

## DISEASE STATUS AT HCT/CT/GT/IST Day 0

**Date of HCT/CT/GT/IST:** \_\_\_\_/ \_ / \_ (YYYY/MM/DD) (or planned date of HCT/CT/GT/IST if patient died before)

### Survival status at HCT/CT/GT/IST:

☐ Alive

Died after conditioning but before HCT/CT/GT/IST

Died after apheresis but before cell infusion

Date of death: \_\_\_\_/ \_\_/ (YYYY/MM/DD)

### Main cause of death:

(check only one main cause)

Relapse or progression/persistent disease	
Secondary malignancy	
CT-related	Select treatment related cause: (select all that apply)
HCT-related	<ul> <li>Infectious complication: (select all that apply)</li> <li>Bacterial infection</li> <li>Viral infection</li> </ul>
GT-related	<ul> <li>Fungal infection</li> <li>Parasitic infection</li> <li>Infection with unknown pathogen</li> </ul>
IST-related (only if IST was a main treatment)	
Unknown	
Other; specify:	

#### Was an autopsy performed?

🗌 No

☐ Yes

Unknown



PATIENT STATUS (All Diagnoses)										
Performance status at initiation of HCT/CT/GT/IST (choose only one):										
Type of scale use	ed:		Score:							
☐ Karnofsky ☐ Lansky	□ 10	20	□ 30	□ 40	50	□ 60	70	80	09 🗌	□ 100
	0		1	2	<u>□</u> 3		4			
Patient weight at initiation of HCT/CT/GT/IST: kg										
Patient height at initiation of HCT/CT/GT/IST: cm										
Patient age at initiation of HCT/CT/GT/IST: years										
Patient EBV status:       Patient CMV status:         Negative       Negative										

Negative
Negative
Positive
Positive
Positive
Not evaluated
Not evaluated
Unknown
Unknown



Definition:

Treatment Type HCT	СТ 🗌	🗌 GT	🗌 IST	Other

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

### **COMORBIDITY INDEX**

Sorror et al., Blood, 2005 Oct 15; 106(8): 2912-2919: http://www.ncbi.nlm.nih.gov/pmc/articles/PMC1895304

# Was there any <u>clinically significant</u> co-existing disease or organ impairment <u>as listed below</u> at time of patient assessment prior to the preparative regimen?

🗌 No

☐ Yes (indicate each comorbidity below)

Unknown

### COMORBIDITY:

Solid tumour, previously present	Treated at any time point in the patient's past history, excluding non-melanoma skin cancer. Indicate type:	🗌 No	🗌 Yes	□ Not evaluated
Inflammatory bowel disease	Crohn's disease or ulcerative colitis	🗌 No	🗌 Yes	☐ Not evaluated
Rheumatologic	SLE, RA, polymyositis, mixed CTD or polymyalgia rheumatica	🗌 No	🗌 Yes	Not evaluated
Infection	Requiring continuation of antimicrobial treatment after day 0	🗌 No	🗌 Yes	□ Not evaluated
Diabetes	Requiring treatment with insulin or oral hypoglycaemics but not diet alone	🗌 No	🗌 Yes	Not evaluated
Renal: moderate/severe	Serum creatinine > 2 mg/dL or >177 $\mu$ mol/L, on dialysis, or prior renal transplantation	🗌 No	☐ Yes	□ Not evaluated
Hepatic	Mild: Chronic hepatitis, bilirubin between Upper Limit Normal (ULN) and 1.5 × ULN, or AST/ALT between ULN and 2.5 × ULN Moderate/severe: Liver cirrhosis, bilirubin greater than 1.5 × ULN, or AST/ALT greater than 2.5 × ULN		erate/severe valuated	
Arrhythmia	Atrial fibrillation or flutter, sick sinus syndrome, or ventricular arrhythmias	🗌 No	🗌 Yes	□ Not evaluated
Cardiac	Coronary artery disease, congestive heart failure, myocardial infarction, $EF \le 50\%$ , or shortening fraction in children (<28%)	🗌 No	🗌 Yes	Not evaluated
Cerebrovascular disease	Transient ischaemic attack or cerebrovascular accident	🗌 No	☐ Yes	□ Not evaluated
Heart valve disease	Except mitral valve prolapse	🗌 No	🗌 Yes	Not evaluated
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	☐ No ☐ Mode ☐ Seve ☐ Not e		
Obesity	Patients with body mass index > 35 kg/m <sup>2</sup>	🗌 No	🗌 Yes	Not evaluated
Peptic ulcer	Requiring treatment	□ No	☐ Yes	□ Not evaluated
Psychiatric disturbance	Depression or anxiety requiring psychiatric consultation or treatment	🗌 No	🗌 Yes	Not evaluated



