Accuracy of $^{18}$F-fluorodeoxyglucose-positron emission tomography/computed tomography in the staging of newly diagnosed nasopharyngeal carcinoma: a systematic review and meta-analysis

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Radiol Oncol 2014; 48(4): 331-338.

Received: 9 October, 2013
Accepted: 14 January, 2014

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Disclosure: No potential conflicts of interest were disclosed.

Background. The specific role of $^{18}$F-fluorodeoxyglucose-positron emission tomography/computed tomography (FDG-PET/CT) in staging of nasopharyngeal carcinoma (NPC) remains to be validated. A systematic review and meta-analysis were performed to assess the accuracy of staging FDG-PET/CT for newly diagnosed NPC.

Methods. We searched various biomedical databases and conference proceedings for relevant studies. We determined the pooled sensitivities and specificities, diagnostic odds ratios (DOR) and constructed summary receiver operating characteristic (SROC) curves using the hierarchical regression model.

Results. 15 relevant studies including 851 patients were identified. Five addressed primary tumor (T), nine addressed regional lymph nodes (N) and seven addressed distant metastasis (M). The combined sensitivity estimate for FDG-PET/CT in T classification was 0.77 (95% confidence interval [CI] 0.59-0.95). For N classification, combined sensitivity was 0.84 (95% CI 0.76-0.91), specificity was 0.90 (95% CI 0.83-0.97), DOR was 82.4 (23.2-292.6) and Q*-index was 0.90. For M classification, the combined sensitivity estimate was 0.87 (95% CI 0.74-1.00), specificity was 0.98 (95% CI 0.96-1.00), DOR was 120.9 (43.0-340.0) and Q*-index was 0.89.

Conclusion. FDG-PET/CT showed good accuracy in N and M but not T classification for newly diagnosed NPC. FDG-PET/CT, together with Magnetic resonance imaging (MRI) of the nasopharynx, should be part of the routine staging investigations.

Key words: nasopharyngeal carcinoma; PET/CT; staging; accuracy; meta-analysis

Introduction

In 2008, there were approximately 84400 new cases of nasopharyngeal carcinoma (NPC) and 51600 deaths from the disease worldwide.¹ The geographical disparities in the burden of NPC are noteworthy, with incidence rates highest in East and Southeast Asia and lowest in Central America.¹

NPC may spread locally to involve the parapharyngeal soft tissue, base of skull or intracranial structures. The nasopharynx has a rich lymphatic plexus; 75% of patients present with enlarged cer-
vical nodes, 80% of whom have bilateral involvement. NPC has a relatively high incidence of systemic metastasis (up to 41%) when compared with the other head and neck tumors (5%–24%). The most common sites of metastases are bone (20%), lung (13%), and liver (9%).

NPC is staged non-surgically and treated primarily with radiotherapy (with or without chemotherapy). Accurate staging is essential as it influences the choice of treatment modalities, radiotherapy planning and prognosis. Combined modality treatment, as well as larger treatment volumes, invariably leads to greater toxicities. Although FDG-PET/CT is sometimes used in the clinical management of NPC in preference to other imaging modalities, such computed tomography or bone scans, the magnitude of benefit of using FDG-PET/CT, if any, is unclear. Indications for its use in the clinic have been rather empirical than standardized in many centres, often in the setting of a diagnostic dilemma affecting treatment options after the use of conventional imaging modalities.

The American Joint Committee on Cancer (AJCC) T (Primary tumor) N (Regional lymph nodes) M (Distant Metastasis) system is one of the most widely used staging system internationally. Conventional staging modalities may include MRI of the head and neck, contrast enhanced CT scans, liver ultrasound (US) and whole body radionuclide bone scan (WBBS). For M classification, one series reported the sensitivity and specificity for conventional workup (chest X-Ray, liver US, WBBS) to be 0.33 and 0.90 respectively; the same series reported the sensitivity and specificity for conventional workup in TNM staging of newly diagnosed treatment naïve NPC patients, with reference to conventional modalities and/or clinical follow up.

