WG3 – Pre-market: Software as a Medical Device (SaMD)

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AHWP TC Meeting 7 Dec 2017, New Delhi





Updates

• No. of WG members: 12 (Exclude Chair & Co-chair)



Proposed Work Plan 2015 - 2017

Priority	Work Item	Deliverables	Action Plan and Timeline	
1	Guidance document on Qualification of Medical Device Software The White paper on this topic that was prepared by the earlier WG1 will be the foundation for this. The appropriate aspects from the recent IMDRF document on Software as Medical Device (SaMD)will be kneaded with the existing white paper to develop this AHWP document	Guidance document	Q3 2015	
2	Risk Classification of Medical Device Software / SaMD — To draw reference from the IMDRF SaMD workgroup and also to develop a AHWP document with adequate examples to illustrate and clarify on risk classification of software MDs	Guidance document	Q1 2016 (First Draft)	



Proposed Work Plan 2015 - 2017

Priority	Work Item	Deliverables	Action Plan and Timeline
3	 White paper / Position paper on Pre-market initial Submission format for SaMD To draw up a white paper or position paper for AHWP TC covering the pre-market submission format for SaMD highlighting the need for considering approaches different from those in practice for traditional MDs 	White paper / Position paper	Q4 2016 (First draft)



WG Achievement and Progress Update

No.	Work Item	Status	Achievements
1	Guidance document on Qualification of Medical Device Software	Completed	Endorsed in the AHWP Annual Meeting in 2015
2	Guidance document on Risk Categorisation of Software as a Medical Device	Completed	Endorsed in the AHWP Annual Meeting in 2016
3	White paper on SaMD Pre- market Submission Requirement	Working Draft ready for public consultation	Consolidation of regulatory requirement completed. Working draft ready for public consultation.
4	Proposed new item: White paper on Cyber Security for SaMD	First draft in Q2 2018 (Review of the new standards in progress)	
5	White paper on SaMD change management – Requirements and Processes	First draft in Q2 2018	



White paper on SaMD Pre-market Submission Requirement

- Working draft ready for public consultation

Scope of document:

This document provides a snap shot of the pre-market submission requirements for some regulatory bodies and jurisdictions such as Australia TGA, China CFDA/CMDE, the European Union, Health Canada, Korea MFDS, MHLW Japan and the US FDA. The information collated is with reference to their published guidelines for medical software regulation and pre-market submission requirements.



White paper on SaMD Pre-market Submission Requirement

Objective of document:

The main aim of this white paper is to summarize the current regulatory environment around the world, by including the harmonized view on pre-market submission requirement across jurisdictions, for next development of AHWP guidelines which can serve as member economies' key reference in establishing in a consistent way, an economic and effective approach to the control of medical software in the interest of public health and in the continued innovation of medical software development.

 Highly evolving area with new guidelines and standards constantly being developed or revised

Summaries of SaMD specific Pre-Market Submission Requirements (Subject to consultation and approval) **US FDA** EU Health **KR MFDS** Doc \ Economy Japan **Australia China CFDA** Canada TGA 'ION **Level of Concern** Yes Ν No SaMD Yes Yes No Ν **Software Description including Platform** SaMD Yes Yes Yes Specific Yes Yes and Operation Environment specific Guidance **Device Hazard Analysis / Risk Assessment** Guidanc Yes issued Yes Yes Yes Yes e issued **Software Requirement Specifications** Yes Ν Ν Yes Yes (SRS) **Architecture Design Chart** Yes (Not required Ν Ν Yes Yes for Minor Concern) Yes (Not required **Software Design Specification (SDS)** Ν Ν Yes Yes for Minor Concern)

Traceability Analysis	Yes		N	N		Yes	Yes
Software Development Environment	Yes (Not required		N	N		Yes	Yes
Description	for Minor Concern)						
Verification & Validation Documentation	Yes		Yes	Yes		Yes	Yes
Revision level History	Yes		N	N		Yes	Yes
Unresolved Anomalies (Bugs or Defects)	Yes (Not required		N	N		Yes	N
	for Minor Concern)						
Software Configuration Management	N		N	N		N	Yes
Medical Device - Software Development	N		N	Yes.		Yes.	Yes. IEC62304
Life Cycle (SDLC) standards				IEC62304 /		IEC62304 /	
				JIS T 2304		YY/T 0664	
Other Non-SaMD specific requirements							
Instruction for use	Yes	Yes	Yes	Yes	Yes	Yes	Yes
Intended Use & Indication for Use	Yes	Yes	Yes	Yes	Yes	Yes	Yes
Contradictions	Yes	Yes	Yes	N	Yes	Yes	Yes
Market History	Not mentioned	Yes	Yes	N	Yes	Yes	N
Clinical Evaluations / Trial / Studies	Yes	Yes	Yes	Yes	Yes	Yes	Yes
Labeling	Yes	Yes	Yes	Yes	Yes	Yes	Yes
Essential Principal / Essential	N	Yes	Yes	Yes	Yes	N	N
Requirements							8



White paper on SaMD Pre-market Submission Requirement

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Thank You