Are Dominant Strictures in Primary Sclerosing Cholangitis a Risk Factor for Cholangiocarcinoma?

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Abstract

Purpose of Review Cholangiocarcinoma is a devastating, unpredictable complication of large duct primary sclerosing cholangitis (PSC), which occurs in 5–15% of patients. The aim of this review is to discuss whether dominant strictures (DS) occurring in the larger bile ducts in PSC are a risk factor for the development of cholangiocarcinoma.

Recent Findings The development of DS is related to specific genetic polymorphisms affecting the innate immune system and the microbiome. In a recent study, the mean survival of PSC patients with DS was much worse (13.7 years) than for those without a DS (23 years). Survival difference was related to a 26% risk of cholangiocarcinoma, which developed only in those with DS. Half of the patients with cholangiocarcinoma presented within 4 months of the diagnosis of PSC. In another study, the risk of developing cholangiocarcinoma was directly related to the presence of underlying IBD, although this remains controversial. Efforts are being made towards surveying for cholangiocarcinoma including magnetic resonance imaging, endoscopic surveillance and serum tumour markers, but so far, an effective surveillance strategy has not been identified. DS should be treated endoscopically in the setting of symptoms, and there is limited evidence to suggest this may impact protectively on progression to cholangiocarcinoma.

Summary It is established that the presence of symptomatic DS occurring in the larger bile ducts in PSC can be the first presentation of cholangiocarcinoma. There is an increasing body of evidence that even when proven to be benign, dominant biliary strictures predispose to the future development of cholangiocarcinoma. Regular surveillance should be targeted at this selected high-risk group of PSC patients.

Keywords Primary sclerosing cholangitis · Cholangiocarcinoma · Dominant stricture · Inflammatory bowel disease · Cholangioscopy

Introduction

Primary sclerosing cholangitis (PSC) is a rare, progressive, cholestatic liver disease characterised by inflammation, stricturing and obliterative fibrosis of the biliary system, ultimately leading to biliary cirrhosis, portal hypertension and eventually hepatic failure in the majority of patients [1]. The clinical course of large duct PSC is highly variable and unpredictable [1, 2]. Whilst the median survival from presentation to death or liver transplantation in symptomatic patients is approximately 10 to 12 years, 75% of asymptomatic patients will survive 15 years or more. A recent Dutch study has shown an overall median survival of 22 years in an unselected population of PSC patients [3].

In the past, the majority of patients with PSC died of hepatic failure following deepening, cholestatic jaundice. However, over the last 20 years with the advent of successful transplantation, the majority (40–50%) of patients with large duct PSC die of malignancy, either hepatobiliary (cholangiocarcinoma, gall bladder cancer or hepatoma in cirrhotic...
patients) or colonic adenocarcinoma in PSC patients with associated inflammatory bowel disease (IBD) [4]. The reason for the high malignancy rate is probably explained by chronic inflammation in the biliary system and the colon, although whether PSC patients have a particular genetic susceptibility to develop cancer is unclear.

It is apparent that, despite the strong association with non-smoking, large duct PSC is a premalignant disease. In contrast, small duct PSC has not been associated with an increased risk of either biliary or colonic neoplasia and has a much better prognosis [5].

Hepatobiliary Cancer

As stated above, approximately 5 to 15% of patients with large duct PSC die from the development of bile duct carcinoma, which often follows a very aggressive course [6–8]. Approximately 0.5 to 1.0% of patients with large duct PSC will develop cholangiocarcinoma (CCA) or gall bladder cancer every year [8].

Unfortunately, at present, there are no factors that will predict which patients will develop this cancer. Tumour markers such as carcinogenic embryonic antigen (CEA) and carbohydrate antigen 19-9 (CA 19-9) have been investigated as potential serum markers of the development of bile duct cancer in PSC. Although some centres have found elevations in serum CA 19-9 a useful predictor, these results have not been confirmed in other units [8].

The median survival after the diagnosis of cholangiocarcinoma in PSC is only 9–12 months, although liver transplantation after chemotherapy is possible in selected patients with small tumours (<3.0 cm) and no evidence of metastatic disease [3, 9]. In this highly selected group, the survival is comparable with PSC without cholangiocarcinoma, viz. a 65–80% 5-year survival post transplantation [10, 11]. In PSC patients who are found to have unsuspected bile duct cancers at the time of liver transplantation, there is an approximately 35% 5-year survival [12]. Thus, there is clearly an unmet need to identify subgroups of PSC patients at particular high risk of developing cholangiocarcinoma, where heightened surveillance may be helpful in identifying cholangiocarcinoma at an early treatable stage.

