	Contro Idontification	
	Centre Identification	
EBMT Code (CIC):	Contact person:	
Hospital: Unit:	Email:	
	Patient Data	
Date of this report: yyyy - mm - dd	First transplant for this patient?: 🗌 Y	es 🗌 No
Patient following national / international study /	' trial:	
□ No □ Yes: Name of study / trial	Ur	lknown
Hospital Unique Patient Number or Code (UPN) Compulsory, registrations will not be accepted withou 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	Female
Prir	mary Disease Diagnosis	
Date of initial diagnosis: yyyy - mm - dd PRIMARY DISEASE DIAGNOSIS (CHECK THE DISEAS		RMED)
Acute Leukaemia		·
 Acute Myelogenous Leukaemia (AML) related Precursor Neoplasms 	 Myeloma/Plasma cell disorder Solid Tumour Myelodysplastic syndromes / 	 Histiocytic disorders Autoimmune disease Juvenile Idiopathic Arthritis

vvvv - mm - d	ld
yyyy mmaa	u

AUTOIMMUNE DISORDERS (main disease code 10)	

CONNECTIVE TISSUE DISEASE

Classification: Systemic sclerosis (SS) Involvement/Clinical problem diffuse cutaneous limited cutaneous SSc sine scleroderma Mixed Connective Tissue Disease (MCTD) other, specify:_____ Systemic lupus erythematosus (SLE) Polymyositis- dermatomyositis Sjögren syndrome Antiphospholipid syndrome Other type of connective tissue disease, specify:_____ Date of this HSCT: yyyy - mm - dd

AUTOIMMUNE DISORDERS (main disease code 10)

VASCULITIS / ARTHRITIS / NEUROLOGICAL

AUTOIMMUNE DISORDERS – VASCULITIS

Wegener granulomatosis	
Classical polyarteritis nodosa	
Microscopic polyarteritis nodosa	
Churg-Strauss	
Giant cell arteritis	
Takayasu	
Behçet syndrome	
Overlap necrotising arteritis	
Other, specify:	
Date of this HSCT: yyyy - mm - dd	
AUTOIMMUNE DISORDERS – ARTHRITIS	
Rheumatoid arthritis	
Psoriatic arthritis/psoriasis	
Juvenile idiopathic arthritis (JIA), systemic (Stills disease)	
Juvenile idiopathic arthritis (JIA), articular: Onset	Oligoarticular
	Polyarticular
Juvenile idiopathic arthritis: other, specify:	
Other arthritis:	
Date of this HSCT:	

AUTOIMMUNE DISORDERS – NEUROLOGICAL DISEASES

MULTIPLE SCLEROSIS
Myasthenia gravis
Amyotrophic lateral sclerosis (ALS)
Chronic inflammatory demyelinating polyneuropathy (CIDP)
Neuromyelitis Optica (NMO)
Other autoimmune neurological disorder, specify:
Date of this HSCT:
yyyy - mm - dd

CIC:

AUTOIMMUNE DISORDERS (main disease code 10)

OTHER AUTOIMMUNE DISORDERS

HAEMATOLOGICAL DISEASES

Idiopathic t	hrombocytopenic purpura (ITP)
Haemolytic	anaemia
Evan syndr	ome
🗌 Autoimmu	ne lymphoproliferative syndrome (primary diagnosis, not subsequent to transplant)
Other haen	natological autoimmune disease, specify:
Date of this HS	CT: <i>yyyy - mm - dd</i>

BOWEL DISEASE

Crohn's disease	
Ulcerative colitis	
Other autoimmune bowel disease, specify:	
Date of this HSCT:	

OTHER AUTOIMMUNE

Grave's disease
other autoimmune, specify:
Date of this HSCT:

		Hospital UPN	I:		Patier	nt UIC		HS	SCT Date:	yyyy - mm - dd
					HSC	CT				
Performar	nce score	syste	m useo	d 🗌 Ka 🗌 La	rnofsky nsky					
Score		□ 20 □				□ 60	□ 70	□ 80	90	□ 100
Weight (kg):	Height	(cm):							

