	Centre Identification	
EBMT Code (CIC):	Contact person:	
Hospital: Unit:	Email:	
	Patient Data	
Date of this report:	First transplant for this patient?: 🗌 Y	′es □No
Patient following national / international study /	trial:	
□ No □ Yes: Name of study / trial	Ur	nknown
Hospital Unique Patient Number or Code (UPN) Compulsory, registrations will not be accepted without All transplants performed in the same patient must be the patient and <u>not</u> to the transplant.	ut this item.	tion number or code as this belongs to
Initials: (first name(s) _	_family name(s))	
Date of birth:	Sex: DMale (at birth)	Emale
Prir	mary Disease Diagnosis	
Date of initial diagnosis:		
PRIMARY DISEASE DIAGNOSIS (CHECK THE DISEAS	SE FOR WHICH THIS TRANSPLANT WAS PERFO	RMED)
<ul> <li>Acute Leukaemia</li> <li>Acute Myelogenous Leukaemia (AML) related Precursor Neoplasms</li> <li>Precursor Lymphoid Neoplasms (old ALL)</li> <li>Therapy related myeloid neoplasms (old Secondary Acute Leukaemia)</li> <li>Chronic Leukaemia</li> <li>Chronic Myeloid Leukaemia (CML)</li> <li>Chronic Lymphocytic Leukaemia (CLL)</li> <li>Lymphoma</li> <li>Non Hodgkin</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	I: Patient UIC		HSCT Date:	yyyy - mm - dd			
	CHRONIC	LEUKAEMIAS (main						
	Chronic Lymphocytic leukaemias (CLL)							
		Disease	· · · · · ·					
Date of Initial D	iagnosis							
Classification: Chronic lym Richter's sy	Date of Initial Diagnosis         yyyy - mm - dd         Classification:         Chronic lymphocytic leukaemia (CLL)/small lymphocytic lymphoma         Richter's syndrome         Transformed from a previously known CLL         Yes       : Date of original CLL diagnosis         yyyy - mm - dd							
	🗌 No :Prim	ary Richter (without previous kno	own diagnosis of CLL)					
	(	Chromosome Analys	is at Diagnosis					
Chromosome An	nalysis (All methods ir Abormal		Unknown					
Trisomy Del 13q <sup>2</sup> Del 11q <sup>2</sup> del(17p) Other, s	14 22-23		AbsentPresentAbsentPresentAbsentPresentAbsentPresentAbsentPresentAbsentPresent	<ul> <li>Not evalua</li> <li>Not evalua</li> <li>Not evalua</li> <li>Not evalua</li> </ul>	ted ted			
		Molecular Markers	at Diagnosis					
Molecular marke	ers							
TP53 muta	ations 🗌 Absent	Present	Not Evaluated	🗌 Unkn	own			
		Treatment Pre	-HSCT					
Treatment pre-HSCT (primary treatment) <ul> <li>No</li> <li>Yes Date Treatment started</li> </ul>								
Regi	men	Date started	Date ended					
		yyyy - mm - dd	yyyy - mm - dd					
		Status at H						
Date of this HSC	CT:							
STATUS	CT: yyyy - mm - dd	Minimal residual di	sease (MRD) (by FACS (	or PCR)				

51A105	minimal residual disease (MRD) (by race of ren)					
<ul><li>Complete remission (CR)</li><li>Partial remission (PR)</li></ul>	Negative	Positive	Not evaluated			
<ul> <li>Stable disease (SD)</li> <li>Untreated Relapse</li> <li>Progression (PD)</li> <li>Never treated</li> </ul>						

CIC:		Hospital UPN:		Patier	nt UIC		H	SCT Date:	yyyy - mm - dd
				HSC	CT				
Performa	nce score	system us		arnofsky Insky					
Score	□ 10	□ 20 □ 30	□ 40	□ 50	□ 60	□ 70	□ 80	□ 90	□ 100
Weight (kg	):	Height (cm):							

