

# GUIDANCE FOR COMPLETING SFDA MDMA APPLICATION FORM

## JAPANESE JURISDICTION (JP)

Medical Device Sector  
Registration & Licenses Department

Revision: 20/06/2017  
V 1.0

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## Section 1. Manufacturer (Japan)

| No  | SFDA Question              | Do<br>Check<br>Warning | Task   |
|-----|----------------------------|------------------------|--|
| 1.1 | Manufacturer               | Do                     | <ul style="list-style-type: none"> <li>Select the name of the manufacturer from the drop list</li> </ul>   |
| 1.2 | Legal<br>Manufacturer      | Do                     | <ul style="list-style-type: none"> <li>Select the name of the <b>Legal manufacturer.</b></li> </ul>  |
|     |                            | Check                  | <ul style="list-style-type: none"> <li>The name and address of the devices' manufacturer in this application must concur with sections:                             <ul style="list-style-type: none"> <li>2.1.10 Labeling</li> <li>2.1.11 IFU</li> <li>2.3 A/C Power Supply Statement – if applicable</li> <li>2.4 Environmental Statements</li> <li>2.5 The provided documents in this section</li> <li>5.3 EC certificates</li> <li>5.3 Recent Audit report</li> <li>5.3 Other Certificates as required by the device class</li> <li>5.4 Declaration of conformity</li> <li>6.3 QMS Certificate</li> <li>7.1 Regulatory Compliance Attestation</li> </ul> </li> </ul> |
|     |                            | Warning                | <ul style="list-style-type: none"> <li>A common error is to select the device manufacturing site address, rather than the manufacturer address.</li> <li>If the manufacturer has two addresses a postal address and a Site address, please provide attested letter from the manufacturer explaining that there are two addresses – insert the letter in 2.1.10</li> </ul>  |
| 1.3 | Medical Device<br>Category | Do                     | <ul style="list-style-type: none"> <li>Use SFDA Drop-down list of 17 Categories</li> </ul>   |

## Section 2. General info. (Japan)

| No                     | SFDA Question   | Do<br>Check<br>Warning | Task   |                     |                  |                  |              |                        |                              |     |        |
|------------------------|---|------------------------|--|---------------------|------------------|------------------|--------------|------------------------|------------------------------|-----|--------|
| 2.1                    | Details of the medical devices applying for marketing authorization     | Do                     | <ul style="list-style-type: none"> <li>Insert the list of devices in the application; make sure NOT to list the MODELS separately as different devices.</li> <li>For IVD, List the kit as a single item in section 2.1 for labels &amp; IFU, all the labels of the reagent in the kit must be provided.</li> </ul>   |                     |                  |                  |              |                        |                              |     |        |
|                        |   | Check                  | <ul style="list-style-type: none"> <li>If the application contains multiple products then check that the listed products are meeting the bundling criteria. Refer to SFDA bundling rules in MDS-G7: <a href="http://www.sfda.gov.sa/en/medicaldevices/regulations/DocLib/MDS-G7.pdf">http://www.sfda.gov.sa/en/medicaldevices/regulations/DocLib/MDS-G7.pdf</a></li> <li>Cross check the list of the devices against: Labels (2.1.10) IFU (2.1.11)</li> <li>If the device has multiple models, for example male urinary catheters of different sizes, the applicant should include these in one line: <table border="1"> <thead> <tr> <th>Product description</th> <th>Intended Purpose</th> <th>Trade/Brand Name</th> <th>Model Number</th> </tr> </thead> <tbody> <tr> <td rowspan="3">Male urinary Catheters</td> <td rowspan="3">Drain urine from the bladder</td> <td rowspan="3">XYZ</td> <td>123-40</td> </tr> <tr> <td>123-50</td> </tr> <tr> <td>123-60</td> </tr> </tbody> </table> </li> </ul> | Product description | Intended Purpose | Trade/Brand Name | Model Number | Male urinary Catheters | Drain urine from the bladder | XYZ | 123-40 |
| Product description    | Intended Purpose  | Trade/Brand Name       | Model Number   |                     |                  |                  |              |                        |                              |     |        |
| Male urinary Catheters | Drain urine from the bladder  | XYZ                    | 123-40   |                     |                  |                  |              |                        |                              |     |        |
|                        |   |                        | 123-50   |                     |                  |                  |              |                        |                              |     |        |
|                        |   |                        | 123-60   |                     |                  |                  |              |                        |                              |     |        |
| 2.1.1                  | Product Brief Description (This field will appear on the MDMA printout) | Do                     | <ul style="list-style-type: none"> <li>Insert the product brief description.</li> <li><b>Note:</b> The product description will be printed on the MDMA license issued by the SFDA.</li> </ul>  |                     |                  |                  |              |                        |                              |     |        |
|                        |   | Check                  | <ul style="list-style-type: none"> <li>The product description must be precise and informative (Maximum of 100 Characters including spaces).</li> <li>The description must be in English only, no commas (acceptable if it makes sense "part of the sentence" e.g.: infant, pediatric &amp; adult ventilator), clear &amp; accurate.</li> <li>Make sure there is no spelling errors.</li> </ul>  |                     |                  |                  |              |                        |                              |     |        |