Dominant Strictures in PSC

Patients with large duct PSC may develop progressive jaundice, worsening of their liver biochemistry, or symptoms of cholangitis, prompting investigation for a dominant stricture (DS). A DS in PSC is defined cholangiographically as a stricture less than 1.5-mm diameter in the common bile duct, or less than 1 mm in the left or right main hepatic ducts [13]. (See Fig. 1.)

Approximately 10–62% of patients with PSC will develop a DS at some stage in their disease [14••, 15]. The development of dominant bile strictures in PSC has been associated with

CD14 receptor signalling. CD14 is known to be a key mediator of the innate immune system, and its common genetic variant has been associated with the development of both alcoholic liver disease and non-alcoholic steatohepatitis [16, 17]. The variant CD14-260C>T polymorphism was associated with the development of DS and an increased risk of cholangitis. It appears from this data that the innate immune response may be important during biliary stricture formation [18].

The importance of genetic factors in the development of DS in PSC was further emphasised by a recent genome-wide association study which identified the FUT2 secretor status and genotype, the single-nucleotide polymorphism rs601338, as a potential genetic risk factor in PSC, which significantly influences biliary bacterial composition [19]. Presence of this genotype in PSC has been strongly associated with episodes of cholangitis, fungobilia and the incidence of dominant stenosis [20•].

The Effect of Dominant Stricture on Prognosis and the Risk of Cholangiocarcinoma

The presence of a DS has been associated with a worsened long-term prognosis. This was demonstrated in a 25-year study of 128 PSC patients from a tertiary referral centre in the UK [14••]. In this selected population, the mean survival of those with DS was much worse (13.7 years) than for those
without a DS (23 years). It is noteworthy that much of the survival difference was related to a 26% risk of cholangiocarcinoma, which developed only in those with DS. Half of the patients with cholangiocarcinoma presented within 4 months of the diagnosis of PSC, emphasising the importance of thorough evaluation of new DS.

These findings have been confirmed from a German study, namely that the presence of DS in PSC is associated with a worse prognosis and an increased risk of carcinomas both in the bile duct and colon [21]. Furthermore, this study suggested that the risk of developing cholangiocarcinoma (and colonic cancer) was directly related to the presence of underlying IBD [22]. In this study, a group of 171 patients with PSC were followed prospectively for as long as 20 years; 20 had DS at entry and a further 77 patients developed a dominant stenosis over the course of follow-up. These patients were treated endoscopically by repeated balloon dilatations, and five patients were temporarily stented. In patients with DS without IBD, no cholangiocarcinoma occurred and the actuarial survival free of transplantation was 77.8% at 18 years. In contrast, the 18-year survival was only 23% in the PSC with a DS and IBD, and six patients in this group developed a cholangiocarcinoma and two patients gall bladder over the 20-year follow-up. The presence of IBD had no impact on survival in those without a DS. In the DS group, bacteria in the bile had no effect on survival whereas Candida in the bile was associated with reduced survival.

The findings relating to the effect of IBD on the natural history of DS were not confirmed in a cohort of 241 Dutch PSC patients followed for a mean of 6 years [23]. The main difference was that 11 malignancies (seven colorectal cancer and four biliary malignancies) were observed in the group without DS (irrespective of IBD status). Overall, the cancer risk in their cohort appeared to be more dependant on the presence of a DS rather than the combination of concurrent IBD and a dominant biliary stricture.

Taken together, the limited data suggest that the presence of a DS, regardless of whether it occurs at the diagnosis of PSC or develops during the course of follow-up, represents an increased risk of both cholangiocarcinoma and reduced survival. A summary of these studies pertaining to the risk of DS leading to cholangiocarcinoma is presented in Table 1.

### Table 1  
Studies which address the risk of the presence of a dominant stricture on development of cholangiocarcinoma

| Study                           | Country     | Length of FU (mean) | Number of patients studied | Presence of DS, n (%) | Diagnosis of CCA during FU |
|---------------------------------|-------------|---------------------|---------------------------|-----------------------|---------------------------|
| Rudolph G. et al., 2009 [21]    | Germany     | 6.9 years           | 171                       | 97 (56.7)             | 6 (6.2)                   |
| Janse M. et al., 2012 [23]      | Netherlands | 6.2 years           | 241                       | 77 (31.9)             | 9 (11.7)                   |
| Chapman M. et al., 2009 [14]    | UK          | 9.8 years           | 128                       | 80 (62.5)             | 21 (26.3)                  |

