Preparing for the NASH epidemic: A call to action

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Nonalcoholic fatty liver disease (NAFLD) and nonalcoholic steatohepatitis (NASH) are common conditions with a rising burden. Yet there are significant management gaps between clinical guidelines and practice in patients with NAFLD and NASH. Further, there is no single global guiding strategy for the management of NAFLD and NASH. The American Gastroenterological Association, in collaboration with 7 professional associations, convened an international conference comprising 32 experts in gastroenterology, hepatology, endocrinology, and primary care providers from the United States, Europe, Asia, and Australia. Conference content was informed by the results of a national NASH Needs Assessment Survey. The participants reviewed and discussed published literature on global burden, screening, risk stratification, diagnosis, and management of individuals with NAFLD, including those with NASH. Participants identified promising approaches for clinical practice and prepared a comprehensive, unified strategy for primary care providers and relevant specialists encompassing the full spectrum of NAFLD/NASH care. They also identified specific high-yield targets for clinical research and called for a unified, international public health response to NAFLD and NASH.

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published guidance and clinical practice in patients with NAFLD and NASH [8,9]. Much of this disparity could come from a lack of recognition of the importance of NAFLD/NASH and an absence of a unified strategy that encompasses all disciplines involved in managing these patients across the full disease spectrum.

To address this need, the American Gastroenterological Association (AGA) conducted a needs assessment survey of health professionals likely to be engaged in managing adult patients with NAFLD/NASH, followed by a virtual conference of international experts representing 7 professional societies to review the current research and outline the future agenda for clinical practice, research, and policy. The overarching goal was to call for a unified, international public health response to NAFLD and NASH. This report summarizes the results from the survey and the virtual conference. “Preparing for a NASH Epidemic: A Call for Action.” Although NAFLD is an important and growing problem in children, the current effort was limited to adults with NAFLD and NASH. Therefore, we do not cover pediatric NAFLD in this report.

1. Nonalcoholic Steatohepatitis Needs Assessment Survey

The NASH Needs Assessment Survey was conducted in May 2020. The survey sought to assess participants’ knowledge related to screening, diagnosis, and management of NAFLD and NASH; compare current diagnostic and treatment patterns with the most recent practice guidance on NAFLD/NASH; and identify the educational needs that could serve as targets to improve implementation of guideline-based treatment of NAFLD and NASH. The survey included 24 questions regarding screening, diagnosing, and managing NASH (see Supplementary Material for the full survey). In total, 751 gastroenterologists, hepatologists, endocrinologists, and PCPs from 46 states across the United States completed the survey. More than 50% of survey participants were PCPs. Respondents had spent an average of 19.5 years in practice (range, 2–35 years).

The survey revealed significant gaps in knowledge about who to screen and how to diagnose and treat patients at high risk for NASH, including disparities between published practice guidance and clinical practice (Table 1). Most respondents (67%) from all practice types were aware that up to one-quarter of the general population may have NAFLD. However, there were shortfalls in the knowledge about prevalence in several high-risk groups. For example, only 35% of all respondents—including 28% of endocrinologists, 32% of PCPs, and 46% of gastroenterologists/hepatologists—recognized that almost all patients with severe obesity are likely to have NAFLD. Only 49% of endocrinologists and 45% of PCPs recognized that NAFLD is very common in patients with type 2 diabetes (T2D) (Table 1).

Most participants reported that they screen patients with abnormal liver chemistries (96%), those with T2D (87%), and those who are older than 50 years with hypertension and hyperlipidemia (70%) for the presence of NAFLD. Most were also aware of the best practices in the initial evaluation of patients with suspected NAFLD, including the need to exclude competing etiologies (96%) and evaluation for commonly associated comorbidities, such as T2D, obesity, and dyslipidemia (96%). However, only 41% recognized that initial evaluation of patients with suspected NAFLD should not include cross-sectional abdominal imaging (e.g., contrast-enhanced computed tomography) to screen for HCC. There were no significant differences in the responses among gastroenterologists/hepatologists, endocrinologists, and PCPs.

More than 80% of participants were aware that noninvasive tests, including the NAFLD fibrosis score, Fibrosis-4 Index, and imaging-based tests, such as vibration-controlled transient elastography or magnetic

### Table 1

<table>
<thead>
<tr>
<th>Variable</th>
<th>All participants (n = 751)</th>
<th>Gastroenterologists/hepatologists (n = 175)</th>
<th>Endocrinologists (n = 175)</th>
<th>Primary care (n = 401)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Proportions of the key patient groups likely to have NAFLD</td>
<td></td>
<td></td>
<td></td>
<td></td>
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<tr>
<td>Patients with severe obesity</td>
<td>35</td>
<td>46</td>
<td>28</td>
<td>32</td>
</tr>
<tr>
<td>With T2D</td>
<td>50</td>
<td>62</td>
<td>49</td>
<td>45</td>
</tr>
<tr>
<td>With dyslipidemia</td>
<td>40</td>
<td>47</td>
<td>41</td>
<td>36</td>
</tr>
<tr>
<td>General population</td>
<td>67</td>
<td>79</td>
<td>65</td>
<td>62</td>
</tr>
<tr>
<td>Patient groups that should be screened for NAFLD</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Patients with abnormal liver chemistry</td>
<td>96</td>
<td>97</td>
<td>97</td>
<td>85</td>
</tr>
<tr>
<td>Patients with T2D</td>
<td>87</td>
<td>88</td>
<td>94</td>
<td>83</td>
</tr>
<tr>
<td>Patients older than 50 y who have hypertension and hyperlipidemia</td>
<td>70</td>
<td>71</td>
<td>73</td>
<td>67</td>
</tr>
<tr>
<td>Approaches to the initial evaluation of the patient with suspected NAFLD</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Exclude competing etiologies for steatosis and coexisting common chronic liver disease</td>
<td>96</td>
<td>95</td>
<td>95</td>
<td>97</td>
</tr>
<tr>
<td>Consider the presence of commonly associated comorbidities, such as obesity, dyslipidemia, insulin resistance, or diabetes</td>
<td>95</td>
<td>97</td>
<td>93</td>
<td>95</td>
</tr>
<tr>
<td>Cross-sectional abdominal imaging (such as contrast-enhanced CT scan) to screen for HCC</td>
<td>41</td>
<td>50</td>
<td>39</td>
<td>38</td>
</tr>
<tr>
<td>Knowledge about strategies for noninvasive diagnosis of steatohepatitis and advanced fibrosis in NAFLD</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>NAFLD fibrosis score or Fibrosis-4 Index are useful tools for identifying NAFLD patients with high likelihood of advanced fibrosis</td>
<td>82</td>
<td>94</td>
<td>86</td>
<td>75</td>
</tr>
<tr>
<td>VCTE (FibroScan) or MRE (imaging) are useful tools for identifying advanced fibrosis in patients with NAFLD</td>
<td>81</td>
<td>93</td>
<td>85</td>
<td>74</td>
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<tr>
<td>Abdominal ultrasound is a useful tool for identifying NAFLD patients with steatohepatitis</td>
<td>16</td>
<td>29</td>
<td>18</td>
<td>9</td>
</tr>
<tr>
<td>Appropriateness of treatments for NASH</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>GLP-1 agonists</td>
<td>16</td>
<td>21</td>
<td>15</td>
<td>15</td>
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<tr>
<td>Metformin</td>
<td>17</td>
<td>33</td>
<td>17</td>
<td>11</td>
</tr>
<tr>
<td>Obeticholic acid</td>
<td>15</td>
<td>33</td>
<td>13</td>
<td>9</td>
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<tr>
<td>Omega-3 fatty acids</td>
<td>23</td>
<td>37</td>
<td>23</td>
<td>16</td>
</tr>
<tr>
<td>Pioglitazone&lt;sup&gt;a&lt;/sup&gt;</td>
<td>53</td>
<td>53</td>
<td>77</td>
<td>42</td>
</tr>
<tr>
<td>Ursodeoxycholic acid</td>
<td>22</td>
<td>49</td>
<td>17</td>
<td>12</td>
</tr>
<tr>
<td>Vitamin E for nondiabetic adults&lt;sup&gt;a&lt;/sup&gt;</td>
<td>40</td>
<td>71</td>
<td>51</td>
<td>38</td>
</tr>
</tbody>
</table>

