Instructions for Pre-Transplant Essential Data (Pre-TED) Form (Revision 9)

This section of the CIBMTR Forms Instruction Manual is intended to be a resource for completing the Pre-Transplant Essential Data (Pre-TED) Form (Revision 9).

E-mail comments regarding the content of the CIBMTR Forms Instruction Manual to: CIBMTRFormsManualComments@nmdp.org. Comments will be considered for future manual updates and revisions. For questions that require an immediate response, contact the CIBMTR Customer Service Center.

Pre-Transplant Essential Data (Pre-TED)

The Pre-TED Form is now required for all transplants, including subsequent transplants on the comprehensive report form track.

All transplant centers participating in the CIBMTR must submit a Pre-TED (2400) Form for each allogeneic (related or unrelated) hematopoietic cell transplant (HCT). The Pre-TED is a requirement of the SCTOD for all United States transplant centers when either the stem cell donation or the transplant occurs within the United States. For more information regarding the SCTOD, see General Instructions, Stem Cell Therapeutics Outcomes Database.

Although data regarding recipients receiving autologous HCTs are not required to be submitted as part of the C.W. Bill Young Transplant Program, the CIBMTR is highly committed to collecting data on these recipients for research studies. Centers choosing to report autologous data to the CIBMTR must report on all autologous transplants performed at their center. For more information regarding data reporting for autologous HCT, see HCT.

The Pre-TED may be submitted to the CIBMTR up to two weeks prior to the start of the recipient’s preparative regimen. The Pre-TED is due the day of the HCT (day 0) and is past due if not received by that date.

Helpful Hint
In order to avoid having to make changes to the HCT date, complete the data for the Pre-TED (in FormsNet3SM or on paper), but do not submit the form until the first dose of the preparative regimen is given.

**Consent Status and Baseline Forms**

There has been a change to the functionality of submitting the Pre-Transplant Essential Data (2400), Pre-Transplant Essential Data Disease Classification (2402), and Pre-Cellular Therapy Essential Data (4000) forms. If a consent status has not yet been reported for a recipient, the edit form icon will appear disabled (see Figure 1 below). When the user hovers over the icon, it will display that consent has not yet been reported for that recipient (see Figure 2 below). The user should go to the Consent Tool (see Navigation to the Consent Tool) and document the recipient’s consent status in order to enable the edit icon and allow for completion of the form.

**Figure 1. Disabled Edit Form Icon**

![Disabled Edit Form Icon](image1)

**Figure 2. Hovered Text, Consent Not Yet Reported**

![Hovered Text, Consent Not Yet Reported](image2)

**For recipients receiving a subsequent HCT:**

Transplant centers must submit a Pre-TED for all subsequent HCTs; this includes recipients assigned to the TED Forms and the Comprehensive Report Forms by the form selection algorithm.

For the majority of subsequent HCTs, the recipient will remain on the original follow-up form track assigned by the form selection algorithm. For more information regarding center type and the form selection algorithm, see Section 1 in the Center Reference Guide. A recipient may need to change tracks if enrolled on a study that requires comprehensive forms.
For recipients of multiple transplants, transplant centers are not granted access to the new Pre-TED Form in FormsNet3SM until the Post-TED (Form 2450) or Post-Infusion Data Form (Form 2100) from the previous transplant has been completed.

Transplant centers can use the FormsNet3SM application to determine if a Pre-TED is due by either: 1) accessing the Forms Due Report, or 2) entering the recipient’s unique ID (CRID) in the Recipient Forms Due field.

Links to Sections of the Form:
Q1-23: Recipient Information
Q24-43: Hematopoietic Cellular Transplant (HCT) and Cellular Therapy
Q44-82: Donor Information
Q83-86: Clinical Status of Recipient Prior to the Preparative Regimen (Conditioning)
Q87-120: Comorbid Conditions
Q121-135: Pre-HCT Preparative Regimen (Conditioning)
Q136-144: Additional Drugs Given in the Peri-Transplant Period
Q145-147: GVHD Prophylaxis
Q148-150: Post-HCT Disease Therapy Planned as of Day 0
Q151: Prior Exposure: Potential Study Eligibility

Manual Updates:
Sections of the Forms Instruction Manual are frequently updated. The most recent updates to the manual can be found below. For additional information, select the manual section and review the updated text.

If you need to reference the historical Manual Change History for this form, please click here or reference the retired manual section on the Retired Forms Manuals webpage.

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**Q1-23: Recipient Information**

**Question 1: Date of Birth**
The date of birth is automatically populated based on the value reported in the CRID Assignment tool in FormsNet3. Verify that the date of birth is correct. If an error is noted, correct the CRID Assignment tool and verify that the date of birth has been updated on the Pre-TED Form.
Question 2: Sex
The recipient’s sex is automatically populated based on the value reported in the CRID Assignment tool in FormsNet3. Verify that the recipient’s sex is correct. If an error is noted, correct the CRID Assignment tool and verify that the recipient’s sex has been updated on the Pre-TED Form.

Question 3: Ethnicity
The recipient’s ethnicity is automatically populated based on the value reported in the CRID Assignment tool in FormsNet3. Verify that the recipient’s ethnicity is correct. If an error is noted, correct the CRID Assignment tool and verify that the recipient’s ethnicity has been updated on the Pre-TED Form.

Question 4: Race
The recipient’s race is automatically populated based on the value reported in the CRID Assignment tool in FormsNet3. Verify that the recipient’s race is correct. If an error is noted, correct the CRID Assignment tool and verify that the recipient’s race has been updated on the Pre-TED Form.

Question 5: Race Detail
The recipient’s race detail is automatically populated based on the value reported in the CRID Assignment tool in FormsNet3. Verify that the recipient’s race detail is correct. If an error is noted, correct the CRID Assignment tool and verify that the recipient’s race detail has been updated on the Pre-TED Form.

Question 6: Country of primary residence
Select the recipient’s country of residence. If the recipient’s country of primary residence is Brazil, continue with question 7. If the recipient’s country of primary residence is Canada, continue with question 8. If the recipient’s country of primary residence is the United States, continue with question 9. If the recipient’s country of primary residence is not Brazil, Canada, or the United States, continue with question 10.

Question 7: State of residence of recipient (for residents of Brazil)
If Brazil was selected as the recipient’s primary country of residence, enter the recipient’s state of permanent residence at the time of transplant.

Question 8: Providence or territory of residence of recipients (for residents of Canada)
If Canada was selected as the recipient’s primary country of residence, enter the recipient’s providence or territory of permanent residence at the time of transplant.

Question 9: State of residence of recipients (for residents of USA)
If the United States was selected as the recipient’s primary country of residence, enter the recipient’s state of permanent residence at the time of transplant.

Question 10. NMDP Recipient ID (RID):
The NMDP RID is automatically populated based on the value reported on the CRID Assignment Form (2804). Verify that the NMDP RID is correct. If an error is noted, correct Form 2804 and verify that the NMDP RID has been updated on the Pre-TED Form.

**Question 11: ZIP or postal code for place of recipient’s residence (USA recipients only)**
Enter the five-digit ZIP code in which the recipient resides. Only five digits are required; however, if the ZIP+4 (nine digit) code is available, please report it in this field. The zip or postal code is required for USA residents.

The postal code is optional for Canadian residents. The question can be answered or left blank without error for Canadian residents.

**Question 12: Specify blood type: (for allogeneic HCTs only)**
Indicate the recipient’s blood type as A, B, AB, or O. Blood type is an important characteristic in allogeneic transplant because products may require manipulation to minimize the risk of immune reaction due to incompatibility.

**Question 13: Specify Rh factor: (for allogeneic HCTs only)**
Indicate the recipient’s Rh (rhesus) factor. The Rh factor is an important characteristic in allogeneic transplant because product may require manipulation to minimize the risk of immune reaction due to incompatibility.

**Question 14: Has the recipient signed an IRB / ethics committee (or similar body) – approved consent form to donate research blood samples to the NMDP / CIBMTR?**
The Research Sample Repository contains blood samples from unrelated recipients and/or their adult volunteer donors or cord blood units. Related allogeneic recipients and/or donors will participate at selected transplant centers.

The primary objective of the Research Repository is to make blood samples available for research studies related to histocompatibility and hematopoietic cellular transplantation.

Studies in which these data may be used include:

- Improving the understanding of tissue matching for hematopoietic cellular donors and recipients.
- Determining and evaluating the factors that affect transplant outcomes.
- Studying the distribution of HLA tissue types in different populations (e.g., study tissue typing differences between different racial and ethnic populations to help develop methods to improve tissue matching between donors and recipients, including testing of rare HLA types).
Indicate if the recipient signed an IRB-approved consent form to donate research blood samples to the NMDP / CIBMTR. If **Yes (recipient consented)**, continue with question 15. If **No (recipient declined)**, **Not approached**, or **Not applicable (center not participating)**, continue with question 18.

Blood samples are not submitted for subsequent transplants; however, this question is asked for subsequent transplants. If the recipient previously consented to submit research blood samples to NMDP / CIBMTR, select **Yes (recipient consented)**.

**Question 15: Date form was signed**
Report the date the research sample consent form was signed by the recipient. Do not report the date that the witness or health care professional signed the consent form.

**Questions 16-17: Did the recipient submit a research sample to the NMDP / CIBMTR repository? (Related donors only)**
There are a select number of transplant center participating in the Related Specimen Repository. If your center is one of the participating centers, and the recipient provided a research sample, select **Yes** and provide the recipient ID in question 17. The ID number is located on the bar code that is attached to the sample tube.

If the recipient did not provide a research sample, select **No** and continue with question 18.

**Question 18: Is the recipient participating in a clinical trial? (Clinical trials sponsors that uses CIBMTR forms to capture outcomes data)**
Indicate if the recipient is a registered participant with BMT-CTN, RCI-BMT, USIDNET, COG, and/or another clinical trial sponsor that uses CIBMTR forms to capture outcomes data. If **Yes**, continue with question 19. If **No**, continue with question 24.

- **BMT-CTN**: Blood and Marrow Transplant Clinical Trials Network
- **RCI-BMT**: Resource for Clinical Investigation in Blood and Marrow Transplant
- **PIDTC**: Primary Immune Deficiency Treatment Consortium
- **USIDNET**: United States Immunodeficiency Network
- **COG**: Children’s Oncology Group

**NOTE: Questions 19 – 23 Reporting Participation in More Than One Study**

**FormsNetSM application**: Complete questions 19 – 23 for each study the recipient is participating in by adding an additional instance in the FormsNet application.

**Paper form submission**: Copy questions 19 – 23 and complete for each study in which the recipient is participating.

If the participant is enrolled in multiple studies, even if from the same sponsor, report each study separately.

**Questions 19-20: Study Sponsor**
Select the study sponsor of the clinical trial the recipient is participating in. If the participant is enrolled in multiple studies, even if from the same sponsor, report each study separately.

If the study sponsor is reported as **BMT-CTN**, **RCI-BMT**, or **PIDTC**, continue with question 21.

If the study sponsor is reported as **USIDNET** or **COG**, continue with question 22.

If **Other sponsor** is reported, specify the study sponsor in question 20 and continue with question 22.

