

Hemovigilance Module Adverse Reaction Allergic Transfusion Reaction

*Required for saving

*Facility ID#: NHSN Adverse	Reaction #:
Patient Information	
*Patient ID:	*Date of Birth:/
*Sex at Birth: □M □F □Unknown	*Gender Identity (Specify):
Social Security #: Sec	condary ID: Medicare #:
Last Name: First	st Name: Middle Name:
Ethnicity Hispanic or Latino N	Not Hispanic or Not Latino
Race	ve 🗌 Asian 🔲 Black or African American
☐ Native Hawaiian/Other Pacif	fic Islander
*Blood Group: ☐ A- ☐ A+ ☐ B- ☐ E	B+
☐ Transitional ABO / Rh +	☐ Transitional ABO / Rh - ☐ Transitional ABO / Transitional Rh
☐ Group A/Transitional Rh ☐ Group B/Tran	nsitional Rh Group O/Transitional Rh Group AB/Transitional Rh
Patient Medical History	
List the patient's admitting diagnosis. (Us	se ICD-10 Diagnostic codes/descriptions)
Code: Descri	ption:
	ption:
Code: Descri	ption:
List the patient's underlying indication for	transfusion. (Use ICD-10 Diagnostic codes/descriptions)
Code: Descri	ption:
Code: Descri	ption:
Code: Descri	ption:
List the patient's comorbid conditions at t reaction. (Use ICD-10 Diagnostic codes/	the time of the transfusion related to the adverse UNKNOWN (descriptions)
Code: Descri	ption:
Code: Descri	ption:
Code: Descri	iption:
will be held in strict confidence, will be used only for the purposes stated, and 304, 306 and 308(d) of the Public Health Service Act (42 USC 242b, 242k, an	in this surveillance system that would permit identification of any individual or institution is collected with a guarantee that it will not otherwise be disclosed or released without the consent of the individual, or the institution in accordance with Sections at 242m(d)). ge 20 minutes per response, including the time for reviewing instructions, searching existing data sources, gathering, and

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 H21-8, Atlanta, GA 30333, ATTN: PRA (0920-0666).



•	medical procedure including past procedures and procedures to be ent hospital or outpatient stay. (Use ICD-10 Procedure	De UNKNOWN NONE		
Code:	Description:			
Code:				
Code:				
Additional Information				
Transfusion History				
Has the patient received a	a previous transfusion?	UNKNOWN		
Blood Product:	☐ WB ☐ RBC ☐ Platelet ☐ Plasma ☐ Cryoprecipitat	e Granulocyte		
Date of Transfusion:	/			
Was the patient's advers	se reaction transfusion-related?	Э		
If yes, provide information	on about the transfusion adverse reaction.			
Type of transfusion adve	erse reaction:	STR		
☐ HTR ☐ TTI	☐ PTP ☐ TACO ☐ TAD ☐ TA-GVHD ☐ TRALI	☐ UNKNOWN		
☐ OTHER Spec	cify			
Reaction Details				
*Date reaction occurred:	_// *Time reaction occurred: : : Tin	ne unknown		
*Facility location where pat	tient was transfused:			
Is this reaction associated with	th an incident?			
Investigation Results				
* Allergic reaction, inclu	uding anaphylaxis			
*Case Definition				
Check the following that o	occurred during or within 4 hours of cessation of transfusion:			
☐ Conjunctival edema ☐ Edema of lips, tongue and uvula ☐ Localized angioedema ☐ Hypotension				
Erythema and edema	a of the periorbital area Respiratory distress; bronchospasm	☐ Urticaria		
☐ Generalized flushing	☐ Maculopapular rash ☐ Pruritus			
Other signs and symptoms	: (check all that apply)			
Generalized:	☐ Chills/rigors ☐ Fever ☐ Nausea	/vomiting		
Cardiovascular:	Shock			
Cutaneous:	☐ Jaundice			
Hemolysis/Hemorrhage:	☐ Disseminated intravascular coagulation ☐ Hemogl ☐ Positive antibody screen	obinemia		
Pain:	☐ Abdominal pain ☐ Back pain ☐ Flank pain	☐ Infusion site pain		
Renal:	☐ Hematuria ☐ Hemoglobinuria ☐ Oliguria			
Respiratory:	☐ Bilateral infiltrates on chest x-ray ☐ Cough ☐ Hypoxemia ☐ Shortness of brea			
Other: (specify)				