EBMT Centre Identification Code (CIC):	Treatment Type HCT CT GT IST Oth	er
Hospital Unique Patient Number (UPN):		
Patient Number in EBMT Registry:	Treatment Date / _ / (YYY/MM/DD)	

### **COMORBIDITY INDEX continued**

Sorror et al., Blood, 2005 Oct 15; 106(8): 2912-2919: http://www.ncbi.nlm.nih.gov/pmc/articles/PMC1895304

#### Was there any additional <u>major</u> clinical abnormality not listed above and present prior to the preparative regimen?

□ No

Yes; specify: \_\_\_\_\_

#### Are there any autoimmune diseases?

All autoimmune diseases listed on the autoimmune disease form must be considered. However, note that there may be additional diseases not listed on the form. If these additional indications should be reported, it should be based on the clinical judgement of the investigator at the centre.

🗌 No

Yes; specify:

Date of autoimmune disease diagnosis: \_ \_ \_ / \_ / \_ (YYY/MM/DD) Unknown



### SARS-CoV-2 RELATED QUESTIONS

Did the patient have a <u>symptomatic</u> SARS-CoV-2 infection (positive PCR or antigen test) in the 3 months prior to the day of HCT/CT/GT/IST treatment? Note: do not report here if the infection was asymptomatic.

	No
--	----

□ Yes; Date: \_ \_ / \_ / \_ (YYYY/MM/DD) □ Unknown

Unknown

□ Not evaluated

Did the patient have an ongoing SARS-CoV-2 infection (positive PCR or antigen test) at initiation of HCT/CT/GT/IST (including potential conditioning regimen)?

🗌 No

🗌 Yes

Unknown

☐ Not evaluated

### END OF GENERAL SECTION

TO COMPLETE DISEASE STATUS AT HCT/CT/GT/IST FORM, PLEASE FILL IN THE

DIAGNOSIS-SPECIFIC QUESTIONS IN THE RELEVANT SECTION BELOW.



### Status at HCT/CT/GT/IST treatment

Complete only for one main indication diagnosis for which this HCT/CT/GT/IST is given.

Acute leukaemias	Go to page 7
Chronic leukaemias - Chronic Myeloid Leukaemias (CML)	Go to page 9
Chronic leukaemias - Chronic Lymphocytic Leukaemias (CLL)	Go to page 10
Chronic leukaemias - Prolymphocytic (PLL) and Other Chronic Leukaemias	Go to page 11
Lymphomas	Go to page 12
Myelodysplastic Neoplasms (MDS)	Go to page 14
MDS/MPN Overlap Syndromes	Go to page 15
Myeloproliferative Neoplasms (MPN)	Go to page 16
Plasma Cell Neoplasms (PCN)	Go to page 18
Solid Tumours	Go to page 19
Autoimmune Diseases	Go to page 20
Haemoglobinopathies	Go to page 21
Inborn errors	Go to page 23
Bone Marrow Failure Syndromes (BMF) including Aplastic Anaemia (AA)	Go to page 25

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ACUTE LEUKAEMIAS
Status at HCT/CT/GT/IST treatment

