FDG-PET/CT in lymphoma

Maria M D’souza, Abhinav Jaimini, Abhishek Bansal, Madhavi Tripathi, Rajnish Sharma, Anupam Mondal, Rajendra Prashad Tripathi
Departments of PET Imaging, Institute of Nuclear Medicine and Allied Sciences, Brig. SK Majumdar Marg, ‘Nuclear Medicine, AIIMS, New Delhi, India

Correspondence: Dr. Maria M D’souza, Division of PET Imaging, INMAS, Brig. SK Majumdar Marg, Delhi - 110 054, India.
E-mail: maria.md@rediffmail.com

Abstract

Lymphomas are a heterogeneous group of diseases that arise from the constituent cells of the immune system or from their precursors. 18F-fludeoxyglucose positron emission tomography/computed tomography (18F-FDG PET/CT) is now the cornerstone of staging procedures in the state-of-the-art management of Hodgkin’s disease and aggressive non-Hodgkin’s lymphoma. It plays an important role in staging, restaging, prognostication, planning appropriate treatment strategies, monitoring therapy, and detecting recurrence. However, its role in indolent lymphomas is still unclear and calls for further investigational trials. The protean PET/CT manifestations of lymphoma necessitate a familiarity with the spectrum of imaging findings to enable accurate diagnosis. A meticulous evaluation of PET/CT findings, an understanding of its role in the management of lymphomas, and knowledge of its limitations are mandatory for the optimal utilization of this technique.

Key words: Imaging; lymphoma; PET/CT

Introduction

Lymphomas are a heterogeneous group of diseases that arise from the constituent cells of the immune system or from their precursors. They are known to arise from virtually any organ or tissue in the body. 18F-fludeoxyglucose positron emission tomography/computed tomography (18F-FDG PET/CT) has become a standard procedure for the evaluation of lymphomas. Their protean imaging manifestations necessitate familiarity with the spectrum of imaging findings to enable accurate diagnosis. An understanding of the role of FDG PET/CT in the management of lymphomas and knowledge of its limitations is mandatory for the optimal utilization of this technique.

Lymphomas are broadly classified into two main groups: Non-Hodgkin’s lymphoma (NHL) and Hodgkin’s lymphoma (HL). NHL is more common and represents nearly 85% of lymphomas.[3] Based on histological characteristics and tumor cell phenotype, HL is subdivided into the classical and non-classical types. The former has four histological subtypes – nodular sclerosis (the most common), lymphocyte predominance (with the most favorable prognosis), mixed cellularity, and lymphocyte depletion, while the latter includes the nodular lymphocyte predominant type.[4] NHL can broadly be classified into two prognostic groups: The aggressive and the indolent lymphomas, the latter group having a better prognosis, with a median survival spanning about 10 years. Of the aggressive lymphomas, diffuse large B-cell lymphoma (DLBCL) is the most common, accounting for about 30% of cases, followed by mantle cell lymphoma (MCL) and adult T-cell leukemia/lymphoma, which represent 6 and 8% of NHL cases, respectively. The most common type of indolent lymphoma is the follicular lymphoma (FL) which represents 22% of NHL cases, followed by marginal zone lymphoma (MZL) and small-cell lymphocytic lymphoma (SLL) representing 6 and 8% of cases, respectively.[3]

PET/CT in Initial Staging

Staging of lymphomas is a vital pre-requisite to their appropriate therapeutic management and prognostication.
This is based on the Ann Arbor system, to which has been added a definition of bulky disease called the Cotswold modification.\[9\] Imaging modalities have a fundamental role in the staging of lymphomas. CT is the most commonly used imaging modality for staging malignant lymphoma because of its widespread availability and relatively low cost.\[6\] However, CT lacks functional information, which impedes identification of disease in normal-sized organs and detection of lesions that have poor contrast with the surrounding tissue.\[7\] Another weakness of CT is that it is not reliable in the detection of bone marrow disease, which, if present, by definition indicates stage IV disease.\[8\] CT also subjects the patient to ionizing radiation, which may induce second cancers.\[7\] Each CT scan, covering the neck, thorax, abdomen, and pelvis, is associated with an effective dose of approximately 20-25 mSv.\[7\] This is different from that of a routine diagnostic whole-body CECT. The effective dose in a stand-alone whole-body PET study is approximately 3.3-7.6 mSv per examination.\[9\] The greatest contributor to the radiation burden incurred is thus from CT, which has contributed to doses ranging from 2.7 to 54.2 mSv, depending on individual protocols.\[11\] The ALARA (as low as reasonably achievable) principle must be borne in mind while performing a PET/CT study, to reduce the radiation dose, without sacrificing diagnostic information.

