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CHAPTER 1

NETWORK ORGANIZATION
1. NETWORK ORGANIZATION

1.1. Mission Statement and Organizational Overview

The Blood and Marrow Transplant Clinical Trials Network (BMT CTN) was established in 2001 to conduct large multi-institutional clinical trials addressing important issues in hematopoietic cell transplantation (HCT) thereby furthering understanding of the best possible treatment approaches.

Participating investigators in the BMT CTN collaborate through an organization designed to maintain continuity of operations and to facilitate effective communication and cooperation among the units.

The Statistical Center of the Center for International Blood and Marrow Transplant Research (CIBMTR®) at the Medical College of Wisconsin, the Coordinating Center of the National Marrow Donor Program® (NMDP)/Be The Match, and The Emmes Corporation™ (Emmes) comprise the Data and Coordinating Center (DCC).

The National Heart, Lung, and Blood Institute (NHLBI) Project Officer, the National Cancer Institute (NCI) Project Officer, and Principal Investigators from the participating Core Clinical Centers and the DCC comprise the Steering Committee, which is responsible for the prioritization, design, execution, and analysis of Network studies.

BMT CTN trials are conducted in both Core and Affiliate Centers. The latter access trials through the DCC. Selected Affiliate Centers are also represented on the Steering Committee.

The Administrative Manual of Procedures (MOP) describes the Network organization, study policies, and participating center procedures. This chapter provides detailed description of the BMT CTN's organizational structure and defines the roles and purposes of the collaborating units.

The success of a multi-center endeavor depends on the cooperation of the staff in all participating units to perform their tasks and responsibilities in an efficient, effective, and timely manner. The participating units in the BMT CTN (i.e., participating clinical centers, DCC, and Project Officers) are posted on the BMT CTN SharePoint website.

1.2. Steering Committee

The BMT CTN Steering Committee is responsible for the operation of the Network. The Steering Committee formulates and implements all policy decisions related to the work of the BMT CTN and establishes its scientific agenda.

The Steering Committee consists of the following voting members:

- Core Clinical Center Principal Investigators (20) including (one vote each)
  - Chairperson
  - Chair-Elect
  - Vice-Chair
  - Immediate Past-Chair
Representatives of Affiliate Centers selected by the Steering Committee for exemplary performance (see Section 1.3.2.3)

- NHLBI Project Officer
- NCI Project Officer
- DCC Principal Investigators (3) (The Principal Investigator from the CIBMTR is the only voting member)
- NCI’s National Clinical Trials Network (NCTN) BMT Committee Representatives from ECOG-ACRIN, SWOG, Alliance for Clinical Trials in Oncology (Alliance) and Children’s Oncology Group (COG)

The following observers may attend and participate in Steering Committee meetings but do not have voting privileges:

- Other NHLBI and NCI staff members
- Co-Investigators from participating Core and Affiliate Centers
- Clinical Research Associates/Coordinators from participating Core and Affiliate Centers
- Other DCC staff
- Other relevant consultants and contributors by invitation for specific agenda items
- Executive sessions of the Steering Committee are limited to voting members and others specifically invited by the Chair

The terms of the Steering Committee members and Chairperson are as follows:

- Steering Committee Membership is indefinite dependent upon continuing participation of the member’s institution in the Network
- Chairperson is presented by the Nominating Committee for Steering Committee approval and is constrained by the following conventions:
  1. Six-year total term, served in the subsequent order:
     - Two years as Vice-Chair
     - One year as Chair-Elect
     - Two years as Chairperson
     - One year as Immediate Past-Chair
  2. Can serve more than one term but not consecutive terms

The functions of the Steering Committee include:

- Develop Manuals of Procedures (Administrative and Technical)
- Ratify major changes in the Manuals of Procedures
- Review Concept Proposals and appoint Protocol Teams
• Prioritize protocols and set timelines for implementation
• Recommend to the NHLBI and NCI Project Officers changes or modifications in BMT CTN protocols that may be necessary or desirable (but not based on Data and Safety Monitoring Board (DSMB) reports)
• Advise and assist the DCC and the Technical Committees on operational matters
• Resolve operational problems brought to the Executive Committee by Investigators, Clinical Research Associates/Coordinators, laboratories/repositories, or the DCC
• Monitor the performance of all participating centers based on information provided by the DCC. This evaluation includes assessment of accrual, the quality of data reported by center staff, laboratory sample compliance and adherence to all protocols. The Steering Committee advises the NHLBI and NCI Project Officers on the performance of participating centers and may recommend that NHLBI and/or NCI invite new participants or terminate centers showing unsatisfactory performance (see Chapter 5 for additional details).
• Assure study results are reported in the scientific literature in a timely manner
• Review decisions and recommendations of the Publications Committee, as needed
• Assume other responsibilities at the request of the BMT CTN Chairperson or the NHLBI or NCI Project Officers

The Steering Committee meets in person three times a year to monitor the progress of BMT CTN studies and consider special issues that may arise. Additional meetings and conference calls are held as necessary. The Steering Committee does not have access to blinded data from BMT CTN studies.

1.2.1. Steering Committee Chairperson

The Steering Committee elects a Chairperson who is primarily responsible for the scientific direction and administration of the BMT CTN. The Chairperson:
• Develops and maintains, with advice from Steering Committee members, an organizational structure that meets the needs of the Network studies, NHLBI and NCI
• Remains informed of all operational aspects of the studies and, working within the developed organization, formulates policy and takes necessary actions to ensure the smooth operation of all studies
• Collaborates on data monitoring and other issues of importance to the overall conduct of the studies
• Appoints individuals from BMT CTN Core and Affiliate centers to appropriate positions and committees
• Chairs the Steering Committee meetings
• Reviews potentially competing studies at participating centers
Elections are held one year prior to the end of the previous Chairperson's term so that the elected individual may serve two years as Vice-Chair. If the BMT CTN Chairperson is unable to serve because of resignation, serious illness or death the Chair-Elect or immediate Past-Chair will assume Acting Chairperson responsibilities. If the BMT CTN Chairperson is unable to fulfill this obligation for a limited period (up to six months), the Chair-Elect or immediate Past-Chair will serve as Acting Chairperson, and a new election for Chair-Elect will not be held; if the period of time exceeds six months, an election for a new Chair-Elect will be held. Similarly, if the Chair-Elect, Vice-Chair or Immediate Past-Chair are unable to fulfill the remainder of their term for more than six months, an election will be held to fill an appropriate leadership position to ensure there are three Steering Committee leaders in place.

1.3. Clinical Centers

1.3.1. Core Clinical Centers

There are 20 Core Clinical Centers (some of which are consortia of two or more centers) with cooperative agreements from the NHLBI and NCI to participate in the BMT CTN. Core Center Principal Investigators have voting representation on the Steering Committee (see Section 1.2) and have responsibility for chairing Protocol Teams, Administrative Committees, and Technical Committees (see Section 1.3.2.3). If a Core Center’s Principal Investigator is unable to serve because of resignation, serious illness or death, the Center will nominate a new Principal Investigator (PI) with approval by the NHLBI.

Participating Core Clinical Centers are responsible for recruiting, examining, and treating study participants and for collecting all clinical, laboratory, demographic, and other data required by each BMT CTN study. The Principal Investigator for each Core Clinical Center is directly responsible for ensuring that all aspects of BMT CTN protocols are followed. Other key center staff includes other physicians, Co-Investigators, Clinical Research Associates/Coordinators (CRAs/CRCs) and related staff. The Principal Investigator of the Core Clinical Center may designate another individual at his or her center to serve as Lead Investigator for specific BMT CTN studies. This individual then assumes responsibility for ensuring that all aspects of the relevant protocol(s) are followed.

Core Clinical Center staff carry out the provisions of the Manuals of Procedures and BMT CTN protocol(s). They are responsible for registering and maintaining follow-up of all enrolled study participants. The responsibilities of the Principal Investigator and Clinical Research Coordinator are further defined in Chapter 9.

1.3.2. Affiliate Clinical Centers

The BMT CTN is a national resource for the advancement of knowledge and understanding in the field of HCT. Participation in Network trials may be open to qualified centers other than Core Clinical Centers through subcontracts with the DCC.

1.3.2.1. Participation in Protocols

BMT CTN protocols will be open to Affiliate Centers who:
• Meet the center qualifications required for the protocol and are approved by the Protocol Team
• Are either FACT-accredited (or pending), or an NMDP/Be The Match participant, or an approved transplant center in an NCI-funded National Clinical Trials Network (NCTN) Group
• Agree to register all transplant recipients (both on and off protocol) through the CIBMTR Statistical Center for the duration of the protocol
• Meet quality assurance standards of the Network

The DCC will actively recruit appropriate Affiliate Centers for Network protocols within the limits of financial resources and in accordance with accrual needs of each protocol. Affiliate Centers will be subject to the same quality assurance procedures as Core Centers. Per patient fees for Core and Affiliate Centers will be decided on a per protocol basis.

Each Protocol Team is encouraged to have one representative from an Affiliate Center that is committed to enrolling significant numbers of study participants on the relevant trial. Selection of an appropriate Affiliate Center representative is at the discretion of the Protocol Chair in consultation with NHLBI/NCI representatives. The Protocol Team is formed after the concept report is approved by the Steering Committee. Affiliate Center investigators are also eligible to serve as Protocol Team Co-Chair, if they propose or contribute significantly to the Study Concept.

1.3.2.2. Participation in Committees

Individuals with relevant expertise from Affiliate Centers may be nominated to participate in Technical Committees. Terms are the same as for individuals from Core Centers. Technical Committees are encouraged to have at least one Affiliate Center representative.

1.3.2.3. Representation on Steering Committee

Affiliate Centers are typically not represented on the Steering Committee unless selected for this privilege to formally recognize exemplary performance in advancing the mission of the BMT CTN.

• An Investigator/PI from any Affiliate Center which enrolls patients on two or more BMT CTN trials with total accrual of at least 12 patients during the previous year is eligible to be named to the Steering Committee. At least 50% of the patients must be enrolled on BMT CTN-led studies.

• The term of membership for Affiliate Center representatives is two years, with the possibility of annual renewal if enrollment continues to be at least 12 patients per year with participation and enrollment to at least two studies, and as long as audited data meet Network requirements

• These positions hold full voting and participation privileges

• These members, or designee from their centers, are eligible to chair both Technical and Protocol Committees (see Section 1.3.2)
• The BMT CTN will reimburse travel expenses for one investigator for each eligible Affiliate Center to attend Steering Committee meetings (other than those held in conjunction with national meetings)

1.4. **Data and Coordinating Center (DCC)**

The DCC plays a key role in developing and facilitating study protocols and is responsible for statistical planning and the collection of quality data from participating clinical centers. DCC functions are performed by a consortium that includes:

- The Statistical Center of the CIBMTR
- The Coordinating Center of the NMDP/Be The Match
- The Emmes Corporation

These three organizations have both separate and overlapping responsibilities for BMT CTN operations as shown by Exhibit 1-4-1 and Exhibit 1-4-2.

DCC responsibilities include the following:

- Maintaining a computerized roster of participants with relevant contact information, BMT CTN roles and organizational affiliations
- Collaborating with the Steering Committee in developing study protocols, procedures, reports, manuscripts and Manuals of Procedures
- Scheduling meetings and conference calls, determining site locations for meetings and providing travel arrangements for meetings
- Coordinating communications among participating centers
- Coordinating and supporting the work of Protocol Teams, including:
  - Designating a Protocol Statistician for each BMT CTN study who has primary responsibility for statistical design and analysis
  - Designating a Protocol Officer for each BMT CTN study who has primary responsibility for keeping the Protocol Team informed about the progress of the trial and providing scientific oversight to the protocol development process
  - Designating a Medical Monitor for each BMT CTN study who reviews safety issues, including protocol-specified safety provisions (e.g., assessment of stopping rules) and unexpected serious adverse events (SAEs)
  - Designating a Protocol Coordinator for each BMT CTN study who has responsibility for overseeing all aspects of developing the protocol document and serves as primary site liaison
  - Designating a Patient and Health Professional Services representative to coordinate development of patient education materials and developing resource materials to handle questions regarding BMT CTN studies
• Coordinating the implementation of BMT CTN studies including:
  - Developing master agreements and protocol riders with Core and Affiliate Centers
  - Identifying, developing contracts and coordinating communications with industry contributors for BMT CTN-led studies
  - Identifying, developing contracts and coordinating communications with suitable pharmacies, laboratories and repositories for support of BMT CTN studies
  - Coordinating the training and certification of clinical center staff in standardized data collection and BMT CTN quality control procedures
  - Reviewing all data submitted on standardized BMT CTN case report forms for completeness and accuracy
  - Communicating with participating centers regarding missing, delayed, incomplete, or erroneous data
  - Monitoring adverse events and participating center reporting
  - Preparing periodic reports on the performance of participating centers
  - Creating computerized data files for BMT CTN data
  - Analyzing study data

• Providing support to the Protocol Review Committee (PRC) and DSMB

• Assisting in preparing scientific reports for publication
<table>
<thead>
<tr>
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<th>CIBMTR</th>
<th>NMDP/BeTheMatch</th>
<th>Emmes</th>
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<tr>
<td><strong>Administrative Functions</strong></td>
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<tr>
<td>Provide overall scientific/administrative leadership</td>
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<td>Develop statistical methodology&lt;sup&gt;1&lt;/sup&gt;</td>
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<td>Recruit, manage, and train pool of physician Medical Monitors</td>
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<td>Develop Manuals of Procedures / Standard Operating Procedures (SOPs)</td>
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<td>Facilitate meeting logistics (including site location, travel arrangements, conference calling, travel reimbursement)</td>
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<td>Coordinate meeting materials&lt;sup&gt;2&lt;/sup&gt;</td>
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<td>Manage general and study-specific electronic communications (including clinicaltrials.gov posting, numbered memoranda, websites)</td>
<td>Lead</td>
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<td>Maintain master rosters</td>
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<td>Prepare protocol budgets and track protocol-specific financials</td>
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<td>Monitor overall budget and subcontracts</td>
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<td><strong>Trials Development &amp; Management</strong></td>
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<td>Develop / review concepts&lt;sup&gt;3&lt;/sup&gt;</td>
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<td>Develop protocols&lt;sup&gt;3&lt;/sup&gt;</td>
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<td><strong>Protocol Team</strong></td>
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<td>Serve as Protocol Officer</td>
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<td>Serve as Protocol Coordinator</td>
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<td><strong>Protocol Implementation</strong></td>
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<td>Manage protocol document and all amendments</td>
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<tr>
<td>Identify centers</td>
<td>Lead</td>
<td></td>
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<tr>
<td>Qualify centers (certify centers’ ability to execute protocol)</td>
<td>Lead</td>
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<tr>
<td>Contract with centers</td>
<td>Lead</td>
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<tr>
<td>Identify and contract laboratories / repositories</td>
<td>Lead</td>
<td></td>
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<tr>
<td>Template regulatory forms (1572, financial disclosure, site delegation log)</td>
<td>Lead</td>
<td></td>
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<tr>
<td>Manage data management system (including registration, Web-based data entry, database design, study archive backup, contingency plans)</td>
<td>Lead</td>
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<tr>
<td>Develop Case Report Forms</td>
<td>Lead</td>
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<tr>
<td>Coordinate laboratory and repository functions</td>
<td>Lead</td>
<td></td>
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<tr>
<td>Manage investigational product distribution&lt;sup&gt;4&lt;/sup&gt;</td>
<td>Shared</td>
<td>Shared</td>
<td></td>
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<tr>
<td>Prepare and submit IND/IDE applications and reports to the FDA</td>
<td>Lead</td>
<td></td>
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<tr>
<td>DCC Member Responsibilities</td>
<td>CIBMTR</td>
<td>NMDP/Be The Match</td>
<td>Emmes</td>
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<td>-------------------------------------------------------------------------------------------</td>
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<tr>
<td>Prepare materials and provide Protocol Review Committee and Data and Safety Monitoring Board meeting support</td>
<td></td>
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<td>Lead</td>
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<tr>
<td>Manage site activation process</td>
<td></td>
<td></td>
<td>Lead</td>
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<tr>
<td>Train site personnel</td>
<td></td>
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<td>Lead</td>
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<tr>
<td>Develop informed consent forms and patient materials</td>
<td></td>
<td></td>
<td>Lead</td>
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<tr>
<td>Develop study-specific handbooks and / or SOPs for processes, lab samples, investigational product, and data management</td>
<td>Shared</td>
<td>Shared</td>
<td>Shared</td>
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<tr>
<td>Monitor adverse events, toxicities, and other safety endpoints</td>
<td></td>
<td></td>
<td>Lead</td>
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<tr>
<td>Develop and implement accrual plan</td>
<td>Shared</td>
<td>Shared</td>
<td>Shared</td>
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<tr>
<td>Review performance of centers</td>
<td>Shared</td>
<td>Shared</td>
<td>Shared</td>
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<tr>
<td>Monitor accrual</td>
<td>Shared</td>
<td>Shared</td>
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<tr>
<td>Develop site monitoring plan, conduct monitoring visits, write monitoring reports, and manage corrective action plans</td>
<td></td>
<td></td>
<td>Lead</td>
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<tr>
<td>Monitor data accuracy and conduct data review sessions</td>
<td></td>
<td></td>
<td>Lead</td>
</tr>
<tr>
<td>Prepare reports / manuscripts / coordinate dissemination of results</td>
<td>Shared</td>
<td>Shared</td>
<td>Shared</td>
</tr>
</tbody>
</table>

1 ~75% of PhD Protocol Statisticians are CIBMTR staff members and 25% Emmes staff members; Emmes MS-level statisticians provide primary support for data set preparation and Data Safety and Monitoring Board reports. CIBMTR MS-level statisticians help with data transfer, concept evaluation, accrual plans and assessment of ongoing accrual.

2 NMDP/Be The Match staff members develop agendas and supporting materials and finalize minutes for Steering and Executive Committee meetings and DCC calls; Emmes staff members develop agendas, supporting materials, and reports for the Protocol Review Committee and Data and Safety Monitoring Boards. Both organizations contribute to protocol team call agendas, materials, and minutes.

3 Key personnel from all three entities review protocol concepts. The CIBMTR provides HCT data to assess feasibility; Emmes and CIBMTR statisticians draft statistical plans. Protocol teams include a CIBMTR Protocol Officer, Emmes Protocol Coordinator, and CIBMTR or Emmes Statistician with support from Emmes Safety Monitors and from NMDP/Be The Match Contracts, Patient Services, and Immunobiology staff members.

4 NMDP/Be The Match develops and executes agreements for investigational agents. NMDP/Be The Match and Emmes work together to implement distribution of investigational agents as per the contract and project sites’ needs.

5 NMDP/Be The Match prepares study-specific handbooks and / or SOPs for lab samples and investigational products as well as patient-specific materials. Emmes prepares study-specific site SOPs and activation materials.

6 The accrual plan is drafted by the NMDP/Be The Match Project Manager / Accrual Coordinator based on projected accrual rates from Core and Affiliate Centers, data from the CIBMTR Research Database, and input from the Protocol Team.

7 Accrual, data delinquency, missing values, and data queries in AdvantageEDC (which will transition to Advantage eClinical in the next funding period), data discrepancies, and major and minor protocol violations are tracked by Emmes. NMDP/Be The Match monitors sample collections and lab testing compliance. The CIBMTR monitors data submitted via FormsNet to the CIBMTR which is not captured by AdvantageEDC. Emmes prepares annual center performance reports and NMDP/Be The Match and Emmes prepare quarterly center performance reports that are sent to Core Centers.

8 Emmes monitors and posts daily accrual to each protocol by center. The NMDP/Be The Match Project Manager / Accrual Coordinator surveys centers for estimated accrual to each study and then monitors center accrual against projections. The CIBMTR uses its Research Database to assess and address accrual issues.

9 The CIBMTR Protocol Officer also has a key role in coordinating the Endpoint Review Committee.

10 The Protocol Coordinator, Officer, and Statistician all contribute to preparation of presentations and publications. Emmes provides administrative support to the Publications Committee in its oversight of the publication process. NMDP/Be The Match Contracts staff assures proper acknowledgement of trial contributors. CIBMTR, NMDP/Be The Match and Emmes staff members coordinate, compile, and distribute the annual BMT CTN Progress Report to the research community and the public to provide them with results from presented and published studies and updated information on protocol activity.
1.4.1. The Statistical Center of the CIBMTR
The CIBMTR, located at the Medical College of Wisconsin in Milwaukee and at the NMDP/Be The Match in Minneapolis, Minnesota, is responsible for overall scientific administrative leadership; concept development and review; development of BMT CTN protocols; and, development of reports and manuscripts. CIBMTR staff includes active HCT physicians and professionals with biostatistics, epidemiology, hematology, oncology, statistics, clinical trials and data analysis experience.

CIBMTR staff has responsibility for developing statistical designs and establishing operational and analytical methodology and analyzing data. CIBMTR staff is also responsible for monitoring accrual and identifying potential Affiliate Centers for participation in specific studies. Some specific functions of the CIBMTR staff are:

- Collaborating with the other members of the DCC in developing study procedures, forms, reports (including the BMT CTN Annual Progress Report), manuscripts, MOPs, and BMT CTN protocols
- Assisting in preparation of scientific reports for publication
- Designating a Protocol Statistician who has primary responsibility for design and analysis of studies
- Designating a Protocol Officer and/or Medical Monitor for specific BMT CTN studies as required

Additional details of CIBMTR activities are included in the Standard Operating Procedures (SOPs) maintained at the CIBMTR.

1.4.2. The Coordinating Center of the National Marrow Donor Program (NMDP)/Be The Match
The NMDP/Be The Match, located in Minneapolis, Minnesota, is responsible for development of protocols, medical monitoring duties, development of trial participant advocacy plans, contracting centers, laboratories, repositories, other suppliers, and participation in contract development for Affiliate Centers. NMDP/Be The Match staff includes HCT physicians, professionals in contracts, finance, data management, donor and recipient advocacy and transplant medicine. Some specific functions of the NMDP/Be The Match staff are:

- Collaborating with the other members of the DCC in development of study procedures, reports, manuscripts, MOPs, and study protocols
- Identifying site locations, schedules and providing travel arrangements for meetings
- Scheduling conference calls
- Developing contracts with Affiliate Centers
- Identifying suitable laboratories and repositories for other suppliers for support of BMT CTN studies
- Developing contracts with identified laboratories, pharmacies, repositories and other suppliers
• Developing contracts with third party contributors
• Providing a Laboratory/Repository Manager and a Project Manager (who also serves as an Accrual Coordinator)
• Providing a Protocol Officer and/or Medical Monitor for specific BMT CTN studies as required
• Developing patient and physician educational materials

Additional details of NMDP/Be The Match activities are included in the SOPs maintained at the NMDP/Be The Match.

1.4.3 The Emmes Corporation

The Emmes Corporation, located in Rockville, Maryland, is responsible for developing BMT CTN MOPs, study protocols, and statistical designs, establishing operational and analytical methodology, coordinating study activities, and analyzing data. Emmes is also responsible for developing case report forms, collecting, editing, and storing all data received from participating centers. Emmes staff includes professionals in biostatistics, epidemiology, clinical trials, data processing, administration, and communication coordination.

Some of the specific functions of the Emmes staff are:

• Collaborating with the other members of the DCC in developing study procedures, forms, reports, manuscripts, MOPs, and BMT CTN protocols
• Coordinating communications among participating centers
• Coordinating communications among laboratories, pharmacies, repositories, contributors, and vendors
• Preparing administrative, recruitment, technical, and statistical reports for meetings
• Coordinating the training and certification of clinical center staff in standardized data collection and BMT CTN quality control procedures
• Reviewing all data submitted on standardized BMT CTN case report forms for completeness and accuracy
• Creating computerized data files for BMT CTN data
• Communicating with participating centers regarding missing, delayed, incomplete, or erroneous data and generating related queries
• Monitoring adverse events and verifying that they have been reported appropriately
• Preparing periodic reports on the performance of participating centers
• Preparing meeting materials and providing on-site meeting support for BMT CTN Steering Committee meetings
• Preparing materials for the PRC
• Analyzing study data and preparing reports for the DSMB
• Providing an Operational Statistician with responsibility for design and analysis, statistical monitoring programs, and participation in Endpoint Review Committee of specific protocols
• Providing a Protocol Coordinator, Data Manager and Safety Monitor to coordinate all aspects of a protocol (the Protocol Coordinator may serve as the Data Manager)
• Providing a Project Director
• Assisting in preparing scientific reports for publication

Additional details of Emmes’ activities are included in the SOPs maintained at Emmes.

1.5. National Institutes of Health (NIH) Program Offices

The BMT CTN Program Offices are located in the Division of Blood Diseases and Resources, National Heart, Lung, and Blood Institute (NHLBI) and in the Division of Cancer Treatment and Diagnosis, National Cancer Institute (NCI).

1.5.1. National Heart, Lung, and Blood Institute

The NHLBI is responsible for organizing and providing support for the BMT CTN in accordance with the allocation of resources provided for this program. The NHLBI Project Team consists of NHLBI staff from the Division of Blood Diseases and Resources. The Office of Biostatistics Research, Division of Cardiovascular Sciences, provides statistical expertise and performs diverse functions in planning, designing, implementing and analyzing NHLBI-sponsored studies to the Principal Investigators and DCC.

The NHLBI Project Officer has substantial responsibilities in protocol development, quality control, interim data and safety monitoring, final data analysis and interpretation, preparation of publications, collaboration with awardees, and coordination and performance monitoring. The NHLBI Project Officer may have lead responsibilities in the preparation of some publications and assists in calculation of capitation budget rates. The Project Officer is an active and fully participating member of the Steering Committee.

1.5.2. National Cancer Institute

The NCI is responsible for organizing and providing support for the BMT CTN in accordance with the allocation of resources provided for this program. The NCI Project Team consists of NCI staff from the Division of Cancer Treatment and Diagnosis.

The NCI Project Officer has substantial responsibilities in protocol development, quality control, interim data and safety monitoring, final data analysis and interpretation, preparation of publications, collaboration with awardees, and coordination and performance monitoring. The NCI Project Officer may have lead responsibilities in the preparation of some publications and assists in calculation of capitation budget rates. The Project Officer is an active and fully participating member of the Steering Committee.
1.6. Study Administration

The organizational structure for developing, implementing and completing a BMT CTN study is characterized in Exhibit 1-6.

Exhibit 1-6

1.6.1. Protocol Team

A Protocol Team is appointed for each approved Study Concept. The Protocol Team has primary responsibility for the development of protocol documents. The Team consists of:

- Protocol Chair(s)
- Core and Affiliate Clinical Center Investigators (3 or more)
- Protocol Officer (DCC physician)
- Primary Protocol Statistician (DCC statistician)
• Protocol Coordinator (DCC Coordinator)
• NHLBI and/or NCI representatives (1 each)
• NHLBI Statistician (1)
• DCC Business Representative
• Ad hoc members as deemed necessary by the Protocol Chair

Selection and terms of service are:
• Non-DCC and non-NIH members are appointed by the Executive Committee
• DCC members are appointed by DCC Principal Investigator
• NIH members are appointed by NHLBI and NCI Project Officers
• Protocol team members serve until the study is completed, analyzed and presented or decision is made to discontinue development

1.6.2. Protocol Review Committee
The PRC is appointed by the NHLBI. The PRC is responsible for providing peer review for final draft protocols including Informed Consent(s) and either approving a protocol and consent form(s) or requesting that changes and/or clarifications are made. If the PRC approves the protocol and informed consent form(s), an independent DSMB will review the study protocol, consent form(s), and monitoring plans, focusing on data quality and safety assurance. The DSMB must approve the protocol before distribution to applicable Institutional Review Boards (IRBs) of record, and FDA if appropriate.

The PRC assesses the scientific merit of each protocol and consent form(s) as follows:
• Importance of the question to be addressed
• Need for a multi-center network to meet objectives
• Merit of experimental design, including appropriate controls
• Availability of adequate resources, including medications
• Adequacy and safety of study participant population and number of study participants, including appropriate representation of minorities, women, and children (if applicable)
• Appropriate recruitment strategies
• Adequacy of proposed plans for data acquisition, transfer, management and analysis
• Adequacy of quality control of data collection and monitoring and overall coordination of protocol management
• Description of appropriate plans to train center personnel to accomplish proposed research goals
The PRC includes a Chairperson and members whose experience reflects areas of expertise necessary to evaluate the scientific merit and design of BMT CTN protocols. Consultants may be added on an *ad hoc* basis to the Committee if greater representation of expertise in a specific scientific area is needed.

The PRC Executive Secretary coordinates the review of each study protocol. The Executive Secretary is an NHLBI staff member, other than the Project Officer, and is responsible for working with the PRC Chairperson to ensure the effective and efficient review of research design, specific aims, and outcomes; appropriateness of methods of intervention, measurement, and analysis; and, recommendations for monitoring of safety and data quality.

Individuals are invited to serve on the PRC by the NHLBI Director. Members are required to complete a Conflict of Interest Certification for review and acceptance by the NHLBI prior to serving on the Committee. At the beginning of all meetings the PRC Chairperson or the Executive Secretary will verbally remind the members of the importance of avoiding conflicts of interest and that members must notify the Executive Secretary promptly if any changes occur which may pose a potential conflict of interest.

### 1.6.3 Data and Safety Monitoring Board

Due to the quantity of BMT CTN trials, the BMT CTN has two appointed DSMBs. Each DSMB is an independent board appointed by the NHLBI and/or NCI. Each DSMB is composed of a Chairperson and members with expertise in biostatistics, clinical trials, bioethics, and the specific research area(s) of the Network studies. Consultants may be added to the DSMB to have greater representation of expertise in the relevant scientific fields. All standing members of a DSMB may vote. Consultants have the same voting rights as an official DSMB member when reviewing the protocol.

Each board meets semi-annually or more often if necessary. Members are required to complete a Conflict of Interest Certification for review and acceptance by the NHLBI prior to serving on the Committee. At the beginning of all meetings the DSMB Chairperson or the Executive Secretary will verbally remind the members of the importance of avoiding conflicts of interest and that members must notify the Executive Secretary promptly if any changes occur which may pose a potential conflict of interest.

After PRC approval, the DSMB must approve the protocol and consent form(s) before distribution to IRB(s) of record. Thereafter, the principal role of the DSMB is to regularly monitor the data from the clinical trial, review and assess the performance of its operations, and make recommendations, as appropriate, to the NIH with respect to:

- Benefits/risks ratio of procedures and the burden under which the study participants are placed
- Completeness, quality, and analysis of measurements that are made
- Performance of individual centers (including possible recommendations on actions to be taken regarding any center that performs unsatisfactorily)
- Interim results of the study for evidence of efficacy or adverse effects
• Possible early termination of the study because of early attainment of study objectives, efficacy and/or safety concerns – if applicable, inadequate performance or accrual.

• Desirability of proceeding to the full-scale trial at the completion of the feasibility phase, when applicable

• Possible modifications/amendments to the study protocol and/or consent form(s)

NHLBI appoints an Executive Secretary for the two DSMBs. The Executive Secretary is an NHLBI staff member, other than the Project Officer and PRC Executive Secretary, and is responsible for working with the DSMB Chairperson to ensure the effective and efficient review of research design, specific aims, and outcomes; appropriateness of methods of intervention, measurement, and analysis; and, recommendations for monitoring of safety and data quality. In addition, the Executive Secretary is responsible for scheduling calls and meetings, and maintaining meeting minutes.

1.6.4. NMDP IRB

Predicated on the single IRB mandate from NIH (Policy on the Use of a Single Institutional Review Board for Multi-site Research; June 21, 2016) and the anticipated revised U.S. Common Rule (Federal Policy for the Protection of Human Subjects; January 19, 2017), the BMT CTN DCC and Core Clinical Centers grant renewal included the provision that the BMT CTN will utilize a single IRB of record with the start of the new grant period on July 1, 2017. The NMDP IRB was selected to serve as the Network’s single IRB and will be used for all BMT CTN protocols released after July 1, 2017 and is also used by some of the centers participating in the BMT CTN 1501 and 1503 NMDP IRB Pilot Project. The only exception is for centers outside the U.S. who will still follow their country’s regulatory requirements for ethical board review.

The NMDP is fully accredited by the Association for the Accreditation of Human Research Protection Programs. The NMDP IRB maintains membership that satisfies the requirements of 45 CFR 46 and 21 CFR 56. The members of the NMDP IRB are a diverse group of distinguished healthcare professionals, donor advocates, and patient advocates with expertise in bone marrow transplantation and hematology/oncology. The majority of members are not affiliated with NMDP.

The NMDP IRB conducts the study-specific reviews as required by the regulations. This includes initial review, continuing review, and review of amendments/modifications to previously approved research. In addition, any other study-specific documents submitted to the NMDP IRB are reviewed per the NMDP IRB Standard Operating Procedures (SOPs) and federal regulations. Administrative functions of the NMDP IRB are managed by NMDP Human Research Protection Program staff members.

The NMDP IRB meets monthly. Board members are required to disclose conflicts of interest per the NMDP IRB SOPs. At the beginning of each study review members are also asked if they have a conflict of interest and, if they do, are recused from the review.
1.7. Administrative Committees

1.7.1. Executive Committee

Membership is by virtue of the following roles in the Network:

- Steering Committee Chairperson (1)
- Steering Committee Vice-Chair, Chair-Elect and/or Immediate Past-Chair (2)
- NHLBI and NCI Project Officers (2)
- DCC Principal Investigators (3)

The Executive Committee is responsible for developing Steering Committee agendas and promulgating recommendations for consideration by the Steering Committee. The Executive Committee will also provide direction between meetings of the Steering Committee, and review/approve all ancillary studies. The Executive Committee will provide initial review of proposals submitted for consideration by the BMT CTN and will approve development of a proposed Study Concept. The Executive Committee participates meets monthly via conference calls.

1.7.2. Publication/Presentation Committee

The Publication/Presentation Committee will consist of up to 10 members (excluding ex-officio):

- DCC Principal Investigators (ex officio)
- Core Clinical Center Principal Investigators (appointed)
- Affiliate Clinical Center representative (appointed)
- NHLBI and/or NCI Project Officers (ex officio)

The terms of membership and selection are as follows:

- Core and Affiliate Center members are identified from a slate of candidates put forth by the Nominating Committee and are approved by the Steering Committee
- Core Center representatives are elected for a three-year term and may serve a second three-year term
- Affiliate Center representative are elected for a three-year term
- Two Co-Chairs are elected by the Executive or Nominating Committee from the Core Clinical Center members for a two-year term; may serve more than one term but not two consecutive terms

The Publication/Presentation Committee is responsible for developing publication and presentation policies. All policies must be approved by the Steering Committee before implementation. The Committee reviews all proposed publications and presentations to ensure protection of proprietary information and study participant confidentiality and to determine the public impact of publication and/or presentation of incomplete or premature results.
No participating institution may present or publish individual findings from work performed on study protocols without approval of the Publication/Presentation Committee, NHLBI and NCI.

1.8. Technical Committees

Technical committees are formed to address specific areas of Network activity. A Chairperson is elected by the Executive or Nominating Committee. Additional members are approved by the Steering Committee (see below). NHLBI and NCI Project Team staff, Steering Committee Chairperson, and DCC staff will be additional members on these committees. Each committee is limited to a maximum of ten non-DCC, non-NIH members.

The membership of each committee will be comprised of members as follows:

- DCC Representative (ex officio)
- Core Clinical Center members (appointed)
- Affiliate Center members (appointed)
- NHLBI and NCI Project Officers (ex officio) or their representatives
- BMT CTN 1202 Biomarkers Protocol Chairs (ex-officio: Biomarkers Committee)

The terms and selection of members is as follows:

- Core and Affiliate Center members are identified from a slate of candidates put forth by the Nominating Committee and approved by the Steering Committee
- Representatives are appointed for a three-year term

The current list of Technical Committees includes:

- Biomarkers: The purpose of the Biomarkers Committee is to establish principles for specimen collection (including their use in Ancillary Studies), to review new and current studies for opportunities to collect biological specimens for analysis of potential prognostic markers and to make protocol recommendations regarding immune reconstitution. The Protocol Chairs of the 1202 Biomarkers protocol will serve as ex-officio members on the Committee.
- Clinical Research Associates (CRA) Committee: The purpose of the CRA Committee is to assist in development of case report forms and data collection systems for specific protocols, review and help resolve logistical issues with respect to protocol implementation (e.g., enrollment procedures, study treatment, study procedures, required observations, shipping and receipt of specimen samples and/or study drugs), and review of educational materials for CRAs/CRCs at participating clinical centers. Their findings are provided to the study teams.
- Pharmacy: The purpose of the Pharmacy Committee is to review all BMT CTN protocols for use, administration and potential interactions of pharmaceuticals, develop a pharmacopoeia for commonly used drugs on BMT CTN protocols, and advise Protocol Teams on possible ancillary studies, e.g. pharmacokinetics.
• Special Populations (Pediatric/Human Subjects): The purpose of the Special Populations Committee is to ensure that women, children and minority study participants are considered for inclusion in all appropriate investigative protocols, including those involving non-malignant marrow disorders, developed by the BMT CTN. The Committee is also charged with the responsibility of ensuring that where pediatric study participants are included, consideration for the differences in obtaining informed consent, patient care, and monitoring are appropriately addressed. The committee provides their recommendations to the study team.

• Toxicity and Supportive Care: The purpose of the Toxicity and Supportive Care Committee is to define methods for evaluation of adverse events and toxicities after transplantation; review the evaluation and monitoring requirements for toxicities on BMT CTN protocols; review routine supportive care practices (as determined by this Committee) which are being considered during protocol development to avoid requiring a protocol specific criterion that may limit patient accrual; and, make recommendations to the Protocol Team on itemssuch as:
  - Infection prophylaxis/surveillance
  - Growth factor use
  - Blood product support

1.8.1 Protocol Review Responsibility

The following technical committees must review all BMT CTN protocols at some point in their development but no later than time of submission to the PRC:

• Biomarkers
• CRA
• Pharmacy, if applicable
• Special Populations (Pediatrics/Human Subjects)
• Toxicity and Supportive Care

Each technical committee has a primary DCC representative appointed by the DCC PI who has responsibility for coordinating committee conference calls and meetings. The DCC representative works with the Committee Chair to prepare and distribute an agenda and minutes for each meeting and ensure that the committee’s findings are communicated to the protocol team.

1.9 Ad Hoc Committees

Additional administrative and technical committees are convened as needed. For example, a Nominating Committee is formed to propose candidates for open committee positions. Several technical committees are also convened as needed to discuss new study concepts, provide an update to the Steering Committee on recent advances in the field, or provide input into the Network’s Technical Manual of Procedures. These ad hoc committees include GVHD, Graft Characterization, Cellular Therapy, Late Effects/QOL, Myeloma and Infectious Disease.
1.10. Collaboration with NCTN Groups

1.10.1. Cross-Networks Collaboration

To enhance communication and partnership with the NCI-funded NCTN Groups, the BMT CTN has included NCTN Group representatives on the BMT CTN Steering Committee (see section 1.2.). In addition, there are appointed BMT CTN representatives that serve on the NCTN Disease-Specific Steering Committees. These investigators are responsible for representing the interests of the BMT CTN and the role it plays within the U.S. BMT community. They represent the Network’s scientific agenda and keep the NCTN apprised of current and planned BMT CTN studies and initiatives relevant to the particular committee.

1.10.2. Standard Collaboration Practices

To assure fairness and standardization of practices to be followed when collaborating with the NCTN Groups, the NHLBI and NCI have established a set of guidelines to follow in the case of BMT CTN-led trials or NCTN Group-led trials.

Major areas defined in these documents include:

- Preserving the enrollment credit system of the NCI
- Establishing a collaborative per-patient payment system
- Opening trials through the NCI Clinical Trial Support Unit (CTSU)
- Expediting review by a single scientific review committee, either the NHLBI-appointed PRC or the NCI Cancer Therapy Evaluation Program PRC, and the leading group’s DSMB
CHAPTER 2

STUDY CONCEPT DEVELOPMENT AND APPROVAL
2. STUDY CONCEPT DEVELOPMENT AND APPROVAL

2.1. Developing a Proposed Study Concept

Proposals for clinical trials may be submitted from members of Core or Affiliate Clinical Centers or others outside the Network. The efficient development of a proposed study idea into a document is facilitated by the DCC, primarily the CIBMTR, in advance of the BMT CTN Steering Committee review. This chapter provides detailed descriptions of the process, timeline and documentation required to develop protocol ideas received by the BMT CTN.

2.2. Contents of a Study Concept Form

All ideas for studies must be submitted to the DCC on a Study Concept Form. The Form is posted on the BMT CTN publicwebsite. The Study Concept Form includes the following information:

- Submitting individual's name and affiliation
- Submitting individual's contact information
- Proposed study title
- Preliminary data and background
- Hypotheses to be tested
- Primary outcome
- Secondary outcomes
- Potential for laboratory, Quality of Life (QOL), and/or ancillary studies
- Study design, accrual and follow-up periods
- Patient population, including diagnosis and disease state, type of transplant, other inclusion and exclusion criteria
- Proposed intervention
- Proposed control therapy (if applicable)
- Statistical section including sample size calculations, and important variables for consideration in stratification (if applicable)
- Necessity for a FDA Investigational New Drug (IND) or Investigational Device Exemption (IDE)
- Central Pharmacy requirements
- Central Lab requirements
- Special specimen collection consideration
- Possible supplemental funding sources
- Potential Networks/NCTN Group involvement
2.3. Other Considerations of a Study Concept Form

2.3.1. Feasibility

A CIBMTR statistician and DCC physician will apply the primary inclusion and exclusion criteria of the study proposal to the CIBMTR and/or NMDP/Be The Match database(s) to determine the number of participants recorded in the database that would meet the primary eligibility criteria of the trial that have received Clinical Centers. Modifications to the inclusion and/or exclusion criteria may be proposed to increase the number of potentially available study participants.

2.3.2. IND or IDE Requirements

As part of the preparation of the Study Concept Report, the DCC identifies any aspect of the protocol that may require an IND or IDE from the FDA. The Report will also specify whether an IND or IDE is currently held by another party.

2.3.3. Competing Protocols

The DCC will review active and proposed clinical trials that may compete for the proposed study participant population. The DCC will describe the impact of the competing study on accrual to the proposed study based on accrual targets for the competing study.

The NCI NCTN Group BMT Committee Chairs (see Section 1.2) will advise the DCC of any NCI NCTN Group planned or active trials that may compete with the proposed study.

2.3.4. Central Pharmacy/BMT CTN Specimen Repository/Specialized or Centralized Laboratory Testing

The investigator proposing the study, in consultation with the DCC, will:

- Address the need for a central pharmacy
- Identify any samples that need to be collected for central review or for storage in the BMT CTN repository
- Determine whether centralized or specialized laboratory services are required to conduct the trial

Upon receipt of a Study Concept Form, DCC staff is responsible for:

- Contacting the submitting individual(s) for missing information or for clarifications
- Preparing and forwarding the completed Study Concept Form to the BMT CTN Executive Committee for review on the monthly Executive Committee conference call.

The Executive Committee reviews a Study Concept Report to determine whether the proposed study is consistent with the overall mission of the BMT CTN, has no major conflicts with active BMT CTN protocols or BMT CTN protocols in development, and poses no major conflicts of interest. It is expected that most proposals will be approved by the Executive Committee and forwarded to the Steering Committee for determination of scientific merit, feasibility and willingness to participate (see section 2.4).
2.4. Study Concept Form Review and Prioritization

Once approved by the Executive Committee, the proposed study will be discussed at the next Steering Committee meeting.

The investigator (or his/her designee) submitting the proposal presents the proposal to the Steering Committee; multiple presentations may be required. The Steering Committee reviews the proposal for scientific merit and alignment with scientific direction of the BMT CTN, feasibility and willingness to participate.

The relative importance of the following three measures is evaluated by the Steering Committee. It is expected that scientific merit will be the primary consideration after which the remaining two areas will be considered equally.

1. Scientific Merit: The scientific merit of the proposed study will be rated using the NIH scale for grant/contract applications. Assessment of scientific merit will include consideration of the BMT CTN scientific direction.

2. Feasibility: The feasibility of the proposed study will be based on the potential for accrual and ability of centers to conduct the proposed treatment interventions (e.g., studies requiring specialized graft manipulations available at only one center may be viewed as less feasible than a trial utilizing only standard graft manipulations). Budget considerations are also important in assessing feasibility.

3. Willingness to Participate: It is anticipated that Core Centers will enroll the majority of study participants on BMT CTN trials. The willingness of Core Center participation is a key factor in assessing any proposal.

2.5. Review Cycle

Proposals for BMT CTN clinical trials may be submitted at any time. The review will be conducted in a timely fashion after receipt of the Study Concept Form.

The Steering Committee reviews Study Concept Forms during regularly scheduled Steering Committee in-person meetings or teleconferences. This does not guarantee a study will be conducted. A study proposal must be approved by a majority of Steering Committee members and achieve a sufficiently high priority score to be scheduled for development.
CHAPTER 3

PROCEDURES FOR IMPLEMENTING APPROVED STUDY CONCEPTS
3. PROCEDURES FOR IMPLEMENTING APPROVED STUDY CONCEPTS

3.1. Tasks Following Approval of a Study Concept

Approval of a Study Concept by the Steering Committee requires that a number of activities be initiated. While outlined below as separate activities, in practice, they are carried out concurrently to the extent possible. Tasks include:

- Assignment of a study number
- Establishment of a Protocol Team
- Development of the protocol document and draft informed consent form(s)
- Consideration of regulatory requirements
- Consideration of contributor and/or collaborator requirements
- Identification of sites
- Identification and procurement of services that will be required from outside (non-Network) providers
- Preparation of a study budget
- Development of an accrual plan
- Preparation of study-related educational materials, if applicable
- Development of Case Report Forms and implementation of data system
- Development of protocol-specific site training
- Development of patient-specific materials
- Development of a Research Sample Information Guide
- Development of a Pharmacy Guide, if appropriate

3.2. Establishment of a Protocol Team

Each approved Study Concept has a Protocol Team responsible for protocol development, oversight of the trial, and analysis and publication of the study results. The Protocol Team includes the following members:

- Protocol Chair(s)
- Core and Affiliate Center Representatives
- Protocol Officer
- Protocol Statistician
- Protocol Coordinator
- NHLBI and/or NCI Representative
- NHLBI Statistician
- Other members (e.g., DCC Business Representative; NCI funded NCTN Group representatives; and ad hoc members as deemed necessary by the Protocol Chair)

Protocol Team Member Application Process for Core and Affiliate Center Representatives:
- Investigators must be nominated or endorsed by a Core center PI, DCC leadership, or by BMT CTN disease/discipline committee leadership
- Potential study members must formally apply to participate by submitting an NIH biosketch with a statement indicating pertinent expertise and commitment to study team; for junior investigators, identification of a senior mentor is encouraged.

Protocol Team Member Selection Process:
- Study team member applications will be reviewed by the Network leadership, to include, the Network Chair, Vice-Chair, Chair-Elect or Immediate Past Chair, DCC PI and co-PIs, and, if applicable, Chair of the relevant disease/discipline-specific committee.
- Final team members will be selected based on the following criteria:
  - Prior involvement in developing the study concept
  - Relevant expertise
  - Junior investigator with well-established senior mentor
  - Diversity
    - Centers previously under-represented
    - Likelihood to make time commitment
    - Academic rank (mix of junior and senior members)
    - Unique skill sets
    - Affiliate Center representation

Protocol Chair(s): The Steering Committee Chairperson appoints one to four Protocol Chairs, in collaboration with the Past-Chair, Chair-Elect, ViceChair and the DCC Principal Investigators. One of the Protocol Chairs is typically the person who submitted the Study Concept. Protocol Chairs must have experience with the patient population and therapy being studied and commit their center to enrolling a significant percentage of their eligible patients on the study. The Protocol Chair(s) has (have) primary responsibility for:
- The study throughout its course, including but not limited to, chairing Protocol Team meetings, reviewing protocol drafts and amendments prepared after discussions of the Protocol Team
- Presenting progress reports on protocol development, implementation and progress to the Steering Committee
- Leading Investigators Meetings for the protocol
- Leading the scientific portion of Site Initiation webcasts for the protocol
- Addressing issues raised by the Protocol Officer and Protocol Coordinator; investigators at participating centers; and the PRC, DSMB and NMDP IRB in consultation with Steering Committee leadership
• Reviewing and approving all versions of the protocol, informed consent(s), and summary of changes documents before and after submission to the PRC, DSMB, and NMDP IRB (and FDA, if appropriate), and prior to release of the official version to sites
• Presenting protocol to the PRC and DSMB
• Preparing Frequently Asked Questions Document for the PRC
• Responding to queries from the Protocol Coordinator on behalf of site personnel in a timely manner
• Attending a minimum of 70% of the Protocol Team calls throughout the development, implementation, maintenance and endpoint review stages. A Protocol Chair cannot appoint another person as a substitute for Protocol Team calls. The Chair is responsible for reviewing the team minutes for a missed call upon receipt and providing feedback via email.
• Ensuring their center is among the top 4 ranked accruing centers, or has accrual ≥ 80% of their DCC-approved accrual projection. Since accrual is integral to successful and timely completion of BMT CTN studies, accrual performance may impact Protocol Chair status, primary manuscript authorship order and/or center performance score.

Core and Affiliate Center Representatives: Approximately 6-10 Core and Affiliate Clinical Center representatives will serve as co-investigators on the Protocol Team to help develop the protocol document, supervise the study throughout its course and interpret and present study results. Specific responsibilities include:
• Addressing issues raised by the Protocol Officer, Protocol Chairs, fellow Protocol Team members and Protocol Coordinator, investigators at participating centers, the PRC, DSMB and NMDP IRB (and FDA, if appropriate) in consultation with Steering Committee leadership
• Participating in the review of all changes to the protocol before and after submission to the PRC, DSMB and NMDP IRB (and FDA, if appropriate) and prior to release of the official version to sites
• Attending a minimum of 70% of the Protocol Team calls throughout the development, implementation and maintenance stages of the protocol. A Protocol Team member cannot appoint another person as a substitute for Protocol Team calls. The member is responsible for reviewing the team minutes for a missed call upon receipt and providing feedback via email. Ensuring their center accrues ≥ 80% of their DCC-approved accrual projection. Since accrual is integral to successful and timely completion of BMT CTN studies, accrual performance may impact protocol team member status and/or center performance score.

Protocol Officer: Responsibilities of the Protocol Officer include:
• Working closely with the Protocol Coordinator and Protocol Chair(s) in developing the protocol and consent documents, responding to center queries, preparing materials/agenda for meetings and teleconferences, reviewing CRFs, etc.
• Providing an additional level of scrutiny of the proposed protocol specifications, specifically addressing whether they are sufficient to accomplish the scientific objectives of the study
• Monitoring delays in protocol development/review/accredual and addressing obstacles to progress promptly in consultation with the Protocol Chair(s)
• Communicating with the Protocol Chair(s) and Protocol Coordinator regarding the status and progress of the trial
• Presenting protocol progress to the Executive Committee
• Assisting in preparation of materials for the PRC, DSMB and NMDP IRB (and FDA, if appropriate)
• Review safety sections of protocols and help develop appropriate safety stopping rules as needed
• Assists with protocol amendments

Protocol Statistician: The Protocol Statistician has primary responsibility for the study’s statistical design and analysis. For some protocols, a secondary Protocol Statistician may also be designated. Both statisticians are from the DCC and have PhDs. The Protocol Statistician:
• Interacts and communicates with the Secondary Protocol Statistician, if applicable, on a regular basis regarding issues of statistical design and for secondary review of the final study analysis plan and analysis
• Assists in preparing materials for the PRC, DSMB, NMDP IRB and FDA (if appropriate)
• Writes the Statistical Considerations protocol chapter and any supporting appendices if applicable
• Assists with development and review of case report forms to ensure appropriate data is collected in an analyzable matter.
• Assists with protocol amendments if related to statistical considerations or endpoints/endpoint definitions
• Interacts and coordinates with the Operational Statistician for the preparation of the Endpoint Review Charter (see section 3.4.5).

Protocol Coordinator: The Protocol Coordinator is generally a member of the Emmes Data Management/Protocol Monitor staff. The Protocol Coordinator has responsibility for:
• Overseeing all aspects of development of the Protocol Document from initial concept though final draft including internal and external reviews, revisions, approvals, and final dissemination of initial protocol and subsequent amendments
• Preparing revised drafts of the protocol pursuant to discussions of the Protocol Team and ensuring completeness and internal consistency of the protocol and conformity with standards of protocol production and BMT CTN DCC policies and procedures
• Preparing agendas, minutes, regulatory documents, as needed, and protocol documents for meetings of the Protocol Team, Steering Committee, PRC, DSMB, NMDP IRB (and FDA, if applicable)

NHLBI and/or NCI Representative(s): Each Protocol Team will include two NIH representatives, one from NHLBI and one from NCI. In addition, an NHLBI statistician may participate.

Other members (e.g., DCC Business Representative, NCI funded NCTN Group representative if the study is being done in collaboration) are appointed to the Protocol Team, as necessary, for specific expertise related to the study.

3.3. Development of a Protocol

The Protocol Team will meet by teleconference as soon as possible after formation of the team and at frequent intervals thereafter. It is recommended that Protocol Team meet in person for a one-day meeting to expedite the protocol development process. The day/time of the teleconferences and recommended in person meeting are scheduled by the NMDP Project Coordinator. Each Protocol Team member is assigned access to the BMT CTN SharePoint website for electronic communications regarding the protocol.

3.3.1. First Protocol Draft/Outline

• The Protocol Team is responsible for generating the first draft of the protocol based on the Study Concept Report and comments from the Steering Committee. Many of the critical elements of the protocol will be contained in the Study Concept Report. The Protocol Document will follow the BMT CTN protocol template (which is posted on the BMT CTN SharePoint website): protocol synopsis, background and rationale, eligibility criteria, study design, study treatment, study endpoints, patient enrollment and evaluations, and statistical considerations; and Appendices including human subject, laboratory procedures and references. The informed consent and assent forms are drafted by a member of the NMDP Patient and Health Professionals Services team after discussion with the protocol team and/or Protocol Coordinator. The Protocol Document is the primary study document used to guide conduct of the trial.

In advance of the first Protocol Team teleconference, the Protocol Chair(s), if needed, with the assistance of the Protocol Officer and Coordinator, prepares a first protocol draft/outline for distribution to the members of the Protocol Team. Prior to distribution, the following occur:

• The Protocol Officer reviews the draft/outline for medical/safety issues, agreement with the Study Concept Report, and potential protocol design and feasibility issues.

• The Protocol Statistician identifies statistical issues that require Protocol Team input.

3.3.2. Working Draft Protocol

After the first Protocol Team conference call, the Protocol Coordinator with assistance from the Protocol Chair(s) and/or Protocol Officer incorporates suggested revisions. This Working Draft Protocol is then circulated to the Protocol Team for further discussion and comment. Multiple
iterations of this process are generally necessary to develop a Protocol Document that is ready for presentation to the Steering Committee.

The Protocol Document includes:

- Synopsis (including protocol schema) that concisely outlines the study design and objectives
- Background, scientific rationale, and objectives for the study
- Detailed eligibility (inclusion/exclusion) requirements for participation
- Detailed description of the treatment plans and supportive care measures
- Detailed description of expected risks associated with the treatment plan
- Accurate clinical definition of all primary and secondary endpoints
- Details of registration and randomization procedures, if applicable
- Required clinical exams and specimen submission schedules
- Follow-up schedules and requirements for patient monitoring
- Unexpected Grades 3-5 adverse events reporting procedures
- Additional adverse event reporting requirements for study drug or device, if provided by industry partner
- Detailed Statistical Considerations section including a description of the experimental design and analysis plan, sample size estimates, randomization, analysis plan of primary and secondary outcome measures, plans for interim analyses (if required) and stopping guidelines. Sample size considerations will include power, baseline rate and precision.
- Description of the risk-based endpoint review process
- References laboratory procedures and human subjects considerations
- Other appendices and miscellaneous procedures (e.g., those for blinding study treatments)

The informed consent documents (including the assent if applicable) are prepared during protocol development but are maintained as separate document(s). Essential elements of the Informed Consent include:

- Statement that the study involves research
- Purpose of the study/research
- Sponsor of study
- Expected duration of the subject’s participation
- Number of subjects
- Description of procedures
- Description of foreseeable risks and discomforts including death if appropriate
- Risks to the unborn; prescribed birth control for males and females
• Benefits to subject
• Benefits to others
• Alternative procedures or treatments
• Degree to which confidentiality of records will be maintained
• Clinicaltrials.gov language
• CIBMTR data collection language
• Record retention
• Inspection of records
• Certificate of Confidentiality Language
• Contact Name/Number for questions about research/study, subject rights and research-related injury
• Medical treatment for study/research-related injury
• Liability for study/research-related injury including that the sponsor (NHLBI/NCI will not pay)
• Participation is voluntary
• Refusal to participate will not be held against you
• Discontinue and/or terminate participation – by subject or others
• Compensation
• Costs and Reimbursements
• New findings
• HIPAA language
• Research samples (required and optional)

Specific language is required for some of the elements as described below:

• Clinicaltrials.gov language must be inserted verbatim: A description of this clinical trial will be available on [http://www.ClinicalTrials.gov](http://www.ClinicalTrials.gov) as required by U.S. Law. This Web site will not include information that will identify you. At most, the Web site will include a summary of results. You can search this web site at any time.

• Certificate of Confidentiality language must be inserted verbatim in the section titled “Who will see my medical information”: Your privacy is very important to us. The study doctors will make every effort to protect it. The study doctors have a privacy permit to help protect your records if there is a court case. However, some of your medical information may be given out if required by law. If this should happen, the study doctors will do their best to make sure that any information that goes out to others will not identify you.

• CIBMTR Data collection language: Data regarding your clinical situation, including follow up [after insert number of years of protocol follow up] may be obtained by the BMT BMT CTN from the CIBMTR, which captures information on all US transplants.
Other considerations generally include handling of laboratory samples and repository use issues, budget preparation/management and development of accrual plans (see Sections 3.5, 3.6).

Early in the process, the Laboratory/Repository Manager, in collaboration with the Biomarkers Committee, works with the Protocol Team to identify biological samples that may be required to address the basic study question. Consultation with the Protocol Team will include discussions relevant to assessment of tests considered standard of practice versus those designated specifically for research.

Steering Committee feedback is requested to identify barriers to site participation and to recommend possible changes that may facilitate participation. The Steering Committee will review required clinical and laboratory assessments and compare to accepted standards of practice. Recommendations from the Steering Committee are discussed by the Protocol Team and changes are then incorporated into the Protocol as necessary.

3.3.3 Circulating Draft Protocol
The Protocol Coordinator, with the assistance of the Administrative Coordinator, prepares the Circulating Draft Protocol which includes a version number and date. The Protocol Team must approve the Circulating Draft Protocol prior to distribution to the Steering Committee for review and approval.

3.3.4 Steering Committee Approval
Two weeks in advance of Steering Committee review, two to three Steering Committee members are designated by the Steering Committee Chairperson to review the circulating draft protocol document prior to presentation to the Committee. These reviewers assess details within the document, complete a Steering Committee Reviewer Checklist (posted on the BMT CTN SharePoint website) and provide comments regarding significance, innovation, approach, investigator(s), environment and overall assessment of the study.

Following the presentation to the Steering Committee and subsequent discussion, committee members will accept, recommend changes or reject the protocol. A Circulating Draft Protocol may require multiple presentations to the Steering Committee prior to final Steering Committee approval at which time “draft” is removed from the document.

Once the Steering Committee has approved the protocol, it is reviewed/edited by the Protocol Team and then preparation begins for PRC review and approval (see section 3.4.4.).

3.4 Activities Related to Protocol Development/Implementation
Once the Working Draft Protocol is available, other protocol specific matters may require coordination. These are as follows:

- External services
- Regulatory requirements
- Site identification
• Accrual initiatives
• Repository/laboratory compliance
• Protocol specific site training
• Participant materials

3.4.1 External Services
The Protocol Chair, Protocol Officer, Protocol Coordinator and Business Representative review a list of all study participant care measures and investigations that are included in the protocol to determine which, if any, aspects of the protocol require contracting with external service providers. Examples include centralized pharmacy for acquisition and distribution of study medications, research or other central laboratories, radiation quality control services, etc.

3.4.2 Regulatory Requirements
The Protocol Chair, Protocol Coordinator and the Protocol Officer are responsible for addressing regulatory requirements. These include the FDA requirements (if appropriate):
• Communicating with the FDA and other regulatory agencies identifying the need for IND or IDE
• Coordinating the submission process with NHLBI and/or NCI/NIH/industry
• Completing an IND/IDE application if required
• Waiting 30 days after submission prior to releasing protocol to participating sites (to be assured that there is not a clinical hold)

3.4.3 Site Identification
The following steps, though usually performed in a logical sequence, represent processes that may occur throughout all protocol implementation phases, from development through accrual completion.

Once the Working Draft Protocol is available, the typical sequence is as follows:
• A protocol synopsis is distributed to Core Centers surveying their interest in participation. If interested, the centers are asked to provide:
  – Contact information for PI and Key Contact
  – The numbers of patients they project to enroll annually
  – Number of competing protocols at their center, if applicable
• Affiliate Centers are identified (and contacted, if necessary) if they have previously enrolled large numbers of patients on other protocols or have a significant patient population that would be eligible for the study as identified from CIBMTR data). The majority of BMT CTN protocols require Affiliate Center participation.
• If even more sites are required, the Protocol Coordinator arranges for a notice to be posted on the BMT CTN public website along with the Protocol Synopsis announcing the potential study and requesting interested centers that meet the requirements to complete the
Application for Participation as a Affiliate Clinical Center in the BMT CTN and to contact the DCC.

- Short form application acceptable if Center has previously participated in a BMT CTN trial
- Long form application required if there is no previous participation

- If additional sites are still required, the Protocol Officer and Protocol Coordinator identify potential Affiliate Centers for direct contact by the Protocol Chair(s)
- If applicable, Central Operations Offices of NCI NCTN Groups are asked for a listing of their transplant centers
- Occasionally international centers or networks may request to participate in BMT CTN studies. Regulatory, activation and study logistics requirements for international centers may vary by study and country. In addition to the Protocol Team, international centers must be approved by NHLBI and receive U.S. State Department clearance.

During a Protocol Team conference call, the Protocol Team considers Affiliate Center participation based on protocol-specific criteria such as:

- Study sample size, study participant population and type of transplant
- Need for special center training for study-specific procedures (e.g., graft manipulation)
- Prior experience with BMT CTN studies including accrual history and retention rates

The DCC Business Representative receives all submitted applications and provides each Affiliate Center written notification indicating their application has been accepted, not selected or placed on “hold” for future considerations.

If not selected as a Study Site, an Affiliate Center may appeal the decision to the Executive Committee. The decision of the Executive Committee regarding Center participation is final.

3.4.4 Protocol Review Committee (PRC) Review

The PRC is a standing committee appointed by the NHLBI (see Chapter 1). Prior to scheduled meetings, the NHLBI Executive Secretary provides the DCC with a roster of PRC members and NIH participants. The Protocol Chair(s), Protocol Officer and Statistician and senior members of the DCC represent the Protocol Team at the PRC meetings.

Protocol reviews are typically held by conference calls. Dates and times are coordinated by the NHLBI Executive Secretary and the DCC. The PRC Chair and Executive Secretary prepare an agenda in consultation with the DCC. In general, the agenda includes:

- Short presentation of the study by Protocol Team representatives
- Open discussion between PRC members and Protocol Team representatives
- Closed executive session (limited to PRC members, the Executive Secretary and other NIH staff)
• Open session listing the PRC recommendations (subject to approval by the NHLBI Office of the Director)

Three weeks in advance of the meeting the DCC prepares the following materials for distribution to the PRC by the Executive Secretary:

• Steering Committee approved protocol document
• “Frequently Asked Questions (FAQs)” document
• Accrual plan
• The PRC Reviewer Checklist to be completed by each reviewer and returned to the Executive Secretary prior to the teleconference regarding the study design and feasibility:
  o Importance of the question being addressed
  o Need for multi-center network to meet objectives
  o Merit of experimental design, including appropriate controls, treatment plan, study endpoints, and sample size
  o Study feasibility including appropriate per patient budget for network study and adequate resources including medications
  o Appropriate recruitment strategies
  o Adequacy of patient population and number of patients, including appropriate representation of minorities, women, and children

The PRC Executive Secretary is responsible for compiling the completed PRC Reviewer Checklists and distributing their comments to the DCC. The Protocol Coordinator will work with the Protocol Team representatives to prepare a written response for distribution by the Executive Secretary to the PRC members at least 24 hours prior to the scheduled meeting. The responses and any additional questions will be discussed during the Open Session of the PRC meeting. A closed session with the PRC members will be held to discuss the responses and determine if there are any remaining concerns. At the end of the teleconference, the PRC Chair will provide a verbal list of recommendations which are considered “unofficial” until approved by the NHLBI Office of the Director.

The NHLBI Executive Secretary prepares the final meeting minutes. After approval by the PRC Chair, the minutes of the PRC meeting are sent to the NHLBI Office of the Director for final approval. The Executive Secretary provides a copy of the Director-approved document to the DCC for distribution to the Protocol Team.

If subsequent PRC reviews are required, revised materials are prepared and include:

• Revised or completed protocol document
• Protocol document noting substantive changes since prior review
• Protocol Team written response to the PRC recommendations/comments
After PRC approval, the Protocol Document with changes recommended by the PRC is prepared for submission to the DSMB for review.

In addition, the following Technical Committees review the protocol at this time.

- Biomarkers
- Clinical Research Associate
- Pharmacy, if applicable
- Special Populations (Pediatrics/Human Subjects)
- Toxicity and Supportive Care

### 3.4.5 DSMB Review

The DSMB is a standing committee appointed by the NHLBI (see Section 1.6.3). New protocol reviews are generally presented during regularly scheduled teleconferences in the spring and fall. These are coordinated by the NHLBI Executive Secretary with assistance of the DCC. Ad hoc meetings may be arranged in the event the scheduled teleconferences and meetings are not scheduled in the near future. The DSMB Chair and Executive Secretary prepare an agenda in consultation with the DCC. In general, the agenda includes:

- Short presentation of the study by Protocol Team representatives
- Open discussion between DSMB members and Protocol Team representatives
- Closed executive session (limited to DSMB members, the Executive Secretary and other NIH staff)

The following materials are provided to the DSMB three weeks prior to scheduled meetings:

- Final PRC approved protocol document
- FAQs document (updated to address questions posed by the PRC if appropriate)
- Accrual plan
- PRC recommendations/comments and Protocol Team responses
- DSMB reviewer checklist to collect comments, issues of concern, requests for additional information or clarification in the following areas:
  - Study objectives and feasibility
  - Study design and statistical approach
  - Study operations
  - Human subjects protection
  - Overall approval recommendation

If applicable, the protocol will also be submitted for FDA review at the time of the initial submission to the DSMB.
The NHLBI Executive Secretary is responsible for compiling the completed DSMB reviewers’ checklists and distributing their comments to the DCC. The Protocol Coordinator will work with the Protocol Team representatives to prepare a written response for distribution by the Executive Secretary to the DSMB members at least 24 hours prior to the scheduled meeting/teleconference.

The Executive Secretary prepares the meeting minutes. After approval by the DSMB Chair, the minutes of the DSMB meeting are sent to the NHLBI Office of the Director for final approval. The Executive Secretary provides a copy of the Director approved document to the DCC for distribution to the Protocol Team.

If subsequent DSMB reviews are required, revised materials are prepared and include:

- Revised or completed protocol document
- Protocol document noting substantive changes since prior review
- Protocol Team written response to the DSMB recommendations

Director-approved documentation of the DSMB minutes are required for each subsequent DSMB review and are distributed by the Executive Secretary to the DCC. The DSMB minutes along with FDA approval of the protocol, if appropriate, are required prior to submission to the NMDP IRB.

3.4.6 NMDP IRB Review

The NMDP IRB serves as the single IRB for all BMT CTN studies released after July 1, 2017. The members of the NMDP IRB are a diverse group of distinguished healthcare professionals, donor advocates, and patient advocates with expertise in bone marrow transplantation and hematology/oncology.

The DCC provides the following materials to the NMDP IRB three weeks prior to scheduled monthly meetings:

- NMDP IRB Initial Application Form
- Final DSMB (and FDA, if applicable) approved protocol document
- Informed consent and assent (if applicable) documents
- Subject recruitment or educational study materials and materials/instruments/forms to be completed by subjects, if applicable
- PRC and DSMB approval documentation
- Principal Investigator CV and documentation of protection of human research training
- Other materials, if applicable (e.g., Investigators Brochure)

The NMDP IRB informs the DCC if the study is approved, approved with stipulations, or not approved. If stipulations or resubmission is required, the DCC will prepare the response with input from the Protocol Team.

As a final step once the protocol is approved by the NMDP IRB, the Protocol Chair(s), Protocol Officer and Protocol Coordinator must review the protocol document for completeness and
accuracy and complete a Protocol Team Reviewer Checklist. Two Protocol Chairs or one Chair and the Officer must sign the front page of the document indicating their approval of the document. Once complete, participating centers will be notified of the new NMDP IRB-approved protocol and informed consent documents by a Numbered Memorandum distributed via e-mail from the DCC. The protocol will be posted on the both the BMT CTN public website and the BMT CTN SharePoint website. The Informed Consent is posted as a Word document on the SharePoint website for institute-specific formatting. No changes to the protocol document by a participating center are allowed with the exception of reformatting the Informed Consent and Assent documents to incorporate institution-specific language. The center sends the modified consent and assent (if applicable) documents to the NMDP IRB. If the consent/assent documents are translated into another language, the translated version must be provided to the NMDP IRB along with translator certification documentation. The NMDP IRB administrator will review the consent document(s), and if approved, sent final document(s) back to the Center with NMDP IRB approval recorded on the documents. If not approved, the center will be notified they need to revise and resubmit the documents.

3.4.7 Medical Monitor Assignment

The DCC assigns a Medical Monitor to each study when it is activated. Medical Monitors are transplant physicians familiar with the conduct of clinical research and regulatory requirements for safety monitoring and reporting. The Network’s Medical Monitors hold teleconferences quarterly to ensure uniformity in assessing adverse events and to share updates on regulatory requirements. As a matter of policy and to avoid bias, these physicians may not enroll patients or care for patients enrolled in a study on which they serve as Medical Monitor.

The Medical Monitor has primary responsibility for reviewing safety issues, including protocol-specified safety provisions (e.g., assessment of stopping rules) and unexpected events. A primary and alternate Medical Monitor is assigned for each study. Should a study require a disease-specific Medical Monitor (e.g., HIV or sickle cell disease), a primary transplant and disease-specific Medical Monitor will be assigned along with an alternate transplant Medical Monitor. The Medical Monitor is a physician:

- Trained in transplant or the disease-specific area at an institution not participating in the study.
- Trained in the protocol safety reporting requirements by the DCC
- Reviews reportable adverse events real-time and other safety concerns identified by the DCC
- Reviews of protocol-specified provisions periodically

3.5 Development of Protocol Endpoint Review Process and Charter

The mission of the Protocol Endpoint Review is to conduct a review of the endpoints and assess subject eligibility and protocol compliance. When possible, the review should be done in a blinded manner. The Protocol Coordinator and Operational Statistician will prepare the review materials for the committee.

The Endpoint Review Committee (ERC) is nominated by the Protocol Chair(s). Typically, the Committee consists of:
- Protocol Chair(s)
- Two or more Investigators that contributed to the development and implementation of the protocol (measured by Protocol Team call attendance and/or center accrual) or Junior Investigators from high-enrolling centers
- Protocol Officer
- Primary Protocol Statistician
- Operational Statistician
- Protocol Coordinator
- Ad hoc members as deemed necessary by the Protocol Chair(s)

The members of the ERC are expected to attend a minimum of 70% of the ERC teleconferences and to provide their written review of cases prior to each call, including calls that they are unable to attend. Unsatisfactory participation may result in removal from the ERC.

The Protocol Team should include a description of the risk-based endpoint review process in the protocol and develop the Protocol Endpoint Review Charter before the study is open to accrual. The charter should outline the primary and secondary endpoints to be evaluated and detail the endpoints to be adjudicated, process for adjudication, procedures for discrepancy resolution, and recording of results. The charter must be aligned with the study protocol and the statistical plan and developed in conjunction with the Case Report Forms and Study Database (see Section 3.6). The charter should be reviewed by the Protocol Team on an annual basis.

### 3.6. Development of Case Report Forms and Study Database

BMT CTN Case Report Forms (CRFs) development begins when the Working Draft Protocol is available. The Protocol Coordinator is responsible for the following items with regards to CRF and database development:

- Drafting the initial version of the BMT CTN CRFs based on the working draft of the protocol and the Endpoint Review Charter and identifying core forms to be used and required protocol-specific forms, if necessary
- Reviewing the draft forms with statistical staff or senior staff members to assess accuracy and comprehensiveness of collecting data for analysis of primary and secondary endpoints
- Reviewing the draft forms with the Protocol Team to assess accuracy and comprehensiveness of collecting data for analysis of primary and secondary endpoints
- Determining specific data to be provided by the CIBMTR and verifying that the CIBMTR data will meet the requirements of the statistical plan and Endpoint Review Charter
- Reviewing CRFs with the CRA Committee
- Drafting secondary versions based on Protocol Team and CRA Committee recommendations
- Piloting CRFs when necessary
- Obtaining statistical staff or senior staff member’s approval of CRFs
• Finalizing CRFs and updating the data system

NOTE: Data collected by CIBMTR supplement BMT CTN data. The CIBMTR collects data at two levels: 1) Transplant Essential Data level and; 2) Comprehensive Report Form level. The majority of BMT CTN studies require CIBMTR Comprehensive Report Form level data.

The methodology of the software industry, specifically the Software Development Life Cycle (SDLC), is employed when updating the data system for release of new or modified CRFs:

• Requirements are defined prior to the initiation of work on the data entry system and an implementation schedule/timeline is agreed upon by all involved staff
• The functionality is implemented by the Protocol Coordinator, an Systems Analyst and/or Programmer
• Validation and verification is performed on all work and all identified issues/defects are addressed before release to the transplant centers
• All work is documented to ensure compliance with industry standards and to provide a history of changes made to the data system

The use of Common Data Elements (CDEs) promotes the understanding and sharing of cancer research information. The comprehensive set of standardized metadata descriptors for cancer research data defined in the Cancer Data Standards Repository (caDSR) was utilized to review and analyze CRFs employed by the BMT CTN. The BMT CTN continues incorporation of CDEs to facilitate interoperability with other research trial efforts including data exchange with the NCI NCTN Groups.

3.6.1. Accrual Initiatives

A critical challenge of the Network is timely accrual to all active protocols. Accrual is the highest priority of the Network. The Accrual Coordinator:

• Participates in the development and implementation of new trials
• Prepares an accrual plan for each new protocol prior to launch
• Sets and tracks activation target dates
• Monitors quarterly accrual reports
• Assists in identification of potential Affiliate Centers
• Coordinates and oversees patient accrual strategies
• Participates in resolving accrual problems
• Helps develop and participates in educational sessions for clinical center coordinators
• Assists in coordination of educational webinars
Other related accrual activities may include collaborating with NCI-funded NCTN Groups and their Operations Offices, working with external Networks having interest in questions specific to BMT CTN study or access to targeted patient populations.

Throughout the process, a representative from the NMDP/Be The Match Patient and Health Professional Services is consulted if there are unique needs for study-related patient educational materials. This representative collaborates with the Accrual Coordinator and the Protocol Team in developing materials as needed:

- Educational materials
- A description of the study to be made available in brochure form and posted on the Network’s public website (and other appropriate websites)
- Advertisements for study participant recruitment
- Web postings for study participant recruitment
- A series of “Frequently Asked Questions” (FAQs) for patients (vs. the FAQs developed for the PRC)
- Responses for the Patient And Health Professional Services staff to use when addressing queries regarding the study as it is publicized

All subject recruitment and educational materials are submitted to the NMDP IRB for approval with the final approved Protocol Document and Consent forms.

3.6.2 Repository/Laboratory Compliance Considerations

The BMT CTN has invested substantial efforts into defining the critical scientific objectives related to the collection, biorepository storage and utilization of patient and donor biologic research specimens. A dedicated Laboratory/Repository Manager is a DCC staff member who coordinates these critical activities, some with in conjunction with the Biomarkers Committee:

- Participates on the Biomarkers Committee as ex-officio member
- Assists in establishing principles for specimen collection (including their use in Ancillary Studies)
- Reviews protocol drafts (see Section 3.2.2) to identify specific study-related tests and required biospecimens, including those with potential as prognostic markers
- Prepares quarterly Laboratory Compliance Reports and makes recommendations to the Executive Committee for annual Center Performance Evaluations
- Assists participating centers in resolving specimen discrepancy problems
- Serves as a resource in the preparation of the protocol specific Laboratory Sample Information Guide

3.6.3 Protocol Specific Site Training

During the development of the Working Draft Protocol, the Protocol Chair(s), Protocol Coordinator and Protocol Officer will identify site training requirements for a specific study. This
may be necessary for patient evaluations (e.g., grading GVHD), use of a new device (e.g., graft manipulation), specific stem cell lab procedures, or web-based CRF completion.

For areas requiring special training, the Protocol Coordinator will identify appropriate individuals to conduct the training. If related to a particular device, the manufacturer of the device will be considered as a potential source of training personnel. For web-based data collection issues, the Protocol Coordinator will coordinate the necessary training.

When such training requires contracting with outside organizations, the name, contact information and required activity will be forwarded to the DCC Business Representative (or designee) who will initiate subcontract discussions with the potential provider. In addition, the Business Representative will use this information when determining the study budget.

See additional information regarding training in Chapter 4.

3.7. Protocol Budget and Management of Contributions

3.7.1. Protocol Budget Preparation and Revisions

The DCC Finance and Contracts Department at NMDP/Be The Match (collectively referred to as the “Business Representative”) are responsible for protocol budget development and coordination with the NIH regarding the management of the overall clinical care funds.

3.7.1.1. Protocol specific budgets

Budget Preparation: When a Working Draft Protocol is available, the Business Representative, Protocol Chair(s), Protocol Officer and Protocol Coordinator, as well as any other key parties if necessary, meet via teleconference to initiate budget preparation. The following materials (found on the BMT CTN SharePoint website in the “Budget and Contracts” folder) are reviewed during this meeting:

- Protocol Budget
- Study Drug Budget
- Laboratory Budget
- Product or Services

The Business Representative drafts a budget based on this information. Standard of care tests, medications, pharmacy costs, etc. are included in the budget.

The protocol draft budget includes the following assumptions:

- Projected enrollment
- Number and type of Clinical Centers (e.g., Core/Consortia Centers, Affiliate Centers, NCI-funded NCTN Groups, etc.)
- DCC labor projections (anticipated contributions, e.g., central laboratory, central pharmacy, packaging and shipping)
• Per-patient budgets (Clinical Center payment)
  - Labor hours for the physician, co-investigator, clinical research coordinator and/or data manager
  - Protocol specified non-standard of care assessments or procedures
  - Shipping costs and supplies used at the Clinical Centers
  - Specialized evaluations (such as MRIs, MRAs, neurocognitive tests, etc.)
  - Other considerations (e.g., studies done with NCI-funded NCTN Groups)

3.7.1.2. Budget review and approval process

The draft budget will be reviewed by the Protocol Chair(s) and the Protocol Officer. The Steering Committee must approve the budget prior to the protocol PRC submission. The final budget must be reviewed by the Business Representative and Protocol Officer prior to submission to the NHLBI and NCI for approval. When approved, a summary of the per patient fee will be posted on the BMT CTN SharePoint Website.

The Business Representative completes the following budget-related actions after Protocol Officer approval:

• Proceeds with the request for proposals and contracts for centralized services
• Finalizes contributions
• Reviews the budget for any final revisions
• Executes contracts and Clinical Study Protocol Riders with the participating Clinical Centers and NCTN Groups

3.7.1.3. Budget revisions

If a protocol specific budget is adjusted due to contributions or a significant change in expenditures, a revised budget will be circulated and approved as described above. After approval, the revised per patient fee summary, if applicable, will be posted on the BMT CTN SharePoint website and the revised budget will be circulated to all appropriate parties by the Business Representative.

If protocol amendments occur, the Protocol Chair(s), Protocol Officer, Protocol Coordinator and Business Representative are responsible for reviewing the amendment to determine if modifications affect the budget. If budget adjustments are required, the procedures described above will be followed.

When all milestone payments of a protocol have been made or can be projected, the overall BMT CTN budget is adjusted to reflect the true cost of the study. Additionally, the DCC provides NIH (NHLBI and NCI) the budgeted and actual protocol costs.

3.7.2. Per Patient Fee

3.7.2.1. Basis of payment

The payment schedule shall be determined by the Protocol Chair(s) and the DCC, and approved by NHLBI and NCI. Examples include:
• Entire payment upon enrollment
• Milestone basis:
  o 50% upon patient enrollment
  o 25% at study day 180 based on submission and approval of data
  o 25% upon receipt of complete and acceptable data for a study participant
• Other payment schedules as appropriate

The per-patient budgets are estimates of labor time effort, materials, etc. and used for budget development purposes. The centers may apply the per-patient funds however they deem appropriate to perform the protocol-specific tasks as BMT CTN does not govern the use of per-patient funds the clinical centers receive.

3.7.2.2. Payment guidelines for collaborative studies with NCI NCTN Groups

Consensus guidelines developed by NCI and NHLBI are followed for studies done in collaboration between BMT CTN and NCI NCTN Groups. Payment is consistent with these guidelines, NCI NCTN Group policies and BMT CTN policies.

• For BMT CTN led trials done in collaboration with the NCI NCTN Groups, the relevant Group (or as applicable the Cancer Trials Support Unit (CTSU)) contributes their standard per-accrual fee for patients enrolled from their sites
• For NCI NCTN Group led trials, the BMT CTN will pay its Network centers the NCI standard per-accrual Lead Academic Participating Sites fee for patients enrolled from a BMT CTN Center. Payment typically occurs at the time of patient enrollment.

3.7.3. Contributions

The Protocol Team may identify potential contributors. Once identified, the Protocol Chair or designee shall work with the DCC Business Representative to identify the best method of contacting the potential contributors. All third party contributions must comply to the “Third Party Involvement in NHLBI-Supported Clinical Trials and other Population-Based Studies: Awardee/Contractor Third Party Related Issues” found at:

http://www.nhlbi.nih.gov/funding/policies/thirdparty.htm

The Business Representative informs the NHLBI Project Officer that contributions are being sought for a particular protocol and obtains permission to proceed.

The Business Representative prepares a Memorandum of Agreement (MOA) utilizing the NHLBI approved template if possible and negotiates the MOA with the potential contributor(s) in consultation with the Protocol Chair(s) or other designees. Upon completion of negotiations, but prior to execution of the MOA, the Business Representative completes the NHLBI checklist, and obtains appropriate DCC and government representative signatures. The MOA may then be executed with the Contributor(s).
3.7.3.1. Contributor Agreements

The MOA with a BMT CTN Contributor should reflect, at a minimum, the following:

- Contributor has no influence on the governance or conduct of the Study, or in the analysis, interpretation, or reporting of its results; except in limited circumstances when Contributor provides support in development of study design and study conduct activities, such activities will be reflected in the MOA;

- Contributor has no commitments with the study investigators that relate to the subject matter of the study as to intellectual or tangible property or to other issues that conflict with Public Health Service policy on grants or contracts;

- There is reasonable evidence that a Contributor’s involvement will not create any conflict of interest issues with the study investigators, the National Institutes of Health (NIH) and its Institutes or employees, or any appearance of any such conflicts of interest;

- Language reflecting any resulting inventions must adhere to the “Bayh-Dole Act” - 35 U.S.C. § 200-212. In addition, the BMT CTN will recognize that:
  - Contributor owns rights to their inventions.
  - Any inventions made jointly by employees or agents of BMT CTN and Contributor will be jointly owned.
  - If a Contributor insists on language that the sites must adhere to, and the BMT CTN and government agrees, such language will be incorporated into the study specific CSPRs; and

- In exchange for financial or in-kind support, the BMT CTN, upon approval from the NHLBI, can provide reports on trial status, i.e., accrual reports, BMT CTN’s Annual Report, manuscript review, etc. Requests for additional information must be approved by BMT CTN leadership, NHLBI and NCI.

Financial contributions are held at the NMDP/Be The Match in a designated account, to be utilized for costs related to that protocol, unless otherwise approved by the contributor.

Questions regarding the budget and contracting process should be directed to bmtctn@NMDP.org.

3.8. Procurement Guidelines for BMT CTN

The DCC is responsible for procuring services as required by the clinical protocols such as correlative laboratory studies, storage of laboratory samples, drug acquisitions and distributions, etc. The DCC Business Representative is responsible for placing Requests for Proposals (RFPs) or Requests for Quotations (RFQs), negotiating pricing, terms and conditions, and for placing contracts or purchase orders for the procurement of goods and services for the Network.

Two approaches to the procurement of services are used by the BMT CTN: i) full and open competition, or ii) limited competition (select or sole source). Open competition is the preferred method in accordance with 45 CFR 74.43, which states “All procurement transactions shall be conducted in a manner to provide to the maximum extent practical, open and free competition.”
Applicable procurement requirements, such as 45 CFR 74.40 through 74.48 are followed for both approaches.

When an open competition is used, the Business Representative will develop, issue, and manage RFPs/RFQs. The Protocol Chair or his/her designee will provide assistance with identifying potential suppliers, developing the statement of work, selecting experts to serve on the proposal review committee and establishing the technical review criteria. Persons involved in the procurement process will not disclose any information about the procurement to prospective offerors unless such disclosure is authorized by the DCC.

Under certain circumstances, using other than full and open competition (e.g., select or sole source awards) may be appropriate. Circumstances that may justify the use of other than full and open competition include: (a) the availability of only one or a limited number of qualified sources; (b) unusual and compelling urgency; and (c) the need to maintain an adequate base of suppliers.

Select or single source procurements over the simplified acquisition threshold (currently set at $150,000), must have sufficient documentation to justify the procurement. To initiate a select or sole source procurement, the Protocol Chair(s) or his/her designee must complete a Selected/Sole Source Justification form and contact the DCC Business Representative for review of the form. Next, the form is distributed to the BMT CTN Executive Committee for final approval or rejection. If approved, the DCC Business Representative proceeds accordingly. An RFP or RFQ will be issued to the selected source(s) or sole source, and in general, offerors will still be required to submit cost or pricing information.

Please direct questions regarding subcontracts or agreements for the BMT CTN to the Business Representatives at bmtctn@NMDP.org
CHAPTER 4

PROCEDURES FOR APPROVAL OF PROTOCOL AMENDMENTS
4. PROCEDURES FOR APPROVAL OF PROTOCOL AMENDMENTS

Approved protocols may require amendments to address issues that arise during study conduct (e.g., eligibility changes, toxicity review). In addition, new information may arise from other studies that affect the conduct of an ongoing study (e.g., new FDA or CDC guidelines). This chapter defines the steps for preparing amendments for approved and/or active protocols.

4.1. Proposal of a Protocol Amendment

Proposals for protocol amendments may come from the Protocol Team, DCC, Steering Committee, DSMB, NMDP IRB, participating clinical centers, NIH or the FDA. All proposals should be submitted to the Protocol Coordinator.

The Protocol Coordinator is responsible for:

- Recording the requested modification
- Contacting the submitting individual for any clarification
- Distributing the proposed amendment and/or document with proposed changes and the rationale to the Protocol Officer and the Protocol Team
- Assigning and maintaining protocol version numbers, copies of amendments and documents of changes in collaboration with the Administrative Coordinator

The Protocol Officer may contact the submitting individual to further discuss and clarify the proposed amendment. Additionally, changes recommended by the DSMB should be discussed with the NHLBI Project Officer.

4.2. Review of Proposal for Protocol Amendment

The Protocol Officer and Protocol Statistician will review the proposed amendment and the rationale and may discuss the changes with the Protocol Chair(s) and DCC Leadership. A decision regarding the need for a protocol amendment is the responsibility of the Protocol Officer in consultation with the Protocol Team.

Once approved by the Protocol Team, input may be solicited from the Steering Committee. If there is a disagreement regarding the protocol amendment, the Executive Committee will review the justification and rationale for the amendment and arbitrate. The decision of the Executive Committee to proceed with a proposed protocol amendment is binding.

4.3. Finalizing the Protocol Amendment

All proposed protocol amendments containing substantive changes must be reviewed, recommended for approval and signed off by the NHLBI Office of the Director before submission to the NMDP IRB and distribution to Clinical Centers. The DSMB review may take place at a DSMB meeting, on an arranged conference call, or by ballot.
Amendment proposals are distributed through the DSMB Executive Secretary, or his/her designee, following the process for new protocol reviews. Amendment proposals distributed to the DSMB for review three weeks prior to the scheduled DSMB review should include the following:

- An indication of old and new version numbers
- A summary of protocol and informed consent changes (if any) with “change document”
- A revised protocol with changes highlighted may also be included
- A rationale for changes may be provided
- A ballot to collect reviewer’s comments

The NHLBI Executive Secretary combines the reviewers’ comments into one document and forwards the document to the DCC/Emmes within 2-4 business days prior to the scheduled review. The DCC/Emmes then arranges for the Protocol Chair(s) and/or Protocol Officer to prepare a written response to be returned to the Executive Secretary at least one business day prior to the scheduled review. The scheduled review is then conducted during a teleconference or in-person meeting.

Final approval of the protocol amendment by the DSMB and the NHLBI Office of the Director is documented in the DSMB minutes supplied by the NHLBI within 14 business days of the review.

For BMT CTN studies under jurisdiction of the NMDP IRB, protocol amendments require NMDP IRB or IRB administrative review. Amendment proposals submitted to the NMDP IRB for review should include the following:

- NMDP IRB Request for Amendment form
- DSMB approval documentation
- Record of revision
- Protocol document, if revised
- Consent and assent (if applicable) documents, if revised

The NMDP IRB informs the DCC if the amendment is approved, approved with stipulations, or not approved. If stipulations or resubmission is required, the DCC will prepare the response with input from the Protocol Team.

As a final step once the protocol is approved by the DSMB and NMDP IRB (if applicable), the Protocol Chair(s), Protocol Officer and Protocol Coordinator must review the protocol document for completeness and accuracy and complete the Protocol Team Reviewer Checklist. Two Protocol Chairs or one Chair and the Officer must sign the front page of the document indicating their approval of the document. Participating centers will be notified of the approved protocol amendment by a Numbered Memorandum distributed via e-mail from the DCC. Amendment documents may be included with the e-mail announcement and will be posted on the BMT CTN SharePoint and public websites. The documents will include:
A summary of protocol changes
A revised protocol with changes red-lined
Revised Consent/assent forms with changes red-lined
A clean second version with the changes fully incorporated into the document
A rationale for changes may be provided
NMDP IRB approval, if applicable

Other types of amendments containing non-substantive changes may be released to Centers or submitted to the NMDP IRB for NMDP IRB for administrative review before DSMB notification or deliberation with permission of the NHLBI Project Officer. These changes will be reviewed at the next regularly scheduled DSMB meeting or conference call. The DCC PIs will determine the appropriate review process in collaboration with the NHLBI Project Officer.

4.4. Regulatory Authorities and Documents
Protocols under an IND/IDE require formal submission of the amendment to the FDA. This is done before or simultaneously with the DSMB submission or NMDP IRB/institutional IRB submission. Centers will use their local IRB for BMT CTN protocols released prior to July 1, 2017 for the duration of the study. The NMDP IRB will be used for all BMT CTN protocols released after July 1, 2017 and also for centers participating in the BMT CTN 1501 and 1503 NMDP IRB Pilot Project.

Local IRB Review
Participating Centers must obtain IRB approval of all protocol amendments for applicable protocols. After receiving IRB approvals, each Center must submit the IRB approval letter and approved consent form, if applicable, for the protocol amendment to the Protocol Coordinator. The Protocol Coordinator records the date of the protocol amendment approval, and consent if applicable, from each participating Center.

NMDP IRB Review
The DCC will distribute the NMDP IRB approval letter to Participating Centers via Numbered Memorandum. If the amendment included updates to the consent and assent (if applicable) documents, the Center must incorporate institution-specific language (as approved by the NMDP IRB, see Section 7.1) into the documents and submit to the DCC Protocol Coordinator for preview. Once approved, the Center sends the modified consent and assent (if applicable) documents to the NMDP IRB along with documentation of DCC Protocol Coordinator approval. If the consent/assent documents are translated into another language, the translated version must be provided to the NMDP IRB along with translator certification documentation. The NMDP IRB administrator will review the consent document(s), and if approved, send final document(s) back to the Center with NMDP IRB approval recorded on the documents and cc the Protocol Coordinator. If not approved, the Center will be notified they need to revise and resubmit the documents.
4.5. Other Study Related Revisions

4.5.1 Case Report Form Revisions

Case Report Forms (CRFs) and the study data file may require modification as a result of a protocol amendment. The Protocol Coordinator takes the following steps regarding CRFs and/or study data file revisions:

- Reviews the final amendment to determine if any changes to CFRs are required
- Drafts changes to the CRFs
- Reviews changes with aDCC PI, Protocol Statistician or Protocol Officer and Protocol Team, if necessary
- Implements changes and coordinates a software release with the Clinical Systems Analyst and/or programmer

4.5.2 Protocol Budget Revisions

The Protocol Chair(s), Protocol Officer, and Protocol Coordinator, in consultation with the Business Representative, will be responsible for reviewing the protocol amendment to determine if any changes will affect the protocol budget. If changes are required, the Business Representative will draft a revised budget. The revised budget will be routed in accordance with Chapter 3. If the per patient fee is affected, the budget summary will be circulated to all appropriate parties and posted to the BMT CTN SharePoint website by the Business Representative.

4.5.3 BMT CTN Services Revisions

The Protocol Chair(s), Protocol Officer and Protocol Coordinator are responsible for reviewing the amendment to determine if any changes that have been made affect the use of additional BMT CTN Services. If changes are required, the Protocol Coordinator notifies the BMT CTN Business Representative who re-negotiates contracts with the service provider and revises the budget as required.

4.5.4 Other

Other materials must be reviewed and appropriately revised to address the protocol modifications. These items include the Data Management Handbook and Users Guide, Forms Guide, Laboratory Sample Information Guide, Site Monitoring Plan, study initiation materials, various patient educational materials, study postings or Frequently Asked Questions (FAQs). DCC staff members such as the Protocol Coordinator, Protocol Officer as well as the Accrual Coordinator and the Laboratory/Repository Manager, all share responsibility in revising relevant sections within these documents. When necessary, the Patient and Health Professional Services Representative is consulted to participate in preparing revisions to the Informed Consent. Senior DCC staff members, and the Protocol Team as required, will review any revised information or documents prior to their implementation.

4.6. Additional Site Training

The Protocol Coordinator reviews the protocol amendment to determine whether any additional center training is required. The Protocol Coordinator is responsible for all training related to
protocol and CRF revisions. If other training requirements are identified, the Protocol Coordinator will work with the Protocol Officer to identify appropriate individuals to conduct the training. If related to a particular device, the manufacturer of the device will be considered as a potential source of training personnel. When such training requires contracting with outside organizations, the Protocol Officer will discuss with the DCC PIs, and forward the name, contact information and required activity to the Business Representative who will initiate subcontract discussions with the potential provider and determine the impact on the study budget.

4.7. Amendments for Collaborative Studies involving NCI NCTN Groups

These studies fall into two categories:

- Led by the BMT CTN
- Led by NCTN Group

For those studies led by the BMT CTN:

- Amendment materials are distributed by the Protocol Coordinator as early as possible to participating Centers and to the Central Operations Office of the appropriate NCTN Group
- Prior to release of the amended protocol, the NCTN Group follows their standard operating procedures for amendment review and approval

For those studies led by the collaborating NCTN Group:

- Amendment materials are provided to the DCC by the relevant Central Operations Office
- The Administrative Coordinator then circulates materials to all participating BMT CTN Clinical Centers per standard practice as noted above

Studies posted in the Clinical Trials Support Unit (CTSU) system will follow the CTSU amendment procedures.
CHAPTER 5

SITE MONITORING
5. SITE MONITORING

As part of the Quality Assurance plan and in full agreement with the NIH policy that states all clinical trials require monitoring to ensure the safety of study participants and the validity and integrity of the data (NIH Guide, NIH Policy for Data and Safety Monitoring, June 10, 1998), monitoring will be a continuous, ongoing and multifaceted process. This includes external review by the DSMB and IRBs, as well as internal data quality control, review and evaluation. Site monitoring visits are central to this process, and will include reporting to appropriate individuals with oversight responsibilities.

5.1. Initiation Site Visits

Prior to protocol implementation, the DCC Protocol Coordinator arranges either an initiation site visit or activation call with key center personnel to review all relevant materials and processes for implementation of the protocol. Multi-site activation calls may be held to ensure outreach to all appropriate personnel. A Protocol Officer or Protocol Chair in attendance on the call is highly recommended.

Prior to study initiation, each clinical site, pharmacy, and laboratory will be assessed to ensure each facility possesses the following:

- Adequate facilities and equipment to conduct the studies
- Site personnel adequately knowledgeable and trained in protocol(s) requirements, study policies, procedures and the data entry system
- Regulatory binder/file(s) have all required regulatory documents
- Adequate processes are established to protect the rights and safety of all study participants involved in the protocol

During the site activation call, the Protocol Coordinator will evaluate the pharmacy and clinical unit, patient record storage area, and computer facilities as well as ensure that adequate communication among all center staff is in place. Subsequent to the start of any study, all clinical sites will also be evaluated following these same procedures.

Laboratories will also be evaluated to confirm that the laboratory possesses the necessary professional certifications and licenses, and that laboratory operations quality assurance programs, assays, equipment and staff meet study requirements.

Site activation calls specific to pharmacists, lab coordinators, and/or clinic coordinators may also be held in the event additional training is required.

5.2. Follow-up Monitoring Visits

The DCC will conduct on-site monitoring visits periodically at the participating clinical centers, laboratories and pharmacies. The visits are conducted by a Protocol Monitor(s) or Clinical Research Associate(s) [PM(s)/CRA(s)] from the DCC and may be accompanied, as appropriate, by other DCC or NIH representatives. The purpose of the site visit is to ensure compliance with protocol requirements, regulatory requirements, study treatment, laboratory procedures, sample
acquisition, data submission and study policies and procedures. These visits are also used to exchange information regarding protocol adherence, review clinic, laboratory and pharmacy operations, provide training, check drug accountability, assess compliance with IRB reporting procedures for serious adverse events and violations/deviations, and discuss any problems encountered regarding implementation or compliance with the protocol design. A data audit comparing source documents to submitted data is performed during these visits. In addition, an overall assessment of management, coordination, and communications of the study site is conducted. Consequently, information can be easily exchanged, mutual problems can be resolved and study quality can be maximized.

On-site monitoring visits are conducted at least once every three years. Frequency of visits is dependent on accrual, site performance, number of patients enrolled, data quality, staff turnover and industry contributor and/or sponsor requirements. The purpose of the visit is to enhance data quality, ensure study integrity, satisfy regulatory requirements, and evaluate and improve site performance. An agenda for the site visit will be provided to the Principal Investigator(s) and site staff at least four weeks in advance of the visit.

The PM(s)/CRA(s) will hold a summary meeting, preferably at the end of the visit, with the Principal Investigator(s), the Lead Investigator and the site’s CRA(s)/CRC(s) to discuss the monitor’s observations, review any problems identified, and provide a preliminary report of the visit to the site. Within one week, an e-mail Memorandum is distributed by the PM(s)/CRA(s) to the center; any significant observations are noted in the Memorandum. In addition, a formal written report of the site visit is prepared by the PM(s)/CRA(s) and typically distributed within 60 days of the site visit. If site-specific problems of a serious nature are identified (e.g., failure to obtain informed consent, failure to have enrolled study participants sign the most current IRB-approved Informed Consent, enrollment of ineligible study participants, pharmacy or product administration errors, high data audit error rate), the Principal Investigator(s) may be requested to submit a corrective action plan for review by the DCC. Once all Action Items noted in the formal written Data Audit Site Visit Report have been resolved, the PM/CRA will issue an e-mail Memorandum closing out the site visit.

Industry-sponsored studies may have different monitoring requirements which will be defined in a study specific MOP, as an addendum to the BMT CTN Administrative MOP, or in the study specific Site Monitoring Plan.

5.3. Data Quality Assurance

Database quality will be maintained through a variety of analyses that target anomalies, delinquent data and key entry errors. Reports summarizing anomalies found are transmitted to the transplant centers for resolution. They are also posted on the AdvantageEDC℠ and Advantage eClinical℠ web sites. The DCC also uses this process to analyze the frequency of errors according to type to determine if certain types of errors are recurrent. Modifications to the data entry system or retraining of centers’ CRAs/CRCs will be employed for errors occurring frequently across transplant centers. If errors are localized within a transplant center, steps will be taken to resolve the problems by additional training to the center.
5.3.1 Data Review

The DCC Protocol Coordinator and/or Data Manager reviews data forms submitted by transplant center CRA(s)/CRC(s) for completeness, internal consistency, protocol compliance and adherence to the MOPs. In addition, “Missing Values Reports” and “Missing Forms Reports” are available through AdvantageEDC and Advantage eClinical and are dynamically updated. At minimum on a monthly basis, computer generated queries will be posted for each center. The queries will identify incomplete, questionable, or inconsistent data. Each center must either correct the data through AdvantageEDC and Advantage eClinical or provide an explanation on the validity of the existing data to the appropriate DCC Protocol Coordinator.

Transplant center CRA(s)/CRC(s) are expected to carefully check all data for completeness and consistency. Numerical values, such as hematological values are cross-checked to ensure accuracy. Validation may be required from the site to verify data. This is further elaborated upon in Chapter 9.

5.3.2 Missing Forms

Delinquent forms in AdvantageEDC and Advantage eClinical will be identified and compared to an exception list. All missing forms will be identified by form type for each study participant enrolled in a protocol. The web-based data entry system, AdvantageEDC and Advantage eClinical, will provide a table that summarizes all forms submitted and past due forms listed by study participant. After a form is entered, the list of forms submitted and outstanding forms is updated. A missing form will continue to be requested until the form is transmitted or until an exception is granted and entered into the missing forms exception file. Transplant Center CRA(s)/CRC(s) are required to review these tables for all study participants on a frequent basis.

Centers participating in BMT CTN trials must provide TED level data to the CIBMTR on all consecutive hematopoietic stem cell transplants performed at their institution during the period they are actively enrolling patients. Additionally, each center must then submit a Comprehensive Report Form for each patient participating on a BMT CTN trial (unless otherwise specified in the protocol) and follow the reporting requirements as outlined in the protocol-specific Form Submission Schedule. The BMT CTN Data Coordinator (located at the CIBMTR-Milwaukee campus), routinely works with transplant centers to obtain any missing and/or inconsistent CIBMTR data, including the collection of outstanding report forms.

5.3.3 Evaluation of Center Performance

The success of a multi-center network depends on high quality performance from the participating sites and careful coordination of effort. It will be the responsibility of the DCC to provide analyses and periodic reports on site performance with oversight from the NHLBI, NCI, DSMB, NMDP IRB and Steering Committee. The DCC is responsible for conducting site monitoring visits and for the administrative and statistical aspects of site evaluation. Reports are prepared and submitted to the NHLBI and Steering Committee according to specified guidelines. For the evaluation process to be successful, it is important to maintain open lines of communication among all parties, periodically review the common goals in order to maintain the highest degree of study integrity and ensure protection of human study participants within an environment that strives for continuous improvement of processes and operations.
Accrual reports for each protocol are prepared by the DCC, posted on the BMT CTN SharePoint website and updated nightly. Centers not meeting accrual goals will be contacted by the DCC and/or Protocol Team and Steering Committee Chairperson to determine the cause of slow accrual and if any corrective processes would improve accrual. The NHLBI and NCI will be informed of center-specific barriers to accrual.

Summaries of missing forms and days past due and data audit error rates, by center, will be provided to the DSMB at each meeting. The DCC will contact centers with serious delinquencies to resolve any training or staffing issues. After each data audit, the error rate by center will also be provided to the DSMB for review.

Periodic reports of center performance are provided to NHLBI and NCI. Serious violations, such as failure to obtain informed consent, enrollment of ineligible study participants, treatment or pharmacy errors, etc. however, will result in prompt notification to NHLBI and NCI. The DCC will analyze each serious violation to determine the impact of the error on study integrity. The issue will be discussed with the center and the center PI will be responsible for supplying a written explanation of the violation and corrective action taken.

Remote Monitoring Activity
In addition to site visits, the DCC routinely monitors accrual reports, CRFs, missing forms and responses to queries, critical data reporting, distribution of data, AE/SAE trending, incidence of protocol deviations/violations and occurrences. The Protocol Coordinator(s) has responsibility for identifying sites with problems in these areas and referring them to the Protocol Chair(s), Project Officers. The Protocol Chair(s) and Project Officer will determine whether corrective action is indicated. The corrective action may include, but not be limited to, discussion with the Principal Investigator, additional training of site personnel, a site visit, or referral to the Executive Committee.

Final Visit
For some studies, it may be appropriate to conduct final closeout visits, which occur after the study is completed and all data and finalized case reports are submitted. During this visit, the monitoring team determines study completion status. The review of regulatory compliance and documentation is done. The DCC policy for maintaining records is also reviewed with the site. The return of supplies and/or study medication(s) is also completed. After the visit, a final report, indicating the completion of the study, will be prepared.

5.3.4 Center Performance Reports
Effective scientific progress within the Network requires ongoing self-evaluation of BMT CTN procedures to maximize efficiency and enhance the scientific agenda. The BMT CTN provides Quarterly and Annual Center Performance Reports for each Core Center. The purpose of the report is to provide reward and encouragement for activities that further the BMT CTN goals; corrective advice for underperforming centers; and, methods for recommending a Center’s removal if their performance is not meeting the specified requirements. A copy of the Core Center Performance Evaluation Tool and Rating Schema is posted on the BMT CTN SharePoint website.
Evaluation of Center performance will include attention to both protocol-specific activities as well as intellectual and administrative participation in the Network. Recognition and reward of superior performance will be emphasized to encourage achievement as well as objective critique and discipline for inadequate participation.

Protocol-Specific Performance
The primary opportunity to advance the BMT CTN scientific agenda is timely and efficient opening of protocols and vigorous accrual of subjects for each study. Recognizing that centers of varying size may have differing patient base as well as institutional scientific commitments to other funded or developing studies, simple measures of periodic accrual may be insufficient.

The Annual Center Performance Report evaluates the following:

- **Network Scientific and Administrative Participation**
  Activities that directly foster the BMT CTN goals are recognized. Intellectual contribution to the Network can include participation as a Protocol Chair of a BMT CTN protocol; active membership on a Protocol Team including participation in team calls (for call participation credit, leadership or active participation on a BMT CTN Technical Committee; authorship or revision of components of the BMT CTN MOPs; election to a BMT CTN leadership position; and/or active scientific collaboration with the Network through study proposals, ancillary studies, scientific publications germane to BMT CTN protocols and supplemental grant applications and awards which extend the scientific leverage of Network projects and funding. Of note, a Protocol Chair or Team member cannot appoint another person as a substitute for Protocol Team calls. The member is responsible for reviewing the team minutes for a missed call upon receipt and indicate his/her understanding/agreement and stance via email.

- **Protocol Initiation and Activation**
  Protocol initiation, activation and enrollment will be monitored from the time of protocol release to submission of Informed Consent for preview; prompt preparation of needed revisions and approval; and, time to open each protocol at the Center.

- **Accrual**
  Commitment of patients to each study will be determined by pre-initiation polling of each center. Quantitative measures of accrual at each center will be judged against the benchmarks of the pre-study commitment; the number enrolled per year; the percent of center’s patients eligible who are enrolled adjusted as possible for the percent of patients who are not enrolled due to participation in a local, institutional competing trial. Centers are strongly encouraged to maintain commitment to each trial and avoid developing competing trials that directly reduce Network participation.

- **Study Procedures**
  Specific enumeration of protocol violations will be monitored. Major violations (e.g., enrollment of ineligible patient or serious deviation from protocol specified activity) will be reviewed on a continuous basis while minor violations (deviations from procedure that do not compromise the patients’ safety or the study endpoint) will be tracked and reviewed semi-annually.
• Data Quality
  Overdue missing forms and data audit error rate will be quantified and reviewed semi-
  annually. Excess delay in data submission or data errors will be interpreted as reflecting
  insufficient staff training, supervision or commitment of the center PI to adhere to the
  Network goals.

• Laboratory Evaluation and Compliance
  Laboratory compliance, including sample collection, shipment and completion of
  appropriate CRFs are routinely monitored by the Laboratory/Repository Manager and
  support staff. The DCC provides complete instructional materials, training and educational
  opportunities for CRAs/CRCs to assist with the logistical procedures related to research
  sample collection, storage, testing and shipment.

Critique and Discipline
For Core Centers, minimum criteria for participation will include satisfactory performance in all
the above categories. Data for each assessment will be collected by the DCC through the data
tracking system that prospectively monitors each of these performance measures. Each center’s
performance will be reviewed annually by the BMT CTN Executive Committee. Inadequate
performance will be documented and reviewed with the Center PI by the Steering Committee
Chairperson and as needed, including one or more of the three BMT CTN DCC PIs. Failure to
improve performance documented at a 6-month re-review may lead to recommendation that a
Center be withdrawn from the Network.

Recognition and Reward for Superior BMT CTN Participation
Accrual over that expected, BMT CTN special projects and extraordinary efforts to enhance
Network goals will be recognized and rewarded. Performance awards will be determined by the
BMT CTN Executive Committee. Awardees can be nominated by Steering Committee members
or directly recognized by the Executive Committee.

Participation will be recognized by one or more of the following:
• Added authorship on BMT CTN publications, primarily for added accrual or specific
  scientific augmentation of a study’s progress
• Recognition of an Affiliate Center by making a representative an active voting member of
  the Steering Committee with travel support through the DCC for two years.
• Travel awards for Center research personnel (e.g., data managers, CRAs/CRCs,
  pharmacists) to attend either BMT CTN Steering Committee or Coordinator meetings,
  professional advancement meetings/courses, or the Tandem BMT meetings.

5.3.5 Protocol Violations and Deviations
Protocol violations and deviations are planned or unplanned departures from the IRB-approved
protocol, informed consent and/or study materials. A protocol violation is a serious noncompliance
that may affect the participant’s rights, safety, or well-being or the completeness, accuracy and
integrity of the study data. A protocol deviation is a less serious non-compliance.
5.3.5.1. Protocol Violation

A protocol violation is any change, divergence or departure from the IRB approved study protocol, consent document, or study materials that affect the subject's rights, safety, or welfare; or the completeness, accuracy and integrity of the study data. It may be a result of an error, fraud or misconduct and/or result in the exclusion of a patient from the study.

If the noncompliance meets any of the criteria below, it is considered a protocol violation. A single noncompliance may meet more than one criterion below. The list is not exhaustive.

- The noncompliance has harmed or posed a new, significant or increased risk of harm to the study participant
  - The participant received an incorrect dose(s) or dosing schedule of the study drug that may pose increased risk to the participant
  - Unauthorized manipulation of the study product or its storage, handling or administration that may pose increased risk to the participant
  - The participant met the withdrawal criteria during the study but was not withdrawn and as a result may be subject to additional or increased risk
  - The participant received an excluded concomitant medication that may pose increased risk to the participant
  - The protocol’s SOPs and/or required procedures were not followed resulting in the potential of increased risk to the participant
  - Study drug not held, reduced or discontinued as instructed by protocol

- The noncompliance compromises the scientific integrity of the data collected for the study
  - The participant was enrolled/randomized but did not meet the eligibility criteria
  - The participant did not receive the study-prescribed treatment assignment
  - The participant was not treated per protocol procedures that specifically relate to primary endpoints/outcomes of the trial (e.g., a repeat biopsy was not conducted within the protocol-specified window prior to the patient beginning study procedures)
  - Key data and/or samples were lost
  - A serious adverse event was not reported within the appropriate timeframe
  - The participant’s blinded treatment assignment was exposed to the participant and/or site staff without necessary approval for unblinding
  - The site stopped reporting data for a participant that did not specifically withdraw consent for data collection
  - The site stopped reporting data for a participant that did not meet the protocol requirements to do such
  - The investigator and/or the participant failed to comply with study requirements
  - Initiation of study treatment prior to enrollment
  - Any delay in study timeline; e.g., transplant delayed to donor availability
• Using procedures different from those specified in the protocol for assessing critical endpoints such as disease stage, relapse, progression, engraftment, chimerism etc

• The noncompliance is a breach of human subject protection regulations, policies or procedures, Good Clinical Practice, and/or or FDA regulations

  • Lapse in IRB approval
  • Failure to obtain informed consent prior to initiation of study procedures
  • Collecting optional research samples who participants who did not provide consent for the samples
  • Inadequate informed consent process
  • Participant consented via an expired consent
  • Consent form not properly signed, initialed and or dated by patient, witness and/or guardian if applicable
  • Falsifying research or medical records
  • Performing tests or procedures beyond the professional’s scope/credentials
  • Working under expired professional license or certification
  • Repeated deviations and minor deviations
  • Inadequate record keeping
  • Data breach in which participant’s patient health information was exposed or risked exposure via unsecured email, database etc.
  • Failure to produce or maintain proper documentation
  • Any other deviation to the protocol that the DCC or Principal Investigator considers significant

5.3.5.2. Protocol Deviation

A protocol deviation is less serious non-compliance with the protocol, typically resulting from unforeseen circumstances or in the best interest of the patient. Some of the violations listed that do not pose increased risk to the participant may be considered deviations. Other examples of protocol deviations include, but are not limited to:

• Follow up assessments completed outside of the protocol-specified window
• The participant received an excluded concomitant medication that is believed not to pose additional increased risk
• Co-enrolling a patient another study without pre-approval from the BMT CTN

The BMT CTN does not require reporting of minor protocol deviations to the BMT CTN DCC. Examples of minor deviations include:

• Missed assessments that do not compromise patient safety or a primary or recondary endpoint of the study
• Assessments completed outside of the protocol-specified date range
• The participant missed <10% of the doses of the study drug that does not subject the participant to any increased risks
- Missed collection of laboratory sample (it will be documented in lab compliance reports)

Planned or Requested Deviation
On the rare occasion, the BMT CTN DCC may approve a request from a Clinical Center Investigator to deviate from a protocol-specified eligibility criterion in order to enroll a patient who does not meet one of the eligibility criteria. Typically, approval will be granted for the requested eligibility deviation if it has been already been approved by the DSMB but is pending release to the clinical centers and IRB(s) of record. For other requests, the Protocol Officer will review the request, and if they deem it as unreasonable, the request is denied. If the Protocol Officer deems the request to be reasonable with no anticipated impact to patient safety or study outcome, he/she will consult with the Medical Monitor assigned to the study in addition to the DCC. If the DCC and Medical Monitor agree with the Protocol Officer’s assessment, the deviation will be permitted. If they do not agree, the request is denied. The Protocol Officer will inform the Investigator of the final decision. The Protocol Officer will also keep the Protocol Coordinator and Chairs apprised of the request and final determination. Approved deviations must be reported to the IRB of record. In addition, centers cannot enroll patients with pre-approved eligibility deviations in the database without the assistance of the study’s Protocol Coordinator.

There are commercial partners for some BMT CTN studies who do not allow planned deviations to eligibility criteria. For these studies, requests will be denied without review.

5.3.5.3. Reporting of Protocol Violations and Deviations
Reporting of protocol violations and deviations follow the same procedures. The Principal Investigator or designee is responsible for reporting protocol violations and deviations to the BMT CTN DCC and IRB of record within 5 business days of days of knowledge of the occurrence. Both protocol violations and deviations must be reported on the Protocol Deviation/Violation Form in the clinical database (where available) or via phone or email to the DCC/Protocol Coordinator for the study. At that time, the center will provide the DCC with a complete description of the deviation and a description of the plan to avoid similar deviations in the future. If applicable, the center and DCC will engage in a discussion to determine whether if the participant(s) impacted by the deviation will continue in the study. In addition, the Investigator will notify their IRB or the NMDP IRB according to the IRB-specific reporting requirements. Reporting of protocol deviations is discussed with centers during the site initiation call and documented in the pre-study initiation call report. Centers are instructed to to keep documentation of all protocol deviations.

Industry-sponsored studies may have different requirements for the reporting of protocol violations and deviations. These requirements will be defined in a study specific MOP, as an addendum to the BMT CTN Administrative MOP, or in the study specific Site Monitoring Plan.
CHAPTER 6

ADVERSE EVENT REPORTING
6. ADVERSE EVENT REPORTING

6.1. Definitions

Adverse Event - Any unfavorable and unintended sign (including an abnormal laboratory finding), symptom or disease temporally associated with the use of a medical treatment or procedure regardless of whether it is considered related to the medical treatment or procedure (attribution of definite, probable, possible, unlikely, or unrelated).

Life-Threatening Adverse Event - Any adverse event that places the participant, in view of the investigator, at immediate risk of death from the reaction.

Serious Adverse Event (SAE) - Any adverse event that results in any of the following outcomes: death, a life threatening adverse event, in-patient hospitalization or prolongation of existing hospitalization, a persistent or significant disability/incapacity, or a congenital anomaly/birth defect. Important medical events that may not result in any of the previous outcomes may be considered serious if they jeopardize the participant and require medical or surgical intervention to prevent any of the previously listed outcomes from occurring.

Unanticipated Adverse Device Effect (UADE) - Any serious adverse effect on health or safety, any life-threatening problem or death caused by, or associated with a device, if that effect, problem, or death was not previously identified in nature, severity, or degree of incidence in the application; or any other unanticipated serious problem associated with a device that relates to the rights, safety, or welfare of subjects.

Unanticipated Problem Involving Risks to Subjects or Others (UPIRSO) - any incident, experience, or outcome that meets all of the following criteria:

- Unexpected in terms of nature, severity, or frequency given the research procedures that are described in the protocol-related documents and the characteristics of the subject population being studied.
- Related or possibly related, meaning there is a reasonable possibility that the incident, experience or outcome may have been caused by participation in the research.
- Suggests that the research places subjects or others at a greater risk of harm (including physical, psychological, economic, or social harm) than was previously known or recognized.

Unexpected Adverse Event - Any adverse event, the specificity or severity of which is NOT listed in the study protocol, product inserts or informed consent document.

Expected Adverse Event – Any adverse event that is listed in the study protocol product insert or informed consent document as observed with the interventions being tested in the clinical trial. Additionally, expectedness for transplant treatment studies is associated with what is observed after a hematopoietic cell transplantation. For example: grade 4 neutropenia is observed in virtually all HCTs, and included among the expected effects of the transplant.
Attribution - The determination of whether an adverse event is related to a medical treatment or procedure. Attribution categories:

**Definite**
The adverse event is *clearly related* to the study drug/device/procedure/treatment(s).

**Probable**
The adverse event is *likely related* to the study drug/device/procedure/treatment.

*For BMT CTN studies:* the adverse event is not likely to be caused by the subject’s underlying medical condition or other concomitant therapy, and the nature of the adverse event or the temporal relationship between the onset of the adverse event and study drug/device/procedure/treatment administration lead the investigator to believe that there is a reasonable chance of causal relationship.

**Possible**
The adverse event *may be related* to the study drug/device/procedure/treatment(s).

*For BMT CTN studies:* the adverse event is not likely to be caused by the subject’s underlying medical condition or other concomitant therapy, and the nature of the adverse event or the temporal relationship between the onset of the adverse event and study drug/device/procedure/treatment administration lead the investigator to believe that there is a reasonable chance of causal relationship.

**Unlikely**
The adverse event is *doubtfully related* to the study drug/device/procedure/treatment(s).

**Unrelated**
The adverse event is *clearly NOT related* to the study drug/device/procedure/treatment(s).

*For BMT CTN studies:* the adverse event is most plausibly explained by the subject’s underlying medical condition or other concomitant therapy, or the adverse event has no plausible biological relationship to study drug/device/treatment.

**Common Terminology Criteria Adverse Events (CTCAE)** – a descriptive terminology developed by the National Cancer Institute (NCI) for use in reporting adverse events. The CTCAE includes a grading (severity) scale for each adverse event term. Exhibits 6-1-1 and 6-1-2 provide reporting requirements for BMT-related complex/multi-component events. A copy of the current CTCAE guidelines is located at [https://ctep.cancer.gov/](https://ctep.cancer.gov/).

**Grade** – Severity of the adverse event. Grades were developed using the following guidelines:

- **Grade 0** – No adverse event or within normal limits
- **Grade 1** – Mild adverse event
- **Grade 2** – Moderate adverse event
- **Grade 3** – Severe adverse event
4 – Life-threatening or disabling adverse event
5 – Fatal adverse event

6.2. Adverse Event Reporting Requirements

6.2.1. Unexpected Adverse Events

For the purpose of continuity, any reporting requirements for Unexpected Adverse Events will also include Unanticipated Adverse Device Effects (UADEs), unless a difference is explicitly stated.

Exhibit 6-2-1 provides unexpected adverse event reporting requirements for study centers participating in a BMT CTN study.

Reporting requirements are calibrated to the severity or seriousness of the event and the perceived expectedness of the event to the study drug/device/procedure/treatment, irrespective of the attribution of the event to the study drug/device/procedure/treatment. Each protocol provides specifics on the type of unexpected events required to be reported (i.e. unexpected grade 3-5 AEs or unexpected SAEs). All reportable unexpected adverse events must be reported to the BMT CTN DCC in an expedited manner from the time of enrollment, unless specified otherwise in the protocol.

Adverse events should be reported using CTCAE terminology and severity scales. In general, investigators should report adverse events as diseases or syndromes whenever possible, instead of reporting individual component symptoms, signs, laboratory abnormalities, and sequelae. Each reportable unexpected adverse event should have a single adverse event description. All reportable adverse events will be coded in the Medical Dictionary for Regulatory Activities (MedDRA) by the Safety Monitor.

Exhibit 6-2-1

**REPORTING UNEXPECTED ADVERSE EVENTS ON A BMT CTN STUDY**

<table>
<thead>
<tr>
<th>SEVERITY GRADE</th>
<th>ATTRIBUTION</th>
<th>TRANSPLANT CENTER REPORTING REQUIREMENTS</th>
</tr>
</thead>
<tbody>
<tr>
<td>5 - Fatal</td>
<td>All attributions</td>
<td>Submit the Adverse Event form and a summary of the event to the DCC within 24 hours of the event. For Grade 5, also submit study death form to the DCC.</td>
</tr>
<tr>
<td>4 - Life-threatening or Disabling</td>
<td></td>
<td>Submit all completed AE forms to the DCC within 4 working days of learning of the event. For Grade 5, the summary should include potential contributing causes of death.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Information reported for the adverse event must include: Name of adverse event, date of first onset, peak severity, relationship to study drug/device/procedure/treatment,</td>
</tr>
</tbody>
</table>
resolution date, actions taken with respect to administration of study drug/device/procedure/treatment, and other treatment for the adverse event.

<table>
<thead>
<tr>
<th>3 – Severe</th>
<th>All attributions</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Submit the Adverse Event form and a summary of the event to the DCC within 3 working days of the adverse event.</td>
</tr>
<tr>
<td></td>
<td>Submit all completed AE forms to DCC within 4 working days of learning of the event.</td>
</tr>
<tr>
<td></td>
<td>Information reported for the adverse event must include: Name of adverse event, date of first onset, peak severity, relationship to study drug/device/procedure/treatment, resolution date, actions taken with respect to administration of study drug/device/procedure/treatment, and other treatment for the adverse event.</td>
</tr>
<tr>
<td></td>
<td>Multiple recurrences of the same adverse event should be reported separately.</td>
</tr>
</tbody>
</table>

Note: For any adverse event that is ongoing at the time of death, the event should be considered ‘Resolved by Death’ with the date of death as the resolution date.

6.2.2 Expected Adverse Events

Exhibit 6-2-2 provides expected adverse event reporting requirements for study centers participating in a BMT CTN study.

All fatal (Grade 5) expected adverse events will be reported in an expedited manner to the DCC through submission of a Death Form. Deaths do not have to be reported within 24 hours for observational studies. Most protocol-specific life-threatening (Grade 4) and other non-fatal, non-life-threatening expected adverse events will be reported on study forms submitted on a defined forms submission schedule. Life-threatening (Grade 4) adverse events not collected on study forms should be reported in an expedited manner via a web-based AE system.

In addition, each Protocol Team must develop an interim analysis plan using the CTCAE grading scale (or Bearman scale if appropriate) to monitor protocol-specific expected adverse events. The plan will be included in the study protocol.
Exhibit 6-2-2

REPORTING EXPECTED ADVERSE EVENTS ON BMT CTN STUDIES

<table>
<thead>
<tr>
<th>SEVERITY GRADE</th>
<th>ATTRIBUTION</th>
<th>TRANSPLANT CENTER REPORTING REQUIREMENT</th>
</tr>
</thead>
<tbody>
<tr>
<td>5 – Fatal</td>
<td>All attributions</td>
<td>Submit study death form to the DCC within 24 hours of notification of the death. Submit death summaries and/or autopsy reports of the expected adverse event to DCC at the time of the report. The summaries should include potential contributing causes of death. Deaths do not have to be reported within 24 hours for observational studies.</td>
</tr>
<tr>
<td>4 – Life-threatening or disabling</td>
<td>All attributions</td>
<td>Submit study form(s) capturing data on the expected adverse event to the DCC at the form’s scheduled due date. If the event is not captured on a study form, report using the AE system in an expedited manner.</td>
</tr>
<tr>
<td>3 – Severe</td>
<td>All attributions</td>
<td>Submit study form(s) capturing data on the expected adverse event to the DCC at the form’s scheduled due date.</td>
</tr>
</tbody>
</table>

Note: Selected Grade 3-5 events will be tracked and regularly monitored by the DCC and DSMB as specified in protocol-specific monitoring plans.

Any expected event that requires expedited reporting based on the reporting requirements outlined in the protocol should be reported under the same timelines outlined by grade for unexpected adverse event reporting. Reports of secondary primary malignancies (SPMs) if required by protocol should be reported within 3 business days, when reporting is required by protocol.

6.3. Adverse Event Monitoring

6.3.1. Unexpected Adverse Events

Unexpected adverse events will be reported via a web-based AE system. The Safety Monitor will review daily all submitted unexpected adverse events. The Safety Monitor will forward the information to the Medical Monitor for review within 1 business day.

All unexpected adverse events will be reviewed by the Medical Monitor at, or associated with the DCC, within 2 business days of receiving the summary of the adverse event from the Safety Monitor. If the Medical Monitor requires additional information to make his/her assessment, transplant centers will have 4 business days to respond to the request for additional information.

The Medical Monitor has medical expertise relevant to the study protocol and may request the participant's treatment assignment (if a blinded trial) when reviewing the adverse event. The DCC
representative (Safety Monitor) is responsible for notifying the NHLBI Project Officer via a password-protected report sent via email within 7 days for all reportable and Medical Monitor-confirmed unexpected adverse events or of any concerns regarding the frequency or type of adverse event(s) on a study or study treatment arm. The NHLBI Project Officer (or designee) is responsible for reviewing the adverse event materials to determine if the materials are complete. If there are any concerns regarding the type or frequency of the event, the NHLBI Project Officer will request that the DSMB Executive Secretary notify the DSMB Chair. The DSMB Chair and NHLBI Medical Monitor will review the adverse event materials, determine if the information is complete, determine if additional DSMB review is required and make recommendations to the NHLBI concerning continuation of the study. Full documentation of the procedures will be available at the DCC. If the DSMB or NHLBI Medical Monitor requires additional information to make their assessment, the transplant center will have 4 business days to respond to the request for additional information.

The Medical Monitor will review a cumulative line listing of all adverse events reported in an expedited fashion on a quarterly basis (data will be reported in a blinded fashion, when the protocol intervention requires blinding). The Medical Monitor will review the adverse events for safety signals identifiable when reviewing in aggregate. He/She will also review the cumulative listing for events required by protocol to make a recommendation of events to be removed from the active listing (i.e. events that are expected and not otherwise required to be reported by protocol). The Medical Monitor may seek additional guidance from one of the DCC Principal Investigators, based on the expertise required, for their assessments as long as the DCC Principal Investigator’s institution is not participating in the protocol under consideration, and the DCC Principal Investigator is not considered to be otherwise in conflict by the NHLBI or by the Steering Committee. If there any concerns regarding safety, the NHLBI Project Officer will be notified immediately. The DCC or Medical Monitor will provide a written summary of the safety concern. In the event of a transplant and disease-specific Medical Monitor, the transplant Medical Monitor will provide the review.

The DCC will prepare semi-annual summary reports of all reportable adverse events for the NHLBI Project Officer and DSMB. The summary reports will be included in the semi-annual DSMB report. The report is sent as a password-protected document via email.

6.3.2 Expected Adverse Events

The DCC will prepare semi-annual summary reports of all expected adverse events for the NHLBI Project Officer and the DSMB. The summary reports will be included in the semi-annual DSMB report. The report is sent as a password-protected document via email. Grade 3-5 expected adverse events will be reported as defined in the protocol. In general, the Network uses a Core Toxicity Form for all studies. In addition, each Protocol Team augments this with protocol-specific toxicities of interest. All events on the Toxicity Form are a subset of the CTCAE relevant to the particular study. The Toxicity Form is submitted at regular time intervals defined by the Protocol Team.

The Medical Monitor will review events reported on the protocol-specific toxicity form, the GVHD forms and the infection forms semi-annually, when forms are included as a part of the protocol. This review will assess whether there are safety concerns that should be referred to the
DSMB. The Medical Monitor may seek additional guidance from one of the DCC Principal Investigators, based on the expertise required, for their assessments as long as the DCC Principal Investigator's institution is not participating in the protocol under consideration, and the DCC Principal Investigator is not considered to be otherwise in conflict by the NHLBI or by the Steering Committee.

Any concern regarding the type or frequency of a Grade 3-5 expected adverse event will be reported to the NHLBI Project Officer who will determine if referral to the DSMB is warranted. If required, data materials will be provided by the DCC. The DSMB Executive Secretary will arrange for review by the DSMB Chair. The Chair will determine if additional DSMB review is required and make recommendations to the NHLBI concerning continuation of the study.

The DCC will ensure that any additional reporting requirements defined by the NHLBI Project Officer, DSMB Chair and other oversight groups are identified and implemented.

The DCC in collaboration with the NHLBI Project Officer will determine the exact content of these summary reports and the reporting schedule.

6.3.3 Stopping Guidelines
The Protocol Coordinator, Safety Monitor and Emmes Statistician for each protocol will review the adverse events monitored for stopping guidelines on a regular basis to be specified in the protocol, but at least monthly. The Medical Monitor will provide a review quarterly, but may be requested to the review the stopping guidelines more frequently in the event that the guidelines approach the threshold. The medical monitor may request to have an expedited review of the trial by the DSMB based on this review.

6.4 Adverse Event Reporting and Management

Because all or most study participants in BMT CTN trials will be receiving potentially toxic preparative therapy, significant regimen-related toxicity is anticipated. Study CRFs are designed to capture information on these adverse events. Likewise, substantial mortality is anticipated and will be captured via filing of appropriate CRFs. In general, unexpected adverse events may require reporting until a protocol defined event within each protocol occurs (i.e. initiation of second intervention post-progression). Each protocol or other protocol material should be consulted for when adverse event reporting is no longer required. Any adverse event that occurs after the protocol defined event requires reporting only if the AE is felt to be related to the primary BMT CTN protocol.

6.4.1 FDA IND Reporting
If a study is under an FDA Investigational New Drug (IND), all suspected and unexpected fatal or life-threatening adverse events are reported to the FDA within seven calendar days after receipt of the information, following FDA guidelines. All other suspected and unexpected serious adverse events are reported to the FDA via a written report within fifteen days of receipt of the information (21 CFR 312.32). If the Medical Monitor assesses the event to be unrelated to the study, then the event will not require expedited reporting but will be included in a summary report issued annually.
All expected adverse events (i.e., those listed in the informed consent, product inserts, or study materials), including death and graft failures are reported to the FDA via annual reports.

The DCC is responsible for reporting to the FDA for studies under an IND on at least an annual basis.

6.4.2. FDA IDE Reporting

If a study is under an FDA Investigational Drug Exemption (IDE), all unanticipated adverse device effects will be reported to the DCC for assessment. Evaluation of UADEs will be communicated to participating investigators, all reviewing IRBs and the FDA within 10 days of the initial event notification to the DCC. If the Medical Monitor assesses the event to be unrelated to the device, then the event will not require expedited reporting but will be included in a summary report issued annually. All anticipated device effects are reported to the FDA via annual progress reports.

The DCC is responsible for reporting safety information to the FDA for studies under an IDE on at least an annual basis.

6.4.3. Guidelines for Reporting AE to IRBs and OHRP for NHLBI Sponsored Clinical Trials Network

6.4.3.1. Introduction

NHLBI guidelines on reporting adverse events on a clinical trial require that all IRBs) associated with the trial be provided with the following information:

- A description of the DSMB procedures
- Identification of the DSMB members' areas of expertise, excluding names
- Feedback after each DSMB meeting

Guidelines note that the Office of Human Research Protection (OHRP), DHHS issued a memorandum dated May 22, 2000, Continuing Review of DSMB-Monitored Clinical Trials, which authorizes IRBs to rely on current statements from DSMBs that the DSMBs have reviewed study-wide adverse events, interim findings and any recent literature that may be relevant to the research.

In the event that the DSMB identifies UPIRSO, the DCC will prepare a summary report for submission to the OHRP. Once the DCC receives a response from OHRP, Principal Investigators will be notified by the DCC via a memo.

Of note, these procedures are in addition to local IRB and/or the NMDP IRB as appropriate adverse event reporting requirements.

For trials using the NMDP IRB, UPIRSO events identified by the DSMB will be sent by the DCC to the NMDP IRB for final approval and potential submission to OHRP.
6.4.4 Adverse Event Documentation

AEs definitions and procedures for reporting and monitoring AEs will be documented in the study protocol and/or the MOP.

In addition, each Network study should have study-specific safety monitoring outlined in the protocol to ensure adverse events are reported and monitored in a timely manner. The protocol will be reviewed by the DSMB prior to implementation of the study and executed by the Network DCC following study initiation.

6.4.5 Providing Follow-up Information to Applicable IRBs

If the DSMB does not identify any safety or other protocol-related concerns, within 30 days after a DSMB meeting, the NHLBI Project Officer will prepare a Summary Report that will state that:

- A review of outcome data, adverse events, and information relating to study performance (e.g., data timeliness, completeness, and quality) across all centers took place on a given date
- The observed frequency of adverse events did not exceed what was expected and indicated in the informed consent
- A review of recent literature relevant to the research took place
- The DSMB recommended that the study continue without modification of the protocol or informed consent

The report will be sent to the DCC for distribution to each participating center’s PI. It is the responsibility of each PI to forward this information to their local IRB, if applicable. The DCC will distribute to the NMDP IRB, if applicable.

If the DSMB recommends protocol or informed consent changes, and NHLBI Leadership, accepts the recommendations, the NHLBI Project Officer will send those recommendations to the DCC for distribution to the participating PIs. The communication will include the rationale for such changes and any relevant data. The data provided will be determined by the DSMB members and NHLBI staff and will not go beyond what is necessary to explain the actions taken.

To maximize safety of study participants while maintaining the integrity of the ongoing trial, the NHLBI Project Officer will review all communications with NHLBI Leadership before sending them to DCC for distribution. All communication regarding protocol changes will be sent to the DCC within 30 days of the DSMB meeting. Depending on their nature these changes may need to be implemented rapidly. The DSMB recommendations and discussions with NHLBI Leadership will determine the distribution timeline. It is the responsibility of each PI to forward the distributed communication to their local IRB, if applicable. The DCC will distribute to the NMDP IRB, if applicable.
6.4.5.1. Early Phase Multi-Center Clinical Trials

Early Phase trials are not designed to assess clinical benefit. There is less clinical experience with the intervention and usually more concern about possible adverse events. In addition, the outcomes of the trial are not expected to change clinical practice.

If the DSMB does not identify any safety, or other protocol concerns, actual listings of percentages of adverse events may be provided to the local or NMDP IRB. Unless there are safety considerations, this disclosure of adverse event information would be done without separating the data by study group. The NHLBI Project Officer will provide a routine summary report of the DSMB meeting within 30 days of the meeting to the DCC for distribution to participating PIs. It is the responsibility of each PI to forward this information to their local IRB, if applicable. The DCC will distribute to the NMDP IRB, if applicable.

If the DSMB recommends protocol or informed consent changes, and the NHLBI Leadership accepts these recommendations, the NHLBI Project Officer will send those changes to the DCC for distribution to the participating PIs. The communication will include the rationale for such changes and any relevant data. The data provided for early stage trials will generally be more extensive than a late phase trial. The NHLBI Project Officer will already have been disclosing tables of adverse events, with the study groups combined, even if the DSMB had not identified safety concerns. The DSMB and NHLBI Leadership will determine the type and amount of data provided.

To maximize safety of study participants while maintaining the integrity of the ongoing trial, the NHLBI Project Officer will review all communications with NHLBI Leadership before sending them to DCC for distribution. All communication regarding protocol changes will be sent to the DCC within 30 days of the DSMB meeting. Depending on their nature, these changes may need to be implemented rapidly. The DSMB recommendations and discussions with NHLBI Leadership will determine the distribution timeline. It is the responsibility of each PI to forward this information to their local IRB, if applicable. The DCC will distribute to the NMDP IRB, if applicable.

6.4.5.2. Gene Transfer Trials

Gene transfer trials have reporting requirements above and beyond those for trials of other interventions. They must follow the NIH Guidelines for Research Involving Recombinant DNA Molecules.

The NHLBI has a DSMB that oversees participant safety aspects of all gene transfer studies that it supports. Therefore, regardless of whether the individual study has its own data monitoring group, the NHLBI Gene Therapy DSMB will review all data.

Information on reporting adverse events and other data on gene transfer trials will be supplied by the NHLBI Project Office during development of the study protocol.
6.4.6 Requests from an IRB for Additional Information

If an IRB, whether for a data coordinating center or a clinic, requests information beyond what was agreed to at the beginning of the trial and beyond the above guidelines, the Protocol Officer will immediately inform NHLBI Leadership. Decisions as to how to handle each request will be made jointly, and will depend on the nature of the request, whether or not the DSMB had identified safety concerns, the kind of trial, the stage of the trial, and perhaps whether the IRB is for a coordinating center or a clinic. Possible options are:

- The NHLBI Project Officer will send a letter to the PI stating that the DSMB is carefully monitoring outcomes and adverse events and that if anything occurs that would alter the protocol or informed consent, each investigator and IRB will be informed promptly. The NHLBI Project Officer will ask the PI to discuss this letter with their IRB.

- For late phase trials, the IRB chairperson will be provided with adverse event data, without breaking down the information by treatment group. The IRB chairperson and one other member of the IRB would be authorized to see these data, in order to assure them that serious, unexpected events are infrequent. These data would be for the entire study, regardless of whether the IRB is for a coordinating center or a single clinic. (Note that for early phase trials, this information would generally already be provided to the IRBs.)

If neither of the above options is acceptable to the IRB, NHLBI Leadership will discuss whether or not to provide the data requested by the IRB. If it is decided that data will not be provided and the IRB still objects, participation of the center in the study may cease.
CHAPTER 7

HUMAN SUBJECTS PROTECTION
AND
REGULATORY PROCEDURES
7. HUMAN SUBJECT PROTECTION AND REGULATORY PROCEDURES

7.1. Institutional Review Board

Each transplant center requires IRB review and approval prior to participation in a protocol. Centers will use their local IRB for BMT CTN protocols released prior to July 1, 2017 for the duration of the study. The NMDP IRB will be used for all BMT CTN protocols released after July 1, 2017 and also for centers participating in the BMT CTN 1501 and 1503 NMDP IRB Pilot Project.

Local IRB

Documentation of the IRB review must be available at the transplant center and submitted to the DCC along with a copy of the IRB-approved Informed Consent prior to study initiation. Each center’s IRB of record is responsible for reviewing the protocol for continuation on an annual basis or as defined by its IRB.

On an annual basis, or as defined by each center’s IRB of record, the IRB will review the protocol for continuation. The status of annual IRB reviews will be maintained by the transplant center and a copy forwarded to the DCC. The DCC will provide a reminder letter to the transplant center at 60 days and 14 days prior to IRB expiration to facilitate timely renewal. In addition, a notification will be sent to the transplant center on the day of expiration, informing them that their enrollment is suspended until they receive IRB approval. A final reminder is sent 14 days post expiration, notifying centers that they will be de-activated if they do not receive IRB approval.

NMDP IRB

The NMDP will serve as the single IRB for all BMT CTN protocols released after July 1, 2017. All BMT CTN Core, Consortia and Affiliate Centers must enroll in the NMDP Single IRB prior to opening a new study. Enrollment steps are detailed in the NMDP IRB Manual for Local Institutions Using the NMDP IRB as a Single IRB. Study-specific requirements to be completed after enrollment for each new study are also outlined in the manual. Centers should contact NMDP Human Research Protection Program staff members for more information: NMDPSIRB@NMDP.org.

Documentation of current NMDP IRB review and approval must be available at the transplant center along with a copy of the NMDP IRB-approved institutional informed consent and assent (if applicable) documents prior to study initiation. The NMDP IRB annual review documentation will be distributed by the DCC via Numbered Memorandum.

7.2. Health Insurance Portability and Accountability Act (HIPAA)

It is the responsibility of each participating transplant center to understand their institution’s requirements under the Health Insurance Portability and Accountability Act (HIPAA). This website is an additional resource to provide guidance in assuring compliance: http://aspe.hhs.gov/admn simp/.

7.3. Office of Human Research Protections (OHRP) Institutional Assurances

Each participating institution must have a current Federal Wide Assurance (FWA) on file with OHRP and provide documentation of this number to the DCC. Transplant centers that are
identified as potential new participating sites and do not have a FWA, are required to obtain one from the Office of Human Research Protections, prior to site activation.

### 7.4. Participation of Women, Ethnic Minorities and Children

NIH sponsored clinical trials and the Code of Federal Regulations state that study participants enrolled in clinical trials must include women, ethnic minorities and children to facilitate potential benefit to all persons at risk for a particular disease, disorder or condition under investigation (45 CRF 46).

Research sponsored by NIH must include children (individuals under the age of 21) unless there are scientific and/or ethical reasons for exclusion. It is the responsibility of the Special Populations (Pediatrics/Human Subjects) Committee to review each BMT CTN protocol to ensure that children, women and minorities are included to the fullest extent possible. Sites should make every effort to include these populations in subject recruitment activities.

### 7.5. Site Regulatory Documents

In order for a site to be activated for participation in a BMT CTN clinical trial that is under an IND or IDE, the following materials must be submitted to the DCC for entry into a regulatory tracking system by the Regulatory Coordinator:

- **Form FDA 1572 for IND studies or Investigator Agreement** for IDE and non-IND/IDE studies - Statement of clinical center’s Principal Investigator
- **Financial Disclosure** - provides information regarding the financial agreement between an investigator/institution listed on the Form FDA 1572 and/or contributor/collaborator for the trial
- **Curriculum Vitae** – a current CV for the Principal Investigator and Co- Investigator(s)
- **Medical Licenses** - current professional licenses for select personnel
- **Delegation of Authority Log** – the log documents the responsibilities assigned to the research team members and their dates of involvement in the project
- **Letter of Local IRB approval for each protocol, if applicable** - Sites will be required to submit an initial approval letter from the IRB prior to enrollment of any participation. In addition, sites will also submit annual documentation of IRB review and approval. For studies reviewed by the NMDP IRB, the DCC will distribute NMDP IRB review and approval documentation via Numbered Memorandum.
- **Local or NMDP IRB Approved Informed Consent Document** - must be submitted for initial approval of each protocol and all amendments thereafter
- **Federal Wide Assurance** number

Additional items are required for activation as described in Section 9.4.
7.6. IND or IDE Application

Filing of an Investigational New Drug (IND) or Investigational Device Exemption (IDE) application with the FDA is required whenever clinical investigations of an unapproved new drug or biological product or new device are to be studied. In the BMT CTN, an industry contributor providing the product under investigation may request that the company, NHLBI or a BMT CTN Investigator serve as the IND or IDE sponsor. It is expected that the DCC will provide technical and administrative services to prepare, distribute, and track the IND/IDE application and assume the reporting obligations of the IND/IDE sponsor unless contractual arrangements specify otherwise. Similarly, the DCC will assume responsibility for coordination and review of annual IND/IDE reports to the FDA and maintain electronic copies of all correspondence.
CHAPTER 8

PUBLICATIONS, ABSTRACTS AND PRESENTATIONS
8. PUBLICATIONS, ABSTRACTS AND PRESENTATIONS

8.1. Policy Statement

Research activities of the BMT CTN are intended to contribute knowledge to the field of hematopoietic cell transplantation. Definitive contributions are made through publications in peer-reviewed literature. Abstracts, public presentations, electronic postings and data sharing also contribute to public knowledge, but do not substitute for peer-reviewed publications and are given less scrutiny.

BMT CTN will comply with NIH public access policies including submission of study results and final manuscripts (e.g., clinicaltrials.gov, PubMed central, etc.). The BMT CTN will comply with all journal requirements (e.g., modified copyright transfer agreements, disclaimers, etc.) as relates to NIH policy.

Types of BMT CTN papers include:

- Primary report of data
- Report of secondary analyses
- Report of unplanned analyses from one or more studies
- Report of technical or administrative committees issues or analyses
- Report of ancillary study manuscripts, presentations or abstracts

8.1.1. Oral Presentations Related to BMT CTN Studies

Oral presentations to local groups, which are limited to the design or rationale of the BMT CTN sponsored protocols, are exempt from the review policy described above. Individuals making such presentations must confirm with the Publications Committee that the material will not be recorded, published or represented without Committee approval.

8.2. The Role of the Publications Committee

The Publications Committee is responsible for developing publication and presentation policies. All policies must be approved by the Steering Committee before implementation.

The Publications Committee reviews all proposed publications and presentations to assess scientific rigor and relevance to the BMT CTN mission. This review process ensures protection of proprietary information and study participant confidentiality and assesses the public impact of publication and/or presentation.

No participating institution, BMT CTN Technical Committee, Protocol Team or other individual may present or publish individual findings from work performed on study protocols or work related to BMT CTN meetings and conference calls without review of the Publications Committee, NHLBI and NCI. This includes methodologic or position papers related to BMT CTN protocol development or operations.
8.2.1. Membership

Members are identified from a slate of candidates put forth by the Nominating Committee. The Steering Committee approves these selected individuals. At least one member should represent an Affiliate Center.

8.2.2. Amendments to Publication/Presentation Committee Guidelines

Changes to the above policies will be subject to review, amendment and approval by the Steering Committee.

8.3. Review Timeline

Abstracts, presentations, and proposed publications relating to data obtained from BMT CTN protocols or to activities of BMT CTN Committees or Protocol Teams are to be submitted to the DCC. The DCC will distribute abstracts, presentations and proposed publications to the Publications Committee for review. The Committee will have five (5) business days to make recommendations to the Corresponding Author, concerning abstracts and presentations and ten (10) business days to make recommendations to the Corresponding Author concerning publications. If an expedited review is necessary, the Chairs may determine their review will suffice.

If Publication Committee members have concerns about the submitted materials and/or appropriateness of data, the Committee, submits questions and/or recommendations within five days to the DCC to forward to the Corresponding/Senior Author for resolution. The Committee, at its discretion, may choose to submit questions and/or recommendations to the Steering Committee for resolution if they determine that Steering Committee involvement is required. The Steering Committee is required to provide any responses to the Committee within ten (10) business days.

8.4. Primary Results Manuscript

Manuscripts reporting the results of each BMT CTN trial or BMT CTN methodologic or position papers are prepared and submitted in a timely manner. No clinical trial results are released, presented or published without review from the Publications Committee, NHLBI and NCI.

8.4.1. Data Analysis

The statistical analysis of trial data is performed by the DCC. Final decisions about patient outcomes and endpoints are the responsibility of the protocol team and/or Endpoint Review Committee and is documented in a Statistical Analysis Plan (SAP) if different from the protocol. Upon completion of the statistical analysis, the DCC issues the Data Analysis Report of the study. In general, the Data Analysis Report is available within two months of locking the trial dataset. The Protocol Chair(s) will monitor progress toward completion of the Data Analysis Report. The Protocol Chair(s) may ask the Publications Committee Chairs to assist with addressing delays in the completion of the Data Analysis Report.
8.4.2. Writing Responsibilities

Completion of the primary study manuscript is the responsibility of the Protocol Chair(s) or designee(s). The first draft manuscript is completed within three (3) months of receiving the Data Analysis Report and necessary supplemental analyses. Co-authors shall have access to the study Data Analysis Report and shall be afforded ample opportunity to contribute to completion of the manuscript.

If the Protocol Chair(s) or designee is unable to complete the first-draft manuscript in a timely fashion, the DCC has the responsibility to address the delay. If necessary, the DCC Principal Investigator may re-assign first-draft responsibility to another author who will become the Lead Author on the manuscript.

Upon completion, the Lead Author distributes the first draft manuscript to all co-authors. Author comments are used to generate subsequent drafts and the final manuscript. Authors should settle differences in interpretation by discussion and consensus whenever possible. If consensus cannot be achieved, the decision is made by majority vote. If necessary, the BMT CTN Steering Committee will adjudicate. In general, the time from completion of the first draft manuscript to the final manuscript submission should not exceed six (6) months.

8.4.3. Manuscript Requirements

All manuscripts of the BMT CTN are subject to the following requirements:

- **Titles**
  - The manuscript title should include the words “Blood and Marrow Transplant Clinical Trials Network,” if permitted by the Journal.
  - If not permitted in the title, manuscript text should include language such as “…. submitted on behalf of the Blood and Marrow Transplant Clinical Trials Network (BMT CTN).” The Methods section is the preferred section for this reference.

- **Authors**
  - Authors are encouraged to list BMT CTN as a secondary affiliation after their institution.

- **Acknowledgments**
  - General
    Other investigators not part of the authorship list but who have made significant contribution to the conduct of the study, as well as staff members from a Clinical Center, DCC or NIH, are noted. Each primary study manuscript must include a listing of all participating clinical centers and the responsible study physician at that center.

  - Government Sponsors
    Each manuscript must acknowledge all NIH funding sources for the study, including in every instance, funding from NHLBI and NCI to the DCC and the participating Clinical Centers using the following language: “The Blood and Marrow Transplant Clinical Trials Network is supported in part by grant #U10HL069294 from the National Heart, Lung, and Blood Institute and the National Cancer Institute.”
− Other Networks/Cooperative Group involvement
  − Manuscripts for BMT CTN-led studies involving collaboration with other Networks or Groups must acknowledge the Network/Group as specified by the Primary Network/Group Investigator and Publications Committee Contact involved with the study.
  − Manuscripts for studies endorsed by the BMT CTN but led by other Networks or Groups must acknowledge the BMT CTN appropriately. The DCC Business Representative will inform the Primary Network/Group Representative and Investigator of the procedure for publication review and acknowledgments.

− Non-government Support
  The Contributors must be acknowledged in concordance with the active Memorandum of Agreement (MOA). Generally contributors are provided a draft copy of all publications and allowed at least thirty (30) days to review the information and provide comments as detailed in the MOA.

  • The DCC Business Representative reviews materials in advance to confirm that contributors are accurately acknowledged
  • Protocol-specific and Cooperative Group Publication Instructions are located on the BMT CTN SharePoint website homepage, under Documents\Author Resources.

8.4.4. Approvals and Submission

  • The First Author and the DCC are responsible for submission of the final manuscript and for obtaining written approvals from all authors as well as the representatives of the DCC, NHLBI and NCI. The DCC uses an internal abstract and manuscript review checklists for this purpose prior to submission of materials to assure that all required approvals are obtained and that contributors and/or sponsors are acknowledged accurately. In addition, the BMT CTN Lead or Corresponding Author will conduct final review of the manuscript to ensure accurate and complete acknowledgments. For BMT CTN studies, the BMT CTN Lead Author is the First Author; for cooperative group led studies, the BMT CTN Lead Author will be assigned and may not be the First Author.

The Lead Author is responsible for obtaining approval from all co-authors. In addition the Lead Author should send all submission materials to the DCC for circulation to the following people 30 days prior to the planned manuscript submission (7 days for presentations/abstracts):

  − DCC co-PIs
  − Publications Committee
  − DCC Business Representative
  − NHLBI/NCI Program Officers
  − Primary Network/Cooperative Group and Publications Committee Contact representative, if applicable
  − Contributors, if applicable
The Corresponding Author should ensure that each author and the DCC, NHLBI and NCI representatives receives a final copy of the submitted manuscript, abstract or presentation.

Review, correction and return of galley proofs are the joint responsibility of the Lead and Senior Authors and the Protocol Statistician. For collaborative studies led by the BMT CTN, the Primary Network/ NCTN Group and Publications Committee Contact representative should also review the galley proofs. For NCTN Group led studies, the BMT CTN Lead Author will provide BMT CTN acknowledgment requirements to the First Author and request to review the final manuscript proof, or any deviations from the BMT CTN acknowledgment requirements, prior to manuscript submission.

8.4.5. Timelines

To ensure timely publication of study results, the timelines below should be followed:

- Dataset closure to Data Analysis Report – two (2) months
- Data Analysis Report to first draft manuscript – three (3) months
- First draft manuscript to submission – six (6) months

8.5. Abstracts, Public Presentations, Electronic Postings

8.5.1. General

Abstracts, public presentations of study data and electronic postings of study data must follow the same processes outlined for primary publications as noted above, but do not require the same levels of scrutiny and are handled expeditiously. The Publications Committee Chairs, or at the Chairs’ discretion, the Publications Committee, may approve requests pending NHLBI Program Office approval for abstract submission, public data presentation or electronic data posting. The DCC has the right to review and prioritize approvals based upon the availability of required resources.

8.5.2. Abstracts

All abstract authors must have made substantive contributions to the study. Acknowledgments of funding sources are not required in the abstract text but must be included in the presentation slides or poster materials, if the abstract is accepted.

Proposals for abstract submissions should be initiated well in advance of the abstract submission deadline. The Publications Committee prioritizes their review process in the following way: peer-reviewed publications receive higher priority than abstracts, public presentations and electronic postings.

8.5.3. Public Presentations and Electronic Postings of Study Data

In general, a complete set of summary slides is prepared by the Lead and Senior Author and Protocol Statistician upon completion and analysis of each BMT CTN study. Individual investigators who wish to prepare additional slides for public presentation must utilize data from the final Data Analysis Report. Such presentations must also be submitted to the DCC for submission to the Publications Committee Chairs, their designee, or at the Chairs’ discretion, the entire Committee, for review and approval. Each such presentation must also acknowledge the
authors, funding agencies, contributors and the participating Core and Affiliate Centers. Electronic posting of study data must not occur prior to publication of the study’s primary manuscript in a peer-reviewed journal.

8.6. Secondary Manuscripts

Manuscripts arising from the study outside the primary endpoints are considered secondary manuscripts. These include:

- Reports of secondary analyses pre-specified in a protocol
- Reports of other secondary analyses from one or more BMT CTN trials
- Ancillary studies
- Laboratory studies
- Methodological papers
- Position papers
- Other technical reports

All secondary manuscripts must follow the same review process as a primary results manuscript as detailed in Section 8.4.

8.7. Authorship Guidelines

Authorship guidelines are based on fairness, inclusion and stimulation of participation and compliance. Author rewards focus on intellectual input and effort extended during the lifecycle of the trial. The Publications Committee is responsible for the development and modification of guidelines for determining authorship. Authorship guidelines are ratified by the Steering Committee prior to implementation.

Authorship on BMT CTN publications is a privilege commensurate with both personal and center contribution to the research being presented and the Network as a whole. The primary requirement for authorship of a BMT CTN publication is a substantive contribution to the research effort and is a reward for many aspects of contribution including: (1) membership and active participation in the Protocol Team; (2) active accrual to the protocol; (3) timely and accurate reporting of data; and, (4) active participation in Steering Committee activities. These activities may include participation in the following areas: hypothesis generation, concept development, protocol development, study implementation, subject enrollment, data collection, data analysis, and manuscript preparation and finalization. In most instances, contributions in several areas must occur, and in every instance, authors are expected to contribute to manuscript preparation and finalization.

The fundamental premise of authorship designation is the reward for overall effort. The success of the Network depends on many people and authorship is used to reward participation in Network activities. First and senior authorship positions on the primary and secondary manuscripts should be broadly shared. First and senior authorship on the primary paper does not imply the same
privilege on any subsequent manuscripts resulting from the same completed protocol or related ancillary studies.

8.7.1 Authorship Eligibility Requirements

Each member of the Protocol Team has eligibility for authorship on the final primary manuscript from the study. In addition, each Clinical Center Principal Investigator whose center enrolls at least one subject in the study is eligible for authorship. Additional authors may be invited at the discretion of the Protocol Chair(s), in consultation with the Protocol Team, with approval of the Publications Committee Chairs or full Committee, if needed. Eligibility for authorship does not guarantee authorship. Eligible authors must still make substantive contributions, including careful review and contributions to the draft manuscript.

8.7.2 Establishing the Order of Authorship for Primary Results Manuscript

For each manuscript, the Protocol Chair(s) will recommend the co-authors and the order of authorship based on the Administrative MOP parameters and provides the authorship list to the DCC for approval. Any potential additional authors proposed outside the scope of the Administrative MOP should be reviewed and approved by the Publications Committee with documented reasoning. The DCC will review the authorship list to confirm if authorship is warranted, and request rationale from Chairs if needed. The DCC will also assess Protocol Chair authorship order based on accrual and participation metric posted on the SharePoint website, and get approval/adjudication from the Executive or Publications Committee if needed. If there are multiple Protocol Chairs, two Chairs share first authorship, provided they meet the below requirements and the journal allows this practice. The Publications Committee will adjudicate any disputes, and may also recommend additional authors after review of accrual tables and based on definitive and justifiable, well-documented reasoning.

Approved requirements for author order are as follows:

- **SENIOR (last) AUTHOR REQUIREMENTS**
  - PI of Core Center or DCC or Center that proposed or led the trial
  - Protocol Team member
  - Attended majority of Protocol Team Calls
  - Attended majority of Steering Committee Meetings
  - Author’s center actively participated in the trial
  - Author’s center showed commendable protocol compliance and data submission

- **FIRST AUTHOR REQUIREMENTS**
  - Protocol Team member (preferably Chair or Co-Chair)
  - Attended majority of Protocol Team Calls
  - Writes the first draft of the manuscript
  - Author’s center actively participated in the trial

- **AUTHORSHIP BASED ON PROTOCOL TEAM AND ENDPOINT REVIEW COMMITTEE (ERC) PARTICIPATION**
  - The second and third authors will be the Protocol Officer or Statistician. The order will be determined by the Protocol Chairs.
Each Protocol Team/ERC member will be an author, in order of accrual, if the following criteria are met:

- Attended the majority of Protocol Team/ERC calls
- Author’s center actively participated in the trial
- Author’s center had satisfactory protocol compliance and data submission

If the Protocol Team/ERC member is already represented as first or Last Author, an additional Second Author from that center will be added.

If a center has more than one member represented on the Protocol Team/ERC, only one member is eligible to be an author under this criterion. The other Protocol Team member(s) may be eligible for authorship if they meet any of the additional authorship criteria.

**AUTHORSHIP BASED ON CENTER PARTICIPATION IN ORDER OF ACCRUAL**

- The top five accruing centers will have a single author in order of accrual
- If one of these centers is the First or Last Author, a Second Author from that center will be added
- Author identity may be determined by center PI
- Author’s center must have also had satisfactory protocol compliance and data submission

**BMT CTN PUBLICATIONS LIST ALL CENTERS THAT PARTICIPATED**

- Any center that has enrolled at least one patient will be acknowledged in the publication

**CONSIDERATIONS FOR ADDITIONAL AUTHORS**

- A physician from an Affiliate Center can be named in acknowledgement of overall effort, exceptional accrual and/or supportive input throughout the lifecycle of the study
- An additional physician author from a same center mentioned above can be named if he/she has shown exceptional accrual or intellectual input
- These can be non-physicians from any center
  - Nurses
  - Study coordinators
  - Pharmacists
  - Statisticians

**IF THE JOURNAL OR ABSTRACT GUIDELINES LIMIT THE NUMBER OF AUTHORS, AUTHORS WILL BE DELETED IN THE FOLLOWING ORDER:**

- Remove authors in reverse order of center accrual
- Remove Protocol Team members and top accruing centers in reverse order of center accrual
- Remove Secondary Statistician
- Minimum author assignments will be First, Last, Protocol Officer and Primary Statistician

8.7.3. Establishing the Order of Authorship for Secondary Results or Ancillary Study Manuscripts

Secondary results manuscripts will follow similar authorship guidelines as described above for primary results manuscripts. However, special recognition is given to those who were not
considered for First and Senior Authors on the primary paper. Consideration as added co-authors is extended to other Protocol Team members such as DCC members, clinical research coordinators and junior Protocol Team members, especially those from centers with good accrual rates.

Protocol chairs and officers will assign, as soon as is feasible, a writing committee for secondary manuscripts. Proposals for secondary manuscripts will be reviewed by the Protocol Team and authorship appropriately assigned.

For secondary analyses or ancillary studies conducted outside the purview of the Protocol Team, the Protocol Chairs of the parent study must be invited to participate and co-author the manuscript. An exception to this policy is if the ancillary study uses biorepository specimens and/or clinical data from multiple studies and the analysis is not related to the parent study’s specific design.

8.7.4. Removing Authors
An author may be removed from the final manuscript upon failure to make substantive contributions to the overall project. The Lead Author may request that another author withdraw from authorship. The Publications Committee Chairs, or at the Chairs’ discretion, the Publications Committee shall adjudicate authorship disputes be simple majority rule. If an author is removed, another eligible author may be added.

8.7.5. Authorship on Joint Studies
These studies primarily involve the NCI NCTN Groups. The BMT CTN Steering Committee supports the approach that the two study chairs representing the collaborating groups hold the first and last author positions at the time of manuscript preparation; this should be decided in advance by the Protocol Team.

8.8. Conflict of Interest
In the event a member of the BMT CTN Publications Committee is asked to review a manuscript, publication or presentation in which he/she is listed as an author, they should recuse him/herself from adjudicating the document.

If both the BMT CTN Publications Committee Chairs are included in the author list, they will take responsibility for re-assigning an Interim-Chair to review the manuscript, publication or presentation.

If more than three members of the Publications Committee are listed in the author list, the Publications Committee Chairs will be responsible for assigning ad-hoc reviewers to review the manuscript, publication or presentation.
CHAPTER 9

CLINICAL CENTER PROCEDURES
9. CLINICAL CENTER PROCEDURES

Each BMT CTN Clinical Center is staffed, at a minimum, by a Principal Investigator (PI) and a Clinical Research Associate (CRA) or Clinical Research Coordinator (CRC). There may be additional physicians designated as co-investigators, as well as other administrative and research-related personnel.

9.1. Functions of the Principal Investigator

The responsibilities of the PI, who is a physician with substantial experience in both HCT and the performance of clinical trials, are to:

- Direct the activities of BMT CTN personnel at the Clinical Center
- Coordinate the scientific and administrative operations of the Clinical Center
- Oversee data integrity and participate in quality assurance measures such as initiation, data audit and close-out site visits
- Ensure adherence by Clinical Center personnel to the procedures described in and required by BMT CTN Protocols, protocol-specific SOPs if appropriate, and the BMT CTN MOPs
- Spend sufficient time in the Clinical Center to adequately observe study procedures
- Assure the Clinical Center's fiscal responsibility in the disposition of BMT CTN funds
- Prepare budgets and annual reports
- Obtain Local IRB approval for BMT CTN Protocols, Consent and Assent (if applicable) Forms and study participant materials; or obtain NMDP IRB approval for Consent and Assent (if applicable) Forms.
- Ensure that unexpected Grade 3-5 adverse events are reported in an expedited manner
- Ensure that any expected adverse events that require expedited reporting based on the reporting requirements outlined in the protocol be reported in an expedited manner
- Review and address concerns related to Core Center Performance Reports

9.2. Function of Lead Investigator

The PI may designate a Lead Investigator at his or her center for any BMT CTN protocol. The responsibilities of the Lead Investigator, who is a physician with substantial experience in both HCT and the performance of clinical trials, are to:

- Oversee daily conduct and progress of the study
- Interact with the CRCs to ascertain any implementation difficulties including enrollment, data submission, laboratory procedures, etc.
- Oversee data and participate in quality assurance measures such as initiation and data audit site visits
- Spend sufficient time in the Clinical Center to adequately observe study procedures
• Ensure compliance with BMT CTN Protocols, protocol-specific SOPs if appropriate and the BMT CTN MOPs
• Communicate with the PI and alert the DCC of any issues regarding protocol implementation, compliance or emergent issues relating to subject safety

9.3. Functions of the Clinical Research Associate/ Clinical Research Coordinator

The CRA/CRC is responsible for supervising daily operations in the Clinical Center and serves as primary contact for the study participants as well as for the DCC. The duties of the CRA/CRC are to:

• Ensure that potential BMT CTN study participants receive appropriate information about the study, including the Informed Consent documents
• Register study participants in the BMT CTN using AdvantageEDC℠ (Electronic Data Capture) or Advantage eClinical data capture system
• Ensure that all active research study participants have signed and dated the most current IRB approved Informed Consent
• Schedule participant appointments
• Ensure regulatory compliance
• Handle communications with the DCC regarding participant enrollment, data entry, missing forms, missing values, data anomalies, field discrepancies and data queries
• Notify the DCC as well as the IRB of record if appropriate within 24 hours of a Grade 3-5 Adverse Event
• Notify the DCC as well as the IRB of record if appropriate any expected adverse events that require expedited reporting based on the reporting requirements outlined in the protocol be reported in an expedited manner
• Notify the DCC of changes or impending changes in the Clinical Center personnel, and any changes of address and/or telephone number(s) of the Clinical Center
• Maintain a file of correspondence with the DCC
• Maintain a current transplant center roster of personnel addresses, telephone and fax numbers, e-mail addresses and notify DCC of changes as they occur
• Obtain necessary information about deceased study participants (e.g., death certificates, autopsy reports)
• Ensure compliance with the BMT CTN MOPs, BMT CTN protocols, protocol-specific SOPs if any and Numbered Memorandum
• Ensure the site has the most recently amended and IRB-approved version of BMT CTN protocols
• Check data forms for accuracy and completeness
• Ensure that study participant names, initials, dates of birth, addresses, phone numbers, social security numbers, and any other personal identifiers are removed from all materials sent to the DCC or attached to CRFs in AdvantageEDC or Advantage eClinical
• Submit complete data to the DCC in a timely manner in compliance with the BMT CTN AdvantageEDC User’s Guide, eClinical User’s Guide and protocol-specific Forms Guides
• Respond to data queries from the DCC in a timely manner
• Ensure compliance with laboratory procedures
• Register patients with the CIBMTR by obtaining a CIBMTR Research ID for all patients
• Develop a training system to ensure that personnel performing BMT CTN procedures are properly trained and certified in AdvantageEDC, Advantage eClinical, and GlobalTrace, participate in scheduled conference calls with the Protocol Coordinator(s) and/or CRAs and secure access to the BMT CTN SharePoint website
• Meet with the Protocol Coordinator and/or DCC CRA during site visits at the Clinical Center
• Report irregularities or emergent problems that can affect the data quality to the PI and the Protocol Coordinator
• Perform other duties as defined by the Steering Committee or Technical Subcommittees

Each CRA/CRC will be given access to the AdvantageEDC User's Guide, eClinical User’s Guide and protocol-specific Forms Guides to aid in completing data forms. Additionally, many protocol-specific items are distributed to the CRAs/CRCs including the protocol-specific Laboratory Sample Information Guide, Study Drug Manual etc. Materials may be stored electronically or in paper files.

9.4. Site Activation

Clinical Centers must be activated for enrollment by the DCC for each protocol. A protocol-specific checklist of required items for activation is distributed to Clinical Centers after the protocol has been released. The checklists typically include most of the following items:

• Lab normal values
• Most recent FACT, CAP and CLIA certifications
• FWA number
• Scheduled pre-study site initiation conference call or visit
• Form FDA 1572 or Investigator Agreement
• Roster for protocol personnel
• Delegation of Authority Log
• CV, Financial Disclosure and Professional License of select personnel
• Local IRB approval letter, if applicable
• DCC-approved institutional consent and assent (if applicable) documents prior to submission to Local or NMDP IRB
• Local or NMDP IRB approved institutional consent and assent (if applicable) documents
• AdvantageEDC, Advantage eClinical, and GlobalTrace online training attendance and completion of EDC practicum and GlobalTrace quiz by at least one coordinator
• Human Subjects Protection (HSP) training documentation for protocol personnel
• Good Clinical Practices (GCP) training consistent with principles of the International Conference on Harmonisation E6 (R2) for protocol personnel
• Signed Clinical Study Protocol Rider
• Clinical Site Contact Form for Central Pharmacies and/or Laboratories

Upon review and approval of these and protocol-specific items, if any, by the DCC, a site activation memo will be sent via e-mail to the study staff at the transplant center. This memo will contain information on accessing AdvantageEDC or Advantage eClinical, enrolling patients, and obtaining study-related materials.

9.4.1. Change in Study Staff
If the site plans to add or replace study staff after site activation, the Protocol Coordinator must be informed prior to the start date of the new staff member. The Protocol Coordinator will provide the list of items that need to be submitted, including updated Delegation of Authority Log and documentation of the staff member’s GCP/ICH E6(R2) and Human Subjects Protection (HSP) trainings. Additionally if there is a new PI assigned to the study, they must submit documents including FDA Form 1572 (if applicable), CV, medical license, GCP and HSP training documents and protocol-specific training.

9.5. Recruitment
A recruitment goal will be established for each Clinical Center for every BMT CTN protocol. Each center will develop a plan in order to achieve this goal. The sites will track study candidates along with each study participant after enrollment. Each center should review this plan periodically throughout recruitment in order to determine the effectiveness of the plan. The plan must outline methods to identify and enroll minorities, women and children (if appropriate), in strict adherence to NIH and DHHS policies. If the site is not achieving its recruitment goal in a timely fashion, the plan will be discussed with the DCC and modified.

Each Clinical Center must register study participants through the CIBMTR using the CIBMTR Research ID Assignment Form, which will be used to track both transplants and eligible transplant candidates to assist in meeting the recruitment goals and requirements of the BMT CTN.

Additionally, systematic evaluation of study participants transplanted or treated off protocol will identify potential barriers to accrual and possible protocol modifications to enhance accrual. Particular attention will be paid to differential enrollment rates for minorities, women and children.
9.6. Eligibility Screening

If a study participant appears to be eligible for a BMT CTN protocol, the following steps should be taken:

- The plan of the study should be reviewed with the potential study participant and any questions posed by the study participant should be answered completely.
- Upon decision to participate in the study, the study participant must sign the Informed Consent form document prior to enrollment and study specific procedures.

Once a study participant has been assigned a registration number, the number remains associated with the study participant and will not be reassigned.

9.7. Scheduling Study Participant Appointments

When scheduling appointments, the various time windows must be kept in mind. Efforts should be made to avoid missed visits and to keep scheduled visits as close to the target date as possible.

In the event that a study participant is moving to an area that is not near the Clinical Center, staff should encourage the study participant to return to the Clinical Center for their scheduled or follow up visits. If a study participant is not planning to return to the Clinical Center for a follow-up visit, the CRC should make arrangements to obtain the necessary information through the study participant's primary physician. In addition, the center can make arrangements with the DCC to transfer the participation to another participating center that has IRB approval for the study (see section below). The DCC supports the center in all efforts to collect data in this situation.

9.8. Preventing Dropouts and Missed Contacts

A primary objective of each protocol is to study the clinical course of study participants receiving protocol treatments and medications. To achieve this objective, it is essential that each patient be examined regularly at follow-up visits until the study is completed or until the patient expires. Missing information can bias the results of the study. Although occasional missed visits cannot be prevented, the study could be invalid if there are many missed visits, numerous patient drop-outs, and/or missed specimen draws. When data are incomplete, it is difficult to predict the direction of any bias resulting from the incompleteness. The only correct way to deal with missing information is not to have any. Preventing dropouts and missed visits is a responsibility shared by the entire site staff.

Prior to registering a patient, a line of communication should be established between the site and the patient's primary care physician. The need for long-term follow-up and data collection should be explained and understood by the primary care physician.

9.9. Checking Completed Forms

Before submitting data to the DCC, the CRA/CRC should carefully check all data for completeness and consistency. The CRA/CRC must also ensure that study participant names, initials, dates of birth, phone numbers, addresses, social security numbers, and any other personal identifiers are removed from all materials sent to the DCC or attached to CRFs in AdvantageEDC or Advantage eClinical.
Completeness and consistency: Every effort should be made to complete every field on each data form. Exception requests made be submitted for fields in which the data point will never be available at the center. Each form will be thoroughly reviewed at the DCC. Incomplete and inconsistent items will be queried by the DCC and clarification requested.

Numerical values: Numerical values such as hematologic data have expected ranges defined in AdvantageEDC or Advantage eClinical. If a value is outside the expected range, then a message is displayed to the user at the time of data entry. This type of message is not necessarily evidence of an error, but simply a request to verify that the number is correct.

9.10. Transferring Study Participants

All follow-up reporting requirements of BMT CTN study participants are the responsibility of the Clinical Center that initially registers the study participant. In the event of study participant transfer, it is the responsibility of the CRA/CRC at the originating center to initiate communication with the destination center. A study participant may be transferred from the originating center to another center only if the destination center has the study IRB-approved. In addition, all missing forms and data discrepancies must be resolved. Study participant charts, forms and BMT CTN information will not be forwarded to the destination center until the DCC receives confirmation of study participant transfer. The study participant must sign the destination’s center IRB approved consent prior to undergoing any study specific assessments or procedures. Subsequently, AdvantageEDC or Advantage eClinical reporting requirements of transferred BMT CTN study participants become the responsibility of destination Clinical Center staff.
CHAPTER 10

SHIPPING INSTRUCTIONS
10. BMT CTN SHIPPING INSTRUCTIONS USING FEDERAL EXPRESS

Samples are collected and shipped to repositories and central reference laboratories for many of the BMT CTN protocols. Protocol-specific Research Sample Information Guides are developed for each protocol that requires the collection and shipment of samples to a repository or central reference laboratory. These guides outline the procedure for collecting, processing, labeling, packaging, and shipping samples. The guides are posted on the BMT CTN SharePoint website.

Unless noted otherwise in the protocol-specific Research Sample Information Guide, all samples will be shipped via Federal Express (FedEx). The FedEx account number and BMT CTN protocol identifier (FedEx billing reference) will be provided to participating transplant centers to cover the cost of the shipment. The account number and reference code are listed in the protocol-specific Research Sample Information Guide. The account numbers may only be used for the shipment of samples to the protocol-designated repository or central reference laboratory. Unauthorized use of the account numbers is prohibited. Centers using the account numbers acknowledge that any unauthorized use will be subject to reimbursement.

10.1. Specimen Packaging Guidelines for BMT CTN Project

Specimen packaging and shipping guidelines are outlined in each protocol-specific Research Sample Information Guide. All specimens will be shipped in accordance with IATA regulations. See the IATA Dangerous Goods Regulations (59th edition, 2018) for detailed requirements. Note: the requirements may change annually.

- IATA packaging 650 – Biological Substance Category B
- IATA 620 - Infectious Substance, affecting Humans
- IATA 202 – Liquid Nitrogen
- IATA 954 – Dry Ice
- IATA section 5 – Packing
- IATA section 6 Packaging Specifications and Performance Tests
- IATA Section 7 – Markings and Labeling
- IATA Section 8 – Documentation (Shippers Declaration)

The shipper (BMT CTN clinical sites) is responsible for ensuring that packages meet all the Regulations

- Training Requirements are defined in Section 1.5 of IATA Dangerous Goods Regulations and must be provided to staff who package specimens for transport.

The following guidelines are generic. Packaging styles may vary depending on the vendor providing the packaging or the commercial shipper used.
Specimen Types

Infectious substances are substances which are known or are reasonably expected to contain pathogens. Infectious substances are divided into the following categories:

A. **Category A:** An infectious substance which is transported in a form that, when exposure to it occurs, is capable of causing permanent disability, life-threatening or fatal disease in otherwise healthy humans or animals. Refer to IATA Regulations for list of organisms, mostly cultures. These infectious substances must be assigned to UN 2814. The proper shipping name is **Infectious substance, affecting humans.**

B. **Category B:** An infectious substance which does not meet the criteria for inclusion in Category A. Infectious substances in Category B must be assigned to UN 3373. The proper shipping name is **Biological substance Category B.**

C. **Exemptions**
- Substances which do not contain infectious substances or substances which are unlikely to cause disease in humans or animals are not subject to these Regulations unless they meet the criteria for inclusion in another class.
- Dried blood spots, collected by applying a drop of blood onto absorbent material, are not subject to these Regulations.
- Patient specimens for which there is minimal likelihood that pathogens are present are not subject to these Regulations if the specimen is packed in a packaging which will prevent leakage and which is marked with the words **Exempt human specimen.**

<table>
<thead>
<tr>
<th>Exempt Human Specimen</th>
<th>Biological Substance, Category B (IATA 650)</th>
<th>Infectious Substance, affecting Humans (IATA 620)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Routine blood samples that have no indication on medical history or suspicion from physical appearance that the donor would be positive for a pathogen. Examples: routine DR, CT, HR, Work-up, IDM Screening, research, etc.</td>
<td>Samples where there is knowledge of or suspicion of a pathogen. Examples: Samples being shipped for confirmatory testing from an IDM reactive screen, knowledge of an infectious disease from health history or confirmed positive test. Positive CMV result is not included.</td>
<td>Infectious substance which is capable of causing permanent disability, life-threatening or fatal disease in otherwise healthy humans or animals if exposure occurs. Refer to IATA Regulations for list of organisms, mostly cultures.</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Specimen Shipment Requirements:</th>
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<th>Specimen Shipment Requirements:</th>
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### Packaging for Shipment

<table>
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<tr>
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</tr>
</thead>
<tbody>
<tr>
<td>2. Leak-proof secondary packaging. Example: sealable bio-bag or canister. <em>Note: Neither the primary nor the secondary packaging are required to meet the 95 kPa pressure requirements.</em></td>
<td>2. Leak-proof secondary packaging. Example: sealable bio-bag or canister.* <em>Note: Either the primary or the secondary packaging must meet the 95 kPa pressure requirements.</em></td>
<td>2. Leak-proof secondary packaging. Example: sealable bio-bag or canister.* <em>Note: Either the primary or the secondary packaging must meet the 95 kPa pressure requirements.</em></td>
</tr>
<tr>
<td>3. Absorbent materials capable of absorbing the entire contents must be placed between the primary receptacle(s) and the secondary packaging.</td>
<td>3. Absorbent materials capable of absorbing the entire contents must be placed between the primary receptacle(s) and the secondary packaging.</td>
<td>3. Absorbent materials capable of absorbing the entire contents must be placed between the primary receptacle(s) and the secondary packaging.</td>
</tr>
<tr>
<td>4. Multiple fragile receptacles (example: glass blood tubes) must be wrapped in cushioning material to prevent breakage. <em>Note: Bubble wrap or cloth is an example of cushioning material.</em></td>
<td>4. Multiple fragile receptacles (example: glass blood tubes) must be wrapped in cushioning material to prevent breakage. <em>Note: Bubble wrap or cloth is an example of cushioning material.</em></td>
<td>4. Multiple fragile receptacles (example: glass blood tubes) must be either individually wrapped in cushioning material to prevent breakage or separated to prevent contact between them. <em>Note: Bubble wrap or cloth is an example of cushioning material.</em></td>
</tr>
<tr>
<td>5. No contents list is required.</td>
<td>5. An itemized list of contents must be enclosed between the secondary packaging and the outer packaging.</td>
<td>5. An itemized list of contents must be enclosed between the secondary packaging and the outer packaging.</td>
</tr>
<tr>
<td>6. Outer packaging must be of adequate strength for its capacity, mass and intended use. At least one surface must have a minimum dimension of 100 mm x 100 mm (4 inches by 4 inches).</td>
<td>6. A ridged outer packaging must be used. At least one surface must have a minimum dimension of 100 mm x 100 mm (4 inches by 4 inches).</td>
<td>6. A ridged outer packaging must be used. The smallest external dimension must be not less than 100 mm x 100 mm (4 inches by 4 inches).</td>
</tr>
<tr>
<td>Packaging for Shipment</td>
<td>Specimen Shipment Requirements:</td>
<td>Specimen Shipment Requirements:</td>
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<tr>
<td>------------------------</td>
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</tr>
<tr>
<td>7. Outer packaging must be labeled with the words: “Exempt human specimen”.</td>
<td>7. Outer packaging must be labeled with the words: “Biological substance, Category B” in letters at least 6mm high (approx. ¼ inch). The UN3373 symbol must appear on the outer packaging adjacent to the wording.</td>
<td>7. Outer packaging must be labeled with the words: “Infectious substance, affecting humans” in letters at least 6mm high (approx. ¼ inch). The UN2814 symbol must appear on the outer packaging adjacent to the wording. An infectious substance hazard label meeting class 6 infectious substance label requirements is needed. Package must be marked durably and legibly on the outside of the package with the name and telephone number of a responsible person.</td>
</tr>
<tr>
<td>8. Notify FedEx for proper over pack shipping supplies and instructions for pick up.</td>
<td>8. Notify FedEx for proper over pack shipping supplies and instructions for pick up.</td>
<td>8. Notify FedEx for proper over pack shipping supplies and instructions for pick up.</td>
</tr>
<tr>
<td>9. A Shipper Declaration is not required.</td>
<td>9. A Shipper Declaration is not required.</td>
<td>9. A Shipper’s Declaration is required. See Section 8 of the IATA manual for instructions on filling out the Shipper’s Declaration documentation.</td>
</tr>
<tr>
<td>10. The package materials must be validated as a whole by the vendor to meet the requirements for strength under the IATA 650 packaging instructions.</td>
<td>10. The package materials must be validated as a whole by the vendor to meet the requirements for strength under the IATA 620 packaging instructions.</td>
<td></td>
</tr>
<tr>
<td>11. See specific IATA requirements when shipping with other dangerous goods. Dry Ice see IATA954 Liquid Nitrogen IATA 202.</td>
<td>11. See specific IATA requirements when shipping with other dangerous goods. Dry Ice see IATA954 Liquid Nitrogen IATA 202.</td>
<td></td>
</tr>
</tbody>
</table>

**Note:** This category had been called “Diagnostic Specimen” in the past. As of January 1, 2007, the verbiage must be Biological substance, Category B. If you require this grade of packaging, please clearly notify the vendor.

**Important!** The inner packaging material must be compliant with the markings on the outer box. Do not use the Exempt human specimen inner packaging materials with a box labeled Biological substance, Category B even if the sample is actually exempt.
<table>
<thead>
<tr>
<th><strong>Special Shipping Considerations</strong></th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Refrigerated 2 - 8°C</strong></td>
</tr>
<tr>
<td>(Frozen cooling packs or wet ice)</td>
</tr>
<tr>
<td>1. Place secondary packaging inside the outer box.</td>
</tr>
<tr>
<td>2. Place frozen cooling packs or wet ice between the secondary and outer box.</td>
</tr>
<tr>
<td>3. Support must also be provided to maintain the position of the secondary packaging as the ice or cooling pack thaw.</td>
</tr>
<tr>
<td>It is not recommended to use wet ice. If ice is used, the container for the wet ice <strong>MUST</strong> be leak proof.</td>
</tr>
<tr>
<td>The primary and secondary packages must maintain their containment integrity at the temperature of the refrigerant.</td>
</tr>
<tr>
<td><strong>Dry Ice</strong></td>
</tr>
<tr>
<td><strong>Reference IATA 954 packaging and labeling</strong></td>
</tr>
<tr>
<td>1. Place secondary packaging inside outer box.</td>
</tr>
<tr>
<td>2. Place dry ice between secondary packaging and outer box.</td>
</tr>
<tr>
<td>3. Support must also be provided to maintain the position of the secondary packaging as the dry ice dissipates.</td>
</tr>
<tr>
<td>1. Instructions for UN1845 apply.</td>
</tr>
<tr>
<td>2. The primary and secondary packages must maintain their containment integrity at the temperature of the refrigerant as well as pressure integrity if refrigerant is lost.</td>
</tr>
<tr>
<td>3. Dry ice when offered for transport by air, must be in packaging designed and constructed to permit the release of carbon dioxide gas and to prevent a build-up of pressure that could rupture the packaging.</td>
</tr>
<tr>
<td>The net weight of the dry ice must be recorded on the outside of the package.</td>
</tr>
<tr>
<td>5. Dry ice UN 1845 symbol.</td>
</tr>
<tr>
<td>When a Shipper’s declaration is not required, the following must be recorded in the “Nature &amp; Quantity of Goods” box of the Air Waybill in the following order:</td>
</tr>
<tr>
<td>• UN1845</td>
</tr>
<tr>
<td>• Proper Shipping Name (Dry Ice or Carbon dioxide, solid)</td>
</tr>
<tr>
<td>• Class 9</td>
</tr>
<tr>
<td>Net quantity of dry ice in each package.</td>
</tr>
<tr>
<td>Contact FedEx for proper shipment and forms.</td>
</tr>
</tbody>
</table>
### Special Shipping Considerations

<table>
<thead>
<tr>
<th>Liquid Nitrogen</th>
<th>Place secondary packages into appropriate nitrogen shipper.</th>
<th>1. Plastic primary and secondary containers must be capable of withstanding very low temperatures. 2. The primary and secondary packages must maintain their containment integrity at the temperature of the refrigerant as well as pressure integrity if refrigerant is lost.</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Reference IATA 202 packaging and labeling</strong></td>
<td>1. Plastic primary and secondary containers must be capable of withstanding very low temperatures. 2. The primary and secondary packages must maintain their containment integrity at the temperature of the refrigerant as well as pressure integrity if refrigerant is lost.</td>
<td><strong>NOTE:</strong> Insulated packaging containing refrigerated liquid nitrogen fully absorbed in a porous material and intended for transport, at low temperature, of non-dangerous products are not subject to these Regulations provided the design of the insulated packaging would not allow the build-up of pressure within the container and would not permit the release of any refrigerated liquid nitrogen irrespective of the orientation of the insulated packaging. (Dry shippers)</td>
</tr>
<tr>
<td><strong>NOTE:</strong> Insulated packaging containing refrigerated liquid nitrogen fully absorbed in a porous material and intended for transport, at low temperature, of non-dangerous products are not subject to these Regulations provided the design of the insulated packaging would not allow the build-up of pressure within the container and would not permit the release of any refrigerated liquid nitrogen irrespective of the orientation of the insulated packaging. (Dry shippers)</td>
<td>The package must be clearly labeled 1. “Do Not Drop-Handle With Care”. 2. Cryogenic Liquids symbol “Class 2 Non-flammable Gas (Division 2.2).” 3. UN 1977 4. Upright Package orientation symbol = 7.4.4 or 7.4.5 in IATA manual.</td>
<td>Contact FedEx for proper shipment and forms.</td>
</tr>
</tbody>
</table>
CHAPTER 11

ANCILLARY STUDIES

Management of Contracted Laboratories

Genetics Studies: NIH Data Sharing Considerations
11. ANCILLARY STUDIES

An Ancillary Study entails the collection from study participants of data and/or specimens, or the conduct of additional analyses of existing materials or samples that are outside the specific objectives of the primary study. Ancillary Studies may also be characterized by:

- Requiring a separate informed consent form or an “opt-in” section in the parent protocol;
- Placing an additional burden on participants;
- Utilizing biospecimens previously collected for unspecified future research;
- Requiring additional data or biospecimen collection during the course of the clinical trial;
- Being funded by a separate funding mechanism or source.

