Antimitochondrial Antibodies: from Bench to Bedside

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Accepted: 13 September 2021 / Published online: 29 September 2021 © The Author(s), under exclusive licence to Springer Science+Business Media, LLC, part of Springer Nature 2021

Abstract

Anti-mitochondrial antibodies (AMA) are directed against the E2 subunits of the 2-oxo acid dehydrogenase complexes (PDC-E2) and are the typical biomarkers of primary biliary cholangitis (PBC), being present in 90–95% of patients, with increasing sensitivity at increasing titers. Albeit being highly specific for PBC diagnosis, AMA can be detected in less than 1% of healthy subjects, and thus the management subjects with no sign or symptom of liver disease is still a challenge and data concerning clinical risk of developing PBC in this subgroup of patients are controversial. Moreover, AMA can also be detected in patients affected by overlap syndrome, as well as hepatic diseases (i.e., NASH and viral hepatitis), while the association with autoimmune diseases, in particular Sjögren’s syndrome, systemic sclerosis, and systemic lupus erythematosus, is well established. Furthermore, new associations are being identified with inflammatory myositis and heart disease. AMA are directed towards the pyruvate dehydrogenase multi enzyme complex (PDC-E2) subunit, which represents an epithelial specific autoantigen for PBC. This review focuses on the main characteristics of AMA, their association with autoimmune diseases and liver diseases.

Keywords Autoimmunity · Primary biliary cholangitis · Systemic sclerosis · Myositis

Introduction

Autoantibodies represent the hallmark of autoimmune diseases (AID), playing a key role in the first steps of diagnostic approach to AID. Interestingly, autoantibodies may be present years before clinical manifestations develop [1].

Anti-mitochondrial antibodies (AMA) target lipoic acid containing immunodominant epitopes, particularly the E2 subunits of the 2-oxo acid dehydrogenase complexes (PDC-E2). However, AMA may target several antigens within the inner mitochondrial membrane, with no fully clear diagnostic and clinical significance known so far. PDC-E2 and E2 subunits of other mitochondrial autoantigens contain an essential lysine residue within the lipoyl domain to which lipoic acid is covalently attached. This lipoic-lysine bond at position 173 is highly conserved across species and is necessary for antigen recognition [2].

At first, AMA were detected for a non-organ specific ATP-ase associated antigen called M2, consisting in several mitochondrial enzymatic polypeptides of which the dihydrolipoamide acetyltransferase (E2 component) of the pyruvate dehydrogenase multi enzyme complex (PDC-E2) is the most identified by AMA (M2-AMA) [3].

 Afterwards, other subunit-specific AMA were identified all against components of the 2-oxoacid dehydrogenase (2-OADC) family of enzymes within the mitochondrial respiratory chain: the subunits-specific AMA type PDC-E2, BCOADC-E2 (E2 subunit of the branched-chain 2-oxo acid dehydrogenase complex) and OGDC-E2 (E2 subunit of the 2-oxo-glutarate dehydrogenase complex). Besides E2 subunit, E1 and E3 are all subunits of these complexes participating in oxidative phosphorylation and have a great deal of homology, all of them containing the motif DKA, with lipoic acid covalently bound to the lysine (K) residue [4]. Interestingly, PDC includes a fourth component, protein X (PDC-E3 binding protein, PDC-E3BP), which copurifies with E2.

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Detection Methods

The detection of AMA is currently recommended by the use of indirect immunofluorescence (IIF) on HEp-2 cells (Fig. 1) [5]. However, several multiplex assays, using methodological platforms such as enzyme-linked immunosorbent assay (ELISA), are becoming more common in diagnostic laboratories. An ELISA test directed towards M2-AMA showed a lower specificity than IIF, although being slightly more sensitive in biopsy-proven PBC [6]. Moreover, newer ELISA kits recognizing not only the M2 subunit, but also including the 3 E2-subunits as a target, showed better performances [7]. Recently, dot blot has been proposed as a new testing for AMA with good performances [8]. These tests, however, need to be validated in large cohorts, and are therefore not recommended, and the gold standard for AMA detection remains IIF.

