Prospective Assessment of the Performance of a New Fine Needle Biopsy Device for EUS-Guided Sampling of Solid Lesions

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Background/Aims: Endoscopic ultrasound-guided fine needle aspiration (EUS-FNA) remains the most common EUS-guided tissue acquisition technique. This study aimed to evaluate the performance of a new Franseen tip fine needle biopsy (FNB) device for EUS-guided sampling of solid lesions and compare it with the historical FNA technique.

Methods: Acquire® 22 G FNB needle (Boston Scientific Co., Natick, MA, USA) was used for solid tumor sampling (Study group). Tissue was collected for rapid on-site evaluation, and touch and crush preparations were made. Historical EUS-FNA samples obtained using Expect® 22 G FNA needle (Boston Scientific Co.) were used as controls (Control group). All specimens were independently evaluated by two cytopathologists blinded to the formal cytopathological diagnosis.

Results: Mean cell block histology scores were significantly higher ($p=0.046$) in the FNB group (51 samples) despite a significantly lower ($p<0.001$) mean number of passes compared to the FNA group (50 specimens). The overall diagnostic yields for the FNB vs. FNA groups were 96% vs. 88%. The degree of tumor differentiation was adequately assessed in all cell block qualifying lesions in the FNB group. Two patients developed post-FNB abdominal pain.

Conclusions: The new Franseen tip FNB device provides histologically superior and cytologically comparable specimens to those obtained by FNA, but with fewer passes. Clin Endosc 2018;51:576-583

Key words: Tissue acquisition; Endoscopic ultrasound-guided fine needle aspiration; Endoscopic ultrasound-guided fine needle biopsy

INTRODUCTION

Endoscopic ultrasound-guided fine needle aspiration (EUS-FNA) remains the most common EUS-guided tissue acquisition technique. However, its reported sensitivity ranges from 65% to 95%. Several factors can influence the diagnostic yield of any EUS-guided sampling procedure, including the needle type and size, number of passes, lesion-specific characteristics, such as location and size, and, finally, endosonographer’s experience. The presence of rapid on-site cytological evaluation (ROSE) can positively impact the outcome of an EUS-FNA procedure, although its overall impact on the diagnostic yield of malignant lesions has been recently challenged. The diagnostic yield of FNA has been consistently high in pancreatic solid lesions, exceeding 85%, but varies in subepithelial tumors, ranging from 45% to 90%. A newer generation of fine needle biopsy (FNB) needles became available in the recent years and helped improve the tissue yield compared to the first generation of Trucut biopsy needles. A recent meta-analysis demonstrated comparable diagnostic yield to standard FNA needles with fewer passes.

This study aimed to evaluate the overall diagnostic performance of a new FNB device with a Franseen tip equipped with three cutting edges for EUS sampling of solid lesions (Fig. 1). We sought to assess the cytological and histological yield of the FNB samples compared to those of historical FNA sam-
amples (controls) obtained using a similar size needle. Our secondary goals were to compare the number of passes required to achieve a cytological diagnosis by each method and to assess the cytological quality of smears prepared from FNB samples for immediate cytological interpretation. Finally, we report on the adverse events associated with the use of this new FNB device.

MATERIALS AND METHODS

Study design and study population
This single-center study was approved by the Indiana University School of Medicine Institutional Review Board. Consecutive patients referred to a single operator (MAH) for EUS-guided sampling of solid lesions were prospectively enrolled between September 2016 and March 2017. All patients underwent EUS-FNB. ROSE was available in all cases. Patients were excluded if no cell block material was collected or if no standard reference diagnosis was obtainable.

Patient demographics and lesion-related characteristics, including the size and location of target lesions, cytological and pathological findings (in case of lesion surgical resection), and clinical outcomes, were recorded. FNB-related abstracted data included: smear cytology score, cell block histology score, number of passes for smears, number of additional passes for cell block, and total number of passes.

EUS-FNA samples previously accrued from solid lesions by the same endosonographer (MAH) between 2013 and 2015 using the Expect® 22 G FNA needle (Boston Scientific Co., Natick, MA, USA) were identified and used as controls. ROSE was available in all FNA cases, as this is the standard of care at our center, and was performed in the same way described below for the FNB group.

