

September 10, 2021

Dentis Co., Ltd. % April Lee Consultant Withus Group Inc 106 Superior Irvine, California 92620

Re: K210410

Trade/Device Name: s-Clean Pre-Milled Abutment Mini

Regulation Number: 21 CFR 872.3630

Regulation Name: Endosseous Dental Implant Abutment

Regulatory Class: Class II Product Code: NHA Dated: June 10, 2021 Received: June 15, 2021

## Dear April Lee:

We have reviewed your Section 510(k) premarket notification of intent to market the device referenced above and have determined the device is substantially equivalent (for the indications for use stated in the enclosure) to legally marketed predicate devices marketed in interstate commerce prior to May 28, 1976, the enactment date of the Medical Device Amendments, or to devices that have been reclassified in accordance with the provisions of the Federal Food, Drug, and Cosmetic Act (Act) that do not require approval of a premarket approval application (PMA). You may, therefore, market the device, subject to the general controls provisions of the Act. Although this letter refers to your product as a device, please be aware that some cleared products may instead be combination products. The 510(k) Premarket Notification Database located at <a href="https://www.accessdata.fda.gov/scripts/cdrh/cfdocs/cfpmn/pmn.cfm">https://www.accessdata.fda.gov/scripts/cdrh/cfdocs/cfpmn/pmn.cfm</a> identifies combination product submissions. The general controls provisions of the Act include requirements for annual registration, listing of devices, good manufacturing practice, labeling, and prohibitions against misbranding and adulteration. Please note: CDRH does not evaluate information related to contract liability warranties. We remind you, however, that device labeling must be truthful and not misleading.

If your device is classified (see above) into either class II (Special Controls) or class III (PMA), it may be subject to additional controls. Existing major regulations affecting your device can be found in the Code of Federal Regulations, Title 21, Parts 800 to 898. In addition, FDA may publish further announcements concerning your device in the <u>Federal Register</u>.

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Please be advised that FDA's issuance of a substantial equivalence determination does not mean that FDA has made a determination that your device complies with other requirements of the Act or any Federal statutes and regulations administered by other Federal agencies. You must comply with all the Act's requirements, including, but not limited to: registration and listing (21 CFR Part 807); labeling (21 CFR Part 801); medical device reporting (reporting of medical device-related adverse events) (21 CFR 803) for devices or postmarketing safety reporting (21 CFR 4, Subpart B) for combination products (see <a href="https://www.fda.gov/combination-products/guidance-regulatory-information/postmarketing-safety-reporting-combination-products">https://www.fda.gov/combination-products/guidance-regulatory-information/postmarketing-safety-reporting-combination-products</a>); good manufacturing practice requirements as set forth in the quality systems (QS) regulation (21 CFR Part 820) for devices or current good manufacturing practices (21 CFR 4, Subpart A) for combination products; and, if applicable, the electronic product radiation control provisions (Sections 531-542 of the Act); 21 CFR 1000-1050.

Also, please note the regulation entitled, "Misbranding by reference to premarket notification" (21 CFR Part 807.97). For questions regarding the reporting of adverse events under the MDR regulation (21 CFR Part 803), please go to <a href="https://www.fda.gov/medical-devices/medical-device-safety/medical-device-reporting-mdr-how-report-medical-device-problems">https://www.fda.gov/medical-device-problems</a>.

For comprehensive regulatory information about medical devices and radiation-emitting products, including information about labeling regulations, please see Device Advice (<a href="https://www.fda.gov/medical-devices/device-advice-comprehensive-regulatory-assistance">https://www.fda.gov/training-and-continuing-education/cdrh-learn</a>) and CDRH Learn (<a href="https://www.fda.gov/training-and-continuing-education/cdrh-learn">https://www.fda.gov/training-and-continuing-education/cdrh-learn</a>). Additionally, you may contact the Division of Industry and Consumer Education (DICE) to ask a question about a specific regulatory topic. See the DICE website (<a href="https://www.fda.gov/medical-devices/device-advice-comprehensive-regulatory-assistance/contact-us-division-industry-and-consumer-education-dice">https://www.fda.gov/medical-devices/device-advice-comprehensive-regulatory-assistance/contact-us-division-industry-and-consumer-education-dice">https://www.fda.gov/medical-devices/device-advice-comprehensive-regulatory-assistance/contact-us-division-industry-and-consumer-education-dice</a>) for more information or contact DICE by email (<a href="DICE@fda.hhs.gov">DICE@fda.hhs.gov</a>) or phone (1-800-638-2041 or 301-796-7100).

