MODULE 1: MANAGING BLOOD AND BLOOD PRODUCT TRANSFERS
# Module 1

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Scope

This document provides guidance to health providers on better practice in transfers of blood and blood products around Australia.

Additionally it offers tools that health providers can use to comply with relevant standards and accreditation.

Introduction

Under the National Blood Agreement all Governments are committed to providing an adequate, safe, secure and affordable supply of blood products, services and promoting high quality management of blood products.

This module for Blood and Blood Product Transfers aims to assist health providers in meeting the requirement of the Statement on National Stewardship Expectations for the Supply of Blood and Blood Products. This module will assist health providers to develop a guideline or Memorandum of Understanding (MOU) to facilitate blood and blood product transfer arrangements between providers.

The intention is for this guideline to:

- Help identify transfer options for blood and blood products between health providers, including between public and private.
- Outline how to ensure that acceptable temperature ranges for blood and blood products are maintained during transportation.
- Outline how health providers can ensure blood has been stored and transported correctly before accepting a transfer.
- Provide a uniform process for transferring blood and blood products.
- Foster a culture throughout the blood sector that strives to transfuse all units before expiry, noting that some discards are appropriate but not inevitable.

The term health provider in this document refers to an organisation providing health services to the Australian community and approved by the NBA to receive blood and/or blood products through the NBA supply contracts.

This guideline covers blood and blood products as outlined in the National Safety and Quality Health Service (NSQHS) Standard 7, Blood and Blood Products Standard, including:

<table>
<thead>
<tr>
<th>Fresh blood components</th>
<th>Plasma-derivatives and recombinant products</th>
</tr>
</thead>
<tbody>
<tr>
<td>• red blood cells</td>
<td>• albumin</td>
</tr>
<tr>
<td>• platelets</td>
<td>• immunoglobulins, including immunoglobulin replacement therapy (e.g. IVIg) and hyperimmune globulins</td>
</tr>
<tr>
<td>• clinical fresh frozen plasma</td>
<td>• clotting factors.</td>
</tr>
<tr>
<td>• cryoprecipitate</td>
<td></td>
</tr>
<tr>
<td>• cryodepleted plasma</td>
<td></td>
</tr>
</tbody>
</table>
Preparation

WHY SET UP A TRANSFER NETWORK?

All health providers are required to identify and develop processes that maximise the appropriate use and minimise the wastage of blood and blood products in line with the statement on *National Stewardship Expectations for the Supply of Blood and Blood Products* issued by the Australian Health Ministers.

Implementing a blood and blood product transfer agreement can assist laboratories to:

- enhance the availability of blood and blood product
- manage a limited resource
- reduce unnecessary wastage by transferring blood and blood product to a health provider where it is more likely to be used appropriately.
WHAT DO YOU NEED TO CONSIDER?

IDENTIFY POSSIBLE PARTICIPATING TRANSFER HEALTH PROVIDERS

Consider the following factors:

- Other health providers with whom you could set up a transfer arrangement. This could include the following:
  - health providers you currently have informal arrangements with;
  - health providers located in your local area that you can approach to set up a transfer arrangement;
  - health providers within your organisation located in other suburbs or health networks.

- A larger laboratory or health provider that you could transfer your product to, to enhance the possibility of its use before expiry.
  - Larger health providers service a varied patient group and are more likely to utilise a range of products before expiry.

- If you are a large laboratory or health provider consider receiving transfers from smaller facilities to enhance the possibility product is used before expiry.
  - As a large laboratory or health provider you are more likely to utilise a product before expiry. You could consider reducing your usual order depending on the amount of product you are aware will be transferred in.
  - You will need to continually consider the impact on your inventory levels and whether or not you can use the transferred product.

- If you are a smaller laboratory consider reviewing your current inventory if frequent transfers are required.

- If you have an established hub and spoke arrangement consider including other health providers outside these arrangements or your organisation.
  - For example, a public local health network hub and spoke could include smaller private health providers in the local area.
  - Alternatively, you could consider becoming a hub if you are a large laboratory with a high blood and blood product turnover. This will allow you to manage your own inventory and that of smaller regional facilities with a low turnover. In turn, smaller sites can operate with a lower inventory and still maximise blood use before expiry.

- The proximity of the other health providers to your site.
• Proximity will factor into transport or courier costs and have an effect on validated shipper configuration requirements. Data loggers are currently recommended for all shipments beyond the Blood Service validated transport times when using their shippers.\(^{10}\)

• Great distance does not mean a transfer arrangement is not possible. A number of local health networks have transfer or hub and spoke arrangements between a large metropolitan laboratory and smaller regional and remote sites. Examples include Hunter Area Pathology Service in NSW, BloodMove South Australia and Pathology Queensland. For more information see the NBA website at www.blood.gov.au/case-studies.

• Transfer arrangement between public and private health providers.
  • Transfer arrangements can work with health providers from different organisations, local health networks, pathology organisations and across the public and private health sectors. BloodMove in South Australia is an example where there is a formal arrangement to transfer blood and blood product between public and private health providers.
  • Public health providers who work with a devolved blood budget may wish to discuss the impact of a transfer agreement with the relevant manager of blood budgets within their jurisdiction.
  • Suitable options could include exchange of supply with short for long expiry and using existing courier networks. Any issues that may arise with these (or any other proposed options) should be considered, agreed and documented by all parties.
  • Transfers between National Association of Testing Authorities (NATA) accredited health providers with blood fridges that are compliant with \textit{AS3864 Medical refrigeration equipment – For the storage of blood and blood products}\(^{6}\) are relatively easy to set up and should provide assurance that blood and blood products are maintained within manufacturer’s temperature specifications.

\section*{ACCREDITATION}

Participating health providers are responsible for maintaining the necessary accreditation, standards and legislation (for example as outlined by NATA or Standard 7). This should provide assurance to receiving health providers that any transferred blood and blood products they receive have been stored appropriately. Include a point in your MOU outlining responsibilities of all parties if accreditation requirements are not met.
REVIEW INVENTORY MANAGEMENT PRACTICE AT PARTICIPATING PROVIDERS

It is recommended that health providers review the current inventory management practice with each product at each laboratory or site participating in the MOU. 10 Tips to Help Manage Your Blood Product Inventory located within Managing Blood and Blood Product Inventory can provide practical advice to appropriately manage blood and blood product inventory.

