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***Retrospective Cohort Study***

**Endoscopic ultrasound-guided through-the-needle microforceps biopsy and needle-based confocal laser-endomicroscopy increase detection of potentially malignant pancreatic cystic lesions: A single-center study**

Robles-Medranda C *et al*.Novel EUS techniques in pancreatic cysts

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**Abstract**

BACKGROUND

Currently, there is insufficient data about the accuracy in the diagnosing of pancreatic cystic lesions (PCLs), especially with novel endoscopic techniques such as with direct intracystic micro-forceps biopsy (mFB) and needle-based confocal laser-endomicroscopy (nCLE).

AIM

To compare the accuracy of endoscopic ultrasound (EUS) and associated techniques for the detection of potentially malignant PCLs: EUS-guided fine needle aspiration (EUS-FNA), contrast-enhanced EUS (CE-EUS), EUS-guided fiberoptic probe cystoscopy (cystoscopy), mFB, and nCLE.

METHODS

This was a single-center, retrospective study. We identified patients who had undergone EUS, with or without additional diagnostic techniques, and had been diagnosed with PCLs. We determined agreement among malignancy after 24-mo follow-up findings with detection of potentially malignant PCLs *via* the EUS-guided techniques and/or EUS-guided biopsy when available (EUS malignancy detection).

RESULTS

A total of 129 patients were included, with EUS performed alone in 47/129. In 82/129 patients, EUS procedures were performed with additional EUS-FNA (21/82), CE-EUS (20/82), cystoscopy (27/82), mFB (36/82), nCLE (44/82). Agreement between EUS malignancy detection and the 24-mo follow-up findings was higher when associated with additional diagnostic techniques than EUS alone [62/82 (75.6%) *vs* 8/47 (17%); OR 4.35, 95%CI: 2.70-7.37; *P <* 0.001]. The highest malignancy detection accuracy was reached when nCLE and direct intracystic mFB were both performed, with a sensitivity, specificity, positive predictive value, negative predictive value and observed agreement of 100%, 89.4%, 77.8%, 100% and 92.3%, respectively (*P <* 0.001 compared with EUS-alone).

CONCLUSION

The combined use of EUS-guided mFB and nCLE improves detection of potentially malignant PCLs compared with EUS-alone, EUS-FNA, CE-EUS or cystoscopy.

**Key Words:** Pancreatic cysts; Endoscopic ultrasound-guided fine-needle aspiration; Confocal microscopy; Image-guided biopsy

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**Core Tip:** This retrospective study compared the accuracy of endoscopic ultrasound (EUS) and associated techniques such as EUS-guided fine needle aspiration (EUS-FNA), contrast-enhanced EUS (CE-EUS), EUS-guided fiberoptic probe cystoscopy (cystoscopy), EUS-guided direct intracystic micro-forceps biopsy (mFB), and EUS-guided needle-based confocal laser-endomicroscopy (nCLE) for the detection of potentially malignant pancreatic cystic lesions (PCLs) in 129 patients. Patients were allocated to three cohorts: those evaluated *via* EUS alone; *via* EUS-FNA, CE-EUS and/or cystoscopy; and with mFB plus nCLE. We observed that combining EUS, mFB, and nCLE had a statistically significant improved detection of potentially malignant PCLs compared to any of the evaluated techniques alone.

**INTRODUCTION**

The incidence of pancreatic cystic lesions (PCLs) is rising mainly in elderly patients[1]. Therefore, early detection of potentially malignant PCLs increases the possibility of a curative approach. Current American Gastroenterological Association guideline recommends magnetic resonance imaging (MRI) or magnetic resonance cholangiopancreatography (MRCP) to assess PCLs[2]. For the same purpose, the revised Fukuoka guideline recommend computerized tomography (CT), MRI or MRCP, keeping endoscopic ultrasound guided fine-needle aspiration (EUS-FNA) for intraductal papillary mucinous neoplasm (IPMN) evaluation[3]. Nevertheless, both guidelines showed an unsatisfactory pooled sensitivity for malignant PCLs of 64% and 59%, respectively[4].

EUS is the most sensitive diagnostic method for detecting potentially malignant pancreatic lesions with an 88.5% sensitivity; yet it holds a 52.9% specificity and a higher inter-observer variability. Thus, EUS alone has very low diagnosability capacity[5-7]. Similarly, a considerable number of PCLs cannot be characterized by CT, MRI or MRCP alone[8,9]. EUS-guided diagnostics techniques increase EUS accuracy for differentiating PCLs, namely: (1) EUS-FNA; (2) contrast-enhanced EUS (CE-EUS); (3) fiberoptic probe cystoscopy (cystoscopy); (4) EUS-guided through-the-needle direct intracystic micro forceps biopsy (mFB); and (5) EUS-guided confocal laser endomicroscopy (nCLE)[9].

EUS-FNA allows biopsy of suspicious lesions and cytological and biochemical cystic fluid analysis[7]. Whereas, CE-EUS help to differentiate between solid *vs* PCLs, by detecting enhanced septa or nodules present within cystic lesions[10]. Through-the-needle fiberoptic probe cystoscopy requires a 19-gauge needle guided by EUS to locate and enter the PCL. Then, the preloaded fiberoptic probe is advanced, allowing visualization of the cyst content as cystic wall features[11]. The microforceps device samples tissue from the cyst’s wall, septations, and/or mural nodules and thus increase cellular yield[12]. Furthermore, nCLE characterizes PCLs type by imaging the intact cyst architecture, targeting abnormal areas and reducing unnecessary sampling of surrounding tissue, with a diagnostic accuracy of 80% to 95%[8].

Given the poor prognosis of malignant pancreatic lesions, determining the best diagnostic approach for early detection of potential malignancy among the variety of newly available EUS-related technology is essential. Therefore, we aimed to compare the accuracy of EUS for detection of potentially malignant PCLs when it is performed alone, EUS-FNA, CE-EUS or cystoscopy and associated with novel EUS-related techniques: mFB and nCLE. We hypothesize that EUS-guided through-the-needle mFB and nCLE may increase malignancy detection during EUS assessment of pancreatic cysts.

**MATERIALS AND METHODS**

***Study design***

The following is an observational, analytic, longitudinal, retrospective cohort and single-center study performed at the Instituto Ecuatoriano de Enfermedades Digestivas (IECED), a tertiary center in Ecuador. The study protocol and informed consent documents were approved by the institutional review board, and the study was conducted in accordance with the Declaration of Helsinki. Selected patients signed corresponding informed written consent for healthcare purposes.

***Population selection***

Records from patients older than 18 years of age who underwent EUS at IECED from January 2013 to March 2018 were extracted from the institutional database. Cases with non-pancreatic lesions were excluded. Patients were allocated to three cohorts: (1) patients who had been evaluated *via* EUS alone; (2) patients who had been evaluated with EUS-FNA, CE-EUS and/or cystoscopy; and (3) those evaluated with novel EUS-related techniques: mFB and nCLE.

