Public Comment Proposal

Extra Vessels: Reducing Reporting Burdens and Clarifying Policies

OPTN/UNOS Operations and Safety Committee

Contents

Executive Summary 1
Is the sponsoring Committee requesting specific feedback or input about the proposal? 2
What problem will this proposal address? 2
Why should you support this proposal? 3
   How was this proposal developed? 4
   How well does this proposal address the problem statement? 7
Which populations are impacted by this proposal? 8
How does this proposal impact the OPTN Strategic Plan? 8
How will the OPTN implement this proposal? 9
How will members implement this proposal? 9
   Transplant Hospitals 9
   OPOs 9
      Will this proposal require members to submit additional data? 9
How will members be evaluated for compliance with this proposal? 9
How will the sponsoring Committee evaluate whether this proposal was successful post implementation? 10
Policy or Bylaws Language 14
Extra Vessels: Reducing Reporting Burdens and Clarifying Policies

Affected Policies: 1.2 (Definitions); 2.7.A (Exceptions to HIV Screening); 2.15.C (Organ Procurement Procedures); 2.15.D (Required Tissue Typing and Blood Type Verification Materials); 2.15.E (Authorization Requirement); 5.4.B (Order of Allocation); 5.5.C (OPO Requirements for Positive HIV Results); 5.9 (Released Organs); 9.8.A (Segmental Transplant and Allocation of Liver Segments); 14.8 (Packaging, Labeling, and Transporting of Living Donor Organs, Vessels, and Tissue Typing Materials); 14.8.A Living Donor Extra Vessel Recovery and Storage; 14.8.B (Recovery and Storage of Vessels from Living Donors); 15.3 (Informed Consent of Transmissible Disease Risk); 15.4.B (Host OPO Requirements for Reporting Post Procurement Discovery of Recipient Disease or Malignancy); 16.(Organ and Vessel Packaging, Labeling, Shipping, and Storage); 16.1 (Packaging and Labeling Requirements for Living Donor Organs and Vessels); 16.2 (Packaging and Labeling Responsibilities); 16.3 (Packaging and Labeling); 16.3.A (Internal Packaging); 16.3.D (Internal Labeling of Vessels Packaged Separately from Other Organs); 16.3.E.i (Disposable Shipping Box); 16.3.Eiii (Cooler); 16.4 (Documentation Accompanying the Organ or Vessel); 16.4.A (Organ Packaging Documentation Requirements); 16.4.B (Vessel Documentation); 16.5 (Verification and Recording of Information before Shipping); 16.6 (Vessel Recovery, Transplant and Storage); 16.6.A (Deceased Donor Vessel Recovery and Transplant Use); 16.6.B (Vessel Storage); 16.6.C (Blood Type Verification Prior to Transplant of Deceased Donor Vessels); 16.6.D (Recovery and Storage of Vessels from Living Donors); 16.6.E (Blood Type Verification Prior to Transplant of Living Donor Vessels)

Sponsoring Committee: Operations and Safety Committee
Public Comment Period: January 22, 2018 – March 23, 2018

Executive Summary

This proposal would change requirements when extra vessels are shared among transplant hospitals. Members would no longer need to submit a justification to the Membership and Professional Standards Committee (MPSC). Instead, they will report sharing to the OPTN Contractor through the existing extra vessels reporting system in UNet™ implemented in August 2015. The justification requirement is no longer needed. Reporting sharing through UNet™ is already occurring and assures tracking capabilities. The justification reviews have not found any associated policy violations. The requirement creates unnecessary burden without benefit for transplant hospitals, the MPSC, and staff. Proposed IT programming will allow OPOs to view extra vessel dispositions from donors that they recovered.

This proposal would change extra vessels policy labeling requirements for infectious disease results by narrowing labeling from “all” to only “HIV, hepatitis B (HBV), and hepatitis C (HCV)” results. This will facilitate aligning test results and names among OPTN Contractor IT systems (e.g. DonorNet®, TransNet™) and the label that currently have inconsistencies. A TransNet barcode will be added to the label to allow scanning and accessing all infectious disease results available in DonorNet.

This proposal will align policy language with the Final Rule indicating that vessels (including extra vessels) are considered part of the organ with which they are recovered and subject to applicable
requirements. Some current policies need clarifications, exclusions, or deletions to fit within the federal regulation logic and framework.

Is the sponsoring Committee requesting specific feedback or input about the proposal?

The Committee is seeking feedback on what additional infectious disease testing is conducted due to donor travel history or other local protocols but is not mandated by national policy (e.g. Strongyloides) so that these tests can be considered as possible optional additions in DonorNet.

What problem will this proposal address?

This project addresses three problem areas.

1. Change extra vessel sharing requirements
2. Change Extra Vessel Label Policy Requirements and Align DonorNet and Extra Vessels Label
3. Align OPTN Extra Vessel Policies with Final Rule

Part One: Change Extra Vessel Sharing Requirements

First, current policy requires receiving transplant hospitals to submit a justification to the MPSC when extra vessels are shared among OPTN transplant programs. The existing requirement, first enacted in 2005, results in the MPSC now reviewing over 50 justifications each year. During 2016-17, 115 justifications were reviewed. No member violations have ever resulted from these reviews. The review is not an efficient use of member, staff, or volunteer time. In August 2015, a comprehensive extra vessel tracking and reporting system was implemented in the OPTN Contractor’s UNet® data collection system as part of the Transplant Information Electronic Data Interchange (TIEDI®). This system provides information needed for monitoring of shared extra vessels. Current policy is outdated, no longer needed, and should be amended. Changing the requirement will reduce both unnecessary burden while maintaining safety through another existing system already being used.

Part Two: Change Extra Vessel Label Policy Requirements and Align DonorNet and Extra Vessels Label

Second, the transplant community has questioned differences between various extra vessel infectious disease reporting mechanisms and requested that they be changed. Current policy requires that all infectious disease testing results be printed on the extra vessels label that is attached to the outermost layer of the triple sterile barrier. With additional testing being completed for emerging or endemic agents (e.g. West Nile Virus, Zika, Chagas), managing the ability to report all infectious disease results on the extra vessels labels becomes complex. Members do not record these extra tests on the infectious diseases tab in DonorNet. Users can type up to three "other" test results in TransNet but at times this is not enough as more than three additional tests are being performed. Currently DonorNet and the extra vessels label are not consistent in either the names of the infectious disease tests nor the test result choices. TransNet, which is now mandatory for OPO use for all labeling and packaging, prints the DonorNet tests that are currently on the extra vessels label. Differences between test names and result options creates confusion. The policy, as well as all tools used to communicate infectious disease results, must be re-examined and updated for clarity and consistency to meet identified concerns.

Part Three: Align OPTN Extra Vessel Policies with Final Rule

Third, the Final Rule states that OPTN organ policies also apply to vessels. Section §121.2 of the Final Rule contains the following definition for an organ: "Organ means a human kidney, liver, heart, lung, pancreas, intestine (including the esophagus, stomach, small and/or large intestine, or any portion of the gastrointestinal tract) or vascularized composite allograft (defined in this section). Blood vessels recovered from an organ donor during the recovery of such organ(s) are considered part of an organ with which they are procured for purposes of this part if the vessels are intended for use in organ transplantation and labeled "For use in organ transplantation only." In addition, §121.7, part (e) (Identification of organ recipient) states “A blood vessel that is considered part of an organ under this part shall be subject to the allocation requirements and policies pertaining to the organ with which the blood
vessel is procured until and unless the transplant center receiving the organ determines that the blood vessel is not needed for the transplantation of that organ.”¹ These clarifications were added when authority to oversee extra vessels used in transplantation were transferred from the Food and Drug Administration (FDA) to Health Resources and Services Administration (HRSA) following a 2006² proposal that was finalized in 2007.³ Using the Final Rule definitions, all “extra vessels” are types of “vessels” and all “vessels” are considered part of “organs” (Figure 1).

