

DECLARATION OF CONFORMITY

Ultradent Products, Inc. has evaluated the following product by using the Conformity Assessment Procedure of Annex II of the Medical Device Directive 93/42/EEC, as amended by 2007/47/EEC:

Ultra-Etch 35%

and confirms in sole responsibility that the product is compliant with the Essential Requirements of Annex I of the Medical Device Directive 93/42/EEC. Technical documentation is located in the Regulatory Affairs Department.

This product system is classified as Class IIa medical device according to the Medical Device Directive 93/42/EEC, Annex IX, Section III Classification 2.2, Rule 6

UMDNS Code: 17737, Dental Etching Liquids GMDN Code: 36153, Dental etching composite **Notified Body: EC Representative:** Ultradent Products GmbH TÜV Nord Cert GmbH Unternehmensgruppe TüV Nord Am Westhover Berg 30 Langemarckstraße 20 51149 Cologne 45141 Essen, Germany Germany ID No. 0044 02 Dec 2020 Karen Kakunes RN, BSN Regulatory Affairs Management State of Utah County of Salt Lake Subscribed and sworn to before me on this 2 day of <u>December</u> 20 20 Army Henderin- Nulm

This document is in force as long as the following EC certificates are valid:

EC Certificate 44 232 090234 valid through 26 May 2024