

Pre-Transplant Essential Data

CIBMTR Use Only	OMB No: 0915-0310 Expiration Date: 10/31/2022						
Sequence Number: Date Received:	Public Burden Statement: The purpose of the data collection is to fulfill the legislative mandate to establish and maintain a standardized database of allogeneic marrow and cord blood transplants performed in the United States or using a donor from the United States. The data collected also meets the C.W. Bill Young Cell Transplantation Program requirements to provide relevant scientific information not containing individually identifiable information available to the public in the form of summaries and data sets. An agency may not conduct or sponsor, and a person is not required to respond to, a collection of information collection is 0915-0310 and it is valid until 10/31/2022. This information collection is 0915-0310 and it is valid until 10/31/2022. This information collection is voluntary under The Stem Cell Therapeutic and Research Reauthorization Act of 2015, Public Law (Pub. L.) 109–129, as amended by the Stem Cell Therapeutic and Research Reauthorization Act of 2015, Public Law 111–264 (the Act) and the Stem Cell Therapeutic and Research Reauthorization Act of 2015, Public Law 114-104. Public reporting burden for this collection of information, seating 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. Research Reauthorization Act of 2015, Public Law 114-104. Public						
Center Identification CIBMTR Center Number:	paperwork@hrsa.gov.						
EBMT Code (CIC):							
CIBMTR Research ID (CRID):							

YYYY MM DD

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CIBMTR Center Number: ____ ___ ___ ___

CIBMTR Research ID: ____ ___ ___ ___ ___ ___ ___ ___ ___

Recipient Information

- 2. Sex
 - □ Male
 - □ Female
- 3. Ethnicity
 - □ Hispanic or Latino
 - Not Hispanic or Latino
 - □ Not applicable (not a resident of the USA)
 - □ Unknown
- 4. Race (check all that apply)
 - □ White Go to question 5
 - Black or African American– Go to question 5
 - Asian– Go to question 5
 - American Indian or Alaska Native– Go to question 5
 - □ Native Hawaiian or Other Pacific Islander– Go to question 5
 - □ Not reported *Go to question 6*
 - Unknown– Go to question 6
 - 5. Race detail (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
 - □ African American

CIBMTR Center Number: ____ ___ ___ ___

- 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
- □ Korean
- □ Chinese
- □ Vietnamese
- Other Southeast Asian
- □ Guamanian
- Hawaiian
- □ Samoan
- Other Pacific Islander
- □ Unknown
- 6. Country of primary residence
 - □ Afghanistan

GhanaGibraltar

□ Greece

□ Greenland

Guadeloupe

□ Grenada

Guam

□ Guatemala

□ Guernsey

□ Guinea-Bissau

Heard Island and McDonald

Guyana

Islands

□ Holy See

Haiti

□ Guinea

- Aland Islands
- Albania
- Algeria
- American Samoa
- □ Andorra
- Angola
- □ Anguilla
- Antarctica
- Antigua and Barbuda
- Argentina
- □ Armenia
- ☐ Aruba☐ Australia
- ☐ Australia☐ Austria

- Palau
- D Palestine, State of
- Panama
- Papua New Guinea
- Paraguay
- Peru
- Philippines
- Pitcairn Islands
- Poland
- Portugal
- Puerto Rico
- Qatar
- □ Reunion
- Romania
- Russia

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CIBMTR Center Number: ____ ___ ___

CIBMTR Research ID: ____ ___ ___ ___ ___ ___

Hong Kong

Hungary

Iceland

Indonesia

India

Iran

Ireland

Iraq

- Azerbaijan
- Bahamas
- Bahrain
- □ Bangladesh
- Barbados
- Belarus
- Belgium

- Benin

- Belize

Bermuda

Bhutan

Bolivia

Saba

Botswana

Bouvet Island

Bulgaria

Burundi

□ Cameroon

Chad

Chile

China

Colombia

Cambodia

Cape Verde

Cayman Islands

Central African Republic

Christmas Island

Cocos (Keeling) Islands

Burkina Faso

Bonaire, Sint Eustatius and

Bosnia and Herzegovina

Brazil - Go to question 7

British Virgin Islands

Brunei Darussalam

British Indian Ocean Territory

Canada - Go to question 8

- Isle of Man
 - Israel
 - Italv
 - Jamaica
 - Japan
 - Jersey
 - Jordan
 - Kazakhstan
 - Kenya
 - Kiribati
 - Kuwait
 - Kyrgyzstan
 - Laos
 - Latvia
 - Lebanon
 - Lesotho
 - Liberia
 - Libya
 - Liechtenstein
 - Lithuania
 - Luxembourg
 - Macau
 - Macedonia
 - Madagascar

