

Pre-Transplant Essential Data

CIB MTR	OMB No: 0915-0
Use Only	Expiration Date: 10/31/2022
Sequ ence Num ber:	Public Burden Statement: Public Burden Statement: Public Burden Statemen The purpose of th data collection is I fulfill the legislativ mandate to establ and maintain a standardized data of allogeneic mar and cord blood transplants perfor in the United Stat using a donor fror
Date Rece ived:	United States. The data collected als

Center Identification	
CIBMTR Center Number:	
EBMT Code (CIC):	
Recipient Identification	
CIBMTR Research ID (CRID):	
Event date: / /	
	_YYYYMM DD

					Recipie
1.	Date	of birth			
			YYYY	MM	DD
2.	Sex				
		Male			
		Fema	ale		
3.	Ethn				
			anic or Latino		
			Hispanic or Latino	aident of the	UCA
		Unkn	applicable <i>(not a re</i>	Siderit of the	USA)
		Ulikii	lowii		
4.	Race	e (chec	k all that apply)		
		White	e – Go to question	າ 5.	
		Black	or African Americ	an– Go to q	uestion 5.
		Asiar	n– Go to question	5.	
		Amer	rican Indian or Alas	ska Native– (Go to ques
		Nativ	e Hawaiian or Oth	er Pacific Isl	ander– Go
		Not re	eported – Go to q	uestion 6.	
		Unkn	own– Go to ques	tion 6.	
	5.	Paco	detail (check all th	at apply)	
	5.		Eastern Europea		
			Mediterranean	"	
			Middle Eastern		
			North Coast of A	frica	
			North American	поа	
			Northern Europe	an	
			Western Europe		
			White Caribbean		
		_	Janostan		

CIBMTR Research ID: ___ __ __ __ __

11.	Zip o	r posta	al code for place of recipient's residence (USA recipients only):	
10.	NMD	P Rec	ipient ID (RID):	
	9.	Sta	ate of residence of recipient (for residents of USA)	
	8.		ovince or territory of residence of recipient (for residents of Canada)estion 10.	Go to
	7.	Sta	ate of residence of recipient (for residents of Brazil)	$_$ - Go to question 10.
	6.		Country of primary residence	
			OHKHOWH	
			Unknown	
			Samoan Other Pacific Islander	
			Hawaiian	
			Guamanian	
			Other Southeast Asian	
			Vietnamese	
			Chinese	
			Korean	*
			Japanese	
			Filipino (Pilipino)	
			South Asian	
			Caribbean Indian	
			American Indian, South or Central America	
			North American Indian	
			Alaskan Native or Aleut	
			Other Black	
			Black South or Central American	
			Black Caribbean	
			African American	
			African	
			Other White	
			White South or Central American	

(last 4 digits optional)

CIBMTR Research ID: ______

CIBN	ITR C	enter l	Number:	CIBMTR Research ID:			
12.	Spe	cify blo	ood type (of recipient)	(For allogeneic HCTs only)			
		4					
	E	3					
		AΒ					
)					
13.	Spec	cify Rh	factor (of recipient)	(For allogeneic HCTs only)			
	F	ositiv	е				
	1	Negativ	ve				
14.			cipient signed an IRE DP / CIBMTR?	3 / ethics committee (or similar body) approved consent form for submitting research data			
		Yes	(recipient consented) – Go to question 15.			
		No (recipient declined) –	Go to question 17.			
		Not	approached – Go to	question 17.			
		15. futu	ire research?	Did the recipient give permission to be directly contacted by CIBMTR for			
			Yes (recipient prov	vided permission) – Go to question 16.			
			No (recipient decli	ned) – Go to question 17.			
		16.		Date form was signed:			
				YYYY MM DD			
17.			cipient signed an IRE the NMDP / CIBMT	3 / ethics committee (or similar body) approved consent form to donate research blood R?			
		Yes	(recipient consented	/) – Go to question 18.			
		No (recipient declined) -	Go to question 21.			
		Not approached - Go to question 21.					
		Not a	applicable (center no	ot participating) - Go to question 21.			
	18.	Date	e form was signed: _				
				YYYY MM DD			
	19.	Did t	the recipient submit a	a research sample to the NMDP/CIBMTR repository? (Related donors only)			
			Yes – Go to ques	tion 20.			
			No – Go to quest	ion 21.			
			20.	Research sample recipient ID:			

