Non-alcoholic fatty liver diseases: current challenges and future directions

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Abstract: Non-alcoholic fatty liver diseases (NAFLD) is rapidly becoming the most common cause of chronic liver disease in Western Countries, and a similar trend is expected in Eastern Countries within the next years. This review focusses on the definition of NAFLD and NASH, possible screening mechanisms and the question who should be screened. Still there is a need for non-invasive diagnostic tools and biomarkers for NASH that can quickly and easily diagnose the severity of NAFLD, monitor liver changes, and identify high risk patients. In addition, treatment strategies are discussed as well as the clientele, who should be treated. There are currently no drugs approved for NAFLD. Successful clinical studies with e.g., obeticholic acid and new substances (e.g., cenicriviroc with anti-inflammatory activity) have already been published. If weight-reducing diets and a change in lifestyle fail in the case of severe obesity, bariatric surgery (e.g., gastric bypass or stomach reduction) should be considered. In the case of manifest type 2 diabetes, metformin can be used as an oral antidiabetic of first choice, and GLP-1 agonists have shown beneficial effects on NAFLD. However, up to now the prevention of overweight and lack of exercise targets the most important risk factors. This review aims to identify therapy relevant risk factors, management strategies, and open questions concerning NAFLD patients.

Keywords: Non-alcoholic fatty liver diseases (NAFLD); NASH; liver; diabetes; metabolic

Submitted May 06, 2020. Accepted for publication Aug 17, 2020.

doi: 10.21037/atm-20-3760

View this article at: http://dx.doi.org/10.21037/atm-20-3760

“An oversized goose liver, no matter how good it tastes, always requires a sick goose.” (Clemens Brentano, 1778–1842).

Introduction

Non-alcoholic fatty liver diseases (NAFLD) affect about a quarter of the total population in industrialized countries. NAFLD is closely related to demographic changes (aging of society), unhealthy lifestyle (overeating, lack of exercise) and cardiovascular-metabolic diseases (type 2 diabetes, arterial hypertension, dyslipidemia). Very fast, NAFLD became the most common cause of chronic liver disease in Western Countries. We expect a similar trend in Eastern Countries over the next few years (1,2). The SHIP study (Study of Health in Pomerania) calculated a NAFLD prevalence of approximately 30% for the age group between 20 and 79 years (3). The range of NAFLD includes non-alcoholic pure (bland) fatty liver (NAFL), steatohepatitis (so called NASH), NASH fibrosis, and liver cirrhosis (NASH cirrhosis) with its associated complications.

By now NAFLD has an estimated prevalence in the Western world of around 25% (4). About 20% of these patients experience NASH (5). The progression of NASH is linked to liver cell stress, consecutive inflammation and fibrosis. Furthermore, the development of cirrhosis with portal hypertension and end stage liver disease is possible. In addition, NASH is an important risk factor with regard

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to hepatocellular carcinoma (HCC). End-stage NASH-associated liver disease is expected to represent the highest proportion of patients who will be listed for liver transplant in the future (6). The disease—although genetic factors have also been identified (7)—is seen as the result of hyperalimentation and as a hepatic manifestation of the so-called metabolic syndrome (8).

In addition to the exclusion of other specific liver diseases, typical clinical, imaging and/or histological characteristics are required for the correct diagnosis. The latter include steatosis, hepatocellular ballooning, lobular inflammation and fibrosis (9). The clinical symptoms of non-cirrhotic NAFLD are usually non-specific. Awareness of the disease and the associated risks are also low (10). The pathogenesis and the natural course of NAFLD are becoming increasingly better understood, even if the heterogeneity of the patients and the multifactorial genesis make it difficult to estimate the individual prognosis. The risk of progression to complications of liver disease (cirrhosis, portal hypertension, liver cancer) is strongly related to the degree of fibrosis. In several prospective observational studies, the degree of fibrosis was associated with total and liver-related lethality (11,12).

A further increase in NAFLD in the sense of the epidemic of obesity, especially among adolescents and younger patients, can be expected. NAFL is present in 15–20% of the normal population, NASH is observed in 2–3% (13). The figures for obesity and diabetes are correspondingly higher: 50–75% have NAFL, 20–50% have NASH and 2–3% already have cirrhosis of the liver (14). Overall and liver related mortality are shown in Figure 1.

According to the study “The heavy burden of obesity-The Economics of prevention” in 34 OECD countries, more than half of the population are already overweight, and one in four is pathologically overweight (15). Between 2010 and 2016, the proportion of obese adults in the OECD countries rose from 21% to 24%—this corresponds to 50 million people. Just under one in four adults in Germany is obese, in Austria and Switzerland about one in five. Overweight children are particularly hard hit: they do worse at school, have higher absenteeism and less chances of a higher education. They are also bullied up to three times more often than non-obese, which may contribute to their poor school performance (15).

