Liver Team:

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Liver Transplant OR Set-up and Preparation

General Game Plan:

1. General via endotracheal tube.
2. ALWAYS preoxygenate!
3. One adequately running peripheral IV is usually sufficient to start the case.
4. Rapid sequence or modified rapid sequence induction with Propofol and Rocuronium or Succinylcholine.
5. Maintenance with Des/ Iso with fentanyl and cisatracurium infusions
6. Consider premed with Midazolam if the patient is very anxious and has no signs of encephalopathy.

Overview of Setup

1. Medications to Order
2. Liver table and auxiliary equipment
3. Infusions
4. Back table medications/ induction
5. Airway
6. Lines
7. Monitors

1. Medications to Order
   - Order 3 medications from pharmacy as soon as you start setting up
     - Albumin 5%, 3L (500ml x6)
     - Octreotide 500mcg/vial. One vial for most livers, two for living donor
     - Insulin 1unit/ml, 100 unit bag
   - Ask your tech to pick up

2. Liver table and auxiliary equipment
• 8 syringe pumps, 1L NS carrier on micro-dripper, 8 stopcocks, extension, built by anesthesia techs  
• Blue top, Purple top tubes for CBC, Coags and lots of ABG syringes.

• 3 hot lines with Plasma-Lyte (2 on liver table, plus 3rd on IV pole to patient’s right)  
• Liver auxiliary cart should be in the room  
  o Contains extra medication, syringes, equipment for lines  
• Triple transducer  
• Zoll pads for defibrillator  
• Radiometer ABG machine—make sure it’s plugged in (lives in back room off of OR 12)  
• Ultrasound  
• Consider TEE probe

3. Infusions
• **Phenylephrine**: 10mg in 100ml NS to make 100mcg/ml. Draw up in a 60ml syringe for infusion. Set pump at 25mcg/min.  
  Also draw up a 10ml syringe for the back table.
• **Norepinephrine**: 4mg in 250ml NS to make 16mcg/ml. Draw up in 60ml syringe. Set pump at 5mcg/min.  
  Also draw up a 10 ml syringe for the back table
• **Calcium**: Draw up 3 vials of 100mg/ml in a 30ml syringe. Set infusion at 500mg/hr. Also draw up 10ml for back table.
• **Fentanyl**: 10ml of 50mcg/ml. Use for induction, then set pump at 100mcg/hr.  
• **Cisatracurium**: 10ml of 2mg/ml. Set infusion at 1mcg/Kg/min  
• **Octreotide**: Used to reduce Portal HTN. Each vial is usually 500mcg. Dilute 1 vial in 50ml NS to make 10mcg/ml. Set infusion at 100mcg/hr. Then go to options—bolus and set a 100mcg bolus over 15 mins. When the bolus is done, the pump will automatically start the infusion if you set it this way.

Variations:
• **Vasopressin 1unit/ml (20ml)**, for sicker patients, omit phenylephrine  
• **Furosemide 10mg/ml (10ml)**, for volume overload, CKD  
• Use Alaris caps on the syringes to make it easy to refill when needed.  
• The infusion line connects to the side port of introducer, VIP port or triple lumen catheter.
4. **Back table medications/induction**
   - **Antibiotic:** Zosyn 3.375g over 30 minutes, every 2 hours
   - **Methylprednisolone (Solumedrol)** 500mg IV over 30 minutes
     - In 100ml NS with minidripper, or run in syringe pump
     - **Give this thru a separate port since it tends to precipitate with other infusions.**
   - **Induction medications**
     - Propofol
     - Lidocaine
     - Rocuronium, rarely succinylcholine
     - Esmolol
     - Fentanyl
   - **Additionally:**
     - Epinephrine 10mcg/ml
     - Norepinephrine 16mcg/ml
     - Phenylephrine 100mcg/ml
     - Calcium 100mg/ml
     - Glycopyrrolate 0.2mg/ml
   - **Verify:**
     - Code-dose epinephrine 100mcg/ml
     - Ample supply of calcium chloride 100mg/ml
     - Vasopressin
     - Bicarbonate
     - Atropine

