

EBMT Centre Identification Code (CIC):	Treatment Type	□ нст	
Hospital Unique Patient Number (UPN):			
Patient Number in EBMT Registry:	Treatment Date _	//	_(YYYY/MM/DD)

ALLOGENEIC HAEMATOPOIETIC CELL TRANSPLANTATION (HCT) Day 0

Date of this HCT://(YYYY/MM/DD) (or planned date of HCT if patient died before treatment))			
Centre where this HCT took place:				
Patient UPN for this treatment:				
Team or unit where treatment took place (select all to	hat apply):			
Adults Pediatrics Haematology 0	Oncology 🔲] Allograft	☐ Autograft	Other; specify:
Unit number: Not applicable				
Indication diagnosis for this HCT: (make sure the indication diagnosis has been registered	first, using the	relevant diagno	sis form)	
Chronological number of this treatment: (Include all types of treatments for this patient, e.g. HCT,	CT, GT, IST)			
Chronological number of this HCT: Chronological number of this allogeneic HCT: (Include all HCTs this patient received in the past)				



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Complete this section only if the <u>chronological number of the treatment is >1</u> for this patient.
<u>If > 1:</u>
Reason for this HCT:
☐ Indication diagnosis
Relapse/progression after previous treatment (HCT/CT/GT/IST)
☐ Complication after previous treatment (HCT/CT/GT/IST)
☐ Primary graft failure
☐ Secondary graft failure
☐ Secondary malignancy
Other; specify:
Date of the last treatment before this one: I I (YYYY/MM/DD)
Type of the last treatment before this one:
☐ Autologous HCT
☐ Allogeneic HCT
Cellular therapy (CT)
☐ Immunosuppressive treatment (IST)
☐ Gene therapy (GT)
Was the last treatment performed at another institution?
□ No
Yes: CIC (if known):
Name of institution:
City:
Submit the relevant follow-up form for the previous HCT/CT/GT/IST using the follow up assessment date before this HCT. It is required to capture relapse data and other events between transplants/cellular therapies.

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Yes:

ЕВМТ	Hospital Unique Patient Number (UPN): Patient Number in EBMT Registry:		Treatment Date _	//(YYYY/MM/DD)	
		DONOR & GRAFT			
Is this HCT p	art of a (planned) multiple (sequent	tial) graft program/prot	ocol?		
☐ No					

If this is the first allogeneic HCT for this patient, complete the patient HLA section in the database.

Chronological number of this HCT as part of multiple (sequential) graft program/protocol for this patient:

Multiple	donors (including multiple CB units):
☐ No	
☐ Yes:	Number of donors:



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DONOR INFORMATION --- Donor ___ (number)---

Copy and fill-in this section as many times as necessary, marking if it refers to Donor 1, 2, etc.

☐ No	(complete only fields	aving their data in the EBMT regis marked with '*' on pages 4-8)	stry?	
☐ Ye	S			
Date o	of birth: / /	(YYYY/MM/DD)		
(year o	of birth is a mandatory	field)	(Skip if the source of ste	om cells is cord blood)
*Age (option	at time of donation: _ nal)	years	*Age in months:	vas younger than 2 years)
*Sex (at birth): lle			
☐ Fe	male			
Dono	r Identification:			
	Donor ID given by the	e treating centre (mandatory):		
	Global registration id	entifier for donors (GRID):		
	ION code of the Done	or Registry or Cord Blood Bank (ma	andatory):	
	EuroCord code for th	e Cord Blood Bank (if applicable): _		
	Name of Donor Regis	stry or Cord Blood Bank:		
	Donor ID given by the	e Donor Registry or Cord Blood Bar	nk:	_
	Patient ID given by the	ne Donor Registry or Cord Blood Ba	ank:	_
*Dono A B AB O	or blood group:	*Donor rhesus factor: Negative Positive	*Donor EBV status: Negative Positive Not evaluated Unknown	*Donor CMV status: Negative Positive Not evaluated Unknown
*Is do No Ye		Sickle cell disease only)		
*Is do)	nked disease? (Inborn Errors only)		
☐ No		ore than one stem cell product:		
☐ Ye	s: *Number of differe	ent stem cell products from this c	donor:	
	(If 2 products e.g. E	BM and PM, complete 'Donor 1 - Pro	oduct Number 1 and 2' on p	page 5)



