

Hematopoietic Cellular Transplant (HCT) Infusion

Registry Use Only	OMB No: 0915-0310 Expiration Date: 1/31/2020	
Sequence Number:	Public Burden Statement: An agency may not conduct or sponsor, and a person is	
Date Received:	not required to respond to, a collection of information unless it displays a currently valid OMB control number. The OMB control number for this project is 0915-0310. Public reporting burden for this collection of information, in combination with the IDM Form 2004 and HLA Typing Form 2005, is estimated to average 1.5 hours per response, including the time for reviewing instructions, searching existing data sources, and completing and reviewing the collection of information. Send comme regarding this burden estimate or any other aspect of this collection of information including suggestions for reducing this burden, to HRSA Reports Clearance Office 5600 Fishers Lane, Room 10-33, Rockville, Maryland, 20857. Expiration date: 1/31/2020	
CIBMTR Center Number:		
CIBMTR Recipient ID:		
Event Date://		
HCT type (check only one) ☐ Autologous ☐ Allogeneid	c, unrelated	
Product type (check only one) Bone marrow PBSC Single cord blood unit Other product. Specify:		
NMDP Product: ☐ Yes ☐ No		
Product Identifiers:		
NMDP cord blood unit ID:		
NMDP donor ID:		
Non-NMDP unrelated donor ID:		
Non-NMDP cord blood unit ID:		
GRID (optional):		
ISBT DIN:		
Registry or UCB Bank ID:		
Donor DOB://MM / _DD		
Donor Age:	d) ☐ Years	
Donor Sex: ☐ Male ☐ Female		

CIBM	TR Center Number:	: CIBMTR Recipient ID:
A se	eries of collections sh	HCT product is infused, each product type must be analyzed and reported separately. ould be considered a single product when they are all from the same donor and use the same collection method lization, if applicable), even if the collections are performed on different days.
Pre	-Collection Therapy	
1.	donors only Yes	growth and mobilizing factors, prior to any stem cell harvest, to enhance the product collection for this HCT? Allogeneic 2. Specify growth and mobilizing factor(s) (Check all that apply)
	□ No	G-CSF (filgrastim, Neupogen) Pegylated G-CSF (pegfilgrastim, Neulasta) Plerixafor (Mozobil) Other growth or mobilizing factor(s) 3. Specify other growth or mobilizing factor(s):
		——————————————————————————————————————
Pro	duct Collection	
4.		or other agents added to the product between collection and infusion? 6. Specify anticoagulant(s): (check all that apply) Acid citrate dextrose (ACD, ACD-A) Citrate phosphate dextrose (CPD, CPD-A) Ethylenediaminetetraacetic acid (EDTA) Heparin Other 7. Specify other anticoagulant:
Pro	duct Transport and Re	occint
FIU		
8.	Was this product colled	9. Date of receipt of product at your facility:///

CIBMTR Center Number:	CIBMTR Recipient ID:
	11. Specify the shipping environment of the product(s) Room temperature Cooled (refrigerator temperature, not frozen) Frozen (cryopreserved) Other shipping environment 12. Specify other shipping environment: - If product is cord blood, go to question 13; all other products
	go to question 22 13. Was there any indication that the environment within the shipper was outside the expected temperature range for this product at any time during shipment? ☐ Yes ☐ No 14 Were the secondary containers (e.g., insulated shipping containers and unit cassette) intact when they arrived at your center?
	☐ Yes ☐ No
	15. Was the cord blood unit stored at your center prior to thawing? (Cord blood units only) Yes →
	Report the total number of cells (not cells per kilogram) prior to cryopreservation: (Information provided for the unit by the cord blood bank).
	19. Total nucleated cells: • x 10 (Includes nucleated recand nucleated white cells) (Cord blood units only)
	20. CD34+ cells (cord blood units only) Done

ואוטוע	TR Center Number:	CIBMTR Recipient ID:
Prod	duct Processing / Man	ipulation
22.	_	ed from a cryopreserved state prior to infusion?
	☐ Yes — ➤ ☐ No	23. Was the entire product thawed? ☐ Yes ☐ No → 24. Specify the percent of the product that was thawed? (Cord blood units only) ☐ 80% ☐ 20% ☐ Other percent → 25. Specify other percent: %
		26. Date thawing process initiated:////
		27. Time at initiation of thaw (24-hour clock): : :
		28. Time of thaw completion (24-hour clock): : :
		29. What method was used to thaw the product? Waterbath Electric warmer Other method — 30. Specify other method:
		31. Did any incidents, or product complaints occur while preparing or thawing the product? yes no
32.	_	essed prior to infusion?
	☐ Yes ———————————————————————————————————	33. Specify processing: (check all that apply) Buffy coat enriched (buffy coat preparation) Diluted Plasma reduced RBC reduced Washed
34.	Was the product mani	ipulated prior to infusion?
	☐ Yes → No	35. Specify manipulations performed: (check all that apply) Antibodies - Go to question 36 Ex-vivo expansion - Go to question 36 Genetic manipulation (gene transfer / transduction) - Go to question 36 CD34 enriched (CD34+ selection) - Go to question 36 Ex-vivo T-cell depletion - Go to question 38 Other manipulation - Go to question 40

