CIC:	Hospital UPN:	Patient UIC	HSCT Date:				
	HSCT - Min	imum Essential I					
	Centre Identification						
	Unit:						
		Patient Data					
Date of this report: First transplant for this patient?: Yes No Yes No Yes: Name of study / trial: Unknown Unknown							
	(first name(s)	Sex:	☐ Female				
	Prir	nary Disease Diagnosis					
	nosis: yyyy - mm - dd DIAGNOSIS (CHECK THE DISEAS	E FOR WHICH THIS TRANSPLANT WAS PERFO	ORMED)				
related Pre Precursor L Therapy relate Secondary Act Chronic Leuka	ogenous Leukaemia (AML) cursor Neoplasms ymphoid Neoplasms (old ALL) ed myeloid neoplasms (old ute Leukaemia) nemia reloid Leukaemia (CML) nphocytic Leukaemia (CLL) in	 Myeloma/Plasma cell disorder Solid Tumour Myelodysplastic syndromes / Myeloproliferative neoplasm MDS MDS/MPN Myeloproliferative neoplasm Bone marrow failure including Aplastic anaemia Inherited disorders Primary immune deficiencies Metabolic disorders 	 ☐ Histiocytic disorders ☐ Autoimmune disease ☐ Juvenile Idiopathic Arthritis ☐ Multiple Sclerosis ☐ Systemic Lupus ☐ Systemic Sclerosis ☐ Haemoglobinopathy 				

CIC:	Hospital UPN:	Patient UIC	HSCT Date:	yyyy - mm - dd
	LYMP	HOMAS (main disease code		
		Hodgkin Lymphomas		
		Disease		
Date of Initial Diagnosis:	yyyy - mm - dd			
Classification:				
☐ Nodular lymphocyte p				
Classical predominantOther , specify:				

CIC:	Hospital UPN:	Patient UIC	HS0	CT Date:	yyyy - mm - dd			
		ALL LY/MDLIONAAO			yyyy - mm - dd			
		ALL LYMPHOMAS						
	Treatment Pre-HSCT							
Treatment pre-HSCT	Enter first day of	treatment and mark all drugs from	that date until co	nditioning	1			
Yes Date of treatmer	nt 	-						
Drugs given								
Antibodies:	Alemtuzumab (MabCampath) (CD52)						
	☐ Brentuximab (A	dcetris) (CD30)						
	Obinutuzumab	(Gyzeva) (CD20)						
	Ofatumumab (A	Azerra) (CD20)						
	☐ Rituximab (Mak	othera) (CD20)						
	other antibody,	specify						
Radioimmunotherapy:	☐ Bexxar (CD20) (radiolabelled MoAB)						
	Zevalin (CD20) ((radiolabelled MoAB)	Relapse	/progres	sion under this drug			
			Yes	No U	nknown			
Specific inhibitors:	☐ ABT-199 (BCL2-	Inhibitor)						
	Crizotinib (ALK-	Inhibitor)						
	CC-292 (B cell re	eceptor kinase inhibitor)						
	☐ Ibrutinib (B cell	receptor kinase inhibitor)						
		receptor kinase inhibitor)						
	·	specify						
Other:	☐ Bortezomib (Ve	lcade)						
	Lenalidomide (F	Revlimid)						

Other, specify

CIC:	Hospital UPN:	Pa	tient UIC	HSCT [Date:
	<u>'</u>				Date:
		ALL LYI	MPHOMAS		
		Status	at HSCT		
Date of this HSCT:	yyyy - mm - dd				
Number of prior lines of	f treatment	□ 1 □ 2	3 or more	e: none	Unknown
(since diagnosis if 1st trans	splant, or since last reporte	d transplant)			
Technique used for	disease assessmer	nt:			
	CT scan done	□ No [Yes		
	PET	Negative	Positive	☐ Not evaluated	
STATUS					
Never treated					
Complete remissio					
Unconfirm		Confirmed	ios of unknown signi	ificanca	
		ersistent scan abnormaliti	es of unknown signi	incance	
	PR) – (with or without a	prior CR)			
Stable disease	//				
		untreated progression (fro			
Disease status unk		ncluding primary refractor	ry disease		
Discuse status unik	anown				
Was this patient refrac	tory to any line of chem	notherapy before this HSC	T? No	Yes	
Number of Complete (Count all CR including th		e patient prior to this HSC	T:		
Number of Partial remi		the patient prior to this H	SCT:		

