



#### **EC Certificate**

## **Full Quality Assurance System according to** Medical Devices Directive 93/42/EEC Annex-II Section 3

Certificate Number: 1984-MDD-15-320

We hereby declare that an examination of the under mentioned full quality assurance system has been carried out following the requirements of the national legislation to which the undersigned is subjected, transposing annex II (with the exemption of section 4) of the Directive 93/42/EEC on medical devices. We certify that the full quality assurance system conforms with the relevant provisions of the aforementioned directive.

#### Organization:

### OSİMPLANT TIBBİ MALZEMELER VE MEDİKAL TICARET LIMITED SIRKETI

Mustafa Kemal Mah. 2133. Sk. No: 4/2 Çankaya /Ankara/Turkey

Products: Sterile and Non Sterile Spinal Fixation Systems, Sterile and Non Sterile Spinal Cages, Sterile Vertebroplasty and Kyphoplasty Kits

The products defined at the enclosure which is the part of this certificate and contains two pages. The certificate is valid till expiration date, subject to successful completion of periodical surveillance audits. Please contact Kiwa for details.

Report Number:

M.4306.08

Date of first issue: 02 January 2015

Date of last issue:

02 December 2020

Revision Number:

04

**Expiry Date:** 

27 May 2024

02 December 2020, Istanbul, Turkey

Muhteşem Gökhan Yücel Head of Notified Body

Kiwa Belgelendirme Hizmetleri A.Ş. ITOSB 9. Cad. No:15 Tepeören, Tuzla, Istanbul, Turkey Tel.: +90 216 593 25 75 , Fax: +90 216 593 25 74 Web: www.kiwa.com.tr , e-mail: posta@kiwa.com







**Enclosure of the EC Certificate:** 

Page 1/2

Full Quality Assurance System according to
Medical Devices Directive 93/42/EEC Annex-II Section 3
Certificate Number: 1984-MDD-15-320, Revision Number: 04

Concerned medical devices;

**Product:** Sterile and Non Sterile Spinal Fixation Systems

OSI Spinal Fixation System	
ON PLUS Spinal Fixation System	
VESTA Cement Injectable Screw Spinal Fixation System	
JUVE Pediatric Spinal Fixation System	
CERES Minimally Invasive, Percutaneous Screw Spinal Fixation System	
STRATOS Interspinous Fixator Spinal Fixation System	
PORTHOS Posterior Cervical Spinal Fixation System	
ULTIO Laminoplasty Posterior Cervical Spinal Fixation System	
AURA Anterior Cervical Plate System	
ATLAS Odontoid Screw	
ATHENA MIS and Open Modular Pedicle Screw System	

**Product: Sterile and Non Sterile Spinal Cages** 

ARION Expandable Bladed Cervical PEEK Cage

**EOS Bladed Cervical PEEK Cage** 

**BIA Cervical PEEK Cage** 

TITANOPEEK ACF Stand Alone Cervical Cage System

ARIA Expandable PEEK PLIF Cage

FIDES Angular PEEK TLIF Cage

02 December 2020, Istanbul, Turkey

Muhteşem Gökhan Yücel Head of Notified Body

Kiwa Belgelendirme Hizmetleri A.Ş. ITOSB 9. Cad. No:15 Tepeören, Tuzla, Istanbul, Turkey Tel.: +90 216 593 25 75 , Fax: +90 216 593 25 74 Web: www.kiwa.com.tr , e-mail: posta@kiwa.com







**Enclosure of the EC Certificate:** 

Page 2/2

Full Quality Assurance System according to Medical Devices Directive 93/42/EEC Annex-II Section 3 Certificate Number: 1984-MDD-15-320, Revision Number: 04

Concerned medical devices;

Product: Sterile and Non Sterile Spinal Cages

TALOS PEEK TLIF Cage	
ZELOS PEEK-TITANIUM PLIF Cage	
SENTINUS-C Cervical Corpectomy Mesh Cage	
SENTINUS-L Lumbar Corpectomy Mesh Cage	
X-XP Corpectomy Cage	

Product: Sterile Vertebroplasty and Kyphoplasty Kits

ON PLUS Gauge	
ON PLUS Needle Beveled Type	
ON PLUS Bone Filler	
ON PLUS Osteo Introducer	
Kischner Wire	
ON PLUS Spacer (Drill)	
ILOS Spine Kit	