The National Comprehensive Cancer Network (NCCN) guidelines recommend gadolinium-enhanced MRI of the nasopharynx and neck as well as CT scan (if indicated, for T and N classifications). It recommends imaging of distant metastases in the chest, liver and bones (which may include PET scan and/or CT) for patients with N2-3 disease. It also suggests that FDG-PET/CT scan may be considered for patients with Stage III and IV disease.

The use of FDG-PET/CT has superseded stand-alone FDG-PET studies, by offering both functional and anatomic imaging, for the initial staging and post-treatment assessments for a wide range of cancers. Published individual studies in the medical literature have reported increased accuracy especially in detection of metastases but are less conclusive on local and regional staging. The role of FDG-PET/CT in the overall staging of pre-treated NPC remains to be validated. To our knowledge, only one systematic review and meta-analysis of six studies examining the accuracy of FDG-PET/CT in detection of distant metastasis in pre-treated NPC showed it to have a high sensitivity of 88% and specificity of 97% for M classification. However, there were some limitations of this meta-analysis. Firstly, it did not address the accuracy of PET/CT scan for T and N classifications. Additionally, it excluded several publications in languages other than English and finally, new data have been published since the meta-analysis.

The aim of our study was to perform a systematic review and meta-analysis of all relevant publications to determine the accuracy of FDG-PET/CT in the TNM staging of newly diagnosed treatment naïve NPC patients, with reference to conventional modalities and/or clinical follow up.

### Materials and methods

#### Identification and eligibility of relevant studies

We included studies, without language restriction, that determined the sensitivity and specificity of FDG-PET/CT for TNM staging of pre-treated (biopsy proven) nasopharyngeal cancer, when compared to conventional staging modalities (i.e. MRI or CT scan of head and neck for T and N classifications, biopsy or clinical follow up of suspected metastases to regional lymph nodes or distant sites).

We searched MEDLINE, Cochrane CENTRAL register of controlled trials, Cochrane Database of systematic reviews, Chinese national knowledge infrastructure (CNKI) and China Biomedical Literature Disc (CBMDisc) from date of inception to September 2011 and meeting proceedings of American Society for Radiation Oncology (ASTRO) and American Society of Clinical Oncology (ASCO) from 2000 to September 2011.

We used a search algorithm that included the following terms: (1) PET OR 18F-FDG PET OR positron emission tomography; (2) nasopharyngeal cancer OR nasopharyngeal carcinoma OR cancer of the nasopharynx OR lymphoepithelioma; (3) staging OR detection OR lymph node OR metastasis OR TNM.

FDG-PET only studies were excluded. For N and M classifications, studies that did not provide sufficient information to construct 2 x 2 table for sensitivity and specificity calculations were excluded. For T classification, we chose to analyze the sensitivity of FDG-PET/CT, in comparison to the
reference standard. (i.e verifying false positive and true negative results in a non-surgically staged tumor would be impossible, and likely not reported in published studies).

The most recent publication was chosen when data was presented in more than one publication.

Two reviewers (B.V and S.Y.Y) independently judged study eligibility and disagreements were resolved by discussion and if necessary by a third reviewer (L.K.M).

**Data extraction**

Two reviewers (B.V and S.Y.Y) extracted data from each eligible study independently using a standardized data extraction form and any disagreements were resolved by discussion or by appeal to a third reviewer (L.K.M).

Reviewers were not blinded with regard to information about the journal name, the authors, country of origin or the year of publication; as this has been shown to be unnecessary. In addition, we recorded the following information: study design (retrospective/prospective), sample size, age and gender distribution, stage of patients included and reference tests used to define extent of disease. Publications looking at more than one aspect of the study’s methodology. Sensitivity analyses were performed after exclusion of retrospective studies, which utilized MRI to be the only acceptable reference standard. We performed a subgroup analysis considering studies, which utilized MRI to be the only acceptable reference test (versus MRI or CT or clinical findings). This may be viewed as a non-inferiority comparison or concordance of FDG-PET/CT to MRI.

