Outcomes of Adding Induction Chemotherapy to Concurrent Chemotherapy for Nasopharyngeal Carcinoma Patients with Moderate-Risk in the Intensity-Modulated Radiotherapy Era

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Background: The aim of this study was to evaluate the efficacy of induction chemotherapy (IC) in nasopharyngeal carcinoma (NPC) patients with moderate-risk treated with intensity-modulated radiotherapy (IMRT).

Methods: We retrospectively assessed 506 patients with T1-2N1M0 or T3-4N0-1M0 NPC (according to the 2010 UICC/AJCC staging system) who received concurrent chemoradiotherapy (CCRT) with or without IC at a single center in China between 2005 and 2010. Survival outcomes were compared between the IC + CCRT and CCRT groups using the Kaplan–Meier method, Log-rank test and a Cox regression model.

Results: Among the 506 patients, CCRT alone resulted in equivalent overall survival (86.8% vs 89.8%, p=0.756), progression-free survival (79.6% vs 79.6%, p=0.364) and distant metastasis-free survival (88.0% vs 89.8%, p=0.407) to IC plus CCRT. In multivariate analysis, IC did not lower the risk of death (HR 0.76, 95% CI 0.46–1.25, p=0.278), progression (HR 0.78, 95% CI 0.51–1.19, p=0.244), locoregional relapse-free survival (90.2% vs 87.0%, p=0.364) and distant metastasis-free survival (88.0% vs 89.8%, p=0.407) to IC plus CCRT. In stratified analysis based on N category (N0 vs N1) and EBV DNA (> vs ≤4000 copies/mL), the use of IC should be carefully considered in these patients, though additional prospective trials are warranted to confirm the conclusions of this study.

Conclusion: Addition of IC to CCRT does not improve survival outcomes in moderate-risk NPC; the use of IC should be carefully considered in these patients, though additional prospective trials are warranted to confirm the conclusions of this study.

Keywords: nasopharyngeal carcinoma, induction chemotherapy, prognosis, survival

Introduction
Nasopharyngeal carcinoma (NPC) is endemic throughout South China, Southeastern Asia, and North Africa, with an annual incidence of approximately 15–30 per 100,000 in Southern China. The primary treatment modality for NPC is radiotherapy. Concurrent chemoradiotherapy (CCRT) is recommended as the main treatment for locoregionally advanced NPC in the National Comprehensive Cancer Network (NCCN) guidelines.

Distant metastasis is now the predominant failure pattern. In an attempt to reduce the rate of distant metastasis and further improve OS, several groups have explored different strategies in the IMRT era, such as adding induction chemotherapy (IC) or adjuvant chemotherapy (AC) to CCRT. The role of induction chemotherapy has been
explored, and the results of clinical trials of IC are also controversial. Three randomized Phase III trial from a NPC-endemic area reported the encouraging result that addition of IC to CCRT significantly improved distant metastasis-free survival (DMFS) and OS in locoregionally advanced NPC.5–7 However, no benefit of survivals from induction chemotherapy adding to CCRT was observed in the other clinical trials.8,9 In consideration of these inconsistent clinical trial results, controversy exists regarding the value of IC in NPC. TPF is a category I option, but only for EBV-associated disease.9 Results from three systematic reviews had inconsistent results when evaluating the impact on survival outcomes, with two reviews showing that induction chemotherapy prior to systemic therapy/RT had superior OS and PFS rates, compared to systemic therapy/RT alone.10,11 while another review showed that induction chemotherapy prior to systemic therapy/RT did not have better survival outcomes than systemic therapy/RT alone or systemic therapy/RT followed by adjuvant chemotherapy.12 Expert groups (eg, ESMO, NCI) differ in their clinical practice guidelines regarding the use of induction chemotherapy for these patients,13 and the NCCN expert panel could not reach uniform consensus in this regard. Recent evidence has indicated that more than 75% of distant failures are concentrated in the advanced N-stage (stage N2–N3).14 Furthermore, high incidences of grade 3/4 adverse events were observed during the IC+CCRT groups.5–7 So in the real world, the role of IC must be reconsidered, especially those patients with stage T1-2N1M0 or T3-4N0-1M0, which were excluded in the clinical trial. In clinical practice, these patients are considered to have a moderate risk of distant metastasis compared to stage T1-2N0M0 (low risk of metastasis) and stage Tany N2-3M0 (high risk of metastasis). Therefore, we performed this study to compare the clinical outcomes of patients with stage T1-2N1M0 and T3-4N0-1M0 NPC treated with IC followed by CCRT or CCRT alone.

