1 Current therapies and new developments in NASH

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ABSTRACT
Non-alcoholic steatohepatitis is becoming the most important aetiology for advanced liver disease. There has been important progress in the field in recent years and the complexity of the pathophysiology of NASH is better understood. Multiple non-invasive circulating and imaging biomarkers have been tested. The importance of lifestyle has been recognised and several drugs are being tested in clinical trials. This review addresses the challenges that healthcare professionals face in the management of NASH patients.

INTRODUCTION
Non-alcoholic steatohepatitis (NASH) is a fast-growing and highly prevalent threat to health, set to become a major cause of liver cancer and transplant. NASH, an advanced form of non-alcoholic fatty liver disease (NAFLD), is strongly linked to obesity and metabolic disorders.

There are no licensed therapies for NASH, despite its prevalence and clinical significance. While a number of drugs are in development, difficulties in staging and identification of accurate biomarkers have hampered the approval of effective therapeutics. The result is a reliance on lifestyle measures, primarily diet.

We brought together nine experts to discuss hot topics and current developments in NASH. Here, we provide an overview of the disease and consider how new technologies, from AI to novel biomarkers to gut microbiome signatures, could drive the introduction of new therapies.

We define the term NAFLD as representing the whole spectrum of fatty liver disease, including NAFLA/steatoisostasis and NASH/steatohepatitis.

Epidemiology of NAFLD/NASH

Key points
► NAFLD is highly prevalent and increasing, with strong links to obesity and diabetes.
► The impact of NAFLD is increasingly seen in severe liver disease and mortality.

NAFLD affects at least a quarter of the global population. This figure may be an underestimate, given that obesity rates globally have been rising since the 1970s.

NAFLD is likely to have exceeded 30% prevalence in most middle-income and high-income countries. The most recent data are from Asia and suggest that NAFLD affects 30% of the Asian population, topping 40% in some territories.

A case for screening?
NAFLD is closely linked with type 2 diabetes and obesity. In one study from China which used transient elastography to screen more than 1900 people with type 2 diabetes, 73% of people had NAFLD, rising to 95% of those who had a body mass index (BMI) of 30 or over. Some 35% of those with both risk factors had high liver stiffness suggestive of advanced fibrosis. This suggests a case for screening people with diabetes or obesity for NAFLD, as recommended by current Asian Pacific guidelines.

NAFLD is not, however, confined to the obese population. In the USA, 43% of people with NAFLD are not obese, rising to 71% in Sweden.

Figures on incidence of NAFLD are not easy to come by, but a study from Hong Kong using serial proton-MR spectroscopy to screen community participants with no liver disease at baseline found 13.8% developed fatty liver over 4 years, giving an incidence of 3.7 per 100 person-years. In contrast, a study using serial transient elastography examinations showed that 52% of patients with type 2 diabetes developed incident fatty liver in 3 years.

Therefore, current European and Asian Pacific guidelines support screening people with diabetes or obesity for NAFLD. That said, the optimal screening strategy remains unclear and will depend on the local clinical practice, referral pathway and availability of different tests. Recently, a consensus paper commissioned by the American Gastroenterological Association recommends the use of the Fibrosis-4 index (FIB-4) followed by specific fibrosis tests to screen for advanced liver disease in patients at risk of NAFLD.

Clinical impact
The impact of NAFLD should not be underestimated. It is the fastest-growing indication for liver transplantation in the USA, accounting for more than 25% of transplants, compared with around 3% in 2002. It is also the fastest-growing cause of liver cancer among candidates for liver transplant. Incidence of decompensated liver cirrhosis and liver-related deaths are projected to rise in coming decades. While the effect of NAFLD on mortality has been masked by cardiovascular disease, we can expect to see more people with severe disease dying of liver-related causes in future. That being said, cardiovascular disease and extrahepatic malignancies will remain the leading causes of death in the vast majority of patients with NAFLD in the foreseeable future. Clinicians taking care of patients with NAFLD should be aware of the association and provide proper care of the cardiometabolic...
conditions and screening for cancers according to current guidelines.

Across the world—but especially in South America, the Middle East and East Asia—childhood obesity has risen fast since 2000.16 Unless the trend can be reversed, these children are growing up with risk factors for serious liver disease as adults.

One hot question is whether the terminology of NAFLD should be changed, given that many people have liver damage both from fatty liver and alcohol. A new name of MAFLD—metabolic-associated liver disease—has been proposed, to include people with fatty liver and metabolic risk factors.17 The proposal is under discussion by professional bodies worldwide.

While it might include more people who have coexisting alcohol-related damage, it would exclude those with fatty liver who have not yet developed metabolic risk factors.18 As such, it is likely to affect understanding of the epidemiology of fatty liver disease.

Pathophysiology of NAFLD

Key points

► NAFLD is a complex disease driven by lipotoxicity, insulin resistance and activation of inflammatory and immune pathways, closely linked to metabolic disorders.

► There is increasing evidence of a link between the gut microbiome and development of both insulin resistance and NASH.

NAFLD is a complex disease which involves many of the systems of the body.

The development of NAFLD comes about through a process of multiple parallel ‘hits’ which put stress on the liver.19 External factors which raise the risk include unhealthy diet such as fructose overconsumption20 and lack of exercise, while genetic factors also affect people’s chances of developing the condition.21

Lipotoxicity and inflammatory pathways

Lipogenesis is a key factor in the early stage of development of NAFLD, driven mainly by increased delivery to the liver of free fatty acids (FFAs) from diet and from the adipose tissue and via increased de novo lipogenesis in the liver.22

An overload of FFAs and production of intermediaries such as ceramides and phospholipids in liver cells may result in apoptosis and liver damage. FFAs seem to be among the most important lipids involved in insulin signalling.23 Consequently, lipotoxicity might link metabolism with the development of inflammation and fibrosis in the liver. Unresolved inflammation frequently results in fibrosis and indeed fibrosis stage is by far the most relevant predictor of overall and disease-specific mortality.24 25

Inflammation is an important aspect of metabolic disorders. All adipose tissue contributes substantially to systemic and liver inflammation observed in NAFLD. Successful weight loss after bariatric surgery results in a massive reduction of IL-6 and TNF, especially in the subcutaneous adipose tissue.26 In addition, we can see from animal studies that lack of interleukin-1 successfully inhibits the transformation of steatosis to steatohepatitis and liver fibrosis.27

Many diverse inflammatory hits are able to affect insulin signalling with receptor activator of nuclear factor and TNF and IL-6 just being some examples.28 Conversely, adiponectin has a protective effect in almost all metabolic pathways affecting organs.29

Many more pathways such as nuclear receptors, bile acids or fibroblast growth factors have evolved in the past decade as key pathophysiological players in NAFLD and related pathologies.30–32

In addition to inflammatory pathways, various genetic polymorphisms affect the development and progression of NAFLD.24 Almost all identified genetic polymorphisms in NAFLD affect lipid pathways, highlighting the crucial role of metabolic pathways in this disorder.

