

## CERTIFICATE Production Quality Assurance Medical Devices Directive 93/42/EEC Annex V

Company Name	: Berika Teknoloji Medikal İmalat İth. Ihr. Tic. Ltd. Şti.	
Company Address	: Alakova Mah. Karaman Cad. No	5:812 Meram KONYA / TURKEY
Related Directives and Annex	: 93/42/EEC Medical Devices Direc	ctive - Annex V
Product	: Sterile Single Use Syringe - Class II - 1 Ml. 3 Pieces Steril With Needle/Wi - 2 Ml. 3 Pieces Steril With Needle/Wi - 3 Ml. 3 Pieces Steril With Needle/Wi - 5 Ml. 3 Pieces Steril With Needle/Wi - 10 Ml. 3 Pieces Steril With Needle/Wi - 20 Ml. 3 Pieces Steril With Needle/Wi - 50Ml. 3 Pieces Steril With Needle/Wi - 700 - 200	ithout Needle ithout Needle ithout Needle Vithout Needle Vithout Needle ithout Needle eedles - Class IIa
GMDN	: 47017, 59230, 58095	
Certificate Number	: M.2016.106.6915	
Report Number	: MD.3159.IB	APPROV
Initial Assessment Date	: 16.07.2016	Dagen.
Registration Date	: 05.08.2016	UDEM International Continication
Recertification Assessment Date		Auditing Training Centre Industry
Reissue Date / No	: 24.02.2020/01	and Trade Inc. Co.
Revision Date /No	:-	
Expiry Date	: 27.05.2024	*
UDEM hereby declares that the requirements of Annex V of the directive 93/42/EEC have been met for the listed products. The above named manufacturer has established and applies a quality assurance system, which is subject to periodic surveillance audits, defined by Annex V, section 4 of the dorementioned directive. UDEMs responsibility for class I devices covered by the EC sertificate is limited to manufacturing issues related to safeguarding and maintaining sterile conditions, if the dev,ce is sterile; and manufacturing issues related to product's conformity with		

products. The above named manufacturer has established and applies a quality assurance system, which is subject to periodic surveillance audits, defined by Annex V, section 4 of the aforementioned directive. UDEN's responsibility for class I devices covered by the EC settificate is limited to manufacturing issues related to safeguarding and maintaining sterile conditions, if the dev.ce is sterile; and manufacturing issues related to product's conformity with international Certification Auditing Training Centre Industry and Trade Inc. Co. to whom it must be returned upon request. The above named company and UDEN must keep a copy of this certificate for 5 years from the registration of the certificate. Usage of the CE mark is under the responsibility of the manufacturer with the completion of EC Declaration of Conformity. The above mentioned company must notify all changes related with the approved product to UDEN. If UDEN will not renew the expiry date of this certificate in question, the mentioned company should stop placing the product on the market. The validity of the certificate can be checked through www.udem.com. tr.

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