Examples of statements for custom-made medical devices

The following products are listed for guidance only. Some of these products will also be available as mass-produced, rather than custom-made medical devices and must be classified according to Part II of the <u>UK Medical Devices Regulations 2002</u> (as amended) (UK MDR 2002), Annex IX as modified by Part I of Schedule 2A to the UK MDR 2002).

Maxillofacial

The patients for these types of devices are predominantly seen within a clinical environment. The easiest and simplest way of communicating to the patients that the statement is available to them if they wish to have it would be to place posters within the clinics themselves. As these patients' records stay with them throughout their treatment the statement itself could be placed on the patient's medical files and would be available to them if requested. The industry Journal could be used to alert the health professionals of their responsibility in this area. Below is a sample of a statement provided by Addenbrookes Maxillofacial Unit, containing all of the elements needed in the statement, which could be used by a maxillofacial manufacturer.

Custom Made Medical Device Do Operator : Forename: Peter Surname: N Address : Maxillofacial Laboratory (Box 4	owak A	Maxillofsciel Laboratory (Box 47), AddenBrooke's Hospital N.H.S.Trust, Hills Road, Cambridge CB2 2QQ			
Addenbrooke's Hospital NHS T	rust				
Hills Road	T T	Tel 01223 216636			
Cambridge		Fax 01223 216708			
Cambs CB2 2QQ	N	M.D.A. Reg. No. CA002628			
Patients Forename: Tester Patients Surna Date: 30/04/2009		Your Ref: 0123456 Our Ref: 006245			
Reviewed and accepted subject to sight of Date: 90/04/2009	positive model. Signed:				
Reasons not signed if construction is to pr	opeed:				
Device(s): Obturator clear baseplate U/					
Prescription: See operators instruction request sheet.					
N-1-1-1	Supplier	Use by Date	Batch No.	CE Mark	
Material	K.C.Smth. Morrouthshire	-	088021	Yes	
Stainfees stool vire 0.5mm dis		31/03/2010	Lat 06CP01	Yes	
Acrylic H/C C&J Delux powder (velned)	Chaperlin & Jacobs	30/09/2010	Let 20136	Yes	
Acrylic Universal Denture liquid clear	Brecon Ltd	0000000		1	
				-	
				-	
				+	
				-	
b. Reasons not fully met:					
Signed:	Date disp	atched: 01/05/200	9		
				-	
ESSENTIAL REQUIREMENTS Madisfracial Laboratory (Size A7), Addeniil rooke Hitle Rood, Cambridge CRI 2040	's Hospilal N.H.S.Trust,	This statement of been offered to no			
M.D.A. Reg. No. CA003026 This applieroe Production. e. is a CUSTOM MADE DEVICE.	s, oheractedation, properties and testures	☐ Accepted ☐ Declined			
specified on the prescription. Any relevant con-	to the section of the	Signed:			
 is for exclusive use by the nan d. Conforms to the essential requirements set on 	high patients. The Section Spectrum 98/42 EEC.	Guardian:			
THIS DEVICE IS SUPPLIED IN A NON STERIL	Date:				

Artificial eyes

The National Artificial Eye Service (NAES) operates clinics around the country where patients are seen by their clinicians and the device is then fitted at the clinic when it has been made by the production laboratory in Blackpool. The patient's records stay with NAES throughout their treatment period and the statement could be placed on the medical file and accessed when the patient requests to see it. A poster campaign would also work within this environment and could be placed in clinics across the country. Below is a sample of a statement by the eye service which contains all of the elements needed in the statement.

The device conforms to the essential requirements (Part II of the UK MDR 2002, Annex I [as modified by Part II of Schedule 2A to the UK MDR 2002])

signed	Print name	
Date		
	conformity has been offered to the named patient.	
□ Accepted □ Dec	lined	
Signed		
Guardian		
Date		

Dental devices

Custom-made dental devices are made by dental laboratories by prescription from dentists. The statement in this instance is dispatched by the laboratory (manufacturer) to the dentist with the device. Unlike the previous examples the patient's dental records do not provide a history of treatment and prescriptions because they do not move with the patient when he/she changes dentist. Nevertheless, the dentist who prescribes and fits the appliance will be the 'health professional' responsible for making the patient aware of the availability of the statement and supplying it on request. To make dentists aware of this obligation, relevant dental journals could carry news items on this requirement change and a press statement from the relevant professional bodies (GDC & BDA) would also ensure dental professionals were aware of their obligation.

There are a number of ways to inform patients about the statement and how they can obtain it. These range from verbal (at the end of the treatment), posters in the surgery, or a note on the receipt for the treatment charge. If the statement is requested then a signature from the patient could be requested by the dentist as proof of fulfilling their obligation. A reference may also be made in the leaflet the Department issues on dental charges.

