CIC:	Hospital UPN:	Patient UIC	HSCT Date: yyyy - mm - dd			
	HSCT - Min	imum Essential REGISTRATION - DAY 0				
		Centre Identification				
	Unit:	_				
		Patient Data				
Date of this report: First transplant for this patient?: Yes No  yyyy - mm - dd  Patient following national / international study / trial:  No Yes: Name of study / trial Unknown  Hospital Unique Patient Number or Code (UPN)  Compulsory, registrations will not be accepted without this item.  All transplants performed in the same patient must be registered with the same patient identification number or code as this belongs to the patient and not to the transplant.  Initials: (first name(s) _family name(s))						
Date of birth:	yyyy - mm - dd	Sex: Male	Female			
	Prir					
Primary Disease Diagnosis  Date of initial diagnosis:						
Other diagnosis, specify:						

CIC:	Hospital UPN:	Patient UIC		HSCT Date:				
	ACUTE LEUKAEMIAS (main disease code 1)  Acute Myeloid leukaemia (AML) (1 of 4)							
		Disease						
Date of Initial Diagnos	yyyy - mm - dd							
AML with inv(16)(p: Acute promyelocyti AML with t(9;11) (p AML with t(6;9) (p2 AML with inv(3) (q2 AML (megakaryobla AML with myelodys Was there a previ No > Co. Yes > Fill Predisposing Co AML not otherwise cate AML with minimal co AML with maturation AML with maturation	netic abnormalities 22;q22); RUNX1-RUNX1T1 13.1;q22) or t(16;16)(p13.1;q22); c leukaemia with t(15;17)(q22;q1:22;q23); MLLT3-MLL 3;q24); DEK-NUP214 21;q26.2) or t(3;3) (q21;q26.2); RPiastic) with t(1;22) (p13;q13); RBM eplasia related changes (old "cous diagnosis of MDS or MDS/MPintinue to Predisposing condition below to the MYELODYPLASTIC SYNDROmodition below eggorised (NOS) differentiation (FAB M0) tration (FAB M1)	2); PML/RARA N1-EVI1 15-MKL1 Acute leukaemia transforme N? v ME (MDS) or MDS/MPN						
Acute erythroid leul Acute megakaryobli Acute basophilic leu Acute panmyelosis Myeloid sarcoma (6 Myeloid proliferatio Blastic plasmacytoid Therapy related my	kaemia (FAB M6) astic leukaemia (FAB M7) ukaemia	cute Leukaemia'')	DS/MPN.					
	Pred	disposing Cor	ndition?					
Skip this question if the	AML is a Therapy related neoplasi	ia						
Did the recipient have prior to the diagnosis	e a predisposing condition of leukaemia?	□ No	Fa	olastic anaemia Inconi anaemia oom syndrome nknown				
Donor Cell Leukaemia?								
IF THE PATIENT HAS RECEIVED AN ALLOGRAFT PRIOR TO THE DIAGNOSIS OF ACUTE LEUKAEMIA, ANSWER THE FOLLOWING QUESTION  Is this a donor cell leukaemia								

