		Patient UIC	HSCT Date: yyyy - mm - dd
	HSCT - Min	imum Essential I	
		Centre Identification	
	Unit:	<del></del>	
		Patient Data	
Patient following natio  No Yes: Na  Hospital Unique Patien Compulsory, registrations All transplants performed the patient and not to the	e transplant.	ut this item. e registered with the same patient identific	Jnknown
_	(first name(s) _	_	
Date of birth:	y - mm - dd	Sex:	☐ Female
	Prir	mary Disease Diagnosis	
	S:  yyyy - mm - dd  GNOSIS (CHECK THE DISEAS	EE FOR WHICH THIS TRANSPLANT WAS PERFO	ORMED)
related Precurs Precursor Lymp Therapy related n Secondary Acute Chronic Leukaem Chronic Myeloi	ohoid Neoplasms (old ALL) nyeloid neoplasms (old Leukaemia) ia d Leukaemia (CML) ocytic Leukaemia (CLL)	<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: yyyy - mm - dd
SOLID TUMOURS (main disease code 5)
Disease
Date of initial diagnosis
Bone sarcoma (excluding Ewing sarcoma/PNET)  Breast  Central nervous system tumours (include CNS PNET)  Colorectal  Ewing sarcoma (ES)/PNET, extra-skeletal  Ewing sarcoma(ES)/PNET, skeletal  Germ cell tumour, extragonadal only  Head and neck  Hepatobiliary  Kidney cancer excluding Wilm's tumour  Lung cancer, non-small cell  Lung cancer, small cell  Medulloblastoma  Medulloblastoma  Medulloblastoma  Medulloblastoma  Medulloblastoma  Melanoma  Melanom  Neuroblastoma  Melanoma  Melanom  Melanom  Melanom
TNM classification  Type: Clinical Pathological  0 1 2 3 4 X Not evaluated Unknown  Tumour Clinical Clinical Pathological  Nodes  Metastases* Clinical Pathological Clinical C
Breast carcinoma only
Receptor status:  Estrogen (ER):
Germ cell tumours only
Histological classification  Seminoma Non-seminoma  Site of origin  Gonadal Extragonadal: retroperitoneal mediastinal Other sites specify:

CIC:	Hospital UPN:	Patient UIC		HSCT Date:	yyyy - mm - dd
	SOLID TU	JMOURS (main	disease code 5		.,,,
		Status At HS	SCT		
Date of this HSCT:	yyyy - mm - dd				
Germ cell tumours					
Risk category at dise	ase recurrence (or platinu	m refractoriness) fo	llowing first line CT		
☐ Very low ☐	Low Intermediate	High	☐ Very High	Not evaluated	
	e/no response  nission (CR)  firmed (CRU*)  sponse with persistent scan abnorm  med	nalities of unknown	NUMBER  1st 2nd 3rd or higher  NUMBER  1st 2nd 3rd or higher	SENSITIVITY TO CHI Sensitive Resistan Untreate	e t
Progressive d	isease (PD)				
Organs involved (co	mplete only if not in CR)				
☐ Nodes ☐ CNS ☐ Liver ☐ Other, specify:			Bone Lung Soft Tissue		

CIC: Hos	pital UPN:	Patient UIC	HSCT Date:	yyyy -	mm - d	d
		HSCT		,,,,		
Performance score  Score		, 50	30 <sup>□</sup> 90 <sup>□</sup>	□ 100	ı	
	Como	rbidity Index				
orror et al., Blood, 2005 Oct 15;	106(8): 2912-2919: http://ww	ww.ncbi.nlm.nih.gov/pmc/articles,	/PMC1895304/			
Vas there any <i>clinically significan</i> preparative regimen?  No Yes	<b>nt</b> co-existing disease or organ	impairment at time of patient ass	essment just prior	to the		
Comorbidity		Definitions		No	Yes	N/E
Solid tumour, previously present	melanoma skin cancer	the patient's past history, excludir	ng non-			
nfammatan, hawal disaasa	Indicate type					
nfammatory bowel disease	Crohn's disease or ulcerative					
Rheumatologic	SLE, RA, polymyositis, mixed	CTD, or polymyalgia rheumatica		Ш		
nfection	Requiring continuation of ar	ntimicrobial treatment after day 0				
Diabetes	Requiring treatment with indiet alone	sulin or oral hypoglycaemics but n	ot			
Renal: moderate/severe	Serum creatinine > 2 mg/dL transplantation	or >177 μmol/L, on dialysis, or pri	or renal			
Hepatic: mild moderate/ severe	ULN, or AST/ALT between U	etween Upper Limit Normal (ULN LN and 2.5 × ULN ter than 1.5 × ULN, or AST/ALT gre	-			
Arrhythmia		ick sinus syndrome, or ventricular				
Cardiac	Coronary artery disease, cor 50%, or shortening fraction	ngestive heart failure, myocardial i in children (<28%)	infarction, EF ≤			
Cerebrovascular disease	Transient ischemic attack or	cerebrovascular accident				
Heart valve disease	Except mitral valve prolapse	2				
Pulmonary: moderate	DLco and/or FEV1 66-80% o	r dyspnoea on slight activity				
severe	DLco and/or FEV1 ≤ 65% or	dyspnoea at rest or requiring oxyg	gen			
Dbesity	Patients with a body mass in	ndex > 35 kg/m2				
Peptic ulcer	Requiring treatment					
Psychiatric disturbance	Depression or anxiety requir	ring psychiatric consultation or tre	atment			
				II.		

