

Hemovigilance Module Adverse Reaction Infection

*Required for saving							
*Facility ID#: NHSN Adverse Reaction #:							
Patient Information							
	*Gender: M F Other *Date of Birth://						
Social Security #:	Secondary ID: Medicare #:						
Last Name:	First Name: Middle Name:						
Ethnicity Hispanic or Latino	☐ Not Hispanic or Not Latino						
Race American Indian/Alaska							
☐ Native Hawaiian/Other F	-						
<u>-</u>	□B+ □ AB- □ AB+ □ O- □ O+ □ Blood type not done						
	+ Transitional ABO / Rh - Transitional ABO / Transitional Rh						
	/Transitional Rh Group O/Transitional Rh Group AB/Transitional Rh						
Patient Medical History							
,	. (Use ICD-10 Diagnostic codes/descriptions)						
Code: De	escription:						
Code: De	escription:						
Code: De	escription:						
List the patient's underlying indication	n for transfusion. (Use ICD-10 Diagnostic codes/descriptions)						
Code: De	escription:						
Code: De	escription:						
Code: De	escription:						
List the patient's comorbid conditions reaction. (Use ICD-10 Diagnostic cod	s at the time of the transfusion related to the adverse UNKNOWN des/descriptions)						
Code: De	escription:						
	escription:						
	escription:						
of any individual or institution is collected with a stated, and will not otherwise be disclosed or re	provided information obtained in this surveillance system that would permit identification a guarantee that it will be held in strict confidence, will be used only for the purposes released without the consent of the individual, or the institution in accordance with ealth Service Act (42 USC 242b, 242k, and 242m(d)).						
reviewing instructions, searching existing data collection of information. An agency may not cunless it displays a currently valid OMB control	ormation is estimated to average 20 minutes per response, including the time for sources, gathering and maintaining the data needed, and completing and reviewing the conduct or sponsor, and a person is not required to respond to a collection of information of number. Send comments regarding this burden estimate or any other aspect of this is for reducing this burden to CDC, Reports Clearance Officer, 1600 Clifton Rd., MS D-74,						