Maldives

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Malawi

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Malaysia

- □ Honduras
- Rwanda
 - Saint Barthelemy
 - Saint Helena

Saint Lucia

п Saint Kitts and Nevis

□ Saint Martin, French

Grenadines

Samoa

San Marino

□ Saudi Arabia

Senegal

Serbia

Sevchelles

Sierra Leone

Sint Maarten, Dutch

Slovak Republic

Solomon Islands

Sandwich Islands

South Georgia and the South

Singapore

Slovenia

Somalia

South Korea

Spain

Sudan

Sri Lanka

Suriname

Swaziland

Switzerland

Sweden

Syria

Svalbard and Jan Mayen

South Sudan

South Africa

Saint Pierre and Miquelon

Saint Vincent and the

Sao Tome and Principe

CIBMTR Center Number: ____ ___ ___ ___

	Comoros	Mali		Taiwan
	Congo, Democratic Republic	Malta		Tajikistan
_	of the	Marshall Islands		Tanzania
	Congo, Republic of the	Martinique		Thailand
	Cook Islands	Mauritania		Timor-Leste
	Costa Rica	Mauritius		Togo
	Cote d'Ivoire	Mayotte		Tokelau
	Croatia	Mexico		Tonga
	Cuba	Micronesia		Trinidad and Tobago
	Curacao	Moldova		Tunisia
	Cyprus	Monaco		Turkey
	Czech Republic	Mongolia		Turkmenistan
	Denmark	Montenegro		Turks and Caicos Islands
	Djibouti	Montserrat		Tuvalu
	Dominica	Morocco		Uganda
	Dominican Republic	Mozambique		Ukraine
	Ecuador	Myanmar		United Arab Emirates
	Egypt	Namibia		United Kingdom (England,
	El Salvador	Nauru	_	Wales, Scotland, Northern
	Equatorial Guinea	Nepal		Ireland)
	Eritrea	Netherlands		United States - Go to question 9
	Estonia	Netherlands Antilles		United States Minor Outlying
	Ethiopia	New Caledonia		Islands
	Falkland Islands	New Zealand		United States Virgin Islands
	Faroe Islands	Nicaragua		Uruguay
	Fiji	Niger		Uzbekistan
	Finland	Nigeria		Vanuatu
	France	Niue		Venezuela
	French Guiana	Norfolk Island		Vietnam
	French Polynesia	North Korea		Wallis and Futuna Islands
	French Southern Territories			Western Sahara
	Gabon	Northern Mariana Islands		Yemen
	Gambia	Norway		Zambia
	Georgia	Oman		Zimbabwe
	Germany	Pakistan		
	,			

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CIBMTR Cer	nter Number:	CIBMTR Research ID:							
7.	State of residence of recipient (for 10	residents of Brazil)	Go to question						
	□ Acre	Maranhão	Rio de Janeiro						
	Alagoas	Mato Grosso	Rio Grande do Norte						
	Amapá	Mato Grosso do Sul	□ Rio Grande do Sul						
	Amazonas	Minas Gerais	Rondônia						
	Bahia	Pará	Roraima						
	Ceará	🗆 Paraíba	Santa Catarina						
	Distrito Federal	Paraná	São Paulo						
	Espírito Santo	□ Pernambuc	□ Sergipe						
	☐ Goiás	□ Piauí	□ Tocantins						
8.	Province or territory of residence of <i>question 10</i>	of recipient (for residents of Canada) _	Go to						
	Provinces		Territories						
	□ Alberta	Nova Scotia	Northwest Territories						
	British Columbia	Ontario	Nunavut						
	Manitoba	Prince Edward Island	□ Yukon						
	New Brunswick	□ Quebec							
	Newfoundland and Labrador	□ Saskatchewan							

9.	State of residence of recipient (for	residents of USA)	
	Alabama	□ Kentucky	North Dakota
	□ Alaska	🗆 Louisiana	🗆 Ohio
	Arizona	□ Maine	Oklahoma
	□ Arkansas	□ Maryland	Oregon
	California	□ Massachusetts	Pennsylvania
	□ Colorado	Michigan	□ Rhode Island
		Minnesota	South Carolina
	Delaware	Mississippi	South Dakota
	District of Columbia	□ Missouri	Tennessee

