

## DECLARATION OF CONFORMITY

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

### VALO Cordless and VALO Grand

and confirms 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 I medical device according to the Medical Device Directive 93/42/EEC, Annex IX, Section III Classification, 3.2 Rule 12

**UMDNS Code:** 16386, Lights, Dental Activator

**GMDN Code:** 35775, Light, Dental, polymerization activator

**EC Representative:**

Ultradent Products GmbH  
Am Westhover Berg 30  
51149 Cologne  
Germany



Adam Black  
Regulatory Affairs Manager

23 Nov 2020  
Date

State of Utah  
County of Salt Lake

Subscribed and sworn to before me on this 23 day of November 2020

By Adam Black

Amy Henderson-Nielson  
Notary Public



This document is in force as long as the following ISO 13485 certificates are valid:

ISO 13485:2016 Certificate 19-1612-Q valid through 02-Aug-2023  
acc. to ISO 13485:2016 Certificate 19-1613-M valid through 02-Aug-2023