

Improving the results of pancreatic endoscopic ultrasound-guided fine needle aspiration in daily practice: keep it simple

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submitted

23. December 2014

accepted after revision

7. January 2015

Bibliography

DOI <http://dx.doi.org/10.1055/s-0034-1391416>
Published online: 11.2.2015
Endoscopy International Open 2015; 3: E138–E139
© Georg Thieme Verlag KG
Stuttgart · New York
E-ISSN 2196-9736

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Endoscopic ultrasound-guided fine needle aspiration (EUS-FNA) is an indispensable tool for tissue acquisition. The National Comprehensive Cancer Network has incorporated EUS-FNA cytology in its diagnostic algorithm for pancreatic cancer [1]. EUS-FNA is a multi-step procedure involving not only best practices in technique but also in specimen collection and processing; thus, rapid on-site evaluation (ROSE), needle selection, and sampling technique determine the outcomes of EUS-FNA diagnosis of pancreatic cancer. The availability of on-site cytopathology services increases diagnostic accuracy and decreases the number of suboptimal specimens such that the non-diagnostic rate has been reported to be as low as 1% for FNAs with ROSE, whereas it is 20% without ROSE [2,3]. However, ROSE is not available at many centers because of the time needed, cost, and inadequate financial reimbursement. Hence, most centers depend on off-site laboratory interpretation of FNA specimens.

In this issue of the journal, Schneider and colleagues from Germany report on 63 patients who underwent EUS-FNA of pancreatic lesions in two phases: In phase I, the in-house pathology laboratory evaluated the FNA aspirates, and in phase II, a specialized outside cytology laboratory evaluated them [4]. The diagnostic sensitivity in phase II improved to 91.4% from 38.5% found in phase I, which emphasizes the importance of specialized diagnostic services in evaluating FNA specimens. In a retrospective study of 876 EUS-FNA specimens obtained from a tertiary referral center, the overall cumulative sum charts for both the non-diagnostic error rate and diagnostic error rate showed a short learning period and improved to an acceptable level at case numbers 121 and 97, respectively [5]. These findings suggest that like the practice of EUS-FNA where diagnosis improves with the endosonographer's experience [6], the accuracy of specimen interpretation improves with the cytopathologist's experience.

Therefore, high-volume centers practicing EUS-FNA must implement ROSE or the services of an off-site cytopathology laboratory to obtain the best diagnostic outcomes.

In addition to specimen interpretation as performed by cytopathologists, the endosonographers must be familiar with the techniques for handling and processing tissues to optimize specimens sent for off-site evaluation. Assigning one person to handle the aspirated materials establishes a protocol for consistent sample preparation and minimizes artifacts resulting from improper tissue handling and collection. Although most endosonographers do not routinely practice in-room cytomorphological analysis, it has been shown that microscopic evaluation of smears by endosonographers without access to ROSE improves diagnostic accuracy [7]. In a recent study, we showed that a short, intensive EUS cytopathology course for endosonographers provided effective training in cytopathological interpretation [8]. In our opinion, consideration must be given to incorporating basic cytopathology during an EUS fellowship to improve the endosonographer's performance of FNA procedures.

There are several technical issues that are critical to improving the results of EUS-FNA of pancreatic lesions. These include needle selection, FNA technique, use of suction or stylet, and the number of passes. Most masses in the pancreas originate in the head region and require transduodenal sampling. This mandates the need for a flexible, thin caliber needle that can easily exit the biopsy channel of the echoendoscope; that is, a needle with an angled tip used during sampling of head and uncinate masses. Limited evidence suggests that a 25G needle is suitable for sampling pancreatic masses. A recent meta-analysis compared the 22G and 25G needles for EUS-FNA of pancreatic masses and found that use of the 25G needle yielded a higher diagnostic sensitivity than the 22G needle [9]. However, if the endosonographer

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must rely on only the cell block for off-site diagnosis, the 25G needle may not be optimal. In a recent randomized trial, we demonstrated that the 25G needle yielded a diagnostic cell block in only 80.5% of patients irrespective of the number of FNA passes performed [10]. Therefore, in most centers in Europe and Asia, where ROSE is not available, the 22G needle is used in conjunction with suction for better tissue procurement. There has been growing interest in the use of core biopsy needles for histological tissue procurement when sampling pancreatic masses. However, in a recent meta-analysis, it was evident that the performance of the ProCore biopsy needle was not superior to the performance of the standard FNA needle either for cytological or histological assessment [11].

Although many techniques have been suggested for tissue procurement, our recommendation is to keep the procedure simple! We recommend fanning the needle when sampling a mass. This ensures not only the procurement of good quality tissue that is representative of the lesion (minimizing sampling error) but also allows diagnosis with fewer passes [12]. In addition, we do not recommend the routine use of suction during EUS-FNA because it increases the amount of blood in the specimen. Suction may be used when sampling a mass in the setting of chronic pancreatitis when tissue yield can be minimal, and there is a growing body of evidence that suction may augment the performance of the 25G needle. In a randomized trial comparing high negative pressure suction (50 mL negative pressure) with normal negative pressure suction (10 mL negative pressure) for EUS-FNA of pancreatic masses using the 25G needle, a higher proportion of diagnostically adequate samples were obtained using high negative pressure suction (90.0% vs 72.2%, $P=0.003$) [13]. We do not recommend the routine use of a stylet during EUS-FNA because it prolongs the procedure, increases the amount of blood in the specimen, and does not improve the accuracy of the diagnosis. Finally, when sampling pancreatic masses, one reaches a point of “diminishing returns” after seven passes [14]. The most common reason for non-diagnostic interpretation is inadequate sampling and, hence, a sufficient number of passes must be performed to yield an adequate sample.

Therefore, our recommendations to endosonographers for superior outcomes during EUS-FNA of the pancreas include: 1) use of a 25G needle, 2) use of a 19/22G needle if the diagnosis is to be established only by cell block or when ROSE is not available, 3) use of the fanning technique, 4) avoidance of the routine use of suction and stylet, 5) procuring an adequate quantity of the specimen, 6) learning to self-assess for diagnostic adequacy of FNA specimens if ROSE is not available, and 7) hiring the services of an expert cytopathologist who can provide off-site interpretations. EUS-FNA is an art — just keep it simple!

Abbreviations



EUS	endoscopic ultrasound
FNA	fine needle aspiration
ROSE	rapid on-site evaluation

Competing interests: None

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