NOTE: Data represent percentages of participants who answered the item correctly.

CT, computed tomography; MRE, magnetic resonance elastography; VCTE, vibration-controlled transient elastography.

<sup>a</sup> The estimates for pioglitazone and vitamin E indicate percentages of participants who would consider treatment overall (with or without liver biopsy).
resonance elastography, are clinically useful tools for identifying NAFLD/NASH patients with a high likelihood of advanced liver fibrosis. However, 78% also thought that abdominal ultrasound can identify NAFLD patients with NASH.

Most participants were aware that 7%–10% weight loss is recommended for patients with NAFLD, but fewer than half of the participants were aware that pioglitazone or vitamin E can be recommended as treatment in select patients with NASH. Most respondents (>80%) wanted more education about screening, diagnosis, and treatment of NAFLD/NASH.

2. A call-to-action conference

To address these knowledge gaps, the AGA convened a virtual conference of international experts in gastroenterology, hepatology, endocrinology, obesity management, and primary care on July 10, 2020. Participants represented key opinion leaders from 8 professional societies, and practiced in the United States, Europe, Australia, and Asia. See the Supplementary Material for the names and affiliations of all participants.

In a series of preconference meetings conducted over 2 months (May and June 2020), these key opinion leaders met and discussed the most important and potentially controversial aspects of the current NAFLD/NASH landscape, including epidemiology, risk factors, screening, diagnosis, and management issues. Formal presentations by each participant followed during the 1-day conference, which included the best-available evidence about their topic. Subsequent to the meeting, workgroups ( predefined by subject) reviewed, discussed, and collated a summary from all presentations in their respective sections, followed by an internal review of the summary from all workgroup members. The final manuscript (including summaries from each workgroup) was then submitted to the full group for a second round of input and approval. The sections here detail the discussion, conclusions, and recommendations for clinical practice and future research that emerged from this process.

3. Burden of nonalcoholic fatty liver disease and nonalcoholic steatohepatitis

The clinical burden of both NAFLD overall and NASH specifically has increased steadily since the 1980s. NAFLD currently affects 25% of the global population and ~60% of patients with T2D [10]. Studies evaluating the prevalence of NASH suggest that it may involve an estimated 1.5%–6.5% of the general population and as many as 37% of people with T2D [10]. Prevalence of NASH is expected to increase by 63% between 2015 and 2030 [11]. Although these numbers seem substantially lower than those for NAFLD overall, they still translate to 4.9 million to 21 million Americans and more than 100 million individuals worldwide. Modeling data estimate that the number of patients with NAFLD-related advanced fibrosis will likely double by 2030, resulting in 800,000 liver-related deaths [11].

NASH is already the number 1 indication for liver transplantation in women, patients older than 54 years, and Medicare recipients [12]. Beyond the significant impairment of quality of life experienced by individuals with NASH and advanced fibrosis [10,13], Younossi et al [14] estimated in 2017 that the overall lifetime direct costs of NASH in the United States would be $222.6 billion, and approximately $95.4 billion estimated in 2017 that the overall lifetime direct costs of NASH in the United States would be $222.6 billion, and approximately $95.4 billion.

In 2020, for patients with NAFLD, the average lifetime direct cost of NASH was $60,000 (95% CI, $22,800–$115,000) for stage 1; $420,300 (95% CI, $35,100–$510,340) for stage 4; and $1,350,000 (95% CI, $420,300–$3,450,000) for stage 5. These results were more pronounced for risk of liver-related mortality, which increased exponentially with each increase in fibrosis stage, from an RR of 1.41 (95% CI, 0.17–11.95) for stage 1 to an RR of 9.57 (95% CI, 1.67–54.93) for stage 2, and an RR of 42.30 (95% CI, 3.51–510.34) for stage 4 fibrosis [5].

Notably, fibrogenesis does not proceed linearly from simple fatty liver to NASH to cirrhosis, but progresses and regresses in up to 30% of patients during a mean period of 5 years [20]. Furthermore, many patients with isolated hepatic steatosis, previously thought to be benign, are likely to progress to NASH [20]. On average, patients with NASH and NASH progress 1 stage of fibrosis every 7 and 14 years, respectively [21]. Older age, visceral obesity, T2D, and hypertension are associated with fibrosis progression [21,22]. T2D and number of metabolic comorbidities are also associated with an increased risk of liver-related mortality and HCC [23,24]. The severity of steatosis, however, has a modest (if any) correlation with the severity of liver histology [25], and the relationship between severity of steatosis and cardiovascular disease remains unclear.

4. Risk factors for nonalcoholic fatty liver disease, nonalcoholic steatohepatitis, and related complications

Patients with obesity or T2D are at a higher risk of developing NAFLD/NASH [15,16]. Conversely, patients with NAFLD are at an increased risk of T2D [17]. NAFLD and especially NASH are independently associated with several liver-related complications, including cirrhosis, HCC, and liver-related mortality. Patients with NAFLD also have a 2-fold increase in risk of cardiovascular disease [18,19]. Indeed, individuals with NAFLD/NASH are twice as likely to die of cardiovascular disease as liver disease [17]. The risk of cardiovascular disease in NAFLD is not completely explained by the shared risk factors, and might be related in part to abnormalities of cardiac structure and function [17].

In patients with NAFLD, the strongest histologic determinant of hepatic and overall outcomes is the presence and stage of fibrosis, although the presence of NASH is the driving force for fibrosis development. Patients with histologic evidence of fibrosis higher than stage 2 are at higher risk for adverse outcomes (hepatic decompensation, HCC, and liver-related mortality), and this risk increases as fibrosis advances to cirrhosis [5]. Specifically, a recent meta-analysis found that, compared to NAFLD patients with no fibrosis (stage 0), patients with fibrosis were at an increased risk for all-cause mortality, and this risk increased with the stage of fibrosis: stage 1: risk ratio (RR) vs stage 0, 1.58 (95% confidence interval [CI], 1.19–2.11); stage 2: RR, 2.52 (95% CI, 1.85–3.42); stage 3: RR, 3.48 (95% CI, 2.51–4.83); and stage 4: RR, 6.40 (95% CI, 4.11–9.95). The results were more pronounced for risk of liver-related mortality, which increased exponentially with each increase in fibrosis stage, from an RR of 1.41 (95% CI, 0.17–11.95) for stage 1 to an RR of 9.57 (95% CI, 1.67–54.93) for stage 2, and an RR of 42.30 (95% CI, 3.51–510.34) for stage 4 fibrosis [5].