Materials and Methods

Patients

Inclusion criteria were: (1) NPC confirmed histologically via nasopharyngeal biopsy; (2) no distant metastasis; (3) no prior treatment; (4) no previous malignancy or other concomitant malignant disease; (5) available pretreatment EBV DNA data, (6) restaged as T1-2N1M0 or T3-4N0-1M0 according to the 8th the American Joint Committee on Cancer (AJCC) staging system for NPC5 and (7) treatment with radical IMRT; 506 patients treated with IC plus CCRT or CCRT alone between 2005 and 2010 were included. Patients were excluded if their NPC was in stage T1-2N0M0 or Tany N2-3M0 or Tany Nany M1; if they had a second primary tumor or history of malignant tumors; if they were receiving adjuvant chemotherapy or radiotherapy alone; or if pretreatment EBV DNA data were unavailable. This retrospective study was approved by the Clinical Research Ethics Committee of Sun Yat-Sen University Cancer Canter. Patient records were anonymized prior to analysis. Informed consent was not required for this research due to the anonymized patient data.

Pretreatment evaluations included a detailed medical history, physical examination and performance status assessment. Routine workup included magnetic resonance imaging (MRI) of the nasopharynx and neck, chest radiograph, bone scintigraphy and abdominal ultrasonography for all patients. 20.1% (102/506) patients also underwent positron emission tomography/computed tomography (PET/CT) examination. The detection of EBV DNA copy was as previously described.15 64.0% (324/506) patients had a positive EBV DNA copy.

Treatment

All patients received definitive IMRT with concurrent chemotherapy; 40.1% (203/506) received IC before concurrent chemoradiotherapy. The cumulative radiation doses were 68 Gy or greater to the primary tumor, 60–66 Gy to involved cervical lymph nodes and 50 Gy or greater to potential sites of local infiltration and bilateral cervical lymphatics, in 30–33 fractions. The planning technique, target volume delineation and evaluation of IMRT planning were as previously described.4,16

The IC regimens included PF (80 mg/m2 cisplatin on day 1 and 800 mg/m2/d fluorouracil civ on days 1–5), TP (75 mg/m2 docetaxel on day 1 and 75 mg/m2 cisplatin on day 1), or TPF (60 mg/m2 docetaxel on day 1, 60 mg/m2 cisplatin on day 1, 600 mg/m2/d fluorouracil civ on days 1–5) for 2–3 courses. Concurrent chemotherapy was mainly 80–100 mg/m2 of a cisplatin- or nedaplatin-based regimen given every three weeks for two to three cycles, or 30–40 mg/m2 of the cisplatin- or nedaplatin-based regimen or 20–30 mg/m2 of a docetaxel-based regimen given weekly for up to seven cycles. Patients who received only one cycle of IC or concurrent chemotherapy were excluded. In the IC group, 46.8% (95/203) of patients received TPF, 30.5% (62/203) received TP and 22.7% (46/203) received PF.
Follow-Up
All patients were followed up at regular intervals after completing treatment: every 3 months in the first 3 years, every 6 months for the next 2 years, then annually.

