

☐ Yes

☐ Unknown

EBMT Centre Identification Code (CIC):	Treatn
Hospital Unique Patient Number (UPN):	
Patient Number in FRMT Registry:	Treatn

Treatment Type	☐ IST	
Treatment Date	1 1	(VVVV/MM/DD)

IMMUNOSUPPRESSIVE TREATMENT (IST) --- Annual/Unscheduled Follow-Up ---

SURVIVAL STATUS				
Date of follow-up://(YYYY/MM/DD) (if patient died: date of death. If patient is lost to follow up: date	last seen)			
Survival status: Alive Dead Lost to follow-up Date of the last IST for this patient://(YYYY//////////////////////////	MM/DD)			
Main cause of death: (check only one main cause)				
☐ Relapse or progression/persistent disease				
☐ Secondary malignancy				
☐ IST-related	Select treatment related cause: (select all that apply) Graft versus Host Disease Non-infectious complication Infectious complication (select all that apply)			
☐ HCT-related	☐ Bacterial infection ☐ Viral infection ☐ Fungal infection ☐ Parasitic infection ☐ Infection with unknown pathogen			
Other; specify:				
Unknown				
Was an autopsy performed? □ No				



 □ Not evaluated ☐ Unknown

	EBMT Centre Identification Code (CIC):	Treatment Type
EBMT	Hospital Unique Patient Number (UPN): Patient Number in EBMT Registry:	Treatment Date // _ (YYYY/MM/DD)
	DECT DECD 110	-
	BEST RESPONS (Complete only for the first an	
Complete Partial ren Haematol Stable dis Relapse / Not evalua Unknown	se after this IST (even if the response got worse again after remission (CR) mission (PR) ogical improvement (HI); NIH partial response ease (no change, no response/loss of response) Progression ated sponse first observed:II (YYYY/MM/DI	
	TRANSFUSION	s
RBC:	☐ Unknown Instrusions given since last follow-up: ☐ No Platelets irradiated: ☐ No ☐ 20 - 50 units ☐ Ye	o es nknown Yes Unknown
	FIRST RELAPSE AFT	ER IST
First relapse ☐ No	s section only fo <u>r the first</u> relapse after this IST. Iprogression of Aplastic Anaemia (detected by any medee of relapse/progression: / / (YYYY/MM/I	
	DISEASE STATUS AT THIS	FOLLOW-UP
☐ Complete ☐ Partial rem ☐ Haematolo ☐ Stable dise	s this follow-up: remission (CR) nission (PR) ogical improvement (HI); NIH Partial Response ease (no change, no response/loss of response) Progression	

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EBMT Centre Identification Code (CIC):	Treatment Type	☐ IST	
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Patient Number in EBMT Registry:	Treatment Date _	//	_(YYYY/MM/DD)

COMPLICATIO	NS SINCE LAST FOLLOW-UP
Adverse events/non-infectious complications grade	3-5 observed (based on CTCAE grades):
☐ No☐ Yes (provide details in the table below)	
Idiopathic pneumonia syndrome	
Complication observed during this follow-up period?	—
	☐ Yes: ☐ Newly developed ☐ Ongoing since previous assessment☐ Unknown
Maximum CTCAE grade observed during this period:	
Onset date (YYYY/MM/DD):/ Ur	known only if newly developed
Resolved: No	
☐ Yes; Stop date (YYYY/MM/DD):	_// Unknown
☐ Unknown	
Veno-occlusive disease (VOD)	
Complication observed during this follow-up period?	—
	☐ Yes: ☐ Newly developed ☐ Ongoing since previous assessment☐ Unknown
Maximum CTCAE grade observed during this period:	☐ Mild ☐ Moderate ☐ Fatal
	Severe Unknown
Onset date (YYYY/MM/DD):/ _ Unk Resolved: ☐ No	nown Only if newly developed
☐ Yes; Stop date (YYYY/MM/DD):	/ / Unknown
Unknown	
Cataract	
Complication observed during this follow-up period?	□ No*
	Yes: Newly developed Ongoing since previous assessment
Maximum CTCAE grade cheened during this paried	Unknown
Maximum CTCAE grade observed during this period:	
Onset date (YYYY/MM/DD):/ Unkir Resolved: No	own Only if newly developed
☐ Yes; Stop date (YYYY/MM/DD):	_// Unknown
Unknown	
Haemorrhagic cystitis, non-infectious	
Complication observed during this follow-up period?	□ No*
	☐ Yes: ☐ Newly developed ☐ Ongoing since previous assessment☐ Unknown
Maximum CTCAE grade observed during this period:	
Onset date (YYYY/MM/DD):/ _ Unkr	nown Only if newly developed
Resolved: No	
☐ Yes; Stop date (YYYY/MM/DD):	_// Unknown
Unknown	