Status:	
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Primary induction failure	
□ 1 <sup>st</sup> complete haematological remission (CR)	
1 <sup>st</sup> relapse	
2 <sup>nd</sup> complete haematological remission (CR)	
2 <sup>nd</sup> relapse	
3 <sup>rd</sup> or higher complete haematological remission (CR)	
☐ 3 <sup>rd</sup> or higher relapse	
🗌 Non blastic pancytopenia	
Not evaluated	
umber of induction courses: Unknown Only for patient in Primary Induction failure or in 1st complete remission)	
one marrow burden (% blasts): % 🔲 Not evaluated 🗍 Unknown	
If the precise blast count is not available, please indicate whether it is:	
$\Box \le 5\%$ $\Box > 5\%$ $\Box$ Not evaluated $\Box$ Unknown	
patient was in complete remission:	
Date of first complete remission:I (YYY/MM/DD) Unknown	
patient was in relapse:	
Date of first relapse: / _ / / (YYY/MM/DD) 🔲 Unknown	
<b>Date of the last relapse before this treatment:</b> <i>I I</i> (YYYY/MM/DD) Unknown (if more than 1 relapse before HCT/CT/GT/IST)	
<b>CD19 expression at the last relapse:</b> Negative Positive Not evaluated (Only for B lymphoblastic leukaemia/lymphoma and Mixed phenotype, if the main treatment is a Cellular Therapy)	



				EMIAS continued T/GT/IST treatment
Involvement at time				
Medullary:	□ No [	🗌 Yes		
Extramedullary:	□ No [	Yes	🔲 Unknown	
Organs involved at ti	me of treatme	ent:		
Skin:	🗌 No	C	Yes	□ Not evaluated
CNS:	🗌 No	-	Yes	□ Not evaluated
Testes/Ovaries:		_	Yes	□ Not evaluated
Other; specify:	No	L	Yes	
Complete this sectiMinimal residual $\Box$ Negative $\Box$ Positive $\Box$ Not evaluatedDate MRD statusSensitivity of MR $\subseteq 10^{-6}$ $\subseteq 10^{-5}$ $\subseteq 10^{-3}$ $\Box$ Other; specify: $\Box$ UnknownMethod used:(select all that app) $\Box$ PCR $\Box$ Flow cytometry	disease (MRI evaluated: _ D assay:	D) at initia	tion of treatme	
<ul> <li>NGS</li> <li>Other; specify:</li> </ul>				
Unknown				

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	<b>Chronic Myeloid</b>	LEUKAEMIAS Leukaemias (CML) T/GT/IST treatment	
Status:			
Chronic phase (CP)			
<u>Number:</u> □ 1 <sup>st</sup>	Haematological remission:	<u>Cytogenetic remission:</u> ☐ No	Molecular remission:
□ 2 <sup>nd</sup>	🗌 Yes	Yes	Yes
☐ 3 <sup>rd</sup> or higher	Not evaluated	Not evaluated	☐ Not evaluated
🗌 Unknown	Unknown	Unknown	Unknown
Accelerated phase			
<u>Number:</u> □ 1 <sup>st</sup>			
$\Box 2^{nd}$			
☐ <sup>–</sup> ☐ 3 <sup>rd</sup> or higher			
☐ Blast crisis			
Number:			
□ 2 <sup>nd</sup>			
☐ 3 <sup>rd</sup> or higher			
🗌 Unknown			
☐ Not evaluated			

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CHRONIC LEUKAEMIAS
Chronic Lymphocytic Leukaemias (CLL)
Status at HCT/CT/GT/IST treatment

#### Status:

Complete remission (CR)	
Partial remission (PR)	
Stable disease (no change,	no response/loss of response)
Relapse (untreated)	
Progressive disease (PD):	Sensitive to last regimen
	Resistant to last regimen
	Unknown
Never treated	
Unknown	
<ul> <li>Negative</li> <li>Positive</li> <li>Not evaluated</li> </ul>	(MRD) at initiation of treatment: ed://(YYYY/MM/DD)



### CHRONIC LEUKAEMIAS Prolymphocytic (PLL) and Other Chronic Leukaemias Status at HCT/CT/GT/IST treatment

### Status:

Complete remission (	(CR)
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- Partial remission (PR)
- Stable disease (no change, no response/loss of response)
- Relapse (untreated)
- Progressive disease (PD): Sensitive to last regimen

