

Hemovigilance Module Adverse Reaction Delayed Hemolytic Transfusion Reaction

*Required for saving		
*Facility ID#: N	HSN Adverse Reaction #:	
Patient Information		
*Patient ID:		*Date of Birth://
*Sex at Birth: M F U		*Gender Identity (Specify):
Social Security #:	Secondary ID:	Medicare #:
Last Name:	First Name:	Middle Name:
Ethnicity 🗌 Hispanic or La	tino	ino
Race 🗌 American India	an/Alaska Native 🗌 Asian	Black or African American
🗌 Native Hawaii	an/Other Pacific Islander	White
*Blood Group: 🗌 A- 🗌 A	+ 🗌 B- 🗌 B+ 🗌 AB- 🗌 AB+	O- O+ Blood type not done
Transitional	ABO / Rh +	O / Rh -
Group A/Transitional Rh] Group B/Transitional Rh 🛛 Group C	D/Transitional Rh 🛛 Group AB/Transitional Rh
Patient Medical History		
List the patient's admitting	diagnosis. (Use ICD-10 Diagnostic co	odes/descriptions)
Code:	Description:	
Code:		
Code:		
List the patient's underlying	g indication for transfusion. (Use ICD-	-10 Diagnostic codes/descriptions)
Code:	Description:	
Code:		
Code:		
	conditions at the time of the transfusi	
Code:	Description:	
Code:	Description:	
Code:	Description:	
or institution is collected with a guarant	ee that it will be held in strict confidence, will be u ent of the individual, or the institution in accordan	eillance system that would permit identification of any individual used only for the purposes stated, and will not otherwise be use with Sections 304, 306 and 308(d) of the Public Health
searching existing data sources, gather may not conduct or sponsor, and a per- number. Send comments regarding thi	ring, and maintaining the data needed, and comp son is not required to respond to a collection of ir	tes per response, including the time for reviewing instructions, obleting and reviewing the collection of information. An agency information unless it displays a currently valid OMB control lection of information, including suggestions for reducing this 0333, ATTN: PRA (0920-0666).



List the patient's relevant n performed during the curre codes/descriptions)	UNKNOWN			
Code:	_ Description:			
Code:	_ Description:			
Code:				
Additional Information				
Transfusion History				
Has the patient received a	previous transfusion?	KNOWN		
Blood Product:] WB 🗌 RBC 🔲 Platelet 🗌 Plasma 🗌 Cryoprecipitate	Granulocyte		
Date of Transfusion:				
Was the patient's advers	e reaction transfusion-related?			
If yes, provide informatio	n about the transfusion adverse reaction.			
Type of transfusion adve	rse reaction:	🗌 FNHTR		
🗌 HTR 🛛 🗌 TTI	PTP TACO TAD TA-GVHD TRALI			
OTHER Speci	ify			
Reaction Details				
*Date reaction occurred:	// *Time reaction occurred:: [] Time u	nknown		
*Facility location where pati	ient was transfused:			
Is this reaction associated wit	h an incident? 🛛 🗌 Yes 🗌 No 🛛 If Yes, Incident #:			
Investigation Results (O	nly answer questions listed under the selected reaction type.)			
	sfusion reaction (DHTR)			
• •	Non-immune (specify)			
*Case Definition				
	accurred between 24 betweend 29 days ofter acception of transfur	ion		
	occurred between 24 hours and 28 days after cessation of transfus	ion.		
Positive direct antigle				
Newly-identified red blood cell alloantibody in recipient serum				
Positive elution test with alloantibody present on the transfused red blood cells				
Inadequate rise of post-transfusion hemoglobin level or rapid fall in hemoglobin back to pre-transfusion levels				
Otherwise unexplained appearance of spherocytes				
	ed appearance of spherocytes			
Check all that apply:				
Check all that apply:				
Incomplete laborator		not sufficient		
Incomplete laborator	y evidence but reported symptoms, test results, and/or available information are	not sufficient		
 Incomplete laborator DHTR is suspected, 	y evidence but reported symptoms, test results, and/or available information are check all that apply)	not sufficient a/vomiting		
 Incomplete laborator DHTR is suspected, Other signs and symptoms: (c) 	y evidence but reported symptoms, test results, and/or available information are <u>check all that apply)</u>			
 Incomplete laborator DHTR is suspected, Other signs and symptoms: (c Generalized: 	y evidence but reported symptoms, test results, and/or available information are check all that apply) Chills/rigors Ever Nause Blood pressure decrease Shock Edema Flushing Jaundi	a/vomiting		
 Incomplete laborator DHTR is suspected, Other signs and symptoms: (c Generalized: Cardiovascular: Cutaneous: 	y evidence but reported symptoms, test results, and/or available information are check all that apply) Chills/rigors Fever Nauses Blood pressure decrease Shock Edema Flushing Jaundi Other rash Pruritus (itching) Urticar	a/vomiting ce ia (hives)		
 Incomplete laborator DHTR is suspected, Other signs and symptoms: (c Generalized: Cardiovascular: 	y evidence but reported symptoms, test results, and/or available information are check all that apply) Chills/rigors Fever Nause Blood pressure decrease Shock Edema Flushing Jaundi Other rash Pruritus (itching) Urticar Disseminated intravascular coagulation	a/vomiting		



