Primary sclerosing cholangitis (PSC) is a chronic inflammatory, large duct cholangiopathy that results in fibrotic strictures and dilations of the intra- and extrahepatic bile ducts. Unlike the more common small bile duct disease, primary biliary cirrhosis (PBC), PSC occurs predominantly in men and is commonly associated with inflammatory bowel disease. While the pathogenesis of PSC has not been fully elucidated, emerging evidence supports roles for the innate and adaptive immune systems, and genome-wide analyses have identified several genetic associations. Using the best available evidence, the present review summarizes the current understanding of the diagnosis, pathogenesis and management of PSC. Despite its rarity, there is an urgent need for collaborative research efforts to advance therapeutic options for PSC beyond liver transplantation.

**Key Words:** Diagnosis; Liver transplantation; Pathogenesis; Primary sclerosing cholangitis

Management of primary sclerosing cholangitis: Conventions and controversies

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Primary sclerosing cholangitis (PSC) is a chronic inflammatory, large duct cholangiopathy that results in fibrotic strictures and dilations of the intra- and extrahepatic bile ducts. Unlike the more common small bile duct disease, primary biliary cirrhosis (PBC), PSC occurs predominantly in men and is commonly associated with inflammatory bowel disease (IBD), albeit with a subtle phenotypic difference (eg, rectal sparing, right-sided disease and backwash ileitis) from classic ulcerative colitis (UC) (1). Furthermore, it is clinically associated with cholangitis, biliary cirrhosis and progression to liver failure. An overarching fear for patients arises from an ominous relative increased risk of malignancy, both within the hepatobiliary tree and in the bowel, if associated with IBD.

The fundamental injury seen in PSC is not to hepatocytes, but rather to medium- and large-size bile ducts, with cholangiography demonstrating intra- and/or extrahepatic bile ducts with localized or multifocal strictures and intervening segments of normal or dilated ducts (2). Histologically, concentric periductal fibrosis (‘onion skinning’) is seen, with eventual progression to narrowing and obliteration of small bile ducts. The cholangiocytes demonstrate reactive features including expression of adhesion molecules, inflammatory and profibrogenetic cytokines, and receptors, as well as growth factors stimulating extracellular matrix production, accumulation and proliferation of periductal myofibroblasts. There is an increased prevalence of autoimmune disease in patients and their relatives and, while evidence to support genetic predisposition is emerging, non-genetic factors likely play a significant role in disease risk. The presence of a variety of secondary causes that can mimic the histological and radiological features of PSC (including autoimmune disease, ischemia, infection and toxins) suggests commonality in certain final pathways associated with biliary injury and that the primary insult may be heterogeneous (3).

The profound and deleterious fibro-obliterrative process is clearly present among the desired targets for new treatments. Presently, care is support- and transplantation is offered as a surgical (life-saving) therapy when the patient progresses to end stage (4,5). The choleretic effects of ursodeoxycholic acid (UDCA) are associated with an improvement in liver biochemistry, but does not translate in to a credible survival benefit. Population-based studies from the Northern hemisphere, such as that by Lendvist et al (6) and others (7,8) estimate the incidence and prevalence of PSC to be 1.2 per 100,000 and 16.2 per 100,000, respectively, corresponding to earlier estimates from the Midwestern United States. Little in the way of consensus on the overall pathogenesis of PSC exists, and it remains the chronic liver disease with the greatest unmet clinical need. The present review addresses common concerns for practicing clinicians.

**PATHOGENESIS**

Overall, there is strong emerging evidence to demonstrate roles for both the innate and adaptive immune systems in disease pathogenesis (9). It is hoped that a better understanding of etiology will furnish clinicians and scientists with a greater opportunity to develop effective treatments. Notably, the pathways identified in PSC are distinct from those identified for PBC – a small bile duct lymphocytic cholangitis. Additionally, PSC is much less ‘autoimmune’ than PBC, at least genetically speaking. It is expected that future research will further define risk pathways using novel genetic approaches (eg, the Immunochip Project, fine mapping studies, exome sequencing) and increased sample size (eg, The International PSC Collaboration). Some of the emerging pathogenic themes are highlighted below.

**Lymphocytes and PSC**

The lymphocytic infiltrate of PSC consists primarily of T cells with a predominance of CD4+ T cells with a T helper cell 1 phenotype (portal infiltrate) and CD8+ T cells (lobular infiltrate) (10). In addition to the αβ T cells infiltrating the liver, there are increased proportions of γδ T cells in PSC, and the percentage of γδ T cells in peripheral
blood of patients with PSC is greater than in healthy controls. The coexpression of the interleukin (IL)-2 receptor and CD4350 suggests they have an activated memory phenotype. Although the inflammatory infiltrate of patients with PSC is largely comprised of T cells, other cell types including natural killer (NK) cells, macrophages, B cells and biliary epithelial cells likely contribute to immunopathogenesis. The activation of the innate immune system as a primary event in PSC is also suggested, with disease potentially triggered by bacteria (or more likely their pathogen-associated molecular patterns) that enter the portal circulation through an inflamed permeable intestine.

