OMB No: 0915-Expiration Date:

Public Burden Statement

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Hematopoietic Stem Cell Transplant (HSCT) Infusion

	Registry Use Only
Sequence Number:	
Date Received:	

CIBMTR Center Number:
CIBMTR Recipient ID:
NMDP Donor ID:
NMDP Cord Blood Unit ID: Non-NMDP cord blood unit *
* For non-NMDP cords, see page 11 to report CBU ID and donor demographics.
Today's Date: 20
Date of HSCT for which this form is being completed: Month Day Year
HSCT type: ☐ autologous ☐ allogeneic, ☐ allogeneic, ☐ syngeneic (check only one) unrelated related (identical twin)
Product type: ☐ marrow ☐ PBSC ☐ cord blood ☐ other product, (check only one) ☐ multiple cord specify: blood units infused

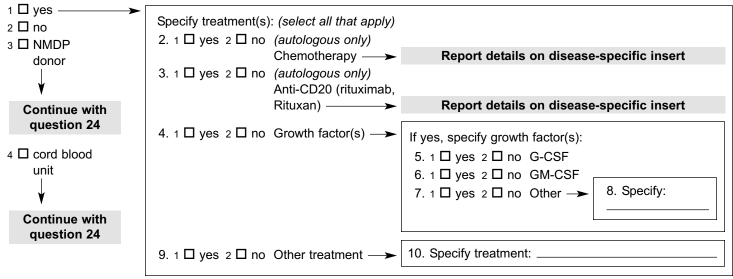
This form must be completed for all recipients who receive a HSCT product. If more than one type of HSCT product is infused, each product type must be analyzed and reported separately.

Questions followed by the symbol \square indicate additional information necessary to complete the question is referenced in the forms instruction manual; $\square A$ indicates an appendix.

A series of collections should be considered a <u>single product</u> when they are all from the same donor and use the same collection method and technique (and mobilization, if applicable), even if the collections are performed on different days.

Pre-Collection Therapy

1. Did the donor receive treatment, prior to any stem cell harvest, to enhance the product collection for this HSCT? (If the HSCT product was from an NMDP donor, or the product is a cord blood unit, then continue with question 20.)



CIBMTR Center Number:	CIBMTR Recipient ID:
Product Collection	
11. Date of product collection	Month Day Year
12. Was more than one colle	ection required for this HSCT? 🕮
1 □ yes ————— 2 □ no	13. Specify the number of subsequent days of collection in this episode:
	Complete a separate product form for each subsequent collection that was not part of this mobilization.
	ded to the product during collection?
1 □ yes — → 2 □ no	Specify anticoagulant(s):
2 ப 110	15. Acid citrate dextrose (ACD)
	1 □ yes
	2 □ no
	16. Citrate phosphate dextrose (CPD)
	1 □ yes
	2 □ no
	17. Heparin
	1 ☐ yes
	2 □ no
	18. Other anticoagulant
	1 ☐ yes — → 19. Specify other anticoagulant:
	2 □ no
Product Transport and	d Receipt
20. Was this product collected	ed off-site and shipped to your facility?
1 ☐ yes ———	
2 □ no	21. Date of receipt of product at your facility:
	Month Day Year
	22. Time of receipt of product (24-hour clock): Hour 1 □ standard time 2 □ daylight savings time
	23. Specify the shipping environment of the product(s):
	1 ☐ frozen gel pack
	2 ☐ frozen cord blood unit(s)
	3 ☐ room temperature per transplant center request
	4 □ other temperature ➤ 24. Specify shipping environment:
	temperature ➤ 24. Specify shipping environment:
	25. (Cord blood product only) Were the secondary containers (e.g., insulated shipping containers and unit cassette) intact when they arrived at your center?
	1 □ yes
	2 □ no

IBMIR Center Number:	CIBINITR RECIPIENT ID:
_	
	26. (Cord blood product only) Was the cord blood unit completely frozen when it arrived at your center?
	1 □ yes 2 □ no
	27. (Cord blood product only) Was the cord blood unit stored at your center prior to thawing?
	1 ☐ yes 2 ☐ no 28. Specify the storage method used for the cord blood unit: 1 ☐ liquid nitrogen 2 ☐ vapor phase 3 ☐ electric freezer
	29. Temperature during storage: — ° C
	30. Date storage started: Month Day Year
roduct Processing	Manipulation
· · · · · · · · · · · · · · · · · · ·	ceived, then cryopreserved at your facility prior to infusion?
1 □ yes 2 □ no	
3 ☐ not applicable, core	
32. Was the product thawe 1 □ yes ————	d from a cryopreserved state prior to infusion?
2 □ no	33. Was the entire product thawed? 1 □ yes
	2 ☐ no — → 34. Was a compartment of the bag thawed?
	1 □ yes 2 □ no
	35. Were there multiple product bags?
	1 ☐ yes — → 36. Specify number of bags thawed:
	2 🗆 110
	37. Date thawing process initiated: Day Year Page Page
	38. Time at initiation of thaw (24-hour clock): ☐ Image: Standard time continue co
	39. Time at completion of thaw (24-hour clock): ☐ Image: Line of the standard time and time
	40. Was the primary container (e.g., cord blood unit bag) intact upon thawing? 1 □ yes 2 □ no
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IBMTR Center Number:	CIBMTR Recipient ID:
	 41. What method was used to thaw the product? 1 □ no wash — thawed at bedside, then infused 2 □ DMSO dilution — thawed in lab (added dextran and albumin), then infused 3 □ washed — thawed in lab (added dextran and albumin), spun and reconsituted in dextran albumin, then infused 4 □ other
	method — > 42. Specify other thaw method:
	43. Did any adverse events or incidents occur while thawing the product? 1 □ yes 2 □ no
14. Was the product manipu	lated prior to infusion?
1 □ yes — ➤ 2 □ no	45. Specify portion manipulated: 1 □ entire product 2 □ portion of product
If autologous product, continue with question 92;	Specify all methods used to manipulate the product: 46. ABO incompatibility (RBC depletion)
if allogeneic product, continue with question 141.	Specify method: 47. 1 yes 2 no Buffy coat preparation 48. 1 yes 2 no Cell separator (i.e., COBE Spectra) 49. 1 yes 2 no Density gradient separation (i.e., Ficoll) 50. 1 yes 2 no Plasma removal 51. 1 yes 2 no Sedimentation (i.e., hetastarch) 52. 1 yes 2 no Other 53. Specify:
	54. Ex-vivo expansion 1 yes 2 no 55. Genetic manipulation (gene transfer / transduction) 1 yes 2 no 56. Volume reduction 1 yes 2 no 57. CD34+ selection
	1 ☐ yes 2 ☐ no 58. Specify cell selection system used: 1 ☐ CliniMACS / CliniMax 2 ☐ Isolex
	3 ☐ other — → 59. Specify system:
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CIBMTR Center Number:																							(С	IE	В	٨	/1	ΤI	₹	F	₹6	е	С	ip	oie	er	nt	IE) :													

