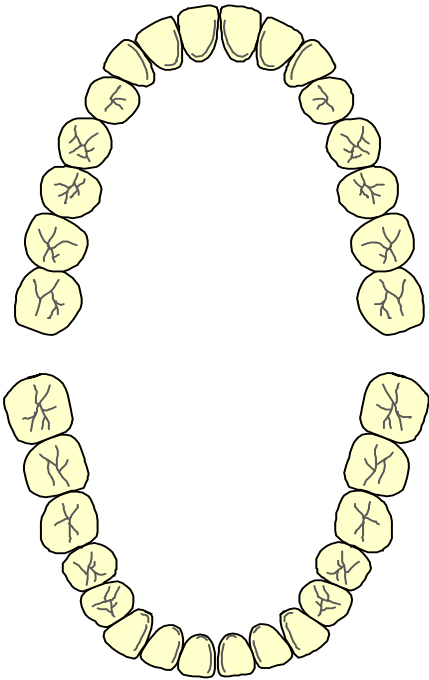


AW Precision Ceramics Unit 3, The Street, Ewhurst, Surrey, GU6 7QD 01483 277070 REGISTERED WITH THE UK COMPETENT AUTHORITY	CUSTOM-MADE DENTAL APPLIANCE PRESCRIPTION <i>Please complete the appropriate sections of this prescription and return to the address opposite.</i>
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Patients' Name/ID:	Name of prescriber:	Clinic name and address:
Date Sent:	Date required:	

INSTRUCTIONS AND AMENDMENTS RECORD



FIELDS BELOW TO BE COMPLETED BY LABORATORY PERSONNEL ONLY	
Approved for manufacture by: Sign:	Approved for release by: Sign:
Details of materials etc supplied by prescriber: Initials:	Details of any model approval by prescriber: Initials:

Your attention is drawn to the following statement: This is a custom-made medical device that has been manufactured to satisfy the design characteristics and properties specified by the prescriber for the above-named patient. This medical device is intended for exclusive use by this patient and conforms to the **general safety and performance requirements** specified in Annex I of the **Medical Devices Regulations**.

This statement does not apply to medical devices that have been repaired and/or refurbished for an individual patient's use.

Storing, handling and instructions for use: It is recommended that before use, this medical device is stored in a clean and safe environment that prevents it from coming into contact with materials, equipment, acids or bleaches that could cause physical or chemical damage to the medical device. The medical device should not be subjected to extremes of temperature during storage. Where applicable, you should take care not to damage the medical device when removing it from its model.

ORIGIN OF MANUFACTURE DECLARATION

This complete appliance has been wholly manufactured within the EU.

PRESCRIBER FEEDBACK:

To enable our dental laboratory to comply with the Medical Devices Regulations for Post Market Surveillance, please inform us of any feedback or issues regarding the enclosed device(s) as soon as possible.