***Endoscopic techniques malignancy criterion for pancreatic cystic lesions***

Due to sparse cellularity of acquired specimens, several complementary clinical, radiological, and imaging techniques are required to achieve PCLs definitive diagnosis. PCLs with potential to progress to malignancy mainly IPMN, mucinous cystic neoplasms (MCN), and neuroendocrine tumors (c-NET) with cystic degeneration. Identifying malignancy features for these lesions with EUS, CE-EUS, cystoscopy, nCLE, FNA, and mFB include the following:

**EUS:** Presenting two out of the three following characteristics was considered as increased risk for malignancy criteria: main pancreatic duct dilation between 5-9 mm (10 mm high risk stigmata for malignancy), PCLs size > 3 cm, and mural nodules presence[3,13].

**CE-EUS:** A thick/hyper-enhancing wall/septum, enhancing solid component within a cyst, or an enhancing mural nodule favors malignancy criterion. Furthermore, there is a radiological correlation between pancreatic duct communication and IPMN diagnosis, but not MCN. Also, main duct type IPMNs hold a higher risk of malignancy transformation than branch duct type IPMNs (up to 68% *vs* 22%, respectively). MCN may show peripheral calcifications within multilocular septate lesions[3,14].

**Cystoscopy:** Cloudy fluid and a smooth cyst wall identify MCN, while finger-like projections and a mucin cloud are perceived with IPMN through single-operator cholangioscopy (SOC)[11,14].

**nCLE:** Prone to malignancy lesions may depict epithelial or vascular patterns in nCLE[5,8,11,13,15]. nCLE Epithelial patterns: MCN show epithelial borders with a flat mosaic appearance (single or multiple layers of epithelial bands). IPMN exhibit dark rings and papillary projections. c-NET portray a trabecular pattern (fibrous bands separating cells nests). nCLE Vascular patterns: MCN, IPMN and cystic-NET may show a branched pattern; IPMN and MCN may also display a rope-ladder pattern[5].

EUS-FNA and EUS-mFB are resources for tissue sample extraction. For these techniques, cytology should be assessed in the context of radiological and clinical findings[3,11,14]. Low and high-grade IPMN dysplasia should be distinguished as the latter may easily become invasive. Low-grade IPMN: may resemble normal gastric epithelium. High-grade IPMN may show a cell size ≤ 12 μm, hypo/hyperchromasia, background necrosis, nuclear irregularity, large single vacuolated cells, and increased nuclear to cytoplasmic ratio[14].

IPMNs histologic examinations exhibit four possible morphologies: gastric (columnar cells lining papillae with basally located nuclei rich in apical mucin), intestinal (similar morphology to colonic villous adenomas with cigar shaped nuclei and variable apical mucin amount), pancreaticobiliary (more complex papillae composed of rounded nuclei cuboidal cells with some prominent nucleoli), and oncocytic (complex papillae lined with round cells with granular eosinophilic cytoplasm and prominent central nucleoli)[3,14].

MCNs also display low and high-grade dysplasia features. While bland mucin-containing epithelium honeycomb sheets are seen with low-grade MCNs, a complex papillary structure with smooth nuclear contour mucin-containing cells, inconspicuous nucleoli, and fine chromatin is found in high-grade MCNs. On histologic examination, MCNs show focally flat o cuboidal lining and tall mucin-containing epithelium, with a densely ovarian-type stroma wall that positively stains for progesterone/estrogen receptors, calretinin, and inhibin[3,14].

C-NET aspirate display classic endocrine morphology (pseudorosettes, isolated, and loosely cohesive groups of round/polygonal cells with finely stippled chromatin round nucleus)[5,11,14,15]. Immunostains (chromogranin, CD10, vimectin, and β-catenin cytoplasmic expression) provide a definitive diagnosis[14].

***Endoscopic techniques methods***

Three experienced endosonographers (C.R-M., J.O., R.V.) performed all EUS evaluations, under general anesthesia with patients in the supine position and use of antibiotic prophylaxis. EUS procedures were performed with a linear-array video echoendoscope (EG-3870 UTK, Pentax Medical, Montalve, NJ, United States) attached to an ultrasound console (HI VISION Avius®, Hitachi Medical Systems, Steinhaus, Switzerland). Indication of EUS-related techniques was based on endosonographers discretion. Although more techniques are available to perform on larger cysts (> 3 cm).

**Endoscopic ultrasound fine needle aspiration:** EUS-FNA was performed with a 19-gauge needle (Expect™ Slimline, Boston Scientific, Malborough, United States) (Figure 1A). The cystic fluid was examined for tumor markers (amylase, lipase, carcinoembryonic antigen levels).

**Contrast enhanced endoscopic ultrasound:** To display cystic wall and nodule vascularization, 4.8 mL of SonoVue® (Braccio, Milan, Italy) was used for CE-EUS. Cystic wall and nodule vascularization were defined as visible contrast enhancer bubble movement within the cystic wall, septum, and nodules (Figure 1B), and were referred for further diagnosis with EUS-FNA.

**Cystoscopy:** Examinations were performed by using a linear-array video echoendoscope attached to an ultrasound console, as previously described. A SOC fiber optic probe (Legacy SpyGlass® fiber optic, Boston Scientific, Marlborough, United States) was inserted through the 19-gauge needle into the cystic cavity to observe the intracystic wall and contents (Figure 1C).

**EUS-guided through-the-needle direct intracystic micro forceps biopsy:** The target lesion was identified under EUS and punctured with a 19-gauge FNA needle. With the needle inside the lesion, the stylet was removed, and the micro forceps (Moray™ micro forceps, STERIS, Mentor, United States) were inserted through the needle for tissue sampling. Two to three bites of biopsy specimens were taken with each pass of the micro forceps. The tissue acquisition was visually confirmed and directly placed on formalin containers for pathologic evaluation.

**EUS-****guided confocal laser endomicroscopy:** After EUS examination, patients were intravenously injected with 5 mL of 10% fluorescein (BioGlo®, Sofar Productos, Bogota, Colombia) 2 to 3 min before nCLE imaging. CLE was performed using the AQ-Flex nCLE miniprobe (Cellvizio, Mauna Kea Technologies, Paris, France). The probe was advanced through the locking device into the 19-gauge needle. The preloaded needle was advanced under EUS guidance into the PCL. The tip of the nCLE probe was placed in contact with the intracystic epithelium, and intracystic endomicroscopic images were captured (Video 1 and Video 2). After image acquisition, the nCLE probe was withdrawn, and the PCL was aspirated.

