy. Neoadjuvant or adjuvant chemotherapy 
consisted of cisplatin (80–100 mg/m²) with either 5-fluo-
rouracil (800–1000 mg/m²) or taxanes (paclitaxel [135– 
175 mg/m²] or docetaxel [70–75 mg/m²]) every 3 weeks 
for 3 cycles. Concomitant chemotherapy consisted of 
cisplatin (80–100 mg/m²) given on weeks 1, 4, and 7 of 
radiotherapy or cisplatin (30–40 mg/m²) given weekly. 
When possible, salvage treatments, such as intracavitary 
brachytherapy, surgery, and chemotherapy, were pro-
vided in the event of documented relapse or persistent 
disease.

Follow-up
Treatment responses and toxicity were assessed in each 
patient every week during treatment, every 2–3 months 
during the first 2 years after treatment, and then every 
3–6 months during the following 3 years. Endoscopy, 
computed tomography (CT), or MRI scans of the head 
and neck were also performed every 3 months in the first 
year and annually thereafter. Additional tests were per-
formed when indicated to evaluate local or distant fail-
ure. The last follow-up time was 30th August 2013.

Statistical analysis
All analyses were performed using SPSS version 16.0 
(SPSS, Chicago, IL, USA). The actuarial rates were esti-
mated using the Kaplan–Meier method, and the survival 
curves were compared using the log-rank test. The fol-
lowing endpoints (measured from the start of treatment 
to the first defined event) were estimated: local relapse-
free survival (LRFS), nodal relapse-free survival (NRFS), 
distant metastasis-free survival (DMFS), disease-free 
(DFS), and OS.

Multivariate analyses with the Cox proportional haz-
ARDS model were used to test for independent prognostic 
fhazards model was also used to calculate the hazard ratios (HRs) 
and 95% confidence intervals (CIs). Host factors (age and 
sex) were included as covariates in all tests. Two-tailed $P$
values <0.05 were considered statistically significant.

Results
Patient characteristics
The median patient age was 43 years (range, 
13–78 years), and the cohort included 580 males and 
169 females (male:female ratio of 3.4:1). Histologically, 
744 (99.3%) patients had World Health Organization 
(WHO) type II/III disease, and 5 (0.7%) had WHO 
type I disease. Positron emission tomography (PET)/ 
CT was performed for 162 (21.6%) patients. Using the 
2009 7th UICC/AJCC staging system, the numbers of 
patients with stage I, II, III, IVa, and IVb disease were 
78 (10.4%), 179 (23.9%), 282 (37.7%), 160 (21.3%), and 
50 (6.7%), respectively. The median follow-up duration 
was 81.4 months (range, 3.1–126.5 months). In total, 
424 (86.2%) of the 492 patients with stage III–IV dis-
ease received chemotherapy. The characteristics of the 
patients are summarized in Table 1.

Dose-volume histogram analysis
Table 2 summarizes the dose-volume histogram (DVH) 
statistics for the target volumes of interest. The maxi-
mum dose ($D_{\text{max}}$) and minimum dose ($D_{\text{min}}$) of radia-
tion were defined as the maximum and minimum dose 
points in the DVH. The radiation dose prescribed to the 
PTV-P was 68 Gy for all patients. Because the proportion 
of the volume that received $<93\%$ of the prescribed dose 
was set to $\leq3\%$, the actual median dose to the PTV-P was 
73.34 Gy.

The dose limitations are $\leq60$ Gy to the brainstem 
and $\leq50$ Gy to the spinal cord. In some advanced 
diseases with extensive extension, we would relax the dose 
limitation to $D_{\text{max}} \leq 63$ Gy. The median $D_{\text{max}}$ for the 
brainstem was 54.16 Gy (range, 31.68–77.65 Gy), and the 
median proportion of the volume in the brainstem that 
received $>60$ Gy was 9.67% (range, 0.03%–26.97%). The median $D_{\text{max}}$ for the spinal cord was 33.96 Gy (range, 
8.29–60.54 Gy), and the median proportion of the vol-
ume in the spinal cord that received $>50$ Gy was 0.97% 
(range, 0.01%–3.42%). However, 3 of the 749 patients 
received irradiation with a $D_{\text{max}} > 63$ Gy, with no serious 
late toxicity for the brainstem. The median doses to the 
left and right parotid glands were 28.20 Gy (range, 11.90– 
49.70 Gy) and 29.98 Gy (range, 21.30–42.02 Gy). The median proportions of the volumes in the left and right
Patterns of treatment failure and survival

Among the 749 patients, 56 (7.5%) relapsed [34 (4.5%) had local relapse only, 15 (2.0%) had nodal relapse only, 7 (1.0%) had both local and nodal relapse], and 129 (17.2%) had distant metastases. One hundred forty-nine (19.9%) patients died. The median time to relapse was 25.2 (range, 7.0–93.8) months, and the median time to distant metastases was 18.9 (range, 2.6–92.1) months.

