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**Liver surgery in cirrhosis and portal hypertension**

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**Abstract**

The prevalence of hepatic cirrhosis in Europe and the United States, currently 250 patients per 100000 inhabitants, is steadily increasing. Thus, we observe a significant increase in patients with cirrhosis and portal hypertension needing liver resections for primary or metastatic lesions. However, extended liver resections in patients with underlying hepatic cirrhosis and portal hypertension still represent a medical challenge in regard to perioperative morbidity, surgical management and postoperative outcome. The Barcelona Clinic Liver Cancer classification recommends to restrict curative liver resections for hepatocellular carcinoma in cirrhotic patients to early tumor stages in patients with Child A cirrhosis not showing portal hypertension. However, during the last two decades, relevant improvements in preoperative diagnostic, perioperative hepatologic and intensive care management as well as in surgical techniques during hepatic resections have rendered even extended liver resections in higher-degree cirrhotic patients with portal hypertension possible. However, there are few standard indications for hepatic resections in cirrhotic patients and risk stratifications have to be performed in an interdisciplinary setting for each individual patient. We here review the indications, the preoperative risk-stratifications, the morbidity and the mortality of extended resections for primary and metastatic lesions in cirrhotic livers. Furthermore, we provide a review of literature on perioperative management in cirrhotic patients needing extrahepatic abdominal surgery and an overview of surgical options in the treatment of hepatic cirrhosis.

**Key words:** Liver resection; cholangiocellular carcinoma; hepatocellular carcinoma; liver metastases; portal hypertension; cirrhosis

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**Core tip:** Liver resections in patients with underlying hepatic cirrhosis and portal hypertension still represent a medical challenge with regard to perioperative morbidity, surgical management and postoperative outcome. However, the increasing incidence of liver cirrhosis on the one hand, and the ongoing improvements in surgical technique and perioperative management on the other hand have up until today rendered even extended hepatic resections in these patients possible. Especially in primary and metastatic liver malignancies, surgery often presents the only curative approach and thus potential short- and long-term benefits and risks have to be evaluated carefully and interdisciplinary from a surgical, oncological and hepatologist point of view.

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**INTRODUCTION**

***Liver cirrhosis and portal hypertension***

Liver cirrhosis, the end-stage of chronic liver fibrosis, shows a prevalence of 250 patients per 100000 inhabitants[[1](#_ENREF_1),[2](#_ENREF_2)]. Pathological alterations of liver parenchyma and hepatic perfusion lead to a decrease in hepatocyte function and an increase in transhepatic perfusion resistance, resulting in portal hypertension. Over time, up to 20% of cirrhotic patients develop a hepatocellular carcinoma[1].

Causes of liver cirrhosis are manifold (Table 1). In Europe and the United States, alcoholic liver disease is the predominant cause of hepatic cirrhosis, followed by chronic viral hepatitis (B/C) and nonalcoholic fatty liver disease. In many African and Asian countries, chronic viral hepatitis remains the main cause of liver cirrhosis[[1](#_ENREF_1)].

Phenotypically, liver cirrhosis can show a variety of symptoms from almost asymptomatic stage to liver failure. In advanced cirrhosis, live-threatening complications are bleeding from esophageal varices, hydroptic decompensation (ascites), spontaneous bacterial peritonitis and hepatic encephalopathy. A first evaluation of the cirrhotic stage can be estimated by the Child-Pugh-Turcotte Score (CPT) using the INR (International Normalized Ratio), the serum bilirubine, the serum albumine, the grade of hepatic encephalopathy and the grade of ascites[[3](#_ENREF_3)]. Although the CPT score is a valuable method to categorize patients into individuals with compensated (CPT A), mildly (CPT B) or severely decompensated (CPT C) cirrhosis, it can be influenced by the subjective assessment of the parameters “ascites” and “encephalopathy”. Furthermore, it does not include the prognostic relevant assessment of renal function.

Another system evaluating the mortality of a patient with liver cirrhosis is the Model of End Stage Liver Disease (MELD) score, currently used for organ allocation in liver transplantation[[4](#_ENREF_4),[5](#_ENREF_5)]. It is reproducibly calculated from creatinine, bilirubin and INR, and does not include less precise parameter such as ascites or varices. Since a continuous deterioration of each individual parameter results in a continuous increase in MELD score, risk discrimination of individual patients is more accurate than using the CPT score[[6](#_ENREF_6),[7](#_ENREF_7)]. However, neither the CPT- nor the MELD-score were originally created for surgical settings. While the CPT classification was developed to estimate the liver-specific overall survival of cirrhotic patients, the MELD-score was developed for estimating the mortality after TIPS (transjugular portosystemic stent shunt) –placement[[4](#_ENREF_4),[5](#_ENREF_5)]. A main drawback in surgical risk stratification is that both scoring systems do not include features of portal hypertension except for ascites in the CPT classification.