Thanks to the improvement of diagnostic methods, nine mitochondrial antigen/antibody patterns —from M1 to M9 — have been described [9] being described not only in hepatic disorders but also in non-hepatic diseases. Interestingly, only M2, M4, M8, and M9 are specific for PBC [10]; however, anti-M4 and anti-M8 positivity probably represents artifacts of the methods used to detect AMA since they are both targets of AMA-M2 and are predictors of an elevated immunological activity of the disease. On the other hand, anti-M9, firstly described in 1984 by Klein and Berg, represents a markers of PBC even in anti-M2 PBC sera [11].

Anti-M1 ad anti-M7 have been detected in infectious disorders such as syphilis or myocarditis, while anti-M3 and M6 are associated with drug allergic reactions; anti-M5 have been detected in several collagen disorders [9]. Both M1 and M5 will be discussed later as they are associated with antiphospholipid syndrome.

Clinical Associations

Anti-mitochondrial Antibodies in Liver Diseases

Primary Biliary Cholangitis

PBC, previously known as primary biliary cirrhosis [12], is a chronic and cholestatic liver disease affecting mostly middle-aged women [13]. It is characterized by progressive, non-suppurative cholangitis involving biliary ducts, which are consequently destructed; the severe inflammation eventually leads to fibrosis and thus to liver cirrhosis with its complications [14]. Once the immune response is established, aberrant autoantigens are expressed on biliary epithelial cells, which may lead to an increased presentation to autoreactive T cells. After autoantigens have been presented, a multi-lineage T and subsequently B cell response develops. When symptomatic, up to 70% of patients present with fatigue or pruritus (usually at night). Rarely the onset of the disease is represented by the complications of cirrhosis such as ascites, jaundice, hepatic encephalopathy, upper gastrointestinal bleeding, or hepatocellular carcinoma [15].

The diagnosis of PBC relies on three main elements: 1) biochemical liver cholestasis with elevation of alkaline phosphatase (ALP) 2) detection of anti-mitochondrial antibody, 2-oxo-glutaric acid dehydrogenase complex (PDC-E2), and PBC-specific anti-nuclear antibodies ANA (anti-GP210, and/or anti-SP100) 3) suggestive histology obtained through liver biopsy [16]. Of note, AMA detection is often associated with a non-specific elevation of serum IgM concentration [10, 17].

The etiology of PBC is still obscure, but many data suggest that a genetic predisposing background together with infective, immunological and environmental risk factors may explain the onset of the disease [18]. Several studies have reported a strong link with HLA alleles, in particular

![Fig. 1](image-url)
DRB1*08, DR3, DPB1*0301, DRB1*08-DQA1*0401-DQB1*04, whereas DRB1*11 and DRB1*13 seem to be protective. Furthermore, non-HLA genes have been associated with PBC development, i.e., genes involved in the regulation of the immune system and antigen presentation, such as IL-1RL1, STAT4, STAT1, and IL12A [19]. Concerning environmental factors, infections are thought to increase the susceptibility to PBC based on molecular mimicry which ensues when an infectious agent, bacterial or viral, presents antigens with a significant amino acid similarity to self-proteins [20]. Furthermore, smoking, the use of hormonal replacement therapy and history of recurrent urinary tract infections have been associated with an increased risk of PBC development. Moreover, xenobiotics, such as nail varnish, hair dye, and cleaning chemicals can break self-tolerance [21].

For unknown reasons PDC-E2 is aberrantly expressed in the luminal surface of bile duct epithelial cells of PBC patients, in a pathogenetic process that has been entitled “autoimmune epithelitis”, shared with Sjögren’s syndrome (Fig. 2) [20]. CD8+ T cells do recognize this epitope and are responsible for subsequent bile duct damage together with accumulation of bile acids, which reach toxic concentrations due to cholestasis [22, 23]. In addition to the T-lymphocyte mediated destruction of small bile ducts, secondary damage to hepatocytes may occur from the accumulation in the liver of increased concentrations of potentially toxic substances, such as bile acids, which are normally secreted into the bile. The naturally occurring bile acids (cholic acid, chenodeoxycholic acid, and deoxycholic acid) are all detergents and can dissolve cell membranes if present in a sufficiently high concentration; such toxic concentrations are often reached in cholestasis [23].

