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**Outcome of endotherapy for pancreas divisum in patients with acute recurrent pancreatitis**

Mariani A *et al*.Outcome of patients with pancreas divisum

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**Abstract**

**AIM:** to assess the rate of relapses of acute pancreatitis (AP) recurrent AP (RAP) and the evolution of endosonographic signs of chronic pancreatitis (CP) in patients with pancreas divisum (PDiv) and RAP.

**Methods:** Over a five-year period, Patients with PDiv and RAP prospectively enrolled were divided into two groups: (1) those with relapses of AP in the year before enrollment were assigned to have endoscopic therapy (recent RAP group); and (2) those free of recurrences were conservatively managed, unless they relapsed during follow-up (previous RAP group). All patients in both groups entered a follow-up protocol that included clinical and biochemical evaluation, pancreatic endoscopic ultrasonography (EUS) every year and after every recurrence of AP, at the same time as endoscopic retrograde cholangiopancreatography (ERCP).

**Results:** Twenty-two were treated by ERCP and 14 were conservatively managed during a mean follow-up of 4.5 ± 1.2 years. In the recent RAP group in whom dorsal duct drainage was achieved, AP still recurred in 11 (57.9%) after the first ERCP, in 6 after the second ERCP (31.6%) and in 5 after the third ERCP (26.3%). Overall, endotherapy was successful 73.7%. There were no cases of recurrences in the previous RAP group. EUS signs of CP developed in 57.9% of treated and 64.3% of untreated patients. EUS signs of CP occurred in 42.8% of patients whose ERCPs were successful and in all those in whom it was unsuccessful (*P* = 0.04). There were no significant differences in the rate of AP recurrences after endotherapy and in the prevalence of EUS signs suggesting CP when comparing patients with dilated and non-dilated dorsal pancreatic ducts within each group.

**Conclusion:** Patients with PDiv and recent episodes of AP can benefit from endoscopic therapy. Effective endotherapy may reduce the risk of developing EUS signs of CP at a rate similar to that seen in patients of previous RAP group, managed conservatively. However, in a subset of patients, endotherapy, although successful, did not prevent the evolution of endosonographic signs of CP.

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**Key words:** Endoscopic retrograde cholangiopancreatography; Magnetic resonance cholangiopancreatography with secretin stimulation; Minor papilla endotherapy; Pancreatic stenting; Endoscopic ultrasonography

**Core tip:** In this paper we compared the outcome of patients with pancreas divisum (PDiv) and recent or previous recurrent acute pancreatitis (RAP) after minor papilla endotherapy or observation, respectively. We confirmed previous findings regarding the benefit of endotherapy in patients with PDiv and RAP. In addition, we showed that effective endotherapy in patients with recent bouts of pancreatitis reduced the risk of developing endoscopic ultrasonography signs of chronic pancreatitis (CP) at a rate similar to that seen in patients without recent episodes of acute pancreatitis who are managed conservatively. However, in a subset of patients, endotherapy, although successful, did not prevent the evolution of endosonographic signs of CP.

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**INTRODUCTION**

Pancreas divisum (PDiv) is the most common congenital variant of the pancreas, in which the ventral and dorsal ducts of the embryonic pancreas fail to fuse during organogenesis[1,2]. Less than 5% of people with PDiv have symptoms related to this altered anatomy[3,4]. However, the prevalence of PDiv amongst patients undergoing endoscopic retrograde cholangiopancreatography (ERCP) for unexplained acute pancreatitis (AP) is much higher, up to 25.6%[5,6]. Several studies have found an increased prevalence of PDiv in patients with recurrent acute pancreatitis (RAP)[5,7-9]. A recent study based on magnetic resonance cholangiopancreatography (MRCP) findings appeared to confirm that PDiv predisposes patients to RAP and chronic pancreatitis (CP)[4].

The mechanism responsible for the development of pain or AP in PDiv is unknown, but outflow obstruction of the pancreatic ductal system associated with stenosis or dysfunction of the minor papillary sphincter has been postulated[10-12]. Persisting outflow obstruction may lead to chronic damage to the gland. However, not all studies support the obstruction theory as the explanation for AP in PDiv[13] .