To recognize NAFLD, the PCP must be aware of the following facts:

1. NAFLD is the one of the most common causes of abnormal liver enzymes, but serum alanine aminotransferase (ALT) and aspartate aminotransferase (AST) can be normal in many cases of NAFLD/NASH at all stages, including in patients with advanced fibrosis [27].

2. Liver fibrosis has been linked to morbidity and reduced overall patient survival [28].

3. NAFLD and fibrosis are reversible with weight loss [29].

4. Alcohol causes fatty liver disease with many histologic features of NAFLD. Although good clinical history is extremely important, one way to differentiate alcoholic from nonalcoholic fatty liver is the AST/ALT ratio, which is generally ≥2 in patients with alcohol as the underlying cause. In certain patients, selective testing for alcohol metabolites may also be appropriate.

Clinical practice guidelines do not recommend screening for NAFLD in the general population, but case finding for NASH and significant fibrosis is advised for key high-risk groups, such as those with moderate...
to severe obesity (body mass index $\geq 35$ kg/m$^2$), T2D of more than 10 years’ duration or in people older than 50 years, or metabolic syndrome [30]. The American Diabetes Association’s 2020 Standards of Medical Care in Diabetes also recommend evaluating patients with prediabetes or T2D with steatosis or elevated ALT for NASH and fibrosis [31].

Diagnosing NAFLD/NASH begins with evaluating patients for alternative or coexisting causes of liver disease, such as viral hepatitis or significant alcohol intake, through history and laboratory testing (Table 2). The accuracy of ultrasound for the detection of moderate and severe steatosis is quite high, $>80\%$ in a meta-analysis compared to that of liver biopsy. However, ultrasound has suboptimal sensitivity for mild steatosis [32,33]. Among patients with a high pretest probability of NAFLD, moving directly to risk stratification without an ultrasound to confirm steatosis may be appropriate.

Although an optimal strategy for risk stratification of individuals with NAFLD/NASH in primary care and specialist clinics remains undefined, the guiding principle is to rule out advanced fibrosis by simple, noninvasive fibrosis scores (such as NAFLD fibrosis score or Fibrosis-4 Index). Patients at intermediate or high risk may require further assessment with a second-line test—elastography, or a serum marker test with direct measures of fibrogenesis (such as enhanced liver fibrosis [34] or fragments of propeptide of type III procollagen [35], and may require referral to a hepatology clinic (Fig. 1). Of note, the enhanced liver fibrosis and propeptide of type III procollagen tests are not approved in the United States, limiting their use in clinical practice. In contrast, elastography-based tests are available and can be used for risk stratification. Several recent studies show that this sequential use of noninvasive tests reduces unnecessary referrals to specialists, increases the detection of advanced fibrosis and cirrhosis, and hence may be cost-effective [36,37].

Once diagnosis and initial risk stratification have been completed, a more detailed assessment of liver fibrosis is essential. Accurate fibrosis staging provides information regarding prognosis, need for pharmacotherapy, intensive lifestyle modification and/or bariatric surgery, and screening/surveillance for varices and HCC. The most commonly used imaging techniques to evaluate fibrosis are vibration-controlled transient elastography and magnetic resonance elastography. Vibration-controlled transient elastography uses ultrasound waves to investigate the presence or absence of advanced fibrosis with a specificity of $92\%$ [38]. Magnetic resonance elastography can identify the intermediate stages of fibrosis more readily, but is not as widely available and is much more costly [30].

Liver biopsy, historically required to diagnosis liver fibrosis and NASH, provides helpful information and should be considered for cases in which there is a diagnostic doubt, such as patients with indeterminate, unreliable, or conflicting noninvasive assessments, or as part of phase 2 or 3 clinical trials. In addition to excluding co-existing liver diseases, liver biopsy allows for assessment of disease activity in the form of lobular and portal inflammation and ballooning degeneration (a marker of liver-cell injury). These 2 processes are thought to be responsible for triggering the development of liver fibrosis.

Assessment of cardiometabolic risk in NAFLD/NASH is also important, especially in patients who are at intermediate to high risk of advanced fibrosis [39]. The Atherosclerotic Cardiovascular Disease risk calculator has been validated in NAFLD patients and provides guidance for statin use [40].

### 4.2. Management

Most patients with NAFLD and many with NASH have a low risk of clinically significant fibrosis and can be managed by PCPs. Because NAFLD is not an isolated disease but a component of cardiometabolic abnormalities typically associated with obesity, the cornerstone of therapy is the same as that for people with obesity and cardiometabolic complications, namely lifestyle-based therapies (altered diet, such as reduced-calorie or Mediterranean diet and regular, moderate physical activity), and replacing obesogenic medications to decrease body weight and improve cardiometabolic health. The magnitude of weight loss correlates with decreases in intrahepatic triglyceride (IHTG) content, hepatocyte ballooning, and hepatic inflammation [29].

IHTG is extraordinarily sensitive to changes in energy balance; even 48 h of a low-calorie diet can decrease IHTG by about 20%, and 7% weight reduction decreases IHTG by approximately 40% [41]. The durability of these acute weight-loss–related changes remains to be determined. Furthermore, hepatic fibrosis is more resistant to weight loss and requires larger amounts ($\geq 10\%$) and possibly longer duration of weight loss to achieve clinically meaningful outcomes. Regular endurance [42,43] or resistance exercise [44] in the absence of weight loss decreases IHTG content only slightly but improves metabolic health. US Food and Drug Administration–approved weight-loss medications can enhance weight loss induced by lifestyle therapy and may contribute to the successful management of patients with NAFLD. Patients at risk of significant fibrosis (based on their clinical profile, blood test panels, and/or imaging) should be referred to a hepatologist to discuss the need for further testing, including biopsy, appropriate follow-up (particularly for those patients with advanced fibrosis/cirrhosis), and possible inclusion in NASH clinical trials (Fig. 1).

Patients with NASH and fibrosis stage 2 or higher are candidates for liver-directed pharmacotherapy (Table 3). Although there are currently no US Food and Drug Administration–approved drugs for treating NASH, vitamin E (800 IU/d) improves steatosis in NASH patients without out T2D [45]. Although randomized controlled trials have not shown similar efficacy in patients with T2D [46], one retrospective study of patients with NASH and either bridging fibrosis or cirrhosis, with or without T2D, associated vitamin E with greater transplant-free survival and lower rates of hepatic decompensation [47].

