Protocol for a Research Sample Repository for Hematopoietic Cell Transplantation, Other Cellular Therapies and Marrow Toxic Injuries (IRB-1991-0002)

Version 13.0 (Amendment) – June 9, 2021

Consent Form Revisions – July 13, 2020 (Effective 7/27/2020)

Version 12.0 (Status Review) - June 14, 2019

Version 11.0 (Continuing) - July 30, 2018

Version 10.0 (Continuing) – July 30, 2017

Version 9.0 (Continuing) – July 30, 2016

Version 8.2 (Continuing) – July 30, 2015

Consent Form Revisions – February 24, 2015

Version 8.1 (Continuing) – July 30, 2014

Version 8.0 (Continuing) - July 30, 2013

Version 7.0 (Continuing) – July 30, 2012

Version 6.0 (Continuing) - July 30, 2011

Version 6.0 (Continuing) - July 30, 2010

Version 5.1 (Continuing) – July 30, 2009

Version 5.0 (Continuing) - July 30, 2008

Version 4.1 (Continuing) – July 30, 2007

Version 4.0 (Amendment) – April 26, 2007 (Effective 6/11/07)

Version 3.0 (Continuing) – July 30, 2006

Version 2.5 (Continuing) – July 30, 2005

Version 2.4 (Amendment) - May 20, 2005

Version 2.3 (Amendment) - May 1, 2005

Version 2.2 (Continuing) – July 30, 2004

Version 2.1 (Amendment) – January 28, 2004 Version 2.0 (Continuing) – October 1, 2003

Version 1.0 – July 2002

Description of Revision	Document/Section(s) Affected	Effective Date
Added "ClinicalTrials.gov" identifier	Protocol: Title page	6/11/2021
1st bullet: "Studies of histocompatibility including HLA typing, analysis of allele and haplotype frequencies in different populations, evaluation of microsatellites and single nucleotide polymorphisms, examination of minor histocompatibility histocompatibility antigens"	Protocol: section 8.1	6/9/21
1st paragraph: "Up to <u>fifty</u> thirty milliliters (<u>50</u> 30 mL) of blood <u>can be</u> are collected from adult. For pediatric patients where the collection of the full sample is medically contraindicated, <u>adjusted volumes will be outlined in study specific documents</u> . as little as one milliliter (1 mL) of blood may be collected."	Protocol: section 4.3	6/9/21
2 nd paragraph, 3 rd & 4 th sentences: " <u>Sample collection</u> specifications will be outlined in study specific documents. These study project specific consent forms"	Protocol: section 4.2	6/9/21

1st paragraph: "Up to <u>fifty</u> thirty milliliters (<u>50</u> 30 mL) of	Protocol: section 4.2	6/9/21
blood <u>can be</u> are collected from adult patients. For pediatric		
patients where the collection of the full sample is medically		
contraindicated, <u>adjusted volumes will be outlined in study</u>		
specific documents. as little as one milliliter (1 mL) of blood		
may be collected. Previously collected tissue samples may be		
requested for events including diagnosis of primary disease		
and or a secondary primary malignancy as well as other		
events listed above."		
Section 4.2 heading: "Post-Transplant or Cellular Therapy	Protocol: section 4.2	6/9/21
Collection of Patient Blood and Tissue Samples"		
3 rd sentence: "full sample is medically contraindicated,	Protocol: section 4.1.2	6/9/21
adjusted volumes will be outlined in study specific	110tocoi. Section 4.1.2	0/ // 21
documents. as little as one milliliter (1 mL) of blood may be		
collected."		
2 nd sentence: "Up to thirty fifty milliliters (30 50 mL) of	Protocol: section 4.1.2	6/9/21
blood can be are collected from adult patients."	11000con section 111.2	0/ 5/ 21
2 nd paragraph, 2 nd sentence: "Up to thirty fifty milliliters (30)	Protocol: section 4.1.1	6/9/21
50 mL) of blood can be are collected from adult donors."	Frotocoi: section 4.1.1	0/9/21
	D (1 (111	6/0/01
2 nd paragraph, 1 st sentence: " Unrelated d Donor samples	Protocol: section 4.1.1	6/9/21
are"	Desta all modium 4.1.1	C/0/21
1st paragraph: "full sample is medically contraindicated,	Protocol: section 4.1.1	6/9/21
adjusted volumes will be outlined in study specific		
documents as little as one milliliter (1 mL) of blood may be		
eollected. Previously collected blood or tissue diagnostic		
samples may be requested for evaluation of primary disease		
for specific studies. Sample collection specifications will be		
outlined in study specific documents each consent specific to		
the research study."		
1 st paragraph, 2 nd sentence: "Up to thirty fifty milliliters (30)	Protocol: section 4.1.1	6/9/21
50 mL) of blood can be are collected"		
"Blood samples should be collected as indicated on the	Protocol: section 4.1	6/9/21
Research Repository Critical Facts Sheet and/or study		
specific consents."		
Section 4.1 heading: "Pre-Transplant or Cellular Therapy	Protocol: section 4.1	6/9/21
Collection of Donor and Patient Blood and Tissue Samples"		
"All U.S. recipients of allogeneic or autologous"	Protocol: section 2.3	6/9/21
Updated Table of Contents to match body of the protocol.	Protocol: Page 2 Table of	6/9/21
epatice ruble of contents to materi body of the protocol.	Contents	0/ // 21
Updated formatting/margins throughout.	Protocol: Throughout	6/9/21
Changed version # to 13.0, 2021 copywrite and revision 19	Protocol: Footer	6/9/21
Changed version # to 13.0	Protocol: Title page	6/9/21
	Record of Revisions: Title Page	V. 21 = 1
Updated Principal Investigator Title: "CIBMTR Senior	Protocol: Title page	6/9/21
Scientific Director, Vice President, Research"	110000000 Time page	V. 21 = 1
New Assent Form: Minor Recipient Assent Form (7 to 11		3/1/21
years of age) SARS-CoV-2 Vaccine Response Study		=, =, = =
New Assent Form: Minor Recipient Assent Form (12 to 17		3/1/21
years of age) SARS-CoV-2 Vaccine Response Study		5, 1, 21
New Consent Form: Adult Research Consent Form and		3/1/21
Parent /Legal Guardian Permission Form Allogeneic or		2, 1, 21
Taront / Dogar Guardian Formission Form / mogenitie of		

Autologous Recipient SARS-CoV-2 Vaccine Response		
Study		
Removal of the Donor ID field from the footer	Donor Adult/Parent Consent Form	2/27/21
Removal of the Donor ID field from the footer	Donor Match Algorithm Consent Form	12/1/20
Section 5: Paragraph 2 [insert <u>center name-location</u> here] Section 5: Added "This research is covered by a Certificate of Confidentiality from the Health Resources and Services Administration (HRSA)." to paragraph 5. Section 6: Cost and Reimbursement language expanded	Consent Forms: Donor Adult/Parent Consent Form; Donor Match Algorithm Consent Form; Recipient Marrow Toxic Injury Adult/Parent Consent Form; Recipient Adult/Parent Consent Form; Recipient Secondary Primary Malignancy Adult/Parent Consent Form.	7/27/20
Replaced "brother or sister" with "family member" through the assent forms.	Assent Forms: Minor Related Donor Assent Form 7-11 and 12- 17	7/27/20
Updated formatting of the page number in the footer)	Assent Forms: Minor Recipient Assent (12-17) and Minor Marrow Toxic Injury Assent Form (12-17)	7/27/20
Re-formatting of consent forms	All consent forms	6/14/19
Merged separate Adult and Parent/Legal Guardian consent forms into one Adult Parent/Legal Guardian consent form	Consent Forms: Adult Research Consent Form and Parent/Legal Guardian Permission Form Allogeneic or Autologous Recipient; Adult Research Consent Form and Parent/Legal Guardian Permission Form Allogeneic Donor; Adult Research Consent Form and Parent/Legal Guardian Permission Form Marrow Toxic Injury Patient	6/14/19
2 nd paragraph, 2 nd sentence: "name and the <u>alpha</u> numeric code."	Protocol: section 10.1	6/14/19
2 nd paragraph, 1st sentence: "only with a <u>n alpha</u> numeric code"	Protocol: section 10.1	6/14/19
1st paragraph, 3rd sentence: "If the subject's sample has already been given to an investigator at the time he/she asked to withdraw from the Research Repository, the investigator will be instructed to remove the subject's sample from their study set. Any unused sample will be destroyed."	Protocol: section 9.0	6/14/19
1st paragraph, 1st sentence: "samples obtained from the NMDP Research Sample Repository"	Protocol: 10.3	6/14/19
Added 5 th paragraph: "Additionally, NMDP and MCW maintain appropriate technical and organizational measures for the adequate protection of the security and privacy of its systems and data. These protections comply with the United States National Institute of Standards and Technology. Security Controls for Federal Information Systems (NIST 800-53), and all other applicable security and data privacy	Protocol: section10.1	6/14/19

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requirements. These safeguards are audited annually by a		
qualified independent auditor; results are reported to		
CIBMTR management for timely resolution."		
Added 4 th paragraph: "All research staff at the CIBMTR and	Protocol: section 10.1	6/14/19
the NMDP maintain up-to-date training in protection of		
human subjects. This training is received through the		
Collaborative IRB Training Initiative (CITI) program. This		
is a web-based training program offered through the		
Biomedical Research Alliance of New York (BRANY)."		
Added 3 rd paragraph: "Access to all information in the	Protocol: section 10.1	6/14/19
Research Sample Repository is tightly controlled with		
passwords and logins at multiple levels. Access to the		
Research Sample Repository is limited to those employees		
who have specific job responsibilities related to the		
repository."		
1st bullet point: "All sample data is stripped of identifying	Protocol: section 8.6	6/14/19
sequences, e.g. Y chromosome, mitochondrial DNA, or other		
unique sequences, dates and detailed demographic data prior		
to submission to dbGAP."		
1st paragraph, 1st bullet point: "NMDP/CIBMTR	Protocol: section 8.4	6/14/19
$\pm \underline{R}$ esearch $\pm \underline{D}$ atabase, subject to provisions of the $\pm \underline{R}$ esearch		
dDatabase protocol"		
Removed section 4.1.2 Related Donor Transplants or	Protocol: section 4.1.2	6/14/19
Cellular Therapies		3, 2 3, 2 3
7 th bullet point: "In cases where the center has designated is	Protocol: section 3.1	6/14/19
relying on the NMDP IRB for this process their IRB of		3, 2 3, 2 3
record for the Research Repository protocol, and an IRB		
Authorization Agreement is in place, the center"		
6 th bullet point, 1 st sentence: "review period <u>if the center</u>	Protocol: section 3.1	6/14/19
has not transitioned the protocol to the 2018 Common Rule	21000000 5000000 511	0/1 1/19
requirements. If the center transitioned the protocol to the		
2018 Common Rule requirements, then the above process is		
only followed when there are amendments to the protocol or		
consent forms."		
4 th bullet point: "submitted to the NMDP IRB office at the	Protocol: section 3.1	6/14/19
NMDP."	110tocon section 3.1	0/11/19
3 rd bullet point: "the local IRB need to must be	Protocol: section 3.1	6/14/19
reviewed"	110tocon section 3.1	0/11/19
1st bullet point: "The NMDP's template protocol"	Protocol: section 3.1	6/14/19
Removed 1st sentence: "The process for local and NMDP	Protocol: section 3.1	6/14/19
IRB approval are as follows."	1 Totocol: Section 3.1	0/14/17
Added 3 rd paragraph: "International centers must follow their	Protocol: section 3.0	6/14/19
own national regulations and provide assurance to the	1 Totocol. Section 3.0	0/14/19
CIBMTR that national regulations are being followed."		
2nd paragraph: "Local IRB review and approval is necessary	Protocol: section 3.0	6/14/19
except in the case of centers that designate the NMDP IRB as	1 10tocor. Section 5.0	0/14/19
their IRB. Centers must use the protocol and consent forms		
provided by the NMDP and submit them to their designated		
IRBs for review and approval. This protocol and its		
associated consent forms are provided to centers on the		
CIBMTR website, www.cibmtr.org."	Durata asla angli ang 2.0	C/14/10
1 st paragraph, added 2 nd sentence: " <u>The center may obtain</u>	Protocol: section 3.0	6/14/19

