Rapid on-site cytopathological examination (ROSE) performed by endosonagrapheers and its improvement in the diagnosis of pancreatic solid lesions1


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ABSTRACT

PURPOSE: To evaluate the diagnosis improvement of EUS-FNA when using ROSE performed by the endosonographer.

METHODS: A retrospective study was conducted. A total of 48 pancreatic solid masses EUS-FNA were divided into two groups according to the availability of on-site cytology (ROSE) – the first 24 patients (group A-without ROSE) and the latter 24 cases (group B-with ROSE). Sensitivity, specificity, positive predictive value, negative predictive value, accuracy, complications and inadequacy rate of EUS-FNA were determined and compared.

RESULTS: Among the 48 EUS-FNA, the overall performance was: sensitivity 82%; specificity 100%; positive predictive value (PPV) 100%; negative predictive value (NPV) 70% and accuracy 87%. The sensitivity of the Group A was 71%, versus 94% in-group B (p=0.61). Moreover, the negative predictive value was 58% versus 87% (p=0.72). The accuracy rate increased from 79% to 96% (p=0.67) in the ROSE group. The number of punctures was similar between the groups. No major complications were reported.

CONCLUSION: Rapid on-site cytopathological examination, even when performed by the endosonographer, may improve the diagnostic performance in the diagnosis of solid pancreatic lesions, regardless of the slight increase in the number of punctures.

Key words: Endosonography. Pancreatic Neoplasms. Cell Biology.
Introduction

Endoscopic ultrasound (EUS) was introduced in clinical practice in 1980 in order to improve the appreciation of the digestive tract walls and adjacent structures. Later, with the development of endoscopic ultrasound-guided fine needle aspiration (EUS-FNA), obtaining tissue specimen was made possible; thereby further improving its accuracy\(^1\)-\(^3\).

EUS is the most accurate method for loco-regional staging of upper gastrointestinal cancers, with well-established clinical impact, influencing decision-making and patient management. Furthermore, EUS-FNA proved to be a safe, accurate, and a reliable diagnostic procedure, with a high diagnostic and therapeutic yield\(^4\). The rapid on-site cytopathological examination (ROSE) during the procedure appears to have a significant impact on EUS-FNA success rates\(^5\).

The objectives of the study are to determine the overall performance of EUS-FNA in the diagnosis of solid pancreatic lesions in a Latin American EUS training center and to evaluate the diagnosis improvement of EUS-FNA when using ROSE performed by the endosonographer.

Methods

This is a single center prospectively enrolled retrospective study, including all patients referred for EUS examinations for pancreatic solid lesions from January 2009 to November 2011. The study was conducted at the French-Brazilian Centre of Endoscopic Ultrasonography (CFBEUS), located at the endoscopy unit of Santa Casa of Sao Paulo Hospital, Brazil.

After obtaining formal informed written consent, an anesthesiologist sedated all patients. EUS-FNA was performed by using an EG-530UT linear echoendoscope and SU-7000 ultrasonic processor (Fujinon, Saitama, Japan) and 22-G EchoTip\(^\text{®}\) needle (Cook Medical Inc, Limerick, Ireland).

During the study period, a total of 963 EUS examinations were performed, of which, 71 EUS-FNA of suspected pancreatic lesions were evaluated, including 48 solid lesions and 23 cystic lesions.

The present article regards pancreatic solid lesions, therefore, only the 48 EUS-FNA of such lesions were included. Single and multiple FNA passes (1 to 7 passes) were done in the first 24 EUS-FNA cases (group A) (Figure 1). For the latter 24 cases (group B), EUS-FNA passes were performed until ROSE evaluation confirmed the presence of a sufficient number of representative lesion cells. The same endosonographer performed the fine needle aspirates and prepared the cytology slides using the Diff-Quik\(^\text{®}\) stain set\(^6\). The tissue specimens obtained were also immediately formalin-fixed for further cell-block study.

FIGURE 1 - EUS-FNA of a pancreatic mass.

The endosonographer had sufficient experience in more than 300 pancreatic and biliar EUS-FNA procedures and underwent formal cytology training with the cytopathologist. This included at least 30 hours of theoretical-practical and review of slides, with the board-certified cytopathologist demonstrating examples of adequate, inadequate, benign and malignant slides (Figure 2).
Rapid on-site cytopathological examination (ROSE) performed by endosonagraphers and its improvement in the diagnosis of pancreatic solid lesions

For the EUS-FNA procedures, the standard suction technique was used, with the 10 mL syringe that is part of the EchoTip® needle kit.

