

Form Approved OMB No. 0920-0666 Exp. Date: 12/31/2026 www.cdc.gov/nhsn

Hemovigilance Module Adverse Reaction Other Transfusion Reaction

*Required for saving

*Facility ID#: NHSN Adverse Reaction #:				
Patient Information				
*Patient ID:		*Date of Birth:	<i></i>	
*Sex at Birth: □M □F □Unknown		*Gender Identity (S	pecify):	
Social Security #: Secon	ndary ID:	Medicare #:		
Last Name: First Name:	Name:	Middle Name:		
Ethnicity Hispanic or Latino No	Hispanic or Not Latino			
Race	Asian [Black or African American	ı	
☐ Native Hawaiian/Other Pacific	Islander [White White		
*Blood Group: ☐ A- ☐ A+ ☐ B- ☐B+	☐ AB- ☐ AB+	O- O+ Blood	type not done	
☐ Transitional ABO / Rh +	☐ Transitional ABO /	Rh - Transitional AB	O / Transitional Rh	
☐ Group A/Transitional Rh ☐ Group B/Transit	ional Rh 🔲 Group O/T	ransitional Rh	3/Transitional Rh	
Patient Medical History				
List the patient's admitting diagnosis. (Use	ICD-10 Diagnostic code	es/descriptions)		
Code: Descripti	on:			
Code: Descripti	on:			
Code: Description:				
List the patient's underlying indication for tr	ansfusion. <i>(Use ICD-10</i>	Diagnostic codes/descriptio	ns)	
Code: Descripti	on:			
Code: Descripti	on:			
Code: Description:				
List the patient's comorbid conditions at the time of the transfusion related to the adverse UNKNOWN reaction. (Use ICD-10 Diagnostic codes/descriptions)				
Code: Descripti			· 	
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 H21-8, Atlanta, GA 30333, ATTN: PRA (0920-0666).				



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List the patient's relevant medical procedure including past procedures and procedures to be performed during the current hospital or outpatient stay. (Use ICD-10 Procedure codes/descriptions)				
Code:	_ Description:			
Code:				
Code:	_ Description:			
Additional Information				
Transfusion History				
Has the patient received a	previous transfusion?			
Blood Product:] WB ☐ RBC ☐ Platelet ☐ Plasma ☐ Cryoprecipitate ☐ Gra	nulocyte		
Date of Transfusion:	/			
Was the patient's advers	se reaction transfusion-related?			
If yes, provide informatio	n about the transfusion adverse reaction.			
	erse reaction:	HTR		
☐ HTR ☐ TTI	☐ PTP ☐ TACO ☐ TAD ☐ TA-GVHD ☐ TRALI ☐ UNK	NOWN		
OTHER Spec	ify			
Reaction Details				
*Date reaction occurred:/ *Time reaction occurred:: Time unknown *Facility location where patient was transfused: Is this reaction associated with an incident? Yes No If Yes, Incident #:				
Investigation Results				
Investigation Results * Other				
* Other				
* Other				
* Other Specify: List tests relevant to reac				
* Other Specify: List tests relevant to reac Test name:	tion investigation:			
* Other Specify: List tests relevant to reac Test name: Test name:	tion investigation: Testing date: Test result: Test result:			
* Other Specify: List tests relevant to reac Test name: Test name: Other signs and symptoms:	tion investigation: Testing date: Test result: Testing date: Test result: (check all that apply)			
* Other Specify: List tests relevant to reac Test name: Test name: Other signs and symptoms: Generalized:	tion investigation: Testing date: Testing date: Test result: Test result: (check all that apply) Chills/rigors Fever Nausea/vomiting			
* Other Specify: List tests relevant to react Test name: Test name: Other signs and symptoms: Generalized: Cardiovascular:	tion investigation: Testing date: Test result: Testing date: Test result: (check all that apply)			
* Other Specify: List tests relevant to reac Test name: Test name: Other signs and symptoms: Generalized:	tion investigation: Testing date: Testing date: Test result: Test result: (check all that apply) Chills/rigors Fever Blood pressure decrease Shock			
* Other Specify: List tests relevant to react Test name: Test name: Other signs and symptoms: Generalized: Cardiovascular: Cutaneous:	tion investigation: Testing date: Testing date: Test result: Test result: (check all that apply) Chills/rigors Fever Nausea/vomiting Blood pressure decrease Shock Bedema Jaundice			
* Other Specify: List tests relevant to react Test name: Test name: Other signs and symptoms: Generalized: Cardiovascular:	tion investigation: Testing date: Testing date: Test result: Test result: (check all that apply) Chills/rigors Fever Nausea/vomiting Blood pressure decrease Shock Edema Flushing Jaundice Other rash Pruritus (itching) Urticaria (hives)			
* Other Specify: List tests relevant to react Test name: Test name: Other signs and symptoms: Generalized: Cardiovascular: Cutaneous:	tion investigation: Testing date: Testing date: Test result: Test result: (check all that apply) Chills/rigors Fever Nausea/vomiting Blood pressure decrease Shock Edema Flushing Jaundice Other rash Pruritus (itching) Urticaria (hives) Biood pressure decrease Hemoglobinemia			
* Other Specify: List tests relevant to reach Test name: Test name: Other signs and symptoms: Generalized: Cardiovascular: Cutaneous: Hemolysis/Hemorrhage:	tion investigation: Testing date: Testing date: Testing date: Test result: (check all that apply) Chills/rigors Fever Nausea/vomiting Blood pressure decrease Shock Edema Flushing Jaundice Other rash Pruritus (itching) Urticaria (hives) Disseminated intravascular coagulation Hemoglobinemia Positive antibody screen			
* Other Specify: List tests relevant to reach Test name: Test name: Other signs and symptoms: Generalized: Cardiovascular: Cutaneous: Hemolysis/Hemorrhage: Pain: Renal:	tion investigation: Testing date: Testing date: Testing date: Test result: (check all that apply) Chills/rigors Fever Nausea/vomiting Blood pressure decrease Shock Edema Flushing Jaundice Other rash Pruritus (itching) Urticaria (hives) Disseminated intravascular coagulation Positive antibody screen Abdominal pain Back pain Flank pain Infusion			
* Other Specify: List tests relevant to reach Test name: Test name: Other signs and symptoms: Generalized: Cardiovascular: Cutaneous: Hemolysis/Hemorrhage: Pain: Renal: Respiratory:	tion investigation: Testing date: Testing date: Testing date: Test result: (check all that apply) Chills/rigors Fever Nausea/vomiting Blood pressure decrease Shock Edema Flushing Jaundice Other rash Pruritus (itching) Urticaria (hives) Disseminated intravascular coagulation Hemoglobinemia Positive antibody screen Abdominal pain Back pain Flank pain Infusion services.			
* Other Specify: List tests relevant to reach Test name: Test name: Other signs and symptoms: Generalized: Cardiovascular: Cutaneous: Hemolysis/Hemorrhage: Pain: Renal:	tion investigation: Testing date: Testing date: Testing date: Test result: (check all that apply) Chills/rigors Fever Nausea/vomiting Blood pressure decrease Shock Edema Flushing Jaundice Other rash Pruritus (itching) Urticaria (hives) Disseminated intravascular coagulation Hemoglobinemia Positive antibody screen Abdominal pain Back pain Flank pain Infusion: Hematuria Hemoglobinuria Oliguria Bilateral infiltrates on chest x-ray Bronchospasm Cough			



