## IRB APPROVAL CERTIFICATION

**Continuing Review - Study** 



IRB Signature

Salus IRB Board 5 #IRB00013544 Roster dated 16September2022

2020

Signature <u>san Taylor-Wood</u> Jean Taylor-Woodbury, RN, MS, A Review and Approval Information Salus IRB Study Number 01015 - 23		Lie Wilson	1/20/2023 Date
Review Process Expedited 5		te Thursday, Janua te Saturday, Janua	ary 19, 2023 ary 20, 2024 at 11:59 PM
This document certifies the IRB's continuing re Approved" to be conducted by the named Prir No content changes were made to the bolded	ncipal Investigator.		
Applied FDA's Guidance: Informed Consent for		Device Studies Usir	ng Leftover Human
	Client Sponsor	New York Biologic New York Biologic	s, Inc.
Study Surplus Sample Collection	Client Sponsor  Address New York 41 Oak G	New York Biologic	s, Inc.
Study Surplus Sample Collection  Principal Investigator Samuel B. Reichberg, MD, PhD	Client Sponsor  Address New York 41 Oak G Southamp	New York Biologic New York Biologic Biologics, Inc.	s, Inc.

## Stipulations of Approval

Collection Protocol

- No subjects may be involved in any study procedure prior to the IRB approval date or after the expiration date, unless otherwise stated in this letter. Investigators and sponsors are responsible for initiating Continuing Review proceedings.
- 2. All protocol modifications must be IRB approved prior to implementation. This includes any addition or change of recruitment materials, change of investigator, or performance site address. (Exception: If necessary to eliminate apparent immediate hazard to subjects.)
- 3. Report to Salus IRB within ten working days of learning if any of the following occur:
  - Unanticipated problems involving risk to human subjects or others;
  - Unanticipated Serious Adverse Events and Safety Reports;
  - Protocol deviations, violations, and exceptions that impact subject welfare or safety or study integrity including changes intended to reduce immediate risk to subjects;
  - Use of an investigational product in an emergency situation; and
  - Claims for compensation or for medical care for research-related injury.
- 4. Advertising and recruitment materials must be approved by Salus IRB prior to use or publication.





## IRB APPROVAL CERTIFICATION

**Continuing Review - Study** 

Salus IRB Board 5 #IRB00013544 Roster dated 16September2022

Copies to:

Ivy Winick, Ivy.Winick@newyorkbiologics.com

END