REVIEW TRANSPORT OPTIONS AND PRACTICES AT PARTICIPATING HEALTH PROVIDERS

Review existing blood courier system

Determine if there is an existing courier service you could utilise. If so, investigate the cost to use this service. If not, investigate courier or taxi options and discuss with management to determine the division of costs.

When taking transport into consideration a cost benefit analysis can be undertaken to understand the workload and financial cost of transferring blood and blood products in and out versus the financial cost and loss of a valuable resource. If the cost of transport outweighs the cost of the blood product/s then it may not be feasible to develop regular transfer arrangements in exceptional circumstances.
All signatories on the MOU may invest in a transport option to reduce costs. Examples of transport options include:

- using an existing health provider courier service;
- engaging hire cars, buses or taxis;
- investigating a courier service that specialises in cold product transfer. This may be especially useful for longer distances and flights.

You may consider entering into a Service Level Agreement with your chosen courier company.

**Review existing shipping configuration for the transfer of blood and blood products**

In accordance with the National Pathology Accreditation Advisory Council blood and blood products must be transported in validated shipping containers. It is important to review what shipping configurations and containers, if any, are currently in use with each health provider participating in the MOU. More information on packing and transport shippers can be found under Packing Requirements, section below.

See *Appendix 1: Transfer arrangement checklist* for a summary of items to consider when setting up formal transfer arrangements.

See *Appendix 2: Example Validation Process* for an outline of a possible shipper configuration and validation process.
Implementation

REQUIREMENTS FOR PACKING AND TRANSPORT

Health providers should agree to a validated method for packing and transporting blood and blood products and document in an MOU.

HANDLING OF BLOOD AND BLOOD PRODUCT

Care must be taken when handling blood and blood products. In particular the following steps should be completed:

- minimal physical handling of all blood and blood products must be practiced to ensure those products are kept within their recommended temperature ranges;
- ensure that when you are handling any red blood cell product that it is not exposed to temperatures outside refrigeration specifications for longer than 30 minutes;
- ensure that when you are handling any frozen blood products outside storage conditions that it is kept on dry ice or frozen ballast within a container to prevent temperature changes;
- the blood and blood product bag and/or packaging integrity must be inspected before sending to another health provider;
- management of out of specification consignments to be included in the MOU.

PACKING REQUIREMENTS

Blood and blood products must be packed for transport in accordance with the validated specifications agreed to between participating health providers. If packing materials such as dry ice are used it is recommended that relevant policies and procedures are developed and appropriate training is provided for staff.

When choosing validated packing methods, you may like to consider implementing existing validated shipper configurations. For example an existing validated blood and blood product packing configuration and transport times is detailed in Australian Red Cross Blood Service Receipt and Use of Blood Service Shippers by External Institutions to Transport Blood and Blood Products and Transportation of blood components and fractionated products. The Blood Service recommend the use of data loggers outside their validated transport times with specific packing configurations.
You may wish to validate your own shipper configurations or the shippers and packing configurations used by the Blood Service for extended time frames. Principles and guidelines on validation can be found in the National Association of Testing Authorities Guidelines for the validation and verification of quantitative and qualitative test methods. An example of one method of shipper validation is located at Appendix 2. Validation data must be documented, reproducible and available to NATA on request.

Consideration of the ambient temperature is important when transporting blood and blood products. There are vast temperature differences across Australia that can affect transport shipper validation times and ensuring blood and blood products remain within manufacturer’s temperature specifications.

It is important to consider all temperature conditions and length of time that may be experienced by the blood or blood product you are packing and transporting. Think about where your blood or blood product is being transferred to and all the environments the shipper may experience. The validation process must be repeated for all possible temperatures the blood and blood product may be exposed to during its transport. Some examples of items to consider when transferring blood and blood product include:

- If the product is going by plane:
  - the cargo hold might not be heated resulting in very low temperatures
  - the items might be held in air conditioned storage before or after the flight
  - the items might be left on the tarmac for some time in extreme hot or cold temperatures.
• If the product is travelling by road:
  • it may travel long distances on a truck that could experience extremes of hot or cold temperatures
  • it might be left on a loading dock for an extended period of time before arriving at the storage site.

Consider conducting initial and ongoing seasonal courier temperature audits and ad hoc quality assurance audits as required. If you are transporting blood or blood product outside of a validated timeframe or condition you must ensure manufacturers’ temperature specifications are met. Where a validated shipper has a minimum number of packs specified, ballast must be used to ensure minimum numbers are maintained.

**TRANSPORT REQUIREMENTS**

Blood and blood products must be transported at the temperature range specified in Table 1.

*Table 1: Transport temperature manufacturer’s requirements*

<table>
<thead>
<tr>
<th>Product</th>
<th>Transport temperature range</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Red blood cells&lt;sup&gt;#&lt;/sup&gt;</td>
<td>2-10°C</td>
<td>All blood refrigerators, including theatre and other holding refrigerators, must comply with current AS3864</td>
</tr>
<tr>
<td>Fresh frozen plasma, cryo-precipitate, cryodepleted plasma&lt;sup&gt;#&lt;/sup&gt;</td>
<td>At or below -25°C</td>
<td>Plasma freezers must comply with current AS3864</td>
</tr>
<tr>
<td>Platelets&lt;sup&gt;#&lt;/sup&gt;</td>
<td>20-24°C</td>
<td>Discontinuation of agitation of platelets during transportation should not exceed 24 hours</td>
</tr>
<tr>
<td>Manufactured Products</td>
<td></td>
<td><strong>See relevant manufacturer’s Product Information</strong> Leaflet.</td>
</tr>
</tbody>
</table>

<sup>#</sup>Adapted from the Australian Red Cross Blood Service (2012). *Blood Component Information, circular of information*

It is recommended that a temperature data logger, or other temperature monitoring device, is utilised in the transport of all blood and blood products. Various temperature monitoring devices are available on the market.
TRANSFER PROCEDURE OUTLINE

There are a number of elements to consider when transferring or receiving blood and blood product. Below are suggestions you may wish to include in your MOU.

