



Medical Device Regulatory Update

Medical Device Control Office
Department of Health
The Government of Hong Kong SAR

5 January 2017



www.mdco.gov.hk



Matters related to Legislative Proposal

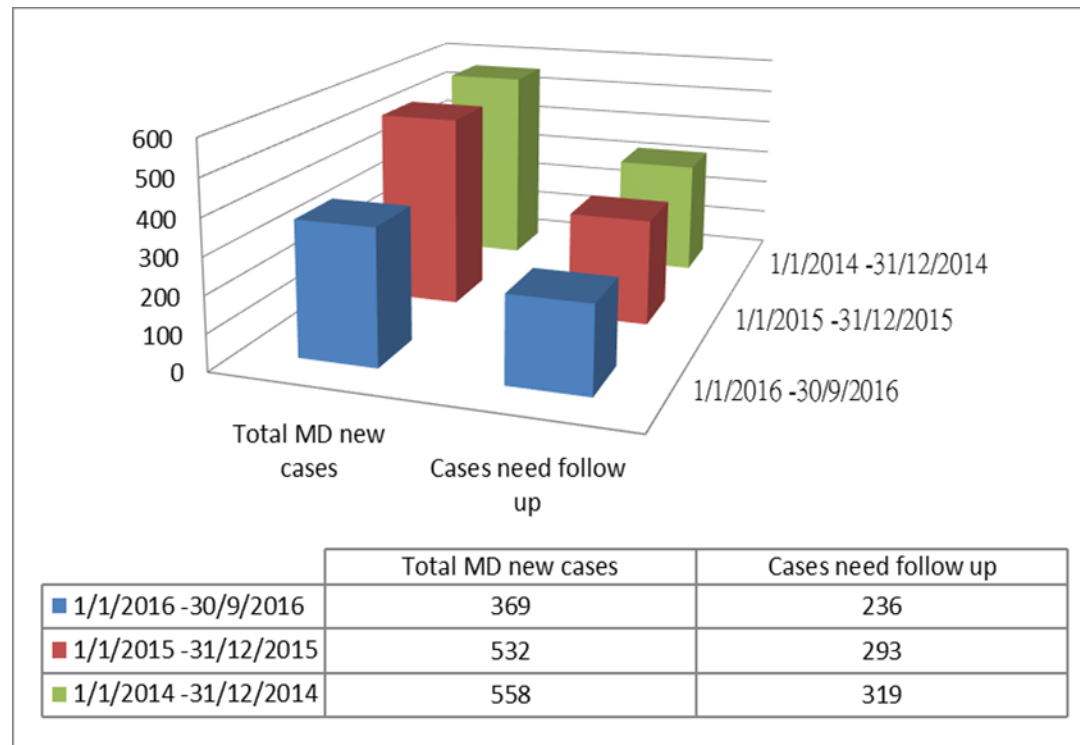
- The consultancy study by the ECRI Institute on the control of use of selected medical devices in Hong Kong has been completed. The consultant's recommendations and the finalised regulatory proposal of medical devices are scheduled to be reported at the Health Panel meeting of the LegCo on 16 Jan 2017.





MD Listing under MDACS

Result of Initial Screening



MD Listing under MDACS



Change - Renewal of listed equipment implemented on 1 January 2017

- Mainly an administrative process, through provision of essential document (business registration & etc)
- Involve a declaration on application details
- Need to submit the overdue change application within a short time period if applicable
- 'Change' will be handled separately and to be submitted when situation arise



MD

Medical Device Control Office Renewal Form for Listed Medical Devices

To: Medical Device Control Office

For official use only
Date Received (dd/mm/yyyy): _____
Application No.: _____

S



Particulars of Application	
1	HKMD No.
	Make
	Model
	Company Name
	Contact Person
	Telephone
	E-mail
2	A copy of valid Business Registration Certificate (number: _____) is enclosed.
3	<u>There are active recalls, field safety corrective actions or adverse incidents (local and worldwide)</u> <input type="checkbox"/> No <input type="checkbox"/> Yes. Details are provided in separate sheets.
4	<u>There is change to the listing details since last approval</u> <input type="checkbox"/> No <input type="checkbox"/> Yes (Please go to 4(i))
	(i) <u>Change application of the devices is submitted to MDCO</u> <input type="checkbox"/> Yes <input type="checkbox"/> No. We will submit change application within 10 calendar days.
Declaration	
We acknowledge that the listing details of this renewal application would be the same as the last approved version. Approval of this renewal application does not imply the approval of any change application in progress.	
We confirm that:	
a. The applicant remains designated as Local Responsible Person by the manufacturer;	
b. The applicant remains aware and complies with all device listing conditions (e.g. Post market surveillance);	
c. All certifications / licences (e.g. ISO 13485 certificate for manufacturing site and recognized marketing approval) relating to the listed device remain valid and will be submitted to MDCO upon request; and	
d. The information contained in this renewal form is true and correct.	
Signature of Applicant:	

Name: _____	
Position: _____	
Date (dd/mm/yyyy): _____	
_____ Company Chop	

MD Listing under MDACS



Possible change

Change in Importer requirement

- Evaluation of the capability of the medical devices manufacturer
- Evaluation of the safety, efficacy and quality of the medical devices to be imported
- Incoming and outgoing goods inspection



MD Listing under MDACS



Possible change

Addition of intended use at website of MDCO.

What is the limitation?:

- length
- style of writing
- promotion (world smallest)
- discrimination (replacing the previous)



Traders under MDACS



Office Address

- Request for information on all relevant addresses of LRP is in progress
- Request for information on all relevant addresses of other traders will follow





Thank you !