|       |   |                |  |
|-------|---|----------------|--|
|       |   | <b>Warning</b> | <ul style="list-style-type: none"> <li>• “Catheter, Urinary” will be rejected whereas “Urinary Catheter” is acceptable.</li> <li>• Do <b>NOT</b> include Brand Names or Company Names <b>UNLESS</b> the brand name is descriptive (Describing the product).</li> </ul> |
| 2.1.2 | Intended Purpose of the medical device type | <b>Do</b>      | <ul style="list-style-type: none"> <li>• Insert the intended purpose.</li> </ul>   |
|       |   | <b>Check</b>   | <ul style="list-style-type: none"> <li>• Typically this is an extract from IFU</li> </ul>  |
| 2.1.3 | Product Trade/Brand Name.                   | <b>Do</b>      | <ul style="list-style-type: none"> <li>• Insert the product Trade/Brand name as it appears on the label.</li> <li>• <b>Note:</b> The product Trade/Brand Name will be printed on the issued MDMA License.</li> </ul>   |
|       |   | <b>Check</b>   | <ul style="list-style-type: none"> <li>• Check it concurs with the product trade/brand name as it appears on the product actual label.</li> </ul>  |
|       |   | <b>Warning</b> | <ul style="list-style-type: none"> <li>• The combination of the Product Description and Trade/Brand Name must be unique for every device listed in the application</li> </ul>  |
| 2.1.4 | Model Name/Number                           | <b>Do</b>      | <ul style="list-style-type: none"> <li>• Insert the model name/number as it appears on the label</li> </ul>  |
|       |   | <b>Check</b>   | <ul style="list-style-type: none"> <li>• If more than one model number listed in the specified section, these models should only differ in color, size, weight, dimensions or shape.</li> </ul>  |
|       |   | <b>Warning</b> | <ul style="list-style-type: none"> <li>• If the product has model/ref number, the brand name must not be repeated in section 2.1</li> </ul>  |
| 2.1.5 | Manufacturer’s Device Identifier Number     | <b>Do</b>      | <ul style="list-style-type: none"> <li>• Insert the Manufacturers Device Identifier Number</li> </ul>  |
|       |   | <b>Check</b>   | <ul style="list-style-type: none"> <li>• Typically this is the REF number, or Product catalogue number. Check it concurs with the product ID number as it appears on the label.</li> </ul>   |

