

Information Collection Domains

Indicates the category of information collection by time period that corresponds to the burden table. For each of the following Domains, there is a corresponding Tab.

1- Pre-Transplant Information Collection

2- Transplant Procedure and Product Information

3- Post-Transplant Periodic Information Collection

Below are the definitions for each column heading.

Column Header Title	Column Header Title Definitions
Information Collection Domain Sub-Type	Identifies a grouping of information collection within an Information Collection Domain. These information collection domain sub types roughly correspond to section/domain headers currently found on CIBMTR data collection instruments.
Information Collection Domain Additional Sub Domain	Additional Sub Domain set recipient, donor, infusion type or product criteria that must be met for an information collection element to be required
Response required if Additional Sub Domain applies	Response options are "yes" or "no". If the criteria noted in Additional sub domain applies, the information collection data element will be applicable and information collection data element responses supplied. Always "yes" when an additional sub domain is present.
Information Collection may be requested at multiple times	Response options are "yes" or "no". Some information may be collected at "multiple" time points or in multiple iterations. A multiple request may occur with a new or duplicate event, new infusion, changes in treatment or outcomes follow up. For example: product analyses at multiple timepoints, chimerism analyses on multiple dates, subsequent neoplasms, co-morbidities, covid infection, Disease Status, Post Transplant Therapy, GVHD, labs and pathology (collected at diagnosis, between diagnosis and infusion, at infusion and during followup)
Current Information Collection Data Element (if applicable)	Depicts the information collection data element currently being requested.
Current Information Collection Data Element Response Option(s)	Depicts the information collection data element response options currently being requested.
Information Collection update:	Notes the type of update. If Blank, there was no change.
	options:
	Addition of Information Requested
	Deletion of Information Requested
	Deletion of Information: Merged to Check all that Apply
	Change/Clarification of Information Requested
	Change/Clarification of Response Options
	Change/Clarification of Information Requested and Response Options
	Data will be captured on Lab Module
Proposed Information Collection Data Element (if applicable)	Depicts the changes to the information collection data element requested in red line format. Rows containing changes are highlighted in Yellow
Proposed Information Collection Data Element Response Option(s)	Depicts the changes to the information collection data element response options in red line format. Rows containing changes are highlighted in yellow.
Rationale for Information Collection Update	The following options identify the change summary:
	options:
	Reduce burden: expanded response options to include responses previously reported manually or created a "check all that apply"
	Be consistent with current clinical landscape, improve transplant outcome data
	Capture data accurately
	Examples added or typographical errors corrected for clarification
	Covid-19 Impact
	Capture additional relevant disease information

[illegible]

Information Collection Domain: Pre-Transplant Information Collection											
Item ID	Time Point	Information Collection Domain Sub-Type	Information Collection Domain Additional Sub-Domain	Response required if Additional Sub Domain applies	Information Collection may be requested multiple times	Current Information Collection Data Element (if applicable)	Current Information Collection Data Element Response Option(s)	Information Collection update:	Proposed Information Collection Data Element (if applicable)	Proposed Information Collection Data Element Response Option(s)	Rationale for Information Collection Update
PRE388	Pre-Transplant	Pre-Transplant Essential Data	Clinical Trial Participants	yes	no	Study Sponsor	BMT CTN, RIC-BMT, PDTC, USDNET, COO, PoSAL, Other sponsor	Change/Clarification of Response Options	Study Sponsor	BMT CTN, NeuBMT CIBMT® CRO Services, PDTC, USDNET, COO, PoSAL, Other sponsor	Capture data accurately
PRE390	Pre-Transplant	Pre-Transplant Essential Data	Clinical Trial Participants	yes	no	Study ID Number	A Representative list of current response options is shown here. This list will change on a frequent basis to accommodate updates - changes in the response options do not affect burden of completing this question. BMT CTN 0301 - Asplastic Anemia.BMT CTN 0601 - Sickle Cell Anemia.BMT CTN 0701 - Follicular Lymphoma.BMT CTN 0702 - Myeloma.BMT CTN 0801 - Chronic GVHD Treatment.BMT CTN 0803 - Auto-HCT in HIV + Patients.RC-BMT 09 - HIV-BMT CTN 0901 - Myeloablative vs. RIC.BMT CTN 0902 - Post-Tx Stress Mgmt.BMT CTN 0903 - Allo-HCT in HIV + Patients.RC-BMT 10 - CBARCBT 10-CHS4MDS-1.RC-BMT 11 - Treos.BMT CTN 1101 - Haplo vs. Double UCB with RIC.BMT CTN 1102 - MDS in older patients.RC-BMT 12 - Mosa.BMT CTN 1202 - Biomelev.BMT CTN 1203 - GVHD Prophylaxis.BMT CTN 1204 - HLA-BMT CTN 1205 - Easy-to-read Consent from EBTC.RC-BMT 13 - TREC.BMT CTN 1301 - CN-Free.BMT CTN 1302 - Allo-BM.BMT CTN 1401 - Myeloma Vaccine.RC-BMT 140-AD0-202.RC-BMT 15 - MMAD.BMT CTN 1501 - Standard Risk GVHD.BMT CTN 1502 - CHAMP Asplastic Anemia.BMT CTN 1503 - STRIDE2.BMT CTN 1504 - AML Maintenance Therapy.BMT CTN 1507 - Haplo-Sickle Cell.RC-BMT 16-CHS4MDS-RC-BMT 16 - HTCC.RC-BMT 17 - CD34.RC-BMT 17-CAS-NA.RC-BMT 17-CAS-SC2.RC-BMT 17 - CSDE.BMT CTN 1703 - PROGRESS II.BMT CTN 1704 - CHARM.BMT CTN 1803 - Haplo-NR Cell.BMT CTN 1903 - HIV T Cell.BMT CTN 1904 - Treos BM Failure Syndrome.BMT CTN 1905 - BEAT-MS (TNO7A).PDTC 4901 - Disorders of the Immune system (SCID).PDTC 4903 - Disorders of the Immune system (SCID).PDTC 4904 - Disorders of the Immune system (WAS).RC-BMT ACCESS.RC-BMT 68 - DS.RC-BMT 900.	Change/Clarification of Response Options	Study ID Number	A Representative list of current response options is shown here. This list will change on a frequent basis to accommodate updates - changes in the response options do not affect burden of completing this question. BMT CTN 0301 - Asplastic Anemia.BMT CTN 0601 - Sickle Cell Anemia.BMT CTN 0701 - Follicular Lymphoma.BMT CTN 0702 - Myeloma.BMT CTN 0801 - Chronic GVHD Treatment.BMT CTN 0803 - Auto-HCT in HIV + Patients.RC-BMT 09 - HIV-BMT CTN 0901 - Myeloablative vs. RIC.BMT CTN 0902 - Post-Tx Stress Mgmt.BMT CTN 0903 - Allo-HCT in HIV + Patients.RC-BMT 10 - CBARCBT 10-CHS4MDS-1.RC-BMT 11 - Treos.BMT CTN 1101 - Haplo vs. Double UCB with RIC.BMT CTN 1102 - MDS in older patients.RC-BMT 12 - Mosa.BMT CTN 1202 - Biomelev.BMT CTN 1203 - GVHD Prophylaxis.BMT CTN 1204 - HLA-BMT CTN 1205 - Easy-to-read Consent from EBTC.RC-BMT 13 - TREC.BMT CTN 1301 - CN-Free.BMT CTN 1302 - Allo-BM.BMT CTN 1401 - Myeloma Vaccine.RC-BMT 140-AD0-202.RC-BMT 15 - MMAD.BMT CTN 1501 - Standard Risk GVHD.BMT CTN 1502 - CHAMP Asplastic Anemia.BMT CTN 1503 - STRIDE2.BMT CTN 1504 - AML Maintenance Therapy.BMT CTN 1507 - Haplo-Sickle Cell.RC-BMT 16-CHS4MDS-RC-BMT 16 - HTCC.RC-BMT 17 - CD34.RC-BMT 17-CAS-NA.RC-BMT 17-CAS-SC2.RC-BMT 17 - CSDE.BMT CTN 1703 - PROGRESS II.BMT CTN 1704 - CHARM.BMT CTN 1803 - Haplo-NR Cell.BMT CTN 1903 - HIV T Cell.BMT CTN 1904 - Treos BM Failure Syndrome.BMT CTN 1905 - BEAT-MS (TNO7A).PDTC 4901 - Disorders of the Immune system (SCID).PDTC 4903 - Disorders of the Immune system (SCID).PDTC 4904 - Disorders of the Immune system (WAS).RC-BMT ACCESS.RC-BMT 68 - DS.RC-BMT 900.	Capture data accurately
PRE437	Pre-Transplant	Pre-Transplant Essential Data	Autologous Transplant	yes	yes	Name of product (gene therapy recipient)	Other name	Change/Clarification of Response Options	Name of product (gene therapy recipient)	Exagamgine autotemex (Zyntigro®), Ectagamgine autotemex (Ectagovon®), Exagamgine autotemex, Other name	Capture data accurately
PRE494	Pre-Transplant	Pre-Transplant Essential Data		no	no	Stomachal filtration rate (SFR)	----- mL/min/1.73-2	Change/Clarification of Response Options	Stomachal filtration rate (SFR)	----- mL/min/1.73-2	Capture data accurately
PRE001	Pre-Transplant	Additional Drug Given in the Post-transplant Period		no	no	ALG, ALS, ATG, ATS, Alemtuzumab, Deffirobide, KGF, Ursodiol, None	(check all that apply)		ALG, ALS, ATG, ATS, Alemtuzumab, Deffirobide, KGF, Ursodiol, None	(check all that apply)	
PRE002	Pre-Transplant	Additional Drug Given in the Post-transplant Period		no	no	Total prescribed dose:	----- mg/kg		Total prescribed dose:	----- mg/kg	
PRE003	Pre-Transplant	Additional Drug Given in the Post-transplant Period		no	no	Specify source	ATGAM (horse),ATG - Fresenius (rabbit),Other,Thymoglobulin (rabbit)		Specify source	ATGAM (horse),ATG - Fresenius (rabbit),Other,Thymoglobulin (rabbit)	
PRE004	Pre-Transplant	Additional Drug Given in the Post-transplant Period		no	no	Specify other source:	open text		Specify other source:	open text	
PRE005	Pre-Transplant	Additional Drug Given in the Post-transplant Period		no	no	Total prescribed dose:	----- mg/m2 ----- mg/kg ----- mg/kg		Total prescribed dose:	----- mg/m2 ----- mg/kg ----- mg/kg	
PRE006	Pre-Transplant	Covid-19 Impact		no	no			Was the HCT impacted for a reason related to the COVID-19 (SARS Cov-2) pandemic?	no,yes		
PRE007	Pre-Transplant	Covid-19 Impact		no	no			Is the HCT date different than the originally intended HCT date?	no,yes		
PRE008	Pre-Transplant	Covid-19 Impact		no	no			Original date of HCT	YYYY/MM/DD		
PRE009	Pre-Transplant	Covid-19 Impact		no	no			Date estimated	checked		
PRE010	Pre-Transplant	Covid-19 Impact		no	no			Is the donor different than the originally intended donor?	no,yes		
PRE011	Pre-Transplant	Covid-19 Impact		no	no			Specify the originally intended donor	unrelated donor, syngeneic (monozygotic twin) , HLA-identical sibling (may include non-monozygotic twin) , HLA-matched other relative (does NOT include a haplo-identical donor), HLA-mismatched relative		
PRE012	Pre-Transplant	Covid-19 Impact		no	no			Is the product type (bone marrow, PBSC, cord blood unit) different than the originally intended product type?	no,yes		
PRE013	Pre-Transplant	Covid-19 Impact		no	no			Specify the originally intended product type	bone marrow,Other product,PBSC, cord blood unit		
PRE014	Pre-Transplant	Covid-19 Impact		no	no			Specify other product type	open text		
PRE015	Pre-Transplant	Covid-19 Impact		no	no			Was the current product thawed from a previously stored state prior to infusion?	no,yes		
PRE016	Pre-Transplant	Covid-19 Impact		no	no			Did the preparative regimen change from the original plan?	no,yes		
PRE017	Pre-Transplant	Covid-19 Impact		no	no			Did the Graft prophylaxis change from the original plan?	no,yes		
PRE018	Pre-Transplant	Disease Classification	Acute Myelogenous Leukemia (AML)	yes	no	Specify method(s) that was used to assess measurable residual disease status (check all that apply)	FISH, Karyotyping, Flow Cytometry, PCR, NGS, Not assessed	Specify method(s) that was used to assess measurable residual disease status (check all that apply)	FISH, Karyotyping, Flow Cytometry, PCR, NGS, Not assessed		
PRE019	Pre-Transplant	Disease Classification	Acute Myelogenous Leukemia (AML)	yes	no	Was measurable residual disease detected by FISH?	no,yes	Was measurable residual disease detected by FISH?	no,yes		
PRE020	Pre-Transplant	Disease Classification	Acute Myelogenous Leukemia (AML)	yes	no	Was measurable residual disease detected by karyotyping assay?	no,yes	Was measurable residual disease detected by karyotyping assay?	no,yes		
PRE021	Pre-Transplant	Disease Classification	Acute Myelogenous Leukemia (AML)	yes	no	Which leukemia phenotype was used for detection (check all the apply)	original leukemia immunophenotype, aberrant phenotype	Which leukemia phenotype was used for detection (check all the apply)	original leukemia immunophenotype, aberrant phenotype		
PRE022	Pre-Transplant	Disease Classification	Acute Myelogenous Leukemia (AML)	yes	no	What is the lower limit of detection (for the original leukemia immunophenotype)	open text	What is the lower limit of detection (for the original leukemia immunophenotype)	open text		
PRE023	Pre-Transplant	Disease Classification	Acute Myelogenous Leukemia (AML)	yes	no	What is the lower limit of detection (for the aberrant phenotype)	open text	What is the lower limit of detection (for the aberrant phenotype)	open text		
PRE024	Pre-Transplant	Disease Classification	Acute Myelogenous Leukemia (AML)	yes	no	Was measurable residual disease detected by flow cytometry?	no,yes	Was measurable residual disease detected by flow cytometry?	no,yes		
PRE025	Pre-Transplant	Disease Classification	Acute Myelogenous Leukemia (AML)	yes	no	Was measurable residual disease detected by PCR?	no,yes	Was measurable residual disease detected by PCR?	no,yes		
PRE026	Pre-Transplant	Disease Classification	Acute Myelogenous Leukemia (AML)	yes	no	Was measurable residual disease detected by NGS?	no,yes	Was measurable residual disease detected by NGS?	no,yes		
PRE027	Pre-Transplant	Disease Classification	Acute Lymphoblastic Leukemia (ALL)	yes	no	Specify method(s) that was used to assess measurable residual disease status (check all that apply)	FISH, Karyotyping, Flow Cytometry, PCR, NGS, Not assessed	Specify method(s) that was used to assess measurable residual disease status (check all that apply)	FISH, Karyotyping, Flow Cytometry, PCR, NGS, Not assessed		
PRE028	Pre-Transplant	Disease Classification	Acute Lymphoblastic Leukemia (ALL)	yes	no	Was measurable residual disease detected by FISH?	no,yes	Was measurable residual disease detected by FISH?	no,yes		
PRE029	Pre-Transplant	Disease Classification	Acute Lymphoblastic Leukemia (ALL)	yes	no	Was measurable residual disease detected by karyotyping assay?	no,yes	Was measurable residual disease detected by karyotyping assay?	no,yes		
PRE030	Pre-Transplant	Disease Classification	Acute Lymphoblastic Leukemia (ALL)	yes	no	Which leukemia phenotype was used for detection (check all the apply)	original leukemia immunophenotype, aberrant phenotype	Which leukemia phenotype was used for detection (check all the apply)	original leukemia immunophenotype, aberrant phenotype		
PRE031	Pre-Transplant	Disease Classification	Acute Lymphoblastic Leukemia (ALL)	yes	no	What is the lower limit of detection (for the original leukemia immunophenotype)	open text	What is the lower limit of detection (for the original leukemia immunophenotype)	open text		
PRE032	Pre-Transplant	Disease Classification	Acute Lymphoblastic Leukemia (ALL)	yes	no	What is the lower limit of detection (for the aberrant phenotype)	open text	What is the lower limit of detection (for the aberrant phenotype)	open text		
PRE033	Pre-Transplant	Disease Classification	Acute Lymphoblastic Leukemia (ALL)	yes	no	Was measurable residual disease detected by flow cytometry?	no,yes	Was measurable residual disease detected by flow cytometry?	no,yes		
PRE034	Pre-Transplant	Disease Classification	Acute Lymphoblastic Leukemia (ALL)	yes	no	Was measurable residual disease detected by PCR?	no,yes	Was measurable residual disease detected by PCR?	no,yes		
PRE035	Pre-Transplant	Disease Classification	Acute Lymphoblastic Leukemia (ALL)	yes	no	Was measurable residual disease detected by NGS?	no,yes	Was measurable residual disease detected by NGS?	no,yes		
PRE036	Pre-Transplant	Disease Classification	Myeloproliferative Neoplasms (MPN)	yes	no	Specify the liver size	----- centimeters	Specify the liver size	----- centimeters		
PRE037	Pre-Transplant	Disease Classification	Myeloproliferative Neoplasms (MPN)	yes	yes	JAK2 Exon 12	Negative,Not done,Positive	JAK2 Exon 12	Negative,Not done,Positive		
PRE038	Pre-Transplant	Disease Classification	Myeloproliferative Neoplasms (MPN)	yes	yes	Specify abnormalities (check all that apply)	del(11q) / 11q,del(12p) / 12p,del(20q) / 20q,del(5q) / 5q,del(7q) / 7q,del(13q) / 13q,dup(11)(p11.3),5-7,Y,Other abnormality,(11,any),(11q23,any),(11p11.2,any),(12q21,any),(16,9),-8,-9	Specify abnormalities (check all that apply)	del(11q) / 11q,del(12p) / 12p,del(20q) / 20q,del(5q) / 5q,del(7q) / 7q,del(13q) / 13q,dup(11)(p11.3),5-7,Y,Other abnormality,(11,any),(11q23,any),(11p11.2,any),(12q21,any),(16,9),-8,-9		
PRE039	Pre-Transplant	Disease Classification	Myeloproliferative Neoplasms (MPN)	yes	yes	Was documentation submitted to the CIBMT® (e.g. FISH report)	No,Yes	Was documentation submitted to the CIBMT® (e.g. FISH report)	No,Yes		
PRE040	Pre-Transplant	Disease Classification	Hodgkin and Non-Hodgkin Lymphoma	yes	no	Assignment of DLBCL (germinal center B-cell type vs. activated B-cell type) subtype was based on	Gene expression profile,immunohistochemistry (e.g. Han's algorithm),Unknown	Assignment of DLBCL (germinal center B-cell type vs. activated B-cell type) subtype was based on	Gene expression profile,immunohistochemistry (e.g. Han's algorithm),Unknown		
PRE041	Pre-Transplant	Disease Classification		no	yes	Date of diagnosis of primary disease for HCT / cellular therapy	YYYY/MM/DD	Date of diagnosis of primary disease for HCT / cellular therapy	YYYY/MM/DD		

Item ID	Time Point	Information Collection Domain Sub-Type	Information Collection Domain Additional Sub-Domain	Response required if additional sub-Domain applies	Information Collection may be requested multiple times	Current Information Collection Data Element (if applicable)	Current Information Collection Data Element Response Option(s)	Information Collection update:	Proposed Information Collection Data Element (if applicable)	Proposed Information Collection Data Element Response Option(s)	Rationale for information Collection Update
PRE077	Pre-Transplant	Disease Classification	Acute Myelogenous Leukemia (AML)	yes	yes	Were cytogenetics tested via FISH?	No,Yes		Were cytogenetics tested via FISH?	No,Yes	
PRE078	Pre-Transplant	Disease Classification	Acute Myelogenous Leukemia (AML)	yes	yes	Results of tests	Abnormalities identified.No abnormalities		Results of tests	Abnormalities identified.No abnormalities	
PRE079	Pre-Transplant	Disease Classification	Acute Myelogenous Leukemia (AML)	yes	yes	International System for Human Cytogenetic Nomenclature (ISCN) compatible string:	open text		International System for Human Cytogenetic Nomenclature (ISCN) compatible string:	open text	
PRE080	Pre-Transplant	Disease Classification	Acute Myelogenous Leukemia (AML)	yes	yes	Specify number of distinct cytogenetic abnormalities	Four or more (4 or more),One (1),Three (3),Two (2)		Specify number of distinct cytogenetic abnormalities	Four or more (4 or more),One (1),Three (3),Two (2)	
PRE081	Pre-Transplant	Disease Classification	Acute Myelogenous Leukemia (AML)	yes	yes	Specify abnormalities (check all that apply)	1,t(2;3) any abnormality,1;2p any abnormality,del(11q) / 11q,del(14q) / 14q,del(17q) / 17q,del(20q) / 20q,del(21q) / 21q,del(3q) / 3q,del(5q) / 5q,del(7q) / 7q,del(9q) / 9q,inv(16),inv(3), 17-18, 5-7,X,-Y,Other abnormality,(t(15;17) and variants,(t(16;16),(3;3),(6;9),(8;21),(9;11),(9;22),+1,+13,+14,+21,+22,+4,+8		Specify abnormalities (check all that apply)	1,t(2;3) any abnormality,1;2p any abnormality,del(11q) / 11q,del(14q) / 14q,del(17q) / 17q,del(20q) / 20q,del(21q) / 21q,del(3q) / 3q,del(5q) / 5q,del(7q) / 7q,del(9q) / 9q,inv(16),inv(3), 17-18, 5-7,X,-Y,Other abnormality,(t(15;17) and variants,(t(16;16),(3;3),(6;9),(8;21),(9;11),(9;22),+1,+13,+14,+21,+22,+4,+8	
PRE082	Pre-Transplant	Disease Classification	Acute Myelogenous Leukemia (AML)	yes	yes	Specify other abnormality:	open text		Specify other abnormality:	open text	
PRE083	Pre-Transplant	Disease Classification	Acute Myelogenous Leukemia (AML)	yes	yes	Were cytogenetics tested via karyotyping?	No,Yes		Were cytogenetics tested via karyotyping?	No,Yes	
PRE084	Pre-Transplant	Disease Classification	Acute Myelogenous Leukemia (AML)	yes	yes	Results of tests	Abnormalities identified.No abnormalities.No evaluable metaphases		Results of tests	Abnormalities identified.No abnormalities.No evaluable metaphases	
PRE085	Pre-Transplant	Disease Classification	Acute Myelogenous Leukemia (AML)	yes	yes	International System for Human Cytogenetic Nomenclature (ISCN) compatible string:	open text		International System for Human Cytogenetic Nomenclature (ISCN) compatible string:	open text	
PRE086	Pre-Transplant	Disease Classification	Acute Myelogenous Leukemia (AML)	yes	yes	Specify number of distinct cytogenetic abnormalities	Four or more (4 or more),One (1),Three (3),Two (2)		Specify number of distinct cytogenetic abnormalities	Four or more (4 or more),One (1),Three (3),Two (2)	
PRE087	Pre-Transplant	Disease Classification	Acute Myelogenous Leukemia (AML)	yes	yes	Specify abnormalities (check all that apply)	1,t(2;3) any abnormality,1;2p any abnormality,del(11q) / 11q,del(14q) / 14q,del(17q) / 17q,del(20q) / 20q,del(21q) / 21q,del(3q) / 3q,del(5q) / 5q,del(7q) / 7q,del(9q) / 9q,inv(16),inv(3), 17-18, 5-7,X,-Y,Other abnormality,(t(15;17) and variants,(t(16;16),(3;3),(6;9),(8;21),(9;11),(9;22),+1,+13,+14,+21,+22,+4,+8		Specify abnormalities (check all that apply)	1,t(2;3) any abnormality,1;2p any abnormality,del(11q) / 11q,del(14q) / 14q,del(17q) / 17q,del(20q) / 20q,del(21q) / 21q,del(3q) / 3q,del(5q) / 5q,del(7q) / 7q,del(9q) / 9q,inv(16),inv(3), 17-18, 5-7,X,-Y,Other abnormality,(t(15;17) and variants,(t(16;16),(3;3),(6;9),(8;21),(9;11),(9;22),+1,+13,+14,+21,+22,+4,+8	
PRE088	Pre-Transplant	Disease Classification	Acute Myelogenous Leukemia (AML)	yes	yes	Specify other abnormality:	open text		Specify other abnormality:	open text	
PRE089	Pre-Transplant	Disease Classification	Acute Myelogenous Leukemia (AML)	yes	yes	Was documentation submitted to the CIBMTR? (e.g. cytogenetic or FISH report)	No,Yes		Was documentation submitted to the CIBMTR? (e.g. cytogenetic or FISH report)	No,Yes	
PRE090	Pre-Transplant	Disease Classification	Acute Myelogenous Leukemia (AML)	yes	yes	Were tests for molecular markers performed? (e.g. PCR, NGS) (between diagnosis and last evaluation)	No,Unknown,yes		Were tests for molecular markers performed? (e.g. PCR, NGS) (between diagnosis or relapse and last evaluation)	No,Unknown,yes	
PRE091	Pre-Transplant	Disease Classification	Acute Myelogenous Leukemia (AML)	yes	yes	CEBPA	Negative,Not Done,Positive		CEBPA	Negative,Not Done,Positive	
PRE092	Pre-Transplant	Disease Classification	Acute Myelogenous Leukemia (AML)	yes	yes	Specify CEBPA mutation	Biallelic (homozygous),Monoallelic (heterozygous),Unknown		Specify CEBPA mutation	Biallelic (double mutant),Monoallelic (single mutant),Unknown	
PRE093	Pre-Transplant	Disease Classification	Acute Myelogenous Leukemia (AML)	yes	yes	FLT3 - TRD (point mutations in D835 or deletions of codon B356)	Negative,Not done,Positive		FLT3 - TRD (point mutations in D835 or deletions of codon B36)	Negative,Not done,Positive	
PRE094	Pre-Transplant	Disease Classification	Acute Myelogenous Leukemia (AML)	yes	yes	FLT3 - ITD mutation	Negative,Not Done,Positive		FLT3 - ITD mutation	Negative,Not Done,Positive	
PRE095	Pre-Transplant	Disease Classification	Acute Myelogenous Leukemia (AML)	yes	yes	FLT3 - ITD allelic ratio	Known,Unknown		FLT3 - ITD allelic ratio	Known,Unknown	
PRE096	Pre-Transplant	Disease Classification	Acute Myelogenous Leukemia (AML)	yes	yes	Specify FLT3 - ITD allelic ratio:	--- / --- = --- / ---		Specify FLT3 - ITD allelic ratio:	--- / --- = --- / ---	
PRE097	Pre-Transplant	Disease Classification	Acute Myelogenous Leukemia (AML)	yes	yes	IDH1	Negative,Not Done,Positive		IDH1	Negative,Not Done,Positive	
PRE098	Pre-Transplant	Disease Classification	Acute Myelogenous Leukemia (AML)	yes	yes	IDH2	Negative,Not Done,Positive		IDH2	Negative,Not Done,Positive	
PRE099	Pre-Transplant	Disease Classification	Acute Myelogenous Leukemia (AML)	yes	yes	KIT	Negative,Not Done,Positive		KIT	Negative,Not Done,Positive	
PRE100	Pre-Transplant	Disease Classification	Acute Myelogenous Leukemia (AML)	yes	yes	NPM1	Negative,Not Done,Positive		NPM1	Negative,Not Done,Positive	
PRE101	Pre-Transplant	Disease Classification	Acute Myelogenous Leukemia (AML)	yes	yes	Other molecular marker	Negative,Not Done,Positive		Other molecular marker	Negative,Not Done,Positive	
PRE102	Pre-Transplant	Disease Classification	Acute Myelogenous Leukemia (AML)	yes	yes	Specify other molecular marker:	open text		Specify other molecular marker:	open text	
PRE103	Pre-Transplant	Disease Classification	Acute Myelogenous Leukemia (AML)	yes	yes	Were cytogenetics tested (karyotyping or FISH)? (at last evaluation)	No,Unknown,yes		Were cytogenetics tested (karyotyping or FISH)? (at last evaluation)	No,Unknown,yes	
PRE104	Pre-Transplant	Disease Classification	Acute Myelogenous Leukemia (AML)	yes	yes	Were cytogenetics tested via FISH?	No,Yes		Were cytogenetics tested via FISH?	No,Yes	
PRE105	Pre-Transplant	Disease Classification	Acute Myelogenous Leukemia (AML)	yes	yes	Results of tests	Abnormalities identified.No abnormalities		Results of tests	Abnormalities identified.No abnormalities	
PRE106	Pre-Transplant	Disease Classification	Acute Myelogenous Leukemia (AML)	yes	yes	International System for Human Cytogenetic Nomenclature (ISCN) compatible string:	open text		International System for Human Cytogenetic Nomenclature (ISCN) compatible string:	open text	
PRE107	Pre-Transplant	Disease Classification	Acute Myelogenous Leukemia (AML)	yes	yes	Specify number of distinct cytogenetic abnormalities	Four or more (4 or more),One (1),Three (3),Two (2)		Specify number of distinct cytogenetic abnormalities	Four or more (4 or more),One (1),Three (3),Two (2)	
PRE108	Pre-Transplant	Disease Classification	Acute Myelogenous Leukemia (AML)	yes	yes	Specify abnormalities (check all that apply)	1,t(2;3) any abnormality,1;2p any abnormality,del(11q) / 11q,del(14q) / 14q,del(17q) / 17q,del(20q) / 20q,del(21q) / 21q,del(3q) / 3q,del(5q) / 5q,del(7q) / 7q,del(9q) / 9q,inv(16),inv(3), 17-18, 5-7,X,-Y,Other abnormality,(t(15;17) and variants,(t(16;16),(3;3),(6;9),(8;21),(9;11),(9;22),+1,+13,+14,+21,+22,+4,+8		Specify abnormalities (check all that apply)	1,t(2;3) any abnormality,1;2p any abnormality,del(11q) / 11q,del(14q) / 14q,del(17q) / 17q,del(20q) / 20q,del(21q) / 21q,del(3q) / 3q,del(5q) / 5q,del(7q) / 7q,del(9q) / 9q,inv(16),inv(3), 17-18, 5-7,X,-Y,Other abnormality,(t(15;17) and variants,(t(16;16),(3;3),(6;9),(8;21),(9;11),(9;22),+1,+13,+14,+21,+22,+4,+8	
PRE109	Pre-Transplant	Disease Classification	Acute Myelogenous Leukemia (AML)	yes	yes	Specify other abnormality:	open text		Specify other abnormality:	open text	
PRE110	Pre-Transplant	Disease Classification	Acute Myelogenous Leukemia (AML)	yes	yes	Were cytogenetics tested via karyotyping?	No,Yes		Were cytogenetics tested via karyotyping?	No,Yes	
PRE111	Pre-Transplant	Disease Classification	Acute Myelogenous Leukemia (AML)	yes	yes	Results of tests	Abnormalities identified.No abnormalities.No evaluable metaphases		Results of tests	Abnormalities identified.No abnormalities.No evaluable metaphases	
PRE112	Pre-Transplant	Disease Classification	Acute Myelogenous Leukemia (AML)	yes	yes	International System for Human Cytogenetic Nomenclature (ISCN) compatible string:	open text		International System for Human Cytogenetic Nomenclature (ISCN) compatible string:	open text	
PRE113	Pre-Transplant	Disease Classification	Acute Myelogenous Leukemia (AML)	yes	yes	Specify number of distinct cytogenetic abnormalities	Four or more (4 or more),One (1),Three (3),Two (2)		Specify number of distinct cytogenetic abnormalities	Four or more (4 or more),One (1),Three (3),Two (2)	
PRE114	Pre-Transplant	Disease Classification	Acute Myelogenous Leukemia (AML)	yes	yes	Specify abnormalities (check all that apply)	1,t(2;3) any abnormality,1;2p any abnormality,del(11q) / 11q,del(14q) / 14q,del(17q) / 17q,del(20q) / 20q,del(21q) / 21q,del(3q) / 3q,del(5q) / 5q,del(7q) / 7q,del(9q) / 9q,inv(16),inv(3), 17-18, 5-7,X,-Y,Other abnormality,(t(15;17) and variants,(t(16;16),(3;3),(6;9),(8;21),(9;11),(9;22),+1,+13,+14,+21,+22,+4,+8		Specify abnormalities (check all that apply)	1,t(2;3) any abnormality,1;2p any abnormality,del(11q) / 11q,del(14q) / 14q,del(17q) / 17q,del(20q) / 20q,del(21q) / 21q,del(3q) / 3q,del(5q) / 5q,del(7q) / 7q,del(9q) / 9q,inv(16),inv(3), 17-18, 5-7,X,-Y,Other abnormality,(t(15;17) and variants,(t(16;16),(3;3),(6;9),(8;21),(9;11),(9;22),+1,+13,+14,+21,+22,+4,+8	
PRE115	Pre-Transplant	Disease Classification	Acute Myelogenous Leukemia (AML)	yes	yes	Specify other abnormality:	open text		Specify other abnormality:	open text	
PRE116	Pre-Transplant	Disease Classification	Acute Myelogenous Leukemia (AML)	yes	yes	Was documentation submitted to the CIBMTR? (e.g. cytogenetic or FISH report)	No,Yes		Was documentation submitted to the CIBMTR? (e.g. cytogenetic or FISH report)	No,Yes	
PRE117	Pre-Transplant	Disease Classification	Acute Myelogenous Leukemia (AML)	yes	yes	Were tests for molecular markers performed?(e.g. PCR, NGS) (at last evaluation)	No,Unknown,yes		Were tests for molecular markers performed?(e.g. PCR, NGS) (at last evaluation)	No,Unknown,yes	
PRE118	Pre-Transplant	Disease Classification	Acute Myelogenous Leukemia (AML)	yes	yes	CEBPA	Negative,Not Done,Positive		CEBPA	Negative,Not Done,Positive	
PRE119	Pre-Transplant	Disease Classification	Acute Myelogenous Leukemia (AML)	yes	yes	Specify CEBPA mutation	Biallelic (homozygous),Monoallelic (heterozygous),Unknown		Specify CEBPA mutation	Biallelic (double mutant),Monoallelic (single mutant),Unknown	
PRE120	Pre-Transplant	Disease Classification	Acute Myelogenous Leukemia (AML)	yes	yes	FLT3 - TRD (point mutations in D835 or deletions of codon B36)	Negative,Not done,Positive		FLT3 - TRD (point mutations in D835 or deletions of codon B36)	Negative,Not done,Positive	