TRANSFERRING OUT BLOOD AND BLOOD PRODUCT

It is important to allow sufficient time for receiving facilities to utilise blood and blood products before expiry. The recommendations below are given to ensure the receiving site has the highest likelihood to use the product before expiry;

- 7 - 14 days before expiry for red blood cells,
- >5 days before expiry of supplier irradiated blood cells,
- 24 hours, or as short as agreed to with the receiving site, before expiry for platelets,
- 1 - 3 months before expiry for manufactured products.

You may wish to introduce a system to improve utilisation of short expiry products. For example, the sending laboratory should phone the receiving laboratory to see if they can use the product prior to expiry.
The receiving health provider should be notified of an impending delivery within a time frame agreed between sites in the MOU [e.g. 24 hours’ notice]. This notification will allow sufficient time for receiving facilities to adjust their own inventory orders from the supplier. If relevant, outline in your MOU multiple hub and spoke arrangements to ensure that if one health provider is unable to accept a transfer another arrangement is in place.

Details of blood and blood product transfers should be recorded in BloodNet prior to transport for facilities with access to BloodNet. The BloodNet transfer receipt form, or equivalent, should then be printed and added to the shipper for transportation with packing date, time and signature of packer. Health providers without access to BloodNet should complete the transfer form in Appendix 6 to accompany their shipment.

It is important to inspect all blood and blood products for prior to packing. You should consider documentation of the inspection in your MOU. A checklist to check blood and blood products is available for receiving sites in Appendix 6.

Your MOU should include the agreed shipping configurations and container specifications for the varied climates your locations will face. This should assure all participating health providers in the MOU that blood and blood products are maintained within the recommended manufacturer’s temperature specifications.

A record of the storage temperature of the products must be available on request at the facility that shipped the product. NATA accredited health providers may need to provide this documentation during their accreditation process. This documentation will assure all participating health providers are storing blood and blood products within the current AS3864 Medical refrigeration equipment – For the storage of blood and blood products. It will also provide assurance that blood and blood products are maintained within manufacturer’s temperature specifications and prevent discards due to unknown storage conditions.

Consider what documentation will be required with transfers in the MOU. Recommended documentation and information is:

- name of shipping health provider and receiving health provider
- identification of components/products shipped (donor numbers) and description of component/product and total number of units transferred
- date and time transfer entered into BloodNet, where applicable
- date and time packed
- identification of person who packed the shipment
- inspection of product appearance attended
- transfer procedure checklist, see Appendix 4
- shipper Packing Slip for sites without BloodNet, example available in Appendix 7

* The above documentation can be recorded on the printout from the BloodNet transfer episode or on the Blood and Blood Product Transfer Form available in Appendix 6 of this document.
LABELLING OF TRANSPORT SHIPPERS

It is important to label blood and blood product transport shippers appropriately. See Appendix 8: Shipper Label for an example. Consider including the following details as a minimum;

- name and phone number of contact person at the receiving site,
- name and address of intended receiving site,
- dispatch time and date,
- dispatching health provider, staff member name and contact details,
- clear and precise instructions to the courier contractor.

RECEIVING BLOOD AND BLOOD PRODUCT

When receiving blood and blood product transferred from another health provider, consider the following:

- Review your current inventory and routine stock orders to account for expected transfers into your facility. This may require you to reduce your routine order. For example, if you know you are receiving 20 units of red blood cells, reduce your order by 20 units unless you know that more is required that day.

- Document the time and date the transferred blood and blood product was received on Appendix 6: Blood and Blood Product Transfer Form and whether temperature specifications have been met. This will ensure you have completed records for accreditation purposes.

- It is the responsibility of the receiving facility to ensure the blood and blood product has been maintained within the manufacturer’s required temperature range before accepting it into your inventory. If in doubt, the product must be quarantined until storage and transport conditions have been verified. Please refer to the Australian & New Zealand Society of Blood Transfusion Guidelines for pretransfusion laboratory practice.

- It is good practice to inspect the shipper and blood product upon receipt as per Appendix 6: Blood and Blood Product Transfer Form. This ensures the integrity of each unit and that patient safety is maintained.

- Enter the acceptance or non-acceptance of product into your laboratory information system as required. To ensure traceability, it is important that all laboratories with access to BloodNet must enter their transfers and discards.

- You should maintain a record of transferred products for auditing and accreditation purposes. This can be completed through BloodNet or manually if the transferring and/or receiving site does not have access to BloodNet. This will allow you to review the transfer arrangements and track if transferred products are utilised, if discards rates of expired product decrease and so on.
REMOTE SITES WITHOUT A LABORATORY OR BLOOD FRIDGE

For non-laboratory remote sites with a blood fridge consider who is responsible for maintaining the fridge. The receipting site is responsible for ensuring that all product received has been kept within manufacturer’s temperature specifications and will need to include access to temperature records in the MOU. You may also want to consider including any maintenance records. Alternatively, you may choose to opt for a signed declaration that all blood and blood product has been maintained within manufacturer’s temperature specifications, by the transferring out site.

For remote sites or facilities without a laboratory you may like to include the following additional documentation with the transferred blood and blood products as an assurance for the receiving facility that blood and blood product have been stored according to manufacturers’ temperature requirements;

- completed paperwork outlining the daily storage temperature checks of the blood fridge or storage area, see Appendix 5: Blood Fridge Maintenance Record;
- a photocopy of the objective graph recorder from the blood fridge demonstrating the temperature range of the place of storage; or
- information from the health provider responsible for maintaining the blood fridge, for example temperature graphs, maintenance records. You may be satisfied with a declaration from the sending health provider.

For remote sites without a blood fridge consider how blood and blood products are stored while onsite and include specific storage methods in your MOU. Examples of inclusions include, but are not limited to;

- the product must remain in a sealed shipper until used,
- the product must be accompanied by a data logger or temperature indicator,
- transfer of patient specific product only for immediate use.

