

Hemovigilance Module Adverse Reaction Infection

*Required for saving *Facility ID#: _____ NHSN Adverse Reaction #: _____ Patient Information *Gender: M F Other *Date of Birth: ___/___ *Patient ID: ____ Social Security #: _____ Secondary ID: _____ Medicare #: _____ Last Name: First Name: Middle Name: ☐ Hispanic or Latino ☐ Not Hispanic or Not Latino Ethnicity ☐ Black or African American Race American Indian/Alaska Native Asian Native Hawaiian/Other Pacific Islander White ***Blood Group:** ☐ A- ☐ A+ ☐ B- ☐ B+ ☐ AB-☐ AB+ ☐ O- ☐ O+ ☐ Blood type not done ☐ Transitional ABO / Rh + ☐ Transitional ABO / Rh -Transitional ABO / Transitional Rh ☐ Group A/Transitional Rh ☐ Group B/Transitional Rh ☐ Group O/Transitional Rh ☐ Group AB/Transitional Rh **Patient Medical History** List the patient's admitting diagnosis. (Use ICD-10 Diagnostic codes/descriptions) Description: Code: _____ Description: Description: Code: List the patient's underlying indication for transfusion. (Use ICD-10 Diagnostic codes/descriptions) Code: Description: Description: Code: _____ Code: Description: ☐ UNKNOWN List the patient's comorbid conditions at the time of the transfusion related to the adverse reaction. (Use ICD-10 Diagnostic codes/descriptions) NONE Description: Code: Description: Code: Description: Assurance of Confidentiality: The voluntarily provided information obtained in this surveillance system that would permit identification of any individual or institution is collected with a guarantee that it will be held in strict confidence, will be used only for the purposes stated, and will not otherwise be disclosed or released without the consent of the individual, or the institution in accordance with Sections 304, 306 and 308(d) of the Public Health Service Act (42 USC 242b, 242k, and 242m(d)). Public reporting burden of this collection of information is estimated to average 20 minutes per response, including the time for reviewing instructions, searching existing data sources, gathering and maintaining the data needed, and completing and reviewing the collection of information. An agency may not conduct or sponsor, and a person is not required to respond to a collection of information unless it displays a currently valid OMB control number. Send comments regarding this burden estimate or any other aspect of this collection of information, including suggestions for reducing this burden to CDC, Reports Clearance Officer, 1600 Clifton Rd., MS D-74, Atlanta, GA 30333 ATTN: PRA (0920-0666).



