Pathological Acute Inflammation in Chronic Pancreatitis

Pavankumar Vijayaraj¹, Biju Pottakkat¹, Raja Kalayarasan¹, Sandip Chandrasekar¹, Surinder Kumar Verma²

¹Department of Surgical Gastroenterology, Jawaharlal Institute of Post graduate Medical Education and Research (JIPMER), Puducherry, India
²Department of Pathology, Jawaharlal Institute of Post graduate Medical Education and Research (JIPMER), Puducherry, India

ABSTRACT

Background Mechanism of pain in Chronic Pancreatitis is not fully understood. Recurrent acute inflammation is one of the proposed hypotheses for pain in chronic pancreatitis. The actual incidence and prevalence of ongoing acute inflammation in chronic pancreatitis patients have been under-noticed in literature. This study aims to examine the prevalence of acute pancreatic inflammation in chronic pancreatitis patients. Methods Fifty patients who underwent surgery for chronic pancreatitis were analyzed. Those with clinical, biochemical or radiological features of acute on chronic pancreatitis were excluded. Intra operative fine needle aspiration cytology was taken from the head and body of pancreas. Pancreatic tissue was sent for histopathological examination in all patients. Results Intraoperative fine needle aspiration cytology from pancreas showed features of acute inflammation in twenty three (46%) cases. Biopsy from pancreas showed features of acute inflammation in 12 patients (24%). 30/50 (60%) patients had some features of acute inflammation in either fine needle aspiration cytology or biopsy. Conclusion Significant proportion of patients with chronic pancreatitis has pathological features of acute inflammation in pancreas despite clinical, biochemical and radiological features showing no evidence of acute pancreatitis. Ongoing pathological acute inflammatory process in the pancreas might be a major cause for initiation and progression of chronic fibrosis in chronic pancreatitis.

INTRODUCTION

Chronic pancreatitis (CP) is a condition characterized by progressive chronic inflammation of the pancreas and its clinical manifestations may vary from patient to patient. Abdominal pain is the most common and significant symptom in CP and is seen in up to 90% of patients. The mechanism of pain in chronic pancreatitis is not fully understood but is probably multifactorial [1]. Recurrent acute inflammation can develop in a chronically diseased pancreas, and this can cause severe pain in patients with CP [2]. The activated enzymes and other injurious substances released during the acute inflammatory process might be responsible for the initiation of pain. Also, there is a possibility that this acute inflammation may be occurring continuously in a chronically inflamed gland until the fibrosis process sets in. The process may be happening irrespective of the clinical manifestation of acute pancreatitis. The actual incidence and prevalence of ongoing acute inflammation in CP have been under-noticed in the literature. [2]

Specimen from the pancreas is required to better understand the incidence and pattern of inflammation in chronic pancreatitis. The pancreas is not a safe organ for percutaneous tissue biopsy given its location deep in the abdomen. Adequate tissue for histological examination can be only obtained in patients who undergo surgical procedures. The pathological study of the cored out tissue obtained for biopsy from the pancreas in Frey’s procedure can be studied but there is a chance that the tissue tends to get distorted and charred owing to handling and usage of electrocautery in the pancreas. Intra-operative fine needle aspiration cytology (FNAC) of the pancreas is commonly used to determine if a lesion is malignant and to determine the type of malignancy. Studies have shown the sensitivity, specificity, positive and negative predictive values, and diagnostic accuracy for intraoperative FNAC comparable to that of biopsy [3, 4, 5]. FNAC has the advantage of studying the cells from a well perfused and unhandled pancreas, but also has the disadvantage of low yield in case of a firm to hard gland with more fibrosis and less cellularity. In this observational study, we attempt to examine the prevalence of acute pancreatic inflammation in patients with CP undergoing surgery using both intraoperative FNAC and histopathological examination of the operative specimen from the pancreas.
MATERIALS AND METHODS

Our study was a prospective observational study performed in 50 consecutive patients who underwent surgery for CP from October 2013 in the Department of Surgical Gastroenterology, Jawaharlal Institute of Postgraduate Medical Education and Research (JIPMER). Ethical clearance was obtained from the Institute ethics committee. A pancreatic protocol contrast enhanced computed tomography (CECT) of abdomen was done in all patients to confirm the diagnoses, assess the characteristics of pancreas and to rule out CP related complications. All proven cases of CP with pain not controlled on medications were included in the study. Those patients with associated pancreatic malignancy or clinical acute on chronic pancreatitis with acute fluid collection on imaging were excluded. Demographic data were collected from all patients at the time of admission, including age, sex, body mass index, history of alcoholism, habit of smoking, clinical presentation, detailed history of abdominal pain, associated diabetes mellitus or steatorrhoea. All patients underwent hemogram, renal and liver function tests. Serum amylase levels were assessed the day prior to surgery. Frey’s procedure was the surgery of choice in all patients except those with associated portal hypertension where lateral pancreatico jejunostomy (LPJ) was the procedure performed. Distal pancreatectomy and hepaticojejunostomy were added in patients with tail predominant disease and concurrent biliary stricture respectively. Izbicki procedure was the preferred procedure in patients with small duct disease (duct diameter<4 mm). Intra operative FNAC was taken using a 21-gauge needle from the head and body of the pancreas after adequate exposure of the pancreas. Detailed intra operative assessment of the pancreas including pancreatic texture, volume, calcifications, main pancreatic duct (MPD) diameter and evidence of associated features like portal hypertension, biliary stricture, pseudoaneurysm, pancreatic head mass and pseudocysts were made and findings recorded. Cored out tissues from the pancreatic head were sent for histopathological examination. Patients with evidence of malignancy on histopathology were excluded from the study. FNAC smears were stained with Papanicolaou (PAP) and May Grunwald Giemsa (MGG) stains. All the FNAC slides were examined by a single pathologist with 26 years of experience. Aspirates showing evidence of neutrophilic infiltrates were taken as features suggesting acute inflammation on chronic pancreatitis. Histopathological slides were also reviewed by the same pathologist to look for features of acute inflammation (Figures 1, 2 and 3).

