Hellenic Association for the Study of the Liver Clinical Practice Guidelines: Autoimmune hepatitis

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Abstract
Autoimmune hepatitis (AIH) is a relatively rare acute or chronic liver disease of unknown etiology characterized by large heterogeneity. Its distribution is global, covering all ages, both sexes and all ethnic groups. The aim of the present Clinical Practice Guidelines (CPG) of the Hellenic Association for the Study of the Liver was to provide updated guidance and help to gastroenterologists, hepatologists, internists and general practitioners for AIH diagnosis and management. AIH diagnosis is based on clinicopathological characteristics: namely, polyclonal hypergammaglobulinemia, particularly of immunoglobulin G (IgG), circulating autoantibodies, interface hepatitis on liver histology, absence of viral hepatitis, and a favorable response to immunosuppression. Clinical manifestations at disease onset are variable, ranging from asymptomatic to the acute/severe form. Aminotransferase and bilirubin levels vary, while the presence of hepatitis at the histological level is a prerequisite for diagnosis. Autoantibodies are the hallmark for AIH diagnosis; therefore, the CPG describe the appropriate serological algorithm for their detection. AIH therapy should aim to achieve complete biochemical (normalization of IgG and aminotransferases) and histological remission. All patients who have active disease, even those with cirrhosis, should be treated with individualized and response-guided induction therapy using prednisolone in combination with azathioprine or mycophenolate mofetil as first-line therapy. Immunosuppression should be given for at least 3 years and for at least 2 years after the achievement of complete biochemical response, while a liver biopsy should be recommended before treatment discontinuation. Current CPG are also provided for several specific conditions and difficult-to-treat patients.

Keywords Autoimmune hepatitis, autoantibodies, clinical practice guidelines, corticosteroids, azathioprine


Introduction and historical review
In 1950, Jan Waldenström was the first to describe a new type of a progressive, usually fatal, chronic hepatitis in young females with endocrine dysfunction, cutaneous striae, acne, polyarthralgias, and high γ-globulins that correlated with abundant plasma cells on liver biopsy [1]. In 1956, Cowling et al [2] introduced the term “lupoid hepatitis”, because the lupus erythematosus cell phenomenon was observed in these patients. However, 10 years later, this term was changed to “autoimmune hepatitis” (AIH), finally accepted by the International AIH Group (IAIHG) as the definite one [3,4].

The aim of the present Clinical Practice Guidelines (CPG) of the Hellenic Association for the Study of the Liver (HASL) was to provide guidance and help to gastroenterologists, hepatologists, internists and general practitioners in the diagnosis and management of this disease, in an attempt to improve care for affected patients. The current statements...
and recommendations were based on the GRADE system for evidence published by Shaneyfelt et al [5] (Suppl. Table 1).

**Epidemiology**

AIH is an acute or more frequently chronic liver disease of unknown etiology that primarily affects females (female/male: 3-4/1) and is characterized by polyclonal hypergammaglobulinemia, particularly of immunoglobulin G (IgG), autoantibodies, interface hepatitis and favorable response to treatment [6-10].

AIH is considered relatively rare, as its prevalence is about 160-180/1,000,000 population in Europe [11-14]. Recently, a large nationwide study in Denmark showed a significant increase in AIH incidence, reaching a prevalence of 350/1,000,000 in women [15]. So far, reliable epidemiological data from Greece are not available; therefore, since 2016 HASL has started a nationwide registry for all retro- and prospective AIH cases.

AIH carries several clinical phenotypes and outcomes according to ethnicity, with patients of Hispanic, Asian or other non-European Caucasian origin demonstrating poor outcomes [16]. These differences are thought to be due to differences in genetic predisposition and triggering agents, but also to complex socioeconomic factors, such as discrepancies in healthcare delivery and failure to diagnose AIH, which finally results in delayed diagnosis [17].

**Clinical manifestations**

**Clinical characteristics**

AIH is a discrete clinical syndrome characterized by considerable demographic, clinical, laboratory and histological heterogeneity (Table 1). Therefore, extended differential diagnosis should be performed, considering the possibility of AIH in any acute or chronic liver disease (Tables 2, 3) [4,6-8,10,18-22]. Both sexes in all ethnic groups can be affected at any age. Disease onset has a bimodal distribution during the childhood/teenage and 4th-6th decades, but recently many patients have been diagnosed at older ages (>65 years) [6-8,11-15,23-25]. The disease may accumulate in first-degree relatives, but the absolute risk is very low. High rates of depression and anxiety have recently been recognized in AIH patients [26,27].

The manifestations are variable, ranging from asymptomatic to acute/severe or even fulminant hepatitis.

**Table 1 Clinical manifestations of autoimmune hepatitis (AIH)**

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>Description</th>
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<tbody>
<tr>
<td><strong>Age at presentation</strong></td>
<td>Any age in both sexes (F/M: 3-4/1) and all ethnic groups can be affected; bimodal distribution with usual peaks in puberty and 4th-6th decades; a substantial proportion of patients, however, are older (&gt;65 years)</td>
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<tr>
<td><strong>Disease onset</strong></td>
<td>From asymptomatic to acute/severe or even fulminant hepatitis</td>
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<tr>
<td>Two thirds of patients present either without any symptom or with an insidious onset (one or more of the following unspecific symptoms: malaise, fatigue, amenorrhea, general ill health, lethargy, anorexia, right upper quadrant pain)</td>
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<td>Acute AIH in 25-40% of patients, presenting either as an acute worsening of chronic AIH or as real acute AIH without findings of chronic disease on liver histology; absence of autoantibodies detection or other usual features is not surprising; response to corticosteroids variable</td>
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<td>At diagnosis 1/3 of patients already have cirrhosis, regardless of the presence or absence of symptoms, suggesting a delay in diagnosis</td>
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<td><strong>Classification</strong></td>
<td>AIH-1: the more frequent type (90% of cases); ANA, SMA or anti-SLA/LP reactivity (the latter often with anti-Ro52 reactivity; potentially more severe); association with HLA DR3, DR4 and DR13; rare treatment failure but variable relapse rates after complete drug cessation and variable need for long-term maintenance therapy</td>
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<tr>
<td>AIH-2: approximately 10% of cases; anti-LKM1, anti-LC1 and rarely anti-LKM3 reactivity; association with HLA DR3 and DR7; onset usually in childhood and young adulthood; from the clinical and histological points of view this type is usually more acute and advanced; frequent treatment failure and frequent relapses after complete drug discontinuation; need for long-term maintenance therapy very common</td>
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<td><strong>Physical findings</strong></td>
<td>From completely normal to signs of chronic liver disease and/or portal hypertension</td>
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<tr>
<td><strong>Complications</strong></td>
<td>HCC rates are significantly lower than in other liver diseases, but it does exist in association with underlying cirrhosis, suggesting surveillance in all AIH-related cirrhotics</td>
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<tr>
<td>Significant treatment-related side-effects are found in 15-25% of patients (most commonly related with long-term corticosteroid use or toxicity and/or intolerance of azathioprine)</td>
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F, female; M, male; ANA, antinuclear antibodies; SMA, smooth muscle antibodies; anti-SLA/LP, antibodies against soluble liver antigens/liver pancreas; HLA, human leukocyte antigens; anti-LKM1, anti-liver/kidney microsomal antibody type-1; anti-LC1, antibodies against liver cytosol type-1 antigen; anti-LKM3, anti-liver/kidney microsomal antibody type-3; HCC, hepatocellular carcinoma