Gut microbiota and NAFLD

More recently, researchers have investigated the role of gut microbiota, trying to identify a specific microbial fingerprint in NAFLD. This research could pave the way for a non-invasive stool test to identify the progression from NAFLD to advanced fibrosis.33 Researchers have identified 8 species of gut microbes which are twice as abundant in advanced fibrosis, and 22 species which are twice as abundant in mild or moderate fibrosis. More recently, a universal gut microbiome signature in NAFLD cirrhosis has also been described.34 Besides the microbiome also the mycobiome might be involved in the pathogenesis especially in advanced NAFLD.35

Two types of bacteria, Prevotella copri and Bacteroides vulgatus, have been shown to drive the production of branched-chain amino acids (BCAA) in the gut. Increased levels of BCAA have been identified in people with insulin resistance, and animal experiments have shown that P. copri induces insulin resistance in mice.36 Whether similar pathways are relevant in human insulin resistance in NAFLD remains to be proven.

The pathophysiology of NASH and NAFLD explains its close correlation with type 2 diabetes and the metabolic syndrome. Presence of either type 2 diabetes or NAFLD at least doubles the risk of developing the other condition. In addition, both diabetes and NAFLD drive an increase in cardiovascular disease (up to double the risk), chronic kidney disease (up to 3.5 times the risk) and intrahepatic and extrahepatic malignancy.27 38

The histology of NASH

Key points

► Liver biopsy remains the reference standard for diagnosis of NASH (steatosis vs steatohepatitis).

► Diagnosis and scoring are distinct and separate steps in the evaluation of NAFLD.

NAFLD encompasses a spectrum of liver lesions, from steatosis to cirrhosis, via NASH. The presence of 5% fat cells in the liver is a prerequisite for diagnosis, based on microscopic examination of liver tissue. Current European guidelines39 state that distinguishing NASH from simple steatosis requires biopsy (see figure 1), and the diagnosis cannot be made by clinical, biochemical or imaging measures.

Liver biopsy can also be used to assess severity (grade) and stage (fibrosis) of disease. This enables enrolment in clinical trials, evaluation of treatment response and may point to comorbidity risk factors. Among the usual limits of liver biopsy, especially its invasiveness, sampling variability is a challenging issue as it represents a tiny fraction (1/50 000) of the whole liver. This strongly supports the use of criteria of biopsy adequacy including length and width of liver specimen. Nevertheless, given the zonation of the disease (starting in centrolobular areas), we may expect less impact of sampling compared with viral hepatitis, for instance.

Scoring systems to identify NASH

Pathologists use two main types of scoring system to assess liver biopsy for NASH:
Unresolved issues in staging and diagnosis

Staging of fibrosis remains an issue—and an important one. Development of fibrosis may be the most relevant histological endpoint to monitor during clinical trials, yet the standard staging system lacks granularity.

In addition, the diagnosis of NASH itself could be refined, with the inclusion of additional key features such as centrilobular fibrosis and portal inflammation.

Circulating biomarkers of NASH

**Key points**
- Non-invasive testing is useful to rule out people unlikely to have advanced liver disease, without need for biopsy.
- Direct fibrosis biomarkers can be used in combination with simple scores such as FIB-4, but there remains a need for more sensitive, specific and better validated biomarkers in NAFLD.

While histology remains the reference standard for diagnosis, prognosis and monitoring treatment response in NAFLD, there are issues around accuracy, reader variation and safety—not to mention the need for less invasive tests.

In primary care or settings where pretest probability of severe disease is low, a test with high negative predictive value to rule out severe disease may be preferable. In secondary and tertiary care, probability of severe disease rises and a test needs higher positive predictive value to select cases for enhanced therapy or surveillance.

There are two clinically relevant liver disease categories in which high quality biomarkers are particularly needed.

Identification of at-risk NASH

Identifying at-risk NASH—where steatohepatitis is active and there is moderate fibrosis (NAS≥4, Fibrosis stage ≥2), suggesting that they are likely to progress to cirrhosis—is particularly important for identifying candidates for treatment.

To date, few blood-based biomarkers have been developed specifically targeting this condition. NS-4 is a composite biomarker comprising microRNA 34A, alpha2 macroglobulin, HbA1c and YKL-40; representing information about insulin resistance, inflammation and fibrogenesis. Research published in 2020 showed it out-performed several other biomarkers to identify this cohort of patients, although these results need further validation.

A study from 2021 by the LITMUS consortium compared the performance of 10 widely available biomarkers in prescreening patients for inclusion in a clinical trial. The researchers compared how many biopsies would be needed and how many eligible patients would be identified per 100 people tested. Performance varied, making it a trade-off decision as to whether to conduct more biopsies, but accept a marginally higher screen failure rate, to identify more eligible patients.

Some have suggested the use of combined MR elastography (MRE) with FIB-4 biomarkers (MEFIB). This is discussed further below in the section ‘imaging biomarkers’.

Identification of advanced fibrosis

People with advanced fibrosis—F3–F4—are likely to see outcomes worsen significantly.

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Dufour J-F, et al. Gut 2022;71:2123–2134. doi:10.1136/gutjnl-2021-326874
Some biomarkers are indirect measures of processes leading to fibrosis, such as markers of inflammation, necrosis and cell death, while others are direct serum markers of collagen components or factors regulating fibrogenesis.50

Risk stratification guides are currently used to direct patients along appropriate pathways. The FIB-4 combines three indirect markers (ALT, AST, platelets) with age, while the NAFLD Fibrosis score combines BMI, albumin and impaired fasting glucose or type 2 diabetes (see table 1). Both systems have a high sensitivity cut-off at the lower end of probability51–55 and exhibit a high negative predictive value to rule out advanced disease.

The most efficient route is to use biomarkers in combination within a care pathway, moving through clinical assessment to non-invasive imaging, to minimise the number of patients that require biopsy (see figure 2).51–61

Direct biomarkers give more granular evidence of fibrosis development and so provide a tractable ‘second-line’ test, that might also aid monitoring of progression/response to treatment. Candidates include markers of extracellular matrix turnover, for example, the PRO-C3 Collagen Neoepitope, which has been incorporated into several risk scores4,42 and the ELF test (see table 1).

The ELF test (combining procollagen III N-terminal peptide, hyaluronic acid, tissue inhibitor of metalloproteinase 1) exhibits a high negative predictive value62 but more modest positive predictive value in most clinical settings. Its use in a care pathway following FIB-4 improved the proportion of patients with advanced fibrosis undergoing specialist referral from 5% to 30%, although it should be noted that 87% of cases with mild disease were eliminated by FIB-4 in that study (see figure 2).64

Comparative studies demonstrate that many fibrosis biomarkers offer only marginal improvement over the performance of FIB-4 in routine care.65

**New markers on the horizon**
Researchers continue to search for new biomarkers in the field. Current areas of interest include use of metabolomic profiling,66 which measures serum lipids and amino acids; transcriptomic profiling gene expression patterns in the liver to guide identification of circulating protein biomarkers67; and measurement of circulating proteins.