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 H	SCT (Alloge	neic)	
		1,60011	301 (/ moge		
☐ Allogeneic  Patient CMV statu	ıs	gative	ve □ Not eval	luated	n
				luated Unknow	II
Multiple donors (including multiple	CB units)	☐ Yes:	Number of donors		
			Donor 1		
HLA MATCH TYPE (DOI  HLA - Identical sibli  Syngeneic (monozy  HLA - Matched oth  HLA - Mismatched	ng <i>(may include non gotic twin)</i> er relative			ıs mismatch oci mismatch	
Donor ID given	by the centre				
<b>HLA</b> MISMATCHES (Mismatched relatives of	BETWEEN DONOR A	ND PATIENT			
Complete num	ber of mismatches i	nside each box			
A E	C DRB1	DQB1 DPB1			
0=match; 1=one misma	tch; 2=2 mismatches; N	/E=not evaluated	Antigenic Allelic		
Unrelated donor					
ION code of the Donor R	• ,				
BMDW code of the Dono Name of Donor Registry/		c (If ION code is of the above codes is	unknown) (up to 4 ch		
Donor centr	.,	able, optional)	. unknown,		
	(i) applie		CB Bank listed above		
			e CB Bank listed above		
Plea	se enter the LABORA	TORY RESULTS WITH	H HLA TYPING into the		
Donor information					
Date of birth	- mm - dd	<u>OR</u> Age	e at time of donation	(if date of birth not pr	
Donor Sex	(at birth)	Male	Female		(0)
Donor CM\	/ status	Negative	Positive	☐ Not evaluated	Unknown
Did this donor provide mor	e than one stem cell	product		_	
	lease fill "Donor 1 ber of different sten				
		•	1 – Product Number 1	AND 2" on next page)	

CIC:	Hospital UPN:	Patient UIC	HSCT Date:	уууу - mm - dd
	Donor	1 - Product Numb	er 1	
If	more than one stem cell product, this is the FIRST prod	uct infused from this donor		
	Source of Stem Cells for <b>this product</b> , select only <b>one</b>			
	Bone marrow Periphe  Cord blood Other:	eral blood		
	Graft manipulation ex-vivo of this product including T-conther than for RBC removal or volume reduction  No	ell depletion		
	☐ Yes Negative: ☐ No ☐ Yes	T-cell (CD3+) depletion (do  T-cell receptor αβ depletion  B-cell depletion (CD19+) by  NK cell depletion by MoAB		
	Positive: No Yes	CD34+ enrichment		
	Genetic manipulation	☐ No ☐ Yes		
	Please enter the LABORATORY RESULTS V	VITH HLA TYPING into the o		
If	more than one stem cell product, this is the SECOND p			
	Cord blood Other:	eral blood		
	Graft manipulation ex-vivo of this product including T-content than for RBC removal or volume reduction  No			
	Yes Negative: No Yes	T-cell (CD3+) depletion (do T-cell receptor αβ depletion B-cell depletion (CD19+) by  NK cell depletion by MoAB		
	Positive: No Yes	CD34+ enrichment		
	Genetic manipulation	☐ No ☐ Yes		

