CIC:	Hospital UPN:	Patient UIC	HSCT Date:					
	HSCT - Mir	nimum Essential REGISTRATION - DAY 0						
	Centre Identification							
	Unit:							
		Patient Data						
□ No □ Yes: Hospital Unique Pati Compulsory, registration	-	′ trial:	Unknown					
Initials: Date of birth:	(first name(s)	_family name(s)) Sex: Male (at birth)	☐ Female					
	Prii	mary Disease Diagnosis						
Acute Leukaem Acute Myelo related Precursor Ly Therapy related Secondary Acute Chronic Leukaem Chronic Myelo	nia ogenous Leukaemia (AML) ursor Neoplasms mphoid Neoplasms (old ALL d myeloid neoplasms (old te Leukaemia) emia eloid Leukaemia (CML) phocytic Leukaemia (CLL)	SE FOR WHICH THIS TRANSPLANT WAS PERI ☐ Myeloma/Plasma cell disorder ☐ Solid Tumour ☐ Myelodysplastic syndromes /	Histiocytic disorders Autoimmune disease Juvenile Idiopathic Arthritis Multiple Sclerosis Systemic Lupus Systemic Sclerosis Haemoglobinopathy					
Other diagnosis	, specify:							

CIC: Hos	pital UPN:	Patient UIC	HSC	CT Date: yyyy - mm - dd
		JKAEMIAS (main relogenous Leuka	disease code 2)	
		Disease	(01112)	
Data of Initial Diagnosis:				
Date of Initial Diagnosis:	yyyy - mm - dd			
Classification: (CMML is not	a CML but MDS/MPN	")		
At least one investigation must	•	resent	luotod	
(, , _		resent		
DCI-abi ^				
	l	reatment Pre-HS	C1	
Treatment pre-HSCT (primary t	reatment)			
□ No - Includes supportive	e care or treatment wit	thout Tyrosine Kinase Inh	ibitor (TKI) or chemothe	rapy
☐ Yes Date Treatment start	ed	-		
	уууу - тт - с	dd		
Tyrosine Kinase Inhibitor (TKI): No			
	☐ Yes	Imatinib mesylate	е	
		Nilotinib		
		Dasatinib		
		Bosutinib		
		Ponatinib		
		Other TKI, specif	fy:	
Other chemotherapy, spec	ify:			
		Status at HSCT	-	
Data of this UCCT.				
Date of this HSCT:	y - mm - dd			
PHASE	NUMBER	TYPE OF REMISSION		<u></u>
☐ Chronic phase (CP)	☐ 1st	HAEMATOLOGICAL	CYTOGENETIC	MOLECULAR
	2nd	☐ No	☐ No	□ No
	3rd or higher	Yes	☐ Yes	Yes
		☐ Not evaluated	☐ Not evaluated	Not evaluated
		Unknown	☐ Not Applicable*	☐ Not Applicable*
			Unknown	Unknown
Accelerated phase	☐ 1st ☐ 2nd			
	3rd or higher			
□ Plact crisis		-		
☐ Blast crisis	☐ 1st ☐ 2nd			
	☐ 3rd or higher			

^{*} No abnormalities detected prior to this time point

CIC: Hosp	oital UPN: Patient UIC H	SCT Date:	уууу -	mm - d	d			
	HSCT							
Performance score system used ☐ Karnofsky ☐ Lansky Score ☐ 10 ☐ 20 ☐ 30 ☐ 40 ☐ 50 ☐ 60 ☐ 70 ☐ 80 ☐ 90 ☐ 100 Weight (kg):								
weight (kg).	neight (chi):							
	Comorbidity Index							
forror et al., Blood, 2005 Oct 15;	106(8): 2912-2919: http://www.ncbi.nlm.nih.gov/pmc/articles/PM	IC1895304/						
Vas there any <i>clinically significar</i> oreparative regimen? No Yes	t co-existing disease or organ impairment at time of patient assessr	ment just prior	to the					
Comorbidity	Definitions		No	Yes	N/E			
Solid tumour, previously present	Treated at any time point in the patient's past history, excluding no melanoma skin cancer	on-						
	Indicate type							
nfammatory bowel disease	Crohn's disease or ulcerative colitis							
Rheumatologic	SLE, RA, polymyositis, mixed CTD, or polymyalgia rheumatica							
nfection	Requiring continuation of antimicrobial treatment after day 0							
Diabetes	Requiring treatment with insulin or oral hypoglycaemics but not diet alone							
Renal: moderate/severe	Serum creatinine > 2 mg/dL or >177 μ mol/L, on dialysis, or prior retransplantation	enal						
Hepatic: mild	Chronic hepatitis, bilirubin between Upper Limit Normal (ULN) and ULN, or AST/ALT between ULN and 2.5 × ULN							
moderate/ severe	Liver cirrhosis, bilirubin greater than 1.5 × ULN, or AST/ALT greate × ULN	r tnan 2.5						
Arrhythmia	Atrial fibrillation or flutter, sick sinus syndrome, or ventricular arrhythmias							
Cardiac	Coronary artery disease, congestive heart failure, myocardial infar 50%, or shortening fraction in children (<28%)	rction, EF ≤						
Cerebrovascular disease	Transient ischemic attack or cerebrovascular accident							
Heart valve disease	Except mitral valve prolapse							
Pulmonary: moderate	DLco and/or FEV1 66-80% or dyspnoea on slight activity							
severe	DLco and/or FEV1 ≤ 65% or dyspnoea at rest or requiring oxygen							
Dbesity	Patients with a body mass index > 35 kg/m2							
Peptic ulcer	Requiring treatment							
Psychiatric disturbance	Depression or anxiety requiring psychiatric consultation or treatm	ent						

