Ulcerative Colitis with and without Primary Sclerosing Cholangitis: Two Different Diseases?

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\textbf{Abstract}

\textbf{Background:} Primary sclerosing cholangitis (PSC) is a chronic cholestatic liver disease of unknown origin and an important hepatobiliary complication of inflammatory bowel diseases (IBD), especially ulcerative colitis (UC). When further differentiated, about 3–8\% of UC patients suffer from PSC, whereas among Crohn’s disease patients the reported prevalence of PSC is probably between 1 and 3.5\%. Although it was reported from Japan that the frequency of PSC in UC was only 34\%, the same registry data indicated that the prevalence was up to 57\% among young patients with PSC even in Japan, which is comparable to the 60–80\% in Europe and the US. Additionally, the clinical features of UC in patients with PSC are different from those in patients without PSC, for instance, rectal sparing and right-sided dominance. \textbf{Summary:} The strong link between atypical IBD and PSC suggests that the pathogenesis of PSC involves pathology of the gut, including abnormal gut microbiota and aberrant activation of mucosal lymphocytes. These seem to be different in UC PSC as compared to the pathology of typical UC. \textbf{Key Messages:} The key to solving the question ‘Is there a difference between East and West?’ are genetic studies, genome-wide association studies of PSC in particular, which have already been performed in the West and are strongly warranted in the East.

\textbf{Background West}

In Europe, especially Northern Europe, about 70–80\% of patients with primary sclerosing cholangitis (PSC) are also diagnosed with inflammatory bowel disease (IBD). This is primarily ulcerative colitis (UC), less frequently Crohn’s disease (CD) [1]. Conversely, abnormalities of the hepatobiliary systems are found in up to 30\% of IBD patients [2].

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Of the liver diseases specifically associated with IBD, PSC is the most common, affecting up to 5% of all IBD patients depending on the geographic region [3]. When further differentiated, about 3–8% of UC patients suffer from PSC [4, 5], while in CD patients the reported prevalence of PSC is probably between 1 and 3.5% [6]. PSC has a clear male predominance with a ratio of 2:1.

Prior to the diagnosis of PSC in a patient with elevated serum markers of cholestasis (alkaline phosphatase and \( \gamma \)-glutamyltransferase), a possible secondary sclerosing cholangitis caused, for example, by infection, ischemia, IgG4-related cholangiopathy, or autoimmune hepatitis needs to be excluded. While PSC may present with symptoms like fever, night sweats, and pruritus, it is important to remember that IBD patients are frequently asymptomatic with regard to their liver disease, and physicians treating IBD patients need to actively investigate for possible associated hepatobiliary conditions [7]. The diagnostic gold standard for PSC remains magnetic resonance cholangiography (MRC), with typical 'string-of-pearls' imaging features of bile duct changes with multifocal strictures and segmental dilatations. MRC findings may be complemented by histology from liver biopsy if imaging is inconclusive. Histologically, PSC can be differentiated by several staging systems. The Ludwig staging system applies 4 stages to describe the histological features of PSC: cholangitis and portal inflammation (stage I), periportal inflammation or fibrosis (stage II), septal fibrosis, and/or bridging necrosis (stage III), and biliary cirrhosis (stage IV). The applicability and prognostic value of the histological scoring was just recently confirmed by de Vries et al. [8]. Patients that present with clinical, biochemical, and histological findings indicating PSC but show a normal cholangiogram in MRC are classified as having small-duct PSC [9]. Small-duct PSC seems to have differing HLA associations, depending on whether it occurs in the context of IBD or not. While small-duct PSC in IBD is associated with HLA-B*08 and thus seems to be an early or mild form of large-duct PSC, small-duct PSC without IBD displays a different association with HLA-DRB1*13:01 and may represent a different entity of inflammatory cholangitis. In general, small-duct PSC may have a better prognosis than large-duct PSC [10].

Currently there is no therapy for PSC available that has been shown to alter the course of the disease. Guidelines in Europe and the US differ regarding the use of ursodeoxycholic acid (UDCA), which has been shown to improve laboratory values and histology in PSC patients when given at doses not higher than 15 mg/kg/day [11, 12]. Unfortunately, UDCA has no proven benefit for transplant-free and overall survival. 24-norUrsodeoxycholic acid is a novel, modified derivative of UDCA currently in clinical trial after promising results in a PSC animal model [13].