The clinical implications of hepatic steatosis run the risk of being underestimated in several ways. This concerns both their possible causes and the variety of the disease spectrum, the (differential) diagnostic pitfalls, the possible prognostic implications as well as new therapy options and perspectives that might be necessary to manage this widespread disease.

Changes in lifestyle, demographic change and increasing complexity of pharmacological therapies lead to an increase in NAFLD. Doctors and patient organizations have to deal with this collectively and individually. The first S2k guideline for NAFLD at all, the German guideline for NAFLD from 2015 expired in February 2020 and is currently being updated (16). Still there exist a lot of open questions, especially concerning drug therapy and screening. The aim of this review is to better cope with the increased challenges to get on with NAFLD, that is identifying therapy relevant risk factors, management strategies, and open questions.

**What is the optimal definition of NAFLD and NASH?**

NAFLD covers a spectrum of liver diseases, which are characterized by increased fat storage in liver cells. The illness includes sole steatosis (NAFL, that is NAFLD), as well as NAFLD hepatitis (nonalcoholic steatohepatitis, NASH), (advanced) NASH fibrosis, cirrhosis, and HCC (17). Between pediatric and adult NAFLD patients there are specific differences in etiology, epidemiology and pathology (18). Diagnostic criteria for NAFLD according to current guidelines include hepatic steatosis verified by imaging or histology (>5%), the exclusion of other causes of fatty liver including alcohol, and in some cases the
existence of insulin resistance (16,19-21). The commonly used acronym is NAFLD (from nonalcoholic fatty liver disease).

Fatty liver or fatty liver hepatitis can be triggered by numerous diseases or causes, e.g., alcoholic liver disease, medication or chemotherapy, hepatitis C virus infection, fat metabolism disorders or malnutrition. These differential diagnoses should be taken into account as there are specific therapeutic consequences for these different etiologies (16).

Alcoholic fatty liver (AFL) and alcoholic fatty liver hepatitis (ASH) are caused by chronic alcohol abuse. In this context heavy alcohol use implies >3 drinks per day in men and >2 drinks in women according to the recent ACG guideline (22). In the US, a beverage containing about 14 g of alcohol equates to one drink, which correlates to 12 ounces of beer (5% weight/volume) or 5 ounces of wine (8–10% weight/volume), or 1.5 ounces of hard liquor (40–45% weight/volume) (22).

According to current guidelines NAFLD is defined as steatotic hepatitis without significant alcohol consumption and without causes of secondary steatosis. By definition, NAFL and NASH are not caused by alcohol consumption. However, since only about 8 percent of the population between the ages of 18 and 59 live abstinent from alcohol (23), the diagnosis of NAFL and NASH depends on the level of alcohol consumption. In the literature, the threshold of significant alcohol consumption is assumed to be 20–40 g per day in men and 10–30 g per day in women (24). The current guidelines of EASL and AASLD assume significant alcohol consumption with daily consumption of more than 30 g for men and more than 20 g for women. Earlier studies had suggested that daily alcohol consumption of more than 60 g in men and more than 30 g in women per se can lead to fatty liver (25). Besides the difficulties in the correct assessment of alcohol intake recent studies show that low to moderate alcohol consumption (in contrast to a higher consumption) is more likely to be associated with a risk reduction for fatty liver (26-28). This evidence (from retrospective data) is still inconclusive because some recently published studies showed that modest alcohol consumption increases hepatic fat without increasing the risk of advanced fibrosis. Prospective data, actually suggest that NAFLD patients with regular alcohol intake, although within the safe thresholds, are at higher risk of liver disease progression, including HCC (29).

The authors of the German guideline recommend distinguishing NAFLD from ALD or mixed forms, with a daily alcohol limit of 10 g for women and 20 g for men (16). According to many experimental studies the combination of different unfavorable influences like alcohol, overweight, and free fatty acids, together leads to common damage on hepatocellular lipid accumulation, induces hepatic inflammation, fibrosis and carcinogenesis. Alcohol in rather low dosis already provokes synergistic pathological effects in combination with obesity (30).

There is significant overlap in the pathways by which NAFLD and ALD cause disease. Some data imply an association between drinking of fewer alcohol and reduced NASH and fibrosis. These studies, however, suffer from significant limitations, which are discussed in detail elsewhere (31). The authors think that there is insufficient evidence for or against moderate alcohol consumption in NAFLD. It is, however, important that NASH as well as advanced fibrosis, delineate hepatic damage including an increase in the risk of HCC, which argues against a moderate alcohol consumption (31). Taken together, counseling NAFLD patients for alcohol abstinence should be maintained.

Combinations of adverse conditions in lifestyle increased the risk for fatty liver, as identified by recently developed predictor algorithm FLI (that is fatty liver index; an algorithm based on waist circumference, body mass index (BMI), triglyceride, and gamma-glutamyl-transferase (GGT). The most striking individual impacts on the likelihood for FLI above 60% were observed for physical inactivity (P<0.0005 for both genders) and alcohol consumption (P<0.0005 for men) (9,19).