Immunoglobulin G directed against biliary epithelial cells has been found in the sera of some patients with PSC, and the binding of such antibodies to biliary epithelium appears to induce production of pro-inflammatory cytokines and upregulation of toll-like receptors.

Animal models of the disease that support ‘infectious’ triggers (eg, Helicobacter pylori) have been speculated by some to be relevant, with added biological corollaries to the sclerosing cholangitis seen in immunosuppressed patients with Cryptosporidial infection. As well, the cholangiographically similar autoimmune pancreatitis/sclerosing cholangitis has been associated with the presence of antibodies against a peptide with homology to an amino acid sequence of the plasmogen-binding protein of H pylori. The presence of perinuclear antineutrophil cytoplasmic antibodies (p-ANCA) is often noted in PSC (as well as other autoimmune liver diseases). Recently, it has been demonstrated that p-ANCA in autoimmune liver disorders react with β-tubulin isotype 5 as autoantigen as well as with its bacterial precursor protein FtsZ, with the implication that the p-ANCA in PSC (as well as other autoimmune liver diseases). Recently, it has been demonstrated that p-ANCA in autoimmune liver disorders react with β-tubulin isotype 5 as autoantigen as well as with its bacterial precursor protein FtsZ, with the implication that the p-ANCA in autoimmune liver diseases probably reflects an abnormal immune response to intestinal microorganisms.

Mouse models of PSC remain limited (11), with the only promising model arising after targeted deletion of the murine biliary transporter MDR2. Secondary effects are recognized in this model, with disrupted tight junctions and basement membranes, bile acid leakage into portal tracts, induction of a portal inflammatory (CD11b, CD47) infiltrate and activation of proinflammatory (eg, tumour necrosis factor-alpha and IL-1β) and profibrogenic cytokines (transforming growth factor-β1). A sclerosing cholangiopathy is also encountered in adult patients with cystic fibrosis, raising the question for a role of cystic fibrosis transmembrane receptor (CFTR) variants in PSC (12,13). In a subset of PSC patients with IBD, an increased prevalence of CFTR abnormalities was observed, and experimental induction of colitis in CFTR knockout mice resulted in the development of bile duct injury (14). While the association between CFTR mutations and human PSC is unclear, in a recent genome scan, the G-protein-coupled bile acid receptor 5 (TGR5) has been identified as a potential candidate gene (15). This molecule is involved in bile acid-induced fluid secretion in biliary epithelial cells, and can be found colocalized with CFTR and the apical sodium-dependent bile salt uptake transporter, suggesting a functional coupling of TGR5 to bile acid uptake and chloride secretion (16).

Intriguingly, TGR5 has also been identified in cholangiocyte cilia, where it might conceivably couple biliary bile acid concentration and composition to ductular bile formation.

Genes and PSC

In 2010, the first genome-wide study in PSC (17) found, by far, the strongest genetic associations to be detected near human leukocyte antigen (HLA)-B at chromosome 6p21. Consistent with this first study, the latest PSC genome-wide association study (18) also found that the strongest associations with PSC were detected at single-nucleotide polymorphisms in the HLA complex at chromosome 6p21, peaking at rs3134792 in HLA-B (P=6.8×10−10). Inclusion of rs3134792 as a covariate demonstrated a complex residual association signal in the vicinity of the HLA class II region, suggesting the presence of multiple causative loci within the region. Previous studies of PSC investigated markers of innate immunity in close proximity to the HLA locus. A key set of NK receptors is the killer-cell immunoglobulin-like receptor (KIR), which bind HLA class I molecules, and the frequency of HLA-Bo4 and HLA-C2, ligands for the inhibitory KIRs 3DL1 and 2DL1, respectively, was significantly reduced in PSC patients compared with controls, suggesting an increase in NK cell activity by decreased inhibition (19). In addition to the decreased suppression of NK cell activity through HLA-KIR interactions, genetic evidence suggests a possible role for the direct activation of NK cells through the major histocompatibility complex class I chain-related (MIC) genes. MICa and MICb are encoded in the HLA region and their proteins activate NKG2D receptors on NK cells. A strong protective effect of the MICA×02 allele has been demonstrated, as has an increased allele frequency of MICA×08 (20). In contrast, a Norwegian study (19) found associations with the MICa51 and MICb24 alleles.

