

Hemovigilance Module Adverse Reaction Hypotensive Transfusion Reaction

*Required for saving	
*Facility ID#: NHS	SN Adverse Reaction #:
Patient Information	
*Patient ID: Social Security #:	Secondary ID: Medicare #:
Last Name:	_
Ethnicity 🗌 Hispanic or Latin	o Not Hispanic or Not Latino
Race American Indian/	Alaska Native Asian Black or African American /Other Pacific Islander White
*Blood Group: 🗌 A- 🗌 A+	B- B+ AB- AB+ O- O+ Blood type not done
	BO / Rh + Transitional ABO / Rh - Transitional ABO / Transitional Rh Group B/Transitional Rh Group O/Transitional Rh Group AB/Transitional Rh
Patient Medical History	
List the patient's admitting dia	agnosis. (Use ICD-10 Diagnostic codes/descriptions)
Code:	Description:
Code:	Description:
Code:	Description:
List the patient's underlying ir	ndication for transfusion. (Use ICD-10 Diagnostic codes/descriptions)
Code:	Description:
Code:	Description:
Code:	Description:
List the patient's comorbid co reaction. (Use ICD-10 Diagno	nditions at the time of the transfusion related to the adverse UNKNOWN
Code:	Description:
Code:	Description:
Code:	Description:
of any individual or institution is collect stated, and will not otherwise be discl	untarily provided information obtained in this surveillance system that would permit identification ted with a guarantee that it will be held in strict confidence, will be used only for the purposes osed or released without the consent of the individual, or the institution in accordance with Public Health Service Act (42 USC 242b, 242k, and 242m(d)).
reviewing instructions, searching exist collection of information. An agency r unless it displays a currently valid OM	on of information is estimated to average 20 minutes per response, including the time for ting data sources, gathering and maintaining the data needed, and completing and reviewing the nay not conduct or sponsor, and a person is not required to respond to a collection of information B control number. Send comments regarding this burden estimate or any other aspect of this ggestions for reducing this burden to CDC, Reports Clearance Officer, 1600 Clifton Rd., MS D-74,

Atlanta, GA 30333 ATTN: PRA (0920-0666).

NHSN National Healthcare Safety Network		Form Approved OMB No. 0920-0666 Exp. Date: 12/31/22 www.cdc.gov/nhsn
	medical procedure including past procedures and procedures to be rent hospital or outpatient stay. (Use ICD-10 Procedure	
Code:	Description:	
Code:	Description:	
Code:	Description:	
Additional Information		
Transfusion History		
Blood Product: Date of Transfusion:	a previous transfusion?	Granulocyte
1	ion about the transfusion adverse reaction.	
Type of transfusion ad	verse reaction:	
Reaction Details		
*Date reaction occurred:_	// *Time reaction occurred:: [_ Time	e unknown
*Facility location where pa	atient was transfused:	
Is this reaction associated v	/ith an incident?	
Investigation Results		
* Hypotensive transf	usion reaction	
*Case Definition		
	d during or within 1 hour of cessation of transfusion:	
	reactions presenting with hypotension are excluded.	
Hypotension		
Check all that apply:		
Hypotension occu apply.	rs, does not meet the criteria above. Other, more specific reaction of	definitions do not
Other signs and symptoms	(check all that apply)	
Generalized:	Chills/rigors Fever Nausea/vomitin	g
Cardiovascular:	Shock	
Cutaneous:	Edema Image: Flushing Image: Jaundice Image: Other rash Image: Pruritus (itching) Image: Urticaria (hives)	1
Hemolysis/Hemorrhage:	 Disseminated intravascular coagulation Hemoglobinemi Positive antibody screen 	а
Pain:		fusion site pain
Renal:		iguria
Respiratory:		bugh



