

# Document Cover Page

|                    |   |
|--------------------|---|
| Document Title:    | 22_System Release Compliance Report for IGMP  |
| Classification:    | CSVP  |
| Type:              | System Release Compliance Report  |
| Document Number:   | CSVP-SSR-001  |
| Version #:         | 39  |
| Document ID:       | CSVP-SRR-IGMP-4.001.006   |
| File Name:         | csvp-srr-igmp-4.001.006.pdf   |
| Status:            | Approved  |
| Effective Date:    | 11/15/22  |
| Review Date:       | / /   |
| Reason for Change: | Improvements<br><ul style="list-style-type: none"><li>• Allow 5 digit MPR and BPR numbers</li></ul> Fixes<br><ul style="list-style-type: none"><li>• All roles can update the watermark and review date on documents in the DMS</li></ul> |

## Reviewers:

Richard Soltero (rs01) 11/15/22 11:01 AM Eastern Standard Time

Angelo Nardone (ANardone01) 11/15/22 11:08 AM Eastern Standard Time


Brandy Irons (bi01) 11/15/22 11:15 AM Eastern Standard Time

## Approvals:

Richard Soltero (rs01) Approved 11/15/22 11:17 AM Eastern Standard Time

Angelo Nardone (ANardone01) Approved 11/15/22 11:46 AM Eastern Standard Time

Brandy Irons (bi01) Approved 11/15/22 11:15 AM Eastern Standard Time

|   |   |                                      |
|---|---|--------------------------------------|
|  | <b>Title: CSVP - System Release / Compliance Report for InstantGMP v4.001.006</b> | Document No: CSVP-SRR-IGMP-4.001.006 |
|   |   | Effective upon last signature date   |
|   |   | Page: 1 of 4                         |

## TABLE OF CONTENTS

|   |          |
|---|----------|
| <b>1. DEFINITIONS AND ACRONYMS.....</b>   | <b>1</b> |
| <b>2. PURPOSE.....</b>                    | <b>1</b> |
| <b>3. SCOPE.....</b>                      | <b>1</b> |
| <b>4. SYSTEM DESCRIPTION.....</b>         | <b>2</b> |
| 4.1. Software and Hardware.....           | 2        |
| <b>5. REFERENCES.....</b>                 | <b>2</b> |
| <b>6. ROLES AND RESPONSIBILITIES.....</b> | <b>3</b> |
| 6.1. System Owner.....                    | 3        |
| 6.2. Information Technology.....          | 3        |
| 6.3. Quality Assurance.....               | 3        |
| <b>7. COMPLIANCE STATEMENT.....</b>       | <b>4</b> |
| <b>8. SYSTEM RELEASE STATEMENT.....</b>   | <b>4</b> |


### 1. Definitions and Acronyms

| <b>Term</b> | <b>Definition</b>  |
|-------------|--|
| GXTest      | Automated Validation Software used for Operation Qualification testing |
| SOP         | Standard Operating Procedures  |
| SRR         | System Release/Commissioning Report                                    |
| MVP         | Master Validation Plan   |

### 2. Purpose

The purpose of this System Release/Commissioning Report is to document the release of the InstantGMP system for use.

#### 1. Scope

|   |   |                                      |
|---|---|--------------------------------------|
|  | <b>Title: CSVP - System Release / Compliance Report for InstantGMP v4.001.006</b> | Document No: CSVP-SRR-IGMP-4.001.006 |
|   |   | Effective upon last signature date   |
|   |   | Page: 2 of 4                         |

The scope of this System Release/Commissioning Report is the release and commissioning of the InstantGMP system.

### 3. System Description

#### 3.1. Software and Hardware


The software operates as a cloud application with the following minimum basic parameters for the software and hardware configuration.

- Application Server
  - Windows Server 2019
  - IIS 10.0 version 1809
    - URL Rewrite 2.0.
    - WebSockets enabled
  - .NET Framework 3.5 and 4.7 – All features enabled
- Database Server
  - Windows Server 2019
  - SQL Server 2019 Standard or Enterprise
- Servers are virtualized and dedicated to InstantGMP, Inc.

