

Федеральное агентство по техническому регулированию и метрологии

НОПСС

Система добровольной сертификации "НОПСС". РОСС RU.31827.04ЖСН1
Орган по сертификации ООО "Невский Альянс". ОГРН 1147847286960 ИНН 7842525530
www.nopss.ru

СЕРТИФИКАТ СООТВЕТСТВИЯ

выдан

Общество с ограниченной ответственностью
«МиниМед»

ИНН 3234007127 / ОГРН 1023202138332

241520, Брянская область, Брянский район, с. Супонево, ул. Шоссейная, д.17А

Подтверждает что система менеджмента качества
соответствует требованиям ГОСТ ISO 9001-2015 (ISO 9001:2015)

При осуществлении работ согласно приложению №1 к настоящему сертификату

Сертификат выдан на основании решения экспертной комиссии

от 24.09.2018

Срок действия до 24 сентября 2021

Номер в едином реестре системы С1256

Руководитель органа
по сертификации:



Подпись

Платонов Б.А.



Настоящий сертификат обязывает организацию поддерживать состояние выполняемых работ в соответствии с вышеуказанным стандартом, что будет находиться под контролем органа по сертификации СДС "НОПСС" и подтверждаться при прохождении ежегодного инспекционного контроля.

Федеральное агентство по техническому регулированию и метрологии

НОПСС

Система добровольной сертификации "НОПСС". РОСС RU.31827.04ЖСН1

Орган по сертификации ООО "Невский Альянс". ОГРН 1147847286960 ИНН 7842525530

www.nopss.ru

ПРИЛОЖЕНИЕ №1

К сертификату соответствия № С1256

Применительно к видам деятельности :

Производство лабораторной посуды, медицинских изделий, приборов и принадлежностей, красителей, реагентов и наборов реагентов для in-vitro диагностики.

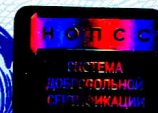
НОПСС

Руководитель органа
по сертификации:



Подпись

Платонов Б.А.



CERTIFICATO N° 505SGQ03

CERTIFICATE N° 505SGQ03

Si certifica che il
this is to certify that

Sistema di Gestione per la Qualità

Quality Management System

messo in atto da
implemented by

NUOVA APTACA S.r.l.

Via Monte Bianco, 4 – IT 20900 MONZA (MB)

nella Sede Operativa di
Operative Unit

Regione Monforte, 30 – IT 14053 CANELLI (AT)

è conforme alla norma
is in compliance with the standard

UNI EN ISO 9001-2015 (ISO 9001-2015)

per i seguenti Processi
concerning the following kinds of Processes

Gestione della fabbricazione ed immissione in commercio di tamponi sterili
per il prelievo di campioni biologici in orifizi naturali e in ambito chirurgico.
Progettazione e fabbricazione di dispositivi medico diagnostici per laboratori di analisi.
Commercializzazione di dispositivi medici e diagnostici in vitro.

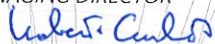
Commercializzazione di articoli da laboratorio

*Management of the manufacturing and placing on the market of sterile tampons for sampling of biological specimens
in natural orifice and in surgical field. Design and manufacturing of diagnostic medical devices for laboratories
of analysis. Marketing of medical and diagnostic devices in vitro. Marketing of laboratory articles.*

Il presente Certificato è soggetto al rispetto delle condizioni stabilite dai Regolamenti per la certificazione in vigore applicabili.
This Certificate shall satisfy the requirements established in the Rules for the certification in force applicable.

In caso di discordanza tra le lingue utilizzate nella traduzione del contenuto del presente certificato, fare riferimento alla lingua italiana.
In cases of discrepancy between the languages used in the translation of the content of this certificate, please refer to the Italian language

L'AMMINISTRATORE DELEGATO
MANAGING DIRECTOR


Dr. Ing. Roberto Cusolito

Data di Prima Emissione
First Issue Date

1998-07-23

Settore IAF 14 - 29

Data di Prima Emissione ITALCERT
First Issue Date ITALCERT

2011-10-30

Data di Rinnovo
Renewal Date

2017-10-30

Data di Scadenza
Expiration Date

2020-10-29



SGQ N° 023A PRD N° 122B
SGA N° 020D ISP N° 075E
PRS N° 097C
Membro degli Accordi di Mutuo Riconoscimento EA, IAF e ILAC
Signatory of EA, IAF and ILAC Mutual Recognition Agreements

DECLARATION OF CONFORMITY

Forlì, 19th April 2012

The device named " ECG SUPERGEL" (internal code 10158) has been produced by the company Ceracarta Spa on the basis of the essential requirements, see enclosure I of the directive 93/42/CEE, as prescribed in attachment VII of the above directive.

