CIC:	Hospital UPN:	Patient UIC	HSCT Date: yyyy - mm - dd
	HSCT - Min	imum Essential I	
		Centre Identification	
	Unit:	<del></del>	
		Patient Data	
	yyyy - mm - dd tional / international study / Name of study / trial		Yes □No Jnknown
Compulsory, registration All transplants perform the patient and not to	the transplant.	it this item. e registered with the same patient identific	ation number or code as this belongs to
	(first name(s) _	_family name(s))	
Date of birth:	yyyy - mm - dd	Sex: Male (at birth)	☐ Female
	Prir	mary Disease Diagnosis	
	osis:yyyy - mm - dd  AGNOSIS (CHECK THE DISEAS	SE FOR WHICH THIS TRANSPLANT WAS PERF	ORMED)
related Prec  Precursor Ly  Therapy related Secondary Acu  Chronic Leukae  Chronic Mye	ogenous Leukaemia (AML) ursor Neoplasms mphoid Neoplasms (old ALL) d myeloid neoplasms (old te Leukaemia) emia eloid Leukaemia (CML) phocytic Leukaemia (CLL) n sease	<ul> <li>Myeloma/Plasma cell disorder</li> <li>Solid Tumour</li> <li>Myelodysplastic syndromes / Myeloproliferative neoplasm</li> <li>MDS</li> <li>MDS/MPN</li> <li>Myeloproliferative neoplasm</li> <li>Bone marrow failure including Aplastic anaemia</li> <li>Inherited disorders</li> <li>Primary immune deficiencies</li> <li>Metabolic disorders</li> </ul>	<ul> <li>☐ Histiocytic disorders</li> <li>☐ Autoimmune disease</li> <li>☐ Juvenile Idiopathic Arthritis</li> <li>☐ Multiple Sclerosis</li> <li>☐ Systemic Lupus</li> <li>☐ Systemic Sclerosis</li> <li>☐ Haemoglobinopathy</li> </ul>

CIC: Hospital UPN: Patient UIC	HSCT Date:
COMBINED MYELODYPLASTIC SYNDROME/ MYELOPI	ROLIFERATIVE NEOPLASM
(MDS/MPN) (main disease of	code 6)
Disease	
Date of initial diagnosisyyyy - mm - dd	
Classification:  Chronic myelomonocytic leukaemia (CMMoL, CMML)  Juvenile myelomonocytic leukaemia (JCMMoL, JMML, JCML, JCMML)  Atypical CML ((t(9;22) negative and BCR-ABL1 negative)	
Therapy related MDS/ MPN:  (Secondary origin)  Yes: Disease related to prior exposure to thera  No  Unknown	peutic drugs or radiation
Chromosome Analysis at Dia	agnosis
Chromosome analysis at diagnosis (All methods including FISH)	
Abnormal Normal Not done or failed	Unknown
If abnormal:	
Complex kariotype:	
You can transcribe the complete karyotype:	
OR	
Indicate below those abnormalities that have been <b>evaluated</b> and whether they w	vere Absentor Present
Abn 1, specify	☐ Absent ☐ Present ☐ Not evaluated
Abn 5, specify	Absent Present Not evaluated
Abn 7, specify	Absent Present Not evaluated
trisomy 8	Absent Present Not evaluated
trisomy 9	☐ Absent ☐ Present ☐ Not evaluated
Del 20	☐ Absent ☐ Present ☐ Not evaluated
Del 13	☐ Absent ☐ Present ☐ Not evaluated
Other, specify	Absent Present Not evaluated
	1
Molecular Markers at Diag	nosis
☐ Not evaluated ☐ Evaluated: Absent ☐ Evaluated: Present ☐ U	Inknown
Indicate below those abnormalities that have been <b>evaluated</b> and whether they v	were Absentor Present
BCR-ABL; molecular product of t(9;22)(q34;q11.2)	Absent Present Not evaluated
JAK2 mutation	Absent Present Not evaluated
FIP1L1-PDGFR	Absent Present Not evaluated
PTPN-11	Absent Present Not evaluated
K-RAS	Absent Present Not evaluated
N-RAS	Absent Present Not evaluated
CBL	Absent Present Not evaluated
Other	Absent Present Not evaluated

