

Title

An international Delphi consensus statement on metabolic dysfunction-associated fatty liver disease and risk of chronic kidney disease

Short title: MAFLD and CKD risk

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Provision of study material or patients: All authors.

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1 **Abstract**

2 With the rising global prevalence of fatty liver disease related to metabolic
3 dysfunction, the association of this common liver condition with chronic kidney
4 disease (CKD) has become increasingly evident. In 2020, the more inclusive term
5 metabolic dysfunction-associated fatty liver disease (MAFLD) was proposed to
6 replace the term non-alcoholic fatty liver disease (NAFLD). The observed association
7 between MAFLD and CKD and our understanding that CKD can be a consequence of
8 underlying metabolic dysfunction support the notion that individuals with MAFLD
9 are at higher risk of having and developing CKD compared with those without
10 MAFLD. However, to date, there is no appropriate guidance on CKD in individuals
11 with MAFLD. Furthermore, there has been little attention paid to the link between
12 MAFLD and CKD in the Nephrology community. Using a Delphi-based approach, a
13 multidisciplinary panel of 50 international experts from 26 countries reached a
14 consensus on some of the open research questions regarding the link between
15 MAFLD and CKD. This Delphi-based consensus statement provided guidance on the
16 epidemiology, mechanisms, management and treatment of MAFLD and CKD, as well
17 as the relationship between the severity of MAFLD and risk of CKD, which establish
18 a framework for the early prevention and management of these two common and
19 interconnected diseases.

20

21

- 1 **Keywords:** metabolic dysfunction-associated fatty liver disease, non-alcoholic fatty
- 2 liver disease, chronic kidney disease, consensus

1 **Introduction**

2 Non-alcoholic fatty liver disease (NAFLD) is the most common liver disease
3 worldwide with a global prevalence of about 25-30%(1,2). NAFLD includes a
4 histological spectrum of liver conditions ranging from simple steatosis (non-alcoholic
5 fatty liver, NAFL) to non-alcoholic steatohepatitis (NASH), advanced fibrosis and
6 cirrhosis(3). NAFLD is always a diagnosis of exclusion in clinical practice; to
7 entertain the diagnosis of NAFLD, clinicians need to exclude “excessive” alcohol
8 consumption and all competing causes of hepatic steatosis. This is despite the fact that
9 the coexistence of NAFLD with other chronic liver diseases (including but not limited
10 to alcohol use disorder) is not rare in clinical practice(4). On the other hand, in the
11 realm of drug development and regulatory approval processes, the definition of a
12 patient population in which the mechanism of the drug can be linked to one
13 underlying dominant pathophysiological process is critical. For these reasons and
14 given the high heterogeneity and stigma around the NAFLD name, in 2020, several
15 experts proposed the new term metabolic dysfunction-associated fatty liver disease
16 (MAFLD)(5,6). A diagnosis of MAFLD is based on evidence of hepatic steatosis (as
17 assessed by liver biopsy, imaging techniques or blood biomarkers/scores) in persons
18 who are overweight or obese or have type 2 diabetes (T2D), or metabolic
19 dysregulation, regardless of the coexistence of excessive alcohol consumption and
20 other chronic liver diseases. The newly proposed definition of MAFLD better
21 emphasises the pathogenic role of metabolic dysfunction in the development of this

1 common liver disease and uses inclusive criteria for diagnosis(7-10). In this article,
2 we explore the definition of MAFLD characterized by the presence of metabolic
3 dysregulation but excluding severe alcohol use or viral-associated liver disease (i.e.,
4 dual aetiology liver disease).

5

6 Growing evidence indicates that NAFLD is associated with an increased risk of
7 having or developing chronic kidney disease (CKD)(11-14), which is an established
8 risk factor for end-stage renal disease, cardiovascular disease and all-cause
9 mortality(15-18). The magnitude of these risks appears to parallel the severity of
10 NAFLD, especially the amount of liver fibrosis(11,19). In contrast, current data on the
11 strength of the association between MAFLD and subsequent risk of CKD is only now
12 being acquired, given its proposed adoption as a clinically-useful entity(20-23).

13 Several epidemiological studies have documented that MAFLD may be even more
14 closely associated with CKD than NAFLD (**Supplementary Table 1**)(24). Sun et al.
15 first reported that in 12,571 individuals with liver ultrasonography data from the Third
16 National Health and Nutrition Examination Survey (NHANES) 1988-1994,
17 individuals with MAFLD had lower values of estimated glomerular filtration rate and
18 a greater prevalence of CKD than those with NAFLD (29.6% vs. 26.6%, $p<0.05$)(25).