Portal hypertension is defined as an increased pressure gradient between portal vein and hepatic veins [= portal-pressure gradient (PPG) also called the porto-caval gradient/porto-systemic gradient]. The hepatic venous pressure gradient (HVPG) is an indirect measurement of the real PPG, which allows obviating the risks linked with the direct puncture of the portal vein[[8](#_ENREF_8)]. Under physiological conditions, the HVPG is 1 to 5 mmHg. Subclinical portal hypertension is defined as a HVPG of 5 to 9 mmHg. Clinically relevant portal hypertension shows a HVPG of ≥ 10 mmHg. Reasons for portal hypertension are shown in Table 2. In western countries, hepatic cirrhosis accounts for > 90% of cases of portal hypertension.

Gold standard for assessing the HVPG is the angio-invasive measurement of the free hepatic venous pressure (FHVP) in the right hepatic vein as well as the wedge hepatic venous pressure (WHVP) at the same location[[8](#_ENREF_8)]. HVPG is then defined as the difference of WHVP and FHVP. Since this invasive method may not be available everywhere and might also cause complications, translational studies are currently performed to establish ultrasound-based elastography for assessing the degree of portal hypertension. So far some promising progress has been made using Fibroscan® measurements. Here, a cut-off value of 13.6 kPA has been described as being 90% sensitive to detect clinically relevant portal hypertension, with excellent specificity at a higher cut-off value of 21.1 kPa. Thus, invasive HVPG measurements to diagnose clinically relevant portal hypertension may be limited to patients without esophageal varices and with a Fibroscan® measurement result between 13.0 and 21.1 kPa[[9](#_ENREF_9)]. Ongoing clinical assessments will further refine this technique[[10](#_ENREF_10),[11](#_ENREF_11)]. Meanwhile, surrogate markers such as thrombocytopenia < 100/nl, splenomegaly > 12cm, the endoscopic confirmation of esophageal varices (that do not occur in HVPG < 10 mmHg[[12](#_ENREF_12)]), clinical signs of collateralization such as the presence of caput medusae and presence as well as extent of ascites are used to define portal hypertension[[13](#_ENREF_13)].

**Preoperative diagnostics**

While taking the medical history of a patient with liver cirrhosis, a main focus should lie on prior episodes of decompensation or variceal hemorrhage, ongoing alcohol abuse and further comorbidities such as congestive gastropathy, cardiovascular complaints (coronary artery sclerosis, cirrhotic cardiomyopathy), pulmonary disorders (chronic obstructive pulmonary disease (COPD), hepato-pulmonary syndrome, pleural effusion), restricted renal function (hepato-renal syndrome), diabetes and malnutrition[[14](#_ENREF_14),[15](#_ENREF_15)]. During physical examination, signs of portal hypertension such as ascites and caput medusae may be seen. Preoperative laboratory screening should include a full blood count (anemia, thrombocytopenia), coagulation tests (INR), renal function parameters (creatinine, GFR, electrolytes) as well as liver function parameters. Radiologic examinations should consist of (if possible: contrast-enhanced) ultrasound (parenchymal liver morphology, focal lesions, liver perfusion, umbilical vein recanalization, elastography if available) and contrast-enhanced CT/MRI scans (morphology of lesions indicating liver resection, staging, morphology of collaterals, volumetric assessment of remnant liver).

Several methods have been established to specifically evaluate the liver function prior to hepatic resections[[16](#_ENREF_16)]. Among them, the LiMaX (maximum liver function capacity) test, which analyses the liver-specific, cytochrome P450-based metabolism of 13C-methacetin[[17](#_ENREF_17),[18](#_ENREF_18)], and the Indocyanine-green-Clearance (ICG-Clearance) test[[19-21](#_ENREF_19)] are most commonly used. Ishizawa and colleagues have reported the combination of ascites, bilirubine and indocyanine-green-clearance to be excellent predictors of resectability for hepatocellular carcinoma (HCC) independent of portal hypertension[[22](#_ENREF_22)]. However, LiMaX-Test and Indocyanine-green-clearance both analyse the global liver function and cannot distinguish the expected post-surgical liver remnant function. A method that may distinguish this remnant liver function is functional scintigraphy and functional MRI using Gd-EOP-DTPA contrast enhancement[[23-25](#_ENREF_23)]. Nonetheless, up until today, little experience of these assays in patients with liver cirrhosis and portal hypertension exists and future prospective studies are warranted.

A critical issue is the remaining liver remnant after resection. In general, CT-based volumetric assessment is used to estimate the volume of liver that remains after resection. Upon liver cirrhosis, a minimum of 40% well-perfused liver tissue must remain in situ to maintain an adequate function[[26](#_ENREF_26)]. However, this is not an absolute value and should be taken with caution.