These two pivotal PSC genome-wide association scans have further identified five PSC risk loci outside the HLA complex, including 2q13, 2q35, 3p21, 10p15 and 13q31. Of these loci, 2q35 and 3p21 are also associated with UC. The causal genes for these regions still need to be definitively confirmed, but the locus on chromosome 2q35 harbours a compelling candidate gene (the bile acid receptor TGR5) as discussed above. BCL2L11 is an interesting candidate for the association at 2q13 because of its role in maintaining immunological tolerance. The BCL2L11 gene encodes the Bcl-2 interacting protein (Bim), which is crucial for maintaining immunological tolerance through induction of apoptosis of autoreactive T-lymphocytes, as well as the deletion of activated T cells after an immune response. Bcl2L11−/− mice spontaneously demonstrate ADEPT infiltrates around intrahepatic bile ducts. IL2RA is of interest in PSC (10p15) because IL2ra−/− mice spontaneously develop intestinal and biliary inflammation. Significant association has also been confirmed for macrophage stimulating protein 1 (MST1) involved in regulation of the production of proinflammatory mediators. Interestingly, a PSC-associated amino acid change has been proposed to influence MST1 receptor interaction, as well as risk of UC and Crohn’s disease. The MST1 locus (3p21) likely represents an important overlapping susceptibility locus for PSC and IBD.

Adhesion molecules and PSC

A specific role for chemokines and their receptors has been highlighted for PSC, a disease often associated with colonic IBD (21-23). In this paradigm, long-lived memory lymphocytes arise as a result of colitis, and express homing receptors that direct subsequent migration not only to the gut but also the liver. Under normal circumstances, gut endothelium expresses a unique adhesion molecule known as mucosal addressin cell adhesion molecule-1 (MADCAM1), which is absent from other vascular beds, and a unique chemokine, CCL25, which is restricted to the small bowel. In PSC, hepatic inflammation leads to the upregulation of hepatic MADCAM1 and CCL25, and increased recruitment of mucosal T cells. Memory cells recirculate between the liver and gut but have the potential to become deleteriously activated by hepatic antigens. When characterized, one in five T cells infiltrating the liver in PSC can be shown to be interferon-γ-producing αβ7+CCR9+ memory/effector T cells, which are present at low frequencies in other liver disease. Such imprinting is dependent on gut dendritic cells because αβ7 and CCR9 are only colocalized on lymphocytes activated in the gut. Recent evidence indicates that gut-associated dendritic cells induce gut-homing receptors on B cells via a mechanism that depends on the vitamin A metabolite retinoic acid. Furthermore, the experimental inability of liver dendritic cells to similarly imprint gut tropism implies that the αβ7+CCR9+ T cell infiltrate in PSC is primed in the gut. The functional relevance of αβ7 and CCR9 expression is supported by observations that both MADCAM1 and CCL25, which are absent from normal liver, are present on hepatic endothelium in liver diseases associated with colitis, and that αβ7+CCR9+ lymphocytes from the livers of patients with PSC bind MADCAM1 and respond to CCL25 in adhesion and migration assays. A murine model has now also provided in vivo evidence that the enterohepatic
circulation of antigen-specific CD8 T cells can generate an auto-
nimmune cholangitis.

CLINICAL MANIFESTATIONS AND GENERAL
DIAGNOSTIC CONSIDERATIONS
PSC is a heterogeneous disease that manifests in a wide variety of
ways across all age groups. While the classic patient is male with IBD
presenting with symptoms in his third to fourth decade of life,
increasingly, with ready access to liver biochemical screening and
magnetic resonance imaging (MRI), patient presentation is spanning
a wider spectrum. Asymptomatic early disease is more frequently
identified while older patients are being seen with chronic disease
into their seventh decade of life and beyond.

Clinical presentations range from an asymptomatic rise in alkali-
line phosphatase levels, to nonspecific fatigue and pruritus, to liver
decompensation with profound jaundice and advanced features of
portal hypertension. With early MRI, incidental diagnosis occurs,
and patients with PSC need not have abnormal liver biochemistry.
Additional complications associated with PSC include cholelithia-
sis, gallbladder carcinoma and colorectal cancer (CRC).

PSC is usually diagnosed on the basis of a chronic cholestatic
derangement in liver enzyme levels (ie, alkaline phosphatase ≥1.5 times
upper limit of normal for at least six months) and fibrotic strictures of
the intra- and/or extrahepatic biliary system. In practical terms, cholangi-
ography by MRI is now, in essence, the primary diagnostic modality, with
blood work and ultrasound used to exclude secondary etiologies that can
mimic PSC. If cholangiography is normal, liver biopsy is indicated if no
alternative explanation for a persistent cholestasis can be unearthed. A
liver biopsy is not required to establish the diagnosis, but is of use when
considering small-duct PSC, or an overlap with autoimmune hepatitis
(AIH). Other laboratory abnormalities associated with PSC but non-
essential for its diagnosis include p-ANCA, hypergammaglobulinemia
and nonspecific antinuclear and smooth muscle autoreactivities.

Cardinal to securing the diagnosis of ‘primary’ sclerosing cholangitis
is the exclusion of secondary causes of cholangitis including, but not
limited to, biliary calculi, cholangiocarcinoma (CCA), biliary tract sur-
gery, Caroli’s disease, biliary toxin exposure, portal vein thrombosis,
ischemic stricturing and alternative liver diseases that can cause biliary
injury (eg, cholestatic drug-induced liver injury). It is particularly import-
ant for the clinician to also consider immunoglobulin G2-associated
sclerosing cholangitis in the differential diagnosis of any patient pre-
senting with biliary strictures given the characteristic high rate of steroid
responsiveness of this entity, thereby averting long-term complications
of strictureing biliary disease (24).

