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Department of Health and Ageing
Therapeutic Goods Administration

Australian Medical Device Requirements Version 4 under the Therapeutic Goods Act 1989

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TGA Health Safety
Regulation



About the Therapeutic Goods Administration (TGA)

- The TGA is a division of the Australian Government Department of Health and Ageing, and is responsible for regulating medicines and medical devices.
- TGA administers the *Therapeutic Goods Act 1989* (the Act), applying a risk management approach designed to ensure therapeutic goods supplied in Australia meet acceptable standards of quality, safety and efficacy (performance), when necessary.
- The work of the TGA is based on applying scientific and clinical expertise to decision-making, to ensure that the benefits to consumers outweigh any risks associated with the use of medicines and medical devices.
- The TGA relies on the public, healthcare professionals and industry to report problems with medicines or medical devices. TGA investigates reports received by it to determine any necessary regulatory action.
- To report a problem with a medicine or medical device, please see the information on the TGA website.

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TABLE OF CONTENTS

BLOCK 1

1.1	INTRODUCTION	1
1.2	PARALLEL IMPORTING - 'UNAUTHORISED' DISTRIBUTORS	7
1.3	PERSONAL IMPORTATION & DIRECT MARKETING	7
1.4	QUARANTINE REQUIREMENTS - AQIS	8
	AQIS Requirements.....	9
1.5	EXPORT OF THERAPEUTIC DEVICES	10
	Export Only Products.....	10
	Export Certification Scheme	10
1.6	POSTMARKET COMPLIANCE PROGRAMS	12
	Incident Reporting Scheme	12
	Compliance Testing.....	12
	Overseas Regulatory Action.....	12
	Post Market Audit Programs	12
1.7	RECALLS	14
1.8	APPLICATION PROCEDURES	15
	Therapeutic Devices Application Form	15
	Enterprise Details Form	15
	Claims for exemption from registration for clinical trial purposes	16
	Evaluation of Grandfathered Devices	16
	Registrable Device Applications	17
	Listable Device Applications	19
1.9	REDUCING ANNUAL CHARGES	20
	Grouping	20
1.10	CHANGES OR VARIATIONS TO DEVICES IN THE ARTG	21
	Is notification of prior approval required?	21
	Change in status of devices in the ARTG	21
	TGA Business Rules	21
1.11	FEES AND CHARGES	23
1.12	CONFIDENTIALITY	23
1.13	RELEASE OF INFORMATION FROM THE ARTG	24
1.14	ADVERTISING / PROMOTION	25
1.15	THERAPEUTIC GOODS ORDERS (TGO) & STANDARDS	26
1.16	PYROGEN / ENDOTOXIN FREE DEVICES	27

1.17	ELECTRICAL SAFETY	29
	Group 1 Compliance Requirements	29
	Group 2 Compliance Requirements	29
	Electromagnetic Compatibility Requirements	31
	Compliance File.....	32
1.18	MUTUAL RECOGNITION AGREEMENT (MRA) WITH EUROPE.....	33
	How The Agreement Will Work	33
	Requirements for the ARTG	34
	Conformity Assessment of Medical Devices.....	34
1.19	GOOD MANUFACTURING PRACTICE	35
1.20	REJECTION OF AN APPLICATION	39
1.21	APPEALS AGAINST DECISIONS	40
1.22	ENFORCEMENT	40
1.23	PENALTIES AND CANCELLATIONS.....	41
1.24	ACCESS TO UNAPPROVED DEVICES	
	CLINICAL TRIALS & SPECIAL ACCESS SCHEMES (SAS)	43
	Clinical Trial Exemption Scheme (CTE)	45
	Clinical Trial Notification Scheme (CTN).....	47
	Special Access Schemes.....	48
	Individual Patient Use (IPU)	48
	Authorised User Access (AUA).....	50