**STATEMENT 1**

- AIH prevalence is increasing in Europe irrespective of sex, ranging from 160-180/1,000,000 inhabitants to as much as 350/1,000,000 in females (II-2)
Table 2 Differential diagnosis of autoimmune hepatitis

<table>
<thead>
<tr>
<th>Condition</th>
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<tr>
<td>Acute or chronic viral hepatitis A, B, C, D, E</td>
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<tr>
<td>Drug-induced liver injury</td>
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<tr>
<td>Alcoholic liver disease</td>
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<tr>
<td>PBC, PSC, AIH/PBC variant, AIH/PSC variant</td>
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<tr>
<td>Wilson's disease</td>
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<tr>
<td>Non-alcoholic fatty liver/steatohepatitis</td>
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<td>Hemochromatosis</td>
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<td>α1-antitrypsin deficiency</td>
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<tr>
<td>Celiac disease</td>
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PBC, primary biliary cholangitis; PSC, primary sclerosing cholangitis; AIH, autoimmune hepatitis

Table 3 Common concurrent autoimmune or immune-mediated diseases in patients with autoimmune hepatitis

<table>
<thead>
<tr>
<th>Disease</th>
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<tr>
<td>Hashimoto thyroiditis – the strongest association</td>
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<td>Grave's disease</td>
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<td>Vitiligo, alopecia, psoriasis</td>
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<tr>
<td>Rheumatoid arthritis</td>
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<tr>
<td>Diabetes mellitus type 1</td>
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<tr>
<td>Inflammatory bowel disease</td>
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<tr>
<td>Systemic lupus erythematosus</td>
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<tr>
<td>Sjögren's syndrome</td>
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<tr>
<td>Celiac disease</td>
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<tr>
<td>Panniculitis, mononeuritis, urticaria pigmentosa, Sweet's syndrome, idiopathic thrombocytopenic purpura, polymyositis, hemolytic anemia, uveitis</td>
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<tr>
<td>Autoimmune polyendocrinopathy-candidiasis ectodermal dystrophy syndrome also known as autoimmune polyendocrinopathy syndrome-type 1</td>
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RECOMMENDATIONS 1-6

- AIH should be considered in any patient with acute or chronic hepatitis, particularly in the presence of high IgG levels, as it has a global distribution at any age in both sexes and in all ethnic groups (II-2)
- Precise and early diagnosis is mandatory, as untreated AIH carries high morbidity and mortality rates (I)
- AIH patients can be considered for screening for concurrent autoimmune diseases, especially autoimmune thyroiditis, since AIH is associated with a reduced quality of life and a broad variety of other autoimmune-mediated conditions (II-2)
- Cirrhosis at diagnosis should be suspected in AIH, as almost 33% of adults and 50% of children with AIH are first diagnosed at the stage of cirrhosis, indicating that they have had subclinical disease for a long time (II-2)
- Acute AIH can be diagnosed presenting as one of the following two clinical forms:
  - acute worsening of previously undiagnosed or misdiagnosed AIH or
  - real (original) acute onset of AIH without chronic lesions on liver histology (II-2)
- AIH can be classified into two types: AIH-1, ANA, SMA and/or anti-SLA/LP positive; and AIH-2, anti-LKM1, anti-LKM3 and/or anti-LC1 positive. Apart from differences in circulating autoantibodies, other differences in the clinical substrate have become apparent that may be helpful to clinicians (II-2)

Specific features

AIH may be first diagnosed during pregnancy or more frequently after delivery (Table 4). Postpartum exacerbations may also occur in AIH patients whose condition improved during pregnancy [37-39]. Interestingly, immunosuppression has probably enabled the occurrence of pregnancy in young females with amenorrhea at presentation due to AIH.

AIH may also develop after the administration of many drugs, supplements and/or herbs, with nitrofurantoin and minocycline being the best documented among diverse cases (Table 4) [40-44]. Drug-induced AIH is a complex and challenging condition characterized clinically and histologically by different phenotypes across the disease spectrum [40,45]. Therefore, the differentiation between drug-induced liver
injury (DILI) and DILI-induced AIH is often difficult [46]. In such cases, the patients’ history is important, as in one third of patients with DILI the clinical features can be associated with hypersensitivity manifestations, such as fever, rash and eosinophilia [47,48]. The follow up can also help in differential diagnosis, as steroid treatment can be discontinued without

Table 4 Specific characteristics of autoimmune hepatitis (AIH)

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>Description</th>
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<tr>
<td><strong>Special conditions</strong></td>
<td>Development of AIH in pregnant women or more frequently in the postpartum period is a rare event but does occur; the disease lessens during pregnancy but postpartum worsening is common; maternal and pregnancy outcomes are similar to those of the general population. AIH development after liver transplantation for conditions other than AIH (de novo AIH). Some AIH patients have PBC or PSC characteristics (AIH-PBC or AIH-PSC variants); in case of cholestatic findings, AMA investigation and cholangiography (particularly in children - autoimmune sclerosing cholangitis) are advised; “Paris criteria” for AIH-PBC variant: presence of at least 2/3 key criteria of each disease; for PBC: 1) ALP ≥2×ULN or γ-GT ≥2×ULN, 2) AMA detection, 3) liver biopsy showing florid bile duct lesions; for AIH: 1) ALT ≥2×ULN, 2) IgG ≥2×ULN or SMA detection, 3) liver biopsy showing moderate or severe perportal or periseptal lymphocytic piecemeal necrosis; IAIHG scoring systems should not be used to define such patients.</td>
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<tr>
<td><strong>Specific characteristics</strong></td>
<td>AIH development after use of drugs, supplements or herbs with nitrofurantoin and minocycline implicated in most cases; other drugs include: oxyphenisatin, ornidazole, methyldopa, diclofenac, interferon-α, atorvastatin, liraglutide, highly active antiretroviral treatment for human immunodeficiency virus and biologics including TNFa blockade agents; often very difficult to differentiate from DILI. AIH development after viral infections (e.g., Epstein-Barr, cytomegalovirus) including HCV; AIH should be strongly taken into account in cases with previous documented viral infections followed by unidentified and prolonged elevation of aminotransferases.</td>
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</table>

*PBC, primary biliary cholangitis; PSC, primary sclerosing cholangitis; AMA, antimitochondrial antibodies; ALP, alkaline phosphatase; γ-GT, γ-glutamyl-transpeptidase; ALT, alanine aminotransferase; IgG, immunoglobulin G, SMA, smooth muscle antibodies; IAIHG, International autoimmune hepatitis group; TNFa, tumor necrosis factor alpha; DILI, drug induced liver injury; HCV, hepatitis C virus; ULN, upper limit of normal*

![Differential diagnostic algorithm between AIH and DILI](image)

Figure 1 Proposed algorithm for the differential diagnosis between autoimmune hepatitis (AIH) and drug-induced liver injury (DILI) in an index case with biochemical hepatitis, positive liver autoimmune serology and hepatitis on liver histology, irrespective of the presence or absence of high IgG levels.

