

Primary Biliary Cholangitis (Primary Biliary Cirrhosis) Diagnosis and Treatment

Primary Biliary Cirrhosis (PBC) is an inflammatory autoimmune reaction leading to destruction of intrahepatic bile ducts. As bile ducts are destroyed, cholestasis results, and over time this may lead to cirrhosis and liver failure. Patients typically have antimitochondrial antibodies present, along with elevated serum alkaline phosphatase and increased cholesterol. Treatment includes ursodiol and liver transplant.



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Diagnosis

Positive Antimitochondrial Antibodies

[Positive Anti-Mitochondrial-factory Ant-tie-body](#)

95% of patients with PBC show increased serum antimitochondrial antibodies, which are the characteristic autoantibodies in this disease. These stimulate autoreactive T cells to accumulate around bile ducts and hepatocytes.

Increased Alkaline Phosphatase (ALK-P)

[Up-arrow Elk-P](#)

Studies will also show increased serum alkaline phosphatase (ALP or ALK-P), which is a marker of cholestasis.

Increased Cholesterol

[Up-arrow Cholesterol-burger](#)

In PBC, and other situations where cholestasis occurs, the ability to transport and breakdown cholesterol is impaired. This yields increased serum cholesterol.

Treatment

Ursodiol

[Usher-doll](#)

This is the only FDA-approved medication to treat PBC, and works by reducing the rate of intestinal cholesterol absorption. It also works to nonsurgically treat gallstones, while limiting symptoms of cholestasis (jaundice, pruritis).

Liver Transplant

[Liver Train-plant](#)

The onset of this disease is insidious. It eventually leads to liver cirrhosis and an increased risk of hepatocellular carcinoma. Liver transplant is the best form of treatment in patients with end-stage liver disease.