Kiwa Belgelendirme Hizmetleri A.Ş. is Notified Body under Council Directive 93/42/EEC concerning medical devices with identification number: 1984

02 December 2020, Istanbul, Turkey

Muhteşem Gökhan Yücel Head of Notified Body

Kiwa Belgelendirme Hizmetleri A.Ş. İTOSB 9. Cad. No:15 Tepeören, Tuzla, İstanbul, Turkey Tel.: +90 216 593 25 75 , Fax: +90 216 593 25 74

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#### **UYGUNLUK BEYANI**

Declaration of Conformity

"OSİMPLANT" markasına ait olarak tamamen kendi sorumluluğumuz altında üretmiş olduğumuz bu ürünlerin 93/42/ EEC yönetmeliğinin gereksinimlerine göre üretilip kontrol edildiğini beyan ederiz. / We declare that, these products which belong to "OSİMPLANT" brand are completely under our responsibility also producted and checked according to the requirements of 93/42/ EEC.

Üretici firma / Manufacturer: Osimplant Tıbbi Malzemeler Medikal Tic.Ltd.Şti.

**Üretici Adresi** / *Manufacturer Address*: Mustafa Kemal Mah. 2133. Sk. No:4/2 Royale Office Çankaya/ Ankara 06510

Ürün Adı / Product Name:

Steril ve Steril Olmayan Spinal Fiksasyon Sistemleri/Sterile and Non-sterile Spinal Fixation Systems

OSI Spinal Fiksasyon Sistemi/ OSI Spinal Fixation System

ON PLUS Spinal Fiksasyon Sistemi / ON PLUS Spinal Fixation System

VESTA Sement Enjekte Edilebilir Vida Spinal Fiksasyon Sistemi / VESTA Cement Injectable Screw Spinal Fixation System

ARTHOS Titanyum Plazma Kaplı Vida Spinal Fiksasyon Sistemi/ ARTHOS Titanium Plasma Coated Screw Spinal Fixation System

JUVE Pediatrik Spinal Fiksasyon Sistemi/ JUVE Pediatric Spinal Fixation System

CERES Minimal İnvaziv, Perkütan Vida Spinal Fiksasyon Sistemi/ CERES Minimally Invasive, Percutaneous Screw Spinal Fixation System

STRATOS Interspinöz Fiksatör Spinal Fiksasyon Sistemi/ STRATOS Interspinous Fixator Spinal Fixation System

PORTHOS Posterior Servikal Spinal Fiksasyon Sistemi/ PORTHOS Posterior Cervical Spinal Fixation System

ULTIO Laminoplasti Posterior Servikal Spinal Fiksasyon Sistemi/ ULTIO Laminoplasty

Posterior Cervical Spinal Fixation System

AURA Anterior Servikal Plak Sistem/ AURA Anterior Cervical Plate System

ATLAS Odontoid Vida/ATLAS Odontoid Screw

Dok. No: SF-A-5-1 Yayın Tarihi: 17.07.2017 Rev. No: 2 Rev. Tarih: 10.10.2019

Sayfa 1/4



HERAKLES Mini Plak Ve Vida Kraniyel Sistemler/ HERAKLES Mini Plate and Screw Cranial Systems
ATHENA MIS Perkütan Spinal Stabilizasyon Sistem/ ATHENA MIS Percutaneous Spinal

Stabilization System

Ürün Tipleri /Product Types:

Steril ve Steril Olmayan Monoaksiyel Vidalar, Steril ve Steril Olmayan Poliaksiyel Vidalar, Steril ve Steril Olmayan Monoaksiyel Redüksiyon Vidalar, Steril ve Steril Olmavan Poliaksiyel Redüksiyon Vidalar, Steril ve Steril Olmayan Monoaksiyel Sement Enjekte Edilebilir Vidalar, Steril ve Steril Olmayan Poliaksiyel Sement Enjekte Edilebilir Vidalar, Steril ve Steril Olmayan Titanyum Plazma Kaplama Vidalar, Steril ve Steril Olmayan Titanyum Plazma Kaplama Redüksiyon Vidalar, Steril ve Steril Olmayan Sakroiliak Poliaksiyal Vidalar, Steril ve Steril Olmayan Perkütan Vidalar, Steril ve Steril Olmayan Minimal Invaziv Vidalar, Steril ve Steril Olmavan Odontoid Vidalar, Steril ve Steril Olmayan Laminoplasti Vidaları, Steril ve Steril Olmayan Servikal Vidalar, Steril ve Steril Olmayan Lamina Kancalar, Steril ve Steril Olmayan Pedikül Kancalar, Steril ve Steril Olmayan Offset Kancalar, Steril ve Steril Olmayan Rodlar, Steril ve Steril Olmayan Konnektörler, Steril ve Steril Olmavan Kortikal Vidalar, Steril ve Steril Olmavan Plaklar, Steril ve Steril Olmavan Mikro Plaklar, Steril ve Steril Olmayan Mikro Vidalar, Steril ve Steril Olmayan Domino Konnektörler, Steril ve Steril Olmayan Non-Sterile Fiksatörler/ Sterile and İnterspinöz Monoaxial Screws, Sterile and Non-Sterile Polyaxial Screws, Sterile and Non-Sterile Monoaxial Reduction Screws, Sterile and Non-Sterile Polyaxial Reduction Screws, Sterile and Non-Sterile Monoaxial Cement Injectable Screws, Sterile and Non-Sterile Polyaxial Cement Injectable Screws, Sterile and Non-Sterile Titanium Plasma Coated Screws, , Sterile and Non-Sterile Titanium Plasma Coated Reduction Screws, Sterile and Non-Sterile Sacroiliac Polyaxial Screws, Sterile and Non- Sterile Percutaneous Screws, Sterile and Non-Sterile Minimal Invasive Screws, Sterile and Non-Sterile Odontoid Screws, Sterile and Non-Sterile Laminoplasty Screws, Sterile and Non-Sterile Cervical Screws, Sterile and Non-Sterile Lamina Hooks, Sterile and Non-Sterile Pedicle Hooks, Sterile and Nonesterile Offset Hooks, Sterile and Non-Sterile Rods, Sterile and Non-Sterile Connectors, Sterile and Non-Sterile Cortica



Screws, Sterile and Non-Sterile Plates, Sterile and Non-Sterile Micro Plates, Sterile and Non-Sterile Micro Screws, Sterile and Non-Sterile Domino Connectors, Sterile and Non-Sterile Interspinous Fixators

Ürün Sınıfı / Class of Product:

Sinif IIb - Kural 8 Class IIb - Rule 8

GMDN Kodu/ GMDN Adı:

61325/ Kemik-vidası iç omurga sabitleme sistemi, steril olmayan 61324/ Kemik-vidası iç omurga sabitleme sistemi, steril 46653/ Spinal fiksasyon plakası biyobozunur olmayan 61533/Interspinöz spinal fiksasyon İmplant 46642/ Kranyofasiyal fiksasyon plakası, biyobozunur olmayan

GMDN Code/GMDN Name:

61325/Bone-screw internal spinal fixation system nonsterile
61324/Bone-screw internal spinal fixation system sterile
46653/Spinal Fixation Plate non-bioabsorbable
61533/Interspinous Spinal Fixation Implant
46642/Craniofacial fixation plate, non-biodegradable

**Uygulanan Standart ve Yönetmelikler** / Applicable Harmonized Standards and directives: Standartlar listesi referans alındı. / Standard list is referanced

**Uygunluk Değerlendirme Yolu:** MDD (93/42/EEC) EK II- Bölüm 3 Conformity Asssesment Route: MDD (93/42/EEC) ANNEX II- Section3

Onaylanmış Kuruluş Adı / Name of Notified Body: Kiwa Belgelendirme Hizmetleri A.Ş.