CIBN	CIBMTR Center Number:			CIBMTR Research ID:		
21.	Is the	e recip	ient participating in a clinical trial	? (clinical trial sponsors that use CIBMTR forms to capture outcom	es data)	
	□ Ye	s - G o	to question 22.			
	□ No	– Go	to question 26.			
	22.	Stud	ly Sponsor			
			BMT CTN - Go to question 2	4.		
			RCI BMT - Go to question 24	ı.		
			PIDTC - Go to question 24.			
			5.			
			COG - Go to question 25.			
			Other sponsor – Go to question	on 23.		
			23. question 25.	Specify other sponsor:	Go to	
			24.	Study ID Number:		
			25.	Subject ID:		
	Сору	ques	tions 2225. to report participa	ation in more than one study.		
				Hematopoietic Cellular Transplant (HCT) and Cellular Therap	ру	
26.			quent HCT planned as part of the nt) (For autologous HCTs only)	e overall treatment protocol? (not as a reaction to post-HCT disease	è	
		Yes	– Go to question 27.			
		No -	- Go to question 28.			
	27.	Spe	cify subsequent HCT planned			
			Autologous			
			Allogeneic			
28.	Has	the re	cipient ever had a prior HCT?			
	□ \	′es – (Go to question 29.			
	<u> </u>	10 – G	o to question 40.			
	29.	Spe	cify the number of prior HCTs:			
	30.	Wer	e all prior HCTs reported to the C	CIBMTR?		

SMIRC	enter i	Number	:	CIBMTR Research ID:	_
			- Go to question 35.		
			Go to question 31.		
		Unkn	own – Go to question	31.	
				Copy and complete questions 31 34. to report all price that have not yet been reported to the CIBMTR	or HC1
			3	31. Date of the prior HCT:	⊐ date
			YYYY	MM DD	
			3	32. Was the prior HCT performed at a different institution?	
			Yes – Go to questio	n 33.	
			No – Go to question	34.	
	Spe	ecify th	e institution that perfo	ormed the last HCT	
			33.	Name:	
				City:	
				State:	
			Country:		
		34.		What was the HPC source for the prior HCT? (check all that apply	v)
			Autologous	What was the Three source for the phor the T. (officer all that appro	,,
			Allogeneic, unrelated		
			Allogeneic, related		
35.	Rea	son for	current HCT		
		Graft	failure / insufficient hen	natopoietic recovery – <i>Go to question 36.</i>	
			stent primary disease–	·	
			rrent primary disease-	·	
			·	er protocol– Go to question 40.	
	_			PTLD and EBV lymphoma) – Go to question 38.	
			ficient chimerism– Go to	o question 40.	
		Othe	– Go to question 39.		

CIBM	ITR C	enter N	Number:		CIBMTR Research ID:			
			36. Go t a	o question 40.	Date of graft failure / rejection:			
				•	YYYY MM DD			
			37. ques	stion 40.	Date of relapse:			
				YYYY	MM DD			
			38. Go t e	o question 40.	Date of secondary malignancy:			
					YYYY MM DD			
			39. 40.		Specify other reason: Go to question			
40.	Has	the red	cipient e	ver had a prior cellular	therapy? (do not include DLIs)			
	□ Y	'es – C	o to qu	uestion 41.				
	☐ No – Go to question 46.			estion 46.				
	□□Unknown– Go to question 46.			to question 46.				
	41.	Were	e all prio	or cellular therapies rep	ported to the CIBMTR?			
			Yes –	Go to question 46.				
			No – (Go to question 42.				
			Unkno	own– Go to question A	46.			
		-	_	complete questions 4 the CIBMTR	1245. to report all prior cellular therapies that have not yet been			
			42.		Date of the prior cellular therapy: YYYY MM DD			
			43.		Was the cellular therapy performed at a different institution?			
				Yes – Go to questio				
				No – Go to questio				
				44.	Name:			
				City:				
				State:				
				Country:				

