REVIEW

Nuclear medicine imaging of gastro-entero-pancreatic neuroendocrine tumors. The key role of cellular differentiation and tumor grade: from theory to clinical practice

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Abstract

Nuclear medicine imaging is a powerful diagnostic tool for the management of patients with gastro-entero-pancreatic neuroendocrine tumors, mainly developed considering some cellular characteristics that are specific to the neuroendocrine phenotype. Hence, overexpression of specific trans membrane receptors as well as the cellular ability to take up, accumulate, and decarboxylate amine precursors have been considered for diagnostic radiotracer development. Moreover, the glycolytic metabolism, which is not a specific energetic pathway of neuroendocrine tumors, has been proposed for radionuclide imaging of neuroendocrine tumors. The results of scintigraphic examinations reflect the pathologic features and tumor metabolic properties, allowing the in vivo characterization of the disease. In this article, the influence of both cellular differentiation and tumor grade in the scintigraphic pattern is reviewed according to the literature data. The relationship between nuclear imaging results and prognosis is also discussed. Despite the existence of a relationship between the results of scintigraphic imaging and cellular differentiation, tumor grade and patient outcome, the mechanism explaining the variability of the results needs further investigation.

Keywords: Neuroendocrine tumor; functional imaging; somatostatin receptor scintigraphy; FDG PET; FDOPA PET; differentiation.

Introduction and rationale

Neuroendocrine tumors (NET) are epithelial neoplasms with neuroendocrine differentiation. NETs are a heterogeneous group of rare tumors originating from peptide- and amine-producing cells of the neuroendocrine system. Their incidence is usually reported at between 40 and 50 cases per million, accounting for approximately 0.5% of all malignant neoplasms. Although most NETs originate from the gastrointestinal tract and pancreas, they only make up 2% of all gastro-entero-pancreatic (GEP) malignancies.

Only a small proportion of a GEPNET is functional. Flushes and diarrhea are the most reported symptoms related to the tumoral secretion of biologically active peptides and amines. The clinical symptomatology of GEPNET is generally associated with the presence of systemic metastatic spread. However, bowel obstruction or incidental detection of hepatic metastases are common circumstances that lead to diagnosis. Extensive tumor bulk and metastatic disease are the major causes of death. However, long patient survival, which is mainly related to the slow-growing tumor characteristics, is not unusual.
Table 1  Main bibliographic data on GEPNET reporting: functional imaging, tumor grade, tumor differentiation or patient outcome results

| Author          | Reference | Patients with GEPNET | SRS | Somatostatin imaging with PET analogs | FDOPA PET | FDG PET | Tumor grade | Cellular differentiation | Patient outcome |
|-----------------|-----------|----------------------|-----|--------------------------------------|-----------|---------|-------------|------------------------|----------------|
| Gabriel et al.  | [22]      | 84                   | 50  | +                                    | +         |         |             |                        |                |
| Buchmann et al. | [23]      | 27                   | 15  | +                                    | +         |         |             |                        |                |
| Srirajaskanthan et al. | [24] | 51                   | 37  | +                                    | +         |         |             |                        |                |
| Koopmans et al. | [33]      | 53                   | 32  | +                                    | +         |         |             |                        |                |
| Montravers et al. | [34]     | 30                   | 23  | +                                    | +         |         |             |                        |                |
| Ambrosini et al. | [44]      | 13                   | 11  | +                                    | +         |         |             |                        |                |
| Ambrosini et al. | [35]      | 13                   | 13  | +                                    | +         |         |             |                        |                |
| Haug et al.     | [45]      | 25                   | 14  | +                                    | +         |         |             |                        |                |
| Campana et al.  | [49]      | 47                   | 41  | +                                    | +         | +       | +           |                        |                |
| Adams et al.    | [50]      | 15                   | 7   | +                                    | +         | +       | +           |                        |                |
| Belhocine et al.| [51]      | 17                   | 11  | +                                    | +         | +       | +           |                        |                |
| Kayani et al.   | [52]      | 38                   | 28  | +                                    | +         | +       | +           |                        |                |
| Binderup et al. | [29]      | 96                   | 81  | +                                    | +         |         |             |                        |                |
| Abgral et al.   | [54]      | 18                   | 15  | +                                    | +         | +       | +           |                        |                |
| Garin et al.    | [56]      | 31                   | 23  | +                                    | +         | +       | +           |                        |                |
| Binderup et al. | [57]      | 98                   | 83  | +                                    | +         | +       | +           |                        |                |
| Asnacios et al. | [58]      | 98                   | 53  | +                                    | +         |         |             |                        |                |

