

**TREATMENT
NON – HCT/CT/GT/IST****(Malignant disorders and Immunoglobulin-related amyloidosis(AL))**

Mobilisation, the preparative regimen (conditioning/lymphodepletion) and GvHD preventive treatment should be reported in the HCT/CT/GT-related forms. Treatment for GvHD and complications should be reported in the designated form, part of the extended dataset.

Do not use this form to report immunosuppressive treatments for bone marrow failures. Use the IST day 0 treatment form

Date treatment started: ____/____/____ (YYYY/MM/DD)**Diagnosis for which this treatment was given:** _____**Reason for this treatment:**

For Acute Leukemia please select Relapse/Progression for haematological relapse /progression (Blast in BM >5% and/or blast in PB >0% and/or extramedullary disease)

☐ Induction☐ Bridging☐ Relapse / Progression☐ Maintenance / preventive treatment:

MRD status at the start of treatment:

☐ CR MRD negative☐ CR MRD positive☐ CR MRD unknown☐ Not applicable - Not in CR☐ Consolidation☐ Other reason this treatment was given; specify: _____**CHEMOTHERAPY / DRUG REGIMEN****Chemotherapy/Drugs:** ☐ No ☐ Yes ☐ Unknown**If patient received chemotherapy/drugs:**

(Do not report each drug start/end date separately)

Start date: ____/____/____ (YYYY/MM/DD) ☐ Unknown

(report earliest start date of chemo/drugs for this treatment)

Treatment stopped: ☐ No☐ Yes; **End date:** ____/____/____ (YYYY/MM/DD) ☐ Unknown

(report latest end date of chemo/drugs for this treatment)

Reason for treatment withdrawal:

(for Chronic Lymphocytic Leukaemia only)

☐ Planned withdrawal☐ Toxicity☐ Progression or insufficient response☐ Other reason; specify _____☐ Unknown☐ Unknown

CHEMOTHERAPY / DRUG REGIMEN

Chemo/Drug regimen*: _____

Extended dataset

(Acute Leukaemia and Lymphoma only)

Number of cycles: ____ ☐ Unknown

(Acute Leukaemia only)

Number of days per cycle: ____ ☐ Unknown

Daily dose: ____ ☐ Unknown

Units: ☐ mg/m² ☐ mg/kg ☐ mg ☐ mg/mL ☐ UI/m²

Chemo/Drug regimen*: _____

Extended dataset

(Acute Leukaemia and Lymphoma only)

Number of cycles: ____ ☐ Unknown

(Acute Leukaemia only)

Number of days per cycle: ____ ☐ Unknown

Daily dose: ____ ☐ Unknown

Units: ☐ mg/m² ☐ mg/kg ☐ mg ☐ mg/mL ☐ UI/m²

Chemo/Drug regimen*: _____

Extended dataset

(Acute Leukaemia and Lymphoma only)

Number of cycles: ____ ☐ Unknown

(Acute Leukaemia only)

Number of days per cycle: ____ ☐ Unknown

Daily dose: ____ ☐ Unknown

Units: ☐ mg/m² ☐ mg/kg ☐ mg ☐ mg/mL ☐ UI/m²

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

Copy and fill-in this page (chemotherapy / drug regimen) as often as necessary within the same line of treatment.

Extended dataset

Intrathecal therapy: ☐ No ☐ Yes ☐ Unknown

(Acute Leukaemia only)

INTERVENTIONS

Radiotherapy: ☐ No ☐ Yes ☐ Unknown

If patient received radiotherapy incl. irradiation:

Start date: ____/____/____ (YYYY/MM/DD) ☐ Unknown

Treatment stopped: ☐ No

☐ Yes; **End date:** ____/____/____ (YYYY/MM/DD) ☐ Unknown

☐ Unknown

Splenic irradiation: ☐ No ☐ Yes ☐ Unknown

(for Myeloproliferative neoplasms only)

If patient received splenic irradiation:

Total prescribed radiation dose as per protocol (Gy): _____ ☐ Unknown

Number of fractions: _____ ☐ Unknown

Number of radiation days: _____ ☐ Unknown

Surgery: ☐ No ☐ Yes ☐ Unknown

If patient underwent surgery:

Date: ____/____/____ (YYYY/MM/DD) ☐ Unknown

Specify the surgery type: _____ ☐ Unknown

Copy and fill-in this section as often as necessary within the same line of treatment.

RESPONSE TO THIS LINE OF TREATMENT

(Disease Specific)

Complete only one section with the main indication diagnosis for which treatment was given.

