Liver Transplantation for Patients with Non-Alcoholic Steato-Hepatitis

First Edition
April 2011

British Transplantation Society Guidelines
## Contents

1. Guideline development ........................................ 3
2. Grading of recommendations .................................. 5
3. Summary of recommendations ................................ 6
4. Abbreviations ..................................................... 11
5. Prevalence of NASH cirrhosis in the UK .................... 12
6. Indications for liver transplantation in NASH-related cirrhosis 14
7. Assessment of operative risk in NAFLD patients undergoing liver transplantation 16
8. Assessment and management of nutritional status during transplant work-up 23
9. Surgical aspects of liver transplantation for patients with NAFLD 27
10. Peri-operative monitoring ..................................... 30
11. Immunosuppression ........................................... 33
12. Post-transplant monitoring of NASH patients and disease recurrence 36
13. Post-transplant management of NAFLD .................... 39
14. Non-alcoholic fatty liver disease in children ................ 43
15. Transplantation for NASH – the patients’ perspective 45
16. Statements of potential conflicts of interest ................. 47
17. References ....................................................... 50
1 Guideline development

NASH is an increasing cause of liver disease necessitating liver transplantation. In patients with advanced NASH, there are often co-existent clinical issues that impact on the outcome of liver transplantation.

There are no guidelines for the assessment and management of patients with NASH undergoing liver transplantation. A group was therefore invited by the Council of the British Transplantation Society (BTS) to prepare guidelines for the management of NASH before and after liver transplantation. The first draft was written by Dr PN Newsome (Senior Lecturer and Consultant Hepatologist, Liver Unit, University Hospital Birmingham NHS Foundation Trust) in Autumn 2010 with contributions from the following guideline group: Dr P Henriksen (Consultant Cardiologist and Honorary Senior Lecturer, Edinburgh Heart Centre, NHS Lothian, University Hospitals Division), Prof CP Day (Professor of Liver Medicine, Institute of Cellular Medicine, Newcastle University), Dr D Thorburn (Consultant Hepatologist, Liver Unit, Royal Free Hospital, London), Mr DF Mirza (Consultant Hepatobiliary and Transplant Surgeon, Liver Unit, University Hospital Birmingham NHS Foundation Trust), Dr JW Ferguson (Consultant Hepatologist and Honorary Senior Lecturer, Liver Unit, University Hospital Birmingham NHS Foundation Trust), Dr G Auzinger (Consultant Intensive Care Medicine, Liver Intensive Therapy Unit, King’s College Hospital London NHS Foundation Trust), Dr M Allison (Consultant Hepatologist, Liver Unit, Department of Medicine, Cambridge University Hospital NHS Foundation Trust), Dr JW Tomlinson (Reader in Endocrinology, Centre for Endocrinology, Diabetes & Metabolism, University of Birmingham), Ms H Manley (British Liver Trust), Dr KJ Simpson (Senior Lecturer in Hepatology, University of Edinburgh & Hon Consultant Physician Scottish Liver Transplantation Unit, Royal Infirmary Edinburgh), Prof SG Hubscher (Leith Professor and Professor of Hepatic Pathology, University of Birmingham and Consultant Histopathologist, University Hospital Birmingham NHS Foundation Trust), Dr C Millson (Consultant Hepatologist, St James’s University Hospital, Leeds), Dr J Oben (Wellcome Trust Senior Lecturer and Consultant Hepatologist, University College London, Centre for Hepatology, Royal Free Hospital, Rowland Hill, London NW3 2PF), Prof JM Neuberger (Associate Medical Director for Organ Donation and Transplantation, NHS Blood and Transplant and Honorary Consultant Physician Queen Elizabeth Hospital Birmingham), Dr PJ McKiernan (Consultant Paediatrician, Liver Unit, Birmingham Children’s Hospital) and Dr JI Wyatt (Consultant Histopathologist, St James’s University Hospital, Leeds).
This followed a systematic review of the literature using retrieval from electronic databases and reading suggestions from colleagues.

The document was revised in the autumn and winter of 2010, principally by Dr PN Newsome and Dr PA Andrews (Chair, BTS Standards Committee). The last date of literature review was November 2010. A draft version was circulated to members of the BTS Council and placed on the BTS website for comment in March 2011. The final version was revised in the light of comments received and published in April 2011.

These guidelines represent consensus opinion from experts in the United Kingdom in the fields of hepatology, transplantation and related disciplines. They represent a snapshot of the evidence available at the time of writing. It is recognised that recommendations are made even when the evidence is weak. It is felt that this is helpful to clinicians in daily practice and is similar to the approach adopted in other BTS guidelines. Although it is believed that the information presented is a fair summary of current evidence and best practice, neither the authors nor the British Transplantation Society can be held responsible for any errors or omissions. The guidelines are not designed to be prescriptive, nor to define a standard of care. Doses of prescribed drugs should always be checked by the responsible clinician according to the relevant information provided by the manufacturers of the drugs.

It is anticipated that these guidelines will be revised in 2015.
2 Grading of recommendations

For each recommendation, assessments have been made of the quality of supporting evidence and the strength of the recommendation. This is in keeping with other national guideline groups (1,2).

For each recommendation, the **quality of evidence** has been graded as one of:

- **Level A** Data derived from multiple randomised clinical trials or meta-analyses.
- **Level B** Data derived from a single randomised trial, or non-randomised studies.
- **Level C** Consensus opinion of experts, case studies, or standard-of-care.

For each recommendation, the **strength of recommendation** has been indicated as one of:

- **Class I** Conditions for which there is evidence and/or general agreement that a given evaluation, procedure or treatment is beneficial and effective.
- **Class II** Conditions for which there is conflicting evidence and/or a divergence of opinion about the usefulness/efficacy of an evaluation, procedure or treatment.
- **Class IIa** Weight of evidence/opinion is in favour of usefulness/efficacy.
- **Class IIb** Usefulness/efficacy is less well established by evidence/opinion.
- **Class III** Conditions for which there is evidence and/or general agreement that an evaluation/procedure/treatment is not useful/effective and in some cases may be harmful.
3 Summary of recommendations

1. Criteria for diagnosis of NASH group should include an established clinical and histological diagnosis of NASH on liver biopsy, or a histological diagnosis of cryptogenic cirrhosis with a clinical phenotype compatible with underlying NASH, as defined by the presence of 3 or more components of the metabolic syndrome prior to LT. (Level C, Class I)

2. Criteria for consideration of listing for liver transplantation in patients with NASH cirrhosis either due to ESLD or presence of hepatocellular carcinoma should be in line with standard national criteria. (Level C, Class I)

3. Although the diagnosis of NASH cirrhosis should not be regarded, in itself, to be a risk factor for poor outcome in the peri-operative period of liver transplantation, cardiovascular risk should be closely considered in patients with NASH as they have more risk factors for cardiovascular disease and are likely to require further non-invasive testing. (Level C, Class I)

4. All NAFLD patients should undergo pre-operative risk stratification to exclude symptomatic coronary artery disease and assessment for the presence of structural heart disease, left ventricular dysfunction and pulmonary hypertension. (Level C, Class I)

5. Cardiovascular risk should be closely considered in patients with NAFLD as many will require further non-invasive testing. (Level C, Class I)

6. Patients unable to achieve 4 METs, or those with at least two intermediate risk factors, should be considered for further cardiac testing. (Level C, Class I)

7. Within a transplant centre, cardiology input should be provided by cardiologists with an interest/experience in the assessment of patients with liver disease. (Level C, Class I)

8. The following moderate risk groups should be discussed early with a cardiologist: (i) patients with chest pain of possible cardiac origin; and (ii) patients receiving treatment for established coronary disease or previous coronary revascularisation. (Level C, Class I)
9. There is insufficient evidence to recommend a single stress test for NAFLD patients undergoing OLT assessment. The choice of test will be in part determined by local expertise. (Level C, Class I)

10. Patients should receive a β-blocker prior to liver transplantation. This should be titrated gradually and not started in the immediate peri-operative period. (Level B, Class IIa)

11. Statins should be started between 30 days and at least 1 week before surgery, or continued in those patients already receiving them. (Level B, Class I)

12. There are no data to support an absolute cut-off for body mass index (BMI) and liver transplantation, although patients with a BMI >40 kg/m² are likely to have an increased post-operative and long-term mortality. BMI should be corrected for the presence of ascites and peripheral oedema. (Level C, Class IIa)

13. Weight loss should not be recommended in all patients with end-stage liver disease due to the risk of protein-calorie malnutrition. (Level C, Class III)

14. For patients with stable cirrhosis and hepatocellular carcinoma, it may be appropriate to try and achieve weight loss before proceeding/whilst waiting for a transplant. (Level C, Class II)

15. All potential NAFLD transplant recipients, including those with apparent obesity, should be assessed by a dietician and supplemental feeding considered if required. (Level C, Class I)

16. Dietary assessment of patients being assessed for liver transplantation should include use of hand-grip strength, anthropometry and/or subjective global assessment to objectively define the patient’s nutritional status and allow supplementation if required. (Level B, Class I)

17. Dietary assessment of patients should be repeated on an annual basis whilst they remain on the waiting list for liver transplantation. (Level B, Class I)

18. Consider bariatric surgery at the time of liver transplantation in recipients with severe morbid obesity, those with failed treatment of obesity or in patients with recurrent disease undergoing retransplantation. (Level C, Class IIa)
19. Consider bariatric surgery in recipients with severe morbid obesity, those with failed treatment of obesity, or in patients developing progressive NASH with fibrosis in the allograft. (Level C, Class IIa)

20. Whilst there is likely to be an increased operative risk, the lack of evidence from controlled clinical trials indicates that no recommendation can be made about the use of intra-operative cardiac output monitoring. (Level B, Class II)

