

Hemovigilance Module Adverse Reaction Transfusion Related Acute Lung Injury

*Required for saving

*Facility ID#:	NHSN Adverse Reaction	on #:
Patient Informa	ation	
*Patient ID:		*Date of Birth://
*Sex at Birth: □M	□F □Unknown	*Gender Identity (Specify):
Social Security #:	Secondary	ID: Medicare #:
Last Name:	First Name	e: Middle Name:
Ethnicity	spanic or Latino	panic or Not Latino
Race	nerican Indian/Alaska Native	Asian Black or African American
☐ Na	ative Hawaiian/Other Pacific Islan	der White
*Blood Group: [☐ A- ☐ A+ ☐ B- ☐B+ ☐	AB-
	Transitional ABO / Rh +	Transitional ABO / Rh - Transitional ABO / Transitional Rh
☐ Group A/Transit	ional Rh 🔲 Group B/Transitional	Rh Group O/Transitional Rh Group AB/Transitional Rh
Patient Medica	I History	
List the patient	's admitting diagnosis. (Use ICD-	10 Diagnostic codes/descriptions)
Code:	Description: _	
Code:	Description: _	
Code:	Description: _	
List the patient	's underlying indication for transfu	sion. (Use ICD-10 Diagnostic codes/descriptions)
Code:	Description: _	
Code:	Description: _	
Code:	Description: _	
	's comorbid conditions at the time ICD-10 Diagnostic codes/descrip	e of the transfusion related to the adverse UNKNOWN (ions)
Code:	Description: _	
Code:	Description: _	
Code:	Description: _	
or institution is collected disclosed or released w Service Act (42 USC 24 Public reporting burden searching existing data may not conduct or sponumber. Send commer	I with a guarantee that it will be held in strict ithout the consent of the individual, or the in 2b, 242k, and 242m(d)). of this collection of information is estimated sources, gathering, and maintaining the dat nsor, and a person is not required to responts regarding this burden estimate or any other.	obtained in this surveillance system that would permit identification of any individual confidence, will be used only for the purposes stated, and will not otherwise be stitution in accordance with Sections 304, 306 and 308(d) of the Public Health to average 20 minutes per response, including the time for reviewing instructions, a needed, and completing and reviewing the collection of information. An agency do to a collection of information unless it displays a currently valid OMB control ner aspect of this collection of information, including suggestions for reducing this 21-8, Atlanta, GA 30333, ATTN: PRA (0920-0666).



ı	erformed during the current hospital					☐ NONE		
(codes/descriptions)							
	Code: Description:							
	Code: Description:							
	Additional Information							
	ansfusion History							
l	Has the patient received a previous tr			_	□ NO □ UNK	KNOWN		
	Blood Product:							
	Date of Transfusion:// UNKNOWN							
	Was the patient's adverse reaction transfusion-related?							
	If yes, provide information about the transfusion adverse reaction.							
	Type of transfusion adverse reaction	n: [Allergic	AHTR D	HTR DSTR	☐ FNHTR		
	☐ HTR ☐ TTI ☐ PTP		D _ TAD	☐ TA-GVHD	☐ TRALI	☐ UNKNOWN		
	OTHER Specify							
Re	action Details							
*Da	ite reaction occurred://	_ *Time	e reaction o	ccurred::_	Time un	known		
*Fa	cility location where patient was tr	ansfuse	d:					
Is this reaction associated with an incident?								
	Investigation Results							
	estigation Results				,			
Inv	restigation Results Transfusion related acute lung in	jury (TF	RALI)					
Inv		ijury (TF	RALI)		Test result positive			
Inv			RALI)	Cognate or	Test result positive No cognate or	Not tested for		
Inv		Not	,	Cognate or cross reacting	Test result positive No cognate or cross reacting	cognate		
Inv			RALI) Negative	Cognate or	Test result positive No cognate or			
Inv	Transfusion related acute lung in	Not Done	,	Cognate or cross reacting	Test result positive No cognate or cross reacting	cognate		
Inv	Transfusion related acute lung in	Not Done	Negative	Cognate or cross reacting	Test result positive No cognate or cross reacting	cognate		
Inv	Donor or unit HLA specificity Donor or unit HNA specificity	Not Done	Negative	Cognate or cross reacting	Test result positive No cognate or cross reacting	cognate		
Inv	Donor or unit HLA specificity Donor or unit HNA specificity Recipient HLA specificity	Not Done	Negative	Cognate or cross reacting	Test result positive No cognate or cross reacting	cognate		
Inv	Donor or unit HLA specificity Donor or unit HNA specificity Recipient HLA specificity Recipient HNA specificity *Case Definition (Check all that app	Not Done Done Doly) ry (ALI)	Negative □ □ □ □ orior to trans	Cognate or cross reacting antigen present	Test result positive No cognate or cross reacting	cognate		
Inv	Donor or unit HLA specificity Donor or unit HNA specificity Recipient HLA specificity Recipient HNA specificity *Case Definition (Check all that application) NO evidence of acute lung injuication ALI onset during or within 6 ho	Not Done Done Doly) ry (ALI)	Negative D orior to transessation of transes	Cognate or cross reacting antigen present	Test result positive No cognate or cross reacting antigen present	cognate		
Inv	Donor or unit HLA specificity Donor or unit HNA specificity Recipient HLA specificity Recipient HNA specificity *Case Definition (Check all that app NO evidence of acute lung inju ALI onset during or within 6 ho Hypoxemia – defined as PaO2	Not Done Doly) ry (ALI) urs of ce /FiO2 les	Negative D orior to transes sation of transes than or equal to the control of the control or equal to the control or equal t	Cognate or cross reacting antigen present Grantigen present Grantige	Test result positive No cognate or cross reacting antigen present	cognate		
Inv	Donor or unit HLA specificity Donor or unit HNA specificity Recipient HLA specificity Recipient HNA specificity *Case Definition (Check all that app NO evidence of acute lung injuent of the property of th	Not Done Doly) ry (ALI) urs of ce /FiO2 lesen satura	Negative D Orior to transes station of transes than or equation less than	Cognate or cross reacting antigen present Grantigen present Grantige	Test result positive No cognate or cross reacting antigen present	cognate		
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Inv	Donor or unit HLA specificity Donor or unit HNA specificity Recipient HLA specificity Recipient HNA specificity *Case Definition (Check all that app NO evidence of acute lung injuent of the property of th	Not Done Doly) ry (ALI) urs of ce /FiO2 lesen satura clinical ee eral infiltr	Negative Deprior to transessation of transes than or equition less that evidence trates	Cognate or cross reacting antigen present Grantigen present Grantige	Test result positive No cognate or cross reacting antigen present	cognate		
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Inv	Donor or unit HLA specificity Donor or unit HNA specificity Recipient HLA specificity Recipient HNA specificity *Case Definition (Check all that application) ALI onset during or within 6 ho Hypoxemia – defined as PaO2 Hypoxemia – defined as Oxyget Hypoxemia – defined as Other Radiographic evidence of bilate No evidence of left atrial hyper	Not Done Doly) ry (ALI) urs of ce /FiO2 les en satura clinical e eral infiltr tension (nat apply)	Negative D Orior to transes sation of transes than or equition less that evidence rates i.e., circulate	Cognate or cross reacting antigen present Grantigen present Grantige	Test result positive No cognate or cross reacting antigen present	cognate		