Item ID	Time Point	Information Collection Domain Sub-Type	Information Collection Domain Additional Sub-Domain	Response required if additional sub-Domain applies	Information Collection may be requested multiple times	Current Information Collection Data Element (if applicable)	Current Information Collection Data Element Response Option(s)	Information Collection update:	Proposed Information Collection Data Element (if applicable)	Proposed Information Collection Data Element Response Option(s)	Rationale for information Collection Update
PRE151	Pre-Transplant	Disease Classification	Acute Lymphoblastic Leukemia (ALL)	yes	yes	Results of tests	Abnormalities identified.No abnormalities		Results of tests	Abnormalities identified.No abnormalities	
PRE152	Pre-Transplant	Disease Classification	Acute Lymphoblastic Leukemia (ALL)	yes	yes	International System for Human Cytogenetic Nomenclature (ISCN) compatible string:	open text		International System for Human Cytogenetic Nomenclature (ISCN) compatible string:	open text	
PRE153	Pre-Transplant	Disease Classification	Acute Lymphoblastic Leukemia (ALL)	yes	yes	Specify number of distinct cytogenetic abnormalities	Four or more (4 or more),One (1),Three (3),Two (2)		Specify number of distinct cytogenetic abnormalities	Four or more (4 or more),One (1),Three (3),Two (2)	
PRE154	Pre-Transplant	Disease Classification	Acute Lymphoblastic Leukemia (ALL)	yes	yes	Specify abnormalities (check all that apply)	11q23) any abnormality,12p any abnormality,9p any abnormality,add(14q),del(12p) / 12p-del(6q) / 6q-del(9p) / 9p-Hyperdiploid (< 50),Hypodiploid (< 46),JAMP21-7,Other abnormality (1;1-19,t(10;14),t(11;14),t(12;21),t(2;8),t(4;11),t(5;14),t(8;14),t(8;22),t(9;22),+17,+21,+4,-8		Specify abnormalities (check all that apply)	11q23) any abnormality,12p any abnormality,9p any abnormality,add(14q),del(12p) / 12p-del(6q) / 6q-del(9p) / 9p-Hyperdiploid (< 50),Hypodiploid (< 46),JAMP21-7,Other abnormality (1;1-19,t(10;14),t(11;14),t(12;21),t(2;8),t(4;11),t(5;14),t(8;14),t(8;22),t(9;22),+17,+21,+4,-8	
PRE155	Pre-Transplant	Disease Classification	Acute Lymphoblastic leukemia (ALL)	yes	yes	Specify other abnormality:	open text		Specify other abnormality:	open text	
PRE156	Pre-Transplant	Disease Classification	Acute Lymphoblastic leukemia (ALL)	yes	yes	Were cytogenetics tested via karyotyping?	No,Yes		Were cytogenetics tested via karyotyping?	No,Yes	
PRE157	Pre-Transplant	Disease Classification	Acute Lymphoblastic leukemia (ALL)	yes	yes	Results of tests	Abnormalities identified.No abnormalities.No evaluable metaphases		Results of tests	Abnormalities identified.No abnormalities.No evaluable metaphases	
PRE158	Pre-Transplant	Disease Classification	Acute Lymphoblastic leukemia (ALL)	yes	yes	International System for Human Cytogenetic Nomenclature (ISCN) compatible string:	open text		International System for Human Cytogenetic Nomenclature (ISCN) compatible string:	open text	
PRE159	Pre-Transplant	Disease Classification	Acute Lymphoblastic leukemia (ALL)	yes	yes	Specify number of distinct cytogenetic abnormalities	Four or more (4 or more),One (1),Three (3),Two (2)		Specify number of distinct cytogenetic abnormalities	Four or more (4 or more),One (1),Three (3),Two (2)	
PRE170	Pre-Transplant	Disease Classification	Acute Lymphoblastic leukemia (ALL)	yes	yes	Specify abnormalities (check all that apply)	11q23) any abnormality,12p any abnormality,9p any abnormality,add(14q),del(12p) / 12p-del(6q) / 6q-del(9p) / 9p-Hyperdiploid (< 50),Hypodiploid (< 46),JAMP21-7,Other abnormality (1;1-19,t(10;14),t(11;14),t(12;21),t(2;8),t(4;11),t(5;14),t(8;14),t(8;22),t(9;22),+17,+21,+4,-8		Specify abnormalities (check all that apply)	11q23) any abnormality,12p any abnormality,9p any abnormality,add(14q),del(12p) / 12p-del(6q) / 6q-del(9p) / 9p-Hyperdiploid (< 50),Hypodiploid (< 46),JAMP21-7,Other abnormality (1;1-19,t(10;14),t(11;14),t(12;21),t(2;8),t(4;11),t(5;14),t(8;14),t(8;22),t(9;22),+17,+21,+4,-8	
PRE171	Pre-Transplant	Disease Classification	Acute Lymphoblastic leukemia (ALL)	yes	yes	Specify other abnormality:	open text		Specify other abnormality:	open text	
PRE172	Pre-Transplant	Disease Classification	Acute Lymphoblastic leukemia (ALL)	yes	yes	Was documentation submitted to the CIBMTR? (e.g. cytogenetic or FISH report)	No,Yes		Was documentation submitted to the CIBMTR? (e.g. cytogenetic or FISH report)	No,Yes	
PRE173	Pre-Transplant	Disease Classification	Acute Lymphoblastic leukemia (ALL)	yes	yes	Were tests for molecular markers performed? (e.g. PCR, NGS) (between diagnosis and last evaluation)	No,Unknown,yes		Were tests for molecular markers performed? (e.g. PCR, NGS) (between diagnosis or relapse and last evaluation)	No,Unknown,yes	
PRE174	Pre-Transplant	Disease Classification	Acute Lymphoblastic leukemia (ALL)	yes	yes	BCR / ABL	Negative,Not Done,Positive		BCR / ABL	Negative,Not Done,Positive	
PRE175	Pre-Transplant	Disease Classification	Acute Lymphoblastic leukemia (ALL)	yes	yes	TEL-AML1 / AML1	Negative,Not Done,Positive		TEL-AML1 / AML1	Negative,Not Done,Positive	
PRE176	Pre-Transplant	Disease Classification	Acute Lymphoblastic leukemia (ALL)	yes	yes	Other molecular marker	Negative,Not Done,Positive		Other molecular marker	Negative,Not Done,Positive	
PRE177	Pre-Transplant	Disease Classification	Acute Lymphoblastic leukemia (ALL)	yes	yes	Specify other molecular marker:	open text		Specify other molecular marker:	open text	
PRE178	Pre-Transplant	Disease Classification	Acute Lymphoblastic leukemia (ALL)	yes	yes	Were cytogenetics tested (karyotyping or FISH)? (at last evaluation)	No,Unknown,yes		Were cytogenetics tested (karyotyping or FISH)? (at last evaluation)	No,Unknown,yes	
PRE179	Pre-Transplant	Disease Classification	Acute Lymphoblastic leukemia (ALL)	yes	yes	Were cytogenetics tested via FISH?	No,Yes		Were cytogenetics tested via FISH?	No,Yes	
PRE180	Pre-Transplant	Disease Classification	Acute Lymphoblastic leukemia (ALL)	yes	yes	Results of tests	Abnormalities identified.No abnormalities		Results of tests	Abnormalities identified.No abnormalities	
PRE181	Pre-Transplant	Disease Classification	Acute Lymphoblastic leukemia (ALL)	yes	yes	International System for Human Cytogenetic Nomenclature (ISCN) compatible string:	open text		International System for Human Cytogenetic Nomenclature (ISCN) compatible string:	open text	
PRE182	Pre-Transplant	Disease Classification	Acute Lymphoblastic Leukemia (ALL)	yes	yes	Specify number of distinct cytogenetic abnormalities	Four or more (4 or more),One (1),Three (3),Two (2)		Specify number of distinct cytogenetic abnormalities	Four or more (4 or more),One (1),Three (3),Two (2)	
PRE183	Pre-Transplant	Disease Classification	Acute Lymphoblastic leukemia (ALL)	yes	yes	Specify abnormalities (check all that apply)	11q23) any abnormality,12p any abnormality,9p any abnormality,add(14q),del(12p) / 12p-del(6q) / 6q-del(9p) / 9p-Hyperdiploid (< 50),Hypodiploid (< 46),JAMP21-7,Other abnormality (1;1-19,t(10;14),t(11;14),t(12;21),t(2;8),t(4;11),t(5;14),t(8;14),t(8;22),t(9;22),+17,+21,+4,-8		Specify abnormalities (check all that apply)	11q23) any abnormality,12p any abnormality,9p any abnormality,add(14q),del(12p) / 12p-del(6q) / 6q-del(9p) / 9p-Hyperdiploid (< 50),Hypodiploid (< 46),JAMP21-7,Other abnormality (1;1-19,t(10;14),t(11;14),t(12;21),t(2;8),t(4;11),t(5;14),t(8;14),t(8;22),t(9;22),+17,+21,+4,-8	
PRE184	Pre-Transplant	Disease Classification	Acute Lymphoblastic leukemia (ALL)	yes	yes	Specify other abnormality:	open text		Specify other abnormality:	open text	
PRE185	Pre-Transplant	Disease Classification	Acute Lymphoblastic leukemia (ALL)	yes	yes	Were cytogenetics tested via karyotyping? (at last evaluation)	No,Yes		Were cytogenetics tested via karyotyping? (at last evaluation)	No,Yes	
PRE186	Pre-Transplant	Disease Classification	Acute Lymphoblastic leukemia (ALL)	yes	yes	Results of tests	Abnormalities identified.No abnormalities.No evaluable metaphases		Results of tests	Abnormalities identified.No abnormalities.No evaluable metaphases	
PRE187	Pre-Transplant	Disease Classification	Acute Lymphoblastic leukemia (ALL)	yes	yes	International System for Human Cytogenetic Nomenclature (ISCN) compatible string:	open text		International System for Human Cytogenetic Nomenclature (ISCN) compatible string:	open text	
PRE188	Pre-Transplant	Disease Classification	Acute Lymphoblastic leukemia (ALL)	yes	yes	Specify number of distinct cytogenetic abnormalities	Four or more (4 or more),One (1),Three (3),Two (2)		Specify number of distinct cytogenetic abnormalities	Four or more (4 or more),One (1),Three (3),Two (2)	
PRE189	Pre-Transplant	Disease Classification	Acute Lymphoblastic leukemia (ALL)	yes	yes	Specify abnormalities (check all that apply)	11q23) any abnormality,12p any abnormality,9p any abnormality,add(14q),del(12p) / 12p-del(6q) / 6q-del(9p) / 9p-Hyperdiploid (< 50),Hypodiploid (< 46),JAMP21-7,Other abnormality (1;1-19,t(10;14),t(11;14),t(12;21),t(2;8),t(4;11),t(5;14),t(8;14),t(8;22),t(9;22),+17,+21,+4,-8		Specify abnormalities (check all that apply)	11q23) any abnormality,12p any abnormality,9p any abnormality,add(14q),del(12p) / 12p-del(6q) / 6q-del(9p) / 9p-Hyperdiploid (< 50),Hypodiploid (< 46),JAMP21-7,Other abnormality (1;1-19,t(10;14),t(11;14),t(12;21),t(2;8),t(4;11),t(5;14),t(8;14),t(8;22),t(9;22),+17,+21,+4,-8	
PRE190	Pre-Transplant	Disease Classification	Acute Lymphoblastic leukemia (ALL)	yes	yes	Specify other abnormality:	open text		Specify other abnormality:	open text	
PRE191	Pre-Transplant	Disease Classification	Acute Lymphoblastic leukemia (ALL)	yes	yes	Was documentation submitted to the CIBMTR? (e.g. cytogenetic or FISH report)	No,Yes		Was documentation submitted to the CIBMTR? (e.g. cytogenetic or FISH report)	No,Yes	
PRE192	Pre-Transplant	Disease Classification	Acute Lymphoblastic leukemia (ALL)	yes	yes	Were tests for molecular markers performed? (e.g. PCR, NGS) (at last evaluation)	No,Unknown,yes		Were tests for molecular markers performed? (e.g. PCR, NGS) (at last evaluation)	No,Unknown,yes	
PRE193	Pre-Transplant	Disease Classification	Acute Lymphoblastic leukemia (ALL)	yes	yes	BCR / ABL	Negative,Not Done,Positive		BCR / ABL	Negative,Not Done,Positive	
PRE194	Pre-Transplant	Disease Classification	Acute Lymphoblastic leukemia (ALL)	yes	yes	TEL-AML1 / AML1	Negative,Not Done,Positive		TEL-AML1 / AML1	Negative,Not Done,Positive	
PRE195	Pre-Transplant	Disease Classification	Acute Lymphoblastic leukemia (ALL)	yes	yes	Other molecular marker	Negative,Not Done,Positive		Other molecular marker	Negative,Not Done,Positive	
PRE196	Pre-Transplant	Disease Classification	Acute Lymphoblastic leukemia (ALL)	yes	yes	Specify other molecular marker:	open text		Specify other molecular marker:	open text	
PRE197	Pre-Transplant	Disease Classification	Acute Lymphoblastic leukemia (ALL)	yes	no	Did the recipient have central nervous system leukemia at any time prior to the start of the preparative regimen / infusion?	No,Unknown,yes		Did the recipient have central nervous system leukemia at any time prior to the start of the preparative regimen / infusion?	No,Unknown,yes	
PRE198	Pre-Transplant	Disease Classification	Acute Lymphoblastic leukemia (ALL)	yes	no	What was the disease status?	1st complete remission (include CR),1st relapse,2nd complete remission,2nd relapse, > 3rd complete remission, >3rd relapse,No treatment,Primary induction failure		What was the disease status?	1st complete remission (include CR),1st relapse,2nd complete remission,2nd relapse, > 3rd complete remission, >3rd relapse,No treatment,Primary induction failure	
PRE199	Pre-Transplant	Disease Classification	Acute Lymphoblastic leukemia (ALL)	yes	no	How many cycles of induction therapy were required to achieve 1st complete remission?	1,2, > 3		How many cycles of induction therapy were required to achieve 1st complete remission?	1,2, > 3	
PRE200	Pre-Transplant	Disease Classification	Acute Lymphoblastic leukemia (ALL)	yes	no	Date of most recent relapse:	YYYY/MM/DD		Date of most recent relapse:	YYYY/MM/DD	
PRE201	Pre-Transplant	Disease Classification	Acute Lymphoblastic leukemia (ALL)	yes	no	Date assessed:	YYYY/MM/DD		Date assessed:	YYYY/MM/DD	
PRE202	Pre-Transplant	Disease Classification	Acute Leukemia Lineage and Other Myeloid Neoplasms	yes	no	Specify acute leukemias of ambiguous lineage and other myeloid neoplasm classification	Acute undifferentiated leukemia,Blastic plasmacytoid dendritic cell neoplasm,Mixed phenotype acute leukemia, B/myeloid, NOS,Mixed phenotype acute leukemia (MPAL) with t(9;22)(q34.1;q11.2), BCR-ABL1,Mixed phenotype acute leukemia with t(6;14q23.3); RMT2A rearranged,Mixed phenotype acute leukemia, T/myeloid, NOS,Other acute leukemia of ambiguous lineage or myeloid neoplasm		Specify acute leukemias of ambiguous lineage and other myeloid neoplasm classification	Acute undifferentiated leukemia,Blastic plasmacytoid dendritic cell neoplasm,Mixed phenotype acute leukemia, B/myeloid, NOS,Mixed phenotype acute leukemia (MPAL) with t(9;22)(q34.1;q11.2), BCR-ABL1,Mixed phenotype acute leukemia with t(6;14q23.3); RMT2A rearranged,Mixed phenotype acute leukemia, T/myeloid, NOS,Other acute leukemia of ambiguous lineage or myeloid neoplasm	

Item ID	Time Point	Information Collection Domain Sub-Type	Information Collection Domain Additional Sub-Domain	Response required if additional sub-Domain applies	Information Collection may be requested multiple times	Current Information Collection Data Element (if applicable)	Current Information Collection Data Element Response Option(s)	Information Collection update:	Proposed Information Collection Data Element (if applicable)	Proposed Information Collection Data Element Response Option(s)	Rationale for information Collection Update
PRE203	Pre-Transplant	Disease Classification	Acute Leukemias of Ambiguous Lineage and Other Myeloid Neoplasms	yes	no	Specify other acute leukemia of ambiguous lineage or myeloid neoplasm:	open text		Specify other acute leukemia of ambiguous lineage or myeloid neoplasm:	open text	
PRE204	Pre-Transplant	Disease Classification	Acute Leukemias of Ambiguous Lineage and Other Myeloid Neoplasms	yes	no	What was the disease status? (based on hematological test results)	1st complete remission (no previous marrow or extramedullary relapse),1st relapse,2nd complete remission,2nd relapse, ≥ 3rd complete remission, ≥ 3rd relapse,No treatment,Primary induction failure		What was the disease status? (based on hematological test results)	1st complete remission (no previous marrow or extramedullary relapse),1st relapse,2nd complete remission,2nd relapse, ≥ 3rd complete remission, ≥ 3rd relapse,No treatment,Primary induction failure	
PRE205	Pre-Transplant	Disease Classification	Acute Leukemias of Ambiguous Lineage and Other Myeloid Neoplasms	yes	no	Date assessed:	YYYY/MM/DD		Date assessed:	YYYY/MM/DD	
PRE206	Pre-Transplant	Disease Classification	Chronic Myelogenous Leukemia (CML)	yes	no	Was therapy given prior to this HCT?	no,yes		Was therapy given prior to this HCT?	no,yes	
PRE207	Pre-Transplant	Disease Classification	Chronic Myelogenous Leukemia (CML)	yes	no	Combination chemotherapy	no,yes		Combination chemotherapy	no,yes	
PRE208	Pre-Transplant	Disease Classification	Chronic Myelogenous Leukemia (CML)	yes	no	Hydroxyurea (Droxia, Hydroa)	no,yes		Hydroxyurea (Droxia, Hydroa)	no,yes	
PRE209	Pre-Transplant	Disease Classification	Chronic Myelogenous Leukemia (CML)	yes	no	Tyrosine kinase inhibitor (e.g.imatinib mesylate, dasatinib, nilotinib)	no,yes		Tyrosine kinase inhibitor (e.g.imatinib mesylate, dasatinib, nilotinib)	no,yes	
PRE210	Pre-Transplant	Disease Classification	Chronic Myelogenous Leukemia (CML)	yes	no	Interferon-α/alpha (Intron, Roferon) (includes PEG)	no,yes		Interferon-α/alpha (Intron, Roferon) (includes PEG)	no,yes	
PRE211	Pre-Transplant	Disease Classification	Chronic Myelogenous Leukemia (CML)	yes	no	Other therapy	no,yes		Other therapy	no,yes	
PRE212	Pre-Transplant	Disease Classification	Chronic Myelogenous Leukemia (CML)	yes	no	Specify other therapy:	open text		Specify other therapy:	open text	
PRE213	Pre-Transplant	Disease Classification	Chronic Myelogenous Leukemia (CML)	yes	no	What was the disease status?	Accelerated phase,Blast phase,Complete hematologic response (CHR) preceded by accelerated phase and/or blast phase,Complete hematologic response (CHR) preceded only by chronic phase,Chronic phase		What was the disease status?	Accelerated phase,Blast phase,Complete hematologic response (CHR) preceded by accelerated phase and/or blast phase,Complete hematologic response (CHR) preceded only by chronic phase,Chronic phase	
PRE214	Pre-Transplant	Disease Classification	Chronic Myelogenous Leukemia (CML)	yes	no	Specify level of response	Complete cytogenetic response (CCR),Complete molecular remission (CMR),Minimal cytogenetic response,Minor cytogenetic response,Major molecular remission (MMR),No cytogenetic response (No Cyt),Partial cytogenetic response (PCyR)		Specify level of response	Complete cytogenetic response (CCR),Complete molecular remission (CMR),Minimal cytogenetic response,Minor cytogenetic response,Major molecular remission (MMR),No cytogenetic response (No Cyt),Partial cytogenetic response (PCyR)	
PRE215	Pre-Transplant	Disease Classification	Chronic Myelogenous Leukemia (CML)	yes	no	Number	1st,2nd,3rd or higher		Number	1st,2nd,3rd or higher	
PRE216	Pre-Transplant	Disease Classification	Chronic Myelogenous Leukemia (CML)	yes	no	Date assessed:	YYYY/MM/DD		Date assessed:	YYYY/MM/DD	
PRE217	Pre-Transplant	Disease Classification	Myelodysplastic Syndrome (MDS)	yes	no	What was the MDS subtype at diagnosis? - If transformed to AML, indicate AML as primary disease; also complete AML Disease Classification questions	Atypical chronic myeloid leukemia (ACML),BCR-ABL1-Chronic myelomonocytic leukemia (CMML),Juvenile myelomonocytic leukemia (JMML)/CMML,Myelodysplastic syndrome with isolated del(5q),Myelodysplastic syndrome with multilineage dysplasia (MDS-M4),MDS / MPN with ring sideroblasts and thrombocytosis (MDS / MPN-RS-1),Myelodysplastic syndrome / myeloproliferative neoplasm,unclassifiable syndrome with single lineage dysplasia (MDS-SL),Myelodysplastic syndrome (MDS),unclassifiable refractory cytopenia of childhood, Myelodysplastic syndrome with excess blasts (MDS-EB) , MDS with excess blasts-1 (MDS-EB-1),MDS with excess blasts-2 (MDS-EB-2), Myelodysplastic syndrome with ring sideroblasts : MDS-RS with multilineage dysplasia (MDS-RS-M4),MDS-RS with single lineage dysplasia (MDS-RS-SL),Myelodysplastic		What was the MDS subtype at diagnosis? - If transformed to AML, indicate AML as primary disease; also complete AML Disease Classification questions	Atypical chronic myeloid leukemia (ACML),BCR-ABL1-Chronic myelomonocytic leukemia (CMML),Juvenile myelomonocytic leukemia (JMML)/CMML,Myelodysplastic syndrome with isolated del(5q),Myelodysplastic syndrome with multilineage dysplasia (MDS-M4),MDS / MPN with ring sideroblasts and thrombocytosis (MDS / MPN-RS-1),Myelodysplastic syndrome / myeloproliferative neoplasm,unclassifiable syndrome with single lineage dysplasia (MDS-SL),Myelodysplastic syndrome (MDS),unclassifiable refractory cytopenia of childhood, Myelodysplastic syndrome with excess blasts (MDS-EB) , MDS with excess blasts-1 (MDS-EB-1),MDS with excess blasts-2 (MDS-EB-2), Myelodysplastic syndrome with ring sideroblasts : MDS-RS with multilineage dysplasia (MDS-RS-M4),MDS-RS with single lineage dysplasia (MDS-RS-SL),Myelodysplastic	
PRE218	Pre-Transplant	Disease Classification	Myelodysplastic Syndrome (MDS)	yes	no	Specify Myelodysplastic syndrome, unclassifiable (MDS-U)	MDS-U with 1% blood blasts,MDS-U based on defining cytogenetic abnormality,MDS-U with single lineage dysplasia and pancytopenia		Specify Myelodysplastic syndrome, unclassifiable (MDS-U)	MDS-U with 1% blood blasts,MDS-U based on defining cytogenetic abnormality,MDS-U with single lineage dysplasia and pancytopenia	
PRE219	Pre-Transplant	Disease Classification	Myelodysplastic Syndrome (MDS)	yes	no	Was documentation submitted to the CIBMTR? (e.g. cytogenetic or FISH report)	No,Yes		Was documentation submitted to the CIBMTR? (e.g. cytogenetic or FISH report)	No,Yes	
PRE220	Pre-Transplant	Disease Classification	Myelodysplastic Syndrome (MDS)	yes	no	Was the disease MDS therapy related?	No,Unknown,yes		Was the disease MDS therapy related?	No,Unknown,yes	
PRE221	Pre-Transplant	Disease Classification	Myelodysplastic Syndrome (MDS)	yes	no	Did the recipient have a predisposing condition?	No,Unknown,yes		Did the recipient have a predisposing condition?	No,Unknown,yes	
PRE222	Pre-Transplant	Disease Classification	Myelodysplastic Syndrome (MDS)	yes	no	Specify condition	Acuteleukemia,DOXA1-associated familial MDS,Fanconi anemia,GATA2 deficiency (including Emberger syndrome, MonoMac syndrome, DCML deficiency),J1,Fraumeni Syndrome,Other condition,Paroxysmal nocturnal hemoglobinuria,Diamond-Blackfan Anemia,RUNX1 deficiency (previously Tamarit platelet disorder with propensity to myeloid malignancies"), SAMD9- associated familial MDS,Bowen-Kim-Diamond syndrome,telomere biology disorder (including Dyskeratosis congenita)		Specify condition	Acuteleukemia,DOXA1-associated familial MDS,Fanconi anemia,GATA2 deficiency (including Emberger syndrome, MonoMac syndrome, DCML deficiency),J1,Fraumeni Syndrome,Other condition,Paroxysmal nocturnal hemoglobinuria,Diamond-Blackfan Anemia,RUNX1 deficiency (previously Tamarit platelet disorder with propensity to myeloid malignancies"), SAMD9- associated familial MDS,Bowen-Kim-Diamond syndrome,telomere biology disorder (including Dyskeratosis congenita)	
PRE223	Pre-Transplant	Disease Classification	Myelodysplastic Syndrome (MDS)	yes	no	Specify other condition:	open text		Specify other condition:	open text	
PRE224	Pre-Transplant	Disease Classification	Myelodysplastic Syndrome (MDS)	yes	yes	Date CBC drawn:	YYYY/MM/DD		Date CBC drawn:	YYYY/MM/DD	
PRE225	Pre-Transplant	Disease Classification	Myelodysplastic Syndrome (MDS)	yes	yes	Blasts in bone marrow	Known,Unknown		Blasts in bone marrow	Known,Unknown	
PRE226	Pre-Transplant	Disease Classification	Myelodysplastic Syndrome (MDS)	yes	yes	Blasts in bone marrow	___ ____ %		Blasts in bone marrow	___ ____ %	
PRE227	Pre-Transplant	Disease Classification	Myelodysplastic Syndrome (MDS)	yes	yes	Were cytogenetics tested (karyotyping or FISH)?	No,Unknown,yes		Were cytogenetics tested (karyotyping or FISH)?	No,Unknown,yes	
PRE228	Pre-Transplant	Disease Classification	Myelodysplastic Syndrome (MDS)	yes	yes	Were cytogenetics tested via FISH?	No,Yes		Were cytogenetics tested via FISH?	No,Yes	
PRE229	Pre-Transplant	Disease Classification	Myelodysplastic Syndrome (MDS)	yes	yes	Sample source	Peripheral blood,Bone marrow		Sample source	Peripheral blood,Bone marrow	
PRE230	Pre-Transplant	Disease Classification	Myelodysplastic Syndrome (MDS)	yes	yes	Results of tests	Abnormalities identified,No abnormalities		Results of tests	Abnormalities identified,No abnormalities	
PRE231	Pre-Transplant	Disease Classification	Myelodysplastic Syndrome (MDS)	yes	yes	International System for Human Cytogenetic Nomenclature (ISCN) compatible string:	open text		International System for Human Cytogenetic Nomenclature (ISCN) compatible string:	open text	
PRE232	Pre-Transplant	Disease Classification	Myelodysplastic Syndrome (MDS)	yes	yes	Specify number of distinct cytogenetic abnormalities	Four or more (4 or more),One (1),Three (3),Two (2)		Specify number of distinct cytogenetic abnormalities	Four or more (4 or more),One (1),Three (3),Two (2)	
PRE233	Pre-Transplant	Disease Classification	Myelodysplastic Syndrome (MDS)	yes	yes	Specify abnormalities (check all that apply)	del(11q) / 11q-del(12p) / 12p-del(20q) / 20q-del(3q) / 3q-del(5q) / 5q-del(7q) / 7q-del(9q) / 9q-del(13q) / 13q-117q,inv(3),13-20,5-7,7-Other abnormality,0(12),0(11),0(12),1(1),0(2,21),0(3-3),0(6,9)+,-t(9-8)		Specify abnormalities (check all that apply)	del(11q) / 11q-del(12p) / 12p-del(20q) / 20q-del(3q) / 3q-del(5q) / 5q-del(7q) / 7q-del(9q) / 9q-del(13q) / 13q-117q,inv(3),13-20,5-7,7-Other abnormality,0(12),0(11),0(12),1(1),0(2,21),0(3-3),0(6,9)+,-t(9-8)	
PRE234	Pre-Transplant	Disease Classification	Myelodysplastic Syndrome (MDS)	yes	yes	Specify other abnormality:	open text		Specify other abnormality:	open text	
PRE235	Pre-Transplant	Disease Classification	Myelodysplastic Syndrome (MDS)	yes	yes	Was documentation submitted to the CIBMTR? (e.g. cytogenetic or FISH report)	No,Yes		Was documentation submitted to the CIBMTR? (e.g. cytogenetic or FISH report)	No,Yes	
PRE236	Pre-Transplant	Disease Classification	Myelodysplastic Syndrome (MDS)	yes	yes	Were cytogenetics tested via karyotyping?	No,Yes		Were cytogenetics tested via karyotyping?	No,Yes	
PRE237	Pre-Transplant	Disease Classification	Myelodysplastic Syndrome (MDS)	yes	yes	Sample source	Peripheral blood,Bone marrow		Sample source	Peripheral blood,Bone marrow	
PRE238	Pre-Transplant	Disease Classification	Myelodysplastic Syndrome (MDS)	yes	yes	Results of tests	Abnormalities identified,No abnormalities,No evaluable metaphases		Results of tests	Abnormalities identified,No abnormalities,No evaluable metaphases	
PRE239	Pre-Transplant	Disease Classification	Myelodysplastic Syndrome (MDS)	yes	yes	International System for Human Cytogenetic Nomenclature (ISCN) compatible string:	open text		International System for Human Cytogenetic Nomenclature (ISCN) compatible string:	open text	
PRE240	Pre-Transplant	Disease Classification	Myelodysplastic Syndrome (MDS)	yes	yes	Specify number of distinct cytogenetic abnormalities	Four or more (4 or more),One (1),Three (3),Two (2)		Specify number of distinct cytogenetic abnormalities	Four or more (4 or more),One (1),Three (3),Two (2)	
PRE241	Pre-Transplant	Disease Classification	Myelodysplastic Syndrome (MDS)	yes	yes	Specify abnormalities (check all that apply)	del(11q) / 11q-del(12p) / 12p-del(20q) / 20q-del(3q) / 3q-del(5q) / 5q-del(7q) / 7q-del(9q) / 9q-del(13q) / 13q-117q,inv(3),13-20,5-7,7-Other abnormality,0(12),0(11),0(12),1(1),0(2,21),0(3-3),0(6,9)+,-t(9-8)		Specify abnormalities (check all that apply)	del(11q) / 11q-del(12p) / 12p-del(20q) / 20q-del(3q) / 3q-del(5q) / 5q-del(7q) / 7q-del(9q) / 9q-del(13q) / 13q-117q,inv(3),13-20,5-7,7-Other abnormality,0(12),0(11),0(12),1(1),0(2,21),0(3-3),0(6,9)+,-t(9-8)	
PRE242	Pre-Transplant	Disease Classification	Myelodysplastic Syndrome (MDS)	yes	yes	Specify other abnormality:	open text		Specify other abnormality:	open text	
PRE243	Pre-Transplant	Disease Classification	Myelodysplastic Syndrome (MDS)	yes	yes	Was documentation submitted to the CIBMTR? (e.g. cytogenetic or FISH report)	No,Yes		Was documentation submitted to the CIBMTR? (e.g. cytogenetic or FISH report)	No,Yes	

Item ID	Time Point	Information Collection Domain Sub-Type	Information Collection Domain Additional Sub-Domain	Response required if additional sub-Domain applies	Information Collection may be requested multiple times	Current Information Collection Data Element (if applicable)	Current Information Collection Data Element Response Option(s)	Information Collection update:	Proposed Information Collection Data Element (if applicable)	Proposed Information Collection Data Element Response Option(s)	Rationale for information Collection Update
PRE244	Pre-Transplant	Disease Classification	Myelodysplastic Syndrome (MDS)	yes	yes	Did the recipient progress or transform to a different MDS subtype or AML between diagnosis and the start of the preparative regimen/ infusion?	No,Yes		Did the recipient progress or transform to a different MDS subtype or AML between diagnosis and the start of the preparative regimen/ infusion?	No,Yes	
PRE245	Pre-Transplant	Disease Classification	Myelodysplastic Syndrome (MDS)	yes	yes	Specify the MDS subtype or AML after transformation	Transformed to AML, Chronic myelomonocytic leukemia (CMML), Myelodysplastic syndrome with isolated del(5q), Myelodysplastic syndrome with multilineage dysplasia (MDS-MLD), MDS / MPN with ring sideroblasts and thrombocytosis (MDS / MPN-RS-T), Myelodysplastic syndrome / myeloproliferative neoplasm, unclassifiable Myelodysplastic syndrome with single lineage dysplasia (MDS-SLD), Myelodysplastic syndrome (MDS), unclassifiable, refractory cytopenia of childhood, Myelodysplastic Syndrome with excess blasts (MDS-EB) , MDS with excess blasts-1 (MDS-EB-1), MDS with excess blasts-2 (MDS-EB-2), Myelodysplastic syndrome with ring sideroblasts , MDS-RS with multilineage dysplasia (MDS-RS-MLD), MDS-RS with single lineage dysplasia (MDS-RS-SLD)		Specify the MDS subtype or AML after transformation	Transformed to AML, Chronic myelomonocytic leukemia (CMML), Myelodysplastic syndrome with isolated del(5q), Myelodysplastic syndrome with multilineage dysplasia (MDS-MLD), MDS / MPN with ring sideroblasts and thrombocytosis (MDS / MPN-RS-T), Myelodysplastic syndrome / myeloproliferative neoplasm, unclassifiable Myelodysplastic syndrome with single lineage dysplasia (MDS-SLD), Myelodysplastic syndrome (MDS), unclassifiable, refractory cytopenia of childhood, Myelodysplastic Syndrome with excess blasts (MDS-EB) , MDS with excess blasts-1 (MDS-EB-1), MDS with excess blasts-2 (MDS-EB-2), Myelodysplastic syndrome with ring sideroblasts , MDS-RS with multilineage dysplasia (MDS-RS-MLD), MDS-RS with single lineage dysplasia (MDS-RS-SLD)	
PRE246	Pre-Transplant	Disease Classification	Myelodysplastic Syndrome (MDS)	yes	yes	Specify Myelodysplastic syndrome, unclassifiable (MDS-U)	MDS-U with 1% blood blast, MDS-U based on defining cytogenetic abnormality, MDS-U with single lineage dysplasia and pancytopenia		Specify Myelodysplastic syndrome, unclassifiable (MDS-U)	MDS-U with 1% blood blast, MDS-U based on defining cytogenetic abnormality, MDS-U with single lineage dysplasia and pancytopenia	
PRE247	Pre-Transplant	Disease Classification	Myelodysplastic Syndrome (MDS)	yes	yes	Specify the date of the most recent transformation:	YYYY/MM/DD		Specify the date of the most recent transformation:	YYYY/MM/DD	
PRE248	Pre-Transplant	Disease Classification	Myelodysplastic Syndrome (MDS)	yes	yes	Date of MDS diagnosis:	YYYY/MM/DD		Date of MDS diagnosis:	YYYY/MM/DD	
PRE249	Pre-Transplant	Disease Classification	Myelodysplastic Syndrome (MDS)	yes	yes	Date CBC drawn:	YYYY/MM/DD		Date CBC drawn:	YYYY/MM/DD	
PRE250	Pre-Transplant	Disease Classification	Myelodysplastic Syndrome (MDS)	yes	yes	Blasts in bone marrow	Known, Unknown		Blasts in bone marrow	Known, Unknown	
PRE251	Pre-Transplant	Disease Classification	Myelodysplastic Syndrome (MDS)	yes	yes	Blasts in bone marrow	___ ____ %		Blasts in bone marrow	___ ____ %	
PRE252	Pre-Transplant	Disease Classification	Myelodysplastic Syndrome (MDS)	yes	yes	Were cytogenetics tested (karyotyping or FISH)?	No, Unknown, yes		Were cytogenetics tested (karyotyping or FISH)?	No, Unknown, yes	
PRE253	Pre-Transplant	Disease Classification	Myelodysplastic Syndrome (MDS)	yes	yes	Were cytogenetics tested via FISH?	No, Yes		Were cytogenetics tested via FISH?	No, Yes	
PRE254	Pre-Transplant	Disease Classification	Myelodysplastic Syndrome (MDS)	yes	yes	Sample source	Peripheral blood, Bone marrow		Sample source	Peripheral blood, Bone marrow	
PRE255	Pre-Transplant	Disease Classification	Myelodysplastic Syndrome (MDS)	yes	yes	Results of tests	Abnormalities identified, No abnormalities		Results of tests	Abnormalities identified, No abnormalities	
PRE256	Pre-Transplant	Disease Classification	Myelodysplastic Syndrome (MDS)	yes	yes	International System for Human Cytogenetic Nomenclature (ISCN) compatible string:	open text		International System for Human Cytogenetic Nomenclature (ISCN) compatible string:	open text	
PRE257	Pre-Transplant	Disease Classification	Myelodysplastic Syndrome (MDS)	yes	yes	Specify number of distinct cytogenetic abnormalities	Four or more (4 or more), One (1), Three (3), Two (2)		Specify number of distinct cytogenetic abnormalities	Four or more (4 or more), One (1), Three (3), Two (2)	
PRE258	Pre-Transplant	Disease Classification	Myelodysplastic Syndrome (MDS)	yes	yes	Specify abnormalities (check all that apply)	del(11q) / t(1q; del(11p) / t(3p; del(3q) / 3q; del(3q) / 3q; del(5q) / 5q; del(7q) / 7q; del(9q) / 9q; del(13q) / 13q; t(17q; inv(3); -13; -20; -5; -7; Y, Other abnormality: t(1;3), t(11;16), t(12;11), t(3;21), t(3;3), t(6;9), +19; -8		Specify abnormalities (check all that apply)	del(11q) / t(1q; del(11p) / t(3p; del(3q) / 3q; del(3q) / 3q; del(5q) / 5q; del(7q) / 7q; del(9q) / 9q; del(13q) / 13q; t(17q; inv(3); -13; -20; -5; -7; Y, Other abnormality: t(1;3), t(11;16), t(12;11), t(3;21), t(3;3), t(6;9), +19; -8	
PRE259	Pre-Transplant	Disease Classification	Myelodysplastic Syndrome (MDS)	yes	yes	Specify other abnormality:	open text		Specify other abnormality:	open text	
PRE260	Pre-Transplant	Disease Classification	Myelodysplastic Syndrome (MDS)	yes	yes	Was documentation submitted to the CIBMTR (e.g. cytogenetic or FISH report)	No, Yes		Was documentation submitted to the CIBMTR (e.g. cytogenetic or FISH report)	No, Yes	
PRE261	Pre-Transplant	Disease Classification	Myelodysplastic Syndrome (MDS)	yes	yes	Were cytogenetics tested via karyotyping?	No, Yes		Were cytogenetics tested via karyotyping?	No, Yes	
PRE262	Pre-Transplant	Disease Classification	Myelodysplastic Syndrome (MDS)	yes	yes	Sample source	Peripheral blood, Bone marrow		Sample source	Peripheral blood, Bone marrow	
PRE263	Pre-Transplant	Disease Classification	Myelodysplastic Syndrome (MDS)	yes	yes	Results of tests	Abnormalities identified, No abnormalities, No evaluable metaphases		Results of tests	Abnormalities identified, No abnormalities, No evaluable metaphases	
PRE264	Pre-Transplant	Disease Classification	Myelodysplastic Syndrome (MDS)	yes	yes	International System for Human Cytogenetic Nomenclature (ISCN) compatible string:	open text		International System for Human Cytogenetic Nomenclature (ISCN) compatible string:	open text	
PRE265	Pre-Transplant	Disease Classification	Myelodysplastic Syndrome (MDS)	yes	yes	Specify number of distinct cytogenetic abnormalities	Four or more (4 or more), One (1), Three (3), Two (2)		Specify number of distinct cytogenetic abnormalities	Four or more (4 or more), One (1), Three (3), Two (2)	
PRE266	Pre-Transplant	Disease Classification	Myelodysplastic Syndrome (MDS)	yes	yes	Specify abnormalities (check all that apply)	del(11q) / t(1q; del(11p) / t(3p; del(3q) / 3q; del(3q) / 3q; del(5q) / 5q; del(7q) / 7q; del(9q) / 9q; del(13q) / 13q; t(17q; inv(3); -13; -20; -5; -7; Y, Other abnormality: t(1;3), t(11;16), t(12;11), t(3;21), t(3;3), t(6;9), +19; -8		Specify abnormalities (check all that apply)	del(11q) / t(1q; del(11p) / t(3p; del(3q) / 3q; del(3q) / 3q; del(5q) / 5q; del(7q) / 7q; del(9q) / 9q; del(13q) / 13q; t(17q; inv(3); -13; -20; -5; -7; Y, Other abnormality: t(1;3), t(11;16), t(12;11), t(3;21), t(3;3), t(6;9), +19; -8	
PRE267	Pre-Transplant	Disease Classification	Myelodysplastic Syndrome (MDS)	yes	yes	Specify other abnormality:	open text		Specify other abnormality:	open text	
PRE268	Pre-Transplant	Disease Classification	Myelodysplastic Syndrome (MDS)	yes	yes	Was documentation submitted to the CIBMTR (e.g. cytogenetic or FISH report)	No, Yes		Was documentation submitted to the CIBMTR (e.g. cytogenetic or FISH report)	No, Yes	
PRE269	Pre-Transplant	Disease Classification	Myelodysplastic Syndrome (MDS)	yes	no	What was the disease status?	Complete remission (CR), Hematologic improvement (HI), Not assessed, No response (NR) / stable disease (SD), Progression from hematologic improvement (Prog from HI), Relapse from complete remission (Rel from CR)		What was the disease status?	Complete remission (CR), Hematologic improvement (HI), Not assessed, No response (NR) / stable disease (SD), Progression from hematologic improvement (Prog from HI), Relapse from complete remission (Rel from CR)	
PRE270	Pre-Transplant	Disease Classification	Myelodysplastic Syndrome (MDS)	yes	no	Specify the cell line examined to determine H status	H-E, H-N, H-P		Specify the cell lines examined to determine H status	H-E, H-N, H-P	
PRE271	Pre-Transplant	Disease Classification	Myelodysplastic Syndrome (MDS)	yes	no	Specify transfusion dependence	Low-transfusion burden (LTB), Non-transfused (NTD)		Specify transfusion dependence	Low-transfusion burden (LTB), Non-transfused (NTD)	
PRE272	Pre-Transplant	Disease Classification	Myelodysplastic Syndrome (MDS)	yes	no	Date assessed:	YYYY/MM/DD		Date assessed:	YYYY/MM/DD	
PRE273	Pre-Transplant	Disease Classification	Myeloproliferative Neoplasms (MPN)	yes	no	What was the MPN subtype at diagnosis?	Chronic eosinophilic leukemia, not otherwise specified (NOS), Primary myelofibrosis (PMF), Chronic neutrophilic leukemia, Essential thrombocythemia, Myeloproliferative neoplasm (MPN), unclassifiable Myeloid / lymphoid neoplasm with FISH/ R rearrangement Myeloid / lymphoid neoplasm with FISH/ R rearrangement Myeloid / lymphoid neoplasm with FISH/ R rearrangement, Polycythemia vera (PCV), Mastocytosis , Cutaneous mastocytosis (CM), Systemic mastocytosis, Mast cell sarcoma (MCS)		What was the MPN subtype at diagnosis?	Chronic eosinophilic leukemia, not otherwise specified (NOS), Primary myelofibrosis (PMF), Chronic neutrophilic leukemia, Essential thrombocythemia, Myeloproliferative neoplasm (MPN), unclassifiable Myeloid / lymphoid neoplasm with FISH/ R rearrangement Myeloid / lymphoid neoplasm with FISH/ R rearrangement Myeloid / lymphoid neoplasm with FISH/ R rearrangement, Polycythemia vera (PCV), Mastocytosis , Cutaneous mastocytosis (CM), Systemic mastocytosis, Mast cell sarcoma (MCS)	
PRE274	Pre-Transplant	Disease Classification	Myeloproliferative Neoplasms (MPN)	yes	no	Specify systemic mastocytosis	Aggressive systemic mastocytosis (ASM), Indolent systemic mastocytosis (ISM), Mast cell leukemia (MCL), Systemic mastocytosis with an associated hematological neoplasm (SM-AHN), Smoldering systemic mastocytosis (SSM)		Specify systemic mastocytosis	Aggressive systemic mastocytosis (ASM), Indolent systemic mastocytosis (ISM), Mast cell leukemia (MCL), Systemic mastocytosis with an associated hematological neoplasm (SM-AHN), Smoldering systemic mastocytosis (SSM)	
PRE275	Pre-Transplant	Disease Classification	Myeloproliferative Neoplasms (MPN)	yes	no	Was documentation submitted to the CIBMTR (e.g. pathology report used for diagnosis)	No, Yes		Was documentation submitted to the CIBMTR (e.g. pathology report used for diagnosis)	No, Yes	
PRE276	Pre-Transplant	Disease Classification	Myeloproliferative Neoplasms (MPN)	yes	yes	Did the recipient have constitutional symptoms in six months before diagnosis? (Symptoms are >10% weight loss in 6 months, night sweats, or unexplained fever higher than 37.5 °C)	No, Unknown, Yes		Did the recipient have constitutional symptoms in six months before diagnosis? (Symptoms are >10% weight loss in 6 months, night sweats, or unexplained fever higher than 37.5 °C)	No, Unknown, Yes	
PRE277	Pre-Transplant	Disease Classification	Myeloproliferative Neoplasms (MPN)	yes	yes	Date CBC drawn:	YYYY/MM/DD		Date CBC drawn:	YYYY/MM/DD	
PRE278	Pre-Transplant	Disease Classification	Myeloproliferative Neoplasms (MPN)	yes	yes	Blasts in bone marrow	Known, Unknown		Blasts in bone marrow	Known, Unknown	
PRE279	Pre-Transplant	Disease Classification	Myeloproliferative Neoplasms (MPN)	yes	yes	Blasts in bone marrow	___ ____ %		Blasts in bone marrow	___ ____ %	
PRE280	Pre-Transplant	Disease Classification	Myeloproliferative Neoplasms (MPN)	yes	yes	Were tests for driver mutations performed?	No, Unknown, Yes		Were tests for driver mutations performed?	No, Unknown, Yes	
PRE281	Pre-Transplant	Disease Classification	Myeloproliferative Neoplasms (MPN)	yes	yes	JAK2	Negative, Not done, Positive		JAK2	Negative, Not done, Positive	
PRE282	Pre-Transplant	Disease Classification	Myeloproliferative Neoplasms (MPN)	yes	yes	JAK2 V617F	Negative, Not done, Positive		JAK2 V617F	Negative, Not done, Positive	
PRE283	Pre-Transplant	Disease Classification	Myeloproliferative Neoplasms (MPN)	yes	yes	JAK2 Exon 12	Negative, Not done, Positive		JAK2 Exon 12	Negative, Not done, Positive	
PRE284	Pre-Transplant	Disease Classification	Myeloproliferative Neoplasms (MPN)	yes	yes	CALR	Negative, Not done, Positive		CALR	Negative, Not done, Positive	
PRE285	Pre-Transplant	Disease Classification	Myeloproliferative Neoplasms (MPN)	yes	yes	CALR type 1	Negative, Not done, Positive		CALR type 1	Negative, Not done, Positive	