### Table 1 Clinicopathologic features of the 749 patients with nasopharyngeal carcinoma (NPC)

| Feature               | No. of patients [cases (%)] |
|-----------------------|-----------------------------|
| Sex                   |                             |
| Male                  | 580 (77.4)                  |
| Female                | 169 (22.6)                  |
| WHO pathology         |                             |
| Type I                | 5 (0.7)                     |
| Type II/III           | 744 (99.3)                  |
| Clinical stage\(a\)  |                             |
| I–II                  | 257 (34.3)                  |
| III–IV                | 492 (65.7)                  |
| T stage\(a\)          |                             |
| T1                    | 177 (23.6)                  |
| T2                    | 140 (18.7)                  |
| T3                    | 264 (35.2)                  |
| T4                    | 168 (22.4)                  |
| N stage\(a\)          |                             |
| N0                    | 184 (24.6)                  |
| N1                    | 409 (54.6)                  |
| N2                    | 106 (14.2)                  |
| N3                    | 50 (6.7)                    |
| Chemotherapy          |                             |
| No chemotherapy       | 214 (28.6)                  |
| Concurrent            | 243 (32.5)                  |
| Induction + concurrent| 246 (32.8)                  |
| Concurrent + adjuvant | 46 (6.1)                    |

WHO World Health Organization

\(a\) According to the 7th edition of the Union for International Cancer Control/American Joint Committee on Cancer (UICC/AJCC) staging system

The 5-year local failure, regional failure, locoregional failure, and distant failure rates for the whole cohort were 5.4, 3.0, 7.4, and 17.4%, respectively (Fig. 1). The 5-year survival rates for the whole cohort were as follows: LRFS, 94.6%; NRFS, 97.0%; DMFS, 82.6%; DFS, 75.1%; and OS, 82.0%. Using the 7th edition of the UICC/AJCC staging system, the 5-year LRFS rates for patients with stage T1–T4 NPC were 98.4, 96.2, 93.0, and 90.5%, respectively (Fig. 2a); the 5-year DMFS rates for patients with stage N0–N3b NPC were 93.3, 84.2, 72.4, 61.9, and 50.7%, respectively (Fig. 2b); and the 5-year OS rates for patients with stage I–IVb NPC were 97.4, 93.8, 81.8, 69.7, and 54.0%, respectively (Fig. 2c).

### Table 2 Dose-volume histogram data for planning target volumes in the 749 patients with NPC

| Variate | Volume (mL) | \(D_{\text{max}}\) (Gy) | \(D_{\text{min}}\) (Gy) | Mean dose (Gy) |
|---------|-------------|--------------------------|--------------------------|-----------------|
| PTV-P   | 51.89 (260–287.48) | 79.91 (70.47–93.95) | 59.79 (50.58–71.16) | 73.11 (68.16–78.48) |
| PTV1    | 97.35 (73.88–494.77) | 79.67 (72.81–86.08) | 58.86 (49.13–67.32) | 67.94 (56.66–74.48) |
| PTV2    | 368.97 (106.36–685.58) | 78.99 (70.04–86.74) | 45.28 (43.12–61.47) | 60.55 (53.64–72.13) |

PTV-P planning target volume of the primary tumor, PTV1 PTV of the high-risk clinical target volume, PTV2 PTV of the low-risk clinical target volume, \(D_{\text{max}}\), maximum dose, \(D_{\text{min}}\), minimum dose. All data are presented as median with range in parentheses

### Prognostic factors

Table 3 lists the factors with independent significance for different endpoints. The TNM stage was consistently the most significant factor for prognosis prediction: T stage for local failure prediction \((P < 0.001)\), and both T and N stages for distant failure, disease failure, and death prediction (all \(P < 0.001)\). None of the factors had a significant influence on nodal failure. An age >50 years was associated with significantly higher rates of distant failure \((HR = 1.530, 95\% \text{ CI} 1.037–2.256, P = 0.032)\), disease failure \((HR = 1.896, 95\% \text{ CI} 1.403–2.561, P < 0.001)\), and death \((HR = 2.482, 95\% \text{ CI} 1.770–3.482, P < 0.001)\).