***Data abstraction***

Demographic, clinic, endoscopic and histopathological and 24-mo follow-up data were obtained from the institutional database and phone calls when necessary. The study endpoint was to determine agreement between detection of potentially malignant in PCLs (EUS malignancy detection) and malignancy after 24-mo follow-up. EUS malignancy detection was defined based on procedure findings (EUS-alone, CE-EUS, cystoscopy and/or nCLE) reported on endoscopic records, as well as EUS-FNA and/or EUS-mFB aquired biopsy results when available. PCLs were classified as malignant (MCN, IPMN and c-NET) according to Fukuoka criteria. This data was recovered by two endoscopists (C.R.M. and H.P-L.). Malignancy after 24-mo follow-up was based on clinical outcomes, endoscopic surveillance, or surgical specimen histopathology when available. This data was recovered by two general practitioners (R.O. and J.B-B.) and a general surgeon (D.C-L.) who were blinded to information concerning to EUS malignancy detection.

***Interobserver agreement***

An offline interobserver analysis (IOA) of the EUS criteria (EUS borders, lobularity, wall, microcyst component, diagnosis, and level of confidence) was performed by three endoscopists (J.O., R.V. and J.N.) using a randomly selected EUS image set (*n* = 111 cases) collected by C.R-M.

***Statistical analysis***

**Technical considerations:** Final database was consolidated and encrypted by M.A-M. Data analysis was performed by IECED Institutional Biostatistician (M.P-T.) using R v.4.0 (R Foundation for Statistical Computing, Vienna, Austria). A *P*-value <0.05 was considered statistically significant.

**Sample size calculation:** We considered a 100% specificity of EUS + nCLE for the prediction of potentially malignant PCLs, with a 35% disease prevalence (6/31 mucinous cystic neoplasm and 5/31 IPMNs) for defining the sample size (16). We estimated a sample size of 25 patients for each cohort, with an α and β-error of 5% and 20% respectively, and an 80% statistical power.

**Descriptive analysis:** Numeric variables were described through the mean ± SD or median (minimum-maximun range) in accordance with statistical distribution (Kolmógorov-Smirnov test). Categorical variables were described with frequency (%), and 95%CI when corresponding. Descriptions about techniques combination was summarized on a Venn Diagram (17).

**Inferential analysis:** Observed agreement between EUS malignancy detection and malignancy after 24-mo follow-up was established. The statistical association between EUS alone or EUS with an additional endoscopic technique *vs* the positive observed agreement described above was determined by binary logistic regression [odds ratio (OR)]. A univariate analysis was performed for each individual technique. Those with a significant association were entered into the multivariate analysis. The overall diagnostic accuracy for malignancy detection was determined for each diagnostic procedure which shown significance on multivariate analysis, considering a 24-mo follow-up as gold standard. Overall diagnostic accuracy comprehended calculation of sensitivity, specificity, positive predictive value (PPV), negative predictive value (NPV), positive likelihood ratio, negative likelihood ratio, and observed agreement. For multivariate analysis discrimination, we estimated the corresponding area under the receiver operating characteristics (AUROC) curves and contrasting using the DeLong’s test for two ROC curves. The IOA of the EUS criteria was performed using Fleiss’ kappa score (κ) calculation and interpreted based on Landis and Koch criteria.

**RESULTS**

***Patient selection***

A total of 2812 patients were referred to our unit for diagnostic EUS along study period. Of these, 856 had pancreatic lesions, of which 129 patients with PCLs were included for analysis (*n* = 129) (Figure 2).

***Baseline characteristics***

The median age of the 129 patients with PCLs was 69 years, and 69.8% patients were female. The most frequent pancreatic cyst location was the head of the pancreas (35.7%). Younger patients were significantly evaluated with EUS and an additional novel technique (mFB and/or nCLE) in comparison to those evaluated with EUS alone, EUS-FNA, CE-EUS or cystoscopy (*P <* 0.001). Cysts size above 30 mm were reported among patients evaluated with EUS and an additional novel technique (46.3%) compared with general cohort (27.1%; *P* < 0.001). There were no statistically significant differences when comparing gender and PCLs location between patients evaluated with EUS alone and those evaluated with EUS plus additional diagnostic techniques (Table 1).

EUS was performed with an additional diagnostic technique in 82/129 patients: EUS-FNA [21/82 (25.6%)], CE-EUS [20/82 (24.4%)], cystoscopy [27/82 (32.9%)], mFB [36/82 (43.9%)], and nCLE [44/82 (53.7%)]. More than one diagnostic technique was performed in a sample proportion (Figure 3). A 100% technical success was reached, with no documented adverse events for any of the performed procedures.

According to the PCLs EUS findings and guided biopsy when available (*n* = 53), potentially malignant PCLs were detected in 81/129 (62.8%) patients, and the most frequent lesion among this group was IPMN [70/129 (54.3%)]. In the nonmalignant group [48/129 (37.2%)], 46 cases were serous cystadenomas (Table 1). Observed agreement between EUS malignancy detection and malignancy after 24-mo follow-up was higher in patients evaluated with EUS plus at least one additional novel technique (mFB and/or nCLE), followed by EUS-FNA, CE-EUS and or cystoscopy; than in patients evaluated with EUS alone [42/55 (80.0%) *vs* 18/27 (66.7%) *vs* 8/47 (17%), respectively; OR 4.35, 95%CI: 2.70-7.37; *P <* 0.001].

***Univariable and multivariable analysis***

Independently, there was a positive statistical association and observed agreement for EUS malignancy detection with cystoscopy, mFB or nCLE, and 24-mo follow-up. EUS-FNA and CE-EUS exhibited a positive but nonsignificant association; whereas EUS alone only presented a negative significantly association [OR 0.066 (0.025-0.157; *P <* 0.001)] when considering the agreement between EUS malignancy detection and malignancy after 24-mo follow-up as an outcome.

Through multivariate analysis, we confirmed that malignancy detection was significantly more accurate with nCLE [OR 8.441 (2.698-33.081; *P <* 0.001)] and mFB [OR 3.425 (1.104-11.682; *P =* 0.038)] than cystoscopy [OR 0.622 (0.125-2.813; *P =* 0.541)] (Table 2).

***Diagnostic accuracy for determining malignancy***

EUS alone was performed in 47 cases and had a sensitivity, specificity, PPV, and NPV of 100%, 3%, 15%, and 100%, respectively. EUS-FNA, CE-EUS, and/or cystoscopy was performed in 28 cases and had a sensitivity, specificity, PPV, and NPV of 91%, 47% 53% and 89%, respectively. EUS with nCLE and mFB yielded similar results for sensitivity (89% *vs* 88%), specificity (86% *vs* 82%), PPV (62% *vs* 58%) and NPV (97% *vs* 96%). When the three techniques were simultaneously performed (EUS with nCLE and mFB, *n* = 26), the diagnostic accuracy analysis showed that the sensitivity, specificity, PPV, and NPV were 100%, 89%, 78%, and 100%, respectively. MCC identified a good correlation between EUS malignancy detection and malignancy after the 24-mo follow-up through different techniques. Nonetheless, EUS paired with nCLE and mFB showed the highest agreement (MCC = 0.83) (Table 3).

