Introduction

Identifying the cause of acute pancreatitis represents a diagnostic challenge in a significant number of patients. Diagnostic etiology is undefined in 10% of patients with a first episode of acute pancreatitis and in 30% with recurrent acute pancreatitis. The initial evaluation should include history of infectious disease, trauma, or recent abdominal surgery; alcohol and drug intake; calcium analysis, lipid profile, ANA and IgG4. Imaging typically includes transcorporeal abdominal ultrasound and possibly cross-sectional imaging. If the cause is not defined following these exams, patients can be considered to have idiopathic acute pancreatitis (IAP) [1–3].
Identifying the cause of pancreatitis in patients with IAP is important, because up to 70% may suffer from recurrent episodes [4], and increased mortality has been described in these individuals [5]. Endoscopic retrograde cholangiopancreatography (ERCP) has been considered as part of the diagnostic pathway, reporting a diagnostic yield reaching up to 80%, but with 10% to 15% incidence of adverse events (AEs) [6,7] including post-ERCP pancreatitis. Another technique is collection of bile for microscopic bile examination (MBE) with a polarized filter. This has a false-negative rate of 29% to 50%, is time consuming, and cannot be completed in up to 20% of patients [8,9].

Given the disadvantages of ERCP and MBE, two techniques have been demonstrated to have a high diagnostic value in IAP: endoscopic ultrasonography (EUS) and magnetic resonance cholangiopancreatography (MRCP). EUS provides an etiologic diagnostic yield of 60% to 80% [3,10–14]. The yield of MRCP ranges from 32% to 90% and improves when associated with secretin stimulation (S-MRCP) [15,16].

The diagnostic yield of both examinations in patients with IAP has been compared in three studies with variable results that have not demonstrated the superiority of either modality [17–19]. In two studies, EUS was compared with MRCP with no secretin stimulation [18,19]. In the third study, ERCP was included in the comparison performed in patients with recurrent IAP [17].

Considering this background, we proposed a unique study comparing the diagnostic yield between EUS or with S-MRCP in patients with IAP.

Patients and methods
We designed a prospective, single-blind, comparative study, offering enrollment to consecutive patients admitted with IAP to a single tertiary University hospital between January 2009 and December 2013; patients were followed for a minimum of 6 months.

Acute pancreatitis was defined as the occurrence of abdominal pain associated with amylase and lipase levels three times and twice over normal value, respectively. We considered IAP when there was no recent history of infection, trauma or surgery; no history of alcohol consumption >60 g of ethanol per day in men and >40 g for women, or drugs implicated in pancreatitis; IgG4 and ANA were normal; triglycerides <500 mg/dL, calcium <10 mg/dL; and two abdominal ultrasound or an ultrasound and computed tomography scan not showing cause of acute pancreatitis.

Patients who met criteria were diagnosed with IAP and were invited to participate in the study. The inclusion criteria in this study were age >18 years, diagnosis of IAP during the period of inclusion, willingness to undergo EUS and S-MRCP, and willingness to be included in the study after the informed consent process.

Patients with Billroth II or total gastrectomy were excluded because the study with EUS could not be completed.

The included patients underwent EUS and S-MRCP no sooner than 4 weeks after hospital discharge for IAP. Examinations were performed by two echoendoscopists and two radiologists, each of whom had a minimum of 3 years of experience.

After each EUS and S-MRCP was performed, a report was delivered to the control researcher. The control researcher was responsible for recording the patients included, scheduling examinations, and recording EUS and S-MRCP results. After both reports were received, the researcher recorded them in the electronic medical record, making them available to the physicians involved in clinical care. The echoendoscopists and radiologists performing the examinations were thus blinded to the results of the other examination.

Data recorded for analysis included information on demographics, prior cholecystectomy, prior episodes of pancreatitis, EUS and S-MRCP findings, and any treatments performed. During the minimum 6-month follow-up period, clinical outcomes and results of new examinations performed were included in defining the final diagnosis.

