



Product Service

## Confirmation Statement on validity of EC Certificate (MDD)

pursuant to Directive 93/42/EEC concerning medical devices

**No. GCQ 066729 0004 Rev. 01**

**Manufacturer:**

**Jiangsu Suyun Medical Materials  
Co., Ltd.**

No.18 Jin Qiao Road  
Dapu Industrial Park  
222002 Lianyungang, Jiangsu Province  
PEOPLE'S REPUBLIC OF CHINA

This Confirmation Statement  
is only valid in combination  
with the following  
EC Certificate (MDD):

**G2S 043337 0044 Rev. 03**

This Confirmation Statement confirms the validity of the aforementioned EC Certificate (MDD).  
It considers clarification of scope statements, scope reductions and changes to the manufacturer  
data initiated 26 May 2021 or later.

The conditions laid down in Article 120 (3) of Regulation (EU) 2017/745 on medical devices for  
placing devices on the market and putting into service apply. For details and confirmation statement  
validity see: [www.tuvsud.com/ps-cert?q=cert:GCQ 066729 0004 Rev. 01](http://www.tuvsud.com/ps-cert?q=cert:GCQ_066729_0004_Rev_01)

**Report No.:**

BJ23088501

**Valid until:**

2024-04-13

Christoph Dicks  
Head of Certification/Notified Body

**Issue Date:** 2023-10-25



Product Service

**Confirmation Statement on validity of EC Certificate (MDD)**

pursuant to Directive 93/42/EEC concerning medical devices

**No. GCQ 066729 0004 Rev. 01**

**Product Category(ies):** Sterile Vaginal Speculum for Single Use, Sterile Gynaecological Collectors for Single Use, Sterile Rectal Catheter for Single Use, Sterile Nasal Speculum for Single Use, Sterile Urine Drainage Bag for Single Use, Sterile Applicator with Cotton Tip, Sterile Tongue Depressor, Sterile Otolaryngological Set for single use, Sterile Gynecological Kit for Single Use, Medical Transport Swab, Sterile Catheter Tip Syringe for Single Use, Sterile Extension Tube for Single Use, Sterile Laryngological Speculum for Single Use, Sterile Umbilical Cord Clamp for Single Use, Sterile Amniohook for Single Use, Ear funnel, Forceps, Ostomy bags and accessories. Sterile Holder Accessories for Single Use (Fastener, Arm Drape).

**Description of Change:**

Change of product name from Transport Swab to Medical Transport Swab.



Benannt durch/Designated by  
Zentralstelle der Länder  
für Gesundheitsschutz  
bei Arzneimitteln und  
Medizinprodukten  
www.zlg.de  
ZLG-BS-244.10.08



Product Service

## EC Certificate

Production Quality Assurance System

Directive 93/42/EEC on Medical Devices (MDD), Annex V

(Devices in class I in sterile conditions, sterilised systems or procedure packs)

**No. G2S 043337 0044 Rev. 03**

### Manufacturer

**Jiangsu Suyun Medical  
Materials Co., Ltd.**

No.1 Medicine Lane, Renmin Rd.

222002 Lianyungang

PEOPLE'S REPUBLIC OF CHINA

### EC-Representative:

Shanghai International Holding Corp. GmbH (Europe)  
Eiffestraße 80, 20537 Hamburg, GERMANY

### Product Category(ies):

**Sterile Non-active Medical devices (For  
detailed information see attachment).**

The Certification Body of TÜV SÜD Product Service GmbH declares that the aforementioned manufacturer has implemented a quality assurance system for manufacture in accordance with MDD Annex V. This quality assurance system covers those aspects of manufacture concerned with securing and maintaining sterile conditions of the respective devices / device categories and conforms to the requirements of this Directive. It is subject to periodical surveillance. See also notes overleaf.

**Report No.:**

BJ1888504

**Valid from:**

2019-05-27

**Valid until:**

2024-04-13

**Date,** 2019-05-27

Stefan Preiß

Head of Certification/Notified Body





Benannt durch/Designated by  
Zentralstelle der Länder  
für Gesundheitsschutz  
bei Arzneimitteln und  
Medizinprodukten  
www.zlg.de  
ZLG-BS-244.10.08



Product Service

## EC Certificate

Production Quality Assurance System

Directive 93/42/EEC on Medical Devices (MDD), Annex V

(Devices in class I in sterile conditions, sterilised systems or procedure packs)

**No. G2S 043337 0044 Rev. 03**

### Facility(ies):

Jiangsu Suyun Medical Materials Co., Ltd.  
No.1 Medicine Lane, Renmin Rd., 222002 Lianyungang,  
PEOPLE'S REPUBLIC OF CHINA