An example of a statement for custom-made dental products provided by the DLA is attached below which contains all of the elements in the statement.

Example A – combined laboratory ticket and patient prescription/information

A N Other Dental Laboratory 44 Wollaton Road Nottingham NG9 2NR 0115 9254888				р	APPL lease comple rescription a site. If you h	ete the and se nave a cripti	end both parts	PTION sections of this to the address vith the use of this	
PATIENT'S NAME			NAME	OF PRES	CRIBER CLINIC NAME AND ADDRESS (if applicable)				
Date sent:			Date rec	quired			Lat	o reference (wh	ere applicable)
Type of Orthodontic Denture Metal ca		sting	Crown & Bridge		Bite raiser	Splint			
е	Obturator	Facial		Body prosthes	iis	Nightguard	ı	Implant	Bleaching tray
Please [Y]		Τ'							

INSTRUCTIONS AND AMENDMENTS RECORD	OUTLINE OF DESIGN REQUIRED
INSTRUCTIONS AND AMENUMENTS RECORD	OUTLINE OF DESIGN REQUIRED
	BY LABORATORY PERSONNEL ONLY Approved for release by:
Approved for manufacture by:	Approved for release by:
Sign:	Sign:
Details of materials etc supplied by prescriber	Details of any model approval by prescriber
Initials:	Initials:
ORIGIN OF MANUFAC	TURE DECLARATION
	wholly manufactured within the EU.
	□ No
(If no, detail manufa	acturing locations below)
1.	
2.	
Your attention is drawn to the following statement: This is to satisfy the design characteristics and properties specified b device is intended for exclusive use by this patient and conformation.	y the prescriber for the above named patient. This medical
****	been repaired and/or refurbished for an individual patient's se.
Storing, handling and instructions for use: It is recomment and safe environment that prevents it from coming into contact could cause physical or chemical damage to the medical devior temperature during storage. Where applicable, you should from its model. Where applicable, instructions on how to use c	It with materials, equipment, acids, alkalies or bleaches that ce. The medical device should not be subjected to extremes take care not to damage the medical device when removing it

For the purposes of the above diagram, please read:

• 'EU' as 'UK'

A N Other Dental Laboratory 44 Wollaton Road Nottingham NG9 2NR 0115 9254888			PATIENT PRESCRIPTION AND CUSTOM MADE APPLIANCE INFORMATION				
REGISTERED WITHT	HEUK COMPETENT AUT	THORITY CADODOO	If you have any queries regarding the fit or performance of your appliance you should contact the prescribing dentist for further information.				
PATIENT'S NAME		NAME OF PRESCRIBER		CLINIC NAME AND ADDRESS			
	DATE OF APPLIANCE IS MANUFACTURE		F TECHNICAL ORT	LAB REFERENCE			
Product Code	Description/T	ype of Appliance	Quantity	Standard of work NHS/Private	Comments		
		DICINI OF MANUEAU		O.H.			
ORIGIN OF MANUFACTURE DECLARATION This complete appliance has been wholly manufactured within the EU.							
			☐ Yes☐ Noacturing locations b	elow)			
	3. 4						
Your attention is dra to satisfy the design of device is intended for	haracteristics and	properties specified b	y the prescriber for t	he above named patie	ent. This medical		
This statement does not apply to medical devices that have been repaired and/or refurbished for an individual patient's use.					ndividual patient's		

For the purposes of the above diagram, please read:

• 'EU' as 'UK'

External prosthetics, orthotics and wheelchairs including their seating Systems

In the case of the above custom-made devices the statement remains with the healthcare professional. A wheelchair can be a one-off custom-made device, but in the majority of cases a seating system manufactured to the profile/needs of the individual user (custom-made device) is fitted to a wheelchair (a medical device) and, as is with external prosthetics and orthotics, each patient receives a product care leaflet. There could be additional information added to the leaflet informing the patient about the statement, and how they can request it. If it is requested, a system could be used to confirm receipt by the patient such as a stamp or a signature. The Patients Association could inform their members of the right for them to ask for the statement and possibly a poster campaign could be used to reach this patient group. An example of a statement has been provided by the British Healthcare Trade Association is attached below and contains all elements in the statement.

DRAFT

Manufacturers name and address:

device and quoting the order number above.

Custom-made medical device

This custom-made orthotic is intended for the exclusive use of (patient's name)
Order number:
Orthotist/prosthetist:
Clinic:
This custom-made orthotic has been manufactured for the above patient to the specification provided by the above named clinician.
The device conforms to the essential requirements (Part II of the UK MDR 2002, Annex I [as modified by Part II of Schedule 2A to the UK MDR 2002]).
SignedPrint name
Date

NB: The patient can obtain a copy of this statement by contacting the manufacturer of the