Nowadays, surgery of the primary tumor is the only curative treatment in patients without metastatic disease\[4\]. Unfortunately, the detection of liver metastases at primary staging generally limits the possibility of complete surgical eradication. Hence, accurate staging is crucial to define the real extent of the disease and to identify patients with inoperable tumors\[5,6\].

Evidence exists that tumor differentiation and grade are related to the clinical behavior of GEPNETs\[7,8\]. Differentiation refers to the extent to which the tumor cells resemble their non-neoplastic counterparts. Well-differentiated GEPNETs present a low mitotic activity, and may have punctate necrosis. Cells are uniform and produce neurosecretory granules. Poorly differentiated neuroendocrine carcinomas show fields of necrosis, less cytoplasmic granularity, irregular nuclei, and marked cellular pleomorphism. The distinction between a well-differentiated and a poorly differentiated NET is one of the most important pathologic assessments: the first are rather indolent, whereas the second are likely to be more aggressive. Grade refers to the degree of biological aggressiveness of the tumor, without taking into account the pathologic differentiation. Low-grade NETs are generally considered as the least aggressive in behavior. On the other hand, high-grade tumors tend to grow rapidly and spread faster than tumors with lower grade. Intermediate grade NETs have a less predictable and moderately aggressive course. The proliferative rate can be assessed as the number of mitoses per unit area of tumor or as the percentage of neoplastic cells immunolabeling for the proliferation marker Ki-67. The Ki-67 index is related to the presence of the Ki-67 protein in the active phases of the cell cycle. Ki-67 is a proliferation marker commonly used in all grading systems and it is currently reported in pathology reports\[9\]. A Ki-67 index less than 2%, between 3% and 20%, and above 20%, is characteristic of low, intermediate, and high-grade GEPNET, respectively. Well-differentiated GEPNETs are classified as either low or intermediate grade, whereas poorly differentiated GEPNETs are classified as high grade in all cases\[7,10\]. The existence of a high-grade well-differentiated tumor is controversial, and does not figure in any expert recommendation or guideline\[11\].

Medical imaging is crucial for the management of NET patients. A multidisciplinary morphofunctional approach has been advocated, associating radiologic and nuclear medicine examinations. Nuclear medicine imaging for NET has been mainly developed considering some cellular characteristics that are specific for the neuroendocrine phenotype. Hence, the overexpression of specific trans membrane receptors as well as the cellular ability to take up, accumulate, and decarboxylate amine precursors have been considered for diagnostic radiotracer development\[12\]. Moreover, the glycolytic metabolism, which is not a specific energetic pathway of NETs, has been proposed for NET radionuclide imaging.

In this article, we briefly review the nuclear medicine diagnostic procedures that are mainly used in clinical practice for NET imaging. The influence of both cellular differentiation and tumor grade in the choice of isotopic examination is examined. The possible relationship between the result of nuclear imaging and patient prognosis is discussed. The main bibliographic data about GEPNETs reported in the article are summarized in Table 1.
Nuclear medicine imaging: a brief overview

Somatostatin receptor-based imaging

Somatostatin is a peptide produced by both paracrine and endocrine cells in the GEP tract. Somatostatin receptors (sstr) are coupled to cell membrane G-proteins. Five sstr subtypes are known, designated sstr1 to 5. In NETs, sstr2 and sstr5 are particularly abundant. Overexpression of receptors as well as internalization of the hormone–receptor complex are the basis of radiolabeled somatostatin imaging modalities. Octreotide conjugated with diethylene-triamine-pentaacetic acid (DTPA) and labeled with 111In is a radiolabeled somatostatin analog with great affinity for sstr2 and widely used in clinical practice.

