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Inflammatory Bowel Disease and Primary Sclerosing Cholangitis

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1. Introduction

Inflammatory bowel disease (IBD) is a chronic condition which is characterized by recurrent immune-mediated inflammation of the gastrointestinal system. IBD is frequently associated with extraintestinal manifestations (EIM) characterized by involvement of multiple organs. EIM occur in 21% to 47% of IBD patients (Navaneethan & Shen, 2010). While some extraintestinal manifestations are encountered more frequently in Crohn’s disease (CD) than in ulcerative colitis (UC), some are encountered equally in CD and UC. While the degree of involvement of the skin, eyes, and joints is parallel to disease activity, hepatobiliary and pulmonary involvement is independent of disease activity and intestinal inflammation (Greenstein AJ, et al., 1996).

Hepatopancreatobiliary (HPB) manifestations are the most frequently encountered EIM in patients with IBD. They can be encountered in various ways:

1. HPB conditions sharing the same pathological mechanisms with IBD (Primary Sclerosing Cholangitis (PSC), small-duct PSC/pericholangitis, and PSC/autoimmune hepatitis overlap, IBD associated acute or chronic idiopathic pancreatitis)
2. HPB conditions that reflect the degree of pathophysiologic damage seen in IBD (Cholelithiasis and portal vein thrombosis)
3. HPB conditions related to side effects of drugs used in treatment of IBD (Drug induced or associated hepatitis, pancreatitis, cirrhosis, Hepatitis B reactivation, hepatosplenic T cell lymphoma)
4. HPB conditions possibly related to IBD (Autoimmune pancreatitis, IgG4-associated cholangitis, fatty liver, hepatic amyloidosis, granulomatous hepatitis, primary biliary cirrhosis) (Navaneethan & Shen, 2010)

The various HPB manifestations and their associations are summarized in Tables 1 and 2. The aim of this chapter is to cover in detail primary sclerosing cholangitis, which is an important and frequently encountered HPB manifestation of ulcerative colitis.

2. The association of Inflammatory Bowel Disease and Primary Sclerosing Cholangitis

PSC is a chronic cholestatic hepatobiliary disease that often develops in the setting of IBD and which affects predominantly young to middle-aged patients (Olsson et al., 1991).
HBP manifestations with a possibly shared pathogenesis and mechanism as IBD | Primary sclerosing cholangitis (PSC)  
Small-duct PSC  
Cholangiocarcinoma  
Autoimmune hepatitis/PSC overlap  
IgG4 associated cholangitis  
Acute and chronic idiopathic pancreatitis

HPB manifestations parallel the pathophysiology associated with IBD | Gallstones  
Portal vein thrombosis and hepatic abscess

HPB manifestations associated with treatment of IBD | Drug induced hepatitis (azathioprine, 6-mercaptopurine, methotrexate, cyclosporine, infliximab)  
Reactivation of hepatitis B (infliximab)  
Drug induced pancreatitis (azathioprine, 6-mercaptopurine), Hepatosplenic T-cell lymphoma

HPB manifestations possibly associated with IBD | Fatty liver  
Hepatic amyloidosis  
Granulomatous hepatitis  
Primary biliary cirrhosis  
Autoimmune pancreatitis

Table 1. Association Between Inflammatory Bowel Disease (IBD) and Hepatopancreatobiliary (HPB) Manifestations. Adapted from Navaneethan & Shen, 2010

| HPB manifestation | Ulcerative Colitis | Crohn’s Disease |
|------------------|--------------------|-----------------|
| Primary sclerosing cholangitis (PSC) | ++ | +(colonic or ileocolonic) |
| Small duct PSC | ++ | + |
| Cholangiocarcinoma | ++ | + |
| Autoimmune hepatitis/PSC overlap | ++ | + |
| IgG4-associated cholangitis | ++ | + |
| Acute and chronic pancreatitis | + | ++ |
| Gall stones | - | ++ |
| Portal vein thrombosis and hepatic abscess | + | ++ |
| Drug induced hepatitis | ++ | ++ |
| Reactivation of hepatitis B (infliximab) | ++ | ++ |
| Drug induced pancreatitis | + | ++ |
| Hepatosplenic T-cell lymphoma | +/- | + |
| Autoimmune pancreatitis | ++ | + |
| Fatty liver | ++ | ++ |
| Hepatic amyloidosis | - | ++ |
| Granulomatous hepatitis | - | ++ |
| Primary biliary cirrhosis | ++ | + |