Comorbidity Index						
Sorror et al., Blood, 2005 Oct 15	orror 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?	Int 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 μ 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					
Arrhythmia	× ULN 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 \leq 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:	Hospi	tal UPN:	Patient UIC	HSCT Date:	
				yyyy - mm - dd	
		Тур	pe of HSCT (Alloge	eneic)	
	eneic				
-	ent CMV status	Negative	🗌 Positive 📄 Not eva	luated 🖂 Unknown	
Mult	iple donors	□ v	Yes: Number of donors		
	iding multiple CB units)				
			Donor 1		
 HLA - Synge HLA - 	I TYPE (DONOR RELAT Identical sibling (may ir neic (monozygotic twin, Matched other relative Mismatched relative:	nclude non-monoz)	ygotic twin) ee of mismatch 🛛 1 HLA loci	us mismatch oci mismatch	
Do	nor ID given by the cen	tre			
HLA N	AISMATCHES BETWEEN ched relatives only)		IENT		
	omplete number of mis	matches inside ea	ach box		
	A B C	DRB1 DQB1			
			Antigenic Allelic		
0=match;	; 1=one mismatch; 2=2 mis	matches; N/E=not e	valuated		
	ated donor				
	the Donor Registry or (
	e of the Donor Registry onor Registry/ CB Bank		f ION code is unknown) (up to 4 cl bove codes is unknown)	naracters)	
	Donor centre name	(if applicable, op			
			gistry or the CB Bank listed above		
	Patient ID give	en by the Donor R	egistry or the CB Bank listed above		
1	Please enter th	e LABORATORY R	ESULTS WITH HLA TYPING into the	database	
Donor inform	nation				
Date of birth	yyyy - mm - dd		<u>OR</u> Age at time of donation	(if date of birth not provided)	
	Donor Sex (at bir	th) 🗌 Male	e 🗌 Female		
	Donor CMV status	Nega	ative Dositive	Not evaluated Unknown	
Did this donor	provide more than one	e stem cell produc	t		
[Yes: Number of diff	erent stem cell pr	uct Number 1″ on next page oducts infused from this donor se fill ″Donor 1 – Product Number 1	AND 2" on next page)	

		product infused from this donor
Source of Stem Cells for this		
Bone marrow	Pe	ripheral blood
Cord blood	Other:	
Graft manipulation ex-vivo of	this product including	g T-cell depletion
other than for RBC removal o		с ,
No No		
Yes Negative	e: 🗌 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 🗌 Ye	
i ositive.		CD34+ enrichment
Constic	nanipulation	── No □ Yes

Donor 1 - Product Number 2			
more than one stem cell product, this is the SECO	ND product infused from this donor		
Source of Stem Cells for this product , select only	one		
Bone marrow	eripheral blood		
Cord blood Other:			
Graft manipulation ex-vivo of this product includin	g T-cell depletion		
other than for RBC removal or volume reduction			
No			
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 🗌 Ye	es		
	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 (DC	DNOR RELATION WITH PATIENT)		
Syngeneic	ntical sibling (may include non-mon (monozygotic twin) Inched other relative matched relative Degree of mismat		
HLA MISMATCHES (Mismatched relatives of	BETWEEN DONOR AND PATIENT		
Complete num	ber of mismatches inside each box		
A	B C DRB1 DQB1 DPB1		
		Antigenic	
		Allelic	
0=match; 1=one misma	tch; 2=2 mismatches; N/E=not evaluated		
Unrelated de	onor		
BMDW code of the Name of Donor Reg		ode is unknown) (up to 4 characters) odes is unknown)	
	centre name (if applicable, optional)		
Patient	D given by the Donor Registry or the CB I ID given by the Donor Registry or the CE Please enter the LABORATORY RESULTS	Bank listed above	
Donor information			
Date of birth _	yyyy - mm - dd	<u>OR</u> Age at time of donation (if da	
Donor Sex (o	nt birth) 🗌 Male 🗌 Fema		
Donor CMV status	Negative Posit	ive 🗌 Not evaluated] Unknown
Did this donor provide	more than one stem cell product		
No Yes: I	(please fill "Donor 1 – Product Numi Number of different stem cell products ir		
_	(If 2 products e.g. BM PB, please fill "Do	nor 1 – Product Number 1 AND 2" on i	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 No
Yes Negative: No Yes:
T-cell (CD3+) depletion (do not use for "Campathbag")
\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