19 Over a 10-year follow-up among 28,890 Japanese individuals, MAFLD also better
20 identified subjects developing CKD, than NAFLD. Furthermore, the addition of
21 MAFLD to traditional CKD risk factors improved discriminatory capacity to diagnose

1 CKD better than NAFLD(26). Similar findings were observed in other large cohorts
2 of Asian individuals (23,27). In contrast, in two prospective cohort studies from USA
3 and China, the MAFLD and NAFLD definitions were both comparable risk factors
4 for CKD(21,28). That said, despite some inconsistencies between research study
5 findings, the MAFLD definition is a landmark in Hepatology bringing about a new
6 way of thinking about fatty liver disease and the relevance of metabolic dysregulation
7 and increased body fat accumulation that has consequences beyond the liver.
8 Importantly, MAFLD brings liver disease into closer alignment with our current
9 understanding of obesity and metabolic syndrome, both of which contribute to
10 development of kidney injury(29). Unfortunately, few outside the field of Hepatology
11 are familiar with the newly-proposed MAFLD terminology and its definition; and
12 there is limited awareness of the link between MAFLD and CKD, amongst the
13 Nephrology community.

14

15 The objective of this study was therefore to build consensus among international
16 experts in the field on the link between MAFLD and CKD using a Delphi-based
17 approach. The consensus statements set out current ideas on the link between
18 MAFLD and CKD in specific areas ranging from epidemiology to mechanisms,
19 management and treatment.

20

21 **Methods**

1 Study design

2 The Delphi method was originally developed at the RAND Corporation (Santa
3 Monica, CA, USA) in the 1950s to forecast the effect of technology on warfare.
4 Today, groups of experts use online tools to anonymously answer questionnaires and
5 receive feedback that represents the “group response” and revise their answers to see
6 whether they can approach expert consensus. Thus, the Delphi method is a structured
7 multistage process which aims to transform expert opinion into group consensus on a
8 given subject(30). The Delphi method can be successfully applied to areas of
9 controversy or when data are inadequate, and involves a series of questionnaires
10 interspersed with controlled feedback(31). In the present study, we used a modified
11 Delphi process via an online survey with the goal of reaching a consensus on the link
12 between MAFLD and the risk of CKD (3). A two-round Delphi survey (i.e. the R1-
13 survey on 15 April 2022, and R2-survey on 16 June 2022) employed a structured
14 interaction in which a multidisciplinary panel of 50 international experts from 26
15 countries evaluated and re-evaluated consensus statements in multiple rounds until
16 agreements were reached (**Figure 1**). The web-based Delphi survey was delivered to
17 each member of the expert panel via email with a secure link using Google forms
18 (link for **R1 survey**: <https://forms.gle/oPNEQqfv53UpsTC59>; for **R2 survey**:
19 <https://forms.gle/tntWm2Nk2s4EeEmg9>). The data collection periods for each survey
20 ranged between one and four weeks. The R1-survey contained four domains and 22
21 draft statements with four-point Likert-type categories for respondents to indicate

1 their level of agreement with the statements (that is, ‘Agree’/ ‘Somewhat agree’/
2 ‘Somewhat disagree’/ ‘Disagree’) (as specified in **Supplementary Table 2**). In the
3 first round, respondents who agreed or somewhat agreed with a statement could
4 provide comments or suggest edits while those who disagreed or somewhat disagreed
5 needed to explain why. Further discussion was undertaken by email to report the
6 results of R1-survey and the comments in R1-survey. The R2-survey reflected
7 suggestions developed from the R1-survey, including revised, merged or deleted
8 statements and, finally, contained 21 statements. Only respondents who completed the
9 R1-survey were eligible to take the R2-survey (**Supplementary Table 3**), and all
10 respondents in the R1-survey participated in the R2-survey. Participants had the
11 option of keeping their first-round ratings or having them re-scored. After the R2-
12 survey, we included summaries of the edits made to each statement from respondents
13 and emailed all respondents to consider their level of agreement or disagreement with
14 the statements. For the Delphi process, the consensus statements were developed by
15 the expert panel and we assigned a grade to each statement and recommendation to
16 indicate the level of agreement utilising a grading system used in other published
17 Delphi studies, in which ‘U’ denotes unanimous (100%) agreement, ‘A’ 90-99%
18 agreement, ‘B’ 78-89% agreement, and ‘C’ 67-77% agreement(3,32). A preliminary
19 consensus draft on these recommendations from the expert panel was sought over a 1-
20 week period via a shared Google document. Any disagreements were resolved
21 through discussion until consensus was reached.

1

2 **Recruitment of expert panel members**

3 Members of the international expert panel (n=50) were selected from the
4 representative Continents. To be included, they were active researchers with expertise
5 in the management of fatty liver and/or kidney diseases.

6 The following criteria were used to select members of the expert panel participating in
7 the Delphi survey:

8 (1) to be corresponding authors of published articles on the association between
9 MAFLD or NAFLD and the risk of CKD.

10 (2) to be representative members from scientific Societies of Nephrology,
11 Hepatology, Endocrinology/Diabetology, and Obesity.

12 (3) to be core members of the NAFLD Consensus Consortium and/or the Improving
13 Global Outcomes (KDIGO) organization.

14 Members of the expert panel were expected to meet at least one of the three
15 aforementioned criteria. To achieve global representation, we selected members from
16 six Continents, i.e. Asia, Europe, North America, South America, Africa and Oceania
17 **(Table 1)**.