Jiangsu Suyun Medical Materials Co., Ltd  
No.18 Jin Qiao Road, Dapu Industrial Park, 222002 Lianyungang,  
Jiangsu Province, PEOPLE'S REPUBLIC OF CHINA

### Attachment:

Sterile Vaginal Speculum for Single Use, Sterile Gynaecological Collectors for Single Use, Sterile Rectal Catheter for Single Use, Sterile Nasal Speculum for Single Use, Sterile Urine Drainage Bag for Single Use, Sterile Applicator with Cotton Tip, Sterile Tongue Depressor, Sterile Otolaryngological Set for single use, Sterile Gynecological Kit for Single Use, Transport Swab, Sterile Catheter Tip Syringe for Single Use, Sterile Extension Tube for Single Use, Sterile Laryngological Speculum for Single Use, Sterile Umbilical Cord Clamp for Single Use, Sterile Amniohook for Single Use, Ear funnel, Forceps, Ostomy bags and accessories. Sterile Holder Accessories for Single Use (Fastener, Arm Drape).



**Add value.  
Inspire trust.**

TÜV SÜD Product Service GmbH · Ridlerstr. 65 · 80339 Munich · Germany

Jiangsu Suyun Medical Materials  
Co., Ltd.  
Dapu Industrial Park  
No.18 Jin Qiao Road  
222002 LIANYUNGANG, JIANGSU PROVINCE  
PEOPLE'S REPUBLIC OF CHINA

Your reference/letter of	Our reference/name	Tel. extension	Email	Date	Page
CBW 66729	GCN-BJ23885A02	+86-10-6590-6186	Xingchun.Li@tuvsud.com	2024-02-06	1 of 8

**TÜV SÜD Product Service GmbH  
Confirmation Letter  
CL 066729 0006 Rev. 00**

**Reference: GCN-BJ23885A02**

To whom it may concern,

**Confirmation of the status of a formal application, written agreement, and appropriate surveillance in the framework of Regulation EU 2023/607 amending Regulations (EU) 2017/745 (in the following referenced as MDR) as regards the transitional provisions for certain medical devices and in vitro diagnostic medical devices.**

With this letter TÜV SÜD Product Service GmbH, designated under MDR and identified by the number 0123 on NANDO, confirms that we have received a formal application in accordance with Section 4.3, first subparagraph of Annex VII of MDR and has signed a written agreement in accordance with Section 4.3, second subparagraph of Annex VII of MDR with the above stated manufacturer with the following

SRN Number: CN-MF-000021132

The devices covered by the formal application and the written agreement mentioned above are identified in the Tables below.

- Table 1 identifies the devices for which an MDR application has been received, written agreement concluded and for which TÜV SÜD Product Service GmbH is also responsible for appropriate surveillance of the corresponding devices under the applicable Directive.
- Table 2 identifies the devices for which an MDR application has been received and a written agreement concluded, but TÜV SÜD Product Service GmbH has not yet taken the responsibility for appropriate surveillance of the corresponding devices under the applicable Directive.

**Registered Office: Munich**  
Trade Register Munich HRB 85 742  
UniCredit Bank AG · BIC HYVEDEMMXXX  
IBAN DE13 7002 0270 0048 8522 11  
VAT ID No. DE129484267  
Information pursuant to § 2 [1] DL-InfoV  
(Germany) at tuvsud.com/imprint

**Supervisory Board:**  
Holger Lindner (Chairman)  
**Board of Management:**  
Walter Reithmaier (CEO)  
Patrick van Welij

**TÜV SÜD Product Service GmbH**  
Ridlerstr. 65  
80339 Munich  
Germany

**tuvsud.com/ps**  
Hotline: +49 89 50084-747





If devices covered by certificates issued under Directive 90/385/EEC (AIMDD) or Directive 93/42/EEC (MDD) that expired after 26 May 2021 and before 20 March 2023, without having been withdrawn, this letter also confirms that

- the manufacturer signed the written agreement under MDR by the date of MDD/AIMDD certificate expiry; or
- provided evidence that a competent authority of a Member State had granted a derogation or exemption from the applicable conformity assessment procedure in accordance with Article 59(1) of MDR or Article 97(1) of the MDR respectively.

