REPORT HIGHLIGHTS

Ali Al dalaan, MBA-IT,PRA,QMS-LA Vice Executive President, Medical Devices Sector

AHWP Technical committee Chair



Asian Harmonization Working Party WORKING TOWARDS MEDICAL DEVICE HARMONIZATION IN ASIA

Current AHWP Membership

AHWP Member Country or Region: 30 (as of Mar 2018)

Brunei Darussalam

Cambodia

Chile

Chinese Taipei

Hong Kong SAR, China

India

Indonesia

Jordan

Kazakhstan

Kingdom of Bahrain

Kingdom of Saudi Arabia

Republic of Korea

Laos

Malaysia

Mongolia

Myanmar

Pakistan

People's Republic of China

Philippines

Republic of Kenya

Singapore

South Africa

State of Kuwait

Sultanate of Oman

Tanzania

Thailand

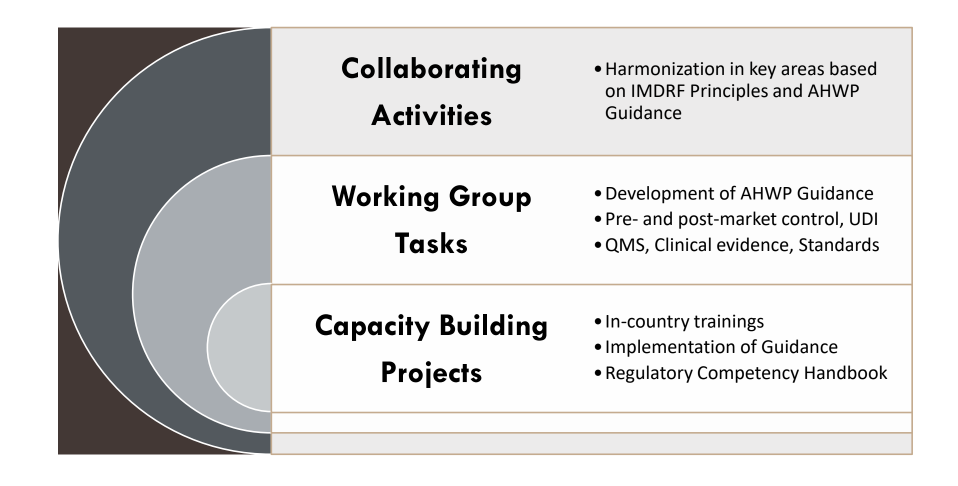
United Arab Emirates

Vietnam

Yemen

Zimbabwe

AHWP TC Strategic Plan



TC Team

TC Office Bearers	Positions
Chair	Mr Ali M Al-Dalaan
Co-Chair	Dr Jeong-Rim Lee
Co-Chair	Mr Alfred Kwek
Secretary	Mr Jack Wong
	Ms Chadaporn Tanakasemsub (Miang)
	Ms Carol Yan
	Ms Soo-Kyeong Shin
Work Groups	Positions
WG1: Pre-market	Chair - Mr. Se-il Park
	Co-Chair - Ms. Kate Hyeong Joo Kim
WG2: Pre-market - IVDD	Chair - Mr. Wen-Wei TSAI
	Co-Chair - Ir. Albert POON
WG3: Pre-market - Software as a Medical Device	Chair - Dr. Abdullatif Alwatban
	Co-Chair - Mr Tony Yip
WG4: Post-market	Chair - Ms. Jennifer MAK
Scope includes post-market aspect of WG 1-3 device categories	Co-Chair - Ms Kitty Mao
WG5: Clinical Evidence for performance & safety	Chair - Ms. Yuwadee PATANAWONG
	Co-Chair - Ms. Sumati Randeo
WG6: Quality Management Systems:	Chair - Mr. Abdullah AL RASHEED
Audit & assessment	Co-Chair - Mr. Vincent LAM Chee-Choong
WG7: Quality Management Systems:	Chair - Ms. Wang Aijun
Operation & implementation	Co-Chair - Mr. Ee Bin Liew
WG8: Standards	Chair - Mrs. Salibiah Yaakop
	Co-Chair – Mr Tony Low
STC (UDI & Nomenclature)	Chair - Mr. Jun Ll
	Co-Chair – Ms Victoria Ou

