

☐ Yes

☐ Unknown

EBMT Centre Identification Code (CIC):	Treatment [*]
Hospital Unique Patient Number (UPN):	
Patient Number in FRMT Registry	Treatment

Treatment Type	☐ IST		
Treatment Date	1 1	(YYYY/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	

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rauent Number in Edivit Kegistry.	Treatment Date / _ / _ (YYYY/MM/DD)

(Сотр	BEST RES	SPONSE first annual follow-up)	
Best response after this IST (even if the restance of Complete remission (CR) Partial remission (PR) Haematological improvement (HI); NIH partiable disease (no change, no response) Relapse / Progression Not evaluated Unknown Date best response first observed:	artial response loss of response)	· ·	vn	
TRANSFUSIONS				
RBC transfusions given since last follow RBC:	r-up: No RBC irradiated:	☐ Yes ☐ Unk☐ No☐ Yes☐ Unknown	nown	
Platelet transfusions given since last followed	low-up:	No 🗌 Yes	☐ Unknown	
Platelets: < 20 units 20 - 50 units > 50 units Unknown	Platelets irradiated	d: No Yes Unknown		



Patient Number in EBMT Registry: Treatment Date/ _/ (YYYY/MM/DD)
FIRST RELAPSE AFTER IST
Complete this section only for the first relapse after this IST.
First relapse/progression of Aplastic Anaemia (detected by any method): No
Yes: Date of relapse/progression://(YYYY/MM/DD) Unknown
DISEASE STATUS AT THIS FOLLOW-UP
Disease status this follow-up: Complete remission (CR) Partial remission (PR) Haematological improvement (HI); NIH Partial Response Stable disease (no change, no response/loss of response) Relapse / Progression Not evaluated Unknown

Adverse events/non-infectious complications grade 3-5 observed (based on CTCAE grades):

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Yes (provide details in the table on the next page)



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^{*} Grade 0-2



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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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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

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^{*} Grade 0-2



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COMPL	LICATIONS	SINCE	LAST	FOLL	LOW-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
☐ fes. ☐ Newly developed ☐ 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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SECONDARY MALIGNANCIES AND AUTOIMMUNE DISORDERS

Did a secondary malignancy or autoimmune disorder occur? ☐ No						
Yes: Was this disease an indication for a subsequent HCT/CT/IST?						
☐ No (complete the non-indicatio☐ Yes (complete the relevant indi	,	orm)				
Unknown						
	BONE MARROV	V INVESTIC	SATION			
one Marrow Investigation:						
Yes: Date of bone marrow investigation	on: / /	(YYYY/M	<i>M/DD)</i> ☐ Unknown			
Type of bone marrow investigati ☐ Cytology ☐ Histology ☐ Both	on:					
Type of dysplasia:						
Erythroid dysplasia Granulocyte dysplasia Megakaryocyte dysplasia	☐ No ☐ No ☐ No	☐ Yes ☐ Yes ☐ Yes	☐ Not evaluated☐ Not evaluated☐ Not evaluated	☐ Unknown ☐ Unknown ☐ Unknown		
Bone marrow assessments:						
Cellularity in the bone marrow aspirate	☐ Acellula ☐ Hypocel ☐ Normoc ☐ Hyperce	llular ellular	☐ Focal cellularit☐ Not evaluated☐ Unknown	ty		
Cellularity in the bone marrow trephine	☐ Acellula ☐ Hypocel ☐ Normoc ☐ Hyperce	llular ellular	☐ Focal cellularit ☐ Not evaluated ☐ Unknown	ty		
Fibrosis on bone marrow biopsy	☐ No ☐ Mild ☐ Moderat ☐ Severe	ie	☐ Not evaluable ☐ Not evaluated ☐ Unknown			
CD34+ cell count percentage (%)		_ %	☐ Not evaluated	☐ Unknown		
		_ %	☐ Not evaluated	☐ Unknown		
Blast count percentage (%)	If the precise b	olast count i	s not available, please in	dicate whether it is:		
	<u></u> ≤ 5%	☐ > <u>{</u>	5% Not evaluated	Unknown		