CIC:	Hospital UPN:	Patient LIIC		HSCT Date:	
CIC	1105pttal 01 14.			niser bate.	yyyy - mm - dd
COMBINED MYELO	ODYPLASTIC SYNDR (MDS/MPN	ROME/ MYELOF ) (main disease		TIVE NEO	PLASM
	S	tatus at HSCT			
Date of this HSCT:	yyyy - mm - dd				
WHO Classification at HS	ст:				
Juvenile myelomonocytic	c leukaemia (CMMoL, CMML) c leukaemia (JCMMoL, JMML, JCML egative and BCR-ABL1 negative)	., JCMML)			
STATUS					
CMML/ Atypical CML					
STATUS			NUMBER		
Treated with chemotherapy:					
Primary refractory phase	(no change)				
Complete remission (CR)			1st 2nd 3rd or high	er	
☐ Improvement but no CR					
			1st		
Relapse (after CR)			2nd 3rd or highe	er	
Relapse (after CR) Progression/worse				er	
Progression/worse	e care or treatment without chemo	therapy)		er	

CIC: H	ospital UPN:	Patient UIC	HSCT Date:	уууу -	mm - d	'd
		HSCT				
Performance score  Score 10 U  Weight (kg):	system used	nsky □ 50 □ 60 □	70 🗆 80 🗆 90	□ 100	)	
	Con	norbidity Index				
forror et al., Blood, 2005 Oct 1		<u> </u>	pmc/articles/PMC1895304/			
Vas there any <i>clinically signific</i> preparative regimen? ☐ No ☐ Yes	cant co-existing disease or or	gan impairment at time c	of patient assessment just prior	to the		
Comorbidity		Definitions		No	Yes	N/E
Solid tumour, previously present	Treated at any time point melanoma skin cancer Indicate type		ory, excluding non-			
nfammatory bowel disease	Crohn's disease or ulcera					
Rheumatologic	SLE, RA, polymyositis, m	ixed CTD, or polymyalgia	rheumatica			
nfection	Requiring continuation of	of antimicrobial treatmen	t after day 0			
Diabetes	Requiring treatment wit diet alone	h insulin or oral hypoglyc	aemics but not			
Renal: moderate/severe	Serum creatinine > 2 mg transplantation	/dL or >177 μmol/L, on d	ialysis, or prior renal			
Hepatic: mild moderate/ severe	ULN, or AST/ALT betwee	en ULN and 2.5 × ULN	Normal (ULN) and 1.5 x the r AST/ALT greater than 2.5			
Arrhythmia		er, sick sinus syndrome, o	r ventricular			
Cardiac	Coronary artery disease, 50%, or shortening fract	=	myocardial infarction, EF ≤			
Cerebrovascular disease	Transient ischemic attac	k or cerebrovascular acci	dent			
Heart valve disease	Except mitral valve prola	apse				
Pulmonary: moderate	DLco and/or FEV1 66-80	% or dyspnoea on slight a	activity			
severe	DLco and/or FEV1 ≤ 65%	or dyspnoea at rest or re	equiring oxygen			
Obesity	Patients with a body ma	ss index > 35 kg/m2				
Peptic ulcer	Requiring treatment					
Psychiatric disturbance	Depression or anxiety re	equiring psychiatric consu	Itation or treatment			

Were there any other major clinical abnormalities prior to the preparative regimen? Specify.....