CIC: Hosp	pital UPN: Patient UIC HSCT	Date:	уууу -	mm - d	d
	HSCT				
Performance score Score		90	100		
Weight (kg):	neight (cm).				
	Comorbidity Index				
forror et al., Blood, 2005 Oct 15;	106(8): 2912-2919: http://www.ncbi.nlm.nih.gov/pmc/articles/PMC18	95304/			
Vas there any <i>clinically significar</i> oreparative regimen? No Yes	ot co-existing disease or organ impairment at time of patient assessmen	t 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 non-melanoma skin cancer				
	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 renal transplantation				
Hepatic: mild	Chronic hepatitis, bilirubin between Upper Limit Normal (ULN) and 1.9 ULN, or AST/ALT between ULN and 2.5 × ULN				
moderate/ severe	Liver cirrhosis, bilirubin greater than 1.5 × ULN, or AST/ALT greater that × ULN	all 2.5			
Arrhythmia	Atrial fibrillation or flutter, sick sinus syndrome, or ventricular arrhythmias				
Cardiac	Coronary artery disease, congestive heart failure, myocardial infarctio 50%, or shortening fraction in children (<28%)	n, 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 treatment				

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

CIC:	Hospital UPN:	Patient UIC	HSCT	Date:			
				yyyy - mm - dd			
Type of HSCT (Allogeneic)							
☐ 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						
HLA 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		aatabase				
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	· 1	
If we are the arrange at an			•	
	n cell product, this is the FIRST pro			
	for this product , select only one	eral blood		
☐ Bone marrow ☐ Cord blood	Other:			
other than for RBC r	ex-vivo of this product including T- emoval or volume reduction Negative: No Yes Genetic manipulation the LABORATORY RESULTS		oAB	
If more than one ster	Donc	or 1 - Product Numbe	r 2	
	for this product , select only one			
☐ Bone marrow	Periph	eral blood		
Cord blood	Other:			
	ex-vivo of this product including T- emoval or volume reduction Negative:		оАВ	
	Positive: No Yes	CD34+ enrichment		
	Genetic manipulation	☐ No ☐ Yes		

Please enter the LABORATORY RESULTS WITH HLA TYPING into the database

CIC:	lospital UPN:	Pat	tient UIC	HSCT Date: yyyy - mm - dd
		Do	onor 2	yyyy - mm - dd
		DC) O	
HLA MATCH TYPE (DONOR REL	ATION WITH PATIENT)	1		
HLA - Identical sibli	ng <i>(may include i</i>	non-monozygotic	twin)	
Syngeneic (mo	nozygotic twin)			
☐ HLA - Matched oth				
HLA - Mismatched	relative Degree o	of mismatch	☐ 1 HLA locus mismatch ☐ >=2 HLA loci mismatch	
			>=2 TILA IOCI IIIISIII atcii	
HLA MISMATCHES BETWEE (Mismatched relatives only)	N DONOR AND PATIEN	IT		
Complete number of m	ismatches inside each	box		
A B C	DRB1 DQB1	DPB1		
		Antiger	nic	
885		Antiger		
		Allelic		
0=match; 1=one mismatch; 2=2 n	nismatches; N/E=not eval	uated		
Unrelated donor				
ION code of the Donor Regis	•			
BMDW code of the Donor Re Name of Donor Registry/ CB		(If ION code is un above codes is u	n/m n.u.m.)	
Donor centre na			ikilowiij	
_	y the Donor Registry o by the Donor Registry			
i atient 15 given	by the bollor negistry	or the eb bank na		
Please e	nter the LABORATORY	RESULTS WITH H	LA TYPING into the database	
Donor information				
Date of birth		<u>OR</u> Age	e at time of donation (if date	of birth not provided)
ууу	y - mm - dd		year(s)	month(s)
Donor Sex (at birth)	Male	Female		
Donor CMV status	Negative	Positive	☐ Not evaluated ☐ U	Jnknown
Did this donor provide more that	ın one stem cell produ	ct		
	fill "Donor 1 – Produ			
	of different stem cell pr Jucts e.a. BM PB. pleas		om this donor Product Number 1 AND 2" on ne	xt paae)
(1) = p100		- , = - ,		- 1- 3-1