Comorbidity Index					
Sorror et al., Blood, 2005 Oct 15; 106(8): 2912-2919: http://www.ncbi.nlm.nih.gov/pmc/articles/PMC1895304/					
Was there any <i>clinically significa</i> preparative regimen?	nt co-existing disease or organ impairment at time of patient assessment 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				
Infammatory bowel disease	Crohn's disease or ulcerative colitis				
Rheumatologic	SLE, RA, polymyositis, mixed CTD, or polymyalgia rheumatica				
Infection	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 $\mu$ mol/L, on dialysis, or prior renal transplantation				
Hepatic: mild moderate/ severe	Chronic hepatitis, bilirubin between Upper Limit Normal (ULN) and 1.5 x the ULN, or AST/ALT between ULN and 2.5 × ULN Liver cirrhosis, bilirubin greater than 1.5 × ULN, or AST/ALT greater than 2.5 × ULN				
Arrhythmia	Atrial fibrillation or flutter, sick sinus syndrome, or ventricular arrhythmias				
Cardiac	Coronary artery disease, congestive heart failure, myocardial infarction, EF ≤ 50%, or shortening fraction in children (<28%)				
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				
Obesity	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:	Hospit	tal UPN:	Patient UIC	HSCT Date:	yyyy - mm - dd		
		-			,,,,,		
Type of HSCT (Allogeneic)							
Allog	eneic						
Pati	ent CMV status	Negative	Positive Not eva	luated 🗌 Unknown			
Mul (incl	tiple donors uding multiple CB units)	🗌 No	Yes: Number of donors				
			Donor 1				
HLA -	H TYPE (DONOR RELAT Identical sibling (may in eneic (monozygotic twin) Matched other relative Mismatched relative:	clude non-monozy	ygotic twin) ee of mismatch 🛛 1 HLA locu	ıs mismatch oci mismatch			
Do	onor ID given by the cent	re					
	MISMATCHES BETWEEN tched relatives only)	DONOR AND PAT	IENT				
	omplete number of misr	natches inside ea	ich box				
	A B C	DRB1 DQB1					
			Antigenic Allelic				
_	h; 1=one mismatch; 2=2 misi	matches; N/E=not ev	valuated				
	lated donor f the Donor Registry or C	B Bank					
	le of the Donor Registry of C		fION code is unknown) (up to 4 ch	aracters)			
	onor Registry/ CB Bank		bove codes is unknown)	,			
	Donor centre name <b>Donor</b> ID given	(if applicable, opt	tional) gistry or the CB Bank listed above				
	Patient ID give	n by the Donor Re	egistry or the CB Bank listed above				
	Please enter the	e LABORATORY RI	ESULTS WITH HLA TYPING into the	database			
Donor inform	mation						
Date of birth	yyyy - mm - dd		<u>OR</u> Age at time of donation	(if date of birth not provided)			
			······	month(s)			
	Donor Sex (at birt						
Did this dono		Nega	_	Not evaluated Unknow	own		
Dia this donor	Yes: Number of diffe	Donor 1 – Produ erent stem cell pro	uct Number 1" on next page oducts infused from this donor				
	(If 2 products e	e.g. BM PB, pleas	e fill "Donor 1 – Product Number 1	AND 2" on next page)			

Donor 1 - Product Number 1					
If more than one stem cell product, this is the FIRST prod	duct infused from this donor				
Source of Stem Cells for this product , select only one					
Bone marrow Periph Periph	eral blood				
Cord blood Other:					
Graft manipulation ex-vivo of this product including T-c	cell depletion				
other than for RBC removal or volume reduction					
No Ves Negative: No Yes					
Yes Negative: No Yes	T-cell (CD3+) depletion (do not use for "Campath in bag")				
	T-cell receptor $\alpha\beta$ depletion				
	B-cell depletion (CD19+) by MoAB				
	NK cell depletion by MoAB				
	□ Other				
Positive: 🗌 No 📄 Yes					
	CD34+ enrichment				
Genetic manipulation	No Yes				