If diabetes is present, the PCP may opt to prescribe a medication for diabetes that can also treat NASH. Although metformin is first-line therapy for the pharmacologic management of T2D, it is not effective in treating NASH [2,25]. Guidelines suggest that clinicians should instead consider using pioglitazone (a thiazolidinedione acting through activation of proliferator-activated receptor–$\gamma$ and $\alpha$–agonism), based on evidence from 5 randomized controlled trials showing that it reverses steatohepatitis in patients with [48–50] and without [45,49] diabetes. In the phase 3 Pioglitazone vs Vitamin E vs Placebo for Treatment of Non-Diabetic Patients with Nonalcoholic Steatohepatitis (PIVENS) trial, pioglitazone led to resolution of steatohepatitis in 47% of patients compared with 21% of patients in the placebo group ($P = .001$; vitamin E in 36%; $P = .05$), although pioglitazone did not meet the prespecified primary end point [45]. Studies of patients with prediabetes or T2D with follow-up for up to 3 years have also consistently reported benefit with pioglitazone treatment [48–50].

Based on these data, the American Association for the Study of Liver Diseases, the European Association for the Study of the Liver, European Association for the Study of Diabetes, and European Association for the

### Table 2

<table>
<thead>
<tr>
<th>History and medical review</th>
<th>Investigations</th>
</tr>
</thead>
<tbody>
<tr>
<td>Obesity</td>
<td>Liver biochemistries (ALT, AST)</td>
</tr>
<tr>
<td>T2D</td>
<td>Exclude/identify other liver diseases*</td>
</tr>
<tr>
<td>Metabolic syndrome</td>
<td>HBV and HCV serology (and viral load)</td>
</tr>
<tr>
<td>Alcohol intake</td>
<td>Auto antibodies (ANA, AMA, ASMA)</td>
</tr>
<tr>
<td>&lt;14 drinks/wk. for women</td>
<td>Serum ferritin, A1AT</td>
</tr>
<tr>
<td>&gt;21 drinks/wk. for men</td>
<td>Liver ultrasound: increased echogenicity</td>
</tr>
<tr>
<td>No known pre-existing liver disease</td>
<td>–</td>
</tr>
</tbody>
</table>

A1AT, $\alpha$ 1 antitrypsin; AMA, antimitochondrial antibody; ANA, antinuclear antibody; ASMA, anti–smooth muscle antibody; HBV, hepatitis B virus; HCV, hepatitis C virus.

* NAFLD can coexist with other chronic liver diseases. Of note, 21% of patients with NAFLD may have elevations in autoantibodies in the absence of autoimmune hepatitis [85], and 20% may have high serum ferritin ($>300$ ng/mL in women and $>450$ ng/mL in men). Elevated serum ferritin is associated with advanced hepatic fibrosis [86] in patients with NAFLD.
Study of Obesity guidelines suggest that pioglitazone can be used for NASH patients with diabetes. The guidelines also state that vitamin E (administered at a daily dose of 800 IU) may be considered in nondiabetic adults with biopsy-proven NASH [2,30]. Pioglitazone can also reduce cardiovascular disease in patients with or without T2D, as reviewed elsewhere, although the US Food and Drug Administration has not approved it for this indication [51,52].

Several GLP-1 receptor agonists and SGLT2 inhibitors, which are increasingly used in T2D, as they reduce cardiovascular risk and promote weight loss, also potentially decrease hepatic steatosis in patients with NAFLD. GLP-1 receptor agonists (dulaglutide, exenatide, liraglutide, and semaglutide) have been tested in patients with T2D and NAFLD, with the most robust evidence to date involving semaglutide [53–57]. A small phase 2 trial (involving 52 patients) that evaluated liraglutide, a synthetic long-acting glucagon-like peptide 1 (GLP-1) receptor agonist available for treating T2D and obesity, resulted in weight loss, resolution of steatohepatitis, and slower progression of fibrosis than placebo, although gastrointestinal adverse effects were common [56].

More recently, a report in 320 patients with biopsy-proven NAFLD offers the strongest evidence for the use of GLP-1 receptor agonists in patients with NAFLD. GLP-1 receptor agonist (dulaglutide, exenatide, liraglutide, and semaglutide) receptor agonist available for treating T2D and obesity, resulted in weight loss, resolution of steatohepatitis, and slower progression of fibrosis than placebo, although gastrointestinal adverse effects were common [56].

Another small recent study found that dulaglutide also reduced liver fat content and transaminases in people with T2D and NAFLD [55]. These findings allow the possibility of treating diabetes, cardiovascular disease, and NASH simultaneously with diabetes medications, such as pioglitazone or a GLP-1 receptor agonist. SGLT-2 inhibitors (canagliflozin, dapagliflozin, empagliflozin, and ertugliflozin) have also been tested in NAFLD, but these studies have been small and do not examine the effect of these agents on liver histology [58].

Despite the promise of antidiabetes medications, the role of improving glycemic control on the natural history of NASH and development of NAFLD/NASH. FIB-4, Fibrosis-4 Index; NFS, NAFLD fibrosis score.

Table 3
Management of patients with nonalcoholic fatty liver disease and nonalcoholic steatohepatitis.

<table>
<thead>
<tr>
<th>Variable</th>
<th>Lifestyle interventiona</th>
<th>Liver-directed pharmacotherapy</th>
<th>Diabetes care (in individuals with diabetes)</th>
<th>Cardiovascular risk reduction</th>
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<tbody>
<tr>
<td>NAFL</td>
<td>Yes</td>
<td>No</td>
<td>Standard of care</td>
<td>Yes</td>
</tr>
<tr>
<td>NASH with fibrosis stage 0 or 1 (F0, F1)</td>
<td>Yes</td>
<td>No</td>
<td>Standard of care</td>
<td>Yes</td>
</tr>
<tr>
<td>NASH with fibrosis stage 2 or 3 (F2, F3)</td>
<td>Yes</td>
<td>Yes</td>
<td>Pioglitazone, GLP-1 receptor agonistsb</td>
<td>Yes</td>
</tr>
<tr>
<td>NASH cirrhosis (F4)</td>
<td>Yes</td>
<td>Yes</td>
<td>Individualizationc</td>
<td>Yes</td>
</tr>
</tbody>
</table>

a All patients require regular physical activity and healthy diet and to avoid excess alcohol intake; weight loss recommended.

b Among GLP-1 receptor agonists, semaglutide has the best evidence of benefit in patients with NASH and fibrosis.

c Evidence for efficacy of pharmacotherapy in patients with NASH cirrhosis is very limited and should be individualized and used with caution.
cirrhosis remains poorly understood, and the role of glycemic control of disease progression in NASH remains to be established. Improving glycemic control with insulin therapy reduces liver steatosis [48], but its impact on liver histology (both NASH or fibrosis) and the natural history of the disease remain unknown. Cross-sectional [59] and longitudinal observational studies [60] do not show a clear correlation between hemoglobin A1c levels over time and liver histology or other clinical outcomes. Lowering hemoglobin A1c levels with pioglitazone treatment for 18 months has been associated with improvement in NASH and slower progression of fibrosis compared to patients with diabetes on placebo but, overall, the histologic response to pioglitazone does not appear to be linked to improved glycemic control, as it is similar in patients with vs those without diabetes [49].