Item ID	Time Point	Information Collection Domain Sub-Type	Information Collection Domain Additional Sub-Domain	Response required if additional sub-Domain applies	Information Collection may be requested multiple times	Current Information Collection Data Element (if applicable)	Current Information Collection Data Element Response Option(s)	Information Collection update:	Proposed Information Collection Data Element (if applicable)	Proposed Information Collection Data Element Response Option(s)	Rationale for information Collection Update
PRE286	Pre-Transplant	Disease Classification	Myeloproliferative Neoplasms (MPN)	yes	yes	CALR type 2	Negative,Not done,Positive		CALR type 2	Negative,Not done,Positive	
PRE287	Pre-Transplant	Disease Classification	Myeloproliferative Neoplasms (MPN)	yes	yes	Not defined	Negative,Not done,Positive		Not defined	Negative,Not done,Positive	
PRE288	Pre-Transplant	Disease Classification	Myeloproliferative Neoplasms (MPN)	yes	yes	MPL	Negative,Not done,Positive		MPL	Negative,Not done,Positive	
PRE289	Pre-Transplant	Disease Classification	Myeloproliferative Neoplasms (MPN)	yes	yes	CSF3R	Negative,Not done,Positive		CSF3R	Negative,Not done,Positive	
PRE290	Pre-Transplant	Disease Classification	Myeloproliferative Neoplasms (MPN)	yes	yes	Was documentation submitted to the CBMTR?	No,Yes		Was documentation submitted to the CBMTR?	No,Yes	
PRE291	Pre-Transplant	Disease Classification	Myeloproliferative Neoplasms (MPN)	yes	yes	Were cytogenetics tested (karyotyping or FISH)?	No,Unknown,yes		Were cytogenetics tested (karyotyping or FISH)?	No,Unknown,yes	
PRE292	Pre-Transplant	Disease Classification	Myeloproliferative Neoplasms (MPN)	yes	yes	Were cytogenetics tested via FISH?	No,Yes		Were cytogenetics tested via FISH?	No,Yes	
PRE293	Pre-Transplant	Disease Classification	Myeloproliferative Neoplasms (MPN)	yes	yes	Sample source	Peripheral blood,Bone marrow		Sample source	Peripheral blood,Bone marrow	
PRE294	Pre-Transplant	Disease Classification	Myeloproliferative Neoplasms (MPN)	yes	yes	Results of tests	Abnormalities identified,No abnormalities		Results of tests	Abnormalities identified,No abnormalities	
PRE295	Pre-Transplant	Disease Classification	Myeloproliferative Neoplasms (MPN)	yes	yes	International System for Human Cytogenetic Nomenclature (ISCN) compatible string	open text		International System for Human Cytogenetic Nomenclature (ISCN) compatible string	open text	
PRE296	Pre-Transplant	Disease Classification	Myeloproliferative Neoplasms (MPN)	yes	yes	Specify number of distinct cytogenetic abnormalities	Four or more (4 or more),One (1),Three (3),Two (2)		Specify number of distinct cytogenetic abnormalities	Four or more (4 or more),One (1),Three (3),Two (2)	
PRE297	Pre-Transplant	Disease Classification	Myeloproliferative Neoplasms (MPN)	yes	yes	Specify abnormalities (check all that apply)	del(11q) / 11q,del(12p) / 12p,del(20q) / 20q,del(5q) / 5q,del(7q) / 7q,del(11q) / 11q,del(13q) / 13q,del(11)(11)(q,mv(3),5-7,Y)Other abnormality,(11,any),(11,q23,any),(11p11.2,any),(11,q21,1,any),(16,9),+8,+9		Specify abnormalities (check all that apply)	del(11q) / 11q,del(12p) / 12p,del(20q) / 20q,del(5q) / 5q,del(7q) / 7q,del(11q) / 11q,del(13q) / 13q,del(11)(11)(q,mv(3),5-7,Y)Other abnormality,(11,any),(11,q23,any),(11p11.2,any),(11,q21,1,any),(16,9),+8,+9	
PRE298	Pre-Transplant	Disease Classification	Myeloproliferative Neoplasms (MPN)	yes	yes	Specify other abnormality:	open text		Specify other abnormality:	open text	
PRE299	Pre-Transplant	Disease Classification	Myeloproliferative Neoplasms (MPN)	yes	yes	Was documentation submitted to the CBMTR? (e.g. FISH report)	No,Yes		Was documentation submitted to the CBMTR? (e.g. FISH report)	No,Yes	
PRE300	Pre-Transplant	Disease Classification	Myeloproliferative Neoplasms (MPN)	yes	yes	Were cytogenetics tested via karyotyping?	No,Yes		Were cytogenetics tested via karyotyping?	No,Yes	
PRE301	Pre-Transplant	Disease Classification	Myeloproliferative Neoplasms (MPN)	yes	yes	Sample source	Peripheral blood,Bone marrow		Sample source	Peripheral blood,Bone marrow	
PRE302	Pre-Transplant	Disease Classification	Myeloproliferative Neoplasms (MPN)	yes	yes	Results of tests	Abnormalities identified,No abnormalities,No evaluable metaphases		Results of tests	Abnormalities identified,No abnormalities,No evaluable metaphases	
PRE303	Pre-Transplant	Disease Classification	Myeloproliferative Neoplasms (MPN)	yes	yes	International System for Human Cytogenetic Nomenclature (ISCN) compatible string	open text		International System for Human Cytogenetic Nomenclature (ISCN) compatible string	open text	
PRE304	Pre-Transplant	Disease Classification	Myeloproliferative Neoplasms (MPN)	yes	yes	Specify number of distinct cytogenetic abnormalities	Four or more (4 or more),One (1),Three (3),Two (2)		Specify number of distinct cytogenetic abnormalities	Four or more (4 or more),One (1),Three (3),Two (2)	
PRE305	Pre-Transplant	Disease Classification	Myeloproliferative Neoplasms (MPN)	yes	yes	Specify abnormalities (check all that apply)	del(11q) / 11q,del(12p) / 12p,del(20q) / 20q,del(5q) / 5q,del(7q) / 7q,del(11q) / 11q,del(13q) / 13q,del(11)(11)(q,mv(3),5-7,Y)Other abnormality,(11,any),(11,q23,any),(11p11.2,any),(11,q21,1,any),(16,9),+8,+9		Specify abnormalities (check all that apply)	del(11q) / 11q,del(12p) / 12p,del(20q) / 20q,del(5q) / 5q,del(7q) / 7q,del(11q) / 11q,del(13q) / 13q,del(11)(11)(q,mv(3),5-7,Y)Other abnormality,(11,any),(11,q23,any),(11p11.2,any),(11,q21,1,any),(16,9),+8,+9	
PRE306	Pre-Transplant	Disease Classification	Myeloproliferative Neoplasms (MPN)	yes	yes	Specify other abnormality:	open text		Specify other abnormality:	open text	
PRE307	Pre-Transplant	Disease Classification	Myeloproliferative Neoplasms (MPN)	yes	yes	Was documentation submitted to the CBMTR? (e.g. karyotyping report)	No,Yes		Was documentation submitted to the CBMTR? (e.g. karyotyping report)	No,Yes	
PRE308	Pre-Transplant	Disease Classification	Myeloproliferative Neoplasms (MPN)	yes	no	Did the recipient progress or transform to a different MPN subtype or AML between diagnosis and the start of the preparative regimen / infusion?	No,Yes		Did the recipient progress or transform to a different MPN subtype or AML between diagnosis and the start of the preparative regimen / infusion?	No,Yes	
PRE309	Pre-Transplant	Disease Classification	Myeloproliferative Neoplasms (MPN)	yes	no	Specify the MPN subtype or AML after transformation	Transformed to AML,Post-essential thrombocythemic myelofibrosis,Post-polycythemic myelofibrosis		Specify the MPN subtype or AML after transformation	Transformed to AML,Post-essential thrombocythemic myelofibrosis,Post-polycythemic myelofibrosis	
PRE310	Pre-Transplant	Disease Classification	Myeloproliferative Neoplasms (MPN)	yes	no	Specify the date of the most recent transformation:	YYYY/MM/DD		Specify the date of the most recent transformation:	YYYY/MM/DD	
PRE311	Pre-Transplant	Disease Classification	Myeloproliferative Neoplasms (MPN)	yes	no	Date of MPN diagnosis:	YYYY/MM/DD		Date of MPN diagnosis:	YYYY/MM/DD	
PRE312	Pre-Transplant	Disease Classification	Myeloproliferative Neoplasms (MPN)	yes	no	Specify transfusion dependence at last evaluation prior to the start of the preparative regimen / infusion	High-transfusion burden (HfB) (< 8 RBCs in 16 weeks; > 4 in 8 weeks),Low-transfusion burden (Lfb) (> 7 RBCs in 16 weeks in at least 2 transfusion episodes; maximum of 3 in 8 weeks),Non-transfused (NTD) (< 8 RBCs in 16 weeks)		Specify transfusion dependence at last evaluation prior to the start of the preparative regimen / infusion	High-transfusion burden (HfB) (< 8 RBCs in 16 weeks; > 4 in 8 weeks),Low-transfusion burden (Lfb) (> 7 RBCs in 16 weeks in at least 2 transfusion episodes; maximum of 3 in 8 weeks),Non-transfused (NTD) (< 8 RBCs in 16 weeks)	
PRE313	Pre-Transplant	Disease Classification	Myeloproliferative Neoplasms (MPN)	yes	yes	Did the recipient have constitutional symptoms in six months before last evaluation prior to the start of the preparative regimen / infusion? (symptoms are >10% weight loss in 6 months, night sweats, or unexplained fever higher than 37.5 °C)	No,Unknown,Yes		Did the recipient have constitutional symptoms in six months before last evaluation prior to the start of the preparative regimen / infusion? (symptoms are >10% weight loss in 6 months, night sweats, or unexplained fever higher than 37.5 °C)	No,Unknown,Yes	
PRE314	Pre-Transplant	Disease Classification	Myeloproliferative Neoplasms (MPN)	yes	no	Did the recipient have splenomegaly at last evaluation prior to the start of the preparative regimen / infusion?	No,Not applicable(splenectomy), Unknown,Yes		Did the recipient have splenomegaly at last evaluation prior to the start of the preparative regimen / infusion?	No,Not applicable(splenectomy), Unknown,Yes	
PRE315	Pre-Transplant	Disease Classification	Myeloproliferative Neoplasms (MPN)	yes	no	Specify the method used to measure spleen size	CT/MRI scan,Physical exam,Ultrasound		Specify the method used to measure spleen size	CT/MRI scan,Physical exam,Ultrasound	
PRE316	Pre-Transplant	Disease Classification	Myeloproliferative Neoplasms (MPN)	yes	no	Specify the spleen size:	____ centimeters below left costal margin		Specify the spleen size:	____ centimeters below left costal margin	
PRE317	Pre-Transplant	Disease Classification	Myeloproliferative Neoplasms (MPN)	yes	no	Specify the spleen size:	____ centimeters		Specify the spleen size:	____ centimeters	
PRE318	Pre-Transplant	Disease Classification	Myeloproliferative Neoplasms (MPN)	yes	no	Did the recipient have hepatomegaly at last evaluation prior to the start of the preparative regimen / infusion?	No,Unknown,yes		Did the recipient have hepatomegaly at last evaluation prior to the start of the preparative regimen / infusion?	No,Unknown,yes	
PRE319	Pre-Transplant	Disease Classification	Myeloproliferative Neoplasms (MPN)	yes	no	Specify the method used to measure liver size	CT/MRI scan,Physical exam,Ultrasound		Specify the method used to measure liver size	CT/MRI scan,Physical exam,Ultrasound	
PRE320	Pre-Transplant	Disease Classification	Myeloproliferative Neoplasms (MPN)	yes	no	Specify the liver size:	____ centimeters below right costal margin		Specify the liver size:	____ centimeters below right costal margin	
PRE321	Pre-Transplant	Disease Classification	Myeloproliferative Neoplasms (MPN)	yes	yes	Date CBC drawn:	YYYY/MM/DD		Date CBC drawn:	YYYY/MM/DD	
PRE322	Pre-Transplant	Disease Classification	Myeloproliferative Neoplasms (MPN)	yes	yes	Blasts in bone marrow	Known,Unknown		Blasts in bone marrow	Known,Unknown	
PRE323	Pre-Transplant	Disease Classification	Myeloproliferative Neoplasms (MPN)	yes	yes	Blasts in bone marrow	____ %		Blasts in bone marrow	____ %	
PRE324	Pre-Transplant	Disease Classification	Myeloproliferative Neoplasms (MPN)	yes	yes	Were tests for driver mutations performed?	No,Unknown,Yes		Were tests for driver mutations performed?	No,Unknown,Yes	
PRE325	Pre-Transplant	Disease Classification	Myeloproliferative Neoplasms (MPN)	yes	yes	JAK2	Negative,Not done,Positive		JAK2	Negative,Not done,Positive	
PRE326	Pre-Transplant	Disease Classification	Myeloproliferative Neoplasms (MPN)	yes	yes	JAK2 V617F	Negative,Not done,Positive		JAK2 V617F	Negative,Not done,Positive	
PRE327	Pre-Transplant	Disease Classification	Myeloproliferative Neoplasms (MPN)	yes	yes	CALR	Negative,Not done,Positive		CALR	Negative,Not done,Positive	
PRE328	Pre-Transplant	Disease Classification	Myeloproliferative Neoplasms (MPN)	yes	yes	CALR type 1	Negative,Not done,Positive		CALR type 1	Negative,Not done,Positive	

Item ID	Time Point	Information Collection Domain Sub-Type	Information Collection Domain Additional Sub-Domain	Response required if additional Sub-Domain applies	Information Collection may be requested multiple times	Current Information Collection Data Element (if applicable)	Current Information Collection Data Element Response Option(s)	Information Collection update:	Proposed Information Collection Data Element (if applicable)	Proposed Information Collection Data Element Response Option(s)	Rationale for information Collection Update
PRE329	Pre-Transplant	Disease Classification	Myeloproliferative Neoplasms (MPN)	yes	yes	CAIR type 2	Negative,Not done,Positive		CAIR type 2	Negative,Not done,Positive	
PRE330	Pre-Transplant	Disease Classification	Myeloproliferative Neoplasms (MPN)	yes	yes	Not defined	Negative,Not done,Positive		Not defined	Negative,Not done,Positive	
PRE331	Pre-Transplant	Disease Classification	Myeloproliferative Neoplasms (MPN)	yes	yes	MPL	Negative,Not done,Positive		MPL	Negative,Not done,Positive	
PRE332	Pre-Transplant	Disease Classification	Myeloproliferative Neoplasms (MPN)	yes	yes	CSF3R	Negative,Not done,Positive		CSF3R	Negative,Not done,Positive	
PRE333	Pre-Transplant	Disease Classification	Myeloproliferative Neoplasms (MPN)	yes	yes	Was documentation submitted to the CBMTB?	No,Yes		Was documentation submitted to the CBMTB?	No,Yes	
PRE334	Pre-Transplant	Disease Classification	Myeloproliferative Neoplasms (MPN)	yes	yes	Were cytogenetics tested (karyotyping or FISH)?	No,Unknown,yes		Were cytogenetics tested (karyotyping or FISH)?	No,Unknown,yes	
PRE335	Pre-Transplant	Disease Classification	Myeloproliferative Neoplasms (MPN)	yes	yes	Were cytogenetics tested via FISH?	No,Yes		Were cytogenetics tested via FISH?	No,Yes	
PRE336	Pre-Transplant	Disease Classification	Myeloproliferative Neoplasms (MPN)	yes	yes	Sample source	Peripheral blood,Bone marrow		Sample source	Peripheral blood,Bone marrow	
PRE337	Pre-Transplant	Disease Classification	Myeloproliferative Neoplasms (MPN)	yes	yes	Results of tests	Abnormalities identified,No abnormalities		Results of tests	Abnormalities identified,No abnormalities	
PRE338	Pre-Transplant	Disease Classification	Myeloproliferative Neoplasms (MPN)	yes	yes	International System for Human Cytogenetic Nomenclature (ISCN) compatible string	open text		International System for Human Cytogenetic Nomenclature (ISCN) compatible string	open text	
PRE339	Pre-Transplant	Disease Classification	Myeloproliferative Neoplasms (MPN)	yes	yes	Specify number of distinct cytogenetic abnormalities	Four or more (4 or more),One (1),Three (3),Two (2)		Specify number of distinct cytogenetic abnormalities	Four or more (4 or more),One (1),Three (3),Two (2)	
PRE340	Pre-Transplant	Disease Classification	Myeloproliferative Neoplasms (MPN)	yes	yes	Specify other abnormality:	open text		Specify other abnormality:	open text	
PRE341	Pre-Transplant	Disease Classification	Myeloproliferative Neoplasms (MPN)	yes	yes	Were cytogenetics tested via karyotyping?	No,Yes		Were cytogenetics tested via karyotyping?	No,Yes	
PRE342	Pre-Transplant	Disease Classification	Myeloproliferative Neoplasms (MPN)	yes	yes	Sample source	Peripheral blood,Bone marrow		Sample source	Peripheral blood,Bone marrow	
PRE343	Pre-Transplant	Disease Classification	Myeloproliferative Neoplasms (MPN)	yes	yes	Results of tests	Abnormalities identified,No abnormalities,No evaluable metaphases		Results of tests	Abnormalities identified,No abnormalities,No evaluable metaphases	
PRE344	Pre-Transplant	Disease Classification	Myeloproliferative Neoplasms (MPN)	yes	yes	International System for Human Cytogenetic Nomenclature (ISCN) compatible string	open text		International System for Human Cytogenetic Nomenclature (ISCN) compatible string	open text	
PRE345	Pre-Transplant	Disease Classification	Myeloproliferative Neoplasms (MPN)	yes	yes	Specify number of distinct cytogenetic abnormalities	Four or more (4 or more),One (1),Three (3),Two (2)		Specify number of distinct cytogenetic abnormalities	Four or more (4 or more),One (1),Three (3),Two (2)	
PRE346	Pre-Transplant	Disease Classification	Myeloproliferative Neoplasms (MPN)	yes	yes	Specify abnormalities (check all that apply)	del(11q) / 11q del(12p) / 12p del(20q) / 20q del(5q) / 5q del(7q) / 7q del(13q) / 13q del(11q) / 11q inv(3) / 5:7-Y/Other abnormality,(11,am),t(11q23,am),t(12p11,2,am),t(13q14,am),t(6,9),t(8,9)		Specify abnormalities (check all that apply)	del(11q) / 11q del(12p) / 12p del(20q) / 20q del(5q) / 5q del(7q) / 7q del(13q) / 13q del(11q) / 11q inv(3) / 5:7-Y/Other abnormality,(11,am),t(11q23,am),t(12p11,2,am),t(13q14,am),t(6,9),t(8,9)	
PRE347	Pre-Transplant	Disease Classification	Myeloproliferative Neoplasms (MPN)	yes	yes	Specify other abnormality:	open text		Specify other abnormality:	open text	
PRE348	Pre-Transplant	Disease Classification	Myeloproliferative Neoplasms (MPN)	yes	yes	Was documentation submitted to the CBMTB? (e.g. karyotyping report)	No,Yes		Was documentation submitted to the CBMTB? (e.g. karyotyping report)	No,Yes	
PRE349	Pre-Transplant	Disease Classification	Myeloproliferative Neoplasms (MPN)	yes	no	What was the disease status?	Clinical improvement (CI),Complete clinical remission (CR),Not assessed,Partial clinical remission (PR),Progressive disease,Relapse,Stable disease (SD)		What was the disease status?	Clinical improvement (CI),Complete clinical remission (CR),Not assessed,Partial clinical remission (PR),Progressive disease,Relapse,Stable disease (SD)	
PRE350	Pre-Transplant	Disease Classification	Myeloproliferative Neoplasms (MPN)	yes	no	Was an anemia response achieved?	No,Yes		Was an anemia response achieved?	No,Yes	
PRE351	Pre-Transplant	Disease Classification	Myeloproliferative Neoplasms (MPN)	yes	no	Was a spleen response achieved?	No,Yes		Was a spleen response achieved?	No,Yes	
PRE352	Pre-Transplant	Disease Classification	Myeloproliferative Neoplasms (MPN)	yes	no	Was a symptom response achieved?	No,Yes		Was a symptom response achieved?	No,Yes	
PRE353	Pre-Transplant	Disease Classification	Myeloproliferative Neoplasms (MPN)	yes	no	Date assessed:	YYYY/MM/DD		Date assessed:	YYYY/MM/DD	
PRE354	Pre-Transplant	Disease Classification	Myeloproliferative Neoplasms (MPN)	yes	no	Specify the cytogenetic response	Complete response (CR: Eradication of pre-existing abnormality,Not assessed,Not applicable,None of the above: Does not meet the CR or PR criteria, Partial response (PR) ≥ 50% reduction in abnormal metaphases,Re-emergence of pre-existing cytogenetic abnormality		Specify the cytogenetic response	Complete response (CR: Eradication of pre-existing abnormality,Not assessed,Not applicable,None of the above: Does not meet the CR or PR criteria, Partial response (PR) ≥ 50% reduction in abnormal metaphases,Re-emergence of pre-existing cytogenetic abnormality	
PRE355	Pre-Transplant	Disease Classification	Myeloproliferative Neoplasms (MPN)	yes	no	Date assessed:	YYYY/MM/DD		Date assessed:	YYYY/MM/DD	
PRE356	Pre-Transplant	Disease Classification	Myeloproliferative Neoplasms (MPN)	yes	no	Specify the molecular response	Complete response (CR: Eradication of pre-existing abnormality,Not assessed,Not applicable,None of the above: Does not meet the CR or PR criteria, Partial response (PR): ≥50% decrease in allele burden,Re-emergence of a pre-existing molecular abnormality		Specify the molecular response	Complete response (CR: Eradication of pre-existing abnormality,Not assessed,Not applicable,None of the above: Does not meet the CR or PR criteria, Partial response (PR): ≥50% decrease in allele burden,Re-emergence of a pre-existing molecular abnormality	
PRE357	Pre-Transplant	Disease Classification	Myeloproliferative Neoplasms (MPN)	yes	no	Date assessed:	YYYY/MM/DD		Date assessed:	YYYY/MM/DD	
PRE358	Pre-Transplant	Disease Classification	Other Leukemia (OL)	yes	no	Specify the other leukemia classification	Chronic lymphocytic leukemia (CLL), NOS,Chronic lymphocytic leukemia (CLL), B-cell / small lymphocytic lymphoma (SLL),Hairy cell leukemia,Hairy cell leukemia variant,Monoclonal B-cell lymphocytosis,Other leukemias,Other leukemias, NOS,PLL, B-cell,Prolymphocytic leukemia (PLL), NOS,PLL, T-cell		Specify the other leukemia classification	Chronic lymphocytic leukemia (CLL), NOS,Chronic lymphocytic leukemia (CLL), B-cell / small lymphocytic lymphoma (SLL),Hairy cell leukemia,Hairy cell leukemia variant,Monoclonal B-cell lymphocytosis,Other leukemias,Other leukemias, NOS,PLL, B-cell,Prolymphocytic leukemia (PLL), NOS,PLL, T-cell	
PRE359	Pre-Transplant	Disease Classification	Other Leukemia (OL)	yes	no	Specify other leukemia:	open text		Specify other leukemia:	open text	
PRE360	Pre-Transplant	Disease Classification	Other Leukemia (OL)	yes	no	Was any 17p abnormality detected?	No,yes		Was any 17p abnormality detected?	No,yes	
PRE361	Pre-Transplant	Disease Classification	Other Leukemia (OL)	yes	no	Did a histologic transformation to diffuse large B-cell lymphoma (Richter syndrome) occur at any time after CLL diagnosis?	No,yes		Did a histologic transformation to diffuse large B-cell lymphoma (Richter syndrome) occur at any time after CLL diagnosis?	No,yes	
PRE362	Pre-Transplant	Disease Classification	Other Leukemia (OL)	yes	no	What was the disease status? (Atypical CLL)	1st complete remission (no previous bone marrow or extramedullary relapse),1st relapse,2nd complete remission,2nd relapse,≥3rd complete remission,≥3rd relapse,No treatment,Primary induction failure		What was the disease status? (Atypical CLL)	1st complete remission (no previous bone marrow or extramedullary relapse),1st relapse,2nd complete remission,2nd relapse,≥3rd complete remission,≥3rd relapse,No treatment,Primary induction failure	
PRE363	Pre-Transplant	Disease Classification	Other Leukemia (OL)	yes	no	What was the disease status? (CLL, PLL, Hairy cell leukemia, Other leukemias)	Complete remission (CR),Not assessed,Untreated,Partial remission (PR),Progressive disease (Prog),Stable disease (SD)		What was the disease status? (CLL, PLL, Hairy cell leukemia, Other leukemias)	Complete remission (CR),Not assessed,Untreated,Partial remission (PR),Progressive disease (Prog),Stable disease (SD)	
PRE364	Pre-Transplant	Disease Classification	Other Leukemia (OL)	yes	no	Date assessed:	YYYY/MM/DD		Date assessed:	YYYY/MM/DD	
PRE365	Pre-Transplant	Disease Classification	Hodgkin and Non-Hodgkin Lymphoma	yes	no	Specify the lymphoma histology	Hodgkin Lymphoma Hodgkin lymphoma, not otherwise specified (150) Lymphocyte depleted (154) Lymphocyte-rich (152) Mixed cellularity (153) Nodular lymphocyte predominant Hodgkin lymphoma (155) Nodular sclerosis (152) Non-Hodgkin Lymphoma B-cell Neoplasms ALK+ large B-cell lymphoma (1833) B-cell lymphoma, unclassifiable, with features intermediate between DLBCL and classical Hodgkin lymphoma (149) Burkitt lymphoma (151) Burkitt-like lymphoma with t(1q aberration) (1834) Diffuse, large B-cell lymphoma: Activated B-cell type (non-GCB) (1821) Diffuse, large B-cell lymphoma: Germinal center B-cell type (1820) Diffuse large B-cell lymphoma (cell of origin unknown) (1507) DLBCL associated with chronic inflammation (1825) Dendroclonal type follicular lymphoma (1815) EBV+ DLCL, NOS (1826) EBV+ mucocutaneous ulcer (1824) Extranodal marginal zone B-cell lymphoma of mucosal associated lymphoid tissue type (MALT) (122) Follicular, mixed, small cleaved and large cell (Grade II follicle center lymphoma) (163) Follicular, predominantly large cell (Grade IIIA follicle center lymphoma) (162) Follicular, predominantly large cell (Grade IIIB follicle center lymphoma) (163) Follicular, predominantly large cell (Grade IIA vs IIB not specified) (1814) Follicular, predominantly small cleaved cell (Grade I follicle center lymphoma) (162) Follicular (grade unknown) (164) HIV+ DLCL, NOS (1826) High-grade B-cell lymphoma, with MYC and BCL2 and/or BCL6 rearrangements (1831) High-grade B-cell lymphoma, NOS (1830) Intraosseous large B-cell lymphoma (136) Large B-cell lymphoma with BCL6 rearrangement (1832) Lymphomatoid granulomatosis (1835) Mantle cell lymphoma (151) Nodal marginal zone B-cell lymphoma (a monocytoid B-cell) (123) Nodular, nodal marginal zone lymphoma (1814) open text	Hodgkin Lymphoma Hodgkin lymphoma, not otherwise specified (150) Lymphocyte depleted (154) Lymphocyte-rich (152) Mixed cellularity (153) Nodular lymphocyte predominant Hodgkin lymphoma (155) Nodular sclerosis (152) Non-Hodgkin Lymphoma B-cell Neoplasms ALK+ large B-cell lymphoma (1833) B-cell lymphoma, unclassifiable, with features intermediate between DLBCL and classical Hodgkin lymphoma (149) Burkitt lymphoma (151) Burkitt-like lymphoma with t(1q aberration) (1834) Diffuse, large B-cell lymphoma: Activated B-cell type (non-GCB) (1821) Diffuse, large B-cell lymphoma: Germinal center B-cell type (1820) Diffuse large B-cell lymphoma (cell of origin unknown) (1507) DLBCL associated with chronic inflammation (1825) Dendroclonal type follicular lymphoma (1815) EBV+ DLCL, NOS (1826) EBV+ mucocutaneous ulcer (1824) Extranodal marginal zone B-cell lymphoma of mucosal associated lymphoid tissue type (MALT) (122) Follicular, mixed, small cleaved and large cell (Grade II follicle center lymphoma) (163) Follicular, predominantly large cell (Grade IIIA follicle center lymphoma) (162) Follicular, predominantly large cell (Grade IIIB follicle center lymphoma) (163) Follicular, predominantly large cell (Grade IIA vs IIB not specified) (1814) Follicular, predominantly small cleaved cell (Grade I follicle center lymphoma) (162) Follicular (grade unknown) (164) HIV+ DLCL, NOS (1826) High-grade B-cell lymphoma, with MYC and BCL2 and/or BCL6 rearrangements (1831) High-grade B-cell lymphoma, NOS (1830) Intraosseous large B-cell lymphoma (136) Large B-cell lymphoma with BCL6 rearrangement (1832) Lymphomatoid granulomatosis (1835) Mantle cell lymphoma (151) Nodal marginal zone B-cell lymphoma (a monocytoid B-cell) (123) Nodular, nodal marginal zone lymphoma (1814) open text			
PRE366	Pre-Transplant	Disease Classification	Hodgkin and Non-Hodgkin Lymphoma	yes	no	Specify other lymphoma histology:	open text		Specify other lymphoma histology:	open text	