Infection

performed during the current	•	• •	•	and procedures to be procedure	
codes/descriptions)					NONE
Code:	Descr	iption:			
Code:	Descr	iption:			
Code:	Descr	iption:			
Additional Information					
Transfusion History					
Has the patient received a pr	evious transfu	sion?	☐ YES	□ NO □ U	NKNOWN
Blood Product:	VB □ RBC	☐ Platelet	☐ Plasma	☐ Cryoprecipitate	☐ Granulocyte
Date of Transfusion:	//	UNK	NOWN		
Was the patient's adverse r	eaction transfu	usion-related?	[☐ YES ☐ NO	
If yes, provide information a	about the trans	fusion adverse	e reaction.		
Type of transfusion adverse	e reaction:	☐ Allergic	\square AHTR	☐ DHTR ☐ DST	R
	PTP TA				
☐ OTHER Specify	·				
Reaction Details					
*Date reaction occurred:/_	/ *Ti	me reaction of	occurred:	:	unknown
*Facility location where patien	ıt was transfu	sed:			
Is this reaction associated with a	an incident?	☐ Yes	☐ No	If Yes, Incident #:	
Investigation Results					
Investigation Results * Infection					
* Infection					? ☐ Yes ☐ No
* Infection *Case Definition	cific pathoge	n performed o			? ☐ Yes ☐ No
* Infection *Case Definition Was a test to detect a spe	cific pathogei	n performed o	on the recipie	ent post-transfusion?	
* Infection *Case Definition Was a test to detect a spe If Yes, positive or reactive	cific pathoge e results? Or	n performed o	on the recipie	ent post-transfusion?	
* Infection *Case Definition Was a test to detect a spe If Yes, positive or reactive Org1	cific pathogei e results? Org cific pathogei	n performed o Yes g2 n performed o	on the recipie No n the donor	ent post-transfusion? Org3 post-donation?	
* Infection *Case Definition Was a test to detect a spe If Yes, positive or reactive Org1 Was a test to detect a spe If Yes, positive or reactive Org1 Org1	cific pathoger e results? Ore cific pathoger e results? Ore	n performed of Yes g2 n performed of Yes g2	on the recipie No No n the donor	ont post-transfusion? Org3 post-donation? Org3	Yes □ No
* Infection *Case Definition Was a test to detect a spe If Yes, positive or reactive Org1 Was a test to detect a spe If Yes, positive or reactive	cific pathoger e results? cific pathoger e results? Ore Cific pathoger	n performed of yes g2 n performed of yes g2 n performed of yes	on the recipie No n the donor	ont post-transfusion? Org3 post-donation? Org3	Yes ☐ No
* Infection *Case Definition Was a test to detect a spe If Yes, positive or reactive Org1 Was a test to detect a spe If Yes, positive or reactive Org1 Was a test to detect a spe transfusion? (i.e., culture, If Yes, positive or reactive	cific pathoger e results? cific pathoger e results? Or cific pathoger serology, NA	n performed of Yes g2 Yes g2 Yes g2 performed of Yes T) Yes	on the recipie No No No No No n the unit po	ont post-transfusion? Org3 post-donation? Org3 Org3 Yes	Yes ☐ No
* Infection *Case Definition Was a test to detect a spe If Yes, positive or reactive Org1 Was a test to detect a spe If Yes, positive or reactive Org1 Was a test to detect a spe transfusion? (i.e., culture,	cific pathoger e results? cific pathoger e results? Or cific pathoger serology, NA	n performed of Yes g2 Yes g2 Yes g2 performed of Yes T) Yes	on the recipie No No No No No n the unit po	ont post-transfusion? Org3 post-donation? Org3 Org3 Yes	Yes No
* Infection *Case Definition Was a test to detect a spe If Yes, positive or reactive Org1 Was a test to detect a spe If Yes, positive or reactive Org1 Was a test to detect a spe transfusion? (i.e., culture, If Yes, positive or reactive Org1 Org1	cific pathoger e results? cific pathoger e results? Orgorific pathoger serology, NA	n performed of Yes g2 Yes g2 Yes g2 performed of Yes T) Yes	on the recipie No No No No No n the unit po	ont post-transfusion? Org3 post-donation? Org3 Org3 Yes	Yes No
* Infection *Case Definition Was a test to detect a spe If Yes, positive or reactive Org1 Was a test to detect a spe If Yes, positive or reactive Org1 Was a test to detect a spe transfusion? (i.e., culture, If Yes, positive or reactive	cific pathoger e results? cific pathoger e results? Orgorisic pathoger serology, NA e results? Orgorisic	n performed of yes g2 n performed of yes g2 n performed of yes T) Yes g2	on the recipie No n the donor No n the unit po	ont post-transfusion? Org3 post-donation? Org3 Ost- Org3 Ost- Org3 Org3	Yes No
* Infection *Case Definition Was a test to detect a spe If Yes, positive or reactive Org1 Was a test to detect a spe If Yes, positive or reactive Org1 Was a test to detect a spe transfusion? (i.e., culture, If Yes, positive or reactive Org1 Check all that apply:	cific pathoger e results? Cific pathoger e results? Orgorisc pathoger serology, NA e results? Orgorisc pathoger serology, NA e results?	n performed of Yes g2 n performed of Yes g2 n performed of Yes g2 g2 Clinical illness	on the recipie No n the donor No n the unit po	ont post-transfusion? Org3 post-donation? Org3 Ost- Org3 Ost- Org3 Org3	Yes No
* Infection *Case Definition Was a test to detect a spe If Yes, positive or reactive Org1 Was a test to detect a spe If Yes, positive or reactive Org1 Was a test to detect a spe transfusion? (i.e., culture, If Yes, positive or reactive Org1 Check all that apply: Temporally associated	cific pathoger e results? Cific pathoger e results? Orgorisc pathoger serology, NA e results? Orgorisc pathoger serology, NA e results?	n performed of Yes g2 n performed of Yes g2 n performed of Yes g2 clinical illness	on the recipie No n the donor No n the unit po	ont post-transfusion? Org3 post-donation? Org3 Ost- Org3 Org3 h infection	Yes No