The writing company Ceracarta located in Via secondo Casadei , 14 Forlì, manufacturer of the product named , " ECG SUPERGEL ", declares under its own responsibility that such a device satisfies all the requirements of directive 93/42/CEE as amended by 2007/47/EC , about medical devices and in particular that:

- the Dispositive in object satisfies the essential requirements as in enclosure I of Directive 93/42/CEE;
- the Dispositive in object must be considered as belonging to Class I;
- the Dispositive in object must be exclusively used together with electro-medical instruments for recording, diagnosis and therapy, which base their functioning upon the measuring of energy flows of electric, magnetic and ultrasound type;
- The manufacturer has prepared and keeps the technical files updated in accordance with enclosure VII, section 3 of the directive itself.
- Such documentation is available at the headquarters of Ceracarta , for any reference by the entitled bodies.

CERACARTA spa

IL PRESIDENTE

Marino Bandini





Общество с ограниченной ответственностью
«Научно-производственная фирма «ВИНАР»
Юр.адрес: 105094, г. Москва, ул. Госпитальный вал, д.5, стр.7А, пом.VIII
Для писем: 105094, г. Москва, а/я 26
тел/факс: (495) 988-76-67, 360-61-46, 360-72-19
http://www.vinar.ru e-mail: main@vinar.ru

Общество с ограниченной ответственностью «Научно-производственная фирма «ВИНАР»

ПАСПОРТ КАЧЕСТВА

Индикаторы химические одноразовые воздушной стерилизации МедИС-В-Винар по ТУ 9398-032-11764404-2004

Регистрационное удостоверение
№ ФСР 2009/05017
от 27.08.2019 г.

Сертификат соответствия
№ РОСС RU. ИМ02.Н18087
от 18.09.2019 г.



Модификация : «МедИС-В-180/60»

Партия 9226119 Дата изготовления Ноябрь 2019 г.

Гарантийный срок 3 года

Условия эксплуатации и хранения в соответствии с инструкцией производителя

Код ОКПД 2 32.50.50.190

Наименование показателя	Норма	Значение
Технические характеристики	ТУ 9398-032-11764404-2004	Соответствует
Соответствие ГОСТ	класс 4 по ГОСТ ISO 11140-1-2011	Соответствует

Ответственный
за контроль качества



ГРОМАКОВСКАЯ О.С.



www.imq.it

**CERTIFICATO N.
CERTIFICATE N. 9124.CRC4**

SI CERTIFICA CHE IL SISTEMA QUALITA' DI
WE HEREBY CERTIFY THAT THE QUALITY SYSTEM OPERATED BY

CERACARTA SPA

VIA SECONDO CASADEI 14 Z.I. VILLA SELVA - 47122 FORLI' (FC)

UNITA' OPERATIVE / OPERATIVE UNITS

VIA SECONDO CASADEI 14 Z.I. VILLA SELVA - 47122 FORLI' (FC)

E' CONFORME ALLA NORMA / IS IN COMPLIANCE WITH THE STANDARD

EN ISO 13485:2012

PER LE SEGUENTI ATTIVITA' / FOR THE FOLLOWING ACTIVITIES

Produzione e stampa di carte speciali e diagrammate per registrazione ad uso medicale anche conto terzi. Produzione e stampa di etichette ad uso medicale. Sviluppo e produzione di creme, gel sterile e non sterile per applicazioni elettrodiagnostiche e ad ultrasuoni, anche conto terzi. Commercializzazione ed immissione in commercio di accessori per applicazioni elettrodiagnostiche, ad ultrasuoni e per strumenti elettromedicali. Sviluppo e produzione di elettrodi per ECG.