The transition timelines in accordance Article 120 (3) of MDR that apply to the devices covered by this letter, subject to the manufacturer's continued compliance to the other conditions specified in Article 120 (3c) of MDR, are shown below:

- 26 May 2026 for Class III custom-made implantable devices
- 31 December 2027 for Class III devices and Class IIb implantable devices (except sutures, staples, dental fillings, dental braces, tooth crowns, screws, wedges, plates, wires, pins, clips and connectors)
- 31 December 2028 for other Class IIb devices, Class IIa, Class I devices placed on the market in sterile condition, measuring function
- 31 December 2028 for devices not requiring the involvement of a notified body under MDD but requiring it under MDR (e.g., class I devices that qualify as re-usable surgical instruments)

We reserve the right to invoice any issuance, copies, amendments and / or changes of the confirmation letter according to effort.

For confirmation letter validity see [www.tuvsud.com/ps-cert?q=cert:CL 066729 0006 Rev. 00](http://www.tuvsud.com/ps-cert?q=cert:CL 066729 0006 Rev. 00)

In case of inquiries please contact [medical\\_devices@tuvsud.com](mailto:medical_devices@tuvsud.com).

On behalf of the Notified Body TÜV SÜD Product Service GmbH,  
06.02.2024

TÜV SÜD Product Service GmbH  
Medical and Health Services

A handwritten signature in black ink, appearing to read 'LmLi'.

Mr. Li Xingchun  
Conformity Assessment Responsible (CARE)

TÜV SÜD Product Service GmbH  
Medical and Health Services

A handwritten signature in blue ink, appearing to read 'A. Fazlija'.

Arianit Fazlija  
2024.02.06 10:23:39 +01'00'

Arianit Fazlija  
Application Reviewer



**Table 1: Devices covered by this letter and for which TÜV SÜD Product Service GmbH is also responsible for appropriate surveillance of the corresponding devices under the applicable Directive:**

<b>Device name or Basic UDI-DI (under MDR application)</b>	<b>MDR Device classification (as proposed by the manufacturer and verified during application review)</b>	<b>If the MDR device is a substitute device, identification of the corresponding MDD/AIMDD device</b>	<b>MDD/AIMDD Certificate Reference(s) of the devices under MDR application, and the NB Identification</b>
<b>Sterile Vaginal Speculum for single use 69332982004</b>	<input type="checkbox"/> Class III <input type="checkbox"/> Class IIb implantable (non-exempted) <input type="checkbox"/> Class IIb / Class IIb implantable (exempted) <input type="checkbox"/> Class IIa <input checked="" type="checkbox"/> Class I devices in sterile condition <input type="checkbox"/> Class I devices with measuring function <input type="checkbox"/> Class III implantable custom-made-device	<input checked="" type="checkbox"/> N/A	<input checked="" type="checkbox"/> Certification as follows: No.G2S 043337 0044 Rev.03; No.GCQ066729 0004 Rev.01; NB#0123
<b>Sterile Gynaecological Collectors for single use 69332982028</b>	<input type="checkbox"/> Class III <input type="checkbox"/> Class IIb implantable (non-exempted) <input type="checkbox"/> Class IIb / Class IIb implantable (exempted) <input type="checkbox"/> Class IIa <input checked="" type="checkbox"/> Class I devices in sterile condition <input type="checkbox"/> Class I devices with measuring function <input type="checkbox"/> Class III implantable custom-made-device	<input checked="" type="checkbox"/> N/A	<input checked="" type="checkbox"/> Certification as follows: No.G2S 043337 0044 Rev.03; No.GCQ066729 0004 Rev.01; NB#0123
<b>Sterile Rectal Catheter for single use 69332982011</b>	<input type="checkbox"/> Class III <input type="checkbox"/> Class IIb implantable (non-exempted) <input type="checkbox"/> Class IIb / Class IIb implantable (exempted) <input type="checkbox"/> Class IIa <input checked="" type="checkbox"/> Class I devices in sterile condition <input type="checkbox"/> Class I devices with measuring function <input type="checkbox"/> Class III implantable custom-made-device	<input checked="" type="checkbox"/> N/A	<input checked="" type="checkbox"/> Certification as follows: No.G2S 043337 0044 Rev.03; No.GCQ066729 0004 Rev.01; NB#0123
<b>Sterile Nasal Speculum for Single Use 69332982016</b>	<input type="checkbox"/> Class III <input type="checkbox"/> Class IIb implantable (non-exempted) <input type="checkbox"/> Class IIb / Class IIb implantable (exempted) <input type="checkbox"/> Class IIa <input checked="" type="checkbox"/> Class I devices in sterile condition	<input checked="" type="checkbox"/> N/A	<input checked="" type="checkbox"/> Certification as follows: No.G2S 043337 0044 Rev.03; No.GCQ066729 0004 Rev.01; NB#0123