CO T call depletion		
60. T-cell depletion	Specify method:	
1 □ yes		o antibody affinity column ——>
2 🗖 110	•	and the decrease and relative
	•	Kepoit the
	05. 1 L yes 2 L II	antibody coated plates antibodies and soybean lectin antibodies used for T-cell
	64 1 □ yes 2 □ n	antibody + complement — depletion at
	65. 1 □ yes 2 □ n	o antibody + toxin — page question 73.
	66. 1 □ yes 2 □ n	immunomagnetic beads — >
	67. 1 □ yes 2 □ n	o elutriation
	68. 1 □ yes 2 □ n	CD34 affinity column plus sheep red blood cell
		rosetting 🕮
71. Other	69. 1 □ yes 2 □ n	o other — > 70. Specify:
manipulation		
1 □ yes ——➤	72. Specify:	
2 □ no		
73. Were antibodies us	ed during product ma	nipulation?
1 □ yes —— >	Charify antibadian	
2 🗖 no	Specify antibodies:	
	74. 1 🗆 yes 2 🗆 n	
	75. 1 □ yes 2 □ n 76. 1 □ yes 2 □ n	
	77. 1 🗆 yes 2 🗆 n	
	77. 1	
	79. 1 🗆 yes 2 🗆 n	
	80. 1 yes 2 n	
	81. 1 yes 2 n	
	· •	o anti TCR alpha / beta (T10-B9)
	83. 1 🗆 yes 2 🗆 n	. , ,
	· •	o other CD3 —> 85. Specify:
	04. 1 L yc3 2 L 11	outlet obe 2 ce. epecity.
	86. 1 □ yes 2 □ n	o anti CD52 — Specify antibodies:
		yes no
		87. 1
		88. 1 □ 2 □ campath-1G 89. 1 □ 2 □ campath-1H
	90. 1 □ yes 2 □ n	
	00. 1 L yC3 2 L II	antibody — 91. Specify:
		, 5 5555,

CIBMTR Center Number: CIBMTR Recipient ID:												
--	--	--	--	--	--	--	--	--	--	--	--	--

Autologous Products Only

The following section refers to autologous products only, including autologous cord blood; if this is not an autologous HSCT, continue with the Product Analysis section at question 141.

92. Were tumor cells detected in the recipient or autologous product prior to HSCT?

□ no	Specify tumor cell detec	tion method used, and s	ite(s) of tumor ce	ells:
	93. Routine			
	histopathology	0 11 11 ()		
	1 □ yes ———	Specify site(s):		0: 1:: 11 1 1
	2 🗖 no			Circulating blood cells
		95. 1 Ll yes 2 Ll no	3 □ not tested	Bone marrow, in the interval between last systemic therapy
	97. Polymerase chain reaction	96. 1 □ yes 2 □ no	3 ☐ not tested	and collection Collected cells, before purging
	(PCR)	Chasify sits(s):		
	1 □ yes	Specify site(s):	o □ not tooted	Circulating blood calls
	2 🗖 no			Circulating blood cells Bone marrow, in the interval
		99. 1 Li yes 2 Li 110	3 Li not tested	between last systemic therapy and collection
	101. Other molecular	100. 1 □ yes 2 □ no	3 ☐ not tested	Collected cells, before purging
	technique	•		
	1 □ yes	102. Specify method: _		
	2 🗖 no	Specify site(s):		
		103. 1 □ ves 2 □ no	3 ☐ not tested	Circulating blood cells
		•		Bone marrow, in the interval
		,		between last systemic therapy and collection
	106. Immunohisto- chemistry	105. 1 ☐ yes 2 ☐ no	3 ☐ not tested	Collected cells, before purging
	1 □ yes ——➤	Specify site(s):		
	2 🗆 no	1 ' ' ' '	3 ☐ not tested	Circulating blood cells
		-		Bone marrow, in the interval
				between last systemic therapy and collection
	110. Cell culture	109. 1 □ yes 2 □ no	3 ☐ not tested	Collected cells, before purging
	technique	Chaoif (aita/a)		
	1 □ yes ———> 2 □ no	Specify site(s):	3 Π not tested	Circulating blood cells
	2 110	-		Bone marrow, in the interval
		112. 1 Li yes 2 Li 110	3 Li not tested	between last systemic therapy and collection
		113. 1 □ yes 2 □ no	3 ☐ not tested	Collected cells, before purging
	114. Other technique			
	1 □ yes ———	115. Specify method: _		
	2 □ no	Specify site(s):		
		116. 1 □ yes 2 □ no	3 ☐ not tested	Circulating blood cells
		· ·		Bone marrow, in the interval between last systemic therapy
				and collection