Taken together NAFLD is per definition the occurrence of hepatic steatosis (fat in the liver) proven by imaging or liver histology. Secondary causes of hepatic fat accumulation (e.g., increased alcohol consumption, certain medications, and other medical conditions) have to be excluded. Risk factors for fatty liver (NAFL, hepatic steatosis) are summarized in Table 1.

Who should be screened for the presence of NAFLD and how should the screening be carried out?

A systematic review showed incidence rates of 31 and 86 per 1,000 person-years based on two Japanese papers, but at the same time a significantly lower incidence rate of 29 cases per 100,000 person-years was reported in an English paper (14). Due to the heterogeneous results, further studies are required in order to make more
precise statements about the incidence of NAFLD. Thus, screening programs should be site specific and incorporate the genetic background. Hispanics and Asians are usually less obese but at higher risk than African Americans, who exhibit a lower degree of NAFLD and a lower NAFLD prevalence.

Prevalent NAFLD cases will increase to about 21%, from 83.1 million [2015] to 100.9 million [2030], while prevalent NASH cases will rise to 63%, from 16.52 million to 27.00 million cases (33). These estimates have shown 28,200 liver-related deaths among the NASH population (7.6% of total deaths) in 2015, increasing to 78,300 or 10.9% of total deaths in 2030 (33). These are already threatening data especially. Screening programs should be adapted to such high number of entities published recently. Incidence and prevalence are increasing especially in risk populations. Estimated data however have to be proven.

A current study indicates a prevalence of fatty liver of 42.2% (34). Markus et al. (3) calculated a (similar) prevalence of 39.7% for the population between 45 and 81 years in Pomerania, northeast of Germany. In the age group between 20 and 79 years, the prevalence of fatty liver was already 29.9%. Overall, the analyzes showed that participants with fatty liver tend to be male, inactive, and smokers.

Among the NAFLD entities, not NAFL but NASH proceeds to cirrhosis and HCC. The epidemic of the overweight and its impact on the increasing prevalence of NAFLD, suggests NASH to be the most common cause of advanced liver disease in coming decades (14).

Musso and coworkers depicted that NAFLD justifies a screening for cardio-metabolic risk and for progressive liver disease. The combination of three noninvasive tests (NAFLD fibrosis score, Cytokeratin 18, elastography) with liver biopsy may optimally individuate patients with NASH, with or without advanced fibrosis (35).

Already in 2007 and 2010, studies by Ruiz and Alisi showed that obese patients with NAFLD have significantly higher endotoxin and LPS levels as well as significantly higher TNF-expression levels in the liver (36,37). However, an increased endotoxin content was not found in a pediatric NASH cohort (38).

Depending on the patients’ predisposition, lipid accumulation could promote lipotoxicity and mitochondrial dysfunction, thus triggering hepatocyte inflammation, death, and fibrosis. In this context free fatty acids and free cholesterol have been ascertained to be toxic. To make the picture more complex, the pathogenesis of NAFLD encompasses pathological connections between the liver and other organs, including the adipose tissue and the gut (39). Nonetheless, presently there is no accurate noninvasive method to detect NASH.

| Main risk factors                      | Drugs                                      | Metabolic diseases (resulting in a NASH-like histology) |
|----------------------------------------|--------------------------------------------|---------------------------------------------------------|
| Insulin resistance                     | Tamoxifen, corticosteroids amiodarone,      | Hypobetalipoproteinemia                                  |
| Obesity                                | methotrexate, estrogens, valproic acid,     | Abetalipoproteinemia                                     |
| Sedentary lifestyle                    | antiretroviral medications                 | Wilson’s disease                                         |
| Type 2 diabetes mellitus               | Carbohydrate excess (e.g., diet and total | Lipodystrophies                                          |
| Hypertension                           | parenteral nutrition)                      | Andersen disease                                         |
| Dyslipidemia                           | Rapid weight loss                          | Weber-Christian syndrome                                 |
|                                        | Altered small bowel anatomy                | Infections                                              |
|                                        | Obesity surgery (e.g., jejunoileal bypass)  | Chronic hepatitis C virus                                |
|                                        | Pancreateoduodenal resection               | Human immunodeficiency virus and acquired immune        |
|                                        | Short gut                                  | deficiency syndrome                                      |
|                                        | Metabolic diseases (resulting in a NASH-like histology) | Emerging associations                               |
|                                        |                                            | Polycystic ovarian syndrome                             |
|                                        |                                            | Hypothyroidism                                          |
|                                        |                                            | Obstructive sleep apnea                                 |
|                                        |                                            | Hypopituitarism                                         |
|                                        |                                            | Hypogonadism                                            |
|                                        |                                            | NAFLD, nonalcoholic fatty liver disease.                |

Table 1 Risk factors for NAFLD [according to (16,21,32)]
Laboratory screening by means of the determination of transaminases is not sensibly possible, since patients with NAFLD predominantly have normal transaminases (40). The prevalence of ultrasound-detected steatosis in the general population is on average around 30% high (14), but the sensitivity of ultrasound for the detection of steatosis is relatively low at around 60% for a screening procedure (41). In addition, sonography is expensive and requires more resources and expertise.