AMA-M2 represent the hallmark of PBC, being present in 90–95% of patients [24], and less than 1% of healthy subjects. Similar to other AID, AMA positivity might arise years before the development of PBC. Interestingly, IF-AMA titers display a predictive value for PBC diagnosis, as titers of 1:320 and 1:640 increase the OR of 4.93 and 7.67, respectively. Studies demonstrate M2-AMA dot-blot to be more specific than IIF-AMA and suggest considering the determination of the number of M2-AMA specificities for PBC diagnosis [8]. Poyatos et al. recently observed that the combination of subunits-specific AMA positive increases both specificity and sensibility for PBC diagnosis (two M2-AMA specificities OR 2.05, $p<0.03$; three M2-AMA

![Fig. 2](image-url) Proposed pathogenetic mechanisms of primary biliary cholangitis APC, antigen presenting cell; PDC-E2, pyruvate dehydrogenase complex; IL, interleukin; IFN, interferon; AMA, anti-mitochondrial antibodies; BEC, biliary epithelial cells

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Nevertheless, AMA testing is not specific for PBC, with AMA positivity up to 1/1000 in general population compared to PBC prevalence up to 0.4/1000 (Table 1) [13]. Thus, management of subject tested positive for AMA with no sign or symptom of liver disease is still a challenge and data concerning clinical risk of developing PBC in this subgroup of patients are controversial. An old report by Metcalf et al. showed that up to 75% of AMA positive patients develop PBC after 18 years, with abnormal cholestatic liver tests at median time of 2 and 5 years, respectively, from the first AMA detection [25]. Other studies report lower rates of PBC diagnosis in AMA positive subjects, but with a mean follow-up of 5.8 years [26, 27]. Both the study designs and the clinical course of the disease, which is slow and progressive, may explain the contrasting results. However, development of overt PBC in AMA positive subject is associated with risk factors such as high titer AMA, increased avidity of PDC-E2, and detection of autoantibodies targeting multiple cell domains [28]. Aside from its high specificity for PBC, AMA-M2 can be detected in a low percentage of autoimmune hepatitis type 1 (AIH) and SSc patients [29]. Although these are often cases of overlapping PBC, the measurement of AMA is certainly relevant for patients suspected of having PBC and may be critical in the prediction of PBC development for patients suffering from other autoimmune manifestations.

Despite their diagnostic value, AMAs have no prognostic value, nor their titer or their subtypes are associated with disease severity and outcome [27, 30, 31]. Since the titer usually remains stable over the years, serial testing during PBC follow-up is not recommended so far and does not have any predictive significance. Thus, even if treatment with ursodeoxycholic acid (UDCA) may decrease AMA titer, there is conflicting evidence that this effect is associated with treatment response [28, 32, 33]. Previous studies also reported that AMA could be detected in nearly all sera of patients transplanted for PBC, without being associated with histological recurrence of PBC [34].

Interestingly, almost 5–10% of PBC patients do not display detectable AMA using routinely methods, and are referred to as “AMA-negative”; remarkably, some cases of AMA negativity by IF result positive by western blot or ELISA [35, 36]. On the other hand, some studies have showed higher rates of ANA and ASMA positivity in the