**Question 21: Study ID Number**
Select the recipient's Study ID number.

**Question 22: Subject ID**
Enter the recipient's USIDNET, COG, or other sponsor Subject ID.

If the recipient is participating in a BMT-CTN study and the EMMES ID is known, enter it here.

If the recipient is participating in an RCI-BMT study, enter the Subject ID given at the time of successful enrollment.

**Question 23: Specify the ClinicalTrials.gov identification number**
All clinical trials are required to be registered on the clinicaltrials.gov website and will have an associated identification number.

Report the identification number – do not include the letters “NCT,” preceding the digits.

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**Q24 – 43: Hematopoietic Cellular Therapy (HCT) and Cellular Therapy**

**Question 24: 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)**
If, at the time of the current HCT, a second (tandem transplant) or subsequent HCT is planned according to the protocol, check **Yes** even if the recipient does not receive the
planned subsequent HCT. The word “planned” should not be interpreted as: if the recipient relapses, then the “plan” is to perform a subsequent HCT.

If Yes, continue with question 25. If No, continue with question 26.

**Question 25: Specify subsequent HCT planned**
Indicate whether the planned subsequent HCT is **Autologous** or **Allogeneic**.

**Question 26: Has the recipient even had a prior HCT?**
Include all HCTs in the recipient’s history, even if the transplants were not performed at your center. The intent is to capture the full picture of the recipient’s treatment history. If Yes, continue question 27. If No, continue with question 28.

**Question 27: Specify the number of prior HCTs**
Enter the number of prior HCTs for the recipient. An HCT event is defined as an infusion of mobilized peripheral blood stem cells (PBSC), bone marrow, or cord blood. For more information on how to distinguish infusion types [example: HCT versus donor cellular infusion (DCI)], see Appendix D.

For recipients who have received a previous HCT (prior to the HCT for which this form is being completed), the following are examples of how to calculate the number of prior HCTs.

**Example 1:** A recipient was previously transplanted under a protocol that included an infusion of cells over multiple days: day 0, day +1 and day +2. This series of infusions is considered one HCT event (as opposed to three HCT events) and should be counted as **HCT Event #1**.

After receiving the infusion, the recipient had relapse of disease. The recipient is scheduled to receive a subsequent HCT including a preparative regimen. This HCT is **HCT Event #2**. One prior HCT should be reported.

**Example 2:** A recipient previously received an *allogeneic* HCT (**HCT Event #1**). Then, due to delayed neutrophil recovery, the recipient received additional cryopreserved *allogeneic* mobilized PBSC from the original donor, without a preparative regimen (i.e., “boost” – **HCT Event #2**).

After receiving the boost, the recipient had relapse of disease. The recipient is scheduled to receive a subsequent allogeneic HCT with preparative regimen (**HCT Event #3**). Two prior HCTs should be reported.

**Example 3:** A recipient previously received an *autologous* HCT (**HCT Event #1**). Then due to delayed neutrophil recovery, the recipient received additional cryopreserved *autologous* cells without a preparative regimen (i.e., “boost” which is not counted as an HCT event because the intent of the autologous infusion is to treat the graft failure).
The boost is successful, but a few years later the recipient develops a new malignancy. The recipient is scheduled to receive a subsequent autologous HCT with preparative regimen (HCT Event #2). One prior HCT should be reported.

If the allogeneic recipient receives an infusion due to poor graft response, count the infusion as a subsequent HCT. The exception to this is “autologous rescue.” Autologous rescue should not be counted as a separate HCT and the data collection forms will not start over (i.e., the forms will continue from the previous HCT).

**Question 28: Prior HCTs reported to the CIBMTR**
If Unknown is selected for question 28, questions 29 – 32 can be answered to report information regarding prior HCTs; however, these questions are not required to be completed.

If Yes or Unknown, continue with question 33. If No, continue with question 29.

**Questions 29 – 32 Reporting Prior HCTs**

- **FormsNet3SM application:** Complete questions 29 – 32 to report all prior HCTs that have not yet been reported to the CIBMTR by adding an additional instance in the FormsNet3SM application.
- **Paper form submission:** Copy questions 29 – 32 and complete for the prior HCT that has not yet been reported to the CIBMTR.

**Question 29: Date of prior HCT**
Report the date (YYYY-MM-DD) of the prior HCT being reported in this instance. If the exact date is unknown and must be estimated, check the "date estimated" box.

For more information regarding reporting partial or unknown dates, see General Instructions, General Guidelines for Completing Forms.

**Question 30: Was the prior HCT performed at a different institution?**
Indicate if the prior HCT being reported in this instance was performed at another institution. If Yes, report the name and address of the institution in question 31. If No, continue with question 32.

**Question 31: Specify the institution that performed the HCT**
Report the name, city, state, and country of the institution where the recipient’s prior HCT being reported in this instance was performed. These data are used to identify and link the recipient’s existence in the database and, if necessary, obtain data from the other institution where the previous infusion was administered.
**Question 32: What was the HPC source for the prior HCT? (check all that apply)**

Report the cell source(s) for the prior HCT being reported in this instance.

An **Autologous** product has cells collected from the recipient for his / her own use.

An unrelated donor (**Allogeneic, unrelated**) is a donor who shares no known ancestry with the recipients. Include adoptive parents / children or stepparents / children.

A related donor (**Allogeneic, related**) is a blood-related relative. This includes monozygotic (identical twins), non-monozygotic (dizygotic, fraternal, non-identical) twins, siblings, parents, aunts, uncles, children, cousins, half-sibling, etc.

**Questions 33 – 37: Reason for current HCT**

Indicate the reason for the current HCT (check only one). If this was a subsequent transplant, verify that this answer is consistent with the reason for the subsequent transplant reported on the previous series of report forms.

- **Graft failure/insufficient hematopoietic recovery:** Additional stem cells are required because there wasn’t any ANC recovery following HCT (primary graft failure), the hematopoietic recovery indefinitely declined after the initial hematopoietic recovery (secondary graft failure), or hematopoietic recovery was deemed insufficient or too slow for survival following previous high-dose therapy and HCT. If autologous cells are infused for this reason, this is considered autologous rescue; in this case, reporting will continue under the prior HCT date and a new Pre-TED form is not required. If the reason is graft failure after initial recovery or insufficient hematopoietic recovery, also complete question 34.

- **Persistent primary disease:** Additional stem cells are required because of the persistent presence of disease pre- and post-transplant (i.e., complete remission was never achieved following the previous transplant).

- **Recurrent primary disease:** Additional stem cells are required because of relapse primary disease (i.e., complete remission was achieved pre- or post-transplant, but the disease relapsed following the previous transplant). If the reason is recurrent primary disease, also complete question 35. Ensure that the date of recurrent primary disease matches the relapse/progression date reported on the previous transplant’s appropriate follow-up form.

- **Planned subsequent HCT, per protocol:** Additional stem cells are given as defined by the protocol for a subsequent transplant/infusion. This includes all planned subsequent transplants (including triple or quadruple transplants). This transplant is not based upon recovery, disease status, or any other assessment.

- **New malignancy (including PTLD and EBV lymphoma):** Additional stem cells are required because the recipient has developed a new malignancy. This does not include a transformation or progression of the original malignancy for which
the recipient was transplanted. If the reason is a new malignancy, also complete question 36, and attach a copy of the pathology report using the “Add Attachment” feature in FormsNet3. Ensure that the date of diagnosis for the new malignancy matches the date of diagnosis for the new malignancy reported on the previous transplant’s appropriate follow-up form.

**Insufficient chimerism:** In the case of a stable, mixed donor chimerism, the infusion of additional cells (usually lymphocytes and not mobilized stem cells) is typically classified as a DCI. Verify with the transplant physician that the cells given should be reported as a subsequent transplant and that stable, mixed chimerism is the reason for the transplant. However, in the case of declining chimerism – when the percentage of donor cells is sequentially decreasing on several studies, indicating possible impending graft failure – additional stem cells are required. Usually the donor chimerism has fallen below 30-50%.

**Other:** If additional stem cells are given for a reason other than the options listed, select Other and complete question 37.

**Question 38:** Has the recipient ever had a prior cellular therapy? (do not include DLIs)
Include all cellular therapy infusions, except DLIs, in the recipient’s history, even if the infusions were not performed at your center. The intent is to capture the full picture of the recipient’s treatment history.

If Yes, continue question 39. If No or Unknown continue with question 44.

**Question 39: Prior cellular therapy reported to the CIBMTR**
If Unknown is selected for question 39, questions 40 – 43 can be answered to report information regarding prior cellular therapies; however, these questions are not required to be completed.

**Question 39: Were all prior cellular therapies reported to the CIBMTR?**
This should include all cellular therapy infusions (except for DLIs) not performed at your center. If the recipient is a transfer patient, you will be able to see all past infusion dates in the Recipient Information Grid in FormsNet3SM. Contact the Customer Service Center if there are questions.

If Yes or Unknown, continue with question 44. If No, continue with question 40.

**Questions 40 – 43: Reporting Prior Cellular Therapies**
**FormsNet3SM application:** Complete questions 40 – 43 to report all prior cellular therapies that have not yet been reported to the CIBMTR by adding an additional instance in the FormsNet3SM application.
**Paper form submission:** Copy questions 40 – 43 and complete for prior cellular therapy that has not yet been reported to the CIBMTR.
Question 40: Date of the prior cellular therapy
Report the date (YYYY-MM-DD) of the prior cellular therapy being reported in this instance.

For information regarding reporting partial or unknown dates, see General Instructions, General Guidelines for Completing Forms.

Question 41: Was the cellular therapy performed at a different institution?
Indicate if the prior cellular therapy being reported in this instance was performed at another institution. If Yes, report the name and address of the institution in question 42. If No, continue with question 43.

Question 42: Specify the institution that performed the cellular therapy
Report the name, city, state, and country of the institution where the recipient’s prior cellular therapy being reported in this instance was performed. These data are used to identify and link the recipient’s existence in the database and, if necessary, obtain data from the other institution where the previous treatment was administered.

Question 43: Specify the source(s) for the prior cellular therapy (check all that apply)
Indicate the cell source(s) for the prior cellular therapy being reported in this instance. If the product is “off the self” or a “third party donor” product obtained from pharmaceutical companies or other corporate entities, donor type should still be identified.

An Autologous product has cells collected from the recipient for his / her own use.

An unrelated donor (Allogeneic, unrelated) is a donor who shares no known ancestry with the recipient. Include adoptive parents / children or step-parents / children.

A related donor (allogeneic or syngeneic, related) is a blood-related relative. This includes monozygotic (identical twins), non-monozygotic (dizygotic, fraternal, non-identical) twins, siblings, parents, aunts, uncles, children, cousins, half-siblings, etc.

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Question 44 – 82: Donor Information

Question 44: Multiple donors?
Indicate if cells from multiple different donors (multiple CBUs, combinations of other products from different donors) are to be used for this HCT. If **Yes**, continue with question 45. If **No**, continue with question 46.

For example, supplemental infusions should be included when determining if multiple donors were used for this HCT event. An infusion of supplemental cells is often given in conjunction with a preparative regimen for HCT. A supplemental infusion is defined as an infusion of cells given prior to clinical day 0 (of an HCT) for any reason other than to produce engraftment.

