PRE-ACQUISITION QUESTIONNAIRE (PAQ Form)

The purpose of this questionnaire is to support the pre-acquisition assessment and approval of proposals to procure Medical Devices and accessories under purchase, exchange, rental, lease, loan, donation or other agreements. Please ensure that all relevant sections have been completed; (questions for which the only available response is 'YES' indicate that this response is a requirement, if the question applies) - and that all supplementary information requested has been provided; (shaded boxes indicate that supplementary information is required).

Note: The term 'Device', as used here, includes equipment and systems; in the case of systems the requirements below apply both to the individual constituent Devices and to the configured system as a whole. (Accessories within the scope of the return need to be identified under 1(d).)

PART I

to be completed by the device Manufacturer or Authorised Representative

PF	ROD	OUCT DETAILS:										
U	DI De	evice Identifier:										
Device Description: (GMDN Code / Group if available)			Weighing S	Scale								
Type:		Make:	Marsden									
		Model:	HM-200D									
Ма	anufa	acturer:	Charder	Charder Electronics Company								
Su	ıpplie	er:	Marsden	Marsden Weighing Machine Group Ltd								
EL	J Aut	horised Representative:	Marsden	Marsden Weighing Machine Group Ltd								
1 a) When was this Model first placed upon the market ?										200	9	
	b)	Is this Model still in production ? NO \square YES \boxtimes if NO, when did production							cease ?			
	c)	Any outstanding Field Safety Corrective Actions / Field Safety Notices ? NO 🗵 YES 🗆 All issued Notices / Alerts a								YES 🔲		
	d)	Does this return cover a range of Model variants ? NO ☒ YES ☐ If YES, list of Models attach Does this return cover Accessories ? NO ☒ YES ☐ If YES, list of Accessories a								YES 🔲		
	e) f)	Does this return cover Acce Has a Device brochure and		haan attachad	to this return 2	NO 🗵	YES 🗀	If YES, list o	of Accessories a	ttached to this	return ?	YES ☐ YES ☒
	')	rias a Device brochare and	Specification	been attached	to this return :							ILS Z
RF	FGU	LATORY COMPLIA	ANCE:									
2	a)	Does the Device meet the E	_	irements of all	Lourrently applica	ahla FC D	iractivas 2				NO \square	YES ⊠
_	b)	Which EC Directive/s apply	•	incincino or un	rearrently applied	abic LC D	irectives :				140	125
	,	Medical Devices Directive				С	lassification	n?		←	(1, 1-m, 1-s / II	a / IIb / III)
		Active Implantable Devices	Directive							1		
		In-Vitro Diagnostics Medical Device Directive						← (gene	ral / self-test / Lis	t-A / List-B)		
		Other/s	044/24/EU N	lan A., tamati	Najabina Tua							
		- which Directive/s? 20	014/31/EU N	on Automatio	c Weighing Ins	trument	5				_	
3	a)	Is the Device CE-Marked, fo		-	,		tives ?				NO 🗆	YES 🖾
	b)	- if YES, have the EC Declar	ration/s of Co	nformity been	attached to this i	return ?						YES 🔲
4		If not CE-marked, (or if 'off-	-label' use is p	proposed for a	CE-marked Devi	ce). then	-					
a) Is this a Medical Device for 'Clinical Investigation' ?						7	NO ⊠	YES				
		- if YES, quote the MHRA 'no objection' reference										VEC 🗖
	if YES, has a copy of the MHRA's notice of 'no objection' been attached to this return ?b) Is this an In-Vitro Diagnostic Medical Device for 'Performance Evaluation'?							NO ⊠	YES YES			
	U)	Is this an In-Vitro Diagnostic Medical Device for 'Performance Evaluation'? - if YES, has a copy of notification to MHRA been attached?								110 🔼	YES 🔲	
c) Is this a 'custom-made' Medical Device ?									NO 🛛	YES 🗌		
		- if YES, name the prescribi	ng Medical Pr	actitioner:]		
d) - if NO to 3(a), and to 4(a) (b) and (c), then provide justification of the Device's status -												
A Height Measure												
5 a) Which EC conformity assessment route/s have been adopted?												
☐ full QA ☐ type examination ☐ product verification								production QA	\			
product QA unit verification internal control (self declaration)						No \square	VEC 🗖					
	b) Has this included Notified Body conformity assessment ? NO ☐ YE - Notified Body identification number & name:						TES 🗌					
	c)	Is the manufacturer current		L	nent system Stan	dards ?					NO 🗆	YES 🗆
	- which Standard/s ? ISO9001:2015 / SGS & 2014/31/EU ← (eg: EN-ISO-9001, 13485, 1											
			GS]		

