





**SECTION 2. MANUFACTURER LICENCES**

**MANDATORY**

**INSTRUCTIONS:** Complete the information in Table 2 below.

**One (1) certified copy (.pdf)** of each licence must also be submitted.

**Translations** - Any original document in languages other than English must be accompanied by an official notarized English translation.

Manufacturers that do not provide valid licences will not be considered.

The submission of invalid licenses will result in the automatic suspension of all of the manufacturer’s products until valid licenses are provided.

Table 2 – Mandatory Licencing

	Formal document no.	Issue date (DD-MMM-YY)	Expiry date (DD-MMM-YY)
<b>MANDATORY:</b> Company license or Registration document			
<b>Optional:</b> Manufacturing license			
<b>Optional:</b> Other:.....			

**SECTION 3. MANUFACTURER CERTIFICATES**

**MANDATORY**

**INSTRUCTIONS:** Complete the information in Table 2 below.

**One (1) certified copy (.pdf) of each certificate** is required.

**Translations** - Any original document in languages other than English must be accompanied by an official notarized English translation.

Manufacturers that do not provide valid certificates will not be considered. Email confirmation of a pending certificate validation will not be accepted.

The submission of invalid certificates will result in the automatic suspension of all manufacturer products until valid certificates are provided.

**A website link to a certification authentication page is mandatory** for certification verification. **PLEASE VERIFY THAT WEBLINK FUNCTIONS.**

**All certification bodies should be ILAC accredited.**

**Certificates expiring within three months of the Annual Review** (i.e. between April to June 2025) must be accompanied by a copy of a renewal request letter.



Table 3 – Mandatory Certification

Requirements per IMD-PQS category	Certificate	Certificate accreditation year (e.g. 2015)	Certification authority (certified by)	Website link for certification verification <b>MANDATORY</b>	Expiry date (DD-MMM-YY)
<b>ALL CATEGORIES</b> <i>Except E008 &amp; E013</i>	ISO 9001:2015*				
E001 E003 E004	ISO 14001:2015*				
E008 E013	ISO 13485:2016*				
Optional	93/42/EEC				
Other approval marks (e.g. CE, UL, FDA or EMA):					

**SECTION 4. CHANGES TO MANUFACTURING INFORMATION** **MANDATORY**

**INSTRUCTIONS:** If no changes have occurred, or are expected to occur, between April 2024 and March 2025, please tick **NO**.  
If changes **HAVE** occurred, manufacturers must submit documentation verifying the amended information.

Table 4 – Changes to manufacturing information

		<i>Please provide details of changes:</i>
Are there any changes to the <b>company name</b> and/or <b>status</b> ?	YES <input type="checkbox"/> NO <input type="checkbox"/>	
Are there any changes to the <b>manufacturing site</b> ?	YES <input type="checkbox"/> NO <input type="checkbox"/>	