



# **MALAYSIAN STANDARD**

**MS 2752:2022**

## **Prosthetic and orthotic devices - Code of practice**

**ICS: 11.040.01**

Descriptors: medical device, medical electrical equipment, code of practice, biomedical engineering, maintenance, services, active medical device, testing, commissioning, acceptance

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## Committee representation

The National Standards Committee on Medical Devices and Facilities for Healthcare (NSC R) under whose authority this Malaysian Standard was developed, comprises representatives from the following organisations:

Association of Malaysian Medical Industries  
Association of Private Hospitals of Malaysia  
Atomic Energy Licensing Board  
Biomedical Engineering Association Malaysia  
Department of Standards Malaysia (Secretariat)  
Federation of Malaysian Manufacturers  
Institute for Medical Research  
Institute of Physics Malaysia (Radiation Physics, Biophysics and Medical Physics Sub-Group)  
Malaysia Medical Device Association  
Malaysian Association of Standards Users  
Malaysian Medical Association  
Malaysian Nuclear Agency  
Malaysian Organisation of Pharmaceutical Industries  
Malaysian Rubber Board  
Malaysian Rubber Council  
Medical Device Authority, Ministry of Health Malaysia  
Pharmaceutical Association of Malaysia  
SIRIM QAS International Sdn Bhd  
Universiti Kebangsaan Malaysia (Faculty of Allied Health Sciences)  
Universiti Teknologi Malaysia (Faculty of Science)

The Technical Committee on Code of Practice for Medical Devices and Healthcare Facilities (TC/R/10) which supervised the development of this Malaysian Standard consists of representatives from the following organisations:

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### **Foreword**

This Malaysian Standard was developed by the Technical Committee on Code of Practice for Medical Devices and Healthcare Facilities (TC/R/10) under the authority of the National Standards Committee on Medical Devices and Facilities for Healthcare (NSC R).

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