Page 1 of 1	Certificate of Conformance (COC) – Surgalign FibreX Demineralized Bone Matrix				
	FORM				
ORIGIN	Document No.	FORM-0305			
BIOLOGICS Life Enhancing Allografts	Parent SOP	SOP-0075			
	Revision	00			
	Effective Date	09 MAR 2021			

DONOR DESCRIPTION				
Donor Number				OB21-0005
Age				53
Gender	The same of the same of the same state of the sa		ers wile w	MALE
CONSENT		YES	NO	NOT DETERMINED
Research/Education		✓	Ш	
International		✓		
ELIGIBILITY DETERMINATION		YES	NO	NOT DETERMINED
EU criteria per EU Directive 2006/17	/EC including HTLV I/II			✓
PRODUCT DESCRIPTION				
Freeze dried, aseptically processed bor terminally sterilized via validated irrad	ne (demineralized bone fibers) cleansed wi iation process. Storage is 15°C to 30°C.	th a validated process tha	t utilizes G	entamicin. Final product is
Product ID Number(s)	OB21-0005	Lot No. R	Lot No. Range -060 th	
Customer Part Number				FBX-0010
Expiration Date				08 JUN 2023
PRODUCT ANALYSIS				
and the second s	ed to microbiological testing at recovery naerobic microorganisms and fungal conta	The state of the s		
The above listed product complies with Administration (FDA) for human tissue	the requirements as set forth by the Amer for transplantation.	rican Association of Tissue	Banks (AA	ATB) and the Food and Drug
TEST				RESULT
Sterility Dose Report				PASS

Residual Moisture DONOR ELIGIBILITY AND SEROLOGICAL TESTING

Residual Calcium

Donor Osteoinductivity Potential

Donor eligibility (screening and testing) is performed in accordance with AATB standards and FDA regulations (21 CFR 1271). Donor screening includes assessment of the medical and social history as well as physician assessment of the donor to assure that no conditions exist that may make the tissue unacceptable for transplantation. Donor eligibility is determined by a Board-Certified Physician (Medical Director).

Communicable disease testing was performed by a laboratory registered with FDA to perform donor testing and certified to perform such testing on human specimens in accordance with the Clinical Laboratory Improvement Amendments of 1988 (CLIA) and 42 CFR Part 493. The testing was conducted using FDA licensed, approved, or cleared donor screening tests for cadaveric specimens where applicable. The records of this testing are maintained at Origin Biologics, 6635 S. Eastern Avenue, Suite 100, Las Vegas, NV 89119. The following required testing was performed and found to be negative or non-reactive:

Antibody to Human Immunodeficiency Virus 1 & 2 (HIV 1 & 2) Human Immunodeficiency Virus Type 1 (HIV-1 NAT) Hepatitis B Core IgG/IgM Antibody (HBcAb) Rapid Plasma Reagin or Serologic Test for Syphilis (RPR or STS) Antibody to Hepatitis C (HCV) Hepatitis C Virus (HCV-NAT) Hepatitis B Surface Antigen (HBsAg) **PASS**

PASS

PASS

Additional tests including, but not limited to, Human T-Cell Lymphotropic Virus Type I & II (HTLV I & II) and HBV NAT may have been performed at the time of donor screening and were found to be acceptable for transplantation.

Quality Representative: Michael Date: Date