21. Moderately tight glucose control (6-10 mmol/l) should be targeted during the early post transplant course in patients of all aetiologies. (Level A, Class II)

22. If started pre-operatively, statin therapy should be continued during the post-operative phase. (Level B, Class I)

23. Alongside immunosuppression with CNI and anti-metabolite, consideration should be given to either a steroid-free regime or early steroid withdrawal (within three months) in patients with NASH. Where steroid-free regimes are used, induction therapy (such as ATG or IL2-R antagonism) should be considered. (Level B, Class IIa)

24. Tacrolimus levels should be <10 ng/ml within the first three months after liver transplantation and 5-8 ng/ml after that to reduce the impact on renal function and dyslipidaemia. Mycophenolate should be used as the preferred anti-metabolite, to permit lower levels of tacrolimus. (Level B, Class IIa)

25. Close follow-up and early recognition and treatment of the recognised consequences of transplantation and immunosuppression (such as weight gain, hypertension, hyperlipidaemia, diabetes and renal impairment) remain the key to preventing excess risk from recurrent NAFLD. (Level C, Class I)

26. Histological examination of the explanted liver should be carried out to confirm the presence of features compatible with end-stage NASH and to exclude features suggesting an alternative diagnosis. (Class I, Level C)
27. The main role of biopsy is to allow diagnosis and staging of liver histopathology. Where NAFLD is the only or dominant pathology, liver allograft biopsies can be scored using the Kleiner classification. Biopsies performed elsewhere should be reviewed at the transplant centre to ensure reproducibility. (Class I, Level C)

28. Post-transplant monitoring of patients should include an initial USS at one year, followed by every two years, looking for the presence of an echobright liver. (Class IIa, Level C)

29. Post-transplant monitoring of patients with echobright liver on USS should include protocol liver biopsies to detect disease recurrence, as liver function tests may be normal. Repeat biopsy should be considered every three years, unless there is a clinical indication for more frequent biopsies. (Class IIa, Level C)

30. Post-transplant patients should receive support, advice and treatment in order achieve a target body mass index of <25 kg/m². This should be in the context of a multidisciplinary team, incorporating dietary modification, exercise intervention and the potential use of pharmacotherapy and surgical intervention where appropriate. (Level C, Class I)

31. Post-transplant patients should be screened for the presence of diabetes and, if present, reviewed regularly for the development of complications. Glycaemic control should be optimised in accordance with NICE guidance. (Level A, Class I)

32. Patients transplanted for NAFLD should be monitored on a 6-monthly basis for risk factors for cardiovascular disease (BP, lipids, HbA1c), which should be addressed with the intention of reducing cardiovascular events. (Level II-3, Level C)

33. A blood pressure target of 140/90 mmHg (130/80 mmHg in patients with diabetes and/or renal dysfunction) should be aimed for (Level A, Class I).

34. Anti-hypertensive agents such as calcium channel blockers or ACE inhibitors should be considered in view of their possible additional effects of abrogating liver fibrosis. (Level C, Class II)
35. A target LDL cholesterol of <2.6 mmol/l is advised as the 10-year cardiovascular event rate exceeds 20% for the liver transplant population. Pravastatin and ezetimibe are preferred agents in view of their demonstrated efficacy and absence of interactions with CNIs. (Level C, Class IIa)

36. There is a need to increase understanding of liver disease and its many causes, to improve patient outcomes and to reduce the stigma many patients experience. (Not Graded)

37. The potential cardiovascular morbidity associated with NASH should be discussed with patients and guidance given on diet and exercise, and sources of support (including psychological support) as part of ongoing management. (Not Graded)

38. Provision of independent pre- and post transplant emotional and psychological counselling and support is very important, along with an opportunity to provide confidential feedback to the transplant team post-operatively. (Not Graded)

39. Patients with NASH should be referred to specialist centres for optimal management and consideration for clinical trials. (Not Graded)
### 4 Abbreviations

<table>
<thead>
<tr>
<th>Abbreviation</th>
<th>Description</th>
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<tbody>
<tr>
<td>ATG</td>
<td>Anti-thymocyte globulin</td>
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<tr>
<td>CKD</td>
<td>Chronic kidney disease</td>
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<tr>
<td>CNI</td>
<td>Calcineurin inhibitor</td>
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<tr>
<td>CPX</td>
<td>Cardiopulmonary exercise</td>
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<tr>
<td>DSE</td>
<td>Dobutamine stress echocardiography</td>
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<tr>
<td>ESLD</td>
<td>End-stage liver disease</td>
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<tr>
<td>HCC</td>
<td>Hepatocellular carcinoma</td>
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<tr>
<td>HCV</td>
<td>Hepatitis C virus</td>
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<tr>
<td>IGB</td>
<td>Intra-gastric balloon</td>
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<tr>
<td>IMS</td>
<td>Immunosuppression</td>
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<tr>
<td>LV</td>
<td>Left ventricular</td>
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<tr>
<td>LT</td>
<td>Liver transplant</td>
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<tr>
<td>MELD</td>
<td>Model for end-stage liver disease</td>
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<tr>
<td>MET</td>
<td>Metabolic equivalent of task</td>
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<tr>
<td>MI</td>
<td>Myocardial infarction</td>
</tr>
<tr>
<td>mPAP</td>
<td>Mean pulmonary artery pressure</td>
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<tr>
<td>mTORi</td>
<td>Mammalian target of rapamycin inhibitor</td>
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<tr>
<td>NAFLD</td>
<td>Non-alcoholic fatty liver disease</td>
</tr>
<tr>
<td>NASH</td>
<td>Non-alcoholic steato-hepatitis</td>
</tr>
<tr>
<td>NHSBT</td>
<td>National Health Service Blood and Transplant</td>
</tr>
<tr>
<td>NODAT</td>
<td>New onset diabetes after transplantation</td>
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<tr>
<td>OLT</td>
<td>Orthotopic liver transplant</td>
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<tr>
<td>PASP</td>
<td>Pulmonary artery systolic pressure</td>
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<tr>
<td>PAC</td>
<td>Pulmonary artery catheter</td>
</tr>
<tr>
<td>PCWP</td>
<td>Pulmonary capillary wedge pressure</td>
</tr>
<tr>
<td>PCM</td>
<td>Protein-calorie malnutrition</td>
</tr>
<tr>
<td>RWMA</td>
<td>Regional wall motion abnormalities</td>
</tr>
<tr>
<td>RV</td>
<td>Right ventricular</td>
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<tr>
<td>SGA</td>
<td>Subjective global assessment</td>
</tr>
<tr>
<td>TOE</td>
<td>Trans-oesophageal echocardiography</td>
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<tr>
<td>TR</td>
<td>Tricuspid regurgitation</td>
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</table>
5 Prevalence of NASH cirrhosis in the UK

Non-alcoholic fatty liver disease (NAFLD) encompasses a spectrum of disease ranging from simple steatosis, to steatohepatitis (NASH) and cirrhosis. NAFLD is closely associated with obesity and represents the hepatic manifestation of the metabolic syndrome. The prevalence of NAFLD has risen rapidly in parallel with the dramatic rise in levels of obesity and diabetes mellitus (3), resulting in it now being the commonest cause of liver disease in the West (4).

NAFLD prevalence

The prevalence of NAFLD is between 20-30% in Western adults (5,6), rising to 90% in extreme obesity (7). NAFLD affects 3% of the general paediatric population, rising to 53% in obese children (8,9), with implications for future disease burden. NASH, the more advanced and clinically important form of NAFLD, has an estimated prevalence of 2-3% in the general population (10) and 37% in the morbidly obese (7). Steatosis was present in 70% of a large cohort of patients with type 2 diabetes (11). The Foresight report predicted that with the alarming growth of obesity, the burden of NAFLD on primary care and liver services will double from a current annual cost of £4.2 billion by 2050 (12).

Association with increased mortality and progression to liver cirrhosis

Patients with a diagnosis of NAFLD have been shown to have a significantly higher overall (13-17) and liver-related (14,15) mortality when compared with an age/sex-matched general population. In patients with NASH, the limited data available point to approximately one third of patients developing progressive fibrosis over a 5 period, with up to 9% developing cirrhosis (18,19). The risk factors for progressive NASH-related liver disease are obesity, type 2 diabetes mellitus, insulin resistance and older age. Current evidence suggests that the natural history of NASH cirrhosis is similar to that of hepatitis C cirrhosis with respect to decompensation (20,21). Hepatocellular carcinoma (HCC), a recognised complication of cirrhosis of a number of aetiologies, is also known to occur in NASH-related cirrhosis, and also rarely in pre-cirrhotic NASH (22). A prospective study on NASH cirrhotics has found a cumulative incidence of 2.6% for the development of HCC in this patient group compared to 4.0% for patients with Hepatitis C virus (HCV) cirrhosis (23). Obesity
and diabetes have been found to be risk factors for the development of HCC in cirrhosis of a variety of aetiologies (24,25). The mortality of patients with NAFLD ranges from 12.6% over 7.6 years follow-up in mixed cohorts to between 20.2-59.5% in secondary care cohorts (13.7-21 years follow-up). In the most recent study with 21 years of follow-up, this corresponded to an excess mortality of 70% (standardised mortality ratio 1.7; 95% CI 1.24-2.25)(16).