-	Cutaneous:	☐ Edema ☐ Other rash	☐ Flushing ☐ Pruritus		aundice Irticaria (hives)						
-	Hemolysis/Hemorrhage:	☐ Disseminated intravascular coagulation ☐ Hemoglobinemia ☐ Positive antibody screen									
-	Pain:		☐ Abdominal pain ☐ Back pain ☐ Flank pain ☐ Infusion site pain								
-	Renal:	☐ Hematuria ☐ Hemoglobinuria ☐ Oliguria ☐ Bilateral infiltrates on chest x-ray ☐ Bronchospasm ☐ Cough ☐ Hypoxemia ☐ Shortness of breath									
-	Respiratory:										
	Other: (specify)										
	*Severity										
	Did the patient receive or ex	perience any of the follo	owing?								
	☐ No treatment required	d [☐ Symptomatic tr	eatment only							
	☐ Hospitalization, inlcu	ding prolonged hospitali			ning reaction						
	☐ Disability and/or inca	·	_	maly or birth defec	` '						
	Other medically impo	rtant conditions	Death	Unknown or	not stated						
	Evidence is clearly in the second sec	osures to the pathogen favor of a cause other the ridence beyond reasonal en the adverse reaction gen in the transfused congen in the donor at the tigen in an additional compen in an additional recipitified pathogen strains a confidence (p<0.05). In the strains are the recipient was not infected with the recipient was infected with the recipient was infected.	could be identified an transfusion, but the transfusion and the transfusion and the transfusion and the transfusion. In the pathogen preceded with this parecise and the transfusion with the pathogen at the transfusion with the pathogen preceded with this parecise and transfusion with the pathogen at the transfusion with the pathogen preceded with this parecise and transfusion with the pathogen preceded with this parecise and transfusion and	ed in the recipient. But transfusion can Buse other than the Bion is unknown or Beame donation. Becular or extended Boathogen at the time Bime of donation. Byere negative for the Byerior to transfusion	transfusion. not stated. e donation. d phenotypic comparison ne of transfusion nis pathogen.						
Mc	odule-generated Designa	<u> </u>									
	TE: Designations for case de		putability will be a	automatically assig	ned in the NHSN						
	olication based on responses										
	*Do you agree with the <u>case definition</u> designation?										
	*Do you agree with the se	verity designation?		□YES	Пио						



^Please indicate your designation									
_	agree with the <u>imput</u> ndicate your designat				⁄ES		NO		
Patient Trea	atment								
Did the patient receive treatment for the transfusion reaction? If yes, select treatment(s): Medication (Select the type of medication) Antipyretics Antihistamines Inotropes/Vasopressors Bronchodilator Diuretics Intravenous Immunoglobulin Intravenous steroids Corticosteroids Antibiotics Antithymocyte globulin Cyclosporin Other									
☐ Volu	me resuscitation (Intr	avenous colloid	s or crystalloids)						
 □ Respiratory support (Select the type of support) □ Mechanical ventilation □ Noninvasive ventilation □ Oxygen □ Renal replacement therapy (Select the type of therapy) □ Hemodialysis □ Peritoneal □ Continuous Veno-Venous Hemofiltration 									
☐ Phle	botomy er Specify:								
Outcome	орсску.								
*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	Component Details								
*Was a particular unit implicated in (i.e., responsible for) the adverse reaction?									
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		o	Implicat ed Unit?	
^IMPLICATED	UNIT								
	☐ ISBT-128								
:	☐ Codabar	☐ Entire unit ☐ Partial unit			□ A-	□ A+	□ B-	Y	
		mL		:	□B+ □ 0-	□ AB- □ O+	□ AB+		
	☐ ISBT-128	☐ Entire unit ☐ Partial unit mL				□ A+	□ B-	N	



<u> </u>			0- 🗆 0+	□ N/A	
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