5. **Airway/Anesthesia machine**
   - Heated, humidified circuit
   - Working laryngoscope
   - Size 7.0 or 7.5 ETT
   - Oral airway
   - Tegaderm for eyes
   - Bite block
   - OG tube

6. **Lines**
   - Standard liver setup includes (usually in this order)
     - 20g left radial arterial line
     - 14g or 16g right upper extremity volume line
     - 7 French left antecubital rapid infusion catheter (RIC)
       - Placed with prep, sterile towels and gloves
8.5 or 9 French introducer with double-lumen insert or MAC catheter in right internal jugular (Typically, if the RIC goes in nicely, introducer is placed; otherwise MAC. Occasionally, trialysis catheter is placed, in which case RIC is usually not needed.)

7. Monitors
   • Standard ASA
     o Typically temp Foley is placed
   • A-line
   • CVP
   • Zoll pads
     o Turn on the defibrillator and check that you have ECG trace through the pads!
   • 3rd pressure channel occasionally used to check portal pressures (routine in living donor cases)

Starting the case:
   • 1 small reliable PIV for induction.
   • Position on table, most likely supine with right arm tucked and left arm out on arm board.
   • IV RSI as above
   • After induction, place left radial arterial 20G catheter. Left arm RIC 7.0 or 8.5 Fr
   • 14G or 16G PIV in R-arm before tucking
   • R-IJ Cordis or MAC depending on other access with or without PAC/TLC
   • Place 18Fr OG after all lines are in.
   • Temperature monitoring is usually done via a temperature foley
   • Temperature control: Heated, humidified circuits, 3 hotlines as above, Forced air warming (the circulator usually takes care of this, you won’t have to ask)
   • Labs: Call our tech for CBC, coags. ABGs are done in OR.
Goals and Objectives for Anesthesia Residents on the Liver Transplant Service

Introduction

Anesthesia residents at UCSF perform anesthesia on the liver transplant service as part of the senior resident rotation and during night call as the senior resident. The liver transplant service is a multidisciplinary service comprised of surgeon, hepatologists, anesthesiologists and nurses. Approximately 150 liver transplants are performed each year, providing each resident with roughly 8 transplants each.

Goals

• Describe the surgical course of a liver transplant.
• Understand the management of a donor hepatectomy.
• Understand the preoperative evaluation of a patient with cirrhosis
• Understand blood component therapy.

Objectives

Interpersonal Communication Skills

• Communicate effectively with other healthcare professionals.
• Demonstrate professionalism and interpersonal/communication skills with patients, families and children
• Communicate with patients and their families in easily understood and culture-sensitive language.
• Work effectively as a member of the liver transplant team
• Maintain comprehensive, timely, and legible medical records

Professionalism

• Demonstrate respect, compassion and integrity.
• Demonstrate a commitment to excellence and on-going professional development.
• Demonstrate a commitment to ethical principles pertaining to confidentiality of patient information, informed consent, and resource utilization.
• Demonstrate sensitivity and responsiveness to patients’ culture, age, gender, and disabilities.
• Demonstrate organizational skills to care for patients in a competent and efficient manner.

Medical Knowledge

Liver Specific
• List what vessels are “cross” clamped during the anhepatic stage
• Define veno-veno bypass.
• Define a “piggy back” transplant.
• Define the “anhepatic period”.
• Distinguish heterotopic transplant from orthotopic transplant.

Cirrhosis

• List the factors that are used to calculate a MELD score.
• List several factors used to determine Childs-Pugh classification.
• Define portopulmonary hypertension.
• Define hepatopulmonary syndrome.
• Define the approximate perioperative mortality in a patient with cirrhosis.
• List elements of coagulation abnormalities in a patient with cirrhosis.

Transfusion

• Define indications for Factor VIIa use.
• Define a dose range for Factor VIIa.
• Define the indication for use of FFP
• Define the indication for use of cryoprecipitate
• Define the indication for use of platelets.
• Define “massive transfusion.”
• Define dilutional coagulopathy and the hemostatic abnormalities associated with it.
• List blood components most associated with citrate toxicity.