The involvement of the nasal cavity, oropharynx, parapharynx, skull base, paranasal sinus, cranial nerve palsy, intracranial extension, infratemporal fossa, intracranial extension, orbit, and hypopharynx (all were T stage-related parameters) occurred in 280 (37.4%), 85 (11.3%), 540 (72.1%), 429 (57.3%), 36 (4.8%), 35 (4.7%), and 3 (0.4%) patients, respectively. To explore the prognostic values of the specific anatomic factors included in the TNM staging system, we analyzed all T stage-related parameters, all N stage-related parameters (i.e., retropharyngeal lymph node involvement as well as the laterality, longest diameter, and Ho’s location of the involved cervical lymph nodes), age, histological type, and chemotherapy modality using univariate and multivariate analyses. With the exception of hypopharyngeal involvement, the univariate analysis revealed that age, histological type, and all T stage- and N stage-related parameters were significantly associated with disease.
failure (all $P < 0.05$). All factors that were significant in univariate analysis were included in the Cox proportion hazards model with the backward elimination of non-significant explanatory variables. Only orbit involvement was found to be significant prognostic factor for local failure in the multivariate analysis ($P = 0.011$). Parapharyngeal involvement, retropharyngeal lymph node involvement, and the laterality, longest diameter, and Ho’s location of the involved cervical lymph nodes were significant prognostic factors for both distant failure and disease failure (all $P < 0.05$). Intracranial extension had significant prognostic value for distant failure ($P = 0.040$) (Table 4).

**Discussion**

The present study, with 5-year LRFS and DMFS rates of 94.6% and 82.6%, confirmed that distant metastasis has surpassed local relapse to be the major cause of failure in NPC in the IMRT era, whereas our previous data from the conventional radiotherapy era demonstrated that both local relapse and distant metastasis were the major causes of failure in NPC, with 5-year LRFS and DMFS rates of 85% and 81% [6]. Although T stage was still a significant predictor for local failure, with the more extensive primary extension associated with poorer local control, the 5-year LRFS rates for patients with NPC at adjacent T stages were not significantly different. T stage is becoming less powerful in segregating patients into various risk groups. Improvements in local control have reduced the significance of known prognostic factors, even the T stage, for local control prediction [9]. It has been recommended that three rather than four T stages should be applied to classify patients in the modern era [19, 20]. In the future, treatment modality may be the only significant predictor for local control in NPC patients; and the most important challenge in the management of NPC should be the exploration of more effective systemic agents to control distant metastasis.
Based on CT and conventional radiotherapy, several studies have systematically investigated prognostic factors for local control in patients with NPC [21–23]. Two studies have demonstrated that skull base extension or cranial nerve palsy could affect local failure-free survival [21, 22], whereas one has confirmed that parapharyngeal tumor extension was an independent prognostic factor for local failure-free survival [23]. Based on MRI and IMRT, no systematic data referring to prognostic factors in patients with NPC were available. Zong et al. [24] focused on skull base extension and found that it did not influence local failure in the IMRT era. Based on the same database presented here, our previous studies reported that skull base extension [25] and parapharyngeal tumor extension [26] were not independent prognostic factors for local control in patients with NPC treated with IMRT. In the current study, we systematically re-evaluated all of the prognostic factors included in the TNM staging system and found that their prognostic values no longer remained significant and that orbit involvement was the only independent prognostic factor for local control.

Improvements in local control can explain these results. First, due to its improved dose conformability and escalation, IMRT eliminates the areas at the edges of the shielding blocks and field borders, such as the sphenoid sinus behind the pituitary shield, parapharyngeal region, and posterior skull base, which would be under-dosed when two-dimensional conventional radiotherapy is applied [27]. Thus, IMRT leads to improved local control in these areas. Second, MRI improves visualization and is superior to CT for identifying parapharyngeal tumor extension, skull base extension, and intracranial involvement [13]. Therefore, a more precise dose of radiation can be delivered to the gross target volume of primary tumor, which ultimately improves local control for these areas. However, orbit extension, which usually occurs as tumor extension from the pterygopalatine fossa via the inferior orbital fissure or directly from the cavernous sinus, is a marker of the most severe extension of NPC [15]. Even with the most sophisticated treatment techniques and intensive use of chemotherapy, some patients with advanced T4 disease that approaches important organs at risk cannot tolerate a radiation dose of 70 Gy, and the local control rate for this subgroup remains low, even after IMRT [28]. Some patients who had tiny intracranial extensions (for example, through the oval foramen or foramen lacerum to cavernous sinus) or cranial nerve palsy and did not have disease approaching important organs at risk were still able to tolerate a radiation dose of 70 Gy and achieve excellent local control. This finding might explain why orbit involvement alone and not intracranial extension or cranial nerve palsy was the only independent prognostic factor for local control.

