

NHS Prosthetics

CosTech Dental Laboratory

Unit 17-18, Lion Business Park, Dering Way, Gravesend, Kent, DA12 2DN

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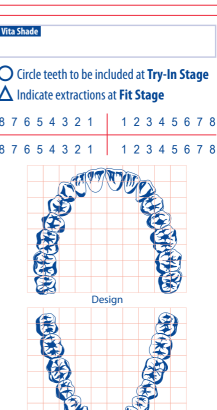
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Barcode for internal use only	

NEW Track this case onli	portal.costech.co.uk				
Patient & Job Details					
Job ID: Patient's Name:					
Chrome Upper Lower	Repair Upper Lower				
Reline Upper Lower	Additions Upper Lower				

3 Due Dates				
Special Tray: U				
Bite Block:				
Try-In:				
Re-Try:				
Finish:	L			

Reline	Upper Lower	Additions Up	per Lower	Fini	S
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CUSTOM MADE DEVICE Certified as conforming to the Medical Device Directive

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 product has been manufactured by either or both CosTech or Laxmi Dental Laboratory (based at Kandivali West, Mumbai, India) with final inspection completed by CosTech acting as UK responsible person.

In the event that the prescriber has supplied some of the materials etc for incorporation in a particular custom made dental appliance then this appliance cannot be guaranteed to fully meet with the applicable relevant essential requirements.

The grounds for placing such a device on the market is that the risk of compromising the patient's health and safety by using materials etc supplied

by the prescriber is assessed as minimal. This risk assessment relies upon a duly qualified medical practitioner's competence and duty to supply materials etc that is either from a CE marked source or from an appropriate European Competent Authority registered manufacturer of custom-made medical devices.

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.

Q.C. APPROVAL