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IRB approval either through their local IRB or delegate		
review to the NMDP IRB through an IRB Authorization		
Agreement."		
1 st paragraph, 1 st sentence: "centers must have obtain IRB-	Protocol: section 3.0	6/14/19
approvedal for the protocol"		
1st bullet point added: "as set forth in 45 CRF 46.408."	Protocol: section 2.5.1	6/14/19
Added 1st paragraph: "The Research Sample Repository	Protocol: section 2.5.1	6/14/19
included pediatric patients and related donors. The		
<u>procedural risk involved in this protocol meets the definition</u>		
of minimal risk set forth in 45 CFR 46.106 (i) "Minimal risk		
means that the probability and magnitude of harm or		
discomfort anticipated in the research are not greater in and		
of themselves than those ordinarily encountered in daily life		
or during the performance of routine physical or		
psychological examinations or tests." Participation on this		
protocol requires a routine blood draw."		
Added 2 nd paragraph: "Non-U.S. centers contributing	Protocol: section 2.5	6/14/19
samples to the Research Sample Repository will provide		
written assurance that the submission of samples has on-		
going oversight by their local Ethics Review Board/Medical		
Ethics Committee and all regulations are followed."		
1 st paragraph, added 3 rd and 4 th sentence: " <u>To confirm that</u>	Protocol: section 2.5	6/14/19
participants have given consent to participate in the Research		
Sample Repository, the first form submitted to the Research		
Database on a participant includes confirmation that the		
participant signed the informed consent document.		
Institutional IRB policies must be followed regarding re-		
consent of minor patients when those patients reach the age		
of majority."		
1st paragraph, 2 nd sentence: "Documentation of assent and of	Protocol: section 2.5	6/14/19
parent legal guardian permission of minor participants, and		
consent for adult participants, must be maintained at the		
center where the participant or their parent or legal guardian		
provided consent to participate. The center where consent is		
obtained is responsible for maintaining the written consent		
form and documentation of the mirror assent decision."		
1 st paragraph, added new 1 st sentence: " <u>In the event of a</u>	Protocol: section 2.4	6/14/19
radiation exposure accident, the NMDP has a radiation injury		
treatment network, whose purpose is to collect data to		
understand the outcomes of patients treated under these		
circumstances."		
2 nd paragraph: "All donors registered on the NMDP Registry,	Protocol: section 2.1	6/14/19
regardless of whether they have been requested to donate a		
product for a patient, are eligible to participate in the		
Research Sample Repository. Registered volunteer donors		
are eligible to participate in the NMDP Research Sample		
Repository if selected for further tissue type characterization		
in research projects designed to enhance the NMDP search		
(match) algorithm. Search algorithm enhancement research		
will entail potential donor match identification, verification		
of donor tissue type (including rare tissue types) and analyses		
to further characterize factors that impact histocompatibility.		
A tissue type is considered rare if it has been found in less		

than one in a million donors on the NMDP registry. Donors		
with rare tissue types will be targeted based on HLA typing		
results reported to the NMDP.	Protocol: section 2.1 & 2.3	C/14/10
1 st paragraph, added last sentence: " <u>This includes adults with</u>	Protocol: section 2.1 & 2.3	6/14/19
and without decision making capacity and children."	D 4 1 2	6/14/10
3 rd paragraph, 1 st bullet point: " <u>Investigate molecular</u>	Protocol: section 1.3	6/14/19
explanations for histocompatibility or clinical outcome		
revealed through analysis of genomic, epigenetic, or other		
biomolecular data; Improve the understanding of tissue		
matching for HSC or cellular therapy donor and recipients		
3 rd paragraph, 1 st sentence: "of studies that the NMDP	Protocol: section 1.3	6/14/19
allows the samples to may be used"		
2 nd paragraph, 1 st sentence: "The NMDP Research	Protocol: section 1.3	6/14/19
Sample"		
1 st paragraph, 3 rd sentence: "donated or received a n	Protocol: section 1.3	6/14/19
allogeneic or autologous HSC transplant"		
1 st paragraph, 1 st sentence: "NMDP established the NMDP	Protocol: section 1.3	6/14/19
Research Sample Repository which is currently located at the		
NMDP Repository Biorepository services"		
Section 1.2: "The International Blood Marrow Transplant	Protocol: section 1.2	6/14/19
Registry (IBMTR), located with the Department of Medicine		
of the Medical College of Wisconsin, was established in		
1972 to monitor and study outcomes of bone marrow		
transplants. In 2004 the NMDP and IBMTR established the		
Center of Blood and Marrow Transplant Research		
(CIBMTR). The CIBMTR is a research affiliation between		
the NMDP and the Medical College of Wisconsin. The		
CIBMTR has both a Minneapolis campus located within the		
NMDP offices and a Milwaukee campus at the Medical		
College of Wisconsin. The NMDP Research Program is		
accomplished through the CIBMTR."		
Added new section 1.2: "Center for International Blood and	Protocol: section 1.2	6/14/19
Marrow Transplant Research®"	110tocon section 1.2	0/11/19
Added second paragraph: "In addition, the Federal contract	Protocol: section 1.1	6/14/19
also recognized that the NMDP could play a critical role in	Trotocon section 1.1	0/11/19
responding to contingency events; primarily radiation and		
chemical exposures occurring either accidentally or resulting		
from military or terrorist actions that cause a marrow toxic		
injury."		
1 st paragraph, 2nd sentence: "matching related donor. As	Protocol: section 1.1	6/14/19
part of the Federal contract the NMDP was required to	1 10tocor. Section 1.1	U/ 14/ 17
collect outcomes data and research samples on patients who		
received a product through NMDP."		
	Ducto colo Thursush sut	C/14/10
Changes references from "NMDP IRB Chair" to "Research	Protocol: Throughout	6/14/19
Repository Principal Investigator"	D 4 1 771 1 4	6/14/10
Changes references from "NMDP Research Sample	Protocol: Throughout	6/14/19
Repository" to "Research Sample Repository"	D 4 1 771	214 4 14 0
Changed references from "HSC" to "HC"	Protocol: Throughout	6/14/19
Changed references from "Recipient" to "Patient"	Protocol: Throughout	6/14/19
Section 10.1 heading: "Coded Sample Inventory and, Links	Protocol: section 10.1	6/14/19
to Personal Identifiers and Staff Training and Access"		
Section 6 heading: "Duration of Sample Storage at the	Protocol: section 6	6/14/19

NMDP Research Sample Repository"		
Added Section 3 Sub-title: 3.1 IRB Approval Process	Protocol: section 3.0	6/14/19
Updated section header 4.1.3 Autologous Transplants or	Protocol: section 4.1.3	6/14/19
Cellular Therapies to 4.1.2	1 Totocoi. Section 4.1.5	0/14/19
Section 4.1.1 heading: "Unrelated and Related Donor	Protocol: section 4.1.1	6/14/19
Transplant or Cellular Therapies"	Trotocon section	0,11,15
Section 2.4 heading: "Individuals Patients With Marrow	Protocol: section 2.4	6/14/19
Toxic Injury Eligibility Criteria"		
Section 2.3 heading: "Hematopoietic Stem Cell or Other	Protocol: section 2.3	6/14/19
Cellular Therapy Recipients Eligibility Criteria"		
Section 2.2 heading: "Cord Blood Units Eligibility Criteria"	Protocol: section 2.2	6/14/19
Section 2.1 heading: "Hematopoietic Stem Cell or Other	Protocol: section 2.1	6/14/19
Cellular Therapy Recipients Eligibility Criteria"		
Section 2 heading: "Eligibility to Participate in the NMDP	Protocol: section 2	6/14/19
Research Sample Repository"		
Section 1.3 heading: "Establishment and Purpose of the	Protocol: section 1.3	6/14/19
NMDP Research Sample Repository"		
Added new section 1.2: "Center for International Blood and	Protocol: section 1.2	6/14/19
Marrow Transplant Research®"		
Section 1.2 renamed section 1.3	Protocol: section 1.2	6/14/19
Updated Table of Contents to match body of the protocol.	Protocol: Page 2 Table of	6/14/19
	Contents	
Changed version # to 12.0 and version date to TBD	Protocol: Title page	6/14/19
	Record of Revisions : Title Page	
Added Title page: "Stephen Spellman, M.S. <u>CIBMTR</u>	Protocol: Title page	6/14/19
Assistant Scientific Director		
Director, Immunobiology Research"		
Removed "Allogeneic" and "Stem" from title of protocol	Protocol: Title Page	6/14/19
	Record of Revisions: Title Page	
Changed footer: GRID (if applicable) line added	Donor Consent Form:	4/29/2019
N. A. (F. M. D.); (A. (F. (7), 11)	Footer	04/04/2010
New Assent Form: Minor Recipient Assent Form (7 to 11		04/24/2019
years of age) Secondary Primary Malignancy		04/04/2010
New Assent Form: Minor Recipient Assent Form (12 to 17		04/24/2019
years of age) Secondary Primary Malignancy New Consent Form: Adult Research Consent Form and		04/24/2019
		04/24/2019
Parent /Legal Guardian Permission Form Allogeneic or Autologous Recipient Secondary Primary Malignancy		
Changed title: "Minor Allogeneic Recipient Assent Form"	Assent Forms:	07/30/2018
Changed title. Willof Anogenete Recipient Assent Form	Recipient Assent 7-11;	07/30/2018
	Recipient Assent 7-11, Recipient Assent 12-17	
3 rd paragraph, added last sentence: "You may be asked to	Assent Forms:	07/30/2018
give another blood sample in the future if something happens	Recipient Assent 12-17	07/30/2010
like your disease comes back, but you don't have to give any	recipioni rissent 12 17	
future blood samples if you don't want to."		
2 nd paragraph, added last sentence: "You might be asked to	Assent Forms:	07/30/2018
give another blood sample in the future, but you don't have	Recipient Assent 7-11	3773072010
to give another one if you don't want to."	F, 22	
Changed title: "Minor Allogeneic Recipient Parent/Legal	Recipient Consent Forms	07/30/2018
Guardian Research Permission Form"	(Title):	
	Minor Recipient Parent/Legal	
	Guardian	
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1st paragraph, added last two sentences: "You may also be asked to contribute a blood sample in the future after a specific clinical event (disease relapse, development of a new cancer or other event). Your participation in this protocol does not obligate you to provide future samples." After 1st paragraph, added:	Recipient Consent Forms (Section II): Adult Recipient; Minor Recipient Parent/Legal Guardian; Recipient Consent Forms	07/30/2018
"This research is covered by a Certificate of Confidentiality from the Federal Government. The researchers with this Certificate may not disclose or use information, documents, or biospecimens that may identify you in any federal, state, or local civil, criminal, administrative, legislative, or other action, suit, or proceeding, or be used as evidence, for example, if there is a court subpoena, unless you have consented for this use. Information, documents, or biospecimens protected by this Certificate cannot be disclosed to anyone else who is not connected with the research except, if there is a federal, state, or local law that requires disclosure (such as to report child abuse or communicable diseases but not for federal, state, or local civil, criminal, administrative, legislative, or other proceedings, see below); if you have consented to the disclosure, including for your medical treatment; or if it is used for other scientific research, as allowed by federal regulations protecting research subjects. The Certificate cannot be used to refuse a request for information from personnel of the United States federal or state government agency sponsoring the project that is needed for auditing or program evaluation by Health Resources and Services Administration (HRSA) which is funding this project or for information that must be disclosed in order to meet the requirements of the federal Food and Drug Administration (FDA). You should understand that a Certificate of Confidentiality does not prevent you from voluntarily releasing information about yourself or your involvement in this research. If you want your research information released to an insurer, medical care provider, or any other person not connected with the research, you must provide consent to allow the researchers to release it."	(Section IV): Adult Recipient; Minor Recipient Parent/Legal Guardian; Marrow Toxic Injury Consent Forms (Section IV): Adult Marrow Toxic Injury; Minor Marrow Toxic Injury Parent/Legal Guardian Donor Consent Forms (Section IV) Adult Allogeneic Donor; Minor Related Donor Parent/Legal Guardian; Adult Registered Donor for Match Algorithm Enhancement;	
1st paragraph: "The Center for International Blood and Marrow Transplant Research (CIBMTR), the a research program collaboration of the National Marrow Donor Program (NMDP)/Be The Match and the Medical College of Wisconsin, invites you to take part in the Research"	Recipient Consent Forms (Section I): Adult Recipient; Minor Recipient Parent/Legal Guardian; Marrow Toxic Injury Consent Forms (Section I): Adult Marrow Toxic Injury; Minor Marrow Toxic Injury Parent/Legal Guardian	07/30/2018