AST, aspartate aminotransferase; ALT, alanine aminotransferase; IgG, immunoglobulin G
relapse in DILI, in contrast to the almost universal relapse after stopping a few-month course of immunosuppressive therapy in the “true” DILI-induced AIH (Fig. 1).

AIH development has also been observed after viral infections [7,8,49,50], including hepatitis C virus (HCV) treated with interferon-α (IFNa) [51], or after liver transplantation for other liver diseases in adults and children (Table 4). This condition has been called de novo AIH and its early recognition seems helpful for avoiding another liver transplantation and improving patients’ long-term survival [52-54].

RECOMMENDATION 7
• AIH should be considered in the appropriate clinical and laboratory setting after use of drugs, supplements or herbs; viral infections; liver transplantation (de novo AIH); and rarely during pregnancy or after delivery (II-3)

Complications

As in other chronic liver diseases, cirrhosis and its consequences, including portal hypertension and hepatocellular carcinoma (HCC), may occur. Therefore, surveillance with ultrasonography every 6 months seems rational in patients with cirrhosis. HCC, however, is developed at significantly lower rates in AIH-associated cirrhosis compared with other causes (Table 1) [14,15,55-58].

RECOMMENDATION 8
• Patients with AIH-related cirrhosis should undergo ultrasonography every 6 months for early HCC detection, as in cirrhosis of other etiologies (II-2)

AIH variants

Some patients present, either concurrently or consecutively, with features of primary biliary cholangitis (PBC) or primary sclerosing cholangitis (PSC), while some PBC or PSC patients may also show AIH characteristics (Table 4) [59]. However, the previous term “overlap” used for years for these entities strongly suggests the simultaneous presence of two distinct diseases, which of course is not the case for many of these patients. Therefore, the term “variant” has recently been proposed by the European Association for the Study of the Liver (EASL), thought to be more precise for these conditions [60].

Worldwide accepted criteria defining these conditions are lacking and hence their diagnosis is usually difficult. Regarding the AIH-PBC variant, its prevalence is about 10% in adults with PBC or AIH [61]. The “Paris criteria” are even nowadays the most often used in everyday clinical practice (Table 4) [61]. Unlike several uncertainties around this issue, the recent EASL guidelines on PBC diagnosis and management recommend that liver histology is mandatory in evaluating patients with this variant of AIH [62]. In particular, liver biopsy seems crucial because of potential therapeutic implications in the PBC cases who do not respond to ursodeoxycholic acid (UDCA), having also disproportionate elevations in alanine aminotransferase (ALT) and/or IgG [62].

The AIH-PSC variant has been reported in 7-14% of mainly young patients with autoimmune liver diseases [59,63-65]. The criteria for AIH-PSC diagnosis are even less well defined than those used in AIH-PBC patients. In routine clinical practice, AIH-PSC diagnosis is based on typical cholangiographic or histological characteristics of PSC, along with AIH characteristics [66]. Interestingly, a specific and unique variant has been reported in 50% of children with AIH, characterized by both AIH and sclerosing cholangitis features; therefore, the term “autoimmune sclerosing cholangitis” was proposed [63]. Magnetic resonance cholangiopancreatography (MRCP) is advised for all children with an initial AIH diagnosis (Table 4) [63,64]. This entity is exceptionally rare in adults with AIH and thus MRCP screening is not justified [67].

STATEMENT 2
• Coexistence of AIH features and cholestatic diseases can be observed at both diagnosis and follow up, but their diagnosis may be problematic because of the lack of internationally accepted criteria (II-2)

RECOMMENDATIONS 9-11
• All children with AIH should undergo at least MRCP to exclude autoimmune sclerosing cholangitis (II-2)
• Adult AIH patients should be considered for MRCP only when cholestatic laboratory manifestations are present (II-3)
• AIH patients showing cholestatic features should be tested for PBC (II-2)

Laboratory investigation

Biochemistry

Bilirubin and aminotransferase levels vary from just above the upper limits of normal (ULN) to very high levels [4,7,8,18,19]. Alkaline phosphatase (ALP) is usually normal or moderately elevated, while γ-glutamyl-transpeptidase (γ-GT) can increase variously [23,34]. Spontaneous normalization of aminotransferases and γ-GT can be observed, although there is usually evidence of continuing inflammatory activity at the histological level. This phenomenon may result
in delay and/or underestimation of AIH diagnosis, as the next AIH hit can be pronounced after many months or years or may even be absolutely asymptomatic, explaining at least partially the presence of cirrhosis in one third of patients at diagnosis. Irrespective of the presence of cirrhosis, the majority of patients have high serum γ-globulins or IgG, a distinctive feature of AIH [19,23]. However, 25-40% of patients with acute AIH have normal IgG [29,68]. It should be noted that the range of “normal” IgG is ample, as it is impractical to have the “real normal ranges” of the respective population where an index case resides.

Liver autoimmune serology

Conventional antibodies

Autoantibodies are the hallmark for AIH diagnosis. Indirect immunofluorescence (IIF), preferably on freshly frozen rodent substrates including kidney, liver and stomach, is the technique of choice for routine screening (Fig. 2) [60]. Significant titers are ≥1:40 in adults, while ≥1:20 for ANA or SMA and ≥1:10 for anti-LKM1 are supportive of AIH diagnosis in children in association with other laboratory and clinical findings [19,60,69]. Other assays, such as ELISA or immunoblotting, are available for anti-LKM1, anti-LKM3, anti-LC1 and anti-SLA/LP testing [7,8,70].

ANA and SMA are not disease-specific and show a wide range of heterogeneity in the IIF pattern on HEP2 cells, together with varying titers [7,8,60]. However, no single ANA staining pattern on HEP2 cells is pathognomonic of AIH or seems to have any clinical and diagnostic implication in everyday practice; for these reasons their use in initial testing is not recommended. Since SMA react to cytoskeletal elements, including F-actin (the major autoantigen of SMA), testing for anti-F-actin antibodies can also be performed by ELISA [7,8,71,72]. However, IIF appears superior compared with ELISA, as testing only by ELISA may result in missing almost 20% of AIH cases [7,8,71-73]. Anti-LKM1/3 and/or anti-LC1 often coexist in AIH-2 but are not disease-specific, as they may be detected in 5-10% of HCV or hepatitis D virus (HDV) infections [7,8,23,51,70-72,74-76]. The major target-autoantigens of anti-LKM1 and anti-LKM3, first described in about 13% of HDV infections, are the cytochrome P4502D6 and family-1 of UDP-glucuronosyltransferases, respectively—although the antigenic sites differ between AIH and HCV or HDV infections; for anti-LC1 the major target-autoantigen is the formiminotransferase cyclodeaminase enzyme [7,8,60,70,72]. Repeated testing is advised in initially seronegative individuals, as conventional autoantibody titers vary during the course of AIH and may be detected later [18,70-72]. Autoantibody titers do not need to be monitored in adults, but should be in children, where they are considered as markers of disease activity [69]. Notably, the previous recommendation of the IAIHG for anti-LKM screening before starting IFNa-based therapies in HCV infections, as IFNa may sometimes unmask or provoke autoimmune liver reactivity and even original AIH, seems not to be rational in the new era of direct acting antivirals (DAAs) [18,51,65,78].