Diagnostic imaging, invasive cholangiography and laboratory
testing for CCA
All patients require hepatobiliary imaging to establish the diagnosis,
characterize the severity and/or complications of the disease and help
exclude other conditions in the differential diagnosis. Ultrasound
remains the first-line test of choice in the initial evaluation of patients
with PSC due to its broad availability, low cost, excellent safety profile
and high sensitivity for the detection of extrahepatic dominant stric-
tures. Ultrasound is important to evaluate the patency of the portal
vein and can also identify very early biliary wall thickening that may
not be evident on MRI. This can help diagnostically in early stages of
the disease (25). MRI with magnetic resonance cholangiopancreatog-
raphy (MRCP) is, as previously stated, also commonly used in the
assessment of PSC, but unlike ultrasound, MRI more reliably detects
the presence of intrahepatic disease. Despite the superior safety of
MRCP over endoscopic retrograde cholangiopancreatography (ERCP),
MRCP has slightly inferior accuracy, in large part due to high interob-
server variability (26). Practically, however, MRCP suffices as the
diagnostic modality of choice, with there being little to be gained
clinically for patients from the added risk of a diagnostic ERCP.
Computed tomography cholangiography is an emerging but not widely
available noninvasive modality for imaging the biliary tree. Preliminary

Management of primary sclerosing cholangitis

| Imaging modality | Advantages | Disadvantages |
|------------------|------------|---------------|
| Ultrasound       | Widely available and inexpensive; no radiation; good visualization of intrahepatic biliary tree | Poor sensitivity for intrahepatic strictures; poor specificity for malignancy |
| Computed tomography | Widely available; specificity for staging malignancy superior to ultrasound | Radiation; computed tomography, cholangiography protocols are not widely used |
| Magnetic resonance cholangiography | Good visualization of intrahepatic and extrahepatic biliary system; no ionizing radiation; no risk of cholangitis/pancreatitis | High interobserver variability in reporting; poor precision for ruling out malignancy; inability to be therapeutic |

TABLE 1 Summary of the advantages and disadvantages of noninvasive imaging modalities in primary sclerosing cholangitis

Unfortunately, tumour markers, such as carbohydrate antigen (CA) 19-9, have suboptimal sensitivity and positive predictive value for the diagnosis of CCA (28). This remains a great challenge for patients and physicians, particularly given the need for high-quality ERCP studies. In the setting of a dominant stricture, it is critical to quickly identify the presence of malignancy because a timely diagnosis of early-stage CCA improves the likelihood of potentially life-saving surgical treat-
ment. Regrettably, the radiological appearance of benign dominant strictures can appear identical to CCA. In such scenarios, tumour markers (ie, CA 19-9) and cytology should be pursued.

Cholangiography is the least invasive and most common means to obtain cytology. It is usually performed endoscopically or percuta-
nearily; the former approach is preferred due to its superior safety profile. The associated risks of ERCP and percutaneous transhepatic cholangiography (PTC) include cholangitis, bleeding, pancreatitis and bowel perforation (29). Due to the potential for complications from ERCP, patients often undergo MRCP in lieu of, or before, invasive cholangiography despite the inferior diagnostic yield of MRCP. However, given that MRCP has a low sensitivity to diagnose CCA, clinicians should preferentially refer patients for ERCP if there is an absence of MRI expertise at their centre, and/or if there is imaging evidence or clinical suspicion for distal dominant stricturing. ERCP or PTC, rather than MRCP, is also more helpful to determine the feasibil-
ity of surgical resection, and is more logical if the patient is jaun-
diced in the absence of end-stage liver disease, given the chance for
intervention.

Due to the risk of bacteremia with ERCP available evidence supports the routine use of prophylactic antibiotics with Gram-negative coverage (30). There are no prospective data to determine whether dilation alone – versus a combination of dilation and stenting – is superior. Patients with dominant strictures amenable to ERCP fre-
quently require several sessions to relieve biliary obstruction, but the treatment has a high success rate in improving symptoms and bio-
chemistry in the majority of patients. PTC has similar efficacy to
**TABLE 2**  
Summary of evidence on medical treatment options for primary sclerosing cholangitis other than ursodeoxycholic acid

| Category                  | Definitive efficacy | Level of evidence* | Reference(s) |
|---------------------------|---------------------|--------------------|--------------|
| Antibiotics               |                     |                    |              |
| Metronidazole             | No                  | 1b                 | 71           |
| Minocycline               | No                  | 1b                 | 72           |
| Immuno-modulatory         |                     |                    |              |
| Corticosteroids           | No                  | 1b                 | 73-75        |
| Methotrexate              | No                  | 1b                 | 76-78        |
| Pencillamine              | No                  | 1b                 | 79           |
| Azathioprine              | No                  | 2c                 | 80           |
| Mycophenolate mofetil     | No                  | 2b                 | 81           |
| Colchicine                | No                  | 1b, 3b             | 75, 82       |
| Etoricoxib                | No                  | 2b                 | 83           |
| Infliximab                | No                  | 1b                 | 84           |
| Cyclosporine              | No                  | 1b                 | 85           |
| Tacrolimus                | No                  | 2b                 | 86           |

*Adapted from the Centre for Evidence Based Medicine website [www.cebm.net](http://www.cebm.net)

ERCP for distal stricture, but is not preferred due to the higher rate of adverse events and the discomfort experienced afterward with external catheters. Furthermore, PTC is technically difficult in the presence of intrahepatic disease.