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	Donor Consent Forms (Section I) Adult Allogeneic Donor; Minor Related Donor Parent/Legal Guardian; Adult Registered Donor for Match Algorithm Enhancement;	
Added entire section 10.2: "Certificate of Confidentiality"	Protocol: Section 10.2	07/30/2018
Added entire section 4.2: "Post-transplant or Cellular	Protocol: Section 4.2	07/30/2018
Therapy Collection of Recipient Blood Samples"		
Added entire section 4.1.3: "Autologous Transplants or	Protocol: Section 4.1.3	07/30/2018
Cellular Therapies"		
1st paragraph, 1st sentence: "Recipient pre-transplant or	Protocol: Section 4.1.2	07/30/2018
<u>cellular therapy</u> samples are collected prior to the"		
1 st paragraph, 1 st sentence: "Recipient <u>pre-transplant or</u> cellular therapy samples are collected prior to the"	Protocol: Section 4.1.1	07/30/2018
Section 4.1 heading: "Pre-Transplant or Cellular Therapy	Protocol: Section 4.1	07/30/2018
Collection of Donor and Recipient Blood Samples"	21000001 Soundar III	0,7,00,2010
1 st paragraph: "All U.S. recipients of allogeneic or	Protocol: Section 2.3	07/30/2018
autologous HSC transplants or cellular therapies"		
Section 2.3 heading: "Hematopoietic Stem Cell	Protocol: Section 2.3	07/30/2018
<u>Transplantation</u> or Other Cellular Therapy Recipients"		
1st paragraph, 3rd sentence: "Blood samples are donated by	Protocol: Section 1.2	07/30/2018
donors, CBUs and recipients who have registered, donated or		
received an allogeneic <u>or autologous</u> HSC transplant or other		
cellular therapy"		05/20/2010
Updated Table of Contents to match body of the protocol.	Protocol: Page 2 Table of	07/30/2018
Classical and a 11.0 and a 12.0 and 12.	Contents	07/20/2019
Changed version # to 11.0 and version date to July 30, 2018	Protocol: Title page	07/30/2018 01/24/2018
Added to end of paragraph: "You and your child will not be paid for taking part in the Research Sample Repository. It	Recipient Consent Forms (Section V):	01/24/2018
will not cost you anything for your child to participate in the	Minor Allogeneic Recipient	
repository. Research samples may be used for commercial	Parent/Legal Guardian;	
projects and profit. If your child's sample is used for	Tarena Begar Guardian,	
development of a commercial product, neither you nor your	Marrow Toxic Injury Consent	
child will share in any profit."	Forms (Section V):	
	Minor Marrow Toxic Injury	
	Parent/Legal Guardian	
	Donor Consent Forms (Section V)	
	Minor Related Donor	
	Parent/Legal Guardian;	
	racin Legai Guardian,	
Added after 2 nd paragraph: "To expand research, it is helpful	Recipient Consent Forms	01/24/2018
for researchers to share information they get from studying	(Section III):	
research samples. They do this by putting the information	Minor Allogeneic Recipient	
into one or more scientific databases, where it is stored along	Parent/Legal Guardian;	
with information from other studies. Researchers can then		
study the combined information to learn even more about	Marrow Toxic Injury Consent	
<u>health and disease.</u>	Forms (Section III):	

Protocol for a Research Sample Repository for Hematopoietic Cell Transplantation, Other Cellular Therapies and Marrow Toxic Injuries (IRB-1991-0002)

If you agree to allow your child to take part in this study, some of your child's genetic and health information may be placed into a scientific database, such as "dbGAP" that is maintained by the National Institutes of Health. A researcher who wants to study the information must apply and be approved to use the database. The CIBMTR restricts the use of the data to studies related to transplant and cellular therapy.

Data submitted to databases, such as dbGAP, may be used in additional research including genomic studies. Researchers with an approved study may be able to see and use your child's information pooled with information from many other individuals, but your child's name and other information that could directly identify your child will never be placed into a scientific database.

Since your child's genetic data is unique to them, there is a chance that someone could trace it back to them. The risk is small, but may grow in the future if you or your child discloses genetic information linked to their identity.

Researchers accessing your child's information will always have a duty to protect their privacy and to keep their information confidential.

You may be concerned that someone could get access to your child's genetic information and that it could be misused.

There are laws in place that make it illegal for an employer or health insurance company to discriminate against an individual based on their genetic information. Further, your child's privacy and the confidentiality of your child's data are very important to CIBMTR, and every effort will be made to protect them.

You or your child will not directly receive any research results or data generated from your child's sample."

Added 2nd paragraph: "The DNA testing may include "whole genome sequencing". Every cell in your child's body contains the genetic code for their DNA. Whole genome sequencing looks at the entire genome, or genetic code. All people have about 99.6% identical genomes. However, everyone is unique, and between any two people there could be about 24 million places where the "spelling" of the code is different. Associating these differences in spelling (gene variants) with differences in transplant or cellular therapy outcomes may help us to understand how these variants are related to disease and treatment success."

Minor Marrow Toxic Injury Parent/Legal Guardian

Donor Consent Forms (Section III)

Minor Related Donor Parent/Legal Guardian;

Recipient Consent Forms (Section II):

Minor Allogeneic Recipient Parent/Legal Guardian;

Marrow Toxic Injury Consent Forms (Section II):

Minor Marrow Toxic Injury Parent/Legal Guardian

Donor Consent Forms (Section II)

Minor Related Donor Parent/Legal Guardian;

01/24/2018

Added to end of paragraph: "You will not be paid for taking part in the Research Sample Repository. It will not cost you anything to participate in the repository. Research samples may be used for commercial projects and profit. If your sample is used for development of a commercial product, you will not share in any profit."	Recipient Consent Forms (Section V): Adult Allogeneic Recipient; Marrow Toxic Injury Consent Forms (Section V): Adult Marrow Toxic Injury; Donor Consent Forms (Section V) Adult Allogeneic Donor; Adult Registered Donor for Match Algorithm Enhancement;	01/24/2018
Added after 2nd paragraph: "To expand research, it is helpful for researchers to share information they get from studying research samples. They do this by putting the information into one or more scientific databases, where it is stored along with information from other studies. Researchers can then study the combined information to learn even more about health and disease. If you agree to take part in this study, some of your genetic and health information may be placed into a scientific database, such as "dbGAP" that is maintained by the National Institutes of Health. A researcher who wants to study the information must apply and be approved to use the database. The CIBMTR restricts the use of the data to studies related to transplant and cellular therapy. Data submitted to databases, such as dbGAP, may be used in additional research including genomic studies. Researchers with an approved study may be able to see and use your information pooled with information from many other individuals, but your name and other information that could directly identify you will never be placed into a scientific database. Since your genetic data is unique to you, there is a chance that someone could trace it back to you. The risk is small, but may grow in the future if you disclose your genetic information will always have a duty to protect your privacy and to keep your information confidential. You may be concerned that someone could get access to your genetic information and that it could be misused. There are laws in place that make it illegal for an employer or health insurance company to discriminate against an individual based on their genetic information. Further, your privacy and the confidentiality of your data are very important to CIBMTR, and every effort will be made to protect them.	Recipient Consent Forms (Section III): Adult Allogeneic Recipient; Marrow Toxic Injury Consent Forms (Section III): Adult Marrow Toxic Injury; Donor Consent Forms (Section III) Adult Allogeneic Donor; Adult Registered Donor for Match Algorithm Enhancement;	01/24/2018