Please enter the LABORATORY RESULTS WITH HLA TYPING into the database

Donor 2 - Product Number 2			
more than one stem cell product, this is the SECOND pro	oduct 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:	 <i>T-cell (CD3+) depletion (do not use for "Campathbag")</i> <i>T-cell receptor αβ depletion</i> 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

CIC: Hospital UPN:	Patient UIC	HSCT Date:	yyyy - 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 If >1 and Allograft, Was the same donor used for all If >1, was last HSCT peformed at another institution?					
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					
Preparative Regimen					
Preparative (conditioning) regimen given? No (Usually Paed Inherited Disorders only) Go to C Yes	GvHD Prophylaxis				
Was this intended to be myeloablative? (allo only)	 Age of recipient Comorbid conditions Prior HSCT Protocol driven Other, specify 				
Drugs 🗌 No 🗌 Yes	Unknown				
(include any active agent be it chemo, monoclonal antibody, polyclonal 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) Animal origin: Horse Rabbit Other specify		mg/m2	mg/kg		
Other, specify		mg/m2	mg/kg		
Busulfan Oral IV Both		mg/m2	mg/kg	mg x hr/L micromol x min/L mg x min/mL	
BCNU		mg/m2	🗌 mg/kg		
Bexxar (radio labelled MoAB)		🗌 mCi	🗌 MBq		
ССЛИ		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		
		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 Specify		🗌 mCi	🗌 MBq		
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

CIC:	Hospital UPN:	Patient UIC	HSCT Date:	vvvv - mm - dd
Total Body Irradiation (TBI)	🗌 No 🗌 Yes	: Total prescribed radiation dose		
, , ,		umber of fractions		-
TLI, TNI, TAI	🗌 No 🗌 Yes	: Total prescribed radiation dos	se as per protocol	Gy
(lymphoid, nodal, abdominal)				
GvHD prophylaxis or pre-	ventive treatment	'Allografts only)		
No Yes				
If Yes: 🗌 Drugs (Immund	osuppressive chemo)			
ALG, ALS Anti CD2 Campatl Systemic Cyclospe Cycloph Etanerce FK 506 Inflixima Methotr Sirolimu Other m	S, ATG, ATS : (given after 25(MoAB in vivo) n (MoAB in vivo; can be " c corticosteroids orine osphamide (given after ept (MoAB in vivo) (Tacrolimus, Prograf) ab (MoAB in vivo) rexate enolate (MMF) s	in the bag") day 0) vivo) , specify	e 🗌 Rabbit 🗍 Other,	specify
	photopheresis (ECP)			
Other, specify				
		Survival Status		
Survival Status on date o	AUCCT			
 Alive De Patient died between Main Cause of Dea Relapse or Progr HSCT Related Ca Unknown Other 	rad administration of the pre th <i>(check only one n</i> ession/Persistent disease use	·····		
GVHD	ry Cause of Death (check as many as appropriate):		
 Interstiti Pulmona Infectior baa vira fun par Un Rejectio History of Haemor Cardiac Gastroin Skin toxi Renal fa Multiple 	tterial al gal asitic known n/Poor graft function of severe Veno occlusive o rhage toxicity nervous system (CNS) tox itestinal (GI) toxicity city	icity		