| 2.1.6                   | Format of medical device identifier number(s) that will appear on labelling for traceability purposes | DO                      | <ul style="list-style-type: none"> <li>Insert the Format of medical device identifier number(s) that will appear on the labeling <u>for traceability purposes</u></li> </ul>  |            |                         |                         |                |
|-------------------------|---|-------------------------|---|------------|-------------------------|-------------------------|----------------|
|                         |   | Check                   | <ul style="list-style-type: none"> <li>Typically this is the LOT number, or Serial number Provide a brief description of how the number is formatted e.g. LOT YYYY-MM-DD (Year-Month-Day)</li> </ul>  |            |                         |                         |                |
| 2.1.7<br>2.1.8<br>2.1.9 | Nomenclature code number<br>GMDN UMDNS<br>Other(e.g. FDA identification number, JMDN)                 | DO                      | <ul style="list-style-type: none"> <li>Insert the nomenclature code number if available.</li> </ul>   |            |                         |                         |                |
| 2.1.10                  | Provide the label(s) affixed to the device or its wrappers when it is supplied to the KSA             | DO                      | <ul style="list-style-type: none"> <li>Attach the device labels for ALL devices listed in section 2.1</li> </ul> <p>A/C Power Supply<br/>If the product is connected to an A/C power supply, the applicant must provide images of the power supply label, so it can be cross-checked with the A/C Power Supply signed declaration provided in section 2.3 (60 Hz supply at nominal values of either 230 or 400 volts)</p> <ul style="list-style-type: none"> <li>For IVD Kits, all the individual reagent labels must be provided.</li> <li>All labels must be in compliance with Japanese Directive office of medical devices Evaluation and licensing in MHLW and national provisions.</li> </ul> |            |                         |                         |                |
|                         |   | Check                   | <ul style="list-style-type: none"> <li>Check that labels provided for <b>ALL</b> the devices listed in section 2.1, including each of the models numbers/REF/Part No. /etc.</li> <li>When the device has range (e.g. sizes) then a representative label is acceptable provided that clearly links one product-size to one product ID number.</li> </ul> <p>Example (Acceptable).:</p> <table border="1"> <thead> <tr> <th>Trade Name</th> <th>REF (Product ID Number)</th> <th>Size (Product variable)</th> </tr> </thead> <tbody> <tr> <td>Medical Device</td> <td>1234</td> <td>5x5cm</td> </tr> </tbody> </table>  | Trade Name | REF (Product ID Number) | Size (Product variable) | Medical Device |
| Trade Name              | REF (Product ID Number)   | Size (Product variable) |   |            |                         |                         |                |
| Medical Device          | 1234  | 5x5cm                   |   |            |                         |                         |                |

|                |                         |                | <table border="1" style="width: 100%; border-collapse: collapse;"> <tr> <td style="width: 33%;">Medical Device</td> <td style="width: 33%;">1236</td> <td style="width: 33%;">10x10cm</td> </tr> </table> <p>The applicant has provided a clear link between each of the product ID numbers and the product sizes/dimensions</p> <ul style="list-style-type: none"> <li>The table must be from the legal Manufacturer and must be signed with name, job title &amp; date.</li> <li>The labels must contain:             <ol style="list-style-type: none"> <li>Device Trade name (See 2.1.3)</li> <li>Device model number (See 2.1.4)</li> <li>Device ID Number (REF) (See 2.1.5)</li> <li>Legal Manufacturers Name &amp; Address (See 1.1 &amp; 1.2)</li> </ol> </li> <li>Labels May contain:             <ol style="list-style-type: none"> <li>LOT or Serial Number</li> <li>Power Supply – if applicable</li> <li>Storage Temperature</li> <li>Expiry Date</li> <li>Date of manufacture</li> <li>Sterile &amp; method – if applicable</li> <li>Single use – if applicable</li> <li>The term “Made in ....” With the country of origin</li> <li>IVD – if applicable</li> <li>IVD Self test – if applicable</li> <li>Rx only – if applicable</li> </ol> </li> <li><b>Note: if the device is for professional use only:</b><br/>It is acceptable if the label provided in English only<br/><u>Reference: SFDA MDS-IR6 Article 9 (C)</u></li> <li><b>Note: If the device is for home use / Self Test (IVD)</b><br/>The label provided for each product must be written in both English &amp; Arabic languages.<br/><u>Reference: SFDA MDS-IR6 Article 9 (C)</u></li> </ul> | Medical Device | 1236                    | 10x10cm        |      |  |      |  |     |
|----------------|-------------------------|----------------|--|----------------|-------------------------|----------------|------|--|------|--|-----|
| Medical Device | 1236                    | 10x10cm        |  |                |                         |                |      |  |      |  |     |
|                |                         | <b>Warning</b> | <ul style="list-style-type: none"> <li>Common error is wrong or missing label.</li> <li>Tables, example (NOT Acceptable)</li> </ul> <table border="1" style="width: 100%; border-collapse: collapse;"> <thead> <tr> <th style="width: 50%;">Trade Name</th> <th style="width: 50%;">REF (Product ID Number)</th> </tr> </thead> <tbody> <tr> <td>Medical Device</td> <td>1234</td> </tr> <tr> <td></td> <td>1236</td> </tr> <tr> <td></td> <td>etc</td> </tr> </tbody> </table> <p>The applicant has NOT provided link between the product ID numbers and the product sizes/dimensions.</p>  | Trade Name     | REF (Product ID Number) | Medical Device | 1234 |  | 1236 |  | etc |
| Trade Name     | REF (Product ID Number) |                |  |                |                         |                |      |  |      |  |     |
| Medical Device | 1234                    |                |  |                |                         |                |      |  |      |  |     |
|                | 1236                    |                |  |                |                         |                |      |  |      |  |     |
|                | etc                     |                |  |                |                         |                |      |  |      |  |     |