In patients diagnosed using CT and treated with conventional radiotherapy, parapharyngeal tumor extension, skull base extension, cranial nerve palsy, retropharyngeal lymph node involvement, and all cervical lymph node-related parameters (i.e., laterality, size, Ho’s location, and...
Table 3 Multivariate analyses of significant factors for various endpoints in the 749 patients with NPC

| Variable                                      | Local failure                  | Regional failure               | Distant failure                | Disease failure                | Death (any cause)                |
|-----------------------------------------------|--------------------------------|--------------------------------|--------------------------------|--------------------------------|----------------------------------|
|                                               | HR (95% CI)                    | P value                        | HR (95% CI)                    | P value                        | HR (95% CI)                    | P value                        |
| T stage (T2, T3, and T4 vs. T1)               | 1.956 (1.371–2.790)            | <0.001                         | 1.243 (0.809–1.912)            | 0.321                          | 1.556 (1.282–1.888)            | <0.001                         | 1.589 (1.363–1.853)            | <0.001                         | 1.583 (1.328–1.886)            | <0.001                         |
| N stage (N1, N2, and N3 vs. N0)               | 1.233 (0.833–1.723)            | 0.218                          | 1.416 (0.928–2.160)            | 0.106                          | 1.741 (1.485–2.041)            | <0.001                         | 1.617 (1.415–1.847)            | <0.001                         | 1.690 (1.455–1.964)            | <0.001                         |
| Age (>50 vs. ≤50 years)                       | 1.011 (0.490–2.086)            | 0.976                          | 1.005 (0.365–2.769)            | 0.992                          | 1.530 (1.037–2.256)            | 0.032                          | 1.896 (1.403–2.561)            | <0.001                         | 2.482 (1.770–3.482)            | <0.001                         |
| Sex (female vs. male)                         | 0.826 (0.378–1.802)            | 0.631                          | 0.797 (0.266–2.386)            | 0.685                          | 1.066 (0.689–1.649)            | 0.776                          | 1.017 (0.713–1.450)            | 0.926                          | 0.900 (0.587–1.379)            | 0.627                          |
| Histology (type II/III vs. type I)            | 1.930E4 (0.000–6.656E258)      | 0.974                          | 2.104E4 (0.000–)               | 0.981                          | 2.313E4 (0.000–4.202E155)      | 0.955                          | 1.397 (0.194–10.043)           | 0.739                          | 1.037 (0.144–7.480)            | 0.971                          |
| Chemotherapy (CCRT, induction with CCRT, and  | 0.921 (0.705–1.202)            | 0.544                          | 0.920 (0.629–1.345)            | 0.666                          | 1.002 (0.870–1.155)            | 0.973                          | 0.947 (0.840–1.068)            | 0.375                          | 0.969 (0.847–1.108)            | 0.644                          |
| CCRT with adjuvant vs. none)                  |                                |                                |                                |                                |                                |                                |                                |                                |                                |                                |

HR hazard ratio, CI confidence interval, CCRT concurrent chemoradiotherapy
The prognosis of a patient with cancer depends on the biological aggressiveness of the tumor, host factors, and therapeutic interventions [9]. However, we explored the prognosis only according to anatomic factors in this study. An increasing number of biological, genetic, and molecular factors, such as plasma/serum Epstein–Barr virus (EBV) DNA [34] and specific microRNAs (miRNAs) [35], as well as the target volume [36, 37] and other non-anatomic factors, have been identified and studied [38]. Such factors may have profound influences on the prognosis for individual patients. However, the extent of local extension, regional lymphatic spread, and distant metastasis as reflected by TNM staging still remain the most important prognostic factors in NPC [9].
Because the definition of parapharyngeal tumor extension has differed substantially across different time points [26, 39], it is difficult to compare the prognostic value of parapharyngeal tumor extension. This was a retrospective study that aimed to evaluate the prognostic value of staging parameters amid the changes in diagnostic and therapeutic methods. Moreover, due to financial factors and some patients’ lack of health insurance, PET/CT was only performed on 130 (25.4%) of the 749 patients in our study. Additional effects of sensitive and specific imaging technologies such as PET on stage migration are expected. With its well-recognized superiority in the detection of nodal metastases and distant metastases [40, 41], there seems to be little doubt that the addition of PET (with or without CT) could contribute to improving staging accuracy and prognostic prediction in NPC.

Conclusions
The key failure pattern for NPC in the IMRT era was distant metastasis. With the changes in diagnostic and therapeutic technologies, the significant prognostic parameters for local control have also altered substantially. Orbit involvement was the only independent prognostic factor for local control, and intracranial extension, parapharyngeal tumor extension, and all N stage-related parameters were independent prognostic factors for DMFS.

Authors’ contributions
Conception and design of the study: YPM, LLT and JM. Acquisition of data: YM and LZL. Analysis and interpretation of the data: YPM, LLT, JM and AHL. Contributed reagents/materials/analysis tools: All authors. Writing and revision of the manuscript: All authors. All authors read and approved the final manuscript.

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Competing interests
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