

## **EU Declaration of Conformity**

We: Water Pik, Inc.

1730 East Prospect Road Fort Collins, CO 80553-0001

Fort Collins, CO 60553-0001

**USA** 

**EC Representative** 

Sofibel SAS

92686 Levallois-Perret Cedex-France

declare under our sole responsibility, using the conformity assessment route through Regulation (EU) 2017/745 of the European Parliament and of the Council of 5 April 2017 on Medical Devices Annex VIII, Rule 5, that the

### Original Tofflemire<sup>®</sup> Matrix Bands

- No.1 Adult Universal (34577-0, 34578-9, 062001-000, 062002-000, 061032-000)
- o No.2 Adult MOD Wide (34579-7, 34580-0)
- o No.3 Adult MOD Narrow (34582-6)
- o No.13 Pedo Universal (34584-2, 34585-0, 062010-000)

### Original Tofflemire® "Dead Soft" Matrix Bands

- o No.1 Adult Universal (34603-6)
- o No.2 Adult MOD Wide (34604-4)
- o No.13 Pedo Universal (34605-2)

#### Getz<sup>®</sup> Contour Bands

- o Regular (061081-000)
- o Ultra-thin (061080-000)

### • Original Tofflemire® II Retainers

- o Universal (20016453)
- o Contra-Angle (20016455)

# Wizard Wedges® Wood Matrix Wedges

- o Package of 100 Assortment of Sizes (061205-000)
- o Box of 500 (061211-000 through 061215-000)

## • Wizard Wedges® Anatomical Wood Matrix Wedges

- o Package of 100 Assortment of Sizes (061315-000)
- o Box of 500 (061306-000 through 061310-000)

classified as **Class I** meet the general safety and performance requirements and are in conformity with the Regulation listed below using the relevant sections of the Regulation and other normative documents:

### Regulation:

Regulation (EU) 2017/745 of the European Parliament and of the Council of 5 April 2017 on Medical Devices



### **Conformity Route:**

In accordance with Section 7 of Article 52 of the medical device regulation, the chosen conformity assessment procedure is declaration of conformity by issuing this EU declaration of conformity referred to in Article 19 after drawing up the technical documentation set out in Annexes II and III.

#### Standards:

- BS EN ISO 13485:2016 Medical Devices Quality Management Systems
  Requirements for Regulatory Purposes
- BS EN ISO 14971: 2012 Medical Devices Application of Risk Management to Medical Devices
- ISO 10993-1:2018 Biological Evaluation of Medical Devices
  - Part 1: Evaluation and Testing Within a Risk Management Process
- BS EN ISO 15223-1:2016 Medical Devices Symbols to be Used with Medical Device Labels, Labelling and Information to be Supplied
  - Part 1: General Requirements
- BS EN 1041:2008 + A1:2013 Information Supplied by the Manufacturer of Medical Devices
- BS EN 1641:2009 Dentistry Medical Devices for Dentistry Materials

Date CE mark was affixed: June 1998

Date

. 10/31/2019

Signature Name

: Robert Vander Vliet

Title

Manager Regulatory and Quality Systems

Supersedes: 7-Dec-2016

End Date: NA