

Hemovigilance Module Adverse Reaction Transfusion Related Acute Lung Injury

*Required for saving]					
*Facility ID#:	NHSN Adverse Re	action #:				
Patient Information						
*Patient ID:			*Date of	f Birth://		
*Sex: M F						
Social Security #:	Secon	dary ID:	Medica	are #:		
Last Name:	First	Name:	Middle	Name:		
Ethnicity (Specify):	Hispanic or Latino	Not Hispanic or Latino	Unknown	Declined to respond		
Race (Select all that apply):	American Indian or Alaska Native Native Hawaiian or Pacific Islander	Asian White	Black or African American Unknown	Middle Eastern or North African Declined to respond		
Preferred Language (Specify from the list provided): Interpreter Needed:YesNoUnknow						
*Blood Group: A- A+ B- B+ AB- AB+ O- O+ Blood type not done Transitional ABO / Rh + Transitional ABO / Rh - Rh Group A/Transitional Rh Group A/Transitional Rh Group A/Transitional Rh						
Patient Medical His	story					
List the patient's ad	mitting diagnosis. (Use I	CD-10 Diagnostic o	odes/descriptions)			
Code:	Description	Description:				
Code: Description:						
	Code: Description:					
List the patient's underlying indication for transfusion. (Use ICD-10 Diagnostic codes/descriptions)						
Code:	Description	on:	_			
Code:						
Code:	Description:					
List the patient's comorbid conditions at the time of the transfusion related to the adverse reaction. (Use ICD-10 Diagnostic codes/descriptions)						
Code:	Description	on:				
Code:						
	Code: Description:					

Assurance of Confidentiality: The voluntarily provided information obtained in this surveillance system that would permit identification of any individual or institution is collected with a guarantee that it will be held in strict confidence, will be used only for the purposes stated, and will not otherwise be disclosed or released without the consent of the individual, or the institution in accordance with Sections 304, 306 and 308(d) of the Public Health Service Act (42 USC 242b, 242k, and 242m(d)). CDC 57.317 Rev. 3, v9.2

Public reporting burden of this collection of information is estimated to average 20 minutes per response, including the time for reviewing instructions, searching existing data sources, gathering, and maintaining the data needed, and completing and reviewing the collection of information. An agency may not conduct or sponsor, and a person is not required to respond to a collection of information unless it displays a currently valid OMB control number. Send comments regarding this burden estimate or any other aspect of this collection of information, including suggestions for reducing this burden to CDC, Reports Clearance Officer, 1600 Clifton Rd., MS H21-8, Atlanta, GA 30333, ATTN: PRA (0920-0666).







pe	t the patient's relevant r formed during the curre des/descriptions)						UNKNOWN NONE
C	Code:	Description:					
C	ode:						
	Code:						
Ad	ditional Information						
	sfusion History						
На	Has the patient received a previous transfusion? YES NO UNKNOWN						
E	Blood Product: WB RBC Platelet Plasma Cryoprecipitate Granulocyte					Granulocyte	
	ate of Transfusion:			UNKN	OWN		
٧	Vas the patient's advers	e reaction tr	ansfusio	n-related?	YE	S NO	
If	yes, provide informatio	n about the	transfusi	on adverse r	eaction.		
Т	ype of transfusion adve	rse reaction	: [Allergic	AHTR D	HTR DSTR	FNHTR
	HTR TTI	PTP [TACO	TAD	TA-GVHD	TRALI	UNKNOWN
	OTHER Speci	ify					<u></u>
Reac	tion Details						
*Date	reaction occurred:	<u> </u>	*Time	reaction oc	curred::	Time unl	known
*Facil	ity location where pat	ient was tra	ınsfused	l:			
Is this	reaction associated wit	h an inciden	ıt?	Yes	No If Yes	s, Incident #:	
Inves	stigation Results						
* Tr	ansfusion related ac	ute lung inj	ury (TR	ALI)			
						Test result positive	
					Cognate or	No cognate or	Not tested for
			Not Done	Negative	cross reacting antigen present	cross reacting antigen present	cognate antigen
	Donor or unit HLA sp	ecificity	Done	rvegative	anagen present	drugeri present	antigen
	Donor or unit HNA sp	•					
	Recipient HLA spec	cificity					
	Recipient HNA spe						
*Case Definition (Check all that apply)							
NO evidence of acute lung injury (ALI) prior to transfusion.							
ALI onset during or within 6 hours of cessation of transfusion							
Hypoxemia – defined as PaO2/FiO2 less than or equal to 300 mm Hg							
Hypoxemia – defined as Oxygen saturation less than 90% on room air							
Hypoxemia – defined as Other clinical evidence							
Radiographic evidence of bilateral infiltrates No evidence of left atrial bypartopsian (i.e., circulatory everlead)							
No evidence of left atrial hypertension (i.e., circulatory overload) Other signs and symptoms: (check all that apply)							
	Generalized: Chills/rigors Fever Nausea/vomiting						
	Cardiovascular: Blood pressure decrease Shock						
С	ardiovascular:						k