Endpoints
Study endpoints were OS, PFS, LRFS and DMFS. OS was defined the time from initiation of treatment to date of death from any cause or last follow-up; PFS to events that included death or disease progression at local, regional or distant sites or last follow-up; LRFS to primary site or neck lymph node relapse or last follow-up; DMFS to first observation of distant metastases or last follow-up.

Statistical Analysis
All statistical analyses were performed using SPSS version 22.0. Categorical variables were compared using the Chi-square test (or Fisher’s exact test, if the expected number was less than five in at least 25% of cells). Survival rates were calculated using the Kaplan–Meier method and the Log-rank test was used to perform paired comparisons between treatment groups using the pair-wise over strata method. Receiver operation characteristic (ROC) curve was applied to calculate the cut-off value of pretreatment EBV DNA for DMFS. Multivariate analyses with the Cox proportional hazards model were used to calculate hazard ratios (HR), 95% confidence intervals (CI) and test the independence of different factors. Two-tailed p-values <0.05 were considered statistically significant; p-value corrections for multiple comparisons were not performed.

Results
Baseline Characteristics and Failure Patterns
Table 1 summarizes the baseline clinical characteristics of the patients: 203 (40.1%) patients received IC followed by CCRT and 303 (59.9%) received CCRT alone. Patients with advanced clinical stage, advanced T category, or high EBV DNA copy number (≥4000 copies/mL) were more likely to receive IC followed by CCRT (Table 1).

The median follow-up duration for the entire cohort was 69.3 months (range, 3.5–130.1 months). By last follow-up, 26 (12.8%) patients in the IC + CCRT group and 29 (9.6%) in the CCRT group had developed locoregional relapse, 22 (10.8%) in the IC + CCRT group and 38 (12.5%) in the CCRT group had developed distant metastasis, and 28 (13.8%) in the IC + CCRT group and 43 (14.1%) in the CCRT group had died.

| Characteristic | IC + CCRT | CCRT | P |
|---------------|-----------|------|---|
| Gender        |           |      |   |
| Male          | 147 (72.4%) | 208 (68.6%) | 0.364 |
| Female        | 56 (27.6%)  | 95 (31.4%)  |   |
| Age (years)   |           |      |   |
| <45           | 114 (56.2%) | 156 (51.5%) | 0.302 |
| ≥45           | 89 (43.8%)  | 147 (48.5%)  |   |
| T category    |           |      |   |
| 1             | 12 (5.9%)  | 27 (8.9%)  | <0.001 |
| 2             | 16 (7.9%)  | 49 (16.2%)  |   |
| 3             | 81 (39.9%) | 154 (50.8%) |   |
| 4             | 94 (46.3%) | 73 (24.1%)  |   |
| N category    |           |      |   |
| 0             | 31 (15.3%) | 48 (15.8%)  | 0.862 |
| 1             | 172 (84.7%) | 255 (84.2%) |   |
| Clinical stage|           |      |   |
| II            | 28 (13.8%) | 78 (25.1%)  | <0.001 |
| III           | 82 (39.9%) | 152 (50.8%) |   |
| IV a          | 93 (46.3%) | 73 (24.1%)  |   |
| EBV DNA       |           |      |   |
| <2000 copies/mL | 73 (36.0%)  | 186 (61.4%)  | <0.001 |
| ≥2000 copies/mL | 130 (64.0%) | 117 (38.6%) |   |

Abbreviations: IC, induction chemotherapy; CCRT, concurrent chemoradiotherapy; EBV DNA, Epstein-Barr virus deoxyribonucleic acid.

Survival Outcomes
The 5-year OS, PFS, locoregional relapse-free survival (LRFS), and DMFS rates for the entire cohort were 87.5%, 79.6%, 88.8%, and 88.7%, respectively. Median follow-up duration was 70.2 months (range, 11.4–130.1 months) and 68.5 months (range, 3.5–112.6 months) for the IC + CCRT and CCRT groups, respectively. The 5-year OS rates for the IC + CCRT and CCRT groups were 88.5% and 86.8%, respectively (p = 0.661, Figure 1A); the 5-year PFS rates were 79.6% and 79.6% (p = 0.756, Figure 1B); the 5-year LRFS rates were 87.0% and 90.2% (p = 0.364, Figure 1C), and the 5-year DMFS rates were 89.8% and 88.0%, respectively (p = 0.407, Figure 1D). There was no significant difference in OS, PFS, LRFS or DMFS between the IC + CCRT and CCRT groups.

