Hemovigilance Module Adverse Reaction Transfusion Associated Circulatory Overload

*Required for saving		-		
*Facility ID#: NHS	SN Adverse Reaction #:			
Patient Information				
*Patient ID:	*Gender: M F	Other *Date of Birth: //		
Sex at Birth: \Box M \Box F \Box Unk	nown			
	Gender Identity (Specify)			
Social Security #:				
Last Name:	First Name: Middle Name:			
Ethnicity Hispanic or Latin	Not Hispanic or Not Lating	0		
Race American Indian	/Alaska Native 📃 Asian	Black or African American		
Native Hawaiian	/Other Pacific Islander	White		
*Blood Group: A- A+	B- B+ AB- AB+	O- O+ Blood type not done		
Group A/Transitional		Transitional Rh Group AB/Transitional Rh		
Rh Patient Medical History		·		
,	agnosis. (Use ICD-10 Diagnostic code	es/descriptions)		
Code:				
Code:				
Code:				
	ndication for transfusion. (Use ICD-10			
Code:	Description:			
Code:				
Code:	Description:			
List the patient's comorbid co reaction. (Use ICD-10 Diagno	nditions at the time of the transfusion ostic codes/descriptions)	related to the adverse UNKNOWN		
Code:	Description:			
Code:	Description:			
Code:	Description:			
is collected with a guarantee that it will be held	l in strict confidence, will be used only for the purpose	stem that would permit identification of any individual or institution es stated, and will not otherwise be disclosed or released without the bublic Health Service Act (42 USC 242b, 242k, and 242m(d)).		
existing data sources, gathering, and maintaini and a person is not required to respond to a col	ng the data needed, and completing and reviewing the lection of information unless it displays a currently va of information, including suggestions for reducing this	onse, including the time for reviewing instructions, searching collection of information. An agency may not conduct or sponsor, alid OMB control number. Send comments regarding this burden s burden to CDC, Reports Clearance Officer, 1600 Clifton Rd., MS		

	edical procedure including past procedures and procedures to be UNKNOWN t hospital or outpatient stay. (Use ICD-10 Procedure NONE
Code:	Description:
Code:	Description:
Code:	Description:
Additional Information	
Transfusion History	
Has the patient received a p	revious transfusion? YES NO UNKNOWN
Blood Product:	WB RBC Platelet Plasma Cryoprecipitate Granulocyte
Date of Transfusion:	
Was the patient's adverse	reaction transfusion-related?
If yes, provide information	about the transfusion adverse reaction.
Type of transfusion advers	se reaction:
HTR TTI	PTP TACO TAD TA-GVHD TRALI UNKNOWN
OTHER Specify	/
Reaction Details	
*Date reaction occurred:/	/ *Time reaction occurred:: Time unknown
*Facility location where patie	
Is this reaction associated with	an incident? Yes No If Yes, Incident #:
Investigation Results	
* Transfusion associate	ed circulatory overload (TACO)
*Case Definition	
	ithin 12 hours of cessation of transfusion (new onset or exacerbation):
	tress (dyspnea, orthopnea, cough)
Elevated brain natriu	retic peptide (BNP)
Elevated central ven	
Evidence of left hear	
Evidence of positive	
	ce of pulmonary edema
Other signs and symptoms:	
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
	Positive antibody screen
Pain:	Abdominal pain Back pain Flank pain pain
Renal:	Hematuria Hemoglobinuria Oliguria
	Bilateral infiltrates on chest x-ray Bronchospasm Cough
Respiratory:	Hypoxemia Shortness of breath

Other: (specify)							
*Severity							
Did the patient receive or experience any of the following?							
No treatment required Symptomatic treatment only							
Hospitalization, inlcuding prolonged hospita	lization	Life-threatening	reaction				
Disability and/or incapacitation	Disability and/or incapacitation Congenital anomaly or birth defect(s) of the fetus						
Other medically important conditions	Death	Unknown or not	stated				
*Imputability							
Which best describes the relationship between the	transfusion and the r	eaction?					
No other explanations for circulatory overload							
Transfusion is a likely contributor to circulato	y overload						
The patient has a history of a pre-existing co	ndition(s) that most lil	kely explains circulat	tory overload.				
Evidence is clearly in favor of a cause other t							
There is conclusive evidence beyond reason							
The relationship between the adverse reaction	n and the transfusior	1 is unknown or not s	stated.				
Did the transfusion occur at your facility?	ES NO						
Does the patient have a history of cardiac insufficie	ncy?						
Yes, the patient has a history of cardiac insu	-		/ overload, but				
transfusion is just as likely to have caused the	•						
Yes, the patient has a history of pre-existing overload.	cardiac insufficiency	that most likely expl	ains circulatory				
No, the patient does not have a history of cardiac insufficiency. Did the patient received other fluids in addition to the transfusion?							
Module-generated Designations							
NOTE: Designations for case definition, severity, and ir			in the NHSN				
application based on responses in the corresponding in	vestigation results se	ection above.					
*Do you agree with the <u>case definition</u> designat	tion?	YES	NO				
^Please indicate your designation							
*Do you agree with the <u>severity</u> designation?		YES	NO				
^Please indicate your designation							
*Do you agree with the <i>imputability</i> designation	ו?	YES	NO				
^Please indicate your designation							
Detiont Treatment							
Patient Treatment							
Did the patient receive treatment for the transfusion	reaction?	YES NO					
If yes, select treatment(s): Medication (Select the type of medication)							
Antipyretics Antihistamines Inotropes/Vasopressors Bronchodilator Diuretics							
Immunoglobulin Intravenous steroids Corticosteroids Antibiotics							
Antithymocyte globulin Cyclosporin Other							

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-Venous Hemofiltration										
Phlebotomy Other Specify:										
Outcome										
*Outcome:	Death M	lajor or long-terr	n sequelae] Minor or no sec	quelae 🗌 Not deter	mined				
Date of	Death:/_	/								
^lf I	ecipient died, relation	·								
	Definite Probabl	e Possib	le Doubtful	Ruled Out	Not determin	ed				
	autopsy performed?	Yes	No							
Component	Details									
	cular unit implicated	d in (i.e., respo		dverse	Yes No	N/A				
Transfusion Start and End	*Component code	Amount transfused at	AUnit number (Required for Infection and	*Unit expiration	*Blood group ated					
Date/Time	(check system used)	reaction onset	TRALI)	Date/Time	of unit	Unit?				
^IMPLICATED										
//	ISBT-128				A- A+ B-					
:	Codabar	Entire unit		/	A- A+ B-	Y				
/		mL			B+ AB- AB+					
:				:	0- 0+ N/A					
//	ISBT-128									
:	Codabar	Entire unit			A- A+ B-	N				
//	·	mL			B+ AB- AB+					
:				::	0- 0+ N/A					
Custom Field	ds									
Label			Label							
		//			//					
Comments										