Renal:	🗌 Hematuria 📃 Hem	noglobinuria] Oliguria	
Respiratory:	Bilateral infiltrates on chest x-ray	·	n 🗌 Cough	
	Hypoxemia Sho	rtness of breath		
Other: (specify)				
*Severity				
	experience any of the following?			
No treatment requi	— • •	natic treatment only		
	cuding prolonged hospitalization	Life-threate	-	
Disability and/or in		tal anomaly or birth defe		
Other medically im	portant conditions Death		or not stated	
*Imputability				
	elationship between the transfusion a for symptoms or newly-identified and			
	tion for symptoms or newly-identified		t transfusion is the most	
likely cause.				
Other explanations for ruled out.	or symptoms or newly-identified antib	ody are more likely, but	transfusion cannot be	
Evidence is clearly in	n favor of a cause other than the trans	sfusion, but transfusion	cannot be excluded.	
There is conclusive e	evidence beyond reasonable doubt of	f a cause other than the	transfusion.	
The relationship betw	veen the adverse reaction and the tra	ansfusion is unknown or	not stated.	
Did the transfusion occur at your facility? YES NO				
Did the transfusion occur a				
	· · · · ·			
Module-generated Desig NOTE: Designations for case	nations definition, severity, and imputability v	vill be automatically ass	igned in the NHSN	
Module-generated Desig NOTE: Designations for case	nations	vill be automatically ass	igned in the NHSN	
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NATIO			LTHC	

Ren	al replacement therap] Hemodialysis 🏾 🗍 F	-			o-Venous Hemo	ofiltratic	'n	Ū	
Phle Othe	botomy er Specify:								
Outcome									
	Death://///////_	•	ion to	death:	Minor or no seq				
Cause	Definite		e [No	_ Doubtful	Ruled Out] Not de 	etermine	ed
Component	Details								
	cular unit implicated	d in (i.e., respo			dverse	_ Yes	3	No [] N/A
Transfusion Start and End Date/Time	*Component code (check system used)	Amount transfused at reaction onset			ood group unit		Implic ated Unit?		
^IMPLICATED	UNIT								
// : //	□ ISBT-128 □ Codabar 	Entire unit Partial unit mL		·	// :	□ A- □B+ □ 0-	□ A+ □ AB- □ O+	□ B- □ AB+ □ N/A	Y
// : //	☐ ISBT-128 ☐ Codabar 	☐ Entire unit ☐ Partial unit mL		·	// :	□ A- □B+ □ 0-	□ A+ □ AB- □ O+	□ B- □ AB+ □ N/A	N
Custom Field	ds								
Label				Label					
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Comments									