ID	Time Point	Information Collection Domain Sub-Type	Information Collection Domain Additional Sub-Domain	Response required if Additional Sub-Domain applies	Information Collection may be requested multiple times	Current Information Collection Data Element (if applicable)	Current Information Collection Data Element Response Option(s)	Information Collection update:	Proposed Information Collection Data Element (if applicable)	Proposed Information Collection Data Element Response Option(s)	Rationale for information Collection Update	
PRE367	Pre-Transplant	Disease Classification	Hodgkin and Non-Hodgkin lymphoma	yes	no	Is the lymphoma histology reported at transplant a transformation from CLL?	no, yes	Is the lymphoma histology reported at transplant a transformation from CLL?	no, yes (Also complete Chronic Lymphocytic Leukemia (CLL))			
PRE368	Pre-Transplant	Disease Classification	Hodgkin and Non-Hodgkin lymphoma	yes	no	Was any 17p abnormality detected?	no, yes	Was any 17p abnormality detected?	no, yes			
PRE369	Pre-Transplant	Disease Classification	Hodgkin and Non-Hodgkin lymphoma	yes	no	Is the lymphoma histology reported at transplant a transformation from a different lymphoma histology? (Not CLL)	No, Yes	Is the lymphoma histology reported at transplant a transformation from a different lymphoma histology? (Not CLL)	No, Yes			
PRE370	Pre-Transplant	Disease Classification	Hodgkin and Non-Hodgkin lymphoma	yes	no	Specify the original lymphoma histology (prior to transformation)	Aggressive NK cell leukemia, Anaplastic large-cell lymphoma (ALCL), ALK negative, Anaplastic large-cell lymphoma (ALCL), ALK positive, Angioimmunoblastic T-cell lymphoma, Adult T-cell lymphoma / leukemia (HTLV) associated, Breast implant associated anaplastic large-cell lymphoma, Burkitt-like lymphoma with 11q aberration, Chronic lymphoproliferative disorder of NK cell origin, Diffuse, large B-cell lymphoma (cell of origin unknown), B-cell lymphoma, unclassifiable, with features intermediate between DLBCL and classical Hodgkin lymphoma, DLBCL associated with chronic inflammation, EBV+ / DLBCL, NOS, Diffuse, large B-cell lymphoma - Germinal center B-cell type, HIV+ DLBCL, NOS, Diffuse, large B-cell lymphoma - Activated B-cell type (non-GBL), EBV+ mucocutaneous ulcer, Interferon-gamma T-cell lymphoma, Extranodal NK / T-cell lymphoma, nasal type, Diffuse large B-cell lymphoma, Interferon-gamma T-cell lymphoma, Pediatric-type follicular lymphoma, Pediatric-type follicular lymphoma, Follicular T-cell lymphoma, Follicular, predominantly large cell (Grade I/IIA follicle center lymphoma), Follicular, predominantly large cell (Grade I/IIA follicle center lymphoma), Follicular, predominantly large cell (Grade I/IIA vs IIII not specified), Follicular, predominantly small cleaved cell (Grade I follicle center lymphoma), Follicular, mixed, small cleaved and large cell (Grade I follicle center lymphoma), Hepatosplenic T-cell lymphoma, High grade B-cell lymphoma, with MYC and BCL2 and/or BCL6 rearrangements, High grade B-cell lymphoma, NOS, Hodgkin lymphoma, not otherwise specified, Infectious mononucleosis, PDL, Intraocular large B-cell lymphoma, Indolent T-cell lymphoproliferative disorder of the GI tract, ALK+ large B-cell lymphoma, Large B-cell lymphoma with HHV-8 rearrangement, Lymphocyte depleted, Lymphocyte-rich, Lymphomatoid granulomatosis, Extranodal marginal zone B-cell lymphoma of mucosal associated lymphoid tissue type (MALT), Mixed cellularity, Primary mediastinal (thymic) large B-cell lymphoma, Monomorphic epithelioid interstitial T-cell lymphoma, Mycosis fungoides, Mycitic cell lymphoma, Nodular lymphocyte predominant Hodgkin lymphoma, Nodal marginal zone B-cell lymphoma, Germans, monocytic B-cell, Nodal peripheral T-cell lymphoma with TTF1 phenotype, Nodular sclerosing, Other T-cell, NK-cell lymphoma, Other B-cell lymphoma, Primary cutaneous CD30+ aggressive epidermotropic cytotoxic T-cell lymphoma, Primary cutaneous CD30+ small / medium T-cell lymphoproliferative disorders (Primary cutaneous anaplastic large-cell lymphoma (C-ALCL), Lymphoid papulosis), Primary cutaneous CD4+ small / medium T-cell lymphoproliferative disorders, Primary cutaneous follicle center lymphoma, Primary cutaneous gamma delta T-cell lymphoma, Primary diffuse, large B-cell lymphoma of the CNS, Primary diffuse, large B-cell lymphoma of the CNS, Primary cutaneous DLBCL, IgG type, Pediatric nodal marginal zone lymphoma, Plasmablastic hyperplasia, PDL, Plasmablastic lymphoma, Primary effusion lymphoma, Peripheral T-cell lymphoma (PTCL), NOS, Pleural follicular hyperplasia, PDL, Classical Hodgkin lymphoma, PDL, Monomorphic, PDL B- and T-NK cell types, Polymorphic, PDL, Splenic B-cell lymphoma / leukemia, undetectable, Splenic diffuse red pulp small B-cell lymphoma, Splenic marginal zone B-cell lymphoma, Burkitt lymphoma, Subcutaneous panniculitis-like T-cell lymphoma, Systemic EBV+ T-cell lymphoma of childhood, Sezary syndrome, T-cell / histiocytic rich large B-cell lymphoma, T-cell large granular lymphocytic leukemia, Waldenström macroglobulinemia / Lymphoplasmacytic lymphoma	Specify the original lymphoma histology (prior to transformation)	Aggressive NK cell leukemia, Anaplastic large-cell lymphoma (ALCL), ALK negative, Anaplastic large-cell lymphoma (ALCL), ALK positive, Angioimmunoblastic T-cell lymphoma, Adult T-cell lymphoma / leukemia (HTLV) associated, Breast implant associated anaplastic large-cell lymphoma, Burkitt-like lymphoma with 11q aberration, Chronic lymphoproliferative disorder of NK cell origin, Diffuse, large B-cell lymphoma (cell of origin unknown), B-cell lymphoma, unclassifiable, with features intermediate between DLBCL and classical Hodgkin lymphoma, DLBCL associated with chronic inflammation, EBV+ / DLBCL, NOS, Diffuse, large B-cell lymphoma - Germinal center B-cell type, HIV+ DLBCL, NOS, Diffuse, large B-cell lymphoma - Activated B-cell type (non-GBL), EBV+ mucocutaneous ulcer, Interferon-gamma T-cell lymphoma, Extranodal NK / T-cell lymphoma, nasal type, Diffuse large B-cell lymphoma, Interferon-gamma T-cell lymphoma, Pediatric-type follicular lymphoma, Pediatric-type follicular lymphoma, Follicular T-cell lymphoma, Follicular, predominantly large cell (Grade I/IIA follicle center lymphoma), Follicular, predominantly large cell (Grade I/IIA follicle center lymphoma), Follicular, predominantly large cell (Grade I/IIA vs IIII not specified), Follicular, predominantly small cleaved cell (Grade I follicle center lymphoma), Follicular, mixed, small cleaved and large cell (Grade I follicle center lymphoma), Hepatosplenic T-cell lymphoma, High grade B-cell lymphoma, with MYC and BCL2 and/or BCL6 rearrangements, High grade B-cell lymphoma, NOS, Hodgkin lymphoma, not otherwise specified, Infectious mononucleosis, PDL, Intraocular large B-cell lymphoma, Indolent T-cell lymphoproliferative disorder of the GI tract, ALK+ large B-cell lymphoma, Large B-cell lymphoma with HHV-8 rearrangement, Lymphocyte depleted, Lymphocyte-rich, Lymphomatoid granulomatosis, Extranodal marginal zone B-cell lymphoma of mucosal associated lymphoid tissue type (MALT), Mixed cellularity, Primary mediastinal (thymic) large B-cell lymphoma, Monomorphic epithelioid interstitial T-cell lymphoma, Mycosis fungoides, Mycitic cell lymphoma, Nodular lymphocyte predominant Hodgkin lymphoma, Nodal marginal zone B-cell lymphoma, Germans, monocytic B-cell, Nodal peripheral T-cell lymphoma with TTF1 phenotype, Nodular sclerosing, Other T-cell, NK-cell lymphoma, Other B-cell lymphoma, Primary cutaneous CD30+ aggressive epidermotropic cytotoxic T-cell lymphoma, Primary cutaneous CD30+ small / medium T-cell lymphoproliferative disorders (Primary cutaneous anaplastic large-cell lymphoma (C-ALCL), Lymphoid papulosis), Primary cutaneous CD4+ small / medium T-cell lymphoproliferative disorders, Primary cutaneous follicle center lymphoma, Primary cutaneous gamma delta T-cell lymphoma, Primary diffuse, large B-cell lymphoma of the CNS, Primary diffuse, large B-cell lymphoma of the CNS, Primary cutaneous DLBCL, IgG type, Pediatric nodal marginal zone lymphoma, Plasmablastic hyperplasia, PDL, Plasmablastic lymphoma, Primary effusion lymphoma, Peripheral T-cell lymphoma (PTCL), NOS, Pleural follicular hyperplasia, PDL, Classical Hodgkin lymphoma, PDL, Monomorphic, PDL B- and T-NK cell types, Polymorphic, PDL, Splenic B-cell lymphoma / leukemia, undetectable, Splenic diffuse red pulp small B-cell lymphoma, Splenic marginal zone B-cell lymphoma, Burkitt lymphoma, Subcutaneous panniculitis-like T-cell lymphoma, Systemic EBV+ T-cell lymphoma of childhood, Sezary syndrome, T-cell / histiocytic rich large B-cell lymphoma, T-cell large granular lymphocytic leukemia, Waldenström macroglobulinemia / Lymphoplasmacytic lymphoma			
PRE371	Pre-Transplant	Disease Classification	Hodgkin and Non-Hodgkin lymphoma	yes	no	Specify other lymphoma histology:	open text	Specify other lymphoma histology:	open text			
PRE372	Pre-Transplant	Disease Classification	Hodgkin and Non-Hodgkin lymphoma	yes	no	Date of original lymphoma diagnosis: (report the date of diagnosis of original lymphoma subtype)	YYYY/MM/DD	Date of original lymphoma diagnosis: (report the date of diagnosis of original lymphoma subtype)	YYYY/MM/DD			
PRE373	Pre-Transplant	Disease Classification	Hodgkin and Non-Hodgkin lymphoma	yes	no	Was a PET (or PET/CT) scan performed? (at last evaluation prior to the start of the preparative regimen / infusion)	no, yes	Was a PET (or PET/CT) scan performed? (at last evaluation prior to the start of the preparative regimen / infusion)	no, yes			
PRE374	Pre-Transplant	Disease Classification	Hodgkin and Non-Hodgkin lymphoma	yes	no	Was the PET (or PET/CT) scan positive for lymphoma involvement at any disease site?	no, yes	Was the PET (or PET/CT) scan positive for lymphoma involvement at any disease site?	no, yes			
PRE375	Pre-Transplant	Disease Classification	Hodgkin and Non-Hodgkin lymphoma	yes	no	Date of PET scan	Known, Unknown	Date of PET scan	Known, Unknown			
PRE376	Pre-Transplant	Disease Classification	Hodgkin and Non-Hodgkin lymphoma	yes	no	Date of PET (or PET/CT) scan:	YYYY/MM/DD	Date of PET (or PET/CT) scan:	YYYY/MM/DD			
PRE377	Pre-Transplant	Disease Classification	Hodgkin and Non-Hodgkin lymphoma	yes	no	Deauville (five-point) score of the PET (or PET/CT) scan	Known, Unknown	Deauville (five-point) score of the PET (or PET/CT) scan	Known, Unknown			
PRE378	Pre-Transplant	Disease Classification	Hodgkin and Non-Hodgkin lymphoma	yes	no	Scale	1: no uptake or no residual uptake 2: slight uptake, but below blood pool (mediastinum) 3: uptake above mediastinal, but below or equal to uptake in the liver 4: uptake slightly to moderately higher than liver 5: markedly increased uptake or any new lesion	Scale	1: no uptake or no residual uptake 2: slight uptake, but below blood pool (mediastinum) 3: uptake above mediastinal, but below or equal to uptake in the liver 4: uptake slightly to moderately higher than liver 5: markedly increased uptake or any new lesion			
PRE379	Pre-Transplant	Disease Classification	Hodgkin and Non-Hodgkin lymphoma	yes	no	What was the disease status?	CR1 - 1st complete remission, no bone marrow or extramedullary relapse prior to transplant, CR2 - 2nd complete remission, CR3+ - 3rd or subsequent complete remission, PR, res - Primary induction failure - resistant, NEVER IN COMPLETE remission but with stable or progressive disease on treatment, PR, rem / PR1 - Primary induction failure - sensitive, NEVER IN COMPLETE remission but with partial remission on treatment, PR, rem - Primary induction failure - sensitively unknown, REL1, sen - 1st relapse - resistant, stable or progressive disease with treatment, REL1, sen - 1st relapse - sensitive, partial remission (if complete remission was achieved, classify as CR2), REL1, sen - 1st relapse - sensitively unknown, REL1, sen - 1st relapse - untreated, includes either bone marrow or extramedullary relapse, REL2, res - 2nd relapse - resistant, stable or progressive disease with treatment, REL2, sen - 2nd relapse - sensitive, partial remission (if complete remission achieved, classify as CR3), REL2, sen - 2nd relapse - sensitively unknown, REL2, sen - 2nd relapse - untreated, includes either bone marrow or extramedullary relapse, REL3, sen - 3rd or subsequent relapse - resistant, stable or progressive disease with treatment, REL3, sen - 3rd or subsequent relapse - sensitive, partial remission (if complete remission achieved, classify as CR3), REL3, sen - 3rd relapse or greater - sensitively unknown, REL3, sen - 3rd or subsequent relapse - untreated, includes either bone marrow or extramedullary relapse, Disease untreated	CR1 - 1st complete remission, no bone marrow or extramedullary relapse prior to transplant, CR2 - 2nd complete remission, CR3+ - 3rd or subsequent complete remission, PR, res - Primary induction failure - resistant, NEVER IN COMPLETE remission but with stable or progressive disease on treatment, PR, rem / PR1 - Primary induction failure - sensitive, NEVER IN COMPLETE remission but with partial remission on treatment, PR, rem - Primary induction failure - sensitively unknown, REL1, sen - 1st relapse - resistant, stable or progressive disease with treatment, REL1, sen - 1st relapse - sensitive, partial remission (if complete remission was achieved, classify as CR2), REL1, sen - 1st relapse - sensitively unknown, REL1, sen - 1st relapse - untreated, includes either bone marrow or extramedullary relapse, REL2, res - 2nd relapse - resistant, stable or progressive disease with treatment, REL2, sen - 2nd relapse - sensitive, partial remission (if complete remission achieved, classify as CR3), REL2, sen - 2nd relapse - sensitively unknown, REL2, sen - 2nd relapse - untreated, includes either bone marrow or extramedullary relapse, REL3, sen - 3rd or subsequent relapse - resistant, stable or progressive disease with treatment, REL3, sen - 3rd or subsequent relapse - sensitive, partial remission (if complete remission achieved, classify as CR3), REL3, sen - 3rd relapse or greater - sensitively unknown, REL3, sen - 3rd or subsequent relapse - untreated, includes either bone marrow or extramedullary relapse, Disease untreated	What was the disease status?	CR1 - 1st complete remission, no bone marrow or extramedullary relapse prior to transplant, CR2 - 2nd complete remission, CR3+ - 3rd or subsequent complete remission, PR, res - Primary induction failure - resistant, NEVER IN COMPLETE remission but with stable or progressive disease on treatment, PR, rem / PR1 - Primary induction failure - sensitive, NEVER IN COMPLETE remission but with partial remission on treatment, PR, rem - Primary induction failure - sensitively unknown, REL1, sen - 1st relapse - resistant, stable or progressive disease with treatment, REL1, sen - 1st relapse - sensitive, partial remission (if complete remission was achieved, classify as CR2), REL1, sen - 1st relapse - sensitively unknown, REL1, sen - 1st relapse - untreated, includes either bone marrow or extramedullary relapse, REL2, res - 2nd relapse - resistant, stable or progressive disease with treatment, REL2, sen - 2nd relapse - sensitive, partial remission (if complete remission achieved, classify as CR3), REL2, sen - 2nd relapse - sensitively unknown, REL2, sen - 2nd relapse - untreated, includes either bone marrow or extramedullary relapse, REL3, sen - 3rd or subsequent relapse - resistant, stable or progressive disease with treatment, REL3, sen - 3rd or subsequent relapse - sensitive, partial remission (if complete remission achieved, classify as CR3), REL3, sen - 3rd relapse or greater - sensitively unknown, REL3, sen - 3rd or subsequent relapse - untreated, includes either bone marrow or extramedullary relapse, Disease untreated		
PRE380	Pre-Transplant	Disease Classification	Hodgkin and Non-Hodgkin lymphoma	yes	no	Total number of lines of therapy received (between diagnosis and HCT / infusion)	1 line 2 lines, 3+ lines	Total number of lines of therapy received (between diagnosis and HCT / infusion)	1 line 2 lines, 3+ lines			
PRE381	Pre-Transplant	Disease Classification	Hodgkin and Non-Hodgkin lymphoma	yes	no	Date assessed:	YYYY/MM/DD	Date assessed:	YYYY/MM/DD			
PRE382	Pre-Transplant	Disease Classification	Multiple Myeloma / Plasma Cell Disorder (PCD)	yes	no	Specify the multiple myeloma/plasma cell disorder (PCD) classification	Amyloidosis, Monoclonal gammopathy of renal significance (MGRS), Multiple myeloma, Multiple myeloma light chain only, Multiple myeloma non-secretory, Osteosclerotic myeloma / POEMS syndrome, Other plasma cell disorder (PCD), Plasma cell leukemia (PCL), Smoldering myeloma, Solitary plasmacytoma	Specify the multiple myeloma/plasma cell disorder (PCD) classification	Amyloidosis, Monoclonal gammopathy of renal significance (MGRS), Multiple myeloma, Multiple myeloma light chain only, Multiple myeloma non-secretory, Osteosclerotic myeloma / POEMS syndrome, Other plasma cell disorder (PCD), Plasma cell leukemia (PCL), Smoldering myeloma, Solitary plasmacytoma			
PRE383	Pre-Transplant	Disease Classification	Multiple Myeloma / Plasma Cell Disorder (PCD)	yes	no	Specify other plasma cell disorder:	open text	Specify other plasma cell disorder:	open text			
PRE384	Pre-Transplant	Disease Classification	Multiple Myeloma / Plasma Cell Disorder (PCD)	yes	no	Specify heavy and/or light chain type (check all that apply)	IgA (heavy chain only), IgA kappa, IgA lambda, IgD (heavy chain only), IgD kappa, IgD lambda, IgE (heavy chain only), IgE kappa, IgE lambda, IgG (heavy chain only), IgG kappa, IgG lambda, IgM (heavy chain only), IgM kappa, IgM lambda, kappa (light chain only), Lambda (light chain only)	Specify heavy and/or light chain type (check all that apply)	IgA (heavy chain only), IgA kappa, IgA lambda, IgD (heavy chain only), IgD kappa, IgD lambda, IgE (heavy chain only), IgE kappa, IgE lambda, IgG (heavy chain only), IgG kappa, IgG lambda, IgM (heavy chain only), IgM kappa, IgM lambda, kappa (light chain only), Lambda (light chain only)			
PRE385	Pre-Transplant	Disease Classification	Multiple Myeloma / Plasma Cell Disorder (PCD)	yes	no	Specify Amyloidosis classification	AA amyloidosis, AA, amyloidosis, AL amyloidosis	Specify Amyloidosis classification	AA amyloidosis, AA, amyloidosis, AL amyloidosis			
PRE386	Pre-Transplant	Disease Classification	Multiple Myeloma / Plasma Cell Disorder (PCD)	yes	no	Select monoclonal gammopathy of renal significance (MGRS) classification	C3 glomerulopathy with monoclonal gammopathy, Crystal-storing histiocytosis, Immunotactoid glomerulopathy (ITGN) / glomerulonephritis with organized monoclonal microtubular immunoglobulin deposits (CGMD), Light chain tancer syndrome, Monoclonal immunoglobulin deposition disease (MIDD), Non-amyloid fibrillary glomerulonephritis, Proliferative glomerulonephritis with monoclonal immunoglobulin C deposits (PGCMID), Prerenal tubulopathy without crystal, Type 1 cryoglobulinemia, glomerulonephritis, Unknown	Select monoclonal gammopathy of renal significance (MGRS) classification	C3 glomerulopathy with monoclonal gammopathy, Crystal-storing histiocytosis, Immunotactoid glomerulopathy (ITGN) / glomerulonephritis with organized monoclonal microtubular immunoglobulin deposits (CGMD), Light chain tancer syndrome, Monoclonal immunoglobulin deposition disease (MIDD), Non-amyloid fibrillary glomerulonephritis, Proliferative glomerulonephritis with monoclonal immunoglobulin C deposits (PGCMID), Prerenal tubulopathy without crystal, Type 1 cryoglobulinemia, glomerulonephritis, Unknown			
PRE387	Pre-Transplant	Disease Classification	Multiple Myeloma / Plasma Cell Disorder (PCD)	yes	no	Select monoclonal immunoglobulin deposition disease (MIDD) subtype	Heavy chain deposition disease (HCDD), Light chain deposition disease (LCDD), Light and heavy chain deposition disease (LHCDD)	Select monoclonal immunoglobulin deposition disease (MIDD) subtype	Heavy chain deposition disease (HCDD), Light chain deposition disease (LCDD), Light and heavy chain deposition disease (LHCDD)			
PRE388	Pre-Transplant	Disease Classification	Multiple Myeloma / Plasma Cell Disorder (PCD)	yes	no	Was documentation submitted to the CIBMTR? (e.g. pathology report)	No, Yes	Was documentation submitted to the CIBMTR? (e.g. pathology report)	No, Yes			
PRE389	Pre-Transplant	Disease Classification	Multiple Myeloma / Plasma Cell Disorder (PCD)	yes	no	Solitary plasmacytoma was	Bone derived, Extramedullary	Solitary plasmacytoma was	Bone derived, Extramedullary			
PRE390	Pre-Transplant	Disease Classification	Multiple Myeloma / Plasma Cell Disorder (PCD)	yes	no	What was the Durie-Salmon staging? (at diagnosis)	Stage I (All of the following: Hgb > 9g/dL, serum calcium normal or < 0.25 mg/dL, bone x-ray normal bone structure (scale 0)), or solitary bone plasmacytoma only, low M-component production rates (lgG < 5g/dL, IgA < 3g/dL, urine light chain M-component on electrophoresis < 4g/24h) - Stage II (Fitting neither Stage I or Stage III) - Stage III (One of more of the following: Hgb < 8.5 g/dL, serum calcium > 12 mg/dL, advanced lytic bone lesions (scale 3); high M-component production rates (lgG > 7g/dL, IgA > 5g/dL, Bence Jones protein > 12g/24h) - Unknown	What was the Durie-Salmon staging? (at diagnosis)	Stage I (All of the following: Hgb > 9g/dL, serum calcium normal or < 0.25 mg/dL, bone x-ray normal bone structure (scale 0)), or solitary bone plasmacytoma only, low M-component production rates (lgG < 5g/dL, IgA < 3g/dL, urine light chain M-component on electrophoresis < 4g/24h) - Stage II (Fitting neither Stage I or Stage III) - Stage III (One of more of the following: Hgb < 8.5 g/dL, serum calcium > 12 mg/dL, advanced lytic bone lesions (scale 3); high M-component production rates (lgG > 7g/dL, IgA > 5g/dL, Bence Jones protein > 12g/24h) - Unknown			
PRE391	Pre-Transplant	Disease Classification	Multiple Myeloma / Plasma Cell Disorder (PCD)	yes	no	What was the Durie-Salmon sub classification? (at diagnosis)	A - relatively normal renal function (serum creatinine < 2.0 mg/dL), B - abnormal renal function (serum creatinine > 2.0 mg/dL)	What was the Durie-Salmon sub classification? (at diagnosis)	A - relatively normal renal function (serum creatinine < 2.0 mg/dL), B - abnormal renal function (serum creatinine > 2.0 mg/dL)			
PRE392	Pre-Transplant	Disease Classification	Multiple Myeloma / Plasma Cell Disorder (PCD)	yes	no	Did the recipient have a preceding or concurrent plasma cell disorder?	No, Yes	Did the recipient have a preceding or concurrent plasma cell disorder?	No, Yes			
PRE393	Pre-Transplant	Disease Classification	Preceding or Concurrent Plasma Cell Disorder	yes	yes	Specify preceding / concurrent disorder	Amyloidosis, Monoclonal gammopathy of renal significance, Monoclonal gammopathy of unknown significance, Multiple myeloma, Multiple myeloma - light chain only, Multiple myeloma - non-secretory, Osteosclerotic myeloma / POEMS syndrome, Other disease, Plasma cell leukemia, Smoldering myeloma, Solitary plasmacytoma	Specify preceding / concurrent disorder	Amyloidosis, Monoclonal gammopathy of renal significance, Monoclonal gammopathy of unknown significance, Multiple myeloma, Multiple myeloma - light chain only, Multiple myeloma - non-secretory, Osteosclerotic myeloma / POEMS syndrome, Other disease, Plasma cell leukemia, Smoldering myeloma, Solitary plasmacytoma			
PRE394	Pre-Transplant	Disease Classification	Preceding or Concurrent Plasma Cell Disorder	yes	yes	Specify other preceding/concurrent disorder:	open text	Specify other preceding/concurrent disorder:	open text			
PRE395	Pre-Transplant	Disease Classification	Preceding or Concurrent Plasma Cell Disorder	yes	yes	Date of diagnosis of preceding / concurrent disorder:	YYYY/MM/DD	Date of diagnosis of preceding / concurrent disorder:	YYYY/MM/DD			
PRE396	Pre-Transplant	Disease Classification	Multiple Myeloma / Plasma Cell Disorder (PCD)	yes	no	Serum beta2 - microglobulin	Known, Unknown	Serum beta2 - microglobulin	Known, Unknown			

Item ID	Time Point	Information Collection Domain Sub-Type	Information Collection Domain Additional Sub-Domain	Response required if additional sub-Domain applies	Information Collection may be requested multiple times	Current Information Collection Data Element (if applicable)	Current Information Collection Data Element Response Option(s)	Information Collection update:	Proposed Information Collection Data Element (if applicable)	Proposed Information Collection Data Element Response Option(s)	Rationale for information Collection Update
PRE484	Pre-Transplant	Disease Classification	Autoimmune Diseases	yes	no	Specify other autoimmune bowel disorder:	open text		Specify other autoimmune bowel disorder:	open text	
PRE485	Pre-Transplant	Disease Classification	Autoimmune Diseases	yes	no	Specify other autoimmune disease:	open text		Specify other autoimmune disease:	open text	
PRE486	Pre-Transplant	Disease Classification	Tolerance Induction Associated with Solid Organ Transplant	yes	no	Specify solid organ transplanted (check all that apply)	Kidney,Liver,Other organ,Pancreas		Specify solid organ transplanted (check all that apply)	Kidney,Liver,Other organ,Pancreas	
PRE487	Pre-Transplant	Disease Classification	Tolerance Induction Associated with Solid Organ Transplant	yes	no	Specify other organ:	open text		Specify other organ:	open text	
PRE488	Pre-Transplant	Disease Classification	Tolerance Induction Associated with Solid Organ Transplant	yes	no	Specify other disease:	open text		Specify other disease:	open text	
PRE489	Pre-Transplant	Disease Classification	Myelodysplastic syndrome (MDS)	yes	yes	WBC	Known,Unknown		WBC	Known,Unknown	
PRE490	Pre-Transplant	Disease Classification	Myelodysplastic syndrome (MDS)	yes	yes	WBC _____ x 10 ⁹ /L (x 10 ⁹ /mm ³) _____ x 10 ⁹ /L	WBC		WBC _____ x 10 ⁹ /L (x 10 ⁹ /mm ³) _____ x 10 ⁹ /L	WBC	
PRE491	Pre-Transplant	Disease Classification	Myelodysplastic syndrome (MDS)	yes	yes	Neutrophils	Known,Unknown		Neutrophils	Known,Unknown	
PRE492	Pre-Transplant	Disease Classification	Myelodysplastic syndrome (MDS)	yes	yes	Neutrophils _____%	Neutrophils _____%		Neutrophils _____%	Neutrophils _____%	
PRE493	Pre-Transplant	Disease Classification	Myelodysplastic syndrome (MDS)	yes	yes	Blasts in blood	Known,Unknown		Blasts in blood	Known,Unknown	
PRE494	Pre-Transplant	Disease Classification	Myelodysplastic syndrome (MDS)	yes	yes	Blasts in blood _____%	Blasts in blood _____%		Blasts in blood _____%	Blasts in blood _____%	
PRE495	Pre-Transplant	Disease Classification	Myelodysplastic syndrome (MDS)	yes	yes	Hemoglobin	Known,Unknown		Hemoglobin	Known,Unknown	
PRE496	Pre-Transplant	Disease Classification	Myelodysplastic syndrome (MDS)	yes	yes	At Diagnostic Hemoglobin _____ g/dL _____ g/L _____ mmol/L	At Diagnostic Hemoglobin _____ g/dL _____ g/L _____ mmol/L		At Diagnostic Hemoglobin _____ g/dL _____ g/L _____ mmol/L	At Diagnostic Hemoglobin _____ g/dL _____ g/L _____ mmol/L	
PRE497	Pre-Transplant	Disease Classification	Myelodysplastic syndrome (MDS)	yes	yes	Were RBCs transfused ≤ 30 days before date of test?	No,Yes		Were RBCs transfused ≤ 30 days before date of test?	No,Yes	
PRE498	Pre-Transplant	Disease Classification	Myelodysplastic syndrome (MDS)	yes	yes	Platelets	Known,Unknown		Platelets	Known,Unknown	
PRE499	Pre-Transplant	Disease Classification	Myelodysplastic syndrome (MDS)	yes	yes	Platelets _____ x 10 ⁹ /L (x 10 ⁹ /mm ³) _____ x 10 ⁹ /L	Platelets _____ x 10 ⁹ /L (x 10 ⁹ /mm ³) _____ x 10 ⁹ /L		Platelets _____ x 10 ⁹ /L (x 10 ⁹ /mm ³) _____ x 10 ⁹ /L	Platelets _____ x 10 ⁹ /L (x 10 ⁹ /mm ³) _____ x 10 ⁹ /L	
PRE500	Pre-Transplant	Disease Classification	Myelodysplastic syndrome (MDS)	yes	yes	Were platelets transfused ≤ 7 days before date of test?	No,Yes		Were platelets transfused ≤ 7 days before date of test?	No,Yes	
PRE501	Pre-Transplant	Disease Classification	Myelodysplastic syndrome (MDS)	yes	yes	WBC	Known,Unknown		WBC	Known,Unknown	
PRE502	Pre-Transplant	Disease Classification	Myelodysplastic syndrome (MDS)	yes	yes	Neutrophils	Known,Unknown		Neutrophils	Known,Unknown	
PRE503	Pre-Transplant	Disease Classification	Myelodysplastic syndrome (MDS)	yes	yes	Neutrophils _____%	Neutrophils _____%		Neutrophils _____%	Neutrophils _____%	
PRE504	Pre-Transplant	Disease Classification	Myelodysplastic syndrome (MDS)	yes	yes	Blasts in blood	Known,Unknown		Blasts in blood	Known,Unknown	
PRE505	Pre-Transplant	Disease Classification	Myelodysplastic syndrome (MDS)	yes	yes	Blasts in blood _____%	Blasts in blood _____%		Blasts in blood _____%	Blasts in blood _____%	
PRE506	Pre-Transplant	Disease Classification	Myelodysplastic syndrome (MDS)	yes	yes	Hemoglobin	Known,Unknown		Hemoglobin	Known,Unknown	
PRE507	Pre-Transplant	Disease Classification	Myelodysplastic syndrome (MDS)	yes	yes	Prior to infusion: Hemoglobin _____ g/dL _____ g/L _____ mmol/L	Prior to infusion: Hemoglobin _____ g/dL _____ g/L _____ mmol/L		Prior to infusion: Hemoglobin _____ g/dL _____ g/L _____ mmol/L	Prior to infusion: Hemoglobin _____ g/dL _____ g/L _____ mmol/L	
PRE508	Pre-Transplant	Disease Classification	Myelodysplastic syndrome (MDS)	yes	yes	Were RBCs transfused ≤ 30 days before date of test?	No,Yes		Were RBCs transfused ≤ 30 days before date of test?	No,Yes	
PRE509	Pre-Transplant	Disease Classification	Myelodysplastic syndrome (MDS)	yes	yes	Platelets	Known,Unknown		Platelets	Known,Unknown	
PRE510	Pre-Transplant	Disease Classification	Myelodysplastic syndrome (MDS)	yes	yes	Platelets _____ x 10 ⁹ /L (x 10 ⁹ /mm ³) _____ x 10 ⁹ /L	Platelets _____ x 10 ⁹ /L (x 10 ⁹ /mm ³) _____ x 10 ⁹ /L		Platelets _____ x 10 ⁹ /L (x 10 ⁹ /mm ³) _____ x 10 ⁹ /L	Platelets _____ x 10 ⁹ /L (x 10 ⁹ /mm ³) _____ x 10 ⁹ /L	
PRE511	Pre-Transplant	Disease Classification	Myelodysplastic syndrome (MDS)	yes	yes	Were platelets transfused ≤ 7 days before date of test?	No,Yes		Were platelets transfused ≤ 7 days before date of test?	No,Yes	
PRE512	Pre-Transplant	Disease Classification	Myeloproliferative Neoplasms (MPN)	yes	yes	WBC	Known,Unknown		WBC	Known,Unknown	
PRE513	Pre-Transplant	Disease Classification	Myeloproliferative Neoplasms (MPN)	yes	yes	WBC _____ x 10 ⁹ /L (x 10 ⁹ /mm ³) _____ x 10 ⁹ /L	WBC _____ x 10 ⁹ /L (x 10 ⁹ /mm ³) _____ x 10 ⁹ /L		WBC _____ x 10 ⁹ /L (x 10 ⁹ /mm ³) _____ x 10 ⁹ /L	WBC _____ x 10 ⁹ /L (x 10 ⁹ /mm ³) _____ x 10 ⁹ /L	
PRE514	Pre-Transplant	Disease Classification	Myeloproliferative Neoplasms (MPN)	yes	yes	Neutrophils	Known,Unknown		Neutrophils	Known,Unknown	
PRE515	Pre-Transplant	Disease Classification	Myeloproliferative Neoplasms (MPN)	yes	yes	Neutrophils _____%	Neutrophils _____%		Neutrophils _____%	Neutrophils _____%	
PRE516	Pre-Transplant	Disease Classification	Myeloproliferative Neoplasms (MPN)	yes	yes	Blasts in blood	Known,Unknown		Blasts in blood	Known,Unknown	
PRE517	Pre-Transplant	Disease Classification	Myeloproliferative Neoplasms (MPN)	yes	yes	Blasts in blood _____%	Blasts in blood _____%		Blasts in blood _____%	Blasts in blood _____%	
PRE518	Pre-Transplant	Disease Classification	Myeloproliferative Neoplasms (MPN)	yes	yes	Hemoglobin	Known,Unknown		Hemoglobin	Known,Unknown	
PRE519	Pre-Transplant	Disease Classification	Myeloproliferative Neoplasms (MPN)	yes	yes	Hemoglobin _____ g/dL _____ g/L _____ mmol/L	Hemoglobin _____ g/dL _____ g/L _____ mmol/L		Hemoglobin _____ g/dL _____ g/L _____ mmol/L	Hemoglobin _____ g/dL _____ g/L _____ mmol/L	
PRE520	Pre-Transplant	Disease Classification	Myeloproliferative Neoplasms (MPN)	yes	yes	Were RBCs transfused ≤ 30 days before date of test?	No,Yes		Were RBCs transfused ≤ 30 days before date of test?	No,Yes	
PRE521	Pre-Transplant	Disease Classification	Myeloproliferative Neoplasms (MPN)	yes	yes	Platelets	Known,Unknown		Platelets	Known,Unknown	
PRE522	Pre-Transplant	Disease Classification	Myeloproliferative Neoplasms (MPN)	yes	yes	Platelets _____ x 10 ⁹ /L (x 10 ⁹ /mm ³) _____ x 10 ⁹ /L	Platelets _____ x 10 ⁹ /L (x 10 ⁹ /mm ³) _____ x 10 ⁹ /L		Platelets _____ x 10 ⁹ /L (x 10 ⁹ /mm ³) _____ x 10 ⁹ /L	Platelets _____ x 10 ⁹ /L (x 10 ⁹ /mm ³) _____ x 10 ⁹ /L	
PRE523	Pre-Transplant	Disease Classification	Myeloproliferative Neoplasms (MPN)	yes	yes	Were platelets transfused ≤ 7 days before date of test?	No,Yes		Were platelets transfused ≤ 7 days before date of test?	No,Yes	
PRE524	Pre-Transplant	Disease Classification	Myeloproliferative Neoplasms (MPN)	yes	yes	WBC	Known,Unknown		WBC	Known,Unknown	
PRE525	Pre-Transplant	Disease Classification	Myeloproliferative Neoplasms (MPN)	yes	yes	WBC _____ x 10 ⁹ /L (x 10 ⁹ /mm ³) _____ x 10 ⁹ /L	WBC _____ x 10 ⁹ /L (x 10 ⁹ /mm ³) _____ x 10 ⁹ /L		WBC _____ x 10 ⁹ /L (x 10 ⁹ /mm ³) _____ x 10 ⁹ /L	WBC _____ x 10 ⁹ /L (x 10 ⁹ /mm ³) _____ x 10 ⁹ /L	
PRE526	Pre-Transplant	Disease Classification	Myeloproliferative Neoplasms (MPN)	yes	yes	Neutrophils	Known,Unknown		Neutrophils	Known,Unknown	

[illegible]

Item ID	Time Point	Information Collection Domain Sub-Type	Information Collection Domain Additional Sub-Domain	Response required if Additional Sub-Domain applies	Information Collection may be requested multiple times	Current Information Collection Data Element (if applicable)	Current Information Collection Data Element Response Option(s)	Information Collection update:	Proposed Information Collection Data Element (if applicable)	Proposed Information Collection Data Element Response Option(s)	Rationale for information Collection Update
PRE277	Pre-Transplant	Pre-Transplant Essential Data		no	no	Were there any co-existing diseases or organ impairment present according to the HCT comorbidity index (HCT-CI) [Source: Sorror, M.L. 2013]. How? Assess comorbidities before hematopoietic cell transplantation. Blood, 121(15), 2854-2863.]	No/Yes		Were there any co-existing diseases or organ impairment present according to the HCT comorbidity index (HCT-CI) [Source: Sorror, M.L. 2013]. How? Assess comorbidities before hematopoietic cell transplantation. Blood, 121(15), 2854-2863.]	No/Yes	
PRE278	Pre-Transplant	Pre-Transplant Essential Data	Comorbid Conditions	yes	no	Specify co-existing diseases or organ impairment (check all that apply) 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: 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 x upper limit of normal, or AST/ALT > 2.5 x 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 10th of the 99th percentile of higher for patients aged 18 years or younger Optic: ulcer - Any history of optic (optic or chiasmal) ulcer confirmed by endoscopic 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 < 60% or dyspnea on light 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 102 Rheumatologic: Any history of a rheumatologic disease (e.g., systemic lupus erythematosus, 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 103	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: 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 x upper limit of normal, or AST/ALT > 2.5 x 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 10th of the 99th percentile of higher for patients aged 18 years or younger Optic: ulcer - Any history of optic (optic or chiasmal) ulcer confirmed by endoscopic 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 < 60% or dyspnea on light 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 102 Rheumatologic: Any history of a rheumatologic disease (e.g., systemic lupus erythematosus, 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 103		Specify co-existing diseases or organ impairment (check all that apply) 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: 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 x upper limit of normal, or AST/ALT > 2.5 x 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 10th of the 99th percentile of higher for patients aged 18 years or younger Optic: ulcer - Any history of optic (optic or chiasmal) ulcer confirmed by endoscopic 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 < 60% or dyspnea on light 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 102 Rheumatologic: Any history of a rheumatologic disease (e.g., systemic lupus erythematosus, 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 103		
PRE279	Pre-Transplant	Pre-Transplant Essential Data	Comorbid Conditions	yes	no	Was the recipient on dialysis immediately prior to start of preparative regimen?	No/Unknown/Yes		Was the recipient on dialysis immediately prior to start of preparative regimen?	No/Unknown/Yes	
PRE280	Pre-Transplant	Pre-Transplant Essential Data	Comorbid Conditions	yes	no	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, esophagus) Genitourinary malignancy (e.g., kidney, bladder, ovary, testicle, genitalia, uterus, cervix, prostate) Acute myeloid leukemia Chronic lymphoblastic leukemia Chronic lymphoblastic leukemia Leukemia lung cancer Lymphoma (includes Hodgkin & non-Hodgkin lymphoma) MDS / MPM Melanoma Multiple myeloma / plasma cell disorder (PCD) Oropharyngeal cancer (e.g., tongue, buccal mucosa) Sarcoma Thyroid cancer Other skin malignancy (basal cell, squamous cell) Other hematologic malignancy Other solid tumor	Breast cancer Central nervous system (CNS) malignancy (e.g., glioblastoma, astrocytoma) Gastrointestinal malignancy (e.g., colon, rectum, stomach, pancreas, intestine, esophagus) Genitourinary malignancy (e.g., kidney, bladder, ovary, testicle, genitalia, uterus, cervix, prostate) Acute myeloid leukemia Chronic lymphoblastic leukemia Chronic lymphoblastic leukemia Leukemia lung cancer Lymphoma (includes Hodgkin & non-Hodgkin lymphoma) MDS / MPM Melanoma Multiple myeloma / plasma cell disorder (PCD) Oropharyngeal cancer (e.g., tongue, buccal mucosa) Sarcoma Thyroid cancer Other skin malignancy (basal cell, squamous cell) Other hematologic malignancy Other solid tumor		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, esophagus) Genitourinary malignancy (e.g., kidney, bladder, ovary, testicle, genitalia, uterus, cervix, prostate) Acute myeloid leukemia Chronic lymphoblastic leukemia Chronic lymphoblastic leukemia Leukemia lung cancer Lymphoma (includes Hodgkin & non-Hodgkin lymphoma) MDS / MPM Melanoma Multiple myeloma / plasma cell disorder (PCD) Oropharyngeal cancer (e.g., tongue, buccal mucosa) Sarcoma Thyroid cancer Other skin malignancy (basal cell, squamous cell) Other hematologic malignancy Other solid tumor		
PRE281	Pre-Transplant	Pre-Transplant Essential Data	Comorbid Conditions	yes	no	Specify other hematologic malignancy: (prior)	open text		Specify other hematologic malignancy: (prior)	open text	
PRE282	Pre-Transplant	Pre-Transplant Essential Data		no	no	Specify other solid tumor: (prior)	open text		Specify other solid tumor: (prior)	open text	
PRE283	Pre-Transplant	Pre-Transplant Essential Data		no	no	Date sample collected:	YYYY/MM/DD		Date sample collected:	YYYY/MM/DD	
PRE284	Pre-Transplant	Pre-Transplant Essential Data		no	no	Upper limit of normal for your institution:	open text		Upper limit of normal for your institution:	open text	
PRE285	Pre-Transplant	Pre-Transplant Essential Data		no	no	Date sample collected:	YYYY/MM/DD		Date sample collected:	YYYY/MM/DD	
PRE286	Pre-Transplant	Pre-Transplant Essential Data		no	no	Did the recipient have a prior solid organ transplant?	No/Yes		Did the recipient have a prior solid organ transplant?	No/Yes	
PRE287	Pre-Transplant	Pre-Transplant Essential Data	Prior Solid Organ Transplant	yes	yes	Specify organ	Bowel,Heart,Kidney(s),Liver,Lung,Other organ,Pancreas		Specify organ	Bowel,Heart,Kidney(s),Liver,Lung,Other organ,Pancreas	
PRE288	Pre-Transplant	Pre-Transplant Essential Data	Prior Solid Organ Transplant	yes	yes	Specify other organ:	open text		Specify other organ:	open text	
PRE289	Pre-Transplant	Pre-Transplant Essential Data	Prior Solid Organ Transplant	yes	yes	Year of prior solid organ transplant:	YYYY		Year of prior solid organ transplant:	YYYY	
PRE290	Pre-Transplant	Pre-Transplant Essential Data			yes	First Name (person completing form):	open text		First Name (person completing form):	open text	
PRE291	Pre-Transplant	Pre-Transplant Essential Data			yes	Last Name:	open text		Last Name:	open text	
PRE292	Pre-Transplant	Pre-Transplant Essential Data			yes	E-mail address:	open text		E-mail address:	open text	
PRE293	Pre-Transplant	Pre-Transplant Essential Data		no	no	Glomerular filtration rate (GFR) before start of preparative regimen (pediatric only)	Known/Unknown		Glomerular filtration rate (GFR) before start of preparative regimen (pediatric only)	Known/Unknown	
PRE295	Pre-Transplant	Pre-Transplant Essential Data		no	no	Serum ferritin (within 4 weeks prior to the start of the preparative regimen, use result closest to the start date)	Known/Unknown		Serum ferritin (within 4 weeks prior to the start of the preparative regimen, use result closest to the start date)	Known/Unknown	
PRE296	Pre-Transplant	Pre-Transplant Essential Data		no	no	Serum ferritin (within 4 weeks prior to the start of the preparative regimen, use result closest to the start date)	----- ng/mL (μg/L)		Serum ferritin (within 4 weeks prior to the start of the preparative regimen, use result closest to the start date)	----- ng/mL (μg/L)	
PRE297	Pre-Transplant	Pre-Transplant Essential Data		no	no	Serum albumin (within 4 weeks prior to the start of the preparative regimen, use result closest to the start date)	Known/Unknown		Serum albumin (within 4 weeks prior to the start of the preparative regimen, use result closest to the start date)	Known/Unknown	
PRE298	Pre-Transplant	Pre-Transplant Essential Data		no	no	Serum albumin (within 4 weeks prior to the start of the preparative regimen, use result closest to the start date)	----- g/dL ----- g/L		Serum albumin (within 4 weeks prior to the start of the preparative regimen, use result closest to the start date)	----- g/dL ----- g/L	
PRE299	Pre-Transplant	Pre-Transplant Essential Data		no	no	Platelets (within 4 weeks prior to the start of the preparative regimen, use result closest to the start date)	Known/Unknown		Platelets (within 4 weeks prior to the start of the preparative regimen, use result closest to the start date)	Known/Unknown	
PRE300	Pre-Transplant	Pre-Transplant Essential Data		no	no	Platelets (within 4 weeks prior to the start of the preparative regimen, use result closest to the start date)	----- x 10 ⁹ /L (x 10 ⁹ /mm ³) ----- x 10 ⁹ /L		Platelets (within 4 weeks prior to the start of the preparative regimen, use result closest to the start date)	----- x 10 ⁹ /L (x 10 ⁹ /mm ³) ----- x 10 ⁹ /L	
PRE301	Pre-Transplant	Pre-Transplant Essential Data		no	no	Were platelets transfused < 7 days before date of test?	No/Unknown/Yes		Were platelets transfused < 7 days before date of test?	No/Unknown/Yes	
PRE302	Pre-Transplant	Prior Exposure: Potential Study Eligibility		no	no	Specify if the recipient received any of the following (at any time prior to HCT / infusion) (check all that apply)	Blinatumomab(Blinxto),Gemtuzumab ozogamicin (Mylotarg),Inotuzumab ozogamicin (Besponsa), Mogamulizumab (Poteligeo) ,None,Other(spea		Specify if the recipient received any of the following (at any time prior to HCT / infusion) (check all that apply)	Blinatumomab(Blinxto),Gemtuzumab ozogamicin (Mylotarg),Inotuzumab ozogamicin (Besponsa), Mogamulizumab (Poteligeo) ,None,Other(spea	