The usefulness of 18F-FDG PET in the initial staging of lymphoma has been demonstrated in several studies. The main advantage over anatomical imaging techniques, such as CT, is its ability to detect metabolic changes in the areas involved with malignant lymphoma before the structural changes become visible. It can detect more lesions than CT [Figure 1] and may lead to a change in the stage of up to 8-20% of patients.\[12,13\] PET may upstage or pick up occult lesions such as splenic [Figure 2], bone marrow [Figure 3], osseous, and with gastrointestinal involvement [Figure 4] which may otherwise be missed on conventional CT.\[14\] A high level of concordance has been observed between the sites of focal FDG uptake in the bone marrow and bone marrow biopsy. In fact, PET scans have demonstrated a high negative predictive value to exclude bone marrow involvement. This holds particularly true for early-stage HL, and may even obviate the need for bone marrow biopsy in this group.\[14\] Another major advantage of FDG-PET/CT is that since it is a whole-body imaging method, it can

---

**Figure 1 (A-G):** Maximal intensity projection (MIP) image (A) in a patient of diffuse large B-cell lymphoma (DLBCL) shows extensive sites of involvement visualized as areas of increased FDG uptake. Transaxial CT images (B, C) show axillary and abdominal lymphadenopathy (thin arrows) and a large subcutaneous nodule (arrowhead). Fused PET/CT images (axial D, E; coronal F; and sagittal G) show sites of bone marrow involvement (thick arrows) in the sternum, spine, and iliac crest, over and above the lesions picked up on CT.

**Figure 2 (A, B):** A case of HL (mixed cellularity type): Spleen appears unremarkable on transaxial CT image (arrow in A). PET/CT fusion image at corresponding site shows diffusely increased FDG uptake (arrow in B).
guide the biopsy from a metabolically active and easily accessible site.

The ability of PET to differentiate indolent from aggressive lymphomas has been the subject of many studies. Routinely, HL and aggressive NHL like DLBCL [Figure 5A, B] and grade III FLs are FDG avid. However, some subtypes of NHL, predominantly the indolent lymphomas such as MZLs [Figures 5C, D, and 6] and peripheral T-cell lymphomas may have low or even no uptake of FDG. Albeit PET/CT provides significantly more accurate information compared to PET and CT for the staging and re-staging of patients with indolent lymphoma, caution is warranted in these histologic subtypes of NHL because a negative FDG PET scan does not necessarily rule out disease; complimentary anatomical imaging [CECT or magnetic resonance imaging (MRI)] is mandatory to increase the detection rate of the lesions. PET/CT also has the potential to detect transformation of a low-grade lymphoma to a more aggressive subtype (Richter transformation) based on the degree of FDG avidity.

The degree of FDG uptake can be expressed quantitatively by means of the Standardized Uptake Value (SUV). It represents the activity in the lesion in μCi/ml corrected for the weight of the patient and the dose of FDG administered. A study by Ngeow et al. showed that an FDG uptake of more than 10 SUV is predictive of an aggressive B-cell lineage or suggestive of the presence of a more aggressive histological component. Another study by Schoder et al. showed that an FDG uptake of more than 10 SUV is predictive of an aggressive B-cell lineage or suggestive of the presence of a more aggressive histological component.

**Figure 3 (A, B):** Transaxial CT image at the level of upper thigh (A) In a case of NHL (DLBCL) appears unremarkable. PET/CT fusion image at the corresponding site (B) intense FDG uptake in the bone marrow of proximal right femoral shaft (arrow) suggestive of marrow involvement.