Cutaneous:	🗌 Edema 🔲 Flushing 🔲 Ja	aundice 🗌 Itching	☐ Hives ☐ Other rash			
Hemolysis/Hemorrhage:	nage: DIC Hemoglobinemia Positive antibody screen					
Pain:	☐ Abdominal pain ☐ Back	pain	☐ Infusion site pain			
Renal:	☐ Hematuria ☐ Hem	oglobinuria	Oliguria			
Respiratory:	☐ Bronchospasm ☐ Cough ☐	Shortness of breath	Other: (specify)			
*Severity						
Did the patient receive of	r experience any of the following?					
☐ No treatment requ	ired Symp	tomatic treatment only				
☐ Hospitalization, in	cuding prolonged hospitalization	Life-thre	eatening reaction			
☐ Disability and/or ir	☐ Disability and/or incapacitation ☐ Congenital anomaly or birth defect(s) of the fetus					
Other medically in	nportant conditions	n 🗌 Unknow	n or not stated			
*Imputability						
Which best describes the	relationship between the transfus	ion and the reaction?				
☐ There are no alterna	ative risk factors for ALI present.					
	f other causes for acute lung injury					
	in favor of a cause other than the t	•				
<u> </u>	evidence beyond reasonable doul					
☐ The relationship bet	ween the adverse reaction and the	e transfusion is unknowi	n or not stated.			
Did the transfusion occur at your facility? ☐ YES ☐ NO						
	<u> </u>					
Module-generated Desig						
NOTE: Designations for case	Inations definition, severity, and imputabili	ty will be automatically a				
NOTE: Designations for case application based on respons	nations definition, severity, and imputabilities in the corresponding investigat	ty will be automatically a ion results section above	e. 			
NOTE: Designations for case application based on respons *Do you agree with the	gnations definition, severity, and imputabilities in the corresponding investigate case definition designation?	ty will be automatically a	e. 			
NOTE: Designations for case application based on respons *Do you agree with the ^Please indicate your de	gnations definition, severity, and imputabilities in the corresponding investigate case definition designation? signation	ty will be automatically a ion results section above	e. S NO			
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*Do you agree with the *Please indicate your de *Do you agree with the *Please indicate your de *Do you agree with the *Please indicate your de *Do you agree with the *Please indicate your de *Do you agree with the *Please indicate your de Patient Treatment Did the patient receive treatment(s) Medication (Select Antipyretics Intravenous In Antithymocyte	definition, severity, and imputabilities in the corresponding investigate case definition designation? signation severity designation? signation imputability designation? signation the type of medication) Antihistamines Inotropes mmunoglobulin Intravenous signation	ty will be automatically a ion results section above	e. S NO S NO S NO D NO D NO D UNKNOWN nchodilator Diuretic	cs		
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☐ Renal replacement therapy (Select the type of therapy)									
_ □	- · · · · ·	Peritoneal [☐ Cor	itinuous Ven	o-Venous Hem	ofiltratio	n		
☐ Phlebotomy									
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	•		Doubtful	☐ Ruled Ou	ıt 🗆	Not determin	ed	
	of death:						_		
Was an	autopsy performed?	Yes	☐ No	•					
Component	Details								
*Was a partic	*Was a particular unit implicated in (i.e., responsible for) the adverse								
Transfusion Start and End Date/Time	*Component code (check system used)	Amount (Requ		number uired for ion and I)	*Unit expiration Date/Time	*Blood group of unit		Implic ated Unit?	
^IMPLICATED	UNIT								
//	☐ ISBT-128								
:	☐ Codabar	☐ Entire unit ☐ Partial unit				□ A-	□ A+ □ B-	Y	
/		mL				□в+	□ AB- □ AB+	'	
:					:	□ o-	□ O+ □ N/A		
	☐ ISBT-128								
:	☐ Codabar	☐ Entire unit				□ A-	□ A+ □ B-	NI NI	
/ /		Partial unitmL				□B+	□ AB- □ AB+	N	
:					:	П 0-	□ O+ □ N/A		
Custom Field	ds								
Label				Label					
		·/	-				//		
Comments									