EUS exams were performed with a radial echoendoscope (EG3670URK, Pentax, Hamburg, Germany). Cholelithiasis and choledocholithiasis were defined by an arch-shaped image with posterior acoustic shadowing in gallbladder or bile duct. Microlithiasis was defined by the occurrence of fleeting, unshadowed, echogenic, point-shaped particles in the gallbladder. Given the clinical implication being identical, the presence of microlithiasis or gallstones were defined as cholelithiasis. We considered a minimum of five diagnostic criteria defined by Sahai [20] to establish a diagnosis of chronic pancreatitis.

S-MRCP was performed using a 1.5 Tesla magnet (Magnetom Avanto, Siemens, Erlangen, Germany) with the patient in a fasting state. The following parameters were employed: TR110, TE4,72, G6 mm for axial T1; TR1000, TE85, G6 mm for axial HASTE and TR4500, TE754, G30 mm with apnea for a heavily T2-weighted cholangiographic sequence with without secretin. Secretin 0.1 U/kg was injected as a bolus, with images obtained every minute for 10 minutes after injection. Five minutes before the S-MRCP, 400 cc of negative oral contrast was administered. Pancreatic ductal and parenchymal morphology, gallbladder or ductal stones, reduced T1 signal intensity, pancreatic secretion, and ductal morphology were evaluated before and after secretin injection as well as response of the sphincter of Oddi. Chronic pancreatitis was diagnosed based on ductal findings according to the Cambridge classification [21].

For the diagnosis of cholelithiasis, we considered the EUS result as definitive. During the follow-up period, some patients underwent cholecystectomy and ERCP. The final diagnosis of cholelithiasis was further refined by these interventions. Likewise, a final diagnosis of chronic pancreatitis may have included data derived from subsequent ERCP and cross-sectional imaging. A final diagnosis of pancreas divisum included subsequent ERCP in some patients. Some patients with pancreatic cystic lesions underwent subsequent resection to refine the final diagnosis. To establish a final diagnosis, we considered all the study results, examinations, and interventions in the follow-up period. Therapeutic decisions were made based on clinical indications by a multidisciplinary team including clinicians, endoscopists, radiologists, and surgeons.
Statistical analysis was performed with SPSS 20. Quantitative descriptive data are provided as mean ± standard deviation and range. The degree of accordance between EUS and S-MRCP was evaluated by using the kappa statistic. Sensitivity, specificity, and positive and negative predictive values were estimated to study diagnostic test validity. In addition, to compare the number of successes between EUS and S-MRCP, success rates were estimated with confidence intervals at 95 %, and the McNemar test for paired samples was used. P<0.05 was considered statistically significant.

The study was approved by our hospital scientific research ethics committee.

Results

All 40 patients admitted with IAP during the study period were invited to enroll; four refused EUS and two had a pacemaker, which precluded S-MRCP. Demographic, clinical, and radiological data from the 34 patients finally enrolled are shown in Table 1.

The mean time between performance of EUS and S-MRCP was 13.82±13 days (range: 2–49). Etiologic findings of EUS and S-MRCP compared with the final diagnosis are shown in Table 2. In four patients, a possible second cause was found on EUS that was not considered responsible for IAP. Patients were followed up for a mean of 31.62±19.20 months (range: 6–60) after EUS and S-MRCP.

Thirteen of 15 patients with cholelithiasis on EUS underwent cholecystectomy. Gallbladder histological examination showed chronic cholecystitis in 12 patients, associated with cholesterol polyps in three and gallbladder adenomyosis in one. The other patient had macroscopically visible stones smaller than 3 mm in the gallbladder. During follow-up, one of these patients suffered recurrent pancreatitis. Another patient developed abdominal pain 2 years after IAP and 1 year later was diagnosed with unresectable pancreatic cancer. One patient who was asymptomatic after cholecystectomy suffered sudden death after 1 year of follow-up. Three patients refused cholecystectomy and remained asymptomatic.