Table 2. HPB Manifestations Associated with IBD. Adapted from Navaneethan & Shen, 2010
The association of primary sclerosing cholangitis (PSC) and ulcerative colitis (UC) was first reported by Smith and Loe in 1965 (Smith & Loe, 1965). The incidence of IBD in patients with PSC is 25-30%. Increasing awareness of association of PSC and UC led to more widespread use of endoscopic retrograde cholangiopancreatography (ERCP) and hence more cases were diagnosed with PSC (Broome & Bergquist, 2006).

The association of PSC and Crohn’s disease, which was first described by Atkinson and Carroll in 1964 (Atkinson & Carroll, 1964), is relatively rare. The incidence of CD in PSC varies between 1.3-14% (Wiesner & LaRusso, 1980; Chapman et al., 1980; Rasmussen et al., 1997; McGarity et al., 1991; Faubion et al., 2001; Loftus et al., 2005; Tobias et al., 1983). The most eminent finding is rectal sparing. Colonic stricture and perianastomotic ulcers are relatively rare in this patient group. Patients with PSC and CD almost always have extensive colitis or ileocolitis, but never have isolated ileitis (Broome & Bergquist, 2006).

About 85-90% of patients with PSC and IBD are comprised of UC patients and the remainder are comprised of patients with Crohn’s colitis or Crohn’s ileocolitis (Olsson et al., 1991).

IBD can be diagnosed at any time throughout the course of PSC, and PSC can develop at any time throughout the course of IBD (Fausa et al., 1991; Broome et al., 1990). However, IBD is often diagnosed many years before the diagnosis of PSC. PSC can occur many years after proctocolectomy for colitis; IBD can be diagnosed many years after liver transplantation for advanced PSC (Fausa et al., 1991; Wiesner & LaRusso, 1980; Aadland et al., 1987; Chapman et al., 1980; Riley et al., 1997). As IBD and PSC can be asymptomatic, the time of diagnosis depends on the diagnostic alertness of the physician (Broome & Bergquist, 2006).

PSC has a variable natural course. Nevertheless, PSC is typically characterized by progressive inflammation, obliterative fibrosis, damage to the intrahepatic and extrahepatic biliary tree and eventually biliary fibrosis, cirrhosis and finally liver failure. PSC is often diagnosed between the 3rd and 5th decades and male to female gender ratio is 2:1 (Chapman et al., 1980; Lee & Kaplan, 1995; Navaneethan & Shen, 2010). Freeman et al. observed that patients with PSC had more extensive endoscopic and histological inflammation of the afferent limb who had restorative proctocolectomy with ileal pouch-anal anastomosis (IPAA) for UC than those with no concurrent PSC (Freeman et al., 2008).

A report from Mayo Clinic has identified characteristic clinical, endoscopic, and histological findings in IBD accompanied by PSC as quiescent colitis, substantial preclinical phase, pancolitis, rectal sparing, backwash ileitis, pouchitis, and colorectal dysplasia/carcinoma (Loftus et al., 2005) (Table 3).

| Clinical, Endoscopic, and Histological Findings that Characterize IBD-PSC |
|---------------------------------------------------------------|
| Quiescent colitis                                            |
| Substantial preclinical phase                                 |
| Pancolitis                                                    |
| Rectal sparing                                                |
| Backwash ileitis                                             |
| Pouchitis                                                     |
| Colorectal dysplasia/carcinoma                                |