CIBM	ITR Ce	enter Number:	CIBMTR Research ID:		
		45. □ Autologous □ Allogeneic, unrelated	Specify the source(s) for the prior cellular therapy (check all that apply)		
		☐ Allogeneic, related			
			Donor Information		
46.	Multip	ole donors?			
		Yes – Go to question 47.			
		No - Go to question 48.			
	47.	Specify number of donors:			
To re	port n	nore than one donor, copy question	s 4883. and complete for each donor.		
48.	Spec	ify donor			
		Autologous			
		Allogeneic, related			
		Allogeneic, unrelated			
	49.		Specify product type (check all that apply)		
		Bone marrow			
		PBSC			
		Single cord blood unit			
		Other product– Go to question 50.			
	50.	Specify other product:			
51.	Is the	product genetically modified?			
		Yes			
		No			
	If aut	ologous, go to question 80			
	If allo	ogeneic related, go to question 52			
	If allo	ogeneic unrelated, go to question 56	.		
	52.	Specify the related donor type			
		☐ Syngeneic (monozygotic twin)	– Go to question 57.		

		HLA-i	dentical sibling (may include non-monozygotic twin) – Go to question 57.	
		HLA-r	matched other relative (does NOT include a haplo-identical donor) - Go to question 53.	
		HLA-r	mismatched relative– Go to question 53 .	
		53.	Specify the biological relationship of the donor to the recipient	
			Mother	
			Father	
			Child	
			Sibling	
			Fraternal twin	
			Maternal aunt	
			Maternal uncle	
			Maternal cousin	
			Paternal aunt	
			Paternal uncle	
			Paternal cousin	
			Grandparent	
			Grandchild	
			Other biological relative – Go to question 54.	
			54. Specify other biological relative:	- Go to
			question 55.	00 10
		55.	Degree of mismatch (related donors only)	
			HLA-mismatched 1 allele– Go to question 57.	
			HLA-mismatched ≥2 alleles (does include haplo-identical donor) – Go to question 57.	
56.	Spec	cify unre	elated donor type	
	_		matched unrelated	
		HLA r	nismatched unrelated	
57.	Did I	NMDP /	Be the Match facilitate the procurement, collection, or transportation of the product?	
		Yes		
		No		
58.	Was	this dor	nor used for any prior HCTs? (for this recipient)	
		Yes) F (F - 9	

ITR C	enter Number:	CIBMTR Research ID:
	□ No	
59.	NMDP cord blood unit ID:	
60.	NMDP donor ID:	— Go to question 63.
61.	Non-NMDP unrelated donor ID	: (not applicable for related donors)
		Go to question 63.
62.	Non-NMDP cord blood unit ID:	(include related and autologous CBUs)
		Go to question 63.
	63.	Global Registration Identifier for Donors (GRID):
	NMDP cord blood unit, go to	
	NMDP donor, go to question	
	Non-NMDP unrelated donor,	go to question 66.
	Non-NMDP cord blood unit,	go to question 64.
6.4	le the ODILID also the IODT DU	Mary had
64.	Is the CBU ID also the ISBT DI ☐ Yes – Go to question 66	
	□ No – Go to question 65	
	☐ Unknown— Go to questi	
	65.	Specify the ISBT DIN number:
66.	Registry or UCB Bank ID:	If 'Other registry' go to 67., otherwise go to question 68.
	67.	Specify other Registry or UCB Bank: Go to question 68
68.	Date of birth (donor / infant)	
	☐ Known – Go to question	n 69.
	☐ Unknown – Go to quest	ion 70.
	69. to question 72.	Date of birth: (donor / infant)

CIBMTR Center Number:				CIBMTR Research ID:					
						YYYY	MM	DD	
			70.			Age (donor)	' infant)		
				Known – C	o to questi	ion 71.			
				Unknown –	Go to ques	stion 72.			
				71. old)		Age: (done	or / infant)	Months (use only if less than 1 year	
						☐ Years			
		72.				Sex (donor /	' infant)		
			Male						
			Fema	le					
73.	Spec	cify blo	od type	(donor) (nor	-NMDP allo	ogeneic donoi	rs only)		
		Α							
		В							
		AB							
		0							
74.	Spec	cifv Rh	factor (ídonor) (non-	NMDP allog	geneic donors	s only)		
		Posi							
		Nega							
75.				dies (IgG or	Total) (Allo	geneic HCTs	only)		
		Read							
	_		reactive						
			termina	te					
		Not o		olo (oord bloo	d unit)				
		NOL 6	аррисац	ole (cord bloo	u uriii)				
76.						ttee (or similar I donors only		consent form to donate research blood	
		Yes	(donor d	consented) –	Go to ques	tion 77.			
		No (donor d	eclined) - Go	to question	า 80.			
		Not a	approac	hed - Go to	question 80).			
		Not applicable (center not participating) - Go to question 80.							

