



ΕΘΝΙΚΟ ΚΕΝΤΡΟ ΑΞΙΟΛΟΓΗΣΗΣ
ΤΗΣ ΠΟΙΟΤΗΤΑΣ & ΤΕΧΝΟΛΟΓΙΑΣ
ΣΤΗΝ ΥΓΕΙΑ Α.Ε.

NATIONAL EVALUATION CENTER
OF QUALITY & TECHNOLOGY
IN HEALTH S.A.

DATE: 18/05/23
REF. NUM.: 76754

Notified Body Statement Letter for audit surveillance of VIOSER S.A PARENTERAL SOLUTIONS INDUSTRY

To whom it may concern,

Confirmation of the status of an appropriate surveillance in the framework of Regulation (EU) 2023/607 of 15 March 2023 amending Regulations (EU) 2017/745 and (EU) 2017/746 as regards the transitional provisions for certain medical devices and in vitro diagnostic medical devices.

This letter confirms that, **EKAPTY**, a Notified Body (NB) designated against Medical Devices Directive (EU) 93/42/EEC (MDD) and identified by the number **0653** on NANDO, has provided a surveillance audit on 03 & 04/05/2023 for the extension of the validity of the certificate numbered **301011036CN** issued in accordance with Directive 93/42/EEC.

During the transitional period devices of the aforementioned certificate can lawfully be placed on the market by the condition that they are subject to the manufacturer's continued compliance to the conditions specified in Article 120.3c of MDR (as amended by (EU) 2023/607).

On behalf of the Notified Body,

Pikrou- Moraitaki Eleftheria

President and Managing Director of EKAPTY SA

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