Sincerely,

Andrew I. Steen
Assistant Director
DHT1B: Division of Dental and ENT Devices
OHT1: Office of Ophthalmic, Anesthesia,
Respiratory, ENT and Dental Devices
Office of Product Evaluation and Quality
Center for Devices and Radiological Health

Enclosure

# DEPARTMENT OF HEALTH AND HUMAN SERVICES Food and Drug Administration

# Indications for Use

510(k) Number (if known)

Form Approved: OMB No. 0910-0120 Expiration Date: 06/30/2023

Expiration Date: 06/30/2023 See PRA Statement below.

K210410							
Device Name							
s-Clean Pre-Milled Abutment Mini							
Indications for Use (Describe)							
Clean Pre-Milled Abutment Mini is intended for use with dental implants as a support for single or multiple tooth							
rostheses in the maxilla or mandible of a partially or fully edentulous patient.							
All digitally designed abutments for use with s-Clean Pre-Milled Abutment Mini are intended to be manufactured at a Dentis validated milling center.							
Time of the (Select one or both, as applicable)							
Type of Use (Select one or both, as applicable)							
Prescription Use (Part 21 CFR 801 Subpart D) Over-The-Counter Use (21 CFR 801 Subpart C)							
CONTINUE ON A SEPARATE PAGE IS NEEDED							

This section applies only to requirements of the Paperwork Reduction Act of 1995.

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## 510(k) Summary

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#### **Device Information**

Trade Name: s-Clean Pre-Milled Abutment MiniCommon Name: Dental Abutment System

• Classification Name: Endosseous dental implant abutment

• Product Code: NHA

• Panel: Dental

Regulation Number: 872.3630

Device Class: Class IIDate prepared: 09/09/2021

## **Predicate Devices:**

**Primary Predicate** 

K181037, DIO CAD/CAM Abutment by DIO CORPORATION

## Reference Device

K200099, s-Clean SQ-SL Implant System Mini by Dentis Co., Ltd. K111364, Haptite Coating Implant System by Dentis Co., Ltd.

#### **Indications for Use:**

s-Clean Pre-Milled Abutment Mini is intended for use with dental implants as a support for single or multiple tooth prostheses in the maxilla or mandible of a partially or fully edentulous patient.

All digitally designed abutments for use with s-Clean Pre-Milled Abutment Mini are intended to be manufactured at a Dentis validated milling center.

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# **Device Description:**

Patient-specific abutment is made from titanium alloy conforming to ASTM F136 titanium abutment to be used in fabricating patient-specific abutments. The subject abutments are indicated for cemented or "Screw-and Cement-Retained Prosthesis(SCRP)" restorations. Each patient-specific abutment is individually prescribed by the clinician.

The diameters of patient-specific abutment are 5.8, 6.8mm and two connection designs (Hex, Non-hex).

Patient-Specific Abutment is compatible with following Implant Systems:

Proprietary Name	Dentis s-Clean s-Line Mini
Compatible Implants (K number)	K210080
Implant diameter size	5.8/6.8
Implant Interface Connection Type/Size (mm)	Internal Connection type / 2.1
Type of Implant-Abutment Connection	Hex/Non-Hex

s-Clean Pre-Milled Abutment Mini are supplied with s-Clean abutment screw Mini, previously cleared in K210080 and provided non-sterile.

#### **Materials:**

• s-Clean Pre-Milled Abutment Mini and s-Clean Abutment Screw s-Line Mini is fabricated from Ti-6Al-4V ELI (Conforming to ASTM Standard F136).

