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To cite this version:
D. Taïeb, D. Rubello, A. Al-Nahhas, M. Calzada, M.C. Marzola, et al.. Modern PET imaging for paragangliomas: Relation to genetic mutations. EJSO - European Journal of Surgical Oncology, WB Saunders, 2011, 37 (8), pp.662. <10.1016/j.ejso.2011.05.004>. <hal-00715503>

HAL Id: hal-00715503
https://hal.archives-ouvertes.fr/hal-00715503
Submitted on 8 Jul 2012

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Modern PET imaging for paragangliomas: relation to genetic mutations

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Running head: New tracers for paragangliomas

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Conflict of interest statement: the authors declare no conflict of interest.
Abstract

Aim and Methods: Paragangliomas (PGL) are neural crest-derived tumours that are found along the autonomic neural network throughout the body and can be multiple and/or metastatic. Nuclear medicine imaging in combination with conventional imaging is required to fully delineate the extent of the disease. The performance of molecular imaging modalities is widely dependent on tumour biology.

Results: In the present paper we discuss the recent publications focused on the role of positron emission tomography (PET) imaging and the relationship between tracer uptake patterns and genetic mutations associated with the disease.

Conclusion: Recent advances in genetic and molecular pathogenesis of PGL have allowed for the identification of new molecular diagnostic and therapeutic radiopharmaceuticals tailored to genetic abnormalities. However, the optimal diagnostic imaging algorithm remains to be determined.

Keywords: Paraganglioma; Genetic mutations; Positron-emission tomography; radiopharmaceuticals
**Introduction**

Paragangliomas (PGL) are part of a family of highly vascularized neural crest-derived neoplasms originating in the adrenal medulla, chemoreceptors or autonomic ganglia. PGL are found throughout the body in association with either the sympathetic or parasympathetic systems. PGL involving the adrenal glands are called phaeochromocytomas. Extra-adrenal PGL can be associated with the parasympathetic system and found in the head, neck and mediastinum. Those associated with the sympathetic system predominate in the retroperitoneum along the thoracolumbar para-aortic region. In contrast to other endocrine tumours, a high proportion (>20%) of patients carry germline mutations that predispose to multifocality.

Six major genes (SDHB, SDHC, SDHD, VHL, RET, NF1) are involved in tumorigenesis. SDHx and VHL gene mutations predispose to adrenal and extra-adrenal PGL whereas RET and NF1 mutations are associated with adrenal PGL (often bilateral). Head and neck (H&N) PGL and tendency for multifocal involvement are more frequently associated with SDHD compared to SDHB; whereas malignancy is strongly associated with SDHB mutations (37.5% vs. 3.1% for SDHD).\(^{1,2}\) Current knowledge of the disorders associated with SDHC mutations is still limited but typically PGL originate from the parasympathetic ganglia.

**Challenges of modern imaging**

It is expected that the early detection of PGL may minimize complications related to mass effect and hormonal excess, facilitate curative treatment and potentially reduce the metastatic spread. One important but unresolved question is whether early detection of very small and biochemically silent PGL in family members would improve the prognosis of patients.

*Functional imaging*
Functional imaging is probably not necessary for most adrenal PGL (sporadic, <5 cm). By contrast, in malignant and multifocal forms (often inherited), the role of pre therapy imaging is crucial for providing accurate staging of the disease. In this respect, nuclear imaging in combination with conventional imaging may be required to fully delineate the extent of the disease. Radiolabelled metaiodobenzylguanidine (MIBG) and somatostatin analogues are the traditional imaging procedures. \[^{[13]I/^{[123]}I}\]MIBG scintigraphy is the most widely used tracer for detecting abdominal PGL, while \[^{[111]}In\]pentetreotide-SPECT can be more sensitive in H\&N PGL. They represent the gold standard functional imaging modalities for sporadic PGL. However, \[^{[13]I/^{[123]}I}\]MIBG and \[^{[111]}In\]pentetreotide scintigraphy demonstrate a suboptimal sensitivity due to the detection limits of conventional gamma camera imaging, although the contribution of SPECT/CT has led to significant improvement in the diagnostic confidence in image interpretation. Therefore, significant underestimation of the disease may occur in patients at risk of multifocality and/or malignancy. More recently, the use of positron emission tomography (PET) imaging in this group of tumours is growing rapidly, paralleled by great efforts towards the development of new tracers.