**Figure 4 (A, B):** Transaxial PET/CT fusion image (B) In a case of DLBCL shows intense FDG uptake in the small bowel loop (arrow). This was picked up only on retrospect (arrow) in the corresponding CT image (A).

**Figure 5 (A-D):** Transaxial CT (A) and PET/CT (B) images in a case of DLBCL showing FDG-avid retroperitoneal lymphadenopathy (thin arrows). Retroperitoneal lymph node involvement (thick arrows) is also visualized on CT (C) in a case of MZL, which shows very poor FDG avidity on the PET/CT fusion image (D).

**Figure 6 (A-C):** A case of low-grade FL: MIP image (A) appears unremarkable. Transaxial CT image (B) an enlarged pelvic lymph node mass (arrow), with poor FDG avidity on the fused PET/CT image (C).
showed that all indolent lymphomas had an SUV of less than 13, while 35% of aggressive lymphomas also had an SUV of less than 13. Despite the overlap between the two, the study concluded that aggressive disease had a higher $^{18}$F-FDG uptake than did indolent lymphomas ($P < 0.01$) and the authors suggested that an SUV of more than 10 confers a higher likelihood for aggressive disease. This has important diagnostic and therapeutic implications. PET/CT plays an important role in arriving at an accurate diagnosis. It can be used to detect sites that may be more accessible for biopsy and guide biopsies to the site of highest FDG uptake, representing the most aggressive site of lymphoma.\textsuperscript{[20]} It may thus play an important role in identifying the foci of aggressive transformation or aggressive histology in those patients who were thought to harbor an indolent lymphoma on previous biopsy.\textsuperscript{[14]} Owing to the differential pattern of FDG avidity, PET can potentially detect two separate clones at the same time. Biopsy of the lesions with intense uptake will help identify a separate clone (if only lesions with moderate uptake have been sampled before). This additional information, which cannot be obtained on conventional morphological imaging, has utmost clinical relevance, as these patients require more aggressive therapy.

PET/CT in Evaluation of Treatment Response

Evaluating therapeutic response is vital to the management of lymphoma patients. This information is a must to tailor the therapy in each patient depending upon the individualized response. With a larger and more effective arsenal of therapeutic regimens currently available, it is important to optimize treatment strategies to achieve maximal disease control with minimal toxicity to the patient. Therapeutic response is assessed based on clinical, histopathologic, and imaging criteria. CT has been hitherto the most commonly used imaging modality. A decrease in size of the tumor mass has been the cornerstone in establishing a good therapeutic response. However, a residual mass picked up on CT may not necessarily be metabolically active. The inclusion of PET/CT has addressed some of the limiting factors of CT, which include: the size criteria for lymph node involvement, the differentiation of unopacified bowel from lesions in the abdomen and pelvis, the inability to distinguish viable tumor from necrotic/fibrotic lesions after therapy, and the characterization of small lesions.\textsuperscript{[21]}

$^{18}$F-FDG PET has been shown to be able to distinguish between post-treatment fibrosis and viable tumor [Figure 7]. In this regard, $^{18}$F-FDG PET has a higher specificity (92% vs. 17%, $P < 0.01$), accuracy (96% vs. 63%, $P < 0.05$), and positive predictive value (94% vs. 60%, $P < 0.05$) than does CT.\textsuperscript{[22]} In a recently published retrospective analysis of 75 patients with HD or aggressive NHL treated with standard chemotherapy regimens with or without radiation therapy whose disease was restaged with PET and CT, a correlation was found between positive findings on restaging PET and clinical relapse.\textsuperscript{[23]} Studies have also documented the utility of PET to assess the response to treatment with radiolabeled antibodies.\textsuperscript{[24]}

PET/CT studies have been found useful to assess the therapeutic response during the course of

![Figure 7 (A-F): A case of mesenteric lymphoma: Top panel - CT images, bottom panel - PET/CT images. Initial scan (Feb 09) (A, B) Before the onset of therapy shows an FDG-avid mesenteric lymph node mass (arrows). The mass has reduced in size, but still persists on CT (C) in the post-therapy scan (Jan 10). Corresponding PET/CT image (D) No uptake indicating complete metabolic resolution. A follow-up scan (July 10) in the absence of therapy shows further reduction in the size of the mass on CT (E), with persistently absent FDG uptake on the PET/CT fusion image (F)](image-url)
chemotherapy [Figure 8]. This is interim PET evaluation or early PET study. Conventional imaging methods are inadequate in this realm as tumor shrinkage takes time, and cannot be used as the basis for early monitoring of response to treatment.

