Systematic Review: Current Evidence in Non-Alcoholic Fatty Liver Disease Lacks Relevance to Patients with Advanced Fibrosis

Richard Parker 1
James Hodson2
Ian AC Rowe3

1. Centre for Liver Research, University of Birmingham United Kingdom
2. Statistics Department, University Hospitals Birmingham NHS Foundation Trust, Birmingham UK
3. Leeds Institute for Data Analysis, University of Leeds, UK

Corresponding author:
Dr Richard Parker
Centre for Liver Research
5th Floor IBR
University of Birmingham
Birmingham B15 2TT
United Kingdom
richardparker@nhs.net
(+44) (0) 7971 118036

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**Author contributions**

RP conceived the idea for this study, performed literature searches, data extraction, analysis and wrote the draft manuscript.

JH assisted with data analysis and reviewed the draft manuscript.

IAR conceived the idea for this study, data extraction and reviewed the draft manuscript.

All authors approved the final manuscript.
Abstract

Objectives
Epidemiological data have shown that individuals with advanced fibrosis are at greatest risk of premature morbidity in NAFLD. Individuals included in clinical trials are often highly selected to remove confounding factors but selection can introduce bias and limit external validity. We examined the external validity of trials in non-alcoholic fatty liver disease (NAFLD) by examining characteristics of participants in observational studies (OS) and randomised controlled trials (RCT) in NAFLD.

Design
A systematic review was performed with structured literature searches for relevant OS and RCT using PubMed and Ovid Embase (1948 - 2016). Identified studies were screened for inclusion by the authors and data extracted. Study populations were compared using t-tests to compare means and variances, in each case weighted by the size of individual studies. Dichotomous data were compared by Chi-squared test.

Results
In total 148 studies were included: 67 RCT and 81 OS including data from 44,860 individuals. Fifteen RCT participants differed from individuals in OS with regard to age, BMI, prevalence of DM, and gender (p<0.001 in each case). The most pronounced differences were seen between RCT participants and patients with advanced fibrosis. Co-morbid conditions prevalent amongst individuals with NAFLD were frequent exclusion criteria in RCT.

Conclusions
The characteristics of participants in randomised controlled trials differ to those of the wider population of individuals with NAFLD. These differences may reduce the utility of trial data to individuals with NAFLD at greatest risk of death.

Key words: Non-alcoholic fatty liver disease, Non-alcoholic steatohepatitis, Systematic review
Non-alcoholic fatty liver disease (NAFLD) is a spectrum of disease, ranging from simple hepatic steatosis through non-alcoholic steatohepatitis (NASH), to cirrhosis (1). NAFLD is common: hepatic steatosis is present in up to one-third of individuals (2) and NASH is seen in approximately 10% (3). Whilst the association of NAFLD per se with mortality is debated (4), large studies have shown that it is the subset of patients with advanced fibrosis who have an increased risk of liver-related morbidity as well as cardiovascular and neoplastic disease (5, 6). Given the high prevalence of NAFLD there is a need for effective therapies to prevent progression of disease and to treat established fibrosis. However, whilst many treatments have been trialed, few have shown categorical benefit in NASH (7).

The external validity of a trial describes its relevance to a population outside of the trial’s participants (8). Systematic analyses of trials in cardiology (9) and respiratory medicine (10, 11) have demonstrated have poor external validity, where commonly occurring medical conditions and age often exclude individuals from participation in trials (12). Poor external validity may promote ineffective treatments or, conversely, limit the acceptance of effective treatments (8). We used systematic review methods to assess the characteristics of participants included in randomised controlled trials (RCT) for treatment of NAFLD and the characteristics of individuals described in observational studies (OS) of NAFLD. This allowed comparison of the two groups and therefore an estimate of the external validity of RCTs performed in NAFLD to date.

**Methodology**

**Literature search**

Literature searching was undertaken using three search strategies, to identify OS for NAFLD, OS reporting advanced fibrosis, and to identify randomised controlled trials in NAFLD. PubMed/MedLine and Ovid Embase were searched (1948 - 2016), using the search terms: (((randomised controlled trial) AND non alcoholic steatohepatitis) OR
NASH) OR NAFLD) OR non-alcoholic fatty liver disease) for RCT, and (((Prevalence[Title]) OR Natural history[Title])) AND (((non-alcoholic fatty liver disease) OR NASH) OR NAFLD) OR non-alcoholic steatohepatitis) and (fibrosis OR histology) AND (NASH OR non-alcoholic steatohepatitis) for OS. Results were limited to human studies and those published in English. The literature search was performed 1/7/2014 and updated on 11/8/2016. To examine ‘grey literature’ for suitable studies the reference lists of included studies were searched for other suitable studies, and papers citing included studies were also reviewed. The title and abstract of papers found by the literature search were reviewed and unsuitable manuscripts or duplicate results excluded. Remaining papers were reviewed independently by two authors (RP, IAC) and disagreements resolved by consensus. This report is the only published account of this protocol.

