



EC DECLARATION OF CONFORMITY

Product: GD21 - POWDER FREE EXAMINATION NITRILE GLOVES

SANTEX 2000 INTERNACIONAL SL in Paseo de la Castellana, 141 plantas 8 y 9, Cuzco IV 28046 Madrid, with SRN: ES-MF-000026359 declares that:

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

The afore mentioned materials and products have been revised in accordance with the self-declaration of conformity requirements for EC marking according to the Medical Devices Regulation (EU) 2017/745, and standards EN 455-1, EN 455-2, EN 455-3, EN 455-4.

In the transformation and production of the film, all current health standards are followed, being the facilities where it has been manufactured certified with the international harmonized standard ISO 13485: 2016.

According annex VIII Regulation (EU) 2017/745, It is a class I medical device and Basic UDI-DI: 84370NITRILOTV.

In Mataró, on July 25, 2022

A handwritten signature in blue ink, consisting of several vertical strokes followed by a diagonal line extending to the right.