Statistical Analysis

The data was entered and analyzed in Statistical Package for the Social Sciences (SPSS) version 16. FNAC and histopathological findings were summarized as proportions.

Figure 1. Photomicrographs of FNAC from pancreas showing neutrophilic infiltrate (thick arrow) scattered around amidst the background of chronic inflammatory cells (thin arrow) [MGG 10X].

Figure 2. Photomicrograph of Papanicolaou stained FNAC smears showing plenty of neutrophils (thick arrow) (PAP 20X).

Figure 3. Histopathological picture showing giant cell reaction (thick arrow) surrounded by fibrosis (angled arrow) nearby a polymorphonuclear infiltrate (thin arrow) suggestive of acute inflammation in a setting of chronic pancreatitis (Hematoxylin and Eosin 20X).
RESULTS

During the study period, 50 patients (36 (72%) males and 14 (28%) females) underwent surgery for chronic pancreatitis. None of the patients had either a recent acute pain episode or elevated serum amylase levels. Median age of the group was 37 (13-68) years. 25 (50%) patients had history of alcoholism and 26 (51%) patients were smokers. 27 (54%) patients were diabetics, and hypoalbuminemia (serum albumin <3.5 gm/dl) was noted in 9 (18%). Median BMI of the patient group was 18.4 (10.5-26.5 kg/m²). All patients had calcifications in their pancreas and 14 (28%) had chronic pseudocysts. 46/50 patients underwent Frey's procedure. Of those 46 patients, additional procedures like distal pancreatectomy, spleen preserving distal pancreatectomy and Roux-en-Y hepaticojejunostomy was done in two, one and six patients respectively. Lateral pancreaticojejunostomy was done in three patients. One patient with small duct disease underwent Izbicki's procedure. Intra operative FNAC was taken in all the patients from the head and body of pancreas before proceeding with the dissection in the pancreas. Pancreatic tissue was taken for biopsy in all patients. The consistency of the pancreas (assessed by the operating surgeon) was found to be firm to hard, firm and soft in 17 (34%), 28 (56%) 5 (10%) patients respectively. Lateral pancreaticojejunostomy was done in three patients. One patient with small duct disease underwent Izbicki's procedure. Intra operative FNAC was taken in all the patients from the head and body of pancreas before proceeding with the dissection in the pancreas. Pancreatic tissue was taken for biopsy in all patients. The consistency of the pancreas (assessed by the operating surgeon) was found to be firm to hard, firm and soft in 17 (34%), 28 (56%) 5 (10%) patients respectively.

Table 1 summarises the findings from FNAC and biopsy. None of the patients had any features of malignancy in either FNAC or histopathology. Out of the study population of 50 patients, 23 patients showed features of acute inflammation in FNAC. Of these 23 patients, five patients showed features of acute inflammation in biopsy as well, while the remaining eighteen patients didn’t show any features of acute inflammation in biopsy. Out of the study population of 50 patients, 12 patients showed features of acute inflammation in biopsy. Of these 12 patients, five patients showed features of acute inflammation in FNAC as well, while the remaining seven patients didn’t show any features of acute inflammation in FNAC. Hence in our study, 30/50 (60%) patients had some feature of acute inflammation in FNAC or biopsy (18 and 7 had acute inflammation in FNAC and biopsy alone respectively and 5 had those features in both). Twenty (40%) patients showed no evidence of acute inflammation.

DISCUSSION

CP is an ongoing chronic inflammatory process of the pancreas leading to the destruction of parenchyma and ducts with development of fibrosis [6]. The pathological hallmarks of chronic pancreatitis are diffuse or focal parenchymal destruction with damage to the acini and islet cells, chronic inflammatory cell infiltration, loss of normal architecture and progressive fibrosis [7]. Multiple studies in the last four decades have observed that recurrent acute pancreatitis leads to CP [8, 9, 10].

