OMB No: 0915-0310

Expiration Date: 10-31-2010

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 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.



# Hematopoietic Stem Cell Transplant (HSCT) Infusion

	Registry Use Only							
Sequence Number:								
Date Received:								

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 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 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.

CIBMTR Center Number:	OMB No: 0915-0310 Expiration Date: 10/31/2010
CIBMTR Recipient ID:	
Specify donor:  1 □ autologous 2 □ NMDP unrelated	
cord blood unit →	NMDP Cord Blood Unit ID:
3 ☐ NMDP unrelated donor — ➤	NMDP Donor ID:
4 ☐ related donor —→  5 ☐ non-NMDP unrelated donor →  6 ☐ non-NMDP cord blood unit — → (include related and autologous CBUs)	Donor's / infant's date of birth:    Month   Day   Year
Today's Date: Month	Day Year
Date of HSCT for which the being completed:	is form is  Month Day  Year
HSCT type: ☐ autologo (check only one)	us □ allogeneic, □ allogeneic, □ syngeneic unrelated related (identical twin)
Product type: ☐ marrow (check only one)	□ PBSC □ cord blood □ other product, specify:

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.

CIBMTR Center Number:	CIBMTR Recipient ID:
Pre-Collection Therap	y
	atment, prior to any stem cell harvest, to enhance the product collection for this HSCT? as from an NMDP donor, or the product is a cord blood unit, then continue with question 20.)
1 ☐ yes — → 2 ☐ no 3 ☐ NMDP donor	Specify treatment(s): (select all that apply)  2. 1 ☐ yes 2 ☐ no (autologous only)
Continue with question 20	4. 1 ☐ yes 2 ☐ no Growth factor(s) —➤ If yes, specify growth factor(s):
4 □ cord blood unit ↓	5. 1 ☐ yes 2 ☐ no G-CSF 6. 1 ☐ yes 2 ☐ no GM-CSF 7. 1 ☐ yes 2 ☐ no Other →  8. Specify:
Continue with question 20	9. 1 ☐ yes 2 ☐ no Other treatment → 10. Specify treatment:
Product Collection  11. Date of product collection  12. Was more than one colle  1 □ yes  2 □ no	n: Day Year  ction required for this HSCT?   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.
14. Were anticoagulants add	ed to the product during collection?
1 □ yes	Specify anticoagulant(s):  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 2  no  19. Specify other anticoagulant: 2  no

CIBMTR Center Number:		CIBMTR Red	cipient ID:										
							'						
Product Transport an	d Receipt												
20. Was this product collect	ed off-site and shipped to	your facility?											
1 □ yes ———							 1						
2 □ no	21. Date of receipt of product at your facility:    Month   Day   Year   Total Activities   Total Activities												
	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 en	vironment:										
	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												
	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	27. (Cord blood product only) Was the cord blood unit stored at your center prior to thawing?											
	1 ☐ yes — ➤ 28. Specify the storage method used for the cord blood unit:												
	2 □ no	1 ☐ liquid nitrogen	metriou use	ed for the	e cora bi	ood unit.							
		2 ☐ vapor phase 3 ☐ electric freezer											
		29. Temperature during storage: — ° C											
		30. Date storage started:    Month   Day   Year											
Product Processing /	Manipulation												
31. Was a fresh product red	•	at your facility prior to inf	fusion?										
1 □ yes													
2 🗖 no													
₃ ☐ not applicable, cord	blood unit												
32. Was the product thawed	from a cryopreserved sta	te prior to infusion?											
1 □ yes ——— <del>&gt;</del> 2 □ no	33. Was the entire prod	uct thawed?											
	1 □ yes												
	2 □ no — →	34. Was a compartmen	t of the bag	thawed'	?								
		1 □ yes											
		2 🗆 no		0									
		35. Were there multiple	product bag	gs?									
		1 □ yes> 2 □ no	36. Speci	fy numb	er of bag	gs thawed	: [	$\square \square \square \square$					
CIBMTR Form 2006 revision 2	 (page 3 of 13) .lune 2009	2 <b></b> 110				-							
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CIBMTR Center Number:	CIBMTR Recipient ID:									
	37. Date thawing process initiated:    Month   Day   Year									
	38. Time at initiation of thaw (24-hour clock):  ☐ I ☐ standard time ☐ Description of them (24-hour clock): ☐ Hour ☐ Hour ☐ daylight savings time									
	39. Time at completion of thaw (24-hour clock):  Hour  1 □ standard time 2 □ daylight savings time									
	<ul> <li>40. Was the primary container (e.g., cord blood unit bag) intact upon thawing?</li> <li>1 □ yes</li> <li>2 □ no</li> <li>41. What method was used to thaw the product?</li> <li>1 □ no wash — thawed at bedside, then infused</li> <li>2 □ DMSO dilution — thawed in lab (added doxtrap and albumin), then infused</li> </ul>									
	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:									
	<ul><li>43. Did any adverse events or incidents occur while thawing the product?</li><li>1 □ yes</li><li>2 □ no</li></ul>									
44. Was the product manip	ulated 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 other method:									
	54. Ex-vivo expansion  1 ☐ yes 2 ☐ no  55. Genetic manipulation (gene transfer / transduction)  1 ☐ yes 2 ☐ no									
CIBMTR Form 2006 revision 2 Copyright © 2009 National Mari	56. Volume reduction  1 ☐ yes 2 ☐ no  (page 4 of 13) June 2009									
Sopringing S 2000 Hational Mail	on Bonor Fogiam and									

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Internal use: Document number F00481 revision 2 Replaces: F00481 version 1.0 July 2007