The goal of endoscopic therapy in PDiv patients is to open the minor sphincter to relieve presumed obstruction to pancreatic exocrine flow[13-15]. Although most published reports have been small case series, it has become clear that the RAP subset of PDiv patients is most likely to respond to endoscopic intervention, usually minor papillotomy[14-18]. However, it is still debated whether endoscopic minor sphincterotomy really improves the outcome and prevents progression to CP in these patients[19].

The aims of this prospective study were to assess the rate of relapses of CP and the evolution of morphological signs of CP in patients with PDiv and RAP.

**MATERIALS AND METHODS**

According to the protocol approved by our institutional review board, all patients with PDiv suffering from RAP referred to our institution were scheduled to undergo routine MRCP with secretin stimulation (ss-MRCP), endoscopic ultrasound (EUS), genetic screening [specifically, for cystic fibrosis transmembrane regulator (CFTR) gene mutation], serum IgG-subclass 4 (IgG-4), and fecal elastase assays. Patients also had a medical consultation if they suffered a recurrence of AP. RAP was defined as two or more episodes of abdominal pain associated with serum amylase and/or lipase levels at least double the upper normal limit (110 IU/L for amylase, 90 IU/L for lipase) are present. AP was classified as mild or severe according to the Atlanta criteria[20].

Over a five-year period (from January, 2005, to December, 2009), all consecutive patients with a history of RAP, without signs suggesting CP on EUS investigation, and PDiv documented by ss-MRCP, were entered into a prospective follow-up study. Patients reporting at least one episode of AP in the year before enrollment were assigned to minor papilla sphincterotomy (recent RAP group), while those with no recurrences in the same period were assigned to observation, unless they relapsed during follow-up (previous RAP group). Patients in both groups had either a dilated or a non-dilated dorsal pancreatic duct. Amongst patients assigned to endotherapy, only those in whom dorsal duct drainage was achieved, were included in the per protocol analysis.

The following data were collected for each patient: age, sex, time of the first attack of AP, number of AP recurrences before and after enrollment, duration of the disease, duration of follow-up and time to development of CP. Patients were excluded for any of the following reasons: pancreatitis associated with known alcohol abuse (≥ 60 g of pure alcohol per day), gallstones, trauma, drug abuse, elevated serum IgG-4, hypertriglyceridemia or hypercalcemia; CFTR gene mutations; decreased fecal elastase activity; CP, known or suspected intraductal papillary mucinous neoplasm or pancreatic cancer, family history of pancreatitis, pancreatic surgery and previous pancreatic sphincterotomy.

Informed consent for diagnostic procedures, blood sampling and data management for scientific purposes was routinely obtained from all patients.

The diagnosis of PDiv was established by MRCP when the dorsal pancreatic duct crossed the common bile duct to drain through the minor duodenal papilla, and was clearly separate from a smaller ventral duct[21]. The diameter of the main (dominant dorsal) pancreatic duct was considered dilated when its caliber was ≥ 3 mm. Ss-MRCP was performed in all patients to confirm the diagnosis of PDiv, to evaluate an abnormal pancreatic juice outflow through the minor papilla[22] and quantify pancreatic exocrine secretion.

***ERCP procedure***

ERCP procedures were done using a side-viewing duodenoscope (Pentax ED-3440T or ED3480TK or ED3680TK, Tokyo, Japan). The minor papilla was cannulated with a metal-tipped catheter (ERCP-1-Cremer, Cook Medical, Winston-Salem, NC, United States) or a pull-type sphincterotome (Mini-Tome MT-20M or Cannula-Tome II CT-20M, Cook Medical, Winston-Salem, NC, United States) which was used in all cases for sphincterotomy (minor papillotomy). Immediately after minor sphincterotomy, 5 French (Fr) gauge, single flanged, plastic pancreatic stents, 2-3 cm long, with a duodenal pigtail (SPSOF-5-3 Cook Medical, Winston-Salem, NC United States), or in some cases unflanged 5Fr stents (ZEPDS-5-2 Cook Medical, Winston-Salem, NC United States), were placed into the dorsal duct to prevent the development of post-ERCP pancreatitis. All patients received pharmacologic prophylaxis against post-ERCP pancreatitis using gabexate mesilate (Foy, Sanofi-Aventis, Milano, Italy)[23,24].

