

Form Approved OMB No. 0920-0666

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## **Hemovigilance Module Adverse Reaction Acute Hemolytic Transfusion Reaction**

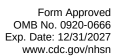
\*Required for saving NHSN Adverse Reaction #: \_\_\_\_\_ \*Facility ID#: \_\_\_\_\_ **Patient Information** \*Patient ID: \*Date of Birth: \_\_\_/\_\_/\_\_ \*Sex: M Secondary ID: Social Security #: Medicare #: First Name: \_\_\_\_\_ Last Name: Middle Name: Not Hispanic or Ethnicity (Specify): Unknown Declined to respond Hispanic or Latino Latino American Indian or Asian Black or African Middle Eastern or North Race (Select all that Alaska Native American African apply): White Native Hawaiian or Unknown Declined to respond Pacific Islander Interpreter Needed: Yes No Preferred Language (Specify from the list provided): Unknow Declined to Respond n \*Blood Group: O+Blood type not done Transitional ABO / Transitional Transitional ABO / Rh + Transitional ABO / Rh -Group A/Transitional 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: 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 UNKNOWN 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)). CDC 57.307 Rev. 3, v9.2

Public reporting burden of this collection of information is estimated to average 22 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:









	edical procedure including past procedures and procedures to be UNKN hospital or outpatient stay. (Use ICD-10 Procedure NONE	
Code:	Description:	
Code:	Description:	
Code:	Description:	
Transfusion History		
	revious transfusion? YES NO UNKNOWN	
Blood Product:	WB RBC Platelet Plasma Cryoprecipitate Granu	locyte
Date of Transfusion:	// UNKNOWN	
Was the patient's adverse เ	reaction transfusion-related? YES NO	
If yes, provide information a	about the transfusion adverse reaction.	
Type of transfusion adverse	e reaction: Allergic AHTR DHTR DSTR FNHT	R
HTR TTI	PTP TACO TAD TA-GVHD TRALI UNKNO	NWC
OTHER Specify	·	
Reaction Details		
*Date reaction occurred:/_	/ *Time reaction occurred:: Time unknown	
*Facility location where patien	nt was transfused:	
Is this reaction associated with a	an incident? Yes No If Yes, Incident #:	
Investigation Results		
* Acute hemolytic transfus		
Immune Antibody: _	Non-immune (specify)	_
*Case Definition		
	ccurred <b>during, or within 24 hours</b> of cessation of transfusion with <i>new</i> onse	
	Chills/rigors	.)
Oliguria/anuria		
Pain and/or oozing at I	IV site Renal failure	
Check all that apply:	Decreased fibrinogen Decreased haptoglobin Elevated bilirubin	
Elevated LDH He	emoglobinemia 🗌 Hemoglobinuria 🔲 Plasma discoloration c/w hemolysis	;
Spherocytes on blood	Positive direct antiglobulin test (DAT) for anti-IgG or anti-C3	
film  Resitive elution test with		
	th alloantibody present on the transfused red blood cells gative, and physical cause (e.g., thermal, osmotic, mechanical, chemical) is	
confirmed.	gative, and physical cause (e.g., thermal, esmolie, mechanical, enemical) is	
Physical cause is exclu	uded but serologic evidence is not sufficient to meet definitive criteria.	
Physical cause is susp	pected and serologic testing is negative.	
AHTR is suspected, bu	ut symptoms, test results, and/or information are not sufficient to confirm react	ion.
Other signs and symptoms: (	check all that apply)	
Generalized:	Nausea/vomiting	
Cardiovascular:	Shock	