You will not directly receive any research results or data		
generated from your sample."		
Added 2 nd paragraph: "The DNA testing may include "whole genome sequencing". Every cell in your body contains the genetic code for your DNA. Whole genome sequencing looks at the entire genome, or genetic code. All people have about 99.6% identical genomes. However, everyone is unique, and between any two people there could be about 24 million places where the "spelling" of the code is different. Associating these differences in spelling (gene variants) with differences in transplant or cellular therapy outcomes may help us to understand how these variants are related to disease and treatment success."	Recipient Consent Forms (Section II): Adult Allogeneic Recipient; Marrow Toxic Injury Consent Forms (Section II): Adult Marrow Toxic Injury; Donor Consent Forms (Section II) Adult Allogeneic Donor; Adult Registered Donor for Match Algorithm Enhancement;	01/24/2018
Added 4 th bullet: "Study the success of transplantation, cellular therapies, or supportive care in the management of marrow toxic injuries."	Protocol: Section 1.2	07/30/2017
Added 7th bullet: "Studies of the success of transplantation, cellular therapies or supportive care in the management of marrow toxic injuries."	Protocol: Section 8.1	07/30/2017
Changed study title to: Protocol for a Research Sample Repository for Allogeneic Hematopoietic Stem Cell Transplantation, Other Cellular Therapies and Marrow Toxic Injuries	Protocol: Title page	07/30/2016
Changed NMDP's address	Protocol: Title page	07/30/2016
Added "cellular therapy" appropriately throughout the protocol	Protocol: throughout	07/30/2016
Changed references from "transplant center" to "treatment center" throughout the protocol	Protocol: throughout	07/30/2016
Paragraph 1, last sentence: "outcomes of unrelated <u>and</u> related donor HCT <u>and cellular therapies:"</u>	Protocol: Section 8.1	07/30/2016
Paragraph 2, 1st sentence: "The CIBMTR is trying to learn more about what makes bone marrow, blood stem cell and cord blood transplants and cellular therapies work well."	Donor Consent Forms (Section I) Adult Allogeneic Donor; Adult Registered Donor for Match Algorithm Enhancement; Minor Related Donor Parent/Legal Guardian; Recipient Consent Forms (Section I): Adult Allogeneic Recipient; Minor Allogeneic Recipient Parent/Legal Guardian;	07/30/2016
Paragraph 2, 2 nd sentence: "Although the exact studies for which Research Repository samples may be used is are not known at this time"	Donor Consent Forms (Section I) Adult Allogeneic Donor; Adult Registered Donor for Match Algorithm Enhancement; Minor Related Donor	07/30/2016

	Parent/Legal Guardian;	
	Recipient Consent Forms	
	(Section I):	
	Adult Allogeneic Recipient;	
	Minor Allogeneic Recipient	
	Parent/Legal Guardian;	
	Marrow Toxic Injury Forms	
	(Section I):	
	Adult Marrow Toxic Injury;	
	Minor Marrow Toxic Injury	
	Parent/Legal Guardian	
Paragraph 2, 2 nd bullet: "Determine and evaluate the factors	Donor Consent Forms (Section	07/30/2016
that affect transplant and cellular therapy outcome;"	I)	
	Adult Allogeneic Donor;	
	Minor Related Donor	
	Parent/Legal Guardian;	
	Recipient Consent Forms	
	(Section I):	
	Adult Allogeneic Recipient;	
	Minor Allogeneic Recipient	
	Parent/Legal Guardian;	
Paragraph 1, 2 nd sentence: "The blood will be collected	Donor Consent Forms (Section	07/30/2016
either just before your bone marrow, blood stem cell, or	II)	
<u>cellular therapy</u> donation"	Adult Allogeneic Donor;	
	Minor Related Donor	
	Parent/Legal Guardian;	
Paragraph 1, 2 nd sentence: "The blood will be collected just	Recipient Consent Forms	07/30/2016
before you start the conditioning regimen to prepare you for	(Section II):	
your transplant or cellular therapy."	Adult Allogeneic Recipient;	
	Minor Allogeneic Recipient	
	Parent/Legal Guardian;	
Paragraph 2, 2 nd sentence: "Your transplant treatment center	Recipient Consent Forms	07/30/2016
Paragraph 2, 2 nd sentence: "Your transplant treatment center and the CIBMTR have procedures in place to keep	Recipient Consent Forms (Section III):	07/30/2016
	Recipient Consent Forms (Section III): Adult Allogeneic Recipient;	07/30/2016
	Recipient Consent Forms (Section III): Adult Allogeneic Recipient; Minor Allogeneic Recipient	07/30/2016
and the CIBMTR have procedures in place to keep	Recipient Consent Forms (Section III): Adult Allogeneic Recipient; Minor Allogeneic Recipient Parent/Legal Guardian;	
and the CIBMTR have procedures in place to keep Paragraph 3, 2 nd sentence: "who need a transplant or	Recipient Consent Forms (Section III): Adult Allogeneic Recipient; Minor Allogeneic Recipient Parent/Legal Guardian; Donor Consent Forms (Section	07/30/2016
and the CIBMTR have procedures in place to keep	Recipient Consent Forms (Section III): Adult Allogeneic Recipient; Minor Allogeneic Recipient Parent/Legal Guardian; Donor Consent Forms (Section III)	
and the CIBMTR have procedures in place to keep Paragraph 3, 2 nd sentence: "who need a transplant or	Recipient Consent Forms (Section III): Adult Allogeneic Recipient; Minor Allogeneic Recipient Parent/Legal Guardian; Donor Consent Forms (Section III) Adult Allogeneic Donor;	
and the CIBMTR have procedures in place to keep Paragraph 3, 2 nd sentence: "who need a transplant or	Recipient Consent Forms (Section III): Adult Allogeneic Recipient; Minor Allogeneic Recipient Parent/Legal Guardian; Donor Consent Forms (Section III) Adult Allogeneic Donor; Adult Registered Donor for	
and the CIBMTR have procedures in place to keep Paragraph 3, 2 nd sentence: "who need a transplant or	Recipient Consent Forms (Section III): Adult Allogeneic Recipient; Minor Allogeneic Recipient Parent/Legal Guardian; Donor Consent Forms (Section III) Adult Allogeneic Donor; Adult Registered Donor for Match Algorithm Enhancement;	
and the CIBMTR have procedures in place to keep Paragraph 3, 2 nd sentence: "who need a transplant or	Recipient Consent Forms (Section III): Adult Allogeneic Recipient; Minor Allogeneic Recipient Parent/Legal Guardian; Donor Consent Forms (Section III) Adult Allogeneic Donor; Adult Registered Donor for Match Algorithm Enhancement; Minor Related Donor	
and the CIBMTR have procedures in place to keep Paragraph 3, 2 nd sentence: "who need a transplant or	Recipient Consent Forms (Section III): Adult Allogeneic Recipient; Minor Allogeneic Recipient Parent/Legal Guardian; Donor Consent Forms (Section III) Adult Allogeneic Donor; Adult Registered Donor for Match Algorithm Enhancement; Minor Related Donor Parent/Legal Guardian;	
and the CIBMTR have procedures in place to keep Paragraph 3, 2 nd sentence: "who need a transplant or	Recipient Consent Forms (Section III): Adult Allogeneic Recipient; Minor Allogeneic Recipient Parent/Legal Guardian; Donor Consent Forms (Section III) Adult Allogeneic Donor; Adult Registered Donor for Match Algorithm Enhancement; Minor Related Donor Parent/Legal Guardian; Recipient Consent Forms	
and the CIBMTR have procedures in place to keep Paragraph 3, 2 nd sentence: "who need a transplant or	Recipient Consent Forms (Section III): Adult Allogeneic Recipient; Minor Allogeneic Recipient Parent/Legal Guardian; Donor Consent Forms (Section III) Adult Allogeneic Donor; Adult Registered Donor for Match Algorithm Enhancement; Minor Related Donor Parent/Legal Guardian; Recipient Consent Forms (Section III):	
and the CIBMTR have procedures in place to keep Paragraph 3, 2 nd sentence: "who need a transplant or	Recipient Consent Forms (Section III): Adult Allogeneic Recipient; Minor Allogeneic Recipient Parent/Legal Guardian; Donor Consent Forms (Section III) Adult Allogeneic Donor; Adult Registered Donor for Match Algorithm Enhancement; Minor Related Donor Parent/Legal Guardian; Recipient Consent Forms (Section III): Adult Allogeneic Recipient;	
and the CIBMTR have procedures in place to keep Paragraph 3, 2 nd sentence: "who need a transplant or	Recipient Consent Forms (Section III): Adult Allogeneic Recipient; Minor Allogeneic Recipient Parent/Legal Guardian; Donor Consent Forms (Section III) Adult Allogeneic Donor; Adult Registered Donor for Match Algorithm Enhancement; Minor Related Donor Parent/Legal Guardian; Recipient Consent Forms (Section III): Adult Allogeneic Recipient; Minor Allogeneic Recipient	
and the CIBMTR have procedures in place to keep Paragraph 3, 2 nd sentence: "who need a transplant or cellular therapy."	Recipient Consent Forms (Section III): Adult Allogeneic Recipient; Minor Allogeneic Recipient Parent/Legal Guardian; Donor Consent Forms (Section III) Adult Allogeneic Donor; Adult Registered Donor for Match Algorithm Enhancement; Minor Related Donor Parent/Legal Guardian; Recipient Consent Forms (Section III): Adult Allogeneic Recipient; Minor Allogeneic Recipient Parent/Legal Guardian;	07/30/2016
and the CIBMTR have procedures in place to keep Paragraph 3, 2 nd sentence: "who need a transplant <u>or cellular therapy.</u> " Paragraph 1, 1 st sentence: "Your transplant <u>treatment</u> center	Recipient Consent Forms (Section III): Adult Allogeneic Recipient; Minor Allogeneic Recipient Parent/Legal Guardian; Donor Consent Forms (Section III) Adult Allogeneic Donor; Adult Registered Donor for Match Algorithm Enhancement; Minor Related Donor Parent/Legal Guardian; Recipient Consent Forms (Section III): Adult Allogeneic Recipient; Minor Allogeneic Recipient Parent/Legal Guardian; Recipient Consent Forms	
and the CIBMTR have procedures in place to keep Paragraph 3, 2 nd sentence: "who need a transplant or cellular therapy."	Recipient Consent Forms (Section III): Adult Allogeneic Recipient; Minor Allogeneic Recipient Parent/Legal Guardian; Donor Consent Forms (Section III) Adult Allogeneic Donor; Adult Registered Donor for Match Algorithm Enhancement; Minor Related Donor Parent/Legal Guardian; Recipient Consent Forms (Section III): Adult Allogeneic Recipient; Minor Allogeneic Recipient Parent/Legal Guardian; Recipient Consent Forms (Section IV):	07/30/2016
and the CIBMTR have procedures in place to keep Paragraph 3, 2 nd sentence: "who need a transplant <u>or cellular therapy.</u> " Paragraph 1, 1 st sentence: "Your transplant <u>treatment</u> center	Recipient Consent Forms (Section III): Adult Allogeneic Recipient; Minor Allogeneic Recipient Parent/Legal Guardian; Donor Consent Forms (Section III) Adult Allogeneic Donor; Adult Registered Donor for Match Algorithm Enhancement; Minor Related Donor Parent/Legal Guardian; Recipient Consent Forms (Section III): Adult Allogeneic Recipient; Minor Allogeneic Recipient Parent/Legal Guardian; Recipient Consent Forms	07/30/2016