Gestione della produzione ed immissione in commercio di elettrodi per ECG. Immissione in commercio di piastre per elettrobisturi e defibrillatori. Commercializzazione di video stampanti, carte fotografiche per video stampanti, stampanti e relativi materiali di consumo ed accessori per uso medicale

Manufacture and print of special recording chart papers for medical use also on behalf of third parties. Manufacture and print of labels for medical use. Development and manufacture of creams, gels sterile and not sterile for electromedical and ultrasound procedures also on behalf of third parties. Trade and placing on the market of accessories for electromedical and ultrasound diagnostic devices and for electromedical equipment. Development and manufacture of electrodes for ECG. Production management and placing on the market of electrodes for ECG. Placing on the market of electrosurgical plates and defibrillation pads. Trade of videoprinters, photographic papers for videoprinters, printers and related consumable and accessories for medical use

Ulteriori informazioni riguardanti l'applicabilità dei requisiti EN ISO 13485:2012 possono essere ottenute consultando l'organizzazione
Further clarifications regarding the applicability of EN ISO 13485:2012 requirements may be obtained by consulting the organization

IL PRESENTE CERTIFICATO E' SOGGETTO AL RISPETTO DEL
REGOLAMENTO PER LA CERTIFICAZIONE DEI SISTEMI DI GESTIONE
THE USE AND THE VALIDITY OF THE CERTIFICATE SHALL SATISFY THE
REQUIREMENTS OF THE RULES FOR CERTIFICATION OF MANAGEMENT SYSTEMS

DATE:	PRIMA CERTIFICAZIONE	EMISSIONE CORRENTE	SCADENZA
	FIRST CERTIFICATION	CURRENT ISSUE	EXPIRY
	1999-07-20	2017-10-13	2020-10-07

L'Organizzazione dovrà ottenere la certificazione secondo la norma ISO 13485:2016 entro il 2019/02/28;
in caso contrario, il presente certificato cesserà la propria validità in tale data
The Organization shall obtain the certification according to ISO 13485:2016 within 2019/02/28;
otherwise the validity of this certificate will expire

IMQ S.p.A. - VIA QUINTILIANO, 43 - 20138 MILANO ITALY
Management Systems Division - Flavio Ormago

Data di scadenza del precedente ciclo di certificazione: 2017-10-07
Data di conclusione dell'audit di rinnovo: 2017-10-11
Data della decisione di rinnovo: 2017-10-13

CISQ is a member of



*IQNet, the association of the world's first
class certification bodies, is the largest
provider of management System
Certification in the world.*

*IQNet is composed of more than 30
bodies and counts over 150 subsidiaries
all over the globe.*



SGQ N°005A, SGA N°006D, SCR N°005F,
SSI N°003G, FSM N°007I, SGE N°006M,
EMAS N°003P, PRD N°005B, PRS N°008C,
ISP N°003E, LAB N°012I, LAT N°002I

Membro degli Accordi di Mutuo Riconoscimento EA, IAF e ILAC
Signatory of EA, IAF and ILAC Mutual Recognition Agreements

La validità del certificato è subordinata a sorveglianza annuale e riesame completo del Sistema di Gestione con periodicità triennale
The validity of the certificate is submitted to annual audit and a reassessment of the entire Management System within three years

CISQ è la Federazione Italiana di
Organismi di Certificazione dei
sistemi di gestione aziendale.

*CISQ is the Italian Federation
of management system
Certification Bodies.*



www.cisq.com

EC Declaration of Conformity

Manufacturer: Durico C&T, Inc.

33, Oedap 6-Gil, Sangju-Si

Gyeongsangbuk-Do 37240 Republic of Korea

Phone: +82-2-525-8405

Fax: +82-2-525-7461

E-mail: info@durico.co.kr, <http://www.durico.co.kr>

European representative: Durico Imaging BVBA

Villastraat 2 C

1830 Machelen, Belgium

Product: Thermal Paper for Video Printer (Super ULSTAR Brand)

Model: ULSTAR-1100HG, ULSTAR-1100HD, ULSTAR-2100HD,
ULSTAR-1100HD mibi, ULSTAR-1100HD MATT, ULSTAR-1100S,
ULSTAR-1100S mibi, ULSTAR-840HG, ULSTAR-840S

Classification: Class I by the rules of Classification Criteria, Annex IX, MDD 93/42/EEC.