Detection of potentially malignant PCLs using EUS alone reached a 51.3% AUROC (*P =* 0.3599; moderate agreement). Meanwhile, EUS-guided mFB, nCLE or/and mFB reached an 87.3% AUROC (*P <* 0.001), 84.8% (*P <* 0.001) and 94.7% (*P <* 0.001), respectively. In addition, nCLE reached a greater AUROC in comparison to EUS alone (*P <* 0.001) (Figure 4A). Moreover, a significantly higher AUROC was described for combined EUS-guided nCLE and mFB in comparison to EUS-FNA/CE-EUS/cystoscopy (94.7% *vs* 69%, *P* = 0.044) (Figure 4B).

***Interobserver agreement***

In the secondary IOA performed by three experienced endoscopists, the κ values in EUS borders, lobularity, wall, microcyst component, diagnosis, and level of confidence were as follows: 0.12 (poor agreement), 0.08 (poor agreement), 0.04 (poor agreement), 0.29 (fair agreement), 0.21 (fair agreement), and 0.06 (poor agreement) respectively.

**DISCUSSION**

Various clinically-available advanced EUS-guided diagnostic techniques have improved the accuracy of malignancy detection among PCLs; however, these techniques are not referenced in current guidelines, with unsatisfactory diagnostic accuracy in the risk stratification of potentially malignant PCLs[4].

To provide guidance on the relative accuracy and effectiveness of these new EUS-related techniques, we compared various additional endoscopic techniques during the EUS evaluation of PCLs. We evaluated the accuracy of EUS alone with more recent EUS-related techniques, namely EUS-FNA, cystoscopy, nCLE, mFB, and CE-EUS and found that the highest level of malignancy detection can be achieved when EUS is combined with both nCLE and direct intracystic mFB.

An increasing number of PCLs have been identified due to the growing use of complementary diagnostic techniques, such as CT and MRI; moreover, the malignancy potential of PCLs vary, and current diagnostic techniques cannot characterize the lesions with precision by their self[18-20]. Due to the malignancy potential, patients with pancreatic neoplasms are recommended to undergo resection therapy; however, for patients with a high risk of postsurgical complications, preoperative determination of malignancy is critical for management guidance.

In our study, EUS alone had a low agreement in comparison to the 24-mo follow-up. Also, in an offline interobserver agreement between three endosonographers, endoscopic criteria showed low agreement between operators, as previously described. Therefore, EUS itself should be complemented with additional endoscopic techniques for a more accurate detection of malignancy in PCLs.

Wang *et al*[21] demonstrated that EUS-FNA can accurately confirm the presence of malignancy but does not perform well at excluding malignant or premalignant pancreatic lesions. This procedure achieved a pooled sensitivity and specificity of 51%, 94%, respectively, for differentiating malignant lesions. In our study, which included 21/129 patients with pancreatic lesions for whom FNA was performed, we found that EUS-FNA did not achieve statistical significance in detecting malignancy with a modest agreement with the 24-mo follow-up; however, this may be due a limited number of cases in our cohort.

The DETECT trial revealed that a combination of through-the-needle cystoscopy and nCLE for PCLs under EUS was feasible, with a sensitivity of 90% for cystoscopy in the clinical diagnosis of MCNs*,* an 80%sensitivity for nCLE, and a 100% sensitivity for the combination of both[11]. In our study, we analyzed both techniques (separately and then combined) and obtained similar results – we obtained a sensitivity of 89% for EUS-guided-nCLE and 88% for EUS-guided through-the-needle cystoscopy; however, the sensitivity of EUS-guided nCLE combined with mFB was 78%. Additionally, in our cohort, we had more heterogenic lesions than in the DETECT trial, which was limited to mucinous lesions.

Haghighi *et al*[8] compared the diagnostic accuracy of nCLE and EUS-FNA, where nCLE was found to have a higher accuracy (87.5%), sensitivity (91.7%), and NPV (93.3%). In our cohort, 44/129 patients underwent nCLE, obtaining similar results (an 86.0% accuracy, an 89% sensitivity, and an NPV of 96%). Konda *et al*[22] reviewed 31 PCLs that were examined using nCLE, and showed a high specificity (100%) and PPV (100%); and an overall accuracy of 71%. In our study, we obtained a higher sensitivity (89%), NPV (96%) and accuracy (86%) probably owing to a higher number of cases.

EUS-nCLE and mFB exhibited an 86.4% and an 83.3% agreement for PCLs malignancy detection, probably due to a better *in vivo* cyst component evaluation and guided tissue acquisition. EUS combined with nCLE and mFB reached the highest AUROC (94.7%), in comparison to independent nCLE (87.3%) and mFB (84.8%). We propose that these techniques should be considered for the diagnostic workup of PCLs.

The main limitation of our study lies in its retrospective design and in establishing an agreement of different endoscopic techniques for determining potential malignancy among different types of PCLs. This resulted in a difficulty in the recovery of different size cysts, where the smaller the cyst, the fewer the diagnostic methods at our disposal for use. On the other hand, larger cysts (specially over 30 mm), allowed us to perform a wider array of diagnostic procedures, including novel techniques. Moreover, these novel endoscopic techniques (*i.e,* nCLE), are costly, limiting their widespread use. Furthermore, these tools require training, which increase the procedure’s startup cost. Despite these limitations, we compared these endoscopic techniques in terms of their ability to detect potential malignancy in patients with PCLs, and not only pancreatic lesions, as with other studies. Finally, as this study was designed in the context of PCLs assessment with EUS, to estimate EUS (and eventual used related techniques) diagnosability of malignancy considering a 24-mo follow-up as gold standard, a prospective diagnostic trial to re-analyse histopathological samples of PCLs after discarding malignancy during follow-up may be warranted to further asses the accuracy in diagnosing high-grade dysplasia/adenocarcinoma in non-malignant PCLs (MCN, IPMN) using the studied endoscopic techniques.

**CONCLUSION**

In conclusion, new EUS technologies such as through-the-needle techniques (direct intracystic mFB combined with nCLE), improve malignancy detection in patients with PCLs. However, multicenter, and cost-benefit studies are recommended to validate these findings.

**ARTICLE HIGHLIGHTS**

***Research background***

Pancreatic cystic lesions (PCLs) incidence is rising mainly in elderly patients. Accurate diagnosing and appropriate management of patients with malignant PCLs, have a positive impact in regards of healthcare expenses and in patients’ quality of life.