Endoscopic ultrasound (EUS) is another minimally invasive, non-surgical technique with outstanding ability to assess distal bile duct lesions and regional lymph nodes. EUS with final-needle aspiration has greater sensitivity and specificity for the diagnosis of distal CCA compared with ERCP and brush cytology. EUS has the added advantage of not contaminating the biliary tree (31).

Emerging invasive, nonsurgical technologies, such as cholangioscopy and intra-ductal ultrasound, hold promise but are not widely available, and more research is needed to assess their role in the arsenal of tests available for evaluation of the biliary system. In many patients with suspected biliary lesions, a staging laparotomy to ultimately determine resectability is required unless noninvasive imaging clearly shows distant metastases.

Because the sensitivity of cytology alone for the diagnosis of CCA is only 35% to 70% (32,33), combining cytology with molecular markers and novel genetic techniques is useful to improve the accuracy of diagnosis of CCA. The challenge, however, remains that these molecular techniques are not widely available and the practicalities of translating such ancillary testing to non-tertiary centres is very unclear. It should also be noted that the presence of high-grade dysplasia alone on brush cytology has high sensitivity, specificity and positive predictive value for the diagnosis of CCA (73%, 95% and 85%, respectively) (34).

Although CA 19-9 levels are frequently elevated in CCA, they are, unfortunately, also elevated in benign pancreaticobiliary conditions (particularly when patients are jaundiced) and other cancers such as adenocarcinoma of the pancreas. This severely limits the utility of measuring CA 19-9 and raises diagnostic problems when values are high but nothing is evident radiologically. The threshold value above which CA 19-9 is a useful diagnostic tool to rule in CCA depends on the clinical scenario in which it is measured. The sensitivity of CA 19-9 above 180 U/mL is only 67%, but the specificity is better at 98% (35).

In patients with cholangitis, a higher cut-off value of CA 19-9 >300 U/mL to 400 U/mL is still associated with a poor sensitivity (41%), but specificity remains high. While measuring CA 19-9 in patients with suspected CCA is reasonable, elevation of tumour marker levels alone is insufficient to establish the diagnosis. However, combining brush cytology with CA 19-9 increases the sensitivity to 88% (35).

Fluorescence in situ hybridization (FISH) and digital image analysis (DIA) are novel techniques to measure DNA proliferation and improve the diagnostic yield of cytology. FISH is a cytological method that uses DNA probes to identify a shortage or excess of chromosomes or chromosomal loci; these molecular derangements are common in CCA. Similarly, DIA quantifies the amount of cellular DNA by measuring intensity of nuclei stained with dye that binds to nuclear DNA. It is debatable as to what extent, if any, that FISH and DIA, in combination with cytology, improve the ability to diagnose CCA. The sensitivity and specificity for DNA aneuploidy in tumour detection is 43% and 96%, respectively (36).

Given that both PSC and CCA are rare, patients with PSC and suspected CCA are likely best served at designated centres of excellence where clinical expertise and novel diagnostic testing are readily available. Practitioners should consider referring PSC patients with malignant appearing stricture(s) with moderate to severe atypia on brush cytology, with or without CA 19-9 above 100 U/L, to a hepatobiliary centre of excellence.

**MEDICAL MANAGEMENT CONSIDERATIONS**

Other than liver transplantation (LT), there are no proven effective treatments for PSC. An array of antimicrobials and immunomodulatory agents have been studied in the past two decades, with no robust demonstration of efficacy (Table 2).

**UDCA**

The role of UDCA in PSC is controversial, and requires further study before evidence-based recommendations can be finalized. As a therapeutic agent, UDCA is purported to have four mechanisms of action:

1. Protection of cholangiocytes against cytotoxic bile acids
2. Promotion of hepatobiliary secretion
3. Prevention of bile acid-induced apoptosis of hepatocytes; and
4. Induction of antioxidants.

Several PSC trials involving UDCA have repeatedly demonstrated an improvement in liver biochemistry, and smaller studies have even shown a tendency toward improved liver histology and cholangiogram appearances of strictures (37,38). Furthermore, in a multicentre trial, Olsson et al (38) reported a tendency toward improved transplant-free survival in patients randomly assigned to 17 mg/kg/day to 23 mg/kg/day of UDCA versus placebo, but statistical significance was not achieved. Additionally, Pardi et al (39) provided evidence for a significant reduction in the risk of colonic dysplasia or CRC in patients with PSC and UC who received UDCA at 17 mg/kg/day to 23 mg/kg/day. The results of this latter study further propelled the routine use of UDCA for patients with PSC and IBD, despite the fact that this trial involving only 52 subjects was never replicated. It should be noted that high-dose UDCA (28 mg/kg/day to 30 mg/kg/day) has been recently shown to increase the risk of colorectal neoplasia (HR 4.44; P=0.02) for unclear mechanisms (34).