For more information on supplemental infusions, see Appendix D.

**Question 45: Specify number of donors**
Report the number of donors used for this HCT. Note that this value should never be “1,” since multiple donors were reported in question 44.

**Questions 46 – 82: Reporting More Than One Donor**

- **FormsNet3SM application**: Complete questions 46 – 82 to report all prior cellular therapies that have not yet been reported to the CIBMTR by adding an additional instance in the FormsNet3SM application.
- **Paper form submission**: Copy questions 46 – 82 and complete for prior cellular therapy that has not yet been reported to the CIBMTR.

**Question 46: Specify donor**
Indicate the donor type for this product and continue with question 49.

An autologous product has cells collected from the recipient for his / her own use.

An unrelated donor (allogeneic, unrelated) is a donor who shares no known ancestry with the recipient. Include adoptive parents / children or step-parents / children.

A related donor (allogeneic or syngeneic, related) is a blood-related relative. This includes monozygotic (identical twins), non-monozygotic (dizygotic, fraternal, non-identical) twins, siblings, parents, aunts, uncles, children, cousins, half-siblings, etc.

**Questions 47 – 48: Specify product type (check all that apply)**
Select from the list of product type(s) for the donor being reported in this instance.

**Questions 47 – 48: Specify Product Type**
Previous CIBMTR forms required two instances to be entered in the donor section when a single donor donated multiple products. **This is no longer required.** Report all product collected from a single donor in the same instance of the donor section.

If “Other product” is indicated, specify the product type in question 48. If your center has a protocol where using “other products” is common, you should consistently be
reporting the same text in the specify field so that the like products can be grouped together.

**Question 49: Is the product genetically modified?**
Genetically modified products include any product where the cells are manipulated via either:
- Gene transfer: a process by which copies of a gene are inserted into living cells in order to induce synthesis of the gene’s product; or
- Transduction: a process by which foreign DNA is introduced into a cell by a virus or viral vector

These techniques alter its gene expression through the insertion of different genes or editing of genes. If more than one product is being infused, indicate if any of the products are genetically modified.

If the infusion is a gene therapy, select **Yes**.

Continue with question 50 if the donor type is a related donor (allogeneic, related).

Continue with question 54 if the donor type is an unrelated donor (allogeneic, unrelated).

Continue with question 77 if the donor type is autologous.

**Question 50: Specify the related donor type**
Indicate the relationship and match between the recipient and the related donor being reported in this instance. When determining the donor’s match/mismatched relationship to the recipient, only consider HLA-A, B, C, and DRB1.

**Syngeneic:**
*Includes:* Monozygotic (identical) twins. Occurs when a single egg is fertilized to form one zygote, which then divides into two separate embryos.  
*Does not include:* Other types of twins or HLA-identical siblings (see below).

Continue with question 55 if relationship and match is **Syngeneic**.

**HLA-identical sibling:**
*Includes:* Non-monozygotic (dizygotic, fraternal, non-identical) twins. Occurs when two eggs are fertilized by two different sperm cells at the same time. This category also includes siblings who aren’t twins but have identical HLA types. The recipient and donor will be allele level matched at HLA-A, B, C, and DRB1.

*Does not include:* Half-siblings (report as **HLA-matched other relatives** if their HLA typing is a match, or **HLA-mismatched relative** if it does not match).

Continue with question 55 if the relationship is **HLA-identical sibling**.
HLA-matched other relative:
Includes: All blood-related relatives, other than siblings, who are HLA matched (e.g.,
parents, aunts, uncles, children, cousins, half-siblings). The recipient and donor will be
allele level matched HLA-A, B, C, and DRB1.

Does not include: Adoptive parents / children or stepparents / children who
are HLA matched.

Continue with question 51 if the relationship is HLA-matched other relative.

HLA-mismatched relative:
Includes: Siblings who are not HLA-identical and all other blood-related relatives who
have at least one HLA mismatch (mismatch can be at the antigen or allele level) (e.g.,
parents, aunts, uncles, children, cousins, half-siblings). The recipient and donor will be
antigen or allele level mismatched at 1 or more loci (HLA-A, B, C, or DRB1).

Does not include: Adoptive parents / children or stepparents / children.

This is the option that should be used for haploidentical transplants.

Continue with question 51 if the relationship is HLA-mismatched relative.

Questions 51 – 52: Specify the biological relationship of the donor to the recipient
Indicate the relationship between the recipient and the related donor being reported in
this instance. If the donor is Other biological relative, specify in question 52 and
continue with question 53.

Question 53: Degree of mismatch (related donors only)
If the donor being reported in this instance is an HLA-mismatched relative, indicate the
degree of mismatch as either HLA-mismatched 1 allele or HLA-mismatched ≥ 2
alleles (does include haploidentical donor) and continue with question 55.

Haploidentical means that one half of the HLA type matches the recipient. This type
of HLA mismatch is common between blood-related parents and children. When
determining the donor’s matched/mismatched relationship to the recipient, only
consider HLA-A, B, C and DRB1.

Question 54: Specify unrelated donor type
Indicate the unrelated donor type. When determining the donor’s match/mismatched
relationship to the recipient, only consider HLA-A, B, C, and DRB1.

Question 55: Did NMDP / Be The Match facilitate the procurement, collection, or
transportation of the product?
Distinguish if the product from the donor being reported in this instance is
an NMDP product or a non-NMDP product. Examples of non-NMDP donor registries
include but are not limited to St. Louis Cord Blood Bank, Anthony Nolan, and StemCyte
International Cord Blood Center. This information is included on the product label, the paperwork accompanying the product, and within the NMDP search/product documentation.

**Question 56: Was this donor used for any prior HCTs? (for this recipient)**
Indicate if the donor reported in question 50 was used for prior HCTs for this recipient. If this is the recipient’s first HCT, select **No**.

If this is an autologous infusion, select **No**.

If this is an unrelated donor (cord blood unit, PBSC, or marrow product), continue with question 59.

If this is a non-NMDP unrelated donor (cord blood unit, PBSC, or marrow product), continue with question 59.

If this is a related donor (PBSC or marrow product), continue with question 60.

**Question 57: Global Registration Identifier for Donors (GRID)**
The Global Registration Identifier for Donors (GRID) was developed by the WMDA to ensure secure, reliable and unambiguous assignment of donors. The GRID standard is a 19-character donor identifier composed of three elements: Issuing Organization Number (ION), Registration Donor Identifier, and Checksum (shown below). This standard will ensure each donor ID is globally unique and will reduce the risk of misidentification of donors or their donations.

**GRID**
The GRID has its own section on the Pre-TED (2400) form now. Therefore, only the 19-character donor identifier needs to be reported. This is essential for proper donor linking and, if done incorrectly, will result in queries being placed on the form.

NOTE: GRIDs from DKMS
If you are receiving a GRID from the DKMS registry, the eighth character is being reported as the letter "O" however, this character should be the number "0". When entering a GRID from the DKMS ensure that the eighth character reported is the number "0".

If the donor is NMDP donor, indicate the GRID number and continue with question 72.

If the donor is a non-NMDP donor, indicate the GRID number and continue with question 63.

**Question 58: NMDP cord blood unit ID**
Report the NMDP Donor ID (e.g., 0000-0000-0). This ID is unique for each donor and is assigned by NMDP. This information is included on the product label, the paperwork accompanying the product, and within the NMDP search/product documentation. Enter the NMDP Donor ID (e.g., 0000-0000-0) and continue with question 72.

**Question 59: Registry donor ID (not applicable for related donors)**
Report the non-NMDP unrelated donor ID. Examples of non-NMDP donor registries include, but are not limited to Anthony Nolan, Australia Bone Marrow Donor Registry, and REDOME. This ID is often located on the product label, the paperwork accompanying the product, and registry-specific search/product documentation. Enter the non-NMDP unrelated donor ID and continue with question 63.

**Question 60: Non-NMDP cord blood unit ID (include related and autologous CBUs)**
Report the non-NMDP cord blood unit ID. Examples of non-NMDP donor registries include but are not limited to: St. Louis Cord Blood Bank and StemCyte International Cord Blood Center. This ID is often located on the product label, the paperwork accompanying the product, and registry-specific search/product documentation. Enter the non-NMDP cord blood ID and continue with question 61.

Note that some cord blood banks can ship their units either through the NMDP or directly to the transplant center. Carefully review the accompanying documentation to determine which is appropriate for your unit. You may wish to consult with your center’s Transplant Coordinator, as he or she will have insight as to how the product was acquired.

**Question 61: Is the CBU ID also the ISBT DIN number?**
Report Yes if the non-NMDP CBU ID is the same as the International Society of Blood Transfusion (ISBT) Donation Identification Number (DIN) and continue with question 63. If the product has an ISBT label on it, the ISBT DIN number is in the upper-left-hand corner and consists of a letter followed by 12 numbers, two sideways numbers, and a letter in a box. Example below:
Please find additional information regarding the ISBT DIN numbers and traceability at http://www.iccbba.org/uploads/22/82/2282aa443bf8a2187880304636814244/IN-003-ISBT-128-for-Blood-Components-An-Introduction-v4.pdf. For example, you may see a barcode with an alphanumeric string below it.

If the CBU ID is not the same as the ISBT DIN number, or it is not known, select No and continue with question 62.

**Question 62: Specify the ISBT DIN number**
Report the ISBT DIN number using the letter, 12 digits, 2 sideways numbers, and the letter in the box.

**NOTE: Registry Code(s)**
- **FormsNet3 application**: Select the appropriate registry code from the drop down directory.
- **Paper form submission**: Use the CIBMTR Hematopoietic Stem Cell Transplant (HCT) Infusion (2006) form to determine the registry’s appropriate match code. Enter the match code listed in brackets.

**Question 63: Registry or UCB Bank ID**
Specify the registry used to obtain the adult donor or umbilical cord blood unit and continue with question 68. The Bone Marrow Donors Worldwide (BMDW) codes have been adopted to avoid submitting the entire name and address of the donor registry. Some common banks that do not list with BMDW have been added to the FormsNet list, including St Louis Cord Blood Bank (SLCBB) and Viacord (VIAC).

The registry code for NMDP donors is USA1 and for NMDP cord units is U1CB.

If the donor was found through DKMS, report the registry that facilitated the HCT. Some registries may be listed more than once with BMDW (one way for marrow/PBSC products and differently for cord blood products). Ensure that the appropriate code for the product was selected because distribution of data depends on the code. If the registry code cannot be determined using the BMDW website, select Other registry and continue to question 65.

**Question 64: Specify other Registry or UCB Bank**
If the BMDW website does not list a match code for the adult donor registry or cord blood bank, provide the registry’s official name in the “Specify other Registry or UCB Bank” field.
Please ensure that the registry you are entering under “other” is not already listed in the pull-down list for question 63. For example, NMDP adult donors, NMDP cords, and New York Cord Bank each have their own entries above in the registry or UCB Bank ID drop down menu.

**Questions 65 – 66: Donor date of birth**
Report if the donor’s / infant’s date of birth is **Known** or **Unknown**. If the donor’s/infant’s date of birth is **Known**, report the date of birth (YYYY-MM-DD) in question 66. If the donor’s/infant’s date of birth is **Unknown**, continue with question 67.