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

CIC:	Hospital UPN	:	Patient UIC	HSCT	
					yyyy - mm - dd
		Type of	HSCT (Alloge	eneic)	
☐ Allogeneic					
Patient CMV statu	ıs N	egative Po	sitive Not eva	luated Unknow	n
Multiple donors (including multiple	CB units)	o Ye	S: Number of donors		
			Donor 1		
HLA MATCH TYPE (DON HLA - Identical sibli Syngeneic (monozy) HLA - Matched other	ng <i>(may include n</i> gotic twin) er relative		match 1 HLA locu	ıs mismatch oci mismatch	
Donor ID given b	by the centre				
HLA MISMATCHES (Mismatched relatives o	BETWEEN DONOR	AND PATIENT			
Complete num	ber of mismatches	inside each box			
A B	C DRE	1 DQB1 DPB1			
0=match; 1=one mismat	ch; 2=2 mismatches,	N/E=not evaluated	Antigenic Allelic		
Unrelated donor					
ION code of the Donor Re	· ,				
BMDW code of the Dono Name of Donor Registry/	- ,		e is unknown) (up to 4 ch	•	
Donor centr	.,	y of the above code	es is unknown)		
	(1) 5.77	<i>icable, optional)</i> Donor Registry or t	he CB Bank listed above		
			the CB Bank listed above		
			/ITH HLA TYPING into the		
Donor information					
Date of birth	 - mm - dd	<u>OR</u>	Age at time of donation	(if date of birth not pi	·
Donor Sex	(at birth)	Male	Female		101(3)
Donor CMV	' status	Negative	Positive	□ Not evaluated	Unknown
Did this donor provide more	e than one stem c	ell product		_	
Yes: Num	ber of different st	em cell products in	ber 1" on next page fused from this donor nor 1 – Product Number 1	AND 2" on next page)	

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

Please enter the LABORATORY RESULTS WITH HLA TYPING into the database

CIC:	Hospital UPN:		Patient UIC	HSCT Date:	уууу - тт - с
			onor 2		
HLA MATCH TYPE (DONOR RELATION WITH PATIEN	T)			
HLA - Id	lentical sibling (may include	e non-monozygot	ric twin)		
Syngen	eic <i>(monozygotic twin)</i>				
	latched other relative				
∐ HLA - M	lismatched relative Degree	e of mismatch	☐ 1 HLA locus misma ☐ >=2 HLA loci misma		
HLA MISMATCH (Mismatched relative	IES BETWEEN DONOR AND PATII es only)	ENT			
Complete no	umber of mismatches inside eac	ch box			
Α	B C DRB1 DQB1	DPB1			
		Antig	genic		
	一一一一	Allel			
0=match: 1=one mis	match; 2=2 mismatches; N/E=not ev		C		
Unrelated					
ION code of the	Donor Registry or CB Bank				
	he Donor Registry or CB Bank			cters)	
Name of Donor R	Registry/ CB Bank (If any of t	he above codes is	unknown)		
Dono	or centre name (if applicable	, optional)			
Donor	ID given by the Donor Registry	or the CB Bank li	sted above		
Patien	t ID given by the Donor Registi	ry or the CB Bank	listed above		
	Please enter the LABORATOR	RY RESULTS WITH	I HLA TYPING into the dat	:abase	
Donor information	1				
Date of birth	yyyy - mm - dd	OR /	Age at time of donation	(if date of birth not provided,)
Donor Sex		☐ Female	yea	r(s)month(s))
			□ Nat avaluated		
Donor CMV status	☐ Negative	☐ Positive		Unknown	
Did this donor provid	de more than one stem cell prod	luct			
☐ No	(please fill "Donor 1 – Prod s: Number of different stem cell		· · · · · · · · · · · · · · · · · · ·		
	(If 2 products e.g. BM PB, ple				