Retrospective studies that utilize prospectively collected data in clinical trials without biospecimens are not considered ancillary studies and are discussed in Chapter 12 of this manual of procedures.

11.1. Classification of Ancillary Study Proposals

Ancillary studies are broadly classified as laboratory or clinical depending on whether they involve the collection and/or analysis of biospecimens. Ancillary studies within the BMT CTN are further classified in two categories:

- Protocol-specific laboratory or clinical studies that are included in the main clinical trial protocol and informed consent but are not essential for assessing the primary endpoint. These studies are reviewed and approved in concert with the review and approval of the protocol as a whole. The processes for developing these studies are discussed within the scope of the Protocol Team. The Procurement Guidelines outlined in Section 3.8 of the MOP set forth the guidelines for selecting a supplier to perform the correlative study test.

- Ancillary studies that are not part of a primary protocol but which will use samples and associated clinical data previously collected or to be prospectively collected in the course of one or more clinical trials or which involve approaching subjects on one or more trials for supplementary data.

11.2. Consideration of Ancillary Study Proposals That Are Not Part of a Primary Protocol

Before ancillary studies that are not an integral part of a protocol can be endorsed by the BMT CTN, proposals must go through the review process delineated below (summarized in Figure 11-1) and be approved by the BMT CTN Executive Committee. A document summarizing the ancillary study proposal and review process detailed in the MOP can be found on both the BMT CTN public and SharePoint websites.

Once a BMT CTN clinical trial dataset is transitioned to the custody of the NHLBI and made available in Biologic Specimen and Data Repository Information Coordinating Center (BioLINCC) as a public research resource, associated biospecimen collections may also be accepted and transitioned to an Open public collection managed by the NHLBI/BioLINCC. All
inquiries regarding biospecimen and clinical data availability, and the subsequent submission of formal study proposals requesting such research materials should be submitted directly to the NHLBI by going to the following BioLINCC website page:

https://biolincc.nhlbi.nih.gov/studies/

11.2.1. Information to Help Plan Ancillary Correlative Laboratory Studies

Since investigators interested in conducting secondary analyses may not be familiar with the specific BMT CTN trials or the Network itself, some general information to allow efficient understanding of the clinical trial specifics, study populations, and data available are posted on the public BMT CTN website to encourage secondary analyses: BMT CTN secondary analyses resources

11.2.2. Process for Initiation, Review and Approval

Proposals for Ancillary Studies must be presented in writing to the BMT CTN DCC and/or Protocol Team by investigators who are known to be qualified in their field or who provide evidence of qualifications relevant to the proposed research question(s). Written proposals must be prepared by using the BMT CTN Ancillary Study Proposal Utilizing Biospecimens Form, which can be found on both the BMT CTN public and Sharepoint websites. The investigator must indicate potential sources of funding for the study, and indicate if funds are currently available or being sought.

For ancillary study proposals limited to one protocol, the initial review is conducted by the Protocol Team. An important factor in the review of Ancillary Study proposals is the determination that the objectives and implementation of the primary study are not compromised. Protocol Team members individually review ancillary study proposals and evaluate them for:

- Significance and appropriateness of the proposed research question;
- Design of the proposed research;
- Qualifications of the investigator(s) to conduct the research;
- Availability of funding and by whom;
- Availability of biospecimens and requested clinical data, if required for the study.

Protocol Team member proposal scoring will be captured using the BMT CTN Ancillary Study Proposal Reviewer Checklist that can be found on the private BMT CTN website. This checklist can be used for both retrospective laboratory studies, or for prospective studies proposed to be conducted in parallel to the primary trial. If the proposal involves collection and/or analysis of biologic specimens, the Protocol Team comments may be summarized and forwarded along with the proposal to the BMT CTN Biomarkers Committee for an additional review.

The Biomarkers Committee provides a second level of review, when necessary, focusing on the laboratory aspects of the proposal in terms of feasibility and scientific merit and putting the proposal in the context of other laboratory studies approved or ongoing on BMT CTN clinical trials. The Biomarkers Committee sends its evaluation back to the Protocol Team for their review. The Protocol Team may amend its evaluation of the proposal based on this critique.
The Protocol Team or designee then presents its recommendations regarding the proposed study to the BMT CTN Steering Committee, followed by the Executive Committee for final approval. Upon final approval the BMT CTN DCC records the decision using either a Parent Study and BMT CTN Leadership Approval Form BMT CTN Biospecimen and Clinical Data Resources Form (internal documents found on the private BMT CTN website), signed by a BMT CTN Executive Committee representative noting the BMT CTN committee making the final study approval.

For ancillary studies involving biospecimens but applying to several protocols, the primary proposal review is conducted by the Biomarkers Committee which then presents its recommendations, together with those of the various Protocol Teams, to the BMT CTN Steering Committee with final approval from the BMT CTN Executive Committee.

Any ancillary study requiring a separate informed consent must also be reviewed and approved by the BMT CTN Data and Safety Monitoring Board (DSMB).

![Figure 11-1: Overview of Ancillary Study Proposals Utilizing Biospecimens](image)

11.2.3. Ancillary Study Implementation

Once approved by the BMT CTN Executive Committee (and DSMB, if necessary), the Ancillary Study principal investigator will be notified of the endorsement by the DCC. Ancillary studies outside the main trial are generally voluntary for individual patient or institution participation. The BMT CTN DCC may elect to assess interest in study participation from centers in order to obtain a sample size matching the one of the parent clinical trial.
11.3. Ancillary Studies Requiring the Collection of Clinical Data

Ancillary studies requiring clinical data and no biospecimens will be implemented as a “clinical” study: requiring real time supplemental data collection on patients being enrolled on BMT CTN clinical trials and who are undergoing scheduled evaluations according to the protocol. Ancillary studies which require collecting additional data during the course of the clinical trial need a separate informed consent. Examples of such ancillary studies include quality of life assessments or other functional assessments that are not part of the main clinical trial. For ancillary studies of this sort, utilizing funding provided by through the BMT CTN DCC, or for ancillary studies utilizing in part BMT CTN Data, that are not part of the primary protocol, the Ancillary Study Principal Investigator’s institution must sign an agreement with the NMDP/Be The Match (acting on behalf of the BMT CTN) outlining the responsibilities, data handling, publishing rights and other details pertinent to the study before receiving data.

11.4. Ancillary Studies Utilizing Biospecimens and Associated Clinical Data

Ancillary Studies using biospecimens (associated most often with some clinical data) will be classified as “laboratory” studies. These studies will be implemented in slightly different ways depending on the categories of studies listed below:

- Requiring previously collected samples that are stored at the NMDP/Be The Match Biorepository
- Requiring real time supplemental sample collection during the course of the clinical trial (planned clinical trial ancillary study)

In all of the situations outlined below, the Ancillary Study Principal Investigator must sign the applicable contractual document (Research Materials Distribution Agreement) with the NMDP/Be The Match (acting on behalf of the BMT CTN), prior to receiving any samples and associated clinical data to conduct the approved research.

11.4.1. Retrieval of BMT CTN Biologic Samples from NHLBI Biorepository/BioLINCC

The NHLBI Biorepository/BioLINCC currently stores and manages biospecimens associated with several BMT CTN protocols for which further protocol-defined research tests are not planned (“true repository specimens”). All pertinent NHLBI policies and procedures will be followed to access these samples. All inquiries for information regarding the availability of and requesting procedures for these “Open” or public research biospecimen collections and associated clinical outcome data should be directed to the NHLBI by going to the following BioLINCC website page: https://biolincc.nhlbi.nih.gov/studies/.

11.4.2. Retrieval of Biologic Samples from NMDP/Be The Match Research Biorepository

The NMDP/Be The Match Biorepository (on behalf of the BMT CTN) has also stored and manages many biospecimens collected in some early BMT CTN protocols, and continues to store and manage the collections associated with all subsequent studies. For many of these studies, all protocol-defined testing has been completed, so the remaining biospecimens are considered “true biorepository specimens”. These biospecimens are made available to all qualified investigators with BMT CTN approved ancillary laboratory studies. For the more recent protocols, biospecimen collections are often associated with protocol-defined or Protocol Team planned correlative
laboratory studies. The biospecimen collections for those studies will be first made available to meet the requirements of these planned studies and then opened up to the BMT community to support future BMT CTN approved ancillary laboratory studies. All pertinent NHLBI policies and procedures will also be followed to access these BMT CTN biospecimens.

The process for formally requesting BMT CTN samples from the NMDP/Be The Match Repository is summarized as follows:

11.4.2.1. Pre-submission Requirements

Those submitting an ancillary study request must first contact the BMT CTN DCC and parent study Protocol Team to initiate a discussion regarding the evaluation/approval of planned studies as previously described.

11.4.2.2. Formal NMDP/Be The Match Sample Request Submission Process Overview

- Once the parent study and BMT CTN oversight committees have approved the ancillary study request, the study investigator is provided with a summary of the selected biospecimens available for the study.

- The Principal Investigator will approve the selected samples, and then a BMT CTN DCC representative on behalf of the PI, submits a formal BMT CTN sample request to the NMDP/Be The Match Research Biorepository.

- The elements for the proprietary study request must be completed. These elements include the center administrative and contact information, shipping information for the biospecimens, and a FedEx account number.

- The following documents comprise the approved application:
  - A signed Parent Study and BMT CTN Leadership Approval Form BMT CTN Biospecimen and Clinical Data Resources Form
  - Research plan summary
  - An electronic manifest listing the approved biospecimens, generated from the parent study representative
  - The full or expedited IRB approval/waiver provided by the investigator for the proposed research

- Selected vials are reserved and a BMT CTN Research Material Distribution Agreement (RMDA) is provided to the investigator, signed, and sent back for storage with the sample/data request application.

- The BMT CTN DCC representative is notified by the NMDP/Be The Match Biorepository when the review of selected sample inventory is completed and asked to provide final approval for pulling the identified study samples from the biospecimen inventory and shipping to the investigator.

11.4.3. Distribution of Sample Information and Associated Clinical Outcome Data

The BMT CTN assigns each study subject with two unique subject identifiers.
• The first identifier is the **Project ID**, which uniquely identifies the subject within the context of a single BMT CTN study (e.g., 0702). This is the primary subject identifier that is used in conjunction with the clinical data that is captured throughout that same single BMT CTN study. This subject identifier is used behind the scenes to uniquely identify protocol-specific clinical data that was collected for that subject.

• The second identifier is the **Patient ID**, which uniquely identifies the subject in the context of any BMT CTN study. This is the primary subject ID that is associated with both the AdvantageEDC data collection forms and any biological research samples that are collected in any BMT CTN study and entered into the GlobalTrace application. In the context of research biospecimens processed and stored by the BMT CTN Repository, the biospecimens will be associated with this unique **Patient ID**.

If a subject is enrolled on more than one BMT CTN protocol (e.g., 1101 and 1202), the **Patient ID** will be used for both clinical data and biologic samples being collected throughout the study. Behind the scenes, protocol-specific **Project IDs** will also be assigned each subject, which will unambiguously identify the protocol-specific clinical data collected from each study.

Biospecimen specific information will be provided by a DCC representative to the ancillary study investigator in the form of an electronic Excel shipping manifest. The sample manifest will contain a listing of the unique sample identifiers along with all pertinent sample-related information (e.g., collection date, sample type, volume, etc.) including the unique **Patient ID** associated with each sample. Associated clinical data sets provided to the investigator for the ancillary study (usually in the form of a SAS file) will contain both the unique **Project ID** and the unique **Patient ID** for each subject. The later identifier will serve as the data element link between the biospecimens listed in the sample manifest and the associated protocol-specific clinical data set.

11.4.4. Ancillary Studies with Real Time Sample Collection

Studies requiring real time collection during the course of the clinical trial, once approved, will be implemented with the primary clinical trial. Participation in the integrated ancillary study by transplant centers may be voluntary in some cases; patient participation is always considered voluntary. If not included within the primary clinical trial protocol and consent form, the PI will be required to develop an ancillary protocol and informed consent that will be reviewed by the NHLBI DSMB and clinical site or NMDP single institutional review boards before implementation. Additionally the Ancillary Study PI must sign an agreement with the NMDP/Be The Match, acting on behalf of the BMT CTN, outlining responsibilities, shipping and handling of samples, publishing rights among other details pertinent to the study prior to receiving any samples.

11.5. Funding for Ancillary Studies

Funding for Ancillary Studies may be from Network resources, an institution, private sources, or the NIH. The Steering Committee may recommend submission of a competitive supplement application to the NIH, in which case the application will be reviewed through the traditional peer review mechanism. If NIH funding is to be sought, the BMT CTN will provide a letter of endorsement signed by the Executive or Steering Committee Chair or a DCC Leadership member.
The available funding or grant application budget for all Ancillary Studies must include support for the BMT CTN DCC to cover data retrieval, selection of biospecimens to be released to the investigator, shipping costs, statistical analysis when provided and perhaps other logistical activities. Coverage for BMT CTN DCC costs will vary depending on the complexity of the Ancillary Study. The DCC will provide an approved budget for DCC costs for all approved studies and any other sub-contractor forms required by a prospective funder.

Any funding received from government agencies or third party funders, outside of the primary BMT CTN grants, shall be reported to the BMT CTN for inclusion in the tally of total contributions to the BMT CTN.

11.6. Ancillary Study Manuscripts

Please refer to Chapter 8 of the BMT CTN MOP (Publications, Abstracts and Presentations).

11.7. Management of Protocol-Defined Contracted Laboratories

The BMT CTN has partnered with numerous University-based investigators and Commercial laboratories in the U.S. to provide laboratory testing services to support protocol-defined laboratory studies. Many of the laboratories were selected in response to a release of a competitive RFP, while others were selected specifically by Protocol Teams for their unique capabilities, investigator expertise and/or documented scientific contributions to the field of BMT research. The BMT CTN DCC has developed and implemented procedures for executing lab service agreements that detail the scope of work required and other responsibilities and considerations related to the services the project laboratories are providing. Ensuring high quality services are provided that will effectively meet the requirements of each correlative laboratory study is of utmost importance. To this end, the BMT CTN DCC has developed and implemented procedures that are summarized below that will serve to develop a productive partnership and effectively support quality monitoring of the testing and final results submitted for analysis in our laboratory studies.

- At project initiation a BMT CTN DCC representative meets with the laboratory project staff to review the scope of work associated with the lab service agreement. The team discusses the collection and distribution of samples, receipt and processing (when necessary) of samples, and the analytical testing that is to be performed by the laboratory. Finally, invoice submission procedures for reimbursement for testing services are reviewed.
- Communication procedures are established between the project laboratory and a DCC representative so that there is a single point of contact for the submission of general questions and technical issues that need to be reviewed and resolved.
- Test result content and formatting is discussed and a final result template is developed. The final result template must be reviewed and approved by the protocol PI.
- Within the first few weeks of research sample testing, completed test data, in the agreed upon format, is submitted to the DCC representative for review and approval by the protocol PI. Any inconsistencies, omissions or quality concerns are resolved prior to any additional testing is performed.
- Project laboratory performance is monitored periodically throughout the course of the project and reports provided to the protocol PI.
A final review of test data is periodically performed during the project, and at the completion of all planned testing.

11.8. Genetic Studies: NIH Data Sharing Considerations

When combined with clinical and other phenotypic data, analysis of whole genome information offers the potential for increased understanding of basic biological processes affecting human health and improvement in the prediction of disease and treatment options. In the interest of moving this area of research forward, the NIH established expectations for sharing genetic data (e.g., SNP data) and phenotype data (e.g., information about disease status, outcomes and subject characteristics that are not individually identifiable) obtained through NIH-funded Genome-Wide Association Studies (GWAS). The NIH GWAS data sharing policy (“GWAS policy”) is the policy that GWAS data obtained with NIH support should be shared through a central repository when such data sharing is compatible with the consent provided by the participant. The NIH policy can be found at: (http://grants.nih.gov/grants/guide/notice-files/NOT-OD-07-088.html).

Many of the ongoing and recently opened BMT CTN studies, as well as future studies currently in development, have incorporated the collection and storage of research biospecimens. It is the desire of the BMT CTN (and NIH funding agencies) that these biospecimens should be found suitable for either protocol-defined laboratory studies or for unrestricted future research (including GWAS or whole exome sequencing (WES) studies) in conjunction with the clinical outcome data associated with the parent study. To that end, the BMT CTN has and will continue to include language in all protocol consents where biospecimens are being collected, so that we can facilitate the NIH data sharing compliance efforts and provide biospecimens that are available for the widest range of possible research opportunities.

The BMT CTN informed consent process and document will make it clear that participants’ DNA isolated from certain stored biospecimens (e.g., whole blood, PBMC, isolated DNA, marrow aspirates, stem cell product) and associated clinical data might be selected for use in a GWAS or WES study that is associated with a future NIH-funded or conducted study. The resulting de-identified genotype and phenotype data will be required to be shared for research purposes through the deposit of both the genotype and phenotype data in the access controlled public NIH Genotype and Phenotype database (dbGaP) managed by the National Center for Biotechnology Information (NCBI; http://www.ncbi.nlm.nih.gov/gap)
CHAPTER 12

SECONDARY DATA ANALYSES
12. SECONDARY DATA ANALYSES

12.1. Definition of Secondary Data Analyses

This chapter outlines procedures and policies for secondary data analyses using BMT CTN study data that may be accessed through the BMT CTN DCC during the “Proprietary Period”. The Proprietary Period is defined as a period of no more than 3 years after the end of the clinical activity (final patient follow-up, etc.) or 2 years after the primary results of the trial has been published, whichever comes first. Once these conditions have been met, the BMT CTN clinical trial data is transferred to the NHLBI for management as a public data resource for future studies. Data requests and proposals for secondary data analyses must then be directed to NHLBI through BioLINCC at https://biolincc.nhlbi.nih.gov/home/.

A “secondary data analysis” is defined as a study that uses already-collected clinical data from BMT CTN studies to look at objectives other than addressed in the original studies. This chapter will not address “ancillary studies” defined as analyses using additional data collected separately in the course of a BMT CTN study (addressed in Chapter 11), nor does it apply to studies using biologic material from BMT CTN studies to address secondary questions. This chapter also does not apply to secondary objectives already outlined in the protocol. Studies that use exclusively CIBMTR clinical data have their own mechanism for review and approval though the CIBMTR working committees. Thus, this chapter addresses research questions that use at least some BMT CTN data from one or more clinical trials.

12.2. Rationale for Conducting Secondary Analyses

The rationale for encouraging secondary analysis of BMT CTN data is to leverage the large amount of resources devoted to conducting the clinical trials into the greatest increase in knowledge. BMT CTN studies generally share a common set of case report forms (CRFs), a standardized process for reviewing key data elements through data review committees, and careful monitoring procedures. Almost all BMT CTN studies require completion of the CIBMTR full length comprehensive report forms, which can be linked to BMT CTN data through the CIBMTR CRID (a unique number given to each transplant recipient) to provide additional data for analysis. For studies that can address important hypotheses through the analysis of BMT CTN data, the efficiency of having the data already collected and available is a great strength. From a Network perspective, the ability to conduct high quality secondary analyses further leverages the NIH resources provided and shows the value of the Network.

12.3. Information to Help Plan Secondary Analyses and Determine Feasibility

Since investigators interested in conducting secondary data analyses may not be familiar with the specific BMT CTN trials or the Network itself, some general data to allow efficient understanding of the clinical trial specifics, study populations, and data available are posted on the public BMT CTN website to encourage secondary analyses: (BMT CTN secondary analyses resources), These resources are organized in a way to allow easy access and maximize utility.

1. BMT CTN protocols, and synopses for completed and open trials are available at the following BMT CTN webpage: BMT CTN Protocols.
2. BMT CTN CRFs for completed protocols will be available for review and will be labeled with the actual variable names to allow unambiguous communication about variables of interest. CRFs for completed BMT CTN protocols can be found at the following webpage: https://web.emmes.com/study/bmt2/casereportformspublic.html

3. Tables summarizing and comparing the baseline and clinical characteristics of the study populations associated with closed studies has been provided at the following BMT CTN webpage: https://web.emmes.com/study/bmt2/public/Secondaryanalysis.html.

4. A link to a listing of the primary manuscripts from completed studies can be found on the home page of the BMT CTN public website.

12.4. Submission and Adjudication of Secondary Clinical Data Analysis Study Proposals

To propose a secondary data analysis study, interested investigators will need to download and complete a BMT CTN Secondary Clinical Data Analysis Proposal Form that can be found on the public BMT CTN website (https://web.emmes.com/study/bmt2/public/Secondaryanalysis.html). The study proposal provides a thorough, yet concise description of the preliminary data and background, primary hypotheses, patient population, BMT CTN protocol number(s), necessary clinical information, funding or potential funding, and whether a BMT CTN or non-BMT CTN statistician will be performing the analysis. The completed study proposal is submitted to the designated BMT CTN DCC contacts, and is initially reviewed by by DCC staff and statisticians for completeness and non-overlap with any previously approved secondary data analyses or any planned secondary analyses in the primary study protocol. Following this initial review, the study proposal is provided to the parent study Protocol Team by the assigned BMT CTN statistician and together they adjudicate the proposal for feasibility and scientific merit. The study proposal, along with Protocol Team/BMT CTN statistician comments and recommendations are then provided to the BMT CTN Executive Committee for final review and an approval determination. If the parent Protocol Team is not active or the secondary analysis involves more than one protocol, then the assigned BMT CTN statistician will submit the proposal directly to the BMT CTN Executive committee to discuss and adjudicate the scientific merit and feasibility of the study. The investigator submitting the proposal may be invited to present the study to the Protocol Team and/or Executive Committee, or the assigned BMT CTN statistician may do this. Questions are discussed with the investigator and clarifications potentially made in the study plan in an iterative process throughout the course of the scientific review of the study proposal. Approval of the proposal will be documented on the Parent Study and BMT CTN Leadership Approval BMT CTN Biospecimen and Clinical Data Resources Form.

12.5. Funding for Secondary Data Analysis Studies

Funding for secondary data analysis studies may come from Network resources, an institution, private sources, or the NIH. If NIH or other grant funding is to be sought for an approved study, the BMT CTN will provide a letter of endorsement signed by the Executive or Steering Committee Chair or a DCC Leadership member.

The available funding or grant application budget for all Secondary Data Analysis studies must include support for the BMT CTN DCC to cover the development of the requested data file, the review of all data elements, the delivery and support related to the use of the dataset by the
investigators. Missing data may need to be addressed and potential supplemental data elements may also need to be considered. In order to perform those tasks related to the development, QA and delivery of study data sets, this time and effort needs to be reimbursed as part of the cost of conducting the study. Coverage for BMT CTN DCC costs will vary depending on the complexity of the Secondary Data Analysis study. Additionally, if the investigator has requested BMT CTN statistician collaboration to perform the study analysis the number of statistical hours needs to be included in this budget. The DCC will provide an approved budget for DCC direct and indirect costs for all approved studies and any other sub-contractor forms required by a prospective funder.

Any funding received from government agencies or third party funders, outside of the primary BMT CTN grants, shall be reported to the BMT CTN for inclusion in the tally of total contributions to the BMT CTN.

12.6. Conducting Approved Secondary Data Analysis Studies

Upon approval of the Secondary Analysis Study, a BMT CTN Research Materials Distribution Agreement (RMDA) for the BMT CTN DCC must be executed prior to receiving data from BMT CTN DCC. This agreement shall be signed by the National Marrow Donor Program on behalf of the BMT CTN, the Principal Investigator and the Principal Investigator’s Institution. Any transfer of de-identified data from the BMT CTN to a local institution for analysis, the institution will abide by all rules established in the data agreement. Local IRB approval for data receipt may be required by the investigator’s institution. IRB approval is not required by BMT CTN, but documentation of IRB approval or a waiver is required before the data are released. If the BMT CTN DCC will be analyzing the data, and no data will be released to the Principal Investigator or Principal investigator’s Institution, the RMDA is not required.

The BMT CTN strongly recommends that approved study principal investigators consider the inclusion of a BMT CTN statistician on the study team, as they are already familiar with the data being provided for the study. Other members to consider including on the study team may include the PI or other key members of the parent Protocol Team. Their intimate knowledge of the primary protocol, the content and quality of the clinical data and their previous work with the analysis of study outcomes and endpoints would likely be of considerable value to the study team.

If a non-BMT CTN statistician is conducting the analysis then involvement of a BMT CTN statistician in a consulting role is highly recommended, since BMT CTN will need to review the analysis prior to presentation or publication.

The secondary analysis Study Team creates a timeline for completion of the secondary analysis and ensure the timeline to be met. This timeline must be agreed to by the BMT CTN statistician creating the dataset and the team conducting the analysis, since there are several key milestones that need to be considered over the course of the study.

12.7. Publications Guidelines

Please see Section 8 which addresses BMT CTN publication policies. Please note that the BMT CTN Publications Committee and NHLBI/NCI program officers must review all manuscripts before submission. See section 8.4.3 for guidelines regarding how to reference NIH funding support for the parent study(ies).