In addition, new technologies are allowing measurement of liver proteases, which do not get released into the circulation.68

### Imaging biomarkers in NASH

**Key points**
- Use of non-invasive imaging assessment is increasingly important to identify high risk NASH, to monitor patients for progression, and to identify response to treatment in clinical trials.
- Long-term studies are needed to assess the association between longitudinal change in biomarkers and long-term clinical outcomes.

### Identification of NAFLD

While liver biopsy is the gold standard for identifying NASH, non-invasive imaging is becoming more important for staging and quantifying disease.

Conventional ultrasound has low negative predictive value so is of limited use, especially for mild steatosis. CT scan lacks specificity and sensitivity, and exposes the patient to ionising radiation.

The controlled attenuation parameter provides a useful indication of NAFLD, but the de facto gold standard has become MRI-derived proton density fat fraction (MRI-PDFF), for its ability to quantify fat content. It is widely used for non-invasive screening for NAFLD in research studies.

Assessment of fibrosis relies on indirect biomarkers, primarily tissue stiffness.69 It is assessed by elastography, with the vibration-controlled transient elastography (FibroScan) imaging machine the most commonly used.70 Studies have shown that patients with higher liver stiffness also have higher liver-related mortality, suggesting that this is a prognostic biomarker.71 MRE has been shown to predict progression to advanced fibrosis (stage 3 or 4) with a higher degree of accuracy than FibroScan.72

Combination of various biomarkers in one imaging session could allow for assessment of degree of steatosis, inflammation and fibrosis, by using both MRE and PDFF.

| Table 1 Circulating fibrosis biomarkers |
|----------------------------------------|
| Test | Description | References |
| Direct Fibrosis Biomarker Panels |
| AST:ALT ratio | AST (IU/L)/ALT (IU/L) | 128 |
| AST to platelet ratio index | AST (IU/L)/platelet count (10^12/L) x 100 | 54 |
| BARD score | Weighted sum of BMI≥28 = 1 point, AST/ALT ratio≥0.8 = 2 points, T2DM=1 | 129 |
| FIB-4 | Age x AST (IU/L)/platelet count (10^12/L) x √ ALT (IU/L) | 130 131 |
| NAFLD fibrosis score | −1.675+0.037 x age (years)+0.094 x BMI (kg/m2)+1.13 x IFG or T2DM (yes=1, no=0)+0.09 x AST/ALT ratio − 0.013 x platelet (10^12/L) − 0.66 x albumin (g/dL) | 53 |
| ELF | ELF=−7.412 + (ln(HA)*0.681) + (ln(PHIIIP)*0.775) + (ln(TIMP1)*0.494) | 60 132 |
| Fibro test | Patented algorithm combining total bilirubin, GGT, α2-macroglobulin, apolipoprotein A1, and haptoglobin, corrected for age and gender. | 133 134 |
| FibroMeter NAFLD | Patented algorithm combining age, body weight, glucose, AST, ALT, ferritin and platelet count | 135 136 |
| ADAPT | ADAPT=exp(log10((age x PRO-C3)/sqrt(Platelets)))x T2DM | 137–139 |
| FIBC3 | FIBC3=−5.939 + (0.053*age) + (0.078*AST) + (0.097*platelets) + (0.071*PRO-C3) | 138 139 |
| ABC3D | Age ≥50 = 1 point, BMI ≥30 = 1 point, platelet Count <200 = 1 point, PRO-C3 ≥15.5 = 1 point, T2DM≥2 points | 138 139 |
| AST:ALT, aspartate transaminase:alanine transaminase; BMI, body mass index; ELF, enhanced liver fibrosis; FIB-4, Fibrosis-4 index; IFG, impaired fasting glucose; NAFLD, non-alcoholic fatty liver disease; T2DM, type 2 diabetes mellitus. | |
Recent advances in clinical practice

before biopsy. The main issue with FAST is low positive predictive value.

Combining MRE with FIB-4 biomarkers (MEFIB) may help. A FIB-4 score of 1.6 or above, combined with MRE of 3.3 kPa or above, yields a 97.1 positive predictive value that a patient has fibrosis stage 2 or above and is therefore a candidate for treatment. The investigators believe a PPV above 90 negates the need for a biopsy altogether in these populations. MEFIB has been shown to be better than FAST in detecting at-risk NASH patients, among a cohort with biopsy-proven NAFLD.

The use of either vibration-controlled transient elastography or MRE, with FIB-4 where needed, could separate out patients with NAFLD into low risk fibrotic NAFLD and high risk for progressive fibrosis.

What is a clinically significant treatment response?

Despite the clear need for treatment, there are no licensed treatments for NASH. However, many drugs are in development. A clinically significant biomarker for response to treatment is crucial to identify effective therapies.

An MRI-PDFF liver fat reduction of 30% from baseline has been shown to correlate with five times higher odds of NASH resolution, compared with those not achieving 30% MRI-PFF reduction.

Researchers have not yet been able to identify a single clinically significant change in MRE that would indicate treatment response, but there are hints—for example, a 15% relative increase from baseline is associated with fibrosis progression.

So far there is some evidence of histological response for four imaging biomarkers: MRI-PDFF, Multiscan cT1, FAST and 15%–20% reduction in MRE, although this needs further confirmation.

Multidisciplinary lifestyle intervention in NASH

Key points

- The best evidence we have for lifestyle intervention in NASH is for weight reduction and the Mediterranean dietary pattern.
- Improved dietary composition by itself, weight loss of 5%–10% of the initial body weight and moderate exercise can make a difference for NASH patients in terms of improvement of all histological features.
- The lack of pharmacological interventions for NASH make lifestyle measures even more important (see figure 3). The challenge is adoption of these measures in the long term.
- The best evidence of benefit comes from the Mediterranean diet—a diet characterised by plentiful intake of olive oil, vegetables, fruits and nuts, legumes, whole grains, fish and seafood, and a low intake of red meat and especially processed meat, along with reduced carbohydrates intake (40% of the calories vs 50%–60% in a typical low-fat diet), especially sugars.

A recent 18-month trial with 294 participants suggests the Mediterranean diet can be enhanced by additional green plants rich in polyphenols. The green-Mediterranean diet reduced hepatic fat by 39% compared with 20% for the Mediterranean diet, despite similar weight loss, and both diets did better than controls provided only with healthy dietary guidelines.