**Need for liver transplantation**

The increase in the number of patients with advanced liver disease secondary to NASH, as well as associated HCC, will impact on the potential future demand for liver transplantation (LT). Analysis of National Health Service Blood and Transplant (NHSBT) data shows that in both 2008 and 2009, 12% of patients placed on the elective LT waiting list were categorised as having NASH cirrhosis, with 14.8% of these individuals listed with HCC. Consistent with the known association with insulin resistance, 49.1% of patients with NASH cirrhosis listed for LT were diabetic compared to 22.1% of all other registered patients (although the criteria for the definition of diabetes were not well defined).
6 Indications for liver transplantation in NASH-related cirrhosis

Given that liver transplantation is a relatively recently identified indication for patients with NASH, data for long term follow-up are more limited when compared with LT for the other causes of chronic liver disease. Nevertheless, data from several American centres suggest that patients with NASH do not have an inferior outcome 1, 2 and 5 years after LT when compared to other aetiologies of liver disease (26-28).

There are no data to suggest that the natural history of cirrhosis after an initial clinical decompensation, or the rate of change of model for end-stage liver disease (MELD) score differ for patients with NASH. Hence, the current recommendation should be that there are no disease-specific indications for listing for LT in this condition and that those currently used, i.e. MELD score and/or specific variant indications (e.g. diuretic resistant/intolerant ascites) should remain. In those patients with hepatocellular carcinoma on the background of NASH-related cirrhosis, there are no data to support modification of the general criteria for listing for LT.

**Recommendation 1:** Criteria for diagnosis of NASH group should include an established clinical and histological diagnosis of NASH on liver biopsy, or a histological diagnosis of cryptogenic cirrhosis with a clinical phenotype compatible with underlying NASH, as defined by the presence of 3 or more components of the metabolic syndrome prior to LT. (Level C, Class I).

**Recommendation 2:** Criteria for consideration of listing for liver transplantation in patients with NASH cirrhosis either due to ESLD or the presence of hepatocellular carcinoma should be in line with standard national criteria. (Level C, Class I)

In an era of significant organ-shortage, with current mortality on the elective liver transplant waiting list being around 18% in the UK, there is a duty to allocate organs responsibly. The national guidelines for adult LT are that patients should only be listed for LT if they have a predicted 5 year survival after transplantation of ≥50% (NHSBT Adult Liver Transplantation Guidelines 2009, accessed 19 February 2011 at www.organdonation.nhs.uk/ukt/about _transplants/organ_allocation/liver/national_protocols_and_guidelines/adults.jsp). Standard exclusion criteria will apply to NAFLD patients being considered for LT, although their strong association with the metabolic syndrome will likely involve a more stringent assessment of cardiovascular risk.
7 Assessment of operative risk in NAFLD patients undergoing liver transplantation

Having identified that a patient may benefit from liver transplantation, the next stage in the process involves an assessment of the risk to the patient of proceeding with the operation. The risks can be divided into short (up to 90 days) and long-term (1 year onwards) to cover the immediate risks of the operation itself and then the longer term survival. In both of these categories there is concern that patients with NASH cirrhosis are at higher risk.

Risk of cardiovascular events

Cardiovascular risk factors are commonly found in patients undergoing LT, with hyperinsulinaemia, type II diabetes, hypertension, hyperlipidaemia and abdominal obesity disproportionately present in NASH patients (11,29). Myocardial disease and dysfunction are common in cirrhotic patients. Indices of myocardial contractility such as left ventricular stroke work are increased at rest (30). A sustained increase in baseline cardiac output leads to cardiac hypertrophy and increases in left ventricular diastolic and left atrial pressures (30). Cirrhotic patients may have a normal resting ejection fraction but their reserve and ability to increase heart rate and cardiac output in response to stress are reduced, predisposing to decompensation during the anhepatic phase and following reperfusion (31,32).

The above is of particular relevance to a significant proportion of patients with NASH-related cirrhosis as they suffer from end-organ damage from lipotoxicity and insulin resistance. The presence of pre-transplant insulin treated diabetes mellitus has an adverse impact on long-term post-transplant survival (33), and many individuals with NASH-related cirrhosis have diabetic microvascular and macrovascular complications. Indeed atherosclerosis has been found to be present in a higher proportion of patients with cirrhosis due to NASH compared to cirrhosis of other aetiologies (34). Other diabetic complications such as proliferative retinopathy, diabetic nephropathy, cardiovascular autonomic neuropathy, and diabetic foot ulcer disease have also been shown to predict an increased morbidity/mortality in diabetic cohorts outside the context of organ transplantation (35-37). A further area of perceived increased risk relevant to the NASH cirrhotic population is that of the effect of obesity on LT outcome, which will be covered in the following section.
Whilst there are data which suggest that NASH is an independent risk factor for cardiovascular disease (38), further large scale prospective studies are needed to draw firm conclusions about the independent hepatic contribution to the increased cardiovascular risk seen in patients with NASH (29). Notably NASH has been reported to independently predict chronic kidney disease (CKD), which may explain part of the contribution to cardiovascular risk (39).

**Recommendation 3:** Although the diagnosis of NASH cirrhosis should not be regarded in itself to be a risk factor for poor outcome in the peri-operative period of liver transplantation, cardiovascular risk should be closely considered in patients with NASH as they have more risk factors for cardiovascular disease and are likely to require further non-invasive testing. (Level C, Class I)

**Cardiovascular event rate after liver transplantation**

Cardiovascular complications are a common cause of early morbidity and mortality after liver transplant (40,41). Analysis of over 21,000 patients undergoing OLT in the European Liver Transplant Registry reported a 3-month mortality of 12% (42), with 8% of deaths secondary to cardiac and a further 7% secondary to cerebrovascular causes. In a study of 413 patients of varying causes of ESLD undergoing LT between 2001 and 2005, the 30-day all cause mortality was 9%, with 7% developing a myocardial infarction (as defined by post-operative troponin elevation) within that time period. A history of previous stroke or coronary disease was associated with a 6.5-fold and 4-fold increased risk of non-fatal MI and death respectively (43).

Pre-operative cardiac risk assessment relies on the presence of clinical risk markers, the patient’s functional capacity, and the extent or risk of the surgical procedure. Liver transplantation is considered intermediate risk surgery when compared to higher risk surgical procedures such as aortic surgery in European and American guidelines for pre-operative cardiac risk assessment (44,45). However, the 1-5% 30-day risk of myocardial infarction and cardiac death quoted for intermediate risk surgery in these guidelines (44) is less than the rate of cardiovascular complications observed in many OLT series (46-48), suggesting that LT surgery should more probably be considered as moderate to high risk.
**Research/Audit recommendations:**

1. It is important to collect data on cardiovascular events for patients undergoing LT with NASH to establish if their event rate differs from cohorts transplanted for other indications.
2. To allow for appropriate comparisons it is important to define what constitutes a cardiovascular event in terms of type of event and time since the operation.

**Pre-operative selection and assessment**

**Phase I: Looking for factors to risk stratify**

An accurate and reliable means of predicting short and long-term post-LT prediction in an individual is clearly desirable, but not currently available. Several factors which confer increased risk have been identified (Table 7.1), but data on their risk when added together is less clear (44). The history should seek to determine the patient’s functional capacity, as assessment of the capacity to perform a range of common daily tasks has been shown to correlate well with maximum oxygen uptake on treadmill testing (49). The metabolic equivalent of task (MET) is an expression of energy expenditure during physical activity as a multiple of resting metabolic rate. One MET is considered as the resting metabolic rate, two METs would represent walking at a slow pace (e.g. 3 km/h) and four METs would represent walking up two sets of stairs.

Pre-operative cardiorespiratory assessment in patients with NASH does not differ to that routinely performed in other potential liver transplant recipients. Standard tests include chest radiography, arterial blood gases, pulmonary function tests, 12 lead ECG, and echocardiogram (with the following parameters recorded: pulmonary artery systolic pressure (PASP), right ventricular (RV) function, left ventricular (LV) function, tricuspid regurgitation (TR) velocity, iso-volumic relaxation time).

A retrospective analysis of patients transplanted with NASH cirrhosis found that the combination of age ≥60 years, BMI ≥30 kg/m², diabetes and hypertension predicted a 50% mortality within one year of transplantation (27). This needs to be confirmed in larger cohorts, but is consistent with another analysis of predictors of increased long-term mortality post-LT (50). An attempt to incorporate co-morbidity factors in a summative fashion to predict long-term outcome after LT using a modified co-morbidity index (Charlson Index) also found that the pre-transplant presence
of more than one defined risk factor for patients undergoing elective first transplantation resulted in a 5-year post-LT survival of <50% (51).

Stratification of patients to various risk categories on the basis of risk markers varies across guidelines, but one unifying principle is the assessment of a patient’s functional state as judged by METs. Both European (44) and American (52) guidelines of cardiovascular evaluation and care for patients undergoing non-cardiac surgery suggest that those with good functional capacity (≥4 METs) without symptoms do not require non-invasive testing prior to LT. This is in contrast to the AASLD guidelines, which recommend that potential liver transplant recipients should undergo further cardiac investigation with dobutamine stress echocardiography and confirmatory angiography if they have any of the following factors: age >50 years, a history of chronic smoking, a past medical or family history of heart disease, or diabetes (53).

In the event that a patient’s functional capacity is reduced to below 4 METs, the presence of between 1 (52) and 3 (44) intermediate risk factors are required to stratify a patient as moderate risk and therefore indicate the need for non-invasive testing.

Acute heart failure, symptomatic valvular disease and significant pulmonary hypertension (mean pulmonary artery pressure >25 mmHg at rest and a pulmonary capillary wedge pressure <15 mmHg (54)) will generally preclude patients being listed for OLT. Some of these conditions may respond to treatment and options are expanding with improved management of heart failure and pulmonary hypertension, with some centres reporting successful combined cardiac and OLT surgery.

