

*Required for saving

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Hemovigilance Module Adverse Reaction Hypotensive Transfusion Reaction

*Facility ID#: NHSN Ac	dverse Reaction #:							
Patient Information								
*Patient ID:	*Gender: M F Oth	ner *Date of Birth://						
Social Security #:	Secondary ID:	Medicare #:						
Last Name:	First Name:	Middle Name:						
Ethnicity Hispanic or Latino	Not Hispanic or Not Latino							
Race American Indian/Alaska Native Asian Black or African American								
Native Hawaiian/Other Pacific Islander White								
		O+ Blood type not done Transitional ABO / Transitional						
Transitional ABO / Group A/Transitional Group	Rh + Transitional ABO / Rh - D B/Transitional Crays O/Trans	Rh						
Rh Rh	Group O/Trans	itional Rh Group AB/Transitional Rh						
Patient Medical History								
List the patient's admitting diagnos	sis. (Use ICD-10 Diagnostic codes/de	scriptions)						
Code:	Description:							
Code:	Description:							
Code:	Description:							
List the patient's underlying indicat	tion for transfusion. (Use ICD-10 Diag	gnostic codes/descriptions)						
Code:	Description:							
	Description:							
	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:							
Code:	Description:							
Code:	Description:							
	ith a guarantee that it will be held in strict co or released without the consent of the indivi							
collection of information. An agency may no unless it displays a currently valid OMB con	ata sources, gathering and maintaining the option of conduct or sponsor, and a person is not retrol number. Send comments regarding thions for reducing this burden to CDC, Repor	utes per response, including the time for data needed, and completing and reviewing the required to respond to a collection of information is burden estimate or any other aspect of this its Clearance Officer, 1600 Clifton Rd., MS D-						







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	medical procedure including past procedures and procedures to be rent hospital or outpatient stay. (Use ICD-10 Procedure	UNKNOWN NONE							
Code:	Description:								
Code:	Description:								
Code:									
Additional Information									
Transfusion History									
Has the patient received	a previous transfusion? YES NO UN	IKNOWN							
Blood Product: WB RBC Platelet Plasma Cryoprecipitate Granulocyte									
Date of Transfusion:/ UNKNOWN									
Was the patient's adverse reaction transfusion-related?									
If yes, provide informat	on about the transfusion adverse reaction.								
Type of transfusion adv	rerse reaction: Allergic AHTR DHTR DSTF	R FNHTR							
HTR	PTP TACO TAD TA-GVHD TRALI	UNKNOWN							
OTHER Spe	cify								
Reaction Details									
*Date reaction occurred:_	_// *Time reaction occurred: : : Time t	unknown							
*Facility location where pa	tient was transfused:								
Is this reaction associated w	rith an incident? Yes No If Yes, Incident #:								
Investigation Results									
* Hypotensive transf	usion reaction								
*Case Definition									
Check all that occurred	during or within 1 hour of cessation of transfusion:								
All other adverse	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 def	initions do not							
Other signs and symptoms:	(check all that apply)								
Generalized:	Chills/rigors Fever Nausea/vomiting								
Cardiovascular:	Shock								
Cutaneous:	Edema Flushing Jaundice								
0 0.10.1.10 0 0.10.1	Other rash Pruritus (itching) Urticaria (hives)								
Hemolysis/Hemorrhage:	Disseminated intravascular coagulation Hemoglobinemia								
	Positive antibody screen								
Pain:		sion site pain							
Renal:	Hematuria Hemoglobinuria Olig								
Respiratory:	Bilateral infiltrates on chest x-ray Bronchospasm Cough								
Į.	Hypoxemia Shortness of breath	9							



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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 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 reaction?							
The patient has no other conditions that could explain hypotension. There are other potential causes present that could explain hypotension, but transfusion is the most likely							
cause.							
Other conditions that could readily explain hypotension are present.							
Evidence is clearly in favor of a cause other than the transfusion, but transfusion cannot be excluded.							
There is conclusive evidence beyond reasonable doubt of a cause other than the transfusion.							
The relationship between the adverse reaction and the transfusion is unknown or not stated.							
How did the patient respond the cessation of transfusion and supportive treatment?							
Responds rapidly (i.e., within 10 minutes) to cessation of transfusion and supportive treatment.							
The patient does not respond rapidly to cessation of transfusion and supportive treatment.							
Did the transfusion occur at your facility? YES NO							
When did the reaction occur in relation to the transfusion?							
Occurs less than 15 minutes after the start of the transfusion.							
Onset is between 15 minutes after start and 1 hour after cessation of transfusion.							
Module-generated Designations 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 <u>case definition</u> designation?							
^Please indicate your designation							
*Do you agree with the <u>severity</u> designation?							
^Please indicate your designation							
*Do you agree with the imputability designation?							
^Please indicate your designation Patient Treatment							
Did the patient receive treatment for the transfusion reaction? Lyes, select treatment(s):							
Medication (Select the type of medication)							
Antipyretics Antihistamines Inotropes/Vasopressors Bronchodilator Diuretics							
Intravenous Intravenous steroids Corticosteroids Antibiotics							



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	nmunoglobulin Antithymocyte globu	ulin Cyclo	osporin	Other					
Volume resuscitation (Intravenous colloids or crystalloids)									
			,						
Respiratory support (Select the type of support) Mechanical ventilation Noninvasive ventilation Oxygen									
Ren	al replacement therap	oy <i>(Select the ty</i> Peritoneal		no-Venous Hemo	ofiltration				
Phlebotomy									
Other Specify:									
Outcome									
*Outcome:		lajor or long-terr	n sequelae	Minor or no sec	quelae 🗌 Not deterr	nined			
Date of									
1 11^	ecipient died, relation Definite Probabl	. —		Ruled Out	Not determine	a d			
	of death:	e Possib	le Doubtful	Ruled Out	Not determine	au			
	autopsy performed?	Yes	No						
*Was a particular unit implicated in (i.e., responsible for) the adverse reaction? Yes No N/A									
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	Implic ated Unit?			
AIMPLICATED	•				•				
^IMPLICATED	UNIT					'			
// ::	UNIT ISBT-128 Codabar	Entire unit		<u> </u>	A- A+ B-	Y			
//	ISBT-128				A- A+ B- B+ AB- AB+ O- O+ N/A	Y			
//	ISBT-128	Partial unit			B+ AB- AB+	Y			
//:::	ISBT-128 Codabar	Partial unitmL Entire unit		:	B+ AB- AB+				
//	ISBT-128 Codabar ISBT-128	Partial unitmL		: !	B+ AB- AB+ O- O+ N/A A- A+ B-	Y N			
	ISBT-128 Codabar ISBT-128	Partial unitmL Entire unit		:	B+ AB- AB+ B- B+ AB- AB+				
	ISBT-128 Codabar ISBT-128 Codabar Codabar	Partial unitmL		:	B+ AB- AB+ O- O+ N/A A- A+ B-				
	ISBT-128 Codabar ISBT-128 Codabar Codabar	Partial unitmL		: :	B+ AB- AB+ B- B+ AB- AB+				
	ISBT-128 Codabar ISBT-128 Codabar Codabar	Partial unitmL		; ; ;	B+ AB- AB+ B- B+ AB- AB+				
	ISBT-128 Codabar ISBT-128 Codabar Codabar	Partial unitmL			B+ AB- AB+ B- B+ AB- AB+				