**Questions 67 – 68: Donor age**
Report if the donor’s / infant’s age is **Known** or **Unknown**. If the donor’s/infant’s age is **Known**, report the donor’s/infant’s age at the time of product collection in question 68. Report the age in months if the donor is less than 1 year old, otherwise report the age in years. If the donor’s/infant’s age at collection is **Unknown**, continue with question 69.

**Question 69: Donor sex**
Indicate the donor’s biological sex as **Male** or **Female**. For cord blood units, report the infant’s sex.

**Question 70: Specify blood type (donor) (non-NMDP allogeneic donors only)**
Indicate the donors’ blood type as **A**, **B**, **AB**, or **O**. Blood type is an important characteristic in allogeneic transplant because products may require manipulation to minimize the risk of immune reaction due to incompatibility.

**Question 71: Specify Rh factor (donor) (non-NMDP allogeneic donors only)**
Indicate the donor’s Rh (rhesus) factor. The Rh factor is an important characteristic in allogeneic transplant because product may require manipulation to minimize the risk of immune reaction due to incompatibility.

**Question 72: Donor CMV-antibodies (IgG or Total) (Allogeneic HCTs only)**
CMV is a common virus that infects 50-80% of adults worldwide and is transmitted from person to person through bodily fluids. The virus that causes CMV is part of the herpes virus family and, like other herpes viruses, CMV may be dormant for a period of time before the virus is activated in the host. CMV infections are usually harmless in a healthy immune system and typically cause only mild symptoms, if any. However, if a person’s immune system is seriously weakened (as in an immunosuppressed stem cell recipient) the virus can have serious consequences such as pneumonia, liver failure, and even death.

Most laboratory reports indicate a positive result as **reactive**, and a negative result as **non-reactive**. Occasionally, laboratory reports show a specific antibody titer. In this case, compare the laboratory result to the reported standards to determine if the result was reactive or non-reactive.
If the laboratory reports the results as "inconclusive" or "equivocal," select Indeterminant.

If the laboratory reports a CMV IgM antibody only, not total IgG/IgM or CMV IgG antibody; report the result as Not done.

If the laboratory reports CMV testing by PCR (DNA detection), report the result as Not done. CMV testing by PCR is used to detect the presence of the CMV virus and does not test for prior exposure.

Indicate the test result documented on the laboratory report as either Reactive, Non-reactive, Indeterminant, Not done, or Not applicable (cord blood unit).

Question 73: Has the donor signed and IRB / ethics committee (or similar body) approved consent form to donate research blood samples to the NMDP / CIBMTR? (Related donors only)

Indicate if the related donor signed an IRB-approved consent form to donate research blood samples to the CIBMTR. If Yes (donor consented), continue with question 74. If No (donor declined), Not approached, or Not applicable (center not participating), continue with question 77.

Questions 74: Date form signed

Report the date the research sample consent form was signed by the related. Do not report the date that the witness or healthcare professional signed the consent form.

Questions 75 – 76: Did the donor submit a research sample to the NMDP / CIBMTR repository? (Related donors only)

There are a select number of transplant centers participating in the Related Specimen Repository. If your center is one of the participating centers, and the recipient provided a research sample, select Yes and provide the recipient ID in question 76. The ID number is located on the bar code that is attached to the sample tube.

If the recipient did not provide a research sample, select No and continue with question 77.

Question 77: Specify number of products infused from this donor

Report the number of products infused from the donor being reported in this instance and selected in question 46.

Single Product: CIBMTR defines a single product (i.e., cellular product) as cells collected from a single donor using the same mobilization cycle and collection method regardless of the number of collection days.
Example 1 (multiple bags): A G-CSF stimulated donor had two PBSC collections on subsequent days. The products collected over the two days were divided into four bags. Although the product is contained in multiple bags, this collection is considered a single product, as there was no change in mobilization technique or collection method.

Example 2 (change in mobilization): A G-CSF stimulated donor had a PBSC collection, but the cell count was poor. Plerixafor (Mozobil) was added as part of the mobilization and the donor was re-collected the following day. As the change in mobilization occurred during the same mobilization cycle, these collections are considered a single product.

Multiple Products: For the purposes of this manual, the CIBMTR defines multiple products as cells collected using more than one mobilization technique and/or collection method.

Example 3 (multiple collection methods): A G-CSF-stimulated donor had a PBSC collection and the product was cryopreserved. One month later the donor had a marrow collection; both products were infused at the time of transplant. Each collection is considered a separate product because different collection methods were used. The number of products infused from this donor is two.

Example 4 (re-mobilization): A G-CSF-stimulated donor had a PBSC collection, but the cell count was poor. No further collections were attempted and a week later the donor was re-mobilized with G-CSF and a second PBSC collection was performed. Each collection is considered a separate product due to the re-mobilization of the donor.

Example 5 (two different product types): A cord blood unit is infused at the same time as marrow. Each product type is considered a separate product. The number of products infused is two.

Question 78: Specify the number of these product intended to achieve hematopoietic engraftment

If infusions of additional cells (not intended to product engraftment) were given as a supplemental infusion either prior to the HCT being reported (i.e., prior to clinical Day 0) or shortly after the HCT being reported, the cells must be reported as a product on the Pre-TED Form (Form 2400, question 80) and on a separate Cellular Therapy Product Form (Form 4003).

If additional cells were infused post-HCT, for any reason other than a subsequent HCT or a supplemental infusion as part of the HCT, they should be reported as cellular therapy on the appropriate follow-up form. Reporting the additional cells
(given pre-HCT and not intended to produce engraftment) on the Form 4003 is the only mechanism the CIBMTR has in place to collect this data and ensure that the quality assurance data is reported to the cord blood banks, if applicable.

Report the number of products administered to achieve hematopoietic engraftment.

**Questions 79 – 80:** The following mobilization questions are for autologous HCT recipients only. If other than autologous, continue with question 81.

**Questions 79 – 80:** What agents were used to mobilize the autologous recipient for this HCT? (check all that apply)

Report if any of the following agents listed were used in the mobilization event(s). If Other agent was used, specify the agent in question 80.

- **G-CSF:** granulocyte colony-stimulating factor, filgrastim, Neupogen®
- **Pegylated G-CSF:** pegfilgrastim, Neulasta®
- **Perlixafor:** Mozobil®
- **Combined with chemotherapy:** Systemic therapies used to enhance the stem cell product may include cyclophosphamide or ICE chemotherapy (Ifosfamide, carboplatin, and etoposide) with or without rituximab.
- **Anti-CD20:** rituximab, Rituxan®

**Questions 81 – 82:** Name of product (gene therapy recipients)

Select Other name and specify the gene therapy product name in question 82.

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**Q83 – 86: Clinical Status of Recipient Prior to the Preparative Regimen (Conditioning)**

**Question 83:** What scale was used to determine the recipient’s functional status?
The CIBMTR uses the Karnofsky / Lansky scale to determine the functional status of the recipient immediately prior to the start of the preparative regimen. The Karnofsky Scale is designed for recipients aged 16 years and older and is not appropriate for children under the age of 16. The Lansky Scale is designed for recipients one year old to less than 16 years old.
If the recipient is less than one year old, leave questions 84-85 blank.

**Questions 84 – 85: Performance score prior to the start of the preparative regimen**

Recipient performance status is a critical data field that has been determined to be essential for all outcome-based studies. The CIBMTR uses the Karnofsky / Lansky scale to determine the functional status of the recipient immediately prior to the start of the preparative regimen. For the purposes of this manual, the term “immediately prior” represents the pre-HCT work-up phase, or approximately one month prior to the start of the preparative regimen. In cases where the pre-transplant work-up occurs in months prior to transplant (i.e., the pre-transplant workup occurs more than one month prior to transplant), a documented performance score may be submitted if the recipient does not have a score closer to the start of the preparative regimen, the recipient receives no additional treatment after the date of assessment, and the recipient’s status does not clearly decline.

Select the appropriate performance scale, Karnofsky or Lansky, based on the recipient’s age. Using this scale, select the score (10-100) that best represents the recipient’s activity status immediately prior to the start of the preparative regimen. For an example of the Karnofsky / Lansky scale, see Appendix L.

If a Karnofsky / Lansky score is not documented in the source documentation (e.g., inpatient progress note, physician’s clinic note), data management professionals should not assign a performance score based on analysis of available documents. Rather, a physician or mid-level health care provider (NPs and PAs) should provide documentation of the performance score. Documentation from an RN who has been trained and authorized to determine performance scores may also be used.

The CIBMTR recognizes that some transplant centers prefer to collect and use the ECOG performance score as opposed to the Karnofsky / Lansky score. Although the ECOG and Karnofsky / Lansky performance score systems are based on similar principles, the scales are not the same. For example, the Karnofsky / Lansky scale is described in 11 categories, whereas the ECOG performance status is reported in six categories. Due to the overlap between the two systems, an ECOG score of “one” can represent either “80” or “90” on the Karnofsky/Lansky scale. For centers that collect only an ECOG performance score, CIBMTR will make the following accommodations when auditing the source data:

- Centers collecting ECOG scores should do so using standard practices to ensure accuracy.
- For the purposes of CIBMTR reporting, conversion of ECOG to Karnofsky/Lansky should follow a standard and consistent practice. This practice should be clear and reproducible.

For more information regarding converting an ECOG score to a Karnofsky / Lansky score, see Appendix L.
Question 86: Recipient CMV-antibodies (IgG or Total)

Report the cytomegalovirus (CMV) status of the recipient immediately prior to the start of the preparative regimen. For the purposes of this manual, the term “immediately prior” represents the pre-HCT work-up phase, or approximately one month prior to the start of the preparative regimen. An exception to this definition would apply to a recipient with a documented history of a “reactive” CMV test result. In this case, the CMV test may not be repeated during the pre-HCT work-up phase. Therefore, a timeframe of greater than one month prior to the start of the preparative regimen is acceptable. In cases where the pre-transplant work-up occurs in months prior to transplant (i.e., the pre-transplant workup occurs more than one month prior to transplant), a CMV assessment may be submitted if the recipient does not have an assessment closer to the start of the preparative regimen.

CMV is a common virus that infects 50-80% of adults worldwide and is transmitted from person to person through bodily fluids. The virus that causes CMV is part of the herpes virus family and, like other herpes viruses, CMV may be dormant for a period of time before the virus is activated in the host. CMV infections are usually harmless in a healthy immune system and typically cause only mild symptoms, if any. However, if a person’s immune system is seriously weakened (as in an immunosuppressed stem cell recipient) the virus can have serious consequences such as pneumonia, liver failure, and even death.

Most laboratory reports indicate a positive result as reactive, and a negative result as non-reactive. Occasionally, laboratory reports show a specific antibody titer. In this case, compare the laboratory result to the reported standards to determine if the result was reactive or non-reactive.

If the laboratory reports a CMV IgM antibody only, not total IgG/IgM or CMV IgG antibody, report the result as Not done.

If the laboratory reports the results as “inconclusive” or “equivocal,” select Indeterminant.

Indicate the test result documented on the laboratory report as either Reactive, Non-reactive, Indeterminant, or Not done.

