

Hemovigilance Module Adverse Reaction Other Transfusion Reaction

*Requi	red for saving					
*Facility ID)#: NHSN Ad	verse Reaction #:				
	nformation					
*Patient ID:		*Gender: M F Other *Date of Birth: / /				
	curity #:	Secondary ID:				
): 	First Name: Middle Name:				
Ethnicity	Hispanic or Latino	Not Hispanic or Not Latino				
Race	American Indian/Alask	a Native 🗌 Asian 🗌 Black	or African Americar	ı		
	Native Hawaiian/Othe	r Pacific Islander	ŕ			
*Blood Gr	roup: 🗌 A- 🗌 A+ 🗌 B-	- 🗌 B+ 🗌 AB- 🗌 AB+ 🗌 O-	O+ Blood	type not done		
	Transitional ABO / F	Rh + 🔄 Transitional ABO / Rh -	Transitional AB	O / Transitional Rh		
Group A	VTransitional Rh 🗌 Group	B/Transitional Rh 🛛 Group O/Transition	al Rh 🛛 Group A	B/Transitional Rh		
Patient M	ledical History					
List the	patient's admitting diagnos	is. (Use ICD-10 Diagnostic codes/descr	iptions)			
Code	: I	Description:				
Code	: I	Description:				
Code	: I	Description:				
List the	patient's underlying indicat	ion for transfusion. (Use ICD-10 Diagnos	stic codes/descriptio	ns)		
Code	: I	Description:				
Code		Description:				
		Description:				
		ns at the time of the transfusion related		UNKNOWN		
Code	: I	Description:				
Code	: I	Description:				
Code	: I	Description:				
of any indivi stated, and	dual or institution is collected wi will not otherwise be disclosed o	y provided information obtained in this surveilla th a guarantee that it will be held in strict confid r released without the consent of the individual Health Service Act (42 USC 242b, 242k, and 24	lence, will be used only I, or the institution in ac	for the purposes		
reviewing in collection of	structions, searching existing da information. An agency may no	nformation is estimated to average 20 minutes ta sources, gathering and maintaining the data of conduct or sponsor, and a person is not requ trol number. Send comments regarding this bu	a needed, and completin ired to respond to a col	ng and reviewing the lection of information		

Atlanta, GA 30333 ATTN: PRA (0920-0666).

collection of information, including suggestions for reducing this burden to CDC, Reports Clearance Officer, 1600 Clifton Rd., MS D-74,