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*Severity			
Did the patient receive or experience any of the	following?		
☐ No treatment required	☐ Symptomatic tre	atment only	
Hospitalization, inlcuding prolonged hospi	talization	Life-threatening	reaction
☐ Disability and/or incapacitation	☐ Congenital anon	naly or birth defect(s)	of the fetus
Other medically important conditions	☐ Death	Unknown or not	stated
*Imputability			
Which best describes the relationship between the			
Conclusive evidence exists that the advers			ion.
☐ Evidence is clearly in favor of attributing the			
Evidence is indeterminate for attributing the			
Evidence is clearly in favor of a cause otheThere is conclusive evidence beyond reason			
☐ The relationship between the adverse reac			
<u> </u>	YES NO	on is unknown or not	siaica.
Module-generated Designations NOTE: Designations for case definition, severity, and	l imputability will be au	utomatically assigned	I in the NHSN
application based on responses in the corresponding			
*Do you agree with the <u>case definition</u> designation	nation?	YES	□ NO
*Do you agree with the <u>severity</u> designation? ^Please indicate your designation		YES	□NO
*Do you agree with the <i>imputability</i> designati	on?	□YES	□ NO
^Please indicate your designation			
Patient Treatment			
Did the patient receive treatment for the transfusio	n reaction?	YES NO	UNKNOWN
If yes, select treatment(s):			
☐ Medication (Select the type of medication)			
	Inotropes/Vasopress		
☐ Intravenous Immunoglobulin ☐ Intra		Corticosteroids	Antibiotics
Antithymocyte globulin Cyclos	porin		
☐ Volume resuscitation (Intravenous colloids	or crystalloids)		
☐ Respiratory support (Select the type of sup ☐ Mechanical ventilation ☐ Noninv	<i>port)</i> asive ventilation [Oxygen	
☐ Renal replacement therapy <i>(Select the type</i> ☐ Hemodialysis ☐ Peritoneal ☐	e of therapy) Continuous Veno-Ve	nous Hemofiltration	
☐ Phlebotomy ☐ Other Specify:			





Outcome								
*Outcome: Death Maj	jor or long-tern	n sequ	elae 🗌	Minor or no seq	uelae	☐ No	t determ	nined
Date of Death:/								
^If recipient died, relations	hip of transfus	ion to	death:					
☐ Definite ☐ Probable	☐ Possibl	е [Doubtful	Ruled Out		Not de	etermine	ed
Cause of death:						_		
Was an autopsy performed?	☐ Yes	☐ No						
Component Details								
*Was a particular unit implicated in (i.e., responsible for) the adverse Pes No N/A								
Transfusion	Amount		number ired for	*Unit				Implic
Start and End *Component code t	transfused at In		on and	expiration	*Blood group		ated	
Date/Time (check system used) r	reaction onset TRALI) Date/Time of unit					Unit?		
^IMPLICATED UNIT				T	ı	T	I	
/ 🔲 ISBT-128								
:	☐ Entire unit ☐ Partial unit				□ A-	□ A+	□ B-	Υ
	mL				B+	☐ AB-	П ав+	ı
					По-	□ /\b	□ N/A	
				·	□ 0-	<u> </u>	∐ IN/A	
/	Entire unit							
Codabar	☐ Partial unit			//	□ A-	□ A+	□ B-	N
	mL				□в+	☐ AB-	☐ AB+	
				:	□ 0-	□ 0+	□ N/A	
Custom Fields								
Label			Label					
		-				/		
Comments								