Table 3. Clinical, Endoscopic, and Histological Findings that Characterize IBD-PSC.
Adapted from Broome & Bergquist, 2006
2.1 Prevalence of IBD in patients with PSC

The prevalence of UC in patients with PSC varies from country to country in the range of 21-80% (Takikawa & Manabe, 1997; Bergquist A et al., 2002). The prevalence of PSC in patients with Crohn’s disease varies between 1.4-3.4% (Rasmussen et al., 1997; Schrumpf et al., 1980; Shepherd et al., 1983). Colitis in PSC is often quiescent or mild. Therefore, all patients with PSC should undergo colonoscopy and multiple biopsies should be taken to estimate the true prevalence (Broome & Bergquist, 2006). An algorithm for screening for IBD in patients with PSC is given in Figure 1.

Fig. 1. Screening for IBD in patients with PSC.
Adapted from Navaneethan U, Shen B, 2010, and Broome U, Bergquist A, 2006

2.2 Prevalence of PSC in patients with IBD

The prevalence of PSC in patients with IBD and persistently abnormal liver function tests is 2.4-7.5% (Olson R, et al., 1991; Aadland E et al., 1987). PSC prevalence is 5.5% in patients with substantial colitis, and 0.5% in patients with distal colitis (Olsson et al., 1991). 1.3 to 14% of patients with Crohn’s disease have PSC (Wiesner & LaRusso 1980; Chapman et al., 1980; Rasmussen et al 1997; McGarity et al., 1991; Faubion et al., 2001; Loftus et al., 2005).

PSC can present as an asymptomatic disease characterized by mild increases in aminotransferases, episodes of normal liver function tests can also occur. The frequency of liver function test screening is decisive in determining the prevalence of this disease. However, there may also be liver enzyme elevations due to autoimmune hepatitis, fatty liver, colonic disease activation, total parenteral nutrition and steroid use (Loftus et al., 1997; Broome et al., 1994). Therefore, liver function tests at presentation may be misleading. Ideally, liver function tests must be assessed when the colonic disease is in the inactive phase (Broome & Bergquist, 2006). An algorithm for PSC screening in a patient with IBD is presented below in Figure 2.

Enlargement of perihepatic lymph nodes is a common finding in PSC (Outwater et al., 1992). The presence of enlarged perihepatic lymph nodes in a patient with UC should alert the
clinician for PSC. A prospective study by Hirche et al. that involved 310 IBD patients showed that the detection of enlarged perihepatic lymph nodes by ultrasonography (US) was a better predictor of PSC when compared to serum parameters alone (Hirche et al., 2004).

Rabinowitz et al., who investigated patients with advanced PSC, found that patients with PSC and UC are predominantly male and the first presentation is often mild liver enzyme elevation. Bile duct involvement is different among patients with PSC plus UC and those with PSC only. While the prevalence of combined intrahepatic and extrahepatic bile duct strictures is 82% in patients with PSC and UC, it is 46% in those with only PSC. Isolated extrahepatic duct involvement is 38% in PSC, and 7% in those with PSC and UC (Rabinowitz et al., 1990). Among 305 Swedish PSC patients, no difference between patients with and without IBD could be found (Broome et al., 1996). Currently, we do not have enough data to arrive at a conclusion that PSC in patients without IBD is an entity different from PSC that accompanies IBD.

Rectal sparing and backwash ileitis are more common in patients with UC and PSC (Loftus et al., 2005). As rectal sparing is common in these patients, rectosigmoidoscopy is not
adequate in demonstrating the association of UC and PSC, a colonoscopy is often recommended (Faubion et al., 2001; Perdigoto et al., 1991). The clinical course of colitis is variable in patients who undergo OLT for PSC. MacLean et al have reported clinical healing in one third of patients, unchanged clinical course in one third, and worsening in the last one third (MacLean et al., 2003).

In UC patients operated on with an ileal pouch-anal anastomosis, nonspecific inflammation of the pouch (pouchitis) is the most frequent long-term complication. Chronic pouchitis is more common in patients with PSC and UC when compared to those with UC only (60% vs 15%) (Penna et al., 1996). Pouchitis continues to occur after OLT in patients with PSC (Zins BJ et al., 1995).