Figure 1: Relationship of Organs, Vessels, and Extra Vessels

Policies inconsistently treat vessels as though they are part of the organ with which it was procured. For example, there are organ-specific policies such as documentation of abnormalities that do not need to apply to extra vessels. Various policies have organ requirements and therefore include extra vessels. The citation to extra vessels is not necessary. Examples of these policies include according to this logic already include extra vessels and do not need to specifically call out extra vessels. Some of these policies include some of the packaging and labeling requirements as well as infectious disease reporting. In other cases, specific exclusions or alterations need to be proposed. An example of this type of proposed change includes allowed emergency use of organs when HIV screening has not been completed. While the extra vessels could be used in the original transplant, they could not be stored without the HIV results. Current policy is silent on this but the proposal will provide clarification.

Why should you support this proposal?

Part One: Change Extra Vessel Sharing Requirements

The proposed policy solution will change the reporting requirement for extra vessels sharing. The member receiving the extra vessels will no longer submit a justification to the MPSC when extra vessels are shared. The member sending the extra vessels will be required to report the sharing in the existing TIEDI system within seven days of the sharing. This solution reduces member, staff, and MPSC volunteer burden for a requirement that has not found any policy violations. The functionality for transplant hospitals to report sharing currently exists in the TIEDI system.

Part Two: Change Extra Vessel Label Policy Requirements and Align DonorNet and Extra Vessels Label

The proposed policy solution will also clarify the current policy requirement that “all infectious disease results” must be on the extra vessels label. It will standardize both tests and results that will be required to be on the label yet provide a more flexible and adaptable solution for accessing other infectious disease test results. This will provide the most up to date results as well as better highlight extra vessels that

¹ OPTN Final Rule https://www.ecfr.gov/cgi-bin/text-idx?SID=bb60e0a7222f4086e88c31211f0277d4&mc=true&node=pt42.1.121&rgn=div5
cannot be stored if not used with the organ with which they were sent. This will facilitate better safety practices and help reduce violations.

**Part Three: Align OPTN Extra Vessel Policies with Final Rule**

The third part of the proposal will clarify other OPTN policies to ensure consistency with the Final Rule. The clarifications will reduce existing policy ambiguities raised by staff analysis as well as member questions. Extra vessel questions have resulted in numerous policy interpretation questions.

**How was this proposal developed?**

This proposal was developed in collaboration with multiple internal and external stakeholders.

**Part One: Change Extra Vessel Sharing Requirements**

The MPSC in consultation with UNOS staff requested that the justification required for extra vessels sharing be reconsidered. In the past two years (2016-2017), 115 justifications have been submitted and reviewed for an average of 58 per year. The justification reviews have not found any safety concerns or policy violations. The Operations and Safety Committee agreed that this requirement was no longer needed. They discussed the need to continue to have a tracking mechanism when extra vessels are shared to facilitate timely communications for safety concerns such as infectious diseases. They propose having shared vessels reported through the existing TIEDI system. The requirement would be for the sending hospital to report within seven days as is the existing policy time frame to report use or destruction. The receiving hospital can currently report extra vessels dispositions regardless of whether the sending hospital has reported the dispositions. Members are currently using this function although it is optional and not currently policy-required for sharing.

**Part Two: Change Extra Vessel Label Policy Requirements and Align DonorNet and Extra Vessels Label**

The TransNet Work Group of the Operations and Safety Committee developed recommendations for the proposed policy changes to the extra vessels label that is placed on the outermost layer of the triple sterile barrier. The TransNet Work Group includes representatives from the Organ Procurement Organization, Transplant Administrators, and Transplant Coordinators Committees as well as end users from both OPOs and transplant hospitals. This group reviewed data and considered three options to rectify the issues identified. Currently, the infectious diseases screening page in DonorNet does not record tests that some OPOs perform due to regional concerns (e.g. Strongyloides). The current policy requires that all infectious disease results be on the label and some OPOs have had difficulty because there are only three “other” spots on the current label. The work group identified these issues and considered three options:

1. Keep policy as is and make the polyplastic labels bigger. Print four versus three TransNet labels.
2. Limit label results to policy-required tests (have bar code scan for additional results)
3. Limit label results to HIV, HBV, and HCV test results (have bar code scan for additional results)

The TransNet Work Group recommended option three. The Ad Hoc Disease Transmission Advisory Committee (DTAC) also recommended option three as well. DTAC members stated that other results (e.g. CMV) are not likely to stop use of extra vessels in an emergency although follow up would need to be done afterwards. The Operations and Safety Committee discussed the recommendations and agreed that the proposal would limit extra vessel label results to HIV, HBV, and HCV test results and then a bar code would be added through TransNet labeling to facilitate a scan for all other results. The results will be the most up-to-date results available in DonorNet.

Reasons in support of this recommendation include that the polyplastic label and TransNet programming could be maintained without the need for frequent changes. Having the barcode scan for the most recent results could help promote use of TransNet among transplant hospitals. There will be the ability to better identify those vessels which cannot be stored. It will be possible to add additional tests in DonorNet if desired without making as many changes downstream that take significant time and resources to implement. The results that are most likely to be pending at recovery will then get a scan for the most
recent results. Currently, there is no requirement to update the results on the extra vessels label or from the TransNet label data.

Another change that the Committee is proposing is to eliminate the “unknown” test result option in UNet and then to modify the extra vessels label accordingly to align five test result options between DonorNet and the extra vessels label. These test result options will be positive, negative, indeterminate, pending, and not done. Currently the vessels label only has four options and one of them is “N/A” which is not a DonorNet option. The choices “indeterminate” and “not done” are currently not on the label but will be added. See Table 1 and Figure 1 below.

<table>
<thead>
<tr>
<th>Test Result Options</th>
<th>Positive</th>
<th>Negative</th>
<th>Pending</th>
<th>Not Done</th>
<th>Indeterminate</th>
<th>Unknown</th>
<th>N/A</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Current</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Donor Net</td>
<td>X</td>
<td></td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
</tr>
<tr>
<td>Extra Vessel Label</td>
<td>X</td>
<td></td>
<td>X</td>
<td></td>
<td></td>
<td></td>
<td>X</td>
</tr>
<tr>
<td><strong>Proposed</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Donor Net</td>
<td>X</td>
<td></td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td></td>
</tr>
<tr>
<td>Extra Vessel Label</td>
<td>X</td>
<td></td>
<td>X</td>
<td>X</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Figure 2: Proposed DRAFT Extra Vessels Polyplastic and TransNet Labels

The Committee recognizes the need to access other test results such as CMV, EBV, as well as other discretionary testing done based on local protocols. The TransNet label that is affixed to the extra vessels
label will have a bar code scan that can access all infectious disease results in DonorNet. The Committee requests specific public comment feedback on what other tests are routinely done (e.g. Strongyloides) so that these can be considered for addition in DonorNet.