18

19 **Findings**

20 Here, we report the final consensus statements along with a summary of the broader
21 relevant literature. Across the two-based Delphi surveys, there was an increase in

1 consensus for all proposed statements. The mean percentage of “agreement”
2 responses increased from 63.9% to 76.1% and “agreement or somewhat agreement”
3 responses increased from 94.3% in the R1-survey to 97.3% in the R2-survey (**Figure**
4 **2**). In the end, there was unanimous “agreement or some agreement” on 12 consensus
5 statements and >85% agreement on 7/12 statements (**Table 2**).

6

7 **Epidemiology of MAFLD and CKD**

8 **Statements 1.1-1.6 (Grade U in 1.1 and 1.5; Grade A in 1.2 to 1.4, 1.6)**

9 Studies using the NAFLD definition have estimated a global prevalence of this
10 condition of about 30% in the general adult population. NAFLD is considered part of
11 a multisystem disease associated with an increased risk of developing not only liver-
12 related complications but also cardiovascular disease(33) and CKD(34). Given this
13 current understanding of the pathogenesis of NAFLD, the term MAFLD focuses
14 attention on the pathogenic role of metabolic dysfunction in the development and
15 progression of this liver disease and its accompanying systemic extra-hepatic
16 complications(35-37).

17

18 Recently, it has been reported that during a median follow-up of 23 years, individuals
19 with MAFLD had a 24% higher risk of cardiovascular mortality (HR 1.24; 95% CI
20 1.01-1.51; p=0.041) and a 17% higher risk of all-cause mortality (HR 1.17; 95% CI
21 1.04-1.32; p<0.01) compared to those without MAFLD(38). It is, therefore, not

1 surprising that MAFLD is associated with a higher prevalence of CKD compared to
2 that observed in the non-MAFLD population. For example, from the cross-sectional
3 NHANES 1999-2002, 2003-2006, 2007-2010 and 2011-2016 cohort databases,
4 individuals with MAFLD had a greater odds of any CKD stage and albuminuria
5 compared with those without MAFLD(28). Using the NHANES 1988-1994 database,
6 the authors reported that compared to the NAFLD or non-metabolic risk NAFLD
7 groups, subjects with MAFLD had lower eGFR values and a higher prevalence of
8 both CKD and abnormal albuminuria(25). Collectively, these findings suggest that
9 MAFLD is associated with a higher risk of CKD compared to subjects with fatty liver
10 but without coexisting metabolic disorders.

11

12 In most published studies, using the term NAFLD, liver disease was associated with a
13 nearly 2-fold increased prevalence of CKD and this association persisted both in
14 patients with T2D and in those without diabetes, even after adjustment for common
15 risk factors for CKD(12,39,40). In a large retrospective cohort study of German
16 individuals with NAFLD, Kaps et al. reported that NAFLD was associated with
17 higher risk of developing CKD over 10 years of follow-up(41). This association
18 remained significant across different age and patient subgroups, such as those with
19 T2D, obesity, hypertension or ischaemic heart disease. In contrast, NAFLD was not
20 independently associated with the future risk for end-stage renal disease (ESRD)
21 requiring haemodialysis. In a study where the MAFLD population was stratified by

1 presence or absence of T2D, individuals with MAFLD and T2D had a higher
2 prevalence of CKD stage ≥ 1 than their counterparts without T2D [odds ratio (OR)
3 1.18 (95%CI: 1.05-1.32), $p < 0.05$] or those with T2D alone [OR 2.09 (95%CI: 1.78-
4 2.46), $p < 0.05$](25). Using the NHANES 2017-2018 database, the authors found that
5 the metabolic comorbidities of MAFLD such as T2D, hypertension and
6 hyperuricemia were all independently associated with CKD(22). Therefore, these
7 findings suggest that MAFLD is associated with CKD in both patients with or without
8 T2D, even after adjustment for common risk factors for CKD.

9

10 Although the association between MAFLD and CKD from cross-sectional studies
11 appears to be strong and consistent, whether MAFLD is also an independent risk
12 factor for CKD remains uncertain. In a cohort study of middle-aged and elderly
13 Chinese subjects without CKD at baseline, the authors found that the incidence rates
14 of CKD in those without fatty liver and those with MAFLD were 8.2% (95%CI 7.3-
15 9.2) and 12.9% (95%CI 11.7-14.1), over a mean follow-up of 4.6-years(21). These
16 authors also found that MAFLD was associated with a higher risk of incident CKD
17 (HR 1.64, 95%CI 1.39-1.94). This finding is consistent with results from an updated
18 meta-analysis of 13 observational studies showing that fatty liver disease was
19 significantly associated with a nearly 1.5-fold increased long-term risk of incident
20 CKD stage ≥ 3 (11). In 268,946 individuals from the NHANES 2009-2015 database,
21 the investigators found that MAFLD identified a higher proportion of individuals at

1 risk of developing CKD than NAFLD over a median follow-up of 5.1 years(27).
2 Similar results were reported in another cohort study with a 10-year follow-up, where
3 the risk for incident CKD was 1.12 [95%CI (1.02-1.26)] in MAFLD individuals, even
4 after adjustment of traditional renal risk factors(26). Moreover, a Mendelian
5 randomization study supported the existence of a causal effect of fatty liver disease on
6 lower eGFR levels and CKD(42). Thus, the aforementioned studies suggest that
7 individuals with MAFLD are at higher risk of new-onset CKD even after adjustment
8 for common cardiometabolic risk factors compared to subjects with fatty liver who do
9 not have metabolic dysregulation.

