

Adult liver transplantation: A UK clinical guideline - part 1: preoperation

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ABSTRACT

REVIEW

Liver transplantation is a highly successful treatment for all types of liver failure, some non-liver failure indications and liver cancer. Most referrals come from secondary care. This first part of a two-part guideline outlines who to refer, and how that referral should be made, including patient details and additional issues such as those relevant to alcohol and drug misuse. The process of liver transplant assessment involves the confirmation of the diagnosis and non-reversibility, an evaluation of comorbidities and exclusion of contraindications. Finally, those making it onto the waiting list require monitoring and optimising. Underpinning this process is a need for good communication between patient, their carers, secondary care and the liver transplant service, synchronised by the transplant coordinator. Managing expectation and balancing the uncertainty of organ availability against the inevitable progression of underlying liver disease requires sensitivity and honesty from all healthcare providers and the assessment of palliative care needs is an integral part of this process.

INTRODUCTION

Over just three decades, UK liver transplantation has evolved from the enthusiastic efforts of a few well-intentioned clinicians to a multidisciplinary, closely scrutinised therapy with 1-year survival rates in excess of 90%.^{1–3} Despite these excellent outcomes, only a small fraction of the increasing numbers of patients dying of end-stage liver disease will be referred to a liver transplant unit (LTU) for this life-saving procedure. This two-part guideline is specifically aimed at non-specialist clinicians caring for patients with acute and chronic liver disease (CLD). The first part examines:

- Who to refer for liver transplant (LT).
- ► How to refer for LT.
- The LT Assessment.
- How to manage the patient on the waiting list.

Part 2 explores the post operative care of the LT recipient.

Further reading includes guidelines from BSG (1999),⁴ BASL (2012),⁵ EASL Guidelines for LT (2015)⁶ and acute liver failure (ALF)⁷ and AASLD guideline for LT (2013).⁸

WHO TO REFER FOR LT

Over 90% of LTs in the UK are performed for CLD, where a gradual destruction of liver tissue results in the familiar picture of jaundice, ascites, encephalopathy with coagulopathy and hypoalbuminaemia. A smaller number will have ALF^{3 7} and an even smaller number are transplanted for a non-failing liver, where there is survival advantage.⁹ This section identifies the reasons, which should prompt either a referral to an LTU or mandate an enquiry.

Acute liver failure

While there are several ALF definitions,⁷ the critical elements for the purpose of this guideline, are the three cardinal features of encephalopathy, jaundice and coagulopathy appearing in a patient who, less than 6 months ago, had no evidence of advanced liver disease. The challenge for the generalist in an AMU or emergency department

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Table 1 When referral/discussion with the LTU is required for	r a patient with ALF ⁷
Paracetamol induced acute liver failure	Non-paracetamol induced acute liver failure
 Arterial pH <7.30 or HCO₃ <18 INR >3.0 on day two or >4.0 thereafter Oliguria and/or AKI Altered level of consciousness Hypoglycaemia Elevated arterial lactate (>4 mmol/L) unresponsive to fluid resuscitation 	 pH <7.30 or HCO₃ <18mmol/I INR >1.8 Oliguria/renal failure or Na <130 mmol/L Encephalopathy, hypoglycaemia or metabolic acidosis Bilirubin >300 umol/L (17.6 mg/dL)

AKI, acute kidney injury; ALF, acute liver failure; CLF, chronic liver failure; INR, international normalised ratio; LT, liver transplantation; LTU, liver transplant __unit.

environment is establishing the diagnosis of severe liver injury quickly and minimising the delay in seeking help.¹⁰ Early discussions with LTU enable decisions on comorbidities or contraindications to LT (see below) to be addressed before hepatic encephalopathy develops and safe transfer becomes too high-risk.

Causes of ALF

Despite changes to packaging, paracetamol (acetaminophen) poisoning remains the the most common cause for ALF in the UK.¹¹ The next most common cause is non-A-to-E hepatitis, then other drug induced liver injuries (prescribed, herbal and proscribed), viral hepatitis and ischaemic hepatitis. Malignancy (primary or secondary), pregnancy (AFLP/HELLP), vascular (including Budd-Chiari Syndrome) and metabolic disorders are rarer causes.

Alcoholic hepatitis is considered in the CLD section, as is acute on chronic liver failure (AoCLF).

When to consider referral/discussion with the LTU in ALF

Once diagnosed, the ALF patient should be managed in an HDU environment and discussed with a LTU (see table 1). Important details include any history of paracetamol ingestion (timing, frequency, 'staggered'), pregnancy, other drugs (prescribed, herbal or proscribed), comorbidity (mental and physical health), laboratory results (including PT, pH, arterial lactate, glucose, renal function, viral screen, autoantibodies and immunoglobulins) and liver imaging. Ideally, patients are safer transferred *before* encephalopathy appears, although there is little published guidance covering this important issue.

Table 1 outlines clinical features in paracetamol and non-paracetamol ALF that correlate with poor outcome and mandate referral. Rarely, non-paracetamol ALF, can present with ascites, deep jaundice and even variceal haemorrhage, where the short history remains the only clue to ALF.