Bariatric surgery is currently the most effective therapy available for obesity. The 2 most common procedures are sleeve gastrectomy and Roux-en-Y gastric bypass. Marked weight loss (approximately 25%–35%) induced by bariatric surgery has profound effects on steatosis, NAFLD activity score, hepatocyte ballooning, and lobular inflammation, and results in NASH resolution in most patients [61,62]. Surgery-induced weight loss also has a considerable therapeutic effect in reducing stages 1 and 2 fibrosis, but is less effective in improving stages 3 and 4. Bariatric endoscopy is emerging as a new treatment for obesity, but the long-term durability of its effects remains to be determined. About 15% weight loss has been reported after therapy with a postprandial gastric aspiration device, which is associated with reduced plasma AST and ALT [63], whereas duodenal mucosal resurfacing has reduced Fibrosis-4 Index scores by mechanisms possibly unrelated to weight reduction [64]. Intragastric balloon placement has also been associated with histologic improvement in individuals with NASH [65], although findings remain preliminary. Patients with advanced liver disease, especially with hepatic decompensation, have higher mortality after bariatric surgery. Overall, more efficacy and safety data are needed before these approaches can be recommended as treatment options for patients with NAFLD and NASH.

Special attention to the management of sedentary behavior, as well as to dyslipidemia, diabetes, and hypertension, is recommended for all individuals with NAFLD [66]. Alcohol consumption should be limited to 2–3 drinks per week in women and 4–5 drinks per week in men and avoided in patients with advanced fibrosis [67,68], although high-quality data on the exact risk of progressive liver disease in patients with advanced fibrosis are still needed. Many PCPs and nonhepatologists discontinue statins when liver enzymes are elevated [9,69,70]. However, numerous studies have also demonstrated that statins are safe and efficacious in patients with NAFLD and NASH, and they can be used to treat dyslipidemia in these patients, including those with compensated cirrhosis. Statins have pleiotropic properties that may be directly beneficial in liver disease. In a meta-analysis of 13 studies, including 3 randomized controlled trials, statin use in cirrhosis was associated with a reduction in hepatic decompensation (hazard ratio, RR, 0.54; 95% CI, 0.46–0.62) and lower mortality (hazard ratio, 0.54; 95% CI, 0.47–0.61) [71]. However, because data remain limited regarding safety and risks of statins in patients with compensated cirrhosis [72,73], statins should be avoided until we have stronger evidence to support their safety in these patients. The AGA clinical practice update provides some guidance and advises against statin use among patients with Child-Pugh class B or C cirrhosis [74]. The underlying rationale is that the generally grave liver-related prognosis of patients with Child-Pugh class B or C cirrhosis makes it unlikely that they will benefit from the cardiovascular benefits associated with lipid-lowering therapy. In a large retrospective cohort study of statins in patients with cirrhosis, the survival benefit did not extend to patients with Child class C cirrhosis [75].

4.3. Emerging tools

Given the high prevalence of NAFLD and the limited patient awareness about this disease, applying artificial intelligence/machine learning tools to the big data repositories of electronic health records holds considerable potential for efficient disease identification and risk stratification [76]. “Machine learning” is a subset of artificial intelligence in which computer algorithms are improved through experience [77]. These tools can produce noninvasive calculated scores by using information about patient demographic and clinical characteristics from both narrative (i.e., free text) and codified (e.g., administrative disease codes and laboratory tests) sources. Artificial intelligence is also being tested to improve the accuracy and reliability of liver histologic interpretation using quantitative scoring systems for NAFLD/NASH radiologic and histopathologic features [78,79]. However, although the availability of noninvasive tests to accurately assess response to treatment beyond histopathology would greatly facilitate the efficient enrollment in NASH treatment clinical trials, existing options still require further validation and eventual acceptance by regulatory agencies.

Several liver-targeted and other potential therapies are also currently under investigation, targeting a broad range of pathologic changes associated with NASH, including insulin resistance, alterations in the microbiome and gut permeability, oxidative stress, apoptosis, lipotoxicity, inflammation, and bile acid metabolism. Given the multiple pathways involved in NASH pathogenesis, combination regimens may ultimately be needed to treat NASH most effectively [80,81].

5. Recommendations

5.1. Develop more sensitive and specific diagnostic methods

The invasive nature and relatively high expense of liver biopsy limit its use and call for more sensitive and specific noninvasive diagnostic methods for NASH. Several novel noninvasive tools with the potential to provide more sensitive and specific diagnosis are currently under development. These include top-down approaches, such as multiotics and narrowing down to the minimum number of molecules that could provide the maximum positive and negative predictive value [82,83].

5.2. Adopt a multidisciplinary approach to nonalcoholic steatohepatitis

Optimal care of patients with NASH may require clinicians from a variety of specialties, including primary care, hepatology, obesity management, and endocrinology, to tackle both the hepatic manifestations of the disease and the comorbid metabolic syndrome and cardiovascular risk, as well as screening and treating other comorbid conditions (e.g., obstructive sleep apnea). When NAFLD progresses to NASH, multidisciplinary, team-based care involving these specialties is crucial. Improving the traditional model of primary, secondary, and tertiary care will require not only developing and validating algorithmic approaches (e.g., who can be managed where and how), but also connectivity and multidirectional referrals among these practice settings. Examining other models of care, such as medical homes either dedicated to NAFLD/NASH or incorporated within similar homes that manage metabolic disease more broadly, could also be valuable in developing care models. These integrated models can create and align expertise and incentives among different specialties.

5.3. Develop clinical care pathways

Developing clinical care pathways that use validated and efficient noninvasive tests and calculators is crucial to a multidisciplinary approach to managing NAFLD/NASH. Clinical care pathways, with careful explication of each step-in screening, diagnosis, and treatment, have been shown to improve the quality of health care delivery in other areas of medicine. Members of the NASH: A Call-to-Action Steering Committee and several other conference participants are currently developing such a pathway for NAFLD/NASH. Rapid and timely dissemination of these pathways to all stakeholders, especially the frontline PCPs,
will be important in developing a systematic approach to managing NAFLD/NASH.

5.4. Pursue a unified, international, public health response

The public health response to NAFLD remains rudimentary. There is no single guiding strategy in the United States or Europe. A survey of 29 European countries highlighted the absence of a concrete NAFLD/NASH management strategy or action plans in every one of these countries [84]. This deficit has even more proximal roots. For example, not all hepatology/gastroenterology societies have clear screening, testing, or referral guidelines for NAFLD/NASH, and existing guidelines often conflict with one another (Table 4). Intersociety collaboration for harmonizing guidelines to optimize screening, diagnosis, and therapy is urgently required. Furthermore, because virtually all current guidance regarding HCC surveillance in NAFLD is derived from the viral hepatitis and alcoholic cirrhosis literature, new data and updated guidelines are needed that are specific to NAFLD/NASH-related cirrhosis. In addition, large cohorts with longitudinal data on clinical course and outcomes, particularly cohorts that allow the transition from childhood through adolescence to adulthood to be evaluated, are needed to inform the science and clinical practice of managing NAFLD/NASH.