Somatostatin receptor scintigraphy (SRS) is routinely indicated for staging purposes and during follow-up in patients with GEPNETs. Good sensitivity and specificity have been reported for SRS, irrespective of tumor site and hormonal secretion. SRS sensitivity is mainly related to the tumoral sstr2 density, decreasing for poorly differentiated tumors. The introduction of multiple-head gamma cameras and tomographic acquisition by single photon emission computed tomography (SPECT) has provided three-dimensional imaging, improving sensitivity. Moreover, the latest generation of high-resolution gamma cameras coupled with multidetector computed tomography (SPECT/CT) allows accurate anatomic localization of functional abnormalities, as well as a reduction in study duration and patient discomfort. Nevertheless, SRS is limited by the unfavorable physical characteristics of 111In, reducing the detection of small lesions, especially in organs with physiologic tracer uptake such as the liver. Accordingly, an inverse relationship between SRS sensitivity and the size of hepatic metastasis has been previously reported. EDDA/HYNIC-Tyr3-octreotide is a 99mTc-radiolabeled peptide that has been developed in Poland for clinical investigations and intraoperative imaging. 99mTc-EDDA/HYNIC-octreotate images revealed more metastatic NETs than 111In-octreotide in NETs who are potentially candidates for peptide receptor radionuclide therapy with 64Cu and/or 177Lu radiolabeled somatostatin analogs.

Catecholamine pathway-based imaging

GEPNETs originate from the amine precursor uptake and decarboxylation (APUD) system. Accordingly, the re-uptake of norepinephrine and the uptake of amine substrates such as dihydroxyphenylalanine (DOPA) have been exploited for nuclear medicine radiotracer development. 123I or 131I radiolabeled meta-iodobenzylguanidine (mIBG) is a structural analog of norepinephrine used for clinical imaging. mIBG uses active transport mechanisms to accumulate in the tumor in secretory vesicles. The sensitivity of mIBG scintigraphy for GEPNET has been reported to be about 50%, which is lower than the sensitivity of SRS. Although 123I-mIBG is less sensitive for GEPNET detection, it remains useful to predict uptake and response to radionuclide therapy with 131I-labeled mIBG. 123I-mIBG scans remain the first choice in nuclear medicine imaging for the management of patients with pheochromocytoma and neuroblastoma.

DOPA is an intermediate product in the synthesis of dopamine. Cellular uptake of DOPA is related to a specific cell membrane transport system. Once internalized, DOPA is decarboxylated to dopamine, and transported into secretory vesicles. For nuclear imaging purposes, DOPA is labeled with 18F (FDOPA) leading to the accumulation of 18F-dopamine in the cell. The high uptake of FDOPA in NETs is the result of the increased synthesis, storage and secretion of biogenic amines. FDOPA PET seems to be superior to morphological imaging and SRS in the localization of both the primary site of the GEPNET and metastatic localizations. Moreover, some authors have suggested that FDOPA PET is a valuable imaging tool for the detection of the primary tumor in patients with metastatic GEPNET of
unknown origin and negative or inconclusive findings at conventional imaging and SRS\(^ {15} \) (Fig. 1). However, its high cost and low availability are still important limiting factors for the routine use of FDOPA in clinical practice.

\(^ {11} \)C-5-Hydroxytryptophan (HTP) is a radiolabeled serotonin precursor representing an FDOPA-alternative related radiotracer. HTP is specifically taken up by carcinoid serotonin-producing tumors, decarboxylated by aromatic amino acid decarboxylase (AADC), and stored in vesicles as \(^ {11} \)C-serotonin and was shown to be more sensitive in imaging small NET lesions\(^ {16-38} \). Uptake, decarboxylation, and secretory vesicle transportation rely on the same systems and enzyme as FDOPA. However, the short half-life of \(^ {11} \)C requires an on-site cyclotron facility, making HTP not widely available for routine clinical use.