Chronic liver failure

CLF occurs on the background of established liver cirrhosis. The typical clinical features include jaundice, ascites, encephalopathy, sarcopenia along with laboratory features, such as hypoalbuminaemia and coagulopathy, often associated with a rising creatinine and hyponatraemia as liver disease advances. Box 1 describes common causes of CLF.¹²

Acute on chronic liver failure

Acute-on-chronic liver failure (AoCLF) is a syndrome characterised by acute decompensation of CLD associated with organ failures and high short-term mortality. Sepsis, active alcoholism and relapse of chronic viral hepatitis are the most common reported precipitating factors, but still only account for perhaps half the cases, the remainder have no identifiable trigger. The poor prognosis mimics that seen in ALF and mandates an expedited triage and consideration for LT. However, while LT remains the definitive treatment, sadly very few prove suitable.¹³

When to consider referral/discussion with the LTU in CLF or AoCLF

Liver transplantation should be considered, when a patient with established liver disease develops any of the typical features of decompensation (figure 1). As decompensation correlates with rising morbidity and mortality, the supervising clinician should reflect on the key considerations outlined below.

Box 1 Causes of chronic liver failure (CLF)

Causes of CLF:

- Alcohol.*
- ► Non-alcoholic fatty liver disease.
- Chronic viral hepatitis (B, C and D).
- Autoimmune liver disease (Primary Biliary Cholangitis, Primary Sclerosing Cholangitis, Autoimmune Hepatitis and overlap syndromes).
- Wilson's disease.
- Genetic haemochromatosis.
- Alpha-1 antitrypsin deficiency.
- Secondary sclerosing cholangitis.
- Congenital hepatic fibrosis and other congenital or hereditary liver diseases.

*Alcoholic hepatitis tends to present acutely, frequently with no history of liver disease. Most patients recover with abstinence, but recent studies have demonstrated excellent outcomes with liver transplant in highly selected patients.^{44 45}

Jaundice Jaundice Ascites Variceal haemorrhage Hepatic encephalopathy

Figure 1 Clinical features of hepatic decompensation.

Key considerations in patients with hepatic decompensation

- 1. Is the decompensation potentially reversible (for example with abstinence in the case of ALD or with anti-virals in untreated chronic viral hepatitis)* or
- 2. If not reversible, is the patient suitable for LT*? or
- 3. Are there any contraindications to LT such as comorbidity which preclude transplant* (see table 5) or
- 4. If not currently suitable for transplant, could a patient become suitable with treatment or an intervention*?

*If in any doubt, seek advice from local LTU

Bear in mind that it is possible for a patient to be too unwell for LT *if* the referral is made too late.

To aid with the assessment of suitability for referral, the UK Model for End-Stage Liver Disease (UKELD) score can be calculated. The UKELD score is devised from patient's INR, serum sodium, creatinine and bilirubin (https://www.odt.nhs.uk/transplantation/ tools-policies-and-guidance/calculators/).¹⁴ UKELD scores \geq 49 indicate survival advantage for LT over conservative management in patients with irreversible decompensation. Patients with decompensated CLD, unsuitable for LT, should have this conclusion documented and receive symptom-directed care.

Non-liver failure indications for liver transplantation

Some patients will still benefit from LT, even though their liver is not failing (i.e, likely to cause death within 12 months). This would include patients with cirrhosis and a UKELD score under 49, such as PBC patients with intractable pruritus (see table 2).^{5 6 8 15}

HOW TO REFER FOR LT

The next step is referral to the nearest LTU (see online supplementary appendix 1). For ALF patients, a telephone referral is obligatory, but for all other indications, a written referral suffices. Email may speed the referral process, but correspondence with a named individual in LTU encourages collaborative dialogue between referring and transplant physicians. Most referrals for CLD patients come from secondary care hepatologists or gastroenterologists, but referrals are welcomed from any source in primary or secondary care.

What will the LTU want to know?

The LTU will require details of the primary liver disease and its complications, comorbidity, compliance issues, alcohol or drug misuse, family support, previous abdominal surgery and cancer.

A sample transplant assessment tool used is attached as online supplementary appendix 2, but shouldn't replace a letter covering the above issues.

Table 2 Other aetiologies (non-liver failure) suitable for LT ^{5 6 8 15}	University In the set of the set (UCC) is
Variant syndromes*	Hepatocellular carcinoma (HCC)†
 Hepatopulmonary syndrome Persistent and intractable pruritus Polycystic liver disease Familial hyperlipidaemia Recurrent cholangitis Familial amyloidosis Hepatic epithelioid haemangioendothelioma Nodular regenerative hyperplasia Hereditary haemorrhagic Telangectasia Ornithine transcarbamylase deficiency Glycogen storage disease: symptomatic or presence of hepatic adenoma(s) Primary hyperoxaluria: presence of renal impairment Porphyria Maple syrup urine disease Portopulmonary hypertension Consider referral if raised mean pulmonary artery pressure (≥2.5 mm Hg), PVR >120 dynes/s/cm⁻⁵: PCWP <15 mm Hg with clinical response to medical therapy 	 Up to 25% of liver transplants in UK have HCC Associated with most CLD (HBV, HCV, ALD, NAFLD, autoimmune liver disease, haemochromatosis) and Aflatoxin ingestion Current LT selection criteria: Single tumour <5 cm in diameter, or Up to five tumours all ≤3 cm, or Single tumour >5 cm and ≤7 cm in diameter if no progression over 6 months (larger HCC's can be 'downstaged' by local therapies and then considered for transplantation) AFP <1000

*A variant syndrome is a patient with chronic liver disease whose UKELD score is <49.