Please enter the LABORATORY RESULTS WITH HLA TYPING into the database

more than one stem cell product, this is the S	ECOND product infused from this donor
Source of Stem Cells for <b>this product</b> , select o	nly one
Bone marrow	] Peripheral blood
Cord blood Other:	
Graft manipulation ex-vivo of this product incl	uding T-cell depletion
other than for RBC removal or volume reductic	
No	
Yes Negative: No	Yes:
	T-cell (CD3+) depletion (do not use for "Campath in bag")
	$\Box$ T-cell receptor $\alpha\beta$ depletion
	B-cell depletion (CD19+) by MoAB
	NK cell depletion by MoAB
	Other
Positive: 🗖 No 🗍	☐ Yes
	CD34+ enrichment
Genetic manipulation	─ No □ Yes

 $\Rightarrow$  Please enter the LABORATORY RESULTS WITH HLA TYPING into the database

CIC:	Hospital UPN:	Patient UIC	HSCT Date:					
		Donor 2	,,,,,					
HLA MATCH TYPE (D	DONOR RELATION WITH PATIE	ENT)						
Syngene	ic (monozygotic twin) atched other relative ismatched relative Degr	nde non-monozygotic twin) ree of mismatch						
<b>HLA</b> MISMATCHE (Mismatched relatives	ES BETWEEN DONOR AND PA' s only)	TIENT						
Complete nu	mber of mismatches inside e	ach box						
A	B C DRB1 DQE	1 DPB1						
		Allelic						
0=match: 1=one misn	natch; 2=2 mismatches; N/E=not							
Unrelated								
ION code of the D	onor Registry or CB Bank							
	e Donor Registry or CB Bank							
Name of Donor Re		f the above codes is unknown)						
		ple, optional)						
		ry or the CB Bank listed above						
		ORY RESULTS WITH HLA TYPING into the da	tabase					
Donor information								
		OR Age at time of donation	(if date of birth not provided)					
	yyyy - mm - dd	<u></u>	r(s)month(s)					
Donor Sex	(at birth) 🗌 Male	Female						
Donor CMV status	Negative	Positive Not evaluated	Unknown					
Did this donor provide	Did this donor provide more than one stem cell product							
No Yes	: Number of different stem ce	oduct Number 1" on next page						
	(If 2 products e.g. BM PB, p	lease fill "Donor 1 – Product Number 1 AND	2″ on next page)					

## Donor 2 - Product Number 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 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 $\alpha\beta$ depletion
	B-cell depletion (CD19+) by MoAB
	NK cell depletion by MoAB
	Other
Positive: 🗌 No 🗌 Yes	
	CD34+ enrichment
Genetic manipulation 🗌 No	☐ Yes

Please enter the LABORATORY RESULTS WITH HLA TYPING into the database

Doll	Donor 2 - Product Number 2					
more than one stem cell product, this is the SECON	D product infused from this donor					
Source of Stem Cells for this product, select only on						
Bone marrow Peripheral blood						
Cord blood Other source						
Graft manipulation ex-vivo including T-Cell depletior						
other than for RBC removal or volume reduction						
Yes Negative: No	Yes:					
	T-cell (CD3+) depletion (do not use for "Campathbag")					
	T-cell receptor $\alpha\beta$ depletion					
	B-cell depletion (CD19+) by MoAB					
	NK cell depletion by MoAB					
	Other					
Positive: 🗌 No 🦳 Yes						
	CD34+ enrichment					
Genetic manipulation	o 🗌 Yes					

Please enter the LABORATORY RESULTS WITH HLA TYPING into the database

CIC: Hospital UPN:	Patient UIC	HSCT Date:	уууу - mm - dd				
HSCT (Continued)							
Chronological number of HSCT for this patient?     If >1, date of last HSCT before this one If >1, type of last HSCT before this one Alloc If >1 and Allograft, Was the same donor used for all pri If >1, was last HSCT peformed at another institution?	or and current HSCTs?	□ No □ Yes IC if known 					
If >1, please submit an <u>Annual follow up form</u> 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 part of a planned multiple (sequential) graft protocol (program)?          No       Yes							
Prepa	rative Regimen						
Preparative (conditioning)       regimen given?         No       (Usually Paed Inherited Disorders only) Go to GvH         Yes	No (Usually Paed Inherited Disorders only) Go to GvHD Prophylaxis						
Was this intended to be myeloablative? (allo only)	<ul> <li>Age of recipient</li> <li>Comorbid conditions</li> <li>Prior HSCT</li> <li>Protocol driven</li> <li>Other, specify</li> </ul>						
Drugs       No       Yes         (include any active agent be it chemo, monoclonal antibody, poly	Unknown clonal antibody, serotherapy, etc.)	)					