The importance of increased phenolic acid intake (from fruits and vegetables, nuts, green tea and coffee) is independently

Figure 2 Proposed primary care pathway versus standard care for patients with NAFLD, adapted from Srivastava et al. EFL, enhanced liver fibrosis; FIB-4, fibrosis-4 index; NAFLD, non-alcoholic fatty liver disease.
Recent advances in clinical practice

Recent advances in clinical practice associated with lower prevalence of insulin resistance and NAFLD and fibrosis measured by fibrosis marker (FibroTest).81

Carbs or fats?
One question is over the role of fats and carbohydrates on liver fat content. Generally, low-carb weight reduction diets are not superior to low-fat diets for liver fat loss, and the patients can freely choose which diet they are capable of maintaining in the long term. A meta-analysis of 11 short-term small sample trials found no significant difference in the percentage change of hepatic fat reduction between low carb and low fat diets.82

Evidence is mounting on the importance of the type of fat eaten, pointing towards the specific harmful effect of saturated fat. Short-term trials83 84 suggest that either low-fat +high carb or high-fat + low carb diets reduce liver fat on a weight-loss diet leading to weight reduction. However, for an isocaloric or hyperenergetic diet high-(mostly saturated) fat + low carb results in increased liver fat, while low-fat + high carb reduces liver fat (if isocaloric) or increases liver fat to a much lower extent (if hyperenergetic).

It can be concluded that the type of fat makes a difference to liver fat. Saturated fats consistently increase liver fat more than polyunsaturated or monounsaturated fat, when total energy intake is similar. This evidence supports the benefits of the Mediterranean diet which is low in saturated fat and high in polyunsaturated and monounsaturated fat.

Ultraprocessed food and drink is a major challenge
The consumption of ultraprocessed foods (UPF) has increased dramatically, accounting for 50%–60% of total daily energy intake in some countries.85 These foods are a major source of added sugar, saturated fats, are energy dense and have low nutritional value.

Consumption of UPFs increases cardiovascular disease, cancer incidence and all-cause mortality.85 103 In a UK Biobank study, including 21 730 participants, UPF consumption was associated with type 2 diabetes incidence.86

A meta-analysis of nine cross-sectional and three cohort studies found higher consumption of UPFs resulted in more overweight, obesity and abdominal obesity, with a clear dose–response association.87

The same relationship can be seen with metabolic syndrome both in the general population and among people with NAFLD. In a cross-sectional study of 789 people, among people with NAFLD, those getting 40% or more of their calories from UPFs had more than three times the chance of having metabolic syndrome.88

Higher consumption of UPFs is linked to food insecurity and low income, since healthy food is more expensive. Indeed, NAFLD is almost 10% more common among those with poor food security.89

One target to reduce NAFLD may be sugar-containing beverages such as fruit juice and sugar-added soft drinks. A cohort study of 1940 infants found that consumption of more than two sugar-containing beverages a day at 1 year of age increased the chances of MRI-assessed NAFLD at age 10 threefold, compared with those consuming fewer than one beverage a day.90

Looking beyond diet, physical activity protects from liver-related mortality and seems to lower the risk conveyed by adiposity.91 Both low physical activity and sedentary behaviour (8 hours and above of sitting time per day) raise the risk of NAFLD, independent of each other.92

NASH therapies in phase III
Key points
► A plethora of drugs are in various stages of research, but only a few have currently entered in phase III.
► Despite promising results so far, outcomes sufficient for full regulatory approval may still be years away.

Despite the rising prevalence and serious potential clinical consequences of NASH, there are currently no treatments licensed for the disease. Treatment relies on lifestyle changes, primarily weight loss.

The complexity of the pathophysiology of the disease provide multiple potential target for drug treatment (figure 4). The FDA endpoints for NASH clinical trials in late stage development focus on the histological endpoints of NASH resolution

Figure 3 Lifestyle recommendations for patients with NAFLD. BMI, body mass index; NAFLD, non-alcoholic fatty liver disease.
Recent advances in clinical practice

without worsening of fibrosis or fibrosis improvement of at least one fibrosis stage without worsening of steatohepatitis. Thus, a repeat liver biopsy at entry and at the end of treatment is required.93

Drugs acting to restore metabolic homeostasis

Peroxisome proliferator-activated receptors (PPARs) are ligand-activated transcription factors of nuclear hormone receptor superfamily comprising of the following three subtypes: PPARα, PPARγ and PPARβ/δ. Activation of PPAR-α reduces triglyceride level and is involved in regulation of energy homeostasis, activation of PPAR-γ causes insulin sensitisation and enhances glucose metabolism, whereas activation of PPAR-β/δ enhances fatty acids metabolism.94

The pan-PPAR agonist lanifibranor has successfully completed a 24-week phase IIb trial with 247 participants, meeting its primary endpoint of a reduction of two points or more on the SAF activity score, with no increase in fibrosis, achieved by 49% of patients on 1200 mg compared with 27% on placebo.95 The drug also hit its secondary endpoint of reducing fibrosis by at least one stage without worsening NASH (42% on 1200 mg vs 24% on placebo).

Lanifibranor is generally well tolerated, although side effects include mild weight gain. It has been given FDA breakthrough therapy designation as the lead drug candidate in this class and a phase III study is underway.

GLP1-receptor agonists

GLP1-receptor agonists have multiple effects on several organs and systems, including the pancreas,96 the central nervous system and the liver.97,98

The GLP1 RA semaglutide has completed a 72-week phase II trial with 320 participants, showing resolution of NASH with no worsening of fibrosis for 56% of patients on 0.4 mg compared with 20% on placebo.99 It was unable to hit its secondary outcome of improvement of fibrosis with no worsening of NASH, although the number of patients with worsening fibrosis was lower in the active treatment arms. The drug induces a significant weight loss. Most common adverse events were gastrointestinal, while the safety in NASH was consistent with the observed profile in other trials and disease areas. A phase III trial has already started.

Drugs acting on the liver

Thyroid hormone receptor beta (THR-β) is crucial for liver homeostasis, through multiple metabolic actions of thyroid hormones. THR-β agonists have been shown to improve lipid metabolism.100

The THR-β agonist resmetirom has completed a phase II trial and is in two phase III 52-week trials. The 36-week phase II study with 125 participants showed reduction of liver fat by an average 30% compared with baseline by 12 weeks, leading to significant resolution of NASH.101 Preliminary results from one of the phase III trials demonstrates positive results in imaging measurements of liver fat and liver stiffness, as well as reduction in LDL-cholesterol and ApoB.102

Obeticholic acid is a selective FXR agonist currently tested in a phase III trial of 1968 participants with NASH and fibrosis F2–F3.104 In an 18-month interim analysis, the drug met the endpoint of improvement by at least one stage in fibrosis with no worsening of NASH, but did not meet the endpoint of NASH resolution. The main adverse event was pruritus and increase in LDL-cholesterol, responsive to statin therapy.

Aramchol, a partial inhibitor of hepatic stearoyl-CoA desaturase (SCD1) has been tested in a phase 2b trial including 247 patients with NASH.105 Aramchol at the highest dose of 600

Figure 4  Potential therapeutic targets for NASH, updated and adapted from Konerman et al.110

Dufour J-F, et al. Gut 2022;71:2123–2134. doi:10.1136/gutjnl-2021-326874
mg did not reach the prespecified significance level for decrease in liver fat by MR spectroscopy at 52 weeks. However, post hoc analyses suggest a potential for improving liver histology in patients with high disease activity and precirrhotic stages of fibrosis.