The sensitivity of FDOPA and HTP PET may be reduced for the detection of well-differentiated pancreatic NET due to high physiologic radiotracer uptake and retention in the whole pancreas. As recently reported by Tessonnier et al.\(^ {39} \) in 4 patients with solitary insulinomas, FDOPA PET identified tumor in only one case. The administration of carbidopa, an efficient inhibitor of the peripheral aromatic amino acid decarboxylase (AADC), has been shown to improve image interpretation for FDOPA and HTP PET by increasing tumoral uptake and lowering physiologic pancreatic uptake\(^ {32,40} \). On the other hand, no final consensus has been reached about the usefulness of AADC inhibition by carbidopa premedication before an FDOPA PET study in patients with insulinomas or \(\beta\)-cell hyperplasia\(^ {41-43} \).

Nowadays, few comparative studies investigating the diagnostic role of FDOPA and \(^ {68} \)Ga-DOTA-NOC or \(^ {68} \)Ga-DOTA-TATE PET/CT in GEPNET are available. Ambrosini et al.\(^ {44} \) reported a higher accuracy of \(^ {68} \)Ga-DOTA-NOC in the detection of primary tumor and metastatic disease compared with FDOPA in I1 patients with GEPNET. The detection rate of primary tumor with FDOPA was lower than previously reported by the same authors\(^ {35} \). These results are confirmed by Haug et al.\(^ {45} \), who reported a clear superiority of \(^ {68} \)Ga-DOTA-TATE to FDOPA PET in 14 patients. Nevertheless, prospective studies including larger and more homogeneous patient series are necessary to confirm the real superiority of PET/CT with somatostatin analogs towards FDOPA PET/CT in the diagnosis and follow-up of patients with GEPNET.

**Glucose metabolism-based imaging**

\(^ {18} \)F-Fluoro-2-deoxy-D-glucose (FDG) is a glucose analog labeled with positron-emitting \(^ {18} \)F, which is taken up into cells by the glucose molecular transporter. Once internalized, FDG is phosphorylated without entering any
Further metabolic processes. Thus, FDG remains trapped within the cell. The widespread clinical use in oncological routine of FDG PET as an imaging tool capable of exploring the differences of glucose metabolic activity between normal and pathologic tissues is justified by its high diagnostic accuracy and therapeutic impact. Nevertheless, limited value of FDG PET/CT is usually reported in the management of NETs, probably because of their low metabolic activity and slow growth. Despite these premises, FDG PET could have a potential value for prognosis stratification in patients with NET, as suggested by Strauss et al. in the early 1990s. Accordingly, NET with increased FDG uptake seem to have a more aggressive behavior with consequent less favorable long-term prognosis.

**Functional characterization of GEPNET by nuclear medicine imaging**

**Imaging results according to differentiation and tumor grade**

The expression of sstr on the cell surface is a landmark of neuroendocrine differentiation. Well-differentiated NETs are characterized by an important sstr density explaining the good sensitivity of nuclear imaging based on radiolabeled somatostatin analogs. On the other hand, less-differentiated tumors lose their ability to express sstr with consequent reduction in the diagnostic interest of SRS. Campana et al. reported a significant correlation between the scintigraphic results and NET differentiation in 47 patients studied by 68Ga-DOTA-NOC PET/CT. They showed that the intensity of radiotracer uptake was higher in well-differentiated tumors. They also demonstrated the relationship between the degree of radiotracer uptake and the clinical and pathologic features of NET. In the same manner, Srirajaskanthan et al. found a decreased sensitivity of 68Ga-DOTA-TATE PET/CT in patients with poorly differentiated NET.

An inverse relationship between FDG uptake and tumor differentiation has been observed. Increased glycolytic metabolism has been associated with high proliferative and poorly differentiated NETs. Conversely, a higher rate of false-negative results of FDG PET was found comparing FDG PET and SRS in patients with well-differentiated GEPNETs.

