

Hemovigilance Module Adverse Reaction Unknown Transfusion Reaction

*Required for saving		
*Facility ID#: NHS	N Adverse Reaction #:	
Patient Information		
*Patient ID:		*Date of Birth://
*Sex at Birth: □M □F □Unkr		*Gender Identity (Specify):
Social Security #:		
Last Name:	First Name:	Middle Name:
Ethnicity Hispanic or Latin	o 🗌 Not Hispanic or Not Latir	o
	Alaska Native 🗌 Asian /Other Pacific Islander	
*Blood Group:	□ B- □B+ □ AB- □ AB+	O- O+ Blood type not done
		/ Rh - Transitional ABO / Transitional Rh /Transitional Rh Group AB/Transitional Rh
Patient Medical History		
List the patient's admitting dia	agnosis. <i>(Use ICD-10 Diagnostic co</i>	odes/descriptions)
Code:	Description:	
Code:	Description:	
Code:	Description:	
List the patient's underlying ir	ndication for transfusion. (Use ICD-	10 Diagnostic codes/descriptions)
Code:	Description:	
Code:	Description:	
Code:	Description:	
List the patient's comorbid co reaction. <i>(Use ICD-10 Diagno</i>	• •	on related to the adverse UNKNOWN
Code:		
Code:	Description:	
Code:	Description:	
or institution is collected with a guarantee of disclosed or released without the consent Service Act (42 USC 242b, 242k, and 242c) Public reporting burden of this collection of searching existing data sources, gathering may not conduct or sponsor, and a person	that it will be held in strict confidence, will be us of the individual, or the institution in accordanc m(d)). f information is estimated to average 20 minute , and maintaining the data needed, and compl is not required to respond to a collection of inf	llance system that would permit identification of any individual sed only for the purposes stated, and will not otherwise be e with Sections 304, 306 and 308(d) of the Public Health es per response, including the time for reviewing instructions, eting and reviewing the collection of information. An agency formation unless it displays a currently valid OMB control ection of information, including suggestions for reducing this
burden to CDC, Reports Clearance Officer	; 1600 Clifton Rd., MS H21-8, Atlanta, GA 303	33, ATTN: PRA (0920-0666).

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List the patient's relevant	
	medical procedure including past procedures and procedures to be UNKNOWN rent hospital or outpatient stay. (Use ICD-10 Procedure NONE
Code:	Description:
Code:	Description:
Code:	Description:
Additional Information	
Transfusion History	
Has the patient received	a previous transfusion?
Blood Product:	WB RBC Platelet Plasma Cryoprecipitate Granulocyte
Date of Transfusion:	
Was the patient's adver	rse reaction transfusion-related?
If yes, provide informati	ion about the transfusion adverse reaction.
Type of transfusion adv	rerse reaction:
🗌 HTR 🔤 TTI	PTP TACO TAD TA-GVHD TRALI UNKNOWN
OTHER Spe	cify
Reaction Details	
*Date reaction occurred:	_// *Time reaction occurred:: Time unknown
*Facility location where pa	tient was transfused:
Is this reaction associated w	ith an incident? Yes No If Yes, Incident #:
Is this reaction associated w Investigation Results	rith an incident? Yes No If Yes, Incident #:
	rith an incident? Yes No If Yes, Incident #:
Investigation Results	rith an incident? Yes No If Yes, Incident #:
Investigation Results	
Investigation Results Diagnosis of case: List tests relevant to rea	
Investigation Results Unknown Diagnosis of case: List tests relevant to rea Test name:	action investigation:
Investigation Results Unknown Diagnosis of case: List tests relevant to rea Test name:	Iction investigation: Testing date: Test result: Testing date: Test result:
Investigation Results Unknown Diagnosis of case: List tests relevant to rea Test name: Test name: Other signs and symptom	Iction investigation: Testing date: Test result: Testing date: Test result:
Investigation Results Unknown Diagnosis of case: List tests relevant to rea Test name: Test name: Other signs and symptom	Iction investigation: Testing date: Test result: Testing date: Test result: ns: (check all that apply)
Investigation Results Unknown Diagnosis of case: List tests relevant to rea Test name: Test name: Other signs and symptom Generalized: Cardiovascular:	Inction investigation:
Investigation Results Unknown Diagnosis of case: List tests relevant to rea Test name: Test name: Other signs and symptom Generalized:	Inction investigation:
Investigation Results Investigation Results Image: Image	Inction investigation:
Investigation Results Unknown Diagnosis of case: List tests relevant to rea Test name: Test name: Other signs and symptom Generalized: Cardiovascular:	Inction investigation:
Investigation Results Investigation Results Image: Image	Inction investigation:
Investigation Results Unknown Diagnosis of case: List tests relevant to rea Test name: Test name: Other signs and symptom Generalized: Cardiovascular: Cutaneous: Hemolysis/Hemorrhage:	Inction investigation: Testing date: Test result:
Investigation Results Unknown Diagnosis of case: List tests relevant to real Test name: Test name: Other signs and symptom Generalized: Cardiovascular: Cutaneous: Hemolysis/Hemorrhage: Pain: Renal:	Inction investigation:
Investigation Results Investigation Results Unknown Diagnosis of case: List tests relevant to real Test name: Test name: Other signs and symptom Generalized: Cardiovascular: Cutaneous: Hemolysis/Hemorrhage: Pain:	Inction investigation: Testing date: Test result:
Investigation Results Unknown Diagnosis of case: List tests relevant to real Test name: Test name: Other signs and symptom Generalized: Cardiovascular: Cutaneous: Hemolysis/Hemorrhage: Pain: Renal:	Inction investigation:

Did the patient receive or experience any of the following?

NATIONAL HEALTHCARE SAFETY NETWORK		Form Approved OMB No. 0920-0666 Exp. Date: 12/31/2026 www.cdc.gov/nhsn
No treatment required Symptomatic treatment	ent only	
Hospitalization, inlcuding prolonged hospitalization	Life-threatenin	g reaction
Disability and/or incapacitation Congenital anomaly	or birth defect(s	s) of the fetus
Other medically important conditions	Unknown or no	ot stated
*Imputability		
Which best describes the relationship between the transfusion and the rea Conclusive evidence exists that the adverse reaction can be attribute Evidence is clearly in favor of attributing the adverse reaction to the t	ed to the transfu	ision.
Evidence is indeterminate for attributing the adverse reaction to the t	ransfusion or a	n alternate cause.
Evidence is clearly in favor of a cause other than the transfusion, but	transfusion ca	nnot be excluded.
There is conclusive evidence beyond reasonable doubt of a cause of	ther than the tra	ansfusion.
The relationship between the adverse reaction and the transfusion is	unknown or no	ot stated.
Did the transfusion occur at your facility?		
Module-generated Designations		
NOTE: Designations for case definition, severity, and imputability will be autom		ed in the NHSN
application based on responses in the corresponding investigation results sect	ion above.	
* Do you agree with the <u>case definition</u> designation? ^Please indicate your designation	☐ YES	
*Do you agree with the <u>severity</u> designation? ^Please indicate your designation	☐ YES	
*Do you agree with the <i>imputability</i> designation? ^Please indicate your designation	☐ YES	□ NO
Patient Treatment		
Did the patient receive treatment for the transfusion reaction? YE If yes, select treatment(s): Medication (Select the type of medication) Antipyretics Antihistamines Inotropes/Vasopressors Intravenous Immunoglobulin Intravenous steroids C Antithymocyte globulin Cyclosporin Other		UNKNOWN
Volume resuscitation (Intravenous colloids or crystalloids)		
 Respiratory support (Select the type of support) Mechanical ventilation Noninvasive ventilation C 	Dxygen	
Renal replacement therapy (Select the type of therapy) Hemodialysis Peritoneal Continuous Veno-Venous	s Hemofiltration	
Phlebotomy Other Specify:		
Outcome		

NATIONAL H							Exp. D	Form App 3 No. 0920 Date: 12/31 ww.cdc.gov	-0666 /2026
*Outcome:	*Outcome: Death Major or long-term sequelae Minor or no sequelae Not determined							nined	
Date of Death://									
^lf ı	recipient died, relation	ship of transfus	ion to	death:					
	Definite Probable	e 🗌 Possibl	e [Doubtful	Ruled Out	t 🗌] Not de	etermine	ed
	of death:								
Was an	autopsy performed?	_ Yes)					
Component									
*Was a partion reaction?	cular unit implicate	d in (i.e., respo	onsib	e for) the a	adverse	🗌 Yes	s 🗌	No [] N/A
				number					
Transfusion Start and End	*Component code	Amount transfused at	(Required for		*Unit expiration *Blood group		n	Implic ated	
Date/Time	(check system used)	reaction onset	TRAL		Date/Time	of un		۲	Unit?
^IMPLICATED	UNIT						-		•
//	□ ISBT-128			·					
:	Codabar	Entire unit				🗆 A-	🗆 A+	🗌 В-	
		Partial unit mL				□в+	🗆 AB-	🗆 AB+	Y
· · · ·							□ 0+		
·				·	· ·				
//	ISBT-128	Entire unit		·		🗆 A-	□ A+	🗆 В-	
i	🗌 Codabar	Partial unit			//				N
//		mL		·		□В+	🗌 АВ-	🗌 AB+	
:				. <u> </u>	l:	0-	0+	□ N/A	
Custom Field	ds								
Label				Label					
		//	-				/	/	
Comments									