10

11 Moderate to advanced stages of CKD may also increase the risk of overall mortality
12 among patients with NAFLD (CKD stages 2-3a: HR=2.31, 95% CI: 1.70-3.15; CKD
13 stages 3b-5: HR=4.83, 95% CI: 2.40-9.71)(43). Interestingly, in that study, mortality
14 risk was significantly increased in NAFLD patients with CKD due to metabolic
15 comorbidities, and not influenced by CKD per se. According to the newly proposed
16 MAFLD definition, most of these NAFLD individuals had MAFLD. In contrast, a
17 small prospective study showed that NAFLD patients with CKD had a higher risk of
18 overall mortality than NAFLD patients without coexisting CKD. However, after
19 adjustment for metabolic comorbidities, this risk was no longer significant(44).

20 Although further studies are needed, the evidence from the current studies indicate
21 that recognition of CKD may increase the risk of overall mortality in patients with

1 MAFLD, and the new term MAFLD improves our ability to identify individuals at
2 higher risk of developing CKD.

3

4 Studies also support a role for NAFLD as a risk factor for CKD in childhood(45,46).
5 For example, in a cohort of 596 children who were overweight or obese, an
6 association between NAFLD and early kidney dysfunction (defined as
7 microalbuminuria or eGFR<90 ml/min/1.73m²) was suggested(45). Other studies
8 indicate that the link between NAFLD and CKD could be modulated by some genetic
9 factors. For example, the risk patatin-like phospholipase domain-containing protein 3
10 (*PNPLA3*) allele may increase the risk of developing both NAFLD and CKD.
11 However, in other studies, carriers of the hydroxysteroid 17-beta dehydrogenase 13
12 (*HSD17B13*) at-risk A gene or the trans-membrane 6 superfamily 2 (*TM6SF2*) 167K
13 allele had higher eGFR levels in patients with NAFLD(47-49). Overall, given that
14 current evidence on the relationship between MAFLD and CKD in childhood is not
15 robust, a specific consensus statement cannot be generated. New data to inform this
16 are eagerly awaited. In our two-round Delphi survey process, about 25% of experts
17 disagreed with the statement in the R1-survey, so this statement was deleted in the
18 R2-survey.

19

20 **Severity of MAFLD and CKD**

21 **Statements 2.1-2.6 (Grade U in 2.3 to 2.5; Grade A in 2.1 to 2.2, 2.6)**

1 As per its definition, the MAFLD criteria are more likely to capture those who have
2 coexisting metabolic comorbidities compared to NAFLD criteria, and to identify
3 individuals with advanced liver fibrosis(50,51). Given the close association between
4 fibrotic fatty liver disease and CKD, it is reasonable to infer that the severity of
5 MAFLD may be closely associated with CKD. Though there are only a few studies
6 exploring the relationship between the severity of MAFLD and risk of CKD, the
7 available evidence suggests that MAFLD individuals with steatohepatitis or advanced
8 fibrosis had a higher prevalence and incidence of CKD than those without advanced
9 fibrosis or those with simple steatosis. An observational study demonstrated that
10 advanced liver fibrosis but not steatosis was associated with abnormal albuminuria in
11 Chinese patients with NAFLD and T2D (all of whom fit the MAFLD definition)(52).
12 In a meta-analysis of 13 observational cohort studies with a median follow-up of 9.7
13 years, Mantovani et al. also showed that imaging-defined NAFLD was associated
14 with a moderately increased risk of incident CKD stage ≥ 3 (random-effects HR 1.43,
15 95% CI 1.33-1.54)(11). Similarly, from 5 small studies with liver histology, the
16 presence of advanced fibrosis (F3/4 stage) was associated with a higher prevalence
17 (random-effects OR 5.20; 95% CI 3.14-8.16) and incidence (random-effects HR 3.29;
18 95%CI 2.3-4.71) of CKD than either non-advanced fibrosis (F0-2) or simple steatosis,
19 respectively(53).
20
21 While evidence for the existence of a significant association between severity of