The final diagnosis was based on the examination of the results of the slides, the cell-block and surgical pathology specimens. Technical failure was considered when there was insufficient aspirated material to determine a final diagnosis, according to the pathologist, and this datum was used to calculate the inadequacy rate.

The definition of complications followed the same criteria used by the American Society for Gastrointestinal Endoscopy (ASGE) and European Society of Gastrointestinal Endoscopy (ESGE) guidelines.

All patients signed an informed consent and the ethics committee of the institution approved the study.

Statistical analysis

Initially, all variables were analyzed descriptively. For quantitative variables, the analysis was done through observation of minimum and maximum values and the calculation of mean values, standard deviations. For qualitative variables, we calculated absolute and relative frequencies. To compare the proportions of failures of the methods we used the Fisher’s Exact Test. To study the efficiency of the methods we analyzed the values of sensitivity, specificity, accuracy, positive and negative predictive values. The significance level used for the tests was 5%.

Results

Among the 48 EUS-FNA of pancreatic solid masses, the overall performance was: sensitivity 82%; specificity 100%; positive predictive value (PPV) 100%; negative predictive value (NPV) 70% and accuracy 87%.

The sensitivity of the Group A was 71%, versus 94% on group B (p=0.61). The specificity of both groups was 100%, as well as the positive predictive value. Moreover, the negative predictive value was 58% versus 87% (p=0.72). The accuracy rate increased from 79% to 96% (p=0.67) on the ROSE group. These results can be appreciated in Table 1.

![Cytology slide: pancreatic adenocarcinoma (Diff-Quik stain®).](image)

**TABLE 1** - Performance for EUS-FNA of pancreatic solid lesions.

<table>
<thead>
<tr>
<th>TABLE 1</th>
<th>SENSITIVITY</th>
<th>ESPECIFICITY</th>
<th>PPV</th>
<th>NPV</th>
<th>ACCURACY</th>
</tr>
</thead>
<tbody>
<tr>
<td>OVERALL PERFORMANCE</td>
<td>82%</td>
<td>100%</td>
<td>100%</td>
<td>70%</td>
<td>87%</td>
</tr>
<tr>
<td>GROUP A (WITHOUT ROSE)</td>
<td>71%</td>
<td>100%</td>
<td>100%</td>
<td>58%</td>
<td>79%</td>
</tr>
<tr>
<td>GROUP B (WITH ROSE)</td>
<td>94%</td>
<td>100%</td>
<td>100%</td>
<td>87%</td>
<td>96%</td>
</tr>
</tbody>
</table>

*PPV = positive predictive value, NPV = negative predictive value

The inadequacy rate (insufficient material) on Group A was 20.8 % (5/24 patients) and 4.15% (1/24 patients) in-group B (p=0.19). All of these patients died of advanced disease and were considered false negative (6/48-12.5%). In-group A, the number of punctures ranged from 1 to 6 with a mean value of 3.5. In-group B, the number of punctures ranged from 3 to 8, with a mean value of 4.3. These results were not statistically significant. There were no major complications in this study.

Two patients in group A (8.3%) and three (12.5%) in group B with immediate mild abdominal pain, which resolved with analgesics and did not prolong hospital stay. Two patients (8.3%) in each group complained of a sore throat for a few days that resolved with oral Benzocaine.

The most common diagnosis is the pancreatic adenocarcinoma, followed by the results of non-neoplastic diseases such as pancreatitis and normal pancreatic tissue (Figure 3).
Discussion

Pancreatic adenocarcinoma is the second most common gastrointestinal malignancy and the fourth leading cause of cancer-related mortality in the United States. EUS has well-defined accuracy in GI malignancies stages and is the most accurate method in establishing the presence of pancreatic lesions, but cannot alone, reliably differentiate benign from malignant lesions. Consequently, pathological examination is often required to establish a definitive diagnosis.  