Practice-based Learning and Improvement

• Perform literature search and retrieve relevant literature.
• Access UNOS website and relevant data.
• Access Licage and Litac and relevant data.

Systems-based Practice

• Demonstrate awareness of the role transplantation plays in the health care system.
• Understand efforts being made to expand organ donation, both living and deceased.
Preoperative Evaluation

The severity of illness and prognosis of patients with chronic liver disease can be estimated by a number of different scoring models including the Childs–Pugh–Turcotte score and the MELD score. The latter is now widely used in the United States for the allocation of organs. It is based on a predicted 3-month mortality for patients awaiting a liver transplant, and uses 3 laboratory values to generate a score, which determines priority. The three laboratory values used are serum bilirubin, serum creatinine, and INR. The format for the 2 systems are as follows:

<table>
<thead>
<tr>
<th>Variable</th>
<th>1 Point</th>
<th>2 Points</th>
<th>3 Points</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bilirubin (mg/dL)</td>
<td>&lt;2</td>
<td>2-3</td>
<td>&gt;3</td>
</tr>
<tr>
<td>Albumin (g/dL)</td>
<td>&gt;3.5</td>
<td>2.8-3.5</td>
<td>&lt;2.8</td>
</tr>
<tr>
<td>Encephalopathy grade</td>
<td>None</td>
<td>1-2</td>
<td>2-4</td>
</tr>
<tr>
<td>Ascites</td>
<td>Absent</td>
<td>Slight</td>
<td>Moderate</td>
</tr>
<tr>
<td>PT (sec prolonged)</td>
<td>&lt;4</td>
<td>4-6</td>
<td>&gt;6</td>
</tr>
</tbody>
</table>

Child-Pugh Score:
Class A: 5-6
Class B: 7-9
Class C: 10-15

MELD ([MELD Calculator](#))
MELD = 0.957 • Ln (Creatinine, mg/dl) + 0.378 • Ln (Bilirubin, mg/dl) + 1.120 • Ln (INR) + 0.643 • Ln (cause of cirrhosis)

UNOS has made the following modifications to the score:
If the patient has been dialyzed twice within the last 7 days, then the value for serum creatinine used should be 4.0
Patients with a diagnosis of liver cancer will be assigned a MELD score based on how advanced the cancer is.
Organ System-Specific Considerations

Neuro
Hepatic encephalopathy should be assessed.

In fulminant liver failure patients, you should look for signs of high ICP. Find out if the patient has had a CT scan looking for cerebral edema. Some patients may have ICP monitoring. Usually, these patients are intubated for coma.

Pulmonary
Gas exchange problems are common. While hepatopulmonary syndrome (intrapulmonary shunting from liver failure) may occur, most causes of gas exchange deficiencies are due to standard causes. These may include atelectasis, decrease FRC from ascites, pneumonia, underlying lung disease, and pleural effusions that are common in these patients. Often the effusions can be drained through the diaphragm after incision, so despite a bad effusion, the problem will not ultimately be severe during surgery.

Pulmonary Hypertension (Portopulmonary syndrome)
This is listed between the pulmonary and cardiac system, because it is a consideration for both organs, and is critically important. Cirrhosis is associated with pulmonary hypertension similar to primary pulmonary hypertension. The cause is not clear. However, these patients may do poorly, and severe enough forms may be a contraindication to transplantation. Pulmonary hypertension is difficult to screen for preoperatively, and can be asymptomatic. All patients will have had a cardiac echo, which should have estimated PA pressures. Even echo may be a poor screening tool, and PA catheters may be placed to specifically look for pulmonary hypertension. We are currently placing PA catheters only when clinically indicated.

Cardiac
Most patients have had some cardiac evaluation. Liver transplantation is extremely stressful surgery, and will not be tolerated by a patient with significant coronary artery disease. Stress tests, e.g. P-Thal, dobutamine echo, are routinely performed. There is a low threshold for getting left and right cardiac caths. MI’s have occurred in these patients. Cardiac rhythm problems are also common. Cardiac work-ups from patients referred from outside are usually found under “Care Everywhere” or “Scanned Outside Documents” in Apex or may need to be requested from the liver transplant service.
Evaluation of cardiac function is also essential. In the vasodilated state of liver failure, cardiac function should be hyperdynamic, with an elevated ejection fraction, often 70-80%. Even a mildly reduced ejection fraction, e.g. 55%, may be distinctly pathologic.