1 NAFLD and risk of prevalent and incident CKD is robust, the association between
2 severity of MAFLD and the risk of having or developing CKD remains
3 uncertain(54,55). In a study from the NHANES-III database, it was reported that
4 MAFLD with increased liver fibrosis scores was strongly associated with a greater
5 risk of having CKD stage ≥ 1 or ≥ 3 and abnormal albuminuria(25). Another small
6 prospective study of T2D patients with and without NAFLD followed for 75 months
7 showed that the presence of NAFLD with high-risk fibrosis (defined as NAFLD
8 fibrosis score >0.181) conferred a greater eGFR reduction (58.7% vs. 37%; $p=0.04$)
9 and higher risk of CKD progression (defined as decrease in $>50\%$ eGFR)
10 ($p<0.001$)(56). In a meta-analysis, participants with T2D and steatohepatitis (where
11 by definition all subjects had MAFLD) there was a 3.8-fold risk of prevalent CKD
12 [95%CI (1.47-9.81), $I^2=0\%$, $n=3,119$ participants] and a 2.5-fold increased risk of
13 incident CKD [95%CI (1.05-6.17), $I^2=0\%$, $n=396$ participants] compared with their
14 counterparts who had simple steatosis(53). Furthermore, in subjects who had T2D and
15 NAFLD with advanced fibrosis (state F3/F4) (subjects all fulfilling the MAFLD
16 criteria), there was a 5.1-fold increased risk of prevalent CKD [95%CI (1.46-17.21),
17 $I^2=0\%$, $n=3,120$ participants] and a 4.2-fold increased risk of incident CKD [95%CI
18 (2.10-8.38), $I^2=0\%$, $n=397$ participants], compared to those subjects with non-
19 advanced fibrosis (stage F0-2)(53). The above-mentioned studies indicate that
20 MAFLD patients with steatohepatitis have a higher prevalence and incidence of CKD
21 compared to those with simple steatosis alone. Further, MAFLD with advanced

1 fibrosis has a higher prevalence and incidence of CKD than MAFLD without
2 advanced fibrosis.

3

4 Transient elastography (TE) is extensively used in clinical practice as a non-invasive
5 technique for measuring liver stiffness, a correlate of liver fibrosis. Consistently, TE
6 identifies a subgroup of NAFLD patients who are at higher risk of developing liver-
7 related clinical events(57-59). Our prior study also showed that the association
8 between liver stiffness (assessed by TE) and risk of abnormal albuminuria was
9 consistent with histological data obtained by liver biopsy(34). A meta-analysis of 7
10 cross-sectional studies also showed that increased liver stiffness was associated with
11 an increased odds for both CKD (OR 2.49, 95%CI 1.89-3.29, $p < 0.001$) and abnormal
12 albuminuria (OR 1.98, 95%CI 1.29-3.05, $p = 0.002$) in patients with NAFLD(60).
13 Another small study from 42 outpatients with established T2D showed that significant
14 liver fibrosis [i.e., defined as liver stiffness $\geq 7.0/6.2$ kPa (medium/extra-large probe)]
15 was associated with an increased likelihood of CKD (OR 4.54, 95%CI 1.24-16.6),
16 independently of common cardiometabolic risk factors(61). Thus, liver stiffness,
17 which is a surrogate of liver fibrosis and inflammation, is independently associated
18 with an increased risk of CKD or albuminuria. While there are no specific studies on
19 patients with MAFLD, data are awaited to better clarify the association between the
20 severity of MAFLD and CKD progression.

21

1 It is important to emphasise that none of the aforementioned studies used renal biopsy
2 to examine the pathology of CKD, so whether MAFLD is associated with a specific
3 type of kidney injury is currently unknown. Moreover, it is also important to highlight
4 that while we identify CKD by using a functional classification of CKD stages based
5 on estimated glomerular filtration rate and proteinuria, we do not have a
6 corresponding scale for evaluating the degree of hepatic function impairment.
7 Recently, Aubert et al. reported that patients with diabetic kidney disease (confirmed
8 by renal biopsy) and advanced liver fibrosis (F3-F4 stages) tended to have a greater
9 annual eGFR decline (-3.27 ± 3.07 vs. -6.29 ± 4.72 ml/min/1.73 m²) compared to those
10 with diabetic kidney disease without advanced liver fibrosis during a 75-month follow
11 up period(56).

12

13 **Mechanisms linking MAFLD with CKD**

14 **Statements 3.1-3.4 (Grade U in 3.1 and 3.4, Grade A in 3.3, Grade B in 3.2)**

15 Current evidence suggests that MAFLD may be an independent risk factor for
16 CKD(29). A large cross-sectional study also showed that the metabolic syndrome and
17 its individual components are independently associated with CKD(62). Therefore, as
18 highlighted in the consensus statements, metabolic dysfunction in MAFLD might be
19 an important mechanistic link between MAFLD and CKD as discussed below.

20

21 Firstly, convincing evidence showed that obesity plays an important role in the

1 development and progression of both MAFLD and CKD(63-66). For example, in a
2 retrospective study evaluating native kidney biopsies, obesity-related kidney disease
3 increased in parallel with the worldwide epidemic of obesity. In that study, 56% of
4 patients had overt proteinuria alone and 44% had overt proteinuria and CKD(67). At a
5 mechanistic level, the renal physiologic responses to obesity include increases in
6 glomerular filtration rate, renal plasma flow, filtration fraction and tubular
7 reabsorption of sodium, which exerts a high fluid shear stress on renal podocytes,
8 thereby promoting maladaptive renal hypertrophy, podocyte detachment and global
9 glomerulosclerosis.