Information Collection Domain: Transplant Procedure and Product Information

Item ID	Time Point	Information Collection Domain Sub-Type	Information Collection Domain Additional Sub Domain	Response required if Additional Sub Domain applies	Information Collection may be requested multiple times	Current Information Collection Data Element (if applicable)	Current Information Collection Data Element Response Option(s)	Information Collection update:	Proposed Information Collection Data Element (if applicable)	Proposed Information Collection Data Element Response Option(s)	Rationale for Information Collection Update
PRO001	Transplant Procedure and Product Information	Confirmation of HLA Typing	Non NMDP Allogeneic or syngeneic Donor / Recipient or Non NMDP Cord Blood Unit Information	yes	no	Registry donor ID:	open text		Registry donor ID:	open text	
PRO002	Transplant Procedure and Product Information	Confirmation of HLA Typing	Non NMDP Allogeneic or syngeneic Donor / Recipient or Non NMDP Cord Blood Unit Information	yes	no	Non-NMDP cord blood unit ID:	open text		Non-NMDP cord blood unit ID:	open text	
PRO003	Transplant Procedure and Product Information	Confirmation of HLA Typing	Non NMDP Allogeneic or syngeneic Donor / Recipient or Non NMDP Cord Blood Unit Information	yes	no	Global Registration Identifier for Donors (GRID)	open text		Global Registration Identifier for Donors (GRID)	open text	
PRO004	Transplant Procedure and Product Information	Confirmation of HLA Typing	Non NMDP Allogeneic or syngeneic Donor / Recipient or Non NMDP Cord Blood Unit Information	yes	no	ISBT DIN:	open text		ISBT DIN:	open text	

Item ID	Time Point	Collection Domain Sub-Type	Information Collection Domain Additional Sub Domain	Response required if Additional Sub Domain applies	Information Collection may be requested multiple times	Current Information Collection Data Element (if applicable)	Current Information Collection Data Element Response Option(s)	Information Collection update:	Proposed Information Collection Data Element (if applicable)	Proposed Information Collection Data Element Response Option(s)	Rationale for Information Collection Update
PRO005	Transplant Procedure and Product Information	Confirmation of HLA Typing	Non NMDP Allogeneic or syngeneic Donor / Recipient or Non NMDP Cord Blood Unit Information	yes	no	Registry or UCB Bank ID	(A) Austrian Bone Marrow Donors, (ACB) Austrian Cord Blood Registry, (ACCB) StemCyte, Inc, (AE) Emirates Bone Marrow Donor Registry, (AM) Armenian Bone Marrow Donor Registry Charitable Trust, (AOCB) University of Colorado Cord Blood Bank, (AR) Argentine CPH Donors Registry, (ARCB) BANCEL - Argentina Cord Blood Bank, (AUCB) Australian Cord Blood Registry, (AUS) Australian / New Zealand Bone Marrow Donor Registry, (B) Marrow Donor Program Belgium, (BCB) Belgium Cord Blood Registry, (BG) Bulgarian Bone Marrow Donor Registry, (BR) INCA/REDOMO, (BSCB) British Bone Marrow Registry - Cord Blood, (CB)		Registry or UCB Bank ID	(A) Austrian Bone Marrow Donors, (ACB) Austrian Cord Blood Registry, (ACCB) StemCyte, Inc, (AE) Emirates Bone Marrow Donor Registry, (AM) Armenian Bone Marrow Donor Registry Charitable Trust, (AOCB) University of Colorado Cord Blood Bank, (AR) Argentine CPH Donors Registry, (ARCB) BANCEL - Argentina Cord Blood Bank, (AUCB) Australian Cord Blood Registry, (AUS) Australian / New Zealand Bone Marrow Donor Registry, (B) Marrow Donor Program Belgium, (BCB) Belgium Cord Blood Registry, (BG) Bulgarian Bone Marrow Donor Registry, (BR) INCA/REDOMO, (BSCB) British Bone Marrow Registry - Cord Blood, (CB) Cord Blood Registry, (CH) Swiss BloodStem Cells - Adult Donors, (CHCB) Swiss Blood Stem Cells - Cord Blood, (CKCB) Celgene Cord Blood Bank, (CN) China Marrow Donor Program (CMDP), (CNCB) Shan Dong Cord Blood Bank, (CND) Canadian Blood Services Bone Marrow Donor Registry, (CS2) Czech National Marrow Donor Registry, (CSCR) Czech Stem Cells Registry, (CY) Cyprus Paraskevaudio Bone Marrow Donor Registry, (CY2) The Cyprus Bone Marrow Donor Registry, (D) ZKRD - Zentrales Knochenmarkspender - Register Deutschland Adult Donors, (DCB) ZKRD - Zentrales Knochenmarkspender - Register Deutschland Cord Blood, (DK) The Danish Bone Marrow Donor Registry, (DK2) Bone Marrow Donors Copenhagen (BMDC), (DUCB) German Branch of the European Cord Blood Bank, (E) REDMO, (ECB) Spanish Cord Blood Registry, (F) France Greffe de Moelle - Adult Donors, (FCB) France Greffe de Moelle - Cord Blood, (FI) Finnish Bone Marrow Donor Registry, (FICB) Finnish Cord Blood Registry, (GB) The Anthony Nolan Trust, (GB3) Welsh Bone Marrow Donor Registry, (GB4) British Bone Marrow Registry, (GR) Unrelated Hematopoietic Stem Cell Donor Registry Greece, (GRCB) Michigan Community Blood Centers Cord Blood Bank, (H) Hungarian Bone Marrow Donor Registry, (HEM) Hema-Quebec, (HK) Hong Kong Bone Marrow Donor Registry, (HR) Croatian Bone Marrow Donor	
PRO006	Transplant Procedure and Product Information	Confirmation of HLA Typing	Non NMDP Allogeneic or syngeneic Donor / Recipient or Non NMDP Cord Blood Unit Information	yes	no	Donor DOB:	YYYY/MM/DD		Donor DOB:	YYYY/MM/DD	
PRO007	Transplant Procedure and Product Information	Confirmation of HLA Typing	Non NMDP Allogeneic or syngeneic Donor / Recipient or Non NMDP Cord Blood Unit Information	yes	no	Donor age:	open text, check "Months" or check "Years"		Donor age:	open text, check "Months" or check "Years"	
PRO008	Transplant Procedure and Product Information	Confirmation of HLA Typing	Non NMDP Allogeneic or syngeneic Donor / Recipient or Non NMDP Cord Blood Unit Information	yes	no	Donor sex	female,male		Donor sex	female,male	

Item ID	Time Point	Center for International Blood Grouping Collection Domain Sub-Type	Information Collection Domain Additional Sub Domain	Response required if Additional Sub Domain applies	Information Collection may be requested multiple times	Current Information Collection Data Element (if applicable)	Current Information Collection Data Element Response Option(s)	Information Collection update:	Proposed Information Collection Data Element (if applicable)	Proposed Information Collection Data Element Response Option(s)	Rationale for Information Collection Update
PRO009	Transplant Procedure and Product Information	Confirmation of HLA Typing	Non NMDP Allogeneic or syngeneic Donor / Recipient or Non NMDP Cord Blood Unit Information	yes	no	Specify the person for whom this typing is being done	Donor,Recipient-final typing		Specify the person for whom this typing is being done	Donor,Recipient-final typing	
PRO010	Transplant Procedure and Product Information	Confirmation of HLA Typing	Non NMDP Allogeneic or syngeneic Donor / Recipient or Non NMDP Cord Blood Unit Information	yes	no	Was documentation submitted to the CIBMTR (e.g. lab report)	No,Yes		Was documentation submitted to the CIBMTR (e.g. lab report)	No,Yes	
PRO011	Transplant Procedure and Product Information	Confirmation of HLA Typing	Non NMDP Allogeneic or syngeneic Donor / Recipient or Non NMDP Cord Blood Unit Information	yes	no	Locus A	Known,Unknown		Locus A	Known,Unknown	
PRO012	Transplant Procedure and Product Information	Confirmation of HLA Typing	Non NMDP Allogeneic or syngeneic Donor / Recipient or Non NMDP Cord Blood Unit Information	yes	no	First A* allele designations:	open text		First A* allele designations:	open text	
PRO013	Transplant Procedure and Product Information	Confirmation of HLA Typing	Non NMDP Allogeneic or syngeneic Donor / Recipient or Non NMDP Cord Blood Unit Information	yes	no	Second A* allele designations:	open text		Second A* allele designations:	open text	
PRO014	Transplant Procedure and Product Information	Confirmation of HLA Typing	Non NMDP Allogeneic or syngeneic Donor / Recipient or Non NMDP Cord Blood Unit Information	yes	no	Locus B	Known,Unknown		Locus B	Known,Unknown	
PRO015	Transplant Procedure and Product Information	Confirmation of HLA Typing	Non NMDP Allogeneic or syngeneic Donor / Recipient or Non NMDP Cord Blood Unit Information	yes	no	First B* allele designations:	open text		First B* allele designations:	open text	

Item ID	Time Point	 Collection Domain Sub-Type	Information Collection Domain Additional Sub Domain	Response required if Additional Sub Domain applies	Information Collection may be requested multiple times	Current Information Collection Data Element (if applicable)	Current Information Collection Data Element Response Option(s)	Information Collection update:	Proposed Information Collection Data Element (if applicable)	Proposed Information Collection Data Element Response Option(s)	Rationale for Information Collection Update
PRO016	Transplant Procedure and Product Information	Confirmation of HLA Typing	Non NMDP Allogeneic or syngeneic Donor / Recipient or Non NMDP Cord Blood Unit Information	yes	no	Second B* allele designations:	open text		Second B* allele designations:	open text	
PRO017	Transplant Procedure and Product Information	Confirmation of HLA Typing	Non NMDP Allogeneic or syngeneic Donor / Recipient or Non NMDP Cord Blood Unit Information	yes	no	Locus C	Known,Unknown		Locus C	Known,Unknown	
PRO018	Transplant Procedure and Product Information	Confirmation of HLA Typing	Non NMDP Allogeneic or syngeneic Donor / Recipient or Non NMDP Cord Blood Unit Information	yes	no	First C* allele designations:	open text		First C* allele designations:	open text	
PRO019	Transplant Procedure and Product Information	Confirmation of HLA Typing	Non NMDP Allogeneic or syngeneic Donor / Recipient or Non NMDP Cord Blood Unit Information	yes	no	Second C* allele designations:	open text		Second C* allele designations:	open text	
PRO020	Transplant Procedure and Product Information	Confirmation of HLA Typing	Non NMDP Allogeneic or syngeneic Donor / Recipient or Non NMDP Cord Blood Unit Information	yes	no	Locus DRB1	Known,Unknown		Locus DRB1	Known,Unknown	
PRO021	Transplant Procedure and Product Information	Confirmation of HLA Typing	Non NMDP Allogeneic or syngeneic Donor / Recipient or Non NMDP Cord Blood Unit Information	yes	no	First DRB1* allele designations:	open text		First DRB1* allele designations:	open text	
PRO022	Transplant Procedure and Product Information	Confirmation of HLA Typing	Non NMDP Allogeneic or syngeneic Donor / Recipient or Non NMDP Cord Blood Unit Information	yes	no	Second DRB1* allele designations:	open text		Second DRB1* allele designations:	open text	

Item ID	Time Point	 Collection Domain Sub-Type	Information Collection Domain Additional Sub Domain	Response required if Additional Sub Domain applies	Information Collection may be requested multiple times	Current Information Collection Data Element (if applicable)	Current Information Collection Data Element Response Option(s)	Information Collection update:	Proposed Information Collection Data Element (if applicable)	Proposed Information Collection Data Element Response Option(s)	Rationale for Information Collection Update
PRO023	Transplant Procedure and Product Information	Confirmation of HLA Typing		no	no	Locus DRB3	Known,Unknown		Locus DRB3	Known,Unknown	
PRO024	Transplant Procedure and Product Information	Confirmation of HLA Typing		no	no	First DRB3* allele designations:	open text		First DRB3* allele designations:	open text	
PRO025	Transplant Procedure and Product Information	Confirmation of HLA Typing		no	no	Second DRB3* allele designations:	open text		Second DRB3* allele designations:	open text	
PRO026	Transplant Procedure and Product Information	Confirmation of HLA Typing		no	no	Locus DRB4	Known,Unknown		Locus DRB4	Known,Unknown	
PRO027	Transplant Procedure and Product Information	Confirmation of HLA Typing		no	no	First DRB4* allele designations:	open text		First DRB4* allele designations:	open text	
PRO028	Transplant Procedure and Product Information	Confirmation of HLA Typing		no	no	Second DRB4* allele designations:	open text		Second DRB4* allele designations:	open text	
PRO029	Transplant Procedure and Product Information	Confirmation of HLA Typing		no	no	Locus DRB5	Known,Unknown		Locus DRB5	Known,Unknown	
PRO030	Transplant Procedure and Product Information	Confirmation of HLA Typing		no	no	First DRB5* allele designations:	open text		First DRB5* allele designations:	open text	
PRO031	Transplant Procedure and Product Information	Confirmation of HLA Typing		no	no	Second DRB5* allele designations:	open text		Second DRB5* allele designations:	open text	
PRO032	Transplant Procedure and Product Information	Confirmation of HLA Typing		no	no	Locus DQB1	Known,Unknown		Locus DQB1	Known,Unknown	
PRO033	Transplant Procedure and Product Information	Confirmation of HLA Typing		no	no	First DQB1* allele designations:	open text		First DQB1* allele designations:	open text	
PRO034	Transplant Procedure and Product Information	Confirmation of HLA Typing		no	no	Second DQB1* allele designations:	open text		Second DQB1* allele designations:	open text	

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PRO035	Transplant Procedure and Product Information	Confirmation of HLA Typing		no	no	Locus DPB1	Known,Unknown		Locus DPB1	Known,Unknown	
PRO036	Transplant Procedure and Product Information	Confirmation of HLA Typing		no	no	First DPB1* allele designations:	open text		First DPB1* allele designations:	open text	
PRO037	Transplant Procedure and Product Information	Confirmation of HLA Typing		no	no	Second DPB1* allele designations:	open text		Second DPB1* allele designations:	open text	
PRO038	Transplant Procedure and Product Information	Confirmation of HLA Typing		no	no	Locus DQA1	Known,Unknown		Locus DQA1	Known,Unknown	
PRO039	Transplant Procedure and Product Information	Confirmation of HLA Typing		no	no	First DQA1* allele designations:	open text		First DQA1* allele designations:	open text	
PRO040	Transplant Procedure and Product Information	Confirmation of HLA Typing		no	no	Second DQA1* allele designations:	open text		Second DQA1* allele designations:	open text	
PRO041	Transplant Procedure and Product Information	Confirmation of HLA Typing		no	no	Locus DPA1	Known,Unknown		Locus DPA1	Known,Unknown	
PRO042	Transplant Procedure and Product Information	Confirmation of HLA Typing		no	no	First DPA1* allele designations:	open text		First DPA1* allele designations:	open text	
PRO043	Transplant Procedure and Product Information	Confirmation of HLA Typing		no	no	Second DPA1* allele designations:	open text		Second DPA1* allele designations:	open text	
PRO044	Transplant Procedure and Product Information	Confirmation of HLA Typing	Non NMDP Allogeneic or syngeneic Donor / Recipient or Non NMDP Cord Blood Unit Information	yes	no	A Antigens. Number of antigens provided	one,two		A Antigens. Number of antigens provided	one,two	

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PRO045	Transplant Procedure and Product Information	Confirmation of HLA Typing	Non NMDP Allogeneic or syngeneic Donor / Recipient or Non NMDP Cord Blood Unit Information	yes	no	Specificity – 1st antigen	A1,A10,A11,A19,A2,A203,A210,A23(9),A24(9),A2403,A25(10),A26(10),A28,A29(19),A3,A30(19),A31(19),A32(19),A33(19),A34(10),A36,A43,A66(10),A68(28),A69(28),A74(19),A80,A9,AX		Specificity – 1st antigen	A1,A10,A11,A19,A2,A203,A210,A23(9),A24(9),A2403,A25(10),A26(10),A28,A29(19),A3,A30(19),A31(19),A32(19),A33(19),A34(10),A36,A43,A66(10),A68(28),A69(28),A74(19),A80,A9,AX	
PRO046	Transplant Procedure and Product Information	Confirmation of HLA Typing	Non NMDP Allogeneic or syngeneic Donor / Recipient or Non NMDP Cord Blood Unit Information	yes	no	Specificity – 2nd antigen	A1,A10,A11,A19,A2,A203,A210,A23(9),A24(9),A2403,A25(10),A26(10),A28,A29(19),A3,A30(19),A31(19),A32(19),A33(19),A34(10),A36,A43,A66(10),A68(28),A69(28),A74(19),A80,A9,AX		Specificity – 2nd antigen	A1,A10,A11,A19,A2,A203,A210,A23(9),A24(9),A2403,A25(10),A26(10),A28,A29(19),A3,A30(19),A31(19),A32(19),A33(19),A34(10),A36,A43,A66(10),A68(28),A69(28),A74(19),A80,A9,AX	
PRO047	Transplant Procedure and Product Information	Confirmation of HLA Typing	Non NMDP Allogeneic or syngeneic Donor / Recipient or Non NMDP Cord Blood Unit Information	yes	no	B Antigens. Number of antigens provided	one,two		B Antigens. Number of antigens provided	one,two	
PRO048	Transplant Procedure and Product Information	Confirmation of HLA Typing	Non NMDP Allogeneic or syngeneic Donor / Recipient or Non NMDP Cord Blood Unit Information	yes	no	Specificity – 1st antigen	B12,B13,B14,B15,B16,B17,B18,B21,B22,B27,B2708,B35,B37,B38(16),B39(16),B3901,B3902,B40,B4005,B41,B42,B44(12),B45(12),B46,B47,B48,B49(21),B5,B50(21),B51(5),B5102,B5103,B52(5),B53,B54(22),B55(22),B56(22),B57(17),B58(17),B59,B60(40),B61(40),B62(15),B63(15),B64(14),B65(14),B67,B7,B70,B703,B71(70),B72(70),B73,B75(15),B76(15),B77(15),B78,B8,B81,B82,BX		Specificity – 1st antigen	B12,B13,B14,B15,B16,B17,B18,B21,B22,B27,B2708,B35,B37,B38(16),B39(16),B3901,B3902,B40,B4005,B41,B42,B44(12),B45(12),B46,B47,B48,B49(21),B5,B50(21),B51(5),B5102,B5103,B52(5),B53,B54(22),B55(22),B56(22),B57(17),B58(17),B59,B60(40),B61(40),B62(15),B63(15),B64(14),B65(14),B67,B7,B70,B703,B71(70),B72(70),B73,B75(15),B76(15),B77(15),B78,B8,B81,B82,BX	

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PRO049	Transplant Procedure and Product Information	Confirmation of HLA Typing	Non NMDP Allogeneic or syngeneic Donor / Recipient or Non NMDP Cord Blood Unit Information	yes	no	Specificity – 2nd antigen	B12,B13,B14,B15,B16,B17,B18,B21,B22,B27,B2708,B35,B37,B38(16),B39(16),B3901,B3902,B40,B4005,B41,B42,B44(12),B45(12),B46,B47,B48,B49(21),B5,B50(21),B51(5),B5102,B5103,B52(5),B53,B54(22),B55(22),B56(22),B57(17),B58(17),B59,B60(40),B61(40),B62(15),B63(15),B64(14),B65(14),B67,B7,B70,B703,B71(70),B72(70),B73,B75(15),B76(15),B77(15),B78,B8,B81,B82,BX		Specificity – 2nd antigen	B12,B13,B14,B15,B16,B17,B18,B21,B22,B27,B2708,B35,B37,B38(16),B39(16),B3901,B3902,B40,B4005,B41,B42,B44(12),B45(12),B46,B47,B48,B49(21),B5,B50(21),B51(5),B5102,B5103,B52(5),B53,B54(22),B55(22),B56(22),B57(17),B58(17),B59,B60(40),B61(40),B62(15),B63(15),B64(14),B65(14),B67,B7,B70,B703,B71(70),B72(70),B73,B75(15),B76(15),B77(15),B78,B8,B81,B82,BX	
PRO050	Transplant Procedure and Product Information	Confirmation of HLA Typing	Non NMDP Allogeneic or syngeneic Donor / Recipient or Non NMDP Cord Blood Unit Information	yes	no	C Antigens. Number of antigens provided	one,two		C Antigens. Number of antigens provided	one,two	
PRO051	Transplant Procedure and Product Information	Confirmation of HLA Typing	Non NMDP Allogeneic or syngeneic Donor / Recipient or Non NMDP Cord Blood Unit Information	yes	no	Specificity – 1st antigen	Cw1,Cw10(W3),Cw2,Cw3,Cw4,Cw5,Cw6,Cw7,Cw8,Cw9(W3),CX		Specificity – 1st antigen	Cw1,Cw10(W3),Cw2,Cw3,Cw4,Cw5,Cw6,Cw7,Cw8,Cw9(W3),CX	
PRO052	Transplant Procedure and Product Information	Confirmation of HLA Typing	Non NMDP Allogeneic or syngeneic Donor / Recipient or Non NMDP Cord Blood Unit Information	yes	no	Specificity – 2nd antigen	Cw1,Cw10(W3),Cw2,Cw3,Cw4,Cw5,Cw6,Cw7,Cw8,Cw9(W3),CX		Specificity – 2nd antigen	Cw1,Cw10(W3),Cw2,Cw3,Cw4,Cw5,Cw6,Cw7,Cw8,Cw9(W3),CX	
PRO053	Transplant Procedure and Product Information	Confirmation of HLA Typing	Non NMDP Allogeneic or syngeneic Donor / Recipient or Non NMDP Cord Blood Unit Information	yes	no	Specificity Bw4 present?	no,yes		Specificity Bw4 present?	no,yes	

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PRO054	Transplant Procedure and Product Information	Confirmation of HLA Typing	Non NMDP Allogeneic or syngeneic Donor / Recipient or Non NMDP Cord Blood Unit Information	yes	no	Specificity Bw6 present?	no,yes		Specificity Bw6 present?	no,yes	
PRO055	Transplant Procedure and Product Information	Confirmation of HLA Typing	Non NMDP Allogeneic or syngeneic Donor / Recipient or Non NMDP Cord Blood Unit Information	yes	no	DR Antigens. Number of antigens provided	one,two		DR Antigens. Number of antigens provided	one,two	
PRO056	Transplant Procedure and Product Information	Confirmation of HLA Typing	Non NMDP Allogeneic or syngeneic Donor / Recipient or Non NMDP Cord Blood Unit Information	yes	no	Specificity – 1st antigen	DR1,DR10,DR103,D R11(5),DR12(5),DR13(6),DR14(6),DR1403,DR1404,DR15(2),DR16(2),DR17(3),D R18(3),DR2,DR3,DR4,DR5,DR6,DR7,DR8,DR9,DRX		Specificity – 1st antigen	DR1,DR10,DR103,DR11(5),DR12(5),DR13(6),DR14(6),DR1403,DR1404,DR15(2),DR16(2),DR17(3),DR18(3),DR2,DR3,DR4,DR5,DR6,DR7,DR8,DR9,DRX	
PRO057	Transplant Procedure and Product Information	Confirmation of HLA Typing	Non NMDP Allogeneic or syngeneic Donor / Recipient or Non NMDP Cord Blood Unit Information	yes	no	Specificity – 2nd antigen	DR1,DR10,DR103,D R11(5),DR12(5),DR13(6),DR14(6),DR1403,DR1404,DR15(2),DR16(2),DR17(3),D R18(3),DR2,DR3,DR4,DR5,DR6,DR7,DR8,DR9,DRX		Specificity – 2nd antigen	DR1,DR10,DR103,DR11(5),DR12(5),DR13(6),DR14(6),DR1403,DR1404,DR15(2),DR16(2),DR17(3),DR18(3),DR2,DR3,DR4,DR5,DR6,DR7,DR8,DR9,DRX	
PRO058	Transplant Procedure and Product Information	Confirmation of HLA Typing	Non NMDP Allogeneic or syngeneic Donor / Recipient or Non NMDP Cord Blood Unit Information	yes	no	Specificity DR51 present?	no,yes		Specificity DR51 present?	no,yes	
PRO059	Transplant Procedure and Product Information	Confirmation of HLA Typing	Non NMDP Allogeneic or syngeneic Donor / Recipient or Non NMDP Cord Blood Unit Information	yes	no	Specificity DR52 present?	no,yes		Specificity DR52 present?	no,yes	

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PRO060	Transplant Procedure and Product Information	Confirmation of HLA Typing	Non NMDP Allogeneic or syngeneic Donor / Recipient or Non NMDP Cord Blood Unit Information	yes	no	Specificity DR53 present?	no,yes		Specificity DR53 present?	no,yes	
PRO061	Transplant Procedure and Product Information	Confirmation of HLA Typing	Non NMDP Allogeneic or syngeneic Donor / Recipient or Non NMDP Cord Blood Unit Information	yes	no	DQ Antigens. Number of antigens provided	one,two		DQ Antigens. Number of antigens provided	one,two	
PRO062	Transplant Procedure and Product Information	Confirmation of HLA Typing	Non NMDP Allogeneic or syngeneic Donor / Recipient or Non NMDP Cord Blood Unit Information	yes	no	Specificity – 1st antigen	DQ1,DQ2,DQ3,DQ4,DQ5(1),DQ6(1),DQ7(3),DQ8(3),DQ9(3),DQX		Specificity – 1st antigen	DQ1,DQ2,DQ3,DQ4,DQ5(1),DQ6(1),DQ7(3),DQ8(3),DQ9(3),DQX	
PRO063	Transplant Procedure and Product Information	Confirmation of HLA Typing	Non NMDP Allogeneic or syngeneic Donor / Recipient or Non NMDP Cord Blood Unit Information	yes	no	Specificity – 2nd antigen	DQ1,DQ2,DQ3,DQ4,DQ5(1),DQ6(1),DQ7(3),DQ8(3),DQ9(3),DQX		Specificity – 2nd antigen	DQ1,DQ2,DQ3,DQ4,DQ5(1),DQ6(1),DQ7(3),DQ8(3),DQ9(3),DQX	
PRO064	Transplant Procedure and Product Information	Confirmation of HLA Typing	Non NMDP Allogeneic or syngeneic Donor / Recipient or Non NMDP Cord Blood Unit Information	yes	no	DP Antigens. Number of antigens provided	one,two		DP Antigens. Number of antigens provided	one,two	
PRO065	Transplant Procedure and Product Information	Confirmation of HLA Typing	Non NMDP Allogeneic or syngeneic Donor / Recipient or Non NMDP Cord Blood Unit Information	yes	no	Specificity – 1st antigen	DPw1,DPw2,DPw3,DPw4,DPw5,DPw6,DPX		Specificity – 1st antigen	DPw1,DPw2,DPw3,DPw4,DPw5,DPw6,DPX	
PRO066	Transplant Procedure and Product Information	Confirmation of HLA Typing	Non NMDP Allogeneic or syngeneic Donor / Recipient or Non NMDP Cord Blood Unit Information	yes	no	Specificity – 2nd antigen	DPw1,DPw2,DPw3,DPw4,DPw5,DPw6,DPX		Specificity – 2nd antigen	DPw1,DPw2,DPw3,DPw4,DPw5,DPw6,DPX	

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PRO067	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Infusion		no	no	HCT type (check only one)	Allogeneic, related,Allogeneic, unrelated,Autologous		HCT type (check only one)	Allogeneic, related,Allogeneic, unrelated,Autologous	
PRO068	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Infusion	Non NMDP Allogeneic Donor / Infant Demographic Information	yes	no	Was the donor ever pregnant?	Not applicable (male donor or cord blood unit) ,No,Unknown, Yes		Was the donor ever pregnant?	Not applicable (male donor or cord blood unit) ,No,Unknown,Yes	
PRO069	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Infusion	Non NMDP Allogeneic Donor / Infant Demographic Information	yes	no	Number of pregnancies	Known,Unknown		Number of pregnancies	Known,Unknown	
PRO070	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Infusion	Non NMDP Allogeneic Donor / Infant Demographic Information	yes	no	Specify number of pregnancies:	open text		Specify number of pregnancies:	open text	
PRO071	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Infusion	Non NMDP Allogeneic Donor / Infant Demographic Information	yes	no	Ethnicity (donor)	Hispanic or Latino,Not applicable (not a resident of the USA),Not Hispanic or Latino,Unknown		Ethnicity (donor)	Hispanic or Latino,Not applicable (not a resident of the USA),Not Hispanic or Latino,Unknown	
PRO072	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Infusion	Non NMDP Allogeneic Donor / Infant Demographic Information	yes	no	Race (donor) (check all that apply)	American Indian or Alaska Native,Asian,Black or African American,Not reported,Native Hawaiian or Other Pacific Islander,Unknown, White		Race (donor) (check all that apply)	American Indian or Alaska Native,Asian,Black or African American,Not reported,Native Hawaiian or Other Pacific Islander,Unknown,White	

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PRO073	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Infusion	Non NMDP Allogeneic Donor / Infant Demographic Information	yes	no	Race detail (donor) (check all that apply)	African American,African (both parents born in Africa),South Asian,American Indian, South or Central America,Alaskan Native or Aleut,North American Indian,Black Caribbean,Caribbean Indian,Other White,Eastern European,Filipino (Pilipino),Guamania n,Hawaiian,Japane s,Korean,Mediterranean,Middle Eastern,North American,North Coast of Africa,Chinese,Nort hern European,Other Pacific Islander,Other Black,Samoan,Black South or Central American,Other Southeast Asian,Unknown,Vie tnamese,White Caribbean,Western European,White South or Central		Race detail (donor) (check all that apply)	African American,African (both parents born in Africa),South Asian,American Indian, South or Central America,Alaskan Native or Aleut,North American Indian,Black Caribbean,Caribbean Indian,Other White,Eastern European,Filipino (Pilipino),Guamanian,Hawaiian,Japanese,Korean,Mediterranean,Middle Eastern,North American,North Coast of Africa,Chinese,Northern European,Other Pacific Islander,Other Black,Samoan,Black South or Central American,Other Southeast Asian,Unknown,Vietnamese,White Caribbean,Western European,White South or Central American	
PRO074	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Infusion	Non NMDP Allogeneic Donor / Infant Demographic Information	yes	no	Was the donor a carrier for potentially transferable genetic diseases?	No,Yes		Was the donor a carrier for potentially transferable genetic diseases?	No,Yes	
PRO075	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Infusion	Non NMDP Allogeneic Donor / Infant Demographic Information	yes	no	Specify potentially transferable genetic disease (check all that apply)	Other hemoglobinopathy, Other disease,Sickle cell anemia,Thalassemia		Specify potentially transferable genetic disease (check all that apply)	Other hemoglobinopathy,Other disease,Sickle cell anemia,Thalassemia	
PRO076	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Infusion	Non NMDP Allogeneic Donor / Infant Demographic Information	yes	no	Specify other disease:	open text		Specify other disease:	open text	

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PRO077	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Infusion	Non NMDP Allogeneic Donor / Infant Demographic Information	yes	no	Was the donor / product tested for other transferable genetic or clonal abnormalities?	No,Unknown,Yes		Was the donor / product tested for other transferable genetic or clonal abnormalities?	No,Unknown,Yes	
PRO078	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Infusion	Non NMDP Allogeneic Donor / Infant Demographic Information	yes	no	Clonal hematopoiesis of indeterminate potential (CHIP)	No,Yes		Clonal hematopoiesis of indeterminate potential (CHIP)	No,Yes	
PRO079	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Infusion	Non NMDP Allogeneic Donor / Infant Demographic Information	yes	no	What was the method of testing used?	open text		What was the method of testing used?	open text	
PRO080	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Infusion	Non NMDP Allogeneic Donor / Infant Demographic Information	yes	no	Monoclonal B-cell lymphocytosis	No,Yes		Monoclonal B-cell lymphocytosis	No,Yes	
PRO081	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Infusion	Non NMDP Allogeneic Donor / Infant Demographic Information	yes	no	Other transferable genetic or clonal abnormality	No,Yes		Other transferable genetic or clonal abnormality	No,Yes	
PRO082	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Infusion	Non NMDP Allogeneic Donor / Infant Demographic Information	yes	no	Specify other transferable genetic or clonal abnormality:	open text		Specify other transferable genetic or clonal abnormality:	open text	
PRO083	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Infusion	Allo Related Donor / Infant Demographic Information	yes	no	Did this donor have a central line placed?	no,yes		Did this donor have a central line placed?	no,yes	
PRO084	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Infusion	Allo Related Donor / Infant Demographic Information	yes	no	Was the donor hospitalized (inpatient) during or after the collection?	no,yes		Was the donor hospitalized (inpatient) during or after the collection?	no,yes	
PRO085	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Infusion	Allo Related Donor / Infant Demographic Information	yes	no	Did the donor experience any life-threatening complications during or after the collection?	no,yes		Did the donor experience any life-threatening complications during or after the collection?	no,yes	
PRO086	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Infusion	Allo Related Donor / Infant Demographic Information	yes	no	Specify:	open text		Specify:	open text	

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PRO087	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Infusion	Allo Related Donor Infant Demographic Information	yes	no	Did the allogeneic donor give one or more autologous transfusion units?	No,Yes		Did the allogeneic donor give one or more autologous transfusion units?	No,Yes	
PRO088	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Infusion	Allo Related Donor / Infant Demographic Information	yes	no	Date of collection:	YYYY/MM/DD		Date of collection:	YYYY/MM/DD	
PRO089	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Infusion	Allo Related Donor Infant Demographic Information	yes	no	Number of units:	open text		Number of units:	open text	
PRO090	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Infusion	Allo Related Donor Infant Demographic Information	yes	no	Did the donor receive blood transfusions as a result of the collection?	Allogeneic transfusions,Autologous transfusions,No		Did the donor receive blood transfusions as a result of the collection?	Allogeneic transfusions,Autologous transfusions,No	
PRO091	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Infusion	Allo Related Donor / Infant Demographic Information	yes	no	Specify number of autologous units:	open text		Specify number of autologous units:	open text	
PRO092	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Infusion	Allo Related Donor Infant Demographic Information	yes	no	Specify number of allogeneic units:	open text		Specify number of allogeneic units:	open text	
PRO093	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Infusion	Allo Related Donor / Infant Demographic Information	yes	no	Did the donor die as a result of the collection?	no,yes		Did the donor die as a result of the collection?	no,yes	
PRO094	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Infusion	Allo Related Donor / Infant Demographic Information	yes	no	Specify cause of death:	open text		Specify cause of death:	open text	
PRO095	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Infusion		no	yes	First Name (person completing form):	open text		First Name (person completing form):	open text	
PRO096	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Infusion		no	yes	Last Name:	open text		Last Name:	open text	
PRO097	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Infusion		no	yes	E-mail address:	open text		E-mail address:	open text	