CIC:	Hospital UPN:	Patient UIC	HSCT Date:	yyyy - mm - dd					
	Don	or 2 - Product Numbe	er 1						
If more than one stor			J. 1						
	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 e	ex-vivo including T-Cell depletion	1							
	emoval or volume reduction								
│	Negative: No	Yes:							
		T-cell (CD3+) depletion (do n							
		T-cell receptor αβ depletion B-cell depletion (CD19+) by I							
		NK cell depletion by MoAB							
Por	sitive: No Yes	- Other							
103	itive: No Yes	CD34+ enrichment							
Genet	ic manipulation N	o 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 sten		D product infused from this donor							
Source of Stem Cells	for this product, select only on	ne e							
Bone marrow	Peripheral blood	-							
Cord blood	Other source								
Graft manipulation e	x-vivo including T-Cell depletion	1							
	emoval or volume reduction								
│	Negative: No	Yes:							
		T-cell (CD3+) depletion (do n							
		T-cell receptor αβ depletion B-cell depletion (CD19+) by I							
		NK cell depletion by MoAB							
		□ Utner							
Pos	itive: No Yes	CD34+ enrichment							
Const	ic manipulation No	_							
Genet	ic manipulation No								

Please enter the LABORATORY RESULTS WITH HLA TYPING into the database

	Hospital UPN:	Patient UIC	HSCT Date:	yyyy - mm - dd				
		HSCT (Continued)						
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 use If >1, was last HSCT peformed at another in:		☐ No ☐ Yes CIC if known					
	If >1, please submit an Annual follow up form before proceeding, giving the date of the subsequent transplant as the date of last contact (This is so we can capture relapse data and other events between transplants).							
HSCT pa	ort of a planned multiple (sequential) gra	aft protocol (program)?						
		Preparative Regimen						
Preparative (conditioning) regimen given? No (Usually Paed Inherited Disorders only) Go to GvHD Prophylaxis Yes								
Was thi		Age of recipientComorbid conditionsPrior HSCTProtocol driven						

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

Specification and dose of the preparative regimen

	TOTAL PRESCRIBED CUMULATIVE DOSE* as per protocol:						
DRU	JG (given before day 0)	DOSE				UNIT	S
	Ara-C (cytarabine)			mg/m2		mg/kg	
	ALG, ATG (ALS/ ATS)			mg/m2	Ē	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	

^{*}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:
Total Body Irradiation (TBI)	NI	□ Vee . Tetel green the deal to the	
Total Body Illadiation (TBI)	☐ No	Yes : Total prescribed radiation dose a	
		Number of fractions	over radiation days
TLI, TNI, TAI	☐ No	Yes: Total prescribed radiation dose	as per protocolGy
(lymphoid, nodal, abdominal)			
GvHD prophylaxis or pre	ventive treatn	ent (Allografts only)	
□ No □ Yes		City (mogregoe cm//	
If Yes: Drugs (Immuno	osuppressive che	00)	
ALG, ALS Anti CD2 Campatl Systemic Cyclospo Cyclopho Etanerce FK 506 Inflixima Methotr Mycoph Sirolimu Other ne Extracorporeal	S, ATG, ATS: (gives) S, (MoAB in vivo) S, (MoAB in vivo) S, (Corticosteroids) Sorine Sosphamide (gives) Sept (MoAB in vivo) S, (Tacrolimus, Program S, (MoAB in vivo) Sexate Senonoclonal antibologent (in vivo), specific photopheresis (an be "in the bag") In after day 0) In after day 0) In after day 0) In after day 0. In after day 0.	Rabbit Other, specify
Other, specify			
		Survival Status	
Survival Status on date of		Gui vivai Gtatas	
Patient died between Main Cause of Dea Relapse or Progr HSCT Related Ca Unknown Other	th (check onlession/Persistent use		
GVHD	i y cause of Bee	(check as many as appropriate).	
Pulmona Infection bac vira fun par Uni Rejectio History o Haemor Cardiac	eterial al gal rasitic known n/Poor graft func of severe Veno oc rhage toxicity nervous system (0 itestinal (GI) toxic city illure	lusive disorder (VOD)	