# Align OPTN Policy with U.S. Public Health Service Guideline, 2020: Donor Specimen Storage Resource

### **Donor blood specimen storage requirements**

<b>Donor Type</b>	Collection Timeframe	Storage Timeframe	OPTN Policy
Deceased	Samples must be collected within 24 hours prior to	Samples must be stored for at least 10 years after the	2.2: OPO responsibilities
	organ procurement.	date of organ transplant.	3/1/21 addition: Timeframe for specimen collection
Living*	Samples must be collected within 24 hours prior to organ recovery.	Samples must be stored for at least 10 years after the date of organ transplant.	14.8.B: Living Donor Specimen Collection and Storage
			6/1/21 addition: Requirement for living donor sample storage and timeframe for specimen collection

#### Why is it necessary to store donor blood specimens?

• To identify if transplant recipient HIV, HBV, HCV infection is donor-derived. As the CDC has previously stated, "Appropriate specimen collection, labeling, transportation, handling, and storage facilitate the accuracy of reported laboratory test results."

#### What are the requirements for the living and deceased donor sample being stored?

- OPTN policy does not have specific requirements for type, number, preparation for the blood specimens to be stored, or for the temperature of sample storage.
- The <u>2020 PHS Guideline</u> recommends two blood specimens be collected, one
  ethylenediaminetetraacetic acid (EDTA) plasma or serum specimen for serologic assays and a
  separate EDTA plasma specimen for NAT.
  - O If it is only feasible to collect one specimen, a plasma specimen in EDTA is optimal (EDTA has the purple or lavender top).<sup>2</sup> The 2013 PHS Guideline notes that "Labeling each specimen with a minimum of two unique identifiers ensures a confidential and unbroken chain of traceability to the identity of the donor and recipient."<sup>3</sup>

<sup>&</sup>lt;sup>1</sup> DL Seem, I Lee, C Umscheid, et al, "Public Health Service Guideline for Reducing Transmission of Human Immunodeficiency Virus (HIV) Hepatitis B Virus (HBV), and Hepatitis C Virus (HCV) Through Organ Transplantation", Public Health Reports, 128 (4), July 2013, 247-343, https://doi.org/10.1177/003335491312800403.

<sup>&</sup>lt;sup>2</sup> Jones, JM, Kracalik, I, Levi, ME "Assessing Solid Organ Donors and Monitoring Transplant Recipients for Human Immunodeficiency Virus, Hepatitis B Virus, and Hepatitis C Virus Infection — U.S. Public Health Service Guideline, 2020" MMWR Recomm Rep 2020;69 (7-8) available at: https://www.cdc.gov/mmwr/volumes/69/rr/rr6904a1.htm.

<sup>&</sup>lt;sup>3</sup> DL Seem. "PHS Guideline"

#### Do the donor samples have to be stored at the living donor hospital or OPO?

- No, OPTN policy does not require that these specimens be stored on-premises. OPOs and living
  donor recovery hospitals are able to contract with an outside facility for specimen storage,
  including histocompatibility (HLA) labs.
  - Many HLA labs store samples long term. The labs are used to storing specimens for long periods of time and have systems in place to manage it.

#### How should samples be prepared for storage?

- Note: OPTN and CDC do not provide information on sample preparation; policy does not require the specimens be stored on premises, only that the OPO or hospital arrange for storage.
- Consult with your laboratory personnel on those who have experience with the proper preparation for long-term storage. Below are some factors to consider:
  - Samples should be spun down to separate out red blood cells.<sup>4</sup>
  - The specimens can be put into plastic vials with added glycerin⁵ or DMSO⁶ before storing.
  - Laboratories do not need to separate out white blood cells and plasma from serum in order to store samples, but can choose to do so.
  - Laboratories are not required to separate out granulocytes and erythrocytes from peripheral blood mononuclear cells (PBMCs), but may do so.
  - It is essential that the plasma be maintained to test for viral nucleic acids if laboratories choose to separate out white blood cells.<sup>7</sup>