The next 6 years
Five phase III studies are expected to complete the trial part having surrogate, histological endpoints before the end of 2024 (aramchol, resmetiron, obeticholic acid, belapectin, lanifibranor, efruxifermin). Completion of clinical outcomes trials are expected in September 2025 for obeticholic acid and in May 2028 for semaglutide. Clinicians will need to be ready to assess which patients to treat and how to treat and monitor treatment, when these drugs finally become available.

Historically, many drugs have failed to hit their histological endpoints.

NASH therapies in Phase II
Key points

► Drugs under development rely on achieving two primary endpoints—resolution of NASH and reduced fibrosis—but many other targets are likely to be beneficial.

► Combinations of drugs may allow targeting of both metabolic dysfunction and liver damage, with injectable or infusion drugs used to initiate therapy in more severe disease and oral drugs providing long-term maintenance therapy.

Key histological endpoints for clinical trials are resolution of NASH (without worsening fibrosis) and reduced fibrosis (without worsening NASH).106

Recent data prove that improvement in fibrosis stage translates into an improvement in clinical outcome.107 We know from studies of bariatric surgery that NASH is reversible, and that as NASH reverses, so fibrosis improves.108 109

The multifactorial nature of NASH means that there are a plethora of targets beyond histological resolution of NASH and reduction in fibrosis which could be beneficial. These could include reduction in lipotoxic fat, weight loss, atherogenic lipid improvement and glycaemic control. The question is whether one drug will be enough to target all these potential endpoints.

The routes that lead to NASH and fibrosis are complex, taking in metabolic and inflammatory pathways, meaning there is a multitude of targets.110 Compounds currently being tested in phase II can be divided into those that aim at five main targets:

► Insulin resistance and lipid metabolism.

► Lipotoxicity and oxidative stress.

► Inflammation and immune activation.

► Cell death.

► Fibrogenesis and collagen turnover.

By far the majority of drugs in development aim at insulin and lipids, although some drugs have multiple effects and can be aimed at more than one target.

Many drugs are still in early phase II studies, looking at non-invasive endpoints, while there are a host of others in later phase II, collecting biopsies for histopathological endpoints that will allow them to progress to phase III. The ability of histopathology to accurately identify endpoints in clinical trials is likely to have a major impact on the future development of this field.

The majority of agents are oral, but there are a significant number of injectable/infusion agents under development, including all hormone therapies. Their place in therapy needs consideration.

Five of the compounds in phase II (efruxifermin(FG-21), pegbelfermin (FG-21), aldafermin (FG-19), pegozafermin (FG-21) and BFK8588A (FG-21)) achieved a reduction in ALT which has been shown to be associated with histological improvement in NAFLD, and these results are comparable to those seen in drugs now in phase III. Four phase II compounds (MET-409 (FXR agonist), pegozafermin, efruxifermin, VK2809 (THR-beta agonist)) show a 30% clinically significant reduction in MRI-PDFF. Two phase II drugs (efruxifermin, aldafermin) show some data indicating improvement in fibrosis, although in the case of aldafermin, later histopathology did not support this.

The same two drugs showed NASH resolution, although in the case of efruxifermin, this was undermined by NASH-resolving weight loss from one of only two patients in the placebo group who had a liver biopsy.

The combination of NASH resolution and fibrosis improvement gives a clearer picture: efruxifermin and aldafermin both demonstrate significant results, with 28% of efruxifermin111 and 22% of aldafermin112 patients achieving both endpoints, compared with no placebo patients. Minimising placebo response by use of duel endpoints may point a way forward.

Both efruxifermin and aldafermin are injectables, with potentially potent effects on histopathology and also metabolic parameters. Practical considerations may limit their use, including poor gastrointestinal tolerability, common to all injectable and infusion drugs in this area. Injection and infusion drugs might be best used for short-term induction therapy for patients with F3 or F4, or for patients with F2 showing risk factors for rapid fibrosis progression or significant metabolic comorbidity.

Better-tolerated oral drugs could then be used for long term treatment or maintenance therapy in people with F1–F3, perhaps in a fixed dose combination.

Combination therapy is likely the future. Some patients will benefit more from a focus on therapies to target metabolic syndrome, with secondary drugs targeting anti-inflammatory and antifibrotic endpoints, while for others with more advanced disease, the position will be reversed. A challenge will be to identify which patients are likely to respond best to which therapies, with the possibility of more personalised medicine in future.

Several phase II trials of combination therapies are in progress, offering the possibility for synergies between drugs.

Cancers and NASH

Key points

► NASH increases the risk of hepatocellular carcinoma (HCC) and of other cancers.

► About half of people with NASH-related HCC do not have cirrhosis and tend to be diagnosed at an advanced cancer stage.

Cardiovascular disease has long been the major cause of death for people with NAFLD,113 reflecting its close links to metabolic disorders. The second cause of death of NAFLD patients is cancer.

A Korean study found an incidence rate of 8.5 per 100 000 years for HCC, compared with 11.9 for breast cancer and 46.2 for colorectal cancer.114 However, HCC is more common among people with NAFLD—the same study showed an incidence rate of 23.1 for people with NAFLD, compared with 0.9 for people without. HCC is not the only cancer of concern. Studies have found people with NAFLD have an increased risk of colorectal cancers, cholangiocarcinoma and cancers of the breast, stomach, pancreas, prostate and oesophagus.115
After adjustment for risk factors, NAFLD conveyed a 17-fold increase in risk of HCC, and also a doubling of risk for breast cancer. This is not simply a reflection of obesity. A US case-control study of 4722 cases of NAFLD showed a significant increase in risk for people with NAFLD, even compared with controls with obesity.

Researchers have identified a novel molecular signature found in HCC associated with NASH. This suggests a specific risk for this type of HCC, not found in non-NASH populations.

One meta-analysis calculated that each unit increase in liver stiffness was associated with an 11% increased risk of HCC, although this included all types of liver disease, not just NAFLD.

While cirrhosis with NAFLD conveys the highest risk of HCC, that does not mean all NAFLD-related HCC patients have cirrhosis. One study of 4406 cases found only 46% had known cirrhosis at time of diagnosis. However, the absolute incidence in non-cirrhotic patients is still low compared with cirrhotic NAFLD patients: incidence for cirrhosis 13.55 or 4.82 per 1000 person-years vs non-cirrhosis 0.39 and 0.04 (each with and without high FIB-4).

Patients with NAFLD are diagnosed at a later stage of HCC than patients with HCV. In a prospective observational study from Italy, 11% of HCV patients with HCC were diagnosed at BCLC stage 0, compared with none of the NAFLD patients. More NAFLD patients were diagnosed at stage B or C.

In post hoc analysis of phase 3 trials testing immunotherapy for HCC, patients with virus-related HCC seemed to fare better than patients with non-viral-related cancer. This was also observed in retrospective analysis of small cohorts.

In addition to the risk factors listed above, some medications lower the risk of developing liver cancer:
- Metformin lowers risk.
- Statins (particularly lipophilic statins such as simvastatin, atorvastatin) decrease risk.
- Aspirin lowers risk.