Infection

	dical procedure including past procedures and procedures to hospital or outpatient stay. (Use ICD-10 Procedure	be UNKNOWN NONE
Code:	Description:	
Code:	Description:	
Code:	Description:	
Additional Information		
Transfusion History		
Has the patient received a pro	evious transfusion?	UNKNOWN
Blood Product:	VB ☐ RBC ☐ Platelet ☐ Plasma ☐ Cryoprecipita	ate Granulocyte
Date of Transfusion:	/	
Was the patient's adverse re	eaction transfusion-related?	NO
• •	about the transfusion adverse reaction.	_
* *		OSTR FNHTR
	PTP TACO TAD TA-GVHD TRALI	
Reaction Details		
	/ *Time reaction occurred:: T	ime unknown
Is this reaction associated with a	an incident? Yes No If Yes, Incident #:	
I IS THIS reaction associated with a	an incident?	
Investigation Results		
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Investigation Results * Infection *Case Definition Was a test to detect a spec	cific pathogen performed on the recipient post-transfus	
Investigation Results * Infection *Case Definition Was a test to detect a specific yes, positive or reactive	cific pathogen performed on the recipient post-transfus	ion?
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Investigation Results * Infection *Case Definition Was a test to detect a specific yes, positive or reactive Org1 Was a test to detect a specific yes, positive or reactive Org1 United Section 1.	cific pathogen performed on the recipient post-transfusion results?	ion?
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Cutaneous:	☐ Edema ☐ Other rash	☐ Flushing	<u> </u>	aundice Irticaria (hives)		
Hemolysis/Hemorrhage:	☐ Disseminated intravascular coagulation ☐ Hemoglobinemia ☐ Positive antibody screen					
Pain:	Abdominal pain	Back pain	☐ Flank pain	☐ Infusion site pain		
Renal:	Hematuria			Dliguria		
	_		☐ Bronchospasi	•		
Respiratory:	☐ Bilateral infiltrates on chest x-ray ☐ Bronchospasm ☐ Cough ☐ Hypoxemia ☐ Shortness of breath					
Other: (specify)	Trypoxernia Grioriness of breath					
*Severity						
Did the patient receive or ex	perience any of the fo	ollowing?				
☐ No treatment required	d	☐ Symptomatic t	reatment only			
☐ Hospitalization, inlcud	ding prolonged hospita	alization	Life-threate	ning reaction		
☐ Disability and/or incap	pacitation	☐ Congenital and	omaly or birth defe	ct(s) of the fetus		
Other medically impo	rtant conditions	Death	Unknown o	r not stated		
*Imputability						
*Imputability Which best describes the relationship between the transfusion and the reaction? No other potential exposures to the pathogen could be identified in the recipient. Evidence is clearly in favor of a cause other than transfusion, but transfusion cannot be excluded. There is conclusive evidence beyond reasonable doubt of a cause other than the transfusion. The relationship between the adverse reaction and the transfusion is unknown or not stated. Check all that apply: Evidence of the pathogen in the transfused component. Evidence of the pathogen in the donor at the time of donation. Evidence of the pathogen in an additional component from the same donation. Evidence of the pathogen in an additional recipient of a component from the same donation. Evidence that the identified pathogen strains are related by molecular or extended phenotypic comparison testing with statistical confidence (p<0.05). Evidence that the transfused component was negative for this pathogen at the time of transfusion Evidence that the donor was negative for this pathogen at the time of donation. Evidence that additional components from the same donation were negative for this pathogen. Evidence that the recipient was not infected with the pathogen prior to transfusion. Did the transfusion occur at your facility? YES NO						
Module-generated Designa	tions					
NOTE: Designations for case de application based on responses	finition, severity, and i			gned in the NHSN		
* Do you agree with the <u>cas</u> ^Please indicate your design		ation?	☐ YES	□ NO		
*Do you agree with the <u>set</u>	verity designation?		☐ YES	□NO		



^Please i	ndicate your designat	ion						<u> </u>
_	agree with the <u>impur</u> ndicate your designat				ES] NO	
Patient Tre	atment							
If yes, sele	ent receive treatment ect treatment(s): lication (Select the type) Antipyretics Are Intravenous Immuno Antithymocyte globu	pe of medication ntihistamines [nglobulin] In	n) Inotropes/Vasop travenous steroids			ator	JNKNO ☐ Di ntibiotic	uretics
☐ Volu	ıme resuscitation (Intr	avenous colloid	s or crystalloids)					
	piratory support <i>(Sele</i>] Mechanical ventilatial replacement theraphemodialysis	on Nonir	nvasive ventilation pe of therapy)	_ ,,	ofiltration			
	· -	-						
	☐ Phlebotomy ☐ Other Specify:							
Outcome								
*Outcome: Death Major or long-term sequelae Minor or no sequelae Not determined Date of Death:// ^If recipient died, relationship of transfusion to death: Definite Probable Possible Doubtful Ruled Out Not determined Cause of death: Was an autopsy performed? Yes No								
Component	Details							
*Was a partic	cular unit implicated	d in (i.e., respo	onsible for) the a	dverse	☐ Yes		No [] N/A
Transfusion Start and End Date/Time	*Component code (check system used)	Amount transfused at reaction onset	^Unit number (Required for Infection and TRALI)	*Unit expiration Date/Time	*Blood of unit)	Implicat ed Unit?
^IMPLICATED	UNIT			1				
	☐ ISBT-128	_						
:	☐ Codabar	☐ Entire unit ☐ Partial unit			□ A-	□ A+	□ B-	Y
		mL			□B+ [☐ AB-	☐ AB+	
:				:	□ o- [□ 0+	□ N/A	
// :	☐ ISBT-128	☐ Entire unit ☐ Partial unit mL			□ A- [□ A+	□ B-	N
/ /				:	□B+ [□ AB-	□ AR+	



::				□ O- □ O+ □ N/A
Custom Field	ds			
Label			Label	
		/	-	
Comments				