^{*} Grade 0-2



EBMT Centre Identification Code (CIC):	Treatment Type	☐ IST		
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COMPLICATIONS SINCE LAST FOLLOW-UP

ARDS, non-infectious
Complication observed during this follow-up period? No*
☐ Yes: ☐ Newly developed ☐ Ongoing since previous assessment
☐ Unknown Maximum CTCAE grade observed during this period: ☐ 3 ☐ 4 ☐ 5 (fatal) ☐ Unknown
Onset date (YYYY/MM/DD):/ Unknown only if newly developed
Resolved: No
☐ Yes; Stop date (YYYY/MM/DD): / _ ☐ Unknown
Unknown
Multiorgan failure, non-infectious
Complication observed during this follow-up period? No*
☐ Yes: ☐ Newly developed ☐ Ongoing since previous assessment☐ Unknown
Maximum CTCAE grade observed during this period: 3 4 5 (fatal) Unknown
Onset date (YYYY/MM/DD):/ _ Unknown Only if newly developed
Resolved: No
☐ Yes; Stop date (YYYY/MM/DD): / _ ☐ Unknown
☐ Unknown
Renal failure (chronic kidney disease, acute kidney injury)
Complication observed during this follow-up period? No*
☐ Yes: ☐ Newly developed ☐ Ongoing since previous assessment
Unknown CTCAE grade chearyed during this period. 2 7 7 7 7 7 7 7 8 7 8 8 8 8
Maximum CTCAE grade observed during this period: 3 4 5 (fatal) Unknown
Onset date (YYYY/MM/DD):/ _ Unknown Only if newly developed Resolved: No
Yes; Stop date (YYYY/MM/DD):/ _ Unknown
☐ Unknown
Haemolytic anaemia due to blood group
Complication observed during this follow-up period? No*
☐ Yes: ☐ Newly developed ☐ Ongoing since previous assessment
☐ Unknown
Maximum CTCAE grade observed during this period: ☐ 3 ☐ 4 ☐ 5 (fatal) ☐ Unknown
Onset date (YYYY/MM/DD): / _ Unknown Only if newly developed Resolved: No
☐ Yes; Stop date (<i>YYYY/MM/DD</i>): / ☐ Unknown
☐ Unknown

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^{*} Grade 0-2



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COMPLICATIONS SINCE LAST FOLLOW-UP

Aseptic bone necrosis
Complication observed during this follow-up period? No*
☐ Yes: ☐ Newly developed ☐ Ongoing since previous assessment
Unknown
Maximum CTCAE grade observed during this period: ☐ 3 ☐ 4 ☐ 5 (fatal) ☐ Unknown
Onset date (YYYY/MM/DD):/ _ Unknown only if newly developed
Resolved: No
Yes; Stop date (YYYY/MM/DD):/ _ Unknown
☐ Unknown
Liver disorder
Complication observed during this follow-up period? No*
☐ Yes: ☐ Newly developed ☐ Ongoing since previous assessment
☐ Unknown
Maximum CTCAE grade observed during this period: ☐ 3 ☐ 4 ☐ 5 (fatal) ☐ Unknown
Onset date (YYYY/MM/DD):/ _ Unknown Only if newly developed
Resolved: No
☐ Yes; Stop date (YYYY/MM/DD):/ _ ☐ Unknown
☐ Unknown
Cardiovascular event
Complication observed during this follow-up period? No*
☐ Yes: ☐ Newly developed ☐ Ongoing since previous assessment
☐ Unknown
Maximum CTCAE grade observed during this period: ☐ 3 ☐ 4 ☐ 5 (fatal) ☐ Unknown
Onset date (YYYY/MM/DD):/ _ Unknown Only if newly developed
Resolved: No
☐ Yes; Stop date (YYYY/MM/DD): / _ ☐ Unknown
☐ Unknown
Stroke
Complication observed during this follow-up period? No*
☐ Yes: ☐ Newly developed ☐ Ongoing since previous assessment
☐ Unknown
Maximum CTCAE grade observed during this period: ☐ 3 ☐ 4 ☐ 5 (fatal) ☐ Unknown
Onset date (YYYY/MM/DD): / Unknown Only if newly developed
Resolved: No
☐ Yes; Stop date (YYYY/MM/DD):/ ☐ Unknown
☐ Unknown

^{*} Grade 0-2



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Hospital Unique Patient Number (UPN):			
Patient Number in EBMT Registry:	Treatment Date	//	(YYYY/MM/DD)

