



Prosthetic

www.ivodentech.co.uk

07900 190000

Ivo Dentech Dental Lab

Ivo Dentech Ltd

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NHS
Prosthetics

Private
Prosthetics

Dentist Account Details

Empty box for Dentist Account Details

Patient & Job Details

Patient's Name: _____ Job ID: _____

Date Started: _____

M F

Vita Shade

Chrome Upper Lower

Reline Upper Lower

Additions Upper Lower

Repair Upper Lower

Acrylic Upper Lower

ComFlexin® Upper Lower

Due Dates

Special Tray: U L _____

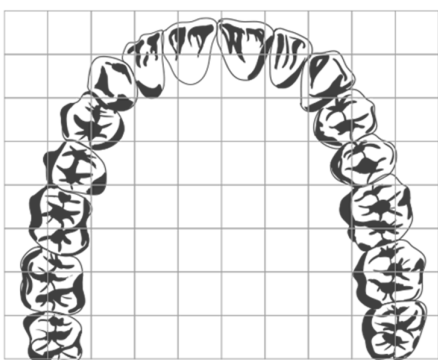
Bite Block: _____

Try-In: _____

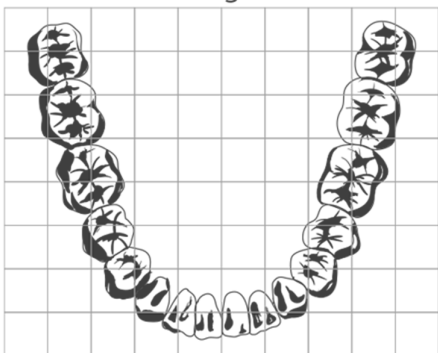
Re-Try: _____

Finish: _____

Please circle teeth to be included



Design



Notes:

CUSTOM MADE DEVICE Certified as conforming to the Medical Device Directive, manufactured and supplied by Ivo Dentech Ltd.

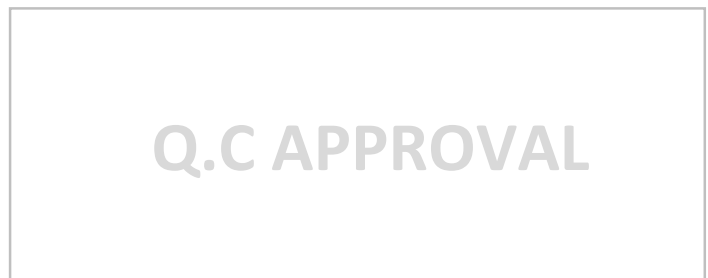
The product here packed is a custom-made device for the named patient stated above and conforms to the essential requirements set out in Annex 1 of the EC Medical Device Directive 93/42/EEC dated June 1993 and if any of these requirements are not fully met the details are documented on reverse or attached and despatched to the user.

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 Dr named above takes responsibility for prescribing the correct material/ alloy in accordance to current regulations in their region.*

For full terms and conditions please see our website on www.ivodentech.co.uk

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.



Please note these items are NOT sterile. MHRA Registration: 8999