CIC:	Hospital UPN:	Patient UIC		HSCT D	oate:	- mm - dd
	A CLITE I ELL	VAENIAC (main diaa	222 224	o 1\	уууу	- mm - aa
		KAEMIAS (main dise		•		
		eloid leukaemia (AML	, ,	,		
	Chromoso	ome Analysis at [	Diagno	sis		
Chromosome analysis a	t diagnosis (All methods	including FISH)				
Done: normal	Done: abnormal	Not done or failed	Unkr	nown		
If abnormal: <b>Com</b>		☐ No ☐ Yes		Unknown		
·	nore abnormalities) nosomal karyotype:	□ No □ Yes		Unknown		
		autosomal monosomy + at least	1 structural a	abnormality	)	
		,				
		OR				
Indicate helow those ahn	ormalities that have he	een <b>evaluated</b> and whethe	r they wer	e <b>Ahsent</b>	or <b>Preser</b>	ıt.
t(15;17)	ormandes that have be	en cvalated and whethe	i	bsent	Present	Not evaluated
t(8;21)				bsent $\Box$	Present [	Not evaluated  Not evaluated
inv(16)/ t(16;16)				bsent	Present	Not evaluated
11q23 abnormality type				bsent	Present [	Not evaluated
Fill only if 11q23 abnormali	ity is Present:					
t(9;11)			A	bsent	Present [	Not evaluated
t(11;19)			A	bsent	Present [	Not evaluated
t(10;11)			A	bsent	Present	Not evaluated
t(6;11)			A	bsent	Present	Not evaluated
Other abn(11q23), speci	fy:					
3q26 (EVI1) abnormality typ			A	bsent	Present [	Not evaluated
Fill only if 3q26 (EVI1) abno	ormality is Present:					
inv(3)/ t(3;3)				bsent	Present	Not evaluated
t(2;3)(p21;q26) Other t(3q26)/EVI1 reari	rangement specify:			lbsent	Present Present	Not evaluated  Not evaluated
t(6;9)	angement, specify			bsent	Present [	Not evaluated  Not evaluated
abn 5 type				bsent	Present	Not evaluated
Fill only if above abn 5 is Pr	resent:					
del (5q)			A	bsent	Present [	Not evaluated
monosomy 5			A	bsent 🗌	Present [	Not evaluated
add(5q)			A	bsent 🗌	Present [	Not evaluated
Other abn(5q); please sp	ecify:		A	bsent 🗌	Present [	Not evaluated
abn 7 type			A	bsent	Present	Not evaluated
Fill only if abn 7 is Present:						
del(7q)			A	bsent	Present	Not evaluated
monosomy 7				bsent	Present [	Not evaluated
add(7q)				bsent	Present _	Not evaluated
Other abn(7q); please sp	ecify:			bsent	Present	Not evaluated
-17				bsent	Present [	Not evaluated
abn(17p)				bsent	Present [	Not evaluated
t(1;22)				bsent	Present [	Not evaluated
trisomy 8			A	bsent	Present	Not evaluated

Other, specify.....

Absent

Present

CIC:	Hospital UPN: Patient UIC	HSCT Date:						
	ACUTE LEUKAEMIAS (main disease code 1)  Primary Acute Myeloid leukaemia (AML) (3 of 4)							
	Molecular Markers at Diagno	osis						
Mole	ecular marker analysis at diagnosis							
	☐ Not evaluated ☐ Evaluated: absent ☐ Evaluated present	Unknown						
	Indicate below those abnormalities that have been evaluated and whether the	ney were Absent or Present						
	AML1-ETO (RUNX1/RUNXT1)  Molecular product of t(8;21)	☐ Absent ☐ Present ☐ Not evaluated						
	CBFB-MYH11 Molecular product of inv(16)(p13.1;q22) or (16;16)(p13.1;q22)	☐ Absent ☐ Present ☐ Not evaluated						
	PML-RARα Molecular product of t(15;17)	☐ Absent ☐ Present ☐ Not evaluated						
Ī	MLL-rearrangement/mutation:	Evaluated at Not evaluated least once						
-	Fill only if 11q23 abnormality is Present:  MLLT3(AF9)-MLL  molecular product of t(9;11)(p22;q23)	☐ Absent ☐ Present ☐ Not evaluated						
-	MLL-PTD (partial tandem duplication)	Absent Present Not evaluated						
-	MLLT4(AF6)-MLL molecular product of t(6;11)(q27;q23)	☐ Absent ☐ Present ☐ Not evaluated						
-	ELL-MLL:  molecular product of t(11;19)(q23;p13.1)	☐ Absent ☐ Present ☐ Not evaluated						
-	MLLT1(ENL)-MLL: molecular product of t(11;19)(q23;p13.3)	☐ Absent ☐ Present ☐ Not evaluated						
-	MLLT10(AF10)-MLL:  molecular product of t(10;11)(p12;q23)	☐ Absent ☐ Present ☐ Not evaluated						
-	Other MLL-rearrangement, specify:	Absent Present Not evaluated						
	DEK-NUP214(CAN) molecular product of translocation t(6;9)(p23;q34)	☐ Absent ☐ Present ☐ Not evaluated						
Ī	RPN1-EVI1  molecular product of inv(3)(q21q26.2) or t(3;3)(q21q26.2)	☐ Absent ☐ Present ☐ Not evaluated						
	RBM15-MKL1  molecular product of translocation t(1;22)(p13;q13)	☐ Absent ☐ Present ☐ Not evaluated						
F	NPM1 mutation	Absent Present Not evaluated						
	CEBPA mutation	☐ Absent ☐ Present ☐ Not evaluated						
	FLT3-ITD (internal tandem duplication)	Absent Present Not evaluated						
	DNMT3A	☐ Absent ☐ Present ☐ Not evaluated						
	ASXL1	Absent Present Not evaluated						
	TP53	☐ Absent ☐ Present ☐ Not evaluated						
	RUNX1	Absent Present Not evaluated						
	c-KIT	Absent Present Not evaluated						
	Other, specify	Absent Present Not evaluated						
	Involvement at Diagnosis							
Invo	Involvement at diagnosis							
	Bone marrow No Yes Not evaluated							
(	CNS No Yes Not evaluated							
T	Testis/ovary							
(	Other No Yes, specify							
	Page 4 AAAL Da	WARD A FORM						