COMPLICATIONS S	SINCE L	AST	FOLL	OW-UP
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Central nervous system (CNS) toxicity
Complication observed during this follow-up period? No*
☐ Yes: ☐ Newly developed ☐ Ongoing since previous assessment
☐ Unknown
Maximum CTCAE grade observed during this period: ☐ 3 ☐ 4 ☐ 5 (fatal) ☐ Unknown
Onset date (YYYY/MM/DD):/ Unknown only if newly developed
Resolved: No
☐ Yes; Stop date (YYYY/MM/DD): / ☐ Unknown
☐ Unknown
Endocrine event
Complication observed during this follow-up period? No*
☐ Yes: ☐ Newly developed ☐ Ongoing since previous assessment
☐ Unknown
Maximum CTCAE grade observed during this period: ☐ 3 ☐ 4 ☐ 5 (fatal) ☐ Unknown
Onset date (YYYY/MM/DD):/ Unknown Only if newly developed
Resolved: No
☐ Yes; Stop date (YYYY/MM/DD): / _ ☐ Unknown
☐ Unknown
Other complication observed during this follow-up period? No*
☐ Yes: ☐ Newly developed ☐ Ongoing since previous assessment
previous assessment
Specify: Consult appendix 1 for a list of complications that should not be reported
Maximum CTCAE grade observed during this period: 3 4 5 (fatal) Unknown
Onset date (YYYY/MM/DD):/ □ Unknown Only if newly developed
Resolved: No
☐ Yes; Stop date (YYYY/MM/DD):/ ☐ Unknown
☐ Unknown

If more other complications occurred, copy and fill-in this table as many times as necessary.

* Grade 0-2



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Patient Number in EBMT Registry:	Treatment Date _	/	(YYYY/MM/DD)

SECONDARY MALIGNANCIES AND AUTOIMMUNE DISORDERS

Did a secondary malignancy or autoimm ☐ No	ıune disorder occur	?		
Yes: Was this disease an indication fo	or a subsequent HC	:T/CT/IST?		
☐ No (complete the non-indicatio☐ Yes (complete the relevant indi	-	n)		
_	BONE MARROW I	NVESTIGATIO	N	
Bone Marrow Investigation:				
Yes: Date of bone marrow investigation	on: / /	(YYYY/MM/DD)	☐ Unknown	
Type of bone marrow investigati Cytology Histology Both				
Type of dysplasia:				
Erythroid dysplasia Granulocyte dysplasia Megakaryocyte dysplasia	No No No No	Yes	Not evaluated Not evaluated Not evaluated	☐ Unknown ☐ Unknown ☐ Unknown
Bone marrow assessments:				
Cellularity in the bone marrow aspirate	☐ Acellular ☐ Hypocellula ☐ Normocellu ☐ Hypercellu	ular	☐ Focal cellularity ☐ Not evaluated ☐ Unknown	y
Cellularity in the bone marrow trephine	☐ Acellular ☐ Hypocellula ☐ Normocellu ☐ Hypercellu	ular	☐ Focal cellularitg☐ Not evaluated☐ Unknown	у
Fibrosis on bone marrow biopsy	☐ No ☐ Mild ☐ Moderate ☐ Severe		☐ Not evaluable ☐ Not evaluated ☐ Unknown	
CD34+ cell count percentage (%)	c	%	☐ Not evaluated	☐ Unknown
Blast count percentage (%)		% st count is not a	☐ Not evaluated	Unknown
	<u></u> ≤ 5%	<u> </u>	☐ Not evaluated	☐ Unknown



Patient Number in EBMT Registry: CHROMOSOME ANA	Treatment Date / (YYYY/MM/DD) ALYSIS
CHROWOSOWE AWA	11313

Chromosome analysis done at follow-up: (Describe results of the most recent complete analysis)	
☐ No ☐ Yes: Output of analysis: ☐ Separate abnormalities	☐ Full karyotype
☐ Yes: Output of analysis: ☐ Separate abnormalities ☐ Unknown	
If chromosome analysis was done:	
What were the results?	
☐ Normal	
☐ Abnormal: number of abnormalities present:	
Date of chromosome analysis: I I (YYYY/MM/DD)	Unknown
or abnormal results, indicate below whether the abnormalities were abse	ent, present or not evaluated.
abn 3	☐ Absent ☐ Present ☐ Not evaluated
del(13q)	☐ Absent ☐ Present ☐ Not evaluated
Monosomy 7	☐ Absent ☐ Present ☐ Not evaluated
Trisomy 8	☐ Absent ☐ Present ☐ Not evaluated
Other; specify:	☐ Absent ☐ Present
OR Transcribe the complete karyotype:	
rranscribe the complete karyotype.	



