# CAB REGISTRATION NUMBER: MDA/CAB-001 VALIDITY: 21/11/2022 - 20/11/2025





TÜV SÜD (MALAYSIA) SDN. BHD. NO. 18 JALAN ASTAKA U8/82 BUKIT JELUTONG 40150 SHAH ALAM SELANGOR DARUL EHSAN TEL: +603-7859 8822 FAX: +603-7859 8824

PERSON RESPONSIBLE: **DR. VINCENT LAM CHEE CHOONG** [vincent.lam@tuv-sud.my]

**CONTACT PERSON: DR. VINCENT LAM CHEE CHOONG** [vincent.lam@tuv-sud.my]

# SCOPE OF REGISTRATION

MDS 7005

Con	Conformity Assessment of Quality Management System (QMS) and Post-Market Surveillance System (PMSS)		
1	ISO 13485	Quality Management System for Medical Devices – Requirements for Regulatory Purpose	
2	GDPMD	Good Distribution Practice for Medical Devices	
Con	formity Assessn	nent of Technical Documentation	
<b>Con</b> 3	<b>iformity Assessn</b> MD 0101	nent of Technical Documentation Non-active devices for anesthesia, emergency and intensive care	
	6		

MD 0107 Bandages and wound dressings MD 0301 6 7 Medical devices referencing the Directive 89/686/EEC on personal protective equipment (PPE)

**Conformity Assessment by Way of Verification** VERIFICATION Conformity Assessment by Way of Verification 8

## CAB REGISTRATION NUMBER: **MDA/CAB-002** VALIDITY: **21/11/2022 – 20/11/2025**

## MEDCERT MALAYSIA SDN. BHD.

NO. 18 3<sup>rd</sup> FLOOR JALAN SS19/1D 47500 SUBANG JAYA SELANGOR DARUL EHSAN TEL: **+603-5131 4773** FAX: **+603-5124 7688** 

PERSON RESPONSIBLE: MR. SIEW YEW KONG [siew.kong@medcert.de]

CONTACT PERSON: MR. SIEW YEW KONG [siew.kong@medcert.de]

Сот	Conformity Assessment of Quality Management System (QMS) and Post-Market Surveillance System (PMSS)	
1	ISO 13485	Quality Management System for Medical Devices – Requirements for Regulatory Purpose
2	GDPMD	Good Distribution Practice for Medical Devices

Con	formity Assessm	ent of Technical Documentation
3	MD 1111	Software
4	MD 1201	Imaging devices utilizing ionizing radiation
5	MD 1202	Imaging devices utilizing non-ionizing radiation
6	MD 1302	Monitoring devices of vital physiological parameters
7	MD 1402	Devices utilizing non-ionizing radiation

Cor	nformity Assessm	ent by Way of Verification
8	VERIFICATION	Conformity Assessment by Way of Verification





# CAB REGISTRATION NUMBER: **MDA/CAB-003** VALIDITY: **21/11/2022 – 20/11/2025**

SGS MALAYSIA SDN. BHD. LOT 3 & 4 PERSIARAN JUBLI PERAK SEKSYEN 22, 40300 SHAH ALAM SELANGOR DARUL EHSAN TEL: +603-7627 0080 FAX: +603-2093 8202





PERSON RESPONSIBLE: MR. KENNY LOOI TUCK KIAN [kenny.looi@sgs.com]

CONTACT PERSON: MR. KAMARRUZAIMISHAM BIN HARUN [zaimie.harun@sgs.com]

# SCOPE OF REGISTRATION

Coi	Conformity Assessment of Quality Management System (QMS) and Post-Market Surveillance System (PMSS)	
1	ISO 13485	Quality Management System for Medical Devices – Requirements for Regulatory Purpose
2	GDPMD	Good Distribution Practice for Medical Devices

Con	formity Assessm	nent of Technical Documentation
3	MD 0101	Non-active devices for anesthesia, emergency and intensive care
4	MD 0102	Non-active devices for injection, infusion, transfusion and dialysis
5	MD 0105	Non-active ophthalmologic devices
6	MD 0108	Non-active medical devices for disinfecting, cleaning, rinsing
7	MD 0204	Non-active soft tissue implants
8	MD 0301	Bandages and wound dressings
9	MD 0303	Other medical devices for wound care
10	MD 1101	Devices for extra-corporal circulation, infusion and haemopheresis
11	IVD 0304	Hereditary disease: phenylketonuria
12	IVD 0307	Tumoral marker: PSA
13	IVD 0404	Molecular biology

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#### SIRIM QAS INTERNATIONAL SDN. BHD.

CAB REGISTRATION NUMBER: **MDA/CAB-004** VALIDITY: **21/11/2022 – 20/11/2025** 





# BLOK 4, 1<sup>st</sup> FLOOR, SIRIM COMPLEX NO. 1 PERSIARAN DATO' MENTERI SEKSYEN 2, 40700 SHAH ALAM SELANGOR DARUL EHSAN TEL: +603-5544 6483 FAX: +603-5544 6763

PERSON RESPONSIBLE: PN. NUR FADHILAH BINTI MUHAMMAD [fadhilah@sirim.my]

> CONTACT PERSON: MR. MD ZAINI BIN MD JAI [mdzaini@sirim.my]

#### SCOPE OF REGISTRATION

Со	Conformity Assessment of Quality Management System (QMS) and Post-Market Surveillance System (PMSS)	
1	ISO 13485	Quality Management System for Medical Devices – Requirements for Regulatory Purpose
2	GDPMD	Good Distribution Practice for Medical Devices

Con	formity Assessm	ent of Technical Documentation
3	MD 0102	Non-active devices for injection, infusion, transfusion and dialysis
4	MD 0107	Contraceptive medical devices
5	MD 0108	Non-active medical devices for disinfecting, cleaning, rinsing
6	MD 0202	Non-active orthopaedic implants
7	MD 0204	Non-active soft tissue implants
8	MD 0301	Bandages and wound dressings
9	MD 0303	Other medical devices for wound care
10	MD 0403	Dental implants
11	MD 1101	Devices for extra-corporal circulation, infusion and haemopheresis
12	MD 1102	Respiratory devices, including hyperbaric chambers for oxygen therapy, inhalation anaesthesia
13	MD 1104	Active surgical devices
14	MD 1106	Active dental devices
15	MD 1107	Active devices for disinfection and sterilisation
16	MD 1109	Active devices for patient positioning and transport
17	MD 1111	Software
18	IVD 0203	Hepatitis B, C and D
19	IVD 0303	Congenital infections: rubella, toxoplasmosis
20	IVD 0307	Tumoral marker: PSA
21	IVD 0401	Clinical chemistry
22	IVD 0404	Molecular biology
23	IVD 0405	Pregnancy and ovulation
24	MDS 7002	Medical devices utilizing tissues of animal origin, including Directive 2003/32/EC
25	MDS 7005	Medical devices referencing the Directive 89/686/EEC on personal protective equipment (PPE)

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# CAB REGISTRATION NUMBER: **MDA/CAB-005** VALIDITY: **21/11/2022 – 20/11/2025**

## BSI SERVICES MALAYSIA SDN. BHD. SUITE 29.01 LEVEL 29 THE GARDENS NORTH TOWER MID VALLEY CITY, LINGKARAN SYED PUTRA 59200 KUALA LUMPUR TEL: +603-9212 9638 FAX: +603-9212 9639



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PERSON RESPONSIBLE: MS. EVELYN CHYE POH YIN [Evelyn.Chye@bsigroup.com]

CONTACT PERSON: MR. RAJAKUMARAN A/L KARNAGARAN [Rajakumaran.Karnagaran@bsigroup.com]

# SCOPE OF REGISTRATION

Co	Conformity Assessment of Quality Management System (QMS) and Post-Market Surveillance System (PMSS)	
1	ISO 13485	Quality Management System for Medical Devices – Requirements for Regulatory Purpose
2	GDPMD	Good Distribution Practice for Medical Devices

Con	nformity Assessm	ent of Technical Documentation
3	MD 0101	Non-active devices for anesthesia, emergency and intensive care
4	****MD 0402	Dental materials
5	MD 1301	Monitoring devices of non-vital physiological parameters
6	MD 1302	Monitoring devices of vital physiological parameters
7	IVD 0401	Clinical chemistry
8	IVD 0404	Molecular biology

\*\*\*\* means approval only for conformity assessment on dental dam.

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## CAB REGISTRATION NUMBER: **MDA/CAB-006** VALIDITY: **11/09/2023 – 10/09/2026**





**DQS CERTIFICATION (M) SDN. BHD.** SUITE 43-3, SETIA AVENUE SU13/S SETIA ALAM 40170 SHAH ALAM

SELANGOR DARUL EHSAN TEL: +603-3342 3259 FAX: +603-3358 3299

PERSON RESPONSIBLE: MR. DANNY NG KIM YAU [danny.ng@dqs.com.my]

CONTACT PERSON: PN. NUR KHAIRUN HANIS BINTI ABU BAKAR [hanis.bakar@dqs.com.my]

Conformity Assessment of Quality Management System (QMS) and Post-Market Surveillance System (PMSS)			
1	ISO 13485	Quality Management System for Medical Devices – Requirements for Regulatory Purpose	
2	GDPMD	Good Distribution Practice for Medical Devices	
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Cor	Conformity Assessment of Technical Documentation		
3	MD 0101	Non-active devices for anesthesia, emergency and intensive care	