**PET tracers**

PGL are characterized by their high density of peptide receptors, their ability to take up amino acids and decarboxylate them into biogenic amines and their high glucose uptake. These features are used for nuclear imaging targeting by using specific and non-specific PET tracers.

Figure 1 shows the mechanism of uptake into PGL cells for four different PET tracers.

\[^{[18]}F\]Fluorodeoxyglucose

\[^{[18]}F\]Fluorodeoxyglucose (\[^{[18]}F\]FDG) is taken up via glucose transporters and phosphorylated by hexokinase into \[^{[18]}F\]FDG-6P. However, increased \[^{18}\]F\]FDG uptake is not
limited to malignancy and many oncogenetic signals are also present in some benign PGL that may result in increased glucose uptake (i.e., mutations in VHL and SDHx genes).\(^5\)

\[^{18}F\text{-dihydroxyphenylalanine}\]

Dihydroxyphenylalanine (DOPA) is a precursor to Dopamine and, \(^{[18}F\)\-dihydroxyphenylalanine (\[^{18}F\)FDOPA) shows characteristically high uptake in PGL cells, which are offspring from the amine precursor uptake and decarboxylation (APUD) stem cells. \(^{[18}F\)FDOPA is transported into cells via sodium-independent large neutral amino acids transporter type 1 (LAT1), and decarboxylated into Fluorodopamine by the enzyme aromatic L-amino acid decarboxylase (AADC) and stored in cytoplasmic neurosecretory granules via vesicular monoamine transporters (VMAT1/2).

\[^{18}F\text{-Fluorodopamine}\]

By contrast, another tracer, \(^{[18}F\)Fluorodopamine (\[^{18}F\)FDA is taken up by to the cytosol via the same cell membrane transporters that take up MIBG (norepinephrine transporter, NET) and stored into vesicles. Dopamine type 2 receptor (D2DR) is also expressed in PGL and may influence the patterns of \(^{[18}F\)FDA uptake.\(^6,7,8\)

\[^{68}Ga\text{-labelled somatostatin analogue peptides}\]

\[^{68}Ga\)-PET has emerged as an alternative to cyclotron-based PET radiopharmaceuticals. \[^{68}Ga\]-labelled somatostatin analogue peptides (\[^{68}Ga\)SSTa) directly bind to somatostatin receptors (SSTR) on the tumour cells surface and are internalized within the receptors. PGL are characterized by their high density in SSTRs, especially the subtype 2 (SSTR2).\(^9\) The mechanisms of uptake and retention of several tracers in tumour cells are depicted in Figure 1 and Table 1.
Preliminary data from PET tracers

Different PET tracers have thus been evaluated in clinical studies: $^{[11]C}$hydroxyephedrine (not discussed), $^{[18]F}$FDG, $^{[18]F}$FDOPA, $^{[18]F}$FDA and $^{[68]Ga}$-labelled somatostatin analogs ($^{[68]Ga}$SSTa). Analysis of the relevant literature is hampered by the frequent mixing of both sympathetic and parasympathetic PGL.

$^{[18]F}$FDG-PET

The performance of $^{[18]F}$FDG-PET has been extensively evaluated only recently in PGL because of its lack of specificity and the initial disappointing results. Several studies have found that $^{[18]F}$FDG-PET was sensitive in the evaluation of PGL. It should be noted that the vast majority of patients had abdominal and or malignant forms, making it impossible to fully extrapolate the results for head and neck (H&N) lesions. By contrast to medullary thyroid carcinomas and digestive endocrine tumours which tend to increase their glucose avidity in the later stages of the disease, $^{[18]F}$FDG-PET should be considered as a new molecular signature of PGL that could occur in the benign form of the disease. Patients with catecholamine-secreting PGL may also have brown adipose tissue activation and uptake of $^{[18]F}$FDG. It is now established that tumours that carry SDHB mutations have higher glucose uptake values (an example is shown in Figure 2). It is probable that these mutations could participate in metabolic reprogramming in tumour cells. Since SDH are mitochondrial proteins of the tricarboxylic acid cycle and the respiratory chain, mutations could impair energy production and in turn induce upregulation of genes involved in angiogenesis, glucose uptake and consumption by the tumour. Transcriptional profiling studies reveal pseudohypoxic signatures in SDHx/VHL tumours that differentiate them from RET/NF1 cases. Glucose transporters (GLUT) may play an important role in mediating tumour glucose uptake [17]. However, overexpression of hypoxic-inducible factor (HIF)-1$\alpha$
target genes is mainly found in VHL tumours due to the decrease in VHL-mediated
degradation of HIF-1α.17 Further studies are thus required to understand the molecular basis
of glucose avidity of SDHx tumours.