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List the patient's relevant r performed during the curre						
codes/descriptions)						
Code:	_ Description:					
Code:	_ Description:		· · · · · · · · · · · · · · · · · · ·			
Code:	_ Description:					
Additional Information						
Transfusion History						
Has the patient received a	•		NKNOWN			
Blood Product:	WB RBC Platelet		Granulocyte			
Date of Transfusion:						
-	e reaction transfusion-related?	YES NO				
	n about the transfusion adverse re		_			
	rse reaction:					
		— —				
	ify					
Reaction Details			_			
	// *Time reaction occ	urred::	unknown			
*Facility location where patient was transfused:						
Is this reaction associated wit		No If Yes, Incident #:				
Is this reaction associated wit						
Is this reaction associated wit Investigation Results * Other						
Is this reaction associated wit Investigation Results * Other		No If Yes, Incident #:				
Is this reaction associated wit Investigation Results * Other	h an incident?	No If Yes, Incident #:				
Is this reaction associated wit Investigation Results * Other Specify:	h an incident?	No If Yes, Incident #:				
Is this reaction associated wit Investigation Results * Other Specify: List tests relevant to reac	h an incident? Yes tion investigation:	No If Yes, Incident #:				
Is this reaction associated wit Investigation Results * Other Specify: List tests relevant to reac Test name:	h an incident? Yes Yes tion investigation: Testing date: Testing date:] No If Yes, Incident #: Test result:				
Is this reaction associated wit Investigation Results * Other Specify: List tests relevant to reac Test name: Test name:	h an incident? Yes Yes tion investigation: Testing date: Testing date:] No If Yes, Incident #: Test result:				
Is this reaction associated wit Investigation Results * Other Specify: List tests relevant to reac Test name: Test name: Other signs and symptoms:	h an incident? Yes tion investigation: Testing date: (check all that apply)	No If Yes, Incident #:				
Is this reaction associated wit Investigation Results * Other Specify: List tests relevant to reac Test name: Test name: Other signs and symptoms: Generalized: Cardiovascular:	h an incident? Yes tion investigation: Testing date: Testing date: (check all that apply) Chills/rigors Fever	No If Yes, Incident #: Test result: Test result: Test result: Nausea/v Shock				
Is this reaction associated wit Investigation Results * Other Specify: List tests relevant to reac Test name: Test name: Other signs and symptoms: Generalized:	h an incident? Yes tion investigation: Testing date: (check all that apply) Chills/rigors Fever Blood pressure decrease Edema Flushir	No If Yes, Incident #: Test result: Test result: Test result: Nausea/v Shock	 			
Is this reaction associated wit Investigation Results * Other Specify: List tests relevant to reac Test name: Test name: Other signs and symptoms: Generalized: Cardiovascular: Cutaneous:	h an incident? Yes tion investigation: Testing date: (check all that apply) Chills/rigors Fever Blood pressure decrease Edema Flushir	No If Yes, Incident #: Test result: Test result: Nausea/v Nausea/v Shock ng Jaundice s (itching) Urticaria (romiting			
Is this reaction associated wit Investigation Results * Other Specify: List tests relevant to reac Test name: Test name: Other signs and symptoms: Generalized: Cardiovascular:	h an incident? Yes Yes tion investigation: Testing date: Testing date: (check all that apply) Chills/rigors Fever Blood pressure decrease Edema Flushir Other rash Pruritu	No If Yes, Incident #: Test result: Test result: Nausea/v Nausea/v Shock ng Jaundice s (itching) Urticaria (romiting			
Is this reaction associated wit Investigation Results * Other Specify: List tests relevant to reac Test name: Test name: Other signs and symptoms: Generalized: Cardiovascular: Cutaneous:	h an incident? Yes Yes tion investigation: Testing date: Testing date: (check all that apply) Chills/rigors Fever Blood pressure decrease Edema Flushir Other rash Pruritu: Disseminated intravascular c	No If Yes, Incident #: Test result: Test result: Nausea/v Shock ng Jaundice s (itching) Urticaria (oagulation Hemoglo	romiting			
Is this reaction associated wit Investigation Results * Other Specify: List tests relevant to reac Test name: Test name: Other signs and symptoms: Generalized: Cardiovascular: Cutaneous: Hemolysis/Hemorrhage:	h an incident? Yes Yes tion investigation: Testing date: Testing date: (check all that apply) Chills/rigors Fever Blood pressure decrease Blood pressure decrease Edema Flushir Other rash Pruritus Disseminated intravascular c Positive antibody screen Abdominal pain Back p	No If Yes, Incident #: Test result: Test result: Nausea/v Shock ng Jaundice s (itching) Urticaria (oagulation Hemoglo	 /omiting (hives) binemia			
Is this reaction associated wit Investigation Results * Other Specify: List tests relevant to reac Test name: Test name: Other signs and symptoms: Generalized: Cardiovascular: Cutaneous: Hemolysis/Hemorrhage: Pain: Renal:	h an incident? Yes Yes Yes Yes Yes Yes Yes Yes	No If Yes, Incident #: Test result: Test result: Nausea/v Nausea/v Shock Nausea/v Shock Nausea/v Shock Nausea/v Shock Nausea/v Shock Nausea/v Shock Nausea/v Shock Nausea/v Shock Nausea/v Shock Nausea/v Shock Nausea/v Shock Nausea/v Shock Nausea/v Shock Nausea/v Shock Sh	 /omiting (hives) binemia			
Is this reaction associated wit Investigation Results * Other Specify: List tests relevant to reac Test name: Test name: Other signs and symptoms: Generalized: Cardiovascular: Cutaneous: Hemolysis/Hemorrhage: Pain:	h an incident? Yes Yes Yes Yes Yes Yes Yes Yes	No If Yes, Incident #: Test result: Test result: Nausea/v Nausea/v Shock Nausea/v Shock Nausea/v Shock Nausea/v Shock Nausea/v Shock Nausea/v Shock Nausea/v Shock Nausea/v Shock Nausea/v Shock Nausea/v Shock Nausea/v Shock Nausea/v Shock Nausea/v Shock Nausea/v Shock Sh	romiting (hives) binemia] Infusion site pain			