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Donor(number) continued
Copy and fill-in this section as many times as necessary, marking if it refers to Donor 1, 2, etc.
*Donor(number) - Product Number 1
If more than one stem cell product, this is the <u>first</u> product collected from this donor.
*Source of stem cells: Bone Marrow Peripheral Blood Cord Blood Other; specify:
(select only one)
*Graft manipulation <i>ex-vivo</i> including T-cell depletion:
(other than for RBC removal or volume reduction)
□ No
*Yes: T-cell (CD3+) depletion (Do not use for "Campath in the bag")
☐ T-cell receptor αβ depletion
☐ B-cell depletion (CD19+) by MoAB
☐ NK cell depletion by MoAB
☐ CD34+ enrichment
Genetic manipulation
Other; specify:
U Other, specify.
*Was the graft cryopreserved prior to infusion? \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \



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DONOR INFORMATION --- Donor __(number) continued ---

Copy and fill-in this section as many times as necessary, marking if it refers to Donor 1, 2, etc.

*Donor(number) - Product Number 2	
If more than one stem cell product , this is the <u>first</u> product collected from this c	donor.
(select only one) *Graft manipulation ex-vivo including T-cell depletion:	Other; specify:
(other than for RBC removal or volume reduction)	
*Yes: T-cell (CD3+) depletion (Do not use for "Campath in the bag")	
T-cell receptor αβ depletion	
☐ B-cell depletion (CD19+) by MoAB	
☐ NK cell depletion by MoAB	
☐ CD34+ enrichment ☐ Genetic manipulation	
Other; specify:	
*Was the graft cryopreserved prior to infusion? No Yes; *Date of cryopreservation:// (YYYY/MM/DD) Unknown Unknown	
	— — — — <u>- 2025-</u> 03-25 — -



EBMT Centre Identification Code (CIC): $___$

Patient Number in EBMT Registry: _____

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Co	ppy and fill-in this		DONOR INFORMATION Donor(number) continued as many times as necessary, marking if it refers to Donor 1, 2, etc.			
Relation between	patient and done	or: □ Rel	elated:			
relation between	patient and don	э	elationship to patient: Syngeneic (monozygotic twin)			
			☐ Sibling (may include non-monozygotic twin)			
			☐ Other related: ☐ Parents			
			☐ Child			
			☐ Aunt/Uncle			
			Cousin			
			Grand Parents			
			Other; specify:			
Related donor:		□ on	nrelated (proceed to next page)			
*Both haplotypes ((for both matched a						
*HLA match type:	☐ *Match (botl	n haplotype	pes matched)			
	*Mismatch: *Method used for patient/donor HLA typing: Molecular (select all that apply) Serology					
			ular typing was done:			
		*Locus:				
		A:	0 (match) 1 2 Not evaluated			
		B:	0 (match) 1 2 Not evaluated			
		C:	0 (match) 1 2 Not evaluated			
		DRB1:	0 (match) 1 2 Not evaluated			
		DQB1:	0 (match) 1 2 Not evaluated			
		DPB1:	0 (match) 1 2 Not evaluated			
if carelogical tuning was done.						
	if serological typing was done: *Locus: *Number of mismatches, antigenic:					
		A:	□ 0 (match) □ 1 □ 2 □ Not evaluated			
		B:	□ 0 (match) □ 1 □ 2 □ Not evaluated □ 0 (match) □ 1 □ 2 □ Not evaluated			
		C:	□ 0 (match) □ 1 □ 2 □ Not evaluated □ 0 (match) □ 1 □ 2 □ Not evaluated			
		DRB1:	□ 0 (match) □ 1 □ 2 □ Not evaluated □ 0 (match) □ 1 □ 2 □ Not evaluated			
		DQB1:	□ 0 (match) □ 1 □ 2 □ Not evaluated □ 0 (match) □ 1 □ 2 □ Not evaluated			
			L			
		DPB1:	☐ 0 (match) ☐ 1 ☐ 2 ☐ Not evaluated			