Lifestyles are also important to mitigate risk. Coffee consumption and regular physical activity are associated with a reduction in the risk of developing HCC. Can this risk be reversed, for example, in the case of bariatric surgery? A study of 98 090 newly diagnosed NAFLD patients with severe obesity, of whom 34% received bariatric surgery, suggests it can. The adjusted risk of any cancer was reduced by 18% after bariatric surgery, and obesity-related cancer risk fell by 25%.

CONCLUSIONS
The challenge of NASH remains to license a treatment for this increasingly prevalent and important condition. There is no shortage of candidate drugs, but difficulties in diagnosis, staging and monitoring the effects of treatment have added unprece-dented complexity to the field.

Technologies to develop accurate, specific and meaningful imaging and circulating biomarkers, along with improvements in reading of biopsies, should allow for better interpretation of drug trial results and likely new drug licenses. Meanwhile, it is important not to forget the real differences that lifestyle measures can make to NASH. This is the reason why it is so important to screen patients at risk.

Correction notice This article has been corrected since it published Online First. Typographical errors have been corrected in the abstract.

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Acknowledgements This paper was written following a virtual round table event organised by BMJ Journals, supported by an unrestricted educational grant from Intercept Pharmaceuticals and Siemens Healthineers. The sponsors had no control or influence over the selection of speakers, the round table or the writing of the paper. Anna Sayburn, an independent professional medical writer, facilitated the meeting and provided medical writing support.

Contributors J-FD elaboration and coordination, writing specific sections and critical review of the manuscript. All the others writing of specific sections and critical review of the manuscript.

Funding RL receives funding support from NCATS (SULTR01442), NIDDK (U01DK061734, U01DK130190, R01DK106419, R01DK121378, R01DK124318, P30DK120515), NHLBI (P01HL147835), and NIAAA (U01AA029019). Other authors have not declared a specific grant for this research from any funding agency in the public, commercial or not-for-profit sectors.

Competing interests J-FD: Advisory committees: Abbvie, Alientis, Allergan, Axcella, Bayer, Bristol-Myers Squibb, Eisai, Enyo Pharma, Falk, Genfit, Gilead Sciences, HepaRegenX, Intercept, Ipsen, Inventiva, Madrigal, Merck, Novartis, Pfizer, Roche, Trillium DX. Speaking and teaching: Bayer, Intercept, Genfit, Royalties: BMJ Journals, Springer, UpToDate. Quentin Ansee: Research Grant Funding: Abbvie, Allergan/Tobira, AstraZeneca, GlaxoSmithKline, GlympseBio, Novartis Pharma AG, Pfizer. Active Research Collaborations (including research supported through the EU IM2 LITMUS Consortium*). Abbvie, Antars Medical*, Allergan/Tobira, AstraZeneca, Boehringer Ingelheim International*, Ella Med, Getz, GlaxoSmithKline, Ipsen, Merck, Novartis, Pfizer, Roche, Trillium DX. Speaking and teaching: Bayer, Intercept, Genfit, Novartis.

REFERENCES
1 Younossi ZM, Stepanova M, Ong J, et al. Nonalcoholic steatohepatitis is the most rapidly increasing indication for liver transplantation in the United States. Clin Gastroenterol Hepatol 2021;19:580–9.
Recent advances in clinical practice

2 Younossi ZM, Koenig AB, Abdelatif D, et al. Global epidemiology of nonalcoholic fatty liver disease-Meta-analytic assessment of prevalence, incidence, and outcomes. *Hepatology* 2016;64:73–84.

3 NCD Risk Factor Collaboration (NCD-RisC). Trends in adult body-mass index in 200 countries from 1975 to 2014: a pooled analysis of 1698 population-based measurement studies with 19.2 million participants. *Lancet* 2016;387:1377–96.

4 Dufour JF, Scherer PE. The global epidemiology of nonalcoholic steatohepatitis (NASH) and associated risk factors– a targeted literature review. *Endocrine and Metabolic Science* 2021;3:10089.

5 Li J, Zou B, Yeo YH, et al. Prevalence, incidence, and outcome of non-alcoholic fatty liver disease in Asia, 1999-2019: a systematic review and meta-analysis. *Lancet Gastroenterol Hepatol* 2019;4:389–98.

6 Knook R, Choi KC, Wong GL-H, et al. Screening diabetic patients for non-alcoholic fatty liver disease with controlled attenuation parameter and liver stiffness measurements: a prospective cohort study. *Gut* 2016;65:1359–68.

7 Wong VW-S, Chan W-K, Chitturi S, et al. Asia-Pacific Working Party on non-alcoholic fatty liver disease guidelines 2017-Part 1: definition, risk factors and assessment. *J Gastroenterol Hepatol* 2018;33:70–85.

8 Ye Q, Zou B, Yeo YH, et al. Global prevalence, incidence, and outcomes of non-obese or lean non-alcoholic fatty liver disease: a systematic review and meta-analysis. *Lancet Gastroenterol Hepatol* 2020;5:739–52.

9 Wong VW-S, Wong GL-H, Yeung DK-W, et al. Incidence of non-alcoholic fatty liver disease in Hong Kong: a population study with paired proton-magnetic resonance spectroscopy. *J Hepatol* 2015;62:182–9.

10 Lee HW, Wong GL-H, Knook R, et al. Serial transient elastography examinations to monitor patients with type 2 diabetes: a prospective cohort study. *Hepatology* 2020;72:1230–41.

11 European association for the study of the liver, European association for the study of diabetes, European association for the study of obesity, EASL-EASD-EASO clinical practice guidelines for the management of non-alcoholic fatty liver disease. *J Hepatol* 2016;64:1388–402.

12 Kanwal F, Shubrok JH, Adams LA, et al. Clinical care pathway for the risk stratification and management of patients with nonalcoholic fatty liver disease. *Gastroenterology* 2021;161:1657–69.

13 Younossi Z, Stepanova M, Ong JP. Global nonalcoholic steatohepatitis council. Nonalcoholic steatohepatitis is the fastest growing cause of hepatocellular carcinoma in liver transplant candidates. *Clin Gastroenterol Hepatol* 2019;17:748–55.

14 Estes C, Anstee QM, Arias-Loste MT, et al. Modeling NAFLD disease burden in China, France, Germany, Italy, Japan, Spain, United Kingdom, and United States for the period 2016-2030. *J Hepatol* 2018;69:896–904.

15 Villar-Gomez E, Caladzill-Bertot L, Wai-Sun Wong V, et al. Fibrosis severity as a determinant of cause-specific mortality in patients with advanced nonalcoholic fatty liver disease: a Multi-National cohort study. *Gastroenterology* 2018;155:443–57.

16 NCD Risk Factor Collaboration (NCD-RisC). Worldwide trends in body-mass index, underweight, overweight, and obesity from 1975 to 2014: a pooled analysis of 216 population-based measurement studies in 188 9 million children, adolescents, and adults. *Lancet* 2017;390:2627–42.

