

February 23, 2022

Prismatik Dentalcraft, Inc. Jiahe Li Regulatory Affairs Specialist 2144 Michelson Drive Irvine, California 92612

Re: K214102

Trade/Device Name: Glidewell Appliance Resin, Hard/Soft

Regulatory Class: Unclassified Product Code: MQC, KMY Dated: December 27, 2021 Received: December 29, 2021

### Dear Jiahe Li:

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>.

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

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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/medical-devices/device-advice-comprehensive-regulatory-assistance</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 (DICE@fda.hhs.gov) or phone (1-800-638-2041 or 301-796-7100).

Sincerely,

For Michael E. Adjodha, M.ChE.
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

Form Approved: 0MB No. 0910-0120

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

510(k) Number (if known) K214102

**Device Name** 

Glidewell<sup>TM</sup> Appliance Resin, Hard/Soft

Indications for Use (Describe)

Glidewell<sup>TM</sup> Appliance Resin, Hard/Soft is indicated for the fabrication of orthodontic and dental appliances such as bite planes, mouthguards, nightguards, splints and repositioners.

Type of Use (Select one or both, as applicable)

X Prescription Use (Part 21 CFR 801 Subpart D)

Over-The-Counter Use (21 CFR 801 Subpart C)

### CONTINUE ON A SEPARATE PAGE IF NEEDED.

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

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K214102

# 510(k) Summary

### I. SUBMITTER

Prismatik Dentalcraft, Inc. 2144 Michelson Drive, Irvine, CA 92612, USA

Primary Contact Person: Jiahe Li, Regulatory Affairs Specialist

Email: Jiahe.Li@glidewelldental.com

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Phone: (949) 222-3531

Date Prepared: December 27, 2021

#### II. DEVICE

Name of Device: Glidewell<sup>TM</sup> Appliance Resin, Hard/Soft

Classification Product Code: MQC Regulatory Class: Unclassified

Common Name: Mouthguard, Prescription; Classification Name: N/A, Pre-Amendment

Subsequent Product Code: KMY

Regulatory Class: Class I

Common Name: Positioner, Tooth, Preformed Classification Name: Preformed tooth positioner

### III. PREDICATE DEVICE

KeyPrint KeySplint Soft Resin (K183598)

### IV. DEVICE DESCRIPTION

Glidewell<sup>TM</sup> Appliance Resin, Hard/Soft is a light-cured resin for fabrication of orthodontic and dental appliances such as bite planes, mouthguards, nightguards, splints and repositioners. The Glidewell<sup>TM</sup> Appliance Resin, Hard and the Glidewell Appliance Resin, Soft can be used in combination to create a dual layer device consisting of a 3D printed hard layer, made from Glidewell<sup>TM</sup> Appliance Resin, Hard, and a soft layer that is molded with Glidewell Appliance Resin, Soft. The Glidewell<sup>TM</sup> Appliance Resin, Hard can also be used alone to create a 3D printed single layer device. The Glidewell<sup>TM</sup> Appliance Resin, Hard is compatible with DLP printers utilizing a wavelength of 385nm.



# V. INDICATIONS FOR USE

Glidewell™ Appliance Resin, Hard/Soft is indicated for the fabrication of orthodontic and dental appliances such as bite planes, mouthguards, nightguards, splints and repositioners.

# VI. COMPARISON OF TECHNOLOGICAL CHARACTERISTICS

Technological Characteristics		Subject Device Glidewell™ Appliance Resin, Hard/Soft	Predicate Device KeyPrint KeySplint Soft Resin (K183598)	Comparison
Manufacturer		Prismatik Dentalcraft, Inc.	Keystone Industries	N/A
Product	Code	MQC, KMY	MQC, KMY	Same
Prescrip	tion Device	Yes	Yes	Same
Intended Use		Resin for orthodontic and dental appliances	Resin for orthodontic and dental appliances	Same
Indications for Use		Glidewell <sup>TM</sup> Appliance Resin, Hard/Soft is indicated for the fabrication of orthodontic and dental appliances such as bite planes, mouthguards, nightguards, splints and repositioners.	The KeyPrint KeySplint Soft device is indicated for the fabrication of orthodontic and dental appliances such as bite planes, mouthguards, nightguards, splints and repositioners.	The indications for use are the same except for the device trade name.
Design Characteristics	General Design	Light-cured resin using 3D printing technology or combination of 3D printing and injection molding to convert the liquid resin to solid orthodontic and dental appliances.	Light-cured resin using 3D printing technology to convert the liquid resin to solid orthodontic and dental appliances.	The subject device uses 3D printing technology to convert the liquid resin to solid orthodontic and dental appliances, same as the predicate device.  The only difference is that the subject device has the additional option of fabricating a dual-layered device through 3D printing of the hard outer layer with the hard resin and injection molding of the soft inner layer with the soft resin.