CIC:

## Specification and dose of the preparative regimen

TOTAL PRESCRIBED CUMULATIVE DOSE* as per protocol:						
DRUG (given before day 0)	DOSE	UNITS				
Ara-C (cytarabine)		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			
		mg/m2	mg/kg			
Campath (AntiCD 52)		mg/m2	mg/kg			
Carboplatin		mg/m2	🗌 mg/kg	mg x hr/L		
				micromol x min/L		
Cisplatin		mg/m2	mg/kg			
		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			
		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			
		☐ mg/m2	└ mg/kg			
L Thiotepa		mg/m2	mg/kg			
L Treosulphan		mg/m2	mg/kg			
Zevalin (radiolabelled MoAB)		∐ mCi	MBq			
Uther 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/m<sup>2</sup> 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)       No       Yes       Total prescribed radiation dose as per protocol       G         Number of fractions       over       radiation d         TLI, TNI, TAI       No       Yes       Total prescribed radiation dose as per protocol       G         Wmphoid, nodal, abdominal)       Gvert       radiation dose as per protocol       G         GvHD prophylaxis or preventive treatment       (Allografts only)       No       Yes         If Yes:       Drugs (Immunosuppressive chemo)       ALG, ALS, ATG, GATS: (given after day 0)Animal origin:       Horse       Rabbit       Other, specify         Anti CD25(MoAB in vivo)       Campath (MoAB in vivo) con be "in the bog")       Systemic corticosteroids       Cyclosporine         Cyclosporine       Cyclosporine       Cyclosporine       Cyclosporine         Cyclosporine       Cyclosporine       Other monoclonal antibody (in vivo), specify       Green controsteroids         Gyrophonelate (MMF)       Sirolimus       Other agent (in vivo), specify       Green controsteroids       Cyclosporine         Cyclosproneal photopheresis (ECP)       Other, specify       Other, specify       Cher, specify         Cher, specify       Dead       Date       Date       Date	y lays Gy					
Number of fractions       over       radiation d         TLI, TNI, TAI       No       Yes       Total prescribed radiation dose as per protocol         (lymphoid, nodal, abdominal)         GvHD prophylaxis or preventive treatment (Allografts only)         No       Yes         If Yes:       Drugs (Immunosuppressive chemo)         ALG, ALS, ATG, ATS: (given after day 0)Animal origin:       Horse         Anti CD25(MoAB in vivo)       Campath (MoAB in vivo; can be "in the bag")         Systemic corticosteroids       Cyclosporine         Cyclosporine       Cyclophosphamide (given after day 0)         Etanercept (MoAB in vivo)       Etanercept (MoAB in vivo)         Methotrexate       Mycophenolate (MMF)         Sirolimus       Other monoclonal antibody (in vivo), specify         Other agent (in vivo), specify       Extracorporeal photopheresis (ECP)         Other, specify       Other, specify         Other, specify       Survival Status on date of HSCT	lays Gy					
TIL, TNI, TAI       No       Yes       : Total prescribed radiation dose as per protocol         (lymphoid, nodal, abdominal)         GVHD prophylaxis or preventive treatment (Allografts only)         No       Yes         If Yes:       Drugs (Immunosuppressive chemo)         ALG, ALS, ATG, ATS : (given after day 0) Animal origin:       Horse       Rabbit       Other, specify         Anti CD25(MoAB in vivo)       Campath (MoAB in vivo; can be "In the bag")       Systemic corticosteroids       Other, specify	Gy					
(lymphoid, nodal, abdominal)         GvHD prophylaxis or preventive treatment (Allografts only)         No       Yes         If Yes:       Drugs (Immunosuppressive chemo)         ALG, ALS, ATG, ATS : (given after day 0)Animal origin:       Horse         Anti CD25(MoAB in vivo)       Campath (MoAB in vivo; can be "in the bag")         Systemic corticosteroids       Cyclosporine         Cyclosporine       Cyclophosphamide (given after day 0)         Etanercept (MoAB in vivo)       FK 506 (Tacrolimus, Prograf)         Infliximab (MoAB in vivo)       Methotrexate         Mycophenolate (MMF)       Sirolimus         Other monoclonal antibody (in vivo) , specify         Extracorporeal photopheresis (ECP)         Other, specify         Survival Status on date of HSCT	·					