17 Eslam M, Newsome PN, Sarin SK, et al. A new definition for metabolic dysfunction-associated fatty liver disease: an international expert consensus statement. *J Hepatol* 2020;73:202–9.

18 Wong VW-S, Wong GL-H, Woo J, et al. Impact of the new definition of metabolic associated fatty liver disease on the epidemiology of the disease. *Clin Gastroenterol Hepatol* 2019;17:2161–71.

19 Tilg H, and Moschen, a, R. Evolution of inflammation in nonalcoholic fatty liver disease: The multiple parallel hits hypothesis. *Hepatology* 2010;52:1836–46.

20 Febril B, Karin M. “Sweet death”: Fructose as a metabolic toxin that targets the gut-liver axis. *Cell Metab* 2021;33:2316–28.

21 Teppe E, Valentin L, Lute E. NAFLD oncopanels: from new variants to the clinic. *J Hepatol* 2020;72:1196–209.

22 Tilg H, Moschen AR. Insulin resistance, inflammation, and non-alcoholic fatty liver disease. *Trends Endocrinol Metab* 2008;19:371–9.

23 Tilg H, Moschen AR, Roden M. NAFLD and diabetes mellitus. *Nat Rev Gastroenterol Hepatol* 2017;14:32–42.

24 Dalal PS, Singh S, Patel J. Increased risk of mortality by fibrosis stage in nonalcoholic fatty liver disease: systematic review and meta-analysis. *Hepatology* 2017;65:1557–65.

25 Taylor RS, Taylor RJ, Bayliss S, et al. Association between fibrosis stage and outcomes of patients with nonalcoholic fatty liver disease: a systematic review and meta-analysis. *Gastroenterology* 2020;158:1611–25.

26 Moschen AR, Mohlar C, Enrich B, et al. Adipose and liver expression of interleukin (IL)-1 family members in morbid obesity and effects of weight loss. *Mol Med* 2016;11:840–5.

27 Kamary Y, Shaish A, Vax E, et al. Lack of interleukin-1x or interleukin-1β inhibits transformation of steatosis to steatohepatitis and liver fibrosis in hypercholesterolemic mice. *J Hepatol* 2011;55:1086–94.
56 Mózes FE, Lee JA, Selvaraj EA, et al. Diagnostic accuracy of non-invasive tests for advanced fibrosis in patients with NAFLD: an individual patient data meta-analysis. Gut 2022;71:1006–1019.

57 Wong VW-S, Vergniol J, Wong GL-H, et al. Diagnosis of fibrosis and cirrhosis using liver stiffness measurement in nonalcoholic fatty liver disease. Hepatology 2010;51:454–62.

58 Siddiqui MS, Vuppalanchi R, Van Natta ML, et al. Vibration-Controlled transient elastography to assess fibrosis and steatosis in patients with nonalcoholic fatty liver disease. Clin Gastroenterol Hepatol 2019;17:156–63.

59 Eddowes PJ, Sasso M, Allison M, et al. Accuracy of FibroScan controlled attenuation parameter and liver stiffness measurement in assessing steatosis and fibrosis in patients with nonalcoholic fatty liver disease. Gastroenterology 2019;156:1711–20.

60 Vai Y, Lee J, Boursier J, et al. Enhanced liver fibrosis test for the non-invasive diagnosis of fibrosis in patients with NAFLD: a systematic review and meta-analysis. J Hepatol 2020;73:252–62.

61 Newsome PN, Crumb R, Davison SM, et al. Guidelines on the management of abnormal liver blood tests. Gut 2018;67:6–19.

62 Boyle M, Timakis D, Schattenberg JM. Performance of the PRO-C3 collagen neo-epitope biomarker in non-alcoholic liver disease. J Hepatol 2019;71:189–98.

63 Daniels SJ, Leeming DJ, Eslam M. ADAPT: an algorithm incorporating pro-C3 accurately identifies patients with NAFLD and advanced fibrosis. Hepatology 2019;69:1075–86.

64 Srivastava A, Gailer R, Tannar S, et al. Prospective evaluation of a primary care referral pathway for patients with non-alcoholic fatty liver disease. J Hepatol 2019;71:371–8.

65 Lee JT et al. OS-243: diagnostic accuracy of blood-based biomarkers for staging fibrosis in NAFLD: phase 1 results of the litmus project. EAS2021.

66 Mayo R, Crespo J, Martinez-Arranz J, et al. Metabolomic-Based noninvasive serum test to diagnose nonalcoholic steatohepatitis: results from discovery and validation cohorts. Hepatol Commun 2018;2:807–8.

67 Govaere O, Cockell S, Timakis D, et al. Transcriptomic profiling across the nonalcoholic fatty liver disease spectrum reveals gene signatures for steatohepatitis and fibrosis. Sci Transl Med 2020;12:eaba448.

68 Cazanave SC, Warren AD, Paucia M, et al. Peptide-Based urinary monitoring of fibrotic nonalcoholic steatohepatitis by mass-barcode activity-based sensors. Sci Transl Med 2021;13:eabbe939.

69 Dalui PS, Srin CB, Loomra R. MRE and MRI for non-invasive quantitative assessment of hepatic steatosis and fibrosis in NAFLD and NASH: clinical trials to clinical practice. J Hepatol 2016;65:1006–16.

70 Tapper EB, Loomra R. Noninvasive imaging biomarker assessment of liver fibrosis by elastography in NAFLD. Nat Rev Gastroenterol Hepatol 2018;15:274–82.

71 Boursier J, Vergniol J, Guillet A, et al. Diagnostic accuracy and prognostic significance of blood tests and liver stiffness measurement by FibroScan in non-alcoholic fatty liver disease. J Hepatol 2016;65:570–8.

72 Loomba R, Wolfson T, Ang B, et al. Magnetic resonance elastography predicts advanced fibrosis in patients with nonalcoholic fatty liver disease: a prospective study, Hepatology 2014;60:1920–8.

73 Newsome PN, Sasso M, Deeks J, et al. FibroScan-AST (FAST) score for the non-invasive identification of patients with non-alcoholic steatohepatitis with significant activity and fibrosis: a prospective derivation and global validation study. Lancet Gastroenterol Hepatol 2020;5:362–73.

74 Nueddin N, Alkhouri N, Brown KA, et al. Driving nonalcoholic steatohepatitis forward using the FibroScan aspartate aminotransferase score, but obey the traffic lights. Hepatology 2020;72:2228–30.

75 Jung J, Loomra RR, Imaji K, et al. MRE combined with RBI-4 (MEFIB) index in detection of candidates for pharmacological treatment of NASH-related fibrosis. Gut 2021;70:1946–53.

76 Tamaki N, Higuchi M, Kurozaki M, et al. Risk difference of Liver-Related and cardiovascular events in liver fibrosis status in nonalcoholic fatty liver disease. Clin Gastroenterol Hepatol 2022;20:1171–1173.e2.