Matching of controls to cases was performed in two stages: first, consecutive samples obtained from solid lesions were included; subsequently, matching of cases and controls by site (pancreatic vs. non-pancreatic) was attempted. Only controls with available cell block histology score and confirmed diagnosis were included using the same reference diagnostic standard of the Study (FNB) group. All specimens were independently evaluated by two expert cytopathologists (SR and HW) to assess the cytological yield (on smears) and histological yield (on cell block preparations) using a standard scoring system reached by a consensus among our cytopathologists prior to the launching of the study (Table 1, Figs. 2-6). The cytopathologists were blinded to the formal cytological diagnosis rendered for clinical care in all FNB cases and controls.

Study definitions
A clear, non-equivocal diagnosis of malignancy was based on neoplastic cyto-histopathology by endoscopic sampling (FNB or FNA) and/or surgical resection. When cytology was non-diagnostic, the diagnosis of malignancy was considered based on neoplastic EUS impression, clinical course, and imaging findings consistent with malignancy at ≥6 months of follow-up.

EUS-FNB technique and ROSE
All procedures were performed by the same experienced endosonographer (MAH). EUS-FNB was performed using a linear array echoendoscope (Olympus America Inc., Center Valley, PA, USA) and a 22 G Acquire® FNB needle (Boston Scientific Co.). Monitored anesthesia care sedation was provided by anesthesiologists for all procedures. The slow-pull technique was used to remove the stylet and was followed

Table 1. Cytological and Histological Scoring Systems Used in the Study

| Scoring System | Cytology criteria (smears) | Histology criteria (cellblock) |
|----------------|---------------------------|-------------------------------|
| Score 1        | 5 or fewer groups (≥10 cells each group) or <50 tumor cells for tumors with dishesive pattern | 5 or fewer groups (≥10 cells each group) or <50 tumor cells for tumors with dishesive pattern |
| Score 2        | 6-10 groups (≥10 cells each group) or 50–100 tumor cells for tumors with dishesive pattern | 6–10 groups (≥10 cells each group) or 50–100 tumor cells for tumors with dishesive pattern |
| Score 3        | >10 groups (≥10 cells each group) or >100 tumor cells for tumors with dishesive pattern | >10 groups (≥10 cells each group) or >100 tumor cells for tumors with dishesive pattern |
by application of 5 mL of negative pressure for 5–10 seconds, during which at least five additional actuations were performed. The fanning technique was used whenever feasible.

Following the needle withdrawal, obtained tissue was processed on site, and touch preparations or crush smears were prepared. Slides were air-dried and stained with modified Wright Giemsa staining for ROSE. If the first smear was negative, the endosonographer performed additional smears until it turned positive, or it was felt that further sampling would not increase the smears yield. All additional visible cores were placed in a cellular preservative (CytoRich Red; Becton Dickinson, Franklin Lakes, NJ, USA) for cell block preparation. Visual inspection of the cell block was performed to ensure that there was sufficient material in terms of the cloudiness of the fluid, and not necessarily relying on the presence of defined whitish cores (some mini-cores can be hard to detect visually and turn the fluid turbid) (Fig. 7).

Follow-up

Procedure-related outcomes were collected prospectively. Our endoscopy unit routinely telephones all patients within 48–72 hours after the procedures to assess for short-term adverse events. Adverse events were recorded according to the

Fig. 2. Fine needle biopsy specimen obtained from a pancreatic head mass. This lesion was sampled using standard fine needle aspiration technique twice prior to referral to our hospital with insufficient cytology. Rare malignant cells seen infiltrating the fibrous stroma. This was assigned a score of 1 on the cell block. The final diagnosis was pancreatic adenocarcinoma (Hematoxylin and eosin, ×100).

Fig. 3. Fine needle biopsy specimen obtained from a pancreatic body mass. This was assigned a score of 2 on cell block. The final diagnosis was pancreatic adenocarcinoma (Hematoxylin and eosin, ×20).

Fig. 4. Fine needle biopsy specimen obtained from a mediastinal mass. This was assigned a score of 3 on cell block. The final diagnosis was poorly differentiated lung cancer (Hematoxylin and eosin, ×40).

