CURRENT PATHOLOGY REPORTING IN NAFLD/

As mentioned above, the reliance on focused

scoring systems alone has several potential weak-

nesses. First, there is no requirement to comment on sample adequacy unless the sponsor and/

or the pathologists involved specified this in the

protocol. As liver biopsy specimens typically repre-

sent approximately 1/50 000th of the entire liver

volume, inadequate biopsy length can greatly affect the quality of assessment.^{16 17} It is generally recom-

mended that all medical liver biopsies should be at

least 25 mm long and of sufficient width (~1.6 mm)

to include at least 10 portal tracts.¹⁸¹⁹ Therefore,

comments on sample adequacy should be required.

one connective tissue stain,^{9 12} deviating from what

medical societies, including the UK Royal College

of Pathologists (RCPath) and the American Associ-

ation for the Study of Liver Diseases, have outlined

as robust practice in clinical care.^{20 21} Up to seven

stains are recommended as certain features cannot

be identified without a particular stain (eg, Shika-

ta's orcein for long-standing cholestasis and a Perls'

stain for iron), any of which could be highly rele-

vant to assessing a patient's response in a trial.

Furthermore, as biopsies are often processed and

stained at multiple sites, the quality of processing

and/or staining should be noted due to difficulties

in assessing disease features if these are suboptimal.

Requiring comments on the quality and utility of

Third, focusing solely on four features alone and

ignoring others can cause major problems. This

narrow focus was implicated in the placing of a

clinical hold on a recent drug trial in which biopsies

were subsequently found to have features atypical

for NASH. It was unclear whether these subjects

had newly developed liver injury or had pre-existing changes and should have been excluded from the

trial in the first place.¹³ ¹⁵ ²² Another major reason

for noting such abnormalities is that response to the

trial drug may be influenced by their presence and

therefore may be relevant to the trial's outcome.

Within the current system, aside from the possibility

that some features will not be recognised without

the appropriate stain, if there are any additional

pathological features, the expectation is that these

will be noted in the 'comment' section. However, in

the absence of any 'comment', it is unclear if there

are no such additional findings, or that there are

but were not detected as a result of the pathologist

staining would address these issues.

Second, trials typically require only H&E and

NASH CLINICAL TRIALS



Improved pathology reporting in NAFLD/NASH for clinical trials

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INTRODUCTION Non-alcoholic fatty liver disease (NAFLD), a

disorder characterised by pathological accumu-

lation of non-visible free fatty acids and visible

triglyceride in hepatocytes, is on the rise globally

in both adult and paediatric populations.¹ Evidence

suggests that 20%-50% of the European Union

and US populations exhibit features of NAFLD,²

driven by higher rates of obesity, insulin resistance

and type 2 diabetes, and metabolic syndrome.³

Additionally, recognition of a growing number of

patients with 'lean NAFLD' who are not obese,

but have high levels of visceral fat, diets high in

fats and carbohydrates, or who have genetic risk

factors, has increased.⁴ Of patients with NAFLD,

6%-55% will have histological signs of non-

alcoholic steatohepatitis (NASH), which if left

unmanaged can lead to cirrhosis and potentially

NAFLD has surpassed viral hepatitis as the leading

cause of chronic liver disease worldwide.⁶ Estimates

suggest that by 2030 NAFLD will overtake hepa-

titis C as the primary cause of liver failure requiring

transplantation and that the number of NAFLD-

related deaths will increase by 178%.⁷ Annual

spending related to NAFLD care is estimated to rise

exponentially from \$103 billion to \$1.005 trillion

in the USA and from €35 billion to €334 billion in

Europe between 2016 and 2025.⁸ While bariatric

surgery and/or weight loss can be effective, they

present their own challenges in delivery. There are

presently no approved drugs on the market; hence,

the number of clinical trials has grown by approxi-

the European Medicines Agency require liver biopsy

for clinical trials as the 'gold standard', diagnosis

and monitoring rely on pathological assessment of a liver biopsy.⁹⁻¹¹ Most trials focus only on the

numerically reported values of the semiguantitative

assessment of four cardinal features of NAFLD/

NASH-steatosis, inflammation, hepatocellular

ballooning and fibrosis-as set out by the Pathology

Committee of the NASH Clinical Research

Network.¹² However, there are a number of poten-

tial problems (see next section), and a recent study

argued that poor reliability of liver biopsy evalu-

ation in NAFLD 'allows improper entry, misclas-

sification, and diminishes treatment effect'.¹³ Two

follow-up editorials from pathologists discussed

this further.^{14 15}

As both the Food and Drug Administration and

mately 60% over the last 10 years.

hepatocellular carcinoma.⁵

► Additional supplemental material is published online only. To view, please visit the journal online (http://dx.doi. org/10.1136/jclinpath-2021-207967).