### Table 1

| Study                  | Year | Country   | Population of study                          | Total subjects | Prevalence | Reference |
|------------------------|------|-----------|----------------------------------------------|----------------|------------|-----------|
| Assassi                | 2009 | USA       | Systemic sclerosis                           | 817            | 7%         | [60]      |
| Cavazzana              | 2011 | Italy     | Systemic sclerosis                           | 201            | 21.4%      | [65]      |
| Ceribelli              | 2018 | Italy     | Systemic sclerosis                           | 85             | 15%        | [29]      |
| Generali               | 2019 | Italy     | Healthy population                           | 2685           | 3.8%, high-titer 1% | Unpublished data |
| Hatzis                 | 2008 | Greece    | Sjogren’s syndrome                           | 410            | 5.1%       | [101]     |
| Imura-Kumada           | 2012 | Japan     | Systemic sclerosis                           | 225            | 16.4%      | [102]     |
| Ranua                  | 2005 | Finland   | Epileptic patients vs. population-based controls | 948 + 579     | 3.9% vs. 1.9% | [103]     |
| Laadhar                | 2007 | Tunisia   | Type 1 diabetes                              | 161            | 0          | [104]     |
| Lazaridis              | 2007 | USA       | First degree PBC relatives                   | 306            | 13.1%      | [105]     |
| Li                     | 2015 | China     | Chronic hepatitis B                          | 325            | 6.8%       | [106]     |
| Li                     | 2006 | China     | Systemic lupus erythematosus                 | 365            | 15.3%      | [71]      |
| Liu                    | 2018 | Meta-analysis | Vitiligo                                         | \             | 0.2%       | [107]     |
| Maeda                  | 2012 | Japan     | Inflammatory myopathies                      | 212            | 11.3%      | [96]      |
| Marconcini             | 2013 | Brazil    | Chronic hepatitis C                          | 66             | 7.6%       | [108]     |
| Mattalia               | 1998 | Italy     | Healthy population                           | 1530           | 0.5%       | [109]     |
| Pelka                  | 2021 | Poland    | Sub-acute cutaneous lupus erythematosus       | 30             | 20%        | [110]     |
| Pope                   | 1999 | Canada    | Systemic sclerosis                           | 61             | 6.6%       | [111]     |
| Ramos-Casals           | 2005 | Spain     | Chronic hepatitis C                          | 237            | 8%         | [112]     |
| Shibata                | 2004 | Japan     | Corporate workers                            | 1714           | 0.64%      | [113]     |
| Beretta-Piccoli        | 2018 | Switzerland | Acute hepatitis E                             | 48             | 2.1%       | [114]     |
| Turchany               | 1997 | Estonia   | Healthy population                           | 1461           | 0.89%      | [115]     |
| Wielosz                | 2016 | Poland    | Systemic sclerosis                           | 86             | 13%        | [116]     |
| Yokokawa               | 2021 | Japan     | Dilated cardiomyopathy                       | 270            | 1.1%       | [117]     |
| Zografos               | 2012 | Greece    | First degree PBC relatives                   | 101            | 18.8%      | [118]     |
AMA negative group as well as lower IgM levels [37, 38]. Current literature has not evidenced any difference between the two groups concerning histology, clinical features, biochemistry, and response to UDCA [39]. Conversely, AMA negative subjects may have a more progressive disease due to delayed diagnosis and UDCA treatment [40]. Concerning ANA, specific immunofluorescence pattern might help to diagnose patients. Anti-sp100, included in the “multiple nuclear dots” (MND) pattern, and anti-gp210, included in the “perinuclear rims pattern”, are both specific for PBC; both correlate with disease severity, too [41].

Despite being rare, there are cases of variations of AMA in PBC, with AMA-positive subjects that may become AMA negative, and vice versa; this could be due to interference with the immune system and the onset of a new AID.

**Overlap Syndromes**

Autoimmune liver diseases, similarly to rheumatic disease, may overlap and present with both hepatocellular and cholangiocellular patterns according to biochemical, histologic, and imaging-based analysis. When left without treatment, these patients show a more progressive course toward liver cirrhosis and failure [19]. Despite PBC is featured with a certain degree of hepatic inflammation, it is usually minimal at histology. However, even after many years from diagnosis, PBC patients on treatment with UDCA (or second line obeticolic acid) may develop increased hepatitis activity; otherwise, features both of PBC and AIH may be observed from the diagnosis. In both cases, the condition is referred as “PBC-AIH overlap syndrome”, defined by the Paris Criteria, and is observed in almost 8–10% of PBC patients (Table 2).

The clues to an overlap syndrome consist of (i) serum alkaline phosphatase level more than twofold the upper limit of normal (ULN) at presentation, which is present in only 20% of patients with AIH; (ii) serum GGT level greater than ULN unimproved or worsened during therapy, (iii) AMA positivity, (iv) histologic findings of bile duct injury or loss, (v) concurrent IBD, and (vi) corticosteroid treatment failure or incomplete response [42].

On the other hand, patients know that AIH may develop signs of PBC; these subjects may test positive for AMA at diagnosis, but titers usually decrease over time. Also, in case of acute hepatitis activity, bile duct damage could be due to inflammation rather than AMA. Besides overlaps, transitions are also possible in rare cases from PBC to AIH, or AIH to PBC [43].

**Non-Alcoholic Steatohepatitis**

Non-alcoholic fatty liver disease (NAFLD) is the most common liver disease nowadays, mainly associated with metabolic syndrome; it is characterized by steatosis, ballooning, and lobular inflammation, typically developing around the central veins [44]. When inflammation develops, NAFLD may evolve to non-alcoholic steatohepatitis (NASH) and eventually to cirrhosis and its complications; however, involvement of the biliary ducts is rare.

Previous studies have reported the detection of autoimmune markers (ANA, AMA, ASMA) in NAFLD, with prevalence ranging from 12 to 48%, being higher than general population [45, 46] [47]. Clinical significance of these findings remains unclear so far, with controversial data concerning the disease course and outcomes.

