A Case Report of False Positive FDG-PET/CT Mediastinal Lymph Node in Oesophageal Adenocarcinoma Revealed to be an Anthracotic and Anthracosilicotic Spindle Cell Pseudotumor (AASCP)

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Abstract

Anthracotic and anthracosilicotic spindle cell proliferation (AASCP) is a rare reactive proliferative entity of phagocytic histiocytes which can affect hilar and mediastinal lymph nodes. Fluorodeoxyglucose-positron emission tomography (FDG-PET) has been used for the clinical diagnosis and staging of oesophageal adenocarcinoma. We report the first case of AASCP exhibiting false positivity on FDG-PET imaging.

A 76-year-old man with distal esophageal adenocarcinoma underwent PET/CT for staging revealing a positive mediastinal lymph node involvement. The patient underwent Endobronchial Ultrasound Fine Needle Aspiration Biopsy (EBUS-FNA) followed by an oesophagectomy with lymph node dissection. The histological diagnosis of the mediastinal lymphadenopathy was AASCP. In mediastinal lymph nodes of esophageal adenocarcinoma, AASCP could be a cause of false-positivity on FDG-PET imaging.

Case Report

We report the case of a 76-year-old male who on investigation for iron deficiency anemia was found to have a distal esophageal adenocarcinoma measuring 2 cm in size and is located at 35 cm from the incisors (Figure 1). His medical history is otherwise positive for type 2 Diabetes Mellitus with a controlled blood sugar and HbA1c level. Social history was significant for living in the Northern Lebanese area of Akkar, which is close to areas of industrial growth and air pollution.

Endoscopic Ultrasound (EUS) showed the tumor to be invading the muscularis propria with negative lymph node T2N0 (Clinical Stage IB). Computed Tomography of the chest and abdomen showed a left 7th rib sclerotic lesion suggestive of metastasis. As a result, a PET/CT whole body with F18-Flurorodeoxyglucose (FDG) scan was performed. The PET/CT scan showed wall thickening and increased activity within the distal esophagus at the site of the known esophageal tumor and an unexpected 1 cm right lower paratracheal (4R) lymph node showing increased radiotracer uptake with a Standardized Uptake Value SUV-max of 4.53 (Figure 2).

Considering the PET/CT results as a positive nodal invasion, the patient wasupstaged to a T2N1 Stage IIB, requiring as a result neoadjuvant treatment. The question whether this is a metastatic node was entertained in our multidisciplinary tumor board meeting and an Endobronchial Ultrasound Biopsy Fine Needle aspiration (EBUS-
FNA) was recommended and performed and turned out to be negative. The patient proceeded to surgery directly and an Ivor Lewis Esophagectomy was performed successfully. The final histopathology staged the disease as T1N0M0 moderately differentiated adenocarcinoma (Grade 2) of the esophagus Stage 1A. The questionable 4R lymph node turned out to be an AASCP.

Figure 1: Distal esophageal tumor measuring 2 cm in size and located at 35 cm from the incisors.

Figure 2: Man with FDG avid lower esophageal cancer and active right paratracheal lymph node (SUVmax=4.53). No evidence of distant metastasis.

The received “level 4R” mediastinal lymph node measured 1.5 x 1.3 x 1.1 cm and had a slightly firm consistency. The cut surface was smooth, tan black and homogenous. Microscopically, on low power view, the lymph node was partially involved by a proliferative lesion in a vague storiform pattern. On high magnification the lesion was composed of bland spindle cells many of which contained variable amounts of anthracotic pigment and interspersed dense collagen deposition.

The other submitted “4R” lymph nodes were also anthracotic. The patient underwent a transbronchial FNA prior to operative resection which was adequate for analysis and negative for adenocarcinoma. It was composed of mature small lymphocytes, normal bronchial epithelium and numerous anthracotic pigment-containing cells either spindle or polygonal with an abundant background anthracotic pigment (Figure 3).

Figure 3: (A) AASCP. H and E stain. Low power view (40X) shows the proliferative process partially involving the lymph node. (B) Prominent fibrosis. (C and D) The spindle nature of the cells along with the prominent anthracosis (100X, 400X).

Immunohistochemically the lesional cells were diffusely positive for CD68, SMA, MSA and Vimentin, focally positive Caldesmin and negative for ALK, CD34, CD21, Calponin, Desmin, S100 and Melan A. The proliferative index (Ki-67) was approximately 10%. Special stains were performed on the submitted tissue; Masson trichrome highlighted the dense collagen fibers. Both melanin and iron stains were negative. Examination of tissue sections with a polarizer light was negative (Figure 4).

Figure 4: (A) Immunohistochemistry staining. Proliferative index (Ki67). (B) Smooth muscle actin (SMA).