Fig. 5. Fine needle aspiration specimen obtained from a liver mass. This was assigned a score of 3 on cell block. The final diagnosis was metastatic esophageal cancer (Hematoxylin and eosin, ×40).
Continuous variables were reported as mean±standard deviation. Categorical variables were presented as numbers and percentages. Sensitivity, specificity, positive predictive value, negative predictive value, and accuracy were calculated. To determine the differences between the groups, the Student's t-test was used for continuous variables and chi-squared or Fisher's exact test was used for categorical variables. The sample size was not estimated a priori as this is the first study conducted with this novel needle. We chose 50 samples in the test group and a similar number of controls as a convenience sample. Two-sided p-values <0.05 were considered statistically significant. Statistical analysis was performed using SPSS 21.0 software for Windows (SPSS Inc., Chicago, IL, USA).

RESULTS

Fifty patients (28 males) underwent EUS-FNB for 51 solid lesions between September 2016 and March 2017 at Indiana University Hospital and comprised the Study group. Lesion and sampling results characteristics are summarized in Table 2. A neoplastic diagnosis was confirmed in the vast majority of patients (48/50; 96%) based on cytology. Resection surgical pathology confirmed the cytological diagnosis in 12 patients (24%).

Fifty FNA specimens from solid lesions were pulled from our cytology archives and were independently reviewed (Control group). Mean cell block histology scores were significantly higher in the FNB group (p=0.046) despite an overall lower mean number of passes compared to the FNA group (p<0.001, Table 2). The overall diagnostic yield of FNB was 96% compared to 88% for the FNA group, but the difference did not reach statistical significance.

The degree of tumor differentiation for primary tumors, origin of metastatic lesions, and mitotic index in neuroendocrine and stromal tumors (n=18) was adequately assessed in
all lesions based on cell block studies in the FNB group (Table 3). The performance characteristics (sensitivity, specificity, and accuracy) of the two sampling methods assessed in this study (FNB vs. FNA) are summarized in Table 4.

### Adverse events
All adverse events were classified as mild, and were recognized during or immediately after the procedure. Two patients (4%) developed post-procedural abdominal pain.

### Table 2. Lesion and Sampling Results Characteristics of the Fine Needle Biopsy and Fine Needle Aspiration Cohorts

| Lesion and sampling characteristics | FNB (n=51) | FNA (n=50) | p-value |
|------------------------------------|-----------|-----------|---------|
| Mean size (maximal dimension, cm)  | 2.98      | 2.90      | NS      |
| Location: Pancreatic (n)           | 23        | 20        | NS      |
| Non pancreatic (n)                 | 28        | 30        |         |
| Cytology scores of smears (mean)   | 2.75      | 2.86      | NS      |
| Number of passes for smears (mean) | 2.07      | 2.44      | 0.056   |
| Cell block histology score (mean)  | 2.32      | 2.02      | 0.046   |
| Number of additional passes for cell block (mean) | 0.74 | 1.68 | <0.001 |
| Total number of passes (mean)      | 2.88      | 3.82      | <0.001  |
| Overall diagnostic yield           | 96%       | 88%       | NS      |

FNB, fine needle biopsy; FNA, fine needle aspiration; NS, non significant.

### Table 3. Confirmed Clinical Outcomes of All 50 Patients Included in the Study

| Site of FNB (n=51) | Final reference diagnosis (n=50) |
|--------------------|----------------------------------|
| Pancreas (n=21)    | Adenocarcinoma (n=9); Chronic pancreatitis (n=1); NET (n=1) |
| Head and Uncinate (n=11) | Adenocarcinoma (n=4); Chronic pancreatitis (n=1); Well differentiated NET (n=1) |
| Body (n=5)         | Adenocarcinoma (n=4); Chronic pancreatitis (n=1); Well differentiated NET (n=1) |
| Tail (n=5)         | Adenocarcinoma (n=4); Well differentiated NET (n=1) |
| SET (n=8)          | Leiomyma (n=3); Metastatic adenocarcinoma (n=4); Large B cell lymphoma (n=1); Metastatic melanoma (n=1) |
| Esophageal (n=3)   | GIST (n=4); GIST (n=1) |
| Gastric (n=4)      | GIST (n=4); GIST (n=1) |
| Duodenum (n=1)     | Metastatic adenocarcinoma (n=4); Metastatic NET (n=1); Metastatic melanoma (n=1) |
| Lymph nodes (n=6)  | Cholangiocarcinoma (n=3); Poorly differentiated carcinoma (n=1); Squamous cell carcinoma (n=1) |
| Liver lesions (n=5) | Metastatic adenocarcinoma (n=3); Metastatic NET (n=1); Metastatic melanoma (n=1) |
| Distal bile duct masses (n=3) | Local recurrence of previously resected pancreatic adenocarcinoma (n=2) |
| Mediastinal masses (n=2) | Cholangiocarcinoma (n=3) |
| Retropertitoneal masses (n=2) | Metastatic lung adenocarcinoma |
| Miscellaneous (n=4) | Adenocarcinoma (linitis plastica); Local recurrence of previously resected rectal adenocarcinoma; Teratoma |
| Gastric wall thickening (n=1) | Adenocarcinoma (linitis plastica) |
| Perirectal mass (n=1) | Local recurrence of previously resected rectal adenocarcinoma |
| Left adrenal mass (n=1) | Metastatic lung adenocarcinoma |
| Pelvic mass (n=1) | Teratoma |