Sarles et al. [11] in their study in 1965 in a set of patients with acute and chronic pancreatitis observed that acute and chronic pancreatitis are two different entities pathologically and acute form rarely progressed to CP. They showed that though both AP and CP can be caused by alcoholism, CP patients tend to be younger than their AP counterparts. Also, they did a careful histological study of the wall of the pseudocyst in CP and showed that the cysts arose not following an acute necrosis of the pancreas but by distension and a blowing out of foci of localized acinar-canalicular dilatation secondary to ductal obstruction. From then onwards multiple studies are debating the pathophysiology of CP and whether recurrent acute inflammation leads to chronic pancreatitis or not.

Ammann and Kloppel in their prospective long-term clinico-morphological study on alcoholic CP correlated the clinical findings to the biopsy of the pancreas at early and advanced CP and showed that there is an evolutionary change from AP to CP with necrosis pre-dominating the acute events while fibrosis sets in finally [12]. Their study showed that morphological features of AP like necrosis and necrotic pseudocysts dominate in early stages while fibrosis paralleled by increasing impairment of pancreatic function was the characteristic findings in the late stages. Morphological findings in early and late ACP suggest that repeated and severe attacks of acute pancreatitis lead to the progression of the disease to CP supporting the necrosis-fibrosis hypothesis. Another study by Kloppel et al [2] on morphology of pseudocysts in CP showed that if peripancreatic fat necrosis become confluent and extensive, as seen in severe acute pancreatitis, pseudocysts will form, while organization of the smaller foci of intrapancreatic fat necrosis that occurs along the interlobular spaces of the pancreas occasionally only leads to pseudocysts but most likely induces fibrosis. A histological study by de Angelis et al. [13] on alcoholic and non-alcoholic CP has shown that inflammatory cells predominantly lymphocytes and macrophages were concentrated at the periphery of a damaged perineurium suggesting a role for inflammatory cells in damaging the perineurium and generating pain. In an autopsy study which attempts to point out classical features of CP to avoid overdiagnosis of CP in the biopsy, Shimizu et al. [14] have shown that a feature of acute inflammation like neutrophil and polymorphonuclear cell

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<tr>
<th>Pathological findings</th>
<th>FNAC-n*(%)</th>
<th>Histopathology-n*(%)</th>
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<tr>
<td>Acute on chronic inflammation</td>
<td>23 (46)</td>
<td>12 (24)</td>
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<tr>
<td>Chronic inflammation only</td>
<td>15 (30)</td>
<td>34 (68)</td>
</tr>
<tr>
<td>Ductal/glandular cells only and no inflammatory cells</td>
<td>2 (4)</td>
<td>3 (6)</td>
</tr>
<tr>
<td>Hemorrhagic/non-diagnostic/ distorted and not suitable for study</td>
<td>10 (20)</td>
<td>1 (2)</td>
</tr>
<tr>
<td>Total</td>
<td>50</td>
<td>50</td>
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*Total number of patients – 50; and 30/50 (60%) patients had some feature of acute inflammation in FNAC or biopsy
infiltrates in the periductal region exists in CP and they may suggest recently exacerbated CP. It has been noted in many autopsy series, and operative biopsy studies of CP that a polymorphonuclear cell infiltrate exists in many cases of CP, and it is attributed to recent acute crises in CP [2, 8, 9, 13, 15]. Exact prevalence of acute inflammation in CP glands has not been studied in detail. Moreover, whenever features of acute inflammation were detected in biopsies of tissues from chronic pancreatitis, they have been attributed to a recent acute episode occurring in CP. Our study aimed to study the prevalence of acute inflammation in CP patients who neither had a recent acute pain episode nor elevated serum amylase levels. We were able to detect features of acute inflammation (in either FNAC or biopsy) in around 60% of our patients. This is significantly high compared to the incidence reported in previous operative biopsy studies [2, 8, 13]. Our study supports the finding noted by Ammann and Kloppel [12] that a pathological acute inflammation occurs continuously in a chronically inflamed gland until the fibrosis process sets in. Our study also shows that this acute inflammatory process may be happening irrespective of the external manifestation of an acute episode in a patient, a finding that has not been reported in literature. Twenty percent of our FNACs came as non-diagnostic possibly because of the difficulty in retrieving the cells using a fine needle from the firm to hard pancreas which is mostly constituted by thick fibrotic tissue and less cellularity. Also, the biopsy specimen from the pancreas showed lesser proportion of acute inflammatory features compared to the FNAC samples. Frey’s procedure is done bit by bit coring of the pancreatic tissue using high energy electrocoagulation which usually results in charring of the pancreatic tissue. We feel that this might have affected the interpretation of acute inflammation in biopsy specimens.

CONCLUSION

Significant proportions of patients with CP have pathological features of acute inflammation in pancreas despite clinical, biochemical and radiological features showing no evidence of acute pancreatitis. Ongoing pathological acute inflammatory process in the pancreas might be a major cause for initiation and progression of chronic fibrosis and chronic pancreatitis.

**REFERENCES**


