

# ERROR CORRECTION FORM

Sequence Number:

CIBMTR Recipient ID:

Initials:

Today's Date:

Infusion Date:

CIBMTR Center Number:

## Form 2400 R7.0: Pre-Transplant Essential Data

Center: \_\_\_\_\_ CRID: \_\_\_\_\_

### Key Fields

OMB No: 0915-0310

Expiration Date: 10/31/2022

**Public Burden Statement:** The purpose of the data collection is to fulfill the legislative mandate to establish and maintain a standardized database of allogeneic marrow and cord blood transplants performed in the United States or using a donor from the United States. The data collected also meets the C.W. Bill Young Cell Transplantation Program requirements to provide relevant scientific information not containing individually identifiable information available to the public in the form of summaries and data sets. An agency may not conduct or sponsor, and a person is not required to respond to, a collection of information unless it displays a currently valid OMB control number. The OMB control number for this information collection is 0915-0310 and it is valid until 10/31/2022. This information collection is voluntary under The Stem Cell therapeutic and Research Act of 2005, Public Law (Pub. L.) 109-129, as amended by the Stem Cell Therapeutic and Research Reauthorization Act of 2010, Public Law 111-264 (the Act) and the Stem Cell Therapeutic and Research Reauthorization Act of 2015, Public Law 114-104. Public reporting burden for this collection of information is estimated to average 0.68 hours per response, including the time for reviewing instructions, searching existing data sources, and completing and reviewing the collection of information. Send comments regarding this burden estimate or any other aspect of this collection of information, including suggestions for reducing this burden, to HRSA Reports Clearance Officer, 5600 Fishers Lane, Room 14N136B, Rockville, Maryland, 20857 or paperwork@hrsa.gov.

Sequence Number: \_\_\_\_\_

Date Received: \_\_\_\_ - \_\_\_\_ - \_\_\_\_

#### Center Identification

CIBMTR Center Number: \_\_\_\_\_

EBMT Code (CIC): \_\_\_\_\_

#### Recipient Identification

CIBMTR Research ID: (CRID) \_\_\_\_\_

Event date: \_\_\_\_ - \_\_\_\_ - \_\_\_\_

### Recipient Information

Questions: 1 - 25

1 Date of birth: \_\_\_\_ - \_\_\_\_ - \_\_\_\_

2 Sex  
 male  female

3 Ethnicity  
 Hispanic or Latino  
 Not Hispanic or Latino  
 Not applicable (not a resident of the USA)  
 Unknown

4 Race (check all that apply)  
 White  
 Black or African American  
 Asian  
 American Indian or Alaska Native  
 Native Hawaiian or Other Pacific Islander  
 Not reported  
 Unknown

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**5 Race detail (check all that apply)**

- Eastern European
- Mediterranean
- Middle Eastern
- North Coast of Africa
- North American
- Northern European
- Western European
- White Caribbean
- White South or Central American
- Other White
- African
- African American
- Black Caribbean
- Black South or Central American
- Other Black
- Alaskan Native or Aleut
- North American Indian
- American Indian, South or Central America
- Caribbean Indian
- South Asian
- Filipino (Pilipino)
- Japanese
- Korean
- Chinese
- Vietnamese
- Other Southeast Asian
- Guamanian
- Hawaiian
- Samoan
- Other Pacific Islander
- Unknown

**6 Country of primary residence** \_\_\_\_\_

**7 State of residence of recipient (for residents of Brazil)** \_\_\_\_\_

**8 Province or territory of residence of recipient (for residents of Canada)** \_\_\_\_\_

**9 State of residence of recipient (for residents of USA)** \_\_\_\_\_

**10 NMDP Recipient ID (RID):** \_\_\_\_\_

**11 Zip or postal code for place of recipient's residence: (USA recipients only)** \_\_\_\_\_ (last 4 digits optional)

**12 Specify blood type (of recipient) (For allogeneic HCTs only)**

- A  B  AB  O

**13 Specify Rh factor (of recipient) (For allogeneic HCTs only)**

- Positive  Negative

**14 Has the recipient signed an IRB / ethics committee (or similar body) approved consent form for submitting research data to the NMDP / CIBMTR?**

- Yes (recipient consented)  
 No (recipient declined)  
 Not approached

**15 Did the recipient give permission to be directly contacted by CIBMTR for future research?**

- Yes (recipient provided permission)  
 No (recipient declined)