Conformity Assessment Route: Annex VII, MDD 93/42/EEC

We herewith declare that the above mentioned products meet the provisions of the Council Directive 93/42/EEC for medical devices. All supporting documentation is retained under the premises of the manufacturer.

Place and Date of issue: Korea, May 1, 2019

Signature:



J.W. Kim, President

on behalf of Durico C&T, Inc.



CERTIFICATO

Nr. 50 100 5990/A - Rev.004

Si attesta che / This is to certify that

IL SISTEMA QUALITÀ DI
THE QUALITY SYSTEM OF

FAZZINI S.r.l.

SEDE LEGALE E OPERATIVA:
REGISTERED OFFICE AND OPERATIONAL SITE:

**STRADA STATALE PADANA SUPERIORE 317
IT - 20090 VIMODRONE (MI)**

È CONFORME AI REQUISITI DELLA NORMA
HAS BEEN FOUND TO COMPLY WITH THE REQUIREMENTS OF

UNI EN ISO 9001:2015

QUESTO CERTIFICATO È VALIDO PER IL SEGUENTE CAMPO DI APPLICAZIONE
THIS CERTIFICATE IS VALID FOR THE FOLLOWING SCOPE

**VEDI ALLEGATO 1
SEE ANNEX 1**



SGQ N° 049A

Membro degli Accordi di Mutuo Riconoscimento
EA, IAF e ILAC
Signatory of EA, IAF and ILAC Mutual
Recognition Agreements

Per l'Organismo di Certificazione
For the Certification Body
TÜV Italia S.r.l.

Validità /Validity

Dal / From: **2018-10-01**

Al / To: **2021-06-14**

Andrea Coscia
Direttore Divisione Business Assurance

Data emissione / Printing Date

2018-10-01

PRIMA CERTIFICAZIONE / FIRST CERTIFICATION: 2006-07-02

DATA DI SCADENZA DELL'ULTIMO CICLO DI CERTIFICAZIONE 2018-06-14

EXPIRATION DATE OF THE LAST CERTIFICATION CYCLE 2018-06-14

"LA VALIDITÀ DEL PRESENTE CERTIFICATO È SUBORDINATA A SORVEGLIANZA PERIODICA A 12 MESI E AL RIESAME COMPLETO DEL SISTEMA DI GESTIONE AZIENDALE CON PERIODICITÀ TRIENNALE"

"THE VALIDITY OF THE PRESENT CERTIFICATE DEPENDS ON THE ANNUAL SURVEILLANCE EVERY 12 MONTHS AND ON THE COMPLETE REVIEW OF COMPANY'S MANAGEMENT SYSTEM AFTER THREE-YEARS"



pagina 1 di 1 / *page 1 of 1*

THE CERTIFICATE N 50 100 5990/A - Rev.004 IS VALID FOR THE FOLLOWING SCOPE:

Design, manufacturing management, trade and after sales service of active surgical devices (suction pumps), non-active devices for intensive care (manual suction pumps), active devices for breathing therapy (aerosol) and their accessories. Management of design and manufacture, trade and after sales service of non-active devices with a measuring function (blood pressure monitors, scales), non-active devices for anesthesia, emergency and intensive care (balloons, accessories for breathing, anesthesia and aerosol therapy, immobilizers, laryngoscopes endotracheal, first aid kit, stretchers), non-active devices for orthopedic and rehabilitation (aids for the disabled and rehabilitation), non-active devices surgical instruments, active surgical instruments (electrocautery), non-active devices (devices for hospital and ambulatory for the support and the movement of the patient and accessories, stethoscopes), active non implantable device and related accessories and of active devices for monitoring (electrocardiographs, pulse oximeters, monitors, scales). Trade and after sales service of non-active devices for anesthesia, emergency and intensive care (accessories for breathing and anesthesia, dressing and accessories for withdrawals), active devices for positioning and patient support (operating tables), active devices for respiration (breathing accessories), active devices for disinfection and sterilization (sterilizers), devices for electrosurgery, stimulation or inhibition (stimulators), non-active devices with a measuring function (thermometers), active devices for monitoring (thermometers, blood pressure monitors) and office furnitures
(IAF 19, 29)



Membro degli Accordi di Mutuo Riconoscimento
EA, IAF e ILAC
Signatory of EA, IAF and ILAC Mutual
Recognition Agreements

Per l'Organismo di Certificazione
For the Certification Body
TÜV Italia S.r.l.