**Recommendation 4:** All NAFLD patients should undergo preoperative risk stratification to exclude symptomatic coronary artery disease and assessment for the presence of structural heart disease, left ventricular dysfunction and pulmonary hypertension. (Level C, Class I)

**Recommendation 5:** Cardiovascular risk should be more closely considered in patients with NAFLD as many will require further non-invasive testing. (Level C, Class I)

**Recommendation 6:** Patients unable to achieve 4 METs, or those with at least two intermediate risk factors, should be considered for further cardiac testing. (Level C, Class I)
An algorithm to aid in the investigation of cardiovascular risk patients with NASH cirrhosis being considered for LT is detailed in Figure 7.1.

Table 7.1 Risk markers for cardiac decompensation during OLT surgery

<table>
<thead>
<tr>
<th>Low Risk</th>
<th>Intermediate Risk</th>
<th>High Risk</th>
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<tbody>
<tr>
<td>No CVS risk factors</td>
<td>Angina pectoris</td>
<td>Acute heart failure</td>
</tr>
<tr>
<td>No co-morbidities</td>
<td>Prior myocardial infarction</td>
<td>Symptomatic valvular heart disease</td>
</tr>
<tr>
<td>Good functional capacity (&gt;4 Mets)</td>
<td>Heart failure</td>
<td>Significant pulmonary hypertension</td>
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<td></td>
<td>Stroke/transient ischaemic attack</td>
<td>Recent (within 30 days) acute coronary syndrome and/or evidence of ongoing myocardial ischaemia</td>
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<td></td>
<td>Renal dysfunction (serum creatinine &gt;170 µmol/l or &gt;2 mg/dl or a creatinine clearance of &gt;60 ml/min) or need for renal support</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Diabetes mellitus requiring insulin therapy</td>
<td>Significant cardiac arrhythmias</td>
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**Phase II: Role of cardiac testing**

Assessment of functional capacity can be difficult in patients with ESLD due to NASH because of the high incidence of reduced mobility, physical deconditioning and obesity. Whilst risk stratification will determine the next level of cardiovascular investigation, early discussion with a cardiologist experienced in assessing cardiovascular risk in patients with ESLD should be considered for more complex cases.

**Recommendation 7:** Within a transplant centre, cardiology input should be provided by cardiologists with an interest/experience in the assessment of patients with liver disease. (Level C, Class I)
**Recommendation 8:** The following moderate risk groups should be discussed early with a cardiologist: (i) patients with chest pain of possible cardiac origin; and (ii) patients receiving treatment for established coronary disease or previous coronary revascularisation. (Level C, Class I)

The role of the cardiologist includes, but is not limited to the following: (i) coronary angiography (invasive or by CT); (ii) functional testing with myocardial perfusion imaging; (iii) optimisation of pharmacotherapy; (iv) coronary revascularisation; and (v) advice on peri-operative cardiac management in patients accepted for OLT.

Cardiopulmonary exercise (CPX) testing predicts survival in liver transplant candidates and following liver transplantation in general. In two studies, a threshold level of <60% peak oxygen consumption during CPX independently predicted survival and length of hospital stay (55,56).

Cardiac stress testing is directed towards establishing the presence of obstructive coronary disease, but it appears to have a more limited predictive value for peri-operative cardiac decompensation in this patient population when compared to other high risk surgical groups. Dobutamine stress echocardiography (DSE), although moderately specific for coronary artery disease (CAD), lacks sensitivity in patients undergoing OLT (57,58). Guidelines on pre-operative cardiac risk assessment indicate that patients with multiple cardiac risk markers or poor functional status undergoing high risk surgery are recommended for stress testing by perfusion imaging or DSE (44,45). Negative stress testing is reassuring providing an excellent negative predictive value in patients undergoing major vascular surgery (59,60). The positive predictive value for peri-operative myocardial infarction or death in this high risk surgery group is between 20 and 30% following a positive stress test (60). Perfusion imaging and DSE detect the presence of fixed obstructive coronary stenoses responsible for myocardial ischaemia during increased myocardial workload. The presence of obstructive coronary disease increases the risk of peri-operative myocardial infarction but other factors contribute including changes in coagulation and platelet activation during the peri-operative period (61).

According to Bayesian principles, the positive predictive value of a screening test depends on the prevalence of disease in the population tested. Patients with established vascular disease are more likely to have (asymptomatic) coronary disease than patients with liver cirrhosis. This may partly explain the comparably weak positive predictive value of DSE in liver transplant patients.
Recent studies have also reported poor discrimination for identifying patients at risk of perioperative troponin elevation and cardiac decompensation (58,62). Many transplant centres use DSE as a preoperative screening test and the most recent AASLD guideline recommends DSE in smokers, patients with coronary disease or type II diabetes mellitus (53). A positive test mandates coronary angiography according to this guideline, but patients with non-obstructed coronary arteries may still experience cardiac decompensation during liver transplantation (63). The predictive value of other non-invasive methods such as ⁹⁹Tc-labelled single photon emission computer tomographic studies is likewise poor (64).

Calcification of coronary arteries on CT scanning is correlated with cardiovascular risk factors and the development of coronary atherosclerosis. Coronary calcification is seen in patients with the metabolic syndrome who are assessed for liver transplantation (65). Coronary calcium scoring may be used to further define the 10-year risk of cardiovascular events in asymptomatic populations (66). It does not provide information on the presence of obstructive coronary disease or inducible myocardial ischaemia and there is no evidence to support its use in peri-operative risk assessment.

The above tests may aid in triaging patients in regards to diagnostic coronary investigations. Apart from standard invasive coronary angiography, CT coronary angiography is used with increasing frequency in the assessment of patients with possible cardiac chest pain. Despite the advantage of minimal invasiveness, it is limited in its ability to discriminate obstructive coronary lesions by the presence of coronary calcification and coronary stents. CT coronary angiography has been used to rule out coronary disease in patients undergoing heart valve surgery and non-cardiac surgery. Recent guidelines have suggested an application to further evaluate patients considered at intermediate risk of coronary disease undergoing non-cardiac surgery, but it is currently not a recommended technique for peri-operative risk assessment in non-cardiac surgery (44,67). Further research is required and no recommendation was given for this imaging modality in the most recent joint European Society of Cardiology and European Society of Anaesthesiology guideline on pre-operative cardiac risk assessment (44).

**Recommendation 9:** There is insufficient evidence to recommend a single stress test for NAFLD patients undergoing OLT assessment. The choice of test will be in part determined by local expertise. (Level C, Class I)
Pharmacological risk reduction with β-blockers and statins has been extensively studied. Randomised trials selecting high risk patients, cohort studies, and meta-analyses provide consistent evidence supporting a decrease in cardiac mortality and MI using β-blockers in patients with clinical risk factors undergoing high-risk surgery. Treatment should be titrated and initiated ideally between 30 days and at least 1 week before surgery, with suggested targets of heart rate 60-70 beats/min and systolic blood pressure >100 mmHg (44). Higher dose β-blockade started in the peri-operative period was associated with increased mortality, reinforcing the need for gradual titrated introduction of therapy (68).

**Recommendation 10**: Patients should receive a β-blocker prior to liver transplantation. This should be titrated gradually and not started in the immediate peri-operative period. (Level B, Class IIa)

Similarly, starting statins has been shown to reduce cardiovascular mortality by 44% in meta-analyses in patients undergoing non-cardiac surgery (69). Stopping of statins may also result in a rebound effect and lead to increased cardiovascular complications (70).

**Recommendation 11**: Statins should be started between 30 days and at least 1 week before surgery, or continued in those patients already receiving them. (Level B, Class I)
8 Assessment and management of nutritional status during transplant work-up

Peri-operative morbidity and mortality is increased following major surgical procedures among the obese because of concomitant cardiovascular disease, hyperlipidaemia and pulmonary dysfunction (71). The AASLD practice guideline, among others, suggests that morbid obesity should be considered a contraindication to liver transplantation (53,72).

Influence of obesity on post-operative liver transplant outcome

A consistent drawback with the majority of studies examining the influence of obesity on transplant outcome has been that the amount of ascites/peripheral oedema has not been taken into account when calculating BMI. Table 8.1 details how weight can be adjusted for ascites and peripheral oedema. This is important given that ascites is a marker of disease severity. Of the 12 published studies, only Leonard’s study (73) measured the ascites volume at transplantation and identified those with obesity as opposed to those with marked ascites (72-83). A second drawback is that the conclusions are drawn only from those patients with obesity who have undergone transplantation, so extrapolation from these to all potential candidates may be misleading.

<table>
<thead>
<tr>
<th></th>
<th>Ascites /kg</th>
<th>Peripheral oedema /kg</th>
</tr>
</thead>
<tbody>
<tr>
<td>Minimal</td>
<td>2.2</td>
<td>1</td>
</tr>
<tr>
<td>Moderate</td>
<td>6</td>
<td>5 (knees)</td>
</tr>
<tr>
<td>Severe</td>
<td>14 or more</td>
<td>10 (thigh)</td>
</tr>
</tbody>
</table>

Table 8.1 Correction for Fluid excess in Body Mass (84)

Analysis of the United Network for Organ Sharing (UNOS) database demonstrated that primary graft non-function and immediate, 1-year, and 2-year mortality were significantly higher in the morbidly obese group (p<0.05). In addition, five-year mortality was significantly higher in both
severely and morbidly obese subjects (p<0.05), mostly as a result of adverse cardiovascular events (72). Kaplan-Meier survival was significantly lower in morbidly obese patients, and morbid obesity was an independent predictor of mortality. Further analysis of the UNOS database in 2009 drew broadly similar conclusions, although it suggested that obese patients (BMI >40 mg/kg²) tended to be transplanted with more advanced MELD scores and hence these patients might be best served by receiving a graft at an earlier stage than non-morbidly obese counterparts (83). This is borne out by a large study of 29,000 patients wait-listed for liver transplants which found that obese recipients waited longer and were more likely to be passed over (85). Looking at UK liver transplant outcome data, a high BMI was an independent predictor of a worse outcome at 90 days, but not long-term (86).

