CIT-03

ADVERSE EVENT

Subject ID	Page 1 of 5
Report Number	
A. ADVERSE EVENT	
1. Date of adverse event/	
2. Date site became aware of AE///	
3. Adverse Event Term	
4. Describe event or problem. (Include any details relating to diagnosis.)	
No Yes 5. O Is this an exacerbation of a pre-existing condition (existing prior to enrollment)?	
6. Describe relevant tests/laboratory data, including dates.	
7. Describe other relevant history, including preexisting medical conditions. (e.g., allergies, race, pregnancy, smoking and alcohol use, hepatic/renal dysfunction, etc.)	

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8. Outcomes attributed to adverse event (Check all that apply) (ALL choices below represent an SAE except "None of the above") Death:	
If outcome changes to an SAE during a postcomplete change, Q8a and 8b pop-up. 8a. Date the Adverse Event became a Serious Adverse Event: (dd/mmm/yyyy) 8b. Date the site became aware that the Adverse Event became a Serious Adverse Event: (dd/mmm/yyyy)	
9. IntensityPlease follow the guidelines in the "TCAE in Trials of Adult Pancreatic Islet Transplantation OMild/Grade I OModerate/Grade II OSevere/Grade III OLife-threatening/Grade IV ODeath/Grade V (If question 9 is Death/Grade V, go to question 10)	on"
10. Was/will an autopsy be performed? (select one) O No O Yes Please provide a de-identified copy to the DCC O Unknown	
11. Indicate outcome of the event OContinuing OResolved (or resolved with sequelae) If resolved, give date of resolution (dd/mmm/yy	 yyy)

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No Yes	Page 3 of 5
b. Action taken regarding deoxyspergualin	
O None O Dose reduced	
OInterrupted ODiscontinued	
ODiscontinued ODose increased	

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B. SUSPECT MEDICATION(S)

	Suspect Medication 1	Suspect Medication 2	Suspect Medication 3
1. Name	i.Islet Transplantation □ Purified Human Pancreatic Islets (check if ever received islets) □ Transplant Procedure(check if ever had transplant procedure initiated)	Immunosuppression and infection prophylaxis	Deoxyspergualin
2. Dose	i.		ii.
3. Therapy dates (if unknown, give best estimate)	i. Date of most recent islet transplantation//		ii. Introduction date//iii. Date of last dose//(dd/mmm/yyyy)
4. Diagnosis for use	Type I Diabetes Mellitus	Islet Transplant/Immunosuppression	Islet Transplant/Immunosuppression
5. Event abated after use stopped or dose reduced?	i. O No O Yes O Doesn't apply	ii. O No O Yes O Doesn't apply	iii.O No O Yes O Doesn't apply
6.Event reappeared after reintroduction?	i. O No O Yes O Doesn't apply	ii. O No O Yes O Doesn't apply	iii.O No O Yes O Doesn't apply
7. Lot number	i		ii.
8. Expiration Date (if known)	i. N/A		ii///(dd/mmm/yyyy)

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C. OTHER MEDICATIONS	
What concomitant medications was the subject receiving at the time of the event? (Exclude treatment of event)	
INSTRUCTIONS:	
1. Select the buttons below to add data to the Other Medications text box.	•
O Select to add data that has been entered into the subject's Concomitant Meds eCRF	
O Select to add data that has been entered into the subject's Study Treatment Regimen eC	RF
2. Please review added data carefully for accuracy and modify this form and the Concor Meds eCRF and/or the Study Treatment Regimen eCRF as needed.	nitant
3. If the subject was on insulin therapy at the time of the event , their insulin therapy must be added to the text box below .	,
4. Add any additional medication information, if applicable.	