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CIBMTR Ce	nter Number:		CIBN	MTR Resear	ch ID:		
	Florida		⊐ Mo	Iontana			Texas
	Georgia	٢	⊐ Ne	lebraska			Utah
	🗆 Hawaii	٢	⊐ Ne	levada			Vermont
	🗆 Idaho	C	⊐ Ne	lew Hampshi	re		Virginia
	Illinois	C	⊐ Ne	lew Jersey			Washington
	🗆 Indiana	C	⊐ Ne	lew Mexico			West Virginia
	□ Iowa	C	⊐ Ne	lew York			Wisconsin
	□ Kansas	۵	⊐ No	lorth Carolina	1		Wyoming
	P Recipient ID (RID): postal code for place of recipient' 			ce (USA and	Canada recipients	s on	/y):
12. Specif □ A □ B □ AB □ O		logen	eic ł	HCTs only)			
•	fy Rh factor <i>(of recipient)</i> (For allo	ogene	ic H	ICTs only)			
	ositive egative						
14. Has th	ne recipient signed an IRB / ethics samples to the NMDP / CIBMTR? Yes (recipient consented) – Go to No (recipient declined) - Go to qu Not approached - Go to question Not applicable (center not particip	? (For o que uestic n 18 pating	allog estion on 18) - G	ogeneic HCT on 15 8 Go to questio	s only)	sen	t form to donate research
10.	YYY			 MM	 DD		
16.	Did the recipient submit a researce □ Yes – <i>Go to question 17</i>					ory?	(Related donors only)

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CIBN	ITR C	enter N	lumber: (CIBMTR Research ID:
			No – Go to question 18	
		17.	Research sample recipient ID: _	
18.	ls the data	•	ent participating in a clinical trial?	(clinical trial sponsors that use CIBMTR forms to capture outcomes
	ΠY	es - Go	to question 19	
	ΠN	o – Go	to question 24	
	19.	Stud	/ Sponsor	
			BMT CTN – Go to question 21	
			RCI BMT – Go to question 21	
			PIDTC – Go to question 21	
			USIDNET - Go to question 22	
			COG – Go to question 22	
			Other sponsor – Go to question	n 20
		20.	Specify other sponsor:	- Go to question 22
		21.	Study ID Number:	
		22.	Subject ID:	
		23.	Specify the ClinicalTrials.gov ide	lentification number: NCT

Copy questions 19-23 to report participation in more than one study.

Hematopoietic Cellular Transplant (HCT) and Cellular Therapy

- 24. Is a subsequent HCT planned as part of the overall treatment protocol? (not as a reaction to post-HCT disease assessment) (For autologous HCTs only)
 - □ Yes Go to question 25
 - □ No Go to question 26
 - 25. Specify subsequent HCT planned
 - □ Autologous
 - □ Allogeneic
- 26. Has the recipient ever had a prior HCT?
 - □ Yes Go to question 27
 - □ No Go to question 38

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- 27. Specify the number of prior HCTs: _____
- 28. Were all prior HCTs reported to the CIBMTR?
 - □ Yes Go to question 33
 - □ No Go to question 29
 - Unknown Go to question 33

Copy and complete questions 29- 32 to report all prior HCTs that have not yet been reported to the CIBMTR

- 30. Was the prior HCT performed at a different institution?
 - □ Yes Go to question 31
 - □ No Go to question 32

Specify the institution that performed the last HCT

City:	 	
State:	 	
Country:		

32. What was the HPC source for the prior HCT? (check all that apply)

- □ Autologous
- □ Allogeneic, unrelated
- □ Allogeneic, related

33. Reason for current HCT

- Graft failure / insufficient hematopoietic recovery Go to question 34
- Persistent primary disease- Go to question 38
- □ Recurrent primary disease- *Go to question 35*
- Planned subsequent HCT, per protocol– Go to question 38
- New malignancy (including PTLD and EBV lymphoma) **Go to question 36**
- Insufficient chimerism– Go to question 38
- Other- Go to question 37

CIBMTR Center Number:		:	CIE	BMTR Resea	rch ID: _									
						YYYY	M	М	DD					
		35.	Date	e of relapse: _				_ – G	o to quest	tion 38				
					YYYY	MM	DD							
		36.	Date	e of secondary	y malignancy:			·		– Go to question 38				
						YYYY	Ν	ИM	DD					
		37.	Spe	cify other reas	son:		·	Got	to questio	n 38				
38.	Has	the rec	cipient e	ever had a pri	or cellular therap	by? (do not ind	clude DL	ls)						
	ΠY	/es – G	Go to q	uestion 39										
		10 – G	o to qu	estion 44										
		Unkno	wn– G	o to questio	n 44									
	39.	Were	e all pri	or cellular the	rapies reported t	the CIBMT	સ ?							
			Yes -	- Go to quest	tion 44									
			No –	Go to questi	on 40									
			Unkn	own– Go to c	question 44									
	Copy and comp reported to the		o the CIBMT				ular tl	herapies t	hat have not yet been					
		40.	Date	, or the phore	, cilular tricrapy. <u>-</u>	YYYY	MM	DD	_					
		41.	Was	the cellular t	herapy performe	d at a differer	nt institut	ion?						
				Yes – Go t e	o question 42									
				No – Go to	question 43									
			42.	Name:										
				City:										
				State:										
	Countr			Country: _										
		43.	Spe	cify the source	ne source(s) for the prior cellular therapy (check all that apply)									
				Autologous										
				Allogeneic,	unrelated									
				Allogeneic,	related									