CIC:	Hospital UPN:	Patient U	IIC	HSCT Date:			
ACUTE LEUKAEMIAS(main disease code 1) Primary Acute Myeloid leukaemia (AML) (4 of 4)							
		Status at I	HSCT				
Date of this HSCT:	yyyy - mm - dd						
STATUS		NUMBER	TYPE OF REMISSION				
Primary induction	n failure						
☐ Complete haema	tological remission (CR)	☐ 1st ☐ 2nd ☐ 3rd or higher	CYTOGENETICS REMISSION  No Yes Not Evaluated Not Applicable* Unknown	MOLECULAR REMISSION  No Yes Not Evaluated Not Applicable* Unknown			
Relapse		☐ 1st☐ 2nd☐ 3rd or higher					
	ed prior to this time point e before this HSCT:	yyyy - mm - dd					

CIC: Hosp	pital UPN:	Patient UIC	HSCT Date:	уууу -	mm - d	'd			
HSCT									
Performance score         system used									
	Como	rbidity Index							
forror et al., Blood, 2005 Oct 15;	106(8): 2912-2919: http://w	ww.ncbi.nlm.nih.gov/pmc/	articles/PMC1895304/						
Vas there any <i>clinically significar</i> oreparative regimen?  No Yes	<b>nt</b> co-existing disease or organ	impairment at time of pat	ient assessment just prior	to the					
Comorbidity	ι	Definitions		No	Yes	N/E			
Solid tumour, previously present	Treated at any time point in melanoma skin cancer Indicate type		excluding non-						
nfammatory bowel disease	Crohn's disease or ulcerative								
Rheumatologic	SLE, RA, polymyositis, mixed CTD, or polymyalgia rheumatica								
nfection	Requiring continuation of ar	ntimicrobial treatment afte	er day 0						
Diabetes	Requiring treatment with indiet alone	sulin or oral hypoglycaemio	cs but not						
Renal: moderate/severe	Serum creatinine > 2 mg/dL transplantation	or >177 μmol/L, on dialysis	s, or prior renal						
Hepatic: mild moderate/ severe	Chronic hepatitis, bilirubin button button, or AST/ALT between U Liver cirrhosis, bilirubin greative vuln	LN and 2.5 × ULN							
Arrhythmia	Atrial fibrillation or flutter, s arrhythmias	ick sinus syndrome, or ven	tricular						
Cardiac	Coronary artery disease, cor 50%, or shortening fraction	=	cardial 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 activit	У						
severe	DLco and/or FEV1 ≤ 65% or	dyspnoea at rest or requiri	ng oxygen						
Obesity	Patients with a body mass in	ndex > 35 kg/m2							
Peptic ulcer	Requiring treatment								
Psychiatric disturbance	Depression or anxiety requi	ring psychiatric consultatio	n or treatment						
Managahan ang akhan masa 1914		26	.c.	1					

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

CIC:	Hospital UPN	J:	Patient UIC	HSCT	Date:
					yyyy - mm - dd
		Type of	HSCT (Alloge	eneic)	
☐ Allogeneic					
Patient CMV st	atus 🔲 [	Negative Po	sitive Not eva	lluated Unknow	vn
Multiple donor (including multi		No Ye	S: Number of donors		
			Donor 1		
HLA MATCH TYPE (E	ONOR RELATION W	ITH PATIENT)			
	ibling <i>(may include r</i> ozygotic twin) other relative	non-monozygotic tw.  Degree of mis	match 1 HLA loca	us mismatch oci mismatch	
Donor ID give	en by the centre				
<b>HLA</b> MISMATCH (Mismatched relative	ES BETWEEN DONO	R AND PATIENT			
Complete nu	umber of mismatche	es inside each box			
Α	B C DR	B1 DQB1 DPB1			
0=match; 1=one miss	match; 2=2 mismatches	s; N/E=not evaluated	Antigenic Allelic		
Unrelated dono	r				
ION code of the Dono	<i>G</i> ,				
BMDW code of the Do			e is unknown) (up to 4 cl	haracters)	
Name of Donor Regist		ny of the above code	es is unknown)		
		olicable, optional)	he CB Bank listed above		
			the CB Bank listed above /ITH HLA TYPING into the		
	case effer the LABO	MATORT RESOLIS W	minited into the	database	
<b>Donor information</b> Date of birth	yy - mm - dd	<u>OR</u>	Age at time of donation	(if date of birth not p	provided)
		Male	Fomolo	mc	onth(s)
Donor S	ex (at birth)  MV status		☐ Female☐ Positive	☐ Not evaluated	□ Unknown
Did this donor provide m		Negative	Positive	Not evaluated	
_		-	har 1" on novt nace		
			ber 1" on next page fused from this donor		
			nor 1 – Product Number 1	AND 2" on next page)	