Most responders will become PET negative after two to three cycles of standard chemotherapy. A study of 121 patients with NHL, which assessed the utility of 18F-FDG PET after 2-3 cycles of chemotherapy found that 18F-FDG PET had a high predictive value for progression-free survival and overall survival.[25] Another study evaluated 18F-FDG uptake on PET before and after 1 cycle of chemotherapy in 30 patients with NHL or HD. Negative PET findings in that study were highly predictive of remission, with 87% of the patients in complete remission after a median of 19 months of follow-up.[26] PET/CT has outperformed CT in the evaluation of therapeutic response as early as after 1-2 cycles of chemotherapy.[27] The interim scan, however, may show evidence of partial or incomplete response. One study performed on HL and NHL patients used visual and quantitative analysis of FDG uptake performed before and after 1-2 cycles of chemotherapy.[28] At the end of a 2-year follow-up, they concluded that a 60% reduction in SUV in the interim scan is a reliable criterion to distinguish the responders from the non-responders. A negative scan is thus not mandatory to identify a responder. The Deauville criteria, laid by a consensus committee of nuclear medicine physicians, hematologists, and oncologists recommends a 5-point scale rather than taking a binary decision (namely PET positive or negative). This is based upon visual analysis, with background uptake as reference. International validation studies are on to assess the utility of these criteria.[29] Recognition of early therapeutic response or failure enables the treatment to be adjusted accordingly. Several ongoing trials are under way to assess the impact of treatment adaptation based on early 18F-FDG PET results on overall survival.[30]

With the widespread use of PET, the need to revise the existing guidelines for lymphoma staging was felt.[31] The consensus recommendations regarding the use of FDG-PET for assessment response published by the International Harmonization Project in 2007 state that in routinely FDG-avid lymphomas, such as DLBCL and HL, a Complete Response (CR) is assigned to all patients with a negative PET scan regardless of the presence of a residual mass on CT. In cases in which there is residual disease on PET (PET-positive patients), a partial response, stable disease, or progressive disease can be assigned based on the response shown by CT; the Unconfirmed Complete Response (Cru) category has been eliminated.[32]

Spectrum of PET/CT Findings in Lymphoma

As the name suggests, lymph nodes are the most commonly affected sites in lymphoma – be it cervical, supraclavicular, axillary, mediastinal, abdominal, or inguinal [Figure 9]. Extranodal lymphoma occurs in about 40% of patients with lymphoma.[33] It is observed more frequently in NHL than with HL, and is often intermediate to high grade.[34] The incidence of extranodal lymphoma is higher in patients with AIDS and immunodeficient states.[35,36] Lymphomatous nodes appear discretely enlarged or as soft-tissue masses, with variable FDG avidity depending on the histological type.

Extranodal Abdominal Manifestations

The most commonly involved extranodal abdominal site is the spleen, followed by the liver, gastrointestinal tract, pancreas, abdominal wall, genitourinary tract, adrenal, peritoneal cavity, and biliary tract in that order.[37] Both hepatic and splenic involvement may
occur in the form of diffuse infiltration with or without organomegaly [Figure 10] and as focal nodules. PET/CT outperforms conventional imaging in the evaluation of hepatosplenic involvement. In the gastrointestinal tract, the stomach, small bowel, pharynx, large bowel, and esophagus are involved in decreasing order of frequency.[38] The pattern of involvement may be diffusely infiltrative, polypoidal, ulcerative, or nodular [Figure 11]. The interpretation of positive PET/CT findings in the gastrointestinal tract may be challenging owing to physiological FDG uptake in the large bowel and to a lesser extent in the stomach and small bowel.[39] Diffuse uptake has most often been observed to be associated with normal colonoscopy, whereas focal uptake is often pathological.[40] Simple maneuvers like delayed imaging, negative oral contrast like water, or diluted positive oral contrast agents[41] often help to resolve the issue. Subtle lymphomatous bowel deposits that may be missed on CT are well visualized on PET/CT in aggressive NHL and HL. Pancreatic involvement may be diffuse (resembling pancreatitis) in the form of focal masses or as secondary involvement from adjacent lymph nodes. Renal involvement may be in the form of solitary or multiple masses, diffuse infiltration, and spread from adjacent nodes. Peritoneal lymphomatosis is rare and closely mimics carcinomatosis.[42] It presents as peritoneal nodules, infiltrative masses, and ascites [Figure 12]. PET/CT is highly sensitive in the detection of peritoneal involvement in FDG-avid lymphomas.