Response assessment date: ____/____/____ (YYYY/MM/DD) ☐ Unknown

ACUTE LEUKAEMIAS	<i>Go to page 4</i>
CHRONIC LEUKAEMIAS	<i>Go to page 4</i>
PLASMA CELL NEOPLASMS (PCN)	<i>Go to page 4</i>
MYELOPROLIFERATIVE NEOPLASMS (MPN)	<i>Go to page 5</i>
MYELOYDYSPLASTIC NEOPLASMS (MDS)	<i>Go to page 5</i>
MDS / MPN OVERLAP SYNDROMES	<i>Go to page 5</i>
LYMPHOMAS	<i>Go to page 6</i>
SOLID TUMOURS	<i>Go to page 6</i>
OTHER DIAGNOSIS	<i>Go to page 6</i>

RESPONSE TO THIS LINE OF TREATMENT

Complete only one section with the main indication diagnosis for which treatment was given.

Acute leukaemias (AML, PLN, Other)

☐ Complete remission (CR)
☐ Not in complete remission
☐ Not evaluated
☐ Unknown

Chronic leukaemias (CML, CLL, PLL, Other)

Chronic Myeloid Leukaemia (CML):

☐ Chronic phase (CP); Number: ☐ 1 st ☐ 2 nd ☐ 3 rd or higher ☐ Unknown Haematological remission: ☐ No ☐ Yes ☐ Not evaluated ☐ Unknown Cytogenetic remission: ☐ No ☐ Yes ☐ Not evaluated ☐ Unknown Molecular remission: ☐ No ☐ Yes ☐ Not evaluated ☐ Unknown
☐ Accelerated phase; Number: ☐ 1 st ☐ 2 nd ☐ 3 rd or higher ☐ Unknown
☐ Blast crisis; Number: ☐ 1 st ☐ 2 nd ☐ 3 rd or higher ☐ Unknown
☐ Not evaluated
☐ Unknown

Chronic Lymphocytic Leukaemia (CLL), Prolymphocytic Leukaemia (PLL) and other chronic leukaemias:

☐ Complete remission (CR)
☐ Partial remission (PR)
☐ Progression: ☐ Resistant to last regimen ☐ Sensitive to last regimen ☐ Unknown
☐ Stable disease (no change, no response/loss of response)
☐ Relapse
☐ Not evaluated
☐ Unknown

Plasma cell neoplasms (PCN)

☐ Complete remission (CR)	Number: ☐ 1 st ☐ 2 nd ☐ 3 rd or higher ☐ Unknown
☐ Stringent complete remission (sCR)	
☐ Very good partial remission (VGPR)	
☐ Partial remission (PR)	
☐ Relapse	
☐ Progression	
☐ Stable disease (no change, no response/loss of response)	
☐ Not evaluated	
☐ Unknown	

For AL, CLL and PCN proceed to next page

RESPONSE TO THIS LINE OF TREATMENT**Complete only for AL, CLL and PCN****Leukaemias (AL, CLL) and PCN** (complete only for patient in CR or sCR)**Minimal residual disease (MRD):**

- ☐ Negative
☐ Positive
☐ Not evaluated
☐ Unknown

*Extended dataset***Date MRD status evaluated:** _ _ _ _ / _ _ / _ _ (YYYY/MM/DD) ☐ Unknown**Sensitivity of MRD assay:**

- ☐ $\leq 10^{-6}$
☐ $\leq 10^{-5}$
☐ $\leq 10^{-4}$
☐ $\leq 10^{-3}$
☐ Other; specify: _____
☐ Unknown

Method used:

(select all that apply)

- ☐ PCR
☐ Flow cytometry
☐ NGS
☐ Other; specify: _____
☐ Unknown

Complete only one section with the main indication diagnosis for which treatment was given.**Myeloproliferative neoplasms (MPN), Myelodysplastic neoplasms (MDS), MDS/MPN overlap syndromes**

- | | |
|--|---|
| ☐ 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 or higher
☐ Unknown |
|----------------------------------|---|

☐ Progression/Worsening☐ Not evaluated☐ Unknown

RESPONSE TO THIS LINE OF TREATMENT

continued

Complete only one section with the main indication diagnosis for which treatment was given.

Lymphomas

☐ Chemorefractory/ radiorefractory relapse or progression, including primary refractory disease
☐ Complete remission (CR)
☐ Partial remission (PR)
☐ Stable disease (no change, no response/loss of response)
☐ Untreated relapse (from a previous CR) or progression (from a previous PR)
☐ Not evaluated
☐ Unknown

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

Technique used for disease assessment: ☐ CT scan

☐ MRI

☐ PET-CT

☐ Other; specify: _____

☐ Unknown

Solid tumours

☐ 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)
☐ Not evaluated
☐ Unknown

Other diagnosis

☐ No evidence of disease
☐ Improved
☐ No response
☐ Worse
☐ Not evaluated
☐ Unknown