CIBMTR Center Number:			lumber:	CIBMTR Research ID:		
	77.	Date	form was signed:			
			YYYY	MM DD		
	78.	Did th	ne donor submit a research samp	ole to the NMDP/CIBMTR repository? (Related donors only)		
			Yes – Go to question 79.			
			No – Go to question 80.			
			79.	Research sample donor ID:		
same		ction I		single product when they are all from the same donor and use the oblization, if applicable), even if the collections are performed on		
80.	Speci	fy nun	nber of products infused from this	s donor:		
81.	Speci	fy the	number of these products intend	ed to achieve hematopoietic engraftment:		
Ques	stions	8283	are for autologous HCT recip	ients only. If other than autologous skip to question 84		
82.	What	agent	s were used to mobilize the autol	logous recipient for this HCT? (check all that apply)		
		G-CS	SF (filgrastim, Neupogen)			
		Pegy	lated G-CSF (pegfilgrastim, Neul	asta)		
		Plerix	kafor (Mozobil)			
		Coml	bined with chemotherapy			
			CD20 (rituximab, Rituxan)			
			r agent– Go to question 83.			
	83.	Spec	ify other agent:			
				To report more than one donor, copy questions 4883. and complete for each donor.		
				Clinical Status of Recipient Prior to the Preparative Regimen (Conditioning)		
84.	What	scale	was used to determine the recipi	ent's functional status?		
			ofsky (recipient age ≥ 16 years) -			
			ky (recipient age ≥ 1 year and < 1	•		

CIBMTR Center Number:			umb	er: CIBMTR Research ID:
	Perfo	rmano	ce s	core prior to the preparative regimen:
	85.	Karno	ofsky	Scale (recipient age ≥ 16 years)
			100	Normal; no complaints; no evidence of disease - <i>Go to question 87.</i>
			90	Able to carry on normal activity - Go to question 87.
			80	Normal activity with effort - Go to question 87.
			70	Cares for self; unable to carry on normal activity or to do active work - Go to question 87.
			60	Requires occasional assistance but is able to care for most needs - Go to question 87.
			50	Requires considerable assistance and frequent medical care - Go to question 87.
			40	Disabled; requires special care and assistance - Go to question 87.
			30	Severely disabled; hospitalization indicated, although death not imminent - Go to question 87.
			20	Very sick; hospitalization necessary - Go to question 87.
			10	Moribund; fatal process progressing rapidly - Go to question 87.
	86.	Lansk	ky So	cale (recipient age ≥ 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
			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
87.	Recip	ient Cl	MV-a	antibodies (IgG or Total)
		React		
		Non-r	eact	tive
		Indete	ermii	nate
		Not d	one	
				Comorbid Conditions
<mark>88.</mark>				been infected with COVID-19 (SARS-CoV-2) based on a positive test result at any time prior to the start regimen / infusion?
				to question 89.

		No – Go to question 91.	
	89.	Did the patient require hospitalization for management of COVID-19 (SARS-CoV-2) infection?
		□ Yes	
		□ No	
	90.	Was mechanical ventilation given for COVID-19 (SARS-CoV-2) infection?	
		□ Yes	
		□ No	
91.	Is the	ere a history of mechanical ventilation (excluding COVID-19 (SARS-CoV-2))?	
		Yes	
		No	
92.	Is the	ere a history of invasive fungal infection?	
		Yes	
		No	
93.	Glon	erular filtration rate (GFR) before start of preparative regimen (pediatric only)	
		Known- Go to question 94.	
		Unknown- Go to question 95.	
	94.	Glomerular filtration rate (GFR): mL/min/1.73 ²	
95.		the recipient have known complex congenital heart disease? (corrected or uncorrector PDA repair) (pediatric only)	cted) (excluding simple ASD
		Yes	
		No	
96.	(Sou	there any co-existing diseases or organ impairment present according to the HCT orce: Sorror, M. L. (2013). How I assess comorbidities before hematopoietic cell trans-2863.)	
		Yes- Go to question 97.	
		No- Go to question 103.	
		97. Specify co-existing diseases or organ impair	irment (check all that apply)
		☐ Arrhythmia - Any history of atrial fibrillation or flutter, sick sinus syndro arrhythmias requiring treatment	ome, or ventricular