FNB, fine needle biopsy; NET, neuroendocrine tumor; SET, subepithelial tumor; GIST, gastrointestinal stromal tumor.

* A single patient with metastatic melanoma underwent sampling of liver lesion and a lymph node, both were metastatic.

* Four patients diagnosed eventually with adenocarcinoma based on FNB had previous insufficient fine needle aspiration sampling at other institutions.

### Table 4. Performance Characteristics of the 2 Sampling Methods Assessed in This Study (FNB vs. FNA)

|            | Sensitivity (95% CI) | Specificity (95% CI) | Accuracy |
|------------|----------------------|----------------------|----------|
| FNB group  | 95.9% (86.02% to 99.50%) | 100% (15.81% to 100.00%) | 96%      |
| FNA group  | 87.23% (74.26% to 95.17%) | 100% (29.24% to 100.00%) | 88%      |

FNB, fine needle biopsy; FNA, fine needle aspiration; CI, confidence interval.
pain after pancreas mass FNB (confirmed adenocarcinoma on cytology). This required extended observation in the recovery room for 3 hours in one patient and 23-hour admission to the hospital for observation and pain management in another patient.

In two cases, a hematoma developed outside the gastric and duodenal walls during FNB of the pancreatic body (Fig. 8) and head masses, respectively. Both patients underwent 4 passes each. The hematomas were monitored for 10 minutes and appeared stable on EUS. This did not result in clinically apparent adverse outcomes and both patients were discharged home after 90 minutes of observation without the need for further interventions.

No other adverse event was reported during or immediately after the procedure and up to 72 hours of follow-up.

DISCUSSION

EUS-FNB is increasingly performed to sample solid lesions, but is currently reserved to cases where FNA cytology fails or when tissue architecture is essential for the diagnosis. The performance of earlier Trucut core biopsy devices (QuickCore; Cook Medical, Limerick, Ireland) was suboptimal due to mechanical failures of its spring loading mechanism. A newer needle with unique reversed bevel design was released in 2011 (ProCore; Cook Medical) and remains the most widely studied needle. Superior histologic yield has been demonstrated in one recent randomized study comparing the ProCore to the QuickCore needles. However, multiple studies failed to demonstrate the superiority of this needle for sampling adequacy or diagnostic accuracy over conventional EUS-FNA needles. A recent meta-analysis of nine studies including 576 patients demonstrated no significant difference in diagnostic adequacy (75% vs. 89%), diagnostic accuracy (85% vs. 86%), or rate of histological core specimen acquisition (78% vs. 77%) between the ProCore and standard FNA needles. More recently, the performance of the ProCore needle was compared to that of a newer core biopsy device with a novel fork-tip with opposing bevel design and multiple cutting surfaces (Sharkcore™; Medtronic, Minneapolis, MN, USA). In this study, the opposing bevel needle provided significantly higher sensitivity (71.1% vs. 90.1%) and overall accuracy (74% vs. 92%) for discriminating malignant from benign solid pancreatic masses.

Our study aimed to assess another new core biopsy device with three cutting surfaces and a Franseen shaped tip. We demonstrated that this device provides a histologically superior sample with fewer passes compared to conventional FNA needles of the same size. The superiority of FNB samples have been recently reported with the fork-tip device. The current study agrees with the results of those two studies from two perspectives: first, the novel needle tip design appears to be a significant factor enhancing tissue acquisition; second, this same tip design could carry a slightly higher risk for tissue trauma, resulting in bleeding and post-procedural pain. In our series, as opposed to the results reported by Rodrigues-Pinto et al., bleeding was self-limited in all cases with no further interventions were needed.

