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EU Declaration of Conformity

This European Declaration of Conformity is issued under the sole responsibility of the manufacturer.

MANUFACTURER					
Name of Company	Registered Office	Trading Address	SRN		
Medicare Products Ltd	South Fen Road, Bourne, Lincolnshire, PE10 0DN. UK	Unit B, Dolphin Way, Purfleet, Essex, RM19 1NZ, UK	GB-MF-000011612		

AUTHORIZED REPRESENTATIVE					
Name of Company	Address	SRN	Phone/email		
International Associates	The Black Church, St		+353 16971561		
Auditing & Certification	Mary's Place, Dublin 7,	IE-AR-000002248	EUAR@ie.ia-net.com		
Limited	D07 P4AX Ireland				

Product Name	Code / Catalog Number	Basic UDI-DI
Blue Nitrile Powder Free Examination Glove (Nitrex Extra Sensitive)	GN01	506004079GNPFNS00LX
Intended Purpose	Photo	
Examination gloves for non-invasive use to prevent contact with skin, bodily fluids, and chemicals. Invasive with respect to natural body orifices.		

RISK CLASS FOR MEDICAL DEVICES				
Device Classification Common Specifications / Standards		ications / Standards		
Class:	I	EN455-1:2020	Medical gloves for single use: Freedom from holes	
		EN455-2:2015	Medical gloves for single use: Physical properties	
Rule:	5	EN455-3:2015	Medical gloves for single use: Biological evaluation	
		EN455-4:2009	Medical gloves for single use: Shelf-life determination	

Medicare Products Ltd declares that the above-mentioned product meets the provision of the following EU legislation:

- Medical Devices Regulation (EU) 2017/745
- Is classed as Cat III PPE in conformity with the provisions of Regulation (EU) 2016/425 on personal protective equipment and with National Standards transposing harmonised standards EN 374-1:2016+A1:2018, EN 374-5:2016, and EN ISO 21420:2020.

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- Is identical to the personal protective equipment which is the subject of EU Type-Examination Certificate number 2777/10015-07/E01-01, issued by SATRA Technology Europe Limited, Bracetown Business Park, Clonee, Dublin, D15 YN2P, Republic of Ireland (Notified Body number 2777), according to Annex V (Module B) of Regulation (EU) 2016/425.
- Is subject to the conformity assessment procedure Module C2 set out in Annex VII of Regulation (EU) 2016/425, under the surveillance of the notified body SATRA Technology Europe Limited, Bracetown Business Park, Clonee, Dublin, D15 YN2P, Republic of Ireland (Notified Body number 2777).

COMPANY REPRESENTATIVE: David Langridge

TITLE: Head of Technical SIGNATURE: Dangrudge

PLACE: Bourne, UK **DATE**: 04/03/2022