Validità / *Validity*

Dal / From: **2018-10-01**

Al / To: 2021-06-14

Data emissione / *Printing Date*

2018-10-01

PRIMA CERTIFICAZIONE / FIRST CERTIFICATION: 2006-07-02

DATA DI SCADENZA DELL'ULTIMO CICLO DI CERTIFICAZIONE 2018-06-14

EXPIRATION DATE OF THE LAST CERTIFICATION CYCLE 2018-06-14

“LA VALIDITÀ DEL PRESENTE CERTIFICATO È SUBORDINATA A SORVEGLIANZA PERIODICA A 12 MESI E AL RIESAME COMPLETO DEL SISTEMA DI GESTIONE AZIENDALE CON PERIODICITÀ TRIENNALE”

“THE VALIDITY OF THE PRESENT CERTIFICATE DEPENDS ON THE ANNUAL SURVEILLANCE EVERY 12 MONTHS AND ON THE COMPLETE REVIEW OF COMPANY’S MANAGEMENT SYSTEM AFTER THREE-YEARS.”



CERTIFICATO

Nr. 50 100 5990/B - Rev.006

Si attesta che / This is to certify that

IL SISTEMA QUALITÀ DI
THE QUALITY SYSTEM OF

FAZZINI S.r.l.

SEDE LEGALE E OPERATIVA:
REGISTERED OFFICE AND OPERATIONAL SITE:

**STRADA STATALE PADANA SUPERIORE 317
IT - 20090 VIMODRONE (MI)**

È CONFORME AI REQUISITI DELLA NORMA
HAS BEEN FOUND TO COMPLY WITH THE REQUIREMENTS OF

UNI CEI EN ISO 13485:2016

SISTEMI QUALITÀ – DISPOSITIVI MEDICALI
QUALITY SYSTEMS – MEDICAL DEVICES

QUESTO CERTIFICATO È VALIDO PER IL SEGUENTE CAMPO DI APPLICAZIONE
THIS CERTIFICATE IS VALID FOR THE FOLLOWING SCOPE

**VEDI ALLEGATO 1
SEE ANNEX 1**



SGQ N° 049A

Membro degli Accordi di Mutuo Riconoscimento
EA, IAF e ILAC
Signatory of EA, IAF and ILAC Mutual
Recognition Agreements

Per l'Organismo di Certificazione
For the Certification Body
TÜV Italia S.r.l.

Validità / Validity

Dal / From: **2018-10-01**

Al / To: **2021-06-14**

Andrea Coscia
Direttore Divisione Business Assurance

Data emissione / Printing Date

2018-10-01

PRIMA CERTIFICAZIONE / FIRST CERTIFICATION: 2006-07-02

DATA DI SCADENZA DELL'ULTIMO CICLO DI CERTIFICAZIONE 2018-06-14
EXPIRATION DATE OF THE LAST CERTIFICATION CYCLE 2018-06-14

"LA VALIDITÀ DEL PRESENTE CERTIFICATO È SUBORDINATA A SORVEGLIANZA PERIODICA A 12 MESI E AL RIESAME COMPLETO DEL SISTEMA DI GESTIONE AZIENDALE CON PERIODICITÀ TRIENNALE"

"THE VALIDITY OF THE PRESENT CERTIFICATE DEPENDS ON THE ANNUAL SURVEILLANCE EVERY 12 MONTHS AND ON THE COMPLETE REVIEW OF COMPANY'S MANAGEMENT SYSTEM AFTER THREE-YEARS"



ALLEGATO 1 AL CERTIFICATO NR 50 100 5990/B - Rev.006
ANNEX 1 TO CERTIFICATE NO 50 100 5990/B - Rev.006
 pagina 1 di 1 / page 1 of 1

