

## Hemovigilance Module Adverse Reaction Acute Hemolytic Transfusion Reaction

\*Required for saving \*Facility ID#: \_\_\_\_\_ NHSN Adverse Reaction #: Patient Information Social Security #: Secondary ID: \_\_\_\_\_ Medicare #: \_\_\_\_\_ First Name: Middle Name: Last Name: ☐ Hispanic or Latino ☐ Not Hispanic or Not Latino Ethnicity American Indian/Alaska Native ☐ Asian ☐ Black or African American Race ☐ 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) Code: \_\_\_\_\_ Description: Code: Description: Code: Description: List the patient's underlying indication for transfusion. (Use ICD-10 Diagnostic codes/descriptions) Code: \_\_\_\_ Description: Code: \_\_\_\_\_ Description: Code: Description: 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: 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).



	dical procedure including past procedures and procedures to be UNKNOWN				
codes/descriptions)	hospital or outpatient stay. (Use ICD-10 Procedure				
Code:	Description:				
Code:	Description:				
Code:	Description:				
Additional Information					
Transfusion History					
Has the patient received a pr	revious transfusion?				
Blood Product:	WB ☐ RBC ☐ Platelet ☐ Plasma ☐ Cryoprecipitate ☐ Granulocyte				
Date of Transfusion:	//				
Was the patient's adverse	reaction transfusion-related?				
If yes, provide information	about the transfusion adverse reaction.				
Type of transfusion advers	e reaction:				
	] PTP   TACO   TAD   TA-GVHD   TRALI   UNKNOWN				
☐ OTHER Specify					
Reaction Details					
*Date reaction occurred:/_	/ *Time reaction occurred::				
*Facility location where patier	nt was transfused:				
Is this reaction associated with an incident?   Yes  No  If Yes, Incident #:					
	,				
Investigation Results					
Investigation Results * Acute hemolytic transfus					
	sion reaction (AHTR)				
* Acute hemolytic transfus	sion reaction (AHTR)				
* Acute hemolytic transfus    Immune   Antibody:     *Case Definition	sion reaction (AHTR)				
* Acute hemolytic transfus  Immune Antibody:  *Case Definition Check the following that or	sion reaction (AHTR)  Non-immune (specify)				
* Acute hemolytic transfus  Immune Antibody:  *Case Definition  Check the following that or  Back/flank pain	sion reaction (AHTR)  Non-immune (specify)  ccurred during, or within 24 hours of cessation of transfusion with <i>new</i> onset:				
* Acute hemolytic transfus  Immune Antibody:  *Case Definition  Check the following that or  Back/flank pain Oliguria/anuria	sion reaction (AHTR)  Non-immune (specify)  curred during, or within 24 hours of cessation of transfusion with <i>new</i> onset:  Chills/rigors				
* Acute hemolytic transfus  Immune Antibody:  *Case Definition  Check the following that or  Back/flank pain	Sion reaction (AHTR)  Non-immune (specify)  Ccurred during, or within 24 hours of cessation of transfusion with <i>new</i> onset:  Chills/rigors				
* Acute hemolytic transfus    Immune   Antibody:     *Case Definition   Check the following that or     Back/flank pain   Gransfus     Oliguria/anuria     Pain and/or oozing at I	Sion reaction (AHTR)  Non-immune (specify)  Courred during, or within 24 hours of cessation of transfusion with <i>new</i> onset:  Chills/rigors				
* Acute hemolytic transfus    Immune   Antibody:     *Case Definition   Check the following that or     Back/flank pain     Oliguria/anuria     Pain and/or oozing at I   Check all that apply:       Elevated LDH   He	Sion reaction (AHTR)  Non-immune (specify)  Ccurred during, or within 24 hours of cessation of transfusion with new onset: Chills/rigors				
* Acute hemolytic transfus    Immune Antibody:	Sion reaction (AHTR)  Non-immune (specify)  Courred during, or within 24 hours of cessation of transfusion with <i>new</i> onset:  Chills/rigors				
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* Acute hemolytic transfus    Immune Antibody:	sion reaction (AHTR)  Non-immune (specify)  ccurred during, or within 24 hours of cessation of transfusion with <i>new</i> onset: Chills/rigors   Epistaxis   Disseminated intravascular coagulation (DIC)   Hypotension   Fever   Hematuria (gross visual hemolysis)  V site   Renal failure  Decreased fibrinogen   Decreased haptoglobin   Elevated bilirubin moglobinemia   Hemoglobinuria   Plasma discoloration c/w hemolysis film   Positive direct antiglobulin test (DAT) for anti-IgG or anti-C3 th alloantibody present on the transfused red blood cells gative, and physical cause (e.g., thermal, osmotic, mechanical, chemical) is				
* Acute hemolytic transfus    Immune	sion reaction (AHTR)  Non-immune (specify)  curred during, or within 24 hours of cessation of transfusion with <i>new</i> onset:  Chills/rigors   Epistaxis   Disseminated intravascular coagulation (DIC)   Hypotension   Fever   Hematuria (gross visual hemolysis)  V site   Renal failure  Decreased fibrinogen   Decreased haptoglobin   Elevated bilirubin moglobinemia   Hemoglobinuria   Plasma discoloration c/w hemolysis film   Positive direct antiglobulin test (DAT) for anti-IgG or anti-C3 th alloantibody present on the transfused red blood cells gative, and physical cause (e.g., thermal, osmotic, mechanical, chemical) is uded but serologic evidence is not sufficient to meet definitive criteria.				
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Cutaneous:	☐ Edema	☐ Flushing	☐ Jaundice	`			
	Other rash	Pruritus (itching)	Urticaria (hiv	es)			
Hemolysis/Hemorrhage:	Hemoglobinemia Positive antibody screen						
Pain:	Abdominal pain						
Respiratory:    Bilateral infiltrates on chest x-ray   Bronchospasm   Cough     Shortness of breath   Hypoxemia							
Other: (specify)							
*Severity							
Did the patient receive or ex	perience any of the followi	ng?					
☐ No treatment require	ed 🔲 :	Symptomatic treatment	only				
☐ Hospitalization, inlcu	ıding prolonged hospitaliza	ation Life	e-threatening reaction	n			
☐ Disability and/or inca	apacitation	Congenital anomaly or b	oirth defect(s) of the	fetus			
Other medically impo	ortant conditions	Death Un	known or not stated				
*Imputability							
Which best describes the rel	ationship between the tran	sfusion and the reaction	1?				
☐ ABO or other allotypic	RBC antigen incompatibility	ty is known.					
	•	•	molysis is present.				
<ul> <li>Only transfusion-related (i.e., immune or non-immune) cause of acute hemolysis is present.</li> <li>There are other potential causes present that could explain acute hemolysis, but transfusion is the most likely cause.</li> </ul>							
	hemolysis are more likely,	but transfusion cannot	be ruled out.				
	avor of a cause other than			xcluded.			
<u> </u>	ridence beyond reasonable	·					
	een the adverse reaction ar						
Did the transfusion occur at							
		5 110					
Module-generated Designations for case de		ıtahility will he automatic	rally assigned in the	NHSN			
application based on responses				1111011			
*Do you agree with the <u>ca</u>		_		NO			
^Please indicate your design			0				
*Do you agree with the se		Γ	∃YES □□	NO			
^Please indicate your design		L	_ 125	110			
*Do you agree with the in		Г	☐YES ☐	NO			
^Please indicate your design		L	_ 1631	NO			
Patient Treatment							
				UKNIOVA/NI			
Did the patient receive treatm	ient for the transfusion read	ction? YES	□ NO □ UN	IKNOWN			
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							
☐ Volume resuscitation (Intravenous colloids or crystalloids)							