|        |                                    |                |  |
|--------|------------------------------------|----------------|--|
| 2.1.11 | Provide Instructions For Use "IFU" | <b>DO</b>      | Attach the IFU for ALL devices listed in section 2.1 and it comply with by the Ministry of Health, Labour and Welfare (MHLW) or the Pharmaceutical and Medical Devices Agency (PMDA) in Japan  |
|        |                                    | <b>Check</b>   | <ul style="list-style-type: none"> <li>IFU cover ALL the devices Trade/Brand Names listed in section 2.1</li> <li>Check that legal Manufacturers name &amp; address are printed on the IFU and it concurs with sections 1.1 &amp; 1.2</li> <li>Check it contains Electrical rating –if applicable-.</li> <li><b>Note: if the device is for professional use only</b><br/>It is acceptable if the label provided in English only<br/><u>Reference: SFDA MDS-IR6 Article 9 (C)</u></li> <li><b>Note: If the device is for home use / Self Test (IVD)</b><br/>The label provided for each product must be written in both English &amp; Arabic languages.<br/><u>Reference: SFDA MDS-IR6 Article 9 (C)</u></li> <li>Check it contains any WARNING &amp;/or Precautions to take.</li> <li>Device's models &amp; IDs (must match with 2.1 – if mentioned). If IFU does not cover all models or IDs or Doesn't match with sections 2.1.4 &amp; 2.1.5, a justification is required.</li> <li>Storage Temperature: Min &amp; Max (Where required)</li> <li>Sterility Method (Where required)</li> <li>Single use: (Where required)</li> <li>Power requirement if applicable (60 Hz Supply at nominal values or either 230 or 400 Volts)</li> <li><b>NOTE: IF IT IS NOT RELEVANT TO HAVE AN IFU FOR THE PRODUCT, THEN THE APPLICANT MUST PROVIDE A JUSTIFICATION FROM THE MANUFACTURER &amp; MUST BE SIGNED, JOB TITTLE &amp; DATED.</b></li> </ul> |
|        |                                    | <b>Warning</b> | <ul style="list-style-type: none"> <li>A common error is a wrong or missing information OR missing IFU with no justification</li> </ul>  |



|                       |  |         |  |
|-----------------------|--|---------|--|
| 2.1.12                | List of Accessory  | DO      | <ul style="list-style-type: none"> <li>List the accessory for the device (If applicable)</li> </ul>  |
|                       |  | Check   | <ul style="list-style-type: none"> <li>Accessory Definition: Accessories are devices specifically intended by its legal Manufacturer to be used together with the medical device to achieve its intended purpose.</li> </ul>   |
|                       |  | Warning | <ul style="list-style-type: none"> <li>If the Accessory can be used as a stand-alone medical device, the <b>SFDA</b> do <b>NOT</b> consider it as an Accessory. It must be listed as a Medical Device.</li> </ul>  |
| 2.1.12.1 to 2.1.12.11 |  | DO      | <ul style="list-style-type: none"> <li>Same requirements as sections 2.1.1 to 2.1.11</li> </ul>  |
| 2.2                   | Jurisdiction(s) Where this medical device may be placed on the market. | DO      | <ul style="list-style-type: none"> <li>Make selections as appropriate.</li> <li>Minimumly, Japan (JP) must be selected because this is a JP submission.</li> </ul>   |
| 2.3                   | Statement of Power Supply requirements for KSA Market                  | DO      | <ul style="list-style-type: none"> <li>If the device/accessory is connected to an a/c power supply, complete the statement Template provided (<b>It must be printed on the <u>Manufacturer's Letterhead</u></b>)</li> <li>The statement will confirm the medical device is: <ol style="list-style-type: none"> <li>Designed to operate with a 60 Hertz supply at nominal values of either 230 or 400 Volts.</li> <li>Is fitted with the appropriate a/c power connector.</li> <li>Maintains the required electrical safety conditions.</li> <li>Continues to perform to specification.</li> </ol> <b>(The statement must be signed, job tittle &amp; dated from the manufacturer)</b> </li> <li>If it is <b>NOT</b> applicable select the (N/A) box move to the next section.</li> </ul> |