The importance of correcting for ascites volume was demonstrated by Leonard et al, correction resulting in 11-20% of patients moving into a lower BMI classification (73). The relative risk for mortality increased by 7% for each litre of ascites removed, with the conclusion that corrected BMI is not an independent predictor of patient or graft survival. This would suggest that obesity should not be considered to be an absolute contraindication to liver transplantation in the absence of other relative contraindications. The decision to list a patient will include an assessment of all potential relative contra-indications, such as a BMI of >40 kg/m².

**Recommendation 12:** There are no data to support an absolute cut-off for body mass index (BMI) and liver transplantation, although patients with a BMI >40 kg/m² are likely to have an increased post-operative and long-term mortality. BMI should be corrected for the presence of ascites and peripheral oedema (Level C, Class IIa).

**Weight loss in liver transplant recipients**

There are no data on the impact of weight loss pre-transplant, nor are there any data on intervention studies post liver transplant or strategies for maintaining weight loss in transplant recipients.
Behaviour changes required for liver transplantation

For tobacco, alcohol or illicit substances, a requirement for complete abstinence is widely accepted and enforceable, and adherence can be checked. Food, however, is a basic requirement which cannot be proscribed, and food restriction in patients with end-stage liver disease is potentially harmful. Despite the appearance of obesity, some patients with advanced liver disease will have protein calorie malnutrition (PCM), and therefore should be fully assessed by a dietician for evidence of protein malnutrition. Dietary restriction risks worsening protein calorie malnutrition, and should only be contemplated under careful supervision by a dietician. PCM is associated with reduced graft and patient survival (87). A weight-orientated contract after LT is superficially attractive, to mimic the existing no-alcohol contract, but the practicalities would need careful consideration.

**Recommendation 13:** Weight loss should not be recommended in all patients with end-stage liver disease due to the risk of protein-calorie malnutrition. (Level C, Class III)

**Recommendation 14:** For patients with stable cirrhosis and HCC, it may be appropriate to try and achieve weight loss before proceeding/whilst waiting for a transplant. (Level C, Class II)

Assessment of patient nutritional status

Nutrition management in patients with advanced liver disease and obesity is problematic. Accurate estimation of nutritional status is difficult in patients with advanced liver disease, regardless of whether they are obese or not, in part due to poor inter-observer reproducibility and overestimation of values because of third spacing of fluid. PCM is a condition of body wasting related to dietary deficiency of calories and protein, is found in 65-90% of patients with ESLD (88), and is associated with reduced graft and patient survival (87,89). The detection and management of PCM is therefore an integral part of the dietary assessment.

Skinfold measurement and handgrip strength are simple tests of nutritional status, with the latter proving to be a good predictor of complications in patients with advanced liver disease (90). Other methods include the subjective global assessment (SGA), which combines multiple elements of nutritional assessment to classify the severity of malnutrition (91).
In 1997, the European Society for Clinical Nutrition and Metabolism developed guidelines for meeting nutritional goals in patients with end-stage liver disease (92). This included a diet consisting of 35% fat (15-20% monosaturates, minimal trans fats, reduced saturated fats, <10% total fat), 50% carbohydrate (minimal refined carbohydrate), and 15% protein. For obese patients with end-stage liver disease, the transplant team should be aware that dietary restriction risks enhancing protein calorie malnutrition, and should only be contemplated under careful supervision by a dietician. If safe, calorie restriction should be limited to target no more than 0.5-1 kg (1-2 lb) weight loss per week and to avoid weight loss exceeding 1.5 kg per week. In some circumstances malnutrition may be present despite apparent obesity, in which case supplementation with night time tube feeding may be of value (93). Patients with well preserved liver function and small hepatocellular cancers may benefit from weight loss, but only if the delay does not adversely affect their cancer.

**Recommendation 15:** All potential NAFLD transplant recipients, including those with apparent obesity, should be assessed by a dietician and supplemental feeding considered if required (Level C, Class I)

**Recommendation 16:** Dietary assessment of patients being assessed for liver transplantation should include use of hand-grip strength, anthropometry and/or subjective global assessment to objectively define the patient’s nutritional status, and start supplementation if required. (Level B, Class I)

**Recommendation 17:** Dietary assessment of patients should be repeated on an annual basis whilst they remain on the waiting list for liver transplantation. (Level B, Class I)
9 Surgical aspects of liver transplantation for patients with NAFLD

Peri-operative issues in NAFLD patients undergoing liver transplantation

Patients undergoing liver transplantation for NAFLD pose additional challenges for surgical teams. As the potential donor pool continues to evolve, the use of extended criteria donors in NAFLD recipients needs to be assessed, especially donor grafts with moderately severe steatosis. Such patients are also less likely to be suitable recipients for split liver and living related transplantation as the segmental liver graft is more likely to be relatively small for size for the obese NAFLD recipient. Consequently, patients with NAFLD may have a restricted access to the already severely limited donor organ pool, reducing their chances of undergoing liver transplantation.

Specialist theatre equipment in the form of heavy duty operating tables and retractor systems and additional surgical instruments are required for morbidly obese recipients. Few data exist regarding the use of steatotic donor livers and their impact on the development of post transplant recurrent NAFLD. As disease recurrence after liver transplantation is common and the diagnosis based on liver histology, the performance of a core liver biopsy after reperfusion and prior to abdominal closure is of value to provide baseline liver histology for comparison with late post transplant biopsies. These biopsies may also assist the evaluation of patients with early graft dysfunction, which these grafts may be more predisposed to.

Bariatric surgery at the time of transplantation

Liver transplant surgery provides an opportunity to perform additional bariatric surgical procedures to facilitate weight loss, and also limit the risk of recurrent disease. Such procedures do add an additional risk to what is already a high risk procedure. The presence of portal hypertension or post-reperfusion coagulopathy would be strong reasons to perform the least invasive additional bariatric surgical procedure. As a small proportion of liver transplant patients may require a future Roux-en-Y biliary enteric anastomosis, it is preferable to limit bariatric procedures to non-bypass procedures. Similarly, the use of gastric bands in the setting of advanced liver disease and subsequent post liver transplant immunosuppression may increase the risk of sepsis related to the device.
Almost no evidence exists on the role of bariatric surgery at the time of liver transplantation. It would therefore be logical to consider such an approach only in highly selected situations, possibly in recipients with severe morbid obesity, those with failed treatment of obesity, or in patients with recurrent disease undergoing retransplantation. The choice of procedure should be guided by the status of the recipient at the end of the transplant procedure, the essential requirements being the presence of satisfactory early liver graft function, adequate haemostatic parameters, control of all bleeding, and the availability of local bariatric surgical expertise. The use of less invasive options such as gastric balloons is potentially attractive, as this will keep the additional risks to a minimum. The intra-gastric balloon (IGB) is a safe, endoscopically placed, saline-filled, silicone balloon which can remain in the stomach for up to 6 months (94). After 6 months, the risk of spontaneous deflation with the possibility of distal migration from the stomach increases. As such, centres offering the IGB schedule to remove it after 6 months. There are good efficacy data in non-transplant patients, such that some bariatric services use it as a first line procedure and progress only to formal bariatric surgery if the IGB fails. The weight loss achievable by the IGB has been shown to be up to 32 kg after 6 months. After IGB insertion, patients require a minimal (overnight stay) in hospital but require maintenance on proton pump inhibitors as long as it remains in place.

Portal hypertension is a contra-indication to IGB placement and may limit its utility in early post transplant patients. The rate of spontaneous deflation of the IGB is very low at less than 0.1%. In the unlikely event that the IGB does deflate unexpectedly, methylene blue dye admixed with the saline in the balloon will discoulour the stool and so provide prompt evidence of the leakage of balloon contents. As all liver transplant units have 24-hour on call endoscopic services, emergency IGB removal can be done by highly competent, appropriately trained staff. Nausea and vomiting are other potential complications of the IGB and most patients are maintained at least initially on anti-emetics and anti-spasmodics.

**Recommendation 18:** Consider bariatric surgery at the time of liver transplantation in recipients with severe morbid obesity, those with failed treatment of obesity, or in patients with recurrent disease undergoing retransplantation. (Level C, Class IIa)
**Bariatric surgery after liver transplantation**

The liver transplant recipient who remains morbidly obese or develops recurrent or de novo NAFLD may in the course of time be considered a candidate for bariatric surgery. Little or no data exist on the timing and selection of such patients for bariatric surgery. The vast majority of these patients will be on modest stable maintenance immunosuppression, and ideally would be off corticosteroids or on low dose maintenance corticosteroids. Most such patients are on a combination of an antiproliferative agent (azathioprine or mycophenolate) and a CNI (tacrolimus or ciclosporin). Patients on mTOR inhibitors (sirolimus, everolimus) will need to stop these drugs as they are associated with impaired wound healing, and alternative regimes considered.

There are few data on which to base recommendations for patient and procedure selection for patients undergoing bariatric surgery post liver transplant. Extensive adhesions may make the laparoscopic approach difficult or sometime impossible, with a higher risk of open conversion and its additional risks. The use of intestinal bypass surgery may have an impact on future biliary enteric surgery and may also affect the absorption of immunosuppressive medication. In view of these issues, it is preferable to consider the use of endoscopic placed gastric balloons as the procedure of choice, followed by non-bypass gastric surgery as the next option. Case reports support the feasibility and efficacy of Roux-en-Y gastric bypass for recurrent NASH in liver transplant recipients with morbid obesity (95).