Another disease associated with liver failure with important cardiac considerations is hemochromatosis. Iron infiltration of the heart can impair cardiac contractility and conduction. Hemochromatosis is common, and because of additional diseases such as hepatitis C, it may be missed. In any adult patient with IDDM that developed after cirrhosis, hemochromatosis (bronze diabetes) should be suspected.

GI
History of GI bleeding, ascites, portal hypertension, TIPS performed should be noted. While NPO status is certainly ascertained, we consider that most patients may have full stomachs due to poor emptying with ascites, and possible GI bleeding. A Blakemore tube may be necessary during surgery.

Renal
Hepatorenal syndrome and other causes of renal insufficiency should always be assessed. Some patients may be scheduled for combined liver and kidney transplantation. Because large volumes of FFP may be necessary, some form of fluid removal (diuresis, or CVVH) may be necessary. Factor VIIa can also be used to achieve hemostasis in these patients, but discuss first with the surgical team, since factor VIIa has been associated with thrombosis.

Every electrolyte may be abnormal and should be considered. Hyperkalemia and hypokalemia are both common due to renal failure and diuretic use. Hyponatremia is common due to diuretic use, low intravascular volume and excessive free water retention and/or replacement. Although debated, central pontine myelinolysis may be related to over-rapid correction of hyponatremia, and has been reported in the setting of liver transplantation. Mg$^{2+}$ and ionized Ca$^{2+}$ levels should be noted and corrected prior to the anhepatic phase.

Heme
Obvious focus on coagulation factors, hemoglobin level and platelet count are important. Blood is set up by protocol, but the order to thaw FFP needs to be sent to the blood bank, and in a patient with a severe coagulopathy, factors need to be requested for possible transfusion before line placement. This should be anticipated during the preop evaluation. We currently order 5 RBC, 5 FFP and 2 plt in room before start of the case. More products can be ordered in patients who are severely coagulopathic pre-operatively. Talk to your attending.
The Operation: General Considerations
Liver transplantation is almost always performed as an "orthotopic liver transplantation", meaning the transplanted liver is placed in the same position as the native liver (compare with renal transplantation where the kidney is transplanted into a heterotopic location, the pelvis). Auxiliary liver transplantation is a technique that may be used in the future in the setting of fulminant liver failure. In our pediatric patients, our surgeons use a variety of "reduced-size" grafts: pare-down segments from adult donors, small segments (left lateral segment) from live donors, or "split-livers" - two grafts from one donor. Adult-to-adult living related transplants use donor right hepatic lobe. A "piggy-back" refers to partial occlusion of the IVC, with an end-to-side anastomosis, instead of insertion of donor IVC. However, regardless of the surgical technique, liver transplantation involves an abdominal dissection, removal of the native liver, placement of the liver graft, "reperfusion" and additional work on the graft, and then closure. For a variety of studies and discussions, we break the operation down into three stages. Breaking the procedure into these stages will permit a better delineation of specific anesthetic goals because of the changing physiology.

The pictures below show the difference in anastomoses between a conventional bicaval technique versus a piggyback technique.

![Conventional Technique vs Piggyback Technique](image)

[Nature Reviews Gastroenterology & Hepatology 10, 434-440 (July 2013)]

Living Donor Liver Transplant Recipients:
The set up and preoperative evaluation are the same as for cadaveric transplants. For a nice review, see this [NEJM article](https://www.nejm.org/).
Stage I: "DISSECTION Phase"
This will consist of incision, exposure, dissection, and recipient hepatectomy. In another context, consider it major abdominal surgery in the presence of coagulopathy, portal hypertension, and frequently severe scar tissue from prior hepatobiliary surgery or from post-necrotic hepatitis. The goal of this portion of the surgery is to mobilize the vascular structures around the liver (suprahepatic vena cava, infrahepatic vena cava, portal vein, and hepatic artery) and isolate the common bile duct. In addition, adhesions between the liver, diaphragm and retroperitoneal areas must be dissected to allow full mobilization of the liver within the right upper quadrant. Be wary of the effects of hepatic manipulation on venous return: i.e., surgical induced hypotension. This must be distinguished from other causes of hypotension, which are all too common during liver transplantation.