CIBMTR Center Number: ____ ___ ___ ___ ___

Donor Information

- 44. Multiple donors?
 - □ Yes Go to question 45
 - □ No Go to question 46
 - 45. Specify number of donors: _____

To report more than one donor, copy questions 46-82 and complete for each donor.

- 46. Specify donor
 - □ Autologous
 - □ Allogeneic, related
 - □ Allogeneic, unrelated
- 47. Specify product type (check all that apply)
 - □ Bone marrow
 - □ PBSC
 - □ Single cord blood unit
 - Other product- Go to question 48
 - 48. Specify other product: _____
- 49. Is the product genetically modified? *If autologous, go to question 77. If allogeneic related, go to question 50. If allogeneic unrelated, go to question 54.*
 - □ Yes
 - □ No
 - 50. Specify the related donor type
 - Syngeneic (monozygotic twin) Go to question 55
 - HLA-identical sibling (may include non-monozygotic twin) Go to question 55
 - HLA-matched other relative (does NOT include a haplo-identical donor) Go to question 51
 - HLA-mismatched relative- Go to question 51
 - 51. Specify the biological relationship of the donor to the recipient
 - □ Mother
 - □ Father
 - □ Child
 - □ Sibling

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- □ Fraternal twin
- Maternal aunt
- □ Maternal uncle
- Maternal cousin
- Paternal aunt
- Paternal uncle
- Paternal cousin
- □ Grandparent
- Grandchild
- □ Other biological relative *Go to question 52*
- 52. Specify other biological relative: _______ Go to question 53
- 53. Degree of mismatch (related donors only)
 - HLA-mismatched 1 allele– *Go to question 55*
 - □ HLA-mismatched ≥2 alleles (does include haplo-identical donor) Go to question 55
- 54. Specify unrelated donor type
 - HLA matched unrelated
 - □ HLA mismatched unrelated
- 55. Did NMDP / Be the Match facilitate the procurement, collection, or transportation of the product?
 - □ Yes
 - □ No
- 56. Was this donor used for any prior HCTs? (for this recipient)
 - □ Yes
 - □ No

58. Registry donor ID: (not applicable for related donors)

 	 - Go to (question	63						

59. Non-NMDP cord blood unit ID: (include related and autologous CBUs)

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CIBM	ITR Ce	enter N	umber: CIBMTR Research ID:
	60.	Glob	al Registration Identifier for Donors (GRID):
		NML	DP donor, go to question 72
		Non	-NMDP unrelated donor, go to question 63
	61.	Is the	CBU ID also the ISBT DIN number?
			Yes – Go to question 63
			No – Go to question 62
			Unknown– Go to question 63
		62.	Specify the ISBT DIN number:
	63.	Regis	stry or UCB Bank ID: If 'Other registry' go to 64, otherwise go to question 65
		64.	Specify other Registry or UCB Bank: Go to question 65
	65.	Dono	r date of birth
			Known – Go to question 66
			Unknown – Go to question 67
		66.	Donor date of birth:
			YYYY MM DD
		67.	Donor age
			□ Known – Go to question 68
			Unknown – Go to question 69
			68. Donor age:
			□ Years
	69.	Done	Dr sex
			Male
			Female
70.	Spec	ify bloc	od type (donor) (non-NMDP allogeneic donors only)
		А	
		В	
		AB	
		0	

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71. Specify Rh factor (donor) (non-NMDP allogeneic donors only)

- Positive
- Negative

72. Donor CMV-antibodies (IgG or Total) (Allogeneic HCTs only)

- Reactive
- Non-reactive
- Indeterminate
- Not done
- Not applicable (cord blood unit)
- Has the donor signed an IRB / ethics committee (or similar body) approved consent form to donate research blood 73. samples to the NMDP / CIBMTR? (Related donors only)
 - Yes (donor consented) - Go to question 74
 - No (donor declined) - Go to guestion 77
 - Not approached - Go to question 77
 - Not applicable (center not participating) - Go to question 77
 - 74. Date form was signed: ____ ___ ___ ___ ____ YYYY MM DD
 - 75. Did the donor submit a research sample to the NMDP/CIBMTR repository? (Related donors only)
 - Yes - Go to question 76
 - No - Go to question 77
 - 76.
- Specify number of products infused from this donor: _____ 77.
- 78. Specify the number of these products intended to achieve hematopoietic engraftment: _____

Questions 79-80 are for autologous HCT recipients only.