Sensitive to last regimen
 Resistant to last regimen

Unknown

- □ Never treated
- Unknown



### LYMPHOMAS Status at HCT/CT/GT/IST treatment

itatus:
Chemorefractory relapse or progression, including primary refractory disease Histopathological verification of relapse:
□ Complete remission (CR): □ Confirmed □ Unconfirmed (CRU*) □ Unknown
Number: (achieved prior to this treatment including this one if applicable)
Partial remission (PR); Number: (achieved prior to this treatment including this one if applicable)
☐ Stable disease (no change, no response/loss of response)
Untreated relapse (from a previous CR) or progression (from a previous PR)
Histopathological verification of relapse: 🗌 No 📄 Yes
] Unknown
Not evaluated

\* CRU: Complete response with persistent scan abnormalities of unknown significance

### Technique used for disease assessment:

- CT scan
- D PET
- 🗍 MRI

Unknown

#### Parameters for international prognostic indices at HCT/CT:

Age at treatment:	years (this i	s calculated	d automatically in the database)
LDH levels elevated: (at the start of preparatory regimen)	🗌 No	☐ Yes	□ Not evaluated
Haemoglobin < 120g/L: (at the start of preparatory regimen)	🗌 No	🗌 Yes	□ Not evaluated
White Blood Cell count: (at the start of preparatory regimen)		x 10 <sup>9</sup> /L	
if patient NOT in complete remiss	sion (CR):		
Ann Arbor staging:			III IV Not evaluated
> 1 extranodal site involved:	🗌 No	Yes	□ Not evaluated
> 4 nodal sites involved:	🗌 No	☐ Yes	□ Not evaluated

#### CNS involvement:

- 🗌 No
- ☐ Yes
- ☐ Not evaluated



### **LYMPHOMAS** Status at HCT/CT/GT/IST treatment continued

### Final score:

(only for patients NOT in Complete Remission with LBCL (except Primary large B-cell lymphoma of immune-privileged sites), Mantle cell lymphoma, Follicular lymphoma, Waldenstrom macroglobulinaemia)

<b>IPI:</b> (for LBCL (except Primary large B-cell lymphoma of immune-privileged sites) and FLBL)	<b>MIPI:</b> (for Mantle cell lymphoma)	<b>FLIPI:</b> (for Follicular lymphoma (except FLBL))	<b>ISSWM:</b> (for Waldenstrom macroglobulinaemia)
<ul> <li>Low risk (0-1 score points)</li> <li>Low-intermediate risk (2 score points)</li> <li>High-intermediate risk (3 score points)</li> <li>High risk (4-5 score points)</li> <li>Not evaluated</li> </ul>	<ul> <li>Low risk</li> <li>Intermediate risk</li> <li>High risk</li> <li>Not evaluated</li> </ul>	<ul> <li>Low risk</li> <li>Intermediate risk</li> <li>High risk</li> <li>Not evaluated</li> </ul>	<ul> <li>Low risk (0-1 score points except age &gt; 65)</li> <li>Intermediate risk (2 score points OR age &gt; 65)</li> <li>High risk (3-5 score points)</li> <li>Not evaluated</li> </ul>

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	PLASTIC NEOP			
Classification at treatment (WHO 2022):				
MDS with defining genetic abnormalities:				
$\Box$ MDS with low blasts and is	olated 5 q deletion (MI	DS-5q)		
$\Box$ MDS with low blasts and S	F3B1 mutation (MDS-S	SF3B1)		
☐ MDS with biallelic TP53 ina	activation (MDS-biTP53	3)		
MDS, morphologically defined:				
☐ MDS with low blasts (MDS	-LB)			
☐ MDS, hypoplastic (MDS-h)				
☐ MDS with increased blasts	(MDS-IB1)			
MDS with increased blasts	(MDS-IB2)			
☐ MDS with fibrosis (MDS-f)				
Childhood myelodysplastic neoplasms (MDS)	):			
Childhood MDS with low bl				
Childhood MDS with increa				
Status:				
Complete remission (CR)	<u>Number:</u> 🔲 1st			
	2nd			
☐ 3rd or higher				
□ Unknown				
Improvement but no CR				
Primary refractory phase (no change)				
□ Relapse	<u>Number:</u> 🔲 1st			
	🗌 2nd			
	🔲 3rd o	r higher		
	🗌 Unkn	own		
Progression/Worsening				
Never treated (supportive care or treatment	without chemotherapy)			
Unknown				
☐ Not evaluated				
IPSS-R: ☐ Very Low (≤1.5)	IPSS-M:	☐ Very Low (≤-1.5)		
Low (>1.5 to 3)		☐ Low (>-1.5 to -0.5)		
☐ Intermediate (>3 to 4.5)		$\square$ Moderate Low (>-0.5 to 0)		
☐ High (>4.5 to 6)		$\square$ Moderate High (>0 to 0.5)		
Very High (>6)		☐ High (>0.5 to 1.5)		
		☐ Very High (>1.5)		
—				
Disease_status_HCT_CT_GT_IST_Day0_v2.0	14 of 25		2024-06-10	