Inclusion and exclusion criteria

Papers were included if they were full papers describing RCTs of interventions (lifestyle or pharmacological) in adult patients with any stage of NAFLD, or full papers reporting OS of prevalence or natural history of NAFLD in adults. Instances where the same cohort was described in more than one study were identified and included only once, with the study containing the most data included or the most recent report if descriptions were similar. Studies that had been published as abstracts, and those published in languages other than English were excluded. OS that only included particular groups, for example, studies reporting the prevalence of NAFLD amongst patients undergoing bariatric surgery, were also excluded.

Data extraction

Papers were reviewed and data extracted into a pre-prepared spreadsheet. Mean values for characteristics of trial participants in RCTs and individuals in OS were noted, specifically age, BMI, gender (% of male participants), prevalence of Diabetes mellitus (DM). For RCTs, type of intervention, primary outcome, secondary outcome and exclusion criteria were
recorded, as well as whether the trial reported a positive or negative finding. In the first instance, all OS were included in analysis of NAFLD as a whole, encompassing all stages of disease. Subsequent analyses were undertaken for biopsy-proven NASH and advanced fibrosis, including only OS that used and reported biopsy findings. For measurement of fibrosis in studies including biopsies, the Kleiner/Brunt classification was often used where F3 or F4 was taken to indicate advanced fibrosis, as is usual in the field. In studies that did not use the Kleiner/Brunt classification, advanced fibrosis was considered as bridging fibrosis or cirrhosis. The quality of included studies was assessed using the CONSORT guidelines for RCT (13) and the STROBE guidelines for observational studies (14).

Data synthesis and analysis

Many papers reported mean values for each arm of a trial but not an overall mean. Where this occurred, overall weighted means and variances were produced for the study as a whole, using the formulae \( \frac{\sum_i n_i \bar{x}_i}{\sum_i n_i} \) and \( \frac{\sum_i [n_i - 1] s_i^2}{\sum_i [n_i - 1]} \) respectively. Once every study was represented by a single mean and variance, they were then split by study type (RCT or OS), and overall pooled means and variances calculated using the same approach.

Comparisons were initially made between the study types with Kruskal-Wallis tests to produce an unweighted comparison of reported means or prevalences between the study types. As this does not take the size of studies into account, comparisons were then made using t-tests on the pooled means and variances. Variances were tested with F test, with Welch’s correction used when they differed significantly by study type. Sub-group analyses were performed by stratifying data by geography, and by quality of RCTs.

For the dichotomous outcomes (diabetes mellitus, gender), the overall rates were calculated by based on the total number of individuals with diabetes, or total number of
males in each type of study and calculating a percentage based on the total available data. This again removes the bias of larger studies dominating the data when only reported percentages were considered. Comparisons between study types were then performed using Chi-square tests. A p value of less than 0.05 was considered significant in all analyses. Prism v5.0 (Carlsbad, California USA) was used for statistical analysis.

Results

Literature search
The search strategies resulted in a total of 1165 studies being identified. After removal of irrelevant studies and duplicated data, a total of 143 studies were included (figure 1). The quality of included studies, assessed with reference to the CONSORT and STROBE guidelines for RCT and OS respectively, showed most studies to be of high quality (supplementary figure 1).

Characteristics of included studies
In total eighty-one observational studies were included, including data on a total of 40,014 individuals with NAFLD. Twenty-four studies were population-based studies that described individuals with NAFLD in a general population, and fifty-two described characteristics of patients with NAFLD in secondary or tertiary centres. Seventy-three studies described all stages of NAFLD, including 34,147 individuals (supplementary table 1). Eighteen studies including 2,780 individuals described patients with biopsy-proven NASH (supplementary table 2) and 28 studies with 1938 individuals described characteristics of patients with advanced fibrosis (AF) (supplementary table 3). Sixty-seven RCTs were included. These studies included a total of 4,846 individuals (supplementary table 4).

The methods used to diagnose NAFLD varied. Most population-based studies used imaging techniques, predominantly ultrasound, to define NAFLD whilst the majority of
secondary-care based studies used histology. NASH and AF were usually defined using the system proposed by Kleiner and Brunt (15), although some earlier studies used descriptive terms. In these cases, bridging fibrosis and/or cirrhosis were regarded as advanced fibrosis. Several studies did not report the variables of interest, or did so in such a manner that we were unable to extract data for use in the present study. Available data are summarised in supplementary table 5.

Comparison of Randomised Controlled Trials and Observational Studies in NAFLD
Analysis of unadjusted study-reported means from included studies showed significant variance in age, BMI, prevalence of diabetes and distribution of gender across all study cohorts (figure 2). To compare age and BMI between types of study weighted means were calculated to reflect the relative size of each study and compared with student’s t-test, and absolute prevalence of diabetes and male gender compared with Chi-squared test (table 1).