Device name or Basic UDI-DI (under MDR application)	MDR Device classification (as proposed by the manufacturer and verified during application review)	If the MDR device is a substitute device, identification of the corresponding MDD/AIMDD device	MDD/AIMDD Certificate Reference(s) of the devices under MDR application, and the NB Identification
	<input type="checkbox"/> Class I devices with measuring function <input type="checkbox"/> Class III implantable custom-made-device		
<b>Sterile Urine Drainage Bag for single use</b> <b>69332982019</b>	<input type="checkbox"/> Class III <input type="checkbox"/> Class IIb implantable (non-exempted) <input type="checkbox"/> Class IIb / Class IIb implantable (exempted) <input type="checkbox"/> Class IIa <input checked="" type="checkbox"/> Class I devices in sterile condition <input type="checkbox"/> Class I devices with measuring function <input type="checkbox"/> Class III implantable custom-made-device	<input checked="" type="checkbox"/> N/A	<input checked="" type="checkbox"/> Certification as follows: No.G2S 043337 0044 Rev.03; No.GCQ066729 0004 Rev.01; NB#0123
<b>Sterile Applicator with Cotton Tip</b> <b>69332982066</b>	<input type="checkbox"/> Class III <input type="checkbox"/> Class IIb implantable (non-exempted) <input type="checkbox"/> Class IIb / Class IIb implantable (exempted) <input type="checkbox"/> Class IIa <input checked="" type="checkbox"/> Class I devices in sterile condition <input type="checkbox"/> Class I devices with measuring function <input type="checkbox"/> Class III implantable custom-made-device	<input checked="" type="checkbox"/> N/A	<input checked="" type="checkbox"/> Certification as follows: No.G2S 043337 0044 Rev.03; No.GCQ066729 0004 Rev.01; NB#0123
<b>Sterile Tongue Depressor</b> <b>69332982065</b>	<input type="checkbox"/> Class III <input type="checkbox"/> Class IIb implantable (non-exempted) <input type="checkbox"/> Class IIb / Class IIb implantable (exempted) <input type="checkbox"/> Class IIa <input checked="" type="checkbox"/> Class I devices in sterile condition <input type="checkbox"/> Class I devices with measuring function <input type="checkbox"/> Class III implantable custom-made-device	<input checked="" type="checkbox"/> N/A	<input checked="" type="checkbox"/> Certification as follows: No.G2S 043337 0044 Rev.03; No.GCQ066729 0004 Rev.01; NB#0123
<b>Sterile Otolaryngological Set for Single use</b> <b>69332982069</b>	<input type="checkbox"/> Class III <input type="checkbox"/> Class IIb implantable (non-exempted) <input type="checkbox"/> Class IIb / Class IIb implantable (exempted) <input type="checkbox"/> Class IIa	<input checked="" type="checkbox"/> N/A	<input checked="" type="checkbox"/> Certification as follows: No.G2S 043337 0044 Rev.03; No.GCQ066729 0004 Rev.01; NB#0123





Device name or Basic UDI-DI (under MDR application)	MDR Device classification (as proposed by the manufacturer and verified during application review)	If the MDR device is a substitute device, identification of the corresponding MDD/AIMDD device	MDD/AIMDD Certificate Reference(s) of the devices under MDR application, and the NB Identification
	<input checked="" type="checkbox"/> Class I devices in sterile condition <input type="checkbox"/> Class I devices with measuring function <input type="checkbox"/> Class III implantable custom-made-device		
<b>Sterile Gynecological kit for single Use</b> <b>69332982057</b>	<input type="checkbox"/> Class III <input type="checkbox"/> Class IIb implantable (non-exempted) <input type="checkbox"/> Class IIb / Class IIb implantable (exempted) <input type="checkbox"/> Class IIa <input checked="" type="checkbox"/> Class I devices in sterile condition <input type="checkbox"/> Class I devices with measuring function <input type="checkbox"/> Class III implantable custom-made-device	<input checked="" type="checkbox"/> N/A	<input checked="" type="checkbox"/> Certification as follows: No.G2S 043337 0044 Rev.03; No.GCQ066729 0004 Rev.01; NB#0123
<b>Medical Transport swab</b> <b>69332982073</b>	<input type="checkbox"/> Class III <input type="checkbox"/> Class IIb implantable (non-exempted) <input type="checkbox"/> Class IIb / Class IIb implantable (exempted) <input type="checkbox"/> Class IIa <input checked="" type="checkbox"/> Class I devices in sterile condition <input type="checkbox"/> Class I devices with measuring function <input type="checkbox"/> Class III implantable custom-made-device	<input checked="" type="checkbox"/> N/A	<input checked="" type="checkbox"/> Certification as follows: No.G2S 043337 0044 Rev.03; No.GCQ066729 0004 Rev.01; NB#0123
<b>Sterile Catheter Tip Syringe for Single Use</b> <b>69332982017</b>	<input type="checkbox"/> Class III <input type="checkbox"/> Class IIb implantable (non-exempted) <input type="checkbox"/> Class IIb / Class IIb implantable (exempted) <input type="checkbox"/> Class IIa <input checked="" type="checkbox"/> Class I devices in sterile condition <input type="checkbox"/> Class I devices with measuring function <input type="checkbox"/> Class III implantable custom-made-device	<input checked="" type="checkbox"/> N/A	<input checked="" type="checkbox"/> Certification as follows: No.G2S 043337 0044 Rev.03; No.GCQ066729 0004 Rev.01; NB#0123
<b>Sterile Extension Tube for single use</b> <b>69332982074</b>	<input type="checkbox"/> Class III <input type="checkbox"/> Class IIb implantable (non-exempted) <input type="checkbox"/> Class IIb / Class IIb implantable (exempted)	<input checked="" type="checkbox"/> N/A	<input checked="" type="checkbox"/> Certification as follows: No.G2S 043337 0044 Rev.03; No.GCQ066729 0004 Rev.01; NB#0123