Part Three: Align OPTN Extra Vessel Policies with Final Rule

UNOS staff analyzed OPTN/UNOS policies that mention extra vessels as well as organ policies that could apply to extra vessels in light of the logic in the Final Rule. This analysis was conducted following extra vessels policy interpretation questions. The Final Rule, as modified in 2007, states that vessels (including extra vessels) are subject to allocation requirements and policies for the organ for which they were procured. Not all policies were written using this logic as many policies were developed before it was written. OPTN/UNOS policies and Final Rule terms differ slightly. When the term “vessels” is used in the Final Rule it means both vessels attached to the organ as well as what the OPTN defines as “extra vessels.” In addition, policies sometimes uses the term “vessels” versus “extra vessels” although the intent is governing extra vessels.

Staff considered three options to remedy the inconsistencies.

1. Create a separate vessels policy
2. Develop exclusionary policy where needed (e.g. Vascularized Composite Allografts or VCAs)
3. Keep status quo, but clarify as needed

Staff originally thought a separate policy might be more user friendly, but after testing, this model realized that developing an exclusionary policy made the most sense. In addition, all references to “vessels” were clarified to reflect the true intent of “extra vessels”.

In the proposed changes, the terms “vessels” or “extra vessels” are removed when they should not be in policy since extra vessels are considered part of the organ with which they were recovered. It is very important to note that the policy requirements still apply to extra vessels. It will be important for the transplant community to include extra vessels, unless specifically excluded, when complying with organ policies.

Staff identified several areas where Committee consultation was needed. These areas were discussed by the Committee, and the following decisions were made:

1. No extra vessels exceptions were needed for:
   - Documenting surgical damage or abnormalities
   - Tissue typing specimen requirements
   - Transportation costs

2. Clarifications were added to several policies. These include:
   - Allowing use of extra vessels in HIV exceptions only for the primary non-kidney transplant
   - Extra vessels must only be recovered as part of an organ recovery and not in isolation
   - Extra vessels must be sent as part of an organ following recovery. Although they can be packaged separately, they must accompany an organ.
   - Release of organs policies apply when the organ that is released back to the OPO included extra vessels as part of the organ. However, once it is determined that the extra vessels are not needed for the original primary intended recipient, they can be shared between transplant hospitals (not released back to the OPO)
   - Existing verification policies were combined and clarified to apply to transplant in secondary recipients or organ modification procedures. The infectious disease verification requirement was changed from “all” to “HIV, HBV, and HCV” results for consistency with other proposed changes and because those are the results affecting whether vessel use is prohibited.
   - When PHS increased risk extra vessels are used in an emergent situation and informed consent could not be obtained before the procedure, then the recipient can be informed after their use and followed with the program’s post-transplant increased risk testing protocol
Further details on the proposed changes are documented and summarized in Appendix A.

**How well does this proposal address the problem statement?**

*Part One: Change Extra Vessel Sharing Requirements*

The MPSC now reviews over 50 extra vessels justifications per year. During 2016-2017, they reviewed 115 justifications with no resulting policy violations.

Data show that between January 1, 2016 and June 30, 2017, 14,381 extra vessels dispositions were reported through the TIEDI reporting system implemented in August 2015. During that same period, 99% of donors with at least one organ reported as sent with vessels in DonorNet. Of the dispositions reported, there were 103 reports of extra vessels being sent to another hospital. This demonstrates high acceptance and use of the reporting system. The proposed change from requiring a justification to requiring reporting within TIEDI allows for timely tracking. To some extent, it is already being done by the transplant community.

Between January 1, 2017 and June 30, 2017, there were 4,775 dispositions reported and 444 (9.3%) were outside of the seven-day reporting requirement. The proposed requirement to report within seven days is the same timeframe for reporting use or destruction. The data suggest that awareness efforts may be needed for timelier reporting. The majority of extra vessels (98.3%) are reported as transplanted or destroyed within the 14-day period from recovery date and there was only one case where the extra vessels disposition was reported outside of the possible maximum window (21 days after recovery).

*Part Two: Change Extra Vessel Label Policy Requirements and Align DonorNet and Extra Vessels Label*

The decision to align test result options was made based on community requests and TransNet work group feedback. The decision to remove “unknown” as an DonorNet option was made based on confusion over the term’s definition and very low current use (used for only 10 donors in 2016) that indicates that this change should not have major impacts.

The decision to limit the extra vessels label to HIV, HBV, and HCV results is based on printing results that are available at the time of label generation and limiting results to those that affect storage requirements.

Of the 4,775 vessels reported between January 1, 2017 and June 30, 2017, 299 tested positive for HCV (antibody or NAT), HBV (surface antigen or NAT), or HIV (antibody, combo antigen/antibody, or NAT) as stated by policy. While 96.3% of these vessels were transplanted or properly disposed (288), 3.7% (n=11) were stored in violation of policy.

Data analyzed at time of TransNet infectious disease result validation show that only 1.1% or less of HIV, HBV, and HCV results are pending compared to over 11% for EBV results and up to 15% for “Other” results. The data indicate that 5.4% of extra vessels are NAT positive indicating likely active viremia as well as a storage prohibition if not used in the intended recipient. Table 2 below shows extra vessel label results.

**Table 2: TransNet Infectious Disease Validations from June 1, 2017 to October 24, 2017**

<table>
<thead>
<tr>
<th>Infectious Disease</th>
<th>Indeterminate</th>
<th>Negative</th>
<th>Not Done</th>
<th>Pending</th>
<th>POSITIVE</th>
<th>Unk.</th>
<th>No Data</th>
<th>Total</th>
<th>% Pending*</th>
<th>% Positive*</th>
</tr>
</thead>
<tbody>
<tr>
<td>Anti-CMV</td>
<td>12</td>
<td>1716</td>
<td>1</td>
<td>38</td>
<td>2731</td>
<td>756</td>
<td>5254</td>
<td>0.8</td>
<td>60.7</td>
<td></td>
</tr>
<tr>
<td>Anti-HCV</td>
<td>4103</td>
<td>39</td>
<td>360</td>
<td>752</td>
<td>5254</td>
<td>0.9</td>
<td>8.0</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>EBV-IgG</td>
<td>19</td>
<td>251</td>
<td>198</td>
<td>479</td>
<td>3546</td>
<td>761</td>
<td>5254</td>
<td>11.2</td>
<td>82.6</td>
<td></td>
</tr>
<tr>
<td>EBV-IgM</td>
<td>14</td>
<td>3103</td>
<td>931</td>
<td>402</td>
<td>41</td>
<td>1</td>
<td>762</td>
<td>11.3</td>
<td>1.2</td>
<td></td>
</tr>
<tr>
<td>HBeAb</td>
<td>4261</td>
<td>3</td>
<td>42</td>
<td>195</td>
<td>1</td>
<td>752</td>
<td>5254</td>
<td>0.9</td>
<td>4.3</td>
<td></td>
</tr>
<tr>
<td>HBeAg</td>
<td>4452</td>
<td>2</td>
<td>34</td>
<td>6</td>
<td>760</td>
<td>5254</td>
<td>0.8</td>
<td>0.1</td>
<td></td>
<td></td>
</tr>
<tr>
<td>HIV I/II</td>
<td>4411</td>
<td>42</td>
<td>38</td>
<td>10</td>
<td>1</td>
<td>753</td>
<td>5254</td>
<td>0.9</td>
<td>0.2</td>
<td></td>
</tr>
<tr>
<td>NAT HBV</td>
<td>1</td>
<td>4433</td>
<td>2</td>
<td>51</td>
<td>14</td>
<td>1</td>
<td>752</td>
<td>11.1</td>
<td>0.3</td>
<td></td>
</tr>
</tbody>
</table>
Part Three: Align OPTN Extra Vessel Policies with Final Rule

The staff analysis included all policies that affect organs or extra vessels. Half of the references identified and evaluated needed some change or further action. See Table 3 below.

