

Regulatory Survey ION-2614

The data provided on this page is updated by the organisations.

The WMDA makes no representations or express or implied warranties regarding any information on this page.

Use of any information provided on this site does not and is not intended to create a contractual or other relationship.

General	
Organisation name:	Austrian Bone Marrow Donor Registry
Organisation ION:	ION-2614
Country:	Austria
Year the registry started operations:	1989

Products	comment
Do you provide HPC, Marrow?	yes
Do you provide HPC, Apheresis?	yes
Do you provide HPC, Cord Blood?	yes
Do you provide MNC, Apheresis?	yes
Do you provide NC, Whole Blood?	yes

Product quantity		
Data valid for year	2018	
Number of products	National	International
HPC, Marrow products:	1	2
HPC, Apheresis:	4	9
HPC, Cord:	0	0

License	
Organisation is licensed/accredited by the Competent Authority:	yes
Comment:	The Registry is part of Gesundheit Oesterreich GmbH which was established by Act of Parliament. Owner of Gesundheit Österreich GmbH is the Federal Government, represented by the Ministry of Health.
Name of Competent Authority:	
Date of last inspection:	
Link to website of Competent Authority:	
Can you provide the Legal documentation from the Competent Authority that your organisation is allowed to operate as a registry ?	yes
Is your registry WMDA Certified, WMDA Qualified or WMDA Accredited?	WMDA Qualified
The registry is accredited by any other organisation:	no
If yes, by which organisation?	

Affiliation	
The registry works with the following number of affiliated donor centres:	5 affiliated donor centres
The registry audits its donor centres:	Audits will take place every two years.

The registry works with the following number of affiliated collection centres:	4 collection centres
Ther registry audits its collection centres:	Collection centers will be inspected and licensed by the Federal Office for Safety in Healthcare - Austrian Agency for Health and Food Safety. Licensed tissue establishments will be listed on the EU platform: https://webgate.ec.europa.eu/eucoding/reports/te/index.xhtml
The registry works with the following number of affiliated Cord Blood Banks:	1 cord blood bank
The registry audits its Cord Blood Banks:	
The registry works with the following number of affiliated transplant centres:	5 transplant centres
The registry audits its transplant centres:	Permit required by the local government
The registry works with the following number of affiliated IDM Testing Laboratories:	
The registry audits its IDM Testing Laboratories:	
The registry works with the following number of affiliated HLA/other DNA markers testing laboratories:	
The registry audits its HLA/other DNA markers testing laboratories:	
The registry would be able to provide a full list of name/addresses of each affiliated and their licence/accreditation status, on request:	yes
The Cord Blood Banks are FACT-NetCord accredited:	no
If yes, the following Cord Blood Bank(s) are accredited:	
The registry is able to provide a copy of all the certificates:	
Affiliated centres comply with WMDA Standards and applicable national regulations:	
The registry has requirements for affiliated centres in addition to WMDA Standards and applicable national regulations:	
If yes, what are these requirements?	

Donor policy

All donors are unpaid volunteers:	yes
All donors are informed about donation process and associated risks:	yes
Donors sign a valid informed consent to donate in the presence of a medical doctor/health care personnel/registry staff:	yes
The registry has systems in place to protect and control access to donor/patient records:	yes
The registry maintains donor anonymity:	yes
The registry has detailed donor evaluation and exclusion criteria in place:	yes
The registry has donor evaluation and exclusion criteria that do meet or exceed the WMDA guidelines:	

IDM

IDM	Tested	Method	Days between test and sampling/workup
ALT/ASTALT/AST ratio, De-Ritis-Quotient	-		
ChagasChagas, T. cruzi	-		
CMV IgGCytomegalovirus (CMV) Antibody testing IgG	Yes		
CMV IgMCytomegalovirus (CMV) Antibody testing IgM	Yes		
CMV TotalCytomegalovirus Total	No		
EBV IgGEpstein-Barr Virus Antibody testing IgG	Yes		