There is also a large unmet need for programs that can increase disease awareness in the medical community and the general population. Finally, the closely interlinked nature with related metabolic diseases suggests that reducing the clinical and economic burden of NASH and NAFLD will require fundamental societal changes driven by policies to address failing public health systems and the social determinants of health.

6. Summary and conclusions

The upward trend in NAFLD/NASH incidence and prevalence underscores the importance and urgency of developing and implementing effective screening, diagnosis, and treatment strategies in the United States and globally, particularly among emerging at-risk cohorts, such as patients with diabetes and obesity. This goal cannot be achieved if the different specialties engaged in managing this burgeoning population continue to work in separate silos. The Call-to-Action Meeting described in this report represents one of the first steps needed to align key stakeholders, including PCPs, endocrinologists, diabetologists, obesity medicine specialists, gastroenterologists, and hepatologists, on a collective action plan. Improving the spectrum of care for patients with NAFLD from screening, diagnosis, disease severity stratification, and treatment will require significant changes and innovations in technology, health care delivery, and policy. In addition, optimal care of patients with NAFLD/NASH will require a multidisciplinary team integrating primary care, hepatology, obesity medicine, and endocrinology/diabetology via well-defined care pathways, along with exploration of the high-yield targets for clinical research and practice identified by conference participants. These efforts should help the field move toward a collective strategy with shared goals and objectives that will improve care for the growing population of patients with NAFLD/NASH.

Declaration of competing interest

These authors disclose the following: Jay H. Shubrook has served as an advisor to Sanofi, Eli Lilly, Novo Nordisk, Bayer, and MannKind. Elisabetta Bugianesi has served as a consultant to Gilead, BMS, Boehringer, Intercept, and Innova. Zobair Younossi has received research funding from or served as a consultant for Gilead Sciences, Intercept, BMS, Novo Nordisk, Viking, Terns, Siemens, Shionogi, AbbVie, Madrigal, Merck, Abbott, Alexcell, and Novartis. Yamini Natarajan has received grants from Gilead and Allergan. Mary E. Rinella has served as a consultant for Amgen, Atyllym, Allergan, BMS, Boehringer Ingelheim, Cohersus, CymaBay, Enanta, Fibrogen, Fractyl, Galecto, Gelesis, Genentech, GENFIT, Gilead, Intercept, Lipocine, Madrigal, Merck, Metacrine, NGM Biopharmaceuticals, Novo Nordisk, Novartis, Pfizer, Sagimet, Siemens, Takeda, Terns, Tietis, Viking, 3Bio, 89Bio. Robert H. Eckel has served on advisory boards for Nordo Novik, Prevention Bio, Kaleido, and KOWA, and a scientific advisory committee for PROMINENT (CVOT). Stephen A. Harrison has served as an advisory board/consultant for Akero, Altimune, Arrowhead, Alexcell, B. Riley, Boston Pharma, Cirus, Civi Biotherma, CLDF, Coerper, CymaBay, Echoson, Fibronostics, Foresite Labs, Fortress, Galectin, Gelesis, GENFIT, Gilead, Heipion, Hepion, Hightide Bio, Histolindex, Intercept, Inipharm, Ionis, Kowa, Madrigal, Medpace, Metacrine, Microba, NGM Bio, NorthSea, Novartis, Novo Nordisk, Piper Sandler, Poxel, Prometic, Ridgeline Therapeutics, Sagimet, Sonic Incytes, Terns, Theratech, Viking, 89 Bio; received grant/research support from Alexcell, BMS, Cirus, Civi Biotherma, Conatus, CymaBay, Enyo, Galectin, Calmed, Genentech, GENFIT, Gilead, Heipion, Hightide Bio, Immuron, Intercept, Madrigal, NGM Bio, NorthSea, Novartis, Novo Nordik, Pfizer, Sagimet, Second Genome, Tobira/Allergan, Viking; and has stock/shares (self-managed) in Akero, Cirus, Galectin, GENFIT, Heipion, Histolindex, Metacrine, NGM Bio, and NorthSea. Christos Mantzoros reports grants, personal fees, and other from Coherus Biosciences, grants, personal fees and other from Novo Nordisk, personal fees and nonfinancial support from Ansh, Aegerion, PES, California Walnut Commission, and personal fees from GENFIT, Intercept, Regeneron, CardioMetabolic Health Conference and The Metabolic Institute of America, and Amgen. Samuel Klein is a shareholder of Aspire Bariatrics and has served as a consultant for Pfizer, Novo Nordisk, and Boehringer Ingelheim. Robert H. Eckel has received research funding from Gilead (2014–2015), Merck (2016–2018), and Wallo (2014–2017). Kenneth Cusi has received research support as principal investigator for the University of Florida from Cirus, Echosong, Inventiva, Novartis, Novo Nordisk, Poxel, and Zydus, and is a consultant for Allergan, Astra-Zeneca, Alexcell, BMS,

Table 4  Summary of published nonalcoholic fatty liver disease guidelines.

<table>
<thead>
<tr>
<th>Organization</th>
<th>Year</th>
<th>First-line diagnosis test</th>
<th>When to refer to hepatologist</th>
<th>Noninvasive tests</th>
</tr>
</thead>
<tbody>
<tr>
<td>American Association for the Study of Liver Diseases (AASLD)</td>
<td>2018</td>
<td>Not clear in the guideline</td>
<td>Routine screening for NAFLD in high-risk groups is not recommended</td>
<td>Diagnosis for NASH: liver biopsy Assessment for fibrosis: NFS or FIB-4</td>
</tr>
<tr>
<td>American Gastroenterological Association (AGA)</td>
<td>2012</td>
<td>Routine screening for NAFLD is not recommended</td>
<td>Ultrasound + liver enzymes for patients with risk factors</td>
<td>Metabolic syndrome can be used to target patients for liver biopsy Diagnosis for NASH: liver biopsy Assessment for fibrosis: NFS or FIB-4</td>
</tr>
<tr>
<td>European Association for the Study of the Liver (EASL)</td>
<td>2010</td>
<td>Ultrasound + liver enzymes for patients with risk factors</td>
<td>Refer patients with abnormal liver enzymes or medium—high-risk fibrosis markers to specialist</td>
<td>Not clear in the guideline</td>
</tr>
<tr>
<td>World Gastroenterology Organization (WGO)</td>
<td>2012</td>
<td>Ultrasound + liver enzymes for patients with risk factors</td>
<td>Refer adults with advanced liver fibrosis to a hepatologist</td>
<td>Assessment for advanced fibrosis: enhanced liver fibrosis (every 2–3 y)</td>
</tr>
<tr>
<td>National Institute for Health Care and Excellence (NICE)</td>
<td>2016</td>
<td>Ultrasound + liver enzymes for patients with risk factors</td>
<td>But routine liver function blood tests are not sensitive, and ultrasound is not cost-effective</td>
<td>Assessment for advanced fibrosis: enhanced liver fibrosis (every 2–3 y)</td>
</tr>
</tbody>
</table>

FIB-4, Fibrosis-4 Index; NFS, NAFLD fibrosis score.
Acknowledgments

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References

Appendix A. Supplementary materials

Detailed results from the NASH Needs Assessment Survey are available at: https://nash.gastro.org/survey.