- What agents were used to mobilize the autologous recipient for this HCT? (check all that apply) 79.
 - G-CSF (filgrastim, Neupogen)
 - Pegylated G-CSF (pegfilgrastim, Neulasta)
 - Plerixafor (Mozobil)
 - Combined with chemotherapy

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CIBMTR Center Number: ____ ___ CIBMTR Research ID: ____ ___ ___ ___ ___ ___ ___ ___

- □ Anti-CD20 (rituximab, Rituxan)
- □ Other agent- *Go to question 80*
- 80. Specify other agent: _____
- 81. Name of product: (gene therapy recipients)
 - □ Other name
 - 82. Specify other name: _____

To report more than one donor, copy questions 46-82 and complete for each donor.

Clinical Status of Recipient Prior to the Preparative Regimen (Conditioning)

83. What scale was used to determine the recipient's functional status?

- □ Karnofsky (recipient age \geq 16 years) Go to question 84
- □ Lansky (recipient age \geq 1 year and < 16 years) Go to question 85

Performance score prior to the preparative regimen:

- 84. Karnofsky Scale (recipient age \geq 16 years)
 - 100 Normal; no complaints; no evidence of disease Go to question 86
 - 90 Able to carry on normal activity Go to question 86
 - 80 Normal activity with effort *Go to question 86*
 - 70 Cares for self; unable to carry on normal activity or to do active work *Go to question 86*
 - 60 Requires occasional assistance but is able to care for most needs Go to question 86
 - 50 Requires considerable assistance and frequent medical care Go to question 86
 - D 40 Disabled; requires special care and assistance Go to question 86
 - 30 Severely disabled; hospitalization indicated, although death not imminent Go to question 86
 - 20 Very sick; hospitalization necessary *Go to question 86*
 - D 10 Moribund; fatal process progressing rapidly *Go to question 86*
- 85. Lansky Scale (recipient age \geq 1 year and < 16 years)
 - □ 100 Fully active
 - 90 Minor restriction in physically strenuous play
 - 80 Restricted in strenuous play, tires more easily, otherwise active
 - **70** Both greater restrictions of, and less time spent in, active play

CIBMTR Center Number: _____ CIBMTR Research ID: ____ ___ ___

- 60 Ambulatory up to 50% of time, limited active play with assistance / supervision
- 50 Considerable assistance required for any active play; fully able to engage in quiet play
- □ 40 Able to initiate quiet activities
- □ 30 Needs considerable assistance for quiet activity
- 20 Limited to very passive activity initiated by others (e.g., TV)
- 10 Completely disabled, not even passive play
- 86. Recipient CMV-antibodies (IgG or Total)
 - □ Reactive
 - Non-reactive
 - □ Indeterminate
 - □ Not done

Comorbid Conditions

- 87. Has the patient been infected with COVID-19 (SARS-CoV-2) based on a positive test result at any time prior to the start of the preparative regimen / infusion?
 - □ Yes Go to question 88
 - □ No Go to question 90
 - 88. Did the patient require hospitalization for management of COVID-19 (SARS-CoV-2) infection?
 - □ Yes Go to question 89
 - □ No Go to question 90
 - 89. Was mechanical ventilation used for COVID-19 (SARS-CoV-2) infection?
 - □ Yes
 - □ No
- 90. Is there a history of mechanical ventilation (excluding COVID-19 (SARS-CoV-2))?
 - □ Yes
 - □ No
- 91. Is there a history of invasive fungal infection?
 - □ Yes
 - □ No
- 92. Glomerular filtration rate (GFR) before start of preparative regimen (pediatric only)
 - □ Known- *Go to question* 93
 - Unknown- *Go to question 94*