GvHD prophylaxis or preventive treatment (Allografts only)         No       Yes         If Yes:       Drugs (Immunosuppressive chemo)         ALG, ALS, ATG, ATS : (given after day 0)Animal origin:       Horse         Atti CD25(MoAB in vivo)       Other, specify         Campath (MoAB in vivo; can be "in the bag")       Systemic corticosteroids         Cyclosporine       Cyclosporine         Cyclophosphamide (given after day 0)       Etanercept (MoAB in vivo)         FK 506 (Tacrolimus, Prograf)       Infliximab (MoAB in vivo)         Methotrexate       Mycophenolate (MMF)         Strolimus       Other agent (in vivo), specify         Other agent (in vivo), specify       Deter agent (in vivo), specify         Extracorporeal photopheresis (ECP)       Other, specify         Other, specify       Survival Status on date of HSCT						
No       Yes         If Yes:       Drugs (Immunosuppressive chemo)         ALG, ALS, ATG, ATS : (given after day 0)Animal origin:       Horse       Rabbit       Other, specify         Anti CD25(MOAB in vivo)       Campath (MOAB in vivo; can be "in the bag")       Systemic corticosteroids       Other, specify         Cyclophosphamide       (given after day 0)       Etanercept (MOAB in vivo)       Etanercept (MOAB in vivo)         FK 506       (Tacrolimus, Prograf)       Infliximab (MOAB in vivo)         Methotrexate       Mycophenolate (MMF)         Sirolimus       Other monoclonal antibody (in vivo), specify         Other agent (in vivo), specify       Extracorporeal photopheresis (ECP)         Other, specify       Survival Status on date of HSCT						
No       Yes         If Yes:       Drugs (Immunosuppressive chemo)         ALG, ALS, ATG, ATS : (given after day 0)Animal origin:       Horse       Rabbit       Other, specify         Anti CD25(MOAB in vivo)       Campath (MOAB in vivo; can be "in the bag")       Systemic corticosteroids       Other, specify         Cyclophosphamide       (given after day 0)       Etanercept (MOAB in vivo)       Etanercept (MOAB in vivo)         FK 506       (Tacrolimus, Prograf)       Infliximab (MOAB in vivo)         Methotrexate       Mycophenolate (MMF)         Sirolimus       Other monoclonal antibody (in vivo), specify         Other agent (in vivo), specify       Extracorporeal photopheresis (ECP)         Other, specify       Survival Status on date of HSCT						
ALG, ALG, ALG, ATS : (given after day 0)Animal origin: Horse Rabbit Other, specify Anti CD25(MoAB in vivo) Campath (MoAB in vivo; can be "in the bag") Systemic corticosteroids Cyclosporine Cyclophosphamide (given after day 0) Etanercept (MoAB in vivo) FK 506 (Tacrolimus, Prograf) Infliximab (MoAB in vivo) Methotrexate Mycophenolate (MMF) Sirolimus Other monoclonal antibody (in vivo) , specify Other agent (in vivo), specify Extracorporeal photopheresis (ECP) Other, specify Survival Status on date of HSCT						
Survival Status on date of HSCT						
Survival Status on date of HSCT						
Survival Status on date of HSCT	Survival Status					
<ul> <li>Anve</li></ul>						
Contributory Cause of Death (check as many as appropriate):						
GVHD         Interstitial pneumonitis         Pulmonary toxicity         Infection:         bacterial         viral         fungal         parasitic         Unknown         Rejection/Poor graft function         History of severe Veno occlusive disorder (VOD)         Haemorrhage         Cardiac toxicity         Gastrointestinal (GI) toxicity         Skin toxicity         Renal failure         Multiple organ failure         Other, specify						