Faculty of Health: Medicine, Dentistry and Human Sciences

Peninsula Medical School

2018-11

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http://hdl.handle.net/10026.1/18591

10.1111/apt.14968 Alimentary Pharmacology and Therapeutics Wiley

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Inequity of care provision and outcome disparity in autoimmune hepatitis in the United Kingdom

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Funding information

National Institute for Health Research, Grant/Award Number: BH140575 / PD00401; MRC Clinician Scientist Award, Grant/Award Number: G1002552

Summary

Background: Treatment paradigms in autoimmune hepatitis (AIH) have remained largely unchanged for decades. Studies report \leq 20% of patients have sub-optimal treatment response with most requiring long-term therapy.

Aim: The United Kingdom Autoimmune Hepatitis (UK-AIH) study was established to evaluate current treatment practice and outcomes, determine the unmet needs of patients, and develop and implement improved treatment approaches.

Methods: The United Kingdom Autoimmune Hepatitis study is a cross-sectional cohort study examining secondary care management of prevalent adult patients with a clinical diagnosis of autoimmune hepatitis. Enrolment began in March 2014. Prevalent cases were defined as having been diagnosed and treated for >1 year. Demographic data, biochemistry, treatment history and response, and care location were collected.

Results: In total, 1249 patients were recruited; 635 were cared for in transplant units and 614 in non-transplant centres (81% female with median age at diagnosis 50 years). Overall, 29 treatment regimens were reported and biochemical remission rate was 59%. Remission rates were significantly higher in transplant compared to non-transplant centres (62 vs 55%, P = 0.028). 55% have ongoing corticosteroid exposure; 9% are receiving prednisolone monotherapy. Those aged \leq 20 years at diagnosis were more likely to develop cirrhosis and place of care was associated with an aggressive disease phenotype.

Conclusions: There are significant discrepancies in the care received by patients with autoimmune hepatitis in the UK. A high proportion remains on corticosteroids and there is significant treatment variability. Patients receiving care in transplant centres were more likely to achieve and maintain remission. Overall poor remission rates suggest that there are significant unmet therapeutic needs for patients with autoimmune hepatitis.

JKD and LLW are equal first authors.

MAH and DEJJ are equal last authors.

The Handling Editor for this article was Professor Stephen Harrison, and it was accepted for publication after full peer-review.

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1 | INTRODUCTION

Autoimmune hepatitis (AIH) is a progressive inflammatory condition of the liver that may present in either acute or chronic forms.¹⁻³ If not effectively treated it can progress rapidly to acute liver failure or the development of cirrhosis. With a prevalence of approximately 17 per hundred thousand in Northern European populations the disease affects both adults and children.^{4,5} Treatment paradigms established in the 1970s and 1980s utilising corticosteroids and azathioprine to achieve and sustain remission have remained largely unchanged over several decades.¹ Although case series of patients treated in specialist centres suggest that disease remission (even using the current definition of normalised transaminase and immunoglobulin G [IgG] levels) can be achieved in up to 80% of patients,⁶ there is concern that real world disease outcomes in patients treated across the spectrum of health care settings may be substantially worse.⁷ This leaves patients at risk of progression to end-stage disease for which liver transplantation is the only effective therapy.⁸⁻¹⁰

International treatment guidelines have defined initial management and ongoing treatment models in AIH.⁸⁻¹⁰ All recommend corticosteroids in the form of prednis(ol)one or budesonide with azathioprine. The goals of treatment for patients are biochemical and histological remission, with effective control of symptoms, followed by long-term maintenance of the remission state. Ideally, this should be achieved with minimisation of the dose of corticosteroid with full withdrawal being the goal. Maintenance with azathioprine monotherapy where possible is advised in UK and European Guidelines to minimise corticosteroid side effects and their impact on quality of life. The majority of patients require long-term therapy to prevent relapse^{7,11} and increasing numbers of patients suffer with unpleasant side-effects, poorly controlled disease and a life-long immunosuppression burden.¹²⁻¹⁵

The evidence base for management of patients who are nonresponders to conventional immunosuppression is limited. Alternative immunotherapy, whilst recommended as second- and third-line treatment in patients intolerant of azathioprine, can be variable in efficacy and tolerability.¹⁶⁻²⁰ In AIH, the majority of data relating to treatment outcomes is derived from large referral centres.^{12,21} Even amongst these expert centres, significant differences exist in relation to approach to treatment.²²

In recent years, there has been a growing awareness of inadequacies in service provision for many common liver diseases and attention focused on the public health issues pertaining to the burden of liver disease in the UK.²³⁻²⁵ To date, however, little attention has been given to rarer liver diseases such as primary biliary cholangitis (PBC), primary sclerosing cholangitis (PSC) and AIH in these documents and the unmet needs and requirements of these patient groups, whilst known, have never been properly quantified.^{26,27}

Since the majority of data defining outcome and treatment of AIH are derived from tertiary centres which may not accurately reflect the full spectrum of care delivery we set out to derive a national cohort of patients representing multiple hospital practice settings in the UK. The intention of the United Kingdom Autoimmune Hepatitis (UK-AIH) consortium is to use this platform to define current "real world" practice in the management of AIH within the UK, and to develop, evaluate and implement improved approaches to treatment. The goals of this study are to evaluate current treatment practice and remission rates and determine the real-life unmet clinical needs of patients with AIH.

2 | METHODS

The UK-AIH platform is a UK-wide cross-sectional cohort developed to evaluate the management and outcome of adult patients with AIH in the UK and to facilitate the development, evaluation and implementation of improved therapy. A key aim is to determine the unmet needs of patients with AIH. The UK-AIH patient cohort is comprised of patients 16 years of age or older who carry a clinical diagnosis of AIH. Patients were enrolled from secondary and tertiary hospital settings from March 2014 to March 2017. The cohort described here was of prevalent patients (ie, diagnosis of AIH was typically several years before enrolment into the cohort).

