

Hemovigilance Module Adverse Reaction Transfusion Associated Circulatory Overload

*Required for saving	,	
*Facility ID#: NHSN Ad	lverse Reaction #:	
Patient Information		
*Patient ID:	*Gender: M F Other	
Social Security #:	Secondary ID:	
Last Name:	First Name:	_ Middle Name:
Ethnicity 🗌 Hispanic or Latino	Not Hispanic or Not Latino	
Race 🗌 American Indian/Alask	a Native 🗌 Asian 🗌 Black	k or African American
🗌 Native Hawaiian/Othe	r Pacific Islander	e
*Blood Group: 🗌 A- 🗌 A+ 🗌 B	3- 🗌 B+ 🗌 AB- 🗌 AB+ 🗌 O-	O+ Blood type not done
🗌 Transitional ABO / I	Rh + 🛛 Transitional ABO / Rh -	🗌 Transitional ABO / Transitional Rh
Group A/Transitional Rh	B/Transitional Rh Group O/Transition	nal Rh 🛛 Group AB/Transitional Rh
Patient Medical History		
List the patient's admitting diagnos	sis. (Use ICD-10 Diagnostic codes/descri	iptions)
Code:	Description:	
Code:	Description:	
Code:	Description:	
List the patient's underlying indica	tion for transfusion. (Use ICD-10 Diagnos	stic codes/descriptions)
Code:	Description:	
Code:	Description:	
Code:	Description:	
List the patient's comorbid condition reaction. (Use ICD-10 Diagnostic of the ICD-10 Diagnostic of the term of	ons at the time of the transfusion related codes/descriptions)	to the adverse UNKNOWN
Code:	Description:	
Code:	Description:	
Code:	Description:	
any individual or institution is collected with and will not otherwise be disclosed or relea 306 and 308(d) of the Public Health Service Public reporting burden of this collection of instructions, searching existing data source information. An agency may not conduct or displays a currently valid OMB control number	y provided information obtained in this surveilla a guarantee that it will be held in strict confiden sed without the consent of the individual, or the Act (42 USC 242b, 242k, and 242m(d)). information is estimated to average 20 minutes s, gathering and maintaining the data needed, a sponsor, and a person is not required to respo per. Send comments regarding this burden esti- ing this burden to CDC, Reports Clearance Offi	ce, will be used only for the purposes stated, institution in accordance with Sections 304, per response, including the time for reviewing and completing and reviewing the collection of nd to a collection of information unless it mate or any other aspect of this collection of



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	nedical procedure including past procedures and procedures to be UNKNOWN ent 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 advers	e reaction transfusion-related?		
If yes, provide informatio	n about the transfusion adverse reaction.		
Type of transfusion adve	rse reaction: 🗌 Allergic 🗌 AHTR 🗌 DHTR 🗌 DSTR 🗌 FNHTR		
	□ PTP □ TACO □ TAD □ TA-GVHD □ TRALI □ UNKNOWN		
OTHER Spec	fy		
Reaction Details			
*Date reaction occurred:	// *Time reaction occurred::_ D Time unknown		
*Facility location where pat	ent was transfused:		
Is this reaction associated wit	h an incident? Yes No If Yes, Incident #:		
Investigation Results			
* Transfusion associat	ed circulatory overload (TACO)		
*Case Definition			
Check all that occurred v	within 6 hours of cessation of transfusion (new onset or exacerbation):		
Acute respiratory di	stress (dyspnea, orthopnea, cough)		
Elevated brain natri	uretic peptide (BNP)		
Elevated central ver	nous pressure (CVP)		
Evidence of left hea	rt failure		
Evidence of positive fluid balance			
Radiographic evide	nce of pulmonary edema		
Other signs and symptoms	: (check all that apply)		
Generalized:	Chills/rigors Fever Nausea/vomiting		
Cardiovascular:	Blood pressure decrease Shock		
Cutaneous:	Edema Flushing Jaundice		
	Other rash Pruritus (itching) Urticaria (hives)		
Hemolysis/Hemorrhage:	 Disseminated intravascular coagulation Hemoglobinemia Positive antibody screen 		
Pain:	Abdominal pain Back pain Flank pain Infusion site pain		
Renal:	Hematuria Hemoglobinuria Oliguria		
Respiratory:	Bilateral infiltrates on chest x-ray Bronchospasm Cough		



	🗌 Hypoxemia	Shortness of breat	h		
Other: (specify)					
*Severity					
Did the patient receive or	experience any of the	following?			
🗌 No treatment requ	ired	Symptomatic trea	Itment only		
Hospitalization, inl	cuding prolonged hosp	oitalization	Life-threatening	reaction	
Disability and/or in	capacitation	Congenital anoma	aly or birth defect(s) of the fetus	
Other medically in	portant conditions	Death	Unknown or no	t stated	
*Imputability					
☐ Transfusion is a like ☐ The patient has a hi ☐ Evidence is clearly i	ns for circulatory overlo ly contributor to circula story of a pre-existing o n favor of a cause othe	oad are possible.	kely explains circula but transfusion can	not be excluded.	
The relationship bet	ween the adverse read	ction and the transfusior	n is unknown or not	stated.	
Did the transfusion occur	at your facility?	YES 🗌 NO			
 Does the patient have a history of cardiac insufficiency? Yes, the patient has a history of cardiac insufficiency that could explain the circulatory overload, but transfusion is just as likely to have caused the circulatory overload. Yes, the patient has a history of pre-existing cardiac insufficiency that most likely explains circulatory overload. No, the patient does not have a history of cardiac insufficiency. Did the patient received other fluids in addition to the transfusion? 					
Madula non-metod Desin					
Module-generated Desig NOTE: Designations for case		d imputability will be au	tomatically assigned	d in the NHSN	
application based on response					
* Do you agree with the ^Please indicate your de		nation?	☐ YES		
* Do you agree with the ^Please indicate your de		?	☐ YES	□ NO	
* Do you agree with the ^Please indicate your de		ion?	☐ YES	□ NO	
Patient Treatment					
Antipyretics	the type of medication)] Inotropes/Vasopresso			
Antithymocyte	•	ravenous steroids sporin Other	_ Corticosteroids	Antibiotics	

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🗌 Volu	me resuscitation (Intra	avenous colloid	s or crystalloids)				
Res	piratory support <i>(Sele</i>] Mechanical ventilati	-	<i>pport)</i> vasive ventilation	🗌 Oxygen			
Ren C	al replacement therap] Hemodialysis 🗌 F		be of therapy) Continuous Ven	o-Venous Hemo	ofiltration		
🗌 Othe	botomy er Specify:						
Outcome							
Cause o	Death:/ recipient died, relation Definite Probable of death:	e 🗌 Possibl	ion to death: e	Minor or no sec		ot detern etermine	
	autopsy performed?	Yes					
Component *Was a partic reaction?	Details 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	^A Unit number (Required for Infection and TRALI)	*Unit expiration Date/Time	*Blood grou of unit	р	Implic ated Unit?
^IMPLICATED	UNIT						
! : !!	☐ ISBT-128 ☐ Codabar 	☐ Entire unit ☐ Partial unit mL			□ A- □ A+ □B+ □ AB- □ O- □ O+	□ B- □ AB+ □ N/A	Y
// : // :	☐ ISBT-128 ☐ Codabar ——————————	☐ Entire unit ☐ Partial unit mL		// :	□ A- □ A+ □B+ □ AB- □ O- □ O+	□ B- □ AB+ □ N/A	N
Custom Fields							
Label Comments		<u> </u>	Label		/	<u> </u>	