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					
	rovascular disease -Any history of transient ischemic attack, subarachnoid hemorrhage or ral thrombosis, embolism, or hemorrhage				
Diabe	tes -Requiring treatment with insulin or oral hypoglycemic drugs in the last 4 weeks but not lone				
	valve disease -At least a moderate to severe degree of valve stenosis or insufficiency as mined by Echo; prosthetic mitral or aortic valve; or symptomatic mitral valve prolapse				
limit o	ic, mild - Bilirubin > upper limit of normal to $1.5 \times upper$ limit of normal, or AST/ALT > upper of normal to $2.5 \times upper$ limit of normal at the time of transplant OR any history of hepatitis B patitis C infection				
•	ic, moderate/severe -Liver cirrhosis, bilirubin > 1.5 \times upper limit of normal, or AST/ALT > 2.5 \times limit of normal				
suspi Patie	on -Includes a documented infection, fever of unknown origin, or pulmonary nodules cious for fungal pneumonia or a positive PPD test requiring prophylaxis against tuberculosis its must have started antimicrobial treatment before Day 0 with continuation of antimicrobial ment after Day 0				
Inflam	matory bowel disease -Any history of Crohn's disease or ulcerative colitis requiring treatment				
	ty -Patients older than 18 years with a body mass index (BMI) > 35 kg/m2 prior to the start of tioning or a BMI of the 95th percentile of higher for patients aged 18 years or younger				
-	ulcer -Any history of peptic (gastric or duodenal) ulcer confirmed by endoscopy or radiologic osis requiring treatment				
-	iatric disturbance -Presence of any mood (e.g., depression), anxiety, or other psychiatric der (e.g. bipolar disorder or schizophrenia) requiring continuous treatment in the last 4 weeks				
	onary, moderate -Corrected diffusion capacity of carbon monoxide and/or FEV1 of 66-80% or nea on slight activity attributed to pulmonary disease at transplant				
dyspi	nary, severe -Corrected diffusion capacity of carbon monoxide and/or FEV1 of ≤ 65% or nea at rest attributed to pulmonary disease or the need for intermittent or continuous oxygen g the 4 weeks prior to transplant				
	, moderate / severe -Serum creatinine > 2 mg/dL or > 177 μmol/L; on dialysis during the 4 s prior to transplant; OR prior renal transplantation -go to question 98.				
rheur	matologic -Any history of a rheumatologic disease (e.g., systemic lupus erythematosis, natoid arthritis, polymyositis, mixed connective tissue disease, or polymyalgia rheumatica, requiring treatment. (Do NOT include degenerative joint disease, osteoarthritis)				
	malignancy-Treated at any time point in the patient's past history, other than the primary se for which this infusion is being performed <i>-go to question 99.</i>				
98. regir	Was the recipient on dialysis immediately prior to start of preparative nen?				
	Yes				
	No				
П	Unknown				