†All patients with HCC should be managed within a Liver cancer MDT, which would be expected to recommend referral for liver transplantation as one of the potential 'outcomes'.

AFP, alpha foetoprotein; ALD, alcoholic liver disease; CLD, chronic liver disease; HBV, hepatitis B virus; HCV, hepatitis C virus; NAFLD, non alcoholic fatty liver disease; PCWP, pulmonary capillary wedge pressure; PVR, pulmonary vascular resistance.

Table 3 Local investigations and key information to include in referral letter (more details in online supplementary appendices 3–5)				
Comorbidity (include all details of pathologies, and important negatives)	Investigations (general investigations as below)	Disease-specific investigations		
Cardiovascular	Chest X-ray	PSC: colonoscopy and recent liver imaging		
Respiratory	ECG	PBC/AIH: drug history		
Renal	Echocardiogram	Hepatitis B: screening tests and viral load		
Bone	Oxygen Saturation	Hepatitis C: details of treatment		
HIV	Analysis of Ascites	HCC: recent imaging and MDT discussions		
Obesity	Endoscopy	PLD/PLKD: brain imaging for Berry aneurysms		
Surgical/anaesthetic history Nutritional assessment		Budd-Chiari syndrome: history of shunts.		
Nutrition	Assessment of the performance status	Wilson's disease: details of treatments		
Metabolic syndrome	Up to date blood tests and UKELD	Encephalopathy: brain imaging, ammonia, number connection tests		
Non-hepatic cancer	Adherence/addiction			
Infectious disease		Ascites: number of drains, episodes of SBP		
Social support		Alcohol related liver disease: period of abstinence, engagement with		
Disabilities		addiction services and so on		
Alcohol and substance abuse				
Mental health				
Smoking				

AIH, Autoimmune Hepatitis; ArLD, Alcohol related Liver Disease; HCC, Hepatocellular Carcinoma; MDT, Multi Disciplinary Team; NG, Naso-gastric; NICE, National Institute for Health & Care Excellence; PBC, Primary Biliary Cholangitis; PLD/PLKD, Polycystic Liver Disease/Polycystic Liver & Kidney Disease; PSC, Primary Sclerosing Cholangitis; SBP, Spontaneous Bacterial Peritonitis; TC, Transplant Co-ordinator.

The quality and content of the referral can influence the pace of transplant assessment. This guide is not intended to be exhaustive, but ensures referrers provide critical information including; general investigations required prior to referral (online supplementary appendix 3); disease specific data required by LTU (online supplementary appendix 4) and comorbidities, psychosocial factors and addiction data that may inform the transplant assessment process (online supplementary appendix 5). All summarised in table 3.

Considerations in patients with alcohol and drug-use disorders

Alcohol

Best practice suggests that patients benefit from early referral to, and engagement with, local addiction services. Repeated non-adherence with documented advice to abstain from alcohol is an absolute contraindication to LT, so all discussions regarding the requirement for lifelong abstinence must be documented and the patient informed of the implication.

The UK Liver Advisory Group¹⁶ and National Institute for Health and Clinical Care Excellence (NICE)¹⁷ have recently updated the policies relating to referral of alcohol related liver disease (ArLD) patients for consideration of LT.

Patients:

- Who are alcohol dependent and continue drinking (even at reduced levels) should not be referred. Referral to alcohol services and engagement is mandatory.
- Who, after 3 months of validated abstinence, still have an indication for liver transplant, should be referred. Validation of abstinence includes random blood alcohol

levels, alcohol metabolite testing and support from addiction services.¹⁷

▶ Who are abstinent for <3 months, and positively engaged with addiction services, can be referred if there are issues (nutrition, frailty, etc) that might complicate the assessment, or death from liver disease may occur within 3 months.

Thus, there is no absolute rule governing the period of abstinence, other than patients *must* be abstinent at the time of referral. A pragmatic approach is to advise all patients with a failing liver due to alcohol, to become abstinent and engage with addiction services. If there is no immediate indication for referral for LT (as outlined in NICE guidance) then wait for 3 months to observe and if no improvement occurs, refer for LT. If, at 3 months, there is evidence of ongoing liver recovery, then a further 3-month deferment may optimise liver recovery, and also test the patient's commitment to abstinence. At 6 months of abstinence, little further recovery can be expected and referral for LT is appropriate for any patient who remains in liver failure.

Drug addiction/misuse

Drug testing is part of the assessment for such patients. The use of prescribed methadone or buprenorphine replacement therapy does not preclude assessment for LT. However, current use of non-prescribed controlled drugs, addictive medications or 'designer' alternatives precludes referral.