CIBMTR Center Number:	CIBMTR Recipient ID:
Г	
	36. Specify antibodies used: (check all that apply) Anti CD3 Anti CD4 Anti CD8 Anti CD19 Anti CD34 Anti CD34
	□ α/β Antibody □ Anti CD52 □ Other antibody → 37. Specify other antibody:
	38. Specify T-cell depletion method: Antibody affinity column Immunomagnetic beads Other method — 39. Specify other method:
	40. Specify other cell manipulation:
Product Analysis (All Produc	ets)
	he product preparation phase that the product was analyzed: plood only)
42. Date of product analysis:	
	olus additives :
In this section, report the total	al number of cells (not cells per kilogram) and do not correct for viability.
44. Total nucleated cells (TN	C) (Includes nucleated red and nucleated white cells)
☐ Done ——> ☐ Not done	45. Total nucleated cells: x 10
	46. Viability of cells
	☐ Done → 47. Viability of cells: %
	☐ Not done ☐ Unknown 48. Method of testing cell viability:
	☐ Flow cytometry based ☐ Trypan blue
	Other method — 49. Specify other method:
1	

JIBM	TR Center Number:	CIBMTR Recipient ID:
50.	Nucleated white blood Done Not done	ells 51. Total number of nucleated white blood cells: • x 10
52.	Mononuclear cells Done Not done	53. Total number of mononuclear cells: ● x 10
54.	Nucleated red blood c	;
	☐ Done → ►	55. Total number of nucleated red blood cells: • x 10
56.	CD34+ cells Done Not done	57. Total number of CD34+ cells: • x 10 58. Viability of cells Done → Not done Unknown 59. Viability of cells: % 60. Method of testing cell viability: Flow cytometry based Trypan blue Other method → 61. Specify other method:
62.	CD3+ cells Done Not done	63. Total number of CD3+ cells: • x 10 64. Viability of cells
		□ Done → 65. Viability of cells:

	CIBMTR Recipient ID:
68. CD3+CD4+ cells Done Not done	69. Total number of CD3+CD4+: x 10 70. Viability of cells Done → Not done Unknown 71. Viability of cells: % 72. Method of testing cell viability: Flow cytometry based Trypan blue Other method → 73. Specify other method:
74. CD3+CD8+ cells ☐ Done → ☐ Not done	75. Total number of CD3+CD8+: • x 10 76. Viability of cells
	76. Viability of cells □ Done → □ Not done □ Unknown 77. Viability of cells: % 78. Method of testing cell viability: □ Flow cytometry based □ Trypan blue □ Other method → 79. Specify other method:
80. Were the colony-form Yes No	g units (CFU) assessed after thawing? (cord blood units only) 81. Was there growth?
	82. Total CFU-GM ☐ Done → ☐ Not done 83. Total CFU-GM: • x 10
	84. Total CFU-GEMM ☐ Done → ☐ Not done 85. Total CFU-GEMM: • x 10
	86. Total BFU-E □ Done →

☐ Yes ——— ☐ No	Specify organ	uism code(s):
☐ Pending	89	90 91 92
Unknown		93. Specify organism:
		‡ The codes for "other organism, specify" (codes 198, 209, 219 and 259) she rarely be needed; check with your microbiology lab or HCT physician befor using them.
		Codes for Commonly Reported Organisms
		Bacterial Infections
		☐ 121 Acinetobacter
		☐ 122 Actinomyces
		☐ 123 Bacillus
		☐ 124 Bacteroides (gracillis, uniformis, vulgaris, other species)
		☐ 125 Bordetella pertussis (whooping cough)
		☐ 126 Borrelia (Lyme disease)
		☐ 127 Branhamella or Moraxella catarrhalis (other species)
		128 Campylobacter (all species)
		☐ 129 Capnocytophaga
		☐ 171 Chlamydia pneumoniae
		☐ 172 Other chlamydia, specify
		113 Chlamydia, NOS
		☐ 130 Citrobacter (freundii, other species)
		☐ 131 Clostridium (all species except difficile)
		☐ 132 Clostridium difficile
		☐ 173 Corynebacterium jeikeium
		☐ 133 Corynebacterium (all nondiptheria species)
		☐ 101 Coxiella
		☐ 134 Enterobacter
		☐ 177 Enterococcus, vancomycin resistant (VRE)
		☐ 135 Enterococcus (all species)
		☐ 136 Escherichia (also E. coli)
		☐ 137 Flavimonas oryzihabitans
		☐ 138 Flavobacterium
		☐ 139 Fusobacterium
		☐ 144 Haemophilus (all species, including influenzae)
		☐ 145 Helicobacter pylori
		☐ 146 Klebsiella
		147 Lactobacillus (bulgaricus, acidophilus, other species)
		102 Legionella
		☐ 103 Leptospira
		☐ 148 Leptotrichia buccalis