The prevalence of NAFLD (defined as liver fat concentration ≥6%) was low in type 1 diabetes (8.8%) but high in type 2 diabetes (T2D). In a recently published single center study with T2D patients, NAFLD was associated with several markers of insulin resistance (42). Lai et al. screened for NAFLD and advanced fibrosis among patients suffering from type 2 diabetes. The majority of these 560 patients had non-alcoholic steatohepatitis (about 83%) and ≥F1 fibrosis (87%) while advanced fibrosis was seen in 37% (43). Thus, the prevalence of NAFLD and advanced fibrosis identified by transient elastography is rather high among T2DM patients.

There are some arguments in favor of considering screening in high-risk groups such as diabetics and obese people, in whom NAFLD has a prevalence of up to 70% (44). Missing data with regard to diagnostic criteria, prediction of the course and progress as well as established therapy indications and procedures do not currently allow a recommendation for a screening.

Screening of overweight and obese children in primary care for NAFLD with referral to paediatric gastroenterology has the potential to identify clinically relevant liver pathology (45). However, many children with suspected NAFLD were shown to have liver disease other than NAFLD. Elevation of transaminases was combined with worsening of the disease. Levels of ALT, however, could not discriminate between stages of NAFLD on the individual patient level. Importantly, the screening and referral process followed by liver biopsy was able to identify many obese children with advanced fibrosis that would have otherwise remained undiagnosed. Amongst children with NAFLD, approximately half had steatohepatitis (45).

A liver biopsy is the method of choice to determine the stage of fibrosis. However, due to the risk of complications and the high costs, this screening method is not suitable for use within the general population. Hagström et al. have now examined the performance of 5 non-invasive scoring systems for the prediction of severe liver diseases. The authors collected data from the Swedish Apolipoprotein Mortality Risk cohort on persons 35 to 79 years old from 1985 through 1996. APRI (n=127,302), BARD (n=75,303), FIB-4 (n=126,941), Forns (n=122,419), and the nonalcoholic fatty liver disease (NAFLD) fibrosis score (NFS, n=13,160) were applied. The predictive values of the scoring systems applied decreased with time and were more meaningful in men. All scoring systems were more accurate in attendees with risk factors for NAFLD at baseline, with AUROCs reaching 0.83. The predictive ability was modest. Finally, there is a need for better scoring systems to evaluate the risk of fibrosis in the total population (46).

Lean NAFLD is drawing considerable attention. Leung et al. prospectively followed up 72 lean (BMI <25 kg/m²) and 235 overweight (BMI ≥25 kg/m²) patients with biopsy-proven NAFLD. Lean patients had a lower grade of steatosis and lower stage of fibrosis than overweight patients. In addition, the event-free survival was better in lean patients (47,48).

Taken together screening for NAFLD in adults is currently not recommended for the general population (16). For risk groups, examinations for the existence of a NAFLD can be carried out. Patients with NAFLD who either have clinical symptoms or signs of liver disease or abnormal liver-associated laboratory values should be evaluated for the presence of NASH.

NAFLD patients who have neither clinical symptoms nor signs of liver disease or abnormal liver-associated laboratory values should be evaluated for the presence of cardio-metabolic diseases, alcohol consumption and drug side effects.

Not to determine NAFL but to determine a severe form of NAFLD (NASH, NASH fibrosis, NASH cirrhosis) is necessary to define the associated prognosis. Presence and stage of fibrosis, which is the most important determinant for the severity and progression of NAFLD should be confirmed.

What is an ideal and economical diagnostic tool?

Still, there is a generally accepted need for non-invasive diagnostic tools and biomarkers for NASH and fibrosis that can quickly and easily diagnose the severity of NAFLD, monitor liver changes, and identify high risk patients (21).

Modern biomarkers include prediction models (e.g., NAFLD fibrosis score, FIB-4 index and BARD score) or detection of inflammation (e.g., circulating keratin 18 fragments), or fibrosis (e.g., FibroTest®, ELF™ or
Pro-C3 tests). For clinical use, biomarkers may discriminate between patients with NASH or advanced fibrosis, predict dynamic changes in NASH/fibrosis over time, and provide long-term prognostic information (12).

NAFLD fibrosis score (NFS) as well as FIB-4 index seem to be both, cost-effective and very sensitive tools to identify patients with advanced fibrosis. Fibrosis scores for NAFLD are summarized in Table 2.

Srivastava and colleagues investigated a diagnostic algorithm for NAFLD patients, in order to detect severe fibrosis and cirrhosis. By conducting a prospective cohort study, they analysed whether the sequential use of FIB-4 and ELF in over 3,000 patients reduced avoidable referrals from primary care to secondary care by 81% (50).