Technological Characteristics	Subject Device Glidewell™ Appliance Resin, Hard/Soft	Predicate Device KeyPrint KeySplint Soft Resin (K183598)	Comparison
Material composition	Light cured methacrylate based-resin.	Light cured methacrylate based-resin.	Substantially equivalent
Additive Manufacturing System	Glidewell <sup>TM</sup> Appliance Resin, Hard/Soft is intended to be used in conjunction with an additive Computer- Aided Manufacturing (CAM) and curing system.	The KeyPrint KeySplint Soft device is intended to be used in conjunction with an additive Computer- Aided Manufacturing (CAM) and curing system.	Same
Bench Testing (physical property)	Physical properties according to  ISO 20795-2: Flexural Strength, Flexural Modulus, Water Sorption/Solubilit y  ASTM D638: Tensile Strength, Tensile Modulus, Elongation	Physical properties according to  • ASTM D790: Flexural Properties • ISO 20795-2: Flexural Strength, Flexural Modulus, Water Sorption/Solubility • ASTM D638: Tensile Strength, Tensile Modulus, Elongation	The predicate device was tested for flexural properties according to ASTM D790 in addition to ISO 20795-2.
Biocompatibility	Biocompatible per testing results according to ISO 10993-1	Biocompatible per testing results according to ISO 10993-1	Same
Sterility	Non-Sterile	Non-Sterile	Same

# DETERMINATION OF SUBSTANTIAL EQUIVALENCE

The subject device, Glidewell<sup>TM</sup> Appliance Resin, Hard/Soft, is substantially equivalent to the primary predicate device, KeyPrint KeySplint Soft Resin (K183598) in intended use, indications for use and technological characteristics, including technical specifications/features, material and principles of operation.



The subject device, Glidewell<sup>TM</sup> Appliance Resin, Hard/Soft, has the same intended use as the predicate device, KeyPrint KeySplint Soft Resin (K183598) as material for fabricating prescription mouthguards and preformed tooth positioners. The subject device, Glidewell<sup>TM</sup> Appliance Resin, Hard/Soft, has the same Indications for Use Statement (IFUS) as the predicate device, KeyPrint KeySplint Soft Resin (K183598), except for the device trade name. Both the subject device, Glidewell<sup>TM</sup> Appliance Resin, Hard/Soft, and the predicate device, KeyPrint KeySplint Soft Resin (K183598) are dental resins indicated for the fabrication of orthodontic and dental appliances such as bite planes, mouthguards, nightguards, splints and repositioners.

The subject device, Glidewell<sup>TM</sup> Appliance Resin, Hard/Soft, is substantially equivalent to the predicate device, KeyPrint KeySplint Soft Resin (K183598) in technical specifications/features. The product specifications of the subject device, Glidewell<sup>TM</sup> Appliance Resin, Hard/Soft, is based on benchmarking comparable devices with similar indications for use on the market, including the predicate device, KeyPrint KeySplint Soft Resin (K183598). The same property testing according to ISO 20795-2 and ASTM D638 that applied to the predicate device, KeyPrint KeySplint Soft Resin (K183598), were performed on the subject device, Glidewell<sup>TM</sup> Appliance Resin, Hard/Soft, including flexural strength, flexural modulus, water sorption, water solubility, tensile strength, tensile modulus and elongation. The physical property of the subject device, Glidewell<sup>TM</sup> Appliance Resin, Hard/Soft, passed the threshold of performance criteria in ISO 20795-2 when applicable to the design, and met the same performance criteria used by the predicate device, KeyPrint KeySplint Soft Resin (K183598).

The subject device, Glidewell<sup>TM</sup> Appliance Resin, Hard/Soft and the predicate device, KeyPrint KeySplint Soft Resin (K183598) are similar in material composition. Both the subject device, Glidewell<sup>TM</sup> Appliance Resin, Hard/Soft, and the predicate device, KeyPrint KeySplint Soft Resin (K183598) are light cured methacrylate-based resin. Despite the actual difference that might exist between the formulations, the difference does not affect the safety and effectiveness for the indended use, as verified by the safety and performance testing.

