## Records 🕐

## Adult Intestine Transplant Recipient Follow-Up Worksheet

FORM APPROVED: O.M.B. NO. 0915-0157 Expiration Date: 10/31/2010

Note: These worksheets are provided to function as a guide to what data will be required in the online TIEDI<sup>®</sup> 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<sup>®</sup> 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:	Gender:
HIC:	Tx Date:
Previous Follow-Up:	Previous Px Stat Date:
Transplant Discharge Date:	
State of Permanent Residence:*	
Zip Code: *	-
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Provider Information	
Recipient Center:	
Followup Center:	
Physician Name: *	
NPI#:*	
Follow-up Care Provided By: *	<ul> <li>Transplant Center</li> <li>Non Transplant Center Specialty Physician</li> <li>Primary Care Physician</li> <li>Other Specify</li> </ul>
Specify:	
Donor Information	
UNOS Donor ID #:	
Donor Type:	
Patient Status	
Date: Last Seen, Retransplanted or Death $st$	

Patient Status:\*

O DEAD

© RETRANSPLANTED			
Primary Cause of Death:			
Specify:			
Contributory Cause of Death:			
Specify:			
Contributory Cause of Death:			
Specify:			
Hospitalizations:			
Has the patient been hospitalized since the last patient status date: *	C YES C NO C UNK		
Number of Hospitalizations:	ST=		
Noncompliance:			
Was there evidence of noncompliance with immunosuppression medication during this follow-up period that compromised the patient's recovery:	CYES CNO CUNK		
Functional Status: *			
	No Limitations		
	C Limited Mobility		
Physical Capacity:	Wheelchair bound or more limited		
	Not Applicable (< 1 year old or hospitalized)		
	C Unknown		
Working for income: *	YES NO VINK		
If No, Not Working Due To:			
	Working Full Time		
	Working Part Time due to Demands of Treatment		
	Working Part Time due to Disability		
	Working Part Time due to Insurance Conflict		

	Working Part Time due to Inability to Find Full Time Work		
	Working Part Time due to Patient Choice		
If Yes:	Working Part Time Reason Unknown		
	Working, Part Time vs. Full Time Unknown		
	Within One Grade Level of Peers		
	Delayed Grade Level		
Academic Progress	Special Education		
	Not Applicable < 5 years old/ High School graduate or GED		
	Status Unknown		
	Full academic load		
	Reduced academic load		
Academic Activity Level			
	Not Applicable < 5 years old/ High School graduate or GED		
	Status Unknown		
Primary Insurance at Follow-up: *			
Specify:			
Clinical Information			
Height:	ft. in. cm ST=		
Weight:	lbs. kg ST=		
BMI:	kg/m <sup>2</sup>		
Graft Status: *	Functioning Failed		
If death is indicated for the recipient, and	the death was a result of some other factor unrelated to graft failure, select Functioning.		
TPN Dependent:	C YES C NO		
IV Dependent:	C YES C NO		
Oral Feeding:	C YES C NO		
Tube Feeding:	🤄 YES 🌀 NO		

Date of Failure:				
	C RECURRENT TUMOR			
	C ACUTE REJECTION			
	CHRONIC REJECTION			
Primary Cause of Failure:	C TECHNICAL PROBLEMS			
Finnary Gause of Fanule.				
	C LYMPHOPROLIFERATIVE DISEASE			
	PATIENT NONCOMPLIANCE			
	OTHER SPECIFY			
Other, Specify:				
Diabetes onset during the follow-up period: *	YES NO UNK			
Insulin dependent:	C YES C NO C UNK			
Most Recent Lab date:				
Total Bilirubin:	mg/dl ST=			
Serum Albumin:	mg/dl ST=			
Serum Creatinine: *				
	mg/dl ST=			
	mg/dl ST=     Yes, at least one episode treated with anti-rejection agent			
Did actiont have any coute rejection opioodeo				
Did patient have any acute rejection episodes during the follow-up period: *	Yes, at least one episode treated with anti-rejection agent			
Did patient have any acute rejection episodes during the follow-up period: *	<ul> <li>Yes, at least one episode treated with anti-rejection agent</li> <li>Yes, none treated with additional anti-rejection agent</li> </ul>			
Did patient have any acute rejection episodes during the follow-up period: *	<ul> <li>Yes, at least one episode treated with anti-rejection agent</li> <li>Yes, none treated with additional anti-rejection agent</li> <li>No</li> <li>Unknown</li> </ul>			
Did patient have any acute rejection episodes during the follow-up period: ≭	<ul> <li>Yes, at least one episode treated with anti-rejection agent</li> <li>Yes, none treated with additional anti-rejection agent</li> <li>No</li> <li>Unknown</li> <li>Biopsy not done</li> </ul>			
Did patient have any acute rejection episodes during the follow-up period: * Was biopsy done to confirm acute rejection:	<ul> <li>Yes, at least one episode treated with anti-rejection agent</li> <li>Yes, none treated with additional anti-rejection agent</li> <li>No</li> <li>Unknown</li> <li>Biopsy not done</li> <li>Yes, rejection confirmed</li> </ul>			
during the follow-up period: *	<ul> <li>Yes, at least one episode treated with anti-rejection agent</li> <li>Yes, none treated with additional anti-rejection agent</li> <li>No</li> <li>Unknown</li> <li>Biopsy not done</li> </ul>			