☐ 104 Listeria
☐ 150 Methylobacterium
☐ 151 Micrococcus, NOS
☐ 112 Mycobacterium avium–intracellulare (MAC, MAI)
☐ 174 Mycobacterium species (cheloneae, fortuitum, haemophilum, kansasii, mucogenicum)
☐ 110 Mycobacterium tuberculosis (tuberculosis, Koch bacillus)
☐ 175 Other mycobacterium, specify
☐ 176 Mycobacterium, NOS
☐ 105 Mycoplasma
☐ 152 Neisseria (gonorrhoea, meningitidis, other species)
☐ 106 Nocardia
☐ 153 Pasteurella multocida
☐ 154 Propionibacterium (acnes, avidum, granulosum, other species)
☐ 155 Proteus
☐ 156 Pseudomonas (all species except cepacia & maltophilia)
☐ 157 Pseudomonas or Burkholderia cepacia
☐ 158 Pseudomonas or Stenotrophomonas or Xanthomonas maltophilia
☐ 159 Rhodococcus
☐ 107 Rickettsia
☐ 160 Salmonella (all species)
☐ 161 Serratia marcescens
☐ 162 Shigella
☐ 163 Staphylococcus, coagulase negative (not aureus)
☐ 164 Staphylococcus aureus
☐ 165 Staphylococcus, NOS
☐ 166 Stomatococcus mucilaginosis
☐ 167 Streptococcus (all species except Enterococcus)
☐ 178 Streptococcus pneumoniae
☐ 168 Treponema (syphilis)
☐ 169 Vibrio (all species)
☐ 197 Multiple bacteria at a single site, specify bacterial codes
☐ 198 Other bacteria, specify ‡
☐ 501 Suspected atypical bacterial infection
☐ 502 Suspected bacterial infection
Fungal Infections
☐ 200 Candida, NOS
☐ 201 Candida albicans
☐ 206 Candida guillermondi
☐ 202 Candida krusei
☐ 207 Candida lusitaniae
☐ 203 Candida parapsilosis

CIBMTR Center Numl	ber: CIBMTR Recipient ID:
	☐ 204 Candida tropicalis ☐ 205 Candida (Torulopsis) glabrata
	☐ 209 Other Candida, specify ‡
	☐ 210 Aspergillus, NOS
	☐ 211 Aspergillus flavus
	☐ 212 Aspergillus fumigatus
	☐ 213 Aspergillus niger
	☐ 219 Other Aspergillus, specify ‡
	☐ 220 Cryptococcus species
	230 Fusarium species
	☐ 261 Histoplasmosis
	☐ 240 Zygomycetes, NOS
	☐ 241 Mucormycosis ☐ 242 Rhizopus
	☐ 250 Yeast, NOS
	☐ 259 Other fungus, specify ‡
	☐ 260 Pneumocystis (PCP / PJP)
	☐ 503 Suspected fungal infection
Product Infusion	
94. Date of this produ	act infusion://///
95. Was the entire vo	plume of received product infused?
☐ Yes ———	→
☐ No	96. Specify what happened to the reserved portion: □ Discarded
	☐ Cryopreserved for future use
	Other fate 97. Specify other fate:
98. Time product infu	sion initiated (24-hour clock): : : Standard time
99. Date infusion stop	oped:////
100. Time product infu	ision completed (24-hour clock): : :