1 □ yes ———> 2 □ no	58. Specify manufac	cturer:
2 🗖 110	1 🗆 CliniMACS	/ CliniMax
	2 ☐ Isolex	
	3 □ other ——	→ 59. Specify other manufacturer:
60. T-cell depletion		
1 □ yes>	Specify method:	
2 🗖 no	1 .	o Antibody affinity column —
	· ·	o Antibody coated plates
	· ·	o Antibody coated plates antibodies
		and soybean lectin   used for T-ce
	64 1 □ yes 2 □ n	o Antibody + complement
	-	o Antibody + toxin — <b>question 73</b> .
	66. 1 □ yes 2 □ no	•
	67. 1 □ yes 2 □ no	
	•	o CD34 affinity column plus sheep red blood cell
		rosetting 🖺
	69. 1 □ yes 2 □ n	o Other — 70. Specify other method:
71. Other cell		
manipulation		
1 □ yes>	- 72. Specify other ce	ell manipulation:
2 <b>□</b> no	· = · opcomy outlot oc	
73. Were antibodies ບ	sed during product mai	nipulation?
1 □ yes>		•
2 <b>□</b> no	Specify antibodies:	4 4 000
	74. 1 🗆 yes 2 🗆 no	
		o Anti CD3
	75. 1 🗆 yes 2 🗖 no	
	76. 1 □ yes 2 □ no	o Anti CD4
	76. 1 🗆 yes 2 🗆 no	o Anti CD4 o Anti CD5
	76. 1 🗆 yes 2 🗆 no 77. 1 🗆 yes 2 🗆 no 78. 1 🗆 yes 2 🗆 no	o Anti CD4 o Anti CD5 o Anti CD6
	76. 1  yes 2  ne 77. 1  yes 2  ne 78. 1  yes 2  ne 79. 1  yes 2  ne	o Anti CD4 o Anti CD5 o Anti CD6 o Anti CD7
	76. 1 🗆 yes 2 🗆 no 77. 1 🗆 yes 2 🗆 no 78. 1 🗆 yes 2 🗆 no	o Anti CD4 o Anti CD5 o Anti CD6 o Anti CD7
	76. 1  yes 2  ne 77. 1  yes 2  ne 78. 1  yes 2  ne 79. 1  yes 2  ne	o Anti CD4 o Anti CD5 o Anti CD6 o Anti CD7 o Anti CD8
	76. 1  yes 2  ne 77. 1  yes 2  ne 78. 1  yes 2  ne 79. 1  yes 2  ne 80. 1  yes 2  ne	o Anti CD4 o Anti CD5 o Anti CD6 o Anti CD7 o Anti CD8 o Anti CD8
	76. 1  yes 2  no 77. 1  yes 2  no 78. 1  yes 2  no 79. 1  yes 2  no 80. 1  yes 2  no 81. 1  yes 2  no	o Anti CD4 o Anti CD5 o Anti CD6 o Anti CD7 o Anti CD8 o Anti CD34 o Anti TCR alpha / beta (T10-B9)
	76. 1  yes 2  no 77. 1  yes 2  no 78. 1  yes 2  no 79. 1  yes 2  no 80. 1  yes 2  no 81. 1  yes 2  no 82. 1  yes 2  no	o Anti CD4 o Anti CD5 o Anti CD6 o Anti CD7 o Anti CD8 o Anti CD34 o Anti CD34 o Anti TCR alpha / beta (T10-B9) o OKT-3
	76. 1  yes 2  ne 77. 1  yes 2  ne 78. 1  yes 2  ne 79. 1  yes 2  ne 80. 1  yes 2  ne 81. 1  yes 2  ne 82. 1  yes 2  ne 83. 1  yes 2  ne 83. 1  yes 2  ne	o Anti CD4 o Anti CD5 o Anti CD6 o Anti CD7 o Anti CD8 o Anti CD34 o Anti TCR alpha / beta (T10-B9) o OKT-3
	76. 1  yes 2  no 77. 1  yes 2  no 78. 1  yes 2  no 79. 1  yes 2  no 80. 1  yes 2  no 81. 1  yes 2  no 82. 1  yes 2  no 83. 1  yes 2  no 84. 1  yes 2  no 84. 1  yes 2  no	o Anti CD4 o Anti CD5 o Anti CD6 o Anti CD7 o Anti CD8 o Anti CD34 o Anti TCR alpha / beta (T10-B9) o OKT-3 o Other CD3   85. Specify other CD3:
	76. 1  yes 2  ne 77. 1  yes 2  ne 78. 1  yes 2  ne 79. 1  yes 2  ne 80. 1  yes 2  ne 81. 1  yes 2  ne 82. 1  yes 2  ne 83. 1  yes 2  ne 83. 1  yes 2  ne	o Anti CD4 o Anti CD5 o Anti CD6 o Anti CD7 o Anti CD8 o Anti CD34 o Anti TCR alpha / beta (T10-B9) o OKT-3 o Other CD3   85. Specify other CD3:
	76. 1  yes 2  no 77. 1  yes 2  no 78. 1  yes 2  no 79. 1  yes 2  no 80. 1  yes 2  no 81. 1  yes 2  no 82. 1  yes 2  no 83. 1  yes 2  no 84. 1  yes 2  no 84. 1  yes 2  no	o Anti CD4 o Anti CD5 o Anti CD6 o Anti CD7 o Anti CD8 o Anti CD34 o Anti TCR alpha / beta (T10-B9) o OKT-3 o Other CD3   85. Specify other CD3:
	76. 1  yes 2  no 77. 1  yes 2  no 78. 1  yes 2  no 79. 1  yes 2  no 80. 1  yes 2  no 81. 1  yes 2  no 82. 1  yes 2  no 83. 1  yes 2  no 84. 1  yes 2  no 84. 1  yes 2  no	o Anti CD4 o Anti CD5 o Anti CD6 o Anti CD7 o Anti CD8 o Anti CD34 o Anti TCR alpha / beta (T10-B9) o OKT-3 o Other CD3 →  85. Specify other CD3:  ———————————————————————————————————
	76. 1  yes 2  no 77. 1  yes 2  no 78. 1  yes 2  no 79. 1  yes 2  no 80. 1  yes 2  no 81. 1  yes 2  no 82. 1  yes 2  no 83. 1  yes 2  no 84. 1  yes 2  no 84. 1  yes 2  no	o Anti CD4 o Anti CD5 o Anti CD6 o Anti CD7 o Anti CD8 o Anti CD34 o Anti TCR alpha / beta (T10-B9) o OKT-3 o Other CD3 →  Specify antibodies:  yes no 87. 1 □ 2 □ Campath-NOS 88. 1 □ 2 □ Campath-1G
	76. 1  yes 2  no 77. 1  yes 2  no 78. 1  yes 2  no 79. 1  yes 2  no 80. 1  yes 2  no 81. 1  yes 2  no 82. 1  yes 2  no 83. 1  yes 2  no 84. 1  yes 2  no 84. 1  yes 2  no	o Anti CD4 o Anti CD5 o Anti CD6 o Anti CD7 o Anti CD8 o Anti CD34 o Anti CD34 o Anti TCR alpha / beta (T10-B9) o OKT-3 o Other CD3 →  Specify antibodies:  yes no 87. 1 □ 2 □ Campath-NOS 88. 1 □ 2 □ Campath-1G 89. 1 □ 2 □ Campath-1H

CIBMTR Recipient ID:

CIBMTR Center Number:

CIBMTR Center Number: CIBMTR Recipient ID:												
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### **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?

