



## Announcement

**Subject:** Invitation to One Day Industry Awareness Programme on “Understanding ICMED Certification and New Medical Device Rules, 2017 for Medical Device Manufacturers”

**ICMED:** Regulation of medical devices and their quality systems in India has been limited when compared to other developed nations that typically have very robust medical device approval systems in place.

To help address this gap, the Association of Indian Medical Device Industry (AIMED) in collaboration with the Quality Council of India (QCI) established a voluntary quality certification scheme ICMED for medical devices in India, with two levels of certification – ICMED 9000 and ICMED 13485.

**MDR 2017:** The Ministry of Health and Family Welfare has notified Medical Device Rules, 2017 on 31.01.2017 for regulating manufacturing/import/sale/clinical investigation and classification of medical devices.

The new rules have been framed in conformity with Global Harmonization Task Force (GHTF) framework and conform the best international practices. The new rules seek to remove regulatory bottlenecks to Make in India, facilitate ease of doing business while ensuring availability of better medical devices for patient care and safety.

We invite participants from interested manufacturing industry, medical devices professionals and other stakeholders to register to participate in this one day programme.

**Please Note:** There is no registration fee for this event.

**Venue:** Coimbatore on 9 August 2017

**Venue:** Vadodara on 16 August 2017

Please confirm your nomination by submitting filled-in registration form as per the nomination form attached to Ms. Ajita Srivastava (ajita@qcin.org).



**One day Industry Awareness Programme on “Understanding ICMED Certification  
and New Medical Device Rules, 2017 for Medical Device Manufacturers”**

**Venue: Dr. N.G.P Institute of Technology, Seminar Hall,  
Dr. N.G.P. Nagar, Kalapatti Road, Coimbatore - 641048  
on 09<sup>th</sup> August 2017**

Please complete and return this form **on or before 4<sup>th</sup> August 2017**

By email or fax\* to  
**Ms. Ajita Srivastava**  
QCI Secretariat

**Email:** [ajita@qcin.org](mailto:ajita@qcin.org), **Tel:** +91-11-23378056/57; **Fax** +91-11-23378678

\*Sending by email is preferable. Please print clearly if sending by fax

**NOMINATION FORM**

**A. PERSONAL PARTICULARS**

**Title (please tick)**                      Mr.     Mrs.     Ms.     Dr.

Name	:		
Organization	:		
Position / Designation	:		
Organization Address	:		
City		Postal Code / Zip	
State		Telephone (O)	
Fax		Telephone (R)	
Email Address		Mobile	

**Note:**

- No registration fee is applicable.
- Registration will be done on First-come-first-serve basis.



**One day Industry Awareness Programme on “Understanding ICMED Certification  
and New Medical Device Rules, 2017 for Medical Device Manufacturers”**

**Venue: Hotel Hampton by Hilton Vadodara-Alkapuri,  
14, Friends Co-op Society, Vadodara, Gujarat 390007  
on 16<sup>th</sup> August 2017**

Please complete and return this form **on or before 11<sup>th</sup> August 2017**

By email or fax\* to  
**Ms. Ajita Srivastava**  
QCI Secretariat

**Email:** [ajita@qcin.org](mailto:ajita@qcin.org), **Tel:** +91-11-23378056/57; **Fax** +91-11-23378678

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Organization	:		
Position / Designation	:		
Organization Address	:		
City		Postal Code / Zip	
State		Telephone (O)	
Fax		Telephone (R)	
Email Address		Mobile	

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