***Research motivation***

Currently, there is insufficient data about the accuracy in the diagnosing of PCLs, especially with novel endoscopic techniques. Furthermore, the early detection of potentially malignant PCLs, increases the possibility of a curative approach in said patients.

***Research objectives***

Given the poor prognosis of malignant PCLs, attaining early detection, an accurate diagnosis, and determining the best diagnostic approach with newly available endoscopic techniques, was essential to this study.

***Research methods***

This was a retrospective, single-center study. Patients were allocated to three evaluation cohorts: (1) Endoscopic ultrasound (EUS) alone; (2) EUS- fine needle aspiration, contrast-enhanced-EUS and/or EUS-guided fiberoptic probe cystoscopy (cystoscopy); and; (3) EUS-guided direct intracystic micro-forceps biopsy (mFB) and EUS-guided needle-based confocal laser-endomicroscopy (nCLE); and compared the accuracy of these techniques for the detection of potentially malignant PCLs.

***Research results***

We described that pairing EUS, mFB, and nCLE, had a statistically significant improved detection of potentially malignant PCLs compared to any of the evaluated techniques alone. No adverse events were documented, and a 100% technical success rate was achieved.

***Research conclusions***

In our study, EUS-guided mFB combined with nCLE, improve malignancy detection in patients with PCLs.

***Research perspectives***

To define formal diagnostic and therapeutical guidelines, we encourage researchers to conduct long-term follow-up randomized multicenter and cost-benefit studies, comparing newly available endoscopic techniques for the assessment of PCLs.

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**Footnotes**

**Institutional review board statement:** The study was approved by the Institutional Review Board of Instituto Ecuatoriano de Enfermedades Digestivas.

**Informed consent statement:** All study participants, and their legal guardians, provided informed written consent prior to study enrolment.

**Conflict-of-interest statement:** Robles-Medranda C reports other from Pentax Medical, other from Boston Scientific, other from Steris, other from Medtronic, other from Motus, other from Micro-tech, other from G-Tech Medical Supply, other from CREO Medical, other from Mdconsgroup, outside the submitted work; The other authors declare no conflicts of interest.

**Data sharing statement:** The data that support the findings of this study are openly available by contacting the corresponding author.

**STROBE statement:** The authors have read the STROBE Statement-checklist of items, and the manuscript was prepared and revised according to the STROBE Statement-checklist of items.