Table 3: Staff Analysis of OPTN Policy

<table>
<thead>
<tr>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>No</td>
<td>49</td>
</tr>
<tr>
<td>Yes</td>
<td>39</td>
</tr>
<tr>
<td>Committee consultation needed</td>
<td>6</td>
</tr>
<tr>
<td>Move</td>
<td>5</td>
</tr>
<tr>
<td>Total</td>
<td>99</td>
</tr>
</tbody>
</table>

In addition, there have been over 25 member or staff questions in the past several years resulting in formal analysis for policy interpretation. These data support making changes that are required by the Final Rule and will assist with member implementation.

Which populations are impacted by this proposal?

OPO and transplant hospitals who recover and use extra vessels. All 58 OPOs recover extra vessels. The majority, 156 out of 252 (62%) active transplant hospitals, had at least one extra vessel disposition reported to the OPTN Contractor in 2016.

How does this proposal impact the OPTN Strategic Plan?

1. *Increase the number of transplants*: There is no expected impact to this goal.

2. *Improve equity in access to transplants*: There is no expected impact to this goal.

3. *Improve waitlisted patient, living donor, and transplant recipient outcomes*: There is no expected impact to this goal.

4. *Promote living donor and transplant recipient safety*: The majority of the hours projected by IT will address issue #2 (Change Extra Vessel Label Policy Requirements and Align DonorNet and Extra Vessels Label). This affects both safety and efficiency. Having a scan available through TransNet for all of the most up-to-date infectious disease results for extra vessels will promote transplant recipient safety.
5. **Promote the efficient management of the OPTN:** Reducing unnecessary requirements such as the extra vessels justification will reduce member, staff, and volunteer burden. Clarifying other areas of policy (e.g. infectious disease results on labels) will promote the ability to comply with requirements. Addressing all three issues will improve efficiency.

**How will the OPTN implement this proposal?**

With changes to so many aspects of policy and the procedures they influence, it is likely that an educational effort will be necessary. Instructional Innovations will monitor this proposal as it develops.

This proposal will require the following programming in UNetSM:

1. Retire “unknown” as an option for DonorNet Infectious Disease screening page results. This will cascade to the Deceased Donor Registration (DDR) form and vice versa. No data conversion will be performed for historical records. This is needed to align DonorNet results with proposed revised extra vessels label.

2. Provide test additions to existing list on the Infectious Diseases page in DonorNet pending feedback from public comment.

3. Modify TransNet to new vessels label requirements that will only include HIV, HBV, and HCV results on the printed label but will include a bar-code label that calls back to DonorNet to display all infectious disease results.

4. OPO access to view extra vessels dispositions recorded in TIEDI for extra vessels recovered by the OPO. This is needed for OPOs to comply with infectious disease reporting.

This proposal will also require that the polyplastic extra vessels label be modified. Based on the outcome of the proposal, a draft label will be designed by staff for review by the Operations and Safety and organ Procurement Organization Committees. Once approved, a new label will be ordered and posted as available for purchase on the UNOS store. Members will be provided guidance on when the new label must go into use through the traditional policy notice and OPTN news release.

**How will members implement this proposal?**

**Transplant Hospitals**

Transplant hospitals would have a reduced impact in that they would not have to submit justifications to the MPSC when extra vessels are shared.

**OPOs**

OPOs will have to train staff on changed rules and possible changed data entry in DonorNet. If additional testing results are incorporated into DonorNet, then data vendors will need to be informed to change data mining and exporting practices. OPOs will have to purchase and use the revised extra vessels labels.

**Will this proposal require members to submit additional data?**

This proposal will require that transplant hospitals report sharing of extra vessels within seven days of the sharing. This proposal takes away the burden to submit a justification for sharing so the net effect will be less data submission. This is based on the data principle collection to maintain patient safety where no alternative data source exists.

**How will members be evaluated for compliance with this proposal?**

The proposed language will not require additional routine monitoring of OPTN members. Any data submitted to the OPTN Contractor may be subject to OPTN review, and members are required to provide
documentation as requested. The MPSC will stop reviewing justifications for sharing since they will no longer be required.

How will the sponsoring Committee evaluate whether this proposal was successful post implementation?

This policy will be formally evaluated approximately six months and one year post-implementation. Analyses after one year will be performed at the request of the Committee. The OPTN will monitor the following data to assess the impact of policy change:

1. Trends in the number of extra vessels reported as shared in the Extra Vessel Disposition Reporting Database post policy change vs vessels reported to MPSC pre policy.
2. Trends in the number of patient safety events related to testing and labeling pre vs post policy change.
3. Trends in the number of HCV, HBV, and HIV positive vessels stored in violation, pre vs post policy change.

The policy change, if successful, should not result in significant changes and could capture additional sharing events.
### APPENDIX A: Details on Proposed Policy Changes