**Preoperative risk evaluation**

Since liver cirrhosis and portal hypertension significantly increase perioperative morbidity and mortality[[27](#_ENREF_27),[28](#_ENREF_28)], a thorough pre-operative risk evaluation is recommended. The American Society of Anaesthesiologists Score (ASA-Score), although not originally meant specifically for liver surgery, has been shown to reliably predict 7-d mortality of patients with liver cirrhosis after abdominal surgery[[28](#_ENREF_28)]. In addition, CPT and MELD score are used to predict the mortality after major abdominal surgery. Thus, mortality rates of 10%, 30% and 80% were observed for patients classified as CPT A, CPT B and CPT C, respectively[[29](#_ENREF_29),[30](#_ENREF_30)]. However, recent improvements in surgical technique and perioperative management have significantly reduced mortality especially in CPT A and CPT B patients[[12](#_ENREF_12),[22](#_ENREF_22),[29-33](#_ENREF_29)]. The impact of portal hypertension on perioperative morbidity and long-term survival has lately been discussed controversially. Berzigotti and colleagues have shown a relative risk of 2.0 for 3- and 5-year mortality and a relative risk of 3.0 for perioperative clinical cirrhotic decompensation in patients undergoing liver resection for HCC with and without portal hypertension. Similar results were shown for subgroups of patients undergoing resection of a single small HCC in CPT A cirrhosis[[12](#_ENREF_12)]. In contrast, other reports state that portal hypertension is no prognostic parameter for perioperative mortality and long-term survival in multivariate analyses[[34](#_ENREF_34)]. A matched-pair analysis of 241 cirrhotic patients undergoing liver resections for HCC showed no significant difference in perioperative morbidity and long-term postoperative outcome in patients with or without portal hypertension and identified the MELD-score and the extent of liver surgery as only specific prognostic parameters[[34](#_ENREF_34)]. Using the MELD score as predictor, mortality after abdominal surgery has been described to be 5%-10% for MELD ≤ 11, 25%-54% for MELD 12%-25% and 55%-80% for MELD ≥ 26[[35-37](#_ENREF_35)]. Furthermore, a study analysing liver resections in cirrhotic patients and showing an overall-mortality of 16%, defined a MELD score of 9 as “cut-off” point: none of the patients with less than 9 points died in the perioperative period whereas a 29% mortality rate was observed in patients with 9 or more points[[32](#_ENREF_32)]. Based on these analyses, the Mayo Clinic offers a web-based risk evaluation of 7-, 30-, 90-, 360- and 1800-day mortality for cirrhotic patients after abdominal, cardiac and orthopedic surgery (<http://www.mayoclinic.org/meld/mayomodel9.html>)[[28](#_ENREF_28)]. A synopsis of preoperative risk scores is shown in table 3. In conclusion, elective abdominal surgery in CPT C patients or patients with a MELD > 15 is only feasible with significantly increased morbidity and mortality. In these patients, potential therapeutic alternatives as well as morbidity and mortality in case of conservative therapy have to be carefully assessed and risks *vs* potential benefits evaluated for each individual patient. In CPT A and MELD < 9 patients, elective surgery including liver resections can be performed with adequate risk-to benefit ratio. Patients showing a CPT B classification and a MELD score of 9-15, the risk-to benefit ratio has to be carefully discussed for and with each patient. Of note, portal hypertension is not adequately included in all of these staging systems and has to be taken into consideration in the context of liver resection. In particular, thrombocytopenia as one of the features of portal hypertension has been associated with a significant increase in postoperative major complications, postoperative liver failure and finally with a more than 3-times higher 60-d mortality[[22](#_ENREF_22),[38](#_ENREF_38)].

**Hepatic resection in cirrhotic livers and portal hypertension**

***Indication***

The most frequent indication for hepatic resections in cirrhotic patients with and without portal hypertension is HCC. In addition, cholangiocellular adenocarcinoma (CCA) and liver metastases may be reasons for liver resection upon cirrhosis although several studies indicate that cirrhosis reduces the risk of metastasis formation[[39](#_ENREF_39),[40](#_ENREF_40)]. Benign hepatic lesions are very rare indications for hepatic surgery in these high-risk patients.