2 □ no	Specify tumor cell detection method used, and site(s) of tumor cells:									
	93. Routine									
	histopathology	Chaoify aita(a)								
	1 □ yes — ➤ 2 □ no	Specify site(s):	o □ not tooted	Circulation blood calls						
	2 🗖 110			Circulating blood cells  Bone marrow, in the interval						
	07. Daharana	95. 1 Li yes 2 Li 110	3 Li not tested	between last systemic therapy and collection						
	97. Polymerase chain reaction (PCR)	96. 1 □ yes 2 □ no	3 ☐ not tested	Collected cells, before purging						
	1 □ yes — ➤	Specify site(s):								
	2 🗆 no		3 ☐ not tested	Circulating blood cells						
				Bone marrow, in the interval between last systemic therapy						
	404 Other medicalism	100 4 D vos 3 D no	a □ not tosted	and collection Collected cells, before purging						
	101. Other molecular technique	100. 1 Li yes 2 Li 110	3 🗖 Hot tested	Collected Cells, before purging						
	1 ☐ yes ——➤	102. Specify method: _								
	2 🗖 no	Specify site(s):								
		103. 1 ☐ yes 2 ☐ no	3 ☐ not tested	Circulating blood cells						
		104. 1 ☐ yes 2 ☐ no	3 ☐ not tested	Bone marrow, in the interval between last systemic therapy and collection						
	106. Immunohisto-	105. 1 ☐ yes 2 ☐ no	3 ☐ not tested	Collected cells, before purging						
	chemistry 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	Charify =!t=/=>								
	1 □ yes ———— 2 □ no	Specify site(s):	2 □ not tosted	Circulating blood cells						
	2 110	-		Bone marrow, in the interval						
		112. 1 D you 2 D 110	o 🗖 not toolog	between last systemic therapy and collection						
	44.4 Other technique	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):								
		1 ' ' '	3 ☐ not tested	Circulating blood cells						
		1		Bone marrow, in the interval between last systemic therapy						
	page 6 of 13) June 2009			and collection						

		$\overline{}$	$\overline{}$								т т				$\overline{}$	$\overline{}$		
CIBMTR Center Number:	:						CI	BMTR Re	cipient I	D:								
				1													1 1	
119. Was the product trea	ated	to rem	nove	mali	ignant	cells	s (purged)? (au	tologous p	product (	only)								
1 🔲 yes ———	-	Spec	ifv n	netho	d(s) u	ised:												
2 🗖 no		120. 1 ☐ yes 2 ☐ no Mo						ibody →	121. If	ves. sı	pecify:							
																		_
							4-hydroperoxyd	cyclophosp	hamide	(4HC)								
							Mafosfamide		405.14	•		_						
							Other drug —	<b></b>	125. 11	yes, s	pecity:	<u> </u>						_
				-			Elutriation											
				-			mmunomagne											
		128.	1 ∐	yes	2 🏻 1	no	Toxin ———	<b></b>	129. lf	yes, s	pecify:	: <u> </u>						_
		130.	1 🛮	yes	2 🗖 I	no F	Positive stem c	ell selection	on									
							other than pre											
						of mononuclea		404 14		.,								
							fraction) ——	<b></b>	131. 11	yes, s	pecity	me	tnod	l:				_
		122	<b>,</b> П	V00	۰П.	no (	Other method -		122 1		if-							
		132.	' Ш	yes	2 <b>ப</b> 1	110			133. 11	yes, s	pecity.	•						
		Specify if tumor cells were detected in the graft after purging by each method used:																
			134. 1 ☐ yes 2 ☐ no 3 ☐ not tested Routine histopathology															
				•				□ not tested Polymerase chain reaction (PCR)										
				-				Other molecular technique										
				-				Immunohistochemistry										
				-			not tested			-								
		139.	1 🗆	ves	2 🗆 1	no s	not tested	Other >	440.11									_
			. —	,				J 2	140. Ii	yes, s	pecity:	:						_
	L																	
Product Applysis /	<b>A II</b>	Drod	اميا	tc\ ſ	Υī													
Product Analysis (	AII I	riou	luc	ເວ <i>ງ</i> ⊾														
		Produ	ıct /	Analy	sis at	t 1st	Timepoint		Pro	duct A	nalys	is a	ıt 2r	nd T	imepo	int		
Specify the timepoint in	141	1 □	pro	duct	arrival	ı			162. 1	□ proc	duct a	rriva	al					
the product preparation			-							□ post								
phase that the product					prese		on /		_	-	-cryop		-	tion	/			
was analyzed:					ation [					-	nipulat							
		з 🗖	pos	st-tha	W				3	☐ post	t-thaw	,						
		4 🔲	pos	st-ma	nipula	tion			4	□ post	t-mani	ipula	atior	1				
		5 🗖	at i	nfusio	on (fin	al qu	antity infused)		5	□ at ir	nfusior	າ (fir	nal c	quan	tity info	used)		
						] [		1					7 [				1	
Date of product analysis:	142.					2	2  <i>0</i>		163.					2	0			
		Mor	nth		Day		Year	-		Month	D	ay			Year		-	
Total volume of product:	1/12						1 □ mL		164						1 □ n	ηL		

Total volume of product: 143.

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

		Product Analysi	nt	Product Analysis at 2nd Timepoint					
In this section, report	the t	otal number of o	cells (not cells p	er kilogram).					
		Total Numb	er Expon	ent		Total Numb	er	Expone	ent
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 1	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 teste	ed	172.		% 🗆	not teste	ed
Method of testing cell viability:	152.	1 ☐ 7-AAD 2 ☐ propidium 3 ☐ trypan blue 4 ☐ other meth	•		3 □	7-AAD propidium trypan blu other meth	е		
Specify other method:	153.								
Were the colony-forming units (CFU) assessed after thawing? (cord blood product only)	154.		Continue with qu			,			estion 176 estion 179
Was there growth?	155.	1 □ yes 2 □ 1	no		176. <sub>1</sub> 🗆	] yes 2 □	no		
Total colonies per product:	156.		x 10 <sup>5</sup>	☐ unknown	177.			x 10 <sup>5</sup>	□ unknown
Total CFU-GM:	157.		• x 10 <sup>5</sup>	unknown	178.		•	x 10 <sup>5</sup>	□ unknown
Were cultures performed before infusion to test the product(s) for bacterial o fungal infection? (complete for all cell prod	e r	2 □ no →	Continue with qu			,			estion 180 estion 183
Specify results:	,		☐ negative 3 ☐	unknown	180. 1 <b>C</b>	positive 2	☐ negat	ive ₃□	unknown
Specify organism code(s):	160.				181.				
(see page 9 for codes)					100				
If code 198, 209,	161.				182				

219, or 259, specify

organism:

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 171 Chlamydia pneumoniae 213 Asperaillus niger 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 except difficile) 241 Mucormycosis (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)

‡ 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.