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			99.		Specify prior malignancy (check all that apply)
				Breast cancer	
				Central nervous syster	m (CNS) malignancy (e.g., glioblastoma, astrocytoma)
				Gastrointestinal malign	nancy (e.g., colon, rectum, stomach, pancreas, intestine, esophageal)
				Genitourinary malignar	ncy (e.g., kidney, bladder, ovary, testicle, genitalia, uterus, cervix, prostate)
				Leukemia (includes ac	ute or chronic leukemia)
				Lung cancer	
				Lymphoma (includes F	Hodgkin & non-Hodgkin lymphoma)
				MDS / MPN	
				Melanoma	
				Multiple myeloma / pla	sma cell disorder (PCD)
				Oropharyngeal cancer	(e.g., tongue, buccal mucosa)
				Sarcoma	
				Thyroid cancer	
				Other skin malignancy	(basal cell, squamous)- go to question 100.
				Other hematologic mal	lignancy -go to question 101.
				Other solid tumor, prior	r -go to question 102.
				100.	Specify other skin malignancy: (prior)
				101.	Specify other hematologic malignancy: (prior)
				102.	Specify other solid tumor: (prior)
					Use results within 4 weeks prior to the start of the preparative regimen, report results from the test performed closest to the start date. Biomarkers according to the augmented HCT comorbidity index. (Source: Biol Blood Marrow Transplant. 2015 Aug; 21(8): 1418–1424)
L03.	Serui	m ferritin	(within	n 4 weeks prior to the s	tart of the preparative regimen, use result closest to the start date)
		Known	– Go i	to question 104.	
		Unknov	vn – G	o to question 107.	
	104.			ng/mL (μg/L)	
	105.	Date sa	ample	collected:	
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		YYYY MM	DD		
	106.	. Upper limit of normal for your institution:			
107.	Serur	um albumin (within 4 weeks prior to the start of the	e preparative regimen, use result closest to the start date)		
		Known – Go to question 108.			
		Unknown – Go to question 110.			
	108.	g • □ g/d			
		☐ g/L			
	400				
	109.	. Date sample collected:			
		TTTT IVIIVI			
110.	Platel	telets (within 4 weeks prior to the start of the prepa	arative regimen, use result closest to the start date)		
		Known – Go to question 111.			
		Unknown – Go to question 113.			
		F 1000 (102/ 20		
	111.		10³/mm³)		
		□ x 10 ⁶ /L			
	112.	 Were platelets transfused ≤ 7 days before date 	of test?		
		□ Yes			
		□ No			
		□ Unknown			
113.	Did th	the recipient have a prior solid organ transplant?	·		
		Yes- Go to question 114.			
		No- Go to question 117.			
	114.	. Specify organ:			
		□ Bowel			
		☐ Heart			
		☐ Kidney(s)			
		□ Liver			
		□ Lung(s)			
		□ Pancreas			

CIBMTR Center Number:					CIBMTR Research ID:	
			Other	organ- Go to question 1	15 .	
			115.		Specify other organ:	
	116.	Year	of prior	solid organ transplant: _		
					Copy and complete questions 114116. for each prior solid organ transplant	
					Pre-HCT Preparative Regimen (Conditioning)	
117.	Heigh	nt at ini	tiation (of pre-HCT preparative re	egimen: inches	
					centimeters	
118.	Actua	Actual weight at initiation of pre-HCT preparative regimen: pounds				
119.	Was	a pre-l	HCT pre	eparative regimen prescri	hed?	
		-	-	question 120.		
				question 141.		
	120.				parative regimen (Allogeneic HCTs only)	
				ablative		
				nyeloablative (NST)		
			Reduc	ed intensity (RIC)		
	121.	Was	irradiati	on planned as part of the	pre-HCT preparative regimen?	
			Yes –	Go to question 122.		
			No – C	Go to question 127.		
			122.		What was the prescribed radiation field?	
				Total body – Go to que		
				Total body by intensity-	modulated radiation therapy (IMRT) – Go to question 123.	
				Total lymphoid or nodal	regions – Go to question 123.	
				Thoracoabdominal region	on – Go to question 123.	
			123.] Gy	Total prescribed dose: (dose per fraction x total number of fractions)	

CIBMTR Ce	enter N	umber:		CIBMTR Research ID:		
					ПоСу	
					□ cGy	
		124.		Date started:		
			YYYY	MM DD		
		125.		Was the radiation fractionated?		
			Yes – Go to question 1			
			No – Go to question 12			
			126.	Total number of fractions:		
Indic	ate the	e total r	prescribed cumulative d	lose for the preparative regimen		
127.	Drug	(drop d	own list)			
		Benda	mustine			
		Busulf	an			
		Carbo	platin			
		Carmu	ustine (BCNU)			
		CCNU	(Lomustine)			
		Clofar	abine (Clolar)			
		Cyclop	phosphamide (Cytoxan)			
		Cytara	abine (Ara-C)			
		Etopo	side (VP-16, VePesid)			
		Fludar	rabine			
		Gemci	itabine			
		Ibritum	nomab tiuxetan (Zevalin)			
		Ifosfar	mide			
		Melph	nalan (L-Pam)			
		Methyl	Iprednisolone (Solu-Medr	ol)		
		Pento	statin			
		Propyl	ene glycol-free melphalaı	n (Evomela)		
		Rituxir	nab (Rituxan)			
		Thiote	ера			
		Tositu	momab (Bexxar)			
		Treosu	ulfan			

Other drug -go to question 128.

