

# (archived) CBB Survey 2015 WO-1375

## CBB Survey 2015

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|---------------------------------------------------------------------------------------|--------------------------------|
| <b>1. General Information</b>                                                         |                                |
| Name of CBB                                                                           | CBB Basel                      |
| CBB Director                                                                          | Dimitrios                      |
| CBB Director                                                                          | Tsakiris                       |
| Address                                                                               | Diagnostic Hematology          |
| Address                                                                               | University Hospital Basel      |
| Address                                                                               | Petersgraben 4                 |
| Address                                                                               | CH-4031                        |
| Address                                                                               | Basel                          |
| Phone Number                                                                          | +41612652525                   |
| Date CBB Started Collecting Cord Blood Units<br><small>(month/day/year)</small>       | 06/04/1997                     |
| Number of Public Cord Blood Units                                                     | 2,966                          |
| Planned Number of Public Cord Blood Units Stored in 2015                              | 150                            |
| Lists on BMDW                                                                         | No                             |
| Affiliated with National Stem Cell Donor Registry                                     | Yes                            |
| Registry Affiliation                                                                  | Swiss Blood Stem Cell Registry |
| <b>2. Cord Blood Units in Inventory</b>                                               |                                |
| Current Processing Method                                                             | Liquid+ red auto               |
| Previous Processing Method                                                            | Liquid- red manual             |
| Year Current Process Method Started                                                   | 2000                           |
| Percent of Units Plasma and RBC Reduced (manual)                                      | 4                              |
| Percent of Units Plasma and RBC Reduced (automated)                                   | 96                             |
| <b>3. Accreditations, Licenses and Certifications</b>                                 |                                |
| FACT-Netcord                                                                          | No                             |
| AABB                                                                                  | No                             |
| Competent Authority/ National Health Authority                                        | Yes                            |
| Name of Competent Authority                                                           | BAG                            |
| Audited by National Stem Cell Registry                                                | Yes                            |
| ISO                                                                                   | Yes                            |
| Other                                                                                 | ISO17025 JACIE                 |
| <b>4. Cord Blood Collection</b>                                                       |                                |
| Current Collection Practice<br><small>Is the collection In/Ex -Utero or both?</small> | Ex-utero                       |

|                                                                                                     |                                                      |
|-----------------------------------------------------------------------------------------------------|------------------------------------------------------|
| Current Antiseptic                                                                                  | Betadine                                             |
| Collection Bag                                                                                      | Single needle                                        |
| Agitation during Collection                                                                         | Manual                                               |
| <b>5. Conditioning/Transport from Collection Site to CBB</b>                                        |                                                      |
| Secondary Bag Used                                                                                  | Yes                                                  |
| Transport Conditions                                                                                | Ground transport                                     |
| Temp. for Storage and Transport                                                                     | Temperature between +2 to +8°C                       |
| <b>6. Pre-Processing Evaluation</b>                                                                 |                                                      |
| Completed Prior to Accepting a CBU                                                                  | Medical history, collection report, informed consent |
| External Proficiency Testing for QC of FACS Lab                                                     | UKNEQAS                                              |
| Post Processing/ Pre Freeze CD34+ Cell Count                                                        | Yes                                                  |
| Time from Collection to Processing                                                                  | up to 48H                                            |
| <b>7. Processing and Packaging</b>                                                                  |                                                      |
| Pre Freeze Processing Methods- Unit in Inventory                                                    | SEPAX                                                |
| Pre Freeze Processing Methods- Unit in Inventory                                                    | Volume reduction with HES-Manual                     |
| Pre Freeze Processing Methods- Current                                                              | SEPAX                                                |
| Additives Currently in Use                                                                          | HES                                                  |
| Current Cryopreservation Method                                                                     | BioArchive                                           |
| Current Cryoprotectant Additive                                                                     | DMSO                                                 |
| Current Cryobag                                                                                     | Single bag (one fraction)                            |
| Current Target Cryopreservation Volume (mL)                                                         | 25.0                                                 |
| Current Packaging for Storage                                                                       | Overwrap                                             |
| Current Packaging for Storage                                                                       | Canister                                             |
| Current Packaging for Storage                                                                       | More than one segment                                |
| <b>8. Testing</b>                                                                                   |                                                      |
| Extra Material Currently Stored                                                                     | Cord blood DNA                                       |
| Extra Material Currently Stored                                                                     | Plasma/cord blood                                    |
| Extra Material Currently Stored                                                                     | Maternal DNA                                         |
| Extra Material Currently Stored                                                                     | Maternal plasma/serum                                |
| Current Post Processing Threshold for Accepting a CBU for Public Use<br>TNC                         | 150                                                  |
| Current Post Processing Threshold for Accepting a CBU for Public Use<br>CD34 (10^6) Single Platform | NA                                                   |
| Current Post Processing Threshold for Accepting a CBU for Public Use<br>CD34 (10^6) Double Platform | NA                                                   |
| Current Post Processing Threshold for Accepting a CBU for Public Use<br>CFU-GM                      | NA                                                   |
| Current Post Processing Threshold for Accepting a CBU for Public Use<br>CFU                         | NA                                                   |
| Current Post Processing Threshold for Accepting a CBU for Public Use<br>Viability                   | NA                                                   |
| <b>9. Storage</b>                                                                                   |                                                      |