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- 93. Glomerular filtration rate (GFR): ____ mL/min/1.73²
- 94. Does the recipient have known complex congenital heart disease? (corrected or uncorrected) (excluding simple ASD, VSD, or PDA repair) (pediatric only)
 - □ Yes
 - □ No
- 95. Were there any co-existing diseases or organ impairment present according to the HCT comorbidity index (HCT-CI)? (Source: Sorror, M. L. (2013). How I assess comorbidities before hematopoietic cell transplantation. Blood, 121(15), 2854-2863.)
 - Yes- Go to question 96
 - □ No- Go to question 102
 - 96. Specify co-existing diseases or organ impairment (check all that apply)
 - Arrhythmia Any history of atrial fibrillation or flutter, sick sinus syndrome, or ventricular arrhythmias requiring treatment
 - Cardiac -Any history of coronary artery disease (one or more vessel-coronary artery stenosis requiring medical treatment, stent, or bypass graft), congestive heart failure, myocardial infarction, OR ejection fraction ≤ 50% on the most recent test
 - Cerebrovascular disease -Any history of transient ischemic attack, subarachnoid hemorrhage or cerebral thrombosis, embolism, or hemorrhage
 - Diabetes -Requiring treatment with insulin or oral hypoglycemic drugs in the last 4 weeks but not diet alone
 - Heart valve disease -At least a moderate to severe degree of valve stenosis or insufficiency as determined by Echo; prosthetic mitral or aortic valve; or symptomatic mitral valve prolapse
 - Hepatic, mild Bilirubin > upper limit of normal to 1.5 x upper limit of normal, or AST/ALT > upper limit of normal to 2.5 x upper limit of normal at the time of transplant OR any history of hepatitis B or hepatitis C infection
 - Hepatic, moderate/severe -Liver cirrhosis, bilirubin > 1.5 × upper limit of normal, or AST/ALT > 2.5 × upper limit of normal
 - Infection -Includes a documented infection, fever of unknown origin, or pulmonary nodules suspicious for fungal pneumonia or a positive PPD test requiring prophylaxis against tuberculosis. Patients must have started antimicrobial treatment before Day 0 with continuation of antimicrobial treatment after Day 0
 - Inflammatory bowel disease -Any history of Crohn's disease or ulcerative colitis requiring treatment
 - Obesity -Patients older than 18 years with a body mass index (BMI) > 35 kg/m2 prior to the start of conditioning or a BMI of the 95th percentile of higher for patients aged 18 years or younger
 - Peptic ulcer -Any history of peptic (gastric or duodenal) ulcer confirmed by endoscopy or radiologic diagnosis requiring treatment
 - Psychiatric disturbance -Presence of any mood (e.g., depression), anxiety, or other psychiatric disorder (e.g. bipolar disorder or schizophrenia) requiring continuous treatment in the last 4 weeks

- Pulmonary, moderate -Corrected diffusion capacity of carbon monoxide and/or FEV1 of 66-80% or dyspnea on slight activity attributed to pulmonary disease at transplant
- □ Pulmonary, severe -Corrected diffusion capacity of carbon monoxide and/or FEV1 of ≤ 65% or dyspnea at rest attributed to pulmonary disease or the need for intermittent or continuous oxygen during the 4 weeks prior to transplant
- □ Renal, moderate / severe -Serum creatinine > 2 mg/dL or > 177 µmol/L; on dialysis during the 4 weeks prior to transplant; OR prior renal transplantation -go to question 97
- Rheumatologic -Any history of a rheumatologic disease (e.g., systemic lupus erythematosis, rheumatoid arthritis, polymyositis, mixed connective tissue disease, or polymyalgia rheumatica, etc.) requiring treatment. (Do NOT include degenerative joint disease, osteoarthritis)
- Prior malignancy-Treated at any time point in the patient's past history, other than the primary disease for which this infusion is being performed -go to question 98
- 97. Was the recipient on dialysis immediately prior to start of preparative regimen?
 - □ Yes
 - □ No
 - □ Unknown
- 98. Specify prior malignancy (check all that apply)
 - Breast cancer
 - Central nervous system (CNS) malignancy (e.g., glioblastoma, astrocytoma)
 - Gastrointestinal malignancy (e.g., colon, rectum, stomach, pancreas, intestine, esophageal)
 - Genitourinary malignancy (e.g., kidney, bladder, ovary, testicle, genitalia, uterus, cervix, prostate)
 - Leukemia (includes acute or chronic leukemia)
 - □ Lung cancer
 - Lymphoma (includes Hodgkin & non-Hodgkin lymphoma)
 - □ MDS / MPN
 - Melanoma
 - □ Multiple myeloma / plasma cell disorder (PCD)
 - □ Oropharyngeal cancer (e.g., tongue, buccal mucosa)
 - □ Sarcoma
 - □ Thyroid cancer
 - Other skin malignancy (basal cell, squamous)- go to question 99
 - Other hematologic malignancy -go to question 100
 - Other solid tumor -go to question 101
 - 99. Specify other skin malignancy: (prior)