	Parent/Legal Guardian;	
Paragraph 1, 2 nd sentence: "Your transplant treatment center	Recipient Consent Forms	07/30/2016
and the CIBMTR have procedures in place so that"	(Section IV):	0775072010
and the Cibritte have procedures in place so that	Adult Allogeneic Recipient;	
	Minor Allogeneic Recipient	
	Parent/Legal Guardian;	
Paragraph 2, last sentence: "This will not affect your	Recipient Consent Forms	07/30/2016
relationship with your transplant treatment center or	(Section VI):	07/30/2010
CIBMTR."	Adult Allogeneic Recipient;	
	Minor Allogeneic Recipient	
	Parent/Legal Guardian;	
Paragraph 1, 2 nd sentence: "your bone marrow, blood	Donor Consent Forms (Section	07/30/2016
stem cells, or cellular therapy donation will still be used in a	VII)	
transplant for the intended recipient"	Adult Allogeneic Donor;	
······································	Minor Related Donor	
	Parent/Legal Guardian;	
Paragraph 1, last sentence: "Please call your transplant	Recipient Consent Forms	07/30/2016
treatment center coordinator immediately at"	(Section VIII):	
,	Adult Allogeneic Recipient;	
	Minor Allogeneic Recipient	
	Parent/Legal Guardian;	
Paragraph 1, 1st sentence: "(Transplant Treatment Center	Recipient Consent Forms	07/30/2016
Medical Director) at"	(Section IX):	
	Adult Allogeneic Recipient;	
	Minor Allogeneic Recipient	
	Parent/Legal Guardian;	
Paragraph 1, 1 st sentence: "(Transplant <u>Treatment</u> Center	Recipient Consent Forms	07/30/2016
Coordinator) at"	(Section IX):	
	Adult Allogeneic Recipient;	
	Minor Allogeneic Recipient	
	Parent/Legal Guardian;	
Paragraph 1, 2 nd sentence: "is called a transplant or	Assent Forms:	07/30/2016
<u>cellular therapy</u> ."	Related Donor Assent 7-11	
Paragraph 1, 5 th sentence: "This research project is about		
what makes these kinds of transplants or cellular therapies		
work."		
Paragraph 2, last sentence: "you can still do the transplant		
or cellular therapy donation for your brother or sister."		
Paragraph 4, last sentence: "who are sick and need a		
transplant or cellular therapy."		
D 1 1 Ond 4 11 11 1 1 1	A 4.75	07/20/2016
Paragraph 1, 2 nd sentence: "because you will be donating	Assent Forms:	07/30/2016
stem cells or a cellular therapy product to your brother or	Related Donor Assent 12-17	
sister."		
Paragraph 1, last sentence: "The stem cells <u>or cellular</u>		
therapy product will be collected from"		
Paragraph 2, 1st sentence: "or cord blood transplants and		
cellular therapies work well."		
Paragraph 3, last sentence: "you can still do the transplant		
or cellular therapy donation for your brother or sister."		
Paragraph 5, last sentence: "who need a bone marrow or		
blood stem cell transplant or cellular therapy."		

Paragraph 1, 2 nd sentence: "The research project is about what makes transplants <u>and cellular therapies</u> work." Paragraph 2, 3 rd sentence: "Letting the CIBMTR use your blood is not about your transplant <u>or cellular therapy.</u> " Paragraph 2, last sentence: "You will have a transplant <u>or cellular therapy</u> anyway." Paragraph 4, last sentence: "and need a transplant <u>or cellular therapy</u> ."	Assent Forms: Allo Recipient Assent 7-11	07/30/2016
Paragraph 1, last sentence: "or cord blood transplant or cellular therapy." Paragraph 2, 1st sentence: "or cord blood transplants and cellular therapies work well." Paragraph 2, 2nd sentence: "who have had a transplant or cellular therapy to test different ways of matching" Paragraph 3, last sentence: "You will have a transplant or cellular therapy for your disease" Paragraph 5, last sentence: "blood stem cell transplant or cellular therapy."	Assent Forms: Allo Recipient Assent 12-17	07/30/2016
Deleted the sentence: "The NMDP Registry lists more than 11 million volunteer donors, 204,000 cord blood units (CBU), and has facilitated over 66,000 unrelated HSC transplants."	Protocol: Section 1.1	07/30/2015
Replaced sentence as follows: "As of May 2014, there are over 116,000 unique blood samples in the NMDP Research Sample Repository. Details of the research sample inventory are available at http://www.cibmtr.org/Samples/Inventory/Pages/index.aspx.	Protocol: Section 1.2	07/30/2015
Reworded last bullet point as follows: "In cases where the center has designated the NMDP IRB on their center's FederalWide Assurance as their IRB of record for the Research Repository protocol, and an IRB Authorization Agreement is in place, for the Research Repository protocol, the center does not need to obtain any additional IRB approval.	Protocol: Section 3	07/30/2015
Paragraph 3, last sentence: "Any research project may be proposed for anonymous research, examples include such as: - Studies that look at how certain genetic traits may affect transplant outcomes. - Studies that look at biological factors that may predict relapse after transplant. - Studies that look for the presence of traits linked to other diseases, like diabetes."	Donor Consent Forms (Section I): Adult Allogeneic Donor; Adult Registered Donor for Match Algorithm Enhancement; Minor Related Donor Parent/Legal Guardian; Recipient Consent Forms (Section I): Adult Allogeneic Recipient; Minor Allogeneic Recipient Parent/Legal Guardian; Marrow Toxic Injury Forms (Section I): Adult Marrow Toxic Injury; Minor Marrow Toxic Injury Parent/Legal Guardian	02/24/2015

Branding changes were applied to all consent and assent	Donor Consent Forms:	7/30/14
		//30/14
forms to remove references to NMDP (i.e.,	Adult Allogeneic Donor;	
NMDP/CIBMTR).	Adult Registered Donor for	
	Match Algorithm Enhancement;	
	Minor Related Donor	
	Parent/Legal Guardian;	
	Recipient Consent Forms:	
	Adult Allogeneic Recipient;	
	Minor Allogeneic Recipient	
	Parent/Legal Guardian;	
	Marrow Toxic Injury Forms:	
	Adult Marrow Toxic Injury;	
	Minor Marrow Toxic Injury	
	Parent/Legal Guardian	
	Assent Forms:	
	Related Donor Assent 7-11;	
	Related Donor Assent 12-17	
	Allo Recipient Assent 7-11;	
	Allo Recipient Assent 12-17	
	Marrow Toxic Injury Assent 7-	
	11;	
	Marrow Toxic Injury Assent 12-	
	17	
Title of consent/assent forms: "National Marrow Donor	Donor Consent Forms:	7/30/14
Program® (NMDP) and Center for International Blood and	Adult Allogeneic Donor;	
Marrow Transplant Research® (CIBMTR®) Contribution of a	Adult Registered Donor for	
Blood Sample to the National Marrow Donor Program's	Match Algorithm Enhancement;	
Research Sample Repository"	Minor Related Donor	
	Parent/Legal Guardian;	
	Recipient Consent Forms:	
	Adult Allogeneic Recipient;	
	Minor Allogeneic Recipient	
	Parent/Legal Guardian;	
	Marrow Toxic Injury Forms:	
	Adult Marrow Toxic Injury;	
	Minor Marrow Toxic Injury	
	Parent/Legal Guardian	
	Assent Forms:	
	Related Donor Assent 7-11;	
	Related Donor Assent 12-17	
	Allo Recipient Assent 7-11;	
	Allo Recipient Assent 12-17	
	Marrow Toxic Injury Assent 7-	
	11;	
	Marrow Toxic Injury Assent 12-	
	17	
Paragraph 1, sentence 1: "The National Marrow Donor	Donor Consent Forms (Section	7/30/14
Program (NMDP) and the Center for International Blood and	I):	7/30/11
Marrow Transplant Research (CIBMTR), the research	Adult Allogeneic Donor;	
program of the National Marrow Donor Program	Adult Anogenete Donor, Adult Registered Donor for	
(NMDP)/Be The Match, invites you to take part in the	Match Algorithm Enhancement	
Research Sample Repository."	Recipient Consent	
Research sample repusitory.	Forms(Section I):	
	roffiis(Section 1):	1

	Adult Allogeneic Recipient Marrow Toxic Injury Consent Forms (Section I):	
	Adult Marrow Toxic Injury	
Paragraph 1, sentence 1: "The National Marrow Donor Program (NMDP) and the Center for International Blood and Marrow Transplant Research (CIBMTR), the research	Donor Consent Forms (Section I): Minor Related Donor	7/30/14
program of the National Marrow Donor Program (NMDP)/Be The Match, invites your child to take part in a Research Sample Repository."	Parent/Legal Guardian Recipient Consent Forms (Section I):	
	Minor Allogeneic Recipient Parent/Legal Guardian Marrow Toxic Injury Consent Forms (Section I):	
	Minor Marrow Toxic Injury Parent/Legal Guardian	7/20/14
Deleted the sentence, "In these studies, there will be no way for the sample to be linked to you."	Donor Consent Forms (Section I):	7/30/14
	Adult Allogeneic Donor; Adult Registered Donor for Match Algorithm Enhancement Recipient Consent Forms	
	(Section I): Adult Allogeneic Recipient Marrow Toxic Injury Consent Forms (Section I):	
	Adult Marrow Toxic Injury	
Deleted the sentence, "In these studies, there will be no way	Donor Consent Forms (Section	7/30/14
for the sample to be linked to your child."	I): Minor Related Donor Parent/Legal Guardian	
	Recipient Consent Forms (Section I):	
	Minor Allogeneic Recipient Parent/Legal Guardian	
	Marrow Toxic Injury Consent Forms (Section I):	
	Minor Marrow Toxic Injury Parent/Legal Guardian	
"All research studies using these blood samples must first be approved by a group of scientists within the	Donor Consent Forms (Section II):	7/30/14
NMDP/CIBMTR as well as the Repository Oversight Committee. NMDP The proposed study will also be reviewed the proposed study to make sure the research is	Adult Allogeneic Donor; Adult Registered Donor for Match Algorithm Enhancement;	
consistent with the types of studies described above."	Minor Related Donor Parent/Legal Guardian	
	Recipient Consent Forms (Section II):	
	Adult Allogeneic Recipient; Minor Allogeneic Recipient Parent/Legal Guardian Marrow Toxic Injury Consent	
	Forms (Section II):	

	A 1 1 M CD . T .	
	Adult Marrow Toxic Injury;	
	Minor Marrow Toxic Injury	
	Parent/Legal Guardian	
"Your donor center and the NMDP/CIBMTR has have	Donor Consent Forms (Section	7/30/14
procedures in place"	IV):	
	Adult Allogeneic Donor;	
	Adult Registered Donor for	
	Match Algorithm Enhancement	
"Your child's donor center and the NMDP/CIBMTR has	Donor Consent Forms (Section	7/30/14
have procedures in place"	IV):	
	Minor Related Donor	
	Parent/Legal Guardian	
"Your treatment center and the NMDP/CIBMTR has have	Recipient Consent Forms	7/30/14
procedures in place"	(Section IV):	
	Adult Allogeneic Recipient	
	Marrow Toxic Injury Consent	
	Forms (Section IV):	
	Adult Marrow Toxic Injury	
"Your child's treatment center and the NMDP/CIBMTR has	Recipient Consent forms	7/30/14
have procedures in place"	(Section IV):	
	Minor Allogeneic Recipient	
	Parent/Legal Guardian	
	Marrow Toxic Injury Consent	
	Forms (Section IV):	
	Minor Marrow Toxic Injury	
	Parent/Legal Guardian	
"NMDP Be The Match Donor Advocacy"	Donor Consent Forms (Section	7/30/14
	IX):	
	Adult Allogeneic Donor;	
	Adult Registered Donor for	
	Match Algorithm Enhancement;	
	Minor Related Donor	
	Parent/Legal Guardian	
"with Be the Match® Patient and Health Professional	Recipient Consent Forms	7/30/14
Services"	(Section IX):	
	Adult Allogeneic Recipient;	
	Minor Allogeneic Recipient	
	Parent/Legal Guardian	
	Marrow Toxic Injury Consent	
	Forms (Section VIII):	
	Adult Marrow Toxic Injury;	
	Minor Marrow Toxic Injury	
	Parent/Legal Guardian	
Removed National Marrow Donor Program from title page.	Protocol: Title Page	7/30/14
As of May 2013 2014, there are over 100,000 116,000	Protocol: Section 1.2	7/30/14
unique blood samples		
The NMDP Registry lists more than 11 million volunteer	Protocol: Section 1.1	7/30/14
donors, 190,000 204,000 cord blood units (CBU), and has		
facilitated over 55,000 <u>66,700</u> unrelated HSC transplants.		
Addition:	Protocol: Table of Contents on	7/30/13
8.6 Public Release of Data Generated on Samples11	page 2	,,,50,15
At the end of December 2008, the The NMDP Registry listed	Protocol: Section 1.1	7/30/13
lists more than 7.5 11 million volunteer donors, 28,000	110tucui. Section 1.1	1/30/13
<u> 11515</u> 111010 than 7.3 <u>11</u> 111111011 volunteel donois, 20,000		1

190,000 cord blood units (CBU), and had has facilitated over		
35,000 55,000 unrelated HSC transplants.		
First paragraph: "As of May 2012 2013, there are over 90,000 100,000 unique blood samples"	Protocol: Section 1.2	7/30/13
First paragraph: "Twenty Up to thirty milliliters (20 30 mL) of blood"	Protocol: Section 4.1.1	7/30/13
Second paragraph: "Samples from registered volunteer donors with rare tissue types will be collected at the donor's convenience any time after registration. Twenty Up to thirty milliliters (20 30 mL) of blood are collected from adult donors, except up to fifty (50 mL) may be collected from adult donors with rare tissue types."	Protocol: Section 4.1.1	7/30/13
First paragraph: "Ten Up to thirty milliliters(10 30 mL) of blood"	Protocol: Section 4.1.2	7/30/13
Second paragraph: "For all donors, 10 up to thirty milliliters (10 30 mL) of blood are collected."	Protocol: Section 4.1.2	7/30/13
First paragraph: "Twenty Up to thirty milliliters (20 30 mL) of blood are collected from an adult."	Protocol: Section 4.1.3	7/30/13
"Donor and recipient samples are stored as whole blood, peripheral blood mononuclear cells, cell lysates, extracted DNA, stabilized RNA, serum and plasma."	Protocol: Section 5.1	7/30/13
 Added second paragraph: "In summary, all sample requests must meet the following release criteria prior to distribution of samples: The proposed use of samples falls under the acceptable uses defined under section 8.1 for linked research or 8.2 for anonymous research per NMDP IRB Chair review. The proposed study is deemed scientifically sound, feasible and high impact through acceptance by a CIBMTR Working Committee or The planned use is for reference or quality control material only. The proposed study is approved by the Repository Oversight Committee. Consent status is confirmed in the CIBMTR database." 	Protocol: Section 7.2	7/30/13
First paragraph: "Any research project may be proposed for anonymous research, examples include: • Studies that require self-identified race/ethnicity or other demographically defined healthy and/or disease controls. • Studies that need HLA specific immune stimulators for in vitro assays."	Protocol: Section 8.2	7/30/13
Added Section 8.6:	Protocol: Section 8.6	7/30/13
8,6 Public Release of Data Generated on Samples Research studies using Research Repository samples funded through the National Institutes of Health (NIH) are subject to the public data release policies of the NIH. The deposition of testing data from Research Repository samples into the NIH database of Genotypes and Phenotypes (dbGAP) will be permitted under the following conditions:		

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• All sample data is stripped of identifying sequences, e.g.		
Y chromosome, mitochondrial DNA, or other unique		
sequences, dates and detailed demographic data prior to		
submission to dbGAP.		
Access to sample data through dbGAP is limited to the		
controlled-access data process and use limited to		
research purposes defined in the consent.		
research purposes defined in the consent.		
	D G 17 (G 1	F/20/42
Last paragraph: "In addition, Investigators may conduct"	Donor Consent Forms (Section	7/30/13
and "These studies are not limited to the types of studies	I):	
listed above, or related to transplantation in general. Any	Adult Allogeneic Donor;	
research project may be proposed for anonymous research.	Adult Registered Donor for	
examples include:	Match Algorithm Enhancement;	
Studies that look at how certain genetic traits may	Minor Related Donor	
affect transplant outcomes.	Parent/Legal Guardian;	
Studies that look at biological factors that may	Recipient Consent Forms	
predict relapse after transplant."	(Section I):	
prodet relapse arter dansplant.	Adult Allogeneic Recipient;	
	Minor Allogeneic Recipient	
	Parent/Legal Guardian;	
	Marrow Toxic Injury Forms	
	(Section I):	
	Adult Marrow Toxic Injury;	
	Minor Marrow Toxic Injury	
	Parent/Legal Guardian	
First paragraph, last sentence: "research studies <u>related to</u>	Donor Consent Forms (Section	7/30/13
transplant or other research as defined in this consent form."	X):	
	Adult Allogeneic Donor;	
	Adult Registered Donor for	
	Match Algorithm Enhancement;	
	Minor Related Donor	
	Parent/Legal Guardian;	
	Recipient Consent Forms	
	(Section X):	
	Adult Allogeneic Recipient;	
	Minor Allogeneic Recipient	
	Parent/Legal Guardian;	
	Marrow Toxic Injury Forms	
	(Section IX):	
	Adult Marrow Toxic Injury;	
	Minor Marrow Toxic Injury	
	Parent/Legal Guardian	
First sentence: "will be collected from a vein in your	Donor Consent Forms (Section	7/30/13
child's arm <u>body</u> ."	II):	
	Minor Related Donor	
	Parent/Legal Guardian;	
	Recipient Consent Forms	
	(Section II):	
	Minor Allogeneic Recipient	
	Parent/Legal Guardian;	
	Marrow Toxic Injury Forms	
	(Section II):	
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	I willor wiallow Toxic Hijury	1

	Parent/Legal Guardian	1
T'	Donor Consent Forms (Section	7/30/13
First sentence: "where the blood is taken from your child's arm."	III): Minor Related Donor Parent/Legal Guardian; Recipient Consent Forms (Section III): Minor Allogeneic Recipient Parent/Legal Guardian; Marrow Toxic Injury Forms (Section III): Minor Marrow Toxic Injury Parent/Legal Guardian	//30/13
Paragraph 3, 1 st sentence: "The blood sample will come from a n arm -vein in your body."	Assent Forms: Minor Related Donor (7-11)	7/30/13
Paragraph 3, 1 st sentence: "will be taken from a vein in your arm body." Paragraph 4, 1 st sentence: "When the blood is taken from your arm you will"	Assent Forms: Minor Related Donor (12-17)	7/30/13
Paragraph 3, 3 rd sentence: "There is a chance the blood may have to come from a n arm -vein."	Assent Forms: Minor Allo Recipient (7-11)	7/30/13
Paragraph 3, 1 st sentence: "will be taken from your catheter or from a vein in your arm body." Paragraph 4, 2 nd sentence: "If the blood is taken from a vein in your arm, you will"	Assent Forms: Minor Allo Recipient (12-17)	7/30/13
Paragraph 2,1st sentence: "someone will take a small amount of blood from you r arm ."	Assent Forms: Minor Marrow Toxic Injury (7-11)	7/30/13
Paragraph 3, 1 st sentence: "will be taken from your catheter or from a vein in your arm body." Paragraph 4, 2 nd sentence: "If the blood is taken from a vein in your arm, you will"	Assent Forms: Minor Marrow Toxic Injury (12-17)	7/30/13
Last sentence: "The NMDP/CIBMTR will try hard to make sure has procedures in place so that no one outside"	Donor Consent Forms (Section IV): Adult Allogeneic Donor; Adult Registered Donor with Rare Tissue Type; Minor Related Donor Parent/Legal Guardian; Recipient Consent Forms (Section IV): Adult Allogeneic Recipient; Minor Allogeneic Recipient Parent/Legal Guardian; Marrow Toxic Injury Forms (Section IV): Adult Marrow Toxic Injury; Minor Marrow Toxic Injury; Parent/Legal Guardian	7/30/12