By combining clinical risk factors (dangerous alcohol consumption, elevated transaminases, risk factors of the metabolic syndrome), serum markers, and transient elastography primary care programs were able to identify more patients with advanced liver disease in alcoholic as well as in non-alcoholic liver diseases (51).

Methods have been developed, that detect liver fat (NAFL) using magnetic resonance imaging-derived proton density fat fraction (MRI-PDFF) on routine MRI devices and do not require any additional spectroscopy coils (52). MRI-PDFF is based on the indirect estimate of accompanying changes in fatty liver (T1 deviation, T2 deviation and T2 drop, spectral complexity of fat, deviation when integrating separately acquired water and fat images into a combined image to record the fat fraction), but not of significant influencing factors of the NAFLD, e.g., body mass index, inflammation or iron overload. The etiology of liver disease also has no influence on the detection of fat content. In recent years, MRI-PDFF has shown high diagnostic accuracy with the histological degree of steatosis. It is now recognized as a reference standard in diagnostic radiology (53). So far, results of noninvasive screening procedures are suboptimal. However, MRI-PDFF presents the most accurate modality for detecting hepatic fat or NAFL. On the other side MRE appears to be an accurate method for scaling liver disease. MRI-PDFF in combination with MRE might provide a precise method for stratifying NAFLD patients with regard to their overall risks. Still, total costs and long waiting times for these modern techniques display a major challenge to most physicians and hospitals.

Liver biopsy still remains the current gold standard for diagnosis, despite limitations regarding sampling variability, invasive nature, and high cost (16). Currently, liver biopsy is also considered the gold standard for acquisition of NASH and determination of the fibrosis stage. On the other hand, tissue biopsy includes a high standard deviation with regard to characteristic features of NASH. Kose et al. did not record severe adverse events (e.g., fever, abscessing, anaphylactic shock, organ damage, sepsis, deaths). Despite a certain invasiveness, they considered percutaneous liver biopsy as being safe and with low complications (54). Up to now liver biopsy is the only diagnostic that can clearly differentiate between NAFL, NAFL with low inflammation and NASH (steatosis with lobular and portal inflammation and hepatocellular ballooning), and defining the presence or absence of even low fibrosis (21). For clinical purposes, a simple but robust algorithm for categorizing liver lesions in NAFLD patients was introduced by Bedossa et al. (55). Nevertheless, severe complications and fulminant internal haemorrhage are discussed in patient consent forms for sonography guided percutaneous liver biopsy.

Ultrasound-based transient hepatic elastography has been the first true bedside technique to reproducible screen for liver fibrosis. In comparison to other techniques TE has an excellent interobserver variability, small sampling error, and good reproducibility. If liver fibrosis is suspected, TE should be performed directly after the abdominal ultrasound and routine blood tests. In cases of severe obesity (BMI >30) or ascites, the XL probe should be used (56).

There are no scientific data on risk adaptation or cost-effectiveness of NAFLD diagnostics. Step diagnostics takes parameters into account based on their availability, costs/effort and pretest probability of positive results. Due to the high prevalence of NAFLD, the implementation of a level 1 for the identification of high-risk patients (NASH, NASH fibrosis) who should be further diagnosed, appears to be of particular importance. Studies indicate that this can be improved further despite medical knowledge of the relevance of NAFLD.

In addition, cytokeratin 18 fragments in the blood, also called M30, represent novel biomarkers for the presence of inflammation and apoptosis in NASH. So far circulating cytokeratin 18 fragments are not recommended by guidelines. Noninvasive tests are needed. A better stratification of the patients could for example be achieved by a combination of different markers, e.g., NFS with M30 or NFS with TE, or all three markers together to ensure an accurate identification of high-risk patients with advanced inflammation and fibrosis (57).

Taken together NAFL in most cases is diagnosed initially by ultrasound, quantified by CAP in daily routine and by
Table 2 Fibrosis scores appropriate for NAFLD [according to (16,19,21,32,49) and literature indicated in the right column]