Focusing on the characteristics of UC associated with PSC, it was noted that the course of the IBD is more often benign, with mild-to-quiescent clinical symptoms and possibly even normal endoscopic appearance that might result in underdiagnosis of UC in PSC patients [14]. Yet, pancolitis and rectal sparing are described more frequently, that is, in about 65 and 30%, respectively, of UC PSC patients, compared to 50 and 10% of UC patients without PSC. Backwash ileitis may occur slightly more often in UC PSC (17 vs. 12%), but the data are controversial [4, 15, 16]. Possibly due to the milder and sometimes subclinical course of the IBD despite pancolitis, the risk of colorectal cancer (CRC) seems to be substantially higher in UC PSC. Alternative theories implicate impaired hepatic excretion of bile acids with subsequent accumulation of potentially mutagenic secondary bile acids in the colon [17, 18]. The fact that CRC in UC PSC develops more frequently in the right hemicolon with the highest concentration of secondary bile acids could support this proposed pathogenetic mechanism, as could the potential protective effect of UDCA [16, 19]. CRC in UC PSC seems to develop earlier in the course of the disease, with a mean interval from IBD diagnosis to CRC of 12–20 years compared to 17–44 years in non-PSC IBD, but a possible lead time bias due to earlier screening in UC PSC is being debated.

The striking association of PSC with UC has sparked numerous investigations into the underlying pathogenetic mechanisms, giving rise to several theories that link the two diseases to each other. While both are considered an immune-mediated condition, PSC lacks features of classic autoimmune diseases, beginning with its male predominance and the lack of response to any immunosuppressive therapy. Moreover, no disease-specific autoantibody has been identified for PSC [20].

There is mounting evidence that genetic susceptibility plays an important role in the development of both diseases and may constitute a strong link between them. There is an up to 40-fold increased risk of PSC in 1st-degree relatives when compared to the general population, supporting the concept of genetic susceptibility. Moreover, strong associations with HLA genes have been described earlier. In a recent genome-wide asso-
Cells into the liver cause 'aberrant lymphocyte homing' of gut-restricted T cells, and UC with products from the intestinal microbiota may also cause subsequent portal inflammation. The hepatic overload of bacterial components into the portal circulation and the intestinal barrier in the setting of UC may facilitate a 'leakage' syndrome. The chronically inflamed and disturbed intestinal barrier of UC may contribute to the close relationship between PSC and UC. This is consistent with the much more frequent clinical association [21]. Importantly, the newly identified genetic risk factors account for no more than 10% of the estimated overall PSC susceptibility, again raising the important question of environmental factors for the pathogenesis.

In addition to the firmly established genetic association between PSC and IBD there are the long-standing hypotheses of 'leaky gut' and 'toxic bile', which may likewise contribute to the close relationship between PSC and UC. The chronically inflamed and disturbed intestinal barrier in the setting of UC may facilitate a 'leakage' of bacterial components into the portal circulation and subsequent portal inflammation. The hepatic overload with products from the intestinal microbiota may also cause 'aberrant lymphocyte homing' of gut-restricted T cells into the liver [22, 23]. This immunological mechanism of liver injury may be complemented by an altered bile composition in PSC, leading to a more toxic bile with a resulting local injury to bile duct epithelia.

**Key Messages West**

PSC and UC are strongly associated in Europe and the US. Recent GWAS show a substantial genetic overlap while confirming distinct disease entities. The clinical course of UC with PSC is often oligosymptomatic compared to UC without PSC. Nonetheless, patients with UC PSC exhibit an increased risk for CRC. There currently is no effective therapy to alter the outcome of PSC.

**Conclusions West**

PSC with UC likely represents a distinct disease variant that requires special clinical surveillance. Further research into the gut-liver axis is needed to better understand and potentially treat this disease complex.