BLOCK 2 -REGISTRABLE DEVICES

2.1	INFORMATION APPLICABLE TO ALL REGISTRABLE DEVICES.....	53
	Evaluation Fee Structure	53
	Design / Materials / Testing.....	53
	Manufacture/Quality Control/Sterility.....	54
	Biocompatibility/Preclinical	54
	Human Clinical Data.....	54
	Low Level Evaluation Fee.....	54
	Format of Registered Device Submissions.....	54
	General Details	54
	Risk analysis	55
	Table of Equivalence.....	55
	Commercial History, Regulatory Actions and Regulatory Status	55
	Good Manufacturing Practice	56
	Reporting Conditions	56
	Postmarket Surveillance	56
	Samples	56
	Quarantine	56
	Design / Materials / Testing	57
	Design and Construction	57
	Materials	57
	Labelling	57
	Packaging	57
	Product Information / Instructions for use / Promotional Material.....	58
	Manufacturing / Quality Control / Sterility	58
	Manufacturing Process	58
	Quality Control	58
	Sterility	58
	Biocompatibility / Preclinical	59
	Biological Safety and Biocompatibility Testing.....	59
	Preclinical Studies	59
	Human Clinical Data	59
	Submission Content Style	59
	Application Checklist	61
2.2	GROUPING OF REGISTRABLE DEVICES	62
2.3	EQUIVALENCE - ABRIDGED SUBMISSIONS.....	63
	Sponsors who are not manufacturers	63
	Modification of a Registered Device or a New Device.....	64
2.4	OVERSEAS EVALUATIONS	65
	Approvals from Overseas Regulatory Agencies	65
	US FDA Approvals.....	65
	EU Approval (CE mark) B Registrable Device Applications	65
	EU Approval (CE mark) B Listable Device Applications.....	66
	Overseas approvals which can be used to abridge evaluations	67
2.5	RISK ANALYSIS	73
2.6	STERILITY	74
2.7	BIOLOGICAL SAFETY AND BIOCOMPATIBILITY TESTING	78
2.8	HUMAN CLINICAL DATA	81

HIGH LEVEL REGISTRABLE DEVICES.....	83
2.9 ACTIVE IMPLANTABLE MEDICAL DEVICES (AIMD)	85
General Details.....	86
Risk Analysis.....	86
Table of Equivalence	86
Commercial and Regulatory History.....	87
Good Manufacturing Practice	87
Reporting Conditions	87
Postmarket Surveillance.....	87
Samples	87
Labelling.....	87
Design / Construction	88
Materials	88
Testing	89
Qualification Testing.....	89
For Leads and Implantable Accessories:	90
For Implantable Drug Infusion Pumps	90
Manufacturing	91
For Leads and Implantable Accessories:	91
Quality Control	91
Sterility.....	91
Biological Safety and Biocompatibility	91
For Leads and Implantable Accessories:	91
Preclinical Studies	91
2.10 ANIMAL ORIGIN DEVICES	94
General Details.....	94
Risk Analysis.....	94
Table of Equivalence	94
Commercial and Regulatory History.....	94
Good Manufacturing Practice	94
Reporting Conditions	95
Postmarket Surveillance.....	95
Samples	95
Quarantine	95
Manufacturing.....	95
Source Materials and Risk of Infectivity.....	95
Virus Control	95
Control of Agents Causing Spongiform Encephalopathies.....	97
Biological Safety and Biocompatibility Testing.....	98
Testing - Stability Testing and Shelf Life.....	98
2.11 BREAST PROSTHESES (NOT SALINE OR WATER)	101
General Details.....	101
Risk Analysis.....	101
Table of Equivalence	101
Commercial and Regulatory History.....	101
Good Manufacturing Practice.....	101
Reporting Conditions.....	101
Postmarket Surveillance.....	101
Samples.....	101
Design / Construction	102
Materials	102
Product Information / Instructions for Use / Promotional Material	102
Manufacturing - Validation Testing.....	102
Biological Safety and Biocompatibility Testing.....	103
Human Clinical Data	103