IL CERTIFICATO NR 50 100 5990/B - Rev.006 È VALIDO PER IL SEGUENTE CAMPO DI APPLICAZIONE::
 THE CERTIFICATE N 50 100 5990/B - Rev.006 IS VALID FOR THE FOLLOWING SCOPE:

Progettazione, gestione della fabbricazione, immissione in commercio e assistenza post vendita di dispositivi attivi chirurgici (aspiratori chirurgici), dispositivi non attivi per terapia intensiva (aspiratori chirurgici manuali), dispositivi attivi per la respirazione (aerosol) e loro accessori. Gestione della progettazione e della fabbricazione, immissione in commercio e assistenza post vendita di dispositivi non attivi con funzione di misura (sfigmomanometri, bilance), dispositivi non attivi per l'anestesia e l'emergenza e la cura intensiva (palloni, accessori per respirazione, anestesia ed aerosolterapia, immobilizzatori, laringoscopi endotracheali, set di pronto soccorso, barelle), dispositivi non attivi ortopedici e per la riabilitazione (ausili per disabili e riabilitazione), strumenti chirurgici non attivi, strumenti chirurgici attivi (elettrobisturi), dispositivi non attivi (dispositivi ospedalieri ed ambulatoriali per il supporto e la movimentazione del paziente e accessori, stetoscopi), dispositivi attivi non impiantabili e relativi accessori. Gestione della progettazione e della fabbricazione, immissione in commercio e assistenza post vendita di dispositivi attivi per monitoraggio (elettrocardiografi, pulsossimetri, monitor, bilance). Commercializzazione e assistenza post vendita di dispositivi non attivi per l'anestesia e l'emergenza e la cura intensiva (accessori per respirazione ed anestesia, accessori per medicazioni e per prelievi), dispositivi attivi per il posizionamento ed il trasporto del paziente (tavoli operatori), dispositivi attivi per la respirazione (accessori per respirazione), dispositivi attivi per la disinfezione e sterilizzazione (sterilizzatrici), dispositivi attivi per monitoraggio (termometri, misuratori di pressione), dispositivi per l'elettrochirurgia, la stimolazione o l'inibizione (stimolatori), dispositivi non attivi con funzione di misura (termometri)

Design, manufacturing management, trade and after sales service of active surgical devices (suction pumps), non-active devices for intensive care (manual suction pumps), active devices for breathing therapy (aerosol) and their accessories. Management of design and manufacture, trade and after sales service of non-active devices with a measuring function (blood pressure monitors, scales), non-active devices for anesthesia, emergency and intensive care (balloons, accessories for breathing, anesthesia and aerosol therapy, immobilizers, laryngoscopes endotracheal, first aid kit, stretchers), non-active devices for orthopedic and rehabilitation (aids for the disabled and rehabilitation), non-active surgical instruments, active surgical instruments (electrocautery), non-active devices (devices for hospitals and ambulatory for the support and movement of the patient and accessories, stethoscopes), non-active implantable devices and related accessories. Management of design and manufacture, marketing and after sales service of active devices for monitoring (electrocardiographs, pulse oximeters, monitors, scales). Trade and after sales service of non-active devices for anesthesia, emergency and intensive care (accessories for breathing and anesthesia, dressings and accessories for withdrawals), active devices for positioning and patient transport (operating tables), active devices for respiration (breathing accessories), active devices for disinfection and sterilization (sterilizers), active devices for monitoring (thermometers, blood pressure monitors), devices for electrosurgery, stimulation or inhibition (stimulators), non active devices with a measuring function (thermometers)



SGQ N° 049A

Membro degli Accordi di Mutuo Riconoscimento
 EA, IAF e ILAC
 Signatory of EA, IAF and ILAC Mutual
 Recognition Agreements

Per l'Organismo di Certificazione
 For the Certification Body
TÜV Italia S.r.l.