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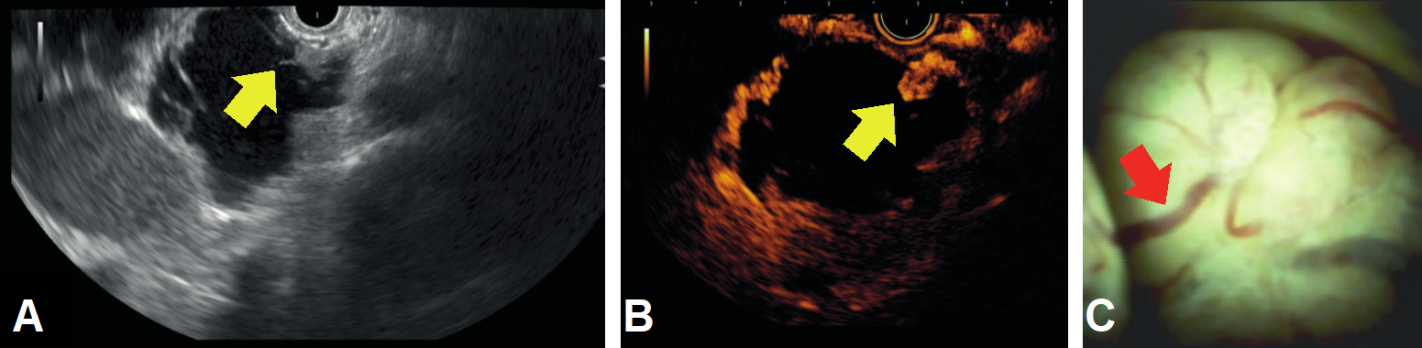
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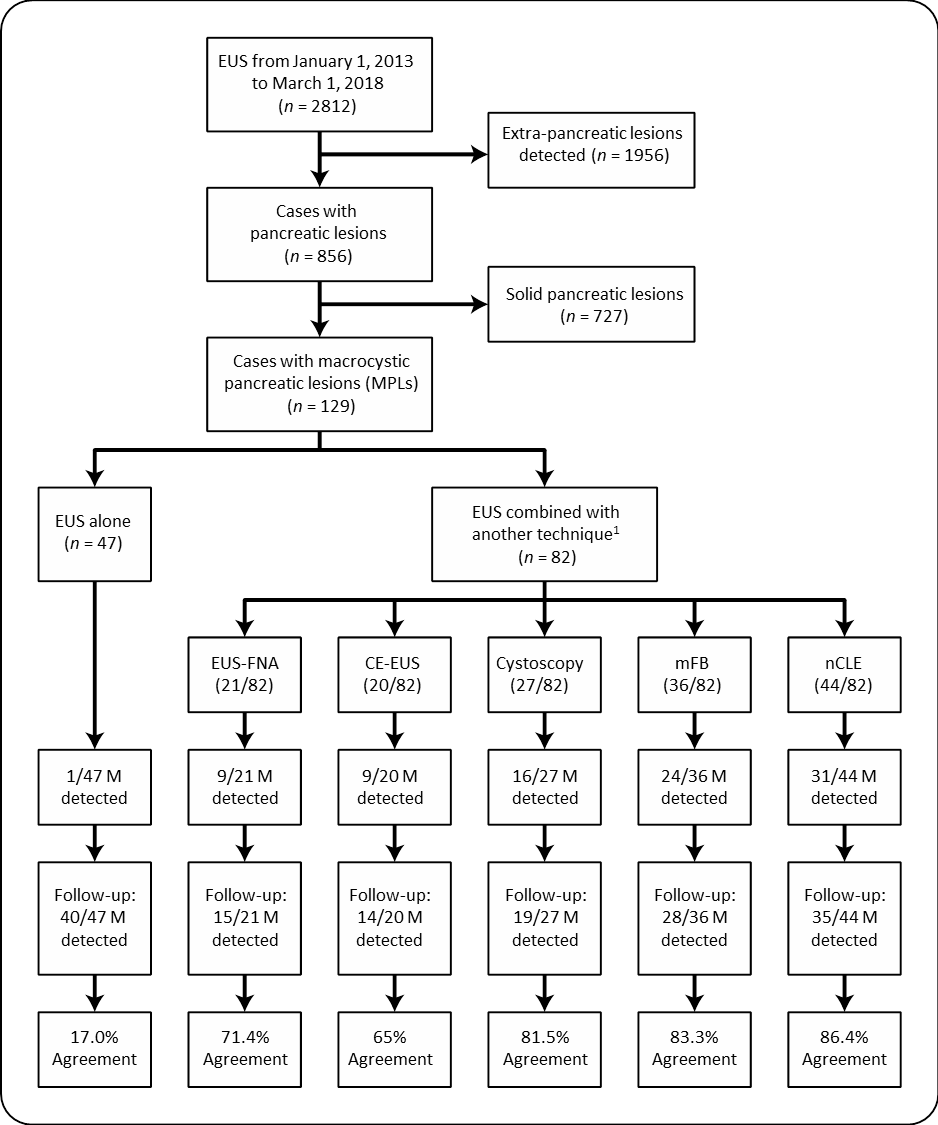
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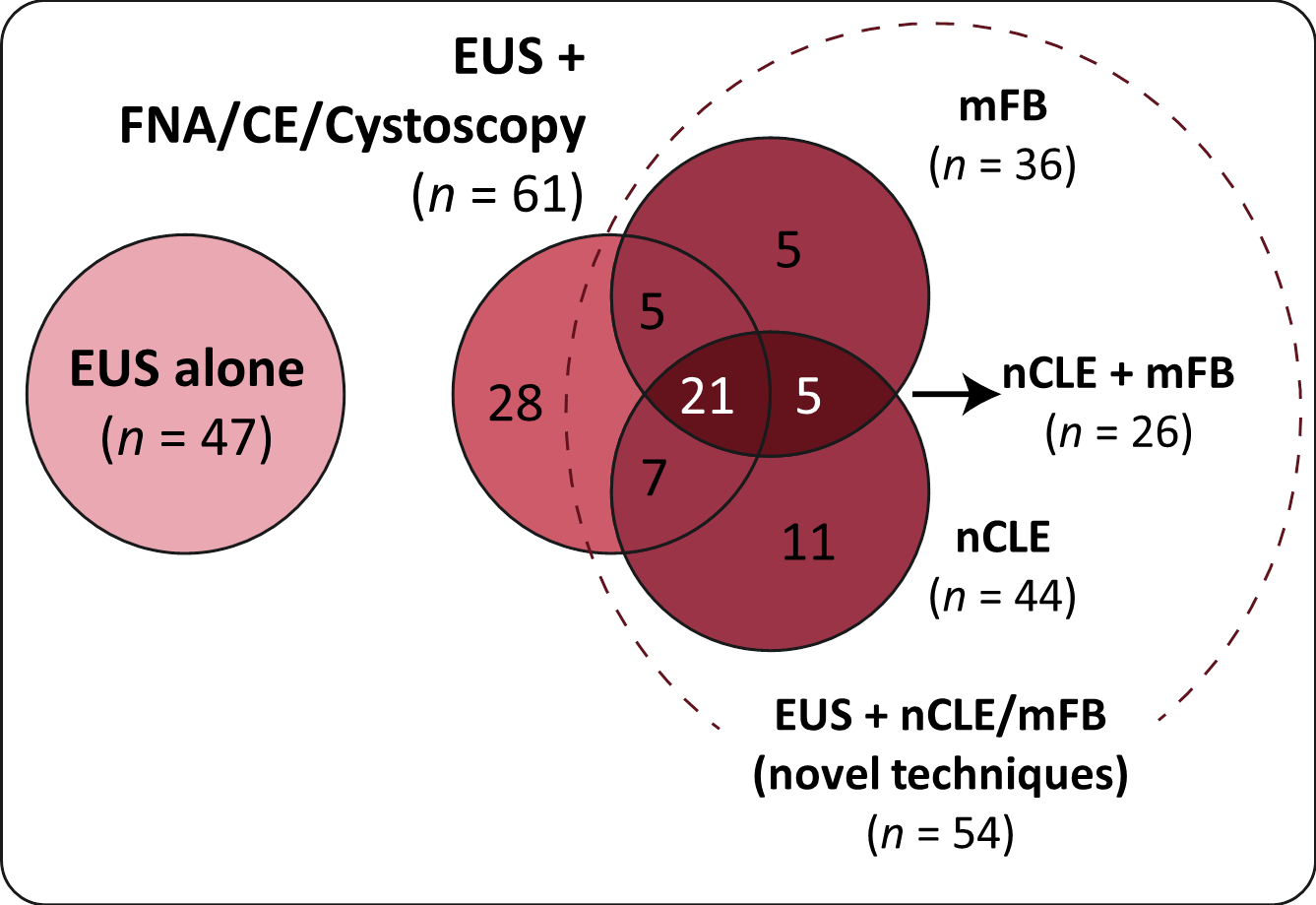
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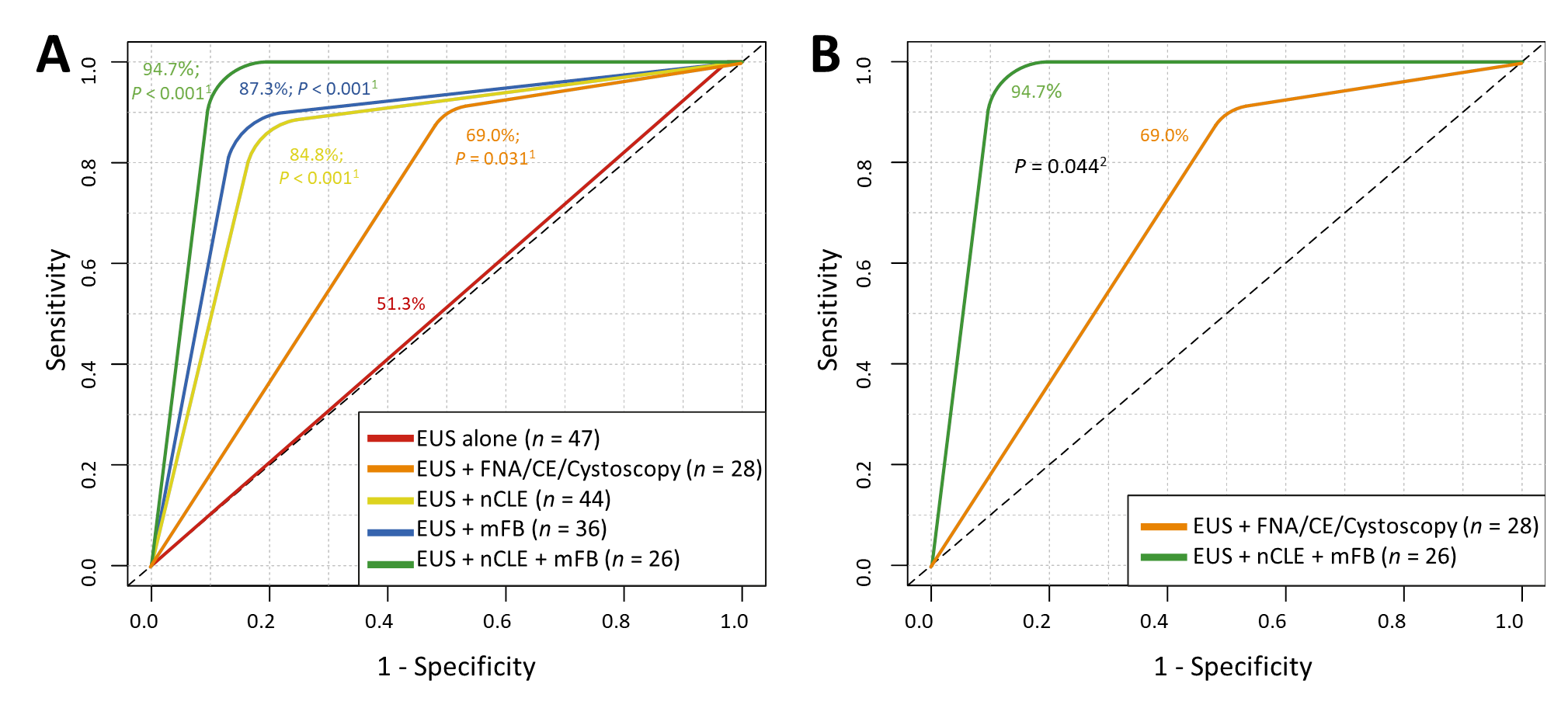
**Figure 1 Case No. 13: A 77 years old woman with a** **pancreatic cyst lesion corresponding to an intraductal papillary mucinous neoplasm.** The lesion exhibited malignancy criteria at endoscopic ultrasound (EUS) and related techniques. A: EUS identifying a 4 cm pancreatic cyst lesion with mural nodules (yellow arrow); B: Mural nodule with hyper-enhancing at EUS (green arrow) shown in contrast-enhanced EUS; C: EUS-guided cystoscopy using a digital probe showing vascularity (red arrow) of a pancreatic macrocystic lesion filled with clear fluid.