502 Suspected bacterial

1 ☐ standard time

2 ☐ daylight savings time

infection

154 Propionibacterium (acnes,

species)

avidum, granulosum, other

#### **Product Infusion**

138 Flavobacterium

136 Escherichia (also E. coli)

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

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

Internal use: Document number F00481 revision 2 Replaces: F00481 version 1.0 July 2007

CIBMTR Center Number:		CIBMTR Recip	nient ID:							
OIBINITY OCINCI IVAINOCI.		OIDWITT TOOK	piciti ib.							
188. Total volume of product	plus additives infused:	ml ell	L							
189. Specify the route of prod	luct infusion:									
1 ☐ intravenous										
2 ☐ intramedullary										
3 ☐ intraperitoneal										
4 □ other route of infusion ———	190. Specify route of in	ıfusion:								
191. Did the volume of infuse	d product include any a	dded agents?								
1 □ yes ———	0									
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	•	ify agent:							.
	·									
199. Was the entire volume o	f product infused?									
1 ☐ yes										
2 □ no ———		pened to the reserved portion	າ:							
	1 discarded	d for future was								
	2 ☐ cryopreserved 3 ☐ other fate →	_								_
		004 0 'r								
		201. Specify:								
		201. Specify:								
The following guestions re	efer to all stem cell pro		us marrov	w or aut	tologo	ous PB	SC proc	ducts	<u> </u>	
		educts except for autologoulogous PBSC product, con								•
If this HSCT used an autol	ogous marrow or auto	ducts except for autologou logous PBSC product, con	itinue wit							•
	ogous marrow or auto	ducts except for autologoulogous PBSC product, con	itinue wit							•
If this HSCT used an autol 202. Were there any adverse	ogous marrow or auto	oducts except for autologour logous PBSC product, controlled with the stem cell information diverse event(s):	usion?	h the si	gnatu Medical		s at que	estio	n 296	
If this HSCT used an autol 202. Were there any adverse 1 □ yes ———	events or incidents assessed. Specify the following an	educts except for autologour logous PBSC product, consciented with the stem cell infective event(s):  Adverse Event	usion?	h the si	gnatu  Medical on?	re line	s at que	estio	n 296	
If this HSCT used an autol 202. Were there any adverse 1 □ yes ———	events or incidents associated Specify the following at 203. 1 \( \text{yes} \) yes \( 2 \) no	educts except for autologouologous PBSC product, consociated with the stem cell information of the diverse event(s):  Adverse Event  Brachycardia	usion? R 204. 1	tequired National Republic Required National Republic Required National Republic Rep	gnatu ledical on? 2 □ no	re line	Resc 5. 1 □ y	estio	n <b>296</b>	<b>o</b>
If this HSCT used an autol 202. Were there any adverse 1 □ yes ———	events or incidents associated Specify the following and 203. 1  yes 2  no 206. 1 yes 2 no	educts except for autologouologous PBSC product, consociated with the stem cell infective event(s):  Adverse event Brachycardia Chest tightness / pain	usion?  R 1 204. 1 [ 207. 1 [	equired Noterventically yes 2	gnatu  Medical on?  2 □ no	208 208	Resc 5. 1 🗆 y 3. 1 🗆 y	estio	n 296	0
If this HSCT used an autol 202. Were there any adverse 1 □ yes ———	events or incidents assessed specify the following at 203. 1  yes 2  no 206. 1 yes 2  no 209. 1 yes 2 no	oducts except for autologouologous PBSC product, consciented with the stem cell infective event(s):  Adverse Event  Brachycardia  Chest tightness / pain  Chills at time of infusion	usion?  R 1 204. 1 [ 207. 1 [	tequired National Republic Required National Republic Required National Republic Rep	gnatu  Medical on?  2 □ no	208 208	Resc 5. 1 □ y	estio	n 296	0
If this HSCT used an autol 202. Were there any adverse 1 □ yes ———	events or incidents associated Specify the following and 203. 1  yes 2  no 206. 1 yes 2 no	ducts except for autologouologous PBSC product, conciated with the stem cell infudverse event(s):  Adverse Event  Brachycardia  Chest tightness / pain  Chills at time of infusion  Fever ≤ 103° F within 24	etinue wit usion? R I 204. 1 E 207. 1 E 210. 1 E	tequired Mantervention yes 2 yes 2 yes 2	Medical on?	205 205 207 217	Resc 5. 1  y 3. 1  y 1. 1  y	estio	n 296	D D D
If this HSCT used an autol 202. Were there any adverse 1 □ yes ———	events or incidents assorting the following are 203. 1  yes 2  no 206. 1  yes 2  no 209. 1  yes 2  no 212. 1  yes 2  no	educts except for autologouologous PBSC product, consciented with the stem cell infectores event(s):  Adverse Event Brachycardia Chest tightness / pain Chills at time of infusion Fever ≤ 103° F within 24 hours of infusion	etinue wit usion? R I 204. 1 E 207. 1 E 210. 1 E	equired Noterventically yes 2	Medical on?	205 206 207 217	Resc 5. 1 🗆 y 3. 1 🗆 y	estio	n 296	D D D
If this HSCT used an autol 202. Were there any adverse 1 □ yes ———	events or incidents assorting the following are 203. 1  yes 2  no 206. 1  yes 2  no 209. 1  yes 2  no 212. 1  yes 2  no	educts except for autologouologous PBSC product, consciented with the stem cell infectores event(s):  Adverse Event  Brachycardia  Chest tightness / pain  Chills at time of infusion  Fever ≤ 103° F within 24  hours of infusion  Fever > 103° F within 24	etinue wit usion? R 204. 1 E 207. 1 E 210. 1 E	equired Material yes 2 yes 2 yes 2	Medical on? 2	200 200 200 211 214	Resc 5. 1  y 3. 1  y 1. 1  y	estio	n 296	D D D
If this HSCT used an autol 202. Were there any adverse 1 □ yes ———	events or incidents assessed specify the following at 203. 1  yes 2 no 206. 1 yes 2 no 209. 1 yes 2 no 212. 1 yes 2 no 215. 1 yes 2 no	oducts except for autologouologous PBSC product, conciled with the stem cell information ociated with the stem cell information continuation ociated with the stem cell information.  Adverse Event Brachycardia Chest tightness / pain Chills at time of infusion Fever ≤ 103° F within 24 hours of infusion Fever > 103° F within 24 hours of infusion	etinue wit usion? R 204. 1 E 207. 1 E 210. 1 E 213. 1 E	equired Montervention yes 2 yes 2 yes 2 yes 2	Medical on? 2    nc 2    nc 2    nc	205 208 207 217 214 217	Resc 5. 1  y 3. 1  y 1. 1  y 4. 1  y	estio	n 296	