If more than one stem cell product, this is the FIRST product infused from this donor	CIC:	Hospital UPN:	Patient UIC	HSCT Date:	yyyy - mm - dd
Source of Stem Cells for this product, select only one   Bone marrow		Dono	r 1 - Product Numb	er 1	
Bone marrow	If more	than one stem cell product, this is the FIRST pro	duct infused from this donor		
Please enter the LABORATORY RESULTS WITH HLA TYPING into the database    Donor 1 - Product Number 2	Sourc	The of Stem Cells for this product, select only one  Bone marrow	eral blood  cell depletion  s:  T-cell (CD3+) depletion (do T-cell receptor αβ depletion B-cell depletion (CD19+) by  NK cell depletion by MoAB Other	n / MoAB	
Donor 1 - Product Number 2  If more than one stem cell product, this is the SECOND product infused from this donor    Source of Stem Cells for this product, select only one		Genetic manipulation	☐ No ☐ Yes		
Source of Stem Cells for this product , select only one  Bone marrow Peripheral blood Cord blood Other:  Graft manipulation ex-vivo of this product 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 "Campath in bag") T-cell receptor αβ depletion B-cell depletion (CD19+) by MoAB  NK cell depletion by MoAB Other  Positive: No Yes  CD34+ enrichment					
Bone marrow		<u></u>	product infused from this donor		
CD34+ enrichment	Graft	Bone marrow Periph  Cord blood Other:  manipulation ex-vivo of this product including T- than for RBC removal or volume reduction  No	cell depletion  s:    T-cell (CD3+) depletion (do   T-cell receptor αβ depletion   B-cell depletion (CD19+) by   NK cell depletion by MoAB	n / MoAB	
Genetic manipulation No Yes		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:	уууу - mm - a
			Onor 2		
HLA MATCH TYPE	(DONOR RELATION WITH PATIEN	IT)			
☐ HLA - Io	dentical sibling (may includ	e non-monozygo	tic twin)		
Syngen		, -	·		
HLA - N	Matched other relative				
HLA - N	Mismatched relative Degree	e of mismatch	☐ 1 HLA locus misma ☐ >=2 HLA loci misma		
HLA MISMATCH (Mismatched relativ	HES BETWEEN DONOR AND PATI les only)	ENT			
Complete n	umber of mismatches inside each	ch box			
Α	B C DRB1 DQB1	DPB1			
		Anti	genic		
		Allel	io		
0=match: 1=one mis	smatch; 2=2 mismatches; N/E=not ev		ic		
Unrelated					
ION code of the	Donor Registry or CB Bank				
	the Donor Registry or CB Bank			cters)	
Name of Donor I	Registry/ CB Bank (If any of t	he above codes i	s unknown)		
Dono	or centre name (if applicable				
Donor	· ID given by the Donor Registry	or the CB Bank I	sted above		
Patier	nt ID given by the Donor Registi	ry or the CB Bank	listed above		
	Please enter the LABORATO	RY RESULTS WIT	H HLA TYPING into the dat	abase	
Donor information	n				
Date of birtl	h yyyy - mm - dd	<u>OR</u>	Age at time of donation	(if date of birth not provided	יט
	<u></u>		yea	r(s)month(s	)
Donor Sex	(at birth) Male	Female			
Donor CMV status	Negative	Positive		Unknown	
Did this donor provi	de more than one stem cell prod	duct			
☐ No	(please fill "Donor 1 – Prodes: Number of different stem cell		·		
	(If 2 products e.g. BM PB, ple			2" on next page)	