*Background East*

PSC is a chronic cholestatic liver disease of unknown origin, characterized by the progressive destruction of bile ducts caused by diffuse inflammation and fibrosis, which eventually leads to liver cirrhosis [24]. PSC is believed to be a rare disease in Asia compared to Europe and North America. In fact, population-based epidemiological data on PSC are lacking in Asia; since 1996 there have been several small case series from India [25], Singapore [26], Korea [27], and Iran [28], each of them enrolling 10–20 patients with PSC. In the study from India [25], 18 patients were presented, and their clinical profiles including age (mean age 39.0 years) and male predominance were comparable to those in the West. IBD were found in 50% of the patients. The outcomes were poor; 4 patients died during the 20.1-month follow-up period. The study from Singapore [26] had enrolled 10 cases with PSC (mean age 50.9 years), confirming the male predominance; however, the outcome was better, with only 1 death during 6.6 years of follow-up. The only epidemiological data available in Asia are data from our group presenting the prevalence of PSC in Japan, as reported in 2014 [29]. According to this work and other previously reported data from the West, incidence and prevalence rates of PSC are summarized in table 1. Except in the study conducted in 1994, the prevalence of PSC was around 5–16 per 100,000 inhabitants in the West, compared to less than 1.0 in Japan. Although the reason for this difference is unclear, genetic factors clearly play a crucial role.

In Japan, we performed questionnaire-based, nationwide surveys for PSC in 2012 and 2015. According to this study [29], several characteristic features of Japanese PSC patients, which had been scarcely demonstrated before by epidemiological studies from other countries, were demonstrated. First, there were 2 peaks in age distribution at diagnosis, that is, in the 3rd and 7th decades of life (fig. 1). Most of the previous epidemiological studies from North America and Europe indicated a single age group for the highest risk of developing PSC [30–33], with the exception of one study in Canada, which suggested two age groups (18–35 and >65 years) as being at higher risk, similar to what has been found in Japanese studies [34]. Second, the prevalence of IBD among PSC patients appeared to be low compared to that in Europe and North America, as described below. The estimated median survival was 13.1 years, and 5-year survival of all patients was 74.5%. Multivariate analysis demonstrated that age (OR = 1.73, 1.09–2.75, p = 0.0206) and total bilirubin (OR = 2.32,
1.47–3.66, p = 0.0003) were independent risk factors for the outcome of PSC in Japan [35].

PSC is an important hepatobiliary complication of UC. According to the reports from the West, PSC is observed as a comorbidity in 2.9–7.6% of patients with UC [7]. In the East, case series from Iran [28] and South Korea [27] demonstrated that the frequency of PSC among UC cases was 4.3% (19/447) and 1.1% (21/1,849), respectively, suggesting a comparable prevalence of PSC among patients with UC. Regarding patients with PSC, the prevalence of UC was striking: up to 80% of PSC patients have concomitant IBD. However, there appears to be a considerable geographical variation. In the study from Singapore [26], IBD was found in 20% of patients with PSC, although only 10 cases were included. In our nationwide study in Japan, however, the prevalence of IBD as a comorbidity was low in Japanese PSC patients [29]. The presence of IBD was restricted to only 34% (68/197) of PSC patients. However, there are considerable differences in the prevalence of IBD between young and elderly patients. Their mean age was 48.1 years, and when comparing patients younger than 48.1 years with elderly patients, the prevalence of IBD was 12% in the elderly and 57% in the young PSC patients, the latter rate being comparable to the prevalence rates in Europe and the US (fig. 2). Taken together, the phenotype of PSC in the younger generation is similar to that in the

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**Table 1. Incidence and prevalence of PSC**