Device name or Basic UDI-DI (under MDR application)	MDR Device classification (as proposed by the manufacturer and verified during application review)	If the MDR device is a substitute device, identification of the corresponding MDD/AIMDD device	MDD/AIMDD Certificate Reference(s) of the devices under MDR application, and the NB Identification
	<input type="checkbox"/> Class IIa <input checked="" type="checkbox"/> Class I devices in sterile condition <input type="checkbox"/> Class I devices with measuring function <input type="checkbox"/> Class III implantable custom-made-device		
<b>Sterile Laryngological Speculum for Single use</b> <b>69332982083</b>	<input type="checkbox"/> Class III <input type="checkbox"/> Class IIb implantable (non-exempted) <input type="checkbox"/> Class IIb / Class IIb implantable (exempted) <input type="checkbox"/> Class IIa <input checked="" type="checkbox"/> Class I devices in sterile condition <input type="checkbox"/> Class I devices with measuring function <input type="checkbox"/> Class III implantable custom-made-device	<input checked="" type="checkbox"/> N/A	<input checked="" type="checkbox"/> Certification as follows: No.G2S 043337 0044 Rev.03; No.GCQ066729 0004 Rev.01; NB#0123
<b>Sterile Umbilical Cord Clamp for single use</b> <b>69332982080</b>	<input type="checkbox"/> Class III <input type="checkbox"/> Class IIb implantable (non-exempted) <input type="checkbox"/> Class IIb / Class IIb implantable (exempted) <input type="checkbox"/> Class IIa <input checked="" type="checkbox"/> Class I devices in sterile condition <input type="checkbox"/> Class I devices with measuring function <input type="checkbox"/> Class III implantable custom-made-device	<input checked="" type="checkbox"/> N/A	<input checked="" type="checkbox"/> Certification as follows: No.G2S 043337 0044 Rev.03; No.GCQ066729 0004 Rev.01; NB#0123
<b>Sterile Amniohook for single use</b> <b>69332982044</b>	<input type="checkbox"/> Class III <input type="checkbox"/> Class IIb implantable (non-exempted) <input type="checkbox"/> Class IIb / Class IIb implantable (exempted) <input type="checkbox"/> Class IIa <input checked="" type="checkbox"/> Class I devices in sterile condition <input type="checkbox"/> Class I devices with measuring function <input type="checkbox"/> Class III implantable custom-made-device	<input checked="" type="checkbox"/> N/A	<input checked="" type="checkbox"/> Certification as follows: No.G2S 043337 0044 Rev.03; No.GCQ066729 0004 Rev.01; NB#0123
<b>Ostomy bags and accessories</b> <b>69332982163</b>	<input type="checkbox"/> Class III <input type="checkbox"/> Class IIb implantable (non-exempted)	<input checked="" type="checkbox"/> N/A	<input checked="" type="checkbox"/> Certification as follows: No.G2S 043337 0044 Rev.03;



