

UNIVERSITY MEDICAL CENTE Lubbock, Texas

Patient Label Here

DISCLOSURE AND CONSENT - MEDICAL AND SURGICAL PROCEDURES

TO THE PATIENT: You have the right as a patient to be informed about your condition and the recommended surgical, medical or diagnostic procedure to be used so that you may make the decision whether or not to undergo the procedure after knowing the risks and hazards involved. This disclosure is not meant to scare or alarm you; it is simply an effort to make you better informed so you may give or withhold your consent to the procedure.

1. I (we) voluntarily request Doctor(s) ______as my physician(s), and such associates, technical assistants and other health care providers as they may deem necessary, to treat my **condition** which has been explained to me (us) as (**lay terms**): Pressure caused by a blockage in the blood flow throughout the liver

2. I (we) understand that the following surgical, medical, and/or diagnostic **procedures** are planned for me and I (we) voluntarily consent and authorize these **procedures** (**lay terms**): <u>Transjugular Intrahepatic</u> Portosystemic Shunt (TIPS) - tube placed in the middle of the liver to reroute the blood flow

Please check appropriate box: Right Left Bilateral Not Applicable

3. I (we) understand that my physician may discover other different conditions which require additional or different procedures than those planned. I (we) authorize my physician, and such associates, technical assistants, and other health care providers to perform such other procedures which are advisable in their professional judgment.

4. Please initial _____Yes____No

I consent to the use of blood and blood products as deemed necessary. I (we) understand that the following risks and hazards may occur in connection with the use of blood and blood products:

- a. Serious infection including but not limited to Hepatitis and HIV which can lead to organ damage and permanent impairment.
- b. Transfusion related injury resulting in impairment of lungs, heart, liver, kidneys and immune system.
- c. Severe allergic reaction, potentially fatal.

5. I (we) understand that no warranty or guarantee has been made to me as to the result or cure.

6. Just as there may be risks and hazards in continuing my present condition without treatment, there are also risks and hazards related to the performance of the surgical, medical, and/or diagnostic procedures planned for me. I (we) realize that common to surgical, medical and/or diagnostic procedures is the potential for infection, blood clots in veins and lungs, hemorrhage, allergic reactions, and even death. I (we) also realize that the following hazards may occur in connection with this particular procedure: Pain, bleeding, infection, injury to or occlusion (blocking) of artery which may require immediate surgery or other intervention, damage to parts of the body supplied by the artery with resulting loss of use or amputation (removal of body part), worsening of the condition for which the procedure is being done, stroke and/or seizure (for procedures involving blood vessels supplying the spine, arms, neck or head), contrast-related temporary blindness or memory loss (for studies of the blood vessels of the brain), paralysis (inability to move) and inflammation of nerves (for procedures involving blood vessels supplying the spine), contrast nephropathy (kidney damage due to the contrast agent used during procedure), thrombosis (blood clot forming at or blocking the blood vessel) at access site or elsewhere, hepatic encephalopathy (confusion/decreased ability to think), liver failure or injury, gallbladder injury, hemorrhage, recurrent ascites (fluid building up in abdomen) and/or bleeding, kidney failure, heart failure, death, failure of procedure or injury to blood vessel requiring stent (small permanent tube placed in blood vessel to keep it open) placement or open surgery, change in procedure to open surgical procedure, failure to place stent/endoluminal graft (stent with fabric covering it), stent migration (stent moves from location in which it was placed, vessel occlusion (blocking)



7. I (we) understand that Do Not Resuscitate (DNR), Allow Natural Death (AND) and all resuscitative restrictions are suspended during the perioperative period and until the post anesthesia recovery period is complete. All resuscitative measures will be determined by the anesthesiologist until the patient is officially discharged from the post anesthesia stage of care.

8. I (we) authorize University Medical Center to preserve for educational and/or research purposes, or for use in grafts in living persons, or to otherwise dispose of any tissue, parts or organs removed except: <u>NONE</u>

9. I (we) consent to the taking of still photographs, motion pictures, videotapes, or closed circuit television during this procedure.

10. I (we) give permission for a corporate medical representative to be present during my procedure on a consultative basis.

11. I (we) have been given an opportunity to ask questions about my condition, alternative forms of anesthesia and treatment, risks of non-treatment, the procedures to be used, and the risks and hazards involved, potential benefits, risks, or side effects, including potential problems related to recuperation and the likelihood of achieving care, treatment, and service goals. I (we) believe that I (we) have sufficient information to give this informed consent.

12. I (we) certify this form has been fully explained to me and that I (we) have read it or have had it read to me, that the blank spaces have been filled in, and that I (we) understand its contents.

IF I (WE) DO NOT CONSENT TO ANY OF THE ABOVE PROVISIONS, THAT PROVISION HAS BEEN CORRECTED.

I have explained the procedure/treatment, including anticipated benefits, significant risks and alternative therapies to the patient or the patient's authorized representative.

	A.M. (P.M.)				
Date	Time	Printed name of provide	er/agent	Signature of provid	der/agent
	A.M. (P.M.)				
Date	Time				
*Patient/Other le	egally responsible person signature		Relationship	(if other than patient)	
*Witness Signat	ure		Printed Name	e	
GI & Out	2 Indiana Avenue, Lubbock, TX patient Services Center 10206 (alth & Wellness Hospital 1101) dress:	Quaker Ave, Lubbock	x TX 79424	treet, Lubbock, T	X 79430
	Address (Street or P.	O. Box)		City, State, Zip Co	ode
Interpretatio	n/ODI (On Demand Interpretin	g) 🗆 Yes 🔲 No	Date/Time	(if used)	
Alternative f	forms of communication used	\Box Yes \Box No	Printed nar	ne of interpreter	Date/Time
Date proced	ure is being performed:				
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Resident and Nurse Consent/Orders Checklist

Instructions for form completion

Note: Enter "not applicable" or "none" in spaces as appropriate. Consent may not contain blanks.

- Section 1: Enter name of physician(s) responsible for procedure and patient's condition in lay terminology. Specific location of procedure must be indicated (e.g. right hand, left inguinal hernia) & **may not be abbreviated.**
- Section 2: Enter name of procedure(s) to be done. Use lay terminology.
- Section 3: The scope and complexity of conditions discovered in the operating room requiring additional surgical procedures should be specific to diagnosis.
- Section 5: Enter risks as discussed with patient.
 - A. Risks for procedures on List A must be included. Other risks may be added by the Physician.
 - B. Procedures on List B or not addressed by the Texas Medical Disclosure panel do not require that specific risks be discussed with the patient. For these procedures, risks may be enumerated or the phrase: "As discussed with patient" entered.
- Section 8: Enter any exceptions to disposal of tissue or state "none".
- Section 9: An additional permit with patient's consent for release is required when a patient may be identified in photographs or on video.

Provider Attestation:	Enter date, time, printed name and signature of provider/agent.
Patient Signature:	Enter date and time patient or responsible person signed consent.
Witness Signature:	Enter signature, printed name and address of competent adult who witnessed the patient or authorized person's signature
Performed Date:	Enter date procedure is being performed. In the event the procedure is NOT performed on the date indicated, staff must cross out, correct the date and initial.

If the patient does **not** consent to a specific provision of the consent, the consent should be rewritten to reflect the procedure that the patient (authorized person) is consenting to have performed.

For additional information on informed consent policies, refer to policy SPP PC-17.

Consent

		No blanks left on consent No medical abbreviations Orders
Procedure Date Procedure	Procedure	Orders
Orders		No blanks left on consent No medical abbreviations

THIS FORM IS NOT PART OF THE MEDICAL RECORD