Participants of “Preparing for the NASH Epidemic: A Call to Action” Initiative

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<table>
<thead>
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<th>Affiliation</th>
<th>Specialty</th>
</tr>
</thead>
<tbody>
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</tr>
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<td>Gastroenterology</td>
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<td>Endocrinology</td>
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<td>Primary care</td>
</tr>
<tr>
<td>Zobair M. Younossi, MD</td>
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<td>Hepatology</td>
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</tbody>
</table>

Nonalcoholic Fatty Liver Disease and Nonalcoholic Steatohepatitis Survey

Thank you for taking the time to participate in this study. The purpose of the study is to determine the current level of physician awareness, familiarity, and practices in the diagnosis and management of nonalcoholic fatty liver disease (NAFLD) and nonalcoholic steatohepatitis (NASH).

NOTE: Text in red indicates correct answer for reference only—no programming implications.

A. Awareness

Q1. To the best of your knowledge, which of the following statements accurately defines nonalcoholic fatty liver disease (NAFLD)? (Please check one)

- Evidence of hepatic steatosis
- Evidence of hepatic steatosis and lack of secondary causes of hepatic fat accumulation
- Evidence of hepatic steatosis with secondary causes of hepatic fat accumulation
- Not sure, would like to receive more information

Q2. To the best of your knowledge, which of the following statements accurately defines nonalcoholic fatty liver (NAFL)? (Please check one)

- Presence of ≥5% hepatic steatosis
- Presence of ≥5% hepatic steatosis with hepatocellular injury
- Presence of ≥5% hepatic steatosis without hepatocellular injury
- Not sure, would like to receive more information
Q3. To the best of your knowledge, which of the following statements accurately defines nonalcoholic steatohepatitis (NASH)? (Please check one)

- Presence of ≥5% hepatic steatosis
- Presence of ≥5% hepatic steatosis with hepatocellular injury
- Presence of ≥5% hepatic steatosis without hepatocellular injury
- Not sure, would like to receive more information

Q4. Roughly what proportion of the following patient groups are likely to have NAFLD? If you are not comfortable answering a question based on your current role or the information you have received to date, please select not sure/need more information.

<table>
<thead>
<tr>
<th>Patients with severe obesity</th>
<th>Almost all</th>
<th>About half</th>
<th>Up to one-quarter</th>
<th>Not sure, would like to receive more information</th>
</tr>
</thead>
<tbody>
<tr>
<td>Patients with type 2 diabetes mellitus</td>
<td>X</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Patients with dyslipidemia</td>
<td>X</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>General population</td>
<td>X</td>
<td></td>
<td></td>
<td></td>
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</table>

Q5. Which one of the following statements is true?

- Men are twice as likely as women to have NAFLD
- Men and women are almost equally likely to have NAFLD
- Women are twice as likely as men to have NAFLD
- Not sure, need more information

Q6. Which of the following is the best estimate of the prevalence rate of NAFLD in patients with type 2 diabetes based on liver ultrasound?

- Less than 25%
- Approximately 55%
- Approximately 75%
- Not sure, need more information

Q7. Statements below describe some potential adverse outcomes that patients with NAFLD or histological NASH may experience. Please indicate whether you believe each statement is true or false based on current evidence. If you are not comfortable answering a question based on your current role or the information you have received to date, please select not sure/need more information.

<table>
<thead>
<tr>
<th>Patients with NAFLD have increased overall mortality compared to matched control populations without NAFLD.</th>
<th>True</th>
<th>False</th>
</tr>
</thead>
<tbody>
<tr>
<td>The most common cause of death in patients with NAFLD is cardiovascular disease, independent of other metabolic comorbidities.</td>
<td>X</td>
<td></td>
</tr>
<tr>
<td>Although liver-related mortality is the 12th leading cause of death in the general population, it is the second or third cause of death among patients with NAFLD.</td>
<td>X</td>
<td></td>
</tr>
<tr>
<td>Cancer-related mortality is among the top 3 causes of death in subjects with NAFLD.</td>
<td>X</td>
<td></td>
</tr>
<tr>
<td>The most important histological feature of NAFLD associated with long-term mortality is advanced fibrosis or cirrhosis.</td>
<td>X</td>
<td></td>
</tr>
</tbody>
</table>

B. Screening and Patient Management

Q8a. Do you screen for NAFLD and/or NASH?

- Yes
- No [GO TO 8b]

Q8b. What are some reasons why you don’t screen for NASH and/or NAFLD? (Please check all that apply)

- I am not familiar with screening procedures for NAFLD/NASH
- NAFLD/NASH are not a priority in my practice
- I do not have time to screen for NAFLD/NASH
- Treatment therapies for NAFLD/NASH are limited
- Prevalence of NAFLD/NASH is low
- NAFLD/NASH is not my specialty
- Other (please specify) __________________

Q9. Do you diagnose NAFLD and/or NASH?

- Yes
- No [GO TO 9a]
Q9a. What are some reasons why you don’t diagnose NASH and/or NAFLD? (Please check all that apply)

- I am not familiar with diagnostic procedures for NAFLD/NASH
- NAFLD/NASH is not my specialty
- Diagnostic procedures are invasive and risky
- Treatment therapies for NAFLD/NASH are limited
- Other (please specify) _______________

Q10. Number of patients with NAFLD seen monthly?

- None
- <5
- 5–10
- 11–20
- >20

Q11. Do you currently manage patients with NAFLD and/or NASH?

- Yes
- No [GO TO Q12]

Q12. IF Q11 = NO: Will you manage new NAFLD and/or NASH Patients?

- Yes
- No, will refer all new patients [GO TO Q12a]

Q12a. What are some reasons why you won’t manage NAFLD/NASH patients? (Please check all that apply)

- I’m not familiar with treatment therapies for NAFLD/NASH
- NAFLD/NASH is not my specialty
- I do not have time to manage patients with NAFLD/NASH
- Other (please specify) ____________________________

C. Diagnosis

This section of the survey asks about NAFLD and NASH diagnosis. We would like to understand your opinions about appropriate clinical practice, even if you are not always the one to actually screen and diagnose patients due to patient referral practices. If you are not comfortable answering a question based on your current role or the information you have received to date, please select not sure/need more information.

Q13. The statements below describe some possible patient groups that should be screened for NAFLD. Please indicate whether you believe each statement is true or false. If you are not comfortable answering a question based on your current role or the information you have received to date, please select not sure/need more information.

<table>
<thead>
<tr>
<th>Statement</th>
<th>True</th>
<th>False</th>
<th>Not sure, need more information</th>
</tr>
</thead>
<tbody>
<tr>
<td>Patients with abnormal liver chemistries</td>
<td>X</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Patients older than 50 years who have hypertension and hyperlipidemia</td>
<td>X</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Patients with type 2 diabetes</td>
<td>X</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Patients with NAFLD family members</td>
<td>X</td>
<td></td>
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</table>

Q14. The statements below describe how to approach an initial evaluation (clinical history, laboratory testing, imaging for confirmation of diagnosis and risk stratification) of the patient with suspected NAFLD. Please indicate whether you believe each statement is true or false. If you are not comfortable answering a question based on your current role or the information you have received to date, please select not sure/need more information.