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- 16 Date form was signed: \_\_\_\_-\_\_\_\_-\_\_\_\_
- 17 Has the recipient signed an IRB / ethics committee (or similar body) approved consent form to donate research blood samples to the NMDP / CIBMTR? **(For allogeneic HCTs only)**
- Yes (recipient consented)
- No (recipient declined)
- Not approached
- Not applicable (center not participating)
- 18 Date form was signed: \_\_\_\_-\_\_\_\_-\_\_\_\_
- 19 Did the recipient submit a research sample to the NMDP/CIBMTR repository? **(Related donors only)**
- yes  no
- 20 Research sample recipient ID: \_\_\_\_\_
- 21 Is the recipient participating in a clinical trial? (clinical trial sponsors that use CIBMTR forms to capture outcomes data)
- yes  no

### Clinical Trials (1)

Questions: 22 - 25

- 22 Study Sponsor: \_\_\_\_\_
- 23 Specify other sponsor: \_\_\_\_\_
- 24 Study ID Number: \_\_\_\_\_
- 25 Subject ID: \_\_\_\_\_

### Hematopoietic Cellular Transplant (HCT) and Cellular Therapy

Questions: 26 - 45

- 26 Is a subsequent HCT planned as part of the overall treatment protocol (not as a reaction to post-HCT disease assessment)? **(For autologous HCTs only)**
- yes  no
- 27 Specify subsequent HCT planned
- Autologous  Allogeneic
- 28 Has the recipient ever had a prior HCT?
- Yes  No
- 29 Specify the number of prior HCTs: \_\_\_\_\_
- 30 Were all prior HCTs reported to the CIBMTR?
- Yes  No  Unknown

### Prior HCTs (1)

Questions: 31 - 34

- 31 Date of the prior HCT: \_\_\_\_-\_\_\_\_-\_\_\_\_  Date estimated
- 32 Was the prior HCT performed at a different institution?
- Yes  No
- Specify the institution that performed the last HCT:**
- 33 Name: \_\_\_\_\_
- City: \_\_\_\_\_
- State: \_\_\_\_\_
- Country: \_\_\_\_\_
- 34 What was the HPC source for the prior HCT? (check all that apply)
- Autologous
- Allogeneic, unrelated
- Allogeneic, related

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### 35 Reason for current HCT

- Graft failure / insufficient hematopoietic recovery
- Persistent primary disease
- Recurrent primary disease
- Planned subsequent HCT, per protocol
- New malignancy (including PTLD and EBV lymphoma)
- Insufficient chimerism
- Other

36 Date of graft failure / rejection: \_\_\_\_ - \_\_\_\_ - \_\_\_\_

37 Date of relapse: \_\_\_\_ - \_\_\_\_ - \_\_\_\_

38 Date of secondary malignancy: \_\_\_\_ - \_\_\_\_ - \_\_\_\_

39 Specify other reason: \_\_\_\_\_

### 40 Has the recipient ever had a prior cellular therapy? (do not include DLIs)

- Yes  No  Unknown

### 41 Were all prior cellular therapies reported to the CIBMTR?

- Yes  No  Unknown

### Prior Cellular Therapies (1)

Questions: 42 - 45

42 Date of the prior cellular therapy: \_\_\_\_ - \_\_\_\_ - \_\_\_\_

43 Was the cellular therapy performed at a different institution?

- Yes  No

44 Name: \_\_\_\_\_

City: \_\_\_\_\_

State: \_\_\_\_\_

Country: \_\_\_\_\_

45 Specify the source(s) for the prior cellular therapy (check all that apply)

- Autologous
- Allogeneic, unrelated
- Allogeneic, related

### Donor Information

Questions: 46 - 83

### 46 Multiple donors?

- yes  no

47 Specify number of donors: \_\_\_\_\_

### Donor Information for this HCT (1)

Questions: 48 - 83

### 48 Specify donor

- Autologous  Allogeneic, related  Allogeneic, unrelated

### 49 Specify product type (check all that apply)

- Bone marrow
- PBSC
- Single cord blood unit
- Other product

50 Specify other product: \_\_\_\_\_

### 51 Is the product genetically modified? If autologous, go to question 80. If allogeneic related, go to question 52. If allogeneic unrelated, go to question 56.