Device name or Basic UDI-DI (under MDR application)	MDR Device classification (as proposed by the manufacturer and verified during application review)	If the MDR device is a substitute device, identification of the corresponding MDD/AIMDD device	MDD/AIMDD Certificate Reference(s) of the devices under MDR application, and the NB Identification
	<input type="checkbox"/> Class IIb / Class IIb implantable (exempted) <input type="checkbox"/> Class IIa <input checked="" type="checkbox"/> Class I devices in sterile condition <input type="checkbox"/> Class I devices with measuring function <input type="checkbox"/> Class III implantable custom-made-device		No.GCQ066729 0004 Rev.01; NB#0123
<b>Forceps 69332982041</b>	<input type="checkbox"/> Class III <input type="checkbox"/> Class IIb implantable (non-exempted) <input type="checkbox"/> Class IIb / Class IIb implantable (exempted) <input type="checkbox"/> Class IIa <input checked="" type="checkbox"/> Class I devices in sterile condition <input type="checkbox"/> Class I devices with measuring function <input type="checkbox"/> Class III implantable custom-made-device	<input checked="" type="checkbox"/> N/A	<input checked="" type="checkbox"/> Certification as follows: No.G2S 043337 0044 Rev.03; No.GCQ066729 0004 Rev.01; NB#0123
<b>Ear funnel 69332982043</b>	<input type="checkbox"/> Class III <input type="checkbox"/> Class IIb implantable (non-exempted) <input type="checkbox"/> Class IIb / Class IIb implantable (exempted) <input type="checkbox"/> Class IIa <input checked="" type="checkbox"/> Class I devices in sterile condition <input type="checkbox"/> Class I devices with measuring function <input type="checkbox"/> Class III implantable custom-made-device	<input checked="" type="checkbox"/> N/A	<input checked="" type="checkbox"/> Certification as follows: No.G2S 043337 0044 Rev.03; No.GCQ066729 0004 Rev.01; NB#0123

**Table 2: Devices covered by this letter and for which TÜV SÜD Product Service GmbH is NOT responsible for appropriate surveillance of the corresponding devices under the applicable Directive:**

Device name or Basic UDI-DI (under MDR application)	MDR Device classification (as proposed by the manufacturer and verified during application review)	If the MDR device is a substitute device, identification of the corresponding MDD/AIMDD device	MDD/AIMDD Certificate Reference(s) of the devices under MDR application, and the NB Identification
<input checked="" type="checkbox"/> N/A	<input checked="" type="checkbox"/> N/A	<input checked="" type="checkbox"/> N/A	<input checked="" type="checkbox"/> N/A



**Confirmation Letter Version History**

Date	TÜV SÜD Product Service GmbH internal reference traceable to each version of the letter	Action
2024/02/06	GCN-BJ23885A02	Initial issue

# DECLARATION OF CONFORMITY TO COUNCIL DIRECTIVE 2007/47/EEC CONCERNING MEDICAL DEVICES

SYC/JY-CE28-0001

Edition:C/0



MANUFACTURER:

JIANGSU SUYUN MEDICAL MATERIALS Co., LTD

No.1 MEDICINE LANE, RENMIN ROAD.222002 LIANYUNGANG, PEOPLE'S REPUBLIC OF CHINA

No.18 JINQIAO ROAD OF DAPU ECONMIC DEVELOPMENT ZONE, LIANYUNGANG, CHINA

MEDICAL DEVICE:

STERILE GYNAECOLOGICAL COLLECTORS FOR SINGLE USE

CERVICAL RAMBRUSH---CODE420211, CERVICAL RAMBRUSH BALL---CODE420222  
CERVICAL SPOON---CODE420501 CERVIX BRUSH PLUSH---CODE420151 CERVIX CELL BRUSH---CODE477701-IIID  
SPATULA WOODEN---CODE 425002, CERVICAL SPATULA---CODE420301-IIIB3,CODE420311-IIIB4  
ENDOMETRIAL SUCTION CURETTE-----CODE420411 (ONE HOLE) CODE420414(FOUR HOLES)

CLASSIFICATION - ANNEX IX:

CLASS I STERILE, RULE 5

CONFORMITY ASSESSMENT ROUTE:

ANNEX V+ VIII

WE, JIANGSU SUYUN MEDICAL MATERIALS Co.,LTD, HEREWITH DECLARE THAT THE STATED MEDICAL DEVICES MEET THE TRANSPOSITION INTO NATIONAL LAW, THE PROVISIONS OF COUNCIL DIRECTIVE 93/42/EEC OF 14 JUNE 1993 CONCERNING MEDICAL DEVICES; INCLUDING, AT 21 MARCH 2010, THE AMENDMENTS BY COUNCIL DIRECTIVE 2007/47/EEC. ALL SUPPORTING DOCUMENTATION IS RETAINED AT THE PREMISES OF THE MANUFACTURER.