Consistent with the results of univariate analysis, multivariate analysis to adjust for various potential prognostic factors revealed IC was not associated with significantly improved OS (HR 0.76, 95% CI 0.46–1.25, p = 0.278),
Lymph node metastasis is associated with a significantly higher rate of distant metastasis compared to patients without lymph node metastasis. Therefore, we conducted subgroup analysis to further investigate the efficacy of IC in N0 and N1 category NPC: 79 (15.6%) patients had N0 category NPC and 427 (84.4%) patients had N1 category NPC.

In N1 category NPC, no significant differences in 5-year OS (89.5% vs 86.0%, \( p = 0.587 \), Figure 2A), DMFS (89.1% vs 87.6%, \( p = 0.494 \), Figure 2B), PFS

**Prognostic Value of IC in N0 and N1 Category NPC**

Lymph node metastasis is associated with a significantly higher rate of distant metastasis compared to patients without lymph node metastasis. Therefore, we conducted subgroup analysis to further investigate the efficacy of IC in N0 and N1 category NPC: 79 (15.6%) patients had N0 category NPC and 427 (84.4%) patients had N1 category NPC.

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Table 2 Summary of Multivariable Analyses of Prognostic Factors

| Endpoint | Variable | HR (95% CI) | P          |
|----------|----------|-------------|------------|
| OS       | Age†     | 1.04 (1.01–1.06) | 0.003      |
|          | Gender†  | 1.12 (0.67–1.86) | 0.660      |
|          | EBV DNA‡ | 1.34 (0.81–2.22) | 0.253      |
|          | T3 vs T1-2 | 0.77 (0.34–1.75) | 0.530      |
|          | T4 vs T1-2 | 1.62 (0.72–3.61) | 0.243      |
|          | N1 vs N0  | 1.09 (0.57–2.07) | 0.794      |
|          | IC + CCRT vs CCRT | 0.76 (0.46–1.25) | 0.278      |
| PFS      | Age†     | 1.02 (0.79–1.83) | 0.302      |
|          | Gender†  | 1.20 (0.79–1.83) | 0.387      |
|          | EBV DNA‡ | 1.29 (0.86–1.96) | 0.221      |
|          | T3 vs T1-2 | 0.81 (0.44–1.50) | 0.498      |
|          | T4 vs T1-2 | 1.59 (0.86–2.92) | 0.139      |
|          | N1 vs N0  | 0.88 (0.52–1.47) | 0.619      |
|          | IC + CCRT vs CCRT | 0.78 (0.51–1.19) | 0.244      |
| LRFS     | Age†     | 1.00 (0.97–1.03) | 0.917      |
|          | Gender†  | 1.42 (0.81–2.48) | 0.219      |
|          | EBV DNA‡ | 1.37 (0.78–2.42) | 0.278      |
|          | T3 vs T1-2 | 0.47 (0.22–1.00) | 0.050      |
|          | T4 vs T1-2 | 0.78 (0.37–1.67) | 0.529      |
|          | N1 vs N0  | 0.66 (0.32–1.32) | 0.237      |
|          | IC + CCRT vs CCRT | 1.06 (0.81–1.42) | 0.651      |
| DMFS     | Age†     | 1.01 (0.98–1.04) | 0.340      |
|          | Gender†  | 0.66 (0.35–1.23) | 0.190      |
|          | EBV DNA‡ | 2.19 (1.24–3.85) | 0.007      |
|          | T3 vs T1-2 | 0.75 (0.37–1.52) | 0.427      |
|          | T4 vs T1-2 | 0.83 (0.39–1.76) | 0.625      |
|          | N1 vs N0  | 1.39 (0.61–3.13) | 0.433      |
|          | IC + CCRT vs CCRT | 0.66 (0.38–1.15) | 0.140      |

Notes: P-values were calculated using an adjusted Cox proportional hazards model.
†Age per year increase. †Women vs men. ‡≥4000 vs <4000 copies/mL.

