**OPTN extra vessel sterile rigid container labeling instructions**

**OPTN policy definition of extra vessel:**
Vessels taken during recovery of deceased or living donor organs with the intent to be used in organ transplantation only. Extra vessels are not connected to the organ. Extra vessels are subject to the same member requirements applying to the organ unless otherwise specified.

**OPTN extra vessel labeling instructions:**
OPTN policy requires that extra vessels be packaged in a triple sterile barrier. There must be **two** labels on each extra vessels package. The rigid container holding the vessels and the outermost layer of the triple sterile barrier must **each** have a completed OPTN extra vessel label. These are two **different** labels. The OPTN contractor distributes the two standardized labels that must be used for this purpose.

**Sterile rigid container label:**
The first label must be affixed to the sterile rigid container. The label must be sterile. The required OPTN contractor standardized label to be placed on the sterile rigid container is shown below. It must be completed in a sterile field using a sterile pen. The results and information recorded must reflect the most up to date information available at recovery.

**Step-by-step instructions**

1. Donor ID from DonorNet®
2. Donor ABO as recorded in DonorNet after second user verification. Include subtype if used for allocation.
3. Date that extra vessel was recovered. DonorNet defines recovery date as the date the donor enters the operating room. Date format should be MM-DD-YYY. (e.g. 03-01-2021)
4. This field must be checked as:
   - “**YES**” if any of the following donor infectious disease tests required by OPTN policy using FDA licensed, approved, or cleared tests are positive:
     - i. For HIV: antibody (anti-HIV) donor screening test, antigen/antibody (Ag/Ab) combination test, or ribonucleic acid (RNA) by donor screening or diagnostic nucleic acid test (NAT).
     - ii. For hepatitis B (HBV): surface antigen (HBsAg) donor screening test or DNA by donor screening or diagnostic NAT.
     - iii. For hepatitis C (HCV): antibody donor screening test (anti-HCV) or RNA by donor screening or diagnostic NAT.
   - “**NO**” if all test results that were conducted in the list above have been received and are **negative**.
   - “**PENDING**” if any tests in the list above were ordered, but the results have not yet been received.
5. This field must be checked as:
   - “**YES**” if the required total HBV core antibody (total anti-HBc or total HbcAb) donor screening test result is positive.
   - “**NO**” if the required total anti-HBc or total HbcAb donor screening test result is negative.
   - “**PENDING**” if the required total anti-HBc or total HbcAb donor screening test has not yet been received.
6. This field must be checked as:
   - “**YES**” if the donor has any risk criteria for acute HIV, HBV, or HCV infection, according to the U.S. Public Health Service (PHS) guidelines.
   - “**NO**” if the donor does not have any risk criteria for acute HIV, HBV, or HCV infection, according to the PHS guidelines.

This label is not the source record for results. Any questions regarding results should be reviewed against source documents.