Treatment Type HCT

Treatment Date _ _ _ / _ _ / _ _ (YYYY/MM/DD)

*Please enter the LABORATORY RESULTS WITH HLA TYPING into the database for all the donors



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EBMT	Hospital Unique Patient Number (UPN): Patient Number in EBMT Registry:			Treatment Date / _ / _ (YYYY/MM/DD)
		DONOR IN		
	Сор	 Donor(nur y and fill-in this section as many times as	=	
Unrelated d	onor	:		
*HLA match ty	pe:	*Method used for patient/donor HLA ty (select all that apply)	· • <u> </u>	Molecular Serology
		if molecular typing was done:	*Locus:	*Number of mismatches, allelic:
			A:	0 (match) 1 2 Not evaluated
			B:	☐ 0 (match) ☐ 1 ☐ 2 ☐ Not evaluated
			C:	□ 0 (match) □ 1 □ 2 □ Not evaluated
			DRB1:	0 (match) 1 2 Not evaluated
			DQB1:	0 (match) 1 2 Not evaluated
			DPB1:	☐ 0 (match) ☐ 1 ☐ 2 ☐ Not evaluated
		if serological typing was done:	*Locus:	*Number of mismatches, antigenic:
			A:	0 (match) 1 2 Not evaluated

Treatment Type HCT

0 (match) 1 2 Not evaluated

☐ 0 (match) ☐ 1 ☐ 2 ☐ Not evaluated □ 0 (match) □ 1 □ 2 □ Not evaluated

□ 0 (match) □ 1 □ 2 □ Not evaluated

☐ 0 (match) ☐ 1 ☐ 2 ☐ Not evaluated

*Please enter the LABORATORY RESULTS WITH HLA TYPING into the database for all the donors

B: C:

DRB1:

DQB1: DPB1:

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☐ Myeloablative conditioning (MAC)

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ADDITIONAL ASSESSMENTS (All diagnoses)
Are there Donor-Specific Antibodies (DSA) against HLA?
□ No
Yes: HLA loci the DSA are directed against: A DRB1 B DQB1 C DPB1
Did the patient have desensibilisation therapy? No
(Haemoglobinopathies only)
Are the DSA red cell antibodies? No (Haemoglobinopathies only) Yes: Are they cross-reacting with the red cells of the donor? No
☐ Not evaluated
Unknown
PREPARATIVE REGIMEN (All Diagnoses)
Preparative (conditioning) regimen given?
□ No
☐ Yes
Drugs given? (any active agent, including chemotherapy, monoclonal antibody, polyclonal antibody, serotherapy, etc.) ☐ No ☐ Yes (provide details in the table on pages 10-11)
What type of conditioning regimen was used?
Reduced intensity conditioning (RIC)

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PREPARATIVE REGIMEN continued

Specification and dose of the preparative regimen:

(Report the total prescribed cumulative dose as per protocol. Multiply daily dose by the number of days; e.g. for Busulfan given 4mg/kg daily for 4days, total dose to report is 16mg/kg.

Report dosages and units only for individual drugs.)