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Paragraph 2: Second sentence, "please contact a Patient Services Coordinator with the NMDP Office of Patient Advocacy Be the Match® Patient Services at"	Recipient Consent Forms (Section IX): Adult Allogeneic Recipient; Minor Allogeneic Recipient Parent/Legal Guardian Marrow Toxic Injury Forms (Section VIII): Adult Marrow Toxic Injury; Minor Marrow Toxic Injury Parent/Legal Guardian	7/30/12
Paragraph 1: Last sentence, "As of May 2009 2012, there are over 46,000 90,000 unique blood samples"	Protocol: Section 1.2	7/30/12
Paragraph 2: Last sentence, "For all donors, 20 Twenty milliliters (20 mL) of blood are collected from adult donors except up to fifty (50 mL) may be collected from adult donors with rare tissue types."	Protocol: Section 4.1.1	7/30/12
Paragraph 1: Second sentence, "This includes, <u>but is not limited to</u> , any of the following"	Protocol: Section 8.1	7/30/12
Added last bullet point, "Studies of global genetic diversity through genome-wide association studies or other techniques to evaluate the impact of other genetic factors on transplant outcome."	Protocol: Section 8.1	7/30/12
Discontinued the consent form "Adult Registered Donor with Rare Tissue Type." (Although effective 7/30/2011, this change was not added to the Record of Revisions until 10/10/2011 due to an oversight.)	Donor Consent Form: Adult Registered Donor with Rare Tissue Type	7/30/11
Paragraph 1: Last sentence, "You are being asked to participate because <u>you</u> have been exposed"	Assent Forms: Minor Marrow Toxic Injury (12-17)	7/30/11
Paragraph 6: 2 nd sentence, "Your doctors or your parents eannot will not make you give the blood sample"	Assent Forms: Minor Related Donor (12-17); Minor Allogeneic Recipient (12-17); Minor Marrow Toxic Injury (12-17)	7/30/11
Paragraph 2: Last sentence, "No identifiable information about your child will be given to the researchers, nor will it be published or presented at scientific meetings." Paragraph 2: Last sentence, "No identifiable information	Donor Consent Forms (Section III): Minor Related Donor Parent/Legal Guardian Recipient Consent Forms (Section III): Minor Allogeneic Recipient Parent/Legal Guardian Marrow Toxic Injury Forms (Section III): Minor Marrow Toxic Injury Parent/Legal Guardian Donor Consent Forms (Section	7/30/11
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	(Section III): Adult Allogeneic Recipient; Marrow Toxic Injury Forms (Section III): Adult Marrow Toxic Injury;	
Added second sentence to last paragraph: "Remember, you can change your mind at any time."	Assent Forms: Minor Allo Recipient (7 to 11); Minor Marrow Toxic Injury (7 to 11)	7/30/10
Added second sentence to last paragraph: "Remember, you can change your mind at any time and still give your special cells to your brother or sister."	Assent Forms: Minor Related Donor (7 to 11)	7/30/10
Second paragraph: "If you agree to be in this research project, a nurse someone will take a small sample amount of blood"	Assent Forms: Minor Marrow Toxic Injury (7 to 11)	7/30/10
Second paragraph: "If you want to be in this research project, your doctor someone will take a small amount"	Assent Forms: Minor Related Donor (7 to 11); Minor Allo Recipient (7 to 11)	7/30/10
Moved the first sentence of the first paragraph, "You are <u>also</u> being invited to be in a research project with the NMDP and CIBMTR." to be the fourth sentence.	Assent Forms: Minor Related Donor (7 to 11)	7/30/10
Paragraph 2: added second sentence "If you wish to contact an independent third party not connected with this study about problems, concerns, questions, information, or input, please contact a Patient Services Coordinator with the NMDP Office of Patient Advocacy at 1-888/999-6743 or patientinfo@nmdp.org."	Recipient Consent Forms (Section IX): Adult Allogeneic Recipient; Minor Allogeneic Recipient Parent/Legal Guardian Marrow Toxic Injury Forms (Section VIII): Adult Marrow Toxic Injury; Minor Marrow Toxic Injury Parent/Legal Guardian	7/30/10
Paragraph 1: Deleted the sentence "NMDP will pay for this treatment."	Donor Consent Forms (Section VIII): Minor Related Donor Parent/Legal Guardian	7/30/10
Paragraph 2; added second sentence "If you wish to contact an independent third party not connected with this study about problems, concerns, questions, information, or input, please contact NMDP Donor Advocacy at 1-800/526-7809, extension 8710."	Donor Consent Forms (Section IX): Adult Related/Unrelated Donor; Minor Related Donor Parent/Legal Guardian; Adult Rare HLA Type Donor	7/30/10
Reworded first sentence as follows: "If you have questions, or complaints about"	Donor Consent Forms (Section IX): Adult Related/Unrelated Donor; Minor Related Donor Parent/Legal Guardian; Adult Rare HLA Type Donor Recipient Consent Forms (Section IX): Adult Allogeneic Recipient; Minor Allogeneic Recipient Parent/Legal Guardian	7/30/10

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Changed year to 2009	Protocol: Title page	7/30/09
Updated number of volunteer donors and transplants	Protocol: Section 1.1	7/30/09
Updated number of unique samples in repository	Protocol: Section 1.2	7/30/09
Added "Transplant Center"	Protocol: Section 4.2	7/30/09
Added line to write in Donor ID on each page of donor	Donor Consent Forms:	7/30/09
consents	Adult Related/Unrelated Donor;	1/30/09
Consents	Adult Match Algorithm Donor;	
	Adult Rare HLA Type Donor	
Added "or tissue" to Section I, second paragraph, second	Donor Consent Forms:	7/30/09
sentence.	Adult Match Algorithm Donor;	1/30/07
somenee.	Adult Rare HLA Type Donor	
Reworded 3 rd bullet in Section I as follows: "Study the	Donor Consent Forms:	7/30/09
distribution of tissue types in different populations; e.g., for	Adult Related/Unrelated Donor;	1130107
example, study tissue typing differences between different	Adult Match Algorithm Donor;	
various racial and ethnic populations, to help develop	Adult Rare HLA Type Donor;	
methods that will to improve tissue matching between donors	Minor Related Donor	
and recipients."	Parent/Legal Guardian;	
and recipions.	Recipient Consent Forms:	
	Adult Allogeneic Recipient;	
	Minor Allogeneic Recipient	
	Parent/Legal Guardian	
Added "and Marrow Toxic Injuries" to title.	Protocol: Title page	7/30/08
Updated number of donors and HSC transplants through	Protocol: Section 1.1	7/30/08
December 2007.	Trotocon Section 1.1	7730700
Updated number of unique blood samples in research	Protocol: Section 1.2	7/30/08
repository through May 2008.	Trottocon Section 1.2	7730700
Added delinked research as fourth type of study where	Protocol: Section 1.2	7/30/08
samples could be used		1,,,,,,,,,
Included use of donors for tissue type characterization and	Protocol: Section 2.1	7/30/08
enhancement of search algorithm.		
Created eligibility of individuals with marrow toxic injury.	Protocol: Section 2.4	7/30/08
Created sample collection from individuals with marrow	Protocol: Section 4.1.3	7/30/08
toxic injury.		
Created sample collection from donors for tissue type	Protocol: Section 4.1.4	7/30/08
characterization and algorithm enhancement.		
Added "and plasma."	Protocol: Section 5.1	7/30/08
Clarified that studies for delinked research follow the same	Protocol: Section 8.2	7/30/08
review/approval process as outlined in Section 7.2 of the		
Protocol		
Created consent form for donors for tissue type	Adult Registered Donor for	7/30/08
characterization and algorithm enhancement.	Match Algorithm Enhancement	
	Consent Form	<u> </u>
Created consent form for adults with marrow toxic injury.	Adult Marrow Toxic Injury	7/30/08
	Research Sample Consent Form	
Created permission form for minors with marrow toxic	Minor Marrow Toxic Injury	7/30/08
injury.	Parent/Legal Guardian	1/30/00
	Permission Form	
Created minor assent form for minors with marrow toxic	Minor Marrow Toxic Injury	7/30/08
injury (7 to 11 years old).	Assent Form (7 to 11 years of	1/30/00
injury (7 to 11 years old).	age)	
Created minor assent form for minors with marrow toxic	Minor Marrow Toxic Injury	7/30/08
injury (12 to 17 years old).	Assent Form (12 to 17 years of	1/30/08
mjury (12 to 17 years old).	Assem Form (12 to 17 years of	

	age)	
Revised wording regarding anonymous research to "In	Consent Forms Section I	7/30/08
addition, investigators may conduct resarch studies with	Consent i ornis section i	7/30/00
stored blood samples that have had all identifiers removed.		
In these studies, there will be no way for the sample to be		
linked to you. NMDP/CIBMTR may allow investigators to		
use these anonymous samples for many other kinds of		
studies. Theses studies are not limited to the types of studies		
listed above, or related to transplantation in general."		
Removed the sentence "Your blood sample will be used to	Consent Forms Section II	7/30/08
look at ways to improve how patients are matched with their	Consent Forms Section II	7/30/08
donors."		
	Consent Forms Section II	7/30/08
Added the sentence "Your blood sample will be frozen and	Consent Forms Section II	1/30/08
stored indefinitely for possible use in future research		
studies."		7 (20) (00)
Added that studies must be reviewed by a group of scientists	Consent Forms Section II	7/30/08
"within NMDP/CIBMTR as well as the Repository		
Oversight Committee."		5 (20) (00)
Removed mention that studies must be reviewed by the	Consent Forms Section II	7/30/08
NMDP IRB and replaced it with "NMDP will also review		
the proposed study"		= 12 0 10 0
Removed the sentence "An IRB is a group of people who	Consent Forms Section II	7/30/08
protect the rights of research participants."		
Clarified "No identifiable information about you" rather than	Consent Forms Section III	7/30/08
just "Your name".		
Added "Center for International Blood and Marrow	Consent Forms	7/30/08
Transplant Research (CIBMTR") to title.		= /= 0 /0 0
Changed "study" to "project."	Minor Assent Forms (7-11)	7/30/08
In last paragraph, changed "want to be in the research	Minor Assent Forms (7-11)	7/30/08
project" to "agree to give a small amount of blood for		
research."		= 12 0 10 0
Throughout form replaced various mentions of "in this	Minor Assent Forms (12-17)	7/30/08
study" to various mentions of "give a blood sample for		
research."		
Removed "The NMDP would like the medical staff at your	Minor Related Donor Assent	7/30/08
donor center to take"	Form (12-17)	
Removed "The NMDP would like your doctor to have one of	Minor Allogeneic Recipient	7/30/08
the medical staff at your hospital take"	Assent Form (12-17)	
Added "including testing of rare HLA types" to the third	Protocol: Section 1.2	7/30/07
bulleted type of research studies		
Clarified that NMDP IRB review is done "administratively"	Protocol: Section 8.0	7/30/07
by the NMDP IRB Chair		
Clarified review of delinked research studies, documenting	Protocol: Section 8.2	7/30/07
NMDP IRB review is done "administratively" by the NMDP		
IRB Chair		
Revisions to section 2.4 <i>Informed Consent</i> , to include	Protocol: Section 2.4	7/30/07
mention of assent and refer to parental consent as		
"permission"		
Revisions to Minor Assent section to state local IRBs are	Protocol: Section 2.4.1	7/30/07
responsible for determination of method to document minor		
assent		
Include caveat that minor must be "capable of providing	Protocol: Section 2.4.1	7/30/07