Patient enrolment into this study was based on what individual clinicians considered an a-priori diagnosis of AIH. Since patients were prevalent and recruited based on the diagnosis made at clinical presentation (typically several years before study enrolment), no attempt was made to calculate the International Autoimmune Hepatitis Group (IAIHG)²⁸ or the simplified IAIHG diagnostic criteria from 2008.²⁹

Prevalent cases were identified from clinical records of enrolling hospitals. To be eligible, patients had to have carried a diagnosis of AIH for more than 1 year in their referring hospital. Demographic and clinical data, including risk factors for progressive liver disease, were collected on standardised data collection forms completed by the local managing clinicians. Current biochemical status including alanine aminotransferase (ALT), aspartate aminotransferase (AST), immunoglobulin G (IgG) levels, current therapy, past treatment history related to corticosteroid dose use in the 12 months prior to inclusion in the study, and treatment flares in the last 12 months were collected. Data in relation to diagnostic liver biopsy and disease progression, as defined by progression to cirrhosis during follow-up, were recorded.

Biochemical remission status at the time of study enrolment (rather than at the time of original diagnosis) was assessed using contemporaneous ALT and IgG values. Upper limits of normal of ALT and IgG were utilised for each recruiting centre. A disease flare was defined as a need to treat an increase in ALT level in the previous 12 months with a higher corticosteroid dose than their maintenance dose or through introduction of corticosteroid therapy in patients on corticosteroid-free maintenance.

The presence of cirrhosis at the time of diagnosis of disease was determined by findings on liver biopsy coupled with imaging criteria. Development of decompensated liver disease and need for liver AP&T Alimentary Pharmacology & Therapeutics -

transplantation following diagnosis were identified as surrogate markers for disease severity and treatment failure. In addition to the information gathered from clinicians, patients completed questionnaires in relation to the prevalence of additional autoimmune disorders, ethnicity, height and weight. A protocol amendment to the study in November 2015 allowed for additional data collection, comprising variables at diagnosis, including autoantibody titres, IgG levels and viral serology including hepatitis A, B, C, E viruses, Cytomegalovirus, and Epstein-Barr virus.

The protocol was approved by the National Health Service (NHS) Health Research Authority (IRAS ID: 144806, REC reference: 14/LO/ 0303) and was conducted in accordance with the International Council for Harmonisation (ICH) Good Clinical Practice (GCP) guidelines and the Declaration of Helsinki. Written informed consent regarding the use of data was obtained. Data were analysed using sPss version 22, GraphPad Prism 7 and sAs 9.4. Nonparametric data are presented as median and range. Continuous variables were described as median, minimum and maximum. Difference between proportions were analysed using the *Z* test. Logistic regression was performed to assess risk factors for cirrhosis development. All *P* values reported are two-sided, and *P* < 0.05 was considered statistically significant.

3 | RESULTS

3.1 Demographics and baseline features

One thousand two hundred and forty-nine patients were enrolled into the study cohort from 44 centres (seven liver transplant centres and 37 nontransplant centres). 635 patients were under the care of transplant units and 614 under nontransplant centres. The majority of patients were female (1006, 81%). Median age at diagnosis of AIH of enrolled patients was 50 years (range 2-86 years). Seventynine (6%) of the patients were diagnosed below the age of 16 with 65/79 (82%) being cared for in transplant units.

Patients managed in transplant centres were diagnosed at a younger age than those managed in nontransplant centres (median 42 years [range 2-86 years]) with 116 (18%) of patients diagnosed at 20 years or younger compared with median age 55 years (range 4-86 years) and 30 (5%) diagnosed at 20 years or younger (P < 0.0001). The duration of follow-up between disease diagnosis and study enrolment was longer in transplant centres with a median of 8 (1-57) years versus 6 (1-41) years.

A summary of baseline characteristics (divided into transplant and nontransplant units) at time of study entry is presented in Table 1. In the whole cohort, median body mass index (BMI) at study recruitment was 28.4 kg/m² (range 15.1-64.0). Figure 1 illustrates the proportion of patients in the study according to age at diagnosis divided into five categories of 20-year age brackets. In keeping with other studies, the majority of patients were diagnosed between the age of 41 and 60 years.^{6,30-33}

Seven hundred and thirty-four of 742 (99%) of patients with available data were abstinent from alcohol or consumed alcohol

TABLE 1 Baseline characteristics of patients at time of study entry (n = 1249) and patient-reported presence of other autoimmune conditions (n = 1192, 596 in both types of unit)

Transplant unitsNon-transplant unitsAge at study entry (y) median (range)52 (17-91)63 (18-95)Weight (kg), median (range)75 (40-169)76 (38-165)Height (cm), median (range)162.5 (132.5-192.5)162.5 (132.5-192.5)Body mass index, BMI (kg/m²), median (range)28 (15-64)29 (16-62)
median (range) 75 (40-169) 76 (38-165) Weight (kg), median (range) 162.5 (132.5-192.5) 162.5 (132.5-192.5) Height (cm), median (range) 162.5 (132.5-192.5) 162.5 (132.5-192.5) Body mass index, BMI (kg/m ²), 28 (15-64) 29 (16-62)
median (range) Height (cm), median (range) Body mass index, BMI (kg/m ²),
median (range) Body mass index, 28 (15-64) 29 (16-62) BMI (kg/m²), 29 (16-62) 29 (16-62)
BMI (kg/m²),
Other autoimmune condition n (%) n (%) P value
Thyroid disease 67 (11) 110 (18.5) 0.0004
Rheumatoid arthritis 29 (4.9) 52 (8.7) 0.008
Primary biliary cholangitis 33 (6) 39 (7) 0.47
Ulcerative colitis or Crohn's disease 33 (5.5) 25 (4.2) 0.281
Coeliac disease 17 (2.9) 24 (4) 0.266
Systemic lupus erythematosus 21 (4) 19 (3.2) 0.748
Sjogren's syndrome 15 (2.5) 24 (4) 0.143
Type 1 diabetes mellitus 15 (2.5) 12 (2) 0.559
Primary sclerosing cholangitis 20 (3.4) 7 (1.2) 0.011
Mixed connective tissue disorder 7 (1.2) 4 (0.7) 0.363
Other autoimmune condition 79 (13.3) 100 (16.8) 0.088

within recommended limits in accordance with UK guidelines prior to 2014³⁴ (below 15 units per week for women and 22 units per week for men). Of those with available data, 69/739 (9%) were current smokers (less than the 19% of the total adult UK population who are reported to be current smokers by Cancer Research UK) and 221/739 (30%) were previous smokers.

Fifty-one patients had undergone liver transplantation at time of accession to the study and have been excluded from further analyses (other than for age at diagnosis and the development of cirrhosis) with the denominators reflecting this.