3. Impact of coexisting PSC on the disease behavior and course of IBD

Patients with PSC and IBD have a different clinical course when compared to IBD patients not complicated with PSC, and this patient group has been defined by the Mayo clinic as a different clinical entity (Loftus et al., 2005). While UC can be diagnosed many years after PSC diagnosis and even after orthotopic liver transplantation (OLT), a patient with UC, who had a proctocolectomy or not, can be diagnosed with PSC many years later (Joo et al., 2009). Patients with UC and PSC more frequently experience rectal sparing, backwash ileitis, pancolitis, colorectal neoplasia when compared to patients with UC alone, and their prognosis is worse (Faubion et al., 2001; Loftus et al., 2005; Loftus et al., 1997; Penna et al.; 1996; Heuschen et al., 2001; Lundkvist & Broome, 1997; Broome et al., 1995; Soetikno et al., 2002). However, there are studies reporting different results. A recent case-control study found no difference between patients with PSC plus IBD and IBD alone in terms of rectal sparing. The same investigators found similar backwash ileitis prevalences in the PSC plus UC group and the UC alone group (Joo et al., 2009).

It was suggested that patients with PSC and UC may more likely run a quiescent course of colitis than UC patients without coexistent PSC. PSC-UC patients have colitis with a lower histopathological grade of inflammation (Joo et al., 2009). In a study from Sweden that included 76 patients with PSC, 7 of 11 asymptomatic patients had a histopathological diagnosis of IBD, and 2 of these 7 patients had colonic dysplasia. In the follow-up, these 2 patients died before developing any IBD symptoms while 3 patients developed IBD symptoms (Broome et al., 1995).

A colonoscopy must be performed at the time of PSC diagnosis for IBD screening (Faubion et al., 2001). Currently, there is no guideline about the necessity and frequency of colonoscopic screening in asymptomatic patients with a normal baseline screening colonoscopy. However, patients with PSC and UC/Crohn’s colitis should undergo annual colonoscopy (Kornbluth & Sachar, 1997). The risk of pouchitis after restorative proctocolectomy is higher in patients with PSC and UC when compared to patients with UC alone (Faubion et al., 2001; Loftus et al., 2005; Penna et al., 1996). On the other hand, there is no correlation between the severity of liver disease and the risk of pouchitis (Penna et al., 1996).

While there are conflicting reports in the literature, it is generally agreed upon that the risk of IBD-associated colonic dysplasia is increased in the presence of PSC. In a meta-analysis, the presence of PSC was an independent risk factor for colorectal dysplasia/cancer in UC patients with an odds ratio of 4.79 (Soetikno et al., 2002). Screening colonoscopy is associated with a survival benefit in patients with PSC and UC or Crohn’s colitis (Rutter et
al., 2004), and a yearly colonoscopy is recommended beginning from the time of PSC diagnosis (Kornbluth & Sachar, 1997). The risk for dysplasia seems to remain high after colectomy in patients with PSC and UC (Ståhlberg et al., 2003). According to histological and flow cytometry studies, atrophy in the pouch mucosa, dysplasia, and DNA aneuploidia are more common in PSC plus UC patients with IPAA when compared to patients with UC only with IPAA (Ståhlberg et al., 2003).

4. The impact of coexisting IBD on the disease behavior and course of PSC

In a liver biopsy study that compared PSC patients with and without accompanying IBD no difference was found in terms of liver histopathology (Ludwig et al., 1981). Two other studies were unable to define any specific clinical or radiological criteria to distinguish patients with UC plus PSC from those with PSC alone (Broome et al., 1996; MacCarty et al., 1985). On the other hand, Rabinovitz et al showed that the first sign of liver disease was often liver enzyme abnormality in PSC accompanied by IBD. In contrast, liver disease more often presented with jaundice, pruritus, and fatigue in patients with PSC alone (Rabinowitz et al., 1990).