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only examining for NAFLD Activity Score (NAS) components, or if the pathologist had observed them but assumed that trial sponsors do not require the information.

Fourth, as histopathology is subjective, interobserver and intraobserver variations in assessing features are inevitable.^{13 23} The fundamental cause of this variability is that definitions of NASH features are not sufficiently specific for different pathologists to identify them reliably. Requiring much more precise and agreed definitions for both recognition and quantification of the features will reduce the degree of variation. As an illustration of the potential, ensuring pathologists within a single trial agree on definitions of features before the start of a trial and/or assess biopsies simultaneously alleviates some of the issues.²⁴

SYNOPTIC REPORTING: COMPREHENSIVE, STRUCTURED REPORTING

To deal with and potentially overcome the above concerns, we propose the most appropriate method for NAFLD trial pathology reporting is the adoption of synoptic reporting, which ensures all features salient to the diagnosis and monitoring of liver disease/ injury and potential treatment effect/resolution are reported.

Synoptic reporting has been adopted in radiology and cancer pathology clinical practice to overcome two main inherent shortfalls of narrative reporting—failure to report all the relevant features and failure to provide a clear final message—resulting in misinterpretation or misreporting of disease features.^{25 26} Typically, synoptic reports offer three levels of organisation and standardisation: a structured format with paragraphs and subheadings; consistent organisation of subheadings, ensuring all required features are described in a logical order; and standardised language and terminology, which enhances the accessibility of reports to non-specialists and reduces ambiguity.²⁷ A key component is the required minimum set of features (data set or checklist) which are judged to be crucial to the accurate assessment of the patient and as such must be addressed in the report.

The implementation of synoptic reporting in clinical practice has been supported by bodies including the RCPath, the Association of Directors of Anatomic and Surgical Pathology, and the College of American Pathologists.²⁸ The use of such mandatory reporting parameters has been shown to improve reporting accuracy and completeness across a range of subspecialties, with more complex studies benefiting most.^{29–32}

IMPLEMENTATION OF SYNOPTIC REPORTING IN NAFLD/ NASH TRIAL PATHOLOGY

We recognise there are several barriers to adoption in NAFLD trial pathology.

First, a key factor is the choice of which features to include/ not include in the standard report: the minimum data set. Two factors are required: good/acceptable reproducibility through precise agreed definitions and clinical relevance through correlation of clinical outcome with each feature. There is no point including a feature which is highly clinically relevant but poorly reproducible; conversely there is no value in including a highly reproducible feature of no clinical relevance. Agreement on what constitutes the minimum data set and which features should be included is a major future task and broader than the argument made here for use of synoptic reporting. However, recommendations issued by bodies including the RCPath could form the basis on which synoptic reporting templates are built; online supplemental appendix 1 shows a potential example of a putative data set. It should be noted that using such a data set does not prevent scoring of the current four cardinal features.

Second, as the flexibility offered by narrative reporting is preferred by some pathologists—it facilitates communications of nuanced diagnoses or microscopic findings—it is important that the adoption of synoptic reports does not diminish a pathologist's ability to flag such findings. Accordingly, the design of the report must allow for this by including free text boxes for relevant comments.

Lastly, but not unimportantly, when considering a trial sponsor's adoption of synoptic reporting in NASH clinical trials, the related increased costs will prove some additional expense. While the burden of increased pathology costs may initially appear high, the financial impact of suboptimal reporting is greater.

CONCLUSIONS

It may be that the time has come to accept that reporting of liver biopsies from individuals with NAFLD who are entered into clinical trials can and should be improved. Although unquestionably invaluable, use of NAS as the only pathological endpoint in trials results in an incomplete assessment of liver disease features, both preintervention and postintervention, which in turn can undermine the outcome of the trial.¹³ To address this, we therefore propose the development, testing and adoption of a more comprehensive, structured reporting style—known as synoptic reporting—for use in clinical trials.

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