If this HSCT used an autol 202. Were there any adverse 1 □ yes ———	events or incidents assorting the following are specify the following are specify the following are specify the following are specify the following are specifically as a specific probability of the following are specifically as a specific probability of the following are specifically as a specific probability of the following are specifically as a specific probability of the following are specifically as a specific probability of the following are specifically as a specific probability of the following are specifically as a specific probability of the following are specifically as a specific probability of the following are specifically as a specific probability of the following are specifically as a specific probability of the following are specifically as a specific probability of the following are specifically as a specific probability of the following are specifically as a specific probability of the following are specifically as a specific probability of the following are specifically as a specific probability of the following are specifically as a specific probability of the following are specifically as a specific probability of the following are specifically as a specific probability of the following are specific probability of the following are specifically as a specific probability of the following are specific probability of the following ar	ducts except for autologouologous PBSC product, conception of the stem cell information of the stem ce	etinue wit usion? R I 204. 1 E 207. 1 E 210. 1 E 213. 1 E 216. 1 E 219. 1 E	equired Materials and sequence of the sequence	Medical on? 2	209 209 211 214 217 220 220	Resc 5. 1  y 8. 1  y 1. 1  y 4. 1  y 7. 1  y 7. 1  y	blved? es 2 es 2 es 2 es 2	n 296	
If this HSCT used an autol 202. Were there any adverse 1 □ yes ———	events or incidents asservents asservents or incidents asservents as a second asservent as a second as	educts except for autologouologous PBSC product, consciented with the stem cell infectores event(s):  Adverse Event Brachycardia Chest tightness / pain Chills at time of infusion Fever ≤ 103° F within 24 hours of infusion Fever > 103° F within 24 hours of infusion Gross hemoglobinuria Headache	204. 1 E 207. 1 E 213. 1 E 216. 1 E 219. 1 E 222. 1 E	equired Material yes 2	Medical on? 2	208 208 211 214 217 220 220 220	Resc 5. 1  y 3. 1  y 1. 1  y 4. 1  y 7. 1  y 7. 1  y 8. 1  y	blved? es 2 es 2 es 2 es 2 es 2	2	
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If this HSCT used an autol 202. Were there any adverse 1 □ yes ———	events or incidents assorting the following are 203. 1 ☐ yes 2 ☐ no 206. 1 ☐ yes 2 ☐ no 209. 1 ☐ yes 2 ☐ no 212. 1 ☐ yes 2 ☐ no 215. 1 ☐ yes 2 ☐ no 221. 1 ☐ yes 2 ☐ no 227. 1 ☐ yes 2 ☐ no 200. 1 ☐ yes 2 ☐	ducts except for autologouologous PBSC product, consciented with the stem cell information diverse event(s):  Adverse Event Brachycardia Chest tightness / pain Chills at time of infusion Fever ≤ 103° F within 24 hours of infusion Fever > 103° F within 24 hours of infusion Gross hemoglobinuria Headache Hives Hypertension	204. 1 E 207. 1 E 210. 1 E 219. 1 E 222. 1 E 228. 1 E	equired Material yes 2	Medical on?  2	200 200 211 212 220 220 220 220 220 220	Resc 5. 1  y 7. 1  y 4. 1  y 7. 1  y	estio	2   no	
If this HSCT used an autol 202. Were there any adverse 1 □ yes ———	events or incidents asserbered property and events or incidents asserbered property and events or incidents asserbered property and events or incidents asserbered property as a second propert	educts except for autologouologous PBSC product, consociated with the stem cell information deciated with the stem cell information consistency and consistency and consistency are stated as a second consistency and consistency are stated as a second consistency and consistency are stated as a second consistency and consistency are stated consistency are stated consistency and consistency are stated consistency are stated consistency and consistency are stated consistency and consistency are stated consistency are stated consistency are stated consistency and consistency are stated consistency are stated consistency and consistency are stated consistency and consistency are stated consistency are stated consistency and consistency are stated consistency and consistency are stated consistency and consistency are stated consistency and consistency are stated consistency are stated consistency are stated consistency are stated consistency and consistency are stated consistency and consistency	204. 1 E 207. 1 E 210. 1 E 219. 1 E 222. 1 E 228. 1 E	equired Mantervention yes 2	Medical on?  2	200 200 211 212 220 220 220 220 220 220	Resc 5. 1  y 3. 1  y 1. 1  y 4. 1  y 7. 1  y 7. 1  y 8. 1  y 6. 1  y	estio	2   no	
If this HSCT used an autol 202. Were there any adverse 1 □ yes ———	events or incidents assorting the following are 203. 1 ☐ yes 2 ☐ no 206. 1 ☐ yes 2 ☐ no 209. 1 ☐ yes 2 ☐ no 212. 1 ☐ yes 2 ☐ no 215. 1 ☐ yes 2 ☐ no 221. 1 ☐ yes 2 ☐ no 227. 1 ☐ yes 2 ☐ no 200. 1 ☐ yes 2 ☐	educts except for autologouologous PBSC product, consciented with the stem cell infectores event(s):  Adverse Event Brachycardia Chest tightness / pain Chills at time of infusion Fever ≤ 103° F within 24 hours of infusion Fever > 103° F within 24 hours of infusion Gross hemoglobinuria Headache Hives Hypertension Hypotension Hypoxia requiring oxygen	204. 1 E 207. 1 E 210. 1 E 213. 1 E 216. 1 E 222. 1 E 225. 1 E 228. 1 E 231. 1 E	equired Marterventic yes 2	Medical on?  Incomplete on the control on the contr	200 200 200 211 20 220 220 220 232	Resconding to the second of th	estio  blved? es 2	2   no	
If this HSCT used an autol 202. Were there any adverse 1 □ yes ———	events or incidents asserbered processes as a second processes as	ducts except for autologouologous PBSC product, consciented with the stem cell information occiated with the stem cell inform	204. 1 E 213. 1 E 216. 1 E 219. 1 E 225. 1 E 228. 1 E 231. 1 E	tequired Montervention yes 2	Medical on?  I no construction on the construction of the construc	200 200 200 217 217 200 220 220 220 220 220 220 220 220 22	Resc 5. 1  y 3. 1  y 4. 1  y 4. 1  y 6. 1  y 6. 1  y 9. 1  y 2. 1  y	estio  blved3 es 2	2	
If this HSCT used an autol 202. Were there any adverse 1 □ yes ———	events or incidents asserbered property and events or incidents asserbered property and events or incidents asserbered property and events or incidents asserbered property as a second propert	aducts except for autologous logous PBSC product, consociated with the stem cell information of the diverse event (s):  Adverse Event Brachycardia Chest tightness / pain Chills at time of infusion Fever ≤ 103° F within 24 hours of infusion Fever > 103° F within 24 hours of infusion Gross hemoglobinuria Headache Hives Hypertension Hypotension Hypoxia requiring oxygen (O₂) support Nausea	204. 1 E 207. 1 E 210. 1 E 213. 1 E 216. 1 E 229. 1 E 225. 1 E 228. 1 E 231. 1 E	equired Marterventic yes 2	Medical on?  Incomplete on the control on the contr	208 208 209 211 212 223 223 223 223 223 223 223 223	Resconding to the second of th	estio  blved? es 2	2   no	