<table>
<thead>
<tr>
<th>Affected Policies</th>
<th>Added word &quot;extra&quot; before &quot;vessels&quot; for clarification</th>
<th>Deleted word &quot;vessels&quot; Policy requirement still applies because it applies to the organ</th>
<th>Added an exclusion for extra vessels</th>
<th>Notes</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.2 Definitions</td>
<td></td>
<td></td>
<td></td>
<td>Revised extra vessels and organ definitions for clarity</td>
</tr>
<tr>
<td>2.7.A Exceptions to HIV Screening</td>
<td></td>
<td></td>
<td></td>
<td>Revised so that although extra vessels may be used with organs (except kidney) not yet screened for HIV in medical emergencies that the extra vessels must not be stored, shared, or used in another recipient prior to HIV screening results</td>
</tr>
<tr>
<td>2.15.C Organ Procurement Procedures</td>
<td>Yes</td>
<td></td>
<td></td>
<td>Abnormalities or surgical damage to extra vessels do not have to be documented</td>
</tr>
<tr>
<td>2.15.D Required Tissue Typing and Blood Type Verification Materials</td>
<td>Yes</td>
<td></td>
<td></td>
<td>Extra vessels procured for transplantation are excluded from minimum tissue typing material requirements.</td>
</tr>
<tr>
<td>2.15.E Authorization Requirement</td>
<td></td>
<td></td>
<td></td>
<td>Clarified that extra vessels may only be recovered with at least one organ. Moved deceased donor authorization language from Policy 16 to this Policy 2.15E</td>
</tr>
<tr>
<td>5.4.B Order of Allocation</td>
<td>Yes</td>
<td></td>
<td></td>
<td>Clarified that extra vessels allocated with an organ but not required for its transplant can be shared and is not subject to other organ reallocation requirements.</td>
</tr>
<tr>
<td>5.5.C OPO Requirements for Positive HIV Results</td>
<td>Yes</td>
<td></td>
<td></td>
<td>Clarified that extra vessels recovered with HIV positive kidneys or livers must only be used for transplantation of these organs and must not be stored.</td>
</tr>
<tr>
<td>5.9 Released Organs</td>
<td>Yes</td>
<td></td>
<td></td>
<td>Clarified that if extra vessels are not used for the recipient, then the transplant hospital may use, share, or store vessels</td>
</tr>
<tr>
<td>9.8.A Segmental Transplant and Allocation of Liver Segments</td>
<td>Yes</td>
<td></td>
<td></td>
<td>This policy appears to apply to attached vessels but might be confused with both and is ultimately not needed.</td>
</tr>
<tr>
<td>14.8 Packaging, Labeling, and Transporting of Living Donor Organs, Vessels, and Tissue Typing Materials</td>
<td>Yes</td>
<td>Yes</td>
<td></td>
<td></td>
</tr>
<tr>
<td>14.8.A Living Donor Extra Vessel Recovery and Transplant</td>
<td>Yes</td>
<td></td>
<td></td>
<td>Modified policy title</td>
</tr>
<tr>
<td>Affected Policies</td>
<td>Added word “extra” before “vessels” for clarification</td>
<td>Deleted word “vessels” Policy requirement still applies because it applies to the organ</td>
<td>Added an exclusion for extra vessels</td>
<td>Notes</td>
</tr>
<tr>
<td>------------------</td>
<td>------------------------------------------------------</td>
<td>---------------------------------------------------------------------------------</td>
<td>-------------------------------------</td>
<td>-------</td>
</tr>
<tr>
<td>15.3 Informed Consent of Transmissible Disease Risk</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>15.4.B Host OPO Requirements for Reporting Post Procurement Discovery of Recipient Disease or Malignancy</td>
<td></td>
<td>Yes</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Policy 16: Organ and Vessel Packaging, Labeling, Shipping, and Storage</td>
<td></td>
<td>Yes</td>
<td></td>
<td></td>
</tr>
<tr>
<td>16.1 Packaging and Labeling Requirements for Living Donor Organs and Vessels</td>
<td></td>
<td>Yes</td>
<td>Yes</td>
<td></td>
</tr>
<tr>
<td>16.2 Packaging and Labeling Responsibilities</td>
<td></td>
<td>Yes</td>
<td></td>
<td></td>
</tr>
<tr>
<td>16.3 Packaging and Labeling</td>
<td></td>
<td>Yes</td>
<td>Yes</td>
<td></td>
</tr>
<tr>
<td>16.3.A Internal Packaging</td>
<td></td>
<td>Yes</td>
<td>Yes</td>
<td></td>
</tr>
<tr>
<td>16.3.D Internal Labeling of Vessels Packaged Separately from Other Organs</td>
<td></td>
<td>Yes</td>
<td></td>
<td></td>
</tr>
<tr>
<td>16.3.E.i Disposable Shipping Box</td>
<td></td>
<td>Yes</td>
<td></td>
<td></td>
</tr>
<tr>
<td>16.3.E.iii Cooler</td>
<td></td>
<td>Yes</td>
<td></td>
<td></td>
</tr>
<tr>
<td>16.4 Documentation Accompanying the Organ or Vessel</td>
<td></td>
<td>Yes</td>
<td></td>
<td></td>
</tr>
<tr>
<td>16.4.A Organ Packaging Documentation Requirements</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>16.5 Verification and Recording of Information before Shipping</td>
<td></td>
<td></td>
<td>Yes</td>
<td></td>
</tr>
<tr>
<td>16.6 Vessel Recovery, Transplant, and Storage</td>
<td></td>
<td></td>
<td>Yes</td>
<td></td>
</tr>
<tr>
<td>16.6.A Deceased Donor Vessel Recovery and Transplant</td>
<td></td>
<td></td>
<td>Substantive change: Deleted requirement to submit justification for extra vessel sharing</td>
<td></td>
</tr>
</tbody>
</table>

Substantive change: Added policy requirements when extra vessels from increased risk donors must be used in an emergency and informed consent could not be obtained beforehand. After transplant, the recipient must be informed and followed with increased risk post-transplant testing.

Deleted unenforceable language about packaging in a "timely" fashion.

Clarified "same external transport container with the organ".

Substantive change: Changed label requirement from "all" infectious disease results to results for HIV, HBV, and HCV testing.

Substantive change: Deleted requirement to submit justification for extra vessel sharing.
<table>
<thead>
<tr>
<th>Affected Policies</th>
<th>Added word “extra” before “vessels” for clarification</th>
<th>Deleted word “vessels” Policy requirement still applies because it applies to the organ</th>
<th>Added an exclusion for extra vessels</th>
<th>Notes</th>
</tr>
</thead>
<tbody>
<tr>
<td>16.6.C Blood Type Verification Prior to Transplant of Deceased Donor Vessels</td>
<td>Yes</td>
<td></td>
<td></td>
<td>Combined into Policy 16.6.C</td>
</tr>
<tr>
<td>16.6.D Recovery and Storage of Vessels from Living Donors</td>
<td></td>
<td></td>
<td></td>
<td>Substantive change: Changed verification requirements regarding infectious disease results from &quot;all&quot; to HIV, HBV, and HCV. Combined living and deceased donor policy and clarified circumstances of verification. Added allowed use of TransNet for consistency with Policy 5.8.</td>
</tr>
<tr>
<td>16.6.E Blood Type Verification Prior to Transplant of Living Donor Vessels</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
Policy or Bylaws Language

Proposed new language is underlined (example) and language that is proposed for removal is struck through (example).

1.2 Definitions

Extra vessels

A vessel taken during procurement of deceased or living donor organs with the intent to be used in organ transplantation only, for vasculature reconstruction or modification of a transplanted organ. Vessels directly attached to the transplantable organ are not considered extra vessels. While extra vessels are routinely taken from areas not immediately connected to the transplantable organ, they are subject to the same member requirements applying to the organ unless otherwise specified.

Organ

A human kidney, liver, heart, lung, pancreas, intestine (including the esophagus, stomach, small or large intestine, or any portion of the gastrointestinal tract), or vascularized composite allograft. Blood vessels, including extra vessels, recovered from an organ donor during the recovery of such organ(s) are considered part of an organ with which they are procured for purposes of this part if the vessels are intended for use in organ transplantation and labeled “For use in organ transplantation only.”

2.7.A Exceptions to HIV Screening Requirement

Exceptions to the HIV screening requirement may be made for organs other than kidneys, when, in the medical judgment of the host OPO and recipient transplant hospital or OPO, an extreme medical emergency warrants the transplantation of an organ that has not been tested for HIV.

In this case the host OPO must do both of the following:

1. Provide all available deceased donor medical and social history to the transplant program.
2. Treat the deceased donor as having an increased risk for disease transmission based on current U.S. Public Health Services (PHS) Guideline.

In this case the receiving transplant hospital must:

- Obtain and document informed consent from the potential transplant recipient or the recipient’s authorized agent before transplantation.
- Obtain HIV screening test results prior to storing, sharing, or using extra vessels in another recipient, according to Policy 16.6: Extra Vessel Recovery, Transplant, and Storage.

2.15.C Organ Procurement Procedures

To ensure organ procurement quality, the host OPO must do all of the following:

1. Ensure that the deceased donor receives medications at appropriate times
2. Document in the deceased donor record any medications administered
3. Begin tissue typing and crossmatching as soon as possible
4. Use standard surgical techniques in a sterile environment
5. Maintain flush solutions, additives, and preservation media at appropriate temperatures
6. Document in the deceased donor record, flush solutions and additives with lot numbers, along with organ anatomy, organ flush characteristics, flush solution amount, flush solution type
7. Document any organ abnormalities, and surgical damage, if any for all organs except extra vessels.

2.15.D Required Tissue Typing and Blood Type Verification Materials

The host OPO must establish a written policy with an OPTN member histocompatibility laboratory that includes specific details of the minimum tissue typing material, type of specimen, medium, and shipping requirements for these items. Extra vessels procured for transplantation are excluded from minimum tissue typing material requirements. Table 2-4 shows the requirements for each organ of this type.