Biochemical data allowing assessment of remission status (defined as normal ALT and IgG at time of study entry) were available for 1114 patients. Of these, 460 (41%) were not in remission at the point of study entry.

3.2 | Immunosuppression regimens

Table 2 summarises treatment regimens, biochemical remission rates and the number of patients with cirrhosis at diagnosis. Patients were stratified according to the treatment regimen. Overall, 29 different treatment regimens were reported, not including dose variations, transplant patients or those not receiving any treatment for their AIH.

Despite treatment guidelines recommending that maintenance corticosteroids not be used, 653 of 1198 patients (55%) were taking long-term corticosteroid therapy as part of their treatment regimen.

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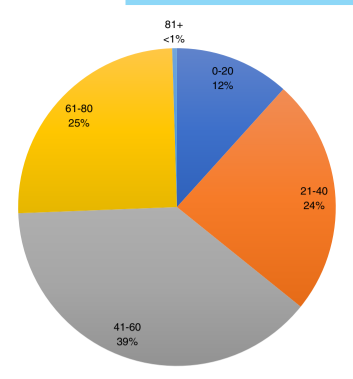


FIGURE 1 Age at diagnosis in 20 y age brackets for whole cohort (n = 1249), n (%). Patients were categorised according to age at diagnosis; 0-20 years, 21-40 y, 41-60 y, 61-80 y, 81 y and over

TABLE 2 Summary of treatment regimens, biochemical remission rates (normal ALT and IgG) and number of patients cirrhotic at diagnosis according to treatment regimen (n = 1198, excluding liver transplant patients)^a

Simplified drug regimen	Number of patients (%)	Number in biochemical remission (%)	Number cirrhotic at diagnosis (%)
Azathioprine/ MP alone	392 (33)	252/362 (70)	82/385 (21)
Azathioprine/MP + prednisolone	316 (26)	175/299 (59)	78/309 (25)
Prednisolone alone	103 (9)	43/85 (51)	28/98 (29)
Budesonide alone	19 (2)	5/19 (26)	2/19 (11)
No immunosuppression	85 (7)	51/79 (65)	15/84 (18)
MMF/MA alone	58 (5)	35/54 (65)	11/57 (19)
MMF/MA + prednisolone	112 (9)	53/107 (50)	34/112 (30)
CNI-containing regimen ^b	56 (5)	16/54 (29)	19/56 (34)
Triple immunosuppression	28 (2)	5/27 (19)	9/27 (33)
Other immunosuppression regimen	53 (4)	24/51 (47)	3/53 (6)

ALT: alanine aminotransferase; IgG: immunoglobulin G; MP: mercaptopurine; MMF: mycophenolate mofetil; MA: myphenolic acid; CNI: calcineurin inhibitor.

^aAll available data presented but some missing data resulting in denominators for the calculation of biochemical remission and numbers cirrhotic at diagnosis being smaller than the total number of included patients. ^bSome patients on a CNI-containing regimen were receiving triple immunosuppression so appear in both groups resulting in n > 1198. **TABLE 3** Details regarding long-term corticosteroid dose and remission rates (n = 1198, excludes transplant patients)^a

Corticosteroid dose	Number of patients (% of cohort)	Number of patients in biochemical remission (%)
≥Prednisolone 10 mg/d ("High dose")	172 (14.4)	57/162 (35.2)
≥Budesonide 6 mg/d ("High dose")	30 (2.5)	7/29 (24)
5 mg > Prednisolone <10 mg/d ("Medium dose")	97 (8.1)	52/91 (57)
Prednisolone ≤5 mg/d ^b ("Low dose or no prednisolone")	929 (77.6)	545/861 (63)
≤ Budesonide 3 mg/d ^b ("Low dose or no prednisolone")	1168 (97.5)	647/1085 (60)

^aAll available data presented but some missing data resulting in denominators for the calculation of biochemical remission being smaller than the total number of included patients.

^bIncluding those on no corticosteroids.

There were 124/1198 (10%) patients on immunosuppression with corticosteroids only (103 [9%] in the form of prednisolone) and of these, 57/106 (54%) patients were not in biochemical remission. Patients requiring lower doses of corticosteroids or on no corticosteroids at all were more likely to be in biochemical remission. These data are summarised in Table 3.

There were 658/1198 (55%) patients taking azathioprine ± prednisolone. The median dose of azathioprine was 1.1 mg/kg/d (range 0.2-2.8). ALT and IgG data were available for 614 patients prescribed azathioprine ± prednisolone, of whom 398 (65%) were in biochemical remission. Biochemical remission rates were lower (256/500 patients, 51% overall) for patients prescribed other treatment regimens. Overall, there were 714 patients taking azathioprine and the dose was available for 696 patients. There were 232 (33%) taking less than the 1 mg/kg/d minimum recommended dose, of whom, 66/ 220 (30%) patients were not in biochemical remission.

A total of 708 patients (59%) were treated with thiopurines either in the form of azathioprine/mercaptopurine alone (392 patients) or in combination with prednisolone (316 patients, 26% of whole cohort). For patients who were taking thiopurine therapy in isolation, 70% (252/362) were in biochemical remission. In contrast, the rate of biochemical remission for patients receiving thiopurine therapy together with prednisolone was significantly lower at 59% (175/299), P = 0.003.

Mycophenolate-based therapy \pm prednisolone (typically used in patients with thiopurine intolerance or nonresponsiveness), were used in 170 patients (14%). 65% (35/54) of patients who received mycophenolate-based therapy (mycophenolate mofetil [MMF] or myphenolic acid [MA]) in isolation were in biochemical remission. However, MMF or MA were used in conjunction with prednisolone in 112 patients (9% of the total cohort, median dose 1000 mg/d [range 250-3000]), and of these, only 50% were in biochemical remission, *P* = 0.066).

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Five per cent of the cohort (59 patients) were receiving a calcineurin inhibitor (CNI) or three drug regimen suggesting difficult to control disease. Of these, only 16/57 (28%) were in biochemical remission and 19/58 (39%) had established cirrhosis at diagnosis. Figure 2 summarises complexity of treatment regimen and likelihood of treatment response.

Seven per cent (85) of patients were receiving no immunosuppression and of these, 65% were in remission with 15 (18%) patients being cirrhotic at diagnosis.