RCT and OS showed statistically significant differences with respect to age (mean age RCT 50.0 years (SEM 0.09) vs. OS 49.4 years (SEM 0.06) student’s t-test p<0.001) and BMI (32.1 kg/m\(^2\) (0.05) vs. 29.3 (0.03), p<0.001) (table 1, figure 3). Prevalence of diabetes and gender were compared by Chi-squared test (table 1). RCT and OS showed statistically significant differences with regard to the prevalence of diabetes (8%, 337 of 4186 participants, vs. 24%, 5561 of 23162 individuals, p<0.001) and gender of participants (55% male, 2617 of 4733 participants, vs. 50%, 12,405 of 24,648 individuals, p<0.001) (table 1, figure 4).

Subgroup analyses were performed to compare individuals with NASH or advanced fibrosis in OS to RCT participants. Age did not differ between RCT and individuals with biopsy-proven NASH (mean age RCT 50.0 years (0.09) vs. 49.9 (0.18), p=0.520) but did differ between RCT and individuals with advanced fibrosis (mean age 54.0 years (0.19), p<0.001) (table 1, figure 3). BMI differed significantly between RCT participants and
individuals with NASH (mean BMI RCT 32.1 (0.04) vs. 32.4 kg/m² (0.09) p=0.009), and also between RCT participants and individuals with advanced fibrosis (33.8 kg/m² (0.08), p<0.001) (table 1, figure 3).

The prevalence of DM and gender also differed significantly between RCT participants and individuals with NASH or advanced fibrosis. In RCTs, 8%, (337 participants) had DM. In observational studies, 39% of individuals with NASH (829 of 2145 participants) and 45% of individuals with advanced fibrosis (782 of 1726 participants) had DM (Chi squared test p<0.001 in each case)(table 1, figure 4). In RCTs, 55% of participants were male compared to 47% of individuals with NASH (1170 of 2476 individuals) and 38% of individuals with advanced fibrosis (712 of 1869 individuals) (Chi squared test p<0.001 in each case) (table 1, figure 4).

Analysis by geography
Ethnicity is associated with marked phenotypic differences in NAFLD (16, 17). In view of this observational studies were stratified based on geographic location (supplementary figure 2). BMI and gender distribution showed significant differences by geographic location (Kruskal-Wallis test p<0.0001 and <0.01 respectively). Accordingly, BMI and gender distribution in observational studies and RCTs were compared by geographical location. Significant differences remained between observational studies and RCTs (t-test of weighted means p<0.001). Again, these differences were most pronounced between RCTs and individuals with advanced fibrosis, with the exception of European data (supplementary figure 3). When considering results for gender, European data showed no differences between populations. In Asia and North America, RCTs contained significantly more men than were observed in epidemiological studies of advanced fibrosis (supplementary figure 4).
Exclusion criteria of RCT in NAFLD

The exclusion criteria of all identified RCT were reviewed. Eighteen trials including 2008 participants (41% of all participants) excluded patients with cirrhosis. Participants with DM were excluded in 17 studies (1144 participants, 24% of total) and exclusion criteria based on medications to treat DM were reported in a further 22 studies (1643 participants, 34% of total). Thus, the presence of DM represented an absolute or relative exclusion criterion in 39 trials including 2,787 participants (58% of all RCT participants). Individuals using drugs to treat dyslipidaemia were excluded in 7 trials (375 participants, 8%).

Discussion

These data, derived from a robust systematic review, show that characteristics of individuals with NAFLD in observational studies differ from those included in RCT. These differences are statistically significant but are often small and may not be clinically significant. However, marked differences exist between RCT cohorts and individuals with advanced fibrosis who are more likely to progress to liver-related morbidity. This is compounded by the frequent exclusion of patients with cirrhosis or diabetes from trials. These differences may limit the application of RCT trial data to high-risk patients with NAFLD.

There are important differences in susceptibility to insulin resistance and fatty liver between individuals of differing ethnicity (2, 16, 18). This was evident when studies were stratified by geography. Importantly, significant differences remained between OS and RCT populations.

An explanation for the differences seen between OS and RCT may lie in the current paradigm that steatohepatitis is required for the development of liver fibrosis. Since this process takes many years to develop, RCTs in patients with NASH have often relied on histological criteria to recruit patients and assess efficacy. For instance the accepted endpoint of an <=2 point improvement in the NAFLD activity score (NAS) without
worsening of fibrosis (19), specifically excludes patients with cirrhosis since there is no way to evaluate whether fibrosis has worsened in this group. Whilst RCT are necessarily different to real life clinical practice, use of this endpoint skews trial populations away from the groups at greatest risk of liver related morbidity and mortality, and towards younger patients with earlier disease. This questions the value of current surrogate endpoints in NAFLD trials and raises important issues regarding the definition of such outcome measures in early phase studies for patients with NASH. In other liver diseases, such as hepatitis B virus infection, when liver disease is treated in patients with cirrhosis there is evidence of a reduction in fibrosis progression (20) and a concurrent reduction in the risk of liver related events. It is important to discover whether this can also be achieved in patients with NASH and advanced fibrosis.