Abbreviations: HR, hazard ratio; CI, confidence interval; EBV DNA, Epstein-Barr virus deoxyribonucleic acid; OS, overall survival; PFS, progression-free survival; LRFS, locoregional relapse-free survival; DMFS, distant metastasis-free survival; IC, induction chemotherapy; CCRT, concurrent chemoradiotherapy.

EBV DNA was 3995 copies/mL for DMFS. Therefore, we conducted stratified analysis of the prognostic value of IC among the 247 (48.8%) patients with ≥4000 copies/mL and 259 (51.2%) patients with <4000 copies/mL.

Among patients with ≥4000 copies/mL, 5-year OS (88.7% vs 82.8%, p = 0.214, Figure 3A), DMFS (82.7% vs 88.1%, p = 0.105, Figure 3B), LRFS (83.9% vs 90.4%, p = 0.249) and PFS (78.0% vs 75.8%, p = 0.441) were comparable between patients receiving CCRT + IC and CCRT. Moreover, among patients with <4000 copies/mL, IC did not significantly improve 5-year OS (88.0% vs 89.2.0%, p = 0.822, Figure 3C), DMFS (92.9% vs 91.4%, p = 0.719, Figure 3D), LRFS (92.6% vs 90.0%, p = 0.692) or PFS (82.4% vs 82.0%, p = 0.891).

Subgroup Analysis of the Prognostic Value of TPF IC

In the latest published Phase III trial, TPF regimen IC significantly improved failure-free survival with acceptable toxicities in locally advanced NPC. Therefore, we compared the survival outcomes of patients receiving TPF regimen IC, other IC regimens and CCRT alone.

Five-year OS (89.1% vs 86.8%, p = 0.739), DMFS (89.0% vs 88.0%, p = 0.593), PFS (77.6% vs 79.6%, p = 0.995) and LRFS (82.4% vs 90.2%, p = 0.111; Table 3) were comparable between patients receiving TPF regimen IC and patients receiving CCRT alone.

Five 5-year OS (88.2% vs 86.8%, p = 0.730), DMFS (90.5% vs 88.0%, p = 0.449), PFS (81.4% vs 79.6%, p = 0.631) and LRFS (91.0% vs 90.2%, p = 0.933) (Table 3) were also comparable between patients receiving IC based on the TP or PF regimens and patients receiving CCRT alone.

We also estimated the prognostic significance of IC for patients with T1-2N1M0 or T3-4N0-1M0 NPC. Compared to CCRT, IC + CCRT alone had no prognostic significance in T1-2N1M0 NPC (p = 0.706 for OS, p = 0.724 for DMFS, p = 0.403 for PFS, p = 0.371 for LRFS) or T3-4N0-1M0 NPC (p = 0.486 for OS, p = 0.309 for DMFS, p = 0.422 for PFS, p = 0.530 for LRFS; Table 3).

Discussion

Recent clinical trials showed the benefit of IC in local advanced NPC, the results in the real world were controversial. Three Bayesian network meta-analyses have shown IC plus CCRT reduces distant metastasis but does not improve OS in NPC compared to CCRT alone.
our study, we focus on those patients with stage T1-2N1M0 or T3-4N0-1M0 NPC, which mostly were excluded in RCT. We found that the addition of IC to radical CCRT did not lower the risk of death, progression, locoregional relapse or distant metastasis. Stratified analysis based on N category and EBV DNA further demonstrated IC did not improve the efficacy of CCRT in terms of reducing distant metastasis.