10

11 Secondly, T2D has a substantial adverse impact on health and increases risk of both
12 kidney and liver diseases. Strong evidence shows that chronic hyperglycaemia is a
13 driving force for the development and progression of MAFLD and CKD, possibly
14 through intraglomerular hypertension induced by glomerular hyperfiltration, increased
15 formation of advanced glycation end-products, microinflammation and subsequent
16 extracellular matrix expansion(68,69). Meanwhile, adipokines may also play
17 important roles in kidney disease progression by promoting maladaptive responses of
18 renal cells to the mechanical forces of hyperfiltration, thereby leading to podocyte
19 depletion, proteinuria, focal segmental glomerulosclerosis and interstitial fibrosis(70).

20

21 Thirdly, abnormal lipid metabolism promotes increased triglyceride and cholesterol

1 ester accumulation in the liver and kidneys(71). Increased lipids accumulate in
2 mesangial cells, which may, in turn, transform to a type of foam cell, which activates
3 insulin growth factor-1 and contributes to the loss of glomerular integrity. More
4 importantly, renal fat accumulation as a result of increased fatty acid synthesis (which
5 is mainly mediated by sterol regulatory element-binding protein 1c [SREBP-1c] and
6 its target enzymes) may induce low-grade inflammation, oxidative stress and
7 increased expression of multiple profibrotic growth factors(72-74). Finally, increased
8 fat accumulation is associated with SREBP expression and activity, thus resulting in
9 the development of renal disease(75). These results provide mechanistic data
10 suggesting that metabolic dysfunction links MAFLD and CKD.

11

12 Findings from genome-wide association studies in large cohorts of well-phenotyped
13 individuals show that the rs738409 C>G SNP encoding the I148M genetic variant of
14 *PNPLA3* accounts for the largest fraction of genetic predisposition to fatty liver
15 disease(76,77). Carriage of this genetic variant has also been associated with an
16 increased risk of liver-related mortality and extrahepatic complications, especially
17 kidney injury(46,78,79). *PNPLA3* is highly expressed both in the liver (by hepatic
18 stellate cells and hepatocytes) and in the kidneys. Studies have shown that individuals
19 with the *PNPLA3* rs738409 GG genotype are more likely to have lower levels of
20 eGFR, and higher prevalence of both abnormal albuminuria and CKD, compared to
21 those carrying the *PNPLA3* rs738409 GC and CC genotypes(46,80-83). Another

1 study showed that this *PNPLA3* genetic variant or other NAFLD-related genetic
2 polymorphisms did not directly contribute to eGFR decline, but that metabolic risk
3 factors were more important(84). However, such study did not retrieve data on
4 albuminuria, so that the CKD diagnosis was based only on eGFR values. Evidence
5 about the association between MAFLD, *PNPLA3* rs738409 variant and CKD is still
6 limited since the data have only accrued for less than 2 years. Further studies are
7 therefore needed to better understand the role of the *PNPLA3* rs738409 variant (or
8 other MAFLD-related genetic polymorphisms) in the development and progression of
9 CKD, and to elucidate the function of the mutant PNPLA3 protein in the kidney.

10

11 Recent studies have unveiled a role for the liver-gut-kidney axis in both health and
12 disease states(85-88). Gut microbiota is thought to be one of the major contributing
13 factors to the pathophysiology of CKD associated with fatty liver. Gut microbiome
14 homeostasis is important for health and its imbalance can lead to bacterial
15 translocation, as well as the release of microbial products like lipopolysaccharide,
16 incosyl sulphate, p-cresyl sulphate and trimethylamine N-oxide (TMAO) into the
17 circulation, where they may contribute to low-grade inflammation. These factors may
18 also increase the risk of both MAFLD and CKD(85,89,90). On the other hand,
19 MAFLD may alter gut microbiota composition and contribute to the development and
20 progression of CKD associated with MAFLD. For instance, gut microbiota
21 metabolizes dietary components such as choline and carnitine to produce TMAO,

1 which may induce kidney and liver injuries. A cohort study of 521 subjects with 5-
2 year follow-up showed that compared to non-CKD individuals, patients with CKD
3 had higher plasma levels of TMAO and that plasma TMAO levels were associated
4 with a near 1.9-fold increase in mortality risk after adjustment for traditional renal risk
5 factors(91). Meanwhile, compared to non-steatotic controls, patients with fatty liver
6 disease had higher plasma TMAO levels, which were positively correlated with serum
7 bile acid concentrations and the mRNA expression of hepatic CYP7A1(92).
8 Experimentally, administration of TMAO to mice induced progressive renal tubulo-
9 interstitial injury and fibrosis, while in mice fed a high-fat diet TMAO administration
10 exacerbated hepatic steatosis by inhibiting hepatic farnesoid X receptor signalling and
11 up-regulating hepatic de novo lipogenesis(92). Although current evidence is
12 inconclusive and further studies are needed, the aforementioned studies suggest that
13 alterations in gut microbiota may be linked to both MAFLD and CKD.