Additional Considerations:

- **Recipients < 6 months**: If the recipient is less than 6 months old, report any positive CMV antibody results as “not done” due to the presence of maternal antibodies. However, in infants greater than 6 months old, positive CMV PCR results indicate a CMV infection, and the results may be reported as “reactive.”
- **Exposure to IVIG**: Exposure to IVIG may result in a false positive CMV antibody result. If the recipient has been exposed to IVIG leading up to HCT (within 3-6 months), indicate the CMV antibody results using the following guidelines:
• If the recipient had a non-reactive CMV antibody result prior to IVIG therapy and then routine CMV PCR results showed no copies of CMV, the CMV antibody may be reported as “non-reactive,” even if the CMV antibody became reactive during IVIG treatment.

• If CMV PCR results quantified copies of CMV DNA (i.e., was positive) during IVIG treatment, the results may be reported as “reactive.”

• If the recipient did not have a CMV antibody test prior to the initiation of IVIG but had a positive antibody test during the IVIG therapy, report “not done.”

• “Not done” should be reported if no CMV antibody tests were done prior to the initiation of IVIG therapy, even if CMV PCR testing was negative during IVIG treatment (because CMV PCR only detects active infection, not prior exposure).

• **Documented history of “reactive” CMV**: In cases where a recipient has a documented history of a “reactive” CMV test and does not have a history of IVIG or blood transfusions from a CMV positive donor, “reactive” should be reported for the CMV status even if the CMV test is repeated during the pre-HCT work-up phase and is “non-reactive”.

• **CMV testing by PCR**: If the laboratory reports CMV testing by PCR (DNA detection) but no CMV antibody testing is done during the pre-transplant work-up or within one month prior to transplant, report the result as “not done.” CMV testing by PCR is used to detect the presence of the CMV virus and does not test for prior exposure.

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**Q87 – 120: Comorbid Conditions**

**Diagnosis of COVID-19 after the start of the preparative regimen**: Questions 87 – 89 are intended to capture COVID-19 (SARS-CoV-2) infections diagnosed prior to the start of the preparative regimen / infusion. If a COVID-19 infection is diagnosed after the start of the preparative regimen, report the COVID-19 diagnosis on the post-infusion follow-up form (2450, 2100, and / or 4100).

**Question 87**: 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?

SARS-CoV-2 is a novel virus belonging to the coronavirus (CoV) family that emerged in December 2019. The disease caused by this new CoV is known as COVID-19 (coronavirus disease 2019). The new virus is highly contagious and was officially
declared a pandemic in March 2020. Transmission is believed to be from person to person through respiratory droplets from coughing and sneezing. Testing for COVID-19 is generally performed on specimens collected from a nasal swab or sputum sample.

Indicate whether or not the patient has ever had a known COVID-19 (SARS-CoV-2) infection, based on a positive test result, at any time prior to the start of the preparative regimen or infusion (if no preparative regimen was given).

If the patient has had a documented COVID-19 (SARS-CoV-2) infection, report Yes and continue with question 88.

If the patient has not had a documented COVID-19 (SARS-CoV-2) infection, report No and continue with question 90.

Question 88: Did the patient require hospitalization for management for COVID-19 (SARS-CoV-2) infection?
Report Yes if the recipient was admitted to the hospital for management of their COVID-19 (SARS-CoV-2) infection. This includes any regular hospital or intensive care unit (ICU) admissions. Otherwise, report No and continue with question 90.

Question 89: Was mechanical ventilation used for COVID-19 (SARS-CoV-2) infection?
The clinical spectrum of COVID-19 varies from asymptomatic or paucisymptomatic forms to clinical conditions characterized by respiratory failure that necessitates mechanical ventilation and support in an intensive care unit (ICU). Mechanical ventilation may impact the recipient’s pulmonary function post-infusion. Report Yes if the recipient was placed on mechanical ventilation for COVID-19 and continue with question 90.

Question 90: Was a vaccine for COVID-19 (SARS-CoV-2) received?
Indicate if the recipient received a vaccine for COVID-19 (one dose without a planned second dose, first dose with planned second dose, second dose, third dose, and / or booster dose) at any time prior to the start of the preparative regimen / infusion. This includes one dose without a planned second dose, first, second, and / or third dose, and a booster dose.

If the recipient did not receive a vaccine for COVID-19 or it is not known if the recipient received a vaccine, select No or Unknown, respectively, and continue with question 95.
Questions 91 – 92: Specify vaccine brand
For the reported dose, specify the vaccine brand the recipient received. If the vaccine brand is not listed, select Other type and specify in question 92.

Questions 93 – 94: Select dose(s) received
For the reported dose, specify the vaccine dose the recipient received prior to the start of the preparative regimen / infusion and report the date when the dose was received. If the exact date is not known, use the process described in the General Instructions, Guidelines for Completing Forms and select Date estimated.

Question 95: Is there a history of mechanical ventilation (excluding COVID-19 (SARS-CoV-2)?
A history of mechanical ventilation may impact the recipient’s pulmonary function post-HCT. Mechanical ventilation is any assisted ventilation on behalf of the recipient. Mechanical ventilation can occur as both an endotracheal tube and ventilator, or as a BIPAP machine with a tight-fitting mask in continuous use. The one exception to BIPAP is CPAP used for sleep apnea, which generally involves overnight use only for patients with documented sleep apnea. Therefore, do not report a CPAP used for sleep apnea, as it does not have the same implications as other forms of mechanical ventilation.

Indications for mechanical ventilation include, but are not limited to:
- Apnea with respiratory arrest (excludes sleep apnea)
- Acute lung injury
- Vital capacity < 15 mL/kg
- Chronic obstructive pulmonary disease (COPD)
- Clinical deterioration
- Respiratory muscle fatigue
- Obtundation or coma
- Hypotension
- Tachypnea or bradypnea

If the recipient was placed on mechanical ventilation at any time prior to this HCT event (excluding mechanical ventilation during surgery) check Yes. If the recipient does not have a history of mechanical ventilation, check No.

Question 96: Is there a history of invasive fungal infection?
Fungal infections play a major role in the clinical outcome of transplant recipients. If the recipient has a history of proven, suspected, or documented invasive fungal infection at any time prior to this HCT, check Yes. If the recipient has not had a history of a proven, suspected, or documented invasive fungal infection, check No. For a subsequent HCT, report any documented significant fungal infections in the recipient’s medical history,
starting with the preparative regimen of the previous HCT to the time prior to the preparative regimen for the current HCT.

Examples of invasive fungal infections include, but are not limited to invasive aspergillosis, zygomycosis and other molds, invasive candidiasis, cryptococcosis, endemic mycosis, other yeasts, and pneumocystosis.

Non-invasive fungal infections such as thrush and nail fungus should not be reported.

For assistance with reporting fungal infections, consult a transplant physician.

**Questions 97 – 98: Glomerular filtration rate (GFR) before start of the preparative regimen (pediatric only)**

The glomerular filtration rate (GFR) estimates how much blood passes through the glomeruli each minute and is used to check how well the kidneys are working. Indicate if the GFR is *Known* or *Unknown*. If the GFR is *Known*, indicate the value for this test.

Testing may be performed multiple times within the pre-transplant work-up period; report the most recent laboratory value obtained. Laboratory values obtained on the first day of the preparative regimen may be reported as long as the blood was drawn before any radiation or systemic therapy was administered.

If the GFR is reported as a range, report the average of the range.

If the GFR is reported as either “< X” or “> X,” report the value as X – 1 or X + 1, respectively.

If the actual GFR result is not available, an estimated GFR may be reported, using the GFR calculator.

**Question 99: Does the recipient have a known complex congenital heart disease? (corrected or uncorrected (excluding simple ASD, VSD, or PDA repair) (pediatric only)**

The intent of this question is to determine the pediatric recipient’s history of any known complex congenital heart disease (corrected or uncorrected). Exceptions for reporting would be any simple ASD, VSD, or PDA repair. Indicate *Yes* if the recipient has known complex congenital heart disease, or *No* if they do not.

**Comorbidities**

Prior to answering question 100, review the list of co-existing disease(s) and/or organ impairments listed under questions 101-106.

**Question 100: 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.)**
Hepatic and Renal Comorbidities

In addition to the guidelines listed on the Pre-TED form, include the following time-specific guidelines when reporting hepatic and renal comorbidities.

**Hepatic Comorbidity:** The assessment of liver function tests (ALT, AST and/or Total Bilirubin) has to include at least 2 values per test on two different days within a period extending between day -24 and the start of the preparative regimen. If only a single value was reported in this time period, use the most recent test performed between days -40 & -25 as the second value. When determining the severity of the hepatic comorbidity, the value closest to the start of the preparative regimen should be used. If the liver function test values closest to the start of the preparative regimen do not meet the criteria specified above, a hepatic comorbidity should not be reported.

**Renal (Moderate/Severe) Comorbidity:** Serum creatinine > 2 mg/dL or > 177 μmol/L, as detected in at least two lab values on two different days within a period extending between day -24 and the start of the preparative regimen. If only a single value was reported in this time period, use the most recent test performed between days -40 & -25 as the second value. If the serum creatinine value closest to the start of the preparative regimen did not meet the criteria specified above, a renal (moderate/severe) comorbidity should not be reported.

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Report **Yes** to question 95 if the recipient has a documented history and/or current diagnosis of any of the following:

- **Documented Medical History**
  - Arrhythmia
  - Cardiac
  - Cerebrovascular disease
  - Inflammatory bowel disease
  - Peptic ulcer
  - Rheumatologic
  - Prior malignancy

- **Current Diagnosis at the Time of Pre-HCT Evaluation**
  - Diabetes
  - Heart valve disease
  - Hepatic, mild
  - Hepatic, moderate/severe
  - Infection
  - Obesity
  - Psychiatric disturbance
  - Pulmonary, moderate
  - Pulmonary, severe
  - Renal, moderate/severe
2 Ejection fraction (EF) ≤ 50% should be reported only if present on most recent test
3 Excluding asymptomatic mitral valve prolapse
4 Including any history of hepatitis B or hepatitis C infection
5 If the PFT lists both a “control” FEV1 and a “post-dilator” FEV1, the “control” FEV1
   should be used to determine if a pulmonary comorbidity is present.
6 Including renal transplantation at any time in the patient’s history

Report all comorbidities including those that are considered complications of the primary
disease for transplant. See examples below.

Examples of complications of the primary disease for transplant that should be reported
as comorbidities:
- A recipient with sickle cell had a stroke prior to HCT, the comorbidity to report
  would be “cerebrovascular disease”.
- A toddler with Hurler Syndrome has cardiomyopathy, cardiac valvular disease
  and an ejection fraction of 45%, the comorbidities to report would be “cardiac” &
  “heart valve disease”.

The intent of this question is to identify serious pre-existing conditions that may have an
effect on the outcome of the HCT. For the purposes of this manual, the term “clinically
significant” refers to conditions that are being treated at the time of pre-HCT evaluation
or are in the recipient’s medical history and could cause complications post-HCT.
Conditions listed in the recipient’s medical history that have been resolved (e.g.,
appendectomy), and/or that would not pose a concern during or after the HCT should
not be reported.

Additionally, for the purposes of this manual, the term “at the time of patient
assessment” is defined as the pre-HCT evaluation period prior to the start of the
preparative regimen. If the recipient does not have a documented history of clinically
significant disease(s) or organ impairment(s), check No and continue with question 107.

For information regarding reporting clinically significant co-existing disease or organ
impairment, see Appendix J.

Questions 101: Specify co-existing disease or organ impairments (check all that
apply)
From the list in question 96, select each clinically significant co-existing disease or
organ impairment for this recipient. The definitions for each of the categories below are
transplantation. Blood, 121(15), 2854-2863.

Arrhythmia: Any history of any type of arrhythmia that has necessitated the
delivery of a specific antiarrhythmic treatment. Examples include, but are not
limited to, bradycardia, tachycardia, atrial fibrillation or flutter, sick sinus
syndrome, and ventricular arrhythmias.
Cardiac (Cardiovascular Disease): Any history of coronary artery disease (one or more vessel coronary artery stenosis requiring medical treatment, stent, or bypass graft), congestive heart failure (regardless of an LVEF >50% at the start of the preparative regimen), myocardial infarction, and / or ejection fraction ≤ 50% (shortening fraction < 26% for pediatric recipients) on the most recent test.

Cerebrovascular disease: Any history of transient ischemic attack, subarachnoid hemorrhage, and / or cerebral thrombosis embolism, or hemorrhage.

Diabetes: Diabetes or steroid-induced hyperglycemia requiring continuous treatment with insulin or oral hypoglycemics in the last 4 weeks.

Heart valve disease: Moderate or severe valve stenosis or insufficiency (mitral, aortic, tricuspid, or pulmonary) as determined by the most recent heart evaluation by an echocardiogram, prosthetic mitral or aortic valve, and / or symptomatic mitral valve prolapse. This does not include a documented medical history of heart valve disease.

Hepatic (mild): Chronic hepatitis, bilirubin > upper limit of normal to 1.5x upper limit of normal, or AST/ALT > upper limit of normal to 2.5x upper limit of normal, or any history of hepatitis B or hepatitis C infection. See note in question 96.

Hepatic (moderate/severe): Liver cirrhosis, bilirubin > 1.5x upper limit of normal, or AST/ALT > 2.5x upper limit of normal. See note in question 96.

Infection: Documented infection, fever of unknown origin, or pulmonary nodules requiring continuation of antimicrobial / antifungal / antiviral treatment after day 0.

Inflammatory bowel disease: Any history of Crohn’s disease or ulcerative colitis requiring treatment.

Obesity: Patients with a body mass index > 35.00 kg/m² or BMI-for-age ≥ 95% (pediatric recipients only) during pre-transplant work-up period. For pediatric recipients, if only the BMI is known, refer to the following link to determine the BMI-for-age: https://www.cdc.gov/growthcharts/.

Peptic ulcer: Any history of peptic ulcer confirmed by endoscopy and requiring treatment.

Psychiatric disturbance: The presence of any mood, anxiety, or other psychiatric disorder requiring continuous treatment during the last four weeks. Examples include, but are not limited to, depression, anxiety, Attention-Deficit Disorder (ADD), Attention-Deficit Hyperactivity Disorder (ADHD), bipolar
disorder, and schizophrenia requiring psychiatric consult or treatment in the last 4 weeks.

**Pulmonary (moderate):** Corrected diffusion capacity of carbon monoxide (e.g., DLCOc, DLCCcorr, DLCO) and/or FEV1 66-80% or dyspnea on slight activity at transplant. Use the Dinakara equation below to determine the DLCOc if only an uncorrected value is provided. For recipients assessed by a postbronchodilator test, only the prebronchodilator FEV1 values are considered for evaluation of pulmonary comorbidity.

**Dinakara Equation:** \( \text{DLCOc} = \frac{\text{uncorrected DLCO}}{0.06965 \times \text{hemoglobin g/dL}} \)

**Pulmonary (severe):** Corrected diffusion capacity of carbon monoxide (e.g., DLCOc, DLCCcorr, DLCO) and/or FEV1 ≤ 65% or dyspnea at rest or requiring oxygen at transplant. Use the Dinakara equation above to determine the DLCOc if only an uncorrected value is provided. or recipients assessed by a postbronchodilator test, only the prebronchodilator FEV1 values are considered for evaluation of pulmonary comorbidity.

**Renal (moderate/severe):** Serum creatinine > 2 mg/dL or > 176.8 μmol/L, or on dialysis at transplant, or prior renal transplantation. See note in question 95. If renal (moderate / severe) comorbidity is selected, complete question 97.

**Rheumatologic:** Any history of systemic lupus erythematosus, rheumatoid arthritis, polymyositis, mixed connective tissue disease, or polymyalgia rheumatica requiring treatment (do NOT include degenerative joint disease, osteoarthritis)

**Prior Malignancy, specify:** Any solid tumor(s) and / or hematologic malignancy(ies) that have been treated at any time point in the patient’s past history. A history of any benign tumor(s) should not be reported.

If the recipient is transplanted for a disease that has transformed from one disease to another, the original malignancy should **not** be reported in this section. Details regarding disease transformation will be captured on the Pre-TED Disease Classification form (Form 2402). For more information regarding disease combinations and transformations, refer to the Common Disease Combinations and Common Disease Transformations tables in the **Primary Disease for HCT** section of the Pre-TED Disease Classification Form (Form 2402).

If prior malignancy, specify is selected, complete question 103.

The physician performing the recipient’s pre-HCT evaluation may use the HCT Co-Morbidity Index (HCT-CI) to document co-morbid conditions (see Appendix J).
Question 102: Was the recipient on dialysis immediately prior to start of preparative regimen?
Indicate if the recipient was dialysis, hemodialysis, or peritoneal dialysis dependent within approximately one month prior to the start of the preparative regimen.

Questions 103 – 106: Specify prior malignancy (check all that apply)
Specify the recipient’s prior solid tumor(s) and / or hematologic malignancy(ies).

If Other skin malignancy is selected, specify the skin malignancy in question 104.

If Other prior hematologic malignancy is selected, specify the hematologic malignancy in question 105.

If Other prior solid tumor is selected, specify the solid tumor in question 106.

Laboratory Values Prior to Start of Preparative Regimen
Complete laboratory values prior to the start of the preparative regimen using the results measured within four weeks prior to the start of the preparative regimen. The following are considered biomarkers according to the augmented HCT comorbidity index.

Questions 107 – 117: Provide last laboratory values recorded just prior to the start of the preparative regimen
These questions are intended to determine the clinical status of the recipient prior to the start of the preparative regimen for stem cell transplantation. Testing may be performed multiple times prior to the start of the preparative regimen; report the most recent laboratory value obtained for each specific test. Laboratory values obtained on the first day of the preparative regimen may be reported as long as the blood was drawn before any radiation or systemic therapy was administered.

For each assessment below, indicate if the result was Known or Unknown prior to the start of the preparative regimen. Indicate the value for each test. If necessary, convert values so they can be reported in the units of measurement available on the form.

Serum ferritin: Ferritin is a protein that stores, transports, and release iron. Iron is toxic to cells, so it is stored within the ferritin protein for use. Ferritin that is too low might be indicative of iron deficiency related anemia. Ferritin that is too high might be indicative of iron overload. It is tracked for some diseases, such as hemaophagocytic lymphohistiocytosis.

Date Sample Collected: Report the date the sample was collected. This date should be before the date of the start of the preparative regimen; however, laboratory values obtained on the first day of the preparative regimen may be reported as long as the blood was drawn before any radiation or systemic therapy was administered.
Upper Limit of Normal for your Institution: Report the upper limit of normal. Normal values may vary by laboratory, so it is important to report the upper limit of normal for each assessment.

**Serum albumin**: Serum albumin is a protein found in the blood. Levels are most often reported on a chemistry panel but may occasionally be found in a separate liver function test report.

Date Sample Collected: Report the date the sample was collected. This date should be before the date of the start of the preparative regimen; however, laboratory values obtained on the first day of the preparative regimen may be reported as long as the blood was drawn before any radiation or systemic therapy was administered.

**Platelets**: Platelet are formed elements within the blood that help with coagulation. A low platelet count, call thrombocytopenia, may lead to easy bleed or bruising. Thrombocytopenia may require platelet transfusions. Indicate if the recipient received a platelet transfusion within 7 days prior to testing.

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**Reporting More Than One Prior Solid Organ Transplant**

**FormsNet3SM application**: Complete questions 18 – 120 for each solid organ transplant by adding an additional instance in the FormsNet3SM application.

**Paper form submission**: Copy questions 118 – 120 and complete for each solid organ transplant.

**Questions 118 – 119**: Did the recipient have a prior solid organ transplant? Indicate if it is **Known** or **Unknown** if the recipient had a prior solid organ transplant. If **Known**, specify the organ transplant. If **Other organ** is reported, specify the organ in question 119. If the recipient did not receive a prior solid organ transplant or it is not known, report **No** and continue with question 121.

**Question 120**: Year of prior solid organ transplant

If a recipient received a solid organ transplant during the reporting period, report the date of the solid organ transplant.

For more information regarding partial or unknown dates, see [General Instructions, General Guidelines for Completing Forms](#).
**Q121 – 135: Pre-HCT Preparative Regimen (Conditioning)**

**Question 121 Height at initiation of pre-HCT preparative regimen**
Report the recipient’s height just prior to the start of the preparative regimen. The intent of this question is to determine the height used when calculating preparative regimen drug doses. This height is usually documented on the transplant orders (for radiation and/or systemic therapy) or admitting orders. Report height to the nearest whole centimeter or inch (round up if 0.5 or greater).

Even if the recipient does not receive a preparative regimen, the height is still required.

**Question 122: Actual weight at initiation of pre-HCT preparative regimen**
Report the recipient’s actual body weight just prior to the start of the preparative regimen. The intent of this question is to report the actual weight at the time the preparative regimen starts (which may be different than the weight used to determine preparative regimen doses). This weight is usually documented on the transplant orders (for radiation and/or systemic therapy) or admitting orders. Report weight to the nearest tenth of a kilogram or pound. Do not report adjusted body weight, lean body weight, or ideal body weight.

Even if the recipient does not receive a preparative regimen, the weight is still required.

**MIBG Therapy:** MIBG therapy given for recipients with neuroblastoma is no longer considered preparative regimen and should not be reported.

**Question 123: Was a pre-HCT preparative regimen prescribed?**
Recipients are generally transplanted under a specific protocol that defines the radiation and/or systemic therapy the recipient is intended to receive as a preparative regimen. This protocol, which may be either a research protocol or standard of care protocol, should be referred to when completing this section.

However, there are instances when a preparative regimen is not given. Examples may include, but are not limited to:
- Primary diagnosis of an immune deficiency.
- Subsequent allogeneic HCT due to loss of, or poor, neutrophil engraftment.

If a preparative regimen is prescribed per protocol, check **Yes**. If a preparative regimen is not prescribed, check **No** and continue with question 132.

For more information regarding the recipient’s preparative regimen, consult a transplant physician or contact CIBMTR Center Support.

**Question 124: Classify the recipient’s prescribed preparative regimen (Allogeneic HCTs only)**
Myeloablative pre-transplant conditioning destroys bone marrow cells using high-dose radiation and/or systemic therapy. It is used to eliminate the recipient's immune system and to leave space in the bone marrow niche for the donated cells. A myeloablative regimen is sometimes used for recipients with non-malignant diseases who require HCT for marrow reconstitution (i.e., immunodeficiencies) or to produce a complete donor chimerism.

Non-myeloablative stem cell transplant (NMA or NST) and reduced-intensity conditioning (RIC) preparative regimens generally use lower doses of radiation and/or systemic therapy to prevent graft rejection and to suppress the recipient’s hematopoietic immune system, but not eliminate it completely. Non-myeloablative protocols rely on the immune cells of the donor to destroy the disease (called graft versus tumor or GVT effect), and typically produces mixed chimerism. NST is a common treatment option for recipients who are older or who have other health problems, as the lower radiation and/or systemic therapy doses are easier for the recipient to tolerate.

In general, RIC includes any regimen that does not meet the criteria for either myeloablative or non-myeloablative regimens.

The determination of the intent of the regimen should be based on the center’s protocol or the opinion of the physician overseeing the care of the recipient. However, if the intent is not specified, the regimen intensity may be reported based on the CIBMTR operational guidelines below.

Table 1. Examples of Myeloablative, Reduced Intensity, and Non-Myeloablative Regimens

<table>
<thead>
<tr>
<th>Myeloablative Regimens</th>
<th>Reduced Intensity and Non-Myeloablative Regimens</th>
</tr>
</thead>
<tbody>
<tr>
<td>• TBI &gt; 500 cGy (single) or &gt; 800 cGy (fractionated)</td>
<td>• TBI ≤ 500 cGy (single) or ≤ 800 cGy (fractionated)</td>
</tr>
<tr>
<td>• Cyclophosphamide + TBI (&gt; 500 cGy (single) or &gt; 800 cGy (fractionated))</td>
<td>• ATG + Cyclophosphamide</td>
</tr>
<tr>
<td>• Cyclophosphamide + Etoposide + TBI (&gt; 500 cGy (single) or &gt; 800 cGy (fractionated))</td>
<td>• BEAM (Carmustine [BCNU], Etoposide, Cytarabine [Ara-C], Melphalan)</td>
</tr>
<tr>
<td>• Busulfan &gt; 7.2 mg/kg IV or &gt; 9.0mg/kg orally</td>
<td>• Busulfan ≤ 7.2 mg/kg IV or ≤ 9.0mg/kg orally</td>
</tr>
<tr>
<td>• Busulfan &gt; 300 mg/m² IV or &gt; 375 mg/m² orally</td>
<td>• Busulfan ≤ 300 mg/m² IV or ≤ 375 mg/m² orally</td>
</tr>
<tr>
<td>• Thiotepa &lt; 10 mg/kg</td>
<td>• Melphalan ≤ 150 mg/m²</td>
</tr>
<tr>
<td>+ Cyclophosphamide</td>
<td>• Fludarabine + Cytarabine</td>
</tr>
<tr>
<td></td>
<td>• Fludarabine + Cyclophosphamide</td>
</tr>
<tr>
<td></td>
<td>• Fludarabine + TBI ≤ 500 cGy (single) or ≤ 800 cGy (fractionated)</td>
</tr>
<tr>
<td></td>
<td>• Thiotepa &lt; 10 mg/kg</td>
</tr>
<tr>
<td></td>
<td>• Treosulfan ≤ 30,000 mg/m² or ≤ 30 g/m²</td>
</tr>
<tr>
<td></td>
<td>• Etoposide + Cyclophosphamide</td>
</tr>
</tbody>
</table>
• Busulfan (>7.2 mg/kg IV or >9.0 mg/kg orally) + Melphalan >150 mg/m²
• Melphalan >150 mg/m²
• Thiotepa ≥ 10 mg/kg
• Treosulfan > 30,000 mg/m² or > 30 g/m²

Preparative Regimen – Intensity
These values represent the total prescribed doses. For example, if a recipient is scheduled to receive Melphalan 100 mg/m² for two days (200 mg/m²), the regimen would be myeloablative because the total prescribed dose is > 150 mg/m².

Indicate whether the intent of the preparative regimen was Myeloablative (to produce marrow ablation or pancytopenia), Non-myeloablative, or Reduced intensity.

Question 125: Was irradiation planned as part of the pre-HCT preparative regimen?
Indicate if irradiation is planned as part of the preparative regimen. If irradiation is not planned, check No and continue with question 131.

Irradiation performed as previous treatment should not be reported in this section. Report irradiation performed as previous treatment on the appropriate Disease Specific Form. Additionally, “radiation boosts,” often given to smaller sites that may have residual malignant cells or to areas that were shielded (i.e., chest wall or lung), should not be reported in this section. Report irradiation boosts administered on the applicable Recipient Baseline Data (2000) Form.

Question 126: What was the prescribed radiation field?
Indicate if the planned irradiation was to Total body, Total body by intensity-modulated radiation therapy (IMRT), Total lymphoid or nodal regions, or Thoracoabdominal region.

Question 127: Total Prescribed dose
Enter the total dose of radiation prescribed. If radiation was prescribed as a single dose, the amount of radiation delivered in the single dose constitutes the total dose. If the radiation was prescribed in fractionated doses, multiply the dose per fraction by the total number of fractions to determine the total dose. Enter the total dose of radiation in either grays (Gy) or centigrays (cGy).

Example:
Radiation Order: TBI, 200 cGy/day for three days (3 doses)
Total dose: 200 cGy x 3 doses = 600 cGy
Report “Total Dose” as: 600 cGy
**Question 128: Date started**
Enter the date the single dose or first fraction of radiation was administered.

**Question 129: Was the radiation fractionated?**
Radiation is either delivered as a single dose or in several treatments (fractions). Radiation is fractionated to increase the loss of diseased cells, as they do not recover as quickly as disease-free cells.

Indicate if the radiation was fractionated. If the radiation was not fractionated, check **No** and continue with question 131.

**Question 130: Total number of fractions**
Enter the total number of fractions (treatments) of radiation that were administered. The recipient may receive more than one fraction per day (hyperfractionation).

The total number of fractions multiplied by the dose per fraction must be equal to the total prescribed dose reported in question 127.

**Reporting Multiple Drugs for Preparative Regimen**
- **FormsNet3SM application**: Complete questions 131 – 135 for each drug given as part of the preparative regimen by adding an additional instance in the FormsNet3SM application.
- **Paper form submission**: Copy questions 131 – 135 and complete for each drug given as part of the preparative regimen.

**Questions 131 – 124: Drugs**

**Preparative Regimen – Drugs**
The following questions report the **prescribed** drug therapy that was part of the preparative regimen. Do not report the dose that was actually given. If the recipient has comprehensive report forms due, the actual dose given will be reported on the Recipient Baseline Form (Form 2000). **Do not include drugs that are intended to offset the side effects of the chemotherapy** (e.g., corticosteroids for nausea, MESNA for hemorrhagic cystitis, etc.).

**Drugs After Transplant**
Occasionally, protocols list drugs that may be given before and after transplant. If the drugs are planned to be given before and after transplant, only the doses given before transplant should be quantified in the preparative regimen section. The doses given after transplant should be reported in the Post-HCT Disease Therapy Planned as of Day 0 or GVHD Prophylaxis section. For example, if bortezomib or rituximab is planned to be given on Days -2, +1, +4, and +7, report the Day -2 dose in the preparative regimen section, and the post-transplant doses as planned post-HCT therapy.

**Drugs during the Peri-Transplant Period**
ATG, alemtuzumab (Campath), defibrotide, KGF, and ursodiol may be given during the
peri-transplant period. Previously, if these drugs were administered prior to Day 0, they were reported in the preparative regimen section of the Pre-TED (2400) Form. However, the Pre-TED (2400) Form has been updated – if these drugs were administered prior to Day 0, report them in Additional Drugs Given in the Peri-Transplant Period section, not in the Pre-HCT Preparative Regimen (Conditioning) section.

The form lists each drug by the generic name. The following website provides the trade names under which generic drugs are manufactured: http://www.rxlist.com/script/main/hp.asp.

The Other drug category should be used only if the drug is not one of the listed options. If an “other” drug is prescribed, list the name of the drug in question 132. Include any intrathecal drugs the recipient received for prophylaxis or treatment of CNS disease within 21 days prior to the start of the preparative regimen. Do not report additional sites of radiation (e.g., cranial boost) in the “other” drug category. If the recipient is assigned to the Comprehensive Report Forms by the form selection algorithm, the additional sites of radiation will be reported on the Recipient Baseline Form (Form 2000). If the recipient is assigned to TED Forms by the form track selection algorithm, the additional sites of radiation will not be reported.

If the Pre-TED is being completed for a subsequent HCT, do not report therapy that was given to treat the recipient’s disease (between the previous and current planned HCTs) in the preparative regimen section.

If there is a change to the chemotherapy preparative regimen (e.g., from busulfan + fludarabine to melphalan + fludarabine) after the Form 2400 has been submitted, return to the Pre-TED (2400) form and make this correction directly in FormsNet3SM to ensure that the chemotherapy reported reflects the actual chemotherapy regimen given.

Question 133: Total prescribed dose
Report the total dose of each drug as prescribed in the preparative regimen section of the HCT protocol. Do not report the prescribed daily dose. Report the drug doses to the nearest tenth. For paper submission, do not modify the number of boxes or include decimal values. The pharmacy record or Medication Administration Record (MAR) should be used for determining the date the drug was started.

Report the dose units as either mg/m², mg/kg, AUC (mg x h/L), AUC (µmol x min/L), or CSS (ng/mL). If the total prescribed dose is reported in a unit other than those listed, convert the dose to the appropriate unit. See the example below or consult with a transplant pharmacist for the appropriate conversion. If drug doses cannot be converted to the unit listed (e.g., Campath), leave the unit field blank, override the error (using “unable to answer”), and attach a copy of the source document to the Pre-TED using the attachment feature in FormsNet3.

Calculating Total Drug Doses

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Drug doses are calculated either by recipient weight in kilograms (kg) or recipient body surface area (BSA) in m². The HCT protocol will specify “x mg/kg” or “x mg/m²” and the total number of doses to be administered.

For example, if the protocol requires cyclophosphamide at 60 mg/kg x 2 days (i.e., 2 doses), the “total prescribed dose” should be reported as “120 mg/kg.”

Pharmacokinetic testing can be used to determine whether the drug concentration in the bloodstream is appropriate to the dose given. This reflects the speed of absorption and elimination of the drug. These tests are usually performed using the first dose of systemic therapy, or a test dose, where multiple samples are drawn at specific time points following the first dose. The samples are sent to a laboratory that performs the testing to determine the drug concentration. If carboplatin was prescribed, indicate if pharmacokinetic testing was performed to determine the preparative regimen dosing. If it is not known whether or not this testing was performed, consult a transplant physician.

A common example of this situation occurs in the use of busulfan. When pharmacokinetic (pK) testing is performed, the ordered busulfan dose can be calculated from either the AUC dose or daily AUC. If an AUC dose is documented, this can be multiplied by the number of ordered doses in order to calculate the ordered busulfan dose. When a daily AUC is documented, this can be multiplied by the number of days in order to calculate the ordered busulfan dose. See the example below for more information.

**Example – Calculating the ordered dose of Busulfan using AUC dose:** The AUC dose in the example below is 2842 uMol x Min, which was prescribed for a total of 5 doses. The total ordered dose of Busulfan in this scenario should be reported as 14,210 uMol x Min.

<table>
<thead>
<tr>
<th>Description</th>
<th>Result</th>
<th>Units</th>
</tr>
</thead>
<tbody>
<tr>
<td>Area Under the Curve (AUC)</td>
<td>2842 for Dose #1</td>
<td>uMol x Min</td>
</tr>
<tr>
<td>AUC Target</td>
<td>Cumulative 21924</td>
<td>uMol x Min</td>
</tr>
<tr>
<td>AUC Estimated Average Exposure</td>
<td>[See comments]</td>
<td>uMol x Min</td>
</tr>
<tr>
<td>Clearance Rate</td>
<td>5.45</td>
<td>ml/min/kg</td>
</tr>
<tr>
<td>Recommended Dosing Type</td>
<td>Q24</td>
<td></td>
</tr>
<tr>
<td>Dose recommended starts at dose #</td>
<td>2</td>
<td></td>
</tr>
<tr>
<td>Dose recommended ends at dose #</td>
<td>5 [See comments] ★</td>
<td></td>
</tr>
</tbody>
</table>

In some cases, a “test dose” of the drug is given before the actual preparative regimen is started, and this dose is used for acquiring drug levels that are used to adjust the dose that will be used in the preparative regimen. In other situations, the first dose of the drug is given in the usual fashion as part of the preparative regimen. After this first dose, serum drug levels are drawn and sent to a reference lab. The drug is continued at the starting dose until the lab results are reported and adjustment is made to later doses.
When a drug is used for the preparative regimen where pharmacokinetics will be tested, it is important to distinguish whether the testing will be done with a “test dose” before beginning the preparative regimen or using the first dose of the preparative regimen. The reporting of the dosing for the CIBMTR forms depends upon this distinction. This helps distinguish whether the dose is part of the therapeutic regimen, or not.

1. A test dose was given > 24 hours prior to the intended therapeutic dosing.
   • **Example:** A patient with AML underwent allogeneic HCT from a sibling; busulfan and cyclophosphamide were used as the preparative regimen. The patient presented to clinic 9 days before the HCT, where a dose of busulfan at 0.5 mg/kg was given intravenously. Blood samples were drawn for the next 6 hours, after which the patient left the clinic. His samples were sent to a lab, results were returned the next day, and an adjusted dose of busulfan was calculated. He returned to the hospital 6 days before HCT and began to receive busulfan at the adjusted dose intravenously for 4 days, followed by cyclophosphamide, and proceeded to receive his cells. Since he received 0.5 mg/kg as a “test dose,” this would not be reported in his total preparative regimen dose.
   
   If a test dose was given, where the dose was distinct from the therapeutic dosing preparative regimen (often 1-2 or more days prior to the initiation of regular dosing), the following should be reported:
   • On the Pre-TED (2400) form, the total prescribed dose per protocol would NOT include the test dose.
   • On the Baseline (2000) form, the start date of the chemotherapy agent should be reported as the date the first therapeutic dose was administered. The actual dose received would NOT include the test dose.

2. The first dose of therapeutic dosing is used for monitoring.
   • **Example:** A patient with MDS received an allogeneic HCT from an unrelated donor; busulfan and fludarabine were used as the preparative regimen. She was admitted to the hospital 7 days before her HCT and received a dose of busulfan at 0.8 mg/kg IV at 6:00 AM. Serum samples were drawn every 30 minutes until the next dose of Busulfan at 0.8 mg/kg IV was given at 12:00 noon. Her blood was sent to a reference lab, and she continued to receive busulfan every 6 hours. On day -6, the lab called with her drug levels, and it was determined that the current dose was correct. No adjustment was made, and she completed all 16 doses of busulfan. Since the dose of busulfan (0.8 mg/kg) that was used for drug testing was ALSO her first dose of the preparative regimen, it should be included in the amount of drug that was given for preparative regimen. The total prescribed dose per protocol should be reported as “13 mg/kg.” (0.8 mg/kg x 16 doses = 12.8 mg/kg rounded to 13 mg/kg).
   
   If the first dose of the preparative regimen was used to determine pharmacokinetics, the following should be reported:
   • On the Pre-TED (2400) form, the total prescribed dose per protocol would include the dose used for monitoring.
On the Baseline (2000) form, the start date of the chemotherapy agent should be reported as the date the first dose was administered. The actual dose received would include the dose used for monitoring.

Test doses must be reported consistently at your center. Since most centers follow a consistent approach to pharmacokinetic testing, it should be straightforward for the center to adopt a consistent approach to the reporting of test doses.

**Question 134: Date started**
Enter the date when the first dose of the preparative regimen drug was administered. The pharmacy record or Medication Administration Record (MAR) should be used for determining the date the drug was started.

**Question 135: Specify administration (busulfan only)**
Report the busulfan administration route as either **Oral**, **IV**, or **Both**.

<table>
<thead>
<tr>
<th>Question Number</th>
<th>Date of Change</th>
<th>Add/Remove/Modify</th>
<th>Description</th>
<th>Reasoning (if applicable)</th>
</tr>
</thead>
<tbody>
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</table>

**Q136 – 144: Additional Drugs Given in the Peri-Transplant Period**
Drugs may be given during the peri-transplant period to prevent transplant-related complications, such as liver injuries or to facilitate engraftment.

**Questions 136 – 144: Drugs**
For each agent listed, indicate whether the drug was administered during the peri-transplant period to prevent transplant-related complications or facilitate engraftment, and any additional question(s) for each drug administered.

**ALG (Anti-Lymphocyte Globulin), ALS (Anti-Lymphocyte Serum), ATG (Anti-Thymocyte Globulin), ATS (Anti-Thymocyte Serum):**
Serum or gamma globulin preparations containing polyclonal immunoglobulins directed against lymphocytes. These drugs are usually prepared from animals immunized against human lymphocytes. Report the total dose *prescribed* in question 137 and the animal source in question 138. If **Other** is selected, specify the source in question 139.

**Alemtuzumab (Campath):** Antibody preparations that are infused in the recipient. Report the total dose *prescribed* to the nearest tenth and specify the units of measurement in question 141.
Defibrotide: Antithrombotic agent used to prevent veno-occlusive disease.

KGF (keratinocyte growth factor): Alternate names: palifermin, Kepivance. KGF acts to stimulate the growth of cells that line the surface of the mouth and intestinal tract. KGF may also be given to treat oral mucositis or as GVHD prophylaxis.

Ursodiol: A naturally occurring bile acid used to dissolve small gall stones and to increase bile flow in patients with primary biliary cirrhosis.

<table>
<thead>
<tr>
<th>Question Number</th>
<th>Date of Change</th>
<th>Add/Remove/Modify</th>
<th>Description</th>
<th>Reasoning (if applicable)</th>
</tr>
</thead>
<tbody>
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</tbody>
</table>

Q145 – 147: GVHD Prophylaxis

The following GVHD prophylaxis questions are to be completed for allogeneic HCTs only. Autologous and syngeneic HCTs continue with question 148.

If ATG or Campath were ordered for GVHD prophylaxis prior to or after Day 0, report these drugs in the Additional Drugs Given in the Peri-Transplant Period section of the Pre-TED. Do not report these drugs in the GVHD prophylaxis section.

Question 145: Was GVHD prophylaxis planned?

After allogeneic HCT, specific immunosuppressive therapy may be administered to prevent GVHD or to immunosuppress the host marrow, thereby promoting engraftment of the donor stem cells. Most transplant centers have specific GVHD prophylaxis protocols and graft rejection protocols. Planned agents a recipient receives as a result of these protocols should be included in this section. This answer does not have to match what is reported on the Post-HCT Follow-Up (2100) Form.

Indicate if GVHD prophylaxis was planned at the time of transplant. If GVHD prophylaxis was not planned at the time of transplant, check No, and continue with question 148.

Questions 146 – 147: Specify drugs / intervention (check all that apply)

The prophylactic drug options listed on the form are intended to be administered in a systemic or oral form. If the recipient received one of the listed drugs in a topical form, select the Other agent option and specify the drug in question 147.

Product Manipulation for GVHD Prophylaxis

In question 146, be sure to report any product manipulation done.
for GVHD Prophylaxis. Product manipulation is not captured anywhere else on this revision of the Pre-TED (2400) Form and any manipulation done for GVHD Prophylaxis should be reported here. An example of product manipulation for GVHD prophylaxis is T-cell depletion.

The Pre-TED Form lists the generic chemotherapy drug names. The following website provides the trade names under which generic drugs are manufactured: http://www.rxlist.com/script/main/hp.asp

If GVHD prophylaxis is used for a syngeneic (monozygotic or identical twin) or autologous HCT, attach a copy of the source document using the attachment feature in FormsNet3SM.

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<th>Reasoning (if applicable)</th>
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| Q148 – 150: Planned Post-HCT Disease Therapy Planned as of Day 0

**Question 148: Is additional post-HCT therapy planned?**
If additional post-HCT therapy is planned according to the protocol or standard of care, check **Yes** even if the recipient does not receive the planned therapy. The word “planned” should not be interpreted as: *if the recipient relapses, then the “plan” is to treat with additional therapy*. If additional post-HCT therapy is not planned per protocol, check **No** and submit the form.

**Planned Post-HCT Therapy**
The following post-HCT planned therapy questions are optional for non-U.S. centers.

**Questions 149 – 150: Specify post-HCT therapy planned (check all that apply)**
Indicate if the options listed on the form are intended to be part of the post-HCT planned therapy according to the protocol or standard of care. Select **Other therapy** for other planned therapies and specify the other therapy in question 150.

Examples of when the **Unknown** option would be used include inclusion in a treatment protocol where a trial drug is used and randomized, or if post-HCT therapy is planned, but the specific therapy intended for use is not known pre-HCT.

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Q151: Prior Exposure: Potential Study Eligibility

Question 151: Specify if the recipient received any of the following (at any time prior to HCT / infusion) (check all that apply)
Indicate if any of the following agents were administered to the patient prior to HCT / infusion:

- **Blinatumomab (Blincyto)**: A monoclonal antibody used to treat B-cell acute lymphoblastic leukemia.
- **Gemtuzumab ozogamicin (Mylotarg)**: An antibody-drug conjugate used to treat CD33 positive acute myeloid leukemia.
- **Inotuzumab ozogamicin (Besponsa)**: An antibody-drug conjugate used to treat B-cell acute lymphoblastic leukemia.
- **Adienne Tepadina®**: A specific brand of thiotepa, an alkylating agent used in the conditioning regimen to treat myeloma, lymphomas, acute leukemia and other malignant and non-malignant hematological diseases.
- **Mogamulizumab (Poteligeo)**: A monoclonal antibody used to treat mycosis fungoides or Sezary syndrome (types of cutaneous T-cell lymphoma). It is also being studied in the treatment of other types of cancer.

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Signature Lines:
The FormsNetSM application will automatically populate the signature data fields, including name and email address of person completing the form and date upon submission of the form.