







Article Content

Title	Regulations for Governing the Management of Medical Device 
Amended Date	2017.07.25
Category	Ministry of Health and Welfare (衛生福利部)
Article 1	The present regulations are in accordance to the Article 13, Paragraph 2 of 1 Pharmaceutical Affairs Act.
Article 2	Medical device are classified into the following classes according to their 1 risks: Class I : Low risk Class II : Medium risk Class III : High risk
Article 3	In accordance to the function, intended use, instruction for use and working principle, medical devices are classified into the following categories: 1. Clinical Chemistry and Clinical Toxicology Devices 2. Hematology and Pathology Devices 3. Immunology and Microbiology Device 4. Anesthesiology Devices 5. Cardiovascular Devices 6. Dental Devices 7. Ear, Nose, and Throat Devices 8. Gastroenterology and Urology Devices 9. General and Plastic Surgery Devices 10. General Hospital and Personal Use Devices 11. Neurological Devices 12. Obstetrical and Gynecological Devices 13. Ophthalmic Devices 14. Orthopedic Devices 15. Physical Medicine Devices 16. Radiology Devices 17. Other Categories Specified by the National Health Competent Authority The classification of medical devices into the above categories is as Annex 1  Annex I.pdf
Article 4	The manufacturing of medical devices shall comply with Part 3 Good Manufactu Practice (GMP) for Medical Device of Pharmaceutical Good Manufacturing Practi Regulations. The manufacturing of medical devices listed in the Annex II of this regulatic comply with Part 3 Chapter 3 Essential Mode of the GMP regulation, except the manufacturing of sterilized devices shall comply with Part 3 Chapter 2 Stand of the GMP regulation.  Annex II.pdf
Article 5	Medical devices that require clinical trial to be performed domestically are in Annex III.  Annex III.pdf
Article 6	Pharmaceutical entity or the general public may approach the National Health Competent Authority, through a payment service, for inquiry of the classifi a medical device and its regulatory control with the provision of the followi i. Instruction for use (or catalogue) and its Chinese translation in details (including the instruction for use, function and working principle). ii. Reference to classification of the product by the European Union or Unite of America iii. Any other information as defined by the National Health Competent Author

Article 7 (deleted)

Article 8 The regulations shall be implemented on the date of announcement, except the manufacturing of Optical Impression Systems for CAD/CAM (Classification Number: F.3661) under the amendment of Article 3 Item 2 Annex 1 on January 7th, 2014, shall comply with this regulation from July 1st, 2014.

The manufacturing of Quality Control Material (assayed and unassayed) (Classification Number: A.1660), Ear, Nose, and Throat Drug Administration Device (Classification Number: G.5220), and Corrective Spectacle Lens (Classification Number: M.5844) under the amendment of Article 3 Item 2 Annex 1 on June 3rd, 2016, shall comply with this regulation from September 1st, 2016.