Liver inflammation induced by NASH may potentially cause immunological cross-reactivity and induce AMA production, thus bile duct damage. As NASH patients may complicate with PBC or other autoimmune liver diseases, current guidelines recommend liver biopsy whenever in the presence of high titers of autoantibodies [48]. However, it is reasonable to consider autoantibody detection an epiphenomenon of NASH rather than a sign of underlying autoimmune liver disease in this subgroup of patients. In 2015, Ravi et al. identified 4 (1%) AMA-positive patients upon a NAFLD cohort

### Table 2: Diagnosis of overlap syndromes

| Laboratory features | Histologic findings |
|---------------------|---------------------|
| **AIH/PBC**         |                     |
| ANA or SMA          | Interface hepatitis |
| Hypergammaglobulinemia | Lymphocytic portal infiltrate |
| Serum IgG level increased | Portal plasma cells |
| Marked serum AST/ALT abnormalities | Destructive cholangitis |
| ALP or GGT > ULN    |                     |
| AMA positive        |                     |
| **AIH/PBC (Paris Criteria)** |             |
| *AIH features (2 of 3):*  | Interface hepatitis (moderate to severe) |
| Serum ALT > fivefold ULN | Destructive cholangitis |
| Serum IgG > twofold ULN or SMA present |                     |
| Interface hepatitis |                     |
| *PBC features (2 of 3):*  |                     |
| Serum ALP > twofold ULN or GGT > fivefold ULN |                     |
| AMA positive        |                     |
| Florid duct lesions |                     |
of 398 patients with known autoimmune profile. Of note, 3 AMA patients had available liver histology, and two of them did not display any feature of PBC and had normal ALP levels; interestingly, AMA, ANA, or ASMA positivity was not associated with any difference in clinical presentation or natural history of NAFLD [49].

Furthermore, several data have been published concerning transient AMA positivity in patients with drug-induced liver injury (DILI). A recent study from Weber et al. shows that AMA positivity in a cohort of 143 patients with DILI is 10%; this subgroup of patients had taken non-steroidal anti-inflammatory drugs more frequently and showed a more severe degree of liver injury, with higher aminotransferases, bilirubin and PT. However, data are scarce so far and future observations are required to look for any correlation with involved drug or clinical/biochemical characteristics [50].

Viral Hepatitis

AMA positivity in the absence of PBC has been observed in viral hepatitis, too. Chronic HCV hepatitis (CHC) has already been associated with the presence of autoantibodies such as ANA, rheumatoid factor, ASMA, AMA, anti-liver-kidney microsomal antibody (LKM), and anti-soluble liver antigen (SLA); however, their clinical significance remains unclear. Literature reports controversial results concerning difference in demographic factors and clinical outcomes between CHC patient with and without autoantibodies [51–55]. Most of these studies showed sustained viral response (SVR) rates to be not significantly different between the two groups except for two studies in which rates were lower in CHC autoantibody positive; one of them showed interferon (IFN)-based regimen to be safe and effective in this setting, even if no patients included had AMA detection [51]. In 2018, Gilman et al. investigated a large cohort of HCV-infected patients tested positive for autoantibodies with the aim to identify potential association with cirrhosis and its complication such as hepatocellular carcinoma (HCC) [56]. Overall, 2.8% of the population resulted AMA positive with no significant difference concerning sex nor ethnicity; also, the presence of an autoantibody had no impact on the prevalence of extrahepatic manifestation or natural history of CHC, except for SVR rates in IFN-based regimen [51, 56].

Antimithocondrial Antibodies Beyond the Liver

One in three PBC patients have a concomitant extrahepatic autoimmune disease, which may include rheumatological, endocrinological, gastrointestinal, pulmonary, or dermatological conditions. Among these, rheumatic diseases show an interesting overlap also in disease pathogenesis and treatment. Besides PBC, AMA positivity has been detected in other autoimmune disorders such as SSc, SLE, and SjS [27]. To note, PBC is associated with at least one concurrent extrahepatic autoimmune disorder in up to 61% of cases [57], with thyroid disease being one of the most common, especially in AMA-positive patients with an overall prevalence of 78% [58]. In a recent meta-analysis, extrahepatic autoimmune disorder is more frequent in AMA-positive PBC patients (88% vs. 84%, p = 0.05) and are associated with better biochemical analysis and early stage of disease at diagnosis [59].