MDS/MPN OVERLAP SYNDROMES
Status at HCT/CT/GT/IST Treatment

### **Classification (WHO 2022):**

Chronic muclemencer tic leukoemic (CMMcL, CMML); CMML authtures C, Mucledysplastic
Chronic myelomonocytic leukaemia (CMMoL, CMML): <b>CMML subtype:</b> Myelodysplastic
Myeloproliferative
CMML subgroup: CMML-1
CMML-2
🗌 Unknown
MDS/MPN with SF3B1 mutation and thrombocytosis
MDS/MPN with neutrophilia (Atypical CML BCR-ABL1 negative)
MDS/MPN with ring sideroblasts and thrombocytosis (MDS/MPN-RS-T)
MDS/MPN not otherwise specified (NOS)

#### Status:

Complete remission (CI	र)	Number:	🗌 1st	
			☐ 2nd	
			☐ 3rd or higher	
			Unknown	
Improvement but no CR				
Primary refractory phase	e (no change)			
🔲 Relapse		<u>Number:</u>	🗌 1st	
			2nd	
			3rd or higher	
			Unknown	
Progression/Worsening				
Never treated (supportiv	e care or treatment wi	thout chem	otherapy)	
Unknown				
□ Not evaluated				
CPSS (for CMML only):	Low		CPSS-Mol (for CMML only):	Low
	 ☐ Intermediate-1			Intermediate-1
	☐ Intermediate-2			□ Intermediate-2
	□ Hiah			🔲 High

🗌 High

Unknown

Unknown



### MYELOPROLIFERATIVE NEOPLASMS (MPN) Status at HCT/CT/GT/IST treatment

### Classification at treatment (WHO 2022):

Primary myelofibrosis (Chronic idiopathic myelofibrosis; fibrosis with myeloid metaplasia)
Secondary myelofibrosis (Transformed to myelofibrosis from PV/ET)
Polycythaemia vera (PV)
Essential or primary thrombocythaemia (ET)
U Juvenile myelomonocytic leukaemia (JCMMoL, JMML, JCML, JCMML)
Hyper eosinophilic syndrome (HES)
Chronic eosinophilic leukaemia (CEL)
Chronic neutrophilic leukaemia
Aggressive systemic mastocytosis
Systemic mastocytosis with an associated haematologic neoplasm (SM-AHD)
Mast cell leukaemia
Mast cell sarcoma
MLN-TK with FGFR1 rearrangement
MLN-TK with PDGFRA rearrangement
MLN-TK with PDGFRB rearrangement
MLN-TK with JAK2 rearragement
MLN-TK with FLT3 rearrangement
MLN-TK with ETV6::ABL1 fusion
Transformed to AML
MPN not otherwise specified (NOS)
Other; specify:

#### Status:

Complete remission (CR)	Number:  1st
	☐ 2nd
	3rd or higher
	Unknown
Improvement but no CR	
Primary refractory phase (no change)	
Relapse	Number: 1st
	☐ 2nd
	3rd or higher
Progression/Worsening	
Never treated (supportive care or treatment wi	thout chemotherapy)
□ Not evaluated	