CIC:	Hospital UPN:	Patient UIC	HSCT	Date:
				yyyy - mm - dd
	Туре	of HSCT (Alloge	eneic)	
☐ Allogeneic				
Patient CMV status	☐ Negative	Positive Not eva	luated Unknow	/n
Multiple donors (including multiple CB	units) No	Yes: Number of donors		
		Donor 1		
HLA MATCH TYPE (DONOR  HLA - Identical sibling (I) Syngeneic (monozygotic HLA - Matched other re HLA - Mismatched related	may include non-monozygot c twin) elative	f mismatch 📗 1 HLA loco	us mismatch oci mismatch	
Donor ID given by th	ne centre			
<b>HLA</b> MISMATCHES BET' (Mismatched relatives only)	WEEN DONOR AND PATIENT			
Complete number	of mismatches inside each b	ох		
A B	C DRB1 DQB1 D	PB1		
0=match; 1=one mismatch; 2	2=2 mismatches; N/E=not evalua	Antigenic  Allelic		
Unrelated donor				
ION code of the Donor Regist	,			
BMDW code of the Donor Re		I code is unknown) (up to 4 ch	naracters)	
Name of Donor Registry/ CB	., ,			
Donor centre na	(1) applicable) options	al) y or the CB Bank listed above		
		ry or the CB Bank listed above		
		TS WITH HLA TYPING into the		
Donor information	ner the Endomnon Negot		adtabase	
Date of birth		OR Age at time of donation	(if date of birth not p	
Donor Sex	(at birth)	Female		Tur(3)
Donor CMV sta	tus Negative	☐ Positive	☐ Not evaluated	Unknown
Did this donor provide more tha	an one stem cell product	_	_	_
No - (pleas	se fill "Donor 1 – Product I of different stem cell produc	• =	AND 2" on next page)	

CIC:	Hospital UPN:	Patient UIC	HSCT Date:	yyyy - mm - dd
	Dono	r 1 - Product Number	r 1	
If more than one st	em cell product, this is the FIRST pro	duct infused from this donor		
Source of Stem Ce		eral blood		
Cord blood  Graft manipulation	Other:n ex-vivo of this product including T-			
other than for RBC	Cremoval or volume reduction			
Yes	Negative:	S:    T-cell (CD3+) depletion (do no     T-cell receptor αβ depletion     B-cell depletion (CD19+) by M   NK cell depletion by MoAB     Other	ІоАВ	<u></u>
	Positive: No Yes	CD34+ enrichment		
	Genetic manipulation	☐ No ☐ Yes		
	Donc	or 1 - Product Numbe	er 2	
If more than one st	em cell product, this is the SECOND		<del>-</del>	
Source of Stem Ce		product infused from this donor		
☐ Bone marro ☐ Cord blood		eral blood		
Cord blood	w Periph	eral blood cell depletion	ІоАВ	
Cord blood  Graft manipulation  other than for RBC	w Periph Other: n ex-vivo of this product including T- cremoval or volume reduction	reral blood  cell depletion  ss:  T-cell (CD3+) depletion (do no  T-cell receptor αβ depletion  B-cell depletion (CD19+) by M	ІоАВ	