AHWP TC PLAN

2018 - 2020

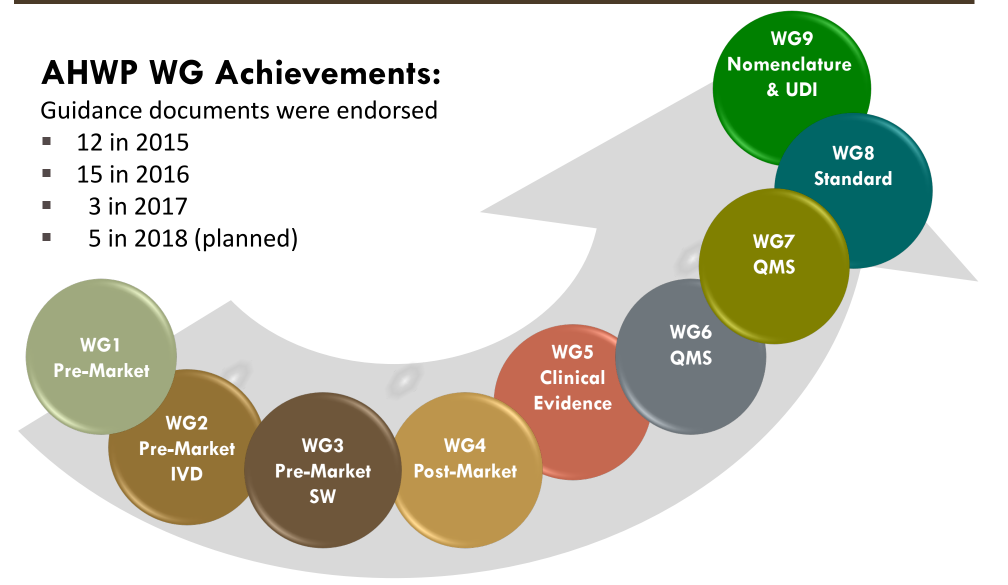
WG Plans for 2018 - 2020 (1)

WG	Tasks	Timeline
WG1	 E-labeling/e-IFU guideline (collaboration with WG2 & WG3) 3D printing handbook update Change management for medical device registration guideline (collaboration with WG2 & WG3) 	Q4, 2018 TBD Q4, 2019
WG2	 E-labeling/e-IFU guideline (collaboration with WG1 and WG3 Change management for medical device registration guideline (collaboration with WG1 & WG3) Guidance document for approval of reagent for instrument family Future trend study & survey: Bridging LDT and IVD 	Q4, 2018 Q4, 2019 Q4, 2020 Q4, 2020
WG3	 White paper on pre-market initial submission format for SaMD E-labeling/e-IFU guideline (collaboration with WG2 & WG3) White paper on cybersecurity for SaMD Change management for medical device registration guideline (collaboration with WG2 & WG3) Guidance document for pre-market submission format for SaMD (draft) 	Q4, 2018 Q4, 2018 Q1, 2019 Q4, 2019
WG4	 Updating the post-market resource centre Gap analysis on the implementation of AHWP guidance among AHWP members Participation in the development works of ISO TC210/WG6 	Ongoing Ongoing Ongoing

WG Plans for 2018 - 2020 (2)

WG	Tasks	Timeline
WG5	 Annual review SWOT analysis of WG5 framework Guidance document on general principles of clinical investigation audit & inspection for medical devices Training: WG5 & AHWP members Survey: country regulations/guidelines and implementation 	Q4, 2018 Q4, 2018 Q4, 2018 Q4, 2019
WG6	 Guidance document on understanding the roles of IMDRF documents concerning auditing (draft) Guidance document on the current best practice in determination of regulatory audit duration (draft) 	Q4, 2018 Q2, 2019
WG7	 Comparison study of new ISO13485 vs QMS requirements in each country QMS consideration for manufacturers and importers for localization 	Q2, 2020 Q4, 2020
WG8	 Guidance document on code of practice for good engineering maintenance management of medical devices Collecting a list of standards used for medical device regulatory purposes that are recognized by AHWP member countries 	TBD TBD
WG9	AHWP UDI reportAHWP UDI rule	TBD TBD

Development & Implementation of AHWP Guidance



Continuous Efforts for Global Harmonization



APEC LSIF RHSC/ Medical Device Vigilance

- Join the Project 'Roadmap to Promote Convergence' and training workshops



IMDRF WG/ UDI & Standards

- Join the International Workshop on UDI, Feb 2018, Brussels
- Participated IMDRF meeting in March, Shanghai, September Beijing