As with recurrences of AP, the Atlanta criteria were used to grade cases of post-ERCP pancreatitis[20]. All patients had plain abdominal x-rays approximately seven days after their pancreatic stent placement to verify its spontaneous migration into the duodenal lumen. Retained stents were removed using a duodenoscope at the earliest time available.

***EUS procedure***

EUS procedures were performed using a linear-scanning echoendoscope (EG-3830UT, EG-3630U, FG-36UX; Pentax, Hamburg, Germany) at 5-10 MHz. EUS procedures were carried out by two experienced endosonographers (MCP, PGA, each with more than 500 EUS procedures/year). The endosonographers were blinded to the clinical findings at enrollment and to the other EUS examinations during follow-up. All pancreatic examinations were reviewed carefully by both endosonographers using the standard nine Wiersema criteria for diagnosing CP[25]. In the event of equivocal EUS findings they reached a consensus agreement. When four or more Wiersema criteria were present, the EUS findings were considered suggestive of chronic pancreatitis[26,27].

***Follow-up protocol***

All patients in both groups entered a follow-up protocol that included: (1) clinical (abdominal pain) and biochemical evaluation including serum pancreatic enzyme levels and a surrogate marker for pancreatic exocrine function (fecal elastase: normal value > 200 µg/g; ScheBo Pancreatic Elastase 1 ELISA kit, ScheBo-Tech, Giessen, Germany); and (2) pancreatic EUS every year and after every recurrence of AP, at the same time as ERCP (*i.e.* under the same sedation).

Patients with a relapse of AP underwent minor papilla sphincterotomy (previous RAP group), or (recent RAP group) a second ERCP with: (1) extension of the previous papillary orifice (if judged inadequate because of difficulty or resistance to passage of a 3- to 5 Fr sphincterotome or catheter through the papillary orifice) and placement of a 7 Fr gauge, 3-7 cm long plastic pancreatic stent (SPSOF-7-3 to 7 or GPSO-7-3 to 7, Cook Medical, Winston-Salem, NC United States), scheduled to be left in place for one month and no longer (short-term stenting) to prevent post-procedural narrowing of the papillary opening and reduce the risk of long-term stent-induced pancreatic ductal changes; and (2) placement of a 7 Fr gauge, 3-7 cm long plastic pancreatic stent, scheduled to be left in place for three months then changed every three months for a year (long-term stenting) in cases not requiring extension of previous sphincterotomy.

When AP still recurred, a third ERCP was performed to place a 7 Fr gauge, 3-7 cm long stent if one was placed for only one month during the second procedure, or a 10 Fr gauge, 3-5 cm long stent (GEPD-10-3 or GEPD-10-5 Cook Medical, Winston-Salem, NC United States) if 7 Fr gauge long-term stenting had already been performed. The stent placed in the dorsal duct was scheduled to be left in place for three months, then changed every three months for a year.

Endotherapy was considered: (1) “successful” if there were no recurrences of AP; (2) “unsuccessful” if AP still recurred; and (3) “failed” if dorsal duct drainage was not achieved, including cannulation failure.

# *Statistical analysis*

Mean ± SD were used for continuous variables, percentages for categorical variables. Groups were compared using the *t*-test or Mann Whitney test for continuous variables and Fisher’s exact test for categorical variables. All differences were considered significant at a two-sided *P* value less than 0.05. Data were analyzed using the SPSS 17.0 system software (Chicago, IL, United States).

# RESULTS

Thirty-six patients entered the protocol study, 22 in the recent RAP group and 14 in the previous RAP group. There were no differences between the two groups in sex, age, number of episodes of AP and duration of the disease before enrollment (Table 1).

In 18/36 patients (50%), ss-MRCP detected dilation of the dorsal pancreatic duct, 12 at baseline and 6 after secretin stimulation. Twelve of these patients were candidates for endotherapy (8 with dorsal duct dilatation at baseline and 4 after secretin) and 6 for observation (4 with dorsal duct dilatation at baseline and 2 after secretin). No morphologic or functional abnormalities of the dorsal pancreatic duct were seen on ss-MRCP in the other 18 patients, ten of whom underwent therapeutic ERCP.