Systemic Sclerosis

Up to 12% of PBC patients have coexisting SSc, and a large proportion of patients with SSc manifest a coexisting autoimmune condition, with PBC being the most frequent autoimmune liver disease, even though in clinical practice a higher percentage of SSc patients may have biochemical liver abnormalities without clinical significance [60–63]. SSc is an autoimmune disease of unknown origin characterized by microvascular damage and progressive fibrosis of skin that in severe cases can affect internal organs such as heart, lungs, and kidneys. The most frequently identified autoantibodies in SSc are anticientromere (ACA) — (30%), anti-topo I (30%), and -RNA pol III (4–20%). However, several other autoantibodies can be identified in SSc patients and they can have an important role for their clinical and prognostic value [64]. It has been recently reported that AMA prevalence might be higher than expected in SSc patients, when sensitive techniques are used. In particular, using immunoprecipitation (IP) to test new and uncommon serum autoantibodies in SSc, it was observed that 15% of SSc sera manifests a novel IP pattern, comprising a set of 4 proteins corresponding to the E1α, E1β, protein X/E3BP, and E2/E3 subunits of PDC recognized by AMA [29]. Even higher prevalence of PBC-related autoantibodies in SSc patients have been reported by ELISA, with up to 25% of patients showing a positivity for anti-MIT3, gp210, and Sp100 antibodies (PBC-Screen), with higher risk of developing PBC or elevation of ALP; however, these tests need further validation [65].

Interestingly, when PBC and SSc coexist they are associated with the limited cutaneous form of SSc and positivity of ACA, and liver disease does not seem to worsen the prognosis of our AMA-positive patients with SSc [61].

Systemic Lupus Erythematosus

Systemic lupus erythematosus (SLE) is a chronic, autoimmune disease affecting women predominantly, especially between puberty and menopause [66]; it is characterized
by a broad spectrum of clinical manifestations; however, its course and organ involvement are unpredictable. Similarly to other AID, the pathogenesis of SLE is multifactorial, with environmental factors acting in genetically predisposed individuals leading to the breaking of self-tolerance and impairment of the immune system; any organ or tissue may potentially be affected. In particular, B cell regulation may be involved and contribute with the production of autoantibodies which play an essential role in SLE, especially when they take part in immune complex formation [67].

SLE is characterized by the expression of autoantibodies to a wide variety of cellular antigens; ANA is for sure the most highly characteristic ones, followed by antibodies to native DNA (anti-dsDNA) and to Sm nuclear antigen (anti-Sm/RNP), all included in the classification criteria. In fact, diagnosis of SLE is made on clinical manifestations and laboratory tests, functional tests and imaging according to either American College of Rheumatology Revised Criteria (ACR 2012) or Systemic Lupus International Collaborating Clinics classification criteria (SLICC 2012) [68–70]. Interestingly, autoantibody specificities overlap between the different clinical manifestations of SLE and positivity for a specific antibody does not necessarily mean that a certain organ will be affected; however, some autoantibodies are associated with peculiar clinical manifestations. Despite not having diagnostic clinical value in SLE patients, AMA are present in this subgroup of patients but with low prevalence. Li et al. found a significant percentage of AMA positivity in their Chinese cohort of SLE patients (approximately 15%), but several other studies on European cohorts did not confirm such significant data maybe due to environmental or genetic background [71].

**Antiphospholipid Syndrome**

Antiphospholipid syndrome (APS) is a syndrome characterized by frequent and often multiple thromboses, both venous and arterial, and pregnancy morbidity; these clinical manifestations are usually associated with moderate thrombocytopenia [72].

Anti-cardiolipin antibodies (aCL) together with lupus anticoagulant (LA) are specific anti-phospholipid antibodies typical of APS may be primary, thus isolated findings, or secondary to other diseases, with SLE being one of the most common ones [73]. Interestingly, AMA-M1 target an antigen identical to cardiolipin, a phospholipid found uniquely in the inner mitochondrial membrane, and were originally identified in syphilis-infected patients [74]. Thus, the presence of AMA-M1/aCL clinically is associated with a greater risk of thrombotic events and thrombocytopenia. AMA-M5 have been identified in only 2% of patients by IIF, but specific antigenic target of these antibodies is undefined so far, even if lack of competition by cardiolipin-containing liposomes suggests, it is different from AMA-M1. They have been detected in patients with APS, SjS, and hemolytic anemia.

