

## **AHWP UPDATE**

Ali M. Aldalaan

#### Vice Executive President, Medical Devices Sector AHWP Chair

IMDRF - 15. 19-21 March 2019 Moscow, Russia



### **Current AHWP Membership**

#### AHWP Member Country or Region: 30 (as of Mar 2019)

Brunei Darussalam	Kingdom of Saudi Arabia	Singapore	
Cambodia	Republic of Korea	South Africa	
Chile	Laos	State of Kuwait	
Chinese Taipei	Malaysia	Sultanate of Oman	
Hong Kong SAR, China	Mongolia	Tanzania	
India	Myanmar	Thailand	
Indonesia	Pakistan	United Arab Emirates	
Jordan	People's Republic of	Vietnam	
Kazakhstan	China	Yemen	
Kingdom of Bahrain	Philippines	Zimbabwe	
Republic of Kenya			

## AHWP TC Strategic Plan

	Collaborating	<ul> <li>Harmonization in key areas based on IMDRF Principles and AHWP</li> </ul>	
	Activities	Guidance	
	Working Group	<ul> <li>Development of AHWP Guidance</li> <li>Pre- and post-market control, UDI</li> </ul>	
	Tasks	• QMS, Clinical evidence, Standards	
	Capacity Building	<ul> <li>In-country trainings</li> <li>Implementation of Guidance</li> </ul>	
	Projects	Regulatory Competency Handbook	

## **AHWP Leadership Team**

#### AHWP Chair: **Mr. Ali Al dalaan,**

Vice Executive President, Medical Devices Sector, Saudi Arabia

#### AHWP Vice-chair:

#### Mr. Guobiao Gao,

Deputy Director General, National Medical Product Administration, China

AHWP Vice-chair: Ms. Tran Quan, Industry

## AHWP TC Team

TC Office Bearers	Positions
Chair	Ms.Sasikala Devi Thangavelu
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

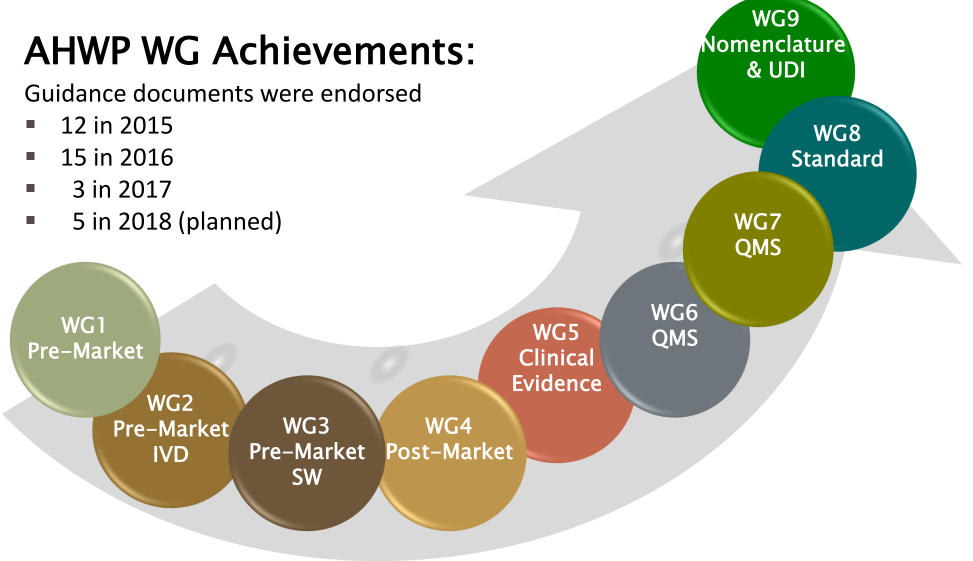
## WG Plans for 2018 - 2020 (1)

WG	Tasks	Timelin e
WG1	<ul> <li>E-labeling/e-IFU guideline (collaboration with WG2 &amp; WG3)</li> <li>3D printing handbook update</li> <li>Change management for medical device registration guideline (collaboration with WG2 &amp; WG3)</li> <li>Al guidance in consideration</li> </ul>	Q4, 2018 TBD Q4, 2019 TBD
WG2	<ul> <li>E-labeling/e-IFU guideline (collaboration with WG1 and WG3</li> <li>Change management for medical device registration guideline (collaboration with WG1 &amp; WG3)</li> <li>Guidance document for approval of reagent for instrument family</li> <li>Future trend study &amp; survey: Bridging LDT and IVD</li> </ul>	Q4, 2018 Q4, 2019 Q4, 2020 Q4, 2020
WG3	<ul> <li>White paper on pre-market initial submission format for SaMD</li> <li>E-labeling/e-IFU guideline (collaboration with WG2 &amp; WG3)</li> <li>White paper on cybersecurity for SaMD</li> <li>Change management for medical device registration guideline (collaboration with WG2 &amp; WG3)</li> <li>Guidance document for pre-market submission format for SaMD</li> </ul>	Q4, 2018 Q4, 2018 Q1, 2019 10