| Score                        | Components                                                                                           | Risk of fibrosis                                      | Literature                                                                                           |
|------------------------------|-------------------------------------------------------------------------------------------------------|-------------------------------------------------------|------------------------------------------------------------------------------------------------------|
| Serum tests                  | Serum glucose, platelet count, albumin, AST/ALT ratio; age, BMI, and diabetes status                   | >0.676 (fibrosis), <-1.455 (no fibrosis)               | Angulo P, Hui JM, Marchesini G, et al. The NAFLD fibrosis score: a noninvasve system that identifies liver fibrosis in patients with NAFLD. Hepatology 2007;45:846-54 |
| FIB-4                        | Age, AST, platelet count, ALT                                                                       | <1.45 (no fibrosis NPV 90%), >3.25 fibrosis (PPV 65%) | Sterling RK, Lissen E, Clumeck N, et. al. Development of a simple noninvasive index to predict significant fibrosis patients with HIV/HCV co-infection. Hepatology 2006;43:1317-25 |
| BARD                         | BMI, diabetes, AST/ALT ratio                                                                        | 0–1: low risk, 2–4: high risk, NPV 97%               | Harrison SA, Oliver D, Arnold HL, et al. Development and validation of a simple NAFLD clinical scoring system for identifying patients without advanced disease. Gut 2008;57:1441-7 |
| Fibro-Test© commercially available | α2-macroglobulin, haptoglobin, apolipoprotein A1, bilirubin, gamma glutamyl transpeptidase (GGT), age, gender | 0–1.0, 0 no fibrosis, 1.0 F4 fibrosis                | Ratziu V, Massard J, Charlotte F, et al. Diagnostic value of biochemical markers (FibroTest-FibroSURE) for the prediction of liver fibrosis in patients with non-alcoholic fatty liver disease. BMC Gastroenterology 2006;6:8. https://www.biopredictive.com/ |
| ELF® enhanced liver fibrosis test commercially available | Hyaluronic acid (HA), procollagen III amino-terminal peptide (PllNP), and tissue inhibitor of matrix metalloproteinase 1 (TIMP-1) | >10.51 advanced fibrosis                             | https://www.nice.org.uk/guidance/ng49/chapter/recommendations |
| Pro-C3® test commercially available | NASH biomarker synthesis of type III collagen, propeptide                                         |                                                       | Gloomba R, Sanyal AJ. The global NAFLD epidemic. Nat Rev Gastroenterol Hepatol 2013;10:686-90. www.bms.com/researchers-and-partners/areas-of-focus. html |
| CK18 M30 (M30-Apopto-sense®) commercially available | Caspase-cleaved CK18 fragments                                                                  | <150 U/L healthy, >200 U/L elevated                  | Fitzpatrick E, Mitry RR, Quaglia A, et al. Serum levels of CK18 M30 and leptin are useful predictors of steatohepatitis and fibrosis in paediatric NAFLD. J Pediatr Gastroenterol Nutr 2010;51:500-6 |
| leucocyte telomere length (LTL) | Fibrosis in T2DM                                                                                     |                                                       | Kim D, Li AA, Ahmed A. Leucocyte telomere shortening is associated with nonalcoholic fatty liver disease-related advanced fibrosis. Liver Int 2018;38:1839-48 |
| Imaging methods              | Liver stiffness measurements                                                                         | <7 kPa normal, <12 kPa severe fibrosis                | Sandrin L, Fourquet B, Hasquenoph JM, et al. Transient elastography: a new noninvasive method for assessment of hepatic fibrosis. Ultrasound Med Biol 2003;29:1705-13 |
| Dynamic liver function test  | Reflects cytochrome P450 1A2 activity                                                                | LiMax values >315 µg/kg/h are considered normal       | Blüthner E, Pape UF, Stockmann M, et al. Assessing Non-Invasive Liver Function in Patients with Intestinal Failure Receiving Total Parenteral Nutrition-Results from the Prospective PN Liver Trial. Nutrients 2020;26:12 |
MR in studies. Fibrosis is diagnosed by liver elastography. NASH could be determined by biopsy or probably by some serum tests. A recently published noninvasive lipidomic serum test assessed by two panels of triglycerides distinguished between NAFLD and NAFL and between NASH and NAFL with high accuracy (58). Thus, metabolomics provides a new technology for noninvasive biomarkers to improve NAFLD diagnosis.

Diagnostic algorithms for differentiating NAFLD by non-invasive tools, including transient hepatic elastography, are presented by Roeb et al. in (16,59).

How should we treat the patients, who have to be treated at all?

NAFL displays hepatic steatosis without evidence of inflammation and increased risk of developing liver-related complications. Patients with NASH, however, should be treated to prevent NASH progression to advanced fibrosis, to prevent cirrhosis and to prevent the development of its hepatic complications.

The therapeutic goal or rationale of NAFLD is to reduce insulin resistance with reduced cardiovascular end organ damage, reduced liver failure (including HCC), and the extension of lifetime. The change in lifestyle in terms of an increased physical activity as well as a moderate weight loss is able to delay or even prevent insulin resistance. Regular physical activity reduces the risk of NAFLD. Increased calorie intake, however, increases the corresponding risk (16).

Hepatic fibrosis displays the main characteristic predicting liver related and all cause mortality in NAFLD. Cenicriviroc, a promising new antifibrotic agent, blocks the CCR5 and CCR2 receptor. The efficacy and safety of cenicriviroc in patients with NASH and liver fibrosis were assessed in the CENTAUR study, a phase 2b, double-blind, randomized, placebo-controlled, multinational study with 289 NASH subjects (60). In the Cenicriviroc group, the fibrosis score remained improved by at least one level, with patients with advanced fibrosis benefiting the most (60). A phase 3 trial (NCT03028740) will have to clarify whether the results are stable.