We have also demonstrated that the smears prepared from such needles do not appear to be inferior to FNA smears for rapid review purposes. This is likely due to the rather fragmented nature of the cores accrued, allowing sheets of cells to smear in thin layers to allow cytological interpretation whether on alcohol fixed or on-site dipped smears.

A recent preliminary descriptive report on this needle demonstrated that diagnostic material for ROSE and histology could be obtained in 97% of patients. Our study demonstrated a higher diagnostic accuracy in the FNB group compared to FNA group, although the difference did not reach statistical significance. This may be related to the fact that we applied strict cytological criteria to assess smears and cell block specimens which relied on manual counting of the smears and cell blocks. Since the majority of sampled lesions were solid pancreatic masses, we drew the conclusions that in such cases, three passes are sufficient for establishing the diagnosis with over 90% diagnostic adequacy.

The yield of EUS-guided tissue acquisition is influenced by many factors. In the current study, we attempted to standardize many of the variables known to impact the outcomes of sampling. For example, a standard FNB sampling technique was performed by the same endosonographer, using the same needle gauge and the same standard cytological processing techniques. In addition, the vast majority of lesions in our FNB cohort were unequivocally neoplastic, and the few with...
negative cytology had sufficient follow-up to rule out a malignant outcome. Finally, our cytopathology team has devised a scoring system that minimizes subjectivity in assessing cellularity of smears and cell block material. This system relies on a manual count of the number of diagnostic cells (most commonly tumors) and not on the presence of any nucleated cell. Similar scoring systems have been developed with variable cell count brackets.\textsuperscript{11}

The diagnostic yield of FNA in pancreatic masses remains high, with a recent meta-analysis showing an EUS-FNA pooled sensitivity and specificity for diagnosing solid pancreatic neoplasms of 85\% and 98\%, respectively.\textsuperscript{22} Despite that, a recent cost-effectiveness analysis from the perspective of a third-party payer showed that FNB was more cost-effective than FNA in sampling both pancreatic and non-pancreatic masses.\textsuperscript{23} Our study demonstrates the utility of routine use of a core biopsy device for solid lesion sampling during EUS.

In addition, this technique appeared to be safe. All adverse events were classified as mild, and were recognized during or immediately after the procedure. Two patients developed post-procedural abdominal pain, and two patients developed hematoma outside the gastric and duodenal walls. No other adverse events were reported during or immediately after the procedure and up to 72 hours of follow-up.

We recognize several limitations to our study. This was not a prospective randomized controlled study. Different solid lesions from different organs were prospectively sampled using the FNB needle only; thus, a direct comparison of the performance of FNB to FNA needles for the same lesion(s) could not be assessed. An important part of the assessment of this new tissue acquisition device was to examine its performance for the purposes of ROSE. As has been endorsed by several studies and key opinion leaders, we continue to rely heavily on ROSE and hence we describe the utility of FNB devices in the context of ROSE. The natural next step is to assess its performance without ROSE, but this study was an early clinical utility study. In our cohort we did not have sufficient number of patients to comment on the performance of this device in specific situations like lymphoma, gastrointestinal stromal tumor, or autoimmune pancreatitis. Finally, we have chosen smears and cell block technique instead of histological exam (typically submitted in formalin). Moreover, our sample size was relatively small, since this is the first study to assess this specific needle platform in a controlled fashion. This could have been underpowered to detect true differences between the FNB and FNA groups in diagnostic accuracy. Despite that, several endpoints, including the total number of passes and the number of passes dedicated to cell block, met the statistical significance. Finally, we confirmed the results of previous reports showing that FNB devices are overall associated with fewer number of passes.\textsuperscript{13} Additionally, we acknowledge that there may be a learning curve to using a new needle platform that was “mastered” during the span of the study.

In conclusion, the new Franseen tip biopsy needle enables sufficient acquisition of histological-quality samples under EUS-guidance. Due to the availability of various sizes and the overall safety of such devices, an FNB exclusive approach to sampling all solid lesions appears feasible. Further studies are needed to characterize the performance of this device in lesions of various origins and locations.

\textbf{Note} 

An abstract of this study was accepted for poster presentation at the Digestive Diseases Week, 2017, Washington, DC and at 2017 American College of Gastroenterology Annual Meeting in Orlando, FL, USA.

\textbf{Conflicts of Interest} 

The authors have no financial conflicts of interest.

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