

Medical Device Production Quality Assurance System Certificate
GB23/00000313

The management system of

Davis, Schottlander & Davis Ltd

Fifth Avenue Letchworth Hertfordshire SG6 2WD United Kingdom
has been assessed and certified as meeting the requirements of

Part II of The Medical Devices Regulations 2002, Annex V [as modified by Part 2 of Schedule 2A to The Medical Devices Regulations 2002]

For the following products

Temporary Crown & Bridge Materials:
Quicktemp 2, Quicktemp Cosmetic,
Quicktemp Cosmetic Glaze & Bond.
Crown & Bridge Materials: Ceramic Powder -
Matchmaker Powders MC, enliven MC
Bonding alloys non-precious Matchmate
NP10 and NP20

Restoratives & Cements:

Light cured composite filling material -
Ultra Fine Glass Composite, Quicktemp Cosmetic
Flowable Composite. Amalgam Alloys –
Starburst 45 Capsulated Alloy.

Prosthetics:

Acrylic Resin Teeth - Enigma, Enigmalive,
Natura, & 4natur.

Prosthetics Materials:

Cobalt chromium alloy - Croform Excel S1.
Pegasus Pourable cold cure denture Base material.
Pegasus Plus repair acrylic.
Pegasus plus denture base
Acrylic Denture Base - Enigma High Base.

Where the above scope includes class IIb or class III medical device(s), a valid Type Examination Certificate according to Annex III [as modified by Part 2 of Schedule 2A to The MDR 2002] is a mandatory requirement for each device in addition to this certificate to place that device on the market

Certification is based on reports numbered GB/PC/07450

Previous certificate number: N/A

Change in between this certificate and previous one: Addition of *Acrylic Denture Base - Enigma High Base* to certificate scope

This certificate is valid from 11 October 2023 until 17 August 2028 and remains valid subject to satisfactory surveillance audits.

Issue 2. Certified since 17 August 2023



Authorised by
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