O1. Specify the route of product infusion: ☐ Intravenous ☐ Intramedullary (Intraosseous)		
Other route of infusion ————	102. Specify other route	of infusion:
ne following questions are applicable to cond NMDP products continue with the signat		DP allogeneic products continue with question 144. Autologou
03. Were there any adverse events or incident	associated with the stem ce	Il infusion?
☐ Yes ———————————————————————————————————	Specify the following a	dverse event(s):
□ NO	104. Brachycardia ☐ Yes → ☐ No	In the Medical Director's judgment, was the adverse event a direct result of the infusion?
	106. Chest tightness / p	ain
	☐ Yes → 107	In the Medical Director's judgment, was the adverse event a direct result of the infusion? Yes No
	108. Chills at time of inf	usion
	☐ Yes → 109	In the Medical Director's judgment, was the adverse event a direct result of the infusion? Yes No
	110. Fever ≤ 103° F wit	nin 24 hours of infusion
	☐ Yes → 111.	In the Medical Director's judgment, was the adverse event a direct result of the infusion? ☐ Yes ☐ No
	112. Fever > 103° F wit	hin 24 hours of infusion
	☐ Yes → 113.	In the Medical Director's judgment, was the adverse event a direct result of the infusion?
		☐ Yes ☐ No
	114. Gross hemoglobin	uria
	☐ Yes → 115.	In the Medical Director's judgment, was the adverse event a direct result of the infusion?
		☐ Yes ☐ No

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	116. Headache ☐ Yes → ☐ No 1174. In the Medical Director's judgment, was the adverse event a direct result of the infusion? ☐ Yes ☐ No
	118. Hives ☐ Yes → ☐ No ☐ No ☐ In the Medical Director's judgment, was the adverse event a direct result of the infusion? ☐ Yes ☐ No
	120. Hypertension ☐ Yes → ☐ No ☐ No ☐ Yes ☐ No ☐ Yes ☐ No ☐ Yes ☐ No
	122. Hypotension ☐ Yes → ☐ No 123. In the Medical Director's judgment, was the adverse event a direct result of the infusion? ☐ Yes ☐ No
	124. Hypoxia requiring oxygen (O₂) support ☐ Yes → ☐ No ☐ No ☐ Yes ☐ No ☐ Yes ☐ No ☐ Yes ☐ No
	126. Nausea Yes No 127. In the Medical Director's judgment, was the adverse event a direct result of the infusion? Yes No
	128. Rigors, mild ☐ Yes → ☐ No ☐ No ☐ Yes ☐ No ☐ Yes ☐ No ☐ Yes ☐ No
	130. Rigors, severe ☐ Yes → ☐ No ☐ No ☐ Yes ☐ No ☐ Yes ☐ No ☐ Yes ☐ No

132. Shortness of breath (SOB)
☐ Yes → ☐ No ☐ No ☐ 133. In the Medical Director's judgment, was the adverse event a direct result of the infusion? ☐ Yes ☐ No
134. Tachycardia ☐ Yes → ☐ No 135. In the Medical Director's judgment, was the adverse event a direct result of the infusion?
☐ Yes ☐ No
136. Vomiting ☐ Yes → ☐ No ☐ No ☐ No ☐ Yes ☐ No ☐ Yes ☐ No ☐ Yes ☐ No
138. Other expected AE
☐ Yes → ☐ No 139. Specify other expected AE: ☐ 140. In the Medical Director's judgment, was the adverse event a direct result of the infusion? ☐ Yes ☐ No
141. Other unexpected AE ☐ Yes →
□ No 142. Specify other unexpected AE: 143. In the Medical Director's judgment, was the adverse event a direct result of the infusion? □ Yes □ No

CIBMTR Center Number:	CIBMTR Recipient ID:
Donor / Infant Demographic Information	
This Donor Demographic Information section (or product was from an NMDP donor or an autolog	juestions 144-170) is to be completed for all non-NMDP allogeneic donors. If the stem cell gous donor, continue with the signature lines.
144. Was the donor ever pregnant? Yes No Unknown Not applicable (male donor or cord blood unit)	145. Number of pregnancies ☐ Known → 146. Specify number of pregnancies:
147. Did this donor have a central line placed?	☐ Yes ☐ No ☐ Unknown
148. Ethnicity (donor) Hispanic or Latino	☐ Not Hispanic or Latino ☐ Not applicable (not a resident of the USA) ☐ Unknown
149. Race (donor) (check all that apply) White Black or African American Asian American Indian or Alaska Native Native Hawaiian or Other Pacific Islander Not reported Unknown Go to Question 151	450. Doos detail (dange) (elegals all that apply)
	150. Race detail (donor) (check all that apply) ☐ Eastern European
	☐ Mediterranean
	☐ Middle Eastern
	☐ North Coast of Africa
	☐ North American
	☐ Northern European
	☐ Western European ☐ White Caribbean
	☐ White South or Central American
	☐ Other White
	☐ African (both parents born in Africa)
	☐ African American
	☐ Black Caribbean ☐ Black South or Central American
	☐ Other Black
	☐ Alaskan Native or Aleut
	☐ North American Indian
	☐ American Indian, South or Central America
	☐ Caribbean Indian
	☐ South Asian
	☐ Filipino (Pilipino) ☐ Japanese