PI	ROE	DUCT COMMITMENT:	
6	a)b)c)d)e)f)	To what date is product support for this Model guaranteed? Does this include training; servicing, repair & availability of parts; supply of consumables / accessories? What is the Device warranty period? Does the manufacturer / supplier have a robust system for notification of Device alerts / upgrades to a named hospital representative? What is the recommended working lifetime for this Device? N/A C (not applicable for disposable Devices) Have details for end-of-life waste management of the Device been attached to this return?	YES ☑ YES ☑ YES ☑
PI	ROE	DUCT SUPPORT:	
7	a) b) c)		YES ☐ YES ☐ YES ☑
_		Commissioning & Deple	yment
8	a) b)	Has a protocol for post-delivery acceptance testing of Device function and safety been attached to this return? Does the Device have particular installation requirements and / or require ancillary services or other prerequisite arrangements? NO - if YES, then have details of all installation requirements been attached to this return?	YES ☐ YES ☐
		Technical S	upport
9	a)	Does the manufacturer or an authorised servicing agent provide a maintenance / repair service ? - if YES, then have details of all service contract options been detailed, fully costed and attached to this return ?	YES ☑ YES ☐
		- where is the servicing facility located ? - are all servicing staff verifiably trained and competency assessed for the servicing tasks that they perform ?	YES 🖾
		- are qualification / competency records of servicing staff available upon request ?	YES 🛛
	b)	Is the servicing organisation currently certified to any management system Standards? - which Standard/s? IS09001:2015 / SGS & 2014/31/EU ← (eg: EN-ISO-9001, 13485,	YES 🛮 17025, etc.)
	c)	- Certification Body: SGS Do the contract alternatives offered in 9(a) include an option for in-house equipment servicing by hospital staff? NO NO	YES 🗆
	c)	- if YES, have details of information / test equipment / tooling / software required for equipment servicing been attached to this return? - if YES, have details of information / test equipment / tooling / software required for equipment servicing been attached to this return?	YES T
_		Decontam	ination
10	a)	What level of Device decontamination / reprocessing is required ? ☐ single-use ☐ cleaning ☐ disinfection ☐ sterilisation	
	b)	If not single-use, have validated decontamination protocol/s been attached to this return?	YES 🛛
	c)	For sterilisable Devices, do these instructions meet the requirements of EN-ISO-17664?	YES 🗆
	d) e)	Have all requirements for special reprocessing equipment, tools and materials been detailed in the attached information? Have any special post-processing Device storage requirements been detailed in the attached information?	YES YES
	f)	Is there a limit to the number of Device reprocessing cycles ? NO YES If YES, what is the limit ?	123
	g) h)	Are Devices uniquely identifiable ? Is this an implantable Device ? NO ☑ YES ☐ YES ☐	
		Data S	ecurity
11	a)	Does the Device store or transmit patient information that will require information governance measures ?	YES 🗌
	L	- if YES, then have details of information capture / encryption / storage / transmission / deletion been attached to this return?	YES 🔲
	b)	Does the Device interface, by wired or wireless connection, with Information technology (IT) equipment or network systems? - if YES, then have details of Device IT software / hardware compatibility requirements been attached to this return? - if YES, then have details of provisions made for Device IT cybersecurity been attached to this return?	YES TYES TYES
		Particular Requir	ements
12	a)	Does the Device present particular hazards that require special safety management measures ? (eg: ionising / non-ionising radiation; contamination / infection; hazardous materials; hazardous mechanical / electrical energy; etc.)	YES 🗆
		- identified hazards:	
	ы	- if YES, then have details of the nature of identified hazards been attached to this return?	YES ☐ YES ☑
	b)	Does the Device require particular performance quality assurance measures ? (eg: calibration, qualification, PoCT controls, etc.) - QA measures: Periodical Calibration check	IES M
		- if YES, then have details of quality assurance requirements been attached to this return ?	YES 🔲