CIC:	Hospital UPN:	Patient UIC	HSCT Date:	yyyy - mm - dd
	Don	or 2 - Product Numbe	er 1	
If mare than one stor	n cell product, this is the FIRST p), i	
	s for this product, select only or			
☐ Bone marrow				
Cord blood	Other source			
Graft manipulation e	ex-vivo including T-Cell depletion	1		
	emoval or volume reduction			
│	Negative: No	Yes:		
		T-cell (CD3+) depletion (do n	ot use for "Campathbag")	
		T-cell receptor αβ depletionB-cell depletion (CD19+) by N	MoAB	
		NK cell depletion by MoAB		
Dog	sitive: No Yes			
100	sitive: No Yes	CD34+ enrichment		
Genet	ic manipulation N	lo Yes		
Please enter	the LABORATORY RESULT	S WITH HLA TYPING into the da	atabase	
	Don	or 2 - Product Numbe	er 2	
If more than one ster		D product infused from this donor		
Source of Stem Cells	s for this product, select only or	ne		
Bone marrow	Peripheral blood	-		
Cord blood	Other source			
Graft manipulation e	ex-vivo including T-Cell depletion	1		
	emoval or volume reduction			
│	Negative: No No	Yes:		
	0 —	T-cell (CD3+) depletion (do n	ot use for "Campathbag")	
		T-cell receptor αβ depletionB-cell depletion (CD19+) by N	MoAB	
		NK cell depletion by MoAB		
		∟ Otner		
Pos	itive: No Yes	CD34+ enrichment		
Genet	ic manipulation N	o Yes		

Please enter the LABORATORY RESULTS WITH HLA TYPING into the database

CIC:	Hospital UPN:	Patient UIC	HSCT Date:	yyyy - mm - dd
		HSCT (Continued)		
	cal 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 used If >1, was last HSCT peformed at another institution.			
	f >1, please submit an Annual follow up to subsequent transplant as the date of last (This is so we can capture relapse data and	contact		
HSCT par	t of a planned multiple (sequential) graft	: protocol (program)?		
	F	Preparative Regimen		
	ive (conditioning) regimen given? No (Usually Paed Inherited Disorders only) (Yes	Go to GvHD Prophylaxis		
	intended to be myeloablative? (allo only	☐ Age of recipient ☐ Comorbid conditions ☐ Prior HSCT ☐ Protocol driven		
Drugs (include a	☐ No ☐ Yes	☐ Unknown ody, polyclonal antibody, serotherapy, et	tc.)	

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)		mg/m2	mg/kg		
ALG, ATG (ALS/ ATS)		mg/m2	mg/kg		
Animal origin: Horse					
Rabbit					
Other, specify					
			□ ma/lea		
Bleomycin Busulfan		☐ mg/m2	☐ mg/kg		
		mg/m2	mg/kg	mg x hr/L micromol x min/L	
☐ Oral ☐ IV ☐ Both				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	
Carbopiatiii		IIIg/IIIZ	□ IIIg/kg	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		
			1		

^{*}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

^{**}AUC = Area under the curve

CIC:	Hospital UPN	:	Patient UIC	HSCT Date:	yyyy - mm - dd
Total Body Irradiation (TRI)		. Takal muses the advantage of the distance of		
Total Body IITadiation (IBI) No		: Total prescribed radiation dose as		
		Nu	umber of fractions c	over ra	adiation days
TLI, TNI, TAI	☐ No	Yes	: Total prescribed radiation dose a	s per protocol	Gy
(lymphoid, nodal, abdomin	al)				
GvHD prophylaxis or	preventive trea	tment (/	Allografts only)		
□ No □ Yes	preventive tree	terricite (
If Yes: Drugs (Imi	nunosuppressive c	hemo)			
ALG Anti Cam Syst Cycl Etar FK 5 Infli Met Myc Siro Oth Oth Extracorporation Extracorporation Extracorporation Extracorporation Campaigness Campai	, ALS, ATG, ATS: (CD25(MoAB in vivor) path (MoAB in vivor) emic corticosteroicosporine ophosphamide (intercept (MoAB in vivor) compatible (in v	given after of o) o; can be "in of one of on	ivo) , specify	∏ Rabbit ☐ Other, sρ	oecify
Other, spe	cify				
			Survival Status		
Main Cause of Relapse or P HSCT Relate Unknown Other Contrib GVH Inte	Dead reen administration Death (check rogression/Persisted Cause utory Cause of D	ent disease eath (c	check as many as appropriate):		
☐ Card ☐ Cen ☐ Gas ☐ Skin ☐ Ren ☐ Mul	diac toxicity tral nervous syster trointestinal (GI) to toxicity al failure tiple organ failure	xicity	city		