For remote sites without a blood fridge, storage and transport requirements and documentation must be detailed in the MOU.

Staff at the remote site should undertake the BloodSafe eLearning “Transporting Blood” module. The receiving laboratory could also consider undertaking additional training for ward/nursing staff at the remote site.
## APPENDIX 1: TRANSFER ESTABLISHMENT AND REVIEW CHECKLIST

<table>
<thead>
<tr>
<th>PROCEDURE</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Identify sites you could transfer to/from.</td>
<td></td>
</tr>
<tr>
<td>Review what blood and blood product you could transfer in/out.</td>
<td></td>
</tr>
<tr>
<td>Review courier or transport possibilities.</td>
<td></td>
</tr>
<tr>
<td>Contact potential participating sites to gauge interest.</td>
<td></td>
</tr>
<tr>
<td>Set up initial meeting with managers and scientists in charge to discuss transferring options.</td>
<td></td>
</tr>
<tr>
<td>Go through the MOU to make it specific for your situation.</td>
<td></td>
</tr>
<tr>
<td>Agree to storage, transport, package, documentation and training requirements.</td>
<td></td>
</tr>
<tr>
<td>Outline acceptable days until expiry that units should be identified for transfer.</td>
<td></td>
</tr>
<tr>
<td>Sign off on agreed MOU.</td>
<td></td>
</tr>
<tr>
<td>Educate staff on arrangements.</td>
<td></td>
</tr>
<tr>
<td>Set start date for transfer arrangement to commence.</td>
<td></td>
</tr>
<tr>
<td>Set date to meet and review transfer arrangement.</td>
<td></td>
</tr>
<tr>
<td>Review MOU arrangement to ensure working for all health providers involved.</td>
<td></td>
</tr>
</tbody>
</table>
APPENDIX 2: VALIDATION PLAN

1. PURPOSE / SCOPE

This document describes the validation of:

- Shipper [enter shipper name] for the use in the inter-hospital/laboratory transport of [Enter components as required].

Responsibilities

<table>
<thead>
<tr>
<th>Position</th>
<th>Responsibility</th>
</tr>
</thead>
<tbody>
<tr>
<td>Senior Scientist</td>
<td>Design validation; analyse results; prepare the report; perform the validation; compile results</td>
</tr>
<tr>
<td>Quality Manager</td>
<td>Authorise validation and approve for implementation</td>
</tr>
</tbody>
</table>

2. REFERENCES

A reference you may like to review is the Council of Europe’s "Guide to the Preparation, Use and Quality Assurance of Blood Components", 16th edition.¹⁸

[List references you outline in this document]

3. MATERIEL (add/delete below as required)

3.1 3x Shipper [enter shipper name].

3.2 Room temperature coolant packs (conditioned at +20 to +24°C for 24hrs prior to use).

3.3 Chilled coolant packs (conditioned at +2 to +6°C for 24hrs prior to use).

3.4 Frozen coolant packs (conditioned at approximately -19°C for 24hrs prior to use).

3.5 Tamper evident labels.

3.6 Cardboard dividers.

3.7 Expired red cells with defaced label "Research Only" or empty dummy packs filled with 275ml saline.

4. EQUIPMENT

Temperature Data Loggers (TDL): [Enter Company Name], [Enter Model No].

<table>
<thead>
<tr>
<th>Temperature Data Logger</th>
<th>Serial Number</th>
<th>Asset Number</th>
</tr>
</thead>
<tbody>
<tr>
<td>TDL1</td>
<td></td>
<td></td>
</tr>
<tr>
<td>TDL2</td>
<td></td>
<td></td>
</tr>
<tr>
<td>TDL3</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
Note: Performance Qualifications for the data loggers have been included in Attachment [X].

Cool Room [Enter name e.g. CR001]
Incubator [Enter name e.g. CR001]

Note: Records for Cool Room [CR001] and Incubator [I001] are located in [Enter location e.g. Engineering Department]

5. ACCEPTANCE CRITERIA

5.1. Temperature maximum does not exceed 10 °C.

5.2. Temperature minimum does not fall below 2 °C.

5.3. There should not be a temperature range difference of ±1 °C between the lowest and highest values for the maximum temperature recorded for each of the data loggers and each of the replications when determining maximum transport time.

5.4. There should not be a temperature range difference of ±1 °C between the lowest and highest values for the minimum temperature recorded for each of the data loggers and each of the replications when determining minimum transport time.

6. PROCEDURE (add/delete below as required)

6.1. Description / Background Information

6.1.1. This laboratory will be validating/revalidating the [enter shipper name] for the transport of red blood cells [or other component] between the following health providers [insert names].

6.1.2. Routine transfer of blood components is undertaken by [enter name]

6.1.3. A review of Bureau of Meteorology for local climatic conditions indicates that minimum environment temperatures do not drop below [enter minimum temperature e.g. 10˚C].

6.1.4. Validation time was set at [x] hours as a review of transport arrangements and non-compliance reports indicate that this would be the worst case scenario the laboratory would experience in the transfer of components.

6.2. Key Variables (add/delete below as required)

6.2.1. Staff will be accessing cool room during low temperature qualification period.

6.2.2. Building air conditioning is switched off from [enter time e.g. 10pm] to [enter time e.g. 6am] during ambient temperature qualification period.
6.3. **Samples** (add/delete below as required, set temperature levels to represent your requirements)

6.3.1. Data logging sample rate set at [x] minute intervals.

6.3.2. Sampling was undertaken over [x] hours for [10 °C - 14°C] low temperature qualification period.

6.3.3. Sampling was undertaken over [x] hours for [20 °C - 24 °C] ambient temperature qualification period.

6.3.4. Sampling was undertaken over [x] hours for [32 °C - 42 °C] high temperature qualification period.

6.4. **Data logger Parameter Settings**

6.4.1. Data logger delay setting set to 15 minutes to allow equilibration.

6.4.2. Data logging sample rate set at [x] minute intervals.

6.4.3. Data logger sample points set at [number of readings to reach 30 hours].

6.5. **Participating centres & personnel**

[Enter Health Provider/Laboratory Name/s], [Enter Location], Senior Scientist, Quality Manager

6.6. **Packing Configuration**

Packing configuration as outlined in Figure 1 below. For each configuration you will need to determine the minimum and maximum number of packs allowed for each configuration to be validated.