Table 2-4: Minimum Typing Materials

<table>
<thead>
<tr>
<th>The host OPO must provide:</th>
<th>For this organ:</th>
</tr>
</thead>
<tbody>
<tr>
<td>One 7 to 10 mL clot red top tube</td>
<td>Any organ</td>
</tr>
<tr>
<td>Two acid-citrate-dextrose (ACD) yellow top tubes</td>
<td>Kidney or pancreas</td>
</tr>
<tr>
<td>If available, one 2 by 4 cm wedge of spleen in culture medium</td>
<td>Kidney or pancreas</td>
</tr>
<tr>
<td>Three to five lymph node samples</td>
<td>Each kidney or pancreas</td>
</tr>
<tr>
<td></td>
<td>Any organ, if the receiving transplant hospital requests and they are available.</td>
</tr>
</tbody>
</table>

The host OPO will provide specimens for tissue typing for all other organs as requested.

2.15.E Deceased Donor Authorization Requirement

Organ recovery teams. The host OPO may only recover organs that they have received authorization to recover. An authorized organ should be recovered if it is transplantable or a transplant recipient is identified for the organ. If an authorized organ is not recovered, the host OPO must document the specific reason for non-recovery.

Extra vessels may only be recovered with at least one organ. To recover and use extra vessels in an organ transplant, the deceased donor authorization forms must include language indicating that the extra vessels will be used for transplant.

This policy does not apply to VCA transplants. Recovery of vascularized composite allografts (VCAs) for transplant must be specifically authorized from individuals authorizing donation whether that be the donor or a surrogate donation decision-maker consistent with applicable state law. The specific authorization for VCA must be documented by the host OPO.

5.4.B Order of Allocation

The process to allocate deceased donor organs occurs with these steps:

1. The match system eliminates candidates who cannot accept the deceased donor based on size or blood type.
2. The match system ranks candidates according to the allocation sequences in the organ allocation policies.
3. OPOs must first offer organs to potential recipients in the order that the potential recipients appear on a match run.
4. If no transplant program on the initial match run accepts the organ, the host OPO may give transplant programs the opportunity to update candidates’ data with the OPTN Contractor. The host OPO must re-execute the match run to allocate the organ.
5. If no transplant program within the DSA or through an approved regional sharing arrangement accepts the organ, the Organ Center will allocate an abdominal organ first regionally and then nationally, according to allocation Policies. The Organ Center will allocate thoracic organs according to Policy 6: Allocation of Hearts and Heart-Lungs and Policy 10: Allocation of Lungs.

6. Extra vessels allocated with an organ but not required for its transplant can be shared according to Policy 16.6.A: Extra Vessel Use and Sharing.

67. Members may export deceased donor organs to hospitals in foreign countries only after offering these organs to all potential recipients on the match run. Members must submit the Organ Export Verification Form to the OPTN Contractor prior to exporting deceased donor organs.

This policy does not apply to VCA transplants; instead, members must allocate VCAs according to Policy 12.2: VCA Allocation.

5.5.C OPO Requirements for Positive HIV Results

If a donor is found to be positive for HIV after any match run has been executed, the host OPO must report the updated information to the OPTN Contractor and do all of the following for each organ being allocated:

1. Stop allocation on the original match run for this donor

2. Re-execute the kidney and liver match runs in order to include only HIV-positive candidates participating in an institutional review board approved research protocol that meets the requirements in the Final Rule regarding the recovery of organs from individuals known to be infected with HIV according to Policy 15.7.A: Requirements for Allocating HIV Positive Deceased Donor Organs

3. Withdraw any pending offers to candidates who are not HIV positive and also participating in an institutional review board approved research protocol that meets the requirements in the OPTN Final Rule according to Policy 15.7.C: Transplant Hospital Requirements for Transplantation of HIV Positive Organs

4. Allocate only kidneys and livers from HIV positive donors. Extra vessels from these donors must only be allocated with the kidneys or liver and must only be used for transplantation of these organs. Members must not store extra vessels from HIV positive donors.

5.9 Released Organs

The transplant surgeon or physician responsible for the care of a candidate will make the final decision whether to transplant the organ.

The transplant program must transplant all accepted, deceased donor organs into the originally designated recipient or release the deceased donor organs back to and notify the host OPO or the OPTN Contractor for further distribution. If a transplant program released an organ, it must explain to the OPTN Contractor the reason for refusing the organ for that candidate. The host OPO must then allocate the organ to other candidates according to the organ-specific policies. The host OPO may delegate this responsibility to the OPTN Contractor or to the OPO serving the candidate transplant program’s DSA.

If extra vessels are not used for the recipient, then the transplant hospital may use, share, or store vessels according to Policy 16: Organ and Extra Vessel Packaging, Labeling, Shipping, and Storage.

9.8.A Segmental Transplant and Allocation of Liver Segments

If a transplant program accepts a liver and performs a segmental transplant, the host OPO must make reasonable attempts to offer the remaining segment according to the adult deceased donor liver match run. If the remaining segment has not been allocated by the time the deceased donor
organ procurement has started, the transplant hospital must offer it to candidates registered with
the transplant program, or any medically appropriate candidate on the waiting list.

The match run will identify a donor’s liver as one with the potential to be split if the donor meets
all the following criteria:

1. Less than 40-years old
2. On a single vasopressor or less
3. Transaminases no greater than three times the normal level
4. Body mass index (BMI) of 28 or less

The deceased donor liver match run will also indicate if potential transplant recipients are willing
to accept a segmental liver transplant.

If the potential transplant recipient that receives the primary whole liver offer ultimately declines
the liver, any subsequent segmental allocation must be relinquished so that the host OPO may
reallocate the whole liver using the liver match run that corresponds to the deceased donor’s age.

The transplant hospital that receives the primary whole liver offer will determine how the liver will
be split and how the vessels are used.

14.8 Packaging, Labeling, and Transporting of Living Donor
Organs, Extra Vessels, and Tissue Typing Materials

Recovery hospitals are responsible for packaging and labeling any living donor organs, or tissue typing
specimens, or vessels that are recovered from living donors according to Policy 16: Organ and Extra
Vessel Packaging, Labeling, Shipping, and Storage when either of the following occurs:

- Living donor organs, or tissue typing specimens, or vessels are recovered and must be transported
  outside the recovery hospital
- A living donor organ, or tissue typing specimens, or vessels require repackaging by a transplant
  hospital for transport outside the transplant hospital

14.8.A Living Donor Extra Vessel Recovery and Transplant Storage

A recovery hospital must only recover extra vessels for transplant if the living donor consents
to the removal of extra vessels for transplant. The extra vessels from a living donor can only
be used for transplant or modification of an organ transplant for the original intended recipient.

14.8.B Living Donors Vessel Storage

Any extra vessels recovered from living donors must be stored according to Policy 16.7: Vessel

15.3 Informed Consent of Transmissible Disease Risk

Transplant programs must obtain specific informed consent before transplant of any organ when any of
the following occurs:

- The donor has a known medical condition that may, in the transplant hospital’s medical judgment, be
  transmissible to the recipient, including HIV.
- The donor meets any of the criteria for increased risk of transmitting HIV, hepatitis B, and hepatitis C
  as specified in the U.S. Public Health Services (PHS) Guideline.
- When a hemodiluted specimen is used for donor HIV, hepatitis B, or hepatitis C screening, according
Exceptions to the informed consent requirement may be made for extra vessels when, in the medical judgment of the transplanting physician, the extra vessels are required for use in an emergency transplant procedure for a recipient other than the original intended recipient. In this case, the transplant hospital must do both of the following post-transplant:

1. Inform the recipient of the use of the extra vessels and the increased risk status
2. Provide follow up to the recipient according to Policy 15.3.B: Donors at Increased Risk for Transmission of Blood-borne Pathogens

Transplant programs must also inform potential candidates of the general risks of potential transmission of malignancies and disease from organ donors, including all of the following information:

1. Deceased donors are evaluated and screened as outlined in Policy 2.3: Evaluating and Screening Potential Deceased Donors.
2. Living Donors are required to undergo screening for the diseases listed in Policy 14.4: Medical Evaluation Requirements for Living Donors.
3. That there is no comprehensive way to screen deceased and living donors for all transmissible diseases.
4. That transmissible diseases and malignancies may be identified after transplant.