<ul> <li>☐ Respiratory support (Select the type of support)</li> <li>☐ Mechanical ventilation</li> <li>☐ Noninvasive ventilation</li> <li>☐ Oxygen</li> </ul>												
☐ Renal replacement therapy (Select the type of therapy) ☐ Hemodialysis ☐ Peritoneal ☐ Continuous Veno-Venous Hemofiltration												
☐ Phlebotomy ☐ Other Specify:												
Outcome												
*Outcome: Date of	Death:/_	ajor or long-terr	·		Minor or no seq	uelae	☐ Not	determ	nined			
	recipient died, relation	•		<u></u>		_	1					
_	Definite Probable	e  Possibl	le [	Doubtful	☐ Ruled Out		Not det	termine	ed			
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 '(Required for Infection and reaction onset TRALI)		*Unit expiration Date/Time	*Blood group of unit		Implic ated Unit?					
^IMPLICATED	UNIT											
/	☐ ISBT-128											
:	☐ Codabar	☐ Entire unit ☐ Partial unit				□ A-	□ A+ □	☐ B-	Y			
/		mL				B+	□ AB- □	☐ AB+	Ť			
:					:	П 0-		□ N/A				
//	☐ ISBT-128											
:	☐ Codabar	☐ Entire unit ☐ Partial unit			/	□ A-	□ A+ □	☐ B-	N			
//		mL				□в+	□ AB-	☐ AB+				
:					:	□ o-	□ O+ [	□ N/A				
Custom Field	ds											
Label				Label								
							/	_/				
Comments												