**Figure 2 Population study flowchart.** 1Numbers of techniques were not mutually exclusive. Endoscopic ultrasound could be combined with more than one other technique, as shown on the illustrated Venn diagram in Figure 3. EUS: Endoscopic ultrasound; EUS-FNA: Endoscopic ultrasound-guided fine needle aspiration; Cystoscopy: Fiberoptic probe cystoscopy; nCLE: Endoscopic ultrasound-guided needle-based confocal laser-endomicroscopy; mFB: Endoscopic ultrasound-guided through-the-needle direct intracystic micro forceps biopsy; CE-EUS: Contrast-enhanced endoscopic ultrasound; M: Malignancy.



**Figure 3 Venn diagram describing distribution of additional diagnostic techniques performed in the studied population.** EUS: Endoscopic ultrasound; EUS-FNA: Endoscopic ultrasound-guided fine needle aspiration; Cystoscopy: Fiberoptic probe cystoscopy; nCLE: Endoscopic ultrasound-guided needle-based confocal laser-endomicroscopy; mFB: Endoscopic ultrasound-guided through-the-needle direct intracystic micro forceps biopsy; CE-EUS: Contrast-enhanced endoscopic ultrasound.



**Figure 4** **Received operating characteristics describing overall diagnostic accuracy of endoscopic ultrasound alone and in addition with fine needle aspiration or contrast-enhanced endoscopic ultrasound,** **needle-based confocal laser-endomicroscopy and/or with** **direct intracystic micro forceps biopsy for detecting malignancy.** A: Comparison among endoscopic ultrasound (EUS) alone *vs* additional diagnostic techniques; B: Comparison among EUS alone *vs* EUS + EUS-guided needle-based confocal laser-endomicroscopy (nCLE) + EUS-guided through-the-needle direct intracystic micro forceps biopsy (mFB). 1DeLong’s test for two received operating characteristics (ROC) curves comparing EUS-alone area under the ROC curve (red line) with EUS + fine needle aspiration (FNA)/contrast-enhanced (CE) (orange line), EUS + nCLE (yellow line), EUS + mFB (blue line) and EUS + nCLE + mFB (green line). **2**DeLong’s test for two ROC curves comparing EUS + FNA/CE (orange line) with EUS + nCLE + mFB (green line). EUS: Endoscopic ultrasound; FNA: Fine needle aspiration; Cystoscopy: Fiberoptic probe cystoscopy; nCLE: Endoscopic ultrasound-guided needle-based confocal laser-endomicroscopy; mFB: Endoscopic ultrasound-guided through-the-needle direct intracystic micro forceps biopsy; CE: Contrast-enhanced.

**Table 1 Baseline characteristics and clinical 24-mo follow-up outcome of included patients**

|  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- |
|  | **Total (*n* = 129)** | **EUS alone (*n* = 47)** | **EUS + FNA/CE/ Cystoscopy (*n* = 28)** | **EUS + mFB/nCLE (novel techniques) (*n* = 54)** | ***P* value** |
| Age (yr), median (range) | 69 (26-97) | 71 (29-97) | 78 (49-92) | 59 (27-97) | < 0.001a |
| Sex (female), *n* (%) | 90 (69.8) | 33 (70.2) | 19 (67.0) | 38 (70.4) | 0.9694b |
| Pancreatic cyst location, *n* (%) |  |  |  |  | 0.6258b |
| Uncinate process | 3 (2.3) |  |  | 3 (5.6) |  |
| Head | 46 (35.7) | 17 (36.2) | 9 (32.1) | 20 (37.0) |  |
| Neck | 13 (10.1) | 3 (6.4) | 4 (14.3) | 6 (11.1) |  |
| Body | 36 (27.9) | 14 (29.8) | 8 (28.6) | 14 (25.9) |  |
| Tail | 31 (24.0) | 13 (27.7) | 7 (25.20) | 11 (20.4) |  |
| Cyst size (mm), *n* (%) |  |  |  |  |  |
| < 10 mm | 33 (25.6) | 29 (61.7) | 1 (3.6) | 3 (5.6) | < 0.001b |
| 10-30 mm | 61 (47.3) | 16 (34.0) | 19 (67.9) | 26 (48.1) |
| > 30 mm | 35 (27.1) | 2 (4.3) | 8 (28.6) | 25 (46.3) |
| Additional endoscopic procedure used for diagnosis1, *n* (%) | | | | | - |
| EUS-FNA | 21 (16.3) |  | 17 (60.7) | 4 (7.4) |  |
| CE-EUS | 20 (15.5) |  | 11 (39.3) | 9 (16.7) |  |
| Cystoscopy | 27 (20.9) |  | 1 (3.6) | 26 (48.1) |  |
| mFB | 36 (27.9) |  |  | 36 (66.7) |  |
| nCLE | 44 (34.1) |  |  | 44 (81.5) |  |
| Pancreatic cyst diagnosis, *n* (%) | | | | | < 0.001b |
| Malignant2 | 81 (62.8) | 46 (97.9) | 19 (67.9) | 16 (29.6) |  |
| Mucinous cystadenocarcinoma | 6 (4.7) | 1 (2.1) | 4 (14.3) | 1 (1.9) |  |
| Mucinous cystadenoma | 4 (3.1) |  | 1 (3.6) | 3 (5.6) |  |
| Intraductal papillary mucinous neoplasm | 70 (54.3) | 45 (95.7) | 14 (50.0) | 11 (20.4) |  |
| Neuroendocrine | 1 (0.8) |  |  | 1 (1.9) |  |
| Non-malignant2 | 48 (37.2) | 1 (2.1) | 9 (32.1) | 38 (70.4) |  |
| Serous cystadenoma | 46 (35.7) | 1 (2.1) | 9 (32.1) | 36 (66.7) |  |
| Pseudocysts | 2 (1.6) |  |  | 2 (3.7) |  |
| 24-mo follow-up, *n* (%) | | | | | 0.0351b |
| Malignant | 28 (21.7) | 7 (14.9) | 11 (39.3) | 10 (18.5) |  |
| Non-malignant | 101 (78.3) | 40 (85.1) | 17 (60.7) | 44 (81.5) |  |
| Positive observed agreement between EUS-guided biopsy *vs* 24-mo follow-up for malignancy detection, *n* (%) | 70 (54.3) | 8 (17.0) | 18 (64.3) | 44 (81.5) | < 0.001b |

