Role of $^{18}$F-FDG PET/CT in Erdheim–Chester Disease in the Era of Multimodality Imaging

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Introduction

Erdheim–Chester disease (ECD) is a rare disease with systemic non-Langerhans cell histiocytosis, the diagnosis of which with conventional imaging modalities is challenging. We describe a case of a 73-year-old woman who was referred with a progressive history of bilateral proptosis. The magnetic resonance imaging (MRI) orbit demonstrated bilateral orbital masses with optic nerve encasement. A subsequent 18F-FDG PET/CT scan showed multi-organ disease with involvement of the orbits, pericardium, aorta, pararenal fascia, and appendicular bones. Metabolically active, easily accessible areas were selected for CT-guided biopsy. The biopsy showed sheets of foamy histiocytes with the expression of CD 68 and CD 163 consistent with a diagnosis of Erdheim–Chester disease. The FDG PET/CT played a pivotal role in establishing the diagnosis with the assessment of disease extent and further guided in the targeted biopsy.

Case History

A 73-year-old woman presented with a history of gradually progressive right proptosis followed by the left with no diminution of vision.

MRI orbit demonstrated bilateral intra and extraconal soft tissue masses causing exophthalmos with optic nerve encasement (►Figs. 1 and 2A–C). Magnetic resonance imaging (MRI) of the brain showed no abnormality. Biopsy of the orbital mass was suggestive of necrobiotic xanthogranuloma. Immunohistochemistry (IHC), however, was not available. She was suspected to have ECD due to retro-orbital involvement. A $^{99m}$Tc-methylene diphosphonate bone scan was performed that revealed a characteristic increase in the

Keywords

► Erdheim–Chester disease
► FDG PET/CT
► multimodality imaging

Abstract

Erdheim–Chester disease is a rare disease with systemic non-Langerhans cell histiocytosis, the diagnosis of which with conventional imaging modalities is challenging. We describe a case of a 73-year-old woman who was referred with a progressive history of bilateral proptosis. The magnetic resonance imaging (MRI) orbit demonstrated bilateral orbital masses with optic nerve encasement. A subsequent 18F-FDG PET/CT scan showed multi-organ disease with involvement of the orbits, pericardium, aorta, pararenal fascia, and appendicular bones. Metabolically active, easily accessible areas were selected for CT-guided biopsy. The biopsy showed sheets of foamy histiocytes with the expression of CD 68 and CD 163 consistent with a diagnosis of Erdheim–Chester disease. The FDG PET/CT played a pivotal role in establishing the diagnosis with the assessment of disease extent and further guided in the targeted biopsy.

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osteoblastic activity in the diaphyses of the long bones of the lower limb centered around the knees (Fig. 3A and 3B) along with mandibular uptake. She received interferon-α 2B and intravenous methylprednisolone. However, her symptoms worsened with sudden diminution of vision and development of bone pains and hence she was referred to our hospital for further evaluation.

An FDG-PET/CT scan was requested to assess for the extraosseous disease.

The PET/CT findings supported the diagnoses of ECD and confirmed multi-organ involvement. Metabolically active, easily accessible areas in the right pararenal region were selected for CT-guided biopsy. A second biopsy was

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**Fig. 1** T2-weighted axial MRI (A) showing intracranial hypointense masses encasing the extraocular muscles (white arrow). T1-weighted axial image (B) demonstrating the isointense nature of orbital masses (white arrow). Post-contrast T1-weighted axial image (C) showing intense enhancement of the intracranial masses (white arrow) with encasement of the optic nerves (yellow arrow). Non-contrast CT (D) demonstrating hypodense masses involving the intra- and extracranial compartments with increased FDG uptake seen in corresponding regions on PET (E) and fusion PET/CT images (F). FDG, fluorodeoxyglucose; PET/CT, positron emission tomography-computed tomography; MRI, magnetic resonance imaging.