CIBMTR Center Number:		CIBMTR Re	cipient ID:											
					•	•	•		•					
119. Was the product treated t	to remove malignant ce	lls (purged)? <i>(autologous p</i>	oroduct only	/)										
1 □ yes	Specify method(s) used	<u>d</u> :												
2 LI 110	120. 1 ☐ yes 2 ☐ no	Monoclonal antibody ->	121. If ye	s, sp	ecify:						_			
		4-hydroperoxycyclophosp	hamide (4I	HC)										
	123. 1 ☐ yes 2 ☐ no 124. 1 ☐ yes 2 ☐ no	Other drug —	125. If ye	s, sp	ecify:									
	126. 1 ☐ yes 2 ☐ no	Elutriation												
	127 1 ☐ yes 2 ☐ no 128. 1 ☐ yes 2 ☐ no	Immunomagnetic column Toxin		c cn	ocify:									
	•		s, sp	ecity.						_				
130. 1 yes 2 no Positive stem cell selection (other than preparation of mononuclear														
	132. 1 ☐ yes 2 ☐ no	Other method —	133. If ye	s, sp	ecify:						_			
	Specify if tumor cells w	ere detected in the graft at	fter nurging	by c	ach n	netho	ל ווכם	q.						
	•	3 ☐ not tested Routine h		-	aonn	ictiio	u usc	u.						
	•	3 ☐ not tested Polymera	-	٠.	on (PC	CR)								
		3 ☐ not tested Other mo			ıe									
	-	3 ☐ not tested Immunoh		-										
	-	3 ☐ not tested Cell cultu	-	ie										
	139. 1 ☐ yes 2 ☐ no	3 ☐ not tested Other ➤	140. If ye	s, sp	ecify:						_			
L														
Product Analysis (All I	Products) 🕮													
Report product analysis reperformed, copy and comp	· ·	·	rformed. If	moı	re tha	n two	ana	lyses w	/ere					
		•			_									
	Product Analysis at 1s	st Timepoint	Produ	ct A	nalys	is at 2	2nd T	imepoi	nt					
Specify the timepoint in the product preparation phase that the product was analyzed:	1 ☐ product arrival 2 ☐ post-processing, pre-cryopreserval manipulation ☐ 3 ☐ post-thaw 4 ☐ post-manipulation 5 ☐ at infusion (final of	1	3 🗆 4 🗖	post- pre-o man post- post-	-proce cryopr ipulati -thaw -mani	essing eserv on 🕮	ation	/ ntity infu	used)					
Date of product analysis: 142.	Month Day	2 0 Year	163. Mor	th	Di	ay	2	O Year						
Total volume of product: 143.	•	1 🗆 mL 2 🗆 g	164.			•		1 □ m 2 □ g						

CIBMTR Center Number:					
-----------------------	--	--	--	--	--

CIBMTR Recipient ID:										
----------------------	--	--	--	--	--	--	--	--	--	--

Product Analysis at 1st Timepoint

Product Analysis at 2nd Timepoint

In this section, report the total	number of cells (ne	ot cells per kilogram).
	Total Number	Exponent

		Total Number	Exponent	Total Numb	er Exponent
Nucleated cells:	144.	•	x 10 not tested	165.	x 10 not tested
Mononucleated cells:	145.	•	x 10 not tested	166.	x 10 not tested
Nucleated red blood cells:	146.	•	x 10 not tested	167.	x 10 not tested
CD34+ cells:	147.	•	x 10 not tested	168.	x 10 not tested
CD3+ cells:	148.	•	x 10 not tested	169.	x 10 not tested
CD4+ cells:	149.	•	x 10 not tested	170.	x 10 not tested
CD8+ cells:	150.	•	x 10 not tested	171.	x 10 not tested
Viability of cells:	151.	%	☐ not tested	172.	% □ not tested
Method of testing cell viability:	152.	1 ☐ 7-AAD 2 ☐ propidium iodid 3 ☐ trypan blue 4 ☐ other method	е	173. 1 ☐ 7-AAD 2 ☐ propidium i 3 ☐ trypan blue 4 ☐ other meth)
Specify other method:	153.			174	
Were the colony-forming units (CFU) assessed after thawing? (cord blood product only)	154.		inue with question 155 inue with question 158		Continue with question 176 Continue with question 179
Was there growth?	155.	1 □ yes 2 □ no		176. 1 □ yes 2 □ r	no
Total colonies per product:	156.	•	x 10 ⁵ □ unknown	177.	• x 10 ⁵ unknown
Total CFU-GM:	157.	•	x 10 ⁵ ☐ unknown	178.	• x 10 ⁵ unknown
Were cultures performed before infusion to test the product(s) for bacterial or fungal infection? (complete for all cell products)	.			•	Continue with question 180 Continue with question 183
Specify results:	159.	1 ☐ positive 2 ☐ ne	egative ₃ □ unknown	180. 1 □ positive 2	☐ negative 3 ☐ unknown
Specify organism code(s): (see page 9 for codes)	160.			181.	
(300 page 0 101 00000)					
If code 198, 209, 219, or 259, specify organism:	161.			182	