EBMT Centre Identification Code (CIC):         Hospital Unique Patient Number (UPN):         Patient Number in EBMT Registry:	
	ERATIVE NEOPLASMS (MPN) CT/CT/GT/IST treatment
Blast count (peripheral blood): % 🔲 Not	evaluated 🔲 Unknown
If the patient was not splenectomized:	
(Palpable) Spleen size: cm (below co	ostal margin) 🔲 Not evaluated 🔲 Unknown
Spleen span on ultrasound or CT scan:	cm (maximum diameter) 🔲 Not evaluated 🔲 Unknown
JAK inhibitor exposure between diagnosis and H	CT/CT/GT/IST treatment:
Yes: Was a JAK inhibitor continued during	conditioning?
□ No	
☐ Yes: Dose: mg/day	
Start date: / / /	(YYYY/MM/DD)
End date: / / / (	YYYY/MM/DD)
Response status:	
Spleen response	
Symptoms response	
☐ Stable disease (no change, no resp	onse/loss of response)
Primary resistance	
□ Not evaluated	
Myelofibrosis only:	
DIPSS at HCT/CT/GT/IST treatment:	MIPSS70 at HCT/CT/GT/IST treatment:
Low risk	Low risk
<ul> <li>Intermediate - 1</li> <li>Intermediate - 2</li> </ul>	☐ Intermediate ☐ High risk
☐ Intermediate - 2 ☐ High risk	☐ Not evaluated
☐ Not evaluated	Unknown
Unknown	

### Secondary myelofibrosis only (post-ET MF, post-PV MF): MYSEC-PM at time of secondary MF diagnosis:

	Low risk
	Intermediate - 1
	Intermediate - 2
	High risk
	Not evaluated
П	Unknown

Other



PLASMA CELL NEOPLASMS (P Status at HCT/CT/GT/IST treatm	-
Status:	
Complete remission (CR)	Number: 1st
Very good partial remission (VGPR)	2nd
Partial remission (PR)	☐ 3rd or higher
	Unknown
Progression	
Stable disease (no change, no response/loss of response)	
Never treated (supportive care or treatment without chemotherapy)	
Unknown	
□ Not evaluated	
Complete this section only if the disease status is CR or sCR         Minimal residual disease (MRD) at initiation of treatment:         Negative         Positive         Not evaluated         Date MRD status evaluated:/ (YYYY/MM/DD)         Sensitivity of MRD assay:         \$\le10^6\$         \$\le10^6\$         \$\le10^4\$         \$\le10^4\$         \$\le10^4\$         \$\le10^4\$         \$\le10^4\$         \$\le10^6\$         \$\le10^6\$         \$\le10^6\$         \$\le10^4\$         \$\le10^4\$	
Was the patient on dialysis at any time before HCT/CT?  No Yes; Start date:/ (YYYY/MM/DD) Unknown	
Did dialysis stop?   No	Unknown
Unknown	



### SOLID TUMOURS Status at HCT/CT/GT/IST treatment

#### Status:

Adjuvant	
Complete remission (CR): Confirmed Unconfirmed Unknown	
First Partial remission	
Partial remission (PR)	
Progressive disease	
🗌 Relapse: 🗌 Resistant 🔲 Sensitive 📋 Unknown	
Stable disease (no change, no response/loss of response)	
Never treated (upfront)	
Not evaluated	

Complete this section only if the disease status <u>is not CR</u>	i i
Organ involvement at time of this treatment:	I I
Nodes below diaphragm	1
Nodes above diaphragm	i
	ł
	ł
Bone	÷
	Ì
Soft tissue	I I
Other; specify:	1
	÷.

### Germ cell tumours only:

Risk category at disease recurrence (or platinum refractoriness) following first line chemotherapy:

Note: according to International Prognostic Factors Study Group classification published in 2010.

