

*GMED certifies that the quality management system developed by*

**OriGen Biomedical, Inc.**  
**7000 Burleson Road, Bldg. D**  
**AUSTIN, TX 78744 UNITED STATES**

**Facility identifier (REPs-generated) : F000671**

*for the activities*

**Conception, Developpement, Fabrication et Distribution de sacs pour culture cellulaire, sacs de congélation et dispositifs associés de transfert de fluides et solutions de cryoconservation.**

*Design, development, manufacture and distribution of cell culture bags, freezing bags and associated fluid transfer devices and cryopreservation solutions.*

*performed on the location(s) of*

**OriGen Biomedical, Inc. 7000 Burleson Road, Bldg. D AUSTIN, TX 78744 USA**

**has been audited and found to conform to the requirements of the international standard ISO 13485 : 2016 and following regulatory requirements**

Australia	Therapeutic Goods (Medical Devices) Regulations, 2002, Schedule 3 Part 1 (excluding Part 1.6) - Full Quality Assurance Procedure
Brazil	RDC ANVISA n. 665/2022 RDC ANVISA n. 551/2021 RDC ANVISA n. 67/2009
Canada	Medical Devices Regulations - Part 1 - SOR 98/282
Japan	MHLW MO 169 PMD Act
United States	21 CFR 820 21 CFR 803 21 CFR 806 21 CFR 807 - -Subparts A to D 21 CFR 821 (where applicable)

**Début de validité / Effective date December 4th, 2024 (included)**

**Valable jusqu'au / Expiry date :December 3rd, 2027 (included)**

**Etabli le / Issued on : October 15th, 2024**



GMED is authorised under the Medical Devices Single Audit Program  
 This certificate is issued according to the rules of GMED Certification  
 The validity of this certificate can be verified on [www.gmed.fr](http://www.gmed.fr)

Renouvelle le certificat 34919-5



DocuSigned by:  
*Béatrice LYS*  
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**On behalf of the President**  
**Béatrice LYS**  
**Technical Director**