IMDRF WG/ Personalized Medical Devices

- Attended IMDRF face to face meeting for Personalized Medical Devices
- * Personalized Medical Devices definitions N49 is approved by MC
- * Now working on another documents for Personalized Medical Devices conformity pathways





IEC/ISO Works

- Drafting: Committees of ISO14971, ISO TR24971, ISO/IEC Guide63, ISO TR20416
- Attending TC meetings: ISO TC210















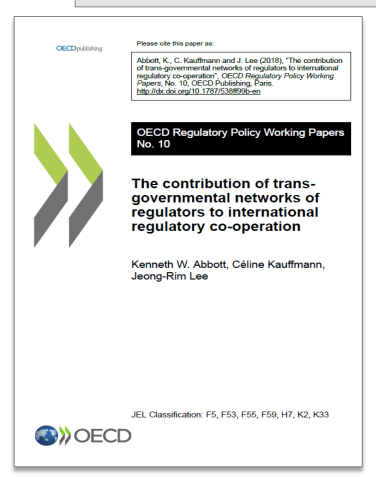






Collaboration with the OECD

Title: The Contribution of Trans-Governmental Networks of Regulators to International Regulatory Co-operation



A Case Study of the AHWP on Medical Devices		
1. Overview	 History Intended objectives of regulatory co-operation Landscape of regulatory actors Collaboration with other IOs 	
2. Governance & Operational Modalities	 AHWP Membership Structure and governance Institutional setup The range of AHWP instruments Implementation mechanism (CBP) Quality mechanism of instruments 	
3. Assessment	- Benefits - Challenges	

- Participation in drafting the 2nd OECD Report (2017 2018)
- Published as an OECD report (September, 2018)

AHWP TC Leaders Meeting in 2018

May 8th - 9th 2018, Beijing, China

- AHWP Technical Committee
 Short-term & long-term Plans
 - Guideline topics and development plans by each WG
 - Development of Competency Handbook by AHWP TC
 - In-country training plans
 - Introduction of OECD case study





AHWP Annual Meeting Plans

- Participation in global events
 (IMDRF, WHO, APEC, OECD, etc)
- Joint workshop plans with liaisons
- Meeting program agenda
- Progress of AHWP website update

AHWP Capacity Building Projects

3 Capacity Building Workshops & 4 In-country Trainings (2015-2017)

- CB Workshops: Thailand Nov'15; Philippines Nov'16; India Dec'17
- In-country Trainings: Indonesia '16; Vietnam '16; Malaysia '17; Kazakhstan '17
- Topics: CSDT for pre-market registration submission, Risk classification, Good distribution practice, QMS audit, SW, Information technology, Post-market considerations

2018



- In-country trainings
- Republic of Kenya (TBD)
- Thailand





Deloitte.

Launch Competency Framework for MedTech Regulators

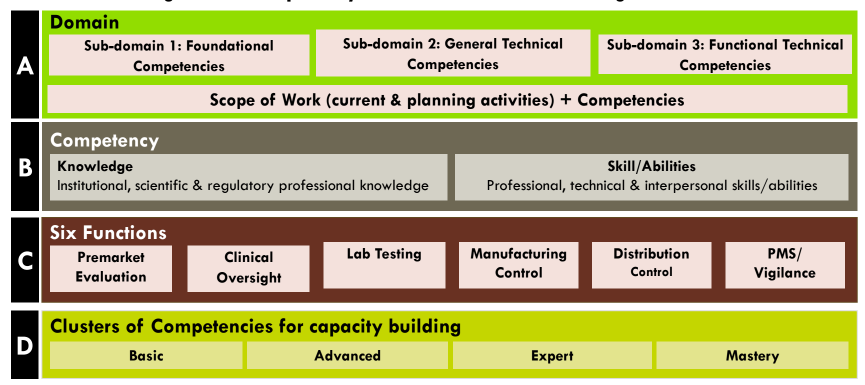
A joint initiative of AHWP, APACMed and Deloitte

Competency Handbook for Medtech Regulators

PROJECT SCOPE:

- AHWP survey for regulators among its 30 member countries and regions
- APACMed launching similar survey among companies to assess satisfaction & expectation

High-Level Competency Framework for MedTech Regulators



Thank you