



Australian Government

Department of Health
Therapeutic Goods Administration

Keith Alderman
Vice President Quality
Musculoskeletal Transplant Foundation
Suite 300 / 125 May Street
Edison NJ 08837

Our Reference: 2014/019664

Dear Mr Alderman

Subject: Issue of GMP certificate MI-2016-CE-01343-1

Please find enclosed the GMP certificate for your manufacturing premises.

The certificate remains valid only if re-inspections are conducted when scheduled by the Therapeutic Goods Administration. The inspection frequency is not a reflection of the expiry date shown on the certificate but is consistent with the re-inspection frequency applicable to Australian manufacturers of the same class of products.

The Therapeutic Goods Administration will contact the relevant sponsor/s to arrange the re-inspection of your facility.

Yours sincerely,

Signed and authorised by

Alyce Maksoud
Senior Inspector
Manufacturing Quality Branch

25 January 2018

Contact: gmp@tga.gov.au, phone +61 2 6221 6881 or fax +61 2 6232 8426



Australian Government

Department of Health
Therapeutic Goods Administration

Certificate of GMP Compliance of a Manufacturer

Certificate Number:

MI-2016-CE-01343-1

Issued to:

Musculoskeletal Transplant Foundation

Manufacturing Site Address:

Suite 300 / 125 May Street
Edison NJ United States Of America

The Therapeutic Goods Administration, the Competent Authority of Australia, confirms that this manufacturer has been inspected following section 32EA(1) of the *Therapeutic Goods Act 1989* in connection with manufacturers of Biologicals (items made from or containing human cells or human tissues) located outside Australia.

From the knowledge gained during inspection of this manufacturer, the latest of, that was conducted on 12 to 16 June 2017, it is considered that the manufacturer complies with the Australian Code of Good Manufacturing Practice for human blood and blood components, human tissues and human cellular therapy products (2013).

This certificate reflects the status of the manufacturing site at the time of the inspection noted above. This certificate remains valid until the expiry date provided that re-inspections are conducted as determined by the Therapeutic Goods Administration as the issuing Authority. This certificate should not be relied upon to reflect the compliance status after the expiry date.

EXPIRY DATE: 16 June 2019

ISSUE DATE: 25 January 2018

This Certificate remains valid only if re-inspections are conducted when scheduled by the Therapeutic Goods Administration.
The authenticity of this Certificate may be verified with the Therapeutic Goods Administration as the issuing authority.

PO Box 100 Woden ACT 2606 ABN 40 939 406 804
Phone: 02 6232 8644 Fax: 02 6203 1605 Email: info@tga.gov.au www.tga.gov.au



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Certificate of GMP Compliance of a Manufacturer

Certificate Number:

MI-2016-CE-01343-1

MANUFACTURING OPERATIONS

This certificate covers the following steps in the manufacture of therapeutic goods at the manufacturing site address specified above.

| Manufacturing Type | Product Category | Manufacturing Step |
|--------------------|---------------------------------|--|
| Human Tissue | Demineralised Bone Matrix (DBM) | Processing Packaging and labelling Storage on site Release for supply |
| Human Tissue | Musculoskeletal Tissue | Processing Packaging and labelling Storage on site Release for supply |
| Human Tissue | Allograft - Bone | Processing Packaging and labelling Storage on site Release for supply |
| Human Tissue | Acellular Human Dermal Matrix | Processing Packaging and labelling Storage on site Release for supply |
| Testing Laboratory | Not Applicable | Testing - Analytical |

This Certificate remains valid only if re-inspections are conducted when scheduled by the Therapeutic Goods Administration.

The authenticity of this Certificate may be verified with the Therapeutic Goods Administration as the issuing authority.