In a randomized placebo-controlled trial, Lindor et al (40) showed that high-dose (28 mg/kg/day to 30 mg/kg/day) UDCA is not associated with improved transplant-free survival and, in fact, UDCA at this dosage may be harmful because subjects in the treatment arm of the study experienced higher frequencies of hepatic decompensation (40).

Although the precise mechanism(s) of adverse events in patients with high-dose UDCA is unknown, it is possible that UDCA has a previously unrecognized therapeutic window (ie, at a very high dose, toxic levels of lithocholic acid may become relevant), and beyond a therapeutic dosage, UDCA may lead to prevention of apoptosis of stellate cells, which in turn could lead to accelerated fibrosis.

Based on the above results, high-dose UDCA at 28 mg/kg/day to 30 mg/kg/day has no role in the management of PSC, but UDCA at 17 mg/kg/day to 23 mg/kg/day may be therapeutic in selected patients pending further trials. It is conceivable that UDCA could modify disease during the early, inflammatory phase of biliary injury before fibrosis has ensued. This has left patients and clinicians confused. While it is clear that UDCA is not a panacea, and that high-dose UDCA is neither effective nor necessarily safe, the role for moderate...
dose (17 mg/kg/day to 23 mg/kg/day) UDCA is unclear. Multisociety guidelines are unable to concur on practical advice. As such, most clinicians now engage the patient in discussion and one practical strategy is to trial UDCA at 17 mg/kg/day to 23 mg/kg/day, and continue it if there is a biochemical response. Although this ‘art of practice’ does not conform to ‘evidence-based’ medicine per se, it does resonate pragmatically with clinicians and patients, and there is no evidence for harm, other than arguably in monetary terms.

Other aspects of PSC management are less controversial and broadly involve either treatment of known complications of the disease or prevention of theoretical complications (Table 3). Modifiable complications include bacterial cholangitis, pruritus, malnutrition and hepatic osteodystrophy. Prevention of CRC and the role of LT will also be discussed.

**Bacterial cholangitis**

Cholangitis is a major cause of morbidity in the PSC population, but it should be borne in mind that it is unusual in early disease and that it is often precipitated by biliary intervention. Treatment of significant bacterial cholangitis begins with patient resuscitation and blood cultures, timely antimicrobial therapy targeting Gram-negative organisms, and subsequent attempt at relieving biliary obstruction with endoscopic or percutaneous cholangiography, if appropriate. However, some patients may experience mild episodes that they are able to manage at home with self-administration of oral ciprofloxacin. It is also appropriate to bear in mind that the classic presentation of pain, fever and leukocytosis need not be present in a patient with PSC, especially the more advanced and extensive their disease. Rarely, even biliary resection may need to be considered for stricturing disease not suitable for therapeutic cholangiography in the patient without advanced hepatic fibrosis (41). In the process of relieving stricturing disease, brushings and/or biopsies should be considered because CCA often presents as an innocent episode of cholangitis. Patients with recurrent cholangitis may benefit from prophylactic antibiotics to prevent septicaemia, with the caveat that usually such patients ultimately progress toward LT.

**Pruritus**

Pruritus is a debilitating symptom that markedly diminishes a patient’s quality of life. In its most classic form, pruritus is characterized by plantopalmar itch, sometimes relentless, and often worse at night. Concomitant depression and anxiety in the most severe of cases is inevitable. In addition to relief of biliary obstruction where appropriate, cholestyramine remains a reasonable first-line agent to prescribe. Additional agents that can be of benefit should be tried sequentially given the absence of an evidence-based algorithmic approach, with discontinuation of noneffective agents. Examples of antipruritic agents for which there is some evidence in cholestatic liver diseases include rifampin, sertraline, hydroxyzine and oral naltrexone. It is also appropriate to bear in mind that the classic presentation of pain, fever and leukocytosis need not be present in a patient with PSC, especially the more advanced and extensive their disease. Rarely, even biliary resection may need to be considered for stricturing disease not suitable for therapeutic cholangiography in the patient without advanced hepatic fibrosis (41). In the process of relieving stricturing disease, brushings and/or biopsies should be considered because CCA often presents as an innocent episode of cholangitis. Patients with recurrent cholangitis may benefit from prophylactic antibiotics to prevent septicaemia, with the caveat that usually such patients ultimately progress toward LT.

**Hepatic osteodystrophy**

Hepatic osteodystrophy is a prevalent complication of end-stage liver disease (ESLD), and particularly of chronic cholestatic liver diseases. Approximately 15% of patients with PSC have osteoporosis at the time of diagnosis, necessitating baseline and serial (ie, every two or three years) bone mineral density testing (44). In addition to appropriate supplementation with calcium and vitamin D, patients with osteoporosis should receive bisphosphonate therapy to prevent further bone loss and fractures as per multisociety osteoporosis guidelines (4).

