

Hemovigilance Module Adverse Reaction Febrile Non-hemolytic Transfusion Reaction

*Required for saving	-	
*Facility ID#: NHSN A	Adverse Reaction #:	
Patient Information		
*Patient ID:	_	*Date of Birth://
*Sex at Birth:	n	*Gender Identity (Specify):
Social Security #:	Medicare #:	
Last Name:	First Name:	Middle Name:
Ethnicity 🗌 Hispanic or Latino	Not Hispanic or Not Latino	
Race American Indian/Alas	ska Native Asian Bl ner Pacific Islander W	
*Blood Group:	B- 🛛 B+ 🗋 AB- 🗌 AB+ 🗌 O	- O+ Blood type not done
	—	Transitional ABO / Transitional Rh itional Rh Group AB/Transitional Rh
Patient Medical History		
List the patient's admitting diagne	osis. (Use ICD-10 Diagnostic codes/de	escriptions)
Code:	Description:	
Code:	Description:	
Code:	Description:	
List the patient's underlying indic	ation for transfusion. (Use ICD-10 Diag	gnostic codes/descriptions)
Code:	Description:	
Code:	Description:	
Code:	Description:	
List the patient's comorbid condit reaction. (Use ICD-10 Diagnostic	tions at the time of the transfusion rela c codes/descriptions)	ted to the adverse UNKNOWN
Code:	Description:	
Code:	Description:	
Code:	Description:	
or institution is collected with a guarantee that i	t will be held in strict confidence, will be used only e individual, or the institution in accordance with Se	
searching existing data sources, gathering, and may not conduct or sponsor, and a person is no number. Send comments regarding this burder	I maintaining the data needed, and completing and ot required to respond to a collection of information	information, including suggestions for reducing this



Febrile Non-hemolytic Transfusion Reaction

List the patient's relevant medical procedure including past procedures and procedures to performed during the current hospital or outpatient stay. (Use ICD-10 Procedure codes/descriptions)				
Code: Description:				
Code: Description:				
Code: Description:				
Additional Information				
Transfusion History				
Has the patient received a previous transfusion?	UNKNOWN			
Blood Product: 🛛 WB 🗌 RBC 🔄 Platelet 🗌 Plasma 🗌 Cryoprecipita	te 🗌 Granulocyte			
Date of Transfusion:// UNKNOWN				
Was the patient's adverse reaction transfusion-related?	10			
If yes, provide information about the transfusion adverse reaction.				
	STR _ FNHTR			
OTHER Specify	· · · · · · · · · · · · · · · · · · ·			
Reaction Details				
*Date reaction occurred:// *Time reaction occurred:: [] Ti	me unknown			
*Facility location where patient was transfused:				
Is this reaction associated with an incident? Yes No If Yes, Incident #:				
Investigation Results				
* Febrile non-hemolytic transfusion reaction (FNHTR)				
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*Case Definition	°F) from pre-			
*Case Definition Check all that occurred during or within 4 hours of cessation of transfusion: Fever (greater than or equal to 38°C/100.4°F oral and a change of at least 1°C/1.8	°F) from pre-			
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🗌 Hyr	ooxemia		Shortness o	f breath	
Other: (specify)					
*Severity					
Did the patient receive or experie	ence any of the fo	ollowing?			
No treatment required		• •	tic treatment on	-	
Hospitalization, inlcuding p	•			hreatening re	
Disability and/or incapacita			l anomaly or birt	. ,	
Other medically important	conditions	Death	Unkn	own or not st	tated
*Imputability					
Which best describes the relation Patient has no other condit					
There are other potential ca likely cause.	auses present th	at could explai	n signs/sympton	ns, but transf	usion is the most
Other present causes are r	nost likely, but tr	ansfusion canr	not be ruled out.		
Evidence is clearly in favor					
There is conclusive evidenc					
The relationship between the	ie adverse reacti	ion and the trar	nsfusion is unkno	own or not st	ated.
Did the transfusion occur at you	ur facility?]YES] NO		
Module-generated Designation	IS				
NOTE: Designations for case definition	on, severity, and				า the NHSN
application based on responses in the		•	_		
* Do you agree with the <u>case de</u> ^Please indicate your designatio		ation?		YES	□ NO
* Do you agree with the <u>severit</u> ^Please indicate your designatio	-			YES	□ NO
*Do you agree with the <i>imputa</i>		on?		YES	
^Please indicate your designatio					
Patient Treatment					
Did the patient receive treatment for	or the transfusion	reaction?	☐ YES	□ NO □	
If yes, select treatment(s):					
Medication (Select the type			_		_
	histamines 🗌 I	•	·	ronchodilator	
Intravenous Immunog		venous steroid	—	steroids	Antibiotics
Antithymocyte globulii	n 🗌 Cyclosp	oorin 🗌	Other		
Volume resuscitation (Intrav	enous colloids c	or crystalloids)			
Respiratory support (Select Mechanical ventilation	<u> </u>	oo <i>rt)</i> asive ventilation	n 🗌 Oxyger	١	
Renal replacement therapy					
🗌 Hemodialysis 🗌 Pe	eritoneal	Continuous Ve	no-Venous Hem	onitration	
CDC 57 211 Day 2 10 2	Daga	2 of 1			

NHSNForm Approved OMB No. 0920-0666NATIONAL HEALTHCARE SAFETY WORKExp. Date: 12/31/2026 www.cdc.gov/nhsn							-0666 /2026		
	ebotomy								
	Outcome								
*Outcome:	Death M	ajor or long-tern	0 600] Minor or no se	مدامینه		determ	nined
Date of		/	n sequ			queide		uetern	inied
	recipient died, relation	, ship of transfus	ion to	death:					
	Definite Probable	•		Doubtful	🗌 Ruled Ou	t 🗌] Not det	termine	d
Cause	of death:								
Was ar	autopsy performed?	🗌 Yes	🗌 No)					
Component									
*Was a partion?	cular unit implicate	d in (i.e., respo	onsibl	e for) the a	dverse	🗌 Ye	s 🗌 N	lo [N/A
Transfusion Start and End Date/Time	*Component code (check system used)	Amount transfused at reaction onset	(Requ	t number uired for ion and I)	*Unit expiration Date/Time	*Bloc of un	od group lit		Implic ated Unit?
^IMPLICATED	UNIT								
//	□ ISBT-128			·					
	Codabar	☐ Entire unit ☐ Partial unit			//	🗆 A-	🗆 A+ 🛛] B-	V
//		mL				□в+	🗆 АВ- 🛛	AB+	
::					:	0-	0+] N/A	
//	□ ISBT-128			·					
:	Codabar	Entire unit			//	🗆 A-	□ A+ [] B-	N
//		mL				□в+	🗆 АВ- 🛛	AB+	
:::					::	□ 0-	□ O+ [] N/A	
Custom Field	Custom Fields								
Label				Label					
		//	-				/	_/	
Comments									