CIBMTR Center Number:			CIBMTR Recipient ID					

Codes for Commonly Reported Organisms Bacterial Infections 139 Fusobacterium 155 Proteus **Fungal Infections** 144 Haemophilus (all species, 156 Pseudomonas (all species 121 Acinetobacter 200 Candida, NOS including influenzae) except cepacia & 122 Actinomyces 201 Candida albicans 145 Helicobacter pylori maltophilia) 206 Candida guillermondi 123 Bacillus 146 Klebsiella 157 Pseudomonas or 124 Bacteroides (gracillis, 202 Candida krusei 147 Lactobacillus (bulgaricus, Burkholderia cepacia uniformis, vulgaris, other 207 Candida Iusitaniae acidophilus, other species) 158 Pseudomonas or species) 203 Candida parapsilosis 102 Legionella Stenotrophomonas or 125 Bordetella pertussis 204 Candida tropicalis 103 Leptospira Xanthomonas maltophilia 205 Candida (Torulopsis) (whooping cough) 148 Leptotrichia buccalis 159 Rhodococcus glabrata 126 Borrelia (Lyme disease) 149 Leuconostoc (all species) 107 Rickettsia 127 Branhamella or Moraxella 209 Other Candida, specify ‡ 104 Listeria 160 Salmonella (all species) catarrhalis (other species) 210 Aspergillus, NOS 150 Methylobacterium 128 Campylobacter (all species) 161 Serratia marcescens 211 Aspergillus flavus 151 Micrococcus, NOS 162 Shigella 129 Capnocytophaga 212 Aspergillus fumigatus 112 Mycobacterium avium-163 Staphylococcus, coagulase 213 Aspergillus niger 171 Chlamydia pneumoniae intracellulare (MAC, MAI) negative (not aureus) 219 Other Aspergillus, specify ‡ 172 Other chlamydia, specify 174 Mycobacterium species 164 Staphylococcus aureus 220 Cryptococcus species 113 Chlamydia, NOS (cheloneae, fortuitum, 165 Staphylococcus, NOS 130 Citrobacter (freundii, other 230 Fusarium species haemophilum, kansasii, 166 Stomatococcus 261 Histoplasmosis species) mucogenicum mucilaginosis 131 Clostridium (all species 240 Zygomycetes, NOS 110 Mycobacterium tuberculosis 167 Streptococcus (all species 241 Mucormycosis except difficile) (tuberculosis, Koch bacillus) except Enterococcus) 132 Clostridium difficile 242 Rhizopus 175 Other mycobacterium, 178 Streptococcus pneumoniae 173 Corynebacterium jeikeium 250 Yeast, NOS specify 168 Treponema (syphilis) 133 Corynebacterium (all non-259 Other fungus, specify ‡ 176 Mycobacterium, NOS 169 Vibrio (all species) diptheria species) 260 Pneumocystis (PCP / PJP) 105 Mycoplasma 197 Multiple bacteria at a single 503 Suspected fungal infection 101 Coxiella 152 Neisseria (gonorrhoea, site, specify bacterial codes 134 Enterobacter meningitidis, other species) 198 Other bacteria, specify ± 177 Enterococcus, vancomycin 501 Suspected atypical bacterial 106 Nocardia resistant (VRE) 153 Pasteurella multocida infection 135 Enterococcus (all species) 154 Propionibacterium (acnes, 502 Suspected bacterial 136 Escherichia (also E. coli)

‡ The codes for "other organism, specify" (codes 198, 209, 219 and 259) should rarely be needed; check with your microbiology lab or HSCT physician before using them.

infection

1 ☐ standard time

2 ☐ daylight savings time

avidum, granulosum, other

species)

Product Infusion

138 Flavobacterium

137 Flavimonas oryzihabitans

183. Was more than one product infused? (e.g., marrow and PBSC, PBSC and cord blood, two different cords, etc.) 1 □ ves 184. Was the product infusion described on this insert intended to produce hematopoietic 2 🗖 no engraftment? 1 ☐ yes 2 🗆 no -185. Date of this product infusion: 2 0 Month Day Year 1 ☐ standard time 186. Time product infusion initiated (24-hour clock): 2 ☐ daylight savings time Minute Hour