5. Pathogenesis

As IBD tends to coexist with other autoimmune diseases such as Type I diabetes mellitus and Graves’ disease (Saarinen S et al., 2000), genetic and immunological mechanisms have been extensively studied (Navaneethan & Shen, 2010). HLA-B8, HLA-DRB1*0301(DR3), HLA-DR3*0101(DRw52a), and HLA-DRB1*0401(DR4) are among genetic variants that are associated with susceptibility to IBD (Farrant et al., 1992; Olerup et al., 1995). Many antibodies have been detected in PSC patients with varying prevalences, including antinuclear antibodies (24-53%), smooth muscle antibodies (13-20%), anti-perinuclear cytoplasmic antibody (65-88%) (Mulder et al., 1993; Bansi et al., 1996; Terjung & Spengler, 2005; Terjung et al., 2000). However, in contrast to other autoimmune diseases, PSC is more frequently encountered in male subjects, and it does not respond to immunosuppressive therapy (O'Mahony & Vierling, 2006). One hypothesis that aims to explain the increased incidence of PSC in IBD patients postulates that the transportation of bacterial endotoxins from the inflamed colonic mucosa to the liver via the portal circulation stimulates the Kupffer cells (Fausa et al., 1991; Aoki et al., 2005). However, another study showed that there is no evidence of altered intestinal permeability or bacterial overgrowth in PSC patients (Björnsson et al., 2000). Other factors that are suspected to play a role in pathogenesis are ANCA autoantigen Beta-tubulin isotype 5 (TBB5) that shares epitopes with microbial antigens and human autoantigens (Erickson, 1995) and leaky gut (Terjung & Spengler, 2009).

6. Diagnosis

Most PSC patients are asymptomatic at the time of diagnosis. Fatigue, pruritus, jaundice, abdominal pain, and weight loss are the presenting symptoms in 10-15% of the patients (Tischendorff et al., 2007; Wiesner et al., 1989). A cholestatic pattern of abnormal liver function tests in the form of elevation in alkaline phosphatase is a biochemical feature for PSC (Rasmussen HH et al., 1997; Talwalkar & Lindor, 2005). The most frequent cause of persistent liver enzyme elevation in a patient with IBD is PSC (Heikius et al., 1997). PSC is also the most frequent cause of persistent liver enzyme elevation in an IBD patient with
proctocolectomy (Navaneethan et al., 2009). Various antibodies may be detected with varying prevalences (antinuclear antibodies 24-53%, antismooth muscle antibodies 13-20%, p-ANCA 65-88%). Nevertheless, the role of these antibodies in diagnosis and discrimination between isolated PSC and PSC accompanied by IBD remains unclear (Mulder et al., 1993; Bansi et al., 1996; Terjung & Spengler, 2005; Terjung et al., 2000).

Diffuse, multifocal strictures involving the medium-sized intrahepatic and/or medium or large-sized extrahepatic ducts demonstrated by cholangiography constitutes the gold standard in the diagnosis of PSC (MacCarty et al., 1983) (Pictures 1 and 2). ERCP may be used both for diagnostic and therapeutic purposes, but is associated with procedure-related complications such as cholangitis and pancreatitis (Moreno & Gores, 2006). Magnetic Resonance Cholangiopancreatography (MRCP) is a non-invasive alternative tool for PSC diagnosis. MRCP has a sensitivity of 80-91%, specificity of 85-99%, and diagnostic accuracy of 83-93% for diagnosis of PSC, all of which are slightly lower than those of ERCP (Fulcher et al., 2000; Angulo et al., 2000; Moff et al., 2006; Berstad et al., 2006; Textor et al., 2002).