Onaylanmış Kuruluş Adresi / Address of Notified Body: (İTOSB) İstanbul Tuzla Organize Sanayi Bölgesi Tepeören Mevkii 34957 Tuzla-İstanbul

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Tel/Phone: +90 216 593 25 75

Türkiye

Faks/Fax: +90 216 593 25 74

Dok. No: SF-A-5-1 Yayın Tarihi: 17.07.2017 Rev. No: 2 Rev. Tarih: 10.10.2019



#### **UYGUNLUK BEYANI**

Declaration of Conformity

"OSİMPLANT" markasına ait olarak tamamen kendi sorumluluğumuz altında üretmiş olduğumuz bu ürünlerin 93/42/ EEC yönetmeliğinin gereksinimlerine göre üretilip kontrol edildiğini beyan ederiz. / We declare that, these products which belong to "OSİMPLANT" brand are completely under our responsibility also producted and checked according to the requirements of 93/42/ EEC

Üretici firma / Manufacturer: Osimplant Tıbbi Malzemeler Medikal Tic.Ltd.Şti.

Üretici Adresi/ Manufacturer Address: Mustafa Kemal Mah. 2133. Sk. 4/2 Royale Office

Çankaya / Ankara-Turkey 06510

Tel / Phone : +90 312 473 82 80 Faks / Fax : +90 312 473 8190
Web: <a href="http://osimplant.com.tr/">http://osimplant.com.tr/</a>
E-Mail: info@osimplant.com.tr

Ürün Adı / Product Name: Steril ve Steril Olmayan Spinal Kafesler/Sterile and Non-

Sterile Spinal Cages

Lomber Kafesler / Lumbar Cages Servikal Kafesler / Cervical Cages

Korpektomi Kafesler / Corpectomy Cages Trabeküler Kafesler/ Trabecular Cages

Ürün Tipleri / Product Types: Steril ve Steril Olmayan Servikal PEEK Kafesler, Steril

ve Steril Olmayan Titanyum Kafesler, Steril ve Steril Olmayan Lomber Interbody Kafesler, Steril ve Steril Olmayan Trabeküler Kafesler, Steril ve Steril Olmayan Genişleyebilir Kafesler, Steril ve Steril Olmayan Bıçaklı Kafesler, Steril ve Steril Olmayan Genişleyebilir Bıçaklı Kafesler, Steril ve Steril Olmayan Servikal Korpektomi Kafesler, Steril ve Steril Olmayan Lomber Korpektomi Kafesler / Sterile and Non-Sterile Cervical PEEK Cages, Sterile and Non-Sterile Titanium Cages, Sterile and Non-Sterile Trabecular Cages, Sterile and Non-Sterile Expandable Cages, Sterile and Non-Sterile Bladed Cages, Sterile and Non-Sterile Expandable Bladed Cages, Sterile and Non-Sterile Cervical Corpectomy Cages, Sterile and Non-Sterile Cervical Corpectomy Cages, Sterile Non-Sterile Cervical Corpectomy Cages, Sterile Non-Sterile Cervical Corpectomy Cages, Sterile Non-Sterile Cervical Corpectomy Cages, Sterile Non-Sterile Cervical Corpectomy Cages, Sterile Non-Sterile Cervical Corpectomy Cages, Sterile Non-Sterile Cervical Corpectomy Cages, Sterile Non-Sterile Cervical Corpectomy Cages, Sterile Non-Sterile Cervical Corpectomy Cages

Sterile Lumbar Corpectomy Cages

NEUROKOS»

Dok. No: TD-SK-4-UB Yayın Tarihi: 17.07.2017 Rev. No: 03 Rev. Tarih: 15.01.2019

Sayfa 1/3



Ürün Sınıfı / Class of Product:

Sinif IIb - Kural 8 Class IIb - Rule 8

GMDN Kodu/ GMDN Adı:

38161/ Spinal Kafes

57805/ Spinal Füzyon Kafesler, Steril Olmayan 60762/ Polimerik Omurga Füzyon Kafesi, Steril Olmayan

60847/ Polimer Omurga Füzyon Kafesi, Steril 38161/ Metalik Spinal Füzyon Kafesler-Steril 57805/ Metalik Spinal Füzyon Kafesler-Steril

Olmayan

GMDN Code/GMDN Name:

38161/Spinal Cage

57805/ Spinal Fusion Cages, Non-Sterile

60762/ Polymeric Spinal Fusion Cage, Non-Sterile

60847/ Polymer Spinal Fusion Cage, Sterile 381611 Metallic Spinal Fusion Cages Sterile 578051 Metallic Spinal Fusion Cages Non-Sterile

Uygulanan Standart ve Yönetmelikler / Applicable Harmonized Standards and directives: Standartlar listesi referans alındı. / Standard list is referanced.