Based on these observations, the mismatch high uptake of somatostatin analogs/low glycolytic metabolism could be considered as the scintigraphic pattern of well-differentiated GEPNETs. Conversely, the scintigraphic pattern low uptake of somatostatin analogs/high glycolytic metabolism could be considered as representative of poorly differentiated tumors.

The potential relationship between the tumor grade and the results of scintigraphic exploration by labeled somatostatin analogs and FDG has still not been completely elucidated. For this purpose, Kayani et al. compared the diagnostic performances of 68Ga-DOTA-TATE and FDG PET/CT in 28 patients with GEPNET and 4 patients with metastatic NET of unknown origin. In this patient cohort, the uptake of 68Ga-DOTA-TATE was globally higher in low-grade NETs (Ki-67 less than 2%) than high-grade tumors. Conversely, lesions with a Ki-67 index above 20% showed a significantly higher FDG uptake. In patients with low-grade NET, only 21 of 123 detected lesions were FDG avid; 120 lesions showed 18Ga-DOTA-TATE uptake. In high-grade NETs, only 5 of 79 detected lesions were positive with 68Ga-DOTA-TATE, whereas 77 were positive for 18F-FDG uptake. Moreover, no lesion exhibited exclusive uptake of either 68Ga-DOTA-TATE in high-grade or FDG in low-grade NETs. In this study, tumor grade was even predictive of scintigraphic pattern.

According to these results, it seems possible to predict in vivo and with good accuracy both tumor grade and cellular differentiation from the analysis of nuclear medicine examinations. Moreover, the 2 scintigraphic patterns previously reported for tumor differentiation (i.e. high uptake of somatostatin analogs/low glycolytic metabolism versus low uptake of somatostatin analogs/high glycolytic metabolism) could also be appropriate for the grade.

Although well-differentiated NETs are often low-grade lesions, the finding of a well-differentiated endocrine carcinoma with a high Ki-67 index is of importance, increasing the interest for proliferative index estimation as a powerful prognostic tool. Currently, the choice of scintigraphic examination is mainly guided by tumor differentiation. Nevertheless, evidence exists about the important role played by tumor aggressiveness independently of the degree of cellular differentiation. For this purpose, Abgral et al. recently investigated the performances of SRS and FDG PET/CT in a rare subgroup of 18 patients with stage IV well-differentiated endocrine carcinoma characterized by a Ki-67 index above or equal to 10%. On a per patient analysis, SRS and FDG PET sensitivity was 83% and 100%, respectively. Taking as a standard the highest number of distinct lesions visualized by at least one imaging method (per lesion analysis), the sensitivities of SRS and FDG PET were 43% and 77%, respectively.
respectively, with a statistically significant superiority of FDG PET for detection of lymph node involvement. According to these results, the authors proposed FDG PET as the first choice scintigraphic method for staging of metastatic well-differentiated endocrine carcinoma with Ki-67 of 10% or higher.

All this evidence suggests that an intimate relationship between tumor differentiation and grade could explain the variability in the results of nuclear medicine examinations. Nevertheless, the exact influence of one on the another has still not been completely elucidated.

Nuclear imaging results for patient outcome prediction

There is a growing body of evidence suggesting the potentiality of FDG PET/CT in the prediction of long-term prognosis in patients with NET. A correlation between the intensity of tumoral radiotracer uptake and survival has been reported. Pasquali et al.\textsuperscript{[55]} first reported the accuracy of FDG PET for identifying NETs with rapidly growing or aggressive behavior, supporting the evidence that an increased glycolytic rate reveals a worse prognosis. More recently, Garin et al.\textsuperscript{[56]} evaluated the prognostic value of FDG PET/CT in a prospective study enrolling 38 patients with metastatic well-differentiated NET. The pancreas or small intestine was the site of the primary tumor in 20 of 38 patients. Histologically, 4 patients had a high-grade tumor and 34 had a low-grade tumor. Overall positive and negative predictive value of FDG PET/CT for the detection of disease progression within the first 6 months of follow-up was 93% and 91%, respectively. FDG PET/CT was found to be more sensitive than pathologic differentiation and Ki-67 in the early prediction of rapidly progressive disease. In 4 cases, there was a discrepancy between the pathologic examination (showing a well-differentiated tumor) and the Ki-67 immunostaining diagnosing a high-grade phenotype.