2.12	DRUG INFUSION SYSTEMS (POWERED)	106
	General Details.....	106
	Risk Analysis.....	106
	Table of Equivalence.....	107
	Commercial and Regulatory History.....	107
	Good Manufacturing Practice.....	107
	Reporting Conditions.....	107
	Postmarket Surveillance.....	107
	Samples.....	107
	Design and Construction.....	108
	Materials.....	109
	Labelling.....	109
	Packaging.....	109
	Product Information / Instructions for Use / Promotional Material.....	109
	Testing - Qualification Testing - Verification.....	110
	Electrical Safety Testing.....	110
	Electromagnetic Compatibility.....	110
	Human Clinical Data.....	110
2.13	EXTRACORPOREAL THERAPY SYSTEMS	113
	General Details.....	113
	Risk Analysis.....	113
	Table of Equivalence.....	113
	Commercial and Regulatory History.....	113
	Good Manufacturing Practice.....	113
	Reporting Conditions.....	114
	Postmarket Surveillance.....	114
	Samples.....	114
	Design / Construction.....	114
	Labelling.....	114
	Sterility.....	114
	Biological Safety and Biocompatibility Testing.....	114
	Human Clinical Data.....	115
2.14	HEART VALVE PROSTHESES	116
	General Details.....	118
	Risk Analysis.....	118
	Table of Equivalence.....	118
	Commercial and Regulatory History.....	119
	Good Manufacturing Practice.....	119
	Reporting Conditions.....	119
	Postmarket Surveillance.....	119
	Samples.....	120
	Design - Additional Requirements.....	120
	Materials - Component Materials.....	120
	Packaging - Shelf Life and Storage Conditions.....	121
	Product Information / Instructions for Use / Promotional Material.....	122
	Testing - Mechanical Testing and Analysis.....	122
	Valved Conduits.....	125
	Manufacturing Process.....	125
	Sterility.....	126
	Preclinical <i>in vivo</i>	127
	Anticalcification Treatment Studies.....	127
	Human Clinical Data.....	128
2.15	HUMAN ORIGIN DEVICES	131
	General Details.....	132
	Risk Analysis.....	132
	Table of Equivalence.....	132
	Commercial and Regulatory History.....	132
	Good Manufacturing Practice.....	132

	Reporting Conditions.....	132
	Postmarket Surveillance.....	133
	Samples.....	133
	Quarantine	133
	Labelling	133
	Product Information / Instructions for Use / Promotional Material	133
	Manufacturing.....	134
	Source Materials And Risk of Infectivity -.....	134
	Biological Safety and Biocompatibility Testing.....	135
	Stability Testing and Shelf Life.....	135
	Additional Requirements for Tissues of CNS or Ophthalmic Origin.....	136
2.16	INTRAOCULAR LENSES (IOL) - REGISTRABLE	139
	General Details.....	139
	Risk Analysis.....	139
	Table of Equivalence	139
	Commercial and Regulatory History.....	140
	Good Manufacturing Practice.....	140
	Reporting Conditions.....	140
	Postmarket Surveillance.....	140
	Samples.....	140
	Design / Construction	140
	Product Information / Instructions for Use / Promotional Material	141
	Testing - Mechanical Data	141
	Manufacturing Process.....	142
	Biological Safety and Biocompatibility Testing.....	142
	Preclinical	142
	Human Clinical Data	143
	IOLs Not Requiring Clinical Data	144
	Minor Modification Parameters.....	144
2.17	INTRAOCULAR VISCOELASTIC FLUIDS (IOF)	148
	General Details.....	148
	Risk Analysis.....	148
	Table of Equivalence	148
	Commercial and Regulatory History.....	148
	Good Manufacturing Practice.....	148
	Reporting Conditions.....	149
	Postmarket Surveillance.....	149
	Samples.....	149
	Biological Safety/Biocompatibility Testing	149
	Human Clinical Data	149
2.18	INTRAUTERINE CONTRACEPTIVE DEVICES (IUCD).....	152
	General Details.....	152
	Risk Analysis.....	152
	Table of Equivalence	152
	Commercial and Regulatory History.....	152
	Good Manufacturing Practice.....	152
	Reporting Conditions.....	152
	Postmarket Surveillance.....	152
	Samples.....	152
	Design / Construction	153
	Labelling	153
	Product Information / Instructions for Use / Promotional Material	153
	Testing	154
	Manufacturing Process.....	154