![Proposed algorithm for routine autoantibody testing in cases with a suspicion of autoimmune hepatitis](image-url)

*Determine also IgG serum levels

**Figure 2** Proposed algorithm for routine autoantibody testing in cases with a suspicion of autoimmune hepatitis

Anti-SLA/LP, antibodies against soluble liver antigens/liver pancreas; ELISA, enzyme linked immunosorbent assay; Abs, autoantibodies; ANA, antinuclear antibodies; SMA, smooth muscle antibodies; anti-LKM1, anti-liver/kidney microsomal antibody type 1; anti-LC1, antibodies against liver cytosol type-1 antigen; pANCA/ANNA, perinuclear anti-neutrophil cytoplasmic antibodies/anti-nuclear neutrophil antibodies; anti-LKM3, anti-liver/kidney microsomal antibody type 3; AIH, autoimmune hepatitis
**Non-conventional antibodies**

Anti-SLA/LP has greater diagnostic value, being the only AIH-specific autoantibody. However, it is only detected in up to a third of patients, usually in strong association with anti-Ro52 antibodies (concordance in 77-98%) [79-82]. The target-autoantigen is a synthase (S) converting O-phosphoseryl-tRNA (Sep) to selenocysteinyl-tRNA (Sec); it is therefore labeled as SepSec5, which in turn resulted in the development of reliable molecular-based assays for its detection [83,84]. Anti-SLA/LP has been associated with more aggressive disease [85], although these findings were not confirmed in recent studies [79,82].

Perinuclear pattern of anti-neutrophil cytoplasmic antibodies (pANCA), also referred to as perinuclear anti-neutrophil nuclear antibodies (p-ANNA), are detected (significant titer ≥1:20) by IIF frequently in AIH-1 patients and may act as an additional pointer towards diagnosis [18,69-72]. Anti-mitochondrial antibodies (AMA), strongly specific for PBC [86], can be detected in 8-12% of AIH patients without, however, any other evidence of chronic cholestatic disease [87-92].

Antibodies against α-actinin, which belongs to the F-actin cross-linking proteins, have been detected in autoimmune diseases such as systemic lupus erythematosus and AIH-1 [93,94]. These antibodies seem to carry particular clinical significance, as they characterize a subgroup of more severe form of AIH, specifically in association with anti-α-fil, while they might be used as predictors of treatment response [95-97]. Antibody against the asialoglycoprotein receptor is another frequently detected autoantibody, which can support diagnosis if patients have tested negative for other autoantibodies [7,8,70-72]. However, its specificity seems low and therefore its routine determination is not recommended [60].

**Liver histology**

The presence of hepatitis is a prerequisite for AIH diagnosis [4,6,10,18,19,60]. Ideally, liver biopsy should be performed before the initiation of treatment, as disease necroinflammatory activity and severity are not always in parallel with biochemistry [6,10,18,19]. Therefore, a pretreatment liver biopsy can provide information on prognosis but also on optimal AIH management; for instance, the potential presence of cirrhosis may influence the choice and dose of immunosuppression and suggest regular screening for complications. The typical lesions consist of interface hepatitis with portal lymphocytic/lymphoplasmacytic cells extending into lobule, hepatocyte rosetting and emperipolesis (etymology from the Greek language), which refers to the intracytoplasmic localization of one cell, usually a lymphocyte, within hepatocytes [19]. It should be noted however, that the abovementioned findings are not pathognomonic for AIH, while the absence of plasma cells observed in almost one third of AIH cases does not rule out the diagnosis [98].

Biopsies performed early in the acute disease show several signs of severe inflammatory activity, such as panlobular hepatitis with parenchymal collapse, presence of portal lymphoid follicles, inflammatory infiltrates enriched by plasma cells, central perivenulitis and pericentral, bridging or massive necrosis resembling those observed in acute DILI [18,28,68,99]. Inflammatory lymphocytic infiltrates of bile ducts have also been reported in approximately 10% of patients—without, however, any other clear manifestation of PBC [100]. At presentation, various fibrosis stages may be seen. Quantitative evaluation of inflammatory activity using the hepatitis activity index (HAI) score seems helpful during therapy and follow up.

Concerning the noninvasive methods, and in particular liver elastography, a recent study established that repeat transient elastography measurement is a reliable tool for AIH monitoring [101]. At present, however, the general belief is that the noninvasive tests are not able to replace liver biopsy at diagnosis and before treatment discontinuation [102].

**Differential diagnosis and diagnostic scores**

The differential diagnosis includes almost all causes of acute and chronic liver diseases as well as celiac disease [8,72,103] (Table 2). In 1999, the IAIHG published a score for AIH...
diagnosis [18] (Table 5). However, it proved quite complex for everyday use, while it was rather unable to safely distinguish AIH from cholestatic syndromes or AIH variants [65,104,105]. In 2008, the IAIHG proposed a simplified score for daily routine clinical practice, which is user-friendly, as it is based on autoantibody detection, IgG, liver histology, and seronegativity for viral hepatitis markers [19] (Table 6). This newer score seems to bear lower sensitivity (95% vs. 100%) but higher specificity and accuracy compared with the original revised score [106-108]. In general, however, physicians should keep in mind that any score should be used only as an aid to AIH diagnosis [109]. This is true, for example, in acute or fulminant AIH cases, AIH variants, children with AIH and DILI cases resembling AIH, as diagnosis by using the abovementioned diagnostic scores may be missed in such cases [28,29,68,99,108,110-112]. In particular, because of poor validation of the scores in the pediatric population, different scores have been suggested for child patients [69].

**Management**

All AIH patients who have active disease, even those with advanced fibrosis or cirrhosis, should receive immunosuppression in an attempt to achieve complete remission and to prevent the progression of liver disease through either maintenance therapy or a sustained remission following treatment withdrawal [6-8,10,33,60,98,113].

**Indications for treatment (Fig. 3)**

Untreated patients with moderate to severe disease (confluent necrosis on biopsy, AST/ALT >5×ULN and IgG >2×ULN) have a poor prognosis, while immunosuppression can improve symptoms and liver biochemistry, including IgG, leading to prolonged survival [114,115]. Spontaneous resolution of AIH may occur and treatment can be withheld. However, because of the fluctuating and unpredictable disease
Azathioprine is then gradually increased according to the response or its potential toxicity up to 1-2 mg/kg/day. Azathioprine alone should not be used as induction therapy as it has been associated with high mortality rates [6-10,60,118].