|     |   |                |   |
|-----|---|----------------|---|
|     |   | <b>Check</b>   | <ul style="list-style-type: none"> <li>The statement is signed, job title &amp; dated from the manufacturer on its letterhead</li> </ul>  |
|     |   | <b>Warning</b> | <ul style="list-style-type: none"> <li>Do <b>NOT</b> alter wording of the SFDA template</li> </ul>  |
| 2.4 | Statement for KSA Environmental Factors   | <b>DO</b>      | <ul style="list-style-type: none"> <li>Complete the statement template provided printed on Manufacturers letterhead and make sure it is signed, job title &amp; dated.</li> </ul>   |
|     |   | <b>Check</b>   | <ul style="list-style-type: none"> <li>The statement is signed, job title &amp; dated from the manufacturer on its letterhead</li> </ul>  |
|     |   | <b>Warning</b> | <ul style="list-style-type: none"> <li>Do <b>NOT</b> alter wording of the SFDA template</li> </ul>  |
|     |   |                |   |
| 2.5 | Statement to correctly stored, transported, installed, maintained & disposed of medical devices | <b>DO</b>      | <ul style="list-style-type: none"> <li>Provide a copy of the manufacturer's instructions to ensure that the medical devices intended to be placed on the KSA market will be correctly stored, transported, installed, maintained &amp; disposed of, and that users can be trained in their proper use and maintenance.</li> <li>The applicant may also provide additional information that they believe is relevant to this request. The additional information must be from the Manufacturer (Letterhead) and must be signed, job title &amp; dated, also, list devices and accessories Trade/Brand Name in this application (See 2.1) or the application number.</li> </ul> |

|     |  |         |   |
|-----|--|---------|---|
| 2.6 | Manufacturers advertising & marketing material intended for use in KSA | DO      | <ul style="list-style-type: none"> <li>Provide a copy of the manufacturer's advertising and marketing material intended for use in the KSA, if <b>NOT AVAILABLE</b> provide an <u>explanation</u> and the <u>date when such material will become available</u>, if <b>NOT REQUIRED</b> provide a <u>justification</u>.</li> <li><b>Note:</b> It is acceptable to provide marketing materials that includes more devices than the listed in 2.1, however, the sections of the marketing material that includes the listed products should be <b>HIGHLIGHTED</b></li> <li><b>Note:</b> A product catalogue (Soft copy or hard copy) is considered marketing material and is acceptable.</li> </ul>  |
|     |  | Check   | <ul style="list-style-type: none"> <li>Marketing literature is provided for at least <b>ALL</b> the devices listed in section 2.1</li> <li>It should contain the manufacturer name</li> <li><b>Note:</b> It is NOT necessary to have the address of the Manufacturer on the Marketing literature.</li> <li>It must contains the document control reference number</li> <li><b>Note: if the device is for professional use only</b><br/>It is acceptable if the marketing material provided in English only<br/><u>Reference:</u> SFDA MDS-IR6 Article 9 (F)</li> <li><b>Note: If the device is for home use / Self Test (IVD)</b><br/>The Marketing material provided must be written in both English &amp; Arabic languages.<br/><u>Reference:</u> SFDA MDS-IR6 Article 9 (F)</li> </ul> |
|     |  | Warning | <ul style="list-style-type: none"> <li>A common error is to state there is no marketing literature when a product catalogue is available</li> <li>A common error is to provide a marketing material without a document control reference number</li> <li>A common error when the Marketing literature includes more devices than the listed in section 2.1 and the relevant devices sections is <b>NOT</b> Highlighted.</li> <li>Marketing literature is <b>NOT</b> acceptable if provided on online-link or website.</li> </ul>  |