	Conformity Assessment by Way of Verification		
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### CAB REGISTRATION NUMBER: **MDA/CAB-007** VALIDITY: **11/09/2023 – 10/09/2026**



# TÜV RHEINLAND MALAYSIA SDN. BHD.

NO. 27 JALAN U8/103 METROPOLITAN BUSINESS PARK BUKIT JELUTONG, 40150 SHAH ALAM SELANGOR DARUL EHSAN TEL: **+603-7859 3000** FAX: **+603-7859 8020** 

PERSON RESPONSIBLE: SITI FAIRUS BINTI SAHUL HAMID [Siti.Hamid@tuv.com]

CONTACT PERSON: SITI NORFARHANAH BINTI MOHAMAD SAIPOL BAHRI [Farhanah.Saipol@tuv.com]

## SCOPE OF REGISTRATION

Cor	Conformity Assessment of Quality Management System (QMS) and Post-Market Surveillance System (PMSS)		
1	ISO 13485	Quality Management System for Medical Devices – Requirements for Regulatory Purpose	
2	GDPMD	Good Distribution Practice for Medical Devices	

Con	Conformity Assessment of Technical Documentation		
3	MD 0101	Non-active devices for anesthesia, emergency and intensive care	
4	MD 0102	Non-active devices for injection, infusion, transfusion and dialysis	
5	MD 0104	Non-active medical devices with measuring function	
6	MD 0106	Non-active instruments	
7	MD 0107	Contraceptive medical devices	
8	MD 0108	Non-active medical devices for disinfecting, cleaning, rinsing	
9	MD 0202	Non-active orthopaedic implants	
10	MD 0301	Bandages and wound dressings	
11	MD 0401	Non-active dental equipment and instruments	
12	MD 0402	Dental materials	
13	IVD 0201	HIV infection (HIV 1 and 2)	
14	IVD 0202	HTLV I and II	
15	IVD 0203	Hepatitis B, C and D	
16	IVD 0307	Tumoral marker: PSA	
17	IVD 0309	Device for self-diagnosis: device for the measurement of blood sugar	
18	IVD 0401	Clinical chemistry	
19	IVD 0403	Immunology	
20	IVD 0405	Pregnancy and ovulation	
21	IVD 0406	Specimen receptacles	
22	MDS 7005	Medical devices referencing the Directive 89/686/EEC on personal protective equipment (PPE)	

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#### CAB REGISTRATION NUMBER: **MDA/CAB-008** VALIDITY: **11/09/2023 – 10/09/2026**



PERSON RESPONSIBLE: MS. SOPHIA HENG CHIEW LING [sophiaheng@tuv-nord.com]

CONTACT PERSON: MS. SOPHIA HENG CHIEW LING [sophiaheng@tuv-nord.com]

# SCOPE OF REGISTRATION

Cor	Conformity Assessment of Quality Management System (QMS) and Post-Market Surveillance System (PMSS)		
1	ISO 13485	Quality Management System for Medical Devices – Requirements for Regulatory Purpose	
2	GDPMD	Good Distribution Practice for Medical Devices	

Con	Conformity Assessment of Technical Documentation		
3	MD 0106	Non-active instruments	
4	MD 0107	Contraceptive medical devices	
5	MD 0108	Non-active medical devices for disinfecting, cleaning, rinsing	
6	MD 1111	Software	
7	MD 1301	Monitoring devices of non-vital physiological parameters	

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#### CARE CERTIFICATION INTERNATIONAL (M) SDN. BHD.

## CAB REGISTRATION NUMBER: **MDA/CAB-009** VALIDITY: **12/02/2024 – 11/02/2027**





NO 16-G, 16-1 JALAN FLORA 1/1, BANDAR RIMBAYU, 42500 TELOK PANGLIMA GARANG SELANGOR DARUL EHSAN TEL: **+603-80732788** FAX: **+603-80732688** 

> PERSON RESPONSIBLE: MR. FLEMING TEO CHIN SIONG [fleming@cciglobe.com]

CONTACT PERSON: MRS. NABILA SETH BINTI MOHD NIVEN [nabila.seth@cciglobe.com]

## SCOPE OF REGISTRATION

Cor	Conformity Assessment of Quality Management System (QMS) and Post-Market Surveillance System (PMSS)	
1	ISO 13485	Quality Management System for Medical Devices – Requirements for Regulatory Purpose
2	GDPMD	Good Distribution Practice for Medical Devices

Con	Conformity Assessment of Technical Documentation	
3	MD 0102	Non-active devices for injection, infusion, transfusion and dialysis
4	MD 0106	Non-active instruments
5	MD 1101	Devices for extra-corporal circulation, infusion and haemopheresis
6	MD 1102	Respiratory devices, including hyperbaric chambers for oxygen therapy, inhalation anesthesia
7	MD 1201	Imaging devices utilizing ionizing radiation
8	MD 1202	Imaging devices utilizing non-ionizing radiation
9	MD 1302	Monitoring devices of vital physiological parameters
10	IVD 0401	Clinical chemistry

 Conformity Assessment by Way of Verification

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 Conformity Assessment by Way of Verification

#### CI INTERNATIONAL CERTIFICATION SDN. BHD.

# CAB REGISTRATION NUMBER: **MDA/CAB-012** VALIDITY: **25/06/2021 – 24/06/2024**



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: MR. OOI SOO KANG [oosk@cimalaysia.com.my]

CONTACT PERSON: MS. LIAU FEI LING [liau@cimalaysia.com.my]

# SCOPE OF REGISTRATION

Conformity Assessment of Quality Management System (QMS) and Post-Market Surveillance System (PMSS)		
1	ISO 13485	Quality Management System for Medical Devices – Requirements for Regulatory Purpose
2	GDPMD	Good Distribution Practice for Medical Devices

		formity Assessm	ent of Technical Documentation
	3	MD 0107	Contraceptive medical devices

# **Conformity Assessment by Way of Verification**

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# KGS CERTIFICATION SDN. BHD.

CAB REGISTRATION NUMBER: **MDA/CAB-013** VALIDITY: **12/11/2021 – 11/11/2024**  NO. 15 LORONG BLM 5/4 BANDAR LAGUNA MERBOK 08000 SUNGAI PETANI KEDAH DARUL AMAN TEL: +604-441 1524 FAX: +604-441 0610

Medical Device AUTHORITY MALAYSIA



PERSON RESPONSIBLE: PN. NACHEYAKALA A/P ELUMALAI [admin@kgscert.com]

CONTACT PERSON: MS. NURUL IZZATI BINTI ABDUL GHANI [admin@kgscert.com]

## SCOPE OF REGISTRATION

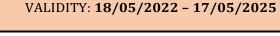
Co	Conformity Assessment of Quality Management System (QMS) and Post-Market Surveillance System (PMSS)	
1	ISO 13485	Quality Management System for Medical Devices – Requirements for Regulatory Purpose
2	GDPMD	Good Distribution Practice for Medical Devices

Con	Conformity Assessment of Technical Documentation		
3	MD 0103	Non-active orthopaedic and rehabilitation devices	
4	MD 0104	Non-active medical devices with measuring function	
5	MD 0105	Non-active ophthalmologic devices	
6	MD 0106	Non-active instruments	
7	MD 0108	Non-active medical devices for disinfecting, cleaning, rinsing	
8	MD 0109	Non-active devices for in vitro fertilization (IVF) and assisted reproductive technologies (ART)	
9	MD 0301	Bandages and wound dressings	
10	MD 1110	Active devices for in vitro fertilization (IVF) and assisted reproductive technologies (ART)	
11	IVD 0403	Immunology	
12	IVD 0404	Molecular biology	
13	IVD 0406	Specimen receptacles	
14	MDS 7005	Medical devices referencing the Directive 89/686/EEC on personal protective equipment (PPE)	

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# DNV GL INTERNATIONAL SDN. BHD.

LEVEL 18 MENARA PRESTIGE NO. 1 JALAN PINANG 50450 KUALA LUMPUR TEL: **+603-2160 1088** FAX: **+603-2160 1099** 



CAB REGISTRATION NUMBER: MDA/CAB-014





PERSON RESPONSIBLE: WAN AZIZUL HAFIZ BIN WAN ABDUL RAHMAN [Wan.Azizul.Rahman@dnv.com]

CONTACT PERSON: WAN AZIZUL HAFIZ BIN WAN ABDUL RAHMAN [Wan.Azizul.Rahman@dnv.com]

# SCOPE OF REGISTRATION

Cor	Conformity Assessment of Quality Management System (QMS) and Post-Market Surveillance System (PMSS)		
1	ISO 13485	Quality Management System for Medical Devices – Requirements for Regulatory Purpose	
2	GDPMD	Good Distribution Practice for Medical Devices	

Con	Conformity Assessment of Technical Documentation		
3	MD 0101	Non-active devices for anesthesia, emergency and intensive care	
4	MD 0102	Non-active devices for injection, infusion, transfusion and dialysis	
5	MD 0103	Non-active orthopaedic and rehabilitation devices	
6	MD 0104	Non-active medical devices with measuring function	
7	MD 0105	Non-active ophthalmologic devices	
8	MD 0106	Non-active instruments	
9	MD 0107	Contraceptive medical devices	
10	MD 0108	Non-active medical devices for disinfecting, cleaning, rinsing	
11	MD 0201	Non-active cardiovascular implants	
12	MD 0202	Non-active orthopaedic implants	
13	MD 0203	Non-active functional implants	
14	MD 0204	Non-active soft tissue implants	
15	MD 0301	Bandages and wound dressings	
16	MD 0302	Suture material and clamps	
17	MD 0303	Other medical devices for wound care	
18	MD 0401	Non-active dental equipment and instruments	
19	MD 0402	Dental materials	
20	MD 0403	Dental implants	
21	MD 1101	Devices for extra-corporal circulation, infusion and haemopheresis	
22	MD 1102	Respiratory devices, including hyperbaric chambers for oxygen therapy, inhalation anesthesia	
23	MD 1103	Devices for stimulation or inhibition	
24	MD 1104	Active surgical devices	
25	MD 1106	Active dental devices	
26	MD 1107	Active devices for disinfection and sterilization	
27	MD 1108	Active rehabilitation devices and active prostheses	
28	MD 1109	Active devices for patient positioning and transport	
29	MD 1110	Active devices for in vitro fertilization (IVF) and assisted reproductive technologies (ART)	
30	MD 1111	Software	
31	MD 1301	Monitoring devices of non-vital physiological parameters	
32	MD 1302	Monitoring devices of vital physiological parameters	

Conformity Assessment by Way of Verification

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# GENUINE DIAMOND SDN. BHD.

CAB REGISTRATION NUMBER: **MDA/CAB-016** VALIDITY: **22/11/2021 – 21/11/2024**  NO. 43B JALAN BP 7/12 BANDAR BUKIT PUCHONG 47120 PUCHONG SELANGOR DARUL EHSAN TEL: **+603-8069 1111** FAX: **+603-8069 1133** 

Medical Device AUTHORITY MALAYSIA



## PERSON RESPONSIBLE: PN. NUR ROSMARINIE BINTI BAHAROM [info@genuinediamond.com.my]

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Со	Conformity Assessment of Quality Management System (QMS) and Post-Market Surveillance System (PMSS)		
1	ISO 13485	Quality Management System for Medical Devices – Requirements for Regulatory Purpose	
2	GDPMD	Good Distribution Practice for Medical Devices	

Con	Conformity Assessment of Technical Documentation			
3	MD 0101	Non-active devices for anesthesia, emergency and intensive care		
4	MD 0102	Non-active devices for injection, infusion, transfusion and dialysis		
5	MD 0104	Non-active medical devices with measuring function		
6	MD 0106	Non-active instruments		
7	MD 0107	Contraceptive medical devices		
8	MD 0108	Non-active medical devices for disinfecting, cleaning, rinsing		
9	MD 0301	Bandages and wound dressings		
10	MD 0302	Suture material and clamps		
11	MD 0303	Other medical devices for wound care		
12	*MD 1101	Devices for extra-corporal circulation, infusion and haemopheresis		
13	MD 1102	Respiratory devices, including hyperbaric chambers for oxygen therapy, inhalation anesthesia		
14	MD 1103	Devices for stimulation or inhibition		
15	MD 1104	Active surgical devices		
16	MD 1105	Active ophthalmologic devices		
17	MD 1106	Active dental devices		
18	MD 1107	Active devices for disinfection and sterilization		
19	MD 1109	Active devices for patient positioning and transport		
20	MD 1201	Imaging devices utilizing ionizing radiation		
21	MD 1202	Imaging devices utilizing non-ionizing radiation		
22	MD 1301	Monitoring devices of non-vital physiological parameters		
23	MD 1302	Monitoring devices of vital physiological parameters		
24	IVD 0101	AB0 system		
25	IVD 0201	HIV infection (HIV 1 and 2)		
26	IVD 0202	HTLV I and II		
27	IVD 0203	Hepatitis B, C and D		
28	IVD 0303	Congenital infections: rubella, toxoplasmosis		
29	IVD 0305	Human infections: cytomegalovirus, chlamydia		
30	IVD 0307	Tumoral marker: PSA		
31	IVD 0401	Clinical chemistry		
32	IVD 0402	Haematology		
33	IVD 0403	Immunology		
34	IVD 0404	Molecular biology		
35	IVD 0405	Pregnancy and ovulation		
36	IVD 0406	Specimen receptacles		
37	MDS 7002	Medical devices utilizing tissues of animal origin, including Directive 2003/32/EC		
38	MDS 7206	IVDs in sterile condition		
39	MDS 7210	IVDs utilizing material of human origin		

**Conformity Assessment by Way of Verification** 

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\*means approval only for conformity assessment on infusion medical devices.

# CAB REGISTRATION NUMBER: **MDA/CAB-019** VALIDITY: **12/11/2021 - 11/11/2024**

ASI CERTIFICATION SDN. BHD. 1ST FLOOR, NO. 87, JALAN NILAM ½ SUBANG HI-TECH INDUSTRIAL PARK 40000 SHAH ALAM SELANGOR DARUL EHSAN TEL: +603-5621 0358





PERSON RESPONSIBLE: MR. CHANDARASEGARAN A/L ARUMUGAM [carumugam1952@gmail.com]

FAX: +603-5621 0358

CONTACT PERSON: MR. CHANDARASEGARAN A/L ARUMUGAM [carumugam1952@gmail.com]

Con	Conformity Assessment of Quality Management System (QMS) and Post-Market Surveillance System (PMSS)		
1	ISO 13485	Quality Management System for Medical Devices – Requirements for Regulatory Purpose	
2	GDPMD	Good Distribution Practice for Medical Devices	
Con	Conformity Assessment of Technical Documentation		
3	MD 0101	Non-active devices for anaesthesia, emergency and intensive care	

3	MD 0101	Non-active devices for anaesthesia, emergency and intensive care
4	MD 0102	Non-active devices for injection, infusion, transfusion and dialysis
5	MD 0103	Non-active orthopaedic and rehabilitation devices
6	MD 0104	Non-active medical devices with measuring function
7	MD 0106	Non-active instruments
8	MD 0108	Non-active medical devices for disinfecting, cleaning, rinsing
9	MD 1101	Devices for extra-corporal circulation, infusion and haemopheresis
10	MD 1102	Respiratory devices, including hyperbaric chambers for oxygen therapy, inhalation anesthesia
11	MD 1103	Devices for stimulation or inhibition
12	MD 1104	Active surgical devices
13	MD 1105	Active ophthalmologic devices
14	MD 1106	Active dental devices
15	MD 1107	Active devices for disinfection and sterilization
16	MD 1108	Active rehabilitation devices and active prostheses
17	MD 1109	Active devices for patient positioning and transport
18	MD 1201	Imaging devices utilizing ionizing radiation
19	MD 1202	Imaging devices utilizing non-ionizing radiation
20	MD 1302	Monitoring devices of vital physiological parameters
21	MD 1401	Devices utilising ionizing radiation
22	MD 1402	Devices utilising non-ionizing radiation
23	MD 1403	Devices for hyperthermia / hypothermia
24	IVD 0309	Device for self-diagnosis: device for the measurement of blood sugar
25	IVD 0404	Molecular biology
26	IVD 0406	Specimen receptacles
27	MDS 7004	Medical devices referencing the Directive 2006/42/EC on machinery
28	MDS 7005	Medical devices referencing the Directive 89/686/EEC on personal protective equipment (PPE)
29	MDS 7206	IVDs in sterile condition
30	MDS 7210	IVDs utilizing material of human origin

Cor	nformity Assessm	ent by Way of Verification
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## CAB REGISTRATION NUMBER: **MDA/CAB-020** VALIDITY: **04/04/2022 – 03/04/2025**

#### MEDIVICE CERTIFICATION SDN. BHD.

U66-1 RED CARPET AVENUE ENCORP STRAND MALL KOTA DAMANSARA PJU 5/22 47810 PETALING JAYA SELANGOR DARUL EHSAN TEL: +603-6150 4007 FAX: +603-6150 4007



PERSON RESPONSIBLE: DR. UNGKU MOHD SHAHRIN BIN UNGKU MOHD ZAMAN [info@medivice.org.my]

CONTACT PERSON: DR. UNGKU MOHD SHAHRIN BIN UNGKU MOHD ZAMAN [info@medivice.org.my]

Coi	Conformity Assessment of Quality Management System (QMS) and Post-Market Surveillance System (PMSS)		
1	ISO 13485	Quality Management System for Medical Devices – Requirements for Regulatory Purpose	
2	GDPMD	Good Distribution Practice for Medical Devices	

Con	Conformity Assessment of Technical Documentation			
3	MD 0101	Non-active devices for anesthesia, emergency and intensive care		
4	MD 0102	Non-active devices for injection, infusion, transfusion and dialysis		
5	MD 0103	Non-active orthopaedic and rehabilitation devices		
6	MD 0104	Non-active medical devices with measuring function		
7	MD 0105	Non-active ophthalmologic devices		
8	MD 0106	Non-active instruments		
9	MD 0107	Contraceptive medical devices		
10	MD 0108	Non-active medical devices for disinfecting, cleaning, rinsing		
11	MD 0109	Non-active devices for in vitro fertilization (IVF) and assisted reproductive technologies (ART)		
12	MD 0301	Bandages and wound dressings		
14	MD 0302	Suture material and clamps		
15	MD 0303	Other medical devices for wound care		
16	MD 0401	Non-active dental equipment and instruments		
17	MD 1101	Devices for extra-corporal circulation, infusion and haemopheresis		
18	MD 1102	Respiratory devices, including hyperbaric chambers for oxygen therapy, inhalation anaesthesia		
19	MD 1103	Devices for stimulation or inhibition		
20	MD 1104	Active surgical devices		
21	MD 1106	Active dental devices		
22	MD 1108	Active rehabilitation devices and active prostheses		
23	MD 1109	Active devices for patient positioning and transport		
24	MD 1301	Monitoring devices of non-vital physiological parameters		
25	MD 1302	Monitoring devices of vital physiological parameters		
26	MD 1403	Devices for hyperthermia / hypothermia		
27	MD 1404	Devices for (extracorporeal) shock-wave therapy (lithotripsy)		
28	IVD 0201	HIV Infection (HIV 1 And 2)		
29	IVD 0203	Hepatitis B, C and D		
30	IVD 0305	Human infections: cytomegalovirus, chlamydia		
31	IVD 0309	Device for self-diagnosis: device for the measurement of blood sugar		
32	IVD 0401	Clinical chemistry		
33	IVD 0402	Haematology		
34	IVD 0403	Immunology		
35	IVD 0404	Molecular biology		

36	IVD 0405	Pregnancy and ovulation
37	IVD 0406	Specimen receptacles
38	MDS 7005	Medical devices referencing the Directive 89/686/EEC on personal protective equipment
39	MDS 7206	IVDs in sterile condition
40	MDS 7210	IVDs utilizing material of human origin

Conformity Assessment by Way of Verification		
41	VERIFICATION	Conformity Assessment by Way of Verification

\*\* means approval only for conformity assessment on aesthetics medical devices.

#### KIWA INTERNATIONAL CERTIFICATIONS SDN. BHD.

CAB REGISTRATION NUMBER: **MDA/CAB-021** VALIDITY: **04/04/2022 – 03/04/2025** 





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CONTACT PERSON: MS. IRMALISA BINTI SAMSURI [kiwa.auditing@gmail.com]

## **SCOPE OF REGISTRATION**

Сог	Conformity Assessment of Quality Management System (QMS) and Post-Market Surveillance System (PMSS)		
1	ISO 13485	Quality Management System for Medical Devices – Requirements for Regulatory Purpose	
2	GDPMD	Good Distribution Practice for Medical Devices	

Con	Conformity Assessment of Technical Documentation		
3	MD 0102	Non-active devices for injection, infusion, transfusion and dialysis	
4	MD 0104	Non-active medical devices with measuring function	
5	MD 0106	Non-active instruments	
6	MD 0108	Non-active medical devices for disinfecting, cleaning, rinsing	
7	MD 0301	Bandages and wound dressings	
8	MD 0303	Other medical devices for wound care	
9	MD 1101	Devices for extra-corporal circulation, infusion and haemopheresis	
10	MD 1107	Active devices for disinfection and sterilization	

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#### AQC TECHNICAL ASSESSORS (M) SDN BHD.

CAB REGISTRATION NUMBER: **MDA/CAB-022** VALIDITY: **17/06/2021-16/06/2024** 





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# SCOPE OF REGISTRATION

Сог	Conformity Assessment of Quality Management System (QMS) and Post-Market Surveillance System (PMSS)		
1	ISO 13485	Quality Management System for Medical Devices – Requirements for Regulatory Purpose	
2	GDPMD	Good Distribution Practice for Medical Devices	