According to the European Association for the Study of the Liver (EASL) and American Association for the Study of the Liver (AASLD) guidelines for HCC therapy, surgical resections are restricted to patients without portal hypertension, patients showing a normal bilirubin level and a singular tumor nodule[[12](#_ENREF_12)]. In addition, the Barcelona Clinic Liver Cancer (BCLC) classification recommends curative liver resections for HCC in cirrhotic patients only in early tumor stages and in CPT A cirrhosis without portal hypertension[[41](#_ENREF_41)]. In contrast, several studies world-wide have shown that HCC resections can be performed in cirrhotic patients with portal hypertension showing acceptable mortality rates with clear oncologic benefits[[22](#_ENREF_22),[33](#_ENREF_33),[34](#_ENREF_34)]. Furthermore, Torzilli and colleagues could show that in centres specialized in complex liver surgery, 50% of patients undergoing liver resection in cirrhosis are resected outside the BCLC criteria with 5-year overall survival rates of 57% and 38% for BCLC B and BCLC C patients, respectively[[42](#_ENREF_42)]. Several authors could show that hepatic resections in patients diagnosed with HCC in advanced cirrhosis result in significantly better long-term survival compared to transarterial chemoembolization[[43-46](#_ENREF_43)] and that tumor size alone should not be an exclusion criteria for curative surgery[[47](#_ENREF_47)]. In a randomized controlled clinical trial, Yin and colleagues performed liver resections or transarterial chemoembolizations in 173 HCC patients exceeding the Milan criteria. The authors described a significantly better long-term survival in patients undergoing liver resection (3-year overall survical rate 51.5% *vs* 18.1%)[[46](#_ENREF_46)]. Thus, advanced liver cirrhosis and portal hypertension should not be considered as absolute contraindication in liver resections but must be evaluated individually for each patient in the context of the hepatic resection planned and further comorbidities of the patient.

***Extent of resection***

In a cirrhotic liver, a minimum of 40% well-perfused liver-volume must remain in situ after resection[[26](#_ENREF_26)]. However, in our institution these radical resections of up to 60% liver volume are restricted to CPT A, MELD < 9 patients with no or only mild portal hypertension (figure 1). Some reports indicate that anatomic hepatic resections for HCC show a trend towards increased disease-free and overall survival compared to non-anatomic resections[[48-50](#_ENREF_48)] whereas others do not see a benefit from anatomic resections[[51-53](#_ENREF_51)]. However, to avoid postoperative liver failure due to insufficient liver remnant, we would prefer hepatic resections to be as parenchymal-sparing as possible. In particular, the latter strategy may enable liver resections in patients with advanced cirrhosis (CPT B) and with (mild) portal hypertension[[54](#_ENREF_54)]. Lately, local ablation using radiofrequency has been shown to be an adequate alternative therapy for small malignant hepatic lesions < 2 cm with similar long-term oncologic outcome as surgical resection but significantly reduced peri-interventional morbidity[[9](#_ENREF_9)]. Thus, for patients with small malignant liver lesions in cirrhosis and portal hypertension local ablation has to be considered as an alternative (and potentially better) option (see figure 2).

***Surgical access***

Standard access for cirrhotic as well as non-cirrhotic patients in liver surgery is the upper abdomen midline incision extended by right-upper quadrant traverse laparotomy (“inversed L-shape”). In cases where right-posterior segmentectomy has to be performed and where clamping of the suprahepatic inferior caval vein is not possible through inversed L-shape incision, a thoraco-abdominal incision is recommended[[55-57](#_ENREF_55)]. In minor resections, a midline incision may suffice. The recanalized umbilical vein must be carefully ligated or preserved if possible. First studies have also established laparoscopic resections in cirrhotic patients. In specialized centres, even extended liver resections can be performed in a minimally invasive technique with removal of the resected liver using a Pfannenstiel incision[[58](#_ENREF_58)].

For parenchymal dissections, all techniques used in non-cirrhotic patients can be used in cirrhotic patients with or without portal hypertension. “Selective” dissection techniques are based on discriminating the higher tissue resistance of blood vessels and bile ducts compared to liver parenchyma. Selective dissection techniques are blunt dissection using a pair of scissors and selective application of clips, water jet dissection and clips, stapler dissection and ultrasound-aspiration (CUSAR) and clips[[59-64](#_ENREF_59)]. In our institution, blunt selective dissection by scissors and clips is the preferred dissection technique. In any case, care must be taken of thorough bleeding control during resection. Post-resection application of fibrin glue or collage fleece has not shown any benefit on blood loss and need for blood transfusions[[65](#_ENREF_65)].

***Pringle manoeuvre***

Blood loss and need for blood transfusions have been defined as independent predictors of postoperative morbidity and mortality[[66-68](#_ENREF_66)]. In cases of problematic blood loss during parenchymal dissection, hilar vascular occlusion can be applied (Pringle manoeuvre). Of note, the Pringle manoeuvre is not recommended in cirrhotic patients but sometimes is necessary to avoid extensive blood loss. If needed, collateral vessels in the liver hilum should be preserved during vascular clamping. If vascular clamping cannot be avoided, selective clamping of a hepatic lobe, section or segment is recommended to limit ischemic and reperfusion damage[[55](#_ENREF_55),[69](#_ENREF_69)]. Moreover, hilar vascular occlusion should be performed as short as possible. In cases where longer occlusion is needed, intermittent occlusion, *e.g.* intervals of 15 min of ischemia, followed by 5 min of reperfusion - has shown to be tolerated better than continuous clamping[70-74]. Total hepatic vascular exclusion, *i.e.* including occlusion of the infra- and suprahepatic caval vein, should be avoided since a significant increase in perioperative morbidity and mortality has been observed even in non-cirrhotic patients[[75](#_ENREF_75)]. A setting were total hepatic vascular exclusion may be necessary is the resection of a hepatocellular carcinoma invading the retrohepatic caval vein and needing (partial) resection and replacement of the retrohepatic caval vein.

