

Form Approved OMB No. 0920-0666 Exp. Date: 12/31/2027 www.cdc.gov/nhsn

Hemovigilance Module Adverse Reaction Delayed Hemolytic Transfusion Reaction

*Required for saving

*Facility ID#:	NHSN Adverse Reaction #:					
Patient Informatio						
*Patient ID: *Date of Birth:/						
*Sex: M F						
Social Security #:	Secondary ID: Medicare #:					
Last Name:	First Name: Middle Name:					
Ethnicity (Specify):	Hispanic or Latino Not Hispanic or Latino Unknown Declined to respond					
Race (Select all that apply):	American Indian or Asian Black or African Middle Eastern or North Alaska Native American African Native Hawaiian or White Unknown Declined to respond Pacific Islander					
Preferred Language (Specify from the list provided): Interpreter Needed: Yes Unknow Declined to Respond n						
*Blood Group: A- A+ B- B+ AB- AB+ O- O+ Blood type not done Transitional ABO / Rh + Transitional ABO / Rh - Rh Group A/Transitional Group B/Transitional Rh Group A/Transitional Rh						
Rh						
Rh Patient Medical Hi	Rh Group O/ Transitional Rif					
Patient Medical Hi	Rh Group O/ Transitional Rif					
Patient Medical Hi	story dmitting diagnosis. (Use ICD-10 Diagnostic codes/descriptions)					
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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 57.309 Rev. 3, v9.2

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).





	edical procedure including past procedures and procedure it hospital or outpatient stay. (Use ICD-10 Procedure	es to be UNKNOWN NONE
Code:	Description:	
Code:	Description:	
Code:	Description:	
Additional Information		
Transfusion History		
Has the patient received a	previous transfusion? YES NO	UNKNOWN
Blood Product:	WB RBC Platelet Plasma Cryopre	cipitate Granulocyte
Date of Transfusion:	// UNKNOWN	
Was the patient's adverse	reaction transfusion-related?	NO
If yes, provide information	about the transfusion adverse reaction.	
Type of transfusion adve	se reaction: Allergic AHTR DHTR	☐ DSTR ☐ FNHTR
HTR TTI	☐ PTP ☐ TACO ☐ TAD ☐ TA-GVHD ☐ TR	ALI UNKNOWN
OTHER Speci	y	
Reaction Details		
*Date reaction occurred:	/ *Time reaction occurred::	Time unknown
*Facility location where pati	ent was transfused:	
Is this reaction associated with	an incident? Yes No If Yes, Incider	nt #:
Investigation Results (Or	ly answer questions listed under the selected reaction type	e.)
* Delayed hemolytic trans	fusion reaction (DHTR)	•
	Non-immune (specify)	
-		
*Case Definition		
Check the following that	occurred between 24 hours and 28 days after cessation of	of transfusion:
Positive direct antigle	bulin test (DAT)	
Newly-identified red	olood cell alloantibody in recipient serum	
Positive elution test v	ith alloantibody present on the transfused red blood cells	
Inadequate rise of pos	t-transfusion hemoglobin level or rapid fall in hemoglobin bac	k to pre-transfusion levels
Otherwise unexplain	ed appearance of spherocytes	
Check all that apply:		
Incomplete laborator	v evidence	
	out reported symptoms, test results, and/or available inforr	nation are not sufficient
Other signs and symptoms: (c		
Generalized:	Chills/rigors Fever	Nausea/vomiting
Cardiovascular:	Blood pressure decrease Shock	
Cutaneous:	Edema Flushing	Jaundice
Hemolysis/Hemorrhage:	Other rash Pruritus (itching)	Urticaria (hives)
	Disseminated intravascular coagulation	Hemoglobinemia