## At what temperature should the donor sample be stored?

- The OPTN and 2020 PHS Guideline do not provide information on optimal storage of serum specimens, but lower-temperature storage (-70 or -80 Celsius) has been shown to improve longterm viral detection for both HIV and HCV.<sup>8,9</sup>
  - The CDC provided additional information on temperature storage in the 2013 PHS
     Guideline: "For archived blood specimens, viral nucleic acid may deteriorate over time

<sup>&</sup>lt;sup>4</sup> Tuck, Melissa K., Daniel W. Chan, David Chia, Andrew K. Godwin, William E. Grizzle, Karl E. Krueger, William Rom, et al. "Standard Operating Procedures for Serum and Plasma Collection: Early Detection Research Network Consensus Statement Standard Operating Procedure Integration Working Group." *Journal of Proteome Research* 8, no. 1 (2008): 113–17. https://doi.org/10.1021/pr800545q.

<sup>&</sup>lt;sup>5</sup> "Best Practices for PBMC Processing," Duke Human Vaccine Institute Immunology, Quality Assessment Cryopreservation Proficiency Testing Program, 2015 ACTG Network Annual Meeting.

https://iqa.center.duke.edu/sites/iqa.center.duke.edu/files/ACTG\_2015\_FINAL\_.pdf

<sup>&</sup>lt;sup>6</sup> Hønge, Bo Langhoff, Mikkel Steen Petersen, Rikke Olesen, Bjarne Kuno Møller, and Christian Erikstrup. "Optimizing Recovery of Frozen Human Peripheral Blood Mononuclear Cells for Flow Cytometry." PLOS ONE 12, no. 11 (2017). https://doi.org/10.1371/journal.pone.0187440.

<sup>&</sup>lt;sup>7</sup> James C. Todd, Sanford, Arthur H., Davidsohn, Israel, Henry, John B. *Clinical diagnosis and management by laboratory methods*, Philadelphia: Saunders, 1979. https://www.ncbi.nlm.nih.gov/nlmcatalog/7901210

<sup>&</sup>lt;sup>8</sup> Halfon, Philippe, Hacène Khiri, Victoria Gerolami, Marc Bourliere, Jean M. Feryn, Pascal Reynier, André Gauthier, Guy Cartouzou, *Impact of various handling and storage conditions on quantitative detection of hepatitis C virus RNA*, Journal of Hepatology, Volume 25, Issue 3 (1996), Pages 307-311, https://doi.org/10.1016/S0168-8278(96)80116-4.

<sup>&</sup>lt;sup>9</sup> Giocchio, Christine C, Xue-Ping Wang, Mark H Kaplan, Gaby Mulligan, Donald Witt, Joseph W Romano, Michael Cronin, and Richard Carroll. "Effects of Specimen Collection, Processing, and Storage Conditions on Stability of Human Immunodeficiency Virus Type I RNA Levels in Plasma." *Journal of Clinical Microbiology* 35, no. 11 (November 1997): 2886–93.

depending on storage conditions. For example, repeated freeze-thaw cycles can cause a moderate reduction in viral nucleic acid levels. Procedures to maximize sample quality include separating specimens that might be used for NAT into multiple aliquots prior to long-term storage, with storage temperature maintained at  $-70^{\circ}$ C or colder. Furthermore, avoiding temperature extremes when archived specimens are shipped for testing inhibits specimen hemolysis, which can result in both false-positive serologic results and false-negative NAT results. Therefore, transporting archived specimens to a testing laboratory on dry ice is a common practice, as well as documenting the specimen quality and condition, with respect to both temperature and hemolysis, upon receipt in the testing laboratory."

The maximum temperature recommended for long-term DNA stability is -27 C, but -80 to -130 C would be more ideal for plasma or serum specimens and is still feasible with a mechanical freezer.<sup>10</sup>

#### **Additional Resources:**

- College of American Pathologists (CAP) provides a <u>database to help find accredited</u> biorepositories.
- <u>CDC Biorepository</u> provides offer sample management expertise to programs, contactable at <u>biorepository@cdc.gov</u>

<sup>10</sup> Rothman, Nathaniel, Jimmie B Vaught, and Marianne K Henderson. Essay. In *Molecular Epidemiology: Principles and Practices*, 23–42. Lyon, France: International Agency for Research on Cancer, World Health Organization, 2011.

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