|                                                                                              |                                      |
|----------------------------------------------------------------------------------------------|--------------------------------------|
| Type of Storage Container Used                                                               | BioArchive                           |
| Monitoring of Storage                                                                        | Centralized alarm system local       |
| Monitoring of Storage                                                                        | Centralized system remote monitoring |
| Monitoring of Storage                                                                        | LN2 level                            |
| <b>10. HLA Typing</b>                                                                        |                                      |
| Current Level of HLA Typing at Time of Listing<br>HLA-A                                      | IR                                   |
| Current Level of HLA Typing at Time of Listing<br>HLA-B                                      | IR                                   |
| Current Level of HLA Typing at Time of Listing<br>HLA-C                                      |                                      |
| Current Level of HLA Typing at Time of Listing<br>HLA-DRB1                                   | IR                                   |
| Current Level of HLA Typing at Time of Listing<br>HLA-DQB1                                   |                                      |
| Current Level of HLA Typing at Time of Listing<br>HLA-DPB1                                   |                                      |
| Accreditation of HLA Lab                                                                     | EFI accredited lab                   |
| Average Turnaround Time for Extended HLA Typing Results<br>in days                           | 7                                    |
| Attached Segment Used for Confirmatory/ Verification Typing                                  | No                                   |
| Units Listed without Attached Segment and have not been Previously Typed on Attached Segment | No                                   |
| Percentage of CBUs that have an Attached Segment                                             | below 50%                            |
| Confirmatory/ Verification Typing on an Attached Segment is Pre-Release Requirement          | No                                   |
| <b>11. Reservation and Cancellation Policies</b>                                             |                                      |
| What Point is a CBU Reserved for a Patient                                                   | HLA typing request                   |
| What Point is a CBU Reserved for a Patient                                                   | Reservation request                  |
| Length of Time a CBU can be Reserved<br>in days                                              | Other                                |
| Length of Time a CBU can be Reserved<br>in days                                              | not defined                          |
| Reservation Fee                                                                              | Yes                                  |
| Reservation Cancellation Fee in Absence of Shipment Request                                  | No                                   |
| Can Reservation be Extended                                                                  | Yes                                  |
| Is a Unit Report Provided on a Unit that is Reserved for Another Patient                     | No                                   |
| Is TC Informed when CBU is Released                                                          | Yes                                  |
| <b>12. Release and Shipment</b>                                                              |                                      |
| Hemoglobinopathy Screening Performed Prior to Release                                        | Yes                                  |
| Criteria to Ship a CBU<br>Viability and Cell Count                                           | NA                                   |
| Criteria to Ship a CBU<br>HLA Identity Testing                                               | Yes                                  |
| Current Packaging for Shipment to TC                                                         | Metal canister                       |
| Current Packaging for Shipment to TC                                                         | Overwrap                             |
| Current Packaging for Shipment to TC                                                         | 2 attached segments                  |

|                                                                                                      |                              |
|------------------------------------------------------------------------------------------------------|------------------------------|
| Time Between Shipment Request and Sending CBU                                                        | 1-2 weeks                    |
| Fee for Shipment Cancellation                                                                        | No                           |
| Dry Shippers Validated to Maintain Temperature of at least -150 for 48 hours Beyond Expected Arrival | Yes                          |
| Electronic Temperature Data Logger on All Dry Shippers                                               | Yes                          |
| Who Selects Transport Company                                                                        | Requesting transplant centre |
| Shape of Transport Container                                                                         | Mushroom                     |
| <b>13. Adverse Events Reporting</b>                                                                  |                              |
| Who are S(P)EARS Reported To<br>Competent Authority                                                  | Yes                          |
| Who are S(P)EARS Reported To<br>National Registry                                                    | Yes                          |
| Who are S(P)EARS Reported To<br>Transplant Center                                                    | Yes                          |

#### 14. Pictures of cord blood units in the inventory

#### 15. Infectious Disease Marker (IDM) CURRENTLY performed.



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#### Holiday Calendar

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| Team Calendars |
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