Conformity Assessment of Technical Documentation		
3	MD 0103	Non-Active Orthopaedic And Rehabilitation Devices
4	MD 0104	Non-Active Medical Devices with Measuring Function
5	MD 0106	Non-active instruments
6	MD 0108	Non-Active Medical Devices for Disinfecting, Cleaning, Rinsing
7	MD 0301	Bandages and Wound Dressings
8	MD 1102	Respiratory Devices, Including Hyperbaric Chambers for Oxygen Therapy, Inhalation Anaesthesia
9	MD 1103	Devices for Stimulation or Inhibition
10	MD 1107	Active devices for disinfection and sterilisation
11	MD 1108	Active Rehabilitation Devices and Active Prostheses
12	MD 1109	Active Devices for Patient Positioning and Transport
13	MD 1402	Devices Utilising Non-Ionizing Radiation
14	IVD 0101	AB0 System
15	IVD 0102	Rhesus (C, C, D, E, E)
16	IVD 0103	Anti-Kell
17	IVD 0201	HIV Infection (HIV 1 And 2)
18	IVD 0202	HTLV I and II
19	IVD 0203	Hepatitis B, C And D
20	IVD 0301	Anti-Duffy And Anti-Kidd
21	IVD 0303	Congenital infections: rubella, toxoplasmosis
22	IVD 0305	Human Infections: Cytomegalovirus, Chlamydia
23	IVD 0307	Tumoral Marker: PSA
24	IVD 0309	Devices for Self-Diagnosis: Device for The Measurement of Blood Sugar
25	IVD 0401	Clinical Chemistry
26	IVD 0402	Haematology
27	IVD 0403	Immunology
28	IVD 0404	Molecular biology
29	IVD 0405	Pregnancy and ovulation
30	IVD 0406	Specimen Receptacles
31	MDS 7004	Medical devices referencing the Directive 2006/42/EC on machinery
32	MDS 7206	IVDs in sterile condition
33	MDS 7207	IVDs utilizing micromechanics

Conformity Assessment by Way of Verification34VERIFICATIONConformity Assessment by Way of Verification

# CAB REGISTRATION NUMBER: **MDA/CAB-023** VALIDITY: **30/08/2022-29/08/2025**





#### NIOSH CERTIFICATION SDN. BHD.

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## SCOPE OF REGISTRATION

Cor	Conformity Assessment of Quality Management System (QMS) and Post-Market Surveillance System (PMSS)		
1	ISO 13485	Quality Management System for Medical Devices – Requirements for Regulatory Purpose	
2	GDPMD	Good Distribution Practice for Medical Devices	

Conformity Assessment of Technical Documentation3MD 0301Bandages and Wound Dressings

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#### PLATINUM SHAUFFMANTZ VERITAS SDN. BHD.

## CAB REGISTRATION NUMBER: **MDA/CAB-024** VALIDITY: **15/08/2023 – 14/08/2026**

Medical Device AUTHORITY MALAYSIA



NO.10, JALAN PENYAJAK U1/45B SEKSYEN U1, TEMASYA GLENMARIE 40150 SHAH ALAM SELANGOR DARUL EHSAN TEL: **+603-5512 9793** FAX: **+603-5518 9793** 

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## **SCOPE OF REGISTRATION**

Со	Conformity Assessment of Quality Management System (QMS) and Post-Market Surveillance System (PMSS)		
1	ISO 13485	Quality Management System for Medical Devices – Requirements for Regulatory Purpose	
2	GDPMD	Good Distribution Practice for Medical Devices	

Con	Conformity Assessment of Technical Documentation		
3	MD 0106	Non-active instruments	
4	MD 0301	Bandages and wound dressings	

# **Conformity Assessment by Way of Verification**

5 VERIFICATION Conformity Assessment by Way of Verification

# LEADER CERTIFICATION SDN. BHD.

CAB REGISTRATION NUMBER: **MDA/CAB-025** VALIDITY: **18/10/2023 - 17/10/2026**  NO 4-3 JALAN SERI PUTRA 1/1 BANDAR SERI PUTRA BANGI 43000 KAJANG SELANGOR DARUL EHSAN TEL: **+603-89127689** FAX: **+603-89127689** 





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## SCOPE OF REGISTRATION

Сог	Conformity Assessment of Quality Management System (QMS) and Post-Market Surveillance System (PMSS)		
1	ISO 13485	Quality Management System for Medical Devices – Requirements for Regulatory Purpose	
2	GDPMD	Good Distribution Practice for Medical Devices	

Con	Conformity Assessment of Technical Documentation		
3	MD 0103	Non-active orthopedic and rehabilitation devices	
4	MD 0106	Non-active instruments	
5	MD 0202	Non-active orthopedic implants	
6	MD 0203	Non-active functional implants	
7	IVD 0309	Device for self-diagnosis: device for the measurement of blood sugar	
8	IVD 0401	Clinical chemistry	
9	IVD 0402	Hematology	
10	IVD 0403	Immunology	
11	IVD 0405	Pregnancy and ovulation	

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< End of List >

Note: Blue-in-colour font means 'new updated information'.

# Section 10(1), Medical Device Act 2012 (Act 737) Regulation 8, Medical Device Regulations 2012

For more enquiries, please contact us:

### **CAB Registration Unit**

Medical Device Authority Ministry of Health Malaysia Aras 5 & 6, Prima 9, Prima Avenue II, Blok 3547, Persiaran APEC, 63000 Cyberjaya, Selangor Tel: **+603 8230 0356 (Mr. Fadhullah) / +603 8230 0372 (Pn. Remee)** Fax: +603 8230 0200 Email: cab.registration@mda.gov.my