Minute

Hour

187. Time product infusion completed (24-hour clock):

CIBMTR Center Number:		CIBMTR Recipier	nt ID:							
188. Total volume of product	plus additives infused:	• mL								
189. Specify the route of prod 1 ☐ intravenous 2 ☐ intramedullary 3 ☐ intraperitoneal 4 ☐ other route of		nfusion:								
infusion —										_
191. Did the volume of infuse 1 □ yes — ➤	ed product include any a	dded agents?								
2 □ no	Specify agent(s) added 192. 1 ☐ yes 2 ☐ no 193. 1 ☐ yes 2 ☐ no 194. 1 ☐ yes 2 ☐ no 195. 1 ☐ yes 2 ☐ no 196. 1 ☐ yes 2 ☐ no 197. 1 ☐ yes 2 ☐ no	ACD Albumin Antibiotic Dextran Heparin	agent:							
199. Was the entire volume o	of product infused?									
1 □ yes 2 □ no ————	200. Specify what happed 1 ☐ discarded 2 ☐ cryopreserved 3 ☐ other fate →									
		oducts except for autologous logous PBSC product, contin					SC prod	lucts) .	
202. Were there any adverse	events or incidents ass	ociated with the stem cell infusion	on?							
1 □ yes	Specify the following a 203. 1 □ yes 2 □ no 206. 1 □ yes 2 □ no	Adverse Event Brachycardia 2	-		n? □ no		Reso . 1 □ ye . 1 □ ye		□n	
	209. 1 yes 2 no 212. 1 yes 2 no	Chills at time of infusion 2 Fever $\leq 103^{\circ}$ F within 24 hours of infusion 2	210. 1 🗆 y	es 2	□ no	211	. 1 🗆 ye	es 2	□ n	0
			216. 1 🗆 y				. 1 □ y€			
	221. 1 yes 2 no 221. 1 yes 2 no 224. 1 yes 2 no 227. 1 yes 2 no 230. 1 yes 2 no 233. 1 yes 2 no	Headache 2 Hives 2 Hypertension 2	219. 1	/es 2 /es 2 /es 2	□ no □ no □ no	223 226 229	. 1 □ ye . 1 □ ye . 1 □ ye . 1 □ ye . 1 □ ye	es 2 es 2	□ n	0
	236. 1 yes 2 no 239. 1 yes 2 no 239. 1 yes 2 no 242. 1 yes 2 no	(O_2) support 2 Nausea 2 Rigors, mild 2	234. 1 🗆 y 237. 1 🗆 y 240. 1 🗀 y 243. 1 🗆 y	/es 2 /es 2	□ no □ no	238 241	. 1 □ ye . 1 □ ye . 1 □ ye . 1 □ ye	es 2	□ n	0

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CIBMTR Center Number:		CIBMTR Recip	pient ID:	
			Required Medical	
	_	Adverse Event	Intervention?	Resolved?
	•	Shortness of breath (SOB)	·	247. 1 ☐ yes 2 ☐ no
	·	•	249. 1 ☐ yes 2 ☐ no	250. 1 ☐ yes 2 ☐ no
	251. 1 ☐ yes 2 ☐ no		252. 1 □ yes 2 □ no	253. 1 ☐ yes 2 ☐ no
	254. 1 ☐ yes 2 ☐ no	Other expected AE 255. Specify:		
			256. 1 ☐ yes 2 ☐ no	257. 1 ☐ yes 2 ☐ no
	258. 1 ☐ yes 2 ☐ no	Other unexpected AE 259. Specify:		
			260. 1 ☐ yes 2 ☐ no	261. 1 ☐ yes 2 ☐ no
	262 In the Medical Dire	ector's judgement, was the a	dverse event a direct res	ult of the infusion?
	1 □ yes	Total o juagomoni, wao ino as	avoido ovoite a amode roc	ant of the initiation.
	2 □ no — →	263. Specify the most likel	y cause of the adverse e	event:
		1 ☐ regimen related		
		2 ☐ product reaction		
		3 ☐ drug reaction		
		4 □ other illness →	264. Specify illness:	
		5 □ other reason →	265. Specify reason: _	
l				
Donor Demographic I	nformation			
NMDP donors, NMDP cord	blood units, and autol	restions 266–287) is to be o ogous marrow or PBSC do donor, continue with quest	onors. If the stem cell p	
Times delici or all autorog	,000	aonon, commus mini quest		
266. Donor's date of birth:	Month Day	Year	date unknown	
267. (Cord blood unit only) Ag	•		age unknown	
260 (Cond blood unit only) No	on NMDD gord blood uni	t identification		
268. (Cord blood unit only) No number (CBU ID):	on-Nivide cord blood uni	it identification		
269. (Cord blood unit only) Is	the CBU ID number also	the ICCBBA ISBT 128 num	ber?	
1				
270. (Cord blood unit only) Na	ame of cord blood bank	providing CBU:		
271. Donor's gender:				
1 □ male				
2 ☐ female ———	272. Was the donor eve	er pregnant?		
	1 □ yes			
	2 □ no 3 □ unknown	273. Specify number of pre	egnancies:	unknown
	4 ☐ not applicable			
	cord blood uni			

CIBMTR Center Number:		CIBMTR Recipient ID:			
274. Donor's blood type and 1 □ A positive 2 □ A negative 3 □ B positive 4 □ B negative 5 □ AB positive 6 □ AB negative 7 □ O positive 8 □ O negative 9 □ unknown	Rh factor:				
275. Did this donor have a ce 1 □ yes 2 □ no 3 □ not applicable, cord blood unit or marrow product	276. Specify the site of the cer 1 ☐ femoral 2 ☐ subclavian 3 ☐ internal jugular	ntral line placement: Specify site:			
278. Donor's ethnicity: 1 ☐ Hispanic or Latino 2 ☐ not Hispanic nor Lat 3 ☐ unknown 279. Donor's race: (Mark the	tino group(s) in which the donor is a	a member. Check all that apply	⁄.)		
White 1 ☐ Eastern European 2 ☐ Mediterranean 3 ☐ Middle Eastern 4 ☐ North Coast of Afric 5 ☐ North American	Black or African American 11	ean 17 American Ind South or Cen America 18 Caribbean Ind	lian, ntral	Native Hawaiian or Other Pacific Islander 26 ☐ Guamanian 27 ☐ Hawaiian 28 ☐ Samoan 29 ☐ Other Pacific Islan	
6 Northern European 7 Western European 8 White Caribbean 9 White South or Central American 10 Other White	American Indian or Alaska Native 15 Alaskan Native of Aleut 16 North American Indian	20 🗆 Filipino (Pilipi 21 🗖 Japanese 22 🗖 Korean	ŕ	Decline 30 □ Donor declines to provide race 31 □ Donor's race unknown	
280. What is the relationship 1 □ sibling 2 □ recipient's child 3 □ other relative → 4 □ unrelated	of the donor to the recipient? 281. Specify the relationship of the parent the local pare				