The primary aim should be the achievement of complete clinical and biochemical remission at the lowest corticosteroid dose, or even complete withdrawal using a rapid manner of corticosteroid tapering (Table 7) in order to avoid corticosteroid-dependent disease and development of several significant side-effects. Azathioprine should be used with caution in pregnancy, malignancies, cytopenias and established thiopurine methyltransferase (TPMT) deficiency.

Many patients (15-25%) develop side-effects or are intolerant to corticosteroid therapy alone, or in combination with azathioprine [121]. In addition, measurement of TPMT activity or genotyping, along with determination azathioprine of metabolite, are time consuming processes and are not widely available, while they neither provide convincing proof of avoidance of toxicity nor predict the treatment response [122-124]. On the other hand, in a review of 11 randomized controlled trials published from 1950-2009, which included 578 AIH patients (363 treatment-naïve), Lamers et al [115] reported a mean 43% remission rate after prednisolone with or without azathioprine treatment, much lower than those reported in the current literature (approximately 65-80%) [125], suggesting that this kind of treatment is far from ideal. In parallel, a large multicenter study in The Netherlands showed that AIH relapse is almost ubiquitous after treatment withdrawal, even though the patients were in long-term remission (>2 years), further supporting concerns regarding the conventional treatment’s lack of long-term efficacy [126].

Because of these concerns, mycophenolate mofetil (MMF) in combination with prednisolone, or budesonide in combination with azathioprine have been used as first-line induction treatment [13,34,127-134]. MMF blocks purine synthesis, inhibits DNA synthesis and has a selective antiproliferative effect on B- and T-cells. MMF has a 5-fold potent inhibitory effect on the type-II isozyme of inosine-5’-monophosphate dehydrogenase, which depletes guanosine nucleotide specifically in activated T- and B-cells without affecting the type-I isozyme and thus results in more powerful and selective immunosuppression with few side-effects [135]. In addition, in patients and experimental animal models, it has been shown that MMF-based immunosuppression

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**RECOMMENDATIONS 20-22**

- AIH therapy should aim to achieve complete biochemical and histological remission in an attempt to prevent potential disease progression (II-2)
- All patients having active disease, even those with advanced fibrosis or cirrhosis, should be treated (I)
- Treatment can be withheld in patients with spontaneous remission for whom a close long-term follow up is advised in order not to miss an AIH exacerbation (III)
could restore the regulatory T-cells [136-138], considered very important in AIH pathogenesis [139-141].

In this context, two uncontrolled real-world prospective studies from Greece [34,128], including the largest numbers of treatment-naïve AIH patients ever published (n=59 and n=109), showed that MMF at a dose of 1.5-2 g/day in two divided doses was safe (discontinuation in 3%) and effective as first-line treatment to induce and maintain response with a rapid steroid sparing effect (Fig. 4). In fact, initial complete response was achieved in 88% [34] and 93.6% [128] in a median time of less than 3 months (significantly shorter compared with conventional azathioprine schedules [23,142]), while on treatment complete remission was achieved in 59.3% [34] and 71.6% [128] of patients, compared with the 26% reported by Muratori et al [23,143] and 43% by Lamers et al [115]. Most importantly, a recent study by Zachou et al showed the highest rates of remission maintenance off treatment (72-75%) ever published, for at least a median of 30 months, accompanied by significant improvement in necroinflammatory activity and stable and/or improved fibrosis at second liver biopsy [128,129]. Similar findings independent of the presence of cirrhosis were reported in another retrospective study (84% response rate) in 29 AIH patients [127].

Budesonide has also been used effectively in a randomized study (9 mg/day) in combination with azathioprine in non-cirrhotic AIH patients [130]. Biochemical remission (IgG normalization was not included in the response criteria) without the typical steroid-induced side-effects was found more frequently in budesonide-treated patients compared with the prednisone-treated group (47% vs. 18%), and side-effects were fewer (28% vs. 53%) [130]. However, response rates and side-effects in the control arm were surprisingly lower and higher, respectively, than in earlier studies, presumably because of the initial fixed-dose and fixed-dose reduction schedule in the prednisone group, whereas in the budesonide group the drug was given at a high dose until a response was

---

**Table 7** Suggested treatment schedule for adults with recent diagnosis of autoimmune hepatitis (e.g., 70 kg; adapted from [60,115])

<table>
<thead>
<tr>
<th>Week</th>
<th>Prednisolone (mg/day)</th>
<th>Azathioprine (mg/day)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>70 (=1mg/kg)</td>
<td>-</td>
</tr>
<tr>
<td>2</td>
<td>60</td>
<td>-</td>
</tr>
<tr>
<td>3</td>
<td>50</td>
<td>50</td>
</tr>
<tr>
<td>4</td>
<td>40</td>
<td>50</td>
</tr>
<tr>
<td>5</td>
<td>30</td>
<td>100**</td>
</tr>
<tr>
<td>6</td>
<td>25</td>
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</tr>
<tr>
<td>7</td>
<td>20</td>
<td>100</td>
</tr>
<tr>
<td>8 + 9</td>
<td>15</td>
<td>100</td>
</tr>
<tr>
<td>10 + 11</td>
<td>12.5</td>
<td>100</td>
</tr>
<tr>
<td>From 12</td>
<td>10*</td>
<td>100</td>
</tr>
</tbody>
</table>

A lower prednisolone dose can be used initially in mild disease or in early relapses during corticosteroids withdrawal (e.g., 0.5-0.7 mg/kg/day). The tapering schedule of corticosteroids should be individualized according to the rapidity of the response and the development of side-effects. *If transaminases are normalized, prednisolone could be reduced to 7.5 mg/day and after 3 months to 5 mg/day, aiming at complete withdrawal after 6-8 months (or after 3-4 months at 2.5 mg/day) according to a personalized assessment of the patient’s risk and response. **Azathioprine dose according to body weight (1-2 mg/kg)
achieved, thus suggesting a therapeutic bias. In addition, data on the histologic response in budesonide-treated patients are still lacking, whereas other small studies or case reports have shown either failure or exacerbation of AIH during budesonide monotherapy, making the advantages of a more expensive regimen as first-line therapy in AIH uncertain [144-146].

**Maintenance therapy**

Patients with mild necroinflammatory activity at initial biopsy, intolerant to azathioprine and have achieved complete biochemical response, can receive prednisolone monotherapy at the lowest dose that can maintain remission. In all other patients, the aim should be prednisolone off

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**Figure 4** Suggested therapeutic algorithm for prednisolone in combination with mycophenolate mofetil (MMF) in treatment-naive AIH patients.

