

Bergamot Sdn Bhd (1153028-V)

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EU DECLARATION OF CONFORMITY

Manufacturer:

Bergamot Sdn Bhd

31, Jalan Balakong Jaya 4, Taman Industri Balakong Jaya, 43300

Seri Kembangan, Selangor, Malaysia

SRN: MY-MF-000004005

Brand:

Bergamot

Product Type & Basic UDI-DI:

Latex Powdered Examination Gloves (9555904LPSMS)

Latex Powder Free Examination Gloves (9555904LPFLY)

Nitrile Powder Free Examination Gloves (9555904NPFMA)

Classification and Rules:

Medical Device Regulations (EU) 2017/745, under Class I, Rule 5

of Annex VIII

European Standards:

EN 455-1:2000, EN 455-2:2015, EN 455-3:2015 and EN 455-

4:2009

EU Authorized Representative:

ASAP INNOVATIONS LIMITED

7, Saggart Lakes, Saggart,

County Dublin, D24PY01, Ireland

SRN: IE-AR-000002548

This EU declaration of conformity is issued under the sole responsibility of manufacturer, Bergamot Sdn. Bhd. We hereby declare that device covered by the present declaration is in conformity with the Medical Device Regulation (EU) 2017/745 and with the above-mentioned standards. All supporting documentation is retained at the premises of the manufacturer.

Place, Date of Declaration:

MALAYSIA, 5th May 2021

Signature:

Mr. Chin Tze Kai

Director