3.3 | Location of care

Of nontransplanted patients (n = 1198), 586 were cared for in transplant units (49%) and 612 in nontransplant units. Treatment regimens utilised in transplant centres were significantly different to those used in nontransplant centres (Table 4). Whilst the total number of combinations of therapy were similar between transplant and nontransplant centres (26 vs 27), patients being cared for in transplant centres were more likely to be on either a triple immunosuppression regimen (P = 0.006) including CNI, anti-CD20 or anti-Tumour Necrosis Factor (TNF) therapy or a regimen including a CNI (P = 0.002).

In nontransplant centres, 405 patients were looked after by a dedicated hepatologist whereas, a gastroenterologist looked after 207 patients. The number of combinations of immunosuppression used by gastroenterologists was fewer than those working as

dedicated hepatologists (22 vs 18). In nontransplant units, there was no statistically significant difference between the use of a CNI or triple immunosuppression between hepatologists and gastroenterologists.

A significantly higher proportion of patients treated in transplant centres were in remission compared to nontransplant centres, 62% versus 55% (P = 0.028). There was no significant difference in disease flares, the proportion of patients maintained on higher doses of corticosteroids (defined in the study protocol as $\geq 10 \text{ mg}$ of prednisolone per day or budesonide $\geq 6 \text{ mg/d}$) or the number who had an attempt to reduce their corticosteroid dose in the previous 12 months between transplant and nontransplant units or care under hepatologist а or gastroenterologist.

3.4 | Development of cirrhosis (includes transplanted patients)

Prevention of the development of cirrhosis in AIH is a major goal of therapy. 289/1223 (24%) of patients were, however, already cirrhotic at the time of diagnosis. Patients diagnosed at 20 years old or younger were significantly more likely to be cirrhotic at diagnosis than patients presenting over the age of 20 (48/141 [34%] vs 241/ 1082 [22%], P = 0.002) although it is unclear whether this represents a more aggressive disease course in children and young adults or a higher likelihood of delayed diagnosis.

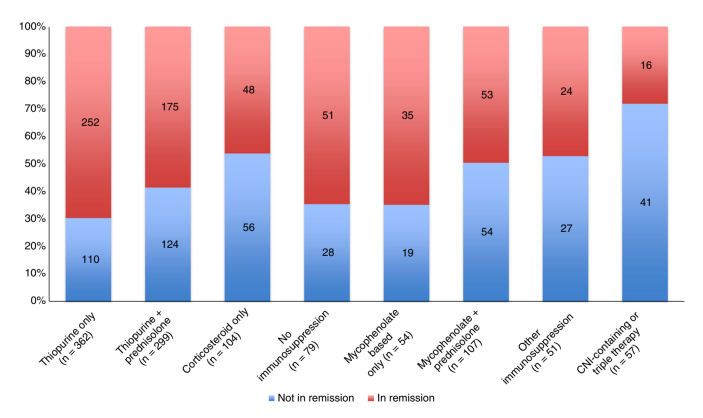


FIGURE 2 Breakdown of patients by remission status according to simplified drug combinations (excluding transplant patients). CNI: calcineurin inhibitor

	Transplant unit (n = 586)	Non-transplant centres (n = 612)	P value	Hepatologist (n = 405)	No hepatologist (n = 207)	P value
Age of patients at diagnosis, median (range)	42 (2-86)	55 (4-86)		55 (2-86)	55 (7-79)	
Number of treatment combinations used	26	27		22	18	
Number of patients on regimen including a CNI	39 (6%)	17 (2.9%)	0.002	8 (1.98%)	9 (4.4%)	0.133
Number of patients on triple immunosuppression	21 (3.6%)	7 (1.14%)	0.006	5 (1.2%)	2 (0.97%)	0.759
Not in biochemical remission (ALT and IgG)	214/562 (38%)	246/552 (44.6%)	0.028	156/366 (42.6%)	90/186 (48.4%)	0.199
Maintenance high-dose corticosteroids during past 12 mo	107/586 (18.3%)	93/612 (15.2%)	0.156	54 (13.3%)	39 (18.8%)	0.085
Attempt to reduce corticosteroid dose during last 12 mo	25/42 (60%)	55/80 (69%)	0.308	35/46 (76.1%)	20/34 (58.8%)	0.101
Previous corticosteroid treatment for disease flare	118/572 (21%)	101/586 (17%)	0.142	67/393 (17%)	34/193 (18%)	0.865
Cirrhotic at diagnosis	145/573 (25.3%)	127/603 (21.1%)	0.085	88/402 (21.9%)	39/201 (19.4%)	0.473

TABLE 4 Comparisons between patients cared for in transplant units and nontransplant centres and, for nontransplant centres, in centres with and without a specialist hepatologist (n = 1198, transplant patients excluded)

CNI: calcineurin inhibitor; ALT: alanine aminotransferase; IgG: immunoglobulin G.

Table 5 summarises risk factors for developing cirrhosis. Patients aged 20 years or younger at diagnosis (including transplant patients) were more likely to develop histological or radiological evidence of cirrhosis during follow-up than patients aged more than 20 years (29% vs 14%, P = 0.0007). The total daily azathioprine and corticosteroid dose were not associated statistically with the development of cirrhosis during follow-up. Whether or not patients were under the care of a hepatologist or gastroenterologist was not associated with disease progression, suggesting that patients with more stable disease were looked after appropriately.

Table 6 shows the uni- and multi-variate analyses for nontransplanted patients. For these analyses, years since diagnosis has been used instead of age at diagnosis to enable exclusion of the transplanted patients, that is, ensuring that the same patient group is used for each variable and that each patient is only included once in the analyses. Increasing years since diagnosis remained significant as a predictor for development of cirrhosis on uni- and multi-variate analysis, Patients in biochemical remission at study entry were less likely to develop cirrhosis, however, this fell short of statistical significance. When examining treatment regimens (compared to standard therapy with azathioprine ± prednisolone), the use of triple immunosuppression or a CNI-containing regimen was associated with a higher probability of developing cirrhosis (P = 0.011) but this effect became just nonsignificant (P = 0.053) on multi-variate analysis when other risk factors were included in the model. Place of care was associated with a more aggressive disease phenotype with transplant units having a greater proportion of patients developing cirrhosis following their initial diagnosis in both the univariate and multiple logistic regression models.

