

CERTIFICATE

Full Quality Assurance System Medical Devices Directive 93/42/EEC Annex II (Excluding Section 4)

M.2019.106.11727-1 Design Examination Certificate Was Prepared for Class III Products Defined in This Certificate.

Company Name	: Shandong Haidike Medical Products Co., Ltd.
Company Address	: Plant No.1, Science and Technology Enterprise Incubator Park, Shan County, Heze City, Shandong Province, China.
Related Directives and Annex	: 93/42/EEC Medical Devices Directive - Annex II
	(Excluding Section 4)
Product	 Sterile Braided Coated Violet or Undyed Absorbable Polyglycolic Acid (PGA) Suture with or without Needle - Class III Sterile Braided Coated Violet or Undyed Absorbable Polyglactin (PGLA) Suture with or without Needle - Class III Sterile Single Use Non-Absorbable Silk Suture with or without needle - Class IIb Sterile Single Use Non-Absorbable Nylon Suture with or without needle - Class IIb Sterile Single Use Non-Absorbable Polyester Suture with or without needle - Class IIb Sterile Single Use Non-Absorbable Polyester Suture with or without needle - Class IIb Sterile Single Use Non-Absorbable Polypropylene Suture with or without needle - Class IIb
GMDN	: 13908, 17471, 13910, 13905, 13906, 13909
	Product Types are attached.
Certificate Number	: M.2019.106.11727
Report Number	: MD.3626.IB
Initial Assessment Date	: 10.02.2018
Registration Date	: 10.04.2019 UDEMInfernational Certification
Revision Date /No	: 22.07.2019/01 Auditing Training Centre Industry
Expiry Date	: 09.04.2024
for the lefed products. The above named manufacture subject to periodic surveillance audits, defined by Anne Lisection 4 an EC design - examination certificate is require	x1. excluding section 4 of the 93/42/EEC Directive have been metric in the setablished and applied a quality assurance system, which is ax1, section 5 of the forementioned directive, According to Annex, ad for placing the Class II devices on the market. UDEMs responsibility to manufacturing issues related to safeguarding and maintaining.

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tor the lated products, line above named manufacturer has established and applied a quality assumace system, which is subject to periodic surveillance audits, defined by Annex II, section 5 of the forementioned directive. According to Annex II, section 5 of the forementioned directive. According to Annex II, section 5 of the forementioned directive. According to Annex II, section 5 of the forementioned directive. According to Annex II, section 5 of the forementioned directive. According to Annex II, section 5 of the forementioned directive. According to Annex II, section 5 of the forementioned directive. According to Annex II, section 5 of the forementioned directive. According to Annex II in the section 5 of the foremention of the Context of the property of UDEM interactional Certification Auditing Training stells conditions. If the device is stells, and manufacturing issues related to product's conformity with methodogical equivements. If it has measurement function. This certificate remains as the property of UDEM interactional Certification Auditing Training Centre Industry and Trade Inc. Co. to whom it must be returned upon request. The above named company and UDEM 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 Contemity. The above mentioned company must notify all changes related with the approved product to UDEM. If UDEM will not renew the validity 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.

Address: Mutlukent Mahallesi 2073 Sokak (Eski 93 Sokak) No:10 Çankaya – Ankara – TURKEY Phone: +90 0312 443 03 90 Fax: +90 0312 443 03 76 E-mail: info@udemltd.com.tr www.udem.com.tr





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ISO 13485 : 2016

Shandong Haidike Medical Products Co., Ltd.

Plant No.1, Science and Technology Enterprise Incubator Park, Shan County, Heze City, Shandong Province, China.

This certificate shows that the medical devices - quality management system (EN ISO 13485:2016) of the above company was approved by PCA Certification for the following scope, the validity of the certificate depends on the company's pass the annual surveillance audits and company's maintenance the related management system conditions according to international accreditation criteria

SCOPE

Design, development, manufacture and sales of Disposable suture needles with non-absorbable threads, sterile syringe for single use, sterile infusion sets for single use, disposable lancets for blood specimen collection, absorbable surgical sutures with or without needle. Medical face mask and Medical protective clothing

GROUP CODE

Α

Certificate No Registration Date Reissue Date Expiry Date Certificate Period Exclusion

: 27.04.2018
: 19.04.2021/01
: 26.04.2024
: 3 Years (From the date of registration)
: 7.5.3

: TC-75059



Management Systems Certification Body

MSCB-103



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PCA Certification Approval

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