

Adult Liver Transplant Recipient Registration Worksheet

FORM APPROVED: O.M.B. NO. 0915-0157 Expiration Date: 12/31/2011

Note: These worksheets are provided to function as a guide to what data will be required in the online TIEDI® 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® 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.

Name:	DOB:
SSN:	Gender:
HIC:	Tx Date:
State of Permanent Residence: **	
Permanent Zip: *	
1	
Recipient Center:	
Surgeon Name: *	
NPI#: *	
NPI#: **	
UNOS Donor ID #:	
Donor Type:	
Primary Diagnosis: *	
Specify:	
Date: Last Seen, Retransplanted or Death ★	
Date. Last Seen, Retransplanted of Death ***	
	LIVING
Patient Status: *	O DEAD
	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: Was patient hospitalized during the last 90 days prior to the transplant admission:	O YES O NO O UNK
Medical Condition at time of transplant: ★	IN INTENSIVE CARE UNIT HOSPITALIZED NOT IN ICU NOT HOSPITALIZED
Patient on Life Support: *	YES NO Ventilator Artificial Liver
Specify:	Other Mechanism, Specify
Functional Status: *	
Physical Capacity:	No Limitations Limited Mobility

	0	Wheelchair bound or more limited
	0	Not Applicable (< 1 year old or hospitalized)
	0	Unknown
Working for income: ★	0	YES NO UNK
If No, Not Working Due To:		
	0	Working Full Time
	0	Working Part Time due to Demands of Treatment
	0	Working Part Time due to Disability
If Yes:	0	Working Part Time due to Insurance Conflict
11 165.	0	Working Part Time due to Inability to Find Full Time Work
	0	Working Part Time due to Patient Choice
	0	Working Part Time Reason Unknown
	0	Working, Part Time vs. Full Time Unknown
	0	Within One Grade Level of Peers
	0	Delayed Grade Level
Academic Progress:	0	Special Education
	0	Not Applicable < 5 years old/ High School graduate or GED
	0	Status Unknown
	0	Full academic load
Academic Activity Level:	0	Reduced academic load
	0	Unable to participate in academics due to disease or condition
	0	Not Applicable < 5 years old/ High School graduate or GED

	Status Unknown		
Source of Payment:			
Primary: *			
Specify:			
Secondary:			
Height: ★	ft. in.	cm	ST=
Weight: *	lbs	kg	ST=
BMI:	kg/m ²	Ü	
Previous Transplants:			
Previous Transplant Organ	Previous Transplant Date	Previous Transplant Gra	aft Fail Date
Viral Detection: HIV Serostatus: ★	Positive Negative Not Done UNK/Cannot Disc	lose	
CMV IgG: *	Negative Not Done UNK/Cannot Disc	lose	
CMV IgM: *	Positive Negative		

	0	Not Done
	0	UNK/Cannot Disclose
	0	Positive
HBV Core Antibody: ★	0	Negative
HBV Core Antibody: A	0	Not Done
	0	UNK/Cannot Disclose
	0	Positive
HBV Surface Antigen: ★	0	Negative
Fiby Surface Artigeti.	0	Not Done
	0	UNK/Cannot Disclose
	0	Positive
HCV Serostatus: ★	0	Negative
Tiov ocrosidads.	0	Not Done
	0	UNK/Cannot Disclose
	0	Positive
EBV Serostatus: ★	0	Negative
	0	Not Done
	0	UNK/Cannot Disclose
Any tolerance induction technique used:	0	YES NO C UNK
Pretransplant Lab Date:		
SGPT/ALT:		U/L ST=
Malignancies between listing and transplant: ★	0	YES NO UNK

This question is NOT applicable for patients receiving living	uonor tra	ansplants who were hever on the waiting list.
		Skin Melanoma
		Skin Non-Melanoma
		CNS Tumor
		Genitourinary
		Breast
If yes, specify type:		Thyroid
		Tongue/Throat/Larynx
		Lung
		Leukemia/Lymphoma
		Liver
		Hepatocellular Carcinoma
		Other, specify
Specify:		
Multiple Organ Recipient		
Were extra vessels used in the transplant procedure: Vessel Donor ID:		
Surgical Procedure:	0	ORTHOTOPIC
	0	HETEROTOPIC
	0	Whole Liver
	0	Partial Liver, remainder not Tx or Living Transplant
Procedure Type:	0	Split Liver
	0	Whole Liver with Pancreas (Technical Reasons)
	0	Partial Liver with Pancreas (Technical Reasons)