4 | DISCUSSION

In this large, nationwide study of "real world" clinical practice in AIH we have demonstrated both significant limitations in the effectiveness of care for AIH and a high degree of variability in practice and quality between unit types. Our first key observation is that the remission rate using standard criteria is only 59%; a figure falling far short of the benchmark figure of 80% demonstrated to be achievable in specialist centres with a specific interest in disease management and structures in place to deliver optimised care.⁶ Second, over 50% of patients with AIH in the UK are receiving ongoing corticosteroid therapy, in spite of multiple clinical practice guidelines suggesting that maintenance should be with thiopurine monotherapy.⁸⁻¹⁰ Third, there was apparent confusion and inconsistency around appropriate treatment with 29 individual treatment regimens described even before allowing for dose variations. Finally, patients receiving care in transplant centres in the UK were more likely to achieve and maintain remission than those looked after in other care locations, despite an apparent bias towards a more severe disease phenotype.

The demographic characteristics and disease associations of the study population were in keeping with previous published series although concurrent PSC was more common in transplant units.^{4,32,35} This suggests that differences in the behaviour of the cohort long term are unlikely to be attributable to the characteristics of the population, but rather, the care received. Risk variation was seen within the cohort with patients presenting at 20 years or younger having the highest rate of cirrhosis at diagnosis and progression to cirrhosis during follow-up.^{36,37} Disease phenotype profile in this group is in keeping with previously published reports of

TABLE 5 Risk factors for developing cirrhosis (either histological and/or radiological) during follow-up from diagnosis to point of study entry (excluding patients who were cirrhotic at diagnosis)

Risk factor		Number developing cirrhosis (%)		Number developing cirrhosis (%)	P value
Age at diagnosis ^a	≤20 y	27/94 (29)	>20 y	104/836 (14)	0.0007
Azathioprine dose	<1 mg/kg/d	17/176 (10)	≥1 mg/kg/d	33/344 (10)	0.984
Corticosteroid dose	≥ prednisolone 10 mg and/or budesonide 6 mg/d	18/140 (13)	Prednisolone ≤5 mg and/or budesonide 3 mg/d ^b	80/901 (9)	0.134
Specialist care (for nontransplant unit)	Hepatologist	28/311 (9)	No hepatologist	7/160 (4)	0.070

^aTaking the event of liver transplantation as evidence of the development of cirrhosis.

^bIncludes patients on no corticosteroid.

TABLE 6 Univariate and multiple logistic regression model of risk factors for developing cirrhosis (either histological and/or radiological) during follow-up from diagnosis to point of study entry (excluding patients who were cirrhotic at diagnosis

Univariate		Multiple		
Odds ratio	P value	Odds ratio	P value	
0.7033 [0.455, 1.087]	0.1129	0.7514 [0.469, 1.203]	0.2335	
1.8688 [0.955, 3.655]	0.0678	1.5573 [0.76, 3.191]	0.2262	
0.9356 [0.383, 2.288]	0.884	0.8025 [0.303, 2.123]	0.6575	
0.9398 [0.535, 1.651]	0.829	0.9743 [0.536, 1.772]	0.9321	
2.8558 [1.272, 6.411]	0.011	2.3512 [0.99, 5.586]	0.0528	
2.3485 [1.484, 3.716]	0.0003	2.0458 [1.258, 3.327]	0.0039	
1.1029 [1.075, 1.131]	0.0001	1.0972 [1.069, 1.126]	0.0001	
	Odds ratio 0.7033 [0.455, 1.087] 1.8688 [0.955, 3.655] 0.9356 [0.383, 2.288] 0.9398 [0.535, 1.651] 2.8558 [1.272, 6.411] 2.3485 [1.484, 3.716]	Odds ratio P value 0.7033 [0.455, 1.087] 0.1129 1.8688 [0.955, 3.655] 0.0678 0.9356 [0.383, 2.288] 0.884 0.9398 [0.535, 1.651] 0.829 2.8558 [1.272, 6.411] 0.011 2.3485 [1.484, 3.716] 0.0003	Odds ratio P value Odds ratio 0.7033 [0.455, 1.087] 0.1129 0.7514 [0.469, 1.203] 1.8688 [0.955, 3.655] 0.0678 1.5573 [0.76, 3.191] 0.9356 [0.383, 2.288] 0.884 0.8025 [0.303, 2.123] 0.9398 [0.535, 1.651] 0.829 0.9743 [0.536, 1.772] 2.8558 [1.272, 6.411] 0.011 2.3512 [0.99, 5.586] 2.3485 [1.484, 3.716] 0.0003 2.0458 [1.258, 3.327]	

CNI: calcineurin inhibitor.

^aStandard therapy = azathioprine ± prednisolone.

children with AIH, particularly, Type 2 AIH associated with detectable anti-Liver Kidney Microsomal (LKM) antibodies in serum.³⁸

The rate of corticosteroid use was high in our cohort with 55% of patients remaining on either prednisolone or budesonide. Recently published data from the UK-AIH study show that the use of corticosteroids is strongly associated with decreased health-related quality of life that is independent of biochemical remission status.³⁹ BMI was high in our cohort, with the median being 28.4 kg/m², and 38% (421/1111) of patients being classified as obese with a BMI > 30 kg/m². This may reflect the overly high use of corticosteroids. Corticosteroid therapy, in conjunction with weight gain, is likely to result in secondary disease development such as the metabolic syndrome, hyperlipidaemia and hypertension in this patient population. What is clear from older data is that when it is possible to entirely withdraw corticosteroids in patients with AIH, an average of 6 kg of weight loss per patient has been reported.¹²

The overall use of thiopurine therapy, either azathioprine or mercaptopurine (±corticosteroid) was 63% (753//1198 patients) of the total patient cohort with 359/753 (48%) taking corticosteroid therapy. This suggests that adherence to established treatment guidelines for AIH patients in the UK (European Association for the Study of the Liver and British Society of Gastroenterology guidelines) is poor. The median dose of azathioprine used in this cohort was only 1.1 mg/kg/d. Thiopurine drug metabolites were not checked in this study (the study protocol precluded this), nor were details of poor treatment tolerance with side effects or adverse events collected. However, we know from published data that an azathioprine dose of up to 2 mg/kg/d can result in enhanced long-term remission rates in AIH with a concomitant ability to withdraw corticosteroids entirely from the treatment regimen for the majority of patients.⁴⁰

