

*Required for saving

Form Approved OMB No. 0920-0666 Exp. Date: 01/31/25 www.cdc.gov/nhsn

Hemovigilance Module Adverse Reaction Transfusion Associated Dyspnea

*Facility ID#: NHSN Adve	rse Reaction #:
Patient Information	
	Gender: M F Other *Date of Birth:/
Social Security #:	Secondary ID: Medicare #:
Last Name:	First Name: Middle Name:
Ethnicity Hispanic or Latino	☐ Not Hispanic or Not Latino
Race	Native Asian Black or African American
☐ Native Hawaiian/Other P	 -
*Blood Group : ☐ A- ☐ A+ ☐ B-	□B+ □ AB- □ AB+ □ O- □ O+ □ Blood type not done
	+
Patient Medical History	
List the patient's admitting diagnosis.	(Use ICD-10 Diagnostic codes/descriptions)
Code: De	scription:
Code: De	scription:
	scription:
List the patient's underlying indication	n for transfusion. (Use ICD-10 Diagnostic codes/descriptions)
Code: De	scription:
Code: De	scription:
	scription:
List the patient's comorbid conditions reaction. (Use ICD-10 Diagnostic cod	at the time of the transfusion related to the adverse UNKNOWN des/descriptions) □ NONE
Code: De	scription:
Code: De	scription:
Code: De	scription:
of any individual or institution is collected with a stated, and will not otherwise be disclosed or respections 304, 306 and 308(d) of the Public Head Public reporting burden of this collection of inforeviewing instructions, searching existing data collection of information. An agency may not counless it displays a currently valid OMB control	rovided information obtained in this surveillance system that would permit identification a guarantee that it will be held in strict confidence, will be used only for the purposes eleased without the consent of the individual, or the institution in accordance with alth Service Act (42 USC 242b, 242k, and 242m(d)). Transition is estimated to average 20 minutes per response, including the time for sources, gathering and maintaining the data needed, and completing and reviewing the onduct or sponsor, and a person is not required to respond to a collection of information number. Send comments regarding this burden estimate or any other aspect of this for reducing this burden to CDC, Reports Clearance Officer, 1600 Clifton Rd., MS D-74,



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Transfusion Associated Dyspnea

· · · · · · · · · · · · · · · · · · ·	nedical procedure including put hospital or outpatient stay	-	-	UNKNOWN NONE
Code:	Description:			
Code:				
Code:				
Additional Information				
Transfusion History				
Blood Product: Date of Transfusion: Was the patient's advers If yes, provide informatio Type of transfusion adve	previous transfusion? WB RBC Platele// Ul e reaction transfusion-related n about the transfusion adve rse reaction: Allergic PTP TACO T	ot	Cryoprecipitate 'ES	FNHTR □ UNKNOWN
Reaction Details	·,			
*Date reaction occurred:	/ / *Time reaction	n occurred: :	Time	e unknown
*Facility location where pat				
Is this reaction associated wit	n an incident?	s 🗌 No 🛮 If Y	es, Incident #:	
Investigation Res	ults			
* Transfusion associated dyspnea (TAD)				
	stress occurring within 24 ho CO, and TRALI definitions a		ransfusion.	
Other signs and symptoms:	(check all that apply)			
Generalized:	☐ Chills/rigors ☐ F	ever 🗌 Na	ausea/vomiting	
Cardiovascular:	☐ Blood pressure decreas	se 🗌 Sh	ock	
Cutaneous:		lushing Pruritus (itching)	☐ Jaundice ☐ Urticaria (hiv	/es)
Hemolysis/Hemorrhage:	☐ Disseminated intravaso	•	☐ Hemoglobin	emia
Pain:	☐ Abdominal pain ☐ E	Back pain 🔲 Fla	ank pain [☐ Infusion site pain
Renal:	☐ Hematuria ☐ F	lemoglobinuria	☐ Oliguria	
Respiratory:	☐ Bilateral infiltrates on ch ☐ Hypoxemia ☐ S	nest x-ray	onchospasm [Cough



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Other: (specify)				
*Severity				
Did the patient receive or experience any of the following?				
☐ No treatment required ☐ Symptomatic treatment only				
☐ Hospitalization, inlcuding prolonged hospitalization [Life-threatening	reaction		
☐ Disability and/or incapacitation ☐ Congenital anomal	ly or birth defect(s) of the fetus		
☐ Other medically important conditions ☐ Death [Unknown or no	t stated		
*Imputability				
Which best describes the relationship between the transfusion and the re	eaction?			
☐ Patient has no other conditions that could explain symptoms.				
☐ There are other potential causes that could explain symptoms, but	transfusion is the	most likely cause.		
Other present causes are most likely, but transfusion cannot be ru	led out.			
$\hfill \square$ Evidence is clearly in favor of a cause other than the transfusion, b	ut transfusion can	not be excluded.		
☐ There is conclusive evidence beyond reasonable doubt of a cause	other than the trai	nsfusion.		
☐ The relationship between the adverse reaction and the transfusion	is unknown or not	stated.		
Did the transfusion occur at your facility? ☐ YES ☐ NO				
Module-generated Designations				
NOTE: Designations for case definition, severity, and imputability will be auto application based on responses in the corresponding investigation results see		d in the NHSN		
*Do you agree with the <u>case definition</u> designation? ^Please indicate your designation	☐ YES	□ NO		
*Do you agree with the <u>severity</u> designation? ^Please indicate your designation	☐ YES	□ NO		
*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? If yes, select treatment(s): Medication (Select the type of medication)	∕ES □ NO	UNKNOWN		
☐ Antipyretics ☐ Antihistamines ☐ Inotropes/Vasopressors	s	tor Diuretics Antibiotics		
☐ 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-Veno ☐ Phlebotomy 	us Hemofiltration	,		





Other Specify:									
Outcome									
*Outcome: Death Major or long-term sequelae Minor or no sequelae Not determined Date of Death:// ^If recipient died, relationship of transfusion to death: Definite Probable Possible Doubtful Ruled Out Not determined Cause of death: Was an autopsy performed? Yes No									
Component									
*Was a partic	cular unit implicate	d in (i.e., resp	onsib	le for) the a	dverse	☐ Ye	s 🗌	No [□ N/A
Transfusion Start and End Date/Time	*Component code (check system used)	Amount transfused at reaction onset	iiiiootioii aiia		*Unit expiration Date/Time	*Blood group of unit			Implic ated Unit?
^IMPLICATED	UNIT								
: : :	☐ ISBT-128 ☐ Codabar	☐ Entire unit ☐ Partial unit mL	 			□ A- □B+ □ O-		□ B- □ AB+ □ N/A	Y
	☐ ISBT-128 ☐ Codabar — — — — —	☐ Entire unit ☐ Partial unit mL		·———		□ A- □B+ □ O-	□ A+ □ AB-	□ B- □ AB+ □ N/A	N
Custom Fields								-	
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Comments									