LOW LEVEL REGISTRABLE DEVICES	157
2.19 BARRIER CONTRACEPTIVE DEVICES	158
General Details.....	158
Risk Analysis.....	158
Commercial and Regulatory History.....	158
Good Manufacturing Practice.....	158
Reporting Conditions.....	158
Postmarket Surveillance.....	158
Samples.....	158
Design / Construction.....	158
Labelling.....	159
Product Information / Instructions for Use / Promotional Material.....	159
Testing - Chemical and Mechanical Properties.....	159
Biological Safety / Biocompatibility Testing - Stability.....	159
Toxicity.....	160
Barrier Properties to Sexually Transmitted Pathogens.....	160
Human Clinical Data.....	160
2.20 BREAST PROSTHESES (SALINE)	163
General Details.....	163
Risk Analysis.....	163
Table of Equivalence.....	163
Commercial and Regulatory History.....	163
Good Manufacturing Practice.....	163
Reporting Conditions.....	163
Postmarket Surveillance.....	163
Samples.....	164
Design / Construction.....	164
Testing - Integrity of the Shell Material.....	164
Testing - Integrity of the Valve.....	164
Biological Safety and Biocompatibility Testing.....	164
Human Clinical Data.....	165
2.21 DISINFECTANTS AND STERILANTS	167
2.22 HIV/HCV IN VITRO DIAGNOSTIC KITS	169
Conditions of Registration.....	169
Use of Unapproved HIV/HCV Test Kits.....	170
Changes to Products.....	170
General Details.....	171
Risk Analysis.....	171
Table of Equivalence.....	171
Commercial and Regulatory History.....	171
Good Manufacturing Practice.....	171
Reporting Conditions.....	171
Postmarket Surveillance.....	171
Quarantine.....	172
Samples.....	172
Performance Testing.....	172
Stage 1: Performance Data.....	172
Clinical Trial Data.....	172
Stage 2: Preliminary Trials at the NRL.....	172
Stage 3: Evaluation.....	173
Design - Labelling.....	173
Shelf Life.....	173
Product Information / Instructions for Use / Promotional Material.....	173
Manufacturing.....	174
Antigen/Antibody/Primer(s) Reagents.....	174
Control Serum Reagents.....	175
Quality Control Tests.....	175

BLOCK 3 - LISTABLE DEVICES

3.1	INFORMATION APPLICABLE TO ALL LISTABLE DEVICES.....	177
	Information to be Supplied for Listing.....	177
	Enterprise Details Form	178
	Application Form.....	178
	Product Details.....	178
	Promotional Material/Instructions for use/Product Information	178
	Labelling Evidence	179
	Standard & Specific Device Conditions	179
	Variations or Changes to Existing Listings	179
	Licencing / GMP Evidence	180
	Test Certificates.....	180
3.2	GROUPING OF LISTABLE DEVICES.....	182
	What is a separate and distinct listable device product?.....	183
	SPECIFIC LISTABLE DEVICE POLICIES.....	184
3.3	ANIMAL DERIVATIVES CONTAINED IN LISTABLE DEVICES	185
	Gelatin-containing Devices	185
3.4	BANDAGES, DRESSINGS AND ALLIED PRODUCTS	186
	Group 1 - Goods To Be Supplied Sterile.....	186
	Group 2 - Goods Required To Be 'Clean' (Supplied Non-Sterile)	187
	Group 3 - Other Products	189
	Group 4 - Related Products Not Required To Be Listed	189
	Site Information File.....	190
3.5	BARIUM LIME	194
	What to Submit for Listing	194
	Specific Conditions.....	194
	Test Certificates	194
3.6	BLOOD BAGS	195
	What to Submit for Listing.....	195
	Differentiating Between Blood Bag Products.....	195
	Grouping Blood Bags.....	195
3.7	CATHETERS (URETHRAL).....	196
	What to Submit for Listing	196
	Specific Conditions.....	196
	Batch Size	196
	Test Certificates	197
3.8	CONDOMS (CONTRACEPTIVE, RUBBER).....	198
	What to Submit for Listing.....	198
	Test Certificates.....	198
	Condoms which Contain a Spermicide or Similar Agent.....	199
	Condoms in Novelty Packaging	199
	Novelty Products	199
	Monitoring for Compliance with the Standard by the TGA	200
3.9	CONTACT LENSES & CONTACT LENS CARE PRODUCTS.....	202
	What to Submit for Listing.....	202
	How Products/Actions Are Categorised	202
	Contact Lenses	203
	Contact Lens Care Products	203
3.10	CONTRAST MEDIA INJECTORS (POWERED).....	204
	What to Submit for Listing	204
	Specific Conditions.....	204