\[ ^{18}\text{F}]\text{FDA} \]

\[^{18}\text{F}]\text{FDA} \text{ is an experimental radiotracer that has been developed and evaluated at the National Institutes of Health in Bethesda.}^{15,19,20,21,22} \[^{18}\text{F}]\text{FDA-PET was found to be more sensitive (76\%) than }^{[18}\text{F}]\text{FDOPA-PET (45\%) and }^{[123]}\text{I-MIBG (57\%)} \text{ in the detection of metastatic deposits. In non-metastatic PGL, differences between imaging modalities did not achieve statistical significance. In their prospective study, Timmers et al recommend }^{[18}\text{F}]\text{FDOPA-PET in patients with an unknown genotype, }^{[18}\text{F}]\text{FDG and PET in SDHB mutation carriers, and }^{[18}\text{F}]\text{FDOPA or }^{[18}\text{F}]\text{FDA PET in non-SDHB patients.}^{20}

\[ ^{18}\text{F}]\text{Fluoride} \]

Since bone metastases frequently occur in metastatic PGL, \[^{18}\text{F}]\text{Fluoride-PET might emerge as a sensitive diagnostic tool and replace bone scintigraphy in these patients.}^{15,23}

\[ ^{18}\text{F}]\text{FDOPA} \]

Several studies have addressed the issues of \[^{18}\text{F}]\text{FDOPA-PET in non-metastatic and metastatic PGL.}^{13,20,24,25,26,27,28,29,30,31,32,33,34} \text{ To date, }^{[18}\text{F}]\text{FDOPA-PET could be the most sensitive tracer for detecting H&N PGL}^{25,27,30,31,35} \text{ (Figure 3). High tumour avidity and favorable signal to noise ratio in the head and neck allows the detection of very small PGL by }^{[18}\text{F}]\text{FDOPA-PET. However, }^{[18}\text{F}]\text{FDOPA-PET should not be considered simply as an MIBG scan with higher sensitivity but rather a new specific radiotracer with its own advantages and limitations. }^{[18}\text{F}]\text{FDOPA-PET/CT is a promising tool for detecting adrenal PGL with sensitivity ranging from 85 to 100\%.}^{29,32} \text{ The main advantage of the technique over }^{[123]}\text{I-}
MIBG and $^{18}$F-FDOPA is that there is low physiological $^{18}$F-FDOPA uptake in healthy adrenal tissue that facilitates the diagnosis of adrenal PGL in patients at risk of bilateral lesions (i.e., MEN2, NF1). Two studies have demonstrated that $^{18}$F-FDOPA could miss abdominal PGL otherwise detected by MIBG scan and/or $^{18}$F-FDG-PET. Fottner et al demonstrated that tumour expression of vesicular monoamine transporter type 1 (VMAT-1) is critical for detection by MIBG-scintigraphy. This is particularly important in the management of SDHD patients who are susceptible to develop both parasympathetic and sympathetic PGL. Charrier et al recently found that $^{18}$F-FDOPA-PET was more sensitive than $^{111}$In-pentetreotide-SPECT in the diagnosis of H&N PGL (96.7% versus 66.7%, respectively). Only one carotid body PGL was missed by $^{18}$F-FDOPA-PET but pathological analysis showed this to be an atypical PGL that was fibrosed. In a recent series of nine patients with H&N PGL (total of 18 lesions), $^{18}$F-FDOPA-PET detected all lesions (18/18) compared to $^{18}$F-FDG (16/18), $^{111}$In-pentetreotide scintigraphy (11/17), $^{18}$F-FDA (8/18) and $^{123}$F-MIBG (5/18). Carbidopa, which decreases the decarboxylation and subsequent renal clearance of DOPA may be used to increase the tumour-to-background ratio of tracer uptake but had a low influence on the number of lesions depicted with $^{18}$F-FDOPA-PET since the vast majority of tumours had high uptake values.