### Section 3. Jurisdiction. (Japan)

| No  | SFDA Question           | Do<br>Check<br>Warning | Task   |
|-----|-------------------------|------------------------|--|
| 3.1 | Desired<br>Jurisdiction | Do                     | Ensure <b>Japanese</b> jurisdiction has been selected. |



### Section 4. Product Categories (Japan)

| No  | SFDA Question                                | Do<br>Check<br>Warning | Task  |
|-----|--|------------------------|---|
| 4.1 | Device Type                                  | Do                     | Select one correct option: <ul style="list-style-type: none"> <li>• IVD</li> <li>• Medical Device</li> </ul>  |
| 4.2 | Device Classification                        | Do                     | Select one correct option: <ul style="list-style-type: none"> <li>• Medical device: <ol style="list-style-type: none"> <li>1- Class I</li> <li>2- Class II</li> <li>3- Class III</li> <li>4- Class IV</li> </ol> </li> <li>• IVD: <ol style="list-style-type: none"> <li>1- Class I</li> <li>2- Class II</li> <li>3- Class III</li> </ol> </li> </ul> |
|     |  | Check                  | The selected Device classification is correct for the devices listed in 2.1, and it must concurs the classification stated in section 5.6 (Manufacturer's Declaration of Conformity to Japanese regulations)  |
| 4.5 | Designated Controlled (only for Class II MD) | Do                     | <ul style="list-style-type: none"> <li>• Select (Yes/No) based on the certificates provided in section 5</li> </ul>   |

## Section 5. Product Verification (Japan)

| No  | SFDA Question  | Do<br>Check<br>Warning | Task  |
|-----|--|------------------------|---|
| 5.1 | Indicate whether the MHLW or Registered Certification Body (RCB) has issued a marketing authorization / certification to the Marketing Approval Holder and provide its identification number | Do                     | <ul style="list-style-type: none"> <li>Select <b>YES</b>. However, For Class I both answers are acceptable depend on the current conformity certification.</li> </ul>   |
| 5.2 | Government Approval (SHONIN)   | DO                     | <ul style="list-style-type: none"> <li>This section is applicable for class II, III and IV. (If NOT applicable type N/A)</li> <li>Type the certificate number provided in section 5.5</li> </ul>  |
| 5.3 | Certification (NINSHO)   | DO                     | <ul style="list-style-type: none"> <li>This section is applicable for class II designated controlled and some of class III (If NOT applicable type N/A).</li> <li>If applicable type certificate number provided in section 5.5.</li> </ul>   |
| 5.4 | Notification (TODOKEDE)  | DO                     | <ul style="list-style-type: none"> <li>This section is applicable for class I.</li> <li>Type the notification number of the approval letter provided in Section 5.5</li> </ul>  |
| 5.5 | Provide copy of the current certificate of conformity  | DO                     | <ul style="list-style-type: none"> <li>Provide a copy of the current certificate of conformity, issued by MHLW or an authorized RCB, that confirms the technical file been reviewed to asses the device's design. Where applicable, the certificate should be accompanied by a translation into English.</li> <li>Providing FSC from MHLW is NOT enough.</li> </ul> |
|     |  | Check                  | <ul style="list-style-type: none"> <li>Check that the approval provided is correct depending on classification of the device &amp; current conformity certification.</li> </ul>   |

|     |   |              |   |
|-----|---|--------------|---|
|     |   |              | <ul style="list-style-type: none"> <li>All approval/certificate must be provided in Japanese with a reliable English translated copy from the certification body or a third party.</li> </ul>   |
| 5.6 | <b>Provide copy of the manufacturer current Declaration of Conformity (DOC) to Japanese regulations</b> | <b>DO</b>    | <ul style="list-style-type: none"> <li>Provide DOC that covers all the listed products in section 2.1</li> <li>Make sure it covers the following: <ol style="list-style-type: none"> <li>1- Device name</li> <li>2- Device classification</li> <li>3- RCB name (if applicable)</li> <li>4- Certificate number (if applicable)</li> <li>5- Laws applied</li> <li>6- Legal Manufacturer (Name &amp; Address)</li> <li>7- Signed, Job title &amp; dated</li> </ol> </li> </ul> |
| 5.7 | <b>Name of RCB involved with assessing the design</b>   | <b>DO</b>    | <ul style="list-style-type: none"> <li>Type the name of the RCB involved in assessing the design</li> </ul>   |
|     |   | <b>Check</b> | <ul style="list-style-type: none"> <li>Check that it is authorized by MHLW</li> </ul>   |

