Dear Sir,

We read with interest the paper of Lytras et al. [1] on the role of computed tomography (CT) and positron emission tomography (PET) in pancreatic cancer. We believe that the conclusions of the paper should be considered cautiously because of several procedure- and instrumentation-related problems.

The positiveness of fluorodeoxyglucose (FDG)-PET in pancreatic cancer is based on the high constitutive glucose uptake and metabolism that are characteristic features of cancer cells [2]. The glucose transporter-1 (GLUT-1) is a member of the GLUT family of facilitative glucose transporters that mediate Na⁺-independent cellular uptake of glucose. GLUT-1 has been reported to be over-expressed in a wide range of human cancers, including pancreatic cancer [3]. Tumoral GLUT-1 expression levels are inversely correlated with prognosis; furthermore GLUT-1 promotes invasive and metastatic potential of pancreatic cancer cell lines [3]. We can assume that less aggressive cancers may be false-negative at FDG-PET scan just because of lower levels of GLUT-1 in cancer cells, or because the desmoplastic reaction, typical of pancreatic cancer, reduces the number of cancer cells per square centimeter within the tumor area. It is known that FDG uptake in most malignant tissues increases with time even after 1 h post injection, while that of benign lesions decreases with time [4]. Both glucose and FDG are used avidly by cellular mediators of inflammation, and false-positive findings in the face of inflammatory changes in the pancreas have been reported [5]. A normal serum concentration of C-reactive protein, a normal leukocyte count and a normal concentration of alkaline phosphatase can help to exclude inflammatory changes [6]. Therefore the interpretation of FDG-PET can be influenced by differences of the procedure and/or of the instrumentation.

First of all it is well known that low standardized uptake values (SUVs) and false-negative FDG-PET scans have been found in hyperglycemic diabetic patients [5–10], with a decrease in sensitivity for the procedure of about 4% [9]. To optimize the results of FDG-PET, blood glucose (BG) levels should be checked and found below 130 mg/dl. A ‘modified SUV’ was also introduced to try to overcome this problem in patients with BG >130 mg/dl [7]. In Lytras et al.’s [1] paper, 10 patients with BG concentrations >10 mmol/l (180.2 mg/dl) were excluded from the comparative analysis, a limit much higher than 130 mg/dl (7.2 mmol/l) suggested by most authors [5–9]. Furthermore, the number of patients with BG levels between 130 and 180 mg/dl was not reported, and no attempt was made to calculate the ‘modified SUV’.

The scan acquisition time is also important for the accuracy of the procedure. Image interpretation in FDG-PET is usually performed for images obtained 1 h post FDG injection [7, 8, 10]. Nakamoto et al. [4] found no significant differences in detection performance of primary tumor between images acquired 1 and 2 h after FDG injection, but the certainty of excluding liver metastases was increased when the 2-hour image was used. In Lytras et al.’s [1] study, scanning started 15 min after FDG injection and continued for 40 min (instead of the usual 10 min acquisition time). This means that scanning started much earlier than generally used, and continued for a quite long time, by this way contributing to a decrease in the accuracy of the procedure.

The interpretation of images obtained with an FDG-PET scan is also crucial. The visual interpretation, as used in the Lytras’ study, is not the best one. Visual analysis is reader dependent and less sensitive than SUV. In the study of Koyama et al. [11], the sensitivity and specificity were 82 and 81% for visual and 93 and 80% for quantitative (SUV) interpretation. Also the use of semi-quantitative analysis methods such as SUV is not yet clearly defined to differentiate malignant from benign pancreatic lesions; a cut-off value of SUV valid for all centers cannot be defined due to the high variability of absolute SUV values among different institutions [12]. However, it is still the most frequently used and reliable method for the interpretation of images obtained with FDG-PET scan.