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PRO098	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Infusion		no	yes	Date:	YYYY/MM/DD		Date:	YYYY/MM/DD	
PRO099	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Infusion Product		no	no	Product type (check only one)	Bone marrow,Other product,PBSC,Singl e cord blood unit		Product type	Bone marrow,Other product,PBSC,Single cord blood unit	
PRO100	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Infusion Product		no	no	Specify:	open text		Specify:	open text	
PRO101	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Infusion Product		no	no	NMDP Product	No,Yes		NMDP Product	No,Yes	
PRO102	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Infusion Product		no	no	NMDP cord blood unit ID:	open text		NMDP cord blood unit ID:	open text	
PRO103	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Infusion Product		no	no	NMDP donor ID:	open text		NMDP donor ID:	open text	
PRO104	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Infusion Product		no	no	Registry donor ID:	open text		Registry donor ID:	open text	
PRO105	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Infusion Product		no	no	Non-NMDP cord blood unit ID:	open text		Non-NMDP cord blood unit ID:	open text	
PRO106	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Infusion Product		no	no	Global Registration Identifier for Donors (GRID)	open text		Global Registration Identifier for Donors (GRID)	open text	
PRO107	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Infusion Product		no	no	ISBT DIN:	open text		ISBT DIN:	open text	

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PRO108	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Infusion Product		no	no	Registry or UCB Bank ID	(A) Austrian Bone Marrow Donors, (ACB) Austrian Cord Blood Registry, (ACCB) StemCyte, Inc., (AE) Emirates Bone Marrow Donor Registry, (AM) Armenian Bone Marrow Donor Registry Charitable Trust, (AOCB) University of Colorado Cord Blood Bank, (AR) Argentine CPH Donors Registry, (ARCB) BANCEL - Argentina Cord Blood Bank, (AUCB) Australian Cord Blood Registry, (AUS) Australian / New Zealand Bone Marrow Donor Registry, (B) Marrow Donor Program Belgium, (BCB) Belgium Cord Blood Registry, (BG) Bulgarian Bone Marrow Donor Registry, (BR) INCA/REDOMO, (BSCB) British Bone Marrow Registry - Cord Blood, (CB)		Registry or UCB Bank ID	(A) Austrian Bone Marrow Donors, (ACB) Austrian Cord Blood Registry, (ACCB) StemCyte, Inc., (AE) Emirates Bone Marrow Donor Registry, (AM) Armenian Bone Marrow Donor Registry Charitable Trust, (AOCB) University of Colorado Cord Blood Bank, (AR) Argentine CPH Donors Registry, (ARCB) BANCEL - Argentina Cord Blood Bank, (AUCB) Australian Cord Blood Registry, (AUS) Australian / New Zealand Bone Marrow Donor Registry, (B) Marrow Donor Program Belgium, (BCB) Belgium Cord Blood Registry, (BG) Bulgarian Bone Marrow Donor Registry, (BR) INCA/REDOMO, (BSCB) British Bone Marrow Registry - Cord Blood, (CB) Cord Blood Registry, (CH) Swiss BloodStem Cells - Adult Donors, (CHCB) Swiss Blood Stem Cells - Cord Blood, (CKCB) Celgene Cord Blood Bank, (CN) China Marrow Donor Program (CMDP), (CNCB) Shan Dong Cord Blood Bank, (CND) Canadian Blood Services Bone Marrow Donor Registry, (CS2) Czech National Marrow Donor Registry, (CSCR) Czech Stem Cells Registry, (CY) Cyprus Paraskevaudio Bone Marrow Donor Registry, (CY2) The Cyprus Bone Marrow Donor Registry, (D) ZKRD - Zentrales Knochenmarkspender - Register Deutschland Adult Donors, (DCB) ZKRD - Zentrales Knochenmarkspender - Register Deutschland Cord Blood, (DK) The Danish Bone Marrow Donor Registry, (DK2) Bone Marrow Donors Copenhagen (BMDC), (DUCB) German Branch of the European Cord Blood Bank, (E) REDMO, (ECB) Spanish Cord Blood Registry, (F) France Greffe de Moelle - Adult Donors, (FCB) France Greffe de Moelle - Cord Blood, (FI) Finnish Bone Marrow Donor Registry, (FICB) Finnish Cord Blood Registry, (GB) The Anthony Nolan Trust, (GB3) Welsh Bone Marrow Donor Registry, (GB4) British Bone Marrow Registry, (GR) Unrelated Hematopoietic Stem Cell Donor Registry Greece, (GRCB) Michigan Community Blood Centers Cord Blood Bank, (H) Hungarian Bone Marrow Donor Registry, (HEM) Hema-Quebec, (HK) Hong Kong Bone Marrow Donor Registry, (HR) Croatian Bone Marrow Donor	
PRO109	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Infusion Product		no	no	Donor DOB:	YYYY/MM/DD		Donor DOB:	YYYY/MM/DD	
PRO110	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Infusion Product		no	no	Donor age:	open text, check "Months" or check "Years"		Donor age:	open text, check "Months" or check "Years"	
PRO111	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Infusion Product		no	no	Donor sex	open text, check "Months" or check "Years"		Donor sex	open text, check "Months" or check "Years"	

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PRO112	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Infusion Product	Allogeneic Donors	yes	no	Did the donor receive growth and mobilizing factors, prior to any stem cell harvest, to enhance the product collection for this HCT?	No,Yes		Did the donor receive growth and mobilizing factors, prior to any stem cell harvest, to enhance the product collection for this HCT?	No,Yes	
PRO113	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Infusion Product	Allogeneic Donors	yes	no	Specify growth and mobilizing factor(s) (check all that apply)	G-CSF (filgrastim, Neupogen),Pegylated G-CSF(pegfilgrastim, Neulasta) , Plerixafor (Mozobil) Other growth or mobilizing factor(s)		Specify growth and mobilizing factor(s) (check all that apply)	G-CSF (filgrastim, Neupogen),Pegylated G-CSF(pegfilgrastim, Neulasta) , Plerixafor (Mozobil) Other growth or mobilizing factor(s)	
PRO114	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Infusion Product	Allogeneic Donors	yes	no	Specify other growth or mobilizing factor(s):	open text		Specify other growth or mobilizing factor(s):	open text	
PRO115	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Infusion Product		no	no	Date of first collection for this mobilization:	YYYY/MM/DD		Date of first collection for this mobilization:	YYYY/MM/DD	
PRO116	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Infusion Product		no	no	Were anticoagulants or other agents added to the product between collection and infusion?	No,Yes		Were anticoagulants or other agents added to the product between collection and infusion?	No,Yes	
PRO117	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Infusion Product		no	no	Specify anticoagulant(s) or other agents (check all that apply)	Acid citrate dextrose (ACD, ACD-A), Citrate phosphate dextrose (CPD, CPD-A), Ethylenediaminetetraacetic acid (EDTA), Heparin, Other agent		Specify anticoagulant(s) or other agents (check all that apply)	Acid citrate dextrose (ACD, ACD-A), Citrate phosphate dextrose (CPD, CPD-A), Ethylenediaminetetraacetic acid (EDTA), Heparin, Other agent	

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PRO118	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Infusion Product		no	no	Specify other agent:	open text		Specify other agent:	open text	
PRO119	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Infusion Product		no	no	Was this product collected off-site and shipped to your facility?	no,yes		Was this product collected off-site and shipped to your facility?	no,yes	
PRO120	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Infusion Product		no	no	Date of receipt of product at your facility:	YYYY/MM/DD		Date of receipt of product at your facility:	YYYY/MM/DD	
PRO121	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Infusion Product		no	no	Time of receipt of product (24-hour clock):	Hour:Minute Check standard time or check daylight savings		Time of receipt of product (24-hour clock):	Hour:Minute Check standard time or check daylight savings	
PRO122	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Infusion Product		no	no	Specify the shipping environment of the product(s)	Room temperature, Cooled (refrigerator temperature, not frozen), Frozen (cryopreserved), Other shipping environment		Specify the shipping environment of the product(s)	Room temperature, Cooled (refrigerated gel pack, refrigerator temperature, not frozen), Frozen (cryopreserved), Other shipping environment	
PRO123	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Infusion Product		no	no	Specify other shipping environment:	open text		Specify other shipping environment:	open text	
PRO124	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Infusion Product		no	no	Was there any indication that the environment within the shipper was outside the expected temperature range for this product at any time during shipment?	no,yes		Was there any indication that the environment within the shipper was outside the expected temperature range for this product at any time during shipment?	no,yes	

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PRO125	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Infusion Product		no	no	Were the secondary containers (e.g., insulated shipping containers and unit cassette) intact when they arrived at your center?	no,yes		Were the secondary containers (e.g., insulated shipping containers and unit cassette) intact when they arrived at your center?	no,yes	
PRO126	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Infusion Product	Cord Blood Product Infusion	yes	no	Was the cord blood unit stored at your center prior to thawing?	no,yes		Was the cord blood unit stored at your center prior to thawing?	no,yes	
PRO127	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Infusion Product	Cord Blood Product Infusion	yes	no	Specify the storage method used for the cord blood unit	Electric freezer,Liquid nitrogen,Vapor phase		Specify the storage method used for the cord blood unit	Electric freezer,Liquid nitrogen,Vapor phase	
PRO128	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Infusion Product	Cord Blood Product Infusion	yes	no	Temperature during storage	< -150 OC , > -150 OC to < -135 OC , > -135 OC to < -80 OC, > -80 OC		Temperature during storage	< -150 OC , > -150 OC to < -135 OC , > -135 OC to < -80 OC, > -80 OC	
PRO129	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Infusion Product		no	no	Date storage started:	YYYY/MM/DD		Date storage started:	YYYY/MM/DD	
PRO130	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Infusion Product	Cord Blood Product Infusion	yes	no	Total nucleated cells: (Includes nucleated red and nucleated white cells)	_____ . ____ x 10 ^{____} (Includes nucleated red and nucleated white cells) (Cord blood units only)		Total nucleated cells: (Includes nucleated red and nucleated white cells)	_____ . ____ x 10 ^{____} (Includes nucleated red and nucleated white cells) (Cord blood units only)	
PRO131	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Infusion Product	Cord Blood Product Infusion	yes	no	CD34+ cells	Done,Not done		CD34+ cells	Done,Not done	
PRO132	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Infusion Product	Cord Blood Product Infusion	yes	no	Total number of CD34+ cells:	_____ . ____ x 10 ^{____} _____		Total number of CD34+ cells:	_____ . ____ x 10 ^{____} _____	

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PRO133	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Infusion Product		no	no	Was the product thawed from a cryopreserved state prior to infusion?	no,yes		Was the product thawed from a cryopreserved state prior to infusion?	no,yes	
PRO134	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Infusion Product		no	no	Was the entire product thawed?	no,yes		Was the entire product thawed?	no,yes	
PRO135	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Infusion Product	Cord Blood Product Infusion	yes	no	Specify the percent of the product that was thawed? (Cord Blood units only)	20%,80%,Other percent		Specify the percent of the product that was thawed? (Cord Blood units only)	20%,80%,Other percent	
PRO136	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Infusion Product	Cord Blood Product Infusion	yes	no	Specify other percent:	__ _%		Specify other percent:	__ _%	
PRO137	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Infusion Product		no	no	Date thawing process initiated:	YYYY/MM/DD		Date thawing process initiated:	YYYY/MM/DD	
PRO138	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Infusion Product		no	no	Time at initiation of thaw (24-hour clock):	Hour:Minute Check "standard time" or "check daylight savings time"		Time at initiation of thaw (24-hour clock):	Hour:Minute Check "standard time" or "check daylight savings time"	
PRO139	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Infusion Product		no	no	Time of thaw completion:	Hour:Minute Check "standard time" or "check daylight savings time"		Time of thaw completion:	Hour:Minute Check "standard time" or "check daylight savings time"	
PRO140	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Infusion Product		no	no	What method was used to thaw the product?	Electric warmer,Other method,Waterbath		What method was used to thaw the product?	Electric warmer,Other method,Waterbath	
PRO141	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Infusion Product		no	no	Specify other method:	open text		Specify other method:	open text	

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PRO142	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Infusion Product		no	no	Did any incidents or product complaints occur while preparing or thawing the product?	No,Yes		Did any incidents or product complaints occur while preparing or thawing the product?	No,Yes	
PRO143	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Infusion Product		no	no	Was the product processed prior to infusion?	No,Yes		Was the product processed prior to infusion?	No,Yes	
PRO144	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Infusion Product		no	no	Specify processing (check all that apply)	Buffy coat enriched (buffy coat preparation) ,Dilute d,Plasma reduced,RBC reduced,Washed		Specify processing (check all that apply)	Buffy coat enriched (buffy coat preparation) ,Diluted,Plasma reduced,RBC reduced,Washed	
PRO145	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Infusion Product		no	no	Was the product manipulated prior to infusion?	no,yes		Was the product manipulated prior to infusion?	no,yes	
PRO146	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Infusion Product		no	no	Specify manipulations performed (check all that apply)	CD34 enriched (CD34+ selection), Ex-vivo expansion, Ex-vivo T-cell depetion, Genetic manipulation (gene transfer / transuction), Other cell manipulation		Specify manipulations performed (check all that apply)	CD34 enriched (CD34+ selection), Ex-vivo expansion, Ex-vivo T-cell depetion, Genetic manipulation (gene transfer / transuction), Other cell manipulation	
PRO147	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Infusion Product		no	no	Specify antibodies used (check all that apply)	Alpha/beta antibody,Anti CD19,Anti CD3,Anti CD4,Anti CD45RA,Anti CD52,Anti CD8,Other antibody		Specify antibodies used (check all that apply)	Alpha/beta antibody,Anti CD19,Anti CD3,Anti CD4,Anti CD45RA,Anti CD52,Anti CD8,Other antibody	
PRO148	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Infusion Product		no	no	Specify other antibody:	open text		Specify other antibody:	open text	

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PRO149	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Infusion Product		no	no	Specify T-cell depletion method	Antibody affinity column,Immunomagnetic beads,Other Method		Specify T-cell depletion method	Antibody affinity column,Immunomagnetic beads,Other Method	
PRO150	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Infusion Product		no	no	Specify other method:	open text		Specify other method:	open text	
PRO151	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Infusion Product		no	no	Specify other cell manipulation:	open text		Specify other cell manipulation:	open text	
PRO152	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Infusion Product		no	yes	Specify the timepoint in the product preparation phase that the product was analyzed	Product arrival (cord blood only) , At infusion (final quantity infused)		Specify the timepoint in the product preparation phase that the product was analyzed	Product arrival (cord blood only) , At infusion (final quantity infused)	
PRO153	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Infusion Product		no	yes	Date of product analysis:	YYYY/MM/DD		Date of product analysis:	YYYY/MM/DD	
PRO154	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Infusion Product		no	yes	Total volume of product plus additives:	_____ _ ml		Total volume of product plus additives:	_____ _ ml	
PRO155	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Infusion Product		no	yes	Total nucleated cells (TNC)	Done,Not done		Total nucleated cells (TNC)	Done,Not done	
PRO156	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Infusion Product		no	yes	Total nucleated cells:	_____ . _____ x 10 _____		Total nucleated cells:	_____ . _____ x 10 _____	
PRO157	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Infusion Product		no	yes	Viability of TNC	Done,Not done,Unknown		Viability of TNC	Done,Not done,Unknown	

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PRO158	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Infusion Product		no	yes	Viability of TNC:	___ %		Viability of TNC:	___ %	
PRO159	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Infusion Product		no	yes	Method of testing TNC viability	Flow cytometry based, Other method, Trypan blue		Method of testing TNC viability	Flow cytometry based (7AAD, AOPI, AOEB), Other method, Trypan blue	
PRO160	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Infusion Product		no	yes	Specify other method:	open text		Specify other method:	open text	
PRO161	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Infusion Product		no	yes	Nucleated white blood cells	Done, Not done		Nucleated white blood cells	Done, Not done	
PRO162	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Infusion Product		no	yes	Total number of nucleated white blood cells:	___ . ___ x 10 ___		Total number of nucleated white blood cells:	___ . ___ x 10 ___	
PRO163	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Infusion Product		no	yes	Mononuclear cells	Done, Not done		Mononuclear cells	Done, Not done	
PRO164	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Infusion Product		no	yes	Total number of mononuclear cells:	___ . ___ x 10 ___		Total number of mononuclear cells:	___ . ___ x 10 ___	
PRO165	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Infusion Product		no	yes	Nucleated red blood cells	Done, Not done		Nucleated red blood cells	Done, Not done	
PRO166	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Infusion Product		no	yes	Total number of nucleated red blood cells:	___ . ___ x 10 ___		Total number of nucleated red blood cells:	___ . ___ x 10 ___	
PRO167	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Infusion Product		no	yes	CD34+ cells	Done, Not done		CD34+ cells	Done, Not done	

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PRO168	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Infusion Product		no	yes	Total number of CD34+ cells:	----- · ----- x 10 -----		Total number of CD34+ cells:	----- · ----- x 10 -----	
PRO169	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Infusion Product		no	yes	Viability of CD34+ cells	Done,Not done,Unknown		Viability of CD34+ cells	Done,Not done,Unknown	
PRO170	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Infusion Product		no	yes	Viability of CD34+ cells:	-----%		Viability of CD34+ cells:	-----%	
PRO171	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Infusion Product		no	yes	Method of testing CD34+ cell viability	Flow cytometry based,Other method,Trypan blue		Method of testing CD34+ cell viability	Flow cytometry based (7AAD, AOPI, AOEB), Other method,Trypan blue	
PRO172	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Infusion Product		no	yes	Specify other method:	open text		Specify other method:	open text	
PRO173	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Infusion Product		no	yes	CD3+ cells	Done,Not done		CD3+ cells	Done,Not done	
PRO174	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Infusion Product		no	yes	Viability of CD3+ cells	Done,Not done,Unknown		Viability of CD3+ cells	Done,Not done,Unknown	
PRO175	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Infusion Product		no	yes	Total number of CD3+ cells:	----- · ----- x 10 -----		Total number of CD3+ cells:	----- · ----- x 10 -----	
PRO176	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Infusion Product		no	yes	Viability of CD3+ cells:	-----%		Viability of CD3+ cells:	-----%	
PRO177	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Infusion Product		no	yes	Method of testing CD3+ cell viability	Flow cytometry based,Other method,Trypan blue		Method of testing CD3+ cell viability	Flow cytometry based (7AAD, AOPI, AOEB), Other method,Trypan blue	

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PRO178	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Infusion Product		no	yes	Specify other method:	open text		Specify other method:	open text	
PRO179	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Infusion Product		no	yes	CD3+CD4+ cells	Done,Not done		CD3+CD4+ cells	Done,Not done	
PRO180	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Infusion Product		no	yes	Total number of CD3+CD4+ cells: _____ x 10 _____			Total number of CD3+CD4+ cells: _____ x 10 _____		
PRO181	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Infusion Product		no	yes	Viability of CD3+CD4+ cells	Done,Not done,Unknown		Viability of CD3+CD4+ cells	Done,Not done,Unknown	
PRO182	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Infusion Product		no	yes	Viability of CD3+CD4+ cells: ____%			Viability of CD3+CD4+ cells: ____%		
PRO183	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Infusion Product		no	yes	Method of testing CD3+CD4+ cell viability	Flow cytometry based,Other method,Trypan blue		Method of testing CD3+CD4+ cell viability	Flow cytometry based (7AAD, AOPI, AOEB), Other method,Trypan blue	
PRO184	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Infusion Product		no	yes	Specify other method:	open text		Specify other method:	open text	
PRO185	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Infusion Product		no	yes	CD3+CD8+ cells	Done,Not done		CD3+CD8+ cells	Done,Not done	
PRO186	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Infusion Product		no	yes	Total number of CD3+CD8+ cells: _____ * ____ x 10 ____			Total number of CD3+CD8+ cells: _____ * ____ x 10 ____		
PRO187	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Infusion Product		no	yes	Viability of CD3+CD8+ cells	Done,Not done,Unknown		Viability of CD3+CD8+ cells	Done,Not done,Unknown	

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PRO188	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Infusion Product		no	yes	Viability of CD3+CD8+ cells:	___ %		Viability of CD3+CD8+ cells:	___ %	
PRO189	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Infusion Product		no	yes	Method of testing CD3+CD8+ cell viability	Flow cytometry based, Other method, Trypan blue		Method of testing CD3+CD8+ cell viability	Flow cytometry based (7AAD, AOPI, AOEB), Other method, Trypan blue	
PRO190	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Infusion Product		no	yes	Specify other method:	open text		Specify other method:	open text	
PRO191	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Product Infusion	Cord Blood Product Infusion	yes	yes	Were the colony-forming units (CFU) assessed after thawing? (cord blood units only)	no, yes		Were the colony-forming units (CFU) assessed after thawing? (cord blood units only)	no, yes	
PRO192	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Product Infusion	Cord Blood Product Infusion	yes	yes	Was there growth?	no, yes		Was there growth?	no, yes	
PRO193	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Product Infusion	Cord Blood Product Infusion	yes	yes	Total CFU-GM	Done, Not done		Indicate which Assessments were Carried out (Check all that apply)	Total CFU-GM, Total CFU-GEMM, Total BFU-E	
PRO194	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Product Infusion	Cord Blood Product Infusion	yes	yes	Total CFU-GM:	_____x10__ —		Total CFU-GM:	_____x10__ —	
PRO195	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Product Infusion	Cord Blood Product Infusion	yes	yes	Total CFU-GEMM:	_____x10__ —		Total CFU-GEMM:	_____x10__ —	
PRO196	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Product Infusion	Cord Blood Product Infusion	yes	yes	Total BFU-E:	_____x10__ —		Total BFU-E:	_____x10__ —	

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PRO197	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Product Infusion		no	yes	Were any positive cultures (for bacterial or fungal infections) obtained from the product at the transplant center? (complete for all cell products)	No,Pending,Unknown,Yes		Were any positive cultures (for bacterial or fungal infections) obtained from the product at the transplant center? (complete for all cell products)	No,Pending,Unknown,Yes	
PRO198	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Product Infusion	Product Analysis	yes	yes	Specify Organism Code(s):	Bacterial Infections: 121 inetobacter (all species), 125 Bordetella pertussis (whooping cough), 128 Campylobacter (all species), 129 Capnocytophaga (all species), 171 Chlamydia (pneumoniae), 130 Citrobacter (freundii, other species), 131 Clostridium (all species except difficile), 132 Clostridium difficile, 173 Corynebacterium jeikeium, 134 Enterobacter (all species), 135 Enterococcus (all species), 177 Enterococcus, vancomycin resistant (VRE), 136 Escherichia (also E. coli), 139 Fusobacterium (all species), 187 Haemophilus influenzae, 188 Haemophilus non-influenzae, 146 Klebsiella (all		Specify Organism Code(s):	Bacterial Infections: 121 inetobacter (all species), 125 Bordetella pertussis (whooping cough), 128 Campylobacter (all species), 129 Capnocytophaga (all species), 171 Chlamydia (pneumoniae), 130 Citrobacter (freundii, other species), 131 Clostridium (all species except difficile), 132 Clostridium difficile, 173 Corynebacterium jeikeium, 134 Enterobacter (all species), 135 Enterococcus (all species), 177 Enterococcus, vancomycin resistant (VRE), 136 Escherichia (also E. coli), 139 Fusobacterium (all species), 187 Haemophilus influenzae, 188 Haemophilus non-influenzae, 146 Klebsiella (all	

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PRO199	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Product Infusion	Product Analysis	yes	yes	Specify Organism Code(s):	Bacterial Infections: 121 inetobacter (all species), 125 Bordetella pertussis (whooping cough), 128 Campylobacter (all species), 129 Capnocytophaga (all species), 171 Chlamydia (pneumoniae), 130 Citrobacter (freundii, other species), 131 Clostridium (all species except difficile), 132 Clostridium difficile, 173 Corynebacterium jeikeium, 134 Enterobacter (all species), 135 Enterococcus (all species), 177 Enterococcus, vancomycin resistant (VRE), 136 Escherichia (also E. coli), 139 Fusobacterium (all species), 187 Haemophilus influenzae, 188 Haemophilus non-influenzae, 146 Klebsiella (all		Specify Organism Code(s):	Bacterial Infections: 121 inetobacter (all species), 125 Bordetella pertussis (whooping cough), 128 Campylobacter (all species), 129 Capnocytophaga (all species), 171 Chlamydia (pneumoniae), 130 Citrobacter (freundii, other species), 131 Clostridium (all species except difficile), 132 Clostridium difficile, 173 Corynebacterium jeikeium, 134 Enterobacter (all species), 135 Enterococcus (all species), 177 Enterococcus, vancomycin resistant (VRE), 136 Escherichia (also E. coli), 139 Fusobacterium (all species), 187 Haemophilus influenzae, 188 Haemophilus non-influenzae, 146 Klebsiella (all species), 147 Lactobacillus (bulgaricus, acidophilus, other species), 189 Legionella pneumophila, 190 Legionella non-pneumophila, 103 Leptospira (all species), 148 Leptotrichia buccalis, 149 Leuconostoc (all species), 104 Listeria monocytogenes, 151 Micrococcus, NOS, 118 Mycobacterium abscessus, 112 Mycobacterium avium - intracellulare (MAC, MAI), 108 Mycobacterium cheloneae, 109 Mycobacterium fortuitum, 114 Mycobacterium haemophilum, 115 Mycobacterium kansasii, 116 Mycobacterium marinum, 117 Mycobacterium mucogenicum, 110 Mycobacterium tuberculosis (tuberculosis, Koch bacillus), 105 Mycoplasma (all species), 183 Neisseria gonorrhoeae, 184 Neisseria meningitidis, 106 Nocardia (all species), 153 Pasteurella multocida, 155 Proteus (all species), 157 Pseudomonas or Burkholderia cepacia, 185 Pseudomonas aeruginosa, 186 Pseudomonas non-aeruginosa, 159 Rhodococcus (all species), 107 Rickettsia (all species), 160 Salmonella (all species), 161 Serratia marcescens, 162 Shigella (all species), 180 Staphylococcus aureus (Methicillin Resistant), 179 Staphylococcus aureus (Methicillin Sensitive), 158 Stenotrophomonas maltophilia, 166 Stomatococcus mucilaginosus, 181 Streptococcus, alpha-hemolytic, 182 Streptococcus, Group B, 178 Streptococcus pneumoniae, 168 Treponema (syphilis), 169 Vibrio (all species) Fungal	

Item ID	Time Point	 Collection Domain Sub-Type	Information Collection Domain Additional Sub Domain	Response required if Additional Sub Domain applies	Information Collection may be requested multiple times	Current Information Collection Data Element (if applicable)	Current Information Collection Data Element Response Option(s)	Information Collection update:	Proposed Information Collection Data Element (if applicable)	Proposed Information Collection Data Element Response Option(s)	Rationale for Information Collection Update
PRO200	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Product Infusion	Product Analysis	yes	yes	Specify Organism Code(s):	Bacterial Infections: 121 inetobacter (all species), 125 Bordetella pertussis (whooping cough), 128 Campylobacter (all species), 129 Capnocytophaga (all species), 171 Chlamydia (pneumoniae), 130 Citrobacter (freundii, other species), 131 Clostridium (all species except difficile), 132 Clostridium difficile, 173 Corynebacterium jeikeium, 134 Enterobacter (all species), 135 Enterococcus (all species), 177 Enterococcus, vancomycin resistant (VRE), 136 Escherichia (also E. coli), 139 Fusobacterium (all species), 187 Haemophilus influenzae, 188 Haemophilus non-influenzae, 146 Klebsiella (all		Specify Organism Code(s):	Bacterial Infections: 121 inetobacter (all species), 125 Bordetella pertussis (whooping cough), 128 Campylobacter (all species), 129 Capnocytophaga (all species), 171 Chlamydia (pneumoniae), 130 Citrobacter (freundii, other species), 131 Clostridium (all species except difficile), 132 Clostridium difficile, 173 Corynebacterium jeikeium, 134 Enterobacter (all species), 135 Enterococcus (all species), 177 Enterococcus, vancomycin resistant (VRE), 136 Escherichia (also E. coli), 139 Fusobacterium (all species), 187 Haemophilus influenzae, 188 Haemophilus non-influenzae, 146 Klebsiella (all species), 147 Lactobacillus (bulgaricus, acidophilus, other species), 189 Legionella pneumophila, 190 Legionella non-pneumophila, 103 Leptospira (all species), 148 Leptotrichia buccalis, 149 Leuconostoc (all species), 104 Listeria monocytogenes, 151 Micrococcus, NOS, 118 Mycobacterium abscessus, 112 Mycobacterium avium - intracellulare (MAC, MAI), 108 Mycobacterium cheloneae, 109 Mycobacterium fortuitum, 114 Mycobacterium haemophilum, 115 Mycobacterium kansasii, 116 Mycobacterium marinum, 117 Mycobacterium mucogenicum, 110 Mycobacterium tuberculosis (tuberculosis, Koch bacillus), 105 Mycoplasma (all species), 183 Neisseria gonorrhoeae, 184 Neisseria meningitidis, 106 Nocardia (all species), 153 Pasteurella multocida, 155 Proteus (all species), 157 Pseudomonas or Burkholderia cepacia, 185 Pseudomonas aeruginosa, 186 Pseudomonas non-aeruginosa, 159 Rhodococcus (all species), 107 Rickettsia (all species), 160 Salmonella (all species), 161 Serratia marcescens, 162 Shigella (all species), 180 Staphylococcus aureus (Methicillin Resistant), 179 Staphylococcus aureus (Methicillin Sensitive), 158 Stenotrophomonas maltophilia, 166 Stomatococcus mucilaginosus, 181 Streptococcus, alpha-hemolytic, 182 Streptococcus, Group B, 178 Streptococcus pneumoniae, 168 Treponema (syphilis), 169 Vibrio (all species) Fungal	

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PRO201	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Product Infusion	Product Analysis	yes	yes	Specify Organism Code(s):	Bacterial Infections: 121 inetobacter (all species), 125 Bordetella pertussis (whooping cough), 128 Campylobacter (all species), 129 Capnocytophaga (all species), 171 Chlamydia (pneumoniae), 130 Citrobacter (freundii, other species), 131 Clostridium (all species except difficile), 132 Clostridium difficile, 173 Corynebacterium jeikeium, 134 Enterobacter (all species), 135 Enterococcus (all species), 177 Enterococcus, vancomycin resistant (VRE), 136 Escherichia (also E. coli), 139 Fusobacterium (all species), 187 Haemophilus influenzae, 188 Haemophilus non-influenzae, 146 Klebsiella (all		Specify Organism Code(s):	Bacterial Infections: 121 inetobacter (all species), 125 Bordetella pertussis (whooping cough), 128 Campylobacter (all species), 129 Capnocytophaga (all species), 171 Chlamydia (pneumoniae), 130 Citrobacter (freundii, other species), 131 Clostridium (all species except difficile), 132 Clostridium difficile, 173 Corynebacterium jeikeium, 134 Enterobacter (all species), 135 Enterococcus (all species), 177 Enterococcus, vancomycin resistant (VRE), 136 Escherichia (also E. coli), 139 Fusobacterium (all species), 187 Haemophilus influenzae, 188 Haemophilus non-influenzae, 146 Klebsiella (all	
PRO202	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Product Infusion		no	yes	Specify organism:	open text		Specify organism:	open text	
PRO203	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Product Infusion		no	yes	Date of this product infusion:	YYYY/MM/DD		Date of this product infusion:	YYYY/MM/DD	
PRO204	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Product Infusion		no	yes	Was the entire volume of received product infused?	no,yes		Was the entire volume of received product infused?	no,yes	
PRO205	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Product Infusion		no	yes	Specify what happened to the reserved portion	cryopreserved for future use,discarded,other fate		Specify what happened to the reserved portion	cryopreserved for future use,discarded,other fate	

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PRO206	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Product Infusion		no	yes	Specify other fate:	open text		Specify other fate:	open text	
PRO207	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Product Infusion		no	yes	Time product infusion initiated (24-hour clock):	Hour:Minute Check "standard time" or "check daylight savings time"		Time product infusion initiated (24-hour clock):	Hour:Minute Check "standard time" or "check daylight savings time"	
PRO208	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Product Infusion		no	yes	Date infusion stopped:	YYYY/MM/DD		Date infusion stopped:	YYYY/MM/DD	
PRO209	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Product Infusion		no	yes	Time product infusion completed (24-hour clock):	Hour:Minute Check "standard time" or "check daylight savings time"		Time product infusion completed (24-hour clock):	Hour:Minute Check "standard time" or "check daylight savings time"	
PRO210	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Product Infusion		no	yes	Specify the route of product infusion (24-hour clock);	Intramedullary,Intravenous,Other route of infusion		Specify the route of product infusion (24-hour clock);	Intramedullary,Intravenous,Other route of infusion	
PRO211	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Product Infusion		no	yes	Specify other route of infusion:	open text		Specify other route of infusion:	open text	
PRO212	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Product Infusion	Cord Blood Product Infusion	yes	no	Were there any adverse events or incidents associated with the stem cell infusion?	no,yes		Were there any adverse events or incidents associated with the stem cell infusion?	no,yes	
PRO213	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Product Infusion	Cord Blood Product Infusion	yes	no	Brachycardia	no,yes		Brachycardia	no,yes	
PRO214	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Product Infusion	Cord Blood Product Infusion	yes	no	In the Medical Director's judgment, was the adverse event a direct result of the infusion?	no,yes		In the Medical Director's judgment, was the adverse event a direct result of the infusion?	no,yes	

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PRO215	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Product Infusion	Cord Blood Product Infusion	yes	no	Chest tightness / pain	no,yes		Chest tightness / pain	no,yes	
PRO216	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Product Infusion	Cord Blood Product Infusion	yes	no	In the Medical Director's judgment, was the adverse event a direct result of the infusion?	no,yes		In the Medical Director's judgment, was the adverse event a direct result of the infusion?	no,yes	
PRO217	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Product Infusion	Cord Blood Product Infusion	yes	no	Chills at time of infusion	no,yes		Chills at time of infusion	no,yes	
PRO218	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Product Infusion	Cord Blood Product Infusion	yes	no	In the Medical Director's judgment, was the adverse event a direct result of the infusion?	no,yes		In the Medical Director's judgment, was the adverse event a direct result of the infusion?	no,yes	
PRO219	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Product Infusion	Cord Blood Product Infusion	yes	no	Fever ≤ 103 °F within 24 hours of infusion	no,yes		Fever ≤ 103 °F within 24 hours of infusion	no,yes	
PRO220	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Product Infusion	Cord Blood Product Infusion	yes	no	In the Medical Director's judgment, was the adverse event a direct result of the infusion?	no,yes		In the Medical Director's judgment, was the adverse event a direct result of the infusion?	no,yes	
PRO221	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Product Infusion	Cord Blood Product Infusion	yes	no	Fever > 103° F within 24 hours of infusion	no,yes		Fever > 103° F within 24 hours of infusion	no,yes	
PRO222	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Product Infusion	Cord Blood Product Infusion	yes	no	In the Medical Director's judgment, was the adverse event a direct result of the infusion?	no,yes		In the Medical Director's judgment, was the adverse event a direct result of the infusion?	no,yes	

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PRO223	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Product Infusion	Cord Blood Product Infusion	yes	no	Gross hemoglobinuria	no,yes		Gross hemoglobinuria	no,yes	
PRO224	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Product Infusion	Cord Blood Product Infusion	yes	no	In the Medical Director's judgment, was the adverse event a direct result of the infusion?	no,yes		In the Medical Director's judgment, was the adverse event a direct result of the infusion?	no,yes	
PRO225	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Product Infusion	Cord Blood Product Infusion	yes	no	Headache	no,yes		Headache	no,yes	
PRO226	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Product Infusion	Cord Blood Product Infusion	yes	no	In the Medical Director's judgment, was the adverse event a direct result of the infusion?	no,yes		In the Medical Director's judgment, was the adverse event a direct result of the infusion?	no,yes	
PRO227	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Product Infusion	Cord Blood Product Infusion	yes	no	Hives	no,yes		Hives	no,yes	
PRO228	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Product Infusion	Cord Blood Product Infusion	yes	no	In the Medical Director's judgment, was the adverse event a direct result of the infusion?	no,yes		In the Medical Director's judgment, was the adverse event a direct result of the infusion?	no,yes	
PRO229	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Product Infusion	Cord Blood Product Infusion	yes	no	Hypertension	no,yes		Hypertension	no,yes	
PRO230	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Product Infusion	Cord Blood Product Infusion	yes	no	In the Medical Director's judgment, was the adverse event a direct result of the infusion?	no,yes		In the Medical Director's judgment, was the adverse event a direct result of the infusion?	no,yes	

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PRO231	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Product Infusion	Cord Blood Product Infusion	yes	no	Hypotension	no,yes		Hypotension	no,yes	
PRO232	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Product Infusion	Cord Blood Product Infusion	yes	no	In the Medical Director's judgment, was the adverse event a direct result of the infusion?	no,yes		In the Medical Director's judgment, was the adverse event a direct result of the infusion?	no,yes	
PRO233	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Product Infusion	Cord Blood Product Infusion	yes	no	Hypoxia requiring oxygen (O ₂) support	no,yes		Hypoxia requiring oxygen (O ₂) support	no,yes	
PRO234	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Product Infusion	Cord Blood Product Infusion	yes	no	In the Medical Director's judgment, was the adverse event a direct result of the infusion?	no,yes		In the Medical Director's judgment, was the adverse event a direct result of the infusion?	no,yes	
PRO235	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Product Infusion	Cord Blood Product Infusion	yes	no	Nausea	no,yes		Nausea	no,yes	
PRO236	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Product Infusion	Cord Blood Product Infusion	yes	no	In the Medical Director's judgment, was the adverse event a direct result of the infusion?	no,yes		In the Medical Director's judgment, was the adverse event a direct result of the infusion?	no,yes	
PRO237	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Product Infusion	Cord Blood Product Infusion	yes	no	Rigors, mild	no,yes		Rigors, mild	no,yes	
PRO238	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Product Infusion	Cord Blood Product Infusion	yes	no	In the Medical Director's judgment, was the adverse event a direct result of the infusion?	no,yes		In the Medical Director's judgment, was the adverse event a direct result of the infusion?	no,yes	

