## Pediatric Liver Transplant Recipient Registration Worksheet

Note: These worksheets are provided to function as a guide to what data will be required in the online  $TIEDI^{(0)}$  application. Currently in the worksheet, a red asterisk is displayed by fields that are required, independent of what other data may be provided. Based on data provided through the online  $TIEDI^{(0)}$  application, additional fields that are dependent on responses provided in these required fields may become required as well. However, since those fields are not required in every case, they are not marked with a red asterisk.

Recipient Information		
Name:	DOB:	
SSN:	Birth sex:	
HIC:	Transplant Date and Time:	
State of Permanent Residence: *		
Permanent Zip: *		
Provider Information		
Recipient Center:		
Surgeon Name: *		
NPI#:*		
· · · · · · · · · · · · · · · · · · ·		
Donor Information UNOS Donor ID #:		
Recovering OPO:		
Donor Type:		
Detiont Status		
Patient Status Primary Diagnosis: *		
Specify:		
Date: Last Seen, Retransplanted or Death *		
Patient Status: *	CLIVING	
	ODEAD	
	RETRANSPLANTED	
Primary Cause of Death:		
Specify:		
Contributory Cause of Death:		
Specify:		
Contributory Cause of Death:		
Specify:		
Transplant Hospitalization:		
Date of Admission to Tx Center:*		
Date of Discharge from Tx Center:		
		ì
Clinical Information : PRETRANSPLANT Medical Condition at time of transplant: *	IN INTENSIVE CARE UNIT	
	HOSPITALIZED NOT IN ICU	
	ONOT HOSPITALIZED	
Patient on Life Support: *	⊖yes ⊖no	
	□ Ventilator	
Specify:	Other Mechanism, Specify	
Functional Status: *		

	_	
Cognitive Development: *	Definite Cognitiv	ive delay/impairment
	Probable Cognit	tive delay/impairment
	Questionable Co	ognitive delay/impairment
	No Cognitive del	elay/impairment
	ONot Assessed	
Motor Development: *	Definite Motor d	delay/impairment
	Probable Motor	delay/impairment
	Questionable Mo	otor delay/impairment
	ONo Motor delay/	/impairment
	Not Assessed	
Academic Progress:*	Within One Grad	de Level of Peers
	Delayed Grade L	Level
	Ospecial Education	on
	Not Applicable, f	too young for school/ High School graduate or GED
	Status Unknown	n Gol
Academic Activity Level: *	Full academic lo	bad
	Reduced academ	mic load
	Ounable to partic	cipate in academics due to disease or condition
	Ounable to partic	cipate regularly due to dialysis
	Not Applicable,	too young for school/ High School graduate or GED
	Status Unknown	
Source of Payment:		
Primary: *		
Specify:		
Height Measurement Date:		
Height: *	ft. in.	cm ST=
Weight Measurement Date:		
Weight: *	lbs	kg ST=
BMI:	kg/m <sup>2</sup>	
Previous Transplants:		
Previous Transplant Organ	Previous Transplant Date	Previous Transplant Graft Fail Date
The three most recent transplants are list emailing unethelpdesk@unos.org.	ed here. Please contact the UNet Help Desk to c	confirm more than three previous transplants by calling 800-978-4334 or by
Viral Detection:		
HIV Serostatus: *	Positive	
	Negative	
	Not Done	
	OUNK/Cannot Dis	sclose
CMV Status*	Positive	
	Negative	
	Not Done	
	UNK/Cannot Dis	sclose
HBV Surface Antibody Total *	Positive	
	Not Done	_
	<b>UNK/Cannot Dis</b>	sciose

HBV Core Antibody: *	Positive	
	Negative	
	ONot Done	
	UNK/Cannot Disclose	
HBV Surface Antigen: *	Positive	
	ONegative	
	ONot Done	
	UNK/Cannot Disclose	
HCV Serostatus: *	Positive	
	ONegative	
	○Not Done	
	UNK/Cannot Disclose	
EBV Serostatus: *	Positive	
	Negative	
	ONot Done	
	UNK/Cannot Disclose	
Vaccination Status:		
Did the recipient receive Hepatitis B vaccines prior to transplant?: $*$		
Reason not vaccinated:	Immunity	
	Medical precaution	
	CTime constraints	
	OPatient objection	
	Product out of stock	
	Other, specify	
Specify:		
NAT Results:		
HIV NAT: *	Positive	
	Negative	
	Not Done	
	UNK/Cannot Disclose	
HBV NAT: *	Positive	
	Negative	
	Not Done	
	UNK/Cannot Disclose	
HCV NAT: *	Positive	
	Onegative Charles Deers	
	Not Done UNK/Cannot Disclose	
	Owk/ Cannot Disclose	
Has the recipient ever had a diagnosis of HCC?* YES	NO	
Clinical Information : TRANSPLANT PROCED	lire	

Multiple Organ Recipient Were extra vessels used in the transplant procedure:

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Procedure Type:	OWhole Liver				
	Partial Liver, remainder not Tx or Living Transplant				
	Split Liver				
	OWhole Liver with Pancreas (Technical Reasons)				
	Partial Liver with Pancreas (Technical Reasons)				
	Split Liver with Pancreas (Technical Reasons)				
Split Type:					
Preservation Information:					
Total Cold Ischemia Time (if pumped, include pump	hrs ST=				
time):*	1115 <b>31</b> =				
Risk Factors:					
Previous Abdominal Surgery: *					
Portal Vein Thrombosis: *					
Transjugular Intrahepatic Portosystemic Shunt: *					
Organ Check-in					
Information:					
Liver Check-In Date: Time:	Military time Time Zone:				
Clinical Information : POST TRANSPLANT					
Pathology Conf. Liver Diag. of Hospital Discharge: *					
Specify:					
Graft Status: *	Functioning Failed				
If death is indicated for the recipient, and the death was a result o Date of Graft Failure:	f some other factor unrelated to graft failure, select Functioning.				
Causes of graft failure:					
Primary Non Function					
Hepatic Artery Thrombosis	YES NO UNK				
Other Vascular Thrombosis					
Portal Vein Thrombosis:					
Hepatic Outflow Obstruction:					
Diffuse Cholangiopathy					
Hepatitis: DeNovo					
Hepatitis: Recurrent	YES NO UNK				
Recurrent Disease (non-Hepatitis)					
Acute Rejection					
Infection					
Other, Specify:					
Did patient have any acute rejection episodes between transplant and discharge: *	Yes, at least one episode treated with anti-rejection agent				
	Yes, none treated with additional anti-rejection agent				
	○No				
Immunosuppressive Information Are any medications given currently for maintenance or					
anti-rejection: *					

Immunosuppressive Medications Copyright © 2023 United Network for Organ Sharing. All rights reserved. OPTN use only. 091423 View Immunosuppressive Medications

## **Definitions Of Immunosuppressive Medications**

For each of the immunosuppressive medications listed, select Ind (Induction), Maint (Maintenance) or AR (Anti-rejection) to indicate all medications that were prescribed for the recipient during the initial transplant hospitalization period, and for what reason. If a medication was not given, leave the associated box(es) blank.

Induction (Ind) immunosuppression includes all medications given for a <u>short finite period</u> in the perioperative period for the purpose of preventing acute rejection. Though the drugs may be continued after discharge for the first 30 days after transplant, it <u>will not</u> be used long-term for immunosuppressive maintenance. Induction agents are usually polyclonal, monoclonal, or IL-2 receptor antibodies (example: methylprednisolone, Campath, Thymoglobulin, or Simulect). Some of these drugs might be used for another finite period for rejection therapy and would be recorded as anti-rejection therapy if used for this reason. For each induction medication indicated, write the <u>total number of days the drug was actually administered</u> in the space provided. For example, if Simulect was given in 2 doses a week apart then the total number of days would be 2, even if the second dose was given after the patient was discharged.

Maintenance (Maint) includes all immunosuppressive medications given before, during or after transplant with the intention to maintain them long-term (example: prednisone, cyclosporine, tacrolimus, mycophenolate mofetil, azathioprine, or Rapamune). This does not include any immunosuppressive medications given to treat rejection episodes, or for induction.

Anti-rejection (AR) immunosuppression includes all immunosuppressive medications given for the purpose of treating an acute rejection episode during the initial post-transplant period or during a specific follow-up period, usually up to 30 days after the diagnosis of acute rejection (example: methylprednisolone, or Thymoglobulin). When switching maintenance drugs (example: from tacrolimus to cyclosporine; or from mycophenolate mofetil to azathioprine) because of rejection, the drugs <u>should not</u> be listed under AR immunosuppression.

If an immunosuppressive medication other than those listed is being administered (e.g., new monoclonal antibodies), select Ind, Maint, or AR next to Other Immunosuppressive Medication field, and enter the full name of the medication in the space provided. Do not list non-immunosuppressive medications.

Drug used for induction, acute rejection, or maintenance					
Steroids (prednisone, methylprednisolone, Solumedrol, Medrol)	Ind.	Days	ST	Maint	AR
Drugs used for induction or acute rejection					
Atgam	Ind.	Days	ST	Maint	AR
Campath (alemtuzumab)					
Cytoxan (cyclophosphamide)			B		
Methotrexate (Folex PFS, Mexate-AQ, Rheumatrex)					
Rituxan (rituximab)					
Simulect (basiliximab)					
Thymoglobulin					
	0				
Drugs primarily used for maintenance	Ind.	Days	ST	Maint	AR
Cyclosporine, select from the following:					
Gengraf					
- Neoral					
- Sandimmune					
- Generic cyclosporine					
Imuran (azathioprine, AZA)					
Leflunomide (LFL)					
Mycophenolic acid, select from the following:					
- CellCept (MMF)					
- Generic MMF (generic CellCept)					
- Myfortic (mycophenolic acid)					
- Generic Myfortic (generic mycophenolic acid)					
mTOR inhibitors, select from the following:					
- Rapamune (sirolimus)					
- Rapantune (siroinnus)					
- Generic sirolimus					

Tacrolimus, select from the following:					
- Astagraf XL (extended release tacrolimus)					
- Envarsus XR (tacrolimus XR)					
- Prograf (tacrolimus)					
- Generic tacrolimus (generic Prograf)					
Other drugs					
	Ind.	Days	ST	Maint	AR
Other immunosuppressive medication, specify:					
Other immunosuppressive medication, specify:					

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