Please enter the LABORATORY RESULTS WITH HLA TYPING into the database

CIC:	Hospital UPN:	Pa	atient UIC	HSCT Date:
		D	onor 2	,,,,
HLA MATCH TYPE (DON	NOR RELATION WITH PATIEN	NT)		
☐ HLA - Identi	ical sibling (may includ	de non-monozygotio	twin)	
Syngeneic	(monozygotic twin)			
HLA - Matcl	hed other relative			
HLA - Mism	atched relative Degre	e of mismatch	☐ 1 HLA locus misma ☐ >=2 HLA loci misma	
<b>HLA</b> MISMATCHES E (Mismatched relatives on	BETWEEN DONOR AND PATI hly)	IENT		
Complete numb	er of mismatches inside ea	ch box		
A B	C DRB1 DQB1	L DPB1		
		Antina		
		Antige	erric	
	$oldsymbol{\sqcup} oldsymbol{\sqcup} oldsymbol{\sqcup} oldsymbol{\sqcup}$	Allelic		
0=match; 1=one mismatc	ch; 2=2 mismatches; N/E=not ev	valuated		
Unrelated dor	nor			
ION code of the Don	or Registry or CB Bank			
				cters)
Name of Donor Regis			unknown)	
Donor ce	ntre name (if applicable	e, optional)		
<b>Donor</b> ID	given by the Donor Registry	y or the CB Bank list	ed above	
Patient	D given by the Donor Regist	try or the CB Bank li	sted above	
>p	lease enter the LABORATO	RY RESULTS WITH	HLA TYPING into the dat	abase
,				
Donor information				
Date of birth	yyyy - mm - dd	<u>OR</u> Ag		(if date of birth not provided)
Donor Sex (at	birth)	Female	yeai	r(s)month(s)
Donor CMV status	Negative	Positive	☐ Not evaluated	Unknown
Did this donor provide m	nore than one stem cell pro	duct		
•	<i>please fill "Donor 1 – Pro</i> umber of different stem cell		· · · · · · · · · · · · · · · · · · ·	
	f 2 products e.g. BM PB, ple			?" on next page)

If more than one stem cell product, this is the FIRST product Infused from this donor	CIC:	Hospital UPN:	Patient UIC	HSCT Date:	уууу - mm - dd
If more than one stem cell product, this is the FIRST product infused from this donor    Source of Stem Cells for this product, select only one   Bone marrow   Peripheral blood   Cord blood   Other source   Graft manipulation ex-vivo including T-Cell depletion other thon for RBC removal or volume reduction   No   Yes   Negative:   No   Yes   CD34+ enrichment   Reduction   B. cell depletion (CD39+) by MoAB   No. cell depletion (CD39+) depletion (CD39+) depletion (CD39+) depletion (CD39+) depletion (CD39+) by MoAB   No. cell		Donor	2 - Product Numb	ar 1	
Source of Stem Cells for this product, select only one   Bone marrow				OI I	
Bone marrow   Peripheral blood   Cord blood   Other source			uct infused from this donor		
Cord blood   Other source   Graft manipulation ex-vivo including T-Cell depletion other than for RBC removal or volume reduction   No   Yes   T-cell (CD3+) depletion (do not use for "Compathbag")   T-cell receptor αβ depletion   No   Yes   Positive:   No   Yes   CD34+ enrichment   Positive:   No   Yes   T-cell receptor αβ depletion   No   Yes   Positive:   No   Yes   CD34+ enrichment   T-cell receptor αβ depletion   No   Yes   T-cell receptor αβ depletion   No   No   No   No   No   No   No	Source o	of Stem Cells for this product, select only one			
Graft manipulation ex-vivo including T-Cell depletion  other than for RBC removal or volume reduction  No					