IC may not provide any value for patients at low risk of distant metastasis, and may even weaken the effect of radiotherapy. In the retrospective study of patients with stage II, IC caused deleterious effect and was a negative factor associated with OS (HR of death = 3.768, 95% CI = 1.117–12.709; P = 0.032), PFS (HR of progression = 2.156, 95% CI = 1.060–4.386; P = 0.034), LRFFS (HR of locoregional relapse = 2.435, 95% CI = 1.009–5.874;
P = 0.048) and also DMFS (HR of metastasis = 2.873, 95% CI = 1.005–8.211; P = 0.049).

IC was reported to improve DFS in stage IV N2-3 NPC compared with CCRT, indicating only selected patients benefit from IC. Advanced N category (N2-3) is well recognized as a risk factor for distant metastasis in the clinic; a clinical trial of patients with T1-4N0-3M0 is currently underway (NCT02512315). Patients with advanced N-stage (stage N2-3) NPC may not all benefit from IC. No significant difference between IC plus CCRT and CCRT was observed (p = 0.831 and 0.608, respectively) in the intermediate and low-risk groups. The results told us that careful consideration of IC must be taken, especially in those patients considered lower distant metastasis risk. In this study, we focused on stage T1-2N1M0 and T3-4N0-1M0 NPC. To identify the subgroups who may benefit from IC, we performed stratified analysis based on N category. However, IC provided no significant survival benefit in

Figure 3 Kaplan–Meier survival curves for patients with stage T1-2N1M0 or T3-4N0-1M0 nasopharyngeal carcinoma treated with IC + CCRT or CCRT alone and stratified by EBV DNA copy number (≥4000 vs <4000 copies/mL). (A and C) Overall survival; (B and D) distant metastasis-free survival.

Abbreviations: IC, induction chemotherapy; CCRT, concurrent chemoradiotherapy; NPC, nasopharyngeal carcinoma; EBV DNA, Epstein-Barr virus deoxyribonucleic acid.
the subgroup analysis. These negative results could be 
attributed to three factors. Firstly, IC may not actually 
reduce distant metastasis. Secondly, the relatively small 
sample sizes in each subgroup may have resulted in low 
statistical power, so differences between subgroups could 
not be detected. Third, several large-scale Phase III trials 
have confirmed adding docetaxel to PF regimen IC pro-
vides a significant clinical benefit. However, a large-
scale propensity-matched study indicated taxane-based IC 
could not improve survival in locoregionally advanced 
NPC, but decreased the risk of distant metastasis in stage 
T4N1-2M0 and stage IVb NPC. In this study, we only 
assessed patients with stage T1-2N1M0 and T3-4N0-1M0 
NPC; 46.8% of patients received TPF and 22.7% received 
PF. The use of different IC chemotherapy regimens in this 
cohort may provide a slight bias towards a negative result. 
However, subgroup analysis was used to reduce the poten-
tial bias associated with different chemotherapy regimens, 
and demonstrated the TPF regimen did not have any sig-
nificant prognostic value. Further studies are required to 
verify these results. Finally, more cycles of IC might be 
essential. Four cycles of IC were effective and well 
tolerated. A trial on 4 cycles of neoadjuvant chemotherapy 
plus concurrent chemoradiation in N2-3 nasopharyngeal 
carcinoma is ongoing (NCT02512315).

