CIC:	Hospital UPN:	Patient UIC	HSCT Date:				
	HSCT - Min	imum Essential C					
		Centre Identification					
	Unit:						
Patient Data							
□ No □ Yes	yyyy - mm - dd national / international study / s: Name of study / trial atient Number or Code (UPN) ations will not be accepted withour med in the same patient must be	U	nknown				
	(first name(s)	_family name(s)) Sex:	Female				
	Prir	mary Disease Diagnosis					
	nosis: yyyy - mm - dd DIAGNOSIS (CHECK THE DISEAS	 SE FOR WHICH THIS TRANSPLANT WAS PERFO	DRMED)				
related Pre Precursor Therapy relat Secondary Ac Chronic Leuk	elogenous Leukaemia (AML) ecursor Neoplasms Lymphoid Neoplasms (old ALL) ted myeloid neoplasms (old cute Leukaemia) aemia yeloid Leukaemia (CML) mphocytic Leukaemia (CLL) kin Disease	 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:	Pati	ent UIC		HSCT Date:	vvvv - mm - dd
	PLASMA CELL DISOR		JDING N	MULTIPLE		
		Disc	ease			
Date of Initial	Diagnosis:					
Classifica				HEAVY CH	IAIN TYPE I IG	GHT CHAIN TYPE
	nyeloma (MM)				IgG	Kappa
	heavy chain and light chain	Check light and h	neavy chain	$types \rightarrow \Box$	IgA	Lambda
	light chain	Check light chain	type only –	·	lgD	
	non-secretory				IgE IgM <i>(not Walder</i>	nstrom)
	ell leukaemia olasmacytoma of bone					,
	amyloidosis					
D POEMS	•					
_	nal light and heavy chain deposition ecify	disease (LCDD/H	CDD)			
	ing for Multiple myeloma only SALMON & DURIE STAGE			ISS	STAGE	
`	(optional)	-				
(F	PLEASE TICK EACH COLUMN)			β2-μς	nlob mg/L)	Albumin (g/L)
	Stage Symptoms			< 3.		>35
] I			< 3.	.5 OR	< 35
] II			3.5 - <	5.5	any
] 111			> 5.	5	any
	Chromosome Analy	sis at Diagno	osis (not	for Primary	y amyloidosi	s)
Chromosom	e analysis at diagnosis (All metho	ods including FISH))			
	Normal	☐ Abnormal		lot done or faile	d □ Unk	nown
If ab	normal:					
	Complex kariotype: (3 or more abnormalities)	No	Yes	∐ Unk	nown	
ou can transc	ribe the complete karyotype:					
	OR					
Indicate belo	ow those abnormalities that have b	een evaluated and	d whether t	hev were Abser	nt or Present	
	Del 13q14		Absent	Present	☐ Not evalua	ted
	t(11;14)		Absent	Present	☐ Not evalua	
	abn 17q		Absent	Present	☐ Not evalua	
	del 17p		Absent	Present	☐ Not evalua	ted
	t(4:14)		Absent	Present	Not evalua	ted
	t(14:16)		Absent	Present	☐ Not evalua	
	1q amplification		Absent	Present	☐ Not evalua	
	myc rearrangement Other, specify		Absent Absent	Present Present	Not evalua Not evalua	
	Molecular Marker			r Primary a		
larker anal	ysis at diagnosis			,	, ,	
Absent	_	Not Evaluated		Unknown		
		Page 2	<u>P</u>	CD_Day 0 Auto N	1ED-A Form	

CIC:	Hospital UPN:	Patient UIC	HSCT Date:	vyyy - mm - dd
PLASMA		INCLUDING MUL in disease code 4)	TIPLE MYELOMA (PC	D)
		Status At HSCT		
Date of this HSCT:	yyyy - mm - dd			
STATUS		NUMBER		
Never treated				
Stringent complete remiss	sion (sCR)	1st		
Complete remission (CR)				
Very good partial remission	on (VGPR)	2nd		
Partial remission (PR)		2nd on bishon		
Relapse from CR (untreate	ed)	3rd or higher		
Progression				
No change / stable disease	е			

CIC: Hosp	oital UPN: Patient UIC HS	CT Date:	уууу -	mm - d	d
	HSCT				
Performance score Score		□ 90 □	100		
Weight (Ng).	Tielgitt (elli).				
	Comorbidity Index				
forror et al., Blood, 2005 Oct 15;	106(8): 2912-2919: http://www.ncbi.nlm.nih.gov/pmc/articles/PMC	1895304/			
Vas there any <i>clinically significan</i> preparative regimen? No Yes	at co-existing disease or organ impairment at time of patient assessm	ent 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 nor melanoma skin cancer	า-			
	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 retransplantation	nal			
Hepatic: mild	Chronic hepatitis, bilirubin between Upper Limit Normal (ULN) and ULN, or AST/ALT between ULN and 2.5 × ULN				
moderate/ severe	Liver cirrhosis, bilirubin greater than 1.5 × ULN, or AST/ALT greater × ULN	tnan 2.5			
Arrhythmia	Atrial fibrillation or flutter, sick sinus syndrome, or ventricular arrhythmias				
Cardiac	Coronary artery disease, congestive heart failure, myocardial infarc 50%, or shortening fraction in children (<28%)	tion, 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 treatme	nt			

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

CIC:	Hospital UPN	I: Patient UIC		HSCT Date:	nm - dd
		Type of HSCT (Aut	ologous)		
	Autologous				
	Source of the Stem cells (check all that apply):	☐ Bone marrow☐ Cord blood	☐ Peripheral bl		
	Graft manipulation ex-vivo other than for RBC removal or	volume reduction			
	☐ No ☐ Yes: Ge	enetic manipulation of the graft:	☐ No ☐ Yes	s:	
	☐ IF AUTOLOGOUS, C	ONTINUE TO "CHRONOLOGICAL NU	JMBER OF HSCT"		

CIC: Hospital UPN:	Patient UIC	HSCT Date: yyyy - mm - dd
ŀ	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, was last HSCT peformed at another institut If >1, please submit an Annual follow up for subsequent transplant as the date of last co (This is so we can capture relapse data and other part of a planned multiple (sequential) graft p	Name of the institution City TM before proceeding, giving the date of the ontact other events between transplants).	
☐ No ☐ Yes		
Pr	reparative Regimen	
Preparative (conditioning) regimen given? No (Usually Paed Inherited Disorders only) Go Yes Prugs No Yes No Yes	☐ Unknown	

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	UNITS				
Ara-C (cytarabine)	2002	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			
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:	Hospit	tal UPN:		Patient UIC	HSCT Da	te: yyyy - mm - dd
Total Body Irradiation (TBI)		No	Yes	: Total prescribed radiation dose as pe		
				umber of fractions ove		
TLI, TNI, TAI lymphoid, nodal, abdominal)		No _				
				Cuminal Status		
		_		Survival Status		
Survival Status on date o		Т				
		stration of t	he pre	parative regimen and date of HSCT		
Main Cause of Dea				nain cause):		
Relapse or Progre	•	-				
☐ HSCT Related Cau						
Unknown						
Other						
Contributor	ry Caus	se of Death	ı (check as many as appropriate):		
☐ GVHD						
Interstiti						
☐ Pulmona ☐ Infection		city				
-	i. terial					
vira						
fun	gal					
	asitic					
	known					
		graft function				
		e Veno occl	usive (disorder (VOD)		
Haemorr						
Cardiac t						
		s system (CN		icity		
		I (GI) toxicity	/			
Skin toxi						
Renal fai		failura				
☐ Multiple						
Other, St	JCUII Y					