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Cutaneous:		Edema	Flushin	ng	Jaur	ndice
		Other rash	Pruritus	s (itching)	Urtic	caria (hives)
Hemolysis/Hemorrhage:	Ļ	Hemoglobinemia	Positive	e antibody scree	en	
Pain:	Ļ	Abdominal pain				
	ıL	Bilateral infiltrates on che	est x-	Bronchosp	asm	Cough
Respiratory:	ray	٦				
	L	Shortness of breath		Hypoxemia	<u> </u>	
Other: (specify)						
*Severity						
Did the patient receive or ex	per	ience any of the following?	<b>,</b>			
No treatment require	ed	Sy	mptomat	tic treatment onl	У	
Hospitalization, inlc	udin	g prolonged hospitalizatio	n	Life-tl	nreatenin	g reaction
Disability and/or inc	ара	citation $oxedsymbol{\Box}$ Co	ngenital	anomaly or birtl	n defect(s	s) of the fetus
Other medically imp	orta	nt conditions De	ath	Unkn	own or no	ot stated
*Imputability						
Which best describes the rel	atio	nship between the transfu	sion and	d the reaction?		
ABO or other allotypic	RB	C antigen incompatibility i	s known	١.		
Only transfusion-relate	ed (	i.e., immune or non-immui	ne) caus	se of acute hemo	olysis is p	resent.
= '	•	causes present that could	•			
	he	molysis are more likely, bu	ıt transfu	usion cannot be	ruled out	`.
		or of a cause other than the				
		nce beyond reasonable do				
		the adverse reaction and				
					WIT OF THO	. Statea.
Did the transfusion occur at	you	r facility? YES	N	10		
Module-generated Design						
NOTE: Designations for case de application based on responses						d in the NHSN
*Do you agree with the <u>ca</u>	ıse	definition designation?			YES	NO
^Please indicate your desig	gnat	ion				
*Do you agree with the <u>se</u>	ever	rity designation?			YES	NO
^Please indicate your design		•				
*Do you agree with the <u>in</u>	ามน	tability designation?			YES	NO
^Please indicate your design	•	•				
Patient Treatment						
Did the patient receive treatm	t	for the transfusion reaction		YES	NO	UNKNOWN
If yes, select treatment(s):	iciit	Tor the transitional reaction				
Medication (Select th	e ty	pe of medication)				
Antipyretics	<b>┐</b> ∧.	 ntihistamines Inotropes/\	/acopros	seore Dr	onchodila	ator Diuretics
Intravenous	_ <b>~</b> I	innstannies motropes/V	ασυμιθδ	33UI3 DI	oriciloulla	
Immunoglobulin		Intravenous	steroids	S Corticos	steroids	Antibiotics

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SAFETY N	Antithymocyte glob	ulin Cycle	osporin	Other	www.cdc.go	
	_ , , ,			Julei		
Vol	ume resuscitation (Int	ravenous colloid	ls or crystalloids)			
Res	spiratory support (Sele					
L	Mechanical ventilat	ion Noni	nvasive ventilation	Oxygen		
Rer	nal replacement thera	oy (Select the ty	_			
	Hemodialysis	Peritoneal	Continuous Ver	no-Venous Hem	ofiltration	
Phle	ebotomy					
Oth	er Specify:					
Outcome				Naissau su sa		
*Outcome:	Death M	lajor or long-terr	n sequelae se	」Minor or no equelae	Not deteri	mined
Date of			·	•		
^If	recipient died, relation	. —				
	Definite Probabl	e Possib	le Doubtful	Ruled Ou	t Not determin	ed
	of death:		□ N.a		<del></del>	
	autopsy performed?	Yes	No			
*Was a parti	t Details cular unit implicate	din (i.a. raen	neible for) the s	dverse		
reaction?	cular unit implicate	u III (I.e., Tespi		uvei se	Yes No	N/A
Transfusion		Amount	^Unit number (Required for	*Unit		Implic
Start and End Date/Time	*Component code (check system used)	transfused at reaction onset	Infection and TRALI)	expiration Date/Time	*Blood group of unit	ated Unit?
^IMPLICATED		reaction onset	INALI)	Daterrine	or unit	Oint.
1 1	ISBT-128					
:	Codabar	Entire unit		, ,	A- A+ B-	
		Partial unit				Y
		mL			B+ AB- AB+	
:	LODE 100			::	O- O+ N/A	
	ISBT-128 Codabar	Entire unit			A- A+ B-	
:	Codabar	Partial unit				N
/		mL			B+ AB- AB+	
:				<u> </u>	O- O+ N/A	
Custom Field	ds					
Label			Label			
		<u> </u>	-			<del></del>
Comments						



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