CIBMTR Center Number:			CIBMTR Recip	oient IE	D:							
283. Was the donor / product	83. Was the donor / product tested for potentially transplantable genetic diseases?											
1 □ yes — ➤ 2 □ no 3 □ unknown	Specify disease(s) teste 284. 1 ☐ yes 2 ☐ no 3 285. 1 ☐ yes 2 ☐ no 3 286. 1 ☐ yes 2 ☐ no 3	Sickle cell an Thalassemia	emia 287. Specify g	genetic	dise	ase: _						
The following questions 2 autologous donor or NMD								odu	ct was f	rom	an	
288. Was the donor hospitaliz 1 □ yes 2 □ no	zed (inpatient) during or a	fter the colle	ction?									
89. Did the donor experience any life-threatening complications during or after the collection? 1 □ yes → 200. Specify complications:												
290. Specify complications:												
291. Did the donor recieve ble 1 □ yes ———————————————————————————————————	292. Was the blood tran 1 □ yes 2 □ no 294. Was the blood tran 1 □ yes 2 □ no	sfusion production sfusion production	uct autologous? y number of un	hits:	ogou	s)?						
296. Did the donor die as a re 1 □ yes ───────────────────────────────────	esult of the collection? 297. Specify cause of de	eath:										
298. (Related donors only) Di	id the recipient submit a r	esearch sam	ple?									
1 □ yes	299. Research sample r	ecipient ID:										
300. (Related donors only) Di 1 □ yes ———————————————————————————————————	id the donor submit a research sample of	-	9?									
302. Signed:		Person com	apleting form									
Please print name:												
Phone number: ()											
Fax number: ()											
E-mail address:												



Confirmation of HLA Typing

Non-NMDP Donor ID: * Non-NMDP Cord Blood Unit ID: * **Registry Use Only** * See page 5 for additional ID boxes or donor date of birth and gender. 2 0 Sequence Today's Date: Number: Year Date of HSCT for which this form is being completed: 🕮 Month Day Year Date Received: HSCT type: ☐ allogeneic, ☐ allogeneic, ☐ syngeneic (identical twin) unrelated related Product type: ☐ marrow ☐ PBSC □ cord blood □ other product, specify: . 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. 1. Please specify the person for whom this typing is being done: 1 ☐ recipient — final typing

CIBMTR Center Number:

CIBMTR Recipient ID:

3	ш	recipient's father — confirmatory typing –	-
4		donor — confirmatory typing	

2 ☐ recipient's mother — confirmatory typing →

- 5 ☐ cord blood unit confirmatory typing
- 6 ☐ maternal HLA typing

7 □ other	>
7 - 001101	

3. Was the recipient's father used as the donor?

2. Was the recipient's mother used as the donor?

1 □ yes

1 □ yes 2 🗖 no

2 🗖 no

4. Specify person and typing: _

HLA Typing by DNA Technology

Six sets of boxes are provided for reporting several possible alleles for each allele at a locus. If the laboratory reports more than six possible alleles, report six of the alleles in the boxes provided and write the remainder of the alleles in the space above or below the boxes for that locus.

5. Is a copy of the lab	report	attached ?
-------------------------	--------	------------

- 1 □ yes
- 2 🗖 no

CIBMTR C	enter Number:		CIBMTR Recipient ID:					
Class I								
Class I	No. of		Allele Designations					
Locus	Alleles Provided							
6. A	1 □ one	First A*						
	2 □ two							
		0 14*						
	3 ☐ not tested	Second A*						
7. B	1 □ one	First B*						
7. 0		1 1100 B						
	2 ☐ two							
	3 ☐ not tested	Second B*						
8. C	1 □ one	First C*						
	2 □ two							
	3 ☐ not tested	Second C*						
Class II		l						
0.000	No. of		Allele Designations					
Locus	Alleles Provided							
9. DRB1	1 □ one	First DRB1*						
	2 □ two							
	2 🗖 two							
	3 ☐ not tested	Second DRB1*						
Class II (C		allele information if	it is available from your laboratory.					
i icase pro	No. of		Allele Designations					
Locus	Alleles Provided		· ······					
10. DRB3		First DRB3*						
	2 □ two							
	3 ☐ none							
	4 ☐ not tested	Second DRB3*						
11. DRB4		First DRB4*						
	2 □ two							
	3 ☐ none							
	4 ☐ not tested	Second DRB4*						

CIB	MTR Ce	enter Number:		CIBMTR Recipient ID:				
Cla	lass II (Optional)							
		No. of		Allele Designations				
12.	Locus DRB5	Alleles Provided 1 □ one 2 □ two	First DRB5*					
		3 ☐ none 4 ☐ not tested	Second DRB5*					
13. DQB1	DQB1	1 □ one 2 □ two	First DQB1*					
			Second DQB1*					
14.	DPB1	1 □ one 2 □ two	First DPB1*					
			Second DPB1*					
15. DQA1	DQA1	1 □ one 2 □ two	First DQA1*					
			Second DQA1*					
16.	DPA1	1 □ one 2 □ two	First DPA1*					
			Second DPA1*					

	_						_			
CIBMTR Center Number:				CIBMTR Recipient ID:						

Antigens Defined by Serologic Typing

Use the following lists when reporting HLA-A and B antigens. Report broad antigens only when your laboratory was not able to confirm typing for a known split antigen.