Minor papilla cannulation and sphincterotomy was successful in 20 of the 22 patients (90.9%) who underwent ERCP. The two cases in which cannulation failed were not included in the per protocol analysis.

***Follow-up***

Thirty-three patients were followed up for a mean of 4.5 ± 1.2 years; range: 2.0-6.7), 19/20 in the recent RAP group and all 14 in the previous RAP group. One patient in whom the second ERCP failed (after initial successful dorsal duct drainage) refused a further ERCP and was lost to follow-up and for this reason excluded in the per protocol analysis.

The mean follow-up time did not significantly differ between treated (4.3 ± 1.3 years) and untreated patients (4.7 ± 1.1 years). In all patients undergoing endotherapy, the mean duration of stenting was 1.17 year (range: 1 mo-2 years). The mean duration of the follow-up after retrieval of the last stent was 2.7 years (range: 1.5-4 years).

***Recurrences of acute pancreatitis***

In the 19 patients in whom dorsal duct drainage was achieved, AP still recurred in 11 (57.9%) after the first ERCP, in 6 after the second ERCP (31.6%) and in 5 after the third ERCP (26.3%). Overall, endotherapy was successful in 14 out of 19 patients (73.7%). There were no AP recurrences in the previous RAP group.

The five patients who still had further recurrences of AP after the third ERCP were followed-up by pancreatic EUS. One patient developed a pseudocyst and underwent pancreaticoduodenectomy. Except for this case, which was classified as severe AP, all the other relapses of pancreatitis were classified as mild disease.

Although the rate of AP recurrences after endotherapy was higher in patients without (37.5%) than in those with main pancreatic duct (MPD) dilatation (18.2%) at the time of enrollment, there was no statistically significant difference (*P* = 0.60) (Table 2).

Considering only the patients treated with long-term stenting, the recurrence rate of pancreatitis after retrieval of the last stent was 50% (5/10 patients).

***Complications***

A total of 58 ERCP procedures were carried out. Complications arose following fourteen of them (24.1%). Amongst the eight patients with major complications, seven developed mild AP (three after the first procedure) and one had bleeding. Minor complications were observed after six other procedures: pancreatic-type pain with normal serum amylase levels in three cases, and serum amylase levels more than three times the upper limit of normal without pain in the other three.

***EUS signs of chronic pancreatitis***

EUS investigation detected findings suggestive of CP in 20 of the 33 patients (60.6%) during follow-up, 11/19 (57.9%) amongst those in the recent RAP group and 9/14 (64.3%) in the previous RAP group (Figure 1): these rates were not significantly different. There were no significant differences in the prevalence of EUS signs suggesting CP when comparing patients with dilated and non-dilated dorsal pancreatic ducts within each group (Table 2). The mean duration of disease between the first attack of AP and the occurrence of EUS signs suggesting CP in the two groups was 6.1 ± 1.4 years ( and did not significantly differ between treated (5.7 ± 1.5) and untreated patients (6.4 ± 1.3).

In the 20 patients who developed EUS signs suggesting CP, the numbers (mean ± SD) of overall EUS criteria at enrolment (1.46 ± 0.52 *vs* 1.62 ± 0.74) and at the end of follow-up (4.46 ± 0.52 *vs* 4.25 ± 0.46) were not significantly different in the recent and previous RAP group, respectively. In each group, the number of EUS criteria detected at the end of follow-up was significantly higher (*P* < 0.001) than at enrollment; in all patients with EUS signs suggestive of CP, this increase was due to both ductal and parenchymal criteria. In these patients the most frequent EUS abnormalities were side branch dilation, hyperechoic MPD margins (ductal criteria), hyperechoic strands and foci (parenchymal criteria).

Amongst patients who underwent endotherapy, there were significantly fewer EUS signs of CP in the successful cases (6/14; 42.8%) than in the unsuccessful ones (5/5; 100%) (*P* =0.04) (Table 3).

EUS signs of CP occurred less frequently after successful endotherapy in patients with recent RAP than in those with previous RAP which did not undergo to endotherapy (42.8% *vs* 64.3%), but the differences were not statistically significant (*P* = 0.45).