*Severity					
Did the patient receive or experience any of the follow	owing?				
☐ No treatment required ☐ Symptomatic treatment only					
Hospitalization, inlcuding prolonged hospita	lization	Life-threatening	g reaction		
☐ Disability and/or incapacitation	Congenital anom	aly or birth defect(s	s) of the fetus		
☐ Other medically important conditions	Death	Unknown or no	ot stated		
*Imputability					
Which best describes the relationship between the	transfusion and the I	reaction?			
 □ No other evidence of environmental, drug or dietary risks. □ There are other potential causes present that could explain acute hemolysis, but transfusion is the most likely cause. □ Other present causes are most likely, but transfusion cannot be ruled out. □ Evidence is clearly in favor of a cause other than the transfusion, but transfusion cannot be excluded. 					
☐ There is conclusive evidence beyond reasona					
☐ The relationship between the adverse reaction	n and the transfusior	n is unknown or not	stated.		
Did the transfusion occur at your facility?	YES NO				
When did the reaction occur in relation to the transf Cocurred during or within 2 hours of cessation Cocurred 2 - 4 hours after cessation of transfer	of transfusion.				
Did the same reaction occur after the transfusion wa	s restarted (rechalle	nge)?	☐ YES ☐ NO		
Module-generated Designations					
	NOTE: Designations for case definition, severity, and imputability will be automatically assigned in the NHSN application based on responses in the corresponding investigation results section above.				
*Do you agree with the <u>case definition</u> designate ^Please indicate your designation	ion?	☐ YES	□ NO		
*Do you agree with the <u>severity</u> designation? ^Please indicate your designation		☐ YES	□NO		
*Do you agree with the <u>imputability</u> designation ^Please indicate your designation		☐ YES	□NO		
Patient Treatment					
Did the patient receive treatment for the transfusion of the select treatment (s): Medication (Select the type of medication) Antipyretics Antihistamines Intravenous Immunoglobulin Intravenous Immunoglobulin Cyclospo	otropes/Vasopresso	YES NO ors Bronchodila Corticosteroids	UNKNOWN ator Diuretics Antibiotics		
☐ Volume resuscitation (Intravenous colloids or	crystalloids)				
☐ Respiratory support (Select the type of support ☐ Mechanical ventilation ☐ Noninvas	ort) ive ventilation] Oxygen			



☐ Renal replacement therapy (Select the type of therapy) ☐ Hemodialysis ☐ Peritoneal ☐ Continuous Veno-Venous Hemofiltration									
☐ Phlebotomy ☐ Other Specify:									
Outcome									
*Outcome: Death Major or long-term sequelae Minor or no sequelae Not determined									
Date of Death:/									
Alf recipient died, relationship of transfusion to death:									
_	Definite	e	e [Doubtful	∏ Ruled Ou	IT	Not determin	ea	
	autopsy performed?	☐ Yes	☐ No)					
	· · ·								
*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	(Requ	number uired for ion and I)	*Unit expiration Date/Time	*Bloc	od group it	Implic ated Unit?	
^IMPLICATED	UNIT								
/	☐ ISBT-128								
:	☐ Codabar	☐ Entire unit ☐ Partial unit	 			□ A-	□ A+ □ B-	Y	
		mL	l — —			□в+	□ AB- □ AB+		
:					::	□ 0-	□ O+ □ N/A		
//	☐ ISBT-128								
:	☐ Codabar	☐ Entire unit ☐ Partial unit		/		□ A- □ A+ □ B-	□ A+ □ B-	N	
//		mL				□В+	□ AB- □ AB+		
:					:	□ o-	□ O+ □ N/A		
Custom Fields									
Label				Label					
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Comments									