### Section 6. Manufacturer's QS Status (Japan)

| No  | SFDA Question  | Do<br>Check<br>Warning | Task   |
|-----|--|------------------------|--|
| 6.1 | Indicate whether the manufacturer of the medical device operates an established quality management system (QMS) that complies with the required conformity assessment procedures | Do                     | <ul style="list-style-type: none"> <li>Select the correct answer</li> </ul>  |
| 6.2 | If YES, indicate the QMS standard used   | Do                     | <ul style="list-style-type: none"> <li>Type the standard used for QMS</li> </ul>   |
| 6.3 | Description of the medical devices covered by the manufacturer's QMS.  | Do                     | <ul style="list-style-type: none"> <li>Type the description and confirm it concur with the notification of conformity in section 6.6.</li> </ul> |
| 6.4 | Indicate the procedures that are included within the manufacturer's QMS.   | Do                     | <ul style="list-style-type: none"> <li>Select one or more, depends on the scope of QMS.</li> </ul>   |



|     |   |       |  |
|-----|---|-------|--|
| 6.5 | Indicate whether the QMS is regularly audited by PMDA, the prefectural government, or a RCB.  | Do    | <ul style="list-style-type: none"> <li>Select Yes OR No depending on the devices listed</li> <li><b>NOTE:</b> "NO" can be selected only if the device is unspecified class I device.</li> </ul>  |
| 6.6 | If YES name the body that undertakes the audit and provide a copy of the current notification of conformity, where applicable, accompanied by a translation into English. | Do    | <ul style="list-style-type: none"> <li>Provide the following if "Yes" was selected in (6.6) if applicable: <ul style="list-style-type: none"> <li>1- Copies of notification of conformity (TSUCHISHO) or certificate of standard conformity (TEKIGOSHO) written in Japanese with English copies from the certification body or a third party.</li> <li>2- Applied Standard: "MO 169" for class I designated controlled II, III and IV. Any other applicable standard for class I undesignated controlled.</li> </ul> </li> </ul> |
|     |   | Check | <ul style="list-style-type: none"> <li>Check the validity of the provided documents.</li> <li>That the device name or product group cover devices in section 2.1.</li> <li>The name &amp; address of the manufacturer is matching section 1.1 / 1.2.</li> </ul>  |

## Section 7. Other National Provisions (Japan)

| No  | SFDA Question   | Do<br>Check<br>Warning | Task  |
|-----|---|------------------------|---|
| 7.1 | <b>Manufacturer Declaration</b>   | <b>Do</b>              | <ul style="list-style-type: none"> <li>Complete the attestation using the SFDA template provided</li> <li>It must be printed on manufacturers Letterhead</li> <li>Select Japan for Japanese based application</li> <li>The attestation letter must be signed, name, job title and dated</li> </ul>  |
| 7.2 | <b>Provide the address of the location where the manufacturer holds technical information to support this attestation</b> | <b>Do</b>              | <ul style="list-style-type: none"> <li>Provide the address of the location where the manufacturer holds the technical information to support this attestation.</li> </ul>   |
|     |   | <b>Check</b>           | The <b>FULL</b> Postal address must be provided (Building, number, road, city, postal code, state, country)   |
| 7.3 | <b>Authorized Representative (AR) Declaration</b>   | <b>Do</b>              | <ul style="list-style-type: none"> <li>Complete the attestation using the SFDA template provided</li> <li>It must be printed on <b>AR / Local Manufacturers (LM)</b> Letterhead</li> <li>Write the correct application number</li> <li>The attestation letter must be signed with name, job title and dated in English and it must match the MDNR information (Arabic &amp; English is Acceptable)</li> </ul> |