**Management of primary sclerosing cholangitis**

| Complication | Management strategy considerations |
|--------------|-----------------------------------|
| Cholangitis  | Antibiotics, therapeutic cholangiography with dilation or stent insertion |
| Pruritus     | Antipruritic agents (see text). Relief of biliary obstruction with therapeutic cholangiography |
| Hepatic osteodystrophy | Calcium and vitamin D supplementation, bone density screening, bisphosphonate therapy for osteoporosis |
| Malnutrition | Protein and calorie supplementation. Screening and treatment of fat soluble vitamin deficiencies |
| Colorectal cancer | Colonoscopy with segmental biopsies for surveillance in those with inflammatory bowel disease |
| Atypical ducts | Review with expert pathologist, specimens with moderate to severe atypia require careful follow-up |

Intravenous bisphosphonates may be an attractive option in subjects with esophageal varices in whom there is theoretical added concern over the risk of pill-induced upper gastrointestinal tract ulceration.

**Malnutrition**

Regardless of etiology, patients with ESLD have a propensity for sarcopenia due to their profound catabolic state. Patients with clinical features of malnutrition benefit from dietary assessment, and protein and calorie supplementation. In addition to protein and calorie malnutrition, patients with advanced cholestatic liver diseases are vulnerable to fat-soluble vitamin deficiencies. Once patients with PSC develop visible jaundice (total bilirubin level >35 mmol/L), they should receive annual surveillance and aggressive replacement of fat-soluble vitamins where indicated. Appropriate replacement of fat-soluble vitamins is crucial to prevent or treat complications such as osteopenia and osteoporosis, myalgia, night blindness, wound healing and muscle cramps, among others.

**Portal hypertension**

Patients with advanced PSC are prone to the garden-variety complications of portal hypertension, including variceal hemorrhage and ascites. Similar to the generic management of ESLD, patients with cirrhotic-stage PSC require routine endoscopy to survey for large varices. A platelet count of <150×10^9/L in a patient with PSC is associated with an OR of 6.3 (95% CI 2.6 to 15.8) to predict large varices (45). Because large varices can occur in the setting of normal range platelet counts in subjects with cholestatic liver diseases (46), any patient with features of portal hypertension, even in the absence of thrombocytopenia, should be considered for endoscopy to avert variceal hemorrhage through banding ligation or nonselective beta blockade.

**Surveillance for CRC**

As previously discussed, PSC is associated with both UC and Crohn’s disease, but the former is a far stronger association. The PSC/UC overlap is considered a unique phenotype of IBD and, as previously stated, is associated with rectal sparing and backwash ileitis (47). In patients with PSC/UC, the RR of colorectal cancer is 10-fold greater than in the general population and, hence, annual surveillance colonoscopy with segmental biopsies to assess for dysplasias is advised (48,49). A meta-analysis of observational studies suggested that 5-aminosalicylic acid agents may be protective against colonic dysplasia and CRC (OR 0.51 [95% CI 0.38 to 0.69]) (50), and should thus be considered in PSC patients with UC. In the setting of colonic dysplasia or severe uncontrolled UC, a colectomy is warranted, although patients with advanced liver dysfunction and portal hypertension (Child-Pugh Class
B or C disease) should be assessed for concomitant LT due to their high risk of hepatic failure with intra-abdominal surgery.

Managing variant syndromes

The diagnosis of variant syndromes associated with PSC, including overlap syndromes, small-duct PSC and sequential syndromes, is problematic due to the lack of standardized diagnostic criteria (51). While an approach to the diagnosis of variant syndromes is beyond the scope of the present review, the clinician should be mindful of a possible overlap with AIH in a PSC patient with a prominent hepatic derangement in their liver enzyme profile not otherwise accounted for, and corroborating histopathological features of AIH. Published literature suggests that 5% to 30% of patients with PSC have concomitant AIH (52-54). Patients with PSC-AIH overlap frequently benefit from immunosuppression to prevent disease progression and liver failure. An overlap with immune-mediated hepatitis is much more common in children (55,56). This condition has been called ‘autoimmune sclerosing cholangitis’ by pediatricians. It describes the situation by which it was identified: approximately 50% of children with classic AIH have cholangiographic features of PSC. Pediatricians are, therefore, likely to manage patients with immunosuppression and UDCA in this setting, although a robust evidence base for this is lacking. If clinicians see patients early in disease who are young, they should be more mindful to the possibility of a steroid responsive component of the disease but must be prudent before making the label.

Another consideration is that patients with PSC may also have a specific variant of the disease, small-duct PSC, diagnosed on the basis of a compatible clinical context and liver biopsy, and normal cholangiogram. Natural history studies suggest that small-duct PSC is merely an early stage of large-duct PSC, and the incidence of this entity is increasing over time (8,57). Patients with small-duct PSC are usually asymptomatic and have minimal foreseeable risk of dire complications such as CCA (57,58).