	CIBMTR Recipient ID:
	☐ Korean ☐ Chinese ☐ Vietnamese ☐ Other Southeast Asian ☐ Guamanian ☐ Hawaiian ☐ Samoan ☐ Other Pacific Islander ☐ Unknown
151. Was the donor a carrier for poter ☐ Yes	ntially transplantable genetic diseases?
□ No	152. Specify potentially transplantable genetic disease: Sickle cell anemia Thalassemia Other hemoglobinopathy
	Other disease — 153. Specify other disease:
	or other transferable genetic or clonal abnormalities?
☐ Yes - Go to question 155 ☐ No - If this is a related dono	or, go to question 160; all other donor types go to signature line d donor, go to question 160; all other donor types go to signature line
☐ Yes - Go to question 155 ☐ No - If this is a related dono	or, go to question 160; all other donor types go to signature line
☐ Yes - Go to question 155 ☐ No - If this is a related dono	or, go to question 160; all other donor types go to signature line d donor, go to question 160; all other donor types go to signature line 155. Clonal hematopoiesis of indeterminate potential (CHIP): Yes 156. What was the method of testing used? 157. Monoclonal B-cell lymphocytosis
☐ Yes - Go to question 155 ☐ No - If this is a related dono	or, go to question 160; all other donor types go to signature line d donor, go to question 160; all other donor types go to signature line 155. Clonal hematopoiesis of indeterminate potential (CHIP): Yes No 156. What was the method of testing used?
Yes - Go to question 155 No - If this is a related dono Unknown - If this is a related The following questions (160–167) a	or, go to question 160; all other donor types go to signature line d donor, go to question 160; all other donor types go to signature line 155. Clonal hematopoiesis of indeterminate potential (CHIP): Yes 156. What was the method of testing used? 157. Monoclonal B-cell lymphocytosis Yes No 158. Other transferable genetic or clonal abnormality Yes 159. Specify other transferable genetic or clonal abnormality:
☐ Yes - Go to question 155 ☐ No - If this is a related dono ☐ Unknown - If this is a related The following questions (160–167) a NMDP unrelated donor, NMDP dono 160. Was the donor hospitalized (inpar	or, go to question 160; all other donor types go to signature line 155. Clonal hematopoiesis of indeterminate potential (CHIP): Yes → 156. What was the method of testing used? No 157. Monoclonal B-cell lymphocytosis Yes No 158. Other transferable genetic or clonal abnormality Yes → No No 159. Specify other transferable genetic or clonal abnormality: Specify other transferable genetic or clonal abnormality: No No No No No No No N

BMTR Center Number:	CIBMTR Recipient ID:	
163. Did the allogeneic donor give one or mo	ore autologous transfusion units?	
Yes ———	164. Date of collection://///	
□ No		
	165. Number of units:	
66. Did the donor receive blood transfusion	is as a result of the collection?	
☐ Autologous transfusions ———	→	
	167. Specify number of autologous units:	
Allogenic transfusions ————————————————————————————————————		
□ No	168. Specify number of allogenic units:	
69. Did the donor die as a result of the colle	ection?	
☐ Yes ☐ No - Go to signature line	170. Specify cause of death:	
□ No - Go to signature line		
First Name (person completing form):		
ast Name:		
-mail address:		
Date:////		



Donor Sex: Male Female

Confirmation of HLA Typing

Registry Use Only Sequence Number: Date Received: Date Received: Date Received: Public Burden Statement: An agency may not conduct or sponsor, and a person is no required to respond to, a collection of information unless it displays a currently valid OMB control number. The OMB control number for this project is 0915-0310. Public reporting burden for this collection of information, in combination with the IDM Form 2004 and HCT Infusion Form 2006, is estimated to average 1.5 hours per response, including the time for reviewing instructions, searching existing data sources, and completing and reviewing the collection of information. Send comments regarding this burden estimate or any other aspect of this collection of information, including suggestions for reducing this burden, to HRSA Reports Clearance Officer, 5600 Fishers Lane Room 10-33, Rockville, Maryland, 20857. Expiration date: 1/31/2020
CIBMTR Center Number:
CIBMTR Research ID:
Event date:////
Product Identifiers:
NMDP cord blood unit ID:
Non-NMDP unrelated donor ID:
Non-NMDP cord blood unit ID:
GRID (optional):
ISBT DIN:
Registry or UCB Bank ID:
Donor DOB:////
Donor Age:

OMB No: 0915-0310 Expiration Date: 1/31/2020

CIBMTR Center Number:	CIBMTR Research	arch ID:	
Donor/Cord Blood Unit Identification			
This form must be completed for all non-NMDP allogeneic or syngeneic donors or recipients, or non-NMDP cord blood units. If the donor, recipient, or cord blood unit was secured through the NMDP, then report HLA typing on the appropriate NMDP forms.			
A separate copy of this form should be completed for each non-NMDP donor, recipient, or cord blood unit. Parental typing (maternal and paternal) should be submitted for all mismatched related donor transplants (CRF track only), if available. Cord blood maternal typing should be submitted for all unrelated cord blood transplants (CRF track only), if available.			
Specify the person for	1. Specify the person for whom this typing is being done:		
HLA Typing by DNA Tech	nology		
Was documentation s	ubmitted to the CIBMTR? (e.g. lab report)	☐ Yes ☐ No	
HLA Alleles Defined by Di typing or Sequence Base	NA Technology (e.g., Sequence Specific Oligonucleotide F d (SBT) typing.)	Probe (SSOP) typing, Sequence Specific Primer (SSP)	
	sed to type for a single allele, combinations of alleles (all ng result. For this reason, the number of digits, as well as		
Transcribe the informatio length which represent a	", " – " or a combination of numbers and letters on the ty n onto the form as directly as possible. The letters are cal combination of possible alleles at a locus. The same allel 2, DRB1*01:01/01:02, DRB1*01:01/02, or DRB1*01:AB).	lled allele codes, and will be 1 or more characters in	
	eported for each locus, unless the individual is presumed be the first allele designation in the first box, and the sec second box blank.		
Class I			
3. Locus A			
☐ Known — → ☐ Unknown	4. First A* allele designations		
	Second A* allele designations		
5. Locus B			
☐ Known ☐ Unknown	6. First B* allele designations		
	Second B* allele designations		
7. Locus C			
☐ Known ☐ Unknown	8. First C* allele designations		
	Second C* allele designations		

SIDIV	TIR Center Number:	CIBMTR Research ID:	
Class II			
9.	Locus DRB1 ☐ Known → ☐ Unknown	10. First DRB1* allele designations	
		Second DRB1* allele designations	
Cla	ss II (Optional)		
		al allele information if it is available from your laboratory	
11.	☐ Known →	12. First DRB3* allele designations	
		Second DRB3* allele designations	
13.	Locus DRB4		
	☐ Known → ►	14. First DRB4* allele designations	
		Second DRB4* allele designations	
15.	Locus DRB5		
	☐ Known ──►	16. First DRB5* allele designations	
		Second DRB5* allele designations	
17.	Locus DQB1		
	☐ Known ——→	18. First DQB1* allele designations	
		Second DQB1* allele designations	

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19. Locus DPB1 ☐ Known ——→		
Unknown	20. First DPB1* allele designations	
	Second DPB1* allele designations	
21. Locus DQA1		
☐ Known — → ☐ Unknown	22. First DQA1* allele designations	
	Second DQA1* allele designations	
23. Locus DPA1 ☐ Known ———		
Unknown	24. First DPA1* allele designations	
	Second DPA1* allele designations	
Antigens Defined by Sero	logic Typing	
Use the following lists wh	en reporting HLA-A and B antigens. Report broad antigens only when your laboratory was not able to confirm	
typing for a known split a		
Instructions for the use of	the "X" Antigen Specificity for Typing By Serology	
"unknown but known to be of assumed to be the same as specificity, the "X" or "blank"	ogically defined "X" antigen specificity: AX, BX, CX, DRX, DPX, and DQX. At this time an "X" specificity is defined as different from the other antigen at that locus." This is different from a blank specificity, which is defined as "unknown but the other antigen at that locus." When comparisons between recipient and donor antigens involve an "X" or "blank" is assumed to be homozygous for the antigen reported at the locus. In other words, the search algorithm treats typings tigens in the same manner as known homozygous typings.	
A Antigens		
25. Number of antigens provided:		
☐ One - Go to question 26, then continue with question 28 ☐ Two - Go to questions 26-27		
	26. Specificity – 1st antigen ☐ A1	
	☐ A203	
	☐ A210 ☐ A3	

CIBM I R Center Number:	CIBMTR Research ID:
	A9
	27. Specificity – 2nd antigen A1

CIBMTR Center Number:	CIBMTR Research ID:
	☐ A31(19)
	☐ A32(19)
	☐ A33(19)
	☐ A34(10)
	☐ A36
	☐ A43
	☐ A66(10)
	☐ A68(28)
	☐ A69(28)
	☐ A74(19)
	☐ A80 —
	□ AX
B Antigens	
28. Number of antigens provided:	
One - Go to question 29, then continue	with question 31
☐ Two - Go to questions 29-30	4
,	
	29. Specificity – 1st antigen
	☐ B5
	□ B7
	☐ B703
	□ B8
	☐ B12
	☐ B13
	☐ B14
	☐ B15
	☐ B16
	☐ B17
	☐ B18
	☐ B21
	☐ B22
	□ B27
	☐ B2708
	☐ B35
	☐ B37
	☐ B38(16)
	☐ B39(16)
	☐ B3901 —
	□ B3902
	☐ B40
	☐ B4005
	☐ B41
	☐ B42

