

**IRB Signature**

Signature Jean Taylor-Woodbury By TM 1/16/2019  
 Jean Taylor-Woodbury, RN, MS, ANP-BC, Chair Teresa Majors, CIP Date

**Review and Approval Information**

E&I Study Number 01015 - 19 Approval Date Tuesday, January 15, 2019  
 Review Process Expedited 5 Expiration Date Monday, January 20, 2020 at 11:59 PM

**This document certifies the IRB's continuing review approval of the items identified under "Documents Approved" to be conducted by the named Principal Investigator.**

**Applied FDA's Guidance: Informed Consent for In Vitro Diagnostic Device Studies Using Leftover Human Specimens that are Not Individually Identifiable.**

**Study**

Surplus Sample Collection Client New York Biologics, Inc.  
 Sponsor New York Biologics, Inc.

**Principal Investigator**

Samuel B. Reichberg, MD, PhD

**Address**

New York Biologics, Inc.  
41 Oak Grove Road  
Southampton, NY 11968

E&I PI Number 17023 - 002

**Performance Sites**

New York Biologics, Inc., 41 Oak Grove Road, Southampton, NY 11968

**Documents Approved**

**Document # Version Date**

Document #	Version	Date
Protocol Description (Cover Letter)		January 11, 2017
Collection Protocol		Revised 01-11-17

**Stipulations of Approval**

- No subjects may be involved in any study procedure prior to the IRB approval date or after the expiration date. Investigators and sponsors are responsible for initiating Continuing Review proceedings.
- 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.)
- Report to E&I within five 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.
- Advertising and recruitment materials must be approved by E&I prior to use or publication.

**Copies to:**

Ivy Winick, nybiologics@gmail.com

This is a multi-sided document.

**E&I Business Office**

14400 East 42<sup>nd</sup> Street, Suite 240  
 Independence, MO 64055  
**Phone (816) 421-0008**  
 Fax (816) 356-2227



**E&I West Coast Board**

5710 Paradise Drive, Suite 11  
 Corte Madera, CA 94925  
**Phone: (415) 485-0717**  
 Fax: (415) 485-0328

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**END**

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