An additional cause for the differences in observational cohorts and trial participants is the stringent exclusion criteria applied in RCT. In particular, diabetes *per se* or use of medications for diabetes is a frequent cause of exclusion of patients from trials, while nearly a quarter of patients (24%) have DM in observational studies of NAFLD and nearly half of patients (45%) with advanced fibrosis have DM. In some trials, for example trials of metformin or thiazolidinediones, limitations on diabetic patients or diabetic medications may be justified but the applicability of these findings to patients with advanced fibrosis is then limited. The design of trials that exclude patients with both diabetes and cirrhosis is thus a major barrier to external validity since many of those patients at the greatest risk of liver related death are not represented in these studies. Recent notable trials in NAFLD have suggested that studies are becoming more inclusive. For example, the recent trial of Elafibranor included around reported that over 30% of participants had diabetes (21).

The American Association for the Study of Liver Disease (AASLD) published a consensus statement regarding trial design in NAFLD (22). This was added to in 2015 by a report of a meeting between representatives of the AASLD the US Food and Drug Administration (FDA) (19). These documents provide guidance that aims to achieve greater consistency in
trial design and to define outcomes relevant to patients. They do not provide recommendations on measures that might strengthen external validity *per se* but do recommend that trials in NAFLD should target specific groups, in particular those at risk of progression to cirrhosis, those with cirrhosis, and post-transplant patients. Advice from regulatory bodies regarding measures to improve of review external validity of trials is either lacking or non-binding.

These findings are similar to studies in other disease areas. In cardiovascular disease advanced patient age and the presence of co-morbidity were identified as important factors limiting external validity (9). The issue of co-morbidity is also important in patients with NAFLD. The association of NAFLD with the metabolic syndrome raises the risk of co-existing cardiovascular disease, prior extra-hepatic cancer, and other obesity related complications (23) that may also impact on recruitment into clinical trials. It is important that these co-morbidities are considered in future trial design. It is likely that inclusion of patients with significant co-morbidities would increase the risk of competing mortality such that in large scale, long duration licensing studies the number of patients required for sufficient statistical power would also be increased. Thus there is a short-term disincentive to include such patients in registration studies. In the long-term however it is preferable that physicians and patients understand the likely benefits and harms of treatment and this can only be achieved through the inclusion of such patients in licensing studies of novel therapeutics for patients with NASH.

This study has several limitations. It is limited by missing data, which also reflects data being presented in a manner that was not suitable for analysis in this study. Nevertheless, the comprehensive systematic review and consequent large numbers studies included goes some way to limiting the impact of these missing data. There is also a risk of bias in the selection of patients for inclusion in observational studies of NASH and advanced fibrosis, through selection for biopsy, which may only be performed in patients who appear at high risk to clinicians. This is also the case in observational studies based in
secondary care where a degree of selection bias may be present. The larger community based studies using non-invasive methods to diagnose NAFLD are less at risk of these biases.

In conclusion, we have identified differences in the characteristics of patients identified in observational studies of NASH, particularly those with advanced fibrosis, and those patients enrolled in RCTs of new therapeutic approaches. These findings, whilst partially explained by the therapies that have been trialed and by the choice of surrogate endpoint, highlight a risk that future studies will have limited external validity.

References

8. Rothwell PM. External Validity of Randomised Controlled Trials:“to Whom Do the Results of This Trial Apply?”. The Lancet. 2005;365(9453):82-93.
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### Calculated prevalence of diabetes, distribution of gender

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<td>48</td>
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**Table 1**: Characteristics of RCT and observational study cohorts. p-value refers to difference of observational studies to RCT cohort.
Figure 1. Flow chart of literature search outcomes.
Figure 2: Distribution of variables between types of study: A age, B BMI, where each point represents the mean value from an individual trial. C prevalence of diabetes and D male gender, each data point represents prevalence within an individual study. Lines at median and interquartile range. *p<0.05, **p<0.001, ***p<0.001 by Kruskal-Wallis test.
Figure 3: Age and BMI of participants in observational studies and RCTs. Data are shown as weighted mean, horizontal lines represent standard deviation. Dashed line at RCT mean. ***p<0.001 by student’s t-test
Figure 4: Prevalence of diabetes and distribution of gender in observational studies and RCTs. ***p<0.001 by Chi-squared test