EBV IgM Epstein-Barr Virus Antibody testing IgM	Yes		
HAV (NAT) Anti-hepatitis A virus nucleic acid testing	-		
HBV (NAT) Hepatitis B nucleic acid testing	Yes		
HBc Ab Hepatitis B core antibody testing	Yes		
HBs Ag Hepatitis B Surface antigen testing	Yes		
HCV (NAT) Hepatitis C nucleic acid testing	Yes		
HCV Ab Hepatitis C antibody testing	Yes		
HEV (NAT) Hepatitis E Virus nucleic acid testing	-		
HIV (NAT) Human Immunodeficiency Virus nucleic acid testing	Yes		
HIV-1 Ab Human Immunodeficiency Virus HIV-1 antibody testing	Yes		
HIV-2 Ab Human Immunodeficiency Virus HIV-2 antibody testing	Yes		
HIV p24 Human Immunodeficiency Virus p24 antigen testing	On request		
HTLV-I Human T-Lymphotropic Virus type I testing	On request		
HTLV-II Human T-Lymphotropic Virus type II testing	On request		
Malaria Malaria	-		
HSV Herpes Simplex Virus	-		
STS Serological tests for syphilis	Yes		
STS FTA-ABS Serological test for syphilis	No		
Toxoplasmosis Toxoplasmosis	-		
VZV Varicella Zoster Virus	-		
WNV-NAT West Nile Virus nucleic acid testing	Yes		
Other tests performed	On request		

Testing

The physical and medical exam at donor workup is performed by a medical doctor:

All donor testing (at workup) for infectious disease is performed in a laboratory certified/licensed by a Competent Authority:

HLA typing for patient specific request is performed in an appropriately accredited laboratory:

Sterility testing is performed on the adult donor product:

Sterility testing is performed on the cord blood product:

Screening questionnaire to exclude communicable disease:

Screening questionnaire to exclude donors with 'high risk' lifestyles:

Donor reliability identified by a medical doctor:

Donor clearance to donate is confirmed by a medical doctor, following as a minimum the donor exclusion criteria in Annex 1 of [EU Directive 2006/17/EC](#) :

The party providing the cell product must exclude donors when:

- They are pregnant;
- They are breastfeeding;
- There is the potential for transmission of inherited conditions;
- There is evidence of any other risk factors for transmissible diseases on the basis of a risk assessment, taking into consideration *Donor travel and exposure history and local infectious disease prevalence*;
- There is presence on the donor's body of physical signs implying a risk of transmissible disease(s);
- There is a history of a disease of unknown aetiology;
- There is a risk of transmission of diseases caused by prions;
- There is systemic infection which is not controlled at the time of donation, including bacterial diseases, systemic viral, fungal or parasitic infections, or significant local infection in the tissues and cells to be donated;
- There is history of chronic, systemic autoimmune disease that could have a detrimental effect on the quality of the Cell Product;
- There is recent history of vaccination with a live attenuated virus where a risk of transmission is considered to exist;
- There is ingestion of, or exposure to, a substance (such as cyanide, lead, mercury, gold) that may be transmitted to recipients in a dose that could endanger their health.

Customs regulations

Are there any customs regulations to follow, or customs paperwork required, to import cell products into the organisation country? If yes, please specify:

Are there any customs regulations to follow, or customs paperwork required, to export cell products from the organisation country? If yes, please specify:

Are there any import regulations to follow, or paperwork required, to import cell products into the organisation country? If yes, please specify:

Are there any export regulations to follow, or paperwork required, to export cell products from the organisation country? If yes, please specify:

Reporting of Serious Adverse Events

Please indicate which of the following schemes for reporting Serious Adverse Events relating to either the Donor or the product the Registry participates in:

- Mandatory National Reporting Scheme

- Voluntary National Reporting Scheme

- WMDA SEAR/SPEAR Reporting Scheme

The registry will notify the receiving Registry within 24 hours of receiving information relating to any serious adverse event that could be considered to affect the patient receiving the product:

Quality management

The registry does maintain Standard Operating Procedures for your work:

The registry would be willing to provide these to WMDA or another registry upon request:

The registry would be willing to provide WMDA or another registry, on request, with copies of any packaging the HPC product will arrive in:

Donor records relating to the medical exam and HPC collection process are the following number(s) of year(s) retained: