

**Production Quality Assurance**  
Directive 93/42/EEC on Medical devices, Annex V

CE Certiso Ltd. (NB 2409) certifies that the following manufacturer's quality management system concerning to the listed devices and device categories meets the requirements of the related requirements of the directive.

Name of the manufacturer:

**BVK Konsalting D.o.o.**

Headquarters: **11000 Beograd, Kraljice Natalije 68/26, Serbia**  
Manufacturing plant: **Dragiše Mišovića 144/1, 32000 Čačak, Serbia**  
Authorised representative: **VEP CONSULTING Kft., Bécsi krt.23, 6722 Szeged, Hungary**

**VEP device**

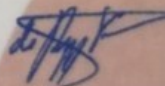
The certificate covers the following devices:

Description of the device	Intended use	Risk class
VEP device	induce and maintain erection in male sexual organ by increasing the blood flow in it	Ila

This certificate is valid only in case of successfully conducted annual surveillance audits.

ID number of the related audit report: 163-CE-170929

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Valter PAPP, Dr.  
General Manager

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