The initial diagnosis of chronic pancreatitis made in EUS was confirmed during follow-up in four of five patients by means of subsequent EUS and functional testing. One patient suffered recurrent episodes of pancreatitis and the other three remained asymptomatic. In the fifth patient, chronic pancreatitis

### Table 1 Demographic, clinical and radiological data of the 34 patients before inclusion in the study.

| Male/female | 17/17 |
| Age | 63.53 ± 12.63 (range: 21–78 years) |
| Abdominal ultrasound | 2.21 ± 1.51 (range: 1–7) |
| Abdominal CT | 1.18±1.31 (range: 0–5) |
| Previous pancreatitis | 1.52 ± 1.06 (range: 1–5) |
| Previous cholecystectomy | 7 patients |
| CT, computed tomography |

### Table 2 Main etiologic findings with EUS and S-MRCP compared with final diagnoses after follow-up (definitive diagnosis).

| Final Diagnosis | Total |
|-----------------|-------|
|                 | Normal | Cholelithiasis | CBD Stones | Pancreas Divisum | IPMT | CP |
| **EUS**         |        |                |           |                 |      |    |
| Normal          | 4      |                | 1         | 1                | 1    | 6  |
| Cholelithiasis  | 15     |                | 2         | 2                | 3    | 15 |
| CBD stones      | 2      |                | 3         | 3                | 3    | 2  |
| Pancreas divisum| 3      |                | 3         | 3                | 3    | 3  |
| IPMT            |        |                | 3         | 3                | 3    | 3  |
| CP              | 1      |                | 1         | 1                | 1    | 4  |
| **S-MRCP**      |        |                |           |                 |      |    |
| Normal          | 4      | 16             | 1         | 0                | 2    | 24 |
| Cholelithiasis  | 0      |                | 1         | 1                | 1    |    |
| CBD stones      | 1      |                | 4         | 4                | 4    | 4  |
| Pancreas divisum| 4      |                | 3         | 3                | 3    | 3  |
| IPMT            |        |                | 3         | 3                | 3    | 3  |
| CP              | 1      |                | 1         | 1                | 1    | 2  |
| **Total**       | 4      | 16             | 2         | 4                | 4    | 4  |

EUS, endoscopic ultrasound; S-MRCP, secretin-enhanced magnetic resonance choangioipancreatography; CBD, common bile duct; IPMT, intraductal papillary mucinous tumor; CP, cholangiopancreatography.
was not confirmed in a subsequent EUS where biliary microthiasis was observed. This patient was considered a false-positive in EUS for chronic pancreatitis and a false-negative for cholelithiasis in the initial EUS.

The two patients with choledocholithiasis underwent ERCP, which confirmed the diagnosis. Both remained asymptomatic thereafter. In addition, three of four patients with pancreas divisum underwent ERCP with minor papilla sphincterotomy and stent placement. One subsequently suffered another episode of pancreatitis. The fourth patient refused ERCP and remained asymptomatic.

One patient with intraductal papillary mucinous tumors (IPMT) showed signs of malignancy on EUS with fine-needle aspiration, which were not identified on S-MRCP. Malignancy was confirmed histologically after surgical resection and the patient remained asymptomatic thereafter. Side-branch IPMT was also confirmed histologically in two more patients after surgical resection. One patient suffered recurrence of pancreatitis. The fourth patient did not undergo surgery because of high surgical risk and the cyst became malignant and the patient died 3 years after diagnosis. The size of these IPMTs ranged between 5 mm and 40 mm and were located in the uncinate process (1), pancreatic head (2) and pancreatic body (1).

In summary, the final diagnoses in this series were cholelithiasis in 16 patients, choledocholithiasis in two, pancreas divisum in four, chronic pancreatitis in four and side-branch IPMT in four patients (▶Table2). In the remaining four patients, the cause of pancreatitis was not identified with EUS or S-MRCP or during follow-up. The diagnostic, therapeutic and follow-up data from the patients included are described in ▶Fig.1.

According to these final results, a correct etiologic diagnosis was made in 27 patients (79.4 %) with EUS and in 10 (29.4 %) with S-MRCP. Nineteen of 24 patients with normal findings on S-MRCP were diagnosed with EUS. Two of six patients with normal EUS were diagnosed with S-MRCP.