**Fig. 2** T2-weighted coronal MRI (A) showing hypointense intracranial masses (white arrows) encasing the optic nerve (yellow arrows). T1-weighted coronal MRI (B) demonstrating orbital masses, which are isointense to the muscle (white arrow). Post-contrast T1-weighted coronal MRI (C) showing avid enhancement of the intracranial masses (white arrow) with encasement of the optic nerves (yellow arrow). T1-weighted sagittal image (D) demonstrating normal hyperintense signal of the neurohypophysis (thin white arrow). MRI, magnetic resonance imaging.
warranted to assess IHC and mutational status, particularly with the worsening of symptoms as targeted therapy was being considered. The biopsy showed sheets of foamy histiocytes (►Fig. 6D), consistent with ECD. Immunohistochemistry showed the expression of CD 68 and CD 163 and was negative for S-100, CD1a, and Langerin. BRAF mutation was negative.

Owing to a negative BRAF mutation status, targeted therapy was not feasible. She was treated with high-dose dexamethasone, cyclophosphamide, and vincristine to which there was an initial response but the patient subsequently succumbed to the cardiac complications of ECD.

Discussion

ECD is a subset of non-Langerhans cell histiocytosis, first described by William Chester and Jakob Erdheim in 1930.6 The diagnosis is often delayed due to the rarity of the disease, multi-organ involvement, and varied presentation.7 Imaging including 18F-FDG PET/CT plays an important role in the diagnosis, assessing the extent and evaluating the treatment response.2–5,7

Up to 96% of patients show skeletal involvement,8,9 like our patient. The 99mTc-methylene diphosphonate bone scan and the FDG-PET showed intense, bilaterally symmetric uptake at the end of long bones centered around the knees—“hot-knee pattern” (►Fig. 3), which is characteristic of ECD.7,10 The CT demonstrated corresponding symmetrical osteosclerosis of the diaphyses of long bones with characteristic sparing of the epiphysis (white arrow). MIP, maximum intensity projection; MRI, magnetic resonance imaging; PET/CT, positron emission tomography-computed tomography.

Orbital involvement is present in up to 38%.8,9,13 Exophthalmos was the initial presentation of our patient. The
initial biopsy suggested necrobiotic xanthogranuloma, which commonly presents as anterior orbital or ocular adnexal lesions. The retro-orbital involvement led to the suspicion of ECD. The MRI revealed classic T1 and T2 hypo-intense multi-compartmental retro-orbital masses encasing the optic nerve (Figs. 1 and 2). PET/CT showed corresponding increased metabolic activity (Fig. 1) with SUV$_{max}$ of 13 similar to other studies.

Imaging plays a vital role in surgical planning, particularly in the orbit. The exquisite soft-tissue contrast of MRI helps planning for surgical decompression, particularly if the optic nerve is at risk. Imaging including PET/CT, which identifies active tissue, helps the ophthalmologist choose the best site for biopsy.

CNS involvement is seen in more than half of cases and is a leading cause of death. The absence of a normal T1 hyper-intense signal of the neurohypophysis, dural, and meningeal involvement and symmetrical T2-hyperintensities of the dentate nucleus have been described.

Cardiovascular involvement is seen in nearly three-fourths of cases and presents either as circumferential infiltration of the adventitia of the blood vessels, commonly involving the aortic arch or as pericardial infiltration or rarely as right-sided myocardial involvement. PET/CT in our patient revealed hypermetabolic hypo-dense, plaque-like soft tissue circumferentially encasing the aorta and its branches (Fig. 4) — described as the "coated aorta sign." It also showed smooth plaque-like thickening.

**Fig. 5** (A–F, clockwise) CECT (A) and PET/CT (B) showing smooth plaque-like hypermetabolic thickening mainly involving the right side of the heart (thin white arrows), pericardial thickening with effusion (bold white arrow), and bilateral pleural effusion (arrowheads). Coronal reformatted CECT (C) and PET/CT (D) images showing plaque-like hypermetabolic soft tissue in relation to the right atrium (white arrows). Axial (E) and coronal images (F) of the CT chest (lung window) demonstrating septal thickening in the lung bases (bold yellow arrows). CECT, contrast-enhanced computed tomography; PET/CT, positron emission tomography-computed tomography.

