

1 Introduction

E-Guard, available as “*E-Guard*” and “*E-Guard Blue*”, is a light-curing material for the additive manufacturing of individual dental bite splints. *E-Guard* has been optimized for use with Desktop Health’s *Einstein* and *Einstein Pro XL* machines and EnvisionTECs *Perfactory*® DDP (*Digital Dental Printer*) series, *Perfactory*® *Vida* and *Perfactory*® *Vida cDLM*, *Perfactory*® *MicroPlusXL*, *Perfactory* *EnvisionOne cDLM*, *Perfactory*® *P4K* series and *Perfactory*® *D4K* machines and may only be used with these machines and the corresponding software systems. *E-Guard* is a medical device classified as class 2 according to 21 CFR part 872. Bite splints from *E-Guard* may only be manufactured by dental technicians and must be inspected by authorized practitioners such as dentists or orthodontists before they are released to the patients.

The following Instruction for Use includes safety and environmental information and the instructions for the manufacturing and post-processing procedures of the product, which must be strictly adhered to.

The minimum approved wall thickness is 3 mm and the maximum approved wall thickness is 10 mm. After several printing processes the product may show slight color changes. However, this does not reduce the quality of the application.

2 Indication for Use / Intended Use

EnvisionTEC’s *E-Guard* is a light-cured resin. It is a polymer used to create removable structures for therapeutic restorations, i.e. bite guards/splints and occlusal night guard/splints using the Additive Manufacturing process. The resin in combination with a scanner, printer, and curing unit make up the system.

3 Contraindication

E-Guard may only be used for the production of dental bite splints. Any deviation from the Instruction for Use can negatively affect the chemical and physical properties. Consequently, the biocompatibility of the bite splint cannot be guaranteed.

E-Guard may not be used for the production of clear aligners, bleaching-splints, sport splints or protrusion- / snoring-splints.

For patient and users:

Patients, users or third parties who come in contact with products from *E-Guard* must be informed about any side effects before use. *E-Guard* products may not be used if there are known allergies to any of the ingredients, otherwise possible side effects such as shortness of breath, gastrointestinal complaints, dizziness, anaphylactic reactions or shocks, itching and tearing (watery) eyes, headaches or reactions of the skin or mucous membrane such as irritation, rash, swelling, inflammation, redness, wheals or blisters or other allergic reactions may occur.

4 Composition

Acrylate/Methylacrylates, methacrylated Oligomers and Monomers, photo initiators, colorants/dyes and absorbers

5 Warning

Do not substitute any of the components of the device system, i.e., device resin materials, scanners, printers, post-curing units, CAD/CAM software, templates and tools. Use only those specifically identified in this labelling. Contact the manufacturer for compatible components.

6 Hazard and Precaution statements

Warning:

Causes skin irritation.

May cause an allergic reaction.

Causes serious eye irritation.

Suspected of damaging fertility.

Wear protective gloves, protective clothing, eyes protection, face protection.

First Aid:

If on skin wash with plenty of water.

If in eyes rinse cautiously with water for several minutes. Remove contact lenses, if present and easy to do. Continue rinsing.

If swallowed, immediately call poison center.

7 Storage conditions and und expiry date

The resin needs to be stored in the original packaging between 41 – 86 °F. While removing the resin it must be protected from exposure to light, as spontaneous polymerization is possible. The bottle must be tightly closed after every usage and material removal. If the material is not used, it must be filled back into the bottle. An expiration date is given on the label of every product bottle. The use of expired material is not permitted. The bite splints need to be protected from exposure to light before the final use and during the not usage and storage.

8 Notes on disposal

Dispose of contents and container in accordance with local regulation.

The manufacture bite splints, which are used on patients must be disposed properly due to the risk of infection (contaminated by substances of human origin).

9 Instruction disinfection and sterilization

If necessary, the bite splints made of *E-Guard* can be disinfected before use with the following disinfectants: Cidex OPA or a 70 % Ethanol-solution. The disinfecting solutions must be used according to the manufacturer instructions.

Bite splints from *E-Guard* cannot be sterilized.


10 Use of software systems and products from other manufacturer

The use of certified software systems for generating the STL-data depends on user's assessments.