The approach to management of AIH was different between transplant and nontransplant centres. While the overall number of treatment regimens used in transplant compared to nontransplant units was similar (26 vs 27), there was a greater likelihood of patients being exposed to an expanded range of novel treatment options in AIH management (eg, CNIs, three drug regimens or biological agents). This more nuanced and individualised approach to care in the transplant centres appears to translate to less fluctuation in the disease, with more patients in biochemical remission, and consequently is likely to be protective in relation to hepatic outcomes.⁶

This "real world" study of patients with AIH demonstrates and quantifies the therapeutic challenges that have been discussed in the literature.^{22,41,42} The poor remission rates that we report suggest that there are significant unmet needs therapeutically for patients with AIH. For the majority of other autoimmune disorders, significant strides have taken place to facilitate corticosteroid-free

regimens. Examples include the use of disease-modifying agents and novel antibodies in diseases such as those utilised in Multiple Sclerosis, Inflammatory Bowel Disease and Rheumatoid Arthritis.43,44 These conditions are exemplars of being beneficiaries of the novel therapeutics explosion. As an orphan disease with potential hard outcomes such as cirrhosis development, death and liver transplantation, there is a pressing need for novel therapeutic approaches and targets in AIH. Some potential targets for AIH treatment have been explored but none have been realised in clinical practice as yet.42 Patients with liver disease, especially those with rare diseases such as AIH, are disadvantaged not just through inertia from the medical community, but are secondarily disadvantaged through an unwillingness of the pharmaceutical industry to offer the use of potentially useful therapeutic agents in clinical trials. As an example, current literature suggests that antibody therapies such as anti-CD20 or anti-TNF therapy have only been utilised in <50 patients worldwide with AIH, and even then, only in the context of late disease. $^{\rm 45,46}$ In contrast, the alternative treatment paradigm should be one of treating early disease aggressively with potential disease modifiers or stoppers, to avoid life-long therapy with corticosteroids and other drugs that have been proven outdated for the majority of other inflammatory autoimmune disorders.

Although we believe our data shed important light on the reality of care for AIH in the UK the study has important limitations. The first is that this is an observational cohort study describing outcomes in practice. This limits the scale of the data capture and will clearly miss any very high-risk patients who died from the disease early in its course and who would thus feature in an incident but not a prevalent cohort. Our approach does avoid, however, the potential for an intensive prospective study to focus attention on, and potentially lead to artificial improvement in the quality of the care being delivered. The second is that, self-evidently, the study relates only to practice in the UK. It would of course be of great interest to repeat the approach in other health care settings to explore whether the same limitations in care are present. Anecdote and the limited data available suggest that they are. The third is that our categorisation into transplant units and nontransplant units was a robust but slightly blunt approach. We have tried to account for centres with a specialist interest in AIH (where excellent results are reported⁶) by sub-dividing nontransplant units into those with and without dedicated hepatologists (data courtesy of Jessica Dyson and Mark Hudson from national survey of liver services). It may be that these centres improve the apparent outcomes in the nontransplant centre group as a whole (as seen with the significantly lower rate of disease flares in centres with a hepatologist), masking the true scale of the care quality divide.

In conclusion, despite its limitations the UK-AIH cohort demonstrates significant discrepancies in care delivery for patients with AIH. It outlines, in particular, the contrast between real world outcomes for a rare disease cohort and the outcomes achieved in clinical trials. It also suggests that the medical community seems comfortable in accepting both suboptimal patient outcomes and largely outmoded therapeutics for the disorder. This cohort provides evidence of the need to enhance adherence to optimal treatment approaches identified in clinical practice guidelines (through education of both clinicians and patients) and a need for more potent, and patient-acceptable, therapies for this important condition. Both areas should be research priorities moving forward.

ACKNOWLEDGEMENTS

We are indebted to all the members of the clinical and research teams in centres who have recruited to UK-AIH and are key to the ongoing success of the study. The principal investigators and recruiting centres are: Dr Shirley English (Aberdeen Royal Infirmary), Dr Graeme Alexander/Dr George Mells (Addenbrooke's Hospital), Dr Debabrata Majumdar (St Peters Hospital), Dr Vinay Sathyanarayana (Barnsley Hospital), Professor John Ramage (Basingstoke & North Hampshire Hospital), Dr Christopher Shorrock (Blackpool Victoria Hospital), Dr James Maggs (Buckinghamshire Hospital), Dr David Elphick (Chesterfield Royal Hospital), Dr Chris Macdonald (Cumberland Infirmary), Professor Matthew Cramp (Derriford Hospital), Dr Joanne Sayer (Doncaster Royal Infirmary), Dr James Jupp (Dorset County Hospital), Dr Jessica Dyson (Freeman Hospital), Dr Coral Hollywood (Gloucestershire Royal Hospital), Dr Alexandra Daley (Heartlands Hospital), Dr Lynsey Corless (Hull Royal Infirmary), Dr Darren Craig (James Cook University Hospital), Dr Jane Collier (John Radcliffe Hospital), Professor Michael Heneghan (King's College Hospital), Dr Sharat Misra (King's Mill Hospital), Dr Chris Corbett (New Cross Hospital), Professor John Dillon (Ninewells Hospital), Dr Simon Rushbrook (Norfolk and Norwich University Hospital), Dr Thomas Lee (North Tyneside General Hospital), Dr Nicholas M Sharaer (Poole Hospital), Dr Kara Rye (Princess Royal Hospital), Dr Andrew Fowell (Queen Alexandra Hospital, Portsmouth), Dr Andrea Broad/Dr Dina Mansour (Queen Elizabeth Hospital, Gateshead), Dr Andy Douds (Queen Elizabeth Hospital, King's Lynn), Dr Stephen Ryder (Queen's Medical Centre), Dr Richard Keld (Royal Albert Edward Infirmary), Dr Earl Williams (Royal Bournemouth Hospital), Dr William Stableforth (Royal Cornwall Hospital), Dr Andrew Austin (Royal Derby Hospital), Professor Dermot Gleeson (Royal Hallamshire Hospital), Dr Kenneth Simpson (Royal Infirmary of Edinburgh), Dr Imran Patanwala (Royal Liverpool University Hospital), Dr Alison Brind (Royal Stoke University Hospital), Dr Shanika de Silva (Russells Hall Hospital), Dr Aqueel Jamil (Salisbury District Hospital), Dr Saket Singhal (Sandwell General Hospital), Dr Chin Lye Ch'ng (Singleton Hospital), Dr Joanne Topping (South Tyneside District Hospital), Dr Mark Wright (Southampton General Hospital), Dr Talal Valliani (Southmead Hospital), Dr Rebecca Jones (St. James's University Hospital, Leeds), Dr Harriet Mitchison (Sunderland Royal Hospital), Dr Douglas Thorburn (The Royal Free Hospital), Professor Aftab Ala (The Royal Surrey County Hospital), Dr Ye Htun Oo (University Hospital Birmingham NHS Foundation Trust), Dr Sushma Sakena/Dr Francisco Porras-Perez (University Hospital of North Durham), Prof Jane Metcalf/Dr Stephen Mitchell (University Hospital of North Tees), Dr Esther Unitt/Dr Victoria Gordon (University Hospitals Coventry & Warwick), and Dr Jeremy Shearman (Warwick Hospital,