	0	Split Liver with Pancreas (Technical Reasons)
Split Type:		
Preservation Information:		
Warm Ischemia Time (include anastomotic time):		min ST=
Total Cold Ischemia Time (if pumped, include pump time): *		hrs ST=
Risk Factors:		
Did Patient receive 5 or more units of packed red blood cells within 48 hours prior to transplantation due to spontaneous portal hypertensive bleeding:	0	YES NO UNK
Spontaneous Bacterial Peritonitis:	0	YES NO UNK
Previous Abdominal Surgery: ★	0	YES NO UNK
Portal Vein Thrombosis: ★	0	YES NO UNK
Transjugular Intrahepatic Portacaval Stint Shunt: *	0	YES NO UNK
Incidental Tumor found at time of Transplant:	0	YES NO C UNK
	0	Hepatocellular Adenoma
	0	Hemangioma
	0	Hemangioendothelioma
	0	Angiomyolipoma
If yes, specify tumor type:	0	Bile Duct Cystadenocarcinoma
	0	Cholangiocarcinoma
	0	Hepatocellular Carcinoma
	0	Hepatoblastoma
	0	Angiosarcoma

	Other Primary Liver Tumor, Specify
Specify:	
Pathology Conf. Liver Diag. of Hospital Discharge: *	
Specify:	
Graft Status: ★	C Functioning Failed
If death is indicated for the recipient, and the death was a r	result of some other factor unrelated to graft failure, select Functioning.
Date of Graft Failure:	
Causes of graft failure:	
Primary Graft Failure	C YES C NO C UNK
Vascular Thrombosis	O YES O NO UNK
	C YES C NO C UNK
	C YES C NO C UNK
	C YES C NO C UNK
Biliary Tract Complication	C YES C NO C UNK
Hepatitis: DeNovo	C YES C NO C UNK
Hepatitis: Recurrent	C YES C NO UNK
Recurrent Disease (non-Hepatitis)	C YES C NO C UNK
Acute Rejection	C YES C NO UNK
Infection	C YES C NO UNK
Other, Specify:	

Discharge Lab Date:

Total Bilirubin:			mg/dl	ST=	
SGPT/ALT:			U/L	ST=	
Serum Albumin:			g/dl	ST=	
Serum Creatinine:			mg/dl	ST=	
INR:				ST=	
	0	Yes, at least one	episode treated with anti-	-rejection agent	
Did patient have any acute rejection episodes between transplant and discharge: ★	0	Yes, none treated	d with additional anti-rejec	ction agent	
	0	No			
	0	Biopsy not done			
Was biopsy done to confirm acute rejection:	0	Yes, rejection cor	nfirmed		
	0	Yes, rejection not	confirmed		
Biological or Anti-viral Therapy:	0	YES O NO C	Unknown/Cannot disc	close	
		Acyclovir (Zovirax)			
		Cytogam (CMV)			
		Gamimune			
		Gammagard			
W.V I I I II II I I I I I I I I I I I		Ganciclovir (Cytovene)			
If Yes, check all that apply:		Valgancyclovir (Va	alcyte)		
		HBIG (Hepatitis B	Immune Globulin)		
		Flu Vaccine (Influe	enza Virus)		
		Lamivudine (Epivir	r) (for treatment of Hepati	tis B)	
		Other, Specify			

	Valacyclovir (Valtrex)
Specify:	
Specify:	
Other therapies:	C YES C NO
	Photopheresis
If Yes, check all that apply:	Plasmapheresis
	Total Lymphoid Irradiation (TLI)
Are any medications given currently for maintenance or anti-rejection: *	C YES C NO
Did the patient participate in any clinical research protocol for immunosuppressive medications:	C YES C NO
If Yes, Specify:	
View Immunosuppressive Medications	
Definitions Of Immunosuppressive Medications	
	Previous Maintenance (Prev Maint), Current Maintenance (Curr Maint) or Anti- d for the recipient during this follow-up period, and for what reason. If a medication was
clinic visit to the current clinic visit, for varying periods of time	pressive medications given during the report period, which covers the period from the last which may be either long-term or intermediate term with a tapering of the dosage until the intenance drug (example: Prednisone, Cyclosporine, Tacrolimus, Mycophenolate Mofetil, nosuppressive medications given to treat rejection episodes.
periods of time which may be either long-term or intermediate	ressive medications given at the current clinic visit to begin in the next report for varying term with a tapering of the dosage until the drug is either eliminated or replaced by cyclosporine, Tacrolimus, Mycophenolate Mofetil, Azathioprine, or Rapamycin). This does t rejection episodes.
since the last clinic visit (example: Methylprednisolone, Atgam Tacrolimus to Cyclosporine; or from Mycophenolate Mofetil to immunosuppression, but should be listed under maintenance	suppressive medications given for the purpose of treating an acute rejection episode , OKT3, or Thymoglobulin). When switching maintenance drugs (example: from Azathioprine) because of rejection, the drugs should not be listed under AR immunosuppression. The defications since the last clinic visit, not just at the time of the current clinic visit.
If an immunosuppressive medication other than those listed is	being administered (e.g., new monoclonal antibodies), select Previous Maint, or Current ld, and enter the full name of the medication in the space provided. Do not list non-

