







List of Reference Countries

These are the regulatory authorities in the reference countries where the **Free Sale Certificate OR Registration Certificate** should be issued including the manufacturer name, product name and stating that the product is classified as a medical device under the directive 93/42/EEC of medical devices.

Regulatory Authority & Reference countries	Emails	Logo on the FSC
Saudi Food and Drug SFDA (Saudi Arabica)	md.rs@sfda.gov.sa	
US Food and Drug FDA (USA)	Medical Devices FDA	
Therapeutic Goods Administration TGA (Australia)	TGA.International@health.gov.au	
Canada's Food and Drugs Act and Regulations (Canada)	Canada's Food and Drugs Act and Regulations - Canada.ca	
Pharmaceuticals and Medical Devices Agency PMDA (Japan)	Regulatory Science/The Science Board/Standard Development Pharmaceuticals and Medical Devices Agency (pmda.go.jp)	



the New Zealand Medicine and Medical Devices Safety Authority MEDSAFE (New Zealand)	device@health.govt.nz	
Swissmedic (Switzerland)	fsc@swissmedic.ch	
Medicines & Healthcare products Regulatory Agency (United Kingdom)	Medicines and Healthcare products Regulatory Agency - GOV.UK (www.gov.uk) devices.regulatory@mhra.gov.uk	
ANSM / CCI Paris (France)	clv@cci-paris-idf.fr	
The Health Product Regulatory Authority HPRA (Ireland)	devices@hpra.ie	
Health and Youth Inspectorate. (Netherland (Holland))	Medical Technology Health and Youth Care Inspectorate (igj.nl) meldpunt@igj.nl	
Danish Health and Medicines Authority (DHMA) (Denmark)	Medical devices (laegemiddelstyrelsen.dk) Send an email	
Federal Agency for Medicines and Health Products (FAMHP) (Belgium)	FAMHP Your medicine and health products welcome@fagg-afmps.be	