Validità /Validity

Dal / From: 2018-10-01

Al / To: 2021-06-14

Andrea Coscia
 Direttore Divisione Business Assurance

Data emissione / Printing Date

2018-10-01

PRIMA CERTIFICAZIONE / FIRST CERTIFICATION: 2006-07-02

DATA DI SCADENZA DELL'ULTIMO CICLO DI CERTIFICAZIONE 2018 -06-14

EXPIRATION DATE OF THE LAST CERTIFICATION CYCLE 2018-06-14

"LA VALIDITÀ DEL PRESENTE CERTIFICATO È SUBORDINATA A SORVEGLIANZA PERIODICA A 12 MESI E AL RIESAME COMPLETO DEL SISTEMA DI
 GESTIONE AZIENDALE CON PERIODICITÀ TRIENNALE"

"THE VALIDITY OF THE PRESENT CERTIFICATE DEPENDS ON THE ANNUAL SURVEILLANCE EVERY 12 MONTHS AND ON THE COMPLETE REVIEW OF
 COMPANY'S MANAGEMENT SYSTEM AFTER THREE-YEARS"

G-CERTI *Certificate*

hereby certifies that

DURICO C&T INC.

33, Oedap 6-gil, Sangju-si, Gyeongsangbuk-do, Korea

has been audited and certified as meeting the requirements & Scope of registration

ISO 9001:2015
Quality Management Systems

**Design, Development, Manufacture and Service of Special Paper
(Thermal Paper, Ink-jet Paper, Photographic Paper, Mat Sheet)**

Certificate No : GK-0233-QC

Valid Period : 22 Jan 2019 ~ 21 Jan 2022

Initial Date : 29 Jan 2010 Issue Date : 17 Jan 2019

Expiry Date : 21 Jan 2022 Code : 07

UIC : MSCB-113-494

Signed for and on behalf of GCERTI
President I.K.Choi



To verify the validity of this certificate please visit : www.gcerti.com
Korea, Seoul, Eunpyeong-gu, Eunpyeong-ro, 88, 15F, Surveillance audits shall be conducted at least once a calendar year, except in recertification years. This is to certify that the Management Systems of this company has been found to conform to the above. If the certified client does not allow surveillance, recertification audits, certificate should be returned to GCERTI. This certificate remains the property of GCERTI and this certificate is recognized by GCERTI.

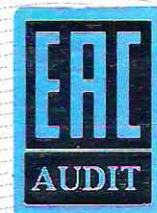




ФЕДЕРАЛЬНОЕ АГЕНТСТВО
ПО ТЕХНИЧЕСКОМУ РЕГУЛИРОВАНИЮ И МЕТРОЛОГИИ
СИСТЕМА ДОБРОВОЛЬНОЙ СЕРТИФИКАЦИИ ГОСТ Р
«EAC AUDIT»

РЕГИСТРАЦИОННЫЙ НОМЕР РОСС RU.32028.04EAC1
ОРГАН ПО СЕРТИФИКАЦИИ ООО «ГОРТЕСТ»
РЕГИСТРАЦИОННЫЙ НОМЕР РОСС RU.32028
ИНН 7717616798 ОГРН 1087746489060

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№003749

СЕРТИФИКАТ СООТВЕТСТВИЯ

Регистрационный номер № 04EAC1.CM.00813

Общество с ограниченной ответственностью «МиниМед»

(наименование лица)

241520, Россия, Брянская область, Брянский район, с. Супонево, ул. Шоссейная, д.17А

(юридический адрес лица)

241520, Россия, Брянская область, Брянский район, с. Супонево, ул. Шоссейная, д.17А

(фактический адрес лица)

ИНН: 3234007127

ОГРН: 1023202138332

НАСТОЯЩИЙ СЕРТИФИКАТ УДОСТОВЕРЯЕТ СООТВЕТСТВИЕ

системы менеджмента качества изделий медицинских Общества с ограниченной ответственностью «МиниМед» требованиям ГОСТ ISO 13485-2017 (ISO 13485:2016) «Изделия медицинские. Системы менеджмента качества. Системные требования для целей регулирования» применительно к Производство лабораторной посуды, медицинских изделий, приборов и принадлежностей, красителей, реагентов и наборов реагентов для in-vitro диагностики

Дата регистрации: 19-03-2019

Срок действия до: 18-03-2022

Руководитель органа
по сертификации:

(подпись)

В. И. Погодин

Председатель
экспертной комиссии:



(подпись)