STANDARDS APPLIED: EN ISO13485:2016,EN ISO 14971:2012,EN ISO 11135-1:2007,EN ISO 11607-1:2017,EN ISO 11607-2:2017,EN ISO 10993-1:2009/AC: 2010,EN ISO 10993-5:2009,EN ISO 10993-7:2008/AC: 2009,EN ISO 10993-10:2013,EN 1041:2008,EN ISO 15223-1:2016,EN 1422:2014,EN ISO 11737-1:2018,EN ISO 14155:2011, EN ISO 11138-2:2017

NOTIFIED BODY:

TÜV SÜD PRODUCT SERVICE GMBH  
RIDLERSTR 65, D-80339 MÜNCHEN, GERMANY

IDENTIFICATION NUMBER

CE 0123

(EC) CERTIFICATE(S):

G2S 043337 0044 Rev.03



EUROPEAN REPRESENTATIVE:

Shanghai International Holding Corp. GmbH (Europe)  
Eiffestrasse, 80, D-20537, Hamburg, Germany Tel: 0086-021-65951371, 0049-40-2513175, Fax: 0049-40-255726

START OF CE-MARKING:

DATE OF FIRST CE MARKING  
17.06.2005

PLACE, DATE OF DECLARATION:

LIANYUNGANG, JIANGSU; 18.09.2019

SIGNATURE:

江苏苏云医疗器材有限公司  
JIANGSU SUYUN MEDICAL MATERIALS CO.,LTD.  
NAME: QIN HONGPING  
POSITION: BOARD OF CHAIRMAN





# Certificate

No. Q5 066729 0001 Rev. 02

**Holder of Certificate:** **Jiangsu Suyun Medical Materials Co., Ltd.**

No.18 Jin Qiao Road  
Dapu Industrial Park  
222002 Lianyungang, Jiangsu Province  
PEOPLE'S REPUBLIC OF CHINA

**Certification Mark:**



**Scope of Certificate:** **Design and development, production and sales of Sterile /Non-sterile Non-active Medical devices (For detailed information see attachment).**

The Certification Body of TÜV SÜD Product Service GmbH certifies that the company mentioned above has established and is maintaining a quality management system, which meets the requirements of the listed standard(s). All applicable requirements of the testing and certification regulation of TÜV SÜD Group have to be complied with. For details and certificate validity see: [www.tuvsud.com/ps-cert?q=cert:Q5 066729 0001 Rev. 02](http://www.tuvsud.com/ps-cert?q=cert:Q5_066729_0001_Rev_02)

**Report No.:** BJ23088501

**Valid from:** 2023-10-25

**Valid until:** 2025-05-31

**Date,** 2023-10-25

Christoph Dicks  
Head of Certification/Notified Body



# Certificate

**No. Q5 066729 0001 Rev. 02**

**Applied Standard(s):**

ISO 13485:2016  
(EN ISO 13485:2016/AC:2018, EN ISO 13485:2016/A11:2021)  
Medical devices - Quality management systems -  
Requirements for regulatory purposes

**Facility(ies):**

**Jiangsu Suyun Medical Materials Co., Ltd.**  
No.18 Jin Qiao Road, Dapu Industrial Park, 222002 Lianyungang,  
Jiangsu Province, PEOPLE'S REPUBLIC OF CHINA

Design and development, production and sales of Sterile /Non-sterile Non-active Medical devices (For detailed information see attachment).

**Jiangsu Suyun Medical Materials Co., Ltd.**

No.1 Medicine Lane, Renmin Rd., 222002 Lianyungang,  
PEOPLE'S REPUBLIC OF CHINA

Production of Sterile /Non-sterile Non-active Medical devices (For detailed information see attachment).