Based on the above findings a diagnosis of “anthracotic and anthracosilicotic spindle-cell pseudotumor” was made. This is a rare reactive process that results in significant fibrosis and collagen deposition and in which there is conversion of macrophages into numerous spindle cells mimicking neoplastic processes. It can occur in the absence of established lung silicosis and may show histologically an infiltrative and destructive growth with extracapsular and destructive spread mimicking sarcoma [7].

The differential diagnosis in this case included; follicular dendritic cell tumor (FDCT), Mycobacterial spindle-cell tumor, Spindle cell melanoma, Kaposi's sarcoma and Intranodal hemorrhagic spindle cell tumor with amianthoid fibers (Palsaded myofibroblastoma), and inflammatory myofibroblastic tumor.
The other diagnostic possibilities were excluded as follows: Negative CD21 ruled out Follicular dendritic cell tumor. Spindle cell melanoma was ruled out by bland cytolgy of the spindle cells and negative S100 and Melan A stain. Intramedullary hemagic spindle cell tumor (Palisaded myofioblastoma) was excluded due to its occurrence mainly in the groin region and the absence of the characteristic amianthoid fibers. Mycobacterial spindle cell tumor and Kaposis sarcoma have histologic features that differ slightly from those in our case. Additionally both entities occur in immunocompromised and or AIDS patients. Our patient hoverer however is immune-competent making these diagnoses untenable.

Discussion

Adenocarcinoma of the esophagus is a malignant epithelial tumor of the esophagus with glandular differentiation arising predominantly from Barrett mucosa in the lower third of the esophagus. At the time of diagnosis, most tumors are advanced with deep infiltration of the esophageal wall [8]. Adenocarcinomas spread first locally and infiltrate the esophageal wall with a possible extension through the esophageal wall into adventitial tissue, and then into adjacent organs or tissues are similar to squamous cell carcinoma. Barrett associated adenocarcinoma metastasizes to para-esophageal and paracardial lymph nodes, those of the lesser curvature of the stomach and the celiac nodes.

The major prognostic factors in adenocarcinoma of the esophagus are the depth of mural invasion and the presence or absence of lymph node or distant metastasis. Conventional staging methods include upper endoscopic gastroduodenoscopy, endoscopic ultrasonography (US), and computed tomography (CT) of the thorax and abdomen. The routine use of integrated positron emission tomography (PET)/CT with 18-fluoro-2-deoxy-d-glucose (FDG) in evaluation of patients with esophageal cancer is increasing, it has been reported to be useful in initial staging, follow-up and assessment of therapeutic response, and detection of recurrent malignancy [9,10].

Locoregional lymph nodes in esophageal cancer depend to some extent on the location of the primary tumor in the esophagus and are normally resected with the primary tumor at the time of esophagectomy [11]. Limitation of PET/CT in the evaluation of nodal metastasis was mainly due to the FDG uptake within periesophageal lymph nodes that are anatomically close to the primary tumor which is difficult to differentiate from uptake within the esophagus itself. This is mainly due to the limited spatial resolution of PET [12]. In addition, PET can result in false negative outcome in cases of microscopic metastatic disease within lymph nodes that may not demonstrate sufficient FDG uptake for detection [13]. Furthermore, FDG uptake within lymph nodes can occur in benign disease such as granulomatous infection (secondary to histoplasmosis or tuberculosis) or Sarcoïdosis.

This entertains the importance of taking a detailed exposure or occupational history from patients presenting with mediastinal lymphadenopathy, which was missednot clearly investigated for in our initial investigationsinitially. In a metanalysis done by van Westreenen et al. PET/CT had a sensitivity of 51% and specificity of 84% for detection of nodal metastases in preoperative staging setting in patient diagnosed with esophageal cancer [14].

No association between adenocarcinoma of the esophagus and AASCP has been reported to date in the literature.

Endoscopic US is superior to PET/CT in detection of locoregional nodal metastases and has the added advantage over PET/CT of allowing biopsy of suspicious lymph nodes at the time of detection [15]. However PET/CT is still considered as a reliable modality in the initial staging and for assessment of treatment response especially in the evaluation of distant metastasis [16]. PET/CT is shown in a recent study to provide incremental staging information, changing management in one third of patients [17].

Conclusion

Although being a rare finding, AASCP which is a benign condition should be considered in the differential diagnosis of nodal involvement in esophageal cancer on PET/CT imaging. PET/CT is still considered as an interesting method in the initial staging and for –assessment of treatment response especially in the evaluation of distant metastasis. However it has a limited role in the evaluation of locoregional nodal involvement in esophageal cancer and its results should be interpreted in conjunction of with endoscopic US and pathologic data.

References