100. Specify other hematologic malignancy: (prior) ____

CIBN	ITR Ce	ter Number: CIBMTR Research ID:
		101. Specify other solid tumor: (prior)
close	est to t	within 4 weeks prior to the start of the preparative regimen, report results from the test performed the start date. Biomarkers according to the augmented HCT comorbidity index. (Source: Biol Blood plant. 2015 Aug; 21(8): 1418–1424)
102.	Seru	ferritin (within 4 weeks prior to the start of the preparative regimen, use result closest to the start date)
		Known – Go to question 103
		Unknown – <i>Go to question 106</i>
	103.	ng/mL (μg/L)
	104.	Date sample collected:
		YYYY MM DD
	105.	Upper limit of normal for your institution:
106.	Seru	albumin (within 4 weeks prior to the start of the preparative regimen, use result closest to the start date)
		Known – Go to question 107
		Unknown – Go to question 109
	107.	• □ g/dL
		□ g/L
	108.	Date sample collected:
		YYYY MM DD
109.	Plate	ts (within 4 weeks prior to the start of the preparative regimen, use result closest to the start date)
		Known – Go to question 110
		Unknown – <i>Go to question 112</i>
	110.	□ □ x 10 ⁹ /L (x 10 ³ /mm ³)
		$\Box \times 10^{6}/L$
	111.	Were platelets transfused \leq 7 days before date of test?
		□ Yes

- 112. Did the recipient have a prior solid organ transplant?
 - □ Yes- Go to question 113
 - □ No- Go to question 116
 - 113. Specify organ:
 - □ Bowel
 - □ Heart
 - □ Kidney(s)
 - □ Liver
 - □ Lung(s)
 - Pancreas
 - Other organ- *Go to question 114*
 - 114. Specify other organ: _____
 - 115. Year of prior solid organ transplant: ____ ___ ___
 - YYYY

Copy and complete questions 113-115 for each prior solid organ transplant

Pre-HCT Preparative Regimen (Conditioning)						
116.	Heigh	nt at ini	itiation of pre-HCT preparative regimen:	□ inches		
117.	17. Actual weight at initiation of pre-HCT preparative regimen: D pounds					
				□ kilograms		
118.	Was a pre-HCT preparative regimen prescribed?					
		Yes – Go to question 119				
	□ No – Go to question 132					
	 119. Classify the recipient's prescribed preparative regimen (Allogeneic HCTs only) Myeloablative Non-myeloablative (NST) 					
			Reduced intensity (RIC)			

- 120. Was irradiation planned as part of the pre-HCT preparative regimen?
 - □ Yes Go to question 121
 - □ No Go to question 126

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CIBMTR Center Nu	mber: CIBMTR Research ID:		
121.	What was the prescribed radiation field?		
	Total body – Go to question 122		
	Total body by intensity-modulated radiation therapy (IMRT) – Go to question 122		
	Total lymphoid or nodal regions – Go to question 122		
	Thoracoabdominal region – Go to question 122		
122.	Total prescribed dose: (dose per fraction x total number of fractions) Gy		
123.	Date started:		

- 124. Was the radiation fractionated?
 - □ Yes Go to question 125
 - □ No Go to question 126
 - 125. Total number of fractions: _____

Indicate the total prescribed cumulative dose for the preparative regimen

126. Drug (drop down list)

- Bendamustine
- □ Busulfan
- Carboplatin
- □ Carmustine (BCNU)
- CCNU (Lomustine)
- □ Clofarabine (Clolar)
- □ Cyclophosphamide (Cytoxan)
- □ Cytarabine (Ara-C)
- Etoposide (VP-16, VePesid)
- □ Fludarabine
- □ Gemcitabine
- □ Ibritumomab tiuxetan (Zevalin)
- □ Ifosfamide
- Melphalan (L-Pam)
- □ Methylprednisolone (Solu-Medrol)
- □ Pentostatin
- Propylene glycol-free melphalan (Evomela)