LT

The only proven curative treatment for PSC is LT. The decision for LT rests on several general considerations: quality of life of the potential recipient; longevity of life of the potential recipient; and organ availability and societal costs. Patients with advanced PSC have a particularly dismal quality of life due to the unique symptoms that afflict them, including pruritus and disproportionate fatigue; such symptoms are not directly captured by scoring systems used to evaluate LT candidates.

In most transplant programs in North America, qualified LT candidates are listed on the basis of the Model for End-stage Liver Disease (MELD) score. The MELD score, based on international normalized ratio, bilirubin and creatinine levels, provides a three-month mortality risk for potential recipients. Patients with the highest MELD score are given priority for LT in an effort to reduce the mortality rate among dates are listed on the basis of the Model for End-stage Liver Disease (MELD) score. The MELD score, based on international normalized ratio, bilirubin and creatinine levels, provides a three-month mortality risk for potential recipients. Patients with the highest MELD score are given priority for LT in an effort to reduce the mortality rate among LT recipients with a history of PSC should be aware that the recurrence rate of PSC after LT is approximately 20% after 10 years (66). To establish the diagnosis of recurrent PSC, causes of nonanastomotic biliary strictures must be excluded, including but not limited to, ischemic injury of the hepatic graft, graft derivation from donation after cardiac death, ABO incompatibility, hepatic artery thrombosis and cytomegalovirus infection. Risk factors for recurrent PSC include IBD, colectomy in patients with overlapping IBD, CCA before LT and history of acute cellular rejection (65,66).

Post-transplant patient and graft survival rates for PSC and other autoimmune liver diseases are outstanding. For instance, in a study by the Mayo Clinic involving 150 consecutive patients with PSC (65), the one- and five-year patient and graft survival rates were 93.7%, 86.4%; and 83.4% and 79%, respectively. However, as part of the informed consent process for LT, patients with PSC should be aware that the recurrence rate of PSC after LT is approximately 20% after 10 years (66). To establish the diagnosis of recurrent PSC, causes of nonanastomotic biliary strictures must be excluded, including but not limited to, ischemic injury of the hepatic graft, graft derivation from donation after cardiac death, ABO incompatibility, hepatic artery thrombosis and cytomegalovirus infection. Risk factors for recurrent PSC include IBD, colectomy in patients with overlapping IBD, CCA before LT and history of acute cellular rejection (65,66).

With the improved survival of patients after LT regardless of underlying etiology, post-transplant malignancy is now a prevailing cause of late morbidity and mortality. Based on prospective data from the liver transplant database of the National Institute of Diabetes and Digestive and Kidney Diseases (67), the risk of skin and nonskin cancer after LT is, in fact, highest in recipients transplanted for PSC; these patients had a 19% incidence of skin cancer and 22% incidence of nonskin malignancy within 12 years of LT. LT recipients with a history of PSC are clearly at higher risk for CRC because of the added risks imposed by concomitant IBD and immunosuppression, but in addition to gastrointestinal malignancies, these patients experience more lymphoproliferative and skin malignancies than patients transplanted for other etiologies (67).

LT for hilar cholangiocarcinoma is an important consideration for the unfortunate patient with PSC-associated early hilar CCA in the context of a research protocol at a specialized CCA centre. Few tumours of hilar CCA are amenable to surgical cure, and the five-year survival rate for resection or transplantation alone is only approximately 30% (68,69). Due to the lack of acceptable treatment options, the Mayo Clinic pioneered a novel protocol using neoadjuvant therapy and LT in 1993. This specialized protocol combines the benefits of radiotherapy, chemosensitization and LT in select patients with unresectable hilar CCA fulfilling specific inclusion criteria (70). While most patients with hilar CCA would be outside of transplantation criteria at diagnosis due to the aggressive nature of the tumour, for
patients with early diagnosis and tumour responsive to neoadjuvant treatments, the one- and five-year patient survival rates are excellent (91% and 73%, respectively) (70).

SUMMARY

Our understanding of the pathogenesis of PSC is evolving rapidly, and the management of PSC is rife with conventions and controversies. To date, there are no proven medical treatments for PSC, and there is an urgent need for collaborative research in this area to provide efficacious therapeutic options to improve the associated high morbidity and mortality. Although high-dose UDCA has been refuted as a safe and effective agent for PSC, the role of low-dose UDCA requires clarification and remains disputed. The management of PSC is largely focused on the prevention and treatment of complications such as cholangitis, pruritus, malnutrition, hepatic osteodystrophy and, where overlap of IBD exists, prevention of CRC. Despite its rarity, and the fact that the majority of patients with PSC will never be affected, CCA is a grave concern in patients with PSC and, hence, a high index of suspicion should be maintained. LT remains the only effective and proven treatment for PSC, with five-year patient survival rates approximating 85% in the current era of transplantation. Select patients with limited-stage hilar CCA may benefit from LT with extensive neoadjuvant therapy, but such a specialized protocol remains experimental.

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