

Hemovigilance Module Adverse Reaction Delayed Hemolytic Transfusion Reaction

*Required for saving NHSN Adverse Reaction #: *Facility ID#: _____ **Patient Information** *Gender: ☐ M ☐ F ☐ Other *Date of Birth: __/__/ *Patient ID: ____ Social Security #: _____ Secondary ID: _____ Medicare #: _____ Last Name: First Name: Middle Name: ☐ Not Hispanic or Not Latino Ethnicity Hispanic or Latino Race American Indian/Alaska Native Asian ☐ Black or African American Native Hawaiian/Other Pacific Islander ☐ White ***Blood Group:** □ A- □ A+ □ B- □ B+ □ AB- □ AB+ □ O- □ O+ ☐ Blood type not done Transitional ABO / Rh + ☐ Transitional ABO / Rh -☐ Transitional ABO / Transitional Rh ☐ Group A/Transitional Rh ☐ Group B/Transitional Rh ☐ Group O/Transitional Rh ☐ Group AB/Transitional Rh Patient Medical 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 transfusion. (Use ICD-10 Diagnostic codes/descriptions) Code: Description: Code: _____ Description: Description: _____ Code: _____ List the patient's comorbid conditions at the time of the transfusion related to the adverse reaction. (Use ICD-10 Diagnostic codes/descriptions) □ NONE Code: _____ Description: Code: _____ Description: 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)). 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 D-74,

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



| | ent hospital or outpatient stay. (Use ICD-10 Procedure NONE | WN |
|---------------------------------|---|------|
| Code: | | |
| Code: | | |
| Code: | Description: | |
| Additional Information | | |
| Transfusion History | | |
| Has the patient received a | previous transfusion? | |
| Blood Product: |] WB □ RBC □ Platelet □ Plasma □ Cryoprecipitate □ Granulo | cyte |
| Date of Transfusion: | // | |
| Was the patient's advers | se reaction transfusion-related? | |
| If yes, provide informatio | on about the transfusion adverse reaction. | |
| Type of transfusion adve | erse reaction: | |
| ☐ HTR ☐ TTI | ☐ PTP ☐ TACO ☐ TAD ☐ TA-GVHD ☐ TRALI ☐ UNKNOV | ٧N |
| ☐ OTHER Spec | ify | |
| Reaction Details | | |
| *Date reaction occurred: | _// *Time reaction occurred: : : Time unknown | |
| *Facility location where pat | ient was transfused: | _ |
| Is this reaction associated wit | th an incident? | |
| Investigation Results (O | only answer questions listed under the selected reaction type.) | |
| | sfusion reaction (DHTR) | |
| | Non-immune (specify) | |
| *Case Definition | | |
| | occurred between 24 hours and 28 days after cessation of transfusion: | |
| Positive direct antigle | · | |
| | blood cell alloantibody in recipient serum | |
| - | with alloantibody present on the transfused red blood cells | |
| | st-transfusion hemoglobin level or rapid fall in hemoglobin back to pre-transfusion leve | عاد |
| | ned appearance of spherocytes | ,10 |
| | iod appearance of apriciocytes | |
| Check all that apply: | | |
| ☐ Incomplete laborator | | |
| ☐ DHTR is suspected, | but reported symptoms, test results, and/or available information are not sufficient | t |
| Other signs and symptoms: (c | | |
| Generalized: | Chills/rigors Fever Nausea/vomiting | |
| Cardiovascular: | Blood pressure decrease Shock | |
| Cutaneous: | ☐ Edema ☐ Flushing ☐ Jaundice ☐ Drugitus (itabing) ☐ Urticogic (biyos) | |
| Hemolysis/Hemorrhage: | ☐ Other rash ☐ Pruritus (itching) ☐ Urticaria (hives) ☐ Disseminated intravascular coagulation ☐ Hemoglobinemia | |
| i ioiiioiyolo/i lolllollllage. | | |