<table>
<thead>
<tr>
<th>Statement</th>
<th>True</th>
<th>False</th>
<th>Not sure, need more information</th>
</tr>
</thead>
<tbody>
<tr>
<td>When evaluating a patient with suspected NAFLD, it is important to exclude competing etiologies for steatosis and coexisting common chronic liver disease.</td>
<td>X</td>
<td></td>
<td></td>
</tr>
<tr>
<td>In patients with suspected NAFLD, persistently high serum ferritin, and increased iron saturation, especially in the context of homozygote and heterozygote C282Y HFE mutation, a liver biopsy should be considered.</td>
<td>X</td>
<td></td>
<td></td>
</tr>
<tr>
<td>High serum titers of autoantibodies in association with &gt;5 upper limit of normal aminotransferases, high globulins, or high total protein to albumin ratio should prompt a workup for autoimmune liver disease.</td>
<td>X</td>
<td></td>
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<tr>
<td>Initial evaluation of patients with suspected NAFLD should include cross-sectional abdominal imaging (such as contrast-enhanced computed tomography scan) to screen for hepatocellular cancer.</td>
<td>X</td>
<td></td>
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</tbody>
</table>
True False Not sure, need more information

Initial evaluation of patients with suspected NAFLD should carefully consider the presence of commonly associated comorbidities such as obesity, dyslipidemia, insulin resistance or diabetes, hypothyroidism, polycystic ovary syndrome, and sleep apnea.

X

Q15. The statements below describe how to approach a noninvasive diagnosis of steatohepatitis and advanced fibrosis in NAFLD. Please indicate whether you believe each statement is true or false. If you are not comfortable answering a question based on your current role or the information you have received to date, please select not sure/need more information.

True False Not sure, need more information

NAFLD fibrosis score (NFS) or Fibrosis-4 Index are clinically useful tools for identifying NAFLD patients with higher likelihood of having advanced fibrosis (stage 2 or higher) or cirrhosis (stage 4).

X

Abdominal ultrasound is a clinically useful tool for identifying NAFLD patients with steatohepatitis.

X

Vibration-controlled transient elastography (FibroScan) or magnetic resonance elastography (imaging) are clinically useful tools for identifying advanced fibrosis in patients with NAFLD.

X

Q16. Please indicate situations when you will consider obtaining a liver biopsy in patients with NAFLD. If you are not comfortable answering a question based on your current role or the information you have received to date, please select not sure/need more information.

Yes No Not sure, need more information

Patients with suspected NAFLD, mildly elevated serum ferritin, and severe obesity

X

The presence of metabolic syndrome, NAFLD fibrosis score or Fibrosis-4 Index, or liver stiffness measured by vibration-controlled transient elastography or magnetic resonance elastography (imaging) suggestive of moderate-to-severe fibrosis

X

Patients with suspected NAFLD in whom competing etiologies for hepatic steatosis and the presence and/or severity of coexisting chronic liver diseases cannot be excluded

X

Q17. Clinical pathology reporting should include which of the following? (Check all that apply)

- Distinction between NAFL (steatosis), NAFL with inflammation, and NASH (steatosis with lobular and portal inflammation and hepatocellular ballooning)
- Commentary on severity (mild, moderate, or severe)
- Use of specific scoring systems, such as NAFLD activity score and/or steatosis, activity, and fibrosis, if deemed appropriate
- Description of the presence of fibrosis, including location, amount, and parenchymal remodeling, if warranted
- All of the above
- Not sure, need more information

D. Patient Management Practices

This section of the survey asks about NAFLD and NASH management and treatment. We would like to understand your opinions about appropriate clinical practice, even if you are not always the one to manage patients and their treatment due to patient referral practices. If you are not comfortable answering a question based on your current role or the information you have received to date, please select not sure/need more information.

Q18. Clinicians have different options about what NASH treatments are appropriate, or if a treatment requires confirmation of NASH first via liver biopsy. Indicate your opinion for each treatment in the grid below. If you are not comfortable answering a question based on your current role or the information you have received to date, please select not sure/need more information. (X – marked per Guidance)

Use without liver biopsy Recommended but only after liver biopsy Not recommended per current practice guidelines Not sure, need more information

Foregut bariatric surgery for otherwise eligible individuals with obesity

X

GLP-1 agonists

X

Metformin

X

Obeticholic acid

X

Omega-3 fatty acids

X

Pioglitazone

X

Ursodeoxycholic acid

X

Weight loss of 7%-10%

X

Vitamin E for nondiabetic adults

X

Vitamin E for diabetic patients

X

Q19. The statements below describe guidance for managing patients with NAFLD or NASH. Please indicate whether you believe each statement is true or false based on current evidence. If you are not comfortable answering a question based on your current role or the information you have received to date, please select not sure/need more information.
NAFLD and NASH patients should avoid heavy alcohol consumption X
Aggressive modification of CVD risk factors should be considered in all patients with NAFLD or NASH X
Statins can be used to treat dyslipidemia in patients with NAFLD or NASH X
Statins can only be used to treat dyslipidemia in patients with NASH X
Statins should be avoided in patients with decompensated cirrhosis X

Q20. The statements below describe guidance for managing patients with NASH. Please indicate whether you believe each statement is true or false based on current evidence. If you are not comfortable answering a question based on your current role or the information you have received to date, please select not sure/need more information.

Patients with NASH cirrhosis should be screened for gastroesophageal varices X
Patients with cirrhosis suspected because of NASH should be considered for hepatocellular carcinoma (HCC) screening X
Routine screening and surveillance should be conducted for hepatocellular carcinoma (HCC) in patients with noncirrhotic NASH X
Liver biopsy should be repeated in patients with NAFL or NASH X
Repeat liver biopsy in NAFL or NASH should be considered on a case-by-case basis X

Q21. Please indicate your level of agreement with the following statements.

NASH should not be ignored because it will impact our health system dramatically in the next few years.
NASH is a frustrating condition because there are no US Food and Drug Administration–approved treatments.
Once effective therapeutic treatments are found, I will be more likely to screen for NASH.
Once effective therapeutic treatments are found, I will be more likely to diagnose NASH.
Once effective therapeutic treatments are found, I will be more likely to treat NASH.
I would like more education on screening for NASH.
I would like more education on diagnosing NASH.
I would like more education on treating NASH.

E. Information Preference and Demographics

Q21. What is your preferred source of information about practice guidance for the diagnosis and management of NAFLD/NASH?

- American Gastroenterological Association
- My own association
- Other (please specify) ____________

Q22. What is your preferred method of receiving more information about NAFLD/NASH? (Please rank the following options in order of preference from 1 to 3, where 1 is most preferred option and 3 is least preferred option.)

- Peer-reviewed journal article
- Online medical education
- Live medical education

Q23. Location of practice (Please check one):

- Solo private practice
- Medical group practice
- Hospital-based practice
- Clinic

Q24. Size of practice (Please check one):

- Solo
- 2–5 physicians
- 6–30 physicians
- 31–100 physicians
- 101 or more physicians