LT ASSESSMENT

On receipt of a referral, the LTU determines whether the assessment requires urgent transfer, elective

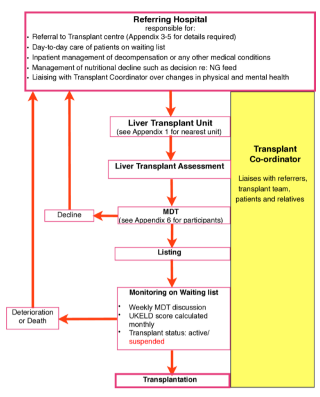


Figure 2 Flow diagram displaying the process of LT assessment from initial referral through workup and listing meeting, to monitoring on the list and either transplant/ death or suspension from the list. The roles and responsibilities of the local unit and the TC are shown. LT, liver transplant; MDT, multi-disciplinary team; NG, naso-gastric; TC, transplant coordinator; UKELD, UK Model for End-Stage Liver Disease

inpatient review, outpatient review or a combination approach. The referral-to-decision times of LTUs should be available for patients and referring hospitals (see figure 2).¹⁸

The aims of the liver transplant assessment are:

- ► To confirm the hepatological diagnosis.
- ► To confirm medical treatment has been optimised.
- ▶ To confirm that LT remains the most appropriate option.
- ► To evaluate mental and physical health comorbidities.
- ► To identify any contraindications.
- ▶ To ensure patients are fully informed of LT.

The transplant assessment is managed by the transplant coordinator (TC) and performed by the core MDT, consisting of physician, surgeon, anaesthetist, social worker, TC and dietician with additional input from pharmacist, addiction specialist, renal physician, oncologist and psychiatrist (see online supplementary appendix 6).

Transplant coordinator

The LT evaluation is supervised by the TC.¹⁹ The TC communicates directly with the patient and family/ support network. This relationship evolves over the assessment process and beyond, depending on progress (figure 2). The TC gains invaluable insight into the candidate.

Table 4 Absolute and relative contra-indications to LT			
Absolute contraindications	Relative contraindications*		
Untreated HIV*	Inadequate social support		
Severe extrahepatic disease with predicted mortality >50% at 5 years including psychiatric disorder	Smoking		
Severe irreversible pulmonary disease	Certain anatomical variants		
Ongoing alcohol misuse	Extensive previous abdominal surgery		
Active illicit drug use	$BMI > 40 \text{ kg/m}^2$		
Certain anatomic variants	Poor clinic attendance and/or adherence		
Ongoing extra-hepatic sepsis*			
Active or previous extra-hepatic malignancy†			
Liver cancer outside criteria*			
*These contra-indications can be temporary and require discussion with			

LTU. tT is considered for patients with neuroendocrine tumours (requires

referral to national panel).⁴⁶

BMI, body mass index; LT, liver transplant; LTU, liver transplant unit.

As up to 40% of assessments are declined, the TC's role includes advising and supporting such individuals, especially those who assumed referral *automatically* implied acceptance onto the waiting list. For patients accepted onto the waiting list, the TC communicates with patient's referring hospital, GP and community services. The TC provides an explanation of what patients can expect while waiting and when called in for LT.

Indications for LT

The transplant hepatologist confirms the primary liver condition (section 2), makes certain that medical treatment has been optimised, and ensures disease specific investigations are completed (see online supplementary appendices 3-5).

Contraindications to LT

Table 4 describes absolute and relative contraindica-tions to LT.

Conclusion of transplant assessment

MDT decisions

Following assessment, candidates are discussed at the MDT meeting with three potential outcomes: decline, defer, or accept. For those declined, the option of a second opinion, from another transplanting centre, is discussed with the patient and carers either by the hepatologist or the TC. Responsibility for such a further referral rests with the patient's local physician or GP.

Some patients require further optimisation such as nutritional or cardio-respiratory input, prior to activation on the waiting list. Such 'deferred' patients require re-discussion, once the additional elements are addressed. The remainder of assessments will be 'accepted' and placed on the LT waiting list. The option of live donation will be discussed, if appropriate.

Consent for LT

Consent for LT for candidates who retain mental capacity includes the general guidance for individuals undergoing any clinical intervention (see www.gmc-uk. org). However, the nature and risks of solid organ transplantation, means the process is more complex. Fuller guidance is given by the British Transplantation Society and National Health Service Blood and Transplant (https://bts.org.uk/wpcontent/uploads/2016/09/12 BTS NHS Consent April 2013-1.pdf). LTU's provide oral and written information concerning the risks and benefits of LT for patients and their carers, incorporating outcomes, donor organ related risks (infective, malignant, autoimmune, metabolic and others), procedure risks, disease recurrence and the need for adherence and life-long follow-up. The right to decline certain organs is carefully discussed (see Part 2, figure 1).

HOW TO MANAGE THE PATIENT ON THE WAITING LIST Waiting list

During 2015–2016, 1161 patients joined the LT waiting list in the UK. At 1-year post-registration, 73% of patients had received a LT, 9% had died waiting or been removed due to deterioration. A further 4% were removed for other reasons such as clinical improvement, non-compliance or at patient's request. The remaining 14% were still waiting.³

The waiting times depend on several factors, including recipient blood group, size and illness severity (i.e. UKELD score). The median waiting time is currently 135 days, though this is shorter for recipients who are blood group AB (56 days), A (84 days) and B (129 days) than blood group O (256 days).³ Details of organ allocation are outlined in Part 2.

Monitoring on the waiting list

Patients listed for LT must be monitored closely for changes in their clinical circumstances. The 'local' team manage *all* the patient's routine care, but as this may impact on their suitability for LT, regular and careful communication with the LTU, via the TC is essential (see figure 2).

