EC-Declaration of Conformity

EC - Declaration of Conformity

NovoFine®

30G 8 mm

31G 6 mm

32G 6 mm

32G 4 mm

Novo Nordisi

s: Find 2 of

EC - Declaration of Conformity

We,

Legal Manufacturer/ Market Authorization Holder

Novo Nordisk A/S Novo Allé 2880 Bagsværd Denmark

Being the manufacturer/distributor within the European Economic Area, declare that this Declaration of Conformity is issued under our sole responsibility and covers the following product(s):

Product Name	Item number (4 or 5-number)	Classification	GMDN Code
NovoFine® 30G 8 mm	5-4325-21 5-4327-16	Class IIa	44127
NovoFine® 31G 6 mm	5-4356-16 5-4356-17	Class IIa	44127
NovoFine® 32G 6 mm	5-4314-18 5-4314-17	Class IIa	44127
NovoFine® 32G 4 mm	5-4401-11	Class IIa	44127

manufactured in the below mentioned production facilities:

Japan	., 2-19-64, Matsubara, Tatebayashi-shi, Gunma, 374-8518,
Nipro (Thailand) Corporation	n Ltd., 10/2 Moo 8, Bangnomko, Sena, Phra Nakhon Si
Ayutthaya, 13110, Thailand	
Needle Manufacturing & Sou	rcing, Stenager Allé, 9800 Hjorring, Denmark

declare that the above is in conformity with the provisions of the Council Directive

European Council Medical Device Directive 93/42/EEC, of 14 June 1993, inclusive amendment 2007/47/EC of 5 September 2007.

The above devices are CE-marked and classified as IIa according to Annex IX, Classification Criteria, rule 6.

Novo Nordisi

The devices have been subject to the conformity procedure laid down in Annex II under the supervision of TÜV SÜD Product Service GmbH, a Notified Body authorized by the German Competent Authority, and carrying the Notified Body number 0123.

Notified Body Address: TÜV SÜD PRODUCT SERVICE GMBH

PS-NAM1-MUC Ridlerstr. 65 80339 Munchen Germany

The following standards have been observed:

EN ISO 13485:2016/AC:2018	Medical devices - Quality management systems - Requirement for regulatory purposes	
EN ISO 14971:2019	Medical devices - Application of risk management to medical devices	
EN ISO 11608-2:2012	Needle-based injection systems for medical use – Requirements and test methods – Part 2: Needles	
EN ISO 14155:2011	Clinical investigation of medical devices for human subjects - good clinical practice	
EN 62366-1:2015	Medical devices – Part 1: Application of usability engineering to medical devices	
ISO 11607-1: 2019	Packaging for terminally sterilized medical devices - Part 1: Requirements for materials, sterile barrier systems and packaging	
ISO 11607-2:2019	Packaging for terminally sterilized medical devices - Part 2: Validation requirements for forming, sealing and assembly processes	
EN ISO 11135:2014/A1:2019	Sterilization of health-care products - Ethylene oxide - Requirements for the development, validation and routine control of a sterilization process for medical devices - Amendment 1: Revision of Annex E, Single batch release (ISO 11135:2014/Amd 1:2018)	
EN ISO 11737-1:2006 + AC:2009 (NMS)	Sterilization of medical devices – Microbiological Methods-Part 1: Determination of a population of microorganisms on products	
EN ISO 11737-1:2018 (NMI)	Sterilization of medical devices – Microbiological Methods-Par 1: Determination of a population of microorganisms on products	

Novo Nordisk A/S Department - 42102 EC-DoC NovoFine® nD ID: 002182803

Date: Version: Status: Page:

24 November 2020 8.0

Final 4 of 4 Novo Nordisi

EN ISO 11737-2: 2009 (NTC)	Sterilization of medical devices - Microbiological methods - Part 2: Tests of sterility performed in the definition, validation and maintenance of a sterilization process	
ISO 11737-2:2019 (NMI)	Sterilization of health care products - Microbiological methods - Part 2: Tests of sterility performed in the definition, validation and maintenance of a sterilization process	
ISO 10993-1:2018	Biological evaluation of medical devices - Part 1: Evaluation and testing within a risk management process	
EN ISO 14644-1:2015	Cleanrooms and associated controlled environments – Part 1: Classification of air cleanliness by particle concentration	
EN ISO 14644-2:2015	Cleanrooms and associated controlled environments — Part 2: Monitoring to provide evidence of cleanroom performance related to air cleanliness by particle concentration	
EN 1041:2008+A1:2013	Information supplied by the manufacturer of medical devices	
EN ISO 15223-1:2016	Medical devices - Symbols to be used with medical device labels, labelling and information to be supplied - Part 1: General requirements	

Location: Herring

On: 2020.11.25 By:

Name & function of person signing

Nirector of Quality.

Certificate of A a ysis



isk®

Novo Nordisk A/S Novo Allé 2880 Eagsvaerd Denmark

Tel +45 4444 8668 Fax +45 4449 0665

NOVOFINE 30G X 8MM

100PCS.

7200165

Product No. Batch No.

20D06M

Order Number

: 7735583-000

Date of Manufacture : 04/2020

Date of Expiry

: 03/2025

Supplier: Nipro Thailand Corporation LTD, Thailand

Method of sterility: Gas (ethylene oxide)

Sterility: Complies with the USP ed. 42 <71> and EN 556.

We hereby declare that the above-mentioned batch was manufactured and controlled in compliance with the GMP regulations and approved according to the quality specifications of Novo Nordisk A/S.

20200923, Hjoerring

LENE JØRGENSEN

Quality Department

The Certificate of Analysis is generated by a computer quality system udes an elec tures ..

304888 _ LNJ

064 HUNGARY