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South Warwickshire NHS Foundation Trust). We would also like to thank Mark Hudson (Consultant Hepatologist at Freeman Hospital, Newcastle), the British Society of Gastroenterology (BSG) and the British Association for the Study of the Liver (BASL) who supported the collection of data regarding the location of hepatologists in the UK.

Declaration of personal interests: SK is a GSK employee and stock-holder. YHO was funded by MRC Clinician Scientist Award (G1002552). LLW, TB, GMH, AWL and MAH have nothing to declare. DEJJ has provided consultancy for Novartis. JKD is supported by the NIHR Newcastle Biomedical Research Centre and the NIHR Rare Diseases Translational Research Collaboration.

Declaration of funding interests: The UK-AIH study was funded by the NIHR Rare Diseases Translational Research Collaboration. This paper presents independent research funded by the NIHR (Newcastle and Birmingham Biomedical Research Centres). The views expressed are those of the author(s) and not necessarily those of the NHS, the NIHR or the Department of Health.

AUTHORSHIP

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Author contributions: LLW, JKD and TB analysed and interpreted the data. JKD, DEJJ, MAH, LLW and TB wrote the first draft of the manuscript. JKD, DEJJ and SK were involved in study concept and design. All authors were involved in the critical revision of the manuscript. All authors approved the final version of the manuscript.

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REFERENCES

- 1. Heneghan MA, Yeoman AD, Verma S, Smith AD, Longhi MS. Autoimmune hepatitis. *Lancet*. 2013;382(9902):1433-1444.
- Manns MP, Lohse AW, Vergani D. Autoimmune hepatitis–Update 2015. J Hepatol. 2015;62(Suppl 1):S100-S111.
- 3. Krawitt EL. Autoimmune hepatitis. N Engl J Med. 2006;354:54-66.
- Al-Chalabi T, Boccato S, Portmann BC, McFarlane IG, Heneghan MA. Autoimmune hepatitis (AIH) in the elderly: a systematic retrospective analysis of a large group of consecutive patients with definite AIH followed at a tertiary referral centre. J Hepatol. 2006;45:575-583.
- Boberg KM, Aadland E, Jahnsen J, Raknerud N, Stiris M, Bell H. Incidence and prevalence of primary biliary cirrhosis, primary sclerosing cholangitis, and autoimmune hepatitis in a Norwegian population. *Scand J Gastroenterol*. 1998;33:99-103.
- 6. Hoeroldt B, McFarlane E, Dube A, et al. Long-term outcomes of patients with autoimmune hepatitis managed at a nontransplant center. *Gastroenterology*. 2011;140:1980-1989.
- Muratori L, Muratori P, Lanzoni G, Ferri S, Lenzi M. Application of the 2010 American Association for the study of liver diseases criteria of remission to a cohort of Italian patients with autoimmune hepatitis. *Hepatology*. 2010;52:1857; author reply 1857-8.
- European Association for the Study of the L. EASL Clinical Practice Guidelines: autoimmune hepatitis. J Hepatol. 2015;63:971-1004.

- Gleeson D, Heneghan MA, British Society of G. British Society of Gastroenterology (BSG) guidelines for management of autoimmune hepatitis. *Gut.* 2011;60:1611-1629.
- 10. Manns MP, Czaja AJ, Gorham JD, et al. Diagnosis and management of autoimmune hepatitis. *Hepatology*. 2010;51:2193-2213.
- 11. van Gerven NM, Verwer BJ, Witte BI, et al. Relapse is almost universal after withdrawal of immunosuppressive medication in patients with autoimmune hepatitis in remission. *J Hepatol.* 2013;58: 141-147.
- Johnson PJ, McFarlane IG, Williams R. Azathioprine for long-term maintenance of remission in autoimmune hepatitis. N Engl J Med. 1995;333:958-963.
- McFarlane E, Walters SJ, Peters J, et al. Quality of life in patients with autoimmune hepatitis measured with the SF-36 health status questionnaire. *Gut.* 2004;53(Suppl 3):A84.
- 14. Schramm C, Wahl I, Weiler-Normann C, et al. Health-related quality of life, depression, and anxiety in patients with autoimmune hepatitis. *J Hepatol.* 2014;60:618-624.
- Sockalingam S, Blank D, Abdelhamid N, Abbey SE, Hirschfield GM. Identifying opportunities to improve management of autoimmune hepatitis: evaluation of drug adherence and psychosocial factors. J Hepatol. 2012;57:1299-1304.
- Hubener S, Oo YH, Than NN, et al. Efficacy of 6-mercaptopurine as second-line treatment for patients with autoimmune hepatitis and azathioprine intolerance. *Clin Gastroenterol Hepatol.* 2016;14:445-453.
- Manns MP, Woynarowski M, Kreisel W, et al. Budesonide induces remission more effectively than prednisone in a controlled trial of patients with autoimmune hepatitis. *Gastroenterology*. 2010;139:1198-1206.
- Sharzehi K, Huang MA, Schreibman IR, Brown KA. Mycophenolate mofetil for the treatment of autoimmune hepatitis in patients refractory or intolerant to conventional therapy. *Can J Gastroenterol*. 2010;24:588-592.
- Than NN, Wiegard C, Weiler-Normann C, et al. Long-term follow-up of patients with difficult to treat type 1 autoimmune hepatitis on Tacrolimus therapy. Scand J Gastroenterol. 2016;51:329-336.
- Baven-Pronk AM, Coenraad MJ, van Buuren HR, et al. The role of mycophenolate mofetil in the management of autoimmune hepatitis and overlap syndromes. *Aliment Pharmacol Ther.* 2011;34:335-343.
- Kirk AP, Jain S, Pocock S, Thomas HC, Sherlock S. Late results of the Royal Free Hospital prospective controlled trial of prednisolone therapy in hepatitis B surface antigen negative chronic active hepatitis. *Gut.* 1980;21:78-83.
- Liberal R, de Boer YS, Andrade RJ, et al. Expert clinical management of autoimmune hepatitis in the real world. *Aliment Pharmacol Ther*. 2017;45:723-732.
- Williams R, Alexander G, Aspinall R, et al. New metrics for the Lancet Standing Commission on Liver Disease in the UK. *Lancet*. 2017;389(10083):2053-2080.
- Williams R, Ashton K, Aspinall R, et al. Implementation of the Lancet Standing Commission on Liver Disease in the UK. *Lancet*. 2015;386 (10008):2098-2111.
- Williams R, Aspinall R, Bellis M, et al. Addressing liver disease in the UK: a blueprint for attaining excellence in health care and reducing premature mortality from lifestyle issues of excess consumption of alcohol, obesity, and viral hepatitis. *Lancet*. 2014;384(9958):1953-1997.
- Dyson JK, Webb G, Hirschfield GM, et al. Unmet clinical need in autoimmune liver diseases. J Hepatol. 2015;62:208-218.
- Sebode M, Hartl J, Vergani D, Lohse AW, International Autoimmune Hepatitis G. Autoimmune hepatitis: from current knowledge and clinical practice to future research agenda. *Liver Int.* 2017;38:15-22.