Comments: Prospective study Association of IBD as well as DS with development of CCA
          Retrospective study No association with IBD Published in letter to editor style only
          Retrospective study 48% of CCA were within 4 months of PSC diagnosis

There are several other studies that examine risk of dominant stricture on overall survival free of liver transplantation, but few studies which specifically address the risk of developing cholangiocarcinoma

Abbreviations: CCA cholangiocarcinoma, DS dominant stricture, FU follow-up, IBD inflammatory bowel disease, PSC primary sclerosing cholangitis
Surveillance for Development of Cholangiocarcinoma in Dominant Strictures in PSC

Surveillance strategies in PSC patients for the detection of cholangiocarcinoma at an early, potentially treatable stage have failed to date. Serum tumour markers such as CEA and CA 19-9 have been investigated as potential serum markers of the development of bile duct cancer in PSC but have a low sensitivity and specificity. Unfortunately, by the time the serum CA 19-9 is significantly persistently elevated, the carcinoma is usually too advanced for curable treatment [25]. Moreover, approximately 7% of the population are unable to express CA 19-9 due to genetic variation in the FUT3 gene [26, 27].

Annual magnetic resonance cholangiography (MRCP) surveillance for cholangiocarcinoma in PSC is widely employed although the technique has not yet proven to be of benefit [8]. Advances in MRCP technology may offer greater sensitivity and specificity than currently obtained with 3D MRCP.

It has been suggested in a recent Finnish study that PSC patients with advanced bile duct disease, which included DS patients, should receive regular surveillance by ERCP and brush cytology every 6 months to 2 years [28]. PSC patients with suspicious changes on ploidy analysis with DNA flow cytometry were referred for liver transplantation to be performed at an early treatable stage. This aggressive approach has not been adopted by many units, as in practice the exclusion of possible cholangiocarcinoma in PSC patients with a DS is often difficult. Despite having a high specificity, brush cytology of biliary strictures performed at ERCP is often non-confirmatory, with unacceptably low levels of sensitivity obtained in most centres. A recent meta-analysis showed a pooled sensitivity of only 43%, making it a poor tool for diagnosis of cholangiocarcinoma [29]. In addition, one must bear in mind the risks posed in employing ERCP in the patient with PSC, not least including the risk of introducing bacteria and the possibility of invoking (recurrent) cholangitis [30].

Fluorescence in situ hybridisation (FISH) analysis has been advocated as increasing the sensitivity and specificity of cytological analysis. Chromosomal polysomy detected by FISH has been shown to identify patients with early cholangiocarcinoma or those patients with a high risk of developing cholangiocarcinoma. Up to 80% of biliary malignancies show abnormalities in chromosomal number (i.e. aneuploidy) and/or structure, which can be detected by FISH [31]. It has recently been demonstrated that patients with serial biliary polysomy have high risk of developing cholangiocarcinoma, although patients with reversion of polysomy (greater than 50% of the PSC patients studied) have a decreased risk [32].

Per-oral video cholangioscopy has been studied as a mode of potentially increasing the sensitivity of diagnosis of cholangiocarcinoma in the setting of DS in PSC. In a prospective study from the Mayo Clinic of 30 patients undergoing ERCP for DS, 4 patients ultimately had cholangiocarcinoma, but the use of cholangioscopy did not improve detection beyond routine ERCP [33]. Other small studies have postulated a potential minor improvement of cholangioscopy on detecting cholangiocarcinoma beyond ERCP [34, 35]. Annual cholangioscopy with biliary biopsies, analogous to yearly colonoscopy for the early detection of colonic carcinoma in PSC, is being studied prospectively in high-risk PSC patients with extrahepatic disease, particularly those patients with DS.

Conclusion

There is an unmet need to identify subgroups of PSC patients at particular high risk of developing cholangiocarcinoma, where heightened surveillance may be helpful in identifying cholangiocarcinoma at an early treatable stage. There is an increasing body of evidence that even when the stricture(s) is proven to be benign, dominant biliary strictures predispose to the future development of cholangiocarcinoma in patients with PSC. Additionally, PSC patients associated with IBD may be at particularly high risk. Currently, there is no proven beneficial form of surveillance strategy for cholangiocarcinoma, and studies are ongoing to address this issue.

Compliance with Ethical Standards

Conflict of Interest Roger W. Chapman and Kate D. Williamson each declare no potential conflicts of interest.

Human and Animal Rights and Informed Consent This article contains no studies with human or animal subjects performed by any of the authors.

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