**Recommendation 19:** Consider bariatric surgery in recipients with severe morbid obesity, those with failed treatment of obesity, or in patients with developing progressive NASH with fibrosis in allograft. (Level C, Class IIa)
10 Peri-operative monitoring

The anhepatic and reperfusion phases during liver transplantation are associated with a high risk of cardiovascular instability. The former occurs due to a sudden reduction in venous return, and the latter because of the effects of the return of cold, acidic and hyperkalaemic preservation fluid into the circulation (96). Release of vasoactive mediators and myocardial depression related to free oxygen radical production are additional pathogenetic mechanisms. The clinical picture is characterised by sudden onset of profound arterial hypotension due to the combined effects of reduction in systemic vascular resistance, increase in pulmonary vascular resistance, and drop in heart rate and cardiac output.

Patients with NAFLD and associated cardiovascular risk factors may be particularly vulnerable to these effects, and advanced haemodynamic monitoring with pulmonary artery catheters (PAC) is routinely performed in many transplant centres. Despite the theoretical advantage of invasive peri-operative monitoring and therapeutic guidance based on PAC data, randomised controlled trials in high risk surgery (97) or in the critically ill (98) have not shown any evidence of benefit. Indeed, there are no controlled data showing an advantage of PAC use, or indeed any other form of advanced haemodynamic monitoring tool during liver transplantation.

Trans-oesophageal echocardiography (TOE) provides immediate and direct assessment of cardiac structures and the functional status of both ventricles. It is an excellent tool to diagnose new onset regional wall motion abnormalities (RWMA) and enables assessment of the filling status of the left ventricle as well as estimation of pulmonary artery pressures. Compared to PAC it is relatively non-invasive and low risk; however, it is operator dependent and requires appropriate training. Whether intra-operative management based on TOE imaging provides any outcome benefit is unknown. The lack of continuous monitoring capability outside the operating theatre in the Intensive Care Unit (ICU) is another potential drawback.

**Recommendation 20:** Whilst there is likely to be an increased operative risk, the lack of evidence from controlled clinical trials indicates that no recommendation can be made about the use of intra-operative cardiac output monitoring. (Level B, Class II)
Peri-operative management of conditions frequently associated with NAFLD

Insulin resistance, diabetes and blood sugar control
A linear relationship exists between blood sugar levels and adverse outcome in hospitalised patients, regardless of the presence of diabetes. A large single centre randomised controlled study published in 2001 showed a 42% relative mortality reduction in surgical ICU patients when blood sugar levels were normalised to 4.4 to 6.1 mmol/l (99). This was achieved by virtue of continuous insulin infusion in combination with hypercaloric parenteral nutrition. The control group only received therapy if glucose level exceeded 11.9 mmol/l. A minority of patients had a history of diabetes (13%) and only 5% of patients were admitted following transplantation, which makes extrapolation of the findings to the post liver transplant setting difficult. Subsequent multicentre studies and a meta-analysis failed to reproduce the results of this trial, and concerns have been raised in regards to the possible harmful effects of tight glucose control through an increased incidence of life threatening hypoglycaemia.

The largest study to date from the ANZICS collaborators (NICE – SUGAR), a multicentre RCT, randomised more than 6,000 patients to tight glucose control of 4.5 to 6 mmol/l vs 10 mmol/l or less. Surprisingly, intensive insulin therapy increased mortality in both medical and surgical patients, perhaps in association with an increased incidence of severe hypoglycaemia (100). In comparison with the earlier trial, patients were primarily enterally fed and they received significantly less calories and insulin; however, the blood sugar level in the control group was comparable, averaging 8 mmol/l. In the absence of evidence in relation to peri-operative blood sugar management in NAFLD and extrapolating from the available data pertaining to ICU patients, moderate glucose control targeting levels between 6 and 10 mmol/l is recommended.

**Recommendation 21**: Moderately tight glucose control (6-10 mmol/l) should be targeted during the early post transplant course in patients of all aetiologies. (Level A, Class II)

NASH and statin use in the peri-operative period

Dyslipidaemia is a common finding in patients with NASH and statins are widely used for this indication. Recent studies have extended the use of statins beyond traditional indications of lipid lowering therapy. Cholesterol-independent therapeutic, pleiotropic effects of statins include anti-
inflammatory and anti-oxidative properties, improvement of endothelial function, and increased nitric oxide bioavailability. In addition to anti-inflammatory properties, statins may interfere with the activation of the coagulation cascade and modulate platelet function. Several retrospective trials and prospective studies in patients undergoing vascular surgery have shown a beneficial effect of statins, with reduction in the incidence of peri-operative myocardial infarction, coupled with an excellent safety profile. Several experimental and observational trials postulate a beneficial role of statins in sepsis, especially in patients with suspected bacterial infection (101). However, these findings have to be confirmed in adequately powered placebo controlled trials before a universal recommendation for statin use in sepsis can be made.

Patients already receiving statins should not have their treatment interrupted peri-operatively or during the intensive care stay to maintain the beneficial pleiotropic effects and reduce the risk of cardiac events (102). As statins can interact with CNIs, careful attention should be paid to the choice and dose of statin used.

**Recommendation 22:** If started preoperatively, statin therapy should be continued during the post-operative phase. (Level B, Class I)
11 Immunosuppression

Life-long immunosuppression (IMS) is required in the great majority of patients following liver transplantation, irrespective of primary indication. Overall, liver allograft recipients have an increased risk of morbidity and mortality from cardiac and cerebrovascular disease, weight gain and diabetes (103). Maintenance IMS regimes are based on either monotherapy with a calcineurin inhibitor (CNI) (ciclosporin or tacrolimus) or an mTOR inhibitor (mTORi) (sirolimus or everolimus); or combination therapy with two or more of corticosteroids, CNI, mTORi and an antimetabolite (usually azathioprine or mycophenolate). The selection of the most appropriate regime for the patient is likely to vary during the course of the lifetime of the graft and will depend on many factors, including the indication, likelihood and consequences of allograft rejection, and co-morbid conditions (104).

Although both recurrent and de novo NAFLD may develop in the allograft, there have been no significant prospective trials evaluating different IMS regimes in patients grafted for NASH. The selection of the most appropriate IMS regime for the liver allograft recipient transplanted for NASH will depend on many factors, including:

- The time after transplant. In many cases, immunosuppression load can be reduced; many centres withdraw steroids by 3 months
- Risk of rejection. This will depend on many factors, including the indication for transplantation, a history of early or late rejection, and the response to increased immunosuppression
- Co-morbid conditions. For example, CNI may be reduced or avoided where there is renal impairment
- Indication. mTORi may be preferred for those with liver cell cancer
- Complications. CNIs may be reduced or avoided when renal impairment develops, and mycophenolate should be avoided when neutropenia develops

Impact of immunosuppressive agents on risk factors for NAFLD

The currently used agents all have differing impacts upon components of the metabolic syndrome, as shown in Table 11.1 (50,103-111).
Table 11.1  Impact of immunosuppressive agents on components of the metabolic syndrome and hepatic steatosis

<table>
<thead>
<tr>
<th>Class of agent</th>
<th>Hypertension</th>
<th>Dyslipidaemia</th>
<th>Diabetes</th>
<th>Weight gain</th>
<th>Hepatic steatosis</th>
</tr>
</thead>
<tbody>
<tr>
<td>Corticosteroids</td>
<td>+</td>
<td>+</td>
<td>++</td>
<td>++</td>
<td>++</td>
</tr>
<tr>
<td>CNI</td>
<td>++</td>
<td>++</td>
<td>++</td>
<td>+</td>
<td>-</td>
</tr>
<tr>
<td>Antimetabolites</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>mTORi</td>
<td>-</td>
<td>+++</td>
<td>+/-</td>
<td>-</td>
<td>-</td>
</tr>
</tbody>
</table>

Ciclosporin and tacrolimus have slightly differing effects on some of these risks: for example, NODAT (new onset diabetes after transplantation) is more common with tacrolimus than ciclosporin (105), while hyperlipidemia is more common with ciclosporin (104). Due to superior patient outcome, tacrolimus is preferred to ciclosporin and is the first line CNI used in clinical practice (112).

Selection of an immunosuppressive regime

The prime goal of any immunosuppression regime must be the maintenance of normal graft function and the prevention of graft damage from immune and other factors, with an acceptable side-effect profile. However, the selection of specific combinations of immunosuppressive agents may allow a reduction of the risk of recurrence of NAFLD in the graft and the associated vascular and metabolic side-effects.

While there are no prospective studies evaluating different immunosuppressive regimes in this context, the following may be considered for those transplanted with or for the metabolic syndrome and/or NAFLD:

- Corticosteroids. Where possible, use either a steroid free regime or early steroid withdrawal. Where steroid free regimes are used, induction therapy (such as anti-thymocyte globulin or IL2-R antagonism) should be considered.
- CNIs. Although tacrolimus may be more strongly associated with diabetes than ciclosporin, it has a better effect on graft outcome.
• mTOR inhibitors. Should be avoided as first line agents, although they may be indicated for those with liver cell cancer or as rescue therapy.

Several studies support use of anti-metabolites such as mycophenolate to allow lower doses of tacrolimus and reduce the impact on diabetes and renal dysfunction (113-115).