*In patients with risk factors (e.g., anti-LKM, anti-LC1, anti-α-actinin, anti-SLA/LP, cirrhosis at diagnosis) the tapering schedule could be applied every 3 weeks. **In relapses (↑AST, ALT ± IgG) prednisolone should be increased to the dose of initial complete response and then tapered, either by increasing the time interval twofold or by decreasing the dose of prednisolone tapering by half at the same time. ***In relapses after corticosteroid withdrawal restart prednisolone at the dose of initial complete response and taper according to **. MMF is given in two divided doses AIH, autoimmune hepatitis; anti-LKM, anti-liver/kidney microsomal antibody; anti-LC1, antibodies against liver cytosol type-1 antigen; anti-SLA/LP, antibodies against soluble liver antigens/liver pancreas; AST, aspartate aminotransferase; ALT, alanine aminotransferase; IgG, immunoglobulin G

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**STATEMENTS 8-9**

- The long-term efficacy of the conventional treatment is uncertain, as 15-25% of patients develop intolerance or side-effects (II-2)
- Long-term biochemical and histological data on budesonide safety and efficacy are lacking (I)

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**RECOMMENDATIONS 23-27**

- Induction treatment for AIH should be individualized and response-guided (II-2)
- Prednisolone at one oral dose of 0.5-1 mg/kg/day in the morning, combined with azathioprine at an initial morning dose of 50 mg/day usually after 2 weeks, if bilirubin is <6 mg/dl, should be the first-line therapy for AIH (I)
- Azathioprine should then be increased up to 1-2 mg/kg/day (maintenance dose) (II-2)
- In specialized AIH centers, prednisolone (0.5-1 mg/kg/day), combined with MMF (2 g/day) from the beginning of treatment if bilirubin is <6 mg/dL, may be used as first-line therapy, since real-world studies showed high rates of remission maintenance off treatment accompanied by improved liver histology (II-3)
- In non-cirrhotic AIH patients, budesonide (9 mg/day) in combination with azathioprine may be used as induction treatment in those with serious comorbidities that might be exacerbated by conventional corticosteroid therapy (II-2)
monotherapy with individualized adjusted azathioprine (2 mg/kg/day), or alternatively MMF doses (1.5-2 g/day) [7,34,115,119,147]. The total duration of immunosuppression should be at least 3 years, with at least the last 2 years having persistent complete biochemical response (normalization of transaminases and IgG). In patients who have received adequate induction and maintenance immunosuppression, but who have not achieved biochemical or histological remission, immunosuppression should not be stopped, as relapse will occur almost universally.

Relapse of the disease

Relapse of AIH is defined as the reappearance of clinical or laboratory markers of active disease (ALT ≥2-3×ULN and/or an increase in IgG, usually preceded by ALT elevation) [60] after achievement of complete remission during the induction therapy, or during maintenance therapy and/or after complete discontinuation of treatment. In this setting, liver biopsy is usually not recommended. Relapses are particularly frequent in patients who suffer from multiple relapses develop more side-effects and are likely to have worse outcomes [149,153,154].

Primary treatment endpoints and discontinuation of therapy

The ideal treatment endpoints are a complete clinical, biochemical and histological response with prolonged off treatment remission [7,60,118,143,150,152,153]. In real-life, however, these endpoints are achieved in a minority of patients who discontinue therapy based on prednisolone alone or in combination with azathioprine [115,143], while data on the same issue in budesonide-treated patients are still lacking. In contrast, real-world studies showed high rates of maintained off-treatment remission accompanied by histological improvement in patients treated with prednisolone in combination with MMF as first-line therapy [34,128,129].

Treatment withdrawal can be suggested only in those patients who have achieved continuous complete biochemical remission for at least the last 2 years of treatment, and especially in those with ALT below half the ULN along with IgG <1200 mg/dL [155]. In these patients, liver biopsy before treatment withdrawal is advisable [156]. However, complete treatment withdrawal is almost impossible in difficult-to-treat patients (see below), including cirrhotics and patients with AIH-2.

Follow-up before and during treatment

Patients under conventional or budesonide/azathioprine schedules should be followed with baseline and weekly tests for transaminases, albumin, prothrombin time, fasting glucose and full blood count during the first month. In patients

<table>
<thead>
<tr>
<th>RECOMMENDATIONS 28-32</th>
</tr>
</thead>
<tbody>
<tr>
<td>- The optimal maintenance treatment should be corticosteroid-free monotherapy with azathioprine, or alternatively MMF (II-2)</td>
</tr>
<tr>
<td>- Low-dose long-term prednisolone monotherapy may be used to maintain remission only in patients with mild disease and azathioprine intolerance who have achieved complete response after induction therapy (II-2)</td>
</tr>
<tr>
<td>- Maintenance therapy should be adjusted to a dosage that can maintain persistent biochemical response (normalization of AST, ALT and IgG) (II-2)</td>
</tr>
<tr>
<td>- Immunosuppression should be given for a total of at least 3 years, and for at least 2 years after the achievement of complete biochemical response (II-2)</td>
</tr>
<tr>
<td>- Maintenance therapy should not be withdrawn without a complete biochemical or histological response (HAI&gt;3) (II-2)</td>
</tr>
</tbody>
</table>
under prednisolone/MMF, the above data could be obtained at 2-3-month intervals, along with IgG determination. After corticosteroid tapering, monitoring intervals can be every 3 months and every 3-6 months during maintenance therapy.

Baseline hepatitis B (HBV) and A serology is recommended before treatment, along with the respective vaccination for those not indicating previous vaccination or virus exposure. For HBV surface antigen-positive (HBsAg) patients, preemptive therapy with either entecavir or tenofovir is strongly recommended according to the EASL and the Hellenic Center for Disease Control and Prevention (HCDCP) CPG [157,158]. Vaccination against Streptococcus pneumoniae, along with yearly influenza vaccination, should also be given to all patients.

Dual energy X-ray absorptiometry scanning assessment before treatment initiation and at 1-5-years intervals seems rational, but there are no specific data to support this assertion. Likewise, though no specific data exist concerning the use of calcium and vitamin D in patients under immunosuppression, such treatment seems reasonable, as in other diseases under corticosteroids.

**Specific conditions and difficult-to-treat patients**

**Non-responders**

Non-response includes a partial or incomplete response and treatment failure (null response), associated or not with an acute/severe form of the disease.

**Partial response**

A partial response means that there is some improvement in clinical, biochemical and histological parameters but without reaching complete remission despite treatment adherence. These patients have abnormal transaminases (usually below 2-3×ULN) or necroinflammatory activity on histology [6,7,10,60].

In partial responders under conventional treatment, an option is to increase azathioprine to 2 mg/kg/day in combination with prednisolone (5-10 mg/kg/day), followed by a repeat liver biopsy after 12-24 months [6,9,10,60]. In patients with a partial response after adequate treatment with budesonide-based schedules, a change to prednisolone (>20 mg/day initially) could be considered [146]. If complete biochemical and histological remission is not achieved, the aim should be either the lowest achievable biochemical activity in parallel with minimum side-effects, or the administration of alternative second-line therapeutic agents [159].

**Treatment failure**

Primary complete biochemical remission after adequate treatment initiation is the rule (90-95% of patients); therefore, reassessment of diagnosis and treatment adherence should be considered in non-responders. Non-response is not well-defined, but usually the absence of a transaminase decline of at least 25% from baseline after 2-3 weeks should be considered as non-response. It is also important to remember that other conditions may develop concurrently during the AIH course, such as viral infections or DILI, which if unrecognized, could be mistakenly regarded as a null response or flares [71,72,160,161].