CIBMTR Center Number:	CIBMTR Research ID:
	☐ B44(12)
	☐ B45(12)
	☐ B46
	☐ B47
	☐ B48
	☐ B49(21)
	☐ B50(21)
	☐ B51(5)
	☐ B5102
	☐ B5103
	☐ B52(5)
	☐ B53
	☐ B54(22)
	☐ B55(22)
	☐ B56(22)
	☐ B57(17)
	☐ B58(17)
	☐ B59
	☐ B60(40)
	☐ B61(40)
	☐ B62(15)
	☐ B63(15)
	☐ B64(14)
	☐ B65(14)
	☐ B67
	☐ B70
	☐ B71(70)
	☐ B72(70)
	☐ B73
	☐ B75(15)
	☐ B76(15)
	☐ B77(15)
	☐ B78
	☐ B81
	☐ B82
	□вх
	30. Specificity – 2nd antigen
	□ B5
	□ B7
	□ B703
	□ 88
	□ B12

CIBMTR Center Number:	CIBMTR Research ID:
	☐ B13
	☐ B14
	☐ B15
	☐ B16
	□ B17
	☐ B18
	☐ B21
	☐ B22 —
	□ B27
	☐ B2708
	☐ B35
	☐ B37
	☐ B38(16)
	☐ B39(16) ☐ B3901
	☐ B3901
	☐ B3902 ☐ B40
	☐ B4005
	☐ B41
	☐ B42
	☐ B44(12)
	☐ B45(12)
	□ B46
	☐ B47
	☐ B48
	☐ B49(21)
	☐ B50(21)
	☐ B51(5)
	☐ B5102
	☐ B5103
	☐ B52(5)
	□ B53
	☐ B54(22)
	☐ B55(22)
	☐ B56(22)
	☐ B57(17)
	☐ B58(17)
	☐ B59
	☐ B60(40)
	☐ B61(40)
	☐ B62(15)
	☐ B63(15)
	☐ B64(14) ☐ B65(14)
	LJ D03(14)

CIBMTR Center Number:	CIBMTR Research ID:
	□ B67 □ B70 □ B71(70) □ B72(70) □ B73 □ B75(15) □ B76(15) □ B77(15) □ B78 □ B81 □ B82 □ BX
Optional Antigen Reporting	
Antigens Defined by Serologic Typing	
C Antigens	
31. Number of antigens provided: ☐ One - Go to question 32, then continue ☐ Two - Go to questions 32-33	32. Specificity – 1st antigen Cw1 Cw2 Cw3 Cw4 Cw5
	□ Cw6 □ Cw7 □ Cw8 □ Cw9(w3) □ Cw10(w3) □ CX
	33. Specificity – 2nd antigen Cw1 Cw2 Cw3 Cw4 Cw5 Cw6

CIBINITR Center Number:	CIBINITR Research ID:		
	☐ Cw7 ☐ Cw8 ☐ Cw9(w3) ☐ Cw10(w3) ☐ CX		
Bw Specificity			
34. Specificity Bw4 present?35. Specificity Bw6 present?		☐ Yes	□ No
DR Antigens			
36. Number of antigens provided: ☐ One - Go to question 37, then continued ☐ Two - Go to questions 37-38	ue with question 39		
	37. Specificity – 1st antigen DR1 DR103 DR2 DR3 DR4 DR5 DR6 DR7 DR8 DR9 DR10 DR11(5) DR12(5) DR13(6) DR14(6) DR1403 DR1404 DR15(2) DR16(2) DR18(3) DRX		
	38. Specificity – 2nd antigen DR1 DR103 DR2		

CIBMTR Center Number:	CIBMTR Research ID:		
	□ DR3 □ DR4 □ DR5 □ DR6 □ DR7 □ DR8 □ DR9 □ DR10 □ DR11(5) □ DR12(5) □ DR13(6) □ DR14(6) □ DR1403 □ DR1404 □ DR15(2) □ DR15(2) □ DR16(2) □ DR17(3) □ DRX		
DR51 Antigen			
39. Specificity DR51 present?		☐ Yes	☐ No
DR52 Antigen			
40. Specificity DR52 present?		Yes	□ No
DR53 Antigen			
41. Specificity DR53 present?		Yes	□ No
DQ Antigens			
42. Number of antigens provided: ☐ One - Go to question 43, then continue ☐ Two - Go to questions 43-44	### ### ##############################		