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Internal use: Document number F00481 revision 2 Replaces: F00481 version 1.0 July 2007

CIBMTR Center Number:		CIBMTR Recip	pient ID:	
		Adverse Event	Required Medical Intervention?	Resolved?
	245. 1 □ yes 2 □ no	Shortness of breath (SOB)	•	·
	•	•	249. 1 ☐ yes 2 ☐ no	•
	251. 1 ☐ yes 2 ☐ no	•	252. 1 ☐ yes 2 ☐ no	o 253. 1 □ yes 2 □ no
	254. 1 ☐ yes 2 ☐ no	Other expected AE 255. Specify:		
			256. 1 □ yes 2 □ no	o 257. 1 □ yes 2 □ no
	258. 1 □ yes 2 □ no	Other unexpected AE 259. Specify:		
			260. 1 □ yes 2 □ no	o 261. 1 □ yes 2 □ no
	262. In the Medical Dire	ector's judgement, was the ac	dverse event a direct re	esult of the infusion?
	2 □ no — →	263. Specify the most likely	y cause of the adverse	event:
		1 ☐ regimen related	•	
		2 ☐ product reaction		
		3 ☐ drug reaction		
		4 □ other illness →	264. Specify illness: _	
		5 ☐ other reason →	265. Specify reason:	
Donor / Infant Demog	•	estions 266–281) is to be o	completed for all non-	NMDP allogeneic donors.
		r or an autologous marrow		
266. (Female donor only) Wa	as the donor ever pregnar	nt?		
1 □ yes	267. Specify number of	pregnancies:	□ unknown	
4 □ not applicable / cord blood unit				
268. 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:			

CIBMTR Center Number:		CIBMTR Recipient ID:							
269. Did this donor have a ce  1 □ yes  2 □ no 3 □ not applicable, cord blood unit or marrow product	270. Specify the site of the central lin  1 ☐ femoral  2 ☐ subclavian  3 ☐ internal jugular	ne placement: fy site:							
272. Donor's ethnicity:  1 ☐ Hispanic or Latino 2 ☐ not Hispanic nor Lat 3 ☐ unknown  273. Donor's race: (Mark the	ino group(s) in which the donor is a mem	nber. Check all that apply.) 🕮A							
White	Black or African American	17 ☐ American Indian, South or Central America	Native Hawaiian or Other Pacific Islander						
<ul> <li>1 ☐ Eastern European</li> <li>2 ☐ Mediterranean</li> <li>3 ☐ Middle Eastern</li> <li>4 ☐ North Coast of Africa</li> <li>5 ☐ North American</li> <li>6 ☐ Northern European</li> </ul>	<ul> <li>11 ☐ African (both parents born in Africa)</li> <li>12 ☐ African American</li> <li>13 ☐ Black Caribbean</li> <li>14 ☐ Black South or Central American</li> </ul>	18 ☐ Caribbean Indian  Asian  19 ☐ South Asian 20 ☐ Filipino (Pilipino)	26 ☐ Guamanian 27 ☐ Hawaiian 28 ☐ Samoan 29 ☐ Other Pacific Islander						
7 ☐ Western European 8 ☐ White Caribbean 9 ☐ White South or Central American 10 ☐ Other White	American Indian or Alaska Native  15 □ Alaskan Native or Aleut 16 □ North American Indian	<ul> <li>21 ☐ Japanese</li> <li>22 ☐ Korean</li> <li>23 ☐ Chinese</li> <li>24 ☐ Vietnamese</li> <li>25 ☐ Other Southeast Asian</li> </ul>	Decline  30 □ Donor declines to provide race  31 □ Donor's race unknown						
274. What is the relationship  1 □ sibling 2 □ recipient's child	of the donor to the recipient?								
3 □ other relative → 4 □ unrelated	275. Specify the relationship of the of the parent  2 □ aunt  3 □ uncle  4 □ cousin  5 □ other  relative → 276. Specie	fy relationship:							
	tested for potentially transplantable g	enetic diseases?							
1 □ yes — → 2 □ no 3 □ unknown	Specify disease(s) tested:  278. 1 ☐ yes 2 ☐ no Sickle cell ar  279. 1 ☐ yes 2 ☐ no Thalassemia  280. 1 ☐ yes 2 ☐ no Other →	-							

		L	
		ogeneic non-NMDP donors. If the s blood unit, then continue with the s	
282. Was the donor hospitaliz  1 □ yes 2 □ no	ed (inpatient) during or a	fter the collection?	
283. Did the donor experience	e any life-threatening con	pplications during or after the collection	n?
1 □ yes	284. Specify complication	ns:	
285. Did the donor receive blo	ood transfusions as a res	ult of the collection?	
1 □ yes ————— 2 □ no	286. Was the blood tran	sfusion product autologous?	
	1 ☐ yes — ➤ 2 ☐ no	287. Specify number of units:	
	288. Was the blood tran	sfusion product allogeneic (homologo	us)?
	1 ☐ yes ———— 2 ☐ no	289. Specify number of units:	
290. Did the donor die as a re	esult of the collection?		
1 □ yes ————		eath:	
2 □ no L			
292. (Related donors only) Did 1 □ yes — ➤	d the recipient submit a r	esearch sample?	
2 □ no	293. Research sample r	ecipient ID:	
294. (Related donors only) Di	d the donor submit a rese	earch sample?	
1 □ yes	295. Research sample of	lonor ID:	
296. Signed:		Person completing form	
Please print name:		Person completing form	
•			
·	·		
F-mail address			

CIBMTR Recipient ID:

CIBMTR Center Number:



## **Confirmation of HLA Typing**

# Sequence Number: Date Received:

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 IDM Form 2004 and HSCT 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.