| Renal: | Hematuria | ☐ Hemoglo | obinuria 🔲 C | Oliguria | | |
|---|---|---------------------|--------------------------|-----------------------|--|--|
| Respiratory: | ☐ Bilateral infiltrates on chest x-ray ☐ Bronchospasm ☐ Cough | | | | | |
| respiratory. | Hypoxemia | ☐ Shortnes | ss of breath | | | |
| Other: (specify) | | | | | | |
| *Severity | | | | | | |
| Did the patient receive or e | experience any of the fol | llowing? | | | | |
| ☐ No treatment requi | red | ☐ Symptomatic | treatment only | | | |
| ☐ Hospitalization, inlo | cuding prolonged hospita | alization | Life-threateni | ng reaction | | |
| ☐ Disability and/or inc | capacitation | ☐ Congenital a | nomaly or birth defect | (s) of the fetus | | |
| Other medically im | portant conditions | ☐ Death | Unknown or r | not stated | | |
| *Imputability | | | | | | |
| Which best describes the re | elationship between the | transfusion and | the reaction? | | | |
| ☐ No other explanation | for symptoms or newly- | identified antibod | ly is present. | | | |
| | tion for symptoms or nev | wly-identified anti | body is present, but tr | ansfusion is the most | | |
| likely cause. | or symptoms or newly-ide | entified antibody | are more likely but tra | ansfusion cannot be | | |
| ruled out. | Toymptomo of nowly las | critined artibody | are more intery, but the | andradion dames be | | |
| ☐ Evidence is clearly in | favor of a cause other t | han the transfusi | on, but transfusion ca | nnot be excluded. | | |
| ☐ There is conclusive evidence beyond reasonable doubt of a cause other than the transfusion. | | | | | | |
| ☐ The relationship betw | een the adverse reactio | n and the transfu | usion is unknown or no | ot stated. | | |
| Did the transfusion occur a | t your facility? | S 🗌 NO | | | | |
| Module-generated Design | nations | | | | | |
| NOTE: Designations for case | | imputability will b | e automatically assign | ned in the NHSN | | |
| application based on response | es in the corresponding i | investigation resu | ults section above. | | | |
| *Do you agree with the <u>c</u> | | ation? | ☐ YES | □NO | | |
| ^Please indicate your des | signation | | | | | |
| *Do you agree with the s | | | ☐ YES | □NO | | |
| ^Please indicate your des | ignation | | | | | |
| *Do you agree with the <u>i</u> | | n? | ☐ YES | □NO | | |
| ^Please indicate your des | ignation | | | | | |
| Patient Treatment | | | | | | |
| Did the patient receive treat | | reaction? | ☐ YES ☐ NO | UNKNOWN | | |
| If yes, select treatment(s): | he type of medication) | | | | | |
| <u> </u> | | notrones//asonr | essors Rronchod | ilator | | |
| ☐ Antipyretics☐ Antihistamines☐ Intravenous Immunoglobulin☐ Intravenous Immunoglobulin | | | | | | |
| ☐ Antithymocyte globulin ☐ Cyclosporin ☐ Other | | | | | | |
| | _ , , | _ | - | | | |
| ∪ volume resuscitation | n (Intravenous colloids o | r crystalloids) | | | | |
| ☐ Respiratory support | (Select the type of supp | ort) | | | | |
| ☐ Mechanical ve | ntilation | sive ventilation | ☐ Oxygen | | | |



| ☐ Renal replacement therapy (Select the type of therapy) ☐ Hemodialysis ☐ Peritoneal ☐ Continuous Veno-Venous Hemofiltration | | | | | | | | | | |
|--|-------------------------|-----------------------------|------|----------------------|------------|----------|-----------|------------|--------|--|
| ☐ Phlebotomy ☐ Other Specify: | | | | | | | | | | |
| Outcome | | | | | | | | | | |
| *Outcome: Death Major or long-term sequelae Minor or no sequelae Not determined | | | | | | | | | | |
| | Date of Death:/ | | | | | | | | | |
| | ecipient died, relation | • | | <u></u> | | _ | | | | |
| | Definite Probable | e | e [| Doubtful | Ruled Out | |] Not de | etermine | ed | |
| | of death: | | | | | | | | | |
| Was an autopsy performed? | | | | | | | | | | |
| Component Details | | | | | | | | | | |
| *Was a particular unit implicated in (i.e., responsible for) the adverse reaction? | | | | | | | | | | |
| Transfusion | | Amount | | number | *Unit | | | | Implic | |
| Start and End | *Component code | transfused at | | iired for ion and | expiration | *Bloo | d grou |) | ated | |
| Date/Time | (check system used) | reaction onset | TRAL | l) | Date/Time | of un | it | | Unit? | |
| ^IMPLICATED | UNIT | Г | | | Г | | I | | | |
| / | ☐ ISBT-128 | | | | | | | | | |
| : | ☐ Codabar | ☐ Entire unit☐ Partial unit | | | / | □ A- | □ A+ | □ B- | Υ | |
| / | | mL | | | | □в+ | ☐ AB- | ☐ AB+ | | |
| : | | | | | :: | □ o- | □ O+ | □ N/A | | |
| // | ☐ ISBT-128 | | | | | | | | | |
| : | ☐ Codabar | ☐ Entire unit | | | / | □ A- | □ A+ | □ B- | N. | |
| / / | | ☐ Partial unit mL | | | | □в+ | ☐ AB- | □ AB+ | N | |
| : | | | | | | П 0- | _ □ 0+ | _ □ N/A | | |
| Custom Field | ds | | | | <u> </u> | <u> </u> | <u> </u> | | | |
| Label | | | | Label | | | | | | |
| Laber Laber | | | | | | | | | | |
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| Comments | | | | | | | | | | |
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