Picture 1. Percutaneous cholangiogram displaying early PSC in a 45-year-old man with ulcerative colitis and elevated results of liver function tests. (Vitellas et al., 2000)

Liver biopsy is useful for the diagnosis of small-duct PSC or pericholangitis in IBD patients with cholestatic enzyme elevation and a normal cholangiogram (Burak et al., 2003). US examination of the liver may be used for screening in patients with liver enzyme elevation. The demonstration of dilated bile ducts by US may guide the clinician for further evaluations such as ERCP or liver biopsy. Nevertheless, there is no published data about the value of abdominal US for screening purposes in PSC (Navaneethan U, Shen B, 2010).

Another clinical scenario that may be encountered throughout the clinical course of PSC is cholangiocarcinoma (Picture 3). Current imaging methods have limited capability in the early diagnosis of cholangiocarcinoma and most patients are diagnosed at advanced stages of the disease. Computerized tomography (CT) and magnetic resonance imaging (MRI) have higher sensitivity than US, but US examination may provide better accuracy than CT and MRI in distinguishing cholangiocarcinoma from underlying PSC (Charatcharoenwitthaya et
al., 2008). When a mass lesion is detected by US, CT or MRI may be performed to assess the extent of the mass.

Picture 2. Percutaneous cholangiogram displaying PSC in a 67-year-old man with ulcerative colitis and jaundice. (Vitellas et al., 2000)

Picture 3. MRCP image of cholangiocarcinoma in a 29-year-old man with ulcerative colitis and primary sclerosing cholangitis who presented with jaundice and abdominal pain. (Vitellas et al., 2000)
ERCP with brush cytology or CT/US guided biopsy of the mass may be required to accurately establish the diagnosis. ERCP is especially useful for the final diagnosis of cholangiocarcinoma, as it provides opportunities such as brush cytology and obtaining a biopsy sample (Charatcharoenwitthaya et al., 2008). The application of advanced cytological techniques such as digital imaging analysis (DIA) and fluorescent in situ hybridization (FISH) in brush cytology and biopsy samples obtained by ERCP may further facilitate the early diagnosis of cholangiocarcinoma (Moreno & Gores, 2006; Rea et al., 2005; Parsi et al., 2008). The sensitivity of combined use of cytology, FISH and DIA is 50-64%, specificity and positive predictive value is 100% (Charatcharoenwitthaya et al., 2008). The National Institutes of Health consensus statement on the role of diagnostic and therapeutic ERCP described the role of ERCP in the diagnosis of biliary cancers in patients with PSC (Cohen et al., 2002). It is not clear whether a different surveillance program is needed in case PSC is accompanied by IBD. An algorithm for cholangiocarcinoma screening in patients with IBD and PSC is given below (Figure 3). The advantages and disadvantages of diagnostic modalities are summarized in Table 4.

![Diagram](Fig. 3. Surveillance of cholangiocarcinoma in patients with IBD-PSC (Adapted from Charatcharoenwitthaya P and Lindor KD, 2007))
Inflammatory Bowel Disease and Primary Sclerosing Cholangitis

| Diagnostic modality | Advantages | Disadvantages |
|---------------------|------------|---------------|
| Endoscopic retrograde cholangiopancreatography (ERCP) | - Can be both diagnostic and therapeutic Gold standard for diagnosis of PSC | - Complications, including post-ERCP pancreatitis. - Procedure associated cholangitis |
| Magnetic resonance retrograde cholangiopancreatography (MRCP) | - Visualizing bile ducts proximal to a complete bile duct obstruction - Useful in patients with gastric bypass or biliary-enteric anastomosis without radiation exposure | - Purely diagnostic - PSC limited to the peripheral ducts - Sensitivity and specificity slightly inferior to ERCP |
| Liver biopsy | - Diagnosis of small duct PSC in suspected patients with a normal cholangiography | - Patchy nature of PSC which leads to sampling error |
| Abdominal ultrasound | Screening for biliary ductal dilatation | Insensitive for diagnosis of PSC |

Table 4. Diagnostic modalities in primary sclerosing cholangitis (Adapted from Navaneethan & Shen 2010)