EBV DNA is the most effective prognostic biomarker for 
guiding the treatment of NPC. High pre-treatment EBV 
DNA can accurately predict distant metastasis. Detectable EBV DNA is associated with poorer DMFS 
compared to undetectable EBV DNA. IC was previously 
reported to improve the survival of patients with high EBV 
DNA. In our study, EBV DNA (≥4000 vs <4000 copies/mL) was an independent prognostic factor for DMFS (HR 2.19, 95% CI 1.24–3.85, p = 0.007). However, IC did not 
provide a significant survival benefit in stratified analysis of 
patients with ≥4000 or <4000 EBV DNA copies/mL, which 
might due to the unbalanced patients. In our enrolled patients, 
64% of patients who received IC had EBV DNA ≥ 4000 
copies/mL, while CCRT was 38%. In clinical, Patients with 
high EBV NDA copies might be treated with greater inten-
sity treatment. However, a propensity-matched analysis of 
stage III–IVb patients with ≥4000 EBV DNA copies/mL 
showed IC did not improve the clinical outcomes of these 
high-risk patients compared to CCRT alone, while IC only 
improved OS in very high-risk patients (N2-3 category with 
EBV DNA ≥ 4000 copies/mL). Hence, the use of IC should 
be carefully considered in patients with stage T1-2N1M0 and 
T3-4N0-1M0, even though some of these patients have 
a high EBV DNA copy number. Further prospective studies 
are needed to more precisely evaluate the role of EBV DNA 
for guiding the treatment of different subgroups of patients 
with NPC. A clinical trial of individualized treatment of 
patients with stage II–IVb NPC based on EBV DNA is 
currently ongoing (NCT02135042).

Table 3 Subgroup Analysis of the Prognostic Value of Induction Chemotherapy

| OS | DMFS |
|----|------|
| **Number of Events** | **Total Number** | **5-Year OS** | **P** | **Number of Events** | **Total Number** | **5-Year DMFS** | **P** |
| TPF Regimen IC | | | | | | | |
| IC + CCRT | 12 | 95 | 89.1% | 0.039 | 10 | 95 | 89.0% | 0.593 |
| CCRT alone | 43 | 303 | 86.8% | | 38 | 303 | 88.0% | |
| TP or PF Regimen IC | | | | | | | |
| IC + CCRT | 16 | 108 | 88.2% | 0.012 | 12 | 108 | 90.5% | 0.449 |
| CCRT alone | 43 | 303 | 86.8% | | 38 | 303 | 88.0% | |
| T1-2N1M0 | | | | | | | |
| IC + CCRT | 3 | 28 | 91.6% | 0.025 | 4 | 28 | 82.9% | 0.724 |
| CCRT alone | 5 | 76 | 86.3% | | 8 | 76 | 87.5% | |
| T3-4N0-1M0 | | | | | | | |
| IC + CCRT | 25 | 175 | 88.2% | 0.048 | 18 | 175 | 90.6% | 0.309 |
| CCRT alone | 38 | 227 | 86.1% | | 30 | 227 | 88.1% | |

Notes: TPF was 60 mg/m² docetaxel on day 1, 60 mg/m² cisplatin on day 1 and 600 mg/m²² fluorouracil civ on days 1–5; TP was 75 mg/m²² docetaxel on day 1 and 75 mg/m²² cisplatin on day 1; PF was 80 mg/m²² cisplatin on day 1 and 800 mg/m²² fluorouracil civ on days 1–5. P-values were calculated using the Log-rank test.