- Yes  No

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**52 Specify the related donor type**

- Syngeneic (monozygotic twin)
- HLA-identical sibling (may include non-monozygotic twin)
- HLA-matched other relative (does NOT include a haplo-identical donor)
- HLA-mismatched relative

**53 Specify the biological relationship of the donor to the recipient**

- Mother
- Father
- Child
- Sibling
- Fraternal twin
- Maternal aunt
- Maternal uncle
- Maternal cousin
- Paternal aunt
- Paternal uncle
- Paternal cousin
- Grandparent
- Grandchild
- Other biological relative

**54 Specify other biological relative:** \_\_\_\_\_

**55 Degree of mismatch (related donors only)**

- HLA-mismatched 1 allele
- HLA-mismatched ≥ 2 alleles (does include haplo-identical donor)

**56 Specify unrelated donor type**

- HLA matched unrelated
- HLA mismatched unrelated

**57 Did NMDP / Be the Match facilitate the procurement, collection, or transportation of the product?**

- Yes  No

**58 Was this donor used for any prior HCTs? (for this recipient)**

- yes  no

**59 NMDP cord blood unit ID:** \_\_\_\_\_

**61 Non-NMDP unrelated donor ID: (not applicable for related donors)** \_\_\_\_\_

**62 Non-NMDP cord blood unit ID: (include related and autologous CBUs)** \_\_\_\_\_

**63 Global Registration Identifier for Donors (GRID)** \_\_\_\_\_

**64 Is the CBU ID also the ISBT DIN number?**

- Yes  No  Unknown

**65 Specify the ISBT DIN number:** \_\_\_\_\_

**66 Registry or UCB Bank ID** \_\_\_\_\_

**67 Specify other Registry or UCB Bank:** \_\_\_\_\_

**68 Date of birth (donor / infant)**

- Known  Unknown

**69 Date of birth: (donor / infant)** \_\_\_\_ - \_\_\_\_ - \_\_\_\_

**70 Age (donor / infant)**

- Known  Unknown

**71 Age: (donor / infant)** \_\_\_\_\_  Months (use only if less than 1 year old)

- years

**72 Sex (donor / infant)**

- male  female

**73 Specify blood type (donor) (non-NMDP allogeneic donors only)**

- A  B  AB  O

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**74 Specify Rh factor (donor) (non-NMDP allogeneic donors only)**

- Positive  Negative

**75 Donor CMV-antibodies (IgG or Total) (Allogeneic HCTs only)**

- Reactive  
 Non-reactive  
 Indeterminate  
 Not done  
 Not applicable (cord blood unit)

**76 Has the donor signed an IRB / ethics committee (or similar body) approved consent form to donate research blood samples to the NMDP / CIBMTR? (Related donors only)**

- Yes (donor consented)  
 No (donor declined)  
 Not approached  
 Not applicable (center not participating)

77 Date form was signed: \_\_\_\_-\_\_\_\_-\_\_\_\_

**78 Did the donor submit a research sample to the NMDP/CIBMTR repository? (Related donors only)**

- yes  no

79 Research sample donor ID: \_\_\_\_\_

80 Specify number of products infused from this donor: \_\_\_\_\_

81 Specify the number of these products intended to achieve hematopoietic engraftment: \_\_\_\_\_

**Questions 82 - 83 are for autologous HCT recipients only. If other than autologous skip to question 84.**

**82 What agents were used to mobilize the autologous recipient for this HCT? (check all that apply)**

- G-CSF (filgrastim, Neupogen)  
 Pegylated G-CSF (pegfilgrastim, Neulasta)  
 Plerixafor (Mozobil)  
 Combined with chemotherapy  
 Anti-CD20 (rituximab, Rituxan)  
 Other agent

83 Specify other agent: \_\_\_\_\_

### Clinical Status of Recipient Prior to the Preparative Regimen (Conditioning)

Questions: 84 - 87

**84 What scale was used to determine the recipient's functional status?**

- Karnofsky (recipient age ≥ 16 years)  
 Lansky (recipient age ≥ 1 year and < 16 years)

**Performance score prior to the preparative regimen:**

85 Karnofsky Scale (recipient age ≥ 16 years) \_\_\_\_\_

86 Lansky Scale (recipient age ≥ 1 year and < 16 years) \_\_\_\_\_

**87 Recipient CMV-antibodies (IgG or Total)**

- Reactive  Non-reactive  Indeterminate  Not done

### Comorbid Conditions

Questions: 88 - 116

**88 Has the patient been infected with COVID-19 (SARS-CoV-2) based on a positive test result at any time prior to the start of the preparative regimen / infusion?**