The choice of instrumentation is of outstanding value for the optimization of the results. The evolution of PET detectors to PET scanners and the progressive improve-
ment of the physical performance of commercial PET scanners [13] render obsolete the results obtained with old instrumentation, like the coincidence detection of gamma-cameras used by Lytras et al. [1]. In oncology, this technique means low resolution and poor detection efficiency.

In conclusion, preoperative diagnosis of pancreatic cancer is useful to reduce the number of aborted resections by less aggressive surgeons, with subsequent reoperative pancreateoduodenectomy in referral centers, and to reduce the rate of inadvertent resections of benign disease by less experienced surgeons [5]. Among newer imaging modalities aimed to avoid these inadequate treatments, 18-FDG-PET is believed by several authors to have an important role in the diagnosis [5,10] and prognosis [8] while others require further evaluation [9]. A consensus conference in Germany concluded that FDG-PET had achieved an established clinical role in the diagnosis and staging of pancreatic cancer [6]. We believe that the conclusions drawn by Lytras et al. [1] on the basis of the comparison between an up-to-date CT and a PET-CT with several procedure and instrumentation bias should be regarded cautiously, especially considering the opposite results and conclusions of a recent paper from Zurich using a last-generation PET-CT [14].

References

Reply

We appreciate the interest shown by Pedrazzoli and colleagues in the paper by Lytras et al. [1].

FDG-PET scans rely on the uptake of glucose by the targeted cells. The glucose transporter 1 (GLUT-1) is a member of the GLUT family of facilitative glucose transporters. Overexpression of GLUT-1 has been demonstrated in 88% of pancreatic cancer (in small numbers) and also in 50% of benign tumours [2]. The association of GLUT-1 and tumour invasiveness has only been shown in vitro at the present time [3].

It is not quite true to say that less aggressive cancers may be FDG-PET scan false-negative as even benign tumours can overexpress GLUT-1 [2]. The key issue is the overcalling of benign lesions to be cancer in the face of inflammation or benign disease [4].

The main differential diagnosis for these patients is an inflammatory mass due to chronic pancreatitis. For these patients the C-reactive protein, leukocyte count and alkaline phosphatase may be normal and therefore cannot be relied upon to rule out inflammatory causes. The serum CA 19-9 may be raised in these patients with inflammation, thus adding to the diagnostic difficulty [5].

There is indeed difficulty in interpreting FDG-PET scans due to different protocols and the effects of body weight and glucose metabolism [6]. Of those patients who had between 130 and 180 mg/dl glucose in our study (total of 11 out of 118 patients, or 9%), 6 were malignant on PET scanning and 4 were non-malignant (1 was indeterminate).

The timing of the PET scan and acquisition in our study was 50 min (not 15 min as written in the paper), then continued for 40 min. This was appropriate for our instrumentation.

SUVs are not appropriate when using gamma-camera PET.

The number of patients involved in this prospective study is large and therefore results from this study are valuable compared to studies of fewer patients with different instrumentation.

The study by Heinrich et al. [6] used a small number of patients and is therefore

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underpowered. Histological diagnosis was available in 52 patients only. It was claimed that PET-CT detected additional distant metastases in 5 and synchronous rectal cancer in 2 patients and that the findings changed the management in 16% of patients, with pancreatic cancer deemed resectable after routine staging (p = 0.031). This study did not utilize contrast-enhanced multi-slice CT scan, which is widely regarded as the gold standard and therefore the conclusions are not valid, as these results could be completely reproducible using CE-CT scan only: a fact acknowledged by the authors themselves. This is the first report describing a coincidence between pancreatic cancer and rectal cancer of 4% and must therefore be treated with considerable caution. The study was not hypothesis driven and was without appropriate power calculations. The conclusions are not supported by the study because of poor design and an invalid comparison.

Advances in technology such as new generation PET-CT scanners and newer contrast agents such as $^{[18}F]\text{FLT}$ will need further assessment [7]. However, the study highlights the importance of an integrated approach between diagnostic and clinical teams for patient management.

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References