The subject device, Glidewell™ Appliance Resin, Hard/Soft, is substantially equivalent to the predicate device, KeyPrint KeySplint Soft Resin (K183598) in terms of principle of operation. Both the subject device, Glidewell™ Appliance Resin, Hard/Soft, and the predicate device, KeyPrint KeySplint Soft Resin (K183598) are light-cured resin in liquid form that through light-mediated conversion can be fabricated into a solid finished device. The finished device has the desired physical properties suitable for orthodontic and dental appliances such as bite planes, mouthguards, nightguards, splints and repositioners.

### VII. PERFORMANCE DATA

Non-clinical data submitted to demonstrate substantial equivalence include:



- Flexural strength and flexural modulus, according to ISO 20795-2
- Water sorption and water solubility, according to ISO 20795-2
- Tensile Strength, Tensile Modulus and Elongation, according to ASTM D638
- Printing Accuracy and Printing Orientation Validation
- Packaging validation
- Biocompatibility

No clinical data is included in this submission.

# Flexural Strength and Flexural Modulus

The subject device, Glidewell<sup>TM</sup> Appliance Resin, Hard/Soft, was tested for flexural strength and flexural modulus in accordance with the test methods outlined in ISO 20795-2. All the testing results met the acceptance criteria. The results of the testing were used to address questions related to substantial equivalence based on differences in technical specifications between the subject device, Glidewell<sup>TM</sup> Appliance Resin, Hard/Soft, and the predicate device, KeyPrint KeySplint Soft Resin (K183598).

## Water Sorption and Water Solubility

The subject device, Glidewell<sup>TM</sup> Appliance Resin, Hard/Soft, was tested for water sorption and water solubility in accordance with ISO 20795-2. All the testing results met the acceptance criteria. The results of the testing were used to address questions related to substantial equivalence based on differences in technical specifications between the subject device, Glidewell<sup>TM</sup> Appliance Resin, Hard/Soft, and the predicate device, KeyPrint KeySplint Soft Resin (K183598).

## Tensile Strength, Tensile Modulus and Elongation

The subject device, Glidewell<sup>TM</sup> Appliance Resin, Hard/Soft, was tested for tensile strength, tensile modulus and elongation in accordance with ASTM D638. All the testing results met the acceptance criteria. The results of the testing were used to address questions related to substantial equivalence based on differences in technical specifications between the subject device, Glidewell<sup>TM</sup> Appliance Resin, Hard/Soft, and the predicate device, KeyPrint KeySplint Soft Resin (K183598).

#### Printing Accuracy and Printing Orientation Validation

Printing accuracy test was performed to validate that the physical output of the additive manufacturing system and procedure for Glidewell<sup>TM</sup> Appliance Resin, Hard are able to meet design input dimensions within the pre-specified tolerance. Printing orientation test was performed to validate that the hard resin printed at different print direction within the build space relative to the device orientation and at different build plate locations are able to meet the same performance criteria. The results met the pre-specified acceptance criteria and demonstrated that the subject device, Glidewell<sup>TM</sup> Appliance Resin, Hard/Soft, can be reliably fabricated at different print directions within the build space and at different build plate locations using the additive manufacturing system and procedure.



# Packaging Validation

Packaging validation was performed for the subject device, Glidewell<sup>TM</sup> Appliance Resin, Hard/Soft. Per ASTM D4169-14, the shipping unit was tested for manual handling, compressive loads, repetitive shocks from vibration, vertical vibration environments, concentrated impacts and secondary manual handling drops. It was determined that Glidewell<sup>TM</sup> Appliance Resin, Hard/Soft with the respective packaging, is suitable for use. The results of the testing were used to address questions related to substantial equivalence based on differences in product packaging between the subject device, Glidewell<sup>TM</sup> Appliance Resin, Hard/Soft, and the predicate device, KeyPrint KeySplint Soft Resin (K183598).

# **Biocompatibility**

The subject device, Glidewell™ Appliance Resin, Hard/Soft, was tested in accordance with ISO 10993-1. Based on the biocompatibility testing results, it was determined that there is no biocompatibility concern for the subject device. The results of the testing were used to address questions related to substantial equivalence based on differences in chemical composition between the subject device, Glidewell™ Appliance Resin, Hard/Soft, and the predicate device, KeyPrint KeySplint Soft Resin (K183598).

## VIII. CONCLUSION

Based on technological characteristics and non-clinical test data included in this submission, the subject device, Glidewell<sup>TM</sup> Appliance Resin, Hard/Soft, has been shown to be substantially equivalent to the predicate device, KeyPrint KeySplint Soft Resin (K183598).