## **DECLARATION OF CONFORMITY**

## **Certificate Number:** ZP 06-014 Flow-metering device\_09-01

This declaration of conformity is issued under the sole responsibility of the

Manufacturer:	GCE, s.r.o.
	Žižkova 381, 583 01 Chotěboř
	CZECH REPUBLIC

We hereby declare that the medical device specified below is in conformity with applicable Regulation MDD 93/42/EEC for medical devices. This declaration is supported by the Quality System approval to ISO 13485:2016 issued by DNV GL PRESAFE AS, valid until May 11, 2021.

Product group:	Medical devices for use with Medical gases	
Product name:	Flow-metering device	
Model:	MEDIMETER (FlowO2 ST, FlowO2 GS, FlowAIR ST, FlowAIR GS,	
	FlowO2 Duo ST, FlowO2 Duo GS, FlowAIR Duo ST)	
Risk Classification:	lla	

The product is in accordance with Annex II (excluding section 4) of the MDD 93/42/EEC and is safe for declared purpose of use under standard conditions. Any modification to the product, not authorized by us, will invalidate this declaration.

Notified Body:	<b>DNV GL PRESAFE AS</b> Veritasveien 3, N-1363 Høvik Norway
Notified Body No.:	2460
EC Certificate number:	10401-2017-CE-CZS-NA-PS Rev. 5.0
Valid until:	May 27, 2024
Name:	Tereza Šnapková
Function:	Regulatory Specialist
Signature:	
Place and date of issue:	Chotěboř, September 22, 2020