other than far RBC removal or volume reduction   No		_			
No					
Yes   Negative:   No   Yes:	I				
T-cell receptor aß depletion   B-cell depletion (CD19+) by MoAB   Not cell depletion (CD19+) by MoAB   Not cell depletion by MoAB   Other   Nother   Nothe					
B-cell depletion (CD19+) by MoAB   NX cell depletion by MoAB   NX cell depletion by MoAB   Other   NX cell depletion   NX cell depletion by MoAB   Other   NX cell depletion   NX cell depletion by MoAB   Other   NX cell depletion   NX cell depletion by MoAB   Other   NX cell depletion   NX cell depletion by MoAB   Other   NX cell depletion   NX cell deple					
Other   Positive:   No   Yes   CD34+ enrichment   Genetic manipulation   No   Yes			_		
Positive:   No   Yes   CD34+ enrichment   Genetic manipulation   No   Yes					
CD34+ enrichment			Utner		
Please enter the LABORATORY RESULTS WITH HLA TYPING into the database    Donor 2 - Product Number 2		Positive: No Yes	CD241 oprichment		
Please enter the LABORATORY RESULTS WITH HLA TYPING into the database    Donor 2 - Product Number 2					
Donor 2 - Product Number 2  If more than one stem cell product, this is the SECOND product infused from this donor    Source of Stem Cells for this product, select only one   Bone marrow   Peripheral blood   Cord blood   Other source   Cord blood   Other source   Peripheral blood   Cord blood   Other source   Cord blood   Other source   Cord blood   Other source   Cord blood   Cord b		Genetic manipulation No	☐ Yes		
If more than one stem cell product, this is the SECOND product infused from this donor    Source of Stem Cells for this product, select only one		Donor	2 - Product Numb	er 2	
Source of Stem Cells for this product, select only one  Bone marrow Peripheral blood Cord blood Other source  Graft manipulation ex-vivo including T-Cell depletion other than for RBC removal or volume reduction No Yes Negative: No Yes:  T-cell (CD3+) depletion (do not use for "Campathbag") T-cell receptor αβ depletion B-cell depletion (CD19+) by MoAB NK cell depletion by MoAB Other  Positive: No Yes  CD34+ enrichment	If more th				
Graft manipulation ex-vivo including T-Cell depletion  other than for RBC removal or volume reduction  No Yes Negative: No Yes:  T-cell (CD3+) depletion (do not use for "Campathbag")  T-cell receptor αβ depletion B-cell depletion (CD19+) by MoAB  NK cell depletion by MoAB  Other  Positive: No Yes  CD34+ enrichment	□ Во	one marrow Peripheral blood			
other than for RBC removal or volume reduction  No Yes Negative: No Yes:  T-cell (CD3+) depletion (do not use for "Campathbag")  T-cell receptor αβ depletion B-cell depletion (CD19+) by MoAB NK cell depletion by MoAB Other  Positive: No Yes  CD34+ enrichment	Graft ma	enipulation ex-vivo including T-Cell depletion			
CD34+ enrichment	other the	an for RBC removal or volume reduction	T-cell (CD3+) depletion (do  T-cell receptor αβ depletio  B-cell depletion (CD19+) by  NK cell depletion by MoAB	n / MoAB	
		Positive: No Yes			
Genetic manipulation No Yes			CD34+ enrichment		
		Genetic manipulation No	Yes		