Very low
Low
Intermediate
High
Very high
Not evaluated



<b>AUTOIMMUNE DISEASES</b>
Status at Mobilisation

Systemic sclerosis only:         SSc subset:         Diffuse cutaneous         Limited cutaneous         Sine scleroderma         Other; specify:         Assessments at time of mobilisation (within 3 months before mobilisation):			
Creatinine Clearance (Cockroft formula): ml/min □ Unknown Proteinuria: g/24hrs □ Unknown Modified Rodnan Skin Score (0-51): □ Unknown DLCO (corrected for Hb):% □ Unknown Mean Pulmonary Arterial Systolic Pressure [PASP] <i>(from right heart catheterisation)</i> : mm Hg GI Involvement: □ No □ Yes □ Not evaluated □ Unknown			
Systemic lupus erythematosus only:         Assessments at time of mobilisation (within 3 months before mobilisation):         SLEDAI-2K Score:          □ Not evaluated         □ Unknown         □         □         □			
Multiple sclerosis only:         Status at time of mobilisation (within 3 months before mobilisation):         Primary progressive         Secondary progressive         Relapsing/remitting         Other; specify:         Assessments at time of mobilisation (within 3 months before mobilisation):         EDSS (1-10):         Number of gadolinium enhancing lesions present on MRI brain scan:			
<u>Crohn's disease only:</u> Assessments at time of mobilisation (within 3 months before mobilisation):			
CDAI (0-700):			

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HAEMOGLOBINOPATHIES Status at HCT/CT/GT/IST treatment			
Ferritin level : ng/mL       Not evaluated       Unknown         Total number of red blood cell units:			
(since the diagnosis or previous HCT/GT) $\square$ 20 to 50 units			
$\Box >50$ units			
Liver study?			
│ □ No │ □ Yes: Liver biopsy performed? □ No			
_	「 F0 (no fibrosis)		
☐ <sup>Yes:</sup> Liver fibrosis (Ishak staging): [	☐ F1 (partial fibrosis)		
	] F2 (general fibrosis)		
[ [ [	] F3 (partial bridging in fibrosis)		
	☐ F4 (general bridging in fibrosis)		
	] F5 (near cirrhosis) ] F6 (cirrhosis)		
	☐ No		
Chronic hepatitis?	_ Yes		
Liver iron concentration assesse	ed? 🗍 No		
	Yes: Iron concentration: mg/g		
	dry weight		
MRI (fibroscan) performed? 🔲 No			
Yes: Liver fibrosis: Absent	☐ Moderate ☐ Severe (bridging cirrhosis)		
Liver iron concentration asse	essed?		
	Yes: Iron concentration: mg/g		
	dry weight		
Was chelation performed regularly?			
No: Estimate the completeness of the chelation therapy administratio	<b>n:</b> %		
Yes: Start date of chelation therapy:// (YYYY/MM/DD)	Unknown		
Cardiac evaluation - cardiac study			
☐ Yes: Cardiac echography ejection fraction: ☐ No ☐ Yes;			
Cardiovascular magnetic resonance (CMR) T2: No Yes;	T2 milliseconds (ms)		



HAEMOGLOBINOPATHIES Status at HCT/CT/GT/IST treatment					
Chronic transfusion program: [	No Did the	e patient receive	hydroxyurea?		
[	Yes				
		Yes: Please spe	ecify the duration of hydroxyurea therapy:	months	
Endocrinopathies pre-existing	to HCT/CT/GT:				
Hypothyroidism	□ No	🗌 Yes	Not evaluated		
Hypoparathyroidism		☐ Yes	Not evaluated		
Diabetes mellitus	□ No	🗌 Yes	Not evaluated		
Osteoporosis	No	☐ Yes	Not evaluated		
Gonadal dysfunction	🔲 No	🗌 Yes	Not evaluated		
Growth impairment	□ No	🗌 Yes	Not evaluated		
Pre-treatment complications	(check all that app	ly)			
-	,,,	<i>,</i> ,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,			
Cerebrovascular disease Abnormal Doppler			☐ Not evaluated		
Stroke		Yes Vos			
Haemorrhage		☐ Yes	Not evaluated		
Arteriopathy		☐ Yes	Not evaluated           Not evaluated		
Moyamoya disease	□ No □ No	☐ Yes □ Yes			
Silent infarcts		☐ Tes	Not evaluated		
	L NO				
Renal involvement					
Microalbumin level	Microalbumin level mg/g 🔲 Not evaluated				
Glomerular filtration rate mL/min/1.73m <sup>2</sup> D Not evaluated					
Avascular necrosis	🗌 No	Yes 🗌 No	ot evaluated		
Hyperhaemolysis or autoimmune No haemolytic anaemia: Yes: Hyperhaemolysis Autoimmune haemolytic anaemia Not evaluated					
Other SCD related complications					
Acute chest syndrome	🗌 No	🗌 Yes	Not evaluated		
Vaso-occlusive crisis	🗌 No	🗌 Yes	Not evaluated		
Priapism	🗌 No	🗌 Yes	Not evaluated		
Pulmonary hypertension	🗌 No	🗌 Yes	Not evaluated		
Chronic lung disease	🗌 No	☐ Yes	☐ Not evaluated		