Recent evidence supports the fact that AMA are represented in the autoantibody repertoire associated with APS, may display different associations with the clinical manifestations of the disease. In particular, anti-DNA-IgM were associated with a lower likelihood of reporting a previous arterial thrombosis, and their levels were not influenced by the duration of the disease, cardiovascular risk factors or medications [75].

Weather AMA detection in SLE patients is associated with future development of PBC remains a question to be answered, with little risk identified so far (<1%), with most of the patients having liver enzymes within the reference limit, or mild laboratory cholestasis [76, 77].

Interestingly, a recent pre-clinical study revealed a greater reactivity of antibodies in SLE sera to intact mitochondria rather than the M2 subunits to the 2-oxo-acid dehydrogenase protein complex, in clear distinction with sera of PBC patients. On the other hand, in the same analysis no significant binding of the PBC sera was found to intact mitochondria meaning PBC target antigens are not easily and immediately accessible [78]. Specifically, AMA-M2 are detected in approximately 10% of SLE patients, with not fully clear functional and clinical significance; so far, literature lacks solid, large cohort-based studies. However, emerging evidence suggests that SLE is characterized by activation of inflammation due to liberation of mitochondria; thus, further identification of mitochondrial antigens target by autoantibodies is needed to understand the role of mitochondrial in autoimmunity.

**Sjögren’s Syndrome**

Sjögren’s syndrome is a chronic, systemic autoimmune disease, which can be primary or secondary to other systemic AID, i.e., RA, SLE, SSc, mixed connective tissue disease, inflammatory muscle disease, autoimmune liver disease, and autoimmune thyroid disease [79, 80]. SjS is the most common co-occurring condition in PBC, in proportion ranging from 3.5 to nearly 100% of cases [57].

The pathogenesis of SjS is currently based on the concept of “autoimmune epithelitis”, similarly to PBC, since in SjS the exocrine glands, especially salivary and lacrimal, are progressively destructed by an immune-mediated process [20, 81]. PBC and SjS share a common immunopathogenesis in which genetics and environmental factors interact to result in disease onset inducing salivary or biliary epithelial cell apoptosis and contributing to the breakdown of tolerance to self-antigen exposed to the apoptotic blebs [20].

The diagnosis of SjS should be suspected in case of sicca syndrome symptoms, and an objective evaluation of eye dryness is obtained with a Schirmer’s test which measures the
normal tear production by the positioning of a strip of sterile filter paper overhanging the lateral third of the lower eyelid of each eye with a ≤ 5 mm considered positive [82].

Systemic extraglandular manifestations are present in 10–15% of patients with SjS, and arise from diverse pathogenetic mechanisms, being the result of the typical lymphocytic infiltration around the epithelium of target organs such as the liver, the kidney, and the bronchi/bronchioles, or the systemic form of the disease that arises from immune-complex deposition, presenting with the clinical picture of vasculitis. Other extra-glandular manifestations include arthralgia and arthritis, and fatigue [81].

Anti-SSB are usually found with anti-SSA antibodies, while the only specificity in 25–40% of cases [83], and are not included in the latest classification criteria [84]. ANA, albeit not specific for SjS, are present in up to 90% of cases, while RF is present in up to 70% of cases [83]. Histologic examination remains the gold standard for the diagnosis of SjS. Among SjS patients without overt autoimmune liver disease, up to 20% have detectable AMA, while 10% of AIH-associated autoantibodies. Interestingly, previous studies reported that the association of PBC with SjS was associated with poorer prognosis and increased mortality [57, 85]. However, a recent study showed that the response and clinical outcomes of UDCA-treated patients with PBC are not adversely affected by concomitant SjS [86].

Other Autoimmune Diseases

There are several other, less recognized diseases that have higher prevalence among patients with PBC. Among others, rheumatoid arthritis has been associated with PBC, but epidemiological data only suggests a prevalence rate between 1.8% and 5.6% [87, 88]. Data from national Danish and Swedish registers have shown an association between PBC and celiac disease, with up to 6% of patients having also celiac disease [89, 90]. In contrast, other European studies from Poland and Italy have shown no prevalence greater than that seen in the general population [91, 92].