Please enter the LABORATORY RESULTS WITH HLA TYPING into the database

CIC:	Ho	spital UPN:		Patient UIC	HSCT Date:	mm - c
				Donor 2	уууу-	mm - u
HLA MATCH TYPE (	DONOR RELAT	TION WITH PATIEN	NT)			
☐ HLA - Id	lentical sibling	(may includ	le non-monozygo	otic twin)		
Syngene	eic <i>(mond</i>	zygotic twin)	, 5	ŕ		
☐ HLA - N	latched other	relative				
☐ HLA - N	lismatched re	ative Degre	e of mismatch	☐ 1 HLA locus misma ☐ >=2 HLA loci mism		
HLA MISMATCH (Mismatched relative		DONOR AND PATI	ENT			
Complete nu	umber of misr	natches inside ea	ch box			
Α	В С	DRB1 DQB1	DPB1			
	пг			dt-		
	片는		Ant	igenic		
			Alle	elic		
0=match; 1=one miss	match; 2=2 misi	matches; N/E=not ev	valuated			
Unrelated	donor					
ION code of the I	Donor Registry	or CB Bank				
BMDW code of t	_	-			acters)	
Name of Donor R	Registry/ CB Ba			is unknown)		
Dono	r centre name	e (if applicable	e, optional)			
Donor	ID given by t	he Donor Registry	or the CB Bank	listed above		
Patien	t ID given by	the Donor Regist	ry or the CB Ban	k listed above		
	Please ento	er the LABORATO	RY RESULTS WIT	TH HLA TYPING into the da	tabase	
Donor information	1					
Date of birth		mm - dd	<u>OR</u>		(if date of birth not provided)	
Donor Sex	(at birth)	☐ Male	Female	yea	nr(s)month(s)	
Donor CMV status		Negative	Positive	Not evaluated	Unknown	
Did this donor provid	de more than	one stem cell pro	duct			
☐ No	(please fil	l "Donor 1 – Pro	duct Number 1	!" on next page		
Yes				d from this donor		
	(If 2 produc	ts e.g. BM PB, ple	ease fill "Donor 1	l – Product Number 1 AND	2" on next page)	