Treatment failure can be seen either in patients with an acute/severe or even fulminant disease form, or in those without such intense severity. Data on patients with acute/severe presentation are scarce, consisting mostly of real-life non-randomized studies with a small number of patients and varying, mostly arbitrary entry criteria, because the precise definition of this form of AIH is still missing [112,162-171]. Therefore, the role and timing of corticosteroids in modifying the outcome of acute/severe AIH remains unclear, as it is ambiguous whether such patients should be given a corticosteroids trial, a priority listing for liver transplantation, or both. Potts and Verma recently reviewed five retrospective studies, each with a small number of patients, including in total 85 patients with AIH-related acute liver failure [170].
nine patients (89.2%) received immunosuppression, mostly oral corticosteroids, and had remission rates of 8.3-50%, while 43.5% underwent or were listed for liver transplantation and 33% died. The largest recent studies in acute/severe AIH come from France (n=104) and Greece (n=42) [112,171]. In the multicenter retrospective French study, an overall survival rate of 90% was reported (median follow up: 2.3 years), although early liver transplantation was required in one third of patients. The beneficial effect of corticosteroids was observed in 66% of patients, mainly in those with low international normalized ratio (INR) at baseline and improvement in liver function during the first week of treatment [171]. In the Greek study, high-dose intravenous corticosteroids were given (prednisolone 1.5 mg/kg/day) in the early stages of acute/severe AIH, defined as an acute presentation with any sign of hepatic encephalopathy and transaminases >10×ULN, INR ≥1.5 and bilirubin ≥4 mg/dL at any time during the acute course [112]. This management appeared to prevent disease deterioration without increasing morbidity and mortality (long-term overall survival without transplantation: 95.2%; median follow up: 5.3 years).

Conclusively, despite the low level of evidence, the available data indicate that all patients with acute/severe AIH should be considered for a corticosteroid trial at the earliest opportunity (the sooner the better), using high doses of prednisolone (>1 mg/kg/day) intravenously [172]. Failure to improve within 7 days should lead to emergency listing for liver transplantation [167]. The prophylactic use of antibiotics/antifungals is not supported by the most recent studies [112,171], but they should be kept in mind and may be justified at an individualized basis.

The other form of treatment failure without intense severity is characterized by minimal, or even no improvement in the clinical and biochemical parameters after several weeks of standard regimen, despite confirmation of diagnosis and treatment adherence. In these patients, determination of active azathioprine metabolites, such as thioguanine nucleotides (TGN), could be helpful in revealing a lack of adherence to treatment or an altered azathioprine metabolism, although the therapeutic range of TGN levels is not precisely known in AIH [173]. As endorsed by EASL, AASLD and the British Society of Gastroenterology GPG [6,10,60], an increase of prednisolone for at least one month and azathioprine at the maximum dosage (1 mg/kg/day intravenously) could be tried, followed by management at expert centers in the case of non-response (II-3).

Non-adherence to treatment is an important problem during long-term follow up, particularly among children and adolescents under corticosteroids, resulting in frequent relapses or flares of AIH [174,175]. The management of non-compliance is difficult; therefore, a multidisciplinary approach is needed, including psychologists, social and youth workers, health carers, committed nurses, and pediatric and adult hepatologists in an attempt to re-motivate the young patient [176].

In addition, high rates of depression and anxiety have recently been reported in AIH patients [27] Thus, considerable attention is needed for those not appropriately followed for the psychiatry compartment, because this behavior could cause an increased frequency of non-adherence, leading to these patients erroneously being considered as non-responders and candidates for alternative therapies, with unexpected consequences [177].
Treatment adherence should be of utmost importance for all patients and especially for children, adolescents and young adults (II-2).

The transition from childhood to adult care should be based on a multidisciplinary approach by special transition services (II-3).

Considerable attention should be given to patients with anxiety or depression in order to be sure that the psychiatric follow up and treatment are appropriate before appraising these patients as non-responders to AIH therapy (III).

Intolerance to and side-effects of treatment

Corticosteroid side-effects are numerous and develop in up to 80% of patients receiving steroid monotherapy for more than 2 years, mainly at doses >15 mg/day. However, corticosteroid discontinuation because of severe adverse events is observed in 15%. Combination with azathioprine is associated with a much lower rate of side-effects [112,147]. In non-cirrhotic prednisolone responders who nevertheless develop side-effects, even though azathioprine has been increased to the highest dose, a switch to budesonide could be suggested.

Azathioprine side-effects occur in about 25% of AIH patients, accompanied by drug withdrawal in approximately 10-15%. In patients intolerant to or with side-effects of azathioprine (bone marrow suppression, nausea, vomiting, pancreatitis, etc.), a switch to MMF (2 g/day) with subsequent prednisolone tapering appears to be an excellent alternative [178-186]. Other alternatives are steroid monotherapy in patients with mild disease, budesonide, mercaptopurine or thioguanine, tacrolimus, cyclosporine, methotrexate, cyclophosphamide, allopurinol, and biologic regimens, including tumor necrosis factor α (TNFa) blockade agents and rituximab [187-194].

Non liver-related comorbidities and aging

Approximately one third of older AIH patients (>65 years) already have cirrhosis or advanced fibrosis at diagnosis, although they are more frequently asymptomatic at presentation compared to younger patients [25,27,33,195]. These patients achieve biochemical response more frequently and have a higher prevalence of concurrent autoimmune diseases compared with younger patients [25,196,197].

In asymptomatic elderly patients with mild disease, the decision about treatment initiation could be based on the presence and the severity of other comorbidities, such as uncontrolled diabetes mellitus, refractory arterial hypertension, established osteoporosis or a previous or current history of psychosis. In this context, it is better to choose a watch-and-wait strategy, but close follow up of these patients is strongly advised. Although there is no convincing data concerning the management of elderly AIH patients with mild disease and no severe comorbidities [33,116,197], it seems rational to initiate immunosuppression using a lower starting prednisolone dose (0.5 mg/kg/day), followed by a more rapid steroid de-escalation schedule in combination with azathioprine (1-2 mg/kg/day). Another attractive option for these patients without cirrhosis may be the use of budesonide (9 mg/day) plus azathioprine. In elderly patients with at least moderate necroinflammatory activity, treatment is recommended, but again the choice of steroid therapy and the tapering schedule should be considered carefully. The same is true for all AIH patients, irrespective of age, who have concurrent severe comorbidities.