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PRO239	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Product Infusion	Cord Blood Product Infusion	yes	no	Rigors, severe	no,yes		Rigors, severe	no,yes	
PRO240	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Product Infusion	Cord Blood Product Infusion	yes	no	In the Medical Director's judgment, was the adverse event a direct result of the infusion?	no,yes		In the Medical Director's judgment, was the adverse event a direct result of the infusion?	no,yes	
PRO241	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Product Infusion	Cord Blood Product Infusion	yes	no	Shortness of breath (SOB)	no,yes		Shortness of breath (SOB)	no,yes	
PRO242	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Product Infusion	Cord Blood Product Infusion	yes	no	In the Medical Director's judgment, was the adverse event a direct result of the infusion?	no,yes		In the Medical Director's judgment, was the adverse event a direct result of the infusion?	no,yes	
PRO243	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Product Infusion	Cord Blood Product Infusion	yes	no	Tachycardia	no,yes		Tachycardia	no,yes	
PRO244	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Product Infusion	Cord Blood Product Infusion	yes	no	In the Medical Director's judgment, was the adverse event a direct result of the infusion?	no,yes		In the Medical Director's judgment, was the adverse event a direct result of the infusion?	no,yes	
PRO245	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Product Infusion	Cord Blood Product Infusion	yes	no	Vomiting	no,yes		Vomiting	no,yes	
PRO246	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Product Infusion	Cord Blood Product Infusion	yes	no	In the Medical Director's judgment, was the adverse event a direct result of the infusion?	no,yes		In the Medical Director's judgment, was the adverse event a direct result of the infusion?	no,yes	

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PRO247	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Product Infusion	Cord Blood Product Infusion	yes	no	Other expected AE	no,yes		Other expected AE	no,yes	
PRO248	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Product Infusion	Cord Blood Product Infusion	yes	no	Specify other expected AE:	open text		Specify other expected AE:	open text	
PRO249	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Product Infusion	Cord Blood Product Infusion	yes	no	In the Medical Director's judgment, was the adverse event a direct result of the infusion?	no,yes		In the Medical Director's judgment, was the adverse event a direct result of the infusion?	no,yes	
PRO250	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Product Infusion	Cord Blood Product Infusion	yes	no	Other unexpected AE	no,yes		Other unexpected AE	no,yes	
PRO251	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Product Infusion	Cord Blood Product Infusion	yes	no	Specify other unexpected AE:	open text		Specify other unexpected AE:	open text	
PRO252	Transplant Procedure and Product Information	Hematopoietic Cellular Transplant (HCT) Product Infusion	Cord Blood Product Infusion	yes	no	In the Medical Director's judgment, was the adverse event a direct result of the infusion?	no,yes		In the Medical Director's judgment, was the adverse event a direct result of the infusion?	no,yes	
PRO253	Transplant Procedure and Product Information	Infectious Disease Markers			yes	Sequence Number:	Auto Filled Field		Sequence Number:	Auto Filled Field	
PRO254	Transplant Procedure and Product Information	Infectious Disease Markers			yes	Date Received:	Auto Filled Field		Date Received:	Auto Filled Field	
PRO255	Transplant Procedure and Product Information	Infectious Disease Markers			yes	CIBMTR Center Number:	Auto Filled Field		CIBMTR Center Number:	Auto Filled Field	
PRO256	Transplant Procedure and Product Information	Infectious Disease Markers			yes	CIBMTR Research ID:	Auto Filled Field		CIBMTR Research ID:	Auto Filled Field	

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PRO257	Transplant Procedure and Product Information	Infectious Disease Markers				Event date:	Auto Filled Field created with CRID		Event date:	Auto Filled Field created with CRID	
PRO258	Transplant Procedure and Product Information	Infectious Disease Markers		no	no	HCT type (check all that apply)	Allogeneic, related,Allogeneic, unrelated		HCT type (check all that apply)	Allogeneic, related,Allogeneic, unrelated	
PRO259	Transplant Procedure and Product Information	Infectious Disease Markers		no	no	Product type (check all that apply)	Bone marrow,Other product,PBSC,Singl e cord blood unit		Product type (check all that apply)	Bone marrow,Other product,PBSC,Single cord blood unit	
PRO260	Transplant Procedure and Product Information	Infectious Disease Markers		no	no	Other product. Specify:	open text		Other product. Specify:	open text	
PRO261	Transplant Procedure and Product Information	Infectious Disease Markers		no	no	Registry donor ID:	open text		Registry donor ID:	open text	
PRO262	Transplant Procedure and Product Information	Infectious Disease Markers		no	no	Non-NMDP cord blood unit ID:	open text		Non-NMDP cord blood unit ID:	open text	
PRO263	Transplant Procedure and Product Information	Infectious Disease Markers		no	no	Global Registration Identifier for Donors (GRID)	open text		Global Registration Identifier for Donors (GRID)	open text	
PRO264	Transplant Procedure and Product Information	Infectious Disease Markers		no	no	ISBT DIN:	open text		ISBT DIN:	open text	

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PRO265	Transplant Procedure and Product Information	Infectious Disease Markers		no	no	Registry or UCB Bank ID	(A) Austrian Bone Marrow Donors, (ACB) Austrian Cord Blood Registry, (ACCB) StemCyte, Inc., (AE) Emirates Bone Marrow Donor Registry, (AM) Armenian Bone Marrow Donor Registry Charitable Trust, (AOCB) University of Colorado Cord Blood Bank, (AR) Argentine CPH Donors Registry, (ARCB) BANCEL - Argentina Cord Blood Bank, (AUCB) Australian Cord Blood Registry, (AUS) Australian / New Zealand Bone Marrow Donor Registry, (B) Marrow Donor Program Belgium, (BCB) Belgium Cord Blood Registry, (BG) Bulgarian Bone Marrow Donor Registry, (BR) INCA/REDOMO, (BSCB) British Bone Marrow Registry - Cord Blood, (CB)		Registry or UCB Bank ID	(A) Austrian Bone Marrow Donors, (ACB) Austrian Cord Blood Registry, (ACCB) StemCyte, Inc., (AE) Emirates Bone Marrow Donor Registry, (AM) Armenian Bone Marrow Donor Registry Charitable Trust, (AOCB) University of Colorado Cord Blood Bank, (AR) Argentine CPH Donors Registry, (ARCB) BANCEL - Argentina Cord Blood Bank, (AUCB) Australian Cord Blood Registry, (AUS) Australian / New Zealand Bone Marrow Donor Registry, (B) Marrow Donor Program Belgium, (BCB) Belgium Cord Blood Registry, (BG) Bulgarian Bone Marrow Donor Registry, (BR) INCA/REDOMO, (BSCB) British Bone Marrow Registry - Cord Blood, (CB) Cord Blood Registry, (CH) Swiss BloodStem Cells - Adult Donors, (CHCB) Swiss Blood Stem Cells - Cord Blood, (CKCB) Celgene Cord Blood Bank, (CN) China Marrow Donor Program (CMDP), (CNCB) Shan Dong Cord Blood Bank, (CND) Canadian Blood Services Bone Marrow Donor Registry, (CS2) Czech National Marrow Donor Registry, (CSCR) Czech Stem Cells Registry, (CY) Cyprus Paraskevaudio Bone Marrow Donor Registry, (CY2) The Cyprus Bone Marrow Donor Registry, (D) ZKRD - Zentrales Knochenmarkspender - Register Deutschland Adult Donors, (DCB) ZKRD - Zentrales Knochenmarkspender - Register Deutschland Cord Blood, (DK) The Danish Bone Marrow Donor Registry, (DK2) Bone Marrow Donors Copenhagen (BMDC), (DUCB) German Branch of the European Cord Blood Bank, (E) REDMO, (ECB) Spanish Cord Blood Registry, (F) France Greffe de Moelle - Adult Donors, (FCB) France Greffe de Moelle - Cord Blood, (FI) Finnish Bone Marrow Donor Registry, (FICB) Finnish Cord Blood Registry, (GB) The Anthony Nolan Trust, (GB3) Welsh Bone Marrow Donor Registry, (GB4) British Bone Marrow Registry, (GR) Unrelated Hematopoietic Stem Cell Donor Registry Greece, (GRCB) Michigan Community Blood Centers Cord Blood Bank, (H) Hungarian Bone Marrow Donor Registry, (HEM) Hema-Quebec, (HK) Hong Kong Bone Marrow Donor Registry, (HR) Croatian Bone Marrow Donor	
PRO266	Transplant Procedure and Product Information	Infectious Disease Markers		no	no	Donor DOB:	YYYY/MM/DD		Donor DOB:	YYYY/MM/DD	
PRO267	Transplant Procedure and Product Information	Infectious Disease Markers		no	no	Donor age:	open text, check "Months" or check "Years"		Donor age:	open text, check "Months" or check "Years"	
PRO268	Transplant Procedure and Product Information	Infectious Disease Markers		no	no	Donor sex	female,male		Donor sex	female,male	
PRO269	Transplant Procedure and Product Information	Infectious Disease Markers	Non NMDP Allogeneic or syngeneic Donor or Non NMDP Cord Blood Unit Information	yes	no	Who is being tested for IDMs?	donor IDM (marrow or PBSC),cord blood unit IDM,maternal IDM (cord blood)		Who is being tested for IDMs?	donor IDM (marrow or PBSC),cord blood unit IDM,maternal IDM (cord blood)	

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PRO270	Transplant Procedure and Product Information	Infectious Disease Markers	Non NMDP Allogeneic or syngeneic Donor or Non NMDP Cord Blood Unit Information	yes	no	HBsAg: (hepatitis B surface antigen)	Non-reactive,Not done,Reactive		HBsAg: (hepatitis B surface antigen)	Non-reactive,Not done,Reactive	
PRO271	Transplant Procedure and Product Information	Infectious Disease Markers	Non NMDP Allogeneic or syngeneic Donor or Non NMDP Cord Blood Unit Information	yes	no	Date sample collected:	YYYY/MM/DD		Date sample collected:	YYYY/MM/DD	
PRO272	Transplant Procedure and Product Information	Infectious Disease Markers	Non NMDP Allogeneic or syngeneic Donor or Non NMDP Cord Blood Unit Information	yes	no	Anti HBC: (hepatitis B core antibody)	Non-reactive,Not done,Reactive		Anti HBC: (hepatitis B core antibody)	Non-reactive,Not done,Reactive	
PRO273	Transplant Procedure and Product Information	Infectious Disease Markers	Non NMDP Allogeneic or syngeneic Donor or Non NMDP Cord Blood Unit Information	yes	no	Date sample collected:	YYYY/MM/DD		Date sample collected:	YYYY/MM/DD	
PRO274	Transplant Procedure and Product Information	Infectious Disease Markers	Non NMDP Allogeneic or syngeneic Donor or Non NMDP Cord Blood Unit Information	yes	no	FDA licensed NAAT testing for HBV	Negative,Not done,Positive		FDA licensed NAAT testing for HBV	Negative,Not done,Positive	
PRO275	Transplant Procedure and Product Information	Infectious Disease Markers	Non NMDP Allogeneic or syngeneic Donor or Non NMDP Cord Blood Unit Information	yes	no	Date sample collected:	YYYY/MM/DD		Date sample collected:	YYYY/MM/DD	
PRO276	Transplant Procedure and Product Information	Infectious Disease Markers	Non NMDP Allogeneic or syngeneic Donor or Non NMDP Cord Blood Unit Information	yes	no	Anti-HCV: (hepatitis C antibody)	Non-reactive,Not done,Reactive		Anti-HCV: (hepatitis C antibody)	Non-reactive,Not done,Reactive	
PRO277	Transplant Procedure and Product Information	Infectious Disease Markers	Non NMDP Allogeneic or syngeneic Donor or Non NMDP Cord Blood Unit Information	yes	no	Date sample collected:	YYYY/MM/DD		Date sample collected:	YYYY/MM/DD	

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PRO278	Transplant Procedure and Product Information	Infectious Disease Markers	Non NMDP Allogeneic or syngeneic Donor or Non NMDP Cord Blood Unit Information	yes	no	FDA licensed NAAT testing for HCV	Negative,Not done,Positive		FDA licensed NAAT testing for HCV	Negative,Not done,Positive	
PRO279	Transplant Procedure and Product Information	Infectious Disease Markers	Non NMDP Allogeneic or syngeneic Donor or Non NMDP Cord Blood Unit Information	yes	no	Date sample collected:	YYYY/MM/DD		Date sample collected:	YYYY/MM/DD	
PRO280	Transplant Procedure and Product Information	Infectious Disease Markers	Non NMDP Allogeneic or syngeneic Donor or Non NMDP Cord Blood Unit Information	yes	no	HIV-1 p24 antigen	Non-reactive,Not done,Not reported,Reactive		HIV-1 p24 antigen	Non-reactive,Not done,Not reported,Reactive	
PRO281	Transplant Procedure and Product Information	Infectious Disease Markers	Non NMDP Allogeneic or syngeneic Donor or Non NMDP Cord Blood Unit Information	yes	no	Date sample collected:	YYYY/MM/DD		Date sample collected:	YYYY/MM/DD	
PRO282	Transplant Procedure and Product Information	Infectious Disease Markers	Non NMDP Allogeneic or syngeneic Donor or Non NMDP Cord Blood Unit Information	yes	no	FDA licensed NAAT testing for HIV-1	Negative,Not done,Positive		FDA licensed NAAT testing for HIV-1	Negative,Not done,Positive	
PRO283	Transplant Procedure and Product Information	Infectious Disease Markers	Non NMDP Allogeneic or syngeneic Donor or Non NMDP Cord Blood Unit Information	yes	no	Date sample collected:	YYYY/MM/DD		Date sample collected:	YYYY/MM/DD	
PRO284	Transplant Procedure and Product Information	Infectious Disease Markers	Non NMDP Allogeneic or syngeneic Donor or Non NMDP Cord Blood Unit Information	yes	no	Anti-HIV 1 and anti-HIV 2*: (antibodies to Human Immunodeficiency Viruses)	Non-reactive,Not done,Not reported,Reactive		Anti-HIV 1 and anti-HIV 2*: (antibodies to Human Immunodeficiency Viruses)	Non-reactive,Not done,Not reported,Reactive	
PRO285	Transplant Procedure and Product Information	Infectious Disease Markers	Non NMDP Allogeneic or syngeneic Donor or Non NMDP Cord Blood Unit Information	yes	no	Date sample collected:	YYYY/MM/DD		Date sample collected:	YYYY/MM/DD	

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PRO286	Transplant Procedure and Product Information	Infectious Disease Markers	Non NMDP Allogeneic or syngeneic Donor or Non NMDP Cord Blood Unit Information	yes	no	Chagas testing	Negative,Not Done,Positive		Chagas testing	Negative,Not Done,Positive	
PRO287	Transplant Procedure and Product Information	Infectious Disease Markers	Non NMDP Allogeneic or syngeneic Donor or Non NMDP Cord Blood Unit Information	yes	no	Date sample collected:	YYYY/MM/DD		Date sample collected:	YYYY/MM/DD	
PRO288	Transplant Procedure and Product Information	Infectious Disease Markers	Non NMDP Allogeneic or syngeneic Donor or Non NMDP Cord Blood Unit Information	yes	no	Anti-HSV (Herpes simplex virus antibody)	Negative,Not Done,Positive		Anti-HSV (Herpes simplex virus antibody)	Negative,Not Done,Positive	
PRO289	Transplant Procedure and Product Information	Infectious Disease Markers	Non NMDP Allogeneic or syngeneic Donor or Non NMDP Cord Blood Unit Information	yes	no	Date sample collected:	YYYY/MM/DD		Date sample collected:	YYYY/MM/DD	
PRO290	Transplant Procedure and Product Information	Infectious Disease Markers	Non NMDP Allogeneic or syngeneic Donor or Non NMDP Cord Blood Unit Information	yes	no	Anti-EBV (Epstein-Barr virus antibody)	Inconclusive,Negative,Not done,Positive		Anti-EBV (Epstein-Barr virus antibody)	Inconclusive,Negative,Not done,Positive	
PRO291	Transplant Procedure and Product Information	Infectious Disease Markers	Non NMDP Allogeneic or syngeneic Donor or Non NMDP Cord Blood Unit Information	yes	no	Date sample collected:	YYYY/MM/DD		Date sample collected:	YYYY/MM/DD	
PRO292	Transplant Procedure and Product Information	Infectious Disease Markers	Non NMDP Allogeneic or syngeneic Donor or Non NMDP Cord Blood Unit Information	yes	no	Anti-VZV (Varicella zoster virus antibody)	Negative,Not Done,Positive		Anti-VZV (Varicella zoster virus antibody)	Negative,Not Done,Positive	
PRO293	Transplant Procedure and Product Information	Infectious Disease Markers	Non NMDP Allogeneic or syngeneic Donor or Non NMDP Cord Blood Unit Information	yes	no	Date sample collected:	YYYY/MM/DD		Date sample collected:	YYYY/MM/DD	

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PRO294	Transplant Procedure and Product Information	Infectious Disease Markers	Non NMDP Allogeneic or syngeneic Donor or Non NMDP Cord Blood Unit Information	yes	no	Other infectious disease marker, specify	no,yes		Other infectious disease marker, specify	no,yes	
PRO295	Transplant Procedure and Product Information	Infectious Disease Markers	Non NMDP Allogeneic or syngeneic Donor or Non NMDP Cord Blood Unit Information	yes	no	Date sample collected:	YYYY/MM/DD		Date sample collected:	YYYY/MM/DD	
PRO296	Transplant Procedure and Product Information	Infectious Disease Markers	Non NMDP Allogeneic or syngeneic Donor or Non NMDP Cord Blood Unit Information	yes	no	Specify test and method:	open text		Specify test and method:	open text	
PRO297	Transplant Procedure and Product Information	Infectious Disease Markers	Non NMDP Allogeneic or syngeneic Donor or Non NMDP Cord Blood Unit Information	yes	no	Specify test results:	open text		Specify test results:	open text	



Information Collection Domain: Post-Transplant Periodic Information Collection

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POST001	Post-Transplant	Post-Transplant Essential Data		no	yes	Sequence Number:	Auto Filled Field		Sequence Number:	Auto Filled Field	
POST002	Post-Transplant	Post-Transplant Essential Data		no	yes	Date Received:	Auto Filled Field		Date Received:	Auto Filled Field	
POST003	Post-Transplant	Post-Transplant Essential Data		no	yes	CIBMTR Center Number:	Auto Filled Field		CIBMTR Center Number:	Auto Filled Field	
POST004	Post-Transplant	Post-Transplant Essential Data		no	yes	CIBMTR Research ID:	Auto Filled Field		CIBMTR Research ID:	Auto Filled Field	
POST005	Post-Transplant	Post-Transplant Essential Data		no	yes	Event date:	Auto Filled Field created with CRID		Event date:	Auto Filled Field created with CRID	
POST006	Post-Transplant	Post-Transplant Essential Data		no	yes	Visit	100 day,1 year,2 years,> 2 years,6 months		Visit	100 day,1 year,2 years,> 2 years,6 months	
POST007	Post-Transplant	Post-Transplant Essential Data		no	yes	Specify:	open text		Specify:	open text	
POST008	Post-Transplant	Post-Transplant Essential Data		no	yes	Date of actual contact with the recipient to determine medical status for this follow-up report:	YYYY/MM/DD		Date of actual contact with the recipient to determine medical status for this follow-up report:	YYYY/MM/DD	
POST009	Post-Transplant	Post-Transplant Essential Data		no	yes	Specify the recipient's survival status at the date of last contact	Alive,Dead		Specify the recipient's survival status at the date of last contact	Alive,Dead (Complete recipient death data)	
POST010	Post-Transplant	Post-Transplant Essential Data		no	yes	Did the recipient receive a subsequent HCT?	no,yes		Did the recipient receive a subsequent HCT?	no,yes	
POST011	Post-Transplant	Post-Transplant Essential Data	Subsequent Transplant	yes	yes	Date of subsequent HCT:	YYYY/MM/DD		Date of subsequent HCT:	YYYY/MM/DD	
POST012	Post-Transplant	Post-Transplant Essential Data	Subsequent Transplant	yes	yes	What was the indication for subsequent HCT?	Graft failure / insufficient hematopoietic recovery,Insufficient chimerism,New malignancy (including PTLD and EBV lymphoma),Other,Persistent primary disease,Planned subsequent HCT, per protocol,Recurrent primary disease		What was the indication for subsequent HCT?	Graft failure / insufficient hematopoietic recovery,Insufficient chimerism,New malignancy (including PTLD and EBV lymphoma),Other,Persistent primary disease,Planned subsequent HCT, per protocol,Recurrent primary disease	
POST013	Post-Transplant	Post-Transplant Essential Data	Subsequent Transplant	yes	yes	Specify other indication:	open text		Specify other indication:	open text	
POST014	Post-Transplant	Post-Transplant Essential Data	Subsequent Transplant	yes	yes	Source of HSCs (check all that apply)	Allogeneic, related,Allogeneic, unrelated,Autologous		Source of HSCs (check all that apply)	Allogeneic, related,Allogeneic, unrelated,Autologous	

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POST015	Post-Transplant	Post-Transplant Essential Data		no	yes	Has the recipient received a cellular therapy? (e.g. CAR-T, DCI)	no,yes		Has the recipient received a cellular therapy? (e.g. CAR-T, DCI)	no,yes	
POST016	Post-Transplant	Post-Transplant Essential Data	Subsequent Transplant	yes	yes				Was this infusion a donor lymphocyte infusion (DLI)?	no,yes	
POST017	Post-Transplant	Post-Transplant Essential Data	Subsequent Transplant	yes	yes				Number of DLIs in this reporting period	___ ___	
POST018	Post-Transplant	Post-Transplant Essential Data	Subsequent Transplant	yes	yes				Are any of the products, associated with this course of cellular therapy, genetically modified?	no, yes	
POST019	Post-Transplant	Post-Transplant Essential Data	Subsequent Transplant	yes	yes	Date of cellular therapy:	YYYY/MM/DD		Date of cellular therapy:	YYYY/MM/DD	
POST020	Post-Transplant	Post-Transplant Essential Data		no	yes	Was there evidence of initial hematopoietic recovery?	No(ANC ≥ 500/mm3 was not achieved) ,Not applicable(ANC never dropped below 500/mm3 at any time after the start of the preparative regimen,Previously reported(recipient's initial hematopoietic recovery was recorded on a previous report) ,Yes(ANC ≥ 500/mm3 achieved and sustained for 3 lab values)		Was there evidence of initial hematopoietic recovery?	No(ANC ≥ 500/mm3 was not achieved) ,Not applicable(ANC never dropped below 500/mm3 at any time after the start of the preparative regimen,Previously reported(recipient's initial hematopoietic recovery was recorded on a previous report) ,Yes(ANC ≥ 500/mm3 achieved and sustained for 3 lab values)	
POST021	Post-Transplant	Post-Transplant Essential Data		no	yes	Date ANC ≥ 500/mm ³ (first of 3 lab values):	YYYY/MM/DD		Date ANC ≥ 500/mm ³ (first of 3 lab values):	YYYY/MM/DD	
POST022	Post-Transplant	Post-Transplant Essential Data		no	yes	Did late graft failure occur?	No,Yes		Did late graft failure occur?	No,Yes	
POST023	Post-Transplant	Post-Transplant Essential Data		no	yes	Was an initial platelet count ≥ 20 x 10 ⁹ /L achieved?	No,Not applicable(Platelet count never dropped below 20 x 10 ⁹ /L) ,Previously reported(≥ 20 x 10 ⁹ /L was achieved and reported previously),Yes		Was an initial platelet count ≥ 20 x 10 ⁹ /L achieved?	No,Not applicable(Platelet count never dropped below 20 x 10 ⁹ /L) ,Previously reported(≥ 20 x 10 ⁹ /L was achieved and reported previously),Yes	
POST024	Post-Transplant	Post-Transplant Essential Data		no	yes	Date platelets ≥ 20 x 10 ⁹ /L:	YYYY/MM/DD		Date platelets ≥ 20 x 10 ⁹ /L:	YYYY/MM/DD	
POST025	Post-Transplant	Post-Transplant Essential Data		no	yes	Did acute GVHD develop?	No,Unknown,Yes		Did acute GVHD develop?	No,Unknown,Yes	
POST026	Post-Transplant	Post-Transplant Essential Data	Graft vs. Host Disease	yes	yes	Date of acute GVHD diagnosis:	YYYY/MM/DD		Date of acute GVHD diagnosis:	YYYY/MM/DD	
POST027	Post-Transplant	Post-Transplant Essential Data	Graft vs. Host Disease	yes	yes	Did acute GVHD persist?	No,Unknown,Yes		Did acute GVHD persist?	No,Unknown,Yes	

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POST028	Post-Transplant	Post-Transplant Essential Data	Graft vs. Host Disease	yes	yes	Overall grade of acute GVHD at diagnosis	I - Rash on ≤ 50% of skin, no liver or gut involvement II - Rash on > 50% of skin, bilirubin 2-3 mg/dL, or diarrhea 500 – 1000 mL/day or persistent nausea or vomiting III - Bilirubin 3-15 mg/dL, or gut stage 2-4 diarrhea > 1000 mL/day or severe abdominal pain with or without ileus IV - Generalized erythroderma with bullous formation, or bilirubin >15 mg/dL Not applicable (acute GVHD present but cannot be graded)		Overall grade of acute GVHD at diagnosis	I - Rash on ≤ 50% of skin, no liver or gut involvement II - Rash on > 50% of skin, bilirubin 2-3 mg/dL, or diarrhea 500 – 1000 mL/day or persistent nausea or vomiting III - Bilirubin 3-15 mg/dL, or gut stage 2-4 diarrhea > 1000 mL/day or severe abdominal pain with or without ileus IV - Generalized erythroderma with bullous formation, or bilirubin >15 mg/dL Not applicable (acute GVHD present but cannot be graded)	
POST029	Post-Transplant	Post-Transplant Essential Data	Graft vs. Host Disease	yes	yes	Skin	Stage 0 – No rash, no rash attributable to acute GVHD Stage 1 – Maculopapular rash, < 25% of body surface Stage 2 – Maculopapular rash, 25–50% of body surface Stage 3 – Generalized erythroderma, > 50% of body surface Stage 4 – Generalized erythroderma with bullae formation and/or desquamation		Skin	Stage 0 – No rash, no rash attributable to acute GVHD Stage 1 – Maculopapular rash, < 25% of body surface Stage 2 – Maculopapular rash, 25–50% of body surface Stage 3 – Generalized erythroderma, > 50% of body surface Stage 4 – Generalized erythroderma with bullae formation and/or desquamation	
POST030	Post-Transplant	Post-Transplant Essential Data	Graft vs. Host Disease	yes	yes	Lower intestinal tract (use mL/day for adult recipients and mL/kg/day for pediatric recipients)	Stage 0 – No diarrhea, no diarrhea attributable to acute GVHD / diarrhea < 500 mL/day (adult), or < 10 mL/kg/day (pediatric) Stage 1 – Diarrhea 500 - 1000 mL/day (adult), or 10 - 19.9 mL/kg/day (pediatric) Stage 2 – Diarrhea 1001 - 1500 mL/day (adult), or 20 - 30 mL/kg/day (pediatric) Stage 3 – Diarrhea > 1500 mL/day (adult), or > 30 mL/kg/day (pediatric) Stage 4 – Severe abdominal pain, with or without ileus, and/or grossly bloody stool		Lower intestinal tract (use mL/day for adult recipients and mL/kg/day for pediatric recipients)	Stage 0 – No diarrhea, no diarrhea attributable to acute GVHD / diarrhea < 500 mL/day (adult), or < 10 mL/kg/day (pediatric) Stage 1 – Diarrhea 500 - 1000 mL/day (adult), or 10 - 19.9 mL/kg/day (pediatric) Stage 2 – Diarrhea 1001 - 1500 mL/day (adult), or 20 - 30 mL/kg/day (pediatric) Stage 3 – Diarrhea > 1500 mL/day (adult), or > 30 mL/kg/day (pediatric) Stage 4 – Severe abdominal pain, with or without ileus, and/or grossly bloody stool	
POST031	Post-Transplant	Post-Transplant Essential Data	Graft vs. Host Disease	yes	yes	Upper intestinal tract	Stage 0 – No persistent nausea or vomiting Stage 1 – Persistent nausea or vomiting		Upper intestinal tract	Stage 0 – No persistent nausea or vomiting Stage 1 – Persistent nausea or vomiting	
POST032	Post-Transplant	Post-Transplant Essential Data	Graft vs. Host Disease	yes	yes	Liver	Stage 0 – No liver acute GVHD / bilirubin < 2.0 mg/dL (< 34 µmol/L) Stage 1 – Bilirubin 2.0–3.0 mg/dL (34–52 µmol/L) Stage 2 – Bilirubin 3.1–6.0 mg/dL (53–103 µmol/L) Stage 3 – Bilirubin 6.1–15.0 mg/dL (104–256 µmol/L) Stage 4 – Bilirubin > 15.0 mg/dL (> 256 µmol/L)		Liver	Stage 0 – No liver acute GVHD / bilirubin < 2.0 mg/dL (< 34 µmol/L) Stage 1 – Bilirubin 2.0–3.0 mg/dL (34–52 µmol/L) Stage 2 – Bilirubin 3.1–6.0 mg/dL (53–103 µmol/L) Stage 3 – Bilirubin 6.1–15.0 mg/dL (104–256 µmol/L) Stage 4 – Bilirubin > 15.0 mg/dL (> 256 µmol/L)	
POST033	Post-Transplant	Post-Transplant Essential Data	Graft vs. Host Disease	yes	yes	Other site(s) involved with acute GVHD	No,Yes		Other site(s) involved with acute GVHD	No,Yes	

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POST034	Post-Transplant	Post-Transplant Essential Data	Graft vs. Host Disease	yes	yes	Specify other site(s):	open text		Specify other site(s):	open text	
POST035	Post-Transplant	Post-Transplant Essential Data	Graft vs. Host Disease	yes	yes	Maximum overall grade of acute GVHD	I - Rash on ≤ 50% of skin, no liver or gut involvement II - Rash on > 50% of skin, bilirubin 2-3 mg/dL, or diarrhea 500 - 1000 mL/day or persistent nausea or vomiting III - Bilirubin 3-15 mg/dL, or gut stage 2-4 diarrhea > 1000 mL/day or severe abdominal pain with or without ileus IV - Generalized erythroderma with bullous formation, or bilirubin >15 mg/dL Not applicable (acute GVHD present but cannot be graded)		Maximum overall grade of acute GVHD	I - Rash on ≤ 50% of skin, no liver or gut involvement II - Rash on > 50% of skin, bilirubin 2-3 mg/dL, or diarrhea 500 - 1000 mL/day or persistent nausea or vomiting III - Bilirubin 3-15 mg/dL, or gut stage 2-4 diarrhea > 1000 mL/day or severe abdominal pain with or without ileus IV - Generalized erythroderma with bullous formation, or bilirubin >15 mg/dL Not applicable (acute GVHD present but cannot be graded)	
POST036	Post-Transplant	Post-Transplant Essential Data	Graft vs. Host Disease	yes	yes	Date maximum overall grade of acute GVHD:	YYYY/MM/DD		First date maximum overall grade of acute GVHD:	YYYY/MM/DD	
POST037	Post-Transplant	Post-Transplant Essential Data	Graft vs. Host Disease	yes	yes	Skin	Stage 0 – No rash, no rash attributable to acute GVHD Stage 1 – Maculopapular rash, < 25% of body surface Stage 2 – Maculopapular rash, 25–50% of body surface Stage 3 – Generalized erythroderma, > 50% of body surface Stage 4 – Generalized erythroderma with bullae formation and/or desquamation		Skin	Stage 0 – No rash, no rash attributable to acute GVHD Stage 1 – Maculopapular rash, < 25% of body surface Stage 2 – Maculopapular rash, 25–50% of body surface Stage 3 – Generalized erythroderma, > 50% of body surface Stage 4 – Generalized erythroderma with bullae formation and/or desquamation	
POST038	Post-Transplant	Post-Transplant Essential Data	Graft vs. Host Disease	yes	yes	Lower intestinal tract (use mL/day for adult recipients and mL/kg/day for pediatric recipients)	Stage 0 – No diarrhea, no diarrhea attributable to acute GVHD / diarrhea < 500 mL/day (adult), or < 10 mL/kg/day (pediatric) Stage 1 – Diarrhea 500 - 1000 mL/day (adult), or 10 - 19.9 mL/kg/day (pediatric) Stage 2 – Diarrhea 1001 - 1500 mL/day (adult), or 20 - 30 mL/kg/day (pediatric) Stage 3 – Diarrhea > 1500 mL/day (adult), or > 30 mL/kg/day (pediatric) Stage 4 – Severe abdominal pain, with or without ileus, and/or grossly bloody stool		Lower intestinal tract (use mL/day for adult recipients and mL/kg/day for pediatric recipients)	Stage 0 – No diarrhea, no diarrhea attributable to acute GVHD / diarrhea < 500 mL/day (adult), or < 10 mL/kg/day (pediatric) Stage 1 – Diarrhea 500 - 1000 mL/day (adult), or 10 - 19.9 mL/kg/day (pediatric) Stage 2 – Diarrhea 1001 - 1500 mL/day (adult), or 20 - 30 mL/kg/day (pediatric) Stage 3 – Diarrhea > 1500 mL/day (adult), or > 30 mL/kg/day (pediatric) Stage 4 – Severe abdominal pain, with or without ileus, and/or grossly bloody stool	
POST039	Post-Transplant	Post-Transplant Essential Data	Graft vs. Host Disease	yes	yes	Upper intestinal tract	Stage 0 – No persistent nausea or vomiting Stage 1 – Persistent nausea or vomiting		Upper intestinal tract	Stage 0 – No persistent nausea or vomiting Stage 1 – Persistent nausea or vomiting	

Item ID	Time Point	Information Collection Domain Sub-Type	Information Collection Domain Additional Sub Domain	Response required if Additional Sub Domain applies	Information Collection may be requested multiple times	Current Information Collection Data Element (if applicable)	Current Information Collection Data Element Response Option(s)	Information Collection update:	Proposed Information Collection Data Element (if applicable)	Proposed Information Collection Data Element Response Option(s)	Rationale for Information Collection Update
POST040	Post-Transplant	Post-Transplant Essential Data	Graft vs. Host Disease	yes	yes	Liver	Stage 0 – No liver acute GVHD / bilirubin < 2.0 mg/dL (< 34 µmol/L) Stage 1 – Bilirubin 2.0–3.0 mg/dL (34–52 µmol/L) Stage 2 – Bilirubin 3.1–6.0 mg/dL (53–103 µmol/L) Stage 3 – Bilirubin 6.1–15.0 mg/dL (104–256 µmol/L) Stage 4 – Bilirubin > 15.0 mg/dL (> 256 µmol/L)		Liver	Stage 0 – No liver acute GVHD / bilirubin < 2.0 mg/dL (< 34 µmol/L) Stage 1 – Bilirubin 2.0–3.0 mg/dL (34–52 µmol/L) Stage 2 – Bilirubin 3.1–6.0 mg/dL (53–103 µmol/L) Stage 3 – Bilirubin 6.1–15.0 mg/dL (104–256 µmol/L) Stage 4 – Bilirubin > 15.0 mg/dL (> 256 µmol/L)	
POST041	Post-Transplant	Post-Transplant Essential Data	Graft vs. Host Disease	yes	yes	Other site(s) involved with acute GVHD	No,Yes		Other site(s) involved with acute GVHD	No,Yes	
POST042	Post-Transplant	Post-Transplant Essential Data	Graft vs. Host Disease	yes	yes	Specify other site(s):	open text		Specify other site(s):	open text	
POST043	Post-Transplant	Post-Transplant Essential Data	Graft vs. Host Disease	yes	yes	Did chronic GVHD develop?	No,Unknown,Yes		Did chronic GVHD develop?	No,Unknown,Yes	
POST044	Post-Transplant	Post-Transplant Essential Data	Graft vs. Host Disease	yes	yes	Date of chronic GVHD diagnosis:	YYYY/MM/DD		Date of chronic GVHD diagnosis:	YYYY/MM/DD	
POST045	Post-Transplant	Post-Transplant Essential Data	Graft vs. Host Disease	yes	yes	Did chronic GVHD persist?	No,Unknown,Yes		Did chronic GVHD persist?	No,Unknown,Yes	
POST046	Post-Transplant	Post-Transplant Essential Data	Graft vs. Host Disease	yes	yes	Maximum grade of chronic GVHD (according to best clinical judgment)	Mild,Moderate,Severe,Unknown		Maximum grade of chronic GVHD (according to best clinical judgment)	Mild,Moderate,Severe,Unknown	
POST047	Post-Transplant	Post-Transplant Essential Data	Graft vs. Host Disease	yes	yes	Date of maximum grade of chronic GVHD:	YYYY/MM/DD		Date of maximum grade of chronic GVHD:	YYYY/MM/DD	
POST048	Post-Transplant	Post-Transplant Essential Data	Graft vs. Host Disease	yes	yes	Specify if chronic GVHD was limited or extensive	Extensive – One or more of the following: – Generalized skin involvement; or, – Liver histology showing chronic aggressive hepatitis, bridging necrosis or cirrhosis; or, – Involvement of eye: Schirmer's test with < 5 mm wetting; or – Involvement of minor salivary glands or oral mucosa demonstrated on labial biopsy; or – Involvement of any other target organ, Limited - Localized skin involvement and/or liver dysfunction		Specify if chronic GVHD was limited or extensive	Extensive – One or more of the following: – Generalized skin involvement; or, – Liver histology showing chronic aggressive hepatitis, bridging necrosis or cirrhosis; or, – Involvement of eye: Schirmer's test with < 5 mm wetting; or – Involvement of minor salivary glands or oral mucosa demonstrated on labial biopsy; or – Involvement of any other target organ, Limited - Localized skin involvement and/or liver dysfunction	
POST049	Post-Transplant	Post-Transplant Essential Data	Graft vs. Host Disease	yes	yes	Is the recipient still taking systemic steroids? (Do not report steroids for adrenal insufficiency, or steroid dose ≤10 mg/day for adults, <0.1 mg/kg/day for children)	No,Not Applicable,Unknown,Yes		Is the recipient still taking systemic steroids? (Do not report steroids for adrenal insufficiency, or steroid dose ≤10 mg/day for adults, <0.1 mg/kg/day for children)	No,Not Applicable,Unknown,Yes	