Uygunluk Değerlendirme Yolu:

MDD (93/42/EEC) EK II - Bölüm 3

Conformity Asssesment Route:

MDD (93/42/EEC) ANNEX II- Section3

Onaylanmış Kuruluş Adı / Name of Notified Body:

Kiwa Belgelendirme Hizmetleri A.Ş.

Onaylanmış Kuruluş Adresi /

(İTOSB) İstanbul Tuzla Organize Sanayi Bölgesi Tepeören Mevkii 34957 Tuzla-İstanbul Türkiye

Address of Notified Body:

Tel/Phone: +90 216 593 25 75

Faks/Fax: +90 216 593 25 74

CE Belge No: 1984-MDD-15-320

CE Certificate No:

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#### UYGUNLUK BEYANI Declaration of Conformity

"OSİMPLANT" markasına ait olarak tamamen kendi sorumluluğumuz altında üretmiş olduğumuz bu ürünlerin 93/42/ EEC yönetmeliğinin gereksinimlerine göre üretilip kontrol edildiğini beyan ederiz. / We declare that, these products which belong to "OSİMPLANT" brand are completely under our responsibility also producted and checked according to the requirements of 93/42/ EEC

Üretici firma / Manufacturer: Osimplant Tıbbi Malzemeler Medikal Tic.Ltd.Şti.

Ofis Adres / Office Address: Mustafa Kemal Mah. 2133. Sk. 4/2 Royale Office Çankaya /

Ankara-Turkey 06510

Üretim ve Tasarım Adresi: İkitelli O.S.B. Mutfak Eşyaları San. Sit. M4 Blok No:47

Başakşehir / İSTANBUL

Tel / Phone: +90 312 473 82 80 Faks / Fax: +90 312 473 8190

Web: <a href="http://osimplant.com.tr/">http://osimplant.com.tr/</a>
E-Mail: <a href="mailto:info@osimplant.com.tr">info@osimplant.com.tr</a>

Ürün Adı / Product Name: ENSTRÜMAN SETLER

INSTRUMENTS SETS

Ürün Sınıfı: Sınıf-I Diğer (steril ve ölçme fonksiyonu olmayan)

Class of Product: Class-I Other (without sterile and measuring function)

Uygunluk Değerlendirme Yolu: MDD (93/42/EEC) EK IX

Conformity Assessment Route: MDD (93/42/EEC) ANNEX IX

Yukarıda belirtilen ürünlerin Avrupa Tıbbi Cihaz Direktifi 93/42/EEC, EK-VII warınca üretildiğini beyan ederiz.

We, the manufacturer hereby declare that, the above mentioned products are manufactured in accordance with European Medical Device Directive 93/42/EEC, ANNEX-VII.

Ürünler Tıbbi Cihaz Direktifi 93/42/EEC, EK-I Temel Gereksinimlerine uygundar ve CE işareti ile piyasaya sürülebilir. Cihazların MDD (Tıbbi Cihaz Direktifi) 93/42/EEC PK VII'e uygun olduğunun değerlendirilmesi münhasır sorumluluğumuz altında gerçeklesmiştir.

Products are suitable for Essential Requirements of Medical Device Directive 93/42/EEC, ANNEX-I and could be marketable with CE mark. The evaluation of the conformity of products

Dok. No: TD-UB Yayın Tarihi: 15.01.2019 Rev. No: 00 Rev. Tarih: -

ustale recommend 2131, Se. No. 2 Vesto (2 81, 0.312, 471.62, 20 Fax: 6, 312, 473.81 Ca Umage VO. 1, 632, 031, 10, 33 Texter Sec. 10, 121.21



according to MDD 93/42/EEC (European Medical Device Directive), ANNEX-VII is made under the responsibility of Osimplant exclusively.

Ayrıca 93/42/EEC, EK-VII'e göre teknik bir dokümantasyon sistemi kurduğumuzu beyan ederiz.

Besides, we hereby declare that we have established a technical documentation system according to 93/42/EEC, ANNEX-VII.