Unexpected Associations

Inflammatory Myositis

In the recent years, AMA have been associated with inflammatory myopathy, a heterogeneous disease entity affecting primarily the skeletal muscles. Myositis-specific antibodies as well as myositis-associated antibodies have been described to date, each associated with distinct clinical serological syndromes [93].

Myositis-specific antibodies are found only in patients with polymyositis, dermatomyositis, and immune-mediated necrotizing myopathies and have a strong association with clinical disease, meaning that patients with these specificities frequently have unique features characteristic of that antibody [93]. Conversely, myositis-associated antibodies are less well-defined, and are generally accepted to be those antibodies found in immune and inflammatory myopathies that can also be found in other AID, such as anti-PM-Scl, Ro52, and U1RNP. Having the antibody does not always correlate with the presence of inflammatory muscle disease, while other clinical associations can be seen [94]. These antibodies can be found alone, or in conjunction with other myositis-specific antibodies, and can also have typical clinical presentations. AMA are rarely present in inflammatory myopathy, but there is a growing recognition for this clinical entity [95].

Furthermore, AMA have been extensively associated with serious cardiac involvement [96, 97]. In a large North American study on inflammatory myopathy, albeit AMA prevalence was extremely low (0.006%), it was associated with a distinct phenotype with severe cardiac involvement, as severe conduction abnormalities and arrhythmias, cardiomyopathy, ventricular dilations, and myocarditis, occurring all in the same patient [97]. Other authors suggested that AMA-associated myopathy mimics sarcoidosis heart involvement, based on magnetic resonance findings, showing left ventricle systolic dysfunction [98]. Of interest, in the case of cardiomyopathy and myocarditis, treatment with immunosuppression could improve cardiac function. Fascinatingly, AMA have been reported in a significant proportion of patients with myocarditis as well as idiopathic dilated cardiomyopathy, where the most abundant protein in the inner mitochondrial membrane has been shown to be an organ specific autoantigen [99].

SARS-CoV-2 Infection

Since early 2020, SARS-CoV-2 infection has dramatically changed our perspectives. Of note, a case of PBC developing during or soon after COVID-19 has been reported [100]. Remarkably, the patient was diagnosed after being admitted for severe COVID-19 with the need of mechanical ventilation and treatment with tocilizumab, ceftiraxone, azithromycin, darunavir/cobicistat, anakinra, remdesivir, and fluconazole. During hospitalization, both GGT and ALP levels, that were normal at admission gradually increased, while serum bilirubin and ALT fluctuated between normal values and slightly elevated. An abdomen ultrasound showed a lightly enlarged liver with moderate steatosis and a mildly enlarged spleen, the liver stiffness measured with Fibroscan was 9.1 kPa. From the immunology standpoint, the patient tested positive for ANA and AMA (M2/BCOADC) at high titer. A liver biopsy was performed and showed florid ductal lesions, therefore leading to the definite diagnosis of PBC.
This was the first reported case of PBC induced by SARS-CoV-2, which is a RNA virus capable of inducing a profound activation of the immune system and also by the previous finding that infection by a human RNA beta-retrovirus, related to the mouse mammary tumor virus, was suggested as a possible trigger for PBC development [100].

Conclusions

The immune system scrupulously controls reactivity against self-antigens, and AID are relatively rare. Importantly, early diagnosis and treatment of AID can prevent morbidity and mortality and therefore specific diagnostic tools are critical. Currently, screening of AID (including PBC) relies heavily on the use of serological tests for the detection of serum autoantibodies, and a better understanding of such tests allows to ensure more efficient diagnosis. In these prospective, some issues regarding the clinical significance of AMAs remain to be addressed. First, determining the pathogenic role of AMA remains a priority for basic and clinical research in PBC. Indeed, although a pathogenic role in PBC has been extensively suggested, conclusive data are not available. Second, clinical implications of AMA presence in asymptomatic subjects remain an area for further discussion. Indeed, AMA represent the hallmark of PBC, being present in more than 95% of patients even years before the development of clinical manifestations and diagnosis. Moreover, AMA are also associated with other AID, in particular SjS, SSc, and SLE, while rarely detectable in healthy subjects. Finally, AMA positivity usually relies on the detection of autoantibodies directed towards the M2 subtype, and further research is needed to characterize the clinical significance of the other subtypes. With IIF being the gold standard for AMA detection, a newer technique validation is warranted to improve the test sensibility and therefore their clinical application.

Declarations

Conflict of Interest The authors declare no competing interests.

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