Subgroups to be analyzed were determined a-priori, with the following reasons:

**T classification.** Contrast enhanced MRI is considered to the current gold standard for soft tissue involvement and intracranial extension. A subgroup analysis was performed considering studies, which utilized MRI to be the only acceptable reference test (versus MRI or CT or clinical findings). This may be viewed as a non-inferiority comparison or concordance of FDG-PET/CT to MRI.

**N classification**. FDG-PET/CT may over or underestimate the involvement of retropharyngeal and parapharyngeal lymph nodes; possibly because of poor distinction from the primary nasopharyngeal tumour. A subgroup analysis was done for studies looking primarily at cervical lymph nodal involvement versus non-cervical lymph nodes (i.e. retro/parapharyngeal). As neck dissection is not part of standard staging, it is unlikely to have histopathology as the reference standard. We performed a subgroup analysis to see if there was a difference between studies that required histology versus those that did not.

**M classification.** We performed a subgroup analysis to determine if there was a difference between studies
T classification. Based on the combined data from five available studies that evaluated the T-classification our analysis revealed a sensitivity of 0.77 (95% CI 0.59-0.95) while no specificity level could be ascertained (Figure 2). Four (of the five) studies did not report false positive results hence preventing us from calculating the specificity for T classification. Subgroup analysis revealed the sensitivity of FDG-PET/CT was lower when compared to MRI alone; however, this was not statistically significant (0.65 vs. 0.86, \(P=0.214\)). The sensitivity results on T classification were similar with exclusion of the two low quality studies, or the two retrospective studies.

N classification. The combined sensitivity estimate for N classification is 0.84 (95% CI 0.76-0.91) and specificity 0.90 (95% CI 0.83-0.97). The pooled DOR for N classification was 82.4 (23.2-292.6). The Q*-index was 0.90 (SE 0.03) (Figure 3). The reference standards used for N classification varied amongst studies. MRI neck was the most frequently used reference standard. Two studies relied on clinical follow up to be their reference standard, and 2 other studies required histological confirmation though fine needle aspiration of involved cervical nodes. One study used contrast enhanced CT to be their reference standard, which is considered to be inferior to MRI. The effect on sensitivity was significantly lower for studies assessing retro/parapharyngeal nodal involvement (0.94 vs. 0.44, \(p<0.001\)) whereas specificity did not differ significantly (0.85 vs. 1.00).
TABLE 1. Characteristics of included studies