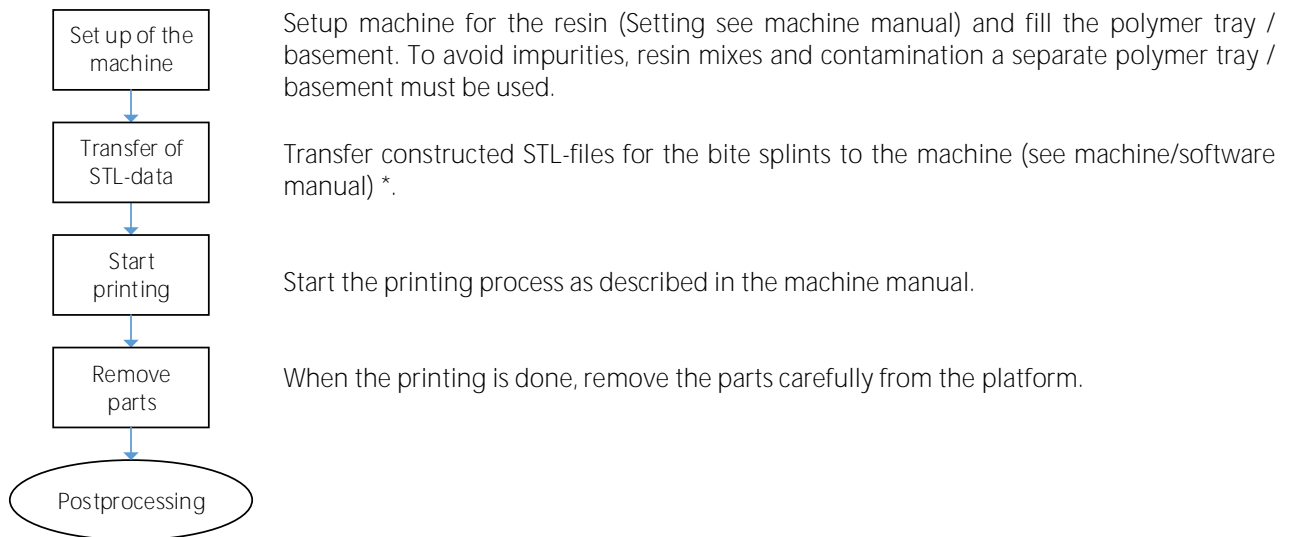
11 Delivery unit, symbol explanation and manufacturer information

Delivery unit: E-Guard is available in containers of 1 kg

Symbol explanation:

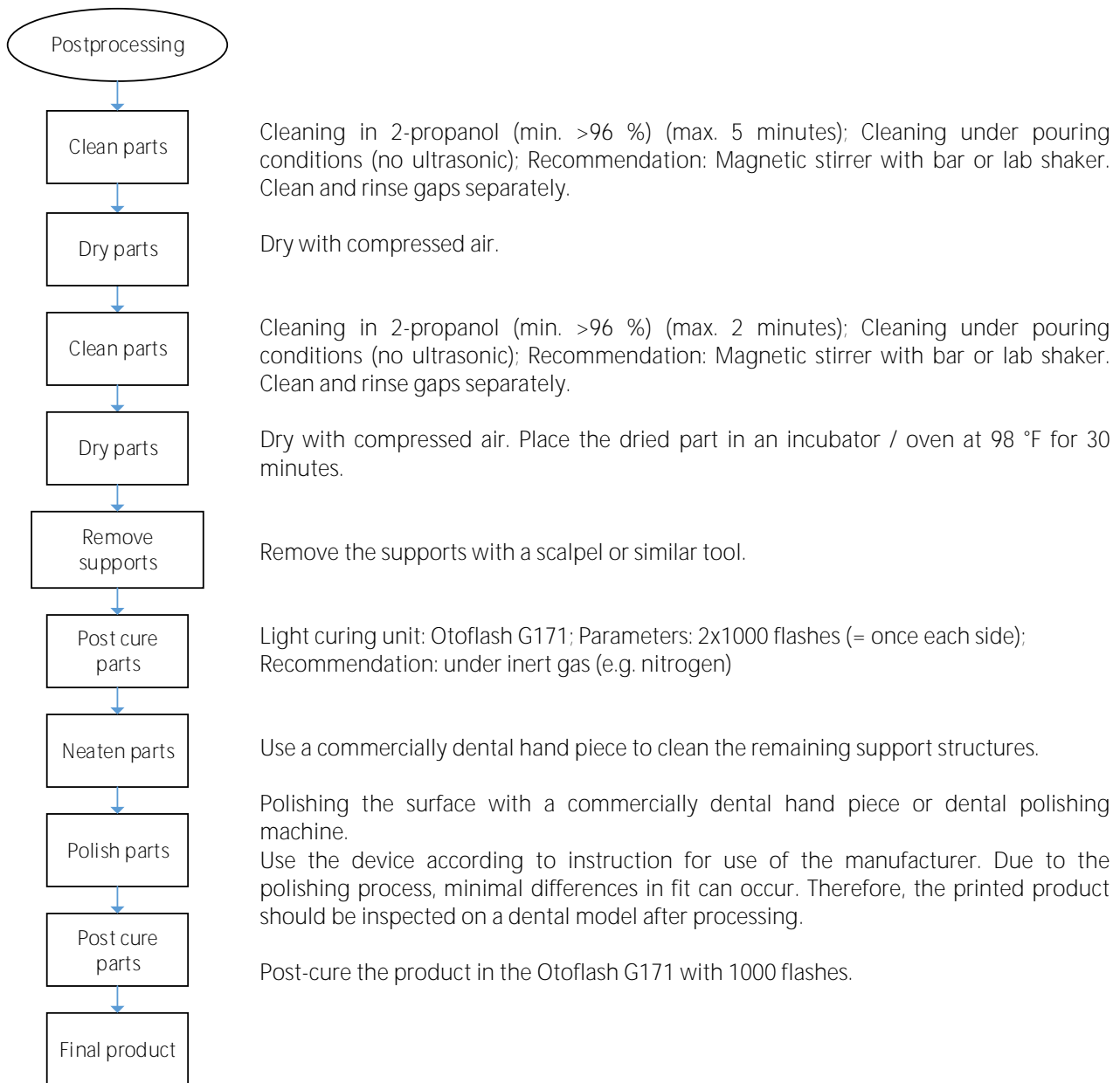
 LOT	Batch number		Protect from sunlight
	Expiration date		Follow Instruction for Use
	Manufacturer		Temperature limit
 REF	Purchase order number		Production date
 Rx Only	Prescription device labeling statement	 UDI	Unique device identification

12 Manufacturing process



*The scanning and construction of patient's STL data is the responsibility of the customer. We recommend that only trained dental personnel perform the scanning and design. Further, a certified software such as, e.g. 3Shape is recommended. To generate the support structures, we recommend the Perfactory® RP Software (version 3.1540.1602 or later), EnvisionOne RP (version 1.0.1165 or later) or the Cambridge Software from 3Shape (version 2015 2650 or later).

13 Postprocessing



The product can now be used on the patient.

Maintain and calibrate equipment according to manufacturer instructions. Using an alternative light source can affect the properties of the final product. If a manual removal of the material after the fitting on a patient is necessary, then the products needs to be post-cured again with 1000 flashes in Otofash G171.

14 Patient cleaning instruction for bite splint

The bite splint can be cleaned by the patient with clear water, a toothbrush and toothpaste. Abrasive or whitening agents in toothpastes can damage the surface of the splint. After cleaning with clear water, the splint should be dried and not soaked in liquid.

15 Reporting undesirable effects

In the event of adverse effects, reactions or similar occurrences arising from the use of this products, including those not listed in this Instruction for Use, these must be reported immediately by opening a support ticket via our website <https://envisiontec.com/> or by contacting your local distributor.

16 Manufacturer

EnvisionTEC GmbH

Brüsseler Str. 51

45968 Gladbeck

Germany

Phone: +49-(0)2043-98750

E-Mail: info@envisiontec.com

Website: www.envisiontec.com

Revision 07, 16.08.2022

Rx Only