**Recommendation 23:** Alongside immunosuppression with CNI and an anti-metabolite, consideration should be given to either a steroid free regime or early steroid withdrawal (within three months) in patients with NASH. Where steroid-free regimes are used, induction therapy (such as ATG or IL2-R antagonism) should be considered. (Level B, Class IIa)

**Recommendation 24:** Tacrolimus levels should be <10 ng/ml within the first three months after liver transplantation and 5-8 ng/ml after that to reduce the impact on renal function and dyslipidaemia. Mycophenolate should be used as the preferred anti-metabolite, to permit lower levels of tacrolimus. (Level B, Class IIa)

It is essential that modification of the IMS regime should be considered in combination with lifestyle and pharmacological approaches to reducing the recurrence and impact of metabolic syndrome. This includes regular monitoring and treatment (where indicated) of diabetes, hypertension and hyperlipidaemia.

**Recommendation 25:** Close follow-up and early recognition and treatment of the recognised consequences of transplantation and immunosuppression (such as weight gain, hypertension, hyperlipidaemia, diabetes and renal impairment) remain the key to preventing excess risk from recurrent NAFLD. (Level C, Class I)
12 Post-transplant monitoring of NASH patients and disease recurrence

NAFLD recurrence in the liver allograft

Many studies have described histological features of recurrent fatty liver disease in patients transplanted for NASH (26,116-123). Similar changes have also been observed in patients transplanted for conditions other than NASH (118,119,124-129). Some of these patients were initially diagnosed as having cryptogenic cirrhosis at the time of listing but were subsequently found to have pre-transplant risk factors for fatty liver disease, whilst others presumably reflect transplant-related risk factors that predispose to the metabolic syndrome.

Recommendation 26: Histological examination of the explanted liver should be carried out to confirm the presence of features compatible with end-stage NASH and to exclude features suggesting an alternative diagnosis. (Class I, Level C)

Histological assessment of NAFLD recurrence

The histological features of NAFLD in the liver allograft are essentially the same as those occurring in the native liver. However, changes seen in post-transplant biopsies may be modified by a number of other complications of liver transplantation. The challenge to the pathologist is to distinguish whether all features are attributable to fatty liver disease, or whether there is an additional post-transplant complication. Conditions that may have overlapping histological features with certain components of NAFLD include rejection, biliary complications and vascular/architectural changes including nodular regenerative hyperplasia (130-132).

Late rejection may lack the typical portal tract features of acute rejection and frequently includes centrilobular necro-inflammatory changes (‘central perivenulitis’) that may be associated with hepatocyte ballooning and with the development of centrilobular fibrosis. Biliary complications are characterised by periportal ductular reaction and fibrosis, both of which have been described as features of NAFLD in the native liver (133). Nodular regenerative hyperplasia is an increasingly recognised finding in late post-transplant biopsies and may be associated with the development of perisinusoidal fibrosis in centrilobular regions (134). The latter sometimes occurs as an isolated finding without conspicuous nodularity. There are also interactions between HCV infection, insulin
resistance and metabolic fatty liver disease, which may be important in the pathogenesis and severity of recurrent HCV infection and de novo NAFLD (127,135-138). In most cases where a dual pathology is suspected, careful histological assessment and clinico-pathological correlation should enable the main cause of graft damage to be identified. For this reason, biopsies performed elsewhere should be reviewed at the transplant centre.

Semi-quantitative scoring systems are increasingly used to assess the severity of NAFLD in the native liver (139). The most widely used systems are those described by Brunt (140) and Kleiner (141), both of which grade the severity of steatosis, hepatocyte ballooning and lobular inflammation, and stage the severity of fibrosis. Although several studies have assessed the histological severity of NAFLD in the liver allograft, the clinical utility of formal scoring systems in this context has not been validated. Problems may arise if there are co-existent graft complications that can mimic some of the NAFLD-related features that are used for histological grading or staging. Scoring should therefore only be carried out if the changes seen can be attributed to NAFLD alone.

**Recommendation 27:** The main role of biopsy is to allow diagnosis and staging of liver histopathology. Where NAFLD is the only or dominant pathology, liver allograft biopsies can be scored using the Kleiner classification. Biopsies performed elsewhere should be reviewed at the transplant centre to ensure reproducibility. (Class I, Level C)

**Frequency and impact of NAFLD recurrence**

Fatty change occurs in 60-100% of patients within a few months of transplantation, and approximately 10-40% develop features of steatohepatitis, albeit usually not severe. Approximately 10% of patients progress to bridging fibrosis or cirrhosis by 10 years post-transplant (28). Likewise, features of fatty liver disease are commonly seen in protocol biopsies from patients transplanted for conditions other than NASH (129,142,143). One third of patients with histological evidence of recurrent NASH have normal liver enzymes, which supports the role of protocol liver biopsies in monitoring disease progression in the liver allograft (121). Whilst disease progression is rapid in occasional cases, resulting in graft failure within 2-3 years of transplantation (117), in the majority of cases recurrent NAFLD does not appear to have a major impact on graft survival in the first 5 years post-transplant (122,144,145). However, the long term
clinical importance of recurrent NASH is not yet known, and data from the pre-transplant setting indicate that NASH confers a worse prognosis to that of simple steatosis over a 10-15 year period. Distinction between NASH and steatosis in the allograft should therefore result in a focus on changes in lifestyle advice and medical therapy.

**Recommendation 28**: Post-transplant monitoring of patients should include an initial USS at one year, followed by every two years, looking for the presence of an echobright liver. (Class IIa, Level C)

**Recommendation 29**: Post-transplant monitoring of patients with echobright liver on USS should include protocol liver biopsies to detect disease recurrence, as liver function tests may be normal. Repeat biopsy should be considered every three years, unless there is a clinical indication for more frequent biopsies. (Class IIa, Level C)

**Factors predicting NAFLD recurrence**

Cumulative steroid dose after LT has been shown to predict recurrence of NASH in the allograft (26), but further work is required to identify whether other factors such as BMI, dyslipidaemia and diabetic status/control are also responsible. Use of ACE inhibitors has been suggested to reduce recurrence post-transplant (127).

**Research/audit recommendation**: It will be important to compare non invasive modalities for the assessment of NASH and fibrosis with liver biopsy post-LT.
13 Post-transplant management of NAFLD

Obesity and weight management

Patients transplanted for NASH are at an increased risk for the subsequent development of features of the metabolic syndrome. New onset obesity is reported in 17-43% of post transplant patients (123,146-148). In addition, hypertension is reported in 35%, IGT/DM 16%, hypercholesterolaemia 18%, and hypertriglyceridaemia in 19% of patients (123). Overt post-transplant metabolic syndrome comprising a constellation of cardiovascular risk factors may be present in up to 58% of patients (104,149,150). Importantly, 26% of deaths in post-transplant NAFLD patients are due to cardiovascular causes (123) and those patients with post-transplant metabolic syndrome have significantly more cardiovascular events than those without metabolic syndrome (30% vs 8%)(149).

Morbidly obese patients (BMI >40 kg/m$^2$) have a higher incidence of primary graft non-function post-transplant, but this does not appear to impact upon 1-2 year graft survival. However, 30-day, 1-year and 2-year mortality are significantly higher in morbidly obese patients. 5 year mortality is increased in both severe (BMI >35 kg/m$^2$) and morbid obesity, but this is not a reflection of intraoperative mortality. Using a proportional hazards model of survival, 2-year survival is independently predicted by morbid obesity and type 2 diabetes (72).

 Whilst it makes clinical sense to optimise weight management and reduce cardiovascular risk, there are currently no published studies that have examined the impact of weight reduction on outcome in patients transplanted for NAFLD. In a single study, >90 minutes of exercise per week was associated with improved quality of life outcomes as measured by the SF-36 questionnaire in post-liver transplant patients, although the precise indication for transplantation was not defined in this cohort (151).

**Recommendation 30:** Post-transplant patients should receive support, advice and treatment in order achieve a target body mass index of <25 kg/m$^2$. This should be in the context of a multidisciplinary team, incorporating dietary modification, exercise intervention and the potential use of pharmacotherapy and surgical intervention where appropriate. (Level C, Class I)
**Insulin resistance and type 2 diabetes**

Studies have reported a variable prevalence of post-transplant diabetes and although some of the variability may be explained by different diagnostic criteria, values range from 13% to 61% (149,152-154). The prevalence of diabetes in the post-transplant group is 5.99 times higher (95% CI 4.15-8.38) than in age and sex matched control populations (155). Insulin resistance has been documented in almost all patients with pre-transplant cirrhosis (156,157). After transplantation, insulin sensitivity improves (156-158), but importantly many patients (in excess of 40%) remain insulin resistant (104).

Patients with diabetes have increased mortality post transplant in comparison with non-diabetic patients (33). Whilst mortality in patients whose diabetes is controlled by diet alone is not increased in comparison with non-diabetic post transplant patients, those patients on oral agents and/or insulin therapy have increased mortality (159). More detailed outcomes studies are urgently needed and there are currently no published data that have examined the impact of glycaemic control on outcome in patients transplanted for NASH.

**Recommendation 31:** Post-transplant patients should be screened for the presence of diabetes and, if present, reviewed regularly for the development of complications. Glycaemic control should be optimised in accordance with NICE guidance. (Level A, Class I)

**Cardiovascular risk**

There is a high incidence of cardiovascular events in liver transplant recipients, with an increased standardised risk compared with a non-transplant population (40). 26% of deaths in post transplant NAFLD patients are due to cardiovascular causes (123). Patients with post transplant metabolic syndrome have significantly more cardiovascular events than those without metabolic syndrome (30% vs 8%) (149).

**Recommendation 32:** Patients transplanted for NAFLD should be closely monitored on a 6-monthly basis for risk factors for cardiovascular disease (BP, lipids, HbA1c), which should be addressed with the intention of reducing cardiovascular events (Level II-3, Level C).
Hypertension

The prevalence of hypertension after OLT is significantly higher than that in the general population (standard prevalence ratio 3.07, 95% confidence interval 2.35-3.93)(155). While accepted JNC-7 definitions of hypertension apply, i.e. BP ≥140/90 mmHg, patients with type 2 diabetes, common in this population, and/or renal dysfunction should have a target BP <130/80 mmHg.

Non-pharmacological measures including weight reduction, salt restriction, physical activity and alcohol restriction are recommended despite the absence of specific trials. The aim of pharmacological therapy is to achieve the target BP and in the setting of liver transplantation there is evidence that calcium channel blockers, β-blockers and ACE inhibitors are efficacious in this respect (160-163). Specific agents may also confer additional benefits, such as reduction of proteinuria (164) and prevention of liver fibrosis (165) with ACE inhibitors, although these are not yet proven (166).

Recommendation 33: A blood pressure target of 140/90 mmHg (130/80 mmHg in patients with diabetes mellitus and/or renal dysfunction) should be aimed for (Level A, Class I).

Recommendation 34: Anti-hypertensive agents such as calcium channel blockers or ACE inhibitors should be considered in view of their possible additional effect of abrogating liver fibrosis. (Level C, Class II)

Dyslipidaemia

The prevalence of post-OLT dyslipidaemia ranges from 66% to 85%, usually with a mixed picture of elevated total cholesterol and triglycerides (146,148,167). Although standard treatment thresholds apply, a target LDL cholesterol of <2.6 mmol/l is advised as the 10-year cardiovascular event rate exceeds 20% for the liver transplant population (167). Therapeutic lifestyle changes are recommended, although not evidence based. Statins have been shown to be efficacious (168,169); however, consideration should be given to drug interactions (e.g. ciclosporin and calcium channel blockers) which can be minimised with the use of pravastatin (103). Ezetimibe has been demonstrated to be effective at reducing serum cholesterol and to have few side effects.
and no interaction with immunosuppressive regimens (170). However, long-term outcome studies are not yet available for this agent.

**Recommendation 35:** A target LDL cholesterol of <2.6 mmol/l is advised as the 10-year cardiovascular event rate exceeds 20% for the liver transplant population. Pravastatin and ezetimibe are preferred agents in view of their demonstrated efficacy and absence of interactions with CNIs (Level C, Class IIa).
14 Non-alcoholic fatty liver disease in children

NAFLD is now the commonest liver disease in childhood, reflecting the global increase in childhood obesity (171,172). NAFLD currently affects approximately 10% of adolescents and up to 40% of obese adolescents (173). The majority of children who have undergone biopsy for suspected NAFLD have some degree of hepatic fibrosis. This is usually mild, but risk factors for increased fibrosis include age, the degree of obesity, male sex, ethnicity and hypopituitarism (171,174).

Increased carotid artery intimal thickness occurs in children with NAFLD, and they have also been shown to have increased mortality (175,176). It is unclear at present whether the presence of NAFLD is itself an independent risk factor for cardiovascular disease in obese children.

There are four published cases of paediatric transplantation for NAFLD (176-178). In two cases the primary aetiology was hypopituitarism, and all four cases had hepatopulmonary syndrome as the immediate indication for transplantation. All survived, but recurrence of NAFLD with significant fibrosis was universal within the first year. In one case this required repeat transplantation after 3 years. Another case was treated with metformin and initial moderate fibrosis resolved 3 years post transplant.

**Pre-transplant assessment**
Investigations should include routine pre-transplant assessment, but in addition should focus on components of the metabolic syndrome including fasting lipid profile, hyperuricaemia, hypertension and evaluation of pituitary function. Specific screening for hepatopulmonary syndrome including, as a minimum, saturation monitoring is indicated. At present there is insufficient evidence to justify formal invasive screening for cardiovascular disease.

**Indications for transplantation**
These are similar to those for end-stage liver disease of any cause.

**Management post transplantation**
No specific evidence guidelines can be recommended; however, in the absence of autoimmune disease, steroid administration should be minimised. It would appear sensible to minimise calcineurin inhibition by the use of anti-metabolites such as mycophenolate or azathioprine.
Monitoring
Evidence based recommendations cannot be given. Good practice would suggest active monitoring for components of the metabolic syndrome and protocol liver biopsies at one, five and ten years post transplant.

De novo post transplant NAFLD
This may well turn out to be a quantitatively more important issue than NAFLD as a primary indication for liver transplantation. There is a high incidence of obesity following transplantation. However, current long term studies suggest that although fibrosis is a common finding in late post-transplant biopsies, this does not appear to be related to the development of fatty liver disease (179,180).
15 Transplantation for NASH – the patients’ perspective

Improving awareness and reducing stigma

Patient feedback suggests that opportunities for early intervention and diagnosis in patients with key risk factors for NAFLD and NASH, such as obesity, diabetes and insulin resistance, are being missed, leading to unnecessary disease progression. Many patients also report a degree of stigma associated with liver disease in the public’s perception. This can be very difficult for patients and families to deal with and can make them feel very isolated. Patients, and in particular family members, say they would have liked to have been forewarned about the symptoms of advanced liver disease and not just been given information on the transplant process. Many mention hepatic encephalopathy as being particularly distressing and feel that understanding the cause would have reduced this distress.

**Recommendation 36:** There is a need to increase understanding of liver disease and its many causes, to improve patient outcomes and reduce the stigma many patients experience. (Not Graded)

Patients diagnosed with NASH, particularly those who do not have obvious symptoms of the metabolic syndrome, report confusion or lack of information on their diagnosis, its cause, and what they can do to manage and improve their condition. With appropriate advice, management and support patients can make lifestyle changes, improve liver health and avoid transplantation. Many patients struggle to maintain lifestyle changes in the long-term. Many receive ongoing monitoring for clinical symptoms but no support for maintaining lifestyle changes.

Some patients believe they will be a ‘different person’ post-transplant and can find it difficult to come to terms with ongoing health problems. Those who suffered from depression, stress or eating disorders prior to transplant can find themselves falling back into old patterns.

It is important that patients understand the cardiovascular risks associated with both NASH and LT and what they can do to reduce these. Family members have a key role in supporting patients pre- and post-transplant, particularly with understanding and managing diet and being aware of symptoms. It is important that they are given information and involved in decisions.
**Recommendation 37:** The potential cardiovascular morbidity associated with NASH should be discussed with patients and guidance given on diet and exercise, and sources of support (including psychological support) as part of ongoing management. (Not Graded)

**Psychological impact**

The emotional and psychological impacts of the transplant process are significant and can be particularly difficult for people without good support from family and friends, or for those who have suffered from depression before transplantation. Patients can find it helpful to be put in touch with people who have previously been through the transplant experience, often as part of support groups. People who receive a transplant say they struggle to put into words the gratitude they feel towards the transplant team and the donor family. They therefore find it difficult to raise any complaints directly with the transplant team. Some patients (and their family members) have difficulty post-transplant in coming to terms with a transplanted organ. Patients also report experiencing flashbacks to their time in intensive care and say that they find it difficult to discuss these feelings with the transplant teams. The need for confidential emotional and psychological support is often raised by patients and carers.

**Recommendation 38:** Provision of independent pre- and post-transplant emotional and psychological counselling and support is very important, along with an opportunity to provide confidential feedback to the transplant team post-operatively. (Not Graded)

This patient feedback was collated from British Liver Trust helpline enquiries, case studies, contacts and transplant support groups.
16 Statements of potential conflicts of interest

Dr M Allison
I have received travel and meeting expenses, but have not received lecture fees or contributed to Advisory Boards from Astellas, Novartis, Pfizer and Roche. I have received no support from other agencies with a known interest in the development of these guidelines.

Dr PA Andrews
I have received travel and meeting expenses, but have not received lecture fees or contributed to Advisory Boards from Astellas, Novartis, Pfizer and Roche. I have received no support from other agencies with a known interest in the development of these guidelines.

Dr G Auzinger
I have received lecture fees from Pfizer (Anti-Infectives Division) and contributed to Advisory Boards from Gilead, MSD and Pfizer. I have received no support from other agencies with a known interest in the development of these guidelines.

Prof CP Day
I have received travel and meeting expenses from Astellas, Novartis and Roche and have been on the Advisory Boards for NASH of Astellas, GSK, Pfizer and Synageva.

Dr JW Ferguson
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Dr P Henriksen. I have received no support from agencies with a known interest in the development of these guidelines. I have no conflicts of interest for this document.

Prof SG Hubscher
I have received no support from agencies with a known interest in the development of these guidelines. I have no conflicts of interest for this document.
Ms H Manley
I have received no support from agencies with a known interest in the development of these guidelines. The British Liver Trust has received support from Astellas and Roche in the form of educational grants to enable the Trust to carry out its mission and activities.

Dr PJ McKiernan
I have received no support from agencies with a known interest in the development of these guidelines. I have no conflicts of interest for this document.

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I have received no support from agencies with a known interest in the development of these guidelines. I have no conflicts of interest for this document.

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Prof JM Neuberger
I have received speaker support from Astellas and Novartis and was a Principal Investigator for a Roche funded clinical trial. I have received no support from other agencies with a known interest in the development of these guidelines.

Dr PN Newsome
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Dr J Oben
I have received no support from agencies with a known interest in the development of these guidelines. I have no conflicts of interest for this document.
Mr S Pollard
I have received travel and meeting expenses from Astellas, Novartis and Roche over the past 18 years and was on the Advisory Board of Novartis until 2002. I have received no support from other agencies with a known interest in the development of these guidelines.

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I have received no support from agencies with a known interest in the development of these guidelines.

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I have received no support from agencies with a known interest in the development of these guidelines. I have no conflicts of interest for this document.
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