Table 4 shows the rate of EUS signs of CP in patients undergoing pancreatic stenting: this was higher in patients undergoing long-term stenting (80%) than those with no stent or only short-term stenting (33.3%) and similar to the observation group (64.3%).

Two patients developed reduced exocrine function, one in the previous RAP group, the other in the recent RAP group (whose ERCP was unsuccessful).

# DISCUSSION

Although only a minority of patients with PDiv suffer life-long symptoms, this anatomical variant is found in up to 20% of patients with RAP[5,28,29]. It is not clear why these patients are at a higher risk for recurrent acute pancreatitis or whether their symptoms are etiologically related to PDiv[30]. In fact, genetic studies have suggested that as many as 10%-20% of patients with PDiv who have pancreatitis carry at least one allele of the *CFTR* gene product[31,32], or a higher frequency of *SPINK1* gene mutation[33], compared with healthy controls, suggesting a multifactorial origin of pancreatitis in these cases.

The obstructive hypothesis has led in the last few years to symptomatic PDiv patients being treated by endoscopic minor papilla sphincterotomy and/or dorsal duct stenting[18,34], which has proved as effective as surgical sphincteroplasty, according to a recent systematic review[35].

As regards the efficacy of successful minor sphincterotomy, it is not known if lowering intra-ductal pressure affects the evolution of CP in these patients.

Our prospective follow-up study of patients with PDiv and RAP without signs of CP aimed to evaluate, over a mean period of 4.5 years, the clinical outcome in those who had or did not have bouts of acute pancreatitis in the year preceding the study after endoscopic therapy or observation, respectively. All patients with relapses of pancreatitis after the enrollment underwent endoscopic therapy. The study also investigated morphologic and functional changes suggesting CP during the follow-up. To our knowledge, this is the first study with these issues up to date.

The endoscopic therapy was successful in approximately 73% of cases, as reported in a recent review[35]. The relapses of acute pancreatitis in patients in whom endotherapy was unsuccessful could have had other unknown causes, possibly involving the pancreatic parenchyma rather than the ductal system, since we excluded any patients with alcohol abuse or with the *CFTR*-gene, but not *SPINK 1*-gene mutations.

EUS is recognized as the best imaging method to obtain high-resolution images of the pancreas. It can detect features of CP in the pancreatic parenchyma and ducts that are not visible by any other imaging modality including ERCP and pancreatic exocrine function tests[36,37]. EUS findings suggestive of CP, according to the Wiersema criteria[25], were seen during the follow-up in 57.9% patients undergone endoscopic therapy and 64.3% in the observation group.

Considering that the mean duration of symptoms before enrolment in the study was approximately 3.5 years, changes consistent with CP occurred in these patients after they had had the disease for six years; this agrees with a previous report[38].

Overall, the frequency of EUS signs of CP was similar in both groups of patients. Dorsal duct dilatation did not predict the EUS findings suggestive of CP in either group, confirming once again that factors other than ductal dilation may be involved in chronic disease in these patients. However, among patients assigned to endotherapy there were significantly fewer CP findings in those with successful compared to unsuccessful ERCP (42.8% *vs* 100%, *P* = 0.04). In cases where endotherapy was successful, there was a lower rate of EUS signs of CP (42.8%) than in the observation group (64.3%), although the difference was not statistically significant. Possibly we investigated two different categories of patients with PD and RAP: one with inactive disease, with a history of recurrent pancreatitis but no episodes in the year before the study, and another with persisting pancreatitis at the time of enrollment, as a consequence of a still active inflammatory process. In the latter group, the ductal drainage obtained by endotherapy may have reduced or at least delayed the evolution toward CP, compared with cases after unsuccessful therapy, and gave results similar to those in the observation group, confirming a possible role of the outflow obstruction in the occurrence of EUS signs suggesting CP. These data are in favor of a therapeutic endoscopic approach in patients with PDiv and a history of recent episodes of pancreatitis, independently of the dorsal ductal dilatation, but they need to be confirmed in further studies because of the small number of patients evaluated and the relatively limited follow-up after the last stent retrieval.