Abbreviations: OS, overall survival; DMFS, distant metastasis-free survival; IC, induction chemotherapy; CCRT, concurrent chemoradiotherapy.
The exploration on IC for local advanced NPC is still ongoing. Whether benefit of IC is uncertain. The incidences of toxicity (a higher risk of neutropenia and weight loss, and so on) were increased proved by the clinical trials. Other than improving distant metastasis control, one of the targets of IC in clinical for oncologist might be to shrink the tumor volume to reduce the side reaction of radiation therapy. A lower risk of mucositis and hearing loss was found in IC+CCRT compared to CCRT. The proportion of patients with eye damage was significantly lower in the IC followed by CCRT alone group than the CCRT group (9.7% versus 16.4%, p= 0.029). Furthermore, an interesting finding from Taiwan which the patients with stage N2-3 accounting 85% showed that IC significantly improved the LRFS (HR=0.74, p=0.026) and failed to reduce distant metastasis (HR=0.82, p=0.2828). So selecting patients with nasopharyngeal carcinoma who benefit from induction chemotherapy was important. A prognostic nomograms based on 8th edition of the UICC staging system found that IC could only improve OS for patients with stage III–IV at intermediate to high risks (including Histological type, T category, N category, plasma Epstein-Barr virus deoxyribonucleic acid, age and neutrophil-lymphocyte ratio). In the past few years, researchers and we focused on the clinical characteristics to select patients who can benefit from IC. More and more attentions are paid on the immune molecules and the immune system. Increased tumour-infiltrating lymphocytes (TIL) concentration predicted response to neoadjuvant chemotherapy in breast cancer, and was also associated with a survival benefit in HER2-positive breast cancer and TNBC. Infiltrating lymphocytes (IL)17A cells infiltration could be used as an independent prognostic biomarker for OS and predictive biomarker for superior response to adjuvant chemotherapy. Stromal immunotype predict survival and benefit from adjuvant chemotherapy in patients with muscle-invasive bladder cancer. TIL, immunotype might predict the response to chemotherapy. In the future, we might need to combine clinical characteristics and immune molecules and the immune system to select NPC patient who can benefit from IC. Further research investigating is warranted.

There are some limitations that must be noted. First, we just focused on the clinical characteristics to select patients who can benefit from IC. In future study, we should pay attention to molecular markers, such as TIL, immune classification and so on. Another limitation was the heterogeneity of the IC regimen and dosage because of the retrospective study design. Finally, the data on cute and late toxicities, treatment-related toxicities were lack due to our failing to collect in our intelligence platform in the current study.

In conclusion, IC + CCRT did not improve any survival outcome in patients with stage T1-2N1M0 or T3-4N0-1M0 NPC receiving IMRT compared with CCRT alone. IC has limited prognostic value in moderate-risk NPC, and should not be routinely prescribed in clinical practice to patients with T1-2N1M0 and T3-4N0-1M0 NPC. However, as this was a retrospective single-center study, prospective studies are warranted to confirm the value of IC in moderate-risk NPC.

### Abbreviations
NPC, nasopharyngeal carcinoma; OS, overall survival; CCRT, concurrent chemoradiotherapy; NCCN, National Comprehensive Cancer Network; IMRT, intensity-modulated radiation therapy; IC, induction chemotherapy; AC, adjuvant chemotherapy; PFS, progression-free survival; DMFS, distant metastasis-free survival; EBV DNA, Epstein-Barr virus deoxyribonucleic acid; AJCC, American Joint Committee on Cancer; MRI, magnetic resonance imaging; LRFS, locoregional relapse-free survival; PET/CT, positron emission tomography/computed tomography; ROC, receiver operating characteristic; HR, hazard ratios; CI, confidence intervals.

### Data Sharing Statement
All data generated or analyzed during the current study are available from the corresponding author upon reasonable request.

### Ethics Statement
This retrospective study was approved by the Clinical Research Ethics Committee of Sun Yat-Sen University Cancer Canter. All procedures performed in studies involving human participants were in accordance with the ethical standards of the institutional and/or national research committee and with the 1964 Helsinki Declaration and its later amendments or comparable ethical standards.

### Informed Consent
Informed consent was not required for this research due to the anonymized patient data.

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**Author Contributions**

ZS, GRZ, JT, XYL and FYX contributed to conception and design of the study; ZS, GRZ, JT, and XYL contributed to acquisition of data, or analysis and interpretation of data; ZS, GRZ, JT, and XYL contributed to drafting the article. ZS, GRZ, and FYX contributed to revising the article critically for important intellectual content. ZS, GRZ, JT, and XYL gave final approval of the version to be published, and agreement to be accountable for all aspects of the work in ensuring that questions related to the accuracy or integrity of any part of the work are appropriately investigated and resolved.

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**Disclosure**

The authors have no competing interests to declare.

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