*Figure 1 adapted from the Australian Red Cross Blood Service *Receipt and Use of Blood Service Shippers by External Institutions to Transport Blood and Blood Products* Red Cell Configuration R1.
6.7. Temperature Data Loggers and Placement

6.7.1. Data loggers equilibrated to required temperature e.g. 10 °C - 14 °C, 20 °C - 24 °C, 32 °C and 52 °C.

6.7.2. Place one logger in-between the red cells or dummy packs.

6.7.3. Place one logger above the red cells or dummy packs.

6.7.4. Attach one logger to the outside of the shipper to measure the environment temperature.

6.7.5. Refer Figure 1 above.

6.8. Temperature Validations

6.8.1. Shipper packed in accordance with Figure 1 was placed in Cool Room CR001 for [as per 6.1.4] for minimum temperature validation. This was repeated on three separate occasions.

6.8.2. Shipper packed in accordance with Figure 1 was placed in Blood Bank Laboratory for [as per 6.1.4] for ambient temperature validation. This was repeated on three separate occasions.

6.8.3. Shipper packed in accordance with Figure 1 was placed in Incubator I001 set at 32 °C for [as per 6.1.4] and then at 42 °C for maximum temperature validations. This was repeated on three separate occasions.

7. RESULTS

7.1. Data logger Performance Qualification

Performance testing was undertaken and completed by [Enter detail e.g. Engineering Department] on [Enter date]. Results of performance testing against a reference thermometer are outlined in the Table 1 below.

Table 1 Reference thermometer performance testing

<table>
<thead>
<tr>
<th>Temperature Data Logger</th>
<th>Recorded Temp °C</th>
<th>Ref Thermometer Recorded Temp °C</th>
<th>Difference °C</th>
</tr>
</thead>
<tbody>
<tr>
<td>TLD1</td>
<td>23.6</td>
<td>23.5</td>
<td>- 0.1</td>
</tr>
<tr>
<td>TLD2</td>
<td>23.6</td>
<td>23.6</td>
<td>0.0</td>
</tr>
<tr>
<td>TLD3</td>
<td>23.6</td>
<td>23.5</td>
<td>- 0.1</td>
</tr>
</tbody>
</table>

Note: Reference Thermometer [Enter Serial No] records can be obtained from [Enter Engineering Department]
7.2. **Minimum Temperature Validation**

7.2.1. Raw data of data logger download is included in Attachment 1.

7.2.2. Summary of results of data logger mapping is in **Table 2** below:

<table>
<thead>
<tr>
<th>Position</th>
<th>Date</th>
<th>Data Logger</th>
<th>Minimum</th>
<th>Maximum</th>
</tr>
</thead>
<tbody>
<tr>
<td>Position 1</td>
<td>[Enter date]</td>
<td>TDL1</td>
<td>[5.7] °C</td>
<td>[7.9] °C</td>
</tr>
<tr>
<td>Position 1</td>
<td>[Enter date]</td>
<td>TDL1</td>
<td>[5.2] °C</td>
<td>[7.2] °C</td>
</tr>
</tbody>
</table>

[Enter other tables as required]

**8. DISCUSSION AND RECOMMENDATIONS**

The performance testing of the four data loggers was undertaken and completed by [Enter name e.g. Engineering Department], an ISO9000 accredited facility, on [enter date]. The results against a reference thermometer showed that no data logger had a variance greater than ± [enter variation e.g. 0.1] °C.

The minimum temperature validation occurred on three separate occasions over a 7 day period from [enter date] to [enter date]. Cool room CR001 decommissioned for maintenance was recommissioned and set to 10 °C - 14 °C for this validation study. The packing configuration and data logger placement is outlined in Figure 1.

The results show that the shipper stored at 10 °C - 14 °C for [x] hours did not drop below [x] °C for the validation period. The variation of minimum temperature across the three validations for each of the data loggers is [0.8] °C within the allowable ±1°C acceptance criteria. The results show that the shipper stored at 10 °C - 14 °C for [x] hours did not exceed 10 °C until [x] hours. The maximum temperature variation across the three validations for each of the data loggers is [enter variation e.g. 1.1] °C and is within the allowable ±1°C acceptance criteria.

[Discuss ambient temperature validation]

[Discuss maximum temperature validation]

It is recommended that Shipper [Enter name] is suitable for the transport of red cells as inter-hospital/laboratory for up to [x] hours. If the transport is expected to exceed [x] hours or if non-contracted transport such as a taxi is required then consignments should include a data logger as part of the packing configuration, to be positioned next to the red cell packs.
9. APPENDICES

Attachment [1]: Raw data download of [data logger] for minimum temperature validations.

Attachment [2]: Cool Room Temperature Map

Attachment [3]: [other documents as required]

10. APPROVALS

<table>
<thead>
<tr>
<th>Name</th>
<th>Signature</th>
<th>Date</th>
</tr>
</thead>
<tbody>
<tr>
<td>Report prepared by:</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Quality Manager Approval</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

End of document

Repeat the process below for each configuration, product and possible temperature exposure range, for example: 0 °C to 4 °C, 4 °C to 24 °C, 24 °C to 40 °C, 40 °C to 52 °C
APPENDIX 3: EXAMPLE MEMORANDUM OF UNDERSTANDING (MOU):

MEMORANDUM OF UNDERSTANDING

for the
Transfer of blood and blood products between the below listed health providers

<ENTER HOSPITAL OR PATHOLOGY SERVICE (PROVIDER)>

<ENTER HOSPITAL OR PATHOLOGY SERVICE (PROVIDER)>

<ENTER HOSPITAL OR PATHOLOGY SERVICE (PROVIDER)>

<ENTER HOSPITAL OR PATHOLOGY SERVICE (PROVIDER)>

<ENTER HOSPITAL OR PATHOLOGY SERVICE (PROVIDER)>

<ENTER HOSPITAL OR PATHOLOGY SERVICE (PROVIDER)>

<ENTER HOSPITAL OR PATHOLOGY SERVICE (PROVIDER)>

<ENTER HOSPITAL OR PATHOLOGY SERVICE (PROVIDER)>

<ENTER HOSPITAL OR PATHOLOGY SERVICE (PROVIDER)>

<ENTER HOSPITAL OR PATHOLOGY SERVICE (PROVIDER)>
1. PARTICIPATING HOSPITALS OR PATHOLOGY SERVICES

The Memorandum of Understanding (MOU) is endorsed by the [e.g. Senior Haematologist/ Senior Scientist/ Laboratory Manager] from each participating facility. The signatories agree to abide by the contents of this MOU.