3.11	DENTAL PRODUCTS	205
	Products Required to be Listed	205
	What to Submit for Listing.....	205
	Dental Products Subject to Standards	205
3.12	DENTAL RESTORATIVE MATERIALS	206
	What to Submit for Listing	206
	Specific Conditions.....	206
	Test Certificates	206
	Labelling	207
	GMP.....	207
	Types and Presentations of Dental Restorative Materials	207
3.13	DEVICES FOR PEOPLE WITH DISABILITIES	209
	Exempt Goods	209
	Excluded Goods.....	209
	Examples of Exempt or Excluded Goods.....	209
3.14	DIAPHRAGMS - CONTRACEPTIVE	211
	What to Submit For Listing	211
	Specific Conditions.....	211
3.15	DISINFECTANTS	212
3.16	DRUG INFUSION INJECTORS	213
	What to Submit for Listing.....	213
3.17	DUCTED AND WIRED SYSTEMS	214
	Electrical Systems.....	214
	Medical Gases	214
	Aspiration Systems	214
	Dialysis Systems.....	214
3.18	ENDOSCOPES AND ENDOSCOPIC ACCESSORIES	215
	What to Submit for Listing.....	215
	Kits	215
	Labels.....	215
	GMP.....	216
	Grouping	216
	Other requirements	216
3.19	GLOVES (EXAMINATION, SURGICAL)	217
	What to Submit for Listing.....	217
	Labelling Requirements	217
	General Requirements.....	218
	Examination Gloves	218
	Presumption of single use.....	218
	Surgeons Gloves	218
	Sterile Gloves.....	218
	Specific Conditions.....	218
	Test Certificate.....	218
3.20	HEARING AND SPEECH AIDS	220
	What to Submit for Listing.....	220
	Electromagnetic Compatability Requirements	221
3.21	IN VITRO DIAGNOSTICS [IVDS] - LISTABLE	222
	Diagnostic instruments and equipment	222
	Diagnostic test kits.....	222
	Antimicrobial test discs.....	222
	What to Submit for Listing.....	223
	Good Manufacturing Practice.....	223
	Specific Conditions.....	223
	Quarantine	224
	Drugs of Dependence	224
	Grouping of Listable IVD'S	225

3.22	NEEDLELESS INJECTORS	226
	What to Submit for Listing.....	226
	Labels.....	226
3.23	INSULIN SYRINGES	227
	What to Submit for Listing.....	227
	Specific Conditions.....	227
	Test Certificates.....	227
3.24	INTRAOCULAR LENSES (IOL) - LISTABLE	228
	IOL's to Which Items 13 and 14 of the Regulations Apply.....	228
	What to Submit for Listing.....	229
	Change in Haptic Configuration, Design or Calibre.....	229
	Comparison to a Single Approved Lens.....	229
	Comparison to Multiple Approved Models.....	233
	Remaining Tests.....	235
3.25	KITS - THERAPEUTIC DEVICES ASSEMBLED AS KITS	236
	Definitions.....	236
	Rules.....	236
	ARTG Procedures.....	239
3.26	ORAL HYGIENE PRODUCTS	240
	Items Declared Not to be Therapeutic Goods.....	240
	Items Regarded as Therapeutic Goods.....	241
	What to Submit for Listing.....	241
	Advertising Claims.....	241
3.27	PENILE IMPLANTS - INFLATABLE	243
	What to Submit for Listing.....	243
	Specific Conditions.....	243
3.28	PODIATRY	244
	What to Submit for Listing.....	244
3.29	SODA LIME	245
	What to Submit for Listing.....	245
	Specific Conditions.....	245
	Test Certificates.....	245
3.30	SUTURES AND LIGATURES (ABSORBABLE & NON-ABSORBABLE)	246
	What to Submit for Listing.....	246
	Specific Conditions.....	246
	Test Certificates.....	246
	Related Products.....	247
3.31	TAMPONS - MENSTRUAL	248
	What to Submit for Listing.....	248
	Packaging, Labelling and Product Information.....	248
	Checklist for Labelling of Tampons.....	248
	Primary Packs.....	249
	Supply Packs.....	249
	Transport Pack.....	250
	Manufacturing Details.....	250
	Information Leaflet.....	250
	ABBREVIATIONS	253
	GLOSSARY	254
	APPENDICES	Refer Volume 2

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