Toward the end of this stage, we prepare for the Anhepatic period by a technique of volume loading if a bicaval clamp is planned - administration of colloid solution and/or blood (titrated to filling pressure, e.g. CVP of 10-15 mmHg) in anticipation of a significant decrease in filling pressures after the caval clamps are applied. However, until just before cross clamp, low CVP is preferred (≈5 mmHg if tolerated) to reduce blood loss. In addition, a general goal during the dissection phase of the operation is to promote a diuresis. Diuresis is usually necessary to allow transfusion of sufficient FFP without creating excessive volume overload and hemodilution.

Stage II: "ANHEPATIC Phase"
This is the period of time when no liver is in the circulation. At UCSF, this is nearly always managed without the use of venovenous bypass, and consequently, this stage has significant metabolic and cardiovascular implications. This is the most challenging part of the anesthetic management.

The hemodynamic considerations depend on the planned surgical procedure. The bicaval technique is more challenging from a hemodynamic standpoint and will be described below:

The stage begins with the applications of multiple vascular clamps: 1) Portal Vein 2) Hepatic Artery 3) Infrahepatic Vena Cava 4) Suprahepatic Vena Cava. The mobilized liver is quickly removed, and the ice-cold liver graft is placed in the field. During the next 30-45 minutes, 3 anastomoses are completed: 1) Suprahepatic vena cava (right beneath the diaphragm, look over and see the hepatic veins draining into the cava of the new liver), 2) Infrahepatic vena cava, and 3) Portal vein - a smaller vessel but critical for providing flow into the liver. Fluid management with the caval clamps on is extremely difficult, and we must anticipate the return of volume when the clamps are released at the completion of the vascular anastomoses. This must be balanced with use of pressors and inotropes. Currently, a staged-release is performed: the suprahepatic caval clamp is released and the anastomosis is inspected and small leaks
repaired. The infrahepatic caval clamp is then released and the anastomosis is inspected and repaired if needed. The filling pressures will normally reflect the establishment of caval continuity. Finally, the portal vein clamp is removed allowing recipient blood flow into the donor liver. At this point the anhepatic period ends and the next phase begins.

**Stage III: "REPERFUSION Phase"

This covers the time from vascular reperfusion of the liver graft until the end of the procedure. The actual moment of reperfusion can have profound cardiovascular effects (i.e. arrhythmias, profound hypotension, and about a 3-5% incidence of hyperkalemic or other cardiac arrest) and anesthetic management is directed to maintaining or recovering cardiovascular stability. Multiple factors probably combine to produce this hemodynamic insult: ionic composition of the preservative solution, hypothermia, metabolic acidosis, "vasoactive" peptides from the gut, other cytokines, sudden atrial stretching in response to the unclamping and reperfusion. Initially, profound bradycardia usually occurs, which can become asystole. This can be treated by glycopyrrolate or atropine, but we usually use epinephrine in small doses, 10-30 µg at a time. After the initial bradycardia, the hypotension appears to be overwhelming vasodilatation (very low SVR) with adequate filling pressures and high cardiac output (15-20 L/min).

Following reperfusion, attention is paid to the diagnosis and management of a significant coagulopathy (dilution/consumption of clotting factors, platelet entrapment, heparin effect, primary fibrinolysis), and resultant bleeding. Platelets should be administered to correct thrombocytopenia. Most of the necessary treatment is FFP. Factor VIIa can be used in patients with significant volume overload, although it may be best to wait some time until fibrinolysis resolves. Antifibrinolytics (Amicar) can be given if fibrinolysis is suspected. Protamine can be administered to treat any heparin effect. Factor VIIa, Amicar, and protamine should be discussed with your attending and the surgeons prior to use. Laboratory analysis and blood product support guide the management of the coagulopathy. Bleeding tends to be of mixed, multifactorial etiology: coagulopathy as mentioned above, extensive raw surfaces above and behind the liver, effects of portal hypertension, and multiple anastomoses. In addition, we monitor evolution and evaluation of liver graft function (PT, base deficit), and recovery from renal ischemia. During this time the surgeons are completing the hepatic artery anastomosis, controlling bleeding, performing a cholecystectomy on the liver graft, and providing for biliary drainage via an end-to-end (duct-to-duct) anastomosis or a choledochojejunostomy with a Roux-en-Y loop.

