Biliary strictures, inducing cholestasis and jaundice, result from an obstructive process that can emerge intraluminally from the biliary epithelium, either a cholangitis or a primary cholangiocarcinoma, or extraluminally from compression by a pancreatic malignancy, a metastatic mass or a regional inflammatory process such as pancreatitis. The list of diseases, benign and malignant, which can produce a biliary stenosis is pretty long, although 80 % are malignant, with pancreatic carcinoma the first in line [1, 2]. However, the diagnosis is often easy when a mass is clearly defined and amenable to endoscopic ultrasound-guided fine-needle aspiration (EUS-FNA), which has nearly 90 % sensitivity [3]. The diagnosis is also straightforward when orthotopic liver transplantation is complicated by a short stricture centered on the bilo-biliary anastomosis, or after a difficult cholecystectomy with bile duct injury and protracted biliary leak. However, when the mass is missing and the clinical history is poor or unclear, the stricture can remain indeterminate for a while, which places physicians in the uncomfortable position of operating for a benign disease with potentially undue morbidity and mortality or leaving an undiagnosed malignancy unchecked. Diagnosis in these cases relies mostly on endoscopic retrograde cholangiopancreatography (ERCP)-based sampling techniques, essentially brush cytology and/or fluoroscopy-guided forceps biopsies, which have a sensitivity of less than 50 % [2, 4].

In this issue of Endoscopy International Open, Moura et al. have tried to improve our understanding of the respective inputs of both approaches by comparing the diagnostic performances of ERCP-guided sampling and EUS-FNA in a group of 50 patients selected for a suspected biliopancreatic malignancy revealed by a biliary stricture. The patient selection process, although claiming only indeterminate strictures were included, understates that most patients had a tissular mass obstructing the bile duct which was amenable to EUS-FNA, after which one can predict that most strictures will not remain indeterminate. However, the strength of the study was to have each patient be his or her own control by performing both EUS-FNA and ERCP sampling during the same sedation. Sampling methods were state-of-the-art, including both suction and the slow-pull capillary method for FNA, and a combination of brushing and biopsies during ERCP with cell-block technique to optimize cytological diagnosis. The authors based the study power on a reasonable hypothesis of a 75 % vs 49 % difference of sensitivity in favor of EUS-FNA. Surprisingly, after surgical resection or a follow-up of at least 6 months, only 2 out of 50 patients (4 %) were found to have benign disease, meaning that patients were highly selected before enrollment with a very strong suspicion of malignancy. Sensitivities for ERCP-guided techniques were, as expected, much lower than that for EUS-FNA, with 61 % for the combined brushing and fluoroscopy-guided biopsies, and short of 40 % for brushings alone, as against nearly 94 % for EUS, whereas the combination of ERCP and EUS slightly increased overall sensitivity to close to 98 %. However, as a consequence of the super selection of patients, negative predictive values were very low, below 10 % for ERCP, at only 40 % for EUS and no more than 67 % for combined EUS and ERCP, a figure closer to what is expected for EUS alone in an unselected patient population with a solid pancreatic mass [3]. The proximal vs distal location of the stricture did not appear to significantly affect the diagnostic performance of either technique, although EUS did slightly better in distal rather than proximal strictures and ERCP did the opposite. A more important finding was that the superiority of EUS was overwhelming in bigger lesions, here considered as larger than 1.5 cm in diameter, whereas both methods were rated equally in smaller lesions, at 50 % sensitivity, although the figure was probably based on a small number
Finally, the superiority of EUS was once again clearly demonstrated in extraductal lesions (ie pancreatic carcinomas) but not so much in intraductal ones (ie primary biliary carcinomas) with an accuracy of 82.4% for EUS vs. 70.6% for ERCP.

The weaknesses of ERCP-guided sampling techniques are well-known: whereas extraductal masses, generally pancreatic or lymphatic, cannot be adequately sampled from a neighboring structure unless they invade its lumen, the superficial scratching of intraductal strictures is also frequently inconclusive because those lesions, whether inflammatory or neoplastic, generally develop a thick and dense fibrous microenvironment within which the cells of interest can be scarce and/or deeply seated [5]. Even fluoroscopy-guided biopsies, although reaping deeper material, are often unsuccessful because targeting is inaccurate and the tangential incidence of the forceps jaws makes biopsies too superficial. On the contrary, EUS-FNA or endoscopic ultrasound-guided fine-needle biopsy (EUS-FNB) is excellent at collecting rich cytology and even microbiopsies, especially with recent refinements in needle designs and sampling techniques [6, 7], but that comes with a caveat: EUS-FNA/FNB requires a target volume, approximately over 500 mm$^3$ or 10 mm in diameter, to be efficient. When the target is too discrete, very thin and purely infiltrative, EUS results rapidly decrease and lead to stricture indetermination. In such cases, ERCP brushings and biopsies remain the first-line diagnostic option, but still leave about 50% of cases indeterminate. This is the place for more advanced techniques, such as ERCP-guided retrograde cholangiography with endoscopically, not fluoroscopically, guided biopsies, which can solve a significant part of the problem of indeterminate biliary strictures [8]. The diagnostic performance of cholangiography in such cases has been estimated at about 70% in recent meta-analyses with single-operator devices [9, 10], but the impact on patient management and outcomes remains to be assessed. The residual 30% of indeterminate biliary strictures will have to be dealt with using either innovative sampling methods or more advanced, molecular-based analyses to overcome current cytological limitations [11, 12].

Competing interests

None

References