AIH variants

Although controlled trials are lacking, the recent EASL CPG for PBC have recommended adding immunosuppression to UDCA in previously well-established PBC cases if at least moderate interface hepatitis is present on liver biopsy [62,198-200]. Notably, these patients appear to respond to lower immunosuppression dosages and maintain remission after treatment withdrawal at higher rates than patients with AIH alone [198,199]. It is uncertain whether AIH patients who develop PBC features will benefit from UDCA administration, but this addition seems logical under real-life conditions, particularly in young patients, because of the potential long-term benefit of UDCA, which may subsequently...
In patients with the AIH-PSC variant, the addition of MMF should be stopped at least 3 months prior to conception. A combination treatment consisting of immunosuppression and UDCA has been suggested in response criteria have recently been published for autoimmune sclerosing cholangitis, the pediatric form of the variant [69]. However, although biochemical and histological parameters may improve, the biliary tract lesions may progress and therefore the outcome of these patients seems worse compared to those with only AIH [63,64]. The combination of UDCA and immunosuppression has also been proposed by EASL CPG for the adults with AIH-PSC variant [201], although the available but inadequate studies show that the long-term prognosis appears worse than for either AIH cases without PSC characteristics or PSC alone [66,202,203].

RECOMMENDATIONS 57-58

- In patients with the AIH-PBC variant, a combination of immunosuppression with UDCA is recommended; alternatively, if AIH is the dominant compartment, immunosuppression only should be started and UDCA could be added if remission is not achieved (III)
- In patients with the AIH-PSC variant, the addition of UDCA to immunosuppressive treatment can be considered (III)

Management of AIH in pregnancy

Large series of pregnant AIH patients from the United Kingdom [37], Germany [204] and Sweden [205] have shown that the new or continued administration of azathioprine had no significant impact on the rate of live birth, termination or miscarriage, or on the gestational period. Interestingly, some unexplained adverse outcomes were found to be associated with anti-SLA/LP and anti-RO52 detection [204], while a higher Cesarean section rate was recorded in Sweden [205], but again without higher stillbirth or fetal malformation rates compared to controls. Taken together, these findings suggest that the continuation of azathioprine in females with well-established AIH after conception appears rational and justified.

As discussed previously, AIH may also present for the first time during pregnancy or more frequently after delivery [37-39]. These cases should be treated as discussed previously for non-pregnant patients. However, as the disease during pregnancy usually has lower activity, a minimal adjustment of immunosuppression (5-10 mg/day prednisolone ± 50-75 mg/day azathioprine) in previously diagnosed patients receiving treatment seems rational. Immunosuppressive treatment should then be increased after delivery to the previous dosages, in order to minimize the risk of flare during post-partum. Complications seem to be more frequent for those patients who have not achieved complete biochemical remission at least one year before conception, while cirrhotic patients carry a high risk of adverse outcomes. Although data regarding breastfeeding in AIH females under azathioprine are limited, the drug is considered safe, despite the fact that small amounts of its metabolite can be detected in breast milk [206].

RECOMMENDATIONS 59-62

- Females with AIH in remission should be advised that they have no contraindication for pregnancy or breastfeeding (II-2)
- Maintenance treatment with prednisolone with or without azathioprine should be continued in previously diagnosed females under therapy (II-2)
- Mild flares can be observed during the first trimester and more frequently especially in the post-partum period, requiring an increase in immunosuppression (II-2)
- MMF should be stopped at least 3 months prior to conception (either in females or males under treatment), as this drug is absolutely contraindicated in pregnancy (II-2)

Management of AIH after liver transplantation

Recurrence of AIH has been reported in 20-25% of patients [207] and is usually managed with the standard prednisolone treatment schedules in combination with azathioprine or MMF [208]. Non-responders in this setting could be aided by the use of sirolimus [209]. De novo AIH has been described in approximately 5% of subjects transplanted for reasons unrelated to AIH [53,207] and its management is identical to that suggested for recurrent AIH [210].

RECOMMENDATION 63

- AIH after liver transplantation, either recurrent or de novo, should be managed by the basic treatment principles of AIH (II-3)

Liver-related comorbidities

In countries with a moderate to high prevalence of HBV or HCV infections, the coexistence of AIH with chronic viral hepatitis is not impossible [211-213]. On the other hand, AIH patients can contract viral hepatitis. In these cases, AIH may be overlooked, as the absence of viral hepatitis markers is one of the 4 essential parameters of the simplified criteria for AIH diagnosis [19]. AIH in this situation seems to be more aggressive and carries a poorer prognosis compared to chronic viral hepatitis alone, as AIH could remain untreated in the long term because of its underdiagnosis/misdiagnosis [212]. In patients with chronic HBV infection and characteristics of AIH at diagnosis, HBV should be treated first according to the EASL and HCDCP CPG.
In AIH patients with concurrent chronic HBV or HCV at diagnosis, the viral infections should be treated first and AIH therapy could be offered after HBV suppression or HCV eradication if the necroinflammatory activity persists (III).

In AIH patients who contracted HBV or HCV infections during the AIH course, antiviral treatment with entecavir/tenofovir or DAAs, respectively, is recommended without stopping immunosuppression (III).

In HIV patients with AIH, treatment should be given on an individualized basis, as standard immunosuppression seems to be safe and effective but carries a risk of life-threatening infections (III).

In AIH patients with NAFLD/NASH, efforts should be made to ensure strict adherence to NAFLD/NASH guidelines and the use of the lowest effective dose of steroids for AIH, although the impact of NAFLD/NASH on treatment outcome and response in AIH is unknown (III).

Concluding remarks

AIH is characterized by genetic, clinical, laboratory, histological and serological heterogeneity and therefore, it might be underdiagnosed or unrecognized. Its diagnosis is based on the presence of polyclonal hypergammaglobulinemia, circulating autoantibodies, interface hepatitis on liver histology, absence of viral hepatitis and a favorable response to immunosuppression. In particular, autoantibodies detection and the interpretation of histological findings are considered the hallmarks for a timely diagnosis. Most of the affected patients are treated very efficiently with at least near normal life expectancy and quite good quality of life. However, many patients still have experience of remarkable morbidity because of delayed or missed diagnosis, drugs intolerance and side-effects, partial treatment response or flares, poor management and poor adherence. The establishment of major centers bearing special expertise on this disease diagnosis and control will be a governing factor in improving management of patients. Also, research on AIH etiology and its underlying pathogenetic mechanisms will be the clue to improve therapy of AIH. So far, most patients need long-life immunosuppressive treatment but patients want cure, not only remission of the disease. In an attempt to achieve curative treatment, we have to work hard together with basic scientists and follow developments in immunology and autoimmunity.

References

effectively than prednisone in a controlled trial of patients with autoimmune hepatitis. Gastroenterology 2010;139:1198-1206.


### Supplementary Table 1 Grading of recommendations

<table>
<thead>
<tr>
<th>Grade</th>
<th>Study Design</th>
</tr>
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<tbody>
<tr>
<td>I</td>
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</tr>
<tr>
<td>II-1</td>
<td>Controlled trials without randomization</td>
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<tr>
<td>II-2</td>
<td>Cohort or case-control analytic studies</td>
</tr>
<tr>
<td>II-3</td>
<td>Multiple time series, dramatic uncontrolled experiments</td>
</tr>
<tr>
<td>III</td>
<td>Opinions of respected authorities, descriptive epidemiology</td>
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*Adapted from: [1]*