Optimisation of patients on the waiting list Managing deterioration

Patients on the LT waiting list frequently present with decompensation to their referring hospital. This decompensation carries a significant mortality risk.²⁰

The Lancet Standing Commission on Liver Disease has highlighted the need to focus on improving the care for acutely ill, hospitalised patients with liver disease.²¹ Ideally, hospitals should have in place a 'care bundle' for patients admitted with decompensated cirrhosis.²²

Management of listed patients in the referring hospital

Patients on LT waiting list require monthly review in gastroenterology/hepatology outpatient clinics for nutritional review, blood-sampling, UKELD calculation and surveillance tests for cirrhosis. The LTU should be appraised of any deterioration, as listing status may need reviewing. Transfer to the LTU for acute deterioration is sometimes necessary, but if not, the confidence of patients and their carers is enhanced when there is open dialogue between the local hospital and the LTU.

Nutrition

Most patients with end-stage cirrhosis are malnourished²³ and malnutrition in recipients predicts poorer outcome after LT.^{24 25} LTUs recommend a nutritional supplement for patients on the waiting list for the beneficial effect on anthropomorphic indices.²⁶ Obesity is increasingly common among patients on the waiting list for LT. Sarcopenic obesity, or severe muscle depletion in the setting of obesity, is reported in almost half of obese patients with cirrhosis and is associated with an increased risk of pre-LT mortality. For patients with compensated cirrhosis, traditional lifestyle modifications are safe, but very low calorie diets (<1000 calories/day) are not safe. For decompensated cirrhotics, low-calorie diets may exacerbate sarcopenia and malnutrition, so such patients should maintain caloric intake with higher protein, nutrient-rich foods.²⁷

Exercise

There is little data on exercise in patients waiting for LT. However, exercise programmes known as 'prehabilitation' can improve functional capacity and postoperative outcomes in patients with liver disease awaiting major surgery.²⁸ Recently, the Birmingham group published a proposal to clarify this issue in patients awaiting LT.²⁹ Most centres advise a sustained exercise target of up to 25 minutes daily, depending on the individual patients pre-existing activity levels and physical impairments. A step-counter (available as an app) is helpful for target setting and the Birmingham team's proposal also incorporates specific advice on functional resistance exercise training that can readily be carried out at home.²⁹

Other lifestyle factors

Lifestyle factors are central to survival both on the waiting list and beyond. Patients should be supported to make positive and sustainable lifestyle behaviour changes.

Active engagement with alcohol services is key for any ArLD patient and this may be a condition for listing. Alcohol use rates of 15%–25% have been reported for patients on LT waiting lists so ongoing monitoring (blood, urine, breath or hair) and support are essential.³⁰ Most centres require random testing for patients listed for ArLD and will liaise with the local hospital to facilitate timing and frequency.

Active smoking increases all-cause mortality at 5 and 10 years after LT³¹ due to additional surgical complications, more cardiovascular disease, sepsis and solid-organ malignancy.^{31–34} Smokers must engage in a smoking cessation programme and if smoking cessation is a condition for listing, patients should expect carbon monoxide breath-testing.

Psychosocial support

Patients listed for LT have high rates of psychological distress and depression which reduces quality of life, adaptive coping and functional status.³⁵ Significant depression reduces pretransplant survival.³⁶ Waiting times, concern about deterioration, organ scarcity and false-alarms contribute to patient anxiety. All patients require screening for depression and management accordingly.

Patients and carers receive education, including attendance at patient information sessions at the time of listing. This should include access to 'expert patients', information on LT services, the patient pathway, support groups and services offering psychological, social and spiritual/cultural support. The LTU have a transplant healthcare professional available 24/7 for telephone advice for patients and carers and ongoing contact with TC, specialist nurses and a social worker, as necessary.

PALLIATIVE CARE AND LT

Liver transplantation is the gold standard treatment for many patients with ALF, CLF and liver cancer, however, almost half of those assessed are declined and 20% patients die while waiting, meaning four people die for every one transplanted. Patients with CLD have a variable level of access to palliative care services, which can vary in quality.^{37 38}

By definition, those listed for LT have advanced disease and therefore, concomitant physical and psychosocial issues. Patients suffer from a high physical symptom burden^{39 40}; symptoms are often complex and dramatic, frequently necessitating hospital admission⁴¹ with a poor evidence base to support the use of many drugs. Patients also suffer

Key points

- Consider all patients with decompensated CLD for liver transplantation
- Discuss all ALF patients with LTU
- Provide LTU with details of primary liver disease and comorbidities in mental & physical health
- Patients on transplant waiting list are the shared responsibility of referrer and LTU
- The palliative care needs of the patient are a critical part of the transplant assessment process.

complex psychosocial issues including the stigma of liver disease, complex socioeconomic background circumstances and uncertainty associated with the potential for re-compensation and the possibility of transplantation.

Maintaining hope while ensuring an appropriate holistic assessment of the patient's needs is challenging for everyone involved in delivering care. Discussions on end-of-life care including preferences, is an essential component in the management of patients with advanced liver disease, including those awaiting LT.

Early palliative care intervention can improve quality of life by improving symptom burden and mood, alongside less aggressive treatment and a reduction in hospitalisation.^{42 43} An assessment of palliative care needs should form an integral part of any transplant assessment process. Collaborative working is essential at a time of such great uncertainty if the overall quality of life for these patients and their carers is to be improved.