### 4. References

The purpose of this section is to serve as an index of all documents referenced for production of this document.

| Document Identification | Document Title                    |
|-------------------------|-----------------------------------|
| CSVP-POL-IGMP           | Computer System Validation Policy |
| CSVP-MVP-IGMP           | Master Validation Plan            |
| CSVP-IQ-IGMP            | Installation Qualification        |
| CSVP-GXTEST-OQT-IGMP    | CSVP-OQT-Test Script-IGMP         |
| CSVP-GXTEST-OQE-IGMP    | CSVP-OQE-Test Script-IGMP         |
| CSVP-VSR-IGMP           | CSVP-Validation Summary report    |

|   |   |                                      |
|---|---|--------------------------------------|
|  | <b>Title: CSVP - System Release / Compliance Report for InstantGMP v4.001.006</b> | Document No: CSVP-SRR-IGMP-4.001.006 |
|   |   | Effective upon last signature date   |
|   |   | Page: 3 of 4                         |

## 5. Roles and Responsibilities

The roles and responsibilities for the IQ/OQ protocols are defined below.

| Functional Group       | Assigned Employee / Company        |
|------------------------|------------------------------------|
| System Owner           | Richard Soltero, Ph.D., InstantGMP |
| Information Technology | Angelo Nardone, InstantGMP         |
| Quality Unit           | Brandy Irons, InstantGMP           |

### 5.1. System Owner

The ultimate responsibility for the controlled development, implementation, maintenance, and validation of the computer or automated system resides with the identified system owner. Responsibilities include:

- Define the business area needs and the intended use of the system.
- Ensure that system development, implementation, and use conform to the principles of the System Development Life Cycle (SDLC) process and the IQ/OQ.
- Ensure that the use and maintenance of the system complies with documented procedures that maintain the system in a validated state.
- Ensure the accuracy and integrity of the data entered into and resident on the system.


### 5.2. Information Technology

The person performing or overseeing the development and technical aspects of the system is responsible for the following items:

- Create a system technical design that satisfies the specified business area and regulatory requirements.
- Ensure that the technical design, construction, testing and delivery of the system conform to principles of the SDLC process and the IQ/OQ protocols.
- Ensure that maintenance and changes to the system are performed according to documented procedures that maintain the validated status of the system.
- Ensure that the products or services supplied by external vendors and/or suppliers adhere to the principles of the SDLC process.

### 5.3. Quality Assurance

The Quality Assurance Unit is that organization or person designated by InstantGMP to be an independent reviewer for all completed and/or executed validation documentation. The Quality Assurance Unit is responsible for the following items:

|   |   |                                      |
|---|---|--------------------------------------|
|  | <b>Title: CSVP - System Release / Compliance Report for InstantGMP v4.001.006</b> | Document No: CSVP-SRR-IGMP-4.001.006 |
|   |   | Effective upon last signature date   |
|   |   | Page: 4 of 4                         |

- Maintain independence and objectivity relative to the system development, implementation, use and validation processes.

## 2. Compliance Statement

This InstantGMP system has been validated to comply with the standards outlined in the following references:

| Document Identification | Document Title   |
|-------------------------|--|
| N/A                     | General Principles of Software Validation; Final Guidance for Industry and FDA Staff                                     |
| 21 CFR Part 11          | Electronic Records; Electronic Signatures  |
| 21 CFR Part 820         | MEDICAL DEVICES - QUALITY SYSTEM REGULATION  |
| 21 CFR Part 111         | CURRENT GOOD MANUFACTURING PRACTICE IN MANUFACTURING, PACKAGING, LABELING, OR HOLDING OPERATIONS FOR DIETARY SUPPLEMENTS |
| 21 CFR Part 211         | CURRENT GOOD MANUFACTURING PRACTICE FOR FINISHED PHARMACEUTICALS   |
| Annex 11                | EudraLex Volume 4, Annex 11: Computerized Systems  |
| ISO 9001                | Quality Management System  |
| ISO 13485               | Medical Devices - Quality Management System  |
| GAMP 5                  | General Automated Manufacturing Practices  |

## 3. System Release Statement

This System Release Statement is written to indicate that the InstantGMP system is accepted by the system owner and is deemed suitable for release. The Business Owner and IT base this release upon the successful completion, review and approval of the Validation deliverables and the other activities required by the Master Validation Plan for the InstantGMP system.