***Cooperation with anaesthesiology***

During hepatic resection, a close cooperation and communication with the anaesthetic team is essential. Since intrahepatic sinusoidal pressure correlates with the pressure in the inferior caval vein, which again correlates with the central venous pressure (CVP), blood loss during parenchymal dissection can be significantly decreased by reducing the CVP to approximately 5 mmHg[[55](#_ENREF_55),[69](#_ENREF_69),[76](#_ENREF_76)]. Reduction of the CVP to lower (even negative) values is not recommended due the increased risk of air embolism[[55](#_ENREF_55)]. Restrictive volume management, reduction of tidal volume and positive end-expiratory pressure, diuretics and reverse Trendelenburg positioning are options to reduce the CVP[[77](#_ENREF_77)]. A surgical method to reduce CVP is partial clamping of the infrahepatic caval vein[[78](#_ENREF_78)]. Finally, in case of increased blood loss during hepatic resection, a decent management including not only blood transfusions but also substitution of non-cellular blood components (*e.g.* fresh frozen plasma, coagulation factors) is mandatory. It has been observed that a restrictive transfusion policy for cirrhotic patients with anemia therapeutically targeted to a Hb between 7-9 g/dL is associated with a better clinical outcome[[79](#_ENREF_79)]. This should be considered in order to avoid an overtransfusion policy that can further increase portal pressure as consequence of high hepatic inflow.

***Drainage***

Routine placement of drainage systems in hepatic resections is controversially discussed. In minor resections of non-cirrhotic livers in patients without portal hypertension, drainage placement is not recommended[[80](#_ENREF_80)]. In major resections, prophylactic drainage placement has been associated with increased septic complications and prolonged time to discharge[[81](#_ENREF_81)]. However, other publications show a significantly reduced intraabdominal pressure for temporary drainage of increased postoperative ascites and improved wound healing after drainage placement[[82](#_ENREF_82),[83](#_ENREF_83)].

**Postoperative Morbidity**

Compared to liver resections in otherwise healthy patients, resections in cirrhotic livers in patients with portal hypertension show a markedly increased postoperative morbidity of 22%-50%[[22](#_ENREF_22),[33](#_ENREF_33),[34](#_ENREF_34),[66](#_ENREF_66)].

***Liver failure***

Postoperative liver failure is the most serious and life-threatening complication after liver resection. Several definitions for “liver failure” have been proposed over the last 20 years. So far, the commonly used definitions, though initially not developed to score liver resections in cirrhotic livers, are the “50-50 criteria”, the International Study Group of Liver Surgery (ISGLS)-criteria and the “peakBili > 7” criteria (see table 4)[[84-86](#_ENREF_84)].

Liver transplantation as only causal therapy of liver failure is often no option in these patients. Treatment must therefore focus on optimal perfusion of remnant liver, adequate infection prophylaxis and intensive care management of fluid balance, electrolytes, coagulation and kidney function.

***Ascites, infctions complications***

Ascites is a major problem after hepatic resections in cirrhotic patients with portal hypertension, even if no postoperative liver failure occurs. Ascites production often suspends after 5-7 d. Supportive therapies consist of diuretics (*e.g.* spironolactone), restrictive volume management and substitution of albumin. In cases of persisting ascites, a microbiological examination is recommended to detect and treat infections early. In individual patients, TIPS placement (alternatively: splenic artery embolization) must be considered.

The rate of infection-associated complications is significantly increased after liver resections in cirrhotic patients with portal hypertension compared to liver resections in otherwise healthy livers[[87](#_ENREF_87)]. Main foci here are pneumonia and superinfected ascites. In suspected infection, no delay in antibiotic treatment must be made. In cases of superinfected ascites, recommended empirical treatment is the use of third generation cephalosporines such as ceftriaxone or cefotaxime[[88](#_ENREF_88),[89](#_ENREF_89)], alternatively the use of carbapenems[[90](#_ENREF_90)]. Prior sampling and microbiological culture of the ascitic fluid is recommended for adequate adaption of the antibiotic regime depending on the antimicrobial resistance patterns. Control sampling and microbiologic culture of the ascitic fluid 48 h after start of antibiotic treatment is recommended.

***Wound healing and postoperative hemorrhage***

Due to the frequently underlying malnutrition and postoperative ascites, the rate of wound healing disorders in cirrhotic patients is significantly higher than in non-cirrhotic patients[[91](#_ENREF_91)]. Some reports indicate that abdominal closure with a running suture prevents wound dehiscence. Moreover, placement of an intraabdominal drainage to prevent fluid collection and to lower the intraabdominal pressure has been proposed although this is discussed controversially (see above).

Postoperative hemorrhage in cirrhotic patients can be caused by superficial wound bleeding, bleeding from the resection site as well as by GI bleedings. Rapid diagnostic assessment and interdisciplinary surgical-medical therapy including supplementation of coagulation products are essential in these potentially life-threatening complications.

***Results after liver resection in cirrhosis and portal hypertension***

A recent analysis of 2046 patients treated in 10 high-volume centres for liver surgery has shown that only 50% of patients were operated within the BCLC recommendations, i.e. BCLC stage 0-A and no portal hypertension. 36% and 14% of patients underwent liver resection in BCLC stage B and C, respectively[[42](#_ENREF_42)]. In this publication, 5-year overall survival rates after HCC resections in BCLC A/B/C cirrhosis were 61%, 57% and 38% and 5-year disease free survival of 21%, 27% and 18%. Thus, the authors conclude that liver resections for HCC can be performed even in higher degrees of cirrhosis with acceptable long-term outcome and that the BCLC/EASL/AASLD guidelines should thus be updated. Furthermore, several publications could show that portal hypertension is no prognostic factor for long-term survival or HCC recurrence in multivariate analyses[[25](#_ENREF_25),[34](#_ENREF_34)]. In contrast, Berzigotti and colleagues determined portal hypertension as an independent factor for decreased long-term survival and increased perioperative decompensation after HCC resection[[12](#_ENREF_12)]. However, since many cirrhotic patients diagnosed with HCC do not qualify for liver transplantation, taking an increased operative risk must carefully be discussed for and with each individual patient in the light of the potential long-term oncologic outcome.

**Surgical therapy of portal hypertension**

Only liver transplantation (LT) is the causal therapy for patients diagnosed with portal hypertension and advanced liver cirrhosis[[92](#_ENREF_92)]. However, not all cirrhotic patients are candidates for LT. In particular, patients suffering from severe complications from portal hypertension such as variceal hemorrhage and subsequent failure of conservative, endoscopic and angiographic interventions are referred to TIPS placement[[93](#_ENREF_93),[94](#_ENREF_94)]. Only if TIPS placement is technically not feasible, shunt surgery may be considered. Interestingly, several studies comparing TIPS and shunt surgery show a significantly reduced recurrence of hemorrhage and less occlusive complications for shunt surgery[[95](#_ENREF_95)]. However, due to the high perioperative morbidity and mortality, shunt surgery should only be considered for CPT A or well-compensated CPT B patients[[96](#_ENREF_96)]. Depending on the region of the portal-venous system requiring relief of vascular pressure and influenced by possible anastomotic sites, several shunting techniques have been developed (see table 5). If possible, only a partial bypass of the portalvenous flow is recommended to minimize the risk of encephalopathy.

More rarely used shunting techniques are the meso-caval side-to-side shunt (+/- interposition graft), the proximal end-to-side splenorenal shunt (Linton), the spleno-renal side-to-side shunt (Cooley), the coronario-cava- end-to-side shunt (Inokuchi) and the mesenterico-portal Rex-Shunt. Because shunt surgery is no longer a common practice, it should only be performed where there is sufficient expertise at specialized centers.

**Conclusion**

In conclusion, hepatic resections in cirrhotic patients with portal hypertension show a significantly increased mortality and morbidity. However, due to ongoing improvements in surgical technique and intensive care management, surgery is feasible in selected patients with adequate long-term outcome. Patients have to be carefully screened for the degree of liver cirrhosis (CPT A-C, MELD-score), grade of portal hypertension (thrombocyte count, splenomegaly, presence of collaterals and varices, elastography, HPVG-measurement) and comorbidities. To us, portal hypertension is not an absolute contraindication for liver resections in cirrhotic patients. A close interaction with anaesthesiology and the critical care team is mandatory for optimal perioperative management. Main indications for liver resections in cirrhotic patients are HCC, but also CCA and, rarely, hepatic metastases. Elective surgery should not only be restricted to patients with CPT A cirrhosis, but should also be offered to very well selected patients with more advanced cirrhosis. During surgery, abdominal access should be made carefully to preserve potential collaterals. Careful dissection to prevent intraoperative blood loss is necessary and the extent of liver resection should be restricted to a minimum. Moreover, the therapeutic alternative of local ablation (radiofrequency ablation, irreversible electroporation among others) by an experienced interventional radiologist should always be considered. In case of surgery, the Pringle-manoeuvre should be avoided but may help to bridge critical situations. During the postoperative period, close interdisciplinary management by the hepatobiliary surgical team, critical care specialists and hepatologists is warranted for optimized stabilisation of these critical patients and for early and appropriate management of complications.

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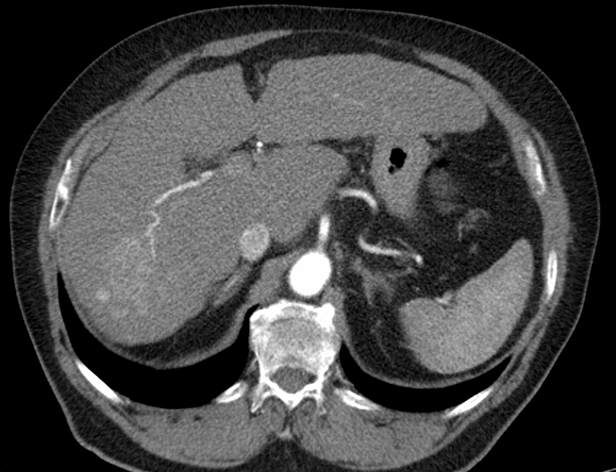
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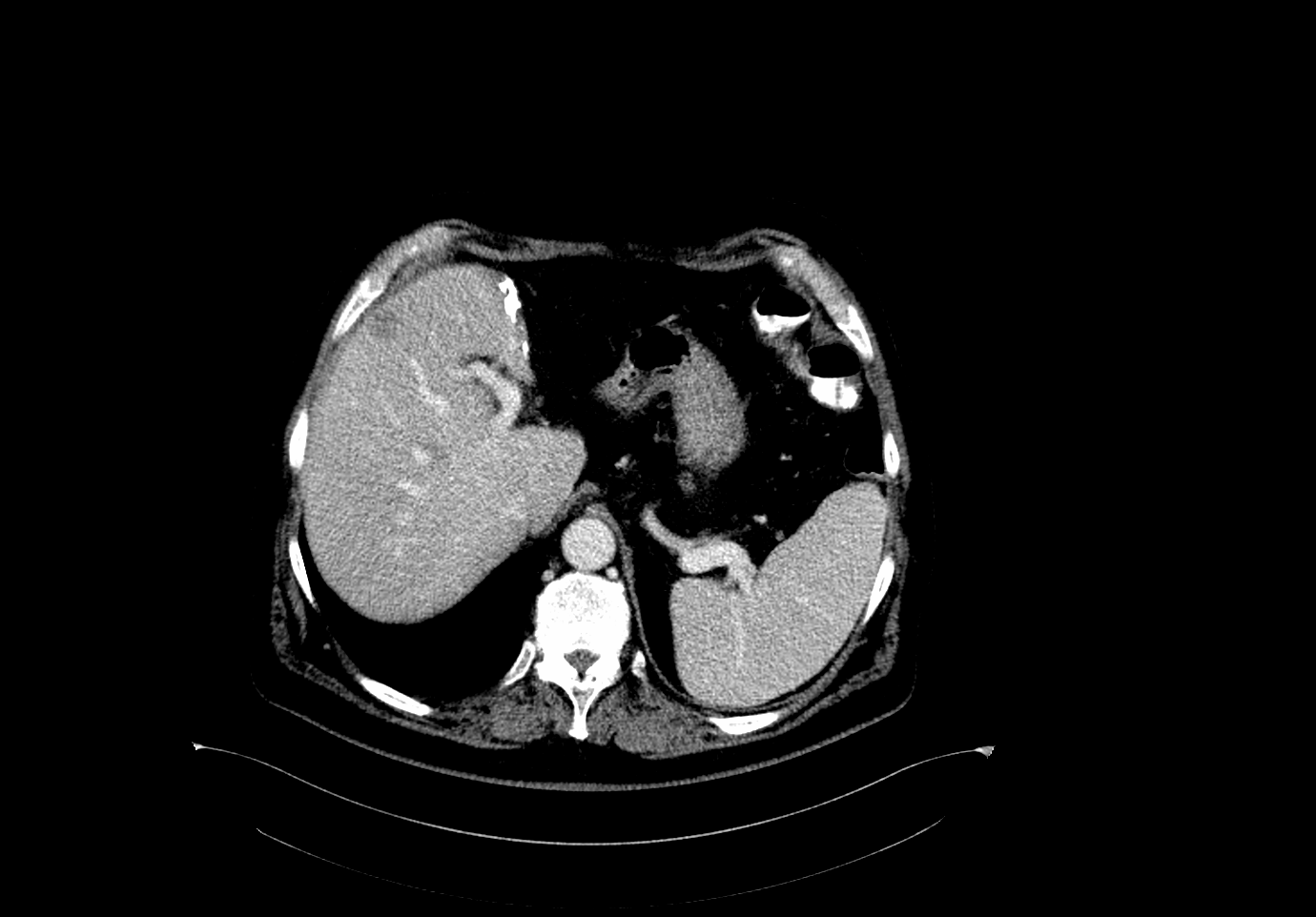
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**Figure 1 Patient with hepatocellular carcinomain macroscopic and biopsy proven liver cirrhosis but without signs of portal hypertension (CTP A, MELD 8, no varices, no splenomegaly, normal thrombocyte count) scheduled for right hepatectomy.**