	T	
assent" and confirm parent/legal guardian permission is		
sufficient if minor lacks capacity to provide assent		7/20/07
Included language addressing use of samples for anonymous research	Consent Forms Section I	7/30/07
Revised mention of Parent/Legal Guardian "consent" to	Legal Guardian Consent Forms	7/30/07
"permission" in title and section statement section		
Added full board name for IRB	Consent Forms Section II	7/30/07
To avoid repetitive language, revised section III stating the	Consent Forms Section III	7/30/07
NMDP/CIBMTR will try hard to avoid a loss of		
confidentiality to read "NMDP/CIBMTR have procedures in		
place to keep your data private"		
Replaced use of "quitting" in two instances to "change your	Consent Forms Section IV	7/30/07
mind" and "this" in the withdrawal language		
Updated site of Repository	Protocol: Section 1.2	6/11/07
Removed the option allowing centers to prepare site specific	Protocol: Section 3	6/11/07
protocol		
Updated section to include two sub-sections, one addressing	Protocol: Section 4.1	6/11/07
collections for unrelated transplants and one addressing		
collections for related transplants		
Included option for cell storage per "dried blood on filter	Protocol: Section 5.1	6/11/07
paper"		
Revised section to allow for NMDP IRB Chair	Protocol: Section 7.2	6/11/07
administrative review for requests for samples – use of		
samples not considered "human research"		
Added documentation that "Requestor will not receive any	Protocol: Section 8.5	6/11/07
identifying information with the samples that could possibly		
be used to link the sample to the contributing individual."		
Added documentation that "The link will never be released	Protocol: Section 10.1	6/11/07
to an investigator."		
Attachment removed – sites no longer allowed to prepare	Protocol: Attachment 1	6/11/07
center specific protocol		
Added CIBMTR to study invitation and referred to	Consent Forms:	6/11/07
NMDP/CIBMTR throughout consent form	Recipient, Legal Guardian/Parent,	
	Minor Assent Forms, Donor	
For inclusion of related donors/recipients the types of studies	Consent Forms:	6/11/07
for which samples may be used revised to state	Recipient, Legal Guardian/Parent,	
"understanding tissue matching of <u>related and</u> unrelated	Minor Assent Forms, Donor,	
donors and recipients	Rare HLA Type	6/11/07
Revised statement regarding NMDP IRB approval to reflect	Consent Forms:	6/11/07
administrative review process "The studies will also be	Recipient, Legal Guardian/Parent,	
reviewed by the NMDP IRB to make sure the research is	Minor Assent Forms, Donor, Rare	
consistent with the types of studies described above. An IRB	HLA Type	
is a group of people who protect the rights of research participants."		
Revised language in Confidentiality section to state the	Consent Forms:	6/11/07
NMDP/CIBMTR will not "intentionally" disclose subject's	Recipient, Legal Guardian/Parent,	0/11/0/
participation and will make every effort to maintain strict	Minor Assent Forms, Donor, Rare	
confidentiality	HLA Type	
Prepared Parent/Legal Guardian consent form and Minor	New Consent Forms	6/11/07
Assent forms for related donors requested to participate	Tion Consent Forms	0/11/0/
Updated the number of unique blood samples included in the	Protocol: Section 1.2	7/30/06
Repository to "over 38,000" as of May 2006	1100001. 5000011.2	,,,50,00
repository to over 30,000 as or iviay 2000	<u>l</u>	

Clarified "parent <u>or legal guardian</u> " as the entity responsible for providing permission for minors to participate	Protocol: Section 2.4.1	7/30/06
Minor modifications clarifying the NMDP IRB Office's role in the IRB approval process for the Repository	Protocol: Section 3.1, 3.2	7/30/06
Updated the process by which requests for samples receive approval from the NMDP/CIBMTR	Protocol: Section 7.2	7/30/06
Revised wording regarding risks of collecting the blood sample from "may cause minor discomfort" to "will likely cause minor discomfort"	Recipient Consent Section III Legal Guardian Consent Section III Rare Tissue Type Donor Consent Section III	7/30/06
Revised wording regarding risks of collecting the blood sample from "you might feel some pain" to "will probably feel some pain"	Minor Assent Form (7-11) Minor Assent Form (12-17)	7/30/06
Revised wording regarding risk of identification of participant from "small risk that someone could find out which blood sample is yours" to small risk that an unauthorized person could find out which blood sample is yours"	Recipient Consent Form Section III Legal Guardian Consent Form Section III Rare Tissue Type Donor Consent Form Section III	7/30/06
Added sentence "It is up to you if you want to participate in the Research Repository"	Recipient Consent Form Section VI Legal Guardian Consent Form Section VI Rare Tissue Type Donor Consent Form Section VI	7/30/06
Corrected voluntary participation and withdrawal language to "you and your child"	Legal Guardian Consent Form Section VI	7/30/06
Removed the phrase "My signature below says that" from the subject's statement of consent	Recipient Consent Form Section X Legal Guardian Consent Form Section X Rare Tissue Type Donor Consent Form Section X	7/30/06
Discontinued use of donor consent combining language regarding participation in Research Repository and Research Database		7/30/06
Prepared separate consent form for donor participation in Research Sample Repository		7/30/06
New Principal Investigator	Protocol	7/30/05
Replaced Dennis Confer, M.D. with space to provide Donor Center Medical Director contact information	Donor Consent Section VIII	7/30/05
Corrected "You do not waive any liability rights for personal injury by signing this form" to "You do not waive any legal rights by signing this form."	Donor Consent Section VIII Donor w/ Rare Tissue Type Section VIII Recipient Consent Section VIII	7/30/05
Identification of the NMDP IRB as the group of people who monitor the use the data and protect the participant's rights.	Donor Consent Section II Recipient Consent Section II Donor w/ Rare Tissue Type Section II	7/30/05
Changed "Legal Guardian Consent" to "Parental /Legal Guardian Signature" and updated signature lines to	Minor Assent Form (7 to 11) Minor Assent Form (12 to 17)	7/30/05

"Parent/Legal Guardian"		
Prepared Parental/Legal Guardian consent form to be used		7/30/05
with the minor assent forms		1/30/03
Protocol was revised, in accordance with the OHRP	Protocol Sections:	5/20/05
"Guidance on Research Involving Coded Private Information	2.2 (revised), 2.4 (revised),	3/20/03
of Biological Specimens" to include a provision to obtain	2.4.2 (new), 4.2 (revised), and	
Cord Blood Unit (CBU specimens).	5.1 (revised)	E /1 /0E
Volume of recipient blood sample reduced from 40mL to	Protocol; Section 4.1, Attachment	5/1/05
20mL (three tablespoons to two tablespoons)	Desirient Consent forms Continu	
	Recipient Consent form; Section	
"C	II Protocol; Section 5.1	5/1/05
"Separation and" removed from section 5.1 header	Protocol; Section 3.1	3/1/03
Removed language stating "When a recipient sample is	Protocol; Section 5.1	5/1/05
received an attempt to made to isolate, collect and store	,	
peripheral blood mononuclear cells, granulocytes and		
serum."		
Removed "on every recipient sample received at the	Protocol; Section 5.2	5/1/05
Repository and" and replaced with "and recipient"	,	
Updated samples in the repository to reflect 32,000 as of	Protocol	7/30/04
April 2004	Section 1.2, Paragraph 1, Line 7	
Revised the volume of donor sample that will be drawn. The	Protocol	7/30/04
volume has been changed from 30mL to 20mL for donors.	Section 4.1 Paragraph 3, Line 4	
Attachment 1 updated to include sample language from	Protocol	7/30/04
revised consent forms and the IRB recommendation to	Attachment 1	
include a section to document the attestation of a counseling		
healthcare professional.		
Added the mention of cord blood transplants to the NMDP	Rare Tissue Type Consent form	7/30/04
goal of research	Section I, Paragraph 2, Line 2,3	
Added the mention of cord blood transplants to the NMDP	Rare Tissue Type Consent form	7/30/04
goal of research.		
Removed "HLA" since the revised language refers to HLA	Rare Tissue Type Consent form	7/30/04
type as "tissue type" only.	Section II, Paragraph 2, Line 2, 3	
Changed the sentence to read "The studies must also be	Rare Tissue Type Consent form	7/30/04
approved by a group of people who monitor the use of your	Section I, Bulleted Point 2	
data and protect your rights."	,	
Changed the statement that the blood sample would not be	Rare Tissue Type Consent form	7/30/04
used in any other research to the more correct statement that	Section VI, Paragraph 1, Line 2	
the sample "will be destroyed".		
Modified the format of the Donor/Subject Statement of	Rare Tissue Type Consent form	7/30/04
Consent	Section X	
Consent form re-written at a more appropriate reading level	Recipient Consent form	7/30/04
Consent form re-written at a more appropriate reading level	Cord Blood Consent form	7/30/04
	D C is	7/20/04
Consent form prepared to combine consent for donors to	Donor Consent form	7/30/04
participate in both the Research Database and Research	New consent form	
Repository studies.	D	7/20/04
Previous consent form for donor participation in Research	Previous consent form for	7/30/04
Repository only, was removed from the study	participation in Research	
	Repository only, now replaced	
	with combined consent form.	<u> </u>

Protocol for a Research Sample Repository for Hematopoietic Cell Transplantation, Other Cellular Therapies and Marrow Toxic Injuries (IRB-1991-0002)

Added "registered" to the following sentence: donors and	Protocol	1/28/04
recipients who have either "registered", donated or been	Section 1.2, Paragraph 1, Line 5	
transplanted		
Modified wording of "There are currently" to "As of July	Protocol	1/28/04
2003 there are over 29,000 unique blood samples in the	Section 1.2, Paragraph 1, Line 7	
NMDP Research Sample Repository."		
Added section to address the types of studies that NMDP	Protocol	1/28/04
allows samples to be used without obtaining additional	Section 1.2, Paragraph 3	
consent from the participants		
Paragraph added indicating registered donors with rare tissue	Protocol	1/28/04
types will be included as eligible participants in the Research	Section 2.1, Paragraph 2	
Repository.		
Collection time for registered donors with rare tissue type	Protocol	1/28/04
added.	Section 4.1, Paragraph 3, Lines 2-	
	4	
Prepared an additional Research Repository consent form to	Consent form titled:	1/28/04
be used by registered donors with rare tissue types	Contribution of a Blood Sample	
, ,	to the National Marrow Donor	
	Program's Research Sample	
	Repository: Registered Donor	
	with Rare Tissue Type Consent	
	Form	
Formatting changes	Protocol	10/1/03
Number of transplants updated Number of stored samples	Recipient/Subject Consent	
updated	Donor/Subject Consent	
	Cord Blood Donor/Subject	
Section added addressing justification of Minor Assent	Protocol Section 2	10/1/03
Section added outlining IRB approval process for centers	Protocol Section 3	10/1/03
contributing a research sample		
Revision to donor samples used for cell transformation	Protocol Section 5.2	10/1/03
Section added to address participant withdrawal from the	Protocol Section 9	10/1/03
Research Repository		
Attachment added defining minimum requirements set forth	Protocol Attachment 1	10/1/03
by the NMDP IRB for centers writing their own protocols		
and consent forms		
Reference to sample being separated into cells and plasma	Sections I, II	10/1/03
removed	Recipient/Subject Consent	
	Donor/Subject Consent	
	Cord Blood Donor/Subject	
	Consent	1
Phrase "ethnic" replaced with "racial and ethnic"	Sections I, II	10/1/03
	Recipient/Subject Consent	
	Donor/Subject Consent	
	Cord Blood Donor/Subject	
	Consent	
Section added to address alternatives to participation	Section VII	10/1/03
	Recipient/Subject Consent	
	Donor/Subject Consent	
	Cord Blood Donor/Subject	
	I C	

Consent

The "Authorization to Use and Disclose Health Information for Research Purposes" removed	Recipient/Subject Consent Donor/Subject Consent Cord Blood Donor/Subject Consent	10/1/03
Statement "The NMDP will pay for this treatment." removed from consent form for recipients	Sections VIII Recipient/Subject Consent	10/1/03
"Unrelated donors" changed to "Cord Blood Units" or "CBUs"	Cord Blood Donor/Subject Consent	10/1/03
Minor Assent for ages 7 to 11 approved		10/1/03