

Update from the World Health Organization







Prequalification of IVDs

PQ listing of the first triple diagnostic test for ANC PQ listing of the first HIV ST by urine

PQ expansion to haemoglobin point-of-care analysers

A new guidance document supporting the post PQ and EUL change request application process has been published and came into force on 1 June 2025

• webinar recording Passcode: 7YK=s?3=

Launch of pilot project for parallel PQ assessment and policy development

Launch of the ePQS portal





WHO Emergency Use Listing for MPXV IVDs

Continued support to MPOX PHEIC response:

- 6 NATs listed, including PoC
- Ag RDTs not eligible for EUL as not recommended by WHO

Date Listed	Manufacturer name	Product name	Product code(s)	Regulatory version	EUL application number	Packaging	Link to access public report
3 October 2024	Abbott Molecular Inc	Alinity m MPXV assay (Alinity m MPXV Amplification (AMP) Kit & Alinity m MPXV Control (CTRL) Kit)	AMP kit: 09R06-095 CTRL kit: 09R06-085	USFDA EUA	MPXV-12644-027-00	Alinity m MPXV Amplification (AMP) Kit: 48 tests/tray, 192 tests/kit Alinity m MPXV Control (CTRL) Kit: 12 tubes per level	Available here
14 October 2024	Roche Molecular Systems, Inc.	cobas MPXV Qualitative assay for use on the cobas 6800/8800 Systems	cobas MPXV: 09863338190 Control kit: 09863320190 Buffer Negative Control Kit: 07002238190, 09051953190	USFDA EUA	MPXV-12647-046-00	cobas MPXV: 192 test cassette Control Kit: 1 x 16 mL Buffer Negative Control Kit: 1 x 16 mL	Available here
28 October 2024	Cepheid	Xpert Mpox	GXMPX-10	USFDA EUA	MPXV-12646-070-00	10 Xpert Mpox Cartridges with Integrated Reaction Tubes Disposable 300 µL Transfer Pipettes: Two bags of 12 per kit	Available here
06 March 2025	KH Medical Co., Ltd	RADIONE Mpox Detection Kit	RP001	Rest of the World	MPXV-13227-214-00	Pre-filled PCR Tube: 4 well- strip tube X 24 EA	Available here
09 May 2025	UStar Biotechnologies Ltd	EasyNAT Monkeypox Virus Assay	U202028-20	CE-Marked	MPXV-13201-208-00	20 cartridges, 1 positive control tube (1.2mL), 1 negative control tube (1.2 mL)	To be published
13 May 2025	UStar Biotechnologies Ltd	PortNAT Monkeypox Virus Test	U202031-20	CE-Marked	MPXV-13202-208-00	Box A: 20 cartridge-A, 20 swabs, 20 zip-lock bag Box B: 20 cartridge-B	To be published





New learning catalogue to upskill regulatory workforce

Health products regulation and prequalification learning catalogue

- WHO has launched a comprehensive learning resource to help countries build stronger systems for overseeing the safety, quality and effectiveness/performance of health products
- includes self-paced e-learning courses, instructor-led training sessions and on-the-job training experiences





Diagnostics coalition and definition

On diagnostics (including IVD and non IVD) Definition and Coalition

Background

- The 76th World Health Assembly on 30 May 2023, approved the WHA76.5 resolution named "<u>Strengthening</u> diagnostics capacity."
- The resolution included a footnote 1:
- "For the purpose of this resolution, the term "diagnostics" includes medical devices used for the diagnosis, screening, monitoring, prediction, staging or surveillance of diseases or health conditions, both in vitro and non-in vitro types."
- · Requested:

(12) to develop and/or update WHO definitions of diagnostics, through a group of experts and public consultations, and to publish revised definitions before the 156th session of the Executive Roard:

Process for definition

1st phase: collect information or definitions, 2024

2nd phase: subgroup of SAGE IVD and STAGMEDEV and WHO Dx.

3rd phase: NSA and consultation in April –May 2025. Address the 75 comments received.

4th phase: to conclude, reviewing all comments. by Q3

Outcome definition (26- August-2025)

- Diagnostics are those medical devices* intended by the manufacturer to be used, either in vitro or non in vitro, alone or in combination, for human beings, for providing information for one or more of the specific medical purposes:
- Diagnosis or monitoring of disease, Investigation; diagnosis, aid to diagnosis, screening, monitoring, predisposition, prognosis, prediction or determination of physiological status.
- *In alignment with the International Medical Device Regulators Forum as: (IMDRF1): Essential Principles of Safety and Performance of Medical Devices and IVD Medical Devices, IMDRF/GRRP WG/N47 FINAL:2024 (Edition 2) published 26 April 2024.





WHO managed network to provide advocacy for strengthened diagnostics capacity

https://www.who.int/initiatives/global-diagnostic-coalition





Medical devices nomenclature

Nomenclature medical devices, reported in the World Health Assembly WHA78.

Using EMDN and GMDN in WHO databases on going updating and expansion of list

In the Medical Devices information system, version 2.1







In the Essential in vitro diagnostics list

HO MeDevIS	MeDevPacks Essential in vitro Diagnostics (EDL) ICD-11 UHCC
World Organi	Health zation
	MEDEVPACKS Medical Devices Packages
Search by keyw	vord or click "enter" to display available filters
Pack types	
	les: A package of single-use medical items needed for one procedure.
Consumable	
Consumabl Kit : Kits are usefintend	e a collection of products, including medical devices, that are packaged together to achieve a con of purpose and are being distributed as medical devices.
Kit : Kits arruse/intend	e a collection of products, including medical devices, that are packaged together to achieve a come of purpose and are to being distributed as medical devices. The product of the produc

In Medical devices packages

https://medevis.who-healthtechnologies.org/

Glucose-6- phosphate dehydrogenase activity (G6PD)

Faecal immunochemical test (FIT)





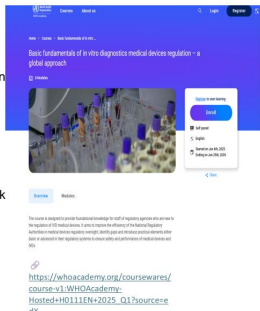
Training materials for IVDs assessors

Finalization of training materials for assessment of IVDs technical files –beginner's courses

New WHO Training Course Launched!

This course provides a foundational overview of IVD regulation, designed for regulators and all stakeholders involved in the lifecycle of in vitro diagnostics.

- The course includes six self-paced modules:
- ✓ An Overview of IVDs: A Regulatory Perspective
- ✓ Quality Management System and Risk Management Devices
- ✓ Performance Studies for IVDs
- √ Labelling Requirements
- ✓ IVD Market Approvals
- ✓ Post-Market Surveillance
- Strengthen your understanding of global regulatory principles and best practices in IVDs.



■ Validation of training materials on assessment of IVDs technical files

- WHO piloted training materials on IVD dossier assessment at the WHO Academy with assessors from 12 NRAs.
- ✓ 23 to 26 July 2025
- ✓ The materials have 17 modules covering key regulatory elements emphasizing the scientific principles necessary for assessing IVD safety and performance.
- ✓ The materials are aligned with WHO and International Medical Device Regulators Forum including use of Table of Contents for compiling IVDs technical.

✓ Finalization in Q4/2025







Supporting manufacturers and regulators for post-market and market surveillance

Regulators

- Renewed focus to strengthen national capacities using WHO global benchmarking tool for medical devices
- In September, WHO will support 7 African countries to self-benchmark themselves against these regulatory functions
 - National Regulatory System (RS);
 - Post-market surveillance, market surveillance and control (PS)
 - Licensing establishments (LI)
- Link to Evaluation of national regulatory systems of medical devices (GBT+ Medical devices)

Industry

- From October 2025, WHO Global Surveillance and Monitoring System for substandard/falsified medical products (GSMS) will allow manufacturers to directly report incidents and FSCA through the GSMS portal
- Portal onboarding will include lessons on IMDRF adverse event reporting terminology





GBT+MD and technical support

Capacity Strengthening:

 Renewed focus on building national regulatory capacity through application of the WHO Global Benchmarking Tool for Medical Devices (GBT+MD) and evaluation of regulatory systems.

WHO-Listed Authorities (WLAs):

 Active discussions across the three levels of WHO to define applicability, criteria, and steps towards listing NRAs as WLAs for medical devices.

WHO CIP Network:

 Recommendation from the 19th ICDRA to extend scope of support to include medical devices.

Link:

 Evaluation of national regulatory systems of medical devices (GBT+ Medical devices)

NATIONAL:

Senegal

- Activity (Sept 2025): Assisted Self-Benchmarking mission
- Objective: Conduct assisted self-benchmarking of Senegal ARP using the WHO GBT+MD
- **Expected Outcome:** Completed self-benchmarking report with documented strengths, gaps, and updated IDP recommendations for further support.

REGIONAL:

African Region

- Activity (Sept 2025): WHO Regional Assisted Self-Benchmarking mission on Medical Device Regulation (GBT+MD) in Africa.
- **Objective:** Build NRA capacity to assess regulatory systems for medical devices with a focus on RS, LI & PS functions.
- **Expected outcome:** Identified strengths, gaps, and priority actions documented in IDPs to guide follow-up support.

Region of the Americas

- Activity (Sept 2025): Webinar on Medical Device Regulation & GBT+MD.
- Objective: Strengthen NRA capacity using WHO frameworks and tools.
- Outcome: Enhanced knowledge of GBT+MD, regulatory frameworks & surveillance.





Upcoming events: UN meeting with manufacturers

- 2025 Joint UNICEF-UNFPA-WHO Meeting with Manufacturers and Suppliers
- 29 September to 3 October 2025. Sessions will be held daily from 11:00 AM to 03:00 PM (CEST)
- Delivering in times of Disruption: Building resilient, quality supply systems for sustainable impact
- Fully virtual event



Thank you!