A major problem related to long-term pancreatic ductal stenting is the occurrence of stent-related ductal changes similar to those observed in CP, already reported in previous studies, especially in cases with non-dilated ducts[39-41].

In our series, despite the frequent use of small, short stents, some ductal changes consistent with CP may have been induced by the long-term stenting. In fact, findings suggesting CP developed during follow-up in 33.3% of patients submitted to minor papilla sphincterotomy without or with short-term stenting, a lower rate than in patients with long-term stenting (80%). However, considering only patients with successful long-term stenting, CP developed in a similar proportion (60%) of untreated patients (64.3%). We do not know whether the combination of ductal and parenchymal lesions suggesting CP observed with EUS in patients with unsuccessful endotherapy depends on the course of an existing undetectable chronic inflammatory process involving the gland, rather than the long-term stenting, or both. This possibility is supported by evidence that up to 53% of patients in studies with idiopathic pancreatitis and PD have an underlying CP[19,42]. In these patients, CP may be the cause rather than the consequence of unsuccessful endotherapy.

In conclusion, this prospective study showed that: (1) in most patients with PDiv suffering from recent repeated episodes of pancreatitis endoscopic ductal drainage had a beneficial symptomatic effect independent of whether there was dorsal duct dilatation; (2) about 60% of patients with either recent (after endotherapy) or previous (observation) episodes of acute pancreatitis developed EUS findings consistent with CP over a six-year period; and (3) patients with recent bouts of acute pancreatitis in whom endotherapy was successful had a significantly lower risk of developing EUS findings consistent with CP than those treated unsuccessfully, but further studies are needed to confirm these results.

**COMMENTS**

***Background***

Pancreas divisum (PDiv) is the most common anatomical congenital variant of pancreatic ductal system. Its role in recurrent acute pancreatitis (RAP) and in chronic pancreatitis (CP) remains controversial.

***Research frontiers***

It is unclear if endoscopic therapy of pancreas divisum affects natural history of patients regarding the rate of recurrences and the evolution of CP.

***Innovations and breakthroughs***

It is the first study that evaluated endoscopic ultrasonography (EUS) sign suggesting CP in patient with PDiv and RAP submitted endotherapy.

***Applications***

The results of the present study confirmed that successful endoscopic treatment of pancreas divisum in patients with recurrent acute pancreatitis could reduce the rate of recurrence and it could also reduce occurrence of EUS signs suggesting CP.

***Terminology***Recurrent was defined as two or more episodes of abdominal pain associated with serum amylase and/or lipase levels at least double the upper normal limit are present. Endotherapy was considered: (1) “successful” if there were no recurrences of AP; (2) “unsuccessful” if AP still recurred; and (3) “failed” if dorsal duct drainage was not achieved, including cannulation failure.

***Peer review***

This paper has done a prospective study using EUS, but further studies are needed to confirm these results.

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**Table 1 Clinical and demographic data of 36 patients with pancreas divisum and acute recurrent pancreatitis at the time of the enrollment into the study**

|  |  |  |  |
| --- | --- | --- | --- |
|  | **Previous RAP**  **(14 pts)** | **Recent RAP**  **(22 pts)** | ***p* value** |
| Male/female | 6/8 | 7/15 | 0.72 |
| Age (yr), mean ± SD | 52.0 ± 12.6 | 55.6 ± 10.4 | 0.35 |
| Episodes of pancreatitis (mean ± SD) | 2.8 ± 1.1 | 3.4 ± 1.2 | 0.14 |
| Duration of the disease (yr), mean ± SD | 3.6 ± 1.9 | 3.4 ± 1.5 | 0.73 |

RAP: recurrent acute pancreatitis.