Item ID	Time Point	Information Collection Domain Sub-Type	Information Collection Domain Additional Sub Domain	Response required if Additional Sub Domain applies	Information Collection may be requested multiple times	Current Information Collection Data Element (if applicable)	Current Information Collection Data Element Response Option(s)	Information Collection update:	Proposed Information Collection Data Element (if applicable)	Proposed Information Collection Data Element Response Option(s)	Rationale for Information Collection Update
POST050	Post-Transplant	Post-Transplant Essential Data	Graft vs. Host Disease	yes	yes	Is the recipient still taking (non-steroid) immunosuppressive agents (including PUVA) for GVHD?	No,Not Applicable,Unknown,Yes		Is the recipient still taking (non-steroid) immunosuppressive agents (including PUVA) for GVHD?	No,Not Applicable,Unknown,Yes	
POST051	Post-Transplant	Post-Transplant Essential Data		no	yes	Was specific therapy used to prevent liver toxicity?	No,Yes		Was specific therapy used to prevent liver toxicity?	No,Yes	
POST052	Post-Transplant	Post-Transplant Essential Data		no	yes	Specify therapy (check all that apply)	Defibrotide,N-acetylcysteine,Other therapy,Tissue plasminogen activator (TPA),Ursodiol		Specify therapy (check all that apply)	Defibrotide,N-acetylcysteine,Other therapy,Tissue plasminogen activator (TPA),Ursodiol, Enoxaparin (Lovenox), Heparin	
POST053	Post-Transplant	Post-Transplant Essential Data		no	yes	Specify other therapy:	open text		Specify other therapy:	open text	
POST054	Post-Transplant	Post-Transplant Essential Data		no	yes	Did veno-occlusive disease (VOD) / sinusoidal obstruction syndrome (SOS) develop?	No,Yes		Did veno-occlusive disease (VOD) / sinusoidal obstruction syndrome (SOS) develop?	No,Yes	
POST055	Post-Transplant	Post-Transplant Essential Data		no	yes	Date of diagnosis:	YYYY/MM/DD		Date of diagnosis:	YYYY/MM/DD	
POST056	Post-Transplant	Post-Transplant Essential Data		no	yes	Did the recipient develop COVID-19 (SARS-CoV-2)?	No,Yes		Did the recipient develop COVID-19 (SARS-CoV-2)?	No,Yes	
POST057	Post-Transplant	Post-Transplant Essential Data		no	yes	Date of diagnosis:	YYYY/MM/DD		Date of diagnosis:	YYYY/MM/DD	
POST058	Post-Transplant	Post-Transplant Essential Data		no	yes	Was a vaccine for COVID-19 (SARS-CoV-2) received?	No,Unknown,Yes		Was a vaccine for COVID-19 (SARS-CoV-2) received?	No,Unknown,Yes	
POST059	Post-Transplant	Post-Transplant Essential Data	Covid-19 Vaccine	yes	yes	Specify vaccine brand	AstraZeneca,Johnson & Johnson,Moderna,Novavax,Other (specify),Pfizer-BioNTech		Specify vaccine brand	AstraZeneca,Johnson & Johnson,Moderna,Novavax,Other (specify),Pfizer-BioNTech	
POST060	Post-Transplant	Post-Transplant Essential Data	Covid-19 Vaccine	yes	yes	Specify other type:	open text		Specify other type:	open text	
POST061	Post-Transplant	Post-Transplant Essential Data	Covid-19 Vaccine	yes	yes	Select dose(s) received	Booster dose,First dose(with planned second dose) ,One dose(without planned second dose) ,Second dose,Third dose		Select dose(s) received	Booster dose,First dose(with planned second dose) ,One dose(without planned second dose) ,Second dose,Third dose	
POST062	Post-Transplant	Post-Transplant Essential Data	Covid-19 Vaccine	yes	yes	Date received:	YYYY/MM/DD		Date received:	YYYY/MM/DD	
POST063	Post-Transplant	Post-Transplant Essential Data	Covid-19 Vaccine	yes	yes	Date estimated	checked		Date estimated	checked	

Item ID	Time Point	Information Collection Domain Sub-Type	Information Collection Domain Additional Sub Domain	Response required if Additional Sub Domain applies	Information Collection may be requested multiple times	Current Information Collection Data Element (if applicable)	Current Information Collection Data Element Response Option(s)	Information Collection update:	Proposed Information Collection Data Element (if applicable)	Proposed Information Collection Data Element Response Option(s)	Rationale for Information Collection Update
POST064	Post-Transplant	Post-Transplant Essential Data		no	yes	Did a new malignancy, myelodysplastic, myeloproliferative, or lymphoproliferative disease / disorder occur that is different from the disease / disorder for which the HCT or cellular therapy was performed?	No,Yes		Did a new malignancy, myelodysplastic, myeloproliferative, or lymphoproliferative disease / disorder occur that is different from the disease / disorder for which the HCT or cellular therapy was performed?	No,Yes (Also complete Subsequent Neoplasms) , previously reported	
POST065	Post-Transplant	Post-Transplant Essential Data	Allogenic Recipients of Cord Blood units, Beta Thalassemia, and/or Sickle Cell Disease	yes	yes	Were chimerism studies performed?	no,yes		Were chimerism studies performed?	no,yes	
POST066	Post-Transplant	Post-Transplant Essential Data	Chimerism Study Performed	yes	yes	Was documentation submitted to the CIBMTR? (e.g. chimerism laboratory reports)	No,Yes		Was documentation submitted to the CIBMTR? (e.g. chimerism laboratory reports)	No,Yes	
POST067	Post-Transplant	Post-Transplant Essential Data	Chimerism Study Performed	yes	yes	Were chimerism studies assessed for more than one donor / multiple donors?	No,Yes		Were chimerism studies assessed for more than one donor / multiple donors?	No,Yes	
POST068	Post-Transplant	Post-Transplant Essential Data	Chimerism Study Performed	yes	yes	Global Registration Identifier for Donors (GRID)	open text		Global Registration Identifier for Donors (GRID)	open text	
POST069	Post-Transplant	Post-Transplant Essential Data	Chimerism Study Performed	yes	yes	NMDP cord blood unit ID:	open text		NMDP cord blood unit ID:	open text	
POST070	Post-Transplant	Post-Transplant Essential Data	Chimerism Study Performed	yes	yes	Registry donor ID:	open text		Registry donor ID:	open text	
POST071	Post-Transplant	Post-Transplant Essential Data	Chimerism Study Performed	yes	yes	Non-NMDP cord blood unit ID:	open text		Non-NMDP cord blood unit ID:	open text	
POST072	Post-Transplant	Post-Transplant Essential Data	Chimerism Study Performed	yes	yes	Date of birth:	YYYY/MM/DD		Donor Date of birth:	YYYY/MM/DD	
POST073	Post-Transplant	Post-Transplant Essential Data	Chimerism Study Performed	yes	yes	Age:	MM ___ __ (if less than 1 year); YY ___ __		Age:	MM ___ __ (if less than 1 year); YY ___ __	
POST074	Post-Transplant	Post-Transplant Essential Data	Chimerism Study Performed	yes	yes	Sex	female,male		Donor Sex	female,male	
POST075	Post-Transplant	Post-Transplant Essential Data	Chimerism Study Performed	yes	yes	Date sample collected:	YYYY/MM/DD		Date sample collected:	YYYY/MM/DD	
POST076	Post-Transplant	Post-Transplant Essential Data	Chimerism Study Performed	yes	yes	Method	Single nucleotide polymorphisms (SNPS) (includes quantitative PCR, real time PCR, sequencing, other), Fluorescent in situ hybridization (FISH) for XX/XY, Karyotyping for XX/XY,vOther, Restriction fragment-length polymorphisms (RFLP), VNTR or STR, micro or mini satellite		Method	Single nucleotide polymorphisms (SNPS) (includes quantitative PCR, real time PCR, sequencing, other), Fluorescent in situ hybridization (FISH) for XX/XY, Karyotyping for XX/XY,vOther, Restriction fragment-length polymorphisms (RFLP), VNTR or STR, micro or mini satellite	

Item ID	Time Point	Information Collection Domain Sub-Type	Information Collection Domain Additional Sub Domain	Response required if Additional Sub Domain applies	Information Collection may be requested multiple times	Current Information Collection Data Element (if applicable)	Current Information Collection Data Element Response Option(s)	Information Collection update:	Proposed Information Collection Data Element (if applicable)	Proposed Information Collection Data Element Response Option(s)	Rationale for Information Collection Update
POST077	Post-Transplant	Post-Transplant Essential Data	Chimerism Study Performed	yes	yes	Specify:	open text		Specify:	open text	
POST078	Post-Transplant	Post-Transplant Essential Data	Chimerism Study Performed	yes	yes	Cell source	Bone marrow,Peripheral blood		Cell source	Bone marrow,Peripheral blood	
POST079	Post-Transplant	Post-Transplant Essential Data	Chimerism Study Performed	yes	yes	Cell type	B-cells,Granulocytes,Hematopoietic progenitor cells,NK cells,Other,Red blood cells,T-cells,Total mononuclear cells,Unsorted / whole		Cell type	B-cells,Granulocytes,Hematopoietic progenitor cells,NK cells,Other,Red blood cells,T-cells,Total mononuclear cells,Unsorted / whole	
POST080	Post-Transplant	Post-Transplant Essential Data	Chimerism Study Performed	yes	yes	Specify:	open text		Specify:	open text	
POST081	Post-Transplant	Post-Transplant Essential Data	Chimerism Study Performed	yes	yes	Total cells examined:	open text		Total cells examined:	open text	
POST082	Post-Transplant	Post-Transplant Essential Data	Chimerism Study Performed	yes	yes	Number of donor cells:	open text		Number of donor cells:	open text	
POST083	Post-Transplant	Post-Transplant Essential Data	Chimerism Study Performed	yes	yes	Percent donor cells:	__ __ __ %		Percent donor cells:	__ __ __ %	
POST084	Post-Transplant	Disease Assessment at the Time of Best Response to HCT		no	yes	Compared to the disease status prior to the preparative regimen, what was the best response to HCT?	Continued complete remission (CCR),Complete remission (CR),Not in complete remission,Not evaluated			Continued complete remission (CCR),Complete remission (CR),Not in complete remission,Not evaluated	
POST085	Post-Transplant	Disease Assessment at the Time of Best Response to HCT		no	yes	Specify disease status if not in complete remission	Disease detected,No disease detected but incomplete evaluation to establish CR		Specify disease status if not in complete remission	Disease detected,No disease detected but incomplete evaluation to establish CR	
POST086	Post-Transplant	Disease Assessment at the Time of Best Response to HCT		no	yes	Was the date of best response previously reported?	no,yes		Was the date of best response previously reported?	no,yes	
POST087	Post-Transplant	Disease Assessment at the Time of Best Response to HCT		no	yes	Date assessed:	YYYY/MM/DD		Date assessed:	YYYY/MM/DD	
POST088	Post-Transplant	Disease Assessment at the Time of Best Response to HCT		no	yes	Was the disease status assessed by molecular testing?	No,Not Applicable,Yes		Was the disease status assessed by molecular testing?	No,Not Applicable,Yes	

Item ID	Time Point	Information Collection Domain Sub-Type	Information Collection Domain Additional Sub Domain	Response required if Additional Sub Domain applies	Information Collection may be requested multiple times	Current Information Collection Data Element (if applicable)	Current Information Collection Data Element Response Option(s)	Information Collection update:	Proposed Information Collection Data Element (if applicable)	Proposed Information Collection Data Element Response Option(s)	Rationale for Information Collection Update
POST089	Post-Transplant	Disease Assessment at the Time of Best Response to HCT		no	yes	Date assessed:	YYYY/MM/DD		Date assessed:	YYYY/MM/DD	
POST090	Post-Transplant	Disease Assessment at the Time of Best Response to HCT		no	yes	Was disease detected?	no,yes		Was disease detected?	no,yes	
POST091	Post-Transplant	Disease Assessment at the Time of Best Response to HCT		no	yes	Was the disease status assessed via flow cytometry?	No,Not Applicable,Yes		Was the disease status assessed via flow cytometry?	No,Not Applicable,Yes	
POST092	Post-Transplant	Disease Assessment at the Time of Best Response to HCT		no	yes	Date assessed:	YYYY/MM/DD		Date assessed:	YYYY/MM/DD	
POST093	Post-Transplant	Disease Assessment at the Time of Best Response to HCT		no	yes	Was disease detected?	no,yes		Was disease detected?	no,yes	
POST094	Post-Transplant	Disease Assessment at the Time of Best Response to HCT		no	yes	Was the disease status assessed by cytogenetic testing? (karyotyping or FISH)	No,Not Applicable,Yes		Was the disease status assessed by cytogenetic testing? (karyotyping or FISH)	No,Not Applicable,Yes	
POST095	Post-Transplant	Disease Assessment at the Time of Best Response to HCT		no	yes	Was the disease status assessed via FISH?	No,Not Applicable,Yes		Was the disease status assessed via FISH?	No,Not Applicable,Yes	
POST096	Post-Transplant	Disease Assessment at the Time of Best Response to HCT		no	yes	Date assessed:	YYYY/MM/DD		Date assessed:	YYYY/MM/DD	
POST097	Post-Transplant	Disease Assessment at the Time of Best Response to HCT		no	yes	Was disease detected?	no,yes		Was disease detected?	no,yes	
POST098	Post-Transplant	Disease Assessment at the Time of Best Response to HCT		no	yes	Was the disease status assessed via karyotyping?	No,Not Applicable,Yes		Was the disease status assessed via karyotyping?	No,Not Applicable,Yes	

Item ID	Time Point	Information Collection Domain Sub-Type	Information Collection Domain Additional Sub Domain	Response required if Additional Sub Domain applies	Information Collection may be requested multiple times	Current Information Collection Data Element (if applicable)	Current Information Collection Data Element Response Option(s)	Information Collection update:	Proposed Information Collection Data Element (if applicable)	Proposed Information Collection Data Element Response Option(s)	Rationale for Information Collection Update
POST099	Post-Transplant	Disease Assessment at the Time of Best Response to HCT		no	yes	Date assessed:	YYYY/MM/DD		Date assessed:	YYYY/MM/DD	
POST100	Post-Transplant	Disease Assessment at the Time of Best Response to HCT		no	yes	Was disease detected?	no,yes		Was disease detected?	no,yes	
POST101	Post-Transplant	Disease Assessment at the Time of Best Response to HCT		no	yes	Was the disease status assessed by radiological assessment? (e.g. PET, MRI, CT)	No,Not Applicable,Yes		Was the disease status assessed by radiological assessment? (e.g. PET, MRI, CT)	No,Not Applicable,Yes	
POST102	Post-Transplant	Disease Assessment at the Time of Best Response to HCT		no	yes	Date assessed:	YYYY/MM/DD		Date assessed:	YYYY/MM/DD	
POST103	Post-Transplant	Disease Assessment at the Time of Best Response to HCT		no	yes	Was disease detected?	no,yes		Was disease detected?	no,yes	
POST104	Post-Transplant	Disease Assessment at the Time of Best Response to HCT		no	yes	Was the disease status assessed by clinical / hematologic assessment?	no,yes		Was the disease status assessed by clinical / hematologic assessment?	no,yes	
POST105	Post-Transplant	Disease Assessment at the Time of Best Response to HCT		no	yes	Date assessed:	YYYY/MM/DD		Date assessed:	YYYY/MM/DD	
POST106	Post-Transplant	Disease Assessment at the Time of Best Response to HCT		no	yes	Was disease detected?	no,yes		Was disease detected?	no,yes	
POST107	Post-Transplant	Post-HCT Therapy		no	yes	Was therapy given for reasons other than relapse, persistent, or progressive disease? (Include any maintenance and consolidation therapy.)	no,yes		Was therapy given for reasons other than relapse, persistent, or progressive disease? (Include any maintenance and consolidation therapy.)	no,yes	
POST108	Post-Transplant	Post-HCT Therapy		no	yes	Specify therapy (check all that apply)	Blinded randomized trial,Cellular therapy,Other therapy,Radiation,Systemic therapy		Specify therapy (check all that apply)	Blinded randomized trial,Cellular therapy,Other therapy,Radiation,Systemic therapy	

Item ID	Time Point	Information Collection Domain Sub-Type	Information Collection Domain Additional Sub Domain	Response required if Additional Sub Domain applies	Information Collection may be requested multiple times	Current Information Collection Data Element (if applicable)	Current Information Collection Data Element Response Option(s)	Information Collection update:	Proposed Information Collection Data Element (if applicable)	Proposed Information Collection Data Element Response Option(s)	Rationale for Information Collection Update
POST109	Post-Transplant	Post-HCT Therapy		no	yes	Specify systemic therapy (check all that apply)	Alemtuzumab,Azacytidine,Blinatumoma b,Bortezomib,Bosutinib,Carfilzomib,Chemotherapy,Dasatinib,Decitabine,Gemtuzumab,Gilteritinib,Ibrutinib,Imatinib mesylate,Ixazomib,Lenalidomide,Lestaurtinib,Midostaurin,Nilotinib,Nivolumab,Other systemic therapy,Pembrolizumab,Pomalidomide,Quizartinib,Rituximab,Sorafenib,Sunitinib,Thalidomide		Specify systemic therapy (check all that apply)	Alemtuzumab,Azacytidine,Blinatumomab,Bortezomib,Bosutinib,Carfilzomib,Dasatinib,Decitabine,Gemtuzumab,Gilteritinib,Ibrutinib,Imatinib mesylate,Ixazomib,Lenalidomide,Lestaurtinib,Midostaurin,Nilotinib,Nivolumab,Other systemic therapy,Pembrolizumab,Pomalidomide,Quizartinib,Rituximab,Sorafenib,Sunitinib,Thalidomide, Brentuximab vendotin, Daratumumab (Darzalex)	
POST110	Post-Transplant	Post-HCT Therapy		no	yes	Specify other systemic therapy:	open text		Specify other systemic therapy:	open text	
POST111	Post-Transplant	Post-HCT Therapy		no	yes	Specify other therapy:	open text		Specify other therapy:	open text	
POST112	Post-Transplant	Post-HCT Therapy		no	yes	Did a fecal microbiota transplant (FMT) occur?	No, Yes		Did a fecal microbiota transplant (FMT) occur?	No, Yes	
POST113	Post-Transplant	Post-HCT Therapy		no	yes				Date of FMT	DD/MM/YY	
POST114	Post-Transplant	Post-HCT Therapy		no	yes				Specify the indication for the FMT	Graft versus host disease (GVHD), Clostridium difficile, Other	
POST115	Post-Transplant	Post-HCT Therapy		no	yes				Specify other indication:	open text	
POST116	Post-Transplant	Relapse or Progression Post-HCT		no	yes	Did the recipient experience a clinical/hematologic relapse or progression post-HCT?	No,Yes		Did the recipient experience a clinical/hematologic relapse or progression post-HCT?	No,Yes	
POST117	Post-Transplant	Relapse or Progression Post-HCT		no	yes	Was the date of the first clinical / hematologic relapse or progression previously reported?	No,Yes (only valid >day 100)		Was the date of the first clinical / hematologic relapse or progression previously reported?	No,Yes (only valid >day 100)	
POST118	Post-Transplant	Relapse or Progression Post-HCT		no	yes	Date first seen:	YYYY/MM/DD		Date first seen:	YYYY/MM/DD	
POST119	Post-Transplant	Relapse or Progression Post-HCT		no	yes	Was intervention given for relapsed, persistent or progressive disease?	No,Yes		Was intervention given for relapsed, persistent or progressive disease?	No,Yes	
POST120	Post-Transplant	Relapse or Progression Post-HCT		no	yes	Specify reason for which intervention was given	Persistent disease,Relapsed / progressive disease		Specify reason for which intervention was given	Persistent disease,Relapsed / progressive disease	
POST121	Post-Transplant	Relapse or Progression Post-HCT		no	yes	Specify the method(s) of detection for which intervention was given (check all that apply)	Clinical and/or hematologic analysis,Cytogenetic Analysis,Disease specific molecular marker,Flow Cytometry,Radiological		Specify the method(s) of detection for which intervention was given (check all that apply)	Clinical and/or hematologic analysis,Cytogenetic Analysis,Disease specific molecular marker,Flow Cytometry,Radiological	
POST122	Post-Transplant	Relapse or Progression Post-HCT		no	yes	Date intervention started:	YYYY/MM/DD		Date intervention started:	YYYY/MM/DD	
POST123	Post-Transplant	Relapse or Progression Post-HCT		no	yes	Specify therapy (check all that apply)	Blinded randomized trial,Cellular therapy,Other therapy,Radiation,Systemic therapy		Specify therapy (check all that apply)	Blinded randomized trial,Cellular therapy,Other therapy,Radiation,Systemic therapy	
POST124	Post-Transplant	Relapse or Progression Post-HCT		no	yes	Specify systemic therapy (check all that apply)	Alemtuzumab,Azacytidine,Blinatumoma b,Bortezomib,Bosutinib,Carfilzomib,Chemotherapy,Dasatinib,Decitabine,Gemtuzumab,Gilteritinib,Ibrutinib,Imatinib mesylate,Ixazomib,Lenalidomide,Lestaurtinib,Midostaurin,Nilotinib,Nivolumab,Other systemic therapy,Pembrolizumab,Pomalidomide,Quizartinib,Rituximab,Sorafenib,Sunitinib,Thalidomide		Specify systemic therapy (check all that apply)	Alemtuzumab,Azacytidine,Blinatumomab,Bortezomib,Bosutinib,Carfilzomib,Chemotherapy,Dasatinib,Decitabine,Gemtuzumab,Gilteritinib,Ibrutinib,Imatinib mesylate,Ixazomib,Lenalidomide,Lestaurtinib,Midostaurin,Nilotinib,Nivolumab,Other systemic therapy,Pembrolizumab,Pomalidomide,Quizartinib,Rituximab,Sorafenib,Sunitinib,Thalidomide, Daratumumb (Darzalex), Venetoclax	

Item ID	Time Point	Information Collection Domain Sub-Type	Information Collection Domain Additional Sub Domain	Response required if Additional Sub Domain applies	Information Collection may be requested multiple times	Current Information Collection Data Element (if applicable)	Current Information Collection Data Element Response Option(s)	Information Collection update:	Proposed Information Collection Data Element (if applicable)	Proposed Information Collection Data Element Response Option(s)	Rationale for Information Collection Update
POST125	Post-Transplant	Relapse or Progression Post-HCT		no	yes	Specify other systemic therapy:	open text		Specify other systemic therapy:	open text	
POST126	Post-Transplant	Relapse or Progression Post-HCT		no	yes	Specify other therapy:	open text		Specify other therapy:	open text	
POST127	Post-Transplant	Current Disease Status		no	yes	What is the current disease status?	Complete remission (CR),Not in complete remission,Not evaluated		What is the current disease status?	Complete remission (CR),Not in complete remission,Not evaluated	
POST128	Post-Transplant	Current Disease Status		no	yes	Specify disease status if not in complete remission	Disease detected,No disease detected but incomplete evaluation to establish CR		Specify disease status if not in complete remission	Disease detected,No disease detected but incomplete evaluation to establish CR	
POST129	Post-Transplant	Current Disease Status		no	yes	Date of most recent disease assessment:	YYYY/MM/DD		Date of -assessment of current disease status	YYYY/MM/DD	
POST130	Post-Transplant	Recipient Death Data	Recipient Death	yes	no				Date of death:	YYYY/MM/DD	
POST131	Post-Transplant	Recipient Death Data	Recipient Death	yes	no				Date estimated	checked	
POST132	Post-Transplant	Recipient Death Data	Recipient Death	yes	no				Was cause of death confirmed by autopsy?	Autopsy pending,No,Unknown,Yes	
POST133	Post-Transplant	Recipient Death Data	Recipient Death	yes	no				Was documentation submitted to the CIBMTR?	No,Yes	
POST134	Post-Transplant	Recipient Death Data	Recipient Death	yes	no	Primary cause of death	Accidental death,Acute GVHD,Adult respiratory distress syndrome (ARDS) (other than IPS),Bacterial infection,Cardiac failure,Chronic GVHD,Central nervous system (CNS) failure,COVID-19 (SARS-CoV-2),Cytokine release syndrome,Diffuse alveolar damage (without hemorrhage), Disseminated intravascular coagulation (DIC),Fungal infection, Gastrointestinal (GI) failure (not liver),Graft rejection or failure, Thrombotic microangiopathy (TMA) (Thrombotic thrombocytopenic purpura (TTP)/Hemolytic Uremic Syndrome (HUS)),Idiopathic pneumonia syndrome (IPS), Liver failure (not VOD),Multiple organ failure,New malignancy,Infection, organism not identified,Other cause, Other infection,Other organ failure,Other pulmonary syndrome (excluding pulmonary hemorrhage),Other vascular,Prior malignancy,Protozoal infection, Pulmonary failure,Recurrence / persistence / progression of disease,Renal failure,Suicide,Thromboembolic, Pneumonitis due to Cytomegalovirus (CMV),Viral infection,Pneumonitis due to other virus,Veno-occlusive disease (VOD) / sinusoidal obstruction syndrome (SOS)		Primary cause of death	Accidental death,Acute GVHD,Adult respiratory distress syndrome (ARDS) (other than IPS),Bacterial infection,Cardiac failure,Chronic GVHD,Central nervous system (CNS) failure,COVID-19 (SARS-CoV-2),Cytokine release syndrome,Diffuse alveolar damage (without hemorrhage),Disseminated intravascular coagulation (DIC),Fungal infection,Gastrointestinal hemorrhage,Gastrointestinal (GI) failure (not liver),Graft rejection or failure,Hemorrhagic cystitis,Thrombotic microangiopathy (TMA) (Thrombotic thrombocytopenic purpura (TTP)/Hemolytic Uremic Syndrome (HUS)),Idiopathic pneumonia syndrome (IPS),Intracranial hemorrhage,Liver failure (not VOD),Multiple organ failure,New malignancy,Infection, organism not identified,Other cause,Other hemorrhage neurotoxicity (ICANS), Other infection,Other organ failure,Other pulmonary syndrome (excluding pulmonary hemorrhage),Other vascular,Prior malignancy,Protozoal infection,Pulmonary hemorrhage,Pulmonary failure,Recurrence / persistence / progression of disease,Renal failure,Suicide,Thromboembolic, Tumor lysis syndrome, Pneumonitis due to Cytomegalovirus (CMV),Viral infection,Pneumonitis due to other virus,Veno-occlusive disease (VOD) / sinusoidal obstruction syndrome (SOS)	
POST135	Post-Transplant	Recipient Death Data	Recipient Death	yes	no	Specify:	open text		Specify:	open text	

Item ID	Time Point	Information Collection Domain Sub-Type	Information Collection Domain Additional Sub Domain	Response required if Additional Sub Domain applies	Information Collection may be requested multiple times	Current Information Collection Data Element (if applicable)	Current Information Collection Data Element Response Option(s)	Information Collection update:	Proposed Information Collection Data Element (if applicable)	Proposed Information Collection Data Element Response Option(s)	Rationale for Information Collection Update
POST136	Post-Transplant	Recipient Death Data	Recipient Death	yes	no	Contributing cause of death	Accidental death,Acute GVHD,Adult respiratory distress syndrome (ARDS) (other than IPS),Bacterial infection,Cardiac failure,Chronic GVHD,Central nervous system (CNS) failure,COVID-19 (SARS-CoV-2),Cytokine release syndrome,Diffuse alveolar damage (without hemorrhage), Disseminated intravascular coagulation (DIC),Fungal infection, Gastrointestinal (GI) failure (not liver),Graft rejection or failure, Thrombotic microangiopathy (TMA) (Thrombotic thrombocytopenic purpura (TTP)/Hemolytic Uremic Syndrome (HUS)),Idiopathic pneumonia syndrome (IPS), Liver failure (not VOD),Multiple organ failure,New malignancy,Infection, organism not identified,Other cause, Other infection,Other organ failure,Other pulmonary syndrome (excluding pulmonary hemorrhage),Other vascular,Prior malignancy,Protozoal infection, Pulmonary failure,Recurrence / persistence / progression of disease,Renal failure,Suicide,Thromboembolic, Pneumonitis due to Cytomegalovirus (CMV),Viral infection,Pneumonitis due to other virus,Veno-occlusive disease (VOD) / sinusoidal obstruction syndrome (SOS)		Contributing cause of death	Accidental death,Acute GVHD,Adult respiratory distress syndrome (ARDS) (other than IPS),Bacterial infection,Cardiac failure,Chronic GVHD,Central nervous system (CNS) failure,COVID-19 (SARS-CoV-2),Cytokine release syndrome,Diffuse alveolar damage (without hemorrhage),Diffuse alveolar hemorrhage (DAH),Disseminated intravascular coagulation (DIC),Fungal infection,Gastrointestinal hemorrhage,Gastrointestinal (GI) failure (not liver),Graft rejection or failure,Hemorrhagic cystitis,Thrombotic microangiopathy (TMA) (Thrombotic thrombocytopenic purpura (TTP)/Hemolytic Uremic Syndrome (HUS)),Idiopathic pneumonia syndrome (IPS),Intracranial hemorrhage,Liver failure (not VOD),Multiple organ failure,New malignancy,Infection, organism not identified,Other cause,Other hemorrhage neurotoxicity (ICANS), Other infection,Other organ failure,Other pulmonary syndrome (excluding pulmonary hemorrhage),Other vascular,Prior malignancy,Protozoal infection,Pulmonary hemorrhage,Pulmonary failure,Recurrence / persistence / progression of disease,Renal failure,Suicide,Thromboembolic, Tumor lysis syndrome, Pneumonitis due to Cytomegalovirus (CMV),Viral infection,Pneumonitis due to other virus,Veno-occlusive disease (VOD) / sinusoidal obstruction syndrome (SOS)	
POST137	Post-Transplant	Recipient Death Data	Recipient Death	yes	no	Specify:	open text		Specify:	open text	
POST138	Post-Transplant	Subsequent Neoplasms	New Malignancy, Lymphoproliferative or Myeloproliferative Disease / Disorder	yes	yes	Specify the new malignancy	Hematologic Malignancy: Acute myeloid leukemia (AML / ANLL), Other leukemia, Myelodysplastic syndrome (MDS), Myeloproliferative neoplasm (MPN), Overlapping myelodysplasia / myeloproliferative neoplasm (MDS / MPN), Hodgkin lymphoma, Non-Hodgkin lymphoma, Clonal cytogenetic abnormality without leukemia or MDS, Uncontrolled proliferation of donor cells without malignant transformation Solid Tumors: Oropharyngeal cancer (e.g. tongue, mouth, throat), Gastrointestinal malignancy (e.g. esophagus, stomach, small intestine, colon, rectum, anus, liver, pancreas), Lung cancer, Melanoma, Squamous cell skin malignancy, Basal cell skin malignancy, Breast cancer, Genitourinary malignancy (e.g. kidney, bladder, cervix, uterus, ovary, prostate, testis), Central nervous system (CNS) malignancy (e.g. meningioma, glioma), Thyroid cancer		Specify the new malignancy	Hematologic Malignancy: Acute myeloid leukemia (AML / ANLL), Acute lymphoblastic leukemia (ALL), Other leukemia, Myelodysplastic syndrome (MDS), Myeloproliferative neoplasm (MPN), Overlapping myelodysplasia / myeloproliferative neoplasm (MDS / MPN), Hodgkin lymphoma, Non-Hodgkin lymphoma, Multiple myeloma / plasma cell neoplasms, Clonal cytogenetic abnormality without leukemia or MDS, Uncontrolled proliferation of donor cells without malignant transformation. Solid Tumors: Bone sarcoma (regardless of site), Soft tissue sarcoma (regardless of site), Oropharyngeal cancer (e.g. tongue, mouth, throat), Gastrointestinal malignancy (e.g. esophagus, stomach, small intestine, colon, rectum, anus, liver, pancreas), Lung cancer, Melanoma, Squamous cell skin malignancy, Basal cell skin malignancy, Breast cancer, Genitourinary malignancy (e.g. kidney, bladder, cervix, uterus, ovary, prostate, testis), Central nervous system (CNS) malignancy (e.g. meningioma, glioma), Thyroid cancer	
POST139	Post-Transplant	Subsequent Neoplasms	New Malignancy, Lymphoproliferative or Myeloproliferative Disease / Disorder	yes	yes				Was post-transplant lymphoproliferative disorder (PTLD) diagnosed?	No,Yes	

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POST140	Post-Transplant	Subsequent Neoplasms	New Malignancy, Lymphoproliferative or Myeloproliferative Disease / Disorder	yes	yes				Specify type of PTLD	Monomorphic,Polymorphic,Unknown	
POST141	Post-Transplant	Subsequent Neoplasms	New Malignancy, Lymphoproliferative or Myeloproliferative Disease / Disorder	yes	yes				Specify oropharyngeal cancer	Mouth,Throat,Tongue, Other oropharyngeal cancer	
POST142	Post-Transplant	Subsequent Neoplasms	New Malignancy, Lymphoproliferative or Myeloproliferative Disease / Disorder	yes	yes				Specify gastrointestinal malignancy	Anus,Colon,Esophagus,Liver ,Pancreas,Rectum,Small Intestine (DUODENUM, JEJUNUM, ILEUM),Stomach, Other gastrointestinal cancer	
POST143	Post-Transplant	Subsequent Neoplasms	New Malignancy, Lymphoproliferative or Myeloproliferative Disease / Disorder	yes	yes				Specify genitourinary malignancy	Bladder,Cervix,Kidney,Ovary,Prostate,Testicle,Uterus, Other genitourary malignancy	
POST144	Post-Transplant	Subsequent Neoplasms	New Malignancy, Lymphoproliferative or Myeloproliferative Disease / Disorder	yes	yes				Specify CNS malignancy	Glioma,Meningioma,Other CNS malignancy	
POST145	Post-Transplant	Subsequent Neoplasms	New Malignancy, Lymphoproliferative or Myeloproliferative Disease / Disorder	yes	yes	Specify other new malignancy:	open text		Specify other new malignancy:	open text	
POST146	Post-Transplant	Subsequent Neoplasms	New Malignancy, Lymphoproliferative or Myeloproliferative Disease / Disorder	yes	yes	Date of diagnosis:	YYYY/MM/DD		Date of diagnosis:	YYYY/MM/DD	
POST147	Post-Transplant	Subsequent Neoplasms	New Malignancy, Lymphoproliferative or Myeloproliferative Disease / Disorder	yes	yes	Was documentation submitted to the CIBMTR?	No,Yes		Was documentation submitted to the CIBMTR?	No,Yes	
POST148	Post-Transplant	Subsequent Neoplasms	New Malignancy, Lymphoproliferative or Myeloproliferative Disease / Disorder	yes	yes	Was the new malignancy donor / cell product derived?	No,Not Done,Yes		Was the new malignancy donor / cell product derived?	No,Not Done,Yes	
POST149	Post-Transplant	Subsequent Neoplasms	New Malignancy, Lymphoproliferative or Myeloproliferative Disease / Disorder	yes	yes	Was documentation submitted to the CIBMTR?	no,yes		Was documentation submitted to the CIBMTR?	no,yes	

Item ID	Time Point	Information Collection Domain Sub-Type	Information Collection Domain Additional Sub Domain	Response required if Additional Sub Domain applies	Information Collection may be requested multiple times	Current Information Collection Data Element (if applicable)	Current Information Collection Data Element Response Option(s)	Information Collection update:	Proposed Information Collection Data Element (if applicable)	Proposed Information Collection Data Element Response Option(s)	Rationale for Information Collection Update
POST150	Post-Transplant	Subsequent Neoplasms	New Malignancy, Lymphoproliferative or Myeloproliferative Disease / Disorder	yes	yes				Was PTLD confirmed by biopsy?	No,Yes	
POST151	Post-Transplant	Subsequent Neoplasms	New Malignancy, Lymphoproliferative or Myeloproliferative Disease / Disorder	yes	yes	Was the pathology of the tumor EBV positive?	no,yes		Was the pathology of the tumor EBV positive?	no,yes	
POST152	Post-Transplant	Subsequent Neoplasms	New Malignancy, Lymphoproliferative or Myeloproliferative Disease / Disorder	yes	yes				Was documentation submitted to the CIBMTR? (e.g. pathology report)	No,Yes	
POST153	Post-Transplant	Subsequent Neoplasms	New Malignancy, Lymphoproliferative or Myeloproliferative Disease / Disorder	yes	yes				Was there EBV reactivation in the blood?	No,Not Done,Yes	
POST154	Post-Transplant	Subsequent Neoplasms	New Malignancy, Lymphoproliferative or Myeloproliferative Disease / Disorder	yes	yes				How was EBV reactivation diagnosed?	Other method,Qualitative PCR of blood,Quantitative PCR of blood	
POST155	Post-Transplant	Subsequent Neoplasms	New Malignancy, Lymphoproliferative or Myeloproliferative Disease / Disorder	yes	yes				Specify other method:	open text	
POST156	Post-Transplant	Subsequent Neoplasms	New Malignancy, Lymphoproliferative or Myeloproliferative Disease / Disorder	yes	yes				Quantitative EBV viral load of blood: At diagnosis	_____copies/ml	
POST157	Post-Transplant	Subsequent Neoplasms	New Malignancy, Lymphoproliferative or Myeloproliferative Disease / Disorder	yes	yes				Was a quantitative PCR of blood performed again after diagnosis?	No,Yes	
POST158	Post-Transplant	Subsequent Neoplasms	New Malignancy, Lymphoproliferative or Myeloproliferative Disease / Disorder	yes	yes				Highest EBV viral load of blood:	_____copies/ml	
POST159	Post-Transplant	Subsequent Neoplasms	New Malignancy, Lymphoproliferative or Myeloproliferative Disease / Disorder	yes	yes				Was there lymphomatous involvement?	No,Yes	

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POST160	Post-Transplant	Subsequent Neoplasms	New Malignancy, Lymphoproliferative or Myeloproliferative Disease / Disorder	yes	yes				Specify sites of PTLD involvement (check all that apply)	Bone marrow,Central nervous system (brain or cerebrospinal fluid),Liver,Lung,Lymph node(s),Other,Spleen	
POST161	Post-Transplant	Subsequent Neoplasms	New Malignancy, Lymphoproliferative or Myeloproliferative Disease / Disorder	yes	yes				Specify other site:	open text	
POST162	Post-Transplant	Subsequent Neoplasms		no	yes	First Name (person completing form):	open text		First Name (person completing form):	open text	
POST163	Post-Transplant	Subsequent Neoplasms		no	yes	Last Name:	open text		Last Name:	open text	
POST164	Post-Transplant	Subsequent Neoplasms		no	yes	E-mail address:	open text		E-mail address:	open text	
POST165	Post-Transplant	Subsequent Neoplasms		no	yes	Date:	YYYY/MM/DD		Date:	YYYY/MM/DD	

Below are pull down options for Column I: Do not delete

Addition of Information Requested

Deletion of Information Requested

Merged to Check all that Apply

Change/Clarification of Information Requested and Response Option

Change/Clarification of Information Requested

Change/Clarification of Response Options

Information Collection Domain Sub-Type will change to Lab

Below are pull down options for Column L: Do not delete

Reduce burden: expanded response options to include responses previously reported manually or created a "check all that apply"

Be consistent with current clinical landscape, improve transplant outcome data
Capture data accurately

Examples added or typographical/grammatical errors corrected for clarification

Covid-19 Impact

Capture additional relevant disease information

Reduce redundancy in data capture

Instruction text (since date of last report) for the individual question is extraneous as it is applied in the overall instructions for questions at this time point of data collection

Instruction text change to remove navigation instructions