Toward the end of the procedure, we shift the anesthetic emphasis to consider postoperative disposition. One option is to provide analgesia and sedation in anticipation of postoperative mechanical ventilation. If the liver is functioning adequately, and the patient is hemodynamically stable with adequate gas exchange,
tracheal extubation should be attempted. This has been very successful in a high percentage of patients. Hypertension may develop toward the end of the procedure in about 20-30% of cases. Relative hypervolemia, activation of endogenous reflexes during the anhepatic period, and possibly activation of the renin/angiotensin or endothelin systems probably combine to produce the relative hypertension. It is important to identify a tendency to hypertension before leaving the operating room and provide treatment since hypertension could promote bleeding at the arterial anastomosis.

**Intraoperative Hemodynamics: A General Depiction**

During the liver transplant procedure a general pattern of hemodynamic changes usually evolve in each patient. This typically reflects the common cardiovascular changes that accompany chronic liver disease, and the manifestations of "reflex responses" to the anhepatic period, and the acute effects of reperfusion as described above. Below, data gathered in one patient during the critical part of the operation are displayed: end of dissection phase, anhepatic period and reperfusion.

Please understand that this is the data from one patient, a 60-year-old woman with primary biliary cirrhosis, with the usual manifestations of end stage liver disease. These values are representative for a typical patient undergoing liver transplantation. Note that the initial cardiac output/index is evidence of a hyperdynamic circulation associated with a decreased systemic vascular resistance. During the anhepatic period, marked reductions in filling pressures and cardiac index do not result in systemic arterial hypotension. The ability to increase the SVR three or four fold results in a relatively stable blood pressure. Not all patients will demonstrate the same reflex responses to the anhepatic period, and support of blood pressure may require additional volume and/or pressor support. Not evident in this data are the acute and transient changes that occur upon reperfusion of the liver graft. Despite a sudden increase in central filling pressures, a period of hypotension (2-3 minutes) usually accompanies graft reperfusion, and usually requires immediate intervention (consider: prophylactic calcium, blood pressure support with phenylephrine or epinephrine.)
<table>
<thead>
<tr>
<th>Stage</th>
<th>HR</th>
<th>MAP</th>
<th>CVP</th>
<th>PA</th>
<th>CO</th>
<th>CI</th>
<th>SVRI</th>
</tr>
</thead>
<tbody>
<tr>
<td>Dissection (II-5)</td>
<td>84</td>
<td>78</td>
<td>5</td>
<td>19</td>
<td>9.9</td>
<td>5.8</td>
<td>589</td>
</tr>
<tr>
<td>Anhepatic (II+10)</td>
<td>99</td>
<td>72</td>
<td>2</td>
<td>4</td>
<td>3.8</td>
<td>2.2</td>
<td>1472</td>
</tr>
<tr>
<td>Anhepatic (III-5)</td>
<td>96</td>
<td>78</td>
<td>5</td>
<td>5</td>
<td>4.3</td>
<td>2.5</td>
<td>1356</td>
</tr>
<tr>
<td>Reperfusion (III+10)</td>
<td>86</td>
<td>72</td>
<td>7</td>
<td>21</td>
<td>11.5</td>
<td>6.7</td>
<td>452</td>
</tr>
<tr>
<td>Reperfusion (III+30)</td>
<td>91</td>
<td>76</td>
<td>8</td>
<td>22</td>
<td>10.0</td>
<td>5.8</td>
<td>543</td>
</tr>
</tbody>
</table>
Anesthesia for Liver Transplantation: The Basic Game Plan

Unlike elective operations, we can never say who we will be transplanting or when the case will start. As mentioned earlier, the cases will be scheduled when a donor liver has been identified. Because of multiple logistic considerations regarding the procurement of the donor liver, do not be surprised if the scheduled "Start" time is changed. It is important to discuss the various risks with the patient, and specific mention should be made of transfusion and post-operative intubation. The anesthesia attending should complete this discussion with the patient.

