



## EC Declaration of Conformity

Doc. No.: MDR\_DOC\_GVYP  
Revision: 0  
Revision Date: 2021-03-01

**Manufacturer:** Pharma Lab International Ltd.  
**Single Registration Number:** TBA  
**Address:** Unit 2102, CC Wu Building, 302-308 Hennessy Road, Wan Chai, Hong Kong  
**European Authorized Representative:** Obelis s.a  
Bd. Général Wahis 53, 1030 Brussels, Belgium  
**Basic UDI-DI:** 366496515101G6  
**Product:** Vinyl Examination Gloves, Powdered  
**Type:** Powdered  
**Product Codes:**

Pharma Lab Int'l Codes	Size
GVYP-S	S
GVYP-M	M
GVYP-L	L

**Classification:** Class I according to Medical Device Regulation 2017/745  
Category III according to PPE Regulation (EU) 2016/425

We hereby declare under sole responsibility that the CE marked products mentioned above conform to the general safety and performance requirement (Annex I) of the Medical Device Regulation 2017/745.

The conformity assessment is based on Annex II and Annex III; Classification accordingly to rule 5, Annex VIII.

**Applied standards:** EN 455-1:2020, EN 455-2:2015, EN 455-3:2015, EN 455-4:2009

We hereby declare under sole responsibility that the CE 2777 marked products mentioned above are in conformity with the provisions of PPE Regulation (EU) 2016/425 and are identical to the PPE, which are the subject of EC certificate of conformity no. 2777/11331-02/E04-0 issued on 01 Feb 2019 by the Notified Body:

**SATRA Technology Europe Limited (Notified Body No: 2777)**

Bracetown Business Park,  
Clonee, D15YN2P,  
Republic of Ireland

These devices are subject to the procedure set out in ANNEX VII of PPE Regulation (EU) 2016/425 under the supervision of the Notified Body:

**SATRA Technology Europe Limited (Notified Body No: 2777)**

Bracetown Business Park,  
Clonee, D15YN2P,  
Republic of Ireland

**Applied standards:** EN 374-1:2016, EN 374-2:2014, EN 16523-1:2015, EN 374-4 :2013, EN ISO 374-5 :2016 and EN ISO 21420:2020

Anthony Guchet  
Managing Director

Issued: Hong Kong, 01 Mar, 2021

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