The transplant program must do both of the following:

1. Explain these risks and obtain informed consent from the potential candidate or candidate’s agent before transplant.
2. Document consent in the potential candidate’s medical record.

15.4.B Host OPO Requirements for Reporting Post-Procurement Discovery of Recipient Disease or Malignancy

If the host OPO is notified that an organ recipient is suspected to have, is confirmed positive for, or dies from a potential transmissible disease, infection, or malignancy and there is substantial concern that it could be from the transplanted organ, then the host OPO must do all the following:

1. Communicate the suspected donor’s and affected organ recipient’s test results and diagnosis that may be relevant to acute patient care, as soon as possible but no more than 24 hours after receipt, to any transplant program patient safety contacts and tissue banks that received organs, vessels or tissue from the donor. This includes any test results that were not available at the time of procurement or that were performed after procurement. The host OPO must document that this information is shared with all receiving transplant programs and tissue banks.
2. Report the event to the OPTN Improving Patient Safety Portal as soon as possible but no more than 24 hours after notification or receipt of recipient test results or diagnosis.

Policy 16: Organ and Extra Vessel Packaging, Labeling, Shipping, and Storage

16.1 Packaging and Labeling Requirements for Living Donor Organs and Extra Vessels

Living donor recovery hospitals are responsible for packaging, labeling, and transporting living donor organs, vessels, and tissue typing samples according to Policy 16, with these differences:
1. Members are not required to use the OPTN organ tracking system for labeling and packaging living donor organs, vessels, and tissue typing samples.

2. When a member repackages a living donor organ, they are not required to notify the member that originally packaged the organ.

3. In addition to the list of documents in Policy 16.4: Documentation Accompanying the Organ or Extra Vessel, living donor organs must contain the blood type source documents, donor informed consent form, and the complete medical record of the living donor. Extra vessels that are shipped separately from living donor organs must include the same documents as are required for shipping living donor organs.

4. Blood samples and tissue typing materials must contain the donor ID and one of the following identifiers: donor date of birth, donor initials, or a locally assigned unique ID. Each sample must contain the donor’s blood type and subtype, the type of tissue, and the date and time when the sample was obtained. The recovery hospital must document in the donor record all unique identifiers used to label blood samples and tissue typing materials.

5. The recovery hospital will provide specimens for tissue typing if requested. The minimum typing materials for living donor kidneys are: two ACD (yellow top) tubes per kidney.

16.2 Packaging and Labeling Responsibilities

The host OPO or recovery hospital is responsible for packaging and labeling organs, and tissue typing material, and vessels that travel outside the recovery facilities. The host OPO or recovery hospital must make reasonable efforts to package and label organs, and tissue typing specimens, and vessels in a timely fashion.

The host OPO must complete labeling and packaging using the OPTN organ tracking system. The OPO must develop and comply with a written protocol for an alternative labeling and packaging process if, for any temporary reason, the OPTN organ tracking system is not used. This written protocol must fulfill all the requirements in Policy 16 and the host OPO must document the reasons the OPTN organ tracking system was not used.

Transplant hospital staff may not leave the operating room without allowing the host OPO to package and label deceased donor organs, and tissue typing specimens, and vessels as required, or the host OPO will be required to submit a report about the event through the OPTN Improving Patient Safety Portal.

If a transplant hospital repackages an organ for transport, it must package, label, and transport the organ according to the requirements in Policy 16, except that the use of the OPTN organ tracking system is not required. The transplant hospital must immediately notify the host OPO of the repackaging.

16.3 Packaging and Labeling

The host OPO must package all organs, and tissue typing material, and vessels in a sterile environment using universal precautions.

The packaged organs from the deceased or living donor’s surgical back table are to be placed directly into the wet iced shipping container. Proper insulation and temperature controlled packaging including adequate ice or refrigeration must be used to protect the organs during transport. The host OPO may either package extra vessels in the same external transport container with the organ or separate from the organs.
The transplant hospital or OPO must use both internal and external transport containers to package a deceased or living donor organ that travels outside of the facility where the organ is recovered.

16.3.A Internal Packaging

A triple sterile barrier must protect organs and vessels. A rigid container must be used as one of these layers when packaging kidneys, pancreas, and/or extra vessels that are packaged separately from the organs. If the rigid container is sterile, it can serve as one layer of the required triple sterile barrier. The use of a rigid container is optional for all other organs.

16.3.D Internal Labeling of Extra Vessels Packaged Separately from Other Organs

The rigid container holding the extra vessels and the outermost layer of the triple sterile barrier must each have a completed OPTN vessel label. The OPTN Contractor distributes standardized labels that must be used for this purpose. The internal label on the outermost layer of the triple sterile barrier must be completed using the OPTN organ tracking system. The labels must include all of the following information according to Table 16-1 below.

### Table 16-1: Required Information on Internal Labels for Extra Vessels

<table>
<thead>
<tr>
<th>This information must be included:</th>
<th>On the rigid container:</th>
<th>On the outermost layer of the triple sterile barrier:</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. Donor ID</td>
<td>•</td>
<td>•</td>
</tr>
<tr>
<td>2. Donor blood type</td>
<td>•</td>
<td>•</td>
</tr>
<tr>
<td>3. Donor blood subtype, if used for allocation</td>
<td>•</td>
<td>•</td>
</tr>
<tr>
<td>4. Recovery date</td>
<td>•</td>
<td>•</td>
</tr>
<tr>
<td>5. Description of the container contents</td>
<td>•</td>
<td>•</td>
</tr>
<tr>
<td>6. That the extra vessel is for use in organ transplantation only</td>
<td>•</td>
<td>•</td>
</tr>
<tr>
<td>7. All infectious disease donor screening test results for all of the following:</td>
<td></td>
<td>•</td>
</tr>
<tr>
<td>a. HIV antibody (anti-HIV)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>b. HIV antigen/antibody (Ag/Ab) combination test</td>
<td></td>
<td></td>
</tr>
<tr>
<td>c. HIV ribonucleic acid (RNA) donor screening or diagnostic NAT</td>
<td></td>
<td></td>
</tr>
<tr>
<td>d. Hepatitis B core antibody (anti-HBc)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>e. Hepatitis B surface antigen (HBsAg)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>f. Hepatitis B NAT</td>
<td></td>
<td></td>
</tr>
<tr>
<td>g. Hepatitis C antibody (anti-HCV)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>h. Hepatitis C ribonucleic acid (RNA) donor screening or diagnostic NAT</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
This information must be included: | On the rigid container: | On the outermost layer of the triple sterile barrier:
---|---|---
8. Whether the extra vessels are from a donor with a positive result (including NAT) for any of the following:
   - Human Immunodeficiency Virus (HIV), Hepatitis C virus (HCV), or Hepatitis B Virus (HBsAg or NAT)
   - Hepatitis B virus (HBcAb)
   - [●]
9. Whether the extra vessels are from a donor that meets the increased risk for disease transmission criteria in the U.S. Public Health Service (PHS) Guideline
   - [●]  [●]

16.3.E.i Disposable Shipping Box
If organs, or tissue typing materials, or vessels are shipped commercially, they must be transported in a new disposable shipping box. Disposable shipping boxes may not be reused and each box must contain all of the following:

1. A closed plastic liner inside the insulated container to encase the cooling material. The liner must be secured and leak-proof.
2. An inner insulated container, 1.5 inches thick, or a container with an equivalent thermal resistance. The container must have proper insulation and enough cooling material to protect the organs during normal conditions of transport.
3. A water-tight, secured, colored, opaque plastic liner between the outer and inner containers. The liner must be secured and leak-proof.
4. An outer container of corrugated plastic or corrugated cardboard, with at least 200 pounds burst strength, that is coated with a water resistant substance.