CIBMTR Center Number:			umber:	CIBI	MTR Research ID:
			Rituximab (Rituxan)		
			Thiotepa		
			Tositumomab (Bexxar)		
			Treosulfan		
			Other drug -go to question 1	27	
		127	Specify other drug:		
	400				
	128.	lotal	prescribed dose:	_·	□ mg/m²
					□ mg/kg
					□ AUC (mg x h/L)
					□ AUC (µmol x min/L)
					□CSS (ng/mL)
	129.	Date s	started:		
			YYYY	MM	DD
130. Specify administration <i>(busulfan only</i> □ Oral)			
			IV		
			Both		
	Сор	y and	complete question 126-130 to	repo	ort each drug given for the preparative regimen
Addit	ional	Drugs	Given in the Peri-Transplant	Perio	d
101					
131.			ATG, ATS • Go to question 132		
			Go to question 135		
		NO - 1	Go to question 135		
	132.	Total	prescribed dose:		mg/kg
	133.	Speci	fy source		
			ATGAM (horse) – Go to ques	tion 1	135
	ATG – Fresenius (rabbit) – Go to question 135				uestion 135
			Thymoglobulin (rabbit) - Go to	ques	stion 135
			Other - Go to question 134		
		134.	Specify other source:		

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CIBMTR Center Number:			CIBMTR Research ID:
135.	Alem	tuzumab (Campath)	
	Yes – Go to question 136		
		No – Go to question 137	
	136.	Total prescribed dose:	🗆 mg/m2
			□ mg/kg
			□mg
137.	Defibrotide		
		Yes	
		No	
138.	138. KGF		
		Yes	
		No	
139.	9. Ursodiol		
		Yes	
		No	

GVHD Prophylaxis

This section is to be completed for allogeneic HCTs only; autologous HCTs continue with question 143.

140. Was GVHD prophylaxis planned?

- Yes Go to question 141
- □ No Go to question 143
- 141. Specify drugs / intervention (check all that apply)
 - □ Abatacept
 - Anti CD 25 (Zenapax, Daclizumab, AntiTAC)
 - Blinded randomized trial
 - □ Bortezomib
 - □ CD34 enriched (CD34+ selection)
 - □ Corticosteroids (systemic)
 - **Cyclophosphamide (Cytoxan)**

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CIBMTR Center Number: ____ ___ ___ ___ ___

CIBMTR Research ID: ____ ___ ___ ___ ___ ___ ___ ___ ____

- Cyclosporine (CSA, Neoral, Sandimmune)
- □ Extra-corporeal photopheresis (ECP)
- Ex-vivo T-cell depletion
- □ Filgotinib
- □ Maraviroc
- □ Methotrexate (MTX) (Amethopterin)
- □ Mycophenolate mofetil (MMF) (CellCept)
- □ Ruxolotinib
- □ Sirolimus (Rapamycin, Rapamune)
- □ Tacrolimus (FK 506)
- Tocilizumab
- □ Other agent-go to question 142
- 142. Specify other agent: _____ (do not report ATG, campath)

Post-HCT Disease Therapy Planned as of Day 0

- 143. Is additional post-HCT therapy planned?
 - □ Yes Go to question 144
 - □ No Go to First Name

Questions 144-145 are optional for non-U.S. centers

- 144. Specify post-HCT therapy planned (check all that apply)
 - □ Azacytidine (Vidaza)
 - □ Blinatumomab
 - Bortezomib (Velcade)
 - Bosutinib
 - Brentuximab
 - Carfilzomib
 - Cellular therapy (e.g. DCI, DLI)
 - Crenolanib
 - Daratumumab
 - Dasatinib
 - Decitabine
 - Elotuzumab
 - □ Enasidenib

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CIBMTR Center Number: ____ ___ ___ ___ ___

- □ Gilteritinib
- □ Ibrutinib
- □ Imatinib mesylate (Gleevec, Glivec)
- □ Intrathecal therapy (chemotherapy)
- □ Ivosidenib
- Ixazomib
- □ Lenalidomide (Revlimid)
- □ Lestaurtinib
- □ Local radiotherapy
- □ Midostaurin
- □ Nilotinib
- Obinutuzumab
- Pacritinib
- Ponatinib
- D Quizartinib
- Rituximab (Rituxan, MabThera)
- □ Sorafenib
- □ Sunitinib
- □ Thalidomide (Thalomid)
- Other therapy- *Go to question 145*
- □ Unknown
- 145. Specify other therapy: _____

Prior Exposure: Potential Study Eligibility

Selecting any option(s) below may generate an additional supplemental form.

- 146. Specify if the recipient received any of the following (at any time prior to HCT / infusion) (check all that apply)
 - Blinatumomab (Blincyto)
 - Gemtuzumab ozogamicin (Mylotarg)
 - □ Inotuzumab ozogamicin (Besponsa)
 - □ Adienne Tepadina[®]
 - □ Mogamulizumab (Poteligeo)
 - □ None of the above

CIBMTR Center Number:			CIBMTR Research ID:
First Name:			
Last Name: _			
E-mail address: _			
Date:			
	YYYY	MM	DD