According to the kappa correlation value, a slight agreement between EUS and S-MRCP was found (kappa = 0.236, SE of kappa = 0.092, 95 % confidence interval: 0.055–0.416). The diagnostic value of EUS was statistically higher than S-MRCP: 79.4 % (CI 95 %: 65 %–94 %) vs 29.4 % (CI 95 %: 13 %–46 %) (P = 0.001).

Figure 1

Diagnostic, therapeutic, and follow-up etiological data from patients with IAP included in the study.
The sensitivity, specificity, and positive p and negative predictive values of EUS and S-MRCP for the etiologic diagnosis of IAP were 90%, 80%, 96%, 57%, 33%, 100%, 100%, and 16%, respectively. In patients with gallbladder (n = 27), the difference in the diagnostic value between EUS and S-MRCP was increased: 81% (CI 95%: 66–97%) of patients were diagnosed with EUS compared to 19% (CI 95%: 3–34%) with S-MRCP (P = 0.0007). In cholecystectomized patients (n = 7), the diagnostic difference disappeared and both examinations were diagnostic in five patients (P = 1).

Discussion

In this study, the value of EUS was significantly higher than that of S-MRCP in the etiologic diagnosis of patients with IAP. The diagnostic values of EUS and MRCP in patients with IAP have been compared prospectively in three studies [17–19]. The first study compared the diagnostic yield of S-MRCP, secretin-enhanced EUS and ERCP in 44 patients with recurrent IAP [17]. The diagnostic values were 65.9%, 79.5%, and 62.8% respectively (P > 0.05).

The second study compared the diagnostic value of EUS and MRCP in 49 patients with IAP [18]. The diagnostic yield of EUS was statistically significantly higher (51% vs 20%; P = 0.001), especially in patients with gallbladder (60% vs 11%; P = 0.008) irrespective of the number of pancreatitis bouts.

The third study described the diagnostic yield of both examinations within a comprehensive diagnostic algorithm for acute pancreatitis [19]. In this study, 128 patients were enrolled and an etiologic diagnosis was found on initial examination in 83 (65%). Thirty-eight of the remaining 45 patients underwent EUS and MRCP, showing a higher diagnostic yield with EUS (39.5% vs 21%) without statistically significant differences (P > 0.09).

There are important methodological differences in our study compared with the previous three studies regarding population included, design, examinations performed, and timing of the examinations. The sample of patients included in our study is similar those of Repiso et al [18] and Thevenot et al [19]. However, Mariani et al [17] included a highly selected sample of patients with recurrent IAP in normal bile duct. These patients represent < 32% of the sample of the other studies.

Moreover, although these studies are prospective and blinded, none explain how this condition was achieved. In our study a control researcher kept the result of each examination unavailable until both were performed. Besides, we wanted to guarantee that the study design did not harm the patient ethically by delaying the diagnostic or therapeutic process.

There are also differences in the examinations performed. Mariani et al [17] added stimulation with secretin to EUS, a technique not widely used in clinical practice, limiting the applicability of the results [18]. Repiso et al [18] and Thevenot et al [19] performed MRCP without secretin stimulation, unlike our study.

With regard to the timing of examinations, Thevenot et al [19] performed most EUS and MRCP on the same day. Repiso et al [18] performed the EUS first, with a mean time difference of 24 ± 15 days with regard to the MRCP, and Mariani et al [17] did not report this information. This time difference may be important because in 40% of patients, biliary sludge disappears completely, a further 40% show a cyclical pattern of occurrence, and in 20%, there is progression to cholelithiasis [2, 22, 23]. Therefore, the diagnosis of biliary microlithiasis may vary depending on the timing with regard to the IAP bout. Although in our study was not possible to perform both examinations on the same day for logistical reasons, we tried to reduce this variability as much as possible. The mean time difference between both examinations was 13.82 ± 13 days (range: 2–49), lower than with Repiso et al [18].