**Table 2 Follow-up results in 33 patients with pancreas divisum and previous and recent recurrent acute pancreatitisaccording to dilatation or non-dilatation of the main pancreatic duct detected at the time of the enrollment into the study *n* (%)**

|  |  |  |  |
| --- | --- | --- | --- |
|  | **Dorsal duct** | **AP recurrences** | **EUS signs of CP** |
| Previous RAP (14 pts) | Dilated (6 pts) | 0 | 4 (66.7)1 |
| Non-dilated (8 pts) | 0 | 5 (62.5) |
| Total | 0b | 9 (64.3)2 |
| Recent RAP (19 pts) | Dilated (11 pts) | 2 (18.2)3 | 7 (63.6)3 |
| Non-dilated (8 pts) | 3 (37.5) | 4 (50) |
| Total | 5 (26.3) | 11 (57.9)4 |

1dilated and non-dilated in the previous RAP group*: p* = ns; 2previous and recent RAP: *P* = NS; b*p* < 0.05, endotherapy *vs* main pancreatic duct dilatation; 3dilated and non-dilated in the recent RAP group: *p* = ns; 4all five patients with AP recurrences during follow-up developed CP. AP: acute pancreatitis; EUS: endoscopic ultrasonography; CP: chronic pancreatitis; RAP: recurrent acute pancreatitis; NS: Not significant.

**Table 3 Demographics, characteristics of the disease and endoscopic ultrasonography signs of chronic pancreatitis development in patients assigned endotherapy able to achieve dorsal duct drainage *n* (%)**

|  |  |  |  |
| --- | --- | --- | --- |
|  | **Successful endotherapy1**  **(14 pts)** | **Unsuccessful endotherapy2**  **(5 pts)** | ***P* value** |
| Male/female | 11/3 | 4/1 | 1 |
| Age (yr), mean ± SD | 55.3 ± 7.9 | 55.0 ± 21.1 | 0.5 |
| Number of episodes of AP before study, yr, mean ± SD | 3.4 (1.3) | 3.4 (1.1) | 0.8 |
| Duration of ARP before enrollment (yr), mean ± SD | 3.2 (1.6) | 3.8 (1.3) | 0.3 |
| Duration of post-enrollment follow-up (yr),  mean ± SD | 4.1 (1.3) | 4.8 (1.1) | 0.7 |
| EUS signs of CP | 6 (42.8) | 5 (100) | 0.04 |
| Duration of disease between the first attack of AP and the diagnosis of CP (yr),  mean ± SD | 5.6 (1.3) | 6.1 (1.7) | 0.7 |

1successful endotherapy: no recurrences of AP in the follow up after one to three ERCP sections; 2unsuccessful endotherapy: recurrences of AP after three ERCP sections. AP: acute pancreatitis; CP: chronic pancreatitis; ERCP: endoscopic retrograde cholangiopancreatography.

**Table 4 endoscopic ultrasonography signs of chronic pancreatitis in patients with pancreas divisum and previous or recent recurrent acute pancreatitisin relation to the pancreatic stenting *n* (%)**

|  |  |  |
| --- | --- | --- |
| **Group** | **Stenting** | **EUS signs of CP** |
| Previous RAP (14 pts) | 0 | 9 (64.3) |
| Recent RAP (19 pts) | No/short-term (9 pts) | 3 (33.3)1 |
| Long-term (10 pts) | 8 (80)2 |
| Total | 11 (57.9) |

1no/short term stenting and long-term stenting: *p* = 0.07; 2Three in 5 successful stenting (60%); 5 in 5 unsuccessful stenting.EUS: endoscopic ultrasonography; CP: chronic pancreatitis.

36 patients

14 previous RAP conservatively managed

22 recent RAP

ERCP in all patients

20/22 successful minor ES (1st ERCP)

ERCP only in case of AP relapse

0 patients

underwent ERCP

ductal drainage was achieved

in 19 out of 20 patients after the 2nd or 3rd ERCP

EUS signs of CP in 9 (64.3%)

2unsuccessful ERCP in 5 pts EUS signs of CP in 5 (100%)

1successful ERCP in 14 pts

EUS signs of CP in 6 (42.8%)

**Figure 1 Occurrence of endoscopic ultrasonography signs indicating chronic pancreatitis in the follow-up of patients with pancreas divisum and recent or previous recurrent acute pancreatitis.** 1successful if dorsal duct drainage was achieved and there were no recurrences of AP; 2unsuccessful if dorsal duct drainage was achieved but AP still recurred. RAP: recurrent acute pancreatitis; ERCP: endoscopic retrograde cholangiopancreatography; AP: acute pancreatitis; EUS: endoscopic ultrasonography; CP: chronic pancreatitis.