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Pain:		Abdominal pain		ack pain		Flank pain			Infusion site pain
Renal:		Hematuria He		Hemoglo	binu	oinuria		ligu	ria
Respiratory:		Bilateral infiltrates on chest x-ray				Bronchospasr	n		Cough
		Hypoxemia		Shortnes	ss of	f breath			
Other: (specify)									
*Severity									
Did the patient receive or e	expe	rience any of the foll	owing	g?					
No treatment requi	red		S	ymptomatic	: trea	atment only			
Hospitalization, inle	cudi	ng prolonged hospita				Life-threat	eni [.]	na r	eaction
Disability and/or inc					nom	aly or birth def		_	
			\equiv	eath	11011	Unknown		. ,	
Other medically im	ipori	ant conditions		ean		OTIKITOWIT	ו וכ	iot s	stateu
*Imputability									
Which best describes the re	alati	onshin hetween the	tranef	usion and t	ho r	eaction?			
		•							
No other explanation					-	•			fucionio the mode
An alternate explanat likely cause.	tion	for symptoms or nev	viy-iae	entified anti	מסמי	y is present, bu	t tra	ans	usion is the most
Other explanations fo	nr ev	mntoms or newly-ide	≥ntifi⊝	d antihody	are	more likely, hut	t trs	nef	usion cannot he
ruled out.	лзу	imptoms of newly-lat	STILLITE	u antibouy	arc	more likely, but	, uc	11131	usion cannot be
Evidence is clearly in	fav	or of a cause other t	han th	ne transfusio	on l	hut transfusion	car	าทดเ	he excluded
There is conclusive e									
_									
The relationship betw	/eer	i the adverse reactio	n and	tne transtu	ISIO	i is unknown or	no	t Sta	atea.
Did the transficien equir of	+	ur fo cilitu ()		NO					
Did the transfusion occur at	ιyo	ur facility? YE	:5	NO					
Module-generated Design	nat	ions							
NOTE: Designations for case of			mputa	ability will be	e au	tomatically ass	ian	ed i	n the NHSN
application based on response			•	•		•			
*Do you agree with the \underline{o}		•	tion?	1		YES			NO
^Please indicate your des	signa	ation							
#D						VEC			
*Do you agree with the s						YES			NO
^Please indicate your des	igna	alion							
*Do you agree with the i	mpi	utability designation	n?			YES			NO
^Please indicate your des	_								
	g								
Patient Treatment									
Did the patient receive treat	mer	nt for the transfusion	reacti	ion?		YES N	10		UNKNOWN
If yes, select treatment(s):			_ 5.01						
Medication (Select the type of medication)									
	- •	,, a sameny							
Antipyretics		Antihistamines Inotr	opes	Vasopress	ors	Bronch	odi	lato	r Diuretics
Intravenous			•	s steroids	Γ	Corticosteroi			Antibiotics
Intraverious		Page 3					40		



SAFEIT N					www.cuc.go	V/1111511				
Immunoglobulin Antithymocyte globulin Cyclosporin Other										
Volume resuscitation (Intravenous colloids or crystalloids)										
Res	spiratory support <i>(Sele</i>		,							
	Mechanical ventilati	ion Noni	nvasive ventilation	Oxygen						
Ren	nal replacement therap Hemodialysis	oy <i>(Select the ty</i> Peritoneal		no-Venous Hemo	ofiltration					
Phle	ebotomy									
Oth	er Specify:			<u> </u>						
Outcome				1						
*Outcome: Date of		ajor or long-teri/ ship of transfus	·	∫ Minor or no quelae	Not determ	mined				
	Definite Probabl	· —		Ruled Out	Not determin	ed				
Cause	of death:		 							
Was an	autopsy performed?	Yes	No							
Component										
*Was a partion?	cular unit implicated	d in (i.e., respo	onsible for) the a	dverse	Yes No	N/A				
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 group	Implic ated Unit?				
^IMPLICATED			,			1				
1 1	ISBT-128									
;	Codabar	Entire unit			A- A+ B-					
1 1		Partial unit mL			B+ AB- AB+	Y				
					O- O+ N/A					
1 1	ISBT-128			·						
:	Codabar	Entire unit		1 1	A- A+ B-					
		Partial unit				N				
//		mL			B+ AB- AB+					
: : _										
Label	us		Label							
Label		' /	Label							
Comments			· ·							



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