<table>
<thead>
<tr>
<th>Class.</th>
<th>Author (year)</th>
<th>Sample size</th>
<th>Design</th>
<th>Age (years)</th>
<th>Male (%)</th>
<th>Patient population</th>
<th>Reference Test</th>
<th>Quadas</th>
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<tbody>
<tr>
<td>T</td>
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<td>20</td>
<td>R</td>
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<td>All comers</td>
<td>nasoscopy and CT/MR</td>
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<td>King (2008)</td>
<td>52</td>
<td>R</td>
<td>50</td>
<td>73</td>
<td>Stage III-IV</td>
<td>MRI</td>
<td>13</td>
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<tr>
<td></td>
<td>Ng (2009)</td>
<td>111</td>
<td>P</td>
<td>48.9</td>
<td>75.6</td>
<td>All comers</td>
<td>MRI</td>
<td>13</td>
</tr>
<tr>
<td></td>
<td>Wu (2011)</td>
<td>12</td>
<td>P</td>
<td>49</td>
<td>66.7</td>
<td>All comers, looking at intracranial and intraorbital extension [T4]</td>
<td>MRI/CT and clinical finding</td>
<td>9</td>
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<tr>
<td></td>
<td>Cai (2011)</td>
<td>25</td>
<td>P</td>
<td>50</td>
<td>64</td>
<td>Locally advanced NPC (at least T3)</td>
<td>MRI/CT and clinical findings</td>
<td>12</td>
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<tr>
<td></td>
<td>N Hu (2005)</td>
<td>105</td>
<td>P</td>
<td>43</td>
<td>78</td>
<td>All comers</td>
<td>Followup for 9 months</td>
<td>10</td>
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<tr>
<td></td>
<td>Su (2006)</td>
<td>53</td>
<td>P</td>
<td>40</td>
<td>68</td>
<td>All comers</td>
<td>MRI – looking at retropharyngeal LN</td>
<td>11</td>
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<tr>
<td></td>
<td>Chen (2006)</td>
<td>20</td>
<td>R</td>
<td>46.3</td>
<td>70</td>
<td>All comers</td>
<td>CT</td>
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<tr>
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<td>Zhang (2006)</td>
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<td>P</td>
<td>NR</td>
<td>79.3</td>
<td>All comers</td>
<td>Followup for 9 months</td>
<td>10</td>
</tr>
<tr>
<td></td>
<td>Tang (2007)</td>
<td>87</td>
<td>P</td>
<td>43</td>
<td>72.8</td>
<td>All comers</td>
<td>MRI - looking at para/retropharyngeal LN</td>
<td>13</td>
</tr>
<tr>
<td></td>
<td>Lin (2008)</td>
<td>68</td>
<td>P</td>
<td>41</td>
<td>58.5</td>
<td>All comers</td>
<td>MRI neck</td>
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<tr>
<td></td>
<td>King (2008)</td>
<td>52</td>
<td>R</td>
<td>50</td>
<td>73</td>
<td>Stage III-IV</td>
<td>MRI neck</td>
<td>12</td>
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<tr>
<td></td>
<td>Ng (2009)</td>
<td>17</td>
<td>P</td>
<td>48.9</td>
<td>75.6</td>
<td>All comers</td>
<td>FNA</td>
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<tr>
<td></td>
<td>Lin (2009)</td>
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<td>R</td>
<td>NR</td>
<td>NR</td>
<td>All comers</td>
<td>FNA</td>
<td>7</td>
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<tr>
<td></td>
<td>M Chen (2006)</td>
<td>20</td>
<td>R</td>
<td>46.3</td>
<td>70</td>
<td>All comers</td>
<td>Histological proof, or clinical followup for 6 months</td>
<td>11</td>
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<tr>
<td></td>
<td>Wang (2007)</td>
<td>18</td>
<td>R</td>
<td>52</td>
<td>60.5</td>
<td>All comers</td>
<td>Histological proof, or clinical followup for 17 months [median]</td>
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<tr>
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<td>King (2008)</td>
<td>52</td>
<td>R</td>
<td>50</td>
<td>73</td>
<td>Stage III-IV</td>
<td>Histological proof, or clinical followup for 12 months</td>
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<tr>
<td></td>
<td>Chua (2009)</td>
<td>78</td>
<td>P</td>
<td>50</td>
<td>74.9</td>
<td>All comers</td>
<td>Histological proof, or clinical followup for 6 months</td>
<td>11</td>
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<tr>
<td></td>
<td>Ng (2009)</td>
<td>150</td>
<td>P</td>
<td>48.1</td>
<td>74</td>
<td>All comers</td>
<td>Histological proof, or clinical followup for 12 months</td>
<td>11</td>
</tr>
<tr>
<td></td>
<td>Lin (2009)</td>
<td>41</td>
<td>R</td>
<td>NR</td>
<td>NR</td>
<td>All comers</td>
<td>Clinical followup [time not specified]</td>
<td>6</td>
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<tr>
<td></td>
<td>Iaguru (2011)</td>
<td>26</td>
<td>R</td>
<td>47.3</td>
<td>69.2</td>
<td>All comers</td>
<td>Clinical followup [time not specified]</td>
<td>9</td>
</tr>
</tbody>
</table>

FNA = fine needle aspirate cytology; M = distant metastasis; N = regional lymph nodes; NPC = nasopharyngeal carcinoma; NR = not reported; P = prospective; R = retrospective; T = primary tumor.