16.3.E.iii Cooler
If a member of the organ recovery team is accompanying the organ to the potential recipient’s transplant hospital, the organs and tissue typing material, and vessels may be transported in a cooler. A cooler may be reused only if it is properly cleaned and sanitized and all labels from previous donor organs are removed.

16.4 Documentation Accompanying the Organ or Extra Vessel

16.4.A Organ Packaging Documentation Requirements
Each external deceased and living donor transport container holding an organ must be sent with all of the following source documentation:

1. Blood type
2. Blood subtype, if used for allocation
3. Infectious disease testing results available at the time of organ packaging
The source documentation must be placed in a watertight container in *either* of the following:

- A location specifically designed for documentation
- Between the inner and external transport containers

For deceased donor organs, the host OPO must label the watertight container. This label must be completed using the OPTN organ tracking system. The label must include the donor ID, blood type, and blood subtype if used for allocation.

### 16.4.B Vessel Documentation

If *extra* vessels are not shipped in the same external transport container as the *other* organs, then the separate *extra vessel external transport* container must include the same complete donor documentation *as the organ.*

### 16.5 Verification and Recording of Information before Shipping

Each OPO or recovery hospital must establish and then implement a protocol for verifying the accuracy of organ and vessel packaging labels by an individual other than the individual initially performing the labeling and documentation.

This verification must occur after completing the required labels and documentation for organs and vessels and the host OPO or recovery hospital must document that verification.

The host OPO must use the OPTN organ tracking system to:

1. Record each item placed into the external organ package
2. Report to the OPTN Contractor that the package is ready for tracking

### 16.6 Extra Vessel Recovery, Transplant, and Storage

#### 16.6.A Deceased Donor Extra Vessel Recovery and Transplant Use and Sharing

To recover and use vessels in an organ transplant, the deceased donor authorization forms must include language indicating that the vessels will be used for transplant. The *Extra vessels can* must only be used for organ transplantation or modification of an organ transplant. Transplant hospitals may share deceased donor *extra vessels* with other transplant hospitals. Extra vessels from a living donor may only be used for transplant or modification of an organ transplant for the original intended recipient and must not be shared. If sharing occurs between transplant hospitals, the receiving transplant hospital must submit a detailed explanation to the OPTN Contractor that justifies why sharing occurred. The Membership and Professional Standards Committee (MPSC) will review the explanation. If the receiving transplant hospital later disposes of any vessels, it must notify the OPTN Contractor.

#### 16.6.B Extra Vessel Storage

Transplant hospitals must not store a donor’s extra vessels if the donor has tested positive for *any* of the following:

- HIV by antibody, antigen, or nucleic acid test (NAT)
- Hepatitis B surface antigen (HBsAg)
• Hepatitis B (HBV) by NAT
• Hepatitis C (HCV) by antibody or NAT

Extra vessels from donors that do not test positive for HIV, HBV, or HCV as above may be stored. When a transplant hospital stores extra vessels it must do all of the following:

1. Use stored extra vessels only for organ transplantation
2. Designate at least one person to monitor extra vessel storage, use, destruction, and reporting
3. Package and label extra vessels as required by Policy 16.3: Packaging and Labeling and Policy 16.4: Documentation Accompanying the Organ or Extra Vessel
4. Store extra vessels in a Food and Drug Administration (FDA) approved preservation solution
5. Store extra vessels in a secured refrigerator with a temperature monitor and maintain the temperature no colder than 2 degrees Celsius and no warmer than 8 degrees Celsius
6. Maintain a log of stored extra vessels
7. Maintain all records relating to the monitoring and use of extra vessels
8. Monitor extra vessels daily and log security and refrigerator temperature checks
9. Destroy unused extra vessels within 14 days after the recovery date
10. Report the extra vessel’s use or destruction to the OPTN Contractor within seven days of the transplant hospital’s use or destruction of the extra vessels

16.6.C Blood Type Verification Prior to Transplant of Extra Deceased Donor Vessels

The transplant hospital must verify the blood type, all infectious disease testing results, container contents, date of expiration, and the Donor ID of the vessels with the blood type and all infectious disease testing results of the recipient prior to transplant. These verifications must be documented and maintained in the recipient medical record.

Transplant hospitals must perform a verification prior to transplant of all extra vessels according to Table 16-2 below.

<table>
<thead>
<tr>
<th>For:</th>
<th>The pre-transplant verification must:</th>
</tr>
</thead>
<tbody>
<tr>
<td>Deceased or living donor extra vessels used in the transplantation of the organ with which they were procured</td>
<td>Be completed according to Policy 5.8: Pre-Transplant Verification</td>
</tr>
<tr>
<td>Any of the following:</td>
<td>Verify all of the following prior to transplant:</td>
</tr>
<tr>
<td>• Deceased donor extra vessels used in the modification of an organ transplant</td>
<td>1. Container contents</td>
</tr>
<tr>
<td>• Deceased donor extra vessels used in the transplantation of another organ</td>
<td>2. Date of expiration</td>
</tr>
<tr>
<td>• Living donor extra vessels used for modification of an organ transplant in the original living donor organ recipient</td>
<td>3. Donor ID</td>
</tr>
<tr>
<td></td>
<td>4. Donor blood type and subtype (if used for allocation)</td>
</tr>
<tr>
<td></td>
<td>5. Donor HIV, hepatitis B (HBV), and hepatitis C (HCV) infectious disease testing results</td>
</tr>
<tr>
<td></td>
<td>6. Recipient blood type</td>
</tr>
<tr>
<td></td>
<td>7. Recipient HIV, hepatitis B (HBV), and hepatitis C (HCV) infectious disease testing results</td>
</tr>
</tbody>
</table>
Transplant hospitals may use the OPTN organ tracking system to assist with completion of this verification. The transplant hospital must document that this verification was completed according to the above requirements.

16.6.D Recovery and Storage of Vessels from Living Donors Reporting Requirements for Extra Vessels

A recovery hospital may only recover extra vessels for transplant if the living donor consents to the removal of extra vessels for transplant. The vessels from a living donor can only be used for transplant or modification of an organ transplant for the original intended recipient and may not share them with anybody else. Transplant hospitals must store vessels recovered according to Policy 16.6.B: Vessel Storage.

Transplant hospitals must report to the OPTN Contractor the disposition of all extra vessels, including their use, sharing, or destruction within seven days of use, sharing, or destruction.

16.6.E Blood Type Verification Prior to Transplant of Living Donor Vessels

Prior to transplant, the recovery hospital must verify all of the following:

1. The living donor’s blood type
2. The living donor’s blood subtype, if used for allocation
3. All infectious disease testing results
4. Container contents
5. Date of expiration
6. Donor ID

The transplant hospital must also verify the blood type and subtype of the intended recipient, if used for allocation, and all infectious disease testing results of the recipient prior to transplant. The documentation of these verifications must be maintained in the recipient medical record.

[Subsequent headings and cross-references to headings affected by the re-numbering of this policy will also be changed as necessary.]