14

15 Studies have identified various immune mechanisms which play a key role in NAFLD
16 pathogenesis, especially triggering low-grade inflammation, and which are rooted in
17 intrahepatic and extrahepatic systems(93). Extrahepatic factors include multiple organ
18 crosstalk between inflammatory signals derived from the gut, adipose tissue, skeletal
19 muscles and bone marrow, and some intrahepatic factors such as the cholangiocytes
20 that are recognised as a potential driver of low-grade inflammation in NAFLD.

21 However, to date, we are uncertain on how specific immune cell subsets interact and

1 how they interact with stromal liver cells during NAFLD development and
2 progression. Even less is known about how immune-mediated molecular mechanisms
3 are implicated in the pathologic interaction between the liver and kidney in MAFLD.
4 It is known that low-grade inflammation plays a key role in the development and
5 progression of CKD. A prospective study of 2,838 Chinese patients with T2D (with or
6 without chronic hepatitis B virus infection who were followed for a median of 3.5
7 years) showed that the presence of liver inflammation was associated with increased
8 risk of end-stage renal disease, and this was independent of other potential
9 confounding factors(94). Finally, emerging evidence supports a potential pathogenic
10 role of the hepato-renal reflex in CKD development which may be triggered by
11 subclinical portal hypertension(95), although further research in this area is needed.

12

13 **Managing and treating MAFLD and CKD**

14 **Statements 4.1-4.5 (Grade U for 4.1-4.5)**

15 Currently, there are no specific treatment guidelines for patients with CKD and
16 MAFLD. However, MAFLD and CKD share multiple cardiometabolic risk factors
17 and therapeutic strategies for MAFLD and CKD should be similar and primarily
18 focussed on improving all coexisting renal and metabolic risk factors.

19

20 Lifestyle intervention (including a hypocaloric diet and regular physical activity) is
21 associated with improvements in both MAFLD and CKD, though the extent of benefit

1 might be different for each disease(96-100). For example, a large prospective study in
2 real-world clinical practice showed that modest (7-10%) and good ($\geq 10\%$) weight
3 reduction induces significant improvements in liver histology in patients with
4 steatohepatitis(101). A recently study that included 261 patients with biopsy-proven
5 NASH also showed that a one-stage reduction in liver fibrosis and resolution of
6 steatohepatitis was associated with an improvement in kidney function
7 parameters(102). Recently, an aerobic exercise intervention study of patients with
8 biopsy-proven MAFLD showed that a 12-week intervention reduced liver fibrosis and
9 hepatocyte ballooning by one stage in 58% ($p=0.034$) and 67% ($p=0.02$) of these
10 patients, respectively(103). Another study including obese patients with T2D and
11 CKD reported that a combined diet and exercise intervention reduced proteinuria
12 compared to a diet only(104). A further study of overweight and obese patients with
13 T2D showed that weight loss improved renal function parameters(105). Therefore, a
14 body of evidence supports the notion that lifestyle interventions play an important role
15 in the prevention and management of both MAFLD and CKD.

16
17 Current evidence indicates that MAFLD and CKD are two risk factors for adverse
18 cardiovascular outcomes and all-cause mortality(106-109) . Increasing evidence
19 recommends that patients with MAFLD should be treated early and aggressively for
20 obesity and other coexisting cardiometabolic risk factors(110,111). Most available
21 drugs that target cardiometabolic risk factors exert their actions either directly or

1 indirectly on glucose and lipid metabolism. Newer classes of glucose-lowering agents,
2 such as GLP-1 receptor agonists (mostly subcutaneous liraglutide and semaglutide)
3 and SGLT2 inhibitors, not only exert some beneficial effects on the liver (especially
4 hepatic steatosis and necro-inflammation), but also have clinically meaningful effects
5 on cardiovascular and kidney outcomes(112-117). Statin use also markedly reduces
6 the risk of fatal and nonfatal CVD events associated with MAFLD(118,119) and may
7 contribute to reduce the risk of MAFLD development(120). Similarly, in patients with
8 CKD not requiring dialysis, statin use decreases the risk of all-cause mortality and
9 major adverse cardiovascular events(121). Therefore, an early and aggressive
10 treatment of coexisting cardiometabolic risk factors will help prevent or slow the
11 development and progression of both MAFLD and CKD.