Е. Д. Курбатова

НАСТОЯЩИЙ СЕРТИФИКАТ ОБЯЗЫВАЕТ ОРГАНИЗАЦИЮ ПОДДЕРЖИВАТЬ СОСТОЯНИЕ ВЫПОЛНЯЕМЫХ РАБОТ В СООТВЕТСТВИИ С
ВЫШЕУКАЗАННЫМИ СТАНДАРТАМИ, ЧТО БУДЕТ НАХОДИТЬСЯ ПОД КОНТРОЛЕМ ОРГАНА ПО СЕРТИФИКАЦИИ СИСТЕМЫ
ДОБРОВОЛЬНОЙ СЕРТИФИКАЦИИ "EAC AUDIT" И ПОДТВЕРЖДАТЬСЯ ПРИ ПРОХОЖДЕНИИ ЕЖЕГОДНОГО ИНСПЕКЦИОННОГО КОНТРОЛЯ

1. Назначение

Коробки стерилизационные круглые КСК-3, КСК-6, КСК-9, КСК-12, КСК-18 (далее – коробки) предназначены для размещения в них перевязочного материала, операционного белья, хирургического инструмента и других предметов медицинского назначения с целью стерилизации их в паровых медицинских стерилизаторах и доставки к месту использования, а также стерильного хранения в течение 3 суток.

Регистрационное удостоверение МЗ РБ № ИМ-7.5619/1004 от 30.03.2010.

Свидетельство о государственной регистрации МЗ Украины № 6184/2007 от 13.03.2012 г.

Регистрационное удостоверение МЗ Казахстана РК-ИМИН-5-№ 001180 от 04.02.2013 г.

Регистрационное удостоверение МЗ Узбекистана № ТТ 13507 от 04.06.2012.

2. Технические характеристики

Наименование параметра	КСК-3	КСК-6	КСК-9	КСК-12	КСК-18
Условный объем, дм ³	3	6	9	12	18
Диаметр, мм	169 _{±5}	234 _{±5}	269 _{±5}	319 _{±5}	374 _{±5}
Масса, кг	0,7±0,1	1,1±0,1	1,3±0,1	1,7±0,15	2,35±0,15

3. Комплект поставки

Коробка, шт. 1
Паспорт, шт. 1
Упаковка, шт. 1

4. Подготовка коробок к работе

До начала применения, коробки должны подвергаться химической очистке дезинфекцией и предстерилизационной обработке в соответствии с действующими инструкциями Минздрава.

5. Свидетельство о приемке, упаковке и продаже

Коробка стерилизационная круглая КСК - 9
соответствует требованиям ТУ 92-00243346-04-93 «Коробки стерилизационные круглые КСК» и признана годной к эксплуатации.



Штамп ОТК
Дата упаковки (изготовления)

Продана

(наименование предприятия торговли)

Дата продажи

(штамп магазина)

(подпись)

Общий срок хранения коробок не ограничен.

6. Гарантии изготовителя

6.1 Гарантийный срок – 24 месяца с даты продажи коробок, но не более 36 месяцев с даты изготовления.

6.2 Гарантийный срок хранения – 24 месяца с даты продажи коробок.

6.3 В течение гарантийного срока изготовитель ремонтирует или заменяет коробки, или их составные части, пришедшие в негодность по вине изготовителя, при предъявлении настоящего паспорта.

Федеральное агентство по техническому регулированию и метрологии

НОПСС

СИСТЕМА ДОБРОВОЛЬНОЙ СЕРТИФИКАЦИИ "НОПСС". РОСС RU.31988.04ЖСН2

Орган по сертификации ООО "Невский Альянс".
ОГРН 1147847286960 ИНН 7842525530

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АКТ

О ПРОХОЖДЕНИИ ЕЖЕГОДНОГО ИНСПЕКЦИОННОГО КОНТРОЛЯ

К СЕРТИФИКАТУ № С1256

ВЫДАН

ООО «МиниМед»

ИНН 3234007127

Настоящий акт удостоверяет, что система менеджмента качества
соответствует требованиям ГОСТ ISO 9001-2015

Сертификат выдан: 24 сентября 2019

Действителен до: 24 сентября 2020

Руководитель органа
по сертификации:



подпись

Платонов Б. А.