7. Natural history

Both symptomatic and asymptomatic PSC patients have decreased survival rates when compared to healthy population (Porayko et al., 1990). Median survival from time of diagnosis to death or liver transplantation was reported as 10 years (Farrant et al., 1991). In a Dutch cohort, transplantation free survival was 18 years on average (Ponsioen et al., 2002). PSC patients are prone to end stage liver disease complications such as esophageal varices, ascites, and hepatic encephalopathy. There are no guidelines on timing of screening for esophageal varices in PSC with or without IBD except for patients with cirrhosis. Because of the patchy nature of PSC, liver biopsy is limited by sampling error. Thus, non-invasive diagnostic methods were explored. A platelet count less than 150 000/mm³ may be a marker for the presence of esophageal varices (Zein et al., 2004). The composite Mayo Risk Score based on age, serum bilirubin, albumin, aspartate aminotransferase and variceal bleeding has been used to assess disease progression and prognosis (Kim et al., 2000). Cholangiocarcinoma is another complication of PSC and annual incidence is 0.6%-1% (Bergquist et al., 2007). It can develop at any stage of PSC, it can present as an intraductal tumor in the biliary system or rarely as a hepatic mass (Fevery et al., 2007). The risks of colorectal cancer and pancreatic cancer are also increased in patients with PSC when compared to the general population (Kim et al., 2000).

8. Management

The presence of accompanying IBD does not change the therapeutic approach in patients with PSC. Currently, there is no treatment option capable of halting progression to PSC or...
modifying the natural course of the disease. Ursodeoxycholic acid (UDCA) is widely used for the treatment of PSC. A randomized trial which tested a UDCA dose of 13-15 mg/kg/day showed normalization in liver enzyme levels, however, there was no improvement of liver histology or liver transplant free survival (Lindor, 1997). In another randomized trial that included 219 patients, a higher dosage of UDCA treatment (17-23 mg/kg/day) did not result in any benefit in terms of mortality, need for OLT, or cholangiocarcinoma risk (Olsson, 2005). A pilot study reported that a very high dosage of UDCA (28-30 mg/kg/day) might improve survival (Cullen et al., 2008), however, a large multicenter randomized controlled trial was terminated prematurely due to side effects in the treatment arm (Lindor et al., 2009).

UDCA treatment can prevent colonic neoplasia in PSC-IBD patients (Tung et al., 2001). On the other hand, high dose UDCA treatment was reported to increase the risk of colon cancer in PSC-UC patients (Eaton et al., 2011).

The treatment of choice in end stage PSC or cholangiocarcinoma-PSC is OLT. The 5 and 10 year survival rates in PSC patients are 85% and 70%, respectively (Rea et al., 2005 and Graziaidei et al., 1999a). On the other hand, the rate of PSC recurrence in the transplanted liver is 20%-25% (Graziaidei et al., 1999b).

**Small-duct PSC:** This entity was previously termed pericholangitis. These patients have biochemical and histopathological features compatible with PSC despite normal cholangiograms (Wee et al., 1985). The Mayo Clinic’s criteria for the diagnosis of small-duct PSC mandates the presence of coexisting IBD (Angulo et al., 2002). On the other hand, European criteria for small duct PSC does not include the presence of IBD (Björnsson et al., 2002 and, Broome et al., 2002). Initial studies with limited follow-up suggested that patients with small-duct PSC have a better prognosis than those with its large-duct counterpart (Angulo P et al., 2002; Björnsson et al., 2002 and, Broome et al., 2002). A large multicenter study with a longer follow-up period showed that the incidence of IBD is 80% in patients with small-duct PSC. Accompanying IBD was UC in 78%, Crohn’s colitis in 21%, and collagenous colitis in one patient (Björnsson et al., 2008).