**Intrathoracic Manifestations**

Intrathoracic disease is noted in 40-50% of patients with NHL at presentation, compared with 85% of those with HL.[43] Pulmonary involvement presents as solitary or multiple nodules [Figure 13] or masses, reticulonodular opacities, and consolidation.[44] Accurate characterization of pulmonary lesions on PET/CT requires diagnostic quality breath-hold CT, preferably high-resolution CT (HRCT). The FDG avidity is variable, depending on the histological type. Mediastinal lymphadenopathy can vary in extent and in size and may manifest as a large solitary mass or as discrete nodes within masses of matted nodes. HL most commonly affects the anterior mediastinal group, while NHL involvement is more diffuse. Interpretation of the PET/CT scan needs to be performed with caution, as FDG uptake is seen in inflammatory lymphadenopathy as well.[45]

Secondary pleural, pericardial [Figure 14], chest wall, and diaphragmatic involvement is also known. Primary cardiac lymphomas are very rare.

**CNS and Head and Neck Lymphomas**

Intracranial lymphoma [Figure 15] which is seen almost exclusively with NHLs manifests as discrete masses or infiltrations involving the corpus callosum, deep gray matter structures, and periventricular regions.[46] Meningeal infiltration may be seen in the form of subarachnoid nodules, diffuse leptomeningeal carcinomatosis, or dural masses. FDG PET has been found useful in the management of AIDS patients with CNS lesions. It is often not possible for CNS lymphomas to be distinguished from toxoplasmosis in the immunocompromised patient on the basis of conventional morphological imaging due to the similarity in appearance. However, CNS lymphomas show high FDG uptake, unlike toxoplasmosis, making it possible to distinguish the two.[47]
This can potentially avoid invasive diagnostic procedures and enable the early initiation of therapy.

Lymphomatous masses of the head and neck may involve the Waldeyer’s ring, including nasopharynx [Figure 16], base of tongue, tonsils [Figure 17], and soft palate. Lymphomatous infiltration of the paranasal sinuses [Figure 18] appears as a nonspecific soft-tissue mass partially or completely obscuring the involved lumen. Destruction of the surrounding bone may be associated. Jaw involvement, which usually occurs in Burkitt’s lymphoma, presents as an FDG-avid lytic lesion with associated soft-tissue component.[1]

**Musculoskeletal Manifestations**

Lymphomas may involve the bone marrow or cortical bone. Several studies have shown that $^{18}$F-FDG PET is accurate in evaluating the presence of bone marrow disease and scores over CT in this regard. However, diffusely increased $^{18}$F-FDG uptake in the bone marrow may be seen owing to marrow hyperplasia post-chemotherapy or due to granulocyte colony-stimulating factors, and thus hamper identification of actual marrow involvement. This increased uptake generally returns to baseline levels by 1 month.[12]
Primary lymphomas frequently involve the appendicular skeleton, whereas secondary lymphomas more frequently affect the spine. They are present as FDG-avid lytic or mixed lytic–sclerotic lesions.

PET/CT is also highly sensitive in picking up lymphomatous deposits in the subcutaneous tissue and in the underlying muscle [Figure 19]. They are visualized as hypermetabolic nodules against the background of normal soft-tissue structures.