If more than one stem cell product, this is the FIRST product Infused from this donor	CIC:	Hospital UPN:	Patient UIC	HSCT Date:	уууу - mm - dd			
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 thon for RBC removal or volume reduction   No   Yes   Negative:   No   Yes   CD34+ enrichment   Reduction   B. cell depletion (CD39+) by MoAB   No. cell depletion (CD39+) depletion (CD39+) depletion (CD39+) depletion (CD39+) depletion (CD39+) by MoAB   No. cell		Donor	2 - Product Numb	ar 1				
Source of Stem Cells for this product, select only one   Bone marrow				OI I				
Bone marrow   Peripheral blood   Cord blood   Other source			uct infused from this donor					
Cord blood   Other source   Graft manipulation ex-vivo including T-Cell depletion other than for RBC removal or volume reduction   No   Yes   T-cell (CD3+) depletion (do not use for "Compathbag")   T-cell receptor αβ depletion   No   Yes   Positive:   No   Yes   CD34+ enrichment   Positive:   No   Yes   T-cell receptor αβ depletion   No   Yes   Positive:   No   Yes   CD34+ enrichment   T-cell receptor αβ depletion   No   Yes   T-cell receptor αβ depletion   No   No   No   No   No   No   No	Source o	of Stem Cells for this product, select only one						
Graft manipulation ex-vivo including T-Cell depletion  other than for RBC removal or volume reduction  No								
other than far RBC removal or volume reduction   No		_						
No								
Yes   Negative:   No   Yes:	I							
T-cell receptor aß depletion   B-cell depletion (CD19+) by MoAB   Not cell depletion (CD19+) by MoAB   Not cell depletion by MoAB   Other   Nother   Nothe		<u></u>						
B-cell depletion (CD19+) by MoAB   NX cell depletion by MoAB   NX cell depletion by MoAB   Other   NX cell depletion   NX cell depletion by MoAB   Other   NX cell depletion   NX cell depletion by MoAB   Other   NX cell depletion   NX cell depletion by MoAB   Other   NX cell depletion   NX cell depletion by MoAB   Other   NX cell depletion   NX cell deple								
Other   Positive:   No   Yes   CD34+ enrichment   Genetic manipulation   No   Yes			_					
Positive:   No   Yes   CD34+ enrichment   Genetic manipulation   No   Yes								
CD34+ enrichment			Utner					
Please enter the LABORATORY RESULTS WITH HLA TYPING into the database    Donor 2 - Product Number 2		Positive: No Yes	CD241 oprichment					
Please enter the LABORATORY RESULTS WITH HLA TYPING into the database    Donor 2 - Product Number 2								
Donor 2 - Product Number 2  If more than one stem cell product, this is the SECOND product infused from this donor    Source of Stem Cells for this product, select only one   Bone marrow   Peripheral blood   Cord blood   Other source   Cord blood   Other source   Peripheral blood   Peripheral blood   Cord blood   Other source   Cord blood   Other source   Cord blood   Other source   Cord blood   Peripheral blood   Cord blood   Other source   Cord blood   Cord blood   Other source   Cord blood   Peripheral blood   Peri		Genetic manipulation No	☐ Yes					
If more than one stem cell product, this is the SECOND product infused from this donor    Source of Stem Cells for this product, select only one		Donor	2 - Product Numb	er 2				
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 αβ depletion B-cell depletion (CD19+) by MoAB NK cell depletion by MoAB Other  Positive: No Yes  CD34+ enrichment	If more th							
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 αβ depletion B-cell depletion (CD19+) by MoAB  NK cell depletion by MoAB  Other  Positive: No Yes  CD34+ enrichment	□ Во	one marrow Peripheral blood						
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 αβ depletion B-cell depletion (CD19+) by MoAB NK cell depletion by MoAB Other  Positive: No Yes  CD34+ enrichment	Graft ma	enipulation ex-vivo including T-Cell depletion						
CD34+ enrichment	other the	an for RBC removal or volume reduction	T-cell (CD3+) depletion (do  T-cell receptor αβ depletio  B-cell depletion (CD19+) by  NK cell depletion by MoAB	n / MoAB				
		Positive: No Yes						
Genetic manipulation No Yes			CD34+ enrichment					
		Genetic manipulation No	Yes					

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	o	aft protocol (program)?						
		Preparative Regimen						
Preparative (conditioning) regimen given?  No (Usually Paed Inherited Disorders only) Go to GvHD Prophylaxis  Yes								
Was thi		<ul><li>Age of recipient</li><li>Comorbid conditions</li><li>Prior HSCT</li><li>Protocol driven</li></ul>						

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			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		П	mg/m2	П	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	Г	mg/kg		
	Other, specify			mg/m2		mg/kg		

<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)	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	5, ATG, ATS: (gives) 5, ATG, ATS: (gives) 6, (MoAB in vivo) 6, corticosteroids 6, corticosteroids 6, corticosteroids 7, cortico	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		Gai 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) NS) toxicity	