## WG Plans for 2018 - 2020 (2)

WG	Tasks	Timeline
WG5	<ul> <li>Annual review SWOT analysis of WG5 framework</li> <li>Guidance document on general principles of clinical investigation audit &amp; inspection for medical devices</li> <li>Training: WG5 &amp; AHWP members</li> <li>Survey: country regulations/guidelines and implementation</li> </ul>	Q4, 2018 Q4, 2018 Q4, 2018 Q4, 2018 Q4, 2019
WG6	<ul> <li>Guidance document on understanding the roles of IMDRF documents concerning auditing (draft)</li> <li>Guidance document on the current best practice in determination of regulatory audit duration (draft)</li> </ul>	Q4, 2018 Q2, 2019
WG7	<ul> <li>Comparison study of new ISO13485 vs QMS requirements in each country</li> <li>QMS consideration for manufacturers and importers for localization</li> </ul>	Q2, 2020 Q4, 2020
WG8	<ul> <li>Guidance document on code of practice for good engineering maintenance management of medical devices</li> <li>Collecting a list of standards used for medical device regulatory purposes that are recognized by AHWP member countries</li> </ul>	TBD TBD 11

### Development & Implementation of AHWP Guidance



## **Continuous Efforts for Global Harmonization**



## Collaboration with the OECD

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

OECD publishing	Please cite this paper as:	A Case St	udy of the AHWP on Medical	]
	Abbott, K., C. Kauffmann and J. Lee (2018), "The contribution of trans-governmental networks of regulators to international regulatory co-operation", <i>OECD Regulatory Policy Working</i> <i>Papers</i> , No. 10, OECD Publishing, Paris. http://dx.doi.org/10.1787/536f99b-en			
	nttp://dx.doi.org/10.1767/536039b-en		– Intended objectives of	
	OECD Regulatory Policy Working Papers No. 10		regulatory	
	NO. 10		co-operation	
	The contribution of trans- governmental networks of		- Landscape of regulatory	
	regulators to international		actors	
	regulatory co-operation		- Collaboration with other IOs	
	Kenneth W. Abbott, Céline Kauffmann, Jeong-Rim Lee	2.	– AHWP Membership	1
		Governance	- Structure and governance	
		&	– Institutional setup	
		Operational	– The range of AHWP	
		Modalities	instruments	
	JEL Classification: F5, F53, F55, F59, H7, K2, K33		– Implementation mechanism	
			(CBP)	
			– Quality mechanism of	
Participatio	on in drafting the 2 <sup>nd</sup> OEC	D Report (2017 -	z់ពូនុទ្សរយាents	
Published	as an OECD report (Septem	1ber <u>32</u> 018)	– Benefits	
		Assessment	– Challenges	3

### 23<sup>rd</sup> AHWP Annual Meeting October 22–25, 2018, Kuala Lumpur, Malaysia



- AHWP Annual Meeting
  - Participation of global organizations
    - (IMDRF, WHO, APEC, OECD, etc)
  - Joint workshop plans with liaisons
  - Strategy for Improvement of Regulatory Capacity, Enforcement



- 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

## **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 d istribution practice, QMS audit, SW, Information technology, Post-market conside rations

# 2018



- In-country trainings
- Republic of Kenya
- Thailand



Launch Competency Framework for MedTech Regulators

A joint initiative of AHWP, APACMed and Deloitte

#### AHWP Capacity Building Bangkok, Thailand Feb 22–23, 2019



## Capacity Building in Thailand

#### *Day 1:*

- ✓ AMDD- Areas of harmonization in place in ASEAN states- summary update
- ✓ Definition of a Medical Device (Recap)
- ✓ Risk Management
- $\checkmark$  Risk classification and grouping
- ✓ CSDT
- ✓ Discussion on 1 application for product registration received by TFDA (Optional)

• 35 Regulators attended training

### Day 2:

- Essential Principles what and why this is necessary
- ✓ Standards application of standards in the context of Essential Principles
- ✓ Post Market (articles in AMDD)
- ✓ Areas of Post market to be applied.



## Capacity Building in Thailand

• Training met my expectation?

~92% Strongly Agree and Agree

Part I: AMDD Summary				
Strongly Agree	Agree	Disagree	Strongly Disagree	No Response
15	6	1	0	0
Part II: Premarket				
Strongly Agree	Agree	Disagree	Strongly Disagree	No Response
13	8	0	0	1
Part III: Post market				
Strongly Agree	Agree	Disagree	Strongly Disagree	No Response
13	6	1	0	2

Note: 22 attendees participated after-training feedback

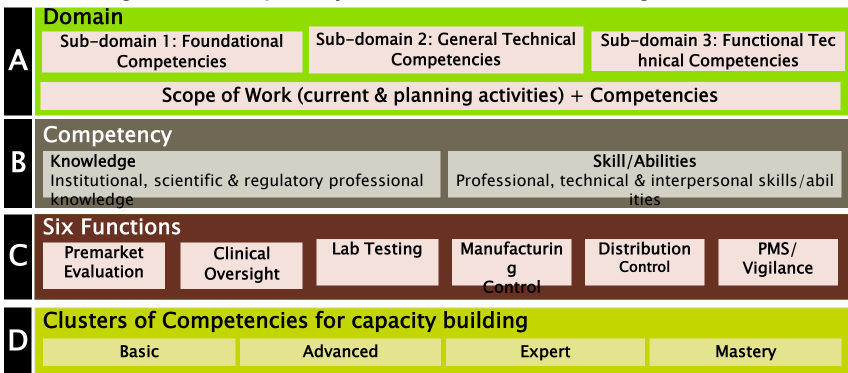
### Competency Handbook for Medtech Regulators

#### **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