<Enter Hospital or Pathology Service (Facility)>
Signature__________________________
Name______________________________
Position___________________________
Date_______________________________

<Enter Hospital or Pathology Service (Facility)>
Signature__________________________
Name______________________________
Position___________________________
Date_______________________________

<Enter Hospital or Pathology Service (Facility)>
Signature__________________________
Name______________________________
Position___________________________
Date_______________________________

<Enter Hospital or Pathology Service (Facility)>
Signature__________________________
Name______________________________
Position___________________________
Date_______________________________
2. CONTACTS

<ENTER HOSPITAL OR PATHOLOGY SERVICE (FACILITY)>
List names, position, contact
List names, position, contact

<ENTER HOSPITAL OR PATHOLOGY SERVICE (FACILITY)>
List names, position, contact
List names, position, contact

<ENTER HOSPITAL OR PATHOLOGY SERVICE (FACILITY)>
List names, position, contact
List names, position, contact

<ENTER HOSPITAL OR PATHOLOGY SERVICE (FACILITY)>
List names, position, contact
List names, position, contact

<ENTER HOSPITAL OR PATHOLOGY SERVICE (FACILITY)>
List names, position, contact
List names, position, contact

<ENTER HOSPITAL OR PATHOLOGY SERVICE (FACILITY)>
List names, position, contact
List names, position, contact

<ENTER HOSPITAL OR PATHOLOGY SERVICE (FACILITY)>
List names, position, contact
List names, position, contact
3. PURPOSE

The purpose of this Memorandum of Understanding (MOU) is to establish cooperation between the above signed health providers for facilitating blood and blood product transfer arrangements between identified facilities. The MOU relates to the Managing Blood and Blood Product Transfers.

The intention for this MOU is to:

- Assist in the reduction of blood and blood product wastage due to expiry or non-use through the transfer of blood and blood products before expiry to enhance the likelihood of usability.
- To provide a uniform process for the transfer of blood and blood products between the participating facilities.
- To ensure that acceptable temperature ranges for blood and blood products are maintained and are demonstrable during storage and transportation.
- That AS3864 compliant blood refrigerators are used for the storage of blood.
- To provide a uniform process for tracking transferred blood and blood products.

4. COORDINATION

The original document and technical and administrative coordination of this MOU will reside with <enter facility name and contact details>. The coordinator will be responsible for the MOU and will communicate with all participating health providers on the activities conducted and information related to the MOU.

5. DEFINITIONS

Sending Health Provider: the health provider that is transferring blood and blood product out of their site.

Receiving Health Provider: the health provider that has agreed to receive the blood and blood product transfers into their site.

Blood product approaching expiry: any product shipped should not have less than the following remaining of the shelf life, unless specifically agreed to by participating health providers in this MOU or in special situations;

- 7-14 days for red blood cells,
- > 5 days before expiry for irradiated blood cells,
- 24 hours or as short as agreed to with the receiving site before expiry for platelets,
- 1-3 months before expiry for manufactured blood products.
6. MEMORANDUM OF UNDERSTANDING REVIEW

<Identify the MOU review responsibilities and timeframe>

For example:

- Review timeframe is every two years,
- Responsibilities include a review of:
  - updated accreditation documents and Australian Standards,
  - MOU participant inventory holdings and blood and blood product usage patterns,
  - inclusion of additional health providers,
  - <list review responsibilities as agreed>.

7. IMPLEMENTATION

Roles and Responsibilities of participating health providers

7.1. Responsibilities for all MOU Participants

Participating health providers are responsible for following the guidance outlined in *Blood and Blood Product Transfers* including the following:

- Maintaining standards and accreditation, where appropriate.
- Meeting all necessary standards and legislation for the storage, handling and transport of blood and blood products as outlined in *Managing Blood and Blood Product Transfers*.
- Participating health providers will ensure that blood components are handled, stored, distributed and transported in a manner that prevents damage, limits deterioration, and meets required standards.
- <Enter additional responsibilities agreed by the participating health providers>

7.2. Sending Health Provider

The sending health provider must: <Identify sending site responsibilities>

For example:

- Contact receiving provider for approval prior to transfer, minimum timeline agreed to is <enter agreed minimum time> hours before arrival of transfer.
- Ensure blood and blood products must have the minimum agreed specified time to expiry as per Section 5. Definitions, unless explicit agreement is acknowledged from receiving site
- Enter transfer into BloodNet (where applicable).
- Enter transfer into your Laboratory Information System (LIS) (where applicable), or manually log where no laboratory is onsite.
Visually inspect all products prior to transferring.

Comply with agreed packing and shipping configuration, specifically:

<enter agreed validated packing configuration>.

Include the Transfer Checklist with either the transfer receipt from BloodNet, OR the Blood and Blood Product Transfer Form (Appendix 6).

For sites without a laboratory include the following documentation as agreed;

- completed Blood Fridge Maintenance Record form, OR
- completed paperwork outlining the daily storage temperature checks of the blood fridge or storage area, AND
- a photocopy of the objective graph recorder from the blood fridge, OR
- information from the health provider responsible for maintain the blood fridge with temperature records, maintenance records or signed declaration.