The most frequent finding in our study, as in that of Repiso et al [18] and Thevenot et al [19], was microlithiasis or biliary cholelithiasis. This finding makes the difference between EUS and S-MRCP, because no cases of biliary microlithiasis were diagnosed with it, as in prior studies [17–19]. Furthermore, it is the most common cause of IAP in most studies, with a incidence as high as 80% [2, 24]. Although MBE has been considered the gold standard for diagnosing biliary microlithiasis, more recent studies have shown better results with EUS [25, 26] and justify the use, along with other authors [17], of this result as a baseline in our study. In addition, the finding of chronic cholecystitis has been associated with the presence of gallstones [27, 28]. In our study, 12 of 13 patients undergoing cholecystectomy had chronic cholecystitis, which supports the diagnosis obtained by EUS.

However, in the study by Mariani et al [17], the most frequent finding was chronic pancreatitis. This is related to the high sample selection, but also to the diagnostic criteria considered to diagnose IAP and chronic pancreatitis. In our study, alcohol consumption > 60 g in men and > 40 g in women was considered to be a possible cause of pancreatitis and we diagnosed chronic pancreatitis with five EUS criteria. This cutoff point provides sensitivity and specificity of 60% and 83%, respectively, a high positive predictive value, an excellent correlation with the ERCP for moderate and severe involvement (κ = 0.82), and a good interobserver correlation (κ = 0.45) [20, 29]. Repiso et al [18] placed the limit at 80 g of ethanol and three diagnostic criteria. Thevenot et al [19] limited consumption to 40 g and did not specify the number of EUS diagnostic criteria for diagnosis of chronic pancreatitis. Mariani et al [17] stated that patients who did not abuse alcohol were enrolled and the diagnostic cutoff was four criteria. The percentage of patients diagnosed of chronic pancreatitis clearly varies depending on these parameters. In our study and that of Thevenot et al [19], chronic pancreatitis was diagnosed in 11.8% and 10.5% of patients while Mariani et al [17] and Repiso et al [18], with less restrictive criteria, report a higher percentage of 25% and 18%, respectively.

The diagnostic yield of MRCP in our study was 29.4%, higher than that of Repiso et al (20%) [18] and Thevenot et al (21%) [19]. This difference is explained by the use of S-MRCP in our study, which improves results of MRCP [16]. The diagnostic yield of S-MRCP in the study by Mariani et al [17] was significantly higher (65.9%), which again can be explained by the
highly selected sample of the patients enrolled and a lower restriction on the alcohol intake limit.

Our study has various limitations. First, the study sample was heterogeneous because patients with and without gallbladder and with a single incident of and recurrent IAP were enrolled. Because of the small sample size, no subpopulation analysis could be performed, which may be regarded as a second limitation. Third, although we tried to minimize the time gap between EUS and S-MRCP, medical reasons or intermittent processes delayed the second examination in some cases, increasing the difference. Finally, taking the result of the baseline EUS and the S-MRCP into consideration may give rise to diagnostic bias. However, EUS is currently considered the gold standard for diagnosis of chronic pancreatitis and MRCP the gold standard for pancreas divisum, and no examination offers better results for diagnosing pancreatic cystic lesions than these two methods. To reduce this diagnostic bias, the histological results and the follow-up data were considered.

On the other hand, the prospective, single-blinded design and use of a control researcher, thereby minimizing biases, are remarkable strengths of the study. Second, comparison of EUS with S-MRCP, which is the best version of MRCP, was made. Finally, the etiological data from long-term follow-up were considered definitive. As a result, the initial diagnostic yield of EUS declined as a patient’s etiological diagnosis changed. This reduction in the initial diagnostic yield during follow-up has been described previously [12].

Conclusion
In conclusion, in our study, the diagnostic value of EUS was significantly higher than that of S-MRCP in patients with IAP. In our opinion and according to our results, a patient with IAP and a gallbladder should undergo EUS as the first examination to search for etiologic diagnosis. If the cause remains unidentified after EUS, S-MRCP should be performed. Further studies with adequate samples should clarify the role of both examinations in cholecystectomized patients with IAP.

Competing interests
Dr. Vila is a consultant for Boston Scientific

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