I۱	1PLE	MENTATION SUPPORT:								
13	a) 1	Is competency-based user training available from the manufacturer or an a	uthorised provider	sed provider ?						
		- if YES, have details of user training offered (amount / content / assessme	YES 🗆							
	b)	Is competency-based technical (equipment servicing) training available from the manufacturer or an authorised provider?								
		- if YES, have details of technical training offered (amount / content / asse	ssment / duration /	location / cost / etc.) been attached	? YES □					
	c) 1	Is competency-based decontamination / reprocessing training available fro	m the manufacture	r or an authorised provider ?	NO ☐ YES ☐					
		- if YES, have details of decontamination training offered (amount / content / assessment / duration / location / cost / etc.) been att								
	d) /	Are qualification / competency records of training providers available upon	request ?		YES □					
	etc.), have details of these been atta	ached? YES								
DI	ECLA	RATION:								
Ple	ase ens	sure that all necessary supplementary information, (as indicated by shaded	boxes 🔲 in the Fo	rm above) accompanies this return.						
	1.c)	All issued Field Safety Notices / Alerts		ATTACHED □	NOT APPLICABLE 🛛					
	1.d)	List of all Model variants covered by this return		ATTACHED □	NOT APPLICABLE ⊠					
	1.e)	List of all Accessories covered by this return		ATTACHED □	NOT APPLICABLE 🛛					
	1.f)	Device brochure / specification		ATTACHED ⊠						
	3.b)	EC Declaration/s of Conformity		ATTACHED □						
	4.a)	MHRA's notice of 'no objection' for Medical Device 'Clinical Investigation'		ATTACHED □	NOT APPLICABLE 🛛					
	4.b)	Notification to MHRA for In-Vitro Diagnostic Medical Device 'Performance	Evaluation'	ATTACHED □	NOT APPLICABLE 🛛					
	6.c)	Warranty details		ATTACHED ⊠						
	6.f)	Details for end-of-life waste management of the Device		ATTACHED □						
	8.a)	Protocol for post-delivery Device acceptance testing		ATTACHED □						
	8.b)	Details of installation requirements		ATTACHED □	NOT APPLICABLE 🛛					
	9.a)	Service support contract options for maintenance / repair		ATTACHED □						
	9.c)	Availability of spare / replacement parts		ATTACHED □	NOT APPLICABLE 🛛					
		Information / test equipment / tooling / software required for Device services	vicing	ATTACHED □	NOT APPLICABLE 🛛					
	10.b)	Validated decontamination protocol/s		ATTACHED □	NOT APPLICABLE 🛛					
	10.d)	Requirements for special reprocessing equipment, tools and materials		ATTACHED □	NOT APPLICABLE ⊠					
	10.e)	Details of special post-processing Device storage requirements	ATTACHED □	NOT APPLICABLE 🛛						
	11.a)	a) Details of patient information capture / encryption / storage / transmission / deletion ATTA			NOT APPLICABLE ⊠					
	11.b)	Details of Device IT software / hardware compatibility requirements	ATTACHED □	NOT APPLICABLE ⊠						
		Details of provisions made for Device IT cybersecurity	Details of provisions made for Device IT cybersecurity							
	12.a)	Details of particular hazards that require special safety management	ATTACHED □	NOT APPLICABLE ⊠						
	12.b)	Details of particular performance quality assurance measures required		ATTACHED □	NOT APPLICABLE 🛛					
	13.a)	Details of user training offered		ATTACHED □	NOT APPLICABLE 🛛					
	13.b)	Details of technical training offered		ATTACHED □	NOT APPLICABLE ⊠					
	13.c)	Details of decontamination training offered		ATTACHED □	NOT APPLICABLE ⊠					
	13.e)	Details of any additional support facilities offered		ATTACHED □	NOT APPLICABLE ⊠					
Νh	en refe	erence is made to this Form and its attachments within the process of obtai	ning the specified p	product/s, we agree that the purchase	er will be entitled to rely					
Jpo	on the c	contents and that subsequent non-compliance with the statements contain	ed herein will entitle	e the purchaser to seek redress.	,					
Name:		Donna Jebson	Donna Jebson							
Р	osition	n: Sales Manager								
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Email:		sales@marsdengroup.co.uk	Telephone:	01709 364296						
S	ignatu	re: D. Jebson	Date:	01/01/2021						
					· 					

PART II

for completion by the device Supplier (eg: Manufacturer, Authorised Representative or other)

Previous sections in PART I provided for general Device information; this PART II addendum provides details specific to particular transaction/s for supply of the product:-

TRA	TRANSACTIONAL:										
14 a)	On wha	at basis will the product be spurchase?	supplied, (including Device exchange?	es for clinical inverges rental/lease? [-	earch) ? an ? 🔲	donation ?				
b)	For sup	ply by loan or donation, oth	ner than Devices for clinic	al investigation /	research -						
	Has a D	Department of Health (DH)	MIA Call-Off Agreement F	orm been attache	ed ?				YES 🔲		
	Is the S	Supplier on the DH Master I	ndemnity Agreement (MI	A) Register ? *				NO 🗆	YES 🗌		
	- if YES	, then quote DH MIA regist	ration number:								
	- if NO, has an Indemnity Insurance Certificate (for local indemnity agreement with the customer) been attached ?										
	(* Note: unregistered Suppliers are advised to register for the MIA Overarching Agreement with the DH)										
c)	c) For supply by loan or donation of Devices for clinical investigation / research -										
	Has confirmation of Health Research Authority (HRA) indemnity approval been attached?								YES 🔲		
d)	d) Is the particular item to be supplied a pre-used product ?								YES 🗌		
	- if YES, has usage and full service history been attached with this return ?								YES 🔲		
Nam	e:	Donna Jebson									
Position:		Sales Manager									
Company:		Marsden Weighing Group									
Address:		Unit 1 Genesis Business Park. Sheffield Rd. Rotherham. S60 1DX									
Signature:		D. Jebson			Date:	01/01/202	1				