$^{68}$Ga/SSTa

In recent years, $^{68}$Ga-DOTA-somatostatin analogue peptides that directly bind to somatostatin receptors on the tumour cells surface have been employed for the evaluation of neuroendocrine tumours with excellent results. Among the different peptides, $^{68}$Ga-DOTA-NOC may offer the advantage over others (TOC, TATE) to target a broader range of somatostatin receptors with high affinity for SSTR2.
Studies have shown the superiority of PET over $[^{111}\text{In}]$pentetreotide-SPECT in the detection of metastases and unknown primary neuroendocrine tumours. PET imaging using $[^{68}\text{Ga}]$SSTa was also more accurate than $[^{18}\text{F}]$FDOPA-PET in non carcinoid tumours. To date, the role of these emerging tracers in PGL is limited to case series.\textsuperscript{37,38,39,40,41,42} Recently, Naji et al found that $[^{68}\text{Ga}]-\text{DOTA-TATE}$ detected more tumour sites than $[^{123}\text{I}]-\text{MIBG}$ in patients with neural crest tumours (including 11 PGL). Only one bone lesion in one patient with adrenal PGL was detected on $[^{123}\text{I}]-\text{MIBG}$ scan but not in $[^{68}\text{Ga}]-\text{DOTA-TATE}$. One additional lesion in a SDHB patient was missed by both modalities but detected with $[^{18}\text{F}]$FDG-PET.\textsuperscript{38}

**Conclusion and perspectives**

The performance of molecular imaging modalities is widely dependent on tumour biology. Recent advances in genetic and molecular pathogenesis of PGL have allowed for the identification of new molecular diagnostic and therapeutic radiopharmaceuticals tailored to genetic abnormalities. However, the optimal diagnostic imaging algorithm remains to be determined.\textsuperscript{35,43}

**Proposed diagnostic algorithm**

According to the literature, $[^{18}\text{F}]$FDOPA-PET should replace $[^{111}\text{In}]$pentetreotide-SPECT as the first-line imaging in H&N PGL, but MIBG scan may be still required in combination with $[^{18}\text{F}]$FDOPA-PET in cases of abdominal and malignant forms. $[^{18}\text{F}]$FDG may add further information in SDHB mutation carriers, especially those with multifocal and/or metastatic disease. If available, $[^{18}\text{F}]$FDA seems to be a highly sensitive tracer in metastatic adrenal PGL and should be the first-line imaging method in non-SDHB inherited patients.

**Perspectives**
The above described tailored algorithm may of course further evolve if a future head to head comparison of $[^{18}\text{F}]$FDOPA-PET and $[^{68}\text{Ga}]$SSTa-PET in patients with parasympathetic PGL offers a new insight. PET imaging should also be considered for delineation of radiotherapy target volumes in H&N PGL, especially after debulking surgery. A summary of clinical indications of selected tracers are listed in Table 2.

The MIBG analogues $[^{124}\text{I}]$-MIBG, meta-$[^{76}\text{Br}]$bromobenzylguanidine ($[^{76}\text{Br}]$-MBBG), and 4-$[^{18}\text{F}]$-fluoro-3-IBG have been suggested and used in few studies but to our knowledge, no clinical studies have yet been reported. Many other peptide receptors are overexpressed in endocrine tumours and thus could represent new emerging targets for diagnosis and therapy. The influence of some new drugs on tracer uptake is also subject to investigations.

One point is clear, the way to image PGL is changing and nuclear medicine has a leading role to play.
Legend to Figures and Tables

Figure 1. Schematic display of mechanisms of uptake and retention of several PET tracers in tumour cells of neural crest origin

\[^{18}\text{F}\]: Fluorine-18; \[^{18}\text{F}\]FDG: \[^{18}\text{F}\]Fluorodeoxyglucose; \[^{18}\text{F}\]FDOPA: \[^{18}\text{F}\]-dihydroxyphenylalanine; \[^{18}\text{F}\]FDA: \[^{18}\text{F}\]Fluorodopamine; \[^{68}\text{Ga}\]SSTa: \[^{68}\text{Ga}\]-labelled somatostatin analogues; HK: Hexokinase, AADC: Aromatic L-amino decarboxylase, SSTR2: Somatostatin receptor type 2; SSTa: Somatostatin analogues; \[^{68}\text{Ga}\]: Gallium-68; GLUT: Glucose transporters; NET: Norepinephrine transporter; LAT1: sodium-independent large neutral amino acids transporter type 1, requires dimerization with CD98 for its targeting to the plasma membrane; VMAT: vesicular monoamine transporter 1 and 2.

Figure 2. \[^{18}\text{F}\]FDG PET in a non metastatic SDHB-related left adrenal paraganglioma.

Marked FDG tumour avidity (SUVmax 50) without any extra-adrenal foci. Axial CT view (A), axial attenuation-corrected \[^{18}\text{F}\]FDG-PET image (B), axial PET/CT fusion image (C), Maximum-Intensity-Projection (MIP) image (D). Pathological examination (E) revealed a 33 mm tumour without any unfavorable histological features (PASS score 0, Ki-67<1%, no mitosis).

Figure 3. \[^{18}\text{F}\]FDOPA PET in a SDHD-patient with multiple paragangliomas in the neck and the abdomen.

Coronal whole-body images showing bilateral neck and abdominal extra-adrenal paragangliomas (A), Maximum-Intensity-Projection (MIP) image (B), Axial (C), coronal (D) and sagittal (E) PET/CT views centered over the para-aortic abdominal parasympathetic paragangliomas.
Table 1. Summary of approved and experimental tracers.

Table 2. Summary of selected tracers used as first and second-line imaging modalities.
<table>
<thead>
<tr>
<th>Current status</th>
<th>Specificity</th>
<th>Targeting pathway</th>
<th>Mechanism of Uptake</th>
<th>Mechanism of retention</th>
</tr>
</thead>
<tbody>
<tr>
<td>Approved</td>
<td>Tumour cells</td>
<td>Glucose uptake, Energy metabolism, Vessels density, Angiogenesis (VEGF-A), Hypoxia</td>
<td>Glucose transporters</td>
<td>Phosphorylation (Hexokinase)</td>
</tr>
<tr>
<td>[123I]-MIBG</td>
<td>Approved</td>
<td>Chromaffin cells</td>
<td>NET</td>
<td>Storage in vesicles (VMATs)</td>
</tr>
<tr>
<td>(*)</td>
<td>Experimental</td>
<td>Catecholamine biosynthesis</td>
<td></td>
<td></td>
</tr>
<tr>
<td>[18F]-FDA</td>
<td>Approved</td>
<td>APUD cells</td>
<td>LAT1</td>
<td>Decarboxylation (AADC)</td>
</tr>
<tr>
<td>[18F]-FDOPA</td>
<td>Approved</td>
<td>Aminoacid uptake and decarboxylation</td>
<td></td>
<td></td>
</tr>
<tr>
<td>[68Ga]-SSTa</td>
<td>Experimental</td>
<td>Endocrine cells</td>
<td>LAT1</td>
<td>Internalization</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Somatostatin uptake and internalization (especially type 2)</td>
<td>SSTRx</td>
<td></td>
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</tr>
</tbody>
</table>

*([11C]-Hydroxyephedrine belongs to this category.

**Trapping is also dependent on the internalization of the tracer.

Table 2

<table>
<thead>
<tr>
<th></th>
<th>Non metastatic abdominal PGL</th>
<th>Non metastatic H&amp;N PGL</th>
<th>Metastatic PGL</th>
</tr>
</thead>
<tbody>
<tr>
<td>1&lt;sup&gt;st&lt;/sup&gt; Line</td>
<td>[123I]-MIBG</td>
<td>[18F]-FDOPA</td>
<td>[123I]-MIBG</td>
</tr>
<tr>
<td>2&lt;sup&gt;nd&lt;/sup&gt; Line</td>
<td>[18F]-FDOPA</td>
<td>[111In]-Pentetreotide&lt;sup&gt;1&lt;/sup&gt;</td>
<td>[18F]-FDG</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>[18F]-FDOPA&lt;sup&gt;3&lt;/sup&gt;</td>
</tr>
<tr>
<td>Experimental tracers</td>
<td>[18F]-FDA</td>
<td>[68Ga]-SSTa</td>
<td>[18F]-FDA</td>
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<tr>
<td></td>
<td></td>
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<td>[68Ga]-SSTa</td>
</tr>
</tbody>
</table>

<sup>1</sup>[111In]-Pentetreotide is performed when diagnosis of H&N PGL remains uncertain on [18F]-FDOPA.

<sup>2</sup>[123I]-MIBG is performed when abdominal tumour sites are suspected on CI, [18F]-FDOPA or in cases of previous history of abdominal surgery for PGL and/or presence of SDHD mutation.

<sup>3</sup>[18F]-FDOPA is more sensitive in non-SDHB than in SDHB-related metastatic PGL.
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