



Tel: (+603) 8230 0332/0356

Portal: [www.mda.gov.my](http://www.mda.gov.my)

Email: [cabregistration@mda.gov.my](mailto:cabregistration@mda.gov.my)



CERTIFICATION  
INTERNATIONAL  
MALAYSIA

CI INTERNATIONAL CERTIFICATION SDN. BHD.  
NO. 37-4 JALAN SP 2/2 TAMAN SERDANG PERDANA  
43000 SERI KEMBANGAN  
SELANGOR DARUL EHSAN  
TEL: +603 – 8942 9001 FAX: +603 – 8942 9002

Person Responsible:-

MS. OOI SOO KANG

[[sook@cimalaysia.com.my](mailto:sook@cimalaysia.com.my)]

Contact Person:-

MS. LIAU FEI LING

[[liau@cimalaysia.com.my](mailto:liau@cimalaysia.com.my)]

Validity MDA/CAB-012: 10/04/2018 – 09/04/2021

## SCOPE OF REGISTRATION

### Conformity Assessment of Quality Management System (QMS)

(1)	GDPMD	Good Distribution Practice for Medical Devices
(2)	ISO 13485	Quality Management Systems for Medical Devices – Requirements for Regulatory Purpose

### Conformity Assessment of Medical Device Technical Areas

(3)	MD 0103	Non-active orthopedic and rehabilitation devices
(4)	MD 0106	Non-active instruments
(5)	MD 0107	Contraceptive medical devices
(6)	MD 0108	Non-active medical devices for disinfecting, cleaning, rinsing
(7)	MD 0202	Non-active orthopedic implants
(8)	MD 0401	Non-active dental equipment and instruments
(9)	MD 0403	Dental implants
(10)	MD 1111	Software

### Conformity Assessment of Medical Device Technical Areas

(11)	VERIFICATION	Conformity Assessment by Way of Verification
------	--------------	--

*Section 10(1), Medical Device Act 2012 (Act 737) and Regulation 8, Medical Device Regulation 2012*