Friedman et al. summarizes the therapeutic targets: metabolic pathways, cell stress and apoptosis, immune targets, fibrosis, and the intestinal microbiome and brown adipose tissue. Thus, sites of drug action—currently in clinical trials—are based on their primary locus of activity. Targets include fatty acids, insulin resistance, immune signals, inflammatory cells caspases, oxidant stress, extracellular matrix, hepatic stellate cells, other key features of fibrogenesis as well as glucose and lipid homeostasis.

There are currently no drugs approved for the NAFLD indication. There are promising phase 3 studies on the FXR agonist obeticholic acid and elafibranor, a dual PPARα/δ agonist. Metformin, thiazolidines and glucagon-like peptide-1 receptor agonists (GLP-1 RA) represent interesting candidates in order to treat type 2 diabetes with NAFLD (61). The risk of developing type 2 diabetes decreases with the improvement of fatty liver and certain therapies (e.g., pioglitazone) are effective both in the treatment of hyperglycemia and in NAFLD (21,24,62). Numerous substances have been and are being analyzed for their effectiveness in NASH in extensive studies. Without claiming to be complete Roeb & Geier provide an up-to-date overview about the different therapeutic approaches at NAFLD and NASH (63). Furthermore, the combination of compounds that addressed various targets of the metabolic pathways involved in NASH progression, e.g., GLP-RA/glucagon receptor agonist and GLP-RA/gastrointestinal peptide agonist, are promising future options (63,64). SGLT2 inhibitors as well as GLP-1 RAs might have a positive effect on the treatment of NAFLD in type 2 diabetics. A systematic review was published recently (65).

NASH is a rapidly growing etiology of end-stage liver disease in the US and elsewhere with significantly higher post-transplant survival compared to HCV (66). Currently NASH is the second leading cause for liver transplantation overall, even the leading cause in in females. Given the rate of increase, NASH will likely rise to become the leading indication for liver transplantation in males as well (66,67).

Therapy of cardiovascular risks and associated metabolic diseases has to be considered because of mortality data. For each therapeutic strategy monitoring as well as an individualized follow-up of the treatment strategies is mandatory. Up to now NAFLD and NASH are considered as one disease. Since we observe different outcomes the management of non-cirrhotic NAFLD cases should be differentiated from cirrhosis in therapy strategies.

Finally, up to now, treatment of this disease is limited to lifestyle modifications.

What is the minimum therapy? What are essential conservative therapeutic measures?

The following recommendations are mentioned in nearly all guidelines (9,16,19,68-70) published so far:
In overweight or obese patients with NAFLD, regardless of the composition of their macronutrients, a weight-reducing diet change should initially be used with the aim of sustainable weight loss of 0.5–1.0 kg per week.

NAFLD patients should be recommended to exercise regularly in the form of endurance and/or strength training.

Patients with compensated NASH cirrhosis should be motivated to undertake a sports program under supervision.

In decompensated NASH cirrhosis patients, muscle therapy should be used to counteract muscle loss.

Patients with NAFLD should be informed about the risks of smoking.

Patients with NAFLD should refrain from drinking alcohol.

Patients with NASH cirrhosis should strictly avoid alcohol.

Drinking of coffee should be recommended due to beneficial effects on the liver and the cardiovascular system.

An indicated statin therapy can be maintained until liver values increase up to 3 times the upper normal value.

Metabolically neutral antihypertensives should be given preference in the case of hypertension requiring treatment—primarily inhibitors of the renin-angiotensin-aldosterone system.

With decompensated liver cirrhosis, vasodilating drugs should not be used.

In the case of manifest type 2 diabetes up to the Child A stage, metformin can be used as an oral antidiabetic drug of choice, even in the case of elevated transaminases.

Vaccination according to respective national vaccination recommendations (e.g., STIKO-, ECDC-, and CDC-guidelines) should be given to patients with chronic liver diseases, especially against hepatitis A, hepatitis B, and influenza (71-73).

Regular monitoring with the aim of early detection of HCC should be carried out according to the recommendations in the Diagnostics section.

**Which pharmacological therapies are permitted?**

Since simple NAFLD, unlike NASH, is not associated with over-mortality or liver-associated morbidity, from a hepatological point of view, even in the absence of large randomized studies, drug therapy is only an option for patients with secured NASH (21).

An overview of NAFLD agents currently under development and their pharmacologic targets is provided by Sumida et al. (61). There are 5 candidates who have been successful in studies, but which will be launched in 2021 at the earliest (obeticholic acid, elafibranor, selonsertib, cenicriviroc, and resmetirom) (74). The results of the FLINT study were recently published, in which the administration of obeticholic acid, a farnesoid X receptor agonist, was associated with a significant improvement in the histological characteristics of NASH including fibrosis compared to placebo (75). A total of 25 mg obeticholic acid significantly improved fibrosis and key components of NASH disease activity among patients with NASH (76). The authors demonstrated histological improvements. The most common adverse event was pruritus (51%) in the obeticholic acid 25 mg group. Pruritus incidence however was generally mild to moderate in severity (76).