7.3. Receiving Health Provider

The receiving health provider must: <Identify receiving site responsibilities>

For example:

- agree to receive the transferred blood or blood product;
- review your current inventory and routine stock orders to account for expected transfers in;
- inspect all packaging of received blood and blood product and do not accept the transfer unless it is intact and packed according to agreed validated shipper configuration;
- document the time and date the product was received;
- document evidence that manufacturer’s temperature specifications have been maintained. If in doubt, quarantine all products until storage, packing and transport conditions can be verified;
- check temperature data logger, if used;
- visually inspect all blood and blood products received;
- record transferred in units into your LIS;
- complete all other documentation as required e.g. group check if transferred from a non-laboratory setting;
  - maintain record of product received by transfer.
8. TRANSPORT LOGISTICS

<Enter transport logistics as agreed by the participating health providers>

For example:

- The agreed packing configuration is as per the Blood Service Validated Shippers.

Refer to:

- *Receipt and Use of Blood Service Shippers by External Institutions to Transport Blood and Blood Products,*
- *Transport Times,*
- *Transportation of blood components and fractionated products.* OR

- The agreed packing configuration is <enter agreed validated packing configuration>.
- Data loggers or temperature monitoring must be used when transport is outside validated shipper times.
- The agreed transport method is:
- <Enter agreement for courier/transport method>,
- <Enter agreement for courier/transport cost>.
APPENDIX 4: TRANSFER PROCEDURE CHECKLIST

<table>
<thead>
<tr>
<th>PROCEDURE</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>1. Routine check for products close to expiry.</td>
<td></td>
</tr>
<tr>
<td>2. Identify and contact possible receiving health provider/s to negotiate</td>
<td></td>
</tr>
<tr>
<td>transfer of close to expiry product.</td>
<td></td>
</tr>
<tr>
<td>3. Record blood or blood product transfer in BloodNet. Print transfer</td>
<td></td>
</tr>
<tr>
<td>record and include with the product.</td>
<td></td>
</tr>
<tr>
<td>4. Record transfer information on Blood and Blood Product Transfer Form**</td>
<td></td>
</tr>
<tr>
<td>** only if BloodNet is not available</td>
<td></td>
</tr>
<tr>
<td>5. Record blood or blood product transfer in the Laboratory Information</td>
<td></td>
</tr>
<tr>
<td>System (LIS)*.</td>
<td></td>
</tr>
<tr>
<td>*If your LIS is interfaced with BloodNet then record entry is not</td>
<td></td>
</tr>
<tr>
<td>required in BloodNet.</td>
<td></td>
</tr>
<tr>
<td>6. Inspection of blood and blood product for abnormal appearance,</td>
<td></td>
</tr>
<tr>
<td>package integrity, leakage and expiry date.</td>
<td></td>
</tr>
<tr>
<td>7. Check blood or blood product is packed according to MOU agreement,</td>
<td></td>
</tr>
<tr>
<td>include date and time packed on transfer form.</td>
<td></td>
</tr>
<tr>
<td>8. Copy of the Blood Fridge Maintenance Record form or temperature graph</td>
<td></td>
</tr>
<tr>
<td>included as per MOU (to be sent when required/requested).</td>
<td></td>
</tr>
<tr>
<td>10. Notify recipient health provider by telephone or email of impending</td>
<td></td>
</tr>
<tr>
<td>delivery.</td>
<td></td>
</tr>
</tbody>
</table>

Checklist completed by:

Name:______________________________ Signature:___________________________
Date and Time:__/__/____
APPENDIX 5: BLOOD FRIDGE MAINTENANCE RECORD

Blood fridge maintenance procedures must be performed according to the schedule above. Record all results on the form.

<table>
<thead>
<tr>
<th>Fridge location / identification</th>
<th>Hospital Name</th>
<th>Asset No.</th>
<th>Month</th>
<th>Year</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**Daily maintenance checks**
- Record blood fridge temperature from digital display or internal thermometer (Acceptable Range 2-6°C) (Record temperature)
- Check temperature recording chart for out of range spikes (√)
- Document reasons for spikes on temperature chart and in Problem Log below.
- Check fridge for blood that can be returned to the transfusion provider.
- Segregate blood and contact transfusion service for return (including advice on packing and transportation of blood as required).
- Initials of staff member performing check (Initials)

| Date | Week | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 | 9 | 10 | 11 | 12 | 13 | 14 | 15 | 16 | 17 | 18 | 19 | 20 | 21 | 22 | 23 | 24 | 25 | 26 | 27 | 28 | 29 | 30 | 31 |
|------|------|---|---|---|---|---|---|---|---|---|----|----|----|----|----|----|----|----|----|----|----|----|----|----|----|----|----|----|----|----|----|----|----|----|----|----|----|----|----|----|----|----|----|----|----|----|

**Maintenance checks** (define period with a minimum requirement as outlined in AS3864) (See procedure on right)
- Change temperature chart (√)
- Audible (√)
- Visible (√)
- Power Loss (√)
- Remote Alarm (√)
- Initials of staff member performing check (Initials)

**BLOOD FRIDGE MAINTENANCE PROCEDURES**

**TEMPERATURE CHART**
- Weekly temperature chart (circular type): Open cover, remove old chart. Check chart for conformance during previous 7 days, date and sign. Date new chart, identify fridge/hospital, sign and place on recorder. Ensure that day and time are coerced with pen recorder position. Adjust if necessary. Ensure pen tip is a touching chart. Close and lock cover. Other types of temperature chart - Check chart conformance daily and change as required.

**ALARM TEST**
- Audible: Push test button and ensure audible alarm (e.g., beep or siren) is functioning. Visible: Push test button and ensure visible alarm (e.g., flashing light) is functioning.
- Power loss: Turn power off at wall switch or main switch if accessible. Ensure alarms function.
- Remote alarm: Push test button and ensure remote alarm system(s) is functioning.

**ALARM HIGH AND LOW TEMPERATURE ACTIVATION CHECKS**
- Some newer fridges have an automated process for these tests. Older fridges require a manual checking process as detailed below. Please refer to the fridge manufacturer’s manual on how to do it.
- Please handle probes carefully for the following checks (once completed carefully replace probes):
  - High temperature alarm: Remove probe from container. Insert into a small container of room temperature water. Ensure an alarm is activated as temperature rises above 5.5°C.
  - Low temperature alarm: Remove probe from container. Insert into a small container of ice water. Ensure an alarm is activated as temperature falls below 2.5°C.