	Hospital UPN:		Patient UIC	HSCT Date:	yyyy - mm - dd
					yyyy - mm - uu
		Donor 2	- Product Number	er 1	
f more than one s	tem cell product, this is	the FIRST product	infused from this donor		
	ells for this product, sel				
Bone marro	ow Peripher	al blood			
☐ Cord blood		!			
Graft manipulatio	n ex-vivo including T-Ce	ll depletion			
	C removal or volume red				
☐ No					
Yes	Negative:	No Yes:			
			T-cell (CD3+) depletion (do r T-cell receptor αβ depletion		
			B-cell depletion (CD19+) by		
			NK cell depletion by MoAB		
			Other		
F	Positive: No	Yes			
			CD34+ enrichment		
	netic manipulation er the LABORATOR	□ No Y RESULTS WIT	☐ Yes H HLA TYPING into the d	latabase	
		Y RESULTS WIT	H HLA TYPING into the d		
Please ent	er the LABORATOR	y results wit	H HLA TYPING into the d		
Please ent	er the LABORATOR'	PRESULTS WIT  Donor 2  the SECOND produ	H HLA TYPING into the d		
Please ent	er the LABORATOR	PRESULTS WIT  Donor 2  the SECOND produ	H HLA TYPING into the d		
Please ent	er the LABORATOR'  tem cell product, this is  ells for this product, sel	PY RESULTS WIT  Donor 2  the SECOND producect only one	H HLA TYPING into the d  - Product Number		
Please ent  f more than one st  Source of Stem Co	er the LABORATOR'  tem cell product, this is  ells for this product, sel	PRESULTS WIT  Donor 2  the SECOND produ	H HLA TYPING into the d  - Product Number		
Please ent  f more than one st  Source of Stem Co  Bone marro  Cord blood	er the LABORATOR'  tem cell product, this is  ells for this product, sel	PY RESULTS WIT  Donor 2  the SECOND producect only one ral blood	H HLA TYPING into the d  - Product Number		
Please ent  f more than one st  Source of Stem Co  Bone marro  Cord blood  Graft manipulatio  other than for RBo	tem cell product, this is  ells for this product, sel  ow Peripher  Other source	PY RESULTS WIT  Donor 2  the SECOND producet only one ral blood	H HLA TYPING into the d  - Product Number		
Please ent  f more than one st  Source of Stem Co  Bone marro  Cord blood  Graft manipulatio  other than for RBo	er the LABORATOR'  tem cell product, this is  ells for this product, sel  w Peripher  Other source  n ex-vivo including T-Ce	PY RESULTS WIT  Donor 2  the SECOND production  all depletion  duction	H HLA TYPING into the d  - Product Number		
Please ent  f more than one st  Source of Stem Co  Bone marro  Cord blood  Graft manipulatio  other than for RBo	er the LABORATOR'  tem cell product, this is  ells for this product, sel  ow Peripher  Other source  n ex-vivo including T-Ce  C removal or volume red	PY RESULTS WIT  Donor 2  the SECOND producet only one ral blood	H HLA TYPING into the d  - Product Number	er 2	
Figure 2 Please ent  Figure 1 Please ent  Figure 2 Please ent  Figure 2 Please ent  Figure 3 Please ent  Figure 3 Please ent  Figure 4 Please ent  Figure 5 Please ent  Figure 4 Please ent  Figure 4 Please ent  Figure 5 Please ent  Figure 4	er the LABORATOR'  tem cell product, this is  ells for this product, sel  w Peripher  Other source  n ex-vivo including T-Ce	PY RESULTS WIT  Donor 2  the SECOND production  all depletion  duction	- Product Number of the description of the descrip	er 2  not use for "Campathbag")	
Figure 2 Please ent  Figure 1 Please ent  Figure 2 Please ent  Figure 2 Please ent  Figure 3 Please ent  Figure 3 Please ent  Figure 4	er the LABORATOR'  tem cell product, this is  ells for this product, sel  w Peripher  Other source  n ex-vivo including T-Ce	PY RESULTS WIT  Donor 2  the SECOND production  all depletion  duction	- Product Number of the description of the descrip	er 2  not use for "Campathbag")	
Figure 2 Please ent  Figure 1 Please ent  Figure 2 Please ent  Figure 2 Please ent  Figure 3 Please ent  Figure 3 Please ent  Figure 4	er the LABORATOR'  tem cell product, this is  ells for this product, sel  w Peripher  Other source  n ex-vivo including T-Ce	PY RESULTS WIT  Donor 2  the SECOND production  all depletion  duction	- Product Number of the description of the descrip	er 2  not use for "Campathbag")	
Please ent  f more than one st  Source of Stem Co  Bone marro  Cord blood  Graft manipulatio  other than for RBo  No  Yes	er the LABORATOR'  tem cell product, this is  ells for this product, sel  w Peripher  Other source  n ex-vivo including T-Ce  C removal or volume rec  Negative:	The SECOND production  The Second production production  The Second production production production  The Second production	- Product Number of the description of the descrip	not use for "Campathbag") MOAB	
Please ent  f more than one st  Source of Stem Co  Bone marro  Cord blood  Graft manipulatio  other than for RBo  No  Yes	er the LABORATOR'  tem cell product, this is  ells for this product, sel  w Peripher  Other source  n ex-vivo including T-Ce  C removal or volume rec  Negative:	PY RESULTS WIT  Donor 2  the SECOND production  all depletion  duction	- Product Number of the description of the descrip	not use for "Campathbag") MOAB	
Figure 2 Please ent  Figure of Stem Co  Bone marro  Cord blood  Graft manipulatio  other than for RBO  Yes	er the LABORATOR'  tem cell product, this is  ells for this product, sel  w Peripher  Other source  n ex-vivo including T-Ce  C removal or volume rec  Negative:	The SECOND production  The Second production production  The Second production production production  The Second production	T-cell (CD3+) depletion (do note that the depletion (CD19+) by NK cell depletion by MoAB Other	not use for "Campathbag") MOAB	

Please enter the LABORATORY RESULTS WITH HLA TYPING into the database

CIC:	Hospital UPN:	Patient UIC	HSCT Date:	
		HSCT (Continued)		
If >1, date of If >1, type of If >1 and A	r of HSCT for this patient?   of last HSCT before this one of last HSCT before this one [ llograft, Was the same donor used fo ast HSCT peformed at another institu			
subseque	nt transplant as the date of last o	rm before proceeding, giving the contact other events between transplants		
HSCT part of a plar	nned multiple (sequential) graft p Yes	orotocol (program)?		
	Р	reparative Regimen		
	ioning) <b>regimen given?</b> ally Paed Inherited Disorders only) Go	o to GvHD Prophylaxis		
Was this intended  Yes	to be myeloablative? (allo only)  No: Reason	Age of recipient Comorbid conditions Prior HSCT Protocol driven		
Drugs (include any active ag	☐ No ☐ Yes gent be it chemo, monoclonal antiboo	☐ Unknown dy, polyclonal antibody, serotherapy, €	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)		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	

<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 or bea	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	