Cenicriviroc is an inhibitor of CCR2 and CCR5 receptor and initially an experimental drug candidate for the treatment of HIV infection. According to Friedman et al. given the urgent need to develop antifibrotic therapies in NASH, these findings warrant phase 3 evaluation (77). After 1 year of Cenicriviroc treatment, twice as many subjects achieved improvement in fibrosis and no worsening of steatohepatitis compared with placebo (77).

Levels of hepatic aminotransferases could be lowered by administration of vitamin E and pioglitazone. In addition, both substances improved hepatic steatosis and inflammation. Improvement of fibrosis scores, however, could not be achieved. Thus, the authors concluded that pioglitazone was no better than placebo with regard to the primary outcome (78).

**Surgical and endoscopic therapeutic procedures**

Adipo-surgery should be offered to patients with a BMI ≥35 kg/m² who are suffering from one or more obesity-associated morbidities, e.g., metabolic syndrome with type 2 diabetes, disorders of fat metabolism, coronary heart failure, high blood pressure, NAFLD or NASH, kidney damage, sleep apnea, obesity-hypoventilation syndrome, gastro-esophageal reflux disease, pseudotumor cerebri, bronchial asthma, chronic varicose veins, urinary incontinence, immobilizing joint disease, fertility restrictions or polycystic
ovarian syndrome, if the conservative therapy is exhaustive. Patients with a BMI ≥40 kg/m² and coexistent type 2 diabetes not only benefit from better glycemic control or reduced antidiabetic medication during a metabolic intervention, but also from sustainable weight reduction. This is superior to conservative therapeutic approaches. Evidence of exhausted conservative therapy in the sense of obesity surgery is not required for the indication of the operation in these patients (9,79,80).

The gastric band, tubular stomach formation, proximal Roux-en-Y gastric bypass and omega-loop gastric bypass may be considered as an obesity surgical or metabolic intervention in patients aged 65 or older. Most of the evidence is available for proximal Roux-en-Y gastric bypass and for gastric tube formation (80).

The gastric balloon can be offered to patients with obesity after failure of conservative therapy and if surgery is refused or if there are contraindications to surgery as part of an appropriate accompanying program. The implantation of the Endobarrier™ can currently only be recommended as part of studies.

There is growing evidence on mutual relationships between NAFLD and type 2 diabetes. For this reason, guidelines including bariatric surgery as well as endoscopic procedures on patients with grade II obesity (BMI >35) or more are currently being created by scientific organizations such as the National Institute for Health and Care Excellence (NICE) in the UK, EASL, EASD and EASO.

Which monitoring and long-term management is optimal?

For the prevention of NAFLD, especially in pediatric particularities, positive recommendation for lifestyle changes with weight loss and increased physical activity as well as the refractive from fructose-containing sweetened foods and drinks, especially in the form of snacks, can be given. Intensity of long term monitoring depends on grade of fibrosis and establishment of cirrhosis. For capacity reasons HCC screening every 6 months is reserved for patients with NASH fibrosis and NASH cirrhosis only. Here, reference is made to the corresponding HCC guidelines from AASLD (69) and EASL (70).

Conclusions & remaining open questions

Meanwhile, NAFLD has become the most common chronic liver disease in industrialized countries. The prevention of overweight and lack of exercise targets the most important risk factors. Physical activity has been shown to improve fatty liver even without reducing body weight. So far, no specific drug therapy has been approved for this illness, neither for NASH nor for NAFLD in general. Despite diverse publications the following questions have not yet been answered. An intensive discussion is required for the time being:

- What is the optimal definition of NAFLD and NASH?
- How can NASH be diagnosed in primary care?
- Who should be followed up and when?
- What is the minimal therapy?
- What are essential conservative therapeutic measures?
- Which monitoring and long-term management is optimal?
- What is the optimal frequency of follow-up examinations?

Acknowledgments

Funding: This work was funded by Deutsche Forschungsgemeinschaft Ro 957/8-1, Ro957/11-1, von-Behring-Röntgen Foundation #66-0008, and UKGM #10/2013.

Footnote

Provenance and Peer Review: This article was commissioned by the Guest Editors (Ralf Weiskirchen and Wolfgang Stremmel) for the series “Unresolved Basic Issues in Hepatology” published in Annals of Translational Medicine. The article was sent for external peer review organized by the Guest Editors and the editorial office.

Peer Review File: Available at http://dx.doi.org/10.21037/atm-20-3760

Conflicts of Interest: The author has completed the ICMJE uniform disclosure form (available at http://dx.doi.org/10.21037/atm-20-3760). The series “Unresolved Basic Issues in Hepatology” was commissioned by the editorial office without any funding or sponsorship. The author has no other conflicts of interest to declare.

Ethical Statement: The author is accountable for all aspects of the work in ensuring that questions related to the accuracy or integrity of any part of the work are
appropriately investigated and resolved.

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Cite this article as: Roeb E. Non-alcoholic fatty liver diseases: current challenges and future directions. Ann Transl Med 2021;9(8):726. doi: 10.21037/atm-20-3760