The presence of IBD does not seem to have an effect on the course of liver disease in patients with small-duct PSC. 12-23% of patients with small-duct PSC progress to large-duct PSC. Cholangiocarcinoma developing in the setting of isolated small-duct PSC has never been reported in the literature, unless it progresses to large-duct PSC (Björnsson et al., 2008). In some patients, OLT may be necessary because of progressive disease and disease recurrence may be encountered in the transplanted liver. If the alkalene phosphatase level is elevated, cholangiogram is normal, and other hepatobiliary diseases are ruled out in an IBD patient, liver biopsy may be required to rule out small-duct PSC (Navaneethan & Shen, 2010).

9. **Disease course of IBD following liver transplantation for PSC**

OLT for PSC or PSC-associated cholangiocarcinoma may affect the clinical course of UC, as corticosteroids and other immunosuppressive agents used following OLT may theoretically improve coexisting UC. However, while some studies report alleviation of UC-associated symptoms after OLT (Saldeen et al., 1999 and Befeler et al., 1998), some studies report exacerbation of symptoms after OLT (Riley et al., 1997; Papatheodoridis et al., 1998 and Verdonk et al., 2006).

The effect of OLT on the natural course of ileal pouch in UC patients undergoing restorative proctocolectomy seems to be minor. In some case series, OLT and subsequent
immunosuppressive therapy did not have a significant effect on the severity of pouchitis (Zins BJ et al.,1995). In a recent study investigating the modification of chronic pouchitis course by PSC and/or OLT performed for PSC, 14 of 32 patients who underwent both IPAA and OLT for PSC experienced chronic pouchitis and in 8 patients PSC recurred, with 4 of them requiring retransplantation (Mathis et al., 2008). Large scale studies are needed to clarify the disease nature of pouchitis and the interaction of it with OLT and OLT related procedures.

10. Course of liver disease following IPAA

IPAA does not appear to affect the disease course of PSC and PSC appears to follow an independent disease course in spite of proctocolectomy (Poritz & Koltun, 2003). In a study of 214 patients with UC undergoing IPAA including 13 (6.1%) with PSC, 4 patients with PSC showed clinical progression, while none of these patients with minor histological changes progressed with a follow-up over 9 years (Mikkola et al., 1995). In a study of PSC in 68 patients with UC with 30 having follow-up examinations, the staging of PSC on liver biopsy after IPAA showed disease progression in 4 (13%), regression in 15 (50%), and stable in 11 (37%) from baseline liver histology at the time of proctocolectomy. Six of the 68 patients (8.8%) developed cholangiocarcinoma. The progression of PSC in patients with minor ductal changes appeared to be uncommon after IPAA surgery (Lepistö et al., 2009).

11. Colorectal Dysplasia and carcinoma in patients with Ulcerative Colitis and Primary Sclerosing Cholangitis

It is well-known that the risk of developing colorectal carcinoma is increased in patients with UC. Disease duration and extensive colitis are two major risk factors for this serious complication. The mechanism of developing malignancy in UC is not fully understood. Recent studies have shown that family history of sporadic colorectal carcinoma (Askling et al., 2001), presence of active inflammation in the mucosa (Rutter et al., 2004), and presence of PSC increase the risk of dysplasia and colorectal carcinoma in patients with UC (Kornfeld et al., 1997; Brentnall et al., 1996). A study from Sweden has shown that the incidence of colorectal dysplasia and carcinoma in patients with UC and PSC at 10, 20, and 25 years is 9%, 31% and 50% respectively (Broome et al., 1995). A meta-analysis of 11 studies has shown that the presence of PSC is an independent risk factor for colorectal dysplasia/carcinoma in patients with UC (Soetikno et al.,2002).

The incidence of colorectal dysplasia/cancer is high after orthotopic liver transplantation (OLT) in patients with UC and PSC (Bleday et al., 1993; Higashi et al., 1990). The risk of developing colorectal carcinoma increases over time after OLT. Vera et al. have identified three risk factors for developing colorectal carcinoma after OLT; colitis duration of more than 10 years, pancolitis, and dysplasia after OLT (Vera et al., 2003). A yearly colonoscopy is adequate in patients with PSC and UC, beginning from the time of PSC diagnosis (Broome & Bergquist, 2006).

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