CIBMTR Center Number:	OMB No: 0915-0310 Expiration Date: 10/31/2010
CIBMTR Recipient ID:	
Specify non-NMDP donor:	
1 ☐ related donor ——➤	Donor's / infant's date of birth:
2 ☐ non-NMDP unrelated donor →	Month Day Year
3 ☐ non-NMDP cord blood unit — ➤ (include related and autologous CBUs)	Donor's / infant's gender:  1 ☐ male 2 ☐ female
	Non-NMDP unrelated donor / cord blood unit ID: (not applicable for related donor)
Today's Date: Month	Day Year
Date of HSCT for which th being completed:	is form is  Month Day  Year
HSCT type: ☐ allogene (check only one) unrelated	
Product type:	□ 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 sl	nould be completed for each non-NMDP donor, recipient, or cord blood unit.
<ol> <li>Specify the person for whom</li> <li>□ recipient — final typing</li> <li>□ recipient's mother —</li> </ol>	this typing is being done:
confirmatory typing ->	<ul> <li>2. Was the recipient's mother used as the donor?</li> <li>1 □ yes</li> <li>2 □ no</li> </ul>
3 ☐ recipient's father —	
confirmatory typing →	3. Was the recipient's father used as the donor?
4 ☐ donor — confirmatory	1 □ yes
typing	2 □ no
5 ☐ cord blood unit — confirmatory typing	
6 ☐ maternal HLA typing г	
7 □ other ———	4. Specify person and typing:

CIB	MTR (	Center Numb	er:						CIB	MTR Red	cipient ID:					
						,	_									
HL	А Ту	ping by D	NA Te	chnol	og	У										
alle	les in t		ove or be esult clar	elow the	e bo	ox for	that locus. orm review	. A lab r	report r	may be a	ttached to					
Cla	ss I															
	Locus									Allele	e Designation	าร				
6.	Α	□ not tested	First A	*												
			Second A	I .												
7.	В	□ not tested	First B	*												
			Second B	I											 	
8.	С	□ not													 	
		tested	First C	*											 	
			Second C												 	
Cla	SS II Locus									Allele	e Designation	าร				
9.	DRB1	□ not tested	Firs DRB1													
			Second DRB1													
			וטוטו												 	

CIBMTR Center Number:		CIBMTR Recipient ID	):					
		_				•		
Class II (Optional) Please provide the optional all	lele information if	it is available from your laboratory. Allele Designat	ions					
10. DRB3 ☐ not Fir	-st							
tested DRB				 			 	
Secor DRB:								
11. DRB4 ☐ not Fir								
tested DRB-	4							
Secon DRB								
12. DRB5 ☐ not Fir	rst T							
tested DRB								
Secor	nd							
DRB	5*							
13. DQB1 ☐ not Fir	-st							
tested DQB								
Secor	nd							
DQB								
14. DPB1 □ not Fir	ret							
tested DPB								
Secor	nd							
DPB								
15. DQA1 □ not Fir	ent							
15. DQA1 ☐ not Fir tested DQA								
Cooor	- d							
Secor DQA								
16. DPA1 □ not Fir	ret							
tested DPA								
0	- d							
Secor DPA								

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										
17. No. of ant 1 □ one		s pro □ tw								
Specificity	A 1st	ntige	n <u>2nd</u>							
A1		01								
A2		02								
A203		03								
A210		04								
A3		05								
A9		06								
A10		07								
A11		80								
A19		09								
A23(9)		10								
A24(9)		11								
A2403		12								
A25(10)		13								
A26(10)		14								
A28		15								
A29(19)		16								
A30(19)		17								
A31(19)		18 19								
A32(19) A33(19)		20								
A34(10)	Н	21	H							
A36	H	22								
A43		23	$\overline{\Box}$							
A66(10)	$\overline{\Box}$	24	_							
A68(28)		25								
A69(28)		26								
A74(19)		27								
A80` ´		28								
AX		99								

		B An	tigens		
		18. Number of a 1 □ one	ntigens provided: 2 ☐ two		
	Antigen		Antigen		Antigen
<b>Specificity</b>	<u>1st</u> <u>2nd</u>	<u>Specificity</u>	<u>1st</u>	<b>Specificity</b>	<u>1st</u> <u>2nd</u>
B5	□ 01 □	B40	□ 21 □	B59	
B7	□ 02 □	B4005	□ 22 □	B60(40)	□ 43 □
B703	□ 03 □	B41	□ 23 □	B61(40)	
B8	□ 04 □	B42	□ 24 □	B62(15)	□ 45 □
B12	□ 05 □	B44(12)	□ 25 □	B63(15)	□ 46 □
B13	□ 06 □	B45(12)	□ 26 □	B64(14)	□ 47 □
B14	□ 07 □	B46	□ 27 □	B65(14)	□ 48 □
B15		B47	□ 28 □	B67	
B16	□ 09 □	B48	□ 29 □	B70	□ 50 □
B17		B49(21)	□ 30 □	B71(70)	□ 51 □
B18		B50(21)	□ 31 □ □	B72(70)	□ 52 □
B21		B51(5)		B73	□ 53 □
B22	□ 13 □ □ 14 □	B5102	□ 33 □ □	B75(15)	□ 54 □
B27 B2708	□ 14 □ □ 59 □	B5103	□ 34 □ □ 35 □	B76(15)	□ 55 □ □ 56 □
	□ 59 □ □ 15 □	B52(5)		B77(15) B78	
B35 B37		B53		В76 В81	□ 57 □ □ 58 □
B38(16)		B54(22) B55(22)		B82	
B39(16)		B56(22)		BX	
B3901		B57(17)		DV	ц 99 ц
B3902		B58(17)			