There was no significant difference in sensitivity or specificity between studies that required histological confirmation (0.93 vs. 0.82, P=0.666; 0.82 vs. 0.91, P=0.533). The sensitivity and specificity results on N classification were similar with exclusion of two low quality studies,10,19 or the three retrospective studies.10,11,19

**M classification.** The combined sensitivity estimate for M classification is 0.87 (95% CI 0.74-1.00), and specificity 0.98 (95% CI 0.96-1.00). The pooled DOR for M classification is 120.9 (43.0-340.0). The Q*-index is 0.92 (SE 0.02) (Figure 4). All studies used either histological proof or clinical follow up (range 6-17 months) to define true positive and true negative lesions. Two studies used clinical follow up alone,10,12 and the duration was not reported. The mean time of follow up for the remaining studies was 12 months. Subgroup analysis did not show any significant differences for pooled sensitivity or specificity (1.00 vs. 0.84, P=0.996; 0.99 vs. 0.98, P=0.531). Sensitivity analysis showed that the results on M classification were similar with exclusion of the three low quality studies,9,10,12 or the five retrospective studies.3-12,19

**Discussion**

This meta-analysis suggests that FDG-PET/CT has excellent sensitivity and specificity compared
to conventional staging modalities for N and M but not for T classification of NPC. We observed that FDG-PET/CT might be less accurate to determine involvement of para/retropharyngeal lymph nodes, although this estimate may be imprecise owing to relatively small number of studies.

Compared to other published meta-analyses investigating the accuracy of FDG-PET/CT, our results showed similar results for M classification but superior results for N classification. Nevertheless, we should note there are intrinsic differences. Kyazs et al. looked at the utility of FDG-PET (without combined CT) for cervical nodal metastasis in squamous cell head and neck cancer, referencing it against surgical specimens. The review did not find good evidence to support the routine use of pretreatment evaluation FDG PET. They reported an overall sensitivity and specificity of 0.79 and 0.86 respectively. The sensitivity was significantly lower in the clinically negative neck (0.50).

The variation in reported results may be due to the improved accuracy of integrated FDG-PET/CT versus stand-alone FDG-PET, differing reference standards (conventional methods versus surgical specimen) and differing primaries (NPC versus non-NPC). Our results did not differ after the inclusion of Chinese language publications for M classification, as previously reported by Xu and colleagues.

The strengths of this study are that it addresses a pragmatic question, incorporates recently published data, includes Chinese language based publications, has a standardized study quality assessment, and has a pre-planned sub-group analysis to address potential sources of heterogeneity. Additionally, sensitivity analyses showed consistent results, suggesting the robustness of the findings.

There are some limitations of this meta-analysis. Firstly, our review was based on published results and not individual patient data. Secondly, the imaging reference standards used for T and N classifications were heterogeneous and subject to interpretation. The follow up time for M classification varied (6-17 months) and there was no consistent follow up strategy. Lastly, the included studies were heterogeneous in design though the majority of the studies were of low-moderate risk of bias based on the QUADAS assessment.

FIGURE 2. Pooled sensitivity for T classification.

FIGURE 3. For N classification: (A) Pooled sensitivity (B) Pooled specificity (C) Pooled diagnostic odds ratio (D) Summary receiver operating characteristic (SROC) curve with Q*-index.
In conclusion, FDG-PET/CT showed good accuracy in N and M but not T classification for newly diagnosed pre-treated NPC. While head and neck MRI is still recommended for T classification, FDG-PET/CT is accurate for clinical staging of regional nodes and distant disease and can be considered as an alternative standard of care wherever available. The diagnostic superiority of FDG-PET/CT over conventional staging modalities for detection of metastatic disease makes for more accurate disease prognostication and optimization of treatment strategy. The additional information derived from the FDG-PET/CT can also potentially aid neck nodal target delineation. FDG-PET/CT, together with MRI of the head and neck, has become part of the routine staging investigations for NPC at our centre. Future research should investigate the accuracy of FDG-PET/MRI as a single staging modality for NPC.

References