Cutaneous Lymphoma

Cutaneous T-cell lymphoma accounts for approximately 6% of all cases of NHL and includes two clinical entities with similar histopathologic features: Mycosis fungoides (MF) and Sézary syndrome (SS). For staging MF [Figure 20] and SS, PET/CT has been found to be more sensitive than CT alone in detecting lymph node involvement, and may thus provide more accurate staging and prognostic information.
Pitfalls of PET/CT

Benign conditions with increased glycolysis, such as infection, inflammation, and granulomatous disease, may also lead to increased FDG uptake,\cite{51} and be mistaken for lymphomatous deposits [Figures 21, 22]. Inflammatory changes secondary to surgery, radiotherapy, and chemotherapy may also lead to false-positive results, if the study is performed soon after these interventions.\cite{52} Further, systemic conditions that lead to generalized lymphadenopathy may mimic malignant lymphoma.\cite{53} Renal involvement may be difficult to assess owing to physiological excretion of the tracer. Additionally, sites of high physiological uptake such as the brain, myocardium, gastrointestinal tract, urinary tract, lymphoid tissue, brown adipose tissue, salivary glands, and thymus may obscure or mimic the presence of tumor deposits. However, the use of CT co-registration largely circumvents this problem [Figure 23]. Bone marrow hyperplasia secondary to chemotherapy with or without cytokines, such as granulocyte colony-stimulating factor, may lead to increased FDG uptake up to 3 weeks after the last dose of cytokines. Similar effects can be seen in the spleen.\cite{54} Hence, attention to the timing of the scan is crucial. FDG PET/CT may be falsely negative in certain histological subtypes of lymphomas.
Figure 21 (A-D): A case of systemic sarcoidosis: MIP image (A) Mediastinal and hilar lymphadenopathy with pulmonary involvement (thick arrow) and abdominal (thin arrow), supraclavicular and inguinal lymphadenopathy (arrowheads). PET/CT images (B, C) Bilateral hilar lymphadenopathy (thick arrow) with reticulonodular lesions along thickened nodular bronchovascular bundles (dotted arrow) and retroperitoneal lymphadenopathy (thin arrow). A provisional diagnosis of lymphoma was made based on the above findings. Biopsy proved it to be a case of sarcoidosis. Note the visualization of better detail on the HRCT image (E) compared to the routine CT image (D).

Figure 22 (A-C): A case of disseminated Koch’s masquerading as lymphoma: MIP image (A) Multiple sites of lymph node involvement (arrows). Transaxial PET/CT fusion images (B, C) sites of increased FDG uptake in the enlarged mediastinal and hilar lymph nodes.

Figure 23 (A-D): A follow-up case of HL (nodular sclerosis type) post-therapy: Multiple sites of FDG uptake (thin arrow) noted on the MIP image (A) in this patient on remission. These FDG-avid sites are well seen on the PET image (B). However, CT image (C) is unremarkable. Fused PET/CT image in inverted gray scale (D) localizes these sites of uptake to brown adipose tissue (arrow).

NHL such as MZLs, peripheral T-cell lymphomas, small lymphocytic lymphomas, and primary FLs. In these cases, conventional morphological imaging techniques such as CT or MRI are a must. Bowel involvement may be negative on FDG PET/CT, especially in low-grade lymphomas. In certain cases such as the marginal zone mucosa-associated lymphoid tissue (MALT) lymphoma, the lesions may evade...
visualization both on FG PET and on CT and be picked up on endoscopy or endoscopic ultrasonography. Another disadvantage of FDG PET/CT is exposure of the patient to ionizing radiation; the effective dose is variable depending upon individual protocols. A meticulous evaluation of PET/CT findings, along with a detailed history, clinical examination, and knowledge of the histological type is a pre-requisite to accurate interpretation.

**Conclusion**

$^{18}$F-FDG PET/CT is now the cornerstone of staging procedures in the state-of-the-art management of HL and aggressive NHL. It plays an important role in staging, restaging, prognostication, planning appropriate treatment strategies, monitoring therapy, and detecting recurrence. The role of $^{18}$F-FDG PET/CT in indolent lymphomas is still unclear and calls for further investigational trials.

**References**


29. D’souza, et al.: FDG-PET/CT in lymphoma

Source of Support: Nil, Conflict of Interest: None declared.