- Yes  No

**89 Did the patient require hospitalization for management of COVID-19 (SARS-CoV-2) infection?**

- Yes  No

**90 Was mechanical ventilation given for COVID-19 (SARS-CoV-2) infection?**

- Yes  No

**91 Is there a history of mechanical ventilation? (excluding COVID-19 (SARS-CoV-2))**

- yes  no

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CRID: \_\_\_\_\_

92 Is there a history of invasive fungal infection?

- Yes  No

93 Glomerular filtration rate (GFR) before start of preparative regimen (**pediatric only**)

- Known  Unknown

94 Glomerular filtration rate (GFR): \_\_\_\_\_ mL/min/1.73<sup>2</sup>

95 Does the recipient have known complex congenital heart disease? (corrected or uncorrected) (excluding simple ASD, VSD, or PDA repair) (**pediatric only**)

- Yes  No

96 Were there any co-existing diseases or organ impairment present according to the HCT comorbidity index (HCT-CI)? *Source: Sorror, M. L. (2013). How I assess comorbidities before hematopoietic cell transplantation. Blood, 121(15), 2854-2863.*

- Yes  No

97 Specify co-existing diseases or organ impairment (check all that apply)

- Arrhythmia - **Any history of atrial fibrillation or flutter, sick sinus syndrome, or ventricular arrhythmias requiring treatment**
- Cardiac - **Any history of coronary artery disease (one or more vessel-coronary artery stenosis requiring medical treatment, stent, or bypass graft), congestive heart failure, myocardial infarction, OR ejection fraction ≤ 50% on the most recent test**
- Cerebrovascular disease - **Any history of transient ischemic attack, subarachnoid hemorrhage or cerebral thrombosis, embolism, or hemorrhage**
- Diabetes - **Requiring treatment with insulin or oral hypoglycemic drugs in the last 4 weeks but not diet alone**
- Heart valve disease - **At least a moderate to severe degree of valve stenosis or insufficiency as determined by Echo; prosthetic mitral or aortic valve; or symptomatic mitral valve prolapse**
- Hepatic, mild - **Bilirubin > upper limit of normal to 1.5 x upper limit of normal, or AST / ALT > upper limit of normal to 2.5 x upper limit of normal at the time of transplant OR any history of hepatitis B or hepatitis C infection**
- Hepatic, moderate / severe - **Liver cirrhosis, bilirubin > 1.5 x upper limit of normal, or AST / ALT > 2.5 x upper limit of normal**
- Infection - **Includes a documented infection, fever of unknown origin, or pulmonary nodules suspicious for fungal pneumonia or a positive PPD test requiring prophylaxis against tuberculosis. Patients must have started antimicrobial treatment before Day 0 with continuation of antimicrobial treatment after Day 0**
- Inflammatory bowel disease - **Any history of Crohn's disease or ulcerative colitis requiring treatment**
- Obesity - **Patients older than 18 years with a body mass index (BMI) > 35 kg/m<sup>2</sup> prior to the start of conditioning or a BMI of the 95th percentile or higher for patients aged 18 years or younger**
- Peptic ulcer - **Any history of peptic (gastric or duodenal) ulcer confirmed by endoscopy or radiologic diagnosis requiring treatment**
- Psychiatric disturbance - **Presence of any mood (e.g., depression), anxiety, or other psychiatric disorder (e.g., bipolar or schizophrenia) requiring continuous treatment in the last 4 weeks**
- Pulmonary, moderate - **Corrected diffusion capacity of carbon monoxide and / or FEV1 of 66 - 80% or dyspnea on slight activity attributed to pulmonary disease at transplant**
- Pulmonary, severe - **Corrected diffusion capacity of carbon monoxide and / or FEV1 of ≤ 65% or dyspnea at rest attributed to pulmonary disease or the need for intermittent or continuous oxygen during the 4 weeks prior to transplant**
- Renal, moderate / severe - **Serum creatinine > 2 mg/dL or > 177 μmol/L; on dialysis during the 4 weeks prior to transplant; OR prior renal transplantation**
- Rheumatologic - **Any history of a rheumatologic disease (e.g., systemic lupus erythematosus, rheumatoid arthritis, polymyositis, mixed connective tissue disease, or polymyalgia rheumatica, etc.) requiring treatment. (Do NOT include degenerative joint disease, osteoarthritis)**
- Prior malignancy - **Treated at any time point in the patient's past history, other than the primary disease for which this infusion is being performed**

98 Was the recipient on dialysis immediately prior to start of preparative regimen?

- Yes  No  Unknown

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**99 Specify prior malignancy (check all that apply)**