77 Hsu C, Caussy C, Imaji K, et al. Magnetic resonance vs transient elastography analysis of patients with nonalcoholic fatty liver disease: a systematic review and pooled analysis of individual participants. Clin Gastroenterol Hepatol 2019;17:630–7.

78 Sline JG, Munagunuru N, Barad A, et al. Change in MRI-PDFF and histologic response in patients with nonalcoholic steatohepatitis: a systematic review and meta-analysis. Clin Gastroenterol Hepatol 2021;19:2274–83.

79 Patel J, Bettencourt R, Cui J, et al. Association of noninvasive quantitative decline in liver fat content on MRI with histologic response in nonalcoholic steatohepatitis. Therap Adv Gastroenterol 2016;9:692–701.

80 Yaskolka Meir A, Rinott E, Tsaaban G, et al. Effect of green-Mediterranean diet on intrahepatic fat: the DIRECT PLUS randomised controlled trial. Gut 2021;70:2085–95.

81 Salomone F, Ivanovský-Wajcman D, Fliss-Isakov N, et al. Higher phenolic acid intake independently associates with lower prevalence of insulin resistance and non-alcoholic fatty liver disease. J Hepatol 2020;2:100069.
Recent advances in clinical practice

113 Angulo P, Kleiner DE, Dam-Larsen S, et al. Liver fibrosis, but no other histologic features, is associated with long-term outcomes of patients with nonalcoholic fatty liver disease. *Gastroenterology* 2015;149:389–97.

114 Kim G-A, Lee HC, Choe J, et al. Association between non-alcoholic fatty liver disease and cancer incidence rate. *J Hepatol* 2017;50:168–82:78:32:94–8.

115 Liu S-M, Ma X-F, Zhao J, et al. Association between non-alcoholic fatty liver disease and extrahepatic cancers: a systematic review and meta-analysis. *Lipids Health Dis* 2020;19:118.

116 Kim G-A, Lee HC, Choe J, et al. Association between non-alcoholic fatty liver disease and cancer incidence rate. *J Hepatol* 2017. doi:10.1016/j.jhep.2017.09.012. [Epub ahead of print: 02 Nov 2017].

117 Allen-AM, Hicks SB, Mara KC, et al. The risk of incident extrahepatic cancers is higher in non-alcoholic fatty liver disease than obesity - A longitudinal cohort study. *J Hepatol* 2019;71:1229–36.

118 Pinyol R, Torrecilla S, Wang H, et al. Molecular characterisation of hepatocellular carcinoma in patients with non-alcoholic steatohepatitis. *J Hepatol* 2021;75:865–78.

119 Singh S, Fuji LL, Murad MH, et al. Liver stiffness is associated with risk of decomposition, liver cancer, and death in patients with chronic liver diseases: a systematic review and meta-analysis. *Clin Gastroenterol Hepatol* 2013;11:1573–84.

120 Sanyal A, Poklepovic A, Moyneur E, et al. Population-Based risk factors and resource utilization for HCC: US perspective. *Curr Med Res Opin* 2010;26:2183–91.

121 Kanw J, Kramer JR, Mapakchi S, et al. Risk of hepatocellular cancer in patients with non-alcoholic fatty liver disease. *Gastroenterology* 2018;153:1828–37.

122 Piscaglia F, Svegliati-Baroni G, Barchetti A, et al. Clinical patterns of hepatocellular carcinoma in nonalcoholic fatty liver disease: a multicenter prospective study. *Hepatology* 2016;63:827–38.

123 Pfister D, Núñez NG, Pinyol R, et al. NASH limits anti-tumour surveillance in immunootherapy-treated HCC. *Nature* 2021;592:450–6.

124 Singh S, Singh PF, Singh AG, et al. Anti-Diabetic medications and the risk of hepatocellular cancer: a systematic review and meta-analysis. *Am J Gastroenterol* 2013;108:881–91.

125 Simon TG, Duberg A-S, Aleman S, et al. Lipophilic statins and risk for hepatocellular carcinoma and death in patients with chronic viral hepatitis: results from a nationwide Swedish population. *Ann Intern Med* 2019;171:318–27.

126 Simon TG, Duberg A-S, Aleman S, et al. Association of aspirin with hepatocellular carcinoma and liver-related mortality. *N Engl J Med Overseas Ed* 2020;382:1018–28.

127 Rustgi VK, Li Y, Gupta K, et al. Bariatric surgery reduces cancer risk in adults with nonalcoholic fatty liver disease and severe obesity. *Gastroenterology* 2021;161:171–84.

128 Sheth SG, Flamm SL, Gordon FD, et al. AST/ALT ratio predicts cirrhosis in patients with chronic hepatitis C virus infection. *Am J Gastroenterol* 1998;93:44–8.

129 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.

130 Shah AG, Lydecker A, Murray K, et al. Comparison of noninvasive markers of fibrosis in patients with nonalcoholic fatty liver disease. *Clin Gastroenterol Hepatol* 2009;7:1104–12.

131 Mozes FE, Lee JA, Selvaraj EA, et al. Diagnostic accuracy of non-invasive tests for advanced fibrosis in patients with NAFLD: an individual patient data meta-analysis. *Gut* 2022;71:1006–19.

132 Guha IN, Parkes J, Roderick P, et al. Noninvasive markers of fibrosis in nonalcoholic fatty liver disease: validating the European liver fibrosis panel and exploring simple markers. *Hepatology* 2008;47:455–60.

133 Ratmansky 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. *BMJ Gastroenterol* 2006;6:6.

134 Vai Y, Lee J, Boursier J, et al. FibroTest for evaluating fibrosis in non-alcoholic fatty liver disease patients: a systematic review and meta-analysis. *J Clin Med* 2021;10. doi:10.3390/jcm10122415. [Epub ahead of print: 29 05 2021].

135 Caleś P, Laine F, Boursier J, et al. Comparison of blood tests for liver fibrosis specific or not to NAFLD. *J Hepatol* 2009;50:165–73.

136 Van Dijk A-M, Vai Y, Maak AL, et al. Systematic review with meta-analyses: diagnostic accuracy of FibroMeter tests in patients with non-alcoholic fatty liver disease. *J Clin Med* 2021;10. doi:10.3390/jcm10132910. [Epub ahead of print: 29 06 2021].

137 Daniels SJ, Leeming DJ, Eslam M, et al. ADAPT: an algorithm incorporating pro-C3 accurately identifies patients with NAFLD and advanced fibrosis. *Hepatology* 2019;69:1075–86.

138 Boyle M, Tkachuk D, Schattenberg JM, et al. Performance of the PRO-C3 collagen neo-epitope biomarker in non-alcoholic fatty liver disease. *HEP Rep* 2019;1:186–98.

139 Mak AL, Lee J, van Dijk A-M, et al. Systematic review with meta-analysis: diagnostic accuracy of pro-C3 for hepatic fibrosis in patients with non-alcoholic fatty liver disease. *Biomedicines* 2021;9. doi:10.3390/biomedicines9121920. [Epub ahead of print: 15 12 2021].