			Hives	
Hemolysis/Hemorrhage:	DIC Hemoglobine	emia Positive antibod	y screen	
Pain:	Abdominal pain	Back pain Flank	pain Infusion site pain	
Renal:	Hematuria	Hemoglobinuria	Oliguria	
Respiratory:	Bronchospasm C	Cough Shortness of bre	eath Other: (specify)	
*Severity				
Did the patient receive or	r experience any of the fol	lowing?		
No treatment requ	uired [Symptomatic treatment	only	
Hospitalization, in	lcuding prolonged hospita	lization Li	fe-threatening reaction	
Disability and/or in	ncapacitation		birth defect(s) of the fetus	
Other medically in	nportant conditions	Death U	nknown or not stated	
*Imputability				
• •	relationship between the	transfusion and the reaction	nn?	
	ative risk factors for ALI p		,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	
	of other causes for acute lu			
		• , ,	ansfusion cannot be excluded.	
		able doubt of a cause othe		
	•	n and the transfusion is ur		
Did the transfusion occur		ES NO	introvir or not stated.	
Module-generated Design		anutahilituwill ba autamati	solly assigned in the NUICNI	
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	case definition designat	ion?	YES NO	
^Please indicate your de				
*Do you agree with the severity designation? YES NO				
^Please indicate your designation				
*Do you agree with the	<u>imputability</u> designation	1?	YES NO	
^Please indicate your de	signation			
Patient Treatment				
Did the patient receive trea	tment for the transfusion r	reaction? YES	NO UNKNOWN	
If yes, select treatment(s):				
Medication (Select the type of medication)				
Antipyretics Antihistamines Inotropes/Vasopressors Bronchodilator Diuretics				
Intravenous Continue				
Immunoglobulin				
Antithymocyte globulin Cyclosporin Other				
Volume resuscitation (Intravenous colloids or crystalloids)				
Respiratory suppor	t (Select the type of suppo	ort)		
Mechanical ventilation Noninvasive ventilation Oxygen				



Renal replacement therapy (Select the type of therapy) Hemodialysis Peritoneal Continuous Veno-Venous Hemofiltration Phlebotomy Other Specify:						
Outcome						
*Outcome: Date of		ajor or long-terr	n sequelae	Minor or no se	quelae Not deter	mined
^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	Details					
*Was a particular unit implicated in (i.e., responsible for) the adverse reaction?						
Transfusion Start and End Date/Time	*Component code (check system used)	Amount transfused at reaction onset	^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 Partial unitmL			A- A+ B- B+ AB- AB+ O- O+ N/A	- Y
:	☐ ISBT-128 ☐ Codabar ☐ — — — — — —	Entire unit Partial unitmL			A- A+ B- B+ AB- AB+ O- O+ N/A	N
Custom Fields						
Label						
Comments						



The state of the s	Form Approved
NHSN	OMB No. 0920-0666
NATIONAL HEALTHCARE	Exp. Date: 12/31/2027
SAFETY NETWORK	www.cdc.gov/nhsn