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CHROMOSOME ANALYSIS				
come analysis done at follow-up:				

Chromosome analysis done at follow-up: (Describe results of the most recent complete analysis)	
NoYes: Output of analysis: ☐ Separate abnormalities☐ Unknown	☐ Full karyotype
If chromosome analysis was done:	
What were the results?	
☐ Normal☐ Abnormal: number of abnormalities present:☐ Failed	
Date of chromosome analysis:	
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:	



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MOLECULAR MARKER ANALYSIS

Molecular marker analysis done at f	ollow	/-up:						
☐ No ☐ Yes ☐ Unknown								
Date of molecular marker analysis (i	if app	licable):	/ _ ·	_ / (YYYY/M	1M/E	DD) 🗌 Unknown		
Indicate below whether the markers wer	re abs	sent, present or	not	evaluated.				
ASXL1		Absent		Present		Not evaluated		Jnknown
BCOR		Absent		Present		Not evaluated	<u> </u>	Jnknown
BCORL1		Absent		Present		Not evaluated		Jnknown
CBL		Absent		Present		Not evaluated	<u> </u>	Jnknown
CSMD1		Absent		Present		Not evaluated		Jnknown
DNMT3A		Absent		Present		Not evaluated		Jnknown
ETV6		Absent		Present		Not evaluated	<u> П</u>	Jnknown
EZH2		Absent		Present		Not evaluated		Jnknown
FLT3		Absent		Present		Not evaluated		Jnknown
GNAS		Absent		Present		Not evaluated		Jnknown
IDH1		Absent		Present		Not evaluated		Jnknown
IDH2		Absent		Present		Not evaluated		Jnknown
JAK2		Absent		Present		Not evaluated		Jnknown
KRAS		Absent		Present		Not evaluated		Jnknown
MPL		Absent		Present		Not evaluated		Jnknown
NPM1		Absent		Present		Not evaluated		Jnknown
NRAS		Absent		Present		Not evaluated		Jnknown
PHF6		Absent		Present		Not evaluated		Jnknown
PIGA		Absent		Present		Not evaluated		Jnknown
PPM1D		Absent		Present		Not evaluated		Jnknown
PTPN11		Absent		Present		Not evaluated		Jnknown
RAD21		Absent		Present		Not evaluated		Jnknown
RUNX1		Absent		Present		Not evaluated		Jnknown
SETBP1		Absent		Present		Not evaluated		Jnknown
SF3B1		Absent		Present		Not evaluated		Jnknown
SRSF2		Absent		Present		Not evaluated		Jnknown
STAG2		Absent		Present		Not evaluated		Jnknown
TET2		Absent		Present		Not evaluated		Jnknown
		Absent		Present		Not evaluated		Jnknown
TP53	,	TP53 mutation	typ	De: ☐ Single hi ☐ Multi hit ☐ Unknowr				
U2AF1		Absent		Present		Not evaluated		Jnknown
ZRSR2		Absent		Present	<u> </u>	Not evaluated		Jnknown
Other; specify:		Absent	_	Present			Ш`	
	ш '		_					



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PNH TESTS SINCE LAST FOLLOW-UP

PNH test done:	
□ No	
Yes: Date of PNH test:/ (YYYY/MM/DD) Unknown	own
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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PNH TESTS	SINCEL	AST FOLL	OW-HD	continued
PIND IESTS	2010CE	ASI EVLI	_UVV-UP	COMMINGE

Clinical manifestation of	ot PNH:		
□ No			
Yes: Date of clinical r	manifestation: $_{}$ / $_{-}$ / $_{-}$	_(YYYY/MM/DD) 🔲 Unkno	own
Anti-complement t	reatment given?		
☐ No			
Yes, complete the	e 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