

# Certificate

acc. to **ISO 13485:2016**

Medical Devices - Quality Management Systems - Requirements for Regulatory Purposes

Certificate Registration No.: **21-1603-Q**

TUV USA, Inc. hereby certifies that the quality management system of the company mentioned below is in conformance with **ISO 13485:2016** for Medical Devices-Quality Management Systems-Requirements for Regulatory Purposes.

**Inovo, Inc.**  
**401 Leonard Blvd. North**  
**Lehigh Acres, FL 33971, USA**

Additional sites covered by QM System: *N/A*

Scope:

**Design and Development, Production, Distribution, Final Testing, and Servicing of Respiratory Devices**

The validity of this certification document can be obtained by contacting the TUV USA, Inc. Office.

TUV USA, Inc. (a Member of the TÜV NORD Group)

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Audit Report Reference No.: **21-3935 TRF**

Certificate Initial Issue Date: **2021-03-26**

Current Cycle Start Date: **2018-06-22**

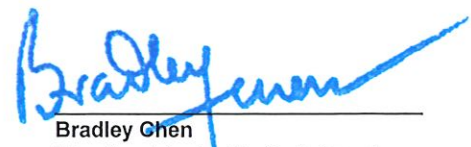
Certificate Revised Date: **2021-03-26**

Effective Date:

**2021-03-26 / ed. 1**

Valid Until:

**2021-06-21**



Bradley Chen  
Vice President - Medical, Americas  
Medical Products Division  
TUV USA, Inc.