These results are consistent with the recent work of Binderup et al.\textsuperscript{[57]} based on a population of 98 patients with NET including 83 GEPNETs, 8 bronchopulmonary carcinoid tumors and 7 primary tumors of unknown origin. In this study, the prognostic value of FDG uptake, the proliferation index, chromogranin A, and liver metastases were assessed. During the 1-year follow-up, 13 of 14 dead patients were FDG PET positive and only one was FDG PET negative. Five of 13 lethal tumors showed a proliferation index below 2%, among which 4 showed FDG uptake. In this study, the positive prognostic value of FDG PET for the prediction of patient outcome was better than the Ki-67 index.

Conversely, positive findings with NET-specific radiotracers seem to be associated with a less aggressive disease and a better patient outcome. Asnacios et al.\textsuperscript{[58]} recently evaluated the correlation between SRS, sstr expression, and prognosis. Consecutive patients with

\textbf{Figure 2}  FDG (A,C) and FDOPA PET/CT (B,E) results performed before liver transplantation in a 49-year-old patient with a history of well-differentiated ileal NET surgically treated (Ki-67, 1%). Intense uptake of FDOPA was observed in multiple and voluminous hepatic metastases. FDG PET showed no or minimal uptake in the liver lesions.
a well-differentiated NET and negative (48 patients) or positive (50 patients) findings at SRS, were compared. The authors reported a better overall survival at 60 months when SRS was positive. Even in the absence of any treatment, SRS proved to be a valuable prognostic tool. Similar results were shown by Campana et al.\cite{49} investigating 47 patients with NET by $^{68}$Ga-DOTA-NOC PET/CT.

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*Figure 3* Results of FDG PET (A,C), SRS-SPECT (B,E) and corresponding non-enhanced CT slice (D) performed in a 63-year-old patient with poorly differentiated rectal NET (Ki-67, 11%). FDG PET allowed the detection of the primary tumor and both lymphatic para-aortic and right-lobe hepatic metastases. On the other hand, SRS showed no pathologic uptake, particularly in the liver (arrowheads).

*Figure 4* SRS (A), FDOPA (B,D) and FDG PET/CT (C,F) results performed for restaging purposes in a 66-year-old patient with a history of well-differentiated colic NET (Ki-67, 2%) previously treated by surgery. SRS only showed faint uptake concerning a peritoneal relapse (*). FDOPA PET showed a clear multifocal peritoneal carcinomatosis (arrowhead) and multiple hepatic metastases. Despite the absence of focal FDG uptake, diffuse and moderate uptake was observed in the pelvic cavity suggesting the shield sign, in keeping with peritoneal carcinomatosis.
Summary and conclusion

Nuclear medicine is a valuable non-invasive diagnostic modality for the staging and follow-up of patients with GEPNET. The results of scintigraphic examinations reflect the pathologic features and metabolic properties of the tumors, making the in vivo characterization of tumor differentiation and grade possible. The 2 scintigraphic patterns previously described (i.e. high uptake of somatostatin analogs/low glycolytic metabolism versus low uptake of somatostatin analogs/high glycolytic metabolism) could be considered as the extremes of all possible results. However, this binary-like classification is not applicable in all situations. A continuum of results combining a variable uptake degree of specific (radiolabeled somatostatin analogs, FDOPA) and non-specific (FDG) radiotracers is usually encountered in daily practice. The uptake intensity of radiolabeled analogs of somatostatin, FDOPA and FDG can vary in different lesions in the same patient mainly reflecting the tumor heterogeneity but also a possible variable degree of tumor differentiation (Fig. 5).

Even if cellular differentiation is commonly related to tumor grade, some exceptions exist, making both assessments necessary[54,57]. The association of multitracer nuclear investigations could play a role in the assessment of tumor differentiation and grade in a non-invasive manner, increasing the diagnostic accuracy and orienting therapeutic strategies[29,52,54,56]. In particular, the assessment of glycolytic metabolism by FDG PET seems to be potentially useful in identifying high-risk patients with more aggressive tumors, especially in those cases where there is disagreement between the pathologic results obtained from the biopsy of a metastatic location and the primary tumor after surgical debulking (Fig. 6).

Figure 5  Multiple hepatic metastases (C,D, contrast-enhanced CT maximum projection intensity and axial slice performed some days after the scintigraphic examinations) in a 57-year-old patient with poorly differentiated pancreatic NET previously treated by surgery. Biopsy confirmed the neuroendocrine nature of one hepatic lesion. The pathologic report showed a well-differentiated NET (Ki-67, <3%). Several hepatic foci of pathologic uptake were detected by FDG PET (A) and SRS (B). Some lesions showed both radiotracer uptake and other exclusive uptake of FDG or labeled somatostatin analogs (arrowhead). The discrepancy between the results of pathologic reports and isotopic procedures strongly suggests different degrees of tumor differentiation in the same patient.
Figure 6  FDOPA PET (A,D), FDG PET (B,F) and enhanced CT (C,E) results of a 65-year-old patient with a duodenal NET and liver metastatic involvement. Biopsy of one hepatic localization showed a well-differentiated carcinoma with Ki-67 index of 10%. The pathologic examination of the primary tumor showed poorly differentiated carcinoma with a Ki-67 index of 10%. Both FDG PET and FDOPA PET showed multiple non-concordant foci of pathologic uptake in the liver (arrowheads).

Figure 7  FDG PET (A,C) and FDOPA PET (B,E) results for a 68-year-old patient with clinical and biological suspicion of NET. FDG PET showed an exclusive pathologic uptake in hepatic segment IV–VIII, corresponding to a metastatic lesion on enhanced CT (D). Biopsy of hepatic metastasis (arrowheads) showed a well-differentiated NET of unknown origin with a Ki-67 index of 3%. FDOPA PET showed the ileal primary tumor (*) and multiple hepatic and lymph node metastases.
The management of patients with NET largely depends on the results of pathologic examination, which allows differentiation of aggressive malignancies from low-grade tumors. Generally, grade and differentiation are determined from a limited tissue sample obtained from biopsy or from partial surgical resection, which might be not truly representative of the whole tumor burden. On the other hand, nuclear medicine provides whole-body imaging, allowing extensive in vivo characterization of the tumor that is potentially useful for guiding biopsy, in particular of FDG-avid lesions that are not clearly detectable on CT or are largely necrotized or are avid in only a part (Fig. 7).

The prediction of patient outcome is a real challenge in clinical routine. Recently, the usefulness of nuclear medicine imaging for predicting patient prognosis has been advocated. Particularly, examinations based on FDG PET and radiolabeled somatostatin analogs may provide prognostic information independently from the pathologic indexes that are commonly accepted and used in daily routine. Moreover, tumor FDG uptake seems to be associated with a worse patient outcome even in well-differentiated or low-grade tumors. Despite the existence of a relationship between the tumor scintigraphic pattern, cellular differentiation, tumor grade and patient outcome, the intimate mechanism explaining the variability of the results has not been completely elucidated[59].

In conclusion, we emphasize the emerging role of nuclear medicine imaging in the management of NET. Hence, a better understanding of tumor biology and pathophysiology seems necessary to improve patient outcome. Prospective studies including larger and homogeneous patient series taking into account both cellular differentiation and tumor grade are necessary to confirm the real place for nuclear medicine in the diagnosis and follow-up of patients with NET.

**Conflict of interest statement**

The authors have no potential competing interests to declare. No funding was provided for this study.

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