aKruskal-Wallis rank sum test.

bPearson's Chi-squared test.

1Additional endoscopic procedures are not mutually exclusive.

2Cases with histopathological confirmation met the Fukuoka criteria.

EUS: Endoscopic ultrasound; EUS-FNA: Endoscopic ultrasound-guided fine needle aspiration; Cystoscopy: Fiberoptic probe cystoscopy; nCLE: Endoscopic ultrasound-guided needle-based confocal laser-endomicroscopy; mFB: Endoscopic ultrasound-guided through-the-needle direct intracystic micro forceps biopsy; CE-EUS: Contrast-enhanced endoscopic ultrasound.

**Table 2 Association between different additional performed techniques *vs* a positive observed agreement for malignancy diagnosis among endoscopic ultrasound and endoscopic ultrasound-related techniques *vs* 24-mo follow-up [OR (95%CI; *P* value)]**

|  |  |  |
| --- | --- | --- |
|  | **Univariate analysis**1 | **Multivariate analysis**1 |
| EUS alone (*n* = 47) | 0.066 (0.025-0.157; < 0.001) |  |
| EUS-FNA (*n* = 21) | 2.409 (0.905-7.182; 0.091) |  |
| CE-EUS (*n* = 20) | 1.694 (0.642-4.811; 0.298) |  |
| Cystoscopy (*n* = 27) | 4.950 (1.862-15.695; 0.003) | 0.622 (0.125-2.813; 0.541) |
| mFB (*n* = 36) | 6.625 (2.667-19.024; < 0.001) | 3.425 (1.104-11.682; 0.038) |
| nCLE (*n* = 44) | 10.489 (4.242-30.125; < 0.001) | 8.441 (2.698-33.081; < 0.001) |

1Positive observed agreement: In 70/129 (54.3%) there was a positive agreement between endoscopic ultrasound *vs* 24-mo follow-up for a malignant and non-malignant diagnosis.

EUS: Endoscopic ultrasound; EUS-FNA: Endoscopic ultrasound-guided fine needle aspiration; Cystoscopy: Fiberoptic probe cystoscopy; nCLE: Endoscopic ultrasound-guided needle-based confocal laser-endomicroscopy; mFB: Endoscopic ultrasound-guided through-the-needle direct intracystic micro forceps biopsy; CE-EUS: Contrast-enhanced endoscopic ultrasound.

**Table 3 Overall diagnostic accuracy for determining malignancy [% (95%CI)]**

|  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- |
|  | **EUS alone (*n* = 47)** | **EUS + FNA/CE/ Cystoscopy (*n* = 28)** | **EUS + mFB (*n* = 36)** | **EUS + nCLE (*n* = 44)** | **EUS + nCLE + mFB (*n* = 26)** |
| Sensitivity | 7/7; 100.0% (59.3-100.0) | 10/11; 90.9% (58.7-99.8) | 7/8; 87.5% (47.3-99.7) | 8/9; 88.8%; (51.8-99.7) | 7/7; 100.0% (59.0-100.0) |
| Specificity | 1/40; 2.5% (0.1-13.2) | 8/17; 47.1% (22.9-72.3) | 23/28; 82.1% (63.1-93.9) | 30/35; 85.7% (69.7-95.2) | 17/19; 89.4% (66.9-98.7) |
| PPV | 7/46; 15.2% (6.3-28.9) | 10/19; 52.6% (28.9-75.6) | 7/12; 58.3% (27.7-84.8) | 8/13; 61.5% (31.6-86.1) | 7/9; 77.8% (40.0-97.1) |
| NPV | 1/1; 100.0% (2.5-100.0) | 8/9; 88.9% (51.8-99.7) | 23/24; 95.8% (78.9-99.8) | 30/31; 97% (83-100) | 17/17; 100.0% (80.5-100.0) |
| PLR | 1.03 (0.98-1.08) | 1.72 (1.06-2.79) | 4.90 (2.12-11.31) | 6.22 (2.68-14.47) | 9.50 (2.56-35.24) |
| NLR | n/a | 0.19 (0.03-1.34) | 0.15 (0.02-0.96) | 0.13 (0.02-0.83) | n/a |
| Observed agreement | 8/47 (17%); *P =* 0.672a | 18/28 (64.3%); *P =* 0.049a | 30/36 (83.3%); *P <* 0.001a | 38/44 (86.4%); *P <* 0.001a | 24/26 (92.3%); *P <* 0.001a |
| MCC | + 0.06 | + 0.40 | + 0.61 | + 0.66 | + 0.83 |
| AU-ROC | 51.3%; *P =* 0.359b | 69.0%; *P =* 0.02b | 84.8%; *P <* 0.001b | 87.3%; *P <* 0.001b | 94.7%; *P <* 0.001b |

aFisher's exact test for count data.

bMann–Whitney U test.

EUS: Endoscopic ultrasound; nCLE: Confocal laser endomicroscopy; mFB: Endoscopic ultrasound-guided through-the-needle direct intracystic micro forceps biopsy; PPV: Positive predictive value; NPV: Negative predictive value; PLR: Positive likelihood ratio; NLR: Negative likelihood ratio; MCC: Matthews correlation coefficient; AU-ROC: Area under the receiver operating characteristics curve; n/a: Not available.