CIBMTR Center Number:	CIBMTR Research ID:
	□ DQ9(3) □ DQX
	44. Specificity – 2nd antigen DQ1 DQ2 DQ3 DQ4 DQ5(1) DQ6(1) DQ7(3) DQ8(3) DQ8(3) DQ9(3) DQX
DP Antigens 45. Number of antigens provided: One - Go to question 46, then continue Two - Go to questions 46-47	ue with signature line
	46. Specificity – 1st antigen DPw1 DPw2 DPw3 DPw4 DPw5 DPw6 DPX
	47. Specificity – 2nd antigen DPw1 DPw2 DPw3 DPw4 DPw5 DPw6 DPX
First Name (person completing form):	
Last Name:	
E-mail address://	



Infectious Disease Markers

	Registry Use Only Sequence Number: Public Burden Statement: 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. The OMB control number for this project is 0915-0310. Public reporting burden for this collection of information, in combination with the HLA Typing Form 2005 and HCT Infusion Form 2006, is estimated to average 1.5 hours per response, including the time for reviewing instructions, searching existing data sources, and completing and reviewing the collection of information. Send comments regarding this burden estimate or any other aspect of this collection of information, including suggestions for reducing this burden, to HRSA Reports Clearance Officer, 5600 Fishers Lane, Room 10-33, Rockville, Maryland, 20857. Expiration date: 1/31/2020
	CIBMTR Center Number:
	CIBMTR Research ID:
	Event date: /
	HCT type (check all that apply): ☐ Allogeneic, unrelated ☐ Allogeneic, related
	Product type (check all that apply): Bone marrow PBSC Single cord blood unit Multiple cord blood units Other product. Specify:
	Product Identifiers:
	NMDP cord blood unit ID:
	Non-NMDP unrelated donor ID:
	Non-NMDP cord blood unit ID:
	GRID (optional):
	ISBT DIN:
	Registry or UCB Bank ID:
	Donor DOB: / / / / /
	Donor Age:
	Donor Sex: Male Female
l	

er Number:	CIBMTR Recipient ID:			
st be completed for all non-NI	MDP allogeneic or syngeneic donors, or non-NMDP cord blood units.			
Donor/Cord Blood Unit Identification				
eing tested for IDMs? r IDM (bone marrow or PBSC) rnal IDM (cord blood) blood unit IDM				
ease Marker (report final test re	esults)			
irus (HBV)				
hepatitus B surface antigen) tive eactive one	3. Date sample collected://////			
tive eactive one				
Nicensed NAAT testing for HBV ve tive one	/ performed? 7. Date sample collected:////			
rus (HCV) (: (hepatitis C antibody)				
eactive ————————————————————————————————————	9. Date sample collected://////			
A licensed NAAT testing for HCV	/ performed?			
one	11. Date sample collected:/ / / DD			
nodeficiency Virus (HIV) 4 antigen: tive eactive eactive eported	13. Date sample collected: / / /			
node 4 an tive eact	tigen: tive			

ואוסוי	TR Center Number:		CIBM I R Recipient ID:
14. Was FDA licensed NAAT testing for HIV-1 performed?			
	□ No □ Not done	15.	Date sample collected:/ / / / DD
16.	Anti-HIV 1 and anti-HIV 2*: (antibodies to Hur * Testing for both HIV antibodies is required.	This tes	sting may be performed as separate tests or done using a combined assay.
	Non-reactive →Not doneNot reported	17.	Date sample collected:/ / / /
Cha			
18.	Chagas testing ☐ Positive → ☐ Negative → ☐	19.	Date sample collected://///
	☐ Not done		
Herp	pes simplex virus (HSV)		
20.	Anti-HSV (Herpes simplex virus antibody)		
	Positive —	21.	Date sample collected: //
	☐ Negative → ► ☐ Not done		YYYY MM DD
Eps	tein-Barr virus (EBV)		
22.	Anti-EBV (Epstein-Barr virus antibody)		
	Positive Negative	23.	Date sample collected: / / /
	☐ Inconclusive →		TTTT IVIIVI DD
	☐ Not done		
Vari	cella zoster virus (VZV)		
24.	Anti-VZV (Varicella zoster virus antibody)		
	Positive	25.	Date sample collected: //
	☐ Negative ☐ Not done		YYYY MM DD

CIBMTR Center Number:	CIBMTR Recipient ID:
Other Infectious Disease Marker	
26. Other infectious disease marker, specify: ☐ Yes →	27 Date sample collected:
□ No	27. Date sample collected://////
	28. Specify test and method:
	29. Specify test results:
	Copy questions 27 - 29 to report multiple other infectious disease markers
First Name:	
Last Name:	
E-mail address:	
Date:///	
TTT WIN DD	