- $I_{\rm L}$ **E Y** AP_&T Alimentary Pharmacology & Therapeutics
- Alvarez F, Berg PA, Bianchi FB, et al. International Autoimmune Hepatitis Group Report: review of criteria for diagnosis of autoimmune hepatitis. J Hepatol. 1999;31:929-938.
- Hennes EM, Zeniya M, Czaja AJ, et al. Simplified criteria for the diagnosis of autoimmune hepatitis. *Hepatology*. 2008;48: 169-176.
- 30. Danielsson Borssen A, Marschall HU, Bergquist A, et al. Epidemiology and causes of death in a Swedish cohort of patients with autoimmune hepatitis. *Scand J Gastroenterol.* 2017;52:1022-1028.
- Gronbaek L, Vilstrup H, Jepsen P. Autoimmune hepatitis in Denmark: incidence, prevalence, prognosis, and causes of death. A nationwide registry-based cohort study. J Hepatol. 2014;60:612-617.
- van Gerven NM, Verwer BJ, Witte BI, et al. Epidemiology and clinical characteristics of autoimmune hepatitis in the Netherlands. *Scand J Gastroenterol*. 2014;49:1245-1254.
- Werner M, Prytz H, Ohlsson B, et al. Epidemiology and the initial presentation of autoimmune hepatitis in Sweden: a nationwide study. Scand J Gastroenterol. 2008;43:1232-1240.
- National Institute for Health and Care Excellence (NICE). Local government briefing - Alcohol - Facts and Figures. 2012; https://www. nice.org.uk/advice/lgb6/chapter/facts-and-figures. Accessed December 2017.
- Wong GW, Yeong T, Lawrence D, Yeoman AD, Verma S, Heneghan MA. Concurrent extrahepatic autoimmunity in autoimmune hepatitis: implications for diagnosis, clinical course and long-term outcomes. *Liver Int.* 2017;37:449-457.
- Ngu JH, Gearry RB, Frampton CM, Stedman CA. Predictors of poor outcome in patients w ith autoimmune hepatitis: a population-based study. *Hepatology*. 2013;57:2399-2406.
- Montano-Loza AJ, Carpenter HA, Czaja AJ. Features associated with treatment failure in type 1 autoimmune hepatitis and predictive value of the model of end-stage liver disease. *Hepatology*. 2007;46:1138-1145.
- Gregorio GV, Portmann B, Reid F, et al. Autoimmune hepatitis in childhood: a 20-year experience. *Hepatology*. 1997;25:541-547.

- Wong LL, Fisher HF, Stocken DD, et al. The impact of autoimmune hepatitis and its treatment on health utility. *Hepatology*. 2018; https://doi.org/10.1002/hep.3003
- Stellon AJ, Keating JJ, Johnson PJ, McFarlane IG, Williams R. Maintenance of remission in autoimmune chronic active hepatitis with azathioprine after corticosteroid withdrawal. *Hepatology*. 1988;8: 781-784.
- Zachou K, Muratori P, Koukoulis GK, et al. Review article: autoimmune hepatitis—current management and challenges. *Aliment Pharmacol Ther*. 2013;38:887-913.
- 42. Czaja AJ. Review article: the management of autoimmune hepatitis beyond consensus guidelines. *Aliment Pharmacol Ther.* 2013;38:343-364.
- 43. Burmester GR, Pope JE. Novel treatment strategies in rheumatoid arthritis. *Lancet*. 2017;389(10086):2338-2348.
- Ontaneda D, Thompson AJ, Fox RJ, Cohen JA. Progressive multiple sclerosis: prospects for disease therapy, repair, and restoration of function. *Lancet.* 2017;389(10076):1357-1366.
- Burak KW, Swain MG, Santodomingo-Garzon T, et al. Rituximab for the treatment of patients with autoimmune hepatitis who are refractory or intolerant to standard therapy. *Can J Gastroenterol*. 2013;27:273-280.
- Weiler-Normann C, Schramm C, Quaas A, et al. Infliximab as a rescue treatment in difficult-to-treat autoimmune hepatitis. J Hepatol. 2013;58:529-534.

How to cite this article: Dyson JK, Wong LL, Bigirumurame T, et al. Inequity of care provision and outcome disparity in autoimmune hepatitis in the United Kingdom. *Aliment Pharmacol Ther*. 2018;48:951–960. <u>https://doi.org/10.1111/</u> apt.14968

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