**BATTERY BACKUP CHECK** – refer to manufacturer’s instructions

**PROBLEM LOG** (Record problems, dates and corrective actions taken (continue on reverse if required)

Copy Sent To: Partner
Transfusion Laboratory: Name: Sign: Date:
Reviewed by Laboratory: Name: Sign: Date:
APPENDIX 6: BLOOD AND BLOOD PRODUCT TRANSFER FORM

For facilities without BloodNet access only

<table>
<thead>
<tr>
<th>From:</th>
<th>Contact phone:</th>
</tr>
</thead>
<tbody>
<tr>
<td>To:</td>
<td>Date: Time:</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Donation Number (if applicable)</th>
<th>Blood Group (if applicable)</th>
<th>Comments (These include any temperature or storage non-compliance issues e.g. outside the 30 minute rule, problems with the blood fridge, any physical damage to the unit)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>2.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>3.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>4.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>5.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>6.</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**STORAGE CONFIRMATION (COMPULSORY)**

Indicate the following checks have occurred by ticking the appropriate box.

- Red Blood Cell Units:
  - Check the Blood Fridge Register to ensure each red cell unit being transferred has been stored appropriately:
    - red cell units have not been removed from the blood fridge longer than 30 minutes at any given time
  - Check the Blood Fridge Maintenance Record to ensure compliance of storage criteria:
    - blood fridge temperature has remained stable within 2 - 6°C degrees during the storage period
    - that the temperature recorded is complete with no missing data

- Other Blood Products:
  - Check that other blood and blood product have been stored as per manufacturer’s temperature specifications
    - If there are any problems with handling and storage of any of these blood or blood products:
      - these MUST be documented in the above comments section next to the appropriate blood or blood product
      - contact the laboratory and inform them of the details

I declare to the best of my knowledge, the above information regarding the handling and storage of the blood and blood product listed above is correct.

Name: Signature: Position:

**RECEIVING LABORATORY USE ONLY**

- Temperature check on receipt: _________ °C
- The above documentation has been completed verifying correct handling and storage of blood and blood products [boxes ticked, signature present]
- Tamper-proof port is intact and no blood is present in the port, for red cells only
- The blood and blood product is intact, not discoloured or has unusual particulate matter [check against other units if necessary]
- Only blood and blood product stored conforming to AS3886 and manufacturer’s temperature specifications have been accepted back into inventory
- Blood and blood products that are not compliant are to be destroyed via medical waste and recorded in LIS and BLOODNET, where available and appropriate

Checked by: Signature: Date: Time:
# APPENDIX 7: SHIPPER PACKING SLIP/ BLOOD CONSIGNMENT RECORD

<table>
<thead>
<tr>
<th>To:</th>
<th>From:</th>
</tr>
</thead>
</table>

## SENDER TO COMPLETE

<table>
<thead>
<tr>
<th>No. of Shippers:</th>
<th>Blood Product:</th>
<th>Qty:</th>
</tr>
</thead>
<tbody>
<tr>
<td>Patient Name:</td>
<td>Packed Date:</td>
<td>Time:</td>
</tr>
<tr>
<td>Sent via:</td>
<td>YOUR Lab courier □</td>
<td>Taxi □</td>
</tr>
<tr>
<td>Other Courier Company Name:</td>
<td>Details:</td>
<td></td>
</tr>
</tbody>
</table>

I have packed this consignment in accordance with the packing configuration:

Signature:  
Dispatched Date:  
Time:  

## RECIPIENT TO COMPLETE  *Please return completed form to Sender*

<table>
<thead>
<tr>
<th>No. of shippers received:</th>
<th>Shipment received unopened and undamaged?</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>YES □</td>
</tr>
</tbody>
</table>

For products listed below

- Is the temperature within the acceptable range?  YES □  NO □
- If product is outside specified temperature range, contact sender immediately for advice.

Received Date:  
Time:  
Unpacked Date:  
Time:  
Signature:  
Date:  

### WARNING

**DO NOT USE products if:**

- The shipper arrives open
- The product is outside the specified temperature range

**ACCEPTABLE TEMPERATURE RANGE**

- Red cells: 2°C to 10°C
- Autologous Blood: 2°C to 10°C
- Platelets: 20°C to 24°C
- Manufactured products as per Product Information

**Laboratory Notes**
APPENDIX 8: SHIPPER LABEL

DELIVER IMMEDIATELY

HUMAN BLOOD PRODUCTS
FOR TRANSFUSION

Do Not Refrigerate This Shipper

Deliver To:
Attention to: [Insert Name and Position]
[Insert name of Transfusion Laboratory]
[Insert location/building name]
[Insert name of Hospital][Insert address]
[Insert Phone Number]
[Insert Fax Number]

Delivered From:
[Insert Name and Position]
[Insert Hospital Name]
[Insert Address]
[Insert Phone Number]
[Insert Fax Number]

CONTENTS
☐ Autologous Blood
☐ Red Cells
☐ Platelets
☐ Thawed FFP
☐ Frozen Plasma Components
☐ Clotting Factors
☐ Immunoglobulins
☐ Albumin
Packed Time: Valid to Time:
Packed Date:
Despatch Date: Time:
Signed:
ACKNOWLEDGEMENTS, ENDORSEMENTS AND REFERENCES
Acknowledgements

The National Blood Authority would like to acknowledge the contributions of the following organisations to the development of these Guidelines:

- Hunter Area Pathology Service, New South Wales
- National Association of Testing Authorities (NATA)
- South Australian Department of Health, BloodMove Program
- Tasmanian Department of Health, Tasmanian Blood Product Network
- Therapeutic Goods Administration (TGA)

Endorsements

- Australian & New Zealand Society of Blood Transfusion Ltd (ANZSBT)
- The Royal College of Pathologists of Australia (RCPA)

References