Deklarasyon Düzenlenme Tarihi: 15/01/2019

Date of Issuance of Declaration:

Şirket Müdürü / Company Manager

Tel 0.312 +73 92 80 Fax: 6 312 473 87 90

Matte VD.: 632 031 76 39 Titzert Sci. No. 210; 1

Mattels No.: 0032031703900016

Referans Kodu/Reference Code	Ürün Adı/Product Name	GMDN Kodu/ <i>GMDN</i> Code
OSI-1020	OSI Spinal Fixation System Instrument Set	16349
PORTHOS-1010	PORTHOS Posterior Cervical System Instrument Set	16349
SENTINUS-1095	SENTINUS Corpectomy Mesh Cage Instrument Set	16349
AURA-1050	AURA Cervical Plate System Instrument Set	16349
X/XP-1060	X-XP Mesh Corpectomy Cage Instrument Set	16349
BIA-1046	BIA Cervical Cage Instrument Set	16349
TITANOPEEK-1031	TITANOPEEK-C Stand Alone Cervical Cage Instrument Set	16349
ARION-1070	ARION Expandable Cervical Bladed Cage Instrument Set	16349
COMBO-1011	ZELOS&FIDES Combined Instrument Set	16349
ULTIO-1040	ULTIO Laminoplasty System Instrument Set	16349
ONPLUS-1021	ON PLUS Spinal Fixation System Instrument Set	16349
HERAKLES-1015	HERAKLES Micro Plate and Screw Instrument Set	16349
JUVE-1030	JUVE Pediatric Spinal System Instrument Set	16349
ATLAS-1022	ATLAS Odontoid Screw Instrument Set	16349
TALOS-1075	TALOS TLIF Cage Instrument Set	16349
EOS-1035	EOS Cervical Bladed Cage Instrument Set	16349
PAN-1033	PAN Cervical Disc Prosthesis Instrument Set	16349
PALES-1023	PALES Transfacet Pedicle Screw System Instrument Set	16349
ARIA-1080	ARIA Expandable PLIF Cage Instrument Set	16349
ZELOS-1085	ZELOS PLIF Cage Instrument Set	16349
FIDES-1065	FIDES Angular TLIF Cage Instrument Set	16349
STRATOS-1032	STRATOS Interspinous Fixation System Instrument Set	16349











## OSİMPLANT TIBBİ MALZEMELER VE MEDİKAL TİCARET LİMİTED ŞİRKETİ

MUSTAFA KEMAL MAHALLESÍ 2133. SOKAK NO: 4/2 ÇANKAYA – ANKARA – TURKEY

with a scope of

### **DESIGN AND PRODUCTION OF SPINAL IMPLANTS**

Medical devices - Quality management systems - Requirements for regulatory purposes

"Following elements of the standard are excluded" "7.5.3" "7.5.4"

EN ISO 13485:2016

Certificate No

: M 10028

Initial Certification Date

: 11 March 2013

Certification Date

: 20 February 2019

**Expiration Date** 

: 19 February 2022

General Manager

Kiwa Certification Services Inc.

ITOSB 9. Cadde No. 15 Tepeören Tuzla - Istanbul - Turkey

Tel: + 90 216 593 25 75 Faks: + 90 216 593 25 74

Web: www.kiwa.com.tr E-mail: info@kiwa.com.tr

Certificate is valid till expiration date, subject to successful completion of periodical surveillance audits.

Please contact above numbers for detailed information.





U. SOCIE







# OSİMPLANT TIBBİ MALZEMELER VE MEDİKAL TİCARET LİMİTED ŞİRKETİ

MUSTAFA KEMAL MAHALLESÍ 2133. SOKAK NO: 4/2 ÇANKAYA - ANKARA - TURKEY

with a scope of

### DESIGN AND PRODUCTION OF SPINAL IMPLANTS

Has established a quality management system in accordance with international standard.

" Following elements of the standard are excluded "
" None "

ISO 9001:2015

Certificate No

: M 10027

Initial Certification Date

: 11 March 2013

Certification Date

: 01 October 2018

**Expiration Date** 

: 30 September 2021





General Manager

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Certificate is valid till expiration date, subject to successful completion of periodical surveillance audits.

Please contact above numbers for detailed information.