- Breast cancer
- Central nervous system (CNS) malignancy (e.g., glioblastoma, astrocytoma)
- Gastrointestinal malignancy (e.g., colon, rectum, stomach, pancreas, intestine, esophageal)
- Genitourinary malignancy (e.g., kidney, bladder, ovary, testicle, genitalia, uterus, cervix, prostate)
- Leukemia (includes acute or chronic leukemia)
- Lung cancer
- Lymphoma (includes Hodgkin & non-Hodgkin lymphoma)
- MDS / MPN
- Melanoma
- Multiple myeloma / plasma cell disorder (PCD)
- Oropharyngeal cancer (e.g., tongue, buccal mucosa)
- Sarcoma
- Thyroid cancer
- Other skin malignancy (basal cell, squamous)
- Other hematologic malignancy
- Other solid tumor

**100** Specify other skin malignancy: (prior) \_\_\_\_\_

**101** Specify other hematologic malignancy: (prior) \_\_\_\_\_

**102** Specify other solid tumor: (prior) \_\_\_\_\_

**Use results within 4 weeks prior to the start of the preparative regimen, report results from the test performed closest to the start date. Biomarkers according to the augmented HCT comorbidity index Source: Biol Blood Marrow Transplant. 2015 Aug; 21(8): 1418-1424.**

**103** Serum ferritin (within 4 weeks prior to the start of the preparative regimen, use result closest to the start date)

- Known  Unknown

**104** \_\_\_\_\_ ng/mL (µg/L)

**105** Date sample collected: \_\_\_\_\_ - \_\_\_\_\_ - \_\_\_\_\_

**106** Upper limit of normal for your institution: \_\_\_\_\_

**107** Serum albumin (within 4 weeks prior to the start of the preparative regimen, use result closest to the start date)

- Known  Unknown

**108** \_\_\_\_\_  g/dL  g/L

**109** Date sample collected: \_\_\_\_\_ - \_\_\_\_\_ - \_\_\_\_\_

**110** Platelets (within 4 weeks prior to the start of the preparative regimen, use result closest to the start date)

- Known  Unknown

**111** \_\_\_\_\_  x 10<sup>9</sup>/L (x 10<sup>3</sup>/mm<sup>3</sup>)

x 10<sup>6</sup>/L

**112** Were platelets transfused ≤ 7 days before date of test?

- Yes  No  Unknown

**113** Did the recipient have a prior solid organ transplant?

- Yes  No

**Prior Solid Organ Transplant (1)**

Questions: 114 - 116

**114** Specify organ

- Bowel  Heart  Kidney(s)  Liver  Lung(s)  Pancreas  Other organ

**115** Specify other organ: \_\_\_\_\_

**116** Year of prior solid organ transplant: \_\_\_\_\_

**Pre-HCT Preparative Regimen (Conditioning)**

Questions: 117 - 131

**117** Height at initiation of pre-HCT preparative regimen: \_\_\_\_\_  inches  centimeters

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118 Actual weight at initiation of pre-HCT preparative regimen: \_\_\_\_\_  pounds  kilograms

119 Was a pre-HCT preparative regimen prescribed?  
 yes  no

120 Classify the recipient's prescribed preparative regimen (**Allogeneic HCTs only**)  
 Myeloablative  
 Non-myeloablative (NST)  
 Reduced intensity (RIC)

121 Was irradiation planned as part of the pre-HCT preparative regimen?  
 yes  no

122 What was the prescribed radiation field?  
 Total body  
 Total body by intensity-modulated radiation therapy (IMRT)  
 Total lymphoid or nodal regions  
 Thoracoabdominal region

123 Total prescribed dose: (dose per fraction x total number of fractions) \_\_\_\_\_  Gy  cGy

124 Date started: \_\_\_\_ - \_\_\_\_ - \_\_\_\_

125 Was the radiation fractionated?  
 yes  no

126 Total number of fractions: \_\_\_\_\_

### Preparative Regimen (1)

Questions: 127 - 131

Indicate the total prescribed cumulative dose for the preparative regimen:

127 Drug \_\_\_\_\_

128 Specify other drug: \_\_\_\_\_

129 Total prescribed dose: \_\_\_\_\_  mg/m<sup>2</sup>  mg/kg  AUC (mg x h/L)  AUC (μmol x min/L)  CSS (ng/mL)

130 Date started: \_\_\_\_ - \_\_\_\_ - \_\_\_\_

131 Specify administration (busulfan only)  
 Oral  IV  Both

### Additional Drugs Given In the Peri-Transplant Period

Questions: 132 - 140

132 ALG, ALS, ATG, ATS  
 yes  no

133 Total prescribed dose: \_\_\_\_\_ mg/kg

134 Specify source  
 ATGAM (horse)  
 ATG - Fresenius (rabbit)  
 Thymoglobulin (rabbit)  
 Other

135 Specify other source: \_\_\_\_\_

136 Alemtuzumab (Campath)  
 yes  no

137 Total prescribed dose: \_\_\_\_\_  mg/m<sup>2</sup>  mg/kg  mg

138 Defibrotide  
 Yes  No

139 KGF  
 Yes  No

140 Ursodiol  
 Yes  No

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### GVHD Prophylaxis

Questions: 141 - 143

This section is to be completed for allogeneic HCTs only; autologous HCTs continue with question 144.

141 Was GVHD prophylaxis planned?

Yes  No

142 Specify drugs / intervention (check all that apply)

- Abatacept
- Anti CD 25 (Zenapax, Daclizumab, AntiTAC)
- Blinded randomized trial
- Bortezomib
- CD34 enriched (CD34+ selection)
- Corticosteroids (systemic)
- Cyclophosphamide (Cytoxan)
- Cyclosporine (CSA, Neoral, Sandimmune)
- Extra-corporeal photopheresis (ECP)
- Ex-vivo T-cell depletion
- Filgotinib
- Maraviroc
- Methotrexate (MTX) (Amethopterin)
- Mycophenolate mofetil (MMF) (Cellcept)
- Ruxolitinib
- Sirolimus (Rapamycin, Rapamune)
- Tacrolimus (FK 506)
- Tocilizumab
- Other agent

143 Specify other agent: \_\_\_\_\_ (do not report ATG, campath)

### Post-HCT Disease Therapy Planned as of Day 0

Questions: 144 - 146

144 Is additional post-HCT therapy planned?

yes  no

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**Questions 145 – 146 are optional for non-U.S. centers**

**145** Specify post-HCT therapy planned (check all that apply)

- Azacytidine (Vidaza)
- Blinatumomab
- Bortezomib (Velcade)
- Bosutinib
- Brentuximab
- Carfilzomib
- Cellular therapy (e.g. DCI, DLI)
- Crenolanib
- Daratumumab
- Dasatinib
- Decitabine
- Elotuzumab
- Enasidenib
- Gilteritinib
- Ibrutinib
- Imanitib mesylate (Gleevec, Glivec)
- Intrathecal therapy (chemotherapy)
- Ivosidenib
- Ixazomib
- Lenalidomide (Revlimid)
- Lestaurtinib
- Local radiotherapy
- Midostaurin
- Nilotinib
- Obinutuzumab
- Pacritinib
- Ponatinib
- Quizartinib
- Rituximab (Rituxan, MabThera)
- Sorafenib
- Sunitinib
- Thalidomide (Thalomid)
- Other therapy
- Unknown

**146** Specify other therapy: \_\_\_\_\_

### Prior Exposure: Potential Study Eligibility

Questions: 147 - 147

Selecting any option(s) below may generate an additional supplemental form.

**147** Specify if the recipient received any of the following (at any time prior to HCT / infusion) (check all that apply)

- Blinatumomab (Blincyto)
- Gemtuzumab ozogamicin (Mylotarg)
- Inotuzumab ozogamicin (Besponsa)
- Adienne Tepadina®
- Mogamulizumab (Poteligeo)
- None of the above

Mail, fax or email this form to Minneapolis. Fax: 612-627-5895. Email: scanform@nmdp.org.  
Retain the original form at the transplant center.

# ERROR CORRECTION FORM

Sequence Number:

--	--	--	--	--	--	--	--	--	--	--	--	--	--	--	--	--	--	--	--

CIBMTR Recipient ID:

--	--	--	--	--	--	--	--	--	--	--	--	--	--	--	--	--	--	--	--

Initials:

--	--

Today's Date:

		2	0		
Month	Day	Year			

Infusion Date:

		2	0		
Month	Day	Year			

CIBMTR Center Number:

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## Form 2400 R7.0: Pre-Transplant Essential Data

Center:

CRID:

First Name: \_\_\_\_\_

Last Name: \_\_\_\_\_

E-mail address: \_\_\_\_\_

Date: \_\_\_\_ - \_\_\_\_ - \_\_\_\_

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