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*Severity					
Did the patient receive or experience any of the following?					
□ No treatment required □ Symptomatic	treatment only				
Hospitalization, inlcuding prolonged hospitalization	Life-threatenin	g reaction			
	omaly or birth defect(
Other medically important conditions	Unknown or no	ot stated			
*Imputability					
Which best describes the relationship between the transfusion and t	the reaction?				
Conclusive evidence exists that the adverse reaction can be a		ision.			
Evidence is clearly in favor of attributing the adverse reaction					
Evidence is indeterminate for attributing the adverse reaction t					
Evidence is clearly in favor of a cause other than the transfusion					
There is conclusive evidence beyond reasonable doubt of a ca					
☐ The relationship between the adverse reaction and the transfu	ision is unknown of no	DI STATEO.			
Did the transfusion occur at your facility?					
Module-generated Designations NOTE: Designations for case definition, severity, and imputability will be automatically assigned in the NHSN application based on responses in the corresponding investigation results section 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 <u>imputability</u> designation? ^ Please indicate your designation	☐ YES				
Patient Treatment					
Did the patient receive treatment for the transfusion reaction? If yes, select treatment(s): Medication (Select the type of medication)	□ YES □ NO				
 Antipyretics Antihistamines Inotropes/Vasopre Intravenous Immunoglobulin Intravenous steroids Antithymocyte globulin Cyclosporin Otherwork 	Corticosteroids	ator Diuretics			
Volume resuscitation (Intravenous colloids or crystalloids)					
 Respiratory support (Select the type of support) Mechanical ventilation Noninvasive ventilation 	□ Oxygen				
 Renal replacement therapy (Select the type of therapy) Hemodialysis Peritoneal Continuous Veno-Year 	Venous Hemofiltration				
Phlebotomy Other Specify:					

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Outcome									
*Outcome: Death Dajor or long-term sequelae Dinor or no sequelae Not determined									
	Date of Death://								
	If recipient died, relationship of transfusion to death:								
	Definite	e 🗌 Possibl	le 🗌 Doubtful	Ruled Ou	it _	Not de	etermine	эd	
	of death:								
vvas an	autopsy performed?	☐ Yes	□ No						
Component									
*Was a partic	*Was a particular unit implicated in (i.e., responsible for) the adverse] N/A		
Transfusion Start and End Date/Time	*Component code (check system used)	Amount transfused at reaction onset	^A Unit number (Required for Infection and TRALI)	*Unit expiration Date/Time	*Blood group of unit			Implic ated Unit?	
^IMPLICATED UNIT									
/ /	□ ISBT-128								
:	☐ Codabar	Entire unit		1 1	🗆 A-	□ A+	🗆 в-		
		Partial unit mL			Пв+	П АВ-	П АВ+	Y	
· · · · · · · · · · · · · · · · · · ·				_		П 0+			
				··					
:	 □ Codabar	Entire unit		1 1	🗆 A-	🗆 A+	🗆 В-		
		□ Partial unit mL			Пв+	П АВ-	П АВ+	N	
· · · ·				_	0-	□ 0+			
Custom Field									
Label			Label						
		II				/	/		
<u></u>									
Comments									