12
13 Hypertension is an established cardiovascular risk factor and a major component of
14 the metabolic syndrome. The coexistence of hypertension and MAFLD has been
15 reported to be common and to increase metabolic and cardiovascular risks(122). The
16 strong association and similar pathogenic profile of MAFLD and hypertension
17 suggests that treatment with antihypertensive agents might be beneficial in
18 hypertensive subjects with MAFLD(123). Although no large randomized controlled
19 trials have specifically investigated the long-term effect of antihypertensive agents on
20 MAFLD, inhibitors of the renin-angiotensin-aldosterone system (RAAS) may be of
21 benefit(124). For example, in a small intervention study of 54 subjects with

1 hypertension and fatty liver disease assigned to receive either valsartan or telmisartan,
2 both treatments led to amelioration of insulin resistance and hepatic fibrosis
3 improvement(123). A meta-analysis of seven interventional studies (1066
4 participants) reported that treatment with RAAS inhibitors may exert beneficial
5 effects on hepatic fibrosis or cirrhosis patients based on effects on liver histological
6 endpoints(125). Another intervention study reported that telmisartan decreased liver
7 fat content and serum free fatty acid levels in hypertensive patients with
8 MAFLD(126). Several studies showed that RAAS inhibitors were associated with
9 beneficial effects on proteinuria and the rate of eGFR decline in patients with
10 CKD(127,128). Similarly, in a cross-sectional study of CKD individuals with or
11 without NAFLD, treatment with RAAS inhibitors was associated with lower liver
12 stiffness in those with NAFLD, compared to those without(129,130). Finally, and
13 more interestingly, treatment with ACE-inhibitors may have beneficial effects on liver
14 fibrosis(131). In a cohort study of 12,327 Asian individuals with NAFLD followed for
15 at least 5 years, the authors found that treatment with ACE-inhibitors (but not with
16 angiotensin II receptor antagonists) in those with hypertension, was associated with a
17 lower risk of developing liver-related events, liver cancers, and cirrhotic
18 complications, especially amongst those with CKD(131). Therefore, treatment with
19 antihypertensive agents, especially RAAS inhibitors (if required), is clinically
20 important in hypertensive patients with MAFLD for decreasing the risk of CKD.
21

1 Taken together, the current evidence from published studies suggest that increased
2 clinical vigilance for the presence of MAFLD should be considered in patients with
3 CKD. Patients with MAFLD and CKD should ideally be managed in teams, though
4 the ideal model of care has not been identified.

5

6 **Study strengths and limitations**

7 Although the Delphi method is a consensus-building initiative, it also comes with
8 strengths and limitations. As an important strength, we employed 50 experts from six
9 Continents and more than 26 countries, comprising hepatologists, nephrologists,
10 endocrinologists, diabetologists and other specialists with extensive research and
11 clinical expertise. Delphi studies often involve a combination of in-person, in-depth
12 deliberation and survey rounds for voting. However, in light of the geographical
13 spread of the panel members and the COVID-19 travel restrictions, we employed
14 alternative modes for group discourse in which members were able to provide written
15 comments on the draft by email and two survey rounds. We incorporated risk factors
16 from the preliminary findings of our review and translated them into Delphi survey
17 statements. We received and incorporated a large volume of open-ended comments
18 across all four data collection components. Such feedback provided a mechanism for
19 reconciling the different views. We however acknowledge that a combination of in-
20 person and written feedbacks might have resulted in more comprehensive
21 contributions overall. The increasing levels of agreement with the consensus

1 statements across the two survey rounds, together with the high level of participation
2 [83.3% (50/60) in the R1-survey and 100% (50/50) in the R2-survey], further
3 strengthens our confidence in the results. The experts' ability to include detailed
4 comments on each of the draft statements enabled us to improve them, as reflected in
5 the increasing level of agreement with the statements in the second round, from
6 93.05% in the R1-survey to 97.8% in the R2-survey. Unlike NAFLD and CKD where
7 after 40 years there has been an organic consensus, for MAFLD and CKD we are just
8 beginning to acquire the relevant data to set a baseline for ongoing improvements in
9 knowledge.

10

11 **Conclusion**

12 MAFLD and CKD are two highly prevalent and interconnected conditions, posing a
13 challenge to global public health. In this Delphi-based consensus statement, several
14 international experts from different countries developed and endorsed a set of
15 consensus statements that provide guidance on the epidemiology, mechanisms,
16 management and treatment of MAFLD and CKD, as well as the relationship between
17 the severity of MAFLD and risk of CKD. These consensus statements establish a
18 framework for the early prevention and management of these two common and
19 interconnected diseases.

20

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12

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31

1 **Figure legends**

2 **Figure 1.** Flow diagram of the Delphi process adopted for the development of
3 consensus statements on MAFLD and the risk of CKD.

4

5 **Figure 2.** Scores for agreement in Delphi process. Scores for agreement by experts in
6 round 1 and round 2 (A); and the total scores for agreement and somewhat agreement
7 of experts in round 1 and round 2 (B).

8

9 **Table legends**

10 **Table 1.** Demographic composition of the expert panel

11 **Table 2.** Consensus statements on MAFLD and risk of CKD

12 **Supplementary table 1.** Comparison between MAFLD and NAFLD for the
13 identification of CKD

14 **Supplementary table 2.** Results of round 1 of the Delphi process

15 **Supplementary table 3.** Results of round 2 of the Delphi process