

Certificate of Registration

Intertek

This is to certify that the quality management system of

Premier Dental Products Company Also DBA Premier Medical

Main Site: 1710 Romano Drive, Plymouth Meeting, Pennsylvania 19462, United States

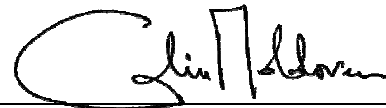
has been assessed and registered by Intertek, a CMDCAS recognized registrar, as conforming to the requirements of

ISO 13485:2003

The quality management system is applicable to

The design and development and manufacture of dental and medical products, including perio-hygiene instruments, restorative materials, dental cements, dental varnishes, endo instruments, implantable RC post, polishing-finishing material, sterile and nonsterile burs, Tracheostomy / Laryngectomy Tubes and Hemostatic Agent/Astringent.

Certificate Number: 9371-12
Initial Certification Date: 12 May 2006
Certificate Effective Date: 12 May 2015
Certificate Expiry Date: 11 May 2018



Calin Moldovean, President

Intertek Testing Services NA, Ltd. – 1829, 32nd avenue, Lachine, QC, H8T 3J1, Canada



In the issuance of this certificate, Intertek assumes no liability to any party other than to the Client, and then only in accordance with the agreed upon Certification Agreement. This certificate's validity is subject to the organization maintaining their system in accordance with Intertek's requirements for systems certification. Validity may be confirmed via email at certificate.validation@intertek.com or by scanning the code to the right with a smartphone.

The certificate remains the property of Intertek, to whom it must be returned upon request.

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