Attachment: Sterile Infusion Sets for Single Use, Sterile Transfusion Sets for Single Use, Sterile Syringe for Single Use, Sterile Vaginal Speculum for Single Use, Sterile Urinary Catheter for Single Use (Nelaton Catheter, Foley Catheter and Silicone Catheter), Sterile Gynaecological Collectors for Single Use, Sterile Rectal Catheter for Single Use, Sterile Scalp Vein Sets for Single Use, Sterile Suction Catheter for Single Use, Sterile Blood Collection Needle for Single Use, Sterile Arterial Venous Fistula Needle for Single Use, Sterile Nasal Oxygen Cannula for Single Use, Sterile Proctoscope for Single Use, Laryngeal Mask Airway, Sterile Feeding Catheter for Single Use, Sterile Endotracheal Tube for Single Use, Sterile Insulin Syringe for Single Use, Sterile Stomach Tube with Guide Wire for Single Use, Sterile Stomach Tube for Single Use (Nasogastric Tube, Levin Tube), Sterile Duodenal Tube for Single Use, Sterile Nasal Speculum for Single Use, Sterile Urine Drainage Bag for Single Use, Sterile Feeding Tube for Single Use, Sterile Specimen Container for Single Use, Sterile Applicator with Cotton Tip, Sterile Tongue Depressor, Sterile Otolaryngological Set for Single Use, Trial Lenses Sets, Disposable Curettes, Sterile Gynecological Kit for Single Use, Medical Transport Swab, Heat and Moisture Exchange (Artificial Nose), Sterile Catheter Tip Syringe for Single Use, Intrauterine Insemination Catheter (IUI Cannula), Disposable Uterine Sound (Hysterometers), Sterile Hypodermic Needle for Single Use, Sterile Extension Tube for Single Use, Medical Disposable Bibs, Medical I.D. Bracelet, Sterile Laryngological Speculum for Single use, Sterile Umbilical Cord Clamp for Single Use, Sterile Dental Injection Needles for Single Use, Sterile Amniohook for Single Use, Sterile Intravascular Catheters for Single Use (I.V. Cannula), Suction catheter Kit, Sterile Incision Retractor for Single Use, Light-resistant Infusion Sets for Single Use, Ostomy bags and accessories, Vacuum Blood Collection Tubes, Medical Alcohol Pads, Forceps, Ear funnel, Disposable Circumcision Stapler, Sterile Holder Accessories for Single Use (Fastener, Arm Drape).





Product Service

# 认证证书

证书号. Q5 066729 0001 Rev. 02

证书持有者：

江苏苏云医疗器材有限公司

中华人民共和国江苏省连云港大浦工业园金桥路18号 222002

认证标志：



认证范围：

设计和开发，生产，销售：无菌/非无菌无源医疗器械  
(详细信息见附件)

认证机构TÜV SÜD产品服务有限公司证明上述公司已经建立并运行了满足所列标准要求的质量管理体系。

TÜV 南德集团检测及认证规则所有适用要求也须得到遵守。详情及证书有效期请见

[www.tuvsud.com/ps-cert?q=cert:Q5 066729 0001 Rev. 02](http://www.tuvsud.com/ps-cert?q=cert:Q5 066729 0001 Rev. 02)

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Christoph Dicks

Head of Certification/Notified Body

# 认证证书

证书号. Q5 066729 0001 Rev. 02

认证标准：

ISO 13485:2016

(EN ISO 13485:2016/AC:2018, EN ISO 13485:2016/A11:2021)

医疗器械 - 质量管理体系 - 用于法规的要求

生产场地：

江苏苏云医疗器材有限公司

中华人民共和国江苏省连云港大浦工业园金桥路18号 222002

设计和开发，生产，销售：无菌/非无菌无源医疗器械

(详细信息见附件)。

江苏苏云医疗器材有限公司

中华人民共和国连云港市人民路制药巷1号 222002

生产：无菌/非无菌无源医疗器械 (详细信息见附件)

附件：一次性使用无菌输液器，一次性使用无菌输血器，一次性使用无菌注射器，一次性使用无菌阴道扩张器，一次性使用无菌导尿管（导尿管、乳胶导尿管、硅胶导尿管），一次性使用无菌妇科取样器，一次性使用无菌肛管，一次性使用无菌静脉输液针，一次性使用无菌吸痰管，一次性使用无菌血样采集针，一次性使用无菌动静脉穿刺针，一次性使用无菌鼻氧管，一次性使用无菌肛门镜，医用喉罩，一次性使用无菌鼻饲管，一次性使用无菌气管插管，一次性使用无菌胰岛素注射器，一次性使用无菌导丝引导型胃肠管，一次性使用无菌胃管（鼻胃管，胃管），一次性使用无菌十二指肠管，一次性使用无菌鼻镜，一次性使用无菌尿袋，一次性使用无菌肠给养器，一次性使用无菌集样杯，无菌医用棉签，无菌医用压舌板，一次性使用无菌耳鼻喉包，验光镜片箱，引流管，一次性使用无菌扩阴器包，医用试管拭子，热湿交换器（人工鼻），一次性使用无菌灌注器，授精管（授精管），一次性使用子宫探测针（子宫探测针），一次性使用无菌注射针，一次性使用无菌延长管，医用口水巾，医用识别带，一次性使用无菌喉镜，一次性使用无菌脐带夹，一次性使用无菌牙科注射针，一次性使用无菌羊水勾，一次性使用无菌静脉留置针（静脉留置针），吸痰包，一次性使用无菌切口牵开保护器，一次性使用避光输液器，造口袋和附件，真空采血管，医用酒精消毒片，耳镜，镊子，一次性使用包皮环切器，一次性使用无菌扶持器配件包（紧固件、手臂）。