COMORBIDITY:

Treatment Date \_ \_ \_ / \_ / \_ (YYY/MM/DD)

### Inborn Errors Status at HCT/CT/GT/IST treatment

### Immune profiling

<b>Test date</b> (within 3 months prior to HCT/CT/GT):	1	1	(YYYY/MM/DD)	Г	1 Unknown
	- '	'	(11111111111000)		

Cell type and test results		Units (for CD4 and CD8, select unit)
T-cells (CD3):	☐ Not evaluated	1/µL
CD4 T-cells (CD4):	☐ Not evaluated	1/µL
CD8 T-cells (CD8):	☐ Not evaluated	1/µL
B-cells (CD19):	☐ Not evaluated	1/µL
NK-cells (CD16/CD56):	☐ Not evaluated	1/µL
Naive CD4 T-cells (CD4/CD45RA):	☐ Not evaluated	🗌 % of CD4 📋 1/μL
Naive CD8 T-cells (CD8/CD45RA):	Not evaluated	🔲 % of CD8 📋 1/μL
lgG:	☐ Not evaluated	Gram/L
IgA:	☐ Not evaluated	Gram/L
IgM:	☐ Not evaluated	Gram/L

### COMORBIDITY INDEX

Inborn Errors of Immunity only

Chronic lung disease	Bronchiectasis, interstitial pneumonitis, GLILD, oxygen dependency, structural lung disease (e.g. pneumatoceles)	🗌 No	Yes	Not evaluated
Previous haematological malignancy	Leukaemia, lymphoma, myelodysplastic syndrome (MDS)	🗌 No	☐ Yes	☐ Not evaluated
Failure to thrive	Weight <3 <sup>rd</sup> percentile or requirement for (par)enteral feeding	🗌 No	🗌 Yes	Not evaluated
Active infection at HCT	Any infection requiring therapy in the immediate pre HCT period	🗌 No	🗌 Yes	☐ Not evaluated
Lymphoproliferation	I.e. splenomegaly, organ specific lymphoproliferation	🗌 No	🗌 Yes	Not evaluated
Pre-HCT organ impairment	Infectious or non-infectious (including neurologic)	🗌 No	🗌 Yes	☐ Not evaluated
Autoimmunity/autoinflammation	Active at HCT (includes patients in remission but on immunomodulatory treatment within 3 months before HCT)	🗌 No	Yes	Not evaluated

Patient admitted in ICU: No Yes Unknown (Patient admitted in ICU in the 3 months before HCT/CT/GT)

Definition:



### Inborn Errors

Status at HCT/CT/GT/IST treatment

### Immunomodulatory treatments

Only report treatments administered in the 3 months before this HCT/CT/GT: (select all that apply)

- IVIG
  SCIG
  Steroids (>0.5 mg/kg/day prednison equivalent)
  Cyclosporine A
  Tacrolimus
  Sirolimus
  Ruxolitinib
  Baricitinib
  Other JAK-inhibitor, specify: \_\_\_\_\_\_
  Leniolisib
  Abatacept
  Anakinra
  Canakinumab
  Etoposide
- 🔲 Interferon gamma
- Etanercept
- □ Infliximab
- U Vedolizumab
- Dupilumab
- Emapalumab
- □ PEG-ADA
- Other drug; specify: \_\_\_\_\_



# Bone marrow failure syndromes (BMF) including Aplastic Anaemia (AA) Status at HCT/CT/GT/IST treatment

Serology			
Ferritin level: ng/mL	☐ Not evaluated	Unknown	