	🗌 Hypoxemia	Shortness of breath		
Other: (specify)				
*•				
* Severity Did the patient receive	or experience any of th	e following?		
No treatment receive			tmont only	
	inlcuding prolonged hos	Symptomatic trea	Life-threatening	n reaction
Disability and/or		•	aly or birth defect(s	-
	important conditions	Death		-
*Imputability	•			
• •	ne relationship between	the transfusion and the	reaction?	
		could explain hypotension		
There are other po cause.	otential causes present	that could explain hypote	ension, but transfu	sion is the most likely
Other conditions t	hat could readily explai	n hypotension are preser	nt.	
Evidence is clearly	y in favor of a cause otl	her than the transfusion,	but transfusion car	not be excluded.
There is conclusiv	ve evidence beyond rea	sonable doubt of a cause	e other than the tra	nsfusion.
The relationship b	etween the adverse rea	action and the transfusion	n is unknown or no	t stated.
How did the patient resp	pond the cessation of tr	ansfusion and supportive	e treatment?	
Responds rapidly	(i.e., within 10 minutes	s) to cessation of transfus	sion and supportive	e treatment.
The patient does	not respond rapidly to o	cessation of transfusion a	and supportive trea	tment.
Did the transfusion occu	ur at your facility?	YES NO		
When did the reaction c	occur in relation to the tr	ransfusion?		
	15 minutes after the sta			
Onset is between	15 minutes after start	and 1 hour after cessatio	n of transfusion.	
Module-generated Des				
NOTE: Designations for cas application based on respor				d in the NHSN
*Do you agree with th ^Please indicate your o		gnation?	☐ YES	□ NO
* Do you agree with th ^Please indicate your c		1?	YES	
* Do you agree with th ^Please indicate your o		ation?	☐ YES	
Patient Treatment	-			
Did the patient receive tre	eatment for the transfus	ion reaction?	YES 🗌 NO	
If yes, select treatment(
	t the type of medication	n)		
Antipyretics	Antihistamines [Inotropes/Vasopresso	rs 🗌 Bronchodila	tor Diuretics

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] Intravenous Immund] Antithymocyte globu	-		Cortico:	steroids	Antibio	tics
🗌 Volu	me resuscitation (Intr	avenous colloid	ls or crystalloids)				
C Res	piratory support <i>(Sele</i>] Mechanical ventilati		upport) nvasive ventilation	Oxyger	n		
	al replacement therap] Hemodialysis		rpe of therapy) Continuous Ven	io-Venous Hen	nofiltratio	n	
	ebotomy er Specify:						
Outcome							
Cause		e	ion to death:	Minor or no se	·	☐ Not dete	
Component							
"was a partic	cular unit implicated	d in (i.e., respo	onsible for) the a	dverse			
reaction?	cular unit implicated	d in (i.e., respo	-	dverse	Yes	s 🗌 No	□ N/A
	*Component code (check system used)	d in (i.e., respo Amount transfused at reaction onset	AUnit number (Required for Infection and TRALI)	dverse *Unit expiration Date/Time		d group	N/A Implic ated Unit?
reaction? Transfusion Start and End	*Component code (check system used)	Amount transfused at	^ Unit number (Required for Infection and	*Unit expiration	*Bloo	d group	Implic ated
reaction? Transfusion Start and End Date/Time	*Component code (check system used)	Amount transfused at	^ Unit number (Required for Infection and	*Unit expiration	*Bloo	d group	- Implic ated Unit?
reaction? Transfusion Start and End Date/Time	*Component code (check system used) UNIT	Amount transfused at reaction onset	^ Unit number (Required for Infection and	*Unit expiration	*Bloo of uni A- B+ 0- A- B+ B+	d group it A+ □ B- AB- □ AB O+ □ N// A+ □ B- AB- □ AB	
reaction? Transfusion Start and End Date/Time	*Component code (check system used) UNIT DISBT-128 Codabar ISBT-128 DISBT-128 Codabar Codabar	Amount transfused at reaction onset	^ Unit number (Required for Infection and	*Unit expiration	*Bloo of uni	d group it A+ □ B- AB- □ AB O+ □ N// A+ □ B- AB- □ AB	High product of the second sec
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