**Fig. 6** Hairy kidney sign—CECT (A) demonstrating perirenal fat infiltration with a characteristic spiculated appearance (thin white arrow) and enhancing pararenal soft tissue masses (bold white arrow). PET/CT (B) demonstrating the hypermetabolic nature of perirenal and pararenal soft tissue thickening (thin white arrows and bold white arrows, respectively). Coronal reformatted MIP image of CECT abdomen (C) showing circumferential soft tissue encircling the mesenteric vessels; which is also seen in images A and B (yellow arrows). Histology (D) showing sheets of foamy histiocytes. CECT, contrast-enhanced computed tomography; MIP, maximum intensity projection.
involving the right side of the heart, together with pericardial thickening and effusion (►Fig. 5A–D).

Cardiovascular involvement indicates a poor prognosis and leads to death in ~60% of cases due to tamponade, arrhythmias, or infarction.8,18 Our patient expired due to cardiovascular complications of ECD.

The lung and pleura are affected in ~40 to 50% of cases8 as smooth inter- and intralobular septal thickening19 as was demonstrated in our patient (►Fig. 5E and 5F).

The retroperitoneum including the kidneys is involved in ~68% of cases.4,8 Our patient had FDG avid soft tissue masses in bilateral perirenal and posterior pararenal spaces (►Fig. 6A and 6B). The hairy kidney sign refers to perirenal fat infiltration seen in ECD with a characteristic spiculated appearance,4,7,8 which was seen quite elegantly on the PET/CT (►Fig. 6A and 6B) with corresponding FDG uptake.

18F-FDG PET/CT assists in differentiating ECD from Langerhans histiocytosis (LCH), Rosai–Dorfman Disease (RDD), IgG4-related disease, idiopathic retroperitoneal fibrosis, and sarcoidosis.8

Generally, LCH manifests earlier than ECD. The bony involvement in LCH is characteristically osteolytic and involves the skull, spine, or long bones asymmetrically unlike the symmetrical osteosclerosis seen in ECD, which involves the ends of the long bones. Cardiac involvement is rare in LCH.20

RDD may show right heart involvement similar to ECD but bony involvement is characteristically osteolytic.20 Lymphadenopathy especially in more than one region with moderate to high 18F-FDG avidity is seen in RDD.

Specific FDG-PET/CT patterns have been described in IgG4-RD—diffusely increased 18F-FDG uptake in the pancreas and salivary glands, patchy lesions in the retroperitoneal region and vascular wall, and multi-organ involvement.21

Retroperitoneal fibrosis (RPF), retro-orbital involvement, and pachymeningitis are common features between ECD and IgG4-RD. RPF associated with ECD generally affects the perirenal areas with resultant proximal ureteric obstruction. RPF associated with IgG4-related disease usually results in distal ureteric obstruction with medial deviantiation.22

Idiopathic RPF shows lower FDG uptake as compared with secondary RPF as seen in ECD.23

Sarcoidosis shows similarity with ECD as far as the CNS and pulmonary findings are considered but the characteristic bony or vascular changes have not been described.8 Sarcoidosis, in turn, is characterized by metabolically active lymphadenopathy.

Although the bone scan is a sensitive tool to assess skeletal involvement in ECD,10 evaluation of the bone marrow is not possible. PET/CT not only evaluates skeletal involvement but also occult visceral, vascular, and bone marrow involvement. FDG-PET hence helps choose the best site for biopsy.

The Annual ECD Medical Symposium Consensus 2020 guidelines recommend multicore biopsies from the most FDG-avid sites that are accessible and safe as histiocytic infiltration may be heterogeneous.24 The right posterior pararenal FDG avid soft tissue masses, which were easily and safely accessible, were chosen for repeat biopsy in our patient.

These guidelines also recommend BRAF mutational status to initiate BRAF V 600E inhibitor therapy.24 As the BRAF mutational status in our patient was negative, targeted therapy was not feasible.

If targeted therapies are unavailable or not indicated, treatment depends on the active disease burden, wherein PET-CT plays an important role.24

Tissue fibrosis can occur in ECD and hence an evaluation of metabolic rather than anatomic treatment response is essential with a follow-up PET-CT 3 to 6 months after the initiation of therapy.24

Hence, 18F-FDG-PET/CT plays an important role in patients with ECD not only in the diagnoses but also in the evaluation of disease extent as a guiding tool for targeted biopsies and assessment for treatment response.

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Conflicts of Interest
None declared.

References