

## **CERTIFICATE**OF REGISTRATION

This is to certify that the management system of:

## Black Tie Medical, Inc. dba Tulip Medical Products

(F004127)

Main Site: 4360 Morena Blvd., Suite 100

San Diego, California, 92117, United States

has been registered by Intertek, an MDSAP recognized auditing organization, as conforming to the requirements of:

## ISO 13485:2016

Australia: Therapeutic Goods (Medical Devices) Regulations, 2002, Schedule 3 Part 1 (excluding Part 1.6)

**Brazil**: Federal Law n. 6360/76; RDC ANVISA n. 16/2013; RDC ANVISA n. 23/2012; RDC ANVISA n. 67/2009; RDC ANVISA n. 56/2001

Canada: Medical Devices Regulations – Part 1- SOR 98/282

United States: 21 CFR 820, 21 CFR 803, 21 CFR 806, 21 CFR 807 (Subparts A to D)

Japan: MHLW Ministerial Ordinance 169, Article 4 to Article 68; PMD Act (as applicable)

## The management system is applicable to:

Design and development, production and distribution of Liposuction, Fat-Harvesting, Tissue-Harvesting, Fluid and Tissue Injection Systems and Accessories. **Certificate Number:** 

0102982

Revision Level: 01

**Initial Certification Date:** 

2020-07-01

**Certification Effective Date:** 

2023-06-30

**Certification Expiry Date:** 

2026-06-30



Calin Moldovean

President, Business Assurance

Intertek Testing Services NA, Inc. 900 Chelmsford Street Lowell, MA, USA 01851