CONCLUSIONS

- All patients with decompensated CLD, should be considered for LT.
- ► All ALF patients should be discussed with the LTU and transferred in timely fashion, if appropriate.
- LT may be considered for variant syndromes, HCC and various other non-liver failure indications after discussion with LT team.
- The LTU require details of diagnosis, comorbidity, nutritional status and frailty in order to complete evaluation.
- Patients on the LT waiting list are shared between the referring centre and LTU. Good communication between the various healthcare professionals is critical. For patients who deteriorate, candid discussions with individuals and their relatives coupled with timely utilisation of palliative care services should optimise outcome for all concerned.

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together the co-ordinators input with the transplant centre and the interaction with secondary care referring centre. Providing insights from both sides of the secondary/tertiary care interaction and then editing the initial contribution of the section to a more manageable section. Ensuring relevance of data collected for the project along with final manuscript editing and approval KM (consultant transplant surgeon) contributed to the section in part 2 on Transplant surgery and outcomes. KM also provided input in Part 1 into section on Transplant assessment and previous surgery. Critical role in planning and acquisition of data for inclusion into guideline, along with evaluation of relevance to project. Final manuscript editing and approval DM (consultant transplant surgeon) co-authored the section on Transplant surgery with KM and post-transplant surgical complications. He also provided input in Part 1 into section on Transplant assessment and previous surgery. Critical role in planning and acquisition of data for inclusion into guideline, along with evaluation of relevance to project. Final manuscript editing and approval JN (consultant transplant hepatologist) wrote the section on Post-Transplant Immunosuppression with the aid of A Considine. Prof Neuberger also contributed to the Part 1 in editorial role, when he helped with the original concept, he provided a critical role following the original guideline production and prior to it's splitting into two halves (Part 1 and 2) and ensuring correct focus was maintained when the manuscript was reduced in size. RP (consultant transplant surgeon) wrote the section on organ allocation and donation. He also made a significant contribution to the post-operative care and complications sections and the pre-op evaluation (Part 1). Critical role in planning and acquisition of data for inclusion into guideline, along with evaluation of relevance to project. Final manuscript editing and approval AP (Liver Pharmacist) contributed to Part 2 immunosuppression section, but also gave useful input into Part 1 and 2 from point of view of medication and drug interactions, particularly with viral hepatitis treatment. Critical role in evaluating data for inclusion into guideline, along with maintain relevance to project. Final manuscript editing and approval WP (Consultant in Palliative Care) authored the section on Palliative Care and Transplantation in Part 1. Her contribution in Part 2 was proof-reading and providing critical input. Critical role in planning and acquisition of data for inclusion into guideline, along with evaluation of relevance to project. Final manuscript editing and approval LS is a Transplant co-ordinator. She contributed to the section on How to refer a patient for liver transplant (Part 1) and gave input into section on Organ allocation (Part 2). Critical role in planning and acquisition of data for inclusion into guideline, along with evaluation of relevance to project. Final manuscript editing and approval KS (consultant transplant hepatologist) wrote the section on 'When to refer' in Part 1, but also gave significant editorial input to entire project, at the time of the section merge and subsequent division into two halves. Critical role in planning and acquisition of data for inclusion into guideline, along with evaluation of relevance to project. Final manuscript editing and approval DT and RW (consultant transplant hepatologists) co-authored the section on Transplant outcomes. They provided critical input into Part 1 as well, with respect to referral process. Critical role in planning and acquisition of data for inclusion into guideline, along with evaluation of relevance to project. Final manuscript editing and approval. DT (consultant transplant hepatologist) co-authored the section on Postoperative care and complications (nonsurgical). DT also supported the entire process by helping the lead author with editing sections and discussion of tables, pictures, deciding on section inclusion, data relevance and final manuscript editing and approval.

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Appendix 1: UK & IRELAND TRANSPLANT CENTRES

DIAGNOS	SIS						
CO-MOR	BIDITY						
BLOOD T	ESTS: (*wi	thin 1 mo	nth of refer	ral)			
Blood Gro				BMI	kg/m ²		
MELD		UKELD		C-P	3	*GFR	mls/min
*Bil	umol/L	*Alb	g/L	*ALP	IU/L	*ALT	U/L
*PT	secs	*Hb	g/L	*PLT	x10 ⁹ /L	*WBC	x10 ⁹ /L
*Creat	umol/L	*Urea	mmol/L	*Na⁺	mmol/L	*K ⁺	mmol/L
Ferritin	ug/L	B12	ng/L	Folate	ug/L		
ANA	0	AMA		SMA			
HBV		HCV		HIV		CMV	
Glucose	mmol/L	tTG		AFP	kU/L	A1AT	
IgM	g/L	IgA	g/L	lgG	g/L		
тѕн	mu/L	J	0	MRSA	0		
*PaO ₂ (F	iO2 %)	kPa	ι	*PaCO	kPa		
	RESPIRATO	ORY etc:			•		
FEV1	L	FVC	L	FEV1/F	VC		
Echocard	iogram: (dat	e / /)				
ECG: (date / /) CXR: (date / /)							
CPEX: (da					,		
-							
IMAGING	i :						
USS Live	r: (date /	/) Port	al vein Pate	nt Y/N, As	cites Y/N, H	CC Y/N	
MRI: (date / /)							
CT: (date							
GASTRO	SCOPY: (da	ate / /)				
DIETICIA	N:						
Notes:				-			
Handgrip kg MAMC cm RFS							
DRUG HI	STORY:	ALC	COHOL/ADI	DICTION	SOCIAL	SUPPOR	T etc:
NOTEO							
NOTES:	NOTES:						
**Color	noscopy wit	hin 12 m	onths if PS	С			
00101				-			

Appendix 2: Transplant Assessment Proforma

INVESTIGATION	WHY IS THE TEST REQUIRED?	
Chest X-ray	Normal/Cardiomegaly/Pulmonary hypertension/Effusion/Metastases (in HCC patients)	
ECG	Normal/Ischaemic changes/Right or left ventricular enlargement/Right heart strain/Conduction block and	
	rhythm disturbance. A prolonged QTc is often seen in cirrhosis.	
Echocardiogram	Echocardiogram is mandated in patients in whom TIPSS is being considered. Note signs of diastolic and	
	valvular dysfunction. Systolic function is often over-exaggerated by the hyperdynamic circulation.	
Oxygen Saturation	Low oxygen saturation can be a feature of hepatopulmonary and/or porto-pulmonary syndromes or may	
	indicate parenchymal lung disease.	
Analysis of Ascites	All patients with ascites should have analysis for protein levels, SAAG gradient, cell counts and differential,	
	cytology and culture (including Tb in at risk patients) and antibiotic prophylaxis offered if indicated.	
Endoscopy	All PSC referrals should have had a recent colonoscopy if safe.	
	All referrals should have had a gastroscopy for varices assessment and consideration of prophylaxis	
Nutritional	Malnutrition and sarcopenia are commonplace in end-stage liver failure. All patients should be assessed	
Assessment	by dieticians for an assessment of their nutritional state	
Assessment of the	Some patients are unfit for transplantation and need extensive pre-conditioning work-up to improve their	
performance status	peri-operative morbidity	
Up to date blood	Patients are stratified on the waiting lists according to their UKELD score so up to date blood work allows	
tests and UKELD	for prioritisation [https://www.odt.nhs.uk/transplantation/tools-policies-and-guidance/calculators/]	
Adherence/Addiction	Any concerns with regard to adherence, clinic attendance and/or engagement should be highlighted at an	
	early stage. See above	

Appendix 3. General investigations to be completed before patients are referred for assessment.

	Disease specific considerations in assessme			
DISEASE	Referring centre	Transplant Centre		
PSC	PSC patients should have had colonoscopy in the last 12 months. The right colon must be visualised as	Transplant indication includes recurrent cholangitis and sepsis. PSC patients with IBD must have IBD optimised as		
	colorectal cancer is increased in this patient group. Recent cross-sectional imaging should be sent	poorly controlled colitis pre-transplant negatively impacts on graft survival.		
PBC/AIH	Include a full drug history including immunosuppression,			
Hepatitis B	Screen all candidates for HBV (HBsAg, HBeAg, HBcAb) All HBV sAg +ves, require viral load and HDV status. Include medication history	 Consider vaccination for all sero-negatives. Previous contact with HBV requires testing for HBV DNA as detectable HBV DNA will require suppression, pre- transplant. Patients transplanted for HBV disease, will receive HBIg and nucleos(t)ide analogues peri- and post transplant, according to local protocols. 		
Hepatitis C	If PCR positive, genotype, viral load and treatment history should be provided. Whilst all patients with a failing liver should be discussed with a LTU, most will recommend eradication therapy prior to surgery for patients with MELD scores <20, and vice versa for MELD score >20	DAAs have reduced the number of patients requiring liver transplant and up to 25% listed with MELD <20 will improve and be-de-listed, after eradication of virus.		
HCC	Recent imaging, notes from HPB MDT discussions, any previous loco-regional therapy (including dates) and information pertaining to response and tumour size/load prior to treatment, must be included Cancer staging and monitoring protocols vary from unit to unit.	called the "extended Milan criteria". The Milan Criteria		
PLD/PLKD	MRI/MR Angiogram of brain to exclude berry aneurysms	Э.		
Alcohol	A six month period of abstinence is recommended	1. All patients referred for liver transplantation receive a full		
related Liver	before listing to optimise liver recovery, and to test the	psycho-social evaluation.		
Disease	patient's commitment to abstinence. However, NICE	2. A further structured substance misuse evaluation with		

Appendix 4. Disease specific considerations in assessment

	recommends referral after 3 months of abstinence to allow for the period of evaluation and waiting and minimise the chance of the patient deteriorating beyond transplantation.	additional psychiatric evaluation is usually carried out. 3. Also, patients will be requested to sign a contract in the presence of family to adhere to abstinence after transplant.	
Budd-Chiari	The LTU will require all records of discussions with	BCS management consists of trying to re-establish	
syndrome	regional HPB unit, historical shunting procedures,	venous drainage of the liver and resorting to transplant	
	surgery etc as well as details of procoagulant disorders	only if stents and shunts have failed. The more severe and	
	tested for.	acute the presentation the more likely that transplantation	
		will be necessary.	
Wilson's	Wilson's disease may present as either acute liver	Patients and family should know that liver transplantation	
disease	failure as well as decompensated chronic liver disease	cures the liver disease and underlying metabolic	
	in patients who did not respond to medical therapy. disturbance, but not the neuropsychological features		
Encephalop	Provide relevant brain imaging (MRI preferably). If diagnostic doubt persists provide EEG reports and/or blood		
athy	ammonia measurements. Detail any hospital admissions with hepatic encephalopathy.		
Ascites	Detail the number and frequency of ascitic drains and whether there has been evidence of SBP. Describe		
	complications such as loculated ascites, hydrothorax and/or haemorrhage.		