A Antigens											
6. No. of antigens provided: 1 □ one 2 □ two											
Specificity	Antigen <u>Specificity 1st 2nd</u>										
A1		01									
A2		02									
A203		03									
A210		04									
A3		05									
A9		06									
A10		07									
A11		80									
A19		09 10									
A23(9) A24(9)		11									
A24(9) A2403		12									
A25(10)		13									
A26(10)		14									
A28		15									
A29(19)		16									
A30(19)		17									
A31(19)		18									
A32(19)		19									
A33(19)		20									
A34(10)		21									
A36		22									
A43		23 24									
A66(10) A68(28)		25									
A69(28)		26	Ä								
A74(19)		27									
A80		28									
AX		99									

		D Am	ti ma ma		
	г		tigens	7	
		7. Number of ar	ntigens provided:		
	Antigen		Antigen		
Specificity	<u>1st</u>	Specificity	<u>1st</u> <u>2nd</u>	Specificity	<u>1st</u>
B5	□ 01 □	B40	□ 21 □	B59	□ 42 □
B7	□ 02 □	B4005	□ 22 □	B60(40)	□ 43 □
B703	□ 03 □	B41	□ 23 □	B61(40)	□ 44 □
B8	□ 04 □	B42	□ 24 □	B62(15)	□ 45 □
B12	□ 05 □	B44(12)	□ 25 □	B63(15)	□ 46 □
B13	□ 06 □	B45(12)	□ 26 □	B64(14)	
B14	□ 07 □	B46	□ 27 □	B65(14)	□ 48 □
B15	□ 08 □	B47	□ 28 □	B67	□ 49 □
B16	□ 09 □	B48	□ 29 □	B70	□ 50 □
B17	□ 10 □	B49(21)	□ 30 □	B71(70)	□ 51 □
B18	11	B50(21)	□ 31 □	B72(70)	□ 52 □
B21	□ 12 □	B51(5)	□ 32 □	B73	□ 53 □
B22	□ 13 □	B5102	□ 33 □	B75(15)	□ 54 □
B27		B5103	□ 34 □	B76(15)	□ 55 □
B2708	□ 59 □	B52(5)	□ 35 □	B77(15)	□ 56 □
B35	□ 15 □	B53	□ 36 □	B78	□ 57 □
B37	□ 16 □	B54(22)	□ 37 □	B81	□ 58 □
B38(16)	□ 17 □	B55(22)		B82	
B39(16)	□ 18 □	B56(22)	□ 39 □ □ 40 □	BX	□ 99 □
B3901		B57(17)			
B3902	□ 20 □	B58(17)	□ 41 □		

CIBMTR Center Number:		CIBMTR Recipient ID:	
Ontional Antigan Banarti			
Optional Antigen Reportion	_	ailable from your laboratory. This in	nformation will be entered into the
NMDP Registry and be available		and the first your laboratory. This is	normation will be entered into the
Antigens Defined by Serologi	c Typing		
C Antigens	DR Antigens	DR51 Antigen	DP Antigens
8. No. of antigens provided: 1 □ one 2 □ two	11. No. of antigens provided: 1 □ one 2 □ two	Specificity Present? 12. DR51 1 ☐ yes 2 ☐ no	16. No. of antigens provided: 1 □ one 2 □ two
Antigen Specificity 1st 2nd	Antigen <u>Specificity</u> 1st 2nd	DR52 Antigen	Antigen <u>Specificity</u> 1st 2nd
Cw1	DR1	Specificity Present? 13. DR52 1 ☐ yes 2 ☐ no	DPw1
Cw3	DR2	DR53 Antigen	DPw3 □ 03 □ □ □ □ □ □ □ □ □ □ □ □ □ □ □ □ □
Cw4	DR4	Specificity Present? 14. DR53 1 ☐ yes 2 ☐ no	DPw5
Cw7 🗆 07 🗆	DR6 □ 07 □	DQ Antigens	DPX
Cw8	DR7 □ 08 □ □ □ DR8 □ 09 □ □ □ DR9 □ 10 □	15. No. of antigens provided: 1 □ one 2 □ two	
CX	DR10 🗆 11 🗆	Antigen	
	DR11(5)	Specificity 1st 2nd DQ1 □ 01 □	
Bw Specificity	DR13(6) □ 14 □	DQ2 🗆 02 🗆	
Specificity Present? 9. Bw4 1 □ yes 2 □ no	DR14(6) □ 15 □ DR1403 □ 16 □	DQ3	
10. Bw6 3 ☐ yes 4 ☐ no	DR1404	DQ5(1) 🗆 05 🗆	
	DR15(2) □ 18 □ □ DR16(2) □ 19 □	DQ6(1)	
	DR16(2) □ 19 □ □ DR17(3) □ 20 □	DQ8(3)	
	DR18(3) □ 81 □	DQ9(3)	
	DRX □ 99 □	DQX 🗆 99 🗆	
17. Signed:	Porcon co	mpleting form	
Please print name:	r erson con		
Phone number: ()	Fax number: ()
E-mail address:			
Continued from page 1			
If multiple cord blood units were in	nfused, record each of the non-N	MDP cord blood unit identification	numbers below:
Non-NMDP Cord Blood Unit ID:			
Non-NMDP Cord Blood Unit ID:			
If non-NMDP donors were used,	record the identification number a	and/or gender and date of birth for	each donor below:
Non-NMDP Donor ID:		— OR — Donor gender: Dono	or date of birth:
		1 □ male 2 □ female	nth Day Year
Non-NMDP Donor ID:		— OR — Donor gender: Dono	or date of birth:
CIBMTR Form 2005 v1.0 (5–5) July 2		1 □ male 2 □ female	
Copyright © 2007 National Marrow D The Medical College of Wisconsin, In	<u> </u>	2 🗖 Terriale Mor	nth Day Year
For internal use only: Document F00xxx vers			