<table>
<thead>
<tr>
<th>First author [Ref.], year</th>
<th>Country</th>
<th>Study period</th>
<th>Method of case ascertainment</th>
<th>Cases, n</th>
<th>Incidencea (95% CI)</th>
<th>Prevalencea (95% CI)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Escorsell [36], 1994</td>
<td>Spain</td>
<td>1984–1988</td>
<td>Questionnaires</td>
<td>43</td>
<td>0.07</td>
<td>0.22</td>
</tr>
<tr>
<td>Berdal [37], 1998</td>
<td>Norway</td>
<td>1985–1994</td>
<td>ICD codes</td>
<td>12</td>
<td>0.7</td>
<td>5.6</td>
</tr>
<tr>
<td>Boberg [38], 1998</td>
<td>Norway</td>
<td>1986–1995</td>
<td>Prospective cohort</td>
<td>17</td>
<td>1.3 (0.8–2.1)</td>
<td>8.5 (2.8–14.2)</td>
</tr>
<tr>
<td>Bambha [39], 2003</td>
<td>USA</td>
<td>1976–2000</td>
<td>Medical record linkage system</td>
<td>22</td>
<td>0.9</td>
<td>13.6</td>
</tr>
<tr>
<td>Card [40], 2008</td>
<td>UK</td>
<td>1987–2002</td>
<td>General Practice Research Database</td>
<td>149</td>
<td>0.41 (0.34–0.48)</td>
<td>3.85 (3.04–4.80)</td>
</tr>
<tr>
<td>Kingham [41], 2004</td>
<td>UK</td>
<td>1984–2003</td>
<td>Prospective cohort</td>
<td>46</td>
<td>0.91</td>
<td>12.7</td>
</tr>
<tr>
<td>Kaplan [43], 2007</td>
<td>Canada</td>
<td>2000–2005</td>
<td>Retrospective cohort</td>
<td>49</td>
<td>0.92</td>
<td>N/A</td>
</tr>
<tr>
<td>Boonstra [44], 2013</td>
<td>The Netherlands</td>
<td>2000–2007</td>
<td>Retrospective cohort</td>
<td>519</td>
<td>0.5</td>
<td>6.0</td>
</tr>
<tr>
<td>Tanaka [45], 2008</td>
<td>Japan</td>
<td>2007</td>
<td>Questionnaires</td>
<td>415</td>
<td>N/A</td>
<td>0.95 (0.61–1.29)</td>
</tr>
</tbody>
</table>

N/A = Not available. a Per 100,000 inhabitants.

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Fig. 1. Age distribution at presentation of patients with PSC in Japan.

Fig. 2. Prevalence of IBD in patients with PSC in Japan. CF = Colonofiberscope.
West, while PSC in the elderly is a somewhat different disease and might not be ‘primary’ but ‘secondary’ sclerosing cholangitis.

It is also of note that, from Japanese experience, the clinical features of UC in patients with PSC are different from those in patients without PSC. It is known that several unique features were noted, such as rectal sparing and right-sided dominance, in UC patients with PSC. In addition, nonspecific or unclassified colitis is frequently found in patients with PSC. Interestingly, this atypical UC or IBD is also reported in a retrospective cohort study of IBD patients in Korea [27], as well as in North American and European countries. Thus, the atypical presentation of IBD in PSC patients appears to be a general phenomenon worldwide. Furthermore, the strong link between atypical IBD and PSC suggests that the pathology of PSC clearly involves pathology of the gut, including abnormal gut microbiota and aberrant activation of mucosal lymphocytes, which seems to be different from the pathology of typical IBD or UC.

Obviously, the key to solving the question ‘Is there a difference between East and West?’ are genetic studies, GWAS of PSC in particular. While GWAS in patients with PSC were performed in the West and they indicated several susceptibility loci contributing to the development of PSC overlapping with those for IBD and other autoimmune diseases, no GWAS have been performed in the East. On another cholestatic liver disease, primary biliary cholangitis, GWAS have been performed both in the East and the West, demonstrating the involvement of similar pathways in the pathogenesis of primary biliary cholangitis, even though the susceptibility genes themselves were different, but there seemed to be no difference in genetic background between East and West. In Japan, GWAS using both young and elderly PSC patients are needed, and results should be analyzed and compared between younger PSC (with high complication rates of IBD) and elderly PSC patients (with low rates of IBD). Clinical experience suggests that the susceptibility loci in young PSC patients are similar to those observed in the West and overlapping with those in IBD or autoimmune diseases as well, while the susceptibility loci in the elderly are quite different or even undetectable.

Key Messages East

Japanese epidemiological data indicate that the prevalence of UC in patients with PSC appears to be low. However, in younger patients with PSC the prevalence of UC is 50–60%, comparable to that in Europe and the US. It is reasonable to assume that in the younger generation, the phenotype of PSC is similar to that in the West. It is also of note that the clinical features of UC in patients with PSC are different from those in patients without PSC, for instance, rectal sparing and right-sided dominance and nonspecific or unclassified colitis.

Conclusions East

Genetic studies – GWAS for PSC in particular – are strongly warranted to solve the question whether there is a difference between PSC in the West and PSC in the East. Clinical experience suggests that the susceptibility loci in young PSC patients are similar to those observed in the West and overlapping with those in IBD or autoimmune diseases as well, while the susceptibility loci in the elderly are quite different or even undetectable.

Disclosure Statement

The authors have nothing to disclose.

References