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# **Summaries of Technology Characteristics**

The subject device is substantially equivalent to the current cleared devices. They are substantially equivalent in intended use, material and connection interfaces to the implants are identical for each individual diameter and connection type. Comparison demonstrating Substantial Equivalence follows:

# <Patient Specific Abutment>

	Subject Device	Pri	mary Predicate Devi	ce	
Applicant	Dentis Co., Ltd.	DIO Corporation			
Trade Name	s-Clean Pre-Milled Abutment Mini	DIC	O CAM/CAM Abutme	ent	
510(k) No.	Not yet assigned	K181037			
Classification Name	Endosseous Dental Implant, Abutment (872.3630)	Endosseous Dental Implant, Abutment (872.3630)			
Product Code	NHA		NHA		
Class	II	II			
Material	Ti-6AL-4V ELI (ASTM F136)	Ti-6AL-4V ELI (ASTM F136)			
Diameter (mm)	CAD/CAM Patient-Specific Abutment : 5.8/6.8	CAD/CAM Patient-Specific Abutment : 3.0/3.3/3.8/4.0/4.5/5.0/5.5/6.0/6.5/7.0			
Sterile	Steam Sterilization by user (Provided Non-Sterile)	Steam Sterilization by user (Provided Non-Sterile)			
Type of Retention	Screw-retained or cement retained	Screw-retained or cement retained			
Abutment Seat	Sits on Taper	Sits on Taper			
Anatomical Site	Oral Cavity	Oral Cavity			
Constructions	Machined	Machined			
		DIO CAD/CAM Abutment is intended for use with dental implants as a support for single or multiple tooth prostheses in the maxilla or mandible of a partially or fully edentulous patient.			
	All digitally designed abutments for use with s-Clean Pre-Milled Abutment Mini are intended to be manufactured at a Dentis validated milling center.	Implant System Compatibility	Implant Diameter (mm)	Platform Diameter (mm)	
		UF(II) Narrow Implant System	3.0/3.3	3.0/3.3	
		UF Sub merged Implant System	3.8/4.0/4.5/5.0/5.5	3.8/4.0/4.5/5.0/5.5	
		Impiant System	/6.0/6.5/7.0	/6.0/6.5/7.0	

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		UF(II) Implant	3.8/4.0/4.5/5.0/5.5	3.8/4.0/4.5/5.0/5.5			
			System	3.6/4.0/4.3/3.0/3.3	3.6/4.0/4.3/3.0/3.3		
		Patient specific abutment is intended for use with the U					
	digitally designed abutn						
	CAD/CAM Abutments are intended to be manufact						
		Co	orporation validated mill	ling center.			
	The subject patient specific abutment is substantially equivalent in designs, dimensions, material, indications, abutment seat, screw seat,						
	anatomical site, connection, and technological characteristics with the identified primary predicate device. The patient specific abutment						
Substantial	is similar in fundamental scientific technology to the predicate.						
Equivalence	The Indications for Use of the subject and primary predicate device are identical other than the compatible implant bodies. This						
-	difference is mitigated by fatigue testing, and identification of reference device for compatible implant bodies. Both the predicate and						
Comparison	subject devices are intended to be milled into patient specific abutments using CAD/CAM technology under the manufacturing control						
	of the sponsor. Any differences in technology characteristics are accompanied by information that demonstrated the device is						
	substantially equivalent as the predicate.						

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#### **Non-Clinical Testing**

The subject device was tested to evaluate its substantial equivalence according to the following standards.

• Fatigue Tests on subject device under the worst-case scenario according to ISO 14801:2016

Below tests were performed for predicate devices and leveraged for the subject device:

- Biocompatibility testing according to ISO 10993-1:2009 abutments referenced in K200099
- End User Sterilization Validation Test Report according to ANSI/AAMI ST79, ISO 17665-1,
   ISO 17665-2, ISO 11737-1, ISO 11737-2, and ISO 11138-1 referenced in K111364
   A worst-case evaluation showed that the previously cleared device was able to be leveraged for the steam sterilization of the subject device

The results of the above tests have met the criteria of the standards and demonstrated the substantial equivalence with the predicate device.

Non-clinical test data was conducted in accordance with the recommendations of FDA Guidance "Class II Special Controls Guidance Document: Root-form Endosseous Dental Implants and Endosseous Dental Implant Abutments", and it consisted of testing finished assembled implant/abutment systems of the worst-case scenario, (smallest diameter with maximum angulation) through fatigue testing.

Clinical testing was not necessary to establish substantial equivalency of the device.

## Conclusion

The s-Clean Pre-Milled Abutment Mini constitutes a substantially equivalent medical device, meeting all the declared requirements of its intended use. This system has the same intended use and fundamental scientific technology as its predicate devices. Therefore, the s-Clean Pre-Milled Abutment Mini and its predicates are substantially equivalent.