As with any operation, there is no ideal anesthetic - there have been no controlled trials comparing different anesthetic agents or techniques during liver transplantation. Anesthetic induction can be performed with Propofol. Desflurane, isoflurane, or Sevoflurane for maintenance is adequate, with opioid (usually fentanyl). Any reasonable anesthetic technique is possible, but must take into consideration the intricacies of intercurrent illness (i.e. hepatic failure to some extent) and surgical requirements (in this case management of the anhepatic phase without the use of venovenous bypass). Central to our anesthetic management is preparation and anticipation: at critical points of the operation we look for and expect certain responses from the patient, and are ready to intervene during those critical moments, mostly during the anhepatic phase and with graft reperfusion.

An arterial line is absolutely essentially. This is usually placed in the left radial artery, because this arm can remain out, while the right arm is tucked. This leads to many fewer problems. Femoral arterial lines can be necessary. A right IJ 8.5 or 9 Fr introducer should be placed, with a PA catheter if indicated, or alternatively with a DLIC. A left antecubital RIC (7.0 Fr) is the preferred volume line, which can be used by the rapid infusion device (Belmont).

Laboratory and Blood Sample Protocol

There is no shortage of intraoperative laboratory tests during this operation. In general, arterial blood samples are sent at baseline and serially during the operation. During critical phases of the operation, labs are usually sent every 30 minutes.

We now have the ability to do point of care ABG analysis, which has eliminated problems with delays. This also provides complete electrolytes, glucose and lactate. In addition, we expect to measure hemoglobin with a Hemocue. CBC and coags are sent to the main laboratory.
Fluid Management

It is fairly common to expect a blood loss of 1-1.5 blood volumes (i.e. 5-8 liters), and the total intravenous fluids administered will be 15-20 liters. This fluid will consist of crystalloid, colloids, blood products, and "rapid infusion" support. The rapid transfusion device is an important part of the anesthetic management and support of the liver transplant patient. At UCSF, we have a technician available for all of our cases, and following the induction of anesthesia, they can prepare 1-2 liters of modified whole blood consisting of washed packed red blood cells (washed to remove K+ and acid) reconstituted with fresh frozen plasma. This solution is administered via a Belmont, usually through a left antecubital 7.0 Fr RIC. We are usually able to obtain flow rates of 300-600 cc/minute, and the perfusionist will administer this volume under the direction of the anesthesia team.

Standard blood products are set up for liver transplant. This includes 5 or 10 u PRBC and 5 or 10 u FFP to be brought to the OR, with 2 or 4 single donor platelet units available on request. Thawing of the FFP is delayed until the start of the transplant is confirmed. The circulating nurse may ask you about thawing FFP. If there are any questions about blood banking issues, refer them to your attending. Blood products are brought to the OR in coolers, and the platelets in a soft blue insulating package. The nurses and perfusionist will check these in for us.
**Anesthetic Record**

There is a "Liver Transplant protocol" on Apex that will insert an (almost) appropriate collection of drugs and fluids for a liver transplant.

Currently Apex does not track “Units” of blood, so it may be easier to write a standard volume for all units, such as 301 ml, which makes reporting units much easier. Please enter unit numbers. Blood from the perfusionist can be totaled at the end, and a memo referring to their record for unit numbers can be made.

Blood gas data should automatically transmit to the anesthesia record.

**Other Useful Links:**

[Licage](#)

[UNOS](#)

[CTDN](#)

[Scientific Registry of Transplant Recipients](#)
Acknowledgements

The original "handbook" has evolved continuously since the first liver transplant at UCSF in 1988, and was created by former members, Jeffrey White, Denna Washington and Scott Kelley.

The online version was created by webmaster John Feiner.

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