



Our Ref : (45) dlm. MDA. 100-1/7/2

Date : 30 December 2019

**OBSOLETE**

**CIRCULAR LETTER OF THE MEDICAL DEVICE AUTHORITY  
NO. 2 YEAR 2018 (REVISION 2)**

**POLICY ON IMPLEMENTATION AND ENFORCEMENT UNDER THE MEDICAL  
DEVICE ACT 2012 (ACT 737):**

**CONTROL OF ORPHANED, OBSOLETE AND DISCONTINUED MEDICAL DEVICE  
IN HOSPITAL, HEALTHCARE FACILITIES INSTITUTION OR ANY RELATED  
FACILITIES**

**PURPOSE**

1) The purpose of this circular is to set the implementation and enforcement under the Medical Device Act 2012 (Act 737) relating to control of orphaned, obsolete and discontinued medical device in hospital, healthcare facilities institution or any related facilities in Malaysia.

**BACKGROUND**

2) Orphaned medical device means an existing medical device in a hospital, healthcare facilities, or any related facilities where the manufacturer or authorized representative has ceased operation.

3) Obsolete medical device means an existing medical device in a hospital, healthcare facilities, or any related facilities which is outdated, or no longer being manufactured due to design changes, evolution of new technologies.

4) Discontinued medical device means an existing medical device in a hospital, healthcare facilities, or any related facilities that is no longer in the distribution.

- 5) Section 5(1) no medical device shall be imported, exported or placed in the market unless the medical device is registered under this Act.
- 6) Section 5(2) any person who contravenes section 5(1) commits an offence and shall, on conviction, be liable to a fine not exceeding two hundred thousand ringgit or to imprisonment for a term not exceeding three years or to both.
- 7) Section 15 (1) states that no establishment shall import, export or place in the market any registered medical device unless it holds an establishment licence granted under this Act.
- 8) Section 77, Act 737, states that, the Minister may, if he considers it consistent with the purposes of this Act or in the interest of public health and safety, by order published in the *Gazette*, exempt any person or medical device from any of the provisions of this Act or any regulations made under this Act for such duration and subject to such conditions as the Minister may specify and he may alter or add the conditions so specified.
- 9) All medical devices must be registered before being imported, exported or placed in Malaysian market. However, there are orphaned, obsolete and discontinued medical devices in hospitals or healthcare facilities institution which is not registered and still being used. Some of these medical devices can not be registered because the technical documentation of the medical device is unavailable and unable to comply with registration requirements.
- 10) Therefore, to impose certain controls to the healthcare facilities institution that are still continue to use such medical devices, as well as ensure that medical services in hospital or healthcare facilities institution are uninterrupted, there is a need to develop a policy to control this medical device.

## **POLICY DECISION FOR IMPLEMENTATION AND ENFORCEMENT**

**11) The Medical Device Authority Meeting No 3/2019 has decided to set the implementation and enforcement control of orphaned, obsolete and discontinued medical device in hospital, healthcare facilities institution or any related facilities as follows:**

- a) An orphaned, obsolete and discontinued medical device is exempted from medical device registration and establishment licence requirement;**
- b) Establishment or healthcare facilities institution which have orphaned, obsolete or discontinued medical devices shall identify and provide the notification / listing to the Medical Device Authority.**
- c) The risk of using orphaned, obsolete and discontinued medical device is under the responsibility of establishment, users and healthcare facilities institution.**
- d) Establishment shall be responsible for post-market issues on any obsolete or discontinued medical device at least in accordance with the projected useful life of the medical device as determined by the manufacturer; and**
- e) Compliance with notification requirement with certain charges as may be specified by the Authority.**

**12) This exemption is implemented administratively before order is published in the Gazette.**

## **USAGE AND EFFECTIVE DATE**

**13) This Circular shall be used as part of requirements under Act 737 and shall be effective from the date it is issued.**

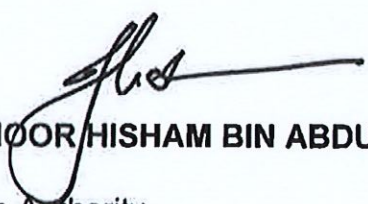
## ENQUIRIES

- 14) Any enquiries relating to this circular can be forwarded to:

Chief Executive  
Medical Device Authority  
Ministry of Health Malaysia  
Level 6, Prima 9, Prima Avenue II,  
Block 3547, Persiaran Apec,  
63000 Cyberjaya, Selangor, MALAYSIA  
Tel.: (+603) 8230 0300, Fax: (+603) 8230 0200  
Email: [mdb@mda.gov.my](mailto:mdb@mda.gov.my)

Thank you

**"BERKHIDMAT UNTUK NEGARA"**



**(DATUK DR NOOR HISHAM BIN ABDULLAH)**  
Chairman  
Medical Device Authority  
Ministry of Health Malaysia