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 short finite period 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 will not be used long-term for immunosuppressive maintenance. Induction agents are usually polyclonal, monoclonal, or IL-2 receptor antibodies (example: Methylprednisolone, Atgam, Thymoglobulin, OKT3, Simulect, or Zenapax). Some of these drugs might be used for another finite period for rejection therapy and would be recorded as rejection therapy if used for this reason. For each induction medication indicated, write the total number of days the drug was actually administered in the space provided. For example, if Simulect or Zenapax 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 for varying periods of time which may be either long-term or intermediate term with a tapering of the dosage until the drug is either eliminated or replaced by another long-term maintenance drug (example: Prednisone, Cyclosporine, Tacrolimus, Mycophenolate Mofetil, Azathioprine, or Rapamycin). 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, Atgam, OKT3, or Thymoglobulin). When switching maintenance drugs (example: from Tacrolimus to Cyclosporine; or from Mycophenolate Mofetil to Azathioprine) because of rejection, the drugs should not be listed under AR immunosuppression, but should be listed under maintenance 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. <u>Do not list non-immunosuppressive medications.</u>

Steroids	Ind.	Days	ST	Maint	AR
(Prednisone, Methylprednisolone, Solumedrol, Medrol, Decadron)			-		
Atgam (ATG)			-		
OKT3 (Orthoclone, Muromonab)			- L	- 🗆	
Thymoglobulin			-		
Simulect - Basiliximab			-		
Zenapax - Daclizumab			-		_
Azathioprine (AZA, Imuran)			-		
EON (Generic Cyclosporine)			-	- П	
Gengraf (Abbott Cyclosporine)			-	_	П
Other generic Cyclosporine, specify brand:			-	-	
Neoral (CyA-NOF)			-		
Sandimmune (Cyclosporine A)			-	_	
CellCept (Mycophenolate Mofetil; MMF)			_		

Generic MMF (Generic CellCept)			
Prograf (Tacrolimus, FK506)			
Generic Tacrolimus (Generic Prograf)			
Advagraf (Tacrolimus Extended or Modified Release)			
Nulojix (Belatacept)			
Sirolimus (RAPA, Rapamycin, Rapamune)			
Myfortic (Mycophenolate Sodium)			
6			_
	Ind. Days	ST Maint AR]
Campath - Alemtuzumab (anti-CD52)	Ind. Days	ST Maint AR	
Campath - Alemtuzumab (anti-CD52) Cyclophosphamide (Cytoxan)	= -	ST Maint AR	
	= -	ST Maint AR	
Cyclophosphamide (Cytoxan)			
Cyclophosphamide (Cytoxan) Leflunomide (LFL, Arava) Methotrexate (Folex, PFS, Mexate-AQ,			
Cyclophosphamide (Cytoxan) Leflunomide (LFL, Arava) Methotrexate (Folex, PFS, Mexate-AQ, Rheumatrex)			
Cyclophosphamide (Cytoxan) Leflunomide (LFL, Arava) Methotrexate (Folex, PFS, Mexate-AQ, Rheumatrex) Other Immunosuppressive Medication, Specify			
Cyclophosphamide (Cytoxan) Leflunomide (LFL, Arava) Methotrexate (Folex, PFS, Mexate-AQ, Rheumatrex) Other Immunosuppressive Medication, Specify			
Cyclophosphamide (Cytoxan) Leflunomide (LFL, Arava) Methotrexate (Folex, PFS, Mexate-AQ, Rheumatrex) Other Immunosuppressive Medication, Specify			
Cyclophosphamide (Cytoxan) Leflunomide (LFL, Arava) Methotrexate (Folex, PFS, Mexate-AQ, Rheumatrex) Other Immunosuppressive Medication, Specify			