Infectious Disease Markers

moonous Bissuss mai		
Pogietry IIco	Non-NMDP Cord Blood Unit ID: *	
Registry Use (* See page 3 for additional ID boxes or donor date of birth and gender.	
Sequence Number:	Today's Date: Day Year	
Data	Date of HSCT for which this form is being completed:	
Date Received:	HSCT type: ☐ allogeneic, ☐ allogeneic, ☐ syngeneic unrelated related (identical twin)	
	Product type: ☐ marrow ☐ PBSC ☐ cord blood ☐ other product, specify:	
		_
	or all non-NMDP allogeneic or syngeneic donors, or non-NMDP cord blood units. was secured through the NMDP, then report IDMs on forms 24 and 50 for allogeneic for cord blood units.	С
 Who is being tested for IDMs? □ donor IDM (marrow or PB\$ □ maternal IDM (cord blood) □ cord blood unit IDM 	SC)	
Infectious Disease Marker	(report final test results) Test Date	
	Month Day Year	
 1 □ reactive 2 □ non-reactive 3 □ testing not performed 	Month Day Year	
Hepatitis B Virus (HBV) 2. HBsAg: (hepatitis B surface and predictive and predic	igen) 3. Month Day Year 2 0	
Hepatitis B Virus (HBV) 2. HBsAg: (hepatitis B surface and predictive and predic	igen) 3. Month Day Year 2 0 pody) (no confirmatory test available) 5. 2 0	
Hepatitis B Virus (HBV) 2. HBsAg: (hepatitis B surface and 1 □ reactive 2 □ non-reactive 3 □ testing not performed 4. Anti HBc: (hepatitis B core antile 1 □ reactive 2 □ non-reactive 3 □ testing not performed Hepatitis C Virus (HCV) 6. Anti-HCV: (hepatitis C antibody 1 □ reactive 2 □ non-reactive	igen) 3. Month Day Year 2 0 pody) (no confirmatory test available) 5. 2 0	

CIBMTR Center Number:

CIBMTR Recipient ID:

Non-NMDP Donor ID: *

CIBMTR Center Number:		CIBMTR Recipient ID:							
		_							
Infectious Disease Ma	arker (report final test results)								
Human Immunodeficiency \ 10. HIV-1 p24 antigen: 1 □ reactive 2 □ non-reactive 3 □ not reported 4 □ not performed; HIV	Virus (HIV) NAT testing performed <i>(skip date)</i>	11	Month .	Day	2	Year O			
12. Was FDA licensed NAT	testing for HIV-1 / HCV performed	?							
1 □ yes — → 2 □ no	Specify results: 13. HIV-1 1 □ positive 2 □ negative 3 □ not reported	14			2	0			
	15. HCV 1 ☐ positive 2 ☐ negative	16			2	0			
•	nmunodeficiency Viruses) ss is required. This testing may be perform	18 ed as separate tests or done using a		assay.	2	0			
Syphilis									
19. STS:1 □ reactive2 □ non-reactive3 □ testing not performe	ed	20).		_	0			
Cytomegalovirus (CMV)									
21. Anti-CMV: (IgG or Total) 1 ☐ reactive 2 ☐ non-reactive 3 ☐ previously reported 4 ☐ testing not performe	reactive, not tested (skip date)	22	2		_	0			
West Nile Virus (WNV)									
23. WNV-NAT testing: 1 □ positive 2 □ negative 3 □ testing not performe 4 □ not applicable	ed	24			_	0			
25. Other infectious disease	marker, specify (e.g., EBV):								
1 □ yes	26. Specify date performed: 27. Specify test and method:				2	0			

CIBINITR Center Number:				CIBMIR Re	cipient iD:					
Infectious Disease Ma				Test Da	ate					
29. Other infectious disease	marker, spec	cify (e.g., EBV)	:			Month	Day	Y	⁄ear	
1 □ yes — ➤ 2 □ no					20					
	31. Specify	test and meth	od:							
	32. Specify	test results: _								
33. Other infectious disease	marker, spec	ify (e.g., EBV)	:							
1 □ yes — ➤ 2 □ no	34. Specify	date performe	ed:					20		
	35. Specify	test and meth	od:							
	36. Specify	test results: _								
37. Signed:				npleting form						
Please print name:										
Phone number: ()									
Fax number: ()									
E-mail address:										
Continued from page 1										
If multiple cord blood units we	ere infused, re	ecord each of t	the non-NN	IDP cord blo	ood unit ide	ntification nu	mbers belo	ow:		
Non-NMDP Cord Blood Unit	ID:									
Non-NMDP Cord Blood Unit	ID:									
If non-NMDP donors were us	ed, record the	e identification	number aı	nd/or gender	and date o	f birth for ea	ch donor b	elow:		
Non-NMDP donor ID:										
— OR —	Donor gender:	male female	Donor o	date of birth:	Month	Day	Year			
Non-NMDP donor ID:										
— OR —	Donor gender:	□ male □ female	Donor o	date of birth:	Month	Day	Year			
						Du v				

Month

Day

Year