Post Transplant Malignancy: *	CYES NO CUNK
Donor Related:	C YES C NO C UNK
Recurrence of Pre-Tx Tumor:	SYES SNO UNK
De Novo Solid Tumor:	C YES C NO C UNK
De Novo Lymphoproliferative disease and Lymphoma:	C YES C NO C UNK

Treatment	
Biological or Anti-viral therapy:	YES ONO Unknown/Cannot disclose
	Acyclovir (Zovirax)
	Cytogam (CMV)
	Gamimune
	Gammagard
	Ganciclovir (Cytovene)
If Yes, check all that apply:	Valgancyclovir (Valcyte)
	HBIG (Hepatitis B Immune Globulin)
	Flu Vaccine (Influenza Virus)
	Lamivudine (Epivir) (for treatment of Hepatitis B)
	Valacyclovir (Valtrex)
	Other, Specify
Specify: *	
Specify:	
Other therapies:	O YES O NO
	Photopheresis
If Yes, check all that apply:	Plasmapheresis
	Total Lymphoid Irradiation (TLI)
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Immunosuppressive Information

Previous Va	alidated	Maintenance	Follow-Up
Medication	s:		

Previous Validated Maintenance Follow-Up Medications:

Were any medications given during the follow- up period for maintenance:	<ul> <li>Yes, same as validated TRR form</li> <li>Yes, same as previous validated report</li> <li>Yes, but different than previous validated report</li> <li>None given</li> </ul>	
Did the physician discontinue all maintenance immunosuppressive medications:	C YES C NO	
Did the patient participate in any clinical research protocol for immunosuppressive medications: Specify: *	YES NO	
Immunosuppressive Medications		
View Immunosuppressive Medications		
Definitions Of Immunosuppressive Follow-Up Medications		
	ed, check <b>Previous Maintenance (Prev Maint)</b> , <b>Current Maintenance (Curr</b> ions that were prescribed for the recipient during this follow-up period, and for associated box(es) blank.	

**Previous Maintenance (Prev Maint)** includes all immunosuppressive medications given during the report period, which covers the period from the last clinic visit to the current clinic visit, 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.

**Current Maintenance (Curr Maint)** includes all immunosuppressive medications given at the current clinic visit to begin in the next report 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.

Anti-rejection (AR) immunosuppression includes all immunosuppressive medications given for the purpose of treating an acute rejection episode since the last clinic visit (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 <u>should not</u> be listed under AR immunosuppression, but <u>should be</u> listed under maintenance immunosuppression.

Note: The Anti-rejection field refers to any anti-rejection medications 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 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.** 

	Prev Maint	Curr Mai	nt AR
Steroids (Prednisone,Methylprednisolone,Solumedrol,Medrol,Decadron)			

Atgam (ATG)		
OKT3 (Orthoclone, Muromonab)		
Thymoglobulin		
Simulect - Basiliximab		
Zenapax - Daclizumab		
Azathioprine (AZA, Imuran)		
EON (Generic Cyclosporine)		
Gengraf (Abbott Cyclosporine)		
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)		
Modified Release Tacrolimus FK506E (MR4)		
Sirolimus (RAPA, Rapamycin, Rapamune)		
Myfortic (Mycophenolate Sodium)		

Other Immunosuppressive Medications			
	Prev Maint	Curr Maint	AR
Campath - Alemtuzumab (anti-CD52)			
Cyclophosphamide (Cytoxan)			
Leflunomide (LFL, Arava)			
Methotrexate (Folex, PFS, Mexate-AQ, Rheumatrex)			
Other Immunosuppressive Medication, Specify			
Rituximab			

Investigational Immunosuppressive Medications			
	Prev Main	t Curr Maint	AR
Everolimus (RAD, Certican)			
Other Immunosuppressive Medication, Specify			

UNOS View Only		
Comments:		