CIBMTR Center Number:			CIBMTR Research ID:		
		128.	Specify other drug:		
	129.	Total prescribed dose:	□ mg/m²		
			□ mg/kg		
			☐ AUC (mg x h/L)		
			□ AUC (µmol x min/L)		
			□CSS (ng/mL)		
	120	Data atartada			
	130.	Date started:			
		1111	IVIIVI DD		
	131.	Specify administration (busulfan on	(y)		
		□ Oral			
		□ IV			
		□ Both			
	Con	word complete supption 127, 121	to veneral each dwar given for the preparative regimen		
	Cop	y and complete question 127131	. to report each drug given for the preparative regimen		
			Additional Drugs Given in the Peri-Transplant Period		
132.	ALG,	ALS, ATG, ATS			
		Yes – Go to question 133.			
		No – Go to question 136.			
	122	Total prescribed dose:	mg/kg		
	100.	Total prescribed dose.			
	134.	Specify source			
		☐ ATGAM (horse) – Go to que	estion 136.		
		☐ ATG – Fresenius (rabbit) – G	Go to question 136.		
		☐ Thymoglobulin (rabbit) – <i>Go</i>	to question 136.		
		□ Other – Go to question 135			
		105	Consolit and a second		
		135.	Specify other source:		
136.	Alemi	uzumab (Campath)			
		Yes – Go to question 137.			
		No – Go to question 138.			
	137.		Total prescribed dose: mg/m2		

CIBMTR Center Number:			umber: CIBMTR Research ID:		
□ mg/kg					
			g		
			□mg		
	138.		Defibrotide		
		Yes			
		No			
	139.		KGF		
		Yes	KGI		
		No			
	_	140			
	140.		Ursodiol		
		Yes			
		No			
			GVHD Prophylaxis		
			This section is to be completed for allogeneic HCTs only; autologous HCTs continue with question 144		
			Tiers continue with question 144		
141.	Was	GVHD	prophylaxis planned?		
		Yes -	Go to question 142.		
		No - 0	Go to question 144.		
	142.	Snaci	fy drugs / intervention (check all that apply)		
	142.		Abatacept		
			Anti CD 25 (Zenapax, Daclizumab, AntiTAC)		
			Bortezomib		
			CD34 enriched (CD34+ selection)		
			Corticosteroids (systemic)		
			Cyclophosphamide (Cytoxan)		
			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					
		Blinded randomized trial					
		Other agent-go to question 1	43.				
		143.	Specify other agent:	(do not report ATG, campath)			
			Post-HCT Disease Therapy	Planned as of Day 0			
144. Is	additio	nal post-HCT therapy planned?					
	Yes -	Go to question 145.					
	No - (Go to First Name					
Questio	ns 145.	146. are optional for non-U.S.	centers				
14	45. Sp	ecify 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					
		Gilteritinib					
		Ibrutinib					

CIBMTR Center Number:			umber:	CIBMTR Research ID:	
			Imatinib mesylate (Gleevec, Gli	vec)	
			Intrathecal therapy (chemothera	ару)	
			Ivosidenib		
			Ixazomib		
			Lenalidomide (Revlimid)		
	□ Lestaurtinib□ Local radiotherapy				
			Midostaurin		
			Nilotinib		
			Obinutuzumab		
			Pacritinib		
	☐ Ponatinib				
			Quizartinib		
	☐ Rituximab (Rituxan, MabTher				
			Sorafenib		
			Sunitinib		
			Thalidomide (Thalomid)		
			Other therapy- Go to question	146.	
			Unknown		
			146.	Specify other therapy:	
				Prior Exposure: Potential Study Eligibility	
				Selecting any option(s) below may generate an additional supplemental form.	
147.	Spec	ify if the	e recipient received any of the fo	ollowing (at any time prior to HCT / infusion) (check all that apply)	
		Blinat	umomab (Blincyto)		
		Gemt	uzumab ozogamicin (Mylotarg)		
		Inotuz	zumab ozogamicin (Besponsa)		
		Adien	ne Tepadina®		
		Moga	mulizumab (Poteligeo)		
		None	of the above		

CIBMTR Center Number:			CIBMT	CIBMTR Research ID:			
First Name:							
Last Name:							
E-mail address	5.						
Date:							
Date							
	YYYY	MM	DD				