Chemotherapy	Dose	Unit
☐ Bendamustine		☐ mg/m² ☐ mg/kg
Bleomycin		mg/m² mg/kg
Busulfan		
Route of administration:		☐ mg/m² ☐ mg/kg
Drug monitoring performed: No Yes; total AUC: mg x hr/L micromol x r mg x min/ml		
Carboplatin		
Drug monitoring performed: No Yes; total AUC: mg x hr/L micromol x m mg x min/mL		☐ mg/m² ☐ mg/kg
☐ Carmustine		☐ mg/m² ☐ mg/kg
☐ Cisplatin		☐ mg/m² ☐ mg/kg
☐ Clofarabine		☐ mg/m² ☐ mg/kg
Corticosteroids: Beclometasone Budesonide Dexamethasone Methylprednisolone Prednisolone		□ mg/m² □ mg/kg □ mg/m² □ mg/kg □ mg/m² □ mg/kg □ mg/m² □ mg/kg □ mg/m² □ mg/kg
Cyclophosphamide		☐ mg/m² ☐ mg/kg
Cytarabine		☐ mg/m² ☐ mg/kg
☐ Daunorubicin		☐ mg/m² ☐ mg/kg
Doxorubicin		☐ mg/m² ☐ mg/kg
☐ Epirubicin		☐ mg/m² ☐ mg/kg
☐ Etoposide		☐ mg/m² ☐ mg/kg
Fludarabine		☐ mg/m² ☐ mg/kg
Gemtuzumab ozogamicin		☐ mg/m² ☐ mg/kg
☐ Ibritumomab tiuxetan		☐ mCi ☐ MBq
☐ Idarubicin		☐ mg/m² ☐ mg/kg



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PREPARATIVE REGIMEN continued

Specification and dose of the preparative regimen:

(Report the total prescribed cumulative dose as per protocol. Multiply daily dose by the number of days; e.g. for Busulfan given 4mg/kg daily for 4days, total dose to report is 16mg/kg.)

4mg/kg daily for 4days, total dose to report is 10mg/kg.)		
Chemotherapy	Dose	Units
☐ Ifosfamide		☐ mg/m² ☐ mg/kg
Imatinib		☐ mg/m² ☐ mg/kg
Lomustine		☐ mg/m² ☐ mg/kg
☐ Melphalan		☐ mg/m² ☐ mg/kg
Mitoxantrone		☐ mg/m² ☐ mg/kg
☐ Paclitaxel		☐ mg/m² ☐ mg/kg
☐ Anti-CD20 antibodies		☐ mg/m² ☐ mg/kg
☐ Teniposide		☐ mg/m² ☐ mg/kg
☐ Thiotepa		☐ mg/m² ☐ mg/kg
☐ Tositumomab		☐ mCi ☐ MBq
☐ Treosulfan		☐ mg/m² ☐ mg/kg
Other; specify*:		☐ mg/m² ☐ mg/kg
		☐ mCi ☐ MBq
Total body irradiation (TBI): No Yes; Total prescribed radiation dose as per protoco Number of fractions: Number of radiation days: Total lymphatic irradiation (TLI): No Yes; Total prescribed radiation dose as per protocol Number of fractions: Number of radiation days:		
Total abdominal irradiation (TAI):		
□ No		
Yes; Total prescribed radiation dose as per protocol	l : Gy	
Number of fractions:		
Number of radiation days:		



Other agent (in vivo); specify*:

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GvHD PREVENTIVE TREATMENT	
GvHD preventive treatment: No Yes: indicate the drugs	
Abatacept	
Alemtuzumab	
Anti-Thymocyte Globulin (ATG) total cumulative dose (mg/kg): H	abbit orse ther; specify:
☐ Basiliximab	
Corticosteroids: Beclometasone Budesonide Dexamethasone Methylprednisolone Prednisolone	
Cyclophosphamide Post Transplant Cyclophosphamide (PTCY) cumulative dose	e (mg/kg): Unknown
Post Transplant Cyclophosphamide (PTCY) timing schedule	 Single dose on day 3 Single dose on day 5 Doses on days 3 and 4 Doses on days 3 and 5 Other, specify:
Cyclosporine	
Etanercept Etanercept	
☐ Everolimus	
☐ Infliximab	
☐ Methotrexate	
Mycophenolate mofetil	
Ruxolitinib	
Sirolimus	
□ Tacrolimus	

*Please consult the LIST OF CHEMOTHERAPY DRUGS/AGENTS AND REGIMENS on the EBMT website for drugs/regimens names

END OF THE ALLO-HCT DAY 0 REPORT
proceed to form DISEASE STATUS AT HCT/CT/GT/IST