EUS-guided fine needle aspiration (EUS-FNA) was first described in 1991 for the evaluation of gastric sub-mucosal lesions and pancreatic cancer. For EUS-FNA of pancreatic lesions, many different needles can be used, nonetheless. The results don’t appear to change much and the complication rate may increase when using a 19G. Therefore we adopted the 22G needle for this study.  

Even though, nowadays different suction techniques may be applied, such as slow-pull, or no suction at all, at the time of the beginning of the protocol, that was not an issue and it was not considered as a variable then.  

There are studies that have examined the improvement of diagnostic ability of EUS-FNA without on-site cytopathologist, but with the immediate microscopic inspection by endosonographers. In a retrospective study, Savoy et al. compared the abilities of endosonographers and cytotechnologists in the immediate evaluation of microscopic inspection. The respective accuracies were of 70% and 89%. In our study, the ROSE-group accuracy was 96%, which can be considered similar to the cytotechnologists on Savoy’s study.  

In contrast, Erickson et al. and Nasuti et al. showed that the absence of ROSE at all, resulted in poor diagnostic accuracy as well as increased procedure time, number of needles used, and overall examination costs. More recently, Iglesias-Garcia et al. demonstrated that ROSE decreased the number of passes with the EUS needle, the inadequacy of the collected specimen, as well as was associated with a significantly higher diagnostic sensitivity (96.2 vs. 78.2%; p=0.002) and overall accuracy (96.8 vs. 86.2%; P=0.013) for malignancy.  

In our study, after ROSE performed by the endosonographer, the sensitivity increased from 71% to 94% (p=0.61), the negative predictive value from 58% to 87% (p=0.72) and the accuracy from 79% to 96% (p=0.67). The insufficient aspirated material decreased from 20.8% to 4.1% (p= 0.19). Although, not statistically significant, ROSE increased the accuracy of the method and reduced technical failure. Iglesias-Garcia et al. observed the same pattern in the previously quoted study, although their inadequacy rates were lower than ours altogether (12.6% without ROSE and 1% with ROSE). We could argue whether the pathologists of their group are better acquainted with pancreatic cytology or that technically the endosonographer of our study is less experienced.

FIGURE 3 - Final histological diagnosis of the punctured lesions.
With that in mind and using our data as a starting point, we have an ongoing prospective study in our center, on ROSE, where we intend to enroll a larger sample size.

Nonetheless, our data is similar to other articles in this field, such as the meta-analysis recently published by Hewitt et al.\textsuperscript{11}. In his study the pooled sensitivity for malignant cytology was 85% and pooled specificity was 98%.

One may wonder how we managed, in a retrospective study, to have exactly 24 patients in each group. The manner we chose to draw the protocol explains that. The idea was to finish the study in three years. When we reached 18 months in the study, we concluded the first group. The number of patients enrolled was 24. Therefore, we decided to perform ROSE, in the following 24 patients. The data acquired was analyzed retrospectively. That’s why, this may be called a prospectively enrolled retrospective study.

In the present article, the appliance of ROSE caused the mean number of punctures to increase from 3.5 to 4.3. Even though this was not significant, it was somehow unexpected, since ROSE is thought to decrease the number of punctures needed to confirm the diagnosis, as suggested by Iglesias-Garcia\textsuperscript{7}. Some may argue that the increase in the number of punctures, might justify the better result by itself, leaving no benefit for the ROSE procedure. We do not believe that, because, the number of punctures did not differ that much (3.5 - 4.3) and was far from being significant.

On the other hand, the apparent improvement in performance and the decrease of inadequate material samples justifies the use of ROSE.

There were no major complications in this study. As previous articles demonstrated, EUS-FNA is a safe procedure with a complication rate of approximately 1-2%. Major complications include infection, bleeding, and acute pancreatitis and are more frequent for EUS-FNA of cystic compared with solid lesions\textsuperscript{7,8,21,22}. The main difference between this study and the others is the fact that, the number of punctures needed for diagnosis with ROSE did not decrease, on the contrary. Nonetheless, the apparent improvement in performance, suggests that this is not an issue, as long as this trend proves to be true. Most likely, larger and multicenter trials might be able to do so.

\section*{Conclusion}

Rapid on-site cytopathological examination, even when performed by the endosonographer, may improve the diagnostic performance in the diagnosis of solid pancreatic lesions, regardless of the slight increase in the number of punctures.

\section*{References}


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