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**Figure 2 Patient with liver metastases in histologcal proven liver cirrhosis with signs of portal hypertension (splenic diameter 15.3 cm, thrombocytopenia 77/nl, hypertensive gastropathy) scheduled for radiofrequency ablation.**

**Table 1 Causes of liver cirrhosis**

|  |
| --- |
| Alcoholic liver disease  Chronic viral hepatitis (hepatitis B and C)  Non-alcoholic fatty liver disease  Primary and secondary biliary cirrhosis  Primary sclerosing cholangitis  Hemochromatosis  Autoimmune hepatitis  Wilson’s disease  α1-Antitrypsin deficiency  Celiac disease  Right-sided heart failure  Granulomatous liver disease  Congenital malformation syndromes |

**Table 2 Most frequent causes of portal hypertension**

|  |
| --- |
| Prehepatic etiology of portal hypertension |
| Portal vein thrombosis |
| External portal vein compression |
| Intrahepatic etiology of portal hypertension |
| Hepatic cirrhosis (of any origin) |
| Congenital hepatic fibrosis |
| Schistosomiasis |
| Idiopathic non-cirrhotic portal hypertension |
| Posthepatic etiology of portal hypertension |
| Budd-Chiari’s syndrome |
| Sinusoidal obstruction syndrome |
| Cirrhose cardiaque |

**Table 3 Preoperative risk stratification models for patients with liver cirrhosis[**[**28**](#_ENREF_28)**-30,35-37]**

|  |  |  |  |  |
| --- | --- | --- | --- | --- |
| **Score** | **Groups** | **Initial function** | **Mortality assessment** | **Mortality after liver resection** |
| ASA | 1-6 | Perioperative risk stratification for any patient | Predictor of 7-d mortality | Not spefically defined |
| CPT | A-C | Overall survival in patients with liver cirrhosis | A: 10%, B: 30%, C: 80%; predictor for 30- and 90-d mortality | A: < 9%, no data for B and C |
| MELD | 0-40 | Mortality of TIPS-placement | 0-11: 5%-10%, 12%-25: 25%-54%, > 26: 55%-80%; predictor for 30- and 90-d mortality | ≤ 8: 0%; > 8: 29% |
| Mayo |  | Mortality after abdominal, orthopedic and cardiac surgery | 7-, 30-, 90-, 360-, 1800-d mortality | Not spefically defined |

ASA: American Society of Anaesthesiologists; CPT: Child-Pugh-Turcotte; MELD: Model of End Stage Liver Disease; TIPS: transjugular portosystemic stent shunt.

**Table 4 Definitions of postoperative liver-failure**

|  |  |  |
| --- | --- | --- |
|  | **Definition** | **Time of scoring** |
| „50-50“ | bilirubin > 50 µmol/L, prothrombin time < 50% (INR > 1.7) 🡪 mortality of 50%. | POD 5 |
| ISGLS | increased INR and hyperbilirubinemia | On or after POD 5 |
| ISGLS A | no intervention necessary |  |
| ISGLS B | non-invasive intervention necessary |  |
| ISGLS C | invasive intervention necessary |  |
| peakBili > 7 | maximum hyperbilirubinemia > 7 mg/dl any day after surgery - predictor for 90-d mortality | any POD |

POD: postoperative day; ISGLS: International Study Group of Liver Surgery.

**Tabele 5 Shunt surgery in portal hypertension and hepatic cirrhosis[93,95,96]**

|  |  |  |
| --- | --- | --- |
| **Shunt** | **Bypass of portalvenous flow** | **Recurrent haemorrhage/complications** |
| Porto-caval; end-to-side | complete | low rate of recurrent haemorrhage (< 5%)  low degree of shunt occlusion  40% encephalopathy  increase in aszites |
| Porto-caval, side-to-side +/- interposition graft | partial | recurrent haemorrhage 5%  low degree of shunt occlusion (5%)  5% encephalopathy |
| distal splenorenal shunt (Warren) | partial | recurrent haemorrhage 5%-8%  shunt occlusion 10%  selective decompression of gastroesophageal varices |