C Antigens	DR Antigens  22. No. of antigens provided:  1 □ one 2 □ two		DR51 Antigen	DP Antigens  27. No. of antigens provided  1 □ one 2 □ two		
9. No. of antigens provided: 1 □ one 2 □ two			Specificity Present? 23. DR51 1 ☐ yes 2 ☐ no			
Antigen         Specificity       1st       2nd         Cw1       □       01       □         Cw2       □       02       □         Cw3       □       03       □         Cw4       □       04       □         Cw5       □       05       □         Cw6       □       06       □         Cw7       □       07       □	Antigen         Specificity       1st       2nd         DR1       □       01       □         DR103       □       02       □         DR2       □       03       □         DR3       □       04       □         DR4       □       05       □         DR5       □       06       □         DR6       □       07       □	Specificity 24. DR52  DR53  Specificity 25. DR53	2 Antigen Present? 1  yes 2 no 3 Antigen Present? 1  yes 2 no Antigens	Specificity DPw1 DPw2 DPw3 DPw4 DPw5 DPw6 DPX	Antigen       1st     2nd       □     01     □       □     02     □       □     03     □       □     04     □       □     05     □       □     06     □       □     99     □	
Cw8	DR7 DR8 DR9	□ 08 □ □ 09 □ □ 10 □ 10 □ 10 □ 10 □ 10 □ 10				
Bw Specificity Specificity Present? 20. Bw4 1 yes 2 no 21. Bw6 1 yes 2 no	DR10 DR11(5) DR12(5) DR13(6) DR14(6) DR1403 DR1404 DR15(2) DR16(2) DR17(3) DR18(3) DRX	□ 11 □ □ □ 12 □ □ 13 □ □ 14 □ □ 15 □ □ 16 □ □ 17 □ □ 18 □ □ 19 □ □ 20 □ □ 81 □ □ 99 □	Specificity DQ1 DQ2 DQ3 DQ4 DQ5(1) DQ6(1) DQ7(3) DQ8(3) DQ9(3) DQX	Antigen  1st 2nd  □ 01 □  □ 02 □  □ 03 □  □ 04 □  □ 05 □  □ 06 □  □ 07 □  □ 08 □  □ 09 □  □ 99 □		
·		Person cor	mpleting form			
Please print name:						
Phone: ()						
Fax: ()						

CIBMTR Recipient ID:

CIBMTR Center Number:



#### Specify non-NMDP donor: **Infectious Disease Markers** 1 ☐ related donor — Donor's / infant's date of birth: 2 ☐ non-NMDP **Registry Use Only** unrelated donor → Month Day Year 3 ☐ non-NMDP cord Donor's / infant's gender: Sequence blood unit -1 ☐ male Number: (include related and 2 female autologous CBUs) Non-NMDP unrelated donor / cord blood unit ID: (not applicable for related donor) Date Received: Public Burden Statement: An agency may not conduct or Today's Date: sponsor, and a person is not required to respond to, a collection of information unless it displays a currently valid OMB control Month Year number. The OMB control number for this project is 0915-0310. Date of HSCT for which this form is Public reporting burden for this collection of information, in 0 combination with the HLA Typing Form 2005 and HSCT Infusion being completed: 🕮 Month Day Year Form 2006, is estimated to average 1.5 hours per response, including the time for reviewing instructions, searching existing HSCT type: ☐ allogeneic, ☐ allogeneic, □ syngeneic data sources, and completing and reviewing the collection of (check only one) unrelated related (identical twin) information. Send comments regarding this burden estimate or any other aspect of this collection of information, including Product type: ☐ marrow ☐ PBSC ☐ cord blood □ other product, suggestions for reducing this burden, to HRSA Reports Clearance Officer, 5600 Fishers Lane, Room (check only one) specify: \_ 10-33, Rockville, Maryland, 20857. This form must be completed for all non-NMDP allogeneic or syngeneic donors, or non-NMDP cord blood units. If the donor or cord blood unit was secured through the NMDP, then report IDMs on forms 24 and 50 for allogeneic donors or through CORD Link for cord blood units. 1. Who is being tested for IDMs? 1 ☐ donor IDM (marrow or PBSC) 2 ☐ maternal IDM (cord blood) 3 ☐ cord blood unit IDM **Infectious Disease Marker** (report final test results) **Test Date** Month Day Year **Hepatitis B Virus (HBV)** 2. HBsAg: (hepatitis B surface antigen) 2 3. 1 ☐ reactive 2 ☐ non-reactive 3 ☐ testing not performed 4. Anti HBc: (hepatitis B core antibody) (no confirmatory test available) 2 0 5. 1 ☐ reactive 2 ☐ non-reactive 3 ☐ testing not performed **Hepatitis C Virus (HCV)** 2 6. Anti-HCV: (hepatitis C antibody) 7. 1 □ reactive 2 ☐ non-reactive

CIBMTR Center Number:

CIBMTR Recipient ID:

CIBMTR Form 2004 revision 2 (page 1 of 3) June 2009
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Internal use: Document number F00479 revision 2 Replaces: F00479 version 1.0 July 2007

3 ☐ testing not performed

Mail this form to your designated campus (Milwaukee or Minneapolis). Retain the original at the transplant center.

OMB No: 0915-0310

Expiration Date: 10/31/2010

CIBMTR Center Number:		CIBMTR Recipient ID:				
Infectious Disease Ma	arker (report final test result	ts)		Test D	ate	
Human T-Lymphotropic Vir  8. Anti-HTLV I / II:  1 □ reactive 2 □ non-reactive 3 □ testing not performe			9. Month	Day	Year 2 0	
Human Immunodeficiency  10. HIV-1 p24 antigen:  1 □ reactive  2 □ non-reactive  3 □ not reported  4 □ not performed; HIV	Virus (HIV)  NAT testing performed (skip)		11.		20	
12. Was FDA licensed NAT	testing for HIV-1 / HCV perfo	ormed?				
1 □ yes ————— 2 □ no	Specify results:  13. HIV-1  1 □ positive 2 □ negative 3 □ not reported	1	4.		20	
	15. HCV  1 ☐ positive 2 ☐ negative	1	6.		20	
•	nmunodeficiency Viruses) ies is required. This testing may be pe	1 erformed as separate tests or done using	8. a combined a	assay.	20	
Syphilis  19. STS:  1 ☐ reactive  2 ☐ non-reactive  3 ☐ testing not performe	ed	2	20.		20	
Cytomegalovirus (CMV)  21. Anti-CMV: (IgG or Total)  1 □ reactive  2 □ non-reactive  3 □ testing not performe		2	22.		20	

CIBMTR Center Number:		CIBMTR Recipient ID:				
Infectious Disease Ma	arker (report final test results)			Test D	ate	
West Nile Virus (WNV)  23. WNV-NAT testing:  1 □ positive 2 □ negative 3 □ testing not performe 4 □ not applicable	ed	24.	Month	Day	20	Year
25. Other infectious disease  1 □ yes → 2 □ no	marker, specify (e.g., EBV):  26. Specify date performed:  27. Specify test and method:  28. Specify test results:				20	
29. Other infectious disease  1 ☐ yes ———————————————————————————————————	marker, specify (e.g., EBV):  30. Specify date performed:  31. Specify test and method:  32. Specify test results:				20	<u>,                                    </u>
33. Other infectious disease  1 □ yes   2 □ no	marker, specify (e.g., EBV):  34. Specify date performed:  35. Specify test and method:  36. Specify test results:				20	
37. Signed:	Person (	completing form				
Please print name:	7 0/30/70					
Phone number: (	)					
·	•					
	)					
F-mail address:						