CIT-()3				STUDYTRE	EATMENT RI	EGIMEN
Subjec	et ID						
							Page 1 of 2
A	. INVESTIGA	TIONAL	AGENT				
	Drug	Date	 e	Total De	ose on this Date (mg)	Add new Entry	
	O DSG		/			·	
		(dd	/mmm/yyyy)				
				<u> </u>			
- B.]	INDUCTION 1	MEDICA	ATIONS				
	Drug	Date		Total Dos	se on this Date (mg)	Add new Entry	
(O ATG	/	_/				
	Other	(dd/mm	m/yyyy)				
C. 8	SUBSEQUENT	TRANS	SPLANT IND	UCTION M	IEDICATION		
	Drug		Date	Total Dos	se on this Date (mg)	Add new Entry	
	O Daclizumab						
	O Basiliximab		(dd/mmm/yyyy)				
L					<u> </u>		
D.	IMMUNUSUI	PPRESS	VE/ANTI-IN	FLAMMA	TORY MEDICATIO	ONS	
	Drug	7	ate	Total D	ose on this Date (mg)	Add new Entry	
							
	O Etanercept	(d	d/mmm/yyyy)				
	Sections A-	D will be	available for .	Induction o	only.		
			ilable for first	-	•		
	Section C w	m be ava	ulable for seco	ona ana ini	rd transplants only.		
E.	MAINTENA	NCE IM	MUNOSUPP	RESSION 1	MEDICATIONS Add new Entry		
	Drug		Total Dose ((mg) / Day	Start Date	Stop Date	
	OTacrolimu	IS					
	O Sirolimus					/	
	O Cyclospoi	rine			(dd/mmm/yyyy)	(dd/mmm/yyyy)	

TROUGH LEVELS						
Drug	Date of Draw	v Trough l	Level (ng/mL)	Add ne	ew Entry	
O Tacrolimus O Sirolimus O Cyclosporine	/(dd/mmm/yyy	yy) Undetec	ctable		•	
G. OTHER MAINT	ENANCE IM	MUNOSUPPRE	SSION MEDIC	CATIONS	-	<i>i</i>
Drug	Т	Total Dose (mg) /	Day Start Date	2	Stop Date	
O Mycophenolate	sodium				/ /	
O Mycophenolate	mofetil		(dd/mn	nm/yyyy)	(dd/mmm/	 ⁄yyyy)
1 ~						
O Other						
If Other		omplete Ma	· ·	col De	viation for	m.
If Other H. INFECTION PR		MEDICATIONS	3	col De	Add new entr	
If Other H. INFECTION PR	OPHYLAXIS		3	col De		
If Other H. INFECTION PR	OPHYLAXIS S=1 tab)* oche)	MEDICATIONS	3		Add new entr	ry
Drug O TMP / SMX (SSO) Clotrimazole (tro	S=1 tab)* oche) ng)	MEDICATIONS	y Start Date		Add new entr	ry
Drug O TMP / SMX (SSO) Clotrimazole (tro	OPHYLAXIS S=1 tab)* ocheo ng) gth TMP = 80m	Total Dose / Day	y Start Date		Add new entr	ry /y)
Drug O TMP / SMX (SSO Clotrimazole (troe) Valganciclovir (note that we will be supported by the control of the	G=1 tab)* oche ng) gth TMP=80m	Total Dose / Day	y Start Date		Add new entropy Stop Date ///_ (dd/mmm/yyy	ry /y)
Drug O TMP / SMX (SSO Clotrimazole (tro O Valganciclovir (notation) *Single Stren I. ANTICOAGULA	G=1 tab)* oche ng) gth TMP=80m	Total Dose / Day	y Start Date // (dd/mmm/		Add new entrestop Date //_(dd/mmm/yyy) Add new entrestop	ry /y)
Drug O TMP / SMX (SSO Clotrimazole (troe) Valganciclovir (note that with the control of the con	G=1 tab)* oche ng) gth TMP=80m	Total Dose / Day	y Start Date // (dd/mmm/	 (yyyy)	Add new entrestop Date //_ (dd/mmm/yyy) Add new entrestop	ry /y)