Supplementary material

Appendix 5: Co-morbidity	Ap	pendix	5: (Co-mo	rbiditv
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CO-MORBIDITY	RELEVANCE
	Cardiomyopathy is seen with cirrhosis and alcohol. Several liver-lung syndromes (HPS/PPHT/HHT) are described. A
Cardiovascular	clinical history, contrast echocardiography and other invasive tests may be necessary.
	Any history of cerebrovascular disease should be sought and described i.e. haemorrhagic/embolic/infarct and
	relevant imaging and investigations supplied.
Respiratory	All respiratory disease and investigations should be described, including any ITU admissions and performance
Respiratory	status. For hypoxic patient, include cross-sectional imaging, lung function, transfer studies and tests looking for
	PEs/Shunts & AVMs
Renal	For patients with chronic kidney disease (CKD) include investigations and treatments. If the eGFR
	<30mls/min/1.73m ² for more than 3 months, a combined liver kidney graft may be necessary. Urinalysis, Albumin to
	Creatinine ration (ACR), renal ultrasound and cultures should be completed in all patients with an eGFR
	<60mls/min/1.73m ² .
Bone	Assess the FRAX score in all patients (https://www.shef.ac.uk/FRAX/tool.jsp)
HIV	HIV on treatment, is not a contraindication to liver transplantation. Enclose details from specialist teams of treatment
	history, adherence and drug prescription.
Obesity	Record BMI and estimate dry BMI if ascites present. BMI>40kg/m ² is a relative contraindication
Surgical/Anaesthetic	All surgical history should be detailed in the referral letter
Nutrition	A dietetic assessment prior to referral is mandatory. Offer appropriate nutritional support (e.g. protein supplements
	and/or NGT feeding)
Metabolic Syndrome	Request an HbA1c (note anHbA1c may be artificially low in patients with chronic anaemia)
	For diabetes, document the duration, time on treatment (including years on insulin therapy), urinalysis, eye
	examination & fundoscopy, vascular complications.
	Record other components of metabolic syndrome such as hypertension and dyslipidaemia.
Non-hepatic cancer	A history of cancer may not be a contraindication
	Details of dates of diagnosis, staging, treatment, prognosis, 5 year survival and correspondence from treating
la fa ati ang Dianang	oncologist
Infectious Disease	Any communicable disease (including TB) in the patient and household contacts should be reported
Social Support	Housing, next of kin, adherence with appointments, tests and therapies should be included in the referral letter
Disabilities	Learning disabilities or deafness/ visual impairment does not prevent assessment for transplantation
Alcohol & substance	Alcohol, illicit and prescribed drug usage including analgesics and sedatives, must be reported
abuse	
Mental health	Prior history and treatment for mental illness, including self-harm should be detailed in the referral letter
Smoking	Active smokers should be advised to stop and be referred to smoking cessation service

The Medical Assessment:		
General health Past medical history, current (non hepatological)		
1		
	Livor diagon	medical issues, medication etc (see Table 3)
.	Liver disease	Confirm history of liver disease, diagnosis,
1		management and current treatment.
1		Disease-specific evaluations (see Table 4)
1		(If hepatocellular cancer present, oncology input)
1	Drug History	To include allergies
	Urine tests	Glucose, protein, drug-screen (if relevant)
ist	Blood tests	Liver tests (non-invasive liver screen, synthetic
og		function), renal function, viral screen blood-typing
t <u>o</u>	Cardio-	ECG, PFTs and echocardiography (if not recently
ра	pulmonary	performed). Further testing, such as stress testing etc
Hepatologist		with advice from cardiologist (see Tables 3)
_	Radiology	CXR, USS liver and CT/MRI depending on indication
		etc (see Tables 3)
	Cancer risk	Breast/Colon/Cervix where appropriate
	Latent infection	CMV status pre-transplant and post transplant
1		
1		prophylaxis
		HBV etc
	F	HIV status and treatment related issues
	Explanation	Explanation of process, all outcomes etc
The Surgical Assessment:		
Sur	gical team	Confirm liver transplant is indicated.
1		Surgical issues: previous abdominal surgery, obesity,
		portal vein compromise, anatomical variants
		Discussion of procedure, risks, complications and
L		organ issues.
The Dietetic Assessment		
	tician	Assess nutritional status, including anthropometry
		Assess patient (and family) understanding of nutritional
		advice.
		Co-ordinate with dietetic service at referring hospital
The	Anaosthatia As-	
The Anaesthetic Assessment Anaesthetist Previous anaesthetic issues.		
Ana	ເຮວແມ່ນເຮັເ	
1		Standard tests include Pulmonary function tests and
		Oxygen saturation.
1		Risk assessment including specific cardiopulmonary
1		issues.
1		May request CPEX or DSE etc
1		Discussion with patient/family over ICU, surgery
		process etc
The Psychosocial Assessment		
Social Worker		Psychosocial issues, family/support mechanisms,
		effect on dependants etc
Add	liction specialist	Tobacco, alcohol and illicit drug dependencies
	chologist	Mental health issues, addiction support etc

Appendix 6: The Liver Transplant Multi-disciplinary Assessment (Coordinated and overseen by Transplant Co-ordinator)