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Patient Number in EBMT Registry:	Treatment Date		(YYYY/MM/DD)

MOLECULAR MARKER ANALYSIS

Molecular marker analysis done at f	ollow-up:			
☐ No				
Yes				
Unknown			–	
Date of molecular marker analysis (<i>MM/DD</i>) ☐ Unknown	
Indicate below whether the markers we	•			
ASXL1	Absent	Present	☐ Not evaluated	Unknown
BCOR	Absent	Present	☐ Not evaluated	Unknown
BCORL1	Absent	Present	☐ Not evaluated	Unknown
CBL	Absent	Present	☐ Not evaluated	Unknown
CSMD1	Absent	Present	☐ Not evaluated	Unknown
DNMT3A	☐ Absent	☐ Present	☐ Not evaluated	Unknown
ETV6	Absent	Present	□ Not evaluated	Unknown
EZH2	Absent	☐ Present	☐ Not evaluated	Unknown
FLT3	Absent	Present	☐ Not evaluated	Unknown
GNAS	Absent	Present	☐ Not evaluated	Unknown
IDH1	Absent	Present	☐ Not evaluated	Unknown
IDH2	Absent	Present	☐ Not evaluated	Unknown
JAK2	Absent	Present	☐ Not evaluated	Unknown
KRAS	Absent	Present	☐ Not evaluated	Unknown
MPL	Absent	Present	□ Not evaluated	Unknown
NPM1	Absent	☐ Present	☐ Not evaluated	Unknown
NRAS	Absent	Present	☐ Not evaluated	Unknown
PHF6	Absent	☐ Present	☐ Not evaluated	Unknown
PIGA	Absent	Present	☐ Not evaluated	Unknown
PPM1D	Absent	Present	☐ Not evaluated	Unknown
PTPN11	Absent	Present	☐ Not evaluated	Unknown
RAD21	Absent	☐ Present	☐ Not evaluated	Unknown
RUNX1	Absent	Present	☐ Not evaluated	Unknown
SETBP1	Absent	Present	☐ Not evaluated	Unknown
SF3B1	Absent	Present	☐ Not evaluated	Unknown
SRSF2	Absent	☐ Present	☐ Not evaluated	Unknown
STAG2	Absent	Present	☐ Not evaluated	Unknown
TET2	Absent	Present	☐ Not evaluated	Unknown
	Absent	Present	☐ Not evaluated	Unknown
TP53	TP53 mutation	n type: 🔲 Single h	nit	
		☐ Multi hit		
U2AF1	Absent	Present	Not evaluated	Unknown
ZRSR2	☐ Absent	☐ Present	☐ Not evaluated	☐ Unknown
Other; specify:	☐ Absent	☐ Present	_	
				



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Patient Number in EBMT Registry:	Treatment Date _	//	_ (YYYY/MM/DD)

PNH TESTS SINCE LAST FOLLOW-UP

PNH test done:	
□ No	
Yes: Date of PNH test:/(YYYY/MM/DD)	☐ Unknown
Unknown	
PNH diagnostics by flow cytometry:	
☐ Clone absent	
Clone present: Size of PNH clone in percentage (%):	
Unknown	
Flow cytometry assessment done on:	
☐ Granulocytes	
RBC	
Both	
Other; specify:	



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Patient Number in EBMT Registry:	Treatment Date _	//	(YYYY/MM/DD)

PNH TESTS	SINCE	AST EOU	OW-HD	continued
PIND IESIS	SHACE F	ASI EUL	LUVV-UP	commuea

Clinical manifestation of PNH:					
□ No					
Yes: Date of clinical manifestation: / (YYYY/MM/DD) Unknown					
Anti-complement treatment given?					
□ No					
☐ Yes, complete the table:					
Drug	New or ongoing	Start date (YYYY/MM/DD) (only if new drug administered)	Treatment stopped/date (YYYY/MM/DD)		
☐ Eculizumab	☐ New drug administration ☐ Ongoing since previous assessment	// Unknown	☐ No ☐ Yes: / ☐ Unknown ☐ Unknown		
☐ Ravalizumab	☐ New drug administration☐ Ongoing since previous assessment	// Unknown	☐ No ☐ Yes:/ ☐ Unknown ☐ Unknown		
☐ Pegcetacoplan	☐ New drug administration☐ Ongoing since previous assessment	// Unknown	□ No □ Yes:// □ Unknown □ Unknown		
Other; specify*:	☐ New drug administration ☐ Ongoing since previous assessment	// Unknown	☐ No ☐ Yes:/ ☐ Unknown ☐ Unknown		

If there were more drugs given during one line of treatment add more copies of this page.

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



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Appendix 1

-- Non-infectious Complications CTCAE term --

No Reporting Required

· Allergic reaction

· All laboratory abnormalities

· All types of pain · Alopecia

· Blurred vision

· Diarrhoea (enteropathy)

· Dry mouth · Dyspepsia Dysphagia

 $\cdot \; \text{Edema}$ $\cdot \ \mathsf{Esophageal} \ \mathsf{stenosis}$

Fatigue

· Flashes

· Gastritis

· Hematologic toxicities

· Hematoma Hypertension

· Injection site reaction · Malaise

· Mucositis · Sore throat $\cdot \ \mathsf{Tinnitus}$ · Vertigo · Weight loss