Please enter the LABORATORY RESULTS WITH HLA TYPING into the database

CIC:	Hospital UPN:	Patient UIC	HSCT Date:	yyyy - mm - dd
		HSCT (Continue	ed)	
Chronolo	gical number of HSCT for this patient?  If >1, date of last HSCT before this one  If >1, type of last HSCT before this one  If >1 and Allograft, Was the same donor u  If >1, was last HSCT peformed at another	institution? No	Yes: CIC if known	
	If >1, please submit an Annual follow subsequent transplant as the date of (This is so we can capture relapse dat	last contact		
HSCT pa	rt of a planned multiple (sequential) a	graft protocol (program)?		
		Preparative Regin	nen	
Prepara	tive (conditioning) regimen given?  No (Usually Paed Inherited Disorders o  Yes	nly) Go to GvHD Prophylaxis		
Was thi	intended to be myeloablative? (allo Yes		ditions	
Drugs (include d	□ No □ Yes		herapy, etc.)	

CIC:	Hospital UPN:	Patient UIC	 HSCT Date:	
			 	yyyy - mm - dd

## Specification and dose of the preparative regimen

TOTAL PRESCRIBED CUMULATIVE DOSE* as per protocol:				
DRUG (given before day 0)	DOSE		UNIT	S
Ara-C (cytarabine)	2002	mg/m2	mg/kg	
ALG, ATG (ALS/ ATS)		mg/m2	mg/kg	
Animal origin: Horse				
Rabbit				
Other, specify				
Bleomycin		mg/m2	☐ mg/kg	
Busulfan		mg/m2	☐ mg/kg	mg x hr/L
Oral IV Both				micromol x min/L mg x min/mL
BCNU		mg/m2	mg/kg	
Bexxar (radio labelled MoAB)		☐ mCi	☐ MBq	
CCNU		mg/m2	mg/kg	
Campath (AntiCD 52)		mg/m2	mg/kg	
☐ Carboplatin		mg/m2	☐ mg/kg	mg x hr/L micromol x min/L mg x min/mL
☐ Cisplatin		mg/m2	☐ mg/kg	
☐ Clofarabine		mg/m2	mg/kg	
Corticosteroids		mg/m2	mg/kg	
☐ Cyclophosphamide		mg/m2	mg/kg	
☐ Daunorubicin		mg/m2	mg/kg	
Doxorubicin (adriamycine)		mg/m2	☐ mg/kg	
Epirubicin		mg/m2	mg/kg	
Etoposide (VP16)		mg/m2	mg/kg	
☐ Fludarabine		mg/m2	mg/kg	
☐ Gemtuzumab		mg/m2	mg/kg	
☐ Idarubicin		mg/m2	mg/kg	
☐ Ifosfamide		mg/m2	☐ mg/kg	
☐ Imatinib mesylate		mg/m2	mg/kg	
☐ Melphalan		mg/m2	mg/kg	
Mitoxantrone		mg/m2	mg/kg	
☐ Paclitaxel		mg/m2	☐ mg/kg	
Rituximab (mabthera, antiCD20)		mg/m2	☐ mg/kg	
☐ Teniposide		☐ mg/m2	☐ mg/kg	
☐ Thiotepa		mg/m2	mg/kg	
☐ Treosulphan		mg/m2	☐ mg/kg	
Zevalin (radiolabelled MoAB)		☐ mCi	MBq	
Other radiolabelled MoAB		☐ mCi	☐ MBq	
Specify				
Other MoAB, specify		mg/m2	mg/kg	
Other, specify		mg/m2	mg/kg	

<sup>\*</sup>Report the total prescribed cumulative dose as per protocol. Multiply daily dose in mg/kg or mg/m² by the number of days; e.g. for Busulfan given 4mg/kg daily for 4days, total dose to report is 16mg/kg

<sup>\*\*</sup>AUC = Area under the curve

CIC:	Hospital UPN:	Patient UIC	HSCT Date:
Total Body Irradiation (TBI)	N.	□ Vee . Tatal grandled . P. C.	
Total Body IITadiation (TBI)	☐ No	Yes : Total prescribed radiation dose as	
		Number of fractions	over radiation days
TLI, TNI, TAI	☐ No	Yes: Total prescribed radiation dose a	as per protocolGy
(lymphoid, nodal, abdominal)			
GvHD prophylaxis or pre	ventive treatn	ent (Allografts only)	
□ No □ Yes	ventive treati	che (mograficom)	
If Yes: Drugs (Immun	osuppressive che	no)	
ALG, ALS Anti CD2 Campati Systemic Cyclospo Cycloph Etanero FK 506 Inflixima Methoto Mycoph Sirolimu Other no	S, ATG, ATS: (giv 25(MoAB in vivo) h (MoAB in vivo; c corticosteroids orine osphamide (give ept (MoAB in vivo; (Tacrolimus, Prog ab (MoAB in vivo; rexate enolate (MMF) s nonoclonal antibo gent (in vivo), spe I photopheresis (	en after day 0) Animal origin: Horse  an be "in the bag")  n after day 0)  raf)  dy (in vivo), specify	Rabbit Other, specify
Other, specify			
		Survival Status	
Survival Status on date o	•	Sulvival Status	
Patient died between  Main Cause of Dea  Relapse or Progr HSCT Related Ca Unknown Other	th (check on ession/Persistent use		
GVHD	ry cause of Dea	check as many as appropriate).	
Pulmona Infection bac vira fur pan Un Rejection History Haemor Cardiac Central	cterial al agal rasitic known n/Poor graft func of severe Veno oc rhage toxicity nervous system (0 atestinal (GI) toxic icity	clusive disorder (VOD) NS) toxicity	