

# Manufacturing and Quality Management System (QMS)

18 - 19 May 2026

#### **WORKSHOP PROGRAMME**

#### **Learning Outcomes**

At the end of this workshop, participants should be able to:

- Describe key regulatory requirements of In Vitro Diagnostic Devices (IVDDs) throughout the product life cycle
- Explain the conformity assessment methods by major regulatory authorities
- Describe the key considerations in change management and post-market activities of IVDDs

#### **Target Audience**

• Regulators, Industry QA/RA professionals, product developers, entrepreneurs and investors with interest in diagnostics, and Academia

#### Pre-reads

- 1. Total Product Lifecycle & Introduction to Risk Management
- 2. Introduction to Regulatory Controls & QMS
- 3. Project Management





## Manufacturing and Quality Management System

### 18-19 May 2026

#### Day 1 (18 May 2026)

Time	Agenda	Speaker/Facilitator
8.00 am	Registration	
8.30 am	Welcome Address	Asst Prof James Leong Head, Health Products & Regulatory Science Centre of Regulatory Excellence (CoRE) Duke-NUS
8.40 am	Introduction on the Singapore Standardisation Programme	Standards Development Organisation (SDO) CoRE, Duke-NUS
9.00 am	Workshop Briefing	Ms Janet Poh Training Manager QARA DxD Hub
9.15 am	Photo taking session	
9.20 am	Introduction to Diagnostics Development Hub	Ms Ho Yuan Lu Vice President Project & Outreach DxD Hub
9.45 am	Dossier preparation and Product registration: IVD and SaMD	Mr Ang Wei Jun Senior Manager QARA DxD Hub
10.45 am	Break	
11.00 am	Manufacturing Transfer and Batch Records: IVD and SaMD	Dr Lin You Bin Senior Manager Molecular Dx DxD Hub  Mr Henry Johnson Susainathan Principal Manager Digital Dx DxD Hub
11.45 am	Case Discussion     Medical Device File (Resolute®/ Inspecta)	Ms Tan Hwee Ee Founder and Director DH RegSys Consulting Pte Ltd
12.30 pm	Lunch	
1.30 pm	QMS application in a Medical Device Organization (MDO)	Ms Tan Hwee Ee DH RegSys Consulting Pte Ltd

<sup>\*</sup>The Programme is accurate as of 11 December 2025 and may be subjected to further refinement, if necessary, before the actual workshop.





2.30 pm	Determining Sites and facility requirements: IVD and SaMD	Dr Lin You Bin Mr Henry Johnson Susainathan DxD Hub
3.15 pm	Break	
3.30 pm	Manufacturing and QMS Case Study: Resolute®	Dr Lin You Bin DxD Hub  Ms Tan Hwee Ee DH RegSys Consulting Pte Ltd
4.45 pm	Knowledge Check	Ms Janet Poh DxD Hub
5.15 pm	Wrap Up and Lessons Learnt	Ms Janet Poh DxD Hub
5.30 pm	End of Day 1	

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#### Day 2 (19 May 2026)

Time	Agenda	Speaker/Facilitator
8.00 am	Registration	
8.30 am	Process validation for scale-up Manufacturing: IVD	<b>Dr Lin You Bin</b> DxD Hub
9.15am	Process validation for scale-up Manufacturing: SaMD	<b>Mr Henry Johnson Susainathan</b> DxD Hub
10.00 am	Tea Break	
	HSA Post-market Regulatory Requirements	Ang Wei Jun DxD Hub
11.00 am	Manufacturing and QMS Case Study: Inspecta	Mr Henry Johnson Susainathan DxD Hub  Ms Tan Hwee Ee DH RegSys Consulting Pte Ltd
12.30 pm	Lunch	
1.30 pm	Gallery Walk Customer related processes: IVD and SaMD	Ms Tan Hwee Ee DH RegSys Consulting Pte Ltd
3.30 pm	Tea break	
3.45 pm	End-of-Course Assessment	Ms Janet Poh DxD Hub
4.45 pm	Closing remarks with Lessons Learnt	Ms Janet Poh DxD Hub
5.30 pm	End of Day 2	